

R Notes for Professionals

400+ pages
of professional hints and tricks

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About

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Chapter 1: Getting started with R Language

Section 1.1: Installing R

You might wish to install [RStudio](#) after you have installed R. RStudio is a development environment for R that simplifies many programming tasks.

Windows only:

[Visual Studio](#) (starting from version 2015 Update 3) now features a development environment for R called [R Tools](#), that includes a live interpreter, IntelliSense, and a debugging module. If you choose this method, you won't have to install R as specified in the following section.

For Windows

1. Go to the [CRAN](#) website, click on download R for Windows, and download the latest version of R.
2. Right-click the installer file and RUN as administrator.
3. Select the operational language for installation.
4. Follow the instructions for installation.

For OSX / macOS

Alternative 1

(0. Ensure [XQuartz](#) is installed)

1. Go to the [CRAN](#) website and download the latest version of R.
2. Open the disk image and run the installer.
3. Follow the instructions for installation.

This will install both R and the R-MacGUI. It will put the GUI in the /Applications/ Folder as R.app where it can either be double-clicked or dragged to the Doc. When a new version is released, the (re)-installation process will overwrite R.app but prior major versions of R will be maintained. The actual R code will be in the /Library/Frameworks/R.Framework/Versions/ directory. Using R within RStudio is also possible and would be using the same R code with a different GUI.

Alternative 2

1. Install homebrew (the missing package manager for macOS) by following the instructions on <https://brew.sh/>
2. `brew install R`

Those choosing the second method should be aware that the maintainer of the Mac fork advises against it, and will not respond to questions about difficulties on the R-SIG-Mac Mailing List.

For Debian, Ubuntu and derivatives

You can get the version of R corresponding to your distro via `apt-get`. However, this version will frequently be quite far behind the most recent version available on CRAN. You can add CRAN to your list of recognized "sources".

```
sudo apt-get install r-base
```

You can get a more recent version directly from CRAN by adding CRAN to your sources list. Follow the [directions](#) from CRAN for more details. Note in particular the need to also execute this so that you can use

`install.packages()`. Linux packages are usually distributed as source files and need compilation:

```
sudo apt-get install r-base-dev
```

For Red Hat and Fedora

```
sudo dnf install R
```

For Archlinux

R is directly available in the Extra package repo.

```
sudo pacman -S r
```

More info on using R under Archlinux can be found on the [ArchWiki R page](#).

Section 1.2: Hello World!

```
"Hello World!"
```

Also, check out the detailed discussion of how, when, whether and why to print a string.

Section 1.3: Getting Help

You can use function `help()` or `?` to access documentations and search for help in R. For even more general searches, you can use `help.search()` or `??`.

```
#For help on the help function of R
help()

#For help on the paste function
help(paste)      #OR
help("paste")    #OR
?paste           #OR
?"paste"
```

Visit <https://www.r-project.org/help.html> for additional information

Section 1.4: Interactive mode and R scripts

The interactive mode

The most basic way to use R is the *interactive* mode. You type commands and immediately get the result from R.

Using R as a calculator

Start R by typing R at the command prompt of your operating system or by executing RGui on Windows. Below you can see a screenshot of an interactive R session on Linux:

```
user:~$ R

R version 3.3.2 (2016-10-31) -- "Sincere Pumpkin Patch"
Copyright (C) 2016 The R Foundation for Statistical Computing
Platform: x86_64-pc-linux-gnu (64-bit)

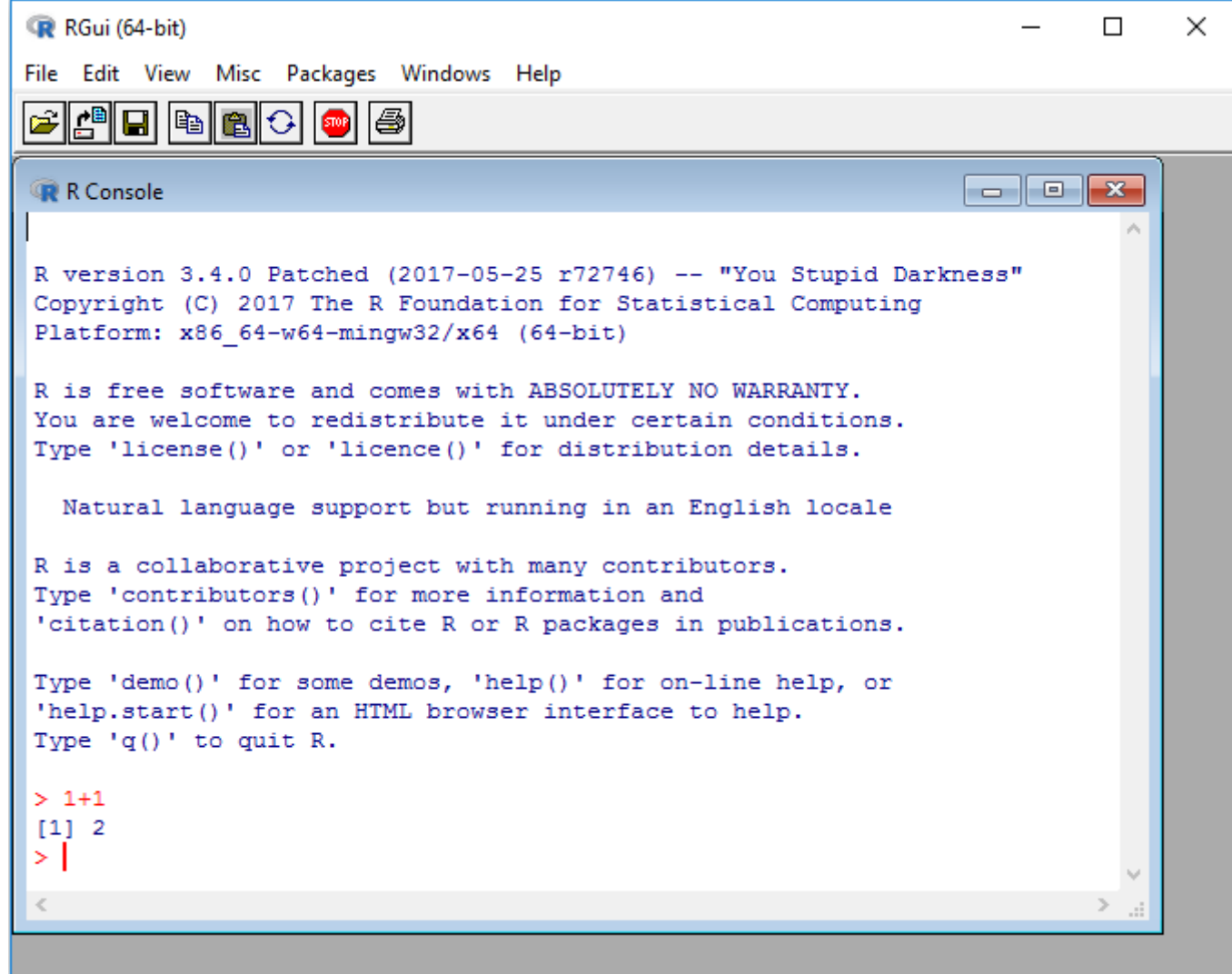
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Tippen Sie 'contributors()' für mehr Information und 'citation()',
um zu erfahren, wie R oder R packages in Publikationen zitiert werden können.

Tippen Sie 'demo()' für einige Demos, 'help()' für on-line Hilfe, oder
'help.start()' für eine HTML Browserschnittstelle zur Hilfe.
Tippen Sie 'q()', um R zu verlassen.

> 1+1
[1] 2
> |
```

This is RGui on Windows, the most basic working environment for R under Windows:



After the > sign, expressions can be typed in. Once an expression is typed, the result is shown by R. In the screenshot above, R is used as a calculator: Type

to immediately see the result, 2. The leading [1] indicates that R returns a vector. In this case, the vector contains only one number (2).

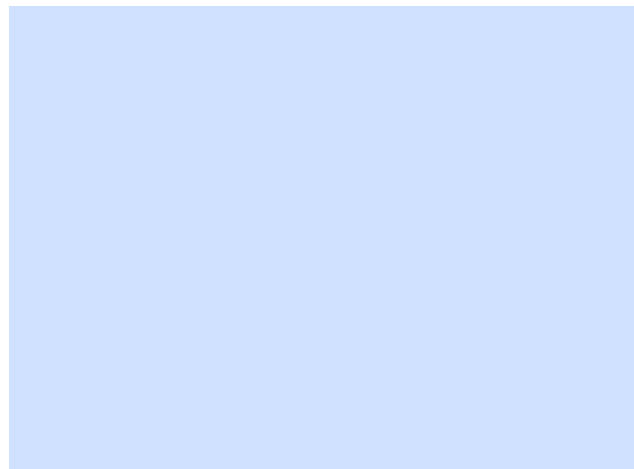
The first plot

R can be used to generate plots. The following example uses the data set **PlantGrowth**, which comes as an example data set along with R

Type int the following all lines into the R prompt which do not start with ##. Lines starting with ## are meant to document the result which R will return.

```
data(PlantGrowth)
str(PlantGrowth)
## 'data.frame':   30 obs. of  2 variables:
## $ weight: num  4.17 5.58 5.18 6.11 4.5 4.61 5.17 4.53 5.33 5.14 ...
## $ group : Factor w/ 3 levels "ctrl","trt1",...: 1 1 1 1 1 1 1 1 1 1 ...
anova(lm(weight ~ group, data = PlantGrowth))
## Analysis of Variance Table
##
## Response: weight
##             Df  Sum Sq Mean Sq F value    Pr(>F)
## group        2  3.7663  1.8832  4.8461 0.01591 *
## Residuals  27 10.4921  0.3886
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
boxplot(weight ~ group, data = PlantGrowth, ylab = "Dry weight")
```

The following plot is created:



data(PlantGrowth) loads the example data set **PlantGrowth**, which is records of dry masses of plants which were subject to two different treatment conditions or no treatment at all (control group). The data set is made available under the name **PlantGrowth**. Such a name is also called a Variable.

To load your own data, the following two documentation pages might be helpful:

- Reading and writing tabular data in plain-text files (CSV, TSV, etc.)
- I/O for foreign tables (Excel, SAS, SPSS, Stata)

str(PlantGrowth) shows information about the data set which was loaded. The output indicates that **PlantGrowth** is a **data.frame**, which is R's name for a table. The **data.frame** contains of two columns and 30 rows. In this case, each row corresponds to one plant. Details of the two columns are shown in the lines starting with \$: The first

column is called `weight` and contains numbers (`num`, the dry weight of the respective plant). The second column, `group`, contains the treatment that the plant was subjected to. This is categorial data, which is called **factor** in R. Read more information about data frames.

To compare the dry masses of the three different groups, a one-way ANOVA is performed using `anova(lm(...))`. `weight ~ group` means "Compare the values of the column `weight`, grouping by the values of the column `group`". This is called a Formula in R. `data = ...` specifies the name of the table where the data can be found.

The result shows, among others, that there exists a significant difference (Column $\text{Pr}(>F)$), $p = 0.01591$) between some of the three groups. Post-hoc tests, like Tukey's Test, must be performed to determine which groups' means differ significantly.

`boxplot(...)` creates a box plot of the data. where the values to be plotted come from. `weight ~ group` means: "Plot the values of the column `weight` versus the values of the column `group`. `ylab = ...` specifies the label of the y axis. More information: Base plotting

Type `q()` or `[Ctrl]-[D]` to exit from the R session.

R scripts

To document your research, it is favourable to save the commands you use for calculation in a file. For that effect, you can create **R scripts**. An R script is a simple text file, containing R commands.

Create a text file with the name `plants.R`, and fill it with the following text, where some commands are familiar from the code block above:

```
data(PlantGrowth)  
  
anova(lm(weight ~ group, data = PlantGrowth))  
  
png("plant_boxplot.png", width = 400, height = 300)  
boxplot(weight ~ group, data = PlantGrowth, ylab = "Dry weight")  
dev.off()
```

Execute the script by typing into your terminal (The terminal of your operating system, **not** an interactive R session like in the previous section!)

```
R --no-save <plant.R >plant_result.txt
```

The file `plant_result.txt` contains the results of your calculation, as if you had typed them into the interactive R prompt. Thereby, your calculations are documented.

The new commands `png` and `dev.off` are used for saving the boxplot to disk. The two commands must enclose the plotting command, as shown in the example above. `png("FILENAME", width = ..., height = ...)` opens a new PNG file with the specified file name, width and height in pixels. `dev.off()` will finish plotting and saves the plot to disk. No output is saved until `dev.off()` is called.

Chapter 2: Variables

Section 2.1: Variables, data structures and basic Operations

In R, data objects are manipulated using named data structures. The names of the objects might be called "variables" although that term does not have a specific meaning in the official R documentation. R names are *case sensitive* and may contain alphanumeric characters(a-z,A-Z,0-9), the dot/period(.) and underscore(_). To create names for the data structures, we have to follow the following rules:

- Names that start with a digit or an underscore (e.g. 1a), or names that are valid numerical expressions (e.g. .11), or names with dashes ('-') or spaces can only be used when they are quoted: `1a` and `.11`. The names will be printed with backticks:

```
list( '.11' = "a")
#$`11`
#[1] "a"
```

- All other combinations of alphanumeric characters, dots and underscores can be used freely, where reference with or without backticks points to the same object.
- Names that begin with . are considered system names and are not always visible using the `ls()`-function.

There is no restriction on the number of characters in a variable name.

Some examples of valid object names are: foobar, foo.bar, foo_bar, .foobar

In R, variables are assigned values using the infix-assignment operator `<-`. The operator `=` can also be used for assigning values to variables, however its proper use is for associating values with parameter names in function calls. Note that omitting spaces around operators may create confusion for users. The expression `a<-1` is parsed as assignment (`a <- 1`) rather than as a logical comparison (`a < -1`).

```
> foo <- 42
> fooEquals = 43
```

So `foo` is assigned the value of 42. Typing `foo` within the console will output 42, while typing `fooEquals` will output 43.

```
> foo
[1] 42
> fooEquals
[1] 43
```

The following command assigns a value to the variable named `x` and prints the value simultaneously:

```
> (x <- 5)
[1] 5
# actually two function calls: first one to `<-`; second one to the `()`-function
> is.function(``)
[1] TRUE # Often used in R help page examples for its side-effect of printing.
```

It is also possible to make assignments to variables using `->`.

```
> 5 -> x
> x
```

```
[1] 5
```

```
>
```

Types of data structures

There are no scalar data types in R. Vectors of length-one act like scalars.

- Vectors: Atomic vectors must be sequence of same-class objects.: a sequence of numbers, or a sequence of logicals or a sequence of characters. `v <- c(2, 3, 7, 10)`, `v2 <- c("a", "b", "c")` are both vectors.
- Matrices: A matrix of numbers, logical or characters. `a <- matrix(data = c(1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12), nrow = 4, ncol = 3, byrow = F)`. Like vectors, matrix must be made of same-class elements. To extract elements from a matrix rows and columns must be specified: `a[1,2]` returns [1] 5 that is the element on the first row, second column.
- Lists: concatenation of different elements `mylist <- list(course = 'stat', date = '04/07/2009', num_isc = 7, num_cons = 6, num_mat = as.character(c(45020, 45679, 46789, 43126, 42345, 47568, 45674)), results = c(30, 19, 29, NA, 25, 26, 27))`. Extracting elements from a list can be done by name (if the list is named) or by index. In the given example `mylist$results` and `mylist[[6]]` obtains the same element. Warning: if you try `mylist[6]`, R won't give you an error, but it extract the result as a list. While `mylist[[6]][2]` is permitted (it gives you 19), `mylist[6][2]` gives you an error.
- data.frame: object with columns that are vectors of equal length, but (possibly) different types. They are not matrices. `exam <- data.frame(matr = as.character(c(45020, 45679, 46789, 43126, 42345, 47568, 45674)), res_S = c(30, 19, 29, NA, 25, 26, 27), res_O = c(3, 3, 1, NA, 3, 2, NA), res_TOT = c(30, 22, 30, NA, 28, 28, 27))`. Columns can be read by name `exam$matr`, `exam[, 'matr']` or by index `exam[1]`, `exam[, 1]`. Rows can also be read by name `exam['rowname']`, or index `exam[1,]`. Dataframes are actually just lists with a particular structure (rownames-attribute and equal length components)

Common operations and some cautionary advice

Default operations are done element by element. See `?Syntax` for the rules of operator precedence. Most operators (and may other functions in base R) have recycling rules that allow arguments of unequal length. Given these objects:

Example objects

```
> a <- 1
> b <- 2
> c <- c(2,3,4)
> d <- c(10,10,10)
> e <- c(1,2,3,4)
> f <- 1:6
> W <- cbind(1:4, 5:8, 9:12)
> Z <- rbind(rep(0,3), 1:3, rep(10,3), c(4,7,1))
```

Some vector operations

```
> a+b # scalar + scalar
[1] 3
> c+d # vector + vector
[1] 12 13 14
> a*b # scalar * scalar
[1] 2
> c*d # vector * vector (componentwise!)
[1] 20 30 40
> c+a # vector + scalar
[1] 3 4 5
> c^2 #
[1] 4 9 16
> exp(c)
```

```
[1] 7.389056 20.085537 54.598150
```

Some vector operation Warnings!

```
> c+e # warning but.. no errors, since recycling is assumed to be desired.  
[1] 3 5 7 6  
Warning message:  
In c + e : longer object length is not a multiple of shorter object length
```

R sums what it can and then reuses the shorter vector to fill in the blanks... The warning was given only because the two vectors have lengths that are not exactly multiples. c+f # no warning whatsoever.

Some Matrix operations Warning!

```
> Z+W # matrix + matrix #(componentwise)  
> Z*W # matrix* matrix#(Standard product is always componentwise)
```

To use a matrix multiply: V %*% W

```
> W + a # matrix+ scalar is still componentwise  
[,1] [,2] [,3]  
[1,] 2 6 10  
[2,] 3 7 11  
[3,] 4 8 12  
[4,] 5 9 13  
  
> W + c # matrix + vector... : no warnings and R does the operation in a column-wise manner  
[,1] [,2] [,3]  
[1,] 3 8 13  
[2,] 5 10 12  
[3,] 7 9 14  
[4,] 6 11 16
```

"Private" variables

A leading dot in a name of a variable or function in R is commonly used to denote that the variable or function is meant to be hidden.

So, declaring the following variables

```
> foo <- 'foo'  
> .foo <- 'bar'
```

And then using the ls function to list objects will only show the first object.

```
> ls()  
[1] "foo"
```

However, passing **all.names** = TRUE to the function will show the 'private' variable

```
> ls(all.names = TRUE)  
[1] ".foo" "foo"
```

Chapter 3: Arithmetic Operators

Section 3.1: Range and addition

Let's take an example of adding a value to a range (as it could be done in a loop for example):

```
3+1:5
```

Gives:

```
[1] 4 5 6 7 8
```

This is because the range operator `:` has higher precedence than addition operator `+`.

What happens during evaluation is as follows:

- `3+1:5`
- `3+c(1, 2, 3, 4, 5)` expansion of the range operator to make a vector of integers.
- `c(4, 5, 6, 7, 8)` Addition of 3 to each member of the vector.

To avoid this behavior you have to tell the R interpreter how you want it to order the operations with `()` like this:

```
(3+1):5
```

Now R will compute what is inside the parentheses before expanding the range and gives:

```
[1] 4 5
```

Section 3.2: Addition and subtraction

The basic math operations are performed mainly on numbers or on vectors (lists of numbers).

1. Using single numbers

We can simple enter the numbers concatenated with `+` for *adding* and `-` for *subtracting*:

```
> 3 + 4.5
# [1] 7.5
> 3 + 4.5 + 2
# [1] 9.5
> 3 + 4.5 + 2 - 3.8
# [1] 5.7
> 3 + NA
#[1] NA
> NA + NA
#[1] NA
> NA - NA
#[1] NA
> NaN - NA
#[1] NaN
> NaN + NA
#[1] NaN
```

We can assign the numbers to *variables* (constants in this case) and do the same operations:

```

> a <- 3; B <- 4.5; cc <- 2; Dd <- 3.8 ;na<-NA;nan<-NaN
> a + B
# [1] 7.5
> a + B + cc
# [1] 9.5
> a + B + cc - Dd
# [1] 5.7
> B-nan
#[1] NaN
> a+na-na
#[1] NA
> a + na
#[1] NA
> B-nan
#[1] NaN
> a+na-na
#[1] NA

```

2. Using vectors

In this case we create vectors of numbers and do the operations using those vectors, or combinations with single numbers. In this case the operation is done considering each element of the vector:

```

> A <- c(3, 4.5, 2, -3.8);
> A
# [1] 3.0 4.5 2.0 -3.8
> A + 2 # Adding a number
# [1] 5.0 6.5 4.0 -1.8
> 8 - A # number less vector
# [1] 5.0 3.5 6.0 11.8
> n <- length(A) #number of elements of vector A
> n
# [1] 4
> A[-n] + A[n] # Add the last element to the same vector without the last element
# [1] -0.8 0.7 -1.8
> A[1:2] + 3 # vector with the first two elements plus a number
# [1] 6.0 7.5
> A[1:2] - A[3:4] # vector with the first two elements less the vector with elements 3 and 4
# [1] 1.0 8.3

```

We can also use the function **sum** to add all elements of a vector:

```

> sum(A)
# [1] 5.7
> sum(-A)
# [1] -5.7
> sum(A[-n]) + A[n]
# [1] 5.7

```

We must take care with *recycling*, which is one of the characteristics of R, a behavior that happens when doing math operations where the length of vectors is different. *Shorter vectors in the expression are recycled as often as need be (perhaps fractionally) until they match the length of the longest vector. In particular a constant is simply repeated.* In this case a Warning is show.

```

> B <- c(3, 5, -3, 2.7, 1.8)
> B
# [1] 3.0 5.0 -3.0 2.7 1.8
> A
# [1] 3.0 4.5 2.0 -3.8

```

```

> A + B # the first element of A is repeated
# [1] 6.0 9.5 -1.0 -1.1 4.8
Warning message:
In A + B : longer object length is not a multiple of shorter object length
> B - A # the first element of A is repeated
# [1] 0.0 0.5 -5.0 6.5 -1.2
Warning message:
In B - A : longer object length is not a multiple of shorter object length

```

In this case the correct procedure will be to consider only the elements of the shorter vector:

```

> B[1:n] + A
# [1] 6.0 9.5 -1.0 -1.1
> B[1:n] - A
# [1] 0.0 0.5 -5.0 6.5

```

When using the **sum** function, again all the elements inside the function are added.

```

> sum(A, B)
# [1] 15.2
> sum(A, -B)
# [1] -3.8
> sum(A)+sum(B)
# [1] 15.2
> sum(A)-sum(B)
# [1] -3.8

```

Chapter 4: Matrices

Matrices store data

Section 4.1: Creating matrices

Under the hood, a matrix is a special kind of vector with two dimensions. Like a vector, a matrix can only have one data class. You can create matrices using the `matrix` function as shown below.

```
matrix(data = 1:6, nrow = 2, ncol = 3)
##      [,1] [,2] [,3]
## [1,]    1    3    5
## [2,]    2    4    6
```

As you can see this gives us a matrix of all numbers from 1 to 6 with two rows and three columns. The `data` parameter takes a vector of values, `nrow` specifies the number of rows in the matrix, and `ncol` specifies the number of columns. By convention the matrix is filled by column. The default behavior can be changed with the `byrow` parameter as shown below:

```
matrix(data = 1:6, nrow = 2, ncol = 3, byrow = TRUE)
##      [,1] [,2] [,3]
## [1,]    1    2    3
## [2,]    4    5    6
```

Matrices do not have to be numeric – any vector can be transformed into a matrix. For example:

```
matrix(data = c(TRUE, TRUE, TRUE, FALSE, FALSE, FALSE), nrow = 3, ncol = 2)
##      [,1] [,2]
## [1,] TRUE FALSE
## [2,] TRUE FALSE
## [3,] TRUE FALSE
matrix(data = c("a", "b", "c", "d", "e", "f"), nrow = 3, ncol = 2)
##      [,1] [,2]
## [1,] "a"  "d"
## [2,] "b"  "e"
## [3,] "c"  "f"
```

Like vectors matrices can be stored as variables and then called later. The rows and columns of a matrix can have names. You can look at these using the functions `rownames` and `colnames`. As shown below, the rows and columns don't initially have names, which is denoted by `NULL`. However, you can assign values to them.

```
mat1 <- matrix(data = 1:6, nrow = 2, ncol = 3, byrow = TRUE)
rownames(mat1)
## NULL
colnames(mat1)
## NULL
rownames(mat1) <- c("Row 1", "Row 2")
colnames(mat1) <- c("Col 1", "Col 2", "Col 3")
mat1
##      Col 1 Col 2 Col 3
## Row 1    1    2    3
## Row 2    4    5    6
```

It is important to note that similarly to vectors, matrices can only have one data type. If you try to specify a matrix with multiple data types the data will be coerced to the higher order data class.

The `class`, `is`, and `as` functions can be used to check and coerce data structures in the same way they were used on the vectors in class 1.

```
class(mat1)
## [1] "matrix"
is.matrix(mat1)
## [1] TRUE
as.vector(mat1)
## [1] 1 4 2 5 3 6
```

Chapter 5: Formula

Section 5.1: The basics of formula

Statistical functions in R make heavy use of the so-called Wilkinson-Rogers formula notation¹.

When running model functions like `lm` for the Linear Regressions, they need a `formula`. This `formula` specifies which regression coefficients shall be estimated.

```
my_formula1 <- formula(mpg ~ wt)
class(my_formula1)
# gives "formula"

mod1 <- lm(my_formula1, data = mtcars)
coef(mod1)
# gives (Intercept)           wt
#             37.285126   -5.344472
```

On the left side of the `~` (LHS) the dependent variable is specified, while the right hand side (RHS) contains the independent variables. Technically the `formula` call above is redundant because the tilde-operator is an infix function that returns an object with `formula` class:

```
form <- mpg ~ wt
class(form)
#[1] "formula"
```

The advantage of the `formula` function over `~` is that it also allows an environment for evaluation to be specified:

```
form_mt <- formula(mpg ~ wt, env = mtcars)
```

In this case, the output shows that a regression coefficient for `wt` is estimated, as well as (per default) an intercept parameter. The intercept can be excluded / forced to be 0 by including `0` or `-1` in the `formula`:

```
coef(lm(mpg ~ 0 + wt, data = mtcars))
coef(lm(mpg ~ wt -1, data = mtcars))
```

Interactions between variables `a` and `b` can be added by including `a:b` to the `formula`:

```
coef(lm(mpg ~ wt:vs, data = mtcars))
```

As it is (from a statistical point of view) generally advisable not have interactions in the model without the main effects, the naive approach would be to expand the `formula` to `a + b + a:b`. This works but can be simplified by writing `a*b`, where the `*` operator indicates factor crossing (when between two factor columns) or multiplication when one or both of the columns are 'numeric':

```
coef(lm(mpg ~ wt*vs, data = mtcars))
```

Using the `*` notation expands a term to include all lower order effects, such that:

```
coef(lm(mpg ~ wt*vs*hp, data = mtcars))
```

will give, in addition to the intercept, 7 regression coefficients. One for the three-way interaction, three for the two-way interactions and three for the main effects.

If one wants, for example, to exclude the three-way interaction, but retain all two-way interactions there are two shorthands. First, using - we can subtract any particular term:

```
coef(lm(mpg ~ wt*vs*hp - wt:vs:hp, data = mtcars))
```

Or, we can use the ^ notation to specify which level of interaction we require:

```
coef(lm(mpg ~ (wt + vs + hp) ^ 2, data = mtcars))
```

Those two formula specifications should create the same model matrix.

Finally, . is shorthand to use all available variables as main effects. In this case, the **data** argument is used to obtain the available variables (which are not on the LHS). Therefore:

```
coef(lm(mpg ~ ., data = mtcars))
```

gives coefficients for the intercept and 10 independent variables. This notation is frequently used in machine learning packages, where one would like to use all variables for prediction or classification. Note that the meaning of . depends on context (see e.g. [?update.formula](#) for a different meaning).

1. G. N. Wilkinson and C. E. Rogers. *Journal of the Royal Statistical Society. Series C (Applied Statistics)* Vol. 22, No. 3 (1973), pp. 392-399

Chapter 6: Reading and writing strings

Section 6.1: Printing and displaying strings

R has several built-in functions that can be used to print or display information, but `print` and `cat` are the most basic. As R is an [interpreted language](#), you can try these out directly in the R console:

```
print("Hello World")
#[1] "Hello World"
cat("Hello World\n")
#Hello World
```

Note the difference in both input and output for the two functions. (Note: there are no quote-characters in the value of x created with `x <- "Hello World"`. They are added by `print` at the output stage.)

`cat` takes one or more character vectors as arguments and prints them to the console. If the character vector has a length greater than 1, arguments are separated by a space (by default):

```
cat(c("hello", "world", "\n"))
#hello world
```

Without the new-line character (\n) the output would be:

```
cat("Hello World")
#Hello World>
```

The prompt for the next command appears immediately after the output. (Some consoles such as RStudio's may automatically append a newline to strings that do not end with a newline.)

`print` is an example of a "generic" function, which means the class of the first argument passed is detected and a class-specific *method* is used to output. For a character vector like `"Hello World"`, the result is similar to the output of `cat`. However, the character string is quoted and a number [1] is output to indicate the first element of a character vector (In this case, the first and only element):

```
print("Hello World")
#[1] "Hello World"
```

This default print method is also what we see when we simply ask R to print a variable. Note how the output of typing s is the same as calling `print(s)` or `print("Hello World")`:

```
s <- "Hello World"
s
#[1] "Hello World"
```

Or even without assigning it to anything:

```
"Hello World"
#[1] "Hello World"
```

If we add another character string as a second element of the vector (using the `c()` function to concatenate the elements together), then the behavior of `print()` looks quite a bit different from that of `cat`:

```
print(c("Hello World", "Here I am.))
```

```
#[1] "Hello World" "Here I am."
```

Observe that the `c()` function does *not* do string-concatenation. (One needs to use `paste` for that purpose.) R shows that the character vector has two elements by quoting them separately. If we have a vector long enough to span multiple lines, R will print the index of the element starting each line, just as it prints [1] at the start of the first line.

```
c("Hello World", "Here I am!", "This next string is really long.")
#[1] "Hello World"                 "Here I am!"
#[3] "This next string is really long."
```

The particular behavior of `print` depends on the *class* of the object passed to the function.

If we call `print` an object with a different class, such as "numeric" or "logical", the quotes are omitted from the output to indicate we are dealing with an object that is not character class:

```
print(1)
#[1] 1
print(TRUE)
#[1] TRUE
```

Factor objects get printed in the same fashion as character variables which often creates ambiguity when console output is used to display objects in SO question bodies. It is rare to use `cat` or `print` except in an interactive context. Explicitly calling `print()` is particularly rare (unless you wanted to suppress the appearance of the quotes or view an object that is returned as `invisible` by a function), as entering `foo` at the console is a shortcut for `print(foo)`. The interactive console of R is known as a REPL, a "read-eval-print-loop". The `cat` function is best saved for special purposes (like writing output to an open file connection). Sometimes it is used inside functions (where calls to `print()` are suppressed), however **using `cat()` inside a function to generate output to the console is bad practice**. The preferred method is to `message()` or `warning()` for intermediate messages; they behave similarly to `cat` but can be optionally suppressed by the end user. The final result should simply be returned so that the user can assign it to store it if necessary.

```
message("hello world")
#hello world
suppressMessages(message("hello world"))
```

Section 6.2: Capture output of operating system command

Functions which return a character vector

Base R has two functions for invoking a system command. Both require an additional parameter to capture the output of the system command.

```
system("top -a -b -n 1", intern = TRUE)
system2("top", "-a -b -n 1", stdout = TRUE)
```

Both return a character vector.

```
[1] "top - 08:52:03 up 70 days, 15:09, 0 users, load average: 0.00, 0.00, 0.00"
[2] "Tasks: 125 total, 1 running, 124 sleeping, 0 stopped, 0 zombie"
[3] "Cpu(s): 0.9%us, 0.3%sy, 0.0%ni, 98.7%id, 0.1%wa, 0.0%hi, 0.0%si, 0.0%st"
[4] "Mem: 12194312k total, 3613292k used, 8581020k free, 216940k buffers"
[5] "Swap: 12582908k total, 2334156k used, 10248752k free, 1682340k cached"
[6] ""
[7] " PID USER      PR NI VIRT  RES  SHR S %CPU %MEM     TIME+ COMMAND"
```

```
[8] "11300 root      20   0 1278m 375m 3696 S  0.0  3.2 124:40.92 trala
[9] " 6093 user1    20   0 1817m 269m 1888 S  0.0  2.3 12:17.96 R
[10] " 4949 user2    20   0 1917m 214m 1888 S  0.0  1.8 11:16.73 R
```

For illustration, the UNIX command `top -a -b -n 1` is used. This is OS specific and may need to be amended to run the examples on your computer.

Package `devtools` has a function to run a system command and capture the output without an additional parameter. It also returns a character vector.

```
devtools::system_output("top", "-a -b -n 1")
```

Functions which return a data frame

The `fread` function in package `data.table` allows to execute a shell command and to read the output like `read.table`. It returns a `data.table` or a `data.frame`.

```
fread("top -a -b -n 1", check.names = TRUE)
     PID    USER PR NI VIRT  RES  SHR S X.CPU X.MEM      TIME. COMMAND
1: 11300    root 20  0 1278m 375m 3696 S    0   3.2 124:40.92  trala
2:  6093   user1 20  0 1817m 269m 1888 S    0   2.3 12:18.56      R
3:  4949   user2 20  0 1917m 214m 1888 S    0   1.8 11:17.33      R
4:  7922   user3 20  0 3094m 131m 1892 S    0   1.1 21:04.95      R
```

Note, that `fread` automatically has skipped the top 6 header lines.

Here the parameter `check.names = TRUE` was added to convert %CPU, %MEM, and TIME+ to syntactically valid column names.

Section 6.3: Reading from or writing to a file connection

Not always we have liberty to read from or write to a local system path. For example if R code streaming map-reduce must need to read and write to file connection. There can be other scenarios as well where one is going beyond local system and with advent of cloud and big data, this is becoming increasingly common. One of the way to do this is in logical sequence.

Establish a file connection to read with `file()` command ("r" is for read mode):

```
conn <- file("/path/example.data", "r") #when file is in local system
conn1 <- file("stdin", "r") #when just standard input/output for files are available
```

As this will establish just file connection, one can read the data from these file connections as follows:

```
line <- readLines(conn, n=1, warn=FALSE)
```

Here we are reading the data from file connection `conn` line by line as `n=1`. one can change value of `n` (say 10, 20 etc.) for reading data blocks for faster reading (10 or 20 lines block read in one go). To read complete file in one go set `n=-1`.

After data processing or say model execution; one can write the results back to file connection using many different commands like `writeLines()`, `cat()` etc. which are capable of writing to a file connection. However all of these commands will leverage file connection established for writing. This could be done using `file()` command as:

```
conn2 <- file("/path/result.data", "w") #when file is in local system  
conn3 <- file("stdout", "w") #when just standard input/output for files are available
```

Then write the data as follows:

```
writeLines("text", conn2, sep = "\n")
```

Chapter 7: String manipulation with stringi package

Section 7.1: Count pattern inside string

With fixed pattern

```
stri_count_fixed("babab", "b")
# [1] 3
stri_count_fixed("babab", "ba")
# [1] 2
stri_count_fixed("babab", "bab")
# [1] 1
```

Natively:

```
length(gregexpr("b", "babab")[[1]])
# [1] 3
length(gregexpr("ba", "babab")[[1]])
# [1] 2
length(gregexpr("bab", "babab")[[1]])
# [1] 1
```

function is vectorized over string and pattern:

```
stri_count_fixed("babab", c("b", "ba"))
# [1] 3 2
stri_count_fixed(c("babab", "bbb", "bca", "abc"), c("b", "ba"))
# [1] 3 0 1 0
```

A base R solution:

```
sapply(c("b", "ba"), function(x) length(gregexpr(x, "babab")[[1]]))
# b ba
# 3 2
```

With regex

First example - find a and any character after

Second example - find a and any digit after

```
stri_count_regex("a1 b2 a3 b4 aa", "a.")
# [1] 3
stri_count_regex("a1 b2 a3 b4 aa", "a\\d")
# [1] 2
```

Section 7.2: Duplicating strings

```
stri_dup("abc", 3)
# [1] "abcabcabc"
```

A base R solution that does the same would look like this:

```
paste0(rep("abc",3),collapse = "")  
# [1] "abcabcabc"
```

Section 7.3: Paste vectors

```
stri_paste(LETTERS, "-", 1:13)  
# [1] "A-1"   "B-2"   "C-3"   "D-4"   "E-5"   "F-6"   "G-7"   "H-8"   "I-9"   "J-10"  "K-11"  "L-12"  "M-13"  
# [14] "N-1"   "O-2"   "P-3"   "Q-4"   "R-5"   "S-6"   "T-7"   "U-8"   "V-9"   "W-10"  "X-11"  "Y-12"  "Z-13"
```

Natively, we could do this in R via:

```
> paste(LETTERS,1:13,sep="-")  
# [1] "A-1"   "B-2"   "C-3"   "D-4"   "E-5"   "F-6"   "G-7"   "H-8"   "I-9"   "J-10"  "K-11"  "L-12"  "M-13"  
# [14] "N-1"   "O-2"   "P-3"   "Q-4"   "R-5"   "S-6"   "T-7"   "U-8"   "V-9"   "W-10"  "X-11"  "Y-12"  "Z-13"
```

Section 7.4: Splitting text by some fixed pattern

Split vector of texts using one pattern:

```
stri_split_fixed(c("To be or not to be.", "This is very short sentence."), " ")  
# [[1]]  
# [1] "To"   "be"   "or"   "not"  "to"   "be."  
#  
# [[2]]  
# [1] "This"      "is"       "very"      "short"     "sentence."
```

Split one text using many patterns:

```
stri_split_fixed("Apples, oranges and pineapllies.",c(" ", ",", ".", "s"))  
# [[1]]  
# [1] "Apples,"      "oranges"      "and"        "pineapllies."  
#  
# [[2]]  
# [1] "Apples"          " oranges and pineapllies."  
#  
# [[3]]  
# [1] "Apple"         ", orange"      " and pineaplle"  "."
```

Chapter 8: Classes

The class of a data-object determines which functions will process its contents. The `class`-attribute is a character vector, and objects can have zero, one or more classes. If there is no class-attribute, there will still be an implicit class determined by an object's `mode`. The class can be inspected with the function `class` and it can be set or modified by the `class<-` function. The S3 class system was established early in S's history. The more complex S4 class system was established later

Section 8.1: Inspect classes

Every object in R is assigned a class. You can use `class()` to find the object's class and `str()` to see its structure, including the classes it contains. For example:

```
class(iris)
[1] "data.frame"

str(iris)
'data.frame': 150 obs. of 5 variables:
 $ Sepal.Length: num 5.1 4.9 4.7 4.6 5 5.4 4.6 5 4.4 4.9 ...
 $ Sepal.Width : num 3.5 3 3.2 3.1 3.6 3.9 3.4 3.4 2.9 3.1 ...
 $ Petal.Length: num 1.4 1.4 1.3 1.5 1.4 1.7 1.4 1.5 1.4 1.5 ...
 $ Petal.Width : num 0.2 0.2 0.2 0.2 0.2 0.4 0.3 0.2 0.2 0.1 ...
 $ Species      : Factor w/ 3 levels "setosa", "versicolor", ...: 1 1 1 1 1 1 1 1 1 1 ...

class(iris$Species)
[1] "factor"
```

We see that `iris` has the class `data.frame` and using `str()` allows us to examine the data inside. The variable `Species` in the `iris` data frame is of class `factor`, in contrast to the other variables which are of class `numeric`. The `str()` function also provides the length of the variables and shows the first couple of observations, while the `class()` function only provides the object's class.

Section 8.2: Vectors and lists

Data in R are stored in vectors. A typical vector is a sequence of values all having the same storage mode (e.g., characters vectors, numeric vectors). See `?atomic` for details on the atomic implicit classes and their corresponding storage modes: `"logical"`, `"integer"`, `"numeric"` (synonym `"double"`), `"complex"`, `"character"` and `"raw"`. Many classes are simply an atomic vector with a `class` attribute on top:

```
x <- 1826
class(x) <- "Date"
x
# [1] "1975-01-01"
x <- as.Date("1970-01-01")
class(x)
#[1] "Date"
is(x, "Date")
#[1] TRUE
is(x, "integer")
#[1] FALSE
is(x, "numeric")
#[1] FALSE
mode(x)
#[1] "numeric"
```

Lists are a special type of vector where each element can be anything, even another list, hence the R term for lists: "recursive vectors":

```
mylist <- list( A = c(5, 6, 7, 8), B = letters[1:10], CC = list( 5, "Z" ) )
```

Lists have two very important uses:

- Since functions can only return a single value, it is common to return complicated results in a list:

```
f <- function(x) list(xplus = x + 10, xsq = x^2)

f(7)
# $xplus
# [1] 17
#
# $xsq
# [1] 49
```

- Lists are also the underlying fundamental class for data frames. Under the hood, a data frame is a list of vectors all having the same length:

```
L <- list(x = 1:2, y = c("A", "B"))
DF <- data.frame(L)
DF
#   x y
# 1 1 A
# 2 2 B
is.list(DF)
# [1] TRUE
```

The other class of recursive vectors is R expressions, which are "language"- objects

Section 8.3: Vectors

The most simple data structure available in R is a vector. You can make vectors of numeric values, logical values, and character strings using the `c()` function. For example:

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

You can also join to vectors using the `c()` function.

```
x <- c(1, 2, 5)
y <- c(3, 4, 6)
z <- c(x, y)
z
## [1] 1 2 5 3 4 6
```

A more elaborate treatment of how to create vectors can be found in the "*Creating vectors*" topic

Chapter 9: Lists

Section 9.1: Introduction to lists

Lists allow users to store multiple elements (like vectors and matrices) under a single object. You can use the [list](#) function to create a list:

```
l1 <- list(c(1, 2, 3), c("a", "b", "c"))
l1
## [[1]]
## [1] 1 2 3
##
## [[2]]
## [1] "a" "b" "c"
```

Notice the vectors that make up the above list are different classes. Lists allow users to group elements of different classes. Each element in a list can also have a name. List names are accessed by the [names](#) function, and are assigned in the same manner row and column names are assigned in a matrix.

```
names(l1)
## NULL
names(l1) <- c("vector1", "vector2")
l1
## $vector1
## [1] 1 2 3
##
## $vector2
## [1] "a" "b" "c"
```

It is often easier and safer to declare the list names when creating the list object.

```
l2 <- list(vec = c(1, 3, 5, 7, 9),
           mat = matrix(data = c(1, 2, 3), nrow = 3))
l2
## $vec
## [1] 1 3 5 7 9
##
## $mat
##      [,1]
## [1,]    1
## [2,]    2
## [3,]    3
names(l2)
## [1] "vec" "mat"
```

Above the list has two elements, named "vec" and "mat," a vector and matrix, respectively.

Section 9.2: Quick Introduction to Lists

In general, most of the objects you would interact with as a user would tend to be a vector; e.g numeric vector, logical vector. These objects can only take in a single type of variable (a numeric vector can only have numbers inside it).

A list would be able to store any type variable in it, making it to the generic object that can store any type of variables we would need.

Example of initializing a list

```
exampleList1 <- list('a', 'b')
exampleList2 <- list(1, 2)
exampleList3 <- list('a', 1, 2)
```

In order to understand the data that was defined in the list, we can use the str function.

```
str(exampleList1)
str(exampleList2)
str(exampleList3)
```

Subsetting of lists distinguishes between extracting a slice of the list, i.e. obtaining a list containing a subset of the elements in the original list, and extracting a single element. Using the [operator commonly used for vectors produces a new list.

```
# Returns List
exampleList3[1]
exampleList3[1:2]
```

To obtain a single element use [[instead.

```
# Returns Character
exampleList3[[1]]
```

List entries may be named:

```
exampleList4 <- list(
  num = 1:3,
  numeric = 0.5,
  char = c('a', 'b')
)
```

The entries in named lists can be accessed by their name instead of their index.

```
exampleList4[['char']]
```

Alternatively the \$ operator can be used to access named elements.

```
exampleList4$num
```

This has the advantage that it is faster to type and may be easier to read but it is important to be aware of a potential pitfall. The \$ operator uses partial matching to identify matching list elements and may produce unexpected results.

```
exampleList5 <- exampleList4[2:3]

exampleList4$num
# c(1, 2, 3)

exampleList5$num
# 0.5

exampleList5[['num']]
# NULL
```

Lists can be particularly useful because they can store objects of different lengths and of various classes.

```
## Numeric vector
exampleVector1 <- c(12, 13, 14)
## Character vector
exampleVector2 <- c("a", "b", "c", "d", "e", "f")
## Matrix
exampleMatrix1 <- matrix(rnorm(4), ncol = 2, nrow = 2)
## List
exampleList3 <- list('a', 1, 2)

exampleList6 <- list(
  num = exampleVector1,
  char = exampleVector2,
  mat = exampleMatrix1,
  list = exampleList3
)
exampleList6
#$num
#[1] 12 13 14
#
#$char
#[1] "a" "b" "c" "d" "e" "f"
#
#$mat
# [,1]      [,2]
#[1,] 0.5013050 -1.88801542
#[2,] 0.4295266  0.09751379
#
#$list
#$list[[1]]
#[1] "a"
#
#$list[[2]]
#[1] 1
#
#$list[[3]]
#[1] 2
```

Section 9.3: Serialization: using lists to pass information

There exist cases in which it is necessary to put data of different types together. In Azure ML for example, it is necessary to pass information from a R script module to another one exclusively through dataframes. Suppose we have a dataframe and a number:

```
> df
    name height      team fun_index title age      desc Y
1  Andrea   195     Lazio      97    6 33  eccecente 1
2    Paja   165 Fiorentina      87    6 31      deciso 1
3    Roro   190     Lazio      65    6 28      strano 0
4  Gioele    70     Lazio     100    0  2  simpatico 1
5   Cacio   170 Juventus      81    3 33       duro 0
6   Edola   171     Lazio      72    5 32 svampito 1
7  Salami   175     Inter      75    3 30 doppipasso 1
8  Braugo   180     Inter      79    5 32      gjn 0
9   Bennu   158 Juventus      80    6 28 esaurito 0
10 Riggio   182     Lazio      92    5 31 certezza 1
11 Giordano   185     Roma      79    5 29      buono 1
```

```
> number <- "42"
```

We can access to this information:

```
> paste(df$name[4], "is a", df3$team[4], "supporter." )
[1] "Gioele is a Lazio supporter."
> paste("The answer to THE question is", number )
[1] "The answer to THE question is 42"
```

In order to put different types of data in a dataframe we have to use the list object and the serialization. In particular we have to put the data in a generic list and then put the list in a particular dataframe:

```
l <- list(df, number)
dataframe_container <- data.frame(out2 = as.integer(serializerserialize(l, connection=NULL)))
```

Once we have stored the information in the dataframe, we need to deserialize it in order to use it:

```
#----- unserialize -----+
unser_obj <- unserialize(as.raw(dataframe_container$out2))
#----- taking back the elements-----+
df_mod      <- unser_obj[1][[1]]
number_mod   <- unser_obj[2][[1]]
```

Then, we can verify that the data are transferred correctly:

```
> paste(df_mod$name[4], "is a", df_mod$team[4], "supporter." )
[1] "Gioele is a Lazio supporter."
> paste("The answer to THE question is", number_mod )
[1] "The answer to THE question is 42"
```

Chapter 10: Hashmaps

Section 10.1: Environments as hash maps

Note: in the subsequent passages, the terms **hash map** and **hash table** are used interchangeably and refer to the [same concept](#), namely, a data structure providing efficient key lookup through use of an internal hash function.

Introduction

Although R does not provide a native hash table structure, similar functionality can be achieved by leveraging the fact that the **environment** object returned from **new.env** (by default) provides hashed key lookups. The following two statements are equivalent, as the hash parameter defaults to TRUE:

```
H <- new.env(hash = TRUE)
H <- new.env()
```

Additionally, one may specify that the internal hash table is pre-allocated with a particular size via the **size** parameter, which has a default value of 29. Like all other R objects, **environments** manage their own memory and will grow in capacity as needed, so while it is not necessary to request a non-default value for **size**, there may be a slight performance advantage in doing so **if** the object will (eventually) contain a very large number of elements. It is worth noting that allocating extra space via **size** does not, in itself, result in an object with a larger memory footprint:

```
object.size(new.env())
# 56 bytes

object.size(new.env(size = 10e4))
# 56 bytes
```

Insertion

Insertion of elements may be done using either of the `[[<-` or `$<-` methods provided for the **environment** class, *but not* by using "single bracket" assignment (`[<-`):

```
H <- new.env()

H[["key"]] <- rnorm(1)

key2 <- "xyz"
H[[key2]] <- data.frame(x = 1:3, y = letters[1:3])

H$another_key <- matrix(rbinom(9, 1, 0.5) > 0, nrow = 3)

H["error"] <- 42
#Error in H["error"] <- 42 :
# object of type 'environment' is not subsettable
```

Like other facets of R, the first method (`object[[key]] <- value`) is generally preferred to the second (`object$key <- value`) because in the former case, a variable maybe be used instead of a literal value (e.g `key2` in the example above).

As is generally the case with hash map implementations, the **environment** object will **not** store duplicate keys. Attempting to insert a key-value pair for an existing key will replace the previously stored value:

```
H[["key3"]] <- "original value"
H[["key3"]] <- "new value"
H[["key3"]]
#[1] "new value"
```

Key Lookup

Likewise, elements may be accessed with [[or \$, but not with [:

```
H[["key"]]
#[1] 1.630631

H[[key2]] ## assuming key2 <- "xyz"
#   x y
# 1 1 a
# 2 2 b
# 3 3 c

H$another_key
# [,1] [,2] [,3]
# [1,] TRUE TRUE TRUE
# [2,] FALSE FALSE FALSE
# [3,] TRUE TRUE TRUE

H[1]
#Error in H[1] : object of type 'environment' is not subsettable
```

Inspecting the Hash Map

Being just an ordinary **environment**, the hash map can be inspected by typical means:

```
names(H)
#[1] "another_key" "xyz"       "key"       "key3"

ls(H)
#[1] "another_key" "key"       "key3"     "xyz"

str(H)
#<environment: 0x7828228>

ls.str(H)
# another_key : logi [1:3, 1:3] TRUE FALSE TRUE TRUE FALSE TRUE ...
# key : num 1.63
# key3 : chr "new value"
# xyz : 'data.frame':   3 obs. of  2 variables:
#   $ x: int 1 2 3
#   $ y: chr "a" "b" "c"
```

Elements can be removed using rm:

```
rm(list = c("key", "key3"), envir = H)

ls.str(H)
# another_key : logi [1:3, 1:3] TRUE FALSE TRUE TRUE FALSE TRUE ...
# xyz : 'data.frame':   3 obs. of  2 variables:
#   $ x: int 1 2 3
```

```
# $ y: chr "a" "b" "c"
```

Flexibility

One of the major benefits of using **environment** objects as hash tables is their ability to store virtually any type of object as a value, *even other environments*:

```
H2 <- new.env()

H2[["a"]] <- LETTERS
H2[["b"]] <- as.list(x = 1:5, y = matrix(rnorm(10), 2))
H2[["c"]] <- head(mtcars, 3)
H2[["d"]] <- Sys.Date()
H2[["e"]] <- Sys.time()
H2[["f"]] <- (function() {
  H3 <- new.env()
  for (i in seq_along(names(H2))) {
    H3[[names(H2)[i]]] <- H2[[names(H2)[i]]]
  }
  H3
})()

ls.str(H2)
# a : chr [1:26] "A" "B" "C" "D" "E" "F" "G" "H" "I" "J" "K" ...
# b : List of 5
# $ : int 1
# $ : int 2
# $ : int 3
# $ : int 4
# $ : int 5
# c : 'data.frame':   3 obs. of  11 variables:
# $ mpg : num  21 21 22.8
# $ cyl : num  6 6 4
# $ disp: num  160 160 108
# $ hp  : num  110 110 93
# $ drat: num  3.9 3.9 3.85
# $ wt   : num  2.62 2.88 2.32
# $ qsec: num  16.5 17 18.6
# $ vs   : num  0 0 1
# $ am   : num  1 1 1
# $ gear: num  4 4 4
# $ carb: num  4 4 1
# d : Date[1:1], format: "2016-08-03"
# e : POSIXct[1:1], format: "2016-08-03 19:25:14"
# f : <environment: 0x91a7cb8>

ls.str(H2$f)
# a : chr [1:26] "A" "B" "C" "D" "E" "F" "G" "H" "I" "J" "K" ...
# b : List of 5
# $ : int 1
# $ : int 2
# $ : int 3
# $ : int 4
# $ : int 5
# c : 'data.frame':   3 obs. of  11 variables:
# $ mpg : num  21 21 22.8
# $ cyl : num  6 6 4
# $ disp: num  160 160 108
# $ hp  : num  110 110 93
# $ drat: num  3.9 3.9 3.85
# $ wt   : num  2.62 2.88 2.32
```

```
# $ qsec: num 16.5 17 18.6
# $ vs : num 0 0 1
# $ am : num 1 1 1
# $ gear: num 4 4 4
# $ carb: num 4 4 1
# d : Date[1:1], format: "2016-08-03"
# e : POSIXct[1:1], format: "2016-08-03 19:25:14"
```

Limitations

One of the major limitations of using `environment` objects as hash maps is that, unlike many aspects of R, vectorization is not supported for element lookup / insertion:

```
names(H2)
#[1] "a" "b" "c" "d" "e" "f"

H2[[c("a", "b")]]
#Error in H2[[c("a", "b")]] :
# wrong arguments for subsetting an environment

Keys <- c("a", "b")
H2[[Keys]]
#Error in H2[[Keys]] : wrong arguments for subsetting an environment
```

Depending on the nature of the data being stored in the object, it may be possible to use `vapply` or `list2env` for assigning many elements at once:

```
E1 <- new.env()
invisible({
  vapply(letters, function(x) {
    E1[[x]] <- rnorm(1)
    logical(0)
  }, FUN.VALUE = logical(0))
})

all.equal(sort(names(E1)), letters)
#[1] TRUE

Keys <- letters
E2 <- list2env(
  setNames(
    as.list(rnorm(26)),
    nm = Keys),
  envir = NULL,
  hash = TRUE
)

all.equal(sort(names(E2)), letters)
#[1] TRUE
```

Neither of the above are particularly concise, but may be preferable to using a `for` loop, etc. when the number of key-value pairs is large.

Section 10.2: package:hash

The [hash package](#) offers a hash structure in R. However, it [terms of timing](#) for both inserts and reads it compares unfavorably to using environments as a hash. This documentation simply acknowledges its existence and provides sample timing code below for the above stated reasons. There is no identified case where hash is an appropriate

solution in R code today.

Consider:

```
# Generic unique string generator
unique_strings <- function(n){
  string_i <- 1
  string_len <- 1
  ans <- character(n)
  chars <- c(letters, LETTERS)
  new_strings <- function(len,pfx){
    for(i in 1:length(chars)){
      if (len == 1){
        ans[string_i] <- paste(pfx,chars[i],sep=' ')
        string_i <- string_i + 1
      } else {
        new_strings(len-1,pfx=paste(pfx,chars[i],sep=' '))
      }
      if (string_i > n) return ()
    }
  }
  while(string_i <= n){
    new_strings(string_len,'')
    string_len <- string_len + 1
  }
  sample(ans)
}

# Generate timings using an environment
timingsEnv <- plyr::adply(2^(10:15), .mar=1, .fun=function(i){
  strings <- unique_strings(i)
  ht1 <- new.env(hash=TRUE)
  lapply(strings, function(s){ ht1[[s]] <- 0L})
  data.frame(
    size=c(i,i),
    seconds=c(
      system.time(for (j in 1:i) ht1[[strings[j]]]==0L)[3]),
    type = c('1_hashedEnv')
  )
})

timingsHash <- plyr::adply(2^(10:15), .mar=1, .fun=function(i){
  strings <- unique_strings(i)
  ht <- hash::hash()
  lapply(strings, function(s) ht[[s]] <- 0L)
  data.frame(
    size=c(i,i),
    seconds=c(
      system.time(for (j in 1:i) ht[[strings[j]]]==0L)[3]),
    type = c('3_stringHash')
  )
})
```

Section 10.3: package:listenv

Although `package:listenv` implements a list-like interface to environments, its performance relative to environments for hash-like purposes is [poor on hash retrieval](#). However, if the indexes are numeric, it can be quite fast on retrieval. However, they have other advantages, e.g. compatibility with `package:future`. Covering this package for that purpose goes beyond the scope of the current topic. However, the timing code provided here can be used in conjunction with the example for `package:hash` for write timings.

```
timingsListEnv <- plyr::adply(2^(10:15), .mar=1, .fun=function(i){  
  strings <- unique_strings(i)  
  le <- listenv::listenv()  
  lapply(strings, function(s) le[[s]] <- 0L)  
  data.frame(  
    size=c(i,i),  
    seconds=c(  
      system.time(for (k in 1:i) le[[k]]==0L)[3]),  
    type = c('2_numericListEnv')  
  )  
})
```

Chapter 11: Creating vectors

Section 11.1: Vectors from build in constants: Sequences of letters & month names

R has a number of build in constants. The following constants are available:

- **LETTERS**: the 26 upper-case letters of the Roman alphabet
- **letters**: the 26 lower-case letters of the Roman alphabet
- **month.abb**: the three-letter abbreviations for the English month names
- **month.name**: the English names for the months of the year
- **pi**: the ratio of the circumference of a circle to its diameter

From the letters and month constants, vectors can be created.

1) Sequences of letters:

```
> letters
[1] "a" "b" "c" "d" "e" "f" "g" "h" "i" "j" "k" "l" "m" "n" "o" "p" "q" "r" "s" "t" "u" "v" "w" "x"
"y" "z"

> LETTERS[7:9]
[1] "G" "H" "I"

> letters[c(1,5,3,2,4)]
[1] "a" "e" "c" "b" "d"
```

2) Sequences of month abbreviations or month names:

```
> month.abb
[1] "Jan" "Feb" "Mar" "Apr" "May" "Jun" "Jul" "Aug" "Sep" "Oct" "Nov" "Dec"

> month.name[1:4]
[1] "January" "February" "March" "April"

> month.abb[c(3,6,9,12)]
[1] "Mar" "Jun" "Sep" "Dec"
```

Section 11.2: Creating named vectors

Named vector can be created in several ways. With c:

```
xc <- c('a' = 5, 'b' = 6, 'c' = 7, 'd' = 8)
```

which results in:

```
> xc
a b c d
5 6 7 8
```

with **list**:

```
x1 <- list('a' = 5, 'b' = 6, 'c' = 7, 'd' = 8)
```

which results in:

```
> x1  
$a  
[1] 5
```

```
$b  
[1] 6
```

```
$c  
[1] 7
```

```
$d  
[1] 8
```

With the **setNames** function, two vectors of the same length can be used to create a named vector:

```
x <- 5:8  
y <- letters[1:4]  
  
xy <- setNames(x, y)
```

which results in a named integer vector:

```
> xy  
a b c d  
5 6 7 8
```

As can be seen, this gives the same result as the **c** method.

You may also use the **names** function to get the same result:

```
xy <- 5:8  
names(xy) <- letters[1:4]
```

With such a vector it is also possibly to select elements by name:

```
> xy[ "c" ]  
c  
7
```

This feature makes it possible to use such a named vector as a look-up vector/table to match the values to values of another vector or column in dataframe. Considering the following dataframe:

```
mydf <- data.frame(let = c('c', 'a', 'b', 'd'))  
  
> mydf  
let  
1 c  
2 a  
3 b  
4 d
```

Suppose you want to create a new variable in the **mydf** dataframe called **num** with the correct values from **xy** in the rows. Using the **match** function the appropriate values from **xy** can be selected:

```
mydf$num <- xy[match(mydf$let, names(xy))]
```

which results in:

```
> mydf
let num
1  c  7
2  a  5
3  b  6
4  d  8
```

Section 11.3: Sequence of numbers

Use the `:` operator to create sequences of numbers, such as for use in vectorizing larger chunks of your code:

```
x <- 1:5
x
## [1] 1 2 3 4 5
```

This works both ways

```
10:4
# [1] 10 9 8 7 6 5 4
```

and even with floating point numbers

```
1.25:5
# [1] 1.25 2.25 3.25 4.25
```

or negatives

```
-4:4
#[1] -4 -3 -2 -1  0  1  2  3  4
```

Section 11.4: seq()

`seq` is a more flexible function than the `:` operator allowing to specify steps other than 1.

The function creates a sequence from the `start` (default is 1) to the end including that number.

You can supply only the end (`to`) parameter

```
seq(5)
# [1] 1 2 3 4 5
```

As well as the start

```
seq(2, 5) # or seq(from=2, to=5)
# [1] 2 3 4 5
```

And finally the step (`by`)

```
seq(2, 5, 0.5) # or seq(from=2, to=5, by=0.5)
# [1] 2.0 2.5 3.0 3.5 4.0 4.5 5.0
```

`seq` can optionally infer the (evenly spaced) steps when alternatively the desired length of the output (`length.out`) is supplied

```
seq(2, 5, length.out = 10)
```

```
# [1] 2.0 2.3 2.6 2.9 3.2 3.5 3.8 4.1 4.4 4.7 5.0
```

If the sequence needs to have the same length as another vector we can use the `along.with` as a shorthand for `length.out = length(x)`

```
x = 1:8
seq(2,5,along.with = x)
# [1] 2.000000 2.428571 2.857143 3.285714 3.714286 4.142857 4.571429 5.000000
```

There are two useful simplified functions in the `seq` family: `seq_along`, `seq_len`, and `seq.int`. `seq_along` and `seq_len` functions construct the natural (counting) numbers from 1 through N where N is determined by the function argument, the length of a vector or list with `seq_along`, and the integer argument with `seq_len`.

```
seq_along(x)
# [1] 1 2 3 4 5 6 7 8
```

Note that `seq_along` returns the indices of an existing object.

```
# counting numbers 1 through 10
seq_len(10)
[1] 1 2 3 4 5 6 7 8 9 10
# indices of existing vector (or list) with seq_along
letters[1:10]
[1] "a" "b" "c" "d" "e" "f" "g" "h" "i" "j"
seq_along(letters[1:10])
[1] 1 2 3 4 5 6 7 8 9 10
```

`seq.int` is the same as `seq` maintained for ancient compatibility.

There is also an old function `sequence` that creates a vector of sequences from a non negative argument.

```
sequence(4)
# [1] 1 2 3 4
sequence(c(3, 2))
# [1] 1 2 3 1 2
sequence(c(3, 2, 5))
# [1] 1 2 3 1 2 1 2 3 4 5
```

Section 11.5: Vectors

Vectors in R can have different types (e.g. integer, logical, character). The most general way of defining a vector is by using the function `vector()`.

```
vector('integer',2) # creates a vector of integers of size 2.
vector('character',2) # creates a vector of characters of size 2.
vector('logical',2) # creates a vector of logicals of size 2.
```

However, in R, the shorthand functions are generally more popular.

```
integer(2) # is the same as vector('integer',2) and creates an integer vector with two elements
character(2) # is the same as vector('integer',2) and creates an character vector with two elements
logical(2) # is the same as vector('logical',2) and creates an logical vector with two elements
```

Creating vectors with values, other than the default values, is also possible. Often the function `c()` is used for this. The `c` is short for combine or concatenate.

```
c(1, 2) # creates a integer vector of two elements: 1 and 2.
c('a', 'b') # creates a character vector of two elements: a and b.
c(T, F) # creates a logical vector of two elements: TRUE and FALSE.
```

Important to note here is that R interprets any integer (e.g. 1) as an integer vector of size one. The same holds for numerics (e.g. 1.1), logicals (e.g. T or F), or characters (e.g. 'a'). Therefore, you are in essence combining vectors, which in turn are vectors.

Pay attention that you always have to combine similar vectors. Otherwise, R will try to convert the vectors in vectors of the same type.

```
c(1,1.1,'a',T) # all types (integer, numeric, character and logical) are converted to the 'lowest' type which is character.
```

Finding elements in vectors can be done with the [operator.

```
vec_int <- c(1,2,3)
vec_char <- c('a','b','c')
vec_int[2] # accessing the second element will return 2
vec_char[2] # accessing the second element will return 'b'
```

This can also be used to change values

```
vec_int[2] <- 5 # change the second value from 2 to 5
vec_int # returns [1] 1 5 3
```

Finally, the : operator (short for the function `seq()`) can be used to quickly create a vector of numbers.

```
vec_int <- 1:10
vec_int # returns [1] 1 2 3 4 5 6 7 8 9 10
```

This can also be used to subset vectors (from easy to more complex subsets)

```
vec_char <- c('a','b','c','d','e')
vec_char[2:4] # returns [1] "b" "c" "d"
vec_char[c(1,3,5)] # returns [1] "a" "c" "e"
```

Section 11.6: Expanding a vector with the `rep()` function

The `rep` function can be used to repeat a vector in a fairly flexible manner.

```
# repeat counting numbers, 1 through 5 twice
rep(1:5, 2)
[1] 1 2 3 4 5 1 2 3 4 5

# repeat vector with incomplete recycling
rep(1:5, 2, length.out=7)
[1] 1 2 3 4 5 1 2
```

The `each` argument is especially useful for expanding a vector of statistics of observational/experimental units into a vector of data.frame with repeated observations of these units.

```
# same except repeat each integer next to each other
rep(1:5, each=2)
[1] 1 1 2 2 3 3 4 4 5 5
```

A nice feature of `rep` regarding involving expansion to such a data structure is that expansion of a vector to an unbalanced panel can be accomplished by replacing the length argument with a vector that dictates the number of times to repeat each element in the vector:

```
# automated length repetition
rep(1:5, 1:5)
[1] 1 2 2 3 3 3 4 4 4 4 5 5 5 5 5
# hand-fed repetition length vector
rep(1:5, c(1,1,1,2,2))
[1] 1 2 3 4 4 5 5
```

This should expose the possibility of allowing an external function to feed the second argument of `rep` in order to dynamically construct a vector that expands according to the data.

As with `seq`, faster, simplified versions of `rep` are `rep_len` and `rep.int`. These drop some attributes that `rep` maintains and so may be most useful in situations where speed is a concern and additional aspects of the repeated vector are unnecessary.

```
# repeat counting numbers, 1 through 5 twice
rep.int(1:5, 2)
[1] 1 2 3 4 5 1 2 3 4 5

# repeat vector with incomplete recycling
rep_len(1:5, length.out=7)
[1] 1 2 3 4 5 1 2
```

Chapter 12: Date and Time

R comes with classes for dates, date-times and time differences; see [?Dates](#), [?DateTimeClasses](#), [?difftime](#) and follow the "See Also" section of those docs for further documentation. Related Docs: Dates and Date-Time Classes.

Section 12.1: Current Date and Time

R is able to access the current date, time and time zone:

```
Sys.Date()           # Returns date as a Date object  
## [1] "2016-07-21"  
  
Sys.time()          # Returns date & time at current locale as a POSIXct object  
## [1] "2016-07-21 10:04:39 CDT"  
  
as.numeric(Sys.time()) # Seconds from UNIX Epoch (1970-01-01 00:00:00 UTC)  
## [1] 1469113479  
  
Sys.timezone()       # Time zone at current location  
## [1] "Australia/Melbourne"
```

Use `OlsonNames()` to view the time zone names in Olson/IANA database on the current system:

```
str(OlsonNames())  
## chr [1:589] "Africa/Abidjan" "Africa/Accra" "Africa/Addis_Ababa" "Africa/Algiers"  
"Africa/Asmara" "Africa/Asmera" "Africa/Bamako" ...
```

Section 12.2: Go to the End of the Month

Let's say we want to go to the last day of the month, this function will help on it:

```
eom <- function(x, p=as.POSIXlt(x)) as.Date(modifyList(p, list(mon=p$mon + 1, mday=0)))
```

Test:

```
x <- seq(as.POSIXct("2000-12-10"), as.POSIXct("2001-05-10"), by="months")  
> data.frame(before=x, after=eom(x))  
  before      after  
1 2000-12-10 2000-12-31  
2 2001-01-10 2001-01-31  
3 2001-02-10 2001-02-28  
4 2001-03-10 2001-03-31  
5 2001-04-10 2001-04-30  
6 2001-05-10 2001-05-31  
>
```

Using a date in a string format:

```
> eom('2000-01-01')  
[1] "2000-01-31"
```

Section 12.3: Go to First Day of the Month

Let's say we want to go to the first day of a given month:

```
date <- as.Date("2017-01-20")
> as.POSIXlt(cut(date, "month"))
[1] "2017-01-01 EST"
```

Section 12.4: Move a date a number of months consistently by months

Let's say we want to move a given date a numof months. We can define the following function, that uses the mondate package:

```
moveNumOfMonths <- function(date, num) {
  as.Date(mondate(date) + num)
}
```

It moves consistently the month part of the date and adjusting the day, in case the date refers to the last day of the month.

For example:

Back one month:

```
> moveNumOfMonths("2017-10-30", -1)
[1] "2017-09-30"
```

Back two months:

```
> moveNumOfMonths("2017-10-30", -2)
[1] "2017-08-30"
```

Forward two months:

```
> moveNumOfMonths("2017-02-28", 2)
[1] "2017-04-30"
```

It moves two months from the last day of February, therefore the last day of April.

Let's see how it works for backward and forward operations when it is the last day of the month:

```
> moveNumOfMonths("2016-11-30", 2)
[1] "2017-01-31"
> moveNumOfMonths("2017-01-31", -2)
[1] "2016-11-30"
```

Because November has 30 days, we get the same date in the backward operation, but:

```
> moveNumOfMonths("2017-01-30", -2)
[1] "2016-11-30"
> moveNumOfMonths("2016-11-30", 2)
[1] "2017-01-31"
```

Because January has 31 days, then moving two months from last day of November will get the last day of January.

Chapter 13: The Date class

Section 13.1: Formatting Dates

To format Dates we use the `format(date, format = "%Y-%m-%d")` function with either the `POSIXct` (given from `as.POSIXct()`) or `POSIXlt` (given from `as.POSIXlt()`)

```
d = as.Date("2016-07-21") # Current Date Time Stamp

format(d, "%a")           # Abbreviated Weekday
## [1] "Thu"

format(d, "%A")           # Full Weekday
## [1] "Thursday"

format(d, "%b")           # Abbreviated Month
## [1] "Jul"

format(d, "%B")           # Full Month
## [1] "July"

format(d, "%m")           # 00-12 Month Format
## [1] "07"

format(d, "%d")           # 00-31 Day Format
## [1] "21"

format(d, "%e")           # 0-31 Day Format
## [1] "21"

format(d, "%y")           # 00-99 Year
## [1] "16"

format(d, "%Y")           # Year with Century
## [1] "2016"
```

For more, see `?strptime`.

Section 13.2: Parsing Strings into Date Objects

R contains a Date class, which is created with `as.Date()`, which takes a string or vector of strings, and if the date is not in ISO 8601 date format YYYY-MM-DD, a formatting string of `strptime`-style tokens.

```
as.Date('2016-08-01')      # in ISO format, so does not require formatting string
## [1] "2016-08-01"

as.Date('05/23/16', format = '%m/%d/%y')
## [1] "2016-05-23"

as.Date('March 23rd, 2016', '%B %d %Y')    # add separators and literals to format
## [1] "2016-03-23"

as.Date(' 2016-08-01 foo')    # leading whitespace and all trailing characters are ignored
## [1] "2016-08-01"

as.Date(c('2016-01-01', '2016-01-02'))
# [1] "2016-01-01" "2016-01-02"
```

Section 13.3: Dates

To coerce a variable to a date use the **as.Date()** function.

```
> x <- as.Date("2016-8-23")
> x
[1] "2016-08-23"
> class(x)
[1] "Date"
```

The **as.Date()** function allows you to provide a format argument. The default is `%Y-%m-%d`, which is Year-month-day.

```
> as.Date("23-8-2016", format = "%d-%m-%Y") # To read in an European-style date
[1] "2016-08-23"
```

The format string can be placed either within a pair of single quotes or double quotes. Dates are usually expressed in a variety of forms such as: `"d-m-yy"` or `"d-m-YYYY"` or `"m-d-yy"` or `"m-d-YYYY"` or `"YYYY-m-d"` or `"YYYY-d-m"`. These formats can also be expressed by replacing `"-"` by `"/"`. Further, dates are also expressed in the forms, say, `"Nov 6, 1986"` or `"November 6, 1986"` or `"6 Nov, 1986"` or `"6 November, 1986"` and so on. The **as.Date()** function accepts all such character strings and when we mention the appropriate format of the string, it always outputs the date in the form `"YYYY-m-d"`.

Suppose we have a date string `"9-6-1962"` in the format `"%d-%m-%Y"`.

```
# 
# It tries to interprets the string as YYYY-m-d
#
> as.Date("9-6-1962")
[1] "0009-06-19"      #interprets as "%Y-%m-%d"
>
as.Date("9/6/1962")
[1] "0009-06-19"      #again interprets as "%Y-%m-%d"
>
# It has no problem in understanding, if the date is in form YYYY-m-d or YYYY/m/d
#
> as.Date("1962-6-9")
[1] "1962-06-09"      # no problem
> as.Date("1962/6/9")
[1] "1962-06-09"      # no problem
>
```

By specifying the correct format of the input string, we can get the desired results. We use the following codes for specifying the formats to the **as.Date()** function.

Format Code	Meaning
<code>%d</code>	day
<code>%m</code>	month
<code>%y</code>	year in 2-digits
<code>%Y</code>	year in 4-digits
<code>%b</code>	abbreviated month in 3 chars
<code>%B</code>	full name of the month

Consider the following example specifying the **format** parameter:

```
> as.Date("9-6-1962", format="%d-%m-%Y")
[1] "1962-06-09"
>
```

The parameter name **format** can be omitted.

```
> as.Date("9-6-1962", "%d-%m-%Y")
[1] "1962-06-09"
>
```

Some times, names of the months abbreviated to the first three characters are used in the writing the dates. In which case we use the format specifier %b.

```
> as.Date("6Nov1962", "%d%b%Y")
[1] "1962-11-06"
>
```

Note that, there are no either '-' or '/' or white spaces between the members in the date string. The format string should exactly match that input string. Consider the following example:

```
> as.Date("6 Nov, 1962", "%d %b, %Y")
[1] "1962-11-06"
>
```

Note that, there is a comma in the date string and hence a comma in the format specification too. If comma is omitted in the format string, it results in an NA. An example usage of %B format specifier is as follows:

```
> as.Date("October 12, 2016", "%B %d, %Y")
[1] "2016-10-12"
>
> as.Date("12 October, 2016", "%d %B, %Y")
[1] "2016-10-12"
>
```

%y format is system specific and hence, should be used with caution. Other parameters used with this function are **origin** and **tz**(time zone).

Chapter 14: Date-time classes (POSIXct and POSIXlt)

R includes two date-time classes -- POSIXct and POSIXlt -- see `?DateTimeClasses`.

Section 14.1: Formatting and printing date-time objects

```
# test date-time object
options(digits.secs = 3)
d = as.POSIXct("2016-08-30 14:18:30.58", tz = "UTC")

format(d, "%S") # 00-61 Second as integer
## [1] "30"

format(d, "%OS") # 00-60.99... Second as fractional
## [1] "30.579"

format(d, "%M") # 00-59 Minute
## [1] "18"

format(d, "%H") # 00-23 Hours
## [1] "14"

format(d, "%I") # 01-12 Hours
## [1] "02"

format(d, "%p") # AM/PM Indicator
## [1] "PM"

format(d, "%z") # Signed offset
## [1] "+0000"

format(d, "%Z") # Time Zone Abbreviation
## [1] "UTC"
```

See `?strptime` for details on the format strings here, as well as other formats.

Section 14.2: Date-time arithmetic

To add/subtract time, use POSIXct, since it stores times in seconds

```
## adding/subtracting times - 60 seconds
as.POSIXct("2016-01-01") + 60
# [1] "2016-01-01 00:01:00 AEDT"

## adding 3 hours, 14 minutes, 15 seconds
as.POSIXct("2016-01-01") + ( (3 * 60 * 60) + (14 * 60) + 15 )
# [1] "2016-01-01 03:14:15 AEDT"
```

More formally, `as.difftime` can be used to specify time periods to add to a date or datetime object. E.g.:

```
as.POSIXct("2016-01-01") +
  as.difftime(3, units="hours") +
  as.difftime(14, units="mins") +
  as.difftime(15, units="secs")
# [1] "2016-01-01 03:14:15 AEDT"
```

To find the difference between dates/times use `difftime()` for differences in seconds, minutes, hours, days or weeks.

```
# using POSIXct objects
difftime(
  as.POSIXct("2016-01-01 12:00:00"),
  as.POSIXct("2016-01-01 11:59:59"),
  unit = "secs")
# Time difference of 1 secs
```

To generate sequences of date-times use `seq.POSIXt()` or simply `seq`.

Section 14.3: Parsing strings into date-time objects

The functions for parsing a string into POSIXct and POSIXlt take similar parameters and return a similar-looking result, but there are differences in how that date-time is stored; see "Remarks."

```
as.POSIXct("11:38",
           format = "%H:%M")                      # time string
#> [1] "2016-07-21 11:38:00 CDT"          # formatting string

strptime("11:38",
         format = "%H:%M")                      # identical, but makes a POSIXlt object
#> [1] "2016-07-21 11:38:00 CDT"

as.POSIXct("11 AM",
           format = "%I %p")                     # identical
#> [1] "2016-07-21 11:00:00 CDT"
```

Note that date and timezone are imputed.

```
as.POSIXct("11:38:22",                      # time string without timezone
           format = "%H:%M:%S",
           tz = "America/New_York")      # set time zone
#> [1] "2016-07-21 11:38:22 EDT"

as.POSIXct("2016-07-21 00:00:00",
           format = "%F %T")                  # shortcut tokens for "%Y-%m-%d" and "%H:%M:%S"
```

See `?strptime` for details on the format strings here.

Notes

Missing elements

- If a date element is not supplied, then that from the current date is used.
- If a time element is not supplied, then that from midnight is used, i.e. 0s.
- If no timezone is supplied in either the string or the `tz` parameter, the local timezone is used.

Time zones

- The accepted values of `tz` depend on the location.
 - CST is given with `"CST6CDT"` or `"America/Chicago"`
- For supported locations and time zones use:
 - In R: `OlsonNames()`
 - Alternatively, try in R: `system("cat $R_HOME/share/zoneinfo/zone.tab")`
- These locations are given by [Internet Assigned Numbers Authority \(IANA\)](#)
 - [List of tz database time zones \(Wikipedia\)](#)

- [IANA TZ Data \(2016e\)](#)

Chapter 15: The character class

Characters are what other languages call 'string vectors.'

Section 15.1: Coercion

To check whether a value is a character use the `is.character()` function. To coerce a variable to a character use the `as.character()` function.

```
x <- "The quick brown fox jumps over the lazy dog"  
class(x)  
[1] "character"  
is.character(x)  
[1] TRUE
```

Note that numerics can be coerced to characters, but attempting to coerce a character to numeric may result in NA.

```
as.numeric("2")  
[1] 2  
as.numeric("fox")  
[1] NA  
Warning message:  
NAs introduced by coercion
```

Chapter 16: Numeric classes and storage modes

Section 16.1: Numeric

Numeric represents integers and doubles and is the default mode assigned to vectors of numbers. The function `is.numeric()` will evaluate whether a vector is numeric. It is important to note that although integers and doubles will pass `is.numeric()`, the function `as.numeric()` will always attempt to convert to type double.

```
x <- 12.3
y <- 12L

#confirm types
typeof(x)
[1] "double"
typeof(y)
[1] "integer"

# confirm both numeric
is.numeric(x)
[1] TRUE
is.numeric(y)
[1] TRUE

# logical to numeric
as.numeric(TRUE)
[1] 1

# While TRUE == 1, it is a double and not an integer
is.integer(as.numeric(TRUE))
[1] FALSE
```

Doubles are R's default numeric value. They are double precision vectors, meaning that they take up 8 bytes of memory for each value in the vector. R has no single precision data type and so all real numbers are stored in the double precision format.

```
is.double(1)
TRUE
is.double(1.0)
TRUE
is.double(1L)
FALSE
```

Integers are whole numbers that can be written without a fractional component. Integers are represented by a number with an L after it. Any number without an L after it will be considered a double.

```
typeof(1)
[1] "double"
class(1)
[1] "numeric"
typeof(1L)
[1] "integer"
class(1L)
[1] "integer"
```

Though in most cases using an integer or double will not matter, sometimes replacing doubles with integers will

consume less memory and operational time. A double vector uses 8 bytes per element while an integer vector uses only 4 bytes per element. As the size of vectors increases, using proper types can dramatically speed up processes.

```
# test speed on lots of arithmetic
microbenchmark(
  for( i in 1:100000){
    2L * i
    10L + i
  },
  for( i in 1:100000){
    2.0 * i
    10.0 + i
  }
)
Unit: milliseconds
                                         expr      min       lq     mean   median      uq
max neval
for (i in 1:1e+05) { 2L * i 10L + i } 40.74775 42.34747 50.70543 42.99120 65.46864
94.11804 100
for (i in 1:1e+05) { 2 * i 10 + i } 41.07807 42.38358 53.52588 44.26364 65.84971
83.00456 100
```

Chapter 17: The logical class

Logical is a mode (and an implicit class) for vectors.

Section 17.1: Logical operators

There are two sorts of logical operators: those that accept and return vectors of any length (elementwise operators: `!`, `&`, `&&`, `||`, `xor()`) and those that only evaluate the first element in each argument (`&&`, `||`). The second sort is primarily used as the cond argument to the `if` function.

Logical Operator	Meaning	Syntax
<code>!</code>	Not	<code>!x</code>
<code>&</code>	element-wise (vectorized) and	<code>x & y</code>
<code>&&</code>	and (single element only)	<code>x && y</code>
<code> </code>	element-wise (vectorized) or	<code>x y</code>
<code> </code>	or (single element only)	<code>x y</code>
<code>xor</code>	element-wise (vectorized) exclusive OR <code>xor(x,y)</code>	

Note that the `||` operator evaluates the left condition and if the left condition is TRUE the right side is never evaluated. This can save time if the first is the result of a complex operation. The `&&` operator will likewise return FALSE without evaluation of the second argument when the first element of the first argument is FALSE.

```
> x <- 5
> x > 6 || stop("X is too small")
Error: X is too small
> x > 3 || stop("X is too small")
[1] TRUE
```

To check whether a value is a logical you can use the `is.logical()` function.

Section 17.2: Coercion

To coerce a variable to a logical use the `as.logical()` function.

```
> x <- 2
> z <- x > 4
> z
[1] FALSE
> class(x)
[1] "numeric"
> as.logical(2)
[1] TRUE
```

When applying `as.numeric()` to a logical, a double will be returned. NA is a logical value and a logical operator with an NA will return NA if the outcome is ambiguous.

Section 17.3: Interpretation of NAs

See Missing values for details.

```
> TRUE & NA
[1] NA
> FALSE & NA
```

```
[1] FALSE
> TRUE || NA
[1] TRUE
> FALSE || NA
[1] NA
```

Chapter 18: Data frames

Section 18.1: Create an empty data.frame

A data.frame is a special kind of list: it is *rectangular*. Each element (column) of the list has same length, and where each row has a "row name". Each column has its own class, but the class of one column can be different from the class of another column (unlike a matrix, where all elements must have the same class).

In principle, a data.frame could have no rows and no columns:

```
> structure(list(character()), class = "data.frame")
NULL
<0 rows> (or 0-length row.names)
```

But this is unusual. It is more common for a data.frame to have many columns and many rows. Here is a data.frame with three rows and two columns (a is numeric class and b is character class):

```
> structure(list(a = 1:3, b = letters[1:3]), class = "data.frame")
[1] a b
<0 rows> (or 0-length row.names)
```

In order for the data.frame to print, we will need to supply some row names. Here we use just the numbers 1:3:

```
> structure(list(a = 1:3, b = letters[1:3]), class = "data.frame", row.names = 1:3)
   a b
1 1 a
2 2 b
3 3 c
```

Now it becomes obvious that we have a data.frame with 3 rows and 2 columns. You can check this using `nrow()`, `ncol()`, and `dim()`:

```
> x <- structure(list(a = numeric(3), b = character(3)), class = "data.frame", row.names = 1:3)
> nrow(x)
[1] 3
> ncol(x)
[1] 2
> dim(x)
[1] 3 2
```

R provides two other functions (besides `structure()`) that can be used to create a data.frame. The first is called, intuitively, `data.frame()`. It checks to make sure that the column names you supplied are valid, that the list elements are all the same length, and supplies some automatically generated row names. This means that the output of `data.frame()` might now always be exactly what you expect:

```
> str(data.frame("a a a" = numeric(3), "b-b-b" = character(3)))
'data.frame': 3 obs. of 2 variables:
 $ a.a.a: num 0 0 0
 $ b.b.b: Factor w/ 1 level "" : 1 1 1
```

The other function is called `as.data.frame()`. This can be used to coerce an object that is not a data.frame into being a data.frame by running it through `data.frame()`. As an example, consider a matrix:

```
> m <- matrix(letters[1:9], nrow = 3)
> m
```

```
[,1] [,2] [,3]
[1,] "a"   "d"   "g"
[2,] "b"   "e"   "h"
[3,] "c"   "f"   "i"
```

And the result:

```
> as.data.frame(m)
  V1 V2 V3
1  a  d  g
2  b  e  h
3  c  f  i
> str(as.data.frame(m))
'data.frame': 3 obs. of 3 variables:
 $ V1: Factor w/ 3 levels "a","b","c": 1 2 3
 $ V2: Factor w/ 3 levels "d","e","f": 1 2 3
 $ V3: Factor w/ 3 levels "g","h","i": 1 2 3
```

Section 18.2: Subsetting rows and columns from a data frame

Syntax for accessing rows and columns: [, [[, and \$

This topic covers the most common syntax to access specific rows and columns of a data frame. These are

- Like a **matrix** with single brackets `data[rows, columns]`
 - Using row and column numbers
 - Using column (and row) names
- Like a **list**:
 - With single brackets `data[columns]` to get a data frame
 - With double brackets `data[[one_column]]` to get a vector
- With \$ for a single column `data$column_name`

We will use the built-in `mtcars` data frame to illustrate.

Like a matrix: `data[rows, columns]`

With numeric indexes

Using the built in data frame `mtcars`, we can extract rows and columns using [] brackets with a comma included. Indices before the comma are rows:

```
# get the first row
mtcars[1, ]
# get the first five rows
mtcars[1:5, ]
```

Similarly, after the comma are columns:

```
# get the first column
mtcars[, 1]
# get the first, third and fifth columns:
mtcars[, c(1, 3, 5)]
```

As shown above, if either rows or columns are left blank, all will be selected. `mtcars[1,]` indicates the first row with *all* the columns.

With column (and row) names

So far, this is identical to how rows and columns of matrices are accessed. With `data.frames`, most of the time it is preferable to use a column name to a column index. This is done by using a `character` with the column name instead of `numeric` with a column number:

```
# get the mpg column  
mtcars[, "mpg"]  
# get the mpg, cyl, and disp columns  
mtcars[, c("mpg", "cyl", "disp")]
```

Though less common, row names can also be used:

```
mtcars["Mazda Rx4", ]
```

Rows and columns together

The row and column arguments can be used together:

```
# first four rows of the mpg column  
mtcars[1:4, "mpg"]  
  
# 2nd and 5th row of the mpg, cyl, and disp columns  
mtcars[c(2, 5), c("mpg", "cyl", "disp")]
```

A warning about dimensions:

When using these methods, if you extract multiple columns, you will get a data frame back. However, if you extract a *single* column, you will get a vector, not a data frame under the default options.

```
## multiple columns returns a data frame  
class(mtcars[, c("mpg", "cyl")])  
# [1] "data.frame"  
## single column returns a vector  
class(mtcars[, "mpg"])  
# [1] "numeric"
```

There are two ways around this. One is to treat the data frame as a list (see below), the other is to add a `drop = FALSE` argument. This tells R to not "drop the unused dimensions":

```
class(mtcars[, "mpg", drop = FALSE])  
# [1] "data.frame"
```

Note that matrices work the same way - by default a single column or row will be a vector, but if you specify `drop = FALSE` you can keep it as a one-column or one-row matrix.

Like a list

Data frames are essentially `lists`, i.e., they are a list of column vectors (that all must have the same length). Lists can be subset using single brackets [for a sub-list, or double brackets [] for a single element.

With single brackets `data[columns]`

When you use single brackets and no commas, you will get column back because data frames are lists of columns.

```
mtcars["mpg"]  
mtcars[c("mpg", "cyl", "disp")]  
my_columns <- c("mpg", "cyl", "hp")  
mtcars[my_columns]
```

Single brackets *like a list* vs. single brackets *like a matrix*

The difference between `data[columns]` and `data[, columns]` is that when treating the `data.frame` as a `list` (no comma in the brackets) the object returned will be a `data.frame`. If you use a comma to treat the `data.frame` like a `matrix` then selecting a single column will return a vector but selecting multiple columns will return a `data.frame`.

```
## When selecting a single column
## like a list will return a data frame
class(mtcars["mpg"])
# [1] "data.frame"
## like a matrix will return a vector
class(mtcars[, "mpg"])
# [1] "numeric"
```

With double brackets `data[[one_column]]`

To extract a single column *as a vector* when treating your `data.frame` as a `list`, you can use double brackets `[[]]`. This will only work for a single column at a time.

```
# extract a single column by name as a vector
mtcars[["mpg"]]

# extract a single column by name as a data frame (as above)
mtcars[ "mpg" ]
```

Using \$ to access columns

A single column can be extracted using the magical shortcut `$` without using a quoted column name:

```
# get the column "mpg"
mtcars$mpg
```

Columns accessed by `$` will always be vectors, not data frames.

Drawbacks of \$ for accessing columns

The `$` can be a convenient shortcut, especially if you are working in an environment (such as RStudio) that will auto-complete the column name in this case. **However**, `$` has drawbacks as well: it uses *non-standard evaluation* to avoid the need for quotes, which means it *will not work* if your column name is stored in a variable.

```
my_column <- "mpg"
# the below will not work
mtcars$my_column
# but these will work
mtcars[, my_column] # vector
mtcars[my_column]   # one-column data frame
mtcars[[my_column]] # vector
```

Due to these concerns, `$` is best used in *interactive* R sessions when your column names are constant. For *programmatic* use, for example in writing a generalizable function that will be used on different data sets with different column names, `$` should be avoided.

Also note that the default behaviour is to use partial matching only when extracting from recursive objects (except environments) by `$`

```
# give you the values of "mpg" column
# as "mtcars" has only one column having name starting with "m"
```

```
mtcars$m  
# will give you "NULL"  
# as "mtcars" has more than one columns having name starting with "d"  
mtcars$d
```

Advanced indexing: negative and logical indices

Whenever we have the option to use numbers for a index, we can also use negative numbers to omit certain indices or a boolean (logical) vector to indicate exactly which items to keep.

Negative indices omit elements

```
mtcars[1, ] # first row  
mtcars[-1, ] # everything but the first row  
mtcars[-(1:10), ] # everything except the first 10 rows
```

Logical vectors indicate specific elements to keep

We can use a condition such as < to generate a logical vector, and extract only the rows that meet the condition:

```
# logical vector indicating TRUE when a row has mpg less than 15  
# FALSE when a row has mpg >= 15  
test <- mtcars$mpg < 15  
  
# extract these rows from the data frame  
mtcars[test, ]
```

We can also bypass the step of saving the intermediate variable

```
# extract all columns for rows where the value of cyl is 4.  
mtcars[mtcars$cyl == 4, ]  
# extract the cyl, mpg, and hp columns where the value of cyl is 4  
mtcars[mtcars$cyl == 4, c("cyl", "mpg", "hp")]
```

Section 18.3: Convenience functions to manipulate data.frames

Some convenience functions to manipulate data.frames are **subset()**, **transform()**, **with()** and **within()**.

subset

The **subset()** function allows you to subset a **data.frame** in a more convenient way (subset also works with other classes):

```
subset(mtcars, subset = cyl == 6, select = c("mpg", "hp"))  
          mpg   hp  
Mazda RX4    21.0 110  
Mazda RX4 Wag 21.0 110  
Hornet 4 Drive 21.4 110  
Valiant      18.1 105  
Merc 280      19.2 123  
Merc 280C     17.8 123  
Ferrari Dino  19.7 175
```

In the code above we asking only for the lines in which cyl == 6 and for the columns mpg and hp. You could achieve the same result using [] with the following code:

```
mtcars[mtcars$cyl == 6, c("mpg", "hp")]
```

transform

The `transform()` function is a convenience function to change columns inside a `data.frame`. For instance the following code adds another column named `mpg2` with the result of `mpg^2` to the `mtcars data.frame`:

```
mtcars <- transform(mtcars, mpg2 = mpg^2)
```

with and within

Both `with()` and `within()` let you evaluate expressions inside the `data.frame` environment, allowing a somewhat cleaner syntax, saving you the use of some \$ or [].

For example, if you want to create, change and/or remove multiple columns in the `airquality data.frame`:

```
aq <- within(airquality, {
  10zone <- log(Ozone) # creates new column
  Month <- factor(month.abb[Month]) # changes Month Column
  cTemp <- round((Temp - 32) * 5/9, 1) # creates new column
  S.cT <- Solar.R / cTemp # creates new column
  rm(Day, Temp) # removes columns
})
```

Section 18.4: Introduction

Data frames are likely the data structure you will used most in your analyses. A data frame is a special kind of list that stores same-length vectors of different classes. You create data frames using the `data.frame` function. The example below shows this by combining a numeric and a character vector into a data frame. It uses the : operator, which will create a vector containing all integers from 1 to 3.

```
df1 <- data.frame(x = 1:3, y = c("a", "b", "c"))
df1
##   x y
## 1 1 a
## 2 2 b
## 3 3 c
class(df1)
## [1] "data.frame"
```

Data frame objects do not print with quotation marks, so the class of the columns is not always obvious.

```
df2 <- data.frame(x = c("1", "2", "3"), y = c("a", "b", "c"))
df2
##   x y
## 1 1 a
## 2 2 b
## 3 3 c
```

Without further investigation, the "x" columns in `df1` and `df2` cannot be differentiated. The `str` function can be used to describe objects with more detail than class.

```
str(df1)
## 'data.frame': 3 obs. of 2 variables:
## $ x: int 1 2 3
## $ y: Factor w/ 3 levels "a","b","c": 1 2 3
str(df2)
## 'data.frame': 3 obs. of 2 variables:
```

```
## $ x: Factor w/ 3 levels "1","2","3": 1 2 3
## $ y: Factor w/ 3 levels "a","b","c": 1 2 3
```

Here you see that df1 is a `data.frame` and has 3 observations of 2 variables, "x" and "y." Then you are told that "x" has the data type integer (not important for this class, but for our purposes it behaves like a numeric) and "y" is a factor with three levels (another data class we are not discussing). ***It is important to note that, by default, data frames coerce characters to factors.*** The default behavior can be changed with the `stringsAsFactors` parameter:

```
df3 <- data.frame(x = 1:3, y = c("a", "b", "c"), stringsAsFactors = FALSE)
str(df3)
## 'data.frame': 3 obs. of 2 variables:
## $ x: int 1 2 3
## $ y: chr "a" "b" "c"
```

Now the "y" column is a character. As mentioned above, each "column" of a data frame must have the same length. Trying to create a data.frame from vectors with different lengths will result in an error. (Try running `data.frame(x = 1:3, y = 1:4)` to see the resulting error.)

As test-cases for data frames, some data is provided by R by default. One of them is `iris`, loaded as follows:

```
mydataframe <- iris
str(mydataframe)
```

Section 18.5: Convert all columns of a data.frame to character class

A common task is to convert all columns of a `data.frame` to character class for ease of manipulation, such as in the cases of sending `data.frames` to a RDBMS or merging `data.frames` containing factors where levels may differ between input `data.frames`.

The best time to do this is when the data is read in - almost all input methods that create `data.frames` have an options `stringsAsFactors` which can be set to FALSE.

If the data has already been created, factor columns can be converted to character columns as shown below.

```
bob <- data.frame(jobs = c("scientist", "analyst"),
                   pay = c(160000, 100000), age = c(30, 25))
str(bob)
```

```
'data.frame': 2 obs. of 3 variables:
$ jobs: Factor w/ 2 levels "analyst","scientist": 2 1
$ pay : num 160000 100000
$ age : num 30 25
```

```
# Convert *all columns* to character
bob[] <- lapply(bob, as.character)
str(bob)
```

```
'data.frame': 2 obs. of 3 variables:
$ jobs: chr "scientist" "analyst"
$ pay : chr "160000" "1e+05"
$ age : chr "30" "25"
```

```
# Convert only factor columns to character
bob[] <- lapply(bob, function(x) {
```

```
if is.factor(x) x <- as.character(x)
return(x)
})
```

Chapter 19: Split function

Section 19.1: Using split in the split-apply-combine paradigm

A popular form of data analysis is [split-apply-combine](#), in which you split your data into groups, apply some sort of processing on each group, and then combine the results.

