Bayesian survival analysis with BUGS

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Abstract

Survival analysis is one of the most important fields of statistics in medicine and biological sciences. In addition, the computational advances in the last decades have favoured the use of Bayesian methods in this context, providing a flexible and powerful alternative to the traditional frequentist approach. The objective of this paper is to summarise some of the most popular Bayesian survival models, such as accelerated failure time, proportional hazards, mixture cure, competing risks, multi-state, frailty, and joint models of longitudinal and survival data. Moreover, an implementation of each presented model is provided using a BUGS syntax that can be run with JAGS from the R programming language. Reference to other Bayesian R-packages are also discussed.

Key Words: Bayesian inference, JAGS, R-packages, time-to-event analysis.

1 Introduction

Survival analysis, sometimes referred to as *failure-time analysis*, is one of the most important fields of statistics, mainly in medicine and biological sciences (Kleinbaum and Klein, 2012; Collett, 2015). Survival times are data that measure follow-up time from a defined starting point to the occurrence of a given event of interest or endpoint, for instance, onset of disease, cure, death, etc. (Kalbfleisch and Prentice, 2002).

Continuous survival times are defined as non-negative random variables, T, whose probabilistic behaviour can be equivalently described by its hazard function, survival function, or density function. The hazard function of T at time t represents the instantaneous rate of the event occurrence for the population group that is still at risk at time t. It is defined as

$$h(t \mid \boldsymbol{\theta}) = \lim_{\Delta t \to 0} \frac{P(t \le T < t + \Delta t \mid T \ge t, \boldsymbol{\theta})}{\Delta t} = \frac{f(t \mid \boldsymbol{\theta})}{S(t \mid \boldsymbol{\theta})}, \quad t > 0,$$
(1)

where θ is a set of unknown quantities, $f(t \mid \theta)$ is the density function of T given θ , and $S(t \mid \theta) = P(T > t \mid \theta)$ is the survival function of T given θ . Note that $h(t \mid \theta) \ge 0$ and in general $\int h(t \mid \theta) dt = \infty$. The hazard function is defined in terms of a conditional probability but is not a probability. It could be interpreted as the approximate instantaneous probability of having the event given that the target individual has not yet experienced it at time t.

Usual relationships between $h(t \mid \boldsymbol{\theta})$, $f(t \mid \boldsymbol{\theta})$, and $S(t \mid \boldsymbol{\theta})$ derived from (1) that will be useful throughout the article include

$$H(t \mid \boldsymbol{\theta}) = \int_0^t h(u \mid \boldsymbol{\theta}) du, \quad t > 0,$$
 (2)

$$S(t \mid \boldsymbol{\theta}) = \exp \left\{ -H(t \mid \boldsymbol{\theta}) \right\}, \quad t > 0, \tag{3}$$

$$f(t \mid \boldsymbol{\theta}) = h(t \mid \boldsymbol{\theta}) \exp \left\{ -H(t \mid \boldsymbol{\theta}) \right\}, \quad t > 0, \tag{4}$$

where $H(t \mid \boldsymbol{\theta})$ denotes the *cumulative hazard function* of T.

Standard statistical techniques cannot usually be applied to survival data because in most cases they are not fully recorded, mainly due to censoring and/or truncation schemes (Klein et al., 2013). Also, normal distributions are usually inappropriate for analyzing survival data at their original scale, since they are positive and often asymmetrical. Distributions such as the Weibull, gamma or log-normal are the ones that now occupy the leading role.

From a Bayesian perspective, censoring mechanisms do not affect the prior distribution but their do affect the likelihood function, which factorises in general as the product of the likelihood function corresponding to each individual i in the sample as $L(\theta) = \prod_{i=1}^{n} L_i(\theta)$. In general terms, censored observations can be classified as *right-censored*, *left-censored* and *interval-censored*. See Klein and Moeschberger (2003) for technical details, interpretations and examples. The contribution of the survival observation t of individual t to the likelihood function, $L_i(\theta)$, depends on the type of censoring as follows:

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Exact survival time: $f_i(t \mid \boldsymbol{\theta})$ Right-censored observation: $S_i(C_r \mid \boldsymbol{\theta})$ Left-censored observation: $1 - S_i(C_l \mid \boldsymbol{\theta})$ Interval-censored observation: $S_i(C_l \mid \boldsymbol{\theta}) - S_i(C_r \mid \boldsymbol{\theta})$

Right censoring here is type I, where the end of study period C_r is known and prefixed before it begins. In the case of left-censored observations, the survival time is known to be below a certain censoring time C_l . When survival times are interval-censored in $[C_l, C_r]$, the survival time is somewhere in this interval.

Due to these peculiar characteristics and their extreme relevance to different scientific subjects, survival models/methods have been extensively developed over the past 50 years, which includes many R-packages*. Hence, in order to illustrate for which situations some survival models are more appropriate than others, we summarise the main models for survival data, such as accelerated failure time (AFT), proportional hazards (PH), mixture cure, competing risks, multi-state, frailty, and joint models of longitudinal and survival data. In particular, the inferential process is performed from a Bayesian perspective, which is an appealing alternative to the traditional frequentist approach, since the Bayesian probability conception allows to measure the uncertainty associated with parameters, models, hypotheses, missing data in probabilistic terms (Ibrahim et al., 2001). So, based on the Bayesian paradigm, we exemplify each survival modelling using Markov chain Monte Carlo (MCMC) methods (Gelman et al., 2013) implemented in BUGS syntax (Gilks et al., 1994) with the support of JAGS (Plummer, 2003) and the rjags (Plummer, 2018) package (version 4-10) for the R language (version 4.0.2) (R Core Team, 2020).

The set of models introduced in this paper is quite comprehensive as it covers the main models in survival analysis. However, there are a number of R-packages to fit different types of Bayesian survival models which can be used in specific contexts. For example, if one is interested in AFT modelling, the bayesSurv package provides implementations of mixed-effects AFT models with various censored data specifications (Komárek, 2018); while the DPpackage package fits nonparametric and semi-parametric models with different prior processes (Jara et al., 2018); and spBayesSurv package includes AFT and PH models (among others) for spatial/non-spatial survival data (Zhou and Hanson, 2018). If a PH specification is preferable, the BMA package implements a Bayesian model averaging approach for Cox PH models (Raftery et al., 2018); the dynsurv package provides time-varying coefficient models for interval-censored and right-censored survival data (Wang et al., 2017); ICBayes package offers semi-parametric regression survival models to interval-censored time-to-event data (Pan et al., 2017); and the spatsury package fits parametric PH spatial survival models (Taylor et al., 2018). For multi-state problems, we highlight the CFC package that implements a cause-specific framework for competing-risk analysis (Sharabiani and Mahani, 2019); and SemiCompRisks package that provides a broader specification using of independent/clustered semi-competing risks models (Lee et al., 2019). When a joint approach of longitudinal and survival data is required, the most popular package is the JMbayes one, where the longitudinal process is modelled through a linear mixed framework, a Cox PH model is assumed for the survival process, and various association structures between both processes are provided (Rizopoulos, 2019). Other generic packages for Bayesian inference such as, for example, Bayesian (Umlauf et al., 2019), RStan (Carpenter et al., 2017; Stan Development Team, 2020) or INLA (Rue et al., 2009) can be used to fit different survival models.

The paper is organised as follows. Sections 2–7 describe the following survival models: AFT, PH, mixture cure, competing risks, multi-state, frailty, and joint models of longitudinal and survival data. In particular, in each one of these sections, we briefly introduce notation and basic concepts of the models under discussion. Then, the survival dataset used (available in an R-package) is described jointly with the BUGS code implementation. Finally, posterior summaries, and graphs of quantities of interest derived from the posterior distribution are provided. To conclude, Section 8 ends with an overview of the Bayesian survival models introduced and implemented in this paper, motivating the use and adaptation of the codes provided. Theoretical and methodological aspects of survival models and Bayesian inference are not discussed in depth in this paper, so we recommend that readers unfamiliar with these topics review specific references, such as Klein et al. (2013) and Gelman et al. (2013).

2 Survival regression models

Regression models focus on the association between time-to-event random variables and covariates (explanatory/predictor variables or risk factors). They allow the comparison of survival times between groups both with regard to the general behaviour of the individuals in each group and prediction for new members of them. The most popular approaches to survival regression models are the accelerated failure time (AFT) and the proportional hazards (PH) models, also known as Cox models (Cox, 1972).

2.1 Accelerated failure time models

AFT models are the survival counterpart of linear models. Survival time T in logarithmic scale is expressed in terms of a linear combination of covariates x with regression coefficients β and a measurement error ξ as follows:

$$\log\left(T\right) = \boldsymbol{x}^{\top}\boldsymbol{\beta} + \sigma\,\epsilon,\tag{5}$$

^{*}https://cran.r-project.org/web/views/Survival.html

where σ is a scale parameter and ϵ an error term usually expressed via a normal, logistic or a standard Gumbel probabilistic distribution. The particular case of a standard Gumbel distribution for ϵ implies a conditional (on β and σ) Weibull survival model for T with shape $\alpha = 1/\sigma$ and scale $\lambda(\beta, \sigma) = \exp\{-x^{\top}\beta/\sigma\}$ parameters (Christensen et al., 2011).

2.1.1 *larynx* dataset

We consider a larynx cancer dataset, referred to as larynx. It is available from the KMsurv package (Klein et al., 2012):

```
R> library("KMsurv")
R> data("larynx")
R> str(larynx)
'data.frame': 90 obs. of 5 variables:
$ stage : int 1 1 1 1 1 1 1 1 1 1 ...
$ time : num 0.6 1.3 2.4 2.5 3.2 3.2 3.3 3.3 3.5 3.5 ...
$ age : int 77 53 45 57 58 51 76 63 43 60 ...
$ diagyr: int 76 71 71 78 74 77 74 77 71 73 ...
$ delta : int 1 1 1 0 1 0 1 0 1 1 ...
```

This dataset provides observations of 90 male larynx-cancer patients, diagnosed and treated in the period 1970–1978 (Kardaun, 1983). The following variables were observed for each patient:

- stage: disease stage (1-4).
- time: time (in months) from first treatment until death, or end of study.
- age: age (in years) at diagnosis of larynx cancer.
- diagyr: year of diagnosis of larynx cancer.
- delta: death indicator (1: if patient died; 0: otherwise).

2.1.2 Model specification

Survival times are analysed through the following accelerated failure time (AFT) model:

$$\log(T) = \beta_1 + \beta_2 I_{(\text{stage}=2)} + \beta_3 I_{(\text{stage}=3)} + \beta_4 I_{(\text{stage}=4)} + \beta_5 \text{age} + \beta_6 \text{diagyr} + \sigma \epsilon, \tag{6}$$

where T represents death time for each individual; β_1 is an intercept; $I_{(stage=\cdot)}$ is an indicator variable for stage=2, 3, 4 with regression coefficients β_2 , β_3 , and β_4 , respectively (stage=1 is considered as the reference category); and β_5 and β_6 are regression coefficients for age and diagyr covariates, respectively. The errors ϵ 's are i.i.d. random variables which follow a standard Gumbel distribution and σ is a scale parameter.

The Bayesian model is completed with the specification of a prior distribution for their corresponding parameters. A non-informative prior independent default scenario is considered. The marginal prior distribution for each regression coefficient β_k , $k=1,\ldots,6$, is elicited as a normal distribution centered at zero and with a small precision, N(0,0.001). A uniform distribution, Un(0,10), is selected as the marginal prior distribution for α (i.e., the Weibull shape parameter). However, the use of another continuous distributions, e.g., Gamma(1,0.001) is also common under a non-informative prior framework Sahu et al. (1997).

2.1.3 Model implementation

Censoring is handled in JAGS using the distribution dinterval which implies the modification of the default time variable and the creation of two new ones, cens and is.censored. For a generic database which deals with observations of different censoring types (i.e., right-censored, left-censored, and/or interval-censored) the specification of these variables should be managed as follows:

- time: survival time is time if the event was observed (delta=1); otherwise, NA.
- cens[i,]: two column matrix with n rows (with n equal to the number of individuals). cens[i,] aggregates for each observation i a vector of two cut points. According to the censoring type cens must be specified as:
 - Uncensored: cens[i,1] = 0 (or any arbitrary value is allowed).
 - Right-censored: cens[i,1] = censoring time.
 - Left-censored: cens[i,2] = censoring time.
 - Interval-censored: cens[i,] = (cens.low, cens.up) (cens.low corresponds with the lower limit and cens.up with the upper limit of the observed interval).
- is.censored: a help ordinal variable to indicate the censoring status. According to the censoring type cens must be specified as:

```
Uncensored: is.censored = 0
Right-censored: is.censored = 1
Left-censored: is.censored = 0
Interval-censored: is.censored = 1
```

Thus, for the larynx dataset which contains uncensored and right-censored observations the following code is dedicated to creating or transforming these variables:

Next, we have scaled the age and diagyr, and encoded the stage as a factor. Then, we have created a design matrix X of these covariates:

```
R> larynx$age <- as.numeric(scale(larynx$age))
R> larynx$diagyr <- as.numeric(scale(larynx$diagyr))
R> larynx$stage <- as.factor(larynx$stage)
R> X <- model.matrix(~ stage + age + diagyr, data = larynx)</pre>
```

Listing 1 shows a generic implementation of an AFT model in BUGS syntax using *larynx* data.

