# Sick-Sicker case study

Supplementary material to the paper: 'A need for change! A coding framework for improving transparency in decision modeling'

DARTH workgroup

# The Sick-Sicker model

In this case-study, we perform a cost-effectiveness analysis (CEA) using a previously published 4-state model called the Sick-Sicker model. (E. Enns et al. 2015) In the Sick-Sicker model, a hypothetical disease affects individuals with an average age of 25 years and results in increased mortality, increased treatment costs and reduced quality of life (QoL). We simulate this hypothetical cohort of 25-year-old individuals over a lifetime (i.e., reaching an age of 100 years old) using 75 annual cycles, represented with n.t. The cohort starts in the "Healthy" health state (denoted "H"). Healthy individuals are at risk of developing the illness, at which point they would transition to the first stage of the disease (the "Sick" health state, denoted "S1"). Sick individuals are at risk of further progressing to a more severe stage (the "Sicker" health state, denoted "S2"), which is a constant probability in this case-study. There is a chance that individuals in the Sick state eventually recover and return back to the Healthy state. However, once an individual reaches the Sicker state, they cannot recover; that is, the probability of transitioning to the Sick or Healthy states from the Sicker state is zero. Individuals in the Healthy state face background mortality that is age-specific (i.e., time-dependent). Sick and Sicker individuals face an increased mortality expressed as a hazard rate ratio (HR) of 3 and 10, respectively, on the background mortality rate. Sick and Sicker individuals also experience increased health care costs and reduced QoL compared to healthy individuals. Once simulated individuals die, they transition to the "Dead" state (denoted "D"), where they remain. Figure 1 shows the state-transition diagram of the Sick-Sicker model. The evolution of the cohort is simulated in one-year discrete-time cycles. Both costs and quality-adjusted life years (QALYs) are discounted at an annual rate of 0.03 %.

Two alternative strategies exist for this hypothetical disease: a no-treatment and a treatment strategy. Under the treatment strategy, Sick and Sicker individuals receive treatment and continue doing so until they recover or die. The cost of the treatment is additional to the cost of being Sick or Sicker for one year. The treatment improves QoL for those individuals who are Sick but has no effect on the QoL of those who are sicker. To evaluate these two alternative strategies, we perform a cost-effectiveness analysis (CEA).

We assume that most of the parameters of the Sick-Sicker model and their uncertainty have been previously estimated and are known to the analyst. However, while we can identify those who are afflicted with the illness through obvious symptoms, we can not easily distinguish those in the Sick state from the those in the Sicker state. Thus, we can not directly estimate state-specific mortality hazard rate ratios, nor do we know the transition probability of progressing from Sick to Sicker. Therefore, we calibrate the model to different epidemiological data. We internally validated the calibrated model by comparing the predicted outputs from the model evaluated at the calibrated parameters against the calibration targets. (Eddy et al. 2012, Goldhaber-Fiebert, Stout, and Goldie (2010))

As part of the CEA, we conducted different deterministic sensitivity analysis (SA), including one-way and two-way SA, and tornado plots. To quantify the effect of parameter uncertainty on decision uncertainty, we conducted a probabilistic sensitivity analysis (PSA) and reported our uncertainty analysis results with a cost-effectiveness acceptability curve (CEAC), cost-effectiveness acceptability frontier (CEAF) and expected loss curves (ELC). We also conducted a value of information (VOI) analysis to determine whether potential future research is needed to reduce parameter uncertainty. All steps of the CEA will be described using the different components of the framework.

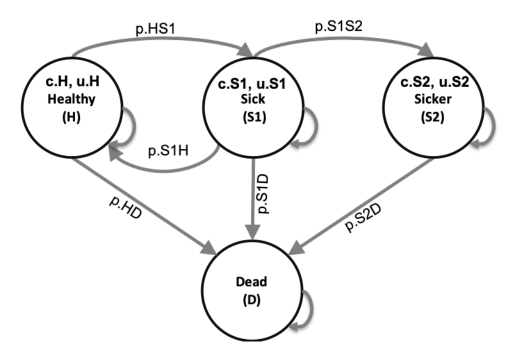


Figure 1: State-transition diagram of the Sick-Sicker model. Healthy individuals can get Sick, die or stay healthy. Sick individuals can recover, transitioning back to healthy, can die, or stay sick. Once individuals are Sicker, they stay Sicker until they die.

