

# Comparing methods for time varying logistic models

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## Introduction

In this vignette, we compare the dynamic logistic model in **dynamichazard** with others methods within the package and methods from the **timereg** and **mgcv** packages. Further this note will serve as an illustration of how to use the **ddhazard** function for the logistic model. We will use the **pbc2** dataset from the **survival** package. The motivation is that the **pbc2** data set is commonly used in survival analysis for illustrations. It is suggested to first read or skim `vignette("ddhazard", "dynamichazard")` to get an introduction to the models and estimation methods in this package.

The note is structured as follows: First, we cover the **pbc2** data set. Then we estimate two static (non-dynamic) logistic regression models using **glm**. This is followed by a fit using a Generalized Additive model with the **gam** function in the **mgcv** package. Next, we will estimate a cox-model with time varying parameters using the **timecox** function in the **timereg** package. Finally, we will end by illustrating the methods in this package for time varying parameters in a logistic regression.

You can install the version of the library used to make this vignettes from github with the **devtools** library as follows:

```
current_version # The string you need to pass devtools::install_github

## [1] "boennecd/dynamichazard@6937596963055ebaddf0baecfb0dd10f065d31c6"

devtools::install_github(current_version)
```

You can also get the latest version on CRAN by calling:

```
install.packages("dynamichazard")
```

## The pbc data set

The **pbc** data set contains data from the Mayo Clinic trial on primary biliary cirrhosis. We use this dataset to compare with results previously found analyzing the data set. We will focus on a the subset of co-variables used in Martinussen & Scheike (2007). The dataset can be created by:

```
# PBC data set from survival with time varying covariates
# Details of tmerge are not important in this scope. The code is included
# to make you able to reproduce the results
# See: https://cran.r-project.org/web/packages/survival/vignettes/timedep.pdf
library(survival)
temp <- subset(pbc, id <= 312, select=c(id, sex, time, status, edema, age))
pbc2 <- tmerge(temp, temp, id=id, death = event(time, status))
pbc2 <- tmerge(pbc2, pbcseq, id=id, albumin = tdc(day, albumin),
              protime = tdc(day, protime), bili = tdc(day, bili))
pbc2 <- pbc2[, c("id", "tstart", "tstop", "death", "sex", "edema",
                "age", "albumin", "protime", "bili")]
```

as described in the vignette *Using Time Dependent Covariates and Time Dependent Coefficients in the Cox Model* in the **survival** package. The resulting data frame is structured as follows:

```
head(pbc2)
```

```
##   id tstart tstop death sex edema  age albumin protime bili
## 1  1      0   192     0  f     1 58.8    2.60    12.2 14.5
## 2  1   192   400     2  f     1 58.8    2.94    11.2 21.3
## 3  2      0   182     0  f     0 56.4    4.14    10.6  1.1
## 4  2   182   365     0  f     0 56.4    3.60    11.0  0.8
## 5  2   365   768     0  f     0 56.4    3.55    11.6  1.0
## 6  2   768  1790     0  f     0 56.4    3.92    10.6  1.9
```

The data set is in the usual start and stop time format used in survival analysis. Each individual in the trial has one or more observations. The `id` column identifies the individual. The `tstart` column indicates when the row is valid from and the `tstop` column indicates when the row is valid to. The `death` column is the outcome which is 2 when the individual dies at `tstop` (1 indicates that the individual gets a transplant). The `sex`, `edema` and `age` are baseline variables while `albumin`, `protime` and `bili` are laboratory values updated at follow ups with the patient. As an example, we can look individual 282:

```
(ex <- pbc2[pbc2$id == 282, ])
```

```
##      id tstart tstop death sex edema  age albumin protime bili
## 1708 282      0   258     0  f     0 33.9    3.52     9.5  1.3
## 1709 282   258   370     0  f     0 33.9    3.39     9.5  1.1
## 1710 282   370   744     0  f     0 33.9    3.28    10.1  1.2
## 1711 282   744  1455     0  f     0 33.9    3.04    10.2  1.3
```

She (`sex` is `f`) has had four laboratory values measured at time 0, 258, 370 and 744. Further, she does not die as the `death` column is zero in the final row.

## Logistic model

We can start of with a simple logistic model where we ignore `tstart` and `tstop` variable using `glm`:

```
glm_simple <- glm(death == 2 ~ age + edema + log(albumin) + log(protime) +
                  log(bili), binomial, pbc2)
```

```
glm_simple$coefficients
```

```
## (Intercept)          age          edema log(albumin) log(protime)    log(bili)
##      -9.3834      0.0537      1.4354      -4.4379       2.9274       1.1925
```

Though, we want to account for the fact that say the second the row of individual 282 has a length of 112 days (370 minus 258) while the fourth row has a length 711 days. A way to incorporate this information is to bin the observations into periods of a given length. Example of such binning is found in Tutz & Schmid (2016) and Shumway (2001).

Say that we set the bin interval lengths to 100 days. Then the first row for `id = 282` will yield three observation: one from 0 to 100, one from 100 to 200 and one from 200 to 300 (since we know that she survives beyond time 300). That is, she survives from time 0 to time 100, survives from time 100 to time 200 etc. We can make such a data frame with the `get_survival_case_weights_and_data` function in this package:

```
library(dynamichazard)
pbc2_big_frame <- get_survival_case_weights_and_data(
  Surv(tstart, tstop, death == 2) ~ age + edema + log(albumin) + log(protime) +
    log(bili), data = pbc2, id = pbc2$id, by = 100, max_T = 3600,
  use_weights = F)
pbc2_big_frame <- pbc2_big_frame$X
```

The code above uses the `Surv` function on the left hand side of the formula. The `Surv` function needs a start time, a stop time and an outcome. The right hand side is as before. The `by` argument specifies the interval length (here 100 days) and the `max_T` specify the last time we want to include. We will comment on `use_weights` argument shortly. As an example, we can look at individual 282 in the new data frame:

```
pb2_big_frame[pb2_big_frame$id == 282, ]
```

```
##      id tstart tstop death sex edema  age albumin protime bili  Y  t
## 282 282      0   258     0  f     0 33.9   3.52    9.5  1.3 FALSE 100
## 590 282      0   258     0  f     0 33.9   3.52    9.5  1.3 FALSE 200
## 703 282      0   258     0  f     0 33.9   3.52    9.5  1.3 FALSE 300
## 1207 282    258   370     0  f     0 33.9   3.39    9.5  1.1 FALSE 400
## 1410 282    370   744     0  f     0 33.9   3.28   10.1  1.2 FALSE 500
## 1666 282    370   744     0  f     0 33.9   3.28   10.1  1.2 FALSE 600
## 1938 282    370   744     0  f     0 33.9   3.28   10.1  1.2 FALSE 700
## 2199 282    370   744     0  f     0 33.9   3.28   10.1  1.2 FALSE 800
## 2550 282    744  1455     0  f     0 33.9   3.04   10.2  1.3 FALSE 900
## 2782 282    744  1455     0  f     0 33.9   3.04   10.2  1.3 FALSE 1000
## 3016 282    744  1455     0  f     0 33.9   3.04   10.2  1.3 FALSE 1100
## 3195 282    744  1455     0  f     0 33.9   3.04   10.2  1.3 FALSE 1200
## 3405 282    744  1455     0  f     0 33.9   3.04   10.2  1.3 FALSE 1300
## 3621 282    744  1455     0  f     0 33.9   3.04   10.2  1.3 FALSE 1400
##      weights
## 282      1
## 590      1
## 703      1
## 1207     1
## 1410     1
## 1666     1
## 1938     1
## 2199     1
## 2550     1
## 2782     1
## 3016     1
## 3195     1
## 3405     1
## 3621     1
```

