Simulation study with logit model

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Intro

This note has four objectives. The first objective is to test how the ddhazard fits compare with a Generalized Additive models (GAM) and a "static" logistic model with simulated data. We will look at the following models/estimation methods from ddhazard function in the dynamichazard package:

- Fits with the Extended Kalman Filters (EKF) with and without extra iterations in the scoring step
- Second order random walks with the EKF estimation method
- Mixture of fixed and time varying effects with the EKF estimation method. Fixed effects are both estimated with the E-step and M-step method described in ddhazard vignette
- Fits using the Unscented Kalman filter (UKF)

The second objective is to show how to estimate various models with the function ddhazard. For this reason, the note contains intermediate R code which is not needed to understand the simulation results. Thus, we will use * in the headers of section to distinguish the content. The headers marked with no * indicates sections with results of simulation or contains important comments. Headers with an * and ** shows increasingly less important code to understand the simulation. Consequently, you can skip to the headers with no * if you are only interested in the results.

The third objective is to illustrate how the various methods performs for out-of-time prediction (forecasting). By out-of-time we mean that we only observe outcomes up to given time, d, and then predict the outcome for future observations at time d+1.

The fourth objective is to show that both the EKF and UKF scales linearly with the number individuals (series).

All method use the logistic link function. We will do three runs of experiments in the following order:

- 1. A Model where all effects are time varying and we use the correct binning intervals
- 2. A model where only one parameter is time varying and we use the correct binning intervals
- 3. A Model where all effects are time varying but we use incorrect binning intervals

where correct or incorrect binning intervals refers to whether or not we bin at the same time where the coefficient are simulated to change. For example, we bin correctly where we simulate the coefficients to change at time $1, 2, \ldots, d$ and we estimate the coefficient at time $1, 2, \ldots, d$. The models will be compared in terms of Brier score, median absolute residuals and standard deviation of the absolute residuals. All metrics will be reported on out-sample data or out-of-time data. All plots will have true coefficients as continuous lines while dashed lines are estimates.

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 to pass devtools::install_github

[1] "boennecd/dynamichazard@8ad8c0701479c79a581e0143a1b01cc12e01d01a"

devtools::install github(current version)

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

install.packages("dynamichazard")

Moreover, you will also find the source code for the vignette at the github page. The note is not meant to be self contained. It is recommend to see ddhazard vignette for an introduction to the models and methods in the dynamichazard package.

Notions

For clarity, here is a list of used notions:

- Run: An experiment with one of the three previously specified settings where we make k simulations with n series in each
- Simulation: One simulation within a run with one set of coefficients $\vec{\beta}_0, \dots, \vec{\beta}_d$ and given number n of series
- Series/individuals: A person/individual either making it to the end of the time of the given simulation or dying at some time during the period
- Coefficients: the entries of the vectors $\vec{\beta}_t$ in a given simulation
- Covariates: vectors \vec{x}_{it} for a given individual at a given time in simulation

Findings

The findings are:

- The UKF method seems to perform well for both small and larger number of series
- Taking multiple iterations in the correction step of the EKF seems to be beneficial
- Specifying a fixed effect as time varying or setting the binning number incorrectly has little effect on the results

You will see that the the estimation sometimes fails. It is worth stressing that it is my experience that you can always do "trail-and-error" with the initial covariance matrix in the state equation, the covariance matrix at time zero and tuning parameters in order to get a model to fit a given dataset. Of course, it is a disadvantage that any given data set may require some tuning by the user. Although as will be shown, tuning by the user is not often needed with data sets like those presented here.

Setup

The following values will be used in the simulation:

```
ns <- c(200, 2000) # Number of series
n_beta <- 5  # Number of covariates
T_max <- 20  # The last time we observe
n_sims <- 100  # Number of simulation in each run

gsub("(^.+)(/dynamichazard.+$)", "...\\2", getwd())</pre>
```

```
## [1] ".../dynamichazard/vignettes/Prebuild"
```

```
source("../../R/test_utils.R")
```

ns is the number of series (individuals) we will estimate in each of the simulation in each of the runs. Thus, we will perform simulations with a total of 200 and 2000 series in each. Each simulation will have n_beta = 5 covariates plus an intercept. Each run will simulate n_sims = 100 times. Finally, we source the test.utils.R file to define the simulation function. You can find this script on the github site. T_max is the number of bins/intervals we observe. Thus, we have 1, 2, ..., T_max + 1 covariate vectors (+1 for the time zero coefficient vector).

Fitting true model

We will make runs for various number of individuals in this section where we estimate a model where all effects are time varying and we use the correct binning intervals. Thus, the only models that are miss specified are the model with one time varying effect (which will be x2) and the model were we use a second order random walk.

do.call function

We will use do.call in this vignette. To my knowledge, do.call is not standard so this section is included to give a brief introduction to do.call for users who are not familiar with do.call. We can take an example with the mean function. We will make the following call where we set na.rm to TRUE:

```
mean(x = c(1, 2, NA, 6), na.rm = T)
## [1] 3
```

This call can also be made as follows do.call:

```
arg_ex \leftarrow list(x = c(1, 2, NA, 6), na.rm = T)
do.call(mean, arg_ex)
```

```
## [1] 3
```

Hence, do.call is useful in situation where we make calls where almost all the arguments are the same. For example, in the setting where we have arguments a1, a2, ..., up to a1000 and we only want to change argument a101 say. This can then be done as follows:

Definition of simulation function

Below, we define a list of default_args (default arguments) to our simulation function which we can later use using do.call.

```
# Default arguments for simulation
default_args <- list(
    n_vars = n_beta, # Number of betas not including intercept
    beta_start = c(-1, -.5, 0, 1.5, 2), # start value of coeffecients
    intercept_start = -5, # start value of intercept
    sds = c(.1, rep(.5, n_beta)), # std. deviations in state equation
    t_max = T_max, # Largest time we observe
    x_range = 1, # range of covariates
    x_mean = .5, # mean of covariates
    tstart_sampl_func = # we randomly draw the start time of each serie</pre>
```

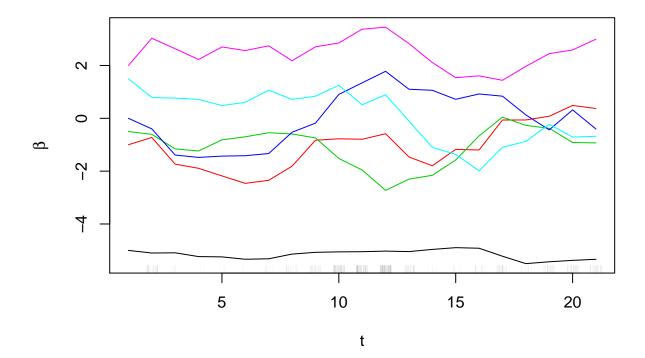
```
function(...) max(runif(1, min = -10, max = 18), 0)
)
```

