

Introduction to R - Part 2

Data Science Skills Day 2022

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Course Description

- Introduction to R - Part 2
Data Science Skills Day

- The vast amount of data produced by evolving information technology requires tools and skills. Among the many tools, R is a free, open-source language for data sciences. R is a programming language that can aid in the process of data analysis. This course is a beginner level, introductory course for R for data analysis. We will learn about R, RStudio (the environment used to work in R), including installation, and apply R for beginner-level data modeling and visualization. By the end of the course, you'll have an introduction to the flexibility of R, different functionalities, and understand how to apply it for basic data exploration.

- Friday 10:00 am – 4 pm EST; online - synchronous.

Material

- Instructor Slides:
 - https://github.com/anjalisilva/DSI_IntroductionToR
 - SlideIntroR2022.pdf
 - SlideIntroR2022_part2.pdf

Covered So Far...

- Data Science Tools By Popularity
 - History of R
 - R and RStudio
 - Options To Work With R
 - Interacting with R
 - Documentation for R
 - R Coding Style
 - R Built-in Functions
 - R Packages: More in 2nd Part
 - R Data Types: Numeric, Logical, Character, Complex
 - Data Structures: Vectors, Matrices, Lists
 - Where To Get Further Help

Data Frames

Data Frames

- Data frames are used to store tabular data in R.

| data frame | 1 | "S" | TRUE |
|------------|-----------|---------|------|
| 7 | "A" | FALSE | |
| 3 | "U" | TRUE | |
| numeric | character | logical | |

Figure: Data frames can store different classes of objects in each column. Figure from: <https://datacarpentry.org/r-socialsci/02-starting-with-data/index.html>

Data Frames

- Data frames are used to store tabular data in R.
- Data frames can store different classes of objects in each column.

```
dataFrameExample <- data.frame(  
  numbers = 1:4,  
  sex = c("M", "M", "F", "F"))  
  
dataFrameExample  
class(dataFrameExample) # "data.frame"  
dim(dataFrameExample) # 4 2  
names(dataFrameExample) # "numbers" "sex"
```

Data Frames

- Data frames can be converted to a matrix using `data.matrix()`.

```
dataMatrix <- data.matrix(dataFrameExample)
class(dataMatrix) # "matrix"
dim(dataMatrix) # 4 2
```

Question 14:

Generate the following information into a data frame.

| | numbers | sex | age | height |
|---|---------|-----|-----|--------|
| 1 | | | | |
| 2 | 1 | M | 30 | 72 |
| 3 | 2 | M | 31 | 70 |
| 4 | 3 | F | 40 | 65 |
| 5 | 4 | F | 35 | 62.4 |

Question 15:

After generating the data frame, you realize the height is recorded in inches but should be changed to centimeters. How would you do this? Note, 1 inch = 2.54 cm.

| | numbers | sex | age | height |
|---|---------|-----|-----|--------|
| 1 | | | | |
| 2 | 1 | M | 30 | 72 |
| 3 | 2 | M | 31 | 70 |
| 4 | 3 | F | 40 | 65 |
| 5 | 4 | F | 35 | 62.4 |

Booleans

Boolean Operators

- Operators > or <:

```
a <- c(1:5)  
a  
a < 2 # TRUE FALSE FALSE FALSE FALSE
```

- Equivalence test:

```
a == 2 # FALSE TRUE FALSE FALSE FALSE
```

Boolean Operators

- & symbol is the "AND" operator:

```
a <- 45  
(a > 40) & (a < 50) # TRUE
```

- | symbol is the "OR" operator:

```
b <- 10  
(b < 20) | (b > 5) # TRUE
```

Any questions?