Notice that three new columns have been added: `Y` which is the outcome, `t` which is the stop time in the bin and `weights` which is the weight to be used in a regression. We see that the first row in the initial data frame for individual 282 has three rows now since the row is in three bins (the bins at times (0,100], (100,200] and (200,300]). We could just use weights instead. This is what we get if we set `use_weights = T`:

```
pb2_small_frame <- get_survival_case_weights_and_data(
  Surv(tstart, tstop, death == 2) ~ age + edema + log(albumin) + log(protime) +
    log(bili), data = pb2, id = pb2$id, by = 100, max_T = 3600,
  use_weights = T)
pb2_small_frame <- pb2_small_frame$X
```

The new data rows for individual 282 looks as follows:

```
pb2_small_frame[pb2_small_frame$id == 282, ]
```

```
##      Y  id tstart tstop death sex edema  age albumin protime bili weights
## 1557 0 282      0   258     0  f     0 33.9   3.52    9.5  1.3      3
## 1558 0 282    258   370     0  f     0 33.9   3.39    9.5  1.1      1
## 1559 0 282    370   744     0  f     0 33.9   3.28   10.1  1.2      4
```

```
## 1560 0 282    744 1455    0  f    0 33.9    3.04    10.2 1.3    6
```

Further, individuals who do die are treated a bit differently. For instance, take individual 268:

```
pb2[pb2$id == 268, ] # the original data
```

```
##      id tstart tstop death sex edema  age albumin protime bili
## 1652 268     0   164     0  f   0.5 55.4    2.75    11  6.4
## 1653 268    164  1191     2  f   0.5 55.4    2.78    11  6.8
```

```
pb2_small_frame[pb2_small_frame$id == 268, ] # new data
```

```
##      Y  id tstart tstop death sex edema  age albumin protime bili weights
## 1506 0 268     0   164     0  f   0.5 55.4    2.75    11  6.4         2
## 1507 0 268    164  1191     2  f   0.5 55.4    2.78    11  6.8         9
## 1707 1 268    164  1191     2  f   0.5 55.4    2.78    11  6.8         1
```

Notice, that we have to add an additional row with weight 1 where  $Y = 1$  as it would be wrong to give a weight of 10 to a the row with  $Y = 1$ . She survives for 11 bins and dies in the 12th bin. Finally, we can fit the model with the `glm` function using either of the two data frames as follows:

```
glm_fit_big <- glm(Y ~ age + edema + log(albumin) + log(protime) +
                  log(bili), binomial, pb2_big_frame)
glm_fit_small <- glm(Y ~ age + edema + log(albumin) + log(protime) +
                    log(bili), binomial, pb2_small_frame,
                    weights = pb2_small_frame$weights) # <-- we use weights
```

We can confirm that the two models give the same estimate:

```
all.equal(glm_fit_big$coefficients, glm_fit_small$coefficients)
```

```
## [1] TRUE
```

Further, the binning affects the estimates as shown below. In particular, it affects the estimates for `edema` and `log(albumin)`. The standard errors from the simple fit are also printed. However, these standard errors do not account for the dependence as we use multiple observations from the each individual.

```
knitr::kable(
  rbind("glm with bins" = glm_fit_big$coefficients,
        "glm without bins" = glm_simple$coefficients,
        "Sds with bins" =
          summary(glm_fit_big)[["coefficients"]][, "Std. Error"],
        "Sds without bins" =
          summary(glm_simple)[["coefficients"]][, "Std. Error"]),
  row.names = T)
```

	(Intercept)	age	edema	log(albumin)	log(protime)	log(bili)
glm with bins	-10.38	0.045	1.019	-3.781	2.937	1.057
glm without bins	-9.38	0.054	1.435	-4.438	2.927	1.193
Sds with bins	2.18	0.010	0.284	0.554	0.777	0.110
Sds without bins	2.25	0.012	0.370	0.624	0.815	0.129

To end this section, you can skip making data frame with `get_survival_case_weights_and_data` by calling the `static_glm` function from `dynamichazard` package. The call below yields the same results as shown:

```
static_glm_fit <- static_glm(
  Surv(tstart, tstop, death == 2) ~ age + edema + log(albumin) + log(protime) +
  log(bili), data = pb2, id = pb2$id, by = 100, max_T = 3600)
```

```
all.equal(static_glm_fit$coefficients, glm_fit_big$coefficients)
```

```
## [1] TRUE
```

For details, see the help file by typing `?static_glm`.

## Generalized Additive Models using mgcv

The first method we will compare with is Generalized Additive Models (GAM) by using the `gam` function in the `mgcv` package. The model we fit is of the form:

$$\begin{aligned} \text{logit}(\pi_i) = & \gamma_{\text{time}} f_{\text{time}}(t_i) + \gamma_{\text{age}} f_{\text{time}}(t_i) a_i + \gamma_{\text{ede}} f_{\text{ede}}(t_i) e_i + \gamma_{\text{alb}} f_{\text{alb}}(t_i) \log a_{it} \\ & + \gamma_{\text{pro}} f_{\text{alb}}(t_i) \log p_{it} + \gamma_{\text{bili}} f_{\text{bili}}(t_i) \log b_{it} \end{aligned}$$

where  $\pi_{it}$  is the probability that the  $i$ 'th individual dies of cancer,  $t$  is the stop time of the bin and  $a_i$ ,  $e_i$ ,  $a_{it}$ ,  $p_{it}$  and  $b_{it}$  are respectively the  $i$ 'th individual's age, edema, albumin, protime and bilirubin. The extra subscript  $t$  is added to refer to the level of the covariate in the bin at time  $t$ . It is important to notice that we will use the same bins as shown previously. In particular, we will use the `pb2_big_frame` data frame from before.  $f$  are a smooth function. We will use cubic regression splines with knots spread evenly through the covariate values. We fit the model with the following call:

```
library(mgcv, quietly = T)
spline_fit <- gam(
  formula = Y ~
    # cr is cubic spline with k knots
    s(t, bs = "cr", k = 3, by = age) +
    s(t, bs = "cr", k = 3, by = edema) +
    s(t, bs = "cr", k = 5, by = log(albumin)) +
    s(t, bs = "cr", k = 5, by = log(protime)) +
    s(t, bs = "cr", k = 5, by = log(bili)),
  family = binomial, data = pb2_big_frame)
```

The above estimates the GAM model where the likelihood is penalized by a L2 norm for each of the spline functions. The tuning parameters are chosen by generalized cross validation. Below are plots of the estimates:

```
plot(spline_fit, scale = 0, page = 1, rug = F)
```