Let $\vec{\beta}_t$ denote the time varying covariates at time t. Then the beta_start is the time 0 values of the coefficients and intercept_start is the starting value of the intercept. The sds are the standard deviations, σ_i , in the state equation. Hence,

$$\beta_{j,t} = \beta_{j,t-1} + \epsilon_{j,t}, \qquad \epsilon_{j,t} \sim N(0, \sigma_j^2)$$

where each margin is independent of the others. The x_mean and x_range defines how the covariate values are simulated. The above setting implies that $x_{itj} = \text{Unif}(0.5 - 1/2, 0.5 + 1/2)$ where x_{itj} is the *i*'th individuals covariate *j* at time *t*. The covariate vector \vec{x}_{it} is updated at time differences of $1 + \eta$ where $\eta \sim \text{Exp}(1)$ and η s are drawn separately for each individual for each covariate vector. The motivation for this behavior is that we can have different covariate update times than our binning time in a given study. For instance, say we are looking at a medical study and the covariates are laboratory values. The time of laboratory values from an individual's visit the doctor can differ from whatever binning periods we use in the state-space model. Further, the time when laboratory values are updated can differ between patients. One might see his doctor every week or so while another only sees his doctor every year.

Below we illustrate how the coefficients vectors from a simulation can look:



The black line is the intercept while the colored lines are the coefficients for the covariates. The lines on the x-axis illustrate when we observe that individuals die. There is one line for each death. Next, we can look at the number of failures in each simulation:

```
# We get a "decent" amount of failures and survivers in some of the simulations
# We use do.call to avoid repeating the above argument list
set.seed(468249)
n_fails_in_sim <- rep(NA_real_, 15)
for(i in seq_along(n_fails_in_sim)){
    sims <- do.call(test_sim_func_logit, c(
        default_args, c(list(n_series = max(ns))))) # Take largest amount of series
    n_fails_in_sim[i] <- sum(sims$res$event)
}
n_fails_in_sim # number of failures in each simulation</pre>
```

91 1321 707 1635 466 1548

83 1596 1577

* Definition of fit functions

349 759 1062

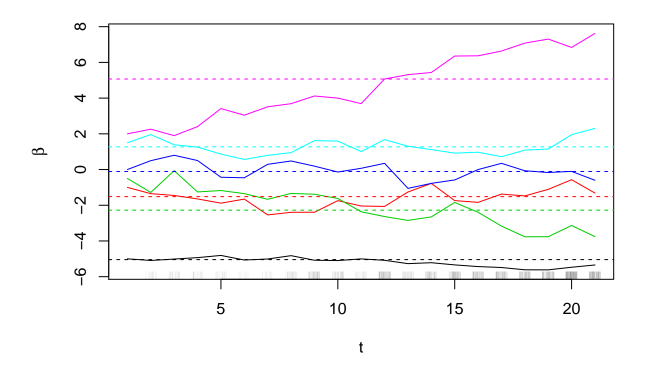
[1] 1132 1822

[15] 1151

We will define functions to estimate the different models with a data frame as the first argument where the data frame is from a test_sim_func_logit call. This will reduce the amount of code later.

** Definition of static fit

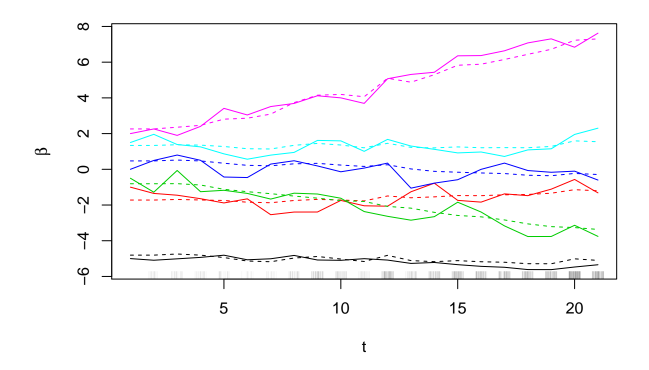
Below, we define function to fit a model where the coefficients are fixed $(\vec{\beta}_t = \vec{\beta})$. It is estimated using glm.fit:

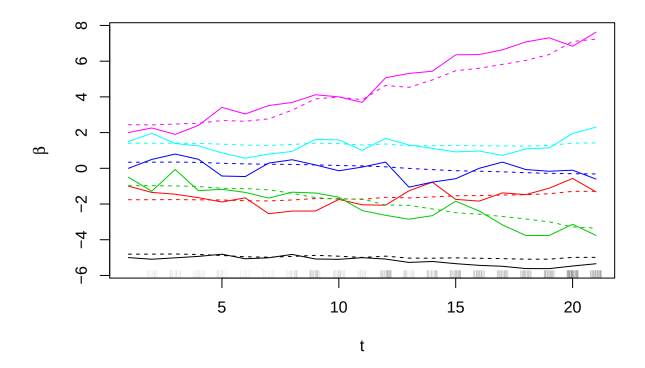


* Definition of ddhazard fit functions

Below, we define a function to fit a first order random walk model with a given learning rate and potential extra iterations in the scoring step (see the ddhazard vignette for details):

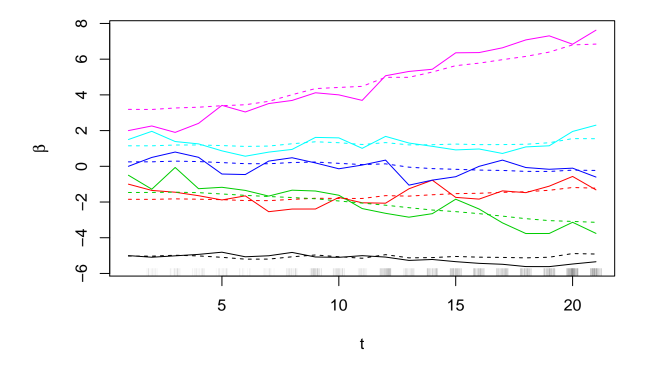
```
library(survival); library(dynamichazard)
# We will use glm.fit for the starting value
options(ddhazard_use_speedglm = FALSE)
# Set up function ddhazard fit function for convenience
      learning rate in correction step
# NR eps: tolerance in correction step. NULL yields no extra iterations
fit_funcs$dd <- function(s = sims$res, LR = 1, NR_eps = NULL)</pre>
 tryCatch({
    ddhazard(
      formula = Surv(tstart, tstop, event) ~ x1 + x2 + x3 + x4 + x5,
      data = s, max_T = T_max, by = 1, id = s$id,
      Q_0 = diag(
        # We set the Q_0 argument lower when we take multiple iterations
        # See the ddhazard vignette under the GMA model for arguments herefor
        if(is.null(NR_eps)) 1000000 else 1,
        n_{\text{beta}} + 1),
      Q = diag(.01, n_beta + 1),
      control = list(LR = LR, NR_eps = NR_eps, eps = 0.01))
  }, error = function(...) NA) # Return NA if fails
fit <- fit_funcs$dd()</pre>
# Plot estimates and actual coffecients
plot_func(ylim = fit$state_vecs)
matplot(fit$state_vecs, col = 1:(n_beta + 1), lty = 2,
        type = "1", add = T)
```





