Simple 3-state Partitioned Survival model in R

The DARTH workgroup

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- Jalal H, Pechlivanoglou P, Krijkamp E, Alarid-Escudero F, Enns E, Hunink MG. An Overview of R in Health Decision Sciences. Med Decis Making. 2017; 37(3): 735-746. https://journals.sagepub.com/doi/abs/10.1177/0272989X16686559
- Krijkamp EM, Alarid-Escudero F, Enns EA, Jalal HJ, Hunink MGM, Pechlivanoglou P. Microsimulation modeling for health decision sciences using R: A tutorial. Med Decis Making. 2018;38(3):400–22. https://journals.sagepub.com/doi/abs/10.1177/0272989X18754513
- Krijkamp EM, Alarid-Escudero F, Enns E, Pechlivanoglou P, Hunink MM, Jalal H. A Multidimensional Array Representation of State-Transition Model Dynamics. BioRxiv 670612 2019.https://www.biorxiv.org/content/10.1101/670612v1

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Change eval to TRUE if you want to knit this document.

```
rm(list = ls())  # clear memory (removes all the variables from the workspace)
```

01 Load packages

```
if (!require('pacman')) install.packages('pacman'); library(pacman) # use this package to
# load (install if required) packages from CRAN
p_load("here", "dplyr", "devtools", "gems", "flexsurv", "survminer", "survHE", "ggplot2", "msm", "igraph
```

02 Load functions

```
source("survival_functions.R")
```

03 Input model parameters

```
v_n
         <- c("healthy", "sick", "dead") # state names
         <- length(v_n)
                                         # No of states
n_s
         <- 50000
                                         # number of simulations
n_i
         <- 1 / 12
                                         # cycle length (a month)
c_1
         <- 30
                                         # number of years (20 years)
n_t
         <- seq(0, n_t, c_1)
                                         # the cycles in years
times
set.seed(2020)
                                         # set the seed
```

Create a transition probability matrix with all transitions indicated and numbered.

Generate data.

```
source("data.R")
head(true_data)
head(sim_data)
head(status)
head(OS_PFS_data)
```

04 Analysis

Showcasing the use of packages survival, flexsurv.

```
fit_KM <- survfit(Surv(time = OS_time, event = OS_status) ~ 1, data = OS_PFS_data,
                      type ="fleming-harrington")
plot(fit_KM, mark.time = T)
# a prettier way of plotting
ggsurvplot(
 fit_KM,
 data = OS_PFS_data,
 size = 1,
                            # change line size
 palette = c("orange2"),  # custom color palettes
 conf.int = TRUE,
                           # Add confidence interval
 pval = TRUE,
                           # Add p-value
                          # Add risk table
 risk.table = TRUE,
 risk.table.height = 0.25, # Useful to change when you have multiple groups
 ggtheme = theme_bw(),
                          # Change ggplot2 theme
                        # Change X-axis label
 xlab = 'Time in days',
 title = "Survival curve for Progression-Free Survival (PFS)",
 subtitle = "Based on Kaplan-Meier estimates"
```

04.1 Partitioned Survival model

04.2 MultiState modeling method 1

• Fit all parametric multistate models simultaneously.

```
# The existing functions in R require the data in a long rather than a wide format
# convert the data in a way that flexsurv understands using the mstate package
                <- msprep(time = sim_data, status = status, trans = tmat )</pre>
data long
data_long$trans <- as.factor(data_long$trans) # convert trans to a factor</pre>
data_long$from <- case_when(data_long$from == 1 ~ "healthy",</pre>
                              data_long$from == 2 ~ "sick",
                              data_long$from == 3 ~ "dead")
                <- case_when(data_long$to == 1 ~ "healthy",
data_long$to
                              data_long$to == 2 ~ "sick",
                              data_long$to == 3 ~ "dead")
# fit all parametric multistate models simultaneously to the data and extract the AIC/BIC
# Select the one with the lowest AIC
fits <- fit.mstate(time ="time", status = "status", trans, data = data_long,
                   times = times, extrapolate = T )
best.fit <- fits[["Loglogistic"]]</pre>
```