Further, we compare the result with static model. Recall that our static model had the following estimates:

```
glm_fit_big$coefficients
```

```
## (Intercept)      age      edema log(albumin) log(protime)  log(bili)
##      -10.384      0.045      1.019      -3.781       2.937       1.057
```

These do seem to correspond with the plots. Further, the intercept in the spline model is:

```
spline_fit$coefficients["(Intercept)"]
```

```
## (Intercept)
##      -10.4
```

which again seems to match. The plot suggest that there may be time varying effects for `bili` particularly

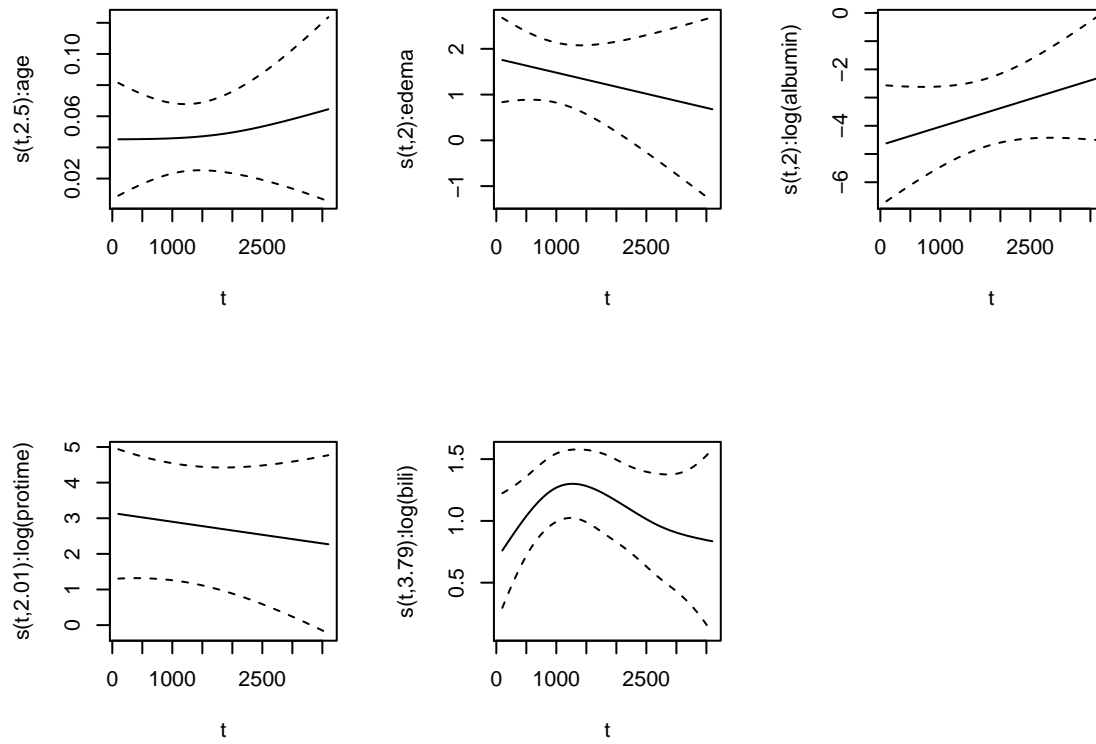


Figure 1: Plots of estimated effects in the GAM model. The effective degree of freedom is noted in the parentheses on the y-axis and is computed given the number knots and final tuning parameter for spline function. For instance,  $s(t, 2.43): \text{age}$  means that the effective degrees of freedom for **age** is 2.43.

## Time varying cox model from `timereg`

Another method we can try is a time varying effects Cox model. We will use the Cox model from the package `timereg` based on Martinussen & Scheike (2007). The model differs from a regular Cox model (e.g. by using the `coxph` function from the `survival` package) by replacing the coefficient  $\beta$  by  $\beta(t)$  such that the instantaneous hazard is:

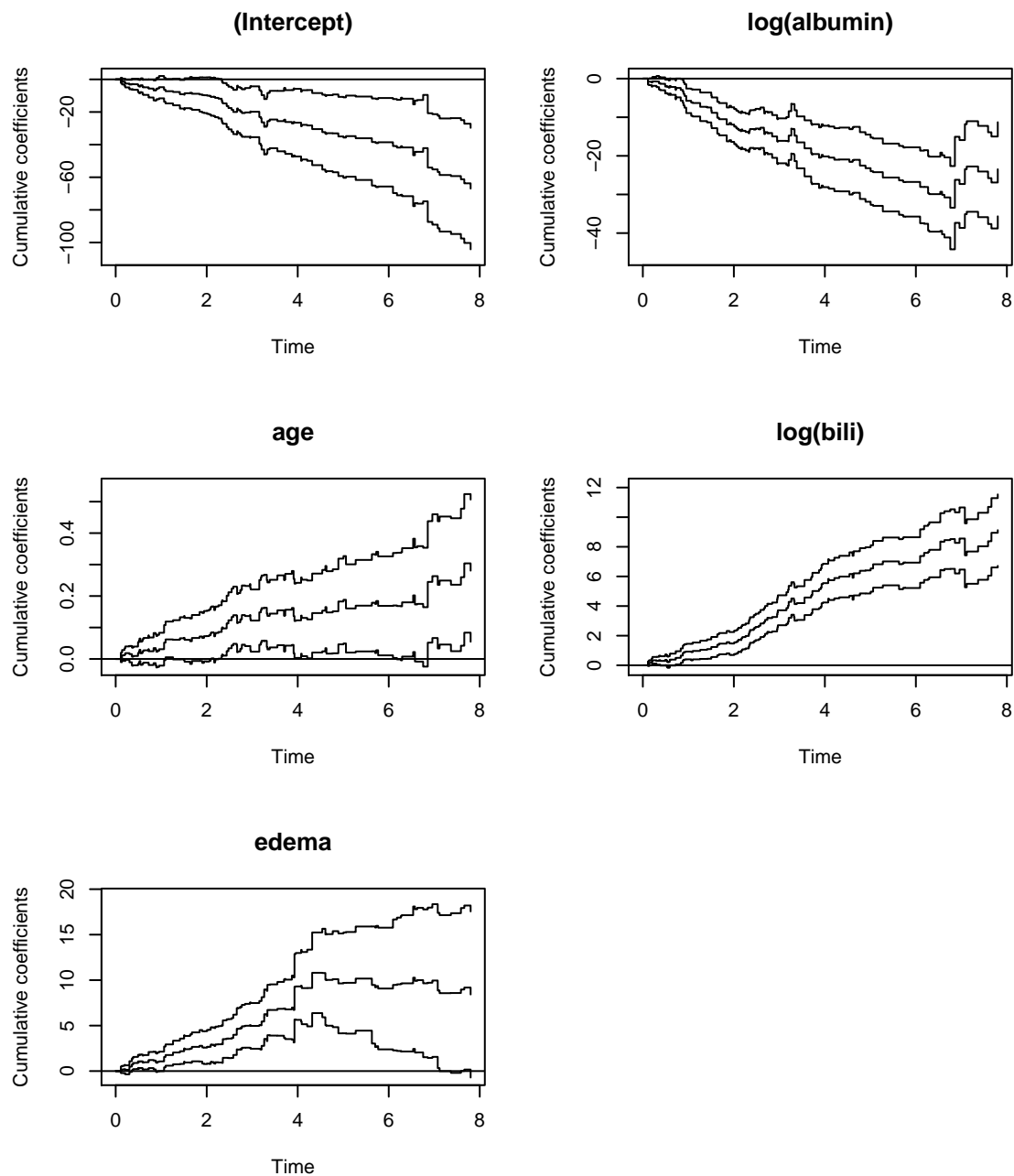
$$\lambda(t) = \lambda_0(t) \exp(\mathbf{x}\beta(t))$$