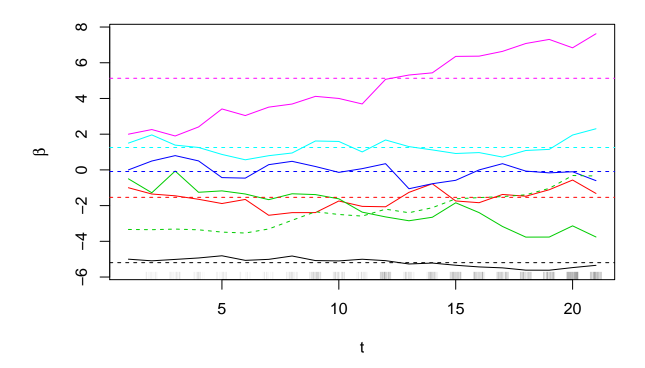
Below, we define a function to fit a first order random walk model with the UKF method:

```
# Fitting with UKF
fit_funcs$dd_UKF <- function(s = sims$res, alpha = 1, beta = 0){</pre>
  tryCatch({
    ddhazard(
      formula = Surv(tstart, tstop, event) ~ x1 + x2 + x3 + x4 + x5,
      data = s, max_T = T_max, by = 1, id = sid,
      Q_0 = diag(1, n_beta + 1), Q = diag(.01, n_beta + 1),
      control = list(
        eps = 0.1,
        alpha = alpha,
                            # Set tuning parameter
        beta = beta,
                            # Set tuning parameter
        method = "UKF"))
                            # Set estimation method (EKF is default)
  }, error = function(...) NA) # Return NA if fails
fit <- fit_funcs$dd_UKF()</pre>
# Look at new plot
plot_func(ylim = fit$state_vecs)
matplot(fit$state_vecs, col = 1:(n_beta + 1), lty = 2,
        type = "l", add = T)
```



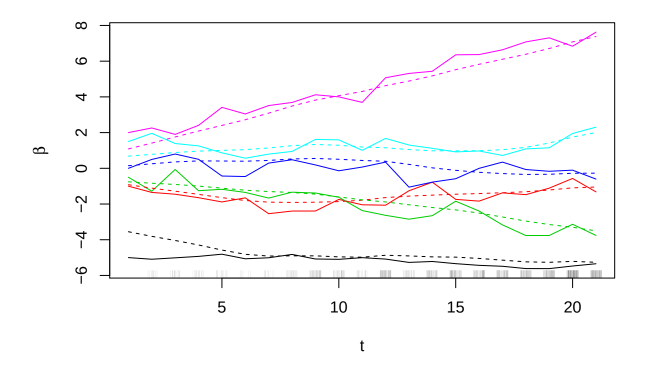
Below, we define a function to estimate a first order random walk model where only one parameter (x2) is time varying:

```
# Fitting with fixed effects
fit_funcs$dd_fixed <- function(</pre>
  s = sims$res, LR = 1, NR_eps = NULL,
  fixed_terms_method = "M_step"){ # The method to use to estimate the fixed
                                   # fixed effects
  tryCatch({
    ddhazard(
      formula = Surv(tstart, tstop, event) ~
        ddFixed(1) +
                                                  # Fix intercept
        ddFixed(x1) + x2 +
                                                  # Note x2 is time varying
        ddFixed(x3) + ddFixed(x4) + ddFixed(x5),
      data = s, max_T = T_max, by = 1, id = sid,
      Q_0 = diag(1, 1), Q = diag(.01, 1),
      control = list(LR = LR, NR_eps = NR_eps, eps = 0.1,
                     fixed_terms_method = fixed_terms_method))
  }, error = function(...) NA) # Return NA if fails
fit <- fit_funcs$dd_fixed()</pre>
# Look at new plot
plot_func(ylim = range(fit$state_vecs, fit$fixed_effects))
matplot(fit$state_vecs, col = 3, lty = 2, type = "1", add = T)
abline(h = fit\fixed_effects, col = c(1:2, 4:6), lty = 2)
```



Next, we define a function to fit the model with a second order random walk:

```
# Fitting with second order
fit_funcs$dd_2_order <- function(s = sims$res, LR = 1, NR_eps = NULL){</pre>
  tryCatch({
    ddhazard(
      formula = Surv(tstart, tstop, event) ~ x1 + x2 + x3 + x4 + x5,
      data = s, max_T = T_max, by = 1, id = sid,
      # Q_0 and Q needs more elements
      Q_0 = diag(c(rep(1, n_beta + 1), rep(0.5, n_beta + 1))),
      Q = diag(c(rep(.01, n_beta + 1))),
      order = 2, # specify the order
      control = list(LR = LR, NR_eps = NR_eps, eps = 0.1))
  }, error = function(...) NA) # Return NA if fails
fit <- fit_funcs$dd_2_order()</pre>
# Look at new plot
plot_func(ylim = fit$state_vecs)
matplot(fit$state_vecs[, 1:6], col = 1:(n_beta + 1), lty = 2,
        type = "1", add = T)
```



