Getting to program in ${\bf R}$

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Programming with ${\bf R}$

- Writing simple functions
- Why programming
- Coding basics
- A few coding tips
- 5 Example: comparing the performances of 2 tests
- Exploring existing code
- Debugging
- Optimisation & Profiling
- Writing R packages
- Writing more advanced functions
- Object systems in F

The best way:

```
my_function <- function(input1, input2) {
  output <- input1 + input2
  return(output)
}

my_function(input1 = 1, input2 = 3)
## [1] 4</pre>
```

The best way:

```
my_function <- function(input1, input2) {
  output <- input1 + input2
  return(output)
}

my_function(input1 = 1, input2 = 3)
## [1] 4</pre>
```

If no return(), then it returns the last row:

```
my_function <- function(input1, input2) {
  input1 + input2
}

my_function(input1 = 1, input2 = 3)
## [1] 4</pre>
```

The best way:

```
my_function <- function(input1, input2) {
  output <- input1 + input2
  return(output)
}

my_function(input1 = 1, input2 = 3)
## [1] 4</pre>
```

If no return(), then it returns the last row:

```
my_function <- function(input1, input2) {
  input1 + input2
}

my_function(input1 = 1, input2 = 3)
## [1] 4</pre>
```

Inline shortcut (useful in *apply()):

```
my_function <- function(input1, input2) input1 + input2
my_function(input1 = 1, input2 = 3)
## [1] 4</pre>
```

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You can set defaults for your inputs:

```
my_function <- function(input1, input2 = 10) {
    output <- input1 + input2
    return(output)
}

my_function(input1 = 1)
## [1] 11
my_function(input1 = 1, input2 = 0)
## [1] 1
my_function(input2 = 1)
## Error in my_function(input2 = 1): argument "input1" is missing, with no default</pre>
```

You can pass optional arguments to another function via "...":

```
my_function <- function(input1, ...) {
  output <- mean(input1, ...)
  return(output)
}

x <- c(1, 2, 3, 4, 5, NA)
my_function(input1 = x)
## [1] NA
my_function(input1 = x, na.rm = TRUE)
## [1] 3</pre>
```

Note: this is particularly useful when designing plotting functions!

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You can print things while the function runs:

```
my_function <- function() {
    print("This function will output 2")
    return(2)
}

my_function()
## [1] "This function will output 2"
## [1] 2</pre>
```

Note: observe also that it is possible to create functions that do not consider any input!

You can return only a single object; so if you need several outputs, you must combine them:

```
my_function <- function(input1, ...) {
  output2 <- max(input1, ...)
  output2 <- max(input1, ...)
  output <- list(output1 = output1, output2 = output2)
  return(output)
}

x <- c(1, 2, 3, 4, 5, NA)
my_function(input1 = x, na.rm = TRUE)

## $output1
## [1] 1
##
##
$output2
## [1] 5</pre>
```

Note: you can also use a vector, a matrix, a data frame...

You can return the output "invisibly":

```
my_function <- function(input1) {
  output <- min(input1)
  return(invisible(output))
}
my_function(input1 = c(1, 2, 3, -4))</pre>
```

You can return the output "invisibly":

```
my_function <- function(input1) {
  output <- min(input1)
  return(invisible(output))
}

my_function(input1 = c(1, 2, 3, -4))</pre>
```

```
foo <- my_function(input1 = c(1, 2, 3, -4))
foo
## [1] -4</pre>
```

You don't have to return something:

```
my_function <- function(input1) {
  the_min <- min(input1)
  print(paste("The minimum is:", the_min))
  return(invisible(NULL))
}

my_function(input1 = c(1, 2, 3, -4))
## [1] "The minimum is: -4"</pre>
```

You don't have to return something:

```
my_function <- function(input1) {
  the_min <- min(input1)
  print(paste("The minimum is:", the_min))
  return(invisible(NULL))
}

my_function(input1 = c(1, 2, 3, -4))
## [1] "The minimum is: -4"</pre>
```

```
foo <- my_function(input1 = c(1, 2, 3, -4))
## [1] "The minimum is: -4"
foo
## NULL</pre>
```

Programming with \boldsymbol{R}

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- 2 Why programming?
- Coding basics
- 4 A few coding tips
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Why writing your own **R** functions?

Using your own functions makes your scripts

- easier to understand
- safer to (re)use
- shorter to write (often)

```
d <- data.frame(proba = c(0.1, 0.5, 0.4), group = factor(c("A", "B", "C")))
with(data = d, (proba[group == "B"] / (1 - proba[group == "B"])) / (proba[group == "A"] / (1 - proba[group == "A"])))
## [1] 9
with(data = d, (proba[group == "C"] / (1 - proba[group == "C"])) / (proba[group == "A"] / (1 - proba[group == "A"])))
## [1] 6
with(data = d, (proba[group == "B"] / (1 - proba[group == "B"])) / (proba[group == "C"] / (1 - proba[group == "C"])))
## [1] 1.5</pre>
```

```
d <- data.frame(proba = c(0.1, 0.5, 0.4), group = factor(c("A", "B", "C")))

vith(data = d, (proba[group == "B"] / (1 - proba[group == "B"])) / (proba[group == "A"] / (1 - proba[group == "A"])))

## [1] 9

with(data = d, (proba[group == "C"] / (1 - proba[group == "C"])) / (proba[group == "A"] / (1 - proba[group == "A"])))

## [1] 6

with(data = d, (proba[group == "B"] / (1 - proba[group == "B"])) / (proba[group == "C"] / (1 - proba[group == "C"])))

## [1] 1.5</pre>

Or

odds_ratio <- function(group1, group2){
    vith(data = d, (proba[group == group1]) / (1 - proba[group == group2])) / (proba[group == group2] / (1 - proba[group == group2])))</pre>
```

odds_ratio(group1 = "B", group2 = "A")

odds ratio(group1 = "C", group2 = "A")

odds_ratio(group1 = "B", group2 = "C")

