



Computergestützte Statistik: Programmieren mit R

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#### Introduction

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package: graphics package: ggplot2

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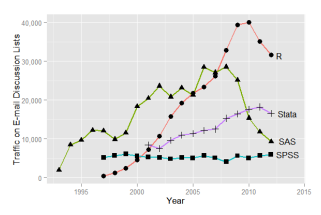
## That's why:

- ► Powerful statistical software
- It allows 'Statistical Programming'
- 'Statistical Programming' allows for automation
- R is the easiest language to speak badly<sup>1</sup>
- ► A vital user community
- Share code for reproducible research
- ► It's free!

<sup>&</sup>lt;sup>1</sup>http://www.r-bloggers.com/r-is-the-easiest-language-to-speak-badly/



## A vital user community:



http://r4stats.com/articles/popularity/



## Write better Code with Fewer Lines.

- Hadley Wickham -

better code: Reproducibility and performance

fewer lines: easy to maintain and share,

no 'copy & paste'



Most of what you will see on the slides, you will find at:

► Course: Roger Peng, Computing for Data Analysis, John Hopkins University, available at https://github.com/DataScienceSpecialization

#### And also in:

- Chamber, John M. (2008): Software for Data Analysis: Programming with R, Springer
- ▶ Matloff, Norman (2012): The Art of R Programming, no starch press
- Wickham, Hadley (2014): Advanced R, CRC Press, available at http://adv-r.had.co.nz/



## General information:

- ▶ http://cran.r-project.org/ and Task Views:
- ► R-bloggers: http://www.r-bloggers.com/

## Getting help:

- ► Rseek: www.rseek.org
- stackoverflow: http://stackoverflow.com/



- ► Fundamentals:
  - Data types
  - Subsetting
  - Missing Values
  - ► Statistics in R
- Graphics
- Data Handling
  - Reading and writing data
  - Data Frames
  - Computing on Data Frames
  - String Manipulation
- Programming
  - Functions
  - Scoping
  - Control Structures
  - \*apply
  - Debugging
  - Object-Orientation (S3)

- Do you need previous knowledge? nope
- What you have to do:
  - Do the exercises! Seriously!
  - ► 5-informal + 3-formal (in groups)
  - In preparation to the next class, try a gentle introductory tutorial: https://www.datacamp.com/courses/introduction-to-r
  - Pass the acceptance test!
- ► How you will be graded: 15% formal-exercises, 85% term paper (10-20 pages)
- ► Find a homework-group. Not more than 2-3 students. Send the names and 'Matr.Nr.' to: Soeren.Pannier@fu-berlin.de before the
- If you don't know anybody in this class, we will find a group for you send us an e-mail!
- Laptops vs. computer lab

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- R can be downloaded from one of the mirror sites in http://cran.r-project.org/mirrors.html.
   You should pick your nearest location.
- "Windows and Mac users most likely want to download the precompiled binaries. Since R is part of many Linux distributions, you should check with your Linux package management system."
- Download R for Linux, Mac or Windows.
- ▶ Install R with the standard settings, since later on we will work with RStudio (IDE).

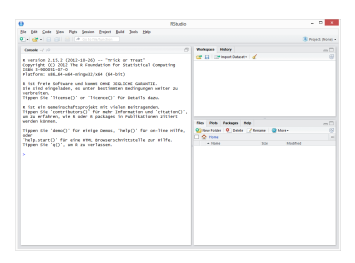
## Installation: RStudio



- RStudio can be downloaded from http://www.rstudio.com/ide/download/.
- ▶ Download the Desktop Version for Linux, Mac or Windows.
- ► Install RStudio. (Make sure you already installed R.)
- ► Start RStudio.

#### Installation: RStudio





Your screen should look like this.

- An R package includes a set of functions and datasets which is not included in the 'base' R System.
- Packages provide additional functionality.
- An exhaustive list of available packages is on CRAN (about 6500): http://cran.r-project.org/web/packages/
- ► There are also many packages associated with the Bioconductor project http://bioconductor.org.



- ▶ Install a new package, e.g.ggplot2: install.packages("ggplot2")
- Sometimes you need to specify more options. For instance, this is the case if you are not an administrator of the computer.
  - lib specifies the directory where you want to store the package
  - repos specifies a list of repositories (CRAN mirrors)
  - dep=T specifies that all the required packages are also downloaded and installed
- Stay up to date: update.packages("ggplot2")
- ► Load packages so that you are able to use them: library("ggplot2")
- Unload packages: detach("package:ggplot2")

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## **Fundamentals**

## Data types

## Atomic data types



R has six basic or "atomic" data types:

- character
- numeric (real numbers)
- integer
- complex
- logical (TRUE/FALSE)
- raw

The most basic object is a vector

- a vector can only contain objects of the same type
- Exception: A list, which is represented as a vector, can contain objects of different classes.

- ▶ Numbers in R are generally treated as numeric objects (i.e. double precision real numbers)
- ▶ If you explicitly want an integer, you need to specify the L suffix
- Example: Entering 1 gives you a numeric object; entering 1L explicitly gives you an integer.
- ▶ There is also a special number Inf which represents infinity; e.g.1/0; Inf can be used in ordinary calculations; e.g. 1/Inf is 0
- ► The value NaN represents an undefined value ("not a number"); e.g. 0/0; NaN can also be thought of as a missing value (more on that later)



## **Entering Input**

x < -1

At the R prompt we type expressions. The <- symbol is the assignment operator.

```
print(x)
## [1] 1
х
## [1] 1
msg <- "hello"
```

The grammar of the language determines whether an expression is complete or not.

```
# Some R code
#x <- ## Incomplete expression
```

The # character indicates a comment. Anything to the right of the # (including the # itself) is ignored.

When a complete expression is entered at the prompt, it is evaluated and the result of the evaluated expression is returned. The result may be auto-printed.

```
# Some R code
x <- 5 ## nothing printed, but Object x is created
       ## auto-printing occurs
х
```

## [1] 5

```
print(x) ## explicit printing
```

## [1] 5

The [1] indicates that x is a vector and 5 is the first element.

```
x <- 1:50
x
```

## [1] 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 ## [24] 24 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40 41 42 43 44 45

## [47] 47 48 49 50

The : operator is used to create integer sequences.

```
length(1:100)
```

## [1] 100

The c() function can be used to create vectors of objects.

```
x <- c(0.5, 0.6) ## numeric
x <- c(TRUE, FALSE) ## logical
x \leftarrow c(T, F)
                ## logical, but harder to read
x <- c("a", "b", "c") ## character
x < -9:29
                     ## integer
x <- c(1+0i, 2+4i) ## complex
```

#### Using the vector() function

```
x <- vector("numeric", length = 10)</pre>
x
```

```
##
    [1] 0 0 0 0 0 0 0 0 0 0
```

## Mixing Objects



## What about the following?

```
c(1.7, "a")  ## character
c(TRUE, 2)  ## numeric
c("a", TRUE)  ## character
c("a", TRUE, 1)  ## character
c("a", c(TRUE, 1))  ## character
```

When different objects are mixed in a vector, *coercion* occurs so that every element in the vector is of the same class.

Objects can be explicitly coerced from one class to another using the as.\* functions, if available.

```
x <- 0:6 class(x)
```

## [1] "integer"

```
as.numeric(x)
```

## [1] 0 1 2 3 4 5 6

```
as.logical(x)
```

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

```
as.character(x)
```

```
## [1] "0" "1" "2" "3" "4" "5" "6"
```

as.complex(x)

Nonsensical coercion results in NAs.

```
x <- c("a", "b", "c")
as.numeric(x)
```

## Warning: NAs introduced by coercion ## [1] NA NA NA

```
as.logical(x)
```

## [1] NA NA NA

#### **Factors**



Factors are used to represent categorical data. Factors can be *unordered* or *ordered*. Internally factor are stored as "labeled" integer vectors.

- ► Factors are treated specially by modelling functions like lm() and glm()
- ▶ Using factors with labels is better than using integers because factors are self-describing; having a variable that has values "Male" and "Female" is better than a variable that has values 1 and 2.

```
x <- factor(c("yes", "yes", "no", "yes", "no"))</pre>
х
## [1] yes yes no yes no
## Levels: no yes
table(x)
## x
## no yes
## 2 3
unclass(x)
## [1] 2 2 1 2 1
## attr(,"levels")
## [1] "no" "yes"
```

#### **Factors**



The order of the levels can be set using the levels argument to factor(). This can be important in linear modelling because the first level is used as the baseline level.

```
x <- factor(c("yes", "yes", "no", "yes", "no"),
           levels = c("yes", "no"))
х
## [1] yes yes no yes no
## Levels: yes no
table(x)
## x
## yes no
##
unclass(x)
## [1] 1 1 2 1 2
## attr(,"levels")
## [1] "yes" "no"
```



The first levels can be set using relevel. Levels can be relabeled and added.

```
## exam
## success failure
## 3 2
```

#### Data Frames



Data frames are used to store tabular data

- ► They are represented as a **special type of list** where every element of the list has to have the same length
- ► Each element of the list can be thought of as a column and the length of each element of the list is the number of rows
- ▶ data frames can store different classes of objects in each column

```
x <- data.frame(eggs = 1:4,
              ham = c(TRUE, TRUE, FALSE, FALSE))
х
##
    eggs ham
        1 TRUE
## 2 2 TRUE
## 3 3 FALSE
## 4 4 FALSE
nrow(x)
## [1] 4
ncol(x)
## [1] 2
dim(x)
## [1] 4 2
```

## Constructing Data Frames



- ▶ str() is helpful to check the variable classes and dimensions of a data frame.
- Character vectors are coerced to factors by default.

```
## $ num : int 1 2 3 4
## $ char : Factor w/ 1 level "a": 1 1 1 1
## $ logic: logi TRUE FALSE TRUE FALSE
```

## Constructing Data Frames



Data frames can be constructed by combining data objects. The elements to be combined should have the same number of rows. If not, R will try to repeat elements.

```
a <- 1:4
b <- c("a", "b")
c <- c(TRUE, FALSE, TRUE, FALSE)
x <- data.frame(num = a, "char" = b, c)
str(x)</pre>
```

```
## 'data.frame': 4 obs. of 3 variables:
## $ num : int 1 2 3 4
## $ char: Factor w/ 2 levels "a","b": 1 2 1 2
## $ c : logi TRUE FALSE TRUE FALSE
```

Lists

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- ▶ Lists are a special type of vector that can contain elements of different data types.
- ▶ They are different from data.frames because the elements can differ in length, so that the structure of the data can not be thought of as being tabular.
- Elements of a list can be lists or data frames.

```
x <- list(1, "a", TRUE)
x

## [[1]]
## [1] 1
##
## [[2]]
## [1] "a"
##
## [[3]]
## [1] TRUE</pre>
```

- Lists can also have names.
- Lists can also be nested and used to represent complex data structures, although this feature should only be used when absolutely necessary:

```
x <- list(a = 1, b = list(1, 2), c = data.frame(x = 1:2))
str(x)
## List of 3</pre>
```

```
## List of 3
## $ a: num 1
## $ b:List of 2
## ..$ : num 1
## ..$ : num 2
## $ c:'data.frame': 2 obs. of 1 variable:
## ..$ x: int [1:2] 1 2
```

## The NULL Object



There is a special object called NULL. It

- ... represents the null object in R
- ... is an object with defined neutral ("null") behavior.
- ... has no type and no modifiable properties
- ... should not be confused with a vector or list of zero length
- ... is a reserved word
- ... is often returned by expressions and functions whose value is undefined

To test for NULL use is.null.

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#### **Fundamentals**

Subsetting

## Subsetting



There are a number of operators that can be used to extract subsets of  ${\sf R}$  objects.

- ▶ [ ] always returns an object of the same class as the original (there is one exception); can be used to select more than one element
- [[]] is used to extract elements of a list or a data frame; it can only be used to extract a single element and the class of the returned object will not necessarily be a list or data frame
- \$ is used to extract elements of a list or data frame by name; semantics are similar to that of [[ ]].

```
x <- c("a", "b", "c", "c", "d", "a")
x[1]
## [1] "a"
x[2]
## [1] "b"
x[1:4]
## [1] "a" "b" "c" "c"
x[x > "a"]
## [1] "b" "c" "c" "d"
u <- x > "a"
u
## [1] FALSE TRUE TRUE TRUE TRUE FALSE
```

x[u]

## [1] "b" "c" "c" "d"

# Subsetting Data Frames



# Subsetting data frames with names:

# Subsetting Data Frames



# Subsetting data frames with positions:

```
x <- data.frame(eggs = 1:4,
                ham = c(TRUE, TRUE, FALSE, FALSE))
x[[1]] #class(x[[1]]) is integer
## [1] 1 2 3 4
x[1] #class(x[1]) is data.frame
##
     eggs
## 4
x[c(1,3), 2]
## [1]
        TRUE FALSE
```

# Subsetting Data Frames



Subsetting data frames with logicals:

```
x <- data.frame(eggs = 1:4,
              ham = c(TRUE, TRUE, FALSE, FALSE))
x[x$eggs > 2, ]
##
            ham
     eggs
        3 FALSE
## 3
## 4 4 FALSE
x[x$ham,]
##
    eggs ham
        1 TRUE
## 2
        2 TRUE
x[c(TRUE, FALSE)]
```

```
##
     eggs
## 1
## 4
```

# Deleting elements of Lists/Data Frames



To delete elements in a list you can simply assign the value NULL to them.

```
x[1] <- NULL
x
## ham
## 1 TRUE
```

## 2 TRUE ## 3 FALSE ## 4 FALSE

For a subset of rows select the rows to keep or use data.frame[-<rows>, ]:

```
x <- x[-(1:2), ]
x
```

## [1] FALSE FALSE

## Missing Values



Missing values are denoted by NA or NaN for undefined mathematical operations.

- ▶ is.na() is used to test objects if they are NA
- ▶ is.nan() is used to test for NaN
- ▶ NA values have a class also, so there are integer NA, character NA, etc.
- ▶ A NaN value is also NA but the converse is not true

## Removing NA values



A common task is to remove missing values (NA).

