Software Documentation

GUTS: Software for the Calculation of the Likelihood Function of the GUTS Model

Version 0.3.2

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GUTS is a software for the fast calculation of the logarithm of the likelihood of an empirical survival model. **GUTS** is currently available as an \mathcal{R} -package. This document describes the software as well as its typical usage.

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Preliminaries

This document was created using "LATEX" and "Sweave" (package SWEAVE, Leisch, 2002) with \mathcal{R} , version 3.0.2 (R Development Core Team, 2013). A function is written function(), a package is written PACKAGE, \mathcal{R} input is marked R: input..., and \mathcal{R} output is marked output. All \mathcal{R} code is set

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in a framed box. Note that it may be required to install the latest version of \mathcal{R} in order to run the software and the examples provided below (see section 5).

```
R: version
platform
                x86_64-apple-darwin10.8.0
                x86 64
arch
                darwin10.8.0
os
                x86_64, darwin10.8.0
system
status
major
                3
                0.2
minor
                2013
vear
month
                09
                25
day
                63987
svn rev
language
                R
version.string R version 3.0.2 (2013-09-25)
nickname
                Frisbee Sailing
```

1 Theoretical Background

GUTS Jager, Albert, Preuss and Ashauer (2011) is a model for survival of organisms, exposed to any kind of quantifiable stress. The time-dependent stressor, C(t), is assumed to cause a time-dependent damage, D(t), which is described by the linear differential equation

$$\dot{D}(t) = k_r (C(t) - D(t)), \tag{1}$$

where k_r is called recovery rate. The damage is the same for all individuals. However, the individuals are assumed to have different thresholds, beyond which the damage increases their probability to die. Thus, the model combines two sources of stochasticity: On the one hand, death is considered a stochastic event, whose probability increases linearly with the damage, once it exceeds a certain threshold. That is, there is stochasticity at individual level. On the other hand, this threshold is assumed to vary stochastically over the population. Thus, there is stochasticity at population level.

The hazard, $h_z(t)$, of an individual with threshold z is determined by the formula

$$h_z(t) = k_k \max(D(t) - z, 0) + h_b,$$
 (2)

where k_k is called *killing rate* and h_b is the *background mortality*. The hazard, in turn, determines the individual's probability to survive until time t, $S_z(t)$, via the linear differential equation

$$\dot{S}_z(t) = -h_z(t)S_z(t). \tag{3}$$

Finally, each individual is assumed to draw its z from a distribution, $f_{\theta}(z)$, on the positive real axis. Hence, the parameter vector of the model reads as

$$\boldsymbol{\theta} = (h_b, k_r, k_k, \dots), \tag{4}$$

where the additional arguments are supposed to determine the distribution $f_{\theta}(z)$.

Combining equations (2) and (3), we find that the probability for an arbitrarily chosen member of the population to survive until time t is given by the formula

$$S_{\theta}(t) = \int \exp\left(-k_k \int_0^t \max(D(\tau) - z, 0)d\tau - h_b t\right) f_{\theta}(z)dz.$$
 (5)

Let $\mathbf{y} = (y_0, y_1, \dots, y_n)$ denote a time series of survivors, counted at times $(t_0 = 0, t_1, \dots, t_n)$, and set $y_{n+1} = 0$. Then, the logarithm of the likelihood, $f(\mathbf{y}|\boldsymbol{\theta})$, of the model output \mathbf{y} given the parameters $\boldsymbol{\theta}$ is, up to $\boldsymbol{\theta}$ -independent terms, given by the formula

$$\ln f(\mathbf{y}|\boldsymbol{\theta}) = \sum_{i=1}^{n+1} (y_{i-1} - y_i) \ln(S_{\boldsymbol{\theta}, i-1} - S_{\boldsymbol{\theta}, i}), \qquad (6)$$

where we have set

$$S_{\boldsymbol{\theta},i} = S_{\boldsymbol{\theta}}(t_i), \quad S_{\boldsymbol{\theta},n+1} = 0.$$
 (7)

2 The Algorithm

The calculation of the log-likelihood requires two numerical integrations (see eq. (5)), and has, therefore, two large numbers, N and M. The following algorithm is of the order $\mathcal{O}(N) + \mathcal{O}(M)$. It is based on the approximation

$$S_{i} = \int \exp\left[-k_{k} \int_{0}^{t_{i}} \max(0, D(\tau) - z) d\tau - h_{b} t_{i}\right] f_{\mathbf{\theta}}(z) dz$$

$$\approx \frac{1}{N} \sum_{j=1}^{N} \exp\left[-k_{k} \Delta \tau \sum_{D_{l} > z_{j}} (D_{l} - z_{j}) - h_{b} t_{i}\right]$$

$$= \frac{1}{N} e^{-h_{b} t_{i}} \left(e^{-k_{k} \Delta \tau (e_{N} - z_{N} f_{N})} + e^{-k_{k} \Delta \tau (e_{N} + e_{N-1} - z_{N-1} (f_{N} + f_{N-1}))} + \dots + e^{-k_{l} \Delta \tau (e_{N} + \dots + e_{1} - z_{1} (f_{N} + \dots + f_{1}))}\right), \quad (8)$$

