Learn R

As a Language

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The R language: "Words" and "sentences"

The desire to economize time and mental effort in arithmetical computations, and to eliminate human liability to error, is probably as old as the science of arithmetic itself.

Howard Aiken *Proposed automatic calculating machine*, 1937; reprinted 1964

1.1 Aims of this chapter

In my experience, for those not familiar with computer programming languages, the best first step in learning the R language is to use it interactively by typing textual commands at the *console* or command line. This will teach not only the syntax and grammar rules, but also give you a glimpse at the advantages and flexibility of this approach to data analysis.

In the first part of the chapter we will use R to do everyday calculations that should be so easy and familiar that you will not need to think about the operations themselves. This easy start will give you a chance to focus on learning how to issue textual commands at the command prompt.

Later in the chapter, you will gradually need to focus more on the R language and its grammar and less on how commands are entered. By the end of the chapter you will be familiar with most of the kinds of "words" used in the R language and you will be able to write simple "sentences" in R.

Along the chapter, I will occasionally show the equivalent of the R code in mathematical notation. If you are not familiar with the mathematical notation, you can safely ignore it, as long as you understand the R code.

1.2 Natural and computer languages

Computer languages have strict rules and interpreters and compilers are unforgiving about errors. They will issue error messages, but in contrast to human readers or listeners, will not guess your intentions and continue. However, computer languages have a much smaller set of words than natural languages, such as English. If you are new to computer programming, understanding the parallels between computer and natural languages may be useful.

One can think of constant values and variables (values stored under a name) as nouns and of operators and functions as verbs. A complete command, or statement, is the equivalent of a natural language sentence: "a comprehensible utterance." The simple statement a+1 has three components: a, a variable, +, an operator and 1 a constant. The statement sqrt(4) has two components, a function sqrt() and a numerical constant 4. We say that "to compute $\sqrt{4}$ we *call* sqrt() with 4 as its *argument*."

Although all values manipulated in a digital computer are stored as *bits* in memory, multiple interpretations are possible. Numbers, letters, logical values, etc., can be encoded into bits and decoded as long as their type or mode is known. The concept of class is not directly related to how values are encoded when stored in computer memory, but instead their interpretation as part of a computer program. We can have, for example, RGB color values, stored as three numbers such as 0, 0, 255, as hexadecimal numbers stored as characters #0000FF, or even use fancy names stored as character strings like "blue". We could create a class for colors using any of these representations, based on two different modes: numeric and character.

In this chapter we will focus on individual program statements, the equivalent of sentences in natural language. In later chapters you will learn how to combine them to create compound statements, the equivalent of natural-language paragraphs, and scripts, the equivalent of essays. You will also learn how to define new verbs, user-defined functions and operators, and new nouns, user-defined classes.

1.3 Numeric values and arithmetic

When working in R with arithmetic expressions, the normal mathematical precedence rules are respected, but parentheses can be used to alter this order. Parentheses can be nested, but in contrast to the usual practice in mathematics, the same parenthesis symbol is used at all nesting levels.

Both in mathematics and programming languages *operator precedence rules* determine which subexpressions are evaluated first and which later. Contrary to primitive electronic calculators, R evaluates numeric expressions containing operators according to the rules of mathematics. In the expression $3 + 2 \times 3$, the product 2×3 has precedence over the addition, and is evaluated first, yielding

as the result of the whole expression, 9. In programming languages, similar rules apply to all operators, even those taking as operands non-numeric values.

It is important to keep in mind that in R trigonometric functions interpret numeric values representing angles as being expressed in radians.

The equivalent of the math expression

$$\frac{3+e^2}{\sin\pi}$$

is, in R, written as follows:

```
(3 + exp(2)) / sin(pi)
## [1] 8.483588e+16
```

It can be seen above that mathematical constants and functions are part of the R language. One thing to remember when translating complex fractions as above into R code, is that in arithmetic expressions the bar of the fraction generates a grouping that alters the normal precedence of operations. In contrast, in an R expression this grouping must be explicitly signaled with additional parentheses.

If you are in doubt about how precedence rules work, you can add parentheses to make sure the order of computations is the one you intend. Redundant parentheses have no effect.

```
1 + 2 * 3

## [1] 7

1 + (2 * 3)

## [1] 7

(1 + 2) * 3

## [1] 9
```

The number of opening (left side) and closing (right side) parentheses must be balanced, and they must be located so that each enclosed term is a valid mathematical expression, i.e., code that can be evaluated to return a value, a value that can be inserted in place of the expression enclosed in parenthesis before evaluating the remaining of the expression. For example, (1 + 2) * 3 after evaluating (1 + 2) becomes 3 * 3 yielding 9. In contrast, (1 +) 2 * 3 is a syntax error as 1 + is incomplete and does not yield a number.

Here results are not shown. These are examples for you to type at the command prompt. In general you should not skip them, as in many cases, as with the statements highlighted with comments in the code chunk below, they have something to teach or demonstrate. You are strongly encouraged to *play*, in other words, create new variations of the examples and execute them to explore how R works.

```
1 + 1
2 * 2
2 + 10 / 5
(2 + 10) / 5
10^2 + 1
sqrt(9)
pi # whole precision not shown when printing
print(pi, digits = 22)
sin(pi) # oops! Read on for explanation.
log(100)
log10(100)
log2(8)
exp(1)
```

Variables are used to store values. After we *assign* a value to a variable, we can use in our code the name of the variable in place of the stored value. The "usual" assignment operator is <-. In R, all names, including variable names, are case sensitive. Variables a and A are two different variables. Variable names can be long in R although it is not a good idea to use very long names. Here I am using very short names, something that is usually also a very bad idea. However, in the examples in this chapter where the stored values have no connection to the real world, simple names emphasize their abstract nature. In the chunk below, a and b are arbitrarily chosen variable names; I could have used names like my.variable.a or outside.temperature if they had been useful to convey information.

```
a <- 1

a + 1

## [1] 2

a

## [1] 1

b <- 10

b <- a + b

b

## [1] 11

3e-2 * 2.0

## [1] 0.06
```

Entering the name of a variable *at the R console* implicitly calls function print() displaying the stored value on the console. The same applies to any other statement entered *at the R console*: print() is implicitly called with the result of executing the statement as its argument.

```
a ## [1] 1

print(a) ## [1] 1

a + 1 ## [1] 2

print(a + 1) ## [1] 2
```

There are some syntactically legal assignment statements that are not very frequently used, but you should be aware that they are valid, as they will not trigger error messages, and may surprise you. The most important thing is to write code consistently. The "backwards" assignment operator \rightarrow and resulting code like 1 \rightarrow a are valid but less frequently used. The use of the equals sign (=) for assignment in place of \leftarrow although valid is discouraged. Chaining assignments as in the first statement below can be used to signal to the human reader that a, b and c are being assigned the same value.

```
a <- b <- c <- 0.0
a
b
c
1 -> a
a
a = 3
```

In R, all numbers belong to mode numeric (we will discuss the concepts of *mode* and *class* in section 1.8 on page 26). We can query if the mode of an object is numeric with function is.numeric().

```
mode(1)
## [1] "numeric"

a <- 1
is.numeric(a)
## [1] TRUE</pre>
```

Because numbers can be stored in different formats, requiring different amounts of computer memory per value, most computing languages implement several different types of numbers. In most cases R's numeric() can be used everywhere that a number is expected. However, in some cases it has advantages to explicitly indicate that we will store or operate on whole numbers, in which case we can use class integer, with integer constants indicated by a trailing capital "L," as in 32L.

```
is.numeric(1L)
## [1] TRUE

is.integer(1L)
## [1] TRUE

is.double(1L)
## [1] FALSE
```

Real numbers are a mathematical abstraction, and do not have an exact equivalent in computers. Instead of Real numbers, computers store and operate on numbers that are restricted to a broad but finite range of values and have a finite resolution. They are called, *floats* (or *floating-point* numbers); in R they go by the name of double and can be created with the constructor double().

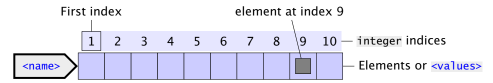
```
is.numeric(1)
## [1] TRUE

is.integer(1)
## [1] FALSE

is.double(1)
## [1] TRUE
```

The name double originates from the C language, in which there are different types of floats available. With the name double used to mean "double-precision floating-point numbers." Similarly, the use of L stems from the long type in C, meaning "long integer numbers."

R's vectors are one-dimensional arrays used to store similar values, e.g., numbers. They are not equivalent to the vectors, commonly used in Physics for describing directional forces, which are symbolized with an arrow as an "accent," such as $\vec{\mathbf{F}}$. In R numeric values and other atomic values are always vector s that can contain zero, one or more elements. The diagram below exemplifies a vector containing ten elements. These elements can be extracted using integer numbers as positional indices, and manipulated as described in more detail in section 1.12 on page 43.



Vectors, in mathematical notation, are similarly represented using positional indexes as subscripts,

$$a_{1...n} = a_1, a_2, \dots a_i, \dots, a_n,$$
 (1.1)

where $a_{1...n}$ is the whole vector and a_1 its first member. The length of $a_{1...n}$ is n as it contains n members. In the diagram above n = 10.

As you have seen above, the results of calculations were printed preceded with [1]. This is the index or position in the vector of the first number (or other value) displayed at the head of the current line. As single values are vectors of length one, when they are printed, they are also preceded with [1].

One can use c() "concatenate" to create a vector from other vectors, including vectors of length 1, or even vectors of length 0, such as the numeric constants in the statements below. We show R code followed by a diagram depicting the vector created.

```
a <- c(3, 1, 2)
a
## [1] 3 1 2

1 2 3 — integer indices
```

numeric values

3

```
b \leftarrow c(4, 5, 0)
## [1] 4 5 0
                 1
                     2
                          3 — integer indices
                     5
                          0
                 4
                                 numeric values
c \leftarrow c(a, b)
## [1] 3 1 2 4 5 0
                                    5
                                         6 — integer indices
                 1
                     2
                          3
                 3
                          2
                               4
                     1
                                    5
                                               numeric values
d \leftarrow c(b, a)
## [1] 4 5 0 3 1 2
                                         6 — integer indices
                 1
                     2
                          3
                                    5
                                4
                      5
                                         2
                                               numeric values
                 4
                          0
                               3
                                    1
```

In the examples above we saved the values returned by the R statements in named variables to reuse them and to help identify the matching diagrams. As shown earlier, values can be also printed at the R console.

```
c(d, numeric()) # concatenate d with a vector of length zero
## [1] 4 5 0 3 1 2
```

Method c() accepts as arguments two or more vectors and concatenates them, one after another. Quite frequently we may need to insert one vector in the middle of another. For this operation, c() is not useful by itself. One could use indexing combined with c(), but this is not needed as R provides a function capable of directly doing this operation. Although it can be used to "insert" values, it is named append(), and by default, it indeed appends one vector at the end of another.

```
append(a, b)
## [1] 3 1 2 4 5 0
```

The output above is the same as for c(a, b), however, append() accepts as an argument an index position after which to "append" its second argument. This results in an *insert* operation when the index points at any position different from the end of the vector.

```
append(a, values = b, after = 2L)
## [1] 3 1 4 5 0 2
```

Both c() and append() can also be used with lists (described in section 1.13 on page 49).

One can create sequences using function seq() or the operator:, or repeat values using function rep(). In this case, I leave to the reader to work out the rules by running these and his/her own examples, with the help of the documentation, available through help(seq) and help(rep).

```
a <- -1:5
a
b <- 5:-1
b
c <- seq(from = -1, to = 1, by = 0.1)
c
d <- rep(-5, 4)
d</pre>
```

Next, something that makes R different from most other programming languages: vectorized arithmetic. Operators and functions that are vectorized accept, as arguments, vectors of arbitrary length, in which case the result returned is equivalent to having applied the same function or operator individually to each element of the vector.

```
a + 1 # we add one to vector a defined above
## [1] 4 2 3

(a + 1) * 2
## [1] 8 4 6

a + b
## [1] 7 6 2

a - a
## [1] 0 0 0
```

As it can be seen in the first line above, another peculiarity of R, is what is frequently called "recycling" of arguments: as vector a is of length 6, but the constant 1 is a vector of length 1, this short constant vector is extended, by recycling its value, into a vector of six ones—i.e., a vector of the same length as the longest vector in the statement, a.

Make sure you understand what calculations are taking place in the chunk above, and also the one below.

```
a <- rep(1, 6)
a
## [1] 1 1 1 1 1 1

a + 1:2
## [1] 2 3 2 3 2 3

a + 1:3
## [1] 2 3 4 2 3 4

a + 1:4

## Warning in a + 1:4: longer object length is not a multiple of shorter object length
## [1] 2 3 4 5 2 3</pre>
```

As mentioned above, a vector can have length zero. Vectors of length zero may seem at first sight quite useless, but in fact they are very useful. They allow the handling of "no input" or "nothing to do" cases as normal cases, which in the absence of vectors of length zero would require to be treated as special cases. Constructors for built in classes like numeric() return vectors of a length given by their first argument, which defaults to zero. I describe here a useful function, length() which returns the length of a vector or list.

```
z <- numeric(0)
z
## numeric(0)
length(z)
## [1] 0

z1 <- numeric()
z1
## numeric(0)

z2 <- numeric(length = 0)
z2
## numeric(0)</pre>
```

Vectors and lists of length zero, behave in most cases, as expected—e.g., they can be concatenated as shown here.

```
length(c(a, numeric(0), b))
## [1] 9
length(c(a, b))
## [1] 9
```

Many functions, such as R's maths functions and operators, will accept numeric vectors of length zero as valid input, returning also a vector of length zero, issuing neither a warning nor an error message. In other words, *these are valid operations* in R.

```
log(numeric(0))
## numeric(0)

5 + numeric(0)

## numeric(0)
```

Even when of length zero, vectors do have to belong to a class acceptable for the operation: 5 + character(0) is an error.

It is possible to *remove* variables from the workspace with rm(). Function 1s() returns a *list* of all objects visible in the current environment, or by supplying a pattern argument, only the objects with names matching the pattern. The pattern is given as a regular expression, with [] enclosing alternative matching characters, ^ and \$, indicating the extremes of the name (start and end, respectively). For

example, " z " matches only the single character 'z' while " z " matches any name starting with 'z'. In contrast " $^[zy]$ " matches both 'z' and 'y' but neither 'zy' nor 'yz', and " $^[a-z]$ " matches any name starting with a lowercase ASCII letter. If you are using RStudio, all objects are listed in the Environment pane, and the search box of the panel can be used to find a given object.

```
ls(pattern="^z$")
## [1] "z"

rm(z)
ls(pattern="^z$")
## character(0)
```

There are some special values available for numbers. NA meaning "not available" is used for missing values. Calculations can also yield the following values NAN "not a number", Inf and -Inf for ∞ and $-\infty$. As you will see below, calculations yielding these values do **not** trigger errors or warnings, as they are arithmetically valid. Inf and -Inf are also valid numerical values for input and constants.

```
a <- NA
a
## [1] NA
-1 / 0
## [1] -Inf

1 / 0
## [1] Inf

Inf / Inf
## [1] NaN

Inf + 4
## [1] Inf

b <- -Inf
b * -1
## [1] Inf
```

Not available (NA) values are very important in the analysis of experimental data, as frequently some observations are missing from an otherwise complete data set due to "accidents" during the course of an experiment. It is important to understand how to interpret NA's. They are simple placeholders for something that is unavailable, in other words, *unknown*.

```
A <- NA
A
## [1] NA
A + 1
## [1] NA
A + Inf
## [1] NA
```

When to use vectors of length zero, and when NAS? Make sure you understand the logic behind the different behavior of functions and operators with respect to NA and numeric() or its equivalent numeric(0). What do they represent? Why NAS are not ignored, while vectors of length zero are?

```
123 + numeric()
123 + NA
```

Model answer: NA is used to signal a value that "was lost" or "was expected" but is unavailable because of some accident. A vector of length zero, represents no values, but within the normal expectations. In particular, if vectors are expected to have a certain length, or if index positions along a vector are meaningful, then using NA is a must.