Listing 1: AFT model in BUGS syntax (file named as **AFT.txt**).

```
model{
    for(i in 1:n){
        # Survival and censoring times
        is.censored[i] ~ dinterval(time[i],cens[i,1])
        time[i] ~ dweib(alpha,lambda[i])
        lambda[i] <- exp(-mu[i] * alpha)
        mu[i] <- inprod(beta[],X[i,])
    }

# Prior distributions
for(l in 1:Nbetas){ beta[1] ~ dnorm(0,0.001) }
alpha ~ dunif(0,10)
}</pre>
```

Once the BUGS syntax and its corresponding variables has been created, JAGS requires specifying some elements to run the MCMC simulation:

- d. jags: a list with all the elements/data specified in the model.
- i. jags: a function that returns a list of initial random values for each model parameters.
- p.jags: a character vector with the parameter names to be monitored/saved.

These elements are defined for our AFT model as follows:

```
R> d.jags <- list(n = nrow(larynx), time = larynx$time, cens = cens, X = X,
+ is.censored = is.censored, Nbetas = ncol(X))
R> i.jags <- function() { list(beta = rnorm(ncol(X)), alpha = runif(1)) }
R> p.jags <- c("beta", "alpha")</pre>
```

Then, MCMC algorithm is run in three steps. Firstly, the JAGS model is compiled by means of the jags.model function available from the rjags package:

```
R> library("rjags")
R> m1 <- jags.model(data = d.jags, file = "AFT.txt", inits = i.jags, n.chains = 3)</pre>
```

The n.chains argument indicates the number of Markov chains selected. Secondly, a burn-in period is considered (here the first 1000 simulations) using the update function:

```
R> update(m1, 1000)
```

Thirdly, the model is run using coda.samples function for a specific number of iterations to monitor (here n.iter=50000) as well as a specific thinning value (here thin=10) in order to reduce autocorrelation in the saved samples:

```
R> res <- coda.samples(m1, variable.names = p.jags, n.iter = 50000, thin = 10)
```

A posterior distributions summary can be obtained through the summary function:

```
R> summary(res)
Iterations = 2010:52000
Thinning interval = 10
Number of chains = 3
Sample size per chain = 5000
1. Empirical mean and standard deviation for each variable,
  plus standard error of the mean:
           Mean
                  SD Naive SE Time-series SE
        1.03426 0.1353 0.001105 0.001365
alpha
beta[1] 2.55344 0.2967 0.002422
                                     0.003868
beta[2] -0.12702 0.4769 0.003894
                                    0.004812
beta[3] -0.64710 0.3628 0.002962
                                    0.004125
beta[4] -1.66226 0.4431 0.003618
                                    0.004878
beta[5] -0.20953 0.1552 0.001267
                                    0.001323
beta[6] 0.07054 0.1622 0.001324
                                    0.001523
2. Quantiles for each variable:
         2.5% 25% 50%
                                   75%
                                           97.5%
alpha
        0.7827 0.93886 1.02986 1.1216 1.31162
beta[1] 2.0427 2.34542 2.52919 2.7382 3.18745
beta[2] -1.0342 -0.44200 -0.13807 0.1745 0.86159
beta[3] -1.3800 -0.88079 -0.63669 -0.4061 0.05082
beta[4] -2.5689 -1.94726 -1.65245 -1.3664 -0.82138
beta[5] -0.5279 -0.31022 -0.20582 -0.1040 0.08472
beta[6] -0.2342 -0.04002 0.06568 0.1766 0.40106
```

Markov chains must reach stationarity and, to check this condition, several convergence diagnostics can be applied. Trace plots (traceplot function) and the calculation of the Gelman-Rubin diagnostic (gelman.diag function) are used for illustrating this issue (Gelman and Rubin, 1992; Brooks and Gelman, 1998) with the coda package (Plummer et al., 2019), which is already loaded as rjags depends on it:

```
R > par(mfrow = c(2,4))
R> traceplot(res)
R> gelman.diag(res)
Potential scale reduction factors:
       Point est. Upper C.I.
alpha
             1
                       1.00
               1
                        1.01
beta[1]
               1
                       1.00
beta[2]
               1
                       1.01
beta[3]
beta[4]
               1
                       1.00
beta[5]
               1
                       1.00
beta[6]
               1
                       1.00
Multivariate psrf
```

Trace plots of the sampled values for each parameter in the chain appear overlapping one another and Gelman-Rubin values are very close to 1, which indicates that convergence has been achieved.

From the densplot function, also available from the coda package, we can draw the marginal posterior distributions (using kernel smoothing) for all the model parameters:

```
R> par(mfrow = c(2,4))
R> densplot(res, xlab = "")
```

Simulation-based Bayesian inference requires using simulated samples to summarise posterior distributions or calculate any relevant quantities of interest. So, posterior samples from the three Markov chains can be merged together using the following code:

```
R> result <- as.mcmc(do.call(rbind, res))</pre>
```

Next, the posterior samples of each parameter are extracted from the result object as follows:

```
R> alpha <- result[,1]; beta1 <- result[,2]; beta2 <- result[,3]; beta3 <- result[,4]
R> beta4 <- result[,5]; beta5 <- result[,6]; beta6 <- result[,7]</pre>
```

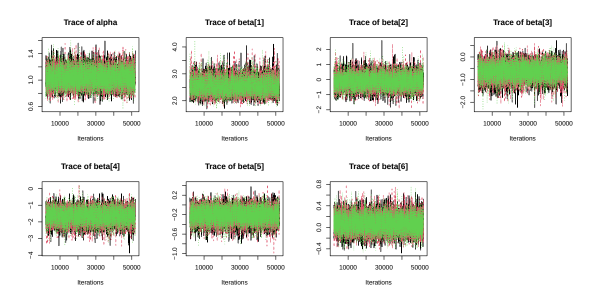


Figure 1: Trace plots for α and β 's of the AFT model (6).

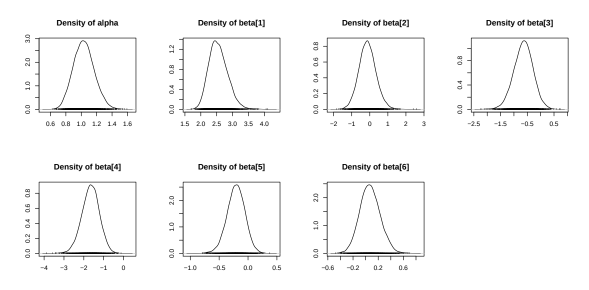


Figure 2: Density plots for α and β 's of the AFT model (6).

A relevant quantity for AFT models is the *relative median* (RM) between two individuals with covariate vectors x_1 and x_2 . This measure compares the median of the survival time between both individuals and is defined as:

$$RM(\boldsymbol{x}_1, \boldsymbol{x}_2, \boldsymbol{\beta}) = \exp\left\{ (\boldsymbol{x}_1 - \boldsymbol{x}_2)^{\top} \boldsymbol{\beta} \right\}.$$

As an illustration, we can easily summarise the posterior distribution of the RM between two men of the same age and diagyr (year of diagnosis) but one in stage=3 and the other in stage=4:

```
R> RM.s3_s4 <- exp(beta3 - beta4)
R> summary(RM.s3_s4)
Iterations = 1:15000
Thinning interval = 1
Number of chains = 1
Sample size per chain = 15000
1. Empirical mean and standard deviation for each variable,
   plus standard error of the mean:
                                    Naive SE Time-series SE
                           SD
          Mean
       3.01520
                                     0.01101
                      1.34859
                                                    0.01179
2. Quantiles for each variable:
      25%
            50% 75% 97.5%
```

1.210 2.093 2.764 3.628 6.308

2.2 Proportional hazards models

The Cox proportional hazards model (Cox, 1972) expresses the hazard function h(t) of the survival time of each individual of the target population as the product of a common baseline hazard function $h_0(t)$, which determines the shape of h(t), and an exponential term which includes the relevant covariates x with regression coefficients β as follows:

$$h(t \mid h_0, \boldsymbol{\beta}) = h_0(t) \exp\left\{ \boldsymbol{x}^{\top} \boldsymbol{\beta} \right\}. \tag{7}$$

The estimation of the regression coefficients in the Cox model under the frequentist approach can be obtained without specifying a model for the baseline hazard function by using partial likelihood methodology (Cox, 1972). This is not the case of Bayesian analysis which in general needs to specify some model for the baseline hazard (Christensen et al., 2011). Depending on the context of the study, baseline hazard misspecification can imply a loss of valuable model information that makes it impossible to fully report the estimation of the outcomes of interest, such as probabilities or survival curves for relevant covariate patterns (Royston, 2011). This is specially important in survival studies where $h_0(t)$ represents the natural course of a disease or an infection, or even the control group when comparing several treatments.

Baseline hazard functions can be defined through parametric or semi-parametric approaches. Parametric models give restricted shapes which do not allow the presence of irregular behaviour. Semi-parametric choices result in more flexible baseline hazard shapes that allow for multimodal patterns such as piecewise constant functions or spline functions. Theoretical and methodological aspects of more flexible approaches to define the baseline hazard function can be found in several specific references such as Ibrahim et al. (2001), Lin et al. (2015), Mitra and Müller (2015), Bogaerts et al. (2017), and Lázaro et al. (2020).

2.2.1 Model specification

The proportional hazards (PH) model implemented here is also illustrated with the *larynx* dataset (see Section 2.1). Survival time for each individual is modelled by means of the hazard function:

$$h(t \mid h_0, \beta) = h_0(t) \exp\left\{\beta_2 \mathbf{I}_{(\text{stage}=2)} + \beta_3 \mathbf{I}_{(\text{stage}=3)} + \beta_4 \mathbf{I}_{(\text{stage}=4)} + \beta_5 \text{age} + \beta_6 \text{diagyr}\right\},\tag{8}$$

with baseline hazard function defined as a mixture of piecewise constant functions, $h_0(t \mid \lambda) = \sum_{k=1}^K \lambda_k \, I_{(a_{k-1},a_k]}(t), \, t > 0$, where $\lambda = (\lambda_1, \ldots, \lambda_K)$ and $I_{(a_{k-1},a_k]}(t)$ is the indicator function defined as 1 when $t \in (a_{k-1},a_k]$ and 0 otherwise. We consider a total number of knots K=3 and an equally-spaced partition of the time axis from $a_0=0$ to $a_4=10.70$ which corresponds to the longest survival time observed. Prior scenario is set under a non-informative independent framework with a N(0,0.001) for β 's and an independent gamma distributions, Ga(0.01,0.01), for each λ .

Piecewise constant baseline hazard functions can accommodate different shapes of the hazard depending on the particular characteristics of the partition of the time axis (See Breslow (1974), Murray et al. (2016), and Lee et al. (2016) for different proposals in the subject). Similarly, there is a wide range of approaches to define marginal prior distributions for h_0 parameters, from prior independence to prior correlation. Correlated scenarios account for shape restrictions and also avoid overfitting and strong irregularities in the estimation process (Lázaro et al., 2020).

The likelihood function of this specific model is not implemented in JAGS, so the "zeros trick" approach using a Poisson distribution has been used to specify it indirectly (Ntzoufras, 2009; Lunn et al., 2012).

2.2.2 Model implementation

The variables age, diagyr, stage, and X are defined as in Section 2.1. In addition, as previously commented, piecewise constant is handled in JAGS considering the "zeros trick" (Ntzoufras, 2009; Lunn et al., 2012). To execute it, it is necessary to reformat the individual times (both observed and right-censored) to build an auxiliary variable (int.obs) that identifies the interval in which the *i*th individual experiences the event of interest. So, the following code is dedicated to define a time axis partition (i.e., intervals) and the int.obs variable:

```
R> # Time axis partition
R> K <- 3 # number of intervals
R> a <- seq(0, max(larynx$time) + 0.001, length.out = K + 1)
R> # int.obs: vector that tells us at which interval each observation is
R> int.obs <- matrix(data = NA, nrow = nrow(larynx), ncol = length(a) - 1)
R> d <- matrix(data = NA, nrow = nrow(larynx), ncol = length(a) - 1)
R> for(i in 1:nrow(larynx)) {
    for(k in 1:(length(a) - 1)) {
        d[i, k] <- ifelse(time[i] - a[k] > 0, 1, 0) * ifelse(a[k + 1] - time[i] > 0, 1, 0)
        int.obs[i, k] <- d[i, k] * k
    }
H    int.obs <- rowSums(int.obs)</pre>
```

Listing 2 shows a generic implementation of a PH piecewise constant model in BUGS syntax using *larynx* data. Once the variables have been defined, a list with all the elements required in the model is created:

```
model {
    for(i in 1:n)
2
       for(k in 1:int.obs[i]){
3
         cond[i,k] \leftarrow step(time[i] - a[k+1])
4
         HH[i,k] \leftarrow cond[i,k] * (a[k+1]-a[k]) * lambda[k] +
                     (1-cond[i,k]) * (time[i]-a[k]) * lambda[k]
      # Cumulative hazard function
      H[i] <- sum(HH[i,1:int.obs[i]])
9
10
    for(i in 1:n){
11
      # Linear predictor
12
       elinpred[i] <- exp(inprod(beta[],X[i,]))
13
      # Log-hazard function
14
      logHaz[i] <- log(lambda[int.obs[i]] * elinpred[i])</pre>
15
      # Log-survival function
16
      logSurv[i] <- -H[i] * elinpred[i]</pre>
17
18
      # Definition of the log-likelihood using zeros trick
19
       phi[i] <- 100000 - delta[i] * logHaz[i] - logSurv[i]
20
       zeros[i] ~ dpois(phi[i])
21
    }
22
23
    # Prior distributions
24
    for(1 in 1:Nbetas) \{ beta[1] \sim dnorm(0,0.001) \}
25
26
    for (k \text{ in } 1:m) \{ lambda[k] \sim dgamma(0.01,0.01) \}
27
```

```
R> d.jags <- list(n = nrow(larynx), m = length(a) - 1, delta = larynx$delta,
+ time = larynx$time, X = X[,-1], a = a, int.obs = int.obs, Nbetas = ncol(X) - 1,
+ zeros = rep(0, nrow(larynx)))</pre>
```

The initial values for each PH model parameter are passed to JAGS using a function that returns a list of random values:

```
R> i.jags <- function(){ list(beta = rnorm(ncol(X) - 1), lambda = runif(3, 0.1)) }
```

The vector of monitored/saved parameters is:

```
R> p.jags <- c("beta", "lambda")</pre>
```

Next, the JAGS model is compiled:

```
R> library("rjags")
R> m2 <- jags.model(data = d.jags, file = "PH.txt", inits = i.jags, n.chains = 3)</pre>
```

We now run the model for 1000 burn-in simulations:

```
R> update(m2, 1000)
```

Finally, the model is run for 50000 additional simulations to keep one in 10 so that a proper thinning is done:

```
R> res <- coda.samples(m2, variable.names = p.jags, n.iter = 50000, n.thin = 10)
```

Similarly to the first example (Section 2.1), numerical and graphical summaries of the model parameters can be obtained using the summary and densplot functions, respectively. Gelman and Rubin's convergence diagnostic can be calculated with the gelman.diag function, and the traceplot function provides a visual way to inspect sampling behaviour and assesses mixing across chains and convergence.

