FRE6871 R in Finance Lecture#1, Spring 2023

Jerzy Pawlowski jp3900@nyu.edu

NYU Tandon School of Engineering

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Welcome Students!

My name is Jerzy Pawlowski jp3900@nyu.edu

I'm an adjunct professor at NYU Tandon because I love teaching and I want to share my professional knowledge with young, enthusiastic students.

I'm interested in applications of *machine learning* to *systematic investing*.

I'm an advocate of *open-source software*, and I share it on GitHub:

My GitHub account

In my finance career, I have worked as a hedge fund portfolio manager, CLO structurer (banker), and quant analyst.

My LinkedIn profile



NYU

OF

Jerzy Pawlowski

Adjunct Professor at NYU Tandon School of Engineering
Greater New York City Area

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Interested in applications of machine learning and high frequency data to systematic investing.



FRE6871 Course Description and Objectives

Course Description

The course will study the applications of the R statistical language to financial data analysis and modeling. The applications will include *classification* for credit scoring, *Monte Carlo simulation* for option pricing and credit portfolio modeling, and *Principal Component Analysis* (*PCA*) for interest rate yield curve modeling. The course will apply statistical techniques, such as *hypothesis testing*. *linear regression*, *logistic regression*, and *bootstrap simulation*.

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Course Objectives

Students will learn through R coding exercises how to:

- Manipulate data structures (vectors, data frames, dates, and time series).
- Download data from external sources, and to scrub and format it.
- Create interactive plots and visualizations.
- Build financial models.
- Perform exception and error handling, and debugging.

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Course Prerequisites

The R language is considered to be challenging, so this course requires some programming experience with other languages such as C++ or Python. Students should also have knowledge of basic statistics (random variables, estimators, hypothesis testing, regression, etc.) The course FRE7241 Algorithmic Portfolio Management is designed as a followup course to FRE6871.

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Homeworks and Tests

Homeworks and Tests

Grading will be based on homeworks and tests. There will be no final exam.

The tests will require writing code, which should run directly when pasted into an R session, and should produce the required output, without any modifications.

Students will be allowed to consult lecture slides, and to copy code from them, and to copy from books or any online sources, but they will be required to provide references to those external sources (such as links or titles and page numbers).

The tests will be closely based on code contained in the lecture slides, so students are encouraged to become very familiar with those slides.

Students will submit their homework and test files only through Brightspace (not emails).

Students will be required to bring their laptop computers to class and run the R Interpreter, and the RStudio Integrated Development Environment (*IDE*), during the lecture.

Homeworks will also include reading assignments designed to help prepare for tests.

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Graduate Assistant

The graduate assistant (GA) will be announced.

The GA will answer questions during office hours, or via *Brightspace* forums, not via emails. Please send emails regarding lecture matters from *Brightspace* (not personal emails).

Tips for Solving Homeworks and Tests

Tips for Solving Homeworks and Tests

The tests will require mostly copying code samples from the lecture slides, making some modifications to them, and combining them with other code samples.

Partial credit will be given even for code that doesn't produce the correct output, but that has elements of code that can be useful for producing the right answer.

So don't leave test assignments unanswered, and instead copy any code samples from the lecture slides that are related to the solution and make sense.

Contact the GA during office hours via text or phone, and submit questions to the GA or to me via Brightspace.

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Please Submit Minimal Working Examples With Your Questions

When submitting questions, please provide a *minimal working example* that produces the error in R, with the following items:

- The complete R code that produces the error, including the seed value for random numbers,
- The version of R (output of command: sessionInfo()), and the versions of R packages,
- The type and version of your operating system (Windows or OSX),
- The dataset file used by the R code,
- The text or screenshots of error messages,

You can read more about producing $minimal\ working\ examples$ here: http://stackoverflow.com/help/mcve http://www.jaredknowles.com/journal/2013/5/27/writing-a-minimal-working-example-mwe-in-r

Course Grading Policies

Numerical Scores

Homeworks and tests will be graded and assigned numerical scores. Each part of homeworks and tests will be graded separately and assigned a numerical score.

Maximum scores will be given only for complete code, that produces the correct output when it's pasted into an R session, without any modifications. As long as the R code uses the required functions and produces the correct output, it will be given full credit.

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Letter Grades

Letter grades for the course will be derived from the cumulative scores obtained for all the homeworks and tests. Very high numerical scores close to the maximum won't guarantee an A letter grade, since grading will also depend on the difficulty of the assignments.

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Plagiarism

Plagiarism (copying from other students) and cheating will be punished.

But copying code from lecture slides, books, or any online sources is allowed and encouraged.

Students must provide references to any external sources from which they copy code (such as links or titles and page numbers).

FRE6871 Course Materials

Lecture Slides

The course will be mostly self-contained, using detailed lecture slides containing extensive, working R code examples.

The course will also utilize data and tutorials which are freely available on the internet.

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FRE6871 Recommended Textbooks

- Statistics and Data Analysis for Financial Engineering by David Ruppert, introduces regression, cointegration, multivariate time series analysis, ARIMA, GARCH, CAPM, and factor models, with examples in R.
- Quantitative Risk Management by Alexander J. McNeil, Rudiger Frey, and Paul Embrechts: review of Value at Risk, factor models, ARMA and GARCH, extreme value theory, and credit risk models.
- Introduction to Statistical Learning by Gareth James, Daniela Witten, Trevor Hastie, and Robert Tibshirani, introduces machine learning techniques using R, but without deep learning.
- Advanced R by Hadley Wickham, is the best book for learning the advanced features of R.
- The Art of R Programming by Norman Matloff, contains a good introduction to R and to some statistical models.

Many textbooks can be downloaded in electronic format from the NYU Library.

FRE6871 Supplementary Textbooks

Supplementary Textbooks

- The books *R* in Action by Robert Kabacoff and *R* for Everyone by Jared Lander, are good introductions to *R* and to statistical models.
- Applied Econometrics with R by Christian Kleiber and Achim Zeileis, introduces advanced statistical models and econometrics.
- Numerical Recipes in C++ by William Press, Saul Teukolsky, William Vetterling, and Brian Flannery, is a
 great reference for linear algebra and numerical methods, implemented in working C++ code.

FRE6871 Supplementary Materials

Notepad++ is a free source code editor for MS Windows, that supports several programming languages, including R.

Notepad++ has a very convenient and fast search and replace function, that allows search and replace in multiple files.

http://notepad-plus-plus.org/



Internal R Help and Documentation

The function help() displays documentation on a function or subject,

Preceding the keyword with a single "?" is equivalent to calling help().

- > # Display documentation on function "getwd"
- > help(getwd)
- > # Equivalent to "help(getwd)"
- > ?getwd

The function help.start() displays a page with links to internal documentation.

 ${\tt R}$ documentation is also available in RGui under the help tab.

The *pdf* files with R documentation are also available directly under:

C:/Program Files/R/R-3.1.2/doc/manual/ (the exact path will depend on the R version.)

> # Open the hypertext documentation

> help.start()



Introduction to R by Venables and R Core Team.

R Online Help and Documentation

R Cheat Sheets

Cheat Sheets are a fast way to find what you want

https://www.rstudio.com/resources/cheatsheets/

R Programming Wikibook

Wikibooks are crowdsourced textbooks

 $http://en.wikibooks.org/wiki/R_Programming/$

R FAQ

Frequently Asked Questions about R

http://cran.r-project.org/doc/FAQ/R-FAQ.html

R-seek Online Search Tool

R-seek allows online searches specific to the R language

http://www.rseek.org/

R-help Mailing List

R-help is a very comprehensive Q&A mailing list

https://stat.ethz.ch/mailman/listinfo/r-help

R-help has archives of past Q&A - search it before you ask

https://stat.ethz.ch/pipermail/r-help/

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R Code Style Guidelines

Please follow the RStudio Style Guide to make your R code more readable.

Please also follow these R code style rules:

- Use the left arrow "<-" for assignment, not the equals sign "=" (to insert "<-" into code, use the *Alt-hyphen* shortcut in Windows),
- Use nouns for variable names and verbs for function names,
- Use a combination of lowercase letters, numbers, and underscores "_" for names of variables and functions,
- Add underscores "_" to names to avoid conflicts with the names of existing R functions and variables,
- Do not use dots "." in names, except in the names of function *methods* (even though R uses them for variables as well).
- Use underscores "_" in file names, instead of spaces,
- Always put a space after a comma, never before it: "x, y" not "x , y",
- Do not put spaces inside or outside parentheses: "if (x > 0)" not "if (x > 0)",
- Surround infix operators (==, +, -, <-, etc.) with spaces: "x > 0" not "x>0" (even though I don't always follow that rule, to save whitespace),
- Add a comment after the closing curly bracket: "} # end my_fun",

You can reformat R code chunks using the *styler* macros in the *RStudio Addins* drop-down menu. You can also reformat whole files with R code by using the *styler* package.

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Stack Exchange

Stack Overflow

Stack Overflow is a Q&A forum for computer programming, and is part of Stack Exchange

http://stackoverflow.com

http://stackoverflow.com/questions/tagged/r

http://stackoverflow.com/tags/r/info

Stack Exchange

Stack Exchange is a family of Q&A forums in a variety of fields

http://stackexchange.com/

http://stackexchange.com/sites#technology

http://quant.stackexchange.com/



RStudio Support

RStudio has extensive online help, Q&A database, and documentation

https://support.rstudio.com/hc/en-us

https://support.rstudio.com/hc/en-us/sections/200107586-Using-RStudio

https://support.rstudio.com/hc/en-us/sections/200148796-Advanced-Topics

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R Online Books and References

Hadley Wickham book Advanced R

The best book for learning the advanced features of R: http://adv-r.had.co.nz/

Cookbook for R by Winston Chang from RStudio

Good plotting, but not interactive: http://www.cookbook-r.com/

Efficient R programming by Colin Gillespie and Robin Lovelace

Good tips for fast R programming: https://csgillespie.github.io/efficientR/programming.html

Endmemo web book

Good, but not interactive: http://www.endmemo.com/program/R/

Quick-R by Robert Kabacoff

Good, but not interactive: http://www.statmethods.net/

R for Beginners by Emmanuel Paradis

Good, basic introduction to R: http://cran.r-project.org/doc/contrib/Paradis-rdebuts_en.pdf

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R Online Interactive Courses

Datacamp Interactive Courses

Datacamp introduction to R: https://www.datacamp.com/courses/introduction-to-r/

Datacamp list of free courses: https://www.datacamp.com/community/open-courses

Datacamp basic statistics in R: https://www.datacamp.com/community/open-courses/basic-statistics

Datacamp computational finance in R:

https://www.datacamp.com/community/open-courses/computational-finance-and-financial-econometrics-with-research and the computational finance and financial-econometrics and the computational finance and the computational finance and financial-econometrics and the computational finance and the computation f

Datacamp machine learning in R:

https://www.datacamp.com/community/open-courses/kaggle-r-tutorial-on-machine-learning and the state of the

Try R

Interactive R tutorial, but rather basic: http://tryr.codeschool.com/

R Blogs and Experts

R-Bloggers

R-Bloggers is an aggregator of blogs dedicated to R

http://www.r-bloggers.com/

Tal Galili is the author of R-Bloggers and has his own excellent blog

http://www.r-statistics.com/

Dirk Eddelbuettel

Dirk is a *Top Answerer* for R questions on Stackoverflow, the author of the Rcpp package, and the CRAN Finance View

http://dirk.eddelbuettel.com/

http://dirk.eddelbuettel.com/code/

http://dirk.eddelbuettel.com/blog/

http://www.rinfinance.com/

Romain Frangois

Romain is an R Enthusiast and Rcpp Hero

http://romainfrancois.blog.free.fr/

http://romainfrancois.blog.free.fr/index.php?tag/graphgallery

http://blog.r-enthusiasts.com/

More R Blogs and Experts

Revolution Analytics Blog

R blog by Revolution Analytics software vendor ${\tt http://blog.revolutionanalytics.com/}$

RStudio Blog

R blog by *RStudio* http://blog.rstudio.org/

GitHub for Hosting Software Projects Online

GitHub is an internet-based online service for hosting repositories of software projects.

 $\it Git Hub$ provides version control using $\it git$ (desved by Linus Torvalds).

Most R projects are now hosted on GitHub.

Google uses GitHub to host its tensorflow library for machine learning:

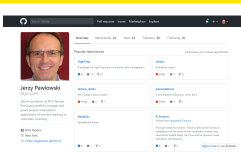
https://github.com/tensorflow/tensorflow

All the FRE-7241 and FRE-6871 lectures are hosted on GitHub:

 $https://github.com/algoquant/lecture_slides$

https://github.com/algoquant

Hosting projects on *Google* is a great way to advertize your skills and network with experts.



What is R?

- An open-source software environment for statistical computing and graphics.
- An interpreted language, that allows interactive code development.
- A functional language where every operator is an R function.
- A very expressive language that can perform complex operations with very few lines of code.
- A language with metaprogramming facilities that allow programming on the language.
- A language written in C/C++, which can easily call other C/C++ programs.
- Can be easily extended with packages (function libraries), providing the latest developments like Machine Learning.
- Supports object-oriented programming with *classes* and *methods*.
- Vectorized functions written in C/C++, allow very fast execution of loops over vector elements.





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Why is R More Difficult Than Other Languages?

 ${\tt R}$ is more difficult than other languages because:



- R is a functional language, which makes its syntax unfamiliar to users of procedural languages like C/C++.
 The huge number of user-created packages makes it difficult to tell
- which are the best for particular applications.

 R can produce very cryptic warning and error messages, because it's a
- programming environment, so it performs many operations quietly, but those can sometimes fail.
- Fixing errors usually requires analyzing the complex structure of the R programming environment.

This course is designed to teach the most useful elements of R for financial analysis, through case studies and examples,

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What are the Best Ways to Use R?

If used properly, R can be fast and interactive:

- Use R as an interface to libraries written in C++, Java, and JavaScript.
- Avoid using too many R function calls (every command in R is a function).
- Avoid using apply() and for() loops for large datasets.
- Use R functions which are compiled C++ code, instead of using interpreted R code.
- Use package data.table for high performance data management.
- Use package shiny for interactive charts of live models running in R.
- Use package dygraphs for interactive time series plots.
- Use package knitr for RMarkdown documents.
- Pre-allocate memory for new objects.
- Write C++ functions in Rcpp and RcppArmadillo.