Any operation, even tests of equality, involving one or more NA's return an NA. In other words, when one input to a calculation is unknown, the result of the calculation is unknown. This means that a special function is needed for testing for the presence of NA values.

```
is.na(c(NA, 1))
## [1] TRUE FALSE
```

In the example above, we can also see that is.na() is vectorized, and that it applies the test to each of the two elements of the vector individually, returning the result as a logical vector of length two.

One thing to be aware of are the consequences of the fact that numbers in computers are almost always stored with finite precision and/or range: the expectations derived from the mathematical definition of Real numbers are not always fulfilled. See the box on page 18 for an in-depth explanation.

```
1 - 1e-20
## [1] 1
```

When comparing integer values these problems do not exist, as integer arithmetic is not affected by loss of precision in calculations restricted to integers. Because of the way integers are stored in the memory of computers, within the representable range, they are stored exactly. One can think of computer integers as a subset of whole numbers restricted to a certain range of values.

```
1L + 3L

## [1] 4

1L * 3L

## [1] 3

1L %/% 3L

## [1] 0

1L %% 3L

## [1] 1

1L / 3L

## [1] 0.3333333
```

The last statement in the example immediately above, using the "usual" division operator yields a floating-point double result, while the integer division operator %/% yields an integer result, and %% returns the remainder from the integer division. If as a result of an operation the result falls outside the range of representable values, the returned value is NA.

```
1000000L * 1000000L * 1000000L: NAS produced by integer overflow ## [1] NA
```

Both doubles and integers are considered numeric. In most situations, conversion is automatic and we do not need to worry about the differences between these two types of numeric values. The next chunk shows returned values that are either TRUE or FALSE. These are logical values that will be discussed in the next section.

```
is.numeric(1L)
## [1] TRUE

is.integer(1L)
## [1] TRUE

is.double(1L)
## [1] FALSE

is.double(1L / 3L)
## [1] TRUE

is.numeric(1L / 3L)
## [1] TRUE
```

Study the variations of the previous example shown below, and explain why the two statements return different values. Hint: 1 is a double constant. You can use is.integer() and is.double() in your explorations.

```
1 * 1000000L * 1000000L
1000000L * 1000000L * 1
```

Both when displaying numbers or as part of computations, we may want to decrease the number of significant digits or the number of digits after the decimal marker. Be aware that in the examples below, even if printing is being done by default, these functions return numeric values that are different from their input and can be stored and used in computations. Function round() is used to round numbers to a certain number of decimal places after or before the decimal marker, while signif() rounds to the requested number of significant digits.

```
round(0.0124567, digits = 3)
## [1] 0.012
signif(0.0124567, digits = 3)
## [1] 0.0125
```

```
round(1789.1234, digits = 3)
## [1] 1789.123

signif(1789.1234, digits = 3)
## [1] 1790

round(1789.1234, digits = -1)
## [1] 1790

a <- 0.12345
b <- round(a, digits = 2)
a == b
## [1] FALSE

a - b
## [1] 0.00345

b
## [1] 0.12</pre>
```

Being digits, the second parameter of these functions, the argument can also be passed by position. However, code is usually easier to understand for humans when parameter names are made explicit.

```
round(0.0124567, digits = 3)
## [1] 0.012
round(0.0124567, 3)
## [1] 0.012
```

Functions trunc() and ceiling() return the non-fractional part of a numeric value as a new numeric value. They differ in how they handle negative values, and neither of them rounds the returned value to the nearest whole number.

- What does value truncation mean? Function trunc() truncates a numeric value, but it does not return an integer.
- Explore how trunc() and ceiling() differ. Test them both with positive and negative values.
- **Advanced** Use function abs() and operators + and to reproduce the output of trunc() and ceiling() for the different inputs.
- Can trunc() and ceiling() be considered type conversion functions in R?

1.4 Logical values and Boolean algebra

What in Mathematics are usually called Boolean values, are called logical values in R. They can have only two values TRUE and FALSE, in addition to NA (not available). They are vectors as all other atomic types in R (by *atomic* we mean that each value is not composed of "parts"). There are also logical operators that allow Boolean algebra. In the chunk below we operate on logical vectors of length one.

```
a <- TRUE
b <- FALSE
mode(a)
## [1] "logical"

a
## [1] TRUE
!a # negation
## [1] FALSE

a && b # logical AND
## [1] FALSE

a || b # logical OR
## [1] TRUE

xor(a, b) # exclusive OR
## [1] TRUE</pre>
```

As with arithmetic operators, vectorization is available with *some* logical operators. The availability of two kinds of logical operators is one of the most troublesome aspects of the R language for beginners. Pairs of "equivalent" logical operators behave differently, use similar syntax and use similar symbols! The vectorized operators have single-character names & and ||, while the non-vectorized ones have double-character names && and ||. There is only one version of the negation operator! that is vectorized. In some, but not all cases, a warning will indicate that there is a possible problem.

```
a <- c(TRUE,FALSE)
b <- c(TRUE,TRUE)
a
## [1] TRUE FALSE

b
## [1] TRUE TRUE

a & b # vectorized AND
## [1] TRUE FALSE

a | b # vectorized OR
## [1] TRUE TRUE

a && b # not vectorized</pre>
```

```
## Warning in a && b: 'length(x) = 2 > 1' in coercion to 'logical(1)'
## Warning in a && b: 'length(x) = 2 > 1' in coercion to 'logical(1)'
## [1] TRUE
a || b # not vectorized
## Warning in a || b: 'length(x) = 2 > 1' in coercion to 'logical(1)'
## [1] TRUE
```

Functions any() and all() take zero or more logical vectors as their arguments, and return a single logical value "summarizing" the logical values in the vectors. Function all() returns TRUE only if all values in the vectors passed as arguments are TRUE, and any() returns TRUE unless all values in the vectors are FALSE.

```
any(a)
## [1] TRUE

all(a)
## [1] FALSE

any(a & b)
## [1] TRUE

all(a & b)
## [1] FALSE
```

Another important thing to know about logical operators is that they "short-cut" evaluation. If the result is known from the first part of the statement, the rest of the statement is not evaluated. Try to understand what happens when you enter the following commands. Short-cut evaluation is useful, as the first condition can be used as a guard protecting a later condition from being evaluated when it would trigger an error.

When using the vectorized operators on vectors of length greater than one, 'short-cut' evaluation still applies for the result obtained at each index position.

```
a & b & NA
## [1] NA FALSE

a & b & C(NA, NA)
## [1] NA FALSE

a | b | C(NA, NA)
## [1] TRUE TRUE
```

Based on the description of "recycling" presented on page 8 for numeric operators, explore how "recycling" works with vectorized logical operators. Create logical vectors of different lengths (including length one) and *play* by writing several code statements with operations on them. To get you started, one example is given below. Execute this example, and then create and run your own, making sure that you understand why the values returned are what they are. Sometimes, you will need to devise several examples or test cases to tease out of R an understanding of how a certain feature of the language works, so do not give up early, and make use of your imagination!

```
x <- c(TRUE, FALSE, TRUE, NA)
x & FALSE
x | c(TRUE, FALSE)</pre>
```

1.5 Comparison operators and operations

Comparison operators return vectors of logical values as results.

```
1.2 > 1.0
## [1] TRUE

1.2 >= 1.0
## [1] TRUE

1.2 == 1.0 # be aware that here we use two = symbols
## [1] FALSE

1.2 != 1.0
## [1] TRUE

1.2 <= 1.0
## [1] FALSE

1.2 < 1.0
## [1] FALSE

1.4 < 1.0
## [1] TRUE
```

These operators can be used on vectors of any length, returning as a result a logical vector as long as the longest operand. In other words, they behave in the same way as the arithmetic operators described on page 8: their arguments are recycled when needed. Hint: if you do not know what to expect as a value for the vector returned by 1:10, execute the statement print(a) after the first code statement below, or, alternatively, 1:10 without saving the result to a variable.

```
a <- 1:10
a > 5

## [1] FALSE FALSE FALSE FALSE FALSE TRUE TRUE TRUE TRUE TRUE

a < 5

## [1] TRUE TRUE TRUE TRUE FALSE FALSE FALSE FALSE FALSE FALSE FALSE

a == 5

## [1] FALSE FALSE FALSE FALSE TRUE FALSE FALSE FALSE FALSE FALSE

all(a > 5)

## [1] TRUE

b <- a > 5

b

## [1] TRUE

any(b)

## [1] TRUE

all(b)

## [1] FALSE
```

Precedence rules also apply to comparison operators and they can be overridden by means of parentheses.

```
a > 2 + 3
## [1] FALSE FALSE FALSE FALSE TRUE TRUE TRUE TRUE TRUE

(a > 2) + 3
## [1] 3 3 4 4 4 4 4 4 4 4 4
```

Use the statement below as a starting point in exploring how precedence works when logical and arithmetic operators are part of the same statement. *Play* with the example by adding parentheses at different positions and based on the returned values, work out the default order of operator precedence used for the evaluation of the example given below.

```
a <- 1:10
a > 3 | a + 2 < 3
```

Again, be aware of "short-cut evaluation". If the result does not depend on the missing value, then the result, TRUE or FALSE is returned. If the presence of the NA makes the end result unknown, then NA is returned.

```
c \leftarrow c(a, NA)
c > 5
## [1] FALSE FALSE FALSE FALSE TRUE TRUE TRUE TRUE TRUE
                                                                     NA
all(c > 5)
## [1] FALSE
any(c > 5)
## [1] TRUE
all(c < 20)
## [1] NA
any(c > 20)
## [1] NA
is.na(a)
## [1] FALSE FALSE FALSE FALSE FALSE FALSE FALSE FALSE FALSE
## [1] FALSE FALSE FALSE FALSE FALSE FALSE FALSE FALSE FALSE TRUE
any(is.na(c))
## [1] TRUE
all(is.na(c))
## [1] FALSE
```

The behavior of many of base-R's functions when NAS are present in their input arguments can be modified. TRUE passed as an argument to parameter na.rm, results in NA values being *removed* from the input **before** the function is applied.

```
all(c < 20)
## [1] NA

any(c > 20)
## [1] NA

all(c < 20, na.rm=TRUE)
## [1] TRUE

any(c > 20, na.rm=TRUE)
## [1] FALSE
```

Here I give some examples for which the finite resolution of computer machine floats, as compared to Real numbers as defined in mathematics, can cause serious problems. In R, numbers that are not integers are stored as *double-precision floats*. In addition to having limits to the largest and smallest numbers that can be represented, the precision of floats is limited by the number of significant digits that can be stored. Precision is usually described by "epsilon" (ϵ) , abbreviated *eps*, defined as the largest value of ϵ for which $1 + \epsilon = 1$. The finite resolution of floats can lead to unexpected results when testing for equality. In the second example below, the result of the subtraction is still exactly 1 due to insufficient resolution.

```
0 - 1e-20

## [1] -1e-20

1 - 1e-20

## [1] 1
```

The finiteness of floats also affects tests of equality, which is more likely to result in errors with important consequences.

```
1e20 == 1 + 1e20

## [1] TRUE

1 == 1 + 1e-20

## [1] TRUE

0 == 1e-20

## [1] FALSE
```

As R can run on different types of computer hardware, the actual machine limits for storing numbers in memory may vary depending on the type of processor and even compiler used to build the R program executable. However, it is possible to obtain these values at run time from the variable .Machine, which is part of the R language. Please see the help page for .Machine for a detailed and up-to-date description of the available constants.

```
.Machine$double.eps
## [1] 2.220446e-16

.Machine$double.neg.eps
## [1] 1.110223e-16

.Machine$double.max
## [1] 1024

.Machine$double.min
## [1] -1022
```

The last two values refer to the exponents of 10, rather than the maximum and minimum size of numbers that can be handled as objects of class double. Values outside these limits are stored as —Inf or Inf and enter arithmetic as infinite values according the mathematical rules.

```
le1026
## [1] Inf

le-1026
## [1] 0

Inf + 1
## [1] Inf

-Inf + 1
## [1] -Inf
```

As integer values are stored in machine memory without loss of precision, epsilon is not defined for integer values.

```
.Machine$integer.max
## [1] 2147483647
2147483699L
## [1] 2147483699
```

In those statements in the chunk below where at least one operand is double the integer operands are *promoted* to double before computation. A similar promotion does not take place when operations are among integer values, resulting in *overflow*, meaning numbers that are too big to be represented as integer values.

```
2147483600L + 99L

## warning in 2147483600L + 99L: NAs produced by integer overflow

## [1] NA

2147483600L + 99

## [1] 2147483699

2147483600L * 2147483600L

## warning in 2147483600L * 2147483600L: NAs produced by integer overflow

## [1] NA

2147483600L * 2147483600

## [1] 4.611686e+18
```

We see next that the exponentiation operator \land forces the promotion of its arguments to double, resulting in no overflow. In contrast, as seen above, the multiplication operator * operates on integers resulting in overflow.

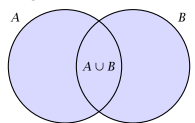
```
2147483600L * 2147483600L * 2147483600L: NAs produced by integer overflow ## [1] NA  
2147483600L^2L  
## [1] 4.611686e+18
```

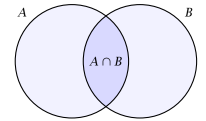
In many situations, when writing programs one should avoid testing for equality of floating point numbers ('floats'). Here we show how to gracefully handle rounding errors. As the example shows, rounding errors may accumulate, and in practice .Machine\$double.eps is not always a good value to safely use in tests for "zero," and a larger value may be needed. Whenever possible according to the logic of the calculations, it is best to test for inequalities, for example using $x \le 1.0$ instead of x = 1.0. If this is not possible, then the tests should be done replacing tests like x = 1.0 with abs (x - 1.0) < eps. Function abs () returns the absolute value, in simpler words, makes all values positive or zero, by changing the sign of negative values, or in mathematical notation |x| = |-x|.

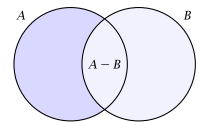
```
a == 0.0 # may not always work
## [1] FALSE FALSE FALSE FALSE FALSE FALSE FALSE FALSE
abs(a) < 1e-15 # is safer
## [1] FALSE FALSE FALSE FALSE FALSE FALSE FALSE FALSE
sin(pi) == 0.0 # angle in radians, not degrees!
## [1] FALSE
sin(2 * pi) == 0.0
## [1] FALSE
abs(sin(pi)) < 1e-15
## [1] TRUE
abs(sin(2 * pi)) < 1e-15
## [1] TRUE
sin(pi)
## [1] 1.224606e-16
sin(2 * pi)
## [1] -2.449213e-16
```

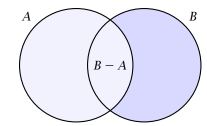
1.6 Sets and set operations

The R language supports set operations on vectors. They can be useful in many different contexts when manipulating and comparing vectors of values. Set algebra operations and their equivalents in mathematical notation and R functions are: union, \cup , union(); intersection, \cap , intersect(); difference (asymmetrical), -, setdiff(); equality test setequal(); membership, is element() and %in%. The first three functions return vector of the same mode as their inputs, and the last three a logical vector. The action of the first three operations is most easily illustrated with Venn diagrams.