for an ordered sample $z_1 < \cdots < z_N$ from $f_{\boldsymbol{\theta}}(z)$, and with $D_l = D(\tau_l)$ on a grid $\tau_0 < \cdots < \tau_{M-1}$. The inner sum in the second line extends over all D_l , for which $\tau_l < t_i$, and we have set $\Delta \tau = t_n/M$. Furthermore,

$$e_j = \sum_{z_j < D_l < z_{j+1}} D_l \,, \tag{9}$$

and

$$f_i = \sharp \{ D_l | z_i < D_l < z_{i+1} \}, \tag{10}$$

for $1 \leq j \leq N$ (Set $z_{N+1} = \infty$).

The corresponding algorithm for the calculation of (6) reads as follows:

- 1. Draw N thresholds from $f_{\theta}(z)$ and order them $z_1 < \cdots < z_N$.
- 2. Refine the grid $t_0 < \cdots < t_n$ to a fine grid $\tau_0 < \cdots < \tau_{M-1}$.
- 3. Set i = 0.
- 4. Solve eq. (1), for $t_i \leq \tau_l \leq t_{i+1}$, using equation

$$D_{l} = D(\tau_{l}) = D(s_{k})e^{-k_{r}(\tau_{l}-s_{k})} + C_{k}\left(1 - e^{-k_{r}(\tau_{l}-s_{k})}\right) + \frac{C_{k+1} - C_{k}}{s_{k+1} - s_{k}}\left(\tau_{l} - s_{k} - k_{r}^{-1} + k_{r}^{-1}e^{-k_{r}(\tau_{l}-s_{k})}\right), \quad (11)$$

for $s_k \leq \tau_l \leq s_{k+1}$.

5. Update (9) and (10), for $1 \le j \le N$. (This can be done in time $\mathcal{O}(1)$, for each D_l .)

6. Calculate S_i using the recursion:

$$F_j = F_{j+1} + f_j,$$
 (12)

$$E_j = E_{j+1} + e_j \,, \tag{13}$$

$$S_{i,j} = S_{i,j+1} + \exp(-k_k \Delta \tau (E_j - F_j z_j)),$$
 (14)

for $j = N - 1, \dots, 1$ and with $S_{i,N} = \exp(-k_k \Delta \tau (E_N - F_N z_N))$ and $F_N = f_N$, $E_N = e_N$. Then,

$$S_i = \frac{1}{N} e^{-h_b t_i} S_{i,1} \,. \tag{15}$$

- 7. Increment i and go to 4.
- 8. Calculate the log-likelihood function according to equation (6).

3 The C++ Class

The **GUTS** class allows to store the time series of exterior concentrations of the stressor, $\mathbf{C} = (C(s_0), \dots, C(s_m))$, the data, i.e., the time series of survivors, $\mathbf{y} = (y(t_0), \dots, y(t_n))$, parameter values, $\boldsymbol{\theta} = (h_b, k_r, k_k, \dots)$, of the model and the distribution, $f_{\boldsymbol{\theta}}(z)$, from which the thresholds of the model are sampled. Furthermore, it provides a method to generate a sample, to calculate damages, the survival probabilities, and the logarithm of the likelihood (see section 2).

Below follows a brief description. Refer to the source code to check for more details.

3.1 Fields of the C++ class

The GUTS C++ class has no public fields. Modifications of an existing GUTS object must therefore be made using setter methods. However, protected fields represent the attributes of an object and can be accessed using getter methods (see section 3.2). Of particular interest are the following attributes (due to programming conventions C++ field names may differ from the mathematical notations above):

- Title: the title of a GUTS experiment. Currently unused
- C: vector of (exterior) concentrations (C_0, C_1, \ldots, C_m) .
- Ct: vector of time points of concentrations $(0 = s_0 < s_1 < \cdots < s_m)$.
- y: vector of survivors (y_0, y_1, \ldots, y_n) .
- yt: vector of time points of survivors $(0 = t_0 < t_1 < \cdots < t_n \le s_m)$.
- par: parameter vector $(\boldsymbol{\theta} = (h_b, k_r, k_k, \dots))$ with the following parameters:
 - 1. background mortality rate (h_b)
 - 2. recovery rate (k_r)
 - 3. killing rate (k_k)

The additional arguments $(par_4...)$ determine parameters of the distribution from which thresholds are sampled. Currently, only the lognormal distribution is implemented, and the additional parameters are its mean and standard deviation. Note that this differs from the implementation in the \mathcal{R} function rlnorm() where the parameters denote mean and standard deviation of the corresponding normal distribution. If attribute dist (see below) is "empirical", parameters for the distribution are ignored.

• M: number of grid points on the time axis for the numerical integration (numerical exactness). Defaults to 10000.

- dist: name of the distribution to sample from (currently implemented "lognormal", or "empirical"). Defaults to "lognormal".
- N: number of threshold samples (numerical exactness). Defaults to 10000.
- z: the actual sample of size N, either generated from dist with parameters from par, or provided as an ascendingly ordered positive numeric vector.
- D: vector of damages generated during the calculation of the survival probabilities.
- S: vector of survival probabilities.
- LL: the loglikelihood.

