SA: Markov Sick-Sicker 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. Med Decis Making. Feb;40(2):242-248. 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("dplyr", "devtools", "scales", "ellipse", "ggplot2", "lazyeval", "igraph", "truncnorm", "ggraph"
# load (install if required) packages from GitHub
# install_github("DARTH-git/dampack", force = TRUE) Uncomment if there is a newer version
p_load_gh("DARTH-git/dampack", "DARTH-git/darthtools")
```

02 Load functions

```
# No function needed
```

03 Input model parameters

 $p_S2D < 1 - exp(-r_S2D)$

```
# Strategy names
v_names_str <- c("No Treatment", "Treatment")</pre>
# Markov model parameters
age
        <- 25
                                     # age at baseline
max_age <- 55
                                     # maximum age of follow up
       <- max_age - age
                                    # time horizon, number of cycles
n_t
       <- c("H", "S1", "S2", "D") # the 4 states of the model: Healthy (H), Sick (S1),
                                     # Sicker (S2), Dead (D)
v init <- c("H" = 1,
            "S1" = 0,
            "S2" = 0,
            "D" = 0)
                                    # initial cohort distribution (everyone allocated to the "healthy"
# Transition probabilities (per cycle)
p_HD
       <- 0.005
                                     # probability to die when healthy
p_HS1
        <- 0.15
                                     # probability to become sick when healthy, conditional on survivin
p_S1H <- 0.5
                                     # probability to become healthy when sick, conditional on survivin
p_S1S2 <- 0.105
                                     # probability to become sicker when sick, conditional on surviving
        <- 3
                                     # hazard ratio of death in sick vs healthy
hr_S1
hr_S2 <- 10
                                     # hazard ratio of death in sicker vs healthy
{	t r}_{	t HD}
       \leftarrow - \log(1 - p_HD)
                                    # rate of death in healthy
r_S1D <- hr_S1 * r_HD
                                     # rate of death in sick
r_S2D \leftarrow hr_S2 * r_HD
                                     # rate of death in sicker
p_S1D \leftarrow 1 - exp(-r_S1D)
                                   # probability to die in sick
```

probability to die in sicker

```
# Cost and utility inputs
       <- 2000
                                      # cost of remaining one cycle in the healthy state
c_H
c S1
        <- 4000
                                      # cost of remaining one cycle in the sick state
       <- 15000
                                     # cost of remaining one cycle in the sicker state
c S2
c_trt <- 12000
                                     # cost of treatment (per cycle)
c D
        <- 0
                                      # cost of being in the death state
                                     # utility when healthy
        <- 1
u_H
u_S1
       <- 0.75
                                     # utility when sick
        <- 0.5
                                      # utility when sicker
u S2
\mathtt{u}_{\mathtt{D}}
        <- 0
                                      # utility when dead
u_trt <- 0.95
                                     # utility when being treated
        <- d_c <- 0.03
                                     # discount rate per cycle equal discount of costs and QALYs by 3%
d_e
        <- length(v_names_str)</pre>
                                      # Number of strategies
{\tt n\_str}
n_states <- length(v_n)</pre>
                                      # number of states
# Discount weights for costs and effects
v_dwc \leftarrow 1 / (1 + d_c) ^ (0:n_t)
v_dwe \leftarrow 1 / (1 + d_e) ^ (0:n_t)
```

Create a state-transition diagram of the cohort model

04 Define and initialize matrices and vectors

04.1 Cohort trace

```
head(m_M_notrt) # show first 6 rows of the matrix

# The cohort starts as healthy
m_M_notrt[1, ] <- m_M_trt[1, ] <- c(1, 0, 0, 0) # initiate first cycle of cohort trace</pre>
```

04.2 Transition probability matrix

Fill in the transition probability matrix:

```
# from Healthy
m_P_notrt["H", "H" ] <- (1 - p_HD) * (1 - p_HS1)
m_P_notrt["H", "S1" ] <- (1 - p_HD) * p_HS1</pre>
m_P_notrt["H", "D" ] <- p_HD</pre>
# from Sick
m_P_notrt["S1", "H"] <- (1 - p_S1D) * p_S1H
m_P_notrt["S1", "S1"] <- (1 - p_S1D) * (1 - (p_S1H + p_S1S2))</pre>
m_P_notrt["S1", "S2"] <- (1 - p_S1D) * p_S1S2</pre>
m_P_notrt["S1", "D" ] <- p_S1D</pre>
# from Sicker
m_P_notrt["S2", "S2"] <- 1 - p_S2D</pre>
m_P_notrt["S2", "D" ] <- p_S2D
# from Dead
m_P_notrt["D", "D" ] <- 1</pre>
# Check that transition probabilities are in [0, 1]
check_transition_probability(m_P_notrt, verbose = TRUE)
# Check that all rows sum to 1
check_sum_of_transition_array(m_P_notrt, n_states = n_states, n_cycles = n_t, verbose = TRUE)
# Overwrite the transition probability matrix for treatment with the no treatment values as they are th
m P trt <- m P notrt
```

05 Run Markov model