# Set up

This report is a supplementary material meant to guide you through the R code of a full functional decision model we use to showcase the DARTH framework described in the manuscript titled A need for change! A coding framework for improving transparency in decision modeling. The code following the framework can be downloaded from the DARTH-git on GitHub (https://github.com/DARTH-git/Decision-Modeling-Framework). We recommend downloading this complete framework as a .zip file containing all directories. Unzip the folder and save it on your computer. The framework is divided into different directories, described in Table 1, that could be accessed from the RStudio project **Decision-Modeling-Framework.Rproj**. In this framework, you will find multiple directories as described in Table 1 of the main manuscript. We refer to the directory names of this framework using *italic* style. This report is created with Markdown and is located in the reports directory of the framework. The figures for the case-study can be found in the figs directory, data required to conduct some of the analyses of the different components are in the data directory and the R scripts with functions, are located in the functions directory. The main R scripts that conduct the analyses of the different components of the framework are stored in the R directory. In this document we do not show all the R code we refer to. Therefore, it is important to follow along while reading this document. To make sure you have all the required packages needed to run all the code, first install all the required packages by running the 00 general-setup.R script in the R directory.

source("R/app0\_packages-setup.R")

# 01 Define model inputs

As described in the main manuscript, in this component we declare all model input variables and set their values. The R script running the analysis of this component is the **01** model-inputs.R file in the R directory.

The input to inform the values is divided in three categories: external, estimated, and calibrated. The majority of the Sick-Sicker model parameters are informed by external data. Only three parameter values

need to be estimated using model calibration.

In this component, we start with the general setup of the model, specifying among others the time horizon, name and number of health states, proportion of the cohort in each of the different health states at the start of the simulation and discount rates. The next step is to specify the external parameters. The initial model parameter values and R variable names are presented in Table 1.

Table 1: Description of the initial parameters with their R name and value of the Sick-Sicker model.

Parameter	R name	Value
Time horizon $(n_t)$	n.t	75 years
Names of health states $(n)$	v.n	H, S1, S2, D
Annual discount rate (costs/QALYs)	d.c/d.e	3%
Annual transition probabilities		
- Disease onset (H to S1)	p.HS1	0.15
- Recovery (S1 to H)	p.S1H	0.5
- Disease progression (S1 to S2) in the time-homogenous model	p.S1S2	0.105
Annual mortality	_	
- All-cause mortality (H to D)	p.HD	age-specific
- Hazard rate ratio of death in S1 vs H	hr.S1	3
- Hazard rate ratio of death in S2 vs H	hr.S2	3
Annual costs		
- Healthy individuals	c.H	\$2,000
- Sick individuals in S1	c.S1	\$4,000
- Sick individuals in S2	c.S2	\$15,000
- Dead individuals	c.D	\$0
- Additional costs of sick individuals treated in S1 or S2	c.Trt	\$12,000
Utility weights		
- Healthy individuals	u.H	1.00
- Sick individuals in S1	u.S1	0.75
- Sick individuals in S2	u.S2	0.50
- Dead individuals	u.D	0.00
Intervention effect		
- Utility for treated individuals in S1	u.Trt	0.95

Age-specific background mortality for healthy individuals is represented by the US population in 2015 and obtained from the Human Mortality database. This information is stored in the 01\_all-cause-mortality.csv file in the data directory. Based on this .csv file a vector with mortality rates by age is created using the f.load\_mort\_data function in the 01\_model-inputs\_functions script. This functions gives us the flexibility to easily import data from other coutries or years.

```
print.function(f.load_mort_data) # print the function
```