Note that in the source code each attribute is prefixed with an m indicating that this is a member variable set by the corresponding method.

3.2 Methods of the C++ Class

The C++ Class has setter methods, getter methods, and methods to compute values and vectors in a **GUTS** object.

The setter methods set up a complete **GUTS** experiment.

- setTitle(string Title): set the title of a GUTS experiment to Title. Currently unused.
- setConcentrations(vector<double> C, vector<double> Ct): set the vectors of concentrations (C) and concentration time points (Ct). Ct must start at 0. C and Ct must have the same length.
- void setSurvivors(vector<int> y, vector<double> yt): set the vectors of survivors (y) and survivor time points (yt). yt must start at 0. y and yt must have the same length.
- setParameters(vector<double> par): set the vector of parameters. See 3.1 for details.
- setTimeGridPoints(int M): set the number of grid point on the time-axis. See 3.1 for details.
- setDistribution(string dist): set the distribution to dist. See 3.1 for details.
- setSampleLength(int N): set the sample length to N. See 3.1 for details.
- setSample(vector<double> z): do not sample, but use the provided sample z instead. Using this method will bypass the sampling procedure. However, the vector is checked for consistency, and a sorted copy is created and assigned to z.

Each getter method is pasted using the prefix "get" plus the variable's name, e.g. getC for getting the vector of concentrations. Note, that in contrast to combined setters (e.g., setConcentrations()) getters are not combined, i.e., there is one getter each for access to the concentrations and the concentration time points. Additional getters are:

- getD(): returns vector<double> D, the vector of damages.
- getS(): returns vector<double> S, the vector of survival probabilities
- getLL(): returns double LL, the loglikelihood
- getErrors: returns vector<bool> Errors, a vector of booleans indicating errors. An error at the position exists if the element at the position is true.
- getErrorMessages(): returns vector<string> ErrorMessages, a vector of strings expressing what error occurred at the position.

The method showObject() prints formatted content of a GUTS object to the console output.

In addition to merely set- and get-methods, a **GUTS** object provides methods for calculating/computing values or vectors.

- calcSample() calculates a sample from parameters and the value of the field N.
- calcSurvivalProbabilities() calculates the survival probabilities. This method is overloaded and available in three variants:
 - 1. without an argument: use the values present in a GUTS object for the calculation
 - 2. with argument vector<double> St: St donate the survivor time points used for the calculation. A vector of the same length is created and filled with 0. Both vectors are supplied to the method setSurvivors(...). Note that this will change the vector of survivors!
 - 3. with argument int Stlength: A vector of integers starting at zero and of length Stlength is created, as well as a corresponding vector of survivors filled with 0. Bot vectors are supplied to the method setSurvivors(). Note that this will change the vector of survivors!
- calcLoglikelihood() calculates the loglikelihood of a properly set up **GUTS** object. The method sets LL.

Two more protected methods are available in **GUTS**:

- doCalcSampleLognormal calculates a sample from the lognormal distribution.
- doCalcSurvivalProbabilities is the work horse for the calculation of the survival probabilities for either of the calcSurvivalProbabilities-version.

However, these are protected methods of the class and may not be called directly.

4 Implementation in R

GUTS is exposed to \mathcal{R} through the deployment of Rcpp (Eddelbuettel and Francois, 2011). GUTS is contained in a module (modguts) and can be used in \mathcal{R} via the S4 reference class Rcpp_GUTS. Except the Title-method, all public setter and calculation methods are exposed to \mathcal{R} and can be used on \mathcal{R} -objects with the appropriate signature. Except the getter of Errors and Title, all getters are exposed to \mathcal{R} as *fields* of an \mathcal{R} -object of class Rcpp_GUTS. The show method (showObject) was rewritten in \mathcal{R} to account for special formatting capabilities of \mathcal{R} compared to console out.

In addition to the methods of the C++ class, the \mathcal{R} -implementation has two S3 generic functions, print() and logLik().

4.1 Dataset Diazinon

ToDo: more to explain here

The \mathcal{R} -package also contains a small data set for use with GUTS. diazinon is a list with 13 slots:

- Description: a short line of description
- C1-C3: 3 vectors of concentrations of diazinon
- Ct1-Ct3: 3 time vectors corresponding to the concentrations vectors
- y1-y3: 3 vectors of survivors
- yt1-yt3: 3 time vectors corresponding to the survivors vectors

```
R: data(diazinon)
R: diazinon
$Description
[1] "3 experiments with survivors exposed to concentrations of diazinon"
 [1] 102.65 97.59 0.00 0.00 103.88 98.19
                                                  0.00 0.00
                                                                 0.00
                                                                        0.00
[1] 100.78 106.32
                   0.00
                           0.00 103.56 95.82
                                                  0.00
                                                         0.00
                                                                 0.00
[1] 100.60 94.61 0.00 0.00 100.58 96.51
                                                  0.00
                                                         9.85
 [1] 0.00 1.02 1.03 2.99 3.01 4.01 4.02 11.01 18.01 22.01
[1] 0.00 1.02 1.03 8.00 8.01 9.00 9.01 15.00 22.01
[1] 0.00 1.02 1.03 16.00 16.01 17.00 17.01 22.01
 [1] 70 66 61 55 31 31 29 26 24 22 21 19 17 14 14 13 11 11 10 9 8 8 8
 [1] 70 65 59 56 54 50 47 46 46 40 23 22 22 21 18 17 17 13 13 13 11 11 11
 [1] 70 65 59 55 53 51 48 46 46 46 44 41 40 40 40 39 38 36 33 28 24 23 19
 [1] 0 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22
$yt2
 [1] 0 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22
  \begin{bmatrix} 1 \end{bmatrix} \quad 0 \quad 1 \quad 2 \quad 3 \quad 4 \quad 5 \quad 6 \quad 7 \quad 8 \quad 9 \ 10 \ 11 \ 12 \ 13 \ 14 \ 15 \ 16 \ 17 \ 18 \ 19 \ 20 \ 21 \ 22
```

4.2 Caveats of the ${\cal R}$ Implementation

1. \mathcal{R} objects of reference classes are references to objects!

Each \mathcal{R} -object created with object <- GUTS\$new() is a reference to a C++-object. Therefore, copying the \mathcal{R} -object would not result in copying the C++-object, but in duplicating the reference:

```
R: x <- new("Rcpp_GUTS")  # a new Rcpp_GUTS object
R: x$setTimeGridPoints(M = 10000)  # set some value
R: y <- x  # copy x to y
R: y$setTimeGridPoints(M = 500)  # set some value to y
R: x$M  # access the value on x

[1] 500
```

Because the reference was copied, we now have access to one and the same C++-object via two \mathcal{R} -objects. Hence, any change on one such reference will change the underlying C++-object and will appear in all other references!

A method to copy an entire GUTS-C++-object is currently not implemented.

2. Serialisation is currently not available.

5 Usage of the R Package

Using **GUTS** may require the user to install the latest version of \mathcal{R} . Refer to section "preliminaries" (page 1) to check, which version was used for the creation of this manual. In addition to the package **GUTS**, users must install the package Rcpp. A convenient way to install **GUTS** in \mathcal{R} is provided through the \mathcal{R} -command:

```
install.packages( 'GUTS', dependencies=TRUE )
```

For the integration of **GUTS** into an MCMC application, users may be required to additionally install MHadaptive (Chivers, 2012). Professional plotting control can be achieved using the functions in package ggplot2 (Wickham, 2009). To plot multiple figures on one page users should install the package grid.

5.1 Basic Usage

The basic usage of **GUTS** is as follows:

```
R: library("GUTS")
R: go1 <- new("Rcpp_GUTS")
R: go1
GUTS object with the following attributes:
Vector of concentrations (C, O elements)
Vector of concentration time points (Ct, O elements)
Vector of survivors (y, 0 elements)
Vector of survivor time points (yt, 0 elements)
Parameters (par, 0 elements)
Time grid points (M): 10000
Distribution (dist) : lognormal
                     : 10000
Sample length (N)
Sample vector (z, 0 elements)
Vector of damages (D, 0 elements)
Vector of survival probabilities (S, O elements)
Loglikelihood (LL): NaN
Messages/warnings:
[1] "C not set up"
                                               "Ct not set up"
[3] "y not set up"
                                               "yt not set up"
[5] "par not set up" "z not available"
[7] "Survival probabilities not calculated" "Loglikelihood not calculated"
R: go1$LL
[1] NaN
R: logLik(go1)
'log Lik.' NA (df=NA)
```

In this example, a "factory-fresh" **GUTS** object (go1) is created. For the creation we can use the function new("Rcpp_GUTS") or the method new(), where the method must be invoked with the *creator*

object GUTS. The object contains default or non-sense values, and the loglikelihood (accessed either by the field LL or using the function logLik()) delivers NA.