- > # Calculate cumulative sum of a vector
- > vectorv <- runif(1e5)
- > # Use compiled function
 > cumsumv <- cumsum(vectorv)</pre>
- > # Use for loop
- > cumsumv2 <- vectorv
- > for (i in 2:NROW(vectorv))
 + cumsumv2[i] <- (cumsumv2[i] + cumsumv2[i-1])</pre>
- > # Compare the two methods
- > all.equal(cumsumv, cumsumv2)
 > # Microbenchmark the two methods
- > # Microbenchmark the two method
- > library(microbenchmark)
 > summary(microbenchmark(
- + cumsum=cumsum(vectorv),
- + loop=for (i in 2:NROW(vectorv))
- vectorv[i] <- (vectorv[i] + vectorv[i-1]),</pre>
- + times=10))[, c(1, 4, 5)]

The R License

 ${\tt R}$ is open-source software released under the GNU General Public License:

http://www.r-project.org/Licenses



Some other \Re packages are released under the Creative Commons Attribution-ShareAlike License:



http://creativecommons.org

Installing R and RStudio

Students will be required to bring their laptop computers to all the lectures, and to run the R Interpreter and RStudio RStudio during the lecture,

Laptop computers will be necessary for following the lectures, and for performing tests,

Students will be required to install and to become proficient with the R Interpreter,



Students can download the R Interpreter from CRAN (Comprehensive R Archive Network):

http://cran.r-project.org/

To invoke the RGui interface, click on:

C:/Program Files/R/R-3.1.2/bin/x64/RGui.exe

Students will be required to install and to become proficient with the *RStudio* Integrated Development Environment (*IDE*),





Using RStudio

```
RStudio
File Edit Code View Plots Session Project Build Tools Help
O • Go to file/function
                                                                                                            Workspace History
 (2) Untitled1" x (3) alphaScripts.R x (4) FRE6811_Lecture_1.Rnw x (4) prototype.Rnw" x (5) knitr_presentation_demo.Rnw x
 Run > Source -

☐ To Console ☐ To Source 
☐ 

  2087 # Run quasi-CEP mode
                                                                                                            22MASS
  2088 cep.ticks <- 0:100 # number of ticks cut off from tail
                                                                                                            installed.packages()
  2089 n.buffer <- 500 # buffer size of ticks fed into model
                                                                                                            packageDescription("MASS")
  2090 model.cep <- model.test
                                                                                                            ?unloadNamespace
  2091 ts.prices <- model.testSprices
                                                                                                            ?library
  2092 cep.signals <- sapply(cep.ticks, function(cep.tick)
                                                                                                            2data
  2093 -
                                                                                                            install.packages("PerformanceAnalytics", repos="http://R-Forge.R-project
  2094
                                cep.prices <- tail(last(ts.prices,-cep.tick), n.buffer)</pre>
                                model.cep <- update.alphaModel(model=model.cep, ts.prices=cep.prices)
  2096
                                model.cep <- recalc.alphaModel(model.cep)
                                                                                                            R. HOME
  2097
                                as.vector(last(model.cep$signals))
                                                                                                            R. home
  2008
                                                                                                            R. home ("home")
  2000
                                                                                                            R. home()
  2100 write.csv(cep.signals, "S:/Data/R_Data/signals.cep.csv")
  2101
        write.csv(model.test$signals, "5:/Data/R_Data/signals.csv")
                                                                                                            ?Startup
                                                                                                                Plots Packages Help
  2105 ### Portfolio Optimization ###
                                                                                                            R: Loading and Listing of Packages * Find in Top
  2107 library(DEoptim)
  2108
                                                                                                             library (base)
  2109 ### Load data
  2110 stock.sectors.prices <- read.csv(paste(alpha.dir, "stock_sectors.csv", sep=""), stringsAsFactors
                                                                                                            Loading and Listing of Packages
  2111 stock.sectors.prices <- xts(stock.sectors.prices[,-1], order.by=as.POSIXIt(stock.sectors.prices[
  2112 ts.rets <- diff(stock.sectors.prices,lag=1)
  2113 ts.rets[1,] <- ts.rets[2,]
                                                                                                            Description
       (
 2113:1 [3] (Untitled) 0
                                                                                                            library and require load add-on packages
 Console Compile PDF ×
                                                                                                            Usage
 C:/Develop/R/Presentations/ @
 Warning in install.packages :
                                                                                                            library(package, help, pos = 2, lib.loc = NULL,
  InternetOpenUrl failed: 'A connection with the server could not be established'
                                                                                                                    character.only = FALSE, logical.return = FALSE,
 warning in install.packages :
                                                                                                                    warn.conflicts = TRUE, quietly = FALSE,
  InternetOpenurl failed: 'A connection with the server could not be established'
                                                                                                                    verbose = getOption("verbose"))
 warning in install.packages :
  unable to access index for repository http://www.stats.ox.ac.uk/pub/RWin/bin/windows/contrib/3.0
                                                                                                            require(package, lib.loc = NULL, quietly = FALSE,
 Installing package into 'C:/Users/Jerzy/Documents/R/win-library/3.0'
                                                                                                                    warn.conflicts = TRUE,
 (as 'lib' is unspecified)
trying URL 'http://R-Forge.R-project.org/bin/windows/contrib/3.0/PerformanceAnalytics_1.1.2.zip'
                                                                                                                    character.only = FALSE)
Content type 'application/zip' length 2205138 bytes (2.1 Mb)
opened URL
                                                                                                            Arguments
 downloaded 2.1 Mb
                                                                                                             package, help the name of a package, given as a name or literal character string, or a character
                                                                                                                           december of the second
```

A First R Session

Variables are created by an assignment operation, and they don't have to be declared.

The standard assignment operator in $\tt R$ is the arrow symbol "<-".

R interretsp text in quotes ("") as character strings.

Text that is not in quotes ("") is interpreted as a *symbol* or *expression*.

Typing a symbol or expression evaluates it.

R uses the hash "#" sign to mark text as comments.

All text after the hash "#" sign is treated as a comment, and is not executed as code.

```
> # "<-" and "=" are valid assignment operators
> myvar <- 3
> # typing a symbol or expression evaluates it
> myvar
[1] 3
> # text in quotes is interpreted as a string
> myvar <- "Hello World!"
> # typing a symbol or expression evaluates it
> myvar
[1] "Hello World!"
> myvar # text after hash is treated as comment
[1] "Hello World!"
```

Exploring an R Session

The function getwd() returns a vector of length 1, with the first element containing a string with the name of the current working directory (cwd).

The function setwd() accepts a character string as input (the name of the directory), and sets the working directory to that string.

R is a functional language, and R commands are functions, so they must be followed by parentheses "()".

```
> getwd() # get cwd
> setwd("/Users/jerzy/Develop/R") # Set cwd
> getwd() # get cwd
```

```
Get system date and time
```

Just the date

```
> Sys.time() # get date and time
[1] "2023-03-20 17:34:35 EDT"
>
> Sys.Date() # get date only
[1] "2023-03-20"
```

The R Workspace

The workspace is the current R working environment, which includes all user-defined objects and the command history.

The function ls() returns names of objects in the R workspace.

The function rm() removes objects from the R workspace.

The workspace can be saved into and loaded back from an .RData file (compressed binary file format).

The function save.image() saves the whole workspace.

The function save() saves just the selected objects.

The function load() reads data from .RData files, and invisibly returns a vector of names of objects created in the workspace.

```
> var1 <- 3  # Define new object
> list all objects in workspace
> # List objects starting with "v"
> ls(pattern=glob2rx("v*"))
> # Remove all objects starting with "v"
> rm(list=ls(pattern=glob2rx("v*")))
> save.inage() # Save workspace to file .RData in cwd
> rm(var1)  # Remove object
> ls() # List objects
> load(".RData")
> ls() # List objects
> var2 <- 5  # Define another object
> save(var1, var2, # Save selected objects
+ file="Wusers/ierry/DevelopOrlecture_slides/data/my_data.RData
```

> loadv <- load(file="/Users/jerzy/Develop/lecture_slides/data/my_data

> rm(list=ls()) # Remove all objects

> ls() # List objects

> ls() # List objects

> loady

The R Workspace (cont.)

When you quit R you'll be prompted "Save workspace image?"

If you answer YES then the workspace will be saved into the .RData file in the cwd,

When you start R again, the workspace will be automatically loaded from the existing .RData file, $\label{eq:loaded}$

q() # quit R session

The function history() displays recent commands,

You can also save and load the command history from a file.

> history(5) # Display last 5 commands

> savehistory(file="myfile") # Default is ".Rhistory"

> loadhistory(file="myfile") # Default is ".Rhistory"

R Session Info

The function sessionInfo() returns information about the current R session.

- R version.
- OS platform,
- locale settings.
- list of packages that are loaded and attached to the search path.
- list of packages that are loaded, but not attached to the search path,

> sessionInfo() # get R version and other session info R version 4.2.1 (2022-06-23) Platform: aarch64-apple-darwin20 (64-bit) Running under: macOS Monterey 12.5.1

Matrix products: default

BLAS: /Library/Frameworks/R.framework/Versions/4.2-arm64/Resource LAPACK: /Library/Frameworks/R.framework/Versions/4.2-arm64/Resource

locale.

[1] en US.UTF-8/en US.UTF-8/en US.UTF-8/C/en US.UTF-8/en US.UTF-8

attached base packages:

datasets stats

[1] graphics grDevices utils other attached packages:

[1] knitr 1.40 HighFreq 0.1

rutils 0.2 dygraphs_1.1 [5] quantmod 0.4.20 TTR 0.24.3 xts 0.12.1 zoo 1.8-10

highr_0.9

xfun 0.32

lifecycle_1.0.3

evaluate_0.16

loaded via a namespace (and not attached): magrittr 2.0.3

[1] Rcpp_1.0.9 rstudioapi_0.13 [5] rlang_1.0.6 fastmap_1.1.0 [9] tools 4.2.1 grid 4.2.1

[13] htmltools_0.5.3 digest_0.6.29 [17] vctrs_0.5.2 curl_4.3.2

[21] stringi_1.7.8 compiler_4.2.1 methods

lattice_

stringr_

cli 3.6.

htmlwidg

glue_1.6

Environment Variables

R uses environment variables to store information about its environment, such as paths to directories containing files used by R (startup, history, OS).

For example the environment variables:

- R_USER and HOME store the R user Home directory,
- R_HOME stores the root directory of the R installation,

The functions Sys.getenv() and Sys.setenv() display and set the values environment variables.

Sys.getenv("env_var") displays the environment variable "env_var".

Sys.setenv("env_var=value") sets the environment variable "env_var" equal to "value".

```
> Sys.getenv()[5:7] # List some environment variables
> 
> Sys.getenv("HOME") # get R user HOME directory
> Sys.setenv(Home="/Users/jerzy/Develop/data") # Set HOME directory
> Sys.getenv("HOME") # get user HOME directory
> Sys.getenv("R_HOME") # get R_HOME directory
> R.home() # get R_HOME directory
> R.home() # get R_HOME directory
> R.home("etc") # get "etc" sub-directory of R_HOME
```

> optionv <- options()

> options(optionv)

> # Restore all options from variable

Global Options Settings

R uses a list of global options which affect how R computes and displays results.

The function options() either sets or displays the values of global *options*.

options("globop") displays the current value of option "globop".

getOption("globop") displays the current value of option "globop".

options(globop=value) sets the option "globop" equal to "value".

```
> # ?options # Long list of global options
> # Interpret strings as characters, not factors
> getOption("stringsAsFactors") # Display option
> options("stringsAsFactors") # Display option
> options(stringsAsFactors=FALSE) # Set option
> # number of digits printed for numeric values
> options(digits=3)
> # control exponential scientific notation of print method
> # positive "scipen" values bias towards fixed notation
> # negative "scipen" values bias towards scientific notation
> options(scipen=100)
> # maximum number of items printed to console
> options(max.print=30)
> # Warning levels options
> # negative - warnings are ignored
> options(warn=-1)
> # zero - warnings are stored and printed after top-confl function
> options(warn=0)
> # One - warnings are printed as they occur
> options(warn=1)
> # two or larger - warnings are turned into errors
> options(warn=2)
> # Save all options in variable
```

Constructing File Paths

Names of *file paths* can be constructed using the function paste().

The function file.path() is similar to paste(), but it also automatically uses the correct file separator for the computer platform.

The function normalizePath() performs tilde-expansions and displays file paths in user-readable format.

```
> # R startup (site) directory
> paste(R.home(), "etc", sep="/")
[i] "/Library/Frameworks/R.framework/Resources/etc"
> file.path(R.home(), "etc") # better way
[i] "/Library/Frameworks/R.framework/Resources/etc"
> # perform tilde-expansions and convert to readable format
> normalizePath(file.path(R.home(), "etc"), winslash="/")
[i] "/Library/Frameworks/R.framework/Versions/4.2-arm64/Resources/e
> normalizePath(R.home("etc"), winslash="/")
```

[1] "/Library/Frameworks/R.framework/Versions/4.2-arm64/Resources/e

R System Directories under Windows

R uses several different directories to search, read, and store files:

- Windows user personal directory: "~" ("%USERPROFILE%/Documents"),
- R user HOME directory (R_USER and Home),
- cwd current working directory the default directory for storing and retrieving user files (such as .Rhistory, .RData, etc.),
- R_HOME root directory of the R installation,
- R startup (site) directory: R_HOME/etc/,

By default, the R user HOME directory is the *Windows* user personal directory.

The cwd is set to the directory from which R is invoked, or the R user HOME directory.

File and Directory Listing Functions

The functions list.files() and dir() return a vector of names of files in a given directory.

The function list.dirs() listv the directories in a given directory.

The function Sys.glob() listv files matching names obtained from wildcard expansion.

```
> sample(dir(), 5) # get 5 file names - dir() listv all files
> sample(dir(pattern="csv"), 5) # List files containing "csv"
> sample(list.files(R.home()), 5) # All files in R.HOME directory
> sample(list.files(R.home("etc")), 5) # All files in "etc" sub-directory
> sample(list.dirs(), 5) # Directories in cwd
> list.dirs(R.home("etc")) # Directories in "etc" sub-directory
> sample(Sys.glob("*.csv"), 5)
> Sys.glob(R.home("etc"))
```

Invoking an R Session in Windows

An R session can run in several different ways:

- In an R terminal (by invoking R.exe or Rterm.exe),
- In an R RGui (by invoking RGui.exe),
- In an RStudio session (or some other IDE),

The initial value of the cwd depends on how the R session is invoked.

If R is invoked:

- from the Windows menu, then cwd is set to the R user HOME directory,
- by clicking on a file (*.R, .RData, etc.), then cwd is set to the file's directory,
- by typing R.exe or Rterm.exe in the command shell (after setting the PATH), then cwd is set to the directory where the command was typed,

> getwd() # get cwd
[1] "/Users/jerzy/Develop/lecture_slides"

R Session Startup

At startup ${\tt R}$ sources (reads) several types of files, in the following order:

- Renviron files defining environment variables,
- Rprofile files containing code executed at R startup,
- RData files containing data to be loaded at R startup,

R sources files from several directories, in the following order:

- R startup directory: Renviron.site and Rprofile.site files,
- cwd directory: .Renviron, .Rprofile, and .RData files,
- HOME user directory (only if no files found in cwd),

The above startup process can be customized by setting environment variables.

```
> # help(Startup) # Description of R session startup mechanism
> # files in R startup directory directory
> dir(normalizePath(file.path(R.home(), "etc"), winslash="/"))
> 
> # *.R* files in cwd directory
> getwd()
> dir(getwd(), all.files=TRUE, pattern="\\.R")
> dir(getwd(), all.files=TRUE, pattern=glob2rx("*.R*"))
```

Data Objects in R

All data objects in R are vectors, or consist of vectors.

Single numbers and character strings are vectors of length "1".

Atomic vectors are homogeneous objects whose elements are all of the same mode (type).

Lists and data frames are recursive (heterogeneous) objects, whose elements can be vectors of different mode

The functions is.atomic() and is.recursive() return logical values depending on whether their arguments are atomic or recursive.

R Data Objects

	Atomic	Recursive
1-dim	Vectors	Lists
2-dim	Matrices	Data frames
n-dim	Arrays	NA

```
> # Single numbers are vectors of length 1
> 1
Γ17 1
> # Character strings are vectors of length 1
> "a"
[1] "a"
> # Strings without quotes are variable names
> a # Variable "a" doesn't exist
Error in eval(expr, envir, enclos): object 'a' not found
> # List elements can have different mode
> list(aa=c("a", "b"), bb=1:5)
$aa
[1] "a" "b"
$bb
[1] 1 2 3 4 5
> data.frame(aa=c("a", "b"), bb=1:2)
  aa bb
1 a 1
> is.atomic(data.frame(aa=c("a", "b"), bb=1:2))
[1] FALSE
> is.recursive(data.frame(aa=c("a", "b"), bb=1:2))
[1] TRUE
```

Type, Mode, and Class of Objects

The *type*, *mode*, and *class* are character strings representing various object properties.

The *type* of an atomic object represents how it's stored in memory (logical, character, integer, double, etc.)

The *mode* of an atomic object is the kind of data it represents (logical, character, numeric, etc.)

The *mode* of an object often coincides with its *type* (except for integer and double types).

Recursive objects (listv, data frames) have both type and mode equal to the recursive type (list).

The *class* of an object is given by either an explicit *class* attribute, or is derived from the object *mode* and its dim attribute (implicit *class*).

The function class() returns the explicit or implicit class of an object.