In Bioinformatics it is usual, for example, to have character vectors of gene tags. We may have a vector for each of a set of different samples, and need to compare them. However, we start by using a more mundane example, everyday shopping, as illustration, followed by explanations.

```
fruits <- c("apple", "pear", "orange", "lemon", "tangerine")
bakery <- c("bread", "buns", "cake", "cookies")
dairy <- c("milk", "butter", "cheese")</pre>
shopping <- c("bread", "butter", "apple", "cheese", "orange")</pre>
intersect(fruits, shopping)
## [1] "apple" "orange"
intersect(bakery, shopping)
## [1] "bread"
intersect(dairy, shopping)
## [1] "butter" "cheese"
"lemon" %in% dairy
## [1] FALSE
"lemon" %in% fruits
## [1] TRUE
dairy %in% shopping
## [1] FALSE TRUE TRUE
setdiff(union(bakery, dairy), shopping)
## [1] "buns" "cake" "cookies" "milk"
```

For explanations we use abstract (symbolic) examples.

```
my.set <- c("a", "b", "c", "b")
```

To test if a given value belongs to a set, we use operator %in% or its function equivalent <code>is.element()</code>. In the algebra of sets notation, this is written $a \in A$, where A is a set and a a member. The second statement shows that the %in% operator is vectorized on its left-hand-side (lhs) operand, returning a logical vector.

```
is.element("a", my.set)
## [1] TRUE

"a" %in% my.set
## [1] TRUE

c("a", "a", "z") %in% my.set
## [1] TRUE TRUE FALSE
```

E Keep in mind that inclusion is an asymmetrical (not reflective) operation among sets. The rhs argument is interpreted as a set, while the lhs argument is interpreted as a vector of values to test for inclusion. In other words, any duplicate member in the lhs will be retained while the rhs is interpreted as a set of unique values. The returned logical vector has the same length as the lhs.

```
my.set %in% "a"
## [1] TRUE FALSE FALSE
```

The negation of inclusion is $a \notin A$, and coded in R by applying the negation operator! to the result of the test done with %in% or function is.element().

```
!is.element("a", my.set)
## [1] FALSE
!"a" %in% my.set
## [1] FALSE
!c("a", "a", "z") %in% my.set
## [1] FALSE FALSE TRUE
```

Although inclusion is a set operation, it is also very useful for the simplification of if()...else statements by replacing multiple tests for alternative constant values of the same mode chained by multiple | operators. A useful property of %in% and is.element() is that they never return NA.

Use operator %in% to write more concisely the following comparisons. Hint: see section 1.4 on page 14 for the difference between | and || operators.

```
x <- c("a", "a", "z")
x == "a" | x == "b" | x == "c" | x == "d"
```

Convert the logical vectors of length 3 into a vector of length one. Hint: see help for functions all() and any().

With unique() we convert a vector of possibly repeated values into a set of unique values. In the algebra of sets, a certain object belongs or not to a set. Consequently, in a set, multiple copies of the same object or value are meaningless.

```
unique(my.set)
## [1] "a" "b" "c"

c("a", "a", "z") %in% unique(my.set)
## [1] TRUE TRUE FALSE
```

In the notation used in algebra of sets, the set union operator is \cup while the intersection operator is \cap . If we have sets A and B, their union is given by $A \cup B$ —in the next three examples, c("a", "a", "z") is a constant, while my.set is a variable.

```
union(c("a", "a", "z"), my.set)
## [1] "a" "z" "b" "c"
```

If we have sets *A* and *B*, their intersection is given by $A \cap B$.

```
intersect(c("a", "a", "z"), my.set)
## [1] "a"
```

What do you expect to be the difference between the values returned by the three statements in the code chunk below? Before running them, write down your expectations about the value each one will return. Only then run the code. Independently of whether your predictions were correct or not, write down an explanation of what each statement's operation is.

```
union(c("a", "a", "z"), my.set)
c(c("a", "a", "z"), my.set)
c("a", "a", "z", my.set)
```

In the algebra of sets notation $A \subseteq B$, where A and B are sets, indicates that A is a subset or equal to B. For a true subset, the notation is $A \subset B$. The operators with the reverse direction are \supseteq and \supset . Implement these four operations in four R statements, and test them on sets (represented by R vectors) with different "overlap" among set members.

All set algebra examples above use character vectors and character constants. This is just the most frequent use case. Sets operations are valid on vectors of any atomic class, including integer, and computed values can be part of statements. In the second and third statements in the next chunk, we need to use additional parentheses to alter the default order of precedence between arithmetic and set operators.

```
9L %in% 2L:4L

## [1] FALSE

9L %in% ((2L:4L) * (2L:4L))

## [1] TRUE

c(1L, 16L) %in% ((2L:4L) * (2L:4L))

## [1] FALSE TRUE
```

Empty sets are an important component of the algebra of sets, in R they are represented as vectors of zero length. Vectors and lists of zero length, which the R language fully supports, can be used to "encode" emptiness also in other contexts. These vectors do belong to a class such as numeric or character and must be compatible with other operands in an expression. By default, constructors for vectors, construct empty vectors.

```
length(integer())
## [1] 0

1L %in% integer()
## [1] FALSE

setdiff(1L:4L, union(1L:4L, integer()))
## integer(0)
```

Character values 25

Although set operators are defined for numeric vectors, rounding errors in 'floats' can result in unexpected results (see section 1.5 on page 18). The next two examples do, however, return the correct answers.

```
9 %in% (2:4)^2
## [1] TRUE

c(1, 5) %in% (1:10)^2
## [1] TRUE FALSE
```

1.7 Character values

Character variables can be used to store any character. Character constants are written by enclosing characters in quotes. There are three types of quotes in the ASCII character set, double quotes ", single quotes ', and back ticks ". The first two types of quotes can be used as delimiters of character constants.

In many computer languages, vectors of characters are distinct from vectors of character strings. In these languages, character vectors store at each index position a single character, while vectors of character strings store at each index position strings of characters of various lengths, such as words or sentences. If you are familiar with C or C++, you need to keep in mind that C's char and R's character are not equivalent and that in R, character vectors are vectors of character strings. In contrast to these other languages, in R there is no predefined class for vectors of individual characters and character constants enclosed in double or single quotes are not different.

Concatenating character vectors of length one does not yield a longer character string, it yields instead a longer vector.

```
a <- 'A'
b <- "bcdefg"
c <- "123"
d <- c(a, b, c)
d
## [1] "A" "bcdefg" "123"
```

Having two different delimiters available makes it possible to choose the type of quotes used as delimiters so that other quotes can be included in a string.

```
a <- "He said 'hello' when he came in"
a
## [1] "He said 'hello' when he came in"
b <- 'He said "hello" when he came in'
b
## [1] "He said \"hello\" when he came in"</pre>
```

The outer quotes are not part of the string, they are "delimiters" used to mark the boundaries. As you can see when b is printed special characters can be represented using "escape sequences". There are several of them, and here we will show just four, new line (\n) and tab (\t), \" the escape code for a quotation mark within a string and \\ the escape code for a single backslash \. We also show here the different behavior of print() and cat(), with cat() *interpreting* the escape sequences and print() displaying them as entered.

```
c <- "abc\ndef\tx\"yz\"\\\tm"
print(c)
## [1] "abc\ndef\tx\"yz\"\\\tm"

cat(c)
## abc
## def x"yz"\ m</pre>
```

The *escape codes* work only in some contexts, as when using cat() to generate the output. For example, the new-line escape (\n) can be embedded in strings used for axis-label, title or label in a plot to split them over two or more lines.

1.8 The 'mode' and 'class' of objects

Variables have a *mode* that depends on what is stored in them. But different from other languages, assignment to a variable of a different mode is allowed and in most cases its mode changes together with its contents. However, there is a restriction that all elements in a vector, array or matrix, must be of the same mode. While this is not required for lists, which can be heterogenous. In practice this means that we can assign an object, such as a vector, with a different mode to a name already in use, but we cannot use indexing to assign an object of a different mode to individual members of a vector, matrix or array. Functions with names starting with is. are tests returning a logical value, TRUE, FALSE or NA. Function mode() returns the mode of an object, as a character string and typeof() returns R's internal type or storage mode.

```
my_var <- 1:5
mode(my_var) # no distinction of integer or double
## [1] "numeric"</pre>
```

```
typeof(my_var)
## [1] "integer"

is.numeric(my_var) # no distinction of integer or double
## [1] TRUE

is.double(my_var)
## [1] FALSE

is.integer(my_var)
## [1] TRUE

is.logical(my_var)
## [1] FALSE

is.character(my_var)
## [1] FALSE

my_var <- "abc"
mode(my_var)
## [1] "character"</pre>
```

While *mode* is a fundamental property, and limited to those modes defined as part of the R language, the concept of *class*, is different in that new classes can be defined in user code. In particular, different R objects of a given mode, such as numeric, can belong to different classes. The use of classes for dispatching functions is discussed in section ?? on page ??, in relation to object-oriented programming in R. Method class() is used to query the class of an object, and method inherits() is used to test if an object belongs to a specific class or not (including "parent" classes, to be later described).

```
class(my_var)
## [1] "character"
inherits(my_var, "character")
## [1] TRUE
inherits(my_var, "numeric")
## [1] FALSE
```

1.9 'Type' conversions

The least-intuitive type conversions are those related to logical values. All others are as one would expect. By convention, functions used to convert objects from one mode to a different one have names starting with as.¹.

```
as.character(1)
## [1] "1"
```

¹Except for some packages in the 'tidyverse' that use names starting with as_ instead of as...

```
as.numeric("1")
## [1] 1

as.logical("TRUE")
## [1] TRUE

as.logical("NA")
## [1] NA
```

Conversion takes place automatically in arithmetic and logical expressions.

```
TRUE + 10
## [1] 11

1 || 0
## [1] TRUE

FALSE | -2:2
## [1] TRUE TRUE FALSE TRUE TRUE
```

There is some flexibility in the conversion from character strings into numeric and logical values. Use the examples below plus your own variations to get an idea of what strings are acceptable and correctly converted and which are not. Do also pay attention at the conversion between numeric and logical values.

```
as.character(3.0e10)
as.numeric("5E+5")
as.numeric("A")
as.numeric(TRUE)
as.numeric(FALSE)
as.logical("T")
as.logical("t")
as.logical("true")
as.logical(100)
as.logical(0)
as.logical(-1)
```

- Compare the values returned by trunc() and as.integer() when applied to a floating point number, such as 12.34. Check for the equality of values, and for the *class* of the returned objects.
- Using conversions, the difference between the length of a character vector and the number of characters composing each member "string" within a vector is obvious.

```
f <- c("1", "2", "3")
length(f)
## [1] 3

g <- "123"
length(g)
## [1] 1

as.numeric(f)
## [1] 1 2 3

as.numeric(g)
## [1] 123</pre>
```

Other functions relevant to the "conversion" of numbers and other values are format(), and sprintf(). These two functions return character strings, instead of numeric or other values, and are useful for printing output. One could think of these functions as advanced conversion functions returning formatted, and possibly combined and annotated, character strings. However, they are usually not considered normal conversion functions, as they are very rarely used in a way that preserves the original precision of the input values. We show here the use of format() and sprintf() with numeric values, but they can also be used with values of other modes.

When using format(), the format used to display numbers is set by passing arguments to several different parameters. As print() calls format() to make numbers *pretty* it accepts the same options.

```
x = c(123.4567890, 1.0)
format(x) # using defaults
## [1] "123.4568" " 1.0000"

format(x[1]) # using defaults
## [1] "123.4568"

format(x[2]) # using defaults
## [1] "1"

format(x, digits = 3, nsmall = 1)
## [1] "123.5" " 1.0"

format(x[1], digits = 3, nsmall = 1)
## [1] "123.5"

format(x[2], digits = 3, nsmall = 1)
## [1] "1.0"

format(x, digits = 3, scientific = TRUE)
## [1] "1.23e+02" "1.00e+00"
```

Function sprintf() is similar to C's function of the same name. The user interface is rather unusual, but very powerful, once one learns the syntax. All the formatting is specified using a character string as template. In this template, placeholders for data and the formatting instructions are embedded using special codes.

These codes start with a percent character. We show in the example below the use of some of these: f is used for numeric values to be formatted according to a "fixed point," while g is used when we set the number of significant digits and e for exponential or *scientific* notation.

```
x = c(123.4567890, 1.0)
sprintf("The numbers are: %4.2f and %.0f", x[1], x[2])
## [1] "The numbers are: 123.46 and 1"

sprintf("The numbers are: %.4g and %.2g", x[1], x[2])
## [1] "The numbers are: 123.5 and 1"

sprintf("The numbers are: %4.2e and %.0e", x[1], x[2])
## [1] "The numbers are: 1.23e+02 and 1e+00"
```

In the template "The numbers are: %4.2f and %.0f", there are two placeholders for numeric values, %4.2f and %.0f, so in addition to the template, we pass two values extracted from the first two positions of vector x. These could have been two different vectors of length one, or even numeric constants. The template itself does not need to be a character constant as in these examples, as a variable can be also passed as argument.

- Function format() may be easier to use, in some cases, but sprintf() is more flexible and powerful. Those with experience in the use of the C language will already know about sprintf() and its use of templates for formatting output. Even if you are familiar with C, look up the help pages for both functions, and practice, by trying to create the same formatted output by means of the two functions. Do also play with these functions with other types of data like integer and character.
- We have above described NA as a single value ignoring modes, but in reality NA s come in various flavors. NA_real_, NA_character_, etc. and NA defaults to an NA of class logical. NA is normally converted on the fly to other modes when needed, so in general NA is all we need to use.

```
a <- c(1, NA)
is.numeric(a[2])
## [1] TRUE

is.numeric(NA)
## [1] FALSE

b <- c("abc", NA)
is.character(b[2])
## [1] TRUE

is.character(NA)
## [1] FALSE

class(NA)
## [1] "logical"

class(NA_character_)
## [1] "character"</pre>
```

```
c <- NA
c(c, 2:3)
## [1] NA 2 3</pre>
```

However, even the statement below works transparently.

```
a[3] <- b[2]
```

1.10 Vector manipulation

If you have read earlier sections of this chapter, you already know how to create a vector. If not, see pages 6–9 before continuing.

In this section we are going to see how to extract or retrieve, replace, and move elements such as a_2 from a vector $a_{1=1...n}$. Elements are extracted using an index enclosed in single square brackets. The index indicates the position in the vector, starting from one, following the usual mathematical tradition. What in maths notation would be a_i , in R is represented as a[i] and the whole vector, by excluding the brackets and indexing vector, as a.

We extract the first 10 elements of the vector letters.

```
a[2]
## [1] "b"
```

Four constant vectors are available in R: letters, LETTERS, month.name and month.abb, of which we used letters in the example above. These vectors are always for English, irrespective of the locale.

```
month.name[6]
## [1] "June"
```

⚠ In R, indexes always start from one, while in some other programming languages such as C and C++, indexes start from zero. It is important to be aware of this difference, as many computation algorithms are valid only under a given indexing convention.

It is possible to extract a subset of the elements of a vector in a single operation, using a vector of indexes. The positions of the extracted elements in the result ("returned value") are determined by the ordering of the members of the vector of indexes—easier to demonstrate than to explain.

```
a[c(3, 2)]
## [1] "c" "b"

a[10:1]
## [1] "j" "i" "h" "g" "f" "e" "d" "c" "b" "a"
```

The length of the indexing vector is not restricted by the length of the indexed vector. However, only numerical indexes that match positions present in the indexed vector can extract values. Those values in the indexing vector pointing to positions that are not present in the indexed vector, result in NAS. This is easier to learn by *playing* with R, than from explanations. Play with R, using the following examples as a starting point.

```
length(a)
a[c(3, 3, 3, 3)]
a[c(10:1, 1:10)]
a[c(1, 11)]
a[11]
```

Have you tried some of your own examples? If not yet, do *play* with additional variations of your own before continuing.