Having a **GUTS** object, users may fill it with more sensible data. One example can be found in the manual page of **GUTS** in \mathcal{R} , and is reproduced here. We use data from the data set diazinon. The data set is included in the package (see section 4.1).

```
R: data("diazinon")
R: go1$setConcentrations(C = diazinon$C1, Ct = diazinon$Ct1)
R: go1$setSurvivors(y = diazinon$y1, yt = diazinon$yt1)
R: go1\$setParameters(par = c(0.05084761, 0.12641525, 1.61840054, 19.09911, 6.495246))
R: go1$setTimeGridPoints(M = 10000)
R: go1$setDistribution(dist = "lognormal")
R: go1$setSampleLength(N = 10000)
R: go1$calcLoglikelihood()
R: go1$LL
[1] -183.2561
R: logLik(go1)
'log Lik.' -183.5762 (df=23)
R: go1
GUTS object with the following attributes:
 ._____
Vector of concentrations (C, 10 elements):
    102.65, 97.59, 0, 0, 103.88, 98.19, 0, 0, 0
Vector of concentration time points (Ct, 10 elements):
   0, 1.02, 1.03, 2.99, 3.01, 4.01, 4.02, 11.01, 18.01, 22.01
Vector of survivors (y, 23 elements):
   70, 66, 61, 55, 31, 31, 29, 26, 24, 22, 21, 19, 17, 14, 14, 13, 11, 11, 10, 9, 8, 8, 8
Vector of survivor time points (yt, 23 elements):
    0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22
Parameters (par. 5 elements):
   0.0508476, 0.1264152, 1.6184005, 19.09911, 6.495246
Time grid points (M): 10000
Distribution (dist) : lognormal
Sample length (N)
                    : 10000
Sample vector (z, 10000 elements):
  Min=4.942491, Max=57.94564, Mean=19.02988, Sigma=6.372639
Vector of damages (D, 10000 elements):
 Min=0, Max=20.45299, Mean=8.556884, Sigma=5.016242
Vector of survival probabilities (S, 23 elements):
    1, 0.9276622, 0.8375214, 0.7954438, 0.4589534, 0.3407472, 0.3235829, 0.3075407, 0.2922939,
   0.277803, 0.2640305, 0.2509408, 0.2385, 0.226676, 0.2154382, 0.2047575, 0.1946064, 0.1849585,
    0.1757889, 0.1670739, 0.1587909, 0.1509186, 0.1434366
Loglikelihood (LL): -183.5762
Messages/warnings: none
```

Compared to the "factory-fresh" **GUTS** object above, all errors disappeared, because the object was set up properly. In addition, the invocation of the calculation of the loglikelihood (method calcLoglikelihood()) also caused the computation of a new sample, a vector of damages, a vector of survival probabilities, and the loglikelihood. The call to the S3 function logLik() re-computed a sample, and hence the resulting likelihood is slightly different from the previous one. Note, that each time the user either calls the method calcLoglikelihood() or the function logLik(), the object's loglikelihood is re-calculated. The recommended way of calculating and accessing the loglikelihood is:

```
R: go1$calcLoglikelihood()  # calculate it
R: go1$LL  # get it

[1] -183.3443
```

5.2 Using GUTS in MCMC Routines

As a real-world example we perform a Bayesian parameter inference (with uniform priors) using the survival data of *Gammarus pulex* exposed to *Diazinon* Ashauer, Hintermeister, Caravatti, Kretschmann and Escher (2010). The data is contained in the data set diazinon, which is contained in the **GUTS** package. In these experiments three different exposure patterns (treatments) have been applied. Since we want to use all the data for the parameter inference, we represent the three exposure patterns by three instances of the GUTS class and use the sum of the three loglikelihoods. For the computation of the MCMC we use the package adaptMCMC by Andreas Scheidegger (andreas.scheidegger@eawag.ch).

First, we set up three different **GUTS** objects. Note that all data is contained in the data set diazinon.

```
R: # new objects
R: tmtA <- new("Rcpp_GUTS")</pre>
R: tmtB <- new("Rcpp_GUTS")</pre>
R: tmtC <- new("Rcpp_GUTS")</pre>
R: # concentrations and concentration time points
R: tmtA$setConcentrations(C = diazinon$C1, Ct = diazinon$Ct1)
R: tmtB$setConcentrations(C = diazinon$C2, Ct = diazinon$Ct2)
R: tmtC$setConcentrations(C = diazinon$C3, Ct = diazinon$Ct3)
R: # survivors and survivor time points
R: tmtA\$setSurvivors(y = diazinon\$y1, yt = diazinon\$yt1)
R: tmtB\$setSurvivors(y = diazinon\$y2, yt = diazinon\$yt2)
R: tmtC$setSurvivors(y = diazinon$y3, yt = diazinon$yt3)
R: # distribution
R: tmtA$setDistribution(dist = "lognormal")
R: tmtB$setDistribution(dist = "lognormal")
R: tmtC$setDistribution(dist = "lognormal")
R: # numercial exactness
R: tmtA$setTimeGridPoints(M = 10000)
R: tmtB$setTimeGridPoints(M = 10000)
R: tmtC$setTimeGridPoints(M = 10000)
R: tmtA\$setSampleLength(N = 10000)
R: tmtB$setSampleLength(N = 10000)
R: tmtC$setSampleLength(N = 10000)
```

We now define a set of starting parameters, and create a function that updates the parameters on either of the three objects and delivers the loglikelihood:

```
R: par.start <- c(0.1, 0.3, 1, 1, 1)
R: loglikeli <- function(par) {
   if ( any(par < 0) ) {
      # Parameters must not be negative
      return(-Inf)
   } else {
      # The loglikelihood of the 3 treatments
      # is just the sum of the individual ones</pre>
```

```
tmtA$setParameters(par)
  tmtB$setParameters(par)
  tmtC$setParameters(par)
  tmtA$calcLoglikelihood()
  tmtB$calcLoglikelihood()
  tmtC$calcLoglikelihood()
  out <- tmtA$LL + tmtB$LL + tmtC$LL
  return(out)
}</pre>
```