## [1] FALSE FALSE TRUE FALSE TRUE FALSE

x[!bad]

## [1] 1 2 4 5

# Removing NA values



# airquality[1:6, ]

```
Ozone Solar.R Wind Temp Month Day
##
## 1
       41
              190 7.4
                         67
       36
              118 8.0
                       72
## 2
                                    3
## 3
       12
              149 12.6 74
       18
              313 11.5
                         62
                                    4
## 4
## 5
       NA
               NA 14.3
                         56
                                    5
## 6
       28
               NA 14.9
                         66
                                5
                                    6
```

# na.omit(airquality)[1:6, ]

| ## |   | Ozone | ${\tt Solar.R}$ | ${\tt Wind}$ | Temp | ${\tt Month}$ | Day |
|----|---|-------|-----------------|--------------|------|---------------|-----|
| ## | 1 | 41    | 190             | 7.4          | 67   | 5             | 1   |
| ## | 2 | 36    | 118             | 8.0          | 72   | 5             | 2   |
| ## | 3 | 12    | 149             | 12.6         | 74   | 5             | 3   |
| ## | 4 | 18    | 313             | 11.5         | 62   | 5             | 4   |
| ## | 7 | 23    | 299             | 8.6          | 65   | 5             | 7   |
| ## | 8 | 19    | 99              | 13.8         | 59   | 5             | 8   |



- Semantics for subsetting a list is equivalent to data frames because they are closely related.
- ► One exception: A list is not meant to represent tabular data, hence subsetting rows/columns is not meaningfull: list[<rows>, <cols>] won't work!
- ▶ Lists can be recursive (elements of a list can be lists). The subsetting for nested elements works *just* like for a normal list:

## [1] 14

▶ The [[ ]] can also take an integer sequence:

```
x[[c(1, 3)]]
```

## [1] 14

```
# not to be confused with: x[c(1, 3)] - lists are vectors!
```

# Partial Matching



Partial matching of names is allowed (but not recommended) with [[ ]] and \$.

```
x \leftarrow list(ham = 1:5, cheese = pi)
x$c
## [1] 3.141593
x[["c"]]
## NULL
x[["c", exact = FALSE]]
## [1] 3.141593
x$h
## [1] 1 2 3 4 5
```

x <- c(x, hohoho="Hohoho") x\$h

## NULL

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# **Vectorized Operations**



Many operations in R are vectorized making code more effcient, concise, and easier to read.

```
x < -1:4; y < -6:9
x + y
```

$$x >= 2$$

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#### Operators

R contains a number of operators. They are listed in the table below.

- +,- Plus, Minus, unary or binary
  - ! NOT, unary
  - : Sequence, binary (in model formulae: interaction)
  - \*, / Multiplication, Division, binary
    - ^ Exponentiation, binary
- %% Modulus, binary
- %/% Integer divide, binary
- % \* % Matrix product, binary
- %in% Matching operator, binary (in model formulae: nesting)
- <,<= Less than, Less than or equal, binary
- >,>= Greater than, Greater than or equal, binary
  - == Equal to, binary
- &, && AND (vectorized, not vectorized), binary
  - |, || OR (vectorized, not vectorized), binary

Further Details: R Language Definition 3.1.4



- &, | The shorter form performs elementwise comparisons in much the same way as arithmetic operators
- &&,|| The longer form evaluates left to right examining only the first element of each vector. Evaluation proceeds only until the result is determined. The longer form is appropriate for programming control-flow and typically preferred in if clauses.

```
c(TRUE, TRUE) & c(TRUE, FALSE)

## [1] TRUE FALSE

c(TRUE, TRUE) && c(TRUE, FALSE)

## [1] TRUE

FALSE && NULL
```

## [1] FALSE

FALSE && NULL

## [1] FALSE

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## R Style Rules



The goal of Style Rules is to make R code easier to read, share, and verify. There is no unique standard. $^2$ 

- 1. File names: end in .r (and have a meaningful name)
- 2. **Identifiers:** should be meaningful; R is case sensitive

```
variables: lowerCamelCase (dataFrame, someData)
```

functions: lowerCamelCase (someFunction, functionName)

constants: lowerCamelCase (i, j, meanOfSomething)

- 3. Line Length: maximum 80 characters
- 4. **Spacing:** Place spaces around all binary operators (=, +, -, <-, etc.).
- 5. Curly Braces: first on same line, last on own line
- 6. Assignment: use <-, not =
- 7. Semicolons: don't use them
- 8. Commenting Guidelines: First, do use comments. All comments begin with # followed by a space; inline comments need two spaces before the #

<sup>&</sup>lt;sup>2</sup>http://journal.r-project.org/archive/2012-2/RJournal\_2012-2\_Baaaath.pdf

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## Graphics

package: graphics
package: ggplot2
package: shiny

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The plotting and graphics engine in R is encapsulated in a few base and recommended packages:

- graphics: contains plotting functions for the 'base' graphing systems, including plot, hist, boxplot and many others.
- ggplot2: introduces a clean syntax for customized graphics using the 'grammar of graphics'. A lot of default settings for legends and colours.
- ▶ lattice: there's a nice introductive tutorial: http://dsarkar.fhcrc.org/lattice-lab/latticeIntro.pdf
- ggvis: interactive graphics based on the 'grammar of graphics'
- ► shiny: interactive applications

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## Basic graphics



- ▶ Base graphics are usually constructed piecewise, with each aspect of the plot handled separately through a series of function calls.
- ► This plotting approach mirrors the thought process.
- Plotting commands are divided into two basic groups:
  - High-level plotting functions
     create a new plot on the graphics device (more on that later), possibly with axes,
     labels, titles and so on.
  - Low-level plotting functions add more information to an existing plot, such as extra points, lines, labels, etc.
- In addition, R maintains a list of graphical parameters par() which can be manipulated to customize your plots.



## High-level plotting functions

- generate a complete plot of the data passed as arguments to the function. Where appropriate, axes, labels and titles are automatically generated (unless you request otherwise).
- (usually) start a new plot, erasing the current plot if necessary.

```
plot() ... generic function; output depends
pairs() ... matrix of scatterplots
smoothScatter() ... color density representation of a 2D-scatterplot
boxplot ... box-and-whisker plot(s) of (grouped) values
barplot() ... dto
qqplot() ... dto
hist() ... histogramm
...
```

Further Functions: An Introduction to R 12.1.3

# The plot() function is a generic function



generic functions: result depends on class or type of the first argument.

- ▶ plot(x):
   if x is a time series → time-series plot
   if x is a numeric vector → plot of the values in the vector against index
   if x is a complex vector → plot of imaginary vs real parts of the vector elements
   if x is a a factor → barplot
- ▶ plot(x, y): if x and y are numeric vectors → scatter plot if x is a factor and y is numeric → grouped boxplot
- ▶ plot(xy): if xy is either a list containing two elements x and y or xy is a two-column matrix → scatter plot / box plot
- ▶ plot(df): if df is a data frame → matrix of scatterplots
- ▶ plot(fx): if fx is a function → a curve corresponding to fx ist plotted
- **.** . . .

## Arguments to high-level plotting functions



add = TRUEsuperimposes plot on current plot (forces the function to act as a low-level graphics function) suppresses the generation of axis (you can customize the axes axes = FALSE

later with axis) log = "x", "y"

log = "xy"Causes the x, y or both axes to be logarithmic.

The type= argument controls the type of plot as follows:

Plot individual points (default) type="p"

Plot lines type="1"

type="b", "o" Plot points connected/overlaid by lines type="h" Plot vertical lines from points to x axis

type="s", "S" Step function plots

type="n" no plotting, axes are still drawn

xlab = string

type=

Axis labels for the x and y axes vlab = string

Figure title, placed at the top of the plot in a large font. main = string sub = string

Sub-title, placed just below the x-axis in a smaller font.

## Low-level plotting functions



Low-level plotting commands can be used to add extra information (such as points, lines or text) to the current plot.

```
points(x, y)
lines(x, y)
                    Adds points or connected lines to the current plot.
text(x, y, ...)
                    Add text to a plot at points given by x, y.
abline(a, b)
abline(h = v)
abline(v = x)
abline(lm.obj)
                    Adds a line of slope b and intercept a to the current plot. h and
                    v add horizontal and vertical lines. The result of a linear modell
                    lm.obj with coefficients a and b may be assigned as well.
legend(...)
                    Adds a legend to the current plot.
title(...)
                    Adds a title main to the top of the current plot.
```

Adds an axis to the current plot on specified the side.

axis(side, ...)

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# Graphics

package: graphics package: ggplot2 package: shiny

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# The elements of the grammar (ggplot2)



- Data and aesthetic mappings:
  - Every plot is based on a data frame
  - ► Aesthetic mappings: data is mapped to aesthetic attributes
- ► Geoms are the graphical representation of a statistic you want to visualize: points, lines, polygons, etc.
- Stats are the statistics you want to calculate for the data. A scatter plot simply uses the identity, a bar plot often uses relative frequencies. A statistic is always associated with a geom.
- Scales map values in the data space to values in the aesthetic space. They define the shape, size or colour and are used to create legends and axes
- ► The coordinate system (coord), typically the Cartesian coordinate system
- Faceting is used to split the data into subsets and then to reproduce the graphic on each subset (see extra slides on graphics for an example)
- ► Theme defines fonts of all labels, the shape of ticks and background. Use the function labs to add labels and a title

## The elements of the grammar



- ▶ Unlike most functions in R there is a naming convention in the package.
  - geom\_<geom-name> to add geoms with statistic
  - stat\_<stat-name> to add a statistic with geom
  - scale\_<aesthetic-name>\_<scale-type> to manipulate scales
  - coord\_<coord-type> to change the coordinate system
  - facet\_<facet-type> to add facets
  - ▶ theme to manipulate text and line elements (which are not geoms)
- This naming conventions mirrors the grammar and is the reason why you need to understand it
- The grammar does not define how a graphic should look like. What is appropriate and inappropriate is up to you
- ▶ ggplot2 only produces static graphics, no movement, no interaction, no 3D
- ▶ The function qplot mimics the syntactical usage of plot

```
str(ggplot)
```

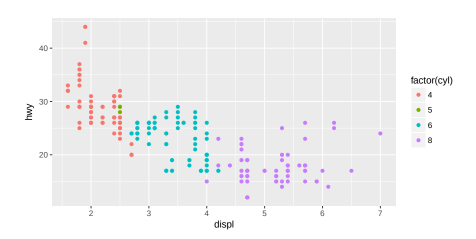
## function (data = NULL, mapping = aes(), ..., environment = parent.fram

- ▶ The fundamental function in ggplot2 is ggplot
- ggplot is the fundament of every plot created by ggplot2, any value specified in ggplot is used as default for any layer. Actually it is simply passed using . . .
- ▶ A layer is the representation of the grammar elements which define the plot
- ▶ A plot can be created with more than one layer. Actually it is created layer-by-layer
- Every component of a plot aesthetics, geoms, stats, scales, etc. can be added using the binary operator '+'



- ▶ Scales map values in the data space to values in the aesthetic space
- aes() simply connects data values (variables) to aesthetic attributes, like x, y, fill, alpha, etc.
- Scales calculate the positions in the Cartesian coordinate system represented by numbers between 0 and 1
- Scales will pick colours and automatically generate legends for aesthetics like 'colour'
- Every scale can be manipulated by using scale\_'aesthetic-name'\_'scale-type'()
- Scales are picked by default according to the class of the data: Numeric data will have a continuous scale, a factor usually uses a discrete scale
  - if 'x' is numeric: scale\_x\_continuous()
  - if 'colour' is a factor: scale\_colour\_discrete()

```
data(mpg)
ggplot(mpg, aes(displ, hwy, colour = factor(cyl))) +
  geom_point()
```



# Example: Mapping and scales



## ...the data:

| manufacturer | model | displ | hwy | cty |
|--------------|-------|-------|-----|-----|
| audi         | a4    | 1.8   | 29  | 4   |
| audi         | a4    | 1.8   | 29  | 4   |
| audi         | a4    | 2.0   | 31  | 4   |
| audi         | a4    | 2.0   | 30  | 4   |
| audi         | a4    | 2.8   | 26  | 6   |
| audi         | a4    | 2.8   | 26  | 6   |
| audi         | a4    | 3.1   | 27  | 6   |
| audi         | a4    | 1.8   | 26  | 4   |
| audi         | a4    | 1.8   | 25  | 4   |
| audi         | a4    | 2.0   | 28  | 4   |

## Example: Mapping and scales



... connection between aesthetic attribute and data:

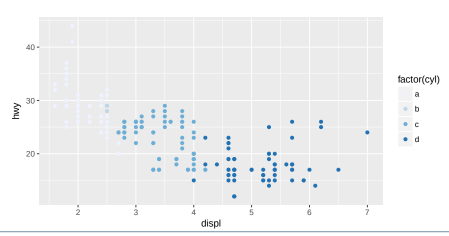
| x(displ) | $\mathbf{y}(hwy)$ | colour(cty |
|----------|-------------------|------------|
| 1.8      | 29                | 4          |
| 1.8      | 29                | 4          |
| 2.0      | 31                | 4          |
| 2.0      | 30                | 4          |
| 2.8      | 26                | 6          |
| 2.8      | 26                | 6          |
| 3.1      | 27                | 6          |
| 1.8      | 26                | 4          |
| 1.8      | 25                | 4          |
| 2.0      | 28                | 4          |
|          |                   |            |



# ... map scales:

| x     | у     | colour  | size | shape |
|-------|-------|---------|------|-------|
| 0.037 | 0.531 | #FF6C91 | 1    | 19    |
| 0.037 | 0.531 | #FF6C91 | 1    | 19    |
| 0.074 | 0.594 | #FF6C91 | 1    | 19    |
| 0.074 | 0.562 | #FF6C91 | 1    | 19    |
| 0.222 | 0.438 | #00C1A9 | 1    | 19    |
| 0.222 | 0.438 | #00C1A9 | 1    | 19    |
| 0.278 | 0.469 | #00C1A9 | 1    | 19    |
| 0.037 | 0.438 | #FF6C91 | 1    | 19    |
| 0.037 | 0.406 | #FF6C91 | 1    | 19    |
| 0.074 | 0.500 | #FF6C91 | 1    | 19    |

```
data(mpg)
ggplot(mpg, aes(displ, hwy, colour = factor(cyl))) +
  geom_point() +
  scale_colour_brewer(palette = 1, labels = letters[1:4])
```



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package: graphic package: ggplot2 package: shiny

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## Shiny



- shiny is a project from RStudio
- ► A framework to create interactive web applications (websites)
  - to do predefined analysis (reports)
  - create dashboards
  - visualize results with interactive elements
- ▶ It uses a lot JavaScript and HTML in the background. Knowledge of these languages can be beneficial but you do not need them for this course



First, let's see an example!