An other function in the **01\_model-inputs\_functions** script, is the f.load\_all\_parms function. This function, which is actually using the f.load\_mort\_data function, loads all parameters for the decision model from multiple sources and creates a list that contains all parameters and their values.

```
print.function(f.load_all_params) # print the function
```

```
## function (file.init = "data/01_init-params.csv", file.mort = "data/01_all-cause-mortality.csv")
## {
##
       df.params.init <- read.csv(file = file.init)</pre>
##
       v.r.mort_by_age <- f.load_mort_data(file = file.mort)</pre>
##
       l.params.all <- with(as.list(df.params.init), {</pre>
##
            v.age.names <- n.age.init:(n.age.init + n.t - 1)</pre>
##
           v.n <- c("H", "S1", "S2", "D")
##
           n.states <- length(v.n)
##
           v.s.init \leftarrow c(H = 1, S1 = 0, S2 = 0, D = 0)
##
            l.params.all <- list(n.age.init = n.age.init, n.t = n.t,</pre>
##
                v.age.names = v.age.names, v.n = v.n, n.states = n.states,
##
                v.s.init = c(H = 1, S1 = 0, S2 = 0, D = 0), v.r.mort_by_age = v.r.mort_by_age)
##
           return(l.params.all)
##
       })
##
       1.params.all <- c(1.params.all, df.params.init)</pre>
## }
## <bytecode: 0x7f987734d460>
```

The f.load\_all\_params function is informed by the arguments file.init and file.mort. The file.init argument is a string with the location and name of the file with initial set of parameters. The inital parameter values for our case-study are stored in the 01\_init-params.csv file located in the data directory. The f.load\_all\_params function read this .csv file into the function environment as a dataframe called, df.params.init.

The file.mort argument is a string with the location and name of the file with mortality data. As described before, in our case-study this is the **01\_all-cause-mortality.csv** file. Within the f.load\_all\_parms function, the f.load\_mort\_data function is used to create a vector with mortality rates from the .csv data.

After loading all the information, the f.load\_all\_params generates a list called,l.params.all, including all parameters for the model including the general setup parametes and the vector of mortality rates. The function also stores the dataframe df.params.init with the initial set of parameters in the list. This is all executed in the in the 01\_model-inputs.R script by running the code below.

For the Sick-Sicker model we do not have to estimated parameters, but we do have three parameters that need to be estimated via model calibration. In this stage of the framework, we simply set these parameters to valid "dummy" values that are compatible with the next phase of the analysis, model implementation, but are ultimately just placeholder values until we conduct the calibration phase. This means that these values will be replaced by the best-fitted calibrated values after we performed the calibration in component 3.

Using a function to create a list of base-case parameters to have all model parameters in a single object is very useful, because this object will have to be updated for the calibration and the different sensitivity analyses in components 3 and 5 of the framework, respectively. Below, we guide you through the components of the function.

#### 02 Model implementation

In this component, we build the backbone of the decision analysis: the implementation of the model. This component is performed by the **02\_simulation-model.R** script. This file itself is not very large. It simply loads some packages, sources the input from component 01, sources the function **f.decision\_model** that is used the capture the dynamic process of the Sick-Sicker example, runs this function and stores the output. The output of the model is the traditional cohort trace, describing how the cohort is distributed among the

different health states over time, which is plotted at the end of this script. This trace will be used in many of the other components.

The function f.decision\_model is defined in the 02\_simulation-model\_functions.R file. As described in the paper, constructing a model as a function at this stage facilitates subsequent stages of the model development and analysis, as these processes will all call the same model function, but pass different parameter values and/or calculate different final outcomes based on the model outputs. In the next part, we will describe the code within the function.

