An alternative representation of state-transition model dynamics - Generic case study using a 3-state model

Appendix of 'Krijkamp EM, Alarid-Escudero F, Enns EA, Hunink MGM, Pechlivanoglou P, Jalal HJ. An alternative representation of state-transition model dynamics'.

2019-02-20

Simple 3-state model

In this simple 3-state model, we simulate a hypothetical cohort of 70-year-old individuals over their remaining lifetime using 50 annual cycles. The cohort start out healthy. The healthy individuals can get sick or they can die. Sick individuals can recover (transitioning back to healthy), stay sick or die. Each health state is associated with utilities and costs. This are the state rewards. In addition to these state rewards, transition rewards apply. There is a sudden decrease of quality of life when just getting sick and the transition to dead is associated with a one-time cost. The state-transition diagram is shown in Figure 1 and all model parameters are described in the table below.

Parameter	R name	Value
Time horizon (n_t)	n.t	50 years
Names of health states (n)	v.n	H, S, D
Annual transition probabilities		
- Disease onset (H to S)	p.HS	0.17
- Recovery (S to H)	p.SH	0.35
Annual mortality		
- All-cause (H to D)	p.HD	0.05
- Disease specific (S to D)	p.SD	0.10
Annual costs		
- Healthy individuals	c.H	\$1,000
- Sick individuals	c.S	\$3,000
- Dead individuals	c.D	\$0
Utility weights		
- Healthy individuals	u.H	1.00
- Sick individuals	u.S	0.60
- Dead individuals	u.D	0.00
Transition rewards		
- Utility decrement of healthy individuals	du.HS	0.10
when transitioning to S		
- Cost of dying	ic.D	\$4,000

01 Initial setup

We start with loading the packages and functions needed.

```
#### 01.1 Load packages and functions ####
library(dplyr)  # to manipulate data
library(reshape2) # to transform data
library(ggplot2) # for nice looking plots
library(scales) # for dollar signs and commas
library(tensorA) # for tensor calculations
```

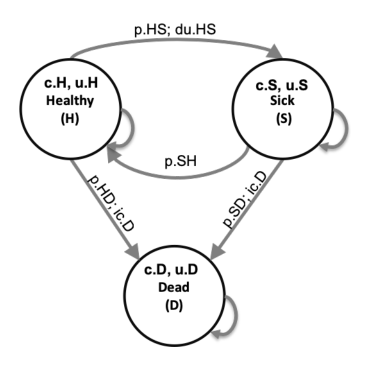


Figure 1: State-transition diagram of the 3-state model

```
#### 01.1.2 Load functions ####
source("../functions/01_model-inputs_functions.R")
source("../functions/02_simulation-model_functions.R")
```

01.2 External parameters

In this section, we specify the staring age of the cohort, the number of cycles, the names of the health states and the initial state vector. The parameters described in the table are combined in a data frame using the f.generate_init_params function.

02 Define and initialize matrices and vectors

In this section, we initialize the matrices and vectors used for storing the data. The transition probability matrix, m.P, is initialized using the function f.create_transition_prob_matrix based on the parameter

values stored in the df.params.init dataframe. The next step is to initialize the state vector s0. All individuals start in the Healthy health state. This state vector is used to inform the first row of the cohort trace matrix m.M.

Equation 1-3

```
#### 02.1 Transition probability matrix ####
# matrix m.P at the first cycle
m.P <- f.create_transition_prob_matrix(df.params = df.params.init)</pre>
#### 02.2 Initial state vector ####
# the cohort start in the Healthy health state
s0 \leftarrow c(H = 1, S = 0, D = 0)
## H S D
## 1 0 0
#### 02.3 Cohort trace
## Create the Markov cohort trace matrix m.M capturing the proportion of the cohort
# in each state at each cycle
m.M <- matrix(0, # initialize cohort trace
              nrow = (n.t + 1), ncol = n.states,
              dimnames = list(0:n.t, v.n))
m.M[1, ] <- s0
                 # store the initial state vector
```

We have determined all parameters for the general set up, we specified our input parameters, initialized all structures and filled the transition probability matrix $\mathtt{m.P}$ and the first row of our cohort trace $\mathtt{m.M}$. The next step is running the Markov model.