[1] 9

[1] 6

[1] 1.5

Still not convinced? Let's compute all pairwise comparisons:

```
with(data = d, (proba[group == "A"] / (1 - proba[group == "A"])) / (proba[group == "A"] / (1 - proba[group == "A"])))
## [1] 1
with(data = d, (proba[group == "B"] / (1 - proba[group == "B"])) / (proba[group == "A"] / (1 - proba[group == "A"])))
## [1] 9
with(data = d, (proba[group == "C"] / (1 - proba[group == "C"])) / (proba[group == "A"] / (1 - proba[group == "A"])))
## [1] 6
with(data = d, (proba[group == "A"] / (1 - proba[group == "A"])) / (proba[group == "B"] / (1 - proba[group == "B"])))
## [1] 0.1111111
with(data = d, (proba[group == "B"] / (1 - proba[group == "B"])) / (proba[group == "B"] / (1 - proba[group == "B"])))
## [1] 1
with(data = d, (proba[group == "C"] / (1 - proba[group == "C"])) / (proba[group == "B"] / (1 - proba[group == "B"])))
## [1] 0.6666667
with(data = d, (proba[group == "A"] / (1 - proba[group == "A"])) / (proba[group == "C"] / (1 - proba[group == "C"])))
## [1] 0.1666667
with(data = d, (proba[group == "B"] / (1 - proba[group == "B"])) / (proba[group == "C"] / (1 - proba[group == "C"])))
## [1] 1.5
with(data = d, (proba[group == "C"] / (1 - proba[group == "C"])) / (proba[group == "C"] / (1 - proba[group == "C"])))
## [1] 1
```

Still not convinced? Let's compute all pairwise comparisons:

```
for (group2 in d$group) {
    for (group1 in d$group) {
        print(paste(group1, group2, odds_ratio(group1 = group1, group2 = group2)))
    }
}

## [1] "A A 1"
## [1] "B A 9"
## [1] "C A 6"
## [1] "B B 1"
## [1] "B B 1"
## [1] "B C 0.6666666666666"
## [1] "A C 0.16666666666666"
## [1] "B C 1.5"
## [1] "C C 1"
```

Note: this is bad code, we will come back on this!

When to write your own functions?

Don't Repeat Yourself

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Control structures: if()

```
i <- 1
a <- 2
if (i == 1) {
  a <- 1
## [1] 1
```

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Control structures: if()

```
i <- 1
a <- 2
if (i == 1) {
  a <- 1
## [1] 1
i <- 5
a <- 2
if (i == 1) {
  a <- 1
} else {
  a <- 2
## [1] 2
```

Note: for help, check ?"if".

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Control structures: loops using for() & while()

```
for (i in 1:5) {
  print(i)
## [1] 1
## [1] 2
## [1] 3
## [1] 4
## [1] 5
i <- 1
while (i < 5) {
  print(i)
  i <- i + 1
## [1] 1
## [1] 2
## [1] 3
```

Note: for help, check ?"for".

[1] 4

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Random number generators

```
runif(5)
## [1] 0.6280441 0.7862230 0.7450039 0.5514726 0.3859306
runif(5)
## [1] 0.4672231 0.5400778 0.9531008 0.8941555 0.4969132
```

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Random number generators

```
runif(5)
## [1] 0.6280441 0.7862230 0.7450039 0.5514726 0.3859306
runif(5)
## [1] 0.4672231 0.5400778 0.9531008 0.8941555 0.4969132
```

You can have reproducible results by setting a seed:

```
set.seed(10132)
runif(5)
## [1] 0.7258731 0.3086039 0.4393603 0.7952702 0.5325773
set.seed(10132)
runif(5)
## [1] 0.7258731 0.3086039 0.4393603 0.7952702 0.5325773
```

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Random number generators

```
runif(5)
## [1] 0.6280441 0.7862230 0.7450039 0.5514726 0.3859306
runif(5)
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You can have reproducible results by setting a seed:

```
set.seed(10132)
runif(5)
## [1] 0.7258731 0.3086039 0.4393603 0.7952702 0.5325773
set.seed(10132)
runif(5)
## [1] 0.7258731 0.3086039 0.4393603 0.7952702 0.5325773
```

Note: check ?Distributions for more distributions.

```
process_factor <- function(x) {
  if (!is.factor(x) & !is.character(x)) {
    stop("Your input for factor should be of class character or factor")
}
  if (!is.factor(x) & is.character(x)) {
    x <- as.factor(x)
    warning("Your input has been converted to factor")
}
  return(x)
}</pre>
```

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```
process_factor <- function(x) {
  if (!is.factor(x) & !is.character(x)) {
    stop("Your input for factor should be of class character or factor")
}
  if (!is.factor(x) & is.character(x)) {
    x <- as.factor(x)
    warning("Your input has been converted to factor")
}
  return(x)
}</pre>
```

```
process_factor(x = factor(c("A", "B")))
## [1] A B
## Levels: A B
```

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process_factor <- function(x) {</pre>

Levels: A B

```
if (!is.factor(x) & !is.character(x)) {
    stop("Your input for factor should be of class character or factor")
  if (!is.factor(x) & is.character(x)) {
    x <- as.factor(x)
    warning("Your input has been converted to factor")
  return(x)
process factor(x = factor(c("A", "B")))
## [1] A B
## Levels: A B
process_factor(x = c("A", "B"))
## Warning in process_factor(x = c("A", "B")): Your input has been converted to factor
## [1] A B
```

```
process factor <- function(x) {
  if (!is.factor(x) & !is.character(x)) {
    stop("Your input for factor should be of class character or factor")
  if (!is.factor(x) & is.character(x)) {
    x <- as.factor(x)
    warning("Your input has been converted to factor")
  return(x)
process factor(x = factor(c("A", "B")))
## [1] A B
## Levels: A B
process_factor(x = c("A", "B"))
## Warning in process_factor(x = c("A", "B")): Your input has been converted to factor
## [1] A B
## Levels: A B
process_factor(x = c(2, 3))
## Error in process factor(x = c(2, 3)): Your input for factor should be of class character or factor
```

Boolean logic in R

! TRUE ## [1] FALSE

```
!FALSE
## [1] TRUE
TRUE == 1
## [1] TRUE
TRUE & TRUE
## [1] TRUE
TRUE & FALSE
## [1] FALSE
FALSE & FALSE
## [1] FALSE
c(TRUE, TRUE) & c(TRUE, FALSE)
## [1] TRUE FALSE
FALSE && "foo"
## [1] FALSE
FALSE & "foo"
## Error in FALSE & "foo": operations are possible only for numeric,
logical or complex types
TRUE && "foo"
## Error in TRUE && "foo": invalid 'y' type in 'x && y'
c(TRUE, TRUE) && c(TRUE, FALSE) ## dangerous
## [1] TRUE
```

```
TRUE | TRUE
## [1] TRUE
TRUE | FALSE
## [1] TRUE
FALSE | FALSE
## [1] FALSE
c(TRUE, TRUE) | c(TRUE, FALSE)
## [1] TRUE TRUE
FALSE || "foo"
## Error in FALSE || "foo": invalid 'y' type in 'x || y'
TRUE | | "foo"
## [1] TRUE
any(c(FALSE, FALSE, FALSE, TRUE))
## [1] TRUE
```

Note: help available at ?"&".