# print.function(f.decision\_model) # print the code of the function

```
## function (1.params.all, verbose = FALSE)
## {
##
                with(as.list(l.params.all), {
##
                         p.HDage <- 1 - exp(-v.r.mort_by_age[(n.age.init + 1) +</pre>
##
                                  0:(n.t - 1)])
##
                         p.S1Dage \leftarrow 1 - exp(-v.r.mort_by_age[(n.age.init + 1) +
##
                                  0:(n.t - 1)] * hr.S1)
##
                         p.S2Dage <- 1 - exp(-v.r.mort_by_age[(n.age.init + 1) +</pre>
##
                                  0:(n.t - 1)] * hr.S2)
##
                         a.P \leftarrow array(0, dim = c(n.states, n.states, n.t), dimnames = list(v.n,
##
                                  v.n, 0:(n.t - 1))
                         a.P["H", "H", ] <- (1 - p.HDage) * (1 - p.HS1)
##
                         a.P["H", "S1", ] <- (1 - p.HDage) * p.HS1
##
                         a.P["H", "D", ] <- p.HDage
##
                         a.P["S1", "H", ] <- (1 - p.S1Dage) * p.S1H
##
                         a.P["S1", "S1", ] \leftarrow (1 - p.S1Dage) * (1 - (p.S1S2 + p.S1Dage)) * (1 - (p.S1S2 + p.S1
##
##
                                  p.S1H))
                         a.P["S1", "S2", ] <- (1 - p.S1Dage) * p.S1S2
##
                         a.P["S1", "D", ] <- p.S1Dage
##
                         a.P["S2", "S2", ] <- 1 - p.S2Dage
##
                         a.P["S2", "D", ] <- p.S2Dage
##
                         a.P["D", "D", ] <- 1
##
                         m.indices.notvalid <- arrayInd(which(a.P < 0 | a.P >
##
                                  1), dim(a.P))
##
##
                         try(if (dim(m.indices.notvalid)[1] != 0) {
##
                                  v.rows.notval <- rownames(a.P)[m.indices.notvalid[,
##
##
                                  v.cols.notval <- colnames(a.P)[m.indices.notvalid[,
##
                                  v.cycles.notval <- dimnames(a.P)[[3]][m.indices.notvalid[,</pre>
##
##
                                            3]]
                                  df.notvalid <- data.frame(`Transition probabilities not valid:` = matrix(pasteO(paste(v...)))</pre>
##
                                            v.cols.notval, sep = "->"), "; at cycle ", v.cycles.notval),
##
##
                                           ncol = 1), check.names = FALSE)
##
                                  if (verbose) {
##
                                           message("Not valid transition probabilities")
##
                                            stop(print(df.notvalid), call. = FALSE)
                                  }
##
##
                         })
                         valid <- apply(a.P, 3, function(x) all.equal(sum(rowSums(x)),</pre>
##
##
                                  n.states))
##
                         if (!isTRUE(all.equal(as.numeric(sum(valid)), as.numeric(n.t)))) {
##
                                  if (verbose) {
##
                                            stop("This is not a valid transition Matrix")
```

```
}
##
           }
##
##
           m.M <- matrix(0, nrow = (n.t + 1), ncol = n.states, dimnames = list(0:n.t,
                v.n))
##
##
           m.M[1, ] <- v.s.init
           for (t in 1:n.t) {
##
                m.M[t + 1, ] \leftarrow m.M[t, ] %*% a.P[, , t]
##
           }
##
##
           return(list(a.P = a.P, m.M = m.M))
##
       })
## }
   <bytecode: 0x7f9877392628>
```

The f.decision\_model function is informed by the argument 1.params.all. Via this argument we give the function a list with with all parameters of the decision model. For the Sick-Sicker model, these parameters are stored in the list 1.params.all, which we passed into the function as shown below.

```
1.out.stm <- f.decision_model(1.params.all = 1.params.all) # run the function</pre>
```

This function itself has all the mathematical equations of the decision models coded inside. It starts by calculating the age-specific transition probabilities from all non-dead states based on the vector of age-specific mortality rates v.r.asr. These parameters will become vectors of length n.t, describing the probability to die for all ages from all non-dead states.