Question 16:

Given the below numeric vector 'numericVec', how can a user check if values are greater than 5?

```
numericVec <- c(1.1, 3, 5.3, 2)
```

Question 17:

Given the below numeric vector 'numericVec', how can a user retrieve the value/s that is/are greater than 5, from the vector?

```
numericVec <- c(1.1, 3, 5.3, 2)
```

Data Import/Export

Data Import/Export

- To see the list of pre-loaded data in R:

```
data(package = "datasets")  
  
AirPassengers # Example dataset  
head(AirPassengers) # see first few entries  
tail(AirPassengers) # see last few entries
```

Data Slicing

- Let us look at another pre-loaded datasets:

```
women # another dataset (last)
?women
dim(women) # 15  2
class(women) # "data.frame"
head(women) # height and weight information

women$height > 60 # slicing
women[women$height > 60, ] # slicing
```

Data Reading/Writing

- Files can be written using functions like `write.csv()`, `write.table()`:

```
getwd() # file will be saved here  
write.csv(x = women, file = "women.csv")  
# saving women dataset in current working directory
```

Data Reading/Writing

- Txt files can be read using `read.table("location of the file")` or `read.csv()`

```
womenNew <- read.csv(file = "women.csv", row.names = 1)  
womenNew
```

```
head(womenNew) # to view first part of object  
tail(womenNew) # to view last part of object  
dim(womenNew) # 15 2  
womenNew[c(1:5), ] # to view first 5 rows  
womenNew[, 1] # to view first column
```

Arithmetic

Arithmetic

- “+” is used for addition:

```
x <- 2.5 + 2  
x # 4.5
```

```
y <- 2:15  
sum(y) # 119  
sum(y[1:3]) # 9
```

Arithmetic

- “-” is used for subtraction:

```
x <- 2.5 - 2  
x # 0.5
```

- “/” is used for division:

```
x <- 2 / 2  
x # 1
```

Arithmetic

- “*” is used for multiplication:

```
x <- 2 * 2  
x # 4
```

- “%*% ” is used for matrix multiplication:

```
a <- matrix(1:6, nrow = 2, ncol = 3) # 2 x 3 matrix  
a  
b <- matrix(7:12, nrow = 3, ncol = 2) # 3 x 2 matrix  
b  
c <- a %*% b  
c # 2 x 2 matrix
```

Any questions?

Functions

Writing Functions

- Functions combine a sequence of expressions that are executed to achieve a goal.
- Can be reused without rewriting the sequence of expressions.
- Take input, *function arguments* and generate output, *return value*.
- When writing functions, ask yourself:
 - *What will the user want to modify in this function?*
 - This will help determine *function arguments*.

Function Interface

- Functions are defined using 'function' assigned to a variable name.

```
firstFunction <- function(argumentOne, argumentTwo) {  
  cat("\n First argument is", argumentOne, "\n")  
  cat("\n Second argument is", argumentTwo, "\n")  
  argumentThree <- argumentOne + argumentTwo  
  cat("\n argumentOne + argumentTwo is",  
      argumentThree, "\n")  
  
  return(argumentThree)  
  # always end with return statement (best practice)  
}
```

Function Interface

- Function names cannot begin with a number.

- To run the function:

```
firstFunction(argumentOne = 2, argumentTwo = 3)
```

- To “call” or “invoke” the function, type function name:

```
firstFunction
```

Function Interface

- Default values play a vital role in R functions and influence user's behaviour.
- Default values should be assigned using “`=`”, and not “`←`”.

```
secondFunction <- function(argOne = 1, argTwo = 3) {  
  cat("\n First argument is", argOne, "\n")  
  cat("\n Second argument is", argTwo, "\n")  
  argThree <- argOne + argTwo  
  
  cat("\n argOne + argTwo is", argThree, "\n")  
  
  return(argThree)  
}
```

Local and Global Variables

- Where variables are defined matters.
- Variables defined within functions are only accessible from within the function.
- Variables declared within a function are called “local”.
- Variables declared outside of functions are called “global”.

Variabls

- If variables are defined outside of functions, globally, they hold that value.

```
argOne <- "Hello"

anotherFunction <- function() {
  argOne <- 10
  return(argOne)
}

anotherFunction()

argOne # What would this return?
```

Variables

- Need to pass data into functions:

```
women
```

```
# Incorrect
dataAnalysis <- function() {
    heightData <- women$height
    return(mean(heightData))
}
dataAnalysis() # will work only for women$height
```

```
# Recommended
dataAnalysis <- function(inputData) {
    meanOfHeight <- mean(inputData)
    return(meanOfHeight)
}
dataAnalysis(inputData = women$height)
# will work for any dataset with height information
```

Any questions?