Next, simulations from the three Markov chains are merged together for inference:

```
R> result <- as.mcmc(do.call(rbind, res))</pre>
```

The posterior samples of each parameter are obtained by:

```
R> beta2 <- result[,1]; beta3 <- result[,2]; beta4 <- result[,3]; beta5 <- result[,4]
R> beta6 <- result[,5]; lambda1 <- result[,6]; lambda2 <- result[,7]; lambda3 <- result[,8]
```

Table 1: Posterior summaries for the PH model parameters.

Parameter	Mean	SD	2.5%	50%	97.5%	$P(\cdot > 0 \mid \text{data})$
β_2 (stage=2)	0.152	0.474	-0.811	0.163	1.050	0.633
β_3 (stage=3)	0.672	0.363	-0.037	0.672	1.388	0.969
β_4 (stage=4)	1.804	0.443	0.924	1.809	2.659	1.000
eta_5 (age)	0.215	0.155	-0.086	0.214	0.523	0.918
eta_6 (diagyr)	-0.042	0.167	-0.368	-0.042	0.288	0.400
λ_1	0.069	0.021	0.035	0.066	0.116	1.000
λ_2	0.104	0.035	0.048	0.100	0.185	1.000
λ_3	0.079	0.064	0.008	0.062	0.246	1.000

Table 1 shows posterior summaries for the PH model parameters using *larynx* data.

The last column of Table 1 contains the posterior probability that the corresponding parameter is positive. A probability equal to 0.5 indicates that a positive value of the parameter is equally likely than a negative one. Once we have the posterior sample of each parameter stored, the calculation of this posterior probability, for example for β_2 , is given by mean (beta2>0).

A relevant quantity for PH models is the *hazard ratio* (HR), also called *relative risk*, between two individuals with covariate vectors x_1 and x_2 . This measure is defined as:

$$\operatorname{HR}(\boldsymbol{x}_1, \boldsymbol{x}_2, h_0, \boldsymbol{\beta}) = \frac{h(t \mid \boldsymbol{x}_1, h_0, \boldsymbol{\beta})}{h(t \mid \boldsymbol{x}_2, h_0, \boldsymbol{\beta})} = \exp\left\{(\boldsymbol{x}_1 - \boldsymbol{x}_2)^{\top} \boldsymbol{\beta}\right\},$$

and it is time independent. As an illustration, we can easily summarise the posterior distribution of the HR between two men of the same age and diagyr (year of diagnosis) but one in stage=3 and the other in stage=4:

```
R> HR.s3_s4 <- exp(beta3 - beta4)
R> summary(HR.s3_s4)
Iterations = 1:15000
Thinning interval = 1
Number of chains = 1
Sample size per chain = 15000
```

 Empirical mean and standard deviation for each variable, plus standard error of the mean:

```
        Mean
        SD
        Naive SE Time-series SE

        0.354210
        0.163810
        0.001338
        0.001338
```

2. Quantiles for each variable: 2.5% 25% 50% 75% 97.5% 0.1384 0.2404 0.3217 0.4297 0.7667

3 Mixture cure models

3.1 Cure models

Cure models deal with target populations in which a part of the individuals cannot experience the event of interest. This type of models has widely been matured as a consequence of the discovery and development of new treatments against cancer. The rationale of considering a cure subpopulation comes from the idea that a successful treatment removes totally the original tumor and the individual cannot experience any recurrence of the disease. These models allow to estimate the probability of cure, a key and valuable outcome in cancer research.

Mixture cure models are the most popular cure models (Berkson and Gage, 1952). They consider the target population as a mixture of susceptible and non-susceptible individuals for the event of interest. Let Z be a cure random variable defined as Z=0 for susceptible and Z=1 for cured or immune individuals. Cure and non cure probabilities are $P(Z=1)=\eta$ and $P(Z=0)=1-\eta$, respectively. The survival function for each individual in the cured and uncured subpopulation, $S_c(t)$ and $S_u(t), t>0$, respectively, is

$$S_u(t) = P(T > t \mid Z = 0), \quad S_c(t) = P(T > t \mid Z = 1),$$
 (9)

and the general survival function for T can be expressed as $S(t) = P(T > t) = \eta + (1 - \eta)S_u(t)$. It is important to point out that $S_u(t)$ is a proper survival function but S(t) is not. It goes to η and not to zero when t goes to infinity.

The effect of a baseline covariate vector \boldsymbol{x} on the cure fraction η for each individual is typically modelled by means of a logistic link function, $\operatorname{logit}(\eta)$, but the probit link or the complementary log-log link could also be used. Covariates for modelling T in the uncured subpopulation are usually considered via Cox models. Cure fraction model is usually known as the *incidence model* and the survival model (i.e., time-to-event T in the uncured subpopulation) as the *latency model* Klein et al. (2013).

3.2 bmt dataset

We consider a bone marrow transplant dataset, referred to as bmt. It is available from the smcure package (Cai et al., 2012):

```
R> library("smcure")
R> data("bmt")
R> str(bmt)
'data.frame': 91 obs. of 3 variables:
$ Time : num   11 14 23 31 32 35 51 59 62 78 ...
$ Status: num   1 1 1 1 1 1 1 1 1 1 ...
$ TRT : num   0 0 0 0 0 0 0 0 0 ...
```

This dataset refers to a bone marrow transplant study for the refractory acute lymphoblastic leukemia patients, in which 91 patients were divided into two treatment groups (Kersey et al., 1987). The following variables were observed for each patient:

- Time: time to death (in days).
- Status: censoring indicator (1: if patient is uncensored; 0: otherwise).
- TRT: treatment group indicator (0: allogeneic; 1: autologous).

3.3 Model specification

The (cure subpopulation) incidence model for each individual is expressed in terms of a logistic regression:

$$logit[\eta(\beta_{C1}, \beta_{C2})] = \beta_{C1} + \beta_{C2} TRT, \tag{10}$$

where β_{C1} represents an intercept and β_{C2} is the regression coefficient for the TRT covariate.

Survival time for each individual in the uncured subpopulation is modelled from a proportional hazards specification:

$$h_u(t \mid h_0, \beta_U) = h_0(t) \exp\left\{\beta_U \text{TRT}\right\},\tag{11}$$

with $h_0(t) = \lambda \alpha t^{\alpha-1}$ specified as a Weibull baseline hazard function, where α and λ are the shape and scale parameters, respectively; and β_U is the regression coefficient for the TRT covariate.

We assume prior independence and specify prior marginal distributions based on non-informative distributions commonly employed in the literature. The β 's follow a N(0, 0.001), while λ and α follow a Gamma(0.01, 0.01) and a Un(0, 10), respectively.

The likelihood function for mixture cure models is not implemented in JAGS, so the "zeros trick" approach using a Poisson distribution has also been used to specify it indirectly (Ntzoufras, 2009; Lunn et al., 2012).

3.4 Model implementation

We have created two design matrices, one for model (10) and another for model (11), with the TRT covariate:

```
R> XC <- model.matrix(~ TRT, data = bmt) # Reference = allogeneic
R> XU <- model.matrix(~ TRT, data = bmt)
R> XU <- matrix(XU[,-1], ncol = 1) # Remove intercept</pre>
```

Listing 3 shows a generic implementation of a mixture cure model in BUGS syntax using bmt data.

Once the variables have been defined, a list with all the elements required in the model is created:

The initial values for each mixture cure model parameter are passed to JAGS using a function that returns a list of random values:

```
R> i.jags <- function() {
+    list(betaC = rnorm(ncol(XC)), betaU = rnorm(ncol(XU)), lambda = runif(1), alpha = runif(1))
+  }</pre>
```

The vector of monitored/saved parameters is:

```
R> p.jags <- c("betaC", "betaU", "alpha", "lambda")</pre>
```

Next, the JAGS model is compiled:

```
R> library("rjags")
R> m3 <- jags.model(data = d.jags, file = "Cure.txt", inits = i.jags, n.chains = 3)</pre>
```

We now run the model for 10000 burn-in simulations:

Listing 3: Mixture cure model in BUGS syntax (file named as Cure.txt).

```
model {
     for(i in 1:n){
2
       # Logistic regression model (cured subpopulation)
       logit (eta[i]) <- inprod (betaC[], XC[i,])
       # PH model (uncured subpopulation)
       # Weibull baseline
       base[i] <- lambda * alpha * pow(t[i], alpha-1)
       elinpred[i] <- exp(inprod(betaU[],XU[i,]))</pre>
       # Log-hazard function
10
       logHaz[i] <- log(base[i] * elinpred[i])</pre>
11
       # Log-survival function
12
       logSurv[i] <- -lambda * pow(t[i], alpha) * elinpred[i]
13
14
       # Definition of the log-likelihood using zeros trick
15
       logLike[i] \leftarrow delta[i] * (log(1-eta[i]) + logHaz[i] + logSurv[i]) +
                      (1-delta[i]) * log(eta[i] + (1-eta[i]) * exp(logSurv[i]))
       phi[i] <- 100000 - logLike[i]
18
       zeros[i] ~ dpois(phi[i])
19
     }
20
21
     # Prior distributions
22
     \begin{array}{lll} for(1 & in & 1:NbetasC) \{ & betaC[1] & \tilde{} & dnorm(0,0.001) \\ for(1 & in & 1:NbetasU) \{ & betaU[1] & \tilde{} & dnorm(0,0.001) \\ \end{array}
23
24
     lambda ~ dgamma(0.01,0.01)
alpha ~ dunif(0,10)
25
26
27
```

```
R> update(m3, 10000)
```

Finally, the model is run for 100000 additional simulations to keep one in 100 so that a proper thinning is done:

```
R> res <- coda.samples(m3, variable.names = p.jags, n.iter = 100000, n.thin = 100)
```

Similarly to the first example (Section 2.1), numerical and graphical summaries of the model parameters can be obtained using the summary and densplot functions, respectively. Gelman and Rubin's convergence diagnostic can be calculated with the gelman.diag function, and the traceplot function provides a visual way to inspect sampling behaviour and assesses mixing across chains and convergence.

Next, simulations from the three Markov chains are merged together for inference:

```
R> result <- as.mcmc(do.call(rbind, res))</pre>
```

The posterior samples of each parameter are obtained by:

```
R> alpha <- result[,1]; betaC1 <- result[,2]; betaC2 <- result[,3]
R> betaU <- result[,4]; lambda <- result[,5]</pre>
```

Table 2 shows posterior summaries for the mixture cure model parameters using bmt data.

Table 2: Posterior summaries for the mixture cure model parameters.

Parameter	Mean	SD	2.5%	50%	97.5%	$P(\cdot > 0 \mid \text{data})$
β_{C1} (intercept)	-1.015	0.349	-1.731	-1.002	-0.365	0.001
$eta_{C2}\left(\mathtt{TRT} ight)$	-0.419	0.519	-1.445	-0.417	0.591	0.208
β_U (TRT)	0.762	0.269	0.239	0.760	1.294	0.998
α	1.143	0.105	0.943	1.140	1.354	1.000
λ	0.002	0.001	0.000	0.002	0.006	1.000

As we discussed earlier, the cure fraction (η) is a relevant quantity for mixture cure models. For allogeneic (TRT=0) or autologous (TRT=1) treated patients, it is modelled as:

$$\eta(\text{TRT}, \beta_{C1}, \beta_{C2}) = \frac{\exp(\beta_{C1} + \beta_{C2}\text{TRT})}{1 + \exp(\beta_{C1} + \beta_{C2}\text{TRT})}.$$

We can easily summarise the posterior distribution of the cure fraction for individuals in both groups:



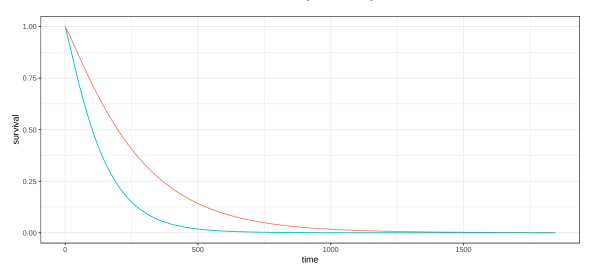


Figure 3: Posterior mean of the uncured survival function from the mixture cure model (11).

```
R> CP.allo <- exp(betaC1) / (1 + exp(betaC1))</pre>
R> CP.auto <- exp(betaC1 + betaC2) / (1 + exp(betaC1 + betaC2))</pre>
R> summary(cbind(CP.allo, CP.auto))
    CP.allo
                       CP.auto
        :0.01604
                    Min.
                            :0.02277
 1st Qu.:0.22426
                    1st Qu.:0.15680
Median :0.26850
                    Median :0.19477
       :0.27146
                            :0.19925
                    Mean
 3rd Qu.:0.31515
                    3rd Qu.: 0.23699
Max.
        :0.65573
                    Max.
                           :0.55898
```

The uncured survival curve based on posterior samples is another relevant information in this type of studies. So, from the posterior samples obtained above, we can summarise the posterior distribution of the mean value of the uncured survival curve for allogeneic and autologous treated patients in a grid of points as follows:

Figure 3 shows the difference between both curves using the code below:

4 Competing risks models

Competing risks occur when the survival process includes more than one cause of failure. In the case of different causes of death it is only possible to report the first event to occur (Putter et al., 2007). There are different approaches for competing risk models: multivariate time to failure model, the cause-specific hazards model, the mixture model, the subdistribution model, and the full specified subdistribution model (Ge and Chen, 2012). We will only consider here the cause-specific hazards model, possibly the most popular of them.