The object *class* is used for method dispatching in the S3 object-oriented programming system in R.

```
> mvvar <- "hello"
> c(typeof(myvar), mode(myvar), class(myvar))
[1] "character" "character" "character"
> mvvar <- 1:5
> c(typeof(myvar), mode(myvar), class(myvar))
[1] "integer" "numeric" "integer"
> myvar <- runif(5)
> c(typeof(myvar), mode(myvar), class(myvar))
[1] "double" "numeric" "numeric"
> myvar <- matrix(1:10, 2, 5)
> c(typeof(myvar), mode(myvar), class(myvar))
[1] "integer" "numeric" "matrix" "array"
> myvar <- matrix(runif(10), 2, 5)
> c(typeof(myvar), mode(myvar), class(myvar))
[1] "double" "numeric" "matrix" "array"
> mvvar <- list(aa=c("a", "b"), bb=1:5)
> c(typeof(myvar), mode(myvar), class(myvar))
[1] "list" "list" "list"
> myvar <- data.frame(aa=c("a", "b"), bb=1:2)
> c(typeof(myvar), mode(myvar), class(myvar))
[1] "list"
                 "list"
                              "data.frame"
```

R Object Attributes

R objects can have different attributes, such as: namesv, dimnames, dimensions, class, etc.

The attributes of an object is a named list of symbol=value pairs.

The function attributes() returns the attributes of an object.

The attributes of an R object can be modified using the "attributes() <- " assignment.

The function structure() adds attributes (specified as symbol=value pairs) to an object, and returns it.

A vector that is assigned an attribute other than namesy is not treated as a vector.

The function is.vector() returns TRUE if its argument is a vector, and returns FALSE otherwise.

```
> # A simple vector has no attributes
> attributes(5:10)
NUIT.T.
> myvar <- c(pi=pi, euler=exp(1), gamma=-digamma(1))
> # Named vector has "namesy" attribute
> attributes(myvar)
$names
[1] "pi"
          "euler" "gamma"
> myvar <- 1:10
> is.vector(myvar) # Is the object a vector?
[1] TRUE
> attributes(myvar) <- list(my_attr="foo")
> myvar
 [1] 1 2 3 4 5 6 7 8 9 10
attr(,"my_attr")
[1] "foo"
> is.vector(myvar) # Is the object a vector?
[1] FALSE
> myvar <- 0
> attributes(myvar) <- list(class="Date")
> mvvar # "Date" object
[1] "1970-01-01"
> structure(0, class="Date") # "Date" object
```

[1] "1970-01-01"

Modifying class Attributes

Objects without an explicit class don't have a class attribute, and the function class() returns the implicit class.

The class of an object can be modified using the "class() <- " assignment.

An object can have a main *class*, and also an inherited class (the class attribute can be a vector of strings).

The function unclass() removes the explicit class attribute from an object.

```
> myvar <- matrix(runif(10), 2, 5)
> class(myvar) # Has implicit class
[1] "matrix" "array"
> # But no explicit "class" attribute
> attributes(mvvar)
$dim
Γ17 2 5
> c(typeof(myvar), mode(myvar), class(myvar))
[1] "double" "numeric" "matrix" "array"
> # Assign explicit "class" attribute
> class(mvvar) <- "mv class"
> class(mvvar) # Has explicit "class"
[1] "my_class"
> # Has explicit "class" attribute
> attributes(myvar)
$dim
Γ17 2 5
$class
[1] "mv class"
> is.matrix(myvar) # Is the object a matrix?
[1] TRUE
> is.vector(mvvar) # Is the object a vector?
[1] FALSE
> attributes(unclass(myvar))
$dim
[1] 2 5
```

Implicit Class of Objects

If an object has no explicit class, then its implicit class is derived from its mode and dim attribute (except for integer vectors which have the implicit class "integer" derived from their type).

If an atomic object has a dim attribute, then its implicit class is matrix or array.

Data frames have no explicit dim attribute, but dim() returns a value, so they have an implicit dim attribute.

```
> # Integer implicit class derived from type
> myvar <- vector(mode="integer", length=10)
> c(typeof(myvar), mode(myvar), class(myvar))
[1] "integer" "numeric" "integer"
> # Numeric implicit class derived from mode
> myvar <- vector(mode="numeric", length=10)
> c(typeof(myvar), mode(myvar), class(myvar))
[1] "double" "numeric" "numeric"
> # Adding dim attribute changes implicit class to matrix
> dim(myvar) <- c(5, 2)
> c(typeof(myvar), mode(myvar), class(myvar))
[1] "double" "numeric" "matrix" "array"
> # Data frames have implicit dim attribute
> myvar <- data.frame(aa=c("a", "b"), bb=1:2)
> c(typeof(myvar), mode(myvar), class(myvar))
[1] "list"
                 "list"
                              "data.frame"
> attributes(myvar)
$names
[1] "aa" "bb"
$class
[1] "data.frame"
$row.names
[1] 1 2
```

> dim(mvvar) [1] 2 2

Object Coercion

Coercion means changing the type, mode, or class of an object, often without changing the underlying data.

Changing the mode of an object can change its class as well, but not always.

Objects can be explicitly coerced using the "as.*" coercion functions.

Most coercion functions strip the attributes from the obiect.

Implicit coercion occurs when objects with different modes are combined into a vector, forcing the elements to have the same mode

Implicit coercion can cause bugs that are difficult to trace.

The rule is that coercion is into larger types (numeric objects are coerced into character strings).

Coercion can introduce bad data, such as NA values.

```
> mvvar <- 1:5
> c(typeof(myvar), mode(myvar), class(myvar))
[1] "integer" "numeric" "integer"
> mode(myvar) <- "character" # Coerce to "character"
> mvvar
[1] "1" "2" "3" "4" "5"
> c(typeof(myvar), mode(myvar), class(myvar))
[1] "character" "character" "character"
> # Explicitly coerce to "character"
> myvar <- as.character(1:5)
> c(typeof(myvar), mode(myvar), class(myvar))
[1] "character" "character" "character"
> matrixv <- matrix(1:10, 2, 5) # Create matrix
> # Explicitly coerce to "character"
> matrixv <- as.character(matrixv)
> c(typeof(matrixv), mode(matrixv), class(matrixv))
[1] "character" "character" "character"
> # Coercion converted matrix to vector
> c(is.matrix(matrixv), is.vector(matrixv))
[1] FALSE TRUE
> as.logical(0:3) # Explicit coercion to "logical"
[1] FALSE TRUE TRUE TRUE
> as.numeric(c(FALSE, TRUE, TRUE, TRUE))
[1] 0 1 1 1
> c(1:3, "a") # Implicit coercion to "character"
[1] "1" "2" "3" "a"
> # Explicit coercion to "numeric"
```

> as.numeric(c(1:3, "a")) [1] 1 2 3 NA

Basic R Objects

The quotation marks "" (or '') around a character string tell R that it's a string, not a variable name.

Vectors are the basic building blocks of R objects.

There are no scalars in R, and single values are stored as vectors of length "1".

A character string is also a vector with a single element, with the first element of the vector containing the string of text.

The colon binary operator ':' produces a vector.

The function c() combines objects into a vector.

The "[1]" symbol means the return value is a vector.

The function is .vector() returns TRUE if its argument is a vector, and returns FALSE otherwise.

- > "Hello World!" # Type some text
- > # hello is a variable name, because it's not in quotes
- > hello # R interretsp "hello" as a variable name
- > is.vector(1) # Single number is a vector
 > is.vector("a") # String is a vector
- > 4:8 # Create a vector
- > # Create vector using c() combine function
- > c(1, 2, 3, 4, 5)
- > # Create vector using c() combine function
- > c("a", "b", "c")
 - > # Create vector using c() combine function
 - > c(1, "b", "c")

Character Strings

Character strings are sequences of characters (and vectors of length one).

The function nchar() returns the length of a string.

Special characters in strings:

"\t" for TAB,

"\n" for new-line.

"\\" for a (single) backslash character

The function cat() concatenates strings and echos them to console, without returning any values.

The function cat() is useful in user-defined functions.

```
> stringv <- "Some string"
> stringv
[1] "Some string"
> stringv[1]
[1] "Some string"
> stringv[2]
[1] NA
> NROW(stringv) # length of vector
> nchar(stringv) # length of string
[1] 11
> # Concatenate and echo to console
> cat("Hello", "World!")
Hello World!
> cat("Enter\ttab")
Enter tab
> cat("Enter\nnewline")
Enter
newline
> cat("Enter\\backslash")
Enter\backslash
```

Manipulating Strings

The function paste() concatenates its arguments into a string, coerces them to characters if needed, and returns the string.

If a vector or list is passed to paste(), together with a collapse string, then paste() concatenates the elements into a string, separated by the collapse string.

The function strsplit() splits the elements of a character vector.

Splitting on the "." character requires surrounding it with brackets: "[.]", or using argument fixed=TRUE.

The function substring() extracts or replaces substrings in a character string.

The recycling rule extends the length to match the longest object.

```
> stringv1 <- "Hello" # Define a character string
> stringv2 <- "World!" # Define a character string
> paste(stringv1, stringv2, sep=" ") # Concatenate and return value
[1] "Hello World!"
> cat(stringv1, stringv2) # Concatenate and echo to console
Hello World!
> paste("a", 1:4, sep="-") # Convert, recycle and concatenate
[1] "a-1" "a-2" "a-3" "a-4"
> paste(c("a1", "a2", "a3"), collapse="+") # Collapse vector to st
[1] "a1+a2+a3"
> paste(list("a1", "a2", "a3"), collapse="+")
[1] "a1+a2+a3"
> paste("Today is", Sys.time()) # Coerce and concatenate strings
[1] "Today is 2023-03-20 17:34:35"
> paste("Today is", format(Sys.time(), "%B-%d-%Y"))
[1] "Today is March-20-2023"
> strsplit("Hello World", split="r") # Split string
FF111
[1] "Hello Wo" "ld"
> strsplit("Hello.World", split="[.]") # Split string
[1] "Hello" "World"
> strsplit("Hello.World", split=".", fixed=TRUE) # Split string
[1] "Hello" "World"
> substring("Hello World", 3, 6) # Extract characters from 3 to 6
```

[1] "110 "

Regular Expressions in R

patterns into regular expressions.

R has Regex functions for pattern matching and replacement.

The function gsub() replaces all matches of a pattern in a string. The function grep() searches for matches of a pattern

in a string. The function glob2rx() converts globbing wildcard

```
> gsub("is", "XX", "is this gratis?") # Replace "is" with "XX"
[1] "XX thXX gratXX?"
> grep("b", c("abc", "xyz", "cba d", "bbb")) # Get indexes
[1] 1 3 4
> grep("b", c("abc", "xyz", "cba d", "bbb"), value=TRUE) # Get val
[1] "abc" "cba d" "bbb"
> glob2rx("abc.*") # Convert globs into regex
[1] "^abc\\."
> glob2rx("*.doc")
[1] "^.*\\.doc$"
```

Vectors

Vectors are the basic building blocks of R objects.

There are no scalars in R, and single values are stored as vectors of length "1".

The function c() combines values into a vector.

The function is.vector() returns TRUE if its argument is a vector, and returns FALSE otherwise.

The object letters is a constant and a vector,

```
> is.vector(1) # Single number is a vector
[1] TRUE
> is.vector("a") # String is a vector
[1] TRUE
> vectory <- c(8, 6, 5, 7) # Create vector
> vectorv
[1] 8 6 5 7
> vectorv[2] # Extract second element
Γ17 6
> # Extract all elements, except the second element
> vectorv[-2]
[1] 8 5 7
> # Create Boolean vector
> c(FALSE, TRUE, TRUE)
[1] FALSE TRUE TRUE
> # Extract second and third elements
> vectorv[c(FALSE, TRUE, TRUE)]
Γ17 6 5
> letters[5:10] # Vector of letters
[1] "e" "f" "g" "h" "i" "i"
> c("a", letters[5:10]) # Combine two vectors of letters
[1] "a" "e" "f" "g" "h" "i" "i"
```

Creating Vectors

The colon operator (":") provides a simple way of creating a numeric vector.

The function vector() returns a vector of the specified *mode*.

The functions seq(), seq_len(), and seq_along() return a sequence (vector) of numbers.

The function rep() replicates an object multiple times.

The functions character() and numeric() return zero-length vectors of the specified *mode* (not to be confused with a NULL object).

Zero length vectors are not the same as NULL objects.

```
> 0:10 # Vector of integers from 0 to 10
[1] 0 1 2 3 4 5 6 7 8 9 10
> vector() # Create empty vector
logical(0)
> vector(mode="numeric", length=10) # Numeric vector of zeros
[1] 0 0 0 0 0 0 0 0 0 0
> seq(10) # Sequence from 1 to 10
 [1] 1 2 3 4 5 6 7 8 9 10
> seq(along=(-5:5)) # Instead of 1:NROW(obj)
 [1] 1 2 3 4 5 6 7 8 9 10 11
> seq_along(c("a", "b", "c")) # Instead of 1:NROW(obj)
[1] 1 2 3
> seq(from=0, to=1, len=11) # Decimals from 0 to 1.0
 [1] 0.0 0.1 0.2 0.3 0.4 0.5 0.6 0.7 0.8 0.9 1.0
> seq(from=0, to=1, by=0.1) # Decimals from 0 to 1.0
 [1] 0.0 0.1 0.2 0.3 0.4 0.5 0.6 0.7 0.8 0.9 1.0
> seq(-2,2, len=11) # 10 numbers from -2 to 2
 [1] -2.0 -1.6 -1.2 -0.8 -0.4 0.0 0.4 0.8 1.2 1.6 2.0
> rep(100, times=5) # Replicate a number
F17 100 100 100 100 100
> character(5) # Create empty character vector
[1] "" "" "" ""
> numeric(5) # Create empty numeric vector
[1] 0 0 0 0 0
> numeric(0) # Create zero-length vector
numeric(0)
```

Arithmetic and Logical Operations on Vectors

Vectors can be multiplied and squared element by element, as if they were single elements.

When vectors are manipulated as if they were single elements, then R automatically performs a loop over the vector elements, and applies the operation element-wise.

This is a very powerful feature of R called *vectorized* arithmetic.

Vectorized arithmetic avoids writing loops and simplifies notation.

Vectors can be combined together and appended.

- > 2*4:8 # Multiply a vector
- > 2*(4:8) # Multiply a vector > 4:8/2 # Divide a vector
- > (0:10)/10 # Divide vector decimals from 0 to 1.0
- > vectorv <- c(8, 6, 5, 7) # Create vector
- > vectorv
- > # Boolean vector TRUE if element is equal to second one
- > vectorv == vectorv[2]
- > # Boolean vector TRUE for elements greater than $\ensuremath{\operatorname{six}}$
- > vectorv > 6
- > 2*vectorv # Multiply all elements by 2
 > vectorv^2 # Square all elements
- > c(11, 5:10) # Combine two vectors
- > c(11, 5:10) # Combine two vectors
- > c(vectorv, 2.0) # Append number to vector

Naming and Manipulating Vectors

Vector elements can be assigned namesv using a list of symbol-value pairs.

The function names() returns the namesv attribute of an object.

The namesv attribute of a vector can be modified by assigning to the names() function ("names() <-" assignment).

The function unname() removes the names attribute.

The function structure() adds attributes (specified as symbol=value pairs) to an object, and returns it.

```
> vectory <- # Create named vector
+ c(pi_const=pi, euler=exp(1), gamma=-digamma(1))
> vectory
pi_const
            euler
           2.718
                    0.577
   3.142
> names(vectory) # Get names of elements
[1] "pi_const" "euler"
                         "gamma"
> vectorv["euler"] # Get element named "euler"
euler
 2.72
> names(vectorv) <- c("pie", "eulery", "gammy") # Rename elements
> vectory
   pie eulery gammy
 3.142 2.718 0.577
> unname(vectorv) # Remove names attribute
[1] 3.142 2.718 0.577
> letters[5:10] # Vector of letters
[1] "e" "f" "g" "h" "i" "j"
> c("a", letters[5:10]) # Combine two vectors of letters
[1] "a" "e" "f" "g" "h" "i" "j"
> # Create named vector
> structure(sample(1:5), names=paste0("el", 1:5))
el1 el2 el3 el4 el5
  2 1 3 4 5
```

Subsetting Vectors

Vector elements can be subset (indexed, referenced) using the "[]" operator.

Vectors can be subset using vectors of:

- positive integers,
- negative integers,
- characters (names),
- Boolean vectors,

Negative integers remove the vector elements.

Subsetting with zero returns a zero-length vector.