- ► Shiny applications know two things: the UI (User-Interface) and the server
- ▶ UI (shinyUI)
  - Defines the layout of the application (bootstrapPage, fluidPage, ...)
  - ▶ Defines interactive elements (numericInput, selectInput)
  - ▶ ...
- ► Server (shinyServer)
  - ► A server is a function you will understand this in time. . .
  - ▶ Defines output elements (renderPlot, renderTable, ...)
  - Also processes input elements
  - ▶ ...
- The server and user-interface can communicate with each other by accessing the objects input and output

# Inputs



When you define an interactive element in the UI an entry is added to the list of inputs

```
sliderInput("adjust", "Select binwidth", 0.1, 5, value = 2)
```

- ▶ Creates the element adjust as element of input. The default value is 2, the range from which values can be selected is between 0.1 and 5
- Inputs can then be used from the server

```
ggplot(airquality, aes(Ozone)) + geom_density(adjust = input$adjust)
```

▶ What can be confusing is that there is no explicit assignment, like input\$adjust <- ..., the manipulation of input is a side-effect of sliderInput

#### Outputs



▶ Outputs are created in the server by explicit modification of output

```
output$plot <- renderPlot({
    ggplot(airquality, aes(Ozone)) +
        geom_density(adjust = input$adjust)
})</pre>
```

- Typically elements in output are objects created by a function called render<Something>
- ▶ Just because we defined an element in output doesn't mean it is visualized by the UI. To do that we need to define where the plot is displayed by calling:

```
plotOutput("plot")
```

▶ This checks output for an element named *plot* and displays it in the UI



- ► Homepage: http://shiny.rstudio.com/
- ► Gallery: http://shiny.rstudio.com/gallery/
- Extensions:
  - http://hrbrmstr.github.io/metricsgraphics
  - http://rstudio.github.io/dygraphs
    - ٠..

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## Data Handling

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There are a few principal functions reading data into R:

- read.table, read.csv, for reading tabular data
- readLines, for reading lines of a text file
- ▶ load, for reading in saved workspaces (\*.Rdata files)



There are analogous functions for writing data to files:

- ▶ write.table
- ▶ writeLines
- save



The read.table function is one of the most commonly used functions for reading data. It has a few important arguments:

- ▶ file, the name of a file, or a connection
- header, logical indicating if the file has a header line
- sep, a string indicating how the columns are seperated
- colClasses, a character vector indicating the class of each column in the dataset
- nrows, the number of rows in the dataset
- comment.char, a character string indicating the comment character
- skip, the number of lines to skip from the beginning
- stringsAsFactors, logical indicating if character variables should be coded as factors



```
x <- read.table("Data/someData.txt")</pre>
str(x)
```

```
## 'data.frame': 100 obs. of 2 variables:
##
    $ someNumbers: num -0.626 0.184 -0.836 1.595 0.33 ...
                 : Factor w/ 2 levels "a", "b": 1 1 1 1 1 1 1 1 1 1 ...
##
    $ aFactor
R will automatically
```

- skip lines that begin with a #
- detect the number of rows
- assign a class to each variable of the table

You can speed up R by defining these things directly - this is very useful for large datasets. read.csv is identical to read.table except that the default seperator is a comma.

## Define the class for each variable manually:



The readLines function can be used to simply read lines of a text file and store them in a character vector.



# readLines can be useful for reading in lines of webpages

```
x <- readLines("http://fu-berlin.de/")
length(x)
## [1] 45</pre>
```

#### x[1:7]

- ## [1] "<!DOCTYPE html><!-- BEGIN Fragment fu-berlin/17091808/default/all</pre>
- ## [2] "<html class=\"ltr\" lang=\"de\"><head><title>Freie UniversitÃ $\Omega$ t H
- ## [3] "<meta charset=\"utf-8\" /><meta content=\"IE=edge\" http-equiv=\"</pre>
- ## [5] "<!-- END Fragment fu-berlin/17091808/views/open\_graph\_image/46016
- ## [6] "<meta content=\"Die Freie UniversitÃ $\alpha$ t zÃ $\alpha$ hlt zu den elf deutsche
- ## [7] "<link href=\"/assets/fu-berlin/favicon-a6b103813c732ebbbff3dd77fc</pre>

# Reading and Writing "Foreign" Data



R is able to read and write from different data sources although not by default. The following packages can be helpful:

- ▶ foreign, Haven read data stored by Minitab, S, SAS, SPSS, Stata, ...
- ▶ RODBC, sqldf or RMySQL, for SQL databases
- ► SPARQL for semantic web
- RCurl for more communication with web resources twitteR as an example for communicating with APIs

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# When working with data you must:

- ► Figure out what you want to do.
- Precisely describe what you want in the form of a computer program.
- ► Execute the code.
- The dplyr package makes each of these steps as fast and easy as (currently) possible.

In the following, we will have a look at the package's (shortened) introductory vignette.

dplyr aims to provide a function for each basic verb of data manipulating:

- filter() (and slice())
- arrange()
- select() (and rename())
- ▶ distinct()
- mutate() (and transmute())
- ▶ summarise()
- sample\_n() and sample\_frac()



```
library(dplyr)
library(nycflights13)
flights
# A tibble: 336.776 x 19
  year month day dep_time sched_dep_time dep_delay arr_time
 <int> <int> <int> <int>
                                    <int>
                                              <dbl>
                                                      <int>
1 2013
                        517
                                      515
                                                        830
 2013
                        533
                                      529
                                                        850
 2013 1
                       542
                                      540
                                                        923
4 2013 1
                       544
                                      545
                                                       1004
 ... with 336,772 more rows, and 12 more variables: sched_arr_time <int>,
   arr_delay <dbl>, carrier <chr>, flight <int>, tailnum <chr>,
   origin <chr>, dest <chr>, air_time <dbl>, distance <dbl>, hour <dbl>,
   minute <dbl>, time_hour <dttm>
```

Data description availabe via ?flights.

```
class(flights)
[1] "tbl df"
                  "t.b1"
                                "data.frame"
```

dplyr can work with data frames as is, but if you're dealing with large data, it's worthwhile to convert them to a tbl\_df: this is a wrapper around a data frame that won't accidentally print a lot of data to the screen.

For example, we can select all flights on July 1st with:

```
filter(flights, month == 7, day == 1)
# A tibble: 966 x 19
  year month day dep_time sched_dep_time dep_delay arr_time
 <int> <int> <int> <int>
                                           <dbl> <int>
                                  <int>
1 2013
                                   2029
                                             212
                                                     236
2 2013 7 1
                                                     344
                                   2359
3 2013 7 1
                       29
                                   2245
                                             104 151
4 2013 7
                       43
                                   2130
                                             193
                                                     322
# ... with 962 more rows. and 12 more variables: sched arr time <int>.
   arr_delay <dbl>, carrier <chr>, flight <int>, tailnum <chr>,
   origin <chr>, dest <chr>, air_time <dbl>, distance <dbl>, hour <dbl>,
   minute <dbl>. time hour <dttm>
```

This is equivalent to the more verbose:

```
flights[flights$month == 7 & flights$day == 1, ]
```



For example, we can select all flights that started early and arrived late:

```
filter(flights, dep_delay < 0 & arr_delay > 0 )
# A tibble: 35.648 x 19
  year month day dep_time sched_dep_time dep_delay arr_time
 <int> <int> <int> <int>
                                 <int>
                                           <dbl>
                                                   <int>
1 2013 1
                      554
                                   558
                                                     740
2 2013 1 1
                      555
                                                    913
                                   600
                                          -5
3 2013 1 1
                      558
                                   600
                                            -2
                                                    753
4 2013 1
                      558
                                   600
                                        -2
                                                     924
 ... with 35,644 more rows, and 12 more variables: sched_arr_time <int>,
   arr_delay <dbl>, carrier <chr>, flight <int>, tailnum <chr>,
# origin <chr>, dest <chr>, air_time <dbl>, distance <dbl>, hour <dbl>,
   minute <dbl>, time_hour <dttm>
```



To select rows by position, use slice():

```
slice(flights, 3:2)
# A tibble: 2 x 19
  year month day dep_time sched_dep_time dep_delay arr_time
 <int> <int> <int> <int>
                                     <int>
                                              <dbl>
                                                       <int>
 2013
                        542
                                       540
                                                          923
2 2013
                        533
                                       529
                                                          850
 ... with 12 more variables: sched_arr_time <int>, arr_delay <dbl>,
   carrier <chr>, flight <int>, tailnum <chr>, origin <chr>, dest <chr>,
   air_time <dbl>, distance <dbl>, hour <dbl>, minute <dbl>,
    time_hour <dttm>
```

# Arrange (reorder) rows with arrange()



```
arrange(flights, year, month, day)
# A tibble: 336,776 x 19
  year month day dep_time sched_dep_time dep_delay arr_time
 <int> <int> <int> <int>
                                     <int>
                                              <dbl>
                                                       <int>
                        517
                                      515
                                                         830
  2013
  2013 1
                        533
                                      529
                                                         850
3 2013 1
                        542
                                      540
                                                         923
  2013
                        544
                                      545
                                                        1004
 ... with 336,772 more rows, and 12 more variables: sched_arr_time <int>,
   arr_delay <dbl>, carrier <chr>, flight <int>, tailnum <chr>,
   origin <chr>, dest <chr>, air_time <dbl>, distance <dbl>, hour <dbl>,
   minute <dbl>, time_hour <dttm>
```



Use desc() to order a column in descending order:

```
arrange(flights, desc(arr_delay))
# A tibble: 336,776 x 19
  year month day dep_time sched_dep_time dep_delay arr_time
 <int> <int> <int>
                                           <dbl>
                    <int>
                                  <int>
                                                   <int>
  2013
                      641
                                    900
                                            1301
                                                    1242
                9
2 2013 6 15 1432
                                   1935
                                           1137 1607
 2013 1 10 1121
                                   1635 1126 1239
  2013
                     1139
                                   1845 1014 1457
               20
 ... with 336.772 more rows, and 12 more variables: sched arr time <int>.
   arr_delay <dbl>, carrier <chr>, flight <int>, tailnum <chr>,
   origin <chr>, dest <chr>, air_time <dbl>, distance <dbl>, hour <dbl>,
   minute <dbl>, time hour <dttm>
```

```
# Select columns by name
select(flights, year, month, day)
# A tibble: 336,776 x 3
    year month day
    <int> <int> <int>
1 2013 1 1
2 2013 1 1
3 2013 1 1
4 2013 1 1
# ... with 336,772 more rows
```



```
# Select all columns between year and day (inclusive)
select(flights, year:day)
# A tibble: 336.776 x 3
  year month
              day
 <int> <int> <int>
1 2013
2 2013
3 2013
4 2013
# ... with 336.772 more rows
# Select all columns except those from year to arr_time (inclusive)
select(flights, -(year:arr_time))
# A tibble: 336.776 x 12
 sched_arr_time arr_delay carrier flight tailnum origin dest air_time
                  <dbl> <chr> <int> <chr> <chr> <chr>
          <int>
                                                             <dbl>
            819
                                 1545
                                       N14228
                                                 EWR.
                                                       IAH
                                                               227
                      11
                             UA
2
            830
                      20
                             UA 1714 N24211 LGA
                                                      IAH
                                                               227
3
            850
                      33
                             AA 1141
                                       N619AA JFK
                                                      MIA
                                                               160
           1022
                     -18
                             B6 725 N804JB JFK
                                                       BQN
                                                               183
 ... with 336,772 more rows, and 4 more variables: distance <dbl>,
   hour <dbl>, minute <dbl>, time_hour <dttm>
```



There are a number of helper functions you can use within select(), like

- starts\_with()
- ends\_with()
- ▶ matches()
- contains().

These let you quickly match larger blocks of variable that meet some criterion. See ?select for more details.

You can rename variables with select() by using named arguments:

But because select() drops all the variables not explicitly mentioned, it's not that useful.



# Instead, use rename():

```
rename(flights, depTime = dep_time)
# A tibble: 336.776 x 19
  year month day depTime sched_dep_time dep_delay arr_time
 <int> <int> <int> <int>
                                <int>
                                        <dbl>
                                                 <int>
1 2013 1 1 517
                                  515
                                                   830
2 2013 1 1 533
                                  529
                                                   850
3 2013 1 1
                  542
                                  540
                                                  923
4 2013 1
              1 544
                                  545
                                                  1004
 ... with 336,772 more rows, and 12 more variables: sched_arr_time <int>,
   arr_delay <dbl>, carrier <chr>, flight <int>, tailnum <chr>,
# origin <chr>, dest <chr>, air_time <dbl>, distance <dbl>, hour <dbl>,
   minute <dbl>, time_hour <dttm>
```



A common use of select() is to find out which values a set of variables takes. In conjunction with distinct() only the unique values are returned.

```
distinct(select(flights, tailnum))
# A tibble: 4,044 x 1
 tailnum
   <chr>
1 N14228
2 N24211
3 N619AA
4 N804.IB
# ... with 4,040 more rows
distinct(select(flights, origin, dest))
# A tibble: 224 x 2
 origin dest
  <chr> <chr>
    EWR. IAH
 LGA IAH
3
  JFK MIA
    JFK
          BQN
 ... with 220 more rows
```

This is very similar to base::unique() but should be much faster.



As well as selecting from the set of existing columns, it's often useful to add new columns that are functions of existing columns. This is the job of the mutate() verb.

```
mutate(flights,
  gain = arr_delay - dep_delay,
  speed = distance / air_time * 60)
# A tibble: 336,776 x 21
  year month day dep_time sched_dep_time dep_delay arr_time
  <int> <int> <int> <int>
                                    <int>
                                              <dbl>
                                                       <int>
  2013
                                      515
                                                         830
                        517
2 2013 1
                        533
                                      529
                                                         850
3 2013 1
                        542
                                      540
                                                         923
  2013
                        544
                                      545
                                                       1004
# ... with 336,772 more rows, and 14 more variables: sched_arr_time <int>,
   arr_delay <dbl>, carrier <chr>, flight <int>, tailnum <chr>,
    origin <chr>, dest <chr>, air_time <dbl>, distance <dbl>, hour <dbl>,
   minute <dbl>, time_hour <dttm>, gain <dbl>, speed <dbl>
```



mutate() allows you to refer to columns that you just created:

```
mutate(flights,
 gain = arr_delay - dep_delay,
 gain_per_hour = gain / (air_time / 60)
# A tibble: 336.776 x 21
  year month day dep_time sched_dep_time dep_delay arr_time
 <int> <int> <int> <int>
                                   <int>
                                            <dbl>
                                                     <int>
 2013 1
                       517
                                     515
                                                       830
2 2013 1 1
                       533
                                     529
                                                       850
3 2013 1 1
                       542
                                     540
                                                       923
4 2013
                       544
                                     545
                                                      1004
 ... with 336.772 more rows, and 14 more variables: sched arr time <int>.
   arr_delay <dbl>, carrier <chr>, flight <int>, tailnum <chr>,
   origin <chr>, dest <chr>, air_time <dbl>, distance <dbl>, hour <dbl>,
   minute <dbl>, time_hour <dttm>, qain <dbl>, qain_per_hour <dbl>
```



If you only want to keep the new variables, use transmute():



The last verb is summarise(), which collapses a data frame to a single row. It's not very useful yet:

```
summarise(flights,
  delay = mean(dep_delay, na.rm = TRUE))
# A tibble: 1 x 1
    delay
    <dbl>
1 12.63907
```

You can use  $sample_n()$  and  $sample_frac()$  to take a random sample of rows

- ▶ a fixed number via sample\_n()
- ▶ a fixed fraction via sample\_frac().

```
sample_n(flights, 10)
# A tibble: 10 x 19
  year month day dep_time sched_dep_time dep_delay arr_time
 <int> <int> <int>
                    <int>
                                 <int>
                                          <dbl>
                                                  <int>
 2013
               21
                 1316
                                  1320
                                                 1630
 2013 11 8
                 2035
                                  2040
                                                   2250
 2013 9 10
                    1859
                                  1905
                                             -6 2127
  2013
              17 1629
                                  1636
                                             -7
                                                   1807
 ... with 6 more rows, and 12 more variables: sched arr time <int>.
   arr_delay <dbl>, carrier <chr>, flight <int>, tailnum <chr>,
   origin <chr>, dest <chr>, air_time <dbl>, distance <dbl>, hour <dbl>,
   minute <dbl>. time hour <dttm>
```



```
sample_frac(flights, 0.01)
# A tibble: 3,368 x 19
  year month day dep_time sched_dep_time dep_delay arr_time
 <int> <int> <int>
                    <int>
                                 <int>
                                          <dbl>
                                                  <int>
  2013
                                  1125
                                             -6 1302
       5 25 1119
 2013 2 21 1719
                                  1700
                                             19 2012
 2013 12 11 1050
                                  1100
                                            -10 1147
  2013 12
               26
                      650
                                   655
                                            -5
                                                    800
# ... with 3,364 more rows, and 12 more variables: sched_arr_time <int>,
   arr_delay <dbl>, carrier <chr>, flight <int>, tailnum <chr>,
   origin <chr>, dest <chr>, air_time <dbl>, distance <dbl>, hour <dbl>,
   minute <dbl>. time hour <dttm>
```

Use replace = TRUE to perform a bootstrap sample, and optionally weight the sample with the weight argument.