Let T_k be the random variable that represents the time-to-event from cause k, for k = 1, ..., K, where K is the total number of different events. The only survival time observed $T = \min\{T_1, T_2, ..., T_K\}$ usually corresponds to the earliest

cause together with their indicator $\delta = k$ when the subsequent individual experiences the event due to cause k. The key concept in a competing risk model is the cause-specific hazard function for cause k, which assesses the hazard of experiencing the event k in the presence of the rest of competing events, and it is defined by:

$$h_k(t) = \lim_{\Delta t \to 0} \frac{P(t \le T < t + \Delta t, \delta = k \mid T \ge t)}{\Delta t}.$$
 (12)

Inference for each $h_k(t)$ considers the observed failure times for cause k as censored observations for the rest of events. Another relevant concept in the competing framework is the cumulative incidence function for cause k, defined as follows:

$$F_k(t) = P(T \le t, \delta = k) = \int_0^t h_k(u) S(u) du,$$
 (13)

where S(t) is the overall survival function. It is typically expressed in terms of the different cause-specific hazard functions in accordance with

$$S(t) = P(T > t) = \exp \left\{ -\sum_{k=1}^{K} \int_{0}^{t} h_{k}(u) du \right\}.$$

The cumulative incidence function is not a proper cumulative distribution function, i.e., the probability that the subsequent individual fails from cause k is $F_k(\infty) = P(\delta = k) \neq 1$. The sub-survival function for cause k, defined as $S_k(t) = P(T > t, \delta = k)$, is also not a proper survival function.

4.1 *okiss* dataset

We consider a stem-cell transplanted patients dataset, referred to as *okiss*. It is available from the compeir package (Grambauer and Neudecker, 2011):

```
R> library("compeir")
R> data("okiss")
R> str(okiss)
'data.frame': 1000 obs. of 4 variables:
$ time : 'times' num 21 6 14 12 11 18 10 8 17 10 ...
..- attr(*, "format")= chr "h:m:s"
$ status: num 2 1 2 2 2 2 7 2 2 2 ...
$ allo : num 1 1 1 1 1 1 0 1 1 ...
$ sex : Factor w/ 2 levels "f", "m": 1 1 1 2 2 2 2 1 2 2 ...
```

This dataset provides information about a sub-sample of 1000 patients enrolled in the ONKO-KISS programme, which is part of the German National Reference Centre for Surveillance of Hospital-Acquired Infections (Dettenkofer et al., 2005). These patients have been treated by peripheral blood stem-cell transplantation, which has become a successful therapy for severe hematologic diseases. After transplantation, patients are neutropenic, i.e., they have a low count of white blood cells, which are the cells that primarily avert infections (Beyersmann et al., 2007). Occurrence of bloodstream infection during neutropenia is a severe complication. The following variables were observed for each patient:

- time: time (in days) of neutropenia until first event.
- status: event status indicator (1: infection; 2: end of neutropenia; 7: death; 11: censored observation).
- allo: transplant type indicator (0: autologous; 1: allogeneic).
- sex: sex of each patient (m: if patient is male; f: if patient is female).

We have redefined the status variable using an auxiliary one (delta) in a matrix format:

```
R> delta <- matrix(c(as.integer(okiss$status == 1), as.integer(okiss$status == 2),
     as.integer(okiss$status == 7)), ncol = 3)
R> head(delta)
     [,1] [,2] [,3]
          1
[1,]
       0
            0
        1
[3,]
        0
            1
                  Ω
[4,]
        0
             1
                  Ω
        0
[5,]
[6,]
```

where the events 1 (infection), 2 (end of neutropenia) and 7 (death) are indicated with a value of 1 in column 1, 2 or 3, respectively, and a row with only 0's represents a censored observation.

4.2 Model specification

Cause-specific hazard functions for infection (k = 1), end of neutropenia (k = 2), and death (k = 3) are modelled from a proportional hazard specification:

$$h_k(t \mid h_{0k}, \beta_k) = h_{0k}(t) \exp\{\beta_{1k} \text{allo} + \beta_{2k} \text{sex}\}, \quad k = 1, 2, 3,$$
 (14)

with $h_{0k}(t) = \lambda_k \, \alpha_k \, t^{\alpha_k - 1}$ for event k specified as a Weibull baseline hazard function, where α_k and λ_k are the shape and scale parameters, respectively; and $\boldsymbol{\beta}_k = (\beta_{1k}, \beta_{2k})^{\top}$ are regression coefficients for the allo and sex covariates, respectively, for k = 1, 2, 3. We assume prior independence and specify prior marginal based on non-informative distributions commonly employed in the literature. The β 's follow a N(0,0.001), while λ 's and α 's follow a Gamma(0.01,0.01) and a Un(0,10), respectively.

4.3 Model implementation

We have created a design matrix X with the covariates allo and sex:

```
R> X <- model.matrix(\tilde{} allo + sex, data = okiss) # Reference = female R> X <- X[,-1] # Remove intercept
```

Listing 4 shows a generic implementation of a competing risks model in BUGS syntax using okiss data.

Listing 4: Competing risks model in BUGS syntax (file named as **CR.txt**).

```
model {
2
    for(i in 1:n){
      for(k in 1:Nrisks){
3
        # Weibull baseline
         base[i,k] \leftarrow lambda[k] * alpha[k] * pow(t[i],alpha[k]-1)
         elinpred [i,k] <- exp(inprod(beta[,k],X[i,])) \\
7
         # Log-hazard functions
        logHaz[i,k] <- log(base[i,k] * elinpred[i,k])</pre>
         # Log-survival functions
9
         logSurv[i,k] <- -lambda[k] * pow(t[i],alpha[k]) * elinpred[i,k]
10
11
12
      # Definition of the log-likelihood using zeros trick
13
      phi[i] <- 100000 - inprod(delta[i,], logHaz[i,]) - sum(logSurv[i,])
14
      zeros[i] ~ dpois(phi[i])
15
    }
16
17
    # Prior distributions
18
    for(k in 1:Nrisks){
19
      for(1 in 1:Nbetas) \{ beta[1,k] ~ dnorm(0,0.001) \}
      lambda[k] ~ dgamma(0.01,0.01)
21
      alpha[k] \sim dunif(0,10)
22
    }
23
24
```

Once the variables have been defined, a list with all the elements required in the model is created:

The initial values for each competing risks model parameter are passed to JAGS using a function that returns a list of random values:

Next, the JAGS model is compiled:

```
R> library("rjags")
R> m4 <- jags.model(data = d.jags, file = "CR.txt", inits = i.jags, n.chains = 3)</pre>
```

We now run the model for 1000 burn-in simulations:

```
R> update(m4, 1000)
```

Finally, the model is run for 10000 additional simulations to keep one in 10 so that a proper thinning is done:

```
R> res <- coda.samples(m4, variable.names = p.jags, n.iter = 10000, n.thin = 10)
```

Similarly to the first example (Section 2.1), numerical and graphical summaries of the model parameters can be obtained using the summary and densplot functions, respectively. Gelman and Rubin's convergence diagnostic can be calculated with the gelman.diag function, and the traceplot function provides a visual way to inspect sampling behaviour and assesses mixing across chains and convergence.

Next, simulations from the three Markov chains are merged together for inference:

```
R> result <- as.mcmc(do.call(rbind, res))</pre>
```

The posterior samples of each parameter are obtained by:

```
R> alpha1 <- result[,1]; alpha2 <- result[,2]; alpha3 <- result[,3]; beta11 <- result[,4]
R> beta21 <- result[,5]; beta12 <- result[,6]; beta22 <- result[,7]; beta13 <- result[,8]
R> beta23 <- result[,9]; lambda1 <- result[,10]; lambda2 <- result[,11]; lambda3 <- result[,12]</pre>
```

Table 3 shows posterior summaries for the competing risks model parameters using okiss data.

Table 3: Posterior summaries for the competing risks model parameters.

Table 3: Posterior summaries for the competing risks model parameters.									
Parameter	Mean	SD	2.5%	50%	97.5%	$P(\cdot > 0 \mid \text{data})$			
Infection (h_1)									
β_{11} (allo)	-0.523	0.151	-0.817	-0.523	-0.225	0.000			
eta_{21} (sex)	0.158	0.148	-0.130	0.156	0.451	0.857			
λ_1	0.014	0.003	0.009	0.014	0.022	1.000			
α_1	1.137	0.070	1.003	1.136	1.278	1.000			
	End of neutropenia (h_2)								
β_{12} (allo)	-1.193	0.075	-1.340	-1.193	-1.046	0.000			
β_{22} (sex)	-0.102	0.073	-0.244	-0.103	0.042	0.081			
λ_2	0.008	0.001	0.006	0.008	0.010	1.000			
α_2	2.033	0.045	1.947	2.034	2.120	1.000			
Death (h ₃)									
β_{13} (allo)	-0.617	0.747	-2.001	-0.650	0.945	0.193			
β_{23} (sex)	0.446	0.730	-0.859	0.408	1.986	0.718			
λ_3	0.000	0.000	0.000	0.000	0.000	1.000			
α_3	2.628	0.418	1.813	2.623	3.466	1.000			

As discussed in Section 4, the cumulative incidence function $F_k(t)$ in (13) is the most appropriate way to analyse the evolution of each cause k over time. For our proportional hazard specification (14), it is given by:

$$F_k(t) = \int_0^t h_k(t \mid h_{0k}, \boldsymbol{\beta}_k) \exp\left\{-\sum_{l=1}^3 \int_0^u h_l(v \mid h_{0l}, \boldsymbol{\beta}_l) \, \mathrm{d}v\right\} \, \mathrm{d}u,\tag{15}$$

where $h_k(t \mid h_{0k}, \boldsymbol{\beta}_k)$ is defined in (14).

The integral in (15) has no closed form, so some approximate method of integration is required. To do this, we first have created a function fk which describes the integrand of (15):

```
R> fk <- function(u.vect, lambda, alpha, beta, x, k) {
    res <- sapply(u.vect, function(u) {
        # Cause-specific hazard
        hk <- lambda[k] * alpha[k] * (u^(alpha[k] - 1)) * exp(sum(unlist(beta[,k]) * x))
        # Cumulative cause-specific hazard
        Hk <- lambda * (rep(u, length(lambda))^alpha) * exp((t(beta) %*% matrix(x, ncol = 1))[,1])
        # Cause-specific hazard x Overall survival
        aux <- hk * exp(-sum(Hk))
        return(aux)
    }
}
return(res) }</pre>
```

Next, we have created a function cif that computes $F_k(t)$ in (15) by integrating out the fk function using the integral function available from the pracma package (Borchers, 2019).

```
R> library("pracma")
R> cif <- function(tt, lambda, alpha, beta, x, k){
+ return(integral(fk, xmin = 0, xmax = tt, method = "Simpson", lambda = lambda, alpha = alpha,
+ beta = beta, x = x, k = k)) }</pre>
```

Finally, we have constructed a function mcmc_cif that takes the output from JAGS (variable obj) and computes $F_k(t)$ in (15) for a vector of times (variable t.pred) using covariates x. Note that mcmc_cif is based on the mclapply function, available from the parallel package to speed computations up (R Core Team, 2020).

```
R> library("parallel")
R> options(mc.cores = detectCores())
R> mcmc_cif <- function(obj, t.pred, x) {</pre>
     var.names <- names(obj)</pre>
     # Indices of beta's, alpha's, and lambda's
    b.idx <- which(substr(var.names, 1, 4) == "beta")</pre>
     a.idx <- which(substr(var.names, 1, 5) == "alpha")</pre>
     1.idx <- which(substr(var.names, 1, 6) == "lambda")</pre>
     # Number of causes and number of covariates
     K <- length(a.idx)</pre>
     n.b <- length(b.idx) / K</pre>
     # Sub-sample to speed up computations
     samples.idx <- sample(1:nrow(obj), 200)</pre>
     res <- lapply(1:K, function(k){
       sapply(t.pred, function(tt){
         aux <- mclapply(samples.idx, function(i){</pre>
            cif(tt, alpha = unlist(obj[i, a.idx]), lambda = unlist(obj[i, l.idx]),
               beta = matrix(unlist(c(res[i, b.idx])), nrow = n.b), x = x, k = k)
         })
         return (mean (unlist (aux)))
       })
     })
     return (res)
```

Hence, we redefine the MCMC output as a data.frame and set a vector of times to evaluate the cumulative incidence function. In this example, we are interested in this function when both covariates are 1 (i.e., allogeneic transplant and male).

```
R> res <- as.data.frame(result)
R> t.pred <- seq(0, 100, by = 2.5)
R> cum_inc <- mcmc_cif(res, t.pred, c(1, 1))</pre>
```

Figure 4, generated with the code below, shows the posterior mean of the cumulative incidence function for a man with an allogeneic transplant for the three types of events considered in the *okiss* data.

```
R> library("ggplot2")
R> df <- data.frame(cif = unlist(cum_inc), time = t.pred,
+ cause = rep(c("infection", "end of neutropenia", "death"), each = length(t.pred)))
R> ggplot(data = df, aes(x = time, y = cif, group = cause)) + geom_line(aes(color = cause)) +
+ ylab("cumulative incidence") + ylim(c(0,1)) + theme_bw() + theme(legend.position = "top")
```

5 Multi-state models

Multi-state models are a class of stochastic processes which account for event history data, with individuals who may experience different events in time. Relevant data are the events and their subsequent survival times. Multi-state models allow for different structures depending on the number and relationships between the states (Andersen and Keiding, 2002). Outputs of interest are the usual in survival analysis (sometimes with a specific vocabulary, e.g., the hazard function which is now called *transition intensity*) to which transition probabilities are added.