Non-base Functions

- Functions have been written by other authors.
- Non-base functions in R can be utilized by downloading R packages.
- To see all the currently loaded packages:

```
search()
```

Non-base Functions

- Popular repositories for Packages:
 - The Comprehensive R Archive Network (CRAN)
 - Link: <https://cran.r-project.org/web/packages/>
 - Bioconductor
 - Link: <https://www.bioconductor.org/packages/release/bioc/>
 - GitHub
- Depending on the source of Package, downloading instructions may differ.

Example

- User wants to do a cluster analysis of the women built-in dataset.
- Assume the user is interested in performing model-based clustering using Gaussian finite mixture models.
 - Do an R search:
`??clustering`
 - Google R packages for model-based clustering.
 - Read blogs on clustering.

Example, continue...

- Get to know about R package ‘mclust’.
- Visit <https://cran.r-project.org/web/packages/mclust/index.html>.
- Click on Reference manual to see all the functions within the package.
- Download package to R session:

```
# Install package from CRAN, case matters!
install.packages("mclust")
library("mclust") # to load and attach
```

Example, continue...

- More details on 'mclust' package (only after downloading):

```
vignette("mclust") # vignette for 'mclust'  
  
?`mclust-package` # get information on package  
  
?mclust # get information on package  
  
ls("package:mclust") # list all functions in package
```

Example, continue...

- Work with 'mclust' package:

```
# Running mclust
MclustResults <- mclust::Mclust(data = women)
str(MclustResults) # provide the structure; list of 15
names(MclustResults)
MclustResults$G # There are four clusters in the dataset

# Citing the package
citation("mclust")
```

Bioconductor

- Bioconductor packages are available from bioconductor.org.
- URL: <https://www.bioconductor.org/install/>.

```
if (! requireNamespace("BiocManager", quietly = TRUE))
install.packages("BiocManager")
```

- Bioconductor packages are listed here:
<https://www.bioconductor.org/packages/release/bioc/>.

Bioconductor



[Home](#) » [Bioconductor 3.11](#) » [3.11 Software Packages](#)

Bioconductor Software Packages

Bioconductor version: Release (3.11)

| Package | Maintainer | Title |
|---------------|--------------------------------|---|
| a4 | Tobias Verbeke, Laure Cougnaud | Automated Affymetrix Array Analysis Umbrella Package |
| a4Base | Tobias Verbeke, Laure Cougnaud | Automated Affymetrix Array Analysis Base Package |
| a4Classif | Tobias Verbeke, Laure Cougnaud | Automated Affymetrix Array Analysis Classification Package |
| a4Core | Tobias Verbeke, Laure Cougnaud | Automated Affymetrix Array Analysis Core Package |
| a4Preproc | Tobias Verbeke, Laure Cougnaud | Automated Affymetrix Array Analysis Preprocessing Package |
| a4Reporting | Tobias Verbeke, Laure Cougnaud | Automated Affymetrix Array Analysis Reporting Package |
| ABAEnrichment | Steffi Grote | Gene expression enrichment in human brain regions |
| ABarray | Yongming Andrew Sun | Microarray QA and statistical data analysis for Applied Biosystems Genome Survey Microrarray (AB1700) gene expression data. |
| abseqR | JiaHong Fong | Reporting and data analysis functionalities for Rep-Seq datasets of antibody libraries |

Packages *

Bioconductor's stable, semi-annual release:

- Analysis [software](#) packages.
 - [Annotation](#) packages.
 - Illustrative [experiment data](#) packages.
 - [Workflow](#) packages.
 - Latest [release announcement](#).

Bioconductor is also available via [Docker](#) and [Amazon Machine Images](#).

Development Version ➤

Bioconductor packages under development:

- Analysis [software](#) packages.
 - [Annotation](#) packages
 - Illustrative [experiment data](#) packages

Developer Resources:

- GIT Log

Example

- User is interested in Bioconductor package GenomicFeatures.
- To load package:

```
BiocManager::install("GenomicFeatures")
library("GenomicFeatures")

# list all functions in the package
ls("package:GenomicFeatures")
```

- More about installation:

<https://www.bioconductor.org/install/>.