Negative indexes have a special meaning; they indicate the positions at which values should be excluded. Be aware that it is *illegal* to mix positive and negative values in the same indexing operation.

```
a[-2]
## [1] "a" "c" "d" "e" "f" "g" "h" "i" "j"

a[-c(3,2)]
## [1] "a" "d" "e" "f" "g" "h" "i" "j"

a[-3:-2]
## [1] "a" "d" "e" "f" "g" "h" "i" "j"
```

Results from indexing with special values and zero may be surprising. Try to build a rule from the examples below, a rule that will help you remember what to expect next time you are confronted with similar statements using "subscripts" which are special values instead of integers larger or equal to one—this is likely to happen sooner or later as these special values can be returned by different R expressions depending on the value of operands or function arguments, some of them described earlier in this chapter.

```
a[]
a[0]
a[numeric(0)]
a[NA]
a[c(1, NA)]
a[NULL]
a[c(1, NULL)]
```

Another way of indexing, which is very handy, but not available in most other programming languages, is indexing with a vector of logical values. The logical vector used for indexing is usually of the same length as the vector from which elements are going to be selected. However, this is not a requirement, because if the logical vector of indexes is shorter than the indexed vector, it is "recycled" as discussed above in relation to other operators.

```
a[TRUE]
## [1] "a" "b" "c" "d" "e" "f" "g" "h" "i" "j"

a[FALSE]
## character(0)

a[c(TRUE, FALSE)]
## [1] "a" "c" "e" "g" "i"

a[c(FALSE, TRUE)]
## [1] "b" "d" "f" "h" "j"

a > "c"
## [1] FALSE FALSE FALSE TRUE TRUE TRUE TRUE TRUE TRUE TRUE

a[a > "c"]
## [1] "d" "e" "f" "g" "h" "i" "j"
```

Indexing with logical vectors is very frequently used in R because comparison operators are vectorized. Comparison operators, when applied to a vector, return a logical vector, a vector that can be used to extract the elements for which the result of the comparison test was TRUE.

The examples in this text box demonstrate additional uses of logical vectors: 1) the logical vector returned by a vectorized comparison can be stored in a variable, and the variable used as a "selector" for extracting a subset of values from the same vector, or from a different vector.

```
a <- letters[1:10]
b <- 1:10
selector <- a > "c"
selector
a[selector]
b[selector]
```

Numerical indexes can be obtained from a logical vector by means of function which().

```
indexes <- which(a > "c")
indexes
a[indexes]
b[indexes]
```

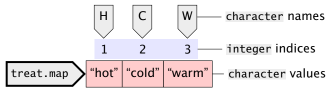
Make sure to understand the examples above. These constructs are very widely used in R because they allow for concise code that is easy to understand once you are familiar with the indexing rules. However, if you do not command these rules, many of these terse statements will be unintelligible to you.

In all earlier examples we have used integer valued indices for extraction of elements. In the vectors used as examples above the elements were anonymous or nameless. In R the elements can be assigned names, and these names used in place of numeric indices to extract the named elements. There is one situation where this is very useful: the mapping of values between two representations.

Let's assume we have a long vector encoding treatments using single letter codes and we want to replace these codes with clearer names.

```
treat <- c("H", "C", "H", "W", "C", "H", "H", "W", "W")
```

We can create a named vector to *map* the single letter codes into some other codes, in this case full words that are easier to understand.



As treat.map is a named vector, we can use the element names as indices for element extraction.

```
treat.map["H"]
## H
## "hot"
```

The indexing vector can be of a different length than the indexed vector, and that the returned value is a new vector of the same length as the indexing vector.

```
treat.new <- treat.map[treat]
treat.new
## H C H W C H H W W
## "hot" "cold" "hot" "hot" "warm" "warm"</pre>
```

where treat.new is a named vector, from which we will frequently want to remove the names.

```
treat.new <- unname(treat.new)
treat.new
## [1] "hot" "cold" "hot" "warm" "cold" "hot" "hot" "warm"
```

It is more common to use named members with lists than with vectors, but in R, in both cases it is possible to use both numeric positional indices and names.

Indexing can be used on either side of an assignment expression. In the chunk below, we use the extraction operator on the left-hand side of the assignments to replace values only at selected positions in the vector. This may look rather esoteric at first sight, but it is just a simple extension of the logic of indexing described above. It works, because the low precedence of the <- operator results in both the left-hand side and the right-hand side being fully evaluated before the assignment takes place. To make the changes to the vectors easier to follow, we use identical vectors with different names for each of these examples.

```
a < -1:10
  [1] 1 2 3 4 5 6 7 8 9 10
a[1] < -99
##
   [1] 99 2 3 4 5 6 7 8 9 10
b <- 1:10
b[c(2,4)] <- -99 # recycling
   [1] 1 -99 3 -99
##
                       5
                                7
                                   8
                                       9 10
                            6
c <- 1:10
c[c(2,4)] \leftarrow c(-99, 99)
  [1] 1 -99 3 99 5
                          6
                                       9 10
d < -1:10
d[TRUE] <- 1 # recycling</pre>
  [1] 1 1 1 1 1 1 1 1 1 1
e <- 1:10
e <- 1 # no recycling
## [1] 1
```

We can also use subscripting on both sides of the assignment operator, for example, to swap two elements.

```
a <- letters[1:10]
a[1:2] <- a[2:1]
a
## [1] "b" "a" "c" "d" "e" "f" "g" "h" "i" "j"
```

Do play with subscripts to your heart's content, really grasping how they work and how they can be used, will be very useful in anything you do in the future with R. Even the contrived example below follows the same simple rules, just study it bit by bit. Hint: the second statement in the chunk below, modifies a, so, when studying variations of this example you will need to recreate a by executing the first statement, each time you run a variation of the second statement.

```
a <- letters[1:10]
a[5:1] <- a[c(TRUE, FALSE)]
a</pre>
```

In R, indexing with positional indexes can be done with integer or numeric values. Numeric values can be floats, but for indexing, only integer values are meaningful. Consequently, double values are converted into integer values when used as indexes. The conversion is done invisibly, but it does slow down computations slightly. When working on big data sets, explicitly using integer values can improve performance.

```
b <- LETTERS[1:10]
b
## [1] "A" "B" "C" "D" "E" "F" "G" "H" "I" "J"

b[1]
## [1] "A"

b[1.1]
## [1] "A"

b[1.9999] # surprise!!
## [1] "A"

b[2]
## [1] "B"</pre>
```

From this experiment, we can learn that if positive indexes are not whole numbers, they are truncated to the next smaller integer.

```
b <- LETTERS[1:10]
b

## [1] "A" "B" "C" "D" "E" "F" "G" "H" "I" "J"

b[-1]

## [1] "B" "C" "D" "E" "F" "G" "H" "I" "J"

b[-1.1]

## [1] "B" "C" "D" "E" "F" "G" "H" "I" "J"

b[-1.9999]

## [1] "B" "C" "D" "E" "F" "G" "H" "I" "J"
```

From this experiment, we can learn that if negative indexes are not whole numbers, they are truncated to the next larger (less negative) integer. In conclusion, double index values behave as if they where sanitized using function trunc().

This example also shows how one can tease out of R its rules through experimentation.

A frequent operation on vectors is sorting them into an increasing or decreasing order. The most direct approach is to use sort().

```
my.vector <- c(10, 4, 22, 1, 4)
sort(my.vector)
## [1] 1  4  4  10  22

sort(my.vector, decreasing = TRUE)
## [1] 22  10  4  4  1</pre>
```

An indirect way of sorting a vector, possibly based on a different vector, is to generate with order() a vector of numerical indexes that can be used to achieve the ordering.

```
order(my.vector)
## [1] 4 2 5 1 3

my.vector[order(my.vector)]
## [1] 1 4 4 10 22

another.vector <- c("ab", "aa", "c", "zy", "e")
another.vector[order(my.vector)]
## [1] "zy" "aa" "e" "ab" "c"</pre>
```

A problem linked to sorting that we may face is counting how many copies of each value are present in a vector. We need to use two functions <code>sort()</code> and <code>rle()</code>. The second of these functions computes <code>run length</code> as used in <code>run length</code> encoding for which <code>rle</code> is an abbreviation. A <code>run</code> is a series of consecutive identical values. As the objective is to count the number of copies of each value present, we need first to sort the vector.

```
my.letters <- letters[c(1,5,10,3,1,4,21,1,10)]
my.letters
## [1] "a" "e" "j" "c" "a" "d" "u" "a" "j"

sort(my.letters)
## [1] "a" "a" "a" "c" "d" "e" "j" "j" "u"

rle(sort(my.letters))
## Run Length Encoding
## lengths: int [1:6] 3 1 1 1 2 1
## values: chr [1:6] "a" "c" "d" "e" "j" "u"</pre>
```

The second and third statements are only to demonstrate the effect of each step. The last statement uses nested function calls to compute the number of copies of each value in the vector.

1.11 Matrices and multidimensional arrays

Vectors have a single dimension, and, as we saw above, we can query their length with method length(). Matrices have two dimensions, which can be queried with dim(), ncol() and nrow(). R arrays can have any number of dimensions, even a single dimension, which can be queried with method dim(). As expected is.vector(), is.matrix() and is.array() can be used to query the class. In mathematical notation a matrix is denoted as follows

$$A_{m \times n} = \begin{bmatrix} a_{11} & a_{12} & \cdots & a_{1j} & \cdots & a_{1n} \\ a_{21} & a_{22} & \cdots & a_{2j} & \cdots & a_{2n} \\ \vdots & \vdots & \ddots & \vdots & & \vdots \\ a_{i1} & a_{i2} & \cdots & a_{ij} & \cdots & a_{in} \\ \vdots & \vdots & & \vdots & \ddots & \vdots \\ a_{m1} & a_{m2} & \cdots & a_{mj} & \cdots & a_{mn} \end{bmatrix}$$

where A represents the whole matrix and the a_{ij} its elements, with i indexing rows and j indexing columns. The two dimensions of the matrix are given by m and n, for rows and columns.

In R we can create a matrix using the matrix() or as.matrix() constructors. The first argument of matrix() is a vector. In the same way as vectors, matrices are homogeneous, all elements are of the same type.

```
matrix(1:15, ncol = 3)
     [,1] [,2] [,3]
       1 2
## [1,]
             6
## [2,]
                12
## [3,]
       3 8 13
## [4,]
## [5,]
matrix(1:15, nrow = 3)
     [,1] [,2] [,3] [,4] [,5]
## [1,] 1 4 7 10 13
      2 5
                 8
                    11
                         14
## [3,] 3 6
```

When a matrix is printed in R the row and column indexes are indicated on the edges left and top margins, in the same way as they would be used to extract whole rows and columns.

When a vector is converted to a matrix, R's default is to allocate the values in the vector to the matrix starting from the leftmost column, and within the column, down from the top. Once the first column is filled, the process continues from the top of the next column, as can be seen above. This order can be changed as you will discover in the playground below.

Check in the help page for the matrix constructor how to use the byrow parameter to alter the default order in which the elements of the vector are allocated to columns and rows of the new matrix.

```
help(matrix)
```

While you are looking at the help page, also consider the default number of columns and rows.

```
matrix(1:15)
```

And to start getting a sense of how to interpret error and warning messages, run the code below and make sure you understand which problem is being reported. Before executing the statement, analyze it and predict what the returned value will be. Afterwards, compare your prediction, to the value actually returned.

```
matrix(1:15, ncol = 2)
```

Subscripting of matrices and arrays is consistent with that used for vectors; we only need to supply an indexing vector, or leave a blank space, for each dimension. A matrix has two dimensions, so to access any element or group of elements, we use two indices. The only complication is that there are two possible orders in which, in principle, indexes could be supplied. In R, indexes for matrices are written "row first." In simpler words, the first index value selects rows, and the second one, columns.

```
A <- matrix(1:20, ncol = 4)
Α
      [,1] [,2] [,3] [,4]
##
## [1,]
       1 6 11
## [2,]
       2 7 12 17
## [3,]
       3 8 13
                  18
## [4,] 4 9 14
                    19
       5 10 15
## [5,]
A[1, 1]
## [1] 1
```

Remind yourself of how indexing of vectors works in R (see section 1.10 on page 31). We will now apply the same rules in two dimensions. The first or leftmost indexing vector corresponds to rows and the second one to columns, so R uses a rows-first convention for indexing. Missing indexing vectors are interpreted as meaning *extract all rows* and *extract all columns*, respectively.

```
A[1, ]
## [1] 1 6 11 16

A[, 1]
## [1] 1 2 3 4 5

A[2:3, c(1,3)]
## [,1] [,2]
## [1,] 2 12
## [2,] 3 13

A[3, 4] <- 99
A
```

```
## [,1] [,2] [,3] [,4]
## [1,]
       1
2
               11
           6
                    16
## [2,]
            7
                12
                    17
        3
           8 13 99
## [3,]
       4
           9 14 19
## [4,]
## [5,]
       5 10 15
                   20
A[4:3, 2:1] \leftarrow A[3:4, 1:2]
      [,1] [,2] [,3] [,4]
## [1,]
       1 6 11 16
        2
            7
               12
                    17
## [2,]
## [3,] 9 4 13
                   99
## [4,]
               14
        8
            3
                    19
## [5,] 5 10 15
```

⚠ Matrices can be indexed as vectors, without triggering an error or warning.

```
A <- matrix(1:20, ncol = 4)
Α
##
      [,1] [,2] [,3] [,4]
## [1,] 1 6 11 16
## [2,]
       2
           7 12
                    17
       3
## [3,]
           8 13 18
           9 14
## [4,]
       4
                   19
               15
       5 10
                    20
## [5,]
dim(A)
## [1] 5 4
A[10]
## [1] 10
A[5, 2]
## [1] 10
```

The next code example demonstrates that indexing as a vector with a single index, always works column-wise even if matrix ${\bf B}$ was created by assigning vector elements by row.

```
B <- matrix(1:20, ncol = 4, byrow = TRUE)</pre>
В
##
       [,1] [,2] [,3] [,4]
                  3
        1
             2
## [1,]
                        4
## [2,]
          5
              6
                    7
                         8
## [3,]
          9
              10
                   11
                        12
## [4,]
         13
              14
                   15
                        16
             18
                  19
## [5,]
        17
                        20
dim(B)
## [1] 5 4
B[10]
## [1] 18
B[5, 2]
## [1] 18
```

In R, a matrix can have a single row, a single column, a single element or no elements. However, in all cases, a matrix will have a *dimensions* attribute of length two defined.

```
my.vector <- 1:6
dim(my.vector)
## NULL

one.col.matrix <- matrix(1:6, ncol = 1)
dim(one.col.matrix)
## [1] 6 1

two.col.matrix <- matrix(1:6, ncol = 2)
dim(two.col.matrix)
## [1] 3 2

one.elem.matrix <- matrix(1, ncol = 1)
dim(one.elem.matrix)
## [1] 1 1

no.elem.matrix <- matrix(numeric(), ncol = 0)
dim(no.elem.matrix)
## [1] 0 0</pre>
```

Arrays are similar to matrices, but can have more than two dimensions, which are specified with the dim argument to the array() constructor.

```
B \leftarrow array(1:27, dim = c(3, 3, 3))
## , , 1
##
     [,1] [,2] [,3]
## [1,] 1 4 7
## [2,] 2 5 8
## [3,]
       3 6
##
## , , 2
##
##
      [,1] [,2] [,3]
## [1,]
       10 13 16
       11
           14
               17
## [2,]
## [3,]
       12
           15
                18
## , , 3
##
##
     [,1] [,2] [,3]
## [1,] 19 22 25
            23
## [2,] 20
                 26
## [3,] 21 24
                 27
B[2, 2, 2]
## [1] 14
```

In the chunk above, the length of the supplied vector is the product of the dimensions, $27 = 3 \times 3 \times 3 = 3^3$. Arrays are printed in slices, with slices across

3rd and higher dimensions printed separately, with their corresponding indexes above each slice and the first two dimensions on the margins of the individual slices, similarly to how matrices are displayed.