Let's consider a data analysis where we want to obtain the two cars with the best miles per gallon (mpg) for each cylinder count (cyl) in the built-in mtcars dataset. First, we split the `mtcars` data frame by the cylinder count:

```
(spl <- split(mtcars, mtcars$cyl))
# $`4`
#           mpg cyl disp hp drat    wt  qsec vs am gear carb
# Datsun 710 22.8   4 108.0 93 3.85 2.320 18.61  1  1    4    1
# Merc 240D  24.4   4 146.7 62 3.69 3.190 20.00  1  0    4    2
# Merc 230   22.8   4 140.8 95 3.92 3.150 22.90  1  0    4    2
# Fiat 128   32.4   4  78.7 66 4.08 2.200 19.47  1  1    4    1
# ...
#
# $`6`
#           mpg cyl disp hp drat    wt  qsec vs am gear carb
# Mazda RX4  21.0   6 160.0 110 3.90 2.620 16.46  0  1    4    4
# Mazda RX4 Wag 21.0   6 160.0 110 3.90 2.875 17.02  0  1    4    4
# Hornet 4 Drive 21.4   6 258.0 110 3.08 3.215 19.44  1  0    3    1
# Valiant    18.1   6 225.0 105 2.76 3.460 20.22  1  0    3    1
# ...
#
# $`8`
#           mpg cyl disp hp drat    wt  qsec vs am gear carb
# Hornet Sportabout 18.7   8 360.0 175 3.15 3.440 17.02  0  0    3    2
# Duster 360     14.3   8 360.0 245 3.21 3.570 15.84  0  0    3    4
# Merc 450SE     16.4   8 275.8 180 3.07 4.070 17.40  0  0    3    3
# Merc 450SL     17.3   8 275.8 180 3.07 3.730 17.60  0  0    3    3
# ...
```

This has returned a list of data frames, one for each cylinder count. As indicated by the output, we could obtain the relevant data frames with `spl$`4``, `spl$`6``, and `spl$`8`` (some might find it more visually appealing to use `spl$"4"` or `spl[["4"]]` instead).

Now, we can use `lapply` to loop through this list, applying our function that extracts the cars with the best 2 mpg values from each of the list elements:

```
(best2 <- lapply(spl, function(x) tail(x[order(x$mpg), ], 2)))
# $`4`
#           mpg cyl disp hp drat    wt  qsec vs am gear carb
# Fiat 128   32.4   4  78.7 66 4.08 2.200 19.47  1  1    4    1
# Toyota Corolla 33.9   4  71.1 65 4.22 1.835 19.90  1  1    4    1
#
# $`6`
#           mpg cyl disp hp drat    wt  qsec vs am gear carb
# Mazda RX4 Wag 21.0   6  160 110 3.90 2.875 17.02  0  1    4    4
# Hornet 4 Drive 21.4   6  258 110 3.08 3.215 19.44  1  0    3    1
#
# $`8`
#           mpg cyl disp hp drat    wt  qsec vs am gear carb
# Hornet Sportabout 18.7   8  360 175 3.15 3.440 17.02  0  0    3    2
# Pontiac Firebird 19.2   8  400 175 3.08 3.845 17.05  0  0    3    2
```

Finally, we can combine everything together using `rbind`. We want to call `rbind(best2[["4"]], best2[["6"]], best2[["8"]])`, but this would be tedious if we had a huge list. As a result, we use:

```
do.call(rbind, best2)
#          mpg cyl disp hp drat wt qsec vs am gear carb
# 4.Fiat 128    32.4   4 78.7 66 4.08 2.200 19.47 1 1 4 1
# 4.Toyota Corolla 33.9   4 71.1 65 4.22 1.835 19.90 1 1 4 1
# 6.Mazda RX4 Wag 21.0   6 160.0 110 3.90 2.875 17.02 0 1 4 4
# 6.Hornet 4 Drive 21.4   6 258.0 110 3.08 3.215 19.44 1 0 3 1
# 8.Hornet Sportabout 18.7   8 360.0 175 3.15 3.440 17.02 0 0 3 2
# 8.Pontiac Firebird 19.2   8 400.0 175 3.08 3.845 17.05 0 0 3 2
```

This returns the result of `rbind` (argument 1, a function) with all the elements of `best2` (argument 2, a list) passed as arguments.

With simple analyses like this one, it can be more compact (and possibly much less readable!) to do the whole split-apply-combine in a single line of code:

```
do.call(rbind, lapply(split(mtcars, mtcars$cyl), function(x) tail(x[order(x$mpg), ], 2)))
```

It is also worth noting that the `lapply(split(x, f), FUN)` combination can be alternatively framed using the `?by` function:

```
by(mtcars, mtcars$cyl, function(x) tail(x[order(x$mpg), ], 2))
do.call(rbind, by(mtcars, mtcars$cyl, function(x) tail(x[order(x$mpg), ], 2)))
```

Section 19.2: Basic usage of split

`split` allows to divide a vector or a data.frame into buckets with regards to a factor/group variables. This ventilation into buckets takes the form of a list, that can then be used to apply group-wise computation (`for` loops or `lapply/sapply`).

First example shows the usage of `split` on a vector:

Consider following vector of letters:

```
testdata <- c("e", "o", "r", "g", "a", "y", "w", "q", "i", "s", "b", "v", "x", "h", "u")
```

Objective is to separate those letters into vowels and consonants, ie split it accordingly to letter type.

Let's first create a grouping vector:

```
vowels <- c('a', 'e', 'i', 'o', 'u', 'y')
letter_type <- ifelse(testdata %in% vowels, "vowels", "consonants")
```

Note that `letter_type` has the same length that our vector `testdata`. Now we can `split` this test data in the two groups, vowels and consonants :

```
split(testdata, letter_type)
#$consonants
#[1] "r" "g" "w" "q" "s" "b" "v" "x" "h"

#$vowels
#[1] "e" "o" "a" "y" "i" "u"
```

Hence, the result is a list which names are coming from our grouping vector/factor `letter_type`.

`split` has also a method to deal with data.frames.

Consider for instance `iris` data:

```
data(iris)
```

By using `split`, one can create a list containing one data.frame per iris specie (variable: Species):

```
> liris <- split(iris, iris$Species)
> names(liris)
[1] "setosa"      "versicolor"   "virginica"
> head(liris$setosa)
  Sepal.Length Sepal.Width Petal.Length Petal.Width Species
1          5.1        3.5         1.4        0.2    setosa
2          4.9        3.0         1.4        0.2    setosa
3          4.7        3.2         1.3        0.2    setosa
4          4.6        3.1         1.5        0.2    setosa
5          5.0        3.6         1.4        0.2    setosa
6          5.4        3.9         1.7        0.4    setosa
```

(contains only data for setosa group).

One example operation would be to compute correlation matrix per iris specie; one would then use `lapply`:

```
> (lcor <- lapply(liris, FUN=function(df) cor(df[,1:4])))

$setosa
  Sepal.Length Sepal.Width Petal.Length Petal.Width
Sepal.Length  1.0000000  0.7425467  0.2671758  0.2780984
Sepal.Width   0.7425467  1.0000000  0.1777000  0.2327520
Petal.Length  0.2671758  0.1777000  1.0000000  0.3316300
Petal.Width   0.2780984  0.2327520  0.3316300  1.0000000

$versicolor
  Sepal.Length Sepal.Width Petal.Length Petal.Width
Sepal.Length  1.0000000  0.5259107  0.7540490  0.5464611
Sepal.Width   0.5259107  1.0000000  0.5605221  0.6639987
Petal.Length  0.7540490  0.5605221  1.0000000  0.7866681
Petal.Width   0.5464611  0.6639987  0.7866681  1.0000000

$virginica
  Sepal.Length Sepal.Width Petal.Length Petal.Width
Sepal.Length  1.0000000  0.4572278  0.8642247  0.2811077
Sepal.Width   0.4572278  1.0000000  0.4010446  0.5377280
Petal.Length  0.8642247  0.4010446  1.0000000  0.3221082
Petal.Width   0.2811077  0.5377280  0.3221082  1.0000000
```

Then we can retrieve per group the best pair of correlated variables: (correlation matrix is reshaped/melted, diagonal is filtered out and selecting best record is performed)

```
> library(reshape)
> (topcor <- lapply(lcor, FUN=function(cormat){
  correlations <- melt(cormat, variable_name="correlation");
  filtered <- correlations[correlations$X1 != correlations$X2,];
  filtered[which.max(filtered$correlation),]
}))
```

```

$setosa
      X1          X2    correlation
2 Sepal.Width Sepal.Length      0.7425467

$versicolor
      X1          X2    correlation
12 Petal.Width Petal.Length     0.7866681

$virginica
      X1          X2    correlation
3 Petal.Length Sepal.Length    0.8642247

```

Note that one computations are performed on such groupwise level, one may be interested in stacking the results, which can be done with:

```

> (result <- do.call("rbind", topcor))

      X1          X2    correlation
setosa   Sepal.Width Sepal.Length      0.7425467
versicolor  Petal.Width Petal.Length     0.7866681
virginica  Petal.Length Sepal.Length    0.8642247

```

Chapter 20: Reading and writing tabular data in plain-text files (CSV, TSV, etc.)

Parameter	Details
file	name of the CSV file to read
header	logical: does the .csv file contain a header row with column names?
sep	character: symbol that separates the cells on each row
quote	character: symbol used to quote character strings
dec	character: symbol used as decimal separator
fill	logical: when TRUE, rows with unequal length are filled with blank fields.
comment.char	character: character used as comment in the csv file. Lines preceded by this character are ignored.
...	extra arguments to be passed to <code>read.table</code>

Section 20.1: Importing .csv files

Importing using base R

Comma separated value files (CSVs) can be imported using `read.csv`, which wraps `read.table`, but uses `sep = ","` to set the delimiter to a comma.

```
# get the file path of a CSV included in R's utils package
csv_path <- system.file("misc", "exDIF.csv", package = "utils")

# path will vary based on installation location
csv_path
## [1] "/Library/Frameworks/R.framework/Resources/library/utils/misc/exDIF.csv"

df <- read.csv(csv_path)

df
##   Var1 Var2
## 1  2.70    A
## 2  3.14    B
## 3 10.00    A
## 4 -7.00    A
```

A user friendly option, `file.choose`, allows to browse through the directories:

```
df <- read.csv(file.choose())
```

Notes

- Unlike `read.table`, `read.csv` defaults to `header = TRUE`, and uses the first row as column names.
- All these functions will convert strings to `factor` class by default unless either `as.is = TRUE` or `stringsAsFactors = FALSE`.
- The `read.csv2` variant defaults to `sep = ";"` and `dec = ","` for use on data from countries where the comma is used as a decimal point and the semicolon as a field separator.

Importing using packages

The `readr` package's `read_csv` function offers much faster performance, a progress bar for large files, and more popular default options than standard `read.csv`, including `stringsAsFactors = FALSE`.

```

library(readr)

df <- read_csv(csv_path)

df
## # A tibble: 4 × 2
##   Var1   Var2
##   <dbl> <chr>
## 1 2.70    A
## 2 3.14    B
## 3 10.00   A
## 4 -7.00   A

```

Section 20.2: Importing with data.table

The `data.table` package introduces the function `fread`. While it is similar to `read.table`, `fread` is usually faster and more flexible, guessing the file's delimiter automatically.

```

# get the file path of a CSV included in R's utils package
csv_path <- system.file("misc", "exDIF.csv", package = "utils")

# path will vary based on R installation location
csv_path
## [1] "/Library/Frameworks/R.framework/Resources/library/utils/misc/exDIF.csv"

dt <- fread(csv_path)

dt
##      Var1 Var2
## 1: 2.70    A
## 2: 3.14    B
## 3: 10.00   A
## 4: -7.00   A

```

Where argument `input` is a string representing:

- the filename (e.g. `"filename.csv"`),
- a shell command that acts on a file (e.g. `"grep 'word' filename"`), or
- the input itself (e.g. `"input1, input2 \n A, B \n C, D"`).

`fread` returns an object of class `data.table` that inherits from class `data.frame`, suitable for use with the `data.table`'s usage of `[]`. To return an ordinary `data.frame`, set the `data.table` parameter to `FALSE`:

```

df <- fread(csv_path, data.table = FALSE)

class(df)
## [1] "data.frame"

df
##      Var1 Var2
## 1: 2.70    A
## 2: 3.14    B
## 3: 10.00   A
## 4: -7.00   A

```

Notes

- `fread` does not have all same options as `read.table`. One missing argument is `na.comment`, which may lead

- in unwanted behaviors if the source file contains #.
- fread uses only " for quote parameter.
- fread uses few (5) lines to guess variables types.

Section 20.3: Exporting .csv files

Exporting using base R

Data can be written to a CSV file using `write.csv()`:

```
write.csv(mtcars, "mtcars.csv")
```

Commonly-specified parameters include `row.names = FALSE` and `na = ""`.

Exporting using packages

`readr::write_csv` is significantly faster than `write.csv` and does not write row names.

```
library(readr)
write_csv(mtcars, "mtcars.csv")
```

Section 20.4: Import multiple csv files

```
files = list.files(pattern=".csv")
data_list = lapply(files, read.table, header = TRUE)
```

This reads every file and adds it to a list. Afterwards, if all data.frames have the same structure they can be combined into one big data.frame:

```
df <- do.call(rbind, data_list)
```

Section 20.5: Importing fixed-width files

Fixed-width files are text files in which columns are not separated by any character delimiter, like , or ;, but rather have a fixed character length (*width*). Data is usually padded with white spaces.

An example:

Column1	Column2	Column3	Column4	Column5
1647	pi	'important'	3.141596	28318
1731	euler	'quite important'	2.718285	43656
1979	answer	'The Answer.'	42	42

Let's assume this data table exists in the local file `constants.txt` in the working directory.

Importing with base R

```
df <- read.fwf('constants.txt', widths = c(8,10,18,7,8), header = FALSE, skip = 1)

df
#>      V1     V2          V3     V4     V5
```

```
#> 1 1647      pi          'important'  3.14159  6.28318
#> 2 1731    euler  'quite important' 2.71828  5.43656
#> 3 1979   answer     'The Answer.'  42         42.0000
```

Note:

- Column titles don't need to be separated by a character (Column4Column5)
- The widths parameter defines the width of each column
- Non-separated headers are not readable with `read.fwf()`

Importing with readr

```
library(readr)

df <- read_fwf('constants.txt',
                 fwf_cols(Year = 8, Name = 10, Importance = 18, Value = 7, Doubled = 8),
                 skip = 1)
df
#> # A tibble: 3 x 5
#>   Year     Name     Importance     Value  Doubled
#>   <int>    <chr>    <chr>       <dbl>    <dbl>
#> 1 1647     pi     'important'  3.14159  6.28318
#> 2 1731    euler  'quite important' 2.71828  5.43656
#> 3 1979   answer     'The Answer.'  42.0000 42.00000
```

Note:

- `readr`'s `fwf_*` helper functions offer alternative ways of specifying column lengths, including automatic guessing (`fwf_empty`)
- `readr` is faster than base R
- Column titles cannot be automatically imported from data file

Chapter 21: Pipe operators (%>% and others)

Ihs

rhs

A value or the magrittr placeholder. A function call using the magrittr semantics

Pipe operators, available in `magrittr`, `dplyr`, and other R packages, process a data-object using a sequence of operations by passing the result of one step as input for the next step using infix-operators rather than the more typical R method of nested function calls.

Note that the intended aim of pipe operators is to increase human readability of written code. See Remarks section for performance considerations.

Section 21.1: Basic use and chaining

The pipe operator, `%>%`, is used to insert an argument into a function. It is not a base feature of the language and can only be used after attaching a package that provides it, such as `magrittr`. The pipe operator takes the left-hand side (LHS) of the pipe and uses it as the first argument of the function on the right-hand side (RHS) of the pipe. For example:

```
library(magrittr)

1:10 %>% mean
# [1] 5.5

# is equivalent to
mean(1:10)
# [1] 5.5
```

The pipe can be used to replace a sequence of function calls. Multiple pipes allow us to read and write the sequence from left to right, rather than from inside to out. For example, suppose we have `years` defined as a factor but want to convert it to a numeric. To prevent possible information loss, we first convert to character and then to numeric:

```
years <- factor(2008:2012)

# nesting
as.numeric(as.character(years))

# piping
years %>% as.character %>% as.numeric
```

If we don't want the LHS (Left Hand Side) used as the *first* argument on the RHS (Right Hand Side), there are workarounds, such as naming the arguments or using `.` to indicate where the piped input goes.

```
# example with grep
# its syntax:
# grep(pattern, x, ignore.case = FALSE, perl = FALSE, fixed = FALSE, useBytes = FALSE)

# note that the `substring` result is the *2nd* argument of grep
grep("Wo", substring("Hello World", 7, 11))

# piping while naming other arguments
"Hello World" %>% substring(7, 11) %>% grep(pattern = "Wo")
```

```

# piping with .
"Hello World" %>% substring(7, 11) %>% grep("Wo", .)

# piping with . and curly braces
"Hello World" %>% substring(7, 11) %>% { c(paste('Hi', .)) }
#[1] "Hi World"

#using LHS multiple times in argument with curly braces and .
"Hello World" %>% substring(7, 11) %>% { c(paste(. , 'Hi', .)) }
#[1] "World Hi World"

```

Section 21.2: Functional sequences

Given a sequence of steps we use repeatedly, it's often handy to store it in a function. Pipes allow for saving such functions in a readable format by starting a sequence with a dot as in:

```
. %>% RHS
```

As an example, suppose we have factor dates and want to extract the year:

```

library(magrittr) # needed to include the pipe operators
library(lubridate)
read_year <- . %>% as.character %>% as.Date %>% year

# Creating a dataset
df <- data.frame(now = "2015-11-11", before = "2012-01-01")
#           now      before
# 1 2015-11-11 2012-01-01

# Example 1: applying `read_year` to a single character-vector
df$now %>% read_year
# [1] 2015

# Example 2: applying `read_year` to all columns of `df`
df %>% lapply(read_year) %>% as.data.frame # implicit `lapply(df, read_year)`
#       now      before
# 1 2015 2012

# Example 3: same as above using `mutate_all`
library(dplyr)
df %>% mutate_all(fun(read_year))
# if an older version of dplyr use `mutate_each`
#       now      before
# 1 2015 2012

```

We can review the composition of the function by typing its name or using functions:

```

read_year
# Functional sequence with the following components:
#
# 1. as.character(.)
# 2. as.Date(.)
# 3. year(.)
#
# Use 'functions' to extract the individual functions.

```

We can also access each function by its position in the sequence:

```
read_year[[2]]
```

```
# function (.)
# as.Date(.)
```

Generally, this approach may be useful when clarity is more important than speed.

Section 21.3: Assignment with %<>%

The `magrittr` package contains a compound assignment infix-operator, `%<>%`, that updates a value by first piping it into one or more `rhs` expressions and then assigning the result. This eliminates the need to type an object name twice (once on each side of the assignment operator `<-`). `%<>%` must be the first infix-operator in a chain:

```
library(magrittr)
library(dplyr)

df <- mtcars
```

Instead of writing

```
df <- df %>% select(1:3) %>% filter(mpg > 20, cyl == 6)
```

or

```
df %>% select(1:3) %>% filter(mpg > 20, cyl == 6) -> df
```

The compound assignment operator will both pipe and reassign `df`:

```
df %<>% select(1:3) %>% filter(mpg > 20, cyl == 6)
```

Section 21.4: Exposing contents with %\$%

The exposition pipe operator, `%$%`, exposes the column names as R symbols within the left-hand side object to the right-hand side expression. This operator is handy when piping into functions that do not have a `data` argument (unlike, say, `lm`) and that don't take a `data.frame` and column names as arguments (most of the main `dplyr` functions).

The exposition pipe operator `%$%` allows a user to avoid breaking a pipeline when needing to refer to column names. For instance, say you want to filter a `data.frame` and then run a correlation test on two columns with `cor.test`:

```
library(magrittr)
library(dplyr)
mtcars %>%
  filter(wt > 2) %$%
  cor.test(hp, mpg)

#>
#> Pearson's product-moment correlation
#>
#> data: hp and mpg
#> t = -5.9546, df = 26, p-value = 2.768e-06
#> alternative hypothesis: true correlation is not equal to 0
#> 95 percent confidence interval:
#> -0.8825498 -0.5393217
#> sample estimates:
#>       cor
```

```
#> -0.7595673
```

Here the standard `%>%` pipe passes the `data.frame` through to `filter()`, while the `%%$` pipe exposes the column names to `cor.test()`.

The exposition pipe works like a pipe-able version of the base R `with()` functions, and the same left-hand side objects are accepted as inputs.

Section 21.5: Creating side effects with `%T>%`

Some functions in R produce a side effect (i.e. saving, printing, plotting, etc) and do not always return a meaningful or desired value.

`%T>%` (tee operator) allows you to forward a value into a side-effect-producing function while keeping the original `lhs` value intact. In other words: the tee operator works like `%>%`, except the return values is `lhs` itself, and not the result of the `rhs` function/expression.

Example: Create, pipe, write, and return an object. If `%>%` were used in place of `%T>%` in this example, then the variable `all_letters` would contain `NULL` rather than the value of the sorted object.

```
all_letters <- c(letters, LETTERS) %>%
  sort %T>%
  write.csv(file = "all_letters.csv")

read.csv("all_letters.csv") %>% head()
#   x
# 1 a
# 2 A
# 3 b
# 4 B
# 5 c
# 6 C
```

Warning: Piping an unnamed object to `save()` will produce an object named `.` when loaded into the workspace with `load()`. However, a workaround using a helper function is possible (which can also be written inline as an anonymous function).

```
all_letters <- c(letters, LETTERS) %>%
  sort %T>%
  save(file = "all_letters.RData")

load("all_letters.RData", e <- new.env())

get("all_letters", envir = e)
# Error in get("all_letters", envir = e) : object 'all_letters' not found

get(".", envir = e)
# [1] "a"  "A"  "b"  "B"  "c"  "C"  "d"  "D"  "e"  "E"  "f"  "F"  "g"  "G"  "h"  "H"  "i"  "I"  "j"  "J"
# [21] "k"  "K"  "l"  "L"  "m"  "M"  "n"  "N"  "o"  "O"  "p"  "P"  "q"  "Q"  "r"  "R"  "s"  "S"  "t"  "T"
# [41] "u"  "U"  "v"  "V"  "w"  "W"  "x"  "X"  "y"  "Y"  "z"  "Z"

# Work-around
save2 <- function(. = ., name, file = stop("'file' must be specified")) {
  assign(name, .)
  call_save <- call("save", ... = name, file = file)
  eval(call_save)
}
```

```
all_letters <- c(letters, LETTERS) %>%
  sort %T>%
  save2("all_letters", "all_letters.RData")
```

Section 21.6: Using the pipe with dplyr and ggplot2

The `%>%` operator can also be used to pipe the dplyr output into ggplot. This creates a unified exploratory data analysis (EDA) pipeline that is easily customizable. This method is faster than doing the aggregations internally in ggplot and has the added benefit of avoiding unnecessary intermediate variables.

```
library(dplyr)
library(ggplot)

diamonds %>%
  filter(depth > 60) %>%
  group_by(cut) %>%
  summarize(mean_price = mean(price)) %>%
  ggplot(aes(x = cut, y = mean_price)) +
  geom_bar(stat = "identity")
```

Chapter 22: Linear Models (Regression)

Parameter	Meaning
formula	a formula in <i>Wilkinson-Rogers</i> notation; response ~ ... where ... contains terms corresponding to variables in the environment or in the data frame specified by the data argument
data	data frame containing the response and predictor variables
subset	a vector specifying a subset of observations to be used: may be expressed as a logical statement in terms of the variables in data
weights	analytical weights (see <i>Weights</i> section above)
na.action	how to handle missing (NA) values: see ?na.action
method	how to perform the fitting. Only choices are "qr" or "model.frame" (the latter returns the model frame without fitting the model, identical to specifying model=TRUE)
model	whether to store the model frame in the fitted object
x	whether to store the model matrix in the fitted object
y	whether to store the model response in the fitted object
qr	whether to store the QR decomposition in the fitted object
singular.ok	whether to allow <i>singular fits</i> , models with collinear predictors (a subset of the coefficients will automatically be set to NA in this case)
contrasts	a list of contrasts to be used for particular factors in the model; see the contrasts.arg argument of ?model.matrix.default . Contrasts can also be set with options() (see the contrasts argument) or by assigning the contrast attributes of a factor (see ?contrasts)
offset	used to specify an <i>a priori</i> known component in the model. May also be specified as part of the formula. See ?model.offset
...	additional arguments to be passed to lower-level fitting functions (lm.fit() or lm.wfit())

Section 22.1: Linear regression on the mtcars dataset

The built-in mtcars data frame contains information about 32 cars, including their weight, fuel efficiency (in miles-per-gallon), speed, etc. (To find out more about the dataset, use **help(mtcars)**).

If we are interested in the relationship between fuel efficiency (mpg) and weight (wt) we may start plotting those variables with:

```
plot(mpg ~ wt, data = mtcars, col=2)
```

The plots shows a (linear) relationship!. Then if we want to perform linear regression to determine the coefficients of a linear model, we would use the **lm** function:

```
fit <- lm(mpg ~ wt, data = mtcars)
```

The ~ here means "explained by", so the formula **mpg ~ wt** means we are predicting mpg as explained by wt. The most helpful way to view the output is with:

```
summary(fit)
```

Which gives the output:

```
Call:  
lm(formula = mpg ~ wt, data = mtcars)  
  
Residuals:  
    Min     1Q Median     3Q    Max  
-2.50 -0.99 -0.12  0.88 14.50
```

```
-4.5432 -2.3647 -0.1252 1.4096 6.8727
```

Coefficients:

	Estimate	Std. Error	t value	Pr(> t)
(Intercept)	37.2851	1.8776	19.858	< 2e-16 ***
wt	-5.3445	0.5591	-9.559	1.29e-10 ***

Signif. codes: 0 ‘***’ 0.001 ‘**’ 0.01 ‘*’ 0.05 ‘.’ 0.1 ‘ ’ 1

Residual standard error: 3.046 on 30 degrees of freedom

Multiple R-squared: 0.7528, Adjusted R-squared: 0.7446

F-statistic: 91.38 on 1 and 30 DF, p-value: 1.294e-10

This provides information about:

- the estimated slope of each coefficient (wt and the y-intercept), which suggests the best-fit prediction of mpg is $37.2851 + (-5.3445) * \text{wt}$
- The p-value of each coefficient, which suggests that the intercept and weight are probably not due to chance
- Overall estimates of fit such as R^2 and adjusted R^2 , which show how much of the variation in mpg is explained by the model

We could add a line to our first plot to show the predicted mpg:

```
abline(fit, col=3, lwd=2)
```

It is also possible to add the equation to that plot. First, get the coefficients with `coef`. Then using `paste0` we collapse the coefficients with appropriate variables and +/-, to built the equation. Finally, we add it to the plot using `mtext`:

```
bs <- round(coef(fit), 3)
lmlab <- paste0("mpg = ", bs[1],
                 ifelse(sign(bs[2]) == 1, " + ", " - "), abs(bs[2]), " wt ")
mtext(lmlab, 3, line=-2)
```

The result is:



Section 22.2: Using the 'predict' function

Once a model is built `predict` is the main function to test with new data. Our example will use the `mtcars` built-in dataset to regress miles per gallon against displacement:

```
my_mdl <- lm(mpg ~ disp, data=mtcars)
my_mdl

Call:
lm(formula = mpg ~ disp, data = mtcars)

Coefficients:
(Intercept)      disp
 29.59985     -0.04122
```

If I had a new data source with displacement I could see the estimated miles per gallon.

```
set.seed(1234)
newdata <- sample(mtcars$disp, 5)
newdata
[1] 258.0 71.1 75.7 145.0 400.0

newdf <- data.frame(disp=newdata)
predict(my_mdl, newdf)
1          2          3          4          5
18.96635 26.66946 26.47987 23.62366 13.11381
```

The most important part of the process is to create a new data frame with the same column names as the original data. In this case, the original data had a column labeled `disp`, I was sure to call the new data that same name.

Caution

Let's look at a few common pitfalls:

1. not using a data.frame in the new object:

```
predict(my_mdl, newdata)
Error in eval(predvars, data, env) :
  numeric 'envir' arg not of length one
```

2. not using same names in new data frame:

```
newdf2 <- data.frame(newdata)
predict(my_mdl, newdf2)
Error in eval(expr, envir, enclos) : object 'disp' not found
```

Accuracy

To check the accuracy of the prediction you will need the actual y values of the new data. In this example, newdf will need a column for 'mpg' and 'disp'.

```
newdf <- data.frame(mpg=mtcars$mpg[1:10], disp=mtcars$disp[1:10])
#   mpg   disp
# 1 21.0 160.0
# 2 21.0 160.0
# 3 22.8 108.0
# 4 21.4 258.0
# 5 18.7 360.0
# 6 18.1 225.0
# 7 14.3 360.0
# 8 24.4 146.7
# 9 22.8 140.8
# 10 19.2 167.6

p <- predict(my_mdl, newdf)

#root mean square error
sqrt(mean((p - newdf$mpg)^2, na.rm=TRUE))
[1] 2.325148
```

Section 22.3: Weighting

Sometimes we want the model to give more weight to some data points or examples than others. This is possible by specifying the weight for the input data while learning the model. There are generally two kinds of scenarios where we might use non-uniform weights over the examples:

- Analytic Weights: Reflect the different levels of precision of different observations. For example, if analyzing data where each observation is the average results from a geographic area, the analytic weight is proportional to the inverse of the estimated variance. Useful when dealing with averages in data by providing a proportional weight given the number of observations. [Source](#)
- Sampling Weights (Inverse Probability Weights - IPW): a statistical technique for calculating statistics standardized to a population different from that in which the data was collected. Study designs with a disparate sampling population and population of target inference (target population) are common in application. Useful when dealing with data that have missing values. [Source](#)

The `lm()` function does analytic weighting. For sampling weights the survey package is used to build a survey design object and run `svyglm()`. By default, the survey package uses sampling weights. (NOTE: `lm()`, and `svyglm()` with family `gaussian()` will all produce the same point estimates, because they both solve for the coefficients by minimizing the weighted least squares. They differ in how standard errors are calculated.)

Test Data

```
data <- structure(list(lexptot = c(9.1595012302023, 9.86330744180814,  
8.92372556833205, 8.58202430280175, 10.1133857229336), progvillm = c(1L,  
1L, 1L, 0L), sexhead = c(1L, 1L, 0L, 1L, 1L), agehead = c(79L,  
43L, 52L, 48L, 35L), weight = c(1.04273509979248, 1.01139605045319,  
1.01139605045319, 1.01139605045319, 0.76305216550827)), .Names = c("lexptot",  
"progvillm", "sexhead", "agehead", "weight"), class = c("tbl_df",  
"tbl", "data.frame"), row.names = c(NA, -5L))
```

Analytic Weights

```
lm.analytic <- lm(lexptot ~ progvillm + sexhead + agehead,  
                   data = data, weight = weight)  
summary(lm.analytic)
```

Output

```
Call:  
lm(formula = lexptot ~ progvillm + sexhead + agehead, data = data,  
  weights = weight)  
  
Weighted Residuals:  
      1       2       3       4       5  
9.249e-02 5.823e-01 0.000e+00 -6.762e-01 -1.527e-16  
  
Coefficients:  
            Estimate Std. Error t value Pr(>|t|)  
(Intercept) 10.016054   1.744293   5.742   0.110  
progvillm   -0.781204   1.344974  -0.581   0.665  
sexhead      0.306742   1.040625   0.295   0.818  
agehead     -0.005983   0.032024  -0.187   0.882  
  
Residual standard error: 0.8971 on 1 degrees of freedom  
Multiple R-squared:  0.467, Adjusted R-squared:  -1.132  
F-statistic: 0.2921 on 3 and 1 DF,  p-value: 0.8386
```

Sampling Weights (IPW)

```
library(survey)  
data$X <- 1:nrow(data) # Create unique id  
  
# Build survey design object with unique id, ipw, and data.frame  
des1 <- svydesign(id = ~X, weights = ~weight, data = data)  
  
# Run glm with survey design object  
prog.lm <- svyglm(lexptot ~ progvillm + sexhead + agehead, design=des1)
```

Output

```
Call:  
svyglm(formula = lexptot ~ progvillm + sexhead + agehead, design = des1)
```

```

Survey design:
svydesign(id = ~X, weights = ~weight, data = data)

Coefficients:
            Estimate Std. Error t value Pr(>|t|) 
(Intercept) 10.016054  0.183942 54.452   0.0117 *
progvillm -0.781204  0.640372 -1.220   0.4371 
sexhead     0.306742  0.397089  0.772   0.5813 
agehead    -0.005983  0.014747 -0.406   0.7546 
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

(Dispersion parameter for gaussian family taken to be 0.2078647)

Number of Fisher Scoring iterations: 2

```

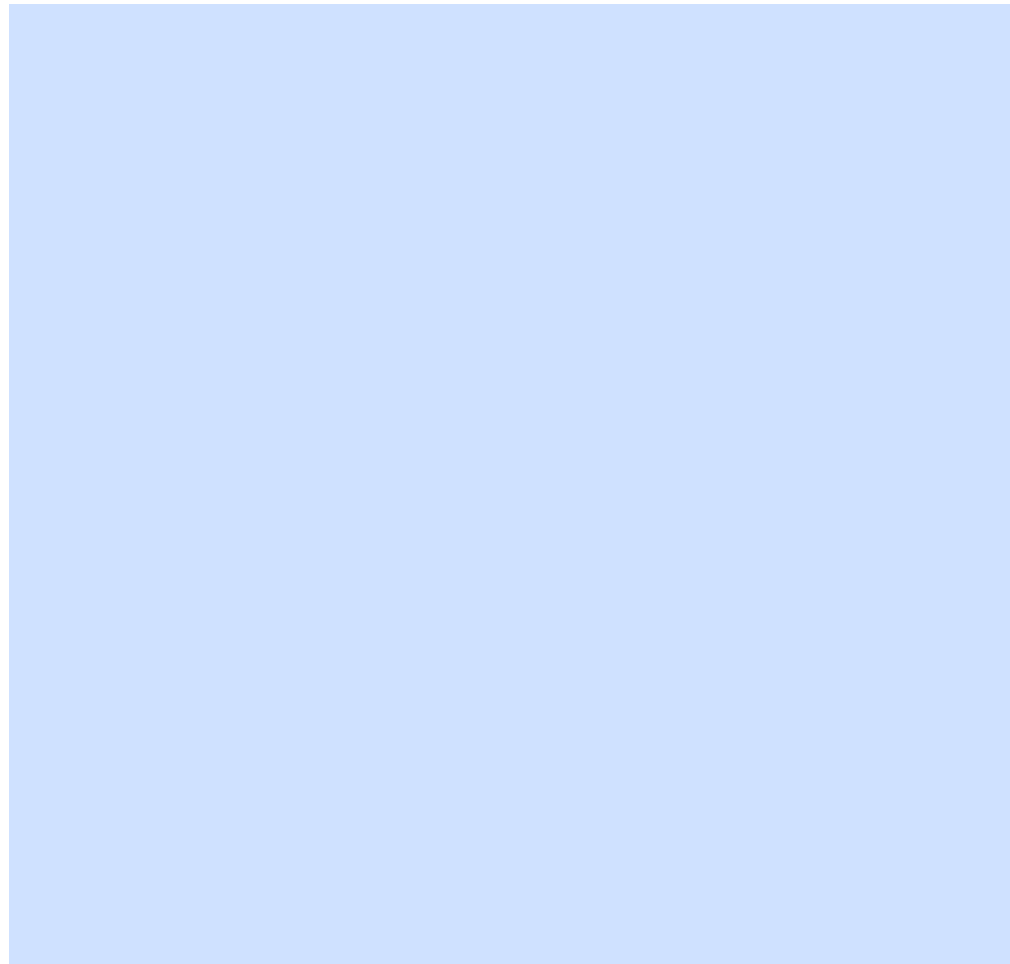
Section 22.4: Checking for nonlinearity with polynomial regression

Sometimes when working with linear regression we need to check for non-linearity in the data. One way to do this is to fit a polynomial model and check whether it fits the data better than a linear model. There are other reasons, such as theoretical, that indicate to fit a quadratic or higher order model because it is believed that the variables relationship is inherently polynomial in nature.

Let's fit a quadratic model for the `mtcars` dataset. For a linear model see Linear regression on the mtcars dataset.

First we make a scatter plot of the variables `mpg` (Miles/gallon), `disp` (Displacement (cu.in.)), and `wt` (Weight (1000 lbs)). The relationship among `mpg` and `disp` appears non-linear.

```
plot(mtcars[,c("mpg", "disp", "wt")])
```



A linear fit will show that disp is not significant.

```
fit0 = lm(mpg ~ wt+disp, mtcars)
summary(fit0)

# Coefficients:
#             Estimate Std. Error t value Pr(>|t|)
#(Intercept) 34.96055   2.16454 16.151 4.91e-16 ***
#wt          -3.35082   1.16413 -2.878  0.00743 **
#disp         -0.01773   0.00919 -1.929  0.06362 .
#---
#Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
#Residual standard error: 2.917 on 29 degrees of freedom
#Multiple R-squared:  0.7809,    Adjusted R-squared:  0.7658
```

Then, to get the result of a quadratic model, we added `I(disp^2)`. The new model appears better when looking at R^2 and all variables are significant.

```
fit1 = lm(mpg ~ wt+disp+I(disp^2), mtcars)
summary(fit1)

# Coefficients:
#             Estimate Std. Error t value Pr(>|t|)
#(Intercept) 41.4019837  2.4266906 17.061 2.5e-16 ***
#wt          -3.4179165  0.9545642 -3.581 0.001278 **
#disp         -0.0823950  0.0182460 -4.516 0.000104 ***
#I(disp^2)    0.0001277  0.0000328  3.892 0.000561 ***
#---
#Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
#Residual standard error: 2.391 on 28 degrees of freedom
```

```
#Multiple R-squared:  0.8578,    Adjusted R-squared:  0.8426
```

As we have three variables, the fitted model is a surface represented by:

```
mpg = 41.4020 - 3.4179 * wt - 0.0824 * disp + 0.0001277 * disp^2
```

Another way to specify polynomial regression is using `poly` with parameter `raw=TRUE`, otherwise *orthogonal polynomials* will be considered (see the `help(ploy)` for more information). We get the same result using:

```
summary(lm(mpg ~ wt+poly(disp, 2, raw=TRUE), mtcars))
```

Finally, what if we need to show a plot of the estimated surface? Well there are many options to make 3D plots in R. Here we use Fit3d from p3dpackage.

```
library(p3d)
Init3d(family="serif", cex = 1)
Plot3d(mpg ~ disp+wt, mtcars)
Axes3d()
Fit3d(fit1)
```



Section 22.5: Plotting The Regression (base)

Continuing on the `mtcars` example, here is a simple way to produce a plot of your linear regression that is potentially suitable for publication.

First fit the linear model and

```
fit <- lm(mpg ~ wt, data = mtcars)
```

Then plot the two variables of interest and add the regression line within the definition domain:

```
plot(mtcars$wt, mtcars$mpg, pch=18, xlab = 'wt', ylab = 'mpg')
lines(c(min(mtcars$wt), max(mtcars$wt)),
as.numeric(predict(fit, data.frame(wt=c(min(mtcars$wt), max(mtcars$wt))))))
```

Almost there! The last step is to add to the plot, the regression equation, the rsquare as well as the correlation coefficient. This is done using the **vector** function:

```
rp = vector('expression', 3)
rp[1] = substitute(expression(italic(y) == MYOTHERVALUE3 + MYOTHERVALUE4 %*% x),
list(MYOTHERVALUE3 = format(fit$coefficients[1], digits = 2),
MYOTHERVALUE4 = format(fit$coefficients[2], digits = 2)))[2]
rp[2] = substitute(expression(italic(R)^2 == MYVALUE),
list(MYVALUE = format(summary(fit)$adj.r.squared, dig=3)))[2]
rp[3] = substitute(expression(Pearson-R == MYOTHERVALUE2),
list(MYOTHERVALUE2 = format(cor(mtcars$wt, mtcars$mpg), digits = 2)))[2]

legend("topright", legend = rp, bty = 'n')
```

Note that you can add any other parameter such as the RMSE by adapting the vector function. Imagine you want a legend with 10 elements. The vector definition would be the following:

```
rp = vector('expression', 10)
```

and you will need to defined r[1].... to r[10]

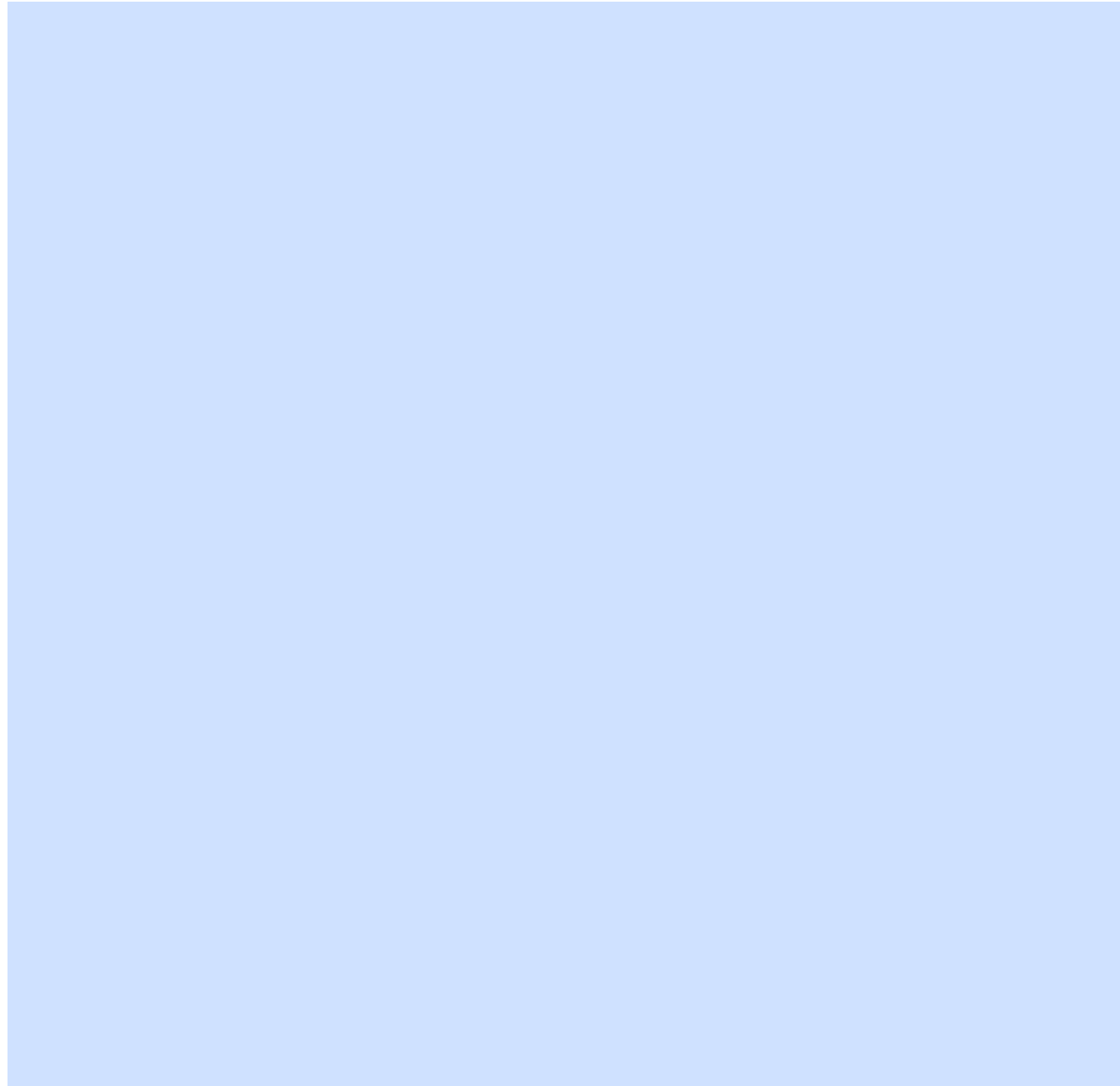
Here is the output:



Section 22.6: Quality assessment

After building a regression model it is important to check the result and decide if the model is appropriate and works well with the data at hand. This can be done by examining the residuals plot as well as other diagnostic plots.

```
# fit the model
fit <- lm(mpg ~ wt, data = mtcars)
#
par(mfrow=c(2,1))
# plot model object
plot(fit, which =1:2)
```



These plots check for two assumptions that were made while building the model:

1. That the expected value of the predicted variable (in this case mpg) is given by a linear combination of the predictors (in this case wt). We expect this estimate to be unbiased. So the residuals should be centered around the mean for all values of the predictors. In this case we see that the residuals tend to be positive at the ends and negative in the middle, suggesting a non-linear relationship between the variables.
2. That the actual predicted variable is normally distributed around its estimate. Thus, the residuals should be

normally distributed. For normally distributed data, the points in a normal Q-Q plot should lie on or close to the diagonal. There is some amount of skew at the ends here.

Chapter 23: data.table

Data.table is a package that extends the functionality of data frames from base R, particularly improving on their performance and syntax. See the package's Docs area at Getting started with data.table for details.

Section 23.1: Creating a data.table

A data.table is an enhanced version of the data.frame class from base R. As such, its `class()` attribute is the vector "data.table" "data.frame" and functions that work on a data.frame will also work with a data.table. There are many ways to create, load or coerce to a data.table.

Build

Don't forget to install and activate the data.table package

```
library(data.table)
```

There is a constructor of the same name:

```
DT <- data.table(  
  x = letters[1:5],  
  y = 1:5,  
  z = (1:5) > 3  
)  
#   x y     z  
# 1: a 1 FALSE  
# 2: b 2 FALSE  
# 3: c 3 FALSE  
# 4: d 4  TRUE  
# 5: e 5  TRUE
```

Unlike `data.frame`, `data.table` will not coerce strings to factors:

```
sapply(DT, class)  
#           x          y          z  
#   "character"  "integer"  "logical"
```

Read in

We can read from a text file:

```
dt <- fread("my_file.csv")
```

Unlike `read.csv`, `fread` will read strings as strings, not as factors.

Modify a data.frame

For efficiency, `data.table` offers a way of altering a `data.frame` or `list` to make a `data.table` in-place (without making a copy or changing its memory location):

```
# example data.frame  
DF <- data.frame(x = letters[1:5], y = 1:5, z = (1:5) > 3)  
# modification  
setDT(DF)
```

Note that we do not `<-` assign the result, since the object `DF` has been modified in-place. The `class` attributes of the

data.frame will be retained:

```
sapply(DF, class)
#      x      y      z
# "factor" "integer" "logical"
```

Coerce object to data.table

If you have a **list**, **data.frame**, or **data.table**, you should use the **setDT** function to convert to a **data.table** because it does the conversion by reference instead of making a copy (which **as.data.table** does). This is important if you are working with large datasets.

If you have another R object (such as a matrix), you must use **as.data.table** to coerce it to a **data.table**.

```
mat <- matrix(0, ncol = 10, nrow = 10)

DT <- as.data.table(mat)
# or
DT <- data.table(mat)
```

Section 23.2: Special symbols in data.table

.SD

.SD refers to the subset of the **data.table** for each group, excluding all columns used in **by**.

.SD along with **lapply** can be used to apply any function to multiple columns by group in a **data.table**

We will continue using the same built-in dataset, **mtcars**:

```
mtcars = data.table(mtcars) # Let's not include rownames to keep things simpler
```

Mean of all columns in the dataset by *number of cylinders*, cyl:

```
mtcars[ , lapply(.SD, mean), by = cyl]

#   cyl      mpg     disp       hp     drat      wt     qsec      vs      am     gear
carb
#1:   6 19.74286 183.3143 122.28571 3.585714 3.117143 17.97714 0.5714286 0.4285714 3.857143
3.428571
#2:   4 26.66364 105.1364  82.63636 4.070909 2.285727 19.13727 0.9090909 0.7272727 4.090909
1.545455
#3:   8 15.10000 353.1000 209.21429 3.229286 3.999214 16.77214 0.0000000 0.1428571 3.285714
3.500000
```

Apart from cyl, there are other categorical columns in the dataset such as vs, am, gear and carb. It doesn't really make sense to take the **mean** of these columns. So let's exclude these columns. This is where **.SDcols** comes into the picture.

.SDcols

.SDcols specifies the columns of the **data.table** that are included in .SD.

Mean of all columns (continuous columns) in the dataset by *number of gears* gear, and *number of cylinders*, cyl, arranged by gear and cyl:

```
# All the continuous variables in the dataset
```

```

cols_chosen <- c("mpg", "disp", "hp", "drat", "wt", "qsec")

mtcars[order(gear, cyl), lapply(.SD, mean), by = .(gear, cyl), .SDcols = cols_chosen]

#   gear cyl   mpg   disp    hp   drat     wt   qsec
#1:   3   4 21.500 120.1000 97.0000 3.700000 2.465000 20.0100
#2:   3   6 19.750 241.5000 107.5000 2.920000 3.337500 19.8300
#3:   3   8 15.050 357.6167 194.1667 3.120833 4.104083 17.1425
#4:   4   4 26.925 102.6250 76.0000 4.110000 2.378125 19.6125
#5:   4   6 19.750 163.8000 116.5000 3.910000 3.093750 17.6700
#6:   5   4 28.200 107.7000 102.0000 4.100000 1.826500 16.8000
#7:   5   6 19.700 145.0000 175.0000 3.620000 2.770000 15.5000
#8:   5   8 15.400 326.0000 299.5000 3.880000 3.370000 14.5500

```

Maybe we don't want to calculate the `mean` by groups. To calculate the mean for all the cars in the dataset, we don't specify the `by` variable.

```

mtcars[ , lapply(.SD, mean), .SDcols = cols_chosen]

#       mpg      disp      hp      drat      wt      qsec
#1: 20.09062 230.7219 146.6875 3.596563 3.21725 17.84875

```

Note:

- It is not necessary to define `cols_chosen` beforehand. `.SDcols` can directly take column names
- `.SDcols` can also directly take a vector of column numbers. In the above example this would be `mtcars[, lapply(.SD, mean), .SDcols = c(1,3:7)]`

.N

`.N` is shorthand for the number of rows in a group.

```

iris[, .(count=.N), by=Species]

#       Species count
#1:     setosa     50
#2: versicolor     50
#3: virginica     50

```

Section 23.3: Adding and modifying columns

`DT[where, select|update|do, by]` syntax is used to work with columns of a data.table.

- The "where" part is the `i` argument
- The "select|update|do" part is the `j` argument

These two arguments are usually passed by position instead of by name.

Our example data below is

```
mtcars = data.table(mtcars, keep.rownames = TRUE)
```

Editing entire columns

Use the `:=` operator inside `j` to assign new columns:

```
mtcars[, mpg_sq := mpg^2]
```

Remove columns by setting to NULL:

```
mtcars[, mpg_sq := NULL]
```

Add multiple columns by using the `:=` operator's multivariate format:

```
mtcars[, `:=`(mpg_sq = mpg^2, wt_sqrt = sqrt(wt))]
# or
mtcars[, c("mpg_sq", "wt_sqrt") := .(mpg^2, sqrt(wt))]
```

If the columns are dependent and must be defined in sequence, one way is:

```
mtcars[, c("mpg_sq", "mpg2_hp") := .(temp1 <- mpg^2, temp1/hp)]
```

The `.()` syntax is used when the right-hand side of LHS `:=` RHS is a list of columns.

For dynamically-determined column names, use parentheses:

```
vn = "mpg_sq"
mtcars[, (vn) := mpg^2]
```

Columns can also be modified with `set`, though this is rarely necessary:

```
set(mtcars, j = "hp_over_wt", v = mtcars$hp/mtcars$wt)
```

Editing subsets of columns

Use the `i` argument to subset to rows "where" edits should be made:

```
mtcars[1:3, newvar := "Hello"]
# or
set(mtcars, j = "newvar", i = 1:3, v = "Hello")
```

As in a data.frame, we can subset using row numbers or logical tests. It is also possible to use a "join" in `i`, but that more complicated task is covered in another example.

Editing column attributes

Functions that edit attributes, such as `levels<-` or `names<-`, actually replace an object with a modified copy. Even if only used on one column in a data.table, the entire object is copied and replaced.

To modify an object without copies, use `setnames` to change the column names of a data.table or data.frame and `setattr` to change an attribute for any object.

```
# Print a message to the console whenever the data.table is copied
tracemem(mtcars)
mtcars[, cyl2 := factor(cyl)]

# Neither of these statements copy the data.table
setnames(mtcars, old = "cyl2", new = "cyl_fac")
setattr(mtcars$cyl_fac, "levels", c("four", "six", "eight"))

# Each of these statements copies the data.table
names(mtcars)[names(mtcars) == "cyl_fac"] <- "cf"
levels(mtcars$cf) <- c("IV", "VI", "VIII")
```

Be aware that these changes are made by reference, so they are *global*. Changing them within one environment

affects the object in all environments.

```
# This function also changes the levels in the global environment
edit_levels <- function(x) setattr(x, "levels", c("low", "med", "high"))
edit_levels(mtcars$cyl_factor)
```

Section 23.4: Writing code compatible with both `data.frame` and `data.table`

Differences in subsetting syntax

A `data.table` is one of several two-dimensional data structures available in R, besides `data.frame`, `matrix` and (2D) `array`. All of these classes use a very similar but not identical syntax for subsetting, the `A[rows, cols]` schema.

Consider the following data stored in a `matrix`, a `data.frame` and a `data.table`:

```
ma <- matrix(rnorm(12), nrow=4, dimnames=list(letters[1:4], c('X', 'Y', 'Z')))
df <- as.data.frame(ma)
dt <- as.data.table(ma)

ma[2:3] #---> returns the 2nd and 3rd items, as if 'ma' were a vector (because it is!)
df[2:3] #---> returns the 2nd and 3rd columns
dt[2:3] #---> returns the 2nd and 3rd rows!
```

If you want to be sure of what will be returned, it is better to be *explicit*.

To get specific `rows`, just add a comma after the range:

```
ma[2:3, ] # \
df[2:3, ] # }---> returns the 2nd and 3rd rows
dt[2:3, ] # /
```

But, if you want to subset `columns`, some cases are interpreted differently. All three can be subset the same way with integer or character indices *not* stored in a variable.

```
ma[, 2:3] # \
df[, 2:3] # \
dt[, 2:3] # }---> returns the 2nd and 3rd columns
ma[, c("Y", "Z")] # /
df[, c("Y", "Z")] # /
dt[, c("Y", "Z")] # /
```

However, they differ for unquoted variable names

```
mycols <- 2:3
ma[, mycols] # \
df[, mycols] # }---> returns the 2nd and 3rd columns
dt[, mycols, with = FALSE] # /

dt[, mycols] # ---> Raises an error
```

In the last case, `mycols` is evaluated as the name of a column. Because `dt` cannot find a column named `mycols`, an error is raised.

Note: For versions of the `data.table` package prior to 1.9.8, this behavior was slightly different. Anything in the column index would have been evaluated using `dt` as an environment. So both `dt[, 2:3]` and `dt[, mycols]` would

return the vector `2:3`. No error would be raised for the second case, because the variable `mycols` does exist in the parent environment.

Strategies for maintaining compatibility with `data.frame` and `data.table`

There are many reasons to write code that is guaranteed to work with `data.frame` and `data.table`. Maybe you are forced to use `data.frame`, or you may need to share some code that you don't know how will be used. So, there are some main strategies for achieving this, in order of convenience:

1. Use syntax that behaves the same for both classes.
2. Use a common function that does the same thing as the shortest syntax.
3. Force `data.table` to behave as `data.frame` (ex.: call the specific method `print.data.frame`).
4. Treat them as `list`, which they ultimately are.
5. Convert the table to a `data.frame` before doing anything (bad idea if it is a huge table).
6. Convert the table to `data.table`, if dependencies are not a concern.

Subset rows. Its simple, just use the `[,]` selector, *with* the comma:

```
A[1:10, ]
A[A$var > 17, ] # A[var > 17, ] just works for data.table
```

Subset columns. If you want a single column, use the `$` or the `[[]]` selector:

```
A$var
colname <- 'var'
A[[colname]]
A[[1]]
```

If you want a uniform way to grab more than one column, it's necessary to appeal a bit:

```
B <- `[ .data.frame`(A, 2:4)

# We can give it a better name
select <- `[.data.frame`
B <- select(A, 2:4)
C <- select(A, c('foo', 'bar'))
```

Subset 'indexed' rows. While `data.frame` has `row.names`, `data.table` has its unique key feature. The best thing is to avoid `row.names` entirely and take advantage of the existing optimizations in the case of `data.table` when possible.

```
B <- A[A$var != 0, ]
# or...
B <- with(A, A[var != 0, ]) # data.table will silently index A by var before subsetting
stuff <- c('a', 'c', 'f')
C <- A[match(stuff, A$name), ] # really worse than: setkey(A); A[stuff, ]
```

Get a 1-column table, get a row as a vector. These are easy with what we have seen until now:

```
B <- select(A, 2)     ##### a table with just the second column
C <- unlist(A[1, ])  ##### the first row as a vector (coerced if necessary)
```

Section 23.5: Setting keys in data.table

Yes, you need to `SETKEY` pre 1.9.6

In the past (pre 1.9.6), your `data.table` was sped up by setting columns as keys to the table, particularly for large tables. [See [intro vignette page 5](#) of September 2015 version, where speed of search was 544 times better.] You may find older code making use of this setting keys with '`setkey`' or setting a '`key=`' column when setting up the table.

```
library(data.table)
DT <- data.table(
  x = letters[1:5],
  y = 5:1,
  z = (1:5) > 3
)

#> DT
#   x y     z
#1: a 5 FALSE
#2: b 4 FALSE
#3: c 3 FALSE
#4: d 2 TRUE
#5: e 1 TRUE
```

Set your key with the `setkey` command. You can have a key with multiple columns.

```
setkey(DT, y)
```

Check your table's key in `tables()`

```
tables()

> tables()
    NAME NROW NCOL MB COLS KEY
[1,] DT      5     3  1 x,y,z y
Total: 1MB
```

Note this will re-sort your data.

```
#> DT
#   x y     z
#1: e 1 TRUE
#2: d 2 TRUE
#3: c 3 FALSE
#4: b 4 FALSE
#5: a 5 FALSE
```

Now it is unnecessary

Prior to v1.9.6 you had to have set a key for certain operations especially joining tables. The developers of `data.table` have sped up and introduced a "`on=`" feature that can replace the dependency on keys. See [SO answer here for a detailed discussion](#).

In Jan 2017, the developers have written a [vignette around secondary indices](#) which explains the "on" syntax and allows for other columns to be identified for fast indexing.

Creating secondary indices?

In a manner similar to key, you can `setindex(DT, key.col)` or `setindexv(DT, "key.col.string")`, where DT is your data.table. Remove all indices with `setindex(DT, NULL)`.

See your secondary indices with `indices(DT)`.

Why secondary indices?

This **does not sort** the table (unlike key), but does allow for quick indexing using the "on" syntax. Note there can be only one key, but you can use multiple secondary indices, which saves having to rekey and resort the table. This will speed up your subsetting when changing the columns you want to subset on.

Recall, in example above y was the key for table DT:

```
DT
# x y      z
# 1: e 1  TRUE
# 2: d 2  TRUE
# 3: c 3 FALSE
# 4: b 4 FALSE
# 5: a 5 FALSE

# Let us set x as index
setindex(DT, x)

# Use indices to see what has been set
indices(DT)
# [1] "x"

# fast subset using index and not keyed column
DT["c", on ="x"]
#x y      z
#1: c 3 FALSE

# old way would have been rekeying DT from y to x, doing subset and
# perhaps keying back to y (now we save two sorts)
# This is a toy example above but would have been more valuable with big data sets
```


Chapter 25: Bar Chart

The purpose of the bar plot is to display the frequencies (or proportions) of levels of a factor variable. For example, a bar plot is used to pictorially display the frequencies (or proportions) of individuals in various socio-economic (factor) groups(levels-high, middle, low). Such a plot will help to provide a visual comparison among the various factor levels.

Section 25.1: barplot() function

In barplot, factor-levels are placed on the x-axis and frequencies (or proportions) of various factor-levels are considered on the y-axis. For each factor-level one bar of uniform width with heights being proportional to factor level frequency (or proportion) is constructed.

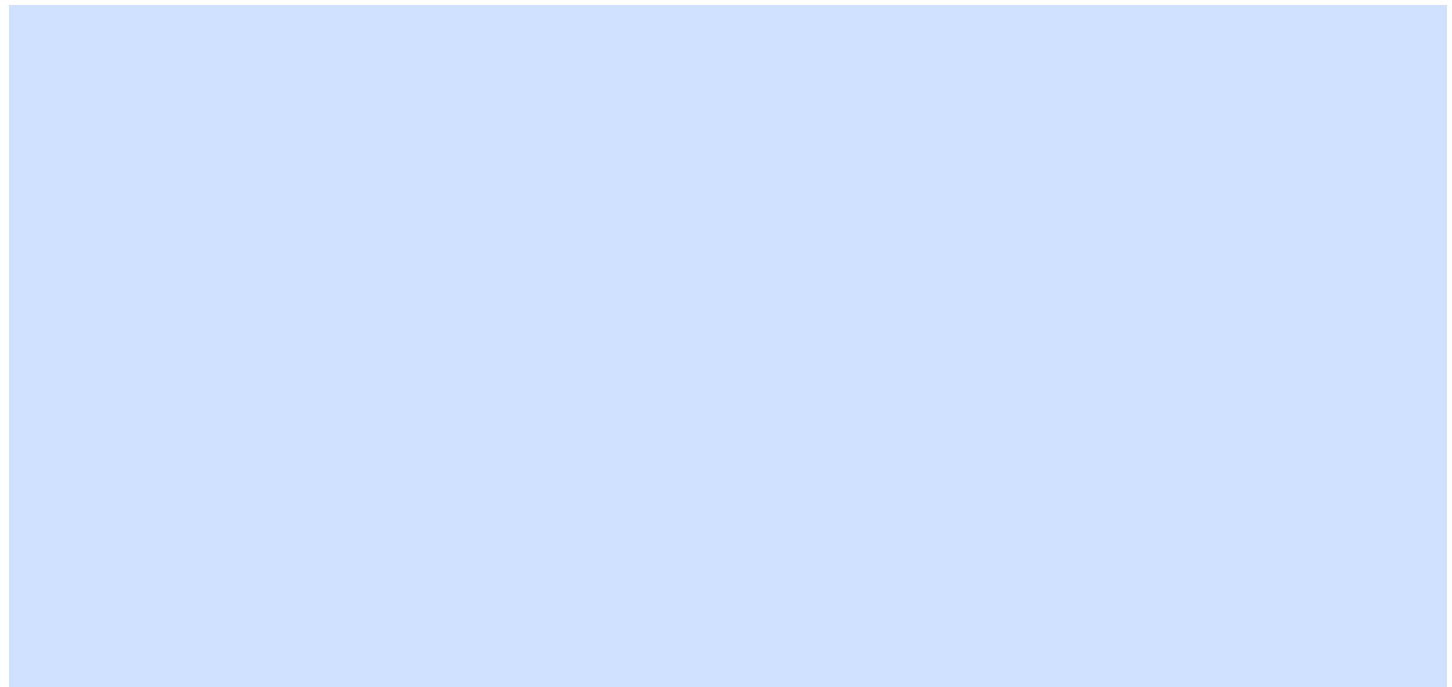
The **barplot()** function is in the graphics package of the R's System Library. The **barplot()** function must be supplied at least one argument. The R help calls this as heights, which must be either vector or a matrix. If it is vector, its members are the various factor-levels.

To illustrate **barplot()**, consider the following data preparation:

```
> grades<-c("A+","A-","B+","B","C")
> Marks<-sample(grades,40,replace=T,prob=c(.2,.3,.25,.15,.1))
> Marks
[1] "A+" "A-" "B+" "A-" "A+" "B" "A+" "B+" "A-" "B" "A+" "A-"
[13] "A-" "B+" "A-" "A-" "A-" "A+" "A-" "A+" "A+" "C" "C"
[25] "B" "C" "B+" "C" "B+" "B+" "B+" "A+" "B+" "A-" "A+" "A-"
[37] "A-" "B" "C" "A+
>
```

A bar chart of the Marks vector is obtained from

```
> barplot(table(Marks),main="Mid-Marks in Algorithms")
```



Notice that, the **barplot()** function places the factor levels on the x-axis in the lexicographical **order** of the levels. Using the parameter names **.arg**, the bars in plot can be placed in the order as stated in the vector, *grades*.

```
# plot to the desired horizontal axis labels  
> barplot(table(Marks),names.arg=grades ,main="Mid-Marks in Algorithms")
```

Colored bars can be drawn using the `col=` parameter.

```
> barplot(table(Marks),names.arg=grades,col = c("lightblue",  
"lightcyan", "lavender", "mistyrose", "cornsilk"),  
main="Mid-Marks in Algorithms")
```

A bar chart with *horizontal bars* can be obtained as follows:

```
> barplot(table(Marks),names.arg=grades,horiz=TRUE,col = c("lightblue",  
"lightcyan", "lavender", "mistyrose", "cornsilk"),
```

```
main="Mid-Marks in Algorithms")
```



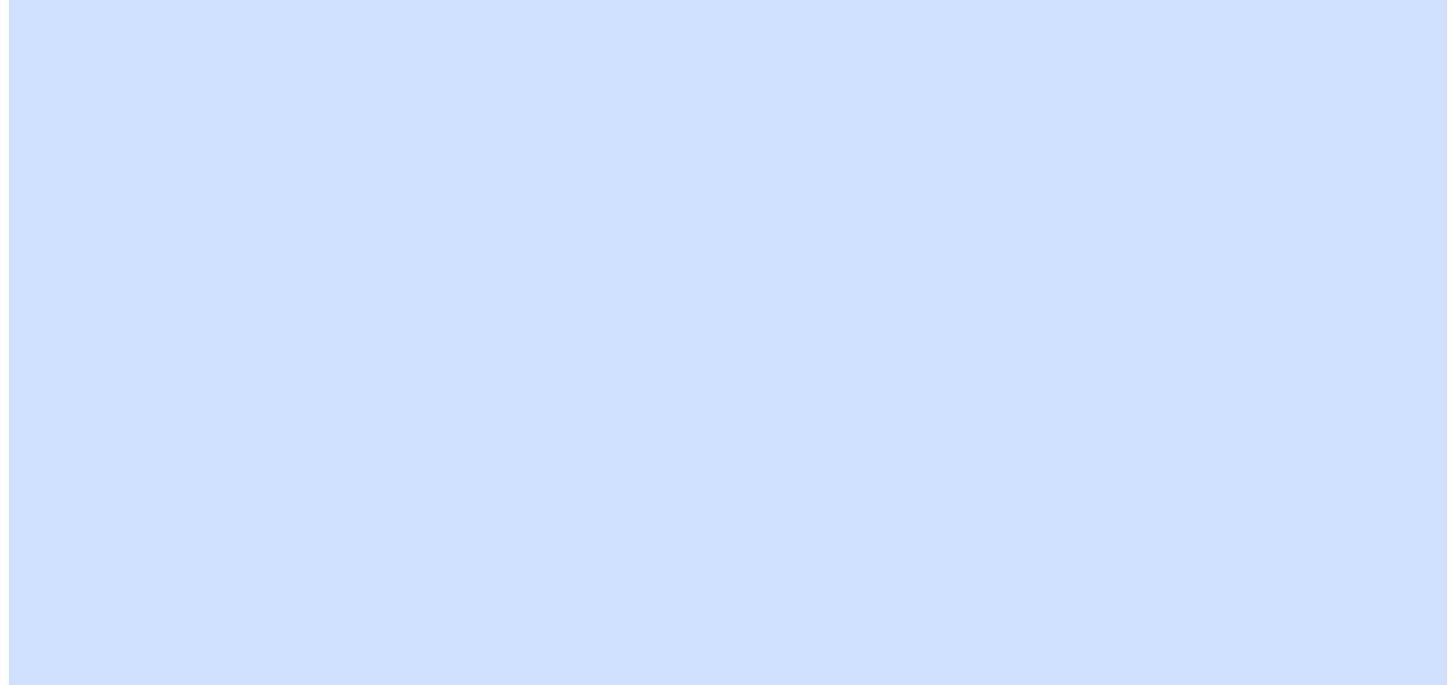
A bar chart with *proportions* on the y-axis can be obtained as follows:

```
> barplot(prop.table(table(Marks)), names.arg=grades, col = c("lightblue",
  "lightcyan", "lavender", "mistyrose", "cornsilk"),
  main="Mid-Marks in Algorithms")
```



The sizes of the factor-level names on the x-axis can be increased using `cex.names` parameter.

```
> barplot(prop.table(table(Marks)), names.arg=grades, col = c("lightblue",
  "lightcyan", "lavender", "mistyrose", "cornsilk"),
  main="Mid-Marks in Algorithms", cex.names=2)
```



The heights parameter of the `barplot()` could be a matrix. For example it could be matrix, where the columns are the various subjects taken in a course, the rows could be the labels of the grades. Consider the following matrix:

```
> gradTab
  Algorithms Operating Systems Discrete Math
A-          13            10            7
A+          10             7            2
B            4             2            14
B+           8            19            12
C            5             2            5
```

To draw a stacked bar, simply use the command:

```
> barplot(gradTab, col = c("lightblue", "lightcyan",
  "lavender", "mistyrose", "cornsilk"), legend.text = grades,
  main="Mid-Marks in Algorithms")
```



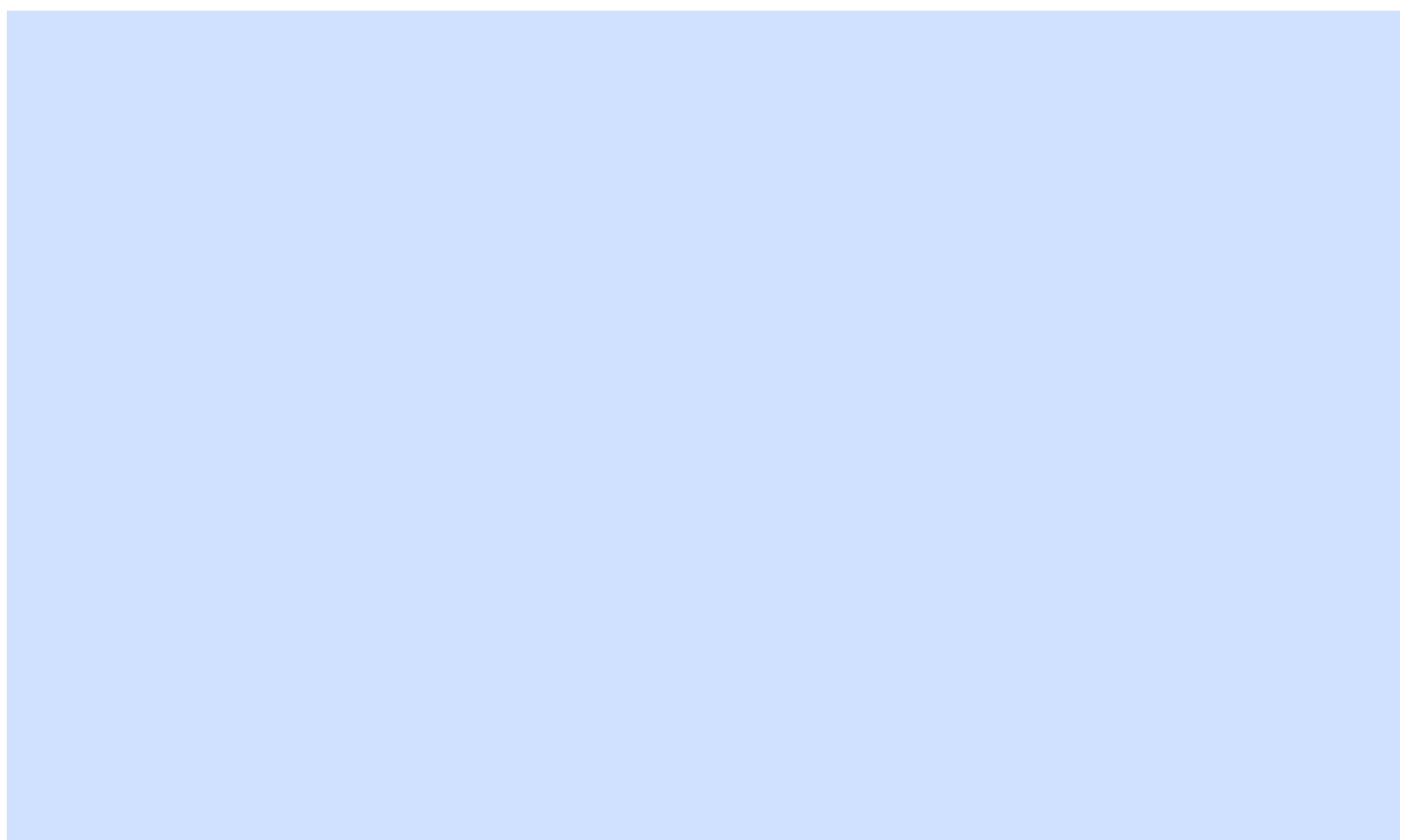
To draw a juxtaposed bars, use the `besides` parameter, as given under:

```
> barplot(gradTab,beside = T,col = c("lightblue","lightcyan",
  "lavender", "mistyrose", "cornsilk"),legend.text = grades,
  main="Mid-Marks in Algorithms")
```



A horizontal bar chart can be obtained using `horiz=T` parameter:

```
> barplot(gradTab,beside = T,horiz=T,col = c("lightblue","lightcyan",
  "lavender", "mistyrose", "cornsilk"),legend.text = grades,
  cex.names=.75,main="Mid-Marks in Algorithms")
```



Chapter 26: Base Plotting

Parameter	Details
x	x-axis variable. May supply either <code>data\$variablex</code> or <code>data[, x]</code>
y	y-axis variable. May supply either <code>data\$variabley</code> or <code>data[, y]</code>
main	Main title of plot
sub	Optional subtitle of plot
xlab	Label for x-axis
ylab	Label for y-axis
pch	Integer or character indicating plotting symbol
col	Integer or string indicating color
type	Type of plot. "p" for points, "l" for lines, "b" for both, "c" for the lines part alone of "b", "o" for both 'overplotted', "h" for 'histogram'-like (or 'high-density') vertical lines, "s" for stair steps, "S" for other steps, "n" for no plotting

Section 26.1: Density plot

A very useful and logical follow-up to histograms would be to plot the smoothed density function of a random variable. A basic plot produced by the command

```
plot(density(rnorm(100)), main="Normal density", xlab="x")
```

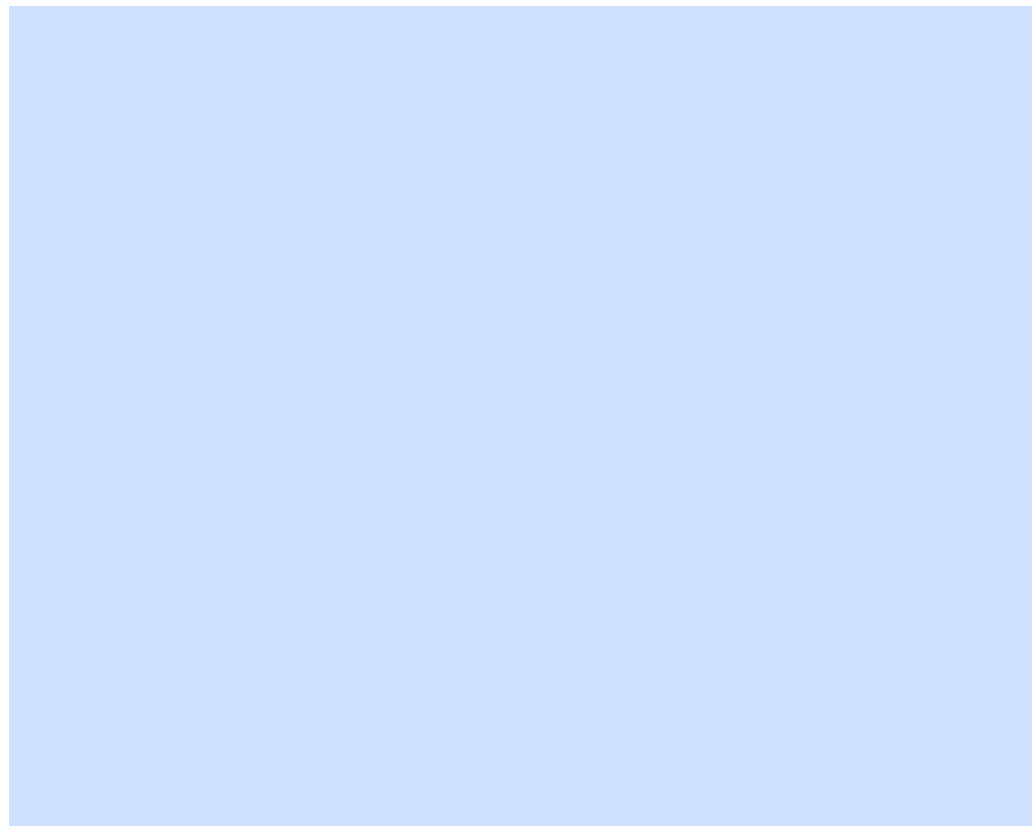
would look like



You can overlay a histogram and a density curve with

```
x=rnorm(100)
hist(x, prob=TRUE, main="Normal density + histogram")
lines(density(x), lty="dotted", col="red")
```

which gives



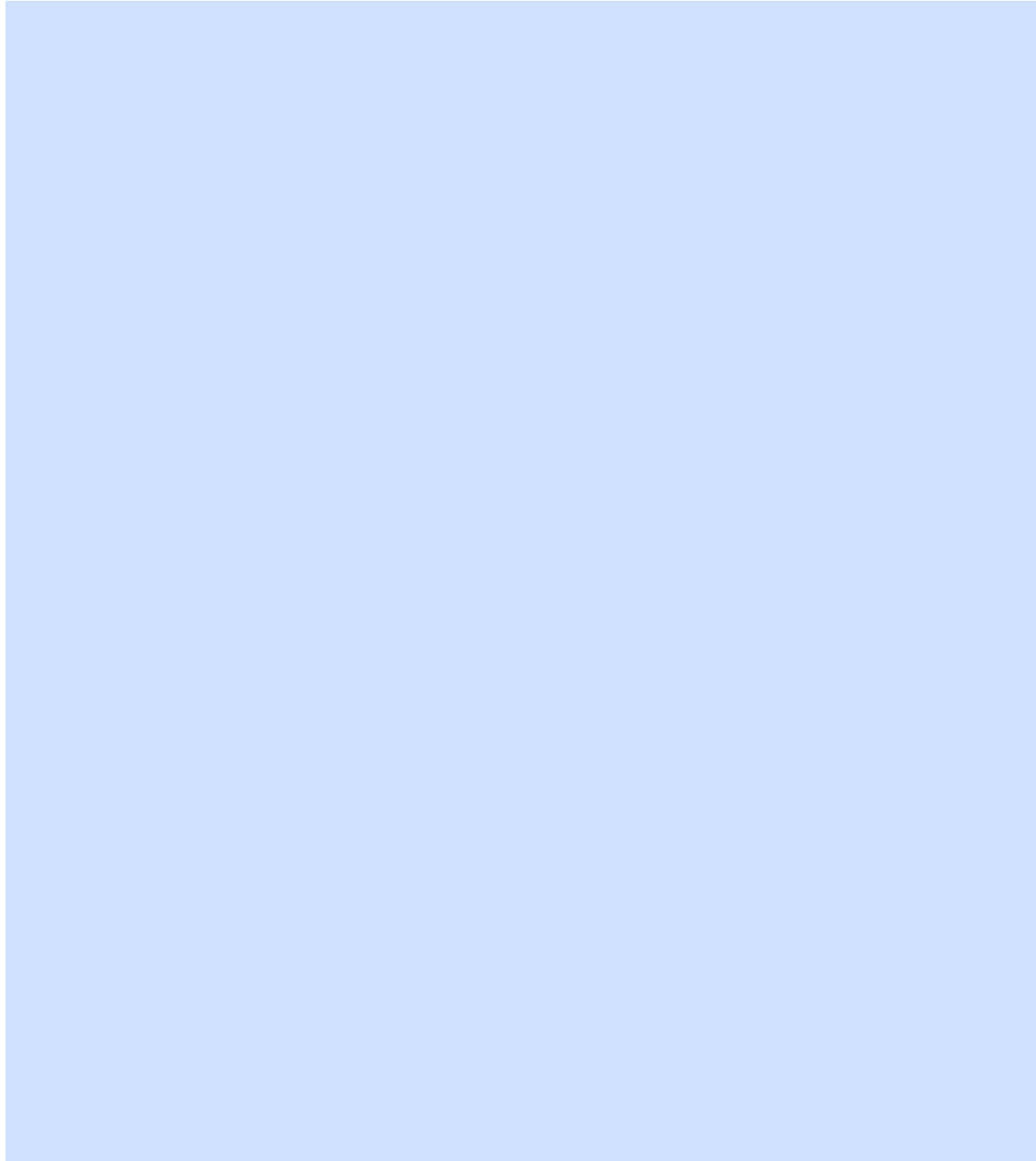
Section 26.2: Combining Plots

It's often useful to combine multiple plot types in one graph (for example a Barplot next to a Scatterplot.) R makes this easy with the help of the functions `par()` and `layout()`.

`par()`

`par` uses the arguments `mfrow` or `mfcol` to create a matrix of `nrows` and `ncols` `c(nrows, ncols)` which will serve as a grid for your plots. The following example shows how to combine four plots in one graph:

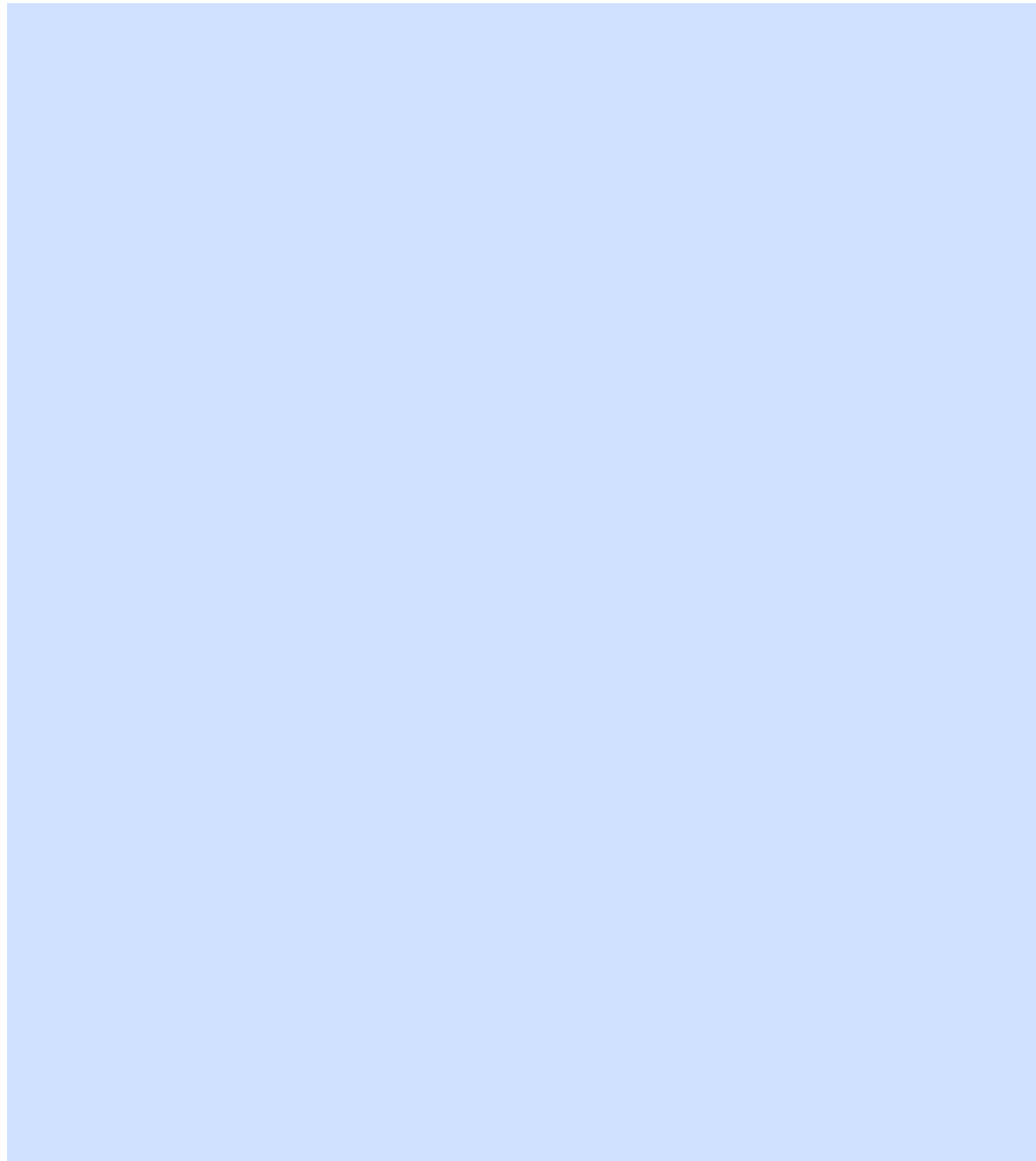
```
par(mfrow=c(2,2))
plot(cars, main="Speed vs. Distance")
hist(cars$speed, main="Histogram of Speed")
boxplot(cars$dist, main="Boxplot of Distance")
boxplot(cars$speed, main="Boxplot of Speed")
```



layout()

The `layout()` is more flexible and allows you to specify the location and the extent of each plot within the final combined graph. This function expects a matrix object as an input:

```
layout(matrix(c(1,1,2,3), 2,2, byrow=T))
hist(cars$speed, main="Histogram of Speed")
boxplot(cars$dist, main="Boxplot of Distance")
boxplot(cars$speed, main="Boxplot of Speed")
```



Section 26.3: Getting Started with R_Plots

- **Scatterplot**

You have two vectors and you want to plot them.

```
x_values <- rnorm(n = 20 , mean = 5 , sd = 8) #20 values generated from Normal(5,8)
y_values <- rbeta(n = 20 , shape1 = 500 , shape2 = 10) #20 values generated from Beta(500,10)
```

If you want to make a plot which has the `y_values` in vertical axis and the `x_values` in horizontal axis, you can use the following commands:

```
plot(x = x_values, y = y_values, type = "p") #standard scatter-plot
plot(x = x_values, y = y_values, type = "l") # plot with lines
```

```
plot(x = x_values, y = y_values, type = "n") # empty plot
```

You can type `?plot()` in the console to read about more options.

- **Boxplot**

You have some variables and you want to examine their Distributions

```
#boxplot is an easy way to see if we have some outliers in the data.
```

```
z<- rbeta(20 , 500 , 10) #generating values from beta distribution  
z[c(19 , 20)] <- c(0.97 , 1.05) # replace the two last values with outliers  
boxplot(z) # the two points are the outliers of variable z.
```

- **Histograms**

Easy way to draw histograms

```
hist(x = x_values) # Histogram for x vector  
hist(x = x_values, breaks = 3) #use breaks to set the numbers of bars you want
```

- **Pie_charts**

If you want to visualize the frequencies of a variable just draw pie

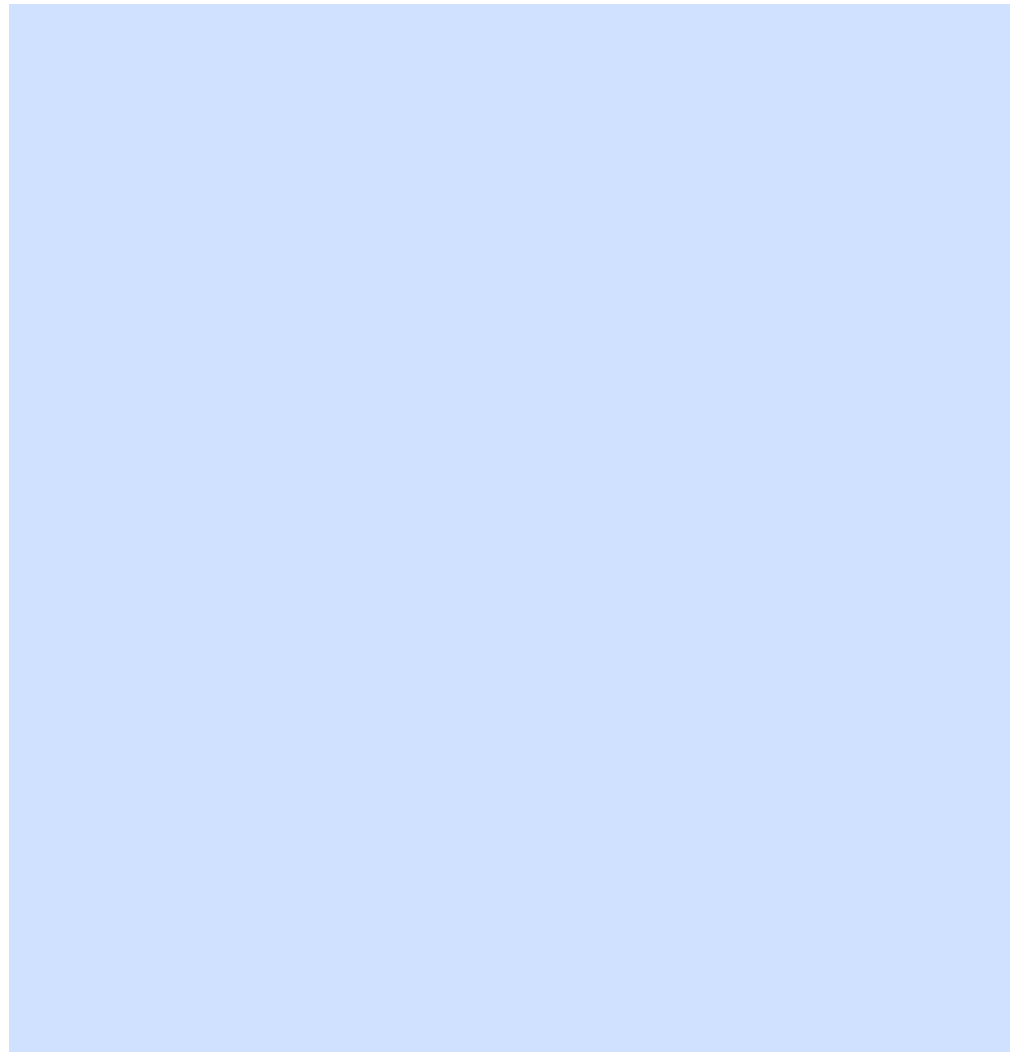
First we have to generate data with frequencies, for example :

```
P <- c(rep('A' , 3) , rep('B' , 10) , rep('C' , 7) )  
t <- table(P) # this is a frequency matrix of variable P  
pie(t) # And this is a visual version of the matrix above
```

Section 26.4: Basic Plot

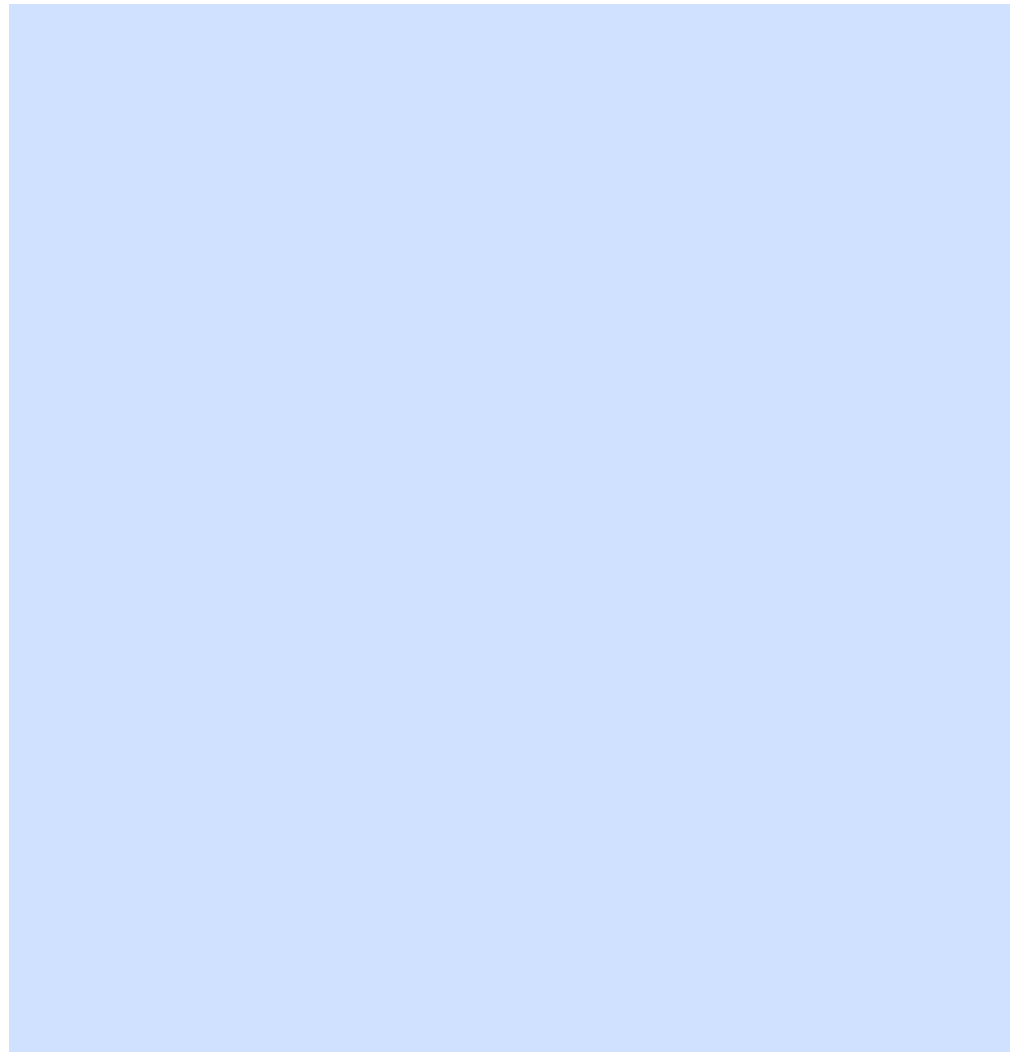
A basic plot is created by calling `plot()`. Here we use the built-in `cars` data frame that contains the speed of cars and the distances taken to stop in the 1920s. (To find out more about the dataset, use `help(cars)`).

```
plot(x = cars$speed, y = cars$dist, pch = 1, col = 1,  
     main = "Distance vs Speed of Cars",  
     xlab = "Speed", ylab = "Distance")
```



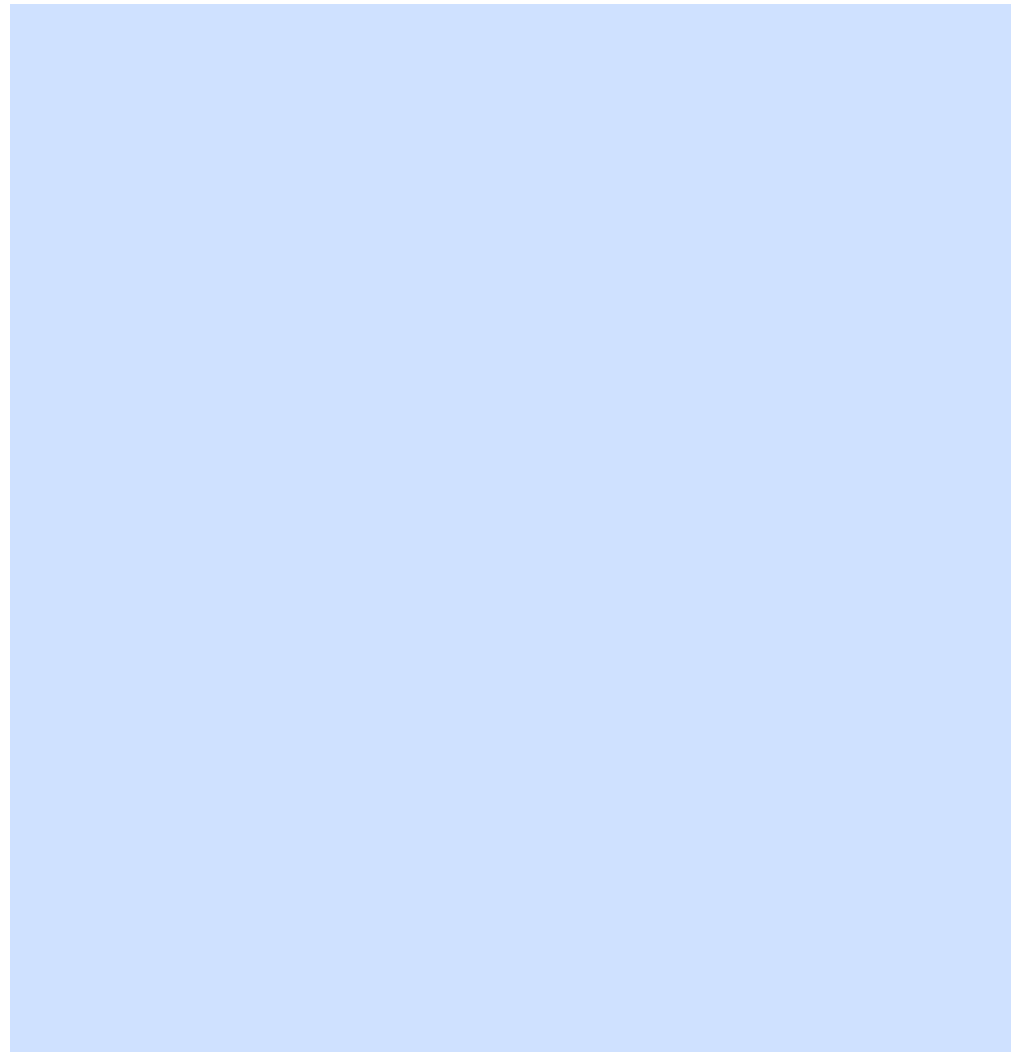
We can use many other variations in the code to get the same result. We can also change the parameters to obtain different results.

```
with(cars, plot(dist~speed, pch = 2, col = 3,
  main = "Distance to stop vs Speed of Cars",
  xlab = "Speed", ylab = "Distance"))
```



Additional features can be added to this plot by calling `points()`, `text()`, `mtext()`, `lines()`, `grid()`, etc.

```
plot(dist~speed, pch = "*", col = "magenta", data=cars,
     main = "Distance to stop vs Speed of Cars",
     xlab = "Speed", ylab = "Distance")
mtext("In the 1920s.")
grid(col="lightblue")
```



Section 26.5: Histograms

Histograms allow for a pseudo-plot of the underlying distribution of the data.

```
hist(ldeaths)
```

```
hist(ldeaths, breaks = 20, freq = F, col = 3)
```



Section 26.6: Matplot

`matplot` is useful for quickly plotting multiple sets of observations from the same object, particularly from a matrix, on the same graph.