Note that if a particular error occurs (e.g., a sample mean smaller than 0), then this functions returns NA. Next, we set the jump standard deviations for the MCMC chain:

```
R: par.names <- c("h.b", "k.r", "k.k", "mean", "sd")
R: sigma <- diag(par.start/10)^2
```

Calculate the adaptive Markov chain using the package adaptMCMC:

```
R: library("adaptMCMC")
R: res.mcmc <- MCMC(p = loglikeli, n = 50000, init = par.start, scale = sigma,
adapt = TRUE, acc.rate = 0.4)
```

Our result can be inspected in various ways, one is to look at the structure:

```
R: str(res.mcmc)
List of 7
 $ samples
                     : num [1:50000, 1:5] 0.1 0.0916 0.0916 0.0916 0.0874 ...
                     : num [1:50000] -7471 -2876 -2876 -2876 -2560 ...
: num [1:5, 1:5] 2.72e-05 8.35e-06 8.98e-04 1.55e-03 -1.04e-03 ...
 $ log.p
 $ cov.jump
                     : num 50000
 $ n.sample
 $ acceptance.rate : num 0.417
 $ adaption
                      : logi TRUE
 $ sampling.parameters:List of 3
  ..$ sample.density:function (par)
  ....- attr(*, "srcref")=Class 'srcref' atomic [1:8] 470 14 486 1 14 1 3 19
  ..... attr(*, "srcfile")=Classes 'srcfilealias', 'srcfile' <environment: 0x1060a6b40>
                   : num 0.4
  ..$ acc.rate
                    : num 0.5
  ..$ gamma
```

5.3 Plotting the Outcomes of the MCMC Inference

Excellent plotting facilities are available through the routines in the package ggplot2 (Wickham, 2009). ggplot2 offers great features for generating object-oriented plots. To plot the trace of the parameter mean one may use the code below. Note that due to file size issues we reduce data to the each 50th iteration. The result is displayed in figure 1):

```
R: library("ggplot2")
R: # prepare a data frame, and select select each 10th observation
R: k <- nrow(res.mcmc$samples)</pre>
R: i \leftarrow seq(1, k, 50)
R: df <- data.frame(i, res.mcmc$samples[i, ])</pre>
R: colnames(df) <- c("iter", par.names)
R: str(df)
'data.frame': 1000 obs. of 6 variables:
 $ iter: num 1 51 101 151 201 251 301 351 401 451 ...
 $ h.b : num 0.1 0.0621 0.0613 0.0516 0.0483 ...
 $ k.r : num  0.3 0.00718 0.00593 0.00616 0.00664 ...
 $ k.k : num 1 1.33 1.34 1.36 1.35 ...
 $ mean: num 1 1.57 1.58 1.58 1.58 ...
 $ sd : num 1 1.31 1.29 1.35 1.38 ...
R: # create the plot object
R: ggp.mean <- ggplot(df, aes(x=iter, y=mean)) +
    geom_line() +
    xlab('iteration') +
    ylab('Value of mean')
```

```
R: ggp.mean
```

In the following we create two plots using ggplot2, the trace plot of the five parameters (figure 2) and the density plots (figure 3). Both statistics should appear on one page each, hence we create a little helper function to set up the page (requires the package grid!). Note, that if one wants only one plot appear on one page (pdf), this helper function and the application of multiplot() is not necessary.

```
R: multiplot <- function(..., plotlist = NULL, cols) {
    require(grid)
    plots <- c(list(...), plotlist)
    numPlots = length(plots)
    plotCols = cols
    plotRows = ceiling(numPlots/plotCols)
    grid.newpage()
    pushViewport(viewport(layout = grid.layout(plotRows, plotCols)))
    vplayout <- function(x, y) {
        viewport(layout.pos.row = x, layout.pos.col = y)
    }
    for (i in 1:numPlots) {
        curRow = ceiling(i/plotCols)
        curCol = (i - 1)%plotCols + 1
        print(plots[[i]], vp = vplayout(curRow, curCol))
    }
}</pre>
```

We now prepare the data frame holding the parameters as well as a column signing the loop in the MCMC. ggplot2 creates objects of plots. Printing these objects results in the final plot. This differs from the standard plot routines in \mathcal{R} , where in most of the cases the plot function plots itself. Note that due to plot size issues we reduce data to each 50th iteration.