How do you use indexes to extract the second element of the original vector, in each of the following matrices and arrays?

```
v <- 1:10
m2c <- matrix(v, ncol = 2)
m2cr <- matrix(v, ncol = 2, byrow = TRUE)
m2r <- matrix(v, nrow = 2)
m2rc <- matrix(v, nrow = 2, byrow = TRUE)

v <- 1:10
a2c <- array(v, dim = c(5, 2))
a2c <- array(v, dim = c(5, 2), dimnames = list(NULL, c("c1", "c2")))
a2r <- array(v, dim = c(2, 5))</pre>
```

Be aware that vectors and one-dimensional arrays are not the same thing, while two-dimensional arrays are matrices.

- 1. Use the different constructors and query methods to explore this, and its consequences.
- Convert a matrix into a vector using unlist() and as.vector() and compare the returned values.

Operators for matrices are available in R, as matrices are used in many statistical algorithms. We will not describe them all here, only t() and some specializations of arithmetic operators. Function t() transposes a matrix, by swapping columns and rows.

```
A \leftarrow matrix(1:20, ncol = 4)
Α
##
      [,1] [,2] [,3] [,4]
## [1,] 1 6 11 16
## [2,]
        2
            7 12 17
## [3,] 3 8 13 18
## [4,] 4 9 14 19
## [5,] 5 10 15
t(A)
##
      [,1] [,2] [,3] [,4] [,5]
       1 2 3 4
6 7 8 9
## [1,]
## [2,]
                         10
        11 12
## [3,]
                13
                     14
                         15
       16 17
                         20
## [4,]
               18
                    19
```

As with vectors, recycling applies to arithmetic operators when applied to matrices.

```
A + 2
## [,1] [,2] [,3] [,4]
```

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```
3 8 13
## [1,]
                      18
## [2,]
             9
                 14
                      19
         5
## [3,]
             10
                 15
                      20
## [4,]
         6
             11
                 16
                      21
## [5,]
         7
             12
                 17
                      22
A * 0:1
       [,1] [,2] [,3] [,4]
##
## [1,]
       0 6 0 16
## [2,]
             0
                 12
## [3,]
        0 8 0
                      18
## [4,]
         4
            0 14
                      0
## [5,]
         0
            10
                 0
                    20
A \times 1:0
##
      [,1] [,2] [,3] [,4]
## [1,]
         1
             0
                 11
## [2,]
         0
              7
                  0
                      17
## [3,]
         3
             0
                 13
                      0
## [4,]
         0
             9
                 0
                      19
## [5,]
         5
              0
                 15
                       0
```

In the examples above with the usual multiplication operator *, the operation described is not a matrix product, but instead, the products between individual elements of the matrix and vectors. Matrix multiplication is indicated by operator ***.

```
B <- matrix(1:16, ncol = 4)
B * B
##
       [,1] [,2] [,3] [,4]
## [1,]
        1 25 81 169
## [2,]
        4 36 100 196
## [3,]
        9 49 121
                     225
## [4,]
        16 64 144
                     256
B %*% B
##
       [,1] [,2] [,3] [,4]
## [1,] 90 202 314
                     426
## [2,] 100 228
                 356
                     484
## [3,]
       110 254
                 398
                      542
## [4,] 120 280
                440
```

Other operators and functions for matrix algebra like cross-product (crossprod()), extracting or replacing the diagonal (diag()) are available in base R. Packages, including 'matrixStats', provide additional functions and operators for matrices.

1.12 Factors

Factors are very important in R. In contrast to other statistical software in which the role of a variable is set when defining a model to be fitted or when setting up a test, in R, models are specified exactly in the same way for ANOVA and regression anal-

ysis, both as *linear models*. The type of model that is fitted is decided by whether the explanatory variable is a factor (giving ANOVA) or a numerical variable (giving regression). This makes a lot of sense, because in most cases, considering an explanatory variable as categorical or not, depends on the quantity stored and/or the design of the experiment or survey. In other words, being categorical is a property of the data. The order of the levels in an unordered **factor** does not affect simple calculations or the values plotted, but as we will see in chapters ?? and ??, it does affect how the output is printed, the order of the levels in the scales and keys of plots, and in some cases how contrasts are applied in significance tests.

In a factor, values indicate discrete unordered categories, most frequently the treatments in an experiment, or categories in a survey. They can be created either from numerical or character vectors. The different possible values are called *levels*. Factors created with factor() are always unordered or categorical. R also supports ordered factors, created with function ordered() with identical user interface. The distinction, however, only affects how they are interpreted in statistical tests as discussed in chapter ??.

When using factor() or ordered() we create a factor from a vector, but this vector can be created on-the-fly and anonymous as shown in this example. When the vector is numeric and no labels are supplied, level labels are character strings matching the numbers. The default ordering of the levels is alphanumerical.

```
factor(x = c(1, 2, 2, 1, 2, 1, 1))
## [1] 1 2 2 1 2 1 1
## Levels: 1 2

ordered(x = c(1, 2, 2, 1, 2, 1, 1))
## [1] 1 2 2 1 2 1 1
## Levels: 1 < 2

factor(x = c(1, 2, 2, 1, 2, 1, 1), ordered = TRUE)
## [1] 1 2 2 1 2 1 1
## Levels: 1 < 2</pre>
```

When the pattern of levels is regular, it is possible to use function g1(), *generate levels*, to construct a factor. Nowadays, it is usual to read data into R from files in which the treatment codes are already available as character strings or numeric values, however, when we need to create a factor within R, g1() can save some typing. In this case instead of passing a vector as argument, we pass a *recipe* to create it: n is the number of levels, and k the number of contiguous repeats (called "replicates" in R documentation) and length the length of the factor to be created.

```
gl(n = 2, k = 5, labels = c("A", "B"))
## [1] A A A A A B B B B B
## Levels: A B

gl(n = 2, k = 1, length = 10, labels = c("A", "B"))
## [1] A B A B A B A B A B
## Levels: A B
```

It is always preferable to use meaningful labels for levels, even if R does not

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require it. Here the vector is stored in a variable named my.vector. In a real data analysis situation in most cases the vector would have been read from a file on disk and would be longer.

```
my.vector <- c("treated", "treated", "control", "control", "control", "treated")
factor(my.vector)
## [1] treated treated control control treated
## Levels: control treated</pre>
```

The ordering of levels is established at the time a factor is created, and by default is alphabetical. This default ordering of levels is frequently not the one needed. We can pass an argument to parameter levels of function factor() to set a different ordering of the levels.

```
factor(x = my.vector, levels = c("treated", "control"))
## [1] treated treated control control treated
## Levels: treated control
```

The labels ("names") of the levels can be set when calling factor(). Two vectors are passed as arguments to parameters levels and labels with levels and matching labels in the same position. The argument passed to levels determines the order of the levels based on their old names or values, and the argument passed to labels gives new names to the levels.

```
factor(x = c(1, 1, 0, 0, 0, 1), levels = c(1, 0), labels = c("treated", "control"))
## [1] treated treated control control treated
## Levels: treated control
```

In the examples above we passed a numeric vector or a character vector as an argument for parameter x of function factor(). It is also possible to pass a factor as an argument to parameter x. This makes it possible to modify the ordering of levels or replace the labels in a factor.

Merging factor levels. We use factor() as shown below, setting the same label for the levels we want to merge.

We can use indexing on factors in the same way as with vectors. In the next example, we use a test returning a logical vector to extract all "controls." We use function <code>levels()</code> to look at the levels of the factors, as with vectors, <code>lengtgh()</code> to query the number of values stored.

```
## [1] treated treated control control control treated
## Levels: control treated
levels(my.factor)
## [1] "control" "treated"
length(my.factor)
## [1] 6
control.factor <- my.factor[my.factor == "control"]</pre>
control.factor
## [1] control control
## Levels: control treated
levels(control.factor) # same as in my.factor
## [1] "control" "treated"
length(control.factor) # shorter than my.factor
## [1] 3
control.factor <- factor(control.factor)</pre>
levels(control.factor) # the unused level was dropped
## [1] "control"
```

It can be seen above that subsetting does not drop unused factor levels, and that factor() can be used to explicitly drop the unused factor levels.

⚠ How to convert factors into numeric vectors is not obvious, even when the factor was created from a numeric vector.

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```
my.vector2 <- rep(3:5, 4)
my.vector2
## [1] 3 4 5 3 4 5 3 4 5 3 4 5

my.factor2 <- factor(my.vector2)
my.factor2
## [1] 3 4 5 3 4 5 3 4 5 3 4 5
## Levels: 3 4 5

as.numeric(my.factor2)
## [1] 1 2 3 1 2 3 1 2 3 1 2 3

as.numeric(as.character(my.factor2))
## [1] 3 4 5 3 4 5 3 4 5 3 4 5</pre>
```

Why is a double conversion needed? Internally, factor values are are stored as running integers starting from one, each distinct integer value corresponding to a level. These underlying integer values are returned by as.numeric() when applied to a factor instead of the level labels. The labels of the factor levels are always stored as character strings, even when these characters are digits. In contrast to as.numeric(), as.character() returns the character labels of the levels for each of the values stored in the factor. If these character strings represent numbers, they can be converted, in a second step, using as.numeric() into the original numeric values. Use of class and mode is described on section 1.8 on page 26, and str() on page 53.

```
class(my.factor2)
## [1] "factor"

mode(my.factor2)
## [1] "numeric"

str(my.factor2)
## Factor w/ 3 levels "3","4","5": 1 2 3 1 2 3 1 2 3 1 ...
```

- Create a factor with levels labeled with words. Create another factor with the levels labeled with the same words, but ordered differently. After this convert both factors to numeric vectors using as.numeric(). Explain why the two numeric vectors differ or not from each other.
- Safely reordering and renaming factor levels. The simplest approach is to use factor() and its levels parameter as shown above. In these more advanced examples we use levels() to retrieve the names of the levels from the factor itself to protect from possible bugs due to typing mistakes, or for changes in the naming conventions used.

Reverse previous order using rev().

```
my.factor2 <- factor(c("treated", "treated", "control", "control", "control", "treated"))
levels(my.factor2)
## [1] "control" "treated"

my.factor2 <- factor(my.factor2, levels = rev(levels(my.factor2)))
levels(my.factor2)
## [1] "treated" "control"</pre>
```

Sort in decreasing order, i.e., opposite to default.

Alter ordering using subscripting; especially useful with three or more levels.

```
my.factor2 <- factor(my.factor2, levels = levels(my.factor2)[c(2, 1)])
levels(my.factor2)
## [1] "control" "treated"</pre>
```

Reordering the levels of a factor based on summary quantities from data stored in a numeric vector is very useful, especially when plotting. Function reorder() can be used in this case. It defaults to using mean() for summaries, but other suitable summary functions, such as median() can be supplied in its place.

```
my.factor3 <- gl(2, 5, labels = c("A", "B"))
my.vector3 <- c(5.6, 7.3, 3.1, 8.7, 6.9, 2.4, 4.5, 2.1, 1.4, 2.0)
my.factor3
## [1] A A A A A B B B B B
## Levels: A B

my.factor3ord <- reorder(my.factor3, my.vector3)
levels(my.factor3ord)
## [1] "B" "A"

my.factor3rev <- reorder(my.factor3, -my.vector3) # a simple trick
levels(my.factor3rev)
## [1] "A" "B"</pre>
```

In the last statement, using the unary negation operator, which is vectorized, allows us to easily reverse the ordering of the levels, while still using the default function, mean(), to summarize the data.

Reordering factor values. It is possible to arrange the values stored in a factor either alphabetically according to the labels of the levels or according to the order of the levels. (The use of rep() is explained on page 8.)

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```
# gl() keeps order of levels
my.factor4 <- gl(4, 3, labels = c("A", "F", "B", "Z"))
my.factor4
as.integer(my.factor4)
# factor() orders levels alphabetically
my.factor5 <- factor(rep(c("A", "F", "B", "Z"), rep(3,4)))
my.factor5
as.integer(my.factor5)
levels(my.factor5)[as.integer(my.factor5)]</pre>
```

We see above that the integer values by which levels in a factor are stored, are equivalent to indices or "subscripts" referencing the vector of labels. Function sort() operates on the values' underlying integers and sorts according to the order of the levels while order() operates on the values' labels and returns a vector of indices that arrange the values alphabetically.

```
sort(my.factor4)
my.factor4[order(my.factor4)]
my.factor4[order(as.integer(my.factor4))]
```

Run the examples in the chunk above and work out why the results differ.

1.13 Lists

Lists' main difference from vectors is, in R, that they can be heterogeneous. While the member elements of a vector must be *atomic* values, any R object can be a list member. In R, the members of a list can be considered as following a sequence, and accessible through numerical indexes, the same as members of vectors. Members of a list as well as members of a vector can be named, and retrieved (indexed) through their names. In practice, named lists are more frequently used than named vectors. Lists are created using function list() similarly as c() is used for vectors. Members of a list can be objects differing both in their class and in their size. Lists can be nested.

In R lists can have as members not only objects storing data on observations and categories, but also function definitions, model formulas, unevaluated expressions, matrices, arrays, and objects of user defined classes.

Differently to languages like C, C++ and Pascal, R does not implement memory pointers or linked lists using pointers.

Our first list has as its members three different vectors, each one belonging to a different class: numeric, character and logical. The three vectors also differ in their length: 6, 1, and 2, respectively.

```
a.list \leftarrow list(x = 1:3, y = "ab", z = c(TRUE, FALSE))
```

```
str(a.list)
## List of 3
   $ x: int [1:3] 1 2 3
   $ y: chr "ab"
   $ z: logi [1:2] TRUE FALSE
names(a.list)
## [1] "x" "y" "z"
                                           character names
                       У
                                Z
                                           integer indices
                       2
                                 3
                           TRUE, FALSE
    a.list
              1, 2, 3
                      "ab"
                                           heterogeneous values
                       "ab"
                               TRUE
    a.list
                2
                              FALSE
                3
```

With lists it is best to use informative names for indexing, as their members are heterogenous usually containing loosely related/connected data. Names make code easier to understand and mistakes more visible. Using names also makes code more robust to future changes in the position of fields in lists created upstream. Below, we use both positional indices and names to highlight the similarities between lists and vectors.

Lists can behave as vectors with heterogeneous elements as members, as we will describe next. Lists can be nested, so tree-like structures are also possible (see section 1.13.2 on page 53).

1.13.1 Member extraction, deletion and insertion

In section 1.10 on page 31 we saw that the extraction operator [] applied to a vector, returns a vector, longer or shorter, possibly of length one, or even length zero. Similarly, applying operator [] to a list returns a list, possibly of different length: a.list["x"] or a.list[1] return a list containing only one member, the numeric vector stored at the first position of a.list. In the last statement above, a.list[c(1, 3)] returns a list of length two as expected.