where each element of  $\beta(t)$  is estimated recursively with an update equation. The update equation is simplified through a first order Taylor expansion of the score function and by adding a smoothness by using weighting depending on time changes with a uniform continuous kernel. For details see Martinussen & Scheike (2007) in chapter 6. The estimation method differs from other alternatives in R that use splines for the time varying effects. The baseline is  $\lambda_0(t) = \exp(\alpha_0(t))$  where  $\alpha_0(t)$  is estimated in a similar way to  $\beta(t)$ . Below we estimate a model similar to the previously fitted models.

We set the last observation time (`max.time`) lower than in the previous model as there are issues with convergence if we do not. For the same reason we specify the effect of `log(protime)` to be constant (non-time varying). The cumulated coefficients  $B(t) = \int_0^t \beta(t)dt$  are plotted to inspect the fitted model. Thus, a constant effect should be roughly linear.

```
library(timereg)
cox_fit<- timecox(Surv(tstart / 365, tstop / 365, death == 2) ~ age + edema +
                  log(albumin) + const(log(protime)) + log(bili), pbc2,
                  start.time=0,
                  max.time = 3000 / 365, # <-- decreased
                  id = pbc2$id, bandwidth = 0.35)

par(mfcol = c(3, 2))
plot(cox_fit)
```



The timecox packages provides two test for whether the coefficient is time varying or not:

```
summary(cox_fit)
```

```
## Multiplicative Hazard Model
##
## Test for nonparametric terms
##
## Test for non-significant effects
##      Supremum-test of significance p-value H_0: B(t)=0
## (Intercept)          5.53          0.000
## age                  3.37          0.012
```

```

## edema                4.65                0.000
## log(albumin)         6.16                0.000
## log(bili)            8.53                0.000
##
## Test for time invariant effects
##           Kolmogorov-Smirnov test p-value H_0:constant effect
## (Intercept)         15.9000                0.218
## age                 0.0785                0.802
## edema               6.1500                0.034
## log(albumin)        13.1000                0.200
## log(bili)           1.0700                0.688
##           Cramer von Mises test p-value H_0:constant effect
## (Intercept)         3.61e+02                0.326
## age                 5.44e-03                0.922
## edema               5.80e+01                0.073
## log(albumin)        3.72e+02                0.214
## log(bili)           2.28e+00                0.585
##
## Parametric terms :
##           Coef.      SE Robust SE      z  P-val lower2.5% upper97.5%
## const(log(protime)) 2.43 0.815      0.966 2.51 0.0121      0.833      4.03
##
## Call:
## timecox(formula = Surv(tstart/365, tstop/365, death == 2) ~ age +
##           edema + log(albumin) + const(log(protime)) + log(bili), data = pbc2,
##           start.time = 0, max.time = 3000/365, id = pbc2$id, bandwidth = 0.35)

```

The above test suggest that only `edema` might be “border line” time-varying.

## Dynamic hazard model

In this section, we will cover the dynamic hazard model with the logistic link function that is implemented in this package. The model is estimated with an EM-algorithm which are from Fahrmeir (1994) and Fahrmeir (1992) when an Extended Kalman Filter is used in the E-step are. Firstly, we will briefly cover the model. See `vignette("ddhazard", "dynamichazard")` for a more detailed explanation of the models. Secondly, we will turn to different ways of designing the model and fitting the model. The idea is that we discretize the outcomes into  $1, 2, \dots, d$  bins. In each bin, we observe whether the individual dies or survives. The state space model we are applying is of the form:

$$\begin{aligned}
\mathbf{y}_t &= \mathbf{z}_t(\boldsymbol{\alpha}_t) + \boldsymbol{\epsilon}_t & \boldsymbol{\epsilon}_t &\sim (\mathbf{0}, \text{Var}(\mathbf{y}_t|\boldsymbol{\alpha}_t)) \\
\boldsymbol{\alpha}_{t+1} &= \mathbf{F}\boldsymbol{\alpha}_t + \mathbf{R}\boldsymbol{\eta}_t & \boldsymbol{\eta}_t &\sim N(\mathbf{0}, \psi_t \mathbf{Q})
\end{aligned}
, \quad t = 1, \dots, d$$

where  $y_{it} \in \{0, 1\}$  is an indicator for whether the  $i$ 'th individual dies in interval  $t$ .  $\dots \sim (a, b)$  denotes a random variable with mean (or mean vector)  $a$  and variance (or co-variance matrix)  $b$ . It needs not to be normally distributed.  $z_{it}(\boldsymbol{\alpha}_t) = h(\boldsymbol{\alpha}_t \mathbf{x}_{it})$  is the non-linear map from state space variables to mean outcomes where  $h$  is the inverse link function. We use the logit model in this note. Thus,  $h(x) = \text{logit}^{-1}(x)$ . The current implementation supports first and second order random walk for the state equation. Further, we define the conditional covariance matrix in the observational equation as  $\mathbf{H}_t(\boldsymbol{\alpha}_t) = \text{Var}(\mathbf{y}_t|\boldsymbol{\alpha}_t)$ .  $\psi_t$  is the length of the bin and is assumed equal for all values of  $t$ .

The unknown parameters are the starting value  $\boldsymbol{\alpha}_0$  and co-variance matrix  $\mathbf{Q}$ . These are estimated with an EM-algorithm. The E-step is calculated using a Extended Kalman filter (EKF), Unscented Kalman filter (UKF) or sequential approximation of the modes. All are followed by a smoothing step. The result is

