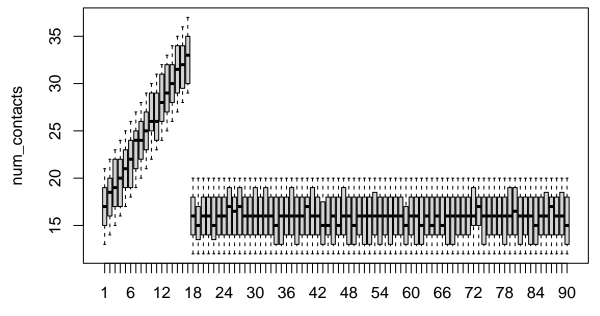
## simid\_workshop\_tutorial\_bootstrap.R

## lwillem

## 2023-02-23

```
# SIMID TUTORIAL: BOOTSTRAP & PARALLEL PROGRAMMING
# This program is free software: you can redistribute it and/or modify
# it under the terms of the GNU General Public License as published by
# the Free Software Foundation, either version 3 of the License, or
# (at your option) any later version.
# Copyright (C) 2023 lwillem, SIMID, UNIVERSITY OF ANTWERP, BELGIUM
# clear workspace
rm(list=ls())
# # set working directory (or open RStudio with this script)
\# setwd("C:\User\path\to\the\rcode\folder") \# WINDOWS
# setwd("/Users/path/to/the/rcode/folder") ## MAC
# SETUP ----
print('SETUP PARAMETERS: START')
## [1] "SETUP PARAMETERS: START"
# set random number generator
rng seed <- 20190524
set.seed(rng_seed)
# e.g. population details
pop_size <- 1e4
           <- 90
max_age
adult_age
           <- 18
gender_male <- 'M'</pre>
gender_female <- 'F'</pre>
         <- c(gender_male, gender_female)</pre>
gender_opt
zipcode_opt <- c(1000,2160,2640,2018,2110,3000,3500,3520,3590,9000)
community_opt <- 1:30</pre>
# e.q. social contact details
mean_num_cnt <- 16
                 # population average
dev_num_cnt <- -4:4
                     # deviance
```

```
# e.g. create population data
pop_data <- data.frame(id</pre>
                                  = 1:pop_size,
                                  = sample(1:max_age,pop_size,replace = T),
                                  = sample(gender_opt,pop_size,replace = T),
                                  = sample(zipcode_opt,pop_size,replace = T),
                        community = sample(community_opt,pop_size,replace = T))
# e.g. sample number of contacts, based on population average
pop_data$num_contacts <- mean_num_cnt + sample(dev_num_cnt,pop_size,replace=T)</pre>
# e.g. children make more contacts, relative to their age
is_child
                                 <- pop_data$age<adult_age</pre>
pop_data$num_contacts[is_child] <- pop_data$num_contacts[is_child] +</pre>
 pop_data$age[is_child]
# explore "number of contacts"
boxplot(num_contacts ~ age, data = pop_data)
```



```
print('SETUP PARAMETERS: COMPLETE')
```