The next part of the function, creates an array that stores the age-specific transition probability matrices in each of the third dimension. The transition probability matrix is a core component of a state-transition cohort model(Iskandar 2018). This matrix contains the probabilities of transitioning from the current health state, indicated by the rows, to the other health states, specified by the columns. Since we have age-specific transition probabilities, the transition probability matrix is different at each cycle. These probabilities are only depending on the age of the cohort, and not on other events; therefore, we can generate all matrices at the start of the model. This results in n.t different age-specific matrices that are stored in an array, called a.P, of dimensions n.s x n.s x n.t. After initializing the array, it is filled with the transition probability stored in the list. When running the model, we can index the correct transition probability matrix corresponding with the current age of the cohort. We then added some sanity checks to make sure that the transition matrices and the transition probabilities are valid. The transition probability matrices stored in the array a.P, for the first three and last cycle, are shown below.

```
1.out.stm$a.P[, , 1:3] # show the first three time-points of a.P
```

```
##
   , 0
##
##
           Η
                  S1
                          S2
                                     D
## H 0.8491385 0.1498480 0.0000000 0.001013486
## S1 0.4984813 0.3938002 0.1046811 0.003037378
## S2 0.0000000 0.0000000 0.9899112 0.010088764
##
##
     1
##
##
                          S2
           Η
                  S1
                                     D
## H 0.8491513 0.1498502 0.0000000 0.0009985012
## S1 0.4985037 0.3938180 0.1046858 0.0029925135
## S2 0.0000000 0.0000000 0.9900597 0.0099402657
##
## , , 2
```

```
##
##
                              S2
            Η
                     S1
                                          D
## H
     0.8490910 0.1498396 0.0000000 0.001069428
  S1 0.4983976 0.3937341 0.1046635 0.003204853
  S2 0.0000000 0.0000000 0.9893570 0.010642959
     1.out.stm$a.P[, , l.params.all$n.t] # show the last
##
            Η
                                         D
                     S1
                               S2
## H
     0.6055199 0.1068564 0.00000000 0.2876237
## S1 0.1807584 0.1427991 0.03795926 0.6384833
## S2 0.0000000 0.0000000 0.03365849 0.9663415
## D 0.0000000 0.0000000 0.00000000 1.0000000
```

By comparing these probability matrices, we observe an increase in the probabilities of transitioning to death from all health states.

After the array is filled, the cohort trace matrix, m.M, of dimensions n.t x n.s is initialized. This matrix will store the state occupation at each point in time. The first row of the matrix is informed by the initial state vector v.s.init. For the remaining points in time, we iteratively multiply the cohort trace with the age-specific transition probability matrix corresponding to the specific cycle obtained by indexing the array a.P appropriately. All the outputs and relevant elements of the decision model are stored in a list, called l.out.stm. This list contains the array of the transition probability matrix for all cycles t and the cohort trace m.M.

```
head(1.out.stm$m.M)
                       show the top part of the cohort trace
##
            Η
                    S1
                               S2
                                            D
## 1 0.8491385 0.1498480 0.00000000 0.001013486
## 2 0.7957468 0.1862564 0.01568695 0.002309774
## 3 0.7684912 0.1925699 0.03501425 0.003924648
## 4 0.7484793 0.1909659 0.05478971 0.005765035
## 5 0.7306193 0.1873106 0.07413838 0.007931783
tail(l.out.stm$m.M)
                      # show the bottom part of the cohort trace
##
                                                 D
               Н
                          S1
                                       S2
## 70 0.009928317 0.0022433565 2.035951e-04 0.9876247
## 71 0.007153415 0.0015935925 1.311619e-04 0.9911218
## 72 0.005058845 0.0011174473 8.674540e-05 0.9937370
## 73 0.003460336 0.0007552206 5.436484e-05 0.9957301
## 74 0.002289594 0.0004937081 3.298632e-05 0.9971837
## 75 0.001475636 0.0003151589 1.985106e-05 0.9981894
```

Using the code below, we can graphically show the model dynamics by plotting the cohort trace. Figure 2 shows the distribution of the cohort among the different health states at each time point.

#### 03 Model calibration

In this component, we calibrate unknown model parameters by matching model outputs to specified calibration targets. Specifically, we calibrate the Sick-Sicker model to match survival, prevalence and the proportion who are Sicker, among all those afflicted (Sick+Sicker). We used a Bayesian calibration approach using the incremental mixture importance sampling (IMIS) algorithm (Steele, R., Raftery A., Emond 2006), which has been used to calibrate health policy models (Raftery, A., Bao (2010), Menzies, Pandya, and Kim (2017), Rutter, C., Ozik, J., DeYoreo, M., Collier (2018)). Bayesian methods allow us to quantify the uncertainty in