• Construct a DES model out of the simultaneously fitted multistate model.

```
source(here("functions", "survival_functions_backend.R"))
# Construct a DES model out of the simultaneously fitted multistate model
DES_data <- sim.fmsm(best.fit, start = 1, t = n_years, trans = tmat, M = n_i)
m_M_DES <- trace.DES(DES_data, n_i = n_i, times = times, tmat = tmat)
m_M_DES1 <- trace.DES1(DES_data, n_i = n_i, times = times, tmat = tmat)
all(m_M_DES == m_M_DES1)
system.time(trace.DES(DES_data, n_i = n_i, times = times, tmat = tmat))
system.time(trace.DES1(DES_data, n_i = n_i, times = times, tmat = tmat))</pre>
```

04.3 MultiState modeling method 2

• Multistate models can be fitted independently for each transition.

```
best.fit_HD <- fit_HD[["Weibull"]]
best.fit_SD <- fit_SD[["Lognormal"]]</pre>
```

- A microsimulation can be fitted instead of a DES.
- more computationally expensive but it provides more freedom to the modeller.
- For the Microsimulation to be run, we need transition probabilities per unit of time.

```
# Extract transition probabilities from the best fitting models
p_HS <- flexsurvreg_prob(object = best.fit_HS, times = times)</pre>
p_HD <- flexsurvreg_prob(object = best.fit_HD, times = times)</pre>
p_SD <- flexsurvreg_prob(object = best.fit_SD, times = times)</pre>
# everyone starts in the "healthy" state and therefore has not spent time in "sick"
v M init <- rep("healthy", times = n i)</pre>
v_Ts_init <- rep(0, n_i) # a vector with the time of being sick at the start of the model
# function that generates the transition probabilities per cycle
Probs <- function(M_t, v_Ts, t) {</pre>
  # Arguments:
    # M_t: health state occupied by at cycle t (character variable)
    # v_Ts: vector with the duration of being sick
    # t:
            current cycle
  # Returns:
    # transition probabilities for that cycle
  # create matrix of state transition probabilities
                 <- matrix(0, nrow = n_s, ncol = n_i)
  # give the state names to the rows
  rownames(m_p_t) <- v_n
  # update m_p_t with the appropriate probabilities
  # transition probabilities when healthy
  m_pt[, M_t == "healthy"] <- rbind(1 - p_HD[t] - p_HS[t], p_HS[t], p_HD[t])
  # transition probabilities when sick
  m_p_t[, M_t == "sick"] \leftarrow rbind(0, 1 - p_SD[v_Ts], p_SD[v_Ts])
  # transition probabilities when dead
  m_p_t[, M_t == "dead"] <- rbind(0, 0, 1)
  return(t(m_p_t))
```

04.3.1 Run Microsimulation

```
m_M <- matrix(nrow = n_i, ncol = length(times) ,</pre>
                  dimnames = list(paste("ind" , 1:n_i, sep = " "),
                                   paste("year", times, sep = " ")))
                                 # initial health state for individual i
  m_M[, 1] <- v_M_init
  v_Ts
           <- v_Ts_init
                                  # initialize time since illness onset for individual i
  # open a loop for time running cycles 1 to n t
  for (t in 1:(length(times)-1)) {
    # calculate the transition probabilities for the cycle based on health state t
    m_p <- Probs(m_M[, t], v_Ts, t)</pre>
    \# sample the current health state and store that state in matrix m\_M
    m_M[, t + 1] <- samplev(m_p, 1)</pre>
    # update time since illness onset for t + 1
    v_Ts \leftarrow ifelse(m_M[, t + 1] == "sick", v_Ts + 1, 0)
    # Display simulation progress
    if(t %in% seq(1,(length(times)),10)) { # display progress every 10%
      cat('\r', paste(round(t/length(times)*100,0), "% done", sep = " "))
    } else if (t == (length(times)-1)) {cat('\r', paste("100% done"))}
  } # close the loop for the time points
  # store the results from the simulation in a list
  results <- list(m_M = m_M)
 return(results) # return the results
} # end of the MicroSim function
# Run the simulation model
Micro_data <- MicroSim(n_i, seed = 1)</pre>
# create the microsimulation trace
m_M_Micro <- t(apply(Micro_data$m_M, 2, function(x) table(factor(x, levels = v_n,</pre>
                                                                   ordered = TRUE))))
m M Micro <- m M Micro / n i
                                 # calculate the proportion of individuals
colnames(m_M_Micro) <- v_n</pre>
rownames(m_M_Micro) <- paste("Cycle", times, sep = " ")</pre>
```

05 Compare all methods