## 3-state Markov model in R

#### with age dependency

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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
- Alarid-Escudero F, Krijkamp EM, Enns EA, Yang A, Hunink MGM Pechlivanoglou P, Jalal H. Cohort State-Transition Models in R: A Tutorial. arXiv:200107824v2. 2020:1-48. http://arxiv.org/abs/2001. 07824
- 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 Mak. 2020;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("diagram", "dampack")
# install_github("DARTH-git/darthtools", force = TRUE) Uncomment if there is a newer version
p_load_gh("DARTH-git/darthtools")
```

## 02 Load functions

```
# No functions needed
```

## 03 Input model parameters

```
## General setup
n_cycles <- 60
                                                # number of cycles
v_names_cycles <- paste("cycle", 0:n_cycles) # cycle names</pre>
v_names_states <- c("Healthy", "Sick", "Dead") # state names</pre>
             <- length(v_names_states)</pre>
                                               # number of health states
n_states
# Discounting factors
d_c <- 0.03
                                                # discount rate for costs
              <- 0.03
                                                # discount rate for QALYs
d_e
# Strategy names
v_names_str <- c("Standard of Care",</pre>
                                               # store the strategy names
                    "Treatment A",
                    "Treatment B")
              <- length(v_names_str)</pre>
                                              # number of strategies
n_str
## Transition probabilities
p_HS_SoC <- 0.05 # probability of becoming sick when healthy, conditional on surviving, under standar
p_HS_trtA <- 0.04 # probability of becoming sick when healthy, conditional on surviving, under treatme
p_HS_trtB <- 0.02 # probability of becoming sick when healthy, conditional on surviving, under treatme
     <- 0.1 # probability of dying
p_HD_min < 0.003 \# probability of dying when healthy at t = 0
p_HD_max <- 0.01 # probability of dying when healthy at t = n_cycles
# probabilities of dying when healthy (age-dependent) - this is now a sequence of numbers
        <- seq(p_HD_min, p_HD_max, length.out = n_cycles)</pre>
## State rewards
```

```
# Costs and utilities
c_H <- 400 # cost of one cycle in healthy state
         <- 1000 # cost of one cycle in sick state
c S
       <- 0 # cost of one cycle in dead state
c D
c_trtA <- 800  # cost of treatment A (per cycle) in healthy state
c_trtB <- 1500 # cost of treatment B (per cycle) in healthy state</pre>
u_H <- 1 # utility when healthy
u_S
       <- 0.5 # utility when sick
u_D
       <- 0 # utility when dead
       <- 0.03 # discount rate per cycle equal discount of costs and QALYs by 3%
d_e
d_c
         <- 0.03 # discount rate per cycle equal discount of costs and QAL</pre>
# Discount weight (equal discounting is assumed for costs and effects)
v_dwc \leftarrow 1 / (1 + d_c) ^ (0:n_cycles)
v_dwe <- 1 / (1 + d_e) ^ (0:n_cycles)</pre>
```

### 04 Define and initialize matrices and vectors

#### 04.1 Cohort trace

#### 04.2 Transition probability array

Fill in the transition probability array:

```
## Standard of Care
# from Healthy
a_P_SoC["Healthy", "Healthy", ] \leftarrow (1 - v_p_HD) * (1 - p_HS_SoC)
a_P_SoC["Healthy", "Sick", ] <- (1 - v_p_HD) * p_HS_SoC
a_P_SoC["Healthy", "Dead", ] <- v_p_HD</pre>
# from Sick
a_P_SoC["Sick", "Sick", ] <- 1 - p_SD
a_P_SoC["Sick", "Dead", ] <- p_SD</pre>
# from Dead
a_P_SoC["Dead", "Dead", ] <- 1</pre>
## Treatment A
a_P_trtA <- a_P_SoC</pre>
a_P_{trtA}["Healthy", "Healthy", ] \leftarrow (1 - v_p_HD) * (1 - p_HS_trtA)
a_P_trtA["Healthy", "Sick", ] <- (1 - v_p_HD) * p_HS_trtA
## Treatment B
a P trtB <- a P SoC
a_P_trtB["Healthy", "Healthy", ] <- (1 - v_p_HD) * (1 - p_HS_trtB)
a_P_trtB["Healthy", "Sick", ] <- (1 - v_p_HD) * p_HS_trtB
```

Check if transition array and probabilities are valid.