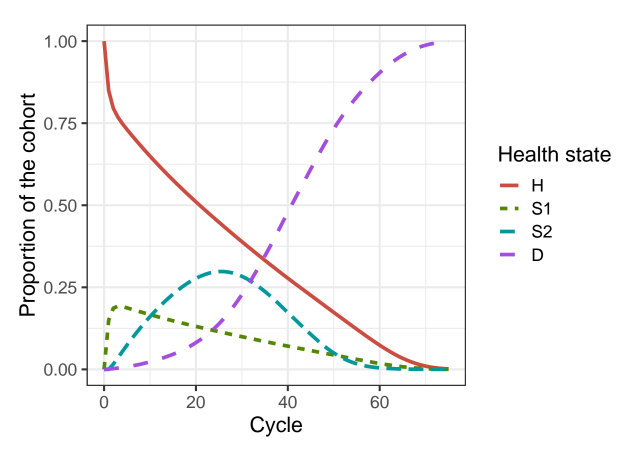


Figure 2: Cohort trace of the Sick-Sicker cohort model

the calibrated parameters even in the presence of non-identifiability (Alarid-Escudero et al. (2018)). This analysis is coded in the  $03\_calibration.R$  file. The target data is stored in the  $03\_calibration-targets.RData$  file. Similar to component 02, in the section **03.1 Load packages, data and functions** we start by loading inputs and functions. In addition, we load the calibration targets data into the R workspace. In the next section, **03.2 Visualize targets**, we plot each of the calibration targets with their confidence intervals.

In section **03.3 Run calibration algorithms**, we set the parameters we like to calibrate to fixed values and test if the function **f.calibration\_out** that produces model outputs corresponding to the calibration targets works. This function takes a vector of parameters that need to be calibrated and a list with all parameters of decision model and computes model outputs to be used for calibration routines.

```
print.function(f.calibration_out) # print the funtions
```

```
## function (v.params.calib, l.params.all)
## {
##
       1.params.all <- f.update_param_list(1.params.all = 1.params.all,</pre>
##
           params.updated = v.params.calib)
##
       1.out.stm <- f.decision_model(1.params.all = 1.params.all)</pre>
       v.os <- 1 - l.out.stm$m.M[, "D"]
##
       v.prev <- rowSums(1.out.stm$m.M[, c("S1", "S2")])/v.os</pre>
##
##
       v.prop.S2 <- 1.out.stm$m.M[, "S2"]/rowSums(1.out.stm$m.M[,</pre>
##
            c("S1", "S2")])
       1.out <- list(Surv = v.os[c(11, 21, 31)], Prev = v.prev[c(11, 21, 31)]
##
##
           21, 31)], PropSicker = v.prop.S2[c(11, 21, 31)])
##
       return(1.out)
## }
```

This function is informed by two argument v.params.calib and l.params.all. The vector v.params.calib contains the values of the three parameters of interest and the list l.params.all contains all parameters of the decision model. The placeholder values are replaced by v.params.calib and with these values the model is evaluated. This is done by running the f.decision\_model function, described in component 02. This results in a new list with output of the model corresponding to the parameter values in the v.params.calib. With this new decision model output, the overall survival, disease prevalence and the proportion of Sicker in the Sick and Sicker states are calculated. The estimated values for these epidemiological outcomes at different timepoints are combined in a list called l.out produced but he f.calibration\_out.

Once we make sure this code works, we specify the calibration parameters in section **03.3.1 Specify** calibration parameters. These include setting the seed for the random number generation, specifying the number of random samples to obtain from the calibrated posterior distribition, the name of the input parameters and the range of these parameters that will inform the prior distributions of the calibrated parameters, and the name of the calibration targets: Surv, Prev, PropSick.

In the next section, **03.3.2 Run IMIS algorithm**, we calibrate the Sick-Sicker model with the IMIS algorithm. For this case-study, we assume a normal likelihood and uniform priors but other could be used. For a more detailed description of IMIS for Bayesian calibration, different likelihood functions and prior distributions, we refer the reader to the tutorial for Bayesian calibration by Menzies et al. (Menzies, Pandya, and Kim 2017). We use the IMIS function from the IMIS package that calls the functions likelihood, sample.prior and prior, to draw samples from the posterior distribution. (Raftery and Le Bao 2012) The functions are specified in the **03\_calibration\_functions.R** file. For the IMIS function, we specify the incremental sample size at each iteration of IMIS, the desired posterior sample size at the resample stage, the maximum number of iterations in IMIS and the number of optimizers which could be 0. The function returns a list, which we call **1.fit.imis**, with the posterior samples, the diagnostic statistics at each IMIS iteration and the centers of Gaussian components(Raftery and Le Bao 2012). We store the posterior samples in the matrix m.calib.post.