03 Traditional cohort trace approach

In this section, we run the Markov model for all cycles. The calculation shown in Equation 3 needs to be performed for all cycles. Therefore, we create a loop starting at t = 1 until t = n.t. The transition probability matrix, m.P is multiplied with the cohort trace m.M[t,], using matrix multiplication, specified in R with %*%, to fill the next row of the m.M[t + 1,]. #### Equation 3

```
for(t in 1:n.t){  # loop through the number of cycles
  # estimate the state vector for the next cycle (t + 1)
  m.M[t + 1, ] <- m.M[t, ] %*% m.P  # Equation 3
}</pre>
```

When printing the first six rows of m.M we see that everyone starts in the Healthy health state and over time the cohort transitions towards sick and dead.

```
head(round(m.M, 3)) # show the first six lines of the Markov cohort trace
```

```
## H S D
## 0 1.000 0.000 0.000
## 1 0.780 0.170 0.050
## 2 0.668 0.226 0.106
## 3 0.600 0.238 0.162
## 4 0.551 0.233 0.216
## 5 0.512 0.222 0.267
```

Running the model using equation 3, results in the traditional cohort trace. This gives us information about state occupation at each cycle. This allows us to apply state rewards (e.g. c.H, u.H, etc.), but it is not possible to include the transition rewards (e.g. du.HS and ic.D). In order to include these rewards, we need to know when individuals made the transition. The dynamics array approach facilitates this and will be explained in the next section.

04 Dynamics array approach

The dynamics array approach starts similar as the traditional cohort trace approach, meaning that the first two sections are identical for the two approaches. The biggest difference between the approaches is the dimensions of the structure to store the model dynamics. In the cohort trace appoarch, we stored all information in matrix m.M of size $n.states \times n.states$, while in the dynamics array approach we add the dimension time resulting in an array with dimensions $n.states \times n.t.$ In R indexing start at 1, therefore, we initialize the array a.A using n.state + 1. This allows us to store the results from cycle 0 until cycle n.t. The initial state vectors s0 is used to inform the initial cycle of the array.

```
a.A <- array(0, dim = c(n.states, n.states, n.t + 1),
dimnames = list(v.n, v.n, 0:n.t)) # initialize array
diag(a.A[, , 1]) <- s0 # store the initial state vector in the diagonal of A
```

Equation 4-7

We are now ready to run the model using the dynamics array appraoch. In this approach, iteratively over time, we multiply the transition probability matrix m.P with array a.A using element-wise mulitplication, indicated with the * in R. The information about all transitions dynamics is stored in the next cycle in a.A.

Equation 4

By printing the first thee cycles of the dynamics array a.A, you can get an impresion of the results. Like in the transition probability matrix m.P, the rows specify in which health state the individual started at the beginning of the cycle, while the columns inform you about where individuals transitioned to. In cycle 0 everyone started in the healthy health states. At cycle 1 we can see that the values look very similar to the transition probabilities. From cycle 2 and onwards the information in a.A becomes more interesting. In cycle 2, we see that 0.78 of the cohort stayed healthy, 0.17 transitioned from Healthy towards Sick and 0.05 of the population died. In addition, we see that 0 of the cohort recovered, 0 stayed Sick and 0.05 died from Sick. All these values sum to 1 since we are still describing what happens to the full cohort over time.

```
a.A[, , 1:3] # shown for two cycles

## , , 0

##

## HSD
```

```
## H 1 0 0
## S 0 0 0
## D O O O
##
##
   , , 1
##
##
        Η
             S
                   D
## H 0.78 0.17 0.05
## S 0.00 0.00 0.00
## D 0.00 0.00 0.00
    , 2
##
##
##
          Η
                  S
## H 0.6084 0.1326 0.039
## S 0.0595 0.0935 0.017
## D 0.0000 0.0000 0.050
sum(a.A[, , 3]) # sum for t = 3
## [1] 1
```

Equation 8

When you sum the values in each column of a.A, the health states towards which the individuals transition to, you get which proportion of the cohort was in each health state for that cycle. The code below shows this for the health state sick.

```
sum(a.A[, "S", 3]) # sum the column of S at t = 3
```

```
## [1] 0.2261
```

By using the colSums function, summing the values in each column of a.A, we can do this for all points in time. By transposing this in turn, we get the traditional cohort trace m.M. We name this matrix, m.M_A, to indicate it is generated via the dynamics array approach and to make it possible toe compare it with the previously generated m.M. When you only do the dynamics array approach you can simply call it m.M.

```
# calculating M from A m.M_A \leftarrow t(colSums(a.A)) # sum over the columns of a.A and transpose
```

Since a Markov model is stochastic, these two approached should give identical results. We check this using the == function. We use rounding on 10 decimals, to avoid wrong FALSE results that have to do with floating point comparison issues. This means that functions allows you to test for equality with a specified difference tolerance.