```
# Check that transition probabilities are in [0, 1]
check_transition_probability(a_P_SoC, verbose = TRUE)
check_transition_probability(a_P_trtA, verbose = TRUE)
check_transition_probability(a_P_trtB, verbose = TRUE)
# Check that all rows sum to 1
check_sum_of_transition_array(a_P_SoC, n_states = n_states, n_cycles = n_cycles, verbose = TRUE)
check_sum_of_transition_array(a_P_trtA, n_states = n_states, n_cycles = n_cycles, verbose = TRUE)
check_sum_of_transition_array(a_P_trtB, n_states = n_states, n_cycles = n_cycles, verbose = TRUE)
```

### 05 Run Markov model

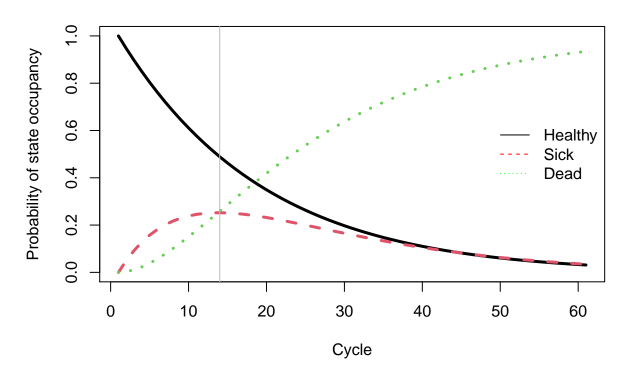
```
for (t in 1:n_cycles){  # loop through the number of cycles
  # estimate the cohort trace for cycle t + 1 using the t-th matrix from the probability array
  m_M_SoC [t + 1, ] <- m_M_SoC [t, ] %*% a_P_SoC [, , t]
  m_M_trtA[t + 1, ] <- m_M_trtA[t, ] %*% a_P_trtA[, , t]
  m_M_trtB[t + 1, ] <- m_M_trtB[t, ] %*% a_P_trtB[, , t]
}
head(m_M_SoC) # print the first few lines of the matrix

## Healthy Sick Dead
## cycle 0 1.0000000 0.00000000 0.00000000
## cycle 1 0.9471500 0.04985000 0.00300000
## cycle 2 0.8969864 0.09207481 0.01093882
## cycle 3 0.8493784 0.12757146 0.02305011
## cycle 4 0.8042016 0.15714071 0.03865771
## cycle 5 0.7613370 0.18149700 0.05716604</pre>
```

# 06 Compute and Plot Epidemiological Outcomes

#### 06.1 Cohort trace

### **Cohort Trace**

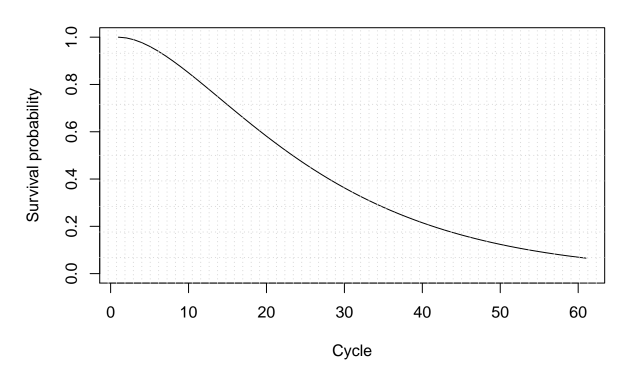


### 06.2 Overall Survival (OS)