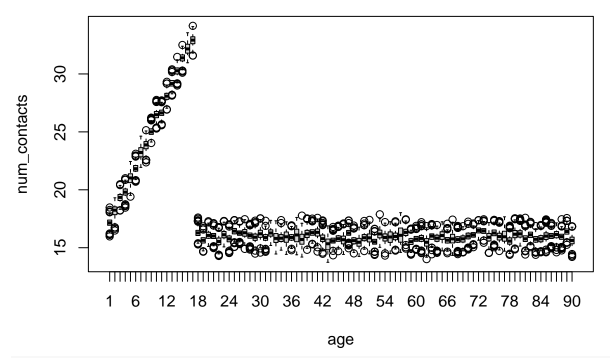
age

## [1] "SETUP PARAMETERS: COMPLETE"

```
# BOOTSTRAP: FOR-LOOP ----
print('BOOTSTRAPS: START')
## [1] "BOOTSTRAPS: START"
num bootstraps <- 500
# for-loop 1
for(i_bootstrap in 1:num_bootstraps){
  # set RGN seed per bootstrap, so the processing sequence does not affect the results
 set.seed(rng_seed + i_bootstrap)
  # bootstrap population
                    <- pop_data[sample(pop_size,replace=T),]</pre>
 pop_bootstrap
  # get mean age by community and gender
  summary_num_contacts <- aggregate(num_contacts ~ age + gender, data = pop_bootstrap, mean)</pre>
  # aggregate results: data.frame
  out_data_frame <- data.frame(i_bootstrap,</pre>
                                   summary_num_contacts)
  # # plot results
  # pdf(file=pasteO('output/bootstrap',i_bootstrap,'.pdf'))
  # plot(out_data_frame$num_contacts)
  # dev.off()
  # # save results to txt file
  # write.table(out_data_frame, file=paste0('output/bootstrap',i_bootstrap,'.txt'),row.names=F,sep=',')
  # # save results to RData file
   \# \ save(out\_data\_frame, file=paste0('output/bootstrap', i\_bootstrap,'.RData')) \\
}
# retrieve data etc
# ....
print('BOOTSTRAPS: COMPLETE')
```

## [1] "BOOTSTRAPS: COMPLETE"

```
# SEQUENTIAL FOREACH ----
print('SEQUENTIAL FOREACH: START')
## [1] "SEQUENTIAL FOREACH: START"
library('foreach')
print(system.time( # start manual profiling
# for-loop 2: return values
bootstrap_out <- foreach(i_bootstrap = 1:num_bootstraps,</pre>
                      .combine = 'rbind') %do%{
 # set RGN seed per bootstrap, so the processing sequence does not affect the results
 set.seed(rng_seed + i_bootstrap)
 # bootstrap population
 pop_bootstrap
                 <- pop_data[sample(pop_size,replace=T),]</pre>
 # get mean age by community and gender
 summary_num_contacts <- aggregate(num_contacts ~ age + gender, data = pop_bootstrap, mean)</pre>
 # aggregate results: data.frame
 out_data_frame <- data.frame(i_bootstrap,</pre>
                                  summary_num_contacts)
 # return results
return(out_data_frame)
)) # end system.time()
     user system elapsed
   7.379 1.282 8.866
##
# plot bootstrap results
boxplot(num_contacts ~ age, data=bootstrap_out)
```



print('BOOTSTRAPS: COMPLETE')

## [1] "BOOTSTRAPS: COMPLETE"

```
# PARALLEL FOREACH ----
print('PARALLEL FOREACH')
## [1] "PARALLEL FOREACH"
# load package 'devtools'
library(devtools)
## Loading required package: usethis
# install (and load) package "simid_rtools" from https://github.com/lwillem/simid_rtools
devtools::install_github("lwillem/simid_rtools",force=F,quiet=T)
# start parallel working nodes
# note: make sure you loaded all required user defined functions at this point
simid.rtools::smd_start_cluster()
print(system.time( # manual profiling
# run loop in parallel
bootstrap_out <- foreach(i_bootstrap = 1:num_bootstraps,</pre>
                        .packages = NULL, # make sure you add the required packages
                        .combine = 'rbind') %dopar%{
   # set RGN seed per bootstrap, so the processing sequence does not affect the results
   set.seed(rng_seed + i_bootstrap)
   # bootstrap population
                      <- pop_data[sample(pop_size,replace=T),]</pre>
  pop_bootstrap
   # get mean age by community and gender
   summary_num_contacts <- aggregate(num_contacts ~ age + gender, data = pop_bootstrap, mean)</pre>
   # aggregate results: data.frame
                    <- data.frame(i_bootstrap,</pre>
  out data frame
                                    summary_num_contacts)
   # return results
  return(out_data_frame)
)) # end system.time()
##
     user system elapsed
    0.309
           0.090
                  2.480
# terminate parallel cluster
simid.rtools::smd_stop_cluster()
print('PARALLEL FOREACH: COMPLETE')
```

## [1] "PARALLEL FOREACH: COMPLETE"