We then expolore this posterior distributions in section **03.4 Exploring posterior distribution**. We start by estimating the posterior mean, median and 95% credible interval, the mode and the maximum-a-posteriori

Table 2: Summary statistics of the posterior distribution

	Mean	2.5%	50%	97.5%	Mode	MAP
p.S1S2	0.1077822	0.0966154	0.107619	0.1206907	0.1071694	0.1078278
hr.S1	2.7151807	1.1007592	2.608970	4.4139330	1.9654838	2.2970418
hr.S2	9.5435354	7.1631569	9.534398	11.8839979	9.2717728	9.8870386

(MAP). To estimate the mode, we use the package modeest. (Poncet 2018) All for these summary statistics are combined in a dataframe called df.posterior.summ. Table XX shows the summary statistics of the posterior distribution.

In section **03.4.2 Visualization of posterior distribution**, we generate a pairwise scatter plot of the calibrated parameters (Figure 4) and a 3D scatter plot of the joint posterior distribution (Figure 3). These figures are saved in the *figs* directory.

Finally, the posterior distribution and MAP estimate from the IMIS calibration are stored in the file 03\_imisoutput. RData. Storing this data as an Rdata file allows to import the data in following sections without needing to re-runn the calibration component.

# 04 Validation

In this section, we check the internal validity of our Sick-Sicker model before we move on to the analysis components. To internally validate the Sick-Sicker model, we compare the model-predicted output evaluated at posterior parameters against the calibration targets. This is all done in the **04\_validation.R** script by loading all previously described functions and the generated calibration targets.

In section **04.2 Compute model-predicted outputs**, we compute the model-predicted outputs for each sample of posterior distribution as well as for the MAP estimate. We then use the function <code>f.data\_sumary</code> to summarize the model-predicted posterior outputs into different summary statistics.

```
print.function(f.data_summary)
```

```
## function (data, varname, groupnames)
## {
##
       require(plyr)
       summary_func <- function(x, col) {</pre>
##
           c(mean = mean(x[[col]], na.rm = TRUE), median = quantile(x[[col]],
##
               probs = 0.5, names = FALSE), sd = sd(x[[col]], na.rm = TRUE),
##
##
               lb = quantile(x[[col]], probs = 0.025, names = FALSE),
               ub = quantile(x[[col]], probs = 0.975, names = FALSE))
##
##
       data_sum <- ddply(data, groupnames, .fun = summary_func,
##
##
           varname)
##
       data sum <- plyr::rename(data sum, c(mean = varname))</pre>
##
       return(data_sum)
## }
## <bytecode: 0x7f9877e62938>
```

This function is informed by three arguments, data, varname and groupnames.

The computation of the model-predicted outputs using the MAP estimate is done by inserting the v.calib.post.map data into the previously described f.calibration\_out function. This function creates a list including the estimated values for survival, prevalence and the proportion of sicker individuals at cycles 10, 20 and 30.

In sections 04.6 Internal validation: Model-predicted outputs vs. targets, we check the internal