Bioconductor

The screenshot shows a Twitter search interface. On the left, there is a sidebar with navigation icons: Home, Explore, Notifications, Messages, Bookmarks, Lists, and Profile. The main area displays a search result for the term "Bioconductor". The top card shows "Bioconductor" with 2,020 tweets. Below this is a user profile card for "Bioconductor" (@Bioconductor). The profile picture is a circular graphic of colored dots. The bio indicates the account is from Roswell Park Cancer Institute and joined in November 2011. It shows 5 Following and 7,028 Followers. There are also "Following" and notification buttons.

Bioconductor
2,020 Tweets

Bioconductor
@Bioconductor

② Roswell Park Cancer Institute ⚡ bioconductor.org
Joined November 2011

5 Following 7,028 Followers

GitHub

- To download from GitHub, need to use 'devtools' package.

```
library("devtools")
devtools::install_github("<GitHubUsername>/<PackageName>",
  build_vignettes = TRUE)
```

- Example: <https://github.com/anjalisilva/MPLNClust>

- To download this GitHub R package:

```
library("devtools")
devtools::install_github("anjalisilva/MPLNClust",
  build_vignettes = TRUE)
library("MPLNClust")
ls("package:MPLNClust") # to see all functions
```

Any questions?

Question 18:

You are interested in R packages for analyzing Covid-19 data for a project you are doing. So you search Google for such R packages.

You come across a package called 'covid19.analytics' which has a GitHub R package

(<https://github.com/mponce0/covid19.analytics>) that was later developed into a CRAN package (<https://cran.r-project.org/web/packages/covid19.analytics/index.html>). How would you download both the GitHub and CRAN package?

R: Graphics

Applying Base Functions

- Package 'graphics' is a standard or base package.

```
library(graphics) # to load and attach

?AirPassengers
# Monthly Airline Passenger Numbers 1949-1960

# plot
plot(AirPassengers)
plot(AirPassengers, type = "p")
plot(AirPassengers, type = "p", main = "Monthly Airline
    Passenger Numbers 1949-1960") # zoom
```

Applying Base Functions

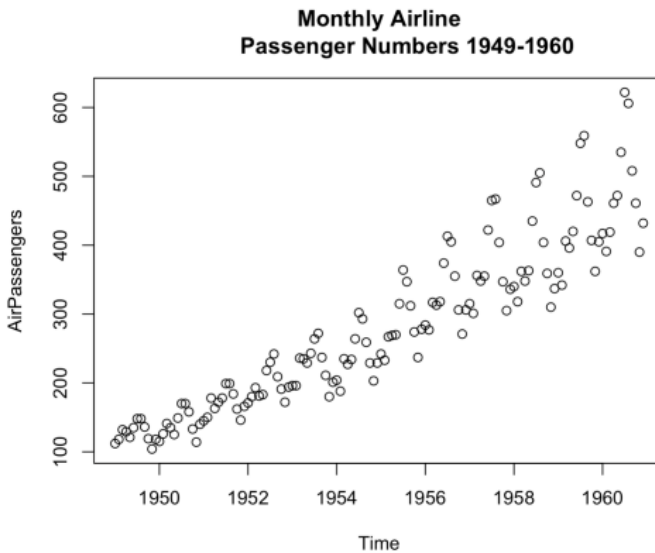


Figure: Plotting AirPassengers dataset as a scatter plot.

Applying Base Functions

- Package 'graphics' is a base package.

```
library(graphics) # to load and attach  
  
# plot  
plot(AirPassengers)  
plot(AirPassengers, type = "l", main = "Monthly Airline  
Passenger Numbers 1949-1960") # zoom
```