We concentrate on the illness-death model (also known as *disability model*) which is a particular multi-state model. This is a relevant model in irreversible diseases where a significant illness' progression increases the risk of a terminal event (Han et al., 2014; Armero et al., 2016a). The underlying stochastic process $\{Z(t), t \geq 0\}$ describes the state of an individual at time t, where t is time from entry into the initial state (state 1). The probabilistic behaviour of the process is determined by the subsequent hazard functions from which transitions probabilities between states, defined as $p_{jk}(s,t\mid\theta)=P(Z(t)=k\mid Z(s)=j,\theta)$, can be derived, where $s\leq t,j$ and k are states, $\sum_{j=k}^3 p_{jk}(s,t\mid\theta)=1$, for k=1,2,3, and θ is the vector of model parameters.

Each transition has its respective hazard function, for example, $h_{12}(t \mid \theta)$ is associated with time T_{12} from state 1 to state 2 $(1 \rightarrow 2)$, while $h_{13}(t \mid \theta)$ and $h_{23}(t \mid \theta)$ represent the hazard functions for the time T_{13} $(1 \rightarrow 3)$ and T_{23} $(2 \rightarrow 4)$

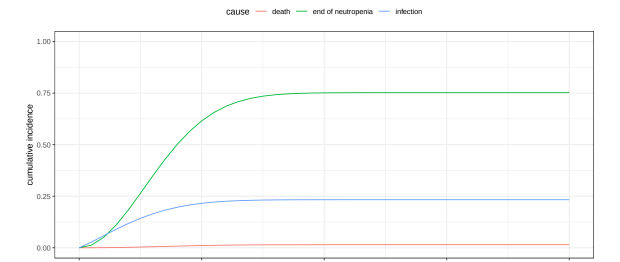


Figure 4: Posterior mean of the cumulative incidence function for allo=1 (allogeneic transplant) and sex=1 (male) from the competing risks model (14).

time

3), respectively. If $h_{23}(t \mid \theta)$ does not depend of the time in which transition $1 \to 2$ occurs, the process will be known as *Markovian*. Otherwise, it is called *semi-Markovian* and transition probabilities $2 \to 3$ will depend on the particular value of T_{12} .

Transition probabilities between states and hazard functions for the semi-Markovian specification are connected as follows (Andersen and Perme, 2008):

$$p_{11}(s,t\mid\boldsymbol{\theta}) = \exp\bigg\{-\int_{s}^{t} \left[h_{12}(u\mid\boldsymbol{\theta}) + h_{13}(u\mid\boldsymbol{\theta})\right] du\bigg\},\tag{16}$$

$$p_{22}(s, t \mid \boldsymbol{\theta}, t_{12}) = \exp \left\{ -\int_{s}^{t} h_{23}(u - t_{12} \mid \boldsymbol{\theta}, t_{12}) \, du \right\}, \quad t_{12} < s, \tag{17}$$

$$p_{12}(s,t \mid \boldsymbol{\theta}) = \int_{s}^{t} p_{11}(s,u \mid \boldsymbol{\theta}) h_{12}(u \mid \boldsymbol{\theta}) p_{22}(u,t \mid \boldsymbol{\theta},u) du,$$
 (18)

$$p_{13}(s, t \mid \boldsymbol{\theta}) = 1 - p_{11}(s, t \mid \boldsymbol{\theta}) - p_{12}(s, t \mid \boldsymbol{\theta}),$$
 (19)

$$p_{23}(s, t \mid \boldsymbol{\theta}) = 1 - p_{22}(s, t \mid \boldsymbol{\theta}),$$
 (20)

$$p_{33}(s,t\mid\boldsymbol{\theta}) = 1, \tag{21}$$

where the marginal probability $p_{22}(s, t \mid \boldsymbol{\theta})$ is obtained by integrating out $p_{22}(s, t \mid \boldsymbol{\theta}, t_{12})$ with regard to the density of T_{12} from 0 to s.

The standard specification of an illness-death model is through the hazard function of the relevant survival times, generally modelled by means of Cox models. Consequently, the specification of prior distributions for θ must be addressed to them in Section 2.2.

5.1 *heart2* dataset

We consider a heart transplant dataset, referred to as *heart2*. It is available from the p3state.msm package (Meira-Machado and Roca-Pardinas, 2012):

```
R> library("p3state.msm")
R> data("heart2")
R> str(heart2)
'data.frame': 103 obs. of 8 variables:
$ times1 : num 50 6 1 36 18 3 51 40 85 12 ...
$ delta : int 0 0 1 1 0 0 1 0 0 1 ...
$ times2 : num 0 0 15 3 0 0 624 0 0 46 ...
$ time : int 50 6 16 39 18 3 675 40 85 58 ...
$ status : int 1 1 1 1 1 1 1 1 1 ...
$ age : num -17.16 3.84 6.3 -7.74 -27.21 ...
$ year : num 0.123 0.255 0.266 0.49 0.608 ...
$ surgery: int 0 0 0 0 0 0 0 0 0 ...
```

This dataset provides information about a sample of 103 patients of the Stanford Heart Transplant Program (Crowley and Hu, 1977). The patients are initially on the waiting list (state 1) and can either be transplanted (state 2, non-terminal event) and then die (state 3, terminal event), or just one or none of them because they continue to be on the waiting list. The following variables were observed for each patient:

- times1: time of transplant/censoring time (state 2).
- delta: transplant indicator (1: yes; 0: no).
- times2: time to death since the transplant/censoring time (state 3).
- time: times1 + times2.
- status: censoring indicator (1: dead; 0: alive).
- age: age 48 years.
- year: year of acceptance (in years after 1 Nov 1967).
- surgery: prior bypass surgery (1: yes; 0: no).

The patients had the following characteristics: 4 were censored for both events (transplant and death), 24 moved from state 1 (waiting list) to state 2 (transplant) and survived, 30 moved from state 1 to state 3 (death) without going through state 2, and 45 moved from state 1 to state 2 and then to state 3.

We have redefined delta and status variables using an auxiliary one (event) in a matrix format:

```
R> event <- matrix(c(heart2$delta, heart2$status * (1 - heart2$delta),
    heart2$delta * heart2$status), ncol = 3)
R> head(event)
     [,1] [,2] [,3]
          1
[1,]
            1
[2,]
[3,]
        1
            0
[4,]
        1
            0
[5,]
        0
            1
[6,]
```

where each row represents the transitions of a patient, in which a 1 in columns 1, 2 and 3 indicates transition $1 \to 2$, $1 \to 3$, and $2 \to 3$, respectively, and a row with only 0's represents a censored observation.

5.2 Model specification

Hazard functions for survival times T_{12} , T_{13} and T_{23} are modelled using a proportional hazard specification:

$$h_{12}(t \mid h_{01}, \beta_1) = h_{01}(t) \exp\{\beta_{11} \operatorname{age} + \beta_{21} \operatorname{year} + \beta_{31} \operatorname{surgery}\}, \quad t > 0,$$
 (22)

$$h_{13}(t \mid h_{02}, \beta_2) = h_{02}(t) \exp\{\beta_{12} \text{age} + \beta_{22} \text{year} + \beta_{32} \text{surgery}\}, t > 0,$$
 (23)

$$h_{23}(t \mid h_{03}, \beta_3, T_{12} = t_{12}) = h_{03}(t - t_{12}) \exp\{\beta_{13} \text{age} + \beta_{23} \text{year} + \beta_{33} \text{surgery}\}, \quad t > t_{12},$$
 (24)

with $h_{0k}(t) = \lambda_k \alpha_k t^{\alpha_k - 1}$ specified as a Weibull baseline hazard function, where α_k and λ_k are the shape and scale parameters, respectively, for k = 1, 2, 3; and $\beta_k = (\beta_{1k}, \beta_{2k}, \beta_{3k})^{\top}$ are regression coefficients for the age, year and surgery covariates, respectively, We assume prior independence and specify prior marginal based on non-informative distributions commonly employed in the literature. The β 's follow a N(0,0.001), while λ 's and α 's follow a Gamma(0.01,0.01) and a Un(0,10), respectively. Note that in (24) we adopt the semi-Markovian specification, but it could also be the Markovian one (Alvares et al., 2019).

5.3 Model implementation

We have created a design matrix X with the covariates age, year and surgery, and defined a time $3 = t - t_{12}$ variable according to semi-Markovian specification:

```
R> X <- model.matrix(~ age + year + surgery, data = heart2)
R> X <- X[,-1] # Remove intercept
R> time3 <- heart2$times2
R> time3[time3 == 0] <- 0.0001</pre>
```

The values of time3 equal to zero have been replaced by 0.0001 to avoid computational problems when calculating $(t-t_{12})^{\alpha_3-1}$.

Listing 5 shows a generic implementation of an illness-death model in BUGS syntax using heart2 data.

Once the variables have been defined, a list with all the elements required in the model is created:

Listing 5: Illness-death model in BUGS syntax (file named as IllDeath.txt).

```
model {
     for(i in 1:n)
2
       # Linear predictor
3
       elinpred[i,1] <- exp(inprod(beta[,1],X[i,]))
4
       elinpred[i,2] \leftarrow exp(inprod(beta[,2],X[i,]))
       elinpred[i,3] \leftarrow exp(inprod(beta[,3],X[i,]))
       # Log-hazard functions
       logHaz[i,1] <- log(lambda[1] * alpha[1] * pow(t1[i],alpha[1]-1) * elinpred[i,1])
       logHaz[i,2] <- log(lambda[2] * alpha[2] * pow(t2[i], alpha[2]-1) * elinpred[i,2])
       logHaz[i,3] <- log(lambda[3] * alpha[3] * pow(t3[i], alpha[3]-1) * elinpred[i,3])
10
       # Log-survival functions
11
       logSurv[i,1] <- -lambda[1] * pow(t1[i], alpha[1]) * elinpred[i,1]
12
       logSurv[i,2] <- -lambda[2] * pow(t2[i], alpha[2]) * elinpred[i,2]
13
       logSurv[i,3] <- -lambda[3] * pow(t3[i],alpha[3]) * elinpred[i,3]
14
15
       # Definition of the log-likelihood using zeros trick
16
       phi[i] <- 100000 - inprod(event[i,],logHaz[i,]) - sum(logSurv[i,])
17
       zeros[i] ~ dpois(phi[i])
18
     }
19
20
     # Prior distributions
21
     for(k in 1:3)
22
       for (1 \text{ in } 1: \text{Nbetas}) \{ \text{ beta} [1, k] \ ^{\sim} \text{ dnorm} (0, 0.001) \}
23
       \begin{array}{ll} lambda[k] & \stackrel{\sim}{\phantom{}} dgamma(0.01,0.01) \\ alpha[k] & \stackrel{\sim}{\phantom{}} dunif(0,10) \end{array}
24
25
26
     }
27
```

```
R> d.jags <- list(n = nrow(heart2), t1 = heart2$times1, t2 = heart2$time, t3 = time3, 
+ X = X, event = event, zeros = rep(0, nrow(heart2)), Nbetas = ncol(X))
```

The initial values for each illness-death model parameter are passed to JAGS using a function that returns a list of random values:

```
R> i.jags <- function() {
+    list(beta = matrix(rnorm(3 * ncol(X)), ncol = 3), lambda = runif(3), alpha = runif(3))
+ }</pre>
```

The vector of monitored/saved parameters is:

```
R> p.jags <- c("beta", "alpha", "lambda")</pre>
```

Next, the JAGS model is compiled:

```
R> library("rjags")
R> m5 <- jags.model(data = d.jags, file = "IllDeath.txt", inits = i.jags, n.chains = 3)</pre>
```

We now run the model for 1000 burn-in simulations:

```
R> update(m5, 1000)
```

Finally, the model is run for 10000 additional simulations to keep one in 10 so that a proper thinning is done:

```
R> res <- coda.samples(m5, variable.names = p.jags, n.iter = 10000, n.thin = 10)
```

Similarly to the first example (Section 2.1), numerical and graphical summaries of the model parameters can be obtained using the summary and densplot functions, respectively. Gelman and Rubin's convergence diagnostic can be calculated with the gelman.diag function, and the traceplot function provides a visual way to inspect sampling behaviour and assesses mixing across chains and convergence.