```
a.list["x"]
## $x
## [1] 1 2 3

a.list$x
## [1] 1 2 3

a.list[1]
## $x
## [1] 1 2 3

a.list[c(1, 3)]
```

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```
## $x
## [1] 1 2 3
##
## $z
## [1] TRUE FALSE
```

As with vectors negative positional indices remove members instead of extracting them. See page 52 for a safer approach to deletion of list members.

```
a.list[-1]
## $y
## [1] "ab"
##
## $z
## [1] TRUE FALSE

a.list[c(-1, -3)]
## $y
## [1] "ab"
```

Using operator [[]] (double square brackets) for indexing a list extracts the element stored in the list, in its original mode. In the example below, a.list[["x"]] and a.list[["x"]] return a numeric vector. We might say that extraction operator [[]] reaches "deeper" into the list than operator [].

```
a.list$x
## [1] 1 2 3

a.list[["x"]]
## [1] 1 2 3

a.list[[1]]
## [1] 1 2 3
```

The operators can be used together as shown below, with a.list[[1]] extracting the vector from a.list and [3] extracting the member at position 3 of the vector.

```
a.list[[1]][3]
## [1] 3
```

Operators can be used on the lhs as well as on the rhs of an assignment, and lists can be empty, i.e, be of length zero. The example below makes use of this to build a list step by step.

```
b.list <- list()
b.list[["x"]] <- 1:3
b.list[["y"]] <- "ab"
b.list[["z"]] <- c(TRUE, FALSE)</pre>
```

Compare b.list to a.list, used for the examples above.

```
b.list[["y"]] <- b.list[["x"]]
```

Compare them again, and try to understand why b.list has changed as it did. Pay also attention to possible changes to the members' names.

Suprisingly, a.list[[c(1, 3)]] returns the value at position 3 in the first member of the list, an operation that normally would be written as a.list[[1]][3] using two extractions, one after another. In most cases statements like the one below will be entered by mistake rather than intentionally, but being valid in R, they will not trigger an error message.

```
a.list[[c(1, 3)]]
## [1] 3
```

Lists, as usually defined in languages like C, are based on pointers to memory locations, with pointers stored at each node. These pointers chain or link the different member nodes (this allows, for example, sorting of lists in place by modifying the pointers). In such implementations, indexing by position is not possible, or at least requires "walking" down the list, node by node. In R, list members can be accessed through positional indexes, similarly to vectors. Of course, insertions and deletions in the middle of a list, shift the position of members and change which member is pointed at by indexes for positions past the modified location. The names, in contrast remain valid.

```
list(a = 1, b = 2, c = 3)[-2]
## $a
## [1] 1
##
## $c
## [1] 3
```

Two frequent simple operations on lists are insertions and deletions. In R, function append() can be used both to append elements at the end of a list and insert elements into the head or any position in the middle of a list.

```
another.list <- append(a.list, list(yy = 1:10, zz = letters[5:1]), 2L)
another.list
## $x
## [1] 1 2 3
##
## $y
## [1] "ab"
##
## $yy
  [1] 1 2 3 4 5 6 7 8 9 10
##
## $zz
## [1] "e" "d" "c" "b" "a"
##
## $z
## [1] TRUE FALSE
```

To delete a member from a list we assign **NULL** to it.

```
a.list$y <- NULL
a.list
```

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```
## $x
## [1] 1 2 3
##
## $z
## [1] TRUE FALSE
```

To investigate the members contained in a list, function str() (*structure*), used above, is convenient, especially when lists have many members. The print() method for the structure formats lists more compactly than function print() applied directly to a list.

```
print(a.list)
## $x
## [1] 1 2 3
##
## $z
## [1] TRUE FALSE
```

1.13.2 Nested lists

Lists can be nested, i.e., lists of lists can be constructed to an arbitrary depth.

```
a.list <- list("a", "aa", 10)
b.list <- list("b", TRUE)
nested.list <- list(A = a.list, B = b.list)
str(nested.list)
## List of 2
## $ A:List of 3
## ..$ : chr "a"
## ..$ : chr "aa"
## ..$ : num 10
## $ B:List of 2
## ..$ : chr "b"
## ..$ : logi TRUE</pre>
```

A nested list can alternatively be constructed within a single statement in which several member lists are created. Here we combine the first three statements in the earlier chunk into a single one.

```
nested.list <- list(A = list("a", "aa", 10), B = list("b", TRUE))
str(nested.list)
## List of 2
## $ A:List of 3
## ..$ : chr "a"
## ..$ : chr "aa"
## ..$ : num 10
## $ B:List of 2
## ..$ : chr "b"
## ..$ : logi TRUE</pre>
```

A list can contain a combination of list and vector members.

```
nested.list <- list(A = list("a", "aa", 10),
B = list("b", TRUE),</pre>
```

```
C = c(1, 3, 9),
D = 4321)

str(nested.list)

## List of 4

## $ A:List of 3

## ..$ : chr "a"

## ..$ : chr "aa"

## ..$ : num 10

## $ B:List of 2

## ..$ : chr "b"

## ..$ : logi TRUE

## $ C: num [1:3] 1 3 9

## $ D: num 4321
```

The logic behind extraction of members of nested lists using indexing is the same as for simple lists, but applied recursively—e.g., nested.list[[2]] extracts the second member of the outermost list, which is another list. As, this is a list, its members can be extracted using again the extraction operator: nested.list[[2]][[1]]. It is important to remember that these concatenated extraction operations are written so that the leftmost operator is applied to the outermost list.

The example above uses the [[]] operator, but the left to right precedence also applies to concatenated calls to [].

What do you expect each of the statements below to return? *Before running the code*, predict what value and of which mode each statement will return. You may use implicit or explicit calls to print(), or calls to str() to visualize the structure of the different objects.

```
nested.list <- list(A = list("a", "aa", "aaa"), B = list("b", "bb"))
str(nested.list)
nested.list[2:1]
nested.list[1]
nested.list[[1]][2]
nested.list[[1]][[2]]
nested.list[2]
nested.list[2][[1]]</pre>
```

When dealing with deep lists, it is sometimes useful to limit the number of levels of nesting returned by str() by means of a numeric argument passed to parameter max.levels.

```
str(nested.list, max.level = 1)
## List of 4
## $ A:List of 3
## $ B:List of 2
## $ C: num [1:3] 1 3 9
## $ D: num 4321
```

Sometimes we need to flatten a list, or a nested structure of lists within lists. Function unlist() is what should be normally used in such cases.

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The list nested.list is a nested system of lists, but all the "terminal" members are character strings. In other words, terminal nodes are all of the same mode, allowing the list to be "flattened" into a character vector.

```
nested.list <- list(A = list("a", "aa", "aaa"), B = list("b", "bb"))</pre>
c.vec <- unlist(nested.list)</pre>
c.vec
##
    A1 A2 A3
                       в1
                            В2
    "a" "aa" "aaa" "b" "bb"
##
is.list(nested.list)
## [1] TRUE
is.list(c.vec)
## [1] FALSE
mode(nested.list)
## [1] "list"
mode(c.vec)
## [1] "character"
names(nested.list)
## [1] "A" "B"
names(c.vec)
## [1] "A1" "A2" "A3" "B1" "B2"
```

The returned value is a vector with named member elements. We use function str() to figure out how this vector relates to the original list. The names, always of mode character, are based on the names of list elements when available, while characters depicting positions as numbers are used for anonymous nodes. We can access the members of the vector either through numeric indexes or names.

```
str(c.vec)
## Named chr [1:5] "a" "aa" "aaa" "b" "bb"
## - attr(*, "names")= chr [1:5] "A1" "A2" "A3" "B1" ...

c.vec[2]
## A2
## "aa"

c.vec["A2"]
## A2
## "aa"
```

Function unlist() has two additional parameters, with default argument values, which we did not modify in the example above. These parameters are recursive and use.names, both of them expecting a logical value as an argument. Modify the statement c.vec <- unlist(c.list), by passing FALSE as an argument to these two parameters, in turn, and in each case, study the value returned and how it differs with respect to the one obtained above.

Function unname() can be used to remove names safely—i.e., without risk of altering the mode or class of the object.

```
unname(c.vec)
## [1] "a" "aa" "aaa" "b"
                                "bb"
unname(nested.list)
## [[1]]
## [[1]][[1]]
## [1] "a"
##
## [[1]][[2]]
## [1] "aa"
##
## [[1]][[3]]
## [1] "aaa"
##
##
## [[2]]
## [[2]][[1]]
## [1] "b"
##
## [[2]][[2]]
## [1] "bb"
```

1.14 Data frames

Data frames are a special type of list, in which each element is a vector or a factor of the same length (or rarely a matrix with the same number of rows as the enclosing data frame). They are created with function <code>data.frame()</code> with a syntax similar to that used for lists—in object-oriented programming we say that data frames are derived from class <code>list</code>. As the expectation is equal length, if vectors of different lengths are supplied as arguments, the shorter vector(s) is/are recycled, possibly several times, until the required full length is reached.

```
a.df <- data.frame(x = 1:6, y = "a", z = c(TRUE, FALSE))
a.df
## x y
## 1 1 a TRUE
## 2 2 a FALSE
## 3 3 a TRUE
## 4 4 a FALSE
## 5 5 a TRUE
## 6 6 a FALSE
str(a.df)
## 'data.frame': 6 obs. of 3 variables:
## $ x: int 1 2 3 4 5 6
## $ y: chr "a" "a" "a" "a" ...
## $ z: logi TRUE FALSE TRUE FALSE TRUE FALSE
class(a.df)
## [1] "data.frame"
mode(a.df)
```

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```
## [1] "list"
is.data.frame(a.df)
## [1] TRUE
is.list(a.df)
## [1] TRUE
```

Extraction of individual member variables or "columns" can be done like in a list with operator [[]].

```
a.df$x
## [1] 1 2 3 4 5 6

a.df[["x"]]
## [1] 1 2 3 4 5 6

a.df[[1]]
## [1] 1 2 3 4 5 6

class(a.df[["x"]])
## [1] "integer"
```

With function class() we can query the class of an R object (see section 1.8 on page 26). As we saw in the two previous chunks, list and data.frame objects belong to two different classes. However, their relationship is based on a hierarchy of classes. We say that class data.frame is derived from class list. Consequently, data frames inherit the methods and characteristics of lists, as long as they have not been hidden by new ones defined for data frames.

In the same way as with lists, we can add members to data frames.

```
a.df$x2 <- 6:1
a.df$x3 <- "b"
str(a.df)
## 'data.frame': 6 obs. of 5 variables:
## $ x : int 1 2 3 4 5 6
## $ y : chr "a" "a" "a" "...
## $ z : logi TRUE FALSE TRUE FALSE TRUE FALSE
## $ x2: int 6 5 4 3 2 1
## $ x3: chr "b" "b" "b" "b" ...</pre>
```

We have added two columns to the data frame, and in the case of column x3 recycling took place. This is where lists and data frames differ substantially in their behavior. In a data frame, although class and mode can be different for different variables (columns), they are required to be vectors or factors of the same length (or a matrix with the same number of rows). In the case of lists, there is no such requirement, and recycling never takes place when adding a node. Compare the values returned below for a.1s, to those in the example above for a.df.

```
a.ls <- list(x = 1:6, y = "a", z = c(TRUE, FALSE))
str(a.ls)
## List of 3
## $ x: int [1:6] 1 2 3 4 5 6
## $ y: chr "a"
## $ z: logi [1:2] TRUE FALSE</pre>
```

```
a.ls$x2 <- 6:1
a.ls$x3 <- "b"
str(a.ls)

## List of 5

## $ x : int [1:6] 1 2 3 4 5 6

## $ y : chr "a"

## $ z : logi [1:2] TRUE FALSE

## $ x2: int [1:6] 6 5 4 3 2 1

## $ x3: chr "b"
```

Usually data frames are created from lists or by passing individual vectors and factors to the constructors. It is also possible to construct data frames starting from matrices, other data frames and named vectors and combinations of them. In these cases additional nuances become important. We give only some examples here, as the details are well described in help(data.frame).

We use a named numeric vector, and a factor. The names are moved from the vector to the rows of the data frame! Consult help(data.frame) for an explanation.

```
my.vector \leftarrow c(one = 1, two = 2, three = 3, four = 4)
my.factor \leftarrow as.factor(c(1, 2, 3, 2))
df1 <- data.frame(my.factor, my.vector)</pre>
df1
##
        my.factor my.vector
         1 1
## one
                          2
## two
               2
                3
                          3
## three
## four
                2
df1$my.vector
## [1] 1 2 3 4
```

If we protect the vector with R's identity function I() the names are not removed from the vector as can be seen by extracting the column from the data frame.

```
df2 <- data.frame(my.factor, I(my.vector))</pre>
df2
##
       my.factor my.vector
## one
       1
                   1
## two
             2
                      2
                      3
## three
             3
## four
df2$my.vector
## one two three four
## 1 2 3 4
```

If we start with a matrix instead of a vector, the matrix is split into separate columns in the data frame. If the matrix has no column names, new ones are created.

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If we protect the matrix with function **I()**, it is not split, and the whole matrix becomes a column in the data frame.

```
df5 <- data.frame(my.factor, I(my.matrix))</pre>
df5
## my.factor my.matrix.1 my.matrix.2 my.matrix.3
## 1
       1 1 5
## 2
          2
                    2
                              6
                                       10
## 3
          3
                    3
                              7
                                       11
          2
                    4
                              8
## 4
                                       12
df5$my.matrix
##
    [,1] [,2] [,3]
## [1,]
      1 5 9
       2
## [2,]
            6
                10
       3
## [3,]
            7
                11
       4 8
## [4,]
```

If we start with a list, each member with a suitable number of elements, each member becomes a column in the data frame. In the case of a too short one, recycling is applied.

If we protect the list, then the list is added in whole, similarly as in a tibble (see chapter ?? starting on page ?? for details about the 'tidiyverse').

```
df7<- data.frame(my.factor, I(my.list))</pre>
##
     my.factor
                  my.list
## a
         1 4, 3, 2, 1
             2 d, c, b, a
## C
             3
             2
## d
df7$my.list
## $a
## [1] 4 3 2 1
## $b
## [1] "d" "c" "b" "a"
##
## $c
## [1] "n"
##
## $d
## [1] "z"
```

What is this exercise about? Do check the documentation carefully and think of uses where the flexibility gained by use of function I() to protect arguments passed to the data.frame() constructor can be useful. In addition, write code to extract individual members of embedded matrices and lists using indexing in a single R statement in each case.

Data frames are extremely important to anyone analyzing or plotting data using R. One can think of data frames as tightly structured work-sheets, or as lists. As you may have guessed from the examples earlier in this section, there are several different ways of accessing columns, rows, and individual observations stored in a data frame. The columns can be treated as members in a list, and can be accessed both by name or index (position). When accessed by name, using \$ or double square brackets, a single column is returned as a vector or factor. In contrast to lists, data frames are always "rectangular" and for this reason the values stored can also be accessed in a way similar to how elements in a matrix are accessed, using two indexes. As we saw for vectors, indexes can be vectors of integer numbers or vectors of logical values. For columns they can, in addition, be vectors of character strings matching the names of the columns. When using indexes it is extremely important to remember that the indexes are always given **row first** in R.

Indexing of data frames can in all cases be done as if they were lists, which is preferable, as it ensures compatibility with regular R lists and with newer implementations of data-frame-like structures like those defined in package 'tibble'. Using this approach, extracting two values from the second and third positions in the first column of a.df is done as follows, using numerical indexes.

```
a.df[[1]][2:3]
## [1] 2 3
```

Or using the column name.