Applying Base Functions

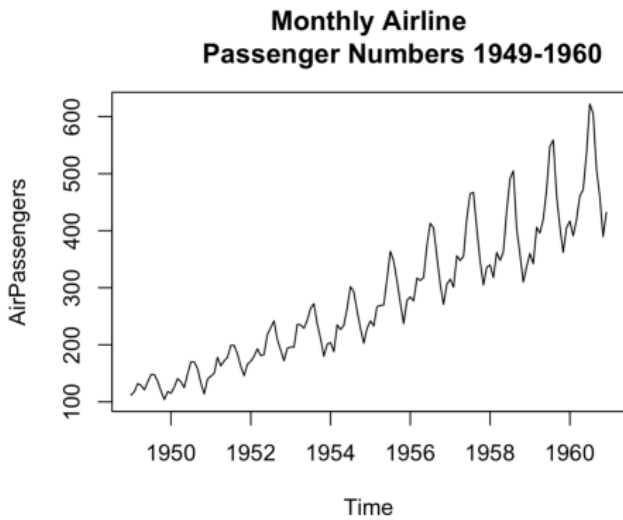


Figure: Plotting AirPassengers dataset as a line plot.

Applying Base Functions

- Package 'graphics' is a base package.

```
library(graphics) # to load and attach  
  
?AirPassengers  
# Monthly Airline Passenger Numbers 1949-1960  
  
# To have multiple plots in one overall plot using  
par(mfrow = c(1, 2))  
hist(AirPassengers) # histogram  
boxplot(AirPassengers) # boxplot
```

Applying Base Functions

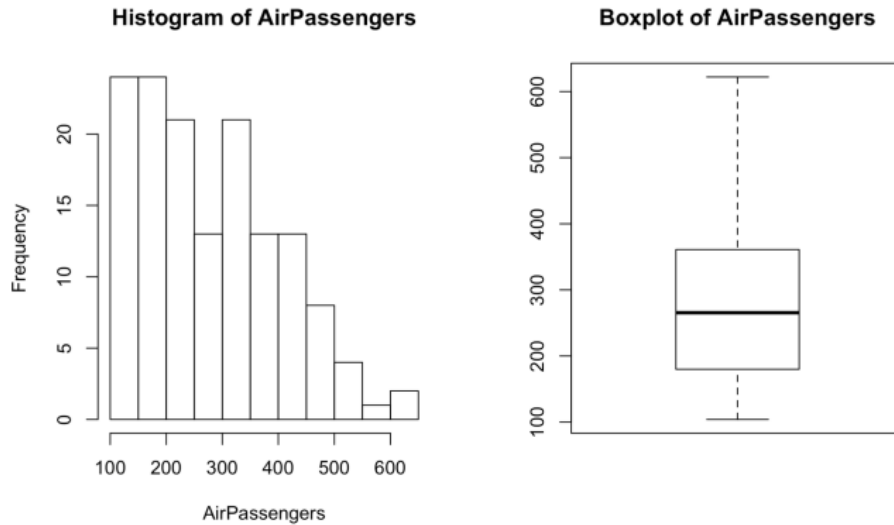


Figure: Output from plotting AirPassengers dataset.

Applying Base Functions

- Package 'graphics' is a base package.

```
library(graphics) # to load and attach  
  
?iris  
?pairs  
  
pairs(iris[, c(1:4)], col = iris$Species,  
      main = "Scatter plot of iris dataset")
```

Applying Base Functions

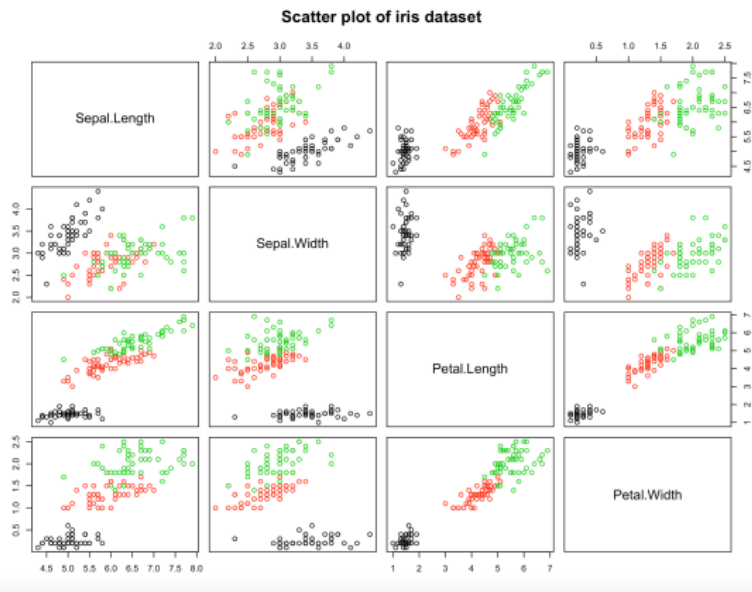


Figure: Output from plotting iris dataset as a pairs plot.