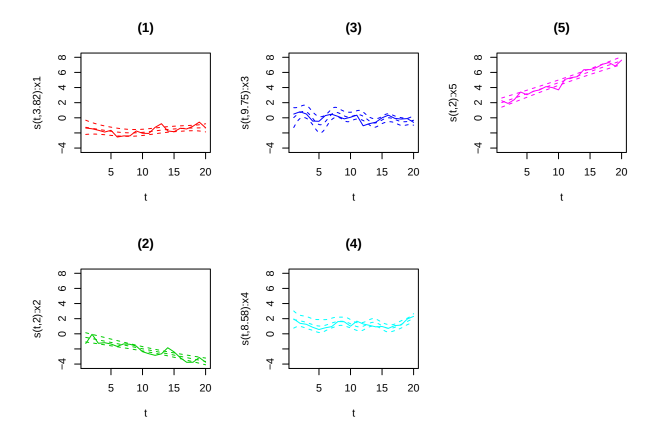
** Definition of GAM fit function

We define the estimation method for the Generalized additive model in the next code snippet. We use bam function from the mgcv package which corresponds to gam but for large datasets.

```
library(mgcv)
fit_funcs$gam <- function(s = sims$res){</pre>
  # get data frame for fitting
  dat_frame <- get_survival_case_weights_and_data(</pre>
    formula = Surv(tstart, tstop, event) ~ x1 + x2 + x3 + x4 + x5,
               data = s, max_T = T_max, by = 1, id = s$id, use_weights = F)$X
  # fit model
  bam(
    formula = Y ~
      # cr is cubic basis with k knots
      s(t, bs = "cr", k = 10, by = x1) +
      s(t, bs = "cr", k = 10, by = x2) +
      s(t, bs = "cr", k = 10, by = x3) +
      s(t, bs = "cr", k = 10, by = x4) +
      s(t, bs = "cr", k = 10, by = x5),
  family = binomial, data = dat_frame,
  method = "GCV.Cp",
  control =
    gam.control(nthreads = parallel::detectCores() - 1)) # Use parallel
```

```
# fit model
fit <- fit_funcs$gam()

# Compare plot
layout(matrix(1:6, nrow = 2))
for(i in 1:n_beta){
   plot(fit, pages = 0, rug = F, col = i + 1, select = i, lty = 2,
        main = paste0("(", i, ")"))
   lines(sims$betas[-1, i + 1], col = i + 1)
}</pre>
```



** Definition of prediction functions

The following code snippets define predictions methods for each of the estimation methods. We start off by defining a split function such that we can sample individuals (series) into a test set and a training test:

```
# Illustrate use
tmp <- split_func()
# No ids intersect in the two sets
length(intersect(tmp$test_dat$id, tmp$fit_dat$id))
## [1] 0
# The union is exactly the number of ids we simulated
length(union(tmp$test_dat$id, tmp$fit_dat$id))</pre>
```

[1] 2000

Having defined the splitting method, we turn to the prediction functions. The idea is to define the brier_funcs\$general function which takes in a prediction function, a fit and a data frame. Next, we then define individual prediction functions for each of the models which will be passed to brier_funcs\$general:

```
# Define general prediction function
brier funcs <- list()</pre>
brier funcs$general <- function(brier func, fit, eval data frame){
  d_frame <- get_survival_case_weights_and_data(</pre>
    formula = Surv(tstart, tstop, event) ~ x1 + x2 + x3 + x4 + x5,
    data = eval_data_frame, max_T = T_max, by = 1, id = eval_data_frame$id,
    use_weights = F)$X
  # Change start and stop times
  d frame$tstart <- d frame$t - 1</pre>
  d_frame$tstop <- d_frame$t</pre>
  # Compute residuals
  resids <- brier_func(fit, d_frame)
  # Return estimates
  with(
    resids,
    list(brier = mean(response^2),
         median abs res = median(abs(response)),
         sd_res = sd(response),
         median_deviance = median(deviance),
         sd_deviance = sd(deviance)))
}
# Prediction method for static model
brier funcs$static <- function(fit, d frame){</pre>
  preds <- predict(fit, newdata = d_frame, type = "response")</pre>
 list(
    response = d_frame$Y - preds,
    deviance =
      d_frame_Y * log(preds) + (1 - d_frame_Y) * log(1 - preds))
}
# Test function
fit <- fit_funcs$static(tmp$fit_dat)</pre>
cols <-
  c("brier", "median_abs_res", "sd_res", "median_deviance", "sd_deviance")
unlist( # in sample stats
```

```
brier_funcs$general(brier_funcs$static, fit, tmp$fit_dat)[cols])
##
             brier median_abs_res
                                              sd res median deviance
                            0.01763
                                             0.20636
                                                             -0.01779
##
           0.04258
##
       sd deviance
##
           0.52205
unlist( # out sample stats
 brier_funcs$general(brier_funcs$static, fit, tmp$test_dat)[cols])
##
             brier median_abs_res
                                              sd res median deviance
##
                            0.01720
                                             0.21768
           0.04741
                                                            -0.01735
##
       sd deviance
##
           0.57509
# Define prediction function for ddhazard model
brier funcs$dd <- function(fit, d frame){</pre>
  preds <- predict(fit, new_data = d_frame, tstart = "tstart", tstop = "tstop")</pre>
  # We truncate as we can get zero-one outcome if the model diverges
 preds$fits <- pmax(pmin(preds$fits, 1 - 1e-14), 1e-14)</pre>
 list(
    response = d_frame$Y - preds$fits,
      d_frame$Y * log(preds$fits) + (1 - d_frame$Y) * log(1 - preds$fits))
}
fit <- fit_funcs$dd(tmp$fit_dat)</pre>
unlist( # in sample stats
 brier_funcs$general(brier_funcs$dd, fit, tmp$fit_dat)[cols])
##
             brier median abs res
                                             sd res median deviance
##
           0.03816
                            0.01664
                                             0.19531
                                                             -0.01678
##
       sd_deviance
           0.48595
unlist( # out sample stats
  brier_funcs$general(brier_funcs$dd, fit, tmp$test_dat)[cols])
##
             brier median_abs_res
                                              sd_res median_deviance
##
           0.04298
                            0.01645
                                             0.20731
                                                             -0.01659
##
       sd_deviance
           0.53966
# Define prediction function for gam model
brier_funcs$gam <- function(fit, d_frame){</pre>
 preds <- predict(fit, newdata = d_frame, type = "response")</pre>
 list(
    response = d_frame$Y - preds,
    deviance =
      d frame\$Y * log(preds) + (1 - d frame<math>\$Y) * log(1 - preds))
}
fit <- fit_funcs$gam(tmp$fit_dat)</pre>
```

```
unlist( # in sample stats
 brier_funcs$general(brier_funcs$gam, fit, tmp$fit_dat)[cols])
             brier median_abs_res
##
                                            sd_res median_deviance
##
           0.03845
                           0.01472
                                            0.19610
                                                           -0.01483
##
       sd deviance
##
           0.50914
unlist( # out sample stats
  brier_funcs$general(brier_funcs$gam, fit, tmp$test_dat)[cols])
##
             brier median abs res
                                            sd res median deviance
##
           0.04308
                           0.01445
                                            0.20748
                                                           -0.01456
##
       sd deviance
##
           0.56195
```

