Untitled

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MiniCourse R packages Day2

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Day 2 contents

- Bioconductor
 - Core packages
 - Core classes
 - Release cycle
 - Submission process
- ► GitHub
 - Package hosting
 - Continuous integration
 - GitHub actions
- Practical: package website with pkgdown

Bioconductor

Bioconductor (or just "Bioc") is "open source software for bioinformatics", based on more than 2000 R packages. It is old enough to have its own Wikipedia page!

It can be installed... from CRAN.

```
if (!requireNamespace("BiocManager", quietly = TRUE))
    install.packages("BiocManager")
BiocManager::install(version = "3.13")
```

Hold on if you are not bioinformatician, we will use it to learn more about *S4* classes.

Bioconductor core packages

Canonical URLs:

```
The core Bioconductor packages provide core classes that you will
find being used in almost every other packages.
# Attaches a lot of stuff !!!
BiocManager::install("GenomicRanges")
library("GenomicRanges")
# Use this in your R Markdown files...
suppressPackageStartupMessages(library("GenomicRanges"))
# Or since R4.1
library("GenomicRanges") |> suppressPackageStartupMessages
```

https://bioconductor.org/packages/nameOfThePackage

Output of library("GenomicRanges")

Note the difference between "loading" and "attaching"
> library("GenomicRanges")

Restarting R session...

> library("GenomicRanges")
Loading required package: stats4

Loading required package: BiocGenerics Loading required package: parallel

Attaching package: 'BiocGenerics'

The following objects are masked from 'package:parallel':

clusterApply, clusterApplyLB, clusterCall, clusterEval(
parApply, parCapply, parLapply, parLapplyLB, parRapply

The following objects are masked from 'package:stats':

Bioconductor core classes

- Use the S4 class system.
- Often names start with capital letters.
- Some of the core classes are in the core packages that have cryptic names.
- Today I will not cover annotation databases or genome objects.

DataFrame

151

class: small-code

```
DataFrame()
```

DataFrame with 0 rows and 0 columns

as	airqualit	y, "Da	atarı	rame",)		
##	DataFrame	with	153	rows	and	6	columns

Solar R Wind ## Nzone teger> <integer> <:

	0_0110	202011		
##	<integer></integer>	<integer></integer>	<numeric></numeric>	<int< th=""></int<>

##		<integer></integer>	<integer></integer>	<numeric></numeric>	<int< th=""></int<>
##	1	Δ1	190	7 Δ	

##	1	41	190	7.4
##	2	36	118	8.0

##	1	41	190	1.4
##	2	36	118	8.0

2 36 118 8.0		-		100	
	ŧ		36	118	8.0

##	2	36	118	8.0

##	2	36	118	8.0
шш	2	10	1.40	10 6

14

# 2	36	118	8.0	
# 3	10	1/0	12.6	

## 3	12	149	12.6	74	5
## /	10	212	11 5	60	_

Temp

67

75

Month

5

## 3	12	149	12.6	74	5
## 4	18	313	11.5	62	5

## 4	18	313	11.5	62	5
## 5	NA	NA	14.3	56	5

## 4	10	313	11.5	02	5
## 5	NA	NA	14.3	56	5

## 4	18	313	11.5	62	t
## 5	NA	NA	14.3	56	Ę

150 NA13.2 77 145

14.3

191

Rle

class: small-code

Run-length encoding (Rle) is a simple and efficient way to compress data.

```
Rle(rpois(1e7, 1e-3))
```

A data structure that I like is a DataFrame of Rle values.

```
airquality |> lapply(Rle) |> DataFrame() |> head(3)
```

```
## DataFrame with 3 rows and 6 columns
    Ozone Solar.R Wind Temp Month
##
           <Rle> <Rle> <Rle> <Rle> <Rle>
##
    <R.1e>
## 1
      41
             190 7.4 67
## 2
    36
             118
                   8 72
## 3
      12
             149 12 6
                        74
```

Other important classes

- GRanges
- ► SummarizedExperiment
- ► MultiAssayExperiment

A few words on the S4 class system

- polymorphic and functional: core function names dispatch on different methods according to the class of their arguments.
- Objects have "slots" accessed with the @ sign.
- copy-on-write / copy-on-modify semantics.
- Useful as data structure but also for type safety