A named vector can be subset using element names.

```
> vectory # Named vector
   pie eulery gammy
 3.142 2.718 0.577
> # Extract second element
> vectory[2]
eulery
  2.72
> # Extract all elements, except the second element
> vectory[-2]
  pie gammy
3.142 0.577
> # Extract zero elements - returns zero-length vector
> vectory[0]
named numeric(0)
> # Extract second and third elements
> vectorv[c(FALSE, TRUE, TRUE)]
eulery gammy
2.718 0.577
> # Extract elements using their names
> vectorv["eulery"]
eulerv
  2.72
> # Extract elements using their names
> vectorv[c("pie", "gammv")]
  pie gammy
3,142 0,577
```

> # Subset whole vector > vectorv[] <- 0

Filtering Vectors

 $\it Filtering$ means extracting elements from a vector that satisfy a logical condition.

When logical comparison operators are applied to vectors, they produce Boolean vectors.

Boolean vectors can then be applied to subset the original vectors, to extract their elements.

The function which() returns the indices of the TRUE elements of a Boolean vector or array.

```
> vectory <- runif(5)
> vectory
[1] 0.749 0.225 0.181 0.365 0.979
> vectory > 0.5 # Boolean vector
[1] TRUE FALSE FALSE FALSE TRUE
> # Boolean vector of elements equal to the second one
> vectorv == vectorv[2]
[1] FALSE TRUE FALSE FALSE FALSE
> # Extract all elements equal to the second one
> vectorv[vectorv == vectorv[2]]
Γ17 0.225
> vectory < 1 # Boolean vector of elements less than one
[1] TRUE TRUE TRUE TRUE TRUE
> # Extract all elements greater than one
> vectorv[vectorv > 1]
numeric(0)
> vectorv[vectorv > 0.5] # Filter elements > 0.5
[1] 0.749 0.979
> which(vectory > 0.5) # Index of elements > 0.5
[1] 1 5
```

> is.vector(factorv)
[1] FALSE

> # Coerce vector to factor > as.factor(1:5) [1] 1 2 3 4 5 Levels: 1 2 3 4 5

> # Coerce factor to character vector
> as.vector(as.factor(1:5))
[1] "1" "2" "3" "4" "5"

Factors

Factors are similar to vectors, but their elements can only take values from a set of *levels*.

Factors are designed for categorical data which can only take certain values.

The function factor() converts a vector into a factor. Factors have two attributes: *class* (equal to "factor")

and levels (the allowed values).

Although factors aren't vectors, the data underlying a factor is an integer vector, called an *encoding vector*.

The function as.numeric() extracts the encoding vector (indices) of a factor.

The function as.vector() coerces a factor to a character vector.

```
> # Create factor vector
> factorv <- factor(c("b", "c", "d", "a". "c". "b"))
> factory
[1] bcdacb
Levels: a b c d
> factory[3]
[1] d
Levels: a b c d
> # Get factor attributes
> attributes(factory)
$levels
[1] "a" "b" "c" "d"
$class
[1] "factor"
> # Get allowed values
> levels(factory)
[1] "a" "b" "c" "d"
> # Get encoding vector
> as.numeric(factory)
[1] 2 3 4 1 3 2
```

Tables of Categorical Data

The function unique() calculates the unique elements of an object.

The function levels() extracts the levels attribute

of a factor (the allowed values).

A contingency table is a matrix that contains the

A contingency table is a matrix that contains the frequency distribution of variables (factors) contained in a set of data.

The function table() calculates the frequency distribution of categorical data.

sapply() applies a function to a vector or a list of objects and returns a vector or a list.

```
> factory
[1] bcdacb
Levels: a b c d
> # Get unique elements
> unique(factorv)
[1] bcda
Levels: a b c d
> # Get levels attribute of the factor
> levels(factory)
[1] "a" "b" "c" "d"
> # Calculate the factor elements from its levels
> levels(factory)[as.numeric(factory)]
[1] "b" "c" "d" "a" "c" "b"
> # Get contingency (frequency) table
> table(factory)
factory
abcd
```

1 2 2 1

Classifying Continuous Numeric Data Into Categories

Numeric data that represents a magnitude, intensity, or score can be classified into categorical data, given a vector of breakpoints.

The breakpoints create intervals that correspond to different categories.

The categories combine elements that have a similar numeric magnitude.

findInterval() returns the indices of the intervals specified by "vec" that contain the elements of "x".

If there's an exact match, then findInterval() returns the same index as function match().

If there's no exact match, then findInterval() finds the element of "vec" that is closest to, but not greater than, the element of "x".

If all the elements of "vec" are greater than the element of "x", then findInterval() returns zero.

args() displays the formal arguments of a function.

```
> # Display the formal arguments of findInterval
> args(findInterval)
function (x, vec, rightmost.closed = FALSE, all.inside = FALSE,
    left.open = FALSE)
NUT.T.
> # Get index of the element of "vec" that matches 5
> findInterval(x=5, vec=c(3, 5, 7))
Γ17 2
> match(5, c(3, 5, 7))
Γ17 2
> # No exact match
> findInterval(x=6, vec=c(3, 5, 7))
Γ17 2
> match(6, c(3, 5, 7))
Γ17 NA
> # Indices of "vec" that match elements of "x"
> findInterval(x=1:8, vec=c(3, 5, 7))
[1] 0 0 1 1 2 2 3 3
> # Return only indices of inside intervals
> findInterval(x=1:8, vec=c(3, 5, 7), all.inside=TRUE)
[1] 1 1 1 1 2 2 2 2
> # make rightmost interval inclusive
> findInterval(x=1:8, vec=c(3, 5, 7), rightmost.closed=TRUE)
[1] 0 0 1 1 2 2 2 3
```

pleasant

very_cold

66.2

45.1

52.8

60.3

pleasant

Classifying Numeric Data Into Categories Example

Temperature can be categorized into "cold". "warm". "hot", etc.

A named numeric vector of breakpoints can be used to convert a temperature into one of the categories.

Breakpoints correspond to categories of the data.

The first *breakpoint* should correspond to the lowest category, and should have a value less than any of the data.

```
> # Named numeric vector of breakpoints
> breakv <- c(freezing=0, very_cold=30, cold=50, pleasant=60, warm=
> breaky
 freezing very_cold
                         cold pleasant
                                             warm
                                                        hot.
                 30
                           50
                                                         90
                                     60
                                               80
> tempv <- runif(10, min=10, max=100)
> feels_like <- names(breakv[findInterval(x=tempv, vec=breakv)])
> names(tempv) <- feels_like
> tempv
               cold freezing very_cold pleasant freezing pleasa
```

28.9

47.4

74.9

24.7

57 / 129

Jerzy Pawlowski (NYU Tandon) FRE6871 Lecture#1 March 20, 2023

Converting Numeric Data Into Factors Using cut()

The function cut() converts a numeric vector into a vector of factors, representing the intervals to which the numeric values belong.

 $\operatorname{cut}()$ divides the range of values into intervals, based on a vector of breaks.

cut() then assigns factors to the numeric values, representing the intervals to which the numeric values belong.

The parameter breaks is a numeric vector of break points that divide the range of values into intervals.

The argument "labels" is a vector of labels for the intervals.

The argument "right" is a Boolean indicating if the intervals should be closed on the right (and open on the left), or vice versa.

cut() can produce the same classification as findInterval(), but findInterval() is faster than cut(), because it's a compiled function.

```
> datav <- sample(0:6) + 0.1
> datav
[1] 0.1 2.1 4.1 6.1 3.1 1.1 5.1
> cut(x=datav, breaks=c(2, 4, 6, 8))
[1] <NA> (2.4] (4.6] (6.8] (2.4] <NA> (4.6]
Levels: (2.4] (4.6] (6.8]
> rbind(datay, cut(x=datay, breaks=c(2, 4, 6, 8)))
      [,1] [,2] [,3] [,4] [,5] [,6] [,7]
datay 0.1 2.1 4.1 6.1 3.1 1.1 5.1
        NA 1.0 2.0 3.0 1.0 NA 2.0
> # cut() replicates findInterval()
> cut(x=1:8, breaks=c(3, 5, 7), labels=1:2, right=FALSE)
[1] <NA> <NA> 1
                  1 2 2
                                 <NA> <NA>
Levels: 1 2
> findInterval(x=1:8, vec=c(3, 5, 7))
[1] 0 0 1 1 2 2 3 3
> # findInterval() is a compiled function, so it's faster than cut(
> vectory <- rnorm(1000)
> summary(microbenchmark(
+ find interval=findInterval(x=vectorv, vec=c(3, 5, 7)).
   cut=cut(x=vectorv, breaks=c(3, 5, 7)),
+ times=10))[, c(1, 4, 5)] # end microbenchmark summary
          expr mean median
1 find_interval 4.46 4.24
           cut. 69.68 61.46
```

Plotting Histograms of Frequency Data

The function hist() calculates and plots a histogram, and returns its data *invisibly*.

The parameter breaks is the number of cells of the histogram.

If the argument freq is TRUE then the frequencies (counts) are plotted, and if it's FALSE then the probability density is plotted (with total area equal to 1).

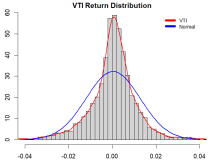
The function ${\tt density}()$ calculates a kernel estimate of the probability density for a sample of data.

The function lines() draws a line through specified points.

```
> returns <- na.omit(rutils::etfenv$returns$VTI)

# Plot histogram
> x11(width=6, height=5)
> par(mar=c(1, 1, 1, 1), oma=c(2, 2, 2, 0))
> madv <- mad(returns)
> histp <- hist(returns, breaks=100,
+ main="", xlim=c(-5*madv, 5*madv),
+ xlab="", vlab="", rfeq=FALSE)</pre>
```

> # Calculate VTI percentage returns



- > # Draw kernel density of histogram
- > lines(density(returns), col="red", lwd=2)
- > # Add density of normal distribution
- > curve(expr=dnorm(x, mean=mean(returns), sd=sd(returns)),
- + add=TRUE, type="1", 1wd=2, col="blue")
- > title(main="VTI Return Distribution", line=0)
- > # Add legend
- > legend("topright", inset=0.05, cex=0.8, title=NULL,
- + leg=c("VTI", "Normal"), bty="n",
- + lwd=6, bg="white", col=c("red", "blue"))
- > # Total area under histogram
- > # lotal area under histogram
- > sum(diff(histp\$breaks) * histp\$density)

Matrices

The function ${\tt matrix}$ () creates a matrix from a vector, and the matrix dimensions.

By default matrix() creates matrices column-wise, unless the argument byrow=TRUE is used.

The elements of matrices can be subset (referenced) using the "[]" operator.

The functions nrow() and ncol() return the number of rows and columns of a matrix.

The functions NROW() and NCOL() also return the number of rows or columns of a matrix, but they can also be applied to vectors, and treat vectors as single column matrices

```
> matrixv <- matrix(5:10, nrow=2, ncol=3) # Create a matrix
> matrixv # By default matrices are constructed column-wise
     [,1] [,2] [,3]
[1,]
        5
[2.]
> # Create a matrix row-wise
> matrix(5:10, nrow=2, byrow=TRUE)
     [,1] [,2] [,3]
[1.]
        5
[2,]
                 10
> matrixv[2, 3] # Extract third element from second row
Γ17 10
> matrixv[2, ] # Extract second row
[1] 6 8 10
> matrixv[, 3] # Extract third column
[1] 9 10
> matrixv[, c(1.3)] # Extract first and third column
     [,1] [,2]
        5
Γ2.1
            10
> matrixv[, -2] # Remove second column
     [,1] [,2]
        5
Γ2.1
           10
> # Subset whole matrix
> matrixv[] <- 0
> # Get the number of rows or columns
> nrow(vectory): ncol(vectory)
NULL
NIII.I.
> NROW(vectory): NCOL(vectory)
Γ17 1000
[1] 1
> nrow(matrixv): ncol(matrixv)
Γ17 2
Γ17 3
> NROW(matrixv); NCOL(matrixv)
```

Matrix Attributes

Arrays are vectors with a dimension attribute.

Matrices are two-dimensional arrays.

The dimension attribute of a matrix is an integer vector of length 2 (nrow, ncol).

The dimnames attribute is a list, with vector elements containing row and column names.

A named matrix can be subset using row and column names

```
> attributes(matrixy) # Get matrix attributes
$dim
[1] 2 3
> dim(matrixv) # Get dimension attribute
Γ17 2 3
> class(matrixy) # Get class attribute
[1] "matrix" "array"
> rownames(matrixv) <- c("row1", "row2") # Rownames attribute
> colnames(matrixy) <- c("col1", "col2", "col3") # Colnames attrib
> matrixv
     col1 col2 col3
row1 0
row2
             0
> matrixv["row2", "col3"] # Third element from second row
> names(matrixy) # Get the names attribute
> dimnames(matrixy) # Get dimnames attribute
[1] "row1" "row2"
[[2]]
[1] "col1" "col2" "col3"
> attributes(matrixy) # Get matrix attributes
$dim
[1] 2 3
$dimnames
$dimnames[[1]]
[1] "row1" "row2"
$dimnames[[2]]
[1] "col1" "col2" "col3"
```

Matrix Subsetting

Matrices can be subset in a similar way as Vectors, either by indices (integers), by characters (names), or Boolean vectors.

Subsetting a matrix to a single row or column produces a vector, unless the parameter "drop=FALSE" is used.

Subsetting with the parameter "drop=FALSE" prevents the implicit coercion and preserves the matrix *class*.

This is an example of implicit coercion in R, which can cause difficult to trace bugs.

```
> matrixv # matrix with column names
     col1 col2 col3
row1
       0
row2
> matrixv[1, ] # Subset rows by index
col1 col2 col3
   0 0
> matrixv[, "col1"] # Subset columns by name
row1 row2
   0
> matrixv[, c(TRUE, FALSE, TRUE)] # Subset columns Boolean vector
     col1 col3
row1
row2
> matrixv[1, ] # Subsetting can produce a vector!
col1 col2 col3
   0 0 0
> class(matrixv); class(matrixv[1, ])
[1] "matrix" "array"
[1] "numeric"
> is.matrix(matrixv[1, ]); is.vector(matrixv[1, ])
[1] FALSE
[1] TRUE
> matrixv[1, , drop=FALSE] # Drop=FALSE preserves matrix
     col1 col2 col3
> class(matrixv[1, , drop=FALSE])
[1] "matrix" "array"
> is.matrix(matrixv[1, , drop=FALSE]); is.vector(matrixv[1, , drop=
[1] TRUE
[1] FALSE
```

Logical Operators

R has the following logical operators:

- "<" less than,</p>
- "<=" less than or equal to,</p>
- ">" greater than,
- ">=" greater than or equal to,
- "==" exactly equal to,
- "!=" not equal to,
- "!x" Not x,
- "x & y" x AND y,
- "x | y" x OR y,

These operators are applied to vectors element-wise.

```
> TRUE | FALSE

> TRUE | NA

> vector1 <- c(2, 4, 6)

> vector1 < 5  # Element-wise comparison

> (vector1 < 5) & (vector1 > 3)

> vector1(vector1 < 5) & (vector1 > 3)]

> vector2 <- c(-10, 0, 10)

> vector1 < vector2

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

> c(FALSE, TRUE, FALSE) | c(TRUE, TRUE, FALSE)

Long Form Logical Operators

R also has two long form logical operators:

- "x && y" x AND y,
- "x || y" x OR y,

These operators differ from the short form operators in two ways:

- They only evaluate the first elements of their vector arguments,
- They short-circuit (stop evaluation as soon as the expression is determined),

Rule of Thumb

Use "&&" and "||" in if-clauses,

```
> c(FALSE, TRUE, FALSE) && c(TRUE, TRUE, FALSE)
> c(FALSE, TRUE, FALSE) || c(TRUE, TRUE, FALSE)
> echo_true <- function() {cat("echo_true\to"); TRUE}
> echo_true <- function() {cat("echo_false(\to"); FALSE})
> echo_true() | echo_false()
> # echo_true()
| echo_false()
| fecho_true()
| echo_false()
| fecho_false()
```

+ }

Arithmetic Operators

Arithmetic *operators* perform arithmetic operations on numeric or complex vectors,

- "+" performs addition,
- "-" performs subtraction,
- "*" performs multiplication,
- "/" performs division,
- "^" and "**" perform exponentiation,

> ?Arithmetic

> 4.7 * 0.5 # Multiplication > 4.7 / 0.5 # division

> 4.7 / U.5 # division

> # Exponentiation > 2**3

> 2^3

Comparing Objects With identical() and all.equal()

The function identical() tests if two objects are exactly the same, and always returns a single logical TRUE or FALSE (never NA or logical vectors).

For atomic arguments identical() often gives the same result as the "==" operator, but it's not synonymous with it in general.

The "==" operator applies the *recycling rule* to vector arguments and returns logical vectors, but identical() doesn't and returns a single logical value.

The function all.equal() tests the equality of two objects to within the square root of the *machine* precision.

The variable .Machine contains information about the numerical characteristics of the computer R is running on, such as the largest double and integer numbers, and the *machine precision*.