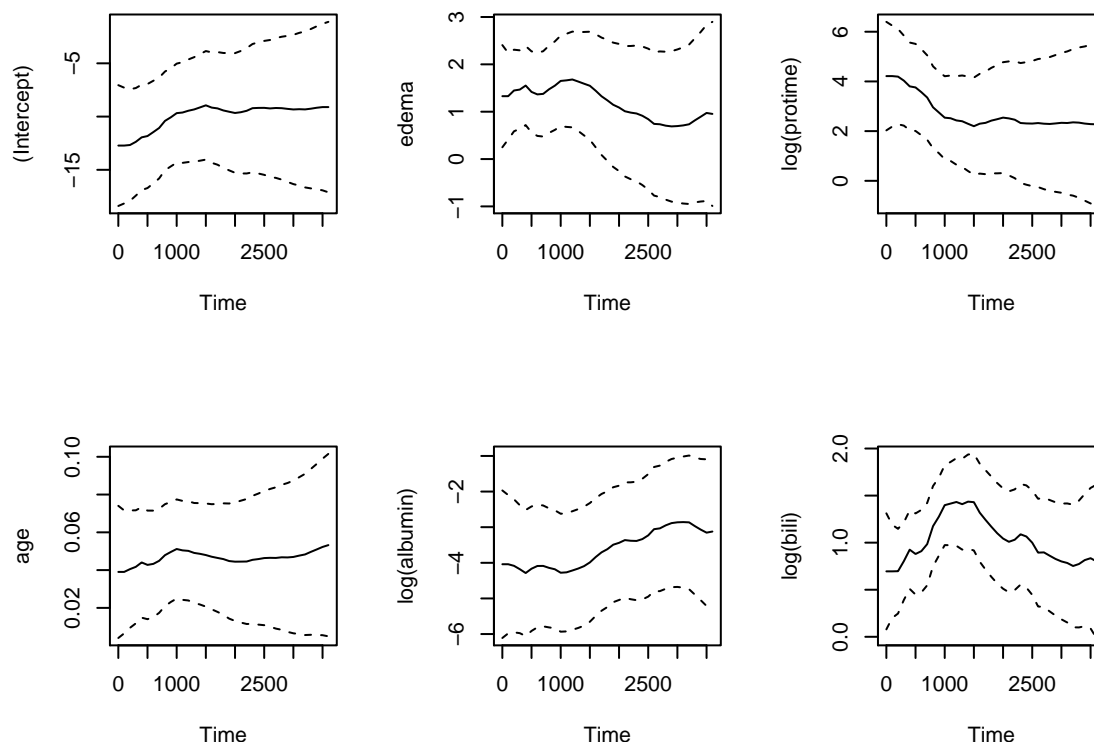
smoothed predictions of  $\alpha_1, \dots, \alpha_d$ , smoothed co-variance matrix and smoothed correlation matrices that we need for the M-step to update  $\alpha_0$  and  $\mathbf{Q}$ .

## Estimation with Extended Kalman Filter

We start by estimating the model using the EKF where we let all coefficients follow a first order random walk:

```
library(dynamichazard)
dd_fit <- ddhazard(Surv(tstart, tstop, death == 2) ~ age + edema +
                  log(albumin) + log(protime) + log(bili), pbc2,
                  id = pbc2$id, by = 100, max_T = 3600,
                  Q_0 = diag(100, 6), Q = diag(0.01, 6),
                  control = list(eps = .001))

plot(dd_fit)
```



The arguments  $\mathbf{Q}_0$  and  $\mathbf{Q}$  corresponds to the co-variance matrix at time zero and the co-variance matrix in the state equation.  $\mathbf{Q}_0$  will remain fixed while  $\mathbf{Q}$  is the starting value in the first iteration of the EM algorithm after which we update  $\mathbf{Q}$ .

Next, we plot the coefficient. That is, we plot the predicted latent variables  $\alpha_0, \dots, \alpha_d$ . Notice that the predicted coefficient are close to the estimates we saw previously for the GAM model.

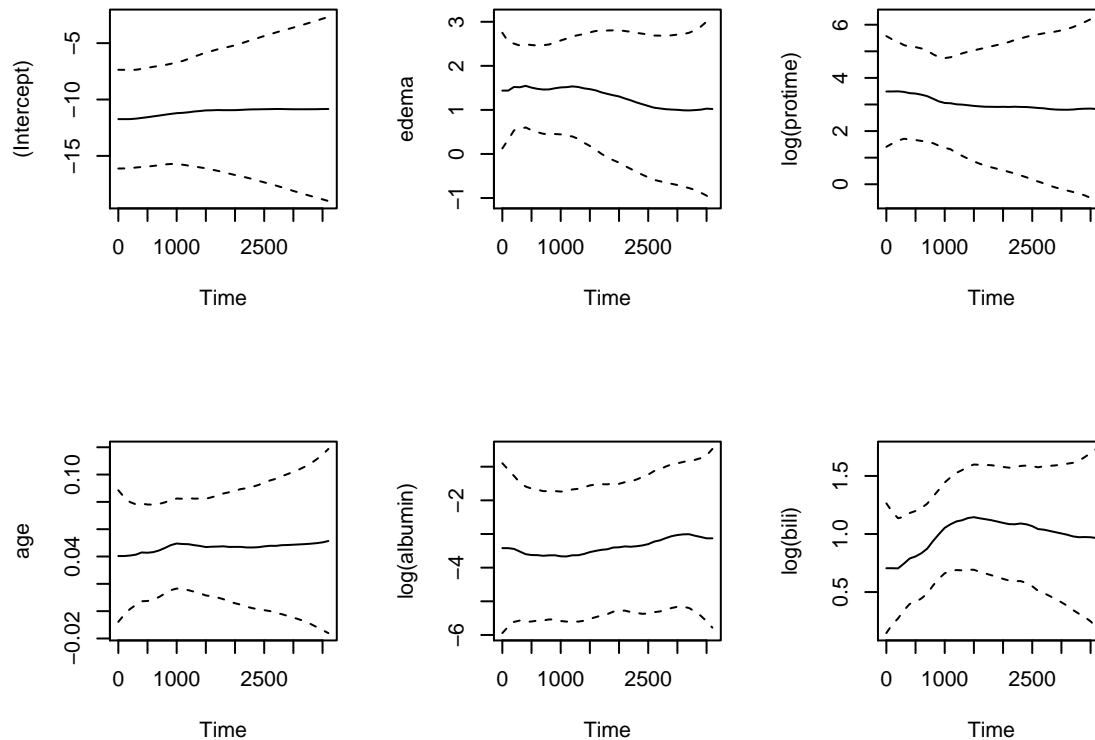
## Extra iteration in the correction step

Another idea is to take extra iterations in the correction step of the EKF. The motivation is that this step has the form of a Newton Rapshon algorithm as pointed out in Fahrmeir (1992) and Fahrmeir (1994). Below, we estimate the model with potentially extra steps in the correction step.

```
# Pre-computed sds of covariates
sds <- c(1, 10, .2, .2, .1, 1)

dd_fit <- ddhazard(
  Surv(tstart, tstop, death == 2) ~ age + edema +
    log(albumin) + log(protime) + log(bili), pbc2,
  id = pbc2$id, by = 100, max_T = 3600,
  Q_0 = diag(10 / sds), Q = diag(0.01 / sds, 6),
  control = list(
    eps = .001,
    NR_eps = 0.0001, # Tolerance in correction step
    LR = .33
  ))

# Plot result
plot(dd_fit)
```