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```
a.df[["x"]][2:3]
## [1] 2 3
```

Matrix-like indexing is done as follows, with the first index indicating rows and the second one indicating columns. This notation allows simultaneous extraction from multiple columns, which is not possible with lists. The value returned is a "smaller" data frame.

```
a.df[2:3, 1:2]

## x y

## 2 2 a

## 3 3 a
```

If the length of the column indexing vector is one, by default the returned single column value is simplified into a vector or factor, which is not consistent with the previous example which returned a data frame. This is not only surprising in everyday use, but can be the source of bugs when coding algorithms in which the length of the second index vector cannot be guaranteed to be always more than one.

```
a.df[2:3, 1]
## [1] 2 3
```

Simplification can be prevented by overriding the default argument of parameter drop.

In contrast, the extraction operator [] of tibbles—defined in package 'tibble'—never simplifies returned one-column tibbles into vectors (see section ?? on page ?? for details on the differences between data frames and tibbles).

```
# first column, a.df[[1]] preferred
a.df[ , 1]
## [1] 1 2 3 4 5 6
# first column, a.df[["x"]] or a.df$x preferred
a.df[ , "x"]
## [1] 1 2 3 4 5 6
# first row
a.df[1, ]
## x y
         z x2 x3
## 1 1 a TRUE 6 b
# first two rows of the third and fourth columns
a.df[1:2, c(FALSE, FALSE, TRUE, TRUE, FALSE)]
##
        z x2
## 1 TRUE 6
## 2 FALSE 5
```

```
# the rows for which z is true
a.df[a.df$z , ]
## x y z x2 x3
## 1 1 a TRUE 6 b
## 3 3 a TRUE 4 b
## 5 5 a TRUE 2 b

# the rows for which x > 3 keeping all columns except the third one
a.df[a.df$x > 3, -3]
## x y x2 x3
## 4 4 a 3 b
## 5 5 a 2 b
## 6 6 a 1 b
```

As explained earlier for vectors (see section 1.10 on page 31), indexing can be present both on the right-hand side and left-hand side of an assignment. The next few examples do assignments to "cells" of a.df, either to one whole column, or individual values. The last statement in the chunk below copies a number from one location to another by using indexing of the same data frame both on the right side and left side of the assignment.

```
a.df[1, 1] \leftarrow 99
a.df
     ху
         z x2 x3
## 1 99 a TRUE 6 b
## 2 2 a FALSE 5 b
## 3 3 a TRUE 4 b
## 4 4 a FALSE 3
## 5 5 a TRUE
## 6 6 a FALSE
               1
a.df[ , 1] \leftarrow -99
a.df
      x y z x2 x3
## 1 -99 a TRUE 6 b
## 2 -99 a FALSE 5 b
## 3 -99 a TRUE 4 b
## 4 -99 a FALSE 3 b
## 5 -99 a TRUE 2 b
## 6 -99 a FALSE 1 b
a.df[["x"]] \leftarrow 123
            z x2 x3
      х у
## 1 123 a TRUE 6 b
## 2 123 a FALSE 5 b
## 3 123 a TRUE 4 b
## 4 123 a FALSE 3 b
## 5 123 a TRUE 2 b
## 6 123 a FALSE 1 b
a.df[1, 1] \leftarrow a.df[6, 4]
           z x2 x3
      ху
## 1 1 a TRUE 6 b
## 2 123 a FALSE 5 b
## 3 123 a TRUE 4 b
## 4 123 a FALSE 3 b
## 5 123 a TRUE 2 b
## 6 123 a FALSE 1 b
```

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We mentioned above that indexing by name can be done either with double square brackets, [[]], or with \$. In the first case the name of the variable or column is given as a character string, enclosed in quotation marks, or as a variable with mode character. When using \$, the name is entered as a constant, without quotation marks, and cannot be a variable.

```
x.list <- list(abcd = 123, xyzw = 789)
x.list[["abcd"]]
## [1] 123

a.var <- "abcd"
x.list[[a.var]]
## [1] 123

x.list$abcd
## [1] 123

x.list$ab
## [1] 123</pre>
```

Both in the case of lists and data frames, when using double square brackets, by default an exact match is required between the name in the object and the name used for indexing. In contrast, with \$, an unambiguous partial match is silently accepted. For interactive use, partial matching is helpful in reducing typing. However, in scripts, and especially R code in packages, it is best to avoid the use of \$ as partial matching to a wrong variable present at a later time, e.g., when someone else revises the script, can lead to very difficult-to-diagnose errors. In addition, as \$ is implemented by first attempting a match to the name and then calling [[]], using \$ for indexing can result in slightly slower performance compared to using [[]]. It is possible to set an R option so that partial matching triggers a warning, which can be very useful when debugging.

1.14.1 Operating within data frames

When the names of data frames are long, complex conditions become awkward to write using indexing—i.e., subscripts. In such cases subset() is handy because evaluation is done in the "environment" of the data frame, i.e., the names of the columns are recognized if entered directly when writing the condition. Function subset() "filters" rows, usually corresponding to observations or experimental units. The condition is computed for each row, and if it returns TRUE, the row is included in the returned data frame, and excluded if FALSE.

What is the behavior of subset() when the condition is NA? Find the answer by writing code to test this, for a case where tests for different rows return NA, TRUE and FALSE.

When calling functions that return a vector, data frame, or other structure, the extraction operators [], [[]], or \$ can be appended to the rightmost parenthesis of the function call, in the same way as to the name of a variable holding the same data.

- When do extraction operators applied to data frames return a vector or factor, and when do they return a data frame?
- In the case of subset() we can select columns directly as shown below, while for most other functions, extraction using operators [], [[]] or \$ is needed.

None of the examples in the last four code chunks alters the original data frame a.df. We can store the returned value using a new name if we want to preserve a.df unchanged, or we can assign the result to a.df, deleting in the process, the previously stored value.

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In the examples above, the names in the expression passed as the second argument to subset() were searched within ad.df and found. However, if not found in the data frame objects with matching names are searched for in the environment. There being no variable A in the data frame a.df, vector A from the environment is silently used in the chunk below resulting in a returned data frame with no rows as A > 3 returns FALSE.

```
A <- 1
subset(a.df, A > 3)
## [1] x y z
## <0 rows> (or 0-length row.names)
```

This also applies to the expression passed as argument to parameter select, here shown as a way of selecting columns based on names stored in a character vector.

The use of subset() is convenient, but more prone to bugs compared to directly using the extraction operator []. This same "cost" to achieving convenience applies to functions like attach() and with() described below. The longer time that a script is expected to be used, adapted and reused, the more careful we should be when using any of these functions. An alternative way of avoiding excessive verbosity is to keep the names of data frames short.

A frequently used way of deleting a column by name from a data frame is to assign NULL to it—i.e., in the same way as members are deleted from lists. This approach modifies a.df in place.

```
aa.df <- a.df
colnames(aa.df)
## [1] "x" "y" "z"

aa.df[["y"]] <- NULL
colnames(aa.df)
## [1] "x" "z"</pre>
```

Alternatively, we can use negative indexing to remove columns from a copy of a data frame. In this example we remove a single column. As base R does not support negative indexing by name with the extraction operator, we need to find the numerical index of the column to delete. (See the examples above using subset() with bare names to delete columns.)

Instead of using the equality test, we can use the operator %in% or function grepl() to create a logical vector useful to delete or select multiple columns in a single statement.

In the previous code chunk we deleted the last column of the data frame a.df. Here is an esoteric trick for you to first untangle how it changes the positions of columns and rows, and then for you to think how and why it can be useful to use indexing with the extraction operator [] on both sides of the assignment operator <-.

```
a.df[1:6, c(1,3)] <- a.df[6:1, c(3,1)]
a.df
```

Although in this last example we used numeric indexes to make it more interesting, in practice, especially in scripts or other code that will be reused, do use column or member names instead of positional indexes whenever possible. This makes code much more reliable, as changes elsewhere in the script could alter the order of columns and *invalidate* numerical indexes. In addition, using meaningful names makes programmers' intentions easier to understand.

1.14.2 Re-arranging columns and rows

The most direct way of changing the order of columns and/or rows in data frames (and matrices and arrays) is to use subscripting as described above. Once we know the original position and target position we can use numerical indexes on both right-hand side and left-hand side of an assignment.

⚠ When using the extraction operator [] on both the left-hand-side and right-hand-side to swap columns, the vectors or factors are swapped, while the names of the columns are not! The same applies to row names, which makes storing important information in them inconvenient and error prone.

To retain the correspondence between column naming and column contents after swapping or rearranging the columns, we need to separately move the names of the columns. This seems counter intuitive, unless we think in terms of positions being named rather than the contents of the columns being linked to the names.

```
my_data_frame.df <- data.frame(A = 1:10, B = 3)
head(my_data_frame.df, 2)</pre>
```

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```
## A B
## 1 1 3
## 2 2 3

my_data_frame.df[ , 1:2] <- my_data_frame.df[ , 2:1]
head(my_data_frame.df, 2)
## A B
## 1 3 1
## 2 3 2

colnames(my_data_frame.df)[1:2] <- colnames(my_data_frame.df)[2:1]
head(my_data_frame.df, 2)
## B A
## 1 3 1
## 2 3 2</pre>
```

Taking into account that order() returns the indexes needed to sort a vector (see page 36), we can use order() to generate the indexes needed to sort rows of a data frame. In this case, the argument to order() is usually a column of the data frame being arranged. However, any vector of suitable length, including the result of applying a function to one or more columns, can be passed as an argument to order(). Function order() is very rarely useful for sorting columns of data frames as it requires a vector across columns as input. In the case of matrix and array this approach can be applied to any of their dimensions as all their elements homogenously belong to one class.

The first task to be completed is to sort a data frame based on the values in one column, using indexing and order(). Create a new data frame and with three numeric columns with three different haphazard sequences of values. Call these columns A, B and C. 1) Sort the rows of the data frame so that the values in A are in decreasing order. 2) Sort the rows of the data frame according to increasing values of the sum of A and B without adding a new column to the data frame or storing the vector of sums in a variable. In other words, do the sorting based on sums calculated on the fly.

Repeat the tasks in the playground immediately above but using factors instead of numeric vectors as columns in the data frame. Hint: revisit the exercise on page 48 were the use of order() on factors is described.

1.14.3 Re-encoding or adding variables

It is common that some variables need to be added to an existing data frame based on existing variables, either as a computed value or based on mapping for example treatments to sample codes already in a data frame. In the second case, named vectors can be used to replace values in a variable or to add a variable to a data frame.

Mapping is possible because the length of the value returned by the extraction operator [] is given by the length of the indexing vector (see section 1.10 on page 31). Although we show toy-like examples, this approach is most useful with data frames containing many rows.

If the existing variable is a character vector or factor, we need to create a named vector with the new values as data and the existing values as names.

```
data.frame(genotype = rep(c("WT", "mutant1", "mutant2"), 2),
            value = c(1.5, 3.2, 4.5, 8.2, 7.4, 6.2))
mutant <- c(WT = FALSE, mutant1 = TRUE, mutant2 = TRUE)</pre>
my.df$mutant <- mutant[my.df$genotype]</pre>
my.df
    genotype value mutant
##
## 1
        WT 1.5 FALSE
## 2 mutant1
              3.2
## 3
     mutant2 4.5
                    TRUE
## 4
          WT 8.2 FALSE
## 5 mutant1 7.4 TRUE
## 6 mutant2 6.2 TRUE
```

If the existing variable is an integer vector, we can use a vector without names, being careful that the positions in the *mapping* vector match the values of the existing variable

```
my.df <- data.frame(individual = rep(1:3, 2),
                   value = c(1.5, 3.2, 4.5, 8.2, 7.4, 6.2))
genotype <- c("WT", "mutant1", "mutant2")</pre>
my.df$genotype <- genotype[my.df$individual]</pre>
mv.df
## individual value genotype
## 1
     1 1.5 WT
## 2
            2
                 3.2 mutant1
## 3
             3
                4.5 mutant2
## 4
             1
                 8.2
## 5
             2
                 7.4
                      mutant1
                      mutant2
## 6
             3
                 6.2
```

Add a variable named genotype to the data frame below so that for individual 4 its value is "wt", for individual 1 its value is "mutant1", and for individual 2 its value is "mutant2".

```
my.df <- data.frame(individual = rep(c(2, 4, 1), 2),
value = c(1.5, 3.2, 4.5, 8.2, 7.4, 6.2))
```

In the case of computing new values from existing variables named vectors are of limited use. Instead, variables in a data frame can be added or modified with R functions transform(), with() and within(). These functions can be thought as convenience functions as the same computations can be done using the extraction operators to access individual variables, in either the lhs, rhs or both lhs and rhs (see section 1.10 on page 31).

In the case of with() only one, possibly compound code statement is affected and this statement is passed as an argument. As before, we need to fully specify the left-hand side of the assignment. The value returned is the one returned by the statement passed as an argument, in the case of compound statements, the value returned by the last contained simple code statement to be executed.

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Consequently, if the intent is to modify the container, assignment to an individual member variable (column in this case) is required.

In this example, column A of my_data_frame.df takes precedence, and the returned value is the expected one.

```
my_data_frame.df$C <- NULL
my_data_frame.df$C <- with(my_data_frame.df, (A + B) / A)
head(my_data_frame.df, 2)
## B A C
## 1 3 1 4.0
## 2 3 2 2.5</pre>
```

In the case of within(), assignments in the argument to its second parameter affect the object returned, which is a copy of the container (In this case, a whole data frame), which still needs to be saved through assignment. Here the intention is to modify it, so we assign it back to the same name, but it could have been assigned to a different name so as not to overwrite the original data frame.

```
my_data_frame.df$C <- NULL
my_data_frame.df <- within(my_data_frame.df, C <- (A + B) / A)
head(my_data_frame.df, 2)
## B A C
## 1 3 1 4.0
## 2 3 2 2.5</pre>
```

In the example above, using within() makes little difference compared to using with() with respect to the amount of typing or clarity, but with multiple member variables being operated upon, as shown below, within() has an advantage resulting in more concise and easier to understand code.

Repeatedly pre-pending the name of a *container* such as a list or data frame to the name of each member variable being accessed can make R code verbose and difficult to understand. Functions attach() and its matching detach() allow us to change where R looks for the names of objects we include in a code statement. When using a long name for a data frame, entering a simple calculation can easily result in a difficult to read statement. (Method head() is used here to limit the displayed value to the first two rows—head() is described in section 1.17 on page 75.)

```
my_data_frame.df <- data.frame(A = 1:10, B = 3)
my_data_frame.df$C <-
    (my_data_frame.df$A + my_data_frame.df$B) / my_data_frame.df$A
head(my_data_frame.df, 2)
## A B C
## 1 1 3 4.0
## 2 2 3 2.5</pre>
```

Using attach() we can alter how R looks up names and consequently simplify the statement. With detach() we can restore the original state. It is important to remember that here we can only simplify the right-hand side of the assignment, while the "destination" of the result of the computation still needs to be fully specified on the left-hand side of the assignment operator. We include below only one statement between attach() and detach() but multiple statements are allowed. Furthermore, if variables with the same name as the columns exist in the search path, these will take precedence, something that can result in bugs or crashes, or as seen below, a message warns that variable A from the global environment will be used instead of column A of the attached my_data_frame.df. The returned value is, of course, not the desired one.

```
my_data_frame.df$C <- NULL
attach(my_data_frame.df)

## The following object is masked _by_ .GlobalEnv:

##

## A

my_data_frame.df$C <- (A + B) / A

detach(my_data_frame.df)
head(my_data_frame.df, 2)

## A B C

## 1 1 3 4

## 2 2 3 4</pre>
```

Use of attach() and detach(), which function as a pair of ON and OFF switches, can result in an undesired after-effect on name lookup if the script terminates after attach() is executed but before detach() is called, as cleanup is not automatic. In contrast, with() and within(), being self-contained, guarantee that cleanup takes place. Consequently, the usual recommendation is to give preference to the use of with() and within() over attach() and detach(). Use of these functions not only saves typing but also makes code more readable.