Next, simulations from the three Markov chains are merged together for inference:

```
R> result <- as.mcmc(do.call(rbind, res))</pre>
```

The posterior samples of each parameter are obtained by:

```
R> alpha1 <- result[,1]; alpha2 <- result[,2]; alpha3 <- result[,3]
R> beta11 <- result[,4]; beta21 <- result[,5]; beta31 <- result[,6]
R> beta12 <- result[,7]; beta22 <- result[,8]; beta32 <- result[,9]
R> beta13 <- result[,10]; beta23 <- result[,11]; beta33 <- result[,12]
R> lambda1 <- result[,13]; lambda2 <- result[,14]; lambda3 <- result[,15]</pre>
```

Table 4: Posterior summaries for the illness-death model parameters.

Parameter	Mean	SD	2.5%	50%	97.5%	$P(\cdot > 0 \mid \text{data})$			
From waiting list to heart transplant (h_{12})									
β_{11}	0.048	0.015	0.020	0.047	0.077	1.000			
β_{21}	0.004	0.069	-0.130	0.004	0.140	0.521			
β_{31}	0.220	0.321	-0.440	0.229	0.826	0.760			
λ_1	0.042	0.016	0.018	0.039	0.079	1.000			
α_1	0.778	0.069	0.645	0.777	0.916	1.000			
	From waiting list to death (h_{13})								
β_{12}	-0.001	0.018	-0.034	-0.002	0.035	0.462			
eta_{22}	-0.243	0.115	-0.476	-0.241	-0.027	0.013			
β_{32}	-0.785	0.672	-2.223	-0.736	0.391	0.109			
λ_2	0.101	0.047	0.034	0.092	0.217	1.000			
$lpha_2$	0.379	0.058	0.272	0.376	0.500	1.000			
	From heart transplant to death (h_{23})								
β_{13}	0.055	0.022	0.014	0.055	0.099	0.997			
β_{23}	-0.008	0.094	-0.196	-0.007	0.174	0.470			
β_{33}	-0.986	0.469	-1.973	-0.961	-0.129	0.011			
λ_3	0.034	0.021	0.008	0.029	0.087	1.000			
α_3	0.602	0.074	0.463	0.598	0.756	1.000			

Table 4 shows posterior summaries for the illness-death model parameters using *heart2* data.

As discussed in Section 5, the transition probabilities (16)–(21) are the most appropriate way to analyse the evolution of each state over time. The implementation of the posterior distribution for these transition probabilities requires auxiliary functions, similar to the calculation of the cumulative incidence function in Section 4. To avoid unnecessary repetitions on how to calculate these quantities of interest, we will omit their implementation here. However, the code to reproduce Figure 5 is available in Appendix A.

6 Frailty models

Regression models include measurable covariates to improve the knowledge of the relevant failure times. However, in most survival studies, there are also individual heterogeneity that are not known or measurable. These elements are known in the statistical framework as *random effects*, but in the context of survival models they are the *frailty* elements (Aalen, 1994). They can approach individual characteristics as well as heterogeneity in groups or clusters (Ibrahim et al., 2001).

The most popular type of frailty models is the multiplicative shared-frailty model. It is a generalisation of the Cox regression model introduced by Clayton (1978) and extensively studied in Hougaard (2000). Let T_i the survival time for each individual in group i with hazard function described by:

$$h_i(t \mid h_0, \boldsymbol{\beta}, w_i) = w_i h_0(t) \exp\left\{\boldsymbol{x_i}^{\top} \boldsymbol{\beta}\right\}, \tag{25}$$

where w_i is the frailty term associated to group i. The usual probabilistic model for the frailty term is a gamma distribution with mean equal to one for identifiability purposes but also the positive stable and log-normal distributions can be considered (Vaupel et al., 1979). In addition, a unity mean can be considered as a neutral frontier because frailty values greater (lower) than one increases (decreases) the individual risk. An alternative way of incorporating a frailty term in the hazard function is via an additive element as follows:

$$h_i(t \mid h_0, \boldsymbol{\beta}, b_i) = h_0(t) \exp \left\{ \boldsymbol{x}_i^{\top} \boldsymbol{\beta} + b_i \right\},$$

where now b_i 's are commonly assumed as normally distributed with zero mean and unknown variance.

The Bayesian framework deals with frailty models in a conceptually simpler way than the frequentist one due to the Bayesian probability conception, introduced in Section 1. Hence, the inclusion of randomness through frailties in a Bayesian perspective does not add any conceptual complexity because the information regarding the risk function is expressed in probabilistic terms through its posterior distribution. Survival modelling with frailty terms is a wide issue of research that applies to all type of regression, competing risks, multivariate survival models, etc. and play a special role in joint models as we will discuss later.

6.1 kidney dataset

We consider a kidney infection dataset, referred to as kidney. It is available from the frailtyHL package (Ha et al., 2018):

- R> library("frailtyHL")
- R> data("kidney")
- R> str(kidney)

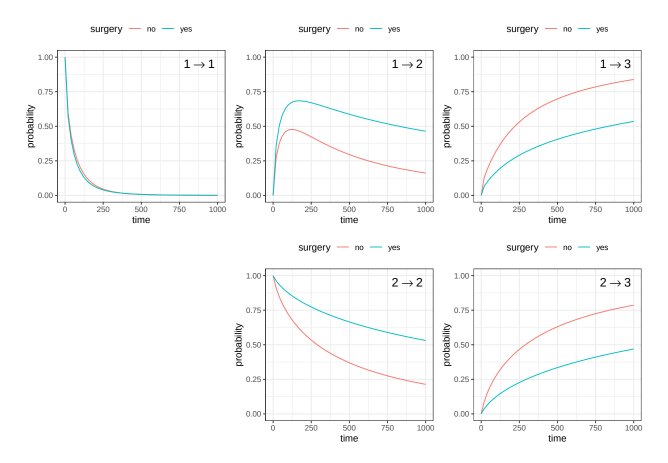


Figure 5: Posterior mean of all probability transitions from the illness-death model (22)–(24) for patients with and without prior bypass surgery and median values of age and year. Graphics on the top correspond to transitions from the initial state in the waiting list. Posterior probabilities for transitions from transplant (bottom row) assume the median time $T_{12}=26$ recorded from the waiting list to heart transplant.

```
'data.frame': 76 obs. of
                          10 variables:
                 1 1 2 2 3 3 4 4 5 5 ...
 $
           num
                 8
                   16 23 13 22 28 447 318 30 12 ...
                 1 1 1 0 1 1 1 1 1 1
 $
  status : num
                 28 28 48 48 32 32 31 32 10 10 ...
           num
                     2 2 1 1 2 2 1 1 ...
           num
                 1 1
  disease: Factor w/ 4 levels "Other", "GN", "AN", ...: 1 1 2 2 1 1 1 1 1 1 ...
                 2.3 2.3 1.9 1.9 1.2 1.2 0.5 0.5 1.5 1.5 ...
         : num
 $
                 0 0 1 1 0 0 0 0 0 0 ...
          : num
                     0 0 0 0 0 0 0 0 ...
 $
  AN
          : num
                 0 0
                 0 0 0 0 0 0 0 0 0 0 ...
           num
```

This dataset consists of times to the first and second recurrences of infection in 38 kidney patients using a portable dialysis machine. Infections can occur at the location of insertion of the catheter. The catheter is later removed if infection occurs and can be removed for other reasons, in which case the observation is censored (McGilchrist and Aisbett, 1991). The following variables were repeated twice for each patient:

- id: patient number.
- time: time (in days) from insertion of the catheter to infection in kidney patients using portable dialysis machine.
- status: censoring indicator (1: if patient is uncensored; 0: otherwise).
- age: age (in years) of each patient.
- sex: sex of each patient (1: if patient is male; 2: if patient is female).
- disease: disease type (GN; AN; PKD; Other).
- frail: frailty estimate from original paper.
- GN: indicator for disease type GN.

- AN: indicator for disease type AN.
- PKD: indicator for disease type PKD.

Note that the status variable equal to zero in any of the recurrences means that until that moment of observation there was no infection. We would also like to mention that we will use a modelling based on the *clock-reset approach* (Kleinbaum and Klein, 2012), in which time starts at zero again after each recurrence.

6.2 Model implementation

Time, in days, from insertion of a catheter in patient *i*th to infection is modelled trough a proportional hazard specification with multiplicative frailties:

$$h_i(t \mid h_0, \boldsymbol{\beta}, w_i) = w_i h_0(t) \exp\left\{\beta_2 \operatorname{sex}\right\}, \tag{26}$$

where $w_i \sim \text{Gamma}(\psi, \psi)$ represents the frailty term for each individual; $h_0(t) = \lambda \alpha t^{\alpha-1}$ is specified as a Weibull baseline hazard function, where α and $\lambda = \exp(\beta_1)$ are the shape and scale parameters, respectively; and β_2 is the regression coefficient for the sex covariate. As the purpose of this modelling is illustrative, the other covariates will not be considered. We assume prior independence and specify prior marginal based on non-informative distributions commonly employed in the literature. The β 's follow a N(0,0.001), while α and ψ follow a Un(0,10) and a Gamma(0.01,0.01), respectively.

Our variable of interest is time, which represents the time from insertion of the catheter to infection. The status covariate plays an important role in the codification of the survival and censoring times:

```
R> # Number of patients and catheters
R> n <- length(unique(kidney$id))
R> J <- 2
R> # Survival and censoring times
R> time <- kidney$time
R> cens <- time
R> time[kidney$status == 0] <- NA # Censored
R> is.censored <- as.numeric(is.na(time))
R> # Matrix format
R> time <- matrix(time, n, J, byrow = TRUE)
R> cens <- matrix(cens, n, J, byrow = TRUE)
R> is.censored <- matrix(is.censored, n, J, byrow = TRUE)</pre>
```

Without loss of generality, we have created a design matrix X with the sex covariate:

```
R> sex <- kidney$sex[seq(1, 2 * n, 2)] - 1 # Reference = male R> X <- model.matrix(~ sex)
```

Listing 6 shows a generic implementation of a frailty model in BUGS syntax using kidney data.

Listing 6: Frailty model in BUGS syntax (file named as **Frailty.txt**).

```
model {
    for(i in 1:n)
       for(j in 1:J)
         # Survival and censoring times
         is.censored[i,j] ~ dinterval(time[i,j],cens[i,j])
         time[i,j] ~ dweib(alpha,lbd[i,j])
         log(lbd[i,j]) \leftarrow inprod(beta[],X[i,]) + log(w[i])
      # Multiplicative frailties
      w[i] ~ dgamma(psi,psi)
10
11
12
    # Prior distributions
13
    for(1 in 1:Nbetas) \{ beta[1] \sim dnorm(0.0,0.001) \}
14
    alpha ~ dunif(0,10)
15
    psi ~ ^{\sim} ~ dgamma (\, 0\,.\,0\,1\,\,,0\,.\,0\,1\,)
16
17
    # Derived quantity
18
    lambda <- exp(beta[1])
19
20
```

6.3 Model estimation: JAGS from R

Once the variables have been defined, a list with all the elements required in the model is created:

```
R> d.jags <- list(n = n, J = J, time = time, cens = cens, X = X, + is.censored = is.censored, Nbetas = ncol(X))
```

The initial values for each frailty model parameter are passed to JAGS using a function that returns a list of random values:

```
R> i.jags <- function(){ list(beta = rnorm(ncol(X)), alpha = runif(1), psi = runif(1)) }
```

The vector of monitored/saved parameters is:

```
R> p.jags <- c("beta", "alpha", "lambda", "psi", "w")</pre>
```

Next, the JAGS model is compiled:

```
R> library("rjags")
R> m6 <- jags.model(data = d.jags, file = "Frailty.txt", inits = i.jags, n.chains = 3)</pre>
```

We now run the model for 10000 burn-in simulations:

```
R> update(m6, 10000)
```

Finally, the model is run for 100000 additional simulations to keep one in 100 so that a proper thinning is done:

```
R> res <- coda.samples(m6, variable.names = p.jags, n.iter = 100000, thin = 100)
```

Similarly to the first example (Section 2.1), numerical and graphical summaries of the model parameters can be obtained using the summary and densplot functions, respectively. Gelman and Rubin's convergence diagnostic can be calculated with the gelman.diag function, and the traceplot function provides a visual way to inspect sampling behaviour and assesses mixing across chains and convergence.

Next, simulations from the three Markov chains are merged together for inference:

```
R> result <- as.mcmc(do.call(rbind, res))</pre>
```

The posterior samples of each parameter are obtained by:

```
R> alpha <- result[,1]; beta2 <- result[,3]; lambda <- result[,4]
R> psi <- result[,5]; w <- result[,6:ncol(result)]</pre>
```

Table 5 shows posterior summaries for the frailty model parameters using kidney.

Table 5: Posterior summaries for the frailty model parameters.

Parameter	Mean	SD	2.5%	50%	97.5%	$P(\cdot > 0 \mid \text{data})$
β_2 (sex)	-1.908	0.555	-3.064	-1.889	-0.876	0.000
α	1.233	0.167	0.929	1.222	1.592	1.000
λ	0.019	0.012	0.004	0.017	0.050	1.000
ψ	2.417	2.101	0.779	1.878	7.844	1.000

The individual survival curve based on posterior samples is a relevant information in this type of studies. So, we can summarise the posterior distribution of the individual survival curve in a grid of points as follows:

Next, we can differentiate the survival curves by sex (code below). Figure 6 shows such curves for all patients in the *kidney* data.