Here is an example of a matrix containing four sets of random draws, each with a different mean.

```
xmat <- cbind(rnorm(100, -3), rnorm(100, -1), rnorm(100, 1), rnorm(100, 3))
head(xmat)
#           [,1]      [,2]      [,3]      [,4]
# [1,] -3.072793 -2.53111494  0.6168063 3.780465
# [2,] -3.702545 -1.42789347 -0.2197196 2.478416
# [3,] -2.890698 -1.88476126  1.9586467 5.268474
# [4,] -3.431133 -2.02626870  1.1153643 3.170689
# [5,] -4.532925  0.02164187  0.9783948 3.162121
# [6,] -2.169391 -1.42699116  0.3214854 4.480305
```

One way to plot all of these observations on the same graph is to do one `plot` call followed by three more `points` or `lines` calls.

```
plot(xmat[,1], type = 'l')
lines(xmat[,2], col = 'red')
lines(xmat[,3], col = 'green')
lines(xmat[,4], col = 'blue')
```



However, this is both tedious, and causes problems because, among other things, by default the axis limits are fixed by `plot` to fit only the first column.

Much more convenient in this situation is to use the `matplot` function, which only requires one call and automatically takes care of axis limits *and* changing the aesthetics for each column to make them distinguishable.

```
matplot(xmat, type = 'l')
```



Note that, by default, `matplot` varies both color (`col`) and linetype (`lty`) because this increases the number of possible combinations before they get repeated. However, any (or both) of these aesthetics can be fixed to a single value...

```
matplot(xmat, type = 'l', col = 'black')
```



...or a custom vector (which will recycle to the number of columns, following standard R vector recycling rules).

```
matplot(xmat, type = 'l', col = c('red', 'green', 'blue', 'orange'))
```



Standard graphical parameters, including `main`, `xlab`, `xmin`, work exactly the same way as for `plot`. For more on those, see `?par`.

Like `plot`, if given only one object, `matplot` assumes it's the y variable and uses the indices for x. However, x and y can be specified explicitly.

```
matplot(x = seq(0, 10, length.out = 100), y = xmat, type='l')
```



In fact, both x and y can be matrices.

```
xes <- cbind(seq(0, 10, length.out = 100),
             seq(2.5, 12.5, length.out = 100),
             seq(5, 15, length.out = 100),
             seq(7.5, 17.5, length.out = 100))
matplot(x = xes, y = xmat, type = 'l')
```



Section 26.7: Empirical Cumulative Distribution Function

A very useful and logical follow-up to histograms and density plots would be the Empirical Cumulative Distribution Function. We can use the function `ecdf()` for this purpose. A basic plot produced by the command

```
plot(ecdf(rnorm(100)), main="Cumulative distribution", xlab="x")
```

would look like



Chapter 27: boxplot

Parameters	Details (source R Documentation)
formula	a formula, such as $y \sim grp$, where y is a numeric vector of data values to be split into groups according to the grouping variable grp (usually a factor).
data	a <code>data.frame</code> (or list) from which the variables in <code>formula</code> should be taken.
subset	an optional vector specifying a subset of observations to be used for plotting.
na.action	a function which indicates what should happen when the data contain NAs. The default is to ignore missing values in either the response or the group.
boxwex	a scale factor to be applied to all boxes. When there are only a few groups, the appearance of the plot can be improved by making the boxes narrower.
plot	if <code>TRUE</code> (the default) then a boxplot is produced. If not, the summaries which the boxplots are based on are returned.
col	if <code>col</code> is non-null it is assumed to contain colors to be used to colour the bodies of the box plots. By default they are in the background colour.

Section 27.1: Create a box-and-whisker plot with `boxplot()` {graphics}

This example use the default `boxplot()` function and the `iris` data frame.

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

Simple boxplot (Sepal.Length)

Create a box-and-whisker graph of a numerical variable

```
boxplot(iris[, 1], xlab="Sepal.Length", ylab="Length(in centimeters)",
        main="Summary Charateristics of Sepal.Length(Iris Data)")
```

Boxplot of sepal length grouped by species

Create a boxplot of a numerical variable grouped by a categorical variable

```
boxplot(Sepal.Length~Species, data = iris)
```

Bring order

To change order of the box in the plot you have to change the order of the categorical variable's levels.

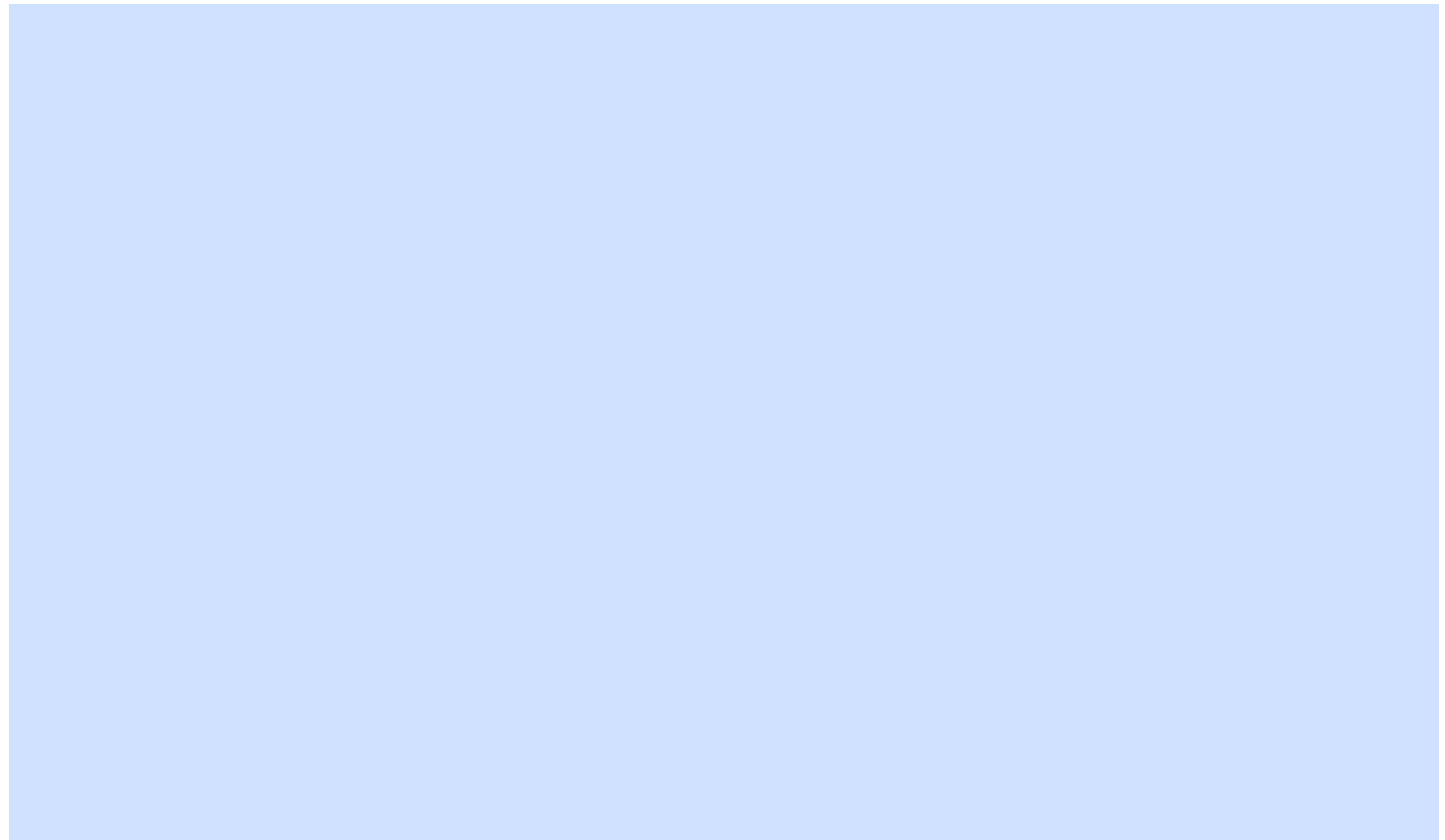
For example if we want to have the order virginica - versicolor - setosa

```
newSpeciesOrder <- factor(iris$Species, levels=c("virginica", "versicolor", "setosa"))
boxplot(Sepal.Length~newSpeciesOrder, data = iris)
```

Change groups names

If you want to specify a better name to your groups you can use the `Names` parameter. It take a vector of the size of the levels of categorical variable

```
boxplot(Sepal.Length~newSpeciesOrder, data = iris, names= c("name1", "name2", "name3"))
```



Small improvements

Color

col : add a vector of the size of the levels of categorical variable

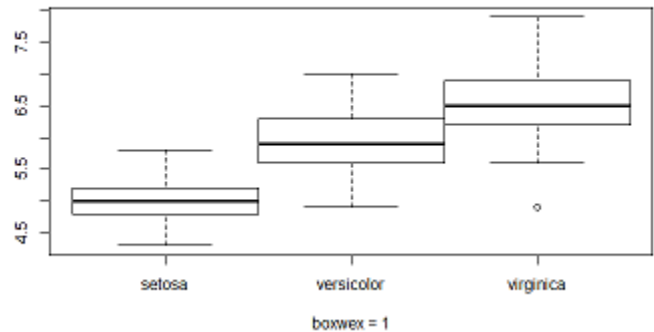
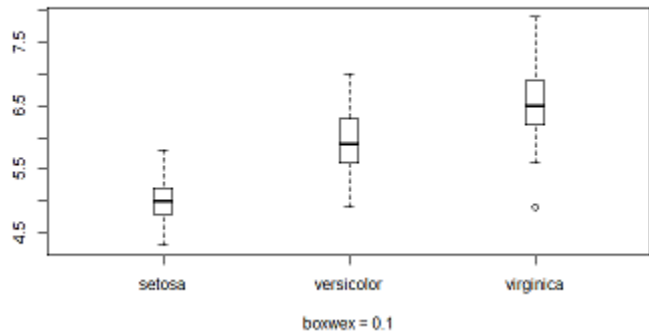
```
boxplot(Sepal.Length~Species, data = iris, col=c("green", "yellow", "orange"))
```

Proximity of the box

boxwex: set the margin between boxes.

Left `boxplot(Sepal.Length~Species, data = iris, boxwex = 0.1)`

Right `boxplot(Sepal.Length~Species, data = iris, boxwex = 1)`



See the summaries which the boxplots are based `plot=FALSE`

To see a summary you have to put the parameter `plot` to FALSE.

Various results are given

```
> boxplot(Sepal.Length~newSpeciesOrder, data = iris, plot=FALSE)
$stats #summary of the numerical variable for the 3 groups
 [,1] [,2] [,3]
[1,] 5.6 4.9 4.3 # extreme value
[2,] 6.2 5.6 4.8 # first quartile limit
[3,] 6.5 5.9 5.0 # median limit
[4,] 6.9 6.3 5.2 # third quartile limit
[5,] 7.9 7.0 5.8 # extreme value

$n #number of observations in each groups
[1] 50 50 50

$conf #extreme value of the notches
 [,1] [,2] [,3]
[1,] 6.343588 5.743588 4.910622
[2,] 6.656412 6.056412 5.089378

$out #extreme value
[1] 4.9

$group #group in which are the extreme value
[1] 1

$names #groups names
[1] "virginica" "versicolor" "setosa"
```

Section 27.2: Additional boxplot style parameters

Box

- `boxlty` - box line type
- `boxlwd` - box line width
- `boxcol` - box line color
- `boxfill` - box fill colors

Median

- medlty - median line type ("blank" for no line)
- medlwd - median line width
- medcol - median line color
- medpch - median point (NA for no symbol)
- medcex - median point size
- medbg - median point background color

Whisker

- whisklty - whisker line type
- whisklwd - whisker line width
- whiskcol - whisker line color

Staple

- staplelty - staple line type
- staplelwd - staple line width
- staplecol - staple line color

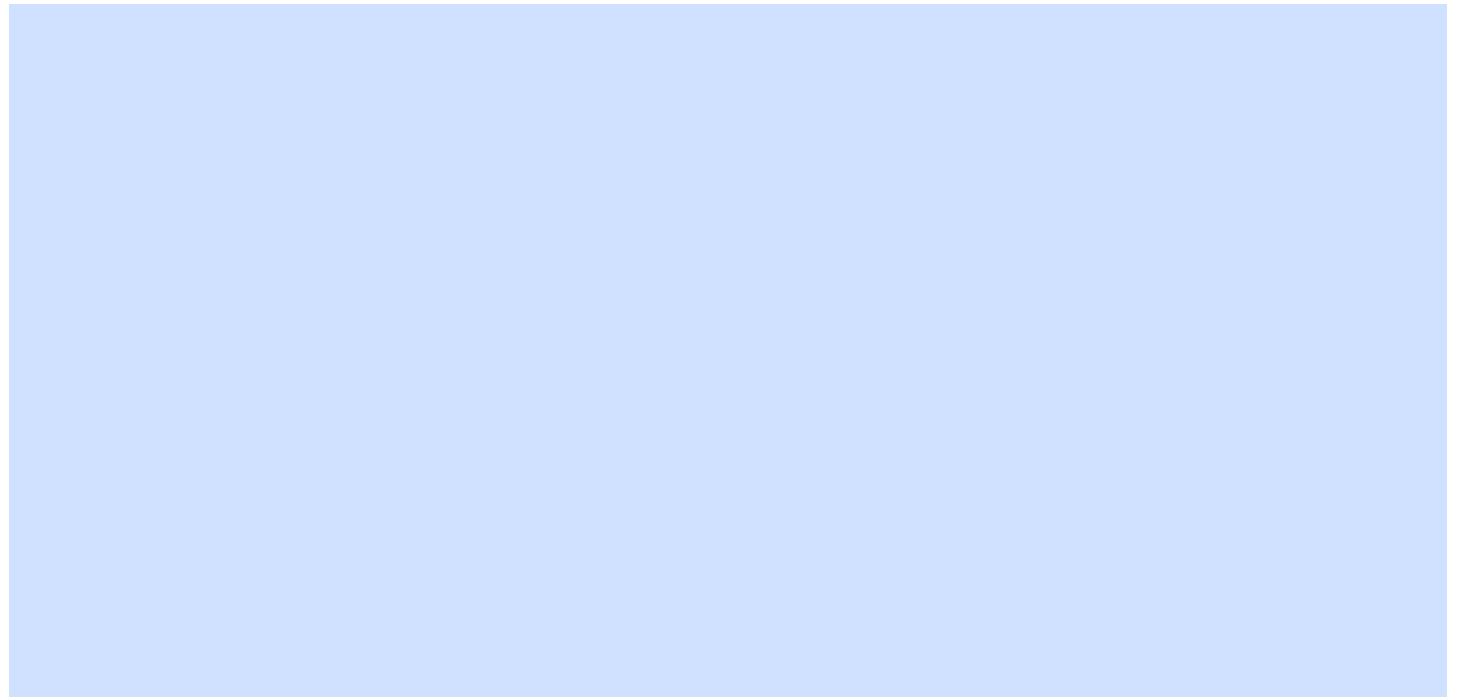
Outliers

- outlty - outlier line type ("blank" for no line)
- outlwd - outlier line width
- outcol - outlier line color
- outpch - outlier point type (NA for no symbol)
- outcex - outlier point size
- outbg - outlier point background color

Example

Default and heavily modified plots side by side

```
par(mfrow=c(1, 2))
# Default
boxplot(Sepal.Length ~ Species, data=iris)
# Modified
boxplot(Sepal.Length ~ Species, data=iris,
        boxlty=2, boxlwd=3, boxfill="cornflowerblue", boxcol="darkblue",
        medlty=2, medlwd=2, medcol="red", medpch=21, medcex=1, medbg="white",
        whisklty=2, whisklwd=3, whiskcol="darkblue",
        staplelty=2, staplelwd=2, staplecol="red",
        outlty=3, outlwd=3, outcol="grey", outpch=NA
    )
```



Chapter 28: ggplot2

Section 28.1: Displaying multiple plots

Display multiple plots in one image with the different facet functions. An advantage of this method is that all axes share the same scale across charts, making it easy to compare them at a glance. We'll use the mpg dataset included in ggplot2.

Wrap charts line by line (attempts to create a square layout):

```
ggplot(mpg, aes(x = displ, y = hwy)) +  
  geom_point() +  
  facet_wrap(~class)
```



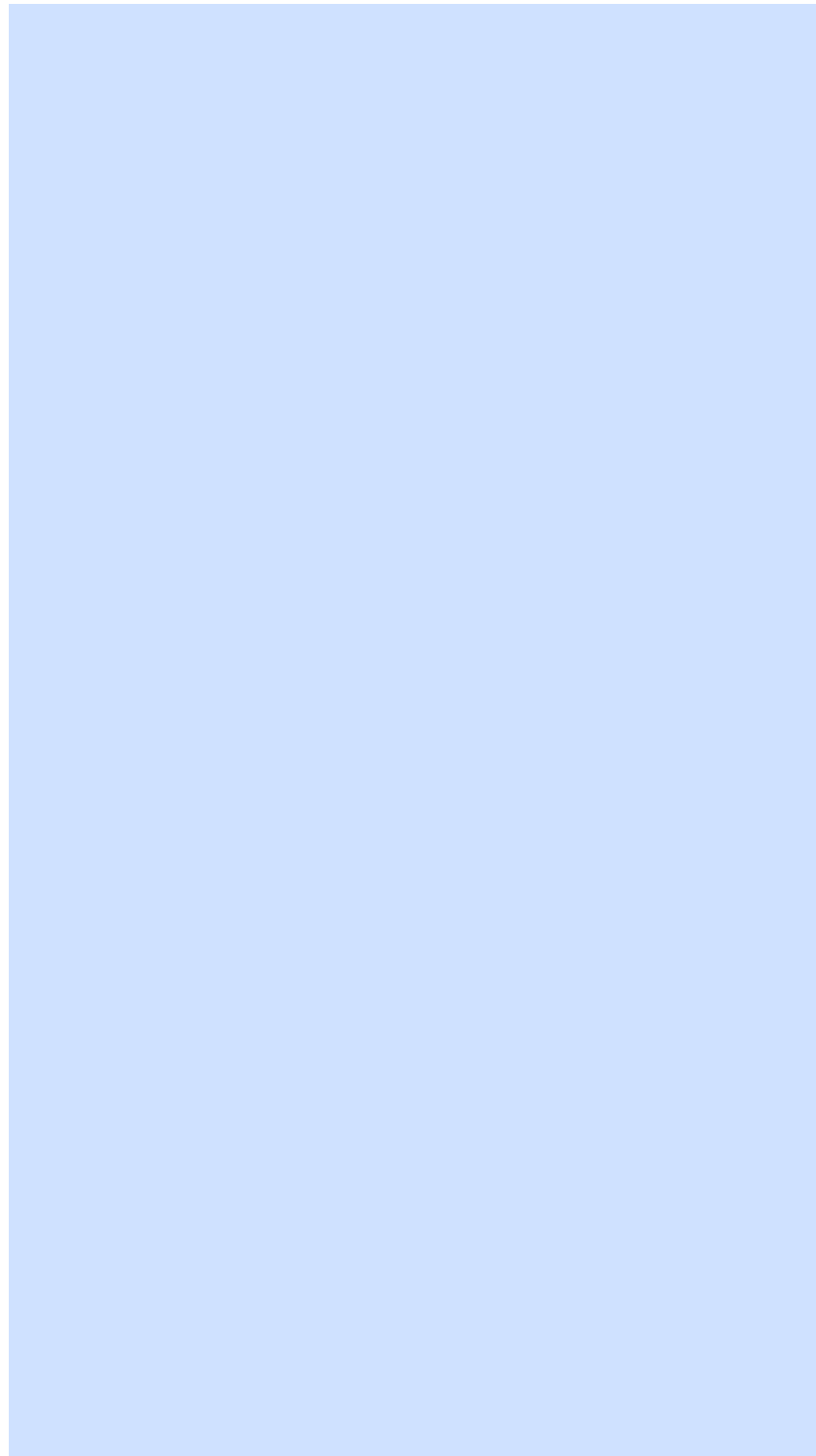
Display multiple charts on one row, multiple columns:

```
ggplot(mpg, aes(x = displ, y = hwy)) +
```

```
geom_point() +  
facet_grid(.~class)
```

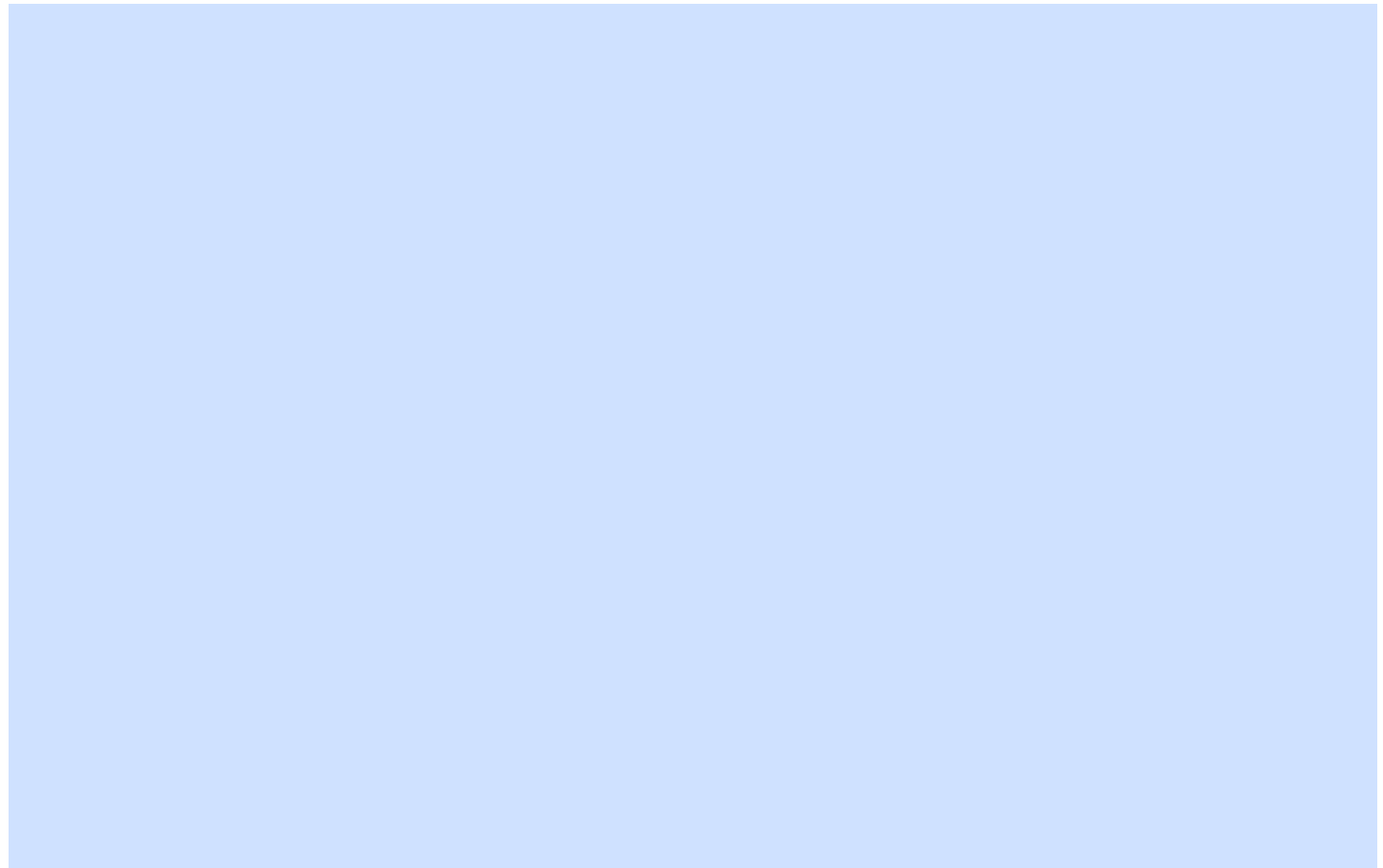
Display multiple charts on one column, multiple rows:

```
ggplot(mpg, aes(x = displ, y = hwy)) +  
geom_point() +  
facet_grid(class~.)
```



Display multiple charts in a grid by 2 variables:

```
ggplot(mpg, aes(x = displ, y = hwy)) +  
  geom_point() +  
  facet_grid(trans~class) # "row" parameter, then "column" parameter
```



Section 28.2: Prepare your data for plotting

ggplot2 works best with a long data frame. The following sample data which represents the prices for sweets on 20 different days, in a format described as wide, because each category has a column.

```
set.seed(47)
sweetsWide <- data.frame(date      = 1:20,
                           chocolate = runif(20, min = 2, max = 4),
                           iceCream  = runif(20, min = 0.5, max = 1),
                           candy     = runif(20, min = 1, max = 3))

head(sweetsWide)
##   date chocolate  iceCream   candy
## 1    1   3.953924 0.5890727 1.117311
## 2    2   2.747832 0.7783982 1.740851
## 3    3   3.523004 0.7578975 2.196754
## 4    4   3.644983 0.5667152 2.875028
## 5    5   3.147089 0.8446417 1.733543
## 6    6   3.382825 0.6900125 1.405674
```

To convert sweetsWide to long format for use with ggplot2, several useful functions from base R, and the packages reshape2, data.table and tidyr (in chronological order) can be used:

```
# reshape from base R
sweetsLong <- reshape(sweetsWide, idvar = 'date', direction = 'long',
                      varying = list(2:4), new.row.names = NULL, times = names(sweetsWide)[-1])

# melt from 'reshape2'
library(reshape2)
sweetsLong <- melt(sweetsWide, id.vars = 'date')
```

```
# melt from 'data.table'  
# which is an optimized & extended version of 'melt' from 'reshape2'  
library(data.table)  
sweetsLong <- melt(setDT(sweetsWide), id.vars = 'date')  
  
# gather from 'tidyR'  
library(tidyR)  
sweetsLong <- gather(sweetsWide, sweet, price, chocolate:candy)
```

The all give a similar result:

```
head(sweetsLong)  
##   date     sweet    price  
## 1     1 chocolate 3.953924  
## 2     2 chocolate 2.747832  
## 3     3 chocolate 3.523004  
## 4     4 chocolate 3.644983  
## 5     5 chocolate 3.147089  
## 6     6 chocolate 3.382825
```

See also Reshaping data between long and wide forms for details on converting data between *long* and *wide* format.

The resulting sweetsLong has one column of prices and one column describing the type of sweet. Now plotting is much simpler:

```
library(ggplot2)  
ggplot(sweetsLong, aes(x = date, y = price, colour = sweet)) + geom_line()
```

Section 28.3: Add horizontal and vertical lines to plot

Add one common horizontal line for all categorical variables

```
# sample data
df <- data.frame(x=c('A', 'B'), y = c(3, 4))

p1 <- ggplot(df, aes(x=x, y=y))
  + geom_bar(position = "dodge", stat = 'identity')
  + theme_bw()

p1 + geom_hline(aes(yintercept=5), colour="#990000", linetype="dashed")
```



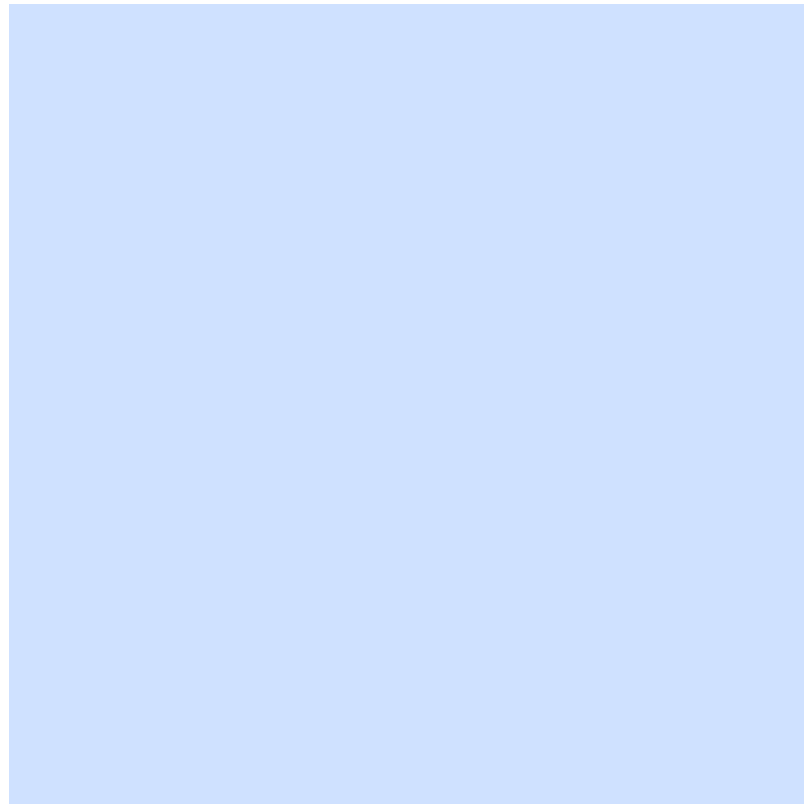
Add one horizontal line for each categorical variable

```
# sample data
df <- data.frame(x=c('A', 'B'), y = c(3, 4))

# add horizontal levels for drawing lines
df$hval <- df$y + 2

p1 <- ggplot(df, aes(x=x, y=y))
  + geom_bar(position = "dodge", stat = 'identity')
  + theme_bw()

p1 + geom_errorbar(aes(y=hval, ymax=hval, ymin=hval), colour="#990000", width=0.75)
```

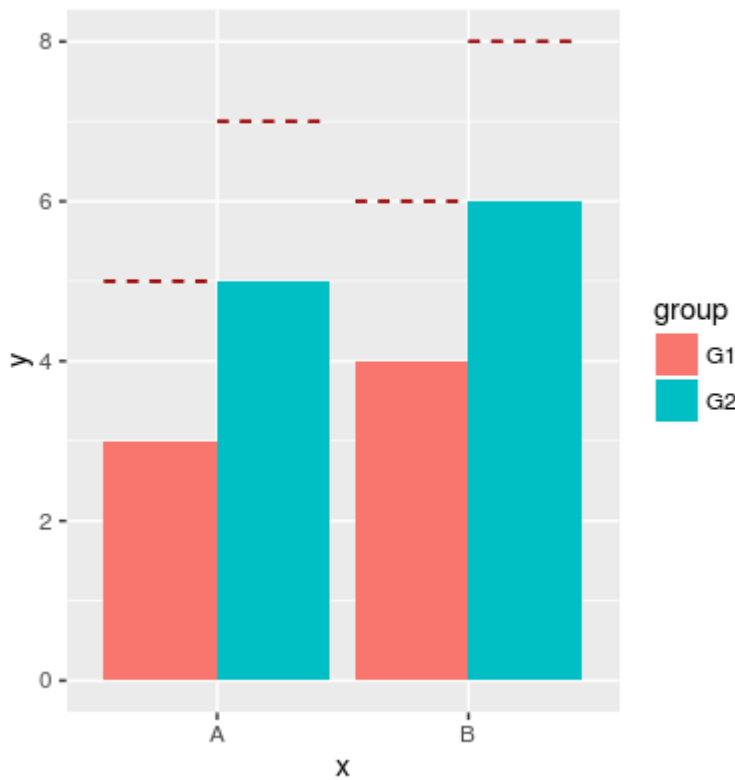


Add horizontal line over grouped bars

```
# sample data
df <- data.frame(x = rep(c('A', 'B'), times=2),
                  group = rep(c('G1', 'G2'), each=2),
                  y = c(3, 4, 5, 6),
                  hval = c(5, 6, 7, 8))

p1 <- ggplot(df, aes(x=x, y=y, fill=group))
  + geom_bar(position="dodge", stat="identity")

p1 + geom_errorbar(aes(y=hval, ymax=hval, ymin=hval),
                    colour="#990000",
                    position = "dodge",
                    linetype = "dashed")
```

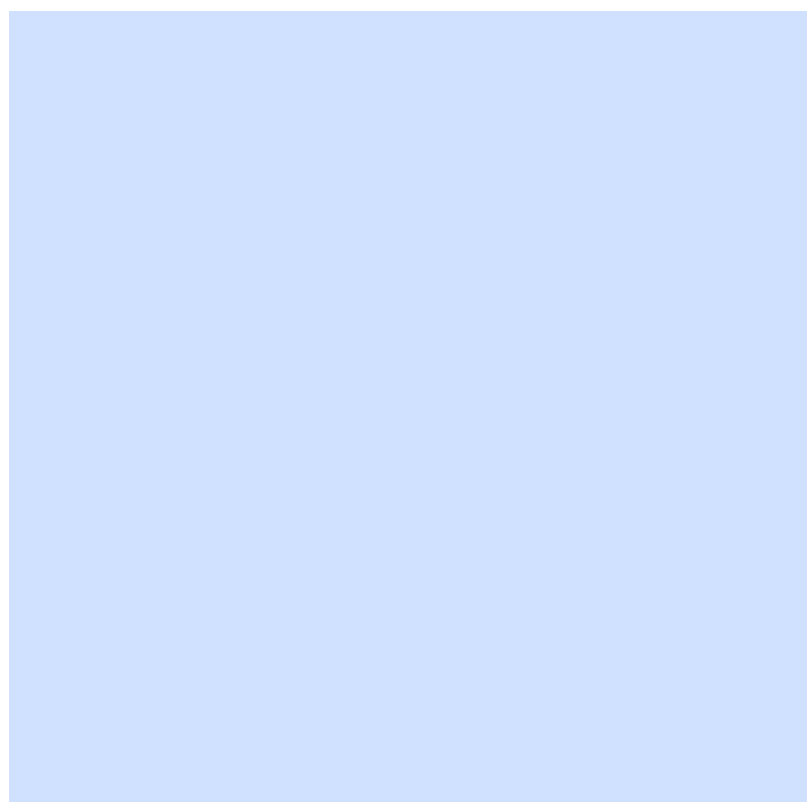


Add vertical line

```
# sample data
df <- data.frame(group=rep(c('A', 'B'), each=20),
                  x = rnorm(40, 5, 2),
                  y = rnorm(40, 10, 2))

p1 <- ggplot(df, aes(x=x, y=y, colour=group)) + geom_point()

p1 + geom_vline(aes(xintercept=5), color="#990000", linetype="dashed")
```



Section 28.4: Scatter Plots

We plot a simple scatter plot using the builtin iris data set as follows:

```
library(ggplot2)
ggplot(iris, aes(x = Petal.Width, y = Petal.Length, color = Species)) +
  geom_point()
```

This gives:



Section 28.5: Produce basic plots with qplot

qplot is intended to be similar to base R `plot()` function, trying to always plot out your data without requiring too much specifications.

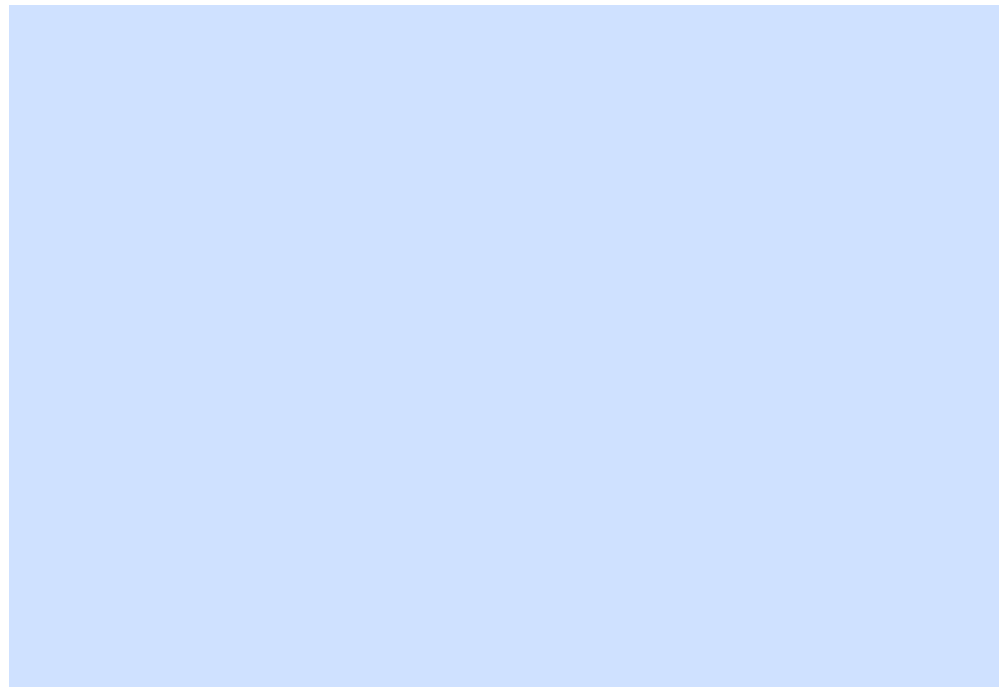
basic qplot

```
qplot(x = disp, y = mpg, data = mtcars)
```



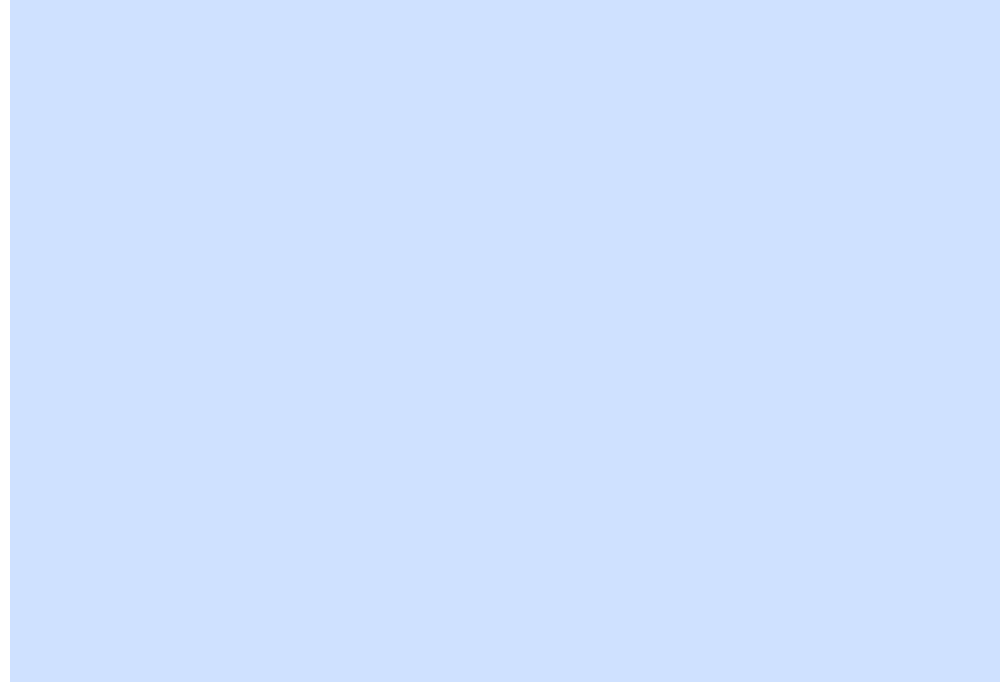
adding colors

```
qplot(x = disp, y = mpg, colour = cyl, data = mtcars)
```



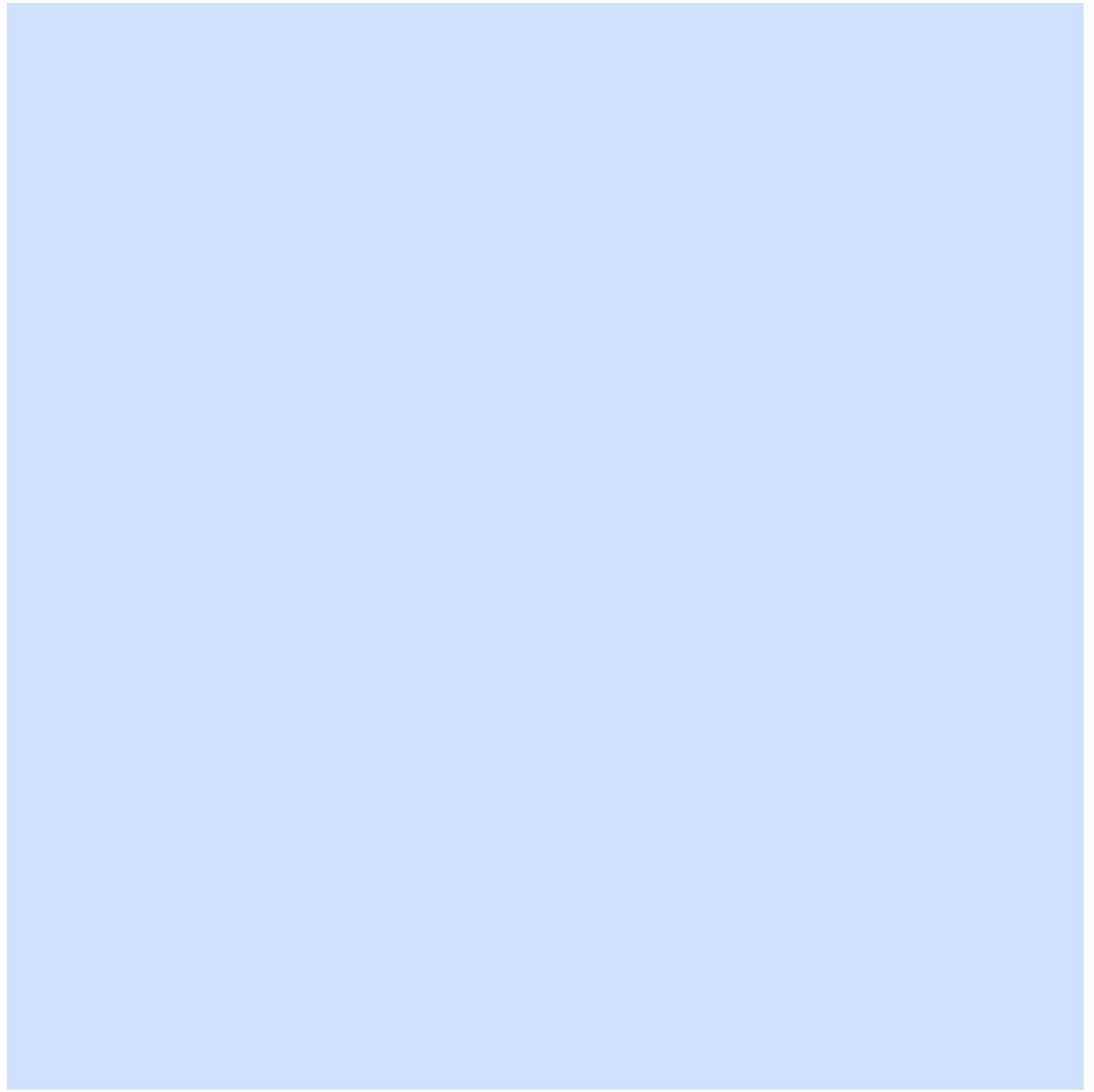
adding a smoother

```
qplot(x = disp, y = mpg, geom = c("point", "smooth"), data = mtcars)
```



Section 28.6: Vertical and Horizontal Bar Chart

```
ggplot(data = diamonds, aes(x = cut, fill = color)) +  
  geom_bar(stat = "count", position = "dodge")
```



it is possible to obtain an horizontal bar chart simply adding coord_flip() aesthetic to the ggplot object:

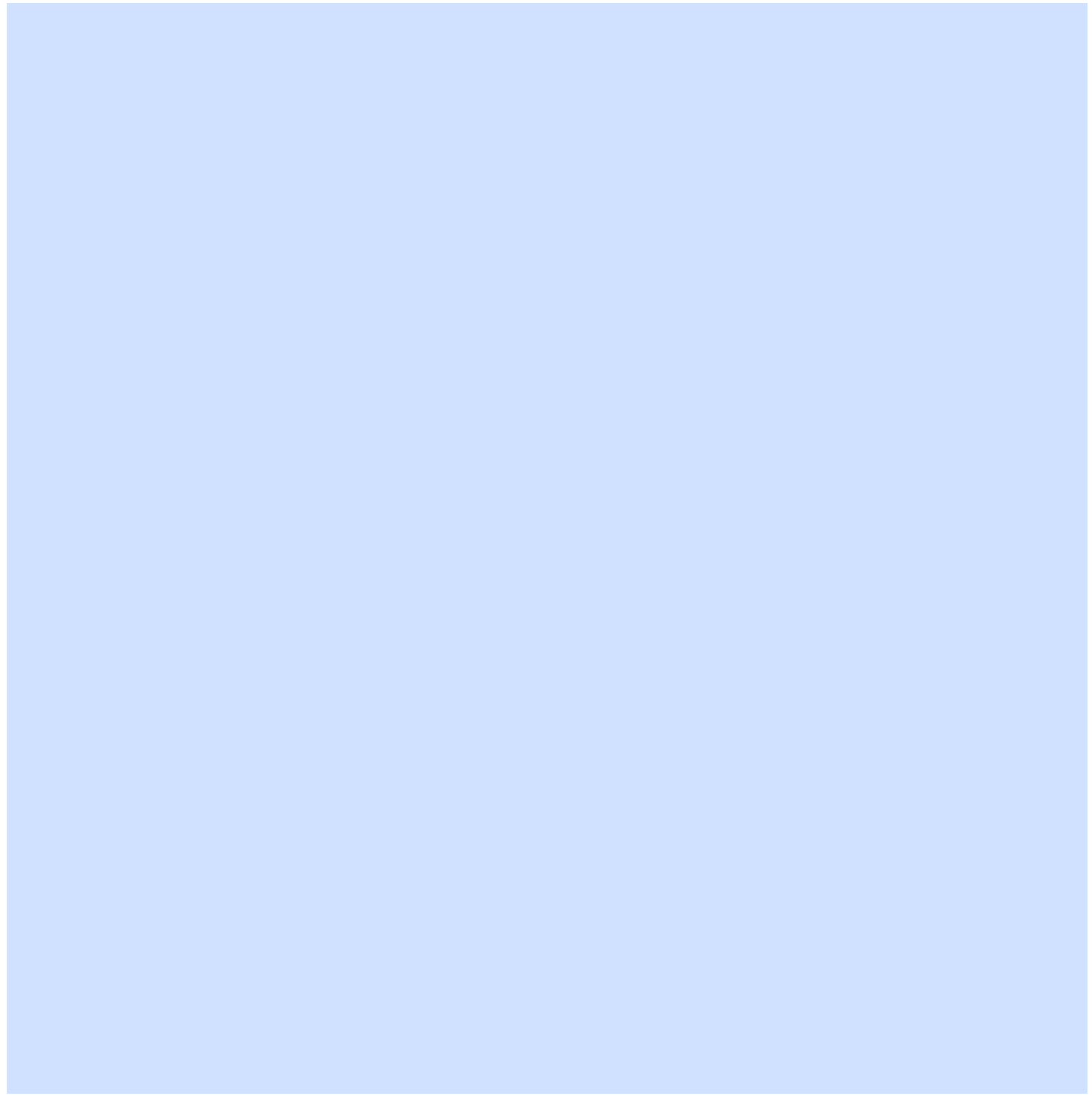
```
ggplot(data = diamonds, aes(x = cut, fill = color)) +  
  geom_bar(stat = "count", position = "dodge") +  
  coord_flip()
```



Section 28.7: Violin plot

Violin plots are kernel density estimates mirrored in the vertical plane. They can be used to visualize several distributions side-by-side, with the mirroring helping to highlight any differences.

```
ggplot(diamonds, aes(cut, price)) +  
  geom_violin()
```



Violin plots are named for their resemblance to the musical instrument, this is particularly visible when they are coupled with an overlaid boxplot. This visualisation then describes the underlying distributions both in terms of Tukey's 5 number summary (as boxplots) and full continuous density estimates (violins).

```
ggplot(diamonds, aes(cut, price)) +  
  geom_violin() +  
  geom_boxplot(width = .1, fill = "black", outlier.shape = NA) +  
  stat_summary(fun.y = "median", geom = "point", col = "white")
```



Chapter 29: Factors

Section 29.1: Consolidating Factor Levels with a List

There are times in which it is desirable to consolidate factor levels into fewer groups, perhaps because of sparse data in one of the categories. It may also occur when you have varying spellings or capitalization of the category names. Consider as an example the factor

```
set.seed(1)
colorful <- sample(c("red", "Red", "RED", "blue", "Blue", "BLUE", "green", "gren"),
                    size = 20,
                    replace = TRUE)
colorful <- factor(colorful)
```

Since R is case-sensitive, a frequency table of this vector would appear as below.

```
table(colorful)
```

colorful	blue	Blue	BLUE	green	gren	red	Red	RED
3	1	4	2	4	1	3	2	

This table, however, doesn't represent the true distribution of the data, and the categories may effectively be reduced to three types: Blue, Green, and Red. Three examples are provided. The first illustrates what seems like an obvious solution, but won't actually provide a solution. The second gives a working solution, but is verbose and computationally expensive. The third is not an obvious solution, but is relatively compact and computationally efficient.

Consolidating levels using `factor(factor_approach)`

```
factor(as.character(colorful),
      levels = c("blue", "Blue", "BLUE", "green", "gren", "red", "Red", "RED"),
      labels = c("Blue", "Blue", "Blue", "Green", "Green", "Red", "Red", "Red"))

[1] Green Blue  Red   Red   Blue  Red   Red   Red   Blue  Red   Green Green Green Blue
    Red   Green
[17] Red   Green Green Red
Levels: Blue Blue Blue Green Green Red Red Red
Warning message:
In `levels<-`(`*tmp*`, value = if (nl == nL) as.character(labels) else paste0(labels, :
  duplicated levels in factors are deprecated
```

Notice that there are duplicated levels. We still have three categories for "Blue", which doesn't complete our task of consolidating levels. Additionally, there is a warning that duplicated levels are deprecated, meaning that this code may generate an error in the future.

Consolidating levels using `ifelse(ifelse_approach)`

```
factor(ifelse(colorful %in% c("blue", "Blue", "BLUE"),
              "Blue",
              ifelse(colorful %in% c("green", "gren"),
                     "Green",
                     "Red")))
```

[1]	Green	Blue	Red	Red	Blue	Red	Red	Red	Blue	Red	Green	Green	Green	Blue
	Red	Green												

```
[17] Red  Green Green Red  
Levels: Blue Green Red
```

This code generates the desired result, but requires the use of nested `ifelse` statements. While there is nothing wrong with this approach, managing nested `ifelse` statements can be a tedious task and must be done carefully.

Consolidating Factors Levels with a List (list_approach)

A less obvious way of consolidating levels is to use a list where the name of each element is the desired category name, and the element is a character vector of the levels in the factor that should map to the desired category. This has the added advantage of working directly on the `levels` attribute of the factor, without having to assign new objects.

```
levels(colorful) <-  
  list("Blue" = c("blue", "Blue", "BLUE"),  
       "Green" = c("green", "gren"),  
       "Red" = c("red", "Red", "RED"))
```

```
[1] Green Blue  Red  Red  Blue  Red  Red  Red  Blue  Red  Green Green Green Blue  
Red  Green  
[17] Red  Green Green Red  
Levels: Blue Green Red
```

Benchmarking each approach

The time required to execute each of these approaches is summarized below. (For the sake of space, the code to generate this summary is not shown)

	expr	min	lq	mean	median	uq	max	neval	cld
factor		78.725	83.256	93.26023	87.5030	97.131	218.899	100	b
ifelse		104.494	107.609	123.53793	113.4145	128.281	254.580	100	c
list_approach		49.557	52.955	60.50756	54.9370	65.132	138.193	100	a

The list approach runs about twice as fast as the `ifelse` approach. However, except in times of very, very large amounts of data, the differences in execution time will likely be measured in either microseconds or milliseconds. With such small time differences, efficiency need not guide the decision of which approach to use. Instead, use an approach that is familiar and comfortable, and which you and your collaborators will understand on future review.

Section 29.2: Basic creation of factors

Factors are one way to represent categorical variables in R. A factor is stored internally as a **vector of integers**. The unique elements of the supplied character vector are known as the *levels* of the factor. By default, if the levels are not supplied by the user, then R will generate the set of unique values in the vector, sort these values alphanumerically, and use them as the levels.

```
charvar <- rep(c("n", "c"), each = 3)  
f <- factor(charvar)  
f  
levels(f)  
  
> f  
[1] n n n c c c  
Levels: c n  
> levels(f)
```

```
[1] "c" "n"
```

If you want to change the ordering of the levels, then one option is to specify the levels manually:

```
levels(factor(charvar, levels = c("n", "c")))

> levels(factor(charvar, levels = c("n", "c")))
[1] "n" "c"
```

Factors have a number of properties. For example, levels can be given labels:

```
> f <- factor(charvar, levels=c("n", "c"), labels=c("Newt", "Capybara"))
> f
[1] Newt      Newt      Newt      Capybara Capybara Capybara
Levels: Newt Capybara
```

Another property that can be assigned is whether the factor is ordered:

```
> Weekdays <- factor(c("Monday", "Wednesday", "Thursday", "Tuesday", "Friday", "Sunday",
  "Saturday"))
> Weekdays
[1] Monday      Wednesday Thursday Tuesday     Friday      Sunday      Saturday
Levels: Friday Monday Saturday Sunday Thursday Tuesday Wednesday
> Weekdays <- factor(Weekdays, levels=c("Monday", "Tuesday", "Wednesday", "Thursday", "Friday",
  "Saturday", "Sunday"), ordered=TRUE)
> Weekdays
[1] Monday      Wednesday Thursday Tuesday     Friday      Sunday      Saturday
Levels: Monday < Tuesday < Wednesday < Thursday < Friday < Saturday < Sunday
```

When a level of the factor is no longer used, you can drop it using the `droplevels()` function:

```
> Weekend <- subset(Weekdays, Weekdays == "Saturday" | Weekdays == "Sunday")
> Weekend
[1] Sunday   Saturday
Levels: Monday < Tuesday < Wednesday < Thursday < Friday < Saturday < Sunday
> Weekend <- droplevels(Weekend)
> Weekend
[1] Sunday   Saturday
Levels: Saturday < Sunday
```

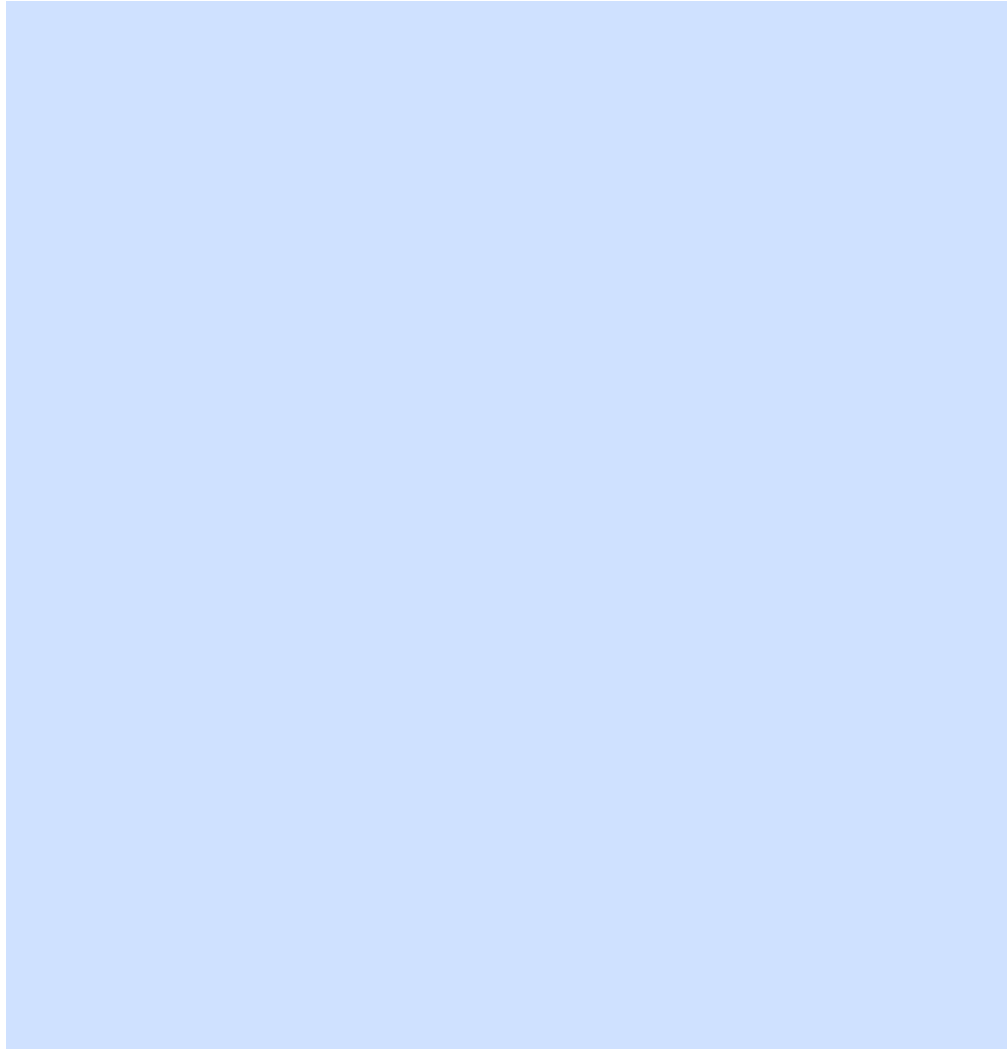
Section 29.3: Changing and reordering factors

When factors are created with defaults, `levels` are formed by `as.character` applied to the inputs and are ordered alphabetically.

```
charvar <- rep(c("W", "n", "c"), times=c(17, 20, 14))
f <- factor(charvar)
levels(f)
# [1] "c" "n" "W"
```

In some situations the treatment of the default ordering of `levels` (alphabetic/lexical order) will be acceptable. For example, if one just wants to `plot` the frequencies, this will be the result:

```
plot(f, col=1:length(levels(f)))
```



But if we want a different ordering of `levels`, we need to specify this in the `levels` or `labels` parameter (taking care that the meaning of "order" here is different from *ordered* factors, see below). There are many alternatives to accomplish that task depending on the situation.

1. Redefine the factor

When it is possible, we can recreate the factor using the `levels` parameter with the order we want.

```
ff <- factor(charvar, levels = c("n", "W", "c"))
levels(ff)
# [1] "n" "W" "c"

gg <- factor(charvar, levels = c("W", "c", "n"))
levels(gg)
# [1] "W" "c" "n"
```

When the input levels are different than the desired output levels, we use the `labels` parameter which causes the `levels` parameter to become a "filter" for acceptable input values, but leaves the final values of "levels" for the factor vector as the argument to `labels`:

```
fm <- factor(as.numeric(f), levels = c(2,3,1),
             labels = c("nn", "WW", "cc"))
levels(fm)
# [1] "nn" "WW" "cc"

fm <- factor(LETTERS[1:6], levels = LETTERS[1:4], # only 'A'-'D' as input
             labels = letters[1:4]) # but assigned to 'a'-'d'
```

```

fm
#[1] a     b     c     d     <NA> <NA>
#Levels: a b c d

```

2. Use `relevel` function

When there is one specific level that needs to be the first we can use `relevel`. This happens, for example, in the context of statistical analysis, when a base category is necessary for testing hypothesis.

```

g<-relevel(f, "n") # moves n to be the first level
levels(g)
# [1] "n" "c" "W"

```

As can be verified f and g are the same

```

all.equal(f, g)
# [1] "Attributes: < Component `levels`: 2 string mismatches >"
all.equal(f, g, check.attributes = F)
# [1] TRUE

```

3. Reordering factors

There are cases when we need to `reorder` the `levels` based on a number, a partial result, a computed statistic, or previous calculations. Let's reorder based on the `frequencies` of the `levels`

```

table(g)
# g
# n   c   W
# 20 14 17

```

The `reorder` function is generic (see `help(reorder)`), but in this context needs: x, in this case the factor; X, a numeric value of the same length as x; and FUN, a function to be applied to X and computed by level of the x, which determines the `levels` order, by default increasing. The result is the same factor with its levels reordered.

```

g.ord <- reorder(g, rep(1, length(g)), FUN=sum) #increasing
levels(g.ord)
# [1] "c" "W" "n"

```

To get de decreasing order we consider negative values (-1)

```

g.ord.d <- reorder(g, rep(-1, length(g)), FUN=sum)
levels(g.ord.d)
# [1] "n" "W" "c"

```

Again the factor is the same as the others.

```

data.frame(f,g,g.ord,g.ord.d)[seq(1,length(g),by=5),] #just same lines
#   f g g.ord g.ord.d
# 1 W W      W      W
# 6 W W      W      W
# 11 W W     W      W
# 16 W W     W      W
# 21 n n     n      n
# 26 n n     n      n
# 31 n n     n      n
# 36 n n     n      n
# 41 c c     c      c

```

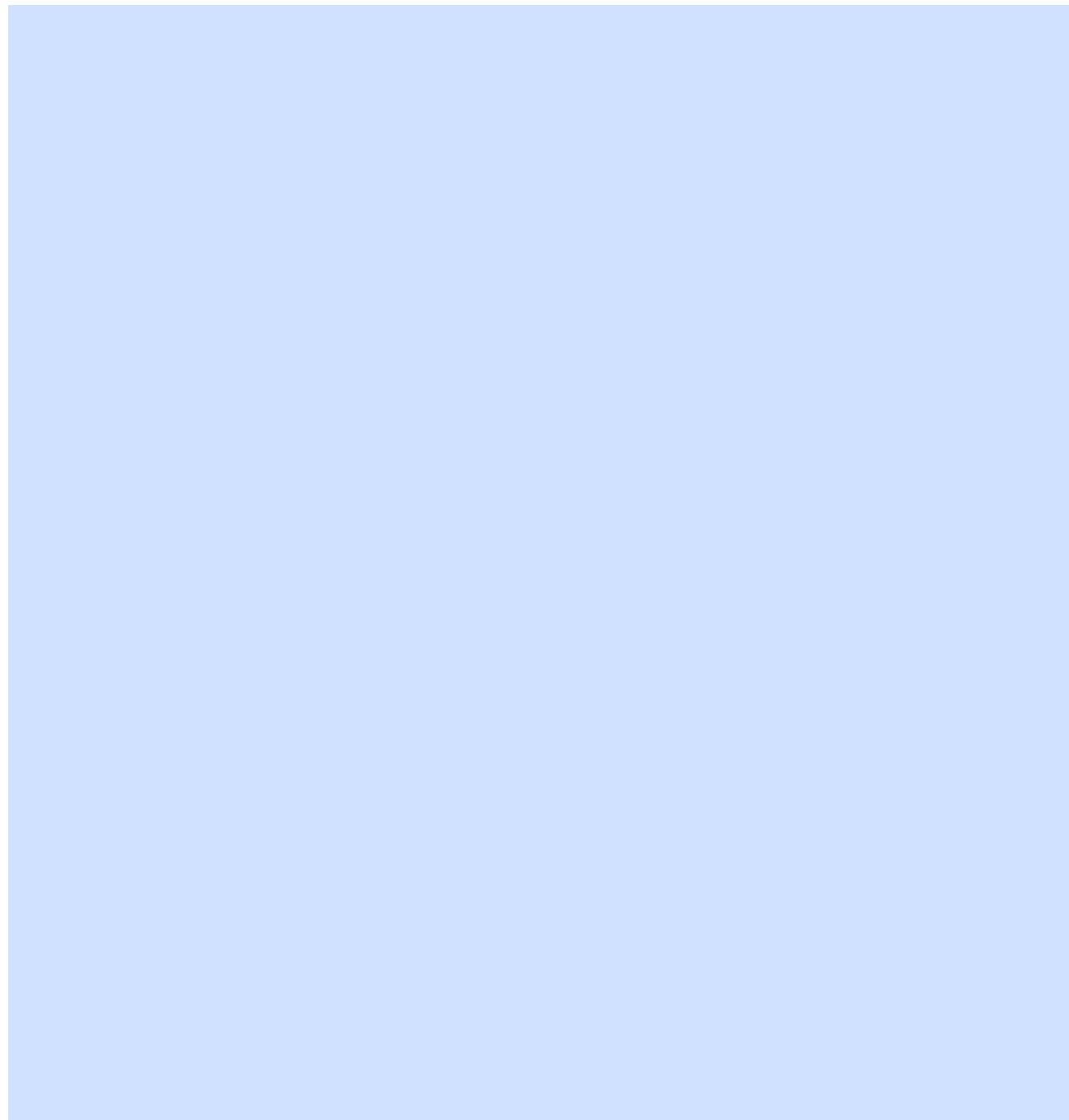
```
# 46 c c      c      c  
# 51 c c      c      c
```

When there is a **quantitative variable** related to the factor variable, we could use other functions to reorder the **levels**. Lets take the **iris** data (`help("iris")` for more information), for reordering the Species factor by using its mean Sepal.Width.

```
miris <- iris #help("iris") # copy the data  
with(miris, tapply(Sepal.Width, Species, mean))  
#   setosa versicolor virginica  
#   3.428     2.770     2.974  
  
miris$Species.o<-with(miris, reorder(Species, -Sepal.Width))  
levels(miris$Species.o)  
# [1] "setosa"    "virginica" "versicolor"
```

The usual **boxplot** (say: `with(miris, boxplot(Petal.Width~Species))`) will show the species in this order: *setosa*, *versicolor*, and *virginica*. But using the ordered factor we get the species ordered by its mean Sepal.Width:

```
boxplot(Petal.Width~Species.o, data = miris,  
        xlab = "Species", ylab = "Petal Width",  
        main = "Iris Data, ordered by mean sepal width", varwidth = TRUE,  
        col = 2:4)
```



Additionally, it is also possible to change the names of `levels`, combine them into groups, or add new `levels`. For that we use the function of the same name `levels`.

```
f1<-f
levels(f1)
# [1] "c" "n" "W"
levels(f1) <- c("upper", "upper", "CAP") #rename and grouping
levels(f1)
# [1] "upper" "CAP"

f2<-f1
levels(f2) <- c("upper", "CAP", "Number") #add Number level, which is empty
levels(f2)
# [1] "upper"   "CAP"     "Number"
f2[length(f2):(length(f2)+5)]<- "Number" # add cases for the new level
table(f2)
# f2
#   upper      CAP Number
#     33       17      6
```

```

f3<-f1
levels(f3) <- list(G1 = "upper", G2 = "CAP", G3 = "Number") # The same using list
levels(f3)
# [1] "G1" "G2" "G3"
f3[length(f3):(length(f3)+6)]<- "G3" ## add cases for the new level
table(f3)
# f3
# G1 G2 G3
# 33 17 7

```

- Ordered factors

Finally, we know that **ordered** factors are different from factors, the first one are used to represent *ordinal data*, and the second one to work with *nominal data*. At first, it does not make sense to change the order of **levels** for ordered factors, but we can change its **labels**.

```

ordvar<-rep(c("Low", "Medium", "High"), times=c(7,2,4))

of<-ordered(ordvar, levels=c("Low", "Medium", "High"))
levels(of)
# [1] "Low"      "Medium"   "High"

of1<-of
levels(of1)<- c("LOW", "MEDIUM", "HIGH")
levels(of1)
# [1] "LOW"      "MEDIUM"   "HIGH"
is.ordered(of1)
# [1] TRUE
of1
# [1] LOW      LOW      LOW      LOW      LOW      LOW      MEDIUM  MEDIUM  HIGH     HIGH     HIGH     HIGH
# Levels: LOW < MEDIUM < HIGH

```

Section 29.4: Rebuilding factors from zero

Problem

Factors are used to represent variables that take values from a set of categories, known as Levels in R. For example, some experiment could be characterized by the energy level of a battery, with four levels: empty, low, normal, and full. Then, for 5 different sampling sites, those levels could be identified, in those terms, as follows:

full, full, normal, empty, low

Typically, in databases or other information sources, the handling of these data is by arbitrary integer indices associated with the categories or levels. If we assume that, for the given example, we would assign, the indices as follows: 1 = empty, 2 = low, 3 = normal, 4 = full, then the 5 samples could be coded as:

4, 4, 3, 1, 2

It could happen that, from your source of information, e.g. a database, you only have the encoded list of integers, and the catalog associating each integer with each level-keyword. How can a factor of R be reconstructed from that information?

Solution

We will simulate a vector of 20 integers that represents the samples, each of which may have one of four different values:

```
set.seed(18)
ii <- sample(1:4, 20, replace=T)
ii
```

```
[1] 4 3 4 1 1 3 2 3 2 1 3 4 1 2 4 1 3 1 4 1
```

The first step is to make a factor, from the previous sequence, in which the levels or categories are exactly the numbers from 1 to 4.

```
fii <- factor(ii, levels=1:4) # it is necessary to indicate the numeric levels
fii
```

```
[1] 4 3 4 1 1 3 2 3 2 1 3 4 1 2 4 1 3 1 4 1
Levels: 1 2 3 4
```

Now simply, you have to *dress* the factor already created with the index tags:

```
levels(fii) <- c("empty", "low", "normal", "full")
fii
```

```
[1] full normal full empty empty normal low normal low empty
[11] normal full empty low full empty normal empty full empty
Levels: empty low normal full
```

Chapter 30: Pattern Matching and Replacement

This topic covers matching string patterns, as well as extracting or replacing them. For details on defining complicated patterns see Regular Expressions.

Section 30.1: Finding Matches

```
# example data
test_sentences <- c("The quick brown fox", "jumps over the lazy dog")
```

Is there a match?

`grep1()` is used to check whether a word or regular expression exists in a string or character vector. The function returns a TRUE/FALSE (or "Boolean") vector.

Notice that we can check each string for the word "fox" and receive a Boolean vector in return.

```
grep1("fox", test_sentences)
#[1] TRUE FALSE
```

Match locations

`grep` takes in a character string and a regular expression. It returns a numeric vector of indexes. This will return which sentence contains the word "fox" in it.

```
grep("fox", test_sentences)
#[1] 1
```

Matched values

To select sentences that match a pattern:

```
# each of the following lines does the job:
test_sentences[grep("fox", test_sentences)]
test_sentences[grep1("fox", test_sentences)]
grep("fox", test_sentences, value = TRUE)
# [1] "The quick brown fox"
```

Details

Since the "fox" pattern is just a word, rather than a regular expression, we could improve performance (with either `grep` or `grep1`) by specifying `fixed = TRUE`.

```
grep("fox", test_sentences, fixed = TRUE)
#[1] 1
```

To select sentences that *don't* match a pattern, one can use `grep` with `invert = TRUE`; or follow subsetting rules with `-grep(...)` or `!grep1(...)`.

In both `grep1(pattern, x)` and `grep(pattern, x)`, the `x` parameter is vectorized, the `pattern` parameter is not. As a result, you cannot use these directly to match `pattern[1]` against `x[1]`, `pattern[2]` against `x[2]`, and so on.

Summary of matches

After performing the e.g. the `grep1` command, maybe you want to get an overview about how many matches where TRUE or FALSE. This is useful e.g. in case of big data sets. In order to do so run the `summary` command:

```
# example data
test_sentences <- c("The quick brown fox", "jumps over the lazy dog")

# find matches
matches <- grep("fox", test_sentences)

# overview
summary(matches)
```

Section 30.2: Single and Global match

When working with regular expressions one modifier for PCRE is g for global match.

In R matching and replacement functions have two version: first match and global match:

- `sub(pattern, replacement, text)` will replace the first occurrence of pattern by replacement in text
- `gsub(pattern, replacement, text)` will do the same as sub but for each occurrence of pattern
- `regexpr(pattern, text)` will return the position of match for the first instance of pattern
- `gregexpr(pattern, text)` will return all matches.

Some random data:

```
set.seed(123)
teststring <- paste0(sample(letters, 20), collapse="")

# teststring
#[1] "htjuwakqxzpgrsbncvyo"
```

Let's see how this works if we want to replace vowels by something else:

```
sub("[aeiou]", " ** HERE WAS A VOWEL** ", teststring)
#[1] "htj ** HERE WAS A VOWEL** wakqxzpgrsbncvyo"

gsub("[aeiou]", " ** HERE WAS A VOWEL** ", teststring)
#[1] "htj ** HERE WAS A VOWEL** w ** HERE WAS A VOWEL** kqxzpgrsbncv ** HERE WAS A VOWEL**  ** HERE
WAS A VOWEL** "
```

Now let's see how we can find a consonant immediately followed by one or more vowel:

```
regexpr("[^aeiou][aeiou]+", teststring)
#[1] 3
#attr(,"match.length")
#[1] 2
#attr(,"useBytes")
#[1] TRUE
```

We have a match on position 3 of the string of length 2, i.e: ju

Now if we want to get all matches:

```
gregexpr("[^aeiou][aeiou]+", teststring)
#[[1]]
#[1] 3 5 19
#attr(,"match.length")
#[1] 2 2 2
```

```
#attr(, "useBytes")
#[1] TRUE
```

All this is really great, but this only give use positions of match and that's not so easy to get what is matched, and here comes `regmatches` it's sole purpose is to extract the string matched from `regexpr`, but it has a different syntax.

Let's save our matches in a variable and then extract them from original string:

```
matches <- gregexpr("[^aeiou][aeiou]+", teststring)
regmatches(teststring, matches)
#[[1]]
#[1] "ju" "wa" "yo"
```

This may sound strange to not have a shortcut, but this allow extraction from another string by the matches of our first one (think comparing two long vector where you know there's a common pattern for the first but not for the second, this allow an easy comparison):

```
teststring2 <- "this is another string to match against"
regmatches(teststring2, matches)
#[[1]]
#[1] "is" " i" "ri"
```

Attention note: by default the pattern is not Perl Compatible Regular Expression, some things like lookarounds are not supported, but each function presented here allow for `perl=TRUE` argument to enable them.

Section 30.3: Making substitutions

```
# example data
test_sentences <- c("The quick brown fox quickly", "jumps over the lazy dog")
```

Let's make the brown fox red:

```
sub("brown", "red", test_sentences)
#[1] "The quick red fox quickly"      "jumps over the lazy dog"
```

Now, let's make the "`fast`" fox act "`fastly`". This won't do it:

```
sub("quick", "fast", test_sentences)
#[1] "The fast red fox quickly"      "jumps over the lazy dog"
```

`sub` only makes the first available replacement, we need `gsub` for global replacement:

```
gsub("quick", "fast", test_sentences)
#[1] "The fast red fox fastly"      "jumps over the lazy dog"
```

See [Modifying strings by substitution](#) for more examples.

Section 30.4: Find matches in big data sets

In case of big data sets, the call of `grep1("fox", test_sentences)` does not perform well. Big data sets are e.g. crawled websites or million of Tweets, etc.

The first acceleration is the usage of the `perl = TRUE` option. Even faster is the option `fixed = TRUE`. A complete example would be:

```
# example data
test_sentences <- c("The quick brown fox", "jumps over the lazy dog")

grep("fox", test_sentences, perl = TRUE)
#[1] TRUE FALSE
```

In case of text mining, often a corpus gets used. A corpus cannot be used directly with `grep1`. Therefore, consider this function:

```
searchCorpus <- function(corpus, pattern) {
  return(tm_index(corpus, FUN = function(x) {
    grep(pattern, x, ignore.case = TRUE, perl = TRUE)
  }))
}
```

Chapter 31: Run-length encoding

Section 31.1: Run-length Encoding with `rle`

Run-length encoding captures the lengths of runs of consecutive elements in a vector. Consider an example vector:

```
dat <- c(1, 2, 2, 2, 3, 1, 4, 4, 1, 1)
```

The `rle` function extracts each run and its length:

```
r <- rle(dat)
r
# Run Length Encoding
#   lengths: int [1:6] 1 3 1 1 2 2
#   values : num [1:6] 1 2 3 1 4 1
```

The values for each run are captured in `r$values`:

```
r$values
# [1] 1 2 3 1 4 1
```

This captures that we first saw a run of 1's, then a run of 2's, then a run of 3's, then a run of 1's, and so on.

The lengths of each run are captured in `r$lengths`:

```
r$lengths
# [1] 1 3 1 1 2 2
```

We see that the initial run of 1's was of length 1, the run of 2's that followed was of length 3, and so on.

Section 31.2: Identifying and grouping by runs in base R

One might want to group their data by the runs of a variable and perform some sort of analysis. Consider the following simple dataset:

```
(dat <- data.frame(x = c(1, 1, 2, 2, 2, 1), y = 1:6))
#   x y
# 1 1 1
# 2 1 2
# 3 2 3
# 4 2 4
# 5 2 5
# 6 1 6
```

The variable `x` has three runs: a run of length 2 with value 1, a run of length 3 with value 2, and a run of length 1 with value 1. We might want to compute the mean value of variable `y` in each of the runs of variable `x` (these mean values are 1.5, 4, and 6).

In base R, we would first compute the run-length encoding of the `x` variable using `rle`:

```
(r <- rle(dat$x))
# Run Length Encoding
#   lengths: int [1:3] 2 3 1
#   values : num [1:3] 1 2 1
```

The next step is to compute the run number of each row of our dataset. We know that the total number of runs is `length(r$lengths)`, and the length of each run is `r$lengths`, so we can compute the run number of each of our runs with `rep`:

```
(run.id <- rep(seq_along(r$lengths), r$lengths))  
# [1] 1 1 2 2 2 3
```

Now we can use `tapply` to compute the mean y value for each run by grouping on the run id:

```
data.frame(x=r$values, meanY=tapply(dat$y, run.id, mean))  
#   x meanY  
# 1 1 1.5  
# 2 2 4.0  
# 3 1 6.0
```

Section 31.3: Run-length encoding to compress and decompress vectors

Long vectors with long runs of the same value can be significantly compressed by storing them in their run-length encoding (the value of each run and the number of times that value is repeated). As an example, consider a vector of length 10 million with a huge number of 1's and only a small number of 0's:

```
set.seed(144)  
dat <- sample(rep(0:1, c(1, 1e5)), 1e7, replace=TRUE)  
table(dat)  
# 0 1  
# 103 9999897
```

Storing 10 million entries will require significant space, but we can instead create a data frame with the run-length encoding of this vector:

```
rle.df <- with(rle(dat), data.frame(values, lengths))  
dim(rle.df)  
# [1] 207 2  
head(rle.df)  
#   values lengths  
# 1     1    52818  
# 2     0      1  
# 3     1   219329  
# 4     0      1  
# 5     1   318306  
# 6     0      1
```

From the run-length encoding, we see that the first 52,818 values in the vector are 1's, followed by a single 0, followed by 219,329 consecutive 1's, followed by a 0, and so on. The run-length encoding only has 207 entries, requiring us to store only 414 values instead of 10 million values. As `rle.df` is a data frame, it can be stored using standard functions like `write.csv`.

Decompressing a vector in run-length encoding can be accomplished in two ways. The first method is to simply call `rep`, passing the `values` element of the run-length encoding as the first argument and the `lengths` element of the run-length encoding as the second argument:

```
decompressed <- rep(rle.df$values, rle.df$lengths)
```

We can confirm that our decompressed data is identical to our original data:

```
identical(decompressed, dat)
# [1] TRUE
```

The second method is to use R's built-in `inverse.rle` function on the `rle` object, for instance:

```
rle.obj <- rle(dat)                                # create a rle object here
class(rle.obj)
# [1] "rle"

dat.inv <- inverse.rle(rle.obj)                      # apply the inverse.rle on the rle object
```

We can confirm again that this produces exactly the original dat:

```
identical(dat.inv, dat)
# [1] TRUE
```

Section 31.4: Identifying and grouping by runs in `data.table`

The `data.table` package provides a convenient way to group by runs in data. Consider the following example data:

```
library(data.table)
(DT <- data.table(x = c(1, 1, 2, 2, 2, 1), y = 1:6))
#    x y
# 1: 1 1
# 2: 1 2
# 3: 2 3
# 4: 2 4
# 5: 2 5
# 6: 1 6
```

The variable `x` has three runs: a run of length 2 with value 1, a run of length 3 with value 2, and a run of length 1 with value 1. We might want to compute the mean value of variable `y` in each of the runs of variable `x` (these mean values are 1.5, 4, and 6).

The `data.table` `rleid` function provides an id indicating the run id of each element of a vector:

```
rleid(DT$x)
# [1] 1 1 2 2 2 3
```

One can then easily group on this run ID and summarize the `y` data:

```
DT[, mean(y), by=.(x, rleid(x))]
#    x rleid V1
# 1: 1     1 1.5
# 2: 2     2 4.0
# 3: 1     3 6.0
```

Chapter 32: Speeding up tough-to-vectorize code

Section 32.1: Speeding tough-to-vectorize for loops with Rcpp

Consider the following tough-to-vectorize for loop, which creates a vector of length `len` where the first element is specified (`first`) and each element `x_i` is equal to `cos(x_{i-1} + 1)`:

```
repeatedCosPlusOne <- function(first, len) {  
  x <- numeric(len)  
  x[1] <- first  
  for (i in 2:len) {  
    x[i] <- cos(x[i-1] + 1)  
  }  
  return(x)  
}
```

This code involves a for loop with a fast operation (`cos(x[i-1]+1)`), which often benefit from vectorization. However, it is not trivial to vectorize this operation with base R, since R does not have a "cumulative cosine of x+1" function.

One possible approach to speeding this function would be to implement it in C++, using the Rcpp package:

```
library(Rcpp)  
cppFunction("NumericVector repeatedCosPlusOneRcpp(double first, int len) {  
  NumericVector x(len);  
  x[0] = first;  
  for (int i=1; i < len; ++i) {  
    x[i] = cos(x[i-1]+1);  
  }  
  return x;  
}")
```

This often provides significant speedups for large computations while yielding the exact same results:

```
all.equal(repeatedCosPlusOne(1, 1e6), repeatedCosPlusOneRcpp(1, 1e6))  
# [1] TRUE  
system.time(repeatedCosPlusOne(1, 1e6))  
#   user   system elapsed  
# 1.274   0.015   1.310  
system.time(repeatedCosPlusOneRcpp(1, 1e6))  
#   user   system elapsed  
# 0.028   0.001   0.030
```

In this case, the Rcpp code generates a vector of length 1 million in 0.03 seconds instead of 1.31 seconds with the base R approach.

Section 32.2: Speeding tough-to-vectorize for loops by byte compiling

Following the Rcpp example in this documentation entry, consider the following tough-to-vectorize function, which creates a vector of length `len` where the first element is specified (`first`) and each element `x_i` is equal to `cos(x_{i-1} + 1)`:

```
repeatedCosPlusOne <- function(first, len) {
```

```

x <- numeric(len)
x[1] <- first
for (i in 2:len) {
  x[i] <- cos(x[i-1] + 1)
}
return(x)
}

```

One simple approach to speeding up such a function without rewriting a single line of code is byte compiling the code using the R `compile` package:

```

library(compiler)
repeatedCosPlusOneCompiled <- cmpfun(repeatedCosPlusOne)

```

The resulting function will often be significantly faster while still returning the same results:

```

all.equal(repeatedCosPlusOne(1, 1e6), repeatedCosPlusOneCompiled(1, 1e6))
# [1] TRUE
system.time(repeatedCosPlusOne(1, 1e6))
#   user  system elapsed
# 1.175  0.014  1.201
system.time(repeatedCosPlusOneCompiled(1, 1e6))
#   user  system elapsed
# 0.339  0.002  0.341

```

In this case, byte compiling sped up the tough-to-vectorize operation on a vector of length 1 million from 1.20 seconds to 0.34 seconds.

Remark

The essence of `repeatedCosPlusOne`, as the cumulative application of a single function, can be expressed more transparently with `Reduce`:

```

iterFunc <- function(init, n, func) {
  funcs <- replicate(n, func)
  Reduce(function(.., f) f(..), funcs, init = init, accumulate = TRUE)
}
repeatedCosPlusOne_vec <- function(first, len) {
  iterFunc(first, len - 1, function(.) cos(. + 1))
}

```

`repeatedCosPlusOne_vec` may be regarded as a "vectorization" of `repeatedCosPlusOne`. However, it can be expected to be *slower* by a factor of 2:

```

library(microbenchmark)
microbenchmark(
  repeatedCosPlusOne(1, 1e4),
  repeatedCosPlusOne_vec(1, 1e4)
)
#> Unit: milliseconds
#>                                              expr      min       lq     mean    median      uq     max neval
cld
#>      repeatedCosPlusOne(1, 10000)  8.349261  9.216724 10.22715 10.23095 11.10817 14.33763   100
a
#>      repeatedCosPlusOne_vec(1, 10000) 14.406291 16.236153 17.55571 17.22295 18.59085 24.37059   100
b

```

Chapter 33: Introduction to Geographical Maps

See also I/O for geographic data

Section 33.1: Basic map-making with `map()` from the package `maps`

The function `map()` from the package `maps` provides a simple starting point for creating maps with R.

A basic world map can be drawn as follows:

```
require(maps)  
map()
```

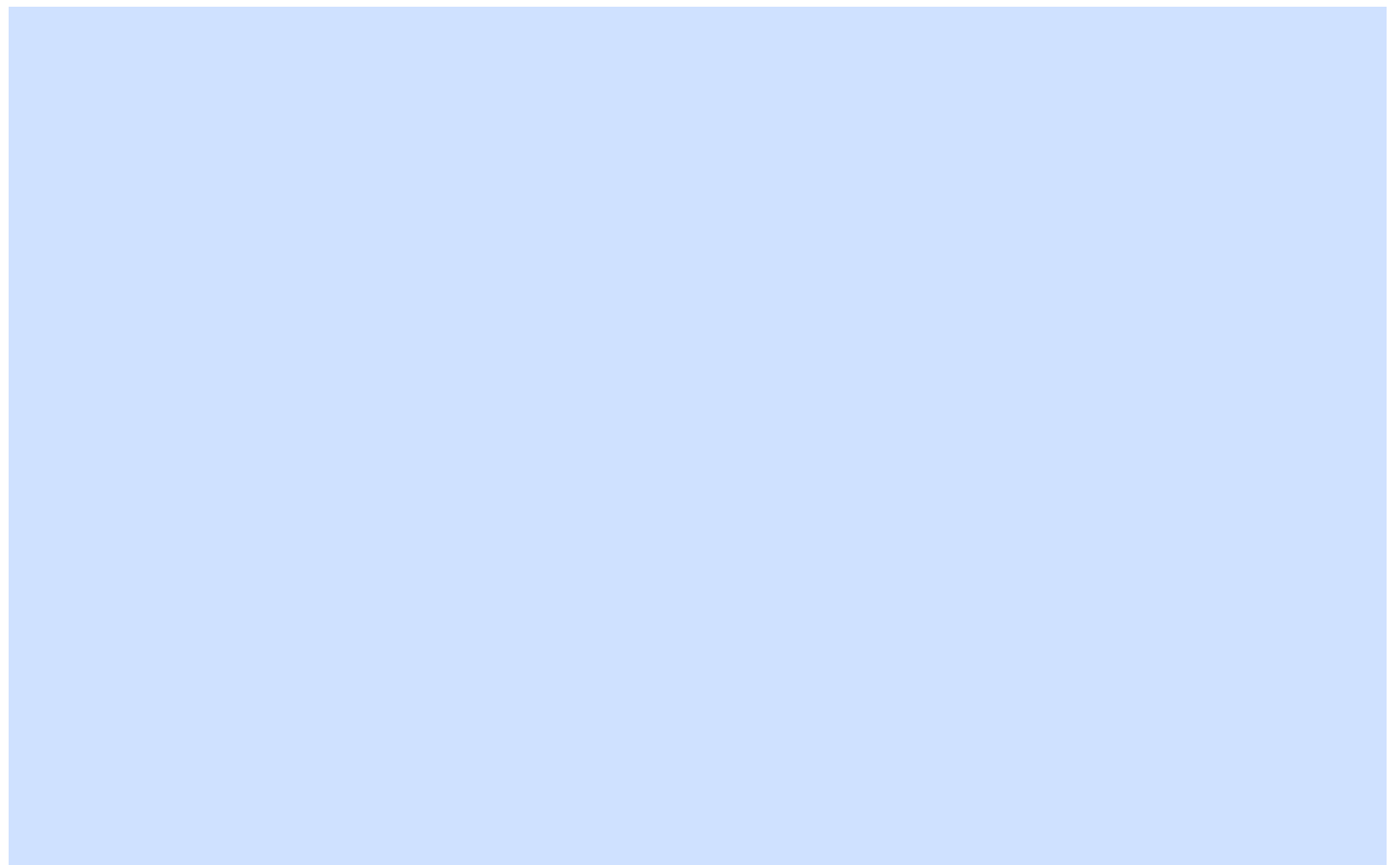
The color of the outline can be changed by setting the color parameter, `col`, to either the character name or hex value of a color:

```
require(maps)  
map(col = "cornflowerblue")
```



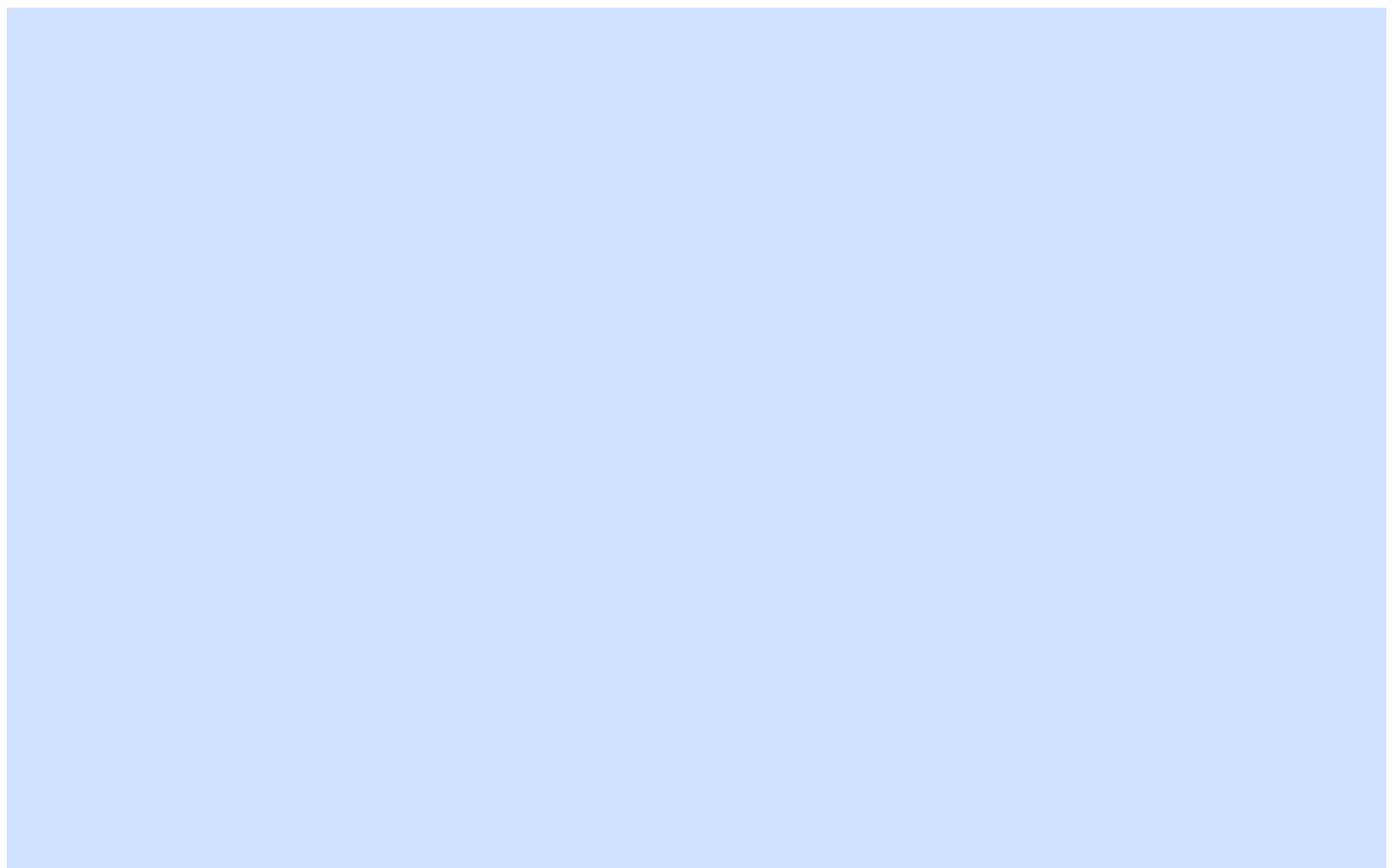
To fill land masses with the color in `col` we can set `fill = TRUE`:

```
require(maps)
map(fill = TRUE, col = c("cornflowerblue"))
```



A vector of any length may be supplied to `col` when `fill = TRUE` is also set:

```
require(maps)
map(fill = TRUE, col = c("cornflowerblue", "limegreen", "hotpink"))
```



In the example above colors from `col` are assigned arbitrarily to polygons in the map representing regions and colors are recycled if there are fewer colors than polygons.

We can also use color coding to represent a statistical variable, which may optionally be described in a legend. A map created as such is known as a "choropleth".

The following choropleth example sets the first argument of `map()`, which is `database` to "`county`" and "`state`" to color code unemployment using data from the built-in datasets `unemp` and `county.fips` while overlaying state lines in white:

```
require(maps)
if(require(mapproj)) {    # mapproj is used for projection="polyconic"
  # color US county map by 2009 unemployment rate
  # match counties to map using FIPS county codes
  # Based on J's solution to the "Choropleth Challenge"
  # Code improvements by Hack-R (hack-r.github.io)

  # load data
  # unemp includes data for some counties not on the "lower 48 states" county
  # map, such as those in Alaska, Hawaii, Puerto Rico, and some tiny Virginia
  # cities
  data(unemp)
  data(county.fips)

  # define color buckets
  colors = c("paleturquoise", "skyblue", "cornflowerblue", "blueviolet", "hotpink", "darkgrey")
  unemp$colorBuckets <- as.numeric(cut(unemp$unemp, c(0, 2, 4, 6, 8, 10)))
```

```

leg.txt <- c("<2%", "2-4%", "4-6%", "6-8%", "8-10%", ">10%")

# align data with map definitions by (partial) matching state, county
# names, which include multiple polygons for some counties
cnty.fips <- county.fips$fips[match(map("county", plot=FALSE)$names,
                                      county.fips$polyname)]
colorsmatched <- unemp$colorBuckets[match(cnty.fips, unemp$fips)]

# draw map
par(mar=c(1, 1, 2, 1) + 0.1)
map("county", col = colors[colorsmatched], fill = TRUE, resolution = 0,
    lty = 0, projection = "polyconic")
map("state", col = "white", fill = FALSE, add = TRUE, lty = 1, lwd = 0.1,
    projection="polyconic")
title("unemployment by county, 2009")
legend("topright", leg.txt, horiz = TRUE, fill = colors, cex=0.6)
}

```



Section 33.2: 50 State Maps and Advanced Choropleths with Google Viz

A common [question](#) is how to juxtapose (combine) physically separate geographical regions on the same map, such as in the case of a choropleth describing all 50 American states (The mainland with Alaska and Hawaii juxtaposed).

Creating an attractive 50 state map is simple when leveraging Google Maps. Interfaces to Google's API include the packages `googleVis`, `ggmap`, and `RgoogleMaps`.