** Definition of multiple simulations function

To make things easier, we define a function that takes in a function to simulate from. Given a function to simulate with, the new function perform n_sims = 100 simulations for each of values ns (200 and 2000):

```
simulate_n_print_res <- function(</pre>
  sim_func, # Function that takes one argment which is number of series
  NR_eps = c(.0001, NA), # Tolerance in scoring step
  file_prefix) # file_prefix for output
  for(n in ns){
    file_name <- pasteO(file_prefix, "_", n, ".RDS")</pre>
    do_compute <- !file.exists(file_name)</pre>
    if(do_compute){
      out \leftarrow array(NA_real_, dim = c(n_sims, 8, 5),
                    dimnames = list(
                      NULL.
                      c("static", "Extra correction", "Single correction",
                        "2 order EKF", "Fixed E-step", "Fixed M-step", "UKF", "gam"),
                      c("Brier", "Abs res", "Sd res", "Dev", "Sd dev")))
      n_failures_and_surviers <- array(</pre>
        NA_{integer}, dim = c(2, n_sims),
        dimnames = list(c("# failures", "# survivers"), NULL))
      #*****
      # Progress bar for inpatient people (me)
      pb <- winProgressBar(</pre>
        paste("Estimating with n =", n), "", 0, n_sims, 50)
      #*****
      for(i in 1:n_sims){
        #****
        info <- sprintf("%.2f%% done", 100 * (i - 1) / n sims)
        setWinProgressBar(pb, i - 1, paste("Estimating with n =", n), info)
        #*****
```

```
# Sample until we get an outcome have sufficient amount of deaths and
# survivers
repeat{
  sims <- sim func(n)
  # We want some survivers and some deaths
  if(sum(sims$res$event) > 25 && n - sum(sims$res$event) > 25)
    break
}
n_failures_and_surviers["# failures", i] <- sum(sims$res$event)</pre>
n_failures_and_surviers["# survivers", i] <- n - sum(sims$res$event)</pre>
# Split data
sim_split <- split_func(sims$res)</pre>
# Fit static model
static_fit <- fit_funcs$static(sim_split$fit_dat)</pre>
# Fit dd model
dd_fits <- list(rep(NA, length(NR_eps)))</pre>
for(k in seq_along(NR_eps)){
  dd_fits[[k]] <- fit_funcs$dd(</pre>
    sim_split$fit_dat,
    NR_eps = if(is.na(NR_eps[k])) NULL else NR_eps[k])
# Fit second order
dd_2_order <- fit_funcs$dd_2_order(sim_split$fit_dat)</pre>
# Fit fixed effect
dd_fixed_E_step <- fit_funcs$dd_fixed(sim_split$fit_dat,</pre>
                                        fixed_terms_method = "E_step")
dd_fixed_M_step <- fit_funcs$dd_fixed(sim_split$fit_dat,</pre>
                                        fixed_terms_method = "M_step")
# UKF fit
dd_UKF <- fit_funcs$dd_UKF(sim_split$fit_dat)</pre>
# Fit gam model
gam_fit <- fit_funcs$gam(sim_split$fit_dat)</pre>
# Evalute on test data
models <- c(list(static_fit), dd_fits,</pre>
             list(dd_2_order, dd_fixed_E_step, dd_fixed_M_step,
                  dd_UKF, gam_fit))
eval_funcs = c(brier_funcs$static,
                replicate(length(dd_fits) + 4, brier_funcs$dd),
                brier_funcs$gam)
for(j in seq_along(models)){
  if(length(models[[j]]) == 1 && is.na(models[[j]]))
```

```
next # We have to skip model fits that failed
        metrics <- brier funcs$general(</pre>
          eval_funcs[[j]], models[[j]], sim_split$test_dat)
        out[i, j, "Brier"] <- metrics$brier</pre>
        out[i, j, "Abs res"] <- metrics$median_abs_res</pre>
        out[i, j, "Sd res"] <- metrics$sd_res</pre>
        out[i, j, "Dev"] <- metrics$median_deviance</pre>
        out[i, j, "Sd dev"] <- metrics$sd_deviance</pre>
      }
    }
    #*****
    close(pb)
    #****
    # Save results
    saveRDS(out, file = file_name)
  # Load results
  out <- readRDS(file = file_name)</pre>
  # Print results
  did_fit <- apply(out[, , 1], 2, function(x) n_sims - sum(is.na(x)))</pre>
  n_cases_all_success <- sum(complete.cases(out[, , 1]))</pre>
  metric_where_all_fit <-</pre>
    t(apply(out[complete.cases(out[, , 1]), , , drop = F], 3, colMeans))
  metric_where_all_fit <- formatC(metric_where_all_fit ,format="f", digits=3)</pre>
  n_cases_all_success <- formatC(n_cases_all_success, format="d")</pre>
  print(knitr::kable(cbind(
    t(metric_where_all_fit), "# succesful fits" = did_fit),
        caption = paste(
  "Mean of metrics with", n/2, "series in test and fit data. 'Abs res' is the median of the absolute:
  align = "r"))
  cat("\n")
  # Make boxplot of std deviance residuals
  par(mar = c(5, 8, 4, 2))
  boxplot(
    out[complete.cases(out[, , 1]), , "Sd dev"],
    main = paste0("Std of deviance residuals w/ ", n/2 ," series"),
    cex.axis = .75, horizontal = TRUE, las = 2,
    ylim = c(min(out[,, "Sd dev"], na.rm = T),
              min(max(out[,, "Sd dev"], na.rm = T), 2)))
}
```

Simulating

We are now able to simulate from the model where all effects are time varying and we use the correct binning intervals with the code below:

```
set.seed(1243)
# Use simulation function
simulate_n_print_res(
    sim_func = function(n)
    do.call(test_sim_func_logit, c(default_args, c(list(n_series = n)))),
    file_prefix = "logit_sim_all_varying")
```

Table 1: Mean of metrics with 100 series in test and fit data. 'Abs res' is the median of the absolute residuals, 'Sd res' is the standard deviation of the residuals, 'Dev' is median of the deviance residuals and 'Sd dev' is the standard deviation of deviance residuals. Only simulations that succeeds for all setups are included. There are 100 of these simulations. The last column shows the number of successful fits for each setup.

	Brier	Abs res	Sd res	Dev	Sd dev	# succesful fits
static	0.043	0.033	0.199	-0.033	0.559	100
Extra correction	0.041	0.030	0.193	-0.030	0.522	100
Single correction	0.042	0.038	0.195	-0.039	0.489	100
2 order EKF	0.042	0.038	0.196	-0.039	0.900	100
Fixed E-step	0.043	0.043	0.198	-0.044	0.499	100
Fixed M-step	0.042	0.030	0.196	-0.031	0.549	100
UKF	0.041	0.029	0.194	-0.030	0.535	100
gam	0.042	0.020	0.196	-0.020	0.634	100

Std of deviance residuals w/ 100 series

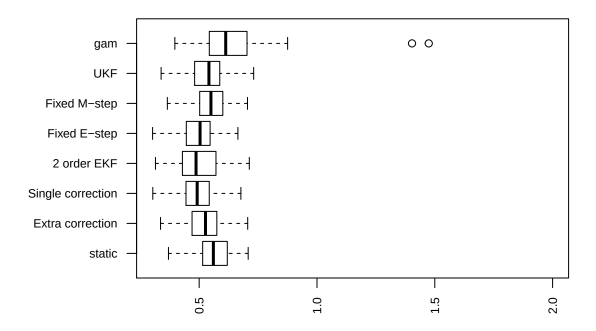
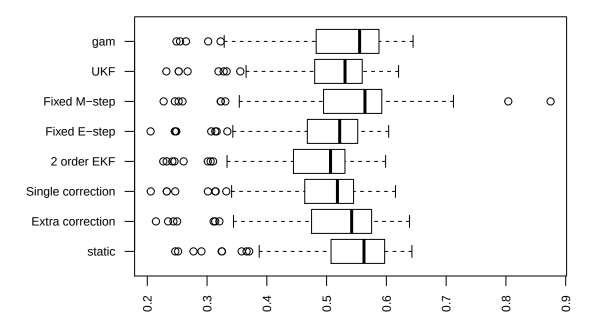


Table 2: Mean of metrics with 1000 series in test and fit data. 'Abs res' is the median of the absolute residuals, 'Sd res' is the standard deviation of the residuals, 'Dev' is median of the deviance residuals and 'Sd dev' is the standard deviation of deviance residuals. Only simulations that succeeds for all setups are included. There are 100 of these simulations. The last column shows the number of successful fits for each setup.