The curves shown in Figure 6 represent the posterior mean of the probability of infection from any catheter insertion at each time considering the two replicates per patient.

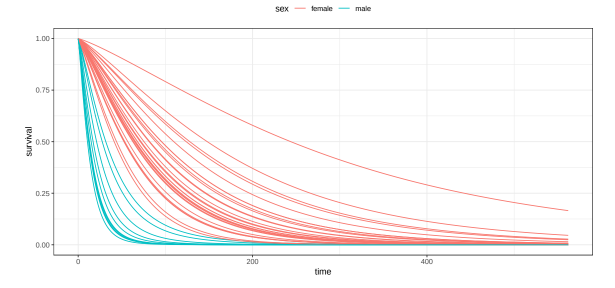


Figure 6: Posterior mean of the individual survival function from the frailty model (26).

7 Joint models of longitudinal and survival data

Joint modelling of longitudinal and time-to-event data is an increasingly productive area of statistical research that examines the association between longitudinal and survival processes (Rizopoulos, 2012). It enhances survival modelling with the inclusion of internal time-dependent covariates as well as longitudinal modelling by allowing for the inclusion of non-ignorable dropout mechanisms through survival tools. Joint models were introduced during the 90s (DeGruttola and Tu, 1994; Tsiatis et al., 1995; Faucett and Thomas, 1996; Wulfsohn and Tsiatis, 1997) and since then, have been applied to a great variety of studies mainly in epidemiological and biomedical areas.

Bayesian joint models assume a full joint distribution for the longitudinal (y) and the survival processes (s) as well as the subject-specific random effects vector (b) and the parameters and hyperparameters (θ) of the model (Armero et al., 2018). Usually, they can be defined as follow:

$$f(y, s, \boldsymbol{b}, \boldsymbol{\theta}) = f(y, s \mid \boldsymbol{b}, \boldsymbol{\theta}) f(\boldsymbol{b} \mid \boldsymbol{\theta}) \pi(\boldsymbol{\theta}), \tag{27}$$

which factorises as the product of the joint conditional distribution $f(y, s \mid b, \theta)$, the conditional distribution $f(b \mid \theta)$ of the random effects, and the prior distribution $\pi(\theta)$. There are different proposals for the specification of the conditional distribution $f(y, s \mid b, \theta)$. The most popular approaches are the share-parameter models and the joint latent class models.

Shared-parameter models are a type of joint models where the longitudinal and time-to-event processes are connected by means of a common set of subject-specific random effects. These models make possible to quantify both the population and individual effects of the underlying longitudinal outcome on the risk of an event and allow to obtain individualised time-dynamic predictions (Wu and Carroll, 1988; Hogan and Laird, 1997, 1998). In particular, this approach postulates conditional independence between the longitudinal and survival processes given the random effects and the parameters:

$$f(y, s \mid \boldsymbol{b}, \boldsymbol{\theta}) = f(y \mid \boldsymbol{b}, \boldsymbol{\theta}) f(s \mid \boldsymbol{b}, \boldsymbol{\theta}).$$

The joint latent class model is based on finite mixtures (Proust-Lima et al., 2014). Heterogeneity among the individuals is classified into a finite number G of homogeneous latent clusters which share the same longitudinal trajectory and the same risk function. Both elements are also conditionally independent within the subsequent latent group as follows:

$$f(y, s \mid L = g, \boldsymbol{b}, \boldsymbol{\theta}) = f(y \mid L = g, \boldsymbol{b}, \boldsymbol{\theta}) f(s \mid L = g, \boldsymbol{\theta}),$$

where L is a random variable that measures the uncertainty on the membership of each individual to each group, usually modelled by means of a multinomial logistic model. These models are possibly the most complex. Predicting observations from models with random effects is not easy. In this case, predicting longitudinal observations will be complicated because the conditional marginal distribution $f(y \mid L = g, b, \theta)$ necessary to obtain the corresponding predictive posterior distribution depends on both random effects and latent groups. On the other hand, the prediction of survival times does seem to be computationally simpler because the conditional distribution $f(s \mid L = g, \theta)$ does not depend on individual random effects.

All these proposals account for a particular type of conditional independence between the longitudinal and the survival processes which facilitates the modelling into longitudinal and survival submodels with various types of connectors. This general structure allows any type of modelling for the survival process such as frailty survival regression models, competing risks with frailties, cure models with frailties as well as linear mixed models or generalised linear mixed models for the longitudinal process (Taylor et al., 2013; Rizopoulos et al., 2015; Armero et al., 2016b; Rué et al., 2017). See Armero (2020) for a short review on Bayesian joint models up to date.

7.1 prothro dataset

We consider a liver cirrhosis dataset, referred to as *prothro* (longitudinal information) and *prothros* (survival information). It is available from the JMbayes package (Rizopoulos, 2019):

```
R> library("JMbayes")
R> data("prothro")
R> data("prothros")
R> str(prothro); str(prothros)
'data.frame': 2968 obs. of 9 variables:
       : num 1 1 1 2 2 2 2 2 2 2 ...
              38 31 27 51 73 90 64 54 58 90 ...
  pro
       : num
  time : num 0 0.244 0.381 0 0.687
  treat: Factor w/ 2 levels "placebo", "prednisone": 2 2 2 2 2 2 2 2 2 ...
  Time : num 0.413 0.413 0.413 6.754 6.754 ...
              0 0.244 0.381 0 0.687 ...
  start: num
              0.244 0.381 0.413 0.687 0.961 ...
  stop : num
 $ death: num
              1 1 1 1 1 1 1 1 1 1 ...
 $ event: num 0 0 1 0 0 0 0 0 0 0 ...
'data.frame': 488 obs. of 4 variables:
        : num 1 2 3 4 5 6 7 8 9 10 ...
 $ Time : num 0.413 6.754 13.394 0.794 0.75 ...
  death: num 1 1 0 1 1 1 1 0 0 1 ...
  treat: Factor w/ 2 levels "placebo", "prednisone": 2 2 2 2 2 1 1 1 1 ...
```

These datasets are part of a placebo-controlled randomised trial on 488 liver cirrhosis patients, where the longitudinal observations of a biomarker (prothrombin) are recorded (Andersen et al., 1993). For our illustrative purpose, only the following variables are relevant:

- id: patient number.
- pro: prothrombin measurements.
- time: time points at which the prothrombin measurements were taken.
- treat: randomised treatment (placebo or prednisone).
- Time: time (in years) from the start of treatment until death or censoring.
- death: censoring indicator (1: if patient is died; 0: otherwise).

7.2 Model implementation

We assume a shared-parameter model specified by a linear mixed-effects model (Laird and Ware, 1982) for the longitudinal response and a Cox model for the survival part. The linear mixed-effects model which describes the subject-specific prothrombin evolution of individual *i* over time is given by:

$$y_i(t) = \beta_{L1} + b_{i1} + (\beta_{L2} + b_{i2})t + \beta_{L3} \text{treat} + \epsilon_i(t), \quad i = 1, \dots, n,$$
 (28)

where $y_i(t)$ represents the prothrombin value at time t for individual i; β_{L1} and β_{L2} are fixed effects for intercept and slope, respectively, with b_{i1} and b_{i2} being the respective individual random effects; β_{L3} is the regression coefficient for the treat covariate; and $\epsilon_i(t)$ is a measurement error for individual i at time t. We assume that the individual random effects, $b_i = (b_{i1}, b_{i2})^{\top}$, given Σ follow a joint bivariate normal distribution with mean vector $(0, 0)^{\top}$ and variance-covariance matrix Σ , and that the errors are conditionally i.i.d. as $(\epsilon_i(t) \mid \sigma^2) \sim N(0, \sigma^2)$, where σ^2 represents the error variance. Random effects and error terms were assumed mutually independent.

Survival time for individual i is modelled from a proportional hazard specification which includes in the exponential term the random effects b_{i1} and b_{i2} in (28) as follows:

$$h_i(t \mid h_0, \beta_S, \gamma, b_i) = h_0(t) \exp\{\beta_{S2} \text{treat} + \gamma(b_{i1} + b_{i2}t)\},$$
 (29)

with $h_0(t) = \lambda \alpha t^{\alpha-1}$ specified as a Weibull baseline hazard function, where α and $\lambda = \exp(\beta_{S1})$ are the shape and scale parameters, respectively; β_{S2} is the regression coefficient for the treat covariate; and γ is an association parameter that measure the strength of the link between the random effects associated to individual i of the longitudinal submodel and their risk of death at time t.

We assume prior independence and specify prior marginal distributions based on non-informative distributions commonly employed in the literature. β_L 's, β_S ' and γ follow a N(0,0.001), while α and σ follow a Un(0,10) and a Un(0,100), respectively, and Σ follows an Inv-Wishart(V, r), where V is a 2×2 identity matrix and r=2 is the degrees-of-freedom parameter. The position of the inverse-Wishart distribution in parameter space is specified by V, while r set the certainty about the prior

information in the scale matrix (Schuurman et al., 2016). The larger the r, the higher the certainty about the information in V, and the more informative is the distribution. Hence, in our application, the least informative specification then results when r=2 (number of random effects), which is the lowest possible number of r. Additionally, V as an identity matrix has the appealing feature that each of the correlations in Σ has, marginally, a uniform prior distribution (Gelman et al., 2013).

Our variable of interest is Time (*prothros* file), which represents the time from the start of treatment until death or censoring. The death=1 variable indicates an uncensored time:

```
R> # Number of patients and number of longitudinal observations per patient
R> n <- nrow(prothros)
R> M <- table(prothro$id)
R> # Survival and censoring times
R> Time <- prothros$Time
R> death <- prothros$death</pre>
```

The (log)prothrombin observations and their respective measurement times (*prothro* file) have been rearranged in matrix format:

We have created the survival design matrix composed of an intercept and a treatment variable:

```
R> treat <- as.numeric(prothros$treat) - 1 # Reference = placebo
R> XS <- model.matrix(~ treat) # Fixed effects</pre>
```

We have split the longitudinal design matrix into two parts, XL (fixed effects) and ZL (random effects):

```
R> XL <- array(1, dim = c(n, max(M), 3)) # Fixed effects R> XL[, , 2] <- time; XL[, , 3] <- treat R> ZL <- array(1, dim = c(n, max(M), 2)) # Random effects R> ZL[, , 2] <- time
```

The survival function for individual i, $S_i(t \mid h_0, \beta_S, \gamma, b_i) = \exp\left\{-\int_0^t h_i(u \mid h_0, \beta_S, \gamma, b_i) du\right\}$ can be efficiently approximated using some Gaussian quadrature method, available from the statmod package (Smyth et al., 2020). For our analysis, we have used 15-point Gauss-Legendre quadrature rule, as is done in Armero et al. (2018):

```
R> # Gauss-Legendre quadrature (15 points)
R> library("statmod")
R> glq <- gauss.quad(15, kind = "legendre")
R> xk <- glq$nodes # Nodes
R> wk <- glq$weights # Weights
R> K <- length(xk) # K-points</pre>
```

Listing 7 shows a generic implementation of a joint model in BUGS syntax using *prothrolprothros* data. Once the variables have been defined, a list with all the elements required in the model is created:

```
R> d.jags <- list(n = n, M = M, Time = Time, XS = XS, log.proth = log.proth, XL = XL, ZL = ZL, 
+ XS = XS, death = death, mub = rep(0, 2), V = diag(1, 2), Nb = 2, zeros = rep(0, n), 
+ NbetasL = dim(XL)[3], NbetasS = ncol(XS), K = length(xk), xk = xk, wk = wk)
```

The variables mub and V represent, respectively, the mean of the random effects normally distributed and the scale matrix of the Wishart distribution which models the precision of the random effects.

The initial values for each joint model parameter are passed to JAGS using a function that returns a list of random values:

```
R> i.jags <- function() {
+    list(betaS = rnorm(ncol(XS)), gamma = rnorm(1), alpha = runif(1),
+    betaL = rnorm(dim(XL)[3]), sigma = runif(1), Omega = diag(runif(2)))
+ }</pre>
```

The vector of monitored/saved parameters is:

```
R> p.jags <- c("betaS", "gamma", "alpha", "lambda", "betaL", "sigma", "Sigma", "b")
```

Next, the JAGS model is compiled:

Listing 7: Joint model in BUGS syntax (file named as JM.txt).

```
model {
    for(i in 1:n)
2
      # Longitudinal observations
3
      for(j in 1:M[i]){
4
          log.proth[i,j] ~ dnorm(mu[i,j],tau)
              mu[i,j] \leftarrow inprod(betaL[],XL[i,j,]) + inprod(b[i,],ZL[i,j,])
7
      # Survival and censoring times
      # Hazard function at integration points
10
      for (j in 1:K)
11
          haz[i,j] <- alpha * pow(Time[i]/2*(xk[j]+1), alpha-1) *
12
13
                       \exp(\inf (beta S[], XS[i,]) + \operatorname{gamma} *(b[i,1] + b[i,2] *(Time[i]/2 *(xk[j]+1))))
14
      # Log-survival function with Gauss-Legendre quadrature
15
      logSurv[i] < -Time[i]/2 * inprod(wk, haz[i,])
16
17
      # Definition of the survival log-likelihood using zeros trick
18
      phi[i] < -100000 - death[i] * log(haz[i,K]) - logSurv[i]
19
      zeros[i] ~ dpois(phi[i])
20
21
      # Random effects
22
      b[i, 1:Nb] ~ dmnorm(mub[], Omega[,])
23
24
25
26
    # Prior distributions
    for(1 in 1:NbetasS) \{ betaS[1] \sim dnorm(0.0,0.001) \}
27
            dnorm (0.0,0.001)
28
    alpha \sim dunif (0,10)
29
    for(1 in 1:NbetasL) \{ betaL[1] \sim dnorm(0.0,0.001) \}
30
    tau <- pow(sigma, -2)
31
    sigma \sim dunif (0,100)
32
    Omega [1:Nb, 1:Nb] dwish (V[,],Nb)
33
    Sigma[1:Nb,1:Nb] < -inverse(Omega[,])
34
35
    # Derived quantity
36
37
    lambda <- exp(betaS[1])
```

```
R> library("rjags")
R> m7 <- jags.model(data = d.jags, file = "JM.txt", inits = i.jags, n.chains = 3)</pre>
```

We now run the model for 1000 burn-in simulations:

```
R> update(m7, 1000)
```

Finally, the model is run for 10000 additional simulations to keep one in 10 so that a proper thinning is done:

```
R> res <- coda.samples(m7, variable.names = p.jags, n.iter = 10000, thin = 10)
```

Similarly to the first example in Section 2.1, numerical and graphical summaries of the model parameters can be obtained using the summary and densplot functions, respectively. Gelman and Rubin's convergence diagnostic can be calculated with the gelman.diag function, and the traceplot function provides a visual way to inspect sampling behaviour and assesses mixing across chains and convergence.