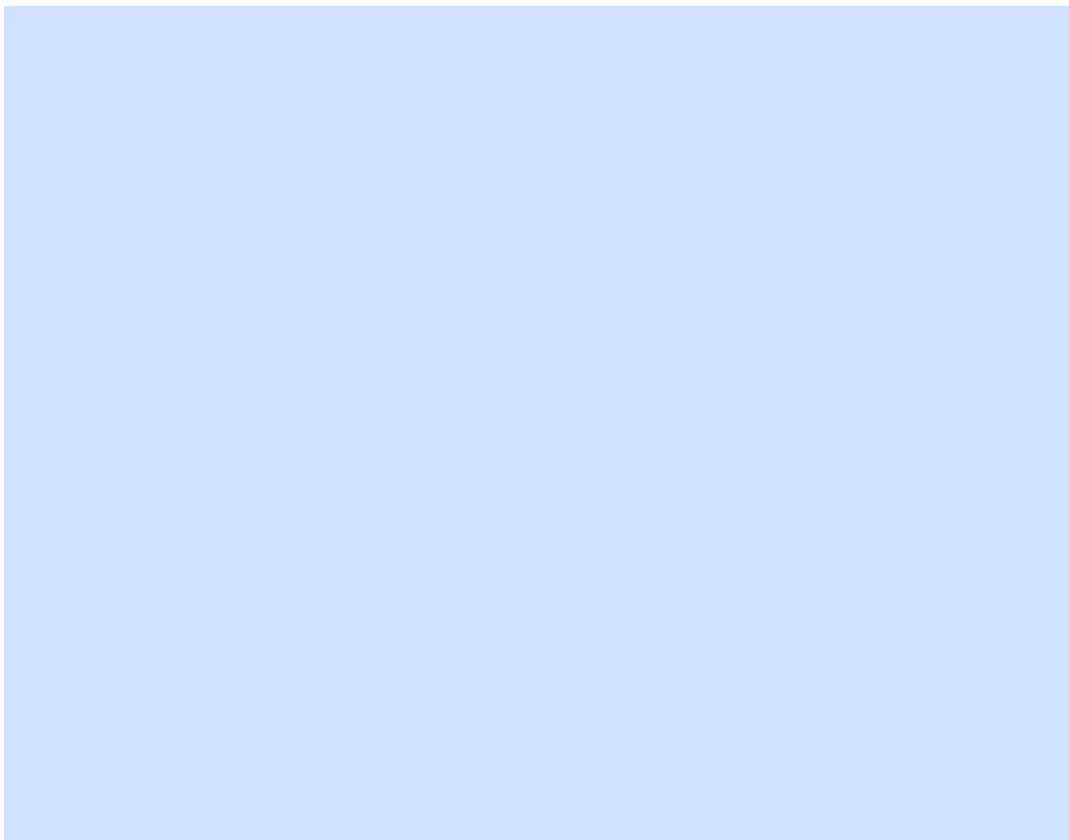
```

require(googleVis)

G4 <- gvisGeoChart(CityPopularity, locationvar='City', colorvar='Popularity',
                     options=list(region='US', height=350,
                                 displayMode='markers',

```

```
        colorAxis="{values:[200,400,600,800],  
                  colors:['red', 'pink', 'orange','green']}")  
    )  
plot(G4)
```



The function `gvisGeoChart()` requires far less coding to create a choropleth compared to older mapping methods, such as `map()` from the package `maps`. The `colorvar` parameter allows easy coloring of a statistical variable, at a level specified by the `locationvar` parameter. The various options passed to `options` as a list allow customization of the map's details such as size (height), shape (markers), and color coding (`colorAxis` and `colors`).

Section 33.3: Interactive plotly maps

The `plotly` package allows many kind of interactive plots, including maps. There are a few ways to create a map in `plotly`. Either supply the map data yourself (via `plot_ly()` or `ggplotly()`), use `plotly`'s "native" mapping capabilities (via `plot_geo()` or `plot_mapbox()`), or even a combination of both. An example of supplying the map yourself would be:

```
library(plotly)  
map_data("county") %>%  
  group_by(group) %>%  
  plot_ly(x = ~long, y = ~lat) %>%  
  add_polygons() %>%  
  layout(  
    xaxis = list(title = "", showgrid = FALSE, showticklabels = FALSE),  
    yaxis = list(title = "", showgrid = FALSE, showticklabels = FALSE)  
)
```

For a combination of both approaches, swap `plot_ly()` for `plot_geo()` or `plot_mapbox()` in the above example. See the [plotly book](#) for more examples.

The next example is a "strictly native" approach that leverages the `layout.geo` attribute to set the aesthetics and zoom level of the map. It also uses the database `world.cities` from `maps` to filter the Brazilian cities and plot them on top of the "native" map.

The main variables: `poph` is a text with the city and its population (which is shown upon mouse hover); `q` is a ordered factor from the population's quantile. `ge` has information for the layout of the maps. See the [package documentation](#) for more information.

```
library(maps)
dfb <- world.cities[world.cities$country.etc=="Brazil",]
library(plotly)
dfb$poph <- paste(dfb$name, "Pop", round(dfb$pop/1e6,2), " millions")
dfb$q <- with(dfb, cut(pop, quantile(pop), include.lowest = T))
levels(dfb$q) <- paste(c("1st", "2nd", "3rd", "4th"), "Quantile")
dfb$q <- as.ordered(dfb$q)

ge <- list(
  scope = 'south america',
  showland = TRUE,
  landcolor = toRGB("gray85"),
  subunitwidth = 1,
  countrywidth = 1,
  subunitcolor = toRGB("white"),
  countrycolor = toRGB("white")
)

plot_geo(dfb, lon = ~long, lat = ~lat, text = ~poph,
  marker = ~list(size = sqrt(pop/10000) + 1, line = list(width = 0)),
  color = ~q, locationmode = 'country names') %>%
  layout(geo = ge, title = 'Populations<br>(Click legend to toggle)')
```



Section 33.4: Making Dynamic HTML Maps with Leaflet

[Leaflet](#) is an open-source JavaScript library for making dynamic maps for the web. RStudio wrote R bindings for Leaflet, available through its [Leaflet package](#), built with [htmlwidgets](#). Leaflet maps integrate well with the [RMarkdown](#) and [Shiny](#) ecosystems.

The interface is piped, using a `leaflet()` function to initialize a map and subsequent functions adding (or removing) map layers. Many kinds of layers are available, from markers with popups to polygons for creating choropleth maps. Variables in the `data.frame` passed to `leaflet()` are accessed via function-style `~` quotation.

To map the [state.name](#) and [state.center](#) datasets:

```
library(leaflet)

data.frame(state.name, state.center) %>%
  leaflet() %>%
  addProviderTiles('Stamen.Watercolor') %>%
  addMarkers(lng = ~x, lat = ~y,
             popup = ~state.name,
             clusterOptions = markerClusterOptions())
```



(Screenshot; click for dynamic version.)

Section 33.5: Dynamic Leaflet maps in Shiny applications

The [Leaflet](#) package is designed to [integrate with Shiny](#)

In the **ui** you call `leafletOutput()` and in the server you call `renderLeaflet()`

```
library(shiny)
library(leaflet)

ui <- fluidPage(
  leafletOutput("my_leaf")
)

server <- function(input, output, session){

  output$my_leaf <- renderLeaflet({

    leaflet() %>%
      addProviderTiles('Hydda.Full') %>%
      setView(lat = -37.8, lng = 144.8, zoom = 10)

  })
}

shinyApp(ui, server)
```

However, reactive inputs that affect the `renderLeaflet` expression will cause the entire map to be redrawn each time the reactive element is updated.

Therefore, to modify a map that's already running you should use the `leafletProxy()` function.

Normally you use leaflet to create the static aspects of the map, and `leafletProxy` to manage the dynamic elements, for example:

```
library(shiny)
library(leaflet)

ui <- fluidPage(
  sliderInput(inputId = "slider",
              label = "values",
              min = 0,
              max = 100,
              value = 0,
              step = 1),
  leafletOutput("my_leaf")
)

server <- function(input, output, session){
  set.seed(123456)
  df <- data.frame(latitude = sample(seq(-38.5, -37.5, by = 0.01), 100),
                    longitude = sample(seq(144.0, 145.0, by = 0.01), 100),
                    value = seq(1,100))

  ## create static element
  output$my_leaf <- renderLeaflet({

    leaflet() %>%
      addProviderTiles('Hydda.Full') %>%
      setView(lat = -37.8, lng = 144.8, zoom = 8)

  })

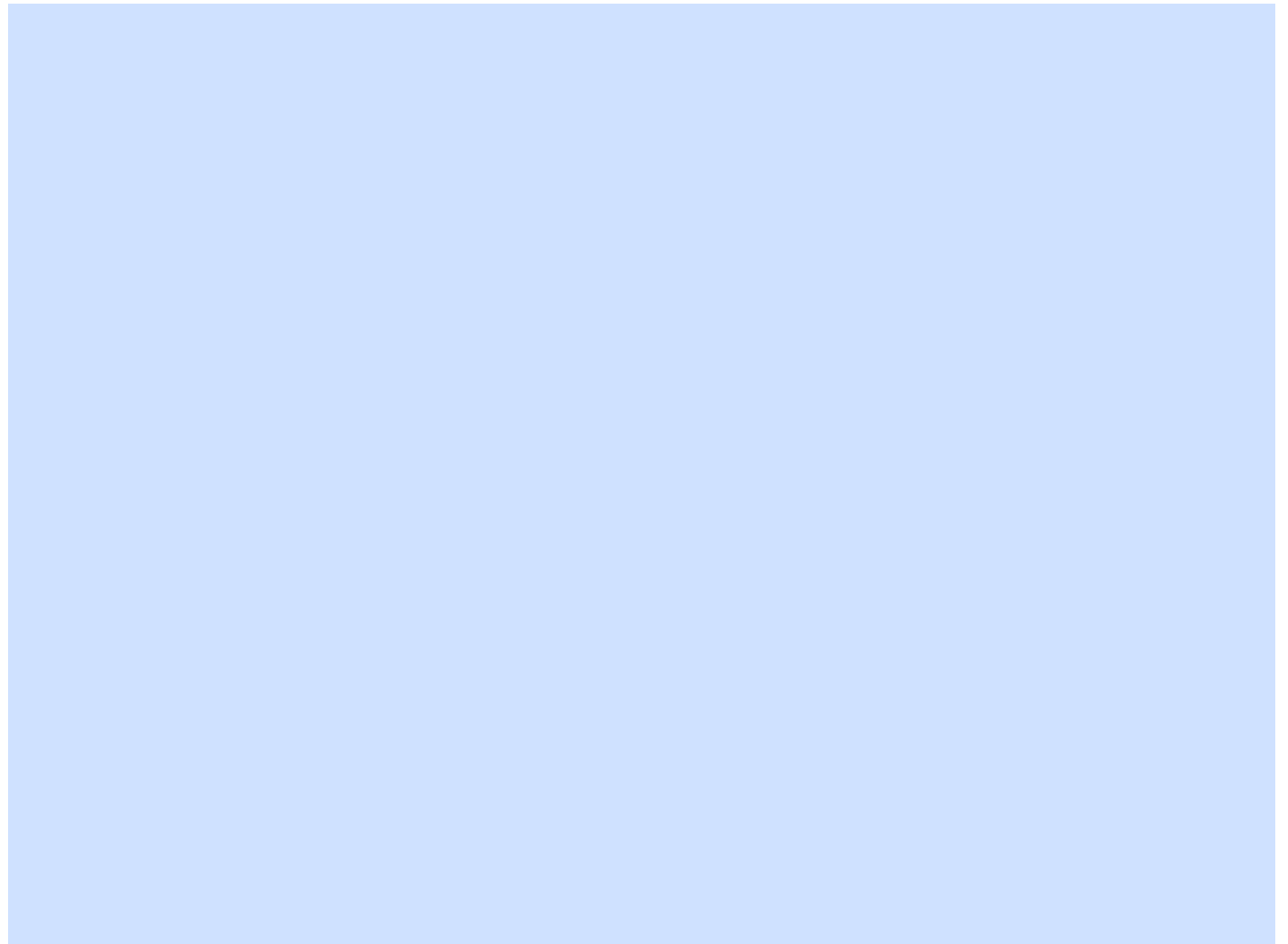
  ## filter data
  df_filtered <- reactive({
    df[df$value >= input$slider, ]
  })

  ## respond to the filtered data
  observe({

    leafletProxy(mapId = "my_leaf", data = df_filtered()) %>%
      clearMarkers() %>% ## clear previous markers
      addMarkers()

  })
}

shinyApp(ui, server)
```



Chapter 34: Set operations

Section 34.1: Set operators for pairs of vectors

Comparing sets

In R, a vector may contain duplicated elements:

```
v = "A"  
w = c("A", "A")
```

However, a set contains only one copy of each element. R treats a vector like a set by taking only its distinct elements, so the two vectors above are regarded as the same:

```
setequal(v, w)  
# TRUE
```

Combining sets

The key functions have natural names:

```
x = c(1, 2, 3)  
y = c(2, 4)  
  
union(x, y)  
# 1 2 3 4  
  
intersect(x, y)  
# 2  
  
setdiff(x, y)  
# 1 3
```

These are all documented on the same page, [?union](#).

Section 34.2: Cartesian or "cross" products of vectors

To find every vector of the form (x, y) where x is drawn from vector X and y from Y , we use `expand.grid`:

```
X = c(1, 1, 2)  
Y = c(4, 5)  
  
expand.grid(X, Y)  
  
#   Var1 Var2  
# 1     1    4  
# 2     1    4  
# 3     2    4  
# 4     1    5  
# 5     1    5  
# 6     2    5
```

The result is a data.frame with one column for each vector passed to it. Often, we want to take the Cartesian product of sets rather than to expand a "grid" of vectors. We can use `unique`, `lapply` and `do.call`:

```
m = do.call(expand.grid, lapply(list(X, Y), unique))
```

```
#   Var1 Var2
# 1   1   4
# 2   2   4
# 3   1   5
# 4   2   5
```

Applying functions to combinations

If you then want to apply a function to each resulting combination $f(x, y)$, it can be added as another column:

```
m$p = with(m, Var1*Var2)
#   Var1 Var2  p
# 1   1   4   4
# 2   2   4   8
# 3   1   5   5
# 4   2   5  10
```

This approach works for as many vectors as we need, but in the special case of two, it is sometimes a better fit to have the result in a matrix, which can be achieved with `outer`:

```
uX = unique(X)
uY = unique(Y)

outer(setNames(uX, uX), setNames(uY, uY), `*`)

#   4   5
# 1 4   5
# 2 8  10
```

For related concepts and tools, see the combinatorics topic.

Section 34.3: Set membership for vectors

The `%in%` operator compares a vector with a set.

```
v = "A"
w = c("A", "A")

w %in% v
# TRUE TRUE

v %in% w
# TRUE
```

Each element on the left is treated individually and tested for membership in the set associated with the vector on the right (consisting of all its distinct elements).

Unlike equality tests, `%in%` always returns TRUE or FALSE:

```
c(1, NA) %in% c(1, 2, 3, 4)
# TRUE FALSE
```

The documentation is at `?`%in%``.

Section 34.4: Make unique / drop duplicates / select distinct

elements from a vector

`unique` drops duplicates so that each element in the result is unique (only appears once):

```
x = c(2, 1, 1, 2, 1)  
  
unique(x)  
# 2 1
```

Values are returned in the order they first appeared.

`duplicated` tags each duplicated element:

```
duplicated(x)  
# FALSE FALSE TRUE TRUE TRUE
```

`anyDuplicated(x) > 0L` is a quick way of checking whether a vector contains any duplicates.

Section 34.5: Measuring set overlaps / Venn diagrams for vectors

To count how many elements of two sets overlap, one could write a custom function:

```
xtab_set <- function(A, B){  
  both     <- union(A, B)  
  inA      <- both %in% A  
  inB      <- both %in% B  
  return(table(inA, inB))  
}  
  
A = 1:20  
B = 10:30  
  
xtab_set(A, B)  
  
#           inB  
# inA      FALSE  TRUE  
# FALSE      0   10  
#  TRUE      9   11
```

A Venn diagram, offered by various packages, can be used to visualize overlap counts across multiple sets.

Chapter 35: tidyverse

Section 35.1: tidyverse: an overview

What is tidyverse?

[tidyverse](#) is the fast and elegant way to turn basic R into an enhanced tool, redesigned by Hadley/Rstudio. The development of all packages included in tidyverse follow the principle rules of [The tidy tools manifesto](#). But first, let the authors describe their masterpiece:

The tidyverse is a set of packages that work in harmony because they share common data representations and API design. The tidyverse package is designed to make it easy to install and load core packages from the tidyverse in a single command.

The best place to learn about all the packages in the tidyverse and how they fit together is R for Data Science. Expect to hear more about the tidyverse in the coming months as I work on improved package websites, making citation easier, and providing a common home for discussions about data analysis with the tidyverse.

([source](#))

How to use it?

Just with the ordinary R packages, you need to install and load the package.

```
install.package("tidyverse")
library("tidyverse")
```

The difference is, on a single command a couple of dozens of packages are installed/loaded. As a bonus, one may rest assured that all the installed/loaded packages are of compatible versions.

What are those packages?

The commonly known and widely used packages:

- [ggplot2](#): advanced data visualisation SO_doc
- [dplyr](#): fast (Rcpp) and coherent approach to data manipulation SO_doc
- [tidyr](#): tools for data tidying SO_doc
- [readr](#): for data import.
- [purrr](#): makes your pure functions purr by completing R's functional programming tools with important features from other languages, in the style of the JS packages underscore.js, lodash and lazy.js.
- [tibble](#): a modern re-imagining of data frames.
- [magrittr](#): piping to make code more readable SO_doc

Packages for manipulating specific data formats:

- [hms](#): easily read times
- [stringr](#): provide a cohesive set of functions designed to make working with strings as easy as possible
- [lubridate](#): advanced date/times manipulations SO_doc
- [forcats](#): advanced work with factors.

Data import:

- [DBI](#): defines a common interface between the R and database management systems (DBMS)
- [haven](#): easily import SPSS, SAS and Stata files SO_doc
- [httr](#): the aim of httr is to provide a wrapper for the curl package, customised to the demands of modern web APIs
- [jsonlite](#): a fast JSON parser and generator optimized for statistical data and the web
- [readxl](#): read.xls and .xlsx files without need for dependency packages SO_doc
- [rvest](#): rvest helps you scrape information from web pages SO_doc
- [xml2](#): for XML

And modelling:

- [modelr](#): provides functions that help you create elegant pipelines when modelling
- [broom](#): easily extract the models into tidy data

Finally, tidyverse suggest the use of:

- [knitr](#): the amazing general-purpose literate programming engine, with lightweight API's designed to give users full control of the output without heavy coding work. SO_docs: one, two
- [rmarkdown](#): Rstudio's package for reproducible programming. SO_docs: one, two, three, four

Section 35.2: Creating `tbl_df`'s

A `tbl_df` (pronounced *tibble diff*) is a variation of a data frame that is often used in tidyverse packages. It is implemented in the [tibble](#) package.

Use the `as_data_frame` function to turn a data frame into a `tbl_df`:

```
library(tibble)
mtcars_tbl <- as_data_frame(mtcars)
```

One of the most notable differences between `data.frames` and `tbl_dfs` is how they print:

```
# A tibble: 32 x 11
  mpg   cyl  disp    hp  drat    wt  qsec    vs    am  gear  carb
* <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl>
1 21.0     6 160.0   110  3.90 2.620 16.46     0     1     4     4
2 21.0     6 160.0   110  3.90 2.875 17.02     0     1     4     4
3 22.8     4 108.0    93  3.85 2.320 18.61     1     1     4     1
4 21.4     6 258.0   110  3.08 3.215 19.44     1     0     3     1
5 18.7     8 360.0   175  3.15 3.440 17.02     0     0     3     2
6 18.1     6 225.0   105  2.76 3.460 20.22     1     0     3     1
7 14.3     8 360.0   245  3.21 3.570 15.84     0     0     3     4
8 24.4     4 146.7    62  3.69 3.190 20.00     1     0     4     2
9 22.8     4 140.8    95  3.92 3.150 22.90     1     0     4     2
10 19.2    6 167.6   123  3.92 3.440 18.30     1     0     4     4
# ... with 22 more rows
```

- The printed output includes a summary of the dimensions of the table (`32 x 11`)
- It includes the type of each column (`dbl`)
- It prints a limited number of rows. (To change this use `options(tibble.print_max = [number])`).

Many functions in the `dplyr` package work naturally with `tbl_dfs`, such as `group_by()`.

Chapter 36: Rcpp

Section 36.1: Extending Rcpp with Plugins

Within C++, one can set different compilation flags using:

```
// [[Rcpp::plugins(name)]]
```

List of the built-in plugins:

```
// built-in C++11 plugin
// [[Rcpp::plugins(cpp11)]]

// built-in C++11 plugin for older g++ compiler
// [[Rcpp::plugins(cpp0x)]]

// built-in C++14 plugin for C++14 standard
// [[Rcpp::plugins(cpp14)]]

// built-in C++1y plugin for C++14 and C++17 standard under development
// [[Rcpp::plugins(cpp1y)]]

// built-in OpenMP++11 plugin
// [[Rcpp::plugins(openmp)]]
```

Section 36.2: Inline Code Compile

Rcpp features two functions that enable code compilation inline and exportation directly into R: `cppFunction()` and `evalCpp()`. A third function called `sourceCpp()` exists to read in C++ code in a separate file though can be used akin to `cppFunction()`.

Below is an example of compiling a C++ function within R. Note the use of `""` to surround the source.

```
# Note - This is R code.
# cppFunction in Rcpp allows for rapid testing.
require(Rcpp)

# Creates a function that multiples each element in a vector
# Returns the modified vector.
cppFunction(
  NumericVector exfun(NumericVector x, int i){
    x = x*i;
    return x;
  })

# Calling function in R
exfun(1:5, 3)
```

To quickly understand a C++ expression use:

```
# Use evalCpp to evaluate C++ expressions
evalCpp("std::numeric_limits<double>::max()")
## [1] 1.797693e+308
```

Section 36.3: Rcpp Attributes

Rcpp Attributes makes the process of working with R and C++ straightforward. The form of attributes take:

```
// [[Rcpp::attribute]]
```

The use of attributes is typically associated with:

```
// [[Rcpp::export]]
```

that is placed directly above a declared function header when reading in a C++ file via `sourceCpp()`.

Below is an example of an external C++ file that uses attributes.

```
// Add code below into C++ file Rcpp_example.cpp

#include <Rcpp.h>
using namespace Rcpp;

// Place the export tag right above function declaration.
// [[Rcpp::export]]
double muRcpp(NumericVector x){

    int n = x.size(); // Size of vector
    double sum = 0; // Sum value

    // For loop, note cpp index shift to 0
    for(int i = 0; i < n; i++){
        // Shorthand for sum = sum + x[i]
        sum += x[i];
    }

    return sum/n; // Obtain and return the Mean
}

// Place dependent functions above call or
// declare the function definition with:
double muRcpp(NumericVector x);

// [[Rcpp::export]]
double varRcpp(NumericVector x, bool bias = true){

    // Calculate the mean using C++ function
    double mean = muRcpp(x);
    double sum = 0;

    int n = x.size();

    for(int i = 0; i < n; i++){
        sum += pow(x[i] - mean, 2.0); // Square
    }

    return sum/(n-bias); // Return variance
}
```

To use this external C++ file within R, we do the following:

```
require(Rcpp)
```

```

# Compile File
sourceCpp("path/to/file/Rcpp_example.cpp")

# Make some sample data
x = 1:5

all.equal(muRcpp(x), mean(x))
## TRUE

all.equal(varRcpp(x), var(x))
## TRUE

```

Section 36.4: Specifying Additional Build Dependencies

To use additional packages within the Rcpp ecosystem, the correct header file may not be `Rcpp.h` but `Rcpp<PACKAGE>.h` (as e.g. for [RcppArmadillo](#)). It typically needs to be imported and then the dependency is stated within

```
// [[Rcpp::depends(Rcpp<PACKAGE>)]]
```

Examples:

```

// Use the RcppArmadillo package
// Requires different header file from Rcpp.h
#include <RcppArmadillo.h>
// [[Rcpp::depends(RcppArmadillo)]]


// Use the RcppEigen package
// Requires different header file from Rcpp.h
#include <RcppEigen.h>
// [[Rcpp::depends(RcppEigen)]]

```

Chapter 37: Random Numbers Generator

Section 37.1: Random permutations

To generate random permutation of 5 numbers:

```
sample(5)
# [1] 4 5 3 1 2
```

To generate random permutation of any vector:

```
sample(10:15)
# [1] 11 15 12 10 14 13
```

One could also use the package `pracma`

```
randperm(a, k)
# Generates one random permutation of k of the elements a, if a is a vector,
# or of 1:a if a is a single integer.
# a: integer or numeric vector of some length n.
# k: integer, smaller as a or length(a).

# Examples
library(pracma)
randperm(1:10, 3)
[1] 3 7 9

randperm(10, 10)
[1] 4 5 10 8 2 7 6 9 3 1

randperm(seq(2, 10, by=2))
[1] 6 4 10 2 8
```

Section 37.2: Generating random numbers using various density functions

Below are examples of generating 5 random numbers using various probability distributions.

Uniform distribution between 0 and 10

```
runif(5, min=0, max=10)
[1] 2.1724399 8.9209930 6.1969249 9.3303321 2.4054102
```

Normal distribution with 0 mean and standard deviation of 1

```
rnorm(5, mean=0, sd=1)
[1] -0.97414402 -0.85722281 -0.08555494 -0.37444299 1.20032409
```

Binomial distribution with 10 trials and success probability of 0.5

```
rbinom(5, size=10, prob=0.5)
[1] 4 3 5 2 3
```

Geometric distribution with 0.2 success probability

```
rgeom(5, prob=0.2)
[1] 14 8 11 1 3
```

Hypergeometric distribution with 3 white balls, 10 black balls and 5 draws

```
rhyper(5, m=3, n=10, k=5)
[1] 2 0 1 1 1
```

Negative Binomial distribution with 10 trials and success probability of 0.8

```
rnbinom(5, size=10, prob=0.8)
[1] 3 1 3 4 2
```

Poisson distribution with mean and variance (lambda) of 2

```
rpois(5, lambda=2)
[1] 2 1 2 3 4
```

Exponential distribution with the rate of 1.5

```
rexp(5, rate=1.5)
[1] 1.8993303 0.4799358 0.5578280 1.5630711 0.6228000
```

Logistic distribution with 0 location and scale of 1

```
rlogis(5, location=0, scale=1)
[1] 0.9498992 -1.0287433 -0.4192311 0.7028510 -1.2095458
```

Chi-squared distribution with 15 degrees of freedom

```
rchisq(5, df=15)
[1] 14.89209 19.36947 10.27745 19.48376 23.32898
```

Beta distribution with shape parameters a=1 and b=0.5

```
rbeta(5, shape1=1, shape2=0.5)
[1] 0.1670306 0.5321586 0.9869520 0.9548993 0.9999737
```

Gamma distribution with shape parameter of 3 and scale=0.5

```
rgamma(5, shape=3, scale=0.5)
[1] 2.2445984 0.7934152 3.2366673 2.2897537 0.8573059
```

Cauchy distribution with 0 location and scale of 1

```
rcauchy(5, location=0, scale=1)
[1] -0.01285116 -0.38918446 8.71016696 10.60293284 -0.68017185
```

Log-normal distribution with 0 mean and standard deviation of 1 (on log scale)

```
rlnorm(5, meanlog=0, sdlog=1)
[1] 0.8725009 2.9433779 0.3329107 2.5976206 2.8171894
```

Weibull distribution with shape parameter of 0.5 and scale of 1

```
rweibull(5, shape=0.5, scale=1)
[1] 0.337599112 1.307774557 7.233985075 5.840429942 0.005751181
```

Wilcoxon distribution with 10 observations in the first sample and 20 in second.

```
rwilcox(5, 10, 20)
[1] 111 88 93 100 124
```

Multinomial distribution with 5 object and 3 boxes using the specified probabilities

```
rmultinom(5, size=5, prob=c(0.1,0.1,0.8))
 [,1] [,2] [,3] [,4] [,5]
[1,]    0    0    1    1    0
[2,]    2    0    1    1    0
[3,]    3    5    3    3    5
```

Section 37.3: Random number generator's reproducibility

When expecting someone to reproduce an R code that has random elements in it, the `set.seed()` function becomes very handy. For example, these two lines will always produce different output (because that is the whole point of random number generators):

```
> sample(1:10, 5)
[1] 6 9 2 7 10
> sample(1:10, 5)
[1] 7 6 1 2 10
```

These two will also produce different outputs:

```
> rnorm(5)
[1] 0.4874291 0.7383247 0.5757814 -0.3053884 1.5117812
> rnorm(5)
[1] 0.38984324 -0.62124058 -2.21469989 1.12493092 -0.04493361
```

However, if we set the seed to something identical in both cases (most people use 1 for simplicity), we get two identical samples:

```
> set.seed(1)
> sample(letters, 2)
[1] "g" "j"
> set.seed(1)
> sample(letters, 2)
[1] "g" "j"
```

and same with, say, `rexp()` draws:

```
> set.seed(1)
> rexp(5)
[1] 0.7551818 1.1816428 0.1457067 0.1397953 0.4360686
> set.seed(1)
> rexp(5)
[1] 0.7551818 1.1816428 0.1457067 0.1397953 0.4360686
```

Chapter 38: Parallel processing

Section 38.1: Parallel processing with parallel package

The base package `parallel` allows parallel computation through forking, sockets, and random-number generation.

Detect the number of cores present on the localhost:

```
parallel::detectCores(all.tests = FALSE, logical = TRUE)
```

Create a cluster of the cores on the localhost:

```
parallelCluster <- parallel::makeCluster(parallel::detectCores())
```

First, a function appropriate for parallelization must be created. Consider the `mtcars` dataset. A regression on `mpg` could be improved by creating a separate regression model for each level of `cyl`.

```
data <- mtcars
yfactor <- 'cyl'
zlevels <- sort(unique(data[[yfactor]]))
datay <- data[, 1]
dataz <- data[, 2]
datax <- data[, 3:11]

fitmodel <- function(zlevel, datax, datay, dataz) {
  glm.fit(x = datax[dataz == zlevel,], y = datay[dataz == zlevel])
}
```

Create a function that can loop through all the possible iterations of `zlevels`. This is still in serial, but is an important step as it determines the exact process that will be parallelized.

```
fitmodel <- function(zlevel, datax, datay, dataz) {
  glm.fit(x = datax[dataz == zlevel,], y = datay[dataz == zlevel])
}

for (zlevel in zlevels) {
  print("*****")
  print(zlevel)
  print(fitmodel(zlevel, datax, datay, dataz))
}
```

Curry this function:

```
worker <- function(zlevel) {
  fitmodel(zlevel, datax, datay, dataz)
}
```

Parallel computing using `parallel` cannot access the global environment. Luckily, each function creates a local environment `parallel` can access. Creation of a wrapper function allows for parallelization. The function to be applied also needs to be placed within the environment.

```
wrapper <- function(datax, datay, dataz) {
  # force evaluation of all parameters not supplied by parallelization apply
  force(datax)
```

```

force(datay)
force(dataz)
# these variables are now in an environment accessible by parallel function

# function to be applied also in the environment
fitmodel <- function(zlevel, datax, datay, dataz) {
  glm.fit(x = datax[dataz == zlevel, ], y = datay[dataz == zlevel])
}

# calling in this environment iterating over single parameter zlevel
worker <- function(zlevel) {
  fitmodel(zlevel, datax, datay, dataz)
}
return(worker)
}

```

Now create a cluster and run the wrapper function.

```

parallelcluster <- parallel::makeCluster(parallel::detectCores())
models <- parallel::parLapply(parallelcluster, zlevels,
                           wrapper(datax, datay, dataz))

```

Always stop the cluster when finished.

```
parallel::stopCluster(parallelcluster)
```

The `parallel` package includes the entire `apply()` family, prefixed with `par`.

Section 38.2: Parallel processing with `foreach` package

The `foreach` package brings the power of parallel processing to R. But before you want to use multi core CPUs you have to assign a multi core cluster. The `doSNOW` package is one possibility.

A simple use of the `foreach` loop is to calculate the sum of the square root and the square of all numbers from 1 to 100000.

```

library(foreach)
library(doSNOW)

cl <- makeCluster(5, type = "SOCK")
registerDoSNOW(cl)

f <- foreach(i = 1:100000, .combine = c, .inorder = F) %dopar% {
  k <- i ** 2 + sqrt(i)
  k
}

```

The structure of the output of `foreach` is controlled by the `.combine` argument. The default output structure is a list. In the code above, `c` is used to return a vector instead. Note that a calculation function (or operator) such as `"+"` may also be used to perform a calculation and return a further processed object.

It is important to mention that the result of each `foreach`-loop is the last call. Thus, in this example `k` will be added to the result.

Parameter	Details
<code>.combine</code>	combine Function. Determines how the results of the loop are combined. Possible values are <code>c</code> , <code>cbind</code> , <code>rbind</code> , <code>"+"</code> , <code>"*"</code> ...

- .inorder if TRUE the result is ordered according to the order of the iteration variable (here `i`). If FALSE the result is not ordered. This can have positive effects on computation time.
- .packages for functions which are provided by any package except base, like e.g. `mass`, `randomForest` or else, you have to provide these packages with `c("mass", "randomForest")`

Section 38.3: Random Number Generation

A major problem with parallelization is the use of RNG as seeds. Random numbers by the number are iterated by the number of operations from either the start of the session or the most recent `set.seed()`. Since parallel processes arise from the same function, it can use the same seed, possibly causing identical results! Calls will run in serial on the different cores, provide no advantage.

A set of seeds must be generated and sent to each parallel process. This is automatically done in some packages (`parallel`, `snow`, etc.), but must be explicitly addressed in others.

```
s <- seed
for (i in 1:numofcores) {
  s <- nextRNGStream(s)
  # send s to worker i as .Random.seed
}
```

Seeds can also be set for reproducibility.

```
clusterSetRNGStream(cl = parallelcluster, iseed)
```

Section 38.4: mparallelDo

The `mparallelDo` package allows for the evaluation of R code asynchronously on Unix-alike (e.g. Linux and MacOSX) operating systems. The underlying philosophy of the package is aligned with the needs of exploratory data analysis rather than coding. For coding asynchrony, consider the `future` package.

Example

Create data

```
data(ToothGrowth)
```

Trigger `mparallelDo` to perform analysis on a fork

```
mparallelDo({glm(len ~ supp * dose, data=ToothGrowth)}, "interactionPredictorModel")
```

Do other things, e.g.

```
binaryPredictorModel <- glm(len ~ supp, data=ToothGrowth)
gaussianPredictorModel <- glm(len ~ dose, data=ToothGrowth)
```

The result from `mparallelDo` returns in your targetEnvironment, e.g. `.GlobalEnv`, when it is complete with a message (by default)

```
summary(interactionPredictorModel)
```

Other Examples

```
# Example of not returning a value until we return to the top level
for (i in 1:10) {
  if (i == 1) {
```

```
    mcpParallelDo({2+2}, targetValue = "output")
}
if (exists("output")) print(i)
}

# Example of getting a value without returning to the top level
for (i in 1:10) {
  if (i == 1) {
    mcpParallelDo({2+2}, targetValue = "output")
  }
  mcpParallelDoCheck()
  if (exists("output")) print(i)
}
```

Chapter 39: Subsetting

Given an R object, we may require separate analysis for one or more parts of the data contained in it. The process of obtaining these parts of the data from a given object is called **subsetting**.

Section 39.1: Data frames

Subsetting a data frame into a smaller data frame can be accomplished the same as subsetting a list.

```
> df3 <- data.frame(x = 1:3, y = c("a", "b", "c"), stringsAsFactors = FALSE)

> df3
##   x y
## 1 1 a
## 2 2 b
## 3 3 c

> df3[1]    # Subset a variable by number
##   x
## 1 1
## 2 2
## 3 3

> df3["x"]  # Subset a variable by name
##   x
## 1 1
## 2 2
## 3 3

> is.data.frame(df3[1])
## TRUE

> is.list(df3[1])
## TRUE
```

Subsetting a dataframe into a column vector can be accomplished using double brackets [[]] or the dollar sign operator \$.

```
> df3[[2]]    # Subset a variable by number using [[ ]]
## [1] "a" "b" "c"

> df3[["y"]]  # Subset a variable by name using [[ ]]
## [1] "a" "b" "c"

> df3$x      # Subset a variable by name using $
## [1] 1 2 3

> typeof(df3$x)
## "integer"

> is.vector(df3$x)
## TRUE
```

Subsetting a data as a two dimensional matrix can be accomplished using i and j terms.

```
> df3[1, 2]    # Subset row and column by number
## [1] "a"
```

```

> df3[1, "y"] # Subset row by number and column by name
## [1] "a"

> df3[2, ]      # Subset entire row by number
##   x y
## 2 2 b

> df3[, 1]      # Subset all first variables
## [1] 1 2 3

> df3[, 1, drop = FALSE]
##   x
## 1 1
## 2 2
## 3 3

```

Note: Subsetting by *j* (column) alone simplifies to the variable's own type, but subsetting by *i* alone returns a **data.frame**, as the different variables may have different types and classes. Setting the **drop** parameter to FALSE keeps the data frame.

```

> is.vector(df3[, 2])
## TRUE

> is.data.frame(df3[2, ])
## TRUE

> is.data.frame(df3[, 2, drop = FALSE])
## TRUE

```

Section 39.2: Atomic vectors

Atomic vectors (which excludes lists and expressions, which are also vectors) are subset using the [operator:

```

# create an example vector
v1 <- c("a", "b", "c", "d")

# select the third element
v1[3]
## [1] "c"

```

The [operator can also take a vector as the argument. For example, to select the first and third elements:

```

v1 <- c("a", "b", "c", "d")

v1[c(1, 3)]
## [1] "a" "c"

```

Some times we may require to omit a particular value from the vector. This can be achieved using a negative sign(-) before the index of that value. For example, to omit the first value from v1, use v1[-1]. This can be extended to more than one value in a straight forward way. For example, v1[-c(1, 3)].

```

> v1[-1]
[1] "b" "c" "d"
> v1[-c(1, 3)]
[1] "b" "d"

```

On some occasions, we would like to know, especially, when the length of the vector is large, index of a particular

value, if it exists:

```
> v1=="c"
[1] FALSE FALSE TRUE FALSE
> which(v1=="c")
[1] 3
```

If the atomic vector has names (a `names` attribute), it can be subset using a character vector of names:

```
v <- 1:3
names(v) <- c("one", "two", "three")

v
##   one   two three
##   1     2     3

v["two"]
## two
## 2
```

The `[` operator can also be used to index atomic vectors, with differences in that it accepts a indexing vector with a length of one and strips any names present:

```
v[[c(1, 2)]]
## Error in v[[c(1, 2)]] :
##   attempt to select more than one element in vectorIndex

v[["two"]]
## [1] 2
```

Vectors can also be subset using a logical vector. In contrast to subsetting with numeric and character vectors, the logical vector used to subset has to be equal to the length of the vector whose elements are extracted, so if a logical vector `y` is used to subset `x`, i.e. `x[y]`, if `length(y) < length(x)` then `y` will be recycled to match `length(x)`:

```
v[c(TRUE, FALSE, TRUE)]
##   one three
##   1     3

v[c(FALSE, TRUE)] # recycled to 'c(FALSE, TRUE, FALSE)'
## two
## 2

v[TRUE] # recycled to 'c(TRUE, TRUE, TRUE)'
##   one   two three
##   1     2     3

v[FALSE] # handy to discard elements but save the vector's type and basic structure
## named integer(0)
```

Section 39.3: Matrices

For each dimension of an object, the `[` operator takes one argument. Vectors have one dimension and take one argument. Matrices and data frames have two dimensions and take two arguments, given as `[i, j]` where `i` is the row and `j` is the column. Indexing starts at 1.

```
## a sample matrix
mat <- matrix(1:6, nrow = 2, dimnames = list(c("row1", "row2"), c("col1", "col2", "col3")))
```

```

mat
#      col1 col2 col3
# row1    1    3    5
# row2    2    4    6

```

`mat[i, j]` is the element in the `i`-th row, `j`-th column of the matrix `mat`. For example, an `i` value of 2 and a `j` value of 1 gives the number in the second row and the first column of the matrix. Omitting `i` or `j` returns all values in that dimension.

```

mat[ , 3]
## row1 row2
##    5    6

mat[1, ]
# col1 col2 col3
#    1    3    5

```

When the matrix has row or column names (not required), these can be used for subsetting:

```

mat[ , 'col1']
# row1 row2
#    1    2

```

By default, the result of a subset will be simplified if possible. If the subset only has one dimension, as in the examples above, the result will be a one-dimensional vector rather than a two-dimensional matrix. This default can be overridden with the `drop = FALSE` argument to `:`:

```

## This selects the first row as a vector
class(mat[1, ])
# [1] "integer"

## Whereas this selects the first row as a 1x3 matrix:
class(mat[1, , drop = F])
# [1] "matrix"

```

Of course, dimensions cannot be dropped if the selection itself has two dimensions:

```

mat[1:2, 2:3] ## A 2x2 matrix
#      col2 col3
# row1    3    5
# row2    4    6

```

Selecting individual matrix entries by their positions

It is also possible to use a Nx2 matrix to select N individual elements from a matrix (like how a coordinate system works). If you wanted to extract, in a vector, the entries of a matrix in the (1st `row`, 1st `column`), (1st `row`, 3rd `column`), (2nd `row`, 3rd `column`), (2nd `row`, 1st `column`) this can be done easily by creating a index matrix with those coordinates and using that to subset the matrix:

```

mat
#      col1 col2 col3
# row1    1    3    5
# row2    2    4    6

ind = rbind(c(1, 1), c(1, 3), c(2, 3), c(2, 1))
ind
#      [,1] [,2]

```

```

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

mat[ind]
# [1] 1 5 6 2

```

In the above example, the 1st column of the `ind` matrix refers to rows in `mat`, the 2nd column of `ind` refers to columns in `mat`.

Section 39.4: Lists

A list can be subset with `[::]`:

```

l1 <- list(c(1, 2, 3), 'two' = c("a", "b", "c"), list(10, 20))
l1
## [[1]]
## [1] 1 2 3
##
## $two
## [1] "a" "b" "c"
##
## [[3]]
## [[3]][[1]]
## [1] 10
##
## [[3]][[2]]
## [1] 20

l1[1]
## [[1]]
## [1] 1 2 3

l1['two']
## $two
## [1] "a" "b" "c"

l1[[2]]
## [1] "a" "b" "c"

l1[[['two']]]
## [1] "a" "b" "c"

```

Note the result of `l1[2]` is still a list, as the `[` operator selects elements of a list, returning a smaller list. The `[[` operator extracts list elements, returning an object of the type of the list element.

Elements can be indexed by number or a character string of the name (if it exists). Multiple elements can be selected with `[` by passing a vector of numbers or strings of names. Indexing with a vector of `length > 1` in `[` and `[[` returns a "list" with the specified elements and a recursive subset (if available), *respectively*:

```

l1[c(3, 1)]
## [[1]]
## [[1]][[1]]
## [1] 10
##
## [[1]][[2]]
## [1] 20
##

```

```
##  
## [2]  
## [1] 1 2 3
```

Compared to:

```
11[[c(3, 1)]]  
## [1] 10
```

which is equivalent to:

```
11[[3]][[1]]  
## [1] 10
```

The \$ operator allows you to select list elements solely by name, but unlike [and [[, does not require quotes. As an infix operator, \$ can only take a single name:

```
l1$two  
## [1] "a" "b" "c"
```

Also, the \$ operator allows for partial matching by default:

```
l1$t  
## [1] "a" "b" "c"
```

in contrast with [[where it needs to be specified whether partial matching is allowed:

```
11[["t"]]  
## NULL  
11[["t", exact = FALSE]]  
## [1] "a" "b" "c"
```

Setting `options(warnPartialMatchDollar = TRUE)`, a "warning" is given when partial matching happens with \$:

```
l1$t  
## [1] "a" "b" "c"  
## Warning message:  
## In l1$t : partial match of 't' to 'two'
```

Section 39.5: Vector indexing

For this example, we will use the vector:

```
> x <- 11:20  
> x  
[1] 11 12 13 14 15 16 17 18 19 20
```

R vectors are 1-indexed, so for example `x[1]` will return 11. We can also extract a sub-vector of x by passing a vector of indices to the bracket operator:

```
> x[c(2, 4, 6)]  
[1] 12 14 16
```

If we pass a vector of negative indices, R will return a sub-vector with the specified indices excluded:

```
> x[c(-1, -3)]
[1] 12 14 15 16 17 18 19 20
```

We can also pass a boolean vector to the bracket operator, in which case it returns a sub-vector corresponding to the coordinates where the indexing vector is TRUE:

```
> x[c(rep(TRUE, 5), rep(FALSE, 5))]
[1] 11 12 13 14 15 16
```

If the indexing vector is shorter than the length of the array, then it will be repeated, as in:

```
> x[c(TRUE, FALSE)]
[1] 11 13 15 17 19
> x[c(TRUE, FALSE, FALSE)]
[1] 11 14 17 20
```

Section 39.6: Other objects

The [and [[] operators are primitive functions that are generic. This means that any *object* in R (specifically `isTRUE(is.object(x))` --i.e. has an explicit "class" attribute) can have its own specified behaviour when subsetted; i.e. has its own *methods* for [and/or [[]].

For example, this is the case with "data.frame" (`is.object(iris)`) objects where `[.data.frame` and `[[.data.frame` methods are defined and they are made to exhibit both "matrix"-like and "list"-like subsetting. With forcing an error when subsetting a "data.frame", we see that, actually, a function `[.data.frame` was called when we -just- used [.

```
iris[invalidArgument, ]
## Error in `[.data.frame`(iris, invalidArgument, ) :
##   object 'invalidArgument' not found
```

Without further details on the current topic, an example[method:

```
x = structure(1:5, class = "myClass")
x[c(3, 2, 4)]
## [1] 3 2 4
'[.myClass' = function(x, i) cat(sprintf("We'd expect '%s[%s]' to be returned but this a custom '[' method and should have a `? [.myClass` help page for its behaviour\n", deparse(substitute(x)), deparse(substitute(i))))
x[c(3, 2, 4)]
## We'd expect 'x[c(3, 2, 4)]' to be returned but this a custom '[' method and should have a `? [.myClass` help page for its behaviour
## NULL
```

We can overcome the method dispatching of [by using the equivalent non-generic `.subset` (and `.subset2` for [[]). This is especially useful and efficient when programming our own "class"es and want to avoid work-arounds (like `unclass(x)`) when computing on our "class"es efficiently (avoiding method dispatch and copying objects):

```
.subset(x, c(3, 2, 4))
## [1] 3 2 4
```

Section 39.7: Elementwise Matrix Operations

Let A and B be two matrices of same dimension. The operators +,-,/,*,^ when used with matrices of same dimension perform the required operations on the corresponding elements of the matrices and return a new

matrix of the same dimension. These operations are usually referred to as element-wise operations.

Operator A op B	Meaning
+	A + B Addition of corresponding elements of A and B
-	A - B Subtracts the elements of B from the corresponding elements of A
/	A / B Divides the elements of A by the corresponding elements of B
*	A * B Multiplies the elements of A by the corresponding elements of B
^	A^(−1) For example, gives a matrix whose elements are reciprocals of A

For "true" matrix multiplication, as seen in *Linear Algebra*, use `%*%`. For example, multiplication of A with B is: A `%*%` B. The dimensional requirements are that the `ncol()` of A be the same as `nrow()` of B

Some Functions used with Matrices

Function	Example	Purpose
<code>nrow()</code>	<code>nrow(A)</code>	determines the number of rows of A
<code>ncol()</code>	<code>ncol(A)</code>	determines the number of columns of A
<code>rownames()</code>	<code>rownames(A)</code>	prints out the row names of the matrix A
<code>colnames()</code>	<code>colnames(A)</code>	prints out the column names of the matrix A
<code>rowMeans()</code>	<code>rowMeans(A)</code>	computes means of each row of the matrix A
<code>colMeans()</code>	<code>colMeans(A)</code>	computes means of each column of the matrix A
<code>upper.tri()</code>	<code>upper.tri(A)</code>	returns a vector whose elements are the upper triangular matrix of square matrix A
<code>lower.tri()</code>	<code>lower.tri(A)</code>	returns a vector whose elements are the lower triangular matrix of square matrix A
<code>det()</code>	<code>det(A)</code>	results in the determinant of the matrix A
<code>solve()</code>	<code>solve(A)</code>	results in the inverse of the non-singular matrix A
<code>diag()</code>	<code>diag(A)</code>	returns a diagonal matrix whose off-diagonal elements are zeros and diagonals are the same as that of the square matrix A
<code>t()</code>	<code>t(A)</code>	returns the transpose of the matrix A
<code>eigen()</code>	<code>eigen(A)</code>	returns the eigenvalues and eigenvectors of the matrix A
<code>is.matrix()</code>	<code>is.matrix(A)</code>	returns TRUE or FALSE depending on whether A is a matrix or not.
<code>as.matrix()</code>	<code>as.matrix(x)</code>	creates a matrix out of the vector x

Chapter 40: Debugging

Section 40.1: Using debug

You can set any function for debugging with `debug`.

```
debug(mean)  
mean(1:3)
```

All subsequent calls to the function will enter debugging mode. You can disable this behavior with `undebbug`.

```
undebbug(mean)  
mean(1:3)
```

If you know you only want to enter the debugging mode of a function once, consider the use of `debugonce`.

```
debugonce(mean)  
mean(1:3)  
mean(1:3)
```

Section 40.2: Using browser

The `browser` function can be used like a breakpoint: code execution will pause at the point it is called. Then user can then inspect variable values, execute arbitrary R code and step through the code line by line.

Once `browser()` is hit in the code the interactive interpreter will start. Any R code can be run as normal, and in addition the following commands are present,

Command	Meaning
c	Exit browser and continue program
f	Finish current loop or function \
n	Step Over (evaluate next statement, stepping over function calls)
s	Step Into (evaluate next statement, stepping into function calls)
where	Print stack trace
r	Invoke "resume" restart
Q	Exit browser and quit

For example we might have a script like,

```
toDebug <- function() {  
  a = 1  
  b = 2  
  
  browser()  
  
  for(i in 1:100) {  
    a = a * b  
  }  
}  
  
toDebug()
```

When running the above script we initially see something like,

```
Called from: toDebug
Browser[1]>
```

We could then interact with the prompt as so,

```
Called from: toDebug
Browser[1]> a
[1] 1
Browser[1]> b
[1] 2
Browse[1]> n
debug at #7: for (i in 1:100) {
  a = a * b
}
Browse[2]> n
debug at #8: a = a * b
Browse[2]> a
[1] 1
Browse[2]> n
debug at #8: a = a * b
Browse[2]> a
[1] 2
Browse[2]> Q
```

`browser()` can also be used as part of a functional chain, like so:

```
mtcars %>% group_by(cyl) %>% {browser()}
```

Chapter 41: Installing packages

Parameter	Details
pkgs	character vector of the names of packages. If repos = NULL, a character vector of file paths.
lib	character vector giving the library directories where to install the packages.
repos	character vector, the base URL(s) of the repositories to use, can be NULL to install from local files
method	download method
destdir	directory where downloaded packages are stored
dependencies	logical indicating whether to also install uninstalled packages which these packages depend on/link to/import/suggest (and so on recursively). Not used if repos = NULL.
...	Arguments to be passed to 'download.file' or to the functions for binary installs on OS X and Windows.

Section 41.1: Install packages from GitHub

To install packages directly from GitHub use the devtools package:

```
library(devtools)
install_github("authorName/repositoryName")
```

To install ggplot2 from github:

```
devtools::install_github("tidyverse/ggplot2")
```

The above command will install the version of ggplot2 that corresponds to the *master* branch. To install from a different branch of a repository use the ref argument to provide the name of the branch. For example, the following command will install the dev_general branch of the googleway package.

```
devtools::install_github("SymbolixAU/googleway", ref = "dev_general")
```

Another option is to use the ghit package. It provides a lightweight alternative for installing packages from github:

```
install.packages("ghit")
ghit::install_github("google/CausalImpact")
```

To install a package that is in a **private** repository on Github, generate a **personal access token** at <http://www.github.com/settings/tokens/> (See ?install_github for documentation on the same). Follow these steps:

1. `install.packages(c("curl", "httr"))`
2. `config = httr::config(ssl_verifypeer = FALSE)`
3. `install.packages("RCurl")`
`options(RCurlOptions = c(getOption("RCurlOptions"), ssl.verifypeer = FALSE, ssl.verifyhost = FALSE))`
4. `getOption("RCurlOptions")`

You should see the following:

```
ssl.verifypeer ssl.verifyhost
```

```
FALSE          FALSE
```

5. `library(httr)`

```
set_config(config(ssl_verifypeer = 0L))
```

This prevents the common error: "Peer certificate cannot be authenticated with given CA certificates"

- Finally, use the following command to install your package seamlessly

```
install_github("username/package_name", auth_token="abc")
```

Alternatively, set an environment variable GITHUB_PAT, using

```
Sys.setenv(GITHUB_PAT = "access_token")
devtools::install_github("organisation/package_name")
```

The PAT generated in Github is only visible once, i.e., when created initially, so its prudent to save that token in .Rprofile. This is also helpful if the organisation has many private repositories.

Section 41.2: Download and install packages from repositories

Packages are collections of R functions, data, and compiled code in a [well-defined format](#). Public (and private) repositories are used to host collections of R packages. The largest collection of R packages is available from CRAN.

Using CRAN

A package can be installed from [CRAN](#) using following code:

```
install.packages("dplyr")
```

Where "dplyr" is referred to as a character vector.

More than one packages can be installed in one go by using the combine function `c()` and passing a series of character vector of package names:

```
install.packages(c("dplyr", "tidy", "ggplot2"))
```

In some cases, `install.packages` may prompt for a CRAN mirror or fail, depending on the value of `getOption("repos")`. To prevent this, specify a [CRAN mirror](#) as repos argument:

```
install.packages("dplyr", repos = "https://cloud.r-project.org/")
```

Using the repos argument it is also possible to install from other repositories. For complete information about all the available options, run `?install.packages`.

Most packages require functions, which were implemented in other packages (e.g. the package `data.table`). In order to install a package (or multiple packages) with all the packages, which are used by this given package, the argument dependencies should be set to TRUE):

```
install.packages("data.table", dependencies = TRUE)
```

Using Bioconductor

[Bioconductor](#) hosts a substantial collection of packages related to Bioinformatics. They provide their own package management centred around the `biocLite` function:

```
## Try http:// if https:// URLs are not supported
source("https://bioconductor.org/biocLite.R")
biocLite()
```

By default this installs a subset of packages that provide the most commonly used functionality. Specific packages can be installed by passing a vector of package names. For example, to install RImmPort from Bioconductor:

```
source("https://bioconductor.org/biocLite.R")
biocLite("RImmPort")
```

Section 41.3: Install package from local source

To install package from local source file:

```
install.packages(path_to_source, repos = NULL, type="source")

install.packages("~/Downloads/dplyr-master.zip", repos=NULL, type="source")
```

Here, path_to_source is absolute path of local source file.

Another command that opens a window to choose downloaded zip or tar.gz source files is:

```
install.packages(file.choose(), repos=NULL)
```

Another possible way is using the GUI based RStudio:

Step 1: Go to **Tools**.

Step 2: Go to **Install Packages**.

Step 3: In the *Install From* set it as **Package Archive File (.zip; .tar.gz)**

Step 4: Then *Browse* find your package file (say crayon_1.3.1.zip) and *after some time (after it shows the Package path and file name in the Package Archive tab)*

Another way to install R package from local source is using `install_local()` function from `devtools` package.

```
library(devtools)
install_local("~/Downloads/dplyr-master.zip")
```

Section 41.4: Install local development version of a package

While working on the development of an R package it is often necessary to install the latest version of the package. This can be achieved by first building a source distribution of the package (on the command line)

```
R CMD build my_package
```

and then installing it in R. Any running R sessions with previous version of the package loaded will need to reload it.

```
unloadNamespace("my_package")
library(my_package)
```

A more convenient approach uses the `devtools` package to simplify the process. In an R session with the working directory set to the package directory

```
devtools::install()
```

will build, install and reload the package.

Section 41.5: Using a CLI package manager -- basic pacman usage

pacman is a simple package manager for R.

pacman allows a user to compactly load all desired packages, installing any which are missing (and their dependencies), with a single command, `p_load`. pacman does not require the user to type quotation marks around a package name. Basic usage is as follows:

```
p_load(data.table, dplyr, ggplot2)
```

The only package requiring a `library`, `require`, or `install.packages` statement with this approach is pacman itself:

```
library(pacman)
p_load(data.table, dplyr, ggplot2)
```

or, equally valid:

```
pacman::p_load(data.table, dplyr, ggplot2)
```

In addition to saving time by requiring less code to manage packages, pacman also facilitates the construction of reproducible code by installing any needed packages if and only if they are not already installed.

Since you may not be sure if pacman is installed in the library of a user who will use your code (or by yourself in future uses of your own code) a best practice is to include a conditional statement to install pacman if it is not already loaded:

```
if(!require(pacman)) install.packages("pacman")
pacman::p_load(data.table, dplyr, ggplot2)
```

Chapter 42: Inspecting packages

Packages build on base R. This document explains how to inspect installed packages and their functionality. Related Docs: [Installing packages](#)

Section 42.1: View Package Version

Conditions: package should be at least installed. If not loaded in the current session, not a problem.

```
## Checking package version which was installed at past or  
## installed currently but not loaded in the current session  
  
packageVersion("seqinr")  
# [1] '3.3.3'  
packageVersion("RWeka")  
# [1] '0.4.29'
```

Section 42.2: View Loaded packages in Current Session

To check the list of loaded packages

```
search()
```

OR

```
(.packages())
```

Section 42.3: View package information

To retrieve information about dplyr package and its functions' descriptions:

```
help(package = "dplyr")
```

No need to load the package first.

Section 42.4: View package's built-in data sets

To see built-in data sets from package dplyr

```
data(package = "dplyr")
```

No need to load the package first.

Section 42.5: List a package's exported functions

To get the list of functions within package dplyr, we first must load the package:

```
library(dplyr)  
ls("package:dplyr")
```

Chapter 43: Creating packages with devtools

This topic will cover the creation of R packages from scratch with the `devtools` package.

Section 43.1: Creating and distributing packages

This is a *compact guide* about how to quickly create an R package from your code. Exhaustive documentations will be linked when available and should be read if you want a deeper knowledge of the situation. See *Remarks* for more resources.

The directory where your code stands will be referred as `./`, and all the commands are meant to be executed from a R prompt in this folder.

Creation of the documentation

The documentation for your code has to be in a format which is very similar to LaTeX.

However, we will use a tool named `roxygen` in order to simplify the process:

```
install.packages("devtools")
library("devtools")
install.packages("roxygen2")
library("roxygen2")
```

The full man page for `roxygen` is available [here](#). It is very similar to `doxygen`.

Here is a practical sample about how to document a function with `roxygen`:

```
#' Increment a variable.
#'
#' Note that the behavior of this function
#' is undefined if `x` is not of class `numeric`.
#'
#' @export
#' @author another guy
#' @name Increment Function
#' @title increment
#'
#' @param x    Variable to increment
#' @return     `x` incremented of 1
#'
#' @seealso   `other_function`
#'
#' @examples
#' increment(3)
#' > 4
increment <- function(x) {
  return (x+1)
}
```

And [here will be the result](#).

It is also recommended to create a vignette (see the topic *Creating vignettes*), which is a full guide about your package.

Construction of the package skeleton

Assuming that your code is written for instance in files `./script1.R` and `./script2.R`, launch the following command in order to create the file tree of your package:

```
package.skeleton(name="MyPackage", code_files=c("script1.R", "script2.R"))
```

Then delete all the files in `./MyPackage/man/`. You have now to compile the documentation:

```
roxygenize("MyPackage")
```

You should also generate a reference manual from your documentation using `R CMD Rd2pdf MyPackage` from a *command prompt* started in `./`.

Edition of the package properties

1. Package description

Modify `./MyPackage/DESCRIPTION` according to your needs. The fields `Package`, `Version`, `License`, `Description`, `Title`, `Author` and `Maintainer` are mandatory, the other are optional.

If your package depends on others packages, specify them in a field named `Depends` (R version < 3.2.0) or `Imports` (R version > 3.2.0).

2. Optional folders

Once you launched the skeleton build, `./MyPackage/` only had `R/` and `man/` subfolders. However, it can have some others:

- `data/`: here you can place the data that your library needs and that isn't code. It must be saved as dataset with the `.RData` extension, and you can load it at runtime with `data()` and `load()`
- `tests/`: all the code files in this folder will be ran at install time. If there is any error, the installation will fail.
- `src/`: for C/C++/Fortran source files you need (using Rcpp...).
- `exec/`: for other executables.
- `misc/`: for barely everything else.

Finalization and build

You can delete `./MyPackage/Read-and-delete-me`.

As it is now, your package is ready to be installed.

You can install it with `devtools::install("MyPackage")`.

To build your package as a source tarball, you need to execute the following command, from a *command prompt* in `./`: `R CMD build MyPackage`

Distribution of your package

Through Github

Simply create a new repository called `MyPackage` and upload everything in `MyPackage/` to the master branch. Here is [an example](#).

Then anyone can install your package from github with devtools:

```
install_package("MyPackage", "your_github_username")
```

Through CRAN

Your package needs to comply to the [CRAN Repository Policy](#). Including but not limited to: your package must be cross-platforms (except some very special cases), it should pass the R CMD check test.

Here is the [submission form](#). You must upload the source tarball.

Section 43.2: Creating vignettes

A vignette is a long-form guide to your package. Function documentation is great if you know the name of the function you need, but it's useless otherwise. A vignette is like a book chapter or an academic paper: it can describe the problem that your package is designed to solve, and then show the reader how to solve it.

Vignettes will be created entirely in markdown.

Requirements

- Rmarkdown: `install.packages("rmarkdown")`
- [Pandoc](#)

Vignette creation

```
devtools::use_vignette("MyVignette", "MyPackage")
```

You can now edit your vignette at `./vignettes/MyVignette.Rmd`.

The text in your vignette is formatted as [Markdown](#).

The only addition to the original Markdown, is a tag that takes R code, runs it, captures the output, and translates it into formatted Markdown:

```
```{r}
Add two numbers together
add <- function(a, b) a + b
add(10, 20)
````
```

Will display as:

```
# Add two numbers together
add <- function(a, b) a + b
add(10, 20)
## [1] 30
```

Thus, all the packages you will use in your vignettes must be listed as dependencies in `./DESCRIPTION`.

Chapter 44: Using pipe assignment in your own package %<>%: How to ?

In order to use the pipe in a user-created package, it must be listed in the NAMESPACE like any other function you choose to import.

Section 44.1: Putting the pipe in a utility-functions file

One option for doing this is to export the pipe from within the package itself. This may be done in the 'traditional' zzz.R or utils.R files that many packages utilise for useful little functions that are not exported as part of the package. For example, putting:

```
#' Pipe operator
#'
#' @name %>%
#' @rdname pipe
#' @keywords internal
#' @export
#' @importFrom magrittr %>%
#' @usage lhs \%>\% rhs
NULL
```

Chapter 45: Arima Models

Section 45.1: Modeling an AR1 Process with Arima

We will model the process

```
#Load the forecast package
library(forecast)

#Generate an AR1 process of length n (from Cowpertwait & Meltcalfe)
# Set up variables
set.seed(1234)
n <- 1000
x <- matrix(0,1000,1)
w <- rnorm(n)

# loop to create x
for (t in 2:n) x[t] <- 0.7 * x[t-1] + w[t]
plot(x,type='l')
```



We will fit an Arima model with autoregressive order 1, 0 degrees of differencing, and an MA order of 0.

```
#Fit an AR1 model using Arima
fit <- Arima(x, order = c(1, 0, 0))
summary(fit)
# Series: x
# ARIMA(1,0,0) with non-zero mean
#
# Coefficients:
#          ar1  intercept
#        0.7040   -0.0842
# s.e.  0.0224    0.1062
#
# sigma^2 estimated as 0.9923: log likelihood=-1415.39
# AIC=2836.79  AICc=2836.81  BIC=2851.51
#
# Training set error measures:
#           ME      RMSE       MAE      MPE      MAPE      MASE       ACF1
# Training set -8.369365e-05 0.9961194 0.7835914 Inf  Inf  0.91488  0.02263595
# Verify that the model captured the true AR parameter
```

Notice that our coefficient is close to the true value from the generated data

```
fit$coef[1]
#      ar1
# 0.7040085

#Verify that the model eliminates the autocorrelation
acf(x)
```

```
acf(fit$resid)
```

```

#Forecast 10 periods
fcst <- forecast(fit, h = 100)
fcst
  Point Forecast     Lo 80     Hi 80     Lo 95     Hi 95
1001  0.282529070 -0.9940493 1.559107 -1.669829 2.234887
1002  0.173976408 -1.3872262 1.735179 -2.213677 2.561630
1003  0.097554408 -1.5869850 1.782094 -2.478726 2.673835
1004  0.043752667 -1.6986831 1.786188 -2.621073 2.708578
1005  0.005875783 -1.7645535 1.776305 -2.701762 2.713514
...
#Call the point predictions
fcst$mean
# Time Series:
# Start = 1001
# End = 1100
# Frequency = 1
[1] 0.282529070 0.173976408 0.097554408 0.043752667 0.005875783 -0.020789866 -0.039562711
-0.052778954
[9] -0.062083302
...

```

```
#Plot the forecast  
plot(fcst)
```

Chapter 46: Distribution Functions

R has many built-in functions to work with probability distributions, with official docs starting at [?Distributions](#).

Section 46.1: Normal distribution

Let's use `*norm` as an example. From the documentation:

```
dnorm(x, mean = 0, sd = 1, log = FALSE)
pnorm(q, mean = 0, sd = 1, lower.tail = TRUE, log.p = FALSE)
qnorm(p, mean = 0, sd = 1, lower.tail = TRUE, log.p = FALSE)
rnorm(n, mean = 0, sd = 1)
```

So if I wanted to know the value of a standard normal distribution at 0, I would do

```
dnorm(0)
```

Which gives us `0.3989423`, a reasonable answer.

In the same way `pnorm(0)` gives `.5`. Again, this makes sense, because half of the distribution is to the left of 0.

`qnorm` will essentially do the opposite of `pnorm`. `qnorm(.5)` gives `0`.

Finally, there's the `rnorm` function:

```
rnorm(10)
```

Will generate 10 samples from standard normal.

If you want to change the parameters of a given distribution, simply change them like so

```
rnorm(10, mean=4, sd= 3)
```

Section 46.2: Binomial Distribution

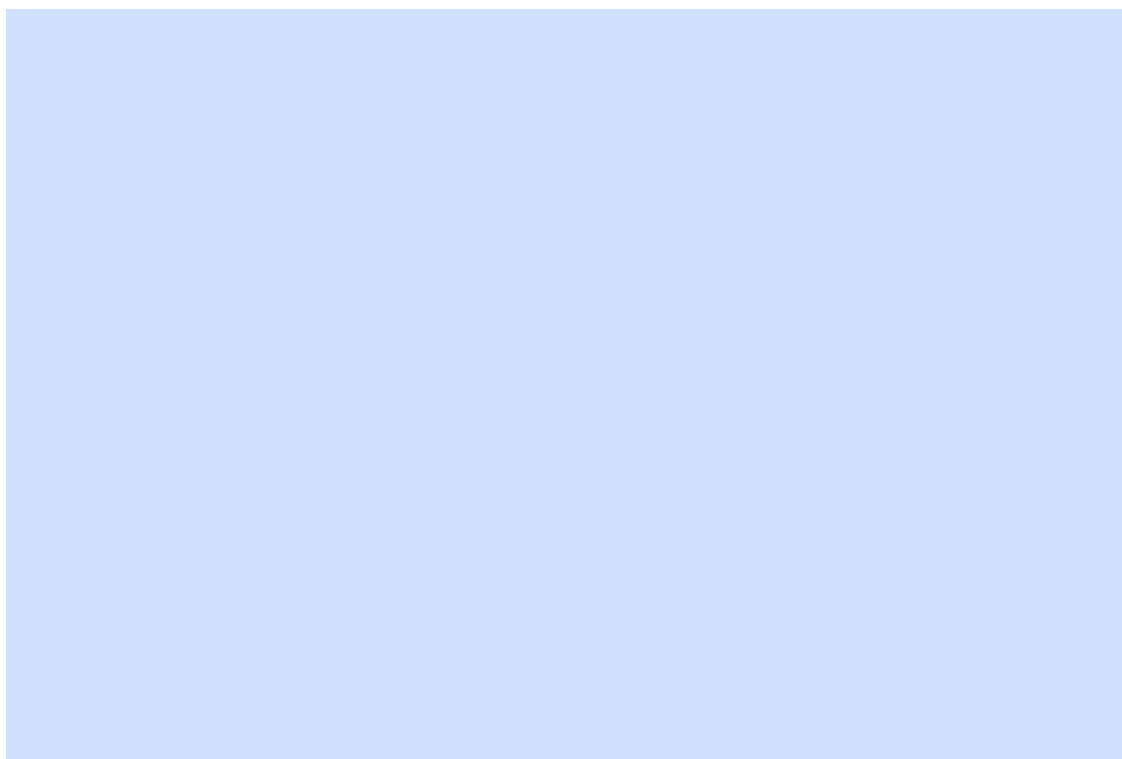
We now illustrate the functions `dbinom`, `pbinom`, `qbinom` and `rbinom` defined for *Binomial distribution*.

The `dbinom()` function gives the probabilities for various values of the binomial variable. Minimally it requires three arguments. The first argument for this function must be a vector of quantiles(the possible values of the random variable X). The second and third arguments are the defining parameters of the distribution, namely, `n`(the number of independent trials) and `p`(the probability of success in each trial). For example, for a binomial distribution with `n = 5`, `p = 0.5`, the possible values for X are `0, 1, 2, 3, 4, 5`. That is, the `dbinom(x, n, p)` function gives the probability values $P(X = x)$ for `x = 0, 1, 2, 3, 4, 5`.

```
#Binom(n = 5, p = 0.5) probabilities
> n <- 5; p<- 0.5; x <- 0:n
> dbinom(x,n,p)
[1] 0.03125 0.15625 0.31250 0.31250 0.15625 0.03125
#To verify the total probability is 1
> sum(dbinom(x,n,p))
[1] 1
>
```

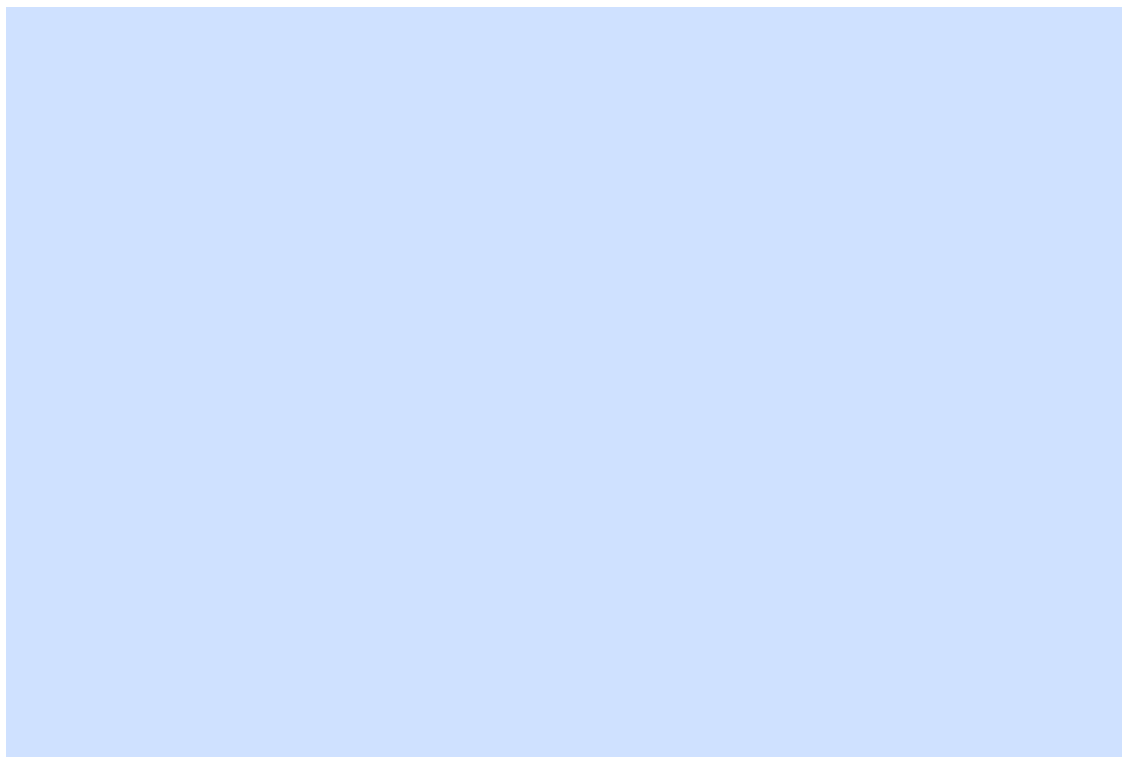
The binomial probability distribution plot can be displayed as in the following figure:

```
> x <- 0:12
> prob <- dbinom(x,12,.5)
> barplot(prob,col = "red",ylim = c(0,.2),names.arg=x,
           main="Binomial Distribution\n(n=12,p=0.5)")
```



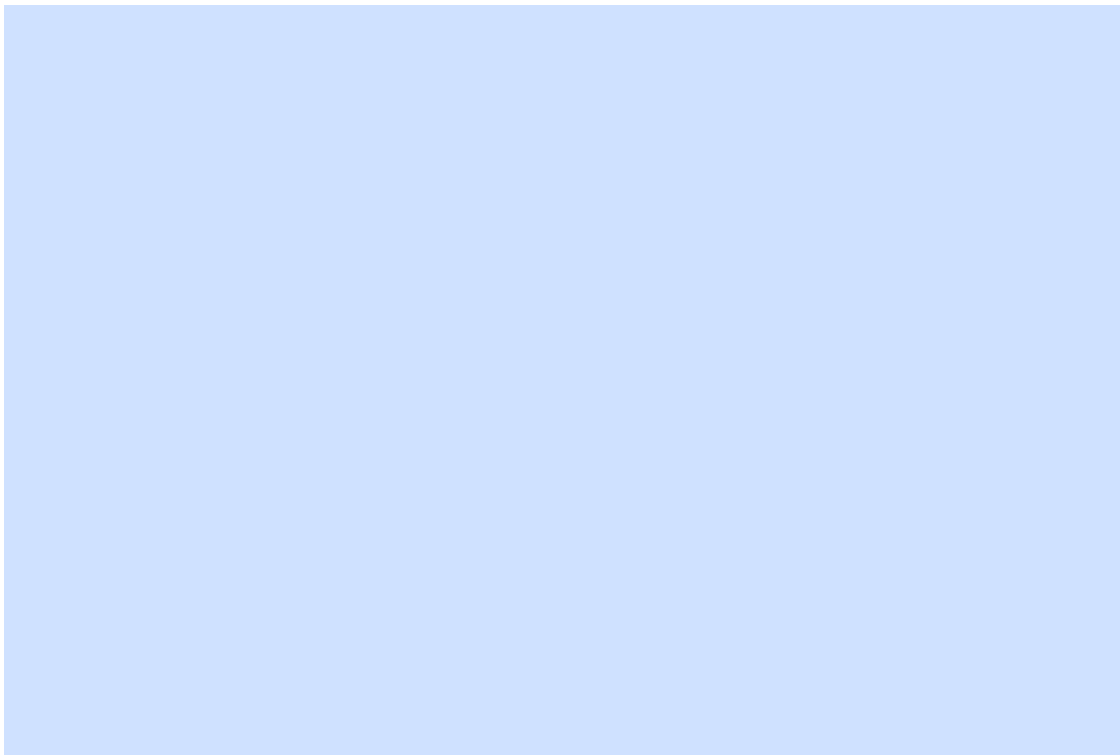
Note that the binomial distribution is symmetric when $p = 0.5$. To demonstrate that the binomial distribution is negatively skewed when p is larger than 0.5 , consider the following example:

```
> n=9; p=.7; x=0:n; prob=dbinom(x,n,p);
> barplot(prob,names.arg = x,main="Binomial Distribution\n(n=9, p=0.7)",col="lightblue")
```



When p is smaller than 0.5 the binomial distribution is positively skewed as shown below.

```
> n=9; p=.3; x=0:n; prob=dbinom(x,n,p);
> barplot(prob,names.arg = x,main="Binomial Distribution\n(n=9, p=0.3)",col="cyan")
```



We will now illustrate the usage of the cumulative distribution function `pbinom()`. This function can be used to calculate probabilities such as $P(X \leq x)$. The first argument to this function is a vector of quantiles(values of x).

```
# Calculating Probabilities
# P(X <= 2) in a Bin(n=5,p=0.5) distribution
> pbinom(2,5,0.5)
[1] 0.5
```

The above probability can also be obtained as follows:

```
# P(X <= 2) = P(X=0) + P(X=1) + P(X=2)
> sum(dbinom(0:2,5,0.5))
[1] 0.5
```

To compute, probabilities of the type: $P(a \leq X \leq b)$

```
# P(3<= X <= 5) = P(X=3) + P(X=4) + P(X=5) in a Bin(n=9,p=0.6) dist
> sum(dbinom(c(3,4,5),9,0.6))
[1] 0.4923556
>
```

Presenting the binomial distribution in the form of a table:

```
> n = 10; p = 0.4; x = 0:n;
> prob = dbinom(x,n,p)
> cdf = pbinom(x,n,p)
> distTable = cbind(x,prob,cdf)
> distTable
      x       prob       cdf
[1,]  0 0.0060466176 0.006046618
[2,]  1 0.0403107840 0.046357402
[3,]  2 0.1209323520 0.167289754
```

```
[4,] 3 0.2149908480 0.382280602
[5,] 4 0.2508226560 0.633103258
[6,] 5 0.2006581248 0.833761382
[7,] 6 0.1114767360 0.945238118
[8,] 7 0.0424673280 0.987705446
[9,] 8 0.0106168320 0.998322278
[10,] 9 0.0015728640 0.999895142
[11,] 10 0.0001048576 1.000000000
>
```

The `rbinom()` is used to generate random samples of specified sizes with a given parameter values.

```
# Simulation
> xVal<-names(table(rbinom(1000,8,.5)))
> barplot(as.vector(table(rbinom(1000,8,.5))),names.arg =xVal,
          main="Simulated Binomial Distribution\n (n=8,p=0.5)")
```



Chapter 47: Shiny

Section 47.1: Create an app

Shiny is an [R](#) package developed by [RStudio](#) that allows the creation of web pages to interactively display the results of an analysis in R.

There are two simple ways to create a Shiny app:

- in one .R file, or
- in two files: ui.R and server.R.

A Shiny app is divided into two parts:

- **ui**: A user interface script, controlling the layout and appearance of the application.
- **server**: A server script which contains code to allow the application to react.

One file

```
library(shiny)

# Create the UI
ui <- shinyUI(fluidPage(
  # Application title
  titlePanel("Hello World!")
))

# Create the server function
server <- shinyServer(function(input, output){})

# Run the app
shinyApp(ui = ui, server = server)
```

Two files

Create ui.R file

```
library(shiny)

# Define UI for application
shinyUI(fluidPage(
  # Application title
  titlePanel("Hello World!")
))
```

Create server.R file

```
library(shiny)

# Define server logic
shinyServer(function(input, output){})
```

Section 47.2: Checkbox Group

Create a group of checkboxes that can be used to toggle multiple choices independently. The server will receive the input as a character vector of the selected values.

```
library(shiny)

ui <- fluidPage(
```

```

checkboxGroupInput("checkGroup1", label = h3("This is a Checkbox group"),
  choices = list("1" = 1, "2" = 2, "3" = 3),
  selected = 1),
fluidRow(column(3, verbatimTextOutput("text_choice")))
)

server <- function(input, output){
  output$text_choice <- renderPrint({
    return(paste0("You have chosen the choice ", input$checkGroup1)))
}
shinyApp(ui = ui, server = server)

```

This is a Checkbox group

- 1
- 2
- 3

[1] "You have chosen the choice 1"

It's possible to change the settings :

- label : title
- choices : selected values
- selected : The initially selected value (NULL for no selection)
- inline : horizontal or vertical
- width

It is also possible to add HTML.

Section 47.3: Radio Button

You can create a set of radio buttons used to select an item from a list.

It's possible to change the settings :

- selected : The initially selected value (character(0) for no selection)
- inline : horizontal or vertical
- width

It is also possible to add HTML.

```

library(shiny)

ui <- fluidPage(
  radioButtons("radio",
    label = HTML('<FONT color="red"><FONT size="5pt">Welcome</FONT></FONT><br> <b>Your
favorite color is red ?</b>'),
    choices = list("TRUE" = 1, "FALSE" = 2),
    selected = 1,
    inline = T,
    width = "100%"),

```

```

fluidRow(column(3, textOutput("value")))

server <- function(input, output){
  output$value <- renderPrint({
    if(input$radio == 1){return('Great !')}
    else{return("Sorry !")}}))

shinyApp(ui = ui, server = server)

```

Welcome

Your favorite color is red ?

TRUE FALSE

[1] "Great !"

Section 47.4: Debugging

`debug()` and `debugonce()` won't work well in the context of most Shiny debugging. However, `browser()` statements inserted in critical places can give you a lot of insight into how your Shiny code is (not) working. See also: Debugging using `browser()`

Showcase mode

[Showcase mode](#) displays your app alongside the code that generates it and highlights lines of code in `server.R` as it runs them.

There are two ways to enable Showcase mode:

- Launch Shiny app with the argument `display.mode = "showcase"`, e.g., `runApp("MyApp", display.mode = "showcase")`.
- Create file called `DESCRIPTION` in your Shiny app folder and add this line in it: `DisplayMode: Showcase`.

Reactive Log Visualizer

[Reactive Log Visualizer](#) provides an interactive browser-based tool for visualizing reactive dependencies and execution in your application. To enable Reactive Log Visualizer, execute `options(shiny.reactlog=TRUE)` in R console and or add that line of code in your `server.R` file. To start Reactive Log Visualizer, hit Ctrl+F3 on Windows or Command+F3 on Mac when your app is running. Use left and right arrow keys to navigate in Reactive Log Visualizer.

Section 47.5: Select box

Create a select list that can be used to choose a single or multiple items from a list of values.

```

library(shiny)

ui <- fluidPage(
  selectInput("id_selectInput",
    label = HTML('<B><FONT size="3">What is your favorite color ?</FONT></B>'),
    multiple = TRUE,
    choices = list("red" = "red", "green" = "green", "blue" = "blue", "yellow" = "yellow"),
    selected = NULL),
  br(), br(),
  fluidRow(column(3, textOutput("text_choice"))))

```

```
server <- function(input, output){  
  output$text_choice <- renderPrint({  
    return(input$id_selectInput)})  
}  
  
shinyApp(ui = ui, server = server)
```

It's possible to change the settings :

- label : title
- choices : selected values
- selected : The initially selected value (NULL for no selection)
- multiple : TRUE or FALSE
- width
- size
- selectize: TRUE or FALSE (for use or not selectize.js, change the display)

It is also possible to add HTML.

Section 47.6: Launch a Shiny app

You can launch an application in several ways, depending on how you create your app. If your app is divided in two files `ui.R` and `server.R` or if all of your app is in one file.

1. Two files app

Your two files `ui.R` and `server.R` have to be in the same folder. You could then launch your app by running in the console the `shinyApp()` function and by passing the path of the directory that contains the Shiny app.

```
shinyApp("path_to_the_folder_containing_the_files")
```

You can also launch the app directly from Rstudio by pressing the **Run App** button that appears on Rstudio when you an `ui.R` or `server.R` file open.

Or you can simply write `runApp()` on the console if your working directory is Shiny App directory.

2. One file app

If you create your in one R file you can also launch it with the `shinyApp()` function.

- inside of your code :

```

library(shiny)

ui <- fluidPage() #Create the ui
server <- function(input, output){} #create the server

shinyApp(ui = ui, server = server) #run the App

```

- in the console by adding path to a .R file containing the Shiny application with the parameter appFile:

```
shinyApp(appFile="path_to_my_R_file_containig_the_app")
```

Section 47.7: Control widgets

| Function | Widget |
|--------------------|--|
| actionButton | Action Button |
| checkboxGroupInput | A group of check boxes |
| checkboxInput | A single check box |
| dateInput | A calendar to aid date selection |
| dateRangeInput | A pair of calendars for selecting a date range |
| fileInput | A file upload control wizard |
| helpText | Help text that can be added to an input form |
| numericInput | A field to enter numbers |
| radioButtons | A set of radio buttons |
| selectInput | A box with choices to select from |
| sliderInput | A slider bar |
| submitButton | A submit button |
| textInput | A field to enter text |

```

library(shiny)

# Create the UI
ui <- shinyUI(fluidPage(
  titlePanel("Basic widgets"),

  fluidRow(
    column(3,
      h3("Buttons"),
      actionButton("action", label = "Action"),
      br(),
      br(),
      submitButton("Submit")),

    column(3,
      h3("Single checkbox"),
      checkboxInput("checkbox", label = "Choice A", value = TRUE)),

    column(3,
      checkboxGroupInput("checkGroup",
        label = h3("Checkbox group"),
        choices = list("Choice 1" = 1,
                      "Choice 2" = 2, "Choice 3" = 3),
        selected = 1)),

    column(3,
      dateInput("date",
                label = h3("Date input")),

```

```

        value = "2014-01-01"))
),
fluidRow(
  column(3,
    dateRangeInput("dates", label = h3("Date range"))),
  column(3,
    fileInput("file", label = h3("File input"))),
  column(3,
    h3("Help text"),
    helpText("Note: help text isn't a true widget,",
            "but it provides an easy way to add text to",
            "accompany other widgets.")),
  column(3,
    numericInput("num",
      label = h3("Numeric input"),
      value = 1))
),
fluidRow(
  column(3,
    radioButtons("radio", label = h3("Radio buttons"),
      choices = list("Choice 1" = 1, "Choice 2" = 2,
                     "Choice 3" = 3), selected = 1)),
  column(3,
    selectInput("select", label = h3("Select box"),
      choices = list("Choice 1" = 1, "Choice 2" = 2,
                     "Choice 3" = 3), selected = 1)),
  column(3,
    sliderInput("slider1", label = h3("Sliders"),
      min = 0, max = 100, value = 50),
    sliderInput("slider2", "",
      min = 0, max = 100, value = c(25, 75)))
),
column(3,
 textInput("text", label = h3("Text input"),
            value = "Enter text..."))
)
))

# Create the server function
server <- shinyServer(function(input, output){})

# Run the app
shinyApp(ui = ui, server = server)

```

Chapter 48: spatial analysis

Section 48.1: Create spatial points from XY data set

When it comes to geographic data, R shows to be a powerful tool for data handling, analysis and visualisation.

Often, spatial data is available as an XY coordinate data set in tabular form. This example will show how to create a spatial data set from an XY data set.

The packages `rgdal` and `sp` provide powerful functions. Spatial data in R can be stored as `Spatial*DataFrame` (where * can be Points, Lines or Polygons).

This example uses data which can be downloaded at [OpenGeocode](#).

At first, the working directory has to be set to the folder of the downloaded CSV data set. Furthermore, the package `rgdal` has to be loaded.

```
setwd("D:/GeocodeExample/")
library(rgdal)
```

Afterwards, the CSV file storing cities and their geographical coordinates is loaded into R as a `data.frame`

```
xy <- read.csv("worldcities.csv", stringsAsFactors = FALSE)
```

Often, it is useful to get a glimpse of the data and its structure (e.g. column names, data types etc.).

```
head(xy)
str(xy)
```

This shows that the latitude and longitude columns are interpreted as character values, since they hold entries like "-33.532". Yet, the later used function `SpatialPointsDataFrame()` which creates the spatial data set requires the coordinate values to be of the data type `numeric`. Thus the two columns have to be converted.

```
xy$latitude <- as.numeric(xy$latitude)
xy$longitude <- as.numeric(xy$longitude)
```

Few of the values cannot be converted into numeric data and thus, NA values are created. They have to be removed.

```
xy <- xy[ !is.na(xy$longitude), ]
```

Finally, the XY data set can be converted into a spatial data set. This requires the coordinates and the specification of the Coordinate Reference System (CRS) in which the coordinates are stored.

```
xySPoints <- SpatialPointsDataFrame(coords = c(xy[,c("longitude", "latitude")]),
proj4string = CRS("+proj=longlat +ellps=WGS84 +datum=WGS84 +no_defs"),
data = xy
)
```

The basic plot function can easily be used to sneak peak the produced spatial points.

```
plot(xySPoints, pch = ".")
```

Section 48.2: Importing a shape file (.shp)

rgdal

ESRI shape files can easily be imported into R by using the function `readOGR()` from the `rgdal` package.

```
library(rgdal)
shp <- readOGR(dsn = "/path/to/your/file", layer = "filename")
```

It is important to know, that the `dsn` must not end with / and the `layer` does not allow the file ending (e.g. .shp)

raster

Another possible way of importing shapefiles is via the `raster` library and the `shapefile` function:

```
library(raster)
shp <- shapefile("path/to/your/file.shp")
```

Note how the path definition is different from the `rgdal` import statement.

tmap

`tmap` package provides a nice wrapper for the `rgdal::readOGR` function.

```
library(tmap)
shp <- read_shape("path/to/your/file.shp")
```

Chapter 49: sqldf

Section 49.1: Basic Usage Examples

`sqldf()` from the package `sqldf` allows the use of SQLite queries to select and manipulate data in R. SQL queries are entered as character strings.

To select the first 10 rows of the "diamonds" dataset from the package `ggplot2`, for example:

```
data("diamonds")
head(diamonds)
```

```
# A tibble: 6 x 10
  carat      cut color clarity depth table price     x     y     z
  <dbl>     <ord> <ord>   <ord> <dbl> <dbl> <int> <dbl> <dbl> <dbl>
1 0.23     Ideal    E     SI2  61.5    55   326  3.95  3.98  2.43
2 0.21     Premium  E     SI1  59.8    61   326  3.89  3.84  2.31
3 0.23     Good    E     VS1  56.9    65   327  4.05  4.07  2.31
4 0.29     Premium I     VS2  62.4    58   334  4.20  4.23  2.63
5 0.31     Good    J     SI2  63.3    58   335  4.34  4.35  2.75
6 0.24 Very Good J     VVS2 62.8    57   336  3.94  3.96  2.48
```

```
require(sqldf)
sqldf("select * from diamonds limit 10")
```

```
carat      cut color clarity depth table price     x     y     z
1 0.23     Ideal    E     SI2  61.5    55   326  3.95  3.98  2.43
2 0.21     Premium  E     SI1  59.8    61   326  3.89  3.84  2.31
3 0.23     Good    E     VS1  56.9    65   327  4.05  4.07  2.31
4 0.29     Premium I     VS2  62.4    58   334  4.20  4.23  2.63
5 0.31     Good    J     SI2  63.3    58   335  4.34  4.35  2.75
6 0.24 Very Good J     VVS2 62.8    57   336  3.94  3.96  2.48
7 0.24 Very Good I     VVS1 62.3    57   336  3.95  3.98  2.47
8 0.26 Very Good H     SI1  61.9    55   337  4.07  4.11  2.53
9 0.22     Fair    E     VS2  65.1    61   337  3.87  3.78  2.49
10 0.23 Very Good H    VS1  59.4    61   338  4.00  4.05  2.39
```

To select the first 10 rows where for the color "E":

```
sqldf("select * from diamonds where color = 'E' limit 10")
```

```
carat      cut color clarity depth table price     x     y     z
1 0.23     Ideal    E     SI2  61.5    55   326  3.95  3.98  2.43
2 0.21     Premium  E     SI1  59.8    61   326  3.89  3.84  2.31
3 0.23     Good    E     VS1  56.9    65   327  4.05  4.07  2.31
4 0.22     Fair    E     VS2  65.1    61   337  3.87  3.78  2.49
5 0.20     Premium E     SI2  60.2    62   345  3.79  3.75  2.27
6 0.32     Premium E     I1   60.9    58   345  4.38  4.42  2.68
7 0.23 Very Good E     VS2  63.8    55   352  3.85  3.92  2.48
8 0.23 Very Good E     VS1  60.7    59   402  3.97  4.01  2.42
9 0.23 Very Good E     VS1  59.5    58   402  4.01  4.06  2.40
10 0.23    Good   E     VS1  64.1    59   402  3.83  3.85  2.46
```

Notice in the example above that quoted strings within the SQL query are quoted using " if the overall query is quoted with "" (this also works in reverse).

Suppose that we wish to add a new column to count the number of Premium cut diamonds over 1 carat:

```
sqlldf("select count(*) from diamonds where carat > 1 and color = 'E'")
```

```
count(*)
1     1892
```

Results of created values can also be returned as new columns:

```
sqlldf("select *, count(*) as cnt_big_E_colored_stones from diamonds where carat > 1 and color = 'E'
group by clarity")
```

| carat | cut | color | clarity | depth | table | price | x | y | z | cnt_big_E_colored_stones |
|--------|-----------|-------|---------|-------|-------|-------|------|------|------|--------------------------|
| 1 1.30 | Fair | E | I1 | 66.5 | 58 | 2571 | 6.79 | 6.75 | 4.50 | 65 |
| 2 1.28 | Ideal | E | IF | 60.7 | 57 | 18700 | 7.09 | 6.99 | 4.27 | 28 |
| 3 2.02 | Very Good | E | SI1 | 59.8 | 59 | 18731 | 8.11 | 8.20 | 4.88 | 499 |
| 4 2.03 | Premium | E | SI2 | 61.5 | 59 | 18477 | 8.24 | 8.16 | 5.04 | 666 |
| 5 1.51 | Ideal | E | VS1 | 61.5 | 57 | 18729 | 7.34 | 7.40 | 4.53 | 158 |
| 6 1.72 | Very Good | E | VS2 | 63.4 | 56 | 18557 | 7.65 | 7.55 | 4.82 | 318 |
| 7 1.20 | Ideal | E | VVS1 | 61.8 | 56 | 16256 | 6.78 | 6.87 | 4.22 | 52 |
| 8 1.55 | Ideal | E | VVS2 | 62.5 | 55 | 18188 | 7.38 | 7.40 | 4.62 | 106 |

If one would be interested what is the **max** price of the diamond **according** to the **cut**:

```
sqlldf("select cut, max(price) from diamonds group by cut")
```

| cut | max(price) |
|-----------|------------|
| Fair | 18574 |
| Good | 18788 |
| Ideal | 18806 |
| Premium | 18823 |
| Very Good | 18818 |

Chapter 50: Code profiling

Section 50.1: Benchmarking using microbenchmark

You can use the [microbenchmark](#) package to conduct "sub-millisecond accurate timing of expression evaluation".

In [this example](#) we are comparing the speeds of six equivalent data.table expressions for updating elements in a group, based on a certain condition.

More specifically:

A data.table with 3 columns: id, time and status. For each id, I want to find the record with the maximum time - then if for that record if the status is true, I want to set it to false if the time is > 7

```
library(microbenchmark)
library(data.table)

set.seed(20160723)
dt <- data.table(id = c(rep(seq(1:10000), each = 10)),
                  time = c(rep(seq(1:10000), 10)),
                  status = c(sample(c(TRUE, FALSE), 10000*10, replace = TRUE)))
setkey(dt, id, time) ## create copies of the data so the 'updates-by-reference' don't affect other
expressions
dt1 <- copy(dt)
dt2 <- copy(dt)
dt3 <- copy(dt)
dt4 <- copy(dt)
dt5 <- copy(dt)
dt6 <- copy(dt)

microbenchmark(

  expression_1 = {
    dt1[ dt1[order(time), .I[.N], by = id]$V1, status := status * time < 7 ]
  },

  expression_2 = {
    dt2[,status := c(.SD[~.N, status], .SD[.N, status * time > 7]), by = id]
  },

  expression_3 = {
    dt3[dt3[,.N, by = id][,cumsum(N)], status := status * time > 7]
  },

  expression_4 = {
    y <- dt4[,SD[.N],by=id]
    dt4[y, status := status & time > 7]
  },

  expression_5 = {
    y <- dt5[, SD[.N, .(time, status)], by = id][time > 7 & status]
    dt5[y, status := FALSE]
  },

  expression_6 = {
    dt6[ dt6[, .I == .I[which.max(time)]], by = id]$V1 & time > 7, status := FALSE]
  },
}
```

```

times = 10L ## specify the number of times each expression is evaluated
)

# Unit: milliseconds
#      expr      min       lq     mean    median      uq     max neval
# expression_1 11.646149 13.201670 16.808399 15.643384 18.78640 26.321346 10
# expression_2 8051.898126 8777.016935 9238.323459 8979.553856 9281.93377 12610.869058 10
# expression_3   3.208773   3.385841   4.207903   4.089515   4.70146   5.654702 10
# expression_4 15.758441 16.247833 20.677038 19.028982 21.04170 36.373153 10
# expression_5 7552.970295 8051.080753 8702.064620 8861.608629 9308.62842 9722.234921 10
# expression_6 18.403105 18.812785 22.427984 21.966764 24.66930 28.607064 10

```

The output shows that in this test expression_3 is the fastest.