```
> numv <- 2
> numv==2
> identical(numv, 2)
> identical(numv. NULL)
> # This doesn't work:
> # numv==NUI.I.
> is.null(numv)
> vectorv <- c(2, 4, 6)
> vectorv==2
> identical(vectorv, 2)
> # numv is equal to "1.0" within machine precision
> numv <- 1.0 + 2*sqrt(.Machine$double.eps)
> all.equal(numv, 1.0)
> # Info machine precision of computer R is running on
> # ?.Machine
> # Machine precision
> .Machine$double.eps
```

match=match(5, vectorv),
which=min(which(vectorv == 5)).

Lookup and Matching Using which() and match()

The function which() returns the indices of the TRUE elements of a Boolean vector or array.

If the argument is an array and arr.ind=TRUE, then which() returns a matrix with rows containing the indices of the TRUE elements.

The functions which.max() and which.min() return the index of the minimum or maximum of a numeric or Boolean vector.

match() returns the index of the vector element that exactly matches its first argument.

If it doesn't find an exact match then it returns NA.

The expressions match(x, vec_tor) and

min(which(vec_tor == x)) produce the same result,
but match() can be faster for large vectors.

```
> vectorv <- sample(1e3, 1e3)
> matrixy <- matrix(vectory, ncol=4)
> which(vectory == 5)
> match(5, vectorv)
> # Equivalent but slower than above
> (1:NROW(vectory))[vectory == 5]
> which(vectory < 5)
> # Find indices of TRUE elements of Roolean matrix
> which((matrixv == 5)|(matrixv == 6), arr.ind=TRUE)
> # Equivalent but slower than above
> arrayInd(which((matrixy == 5)|(matrixy == 6)).
     dim(matrixy), dimnames(matrixy))
> # Find index of largest element
> which.max(vectorv)
> which(vectory == max(vectory))
> # Find index of smallest element
> which.min(vectory)
> # Benchmark match() versus which()
> all.equal(match(5, vectory), min(which(vectory == 5)))
> library(microbenchmark)
> summary(microbenchmark(
```

times=10))[, c(1, 4, 5)] # end microbenchmark summary

Lookup and Matching Using %in% and any()

The binary operator %in% returns a Boolean vector with TRUE values corresponding to elements that have matches.

%in% is a wrapper for match() defined as follows:
"%in%" <- function(x, table) match(x, table,
nomatch=0) > 0.

%in% never returns NA, so it's preferred in if() statements.

any() returns TRUE if at least one element of a Boolean vector is TRUE, and FALSE otherwise.

The function pmatch() performs partial matching of strings.

```
> # Does 5 belong in vectory?
> 5 %in% vectory
> match(5, vectorv, nomatch=0) > 0
> # Does (-5) belong in vectory?
> (-5) %in% vectory
> c(5, -5) %in% vectorv
> match(-5, vectorv)
> # Equivalent to "5 %in% vectorv"
> any(vectorv == 5)
> # Equivalent to "(-5) %in% vectorv"
> any(vectorv == (-5))
> # Any negative values in vectory?
> any(vectorv < 0)
> # Example of use in if() statement
> if (any(vectorv < 2))
+ cat("vector contains small values\n")
> # Partial matching of strings
```

> pmatch("med", c("mean", "median", "mode"))

Finding Closest Match Using findInterval()

The function match() returns the index of the vector element that *exactly* matches its first argument.

If $\mathtt{match}(\tt)$ doesn't find an exact match then it returns \mathtt{NA}

The function findInterval() returns the indices of the intervals specified by "vec" that contain the elements of "x".

If there's an exact match, then findInterval() returns the same index as function match().

If there's no exact match, then findInterval() finds the element of "vec" that is closest to, but not greater than, the element of "x".

- > str(findInterval)
- > # Get index of the element of "vec" that matches 5
- > findInterval(x=5, vec=c(3, 5, 7))
- > match(5, c(3, 5, 7))
 > # No exact match
- > findInterval(x=6, vec=c(3, 5, 7))
- > match(6, c(3, 5, 7))
- > # Indices of "vec" that match elements of "x"
- > findInterval(x=1:8, vec=c(3, 5, 7))
- > # Return only indices of inside intervals
 > findInterval(x=1:8, vec=c(3, 5, 7), all.inside=TRUE)
- > findInterval(x=1:8, vec=c(3, 5, 7), all.inside=TRUE)
 > # Make rightmost interval inclusive
- > findInterval(x=1:8, vec=c(3, 5, 7), rightmost.closed=TRUE)

Assignment Operators

The standard assignment operator in R is "<-".

Both "<-" and "=" are valid assignment operators in R.

The "<-" operator may cause an error if R confuses it with the "<" logical operator.

But they differ in *scope* and *precedence* ("<-" has higher precedence than "=").

The "=" operator is used for named arguments in function calls

When variables are assigned within an argument list using the "=" operator, their *scope* is limited to the function.

Rule of Thumb:

Use "<-" in R scripts and inside functions, Use "=" only in function calls.

```
> numv1 <- 3  # "<-" and "=" are valid assignment operators
> numv1 = 3
> numv1 = 3
> numv1 = 2 <- 3  # "<" operator confused with "<-"
> 2 <- 3  # "<" operator confused with "<-"
> 2 <- 3  # Add space or brackets to avoid confusion
> # "=" assignment within argument list
> median(x=1:10)
> x  # x doesn't exist outside the function
> # "<-" assignment within argument list
> median(x <- 1:10)
> x  # x exists outside the function
```

The assign() Function

The assign() function assigns a value to an object in a specified environment, by referencing it using a character string (name).

assign() can be used to either assign values to existing variables, or to create new variables.

assign() looks for the object name in the specified environment, and assigns a value to it.

If assign() can't find the object name, then it creates it.

assign() expects a character string as its argument.

If a object name is passed to assign(), then it evaluates that object to get the string it contains.

If the object doesn't contain a string, then assign() produces an error.

```
> mvvar <- 1 # Create new object
> assign(x="myvar", value=2) # Assign value to existing object
> myvar
> rm(myvar) # Remove myvar
> assign(x="myvar", value=3) # Create new object from name
> myvar
> # Create new object in new environment
> new_env <- new.env() # Create new environment
> assign("myvar", 3, envir=new_env) # Assign value to name
> ls(new_env) # List objects in "new_env"
> new_env$myvar
> rm(list=ls()) # delete all objects
> symbol <- "myvar" # define symbol containing string "myvar"
> assign(symbol, 1) # Assign value to "myvar"
> 1s()
> myvar
> assign("symbol", "new_var")
> assign(symbol, 1) # Assign value to "new_var"
> 1s()
> symbol <- 10
```

> assign(symbol, 1) # Can't assign to non-string

> identical(DAX, EuStockMarkets[, "DAX"])

Applying assign() to Lists of Names

assign() allows creating new objects from listv or vectors of names (character strings), such as column names.

```
> rm(list=ls()) # delete all objects
> # Create individual vectors from column names of EuStockMarkets
> for (colname in colnames(EuStockMarkets)) {
    # Assign column values to column names
    assign(colname, EuStockMarkets[, colname])
    + } # end for
> ls()
> head(DAX)
> head(EuStockMarkets[, "DAX"])
```

Jerzy Pawlowski (NYU Tandon)

Retrieving Objects Using get()

The function get() accepts a character string and returns the value of the corresponding object in a specified *environment*.

get() retrieves objects that are referenced using character strings, instead of their names.

The functions get() and assign() allow retrieving and assigning values to objects that are referenced using character strings.

The function mget() accepts a vector of strings and returns a list of the corresponding objects.

- > # Create new environment
- > test_env <- new.env()
 > # Pass string as name to create new object
- > assign("myvar1", 2, envir=test_env)
 > # Create new object using \$ string referencing
- > test_env\$myvar2 <- 1
- > # List objects in new environment
- > 1s(test env)
- > # Reference an object by name
- > test_env\$myvar1
- > # Reference an object by string name using get
- > get("myvar1", envir=test_env)
 > # Retrieve and assign value to object
- > assign("myvar1",
- + 2*get("myvar1", envir=test_env),
 + envir=test_env)
- > get("myvar1", envir=test_env)
- > # Return all objects in an environment
- > mget(ls(test_env), envir=test_env)
- > # delete environment
- > rm(test_env)

The Parenthesis "()" and Curly Braces "{}" Operators

The parenthesis "()" and curly braces "{}" operators are used to enclose and to group (combine) expressions.

The parenthesis "()" and curly braces " $\{\}$ " operators are functions, and they return values.

An expression enclosed by the parenthesis "()" operator is evaluated separatately from other expressions, and its result is returned.

Enclosing expressions in parenthesis makes them less ambiguous.

The curly braces "{}" operator can group several expressions, that can be written either on separate lines, or be separated by the semicolon ";" operator.

The curly braces "{}" operator returns the last expression it encloses.

Both the parenthesis "()" and curly braces "{}" operators are functions, and executing them requires a little additional processing time.

The square braces (brackets) "[]" operator subsets (references) the elements of vectors, matrices, and listv.

```
> # expressions enclosed in parenthesis are less ambiguous
> -2:5
> (-2):5
> -(2:5)
> # expressions enclosed in parenthesis are less ambiguous
> -2*3+5
> -2*(3+5)
> # expressions can be separated by semicolons or by lines
> {1+2; 2*3; 1:5}
> # or
> {1+2
+ 2*3
+ 1:5}
> matrixv <- matrix(nr=3, nc=4)
> matrixv <- 0
> # subset whole matrix
> matrixv[] <- 0
> # parenthesis and braces require a little additional processing t
> library(microbenchmark)
> summary(microbenchmark(
    basep=sqrt(rnorm(10000)^2),
   parven=sqrt(((((rnorm(10000)^2))))),
    bra ce=sqrt({{{ffrnorm(10000)^2}}}}),
    times=10))[, c(1, 4, 5)] # end microbenchmark summary
```

The "if () else" Control Statement

R has the familiar "if () $\{...\}$ else $\{...\}$ " statement to control execution flow depending on logical conditions.

The logical conditions must be either a Boolean or numeric type, otherwise an error is produced.

The "else" statement can also be omitted.

"if" statements can be nested using multiple "else if" statements.

```
> numv1 <- 1
>
    if (numv1) { # numeric zero is FALSE, all other numbers are TRUE
+ numv2 <- 4
+ } else if (numv1 == 0) { # 'else if' together on same line
+ numv2 <- 0
+ } else { # 'else' together with curly braces
+ numv2 <- -4
+ } # end if
>
    numv2 <- 0</pre>
```

The switch() Control Statement

The function switch() matches its first argument "EXPR" with one of the symbols in the following arguments, evaluates the corresponding expression, and returns it.

The arguments that follow the first argument "EXPR" should be given as *symbol=value* pairs.

If "EXPR" is a character string, then the expression bound to that symbol is returned by switch().

If "EXPR" is an integer, then switch() returns the expression from that position.

If switch() can't match "EXPR" to any symbol, then it returns NULL invisibly.

Using switch() is a convenient alternative to a cascade of "if () else" statements.

The function match.arg() matches a string to one of the possible values, and returns the matched value, or produces an error if it can't match it.

```
> switch("a", a="aaahh", b="bee", c="see", d=2,
         "else this")
> switch("c", a="aaahh", b="bee", c="see", d=2,
         "else this")
> switch(3, a="aaahh", b="bee", c="see", d=2,
         "else this")
> switch("cc", a="aaahh", b="bee", c="see", d=2,
         "else this")
> # measure of central tendency
> centra lity <- function(input.
      method=c("mean", "mean narm", "median")) {
+ # validate "method" argument
    method <- match.arg(method)
    switch(method.
  mean=mean(input).
+ mean narm=mean(input, na.rm=TRUE).
+ median=median(input))
+ } # end centra_lity
> mvvar <- rnorm(100, mean=2)
> centra litv(mvvar, "mean")
> centra lity(myvar, "mean narm")
> centra_lity(myvar, "median")
```

Iteration Using for() and while() Loops

```
The for() loop statement:
                                                             > color_list <- list("red", "white", "blue")
                                                             > # loop over list
> for (indeks in vectory) {ex pressions}
                                                             > for (some_color in color_list) {
iterates the dummy variable in_dex over the elements
                                                                 print(some_color)
                                                              + } # end for
of the vector or list vec_tor, and evaluates in a loop the
                                                             > # loop over vector
ex_pressions contained in the body of the for() loop.
                                                             > for (indeks in 1:3) {
                                                                 print(color list[[indeks]])
Upon loop exit the dummy variable in_dex is left equal
                                                             + } # end for
to the last element of the vector vec_tor.
                                                             > # while loops require initialization
while() loops start by testing their logical condition,
                                                             > indeks <- 1
and they repeat executing the loop body until that
                                                             > # while loop
                                                             > while (indeks < 4) {
condition is FALSE
                                                                 print(color_list[[indeks]])
                                                                 indeks <- indeks + 1
But while() loops risk producing infinite loops if not
                                                              + } # end while
written properly, so Use Them With Care!
```

Performing Loops Using for() and apply()

The for() loop doesn't return a value, so values calculated in the for() loop body must be assigned to variables in the parent environment, or otherwise they are lost.

The expressions in the for() loop body have access to variables in the parent environment in which the for() loop is executed, and they can modify those variables.

So even though for() loops don't return a value, they can be used to perform calculations on variables in the parent environment, but this is discouraged since it can produce errors that are hard to debug.

Rule of Thumb:

- for() loops are preferred for producing side effects, like plotting or reading and writing data to files,
- apply() loops are preferred for performing calculations which produce vectors or matrices of values,

> fib sea

Fibonacci Sequence Using for() Loop

The *Fibonacci* sequence of integers is defined by the recurrence relation:

$$F_n = F_{n-1} + F_{n-2},$$

 $F1 = 0, F2 = 1,$
 $F_n = 0, 1, 1, 2, 3, 5, 8, 13, ...$

The Fibonacci sequence was invented by the Indian mathematician Virahanka in the 8th century AD, and later described by the Italian mathematician Fibonacci in his famous treatise Liber Abaci.

Very often variables are initialized to NULL before the start of iteration.

A more efficient way to perform iteration is by pre-allocating the vector.

The function numeric() returns an zero length numeric vector.

The function numeric(k) returns a numeric vector of zeros of length k.

```
> # fib_seq <- numeric() # zero length numeric vector
> # pre-allocate vector instead of "growing" it
> fib_seq <- numeric(1)
> fib_seq[1] <- 0 # initialize
> for (i in 3:10) { # perform recurrence loop
+ fib_seq[i] <- fib_seq[i-1] + fib_seq[i-2]
+ } # end for</pre>
```

Allocating Memory to Vectors and Matrices

 ${\tt R}$ automatically allocates memory to new objects as needed during runtime, but at the cost of slowing down calculations.

Allocating memory of the correct mode speeds up calculations by avoiding automatic memory allocation by ${\bf R}.$

The functions character(), integer(), and numeric() return zero-length vectors of the specified mode.

Zero length vectors are not the same as NULL objects.

The function character(k) returns a character vector of empty strings of length k.

The function integer(k) returns a integer vector of zeros of length k.

The function numeric(k) returns a numeric vector of zeros of length k.

The function vector() by default returns a Boolean vector, unless the *mode* is specified.

The function matrix() by default returns a Boolean matrix containing NA values, unless the *mode* is specified.