Applying Base Functions

- Package 'graphics' is a base package.

```
library(graphics) # to load and attach

barplot(as.matrix(iris[, c(1:4)]),
        legend.text = TRUE,
        main = "Bar plot of iris dataset")

# Calculate column sums of iris dataset
colSums(as.matrix(iris[, c(1:4)]))

# Sepal.Length  Sepal.Width Petal.Length  Petal.Width
# 876.5         458.6       563.7       179.9
```

Applying Base Functions

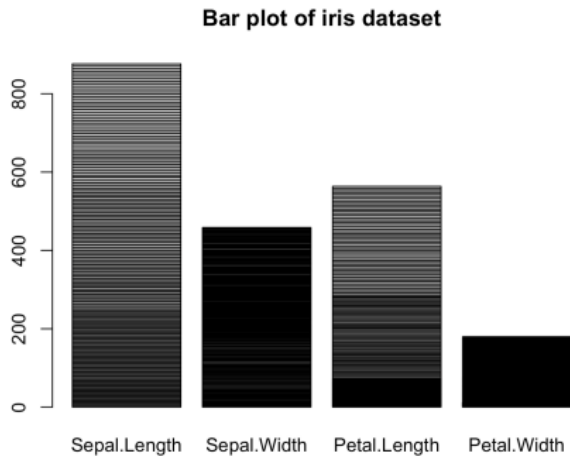


Figure: Output from plotting iris dataset as a bar plot.

R Packages for Plotting

- More resources:

- R Graphics Cookbook: <https://r-graphics.org/>.
- sthda:
<http://www.sthda.com/english/wiki/ggplot2-essentials>.



Any questions?

Question 19:

Anscombes quartet is a set of 4 x,y data sets that were published by Francis Anscombe in a 1973 paper Graphs in statistical analysis. This is available as a standard or base dataset in R.

- Obtain the dataset. What is the size of the dataset? Check the column names of the dataset. Summarise the data by calculating the mean, variance, for each column and the correlation between each pair, eg., (x_1, y_1) , (x_2, y_2) , etc. Then create a scatter plot for each x,y pair of data.

Question 19: continue...

- Anscombe's quartet has four data sets that have nearly identical descriptive statistics, but very different distributions.

| <u>Property</u> | <u>Value</u> |
|----------------------|----------------------------------|
| Mean of X (average) | 9 in all 4 XY plots |
| Sample variance of X | 11 in all four XY plots |
| Mean of Y | 7.50 in all 4 XY plots |
| Sample variance of Y | 4.122 or 4.127 in all 4 XY plots |
| Correlation (r) | 0.816 in all 4 XY plots |

- Original paper:

<https://www.jstor.org/stable/2682899?seq=1>.

More Problems (if time permits)...

Question 20:

Write a function that take user name and print the following to the user: user name, current R version and list of objects currently in memory.

Question 21:

Make a matrix as follows. Replace all numbers greater than 50 with the value of 0. How can you double check your answer? Note: this will be done using random number generator, hence the solution may differ from that of instructor.

```
randMatrix <- matrix(sample(1:100, 100, replace = TRUE),  
                      nrow = 10,  
                      ncol = 10)
```

Question 22:

Plot the ‘ToothGrowth’ dataset available in R as a scatter plot. Show dose vs tooth length. Color by supplemental type. Label the axis with correct unit (if applicable) and put a title.

- ToothGrowth

Question 23:

Dataset ‘trees’ is available on R. Use this dataset and calculate summary statistics for diameter, height and volume for Black Cherry trees. What maybe the best method of plotting all 3 variables (diameter, height and volume) using methods learned today, so the user is able to identify potential trends?

- trees



Summary

Summary

Base R

Cheat Sheet

Getting Help

- Accessing the help files**

mean
Get help of a particular function.

help.search('weighted.mean')
Search the help files for a word or phrase.

help(package = 'dplyr')
Find help for a package.

