

Simple 3-state Partitioned Survival model in R

The DARTH workgroup

Developed by the Decision Analysis in R for Technologies in Health (DARTH) workgroup:

Fernando Alarid-Escudero, PhD (1)

Eva A. Enns, MS, PhD (2)

M.G. Myriam Hunink, MD, PhD (3,4)

Hawre J. Jalal, MD, PhD (5)

Eline M. Krijkamp, MSc (3)

Petros Pechlivanoglou, PhD (6)

Alan Yang, MSc (7)

In collaboration of:

1. Drug Policy Program, Center for Research and Teaching in Economics (CIDE) - CONACyT, Aguascalientes, Mexico
2. University of Minnesota School of Public Health, Minneapolis, MN, USA
3. Erasmus MC, Rotterdam, The Netherlands
4. Harvard T.H. Chan School of Public Health, Boston, USA
5. University of Pittsburgh Graduate School of Public Health, Pittsburgh, PA, USA
6. The Hospital for Sick Children, Toronto and University of Toronto, Toronto ON, Canada
7. The Hospital for Sick Children, Toronto ON, Canada

Please cite our publications when using this code:

- 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>

Copyright 2017, THE HOSPITAL FOR SICK CHILDREN AND THE COLLABORATING INSTITUTIONS. All rights reserved in Canada, the United States and worldwide. Copyright, trademarks, trade names and any and all associated intellectual property are exclusively owned by THE HOSPITAL FOR SICK CHILDREN and the collaborating institutions. These materials may be used, reproduced, modified, distributed and adapted with proper attribution.

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 conveniently
# 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
n_s      <- length(v_n)                  # No of states
n_i      <- 50000                         # number of simulations
c_l      <- 1 / 12                        # cycle length (a month)
n_t      <- 30                           # number of years (20 years)
times    <- seq(0, n_t, c_l)             # the cycles in years
set.seed(2020)                           # set the seed
```

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

```
tmat <- matrix(NA, n_s, n_s, dimnames = list(v_n, v_n))
tmat["healthy", "sick"] <- 1
tmat["healthy", "dead"] <- 2
tmat["sick", "dead"] <- 3

layout.fig <- c(2,1)
plotmat(t(tmat), t(layout.fig), self.cex = 0.5, curve = 0, arr.pos = 0.76,
        latex = T, arr.type = "curved", relsize = 0.85, box.prop=0.8,
        cex = 0.1, box.cex = 0.7, lwd = 1)
```

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
  risk.table = TRUE,            # Add risk table
  risk.table.height = 0.25,      # Useful to change when you have multiple groups
  ggtheme = theme_bw(),         # Change ggplot2 theme
  xlab = 'Time in days',        # Change X-axis label
  title = "Survival curve for Progression-Free Survival (PFS)",
  subtitle = "Based on Kaplan-Meier estimates"
)
```

04.1 Partitioned Survival model

```
# Flexsurv allows parametric fitting of curves
fit_weib <- flexsurvreg(Surv(time = OS_time, event = OS_status) ~ 1, data = OS_PFS_data,
                       dist = "weibull")
plot(fit_weib)

# fit all parametric models to the data and extract the AIC/BIC.
# Select the one with the most appropriate fit
# Repeat for PFS and OS
fit_PFS <- fit.fun(time = "PFS_time", status = "PFS_status", data = OS_PFS_data,
                  times = times, extrapolate = T)
fit_OS <- fit.fun(time = "OS_time", status = "OS_status", data = OS_PFS_data,
                 times = times, extrapolate = T)

best_PFS <- fit_PFS[["Weibull"]]
best_OS <- fit_OS[["Weibull"]]

# construct a partitioned survival model out of the fitted models
m_M_PSM <- partsurv(best_PFS, best_OS, time = times)$trace
```

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
data_long      <- msprep(time = sim_data, status = status, trans = tmat )
data_long$trans <- as.factor(data_long$trans) # convert trans to a factor

data_long$from  <- case_when(data_long$from == 1 ~ "healthy",
                             data_long$from == 2 ~ "sick",
                             data_long$from == 3 ~ "dead")

data_long$to    <- case_when(data_long$to == 1 ~ "healthy",
                             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"]]

```

- 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))

```

04.3 MultiState modeling method 2

- Multistate models can be fitted independently for each transition.

```

# Multistate models can be fitted independently for each transition. This is more flexible!
# Create subsets for each transition
data_HS <- subset(data_long, trans == 1)
data_HD <- subset(data_long, trans == 2)
data_SD <- subset(data_long, trans == 3)

# fit independent models for each transition and pick the one with the lowest AIC
fit_HS <- fit.fun(time = "time", status = "status", data = data_HS, times = times,
                 extrapolate = T)
fit_HD <- fit.fun(time = "time", status = "status", data = data_HD, times = times,
                 extrapolate = T)
fit_SD <- fit.fun(time = "time", status = "status", data = data_SD, times = times,
                 extrapolate = T)

best.fit_HS <- fit_HS[["Gamma"]]

```

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

- 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)
p_HD <- flexsurvreg_prob(object = best.fit_HD, times = times)
p_SD <- flexsurvreg_prob(object = best.fit_SD, times = times)

# everyone starts in the "healthy" state and therefore has not spent time in "sick"
v_M_init <- rep("healthy", times = n_i)
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) {
  # 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
  m_p_t <- 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_p_t[, 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"] <- 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

```
MicroSim <- function(n_i, seed = 1) {
  # Arguments:
  # n_i: number of individuals
  # seed: default is 1

  set.seed(seed) # set the seed

  # m_M is used to store the health state information over time for every individual
  times <- seq(0, n_t, c_l) # the cycles in years
```

```

m_M <- matrix(nrow = n_i, ncol = length(times) ,
              dimnames = list(paste("ind" , 1:n_i, sep = " "),
                              paste("year", times, sep = " ")))

m_M[, 1] <- v_M_init          # initial health state for individual i
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)
  # sample the current health state and store that state in matrix m_M
  m_M[, t + 1] <- samplev(m_p, 1)

  # update time since illness onset for t + 1
  v_Ts <- 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)

# create the microsimulation trace
m_M_Micro <- t(apply(Micro_data$m_M, 2, function(x) table(factor(x, levels = v_n,
                                                                ordered = TRUE)))))

m_M_Micro <- m_M_Micro / n_i      # calculate the proportion of individuals
colnames(m_M_Micro) <- v_n
rownames(m_M_Micro) <- paste("Cycle", times, sep = " ")

```

05 Compare all methods

```

# Calculate trace for the real data
m_M_data <- transitionProbabilities(generate$cohort, times = times)@probabilities

# visually compare all methods
matplot(times,m_M_data, type='l', lty = 1, col = 1, ylab= "proportion of cohort", xlab = "Time",
        main = "Trace comparisons",xlim=c(0,25))
matlines(times, m_M_DES, col = 2, lty = 1)

```

```
matlines(times, m_M_Micro, col = 3, lty = 1)
matlines(times, m_M_PSM, col = 4, lty = 1)
legend("right", c("True Data", "DES", "Microsim", "PSM"),
      col = 1:4, lty = rep(1,4), bty= "n")
```