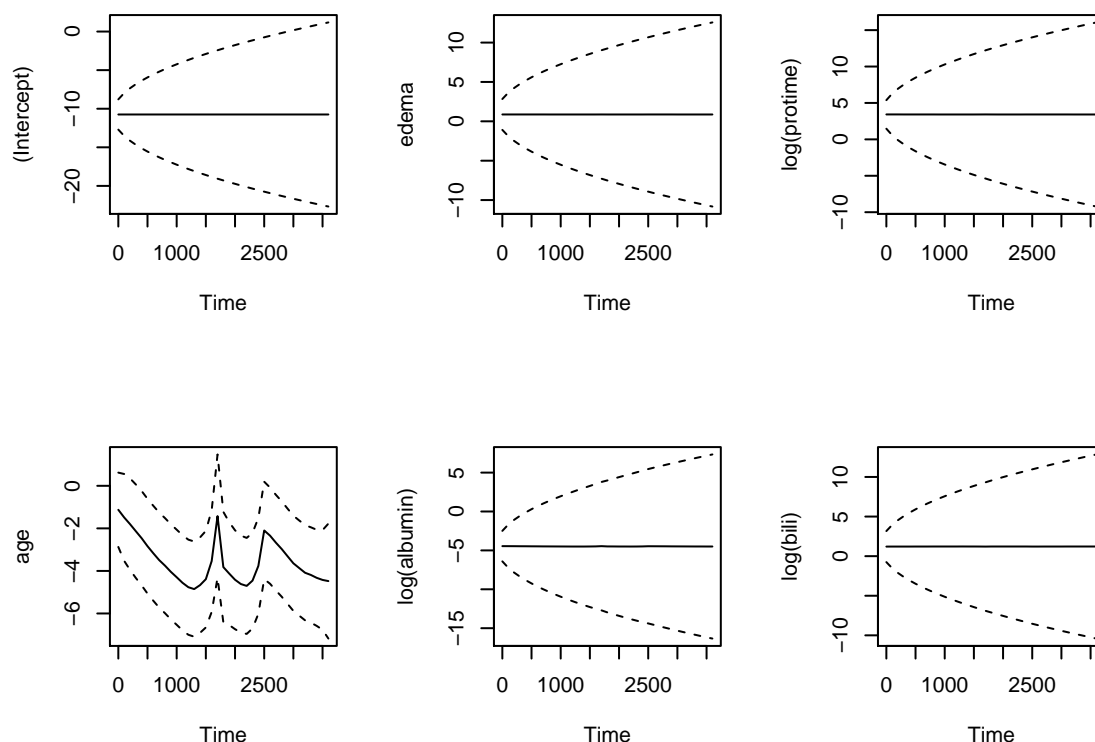
First, we run the code with the `NR_eps` element of the list passed to the `control` argument set to something that is finite. The value is the threshold for the relative change of in the state vector in correction step of the EKF. We end the code above by creating plots of the new estimates.

## Estimation with the Unscented Kalman Filter

Another option is to perform the E-step using an unscented Kalman filter. This is done below. We start by setting the initial co-variance matrix  $\mathbf{Q}_0$  to have large values in the diagonal elements:

```
dd_fit_UKF <- ddhazard(
  Surv(tstart, tstop, death == 2) ~ age +
    edema + log(albumin) + log(protime) + log(bili), pbc2,
  id = pbc2$id, by = 100, max_T = 3600,
  Q_0 = diag(rep(1, 6)), Q = diag(rep(0.01, 6)),
  control = list(method = "UKF", beta = 0, alpha = 1,
    eps = 0.1, n_max = 1e4))

plot(dd_fit_UKF)
```



Clearly, the plots of the estimates are not what we expected. The reason is that  $\mathbf{Q}_0$ 's diagonal entries are quite large. The square root of the diagonal entries are used to form the sigma points in the first iteration. Hence, we mainly get estimates that are either zero or one when  $\mathbf{Q}_0$  is a diagonal matrix with large entries. You can run the code below to see how the algorithm progress:

```
# Not run
tmp_file <- file("pick_some_file_name.txt")
sink(tmp_file)
dd_fit_UKF <- ddhazard(
  Surv(tstart, tstop, death == 2) ~ age +
    edema + log(albumin) + log(protime) + log(bili), pbc2,
  id = pbc2$id, by = 100, max_T = 3600,
```

```

Q_0 = diag(rep(1, 6)), Q = diag(rep(0.01, 6)),
control =
  list(method = "UKF", beta = 0, alpha = 1,
        debug = T)) # <-- prints information in each iteration
sink()
close(tmp_file)

```

It will print quite a lot of information and hence it is recommended to use `sink` to write the output to a file. The main take away is that the conditional co-variance matrices accumulate in each iteration while the state vectors does not move. This motivates us to pick  $\mathbf{Q}$  and  $\mathbf{Q}_0$  more carefully. Our estimates from the EKF was:

```
diag(dd_fit$Q)
```

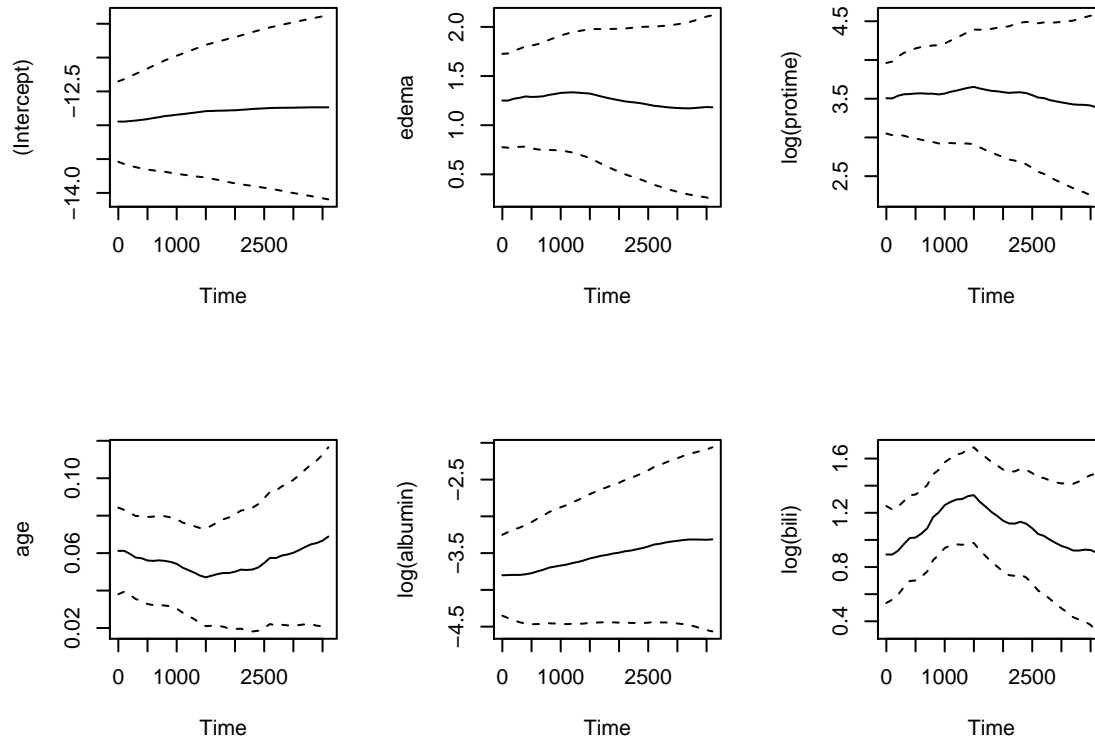
```
## (Intercept)      age      edema log(albumin) log(protime)  log(bili)
##    6.51e-03    1.16e-06    1.18e-03    2.51e-03    1.89e-03    1.91e-04
```

which could motivate us to make the following choices:

```

dd_fit_UKF <- ddhazard(
  Surv(tstart, tstop, death == 2) ~ age +
    edema + log(albumin) + log(protime) + log(bili), pbc2,
  id = pbc2$id, by = 100, max_T = 3600,
  Q_0 = diag(c(0.001, 0.00001, rep(0.001, 4))) * 100, # <-- decreased
  Q = diag(0.0001, 6),                                # <-- decreased
  control =
    list(method = "UKF", beta = 0, alpha = 1, eps = 0.001))
plot(dd_fit_UKF)