You may have noticed that all these functions are very similar:

- ► The first argument is a data frame.
- ► The subsequent arguments describe what to do with it, and you can refer to columns in the data frame directly without using \$.
- ▶ The result is a new data frame

Together these properties make it easy to chain together multiple simple steps to achieve a complex result.

These five functions provide the basis of a language of data manipulation.

At the most basic level, you can only alter a data frame in five useful ways:

- you can reorder the rows (arrange()),
- pick observations (filter() ) and
- variables of interest (select()),
- 4. add new variables that are functions of existing variables (mutate()) or
- 5. collapse many values to a summary (summarise()).

The remainder of the language comes from applying the five functions to different types of data, like to **grouped data**, as described next.

- ► The introduced verbs are useful, but they become really powerful when you combine them with the idea of "group by", repeating the operation individually on groups of observations within the dataset.
- ▶ In dplyr, you use the group\_by() function to describe how to break a dataset down into groups of rows. You can then use the resulting object in exactly the same functions as above; they'll automatically work "by group" when the input is a grouped.

# The verbs are affected by grouping as follows:

- grouped select() is the same as ungrouped select(), excepted that retains grouping variables are always retained.
- grouped arrange() orders first by grouping variables
- mutate() and filter() are most useful in conjunction with window functions (like rank(), or min(x) == x), and are described in detail in vignette("window-function").
- sample\_n() and sample\_frac() sample the specified number/fraction of rows in each group.
- slice() extracts rows within each group.
- summarise() is easy to understand and very useful, and is described in more detail below.

In the following example, we split the complete dataset into individual planes and then summarise each plane by counting the number of flights (count = n()) and computing the average distance (dist = mean(Distance, na.rm = TRUE)) and delay (delay = mean(ArrDelay, na.rm = TRUE)). We then use ggplot2 to display the output.

```
by_tailnum <- group_by(flights, tailnum)</pre>
delay <- summarise(by_tailnum,
 count = n(),
 dist = mean(distance, na.rm = TRUE),
 delay = mean(arr_delay, na.rm = TRUE))
delay <- filter(delay, count > 20, dist < 2000)
delay
# A tibble: 2,962 x 4
 tailnum count dist
                         delay
   <chr> <int> <dbl>
                            <dbl>
1 NOEGMQ 371 676.1887 9.982955
2 N10156 153 757.9477 12.717241
3 N102UW 48 535.8750 2.937500
4 N103US 46 535.1957 -6.934783
# ... with 2,958 more rows
```



You use summarise() with aggregate functions, which take a vector of values, and return a single number. There are many useful functions in base R like min(), max(), mean(), sum(), sd(), median(), and IQR(). dplyr provides a handful of others:

- n(): number of observations in the current group
- ▶ n\_distinct(x): count the number of unique values in x.
- first(x), last(x) and nth(x, n) these work similarly to x[1], x[length(x)], and x[n] but give you more control of the result if the value isn't present.

For example, we could use these to find the number of planes and the number of flights that go to each possible destination:

```
destinations <- group_by(flights, dest)</pre>
summarise (destinations,
 planes = n_distinct(tailnum),
 flights = n()
 A tibble: 105 x 3
  dest planes flights
 <chr> <int>
                 <int>
   ABQ
          108
                  254
   ACK
        58
               265
   AT.B
        172
               439
   ANC
  ... with 101 more rows
```

The dplyr API is functional in the sense that function calls don't have side-effects, and you must always save their results. This doesn't lead to particularly elegant code if you want to do many operations at once. You either have to do it step-by-step:

```
a1 <- group_by(flights, year, month, day)
a2 <- select(a1, arr_delay, dep_delay)
a3 <- summarise(a2,
    arr = mean(arr_delay, na.rm = TRUE),
    dep = mean(dep_delay, na.rm = TRUE))
a4 <- filter(a3, arr > 30 | dep > 30)
```

### Chaining



Or if you don't want to save the intermediate results, you need to wrap the function calls inside each other:

```
filter(
  summarise(
   select(
     group_by(flights, year, month, day),
     arr_delay, dep_delay
    ).
   arr = mean(arr_delay, na.rm = TRUE),
   dep = mean(dep_delay, na.rm = TRUE)
  ),
  arr > 30 \mid dep > 30
 A tibble: 49 x 5
# Groups: year, month [11]
  year month
               day
                       arr
                              dep
  <int> <int> <int> <dbl>
                              <dbl>
 2013 1 16 34.24736 24.61287
2 2013 1 31 32.60285 28.65836
 2013 2 11 36.29009 39.07360
  2013 2 27 31.25249 37.76327
# ... with 45 more rows
```

This is difficult to read because the order of the operations is from inside to out, and the arguments are a long way away from the function. To get around this problem, dplyr provides the %>% operator. x %>% f(y) turns into f(x, y) so you can use it to rewrite multiple operations so you can read from left-to-right, top-to-bottom:

```
flights %>%
 group_by(year, month, day) %>%
 select(arr_delay, dep_delay) %>%
 summarise(
   arr = mean(arr_delay, na.rm = TRUE),
   dep = mean(dep_delay, na.rm = TRUE)
 ) %>%
 filter(arr > 30 | dep > 30)
# A tibble: 49 x 5
# Groups: year, month [11]
  year month
              day
                       arr
                               dep
 <int> <int> <int> <dbl>
                              <dbl>
 2013 1 16 34.24736 24.61287
2 2013 1 31 32.60285 28.65836
3 2013 2 11 36.29009 39.07360
 2013 2 27 31,25249 37,76327
# ... with 45 more rows
```

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# Data Handling

Reading and writing data Computing on Data Frames (with dplyr)

String Manipulation

Programming

# Strings and Characters in Statistical Analysis



Read in data often must be "cleaned" and explicitly coerced to the class of data, that  ${\sf R}$  can deal with.

- Specification of missing values ("99", "NA", ".")
- ▶ Different formats in the same column: ("0.5", "0,5", "1/2")
- ▶ Dates: "2013-05-03 10:14:15", "03.05.2013 10:14:15"
- Contamination of data
- Getting information out of text, for example in text mining, HTML-code and other data represented in textual format



There are some fundamental functions for manipulating strings and character vectors in  $\mathsf{R}$ :

- lacktriangledown paste(), converts all arguments to character and concatenates them to one vector
- strsplit(), splits a string into a list of substring
- grep(), search for pattern in string
- and many related functions

Additional topic: Regular expressions

 ${\tt paste()}\ converts\ any\ arguments\ to\ a\ character\ and\ concatenates\ them:$ 

```
x <- "Some"
paste(x, "String")
## [1] "Some String"
y <- paste(x, "String")</pre>
## [1] "Some String"
z <- c("Some", "More")</pre>
a <- paste(z, "String", sep = "_")
a
## [1] "Some_String" "More_String"
```

strsplit splits a string into several substring:

"String"

```
strsplit(y, split = " ")

## [[1]]
## [1] "Some" "String"

b <- strsplit(a, split = "_")
b

## [[1]]
## [[1] "Some" "String"
##
## [[2]]</pre>
```

## [1] "More"



There are several string constants which can be used with escape sequences. Escape sequences are introduced using a backslash.

```
\" double quote
\n new line
\t tab
\\ backslash
```

see more in the R Language Definition

This is why you have to specify "path/someFile.R" or "path\someFile.R" and not "path\someFile.R".

#### escape sequences



Escape sequences can be stored in character vectors, they will be "evaluated" at the time when R needs to interprete a character - for example when defining file names or paths.

```
newLine <- "new\nline"
print(newLine)
## [1] "new\nline"
cat(newLine)
## new
## line
moreNewLines <- paste(rep("new", 3), "line", sep = "\n")
cat(moreNewLines)
## new
```

## line new

## line new

## line

#### grep

а



If we want to find patterns inside of characters or strings the grep-function family supplies various ways to find, extract or replace substrings. See the help page for grep to find out more.

```
## [1] "Some_String" "More_String"
grep("String", a)
## [1] 1 2
grep("Something", a)
## integer(0)
grepl("Some", a)
## [1]
        TRUE FALSE
sub("String", "Something", a)
## [1] "Some_Something" "More_Something"
```



Select only those columns in dat which start with "x":

dat <- as.data.frame(matrix(1:100, nrow = 10))</pre>

```
names(dat)[1:5] <- paste("x", 1:5, sep = "")
str(dat)
   'data.frame': 10 obs. of 10 variables:
##
    $ x1: int 12345678910
##
    $ x2 : int.
                11 12 13 14 15 16 17 18 19 20
    $ x3 : int
                21 22 23 24 25 26 27 28 29 30
##
##
    $ x4 : int
                31 32 33 34 35 36 37 38 39 40
##
    $ x5 : int
                41 42 43 44 45 46 47 48 49 50
##
    $ V6 : int
                51 52 53 54 55 56 57 58 59 60
                61 62 63 64 65 66 67 68 69 70
##
    $ V7 : int.
    $ V8 : int
                71 72 73 74 75 76 77 78 79 80
##
##
    $ V9 : int.
                81 82 83 84 85 86 87 88 89 90
                91 92 93 94 95 96 97 98 99 100
##
    $ V10: int
dat[1:3, grep("x", names(dat))]
##
     x1 x2 x3 x4 x5
      1 11 21 31 41
     2 12 22 32 42
```

```
names(dat)[1:2] <- c("v1x", "x_Something")</pre>
dat[1:3, grep("x", names(dat))] # that's not what we want
##
    v1x x_Something x3 x4 x5
## 1
                  11 21 31 41
## 2 2
                 12 22 32 42
                  13 23 33 43
## 3
dat %>% select(starts_with("x")) %>% slice(1:3) # dplyr
## # A tibble: 3 x 4
##
    x_Something x3 x4
                                x5
           <int> <int> <int> <int>
##
## 1
              11
                    21
                          31
                                41
## 2
              12
                    22
                          32 42
## 3
              13
                    23
                          33
                              43
```



The previous example showed that just by defining pattern = "x", grep will find any string containing the letter 'x'. Regular expressions can be used to generalize the pattern, hence to abstract it. This is implemented by using special symbols, reserved to match specific patterns. To solve the previous problem, we can modify the pattern:

```
dat[1:3, grep("^x", names(dat))]
```

```
## x_Something x3 x4 x5
## 1 11 21 31 41
## 2 12 22 32 42
## 3 13 23 33 43
```

<sup>&#</sup>x27;^' defines that the pattern to be searched will **begin with** 'x'.



For a further extension we only want to use those variables which start with an 'x' and are followed by a number between 0 and 9:

```
## [1] 3 4 5
```

Now allow 'x' to be upper or lower case:

```
grep("^[Xx][0-9]", names(dat))
```

```
## [1] 3 4 5
```

If you should ever need regular expressions, ?regex offers some information about the implementation and possibilities in R. See Spector (2008) for an introduction.

#### Final Remarks



There are many more topics to cover in the context of data handling. Some remarks if you want or need more:

- See for a focused overview P. Spector (2008): Data Manipulation with R, Springer and/or P. Teetor (2011): R Cookbook, O'Reilly for many good examples with valuable solutions to many typical problems
- ► The package tidyr for data cleaning and reshaping
- ▶ In case that you face difficulties due to really large data sets, the readr package might be helpful: similar to read.table but *much* faster!

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**Functions** 

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#### **Functions**

Wikipedia: A function is a sequence of code that performs a specific task, packaged as a unit. This unit can then be used wherever that particular task should be performed.

A function in R can be characterized by its arguments, the body and the environment in which they are defined. Functions in R are objects of class function and have the following structure:

```
[ someFunction <- ] function( <arguments> ) {
    <expression>
}
```

Functions in R are "first class objects", which means that they can be treated much like any other R object. Importantly,

- ► Functions can be passed as arguments to other functions
- ► Functions can be returned by other functions
- ► Functions can be stored in a list
- Functions can be nested, so that functions can be defined inside of another function

# A function in R is an object of class 'function' and contains three components:

Functions

- ▶ the body, can be accessed with body()
- ▶ the formals, the formal arguments list can be accessed with formals()
- the environment, which determines how variables are found inside the function can be accessed with environment()
- ▶ Printing a function to the console shows all three components
- ▶ If the environment is not specified the global environment is the environment of the function
- ▶ Typically you will add an environment to your functions when creating a package or implicitly when creating a *closure*

**Functions** 



```
f <- function(x) x</pre>
## function(x) x
formals(f)
## $x
body(f)
## x
environment(f)
```

## <environment: R\_GlobalEnv>

**Functions** 

## **Functions**

```
nrow
## function (x)
## dim(x)[1L]
## <bytecode: 0x000000013c44378>
## <environment: namespace:base>
formals(nrow)
## $x
body(nrow)
## dim(x)[1L]
environment(nrow)
## <environment: namespace:base>
```

# Functions



Functions have named arguments which potentially have default values.