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Scope

Every object in **R** belongs to an environment:

```
i <- 1
i <- i + 1
i
## [1] 2
```

```
e1 <- environment()
e1
## <environment: R_GlobalEnv>
get("i", envir = e1)
## [1] 2
```

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Scope

As a rule, anything created inside a function stays inside the function:

```
i <- 1
f <- function(i) {
   i <- i + 1
   return(i)
   }
f(i)
## [1] 2
i
## [1] 1</pre>
```

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i <- 1

This is because the functions have their own environments:

```
f <- function(i) {
    i <- i + 1
    e <- environment()
    return(list(env = e, i = i))
    }

res <- f(i)
res$i
## [1] 2
res$env
## <environment: 0x7febd99c24e0>
```

i <- 1

This is because the functions have their own environments:

```
f <- function(i) {
    i <- i + 1
    e <- environment()
    return(list(env = e, i = i))
    }

res <- f(i)
res$i
## [1] 2
res$env
## <environment: 0x7febd99c24e0>
```

```
get("i", envir = globalenv()) ## globalenv() provides the address of the global environment
## [1] 1
get("i", envir = res$env)
## [1] 2
```

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i <- 1

f <- function(i) { i <- i + 1

get("i", envir = res\$env)

[1] 2

This is because the functions have their own environments:

```
e <- environment()
 return(list(env = e, i = i))
res <- f(i)
res$i
## [1] 2
res$env
## <environment: 0x7febd99c24e0>
get("i", envir = globalenv()) ## qlobalenv() provides the address of the qlobal environment
## [1] 1
```

Challenge: check whether the environment remains the same at each function call or not!

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Environments can be hacked, but it is usually source of troubles:

```
i <- 1
f <- function(i) {
   i <<- i + 1
   return(i)
   }
f(i)
## [1] 1
i
## [1] 2</pre>
```

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Environments can be defined, but it is usually not necessary:

```
i <- 1
f <- function(i) {
    i <- i + 1
    assign("i", i, envir = globalenv())
    return(i)
    }
f(i)
## [1] 2
i
## [1] 2</pre>
```

The best is to use a functional approach:

```
i <- 1
f <- function(i) {
    i <- i + 1
    return(i)
    }
i <- f(i)
i
## [1] 2</pre>
```

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There are a few scoping exceptions:

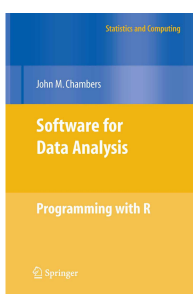
```
i <- 1
for (j in 1:10) {
   i <- i + 1
}
i
## [1] 11</pre>
```

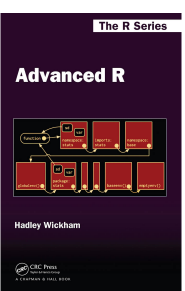
And yet, for is actually a function:

```
i <- 1
`for`(j, 1:10, i <- i + 1)
i
## [1] 11</pre>
```

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For more info:





```
x < -0.7 - 0.4 - 0.3
x == 0
## [1] FALSE
```

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```
x < -0.7 - 0.4 - 0.3
x == 0
## [1] FALSE
print(x, digits = 22)
## [1] -5.551115123125782702118e-17
```

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[3] 0.200000000000000111022 0.3000000000000000444089 [7] 0.600000000000000888178 0.700000000000000666134 [9] 0.8000000000000000444089 0.9000000000000000222045

x < -0.7 - 0.4 - 0.3

x == 0

```
## [1] FALSE
print(x, digits = 22)
## [1] -5.551115123125782702118e-17
print(seq(0, 1, 0.1), digits = 22)
```

```
Note 1: same kind of thing can happen in Excel too (https://support.microsoft.com/en-us/kb/214118).
Note 2: this kind of problem sometimes kills people
```

(http://www-users.math.umn.edu/~arnold/disasters/Patriot-dharan-skeel-siam.pdf).

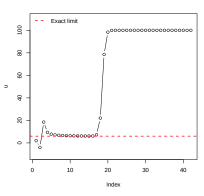
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The errors sometimes add up:

Example of the J.M Muller's Serie

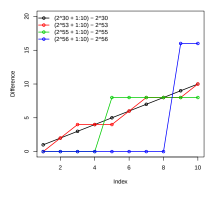
$$u_0 = 2$$
; $u_1 = -4$; $u_{n+1} = 111 - \frac{1130}{u_n} + \frac{3000}{u_n * u_{n-1}}$

```
u \leftarrow c(2, -4)
new.u <- function(u) 111 - 1130/u[length(u)] + 3000/(u[length(u)]*u[length(u) - 1])</pre>
for (i in 1:40) u \leftarrow c(u, new.u(u))
```



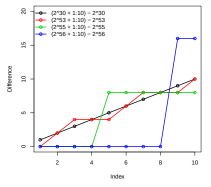
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And it is not just about fractions:



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And it is not just about fractions:



```
as.integer(2~31 - 1) ## upper limit in 32 bits coding
## [1] 2147483647
as.integer(2~31)
## Warning: NAs introduced by coercion to integer range
## [1] NA
```

??"equality"

Solution: beware of floats and use adequate functions, not boolean tests, when performing comparisons!