	Brier	Abs res	Sd res	Dev	Sd dev	# succesful fits
static	0.053	0.050	0.211	-0.053	0.534	100
Extra correction	0.045	0.032	0.198	-0.033	0.510	100
Single correction	0.047	0.038	0.200	-0.039	0.488	100
2 order EKF	0.046	0.042	0.199	-0.043	0.474	100
Fixed E-step	0.048	0.047	0.203	-0.049	0.494	100
Fixed M-step	0.049	0.037	0.205	-0.038	0.536	100
UKF	0.046	0.036	0.199	-0.037	0.503	100
gam	0.046	0.030	0.199	-0.031	0.520	100

Std of deviance residuals w/ 1000 series



Notice that we only compare across methods with mean metrics where all succeeded to fit.

Conclussion on run

All models perform better than the static model in terms of Brier score. The UKF and EKF with extra iterations also does well on the Brier score compared with the GAM model. The finding is slight different if we look at the box plot of the standard deviations of the deviance residuals. The GAM model does worse here compared to the other methods.

Single time varying parameter

In this part, we will look at the performs when only singe coefficient (x2) varies. Thus, we can see if the models where only (x2) is modeled as varying performs performs better.

** Definition of simulation function

We start by defining the simulation function. The main change here is that we only set a single standard deviation and that we set it larger than before:

```
# Use simulation function
set.seed(9999)
sim_one_varying <- function(n){
  test_sim_func_logit(</pre>
```

```
n_series = n,
sds = c(sqrt(3)), # Large variance
is_fixed = c(1:2, 4:6), # All but param three (x2) is fixed

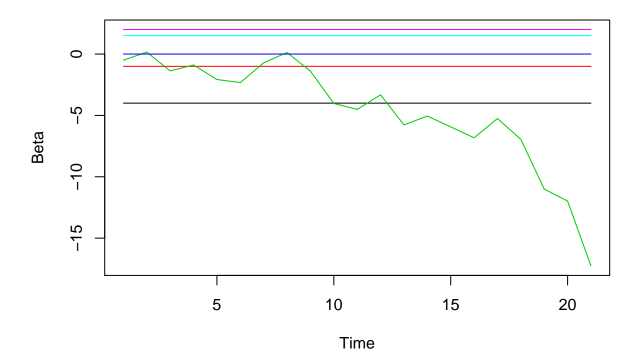
# Same values as before
n_vars = n_beta,
beta_start = c(-1, -.5, 0, 1.5, 2),
intercept_start = -4,
t_max = T_max,
x_range = 1,
x_mean = .5)
}
```

* Illustration of single simulation

```
# We get a more variable number of failures and survivers (we simulate 200
# series)
replicate(10, sum(sim_one_varying(200)$res$event)) # print number of failures

## [1] 197 200 104 187 129 49 200 37 200 62

# Here is an example of a series
tmp <- sim_one_varying(200)
matplot(tmp$betas, type = "l", lty = 1, ylab = "Beta", xlab = "Time")</pre>
```



Simulating

We can simulate with the following call:

```
# Use simulation function
set.seed(8080)
simulate_n_print_res(
   sim_func = sim_one_varying,
   file_prefix = "logit_sim_one_variying")
```

Table 3: Mean of metrics with 100 series in test and fit data. 'Abs res' is the median of the absolute residuals, 'Sd res' is the standard deviation of the residuals, 'Dev' is median of the deviance residuals and 'Sd dev' is the standard deviation of deviance residuals. Only simulations that succeeds for all setups are included. There are 100 of these simulations. The last column shows the number of successful fits for each setup.

	Brier	Abs res	Sd res	Dev	Sd dev	# succesful fits
static	0.036	0.030	0.182	-0.031	0.559	100
Extra correction	0.035	0.026	0.179	-0.026	0.523	100
Single correction	0.040	0.030	0.183	-0.031	0.703	100
2 order EKF	0.036	0.035	0.181	-0.036	0.662	100
Fixed E-step	0.036	0.035	0.181	-0.037	0.514	100
Fixed M-step	0.036	0.024	0.181	-0.025	0.561	100
UKF	0.035	0.026	0.180	-0.027	0.547	100
gam	0.035	0.019	0.180	-0.020	0.573	100

Std of deviance residuals w/ 100 series

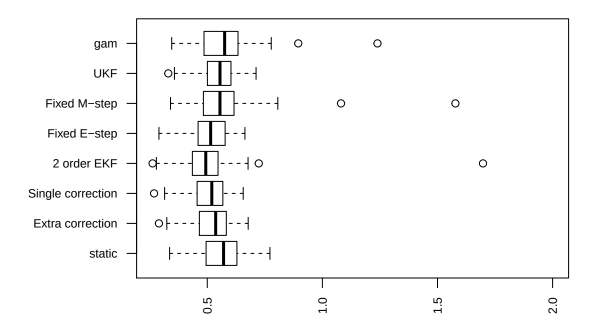
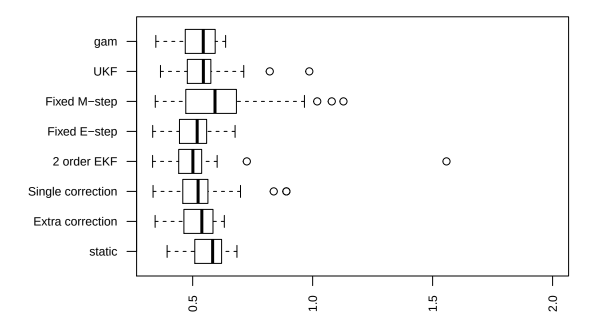


Table 4: Mean of metrics with 1000 series in test and fit data. 'Abs res' is the median of the absolute residuals, 'Sd res' is the standard deviation of the residuals, 'Dev' is median of the deviance residuals and 'Sd dev' is the standard deviation of deviance residuals. Only simulations that succeeds for all setups are included. There are 92 of these simulations. The last column shows the number of successful fits for each setup.