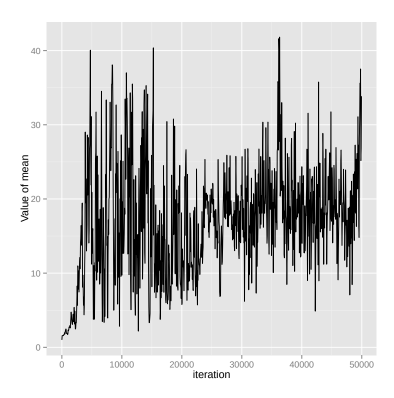


Figure 1: Plot of the trace of the parameter mean using package ggplot2.

```
R: # prepare a data frame
R: k <- nrow(res.mcmc$samples)
R: i \leftarrow seq(1, k, 50)
R: plot.data <- data.frame(i, res.mcmc$samples[i, ])</pre>
R: colnames(plot.data) <- c("iter", par.names)
R: str(plot.data)
'data.frame': 1000 obs. of 6 variables:
 $ iter: num 1 51 101 151 201 251 301 351 401 451 ...
 $ h.b : num 0.1 0.0621 0.0613 0.0516 0.0483 ...
 $ k.r : num 0.3 0.00718 0.00593 0.00616 0.00664 ...
 k.k : num 1 1.33 1.34 1.36 1.35 ...
 $ mean: num 1 1.57 1.58 1.58 1.58 ...
 $ sd : num 1 1.31 1.29 1.35 1.38 ...
R: # five plots for five parameters
R: plot.h.b <- ggplot(plot.data, aes(x=iter, y=h.b)) +
    geom_line() +
    xlab('iter') +
    ylab('Value of h.b')
R: plot.k.r <- ggplot(plot.data, aes(x=iter, y=k.r)) +
     geom_line() +
    xlab('iter') +
    ylab('Value of k.r')
R: plot.k.k \leftarrow ggplot(plot.data, aes(x=iter, y=k.k)) +
    geom_line() +
     xlab('iter') +
    ylab('Value of k.k')
R: plot.mean <- ggplot(plot.data, aes(x=iter, y=mean)) +
```

```
geom_line() +
   xlab('iter') +
   ylab('Value of mean')
R: plot.sd <- ggplot(plot.data, aes(x=iter, y=sd)) +
   geom_line() +
   xlab('iter') +
   ylab('Value of sd')</pre>
```

Now we use the multiplot() function defined above, and plot each ggplot object in the appropriate position.

```
R: multiplot(plot.h.b, plot.k.r, plot.k.k, plot.mean, plot.sd, cols=2)
```

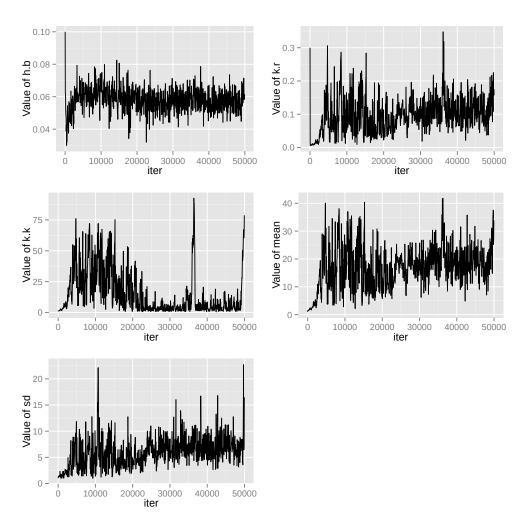


Figure 2: Plot of the trace of the parameters using package ggplot2.

The traces shown in figure 2 suffer from both, burn-in and adaption. Therefore, we cut off the first

20,000 sample points and then produce the density plots.

```
R: # create plot data frame
R: dat <- res.mcmc$samples
R: i <- 20001
R: k <- nrow(dat)
R: plot.data <- data.frame( i:k, dat[i:k, ] )</pre>
R: colnames(plot.data) <- c("iter", par.names)
R: str(plot.data)
'data.frame': 30000 obs. of 6 variables:
 $ iter: int 20001 20002 20003 20004 20005 20006 20007 20008 20009 20010 ...
 $ h.b : num 0.0485 0.0485 0.0528 0.054 0.054 ...
 $ k.r : num  0.0223 0.0223 0.0232 0.0258 0.0258 ...
 $ k.k : num 2.41 2.41 2.13 2.38 2.38 ...
 $ mean: num 5.77 5.77 5.9 6.07 6.07 ...
 $ sd : num 4.86 4.86 4.23 3.69 3.69 ...
R: # five plots for five parameters
R: plot.dens.h.b <- ggplot(plot.data) +
     geom\_density(aes(x = h.b)) +
    xlab('h.b') +
     ylab('Density')
R: plot.dens.k.r <- ggplot(plot.data) +
    geom_density(aes(x = k.r)) +
     xlab('k.r') +
    ylab('Density')
R: plot.dens.k.k <- ggplot(plot.data) +</pre>
     geom\_density(aes(x = k.k)) +
     xlab('k.k') +
     ylab('Density')
R: plot.dens.mean <- ggplot(plot.data) +
    geom_density(aes(x = mean)) +
     xlab('mean') +
    ylab('Density')
R: plot.dens.sd <- ggplot(plot.data) +
    geom\_density(aes(x = sd)) +
     xlab('sd') +
    ylab('Density')
```

```
R: multiplot(plot.dens.h.b, plot.dens.k.r, plot.dens.k.k, plot.dens.mean, plot.dens.sd, cols=2)
```