```
Help pages:
base::all.equal
                              Test if Two Objects are (Nearly) Equal
base::identical
                              Test Objects for Exact Equality
data.table::all.equal
                              Equality Test Between Two Data Tables
datasets::airquality
                              New York Air Quality Measurements
dplyr::all_equal
                              Flexible equality comparison for data frames
all.equal(target = 0, current = x)
## [1] TRUE
```

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Some key advices

- everything you use in the body must pass through the inputs
- everything you output must pass through the return
- try to write functions that you could reuse in other situations

Some key advices

- everything you use in the body must pass through the inputs
- everything you output must pass through the return
- try to write functions that you could reuse in other situations

Bad:

```
dd <- data.frame(proba = c(0.1, 0.5, 0.4), group = factor(c("A", "B", "C")))

odds_ratio <- function(group1, group2){
    with(data = d, (proba[group == group1] / (1 - proba[group == group1])) / (proba[group == group2] / (1 - proba[group == group2])))
}

for (group2 in d$group) {
    for (group1 in d$group) {
        print(paste(group1, group2, odds_ratio(group1 = group1, group2 = group2)))
    }
}</pre>
```

Why is this bad?

A better implementation of functions to compute odd ratios

Better because it is general and based on functions (example 1):

```
odds_ratio <- function(p1, p2){
  OR \leftarrow (p1/(1 - p1)) / (p2/(1 - p2))
 return(OR)
all pairwise <- function(proba, groups){
  groups_id <- unique(groups)</pre>
  results <- matrix(NA, ncol = length(groups id), nrow = length(groups id))
  colnames(results) <- groups_id</pre>
  rownames(results) <- groups_id</pre>
  for (group1 in groups_id) {
   for (group2 in groups_id) {
      results[group1, group2] <- odds_ratio(p1 = proba[groups == group1], p2 = proba[groups == group2])
  return(results)
all pairwise(proba = d$proba, groups = d$group)
## A 1 0.1111111 0.1666667
## B 9 1.0000000 1.5000000
## C 6 0.6666667 1.0000000
```

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A better implementation of functions to compute odd ratios

Better because it is very general and based on functions (example 2):

```
odds_ratio <- function(p1, p2){
  OR \leftarrow (p1/(1 - p1)) / (p2/(1 - p2))
 return(OR)
all_pairwise2 <- function(proba, groups){
  groups_id <- unique(groups)</pre>
  to do <- expand.grid(groups id, groups id)
  OR <- apply(to_do, 1, function(gr) { ## define anonymous function
    odds ratio(p1 = proba[groups == gr[1]], p2 = proba[groups == gr[2]])
  return(data.frame(group1 = to_do[, 1],
                    group2 = to_do[, 2],
                    OR = OR)
all_pairwise2(proba = d$proba, groups = d$group)
     group1 group2
## 1
                 A 1.0000000
## 2
                 A 9.0000000
                A 6.0000000
## 3
                 B 0.1111111
## 5
                 B 1.0000000
## 6
                B 0.6666667
## 7
                 C 0.1666667
## 8
                 C 1.5000000
## 9
                 C 1,0000000
```

Some key advices

- everything you use in the body must pass through the inputs
- everything you output must pass through the return
- try to write functions that you could reuse in other situations
- there are many ways to reach the same outcome; experiment a bit to find something you like/understand

Programming with \boldsymbol{R}

- Writing simple functions
- Why programming
- Coding basics
- 4 A few coding tips
- 5 Example: comparing the performances of 2 tests
- 6 Exploring existing code
- Debugging
- Optimisation & Profiling
- Writing R packages
- Writing more advanced functions
- Object systems in F

Challenge

The t-test and the Mann-Whitney U tests are two tests aiming at comparing 2 groups. We want to compare the risk of false positives and true positives of these two tests in the following conditions:

- assuming that height of males is Gaussian with mean 180 cm and SD 6 cm and that
 the height of females is Gaussian with mean 170 cm and SD 5 cm, how many
 individuals (sex-ratio = 1) do I need to get a power of 80% (risk of false negative =
 20%)?
- assuming that the null hypothesis is true and that you have sampled 20 males and 20 females, what is the probability of false positives for the threshold alpha = 0.05? (And for any threshold between 0 and 1%?)

Programming with ${\bf R}$

- Writing simple functions
- Why programming
- Coding basics
- 4 A few coding tips
- 5 Example: comparing the performances of 2 tests
- 6 Exploring existing code
- Debugging
- Optimisation & Profiling
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Learning by mimicking

Looking at code written by others will teach you

- how their functions work
- how to code
- new functions or packages that could be useful for you

Start by simply typing the function name (without brackets):

```
mosaic::oddsRatio
## function (x, conf.level = 0.95, verbose = !quiet, quiet = TRUE,
## digits = 3)
## {
## orrr(x, conf.level = conf.level, verbose = verbose, digits = digits,
## relrisk = FALSE)
## }
## <br/>
## <converge = verbose = verbo
```

Then, follows the successive calls:

```
mosaic::orrr
## function (x, conf.level = 0.95, verbose = !quiet, quiet = TRUE,
       digits = 3, relrisk = FALSE)
## {
##
       if (any(dim(x) != c(2, 2))) {
##
           stop("expecting something 2 x 2")
##
       names(x) <- NULL.
       row.names(x) <- NULL
##
       colnames(x) <- NULL
##
       rowsums <- rowSums(x)
       p1 <- x[1, 1]/rowsums[1]
       p2 <- x[2, 1]/rowsums[2]
       o1 <- p1/(1 - p1)
       o2 \leftarrow p2/(1 - p2)
##
       RR <- p2/p1
##
       OR <- o2/o1
       crit <- qnorm((1 - conf.level)/2, lower.tail = FALSE)</pre>
       names(RR) <- "RR"
##
##
       log.RR <- log(RR)
       SE.log.RR <- sqrt(sum(x[, 2]/x[, 1]/rowsums))
       log.lower.RR <- log.RR - crit * SE.log.RR
##
##
       log.upper.RR <- log.RR + crit * SE.log.RR
##
       lower.RR <- exp(log.lower.RR)</pre>
##
       upper.RR <- exp(log.upper.RR)
##
       names(OR) <- "OR"
       log.OR <- log(OR)
       SE.log.OR <- sqrt(sum(1/x))
       log.lower.OR <- log.OR - crit * SE.log.OR
##
##
       log.upper.OR <- log.OR + crit * SE.log.OR
       lower.OR <- exp(log.lower.OR)
##
```

Sometimes, the code is not directly displayed...e.g. **R** methods (S3):

```
residuals

## function (object, ...)

## UseMethod("residuals")

## Oytecode: 0x7febd83a07f0>

## <environment: namespace:stats>
```

residuals() is a *generic* function which rely on class specific *methods*:

```
methods(residuals)
    [1] residuals.default*
                                 residuals.glm
    [3] residuals.gls*
                                 residuals.glsStruct*
    [5] residuals.gnls*
                                 residuals.gnlsStruct*
    [7] residuals.HoltWinters*
                                 residuals.isoreg*
    [9] residuals.lm
                                 residuals.lme*
  [11] residuals.lmeStruct*
                                 residuals.lmList*
  [13] residuals.loglm*
                                 residuals nlmeStruct*
## [15] residuals.nls*
                                 residuals.psych*
## [17] residuals.smooth.spline* residuals.tukeyline*
## see '?methods' for accessing help and source code
```

The methods with a * are not exported from their package namespace!

Add the class at the end of the function name to get the code for exported R (S3) methods:

```
residuals.lm
## function (object, type = c("working", "response", "deviance",
##
       "pearson", "partial"), ...)
## {
##
       type <- match.arg(type)
      r <- object$residuals
       res <- switch(type, working = , response = r, deviance = ,
           pearson = if (is.null(object$weights)) r else r * sqrt(object$weights),
           partial = r)
      res <- naresid(object$na.action, res)
       if (type == "partial")
           res <- res + predict(object, type = "terms")
       res
## <bytecode: 0x7febdf95b5e8>
## <environment: namespace:stats>
```

Note: this requires to know the class of the object you work with! You can use class() on your input to figure this out.

It is also possible to get the code of non-exported R methods (S3):

```
residuals.nls
## Error in eval(expr. envir. enclos): object 'residuals.nls' not found
getAnvwhere("residuals.nls") # or getS3method("residuals", "nls")
## A single object matching 'residuals.nls' was found
## It was found in the following places
    registered S3 method for residuals from namespace stats
     namespace:stats
## with value
##
## function (object, type = c("response", "pearson"), ...)
## {
##
       type <- match.arg(type)
       if (type == "pearson") {
           val <- as.vector(object$m$resid())</pre>
           std <- sqrt(sum(val^2)/(length(val) - length(coef(object))))
           val <- val/std
           if (!is.null(object$na.action))
               val <- naresid(object$na.action, val)
##
           attr(val, "label") <- "Standardized residuals"
##
       else {
           val <- as.vector(object$m$lhs() - object$m$fitted())</pre>
##
           if (!is.null(object$na.action))
               val <- naresid(object$na.action, val)
           lab <- "Residuals"
           if (!is.null(aux <- attr(object, "units")$y))
               lab <- paste(lab, aux)
           attr(val, "label") <- lab
##
```

Challenge

Which function actually computes the numbers behind boxplot()?

Challenge

What is the code behind t.test()?

Some functions — the interfaces — call functions that are written in other languages. The source code of these latter functions is not directly visible (spotted as .C(), .Fortran(), .Primitive(), .Internal(), .External()).

```
dnorm
## function (x, mean = 0, sd = 1, log = FALSE)
## .Call(C_dnorm, x, mean, sd, log)
## <br/>
## 
## 
function (x, mean = 0, sd = 1, log = FALSE)
## control (x, mean = 0, sd = 1, log = FALSE)
## control (x, mean = 0, sd = 1, log = FALSE)
## control (x, mean = 0, sd = 1, log = FALSE)
## control (x, mean = 0, sd = 1, log = FALSE)
## control (x, mean = 0, sd = 1, log = FALSE)
## control (x, mean = 0, sd = 1, log = FALSE)
## control (x, mean = 0, sd = 1, log = FALSE)
## control (x, mean = 0, sd = 1, log = FALSE)
## control (x, mean = 0, sd = 1, log = FALSE)
## control (x, mean = 0, sd = 1, log = FALSE)
## control (x, mean = 0, sd = 1, log = FALSE)
## control (x, mean = 0, sd = 1, log = FALSE)
## control (x, mean = 0, sd = 1, log = FALSE)
## control (x, mean = 0, sd = 1, log = FALSE)
## control (x, mean = 0, sd = 1, log = FALSE)
## control (x, mean = 0, sd = 1, log = FALSE)
## control (x, mean = 0, sd = 1, log = FALSE)
## control (x, mean = 0, sd = 1, log = FALSE)
## control (x, mean = 0, sd = 1, log = FALSE)
## control (x, mean = 0, sd = 1, log = FALSE)
## control (x, mean = 0, sd = 1, log = FALSE)
## control (x, mean = 0, sd = 1, log = FALSE)
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## control (x, mean = 0, sd = 1, log = FALSE)
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## control (x, mean = 0, sd = 1, log = FALSE)
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## control (x, mean = 0, sd = 1, log = FALSE)
## control (x, mean = 0, sd = 1, log = FALSE)
## control (x, mean = 0, sd = 1, log = FALSE)
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## control (x, mean = 0, sd = 1, log = FALSE)
## control (x, mean = 0, sd = 1, log = FALSE)
## control (x, mean = 0, sd = 1, log = FALSE)
## control (x, mean = 0, sd = 1, log = FALSE)
## control (x, mean = 0, sd = 1, log = FALSE)
## control (x, mean = 0, sd = 1, log = FALSE)
## control (x, mean = 0, sd = 1, log = 1, lo
```

In these cases, the easiest is to use the read-only mirror for R (https://github.com/wch/r-source) or the relevant package on Github! (here, the answer lies in r-source/src/nmath/dnorm.c)

For more info, check:

https://stackoverflow.com/questions/19226816/how-can-i-view-the-source-code-for-a-function.

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Challenge

What is the code really estimating coefficients behind lm()?

Programming with **R**

- A few coding tips
- **5** Example: comparing the performances of 2 tests
- Debugging

Debugging a faulty function

```
pythagora <- function(x, y) {
    x2 <- x^2
    y2 <- y^2
    hyp <- (x^2 + y^2)^1/2
    return(hyp)
}

pythagora(x = 2, y = 2)
## [1] 4</pre>
```

Debugging a faulty function

```
pythagora <- function(x, y) {
    x2 <- x^2
    y2 <- y^2
    hyp <- (x^2 + y^2)^1/2
    return(hyp)
}

pythagora(x = 2, y = 2)
## [1] 4</pre>
```

```
pythagora <- function(x, y) {
    x2 <- x^2
    y2 <- y^2
    browser()
    hyp <- (x^2 + y^2)^1/2
    return(hyp)
}

pythagora(x = 2, y = 2)</pre>
```

Note: this is very useful but you need to have access to the code!

Debugging a faulty function