```



This is comparable to the fits from the EKF and GAM model. The main points here are:

1. The UKF may work for certain data set. It may require tuning.
2. The algorithm is sensitive to the choice of  $\mathbf{Q}$  and  $\mathbf{Q}_0$ . Further, there is dependence on hyperparameters  $\alpha$ ,  $\kappa$  and  $\beta$  which we have not explored.
3. The output you get by setting `debug` in the list passed to the `control` argument can be useful. Combine this with `sink` because the output may be long.
4. The UKF has shown better performs than the EKF previously. Examples includes Romanenko & Castro (2004), Kandepu, Foss, & Imsland (2008), Julier & Uhlmann (2004), Wan & Van Der Merwe (2000) and chapter 11 of Durbin & Koopman (2012).

## Estimation with rank-one posterior modes approximation

Another method is to use the sequential rank-one approximation of the posterior posterior modes. This is done below. First, we estimate the model with the Extended Kalman filter and then we estimate the model with the rank-one posterior modes approximation method. We estimate both for comparison.

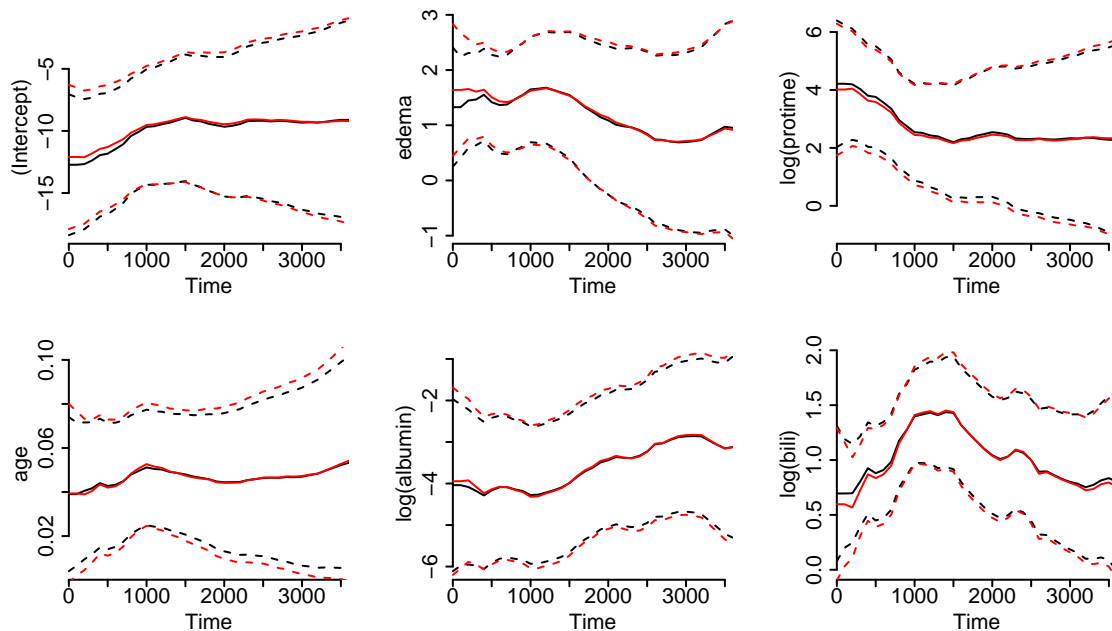
```
dd_fit_EKF <-
  ddhazard(
    Surv(tstart, tstop, death == 2) ~ age + edema +
      log(albumin) + log(protime) + log(bili), pbc2,
    id = pbc2$id, by = 100, max_T = 3600,
    Q_0 = diag(100, 6), Q = diag(0.01, 6),
    control = list(eps = .001))

dd_fit_SMA <-
```

```
ddhazard(
  Surv(tstart, tstop, death == 2) ~ age + edema +
    log(albumin) + log(protime) + log(bili), pbc2,
  id = pbc2$id, by = 100, max_T = 3600,
  Q_0 = diag(100, 6), Q = diag(0.01, 6),
  control = list(
    method = "SMA", # change estimation method
    eps = 0.001))
```

Next, we plot the two sets of predicted coefficients:

```
par(mfcol = c(2, 3))
for(i in 1:6){
  plot(dd_fit_EKF, cov_index = i, col = "black")
  plot(dd_fit_SMA, cov_index = i, col = "red", add = TRUE)
}
```



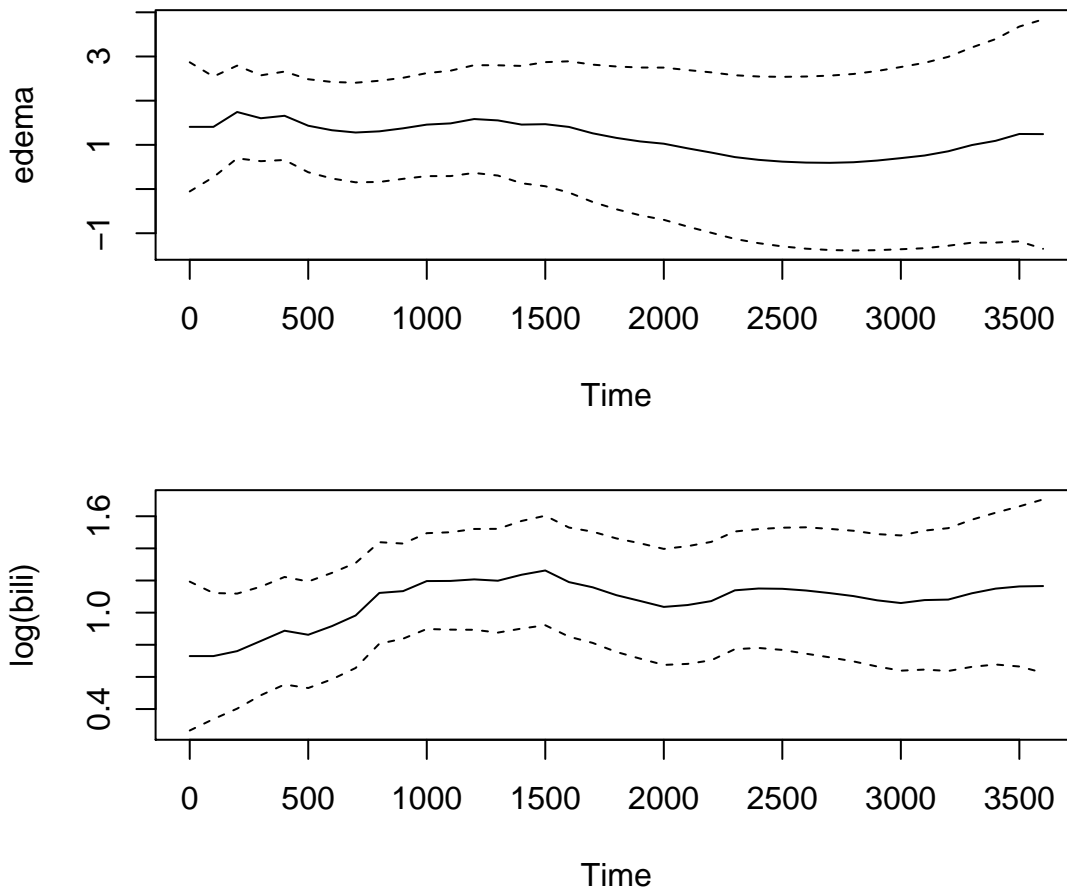
The red curves are from the sequential rank-one approximation. The difference seems minor for this data set.