```
> # Allocate character vector
> character()
> character(5)
> is.character(character(5))
> # Allocate integer vector
> integer()
> integer(5)
> is.integer(integer(5))
> is.numeric(integer(5))
> # Allocate numeric vector
> numeric()
> numeric(5)
> is.integer(numeric(5))
> is.numeric(numeric(5))
> # Allocate Boolean vector
> vector()
> vector(length=5)
> # Allocate numeric vector
> vector(length=5, mode="numeric")
> is.null(vector())
> # Allocate Boolean matrix
> matrix()
> is.null(matrix())
> # Allocate integer matrix
> matrix(NA_integer_, nrow=3, ncol=2)
> is.integer(matrix(NA integer , nrow=3, ncol=2))
> # Allocate numeric matrix
> matrix(NA real , nrow=3, ncol=2)
```

> is.numeric(matrix(NA real , nrow=3, ncol=2))

Logical Operators Applied to Vectors and Matrices

When logical operators are applied to vectors and matrices, they are applied element-wise, producing Boolean vectors and matrices.

```
> vectorv <- sample(1:9)
> vectorv < 5  # Element-wise comparison
> vectorv == 5  # Element-wise comparison
> matrixv <- matrix(vectorv, ncol=3)
> matrixv
> matrixv < 5  # Element-wise comparison
> matrixv == 5  # Element-wise comparison
```

Coercing Vectors Into Matrices

Vectors can be coerced into matrices by adding a dimension attribute.

The dimnames attribute can be assigned a named list to convert it into a named matrix.

The function structure() adds attributes (specified as symbol=value pairs) to an object, and returns it.

```
> matrixv <- 1:6  # Create a vector

> class(matrixv)  # Get its class

> # Is it vector or matrix?

> c(is.vector(matrixv), is.matrix(matrixv))

> structure(matrixv, dim=c(2, 3))  # Matrix object

> # Adding dimension attribute coerces into matrix

> dim(matrixv) <- c(2, 3)

> class(matrixv)  # Get its class

> # Is it vector or matrix?

> c(is.vector(matrixv), is.matrix(matrixv))

> # Assign dimnames attribute

> dimnames(matrixv) <- list(rows=c("row1", "row2"),

+ columns=c("col1", "col2", "col3"))
```

> matrixv

Coercing Matrices Into Other Types

Matrices can be explicitly coerced using the "as.*" coercion functions.

But coercion functions strip the *attributes* from an object.

- > matrixv <- matrix(1:10, 2, 5) # Create matrix
 > matrixv
 - matrixv
- > # as.numeric strips dim attribute from matrix
- > as.numeric(matrixv)
- > # Explicitly coerce to "character"
- > matrixv <- as.character(matrixv)
- > c(typeof(matrixv), mode(matrixv), class(matrixv))
- > # Coercion converted matrix to vector > c(is.matrix(matrixv), is.vector(matrixv))
- C(IS.Matlix(Matlixv), IS.Vector(Matlixv))

Binding Vectors and Matrices Together

Vectors can be bound into matrices using the functions cbind() and rbind().

The *recycling rule* allows operations on vectors of different lengths:

- Vectors are scanned from left to right,
- Shorter vectors are extended in length by recycling their values until they match the length of longer vectors,
- > vector1 <- 1:3 # Define vector > vector2 <- 6:4 # Define vector > # Bind vectors into columns > cbind(vector1, vector2) > # Bind vectors into rows > rbind(vector1, vector2) # Extend to four elements > vector2 <- c(vector2, 7) > # Recycling rule applied > cbind(vector1, vector2) > # Another example of recycling rule > 1:6 + c(10, 20)

Replicating Objects Using rep()

The function rep() replicates vectors and listv a given number of times.

rep() accepts a vector or list "x", and an integer specifying the type and number of replications.

Argument "times" replicates the whole vector a given number of times.

Argument "each" replicates each vector element a given number of times.

Argument "length.out" replicates the whole vector a certain number of times, so that the output vector length is equal to "length.out".

- > # Replicate a single element
- > rep("a", 5)
- > # Replicate the whole vector several times
- > rep(c("a", "b"), 5)
- > rep(c("a", "b"), times=5)
- > # Replicate the first element, then the second, etc.
- > rep(c("a", "b"), each=5) > # Replicate to specified length
- > rep(c("a", "b"), length.out=5)

Multiplying Vectors and Matrices

The multiplication "*" operator performs element-wise (element-by-element) multiplication of vectors and matrices.

By default the matrix elements are multiplied column-wise by the vector elements: the first matrix element in the first column is multiplied by the first vector element, then the second matrix column is multiplied by the remaining vector elements, etc.

The *recycling rule* is applied to the vector elements as needed

The transpose function t() can be applied if we want to perform row-wise multiplication.

But the transpose function t() is very slow for large matrices.

A better choice is to use functions lapply() and do.call().

```
> # Define vector and matrix
> vector1 <- c(2, 4, 3)
> matrixv <- matrix(sample(1:12), ncol=3)
> # Multiply columns of matrix by vector
> vector1*matrixv
> # Or
> matrixv*vector1
> # Multiply rows of matrix by vector
> t(vector1*t(matrixv))
> # Multiply rows of matrix by vector - transpose is very slow
> product <- lapply(1:NCOL(matrixy),
    function(x) vector1[x]*matrixv[, x])
> do.call(cbind, product)
> library(microbenchmark)
> summary(microbenchmark(
    trans=t(vector1*t(matrixv)),
    lapp={
      product <- lapply(1:NCOL(matrixv), function(x) vector1[x]*mat
      do.call(cbind, product)
    times=10))[, c(1, 4, 5)] # end microbenchmark summary
```

Matrix Inner Multiplication

The %*% operator performs inner (scalar) multiplication of vectors and matrices.

Inner multiplication multiplies the rows of one matrix with the columns of another matrix, so that each pair produces a single number:

$$C_{i,j} = \sum_{k=1}^{n} A_{i,k} B_{k,j}$$

Inner multiplication produces a vector or matrix with a reduced dimension.

Inner multiplication requires the dimensions of the matrices to be *conformable* (number of columns in the first matrix must be equal to the number of rows in the second).

The function drop() removes any dimensions of length one.

The functions rowSums() and colSums() calculate the sums of rows and columns, and they're very fast because they pass their data to compiled C++ code.

- > vector2 <- 6:4 # Define vector
- > # Multiply two vectors element-by-element
- > vector1 * vector2
- > # Calculate inner product
- > vector1 %*% vector2
 > # Calculate inner product and drop dimensions
- > drop(vector1 %*% vector2)
- > # Multiply columns of matrix by vector > matrixv %*% vector1 # Single column matrix
- > drop(matrixv %*% vector1) # vector
- > rowSums(t(vector1 * t(matrixv)))
- > # using rowSums() and t() is 10 times slower than %*%
- > library(microbenchmark)
 > summary(microbenchmark(
- + inner=drop(matrixv %*% vector1),
- + transp=rowSums(t(vector1 * t(matrixv))),
- + times=10))[, c(1, 4, 5)] # end microbenchmark summary

Matrix Transpose

The function t() returns the transpose of a matrix.

The function crossprod() also performs inner (scalar) multiplication, exactly the same as the %*% operator, but is slightly faster.

- > # Multiply matrix by vector fails because dimensions aren't confo > vector1 %*% matrixv
- > # Works after transpose
- > drop(vector1 %*% t(matrixv))
- > # Calculate inner product
- > crossprod(vector1, vector2)
 > # Create matrix and vector
- > matrixv <- matrix(1:3000, ncol=3)
- > tmatrixv <- t(matrixv)
- > vectorv <- 1:3
- > # crossprod() is slightly faster than "%*%" operator
- > summary(microbenchmark(
- + cross_prod=crossprod(tmatrixv, vectorv),
- + inner_prod=matrixv %*% vectorv,
- + times=10))[, c(1, 4, 5)] # end microbenchmark summary

Matrix Outer Multiplication

An *outer* product consists of all possible products of pairs of elements of two objects:

$$C_{i,j} = A_i \cdot B_j$$

An *outer* product of a function consists of applying it to all possible pairs of elements of two objects:

$$C_{i,j} = f(A_i, B_j)$$

Outer multiplication produces an object with dimension equal to the sum of the factors' dimensions, and with the number of elements equal to the product of the factors' elements.

The function outer() calculates the *outer* product of two matrices, and by default multiplies the elements of its arguments.

outer() can also calculate the values of a vectorized function of two variables passed to the "FUN" argument.

```
> # Define named vectors
> vector1 <- sample(1:4)
> names(vector1) <- paste0("row", 1:4, "=", vector1)
> vector1
> vector2 <- sample(1:3)
> names(vector2) <- paste0("col", 1:3, "=", vector2)
> vector2
> # Calculate outer product of two vectors
> matrixy <- outer(vector1, vector2)
> matrixy
> # Calculate vectorized function spanned over two vectors
> matrixy <- outer(vector1, vector2,
+ FUN=function(x1, x2) x2*sin(x1))
> matrixy
```

Functions in R.

R functions have three components:

- a list of formal arguments,
- a body containing R code,
- an environment.

An R function plus its environment is referred to as a function closures

The function body should be enclosed in curly braces {}, unless it contains a single command, then it doesn't have to enclosed

The function body doesn't require a return statement, since by default R functions return the last statement evaluated in the body.

args() displays the formal arguments of a function.

```
> # Define a function with two arguments
> testfun <- function(first_arg, second_arg) { # Body
  first_arg + second_arg # Returns last evaluated statement
+ } # end testfun
> testfun(1, 2) # Apply the function
> args(testfun) # Display argument
> # Define function that uses variable from enclosure environment
> testfun <- function(first_arg, second_arg) {
+ first_arg + second_arg + globv
+ } # end testfun
> testfun(3, 2) # error - globv doesn't exist yet!
> globv <- 10 # Create globv
> testfun(3, 2) # Now works
```

Return Values of Functions

value.

The function body doesn't require a return statement, since by default R functions return the last statement evaluated in the body.

return() statements are inserted in logical branches to

terminate function execution and return its intended

> ls() # List objects

Functions That Return invisible

If a return value is wrapped in the function invisible() then the return value isn't printed.

But if the function is assigned to a variable, then its return value is assigned to that variable. invisible() allows creating functions whose return values can be assigned, but which do not print when

values can be assigned, but which do not print when they're not assigned.

The function load() reads data from .RData files, and invisibly returns a vector of names of objects created in the workspace.

Binding Function Arguments

The formal arguments of a function are defined in its argument list.

When a function is called, it's passed a list of actual function arguments.

Formal arguments can be bound to actual arguments

either by name or by position:

- by name: formal arguments are bound to actual arguments with the same name,
- by position: the first formal argument is *bound* to the first actual argument, etc.

Binding by name takes precedence over binding by position: first all the named arguments are bound, then the remaining arguments are bound by position.

Partial argument names are bound to full names.

```
> testfum <- function(first_arg, second_arg) {
    # # Last statement of function is return value
    + first_arg + 2*second_arg
    } # end testfum
    > testfun(first_arg*3, second_arg*2) # Bind by name
    testfun(first_arg*3, second=2) # Partial name binding
    > testfun(3, 2) # Bind by position
    > testfun(3, 2) # Bind by position
    > testfun(second_arg*2, 3) # mixed binding
```

> testfun(3, 2, 1) # Too many arguments
> testfun(2) # Not enough arguments

All the actual arguments must be *bound* to formal arguments, and if not then an "unused argument" error is produced.

If there aren't enough formal arguments, then an "argument is missing" error is produced,

Default Values for Arguments

Formal arguments may be assigned default values, so that when the actual arguments are missing then their default values are used instead.

Default values are often assigned to function parameters, that determine the function's behavior.

Default values can be specified as a vector of strings, representing the possible values of a function's parameter.

The function match.arg() matches a string to one of the possible values, and returns the matched value, or produces an error if it can't match it.

The function str() displays the structure of an R object, for example a function name and its formal arguments.

- > # Function "paste" has two arguments with default values > str(paste)
- > # Default values of arguments can be specified in argument list
 > testfun <- function(first_arg, ratio=1) {</pre>
- + ratio*first_arg
- + } # end testfun
- > testfun(3) # Default value used for second argument
- > testfun(3, 2) # Default value over-ridden
- > # Default values can be a vector of strings
 > testfun <- function(input=c("first_val", "second_val")) {</pre>
- + input <- match.arg(input) # Match to arg list
- + input + } # end testfun
- > testfun("second_val")
- > testfun("se") # Partial name binding
- > testfun("some_val") # Invalid string

Function for Calculating Skew

R provides an easy way for users to write functions.

Formal function arguments can be bound to input variables by position or by name.

If the function arguments are missing then their default value is used.

Functions return the value of the last expression that is evaluated.

datasets is a base package containing various datasets, for example: EuStockMarkets.

The EuStockMarkets dataset contains daily closing prices of european stock indices.

- > # DAX percentage returns
- > returns <- rutils::diffit(log(EuStockMarkets[, 1]))
- > # calc_skew() calculates skew of time series of returns
 > # Default is normal time series
- > calc_skew <- function(returns=rnorm(1000)) {
 - # Number of observations
- nrows <- NROW(returns)
 # # Standardize returns
- + returns <- (returns mean(returns))/sd(returns)
- + # Calculate skew last statement automatically returned + nrows*sum(returns^3)/((nrows-1)*(nrows-2))
- + } # end calc_skew
- > # Calculate skew of DAX returns
- > # Bind arguments by name
- > calc_skew(returns=returns)
 > # Bind arguments by position
- > calc_skew(returns)
- > # Use default value of arguments
- > calc_skew()

The dots "..." Function Argument

The dots "..." function argument is a formal argument without a name, as opposed to the other formal arguments which all have names.

The dots "..." bind with any number of additional arguments, that aren't already bound by name or position to the named arguments.

The dots "..." are used when the number of arguments isn't known in advance, and allows functions to accept an indefinite number of arguments.

The dots "..." are sometimes placed *after* the named arguments, to allow passing of additional parameters into a function.

Functionals often place the dots "..." argument after the named arguments, to allow passing the dots "..." to the function being called by the functional.

```
> str(plot) # Dots for additional plot parameters
> bind_dots <- function(input, ...) {
+ paste(0'*input=", input,
+ ", dots=", paste(..., sep=", "))
+ } # end bind_dots
> bind_dots(1, 2, 3) # "input" bound by position
> bind_dots(2, input=1, 3) # "input" bound by name
> bind_dots(1, 2, 3, foo=10) # Named argument bound to dots
> bind_dots <- function(arg1, arg2, ...) {
+ arg1 * 2*arg2 + sum(...)
+ } # end bind_dots
> bind_dots(3, 2) # Bind arguments by position
> bind_dots(3, 2) # Bind arguments by position
> bind_dots(3, 2, 5, 8) # Extra arguments bound to dots
```

Argument Binding With dots "..." Argument

The dots "..." argument is sometimes placed *before* the named arguments, so that a function can accept an indefinite number of arguments, without binding them by position with the named arguments.

When the dots "..." are placed before the named arguments, the named arguments are often assigned default values, so they don't have to be bound to a value in the call.

Arguments that appear after the dots "..." must be bound by their full name, and can't be partially bound.

```
> str(sum) # Dots before other arguments

> sum(1, 2, 3) # Dots bind before other arguments

> sum(1, 2, NA, 3, na. m=TRUE)

> bind_dots <- function(..., input) {

+ paste(0'input=", input,

+ ", dots=", paste(..., sep=", "))

+ } # end bind_dots

> # Arguments after dots must be bound by full name

> bind_dots(1, 2, 3, input=10)

> bind_dots(1, 2, 3, input=10, foo=4) # Dots bound

> bind_dots(1, 2, 3) # "input" not bound

> bind_dots <- function(..., input=10) {

+ paste(0'input=", input,

+ ", dots=", paste(..., sep=", "))

+ } # end bind_dots
```

> bind_dots(1, 2, 3) # "input" not bound, but has default

Wrapper Functions With dots "..." Argument

Wrapper functions provide a convenient user interface to functions, by assigning default argument values, validating data, and formatting the output.

Wrapper functions are designed to perform the actions of other functions, while reducing their complexity.

The dots "..." argument of the *wrapper* function allows passing additional arguments on to the wrapped function.

Wrapper functions should be used with caution, since wrapping a function creates extra code (overhead), which slows down R.