	Brier	Abs res	Sd res	Dev	Sd dev	# succesful fits
static	0.054	0.055	0.218	-0.058	0.564	100
Extra correction	0.047	0.035	0.205	-0.037	0.519	94
Single correction	0.052	0.043	0.215	-0.045	0.569	99
2 order EKF	0.049	0.047	0.209	-0.049	0.605	98
Fixed E-step	0.053	0.044	0.214	-0.046	0.696	100
Fixed M-step	0.051	0.028	0.212	-0.029	0.600	100
UKF	0.049	0.038	0.209	-0.040	0.535	100
gam	0.048	0.036	0.207	-0.037	0.525	100

Std of deviance residuals w/ 1000 series



Conclussion on run

The main interest here is how the models labeled Fixed ... The results though are similar to what we saw before.

Incorrect binning time

Now, what happens if we get the binning (intervals lengths) wrong? This is the next experiment we will perform. Specifically, we will set the bin length to 0.1 instead 1 when we simulate. Thus, coefficients are updated at time $0, 0.1, 0.2, \ldots$ and whether an individual dies is evaluated at the same times when we simulate. However, the fitted model will still be based on bins of length 1.

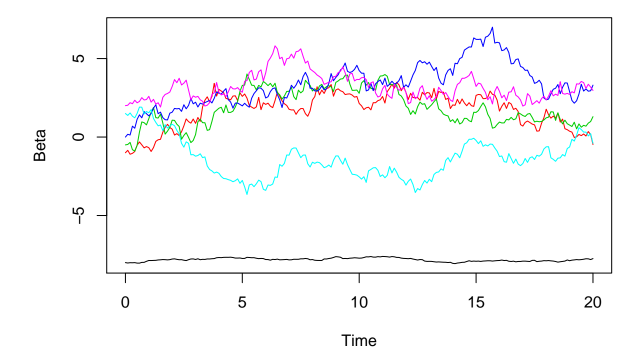
** Definition of simulation function

```
set.seed(9001)
sim_finer_binning <- function(n){
  time_denom = 10 # how much finer do we want to bin?

res <- test_sim_func_logit(
    n_series = n,

# We multiply through appropriately</pre>
```

* Illustration of single simulation



Simulating

We are now able to simulate with the following call:

```
# Use simulation function
set.seed(747)
simulate_n_print_res(
    sim_func = sim_finer_binning,
    file_prefix = "logit_sim_diff_binning")
```

Table 5: Mean of metrics with 100 series in test and fit data. 'Abs res' is the median of the absolute residuals, 'Sd res' is the standard deviation of the residuals, 'Dev' is median of the deviance residuals and 'Sd dev' is the standard deviation of deviance residuals. Only simulations that succeeds for all setups are included. There are 85 of these simulations. The last column shows the number of successful fits for each setup.

	Brier	Abs res	Sd res	Dev	Sd dev	# succesful fits
static	0.031	0.027	0.170	-0.027	0.574	100
Extra correction	0.029	0.019	0.163	-0.020	0.509	85
Single correction	0.031	0.026	0.167	-0.026	0.478	100
2 order EKF	0.031	0.027	0.167	-0.027	1.262	99
Fixed E-step	0.031	0.033	0.168	-0.034	0.514	100
Fixed M-step	0.031	0.020	0.167	-0.020	0.563	100

	Brier	Abs res	Sd res	Dev	Sd dev	# succesful fits
UKF	0.029	0.023	0.165	-0.023	0.518	100
gam	0.030	0.012	0.165	-0.012	0.733	100

Std of deviance residuals w/ 100 series

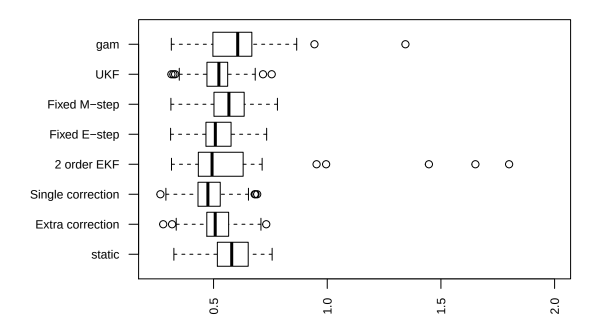
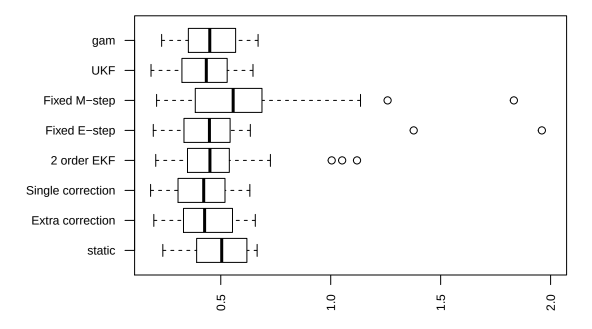


Table 6: Mean of metrics with 1000 series in test and fit data. 'Abs res' is the median of the absolute residuals, 'Sd res' is the standard deviation of the residuals, 'Dev' is median of the deviance residuals and 'Sd dev' is the standard deviation of deviance residuals. Only simulations that succeeds for all setups are included. There are 90 of these simulations. The last column shows the number of successful fits for each setup.

	Brier	Abs res	Sd res	Dev	Sd dev	# successful fits
static	0.030	0.029	0.151	-0.030	0.494	100
Extra correction	0.026	0.015	0.142	-0.016	0.436	92
Single correction	0.030	0.022	0.151	-0.023	0.414	100
2 order EKF	0.028	0.023	0.146	-0.024	0.678	97
Fixed E-step	0.030	0.025	0.151	-0.026	0.620	100
Fixed M-step	0.030	0.015	0.152	-0.015	0.578	99
UKF	0.027	0.019	0.145	-0.020	0.427	100
gam	0.026	0.015	0.143	-0.016	0.456	100

Std of deviance residuals w/ 1000 series



Conclussion on run

The UKF and extra iteration seems to perform well in both settings in terms of Brier score.

Out-of-time prediction

In the following paragraphs, we will investigate how the different estimation method performs when the following period have to be predicted. Thus, we cannot use the GAM model because it uses in-sample splines. Though, we can still use the state-space models as we can predict the following state vector given the previous. Further, we can use the static model to compare with.