- ▶ The formal arguments are the arguments included in the function definition
- Exact, positional and partial matching
- ▶ Not every function call in R makes use of all the formal arguments
- ▶ Function arguments can be *missing* or might have *default* values
- Arguments in terms of other arguments
- Arguments in terms of values inside the function

**Function Arguments** 

### Argument Matching



R functions arguments can be matched positionally or by name. So the following calls to sd are all equivalent:

```
mydata <- rnorm(100) ## Sample from N(0,1)
sd(mydata) ## Compute Standard Deviation
sd(x = mydata)
sd(x = mydata, na.rm = FALSE)
sd(na.rm = FALSE, x = mydata)
sd(na.rm = FALSE, mydata)</pre>
```

Even though it is *legal*, it is not recommended messing around with the order of the arguments too much, since it will lead to confusion.

## Argument Matching



Function arguments can also be partially matched, which is a potential cause for errors and confusion. It can be useful for *interactive work*, though. The order of operations when given an argument is

- 1. Check for exact match for a named argument
- 2. Check for a partial match
- 3. Check for a positional match

 $\label{lem:arguments} \mbox{Arguments in terms of other arguments:}$ 

```
someFunction <- function(x, y = x + 1) x + y
someFunction(2)</pre>
```

### ## [1] 5

Arguments in terms of values inside the function:

```
someFunction <- function(x, y = z + 1) {
   z <- x + 1
   x + y + z
}
someFunction(2)</pre>
```

#### ## [1] 9

Defining default values in terms of values defined inside the function itself is usually bad practise since it is hard to understand the function call without reading the code of the function itself.

The ... argument can be used to pass arguments on to other function calls.

```
myplot <- function(x, y, type = "l", ...) {
    plot(x, y, type = type, ...)
}</pre>
```

Generic functions use  $\dots$  so that extra arguments can be passed to methods (more on this later).

The "..." Argument

The ... argument is also necessary when the number of arguments passed to the function cannot be known in advance.

```
args(paste)
## function (..., sep = " ", collapse = NULL)
## NULL.
args(cat)
## function (..., file = "", sep = " ", fill = FALSE, labels = NULL,
       append = FALSE)
##
## NUT.I.
```

One catch with ... is that any arguments that appear after ... on the argument list must be named explicitly and cannot be partially matched.

```
args(paste)
## function (..., sep = " ", collapse = NULL)
## NULL
paste("a", "b", sep = ":")
## [1] "a:b"
paste("a", "b", se = ":")
## [1] "a b :"
```

### Lazy Evaluation



Arguments to functions are evaluated *lazily*, so they are evaluated only as needed.

```
f <- function(a, b) {
    a^2
}
f(2)</pre>
```

### ## [1] 4

This function never actually uses the argument  $\mathfrak{b}$ , so calling  $\mathfrak{f}(2)$  will not produce an error because 2 gets positionally matched to a.



## Another example:

```
f <- function(a, b) {
  print(a)
  print(b)
}
f(45)</pre>
```

## [1] 45

## Error in print(b): argument "b" is missing, with no default
Notice that 45 got printed first and before the error was triggered. This is because b
did not have to be evaluated until after print(a). Once the function tried to evaluate
print(b) it had to throw an error.

- ► The reason that arguments can be defined in terms of other arguments is the concept of *lazy evaluation*
- ▶ Expressions in a function are only evaluated when they are needed
- ▶ This means that before R tries to evaluate the values of the argument a new environment is created in which the evaluation takes place. At the time when the value for y is needed (when y is evaluated) x is part of the environment

```
someFunction <- function(x, y = x + 1) x + y
someFunction(2)</pre>
```

## [1] 5

In combination with the scoping rules (name masking) it follows that any object outside the function (in the global environment) named x will be masked. Hence R can only find the object named x which is the value of the argument x:

```
x <- 1
someFunction(2)</pre>
```

## [1] 5

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## Replacement Functions

We have already seen that R can distinguish functions from other objects. The same happens for replacement functions:

```
names(x) <- newNames
```

`names<-`

- ▶ When evaluating the expression R will notice, that the left hand side is not a simple name but a function call
- ▶ R will then search for a function called 'names<-'
- ▶ This implies that there are two function definitions for 'names', names and names<-:

```
str(names)
## function (x)
names
## function (x) .Primitive("names")
str(`names<-`)
## function (x, value)
```

- ► These functions act (syntactically) as if they modify their argument
- ▶ Internally they are 'ordinary' functions and are distinguished with a special naming and should return the modified input

```
"functionName<-" <- function(x, value) {
   x[subSet] <- value
   return(x)
}</pre>
```

► Consider the following example:

```
"replacementFunction<-" <- function(x, value) {
    x[2] <- value
    x
}
x <- 1:10
replacementFunction(x) <- -1
x</pre>
```

## [1] 1 -1 3 4 5 6 7 8 9 10

## Replacement Functions



- ▶ Since replacement functions are simply functions the usual function call is also valid
- ► However, when using special names the function name must be referred to using single quotes: '

```
'replacementFunction<-'(x, -2)</pre>
```

```
## [1] 1 -2 3 4 5 6 7 8 9 10
```

► The above function call will not replace the object 'x'. The following expression is much closer to how R understands replacement functions:

```
x <- 'replacementFunction<-'(x, -2)</pre>
```

▶ Note that R only knows one way to interpret a function call and that is: functionName(arguments). When evaluating replacement functions the initial expression: replacementFunction(x) <- value is rewritten as x <- 'replacementFunction<-'(x, value). The purpose of this way to call a function is to add some *sintactic sugar*.

Often it is very useful to combine replacement and subsetting. Expressions like the following are also valid:

```
x <- c(a = 1, b = 2, c = 3)
names(x)[2] <- "two"
x
```

```
## a two c
## 1 2 3
```

This is turned internally into:

```
`*tmp*` <- names(x)
`*tmp*`[2] <- "two"
names(x) <- `*tmp*`
x</pre>
```

This implies that for evaluating the above expressions two functions need to be available, names and names<-. For our replacementFunction<- this is not meaningful.

#### Infix functions



- Most of the time functions are used as 'prefix' operators
- ▶ Binary operators like '+', '-', '==' and even '<-' are also functions
- ► Functions with reserved or illegal names can be referred to using single quotes: '
- ► You can create your own infix operators using the symbol '%' there are already predefined functions in R using this structure: %\*%, %/%, etc.
- ▶ Note that you have to use double quotes to be able to assign values (the function definition) to these variable names see the examples

## Don't try this at home!!!

```
1 + 2
```

```
"+" <- function(x, y) x - y 1 + 2
```

This is why we have to protect ourselves with package creation and namespaces!

## Example: Infix Functions



A more useful example: Create an operator for pasting strings together!

```
"%+%" <- function(x, y) paste(x, y)

"some" %+% "string"
```

### ## [1] "some string"

The naming of functions in this context is more flexible: You are allowed to use all symbols except %.

```
"%paste%" <- function(x, y, ...) paste(x, y, ...)
"new" %paste% "string"</pre>
```

```
## [1] "new string"
```

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# Example: Infix Functions

Note the following function calls are equivalent.

```
## [1] 3
'+'(1, 2)
## [1] 3
and
"new" %paste% "string"
## [1] "new string"
'%paste%'("new", "string")
```

## [1] "new string" and also

```
x < -1
'<-'(x, 1)
х
```

1 + 2

## [1] 1

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## Homework: Infix Functions

As homework: Explain the following function and result!

Write an infix function %and% that behaves like the logical operator &, except that it won't work on vectors. Use only the if-else control structure and the function '=='!

TRUE %and% TRUE

## [1] TRUE

FALSE %and% TRUE

## [1] FALSE

TRUE %and% FALSE

## [1] FALSE

FALSE %and% FALSE

## [1] FALSE

### Functions inside other functions



- ► Functions can be defined anywhere in any environment
- ► Function definitions inside of other functions will define where a defined function is available the scoping rules apply as for any other object
- ▶ If function 'a' is defined inside function 'b' function 'a' will only be available during the evaluation of function 'b'

```
b <- function(x) {
   a <- function(y) y^2
   a(x)
}
b(2)</pre>
```

```
## [1] 4
```

```
exists("a")
```

```
## [1] FALSE
```

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# Scoping Rules

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### Scoping Rules - Outline



- ▶ The scoping rules is the set of rules which define how R finds objects
- ► Lexical scoping and dynamic lookup are the two concepts discussed in the following
- Lexical scoping summarizes the aspects covered by:
  - 1. The Search Path
  - 2. Name Masking
  - 3. Functions vs. Variables and determines how objects are found
- ▶ Dynamic lookup determines when to look for an object



- An environment binds a set of names to values
- ▶ Environments are objects and can have a name and be stored in lists etc.
- ► Environments are very similar to lists. Three fundamental differences:
  - Environments have reference semantics Whenever you modify an environment, you modify every copy
  - 2. Environments have parent if an object is not found in an environment, R will continue to search in the parent, which is an environment and thus has a parent ...
  - 3. Every object in an environment must have a name and the names must be unique
- ▶ Environments can be useful data structures however reference objects should only be used with great care (in R). Most R users are not familiar with reference semantics or do not expect them (in R)
- ► Typically you will only work implicitly with environments (all the time!)

## Environments are objects!

```
e <- new.env()
e$x <- 1
e$x</pre>
```

## [1] 1

```
ls(e)
```

## [1] "x"
And similar to lists!

```
a <- list(x = 1, y = 2)
f <- as.environment(a)
f$y</pre>
```

```
## [1] 2
```

ls(f)

## Exkursus: Environments - reference vs. copy-on-modify



For most objects in R the 'copy-on-modify' semantics are applied. Meaning there is no true modification possible only replacements. Every time an object is modified it is copied first, then modified and then replaces the existing object.

```
a \leftarrow list(a = 1)
b <- a
b$b <- 2
а
```

## \$a ## [1] 1

Environments are reference objects. What we use as objects are pointers to the actual values. Copying an environments means copying the pointer, not the value!

```
e <- new.env()
e$x <- 1
f <- e
f$v <- 2
e$v
```

## [1] 2

## <environment: 0x000000017a74d90>



If a parent is not explicitly set, the default is the environment in which it is created.

```
f <- new.env()
f$x <- 1
parent.env(f)
## <environment: R_GlobalEnv>
The parent can also be set explicitly:
e <- new.env(parent=f)
e$y <- 2
parent.env(e)
## <environment: 0x000000017a74d90>
f
```

### Exkursus: Environments - parent



Every environment has a parent. If an object can not be found in a specific environment the 'search' will be continued in the parent:

```
exists("x", envir = f)
## [1] TRUE
exists("y", envir = e)
## [1] TRUE
exists("x")
## [1] FALSE
exists("x", envir = e)
## [1] TRUE
exists("y", envir = f)
## [1] FALSE
```



- ▶ Every object in an environment must have a name and the names must be unique
- Sub-setting with position and logical vectors is not possible for environments only with names

```
# Try this on your own
f <- as.environment(list(a = 1, b = 2))
e <- as.environment(list(a = 1, 2))
f$a
f[["a"]]
f[1]
f[1]</pre>
```

#### The Search Path



- ▶ Every evaluation in R is made in a specific environment
- ► The workspace is the 'global environment'. It can be accessed using globalenv() or using the build-in object .GlobalEnv
- Try: ls(.GlobalEnv) to see what is in your workspace. The names should correspond to what you see in RStudio
- ► The workspace typically contains the objects you initialized in your current session - the question is then how R can find the build-in functions (like dim() since it is not in your workspace)
- ► To omit the explicit reference to a package from which we want to use a function, the search mechanism for environments is applied

- Since the workspace, the global environment, is an environment it has a parent.
   Typically this is the last package loaded, which itself is again an environment
- ▶ The parent of the last package loaded is the package loaded before
- ▶ The connection via the parent between the loaded packages is the search path
- ► The search path is used to determine the order in which the loaded packages are searched to find a function or any other object
- ▶ The 'end' of the search path is package:base (baseenv())
- ► The parent of package:base is the empty environment, emptyenv(), which itself has no parent, is empty and can not contain any objects: the search ends here



```
search()
  [1] ".GlobalEnv"
                                                 "package:graphics"
                            "package:stats"
   [4] "package:grDevices" "package:utils"
                                                 "package:datasets"
   [7] "Autoloads"
                            "package:base"
library(ggplot2)
search()
## [1] ".GlobalEnv"
                            "package:ggplot2"
                                                 "package:stats"
                            "package:grDevices"
                                                 "package:utils"
   [4] "package:graphics"
```

"Autoloads"

"package:base"

[7] "package:datasets"

### The Search Path



## Searching for objects - the function dim

```
funOfInterest <- "dim"
loadedEnv <- search()
objInEnv <- sapply(as.list(loadedEnv),
 function(env, ...) exists(envir = as.environment(env), ...),
 x = funOfInterest, inherits = F)
loadedEnv[objInEnv]
## [1] "package:base"
Make it a function:
searchForFun <- function(funOfInterest) {</pre>
  loadedEnv <- search()</pre>
  objInEnv <- sapply(as.list(loadedEnv),
    function(env, ...) exists(envir = as.environment(env), ...),
    x = funOfInterest, inherits = F)
  loadedEnv[objInEnv]
```



```
Finding functions:
searchForFun("dim")
## [1] "package:base"
searchForFun("plot")
```

```
## [1] "package:graphics"
```

```
searchForFun("qplot")
## [1] "package:ggplot2"
```

```
x <- 1
```

```
searchForFun("x")
```

```
## [1] ".GlobalEnv"
```

## Name Masking

dim



Whenever there is more than one object with the same name, the object in the 'closest' environment will be selected. An environment is closer if it appears earlier in the search path.

```
## function (x) .Primitive("dim")
dim <- function(x) x
searchForFun("dim")
## [1] ".GlobalEnv" "package:base"
dim</pre>
```

```
## function(x) x
```

Here two functions with the same name can be found. Whenever you call the function dim, R will use the version in your workspace. The function dim in the base package is not overritten, it is simply masked by the function in the workspace.

### Excursus: Environments and functions



- Every call to a function creates a temporal environment in which the function call is evaluated
- ► This guarantees (potentially) that the evaluation does not affect the global environment your workspace
- ▶ This also determines how names are looked for inside a function (we only consider user written functions) if not specified differently the environment is the global environment plus what is passed into the function in form of *formal arguments* and objects created inside the function
- Objects created inside of a function and passed to the function by formal arguments are called local and are only available during the function call



The environment of nrow guarantees that nrow will allways find the 'correct' function named dim which is defined in the base package.

```
x \leftarrow data.frame(V1 = 1:2)
str(x)
## 'data.frame': 2 obs. of 1 variable:
## $ V1: int. 1 2
dim <- function(x) 1
newNrow <- function(x) dim(x)[1L]
nrow(x)
## [1] 2
newNrow(x)
```

## [1] 1



# Consider the following example:

```
x <- 1
someFunction <- function(x) {
    x <- x + 1
    x
}
someFunction(x)</pre>
```

# ## [1] 2

What is the value of x?