References

[data.table - Adding and modifying columns](#)

[data.table - special grouping symbols in data.table](#)

Section 50.2: proc.time()

At its simplest, `proc.time()` gives the total elapsed CPU time in seconds for the current process. Executing it in the console gives the following type of output:

```

proc.time()

#      user      system    elapsed
# 284.507    120.397 515029.305

```

This is particularly useful for benchmarking specific lines of code. For example:

```

t1 <- proc.time()
fibb <- function (n) {
  if (n < 3) {
    return(c(0,1)[n])
  } else {
    return(fibb(n - 2) + fibb(n -1))
  }
}
print("Time one")
print(proc.time() - t1)

t2 <- proc.time()
fibb(30)

print("Time two")
print(proc.time() - t2)

```

This gives the following output:

```

source('~/active-rstudio-document')

# [1] "Time one"
#      user      system    elapsed
#          0          0          0

# [1] "Time two"

```

```
#    user  system elapsed
# 1.534   0.012   1.572
```

`system.time()` is a wrapper for `proc.time()` that returns the elapsed time for a particular command/expression.

```
print(t1 <- system.time(replicate(1000, 12^2)))
##  user  system elapsed
## 0.000   0.000   0.002
```

Note that the returned object, of class `proc.time`, is slightly more complicated than it appears on the surface:

```
str(t1)
## Class 'proc_time'  Named num [1:5] 0 0 0.002 0 0
##  ..- attr(*, "names")= chr [1:5] "user.self" "sys.self" "elapsed" "user.child" ...
```

Section 50.3: Microbenchmark

Microbenchmark is useful for estimating the time taking for otherwise fast procedures. For example, consider estimating the time taken to print hello world.

```
system.time(print("hello world"))

# [1] "hello world"
#    user  system elapsed
#      0      0      0
```

This is because `system.time` is essentially a wrapper function for `proc.time`, which measures in seconds. As printing "hello world" takes less than a second it appears that the time taken is less than a second, however this is not true. To see this we can use the package `microbenchmark`:

```
library(microbenchmark)
microbenchmark(print("hello world"))

# Unit: microseconds
#           expr     min      lq      mean    median      uq      max neval
# print("hello world") 26.336 29.984 44.11637 44.6835 45.415 158.824    100
```

Here we can see after running `print("hello world")` 100 times, the average time taken was in fact 44 microseconds. (Note that running this code will print "hello world" 100 times onto the console.)

We can compare this against an equivalent procedure, `cat("hello world\n")`, to see if it is faster than `print("hello world")`:

```
microbenchmark(cat("hello world\n"))

# Unit: microseconds
#           expr     min      lq      mean    median      uq      max neval
# cat("hello world\\n") 14.093 17.6975 23.73829 19.319 20.996 119.382    100
```

In this case `cat()` is almost twice as fast as `print()`.

Alternatively one can compare two procedures within the same `microbenchmark` call:

```
microbenchmark(print("hello world"), cat("hello world\n"))
# Unit: microseconds
# expr     min      lq      mean    median      uq      max neval
```

```
# print("hello world") 29.122 31.654 39.64255 34.5275 38.852 192.779 100
# cat("hello world\\n") 9.381 12.356 13.83820 12.9930 13.715 52.564 100
```

Section 50.4: System.time

System time gives you the CPU time required to execute a R expression, for example:

```
system.time(print("hello world"))

# [1] "hello world"
#   user  system elapsed
#       0      0      0
```

You can add larger pieces of code through use of braces:

```
system.time({
  library(numbers)
  Primes(1,10^5)
})
```

Or use it to test functions:

```
fibb <- function (n) {
  if (n < 3) {
    return(c(0,1)[n])
  } else {
    return(fibb(n - 2) + fibb(n -1))
  }
}

system.time(fibb(30))
```

Section 50.5: Line Profiling

One package for line profiling is [lineprof](#) which is written and maintained by Hadley Wickham. Here is a quick demonstration of how it works with auto.arima in the forecast package:

```
library(lineprof)
library(forecast)

l <- lineprof(auto.arima(AirPassengers))
shine(l)
```

This will provide you with a shiny app, which allows you to delve deeper into every function call. This enables you to see with ease what is causing your R code to slow down. There is a screenshot of the shiny app below:



Chapter 51: Control flow structures

Section 51.1: Optimal Construction of a For Loop

To illustrate the effect of good for loop construction, we will calculate the mean of each column in four different ways:

1. Using a poorly optimized for loop
2. Using a well optimized for loop
3. Using an `*apply` family of functions
4. Using the `colMeans` function

Each of these options will be shown in code; a comparison of the computational time to execute each option will be shown; and lastly a discussion of the differences will be given.

Poorly optimized for loop

```
column_mean_poor <- NULL
for (i in 1:length(mtcars)){
  column_mean_poor[i] <- mean(mtcars[[i]])
}
```

Well optimized for loop

```
column_mean_optimal <- vector("numeric", length(mtcars))
for (i in seq_along(mtcars)){
  column_mean_optimal <- mean(mtcars[[i]])
}
```

vapply Function

```
column_mean_vapply <- vapply(mtcars, mean, numeric(1))
```

colMeans Function

```
column_mean_colMeans <- colMeans(mtcars)
```

Efficiency comparison

The results of benchmarking these four approaches is shown below (code not displayed)

| Unit: microseconds | expr | min | lq | mean | median | uq | max | neval | cld |
|--------------------|----------|---------|----------|----------|----------|----------|---------|-------|-----|
| | poor | 240.986 | 262.0820 | 287.1125 | 275.8160 | 307.2485 | 442.609 | 100 | d |
| | optimal | 220.313 | 237.4455 | 258.8426 | 247.0735 | 280.9130 | 362.469 | 100 | c |
| | vapply | 107.042 | 109.7320 | 124.4715 | 113.4130 | 132.6695 | 202.473 | 100 | a |
| | colMeans | 155.183 | 161.6955 | 180.2067 | 175.0045 | 194.2605 | 259.958 | 100 | b |

Notice that the optimized `for` loop edged out the poorly constructed for loop. The poorly constructed for loop is constantly increasing the length of the output object, and at each change of the length, R is reevaluating the class of the object.

Some of this overhead burden is removed by the optimized for loop by declaring the type of output object and its length before starting the loop.

In this example, however, the use of an `vapply` function doubles the computational efficiency, largely because we

told R that the result had to be numeric (if any one result were not numeric, an error would be returned).

Use of the `colMeans` function is a touch slower than the `vapply` function. This difference is attributable to some error checks performed in `colMeans` and mainly to the `as.matrix` conversion (because `mtcars` is a `data.frame`) that weren't performed in the `vapply` function.

Section 51.2: Basic For Loop Construction

In this example we will calculate the squared deviance for each column in a data frame, in this case the `mtcars`.

Option A: integer index

```
squared_deviance <- vector("list", length(mtcars))
for (i in seq_along(mtcars)){
  squared_deviance[[i]] <- (mtcars[[i]] - mean(mtcars[[i]]))^2
}
```

`squared_deviance` is an 11 elements list, as expected.

```
class(squared_deviance)
length(squared_deviance)
```

Option B: character index

```
squared_deviance <- vector("list", length(mtcars))
Squared_deviance <- setNames(squared_deviance, names(mtcars))
for (k in names(mtcars)){
  squared_deviance[[k]] <- (mtcars[[k]] - mean(mtcars[[k]]))^2
}
```

What if we want a `data.frame` as a result? Well, there are many options for transforming a list into other objects. However, and maybe the simplest in this case, will be to store the `for` results in a `data.frame`.

```
squared_deviance <- mtcars #copy the original
squared_deviance[TRUE] <- NA #replace with NA or do squared_deviance[,] <- NA
for (i in seq_along(mtcars)){
  squared_deviance[[i]] <- (mtcars[[i]] - mean(mtcars[[i]]))^2
}
dim(squared_deviance)
[1] 32 11
```

The result will be the same event though we use the character option (B).

Section 51.3: The Other Looping Constructs: `while` and `repeat`

R provides two additional looping constructs, `while` and `repeat`, which are typically used in situations where the number of iterations required is indeterminate.

The `while` loop

The general form of a `while` loop is as follows,

```
while (condition) {
  ## do something
  ## in loop body
```

```
}
```

where condition is evaluated prior to entering the loop body. If condition evaluates to TRUE, the code inside of the loop body is executed, and this process repeats until condition evaluates to FALSE (or a **break** statement is reached; see below). Unlike the **for** loop, if a **while** loop uses a variable to perform incremental iterations, the variable must be declared and initialized ahead of time, and must be updated within the loop body. For example, the following loops accomplish the same task:

```
for (i in 0:4) {  
  cat(i, "\n")  
}  
# 0  
# 1  
# 2  
# 3  
# 4  
  
i <- 0  
while (i < 5) {  
  cat(i, "\n")  
  i <- i + 1  
}  
# 0  
# 1  
# 2  
# 3  
# 4
```

In the **while** loop above, the line `i <- i + 1` is necessary to prevent an infinite loop.

Additionally, it is possible to terminate a **while** loop with a call to **break** from inside the loop body:

```
iter <- 0  
while (TRUE) {  
  if (runif(1) < 0.25) {  
    break  
  } else {  
    iter <- iter + 1  
  }  
}  
iter  
#[1] 4
```

In this example, condition is always TRUE, so the only way to terminate the loop is with a call to **break** inside the body. Note that the final value of `iter` will depend on the state of your PRNG when this example is run, and should produce different results (essentially) each time the code is executed.

The **repeat** loop

The **repeat** construct is essentially the same as `while (TRUE) { ## something }`, and has the following form:

```
repeat ({  
  ## do something  
  ## in loop body  
)
```

The extra {} are not required, but the () are. Rewriting the previous example using **repeat**,

```

iter <- 0
repeat ({
  if (runif(1) < 0.25) {
    break
  } else {
    iter <- iter + 1
  }
})
iter
#[1] 2

```

More on `break`

It's important to note that `break` will *only terminate the immediately enclosing loop*. That is, the following is an infinite loop:

```

while (TRUE) {
  while (TRUE) {
    cat("inner loop\n")
    break
  }
  cat("outer loop\n")
}

```

With a little creativity, however, it is possible to break entirely from within a nested loop. As an example, consider the following expression, which, in its current state, will loop infinitely:

```

while (TRUE) {
  cat("outer loop body\n")
  while (TRUE) {
    cat("inner loop body\n")
    x <- runif(1)
    if (x < .3) {
      break
    } else {
      cat(sprintf("x is %.5f\n", x))
    }
  }
}

```

One possibility is to recognize that, unlike `break`, the `return` expression **does** have the ability to return control across multiple levels of enclosing loops. However, since `return` is only valid when used within a function, we cannot simply replace `break` with `return()` above, but also need to wrap the entire expression as an anonymous function:

```

(function() {
  while (TRUE) {
    cat("outer loop body\n")
    while (TRUE) {
      cat("inner loop body\n")
      x <- runif(1)
      if (x < .3) {
        return()
      } else {
        cat(sprintf("x is %.5f\n", x))
      }
    }
  }
})

```

```
)())
```

Alternatively, we can create a dummy variable (`exit`) prior to the expression, and activate it via `<<-` from the inner loop when we are ready to terminate:

```
exit <- FALSE
while (TRUE) {
  cat("outer loop body\n")
  while (TRUE) {
    cat("inner loop body\n")
    x <- runif(1)
    if (x < .3) {
      exit <<- TRUE
      break
    } else {
      cat(sprintf("x is %.5f\n", x))
    }
  }
  if (exit) break
}
```

Chapter 52: Column wise operation

Section 52.1: sum of each column

Suppose we need to do the `sum` of each column in a dataset

```
set.seed(20)
df1 <- data.frame(ID = rep(c("A", "B", "C"), each = 3), V1 = rnorm(9), V2 = rnorm(9))
m1 <- as.matrix(df1[-1])
```

There are many ways to do this. Using base R, the best option would be `colSums`

```
colSums(df1[-1], na.rm = TRUE)
```

Here, we removed the first column as it is non-numeric and did the `sum` of each column, specifying the `na.rm = TRUE` (in case there are any NAs in the dataset)

This also works with `matrix`

```
colSums(m1, na.rm = TRUE)
```

This can be done in a loop with `lapply/sapply/vapply`

```
lapply(df1[-1], sum, na.rm = TRUE)
```

It should be noted that the output is a `list`. If we need a `vector` output

```
sapply(df1[-1], sum, na.rm = TRUE)
```

Or

```
vapply(df1[-1], sum, na.rm = TRUE, numeric(1))
```

For matrices, if we want to loop through columns, then use `apply` with `MARGIN = 1`

```
apply(m1, 2, FUN = sum, na.rm = TRUE)
```

There are ways to do this with packages like `dplyr` or `data.table`

```
library(dplyr)
df1 %>%
  summarise_at(vars(matches("^V\\d+")), sum, na.rm = TRUE)
```

Here, we are passing a regular expression to match the column names that we need to get the `sum` in `summarise_at`. The regex will match all columns that start with V followed by one or more numbers (`\\d+`).

A `data.table` option is

```
library(data.table)
setDT(df1)[, lapply(.SD, sum, na.rm = TRUE), .SDcols = 2:ncol(df1)]
```

We convert the 'data.frame' to 'data.table' (`setDT(df1)`), specified the columns to be applied the function in `.SDcols` and loop through the Subset of Data.table (`.SD`) and get the `sum`.

If we need to use a group by operation, we can do this easily by specifying the group by column/columns

```
df1 %>%
  group_by(ID) %>%
  summarise_at(vars(matches("^V\\d+")), sum, na.rm = TRUE)
```

In cases where we need the **sum** of all the columns, **summarise_each** can be used instead of **summarise_at**

```
df1 %>%
  group_by(ID) %>%
  summarise_each(funs(sum(., na.rm = TRUE)))
```

The **data.table** option is

```
setDT(df1)[, lapply(.SD, sum, na.rm = TRUE), by = ID]
```

Chapter 53: JSON

Section 53.1: JSON to / from R objects

The [jsonlite package](#) is a fast JSON parser and generator optimized for statistical data and the web. The two main functions used to read and write JSON are `fromJSON()` and `toJSON()` respectively, and are designed to work with vectors, matrices and `data.frames`, and streams of JSON from the web.

Create a JSON array from a vector, and vice versa

```
library(jsonlite)

## vector to JSON
toJSON(c(1,2,3))
# [1] 1 2 3

fromJSON('[1,2,3]')
# [1] 1 2 3
```

Create a named JSON array from a list, and vice versa

```
toJSON(list(myVec = c(1,2,3)))
# {"myVec": [1,2,3]}

fromJSON('{"myVec": [1,2,3]}')
# $myVec
# [1] 1 2 3
```

More complex list structures

```
## list structures
lst <- list(a = c(1,2,3),
            b = list(letters[1:6]))

toJSON(lst)
# {"a": [1,2,3], "b": [ "a", "b", "c", "d", "e", "f"]}

fromJSON('{"a": [1,2,3], "b": [ "a", "b", "c", "d", "e", "f"]}' )
# $a
# [1] 1 2 3
#
# $b
# [,1] [,2] [,3] [,4] [,5] [,6]
# [1,] "a"   "b"   "c"   "d"   "e"   "f"
```

Create JSON from a `data.frame`, and vice versa

```
## converting a data.frame to JSON
df <- data.frame(id = seq_along(1:10),
                  val = letters[1:10])

toJSON(df)
#
[{"id":1,"val":"a"}, {"id":2,"val":"b"}, {"id":3,"val":"c"}, {"id":4,"val":"d"}, {"id":5,"val":"e"}, {"id":6,"val":"f"}, {"id":7,"val":"g"}, {"id":8,"val":"h"}, {"id":9,"val":"i"}, {"id":10,"val":"j"}]
```

```
## reading a JSON string
fromJSON('[{"id":1,"val":"a"}, {"id":2,"val":"b"}, {"id":3,"val":"c"}, {"id":4,"val":"d"}, {"id":5,"val":"e"}, {"id":6,"val":"f"}, {"id":7,"val":"g"}, {"id":8,"val":"h"}, {"id":9,"val":"i"}, {"id":10,"val":"j"}]')
#      id val
# 1    1   a
# 2    2   b
# 3    3   c
# 4    4   d
# 5    5   e
# 6    6   f
# 7    7   g
# 8    8   h
# 9    9   i
# 10  10   j
```

Read JSON direct from the internet

```
## Reading JSON from URL
googleway_issues <- fromJSON("https://api.github.com/repos/SymbolixAU/googleway/issues")

googleway_issues$url
# [1] "https://api.github.com/repos/SymbolixAU/googleway/issues/20"
"https://api.github.com/repos/SymbolixAU/googleway/issues/19"
# [3] "https://api.github.com/repos/SymbolixAU/googleway/issues/14"
"https://api.github.com/repos/SymbolixAU/googleway/issues/11"
# [5] "https://api.github.com/repos/SymbolixAU/googleway/issues/9"
"https://api.github.com/repos/SymbolixAU/googleway/issues/5"
# [7] "https://api.github.com/repos/SymbolixAU/googleway/issues/2"
```

Chapter 54: RODBC

Section 54.1: Connecting to Excel Files via RODBC

While RODBC is restricted to Windows computers with compatible architecture between R and any target RDMS, one of its key flexibilities is to work with Excel files as if they were SQL databases.

```
require(RODBC)
con = odbcConnectExcel("myfile.xlsx") # open a connection to the Excel file
sqlTables(con)$TABLE_NAME # show all sheets
df = sqlFetch(con, "Sheet1") # read a sheet
df = sqlQuery(con, "select * from [Sheet1 $]") # read a sheet (alternative SQL syntax)
close(con) # close the connection to the file
```

Section 54.2: SQL Server Management Database connection to get individual table

Another use of RODBC is in connecting with SQL Server Management Database. We need to specify the 'Driver' i.e. SQL Server here, the database name "Atilla" and then use the sqlQuery to extract either the full table or a fraction of it.

```
library(RODBC)
cn <- odbcDriverConnect(connection="Driver={SQL
Server};server=localhost;database=Atilla;trusted_connection=yes;")
tbl <- sqlQuery(cn, 'select top 10 * from table_1')
```

Section 54.3: Connecting to relational databases

```
library(RODBC)
con <- odbcDriverConnect("driver={Sql Server};server=servername;trusted connection=true")
dat <- sqlQuery(con, "select * from table");
close(con)
```

This will connect to a SQL Server instance. For more information on what your connection string should look like, visit connectionstrings.com

Also, since there's no database specified, you should make sure you fully qualify the object you're wanting to query like this databasename.schema.objectname

Chapter 55: lubridate

Section 55.1: Parsing dates and datetimes from strings with lubridate

The lubridate package provides convenient functions to format date and datetime objects from character strings. The functions are permutations of

| Letter | Element to parse | Base R equivalent |
|------------------|------------------|-------------------|
| y | year | %y, %Y |
| m (with y and d) | month | %m, %b, %h, %B |
| d | day | %d, %e |
| h | hour | %H, %I%p |
| m (with h and s) | minute | %M |
| s | seconds | %S |

e.g. `ymd()` for parsing a date with the year followed by the month followed by the day, e.g. "2016-07-22", or `ymd_hms()` for parsing a datetime in the order year, month, day, hours, minutes, seconds, e.g. "2016-07-22 13:04:47".

The functions are able to recognize most separators (such as /, -, and whitespace) without additional arguments. They also work with inconsistent separators.

Dates

The date functions return an object of class Date.

```
library(lubridate)

mdy(c('07/02/2016', '7 / 03 / 2016', '7 / 4 / 16'))
## [1] "2016-07-02" "2016-07-03" "2016-07-04"

ymd(c("20160724", "2016/07/23", "2016-07-25"))      # inconsistent separators
## [1] "2016-07-24" "2016-07-23" "2016-07-25"
```

Datetimes

Utility functions

Datetimes can be parsed using `ymd_hms` variants including `ymd_hm` and `ymd_h`. All datetime functions can accept a `tz` timezone argument akin to that of `as.POSIXct` or `strptime`, but which defaults to "UTC" instead of the local timezone.

The datetime functions return an object of class `POSIXct`.

```
x <- c("20160724 130102", "2016/07/23 14:02:01", "2016-07-25 15:03:00")
ymd_hms(x, tz="EST")
## [1] "2016-07-24 13:01:02 EST" "2016-07-23 14:02:01 EST"
## [3] "2016-07-25 15:03:00 EST"

ymd_hms(x)
## [1] "2016-07-24 13:01:02 UTC" "2016-07-23 14:02:01 UTC"
## [3] "2016-07-25 15:03:00 UTC"
```

Parser functions

`lubridate` also includes three functions for parsing datetimes with a formatting string like `as.POSIXct` or `strptime`:

| Function | Output Class | Formatting strings accepted |
|-------------------------------|--|--|
| <code>parse_date_time</code> | <code>POSIXct</code> | Flexible. Will accept <code>strptime</code> -style with % or <code>lubridate</code> datetime function name style, e.g <code>"ymd hms"</code> . Will accept a vector of orders for heterogeneous data and guess which is appropriate. |
| <code>parse_date_time2</code> | Default <code>POSIXct</code> ; if <code>lt = TRUE</code> , <code>POSIXlt</code> | Strict. Accepts only <code>strptime</code> tokens (with or without %) from a limited set. |
| <code>fast_strptime</code> | Default <code>POSIXlt</code> ; if <code>lt = FALSE</code> , <code>POSIXct</code> | Strict. Accepts only %-delimited <code>strptime</code> tokens with delimiters (-, /, :, etc.) from a limited set. |

```
x <- c('2016-07-22 13:04:47', '07/22/2016 1:04:47 pm')

parse_date_time(x, orders = c('mdy Imsp', 'ymd hms'))
## [1] "2016-07-22 13:04:47 UTC" "2016-07-22 13:04:47 UTC"

x <- c('2016-07-22 13:04:47', '2016-07-22 14:47:58')

parse_date_time2(x, orders = 'Ymd HMS')
## [1] "2016-07-22 13:04:47 UTC" "2016-07-22 14:47:58 UTC"

fast_strptime(x, format = '%Y-%m-%d %H:%M:%S')
## [1] "2016-07-22 13:04:47 UTC" "2016-07-22 14:47:58 UTC"
```

`parse_date_time2` and `fast_strptime` use a fast C parser for efficiency.

See `?parse_date_time` for formatting tokens.

Section 55.2: Difference between period and duration

Unlike durations, periods can be used to accurately model clock times without knowing when events such as leap seconds, leap days, and DST changes occur.

```
start_2012 <- ymd_hms("2012-01-01 12:00:00")
## [1] "2012-01-01 12:00:00 UTC"

# period() considers leap year calculations.
start_2012 + period(1, "years")
## [1] "2013-01-01 12:00:00 UTC"

# Here duration() doesn't consider leap year calculations.
start_2012 + duration(1)
## [1] "2012-12-31 12:00:00 UTC"
```

Section 55.3: Instants

An instant is a specific moment in time. Any date-time object that refers to a moment of time is recognized as an instant. To test if an object is an instant, use `is.instant`.

```
library(lubridate)

today_start <- dmy_hms("22.07.2016 12:00:00", tz = "IST") # default tz="UTC"
today_start
## [1] "2016-07-22 12:00:00 IST"
is.instant(today_start)
## [1] TRUE
```

```

now_dt <- ymd_hms(now(), tz="IST")
now_dt
## [1] "2016-07-22 13:53:09 IST"
is.instant(now_dt)
## [1] TRUE

is.instant("helloworld")
## [1] FALSE
is.instant(60)
## [1] FALSE

```

Section 55.4: Intervals, Durations and Periods

Intervals are simplest way of recording timespans in lubridate. An interval is a span of time that occurs between two specific **instants**.

```

# create interval by subtracting two instants
today_start <- ymd_hms("2016-07-22 12-00-00", tz="IST")
today_start
## [1] "2016-07-22 12:00:00 IST"
today_end <- ymd_hms("2016-07-22 23-59-59", tz="IST")
today_end
## [1] "2016-07-22 23:59:59 IST"
span <- today_end - today_start
span
## Time difference of 11.99972 hours
as.interval(span, today_start)
## [1] 2016-07-22 12:00:00 IST--2016-07-22 23:59:59 IST

# create interval using interval() function
span <- interval(today_start, today_end)
[1] 2016-07-22 12:00:00 IST--2016-07-22 23:59:59 IST

```

Durations measure the exact amount of time that occurs between two instants.

```

duration(60, "seconds")
## [1] "60s"

duration(2, "minutes")
## [1] "120s (~2 minutes)"

```

Note: Units larger than weeks are not used due to their variability.

Durations can be created using dseconds, dminutes and other duration helper functions.
Run ?quick_durations for complete list.

```

dseconds(60)
## [1] "60s"

dhours(2)
## [1] "7200s (~2 hours)"

dyears(1)
## [1] "31536000s (~365 days)"

```

Durations can be subtracted and added to instants to get new instants.

```
today_start + dhours(5)
```

```
## [1] "2016-07-22 17:00:00 IST"
today_start + dhours(5) + dminutes(30) + dseconds(15)
## [1] "2016-07-22 17:30:15 IST"
```

Durations can be created from intervals.

```
as.duration(span)
[1] "43199s (~12 hours)"
```

Periods measure the change in clock time that occurs between two instants.

Periods can be created using period function as well other helper functions like seconds, hours, etc. To get a complete list of period helper functions, Run ?quick_periods.

```
period(1, "hour")
## [1] "1H 0M 0S"

hours(1)
## [1] "1H 0M 0S"

period(6, "months")
## [1] "6m 0d 0H 0M 0S"

months(6)
## [1] "6m 0d 0H 0M 0S"

years(1)
## [1] "1y 0m 0d 0H 0M 0S"
```

is.period function can be used to check if an object is a period.

```
is.period(years(1))
## [1] TRUE

is.period(dydays(1))
## [1] FALSE
```

Section 55.5: Manipulating date and time in lubridate

```
date <- now()
date
## "2016-07-22 03:42:35 IST"

year(date)
## 2016

minute(date)
## 42

wday(date, label = T, abbr = T)
# [1] Fri
# Levels: Sun < Mon < Tues < Wed < Thurs < Fri < Sat

day(date) <- 31
## "2016-07-31 03:42:35 IST"

# If an element is set to a larger value than it supports, the difference
# will roll over into the next higher element
```

```
day(date) <- 32
## "2016-08-01 03:42:35 IST"
```

Section 55.6: Time Zones

`with_tz` returns a date-time as it would appear in a different time zone.

```
nyc_time <- now("America/New_York")
nyc_time
## [1] "2016-07-22 05:49:08 EDT"

# corresponding Europe/Moscow time
with_tz(nyc_time, tzzone = "Europe/Moscow")
## [1] "2016-07-22 12:49:08 MSK"
```

`force_tz` returns a the date-time that has the same clock time as x in the new time zone.

```
nyc_time <- now("America/New_York")
nyc_time
## [1] "2016-07-22 05:49:08 EDT"

force_tz(nyc_time, tzzone = "Europe/Moscow") # only timezone changes
## [1] "2016-07-22 05:49:08 MSK"
```

Section 55.7: Parsing date and time in lubridate

Lubridate provides `ymd()` series of functions for parsing character strings into dates. The letters y, m, and d correspond to the year, month, and day elements of a date-time.

```
mdy("07-21-2016")          # Returns Date
## [1] "2016-07-21"

mdy("07-21-2016", tz = "UTC")      # Returns a vector of class POSIXt
## "2016-07-21 UTC"

dmy("21-07-2016")          # Returns Date
## [1] "2016-07-21"

dmy(c("21.07.2016", "22.07.2016")) # Returns vector of class Date
## [1] "2016-07-21" "2016-07-22"
```

Section 55.8: Rounding dates

```
now_dt <- ymd_hms(now(), tz="IST")
now_dt
## [1] "2016-07-22 13:53:09 IST"
```

`round_date()` takes a date-time object and rounds it to the nearest integer value of the specified time unit.

```
round_date(now_dt, "minute")
## [1] "2016-07-22 13:53:00 IST"

round_date(now_dt, "hour")
```

```
## [1] "2016-07-22 14:00:00 IST"  
  
round_date(now_dt, "year")  
## [1] "2017-01-01 IST"
```

`floor_date()` takes a date-time object and rounds it down to the nearest integer value of the specified time unit.

```
floor_date(now_dt, "minute")  
## [1] "2016-07-22 13:53:00 IST"  
  
floor_date(now_dt, "hour")  
## [1] "2016-07-22 13:00:00 IST"  
  
floor_date(now_dt, "year")  
## [1] "2016-01-01 IST"
```

`ceiling_date()` takes a date-time object and rounds it up to the nearest integer value of the specified time unit.

```
ceiling_date(now_dt, "minute")  
## [1] "2016-07-22 13:54:00 IST"  
  
ceiling_date(now_dt, "hour")  
## [1] "2016-07-22 14:00:00 IST"  
  
ceiling_date(now_dt, "year")  
## [1] "2017-01-01 IST"
```

Chapter 56: Time Series and Forecasting

Section 56.1: Creating a ts object

Time series data can be stored as a `ts` object. `ts` objects contain information about seasonal frequency that is used by ARIMA functions. It also allows for calling of elements in the series by date using the `window` command.

```
#Create a dummy dataset of 100 observations
x <- rnorm(100)

#Convert this vector to a ts object with 100 annual observations
x <- ts(x, start = c(1900), freq = 1)

#Convert this vector to a ts object with 100 monthly observations starting in July
x <- ts(x, start = c(1900, 7), freq = 12)

#Alternatively, the starting observation can be a number:
x <- ts(x, start = 1900.5, freq = 12)

#Convert this vector to a ts object with 100 daily observations and weekly frequency starting in
#the first week of 1900
x <- ts(x, start = c(1900, 1), freq = 7)

#The default plot for a ts object is a line plot
plot(x)

#The window function can call elements or sets of elements by date

#Call the first 4 weeks of 1900
window(x, start = c(1900, 1), end = (1900, 4))

#Call only the 10th week in 1900
window(x, start = c(1900, 10), end = (1900, 10))

#Call all weeks including and after the 10th week of 1900
window(x, start = c(1900, 10))
```

It is possible to create `ts` objects with multiple series:

```
#Create a dummy matrix of 3 series with 100 observations each
x <- cbind(rnorm(100), rnorm(100), rnorm(100))

#Create a multi-series ts with annual observation starting in 1900
x <- ts(x, start = 1900, freq = 1)

#R will draw a plot for each series in the object
plot(x)
```

Section 56.2: Exploratory Data Analysis with time-series data

```
data(AirPassengers)
class(AirPassengers)
```

| 1 "ts"

In the spirit of Exploratory Data Analysis (EDA) a good first step is to look at a plot of your time-series data:

```
plot(AirPassengers) # plot the raw data  
abline(reg=lm(AirPassengers~time(AirPassengers))) # fit a trend line
```

For further EDA we examine cycles across years:

```
cycle(AirPassengers)
```

| | Jan | Feb | Mar | Apr | May | Jun | Jul | Aug | Sep | Oct | Nov | Dec |
|------|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|
| 1949 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 |
| 1950 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 |
| 1951 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 |
| 1952 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 |
| 1953 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 |
| 1954 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 |
| 1955 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 |
| 1956 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 |
| 1957 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 |
| 1958 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 |
| 1959 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 |
| 1960 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 |

```
boxplot(AirPassengers~cycle(AirPassengers)) #Box plot across months to explore seasonal effects
```

Chapter 57: strsplit function

Section 57.1: Introduction

strsplit is a useful function for breaking up a vector into a list on some character pattern. With typical R tools, the whole list can be reincorporated to a data.frame or part of the list might be used in a graphing exercise.

Here is a common usage of **strsplit**: break a character vector along a comma separator:

```
temp <- c("this,that,other", "hat,scarf,food", "woman,man,child")
# get a list split by commas
myList <- strsplit(temp, split=", ")
# print myList
myList
[[1]]
[1] "this"   "that"   "other"

[[2]]
[1] "hat"    "scarf"  "food"

[[3]]
[1] "woman"  "man"   "child"
```

As hinted above, the split argument is not limited to characters, but may follow a pattern dictated by a regular expression. For example, temp2 is identical to temp above except that the separators have been altered for each item. We can take advantage of the fact that the split argument accepts regular expressions to alleviate the irregularity in the vector.

```
temp2 <- c("this, that, other", "hat,scarf ,food", "woman; man ; child")
myList2 <- strsplit(temp2, split=" ?[,;] ?")
myList2
[[1]]
[1] "this"   "that"   "other"

[[2]]
[1] "hat"    "scarf"  "food"

[[3]]
[1] "woman"  "man"   "child"
```

Notes:

1. breaking down the regular expression syntax is out of scope for this example.
2. Sometimes matching regular expressions can slow down a process. As with many R functions that allow the use of regular expressions, the fixed argument is available to tell R to match on the split characters literally.

Chapter 58: Web scraping and parsing

Section 58.1: Basic scraping with rvest

[rvest](#) is a package for web scraping and parsing by Hadley Wickham inspired by Python's [Beautiful Soup](#). It leverages Hadley's [xml2](#) package's [libxml2](#) bindings for HTML parsing.

As part of the tidyverse, rvest is piped. It uses

- `xml2::read_html` to scrape the HTML of a webpage,
- which can then be subset with its `html_node` and `html_nodes` functions using CSS or XPath selectors, and
- parsed to R objects with functions like `html_text` and `html_table`.

To scrape the table of milestones from [the Wikipedia page on R](#), the code would look like

```
library(rvest)

url <- 'https://en.wikipedia.org/wiki/R_(programming_language)'

      # scrape HTML from website
url %>% read_html() %>%
  # select HTML tag with class="wikitable"
  html_node(css = '.wikitable') %>%
  # parse table into data.frame
  html_table() %>%
  # trim for printing
  dplyr::mutate(Description = substr(Description, 1, 70))

##   Release      Date          Description
## 1    0.16 1997-04-23 This is the last alpha version developed primarily by Ihaka
## 2    0.49 1997-04-23 This is the oldest source release which is currently available
## 3    0.60 1997-12-05 R becomes an official part of the GNU Project. The code is now
## 4   0.65.1 1999-10-07 First versions of update.packages and install.packages function
## 5    1.0 2000-02-29 Considered by its developers stable enough for production use
## 6    1.4 2001-12-19 S4 methods are introduced and the first version for Mac OS X
## 7    2.0 2004-10-04 Introduced lazy loading, which enables fast loading of data
## 8    2.1 2005-04-18 Support for UTF-8 encoding, and the beginnings of internationalization
## 9    2.11 2010-04-22 Support for Windows 64 bit systems.
## 10   2.13 2011-04-14 Adding a new compiler function that allows speeding up function
## 11   2.14 2011-10-31 Added mandatory namespaces for packages. Added a new parallel
## 12   2.15 2012-03-30 New load balancing functions. Improved serialization speed for
## 13   3.0 2013-04-03 Support for numeric index values 2^31 and larger on 64 bit systems.
```

While this returns a data.frame, note that as is typical for scraped data, there is still further data cleaning to be done: here, formatting dates, inserting NAs, and so on.

Note that data in a less consistently rectangular format may take looping or other further munging to successfully parse. If the website makes use of jQuery or other means to insert content, `read_html` may be insufficient to scrape, and a more robust scraper like RSelenium may be necessary.

Section 58.2: Using rvest when login is required

A common problem encountered when scrapping a web site is how to enter a userid and password to log into a web site.

In this example which I created to track my answers posted here to stack overflow. The overall flow is to login, go to a web page, collect information, add it to a data frame and then move to the next page.

```

library(rvest)

#Address of the login webpage
login<-
"https://stackoverflow.com/users/login?ssrc=head&returnurl=http%3a%2f%2fstackoverflow.com%2f"

#create a web session with the desired login address
pgsession<-html_session(login)
pgform<-html_form(pgsession)[[2]] #in this case the submit is the 2nd form
filled_form<-set_values(pgform, email="*****", password="*****")
submit_form(pgsession, filled_form)

#pre allocate the final results dataframe.
results<-data.frame()

#loop through all of the pages with the desired info
for (i in 1:5)
{
  #base address of the pages to extract information from
  url<-"http://stackoverflow.com/users/*****?tab=answers&sort=activity&page="
  url<-paste0(url, i)
  page<-jump_to(pgsession, url)

  #collect info on the question votes and question title
  summary<-html_nodes(page, "div .answer-summary")
  question<-matrix(html_text(html_nodes(summary, "div")), trim=TRUE), ncol=2, byrow = TRUE)

  #find date answered, hyperlink and whether it was accepted
  dateans<-html_node(summary, "span") %>% html_attr("title")
  hyperlink<-html_node(summary, "div a") %>% html_attr("href")
  accepted<-html_node(summary, "div") %>% html_attr("class")

  #create temp results then bind to final results
  rtemp<-cbind(question, dateans, accepted, hyperlink)
  results<-rbind(results, rtemp)
}

#Dataframe Clean-up
names(results)<-c("Votes", "Answer", "Date", "Accepted", "HyperLink")
results$Votes<-as.integer(as.character(results$Votes))
results$Accepted<-ifelse(results$Accepted=="answer-votes default", 0, 1)

```

The loop in this case is limited to only 5 pages, this needs to change to fit your application. I replaced the user specific values with ***** , hopefully this will provide some guidance for you problem.


```

sexmale    -2.463377  0.154587 -15.935 < 2e-16 ***
age       -0.042235  0.007415 -5.696 1.23e-08 ***
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' '

```

(Dispersion parameter for binomial family taken to be 1)

```

Null deviance: 1686.8 on 1312 degrees of freedom
Residual deviance: 1165.7 on 1308 degrees of freedom
AIC: 1175.7

```

Number of Fisher Scoring iterations: 5

- The first thing displayed is the call. It is a reminder of the model and the options specified.
- Next we see the deviance residuals, which are a measure of model fit. This part of output shows the distribution of the deviance residuals for individual cases used in the model.
- The next part of the output shows the coefficients, their standard errors, the z-statistic (sometimes called a Wald z-statistic), and the associated p-values.
 - The qualitative variables are "dummified". A modality is considered as the reference. The reference modality can be change with I in the formula.
 - All four predictors are statistically significant at a 0.1 % level.
 - The logistic regression coefficients give the change in the log odds of the outcome for a one unit increase in the predictor variable.
 - To see the *odds ratio* (multiplicative change in the odds of survival per unit increase in a predictor variable), exponentiate the parameter.
 - To see the confidence interval (CI) of the parameter, use `confint`.
- Below the table of coefficients are fit indices, including the null and deviance residuals and the Akaike Information Criterion (AIC), which can be used for comparing model performance.
 - When comparing models fitted by maximum likelihood to the same data, the smaller the AIC, the better the fit.
 - One measure of model fit is the significance of the overall model. This test asks whether the model with predictors fits significantly better than a model with just an intercept (i.e., a null model).

Example of odds ratios:

```

exp(coef(titanic.train)[3])

pclass3rd
0.08797765

```

With this model, compared to the first class, the 3rd class passengers have about a tenth of the odds of survival.

Example of confidence interval for the parameters:

```

confint(titanic.train)

Waiting for profiling to be done...
              2.5 %      97.5 %
(Intercept) 2.89486872 4.23734280
pclass2nd   -1.58986065 -0.75987230
pclass3rd   -2.81987935 -2.05419500

```

```
sexmale      -2.77180962 -2.16528316
age          -0.05695894 -0.02786211
```

Exemple of calculating the significance of the overall model:

The test statistic is distributed chi-squared with degrees of freedom equal to the differences in degrees of freedom between the current and the null model (i.e., the number of predictor variables in the model).

```
with(titanic.train, pchisq(null.deviance - deviance, df.null - df.residual
, lower.tail = FALSE))
[1] 1.892539e-111
```

The p-value is near 0, showing a strongly significant model.

Chapter 60: Reshaping data between long and wide forms

In R, tabular data is stored in data frames. This topic covers the various ways of transforming a single table.

Section 60.1: Reshaping data

Often data comes in tables. Generally one can divide this tabular data in wide and long formats. In a wide format, each variable has its own column.

Person Height [cm] Age [yr]

| | | |
|--------|-----|----|
| Alison | 178 | 20 |
| Bob | 174 | 45 |
| Carl | 182 | 31 |

However, sometimes it is more convenient to have a long format, in which all variables are in one column and the values are in a second column.

Person Variable Value

| | | |
|--------|-------------|-----|
| Alison | Height [cm] | 178 |
| Bob | Height [cm] | 174 |
| Carl | Height [cm] | 182 |
| Alison | Age [yr] | 20 |
| Bob | Age [yr] | 45 |
| Carl | Age [yr] | 31 |

Base R, as well as third party packages can be used to simplify this process. For each of the options, the `mtcars` dataset will be used. By default, this dataset is in a long format. In order for the packages to work, we will insert the row names as the first column.

```
mtcars # shows the dataset  
data <- data.frame(observation=row.names(mtcars), mtcars)
```

Base R

There are two functions in base R that can be used to convert between wide and long format: `stack()` and `unstack()`.

```
long <- stack(data)  
long # this shows the long format  
wide <- unstack(long)  
wide # this shows the wide format
```

However, these functions can become very complex for more advanced use cases. Luckily, there are other options using third party packages.

The tidyr package

This package uses `gather()` to convert from wide to long and `spread()` to convert from long to wide.

```
library(tidyr)  
long <- gather(data, variable, value, 2:12) # where variable is the name of the  
# variable column, value indicates the name of the value column and 2:12 refers to
```

```
# the columns to be converted.
long # shows the long result
wide <- spread(long, variable, value)
wide # shows the wide result (~data)
```

The data.table package

The data.table package extends the reshape2 functions and uses the function `melt()` to go from wide to long and `dcast()` to go from long to wide.

```
library(data.table)
long <- melt(data, 'observation', 2:12, 'variable', 'value')
long # shows the long result
wide <- dcast(long, observation ~ variable)
wide # shows the wide result (~data)
```

Section 60.2: The reshape function

The most flexible base R function for reshaping data is `reshape`. See `?reshape` for its syntax.

```
# create unbalanced longitudinal (panel) data set
set.seed(1234)
df <- data.frame(identifier=rep(1:5, each=3),
                  location=rep(c("up", "down", "left", "up", "center"), each=3),
                  period=rep(1:3, 5), counts=sample(35, 15, replace=TRUE),
                  values=rnorm(15, 5, 10))[-c(4,8,11),]
df
```

| | identifier | location | period | counts | values |
|----|------------|----------|--------|--------|----------|
| 1 | 1 | up | 1 | 4 | 9.186478 |
| 2 | 1 | up | 2 | 22 | 6.431116 |
| 3 | 1 | up | 3 | 22 | 6.334104 |
| 5 | 2 | down | 2 | 31 | 6.161130 |
| 6 | 2 | down | 3 | 23 | 6.583062 |
| 7 | 3 | left | 1 | 1 | 6.513467 |
| 9 | 3 | left | 3 | 24 | 5.199980 |
| 10 | 4 | up | 1 | 18 | 6.093998 |
| 12 | 4 | up | 3 | 20 | 7.628488 |
| 13 | 5 | center | 1 | 10 | 9.573291 |
| 14 | 5 | center | 2 | 33 | 9.156725 |
| 15 | 5 | center | 3 | 11 | 5.228851 |

Note that the data.frame is unbalanced, that is, unit 2 is missing an observation in the first period, while units 3 and 4 are missing observations in the second period. Also, note that there are two variables that vary over the periods: counts and values, and two that do not vary: identifier and location.

Long to Wide

To reshape the data.frame to wide format,

```
# reshape wide on time variable
df.wide <- reshape(df, idvar="identifier", timevar="period",
                    v.names=c("values", "counts"), direction="wide")
df.wide
```

| | identifier | location | values.1 | counts.1 | values.2 | counts.2 | values.3 | counts.3 |
|----|------------|----------|----------|----------|----------|----------|----------|----------|
| 1 | 1 | up | 9.186478 | | 4 | 6.431116 | 22 | 6.334104 |
| 5 | 2 | down | | NA | NA | 6.161130 | 31 | 6.583062 |
| 7 | 3 | left | 6.513467 | | 1 | NA | NA | 5.199980 |
| 10 | 4 | up | 6.093998 | | 18 | NA | NA | 7.628488 |

13

5 center 9.573291

10 9.156725

33 5.228851

11

Notice that the missing time periods are filled in with NAs.

In reshaping wide, the "v.names" argument specifies the columns that vary over time. If the location variable is not necessary, it can be dropped prior to reshaping with the "drop" argument. In dropping the only non-varying / non-id column from the data.frame, the v.names argument becomes unnecessary.

```
reshape(df, idvar="identifier", timevar="period", direction="wide",
       drop="location")
```

Wide to Long

To reshape long with the current df.wide, a minimal syntax is

```
reshape(df.wide, direction="long")
```

However, this is typically trickier:

```
# remove "." separator in df.wide names for counts and values
names(df.wide)[grep("\\\\.", names(df.wide))] <-
  gsub("\\\\.", "", names(df.wide)[grep("\\\\.", names(df.wide))])
```

Now the simple syntax will produce an error about undefined columns.

With column names that are more difficult for the `reshape` function to automatically parse, it is sometimes necessary to add the "varying" argument which tells `reshape` to group particular variables in wide format for the transformation into long format. This argument takes a list of vectors of variable names or indices.

```
reshape(df.wide, idvar="identifier",
       varying=list(c(3,5,7), c(4,6,8)), direction="long")
```

In reshaping long, the "v.names" argument can be provided to rename the resulting varying variables.

Sometimes the specification of "varying" can be avoided by use of the "sep" argument which tells `reshape` what part of the variable name specifies the value argument and which specifies the time argument.

Chapter 61: RMarkdown and knitr presentation

| Parameter | definition |
|-----------|---|
| title | the title of the document |
| author | The author of the document |
| date | The date of the document: Can be "r <code>format(Sys.time(), '%d %B, %Y')</code> " |
| author | The author of the document |
| output | The output format of the document: at least 10 format available. For html document, <code>html_output</code> . For PDF document, <code>pdf_document</code> , .. |

Section 61.1: Adding a footer to an ioslides presentation

Adding a footer is not natively possible. Luckily, we can make use of jQuery and CSS to add a footer to the slides of an ioslides presentation rendered with knitr. First of all we have to include the jQuery plugin. This is done by the line

```
<script src="https://ajax.googleapis.com/ajax/libs/jquery/1.12.2/jquery.min.js"></script>
```

Now we can use jQuery to alter the DOM (*document object model*) of our presentation. In other words: we alter the HTML structure of the document. As soon as the presentation is loaded (`$(document).ready(function() { ... })`), we select all slides, that do not have the class attributes `.title-slide`, `.backdrop`, or `.segue` and add the tag `<footer></footer>` right before each slide is 'closed' (so before `</slide>`). The attribute `label` carries the content that will be displayed later on.

All we have to do now is to layout our footer with CSS:

After each `<footer>` (`footer ::after`):

- display the content of the attribute `label`
- use font size 12
- position the footer (20 pixels from the bottom of the slide and 60 pxs from the left)

(the other properties can be ignored but might have to be modified if the presentation uses a different style template).

```
---
title: "Adding a footer to presentaiton slides"
author: "Martin Schmelzer"
date: "26 Juli 2016"
output: ioslides_presentation
---

```{r setup, include=FALSE}
knitr::opts_chunk$set(echo = FALSE)
```
```

```
## Slide 1  
This is slide 1.  
## Slide 2  
This is slide 2  
# Test  
## Slide 3  
And slide 3.
```

The result will look like this:



Section 61.2: Rstudio example

This is a script saved as .Rmd, on the contrary of r scripts saved as .R.

To knit the script, either use the render function or use the shortcut button in Rstudio.

```
---  
title: "Rstudio exemple of a rmd file"  
author: 'stack user'  
date: "22 July 2016"
```

```
output: html_document
```

```
---
```

The header is used to define the general parameters and the metadata.

```
## R Markdown
```

This is an R Markdown document.

It is a script written in markdown with the possibility to insert chunk of R code in it.
To insert R code, it needs to be encapsulated into inverted quote.

Like that for a long piece of code:

```
```{r cars}
summary(cars)
```
```

And like ``r cat("that")`` for small piece of code.

```
## Including Plots
```

You can also embed plots, for example:

```
```{r echo=FALSE}
plot(pressure)
```
```

Chapter 62: Scope of variables

Section 62.1: Environments and Functions

Variables declared inside a function only exist (unless passed) inside that function.

```
x <- 1

foo <- function(x) {
  y <- 3
  z <- x + y
  return(z)
}

y
```

Error: object 'y' not found

Variables passed into a function and then reassigned are overwritten, *but only inside the function*.

```
foo <- function(x) {
  x <- 2
  y <- 3
  z <- x + y
  return(z)
}

foo(1)
x
```

5
1

Variables assigned in a higher environment than a function exist within that function, without being passed.

```
foo <- function() {
  y <- 3
  z <- x + y
  return(z)
}

foo()
```

4

Section 62.2: Function Exit

The `on.exit()` function is handy for variable clean up if global variables must be assigned.

Some parameters, especially those for graphics, can only be set globally. This small function is common when

creating more specialized plots.

```
new_plot <- function(...) {  
  old_pars <- par(mar = c(5,4,4,2) + .1, mfrow = c(1,1))  
  on.exit(par(old_pars))  
  plot(...)  
}
```

Section 62.3: Sub functions

Functions called within a function (ie subfunctions) must be defined within that function to access any variables defined in the local environment without being passed.

This fails:

```
bar <- function() {  
  z <- x + y  
  return(z)  
}  
  
foo <- function() {  
  y <- 3  
  z <- bar()  
  return(z)  
}  
  
foo()
```

Error in bar() : object 'y' not found

This works:

```
foo <- function() {  
  bar <- function() {  
    z <- x + y  
    return(z)  
  }  
  
  y <- 3  
  z <- bar()  
  return(z)  
}  
  
foo()
```

4

Section 62.4: Global Assignment

Variables can be assigned globally from any environment using `<-.` `bar()` can now access `y`.

```
bar <- function() {
```

```
z <- x + y
return(z)
}

foo <- function() {
  y <<- 3
  z <- bar()
  return(z)
}

foo()
```

4

Global assignment is highly discouraged. Use of a wrapper function or explicitly calling variables from another local environment is greatly preferred.

Section 62.5: Explicit Assignment of Environments and Variables

Environments in R can be explicitly call and named. Variables can be explicitly assigned and call to or from those environments.

A commonly created environment is one which encloses package :base or a subenvironment within package :base.

```
e1 <- new.env(parent = baseenv())
e2 <- new.env(parent = e1)
```

Variables can be explicitly assigned and call to or from those environments.

```
assign("a", 3, envir = e1)
get("a", envir = e1)
get("a", envir = e2)
```

3

3

Since e2 inherits from e1, a is 3 in both e1 and e2. However, assigning a within e2 does not change the value of a in e1.

```
assign("a", 2, envir = e2)
get("a", envir = e2)
get("a", envir = e1)
```

3

2

Chapter 63: Performing a Permutation Test

Section 63.1: A fairly general function

We will use the built in [tooth growth dataset](#). We are interested in whether there is a statistically significant difference in tooth growth when the guinea pigs are given vitamin C vs orange juice.

Here's the full example:

```
teethVC = ToothGrowth[ToothGrowth$supp == 'VC', ]
teethOJ = ToothGrowth[ToothGrowth$supp == 'OJ', ]

permutationTest = function(vectorA, vectorB, testStat){
  N = 10^5
  fullSet = c(vectorA, vectorB)
  lengthA = length(vectorA)
  lengthB = length(vectorB)
  trials <- replicate(N,
    {index <- sample(lengthB + lengthA, size = lengthA, replace = FALSE)
     testStat((fullSet[index]), fullSet[-index]) })
  trials
}
vec1 = teethVC$len;
vec2 = teethOJ$len;
subtractMeans = function(a, b){ return (mean(a) - mean(b)) }
result = permutationTest(vec1, vec2, subtractMeans)
observedMeanDifference = subtractMeans(vec1, vec2)
result = c(result, observedMeanDifference)
hist(result)
abline(v=observedMeanDifference, col = "blue")
pValue = 2*mean(result <= (observedMeanDifference))
pValue
```

After we read in the CSV, we define the function

```
permutationTest = function(vectorA, vectorB, testStat){
  N = 10^5
  fullSet = c(vectorA, vectorB)
  lengthA = length(vectorA)
  lengthB = length(vectorB)
  trials <- replicate(N,
    {index <- sample(lengthB + lengthA, size = lengthA, replace = FALSE)
     testStat((fullSet[index]), fullSet[-index]) })
  trials
}
```

This function takes two vectors, and shuffles their contents together, then performs the function `testStat` on the shuffled vectors. The result of `teststat` is added to `trials`, which is the return value.

It does this `N = 10^5` times. Note that the value `N` could very well have been a parameter to the function.

This leaves us with a new set of data, `trials`, the set of means that might result if there truly is no relationship between the two variables.

Now to define our test statistic:

```
subtractMeans = function(a, b){ return (mean(a) - mean(b))}
```

Perform the test:

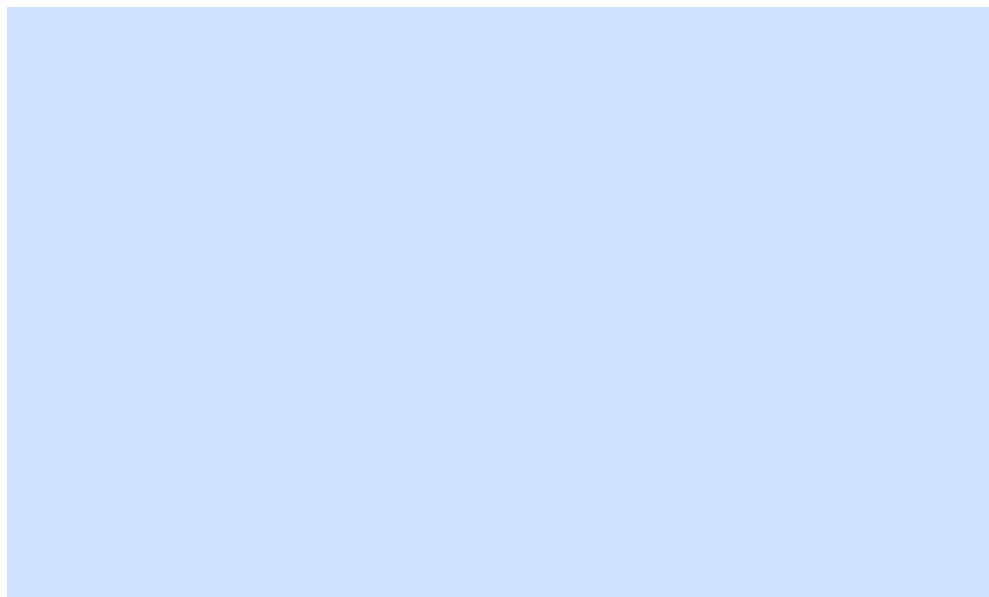
```
result = permutationTest(vec1, vec2, subtractMeans)
```

Calculate our actual observed mean difference:

```
observedMeanDifference = subtractMeans(vec1, vec2)
```

Let's see what our observation looks like on a histogram of our test statistic.

```
hist(result)
abline(v=observedMeanDifference, col = "blue")
```



It doesn't *look* like our observed result is very likely to occur by random chance...

We want to calculate the p-value, the likelihood of the original observed result if there is no relationship between the two variables.

```
pValue = 2*mean(result >= (observedMeanDifference))
```

Let's break that down a bit:

```
result >= (observedMeanDifference)
```

Will create a boolean vector, like:

```
FALSE TRUE FALSE FALSE TRUE FALSE ...
```

With TRUE every time the value of result is greater than or equal to the observedMean.

The function `mean` will interpret this vector as 1 for TRUE and 0 for FALSE, and give us the percentage of 1's in the mix, ie the number of times our shuffled vector mean difference surpassed or equalled what we observed.

Finally, we multiply by 2 because the distribution of our test statistic is highly symmetric, and we really want to know which results are "more extreme" than our observed result.

All that's left is to output the p-value, which turns out to be **0.06093939**. Interpretation of this value is subjective, but I would say that it looks like Vitamin C promotes tooth growth quite a lot more than Orange Juice does.


```

        if (err < smallestError) {
            smallestError = err
            print(paste(depth, rounds, err))
        }
    }

cv <- 30
trainSet <- adultsTrsf[1:trainPortion,]
cvDivider <- floor(nrow(trainSet) / (cv+1))

smallestError <- 100
for (depth in seq(1,10,1)) {
    for (rounds in seq(1,20,1)) {
        totalError <- c()
        indexCount <- 1
        for (cv in seq(1:cv)) {
            # assign chunk to data test
            dataTestIndex <- c((cv * cvDivider):(cv * cvDivider + cvDivider))
            dataTest <- trainSet[dataTestIndex,]
            # everything else to train
            dataTrain <- trainSet[-dataTestIndex,]

            bst <- xgboost(data = as.matrix(dataTrain[,predictors]),
                            label = dataTrain[,outcomeName],
                            max.depth=depth, nround=rounds,
                            objective = "reg:linear", verbose=0)
            gc()
            predictions <- predict(bst, as.matrix(dataTest[,predictors]),

outputmargin=TRUE)

            err <- rmse(as.numeric(dataTest[,outcomeName]), as.numeric(predictions))
            totalError <- c(totalError, err)
        }
        if (mean(totalError) < smallestError) {
            smallestError = mean(totalError)
            print(paste(depth, rounds, smallestError))
        }
    }
}

#####
# Test both models out on full data set

trainSet <- adultsTrsf[ 1:trainPortion,]

# assign everything else to test
testSet <- adultsTrsf[(trainPortion+1):nrow(adultsTrsf),]

bst <- xgboost(data = as.matrix(trainSet[,predictors]),
                label = trainSet[,outcomeName],
                max.depth=4, nround=19, objective = "reg:linear", verbose=0)
pred <- predict(bst, as.matrix(testSet[,predictors])), outputmargin=TRUE)
rmse(as.numeric(testSet[,outcomeName]), as.numeric(pred))

bst <- xgboost(data = as.matrix(trainSet[,predictors]),
                label = trainSet[,outcomeName],
                max.depth=3, nround=20, objective = "reg:linear", verbose=0)
pred <- predict(bst, as.matrix(testSet[,predictors])), outputmargin=TRUE)
rmse(as.numeric(testSet[,outcomeName]), as.numeric(pred))

```

Chapter 65: R code vectorization best practices

Section 65.1: By row operations

The key in vectorizing R code, is to reduce or eliminate "by row operations" or method dispatching of R functions.

That means that when approaching a problem that at first glance requires "by row operations", such as calculating the means of each row, one needs to ask themselves:

- What are the classes of the data sets I'm dealing with?
- Is there an existing compiled code that can achieve this without the need of repetitive evaluation of R functions?
- If not, can I do these operation by columns instead by row?
- Finally, is it worth spending a lot of time on developing complicated vectorized code instead of just running a simple `apply` loop? In other words, is the data big/sophisticated enough that R can't handle it efficiently using a simple loop?

Putting aside the memory pre-allocation issue and growing object in loops, we will focus in this example on how to possibly avoid `apply` loops, method dispatching or re-evaluating R functions within loops.

A standard/easy way of calculating mean by row would be:

| | <code>apply(mtcars, 1, mean)</code> | | | | |
|--------------------|-------------------------------------|-------------------|------------------|-------------------|--|
| Mazda RX4 | Mazda RX4 Wag | Datsun 710 | Hornet 4 Drive | Hornet Sportabout | |
| Valiant | Duster 360 | 23.59818 | 38.73955 | 53.66455 | |
| 29.90727 | 29.98136 | | | | |
| 35.04909 | 59.72000 | | | | |
| Merc 240D | Merc 230 | Merc 280 | Merc 280C | Merc 450SE | |
| Merc 450SL | Merc 450SLC | 31.86000 | 31.78727 | 46.43091 | |
| 24.63455 | 27.23364 | | | | |
| 46.50000 | 46.35000 | | | | |
| Cadillac Fleetwood | Lincoln Continental | Chrysler Imperial | Fiat 128 | Honda Civic | |
| Toyota Corolla | Toyota Corona | 65.97227 | 19.44091 | 17.74227 | |
| 66.23273 | 66.05855 | | | | |
| 18.81409 | 24.88864 | | | | |
| Dodge Challenger | AMC Javelin | Camaro Z28 | Pontiac Firebird | Fiat X1-9 | |
| Porsche 914-2 | Lotus Europa | 58.75273 | 57.37955 | 18.92864 | |
| 47.24091 | 46.00773 | | | | |
| 24.77909 | 24.88027 | | | | |
| Ford Pantera L | Ferrari Dino | Maserati Bora | Volvo 142E | | |
| 60.97182 | 34.50818 | 63.15545 | 26.26273 | | |

But can we do better? Let's see what happened here:

1. First, we converted a `data.frame` to a `matrix`. (Note that this happens within the `apply` function.) This is both inefficient and dangerous. a `matrix` can't hold several column types at a time. Hence, such conversion will probably lead to loss of information and some times to misleading results (compare `apply(iris, 2, class)` with `str(iris)` or with `sapply(iris, class)`).
2. Second of all, we performed an operation repetitively, one time for each row. Meaning, we had to evaluate some R function `nrow(mtcars)` times. In this specific case, `mean` is not a computationally expensive function, hence R could likely easily handle it even for a big data set, but what would happen if we need to calculate the standard deviation by row (which involves an expensive square root operation)? Which brings us to the next point:

3. We evaluated the R function many times, but maybe there already is a compiled version of this operation?

Indeed we could simply do:

| rowMeans(mtcars) | Mazda RX4 | Mazda RX4 Wag | Datsun 710 | Hornet 4 Drive | Hornet Sportabout |
|--------------------|---------------------|-------------------|------------------|----------------|-------------------|
| | Valiant | Duster 360 | | | |
| 29.90727 | 29.98136 | 23.59818 | 38.73955 | 53.66455 | |
| 35.04909 | 59.72000 | | | | |
| Merc 240D | Merc 230 | Merc 280 | Merc 280C | Merc 450SE | |
| Merc 450SL | Merc 450SLC | 31.86000 | 31.78727 | 46.43091 | |
| 24.63455 | 27.23364 | | | | |
| 46.50000 | 46.35000 | | | | |
| Cadillac Fleetwood | Lincoln Continental | Chrysler Imperial | Fiat 128 | Honda Civic | |
| Toyota Corolla | Toyota Corona | 65.97227 | 19.44091 | 17.74227 | |
| 66.23273 | 66.05855 | | | | |
| 18.81409 | 24.88864 | | | | |
| Dodge Challenger | AMC Javelin | Camaro Z28 | Pontiac Firebird | Fiat X1-9 | |
| Porsche 914-2 | Lotus Europa | 58.75273 | 57.37955 | 18.92864 | |
| 47.24091 | 46.00773 | | | | |
| 24.77909 | 24.88027 | | | | |
| Ford Pantera L | Ferrari Dino | Maserati Bora | Volvo 142E | | |
| 60.97182 | 34.50818 | 63.15545 | 26.26273 | | |

This involves no by row operations and therefore no repetitive evaluation of R functions. **However**, we still converted a `data.frame` to a `matrix`. Though `rowMeans` has an error handling mechanism and it won't run on a data set that it can't handle, it's still has an efficiency cost.

```
rowMeans(iris)
Error in rowMeans(iris) : 'x' must be numeric
```

But still, can we do better? We could try instead of a matrix conversion with error handling, a different method that will allow us to use `mtcars` as a vector (because a `data.frame` is essentially a `list` and a `list` is a `vector`).

```
Reduce(`+`, mtcars)/ncol(mtcars)
[1] 29.90727 29.98136 23.59818 38.73955 53.66455 35.04909 59.72000 24.63455 27.23364 31.86000
31.78727 46.43091 46.50000 46.35000 66.23273 66.05855
[17] 65.97227 19.44091 17.74227 18.81409 24.88864 47.24091 46.00773 58.75273 57.37955 18.92864
24.77909 24.88027 60.97182 34.50818 63.15545 26.26273
```

Now for possible speed gain, we lost column names and error handling (including NA handling).

Another example would be calculating mean by group, using base R we could try

```
aggregate(. ~ cyl, mtcars, mean)
cyl      mpg     disp      hp      drat      wt      qsec      vs      am      gear      carb
1   4 26.66364 105.1364 82.63636 4.070909 2.285727 19.13727 0.9090909 0.7272727 4.090909 1.545455
2   6 19.74286 183.3143 122.28571 3.585714 3.117143 17.97714 0.5714286 0.4285714 3.857143 3.428571
3   8 15.10000 353.1000 209.21429 3.229286 3.999214 16.77214 0.0000000 0.1428571 3.285714 3.500000
```

Still, we are basically evaluating an R function in a loop, but the loop is now hidden in an internal C function (it matters little whether it is a C or an R loop).

Could we avoid it? Well there is a compiled function in R called `rowsum`, hence we could do:

```
rowsum(mtcars[-2], mtcars$cyl)/table(mtcars$cyl)
mpg     disp      hp      drat      wt      qsec      vs      am      gear      carb
4 26.66364 105.1364 82.63636 4.070909 2.285727 19.13727 0.9090909 0.7272727 4.090909 1.545455
```

```
6 19.74286 183.3143 122.28571 3.585714 3.117143 17.97714 0.5714286 0.4285714 3.857143 3.428571  
8 15.10000 353.1000 209.21429 3.229286 3.999214 16.77214 0.0000000 0.1428571 3.285714 3.500000
```

Though we had to convert to a matrix first too.

At this point we may question whether our current data structure is the most appropriate one. Is a **data.frame** is the best practice? Or should one just switch to a **matrix** data structure in order to gain efficiency?

By row operations will get more and more expensive (even in matrices) as we start to evaluate expensive functions each time. Lets us consider a variance calculation by row example.

Lets say we have a matrix **m**:

```
set.seed(100)  
m <- matrix(sample(1e2), 10)  
  
m  
[,1] [,2] [,3] [,4] [,5] [,6] [,7] [,8] [,9] [,10]  
[1,]    8   33   39   86   71  100   81   68   89   84  
[2,]   12   16   57   80   32   82   69   11   41   92  
[3,]   62   91   53   13   42   31   60   70   98   79  
[4,]   66   94   29   67   45   59   20   96   64    1  
[5,]   36   63   76    6   10   48   85   75   99    2  
[6,]   18     4   27   19   44   56   37   95   26   40  
[7,]    3   24   21   25   52   51   83   28   49   17  
[8,]   46     5   22   43   47   74   35   97   77   65  
[9,]   55   54   78   34   50   90   30   61   14   58  
[10,]  88   73   38   15    9   72    7   93   23   87
```

One could simply do:

```
apply(m, 1, var)  
[1] 871.6556 957.5111 699.2111 941.4333 1237.3333 641.8222 539.7889 759.4333 500.4889  
1255.6111
```

On the other hand, one could also completely vectorize this operation by following the formula of variance

```
RowVar <- function(x) {  
  rowSums((x - rowMeans(x))^2)/(dim(x)[2] - 1)  
}  
RowVar(m)  
[1] 871.6556 957.5111 699.2111 941.4333 1237.3333 641.8222 539.7889 759.4333 500.4889  
1255.6111
```

Chapter 66: Missing values

When we don't know the value a variable takes, we say its value is missing, indicated by NA.

Section 66.1: Examining missing data

anyNA reports whether any missing values are present; while `is.na` reports missing values elementwise:

```
vec <- c(1, 2, 3, NA, 5)

anyNA(vec)
# [1] TRUE
is.na(vec)
# [1] FALSE FALSE FALSE  TRUE FALSE
```

`is.na` returns a logical vector that is coerced to integer values under arithmetic operations (with FALSE=0, TRUE=1). We can use this to find out how many missing values there are:

```
sum(is.na(vec))
# [1] 1
```

Extending this approach, we can use `colSums` and `is.na` on a data frame to count NAs per column:

```
colSums(is.na(airquality))
#   Ozone Solar.R    Wind     Temp    Month     Day
#     37       7       0       0       0       0
```

The [naniar package](#) (currently on github but not CRAN) offers further tools for exploring missing values.

Section 66.2: Reading and writing data with NA values

When reading tabular datasets with the `read.*` functions, R automatically looks for missing values that look like "NA". However, missing values are not always represented by NA. Sometimes a dot (.), a hyphen(-) or a character-value (e.g.: empty) indicates that a value is NA. The `na.strings` parameter of the `read.*` function can be used to tell R which symbols/characters need to be treated as NA values:

```
read.csv("name_of_csv_file.csv", na.strings = "-")
```

It is also possible to indicate that more than one symbol needs to be read as NA:

```
read.csv('missing.csv', na.strings = c('.','-'))
```

Similarly, NAs can be written with customized strings using the `na` argument to `write.csv`. Other tools for reading and writing tables have similar options.

Section 66.3: Using NAs of different classes

The symbol NA is for a `logical` missing value:

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

This is convenient, since it can easily be coerced to other atomic vector types, and is therefore usually the only NA

you will need:

```
x <- c(1, NA, 1)
class(x[2])
#[1] "numeric"
```

If you do need a single NA value of another type, use NA_character_, NA_integer_, NA_real_ or NA_complex_. For missing values of fancy classes, subsetting with NA_integer_ usually works; for example, to get a missing-value Date:

```
class(Sys.Date())[NA_integer_]
# [1] "Date"
```

Section 66.4: TRUE/FALSE and/or NA

NA is a logical type and a logical operator with an NA will return NA if the outcome is ambiguous. Below, NA OR TRUE evaluates to TRUE because at least one side evaluates to TRUE, however NA OR FALSE returns NA because we do not know whether NA would have been TRUE or FALSE

```
NA | TRUE
# [1] TRUE
# TRUE | TRUE is TRUE and FALSE | TRUE is also TRUE.

NA | FALSE
# [1] NA
# TRUE | FALSE is TRUE but FALSE | FALSE is FALSE.

NA & TRUE
# [1] NA
# TRUE & TRUE is TRUE but FALSE & TRUE is FALSE.

NA & FALSE
# [1] FALSE
# TRUE & FALSE is FALSE and FALSE & FALSE is also FALSE.
```

These properties are helpful if you want to subset a data set based on some columns that contain NA.

```
df <- data.frame(v1=0:9,
                  v2=c(rep(1:2, each=4), NA, NA),
                  v3=c(NA, letters[2:10]))

df[df$v2 == 1 & !is.na(df$v2), ]
#   v1 v2   v3
#1   0  1 <NA>
#2   1  1     b
#3   2  1     c
#4   3  1     d

df[df$v2 == 1, ]
  v1 v2   v3
#1   0  1 <NA>
#2   1  1     b
#3   2  1     c
#4   3  1     d
#NA   NA NA <NA>
#NA.1 NA NA <NA>
```

Chapter 67: Hierarchical Linear Modeling

Section 67.1: basic model fitting

apologies: since I don't know of a channel for discussing/providing feedback on requests for improvement, I'm going to put my question here. Please feel free to point out a better place for this! @DataTx states that this is "completely unclear, incomplete, or has severe formatting problems". Since I don't see any big formatting problems (:-)), a little bit more guidance about what's expected here for improving clarity or completeness, and why what's here is unsalvageable, would be useful.

The primary packages for fitting hierarchical (alternatively "mixed" or "multilevel") linear models in R are `nlme` (older) and `lme4` (newer). These packages differ in many minor ways but should generally result in very similar fitted models.

```
library(nlme)
library(lme4)
m1.nlme <- lme(Reaction~Days, random=~Days|Subject, data=sleepstudy, method="REML")
m1.lme4 <- lmer(Reaction~Days+(Days|Subject), data=sleepstudy, REML=TRUE)
all.equal(fixef(m1.nlme), fixef(m1.lme4))
## [1] TRUE
```

Differences to consider:

- formula syntax is slightly different
- `nlme` is (still) somewhat better documented (e.g. Pinheiro and Bates 2000 *Mixed-effects models in S-PLUS*; however, see Bates *et al.* 2015 *Journal of Statistical Software*/[vignette\("lmer", package="lme4"\)](#) for `lme4`)
- `lme4` is faster and allows easier fitting of crossed random effects
- `nlme` provides p-values for linear mixed models out of the box, `lme4` requires add-on packages such as `lmerTest` or `afex`
- `nlme` allows modeling of heteroscedasticity or residual correlations (in space/time/phylogeny)

The unofficial [GLMM FAQ](#) provides more information, although it is focused on *generalized* linear mixed models (GLMMs).

Chapter 68: *apply family of functions (functionals)

Section 68.1: Using built-in functionals

Built-in functionals: lapply(), sapply(), and mapply()

R comes with built-in functionals, of which perhaps the most well-known are the apply family of functions. Here is a description of some of the most common apply functions:

- **lapply()** = takes a list as an argument and applies the specified function to the list.
- **sapply()** = the same as **lapply()** but attempts to simplify the output to a vector or a matrix.
 - **vapply()** = a variant of **sapply()** in which the output object's type must be specified.
- **mapply()** = like **lapply()** but can pass multiple vectors as input to the specified function. Can be simplified like **sapply()**.
 - **Map()** is an alias to **mapply()** with SIMPLIFY = FALSE.

lapply()

lapply() can be used with two different iterations:

- **lapply(variable, FUN)**
- **lapply(seq_along(variable), FUN)**

```
# Two ways of finding the mean of x
set.seed(1)
df <- data.frame(x = rnorm(25), y = rnorm(25))
lapply(df, mean)
lapply(seq_along(df), function(x) mean(df[[x]]))
```

sapply()

sapply() will attempt to resolve its output to either a vector or a matrix.

```
# Two examples to show the different outputs of sapply()
sapply(letters, print) ## produces a vector
x <- list(a = 1:10, beta = exp(-3:3), logic = c(TRUE, FALSE, FALSE, TRUE))
sapply(x, quantile) ## produces a matrix
```

mapply()

mapply() works much like **lapply()** except it can take multiple vectors as input (hence the m for multivariate).

```
mapply(sum, 1:5, 10:6, 3) # 3 will be "recycled" by mapply
```

Section 68.2: Combining multiple `data.frames` (`lapply`, `mapply`)

In this exercise, we will generate four bootstrap linear regression models and combine the summaries of these models into a single data frame.

```

library(broom)

#* Create the bootstrap data sets
BootData <- lapply(1:4,
  function(i) mtcars[sample(1:nrow(mtcars),
    size = nrow(mtcars),
    replace = TRUE), ])

#* Fit the models
Models <- lapply(BootData,
  function(BD) lm(mpg ~ qsec + wt + factor(am),
    data = BD))

#* Tidy the output into a data.frame
Tidied <- lapply(Models,
  tidy)

#* Give each element in the Tidied list a name
Tidied <- setNames(Tidied, paste0("Boot", seq_along(Tidied)))

```

At this point, we can take two approaches to inserting the names into the data.frame.

```

#* Insert the element name into the summary with `lapply`
#* Requires passing the names attribute to `lapply` and referencing `Tidied` within
#* the applied function.
Described_lapply <-
  lapply(names(Tidied),
    function(nm) cbind(nm, Tidied[[nm]]))

Combined_lapply <- do.call("rbind", Described_lapply)

#* Insert the element name into the summary with `mapply`
#* Allows us to pass the names and the elements as separate arguments.
Described_mapply <-
  mapply(
    function(nm, dframe) cbind(nm, dframe),
    names(Tidied),
    Tidied,
    SIMPLIFY = FALSE)

Combined_mapply <- do.call("rbind", Described_mapply)

```

If you're a fan of `magrittr` style pipes, you can accomplish the entire task in a single chain (though it may not be prudent to do so if you need any of the intermediary objects, such as the model objects themselves):

```

library(magrittr)
library(broom)
Combined <- lapply(1:4,
  function(i) mtcars[sample(1:nrow(mtcars),
    size = nrow(mtcars),
    replace = TRUE), ]) %>%
  lapply(function(BD) lm(mpg ~ qsec + wt + factor(am), data = BD)) %>%
  lapply(tidy) %>%
  setNames(paste0("Boot", seq_along(.))) %>%
  mapply(function(nm, dframe) cbind(nm, dframe),
    nm = names(.),
    dframe =.,
    SIMPLIFY = FALSE) %>%
  do.call("rbind", .)

```

Section 68.3: Bulk File Loading

for a large number of files which may need to be operated on in a similar process and with well structured file names.

firstly a vector of the file names to be accessed must be created, there are multiple options for this:

- Creating the vector manually with `paste0()`

```
files <- paste0("file_", 1:100, ".rds")
```

- Using `list.files()` with a regex search term for the file type, requires knowledge of regular expressions (regex) if other files of same type are in the directory.

```
files <- list.files("./", pattern = "\\.rds$", full.names = TRUE)
```

where X is a vector of part of the files naming format used.

`lapply` will output each response as element of a list.

`readRDS` is specific to .rds files and will change depending on the application of the process.

```
my_file_list <- lapply(files, readRDS)
```

This is not necessarily faster than a for loop from testing but allows all files to be an element of a list without assigning them explicitly.

Finally, we often need to load multiple packages at once. This trick can do it quite easily by applying `library()` to all libraries that we wish to import:

```
lapply(c("jsonlite", "stringr", "igraph"), library, character.only=TRUE)
```

Section 68.4: Using user-defined functionals

User-defined functionals

Users can create their own functionals to varying degrees of complexity. The following examples are from [Functionals](#) by Hadley Wickham:

```
randomise <- function(f) f(runif(1e3))

lapply2 <- function(x, f, ...) {
  out <- vector("list", length(x))
  for (i in seq_along(x)) {
    out[[i]] <- f(x[[i]], ...)
  }
  out
}
```

In the first case, `randomise` accepts a single argument `f`, and calls it on a sample of Uniform random variables. To demonstrate equivalence, we call `set.seed` below:

```
set.seed(123)
randomise(mean)
```

```
#[1] 0.4972778
```

```
set.seed(123)
mean(runif(1e3))
#[1] 0.4972778
```

```
set.seed(123)
randomise(max)
#[1] 0.9994045
```

```
set.seed(123)
max(runif(1e3))
#[1] 0.9994045
```

The second example is a re-implementation of base `:lapply`, which uses functionals to apply an operation (f) to each element in a list (x). The ... parameter allows the user to pass additional arguments to f, such as the `na.rm` option in the `mean` function:

```
lapply(list(c(1, 3, 5), c(2, NA, 6)), mean)
# [[1]]
# [1] 3
#
# [[2]]
# [1] NA

lapply2(list(c(1, 3, 5), c(2, NA, 6)), mean)
# [[1]]
# [1] 3
#
# [[2]]
# [1] NA

lapply(list(c(1, 3, 5), c(2, NA, 6)), mean, na.rm = TRUE)
# [[1]]
# [1] 3
#
# [[2]]
# [1] 4

lapply2(list(c(1, 3, 5), c(2, NA, 6)), mean, na.rm = TRUE)
# [[1]]
# [1] 3
#
# [[2]]
# [1] 4
```

Chapter 69: Text mining

Section 69.1: Scraping Data to build N-gram Word Clouds

The following example utilizes the `tm` text mining package to scrape and mine text data from the web to build word clouds with symbolic shading and ordering.

```
require(RWeka)
require(tau)
require(tm)
require(tm.plugin.webmining)
require(wordcloud)

# Scrape Google Finance -----
googlefinance <- WebCorpus(GoogleFinanceSource("NASDAQ:LFVN"))

# Scrape Google News -----
lv.googlenews <- WebCorpus(GoogleNewsSource("LifeVantage"))
p.googlenews <- WebCorpus(GoogleNewsSource("Protandim"))
ts.googlenews <- WebCorpus(GoogleNewsSource("TrueScience"))

# Scrape NYTimes -----
lv.nytimes <- WebCorpus(NYTimesSource(query = "LifeVantage", appid = nytimes_appid))
p.nytimes <- WebCorpus(NYTimesSource("Protandim", appid = nytimes_appid))
ts.nytimes <- WebCorpus(NYTimesSource("TrueScience", appid = nytimes_appid))

# Scrape Reuters -----
lv.reutersnews <- WebCorpus(ReutersNewsSource("LifeVantage"))
p.reutersnews <- WebCorpus(ReutersNewsSource("Protandim"))
ts.reutersnews <- WebCorpus(ReutersNewsSource("TrueScience"))

# Scrape Yahoo! Finance -----
lv.yahoofinance <- WebCorpus(YahooFinanceSource("LFVN"))

# Scrape Yahoo! News -----
lv.yahoonews <- WebCorpus(YahooNewsSource("LifeVantage"))
p.yahoonews <- WebCorpus(YahooNewsSource("Protandim"))
ts.yahoonews <- WebCorpus(YahooNewsSource("TrueScience"))

# Scrape Yahoo! Inplay -----
lv.yahooinplay <- WebCorpus(YahooInplaySource("LifeVantage"))

# Text Mining the Results -----
corpus <- c(googlefinance, lv.googlenews, p.googlenews, ts.googlenews, lv.yahoofinance,
lv.yahoonews, p.yahoonews,
ts.yahoonews, lv.yahooinplay) #lv.nytimes, p.nytimes, ts.nytimes, lv.reutersnews, p.reutersnews,
ts.reutersnews,

inspect(corpus)
wordlist <- c("lfvn", "lifevantage", "protandim", "truescience", "company", "fiscal", "nasdaq")

ds0.1g <- tm_map(corpus, content_transformer(tolower))
ds1.1g <- tm_map(ds0.1g, content_transformer(removeWords), wordlist)
ds1.1g <- tm_map(ds1.1g, content_transformer(removeWords), stopwords("english"))
ds2.1g <- tm_map(ds1.1g, stripWhitespace)
ds3.1g <- tm_map(ds2.1g, removePunctuation)
ds4.1g <- tm_map(ds3.1g, stemDocument)

tdm.1g <- TermDocumentMatrix(ds4.1g)
dtm.1g <- DocumentTermMatrix(ds4.1g)
```

```

findFreqTerms(tdm.1g, 40)
findFreqTerms(tdm.1g, 60)
findFreqTerms(tdm.1g, 80)
findFreqTerms(tdm.1g, 100)

findAssocs(dtm.1g, "skin", .75)
findAssocs(dtm.1g, "scienc", .5)
findAssocs(dtm.1g, "product", .75)

tdm89.1g <- removeSparseTerms(tdm.1g, 0.89)
tdm9.1g <- removeSparseTerms(tdm.1g, 0.9)
tdm91.1g <- removeSparseTerms(tdm.1g, 0.91)
tdm92.1g <- removeSparseTerms(tdm.1g, 0.92)

tdm2.1g <- tdm92.1g

# Creates a Boolean matrix (counts # docs w/terms, not raw # terms)
tdm3.1g <- inspect(tdm2.1g)
tdm3.1g[tdm3.1g>=1] <- 1

# Transform into a term-term adjacency matrix
termMatrix.1gram <- tdm3.1g %*% t(tdm3.1g)

# inspect terms numbered 5 to 10
termMatrix.1gram[5:10,5:10]
termMatrix.1gram[1:10,1:10]

# Create a WordCloud to Visualize the Text Data -----
notsparse <- tdm2.1g
m = as.matrix(notsparse)
v = sort(rowSums(m),decreasing=TRUE)
d = data.frame(word = names(v),freq=v)

# Create the word cloud
pal = brewer.pal(9,"BuPu")
wordcloud(words = d$word,
          freq = d$freq,
          scale = c(3,.8),
          random.order = F,
          colors = pal)

```

Note the use of `random.order` and a sequential pallet from `RColorBrewer`, which allows the programmer to capture more information in the cloud by assigning meaning to the order and coloring of terms.

Above is the 1-gram case.

We can make a major leap to n-gram word clouds and in doing so we'll see how to make almost any text-mining analysis flexible enough to handle n-grams by transforming our TDM.

The initial difficulty you run into with n-grams in R is that `tm`, the most popular package for text mining, does not inherently support tokenization of bi-grams or n-grams. Tokenization is the process of representing a word, part of a word, or group of words (or symbols) as a single data element called a token.

Fortunately, we have some hacks which allow us to continue using `tm` with an upgraded tokenizer. There's more than one way to achieve this. We can write our own simple tokenizer using the `textcnt()` function from `tau`:

```
tokenize_ngrams <- function(x, n=3)
return(rownames(as.data.frame(unclass(textcnt(x, method="string", n=n)))))
```

or we can invoke `RWeka`'s tokenizer within `tm`:

```
# BigramTokenizer
BigramTokenizer <- function(x) NGramTokenizer(x, Weka_control(min = 2, max = 2))
```

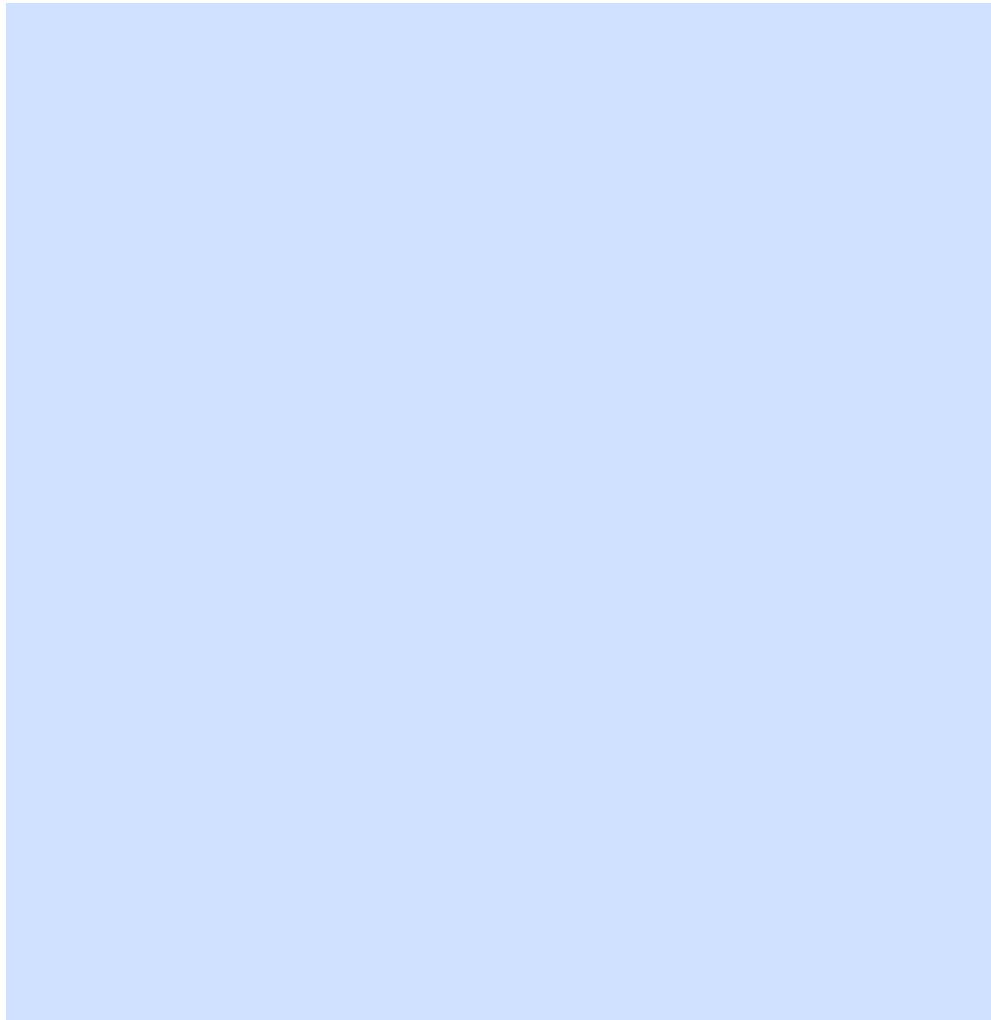
From this point you can proceed much as in the 1-gram case:

```
# Create an n-gram Word Cloud -----
tdm.ng <- TermDocumentMatrix(ds5.1g, control = list(tokenize = BigramTokenizer))
dtm.ng <- DocumentTermMatrix(ds5.1g, control = list(tokenize = BigramTokenizer))

# Try removing sparse terms at a few different levels
tdm89.ng <- removeSparseTerms(tdm.ng, 0.89)
tdm9.ng <- removeSparseTerms(tdm.ng, 0.9)
tdm91.ng <- removeSparseTerms(tdm.ng, 0.91)
tdm92.ng <- removeSparseTerms(tdm.ng, 0.92)
```

```
notsparse <- tdm91.ng
m = as.matrix(notsparse)
v = sort(rowSums(m),decreasing=TRUE)
d = data.frame(word = names(v),freq=v)

# Create the word cloud
pal = brewer.pal(9,"BuPu")
wordcloud(words = d$word,
freq = d$freq,
scale = c(3,.8),
random.order = F,
colors = pal)
```



The example above is [reproduced](#) with permission from Hack-R's data science blog. Additional commentary may be found in the original article.

Chapter 70: ANOVA

Section 70.1: Basic usage of aov()

Analysis of Variance (aov) is used to determine if the means of two or more groups differ significantly from each other. Responses are assumed to be independent of each other, Normally distributed (within each group), and the within-group variances are assumed equal.

In order to complete the analysis data must be in long format (see reshaping data topic). `aov()` is a wrapper around the `lm()` function, using Wilkinson-Rogers formula notation $y \sim f$ where y is the response (independent) variable and f is a factor (categorical) variable representing group membership. *If f is numeric rather than a factor variable, `aov()` will report the results of a linear regression in ANOVA format, which may surprise inexperienced users.*

The `aov()` function uses Type I (sequential) Sum of Squares. This type of Sum of Squares tests all of the (main and interaction) effects sequentially. The result is that the first effect tested is also assigned shared variance between it and other effects in the model. For the results from such a model to be reliable, data should be balanced (all groups are of the same size).

When the assumptions for Type I Sum of Squares do not hold, Type II or Type III Sum of Squares may be applicable. Type II Sum of Squares test each main effect after every other main effect, and thus controls for any overlapping variance. However, Type II Sum of Squares assumes no interaction between the main effects.

Lastly, Type III Sum of Squares tests each main effect after every other main effect *and* every interaction. This makes Type III Sum of Squares a necessity when an interaction is present.

Type II and Type III Sums of Squares are implemented in the `Anova()` function.

Using the `mtcars` data set as an example.

```
mtCarsAnovaModel <- aov(wt ~ factor(cyl), data=mtcars)
```

To view summary of ANOVA model:

```
summary(mtCarsAnovaModel)
```

One can also extract the coefficients of the underlying `lm()` model:

```
coefficients(mtCarsAnovaModel)
```

Section 70.2: Basic usage of Anova()

When dealing with an unbalanced design and/or non-orthogonal contrasts, Type II or Type III Sum of Squares are necessary. The `Anova()` function from the `car` package implements these. Type II Sum of Squares assumes no interaction between main effects. If interactions are assumed, Type III Sum of Squares is appropriate.

The `Anova()` function wraps around the `lm()` function.

Using the `mtcars` data sets as an example, demonstrating the difference between Type II and Type III when an interaction is tested.

```
> Anova(lm(wt ~ factor(cyl)*factor(am), data=mtcars), type = 2)
Anova Table (Type II tests)
```

```
Response: wt
          Sum Sq Df F value    Pr(>F)
factor(cyl)   7.2278  2 11.5266 0.0002606 ***
factor(am)    3.2845  1 10.4758 0.0032895 **
factor(cyl):factor(am) 0.0668  2  0.1065 0.8993714
Residuals     8.1517 26
```

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

```
> Anova(lm(wt ~ factor(cyl)*factor(am), data=mtcars), type = 3)
Anova Table (Type III tests)
```

```
Response: wt
          Sum Sq Df F value    Pr(>F)
(Intercept) 25.8427  1 82.4254 1.524e-09 ***
factor(cyl)  4.0124  2  6.3988 0.005498 **
factor(am)   1.7389  1  5.5463 0.026346 *
factor(cyl):factor(am) 0.0668  2  0.1065 0.899371
Residuals    8.1517 26
```

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

Chapter 71: Raster and Image Analysis

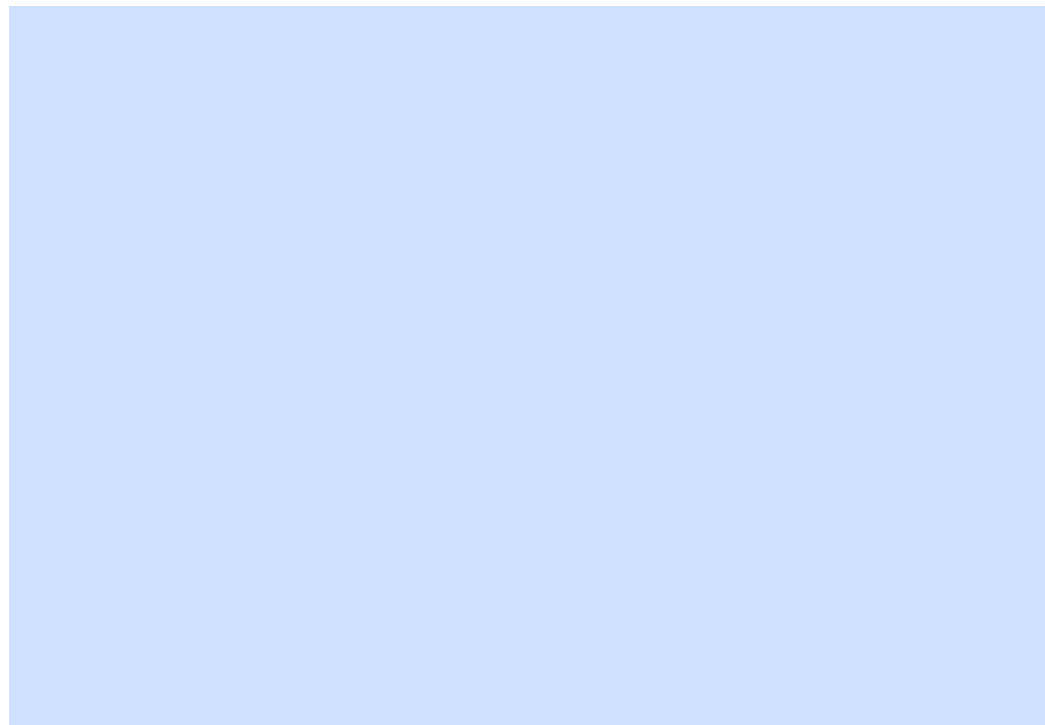
See also I/O for Raster Images

Section 71.1: Calculating GLCM Texture

[Gray Level Co-Occurrence Matrix](#) (Haralick et al. 1973) texture is a powerful image feature for image analysis. The `glcm` package provides a easy-to-use function to calculate such textrual features for `RasterLayer` objects in R.

```
library(glcm)
library(raster)

r <- raster("C:/Program Files/R/R-3.2.3/doc/html/logo.jpg")
plot(r)
```



Calculating GLCM textures in one direction

```
rglcm <- glcm(r,
  window = c(9,9),
  shift = c(1,1),
  statistics = c("mean", "variance", "homogeneity", "contrast",
    "dissimilarity", "entropy", "second_moment")
)
plot(rglcm)
```



Calculation rotation-invariant texture features

The textural features can also be calculated in all 4 directions (0° , 45° , 90° and 135°) and then combined to one rotation-invariant texture. The key for this is the `shift` parameter:

```
rglcm1 <- glcm(r,
  window = c(9,9),
  shift=list(c(0,1), c(1,1), c(1,0), c(1,-1)),
  statistics = c("mean", "variance", "homogeneity", "contrast",
    "dissimilarity", "entropy", "second_moment"))
)
plot(rglcm1)
```



Section 71.2: Mathematical Morphologies

The package `mmand` provides functions for the calculation of Mathematical Morphologies for n-dimensional arrays. With a little workaround, these can also be calculated for raster images.

```
library(raster)
library(mmand)

r <- raster("C:/Program Files/R/R-3.2.3/doc/html/logo.jpg")
plot(r)
```

At first, a kernel (moving window) has to be set with a size (e.g. 9x9) and a shape type (e.g. disc, box or diamond)

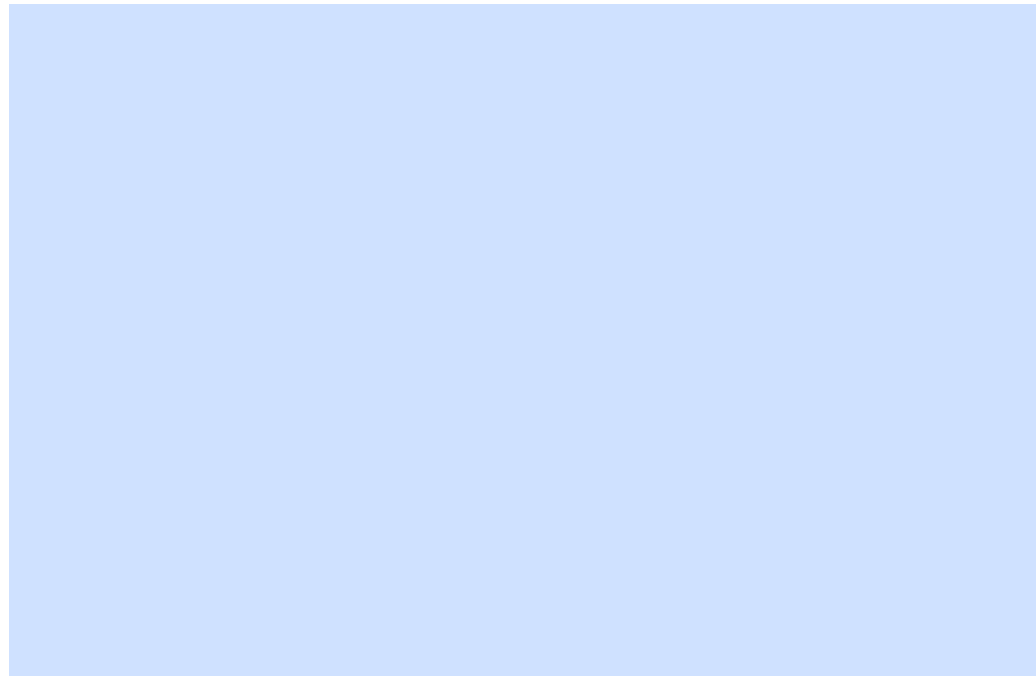
```
sk <- shapeKernel(c(9,9), type="disc")
```

Afterwards, the raster layer has to be converted into an array which is used as input for the erode() function.

```
rArr <- as.array(r, transpose = TRUE)
rErode <- erode(rArr, sk)
rErode <- setValues(r, as.vector(aperm(rErode)))
```

Besides erode(), also the morphological functions dilate(), opening() and closing() can be applied like this.

```
plot(rErode)
```



Chapter 72: Survival analysis

Section 72.1: Random Forest Survival Analysis with randomForestSRC

Just as the [random forest](#) algorithm may be applied to regression and classification tasks, it can also be extended to survival analysis.