```
pythagora <- function(x, y) {
    x2 <- x^2
    y2 <- y^2
    hyp <- (x^2 + y^2)^1/2
    return(hyp)
}

pythagora(x = 2, y = 2)
## [1] 4</pre>
```

```
debug(pythagora)

pythagora(x = 2, y = 2)
undebug(pythagora) ## when you are done, or use debugonce() above, or reload the function
```

Note: this can work on any function without having to mess with the code!

Debugging a faulty function

There are plenty more debugging possibilities out there!

Check:

- the nice possibilities with RStudio: https://support.rstudio.com/hc/en-us/articles/205612627-Debugging-with-RStudio
- ?option and look at error
- ?traceback
- ?trace

Programming with ${\bf R}$

- Writing simple functions
- Why programming?
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- 4 A few coding tips
- **5** Example: comparing the performances of 2 tests
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Remember

"premature optimization is the root of all evil"

Donald Knuth, Computing Surveys, Vol 6, No 4, December 1974

When to optimize?

- optimize only when the code really works
- optimize only if needed

Profiling

You can time functions:

```
library(spaMM)

data("Loaloa")

system.time({
   fullfit <- fitme(maxNDVI ~ 1 + Matern(1|latitude+longitude), data = Loaloa)
})

## user system elapsed
## 11.765 1.816 14.959</pre>
```

Profiling

You can time functions:

```
library(spaMM)

data("Loaloa")

system.time({
   fullfit <- fitme(maxNDVI ~ 1 + Matern(1|latitude+longitude), data = Loaloa)
})

## user system elapsed
## 11.765 1.816 14.959</pre>
```

You can profile the code very precisely using profvis:

```
library(profvis)
profvis({
  fullfit <- fitme(maxNDVI ~ 1 + Matern(1|latitude+longitude), data = Loaloa)
})</pre>
```

Note: try it out!

General tips to improve speed

• limit to number of function calls as much as possible!

```
system.time({
  replicate(le6, runif(1))
  })

## user system elapsed
## 3.664 0.316 4.014
```

```
system.time({
  runif(le6)
  })
## user system elapsed
## 0.027 0.001 0.028
```

Note: it is always best to design functions that can work on vectors and not just on single values.

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General tips to improve speed

- limit to number of function calls as much as possible!
- compare alternative implementations

```
d <- data.frame(proba = runif(500), group = factor(1:500))

system.time({
    all_pairwise(proba = d$proba, groups = d$group)
    })

## user system elapsed
## 7.840 0.446 8.344</pre>
```

```
system.time({
    all_pairwise2(proba = d$proba, groups = d$group)
    })

## user system elapsed
## 7.962 0.368 8.357
```

General tips to improve speed

- limit to number of function calls as much as possible!
- compare alternative implementations
- code bottlenecks in another language and rely on interfaces (advanced)

Programming with ${\bf R}$

- Writing simple functions
- Why programming
- Coding basics
- A few coding tips
- 5 Example: comparing the performances of 2 tests
- Exploring existing code
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What is an ${\bf R}$ package

An R package is only a way to organise one's data, scripts and documentation!

Benefits:

- easy distribution
- thorough check of the code
- incentive to write up some documentation and examples

Note: it is good practice to write a research project as a package!

How to create an R package

- \bullet you can simply use the function package.skeleton() (see help for example)
- you can use the combo RStudio + the R package devtools

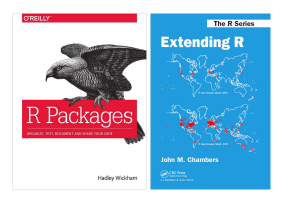
How to create an R package

- you can simply use the function package.skeleton() (see help for example)
- you can use the combo RStudio + the R package devtools

For more info:

https://cran.r-project.org/doc/manuals/r-release/R-exts.html

Or:



Programming with \boldsymbol{R}

- Writing simple functions
- Why programming
- Coding basics
- A few coding tips
- 5 Example: comparing the performances of 2 tests
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- Debugging
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Crude:

```
set.seed(1)
d <- data.frame(foo = runif(10), bar = runif(10))

my_wilcox <- function(var1, var2, data) {
    w <- wilcox.test(x = data[, var1], y = data[, var2]) ## data$var1 would not work!
    return(w)
}

my_wilcox(var1 = "foo", var2 = "bar", data = d)

##
## Wilcoxon rank sum test
##
## data: data[, var1] and data[, var2]
## W = 47, p-value = 0.8534
## alternative hypothesis: true location shift is not equal to 0</pre>
```

Better:

```
set.seed(1)
d <- data.frame(foo = runif(10), bar = runif(10))

my_wilcox <- function(var1, var2, data) {
    w <- eval(substitute(wilcox.test(x = data$var1, y = data$var2)))
    return(w)
}

my_wilcox(var1 = "foo", var2 = "bar", data = d)

##

## Wilcoxon rank sum test

##

## data: d$foo and d$bar

## W = 47, p-value = 0.8534

## alternative hypothesis: true location shift is not equal to 0</pre>
```

quote & substitute generate unevaluated pieces of code. eval can be used to evaluate the piece of code.

```
saved_code <- quote(a + b)
saved_code
## a + b
a <- 2
b <- 3
eval(saved code)
## [1] 5
eval_example <- function(){</pre>
  a <- 5
  b <- 5
  print(eval(saved_code))
  print(eval(saved code, envir = .GlobalEnv))
eval example()
## [1] 10
## [1] 5
```

Note: the difference between quote & substitute is that substitute allows for substitution before the evaluation (useful to play with variable names).

A few examples:

```
foo <- 1
getNameFromObject <- function(i) paste(substitute(i))
getNameFromObject(foo)
## [1] "foo"
getValueFromName <- function(i) get(i)
getValueFromName("foo")
## [1] 1
setObjectFromName <- function(i, x) assign(i, x, envir = globalenv())
setObjectFromName("bar", 2)
bar
## [1] 2</pre>
```

Note: there are plenty more possibilities for working on the language in R, but it would require to dive quite deeper into how R works.

Recursions

The Fibonacci example:

```
fibonacci <- function(x) {
   if (x == 0) return(0)
   if (x == 1) return(1)
   return(fibonacci(x - 1) + fibonacci(x - 2))
}

system.time(foo <- fibonacci(35))

## user system elapsed
## 11.792  0.124  12.000

foo

## [1] 9227465</pre>
```

Note: if you plan to rename functions, you can use "Recall()"!

June 2018

Using C++ code within functions

The package Rcpp allows to run C++ code inside R on the fly!