```
head(m_M_notrt) # show the first 6 lines of the matrix
```

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 <- 1 - m_M_notrt[, "D"]
# alternative way of calculating the OS probability
v_os_notrt <- rowSums(m_M_notrt[, 1:3])
# create a simple plot showing the OS
plot(0:n_t, v_os_notrt, type = 'l',
    ylim = c(0, 1),
    ylab = "Survival probability",
    xlab = "Cycle",
    main = "Overall Survival")
# add grid
grid(nx = n_t, ny = 10, col = "lightgray", lty = "dotted", lwd = par("lwd"),
    equilogs = TRUE)</pre>
```

06.2.1 Life Expectancy (LE)

```
v_le <- sum(v_os_notrt) # summing probability of OS over time (i.e. life expectancy)</pre>
```

06.3 Disease prevalence

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

06.4 Proportion of sick in S1 state

07 Compute Cost-Effectiveness Outcomes

```
# Vectors with costs and utilities by treatment

v_u_notrt <- c(u_H, u_S1, u_S2, u_D)

v_u_trt <- c(u_H, u_trt, u_S2, u_D)

v_c_notrt <- c(c_H, c_S1, c_S2, c_D)

v_c_trt <- c(c_H, c_S1 + c_trt, c_S2 + c_trt, c_D)
```

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(15.6, 16.6))
```

08 Deterministic Sensitivity Analysis

08.1 List of input parameters

Create list 1_params_all with all input probabilities, cost and utilities.

```
l_params_all <- as.list(data.frame(</pre>
         = 0.005, # probability to die when healthy
  p_HD
  p_HS1
         = 0.15,
                    # probability to become sick when healthy, conditional on surviving
 p_S1H = 0.5, # probability to become healthy when sick, conditional on surviving
  p_S1S2 = 0.105, # probability to become sicker when sick, conditional on surviving
 hr_S1 = 3,  # hazard ratio of death in sick vs healthy
hr_S2 = 10,  # hazard ratio of death in sicker vs healthy
       = 2000, # cost of remaining one cycle in the healthy state
  с Н
  c_S1 = 4000, # cost of remaining one cycle in the sick state
         = 15000, # cost of remaining one cycle in the sicker state
  c S2
  c_trt = 12000, # cost of treatment(per cycle)
         = 0,  # cost of being in the death state
= 1,  # utility when healthy
  c_D
 u_H
  u_S1 = 0.75, # utility when sick
  u_S2 = 0.5, # utility when sicker
  u_D = 0,
                  # utility when dead
 u_trt = 0.95, # utility when treated
 d_e = 0.03, # discount factor for effectiveness
         = 0.03 # discount factor for costs
 d_c
))
# store the parameter names into a vector
v_names_params <- names(l_params_all)</pre>
```

08.2 Load Sick-Sicker Markov model function

```
source("Functions_markov_sick-sicker_sol.R")
# Test function
calculate_ce_out(l_params_all)
```

08.3 One-way sensitivity analysis (OWSA)

```
options(scipen = 999) # disabling scientific notation in R
# dataframe containing all parameters, their basecase values, and the min and
# max values of the parameters of interest
df_params_owsa <- data.frame(pars = c("p_S1S2", "c_trt", "u_S1", "u_trt"),</pre>
                             min = c(0.05, 6000, 0.65, 0.80), # min parameter values
                             \max = c(0.155, 18000, 0.85, 0.98) # max parameter values
                                          = df_params_owsa, # dataframe with parameters for OWSA
owsa nmb <- run owsa det(params range
                                                             # list with all parameters
                          params_basecase = l_params_all,
                          nsamp
                                          = 100,
                                                               # number of parameter values
                          FUN
                                          = calculate_ce_out, # function to compute outputs
                                       = c("NMB"), # output to do the OWSA on
= v_names_str, # names of the strategies
                          outcomes
                          strategies
                                          = 120000)
                                                              # extra argument to pass to FUN
                          n_wtp
```

08.3.1 Plot OWSA

08.3.2 Optimal strategy with OWSA

```
owsa_opt_strat(owsa = owsa_nmb, txtsize = 10)
```

08.3.3 Tornado plot

```
owsa_tornado(owsa = owsa_nmb)
```

08.4 Two-way sensitivity analysis (TWSA)

```
FUN = calculate_ce_out, # function to compute outputs
outcomes = c("NMB"), # output to do the TWSA on
strategies = v_names_str, # names of the strategies
n_wtp = 120000) # extra argument to pass to FUN
```

l ##08.4.1 Plot TWSA

```
plot(twsa_nmb)
```

09 Probabilistic Sensitivity Analysis (PSA)