```
> # Wrapper for mean() with default na.rm=TRUE
> my_mean <- function(x, na.rm=TRUE, ...) {
   mean(x=x, na.rm=na.rm, ...)
+ } # end my_mean
> foo <- sample(c(1:10, NA, rep(0.1, t=5)))
> mean(c(foo, NA))
> mean(c(foo, NA), na.rm=TRUE)
> my_mean(c(foo, NA))
> mv mean(c(foo, NA), trim=0.4) # Pass extra argument
> # Wrapper for saving data into default directory
> save data <- function(....
               file=stop("error: no file name").
               my_dir="/Users/jerzy/Develop/data") {
+ # Create file path
   file <- file.path(my_dir, file)
    save(.... file=file)
+ } # end save data
> foo <- 1:10
> save data(foo, file="scratch.RData")
> save data(foo, file="scratch.RData", mv dir="/Users/jerzv/Develop
> # Wrapper for testing negative arguments
> stop_if_neg <- function(input) {
+ if (!is.numeric(input) || input<0)
     stop("argument not numeric or negative")
+ } # end stop_if_neg
> # Wrapper for sqrt()
> my_sqrt <- function(input) {
+ stop_if_neg(input)
+ sqrt(input)
+ } # end my_sqrt
> my_sqrt(2)
> my_sqrt(-2)
> my_sqrt(NA)
```

> sum_dots(1, 2, 3, 4)

Recursive Functions with dots "..." Argument

Recursive functions can also accept the dots "..." argument.

The dots "..." argument can be referenced inside a function by first converting it into a list using "list(...)".

The function missing() returns TRUE if an argument is missing, and FALSE otherwise.

```
> # Recursive function sums its argument list
> sum_dots <- function(input, ...) {
  if (missing(...)) { # Check if dots are empty
     return(input) # just one argument left
   } else {
      input + sum_dots(...) # Sum remaining arguments
   } # end if
+ } # end sum dots
> sum_dots(1, 2, 3, 4)
> # Recursive function sums its argument list
> sum_dots <- function(input, ...) {
   if (NROW(list(...)) == 0) { # Check if dots are empty
     return(input) # just one argument left
   } else {
      input + sum_dots(...) # Sum remaining arguments
    } # end if
+ } # end sum dots
```

Recursive Function for Calculating Fibonacci Sequence

 $\label{lem:Recursive} \textit{Recursive} \text{ functions call themselves in their own body}.$

The *Fibonacci* sequence of integers is defined by the recurrence relation:

$$F_n = F_{n-1} + F_{n-2},$$

$$F1 = 0, F2 = 1,$$

$$F_n = 0, 1, 1, 2, 3, 5, 8, 13, \dots$$

The Fibonacci sequence was invented by Indian mathematicians, and later described by the Italian mathematician Fibonacci in his famous treatise Liber Abaci.

Exploring Functions

If a function name is called alone without arguments, then R displays the function code (but it must be on the search path).

Non-visible objects can't be viewed by calling their name.

The function getAnywhere() displays information about R objects, including non-visible objects.

The function getAnywhere() also displays R objects that aren't on the search path.

- > # Show the function code
- > plot.default > # Display function
- > getAnywhere(plot.default)

4□ > 4回 > 4 = > 4 = > = 990

> environment(func env)

Function Environments

When a function is called, a new *evaluation* environment is created.

The evaluation environment contains the function arguments and locally defined variables.

R evaluates variables inside functions by searching first in the $\it evaluation$ environment, then the $\it enclosure$ environment, then the R search path.

The enclosure of the *evaluation* environment is the environment where the function was defined.

The enclosure of functions defined in the workspace is the *global* environment.

The enclosure of functions defined in packages is the package *namespace*.

Objects defined in the function enclosure can be referenced inside the function.

```
> globv <- 1 # Define a global variable
> ls(environment()) # Get all variables in environment
> func_env <- function() { # Explore function environments
+ locvar <- 1 # Define a local variable
+ cat('objects in evaluation environment:\t',
+ ls(environment()), 'n')
+ cat('objects in enclosing environment:\t',
+ ls(parent.env(environment())), '\n')
+ cat('this is the enclosing environment:')
+ parent.env(environment()) # Return enclosing environment
+ } # end func_env
> func_env()
>
```

> environment(print) # Package namespace is the enclosure

Side effects Using the Super-assignment Operator "<<-"

Function *side effects* are operations on objects outside a function's *evaluation* environment.

The functions plot() and load() are examples of functions that produce *side effects*.

load() reads data from an .RData file, and creates objects in the workspace that are contained in the .RData file.

The super-assignment operator "<<-" allows creating functions that produce *side effects*.

The super-assignment operator "<<-" modifies or creates variables in the *enclosing* environment in which a function was *defined* (*lexical* scoping).

If a function was *defined* in the *global* environment then that's the function's *enclosing* environment, and the "<<-" operator operates on variables in the *global* environment.

```
> rm(list=ls()) # Remove all objects
> ls() # List objects
> # Load objects from file (side effect)
> load(file="my_data.RData")
> ls() # List objects
> globv <- 1 # Define a global variable
> # Explore function scope and side effects
> side effect <- function() {
   cat("global globv =", globv, "\n")
+ # Define local "globv" variable
   globy <- 10
   cat("local globv =", globv, "\n")
   # Re-define the global "globv"
   globy <<- 2
+ cat("local globv =", globv, "\n")
+ } # end side effect
> side effect()
> # Global variable was modified as side effect
> globy
```

Functions as First Class Objects

Functions in R are first class objects, which means they can be treated like any other R object:

- Functions can be passed as arguments to other functions.
- Functions can be nested (defined inside other functions),
- Functions can return functions as their return value.

Higher order functions are R functions that either accept a function as their argument (input) or return a function as their value (output).

- > # Create functional that accepts a function as input argument
- > testfun <- function(func name) { + # Calculates statistic on random numbers
- set.seed(1)
- func_name(runif(1e4)) # Apply the function name
- # end testfun > testfun(mean)
- > testfun(sd)

Functionals

Functionals are functions that accept a function or a function name (string) as one of their input arguments.

Functionals are able to execute function calls using the function names

The function match.fun() returns a function name that is specified by a string.

Functionals that call match.fun() are able to accept a string as a function name, because match.fun() converts it to a function.

match.fun() produces an error condition if it fails to find a function with the specified name.

- > # Functional accepts function name and additional argument
- > testfun <- function(func_name, input) {
- + # Produce function name from argument
- + func_name <- match.fun(func_name)
 + # Execute function call</pre>
- + func_name(input)
- + } # end testfun
- > testfun(sqrt, 4)
- > # String also works because match.fun() converts it to a function
- > testfun("sqrt", 4)
- > str(sum) # Sum() accepts multiple arguments
- > # Functional can't accept indefinite number of arguments
 - > testfun(sum, 1, 2, 3)

Functionals with dots "..." Argument

The dots "..." argument in *functionals* can be used to pass additional arguments to the function being called by the *functional*.

If named values are passed to the dots "..."

argument, then the functional can bind them to the correct formal arguments of the function being called by the functional.

```
> # Functional accepts function name and dots '...' argument > testfun <- function(func_name, ...) {
    func_name <- match.fun(func_name)
    func_name(...) # Execute function call
    } # end testfun
    testfun(sum, 1, 2, 3)
    testfun(sum, 1, 2, NA, 4, 5)
    testfun(sum, 1, 2, NA, 4, 5, na.rm=TRUE)
```

- > # Function with three arguments and dots '...' arguments > testfun <- function(input, param1, param2, ...) {
- + c(input=input, param1=param1, param2=param2, dots=c(...))
- + } # end testfun
- > testfun(1, 2, 3, param2=4, param1=5)
- > testfun(testfun, 1, 2, 3, param2=4, param1=5)
- > testfun(testfun, 1, 2, 3, 4, 5)

Anonymous Functions

R allows defining functions without assigning a name to them. * # Simple anonymous function (them.

Anonymous functions are functions that are not assigned to a name.

Anonymous functions can be passed as arguments to functionals.

2, 3, 4)

Functionals with Anonymous Functions

Anonymous functions can be passed as arguments to functionals.

Anonymous functions can also be used as default values for function arguments.

do.call(func name, list arg[-1])

> # do call() performs same operation as do.call()

do.call(sum. list(1, 2, NA, 3, na.rm=TRUE)),

rutils::do call(sum, list(1, 2, NA, 3), na.rm=TRUE))

> arg list <- list("sum", 1, 2, 3)

+ } # end testfun

> testfun(arg list)

> all.equal(

Executing Function Calls Using the do.call() Functional

The functional do.call() executes a function call using a function name and a list of arguments.

do.call() allows calling a function on arguments that are elements of a list.
do.call() passes the list elements individually, instead of passing the whole list as one argument:

do.call(fum, list) = fum(list[[1]], list[[2]],
...)

do.call() can be called inside other functionals to allow them to execute function calls.

The function ${\tt str}()$ displays the structure of an R object, for example a function name and its formal arguments.

The function do_call() from package *rutils* performs the same operation as do.call(), but using recursion, which is much faster and uses less memory.

```
> str(sum) # Sum() accepts multiple arguments
> # Sum() can't accept list of arguments
> sum(list(1, 2, 3))
> str(do.call) # "what" argument is a function
> # Do.call passes list elements into "sum" individually
> do.call(sum, list(1, 2, 3))
> do.call(sum, list(1, 2, NA, 3))
> do.call(sum, list(1, 2, NA, 3, na.rm=TRUE))
> # Functional accepts list with function name and arguments
> testfum <- function(list_arg) (
+ # Produce function name from argument
+ func_name <- match.fun(list_arg[[[I]])
+ # Execute function call uing do.call()
```

Performing Loops Using the apply() Functionals

An important example of *functionals* are the apply() functionals.

The functional apply() returns the result of applying a function to the rows or columns of an array or matrix.

If ${\tt MARGIN=1}$ then the function will be applied over the matrix ${\tt rows}$,

If MARGIN=2 then the function will be applied over the matrix *columns*.

apply() performs a loop over the list of objects, and can replace "for" loops in R.

```
> str(apply) # Get list of arguments

> # Create a matrix

> matrixv - matrix(6:1, nrow=2, ncol=3)

> matrixv

> # Sum the rows and columns

> rowsumv <- apply(matrixv, 1, sum)

> colsumv <- apply(matrixv, 2, sum)

> matrixv <- chind(c(sum(rowsumv), rowsumv),

+ rbind(colsumv, matrixv))

> dimnames(matrixv) <- list(c("colsumv", "row1", "row2"),

+ c("rowsumv", "col1", "col2", "col3"))

> matrixv
```

The apply() Functional with dots "..." Argument

The dots "..." argument in apply() is designed to pass additional arguments to the function being called by apply().

The additional arguments to apply() must be bound by their full (complete) names.

```
> matrixv[2, 2] <- NA # Introduce NA value

> matrixv

> # Calculate median of columns

> apply(matrixv, 2, median)

> # Calculate median of columns with na.rm=TRUE

> apply(matrixv, 2, median, na.rm=TRUE)
```

```
> str(apply) # Get list of arguments
> matrixv <- matrix(sample(12), nrow=3, ncol=4) # Create a matrix
> matrixv
> apply(matrixv, 2, sort) # Sort matrix columns
> apply(matrixv, 2, sort, decreasing=TRUE) # Sort decreasing order
```

The apply() Functional with Anonymous Functions

The apply() functional combined with anonymous functions can be used to loop over function parameters.

The dots "..." argument in apply() is designed to pass additional arguments to the function being called by apply().

The additional arguments to apply() must be bound by their full (complete) names.

```
> # DAX percentage returns
> returns <- rutils::diffit(log(EuStockMarkets[, 1]))
> library(moments) # Load package moments
> str(moment) # Get list of arguments
> # Apply moment function
> moment(xereturns, order=3)
> # 4x1 matrix of moment orders
> orderv <- as.matrix(1:4)
* # Anonymous function allows looping over function parameters
> apply(X=orderv, MARGIN=1,
* FUNFunction(orderp) {
* moment(x=returns, order=orderp)
+ } * end anonymous function
+ ) # end apply
> * # Another way of passing parameters into moment() function
```

> apply(X=orderv, MARGIN=1, FUN=moment, x=returns)

apply() Calling Functions with Multiple Arguments

When apply() calls a function with multiple arguments, then care must be taken for proper argument binding.

The dots "..." argument in apply() allows passing additional arguments to the function being called by apply().

The additional arguments to apply() must be *bound* by their full (complete) names.

The values of the "X" argument in apply() are bound by position to the first unused argument in the function being called by apply().

```
> # Function with three arguments
> testfun <- function(arg1, arg2, arg3) {
+ c(arg1=arg1, arg2=arg2, arg3=arg3)
+ } # end testfun
> testfun(1, 2, 3)
> datav <- as.matrix(1:4)
> # Pass datav to arg1
> apply(X=datav, MAR=1, FUN=testfun, arg2=2, arg3=3)
```

> # Pass datav to arg2

The lapply() Functional

The functional lapply() is a specialized version of the functional apply().

lapply() applies a function to a list of objects and returns a list

The function unlist() collapses a list with atomic elements into a vector (which can cause type coercion).

Rule of Thumb

It's often better to use lapply(), since apply() and sapply() attempt to coerce their output into a vector or matrix, which may cause them to fail.

- > # Vector of means of numeric columns
- > sapply(iris[, -5], mean)
- > # List of means of numeric columns
 > lapply(iris[, -5], mean)
- > # Lapply using anonymous function
 > unlist(lapply(iris,
- + function(column) {
- + if (is.numeric(column)) mean(column)
 - } # end anonymous function
 -) # end lapply) # end unlist
- > unlist(sapply(iris, function(column) {
- + if (is.numeric(column)) mean(column)}))

The sapply() Functional

The sapply() functional is a specialized version of the apply() functional.

sapply() applies a function to a vector or a list of objects and returns a vector or a list. sapply() tries to return a vector, but if the elements

can't be combined into a vector, then it returns a list.

When sapply() is given a data frame, it interretsp it as a list, and applies the function to each element (column) of the data frame.

sapply() Returning Matrices

If the function called by ${\tt sapply()}$ returns a vector, then ${\tt sapply()}$ returns a matrix, if possible.

The vectors returned by the function are arranged to form columns of the matrix returned by sapply().

But if the function returns vectors of different lengths, then sapply() cannot return a matrix, and returns a list instead.

This behavior of sapply() can cause run-time errors.

The function vapply() is similar to sapply(), but it always attempts to simplify its output to a matrix, and if it can't then it produces an error.

vapply() requires the argument FUN.VALUE that specifes the output format of the function called by vapply().

- > # Vectors form columns of matrix returned by sapply
 > sapply(2:4, function(num) c(el1=num, el2=2*num))
- > # Vectors of different lengths returned as list
- > sapply(2:4, function(num) 1:num)
- > # vapply is similar to sapply
- > vapply(2:4, function(num) c(el1=num, el2=2*num),
- + FUN.VALUE=c(row1=0, row2=0))
- > # vapply produces an error if it can't simplify
 > vapply(2:4, function(num) 1:num,
- + FUN.VALUE=c(row1=0, row2=0))

Normal (Gaussian) Probability Distribution

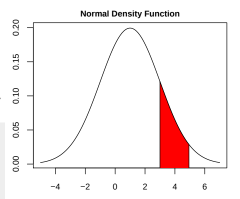
The Normal (Gaussian) probability density function is given by:

$$\phi(x,\mu,\sigma) = \frac{e^{-(x-\mu)^2/2\sigma^2}}{\sigma\sqrt{2\pi}}$$

The Standard Normal distribution $\phi(0,1)$ is a special case of the Normal $\phi(\mu,\sigma)$ with $\mu=0$ and $\sigma=1$.

The function ${\tt dnorm}()$ calculates the Normal probability density.

```
> xvar <- seq(-5, 7, length=100)
> yvar <- dnorm(xvar, mean=1.0, sd=2.0)
> plot(xvar, yvar, type="1", lty="solid",
+ xlab="", ylab="")
> title(main="Normal Density Function", line=0.5)
> startp <- 3; endd <- 5  # Set lower and upper bounds
> # Set polygon base
> subv <- ((xvar >= startp) & (xvar <= endd))
> polygon(c(startp, xvar[subv], endd), # Draw polygon
```



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+ c(-1, vvar[subv], -1), col="red")

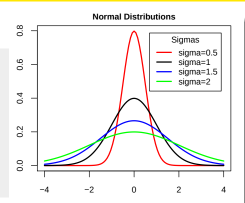
FRE6871 Lecture#1

March 20, 2023

Normal (Gaussian) Probability Distributions

Plots of several *Normal* distributions with different values of σ , using the function curve() for plotting functions given by their name.