More about an object

str(iris)
Get a summary of an object's structure.

class(iris)
Find the class an object belongs to.

Using Packages

- install.packages('dplyr')**
Download and install a package from CRAN.
- library(dplyr)**
Load the package into the session, making all its functions available to use.
- dplyr::select**
Use a particular function from a package.
- data(iris)**
Load a built-in dataset into the environment.

Working Directory

- getwd()**
Find the current working directory (where inputs are found and outputs are sent).
- setwd('C://file/path')**
Change the current working directory.
- Use projects in RStudio to set the working directory to the folder you are working in.**

Vectors

| Creating Vectors | | |
|--------------------------------|-------------|-----------------------------|
| <code>c(2, 4, 6)</code> | 2 4 6 | Join elements into a vector |
| <code>2:6</code> | 2 3 4 5 6 | An integer sequence |
| <code>seq(2, 3, by=0.5)</code> | 2.0 2.5 3.0 | A complex sequence |
| <code>rep(1:2, times=3)</code> | 1 2 1 2 1 2 | Repeat a vector |
| <code>rep(1:2, each=3)</code> | 1 1 1 2 2 2 | Repeat elements of a vector |

Programming

| For Loop | |
|--|--------------|
| <code>for (variable in sequence){</code> | Do something |
| <code>}</code> | |
| <i>Example</i> | |
| <code>for (i in 1:4){</code> | |
| <code>j <- i + 10</code> | |
| <code>print(j)</code> | |
| <code>}</code> | |

While Loop

| While Loop | |
|---------------------------------|--------------|
| <code>while (condition){</code> | Do something |
| <code>}</code> | |
| <i>Example</i> | |
| <code>while (i < 5){</code> | |
| <code>print(i)</code> | |
| <code>i <- i + 1</code> | |
| <code>}</code> | |

Vector Functions

| Vector Functions | |
|-----------------------|--------------------|
| sort(x) | rev(x) |
| Return x sorted. | Return x reversed. |
| table(x) | unique(x) |
| See counts of values. | See unique values. |

If Statements

| If Statements | |
|-------------------------------------|--|
| <code>if (condition){</code> | |
| <code>do something</code> | |
| <code>} else {</code> | |
| <code>do something different</code> | |
| <code>}</code> | |
| <i>Example</i> | |
| <code>if (i > 3){</code> | |
| <code>print("Yes")</code> | |
| <code>} else {</code> | |
| <code>print("No")</code> | |
| <code>}</code> | |

Functions

| Functions | |
|---|--|
| <code>function_name <- function(var){</code> | |
| <code>do something</code> | |
| <code>} return(new_variable)</code> | |
| <i>Example</i> | |
| <code>square <- function(x){</code> | |
| <code>squared <- x*x</code> | |
| <code>return(squared)</code> | |
| <code>}</code> | |

Reading and Writing Data

| Reading and Writing Data | | |
|--|--|---|
| | | Also see the <code>readr</code> package. |
| Input | Output | Description |
| <code>df <- read.table("file.txt")</code> | <code>write.table(df, "file.txt")</code> | Read and write a delimited text file. |
| <code>df <- read.csv("file.csv")</code> | <code>write.csv(df, "file.csv")</code> | Read and write a comma separated value file. This is a special case of <code>readTable</code> / <code>writeTable</code> . |
| <code>load("file.RData")</code> | <code>save(df, file = "file.RData")</code> | Read and write an R data file, a file type specific for R. |

| Conditions | | <code>a == b</code> | <code>Are equal</code> | <code>a > b</code> | <code>Greater than</code> | <code>a >= b</code> | <code>Greater than or equal to</code> | <code>is.na(a)</code> | <code>Is missing</code> |
|-----------------------|--|---------------------|------------------------|-----------------------|---------------------------|------------------------|---------------------------------------|-------------------------|-------------------------|
| | | <code>a != b</code> | <code>Not equal</code> | <code>a < b</code> | <code>Less than</code> | <code>a <= b</code> | <code>Less than or equal to</code> | <code>is.null(a)</code> | <code>Is null</code> |
| <code>a < b</code> | | | | | | | | | |

RStudio® is a trademark of RStudio, Inc. • CC-BY What McNeil • mairihmcneil@gmail.com

Learn more at [web page](#) or [vignette](#) • package version • Updated: 3/15

Figure: Figure from <https://www.rstudio.com/resources/cheatsheets/>.

