(A bit of) Advanced R

Efficient R programming

Julien Chiquet

Université Paris Dauphine

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http://github/jchiquet/CourseAdvancedR





Resources

- Gillespie & Lovelace (2016): efficient R programming
- Wickham (2014)

Part 0: Prerequesties

xpply family, do.call, Reduce

Outline

- 1 Benchmark your code
- 2 Use all your cores when needed
- 3 Good and bad practices in R
- 4 Part 4: Remember that R is object oriented
- 5 Interface with lower-level languages

Quick (and dirty) benchmarking with system.time()

One usually relies on the command system.time(expr) to evaluate the timings:

```
func.one <- function(n) {return(rnorm(n,0,1))}
func.two <- function(n) {return(rpois(n,1))}

n <- 1000
system.time(replicate(100, func.one(n)))</pre>
```

```
## user system elapsed
## 0.012 0.000 0.010

system.time(replicate(100, func.two(n)))
```

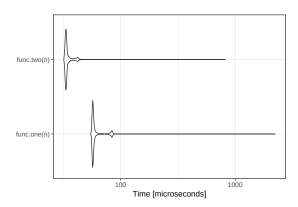
```
## user system elapsed
## 0.008 0.000 0.007
```

Quick benchmarking with microbenchmark

```
func.one <- function(n) {return(rnorm(n,0,1))}
func.two <- function(n) {return(rpois(n,1))}

library(microbenchmark)

n <- 1000
res <- microbenchmark(func.one(n), func.two(n), times=1000)
ggplot2::autoplot(res)</pre>
```



Profile your code

Suppose you want to evaluate which part of the following function is hot:

```
## generate data, center/scale and perform ridge regression
my func <- function(n,p) {
 require (MASS)
  ## draw data
  x <- matrix(rnorm(n*p),n,p)
  y <- rnorm(n)
  ## center/scale
  xs \leftarrow scale(x)
  vs \leftarrow v - mean(v)
  ## return ridge's coefficients
  ridge <- lm.ridge(ys~xs+0,lambda=1)
  return(ridge$coef)
```

Profile your code with base Rprof I

One can rely on the default Rprof function, with somewhat technical outputs

```
Rprof(file="profiling.out", interval=0.05)
res <- my_func(1000,500)
Rprof(NULL)</pre>
```

```
summaryRprof("profiling.out")$by.self
```

```
self.time self.pct total.time total.pct
##
## "La.svd"
                   1.05
                          77.78
                                    1.10
                                            81.48
## "matrix"
                   0.15 11.11
                                    0.15 11.11
## "aperm.default"
                 0.05 3.70
                                    0.05 3.70
## "apply"
                   0.05 3.70
                                    0.05
                                            3.70
## "[.data.frame"
                0.05
                           3.70
                                    0.05
                                            3.70
```

```
summaryRprof("profiling.out")$by.total
```

Profile your code with base Rprof II

##		total.time	total nct	self time	self nct
	"block exec"	1.35	100.00	0.00	0.00
##	"call block"	1.35	100.00	0.00	
##	"eval"	1.35		0.00	
##	"evaluate"	1.35		0.00	
##	"evaluate_call"	1.35	100.00	0.00	0.00
##	"evaluate::evaluate"	1.35	100.00	0.00	0.00
##	"FUN"	1.35		0.00	0.00
##	"handle"	1.35	100.00	0.00	0.00
##	"in_dir"	1.35	100.00	0.00	0.00
##	"knit"	1.35	100.00	0.00	0.00
##	"knitr::knit"	1.35	100.00	0.00	0.00
##	"lapply"	1.35	100.00	0.00	0.00
##	"my_func"	1.35	100.00	0.00	0.00
##	"process_file"	1.35	100.00	0.00	0.00
##	"process_group"	1.35	100.00	0.00	0.00
##	"process_group.block"	1.35	100.00	0.00	0.00
##	"rmarkdown::render"	1.35	100.00	0.00	0.00
##	"timing_fn"	1.35	100.00	0.00	0.00
##	"withCallingHandlers"	1.35	100.00	0.00	0.00
##	"withVisible"	1.35	100.00	0.00	0.00
##	"lm.ridge"	1.15	85.19	0.00	0.00
##	"La.svd"	1.10	81.48	1.05	77.78
##	"svd"	1.10	81.48	0.00	0.00
##	"matrix"	0.15	11.11	0.15	11.11
##	"scale"	0.10	7.41	0.00	0.00
##	"scale.default"	0.10	7.41	0.00	0.00