In the example below a survival model is fit and used for prediction, scoring, and performance analysis using the package `randomForestSRC` [from CRAN](#).

```
require(randomForestSRC)

set.seed(130948) #Other seeds give similar comparative results
x1    <- runif(1000)
y     <- rnorm(1000, mean = x1, sd = .3)
data <- data.frame(x1 = x1, y = y)
head(data)
```

| x1 | y |
|-------------|------------|
| 1 0.9604353 | 1.3549648 |
| 2 0.3771234 | 0.2961592 |
| 3 0.7844242 | 0.6942191 |
| 4 0.9860443 | 1.5348900 |
| 5 0.1942237 | 0.4629535 |
| 6 0.7442532 | -0.0672639 |

```
(modRFSRC <- rfsrc(y ~ x1, data = data, ntree=500, nodesize = 5))
```

```
Sample size: 1000
Number of trees: 500
Minimum terminal node size: 5
Average no. of terminal nodes: 208.258
No. of variables tried at each split: 1
Total no. of variables: 1
Analysis: RF-R
Family: regr
Splitting rule: mse
% variance explained: 32.08
Error rate: 0.11
```

```
x1new    <- runif(10000)
ynew      <- rnorm(10000, mean = x1new, sd = .3)
newdata <- data.frame(x1 = x1new, y = ynew)

survival.results <- predict(modRFSRC, newdata = newdata)
survival.results
```

```
Sample size of test (predict) data: 10000
Number of grow trees: 500
Average no. of grow terminal nodes: 208.258
Total no. of grow variables: 1
Analysis: RF-R
Family: regr
% variance explained: 34.97
Test set error rate: 0.11
```




Section 72.3: Kaplan Meier estimates of survival curves and risk set tables with survminer

Base plot

```
install.packages('survminer')
source("https://bioconductor.org/biocLite.R")
biocLite("RTCGA.clinical") # data for examples
library(RTCGA.clinical)
survivalTCGA(BRCA.clinical, OV.clinical,
             extract.cols = "admin.disease_code") -> BRCAOV.survInfo
library(survival)
fit <- survfit(Surv(times, patient.vital_status) ~ admin.disease_code,
                data = BRCAOV.survInfo)
library(survminer)
ggsurvplot(fit, risk.table = TRUE)
```

More advanced

```
ggsurvplot(  
  fit,                      # survfit object with calculated statistics.  
  risk.table = TRUE,          # show risk table.  
  pval = TRUE,                # show p-value of log-rank test.  
  conf.int = TRUE,             # show confidence intervals for  
                            # point estimates of survival curves.  
  xlim = c(0,2000),            # present narrower X axis, but not affect  
                            # survival estimates.  
  break.time.by = 500,          # break X axis in time intervals by 500.  
  ggtheme = theme_RTCGA(),     # customize plot and risk table with a theme.  
  risk.table.y.text.col = T,    # colour risk table text annotations.  
  risk.table.y.text = FALSE,    # show bars instead of names in text annotations  
                            # in legend of risk table  
)
```



Based on

<http://r-addict.com/2016/05/23/Informative-Survival-Plots.html>

Chapter 73: Fault-tolerant/resilient code

| Parameter | Details |
|-------------------|---|
| expr | In case the "try part" was completed successfully tryCatch will return the last evaluated expression . Hence, the actual value being returned in case everything went well and there is no condition (i.e. a <i>warning</i> or an <i>error</i>) is the return value of readLines . Note that you don't need to explicitly state the return value via return as code in the "try part" is not wrapped inside a function environment (unlike that for the condition handlers for warnings and error below) |
| warning/error/etc | Provide/define a handler function for all the conditions that you want to handle explicitly. AFAIU, you can provide handlers for <i>any</i> type of conditions (not just <i>warnings</i> and <i>errors</i> , but also <i>warning/error/etc custom</i> conditions; see simpleCondition and friends for that) as long as the name of the respective handler function matches the class of the respective condition (see the <i>Details</i> part of the doc for tryCatch). |
| finally | Here goes everything that should be executed at the very end, regardless if the expression in the "try part" succeeded or if there was any condition. If you want more than one expression to be executed, then you need to wrap them in curly brackets, otherwise you could just have written <code>finally = <expression></code> (i.e. the same logic as for "try part"). |

Section 73.1: Using tryCatch()

We're defining a robust version of a function that reads the HTML code from a given URL. *Robust* in the sense that we want it to handle situations where something either goes wrong (*error*) or not quite the way we planned it to (*warning*). The umbrella term for errors and warnings is *condition*

Function definition using **tryCatch**

```
readUrl <- function(url) {
  out <- tryCatch(
    #####
    # Try part: define the expression(s) you want to "try" #
    #####
    {

      # Just to highlight:
      # If you want to use more than one R expression in the "try part"
      # then you'll have to use curly brackets.
      # Otherwise, just write the single expression you want to try and

      message("This is the 'try' part")
      readLines(con = url, warn = FALSE)
    },
    #####
    # Condition handler part: define how you want conditions to be handled #
    #####
    # Handler when a warning occurs:
    warning = function(cond) {
      message(paste("Reading the URL caused a warning:", url))
      message("Here's the original warning message:")
      message(cond)

      # Choose a return value when such a type of condition occurs
      return(NULL)
    },
    # Handler when an error occurs:
    error = function(cond) {
```

```

    message(paste("This seems to be an invalid URL:", url))
    message("Here's the original error message:")
    message(cond)

    # Choose a return value when such a type of condition occurs
    return(NA)
}

#####
# Final part: define what should happen AFTER #
# everything has been tried and/or handled   #
#####

finally = {
  message(paste("Processed URL:", url))
  message("Some message at the end\n")
}
)
return(out)
}

```

Testing things out

Let's define a vector of URLs where one element isn't a valid URL

```

urls <- c(
  "http://stat.ethz.ch/R-manual/R-devel/library/base/html/connections.html",
  "http://en.wikipedia.org/wiki/Xz",
  "I'm no URL"
)

```

And pass this as input to the function we defined above

```

y <- lapply(urls, readUrl)
# Processed URL: http://stat.ethz.ch/R-manual/R-devel/library/base/html/connections.html
# Some message at the end
#
# Processed URL: http://en.wikipedia.org/wiki/Xz
# Some message at the end
#
# URL does not seem to exist: I'm no URL
# Here's the original error message:
# cannot open the connection
# Processed URL: I'm no URL
# Some message at the end
#
# Warning message:
# In file(con, "r") : cannot open file 'I'm no URL': No such file or directory

```

Investigating the output

```

length(y)
# [1] 3

head(y[[1]])
# [1] "<!DOCTYPE html PUBLIC "-//W3C//DTD HTML 4.01 Transitional//EN">"
# [2] "<html><head><title>R: Functions to Manipulate Connections</title>"
# [3] "<meta http-equiv=\"Content-Type\" content=\"text/html; charset=utf-8\">"
# [4] "<link rel=\"stylesheet\" type=\"text/css\" href=\"R.css\">"
# [5] "</head><body>"

```

```
# [6] "
```

```
y[[3]]
```

```
# [1] NA
```

Chapter 74: Reproducible R

With 'Reproducibility' we mean that someone else (perhaps you in the future) can repeat the steps you performed and get the same result. See the [Reproducible Research Task View](#).

Section 74.1: Data reproducibility

`dput()` and `dget()`

The easiest way to share a (preferable small) data frame is to use a basic function `dput()`. It will export an R object in a plain text form.

Note: Before making the example data below, make sure you're in an empty folder you can write to. Run `getwd()` and read `?setwd` if you need to change folders.

```
dput(mtcars, file = 'df.txt')
```

Then, anyone can load the precise R object to their GlobalEnvironment using the `dget()` function.

```
df <- dget('df.txt')
```

For larger R objects, there are a number of ways of saving them reproducibly. See [Input and output](#) .

Section 74.2: Package reproducibility

Package reproducibility is a very common issue in reproducing some R code. When various packages get updated, some interconnections between them may break. The ideal solution for the problem is to reproduce the image of the R code writer's machine on your computer at the date when the code was written. And here comes `checkpoint` package.

Starting from 2014-09-17, the authors of the package make daily copies of the whole CRAN package repository to their own mirror repository -- Microsoft R Archived Network. So, to avoid package reproduciblity issues when creating a reproducible R project, all you need is to:

1. Make sure that all your packages (and R version) are up-to-date.
2. Include `checkpoint::checkpoint('YYYY-MM-DD')` line in your code.

`checkpoint` will create a directory `.checkpoint` in your R_home directory (`"~/"`). To this technical directory it will install all the packages, that are used in your project. That means, `checkpoint` looks through all the `.R` files in your project directory to pick up all the `library()` or `require()` calls and install all the required packages in the form they existed at CRAN on the specified date.

PRO You are freed from the package reproducibility issue.

CONTRA For each specified date you have to download and install all the packages that are used in a certain project that you aim to reproduce. That may take quite a while.

Chapter 75: Fourier Series and Transformations

The Fourier transform decomposes a function of time (a signal) into the frequencies that make it up, similarly to how a musical chord can be expressed as the amplitude (or loudness) of its constituent notes. The Fourier transform of a function of time itself is a complex-valued function of frequency, whose absolute value represents the amount of that frequency present in the original function, and whose complex argument is the phase offset of the basic sinusoid in that frequency.

The Fourier transform is called the frequency domain representation of the original signal. The term Fourier transform refers to both the frequency domain representation and the mathematical operation that associates the frequency domain representation to a function of time. The Fourier transform is not limited to functions of time, but in order to have a unified language, the domain of the original function is commonly referred to as the time domain. For many functions of practical interest one can define an operation that reverses this: the inverse Fourier transformation, also called Fourier synthesis, of a frequency domain representation combines the contributions of all the different frequencies to recover the original function of time.

Linear operations performed in one domain (time or frequency) have corresponding operations in the other domain, which are sometimes easier to perform. The operation of differentiation in the time domain corresponds to multiplication by the frequency, so some differential equations are easier to analyze in the frequency domain. Also, convolution in the time domain corresponds to ordinary multiplication in the frequency domain. Concretely, this means that any linear time-invariant system, such as an electronic filter applied to a signal, can be expressed relatively simply as an operation on frequencies. So significant simplification is often achieved by transforming time functions to the frequency domain, performing the desired operations, and transforming the result back to time.

Harmonic analysis is the systematic study of the relationship between the frequency and time domains, including the kinds of functions or operations that are "simpler" in one or the other, and has deep connections to almost all areas of modern mathematics.

Functions that are localized in the time domain have Fourier transforms that are spread out across the frequency domain and vice versa. The critical case is the Gaussian function, of substantial importance in probability theory and statistics as well as in the study of physical phenomena exhibiting normal distribution (e.g., diffusion), which with appropriate normalizations goes to itself under the Fourier transform. Joseph Fourier introduced the transform in his study of heat transfer, where Gaussian functions appear as solutions of the heat equation.

The Fourier transform can be formally defined as an improper Riemann integral, making it an integral transform, although this definition is not suitable for many applications requiring a more sophisticated integration theory.

For example, many relatively simple applications use the Dirac delta function, which can be treated formally as if it were a function, but the justification requires a mathematically more sophisticated viewpoint. The Fourier transform can also be generalized to functions of several variables on Euclidean space, sending a function of 3-dimensional space to a function of 3-dimensional momentum (or a function of space and time to a function of 4-momentum).

This idea makes the spatial Fourier transform very natural in the study of waves, as well as in quantum mechanics, where it is important to be able to represent wave solutions either as functions either of space or momentum and sometimes both. In general, functions to which Fourier methods are applicable are complex-valued, and possibly vector-valued. Still further generalization is possible to functions on groups, which, besides the original Fourier transform on \mathbb{R} or \mathbb{R}^n (viewed as groups under addition), notably includes the discrete-time Fourier transform (DTFT, group = \mathbb{Z}), the discrete Fourier transform (DFT, group = $\mathbb{Z} \text{ mod } N$) and the Fourier series or circular Fourier transform (group = S^1 , the unit circle ≈ closed finite interval with endpoints identified). The latter is routinely

employed to handle periodic functions. The Fast Fourier transform (FFT) is an algorithm for computing the DFT.

Section 75.1: Fourier Series

Joseph Fourier showed that any periodic wave can be represented by a sum of simple sine waves. This sum is called the Fourier Series. The Fourier Series only holds while the system is linear. If there is, eg, some overflow effect (a threshold where the output remains the same no matter how much input is given), a non-linear effect enters the picture, breaking the sinusoidal wave and the superposition principle.

```
# Sine waves
xs <- seq(-2*pi, 2*pi, pi/100)
wave.1 <- sin(3*xs)
wave.2 <- sin(10*xs)
par(mfrow = c(1, 2))
plot(xs, wave.1, type="l", ylim=c(-1,1)); abline(h=0, lty=3)
plot(xs, wave.2, type="l", ylim=c(-1,1)); abline(h=0, lty=3)

# Complex Wave
wave.3 <- 0.5 * wave.1 + 0.25 * wave.2
plot(xs, wave.3, type="l"); title("Eg complex wave"); abline(h=0, lty=3)
```

```
wave.4 <- wave.3  
wave.4[wave.3>0.5] <- 0.5  
plot(xs,wave.4,type="l",ylim=c(-1.25,1.25))  
title("overflowed, non-linear complex wave")  
abline(h=0,lty=3)
```



Also, the Fourier Series only holds if the waves are periodic, ie, they have a repeating pattern (non periodic waves are dealt by the Fourier Transform, see below). A periodic wave has a frequency f and a wavelength λ (a wavelength is the distance in the medium between the beginning and end of a cycle, $\lambda=v/f_0$, where v is the wave velocity) that are defined by the repeating pattern. A non-periodic wave does not have a frequency or wavelength.

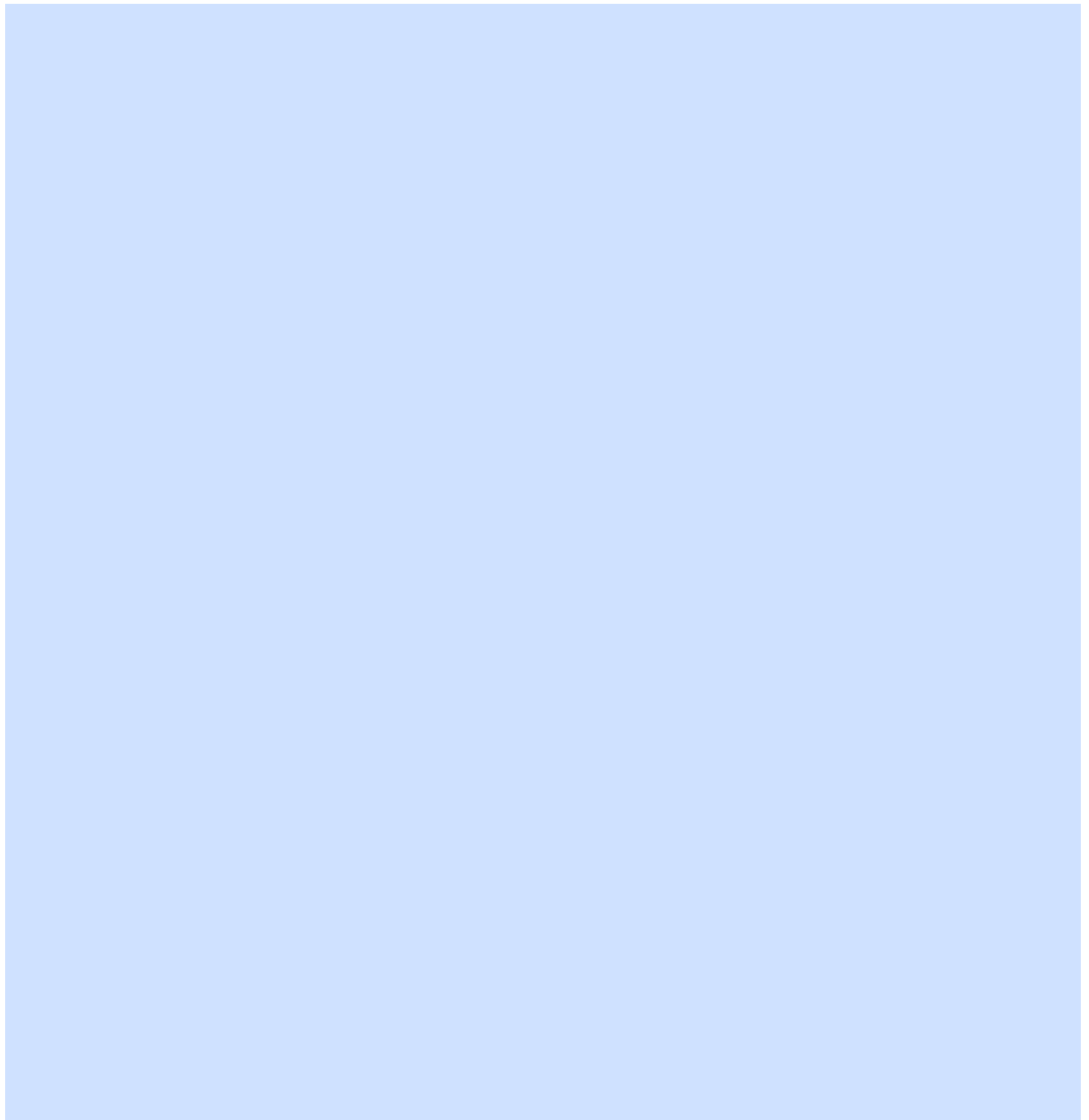
Some concepts:

- The fundamental period, T , is the period of all the samples taken, the time between the first sample and the last
- The sampling rate, sr , is the number of samples taken over a time period (aka acquisition frequency). For simplicity we will make the time interval between samples equal. This time interval is called the sample interval, si , which is the fundamental period time divided by the number of samples N . So, $si=T/N$
- The fundamental frequency, f_0 , which is $1/T$. The fundamental frequency is the frequency of the repeating pattern or how long the wavelength is. In the previous waves, the fundamental frequency was 12π . The

frequencies of the wave components must be integer multiples of the fundamental frequency. f0 is called the first harmonic, the second harmonic is 2*f0, the third is 3*f0, etc.

```
repeat.xs      <- seq(-2*pi, 0, pi/100)
wave.3.repeat <- 0.5*sin(3*repeat.xs) + 0.25*sin(10*repeat.xs)
plot(xs, wave.3, type="l")

title("Repeating pattern")
points(repeat.xs, wave.3.repeat, type="l", col="red");
abline(h=0, v=c(-2*pi, 0), lty=3)
```



Here's a R function for plotting trajectories given a fourier series:

```
plot.fourier <- function(fourier.series, f.0, ts) {
```

```
w <- 2*pi*f.0 trajectory <- sapply(ts, function(t) fourier.series(t,w))
plot(ts, trajectory, type="l", xlab="time", ylab="f(t)");
abline(h=0,lty=3)}
```

Chapter 76: .Rprofile

Section 76.1: .Rprofile - the first chunk of code executed

.Rprofile is a file containing R code that is executed when you launch R from the directory containing the .Rprofile file. The similarly named Rprofile.site, located in R's home directory, is executed by default every time you load R from any directory. Rprofile.site and to a greater extend .Rprofile can be used to initialize an R session with personal preferences and various utility functions that you have defined.

Important note: if you use RStudio, you can have a separate .Rprofile in every RStudio project directory.

Here are some examples of code that you might include in an .Rprofile file.

Setting your R home directory

```
# set R_home
Sys.setenv(R_USER="c:/R_home") # just an example directory
# but don't confuse this with the $R_HOME environment variable.
```

Setting page size options

```
options(papersize="a4")
options(editor="notepad")
options(pager="internal")
```

set the default help type

```
options(help_type="html")
```

set a site library

```
.Library.site <- file.path(chartr("\\", "/", R.home()), "site-library")
```

Set a CRAN mirror

```
local({r <- getOption("repos")
       r["CRAN"] <- "http://my.local.cran"
       options(repos=r)})
```

Setting the location of your library

This will allow you to not have to install all the packages again with each R version update.

```
# library location
.libPaths("c:/R_home/Rpackages/win")
```

Custom shortcuts or functions

Sometimes it is useful to have a shortcut for a long R expression. A common example of this setting an active binding to access the last top-level expression result without having to type out .Last.value:

```
makeActiveBinding(".", function(){.Last.value}, .GlobalEnv)
```

Because .Rprofile is just an R file, it can contain any arbitrary R code.

Pre-loading the most useful packages

This is bad practice and should generally be avoided because it separates package loading code from the scripts where those packages are actually used.

See Also

See `help(Startup)` for all the different startup scripts, and further aspects. In particular, two system-wide Profile files can be loaded as well. The first, Rprofile, may contain global settings, the other file Profile.site may contain local choices the system administrator can make for all users. Both files are found in the `$(RHOME)/etc` directory of the R installation. This directory also contains global files Renviron and Renviron.site which both can be complemented with a local file `~/.Renviron` in the user's home directory.

Section 76.2: .Rprofile example

Startup

```
# Load library setwidth on start - to set the width automatically.  
.First <- function() {  
  library(setwidth)  
  # If 256 color terminal - use library colorout.  
  if (Sys.getenv("TERM") %in% c("xterm-256color", "screen-256color")) {  
    library("colorout")  
  }  
}
```

Options

```
# Select default CRAN mirror for package installation.  
options(repos=c(CRAN="https://cran.gis-lab.info/"))  
  
# Print maximum 1000 elements.  
options(max.print=1000)  
  
# No scientific notation.  
options(scipen=10)  
  
# No graphics in menus.  
options(menu.graphics=FALSE)  
  
# Auto-completion for package names.  
utils::rc.settings(ipck=TRUE)
```

Custom Functions

```
# Invisible environment to mask defined functions  
.env = new.env()  
  
# Quit R without asking to save.  
.env$q <- function (save="no", ...) {  
  quit(save=save, ...)  
}  
  
# Attach the environment to enable functions.  
attach(.env, warn.conflicts=FALSE)
```

Chapter 77: dplyr

Section 77.1: dplyr's single table verbs

[dplyr](#) introduces a grammar of data manipulation in R. It provides a consistent interface to work with data no matter where it is stored: `data.frame`, `data.table`, or a database. The key pieces of `dplyr` are written using `Rcpp`, which makes it very fast for working with in-memory data.

`dplyr`'s philosophy is to have small functions that do one thing well. The five simple functions ([filter](#), `arrange`, [SELECT](#), `mutate`, and `summarise`) can be used to reveal new ways to describe data. When combined with `group_by`, these functions can be used to calculate group wise summary statistics.

Syntax commonalities

All these functions have a similar syntax:

- The first argument to all these functions is always a data frame
- Columns can be referred directly using bare variable names (i.e., without using `$`)
- These functions do not modify the original data itself, i.e., they don't have side effects. Hence, the results should always be saved to an object.

We will use the built-in [mtcars](#) dataset to explore `dplyr`'s single table verbs. Before converting the type of [mtcars](#) to `tbl_df` (since it makes printing cleaner), we add the [rownames](#) of the dataset as a column using `rownames_to_column` function from the [tibble](#) package.

```
library(dplyr) # This documentation was written using version 0.5.0

mtcars_tbl <- as_data_frame(tibble::rownames_to_column(mtcars, "cars"))

# examine the structure of data
head(mtcars_tbl)

# A tibble: 6 x 12
#   cars     mpg cyl disp  hp drat    wt  qsec vs am gear carb
#   <chr> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl>
#1 Mazda RX4  21.0     6   160   110  3.90 2.620 16.46    0   1     4     4
#2 Mazda RX4 Wag 21.0     6   160   110  3.90 2.875 17.02    0   1     4     4
#3 Datsun 710  22.8     4   108    93  3.85 2.320 18.61    1   1     4     1
#4 Hornet 4 Drive 21.4     6   258   110  3.08 3.215 19.44    1   0     3     1
#5 Hornet Sportabout 18.7     8   360   175  3.15 3.440 17.02    0   0     3     2
#6 Valiant    18.1     6   225   105  2.76 3.460 20.22    1   0     3     1
```

filter

[filter](#) helps subset rows that match certain criteria. The first argument is the name of the `data.frame` and the second (and subsequent) arguments are the criteria that filter the data (these criteria should evaluate to either TRUE or FALSE)

Subset all cars that have 4 *cylinders* - cyl:

```
filter(mtcars_tbl, cyl == 4)

# A tibble: 11 x 12
#   cars     mpg cyl disp  hp drat    wt  qsec vs am gear carb
#   <chr> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl>
#1 Datsun 710  22.8     4 108.0    93  3.85 2.320 18.61    1   1     4     1
#2 Merc 240D   24.4     4 146.7    62  3.69 3.190 20.00    1   0     4     2
```

```
#3      Merc 230 22.8      4 140.8    95 3.92 3.150 22.90      1     0     4     2
#4      Fiat 128 32.4      4 78.7    66 4.08 2.200 19.47      1     1     4     1
#5      Honda Civic 30.4      4 75.7    52 4.93 1.615 18.52      1     1     4     2
# ... with 6 more rows
```

We can pass multiple criteria separated by a comma. To subset the cars which have either 4 or 6 *cylinders* - cyl and have 5 *gears* - gear:

```
filter(mtcars_tbl, cyl == 4 | cyl == 6, gear == 5)

# A tibble: 3 x 12
#       cars   mpg   cyl   disp    hp   drat    wt   qsec    vs    am   gear   carb
#   <chr> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl>
#1 Porsche 914-2 26.0     4 120.3    91 4.43 2.140 16.7     0     1     5     2
#2 Lotus Europa 30.4     4 95.1     113 3.77 1.513 16.9     1     1     5     2
#3 Ferrari Dino 19.7     6 145.0    175 3.62 2.770 15.5     0     1     5     6
```

`filter` selects rows based on criteria, to select rows by position, use `slice`. `slice` takes only 2 arguments: the first one is a `data.frame` and the second is integer row values.

To select rows 6 through 9:

```
slice(mtcars_tbl, 6:9)

# A tibble: 4 x 12
#       cars   mpg   cyl   disp    hp   drat    wt   qsec    vs    am   gear   carb
#   <chr> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl>
#1 Valiant 18.1     6 225.0    105 2.76 3.46 20.22     1     0     3     1
#2 Duster 360 14.3     8 360.0    245 3.21 3.57 15.84     0     0     3     4
#3 Merc 240D 24.4     4 146.7     62 3.69 3.19 20.00     1     0     4     2
#4 Merc 230 22.8     4 140.8    95 3.92 3.15 22.90     1     0     4     2
```

Or:

```
slice(mtcars_tbl, -c(1:5, 10:n()))
```

This results in the same output as `slice(mtcars_tbl, 6:9)`

`n()` represents the number of observations in the current group

arrange

`arrange` is used to sort the data by a specified variable(s). Just like the previous verb (and all other functions in `dplyr`), the first argument is a `data.frame`, and consequent arguments are used to sort the data. If more than one variable is passed, the data is first sorted by the first variable, and then by the second variable, and so on..

To order the data by *horsepower* - hp

```
arrange(mtcars_tbl, hp)

# A tibble: 32 x 12
#       cars   mpg   cyl   disp    hp   drat    wt   qsec    vs    am   gear   carb
#   <chr> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl>
#1 Honda Civic 30.4     4 75.7    52 4.93 1.615 18.52     1     1     4     2
#2 Merc 240D 24.4     4 146.7    62 3.69 3.190 20.00     1     0     4     2
#3 Toyota Corolla 33.9     4 71.1    65 4.22 1.835 19.90     1     1     4     1
#4 Fiat 128 32.4     4 78.7    66 4.08 2.200 19.47     1     1     4     1
#5 Fiat X1-9 27.3     4 79.0    66 4.08 1.935 18.90     1     1     4     1
```

```
#6  Porsche 914-2 26.0      4 120.3     91   4.43 2.140 16.70      0     1     5     2
# ... with 26 more rows
```

To arrange the data by *miles per gallon* - mpg in descending order, followed by *number of cylinders* - cyl:

```
arrange(mtcars_tbl, desc(mpg), cyl)

# A tibble: 32 x 12
#       cars   mpg   cyl   disp    hp   drat    wt   qsec    vs    am   gear   carb
#       <chr> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl>
#1 Toyota Corolla 33.9     4  71.1    65  4.22 1.835 19.90     1     1     4     1
#2 Fiat 128     32.4     4  78.7    66  4.08 2.200 19.47     1     1     4     1
#3 Honda Civic  30.4     4  75.7    52  4.93 1.615 18.52     1     1     4     2
#4 Lotus Europa 30.4     4  95.1   113  3.77 1.513 16.90     1     1     5     2
#5 Fiat X1-9    27.3     4  79.0    66  4.08 1.935 18.90     1     1     4     1
#6 Porsche 914-2 26.0     4 120.3     91   4.43 2.140 16.70      0     1     5     2
# ... with 26 more rows
```

select

SELECT is used to select only a subset of variables. To select only mpg, disp, wt, qsec, and vs from mtcars_tbl:

```
SELECT(mtcars_tbl, mpg, disp, wt, qsec, vs)

# A tibble: 32 x 5
#       mpg   disp    wt   qsec    vs
#       <dbl> <dbl> <dbl> <dbl> <dbl>
#1 21.0 160.0 2.620 16.46     0
#2 21.0 160.0 2.875 17.02     0
#3 22.8 108.0 2.320 18.61     1
#4 21.4 258.0 3.215 19.44     1
#5 18.7 360.0 3.440 17.02     0
#6 18.1 225.0 3.460 20.22     1
# ... WITH 26 more ROWS
```

: notation can be used to select consecutive columns. To select columns from **cars** through disp and vs through carb:

```
SELECT(mtcars_tbl, cars:disp, vs:carb)

# A tibble: 32 x 8
#       cars   mpg   cyl   disp    vs    am   gear   carb
#       <chr> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl>
#1 Mazda RX4  21.0     6 160.0     0     1     4     4
#2 Mazda RX4 Wag 21.0     6 160.0     0     1     4     4
#3 Datsun 710  22.8     4 108.0     1     1     4     1
#4 Hornet 4 Drive 21.4     6 258.0     1     0     3     1
#5 Hornet Sportabout 18.7     8 360.0     0     0     3     2
#6 Valiant     18.1     6 225.0     1     0     3     1
# ... WITH 26 more ROWS
```

or **SELECT**(mtcars_tbl, -(hp:qsec))

For datasets that contain several columns, it can be tedious to select several columns by name. To make life easier, there are a number of helper functions (such as `starts_with()`, `ends_with()`, `contains()`, `matches()`, `num_range()`, `one_of()`, and `everything()`) that can be used in **SELECT**. To learn more about how to use them, see `?select_helpers` and `?select`.

Note: While referring to columns directly in **SELECT()**, we use bare column names, but quotes should be used while

referring to columns in helper functions.

To rename columns while selecting:

```
SELECT(mtcars_tbl, cylinders = cyl, displacement = disp)

# A tibble: 32 x 2
#   cylinders displacement
#       <dbl>        <dbl>
#1       6        160.0
#2       6        160.0
#3       4        108.0
#4       6        258.0
#5       8        360.0
#6       6        225.0
# ... WITH 26 more ROWS
```

As expected, this drops all other variables.

To rename columns without dropping other variables, use `rename`:

```
rename(mtcars_tbl, cylinders = cyl, displacement = disp)

# A tibble: 32 x 12
#   cars     mpg cylinders displacement      hp    drat      wt    qsec    vs
#   <chr>   <dbl>        <dbl>        <dbl>   <dbl>   <dbl>   <dbl>   <dbl>   <dbl>
#1 Mazda RX4  21.0         6        160.0    110  3.90 2.620 16.46    0
#2 Mazda RX4 Wag 21.0        6        160.0    110  3.90 2.875 17.02    0
#3 Datsun 710  22.8         4        108.0     93  3.85 2.320 18.61    1
#4 Hornet 4 Drive 21.4        6        258.0    110  3.08 3.215 19.44    1
#5 Hornet Sportabout 18.7        8        360.0    175  3.15 3.440 17.02    0
#6 Valiant    18.1         6        225.0    105  2.76 3.460 20.22    1
# ... with 26 more rows, and 3 more variables: am <dbl>, gear <dbl>, carb <dbl>
```

mutate

`mutate` can be used to add new columns to the data. Like all other functions in `dplyr`, `mutate` doesn't add the newly created columns to the original data. Columns are added at the end of the `data.frame`.

```
mutate(mtcars_tbl, weight_ton = wt/2, weight_pounds = weight_ton * 2000)

# A tibble: 32 x 14
#   cars     mpg   cyl   disp      hp    drat      wt    qsec    vs    am    gear    carb weight_ton
#   <chr>   <dbl> <dbl> <dbl>   <dbl>   <dbl>   <dbl>   <dbl>   <dbl>   <dbl>   <dbl>   <dbl>   <dbl>
#1 Mazda RX4  21.0     6 160.0    110  3.90 2.620 16.46    0     1     4     4     1.3100
#2 Mazda RX4 Wag 21.0     6 160.0    110  3.90 2.875 17.02    0     1     4     4     1.4375
#3 Datsun 710  22.8     4 108.0     93  3.85 2.320 18.61    1     1     4     1     1.1600
#4 Hornet 4 Drive 21.4     6 258.0    110  3.08 3.215 19.44    1     0     3     1     1.6075
#5 Hornet Sportabout 18.7     8 360.0    175  3.15 3.440 17.02    0     0     3     2     1.7200
#6 Valiant    18.1     6 225.0    105  2.76 3.460 20.22    1     0     3     1     1.7300
# ... with 26 more rows
```

Note the use of `weight_ton` while creating `weight_pounds`. Unlike base R, `mutate` allows us to refer to columns that we just created to be used for a subsequent operation.

To retain only the newly created columns, use `transmute` instead of `mutate`:

```
transmute(mtcars_tbl, weight_ton = wt/2, weight_pounds = weight_ton * 2000)

# A tibble: 32 x 2
#   weight_ton    weight_pounds
#       <dbl>        <dbl>
#1     1.3100      2620
#2     1.4375      2875
#3     1.1600      2320
#4     1.6075      3215
#5     1.7200      3440
#6     1.7300      3460
# ... with 26 more rows
```

summarise

`summarise` calculates summary statistics of variables by collapsing multiple values to a single value. It can calculate multiple statistics and we can name these summary columns in the same statement.

To calculate the *mean* and *standard deviation* of `mpg` and `disp` of all cars in the dataset:

```
summarise(mtcars_tbl, mean_mpg = mean(mpg), sd_mpg = sd(mpg),
          mean_disp = mean(disp), sd_disp = sd(disp))

# A tibble: 1 x 4
#   mean_mpg  sd_mpg mean_disp  sd_disp
#       <dbl>    <dbl>      <dbl>    <dbl>
#1 20.09062  6.026948  230.7219 123.9387
```

group_by

`group_by` can be used to perform group wise operations on data. When the verbs defined above are applied on this grouped data, they are automatically applied to each group separately.

To find `mean` and `sd` of `mpg` by `cyl`:

```
by_cyl <- group_by(mtcars_tbl, cyl)
summarise(by_cyl, mean_mpg = mean(mpg), sd_mpg = sd(mpg))

# A tibble: 3 x 3
#   cyl  mean_mpg  sd_mpg
#   <dbl>    <dbl>    <dbl>
#1     4 26.66364 4.509828
#2     6 19.74286 1.453567
#3     8 15.10000 2.560048
```

Putting it all together

We select columns from `cars` through `hp` and `gear`, order the rows by `cyl` and from highest to lowest `mpg`, group the data by `gear`, and finally subset only those cars have `mpg > 20` and `hp > 75`

```
selected <- SELECT(mtcars_tbl, cars:hp, gear)
ordered <- arrange(selected, cyl, DESC(mpg))
by_cyl <- group_by(ordered, gear)
FILTER(by_cyl, mpg > 20, hp > 75)
```

```
SOURCE: LOCAL DATA frame [9 x 6]
```

```
Groups: gear [3]
```

```
#          cars   mpg   cyl  disp    hp  gear
#          <chr> <dbl> <dbl> <dbl> <dbl> <dbl>
#1  Lotus Europa  30.4     4  95.1   113     5
#2  Porsche 914-2  26.0     4 120.3    91     5
#3   Datsun 710   22.8     4 108.0   93     4
#4     Merc 230   22.8     4 140.8   95     4
#5  Toyota Corona  21.5     4 120.1   97     3
# ... WITH 4 more ROWS
```

Maybe we are not interested the intermediate results, we can achieve the same result as above by wrapping the function calls:

```
filter(
  group_by(
    arrange(
      select(
        mtcars_tbl, cars:hp
      ), cyl, desc(mpg)
    ), cyl
  ), mpg > 20, hp > 75
)
```

This can be a little difficult to read. So, dplyr operations can be chained using the pipe `%>%` operator. The above code translates to:

```
mtcars_tbl %>%
  select(cars:hp) %>%
  arrange(cyl, desc(mpg)) %>%
  group_by(cyl) %>%
  filter(mpg > 20, hp > 75)
```

summarise multiple columns

dplyr provides `summarise_all()` to apply functions to all (non-grouping) columns.

To find the number of distinct values for each column:

```
mtcars_tbl %>%
  summarise_all(n_distinct)

# A tibble: 1 x 12
#   cars   mpg   cyl  disp    hp  drat    wt  qsec    vs    am  gear  carb
#   <int> <int>
#1    32    25     3    27    22    22    29    30     2     2     3     6
```

To find the number of distinct values for each column by cyl:

```
mtcars_tbl %>%
  group_by(cyl) %>%
  summarise_all(n_distinct)

# A tibble: 3 x 12
#   cyl   cars   mpg   cyl  disp    hp  drat    wt  qsec    vs    am  gear  carb
#   <dbl> <int> <int> <int> <int> <int> <int> <int> <int> <int> <int>
#1     4    11     9    11    10    10    11    11     2     2     3     2
#2     6     7     6     5     4     5     6     7     2     2     3     3
```

```
#3     8    14    12    11     9    11    13    14     1     2     2     4
```

Note that we just had to add the `group_by` statement and the rest of the code is the same. The output now consists of three rows - one for each unique value of cyl.

To summarise specific multiple columns, use `summarise_at`

```
mtcars_tbl %>%
  group_by(cyl) %>%
  summarise_at(c("mpg", "disp", "hp"), mean)

# A tibble: 3 x 4
#   cyl      mpg      disp       hp
#   <dbl>    <dbl>    <dbl>     <dbl>
#1     4 26.66364 105.1364 82.63636
#2     6 19.74286 183.3143 122.28571
#3     8 15.10000 353.1000 209.21429
```

helper functions (`?select_helpers`) can be used in place of column names to select specific columns

To apply multiple functions, either pass the function names as a character vector:

```
mtcars_tbl %>%
  group_by(cyl) %>%
  summarise_at(c("mpg", "disp", "hp"),
               c("mean", "sd"))
```

or wrap them inside `fun`s:

```
mtcars_tbl %>%
  group_by(cyl) %>%
  summarise_at(c("mpg", "disp", "hp"),
               funs(mean, sd))

# A tibble: 3 x 7
#   cyl mpg_mean disp_mean   hp_mean   mpg_sd   disp_sd   hp_sd
#   <dbl>    <dbl>    <dbl>     <dbl>    <dbl>     <dbl>    <dbl>
#1     4 26.66364 105.1364 82.63636 4.509828 26.87159 20.93453
#2     6 19.74286 183.3143 122.28571 1.453567 41.56246 24.26049
#3     8 15.10000 353.1000 209.21429 2.560048 67.77132 50.97689
```

Column names are now be appended with function names to keep them distinct. In order to change this, pass the name to be appended with the function:

```
mtcars_tbl %>%
  group_by(cyl) %>%
  summarise_at(c("mpg", "disp", "hp"),
               c(Mean = "mean", SD = "sd"))

mtcars_tbl %>%
  group_by(cyl) %>%
  summarise_at(c("mpg", "disp", "hp"),
               funs(Mean = mean, SD = sd))

# A tibble: 3 x 7
#   cyl mpg_Mean disp_Mean   hp_Mean   mpg_SD   disp_SD   hp_SD
#   <dbl>    <dbl>    <dbl>     <dbl>    <dbl>     <dbl>    <dbl>
#1     4 26.66364 105.1364 82.63636 4.509828 26.87159 20.93453
```

```
#2      6 19.74286 183.3143 122.28571 1.453567 41.56246 24.26049
#3      8 15.10000 353.1000 209.21429 2.560048 67.77132 50.97689
```

To select columns conditionally, use `summarise_if`:

Take the `mean` of all columns that are `numeric` grouped by `cyl`:

```
mtcars_tbl %>%
  group_by(cyl) %>%
  summarise_if(is.numeric, mean)

# A tibble: 3 x 11
#   cyl     mpg     disp      hp     drat      wt     qsec
# <dbl>   <dbl>   <dbl>   <dbl>   <dbl>   <dbl>   <dbl>
#1     4 26.66364 105.1364 82.63636 4.070909 2.285727 19.13727
#2     6 19.74286 183.3143 122.28571 3.585714 3.117143 17.97714
#3     8 15.10000 353.1000 209.21429 3.229286 3.999214 16.77214
# ... with 4 more variables: vs <dbl>, am <dbl>, gear <dbl>,
#   carb <dbl>
```

However, some variables are discrete, and `mean` of these variables doesn't make sense.

To take the `mean` of only continuous variables by `cyl`:

```
mtcars_tbl %>%
  group_by(cyl) %>%
  summarise_if(function(x) is.numeric(x) & n_distinct(x) > 6, mean)

# A tibble: 3 x 7
#   cyl     mpg     disp      hp     drat      wt     qsec
# <dbl>   <dbl>   <dbl>   <dbl>   <dbl>   <dbl>   <dbl>
#1     4 26.66364 105.1364 82.63636 4.070909 2.285727 19.13727
#2     6 19.74286 183.3143 122.28571 3.585714 3.117143 17.97714
#3     8 15.10000 353.1000 209.21429 3.229286 3.999214 16.77214
```

Section 77.2: Aggregating with %>% (pipe) operator

The pipe (%>%) operator could be used in combination with dplyr functions. In this example we use the `mtcars` dataset (see `help("mtcars")` for more information) to show how to summarize a data frame, and to add variables to the data with the result of the application of a function.

```
library(dplyr)
library(magrittr)
df <- mtcars
df$cars <- rownames(df) #just add the cars names to the df
df <- df[,c(ncol(df),1:(ncol(df)-1))] # and place the names in the first column
```

1. Summarize the data

To compute statistics we use `summarize` and the appropriate functions. In this case `n()` is used for counting the number of cases.

```
df %>%
  summarize(count=n(), mean_mpg = mean(mpg, na.rm = TRUE),
           min_weight = min(wt), max_weight = max(wt))

# count mean_mpg min_weight max_weight
```

```
#1    32 20.09062      1.513      5.424
```

2. Compute statistics by group

It is possible to compute the statistics by groups of the data. In this case by *Number of cylinders* and *Number of forward gears*

```
df %>%
  group_by(cyl, gear) %>%
  summarize(count=n(), mean_mpg = mean(mpg, na.rm = TRUE),
           min_weight = min(wt), max_weight = max(wt))

# Source: local data frame [8 x 6]
# Groups: cyl [?]
#
#   cyl gear count mean_mpg min_weight max_weight
#   <dbl> <dbl> <int>     <dbl>       <dbl>       <dbl>
#1     4     3     1  21.500     2.465     2.465
#2     4     4     8  26.925     1.615     3.190
#3     4     5     2  28.200     1.513     2.140
#4     6     3     2  19.750     3.215     3.460
#5     6     4     4  19.750     2.620     3.440
#6     6     5     1  19.700     2.770     2.770
#7     8     3    12  15.050     3.435     5.424
#8     8     5     2  15.400     3.170     3.570
```

Section 77.3: Subset Observation (Rows)

`dplyr::filter()` - Select a subset of rows in a data frame that meet a logical criteria:

```
dplyr::filter(iris, Sepal.Length>7)
#   Sepal.Length Sepal.Width Petal.Length Petal.Width Species
# 1         7.1      3.0       5.9      2.1  virginica
# 2         7.6      3.0       6.6      2.1  virginica
# 3         7.3      2.9       6.3      1.8  virginica
# 4         7.2      3.6       6.1      2.5  virginica
# 5         7.7      3.8       6.7      2.2  virginica
# 6         7.7      2.6       6.9      2.3  virginica
# 7         7.7      2.8       6.7      2.0  virginica
# 8         7.2      3.2       6.0      1.8  virginica
# 9         7.2      3.0       5.8      1.6  virginica
# 10        7.4      2.8       6.1      1.9  virginica
# 11        7.9      3.8       6.4      2.0  virginica
# 12        7.7      3.0       6.1      2.3  virginica
```

`dplyr::distinct()` - Remove duplicate rows:

```
distinct(iris, Sepal.Length, .keep_all = TRUE)
#   Sepal.Length Sepal.Width Petal.Length Petal.Width Species
# 1         5.1      3.5       1.4      0.2  setosa
# 2         4.9      3.0       1.4      0.2  setosa
# 3         4.7      3.2       1.3      0.2  setosa
# 4         4.6      3.1       1.5      0.2  setosa
# 5         5.0      3.6       1.4      0.2  setosa
# 6         5.4      3.9       1.7      0.4  setosa
# 7         4.4      2.9       1.4      0.2  setosa
# 8         4.8      3.4       1.6      0.2  setosa
# 9         4.3      3.0       1.1      0.1  setosa
# 10        5.8      4.0       1.2      0.2  setosa
# 11        5.7      4.4       1.5      0.4  setosa
# 12        5.2      3.5       1.5      0.2  setosa
```

```

# 13      5.5      4.2      1.4      0.2    setosa
# 14      4.5      2.3      1.3      0.3    setosa
# 15      5.3      3.7      1.5      0.2    setosa
# 16      7.0      3.2      4.7      1.4 versicolor
# 17      6.4      3.2      4.5      1.5 versicolor
# 18      6.9      3.1      4.9      1.5 versicolor
# 19      6.5      2.8      4.6      1.5 versicolor
# 20      6.3      3.3      4.7      1.6 versicolor
# 21      6.6      2.9      4.6      1.3 versicolor
# 22      5.9      3.0      4.2      1.5 versicolor
# 23      6.0      2.2      4.0      1.0 versicolor
# 24      6.1      2.9      4.7      1.4 versicolor
# 25      5.6      2.9      3.6      1.3 versicolor
# 26      6.7      3.1      4.4      1.4 versicolor
# 27      6.2      2.2      4.5      1.5 versicolor
# 28      6.8      2.8      4.8      1.4 versicolor
# 29      7.1      3.0      5.9      2.1  virginica
# 30      7.6      3.0      6.6      2.1  virginica
# 31      7.3      2.9      6.3      1.8  virginica
# 32      7.2      3.6      6.1      2.5  virginica
# 33      7.7      3.8      6.7      2.2  virginica
# 34      7.4      2.8      6.1      1.9  virginica
# 35      7.9      3.8      6.4      2.0  virginica

```

Section 77.4: Examples of NSE and string variables in dplyr

dplyr uses Non-Standard Evaluation(NSE), which is why we normally can use the variable names without quotes. However, sometimes during the data pipeline, we need to get our variable names from other sources such as a Shiny selection box. In case of functions like **SELECT**, we can just use **select_** to use a string variable to select

```

variable1 <- "Sepal.Length"
variable2 <- "Sepal.Width"
iris %>%
select_(variable1, variable2) %>%
head(n=5)
#   Sepal.Length Sepal.Width
# 1          5.1        3.5
# 2          4.9        3.0
# 3          4.7        3.2
# 4          4.6        3.1
# 5          5.0        3.6

```

But if we want to use other features such as summarize or filter we need to use **interp** function from **lazyeval** package

```

variable1 <- "Sepal.Length"
variable2 <- "Sepal.Width"
variable3 <- "Species"
iris %>%
select_(variable1, variable2, variable3) %>%
group_by_(variable3) %>%
summarize_(mean1 = lazyeval::interp(~mean(var), var = as.name(variable1)), mean2 =
lazyeval::interp(~mean(var), var = as.name(variable2)))
#   Species mean1 mean2
#   <fctr>  <dbl> <dbl>
# 1 setosa   5.006 3.428
# 2 versicolor 5.936 2.770
# 3 virginica  6.588 2.974

```

Chapter 78: caret

caret is an R package that aids in data processing needed for machine learning problems. It stands for classification and regression training. When building models for a real dataset, there are some tasks other than the actual learning algorithm that need to be performed, such as cleaning the data, dealing with incomplete observations, validating our model on a test set, and compare different models.

caret helps in these scenarios, independent of the actual learning algorithms used.

Section 78.1: Preprocessing

Pre-processing in caret is done through the `preProcess()` function. Given a matrix or data frame type object `x`, `preProcess()` applies transformations on the training data which can then be applied to testing data.

The heart of the `preProcess()` function is the `method` argument. Method operations are applied in this order:

1. Zero-variance filter
2. Near-zero variance filter
3. Box-Cox/Yeo-Johnson/exponential transformation
4. Centering
5. Scaling
6. Range
7. Imputation
8. PCA
9. ICA
10. Spatial Sign

Below, we take the mtcars data set and perform centering, scaling, and a spatial sign transform.

```
auto_index <- createDataPartition(mtcars$mpg, p = .8,
                                   list = FALSE,
                                   times = 1)

mt_train <- mtcars[auto_index, ]
mt_test <- mtcars[-auto_index, ]

process_mtcars <- preProcess(mt_train, method = c("center", "scale", "spatialSign"))

mtcars_train_transf <- predict(process_mtcars, mt_train)
mtcars_test_trnf <- predict(process_mtcars, mt_test)
```

Chapter 79: Extracting and Listing Files in Compressed Archives

Section 79.1: Extracting files from a .zip archive

Unzipping a zip archive is done with `unzip` function from the `utils` package (which is included in base R).

```
unzip(zipfile = "bar.zip", exdir = "./foo")
```

This will extract all files in `"bar.zip"` to the `"foo"` directory, which will be created if necessary. Tilde expansion is done automatically from your working directory. Alternatively, you can pass the whole path name to the zipfile.

Chapter 80: Probability Distributions with R

Section 80.1: PDF and PMF for different distributions in R

PMF FOR THE BINOMIAL DISTRIBUTION

Suppose that a fair die is rolled 10 times. What is the probability of throwing exactly two sixes?

You can answer the question using the `dbinom` function:

```
> dbinom(2, 10, 1/6)
[1] 0.29071
```

PMF FOR THE POISSON DISTRIBUTION

The number of sandwich ordered in a restaurant on a given day is known to follow a Poisson distribution with a mean of 20. What is the probability that exactly eighteen sandwich will be ordered tomorrow?

You can answer the question with the `dpois` function:

```
> dpois(18, 20)
[1] 0.08439355
```

PDF FOR THE NORMAL DISTRIBUTION

To find the value of the pdf at $x=2.5$ for a normal distribution with a mean of 5 and a standard deviation of 2, use the command:

```
> dnorm(2.5, mean=5, sd=2)
[1] 0.09132454
```

Chapter 81: R in LaTeX with knitr

| Option | Details |
|------------|---|
| echo | (TRUE/FALSE) - whether to include R source code in the output file |
| message | (TRUE/FALSE) - whether to include messages from the R source execution in the output file |
| warning | (TRUE/FALSE) - whether to include warnings from the R source execution in the output file |
| error | (TRUE/FALSE) - whether to include errors from the R source execution in the output file |
| cache | (TRUE/FALSE) - whether to cache the results of the R source execution |
| fig.width | (numeric) - width of the plot generated by the R source execution |
| fig.height | (numeric) - height of the plot generated by the R source execution |

Section 81.1: R in LaTeX with Knitr and Code Externalization

Knitr is an R package that allows us to intermingle R code with LaTeX code. One way to achieve this is external code chunks. External code chunks allow us to develop/test R Scripts in an R development environment and then include the results in a report. It is a powerful organizational technique. This approach is demonstrated below.

```
# r-noweb-file.Rnw
\documentclass{article}

<<echo=FALSE, cache=FALSE>>=
knitr::opts_chunk$set(echo=FALSE,  cache=TRUE)
knitr::read_chunk('r-file.R')
@

\begin{document}
This is an Rnw file (R noweb). It contains a combination of LaTeX and R.

```

One we have called the `read_chunk` command above we can reference sections of code **in** the `r-file.R` script.

```
<<Chunk1>>=
@
\end{document}
```

When using this approach we keep our code in a separate R file as shown below.

```
## r-file.R
## note the specific comment style of a single pound sign followed by four dashes

# ---- Chunk1 ----

print("This is R Code in an external file")

x <- seq(1:10)
y <- rev(seq(1:10))
plot(x,y)
```

Section 81.2: R in LaTeX with Knitr and Inline Code Chunks

Knitr is an R package that allows us to intermingle R code with LaTeX code. One way to achieve this is inline code chunks. This approach is demonstrated below.

```
# r-noweb-file.Rnw
\documentclass{article}
```

```
\begin{document}
This is an Rnw file (R noweb). It contains a combination of LaTeX and R.
```

```
<<my-label>>=
print("This is an R Code Chunk")
x <- seq(1:10)
@
```

Above **is** an internal code chunk.

We can access **data** created **in any** code chunk **inline with** our LaTeX code like this.

The **length** of **array x is** \Sexpr{length(x)}.

```
\end{document}
```

Section 81.3: R in LaTeX with Knitr and Internal Code Chunks

Knitr is an R package that allows us to intermingle R code with LaTeX code. One way to achieve this is internal code chunks. This approach is demonstrated below.

```
# r-noweb-file.Rnw
\documentclass{article}
\begin{document}
This is an Rnw file (R noweb). It contains a combination of LaTeX and R.
```

```
<<code-chunk-label>>=
print("This is an R Code Chunk")
x <- seq(1:10)
y <- seq(1:10)
plot(x,y) # Brownian motion
@
```

```
\end{document}
```

Chapter 82: Web Crawling in R

Section 82.1: Standard scraping approach using the RCurl package

We try to extract imdb top chart movies and ratings

```
R> library(RCurl)
R> library(XML)
R> url <- "http://www.imdb.com/chart/top"
R> top <- getURL(url)
R> parsed_top <- htmlParse(top, encoding = "UTF-8")
R> top_table <- readHTMLTable(parsed_top)[[1]]
R> head(top_table[1:10, 1:3])
```

| Rank | Title | IMDb Rating |
|------|--|-------------|
| 1 | The Shawshank Redemption (1994) | 9.2 |
| 2 | The Godfather (1972) | 9.2 |
| 3 | The Godfather: Part II (1974) | 9.0 |
| 4 | The Dark Knight (2008) | 8.9 |
| 5 | Pulp Fiction (1994) | 8.9 |
| 6 | The Good, the Bad and the Ugly (1966) | 8.9 |
| 7 | Schindler's List (1993) | 8.9 |
| 8 | 12 Angry Men (1957) | 8.9 |
| 9 | The Lord of the Rings: The Return of the King (2003) | 8.9 |
| 10 | Fight Club (1999) | 8.8 |

Chapter 83: Creating reports with RMarkdown

Section 83.1: Including bibliographies

A bibtex catalogue can easily be included with the YAML option `bibliography`:. A certain style for the bibliography can be added with `biblio-style`:. The references are added at the end of the document.

```
---
title: "Including Bibliography"
author: "John Doe"
output: pdf_document
bibliography: references.bib
---

# Abstract

@R_Core_Team_2016

# References
```

Section 83.2: Including LaTeX Preamble Commands

There are two possible ways of including LaTeX preamble commands (e.g. `\usepackage`) in a RMarkdown document.

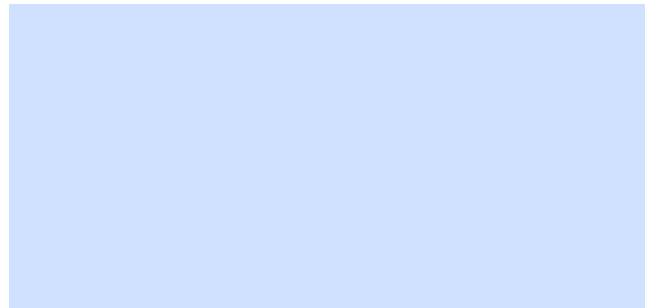
1. Using the YAML option `header-includes`:

```
---
title: "Including LaTeX Preamble Commands in RMarkdown"
header-includes:
- \renewcommand{\familydefault}{cmss}
- \usepackage[cm, slantedGreek]{sfmath}
- \usepackage[T1]{fontenc}
output: pdf_document
---

```{r setup, include=FALSE}
knitr::opts_chunk$set(echo = TRUE, external=T)
```

# Section 1
```

As you can see, this text uses the Computer Modern Font!



2. Including External Commands with `includes`, `in_header`

```
---
title: "Including LaTeX Preamble Commands in RMarkdown"
output:
pdf_document:
includes:
in_header: includes.tex
---

```{r setup, include=FALSE}
knitr::opts_chunk$set(echo = TRUE, external=T)
```

# Section 1

As you can see, this text uses the Computer Modern Font!
```

Here, the content of `includes.tex` are the same three commands we included with `header-includes`.

Writing a whole new template

A possible third option is to write your own LaTeX template and include it with `template`. But this covers a lot more of the structure than only the preamble.

```
---
title: "My Template"
author: "Martin Schmelzer"
output:
pdf_document:
template: myTemplate.tex
---
```

Section 83.3: Printing tables

There are several packages that allow the output of data structures in form of HTML or LaTeX tables. They mostly differ in flexibility.

Here I use the packages:

- knitr
- xtable
- pandoc

For HTML documents

```
---
```

```
title: "Printing Tables"
author: "Martin Schmelzer"
date: "29 Juli 2016"
output: html_document
---
```

```
```{r setup, include=FALSE}
knitr::opts_chunk$set(echo = TRUE)
library(knitr)
library(xtable)
library(pander)
df <- mtcars[1:4,1:4]
```

# Print tables using `kable`
```{r, 'kable'}
kable(df)
```

# Print tables using `xtable`
```{r, 'xtable', results='asis'}
print(xtable(df), type="html")
```

# Print tables using `pander`
```{r, 'pander'}
pander(df)
```
```

For PDF documents

```
---
```

```
title: "Printing Tables"
author: "Martin Schmelzer"
date: "29 Juli 2016"
output: pdf_document
---
```

```
```{r setup, include=FALSE}
knitr::opts_chunk$set(echo = TRUE)
```

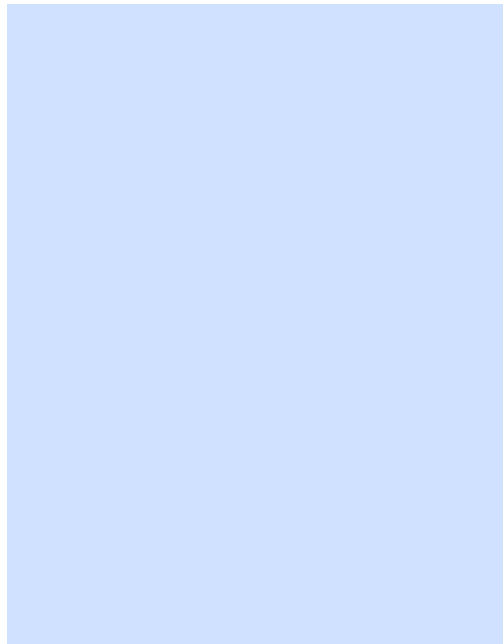
```
library(knitr)
library(xtable)
library(pander)
df <- mtcars[1:4,1:4]
```

# Print tables using `kable`
```{r, 'kable'}
kable(df)
```

# Print tables using `xtable`
```{r, 'xtable', results='asis'}
print(xtable(df, caption="My Table"))
```

# Print tables using `pander`
```{r, 'pander'}
pander(df)
```

```



How can I stop xtable printing the comment ahead of each table?

```
options(xtable.comment = FALSE)
```

Section 83.4: Basic R-markdown document structure

R-markdown code chunks

R-markdown is a markdown file with embedded blocks of R code called *chunks*. There are two types of R code chunks: **inline** and **block**.

Inline chunks are added using the following syntax:

```
`r 2*2`
```

They are evaluated and inserted their output answer in place.

Block chunks have a different syntax:

```
```{r name, echo=TRUE, include=TRUE, ...}
```

2\*2

....

And they come with several possible options. Here are the main ones (but there are many others):

- **echo** (boolean) controls whether the code inside chunk will be included in the document
- **include** (boolean) controls whether the output should be included in the document
- **fig.width** (numeric) sets the width of the output figures
- **fig.height** (numeric) sets the height of the output figures
- **fig.cap** (character) sets the figure captions

They are written in a simple tag=value format like in the example above.

### R-markdown document example

Below is a basic example of R-markdown file illustrating the way R code chunks are embedded inside r-markdown.

```
Title

This is **plain markdown** text.

```{r code, include=FALSE, echo=FALSE}

# Just declare variables

income <- 1000
taxes  <- 125

```

My income is: `r income` dollars and I payed `r taxes` dollars in taxes.

Below is the sum of money I will have left:

```{r gain, include=TRUE, echo=FALSE}

gain <- income-taxes

gain

```

```{r plotOutput, include=TRUE, echo=FALSE, fig.width=6, fig.height=6}

pie(c(income,taxes), label=c("income", "taxes"))

```

```

### Converting R-markdown to other formats

The R knitr package can be used to evaluate R chunks inside R-markdown file and turn it into a regular markdown file.

The following steps are needed in order to turn R-markdown file into pdf/html:

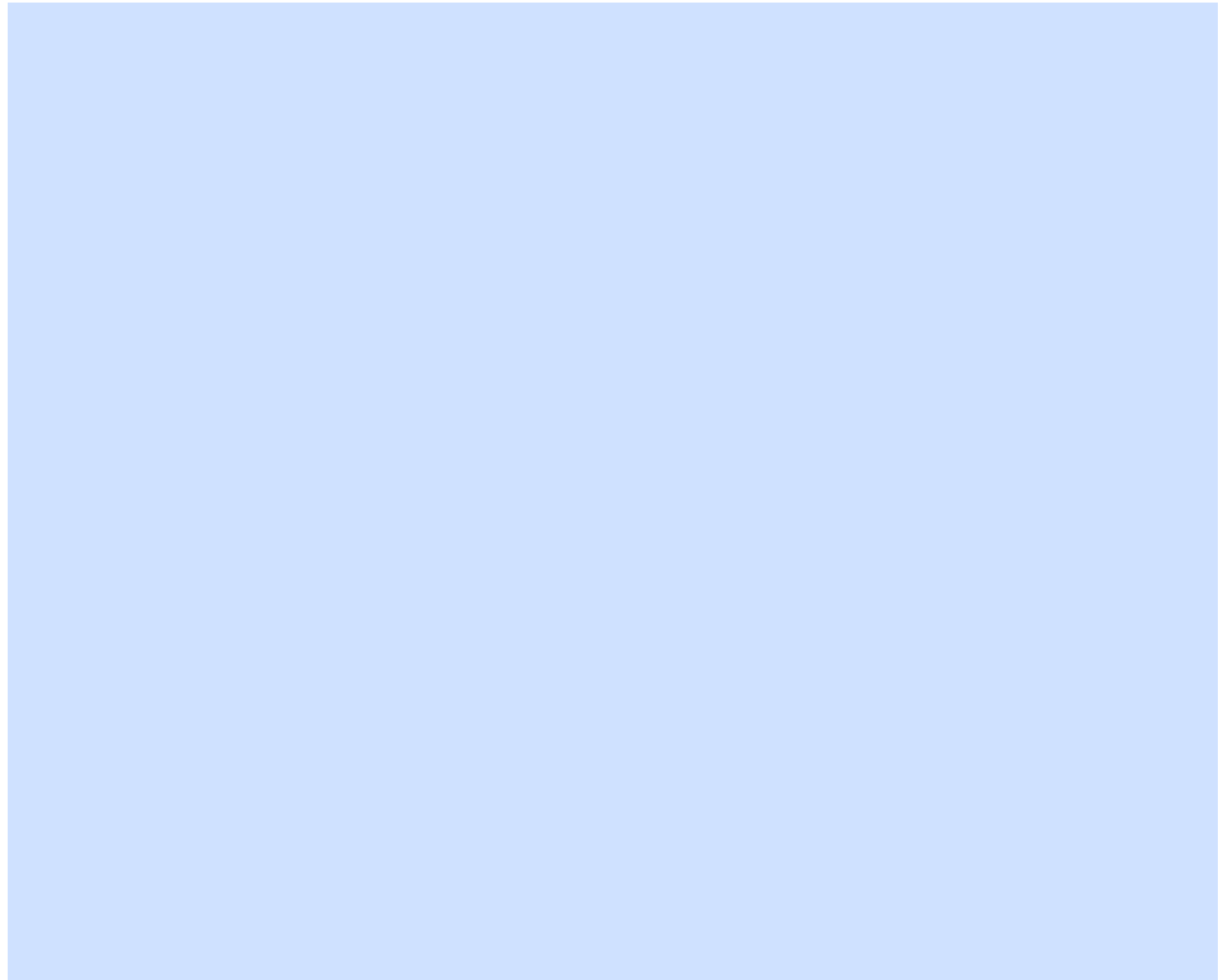
1. Convert R-markdown file to markdown file using knitr.
2. Convert the obtained markdown file to pdf/html using specialized tools like *pandoc*.

In addition to the above knitr package has wrapper functions `knit2html()` and `knit2pdf()` that can be used to produce the final document without the intermediate step of manually converting it to the markdown format:

If the above example file was saved as `income.Rmd` it can be converted to a **pdf** file using the following R commands:

```
library(knitr)
knit2pdf("income.Rmd", "income.pdf")
```

The final document will be similar to the one below.



# Chapter 84: GPU-accelerated computing

## Section 84.1: gpuR gpuMatrix objects

```
library(gpuR) # gpuMatrix objects X <- gpuMatrix(rnorm(100), 10, 10) Y <- gpuMatrix(rnorm(100), 10, 10) # transfer data to GPU when operation called # automatically copied back to CPU Z <- X %*% Y
```

## Section 84.2: gpuR vclMatrix objects

```
library(gpuR) # vclMatrix objects X <- vclMatrix(rnorm(100), 10, 10) Y <- vclMatrix(rnorm(100), 10, 10) # data always on GPU # no data transfer Z <- X %*% Y
```

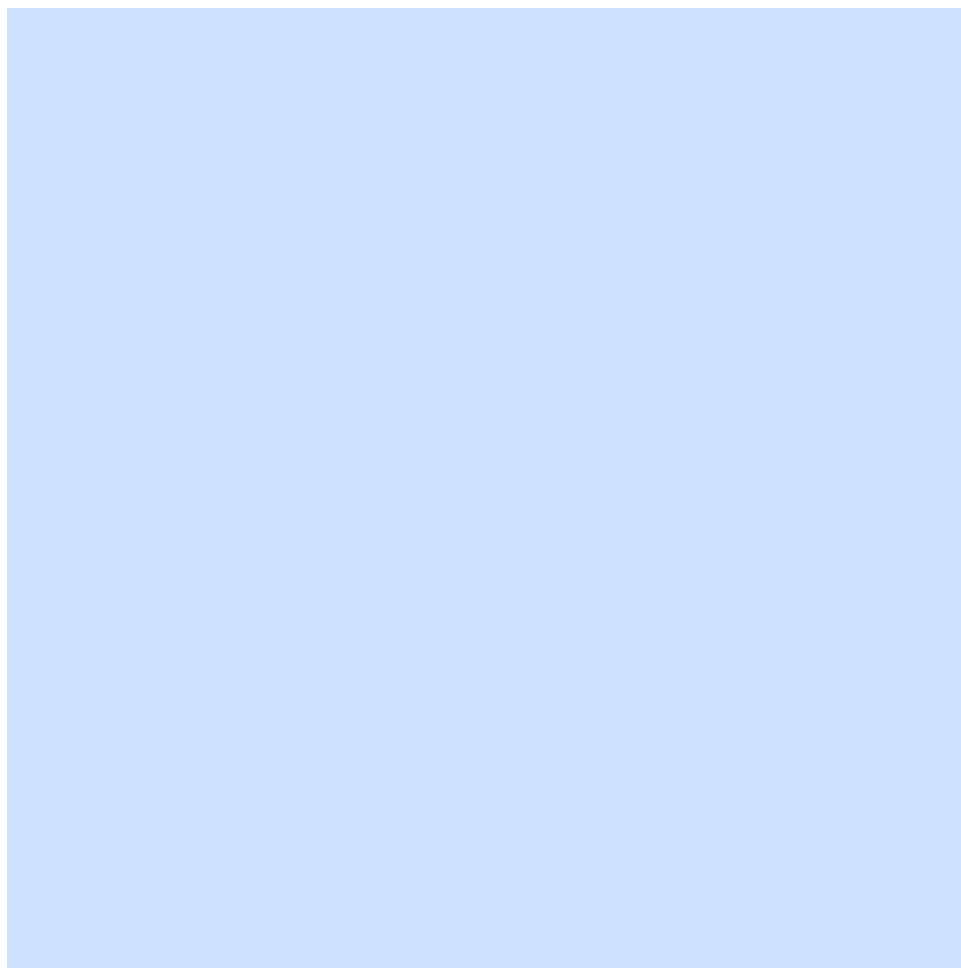
# Chapter 85: heatmap and heatmap.2

## Section 85.1: Examples from the official documentation

stats::heatmap

### Example 1 (Basic usage)

```
require(graphics); require(grDevices)
x <- as.matrix(mtcars)
rc <- rainbow(nrow(x), start = 0, end = .3)
cc <- rainbow(ncol(x), start = 0, end = .3)
hv <- heatmap(x, col = cm.colors(256), scale = "column",
 RowSideColors = rc, ColSideColors = cc, margins = c(5,10),
 xlab = "specification variables", ylab = "Car Models",
 main = "heatmap(<Mtcars data>, ..., scale = \"column\")")
```



```
utils::str(hv) # the two re-ordering index vectors
List of 4
$ rowInd: int [1:32] 31 17 16 15 5 25 29 24 7 6 ...
$ colInd: int [1:11] 2 9 8 11 6 5 10 7 1 4 ...
$ Rowv : NULL
$ Colv : NULL
```

### Example 2 (no column dendrogram (nor reordering) at all)

```
heatmap(x, Colv = NA, col = cm.colors(256), scale = "column",
 RowSideColors = rc, margins = c(5,10),
 xlab = "specification variables", ylab = "Car Models",
 main = "heatmap(<Mtcars data>, ..., scale = \"column\")")
```



### Example 3 ("no nothing")

```
heatmap(x, Rowv = NA, Colv = NA, scale = "column",
 main = "heatmap(*, NA, NA) ~= image(t(x))")
```



#### Example 4 (with reorder())

```
round(Ca <- cor(attitude), 2)
rating complaints privileges learning raises critical advance
rating 1.00 0.83 0.43 0.62 0.59 0.16 0.16
complaints 0.83 1.00 0.56 0.60 0.67 0.19 0.22
privileges 0.43 0.56 1.00 0.49 0.45 0.15 0.34
learning 0.62 0.60 0.49 1.00 0.64 0.12 0.53
raises 0.59 0.67 0.45 0.64 1.00 0.38 0.57
critical 0.16 0.19 0.15 0.12 0.38 1.00 0.28
advance 0.16 0.22 0.34 0.53 0.57 0.28 1.00
symnum(Ca) # simple graphic
rt cm p l rs cr a
rating 1
complaints + 1
privileges . . 1
learning , . . 1
raises . , . , 1
critical . 1
advance . . . 1
attr(,"legend")
[1] 0 ' 0.3 ' 0.6 ' 0.8 '+' 0.9 '*' 0.95 'B' 1
heatmap(Ca, symm = TRUE, margins = c(6,6))
```



### Example 5 (NO reorder())

```
heatmap(Ca, Rowv = FALSE, symm = TRUE, margins = c(6,6))
```



#### **Example 6 (slightly artificial with color bar, without ordering)**

```
cc <- rainbow(nrow(Ca))
heatmap(Ca, Rowv = FALSE, symm = TRUE, RowSideColors = cc, ColSideColors = cc,
 margins = c(6,6))
```



### Example 7 (slightly artificial with color bar, with ordering)

```
heatmap(Ca, symm = TRUE, RowSideColors = cc, ColSideColors = cc,
margins = c(6,6))
```



### Example 8 (For variable clustering, rather use distance based on cor())

```
symnum(cU <- cor(USJudgeRatings))
CO I DM DI CF DE PR F O W PH R
CONT 1
INTG 1
DMNR B 1
DILG + + 1
CFMG + + B 1
DECI + + B B 1
PREP + + B B B 1
FAMI + + B * * B 1
ORAL * * B B * B B 1
WRIT * + B * * B B B 1
PHYS , , + + + + + + + 1
RTEN * * * * * B * B B * 1
attr(,"legend")
[1] 0 ' 0.3 .' 0.6 ',' 0.8 '+' 0.9 '*' 0.95 'B' 1

hU <- heatmap(cU, Rowv = FALSE, symm = TRUE, col = topo.colors(16),
 distfun = function(c) as.dist(1 - c), keep.dendro = TRUE)
```

```

The Correlation matrix with same reordering:
round(100 * cU[hU[[1]], hu[[2]]])
CONT INTG DMNR PHYS DILG CFMG DECI RTEN ORAL WRIT PREP FAMI
CONT 100 -13 -15 5 1 14 9 -3 -1 -4 1 -3
INTG -13 100 96 74 87 81 80 94 91 91 88 87
DMNR -15 96 100 79 84 81 80 94 91 89 86 84
PHYS 5 74 79 100 81 88 87 91 89 86 85 84
DILG 1 87 84 81 100 96 96 93 95 96 98 96
CFMG 14 81 81 88 96 100 98 93 95 94 96 94
DECI 9 80 80 87 96 98 100 92 95 95 96 94
RTEN -3 94 94 91 93 93 92 100 98 97 95 94
ORAL -1 91 91 89 95 95 95 98 100 99 98 98
WRIT -4 91 89 86 96 94 95 97 99 100 99 99
PREP 1 88 86 85 98 96 96 95 98 99 100 99
FAMI -3 87 84 84 96 94 94 94 98 99 99 100

The column dendrogram:
utils::str(hU$Colv)
--[dendrogram w/ 2 branches and 12 members at h = 1.15]
|--leaf "CONT"
`--[dendrogram w/ 2 branches and 11 members at h = 0.258]
|--leaf "INTG"
|--leaf "DMNR"
`--[dendrogram w/ 2 branches and 9 members at h = 0.187]
|--leaf "PHYS"
`--[dendrogram w/ 2 branches and 8 members at h = 0.075]
|--leaf "DILG"
`--[dendrogram w/ 2 branches and 3 members at h = 0.0438]
|--leaf "CFMG"
`--[dendrogram w/ 2 branches and 2 members at h = 0.0189]
|--leaf "DECI"
`--leaf "DECI"

```

```
`--[dendrogram w/ 2 branches and 5 members at h = 0.0584]
|--leaf "RTEN"
`--[dendrogram w/ 2 branches and 4 members at h = 0.0187]
|--[dendrogram w/ 2 branches and 2 members at h = 0.00657]
| |--leaf "ORAL"
| `--leaf "WRIT"
`--[dendrogram w/ 2 branches and 2 members at h = 0.0101]
|--leaf "PREP"
`--leaf "FAMI"
```

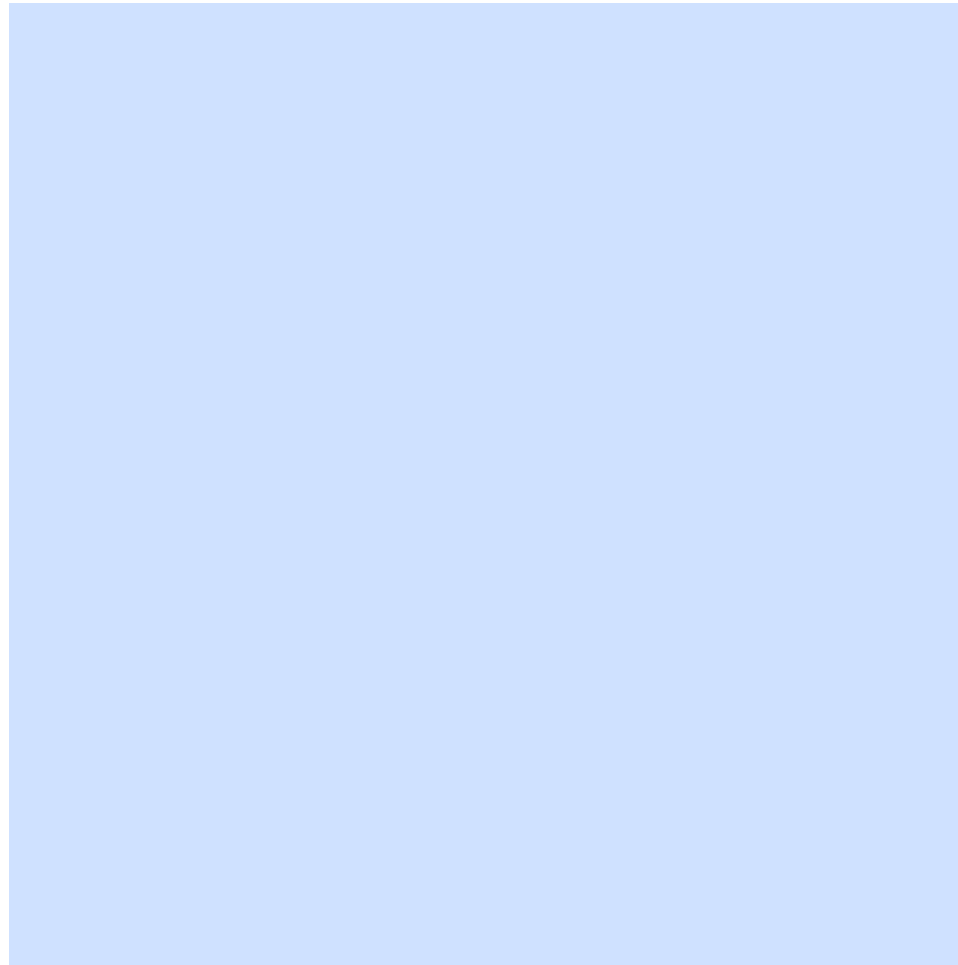
## Section 85.2: Tuning parameters in heatmap.2

Given:

```
x <- as.matrix(mtcars)
```

One can use `heatmap.2` - a more recent optimized version of `heatmap`, by loading the following library:

```
require(gplots)
heatmap.2(x)
```



To add a title, x- or y-label to your heatmap, you need to set the `main`, `xlab` and `ylab`:

```
heatmap.2(x, main = "My main title: Overview of car features", xlab="Car features", ylab = "Car brands")
```

If you wish to define your own color palette for your heatmap, you can set the `col` parameter by using the `colorRampPalette` function:

```
heatmap.2(x, trace="none", key=TRUE, Colv=FALSE, dendrogram = "row", col = colorRampPalette(c("darkblue", "white", "darkred"))(100))
```



As you can notice, the labels on the y axis (the car names) don't fit in the figure. In order to fix this, the user can tune the `margins` parameter:

```
heatmap.2(x, trace="none", key=TRUE, col = colorRampPalette(c("darkblue", "white", "darkred"))(100), margins=c(5,8))
```



Further, we can change the dimensions of each section of our heatmap (the key histogram, the dendograms and the heatmap itself), by tuning `lhei` and `lwid`:



If we only want to show a row(or column) dendrogram, we need to set Colv=FALSE (or Rowv=FALSE) and adjust the dendrogram parameter:

```
heatmap.2(x, trace="none", key=TRUE, Colv=FALSE, dendrogram = "row", col =
colorRampPalette(c("darkblue", "white", "darkred"))(100), margins=c(5,8), lwid = c(5,15), lhei =
c(3,15))
```



For changing the font size of the legend title, labels and axis, the user needs to set `cex.main`, `cex.lab`, `cex.axis` in the `par` list:

```
par(cex.main=1, cex.lab=0.7, cex.axis=0.7)
heatmap.2(x, trace="none", key=TRUE, Colv=FALSE, dendrogram = "row", col =
colorRampPalette(c("darkblue","white","darkred"))(100), margins=c(5,8), lwid = c(5,15), lhei =
c(5,15))
```



# Chapter 86: Network analysis with the igraph package

## Section 86.1: Simple Directed and Non-directed Network Graphing

The igraph package for R is a wonderful tool that can be used to model networks, both real and virtual, with simplicity. This example is meant to demonstrate how to create two simple network graphs using the igraph package within R v.3.2.3.

### Non-Directed Network

The network is created with this piece of code:

```
g<-graph.formula(Node1-Node2, Node1-Node3, Node4-Node1)
plot(g)
```



### Directed Network

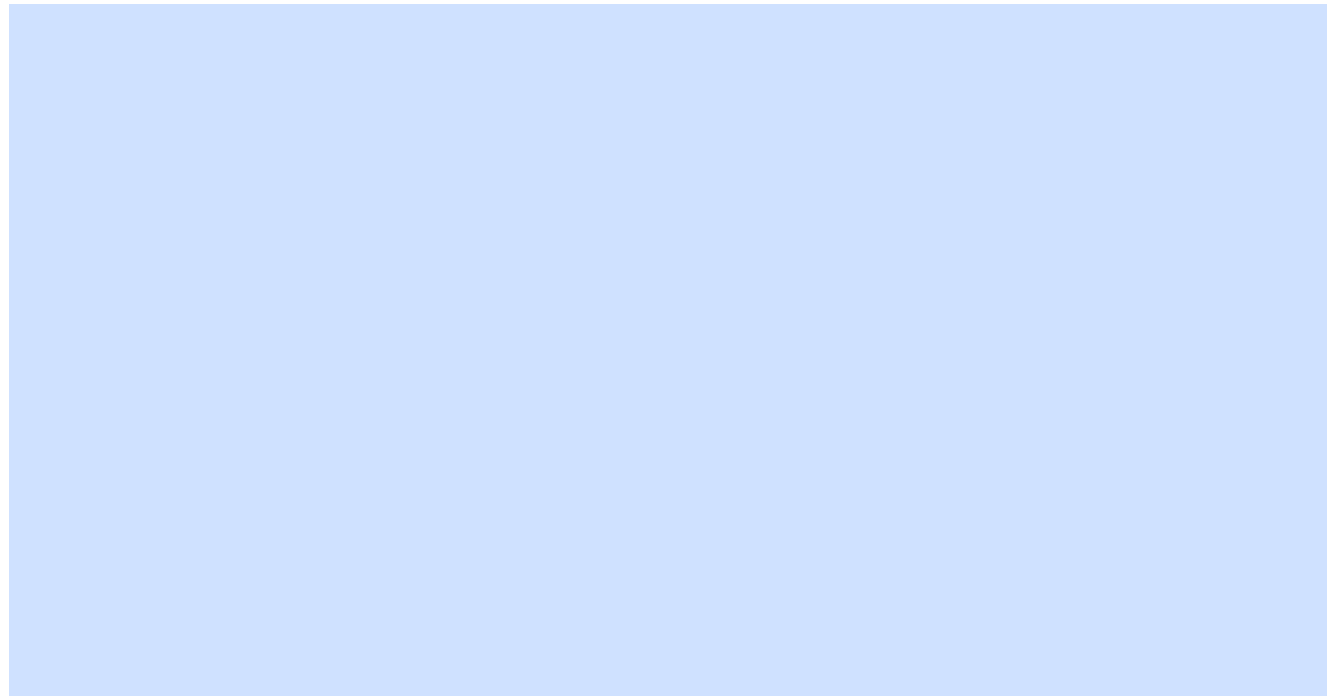
```
dg<-graph.formula(Tom->Mary, Tom->Bill, Tom->Sam, Sue-->Mary, Bill-->Sue)
plot(dg)
```

This code will then generate a network with arrows:



Code example of how to make a double sided arrow:

```
dg<-graph.formula(Tom-->Mary, Tom-->Bill, Tom-->Sam, Sue++>Mary, Bill++>Sue)
plot(dg)
```



# Chapter 87: Functional programming

## Section 87.1: Built-in Higher Order Functions

R has a set of built in higher order functions: [Map](#), [Reduce](#), [Filter](#), [Find](#), [Position](#), [Negate](#).

[Map](#) applies a given function to a list of values:

```
words <- list("this", "is", "an", "example")
Map(toupper, words)
```

[Reduce](#) successively applies a binary function to a list of values in a recursive fashion.

```
Reduce(`*`, 1:10)
```

[Filter](#) given a predicate function and a list of values returns a filtered list containing only values for whom predicate function is TRUE.

```
Filter(is.character, list(1, "a", 2, "b", 3, "c"))
```

[Find](#) given a predicate function and a list of values returns the first value for which the predicate function is TRUE.

```
Find(is.character, list(1, "a", 2, "b", 3, "c"))
```

[Position](#) given a predicate function and a list of values returns the position of the first value in the list for which the predicate function is TRUE.

```
Position(is.character, list(1, "a", 2, "b", 3, "c"))
```

[Negate](#) inverts a predicate function making it return FALSE for values where it returned TRUE and vice versa.

```
is.noncharacter <- Negate(is.character)
is.noncharacter("a")
is.noncharacter(mean)
```

# Chapter 88: Get user input

## Section 88.1: User input in R

Sometimes it can be interesting to have a cross-talk between the user and the program, one example being the [swirl](#) package that had been designed to teach R in R.

One can ask for user input using the `readline` command:

```
name <- readline(prompt = "What is your name?")
```

The user can then give any answer, such as a number, a character, vectors, and scanning the result is here to make sure that the user has given a proper answer. For example:

```
result <- readline(prompt = "What is the result of 1+1?")
while(result!=2){
 readline(prompt = "Wrong answer. What is the result of 1+1?")
}
```

However, it is to be noted that this code be stuck in a never-ending loop, as user input is saved as a character.

We have to coerce it to a number, using `as.numeric`:

```
result <- as.numeric(readline(prompt = "What is the result of 1+1?"))
while(result!=2){
 readline(prompt = "Wrong answer. What is the result of 1+1?")
}
```

# Chapter 89: Spark API (SparkR)

## Section 89.1: Setup Spark context

### Setup Spark context in R

To start working with Sparks distributed dataframes, you must connect your R program with an existing Spark Cluster.

```
library(SparkR)
sc <- sparkR.init() # connection to Spark context
sqlContext <- sparkRSQl.init(sc) # connection to SQL context
```

[Here are infos](#) how to connect your IDE to a Spark cluster.

### Get Spark Cluster

There is an Apache Spark introduction topic with install instructions. Basically, you can employ a Spark Cluster locally via java ([see instructions](#)) or use (non-free) cloud applications (e.g. [Microsoft Azure](#) [topic site], [IBM](#)).

## Section 89.2: Cache data

What:

Caching can optimize computation in Spark. Caching stores data in memory and is a special case of persistence. [Here is explained](#) what happens when you cache an RDD in Spark.

Why:

Basically, caching saves an interim partial result - usually after transformations - of your original data. So, when you use the cached RDD, the already transformed data from memory is accessed without recomputing the earlier transformations.

How:

Here is an example how to quickly access large data (*here 3 GB big csv*) from in-memory storage when accessing it more then once:

```
library(SparkR)
next line is needed for direct csv import:
Sys.setenv('SPARKR_SUBMIT_ARGS'='"--packages" "com.databricks:spark-csv_2.10:1.4.0" "sparkr-
shell"')
sc <- sparkR.init()
sqlContext <- sparkRSQl.init(sc)

loading 3 GB big csv file:
train <- read.df(sqlContext, "/train.csv", source = "com.databricks.spark.csv", inferSchema =
"true")
cache(train)
system.time(head(train))
output: time elapsed: 125 s. This action invokes the caching at this point.
system.time(head(train))
output: time elapsed: 0.2 s (!!)
```

## Section 89.3: Create RDDs (Resilient Distributed Datasets)

**From dataframe:**

```
mtrdd <- createDataFrame(sqlContext, mtcars)
```

**From csv:**

For csv's, you need to add the [csv package](#) to the environment before initiating the Spark context:

```
Sys.setenv('SPARKR_SUBMIT_ARGS'='"--packages" "com.databricks:spark-csv_2.10:1.4.0" "sparkr-shell"') # context for csv import read csv ->
sc <- sparkR.init()
sqlContext <- sparkRSQl.init(sc)
```

Then, you can load the csv either by inferring the data schema of the data in the columns:

```
train <- read.df(sqlContext, "/train.csv", header= true, source = "com.databricks.spark.csv",
inferSchema = true)
```

Or by specifying the data schema beforehand:

```
customSchema <- structType(
 structField("margin", "integer"),
 structField("gross", "integer"),
 structField("name", "string"))

train <- read.df(sqlContext, "/train.csv", header= true, source = "com.databricks.spark.csv",
schema = customSchema)
```

# Chapter 90: Meta: Documentation Guidelines

## Section 90.1: Style

### Prompts

If you want your code to be copy-pastable, remove prompts such as R>, >, or + at the beginning of each new line. Some Docs authors prefer to not make copy-pasting easy, and that is okay.

### Console output

Console output should be clearly distinguished from code. Common approaches include:

- Include prompts on input (as seen when using the console).
- Comment out all output, with # or ## starting each line.
- Print as-is, trusting the leading [1] to make the output stand out from the input.
- Add a blank line between code and console output.

### Assignment

= and <- are fine for assigning R objects. Use white space appropriately to avoid writing code that is difficult to parse, such as x<-1 (ambiguous between x <- 1 and x < -1)

### Code comments

Be sure to explain the purpose and function of the code itself. There isn't any hard-and-fast rule on whether this explanation should be in prose or in code comments. Prose may be more readable and allows for longer explanations, but code comments make for easier copy-pasting. Keep both options in mind.

### Sections

Many examples are short enough to not need sections, but if you use them, start with [H1](#).

## Section 90.2: Making good examples

Most of the guidance for [creating good examples](#) for Q&A carries over into the documentation.

- Make it minimal and get to the point. Complications and digressions are counterproductive.
- Include both working code and prose explaining it. Neither one is sufficient on its own.
- Don't rely on external sources for data. Generate data or use the datasets library if possible:

```
library(help = "datasets")
```

There are some additional considerations in the context of Docs:

- Refer to built-in docs like `?data.frame` whenever relevant. The SO Docs are not an attempt to replace the built-in docs. It is important to make sure new R users know that the built-in docs exist as well as how to find them.
- Move content that applies to multiple examples to the Remarks section.

# Chapter 91: Input and output

## Section 91.1: Reading and writing data frames

Data frames are R's tabular data structure. They can be written to or read from in a variety of ways.

This example illustrates a couple common situations. See the links at the end for other resources.

### Writing

*Before making the example data below, make sure you're in a folder you want to write to. Run `getwd()` to verify the folder you're in and read `?setwd` if you need to change folders.*

```
set.seed(1)
for (i in 1:3)
 write.table(
 data.frame(id = 1:2, v = sample(letters, 2)),
 file = sprintf("file201%s.csv", i)
)
```

Now, we have three similarly-formatted CSV files on disk.

### Reading

We have three similarly-formatted files (from the last section) to read in. Since these files are related, we should store them together after reading in, in a `list`:

```
file_names = c("file2011.csv", "file2012.csv", "file2013.csv")
file_contents = lapply(setNames(file_names, file_names), read.table)

$file2011.csv
id v
1 1 g
2 2 j
#
$file2012.csv
id v
1 1 o
2 2 w
#
$file2013.csv
id v
1 1 f
2 2 w
```

To work with this list of files, first examine the structure with `str(file_contents)`, then read about stacking the list with `?rbind` or iterating over the list with `?lapply`.

### Further resources

Check out `?read.table` and `?write.table` to extend this example. Also:

- R binary formats (for tables and other objects)
- Plain-text table formats
  - comma-delimited CSVs
  - tab-delimited TSVs

- Fixed-width formats
- Language-agnostic binary table formats
  - Feather
- Foreign table and spreadsheet formats
  - SAS
  - SPSS
  - Stata
  - Excel
- Relational database table formats
  - MySQL
  - SQLite
  - PostgreSQL

# Chapter 92: I/O for foreign tables (Excel, SAS, SPSS, Stata)

## Section 92.1: Importing data with rio

A very simple way to import data from many common file formats is with [rio](#). This package provides a function `import()` that wraps many commonly used data import functions, thereby providing a standard interface. It works simply by passing a file name or URL to `import()`:

```
import("example.csv") # comma-separated values
import("example.tsv") # tab-separated values
import("example.dta") # Stata
import("example.sav") # SPSS
import("example.sas7bdat") # SAS
import("example.xlsx") # Excel
```

`import()` can also read from compressed directories, URLs (HTTP or HTTPS), and the clipboard. A comprehensive list of all supported file formats is available on the [rio package github repository](#).

It is even possible to specify some further parameters related to the specific file format you are trying to read, passing them directly within the `import()` function:

```
import("example.csv", format = ",") #for csv file where comma is used as separator
import("example.csv", format = ";") #for csv file where semicolon is used as separator
```

## Section 92.2: Read and write Stata, SPSS and SAS files

The packages `foreign` and `haven` can be used to import and export files from a variety of other statistical packages like Stata, SPSS and SAS and related software. There is a `read` function for each of the supported data types to import the files.

```
loading the packages
library(foreign)
library(haven)
library(readstata13)
library(Hmisc)
```

Some examples for the most common data types:

```
reading Stata files with `foreign`
read.dta("path\to\your\data")
reading Stata files with `haven`
read_dta("path\to\your\data")
```

The `foreign` package can read in stata (.dta) files for versions of Stata 7-12. According to the development page, the `read.dta` is more or less frozen and will not be updated for reading in versions 13+. For more recent versions of Stata, you can use either the `readstata13` package or `haven`. For `readstata13`, the files are

```
reading recent Stata (13+) files with `readstata13`
read.dta13("path\to\your\data")
```

For reading in SPSS and SAS files

```

reading SPSS files with `foreign`
read.spss("path\to\your\data.sav", to.data.frame = TRUE)
reading SPSS files with `haven`
read_spss("path\to\your\data.sav")
read_sav("path\to\your\data.sav")
read_por("path\to\your\data.por")

reading SAS files with `foreign`
read.ssd("path\to\your\data")
reading SAS files with `haven`
read_sas("path\to\your\data")
reading native SAS files with `Hmisc`
sas.get("path\to\your\data") #requires access to saslib
Reading SA XPORT format (*.XPT) files
sasxport.get("path\to\your\data.xpt") # does not require access to SAS executable

```

The SASci package provides functions that will accept SAS SET import code and construct a text file that can be processed with `read.fwf`. It has proved very robust for import of large public-released datasets. Support is at <https://github.com/ajdamico/SASci>

To export data frames to other statistical packages you can use the write functions `write.foreign()`. This will write 2 files, one containing the data and one containing instructions the other package needs to read the data.

```

writing to Stata, SPSS or SAS files with `foreign`
write.foreign(dataframe, datafile, codefile,
 package = c("SPSS", "Stata", "SAS"), ...)
write.foreign(dataframe, "path\to\data\file", "path\to\instruction\file", package = "Stata")

writing to Stata files with `foreign`
write.dta(dataframe, "file", version = 7L,
 convert.dates = TRUE, tz = "GMT",
 convert.factors = c("labels", "string", "numeric", "codes"))

writing to Stata files with `haven`
write_dta(dataframe, "path\to\your\data")

writing to Stata files with `readstata13`
save.dta13(dataframe, file, data.label = NULL, time.stamp = TRUE,
 convert.factors = TRUE, convert.dates = TRUE, tz = "GMT",
 add.rownames = FALSE, compress = FALSE, version = 117,
 convert.underscore = FALSE)

writing to SPSS files with `haven`
write_sav(dataframe, "path\to\your\data")

```

File stored by the SPSS can also be read with `read.spss` in this way:

```

foreign::read.spss('data.sav', to.data.frame=TRUE, use.value.labels=FALSE,
 use.missing=TRUE, reencode='UTF-8')
to.data.frame if TRUE: return a data frame
use.value.labels if TRUE: convert variables with value labels into R factors with those levels
use.missing if TRUE: information on user-defined missing values will be used to set the
corresponding values to NA.
reencode character strings will be re-encoded to the current locale. The default, NA, means to do
so in a UTF-8 locale, only.

```

## Section 92.3: Importing Excel files

There are several R packages to read excel files, each of which using different languages or resources, as

summarized in the following table:

## R package Uses

|           |      |
|-----------|------|
| xlsx      | Java |
| XLconnect | Java |
| openxlsx  | C++  |
| readxl    | C++  |
| RODBC     | ODBC |
| gdata     | Perl |

For the packages that use Java or ODBC it is important to know details about your system because you may have compatibility issues depending on your R version and OS. For instance, if you are using R 64 bits then you also must have Java 64 bits to use xlsx or XLconnect.

Some examples of reading excel files with each package are provided below. Note that many of the packages have the same or very similar function names. Therefore, it is useful to state the package explicitly, like `package::function`. The package openxlsx requires prior installation of RTools.

### Reading excel files with the xlsx package

```
library(xlsx)
```

The index or name of the sheet is required to import.