```
library(Rcpp)
fibRcpp <- cppFunction('
int fibonacci(const int x) {
    if (x == 0) return(0);
    if (x == 1) return(1);
    return (fibonacci(x - 1) + fibonacci(x - 2));
}
')
system.time(foo <- fibRcpp(35)) ## beware: memory issue if x is too large!
## user system elapsed
## 0.066 0.000 0.065
foo
## [1] 9227465</pre>
```

Programming with ${\bf R}$

- Writing simple functions
- Why programming
- Coding basics
- 4 A few coding tips
- 5 Example: comparing the performances of 2 tests
- 6 Exploring existing code
- Debugging
- Optimisation & Profiling
- Writing R packages
- Writing more advanced functions
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Definitions

What is an object?

An instance of a class.

What is a class?

A data structure for which specific methods can be defined.

What is a method?

A function that is designed to work with all objects of a given class.

R is an object-based system

Rule #1:

Everything that exists in \boldsymbol{R} is an object

(John M. Chambers)

Example

res <- 1

```
res
## [1] 1
class(res)
## [1] "numeric"
methods(class = "numeric")
## [1] [
           Γ<-
                                 all.equal Arith
## [5] as.data.frame as.Date
                                 as.POSIXct as.POSIXlt
## [9] as.raster cbind2
                                          Compare
                                 coerce
## [13] confint Logic
                                            rbind2
                                 Ops
## see '?methods' for accessing help and source code
`<-`
## .Primitive("<-")
class(`<-`)
## [1] "function"
methods(class = "function")
## [1] as.data.frame as.list
                                coef
                                             coerce
## [5] coerce<-
                   head
                                plot
                                             print
## [9] tail
## see '?methods' for accessing help and source code
```

Objects are accessed or modified using references

Reference = a name + an environment:

```
ls()
   [1] "a"
                            "all_pairwise"
   [3] "all_pairwise2"
                             "b"
## [5] "bar"
## [7] "do_slow"
                            "e1"
                            "f"
    [9] "eval example"
## [11] "fibonacci"
                            "fibRcpp"
## [13] "foo"
                            "fullfit"
## [15] "getNameFromObject" "getValueFromName"
## [17] "group1"
                             "group2"
## [19] "i"
                            "i"
## [21] "Loaloa"
                             "my_function"
## [23] "mv wilcox"
                            "new.u"
## [25] "odds ratio"
                            "process factor"
## [27] "pythagora"
                            "res"
## [29] "saved code"
                            "setObjectFromName"
## [31] "u"
environmentName(pryr::where("res"))
## [1] "R_GlobalEnv"
```

Note: remember, we discussed environments earlier on!

Objects are accessed or modified using references

Behind one reference there is one memory address:

```
pryr::address(res)
## [1] "0x7febe51ce548"
```

Behind one memory address there can be several references:

```
res2 <- res
pryr::address(res2)
## [1] "0x7febe51ce548"
```

Note: as we will see, the address behind a reference can change during computation.

There are different systems for defining and using objects in R

Native class systems:

- S3 (legacy from S version 3, in base)
- S4 (legacy from S version 4, in core package methods)
- Reference Class (sometimes referred to as R5, in methods)

There are different systems for defining and using objects in R

Native class systems:

- S3 (legacy from S version 3, in base)
- S4 (legacy from S version 4, in core package methods)
- Reference Class (sometimes referred to as R5, in methods)

Additional class systems:

- R6 (in the package R6, one of the most downloaded package on CRAN!)
- ggproto (in the package ggplot2)
- others (proto, ...)

Note: the objects created with one system can contain objects created with another.

The 2 main object-based programming paradigms

Functional (Object Oriented) Programming

- suitable for analytical workflows
- S3, S4
- methods defined outside the objects
- objects are not mutable
- a \xrightarrow{fn} b \xrightarrow{fn} c \xrightarrow{fn} ...

The 2 main object-based programming paradigms

Functional (Object Oriented) Programming

- suitable for analytical workflows
- S3, S4
- methods defined outside the objects
- objects are not mutable
- $\bullet \ a \xrightarrow{fn} b \xrightarrow{fn} c \xrightarrow{fn} \dots$

Encapsulated Object Oriented Programming (aka OOP)

- suitable for data that evolve over time (modularity and re-usability)
- RC, R6
- methods defined inside the objects
- objects are mutable
- $\bullet \ a \xrightarrow{fn} b \ \& \ a \xrightarrow{fn} \varnothing$

Note: actual programming can borrow from multiple paradigms (pure form is difficult).

A simple example using S3

```
addone <- function(x) res + 1
res <- 1
addone(res)
## [1] 2</pre>
```

Alexandre Courtiol (IZW) Getting to program in R June 2018 81/99

A simple example using S4

```
setClass(Class = "S4_obj", slots = list(value = "numeric"))
setGeneric(name = "addone", def = function(object) stop("only for S4_obj")) ## set generic with default behaviour
## [1] "addone"
setMethod(f = "addone", signature = "S4_obj", definition = function(object) {
    object@value <- object@value + 1
    return(object)
    }
    )

resS4 <- new("S4_obj", value = 1)
addone(resS4)
## An object of class "S4_obj"
## Slot "value":
## 511 2</pre>
```

A simple example using RC

A simple example using R6

```
library(R6)
R6_obj <- R6Class(
  public = list(
   value = NA,
   initialize = function(value) self$value <- value.
    addone = function() self$value <- self$value + 1
resR6 <- R6_obj$new(value = 1)
resR6$addone()
resR6$value
## [1] 2
resR6$value <- 1
replicate(10, resR6$addone())
## [1] 2 3 4 5 6 7 8 9 10 11
resR6$value
## [1] 11
```

Encapsulated OOP requires mutability

S3 objects are (generally) not mutable (same for S4):

```
a <- 1:3
a
## [1] 1 2 3
pryr::address(a)
## [1] "0x7febc98a46a0"
```

```
a[2] <- 10
a
## [1] 1 10 3
pryr::address(a)
## [1] "0x7febcb26ac98"
```

```
b <- a
b
## [1] 1 10 3
pryr::address(b)
## [1] "0x7febcb26ac98"
b[2] <- 11
pryr::address(b)
## [1] "0x7febct5031b8"
pryr::address(a)
## [1] "0x7febcb26ac98"
a
## [1] 1 10 3</pre>
```

Encapsulated OOP requires mutability

R6 objects are mutable (same for RC):

```
resR6$value

## [1] 11

pryr::address(resR6)

## [1] "0x7febc7a02c40"
```

```
resR6$value <- 3
resR6$value
## [1] 3
pryr::address(resR6)
## [1] "0x7febc7a02c40"
```