Anjali Silva

Any questions?

71/81

Summary

Figure: Figure from <https://www.rstudio.com/resources/cheatsheets/>.

Next Steps

Resources

- The R Journal: <https://journal.r-project.org/>
- R for Data Science textbook (free): <https://r4ds.had.co.nz/>.
- RStudio Education Series (free):
<https://education.rstudio.com/learn/>
- Bioconductor Workshops:
<https://www.bioconductor.org/help/events/>
- Coursera offers courses via both paid + free options
 - <https://www.coursera.org/learn/r-programming>

Further Readings

- Hackenberger, B. K. (2020) R software: unfriendly but probably the best. *Croatian Medical Journal* 61. URL:
<https://www.ncbi.nlm.nih.gov/pmc/articles/PMC7063554/>.
- Gentleman, R. C., Carey, V. J., Bates, D. M. et al. (2004) Bioconductor: open software development for computational biology and bioinformatics. *Genome Biol* 5(R80). URL:
<https://doi.org/10.1186/gb-2004-5-10-r80>.
- Huber W., Carey V. J., Gentleman R. et al. (2015) Orchestrating high-throughput genomic analysis with Bioconductor. *Nat Methods*. 12(2). URL:
<https://www.ncbi.nlm.nih.gov/pmc/articles/PMC4509590/>.

Resources



R packages for data science

The tidyverse is an opinionated **collection of R packages** designed for data science. All packages share an underlying design philosophy, grammar, and data structures.

Install the complete tidyverse with:

```
install.packages("tidyverse")
```

Figure: Tidyverse R Packages designed for data science. Figure from <https://www.tidyverse.org/>.

Conferences

useR! — International R User Conference



useR! 2022 will be a hybrid conference from June 20 to June 23, with opportunities to participate as an online attendee or to attend in person at Vanderbilt University Medical Center, Nashville, TN, USA. For the latest updates, follow '[@_useRconf](#)' on Twitter, or '[user-conf](#)' on LinkedIn.

Conferences



The image shows the homepage of the BioC2022 conference website. At the top left is the Bioconductor logo. To its right is a navigation bar with links: HOME (which is highlighted in blue), NEWS, REGISTRATION, SPONSORS, SCHEDULE, and ABOUT. Below the navigation bar is a large banner featuring the text "BIOCONDUCTOR CONFERENCE, JULY 27-29, 2022" in white, with "WHERE SOFTWARE AND BIOLOGY CONNECT" underneath. To the right of the text is a hexagonal logo for "BioC 2022" featuring a stylized cityscape with the Space Needle. The background of the banner has faint, overlapping text from previous BioC conferences like "EuroBioC2017", "BioC Asia 2016", and "BioC 2019".

Registration for the BioC2022 conference is now open!

BioC2022 is organized as a hybrid event. This includes in-person gathering in Seattle (Seattle Children's Hospital Cure Building, 1920 Terry Ave, Seattle, WA 98101) and online participation.

Seattle's [conference hotels, restaurants, activities, transportation](#).

User Groups and Meetings

- R User Groups <https://jumpingrivers.github.io/meetingsR/>
 - Greater Toronto Area (GTA) R User Group: <https://www.meetup.com/Greater-Toronto-Area-GTA-R-Users-Group/>
 - Toronto R-Ladies: <https://www.meetup.com/r ladies-toronto/>
- SciNet Virtual Summer Training Program:
https://support.scinet.utoronto.ca/education/go.php/573/index.php/ib/1//p_course/573
- R virtual events: <https://jumpingrivers.github.io/meetingsR/virtual-events.html>

Data Revolution



Regulating the internet giants

The world's most valuable
resource is no longer oil, but
data



Figure: *Economist*, 2017.

Feedback

- Take the time to provide feedback (anonymous):
<https://forms.office.com/r/Dne92Br8US>
- Thank you.