05 Apply state and transition rewards

We now showed how to run a state-transition cohort model using the dynamics array approach and how to interpret or summarize the results. In this section, we demonstrate how to apply state and transition rewards. We start by initiating and filling two matrices for both costs and effects. To create the cost matrix we make use of the functions f.create_transition_reward_matrix_costs. This functions is informed by the df.params.init data frame, including all cost values. Running the function results in a reward matrix for costs, m.R_costs. On the diagonal are the costs for staying one cycle in that state, while the costs off the diagonal are the costs for staying one stycle in that state plus the transition cost associated with that transition. The function f.create_transition_reward_matrix_effects does the same but then for utilities.

```
#### 05.1 Create reward matrices for both costs and effects ####
m.R_costs <- f.create_transition_reward_matrix_costs(df.params = df.params.init)
m.R_effects <- f.create_transition_reward_matrix_effects(df.params = df.params.init)</pre>
```

Equation 9

These rewards matrices look as follow. We see that staying healthy costs \$1000, while someone that transitions from healthy towards dead makes costs of \$5000. In the effects matrix we see that an individual gets a utility of 1 assign for staying healthy, while when the individual transitions towards Sick the decrement of ic.HS is included, resulting in a utility of 0.90.

```
m.R_costs  # show the reward matrix for costs

## H S D

## H 1000 1000 5000

## S 3000 3000 7000

## D O O O

m.R_effects  # show the reward matrix for effects

## H S D

## H 1.0 0.9 1.0

## S 0.6 0.6 0.6

## D 0.0 0.0 0.0
```

Equation 10

In this section we create outcome arrays, a.O, one for costs, called a.O_costs, and one for effects, called a.O_effects. These arrays show the costs and QALYs generated with each transition at each cycle. By iteratively element-wise multiplication of the reward matrices, m.R_costs and m.R_effects, with array a.A we can fill the outcome arrays.

Please note that we are now showing all these steps in a step-wise appraoch, resulting in having a couple of loops for time. This iterative process can be combined within the same timeloop.

Equation 11

The final step is to calculated to the total costs and QALYs. We start by calculating the expected cost and QALYs per cycle. These values, stored in the vectors v.Costs and v.QALYs, in turn we sum these values. This gives us the total expected cost (TC) and QALYs (TE).

```
# calculate the expected costs per cycle
v.Costs <- rowSums(t(colSums(a.0_costs)))
# calculate the expected QALYs per cycle
v.QALYs <- rowSums(t(colSums(a.0_effects)))</pre>
```

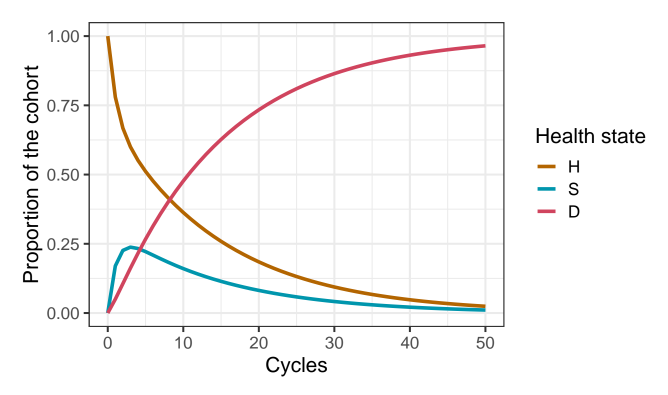


Figure 2: Cohort trace of the model

```
TC <- sum(v.Costs) # calculate the total expected costs

TE <- sum(v.QALYs) # calculate the total expected QALYS

v.Results <- c(TC, TE) # combine the total expected costs and QALYs

names(v.Results) <- c("Costs", "Effect") # name the vector

v.Results # print the results

## Costs Effect

## 28226.5593 14.2835
```

06 Plot cohort trace

The results of a cohort trace are much easier to interpret via a graph. Using the function ggplot can show the proportion of the cohort in each state (y-axis) at each cycle (x-axis).