```
> sigmavs <- c(0.5, 1, 1.5, 2) # Sigma values
> # Create plot colors
> colorv <- c("red", "black", "blue", "green")
> # Create legend labels
> labelv <- paste("sigma", sigmavs, sep="=")
> for (indeks in 1:4) { # Plot four curves
+ curve(expr=dnorm(x, sd=sigmavs[indeks]),
+ xlim=c(-4, 4).
+ xlab="", ylab="", lwd=2,
+ col=colorv[indeks].
+ add=as.logical(indeks-1))
+ } # end for
> # Add title
> title(main="Normal Distributions", line=0.5)
> # Add legend
> legend("topright", inset=0.05, title="Sigmas",
+ labely, cex=0.8, lwd=2, ltv=1, btv="n",
```

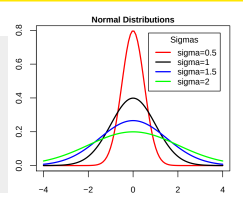


+ col=colory)

Normal Probability Distributions Plotted as Lines

Plots of several Normal distributions with different values of σ .

```
> xvar <- seq(-4, 4, length=100)
> sigmavs <- c(0.5, 1, 1.5, 2) # Sigma values
> # Create plot colors
> colorv <- c("red", "black", "blue", "green")
> # Create legend labels
> labelv <- paste("sigma", sigmavs, sep="=")
> # Plot the first chart
> plot(xvar, dnorm(xvar, sd=sigmavs[1]),
      type="n", xlab="", ylab="",
      main="Normal Distributions")
> # Add lines to plot
> for (indeks in 1:4) {
   lines(xvar, dnorm(xvar, sd=sigmavs[indeks]),
   lwd=2, col=colorv[indeks])
+ } # end for
> # Add legend
> legend("topright", inset=0.05, title="Sigmas",
+ labely, cex=0.8, lwd=2, lty=1, bty="n",
```



+ col=colorv)

The Log-normal Probability Distribution

If x follows the *Normal* distribution $\phi(x, \mu, \sigma)$, then the exponential of x: $y = e^x$ follows the Log-normal distribution $\log \phi()$:

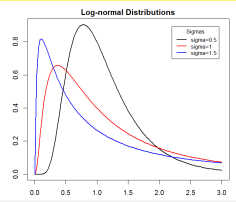
$$\log \phi(y,\mu,\sigma) = \frac{\exp(-(\log y - \mu)^2/2\sigma^2)}{y\sigma\sqrt{2\pi}}$$

With mean equal to: $\bar{y} = \mathbb{E}[y] = \exp(\mu + \sigma^2/2)$, and median equal to: $\tilde{v} = \exp(\mu)$

```
> sigmavs <- c(0.5, 1, 1.5)
> # Create plot colors
> colory <- c("black", "red", "blue")
   Plot all curves
> for (indeks in 1:NROW(sigmays)) {
   curve(expr=dlnorm(x, sdlog=sigmavs[indeks]),
   type="1", xlim=c(0, 3), lwd=2,
   xlab="", vlab="", col=colorv[indeks],
   add=as.logical(indeks-1))
```

end for

> # Standard deviations of log-normal distribution



- > # Add title and legend
- > title(main="Log-normal Distributions", line=0.5)
- > legend("topright", inset=0.05, title="Sigmas",
- + paste("sigma", sigmavs, sep="="),
- + cex=0.8, lwd=2, lty=rep(1, NROW(sigmavs)),
- + col=colory)

Chi-squared Distribution

Let z_1, \ldots, z_k be independent standard *Normal* random variables.

Then the random variable $X = \sum_{i=1}^k z_i^2$ is distributed according to the *Chi-squared* distribution with k degrees of freedom: $X \sim \chi_k^2$, and its probability density function is given by:

$$f(x) = \frac{x^{k/2-1} e^{-x/2}}{2^{k/2} \Gamma(k/2)}$$

The *Chi-squared* distribution with k degrees of freedom has mean equal to k and variance equal to 2k.

```
> # Degrees of freedom

> degf < c(2, 5, 8, 11)

> # Plot four curves in loop

> colorv <- c("red", "black", "blue", "green")

> for (indeks in 1:4) {

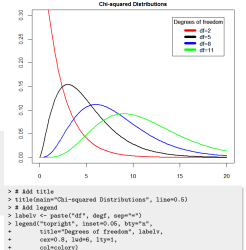
+ curve(expr=dchisq(x, df=degf[indeks]),

+ xlim=c(0, 20), ylim=c(0, 0.3),

+ xlab="", ylab="", col=colorv[indeks],

+ lvd=2. add=s.lorical(indeks=1);
```

+ } # end for



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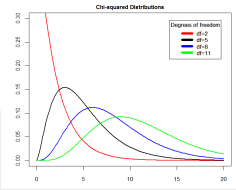
Chi-squared Distribution Plotted as Line

Let z_1, \ldots, z_k be independent standard *Normal* random variables.

Then the random variable $X = \sum_{i=1}^k z_i^2$ is distributed according to the *Chi-squared* distribution with k degrees of freedom: $X \sim \chi_k^2$, and its probability density function is given by:

$$f(x) = \frac{x^{k/2-1} e^{-x/2}}{2^{k/2} \Gamma(k/2)}$$

```
> degf <- c(2, 5, 8, 11) # df values
> # Create plot colors
> colorv <- c("red", "black", "blue", "green")
> # Create legend labels
> labelv <- paste("df", degf, sep="=")
> # Plot an empty chart
> xwar <- seq(0, 20, length=100)
> plot(xvar, dchisq(xvar, df=degf[1]),
+ type="n", xlab="", ylab="", ylim=c(0, 0.3))
> # Add lines to plot
> for (indeks in 1:4) {
+ lines(xvar, dchisq(xvar, df=degf[indeks]),
+ lwd=2, col=colorv[indeks])
+ } # end for
```



> # Add title > title(main="Chi-squared Distributions", line=0.5) > # Add legend > legend("topright", inset=0.05, + title="Degrees of freedom", labely,

cex=0.8, lwd=6, lty=1, bty="n", col=colorv)

Fisher's F-distribution

Let χ_n^2 and χ_n^2 be independent random variables following chi-squared distributions with m and n degrees of freedom.

Then the random variable:

$$F = \frac{\chi_m^2/m}{\chi_n^2/n}$$

Follows the F-distribution with m and n degrees of freedom, with the probability density function:

$$f(F) = \frac{\Gamma((m+n)/2)m^{m/2}n^{n/2}}{\Gamma(m/2)\Gamma(n/2)} \frac{F^{m/2-1}}{(n+mF)^{(m+n)/2}}$$

The F-distribution depends on the ratio F and also on the degrees of freedom, m and n.

The function df() calculates the probability density of

- the F-distribution > # Plot three curves in loop
- > degf <- c(3, 5, 9) # Degrees of freedom
- > colory <- c("black", "red", "blue", "green") > for (indeks in 1:NROW(degf)) {
- + curve(expr=df(x, df1=degf[indeks], df2=3),
- + xlim=c(0, 4), xlab="", vlab="", lwd=2,
- + col=colorv[indeks], add=as.logical(indeks-1))
- + } # end for

— df1=5, df2=3 df1=9, df2=3 0.0 2

F-Distributions

degrees of freedom df1=3 df2=3

- > # Add title
- > title(main="F-Distributions", line=0.5)
- > # Add legend
- > labelv <- paste("df", degf, sep="=")
- > legend("topright", inset=0.05, title="degrees of freedom",
 - labely, cex=0.8, lwd=2, lty=1,
 - col=colory)

Student's t-distribution

Let z_1, \ldots, z_{ν} be independent standard normal random variables, with sample mean: $\bar{z} = \frac{1}{i!} \sum_{i=1}^{\nu} z_i$ $(\mathbb{E}[\bar{z}] = \mu)$ and sample variance:

$$\hat{\sigma}^2 = \frac{1}{\nu - 1} \sum_{i=1}^{\nu} (z_i - \bar{z})^2$$

Then the random variable (t-ratio):

$$t = \frac{\bar{z} - \mu}{\hat{\sigma} / \sqrt{\nu}}$$

Follows the *t-distribution* with ν degrees of freedom. with the probability density function:

$$f(t) = \frac{\Gamma((\nu+1)/2)}{\sqrt{\pi\nu} \, \Gamma(\nu/2)} \, (1 + t^2/\nu)^{-(\nu+1)/2}$$

- > degf <- c(3, 6, 9) # df values
- > colorv <- c("black", "red", "blue", "green")
- > labelv <- c("normal", paste("df", degf, sep="="))
- > # Plot a Normal probability distribution
- > curve(expr=dnorm, xlim=c(-4, 4),
- xlab="", ylab="", lwd=2)
- > for (indeks in 1:3) { # Plot three t-distributions + curve(expr=dt(x, df=degf[indeks]),
- lwd=2, col=colorv[indeks+1], add=TRUE)
- + } # end for

Degrees of freedom normal df=3 df=6 df=9 0.1 0.0 -2 0 2

t-distributions

- > # Add title
- > title(main="t-distributions", line=0.5)
- > # Add legend
- > legend("topright", inset=0.05, bty="n",
- title="Degrees\n of freedom", labely,
 - cex=0.8, lwd=6, lty=1, col=colorv)

Student's t-distribution Plotted as Line

Let z_1, \ldots, z_{ν} be independent standard normal random variables, with sample mean: $\bar{z} = \frac{1}{\nu} \sum_{i=1}^{\nu} z_i$ ($\mathbb{E}[\bar{z}] = \mu$) and sample variance:

$$\hat{\sigma}^2 = \frac{1}{\nu - 1} \sum_{i=1}^{\nu} (z_i - \bar{z})^2$$

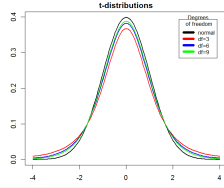
Then the random variable (t-ratio):

$$t = \frac{\bar{z} - \mu}{\hat{\sigma} / \sqrt{\nu}}$$

Follows the t-distribution with ν degrees of freedom, with the probability density function:

$$f(t) = rac{\Gamma((
u+1)/2)}{\sqrt{\pi
u}} (1+t^2/
u)^{-(
u+1)/2}$$

- > xvar <- seq(-4, 4, length=100)
- > degf <- c(3, 6, 9) # df values
- > colorv <- c("black", "red", "blue", "green")
 > labelv <- c("normal", paste("df", degf, sep="="))</pre>
- > # Plot chart of normal distribution
- > # Plot chart of normal distribution
 > plot(xvar, dnorm(xvar), type="1",
- lwd=2, xlab="", ylab="")
- > for (indeks in 1:3) { # Add lines for t-distributions + lines(xvar, dt(xvar, df=degf[indeks]),
- | lines(xvar, dt(xvar, dt=degi[indexs]),
- + lwd=2, col=colorv[indeks+1])
- + } # end for



- > # Add title
- > title(main="t-distributions", line=0.5)
- > # Add legend
- > legend("topright", inset=0.05, bty="n",
- + title="Degrees\n of freedom", labelv,
 - cex=0.8, lwd=6, lty=1, col=colorv)

Cauchy Distribution

The Cauchy distribution is Student's *t-distribution* with one degree of freedom $\nu=1$, with the probability density function:

$$f(x) = \frac{1}{\pi\sigma} \frac{1}{((x-\mu)/\sigma)^2 + 1}$$

Where μ is the location parameter (equal to the mean) and σ is the scale parameter.

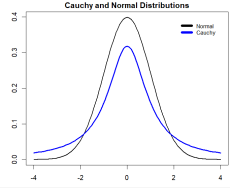
Since the *Cauchy* distribution has an infinite standard deviation, its measure of dispersion is the *interquartile* range (IQR), which is equal to σ .

The interquartile range is a robust measure of dispersion (scale), defined as the difference between the 75th minus the 25th percentiles.

The function dcauchy() calculates the *Cauchy* probability density.

The probability density of the Cauchy distribution decreases as the second power for large values of x:

$$f(x) \propto 1/x^2$$



- > # Plot the Normal and Cauchy probability distributions
- > curve(expr=dnorm, xlim=c(-4, 4), xlab="", ylab="", lwd=2)
- > curve(expr=dcauchy, lwd=3, col="blue", add=TRUE)
- > # Add title

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- > title(main="Cauchy and Normal Distributions", line=0.5)
- > # Add legend
- > legend("topright", inset=0.05, bty="n",
- + title=NULL,leg=c("Normal", "Cauchy"),
- + cex=0.8, lwd=6, lty=1, col=c("black", "blue"))

Pareto Distribution and Zipf's Law

The probability density of Student's t-distribution decreases as a power for large values of x:

$$f(x) \propto |x|^{-(\nu+1)}$$

The probability density of the Pareto distribution decreases as a power of the random variable x:

$$f(x) = \alpha x^{-(\alpha+1)}$$

For x > 1 and decay parameter $\alpha > 1$.

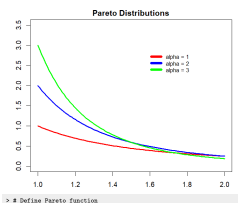
The mean μ and variance σ^2 of the *Pareto* distribution are equal to:

$$\mu = \frac{\alpha}{\alpha - 1}$$
 $\sigma^2 = \frac{\alpha}{(\alpha - 1)^2(\alpha - 2)}$

Zipf's law is analogous to the Pareto distribution, and applies to discrete variables.

Zipf's law states that the frequency f of a given value is inversely proportional to its rank n in the frequency table: $f(n) \propto n^{-s}$.

For example, Zipf's law applies to the frequency of words in a natural language.



```
> pare to <- function(x, alpha)
    alpha*x^(-alpha-1)
> colorv <- c("red", "blue", "green")
> alpha s <- c(1.0, 2.0, 3.0)
> for (indeks in 1:3) { # Plot three curves
    curve(expr=pare to(x, alpha s[indeks]),
    xlim=c(1, 2), vlim=c(0,0, 3,5),
    xlab="", vlab="", lwd=3, col=colorv[indeks],
    add=as.logical(indeks-1))
     # end for
> # Add title and legend
> title(main="Pareto Distributions", line=0.5)
```

Poisson Probability Distribution

The Poisson distribution gives the probability of the number of events observed in an interval of space or time.

The *Poisson* probability function is given by:

$$f(n;\lambda) = \frac{\lambda^n \cdot e^{-\lambda}}{n!}$$

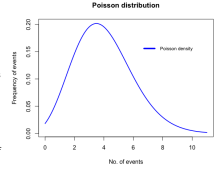
The Poisson random variable n is the number of events observed in the interval

The parameter λ is the average number of events that are observed in the interval.

An example of a *Poisson* distribution is the number of mail items received each day.

The function dpois() returns the probability density of the Poisson distribution.

The function rpois() returns random numbers following the Poisson distribution.



```
> # Poisson frequency
```

> eventy <- 0:11 # Poisson events

> poissonf <- dpois(eventv, lambda=4)

> names(poissonf) <- as.character(eventy)

> # Poisson function

> poissonfun <- function(x, lambda) {exp(-lambda)*lambda^x/factorial(x)}

> curve(expr=poissonfun(x, lambda=4), xlim=c(0, 11), main="Poisson distribution",

+ xlab="No. of events", vlab="Frequency of events", lwd=2, col="blue")

> legend(x="topright", legend="Poisson density", title="", bty="n",

+ inset=0.05, cex=0.8, bg="white", lwd=6, ltv=1, col="blue")

Homework Assignment

Required

- Study all the lecture slides in FRE6871_Lecture_1.pdf, and run all the code in FRE6871_Lecture_1.R,
- Study the RStudio Style Guide.

Recommended

• Read about the Vasicek single factor model in Vasicek Portfolio Default Distribution.pdf, BOE Credit Risk Models.pdf, BIS Bank Capital Model.pdf, and in Elizalde CDO Vasicek Credit Model.pdf.