Next, we address the best fit. The result of the MCMC was saved in object res.mcmc. This object is a list with various attributes. Two attributes are of particular interest here: The vector of loglikelihoods saved during the MCMC (log.p), and the "vector" of parameters (saved as a matrix samples). Each of these two objects has a length equal to the iterations' number (here: n = 50,000). The "maximum probable parameters" can be found at the position of the maximum loglikelihood:

```
R: best.fit.pos <- which.max(res.mcmc$log.p)
R: best.fit.pars <- res.mcmc$samples[best.fit.pos, ]
```

We set the parameters of each of the three experimental **GUTS** objects the parameters of the best fit, and calculate the survival probabilities accordingly:

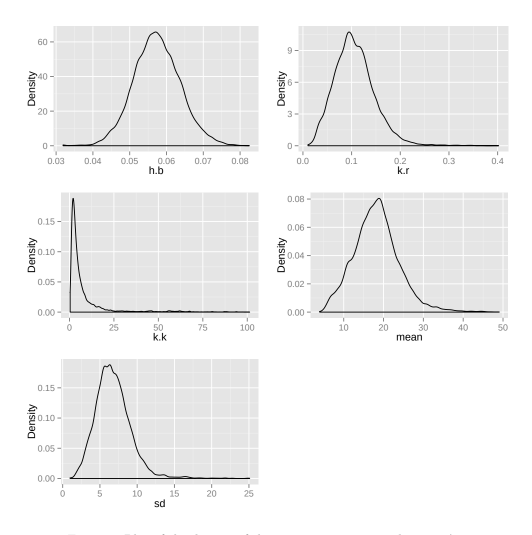


Figure 3: Plot of the density of the parameters using package ggplot2.

```
R: tmtA$setParameters(best.fit.pars)
R: tmtB$setParameters(best.fit.pars)
R: tmtC$setParameters(best.fit.pars)
R: tmtA$calcSurvivalProbabilities()
R: tmtB$calcSurvivalProbabilities()
R: tmtC$calcSurvivalProbabilities()
```

Now, we create three plots, one for each experiment. In each plot, we want to display the measured number of deaths $(y_i - y_{i-1})$ compared to the predicted number of deaths $(y_0(S_i - S_{i-1}))$.

Figure 4 shows the results for the three experiments.

```
R: # prepare data frames
R: plot.data.A <- data.frame(
"var" = rep( c("measured", "predicted"), each=length(tmtA$y) ),
```

```
"yt" = rep(tmtA$yt, 2),
     "y"
          = c(
              -diff(tmtA$y), tmtA$y[length(tmtA$y)],
              c(-diff(tmtA$S), tmtA$S[length(tmtA$S)]) * max(tmtA$y)
R: plot.data.B <- data.frame(
     "var" = rep( c("measured", "predicted"), each=length(tmtB$y) ),
     "yt" = rep(tmtB$yt, 2),
          = c(
            -diff(tmtB$y), tmtB$y[length(tmtB$y)],
             c(-diff(tmtB$S), tmtB$S[length(tmtB$S)]) * max(tmtB$y)
R: plot.data.C <- data.frame(
     "var" = rep( c("measured", "predicted"), each=length(tmtC$y) ),
     "yt" = rep(tmtC\$yt, 2),
          = c(
             -diff(tmtC$y), tmtC$y[length(tmtC$y)],
             c(-diff(tmtC\$S), tmtC\$S[length(tmtC\$S)]) * max(tmtC\$y)
R: # generate 3 plots using ggplot
R: plot.bf.A <- ggplot(plot.data.A, aes(x=yt, y=y, group=var, shape=var, colour=var)) + \frac{1}{2}
     geom_point() +
     xlab('survivor time points in experiment A') +
     ylab('deaths')
R: plot.bf.B <- ggplot(plot.data.B, aes(x=yt, y=y, group=var, shape=var, colour=var)) +
     geom_point() +
     xlab('survivor time points in experiment B') +
     ylab('deaths')
R: \ plot.bf. C \leftarrow ggplot(plot.data.C, \ aes(x=yt, \ y=y, \ group=var, \ shape=var, \ colour=var)) \ + \\
     geom_point() +
     xlab('survivor time points in experiment C') +
     ylab('deaths')
```

Again, we use the function multiplot() from above to merge all plots.

```
R: multiplot(plot.bf.A, plot.bf.B, plot.bf.C, cols=1)
```

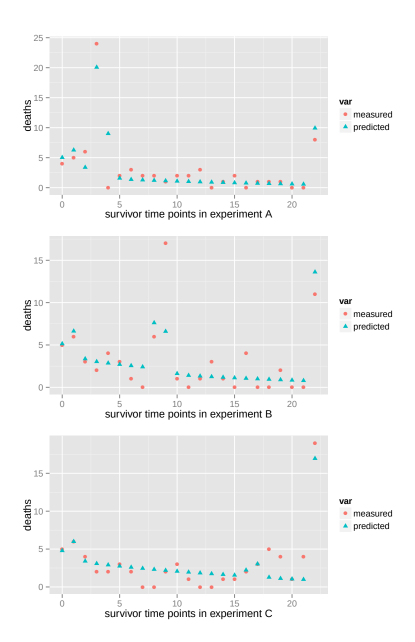


Figure 4: Plot of the best fit using package ggplot2.

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