```
v_os <- 1 - m_M_SoC[, "Dead"]  # calculate the overall survival (OS) probability
v_os <- rowSums(m_M_SoC[, 1:2])  # alternative way of calculating the OS probability

plot(v_os, type = 'l',
    ylim = c(0, 1),
    ylab = "Survival probability",
    xlab = "Cycle",
    main = "Overall Survival")  # create a simple plot showing the OS</pre>
```

# **Overall Survival**

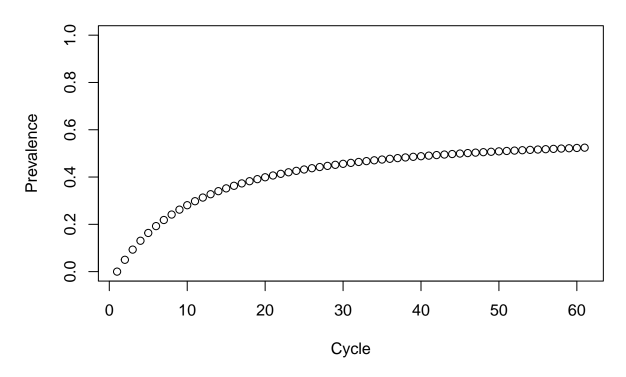


# 06.2.1 Life Expectancy (LE)

```
v_le <- sum(v_os) # summing probability of OS over time (i.e. life expectancy)
```

# 06.3 Disease prevalence

# Disease prevalence



## 07 Compute Cost-Effectiveness Outcomes

#### 07.1 Mean Costs and QALYs

```
# per cycle
# calculate expected costs by multiplying cohort trace with the cost vector for the different health st
v_tc_SoC <- m_M_SoC %*% c(c_H, c_S, c_D)
v_tc_trtA <- m_M_trtA %*% c(c_H + c_trtA, c_S, c_D)
v_tc_trtB <- m_M_trtB %*% c(c_H + c_trtB, c_S, c_D)

# calculate expected QALYs by multiplying cohort trace with the utilities for the different health stat
v_tu_SoC <- m_M_SoC %*% c(u_H, u_S, u_D)
v_tu_trtA <- m_M_trtA %*% c(u_H, u_S, u_D)
v_tu_trtB <- m_M_trtB %*% c(u_H, u_S, u_D)</pre>
```

#### 07.2 Discounted Mean Costs and QALYs

```
# Discount costs by multiplying the cost vector with discount weights (v_dw)
tc_d_SoC <- t(v_tc_SoC) %*% v_dwc
tc_d_trtA <- t(v_tc_trtA) %*% v_dwc
tc_d_trtB <- t(v_tc_trtB) %*% v_dwc

# Discount QALYS by multiplying the QALYs vector with discount weights (v_dw)
tu_d_SoC <- t(v_tu_SoC) %*% v_dwe</pre>
```

```
tu_d_trtA <- t(v_tu_trtA) %*% v_dwe</pre>
tu_d_trtB <- t(v_tu_trtB) %*% v_dwe</pre>
# Store them into a vector
v_tc_d <- c(tc_d_SoC, tc_d_trtA, tc_d_trtB)</pre>
v_tu_d <- c(tu_d_SoC, tu_d_trtA, tu_d_trtB)</pre>
# Dataframe with discounted costs and effectiveness
df_ce <- data.frame(Strategy = v_names_str,</pre>
                     Cost = v_tc_d,
                     Effect = v_tu_d)
df_ce
##
             Strategy
                            Cost
                                   Effect
## 1 Standard of Care 9504.301 14.50646
## 2
          Treatment A 20648.861 15.82191
## 3
          Treatment B 37325.926 19.57368
```

#### 07.3 Compute ICERs of the Markov model

```
Strategy
                         Cost
                                Effect Inc_Cost Inc_Effect
                                                              ICER Status
## 1 Standard of Care 9504.301 14.50646
                                            NA
                                                       NA
                                                                NA
                                                                      ND
         Treatment B 37325.926 19.57368 27821.63
                                                 5.067224 5490.507
                                                                      ND
## 3
         Treatment A 20648.861 15.82191
                                          NA
                                                                      ED
                                                       NA
                                                                NA
```

#### 07.4 Plot frontier of the Markov model

```
plot(df_cea, effect_units = "QALYs")
```