Learn more with https://adv-r.hadley.nz/s4.html

Let's extend a S4 class

It is strongly recommended to re-use the core Bioc classes in packages designed for Bioconductor.

```
library(methods)
setClass("BetterList", contains = "SimpleList")
setMethod("show", "BetterList", function(object) {
   callNextMethod()
   cat("This superior version of the SimpleList class is brown of the Simple class class is brown of the Simple class cl
```

This superior version of the SimpleList class is brough

It also works on S3 classes

##

It is a matter of taste whether to do so or not.

```
setClass("BetterList2", contains = "list")
setMethod("show", "BetterList2", function(object) {
   callNextMethod()
   cat("This superior version of the list class is also brow
})
list(c=3) |> as("BetterList2")

## An object of class "BetterList2"

## [[1]]
## [1] 3
```

This superior version of the list class is also brought

Use for type safety

```
setClass("ListOfChars", contains = "SimpleList", validity =
  # Actually not so safe, what is the list contains sublis
  all(sapply(object, is.character))
})
setMethod("show", "ListOfChars", function(object) {
  callNextMethod()
  cat("This safer version of the SimpleList class is surely
})
SimpleList("haha", "hoho") |> as("ListOfChars") |> validOb
## [1] TRUE
# Try this!
# SimpleList("haha", 1) |> as("ListOfChars") |> validObjec
```

When to Depend or Import Bioc packages

- ▶ If you use their functions internally, Import the packages in the DESCRIPTION file and import their functions in NAMESPACE. Typical examples: IRanges, S4Vectors.
- ▶ If you want their functions to be easily available to their users, just Depend on the packages in the DESCRIPTION file. Typycal examples: GenomicRanges, SummarizedExperiment, ggplot2.

Tip for easier debugging of your package. class: small-code

Do the ground work in a S3 function, and wrap it in the S4 system.

```
setGeneric("countRows", function(x) standardGeneric("countle
```

```
## [1] "countRows"
.countRows <- function(x) {
  if (nrow(x) > 0) cat("Owow, there are rows \n")
  cat("I found ", nrow(x), "rows.\n")
  cat("I think I finished counting\n")
```

```
cat("I found ", nrow(x), "rows.\n")
cat("I think I finished counting\n")
}
setMethod("countRows", "DataFrame", .countRows)
airquality |> DataFrame() |> countRows()
## Owow, there are rows
```

```
## I found 153 rows.
## I think I finished counting
And now you can set the debugger to jump straight in your code
```

More on Bioconductor

- Releases twice a year, shortly after R releases.
- Maintains a release and a devel branch in parallel. Version numbers are even and odd respectively. http://www.bioconductor.org/developers/how-to/versionnumbering/
- Support site: https://support.bioconductor.org/
- Browse all soure code at https://code.bioconductor.org/
- Submission process takes place on GitHub by oppening an issue on Bioconductor/Contributions

GitHub

- ▶ Not the only hosting platform, there is also GitLab, etc.
- Useful to manage source code of course, but also
- There are functions to install a package directly from a GitHub repository.

remotes::install_github("user/repo"), remotes::install_bio

Reminder: never use passowrdless SSH keys, use ssh-add instead (you might need to run eval \$(ssh-agent) first).
 Maye your keys transiently available on Deigo with ssh -A.

Get a Zenodo DOI for packages via Git repositories

CRAN or Bioconductor, and journals provide DOIs. How about non-peer-reviewed packages ?

You can get DOIs from Zenodo, or alternatively Dryad, figshare, etc.

Zenodo has a good integration with GitHub. Random example. Documentation

GitHub actions

GitHub actions for regression testing: example with the charles-plessy/CAGEr package.

usethis::use_github_action()

biocthis::use_bioc_github_action()

Pkgdown

GitHub actions are also used for building GitHub pages.

The website of pkgdown is of course built with pkgdown, so let's have a look.

https://pkgdown.r-lib.org/

https://pkgdown.r-lib.org/articles/linking.html

Example with the oist/GenomicBreaks package

Now, let's try it together!