1.15 Attributes of R objects

R objects can have attributes. Attributes are named slots normally used to store ancillary data such as object properties. There are no restrictions on the class of what is assigned to an attribute. They are used by R itself to store things like column names in data frames and labels of factor levels. All these attributes are

visible to user code, and user code can read and write objects' attributes. However, they are rarely displayed explicitly when an object is printed. They can be also used to store metadata accompanying the data stored in an object, which is important for reproducible research and data sharing.

Attribute "comment" is meant to be set by users to store a character string—e.g., to store metadata as text together with data. As comments are frequently used, R has functions for accessing and setting comments.

```
a.df <- data.frame(x = 1:6, y = "a", z = c(TRUE, FALSE))
comment(a.df)
## NULL

comment(a.df) <- "this is stored as a comment"
comment(a.df)
## [1] "this is stored as a comment"</pre>
```

Methods like names(), dim() or levels() return values retrieved from attributes stored in R objects, and methods like names()<-, dim()<- or levels()<- set (or unset with NULL) the value of the respective attributes. Specific query and set methods do not exist for all attributes. Methods attr(), attr()<- and attributes() can be used with any attribute. With attr() we access, and with attr()<- we set individual attributes by name. With attributes() we retrieve all attributes of an object as a named list. In addition, method str() displays all components and structure of R objects including their attributes.

Continuing with the previous example, we can retrieve and set the comment using these functions. In the second statement we delete the value stored in the "comment" attribute by assigning NULL to it.

```
attr(a.df, "comment")
## [1] "this is stored as a comment"

attr(a.df, "comment") <- NULL
attr(a.df, "comment")
## NULL

comment(a.df) # same as previous line
## NULL</pre>
```

The "names" attribute of a.df was set by the data.frame() constructor when it was created above. In the next example, in the first statement we retrieve the names, and implicitly print them. In the second statement, read from right to left, we retrieve the names, convert them to upper case and save them back to the same attribute.

```
names(a.df)
## [1] "x" "y" "z"

names(a.df) <- toupper(names(a.df))
names(a.df)
## [1] "X" "Y" "z"

attr(a.df, "names") # same as previous line
## [1] "x" "Y" "z"</pre>
```

We can add a new attribute, under our own control, as long as its name does not clash with that of existing attributes.

```
attr(a.df, "my.attribute") <- "this is stored in my attribute"
attributes(a.df)
## $names
## [1] "X" "Y" "Z"
##
## $class
## [1] "data.frame"
##
## $row.names
## [1] 1 2 3 4 5 6
##
## $my.attribute
## [1] "this is stored in my attribute"</pre>
```

There is no restriction to the creation, setting, resetting and reading of attributes, but not all methods and operators that can be used to modify objects will preserve non-standard attributes. This can be a problem when using some R packages, such as some popular packages from the 'tidyverse'. So, using private attributes is a double-edged sword that usually is worthwhile considering only when designing a new class together with the corresponding methods for it. A good example of extensive use of class-specific attributes are the values returned by model fitting functions like Im() (see section ?? on page ??).

Even the class of S3 objects is stored as an attribute that is accessible as any other attribute—this is in contrast to the mode and atomic class of an object. Object-oriented programming in R is explained in section ?? on page ??.

```
numbers <- 1:10
mode(numbers)
## [1] "numeric"
class(numbers)
## [1] "integer"
attributes(numbers)
## NULL
a.factor <- factor(numbers)</pre>
mode(a.factor)
## [1] "numeric"
class(a.factor)
## [1] "factor"
attributes(a.factor)
## $levels
## [1] "1"
             "2" "3" "4" "5" "6" "7" "8" "9"
##
## $class
## [1] "factor"
```

1.16 Saving and loading data

1.16.1 Data sets in R and packages

To be able to present more meaningful examples, we need some real data. Here we use cars, one of the many data sets included in base R. Function data() is used to load data objects that are included in R or contained in packages. It is also possible to import data saved in files with *foreign* formats, defined by other software or commonly used for data exchange. Package 'foreign', included in the R distribution, as well as contributed packages make available functions capable of reading and decoding various foreign formats. How to read or import "foreign" data is discussed in R documentation in *R Data Import/Export*, and in this book, in chapter ?? starting on page ??. It is also good to keep in mind that in R, URLs (Uniform Resource Locators) are accepted as arguments to the file or path parameter of many functions (see section ?? starting on page ??).

In the next example we load data included in R as R objects by calling function data(). The loaded R object cars is a data frame.

```
data(cars)
```

Once we have a data set available, the first step is usually to explore it, and we will do this with cars in section 1.17 on page 75.

1.16.2 .rda files

By default, at the end of a session, the current workspace containing the results of your work is saved into a file called <code>.RData.</code> In addition to saving the whole workspace, it is possible to save one or more R objects present in the workspace to disk using the same file format (with file name tag <code>.rda</code> or <code>.Rda</code>). One or more objects, belonging to any mode or class can be saved into a single file using function <code>save()</code>. Reading the file restores all the saved objects into the current workspace with their original names. These files are portable across most R versions—i.e., old formats can be read and written by newer versions of R, although the newer, default format may be not readable with earlier R versions. Whether compression is used, and whether the "binary" data is encoded into ASCII characters, allowing maximum portability at the expense of increased size can be controlled by passing suitable arguments to <code>save()</code>.

We create a data frame object and then save it to a file.

We delete the data frame object and confirm that it is no longer present in the workspace.

```
rm(my.df)
ls(pattern = "my.df")
## character(0)
```

We read the file we earlier saved to restore the object.

The default format used is binary and compressed, which results in smaller files.

In the example above, only one object was saved, but one can simply give the names of additional objects as arguments. Just try saving more than one data frame to the same file. Then the data frames plus a few vectors. After creating each file, clear the workspace and then restore from the file the objects you saved.

Sometimes it is easier to supply the names of the objects to be saved as a vector of character strings passed as an argument to parameter list. One case is when wanting to save a group of objects based on their names. We can use ls() to list the names of objects matching a simple pattern or a complex regular expression. The example below does this in two steps, first saving a character vector with the names of the objects matching a pattern, and then using this saved vector as an argument to save's list parameter.

```
objcts <- ls(pattern = "*.df")
save(list = objcts, file = "my-df1.rda")</pre>
```

The two statements above can be combined into a single statement by nesting the function calls.

```
save(list = ls(pattern = "*.df"), file = "my-df1.rda")
```

Practice using different patterns with 1s(). You do not need to save the objects to a file. Just have a look at the list of object names returned.

As a coda, we show how to clean up by deleting the two files we created. Function unlink() can be used to delete any files for which the user has enough rights.

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```
unlink(c("my-df.rda", "my-df1.rda"))
```

1.16.3 .rds files

The RDS format can be used to save individual objects instead of multiple objects (usually using file name tag .rds). They are read and saved with functions readRDS() and saveRDS(), respectively. When RDS files are read, different from when RDA files are loaded, we need to assign the object read to a possibly different name for it to added to the search pass. Of course, it is also possible to use the returned object as an argument to a function or in an expression without saving it to a variable.

```
saveRDS(my.df, "my-df.rds")
```

If we read the file, by default the read R object will be printed at the console.

In the next example we assign the read object to a different name, and check that the object read is identical to the one saved.

```
my_read.df <- readRDS("my-df.rds")
identical(my.df, my_read.df)
## [1] TRUE</pre>
```

As above, we clean up by deleting the file.

```
unlink("my-df.rds")
```

1.17 Looking at data

There are several functions in R that let us obtain different views into objects. Function print() is useful for small data sets, or objects. Especially in the case of large data frames, we need to explore them step by step. In the case of named components, we can obtain their names with colnames(), rownames(), and names(). If a data frame contains many rows of observations, head() and tail() allow us to easily restrict the number of rows printed. Functions nrow() and ncol() return the number of rows and columns in the data frame (also applicable to matrices but not to lists or vectors where we use length()). As mentioned earlier, function str() concisely displays the structure of R objects.

```
class(cars)
## [1] "data.frame"
head(cars)
## speed dist
## 1
       4
        4
## 2
            10
## 3
        7
            4
           22
## 4
## 5
        8 16
## 6
        9 10
tail(cars)
     speed dist
## 45
      23 54
        24 70
## 46
## 47
       24 92
## 48
        24
            93
## 49
        24 120
## 50
        25
nrow(cars)
## [1] 50
ncol(cars)
## [1] 2
names(cars)
## [1] "speed" "dist"
colnames(cars)
## [1] "speed" "dist"
head(rownames(cars))
## [1] "1" "2" "3" "4" "5" "6"
str(cars)
## 'data.frame': 50 obs. of 2 variables:
## $ speed: num 4 4 7 7 8 9 10 10 10 11 ...
## $ dist : num 2 10 4 22 16 10 18 26 34 17 ...
```

Look up the help pages for head() and tail(), and edit the code above to print only the first two lines, or only the last three lines of cars, respectively.

The different columns of a data frame can be factors, or vectors of various modes (e.g., numeric, logical, character, etc.) (see section 1.14 on page 56). To explore the mode of the columns of cars, we can use an *apply* function. In the present case, we want to apply function class() to each column of the data frame cars. (Apply functions are described in section ?? on page ??.)

```
sapply(X = cars, FUN = class)
## speed dist
## "numeric" "numeric"
```

The statement above returns a vector of character strings, with the mode of

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each column. Each element of the vector is named according to the name of the corresponding "column" in the data frame. For this same statement to be used with any other data frame or list, we need only to substitute the name of the object, the argument to the first parameter called x, to the one of current interest.

Data set airquality contains data from air quality measurements in New York, and, being included in the R distribution, can be loaded with data(airquality). Load it, and repeat the steps above, to learn what variables (columns) it contains, their classes, the number of rows, etc.

Although the R language allows data frame columns of class matrix, their use is infrequent. On the other hand, columns belonging to class list are disallowed in data frames. The reverse is true for tibbles (described in section ?? on page ??).

Function summary() can be used to obtain a summary from objects of most R classes, including data frames. We can also use sapply(), lapply() or vapply() to apply any suitable function to individual columns.

```
summary(cars)
##
       speed
                      dist
## Min. : 4.0 Min. : 2.00
  1st Qu.:12.0 1st Qu.: 26.00
## Median: 15.0 Median: 36.00
##
   Mean
         :15.4
                 Mean : 42.98
                 3rd Qu.: 56.00
##
   3rd Qu.:19.0
          :25.0
   Max.
                 Max. :120.00
sapply(cars, range)
      speed dist
## [1,]
          4
## [2,]
          25 120
```

Obtain the summary of airquality with function summary(), but in addition, write code with an *apply* function to count the number of non-missing values in each column. Hint: using sum() on a logical vector returns the count of TRUE values as TRUE, and FALSE are transparently converted into numeric 1 and 0, respectively, when logical values are used in arithmetic expressions.

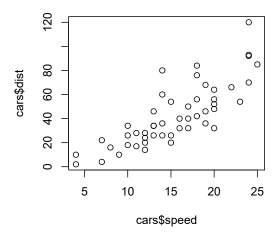
1.18 Plotting

The base-R generic method plot() can be used to plot different data. It is a generic method that has specializations suitable for different kinds of objects (see section ?? on page ?? for a brief introduction to objects, classes and methods). In this section we only very briefly demonstrate the use of the most common base-R graphics functions. They are well described in the book *R Graphics* (Murrell 2019). We will not describe the Lattice (based on S's Trellis) approach to plotting (Sarkar 2008).

Instead we describe in detail the use of the *grammar of graphics* and plotting with package 'ggplot2' in chapter ?? starting on page ??.

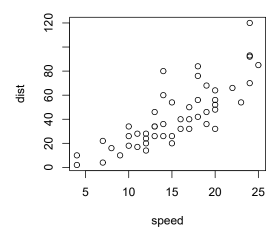
It is possible to pass two variables (here columns from a data frame) directly as arguments to the x and y parameters of \overline{plot} ().

plot(x = cars\$speed, y = cars\$dist)



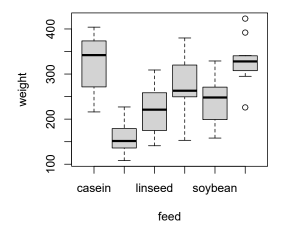
It is also possible, and usually more convenient, to use a *formula* to specify the variables to be plotted on the x and y axes, passing additionally as an argument to parameter data the name of the data frame containing these variables. The formula dist \sim speed, is read as dist explained by speed—i.e., dist is mapped to the y-axis as the dependent variable and speed to the x-axis as the independent variable.

plot(dist ~ speed, data = cars)



Within R there exist different specializations, or "flavors," of method plot() that become active depending on the class of the variables passed as arguments: passing two numerical variables results in a scatter plot as seen above. In contrast passing one factor and one numeric variable to plot() results in a box-and-whiskers plot being produced. To exemplify this we need to use a different data set, here chickwts as cars does not contain any factors. Use help("chickwts") to learn more about this data set, also included in R.

plot(weight ~ feed, data = chickwts)



Method plot() and variants defined in R, when used for plotting return their graphical output to a *graphical output device*.

When R is used interactively, a software device is opened automatically to output the graphical output to a physical device, usually the computer screen. The name of the R software device used may depend on the operating system (e.g., MS-Windows or Linux), or on the IDE (e.g., RStudio).

In R, software graphical devices not necessarily generate output on a physical device like a printer, as several of these devices translate the plotting commands into a file format and save it to disk. Several different graphical devices are available in R and they differ in the kind of output they produce: raster files (e.g., TIFF, PNG and JPEG formats), vector graphics files (e.g., SVG, EPS and PDF), or output to a physical device like the screen of a computer. Additional devices are available through contributed R packages.

Devices follow the paradigm of ON and OFF switches, opening and closing a destination for print(), plot() and related functions. Some devices producing a file as output, can save their output one plot at a time to single-page graphic files or only when the device is closed, possibly as a multi-page file.

When opening a device the user supplies additional information. For the PDF device that produces output in a vector-graphics format, width and height of the output are specified in *inches*. A default file name is used unless we pass a character string as an argument to parameter file.

```
pdf(file = "output/my-file.pdf", width = 6, height = 5, onefile = TRUE)
plot(dist ~ speed, data = cars)
plot(weight ~ feed, data = chickwts)
dev.off()
## cairo_pdf
## 2
```

Raster devices return bitmaps and width and height are specified in *pixels*.

```
png(file = "output/my-file.png", width = 600, height = 500)
plot(weight ~ feed, data = chickwts)
dev.off()
## cairo_pdf
## 2
```

The approach of direct output to a software device is used in base R, and the addition of plot components, as shown below, is done directly to the output device.

```
png(file = "output/my-file.png", width = 600, height = 500)
plot(dist ~ speed, data = cars)
text(x = 10, y = 110, labels = "some texts to be added")
dev.off()
## cairo_pdf
## 2
```

1 This is not the only approach available. As we will see in chapter **??** starting on page **??**, an alternative approach is to build a *plot object* as a list of member components that is later rendered as a whole on a graphical device by calling print() once.

1.19 Further reading

For further reading on the aspects of R discussed in the current chapter, I suggest the books *R Programming for Data Science* (Peng) and *The Art of R Programming: A Tour of Statistical Software Design* (Matloff).

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