```
resR6_bis <- resR6
pryr::address(resR6_bis)
## [1] "0x7febc7a02c40"

resR6_bis$value
## [1] 3
resR6_bis$value <- 4
resR6_bis$value
## [1] 4

resR6$value
## [1] 4
```

86 / 99

Note: mutability can be prevented if necessary.

Example: An simple Individual Based Models (IBM)

Context:

- two age classes: children (0–14 yrs), adults (15+ yrs)
- children die at an average rate of 15 deaths per 1000 per year
- adults die at an average rate of 5 deaths per 1000 per year
- children do not reproduce
- adults reproduce at an average rate of 10 births per 1000 per year

Question:

Starting with 1000 individuals (with age following a uniform distribution between 0 and 40 yrs), what is the number of children and adults after 50 years?

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Functional way using S3: functions

```
death <- function(pop) {
    death_children <- rbinom(n = length(pop), size = 1, prob = 15/1000)
    death_adults <- rbinom(n = length(pop)), size = 1, prob = 5/1000)
    alive <- rep(1, length(pop))
    alive[pop < 15] <- 1 - death_children[pop < 15]
    alive[pop > 14] <- 1 - death_adults[pop > 14]
    pop <- pop[alive == 1]
    return(pop)
}
birth <- function(pop) {
    adults <- pop[pop > 14]
    babies_nb <- sum(rbinom(n = length(adults), size = 1, prob = 10/1000))
    babies <- rep(0, babies_nb)
    pop <- c(pop, babies)
    return(pop)
}
age <- function(pop) pop <- pop + 1</pre>
```

Functional way using S3: run

```
pop <- round(runif(1000, min = 0, max = 40))

for (i in 1:50) {
    pop <- birth(pop)
    pop <- death(pop)
    pop <- age(pop)
    }

table(pop > 14)

##

## FALSE TRUE

## 111 1032
```

OOP way using R6: definition of the class individual

```
individual <- R6Class(
   public = list(
        age = NA,
        alive = 1,
        initialize = function(age = 0) {self$age <- age},
        die = function() {
        if (self$age < 15 & self$alive) self$alive <- 1 - rbinom(n = 1, size = 1, prob = 15/1000)
        if (self$age > 14 & self$alive) self$alive <- 1 - rbinom(n = 1, size = 1, prob = 5/1000)
    },
    reproduce = function() {
        ifelse(self$alive == 1 & self$age > 14, rbinom(n = 1, size = 1, prob = 10/1000), FALSE)
    },
    aging = function() {
        if (self$alive == 1) {self$age <- self$age + 1; self$die()}
    }
}</pre>
```

OOP way using R6: test

You can test things before creating the population!

```
alex <- individual$new()
                                                                        for (i in 1:200) alex$aging()
alex
                                                                        alex
## <R6>
                                                                        ## <R6>
     Public:
                                                                             Public:
       age: 0
                                                                               age: 59
                                                                        ##
       aging: function ()
                                                                               aging: function ()
       alive: 1
                                                                        ##
                                                                               alive: 0
       clone: function (deep = FALSE)
                                                                               clone: function (deep = FALSE)
       die: function ()
                                                                               die: function ()
##
                                                                        ##
##
       initialize: function (age = 0)
                                                                        ##
                                                                               initialize: function (age = 0)
       reproduce: function ()
                                                                        ##
                                                                               reproduce: function ()
```

OOP way using R6: definition of the class population

```
population <- R6Class(
  public = list(
    individuals = list(),
    initialize = function(N = 1000) {
        for (i in 1:N)
          self$individuals[[i]] <- individual$new(age = round(runif(n = 1, min = 0, max = 40)))
    repro = function() {
      for (i in 1:length(self$individuals))
        if (self$individuals[[i]]$reproduce() == TRUE)
          self$individuals[[length(self$individuals) + 1]] <- individual$new()
    death = function(){
      alive <- sapply(self$individuals, function(i) i[["alive"]])</pre>
      self$individuals[!alive] <- NULL
    aging = function(){
      for (i in 1:length(self$individuals)) self$individuals[[i]]$aging()
     },
    year = function() {self$repro(); self$death(); self$aging()},
    count = function() table(sapply(self$individuals, function(i) i[["age"]]) > 14)
```

OOP way using R6: run

```
pop <- population$new()
for (i in 1:50) pop$year()
pop$count()
##
## FALSE TRUE
## 130 980</pre>
```

Note: I did not handle the possible population crash, so it may crash :-/

Our Individual Based Models (IBM): UPDATE

Setup:

- two age classes: children (0–14 yrs), adults (15+ yrs)
- children die at an average rate of 15 deaths per 1000 per year
- adults die at an average rate of 5 deaths per 1000 per year
- children do not reproduce
- adults reproduce at an average rate of 10 births per 1000 per year
- two sexes, females do not reproduce after 45 yrs (males do not reproduce)

Question:

Starting with 1000 individuals (with age following a uniform distribution between 0 and 40 yrs), what is the number of children and adults after 50 years?

OOP way using R6: re-definition of the class individual

```
individual <- R6Class(
 public = list(
   age = NA.
   alive = 1.
   sex = NA,
   initialize = function(age = 0) {
      self$age <- age
     self$sex <- ifelse(runif(1) < 0.5, "male", "female")</pre>
                                                                            ##n.e.u
   die = function() {
     if (self$age < 15 & self$alive) self$alive <- 1 - rbinom(n = 1, size = 1, prob = 15/1000)
     if (self$age > 14 & self$alive) self$alive <- 1 - rbinom(n = 1, size = 1, prob = 5/1000)
   reproduce = function() {
      ifelse(self$alive == 1 &
               (self$age > 14 & self$age < 44 & self$sex == "female"),
                                                                            ##n.ew
             rbinom(n = 1, size = 1, prob = 10/1000), FALSE)
   aging = function() {
      if (self$alive == 1) {self$age <- self$age + 1; self$die()}
```

Note: there is no need to redefine the class population!

OOP way using R6: re-run

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Functional way using S3: re-defining functions

We would have to recode everything ...

Pros and cons of R6 for IBM

Pros

- clearer structure (see butterfly example)
- easier to modify once existing
- easier to share classes between projects, packages...
- easier to translate to C++

Cons

- initially difficult for those knowing mostly S3
- much slower (cost can somewhat be reduced with some tweaks)

98 / 99

• additional issues to take care (e.g. side effects)

A more advanced simulation model using R6

?Butterfly

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