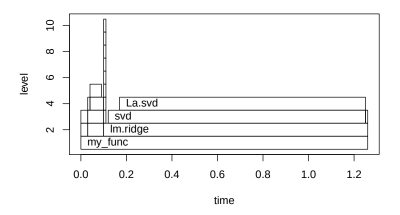
Profile your code with base Rprof III

```
"aperm.default"
                                0.05
                                           3.70
                                                     0.05
                                                               3.70
  "apply"
                                0.05
                                           3.70
                                                     0.05
                                                               3.70
## "[.data.frame"
                                0.05
                                          3.70
                                                     0.05
                                                              3.70
##
                                0.05
                                          3.70
                                                     0.00
                                                              0.00
  "aperm"
                                0.05
                                          3.70
                                                     0.00
                                                               0.00
  "eval.parent"
                                0.05
                                           3.70
                                                     0.00
                                                              0.00
## ".External2"
                                0.05
                                          3.70
                                                     0.00
                                                              0.00
## "model.frame.default"
                                0.05
                                          3.70
                                                     0.00
                                                              0.00
  "na.omit"
                                0.05
                                          3.70
                                                     0.00
                                                              0.00
## "na.omit.data.frame"
                                0.05
                                          3.70
                                                     0.00
                                                              0.00
## "stats::model.frame"
                                0.05
                                           3.70
                                                     0.00
                                                              0.00
## "sweep"
                                0.05
                                           3.70
                                                     0.00
                                                               0.00
```

Profile your code with profr

The *profr* package is maybe a little easier to understand...

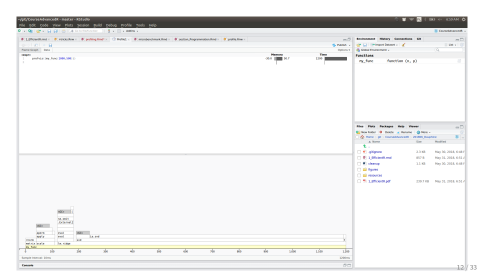
```
library(profr)
profiling <- profr({my_func(1000,500)}, interval = 0.01)
plot(profiling)</pre>
```



Profile your code within R Studiow with profvis

Profvis integrates the profiling to the Rstudio API

```
library(profvis)
profvis({my_func(1000,500)})
```



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Parallel computing

Usual Roadmap

- lacktriangledown Start up and intialize M 'worker' processes
- Send data required for each task to the workers
- $\textbf{ § Split the task into } M \text{ roughly equally-sized chunks and send them (including the R code needed) to the workers$
- 4 Wait for all the workers to complete their tasks, and ask them for their results
- **5** Repeat steps (2–4) for any further tasks
- 6 Shut down the worker processes

Socketing vs Forking

Two approaches achieving the same goal

The socket approach

- launches a new version of R on each core
- connection is done via networking all happening on your own computer

The forking approach

- copies the entire current version of R and moves it to a new core
- several processes acheive the same task resulting in different outputs
- → Forking is only possible on Unix systems (Linux, Mac OS)

Parallel computing with parallel

Package parallel

- merge of packages multicore and snow
- included in base R and maintained by the R Core team

Check your computer

```
library(parallel) ## embedded with R since version 2.9 or something
cores <- detectCores() ## How many cores do I have?
print(cores)</pre>
```

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

→ parallel features both socketing (parLapply) and forking (mclapply)

Forking approach with parallel::mclapply

Very easy: use parallel features as soon as you do simulations!