# Consider the following example:

```
x <- 1
someFunction <- function(x) {
  y <- x + 1
  y
}
someFunction(x)</pre>
```

```
## [1] 2
What is the value of y?
```

### Excursus: Environments and functions



# Consider the following examples:

```
x <- 1 someFunction <- function() x
```

What is the result of a call to someFunction?

```
rm(list=ls())
x <- 1
someFunction <- function() x * y</pre>
```

What is the result of a call to someFunction?



The same rules apply if you define functions inside other functions:

```
x <- 1
someFunction <- function() {
  y <- 2
  otherFunction <- function() {
    z <- 3
    c(x, y, z)
  }
  otherFunction()
}</pre>
```

What is the result of a call to someFunction?



What is the result of the second call to someFunction?

```
rm(list = ls())
someFunction <- function() {
   if (!exists("a")) {
      a <- 1
   } else {
      a <- a + 1
   }
   print(a)
}</pre>
```

Every time a function is called a new environment is created for the evaluation. The object a will only be created locally.



More name masking as homework: What is the return value of f(10)?

```
f <- function(x) {
  f <- function(x) {
    f <- function(x) {
       x ^ 2
    }
    f(x) + 1
}
f(x) * 2
}
f(10)</pre>
```

### Functions vs. variables



If you search for an object from a context from which it is obvious that you are searching a function (a function call like someFunction(2) the search is continued until R finds a *function* ignoring other objects with the same name. Thus the following code is valid:

```
c <- 1
c(c, c, 2)
```

### ## [1] 1 1 2

Be aware that the lookup for the value 'c' and the function 'c' is possible since both objects named 'c' are defined in different environments. This is not possible when working in the global environment (names in an environment must be unique!):

```
f <- 10
f <- function(x) x
f(f)</pre>
```

```
## function(x) x
```

Of course this may get very confusing, proper naming is preventive ... If we wanted to rewrite the function searchForFun defined previously in order to only search for functions, we would have to use match.fun instead of exists. Try this at home!



- ▶ Lexical scoping determines where to look for values, not when to look for them
- ▶ R will search for objects when a function is run, not when it is created
- This means that a function can have different return values which depend on objects defined in the environment in which it is created and run
- This means an important property of well defined functions is violated self containment
- ► The property of self containment is fulfilled if the return value of a function depends only on the input values (formal arguments)
- You want to write self contained functions. If you don't it is very easy to produce unexpected results

## Dynamic lookup



 $\ensuremath{\mathsf{R}}$  will search for objects when a function is run, not when it is created

```
rm(list=ls())
someFunction <- function(x) x + y</pre>
```

The above expression is valid since R will not try to find the value of y until someFunction is called. Depending on the value of y the function someFunction will have a different return value:

```
y <- 1
someFunction(1)</pre>
```

## [1] 2

```
y <- 2 someFunction(1)
```

## [1] 3

In this definition someFunction is not self containing - such behaviour should be avoided if possible

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#### Control Structures



Control structures in R allow you to control the flow of execution of the program, depending on runtime conditions. Common structures are

if, else: testing a condition

for: execute a loop a fixed number of times

while: execute a loop while a condition is true

repeat: execute an infinite loop

break: break the execution of a loop

next: skip an interation of a loop

Most control structures are not used in interactive sessions, but rather when writing functions or longer expresissons.

```
if(<condition>) {
    <do something>
} else {
    <do something else>
}

if(<condition1>) {
    <do something>
} else if(<condition2>) {
    <do something different>
} else {
    <do something different>
}
```

This is a valid if/else structure.

So is this one.

$$y < -if(x > 3)$$
 { 10 } else { 0 }

Of course, the else clause is not necessary.

```
if(<condition1>) {
}
if(<condition2>) {
}
```

### Control Structures: for



for loops take a *loop variable* and assign it successive values from a sequence or a vector. For loops are most commonly used for iterating over the elements of an object (list, vector, etc.)

```
for(i in 1:10) {
      print(i)
}
```

This loop takes the i variable and in each iteration of the loop gives it the successive values contained in vector 1:10 and then exits.

These four loops have the same behavior.

```
x <- c("a", "b", "c", "d")
for(i in 1:4) {
    print(x[i])
}

for(i in seq_along(x)) {
    print(x[i])
}

for(letter in x) {
    print(letter)
}

for(i in x) print(i)</pre>
```

### Nested for loops



for loops can be nested

```
x <-matrix(data = 1:6, ncol = 2, ncol = 3, byrow = TRUE))
for(i in seq_len(nrow(x))) {
    for(j in seq_len(ncol(x))) {
        print(x[i, j])
    }
}</pre>
```

Be careful with nesting. Nesting beyond 2-3 levels is often very diffcult to read/understand.

### Control Structures: while



While loops begin by testing a condition. If it is true, then they execute the loop body. Once the loop body is executed, the condition is tested again, and so forth.

```
number <- 0
while(number < 10) {
    print(number)
    number <- number + 1
}</pre>
```

While loops can potentially result in infinite loops if not written properly. Use with care!

Sometimes there will be more than one condition in the test.

```
 z < -5  while(z >= 3 && z <= 10) { print(z) coin <- rbinom(n = 1, size = 1, prob = 0.5) } 
 if(coin == 1) { ## random walk z <- z + 1 } else { z <- z - 1 } 
 }
```

Conditions are always evaluated from left to right.

### Control Structures: repeat



repeat initiates an infinite loop. The only way to exit a repeat loop is to call break.

```
 \begin{array}{l} \text{x0} < -1 \\ \text{tol} < -1\text{e} - 8 \\ \\ \text{repeat } \{ \\ \text{x1} < -\text{ computeEstimate}(\text{x0}) \\ \\ \text{if}(\text{abs}(\text{x1} - \text{x0}) < \text{tol}) \ \{ \\ \\ \text{break} \\ \} \ \text{else} \ \{ \\ \\ \text{x0} < - \text{x1} \\ \} \\ \} \end{array}
```

This loop is a bit dangerous because there is no guarantee it will stop. Better to set a hard limit on the number of iterations (e.g. using a for loop) and then report whether convergence was achieved or not.



 ${\tt next}$  is used to skip an iteration of a loop

#### Control Structures



# Summary

- Control structures like if, while and for allow you to control the flow of an R program
- $\,\blacktriangleright\,$  Infinite loops should generally be avoided, even if they are theoretically correct
- Control structures mentiond here are primarily useful for writing programs;
- ► For command-line interactive work and "advanced" code, the \*apply functions are more useful and advisable.

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Writing for, while loops is useful when programming but not particularly easy when working interactively on the command line. There are some functions which implement looping to make life easier.

lapply Loop over a list and evaluate a function on each element

sapply Same as lapply but tries to simplify the result

mapply Multivariate version of lapply

An auxiliary function split is also useful, particularly in conjunction with lapply.



lapply takes three arguments: a list X, a function (or the name of a function) FUN, and other arguments via its ... argument. If X is not a list, it will be coerced to a list using as.list.

# lapply

```
## function (X, FUN, ...)
## {
## FUN <- match.fun(FUN)
## if (!is.vector(X) || is.object(X))
## X <- as.list(X)
## .Internal(lapply(X, FUN))
## }
## <bytecode: 0x0000000011e3e080>
## <environment: namespace:base>
```

The actual looping is done internally in C code which makes it fast.

lapply always returns a list, regardless of the class of the input

```
lapply(x, mean)
## $a
## [1] 3
##
##
   $Ъ
## [1] -0.5025504
```

 $x \leftarrow list(a = 1:5, b = rnorm(10))$ 

```
x <- list(a = 1:4,
         b = rnorm(10),
         c = rnorm(20, 1),
          d = rnorm(100, 5))
lapply(x, mean)
## $a
## [1] 2.5
##
## $b
## [1] -0.0647194
##
## $c
## [1] 1.185596
##
## $d
```

## [1] 4.796942

```
x < -1:4
lapply(x, runif)
## [[1]]
## [1] 0.4734999
##
## [[2]]
  [1] 0.3191864 0.4539071
##
## [[3]]
   [1] 0.0649204 0.3284327 0.9000433
##
## [[4]]
## [1] 0.2830072 0.1403274 0.9073102 0.0922993
```

```
x < -1:4
lapply(x, runif, min = 0, max = 10)
## [[1]]
## [1] 4.460217
##
## [[2]]
## [1] 2.645574 7.867302
##
## [[3]]
   [1] 0.8756886 4.0346447 4.5062040
##
## [[4]]
## [1] 2.236603 8.967762 6.456188 7.043500
```

```
x < -1:4
lapply(x, runif, min = 0, n = 10)
## [[1]]
    [1] 0.9497713 0.7705003 0.7727400 0.3789881 0.9652450 0.5760081 0.771
##
    [8] 0.8329985 0.8637544 0.7529941
##
## [[2]]
##
    [1] 1.854488368 0.009708281 0.257603093 0.454071679 0.624560753
##
    [6] 1.859912145 1.591564231 0.704509218 1.128583749 1.847928698
##
## [[3]]
    [1] 0.5820046 1.2259634 1.2273919 1.3487113 2.6360802 0.9046120 2.315
##
##
    [8] 0.1032324 0.9962058 2.7800568
##
## [[4]]
    [1] 0.1910453 0.9689292 2.4099732 3.4518710 1.8079039 1.5293630 0.897
##
##
    [8] 1.5784680 2.2590198 0.2001869
```

lapply and friends make heavy use of anonymous functions.

```
x <- list(a = matrix(data = 1:4, nrow = 2, ncol = 2, byrow = FALSE),
        b = matrix(data = 1:6, nrow = 3, ncol = 2, byrow = TRUE))
х
## $a
       [,1] [,2]
##
## [1,] 1
## [2.] 2
##
## $b
       [,1] [,2]
##
## [1,] 1
## [2,] 3 4
## [3,] 5
               6
```

An anonymous function for extracting the first column of each matrix.

```
lapply(x, function(y) y[ ,1])
## $a
## [1] 1 2
##
## $b
## [1] 1 3 5
```



sapply will try to simplify the result of lapply if possible.

- ▶ If the result is a list where every element is length 1, then a vector is returned.
- $\triangleright$  If the result is a list where every element is a vector of the same length (>1), a matrix is returned.
- ▶ If it can't figure things out, a list is returned.
- Check the manual for the 'sapply' argument 'simplify'.

```
## Warning in mean.default(x): argument is not numeric or logical: return
## NA
## [1] NA
```

split takes a vector or other objects and splits it into groups determined by a factor or list of factors.

```
str(split)
```

```
## function (x, f, drop = FALSE, ...)
            x is usually a data frame
             f is a factor (or coerced to one) or a list of factors
         drop indicates if empty factor levels should be dropped
           ... further potential arguments passed to methods.
```

# Splitting a Data Frame



```
head(airquality, n=3)
##
     Ozone Solar.R Wind Temp Month Day
## 1
        41
               190
                     7.4
                           67
                                  5
                                       1
## 2
        36
               118 8.0
                           72
        12
               149 12.6 74
## 3
s <- split(airquality[, c("Ozone", "Solar.R", "Wind")],
          f = airquality$Month)
sapply(s, colMeans)
##
                   5
                             6
## Ozone
                 NA
                            NA
                                        NA
                                                 NA
                                                           NA
## Solar.R.
                 NA 190.16667 216.483871
                                                 NA 167.4333
## Wind
           11.62258
                      10.26667 8.941935 8.793548
                                                     10.1800
sapply(s, colMeans, na.rm=TRUE)
                    5
                                                      8
##
                              6
            23.61538
                       29.44444
                                 59.115385
                                             59.961538
   Ozone
```

Solar R 181, 29630 190, 16667 216, 483871 171, 857143 167, 43333

11.62258 10.26667

8.793548

10.18000

8.941935

## Wind



Another common idiom is to split-apply-and-combine data.

```
dataList <- lapply(split(x, f), quantile) #Split-Apply
do.call(cbind, dataList) #Combine the result</pre>
```

```
## 0% -2.2879731 0.007736768 -0.1886691

## 25% -0.5053897 0.391566486 0.3150691

## 50% -0.1151033 0.615670335 0.6559077

## 75% 0.8399117 0.754152272 2.2661981

## 100% 1.8631307 0.809778480 2.6267326
```

# split

This is equivalent to:

```
sapply(split(x, f), quantile)
```

```
##
## 0%
       -2.2879731 0.007736768 -0.1886691
## 25% -0.5053897 0.391566486 0.3150691
## 50%
       -0.1151033 0.615670335 0.6559077
## 75% 0.8399117 0.754152272 2.2661981
## 100% 1.8631307 0.809778480 2.6267326
```

However, sapply automates the "combine"-step - it may not allways be what you expect!

Use do.call(FUN, list) to control the output structure.

# supplement strsplit, lapply and do.call



Split a character vector and recombine the results as a  ${\tt data.frame}$ 

```
names <- c("Adam Riese", "Albert Einstein", "Oliver Kahn")
namesList <- strsplit(names, split = " ")</pre>
str(namesList)
## List of 3
## $ : chr [1:2] "Adam" "Riese"
## $ : chr [1:2] "Albert" "Einstein"
## $ : chr [1:2] "Oliver" "Kahn"
namesList <- lapply(namesList,
                   function(x) data.frame(firstName = x[[1]],
                                        familyName = x[[2]])
do.call(rbind, namesList)
##
```

```
## firstName familyName
## 1 Adam Riese
## 2 Albert Einstein
## 3 Oliver Kahn
```



mapply is a multivariate apply of sorts which applies a function in parallel over a set of arguments.

```
str(mapply)
## function (FUN, ..., MoreArgs = NULL, SIMPLIFY = TRUE, USE.NAMES = TRUE
```

... contains arguments to apply over

FUN is a function to apply

MoreArgs is a list of other arguments to FUN

SIMPLIFY indicates whether the result should be simplified

USE. NAMES if the first ... argument has names, use that names

```
## the hard way
lHard \leftarrow list(rep(1, 4), rep(2, 3), rep(3, 2), rep(4, 1))
## the smart way
lSmart <- mapply(rep, 1:4, 4:1)</pre>
1Smart
## [[1]]
## [1] 1 1 1 1
##
## [[2]]
## [1] 2 2 2
##
## [[3]]
## [1] 3 3
##
## [[4]]
## [1] 4
```



```
noise <- function(n, mean, sd) {
    rnorm(n, mean, sd) }
    noise(5, 1, 2)

## [1] -2.0643429  2.8075691 -0.3190682  1.6783004  2.6576145

noise(1:4, 1:4*10, 2)
```

7.630958 19.802885 26.705537 37.984838

Г17



```
set.seed(15061969)
mapply(noise, 1:4, 1:4*10, 2)
## [[1]]
## [1] 7.44371
##
## [[2]]
   [1] 19.65748 18.59855
##
## [[3]]
   [1] 30.27678 32.84333 27.59776
##
## [[4]]
   [1] 39.69425 39.72227 39.95984 40.35376
```



# Which is the same as

```
set.seed(15061969)
list(noise(1, 10, 2), noise(2, 20, 2),
    noise(3, 30, 2), noise(4, 40, 2))
## [[1]]
## [1] 7.44371
##
## [[2]]
## [1] 19.65748 18.59855
##
## [[3]]
## [1] 30.27678 32.84333 27.59776
##
## [[4]]
   [1] 39.69425 39.72227 39.95984 40.35376
```

# Vectorizing a Function internally

```
Freie Universität Berlin
```

```
noise <- function(n, mean, sd) {</pre>
 mapply(rnorm, n, mean, sd)
set.seed(29081975)
noise(1:4, 1:4*10, 2)
## [[1]]
## [1] 6.873807
##
## [[2]]
## [1] 24.88267 18.17274
##
## [[3]]
   [1] 29.15017 29.93314 28.98341
##
## [[4]]
   [1] 42.12792 39.37629 37.73670 39.20634
```



# \*apply vs for loops

- \*apply functions are fast
- ▶ there are special packages so that \*apply functions can be calculated parallel on multicore systems  $\rightarrow$  super-fast
- but:

each element is calculated separately; calculations that depend on the outcome of previous elements are not possible with \*apply but with for loops.