```
xlsx::read.xlsx("Book1.xlsx", sheetIndex=1)
xlsx::read.xlsx("Book1.xlsx", sheetName="Sheet1")
```

### Reading Excel files with the XLconnect package

```
library(XLConnect)
wb <- XLConnect::loadWorkbook("Book1.xlsx")

Either, if Book1.xlsx has a sheet called "Sheet1":
sheet1 <- XLConnect::readWorksheet(wb, "Sheet1")
Or, more generally, just get the first sheet in Book1.xlsx:
sheet1 <- XLConnect::readWorksheet(wb, getSheets(wb)[1])
```

XLConnect automatically imports the pre-defined Excel cell-styles embedded in Book1.xlsx. This is useful when you wish to format your workbook object and export a perfectly formatted Excel document. Firstly, you will need to create the desired cell formats in Book1.xlsx and save them, for example, as `myHeader`, `myBody` and `myPcts`. Then, after loading the workbook in R (see above):

```
Headerstyle <- XLConnect::getCellStyle(wb, "myHeader")
Bodystyle <- XLConnect::getCellStyle(wb, "myBody")
Pctsstyle <- XLConnect::getCellStyle(wb, "myPcts")
```

The cell styles are now saved in your R environment. In order to assign the cell styles to certain ranges of your data, you need to define the range and then assign the style:

```
Headerrange <- expand.grid(row = 1, col = 1:8)
Bodyrange <- expand.grid(row = 2:6, col = c(1:5, 8))
Pctrange <- expand.grid(row = 2:6, col = c(6, 7))

XLConnect::setCellStyle(wb, sheet = "sheet1", row = Headerrange$row,
 col = Headerrange$col, cellstyle = Headerstyle)
XLConnect::setCellStyle(wb, sheet = "sheet1", row = Bodyrange$row,
```

```
 col = Bodyrange$col, cellstyle = Bodystyle)
XLConnect::setCellStyle(wb, sheet = "sheet1", row = Pctrange$row,
 col = Pctrange$col, cellstyle = Pctsstyle)
```

Note that XLConnect is easy, but can become extremely slow in formatting. A much faster, but more cumbersome formatting option is offered by openxlsx.

## Reading excel files with the openxlsx package

Excel files can be imported with package openxlsx

```
library(openxlsx)

openxlsx::read.xlsx("spreadsheet1.xlsx", colNames=TRUE, rowNames=TRUE)

#colNames: If TRUE, the first row of data will be used as column names.
#rowNames: If TRUE, first column of data will be used as row names.
```

The sheet, which should be read into R can be selected either by providing its position in the sheet argument:

```
openxlsx::read.xlsx("spreadsheet1.xlsx", sheet = 1)
```

or by declaring its name:

```
openxlsx::read.xlsx("spreadsheet1.xlsx", sheet = "Sheet1")
```

Additionally, openxlsx can detect date columns in a read sheet. In order to allow automatic detection of dates, an argument detectDates should be set to TRUE:

```
openxlsx::read.xlsx("spreadsheet1.xlsx", sheet = "Sheet1", detectDates= TRUE)
```

## Reading excel files with the readxl package

Excel files can be imported as a data frame into R using the readxl package.

```
library(readxl)
```

It can read both .xls and .xlsx files.

```
readxl::read_excel("spreadsheet1.xls")
readxl::read_excel("spreadsheet2.xlsx")
```

The sheet to be imported can be specified by number or name.

```
readxl::read_excel("spreadsheet.xls", sheet = 1)
readxl::read_excel("spreadsheet.xls", sheet = "summary")
```

The argument col\_names = TRUE sets the first row as the column names.

```
readxl::read_excel("spreadsheet.xls", sheet = 1, col_names = TRUE)
```

The argument col\_types can be used to specify the column types in the data as a vector.

```
readxl::read_excel("spreadsheet.xls", sheet = 1, col_names = TRUE,
 col_types = c("text", "date", "numeric", "numeric"))
```

## Reading excel files with the RODBC package

Excel files can be read using the ODBC Excel Driver that interfaces with Windows' Access Database Engine (ACE), formerly JET. With the RODBC package, R can connect to this driver and directly query workbooks. Worksheets are assumed to maintain column headers in first row with data in organized columns of similar types. **NOTE:** This approach is limited to only Windows/PC machines as JET/ACE are installed .dll files and not available on other operating systems.

```
library(RODBC)

xlconn <- odbcDriverConnect('Driver={Microsoft Excel Driver (*.xls, *.xlsx, *.xlsm, *.xlsb)};
 DBQ=C:\\Path\\To\\Workbook.xlsx')

df <- sqlQuery(xlconn, "SELECT * FROM [SheetName$]")
close(xlconn)
```

Connecting with an SQL engine in this approach, Excel worksheets can be queried similar to database tables including JOIN and UNION operations. Syntax follows the JET/ACE SQL dialect. **NOTE:** Only data access DML statements, specifically **SELECT** can be run on workbooks, considered not updateable queries.

```
joindf <- sqlQuery(xlconn, "SELECT t1.*, t2.* FROM [Sheet1$] t1
 INNER JOIN [Sheet2$] t2
 ON t1.[ID] = t2.[ID]")

uniondf <- sqlQuery(xlconn, "SELECT * FROM [Sheet1$]
 UNION
 SELECT * FROM [Sheet2$]")
```

Even other workbooks can be queried from the same ODBC channel pointing to a current workbook:

```
otherwkbkdf <- sqlQuery(xlconn, "SELECT * FROM
 [Excel 12.0 Xml;HDR=Yes;
 Database=C:\\Path\\To\\Other\\Workbook.xlsx].[Sheet1$];")
```

### Reading excel files with the gdata package

[example here](#)

## Section 92.4: Import or Export of Feather file

[Feather](#) is an implementation of [Apache Arrow](#) designed to store data frames in a language agnostic manner while maintaining metadata (e.g. date classes), increasing interoperability between Python and R. Reading a feather file will produce a tibble, not a standard data.frame.

```
library(feather)

path <- "filename.feather"
df <- mtcars

write_feather(df, path)

df2 <- read_feather(path)

head(df2)
A tibble: 6 x 11
mpg cyl disp hp drat wt qsec vs am gear carb
<dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl>
1 21.0 6 160 110 3.90 2.620 16.46 0 1 4 4
2 21.0 6 160 110 3.90 2.875 17.02 0 1 4 4
3 22.8 4 108 93 3.85 2.320 18.61 1 1 4 1
```

```
4 21.4 6 258 110 3.08 3.215 19.44 1 0 3 1
5 18.7 8 360 175 3.15 3.440 17.02 0 0 3 2
6 18.1 6 225 105 2.76 3.460 20.22 1 0 3 1

head(df)
mpg cyl disp hp drat wt qsec vs am gear carb
Mazda RX4 21.0 6 160 110 3.90 2.620 16.46 0 1 4 4
Mazda RX4 Wag 21.0 6 160 110 3.90 2.875 17.02 0 1 4 4
Datsun 710 22.8 4 108 93 3.85 2.320 18.61 1 1 4 1
Hornet 4 Drive 21.4 6 258 110 3.08 3.215 19.44 1 0 3 1
Hornet Sportabout 18.7 8 360 175 3.15 3.440 17.02 0 0 3 2
Valiant 18.1 6 225 105 2.76 3.460 20.22 1 0 3 1
```

The current documentation contains this warning:

Note to users: Feather should be treated as alpha software. In particular, the file format is likely to evolve over the coming year. Do not use Feather for long-term data storage.

# Chapter 93: I/O for database tables

## Section 93.1: Reading Data from MySQL Databases

### General

Using the package [RMySQL](#) we can easily query MySQL as well as MariaDB databases and store the result in an R dataframe:

```
library(RMySQL)

mydb <- dbConnect(MySQL(), user='user', password='password', dbname='dbname', host='127.0.0.1')

queryString <- "SELECT * FROM table1 t1 JOIN table2 t2 on t1.id=t2.id"
query <- dbSendQuery(mydb, queryString)
data <- fetch(query, n=-1) # n=-1 to return all results
```

### Using limits

It is also possible to define a limit, e.g. getting only the first 100,000 rows. In order to do so, just change the SQL query regarding the desired limit. The mentioned package will consider these options. Example:

```
queryString <- "SELECT * FROM table1 limit 100000"
```

## Section 93.2: Reading Data from MongoDB Databases

In order to load data from a MongoDB database into an R dataframe, use the library [MongoLite](#):

```
Use MongoLite library:
#install.packages("mongolite")
library(jsonlite)
library(mongolite)

Connect to the database and the desired collection as root:
db <- mongo(collection = "Tweets", db = "TweetCollector", url =
"mongodb://USERNAME:PASSWORD@HOSTNAME")

Read the desired documents i.e. Tweets inside one dataframe:
documents <- db$find(limit = 100000, skip = 0, fields = '{ "_id" : false, "Text" : true }')
```

The code connects to the server HOSTNAME as USERNAME with PASSWORD, tries to open the database TweetCollector and read the collection Tweets. The query tries to read the field i.e. column Text.

The results is a dataframe with columns as the yielded data set. In case of this example, the dataframe contains the column Text, e.g. documents\$Text.

# Chapter 94: I/O for geographic data (shapefiles, etc.)

See also Introduction to Geographical Maps and Input and Output

## Section 94.1: Import and Export Shapefiles

With the `rgdal` package it is possible to import and export shapefiles with R. The function `readOGR` can be used to imports shapefiles. If you want to import a file from e.g. ArcGIS the first argument `dsn` is the path to the folder which contains the shapefile. `layer` is the name of the shapefile without the file ending (just `map` and not `map.shp`).

```
library(rgdal)
readOGR(dsn = "path\to\the\folder\containing\the\shapefile", layer = "map")
```

To export a shapefile use the `writeOGR` function. The first argument is the spatial object produced in R. `dsn` and `layer` are the same as above. The obligatory 4. argument is the driver used to generate the shapefile. The function `ogrDrivers()` lists all available drivers. If you want to export a shapefile to ArcGis or QGis you could use `driver = "ESRI Shapefile"`.

```
writeOGR(Rmap, dsn = "path\to\the\folder\containing\the\shapefile", layer = "map",
 driver = "ESRI Shapefile")
```

`tmap` package has a very convenient function `read_shape()`, which is a wrapper for `rgdal::readOGR()`. The `read_shape()` function simplifies the process of importing a shapefile a lot. On the downside, `tmap` is quite heavy.

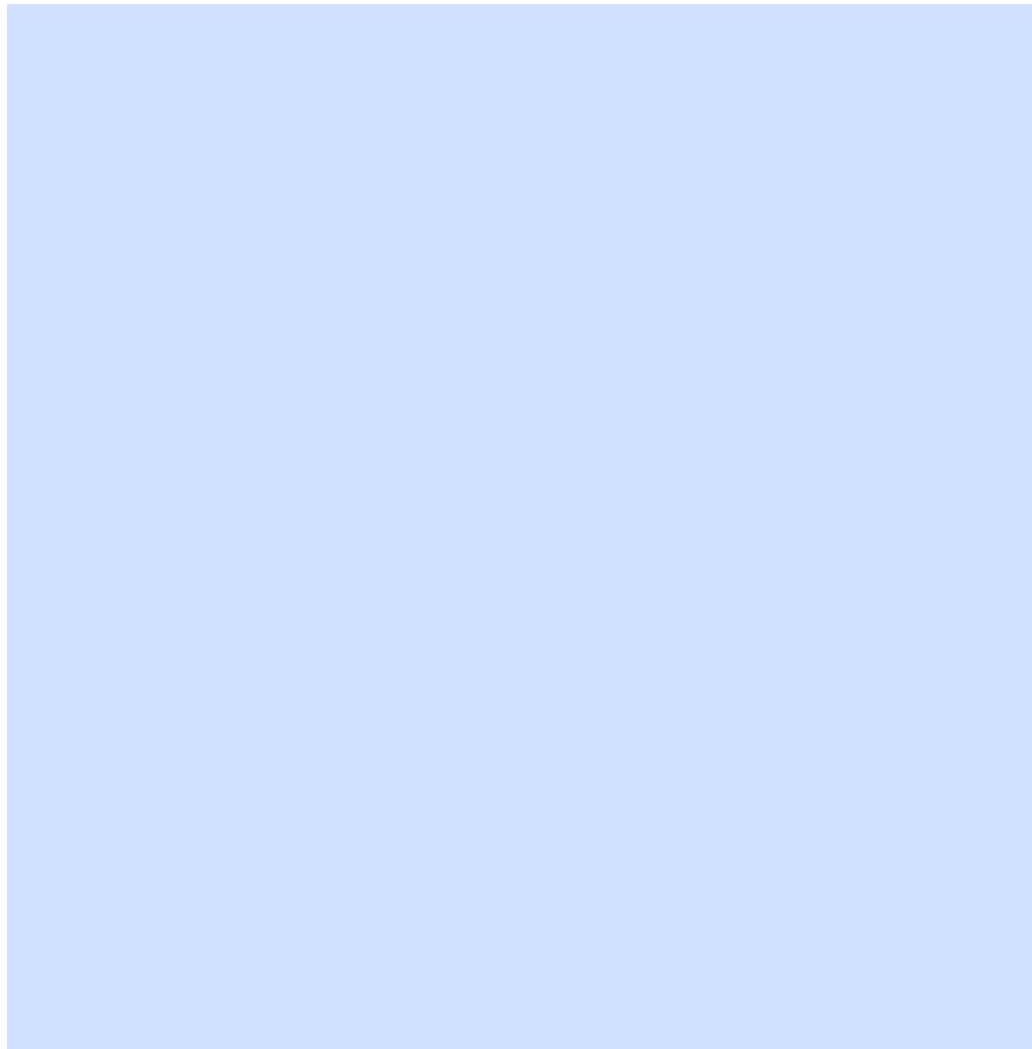
# Chapter 95: I/O for raster images

See also Raster and Image Analysis and Input and Output

## Section 95.1: Load a multilayer raster

The R-Logo is a multilayer raster file (red, green, blue)

```
library(raster)
r <- stack("C:/Program Files/R/R-3.2.3/doc/html/logo.jpg")
plot(r)
```



The individual layers of the `RasterStack` object can be addressed by `[[]]`.

```
plot(r[[1]])
```



# Chapter 96: I/O for R's binary format

## Section 96.1: Rds and RData (Rda) files

.rds and .Rdata (also known as .rda) files can be used to store R objects in a format native to R. There are multiple advantages of saving this way when contrasted with non-native storage approaches, e.g. `write.table`:

- It is faster to restore the data to R
- It keeps R specific information encoded in the data (e.g., attributes, variable types, etc).

`saveRDS/readRDS` only handle a single R object. However, they are more flexible than the multi-object storage approach in that the object name of the restored object need not be the same as the object name when the object was stored.

Using an .rds file, for example, saving the `iris` dataset we would use:

```
saveRDS(object = iris, file = "my_data_frame.rds")
```

To load it data back in:

```
iris2 <- readRDS(file = "my_data_frame.rds")
```

To save a multiple objects we can use `save()` and output as .Rdata.

Example, to save 2 dataframes: iris and cars

```
save(iris, cars, file = "myIrisAndCarsData.Rdata")
```

To load:

```
load("myIrisAndCarsData.Rdata")
```

## Section 96.2: Environments

The functions `save` and `load` allow us to specify the environment where the object will be hosted:

```
save(iris, cars, file = "myIrisAndCarsData.Rdata", envir = foo <- new.env())
load("myIrisAndCarsData.Rdata", envir = foo)
foo$cars

save(iris, cars, file = "myIrisAndCarsData.Rdata", envir = foo <- new.env())
load("myIrisAndCarsData.Rdata", envir = foo)
foo$cars
```

# Chapter 97: Recycling

## Section 97.1: Recycling use in subsetting

Recycling can be used in a clever way to simplify code.

### Subsetting

If we want to keep every third element of a vector we can do the following:

```
my_vec <- c(1,2,3,4,5,6,7,8,9,10)
my_vec[c(TRUE, FALSE)]
[1] 1 3 5 7 9
```

Here the logical expression was expanded to the length of the vector.

We can also perform comparisons using recycling:

```
my_vec <- c("foo", "bar", "soap", "mix")
my_vec == "bar"
[1] FALSE TRUE FALSE FALSE
```

Here "bar" gets recycled.

# Chapter 98: Expression: parse + eval

## Section 98.1: Execute code in string format

In this example, we want to execute code which is stored in a string format.

```
the string
str <- "1+1"

A string is not an expression.
is.expression(str)
[1] FALSE

eval(str)
[1] "1+1"

parse convert string into expressions
parsed.str <- parse(text="1+1")

is.expression(parsed.str)
[1] TRUE

eval(parsed.str)
[1] 2
```

# Chapter 99: Regular Expression Syntax in R

This document introduces the basics of regular expressions as used in R. For more information about R's regular expression syntax, see [?regex](#). For a comprehensive list of regular expression operators, see [this ICU guide on regular expressions](#).

## Section 99.1: Use `grep` to find a string in a character vector

```
General syntax:
grep(<pattern>, <character vector>)

mystring <- c('The number 5',
 'The number 8',
 '1 is the loneliest number',
 'Company, 3 is',
 'Git SSH tag is git@github.com',
 'My personal site is www.personal.org',
 'path/to/my/file')

grep('5', mystring)
[1] 1
grep('@', mystring)
[1] 5
grep('number', mystring)
[1] 1 2 3
```

x|y means look for "x" or "y"

```
grep('5|8', mystring)
[1] 1 2
grep('com|org', mystring)
[1] 5 6
```

. is a special character in Regex. It means "match any character"

```
grep('The number .', mystring)
[1] 1 2
```

Be careful when trying to match dots!

```
tricky <- c('www.personal.org', 'My friend is a cyborg')
grep('.org', tricky)
[1] 1 2
```

To match a literal character, you have to escape the string with a backslash (\). However, R tries to look for escape characters when creating strings, so you actually need to escape the backslash itself (i.e. you need to *double escape* regular expression characters.)

```
grep('\.org', tricky)
Error: '\.' is an unrecognized escape in character string starting "'\.'"
grep('\\.org', tricky)
[1] 1
```

If you want to match one of several characters, you can wrap those characters in brackets ([ ])

```
grep('[13]', mystring)
```

```
[1] 3 4
grep('[@/]', mystring)
[1] 5 7
```

It may be useful to indicate character sequences. E.g. `[0-4]` will match 0, 1, 2, 3, or 4, `[A-Z]` will match any uppercase letter, `[A-z]` will match any uppercase or lowercase letter, and `[A-z0-9]` will match any letter or number (i.e. all alphanumeric characters)

```
grep('[0-4]', mystring)
[1] 3 4
grep('[A-Z]', mystring)
[1] 1 2 4 5 6
```

R also has several shortcut classes that can be used in brackets. For instance, `[:lower:]` is short for `a-z`, `[:upper:]` is short for `A-Z`, `[:alpha:]` is `A-z`, `[:digit:]` is `0-9`, and `[:alnum:]` is `A-z0-9`. Note that these *whole expressions* must be used inside brackets; for instance, to match a single digit, you can use `[[ :digit:]]` (note the double brackets). As another example, `[@[ :digit:]/]` will match the characters `@`, `/` or `0-9`.

```
grep('[[:digit:]]', mystring)
[1] 1 2 3 4
grep('@[:digit:]/', mystring)
[1] 1 2 3 4 5 7
```

Brackets can also be used to negate a match with a carat (^). For instance, `[^5]` will match any character other than "5".

```
grep('The number [^5]', mystring)
[1] 2
```

# Chapter 100: Regular Expressions (regex)

Regular expressions (also called "regex" or "regexp") define patterns that can be matched against a string. Type `?regex` for the official R documentation and see the Regex Docs for more details. The most important 'gotcha' that will not be learned in the SO regex/topics is that most R-regex functions need the use of paired backslashes to escape in a pattern parameter.

## Section 100.1: Differences between Perl and POSIX regex

There are two ever-so-slightly different engines of regular expressions implemented in R. The default is called POSIX-consistent; all regex functions in R are also equipped with an option to turn on the latter type: `perl = TRUE`.

### Look-ahead/look-behind

`perl = TRUE` enables look-ahead and look-behind in regular expressions.

- `"(?<=A)B"` matches an appearance of the letter B *only if* it's preceded by A, i.e. `"ABACADABRA"` would be matched, but `"abacadabra"` and `"aBacadabra"` would not.

## Section 100.2: Validate a date in a "YYYYMMDD" format

It is a common practice to name files using the date as prefix in the following format: YYYYMMDD, for example: `20170101_results.csv`. A date in such string format can be verified using the following regular expression:

```
\d{4}(0[1-9]|1[012])(0[1-9]|12)[0-9]|3[01])
```

The above expression considers dates from year: 0000–9999, months between: 01–12 and days 01–31.

For example:

```
> grep("\\d{4}(0[1-9]|1[012])(0[1-9]|12)[0-9]|3[01])", "20170101")
[1] TRUE
> grep("\\d{4}(0[1-9]|1[012])(0[1-9]|12)[0-9]|3[01])", "20171206")
[1] TRUE
> grep("\\d{4}(0[1-9]|1[012])(0[1-9]|12)[0-9]|3[01])", "29991231")
[1] TRUE
```

**Note:** It validates the date syntax, but we can have a wrong date with a valid syntax, for example: `20170229` (2017 it is not a leap year).

```
> grep("\\d{4}(0[1-9]|1[012])(0[1-9]|12)[0-9]|3[01])", "20170229")
[1] TRUE
```

If you want to validate a date, it can be done via this user defined function:

```
is.Date <- function(x) {return(!is.na(as.Date(as.character(x), format = '%Y%m%d')))}
```

Then

```
> is.Date(c("20170229", "20170101", 20170101))
[1] FALSE TRUE TRUE
```

## Section 100.3: Escaping characters in R regex patterns

Since both R and regex share the escape character , "\\", building correct patterns for `grep`, `sub`, `gsub` or any other function that accepts a pattern argument will often need pairing of backslashes. If you build a three item character vector in which one items has a linefeed, another a tab character and one neither, and the desire is to turn either the linefeed or the tab into 4-spaces then a single backslash is needed for the construction, but paired backslashes for matching:

```
x <- c("a\nb", "c\tb", "e f")
x # how it's stored
[1] "a\nb" "c\tb" "e f"
cat(x) # how it will be seen with cat
#a
#b c d e f

gsub(patt="\n|\t", repl=" ", x)
#[1] "a b" "c d" "e f"
```

Note that the pattern argument (which is optional if it appears first and only needs partial spelling) is the only argument to require this doubling or pairing. The replacement argument does not require the doubling of characters needing to be escaped. If you wanted all the linefeeds and 4-space occurrences replaced with tabs it would be:

```
gsub("\n| ", "\t", x)
#[1] "a\tb" "c\tb" "e\tf"
```

## Section 100.4: Validate US States postal abbreviations

The following regex includes 50 states and also Commonwealth/Territory (see [www.50states.com](http://www.50states.com)):

```
regex <-
"(A[LKSZR])|(C[AOT])|(D[EC])|(F[ML])|(G[AU])|(HI)|(I[DLNA])|(K[SY])|(LA)|(M[EHDAINSOT])|(N[EVHJMYCD])|(MP)|(O[HKR])|(P[WAR])|(RI)|(S[CD])|(T[NX])|(UT)|(V[TIA])|(W[AVIY])"
```

For example:

```
> test <- c("AL", "AZ", "AR", "AJ", "AS", "DC", "FM", "GU", "PW", "FL", "AJ", "AP")
> grep1(us.states.pattern, test)
[1] TRUE TRUE TRUE FALSE TRUE TRUE TRUE TRUE TRUE FALSE FALSE
>
```

### Note:

If you want to verify only the 50 States, then we recommend to use the R-dataset: `state.abb` from `state`, for example:

```
> data(state)
> test %in% state.abb
[1] TRUE TRUE TRUE FALSE FALSE FALSE FALSE FALSE TRUE FALSE FALSE
```

We get TRUE only for 50-States abbreviations: AL, AZ, AR, FL.

## Section 100.5: Validate US phone numbers

The following regular expression:

```
us.phones.regex <- "^\s*(\+\s*|-?\s+)*[0-9]{3}\s*-?\s*[0-9]{3}\s*-?\s*[0-9]{4}$"
```

Validates a phone number in the form of: +1-xxx-xxx-xxxx, including optional leading/trailing blanks at the beginning/end of each group of numbers, but not in the middle, for example: +1-xxx-xxx-xx xx is not valid. The - delimiter can be replaced by blanks: xxx xxx xxx or without delimiter: xxxxxxxxxxxx. The +1 prefix is optional.

Let's check it:

```
us.phones.regex <- "^\s*(\+\s*|-?\s+)*[0-9]{3}\s*-?\s*[0-9]{3}\s*-?\s*[0-9]{4}$"

phones.OK <- c("305-123-4567", "305 123 4567", "+1-786-123-4567",
 "+1 786 123 4567", "7861234567", "786 - 123 4567", "+ 1 786 - 123 4567")

phones.NOK <- c("124-456-78901", "124-456-789", "124-456-78 90",
 "124-45 6-7890", "12 4-456-7890")
```

Valid cases:

```
> grep1(us.phones.regex, phones.OK)
[1] TRUE TRUE TRUE TRUE TRUE TRUE
>
```

Invalid cases:

```
> grep1(us.phones.regex, phones.NOK)
[1] FALSE FALSE FALSE FALSE FALSE
>
```

#### Note:

- \s Matches any space, tab or newline character

# Chapter 101: Combinatorics

## Section 101.1: Enumerating combinations of a specified length

### Without replacement

With `combn`, each vector appears in a column:

```
combn(LETTERS, 3)

Showing only first 10.
[,1] [,2] [,3] [,4] [,5] [,6] [,7] [,8] [,9] [,10]
[1,] "A" "A" "A" "A" "A" "A" "A" "A" "A" "A"
[2,] "B" "B" "B" "B" "B" "B" "B" "B" "B" "B"
[3,] "C" "D" "E" "F" "G" "H" "I" "J" "K" "L"
```

### With replacement

With `expand.grid`, each vector appears in a row:

```
expand.grid(LETTERS, LETTERS, LETTERS)
or
do.call(expand.grid, rep(list(LETTERS), 3))

Showing only first 10.
Var1 Var2 Var3
1 A A A
2 B A A
3 C A A
4 D A A
5 E A A
6 F A A
7 G A A
8 H A A
9 I A A
10 J A A
```

For the special case of pairs, `outer` can be used, putting each vector into a cell:

```
FUN here is used as a function executed on each resulting pair.
in this case it's string concatenation.
outer(LETTERS, LETTERS, FUN=paste0)

Showing only first 10 rows and columns
[,1] [,2] [,3] [,4] [,5] [,6] [,7] [,8] [,9] [,10]
[1,] "AA" "AB" "AC" "AD" "AE" "AF" "AG" "AH" "AI" "AJ"
[2,] "BA" "BB" "BC" "BD" "BE" "BF" "BG" "BH" "BI" "BJ"
[3,] "CA" "CB" "CC" "CD" "CE" "CF" "CG" "CH" "CI" "CJ"
[4,] "DA" "DB" "DC" "DD" "DE" "DF" "DG" "DH" "DI" "DJ"
[5,] "EA" "EB" "EC" "ED" "EE" "EF" "EG" "EH" "EI" "EJ"
[6,] "FA" "FB" "FC" "FD" "FE" "FF" "FG" "FH" "FI" "FJ"
[7,] "GA" "GB" "GC" "GD" "GE" "GF" "GG" "GH" "GI" "GJ"
[8,] "HA" "HB" "HC" "HD" "HE" "HF" "HG" "HH" "HI" "HJ"
[9,] "IA" "IB" "IC" "ID" "IE" "IF" "IG" "IH" "II" "IJ"
[10,] "JA" "JB" "JC" "JD" "JE" "JF" "JG" "JH" "JI" "JJ"
```

## Section 101.2: Counting combinations of a specified length

### Without replacement

```
choose(length(LETTERS), 5)
[1] 65780
```

### With replacement

```
length(letters)^5
[1] 11881376
```

# Chapter 102: Solving ODEs in R

## Parameter Details

|        |                                                                                             |
|--------|---------------------------------------------------------------------------------------------|
| y      | (named) numeric vector: the initial (state) values for the ODE system                       |
| times  | time sequence for which output is wanted; the first value of times must be the initial time |
| func   | name of the function that computes the values of the derivatives in the ODE system          |
| parms  | (named) numeric vector: parameters passed to func                                           |
| method | the integrator to use, by default: lsoda                                                    |

## Section 102.1: The Lorenz model

The Lorenz model describes the dynamics of three state variables, X, Y and Z. The model equations are:



The initial conditions are:



and a, b and c are three parameters with



```
library(deSolve)

Define R-function

Lorenz <- function (t, y, parms) {
with(as.list(c(y, parms)), {
dX <- a * X + Y * Z
dY <- b * (Y - Z)
dZ <- -X * Y + c * Y - Z

return(list(c(dX, dY, dZ)))
})
}

Define parameters and variables
```

```

parms <- c(a = -8/3, b = -10, c = 28)
yini <- c(X = 1, Y = 1, Z = 1)
times <- seq(from = 0, to = 100, by = 0.01)

Solve the ODEs

out <- ode(y = yini, times = times, func = Lorenz, parms = parms)

Plot the results

plot(out, lwd = 2)
plot(out[, "X"], out[, "Y"],
type = "l", xlab = "X",
ylab = "Y", main = "butterfly")

```



## Section 102.2: Lotka-Volterra or: Prey vs. predator

```

library(deSolve)

Define R-function

```

```

LV <- function(t, y, parms) {
with(as.list(c(y, parms)), {
dP <- rG * P * (1 - P/K) - rI * P * C
dC <- rI * P * C * AE - rM * C
return(list(c(dP, dC), sum = C+P))
})
}

Define parameters and variables

parms <- c(rI = 0.2, rG = 1.0, rM = 0.2, AE = 0.5, K = 10)
yini <- c(P = 1, C = 2)
times <- seq(from = 0, to = 200, by = 1)

Solve the ODEs

out <- ode(y = yini, times = times, func = LV, parms = parms)

Plot the results

matplot(out[,1], out[,2:4], type = "l", xlab = "time", ylab = "Conc",
main = "Lotka-Volterra", lwd = 2)
legend("topright", c("prey", "predator", "sum"), col = 1:3, lty = 1:3)

```

## Section 102.3: ODEs in compiled languages - definition in R

```
library(deSolve)

Define parameters and variables

eps <- 0.01;
M <- 10
k <- M * eps^2/2
L <- 1
L0 <- 0.5
r <- 0.1
w <- 10
g <- 1

parameter <- c(eps = eps, M = M, k = k, L = L, L0 = L0, r = r, w = w, g = g)

yini <- c(xl = 0, yl = L0, xr = L, yr = L0,
ul = -L0/L, vl = 0,
ur = -L0/L, vr = 0,
lam1 = 0, lam2 = 0)

times <- seq(from = 0, to = 3, by = 0.01)

Define R-function

caraxis_R <- function(t, y, parms) {
with(as.list(c(y, parms)), {

yb <- r * sin(w * t)
xb <- sqrt(L * L - yb * yb)
Ll <- sqrt(xl^2 + yl^2)
Lr <- sqrt((xr - xb)^2 + (yr - yb)^2)

dxl <- ul; dyl <- vl; dxr <- ur; dyr <- vr

dul <- (L0-Ll) * xl/Ll + 2 * lam2 * (xl-xr) + lam1*xb
dvl <- (L0-Ll) * yl/Ll + 2 * lam2 * (yl-yr) + lam1*yb - k * g

dur <- (L0-Lr) * (xr-xb)/Lr - 2 * lam2 * (xl-xr)
dvr <- (L0-Lr) * (yr-yb)/Lr - 2 * lam2 * (yl-yr) - k * g

c1 <- xb * xl + yb * yl
c2 <- (xl - xr)^2 + (yl - yr)^2 - L * L

return(list(c(dxl, dyl, dxr, dyr, dul, dvl, dur, dvr, c1, c2)))
})}
}
```

## Section 102.4: ODEs in compiled languages - definition in C

```
sink("caraxis_C.c")
cat("
/* suitable names for parameters and state variables */

#include <R.h>
```

```

#include <math.h>
static double parms[8];

#define eps parms[0]
#define m parms[1]
#define k parms[2]
#define L parms[3]
#define L0 parms[4]
#define r parms[5]
#define w parms[6]
#define g parms[7]

/*-----
initialising the parameter common block
-----*/
void init_C(void (* daeparms)(int *, double *)) {
 int N = 8;
 daeparms(&N, parms);
}
/* Compartments */

#define xl y[0]
#define yl y[1]
#define xr y[2]
#define yr y[3]
#define lam1 y[8]
#define lam2 y[9]

/*-----
the residual function
-----*/
void caraxis_C (int *neq, double *t, double *y, double *ydot,
 double *yout, int* ip)
{
 double yb, xb, Lr, Ll;

 yb = r * sin(w * *t) ;
 xb = sqrt(L * L - yb * yb);
 Ll = sqrt(xl * xl + yl * yl) ;
 Lr = sqrt((xr-xb)*(xr-xb) + (yr-yb)*(yr-yb));

 ydot[0] = y[4];
 ydot[1] = y[5];
 ydot[2] = y[6];
 ydot[3] = y[7];

 ydot[4] = (L0-Ll) * xl/Ll + lam1*xb + 2*lam2*(xl-xr) ;
 ydot[5] = (L0-Ll) * yl/Ll + lam1*yb + 2*lam2*(yl-yr) - k*g;
 ydot[6] = (L0-Lr) * (xr-xb)/Lr - 2*lam2*(xl-xr) ;
 ydot[7] = (L0-Lr) * (yr-yb)/Lr - 2*lam2*(yl-yr) - k*g ;

 ydot[8] = xb * xl + yb * yl;
 ydot[9] = (xl-xr) * (xl-xr) + (yl-yr) * (yl-yr) - L*L;

}
", fill = TRUE)
sink()
system("R CMD SHLIB caraxis_C.c")
dyn.load(paste("caraxis_C", .Platform$dynlib.ext, sep = ""))
dllname_C <- dyn.load(paste("caraxis_C", .Platform$dynlib.ext, sep = ""))[[1]]

```

## Section 102.5: ODEs in compiled languages - definition in fortran

```
sink("caraxis_fortran.f")
cat("
c-----
c Initialiser for parameter common block
c-----
subroutine init_fortran(daeparms)

external daeparms
integer, parameter :: N = 8
double precision parms(N)
common /myparms/parms

call daeparms(N, parms)
return
end

c-----
c rate of change
c-----
subroutine caraxis_fortran(neq, t, y, ydot, out, ip)
implicit none
integer neq, IP(*)
double precision t, y(neq), ydot(neq), out(*)
double precision eps, M, k, L, L0, r, w, g
common /myparms/ eps, M, k, L, L0, r, w, g

double precision xl, yl, xr, yr, ul, vl, ur, vr, lam1, lam2
double precision yb, xb, Ll, Lr, dxl, dyl, dxr, dyr
double precision dul, dvl, dur, dvr, c1, c2

c expand state variables
xl = y(1)
yl = y(2)
xr = y(3)
yr = y(4)
ul = y(5)
vl = y(6)
ur = y(7)
vr = y(8)
lam1 = y(9)
lam2 = y(10)

yb = r * sin(w * t)
xb = sqrt(L * L - yb * yb)
Ll = sqrt(xl**2 + yl**2)
Lr = sqrt((xr - xb)**2 + (yr - yb)**2)

dxl = ul
dyl = vl
dxr = ur
dyr = vr

dul = (L0-Ll) * xl/Ll + 2 * lam2 * (xl-xr) + lam1*xb
dvl = (L0-Ll) * yl/Ll + 2 * lam2 * (yl-yr) + lam1*yb - k*g
dur = (L0-Lr) * (xr-xb)/Lr - 2 * lam2 * (xl-xr)
dvr = (L0-Lr) * (yr-yb)/Lr - 2 * lam2 * (yl-yr) - k*g

c1 = xb * xl + yb * yl
c2 = (xl - xr)**2 + (yl - yr)**2 - L * L
```

```

c function values in ydot
ydot(1) = dxl
ydot(2) = dyl
ydot(3) = dxr
ydot(4) = dyr
ydot(5) = dul
ydot(6) = dvl
ydot(7) = dur
ydot(8) = dvr
ydot(9) = c1
ydot(10) = c2
return
end
", fill = TRUE)

sink()
system("R CMD SHLIB caraxis_fortran.f")
dyn.load(paste("caraxis_fortran", .Platform$dynlib.ext, sep = ""))
dllname_fortran <- dyn.load(paste("caraxis_fortran", .Platform$dynlib.ext, sep = ""))[[1]]

```

## Section 102.6: ODEs in compiled languages - a benchmark test

When you compiled and loaded the code in the three examples before (ODEs in compiled languages - definition in R, ODEs in compiled languages - definition in C and ODEs in compiled languages - definition in fortran) you are able to run a benchmark test.

```

library(microbenchmark)

R <- function(){
 out <- ode(y = yini, times = times, func = caraxis_R,
 parms = parameter)
}

C <- function(){
 out <- ode(y = yini, times = times, func = "caraxis_C",
 initfunc = "init_C", parms = parameter,
 dllname = dllname_C)
}

fortran <- function(){
 out <- ode(y = yini, times = times, func = "caraxis_fortran",
 initfunc = "init_fortran", parms = parameter,
 dllname = dllname_fortran)
}

```

Check if results are equal:

```

all.equal(tail(R()), tail(fortran()))
all.equal(R()[,2], fortran()[,2])
all.equal(R()[,2], C()[,2])

```

Make a benchmark (Note: On your machine the times are, of course, different):

```

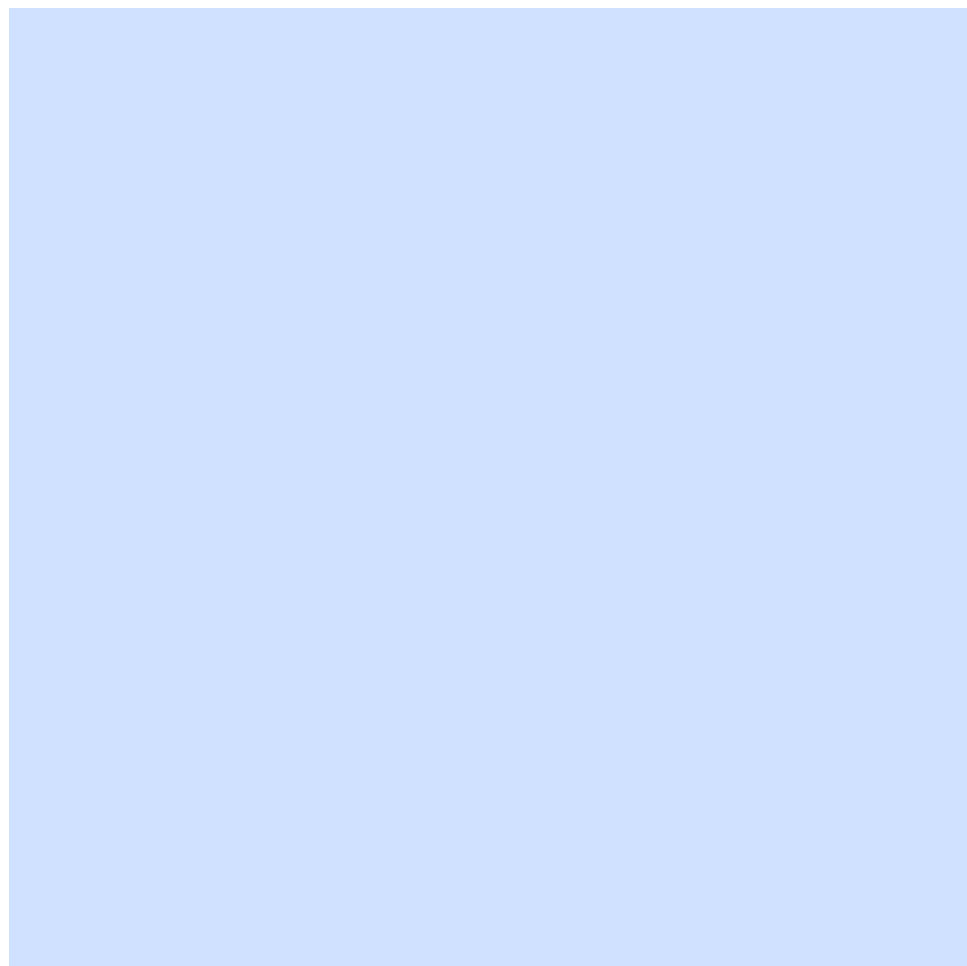
bench <- microbenchmark::microbenchmark(
 R(),
 fortran(),
 C(),
 times = 1000
)

```

```
)
```

```
summary(bench)
```

| expr      | min       | lq        | mean       | median     | uq         | max        | neval | cld |
|-----------|-----------|-----------|------------|------------|------------|------------|-------|-----|
| R()       | 31508.928 | 33651.541 | 36747.8733 | 36062.2475 | 37546.8025 | 132996.564 | 1000  | b   |
| fortran() | 570.674   | 596.700   | 686.1084   | 637.4605   | 730.1775   | 4256.555   | 1000  | a   |
| C()       | 562.163   | 590.377   | 673.6124   | 625.0700   | 723.8460   | 5914.347   | 1000  | a   |



We see clearly, that R is slow in contrast to the definition in C and fortran. For big models it's worth to translate the problem in a compiled language. The package `c0de` is one possibility to translate ODEs from R to C.

# Chapter 103: Feature Selection in R -- Removing Extraneous Features

## Section 103.1: Removing features with zero or near-zero variance

A feature that has near zero variance is a good candidate for removal.

You can manually detect numerical variance below your own threshold:

```
data("GermanCredit")
variances<-apply(GermanCredit, 2, var)
variances[which(variances<=0.0025)]
```

Or, you can use the caret package to find near zero variance. An advantage here is that it defines near zero variance not in the numerical calculation of variance, but rather as a function of rarity:

"nearZeroVar diagnoses predictors that have one unique value (i.e. are zero variance predictors) or predictors that are have both of the following characteristics: they have very few unique values relative to the number of samples and the ratio of the frequency of the most common value to the frequency of the second most common value is large..."

```
library(caret)
names(GermanCredit)[nearZeroVar(GermanCredit)]
```

## Section 103.2: Removing features with high numbers of NA

If a feature is largely lacking data, it is a good candidate for removal:

```
library(VIM)
data(sleep)
colMeans(is.na(sleep))

 BodyWgt BrainWgt NonD Dream Sleep Span Gest
0.00000000 0.00000000 0.22580645 0.19354839 0.06451613 0.06451613 0.06451613
 Pred Exp Danger
0.00000000 0.00000000 0.00000000
```

In this case, we may want to remove NonD and Dream, which each have around 20% missing values (your cutoff may vary)

## Section 103.3: Removing closely correlated features

Closely correlated features may add variance to your model, and removing one of a correlated pair might help reduce that. There are lots of ways to detect correlation. Here's one:

```
library(purrr) # in order to use keep()

select correlatable vars
toCorrelate<-mtcars %>% keep(is.numeric)

calculate correlation matrix
```

```

correlationMatrix <- cor(toCorrelate)

pick only one out of each highly correlated pair's mirror image
correlationMatrix[upper.tri(correlationMatrix)]<-0

and I don't remove the highly-correlated-with-itself group
diag(correlationMatrix)<-0

find features that are highly correlated with another feature at the +- 0.85 level
apply(correlationMatrix,2, function(x) any(abs(x)>=0.85))

mpg cyl disp hp drat wt qsec vs am gear carb
TRUE TRUE TRUE FALSE FALSE FALSE FALSE FALSE FALSE FALSE FALSE

```

I'll want to look at what MPG is correlated to so strongly, and decide what to keep and what to toss. Same for cyl and disp. Alternatively, I might need to combine some strongly correlated features.

# Chapter 104: Bibliography in RMD

| Parameter in YAML header | Detail                               |
|--------------------------|--------------------------------------|
| toc                      | table of contents                    |
| number_sections          | numbering the sections automatically |
| bibliography             | path to the bibliography file        |
| cs1                      | path to the style file               |

## Section 104.1: Specifying a bibliography and cite authors

The most important part of your RMD file is the YAML header. For writing an academic paper, I suggest to use PDF output, numbered sections and a table of content (toc).

```

```

```
title: "Writing an academic paper in R"
author: "Author"
date: "Date"
output:
pdf_document:
number_sections: yes
toc: yes
bibliography: bibliography.bib

```

In this example, our file `bibliography.bib` looks like this:

```
@ARTICLE{Meyer2000,
AUTHOR="Bernd Meyer",
TITLE="A constraint-based framework for diagrammatic reasoning",
JOURNAL="Applied Artificial Intelligence",
VOLUME= "14",
ISSUE = "4",
PAGES= "327--344",
YEAR=2000
}
```

To cite an author mentioned in your .bib file write @ and the bibkey, e.g. `Meyer2000`.

```
Introduction

`@Meyer2000` results in @Meyer2000.

`@Meyer2000 [p. 328]` results in @Meyer2000 [p. 328]

`[@Meyer2000]` results in [@Meyer2000]

`[-@Meyer2000]` results in [-@Meyer2000]

Summary

References
```

Rendering the RMD file via RStudio (Ctrl+Shift+K) or via console `rmarkdown::render("<path-to-your-RMD-file>")` results in the following output:



## Section 104.2: Inline references

If you have no \*.bib file, you can use a references field in the document's YAML metadata. This should include an array of YAML-encoded references, for example:

```

title: "Writing an academic paper in R"
author: "Author"
date: "Date"
output:
pdf_document:
number_sections: yes
toc: yes
references:
- id: Meyer2000
title: A Constraint-Based Framework for Diagrammatic Reasoning
author:
- family: Meyer
given: Bernd
volume: 14
issue: 4
publisher: Applied Artificial Intelligence
page: 327-344
type: article-journal
issued:
year: 2000

Introduction

`@Meyer2000` results in @Meyer2000.

`@Meyer2000 [p. 328]` results in @Meyer2000 [p. 328]

`[@Meyer2000]` results in [@Meyer2000]

`[-@Meyer2000]` results in [-@Meyer2000]

Summary

References

```

Rendering this file results in the same output as in example "Specifying a bibliography".

## Section 104.3: Citation styles

By default, pandoc will use a Chicago author-date format for citations and references. To use another style, you will need to specify a CSL 1.0 style file in the csl metadata field. In the following a often used citation style, the elsevier style, is presented (download at <https://github.com/citation-style-language/styles> ). The style-file has to be stored in the same directory as the RMD file OR the absolute path to the file has to be submitted.

To use another style then the default one, the following code is used:

```

title: "Writing an academic paper in R"
author: "Author"
date: "Date"
output:
pdf_document:
number_sections: yes
toc: yes
bibliography: bibliography.bib
csl: elsevier-harvard.csl

```

```
Introduction
```

```
`@Meyer2000` results in @Meyer2000.
`@Meyer2000 [p. 328]` results in @Meyer2000 [p. 328]
`[@Meyer2000]` results in [@Meyer2000]
`[-@Meyer2000]` results in [-@Meyer2000]
```

```
Summary
```

```
Reference
```



Notice the differences to the output of example "Specifying a bibliography and cite authors"

# Chapter 105: Writing functions in R

## Section 105.1: Anonymous functions

An anonymous function is, as the name implies, not assigned a name. This can be useful when the function is a part of a larger operation, but in itself does not take much place. One frequent use-case for anonymous functions is within the `*apply` family of Base functions.

Calculate the root mean square for each column in a `data.frame`:

```
df <- data.frame(first=5:9, second=(0:4)^2, third=-1:3)
apply(df, 2, function(x) { sqrt(sum(x^2)) })
 first second third
15.968719 18.814888 3.872983
```

Create a sequence of step-length one from the smallest to the largest value for each row in a matrix.

```
x <- sample(1:6, 12, replace=TRUE)
mat <- matrix(x, nrow=3)

apply(mat, 1, function(x) { seq(min(x), max(x)) })
```

An anonymous function can also stand on its own:

```
(function() { 1 })()
[1] 1
```

is equivalent to

```
f <- function() { 1 }
f()
[1] 1
```

## Section 105.2: RStudio code snippets

This is just a small hack for those who use self-defined functions often.

Type "fun" RStudio IDE and hit TAB.

The result will be a skeleton of a new function.

```
name <- function(variables) {
}
```

One can easily define their own snippet template, i.e. like the one below

```

name <- function(df, x, y) {
 require(tidyverse)
 out <-
 return(out)
}

```

The option is Edit Snippets in the Global Options -> Code menu.

## Section 105.3: Named functions

R is full of functions, it is after all a [functional programming language](#), but sometimes the precise function you need isn't provided in the Base resources. You could conceivably install a package containing the function, but maybe your requirements are just so specific that no pre-made function fits the bill? Then you're left with the option of making your own.

A function can be very simple, to the point of being pretty much pointless. It doesn't even need to take an argument:

```

one <- function() { 1 }
one()
[1] 1

two <- function() { 1 + 1 }
two()
[1] 2

```

What's between the curly braces { } is the function proper. As long as you can fit everything on a single line they aren't strictly needed, but can be useful to keep things organized.

A function can be very simple, yet highly specific. This function takes as input a vector (vec in this example) and outputs the same vector with the vector's length (6 in this case) subtracted from each of the vector's elements.

```

vec <- 4:9
subtract.length <- function(x) { x - length(x) }
subtract.length(vec)
[1] -2 -1 0 1 2 3

```

Notice that `length()` is in itself a pre-supplied (i.e. *Base*) function. You can of course use a previously self-made function within another self-made function, as well as assign variables and perform other operations while spanning several lines:

```

vec2 <- (4:7)/2

msdf <- function(x, multiplier=4) {
 mult <- x * multiplier
 subl <- subtract.length(x)
 data.frame(mult, subl)
}

msdf(vec2, 5)
 mult subl
1 10.0 -2.0
2 12.5 -1.5
3 15.0 -1.0
4 17.5 -0.5

```

`multiplier=4` makes sure that 4 is the default value of the argument `multiplier`, if no value is given when calling

the function 4 is what will be used.

The above are all examples of *named* functions, so called simply because they have been given names (one, two, subtract.length etc.)

# Chapter 106: Color schemes for graphics

## Section 106.1: viridis - print and colorblind friendly palettes

Viridis (named after the [chromis viridis fish](#)) is a recently [developed color scheme for the Python library matplotlib](#) (the video presentation by the link explains how the color scheme was developed and what are its main advantages). It is seamlessly ported to R.

There are 4 variants of color schemes: `magma`, `plasma`, `inferno`, and `viridis` (default). They are chosen with the `option` parameter and are coded as A, B, C, and D, correspondingly. To have an impression of the 4 color schemes, look at the maps:



([image source](#))

The package can be installed from [CRAN](#) or [github](#).

The [vignette](#) for `viridis` package is just brilliant.

Nice feature of the `viridis` color scheme is integration with `ggplot2`. Within the package two `ggplot2`-specific functions are defined: `scale_color_viridis()` and `scale_fill_viridis()`. See the example below:

```
library(viridis)
library(ggplot2)

gg1 <- ggplot(mtcars) +
```

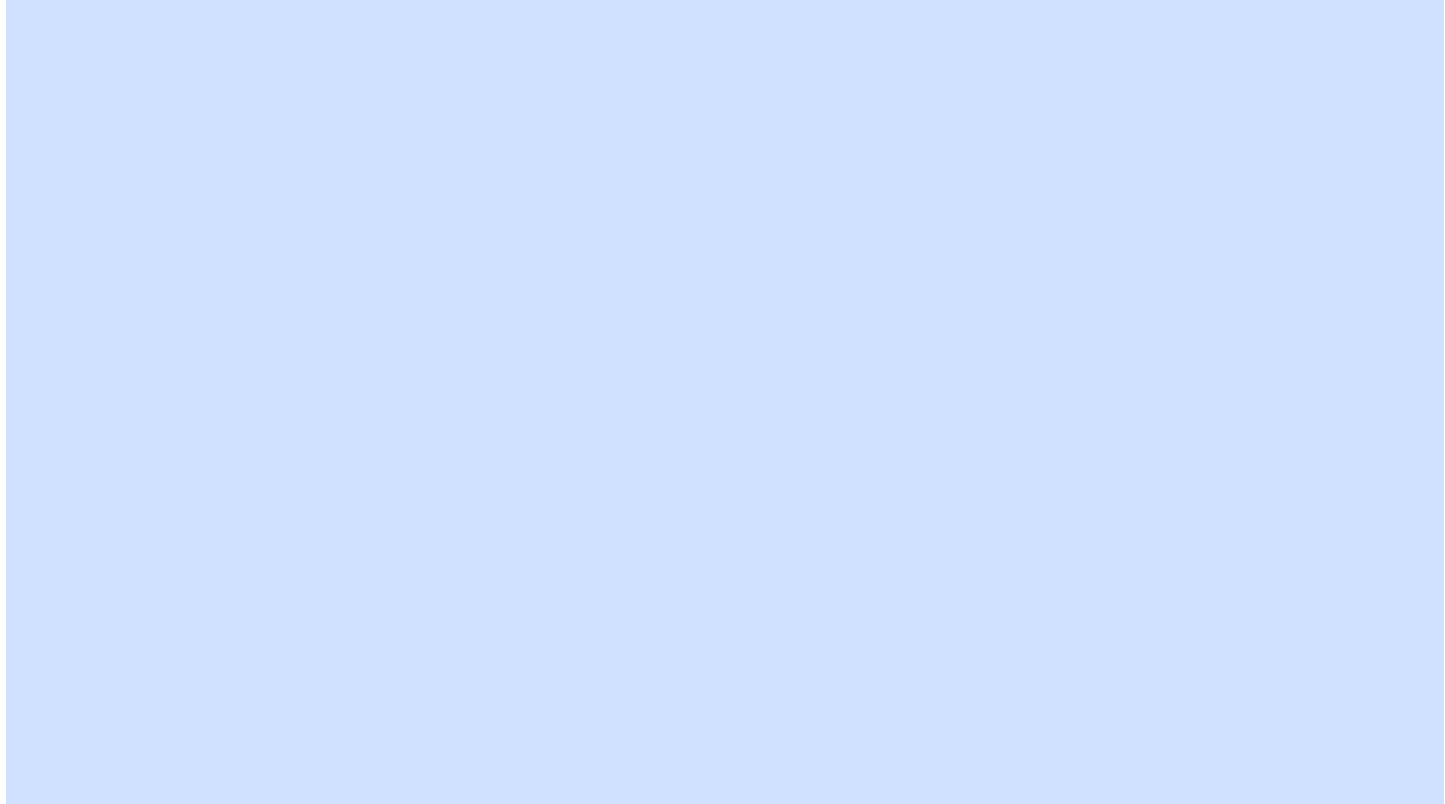
```

geom_point(aes(x = mpg, y = hp, color = disp), size = 3) +
 scale_color_viridis(option = "B") +
 theme_minimal() +
 theme(legend.position = c(.8,.8))

gg2 <- ggplot(mtcars) +
 geom_violin(aes(x = factor(cyl), y = hp, fill = factor(cyl))) +
 scale_fill_viridis(discrete = T) +
 theme_minimal() +
 theme(legend.position = 'none')

library(cowplot)
output <- plot_grid(gg1,gg2, labels = c('B','D'),label_size = 20)
print(output)

```



## Section 106.2: A handy function to glimpse a vector of colors

Quite often there is a need to glimpse the chosen color palette. One elegant solution is the following self defined function:

```

color_glimpse <- function(colors_string){
 n <- length(colors_string)
 hist(1:n, breaks=0:n, col=colors_string)
}

```

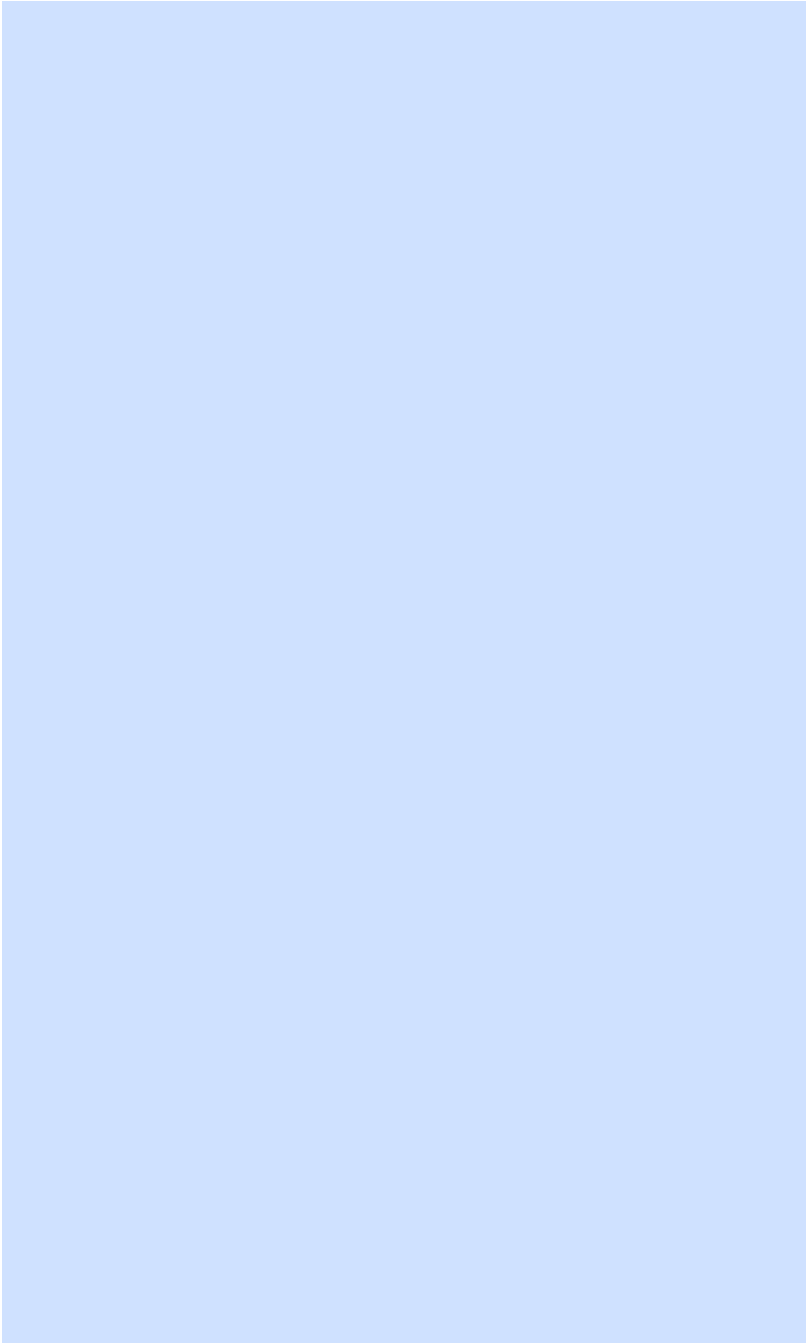
### An example of use

```
color_glimpse(blues9)
```



## Section 106.3: colorspace - click&drag interface for colors

The package `colorspace` provides GUI for selecting a palette. On the call of `choose_palette()` function the following window pops-up:



When the palette is chosen, just hit OK and do not forget to store the output in a variable, e.g. `pal`.

```
pal <- choose_palette()
```

The output is a function that takes `n` (number) as input and produces a color vector of length `n` according to the selected palette.

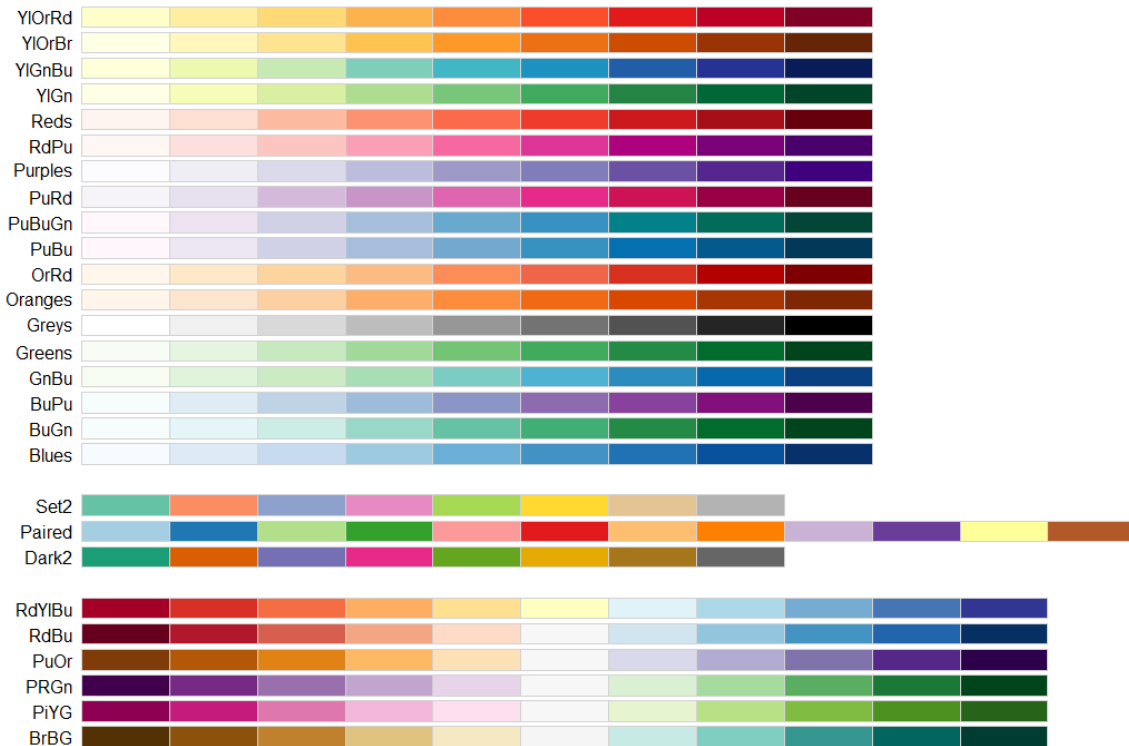
```
pal(10)
[1] "#023FA5" "#6371AF" "#959CC3" "#BEC1D4" "#DBDCE0" "#E0DBDC" "#D6BCC0" "#C6909A" "#AE5A6D"
 "#8E063B"
```

## Section 106.4: Colorblind-friendly palettes

Even though colorblind people can recognize a wide range of colors, it might be hard to differentiate between certain colors.

RColorBrewer provides colorblind-friendly palettes:

```
library(RColorBrewer)
display.brewer.all(colorblindFriendly = T)
```



The [Color Universal Design](#) from the University of Tokyo proposes the following palettes:

```
#palette using grey
cbPalette <- c("#999999", "#E69F00", "#56B4E9", "#009E73", "#F0E442", "#0072B2", "#D55E00",
 "#CC79A7")

#palette using black
cbbPalette <- c("#000000", "#E69F00", "#56B4E9", "#009E73", "#F0E442", "#0072B2", "#D55E00",
 "#CC79A7")
```

## Section 106.5: RColorBrewer

[ColorBrewer](#) project is a very popular tool to select harmoniously matching color palettes. RColorBrewer is a port of the project for R and provides also colorblind-friendly palettes.

### An example of use

```
colors_vec <- brewer.pal(5, name = 'BrBG')
print(colors_vec)
[1] "#A6611A" "#DFC27D" "#F5F5F5" "#80CDC1" "#018571"
```

RColorBrewer creates coloring options for ggplot2: scale\_color\_brewer and scale\_fill\_brewer.

```
library(ggplot2)
```

```
ggplot(mtcars)+
 geom_point(aes(x = mpg, y = hp, color = factor(cyl)), size = 3)+
 scale_color_brewer(palette = 'Greens')+
 theme_minimal()+
 theme(legend.position = c(.8, .8))
```

## Section 106.6: basic R color functions

Function `colors()` lists all the color names that are recognized by R. There is [a nice PDF](#) where one can actually see those colors.

`colorRampPalette` creates a function that interpolate a set of given colors to create new color palettes. This output function takes n (number) as input and produces a color vector of length n interpolating the initial colors.

```
pal <- colorRampPalette(c('white', 'red'))
pal(5)
[1] "#FFFFFF" "#FFBFBF" "#FF7F7F" "#FF3F3F" "#FF0000"
```

Any specific color may be produced with an `rgb()` function:

```
rgb(0,1,0)
```

produces green color.

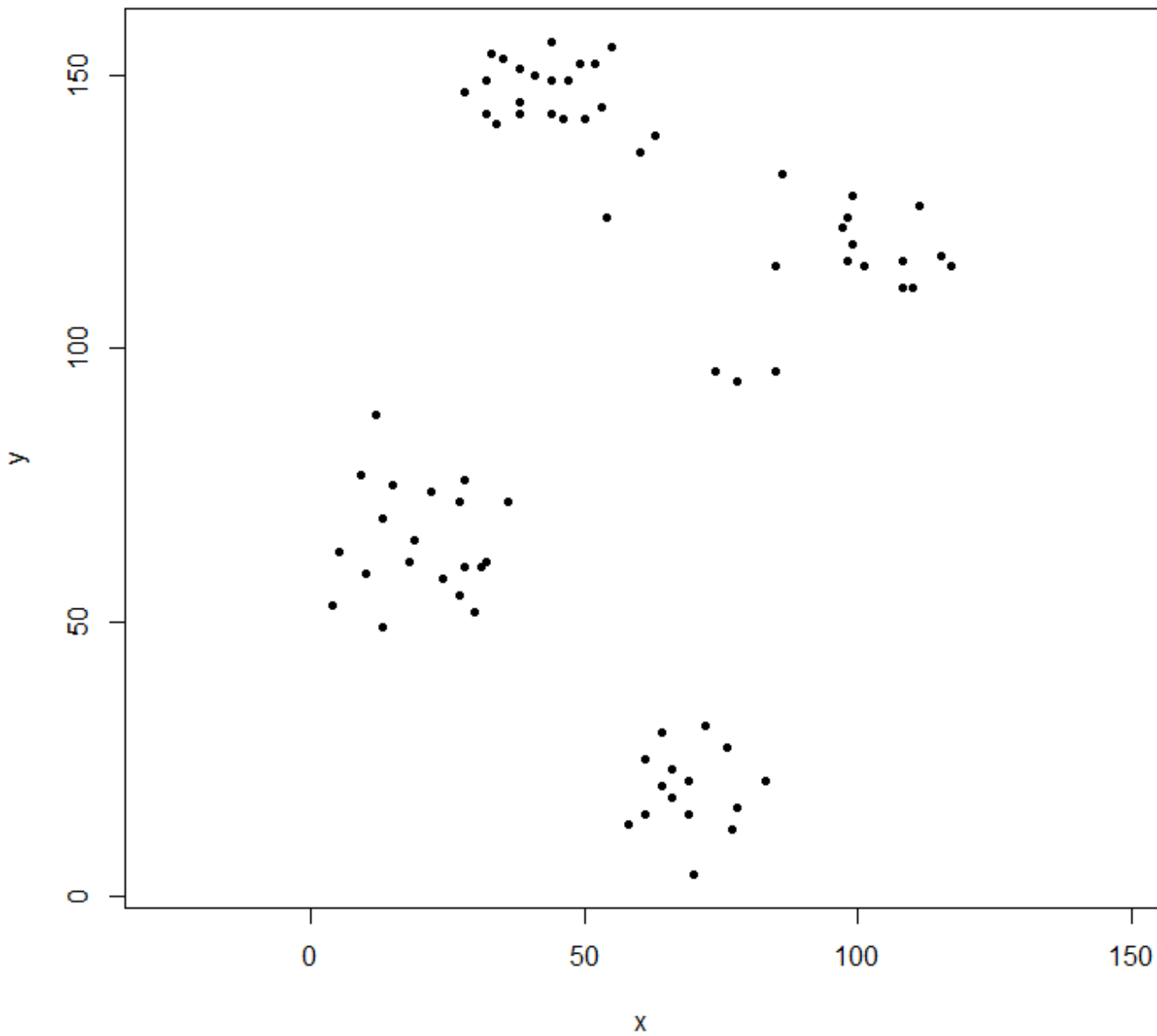
# Chapter 107: Hierarchical clustering with hclust

The stats package provides the **hclust** function to perform hierarchical clustering.

## Section 107.1: Example 1 - Basic use of hclust, display of dendrogram, plot clusters

The cluster library contains the ruspini data - a standard set of data for illustrating cluster analysis.

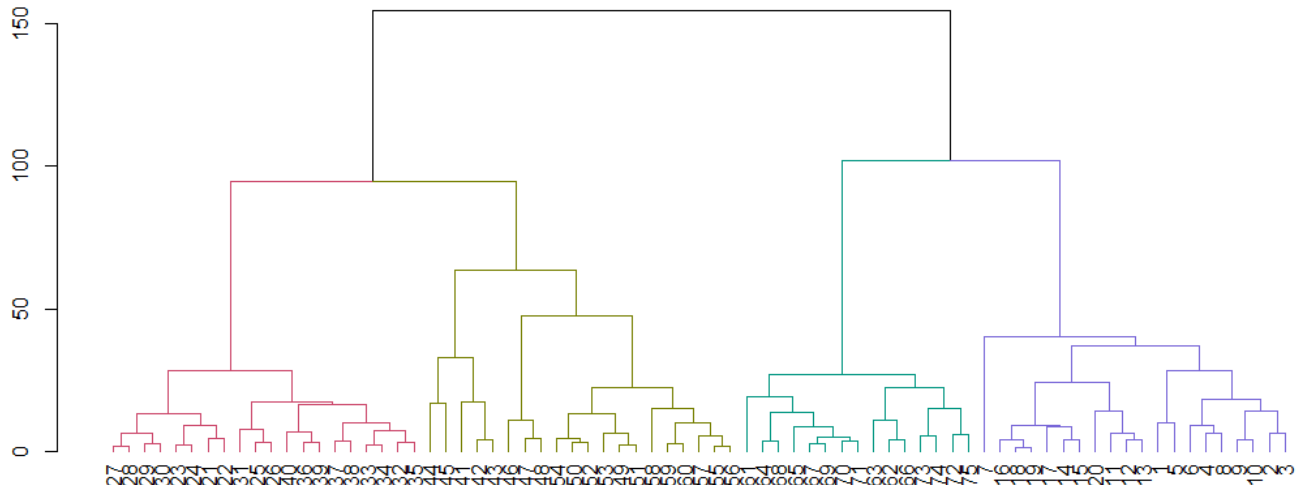
```
library(cluster) ## to get the ruspini data
plot(ruspini, asp=1, pch=20) ## take a look at the data
```



hclust expects a distance matrix, not the original data. We compute the tree using the default parameters and

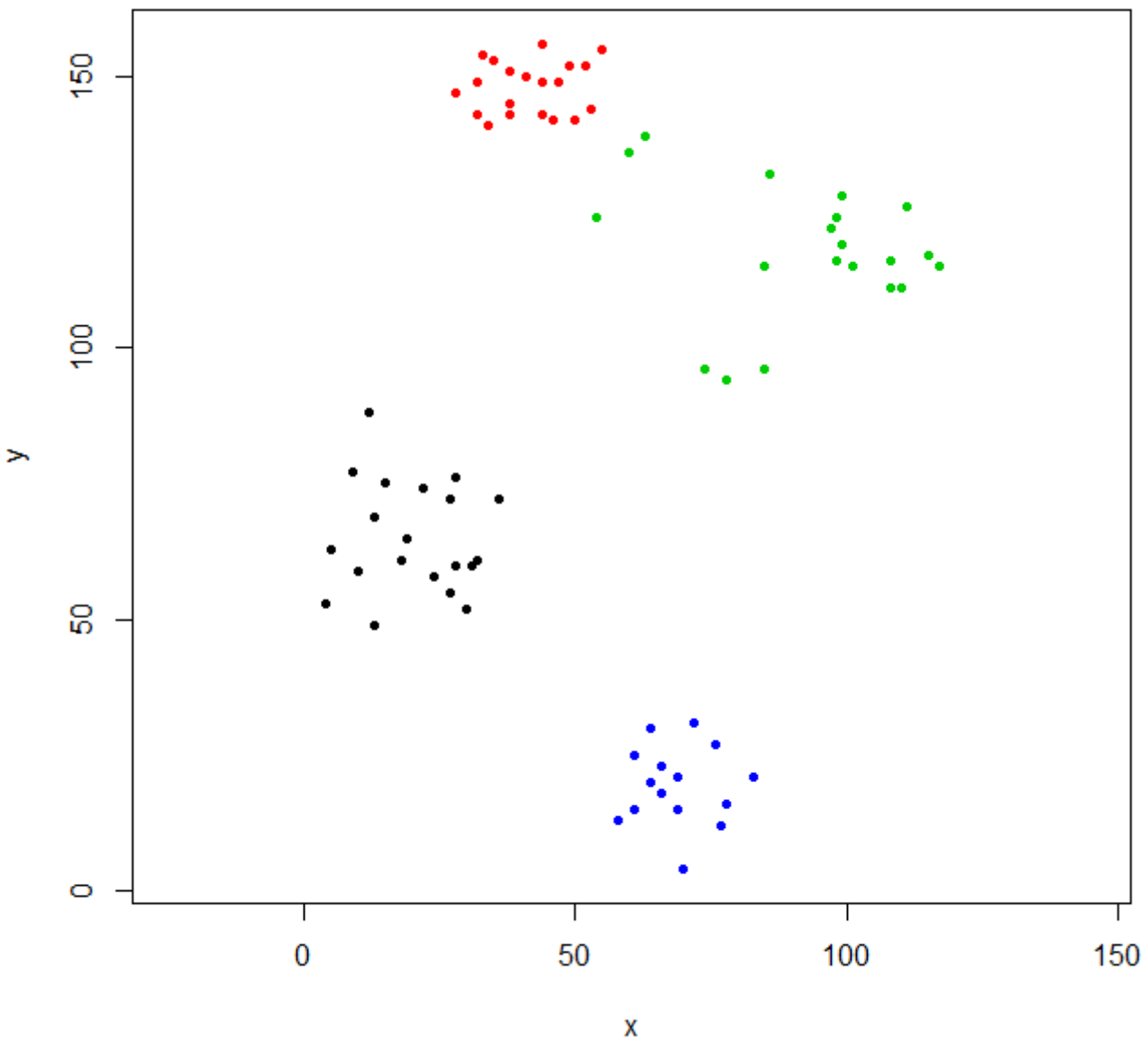
display it. The hang parameter lines up all of the leaves of the tree along the baseline.

```
ruspini_hc_defaults <- hclust(dist(ruspini))
dend <- as.dendrogram(ruspini_hc_defaults)
if(!require(dendextend)) install.packages("dendextend"); library(dendextend)
dend <- color_branches(dend, k = 4)
plot(dend)
```



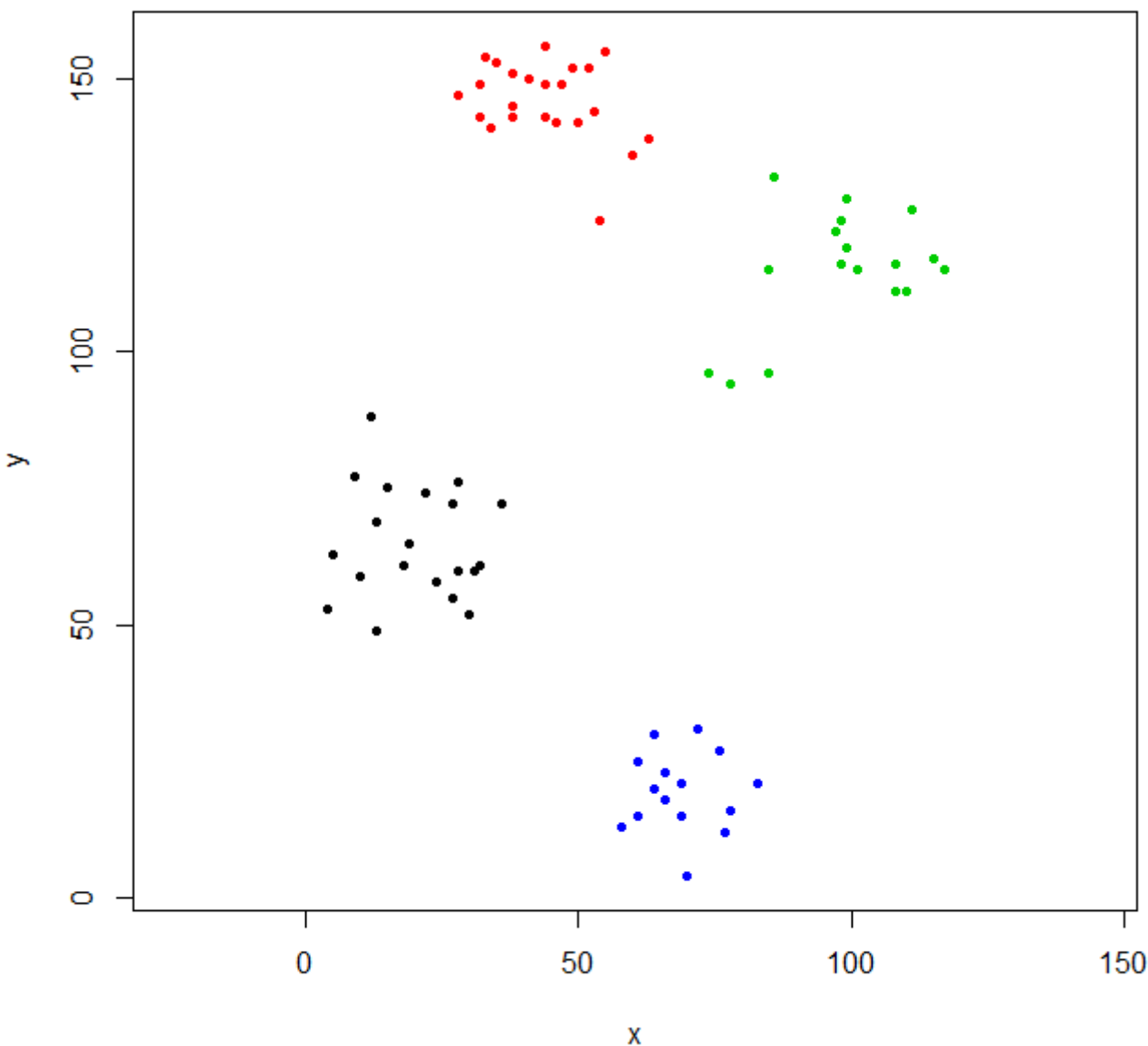
Cut the tree to give four clusters and replot the data coloring the points by cluster. k is the desired number of clusters.

```
rhc_def_4 = cutree(ruspini_hc_defaults,k=4)
plot(ruspini, pch=20, asp=1, col=rhc_def_4)
```



This clustering is a little odd. We can get a better clustering by scaling the data first.

```
scaled_ruspini_hc_defaults = hclust(dist(scale(ruspini)))
srhc_def_4 = cutree(scaled_ruspini_hc_defaults, 4)
plot(ruspini, pch=20, asp=1, col=srhc_def_4)
```



The default dissimilarity measure for comparing clusters is "complete". You can specify a different measure with the method parameter.

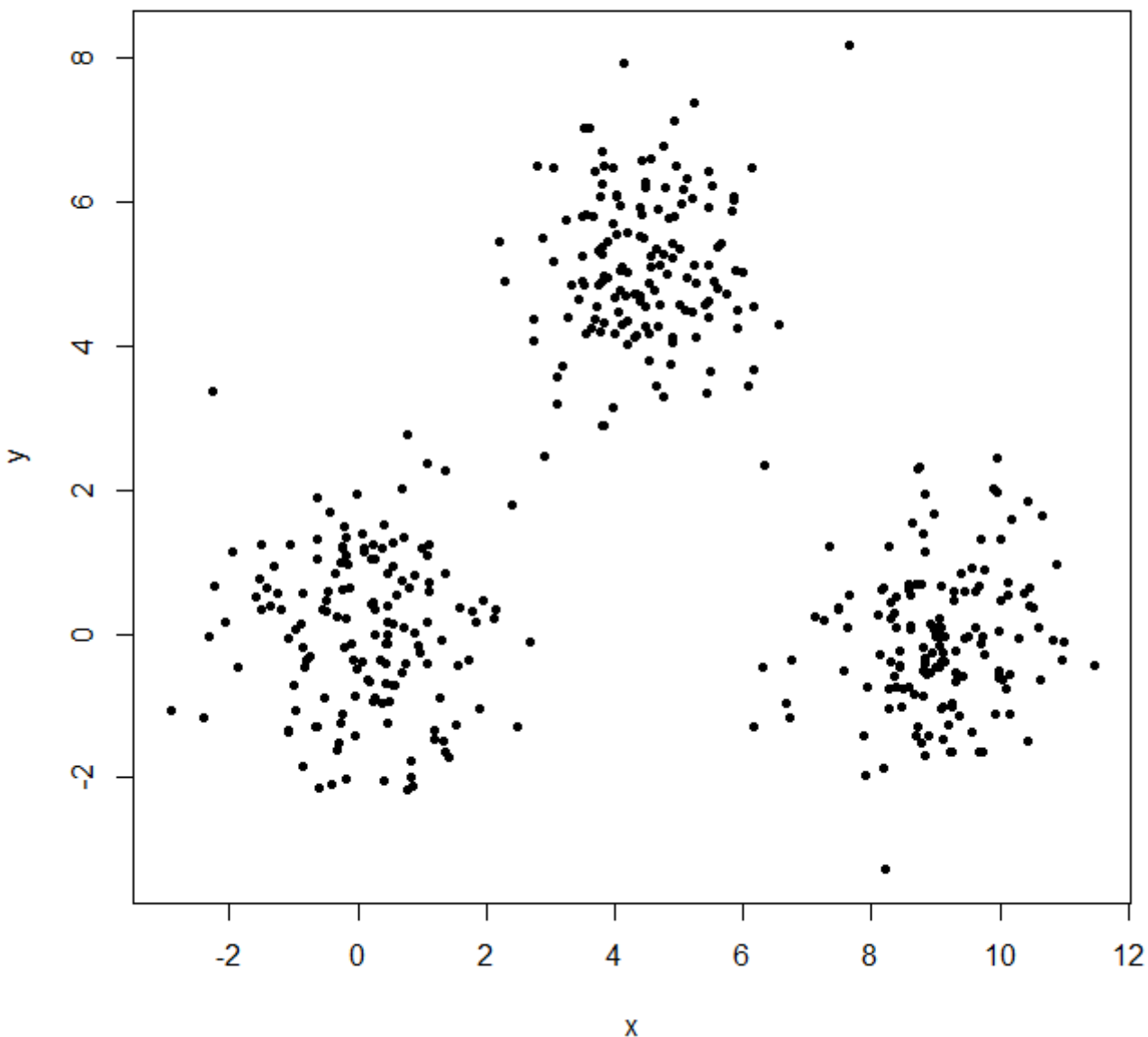
```
ruspini_hc_single = hclust(dist(ruspini), method="single")
```

## Section 107.2: Example 2 - hclust and outliers

With hierarchical clustering, outliers often show up as one-point clusters.

Generate three Gaussian distributions to illustrate the effect of outliers.

```
set.seed(656)
x = c(rnorm(150, 0, 1), rnorm(150, 9, 1), rnorm(150, 4.5, 1))
y = c(rnorm(150, 0, 1), rnorm(150, 0, 1), rnorm(150, 5, 1))
XYdf = data.frame(x,y)
plot(XYdf, pch=20)
```

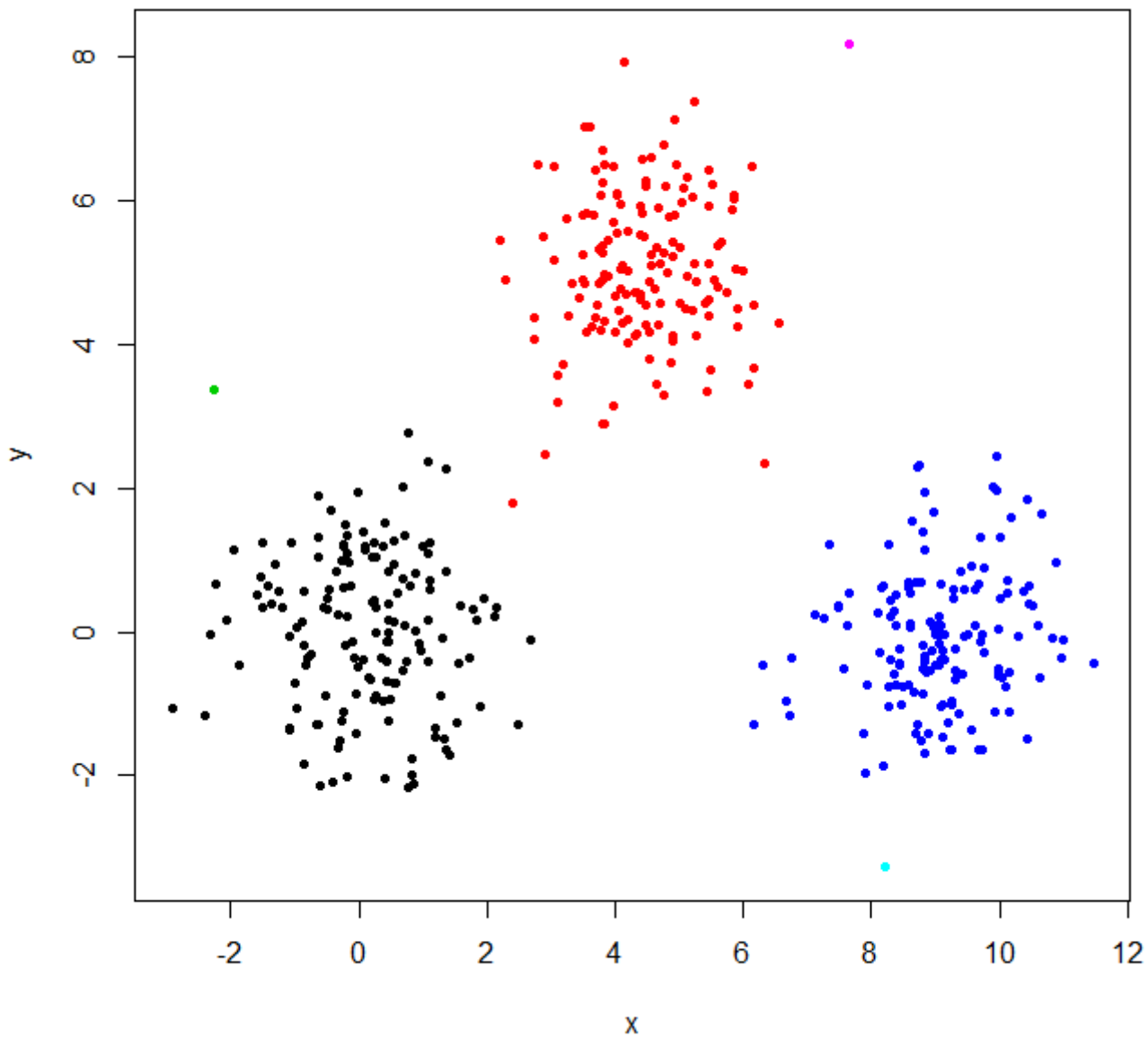


Build the cluster structure, split it into three cluster.

```
XY_sing = hclust(dist(XYdf), method="single")
XYs3 = cutree(XY_sing, k=3)
table(XYs3)
XYs3
 1 2 3
448 1 1
```

hclust found two outliers and put everything else into one big cluster. To get the "real" clusters, you may need to set k higher.

```
XYs6 = cutree(XY_sing, k=6)
table(XYs6)
XYs6
 1 2 3 4 5 6
148 150 1 149 1 1
plot(XYdf, pch=20, col=XYs6)
```



This [StackOverflow post](#) has some guidance on how to pick the number of clusters, but be aware of this behavior in hierarchical clustering.

# Chapter 108: Random Forest Algorithm

RandomForest is an ensemble method for classification or regression that reduces the chance of overfitting the data. Details of the method can be found in the [Wikipedia article on Random Forests](#). The main implementation for R is in the randomForest package, but there are other implementations. See the [CRAN view on Machine Learning](#).

## Section 108.1: Basic examples - Classification and Regression

```
Used for both Classification and Regression examples
library(randomForest)
library(car) ## For the Soils data
data(Soils)

#####
RF Classification Example
set.seed(656) ## for reproducibility
S_RF_Class = randomForest(Gp ~ ., data=Soils[,c(4,6:14)])
Gp_RF = predict(S_RF_Class, Soils[,6:14])
length(which(Gp_RF != Soils$Gp)) ## No Errors

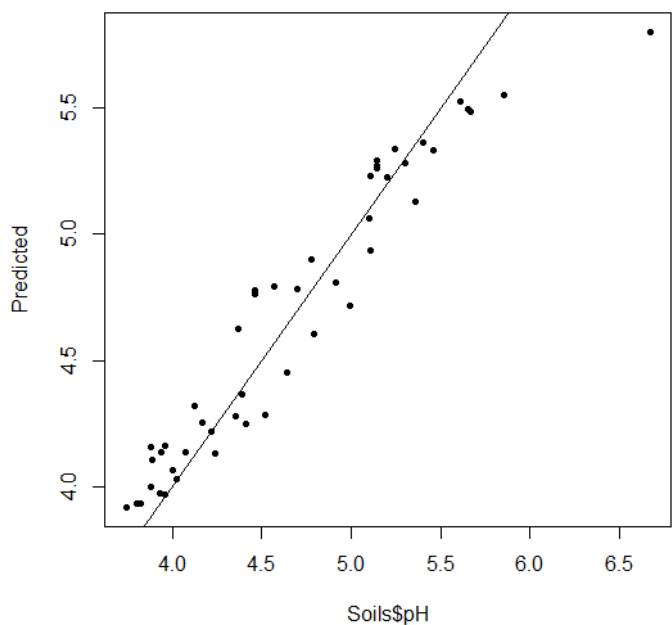
Naive Bayes for comparison
library(e1071)
S_NB = naiveBayes(Soils[,6:14], Soils[,4])
Gp_NB = predict(S_NB, Soils[,6:14], type="class")
length(which(Gp_NB != Soils$Gp)) ## 6 Errors
```

This example tested on the training data, but illustrates that RF can make very good models.

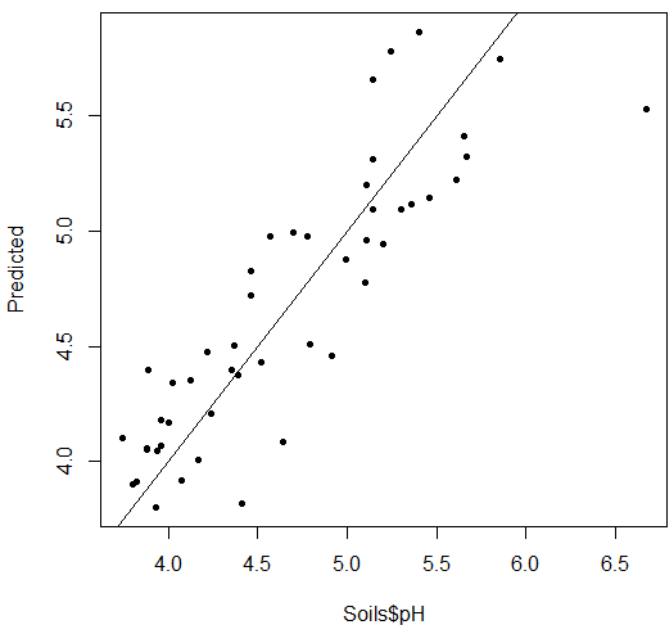
```
#####
RF Regression Example
set.seed(656) ## for reproducibility
S_RF_Reg = randomForest(pH ~ ., data=Soils[,6:14])
pH_RF = predict(S_RF_Reg, Soils[,6:14])

Compare Predictions with Actual values for RF and Linear Model
S_LM = lm(pH ~ ., data=Soils[,6:14])
pH_LM = predict(S_LM, Soils[,6:14])
par(mfrow=c(1,2))
plot(Soils$pH, pH_RF, pch=20, ylab="Predicted", main="Random Forest")
abline(0,1)
plot(Soils$pH, pH_LM, pch=20, ylab="Predicted", main="Linear Model")
abline(0,1)
```

**Random Forest**



**Linear Model**



# Chapter 109: RESTful R Services

[OpenCPU](#) uses standard R packaging to develop, ship and deploy web applications.

## Section 109.1: opencpu Apps

The official website contain good exemple of apps: <https://www.opencpu.org/apps.html>

The following code is used to serve a R session:

```
library(opencpu)
opencpu$start(port = 5936)
```

After this code is executed, you can use URLs to access the functions of the R session. The result could be XML, html, JSON or some other defined formats.

For exemple, the previous R session can be accessed by a cURL call:

```
#curl uses http post method for -X POST or -d "arg=value"
curl http://localhost:5936/opcpu/library/MASS/scripts/ch01.R -X POST
curl http://localhost:5936/opcpu/library/stats/R/rnorm -d "n=10&mean=5"
```

The call is asynchronous, meaning that the R session is not blocked while waiting for the call to finish (contrary to shiny).

The call result is kept in a temporary session stored in /ocpu/tmp/

An exemple of how to retrieve the temporary session:

```
curl https://public.opencpu.org/opcpu/library/stats/R/rnorm -d n=5
/ocpu/tmp/x009f9e7630/R/.val
/ocpu/tmp/x009f9e7630/stdout
/ocpu/tmp/x009f9e7630/source
/ocpu/tmp/x009f9e7630/console
/ocpu/tmp/x009f9e7630/info
```

x009f9e7630 is the name of the session.

Pointing to /ocpu/tmp/x009f9e7630/R/.val will return the value resulting of `rnorm(5)`,  
/ocpu/tmp/x009f9e7630/R/console will return the content of the console of `rnorm(5)`, etc..

# Chapter 110: Machine learning

## Section 110.1: Creating a Random Forest model

One example of machine learning algorithms is the Random Forest algorithm (Breiman, L. (2001). Random Forests. *Machine Learning* 45(5), p. 5-32). This algorithm is implemented in R according to Breiman's original Fortran implementation in the `randomForest` package.

Random Forest classifier objects can be created in R by preparing the class variable as `factor`, which is already apparent in the `iris` data set. Therefore we can easily create a Random Forest by:

```
library(randomForest)

rf <- randomForest(x = iris[, 1:4],
 y = iris$Species,
 ntree = 500,
 do.trace = 100)

rf

Call:
randomForest(x = iris[, 1:4], y = iris$Species, ntree = 500, do.trace = 100)
Type of random forest: classification
Number of trees: 500
No. of variables tried at each split: 2
#
OOB estimate of error rate: 4%
Confusion matrix:
setosa versicolor virginica class.error
setosa 50 0 0 0.00
versicolor 0 47 3 0.06
virginica 0 3 47 0.06
```

### parameters Description

- x a data frame holding the describing variables of the classes
- y the classes of the individual observations. If this vector is `factor`, a classification model is created, if not a regression model is created.
- ntree The number of individual CART trees built
- do.trace every  $i$ th step, the out-of-the-box errors overall and for each class are returned

# Chapter 111: Using texreg to export models in a paper-ready way

The texreg package helps to export a model (or several models) in a neat paper-ready way. The result may be exported as HTML or .doc (MS Office Word).

## Section 111.1: Printing linear regression results

```
models
fit1 <- lm(mpg ~ wt, data = mtcars)
fit2 <- lm(mpg ~ wt+hp, data = mtcars)
fit3 <- lm(mpg ~ wt+hp+cyl, data = mtcars)

export to html
texreg::htmlreg(list(fit1, fit2, fit3), file='models.html')

export to doc
texreg::htmlreg(list(fit1, fit2, fit3), file='models.doc')
```

The result looks like a table in a paper.

|             | Model 1            | Model 2            | Model 3            |
|-------------|--------------------|--------------------|--------------------|
| (Intercept) | 37.29***<br>(1.88) | 37.23***<br>(1.60) | 38.75***<br>(1.79) |
| wt          | -5.34***<br>(0.56) | -3.88***<br>(0.63) | -3.17***<br>(0.74) |
| hp          |                    | -0.03**<br>(0.01)  | -0.02<br>(0.01)    |
| cyl         |                    |                    | -0.94<br>(0.55)    |
| R2          | 0.75               | 0.83               | 0.84               |
| Adj. R2     | 0.74               | 0.81               | 0.83               |
| Num. obs.   | 32                 | 32                 | 32                 |
| RMSE        | 3.05               | 2.59               | 2.51               |

\*\*\*p < 0.001, \*\*p < 0.01, \*p < 0.05

### Statistical models

There are several additional handy parameters in `texreg::htmlreg()` function. Here is a use case for the most helpful parameters.