```
# Function to generate PSA input dataset
gen_psa <- function(n_sim = 1000, seed = 071818){
 set.seed(seed) # set a seed to be able to reproduce the same results
 df_psa <- data.frame(</pre>
    # Transition probabilities (per cycle)
   p_HS1 = rbeta(n_sim, shape1 = 30, shape2 = 170),  # probability to become sick when healthy
   p_S1H = rbeta(n_sim, shape1 = 60, shape2 = 60) , # probability to become healthy when sick
   p_S1S2 = rbeta(n_sim, shape1 = 84, shape2 = 716), # probability to become sicker when sick
   p_HD = rbeta(n_sim, shape1 = 10, shape2 = 1990), # probability to die when healthy
   hr_S1 = rlnorm(n_sim, meanlog = log(3), sdlog = 0.01), # rate ratio of death in S1 vs healthy
   hr S2
          = rlnorm(n_sim, meanlog = log(10), sdlog = 0.02), # rate ratio of death in S2 vs healthy
   # State rewards
   # Costs
   c_H = rgamma(n_sim, shape = 100, scale = 20), # cost of remaining one cycle in state H
   c_S1 = rgamma(n_sim, shape = 177.8, scale = 22.5), # cost of remaining one cycle in state S1
   c S2 = rgamma(n sim, shape = 225, scale = 66.7), # cost of remaining one cycle in state S2
   c Trt = rgamma(n sim, shape = 73.5, scale = 163.3), # cost of treatment (per cycle)
   c_D = 0,
                                                       # cost of being in the death state
   # Utilities
   u_H = rbeta(n_sim, shape1 = 200, shape2 = 3),
                                                       # utility when healthy
   u_S1 = rbeta(n_sim, shape1 = 130, shape2 = 45),
                                                       # utility when sick
   u_S2 = rbeta(n_sim, shape1 = 230, shape2 = 230),
                                                     # utility when sicker
                                                       # utility when dead
   u_Trt = rbeta(n_sim, shape1 = 300, shape2 = 15),
                                                       # utility when being treated
   d_e = 0.03,
                                                       # discount factor for effectiveness
   d_c = 0.03
                                                       # discount factor for costs
   return(df_psa)
}
# Try it
gen_psa(10)
# Number of simulations
n_sim <- 1000
# Generate PSA input dataset
df_psa_input <- gen_psa(n_sim = n_sim)</pre>
```

```
# First six observations
head(df_psa_input)
# Histogram of parameters
ggplot(melt(df_psa_input, variable.name = "Parameter"), aes(x = value)) +
       facet_wrap(~Parameter, scales = "free") +
       geom_histogram(aes(y = ..density..)) +
       scale_x_continuous(breaks = scales::pretty_breaks(n = 3)) +
       theme_bw(base_size = 16) +
       theme(axis.text = element_text(size=6))
# Initialize dataframes with PSA output
# Dataframe of costs
df_c <- as.data.frame(matrix(0,</pre>
                       nrow = n_sim,
                       ncol = n_str))
colnames(df_c) <- v_names_str</pre>
# Dataframe of effectiveness
df_e <- as.data.frame(matrix(0,</pre>
                       nrow = n_sim,
                       ncol = n_str))
colnames(df_e) <- v_names_str</pre>
```

09.1 Conduct probabilistic sensitivity analysis

```
# Run Markov model on each parameter set of PSA input dataset
for(i in 1:n_sim){
    l_out_temp <- calculate_ce_out(df_psa_input[i, ])
    df_c[i, ] <- l_out_temp$Cost
    df_e[i, ] <- l_out_temp$Effect
    # Display simulation progress
    if(i/(n_sim/10) == round(i/(n_sim/10), 0)) { # display progress every 10%
        cat('\r', paste(i/n_sim * 100, "% done", sep = " "))
    }
}</pre>
```

09.2 Create PSA object for dampack

09.2.1 Save PSA objects

09.3 Create probabilistic analysis graphs

```
load(file = "markov_sick-sicker_PSA_dataset.RData")

Vector with willingness-to-pay (WTP) thresholds.

v_wtp <- seq(0, 200000, by = 10000)</pre>
```

09.3.1 Cost-Effectiveness Scatter plot

```
plot(l_psa)
```

09.4 Conduct CEA with probabilistic output

09.4.1 Plot cost-effectiveness frontier

```
plot(df_cea_psa)
```

09.4.2 Cost-effectiveness acceptability curves (CEACs) and frontier (CEAF)

```
ceac_obj <- ceac(wtp = v_wtp, psa = l_psa)
# Regions of highest probability of cost-effectiveness for each strategy
summary(ceac_obj)
# CEAC & CEAF plot
plot(ceac_obj)</pre>
```

09.4.3 Expected Loss Curves (ELCs)

The expected loss is the the quantification of the foregone benefits when choosing a suboptimal strategy given current evidence.

```
elc_obj <- calc_exp_loss(wtp = v_wtp, psa = l_psa)
elc_obj
# ELC plot
plot(elc_obj, log_y = FALSE)</pre>
```

09.4.4 Expected value of perfect information (EVPI)

```
evpi <- calc_evpi(wtp = v_wtp, psa = l_psa)
# EVPI plot
plot(evpi, effect_units = "QALY")</pre>
```