- Let's assume we want to define a mean for the *character* type in R.
- ▶ As mean we simply define the mean length of each element in a vector of type character. The length of an element can be defined as the number of characters.

```
characterVector <- c("some", "more", "text", "and", "different", "nchar")</pre>
meanCharacter <- function(x) mean(nchar(x))
meanCharacter(characterVector)
```

#### ## [1] 4.833333

S3: Motivation

- ▶ There is one obstacle with the definition of meanCharacter, the name. With every function we define we have to memorize one more function name and also we have to find new useful names (which is not easy!).
- ▶ How is a potential user (you in 2 weeks) ever going to understand and know about all the functions you defined? Not at all.

- ▶ If the name is a problem let's define a new mean function.
- ▶ One thing to keep in mind, however, is that we should preserve the behaviour of the original mean function.

```
mean <- function(x) {
  if (is.character(x)) {
    base::mean(nchar(x))
  } else {
    base::mean(x)
  }
}
mean(characterVector)</pre>
```

#### ## [1] 4.833333

- Every time we want to define a mean function for a new data type we have to add more if clauses to the definition. In the long run this strategy is going to be a mess.
- Also we can not extend functionality, we have to change it. Extending is good, changing is bad.

- ▶ The basic idea of the S3 class system is that it should be possible to extend the functionality of a generic vocabulary without adding new words (function names).
- ▶ This is a convenience. You can throw any statistical model and data type into the summary function. And somehow (allmost) for all data types (linear models, data frames, etc.) summary knows what to do.
- ► Even more, summary knows about data types from different packages, although the original author had no way of anticipating these types.
- ▶ To understand this we need to answer the three following questions:
  - ▶ What is a S3 class?
  - What is a generic function?
  - ▶ What is a method?



- ▶ S3 classes serve a simple purpose, to give different or new data types a name.
- ▶ This is done by adding an attribute to the data with the name of the class.

```
class(1:10)
## [1] "integer"
class("a")
## [1] "character"
dat \leftarrow list(c(1, 2), c(2, 3), c(2, 4))
class(dat) <- "rational"
str(dat)
## List of 3
    $ : num [1:2] 1 2
##
    $ : num [1:2] 2 3
```

##

##

\$ : num [1:2] 2 4

- attr(\*, "class")= chr "rational"



▶ Typically you do not assign the class *interactively* but you return data with a class attribute from a constructor function (e.g. numeric, list, lm).

```
rational <- function(num, denom) {
    rat <- mapply(c, num, denom, SIMPLIFY = FALSE)
    class(rat) <- "rational"
    rat
}
str(rational(c(1, 2, 2), c(2, 3, 4)))

## List of 3
## $ : num [1:2] 1 2
## $ : num [1:2] 2 3
## $ : num [1:2] 2 4
## - attr(*. "class") = chr "rational"</pre>
```

- ▶ rational is a constructor function for instances of class rational.
- It is named constructor because it knows how to construct an object of class rational.
- ▶ There is no formal definition of the class, you simply say a list is of class *X*.
- ▶ In most scenarios the *list* type is used as the basic data structure to compose new data (e.g. data.frame and lm).
- ▶ So what is the great benefit of defining new classes? An example: Printing an object of class *rational* to the console results in verbose information. Now we can fix this:

```
print.rational <- function(x, ...) cat(sapply(x, paste, collapse = "/")) rational(c(1, 2, 2), c(2, 3, 4))
```

## 1/2 2/3 2/4

#### Generic functions and methods



- ► A generic functions is a function which only purpose is to find the appropriate method given the class of its first argument.
- ▶ The overall purpose is, that you have a function name, say mean, and this function somehow figures out how the mean is defined for a given data type.
- The function print and mean are generic functions. print will find the correct print method to print things to the console. And mean searches for the correct mean method for a given data type.
- ► Methods are defined by a naming convention: <generic>.<class>

```
print.rational <- function(x, ...) cat(sapply(x, paste, collapse = "/"))</pre>
```

▶ We say print.rational is the print method for objects of class *rational*. There is a print method for most of the data types in R which define how output is printed to the console.



▶ You are not restricted on a given set of generic functions, you can define them.

```
mean <- function(x, ...) UseMethod("mean")</pre>
```

- ▶ This is it. UseMethod is used to start searching for methods. It will search for a function called mean. <class(x)> and passes the arguments to this method. This search is also called method dispatch.
- ▶ If no method for a given class can be found a *default* method is called if it exists:

```
mean("a")
## Warning in mean.default("a"): argument is not numeric or logical: retu
## NA
```

- ## [1] NA
  - ▶ You define default methods with a naming convention: <generic>.default

► We return to our initial example of defining means. We discovered that if-else constructs may be problematic when the number of branches can grow. S3 offers a solution:

```
mean.character <- function(x, ...) mean(nchar(x), ...)
mean.rational <- function(x, ...) mean(sapply(x, function(e) e[1] / e[2], ...))
mean(characterVector)

## [1] 4.833333

num <- 1:6
denom <- 11:16
mean(rational(num, denom))</pre>
```

- ▶ Defining generics: generic <- function(<args>) UseMethod("generic")
- ▶ Defining methods: generic.<class> <- function(<args>) ...
- ▶ Defining default method: generic.default <- function(<args>) ...
- ▶ The arguments of methods need to include all arguments of the generic even if they are not used. Methods can have more arguments than the generic, though.
- Defining a class:

```
myClass <- function(<args>) {
    ...
    class(out) <- "myClass"
    out
}</pre>
```



- ▶ The S3 class system was introduced in version 3 of the S software, hence its name.
- Essentially it is a naming convention.
- Methods are associated with generic functions (<generic>.<class>).
- S3 classes are defined by adding an attribute to any object in R.
- It is a special form of object-orientation, but not to be confused with other implementations in other languages which can be very different!
- One of the main benefits of the S3 system is that you do not have to remember to many function names but can rely on a generic vocabulary.
- ► To find out more about which generic functions exist see the help pages for .S3methods.

#### More final remarks



- ▶ Like in many other languages there was a need to extend the simple class system of S3.
- ▶ Hence a new version of the S software (version 4) introduced a novel system for object orientation, S4.
- ▶ Main features in contrast to S3 are that S4 has formal class definitions. So if you see an object of class Im you can be sure about its properties. Method dispatch for more than one argument is possible in S4, in S3 only the first is used.
- Also there was a need in the community to support a system similar to languages like Java. This is implemented in the function setRefClass.
- ▶ The R community absolutely does not agree on how object-orientation should be implemented. An indication for that is the variety of packages on CRAN which implement different class systems (e.g. methods, R6, R.oo and proto are frequently used).
- ▶ Should you only plan to do interactive analysis in R then it is save to say that it is sufficient to know about S3.

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The \*apply Functions

Packages

#### Motivation



- Reusable code: why don't we just write scripts?
- Stability: dependencies and namespaces vs. library
- ▶ Documentation: dedicated and structured documentation vs. comments inline
- ► Some minimal standards for publishing code: R CMD CHECK

Trustworthy Software: The Prime Directive - Chambers (2008)

### Minimal package structure



- ► To initialize a package use package.skeleton() or devtools::create() or the RStudio IDE
- Must have ingredients:
  - ► Folder with \*.R-files, named 'R'
  - ► DESCRIPTION-file
    - Package, Version, License, Description, Title, Author, Maintainer
  - ▶ NAMESPACE-file
    - export, exportPattern, import, etc.

- ► The folder 'R' is mandatory and has to be located in the root of the package project
- ► All files in the folder 'R' have the ending \*.R or \*.r
- All files in 'R' are simply sourced in alphanumeric order
- All R objects defined in these files are part of the package. Typically you only find functions, data is stored separately

```
mylm <- function(y, X) {
   solve(crossprod(X), crossprod(X, y))
}</pre>
```

Freie Universität

Package: mylm Version: 0.1.1 License: MIT

DESCRIPTION

Description: Test-Package Title: mylm Test-Package Author: Sebastian Warnholz

Maintainer: <Sebastian.Warnholz@fu-berlin.de>

### NAMESPACE & Dependencies



- ▶ Defines the namespace of a package: think of it as the search path for a function living in a package
- ▶ Depends/Imports are fields in the DESCRIPTION: Other needed packages
  - To ensure that all dependencies are available
  - Packages listed in Depends are attached to the search path when your package is loaded. This should be avoided because of potential naming conflicts.
- ▶ import (command in NAMESPACE): Single objects (e.g. functions) or packages which should be available *inside* the package
- export (command in NAMESPACE): Which functions or objects do you want to make available to the user

Possible statements in NAMESPACE:

```
export(<functionName>)
exportPattern("^[^\\.]")
import(<packageName>)
importFrom(<packageName>, <functionName>)
S3method(<genericName>, <className>)
useDynLib(<libName>)
```

The NAMESPACE-file will be generated on the fly by a package called roxygen2.

Packages

- ▶ In a package all *exported* objects have to be documented
- Documentation files are \*.Rd-files located in the folder 'man'
- Those files are used to build a HTML and PDF documentation
- ▶ The package roxygen2 simplifies writing the documentation dramatically

Packages

- ► The \*.Rd-files do not have to be created manually
- ► Also the NAMESPACE file is updated automatically

Documentation

#### Documentation for mylm



```
My linear model
# '
#' @description Computes coefficients of a linear model
#' @param y dependent variable
#' Oparam X design matrix
# '
#' Oreturn An object of class \code{mylm}
#' @details More detailes
# '
#' @export
#' @examples mylm(rnorm(10), rnorm(10))
mylm <- function(y, X) {
  beta <- solve(crossprod(X), crossprod(X, y))
  class(beta) <- "mylm"</pre>
  beta
```

#### Remarks



- ▶ Best Practice:
  - ► Run R CMD CHECK frequently
  - ► Tests, vignettes, version control, continuous integration
- ▶ Very helpful packages: roxygen2, devtools, testthat
- Resources: r-pkgs, Tutorial

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OOP: S3

FU-BerlincRi-Brogramming ramming





- ▶ Let's assume we want to define a mean for the *character* type in R.
- As mean we simply define the mean length of each element in a vector of type character. The length of an element can be defined as the number of characters.

```
characterVector <- c("some", "more", "text", "and", "different", "nchar")
meanCharacter <- function(x) mean(nchar(x))
meanCharacter(characterVector)</pre>
```

#### ## [1] 4.833333

- ▶ There is one obstacle with the definition of meanCharacter, the name. With every function we define we have to memorize one more function name and also we have to find new useful names (which is not easy!).
- ▶ How is a potential user (you in 2 weeks) ever going to understand and know about all the functions you defined? Not at all.

- ▶ If the name is a problem let's define a new mean function.
- ▶ One thing to keep in mind, however, is that we should preserve the behaviour of the original mean function.

```
mean <- function(x) {
  if (is.character(x)) {
    base::mean(nchar(x))
  } else {
    base::mean(x)
  }
}
mean(characterVector)</pre>
```

#### ## [1] 4.833333

- Every time we want to define a mean function for a new data type we have to add more if clauses to the definition. In the long run this strategy is going to be a mess.
- Also we can not extend functionality, we have to change it. Extending is good, changing is bad.

- ▶ The basic idea of the S3 class system is that it should be possible to extend the functionality of a generic vocabulary without adding new words (function names).
- ▶ This is a convenience. You can throw any statistical model and data type into the summary function. And somehow (allmost) for all data types (linear models, data frames, etc.) summary knows what to do.
- ▶ Even more, summary knows about data types from different packages, although the original author had no way of anticipating these types.
- ► To understand this we need to answer the three following questions:
  - ▶ What is a S3 class?
  - What is a generic function?
  - ▶ What is a method?

- ▶ S3 classes serve a simple purpose, to give different or new data types a name.
- ▶ This is done by adding an attribute to the data with the name of the class.

```
class(1:10)
## [1] "integer"
class("a")
## [1] "character"
dat \leftarrow list(c(1, 2), c(2, 3), c(2, 4))
class(dat) <- "rational"
str(dat)
## List of 3
    $ : num [1:2] 1 2
##
    $ : num [1:2] 2 3
```

##

##

\$ : num [1:2] 2 4

- attr(\*, "class")= chr "rational"

▶ Typically you do not assign the class *interactively* but you return data with a class attribute from a constructor function (e.g. numeric, list, lm).

```
rational <- function(num, denom) {
    rat <- mapply(c, num, denom, SIMPLIFY = FALSE)
    class(rat) <- "rational"
    rat
}
str(rational(c(1, 2, 2), c(2, 3, 4)))

## List of 3
## $ : num [1:2] 1 2
## $ : num [1:2] 2 3
## $ : num [1:2] 2 4</pre>
```

- attr(\*, "class")= chr "rational"

##

- rational is a constructor function for instances of class rational
- ▶ It is named *constructor* because it knows how to construct an object of class rational.
- ▶ There is no formal definition of the class, you simply say a list is of class X.
- ▶ In most scenarios the *list* type is used as the basic data structure to compose new data (e.g. data.frame and lm).
- ▶ So what is the great benefit of defining new classes? An example: Printing an object of class rational to the console results in verbose information. Now we can fix this:

```
print.rational <- function(x, ...) cat(sapply(x, paste, collapse = "/"))</pre>
rational(c(1, 2, 2), c(2, 3, 4))
```

## 1/2 2/3 2/4

OOP: \$3

- A generic functions is a function which only purpose is to find the appropriate method given the class of its first argument.
- ▶ The overall purpose is, that you have a function name, say mean, and this function somehow figures out how the mean is defined for a given data type.
- ▶ The function print and mean are generic functions. print will find the correct print method to print things to the console. And mean searches for the correct mean method for a given data type.
- ▶ Methods are defined by a naming convention: <generic>.<class>

```
print.rational <- function(x, ...) cat(sapply(x, paste, collapse = "/"))</pre>
```

▶ We say print.rational is the print method for objects of class *rational*. There is a print method for most of the data types in R which define how output is printed to the console.