```
export to html
texreg::htmlreg(list(fit1, fit2, fit3), file='models.html',
 single.row = T,
 custom.model.names = LETTERS[1:3],
 leading.zero = F,
 digits = 3)
```

Which result in a table like this

|             | <b>A</b>          | <b>B</b>          | <b>C</b>          |
|-------------|-------------------|-------------------|-------------------|
| (Intercept) | 37.285 (1.878)*** | 37.227 (1.599)*** | 38.752 (1.787)*** |
| wt          | -5.344 (0.559)*** | -3.878 (0.633)*** | -3.167 (0.741)*** |
| hp          |                   | -0.032 (0.009)**  | -0.018 (0.012)    |
| cyl         |                   |                   | -0.942 (0.551)    |
| R2          | 0.753             | 0.827             | 0.843             |
| Adj. R2     | 0.745             | 0.815             | 0.826             |
| Num. obs.   | 32                | 32                | 32                |
| RMSE        | 3.046             | 2.593             | 2.512             |

\*\*\*p < 0.001, \*\*p < 0.01, \*p < 0.05

### Statistical models

# Chapter 112: Publishing

There are many ways of formatting R code, tables and graphs for publishing.

## Section 112.1: Formatting tables

Here, "table" is meant broadly (covering `data.frame`, `table`,

### Printing to plain text

Printing (as seen in the console) might suffice for a plain-text document to be viewed in monospaced font:

*Note: Before making the example data below, make sure you're in an empty folder you can write to. Run `getwd()` and read `?setwd` if you need to change folders.*

```
..w = options()$width
options(width = 500) # reduce text wrapping
sink(file = "mytab.txt")
 summary(mtcars)
sink()
options(width = ..w)
rm(..w)
```

### Printing delimited tables

Writing to CSV (or another common format) and then opening in a spreadsheet editor to apply finishing touches is another option:

*Note: Before making the example data below, make sure you're in an empty folder you can write to. Run `getwd()` and read `?setwd` if you need to change folders.*

```
write.csv(mtcars, file="mytab.csv")
```

### Further resources

- `knitr::kable`
- `stargazer`
- `tables::tabular`
- `texreg`
- `xtable`

## Section 112.2: Formatting entire documents

`Sweave` from the `utils` package allows for formatting code, prose, graphs and tables together in a LaTeX document.

### Further Resources

- Knitr and RMarkdown

# Chapter 113: Implement State Machine Pattern using S4 Class

[Finite States Machine](#) concepts are usually implemented under Object Oriented Programming (OOP) languages, for example using [Java language, based on the State pattern](#) defined in GOF (refers to the book: "Design Patterns").

R provides several mechanisms to simulate the OO paradigm, let's apply [S4 Object System](#) for implementing this pattern.

## Section 113.1: Parsing Lines using State Machine

Let's apply the [State Machine pattern](#) for parsing lines with the specific pattern using S4 Class feature from R.

### PROBLEM ENUNCIATION

We need to parse a file where each line provides information about a person, using a delimiter (" ; "), but some information provided is optional, and instead of providing an empty field, it is missing. On each line we can have the following information: Name ; [Address ; ]Phone. Where the address information is optional, sometimes we have it and sometimes don't, for example:

```
GREGORY BROWN; 25 NE 25TH; +1-786-987-6543
DAVID SMITH;786-123-4567
ALAN PEREZ; 25 SE 50TH; +1-786-987-5553
```

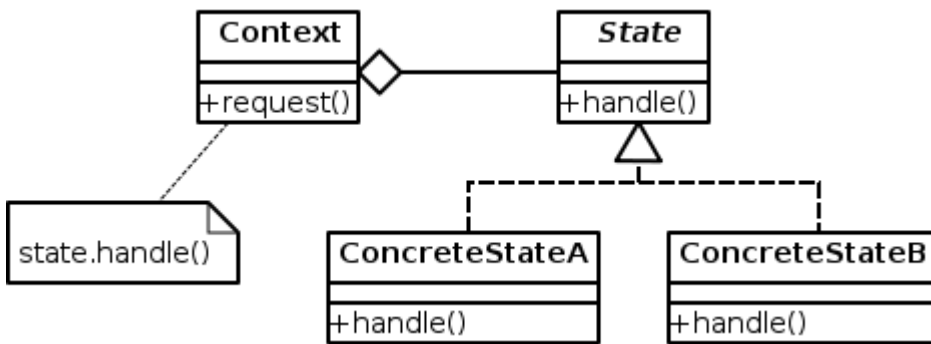
The second line does not provide address information. Therefore the number of delimiters may be different like in this case with one delimiter and for the other lines two delimiters. Because the number of delimiters may vary, one way to attack this problem is to recognize the presence or not of a given field based on its pattern. In such case we can use a [regular expression](#) for identifying such patterns. For example:

- **Name:** `^([A-Z] ?\\s+)* *[A-Z]+(\\s+[A-Z]{1,2}\\ .?, ? +)*[A-Z]+((-|\\s+)[A-Z]+)*$`. For example: RAFAEL REAL, DAVID R. SMITH, ERNESTO PEREZ GONZALEZ, O' CONNOR BROWN, LUIS PEREZ-MENA, etc.
- **Address:** `^\\s[0-9]{1,4}(\\s+[A-Z]{1,2}[0-9]{1,2}[A-Z]{1,2}|[A-Z\\s0-9]+)$`. For example: 11020 LE JEUNE ROAD, 87 SW 27TH. For the sake of simplicity we don't include here the zipcode, city, state, but I can be included in this field or adding additional fields.
- **Phone:** `^\\s*(\\+(-\\s+))*[0-9]{3}(-|\\s+)[0-9]{3}(-|\\s+)[0-9]{4}$`. For example: 305-123-4567, 305 123 4567, +1-786-123-4567.

### Notes:

- I am considering the most common pattern of US addresses and phones, it can be easily extended to consider more general situations.
- In R the sign "`\`" has special meaning for character variables, therefore we need to escape it.
- In order to simplify the process of defining regular expressions a good recommendation is to use the following web page: [regex101.com](#), so you can play with it, with a given example, until you get the expected result for all possible combinations.

The idea is to identify each line field based on previously defined patterns. The State pattern defines the following entities (classes) that collaborate to control the specific behavior (The State Pattern is a behavior pattern):



Let's describe each element considering the context of our problem:

- **Context:** Stores the context information of the parsing process, i.e. the current state and handles the entire State Machine Process. For each state, an action is executed (`handle()`), but the context delegates it, based on the state, on the action method defined for a particular state (`handle()` from State class). It defines the interface of interest to clients. Our Context class can be defined like this:
  - Attributes: `state`
  - Methods: `handle()`, ...
- **State:** The abstract class that represents any state of the State Machine. It defines an interface for encapsulating the behavior associated with a particular state of the context. It can be defined like this:
  - Attributes: `name`, `pattern`
  - Methods: `doAction()`, `isState` (using `pattern` attribute verify whether the input argument belong to this state pattern or not), ...
- **Concrete States (state sub-classes):** Each subclass of the class State that implements a behavior associated with a state of the Context. Our sub-classes are: `InitState`, `NameState`, `AddressState`, `PhoneState`. Such classes just implements the generic method using the specific logic for such states. No additional attributes are required.

**Note:** It is a matter of preference how to name the method that carries out the action, `handle()`, `doAction()` or `goNext()`. The method name `doAction()` can be the same for both classes (State or Context) we preferred to name as `handle()` in the Context class for avoiding a confusion when defining two generic methods with the same input arguments, but different class.

## PERSON CLASS

Using the S4 syntax we can define a Person class like this:

```

setClass(Class = "Person",
 slots = c(name = "character", address = "character", phone = "character")
)

```

It is a good recommendation to initialize the class attributes. The `setClass` documentation suggests using a generic method labeled as "`initialize`", instead of using deprecated attributes such as: `prototype`, `representation`.

```

setMethod("initialize", "Person",
 definition = function(.Object, name = NA_character_,
 address = NA_character_, phone = NA_character_) {
 .Object@name <- name
 .Object@address <- address
 .Object@phone <- phone
 .Object
 }
)

```

Because the `initialize` method is already a standard generic method of package `methods`, we need to respect the

original argument definition. We can verify it typing on R prompt:

```
> initialize
```

It returns the entire function definition, you can see at the top who the function is defined like:

```
function (.Object, ...) {....}
```

Therefore when we use `setMethod` we need to follow *exactly* the same syntax (`.Object`).

Another existing generic method is `show`, it is equivalent `toString()` method from Java and it is a good idea to have a specific implementation for class domain:

```
setMethod("show", signature = "Person",
 definition = function(object) {
 info <- sprintf("%s@[name='%s', address='%s', phone='%s']",
 class(object), object@name, object@address, object@phone)
 cat(info)
 invisible(NULL)
 }
)
```

**Note:** We use the same convention as in the default `toString()` Java implementation.

Let's say we want to save the parsed information (a list of Person objects) into a dataset, then we should be able first to convert a list of objects to into something the R can transform (for example coerce the object as a list). We can define the following additional method (for more detail about this see the [post](#))

```
setGeneric(name = "as.list", signature = c('x'),
 def = function(x) standardGeneric("as.list"))

Suggestion taken from here:
#
http://stackoverflow.com/questions/30386009/how-to-extend-as-list-in-a-canonical-way-to-s4-objects
setMethod("as.list", signature = "Person",
 definition = function(x) {
 mapply(function(y) {
 #apply as.list if the slot is again an user-defined object
 #therefore, as.list gets applied recursively
 if (inherits(slot(x,y), "Person")) {
 as.list(slot(x,y))
 } else {
 #otherwise just return the slot
 slot(x,y)
 }
 },
 slotNames(class(x)),
 SIMPLIFY=FALSE)
 }
)
```

R does not provide a sugar syntax for OO because the language was initially conceived to provide valuable functions for Statisticians. Therefore each user method requires two parts: 1) the Definition part (via `setGeneric`) and 2) the implementation part (via `setMethod`). Like in the above example.

## STATE CLASS

Following S4 syntax, let's define the abstract State class.

```

setClass(Class = "State", slots = c(name = "character", pattern = "character"))

setMethod("initialize", "State",
 definition = function(.Object, name = NA_character_, pattern = NA_character_) {
 .Object@name <- name
 .Object@pattern <- pattern
 .Object
 }
)

setMethod("show", signature = "State",
 definition = function(object) {
 info <- sprintf("%s@[name='%s', pattern='%s']", class(object),
 object@name, object@pattern)
 cat(info)
 invisible(NULL)
 }
)

setGeneric(name = "isState", signature = c('obj', 'input'),
 def = function(obj, input) standardGeneric("isState"))

setGeneric(name = "doAction", signature = c('obj', 'input', 'context'),
 def = function(obj, input, context) standardGeneric("doAction"))

```

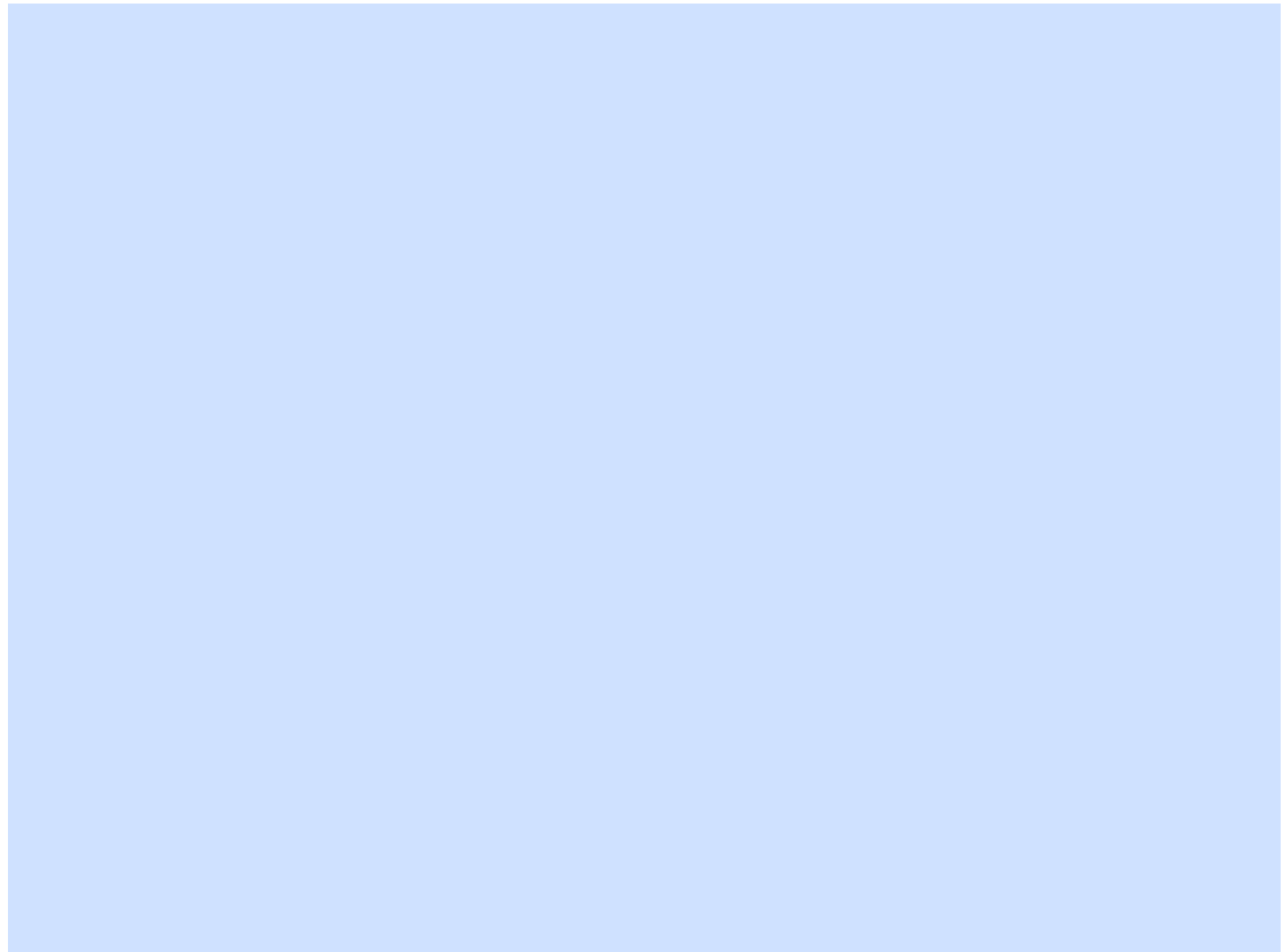
Every sub-class from State will have associated a name and pattern, but also a way to identify whether a given input belongs to this state or not (isState() method), and also implement the corresponding actions for this state (doAction() method).

In order to understand the process, let's define the transition matrix for each state based on the input received:

| <b>Input/Current State</b> | <b>Init</b> | <b>Name</b> | <b>Address</b> | <b>Phone</b> |
|----------------------------|-------------|-------------|----------------|--------------|
| Name                       |             | Name        |                |              |
| Address                    |             |             | Address        |              |
| Phone                      |             |             | Phone          | Phone        |
| End                        |             |             |                | End          |

**Note:** The cell [row, col]=[i, j] represents the destination state for the current state j, when it receives the input i.

It means that under the state Name it can receive two inputs: an address or a phone number. Another way to represents the transaction table is using the following [UML State Machine](#) diagram:



Let's implement each particular state as a sub-state of the class State

## STATE SUB-CLASSES

### Init State:

The initial state will be implemented via the following class:

```
setClass("InitState", contains = "State")

setMethod("initialize", "InitState",
 definition = function(.Object, name = "init", pattern = NA_character_) {
 .Object@name <- name
 .Object@pattern <- pattern
 .Object
 }
)

setMethod("show", signature = "InitState",
 definition = function(object) {
 callNextMethod()
 }
)
```

In R to indicate a class is a sub-class of other class is using the attribute `contains` and indicating the class name of the parent class.

Because the sub-classes just implement the generic methods, without adding additional attributes, then the **show** method, just call the equivalent method from the upper class (via method: **callNextMethod()**)

The initial state does not have associated a pattern, it just represents the beginning of the process, then we initialize the class with an NA value.

Now lets to implement the generic methods from the State class:

```
setMethod(f = "isState", signature = "InitState",
 definition = function(obj, input) {
 nameState <- new("NameState")
 result <- isState(nameState, input)
 return(result)
 }
)
```

For this particular state (without pattern), the idea it just initializes the parsing process expecting the first field will be a name, otherwise it will be an error.

```
setMethod(f = "doAction", signature = "InitState",
 definition = function(obj, input, context) {
 nameState <- new("NameState")
 if (isState(nameState, input)) {
 person <- context@person
 person@name <- trimws(input)
 context@person <- person
 context@state <- nameState
 } else {
 msg <- sprintf("The input argument: '%s' cannot be identified", input)
 stop(msg)
 }
 return(context)
 }
)
```

The doAction method provides the transition and updates the context with the information extracted. Here we are accessing to context information via the @-operator. Instead, we can define **get**/set methods, to encapsulate this process (as it is mandated in OO best practices: encapsulation), but that would add four more methods per get-set without adding value for the purpose of this example.

It is a good recommendation in all doAction implementation, to add a safeguard when the input argument is not properly identified.

## Name State

Here is the definition of this class definition:

```
setClass ("NameState", contains = "State")

setMethod("initialize","NameState",
 definition=function(.Object, name="name",
 pattern = "^([A-Z]'?\\s+)* *[A-Z]+(\\s+[A-Z]{1,2}\\.|.? +)*[A-Z]+((-|\\s+)[A-Z]+)*$") {
 .Object@pattern <- pattern
 .Object@name <- name
 .Object
 }
)
```

```

setMethod("show", signature = "NameState",
definition = function(object) {
 callNextMethod()
}
)

```

We use the function `grepl` for verifying the input belongs to a given pattern.

```

setMethod(f="isState", signature="NameState",
definition=function(obj, input) {
 result <- grepl(obj@pattern, input, perl=TRUE)
 return(result)
}
)

```

Now we define the action to carry out for a given state:

```

setMethod(f = "doAction", signature = "NameState",
definition=function(obj, input, context) {
 addressState <- new("AddressState")
 phoneState <- new("PhoneState")
 person <- context@person
 if (isState(addressState, input)) {
 person@address <- trimws(input)
 context@person <- person
 context@state <- addressState
 } else if (isState(phoneState, input)) {
 person@phone <- trimws(input)
 context@person <- person
 context@state <- phoneState
 } else {
 msg <- sprintf("The input argument: '%s' cannot be identified", input)
 stop(msg)
 }
 return(context)
}
)

```

Here we consider two possible transitions: one for Address state and the other one for Phone state. In all cases we update the context information:

- The `person` information: address or phone with the input argument.
- The state of the process

The way to identify the state is to invoke the method: `isState()` for a particular state. We create a default specific states (`addressState`, `phoneState`) and then ask for a particular validation.

The logic for the other sub-classes (one per state) implementation is very similar.

## Address State

```

setClass("AddressState", contains = "State")

setMethod("initialize", "AddressState",
definition = function(.Object, name="address",
 pattern = "^\s[0-9]{1,4}(\s+[A-Z]{1,2}[0-9]{1,2}[A-Z]{1,2}|[A-Z]\s[0-9]+)$") {
 .Object@pattern <- pattern
 .Object@name <- name
 .Object
}
)

```

```

 }
)

setMethod("show", signature = "AddressState",
 definition = function(object) {
 callNextMethod()
 }
)

setMethod(f="isState", signature="AddressState",
 definition=function(obj, input) {
 result <- grep1(obj@pattern, input, perl=TRUE)
 return(result)
 }
)

setMethod(f = "doAction", "AddressState",
 definition=function(obj, input, context) {
 phoneState <- new("PhoneState")
 if (isState(phoneState, input)) {
 person <- context@person
 person@phone <- trimws(input)
 context@person <- person
 context@state <- phoneState
 } else {
 msg <- sprintf("The input argument: '%s' cannot be identified", input)
 stop(msg)
 }
 return(context)
 }
)

```

## Phone State

```

setClass("PhoneState", contains = "State")

setMethod("initialize", "PhoneState",
 definition = function(.Object, name = "phone",
 pattern = "^\s*(\+|-)\s*\d{3}(\d{2})\s*\d{2}\s*\d{4}$") {
 .Object@pattern <- pattern
 .Object@name <- name
 .Object
 }
)

setMethod("show", signature = "PhoneState",
 definition = function(object) {
 callNextMethod()
 }
)

setMethod(f = "isState", signature = "PhoneState",
 definition = function(obj, input) {
 result <- grep1(obj@pattern, input, perl = TRUE)
 return(result)
 }
)

```

Here is where we add the person information into the list of persons of the context.

```
setMethod(f = "doAction", "PhoneState",
```

```

definition = function(obj, input, context) {
 context <- addPerson(context, context@person)
 context@state <- new("InitState")
 return(context)
}
)

```

## CONTEXT CLASS

Now let's explain the Context class implementation. We can define it considering the following attributes:

```

setClass(Class = "Context",
 slots = c(state = "State", persons = "list", person = "Person")
)

```

Where

- state: The current state of the process
- person: The current person, it represents the information we have already parsed from the current line.
- persons: The list of parsed persons processed.

**Note:** Optionally, we can add a name to identify the context by name in case we are working with more than one parser type.

```

setMethod(f="initialize", signature="Context",
 definition = function(.Object) {
 .Object@state <- new("InitState")
 .Object@persons <- listperson <- new("Person")
 return(.Object)
 }
)

setMethod("show", signature = "Context",
 definition = function(object) {
 cat("An object of class ", class(object), "\n", sep = "")
 info <- sprintf("[state='%s', persons='%s', person='%s']", object@state,
 toString(object@persons), object@person)
 cat(info)
 invisible(NULL)
 }
)

setGeneric(name = "handle", signature = c('obj', 'input', 'context'),
 def = function(obj, input, context) standardGeneric("handle"))

setGeneric(name = "addPerson", signature = c('obj', 'person'),
 def = function(obj, person) standardGeneric("addPerson"))

setGeneric(name = "parseLine", signature = c('obj', 's'),
 def = function(obj, s) standardGeneric("parseLine"))

setGeneric(name = "parseLines", signature = c('obj', 's'),
 def = function(obj, s) standardGeneric("parseLines"))

setGeneric(name = "as.df", signature = c('obj'),
 def = function(obj) standardGeneric("as.df"))

```

With such generic methods, we control the entire behavior of the parsing process:

- `handle()`: Will invoke the particular `doAction()` method of the current state.
- `addPerson`: Once we reach the end state, we need to add a `person` to the list of persons we have parsed.
- `parseLine()`: Parse a single line
- `parseLines()`: Parse multiple lines (an array of lines)
- `as.df()`: Extract the information from persons list into a data frame object.

Let's go on now with the corresponding implementations:

`handle()` method, delegates on `doAction()` method from the current state of the context:

```
setMethod(f = "handle", signature = "Context",
 definition = function(obj, input) {
 obj <- doAction(obj@state, input, obj)
 return(obj)
 }
)

setMethod(f = "addPerson", signature = "Context",
 definition = function(obj, person) {
 obj@persons <- c(obj@persons, person)
 return(obj)
 }
)
```

First, we split the original line in an array using the delimiter to identify each element via the R-function `strsplit()`, then iterate for each element as an input value for a given state. The `handle()` method returns again the context with the updated information (`state`, `person`, `persons` attribute).

```
setMethod(f = "parseLine", signature = "Context",
 definition = function(obj, s) {
 elements <- strsplit(s, ";")[[1]]
 # Adding an empty field for considering the end state.
 elements <- c(elements, "")
 n <- length(elements)
 input <- NULL
 for (i in 1:n)) {
 input <- elements[i]
 obj <- handle(obj, input)
 }
 return(obj@person)
 }
)
```

Because R makes a copy of the input argument, we need to return the context (`obj`):

```
setMethod(f = "parseLines", signature = "Context",
 definition = function(obj, s) {
 n <- length(s)
 listOfPersons <- list()
 for (i in 1:n)) {
 ipersons <- parseLine(obj, s[i])
 listOfPersons[[i]] <- ipersons
 }
 obj@persons <- listOfPersons
 return(obj)
 }
)
```

The attribute `persons` is a list of instance of S4 Person class. This something cannot be coerced to any standard type because R does not know of to treat an instance of a user defined class. The solution is to convert a Person into a list, using the `as.list` method previously defined. Then we can apply this function to each element of the list `persons`, via the `lapply()` function. Then in the next invocation to `lapply()` function, now applies the `data.frame` function for converting each element of the `persons.list` into a data frame. Finally, the `rbind()` function is called for adding each element converted as a new row of the data frame generated (for more detail about this see this [post](#))

```
Sugestion taken from this post:
http://stackoverflow.com/questions/4227223/r-list-to-data-frame
setMethod(f = "as.df", signature = "Context",
 definition = function(obj) {
 persons <- obj@persons
 persons.list <- lapply(persons, as.list)
 persons.ds <- do.call(rbind, lapply(persons.list, data.frame, stringsAsFactors = FALSE))
 return(persons.ds)
 }
)
```

## PUTTING ALL TOGETHER

Finally, lets to test the entire solution. Define the lines to parse where for the second line the address information is missing.

```
s <- c(
 "GREGORY BROWN; 25 NE 25TH; +1-786-987-6543",
 "DAVID SMITH;786-123-4567",
 "ALAN PEREZ; 25 SE 50TH; +1-786-987-5553"
)
```

Now we initialize the context, and parse the lines:

```
context <- new("Context")
context <- parseLines(context, s)
```

Finally obtain the corresponding dataset and print it:

```
df <- as.df(context)
> df
 name address phone
1 GREGORY BROWN 25 NE 25TH +1-786-987-6543
2 DAVID SMITH <NA> 786-123-4567
3 ALAN PEREZ 25 SE 50TH +1-786-987-5553
```

Let's test now the `show` methods:

```
> show(context@persons[[1]])
Person@[name='GREGORY BROWN', address='25 NE 25TH', phone='+1-786-987-6543']
```

And for some sub-state:

```
>show(new("PhoneState"))
PhoneState@[name='phone', pattern='^\\s*(\\+1(-|\\s+))*[0-9]{3}(-|\\s+)[0-9]{3}(-|\\s+)[0-9]{4}$']
```

Finally, test the `as.list()` method:

```
> as.list(context@persons[[1]])
$name
[1] "GREGORY BROWN"

$address
[1] "25 NE 25TH"

$phone
[1] "+1-786-987-6543"

>
```

## CONCLUSION

This example shows how to implement the State pattern, using one of the available mechanisms from R for using the OO paradigm. Nevertheless, the R OO solution is not user-friendly and differs so much from other OOP languages. You need to switch your mindset because the syntax is completely different, it reminds more the functional programming paradigm. For example instead of: `object.setID("A1")` as in Java/C#, for R you have to invoke the method in this way: `setID(object, "A1")`. Therefore you always have to include the object as an input argument to provide the context of the function. On the same way, there is no special `this` class attribute and either a `"."` notation for accessing methods or attributes of the given class. It is more error prompt because to refer a class or methods is done via attribute value (`"Person"`, `"isState"`, etc.).

Said the above, S4 class solution, requires much more lines of codes than a traditional Java/C# languages for doing simple tasks. Anyway, the State Pattern is a good and generic solution for such kind of problems. It simplifies the process delegating the logic into a particular state. Instead of having a big `if-else` block for controlling all situations, we have smaller `if-else` blocks inside on each State sub-class implementation for implementing the action to carry out in each state.

**Attachment:** [Here](#) you can download the entire script.

Any suggestion is welcome.

# Chapter 114: Reshape using tidyr

tidyr has two tools for reshaping data: gather (wide to long) and spread (long to wide).

See Reshaping data for other options.

## Section 114.1: Reshape from long to wide format with spread()

```
library(tidyr)

example data
set.seed(123)
df <- data.frame(
 name = rep(c("firstName", "secondName"), each=4),
 numbers = rep(1:4, 2),
 value = rnorm(8)
)
df
#> #> name numbers value
#> # 1 firstName 1 -0.56047565
#> # 2 firstName 2 -0.23017749
#> # 3 firstName 3 1.55870831
#> # 4 firstName 4 0.07050839
#> # 5 secondName 1 0.12928774
#> # 6 secondName 2 1.71506499
#> # 7 secondName 3 0.46091621
#> # 8 secondName 4 -1.26506123
```

We can "spread" the 'numbers' column, into separate columns:

```
spread(data = df,
 key = numbers,
 value = value)
#> #> name 1 2 3 4
#> # 1 firstName -0.5604756 -0.2301775 1.5587083 0.07050839
#> # 2 secondName 0.1292877 1.7150650 0.4609162 -1.26506123
```

Or spread the 'name' column into separate columns:

```
spread(data = df,
 key = name,
 value = value)
#> #> numbers firstName secondName
#> # 1 1 -0.56047565 0.1292877
#> # 2 2 -0.23017749 1.7150650
#> # 3 3 1.55870831 0.4609162
#> # 4 4 0.07050839 -1.2650612
```

## Section 114.2: Reshape from wide to long format with gather()

```
library(tidyr)

example data
df <- read.table(text = " numbers firstName secondName
1 1 1.5862639 0.4087477
2 2 0.1499581 0.9963923")
```

```
3 3 0.4117353 0.3740009
4 4 -0.4926862 0.4437916", header = T)
df
numbers firstName secondName
1 1 1.5862639 0.4087477
2 2 0.1499581 0.9963923
3 3 0.4117353 0.3740009
4 4 -0.4926862 0.4437916
```

We can gather the columns together using 'numbers' as the key column:

```
gather(data = df,
 key = numbers,
 value = myValue)
numbers numbers myValue
1 1 firstName 1.5862639
2 2 firstName 0.1499581
3 3 firstName 0.4117353
4 4 firstName -0.4926862
5 1 secondName 0.4087477
6 2 secondName 0.9963923
7 3 secondName 0.3740009
8 4 secondName 0.4437916
```

# Chapter 115: Modifying strings by substitution

**sub** and **gsub** are used to edit strings using patterns. See Pattern Matching and Replacement for more on related functions and Regular Expressions for how to build a pattern.

## Section 115.1: Rearrange character strings using capture groups

If you want to change the order of a character strings you can use parentheses in the pattern to group parts of the string together. These groups can in the replacement argument be addressed using consecutive numbers.

The following example shows how you can reorder a vector of names of the form "surname, forename" into a vector of the form "forename surname".

```
library(randomNames)
set.seed(1)

strings <- randomNames(5)
strings
[1] "Sigg, Zachary" "Holt, Jake" "Ortega, Sandra" "De La Torre, Nichole"
[5] "Perkins, Donovan"

sub("^(.+), \\\s(.+)$", "\\\\2 \\\\1", strings)
[1] "Zachary Sigg" "Jake Holt" "Sandra Ortega" "Nichole De La Torre"
[5] "Donovan Perkins"
```

If you only need the surname you could just address the first pairs of parentheses.

```
sub("^(.+), \\\s(.+)", "\\\\1", strings)
[1] "Sigg" "Holt" "Ortega" "De La Torre" "Perkins"
```

## Section 115.2: Eliminate duplicated consecutive elements

Let's say we want to eliminate duplicated subsequence element from a string (it can be more than one). For example:

```
2,14,14,14,19
```

and convert it into:

```
2,14,19
```

Using **gsub**, we can achieve it:

```
gsub("(\\d+)(, \\d+)+", "\\1", "2,14,14,14,19")
[1] "2,14,19"
```

It works also for more than one different repetition, for example:

```
> gsub("(\\d+)(, \\d+)+", "\\1", "2,14,14,14,19,19,20,21")
[1] "2,14,19,20,21"
```

Let's explain the regular expression:

1. `(\\d+)`: A group 1 delimited by () and finds any digit (at least one). Remember we need to use the double backslash (`\`) here because for a character variable a backslash represents special escape character for literal string delimiters (`\"` or `\'`). `\d\` is equivalent to: [0-9].
2. `,`: A punctuation sign: , (we can include spaces or any other delimiter)
3. `\\1`: An identical string to the group 1, i.e.: the repeated number. If that doesn't happen, then the pattern doesn't match.

Let's try a similar situation: eliminate consecutive repeated words:

```
one, two, two, three, four, four, five, six
```

Then, just replace `\d` by `\w`, where `\w` matches any word character, including: any letter, digit or underscore. It is equivalent to [a-zA-Z0-9\_]:

```
> gsub("(\\w+)(,\\w1)+", "\\1", "one, two, two, three, four, four, five, six")
[1] "one, two, three, four, five, six"
>
```

Then, the above pattern includes as a particular case duplicated digits case.

# Chapter 116: Non-standard evaluation and standard evaluation

Dplyr and many modern libraries in R use non-standard evaluation (NSE) for interactive programming and standard evaluation (SE) for programming<sup>1</sup>.

For instance, the `summarise()` function use non-standard evaluation but relies on the `summarise_()` which uses standard evaluation.

The `lazyeval` library makes it easy to turn standard evaluation function into NSE functions.

## Section 116.1: Examples with standard dplyr verbs

NSE functions should be used in interactive programming. However, when developing new functions in a new package, it's better to use SE version.

Load dplyr and lazyeval :

```
library(dplyr)
library(lazyeval)
```

### Filtering

NSE version

```
filter(mtcars, cyl == 8)
filter(mtcars, cyl < 6)
filter(mtcars, cyl < 6 & vs == 1)
```

SE version (to be use when programming functions in a new package)

```
filter_(mtcars, .dots = list(~ cyl == 8))
filter_(mtcars, .dots = list(~ cyl < 6))
filter_(mtcars, .dots = list(~ cyl < 6, ~ vs == 1))
```

### Summarise

NSE version

```
summarise(mtcars, mean(disp))
summarise(mtcars, mean_disp = mean(disp))
```

SE version

```
summarise_(mtcars, .dots = lazyeval::interp(~ mean(x), x = quote(disp)))
summarise_(mtcars, .dots = setNames(list(lazyeval::interp(~ mean(x), x = quote(disp))),
"mean_disp"))
summarise_(mtcars, .dots = list("mean_disp" = lazyeval::interp(~ mean(x), x = quote(disp))))
```

### Mutate

NSE version

```
mutate(mtcars, displ_l = disp / 61.0237)
```

```
mutate_(
 .data = mtcars,
 .dots = list(
 "displ_l" = lazyeval::interp(
 ~ x / 61.0237, x = quote(disp)
)
)
)
```

# Chapter 117: Randomization

The R language is commonly used for statistical analysis. As such, it contains a robust set of options for randomization. For specific information on sampling from probability distributions, see the documentation for distribution functions.

## Section 117.1: Random draws and permutations

The `sample` command can be used to simulate classic probability problems like drawing from an urn with and without replacement, or creating random permutations.

Note that throughout this example, `set.seed` is used to ensure that the example code is reproducible. However, `sample` will work without explicitly calling `set.seed`.

### Random permutation

In the simplest form, `sample` creates a random permutation of a vector of integers. This can be accomplished with:

```
set.seed(1251)
sample(x = 10)

[1] 7 1 4 8 6 3 10 5 2 9
```

When given no other arguments, `sample` returns a random permutation of the vector from 1 to x. This can be useful when trying to randomize the order of the rows in a data frame. This is a common task when creating randomization tables for trials, or when selecting a random subset of rows for analysis.

```
library(datasets)
set.seed(1171)
iris_rand <- iris[sample(x = 1:nrow(iris)),]

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

> head(iris_rand)
 Sepal.Length Sepal.Width Petal.Length Petal.Width Species
145 6.7 3.3 5.7 2.5 virginica
5 5.0 3.6 1.4 0.2 setosa
85 5.4 3.0 4.5 1.5 versicolor
137 6.3 3.4 5.6 2.4 virginica
128 6.1 3.0 4.9 1.8 virginica
105 6.5 3.0 5.8 2.2 virginica
```

### Draws without Replacement

Using `sample`, we can also simulate drawing from a set with and without replacement. To sample without replacement (the default), you must provide sample with a set to be drawn from and the number of draws. The set to be drawn from is given as a vector.

```
set.seed(7043)
sample(x = LETTERS, size = 7)
```

```
[1] "S" "P" "J" "F" "Z" "G" "R"
```

Note that if the argument to `size` is the same as the length of the argument to `x`, you are creating a random permutation. Also note that you cannot specify a size greater than the length of `x` when doing sampling without replacement.

```
set.seed(7305)
sample(x = letters, size = 26)

[1] "x" "z" "y" "i" "k" "f" "d" "s" "g" "v" "j" "o" "e" "c" "m" "n" "h" "u" "a" "b" "l" "r" "w" "t"
"q" "p"

sample(x = letters, size = 30)
Error in sample.int(length(x), size, replace, prob) :
 cannot take a sample larger than the population when 'replace = FALSE'
```

This brings us to drawing with replacement.

## Draws with Replacement

To make random draws from a set with replacement, you use the `replace` argument to `sample`. By default, `replace` is `FALSE`. Setting it to `TRUE` means that each element of the set being drawn from may appear more than once in the final result.

```
set.seed(5062)
sample(x = c("A", "B", "C", "D"), size = 8, replace = TRUE)

[1] "D" "C" "D" "B" "A" "A" "A" "A"
```

## Changing Draw Probabilities

By default, when you use `sample`, it assumes that the probability of picking each element is the same. Consider it as a basic "urn" problem. The code below is equivalent to drawing a colored marble out of an urn 20 times, writing down the color, and then putting the marble back in the urn. The urn contains one red, one blue, and one green marble, meaning that the probability of drawing each color is  $1/3$ .

```
set.seed(6472)
sample(x = c("Red", "Blue", "Green"),
 size = 20,
 replace = TRUE)
```

Suppose that, instead, we wanted to perform the same task, but our urn contains 2 red marbles, 1 blue marble, and 1 green marble. One option would be to change the argument we send to `x` to add an additional Red. However, a better choice is to use the `prob` argument to `sample`.

The `prob` argument accepts a vector with the probability of drawing each element. In our example above, the probability of drawing a red marble would be  $1/2$ , while the probability of drawing a blue or a green marble would be  $1/4$ .

```
set.seed(28432)
sample(x = c("Red", "Blue", "Green"),
 size = 20,
 replace = TRUE,
 prob = c(0.50, 0.25, 0.25))
```

Counter-intuitively, the argument given to `prob` does not need to sum to 1. R will always transform the given arguments into probabilities that total to 1. For instance, consider our above example of 2 Red, 1 Blue, and 1 Green.

You can achieve the same results as our previous code using those numbers:

```
set.seed(28432)
frac_prob_example <- sample(x = c("Red", "Blue", "Green"),
 size = 200,
 replace = TRUE,
 prob = c(0.50, 0.25, 0.25))

set.seed(28432)
numeric_prob_example <- sample(x = c("Red", "Blue", "Green"),
 size = 200,
 replace = TRUE,
 prob = c(2, 1, 1))

> identical(frac_prob_example, numeric_prob_example)
[1] TRUE
```

The major restriction is that you cannot set all the probabilities to be zero, and none of them can be less than zero.

You can also utilize prob when `replace` is set to FALSE. In that situation, after each element is drawn, the proportions of the prob values for the remaining elements give the probability for the next draw. In this situation, you must have enough non-zero probabilities to reach the size of the sample you are drawing. For example:

```
set.seed(21741)
sample(x = c("Red", "Blue", "Green"),
 size = 2,
 replace = FALSE,
 prob = c(0.8, 0.19, 0.01))
```

In this example, Red is drawn in the first draw (as the first element). There was an 80% chance of Red being drawn, a 19% chance of Blue being drawn, and a 1% chance of Green being drawn.

For the next draw, Red is no longer in the urn. The total of the probabilities among the remaining items is 20% (19% for Blue and 1% for Green). For that draw, there is a 95% chance the item will be Blue (19/20) and a 5% chance it will be Green (1/20).

## Section 117.2: Setting the seed

The `set.seed` function is used to set the random seed for all randomization functions. If you are using R to create a randomization that you want to be able to reproduce, you should use `set.seed` first.

```
set.seed(1643)
samp1 <- sample(x = 1:5, size = 200, replace = TRUE)

set.seed(1643)
samp2 <- sample(x = 1:5, size = 200, replace = TRUE)

> identical(x = samp1, y = samp2)
[1] TRUE
```

Note that parallel processing requires special treatment of the random seed, described more elsewhere.

# Chapter 118: Object-Oriented Programming in R

This documentation page describes the four object systems in R and their high-level similarities and differences. Greater detail on each individual system can be found on its own topic page.

The four systems are: S3, S4, Reference Classes, and S6.

## Section 118.1: S3

The S3 object system is a very simple OO system in R.

Every object has an S3 class. It can be get (got?) with the function `class`.

```
> class(3)
[1] "numeric"
```

It can also be set with the function `class`:

```
> bicycle <- 2
> class(bicycle) <- 'vehicle'
> class(bicycle)
[1] "vehicle"
```

It can also be set with the function `attr`:

```
> velocipede <- 2
> attr(velocipede, 'class') <- 'vehicle'
> class(velocipede)
[1] "vehicle"
```

An object can have many classes:

```
> class(x = bicycle) <- c('human-powered vehicle', class(x = bicycle))
> class(x = bicycle)
[1] "human-powered vehicle" "vehicle"
```

When using a generic function, R uses the first element of the class that has an available generic.

For example:

```
> summary.vehicle <- function(object, ...) {
+ message('this is a vehicle')
+ }
> summary(object = my_bike)
this is a vehicle
```

But if we now define a `summary.bicycle`:

```
> summary.bicycle <- function(object, ...) {
+ message('this is a bicycle')
+ }
> summary(object = my_bike)
this is a bicycle
```

# Chapter 119: Coercion

Coercion happens in R when the type of objects are changed during computation either implicitly or by using functions for explicit coercion (such as `as.numeric`, `as.data.frame`, etc.).

## Section 119.1: Implicit Coercion

Coercion happens with data types in R, often implicitly, so that the data can accommodate all the values. For example,

```
x = 1:3
x
[1] 1 2 3
typeof(x)
#[1] "integer"

x[2] = "hi"
x
#[1] "1" "hi" "3"
typeof(x)
#[1] "character"
```

Notice that at first, `x` is of type `integer`. But when we assigned `x[2] = "hi"`, all the elements of `x` were coerced into `character` as vectors in R can only hold data of single type.

# Chapter 120: Standardize analyses by writing standalone R scripts

If you want to routinely apply an R analysis to a lot of separate data files, or provide a repeatable analysis method to other people, an executable R script is a user-friendly way to do so. Instead of you or your user having to call R and execute your script inside R via `source( . )` or a function call, your user may simply call the script itself as if it was a program.

## Section 120.1: The basic structure of standalone R program and how to call it

### The first standalone R script

Standalone R scripts are not executed by the program R (R.exe under Windows), but by a program called Rscript (Rscript.exe), which is included in your R installation by default.

To hint at this fact, standalone R scripts start with a special line called **Shebang line**, which holds the following content: `#!/usr/bin/env Rscript`. Under Windows, an additional measure is needed, which is detailed later.

The following simple standalone R script saves a histogram under the file name "hist.png" from numbers it receives as input:

```
#!/usr/bin/env Rscript

User message (\n = end the line)
cat("Input numbers, separated by space:\n")
Read user input as one string (n=1 -> Read only one line)
input <- readLines(file('stdin'), n=1)
Split the string at each space (\s == any space)
input <- strsplit(input, "\s")[[1]]
convert the obtained vector of strings to numbers
input <- as.numeric(input)

Open the output picture file
png("hist.png", width=400, height=300)
Draw the histogram
hist(input)
Close the output file
dev.off()
```

You can see several key elements of a standalone R script. In the first line, you see the Shebang line. Followed by that, `cat("....\n")` is used to print a message to the user. Use `file("stdin")` whenever you want to specify "User input on console" as a data origin. This can be used instead of a file name in several data reading functions (`scan`, `read.table`, `read.csv`,...). After the user input is converted from strings to numbers, the plotting begins. There, it can be seen, that plotting commands which are meant to be written to a file must be enclosed in two commands. These are in this case `png( . )` and `dev.off()`. The first function depends on the desired output file format (other common choices being `jpeg( . )` and `pdf( . )`). The second function, `dev.off()` is always required. It writes the plot to the file and ends the plotting process.

### Preparing a standalone R script

#### Linux/Mac

The standalone script's file must first be made executable. This can happen by right-clicking the file, opening "Properties" in the opening menu and checking the "Executable" checkbox in the "Permissions" tab. Alternatively,

the command

```
chmod +x PATH/TO/SCRIPT/SCRIPTNAME.R
```

can be called in a Terminal.

## Windows

For each standalone script, a batch file must be written with the following contents:

```
"C:\Program Files\R-XXXXXX\bin\Rscript.exe" "%~dp0\XXXXXX.R" %*
```

A batch file is a normal text file, but which has a \*.bat extension except a \*.txt extension. Create it using a text editor like notepad (not Word) or similar and put the file name into quotation marks "FILENAME.bat") in the save dialog. To edit an existing batch file, right-click on it and select "Edit".

You have to adapt the code shown above everywhere XXX... is written:

- Insert the correct folder where your R installation resides
- Insert the correct name of your script and place it into the same directory as this batch file.

Explanation of the elements in the code: The first part "C:\...\Rscript.exe" tells Windows where to find the Rscript.exe program. The second part "%~dp0\XXX.R" tells Rscript to execute the R script you've written which resides in the same folder as the batch file (%dp0 stands for the batch file folder). Finally, %\* forwards any command line arguments you give to the batch file to the R script.

If you double-click on the batch file, the R script is executed. If you drag files on the batch file, the corresponding file names are given to the R script as command line arguments.

## Section 120.2: Using littler to execute R scripts

[littler](#) (pronounced *little r*) ([cran](#)) provides, besides other features, two possibilities to run R scripts from the command line with littler's r command (when one works with Linux or MacOS).

### Installing littler

**From R:**

```
install.packages("littler")
```

The path of r is printed in the terminal, like

```
You could link to the 'r' binary installed in
'/home/*USER*/R/x86_64-pc-linux-gnu-library/3.4/littler/bin/r'
from '/usr/local/bin' in order to use 'r' for scripting.
```

To be able to call r from the system's command line, a symlink is needed:

```
ln -s /home/*USER*/R/x86_64-pc-linux-gnu-library/3.4/littler/bin/r /usr/local/bin/r
```

**Using apt-get (Debian, Ubuntu):**

```
sudo apt-get install littler
```

### Using littler with standard .r scripts

With r from littler it is possible to execute standalone R scripts without any changes to the script. Example script:

```

User message (\n = end the line)
cat("Input numbers, separated by space:\n")
Read user input as one string (n=1 -> Read only one line)
input <- readLines(file('stdin'), n=1)
Split the string at each space (\s == any space)
input <- strsplit(input, "\s")[[1]]
convert the obtained vector of strings to numbers
input <- as.numeric(input)

Open the output picture file
png("hist.png", width=400, height=300)
Draw the histogram
hist(input)
Close the output file
dev.off()

```

Note that no shebang is at the top of the scripts. When saved as for example `hist.r`, it is directly callable from the system command:

```
r hist.r
```

### Using littler on *shebanged* scripts

It is also possible to create executable R scripts with littler, with the use of the shebang

```
#!/usr/bin/env r
```

at the top of the script. The corresponding R script has to be made executable with `chmod +X /path/to/script.r` and is directly callable from the system terminal.

# Chapter 121: Analyze tweets with R

(Optional) Every topic has a focus. Tell the readers what they will find here and let future contributors know what belongs.

## Section 121.1: Download Tweets

The first think you need to do is to download tweets. You need to Setup your tweeter account. Much Information can be found in Internet on how to do it. The following two links were useful for my Setup (last checked in May 2017)

In particular I found the following two links useful (last checked in May 2017):

[Link 1](#)

[Link 2](#)

### R Libraries

You will need the following R packages

```
library("devtools")
library("twitteR")
library("ROAuth")
```

Supposing you have your keys You have to run the following code

```
api_key <- XXXXXXXXXXXXXXXXXXXXXXXX
api_secret <- XXXXXXXXXXXXXXXXXXXXXXXX
access_token <- XXXXXXXXXXXXXXXXXXXXXXXX
access_token_secret <- XXXXXXXXXXXXXXXXXXXXXXXX

setup_twitter_oauth(api_key, api_secret)
```

Change XXXXXXXXXXXXXXXXXXXXXXXX to your keys (if you have Setup your tweeter account you know which keys I mean).

Let's now suppose we want to download tweets on coffee. The following code will do it

```
search.string <- "#coffee"
no.of.tweets <- 1000

c_tweets <- searchTwitter(search.string, n=no.of.tweets, lang="en")
```

You will get 1000 tweets on "coffee".

## Section 121.2: Get text of tweets

Now we need to access the text of the tweets. So we do it in this way (we also need to clean up the tweets from special characters that for now we don't need, like emoticons with the sapply function.)

```
coffee_tweets = sapply(c_tweets, function(t) t$getText())
```

```
coffee_tweets <- sapply(coffee_tweets, function(row) iconv(row, "latin1", "ASCII", sub=""))
```

and you can check your tweets with the **head** function.

```
head(coffee_tweets)
```

# Chapter 122: Natural language processing

Natural language processing (NLP) is the field of computer sciences focused on retrieving information from textual input generated by human beings.

## Section 122.1: Create a term frequency matrix

The simplest approach to the problem (and the most commonly used so far) is to split sentences into *tokens*. Simplifying, *words* have abstract and subjective meanings to the people using and receiving them, *tokens* have an objective interpretation: an ordered sequence of characters (or bytes). Once sentences are split, the order of the token is disregarded. This approach to the problem is known as **bag of words** model.

A **term frequency** is a dictionary, in which to each token is assigned a *weight*. In the first example, we construct a term frequency matrix from a corpus **corpus** (a collection of **documents**) with the R package **tm**.

```
require(tm)
doc1 <- "drugs hospitals doctors"
doc2 <- "smog pollution environment"
doc3 <- "doctors hospitals healthcare"
doc4 <- "pollution environment water"
corpus <- c(doc1, doc2, doc3, doc4)
tm_corpus <- Corpus(VectorSource(corpus))
```

In this example, we created a corpus of class **Corpus** defined by the package **tm** with two functions **Corpus** and **VectorSource**, which returns a **VectorSource** object from a character vector. The object **tm\_corpus** is a list our documents with additional (and optional) metadata to describe each document.

```
str(tm_corpus)
List of 4
 $ 1:List of 2
 ..$ content: chr "drugs hospitals doctors"
 ..$ meta :List of 7
 ...$ author : chr(0)
 ...$ timestamp: POSIXlt[1:1], format: "2017-06-03 00:31:34"
 ...$ description : chr(0)
 ...$ heading : chr(0)
 ...$ id : chr "1"
 ...$ language : chr "en"
 ...$ origin : chr(0)
 ...- attr(*, "class")= chr "TextDocumentMeta"
 - attr(*, "class")= chr [1:2] "PlainTextDocument" "TextDocument"
 [truncated]
```

Once we have a **Corpus**, we can proceed to preprocess the tokens contained in the **Corpus** to improve the quality of the final output (the term frequency matrix). To do this we use the **tm** function **tm\_map**, which similarly to the **apply** family of functions, transform the documents in the corpus by applying a function to each document.

```
tm_corpus <- tm_map(tm_corpus, tolower)
tm_corpus <- tm_map(tm_corpus, removeWords, stopwords("english"))
tm_corpus <- tm_map(tm_corpus, removeNumbers)
tm_corpus <- tm_map(tm_corpus, PlainTextDocument)
tm_corpus <- tm_map(tm_corpus, stemDocument, language="english")
tm_corpus <- tm_map(tm_corpus, stripWhitespace)
tm_corpus <- tm_map(tm_corpus, PlainTextDocument)
```

Following these transformations, we finally create the term frequency matrix with

```
tdm <- TermDocumentMatrix(tm_corpus)
```

which gives a

```
<<TermDocumentMatrix (terms: 8, documents: 4)>>
Non-/sparse entries: 12/20
Sparsity : 62%
Maximal term length: 9
Weighting : term frequency (tf)
```

that we can view by transforming it to a matrix

```
as.matrix(tdm)
```

| Terms     | Docs         |              |              |              |
|-----------|--------------|--------------|--------------|--------------|
|           | character(0) | character(0) | character(0) | character(0) |
| doctor    | 1            | 0            | 1            | 0            |
| drug      | 1            | 0            | 0            | 0            |
| environ   | 0            | 1            | 0            | 1            |
| healthcar | 0            | 0            | 1            | 0            |
| hospit    | 1            | 0            | 1            | 0            |
| pollut    | 0            | 1            | 0            | 1            |
| smog      | 0            | 1            | 0            | 0            |
| water     | 0            | 0            | 0            | 1            |

Each row represents the frequency of each token - that as you noticed have been stemmed (e.g. **environment** to **environ**) - in each document (4 documents, 4 columns).

In the previous lines, we have weighted each pair token/document with the absolute frequency (i.e. the number of instances of the token that appear in the document).

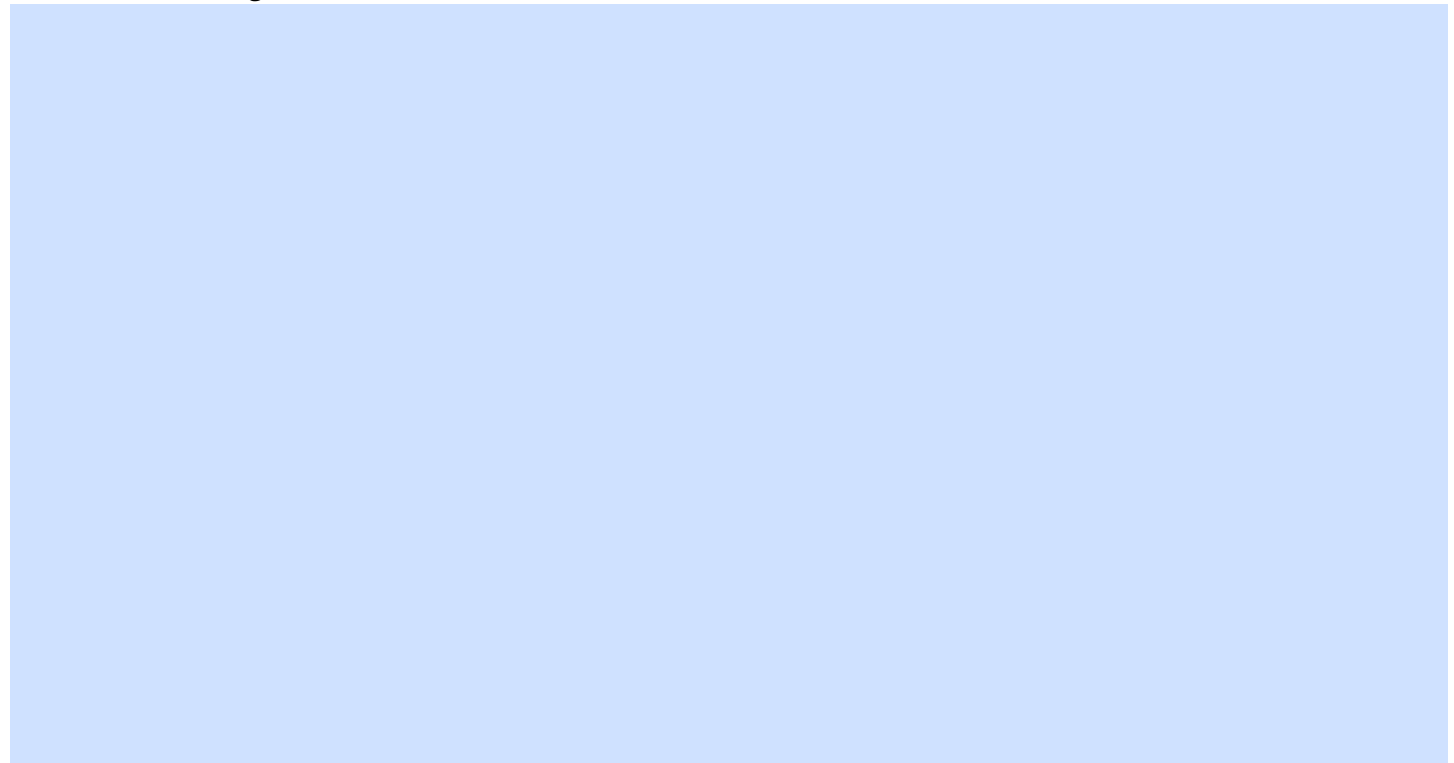
# Chapter 123: R Markdown Notebooks (from RStudio)

An R Notebook is an R Markdown document with chunks that can be executed independently and interactively, with output visible immediately beneath the input. They are similar to R Markdown documents with the exception of results being displayed in the R Notebook creation/edit mode rather than in the rendered output. **Note:** R Notebooks are new feature of RStudio and are only available in version 1.0 or higher of RStudio.

## Section 123.1: Creating a Notebook

You can create a new notebook in RStudio with the menu command File -> New File -> R Notebook

If you don't see the option for R Notebook, then you need to update your version of RStudio. For installation of RStudio follow this guide



## Section 123.2: Inserting Chunks

Chunks are pieces of code that can be executed interactively. In-order to insert a new chunk by clicking on the **Insert** button present on the notebook toolbar and select your desired code platform (R in this case, since we want to write R code). Alternatively we can use keyboard shortcuts to insert a new chunk **Ctrl + Alt + I (OS X: Cmd + Option + I)**



## Section 123.3: Executing Chunk Code

You can run the current chunk by clicking **Run current Chunk (green play button)** present on the right side of the chunk. Alternatively we can use keyboard shortcut **Ctrl + Shift + Enter (OS X: Cmd + Shift + Enter)**

The output from all the lines in the chunk will appear beneath the chunk.

### Splitting Code into Chunks

Since a chunk produces its output beneath the chunk, when having multiple lines of code in a single chunk that produces multiple outputs it is often helpful to split into multiple chunks such that each chunk produces one output.

To do this, select the code to you want to split into a new chunk and press **Ctrl + Alt + I (OS X: Cmd + Option + I)**



## Section 123.4: Execution Progress

When you execute code in a notebook, an indicator will appear in the gutter to show you execution progress. Lines of code which have been sent to R are marked with dark green; lines which have not yet been sent to R are marked with light green.

### Executing Multiple Chunks

Running or Re-Running individual chunks by pressing Run for all the chunks present in a document can be painful. We can use **Run All** from the Insert menu in the toolbar to Run all the chunks present in the notebook. Keyboard shortcut is **Ctrl + Alt + R (OS X: Cmd + Option + R)**

There's also a option **Restart R and Run All Chunks** command (available in the Run menu on the editor toolbar), which gives you a fresh R session prior to running all the chunks.

We also have options like **Run All Chunks Above** and **Run All Chunks Below** to run chunks Above or Below from a selected chunk.



## Section 123.5: Preview Output

Before rendering the final version of a notebook we can preview the output. Click on the **Preview** button on the toolbar and select the desired output format.

You can change the type of output by using the output options as "pdf\_document" or "html\_notebook"

## Section 123.6: Saving and Sharing

When a notebook .Rmd is saved, an .nb.html file is created alongside it. This file is a self-contained HTML file which contains both a rendered copy of the notebook with all current chunk outputs (suitable for display on a website) and a copy of the notebook .Rmd itself.

More info can be found at [RStudio docs](#)

# Chapter 124: Aggregating data frames

Aggregation is one of the most common uses for R. There are several ways to do so in R, which we will illustrate here.

## Section 124.1: Aggregating with data.table

Grouping with the data.table package is done using the syntax `dt[i, j, by]` Which can be read out loud as: "Take *dt*, subset rows using *i*, then calculate *j*, grouped by *by*." Within the *dt* statement, multiple calculations or groups should be put in a list. Since an alias for `list()` is `.()`, both can be used interchangeably. In the examples below we use `.()`.

### CODE:

```
Aggregating with data.table
library(data.table)

dt = data.table(group=c("Group 1", "Group 1", "Group 2", "Group 2", "Group 2"),
 subgroup = c("A", "A", "A", "A", "B"),
 value = c(2, 2.5, 1, 2, 1.5))
print(dt)

sum, grouping by one column
dt[, .(value=sum(value)), group]

mean, grouping by one column
dt[, .(value=mean(value)), group]

sum, grouping by multiple columns
dt[, .(value=sum(value)), .(group, subgroup)]

custom function, grouping by one column
in this example we want the sum of all values larger than 2 per group.
dt[, .(value=sum(value[value>2])), group]
```

### OUTPUT:

```
> # Aggregating with data.table
> library(data.table)
>
> dt = data.table(group=c("Group 1", "Group 1", "Group 2", "Group 2", "Group 2"),
+ subgroup = c("A", "A", "A", "A", "B"),
+ value = c(2, 2.5, 1, 2, 1.5))
> print(dt)
 group subgroup value
1: Group 1 A 2.0
2: Group 1 A 2.5
3: Group 2 A 1.0
4: Group 2 A 2.0
5: Group 2 B 1.5
>
> # sum, grouping by one column
> dt[, .(value=sum(value)), group]
 group value
1: Group 1 4.5
2: Group 2 4.5
>
> # mean, grouping by one column
> dt[, .(value=mean(value)), group]
 group value
```

```

1: Group 1 2.25
2: Group 2 1.50
>
> # sum, grouping by multiple columns
> dt[,.(value=sum(value)),.(group,subgroup)]
 group subgroup value
1: Group 1 A 4.5
2: Group 2 A 3.0
3: Group 2 B 1.5
>
> # custom function, grouping by one column
> # in this example we want the sum of all values larger than 2 per group.
> dt[,.(value=sum(value[value>2])),group]
 group value
1: Group 1 2.5
2: Group 2 0.0

```

## Section 124.2: Aggregating with base R

For this, we will use the function `aggregate`, which can be used as follows:

```
aggregate(formula, function, data)
```

The following code shows various ways of using the `aggregate` function.

**CODE:**

```

df = data.frame(group=c("Group 1", "Group 1", "Group 2", "Group 2", "Group 2"),
 subgroup = c("A", "A", "A", "A", "B"), value = c(2, 2.5, 1, 2, 1.5))

sum, grouping by one column
aggregate(value~group, FUN=sum, data=df)

mean, grouping by one column
aggregate(value~group, FUN=mean, data=df)

sum, grouping by multiple columns
aggregate(value~group+subgroup, FUN=sum, data=df)

custom function, grouping by one column
in this example we want the sum of all values larger than 2 per group.
aggregate(value~group, FUN=function(x) sum(x[x>2]), data=df)

```

**OUTPUT:**

```

> df = data.frame(group=c("Group 1", "Group 1", "Group 2", "Group 2", "Group 2"),
 subgroup = c("A", "A", "A", "A", "B"), value = c(2, 2.5, 1, 2, 1.5))
> print(df)
 group subgroup value
1 Group 1 A 2.0
2 Group 1 A 2.5
3 Group 2 A 1.0
4 Group 2 A 2.0
5 Group 2 B 1.5
>
> # sum, grouping by one column
> aggregate(value~group, FUN=sum, data=df)
 group value
1 Group 1 4.5

```

```

2 Group 2 4.5
>
> # mean, grouping by one column
> aggregate(value~group, FUN=mean, data=df)
 group value
1 Group 1 2.25
2 Group 2 1.50
>
> # sum, grouping by multiple columns
> aggregate(value~group+subgroup,FUN=sum,data=df)
 group subgroup value
1 Group 1 A 4.5
2 Group 2 A 3.0
3 Group 2 B 1.5
>
> # custom function, grouping by one column
> # in this example we want the sum of all values larger than 2 per group.
> aggregate(value~group, FUN=function(x) sum(x[x>2]), data=df)
 group value
1 Group 1 2.5
2 Group 2 0.0

```

## Section 124.3: Aggregating with dplyr

Aggregating with dplyr is easy! You can use the `group_by()` and the `summarize()` functions for this. Some examples are given below.

### CODE:

```

Aggregating with dplyr
library(dplyr)

df = data.frame(group=c("Group 1","Group 1","Group 2","Group 2","Group 2"),
 subgroup = c("A","A","A","A","B"),
 value = c(2,2.5,1,2,1.5))
print(df)

sum, grouping by one column
df %>% group_by(group) %>% summarize(value = sum(value)) %>% as.data.frame()

mean, grouping by one column
df %>% group_by(group) %>% summarize(value = mean(value)) %>% as.data.frame()

sum, grouping by multiple columns
df %>% group_by(group,subgroup) %>% summarize(value = sum(value)) %>% as.data.frame()

custom function, grouping by one column
in this example we want the sum of all values larger than 2 per group.
df %>% group_by(group) %>% summarize(value = sum(value[value>2])) %>% as.data.frame()

```

### OUTPUT:

```

> library(dplyr)
>
> df = data.frame(group=c("Group 1","Group 1","Group 2","Group 2","Group 2"),
> subgroup = c("A","A","A","A","B"),
> value = c(2,2.5,1,2,1.5))
> print(df)
 group subgroup value
1 Group 1 A 2.0
2 Group 1 A 2.5
3 Group 2 A 1.0

```

```

4 Group 2 A 2.0
5 Group 2 B 1.5
>
> # sum, grouping by one column
> df %>% group_by(group) %>% summarize(value = sum(value)) %>% as.data.frame()
 group value
1 Group 1 4.5
2 Group 2 4.5
>
> # mean, grouping by one column
> df %>% group_by(group) %>% summarize(value = mean(value)) %>% as.data.frame()
 group value
1 Group 1 2.25
2 Group 2 1.50
>
> # sum, grouping by multiple columns
> df %>% group_by(group,subgroup) %>% summarize(value = sum(value)) %>% as.data.frame()
 group subgroup value
1 Group 1 A 4.5
2 Group 2 A 3.0
3 Group 2 B 1.5
>
> # custom function, grouping by one column
> # in this example we want the sum of all values larger than 2 per group.
> df %>% group_by(group) %>% summarize(value = sum(value[value>2])) %>% as.data.frame()
 group value
1 Group 1 2.5
2 Group 2 0.0

```

# Chapter 125: Data acquisition

Get data directly into an R session. One of the nice features of R is the ease of data acquisition. There are several ways data dissemination using R packages.

## Section 125.1: Built-in datasets

R has a vast collection of built-in datasets. Usually, they are used for teaching purposes to create quick and easily reproducible examples. There is a nice web-page listing the built-in datasets:

<https://vincentarelbundock.github.io/Rdatasets/datasets.html>

### Example

Swiss Fertility and Socioeconomic Indicators (1888) Data. Let's check the difference in fertility based on rurality and domination of Catholic population.

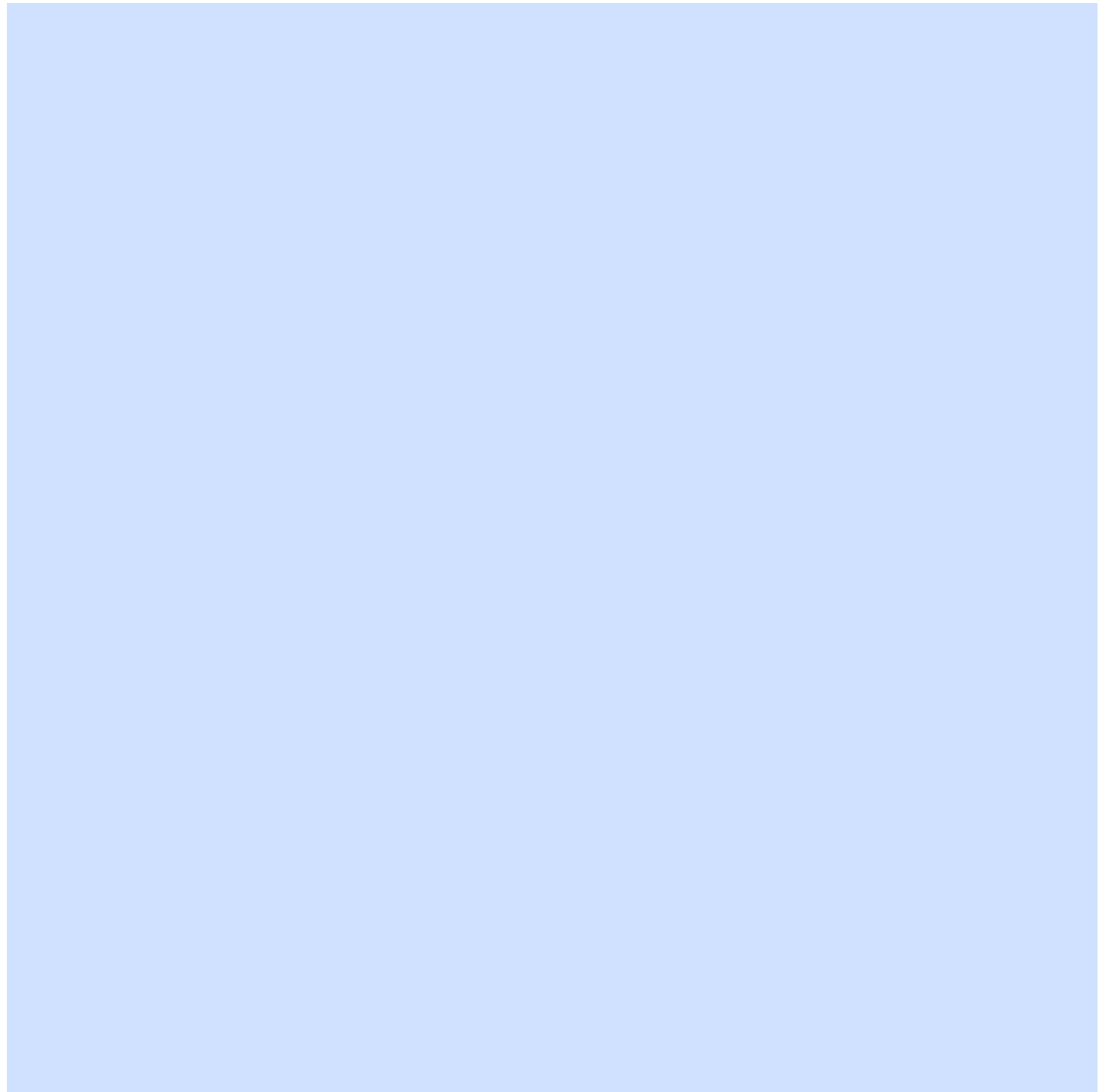
```
library(tidyverse)

swiss %>%
 ggplot(aes(x = Agriculture, y = Fertility,
 color = Catholic > 50)) +
 geom_point() +
 stat_ellipse()
```

## Section 125.2: Packages to access open databases

Numerous packages are created specifically to access some databases. Using them can save a bunch of time on





## Section 125.3: Packages to access restricted data

### Human Mortality Database

[Human Mortality Database](#) is a project of the [Max Planck Institute for Demographic Research](#) that gathers and pre-process human mortality data for those countries, where more or less reliable statistics is available.

```
load required packages
library(tidyverse)
library(extrafont)
library(HMDHDFplus)

country <- getHMDcountries()

exposures <- list()
for (i in 1: length(country)) {
```







## Section 125.4: Datasets within packages

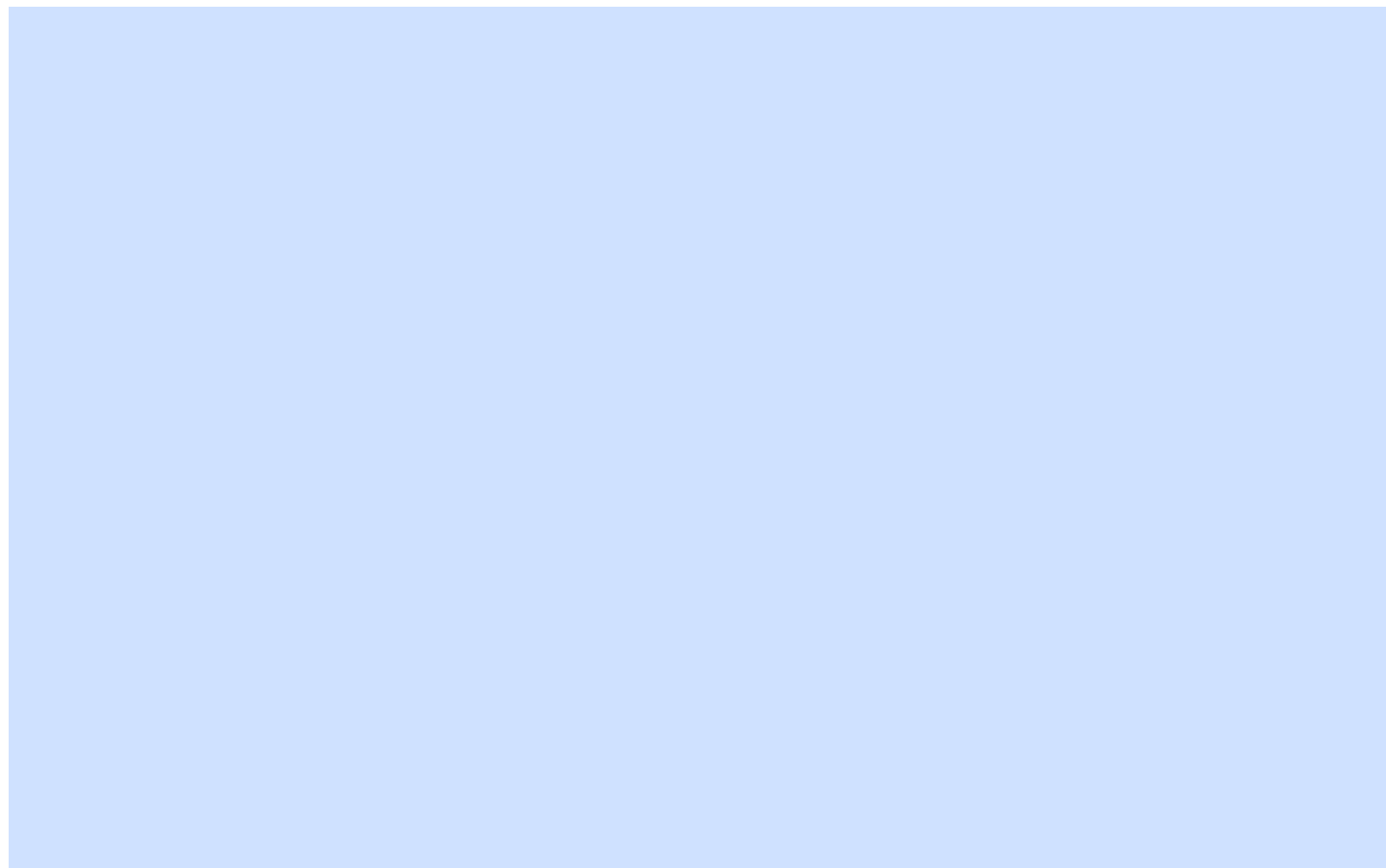
There are packages that include data or are created specifically to disseminate datasets. When such a package is loaded (`library(pkg)`), the attached datasets become available either as R objects; or they need to be called with the `data()` function.

### Gapminder

A nice dataset on the development of countries.

```
library(tidyverse)
library(gapminder)

gapminder %>%
 ggplot(aes(x = year, y = lifeExp,
 color = continent)) +
 geom_jitter(size = 1, alpha = .2, width = .75) +
 stat_summary(geom = "path", fun.y = mean, size = 1) +
 theme_minimal()
```



### World Population Prospects 2015 - United Nations Population Department

Let's see how the world has converged in male life expectancy at birth over 1950-2015.

```
library(tidyverse)
library(forcats)
library(wpp2015)
library(ggjoy)
library(viridis)
```

```

library(extrafont)

data(UNlocations)

countries <- UNlocations %>%
 filter(location_type == 4) %>%
 transmute(name = name %>% paste()) %>%
 as_vector()

data(e0M)

e0M %>%
 filter(country %in% countries) %>%
 select(-last.observed) %>%
 gather(period, value, 3:15) %>%
 ggplot(aes(x = value, y = period %>% fct_rev()))+
 geom_joy(aes(fill = period))+
 scale_fill_viridis(discrete = T, option = "B", direction = -1,
 begin = .1, end = .9)+
 labs(x = "Male life expectancy at birth",
 y = "Period",
 title = "The world convergence in male life expectancy at birth since 1950",
 subtitle = "Data: UNPD World Population Prospects 2015 Revision",
 caption = "ikashnitsky.github.io")+
 theme_minimal(base_family = "Roboto Condensed", base_size = 15)+
 theme(legend.position = "none")

```

# Chapter 126: R memento by examples

This topic is meant to be a memento about the R language without any text, with self-explanatory examples.

Each example is meant to be as succinct as possible.

## Section 126.1: Plotting (using plot)

```
Creates a 1 row - 2 columns format
par(mfrow=c(1,2))

plot(rnorm(100), main = "Graph 1", ylab = "Normal distribution")
grid()
legend(x = 40, y = -1, legend = "A legend")

plot(rnorm(100), main = "Graph 2", type = "l")
abline(v = 50)
```

Result:



## Section 126.2: Commonly used functions

```
Create 100 standard normals in a vector
x <- rnorm(100, mean = 0, sd = 1)

Find the lenght of a vector
length(x)

Compute the mean
mean(x)

Compute the standard deviation
sd(x)

Compute the median value
median(x)
```

```

Compute the range (min, max)
range(x)

Sum an iterable
sum(x)

Cumulative sum (x[1], x[1]+x[2], ...)
cumsum(x)

Display the first 3 elements
head(3, x)

Display min, 1st quartile, median, mean, 3rd quartile, max
summary(x)

Compute successive difference between elements
diff(x)

Create a range from 1 to 10 step 1
1:10

Create a range from 1 to 10 step 0.1
seq(1, 10, 0.1)

Print a string
print("hello world")

```

## Section 126.3: Data types

### Vectors

```

a <- c(1, 2, 3)
b <- c(4, 5, 6)
mean_ab <- (a + b) / 2

d <- c(1, 0, 1)
only_1_3 <- a[d == 1]

```

### Matrices

```

mat <- matrix(c(1,2,3,4), nrow = 2, ncol = 2)
dimnames(mat) <- list(c(), c("a", "b", "c"))
mat[,] == mat

```

### Dataframes

```

df <- data.frame(qualifiers = c("Buy", "Sell", "Sell"),
 symbols = c("AAPL", "MSFT", "GOOGL"),
 values = c(326.0, 598.3, 201.5))
df$symbols == df[[2]]
df$symbols == df[["symbols"]]
df[[2, 1]] == "AAPL"

```

### Lists

```

l <- list(a = 500, "aaa", 98.2)
length(l) == 3
class(l[1]) == "list"
class(l[[1]]) == "numeric"
class(l$a) == "numeric"

```

### Environments

```

env <- new.env()
env[["foo"]] = "bar"
env2 <- env

```

```
env2[["foo"]] = "BAR"

env[["foo"]] == "BAR"
get("foo", envir = env) == "BAR"
rm("foo", envir = env)
env[["foo"]] == NULL
```

# Chapter 127: Updating R version

Installing or Updating your Software will give access to new features and bug fixes. Updating your R installation can be done in a couple of ways. One Simple way is go to [R website](https://cran.r-project.org/) and download the latest version for your system.

## Section 127.1: Installing from R Website

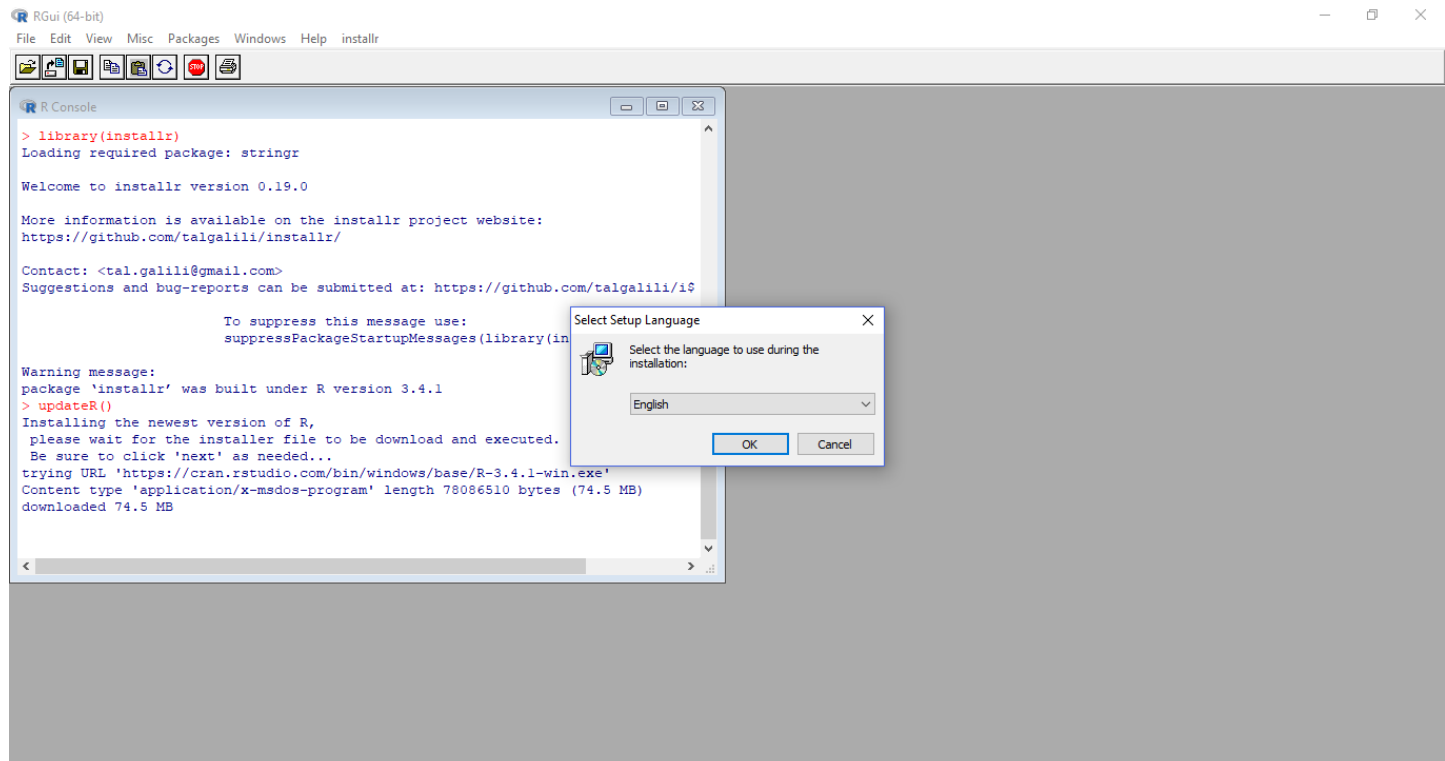
To get the latest release go to <https://cran.r-project.org/> and download the file for your operating system. Open the downloaded file and follow the on-screen installation steps. All the settings can be left on default unless you want to change a certain behaviour.

## Section 127.2: Updating from within R using installr Package

You can also update R from within R by using a handy package called **installr**.

Open R Console (NOT RStudio, this doesn't work from RStudio) and run the following code to install the package and initiate update.

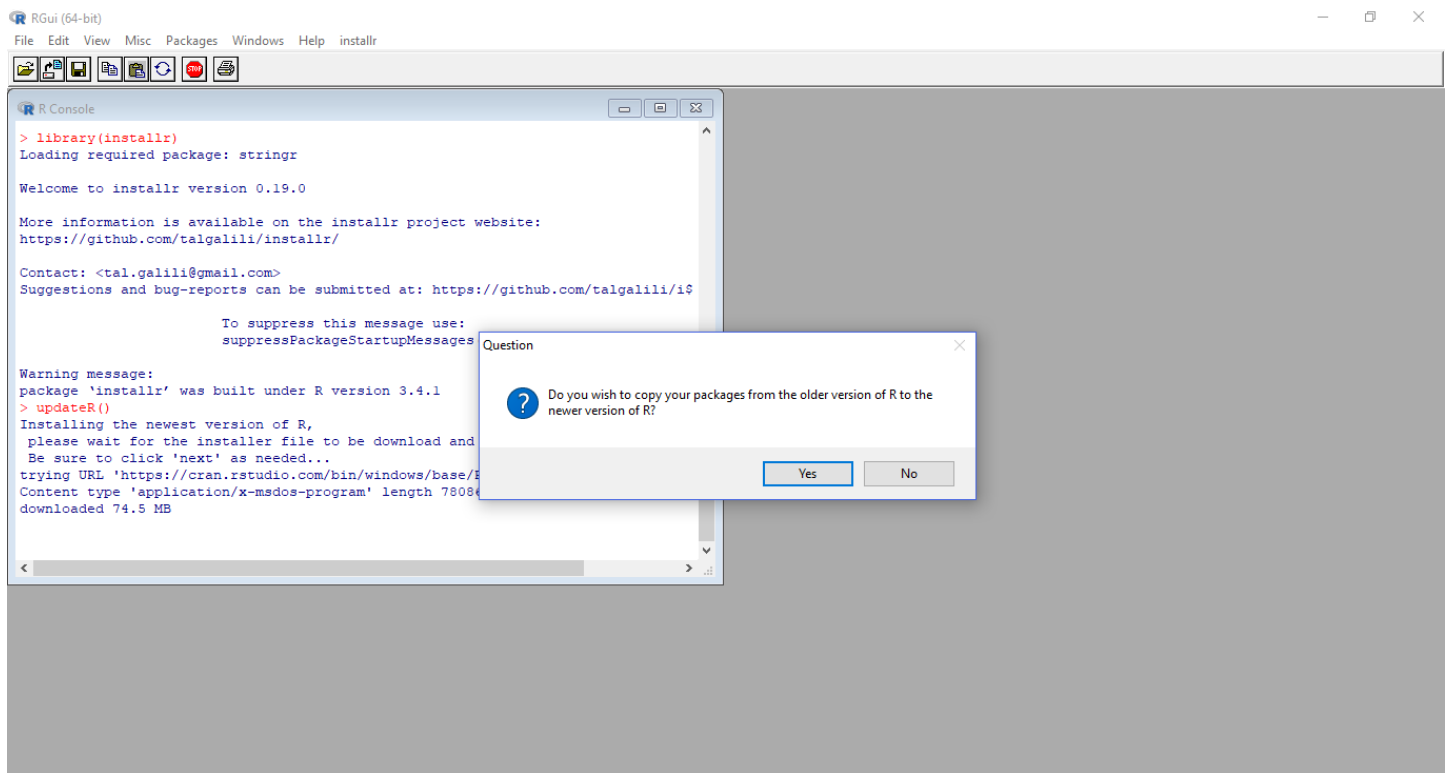
```
install.packages("installr")
library("installr")
updateR()
```



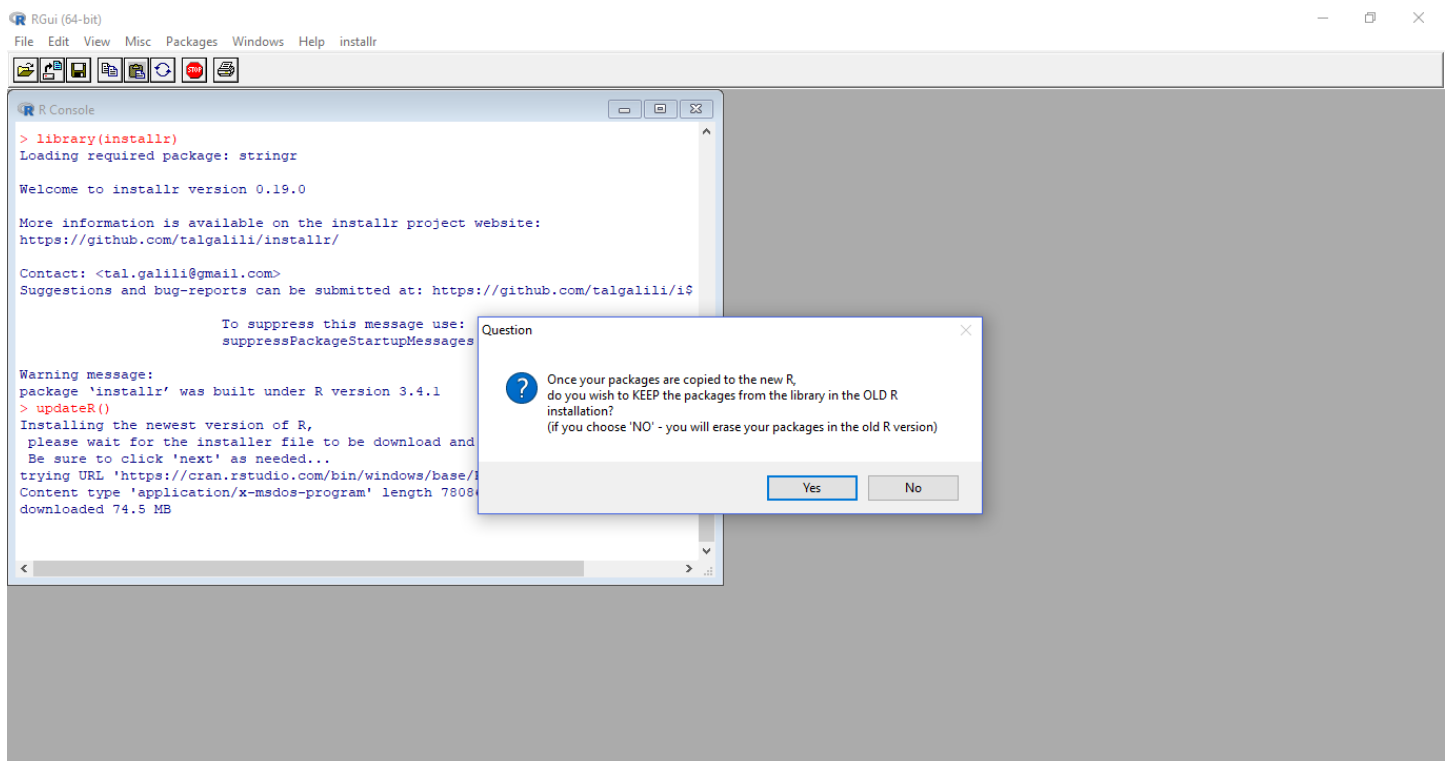
## Section 127.3: Deciding on the old packages

Once the installation is finished click the Finish button.

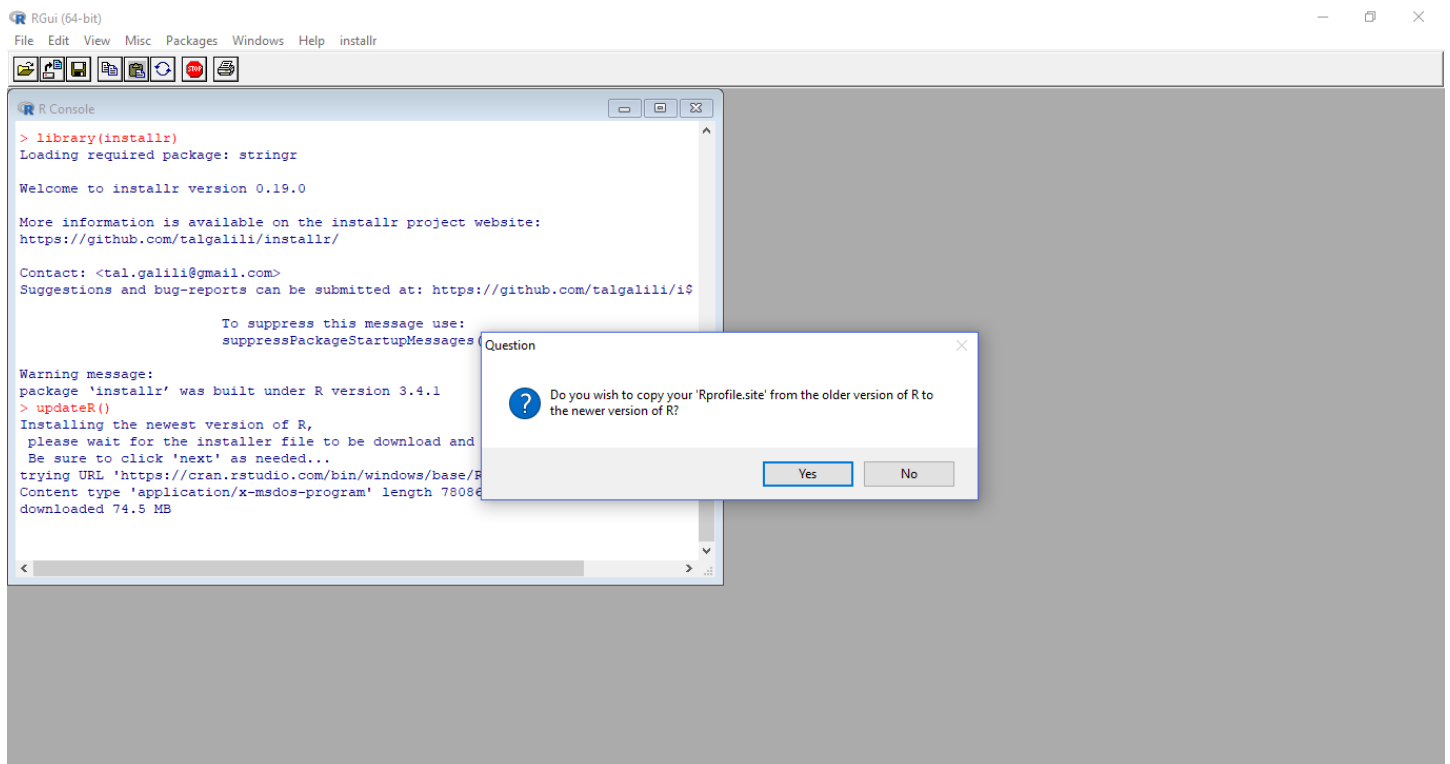
Now it asks if you want to copy your packages fro the older version of R to Newer version of R. Once you choose yes all the package are copied to the newer version of R.



After that you can choose if you still want to keep the old packages or delete.

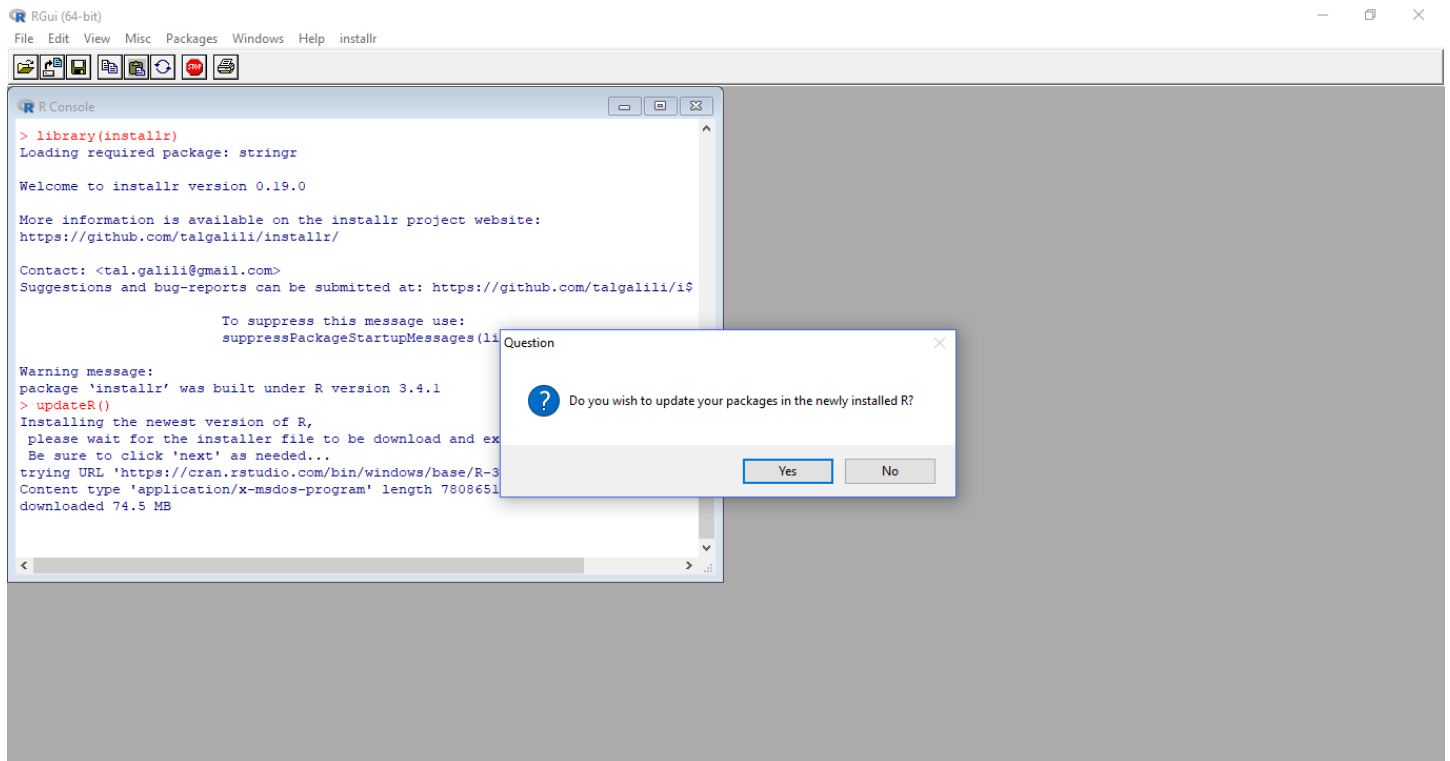


You can even move your Rprofile.site from older version to keep all your customised settings.



## Section 127.4: Updating Packages

You can update your installed packages once the updating of R is done.



Once its done Restart R and enjoy exploring.

## Section 127.5: Check R Version

You can check R Version using the console

```
version
```

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