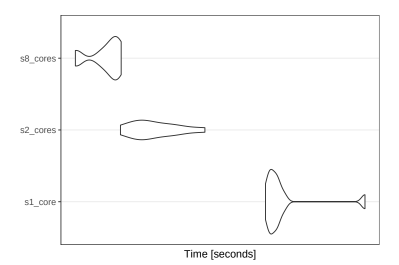
Example: estimates the test error from ridge regression

```
one.simu <- function(i) {
    ## draw data
    n <- 1000; p <- 500
    x <- matrix(rnorm(n*p),n,p) ; y <- rnorm(n)
    ## return ridge's coefficients
    train <- 1:floor(n/2)
    test <- setdiff(1:n,train)
    ridge <- MASS::lm.ridge(y~x+0,lambda=1,subset=train)
    err <- (y[test] - x[test, ] %*% ridge$coef )^2
    return(list(err = mean(err), sd = sd(err)))
}</pre>
```

```
head(do.call(rbind, mclapply(1:8, one.simu, mc.cores = cores)), n = 3)
```

```
## err sd
## [1,] 14.45796 19.79301
## [2,] 10.3222 14.86089
## [3,] 12.15729 16.8608
```

Forking approach with parallel::mclapply (cont'd)



Socket approach with parallel::parLapply

Windows users need a bit more code to make it work

A possible option: export from base workspace

```
cl <- makeCluster(4)
clusterExport(cl,"one.simu")
res <- parSapply(cl, 1:8, one.simu) # several parLapply call are possible
stopCluster(cl)
res</pre>
```

```
## [,1] [,2] [,3] [,4] [,5] [,6] [,7]

## err 11.86539 11.26563 8.845916 14.37104 12.91292 13.25386 12.50689

## sd 16.6455 15.63073 12.32548 18.74685 18.49328 17.62274 16.61861

## [,8]

## err 8.508118

## sd 12.48525
```

Parallel computing with parallel: final remarks

- Parallelize piece of code complex enough
- Do not choose stupidly the number of cores
- Screen outputs are lost in Rstudio: use pbmcapply (progress bar)

```
pbmcapply::pbmcmapply(1:8, FUN = one.simu, mc.cores = 2)
```

```
## [,1] [,2] [,3] [,4] [,5] [,6] [,7] [,8]
## err 10.72632 14.25714 9.77985 9.589137 10.90891 11.8773 10.73228 11.2432
## sd 13.99144 18.25551 13.08587 14.26683 15.5244 15.03568 16.37883 15.9956
```

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Vectorize any algebraic operation I

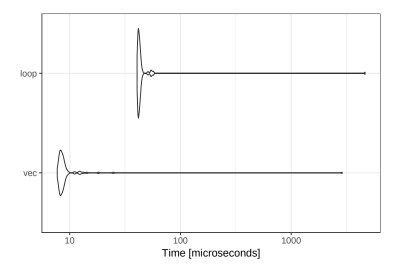
```
Example: compute \exp(x) = \sum_{k=0}^{n} \frac{x^k}{k!}
```

```
## the good way
exp_vec <- function(x, n) {
    res <- sum(x^(0:n)/c(1,cumprod(1:n)))
    res
}

## the sad/bad/less readable way
exp_loop <- function(x, n) {
    res <- 1
    for (k in 1:n) res <- res + 2^k/factorial(k)
    res
}</pre>
```

```
autoplot(microbenchmark(vec = exp_vec(2, 100), loop = exp_loop(2, 100)))
```

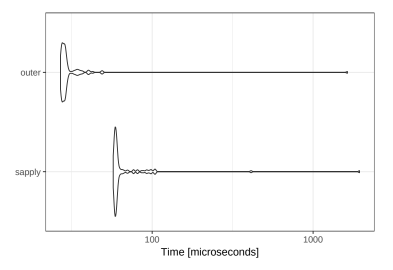
Vectorize any algebraic operation II



Vectorize, even for non-algebraic operation I

```
month_year_apply <- function(year) {</pre>
  sapply(month.name, function(month) paste(month, year, sep = " "))
month year outer <- function(year) {
  outer(month.name, year, FUN = paste, sep = ' ')
head(month year outer(c(2010, 2013)), 3)
##
  [,1]
                        [,2]
## [1,] "January_2010" "January_2013"
## [2,] "February_2010" "February_2013"
## [3,] "March 2010" "March 2013"
autoplot(microbenchmark(
  sapply = month year apply(c(2011, 2013)),
  outer = month_year_outer(c(2011, 2013)),
  times = 100)
```