### More on generic functions



▶ You are not restricted on a given set of generic functions, you can define them.

```
mean <- function(x, ...) UseMethod("mean")</pre>
```

- ▶ This is it. UseMethod is used to start searching for methods. It will search for a function called mean. <class(x)> and passes the arguments to this method. This search is also called method dispatch.
- ▶ If no method for a given class can be found a *default* method is called if it exists:

```
mean("a")
## Warning in mean.default("a"): argument is not numeric or logical: retu
  NΑ
```

- ▶ You define default methods with a naming convention: <generic>.default

## [1] NA

► We return to our initial example of defining means. We discovered that if-else constructs may be problematic when the number of branches can grow. S3 offers a solution:

```
mean.character <- function(x, ...) mean(nchar(x), ...)
mean.rational <- function(x, ...) mean(sapply(x, function(e) e[1] / e[2], ...))
mean(characterVector)

## [1] 4.833333

num <- 1:6
denom <- 11:16
mean(rational(num, denom))</pre>
```

- ▶ Defining generics: generic <- function(<args>) UseMethod("generic")
- ▶ Defining methods: generic.<class> <- function(<args>) ...
- ▶ Defining default method: generic.default <- function(<args>) ...

OOP: \$3

- ▶ The arguments of methods need to include all arguments of the generic even if they are not used. Methods can have more arguments than the generic, though.
- Defining a class:

```
myClass <- function(<args>) {
    class(out) <- "myClass"</pre>
    0111.
```

#### Final remarks



- ▶ The S3 class system was introduced in version 3 of the S software, hence its name.
- Essentially it is a naming convention.
- Methods are associated with generic functions (<generic>.<class>).
- ▶ S3 classes are defined by adding an attribute to any object in R.
- It is a special form of object-orientation, but not to be confused with other implementations in other languages which can be very different!
- One of the main benefits of the S3 system is that you do not have to remember to many function names but can rely on a generic vocabulary.
- To find out more about which generic functions exist see the help pages for .S3methods.



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At it's heart R is a Functional Programming Language. This means that functions are First Class Citizens, i.e. functions can be treated like any other object in R. The following will address the implications of this statement:

- ► Functions can be defined without a name (anonymous function)
- ► Functions can be defined anywhere, including inside other functions
- 'Functionals' Like any other value, they can be passed as parameters to functions
- 'Closure' Like any other value, they can be returned as results from functions
- ▶ Like any other value, they can be stored in lists
- ▶ As for other values, there exists a set of operators to compose functions

Function composition, the last bullet, is an important technique in functional programming. However, this is beyond the scope of this course. Understanding all other implications is fundamental to function composition and will already enable you to accomplish any task you'll ever encounter (in fewer lines).

- 1. Functions which take functions as parameters functional
- 2. Functions which return functions closure

#### Functional:

```
 \begin{array}{ll} \mbox{functional} < - \mbox{ function}(f, \ ...) \ f(...) \\ \mbox{functional}(\mbox{mean, runif}(1e3)) \\ \mbox{functional}(\mbox{sum, runif}(1e3)) \end{array}
```

#### Closure:

```
closure < - function(x) function(y) x + y closure(1)(2) closure(2)(3)
```

# Programming

The \*apply Functions

# Evaluation - Outline





#### Homework: Evaluation



Rewrite the following expressions as it has been done for line 1 to 2:

- 1. (1:10)[2]
- 2. sum(1+1:10)
- 3. 1:10^2
- 4. x <- 1:10+2^2

The rewritten expressions can be said to be evaluated from left to right. The functions which you can rewrite are '(', ':', '[', '+', '^' and '<-'. Check your results in R. What is meant by the phrase: "Everything in R is a call to a function!"?

## Outline

# Programming

The \*apply Functions

#### Overview



# **Debugging** in general:

- running a program step by step
- pausing a program to examine the current state
- tracking the values of some variables
- modify the program while it is running

### Since R is an *interpreter* language, debugging in R means **debugging functions**:

- running a function step by step
- pausing a function to examine the current state
- ► tracking the values of some variables inside<sup>3</sup> functions
- modify the function while it is running

<sup>&</sup>lt;sup>3</sup>variables that (only) exist inside the environment of the function

# Debugging in R



Functions for debugging (package: base):

prints the call stack of the last uncaught error traceback

flags a function for debugging debug

interrupts the execution of an expression and allows the inspection of browser

the environment

allows to insert debugging code into an existing function trace

allows to browse directly on any of the currently active function calls recover

#### Motivation



As with programs written in any other language, functions written in R can contain unforseen problems which lead to failure.

The **purpose of the debugging** tools is to help the programmer find these problems quickly and effciently.<sup>1</sup>

<sup>&</sup>lt;sup>1</sup>Roger Peng, An Introduction to the Interactive Debugging Tools in R, (2002)

# Problem Reporting in R



### Two kind of problems:

warnings do not halt the execution of a function.

"Something unusual happened during the execution of this function, but the function was nevertheless able to execute to completion."

errors are problems that are fatal and result in a complete halt in the execution, because the function simply cannot execute to completion due to the problem.

Problem Reporting in R

# Example:

```
> message <- function(x) {
+ if(x > 0)
+ print("Hello")
+ else
+ print("Goodbye")
+ }
> x < - \log(-1)
Warning message:
In log(-1): NaNs produced
> message(x)
Error in if (x > 0) print("Hello") else print("Goodbye") :
missing value where TRUE/FALSE needed
```

General remark: use robust code that checks for input errors.

The traceback function prints the list of functions which were called before the error occurred.

```
> x <- log(-1)
Warning message:
In log(-1) : NaNs produced
> message(x)
Error in if (x > 0) print("Hello") else print("Goodbye") :
missing value where TRUE/FALSE needed
> traceback()
1: message(x)
```

traceback shows in which function the error occurred. Since only one function was in fact called, this information is not very useful.



```
> f <- function(x) x - g(x)
> g <- function(y) y * h(y)
> h <- function(z) {
+ r <- log(z)
+ if (r < 10)
+ r^2
+ else r^3
+ }
> f(-10)
Error in if (r < 10) r^2 else r^3 : missing value where TRUE/FALSE needed
In addition: Warning message:
In log(z) : NaNs produced</pre>
```

Where did f(-10) fail?

```
> traceback()
3: h(y) at #1
2: g(x) at #1
1: f(-10)
```

traceback shows that the error occurred during evaluation of h(y).



```
> set.seed(100)
> xList <- as.list(rpois(1000, lambda = 5) - 1)</pre>
> lapply(xList, f)
Error in if (r < 10) r^2 else r^3 : missing value where TRUE/FALSE needed
In addition: Warning message:
In log(z): NaNs produced
```

Which element of xList caused the error?

```
> traceback()
4: h(y) at #1
3: g(x) at #1
2: FUN(X[[417L]], ...)
1: lapply(xList, f)
```

traceback shows that evaluation of element xList[[417]] caused the error. Only 0:8% of the list elements cause an error:

```
> which(unlist(xList)<0)</pre>
[1] 417 498 516 559 719 733 903 997
```

## Debugging Tools: debug



Where in the function h did the error occur? - Use debug to find out.

debug(h) ...

- ▶ allows to step through the function h line by line
- alters the way h is executed
- flags the function h for debugging

When a flagged function is called ...

- ▶ the body of the function is printed
- ▶ a *browser* command line opens in the console
- each statement in the function is executed one at a time
- ▶ the user can control when each statement gets executed
- the user interacts with the environment of the function

# Debugging Tools: debug - Example 2 contd.



What happens, if function h is called after flagging?

```
> debug(h)
> f(xList[[417]])
debugging in: h(y)
debug at #1: {
r < -log(z)
if (r < 10)
r^2
else r^3
Browse [2]>
```

Now, we can interact via the browser with the environment of the function:

```
Browse[2] > ls()
[1] "z"
Browse[2]> summary(z)
Min. 1st Qu. Median Mean 3rd Qu. Max.
-1 -1 -1 -1 -1
Browse[2] > str(z)
nıım -1
Browse [2]>
```

### Debugging Tools: debug



The four basic debugging commands inside the browser:

| c, cont | exit the browser and continue execution at the next statement           |
|---------|---|
| n       | enter the step-through debugger if the function is interpreted          |
| where   | print a stack trace of all active function calls                        |
| Q       | exit the browser and the current evaluation and return to the top-level |
|         | prompt  |

Besides the four basic debugging commands, all other R commands (including assignments) are allowed.

New objects are created in the local environment of the debugged function and will disappear when the debugger finishes.

If you have objects in your environment with the names n, c, or Q, then you must explicitly use the print function to print their values (i.e. print(n) or print(c)).



# browsing function h

```
Browse[2]> where
where 1 at #1: h(y)
where 2 at #1: q(x)
where 3: f(xList[[417]])
Browse[2]> n
debug at #2: r \leftarrow log(z)
Browse[2]> n
debug at #3: if (r < 10) r^2 else r^3
Browse[2] > ls()
[1] "r" "z"
Warning message:
    In log(z): NaNs produced
Browse[2]> r < 10
Γ17 NA
Browse[2]> n
Error in if (r < 10) r<sup>2</sup> else r<sup>3</sup> : missing value where TRUE/FALSE needed
```



# Manual debugging: explicit calls to browser

```
> h <- function(z) {
    + r < - log(z)
    + browser()
   + if (r < 10)
        + r^2
    + else r^3
    + }
> f(-10)
Called from: h(y)
Browse[1] > ls()
[1] "r" "z"
Warning message:
    In log(z): NaNs produced
Browse[1]> r
[1] NaN
```

Caution: do not forget to remove the call to browser() after debugging.

Modify code temporarily with trace: - trace makes minor modi cations to existing functions on the fly. - The traced functions are only modi ed indirectly without re-sourcing them. - Since base functions cannot be edited by the user, trace may be the only option available for making modifications.

```
> str(trace)
function (what, tracer, at, ...)
```

what name of function to be traced

tracer code to be inserted (name of function or unevaluated expression)

- at the position where the code will be inserted
- ... is for a lot more arguments available for trace, see ?trace



```
Let's use trace with function h:
```

```
```r
> h <- function(z) {
+ r < - log(z)
+ if (r < 10)
    + r^2
+ else r^3
+ }
> as.list(body(h))
[[1]]
.1.
[[2]]
r < -log(z)
[[3]]
if (r < 10) r^2 else r^3
> ## We can set a conditional break point using the if-statement
> trace(what = h, tracer = quote(if(is.nan(r)) browser()),
        + at = 3, print = FALSE)
[1] "h"
```

Debugging Tools: trace - Example 2 contd.

```
> h
Object with tracing code, class "functionWithTrace"
Original definition:
    function(z) {
        r < -log(z)
        if (r < 10)
            r^2
        else r<sup>3</sup>
# (to see the tracing code, look at body(object))
> body(h)
    r < -log(z)
         .doTrace(if (is.nan(r))
             browser(), "step 3")
        if (r < 10)
            r^2
        else r<sup>3</sup>
```

continued ...



# continued ...

```
> f(1)
[1] 1
> f(-10)
Called from: eval(expr, envir, enclos)
Browse[1] > ls()
[1] "r" "z"
Warning message:
   In log(z): NaNs produced
Browse[1]> Q
```

#### A call to untrace cancels the tracing:

```
> untrace(f)
> f(-10)
Error in if (r < 10) r<sup>2</sup> else r<sup>3</sup> : missing value where TRUE/FALSE needed
In addition: Warning message:
    In log(z): NaNs produced
```

The recover function helps in situations where you want to browse functions several functions in the stack:

```
> ## We set a conditional break point for the function recover
    > trace(what = h, tracer = quote(if(is.nan(r)) recover()),
            + at = 3, print = FALSE)
[1] "h"
> body(h)
    r < -log(z)
        .doTrace(if (is.nan(r))
            recover())
        if (r < 10)
            r^2
        else r<sup>3</sup>
```

# Debugging Tools: recover - Example 2 contd.



continued ...

```
> f(-10)
Enter a frame number, or 0 to exit
1: f(-10)
2: #1: a(x)
3: #1: h(y)
Selection: 2 ## Browse the g function
Called from: eval.parent(expr0bj)
Browse[1] > ls()
[1] "v"
Warning message:
    In log(z): NaNs produced
Browse[1]> y
Γ17 -10
Browse[1]> c
Enter a frame number, or 0 to exit
1: f(-10)
2: #1: g(x)
3: #1: h(y)
Selection: 0 ## Exit the recover function
Error in if (r < 10) r<sup>2</sup> else r<sup>3</sup> : missing value where TRUE/FALSE needed
```

Debugging Tools: recover - Example 3



**Problem**: how to browse functions inside other functions?

```
> f1 <- function(x)  {
    g1 <- function(y) {</pre>
        h1 <- function(z) {
           r < -log(z)
           if (r < 10) r^2 else r^3
        y * h1(y)
   x - g1(x)
> f1(-10)
Error in if (r < 10) r^2 else r^3 ...
> traceback()
3: h1(y) at #7
> trace(what = h1, tracer = quote(if(is.nan(r)) recover()),
+ at = 3, print = FALSE)
Error in methods::.TraceWithMethods(what = h1, tracer = quote(if (is.nan(object |
```



# More general approach to use recover:

```
> options()$error
NULL
options(error = recover)
f1(-10)
Error in if (r < 10) r^2 else r^3 : missing value where TRUE/FALSE needed
In addition: Warning message:
In log(z) : NaNs produced
Enter a frame number, or 0 to exit
1: f1(-10)
2: #9: g1(x)
3: #7: h1(y)
Selection:</pre>
```

Debugging Tools: recover - Example 3

# Even more general approach to use recover to treat warnings:

```
> options()$warn
[1] 0
> options(warn = 2)
> f1(-10)
Error in log(z): (converted from warning) NaNs produced
Enter a frame number, or 0 to exit
1: f1(-10)
2: #9: q1(x)
3: #7: h1(u)
4: #4: .signalSimpleWarning("NaNs produced", quote(log(z)))
5: withRestarts({
.Internal(.signalCondition(simpleWarning(msg, call), msg, call))
.Internal(.dfltWar
6: withOneRestart(expr, restarts[[1]])
7: doWithOneRestart(return(expr), restart)
Selection:
```

# Debugging in R



# Summary

traceback prints the call stack of the last uncaught error

debug flags a function for debugging

browser interrupts the execution of an expression and allows the inspection of

the environment

trace allows to insert debugging code into an existing function

recover allows to browse directly on **any** of the currently active function calls;

with options(warn=2, error=recover) warnings can be debugged

as well