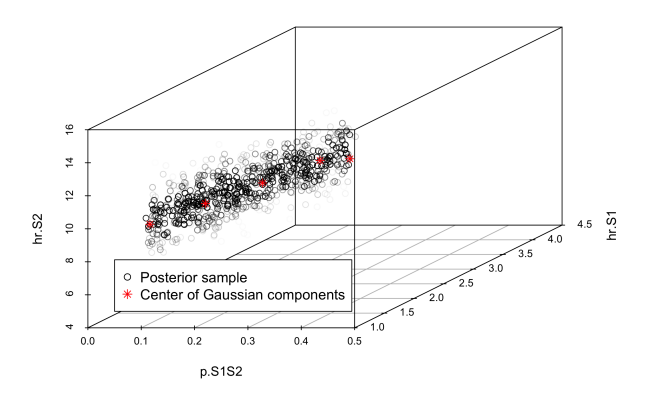


Figure 3: Joint posterior distribution

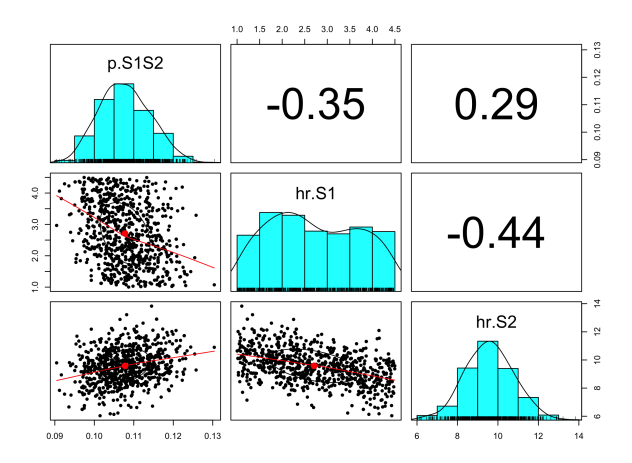


Figure 4: Pairwise posterior distribution of calibrated parameters

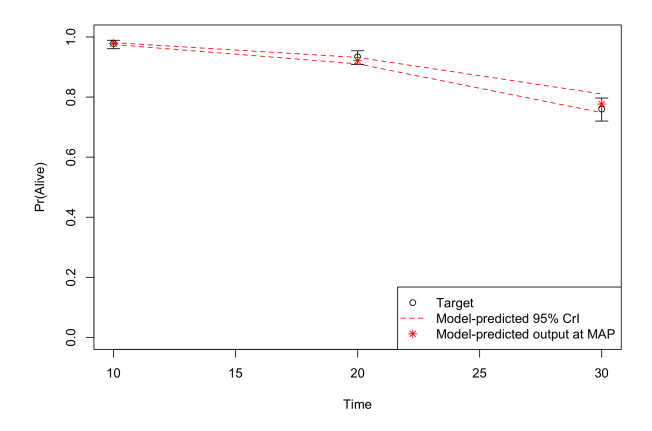


Figure 5: Survival data: Model-predicted outputs vs targets

validation by plotting the model-predicted outputs against the calibration targets Figure (5-7). The generated plots are saved as .png files, which could be used in reports later on without the need of re-running the code.

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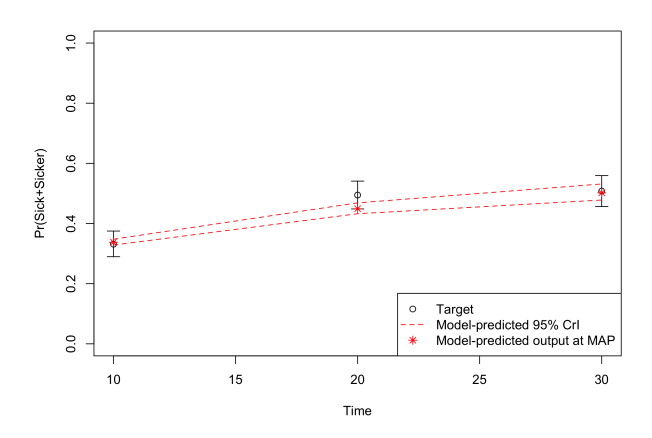


Figure 6: Prevalence data of sick individuals: Model-predicted output vs targets

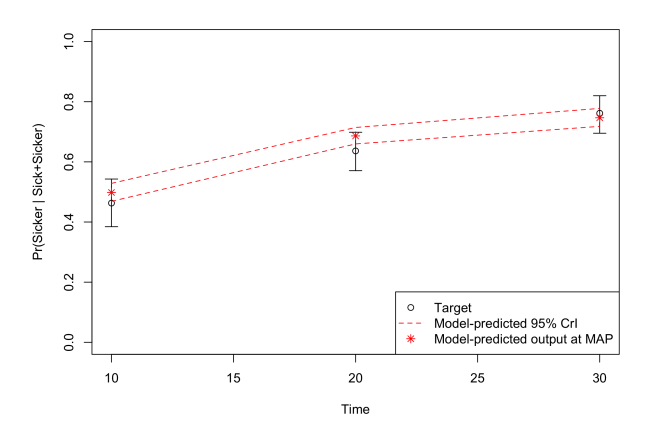


Figure 7: Proportion who are Sicker, among all those afflicted (Sick + Sicker): Model-predicted output

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