Vectorize, even for non-algebraic operation II

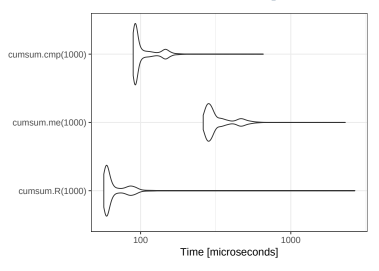


Compile your functions with base::compiler |

If you cannot avoid a loop, you will save some time

```
cumsum.R <- function(n) {</pre>
  x \leftarrow rnorm(n)
  cumsum(x)
cumsum.me <- function(n) {</pre>
  x \leftarrow rnorm(n)
  res <- 0
  for (i in 1:length(x))
    res <- res + x[i]
  res
cumsum.cmp <- compiler::cmpfun(cumsum.me)</pre>
autoplot(
  microbenchmark(
    cumsum.R(1000).
    cumsum.me(1000).
    cumsum.cmp(1000),
    times=1000)
```

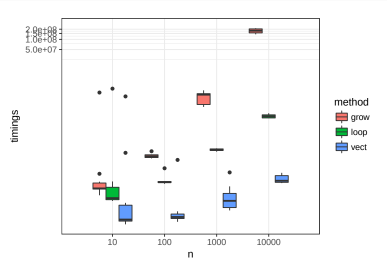
Compile your functions with base::compiler II



→ Can be set automotically with compiler::enableJIT(3)

Preallocate whenever it is possible

```
grow <- function(n) {vec <- numeric(0); for (i in 1:n) vec <- c(vec,i)}
loop <- function(n) {vec <- numeric(n); for (i in 1:n) vec[i] <- i}
vect <- function(n) {1:n}</pre>
```



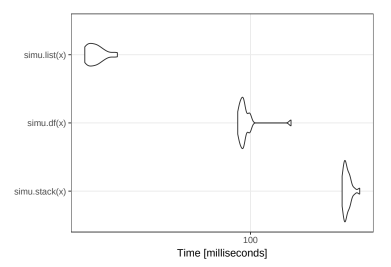
Do not stack objects I

Even if it is tempting when the final size is unknown.

```
simu.stack <- function(x) { ## x is a n x p matrix
  out <- data.frame(mean = numeric(0), sd = numeric(0))
 for (i in 1:n)
    out <- rbind(out, data.frame(mean = mean(x[i,]), sd = sd(x[i,])))
  return(out)
simu.df <- function(x) {
  out <- data.frame(mean = numeric(n), sd = numeric(n))</pre>
  for (i in 1:n)
    out[i, ] \leftarrow c(mean = mean(x[i,]), sd = sd(x[i,]))
 return(out)
simu.list <- function(x) {
  my.list \leftarrow lapply(1:n, function(i) c(mean(x[i,]), sd(x[i,])))
  out <- data.frame(do.call(rbind, mv.list))
  colnames(out) <- c("mean"."sd")</pre>
  return(out)
```

```
n <- 1000; p <- 10; x <- matrix(rnorm(n*p), n, p)
autoplot(microbenchmark(simu.stack(x), simu.df(x), simu.list(x), times=20))</pre>
```

Do not stack objects II



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References

Gillespie, C., & Lovelace, R. (2016). *Efficient r programming*. "O'Reilly Media, Inc." Retrieved from https://bookdown.org/csgillespie/efficientR/

Wickham, H. (2014). *Advanced r.* CRC Press. Retrieved from http://adv-r.had.co.nz/