Next, simulations from the three Markov chains are merged together for inference:

```
R> result <- as.mcmc(do.call(rbind, res))</pre>
```

The posterior samples of each parameter are obtained by:

```
R> Sigma2.11 <- result[,1]; Sigma2.12 <- result[,2]; Sigma2.22 <- result[,4]
R> alpha <- result[,5]; b1 <- result[,6:(n+5)]; b2 <- result[,(n+6):(2*n+5)]
R> betaL1 <- result[,(2*n+6)]; betaL2 <- result[,(2*n+7)]; betaL3 <- result[,(2*n+8)]
R> betaS2 <- result[,(2*n+10)]; gamma <- result[,(2*n+11)]
R> lambda <- result[,(2*n+12)]; sigma <- result[,(2*n+13)]</pre>
```

Table 6: Posterior summaries for the joint model parameters.

Parameter	Mean	SD	2.5%	50%	97.5%	$P(\cdot > 0 \mid \text{data})$
β_{S2} (treat)	0.073	0.138	-0.191	0.071	0.343	0.703
γ (assoc)	-2.269	0.180	-2.640	-2.264	-1.923	0.000
λ	0.187	0.023	0.145	0.186	0.233	1.000
α	0.934	0.049	0.841	0.934	1.034	1.000
β_{L1} (intercept)	4.276	0.021	4.235	4.276	4.318	1.000
$eta_{L2}\left(exttt{slope} ight)$	-0.004	0.007	-0.018	-0.004	-0.010	0.301
eta_{L3} (treat)	-0.099	0.030	-0.159	-0.098	-0.040	0.000
σ	0.258	0.004	0.250	0.258	0.265	1.000
Σ_{11}	0.098	0.008	0.083	0.098	0.116	1.000
Σ_{22}	0.013	0.001	0.011	0.013	0.017	1.000
Σ_{12}	-0.003	0.003	-0.009	-0.003	0.003	0.145

Table 6 shows posterior summaries for the joint model parameters using prothro/prothros data.

Most of the parameters are interpreted similarly to the ones in previous examples. However, the association parameter, γ , plays an important role in this type of models. In our illustration, the posterior mean of γ is negative, -2.269, and $P(\gamma > 0 \mid \text{data}) = 0$, indicating a strong negative association of the prothrombin measurements with respect to vital status. In other words, a negative value for γ means that low values or decreasing trends of prothrombin increase the risk of death.

8 Conclusions

The analysis of time until an event of interest requires a suitable and flexible modelling and has applications in several applied fields. The BUGS language offers the opportunity of easily use and adapt Bayesian hierarchical models without the need to manually implement Markov chain Monte Carlo methods. Hence, this paper has summarised some of the most popular survival models and has focused on the Bayesian paradigm to make the inferential procedure. Furthermore, for each of the models proposed we have provided the codes in BUGS syntax, so that model can be fit with the support of the rjags package from the R language.

We have discussed all the implementation details of the following Bayesian survival models: accelerated failure time, proportional hazards, mixture cure, competing risks, multi-state, frailty, and joint models of longitudinal and survival data. Moreover, the computation of quantities of interest derived from posterior samples has been provided as well as some graphs that assist in the interpretation of results and decision making. The paper has also briefly presented other Bayesian R-packages that handle time-to-event data.

In conclusion, we hope this paper will encourage researchers who use survival models make their analyses based on the Bayesian paradigm from the BUGS codes we have provided and easily adapt them to novel settings. In addition, the descriptions of the survival models provided herein could also be used as a guidance to implement these models using other similar languages such as, for example, Stan (Carpenter et al., 2017).

Acknowledgments

This work was partially supported by FONDECYT (Chile), Grant Number: 11190018; Consejería de Educación, Cultura y Deportes (JCCM, Spain), Grant Number: PPIC-2014-001-P; FEDER, Grant Number: SBPLY/17/180501/000491; and Ministerio de Ciencia e Innovación (MCI, Spain), Grant Number: PID2019-106341GB-I00.

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A Implementation of the transition probabilities of the semi-Markov multi-state model based on posterior samples

Figure 5 was generated from the transition probabilities (16)–(21) based on posterior samples of the illness-death model parameter (22)–(24). In particular, we set a grid of time values to evaluate the transition probabilities and used the median values of age and year covariates. In this example, we are interested in the transition probabilities for patients with and without prior bypass surgery (i.e., surgery equal to 1 or 0, respectively):

```
R> grid <- seq(0, 1000, length.out = 51)
R> \times0 <- c(median(heart2$age), median(heart2$year), 0)
R> \times1 <- c(median(heart2$age), median(heart2$year), 1)
```

For our Weibull baseline hazard specification, the transition probability $p_{11}(s, t \mid \theta)$ (see Equation 16) can be calculated analytically and is expressed by:

$$p_{11}(s,t\mid\boldsymbol{\theta}) = \exp\left\{-\lambda_1\eta_1[t^{\alpha_1} - s^{\alpha_1}] - \lambda_3\eta_3[t^{\alpha_3} - s^{\alpha_3}]\right\},\tag{30}$$

where $\eta_k = \exp(\mathbf{x}^{\top} \boldsymbol{\beta}_k)$, for k = 1, 3. Equation (30) is implemented as follows:

```
R> p11_s_t <- function(tt, ss, l1, a1, b1, l2, a2, b2, x) {
        H1 <- l1 * exp(sum(b1 * x)) * (tt^al - ss^al)
        H2 <- l2 * exp(sum(b2 * x)) * (tt^a2 - ss^a2)
        return(exp(-H1 - H2))
        + }</pre>
```

Next, we have created a function mcmc_p11_s_t that takes posterior samples of each parameter from JAGS and computes $p_{11}(s, t \mid \theta)$ in (30) for a grid of time values (variable grid) using covariates x (here x0 or x1). Note that mcmc_p11_s_t is based on the mclapply function, available from the parallel package to speed computations up (R Core Team, 2020).

As previously mentioned, we are interested in the transition probabilities for patients with and without prior bypass surgery. In particular, $p_{11}(s, t \mid \theta)$ for s = 0 can be calculated as follows:

Finally, these transition probabilities for x0 and x1 are plotted in Figure 5 using the following code:

The transition probability $p_{22}(s, t \mid \theta)$ (see Equation 17) using the Weibull baseline hazard specification is written as:

$$p_{22}(s,t \mid \boldsymbol{\theta}) = \frac{\lambda_1 \alpha_1 \eta_1}{1 - \exp\left\{-\lambda_1 \eta_1 s^{\alpha_1}\right\}} \int_0^s u^{\alpha_1 - 1} \exp\left\{-\lambda_1 \eta_1 u^{\alpha_1} - \lambda_3 \eta_3 \left[(t - u)^{\alpha_3} - (s - u)^{\alpha_3} \right] \right\} du, \tag{31}$$

where $\eta_k = \exp(\mathbf{x}^{\top} \boldsymbol{\beta}_k)$, for k = 1, 3. The integral in (31) has no closed form, so some approximate method of integration is required. To do this, we first have created a function int_p22 to calculate this integral:

```
R> int_p22 <- function(u, tt, ss, l1, a1, b1, l3, a3, b3, x){
+ F1 <- 1 - exp(-l1 * exp(sum(b1 * x)) * ss^al)
+ u1 <- ifelse(u > 0, u^(a1 - 1), 0)
+ h1 <- l1 * a1 * exp(sum(b1 * x)) * u1
+ H1 <- l1 * exp(sum(b1 * x)) * u^a1
+ H3 <- l3 * exp(sum(b3 * x)) * ((tt - u)^a3 - (ss - u)^a3)
+ return(h1 * exp(-H1 - H3) / F1)
+ }</pre>
```

Next, we have created a function p22_s_t that computes $p_{22}(s, t \mid \theta)$ in (31) by integrating out the int_p22 function using the integral function available from the pracma package (Borchers, 2019).

Analogous to the previous case, we have created a function mcmc_p22_s_t that takes posterior samples of each parameter from JAGS and computes $p_{22}(s, t \mid \theta)$ in (31) for a grid of time values (variable grid) using covariates x (here x0 or x1).

Note that the transition time from state 1 to state 2 prevents the case s=0 in $p_{22}(s,t\mid \boldsymbol{\theta})$. To realistically get around this problem, we have set s as the median time of patients who transitioned from state 1 to 2 (s=26) and add this amount to the grid of time values. So, we have calculated the transition probabilities (31) for x0 and x1:

Finally, these transition probabilities are plotted in Figure 5 using the following code:

The transition probability $p_{12}(s, t \mid \theta)$ (see Equation 18) using the Weibull baseline hazard specification is written as:

$$p_{12}(s,t \mid \boldsymbol{\theta}) = \lambda_1 \alpha_1 \eta_1 \int_s^t u^{\alpha_1 - 1} \exp\left\{ -\lambda_1 \eta_1 \left[u^{\alpha_1} - s^{\alpha_1} \right] - \lambda_2 \eta_2 \left[u^{\alpha_2} - s^{\alpha_2} \right] - \lambda_3 \eta_3 (t - u)^{\alpha_3} \right\} du, \tag{32}$$

where $\eta_k = \exp(x^\top \beta_k)$, for k = 1, 2, 3. Analogous to the previous case, the integral in (32) has no closed form, so some approximate method of integration is required. To do this, we first have created a function int_p12 to calculate this integral:

```
R> int_p12 <- function(u, tt, ss, l1, a1, b1, l2, a2, b2, l3, a3, b3, x){
+    u1 <- ifelse(u > 0, u^(a1 - 1), 0)
+    h1 <- l1 * a1 * exp(sum(b1 * x)) * u1
+    H1 <- l1 * exp(sum(b1 * x)) * (u^a1 - ss^a1)
+    H2 <- l2 * exp(sum(b2 * x)) * (u^a2 - ss^a2)
+    H3 <- l3 * exp(sum(b3 * x)) * (tt - u)^a3
+    return(h1 * exp(-H1 - H2 - H3))
+ }</pre>
```

Next, we have created a function p12_s_t that computes $p_{12}(s, t \mid \theta)$ in (32) by integrating out the int_p12 function.

As a last step, we have created a function mcmc_p12_s_t that takes posterior samples of each parameter from JAGS and computes $p_{12}(s, t \mid \theta)$ in (32) for a grid of time values (variable grid) using covariates x (here x0 or x1).

```
R> mcmc_p12_s_t <- function(t.pred, s.pred, l1, a1, b1, l2, a2, b2, l3, a3, b3, x){
    # Sub-sample to speed up computations
    samples.idx <- sample(1:length(l1), 200)
    res <- sapply(t.pred, function(tt){
        aux <- mclapply(samples.idx, function(i){
            p12_s_t(tt, ss = s.pred, l1 = l1[i], a1 = a1[i], b1 = b1[i,], l2 = l2[i],
            a2 = a2[i], b2 = b2[i,], l3 = l3[i], a3 = a3[i], b3 = b3[i,], x = x)
    })
    return(mean(unlist(aux)))
    })
    return(res)
}</pre>
```

The transition probabilities $p_{12}(s, t \mid \boldsymbol{\theta})$ of x0 and x1 for s = 0 can be calculated as follows:

Finally, these transition probabilities are plotted in Figure 5 using the following code:

As shown in (19) and (20), $p_{13}(s, t \mid \boldsymbol{\theta})$ and $p_{23}(s, t \mid \boldsymbol{\theta})$ are derived from the previously calculated transition probabilities. So, we can get them for $x \in \mathbb{R}$ 0 and $x \in \mathbb{R}$ 1 as follows:

```
R> p13_0_t_surgery0 <- 1 - p11_0_t_surgery0 - p12_0_t_surgery0
R> p13_0_t_surgery1 <- 1 - p11_0_t_surgery1 - p12_0_t_surgery1
R> p23_m_t_surgery0 <- 1 - p22_m_t_surgery0
R> p23_m_t_surgery1 <- 1 - p22_m_t_surgery1</pre>
```

These transition probabilities are plotted in Figure 5 using the following code:

To show these graphs in two rows and three columns, as in Figure 5, we have used the grid.arrange function available from the gridExtra package (Auguie and Antonov, 2017).