## Estimation with fixed effects

We may want to fit a model where we assume that some of the co-variables are fixed. For instance, we may want to fit a model where `age`, the intercept, `log(protime)` and `log(albumin)` are fixed. We fix these based on the previous plots where the effects seems no to be time-varying. The model can be fitted as by the following call:

```
dd_fit <- ddhazard(
  Surv(tstart, tstop, death == 2) ~ ddFixed_intercept() +
    ddFixed(age) + ddFixed(log(albumin)) + edema + ddFixed(log(protime)) + log(bili),
  pbc2, id = pbc2$id, by = 100, max_T = 3600,
  Q_0 = diag(100, 2), Q = diag(0.01, 2),
  control = list(eps = .001))
```

```
plot(dd_fit)
```



The predicted curves seems similar. Moreover, the fixed effects are in agreement with the previous fits (they are printed below):

```
dd_fit$fixed_effects
```

```
## (Intercept)      age log(albumin) log(protime)
##      -10.718      0.046      -3.678       3.010
```

## Second order random walk

We will end by fitting a second order random walk to model where only the `log(bili)` effect is time-varying. The motivation is that the second order random walk tend to diverge more easily especially for small data sets. Further, the previous models seems to suggest that it is the only covariate where we may have a time-varying coefficient. We will comparer estimates with the Extended Kalman Filter and the sequential rank-one approximation. First, we make the two estimations:

```
# Define formula
form <- Surv(tstart, tstop, death == 2) ~
      ddFixed_intercept() + ddFixed(age) +
      ddFixed(edema) + ddFixed(log(albumin)) +
      ddFixed(log(protime)) + log(bili)
```

```

# Fit models
dd_fit_EKF <-
  ddhazard(form,
    pbc2, id = pbc2$id, by = 100, max_T = 3600,

    order = 2,          # <-- second order
    Q_0 = diag(100, 2), # <-- needs more elements
    Q = .00001,         # <-- decreased

    control = list(eps = .001))

dd_fit_post <-
  ddhazard(form,
    pbc2, id = pbc2$id, by = 100, max_T = 3600,
    order = 2,
    Q_0 = diag(100, 2),
    Q = .00001,
    control = list(eps = .001, method = "SMA"))

```

We have to decrease the starting value of  $Q$  in the above to get the methods to converge. The fixed effects estimates are not to far from each other:

```

rbind(
  "EKF" = dd_fit_EKF$fixed_effects,
  "Posterior approximation" = dd_fit_post$fixed_effects)

```

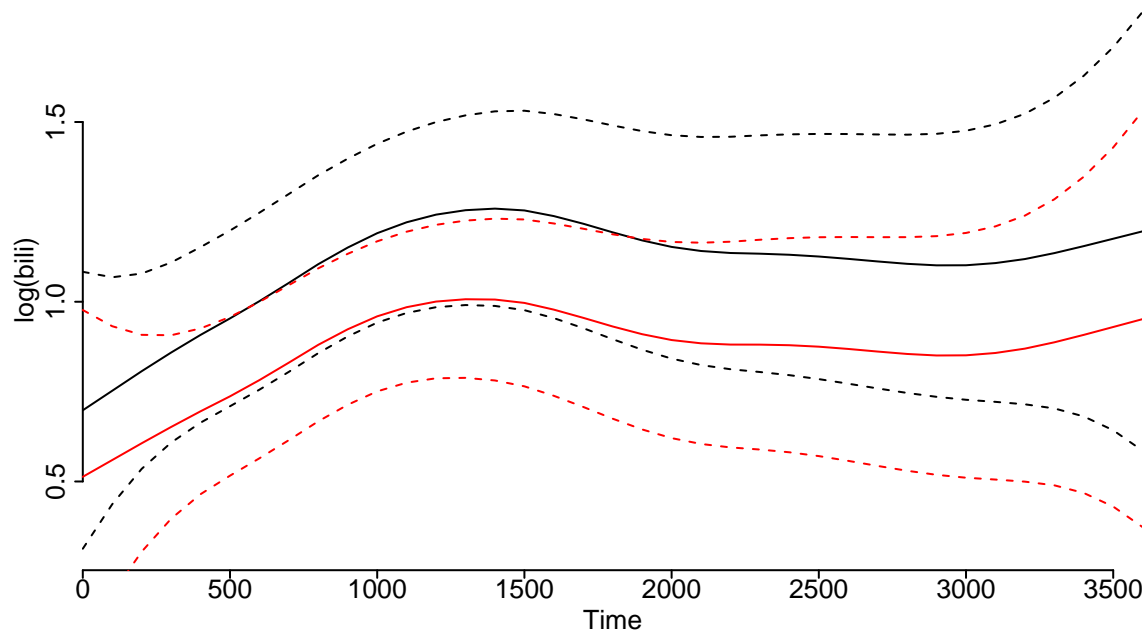
	(Intercept)	age	edema	log(albumin)	log(protime)
## EKF	-11.3	0.0481	1.41	-3.55	3.11
## Posterior approximation	-10.0	0.0344	1.52	-3.13	2.92

The predicted curves are also not to far from each other and the shapes are almost identical:

```

plot(dd_fit_EKF)
plot(dd_fit_post, col = "red", add = T)

```



We see that the predicted curve is more smooth as expected with a second order random walk. Further, we can confirm that the fixed effects are comparable with our previous fits.

## Summary

We have estimated a model using Generalized additive models with the `mgcv` package and a time-varying Cox model with the `timereg` package. Further, we have illustrated how the `ddhazard` function in the `dynamichazard` package can be used. All the fits have been comparable with the Generalized Additive model.

An unaddressed question is why you should this package. Some of the advantageous of the state space model here are among other:

1. You can extrapolate beyond the last observation time. An example hereof could be any time series where the underlying time is the calendar time such a medical trail where we may suspect a different effects of a drug in 2015 than in 1990.
2. The implementation scale well in computation time with the number of individuals.
3. All estimation is made in `c++` with use of the `Armadillo` library. Thus, an optimized version of `Blas` or `Lapack` can decrease the computation time.

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