## Markov Sick-Sicker model in R

with dependency for time-since model start AND with state-residency dependency

#### 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. Med Decis Making. 2020 Online first. https://doi.org/10.1177/0272989X19893973

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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 conveniently
# load (install if required) packages from CRAN
p_load("here", "dplyr", "devtools", "scales", "ellipse", "ggplot2", "lazyeval", "igraph", "truncnorm",
# load (install if required) packages from GitHub
# install_github("DARTH-git/dampack", force = TRUE) Uncomment if there is a newer version
# install_github("DARTH-git/darthtools", force = TRUE) Uncomment if there is a newer version
p_load_gh("DARTH-git/dampack", "DARTH-git/darthtools")
```

#### 02 Load functions

```
# No functions needed
```

### 03 Input model parameters

```
# Strategy names
v_names_str <- c("No Treatment", "Treatment")</pre>
# Number of strategies
n_str <- length(v_names_str)</pre>
# Markov model parameters
age <- 25
                                     # age at baseline
max_age <- 55
                                    # maximum age of follow up
n_t <- max_age - age
                                   # time horizon, number of cycles
       <- c("H", "S1", "S2", "D") # the 4 states of the model: Healthy (H), Sick (S1),
                                    # Sicker (S2), Dead (D)
n_states <- length(v_n)</pre>
                                    # number of health states
# Tunnels
n_tunnel_size <- n_t</pre>
# Sick state
v_Sick_tunnels <- paste("S1_", seq(1, n_tunnel_size), "Yr", sep = "")</pre>
### Create variables for time-dependent model
                 <- c("H", v_Sick_tunnels, "S2", "D") # state names
n_states_tunnels <- length(v_n_tunnels)</pre>
                                                         # number of states
# Transition probabilities (per cycle) and hazard ratios
# Read age-specific mortality rates from csv file
lt_usa_2005 <- read.csv("HMD_USA_Mx_2015.csv")</pre>
```

```
v_r_HD <- lt_usa_2005 %>%
         filter(Age >= age & Age <= (max_age-1)) %>%
          select(Total) %>%
         as.matrix()
p_HD <- 1 - exp(- v_r_HD) # probability to die when healthy
p_HS1 <- 0.15
                                     # probability to become sick when healthy
p_S1H <- 0.5
                                      # probability to become healthy when sick
# Weibull parameters
1 <- 0.08 # scale
      <- 1.1 # shape
# Weibull function
p_S1S2 <- l*g*(1:n_tunnel_size)^{g-1} # probability to become sicker when sick
                                      # (time-dependent)
hr_S1 <- 3
                                      # hazard ratio of death in sick vs healthy
hr_S2 <- 10
                                      # hazard ratio of death in sicker vs healthy
r_{HD} \leftarrow -\log(1 - p_{HD})
                                  # rate of death in healthy
r_S1D \leftarrow hr_S1 * r_HD
                                     # rate of death in sick
r_S2D <- hr_S2 * r_HD
                                     # rate of death in sicker
                                # probability to die in sick
p_S1D \leftarrow 1 - exp(-r_S1D)
p_S2D \leftarrow 1 - exp(-r_S2D)
                                   # probability to die in sicker
# Cost and utility inputs
c_H <- 2000
                                   # cost of remaining one cycle in the healthy state
c S1 <- 4000
                                   # cost of remaining one cycle in the sick state
c S2 <- 15000
                                   # cost of remaining one cycle in the sicker state
                                  # cost of treatment(per cycle)
c_trt <- 12000
c_D <- 0
                                  # cost of being in the death state
      <- 1
                                  # utility when healthy
\mathtt{u}_{-}\mathtt{H}
u_S1 <- 0.75
                                   # utility when sick
                                 # utility when sicker
u_S2 <- 0.5
u_D <- 0
                                   # utility when dead
u_trt <- 0.95
                                   # utility when being treated
# Discounting factor
       <- 0.03
                                    # equal discount of costs and QALYs by 3%
\# calculate discount weights for costs for each cycle based on discount rate d_c
v_dwc \leftarrow 1 / (1 + d_r) \hat{ } (0:n_t)
\# calculate discount weights for effectiveness for each cycle based on discount rate d_{\_}e
v_dwe < 1 / (1 + d_r) ^ (0:n_t)
```

## 04 Define and initialize matrices and vectors

#### 04.1 Cohort trace

```
dimnames = list(paste("cycle", 0:n_t, sep = " "), v_n_tunnels))
head(m_M_notrt) # show first 6 rows of the matrix

# The cohort starts as healthy
# initialize first cycle of Markov trace accounting for the tunnels
m_M_notrt[1, ] <- m_M_trt[1, ] <- c(1, rep(0, n_tunnel_size), 0, 0)</pre>
```

#### 04.2 Transition probability array

```
# create the transition probability array for NO treatment
a_P_notrt <- array(0,  # Create 3-D array
dim = c(n_states_tunnels, n_states_tunnels, n_t),
dimnames = list(v_n_tunnels, v_n_tunnels, 0:(n_t-1))) # name dimensions</pre>
```

Fill in the transition probability array:

```
# from Healthy
a_P_notrt["H", "H", ]
                                    \leftarrow (1 - p_HD) * (1 - p_HS1)
a_P_notrt["H", v_Sick_tunnels[1], ] <- (1 - p_HD) * p_HS1
a_P_notrt["H", "D",]
                                    <- p_HD
# from Sick
for(i in 1:(n_tunnel_size - 1)){
 a_P_notrt[v_Sick_tunnels[i], "H", ] <- (1 - p_S1D) * p_S1H
  a P notrt[v Sick tunnels[i], v Sick tunnels[i + 1], ] <-
    (1 - p_S1D) * (1 - (p_S1H + p_S1S2[i]))
  a_P_notrt[v_Sick_tunnels[i], "S2", ] \leftarrow (1 - p_S1D) * p_S1S2[i]
  a_P_notrt[v_Sick_tunnels[i], "D", ] <- p_S1D
a_P_notrt[v_Sick_tunnels[n_tunnel_size], "H", ] <- (1 - p_S1D) * p_S1H
a_P_notrt[v_Sick_tunnels[n_tunnel_size], v_Sick_tunnels[n_tunnel_size], ] <-
  (1 - p_S1D) * (1 - (p_S1H + p_S1S2[n_tunnel_size]))
a_P_notrt[v_Sick_tunnels[n_tunnel_size], "S2", ] <- (1 - p_S1D) * p_S1S2[n_tunnel_size]
a_P_notrt[v_Sick_tunnels[n_tunnel_size], "D", ] <- p_S1D</pre>
# from Sicker
a_P_notrt["S2", "S2", ] <- 1 - p_S2D
a_P_notrt["S2", "D", ] <- p_S2D
# from Dead
a_P_notrt["D", "D", ] <- 1</pre>
# Check that transition probabilities are in [0, 1]
check_transition_probability(a_P_notrt, verbose = TRUE)
# Check that all rows sum to 1
check_sum_of_transition_array(a_P_notrt, n_states = n_states_tunnels, n_t = n_t, verbose = TRUE)
# create transition probability matrix for treatment same as NO treatment
a_P_trt <- a_P_notrt</pre>
```

### 05 Run Markov model

```
for (t in 1:n_t){ # loop through the number of cycles
  m_M_notrt[t + 1, ] <- t(m_M_notrt[t, ]) %*% a_P_notrt[ , , t] # estimate the Markov
                                                                  # trace for cycle the
                                                                  # next cycle (t + 1)
 m_M_trt[t + 1, ] <- t(m_M_trt[t, ]) %*% a_P_trt[, , t] # estimate the Markov
                                                                  # trace for cycle the
                                                                  # next cycle (t + 1)
} # close the loop
head(m_M_notrt) # show the first 6 lines of the matrix
# create aggregated traces
m_M_td_notrt <- cbind(H = m_M_notrt[, "H"],</pre>
                      S1 = rowSums(m_M_notrt[, 2:(n_tunnel_size +1)]),
                      S2 = m_M_{\text{notrt}}[, "S2"],
                      D = m_M_notrt[, "D"])
head(m M td notrt)
m_M_td_trt <- cbind(H = m_M_trt[, "H"],</pre>
                      S1 = rowSums(m_M_trt[, 2:(n_tunnel_size +1)]),
                      S2 = m_M_trt[, "S2"],
                      D = m_M_{trt}[, "D"])
head(m_M_td_trt)
```

# 06 Compute and Plot Epidemiological Outcomes

#### 06.1 Cohort trace

#### 06.2 Overall Survival (OS)

```
# calculate the overall survival (OS) probability for no treatment
v_os_notrt_tunnels <- 1 - m_M_notrt[, "D"]
# alternative way of calculating the OS probability
v_os_notrt_tunnels <- rowSums(m_M_notrt[, 1:3])
# create a simple plot showing the OS
plot(age:max_age, v_os_notrt_tunnels, type = 'l',</pre>
```

### 06.2.1 Life Expectancy (LE)

### 06.3 Disease prevalence

```
v_prev_tunnels <- rowSums(m_M_td_notrt[, c("S1", "S2")]) / v_os_notrt_tunnels
plot(v_prev_tunnels,
    ylim = c(0, 1),
    ylab = "Prevalence",
    xlab = "Cycle",
    main = "Disease prevalence")</pre>
```

#### 06.4 ratio of sick(S1) vs sicker(S2)

# 07 Compute Cost-Effectiveness Outcomes

### 07.1 Mean Costs and QALYs for Treatment and NO Treatment

### 07.2 Discounted Mean Costs and QALYs

### 07.3 Compute ICERs of the Markov model

#### 07.4 Plot frontier of the Markov model

```
plot(df_cea, effect_units = "QALYs", xlim=c(17,18))
```