** Define simulation and data splitting function

We start by defining a simulation function and a function to split the data into the first time period which we will use for estimation and the later time period which we will use for the test:

```
# Define simulation function
out_sample_args <- default_args
out_sample_args$t_max <- 21

sim_func <- function(n_series = 200)
    do.call(test_sim_func_logit, c(list(n_series = n_series), out_sample_args))</pre>
```

```
# Define split function
split_data_func <- function(d_frame, split_time = 20){
    # Find data before split_time and set event flag and stop time
    in_sample <- d_frame[d_frame$tstart < split_time, ]
    in_sample$event <- in_sample$event & in_sample$tstop <= split_time
    in_sample$tstop <- pmin(in_sample$tstop, split_time)

# Find data that ends after split_time and set start time
    out_sample <- d_frame[split_time < d_frame$tstop, ]
    out_sample$tstart <- pmax(out_sample$tstart, split_time)

# Return
    list(in_sample = in_sample, out_sample = out_sample)
}</pre>
```

We extend the period (t_max) by one which is the only difference in the simulation. Notice that individuals can be in both estimation data and test data. Any failure beyond time 20 will only count as a failure in the test data. Thus, we need to change the event flag for these in the in_sample data if the stop time is beyond time 20. Below, we illustrate how this looks for an individual who do die beyond time 20:

```
# Illustrate with example
set.seed(1119)
tmp <- sim_func()</pre>
# Illustrate for individual 25
tmp$res[tmp$res$id == 25, ]
##
       id tstart tstop event
                                         x2
                                                 x3
                                                         x4
                                                                 x5
                                 x1
## 146 25
            0.00 1.31
                           0 0.6627 0.98270 0.5245 0.86038 0.04009
## 147 25
            1.31
                  4.33
                           0 0.8070 0.20333 0.8489 0.23904 0.14597
## 148 25
            4.33 6.06
                           0 0.2007 0.88774 0.8260 0.96500 0.62327
## 149 25
            6.06 7.29
                           0 0.1358 0.46823 0.1891 0.01255 0.04520
                           0 0.3609 0.74557 0.8190 0.67799 0.04683
## 150 25
            7.29 9.24
## 151 25
            9.24 12.00
                           0 0.2015 0.69450 0.5960 0.07063 0.08570
## 152 25
           12.00 13.66
                           0 0.9212 0.09218 0.5132 0.54526 0.90543
## 153 25
                           0 0.6043 0.69848 0.4787 0.92423 0.84058
           13.66 15.59
          15.59 16.61
## 154 25
                           0 0.1504 0.14431 0.9336 0.10567 0.98583
## 155 25
           16.61 19.48
                           0 0.6359 0.04438 0.6532 0.85826 0.17005
## 156 25
          19.48 21.00
                           1 0.6404 0.52934 0.3124 0.70302 0.72475
# Split data
d split <- split data func(tmp$res)</pre>
# In sample data (notice event flag is changed and last tstop)
d_split$in_sample[d_split$in_sample$id == 25, ]
       id tstart tstop event
                                 x1
                                         x2
                                                 xЗ
                                                                 x5
## 146 25
            0.00 1.31 FALSE 0.6627 0.98270 0.5245 0.86038 0.04009
## 147 25
            1.31 4.33 FALSE 0.8070 0.20333 0.8489 0.23904 0.14597
## 148 25
            4.33 6.06 FALSE 0.2007 0.88774 0.8260 0.96500 0.62327
## 149 25
            6.06 7.29 FALSE 0.1358 0.46823 0.1891 0.01255 0.04520
## 150 25
            7.29
                  9.24 FALSE 0.3609 0.74557 0.8190 0.67799 0.04683
## 151 25
            9.24 12.00 FALSE 0.2015 0.69450 0.5960 0.07063 0.08570
           12.00 13.66 FALSE 0.9212 0.09218 0.5132 0.54526 0.90543
## 152 25
## 153 25
           13.66 15.59 FALSE 0.6043 0.69848 0.4787 0.92423 0.84058
```

```
## 154 25 15.59 16.61 FALSE 0.1504 0.14431 0.9336 0.10567 0.98583
## 155 25 16.61 19.48 FALSE 0.6359 0.04438 0.6532 0.85826 0.17005
## 156 25 19.48 20.00 FALSE 0.6404 0.52934 0.3124 0.70302 0.72475
# Out sample data (notice tstart is changed)
d_split$out_sample[d_split$out_sample$id == 25, ]
##
       id tstart tstop event
                                x1
                                       x2
                                              x3
                                                    x4
                                                           x5
## 156 25
             20
                   21
                       1 0.6404 0.5293 0.3124 0.703 0.7247
```

Simulation

We can now run the simulation with the following code. We end the code by printing the mean Brier score for the test data:

```
# Setup
N <- 100
                                              # number of simulations
n <- 1000
                                              # number of series
out <- matrix(NA_real_, nrow = N, ncol = 4) # matrix for output
# Run simulation
set.seed(42)
for(i in 1:N){
  # Simulate data and split
 repeat{
    sims <- sim func(n)
    # We want some survivers and some deaths
    if(sum(sims$res$event) > 50 && n - sum(sims$res$event) > 50)
      break
  }
  d_split <- split_data_func(sims$res)</pre>
  # Estimate models
  static_fit <- fit_funcs$static(d_split$in_sample)</pre>
  ekf_fit <- fit_funcs$dd(d_split$in_sample)</pre>
  ekf_extra_fit <- fit_funcs$dd(d_split$in_sample, NR_eps = .01)</pre>
  ukf_fit <- fit_funcs$dd_UKF(d_split$in_sample)</pre>
  # Predict outcome
  error <- list(
     static =
       predict(static_fit, d_split$out_sample, type = "response"),
     ekf = if(is.na(ekf_fit)) NA else
       predict(ekf_fit, new_data = d_split$out_sample,
               tstart = "tstart", tstop = "tstop")$fits,
     ekf_extra = if(is.na(ekf_extra_fit)) NA else
       predict(ekf_extra_fit, new_data = d_split$out_sample,
               tstart = "tstart", tstop = "tstop")$fits,
     ukf = if(is.na(ukf_fit)) NA else
       predict(ukf_fit, new_data = d_split$out_sample,
```

```
tstart = "tstart", tstop = "tstop")$fits)
  # Compute Brier score
  error <- unlist(lapply(</pre>
    error, function(x) if(is.na(x)) NA else
      mean.default((x - d_split$out_sample$event)^2)))
  # Save results
  out[i, ] <- error</pre>
# Print mean for cases where all could fit
colnames(out) <- c("Static", "EKF", "EKF with extra correction", "UKF")</pre>
colMeans(out[complete.cases(out), ])
##
                       Static
                                                      EKF
##
                      0.05389
                                                 0.04472
## EKF with extra correction
                                                      UKF
##
                      0.04484
                                                 0.04510
# Print median
apply(out[complete.cases(out), ], 2, median)
                                                      EKF
##
                       Static
                      0.02496
                                                 0.02463
## EKF with extra correction
                                                      UKF
                      0.02465
##
                                                 0.02493
# Print number of cases where all methods succeed to estimate
sum(complete.cases(out))
```

[1] 100

Above, we do 100 simulations with 1000 series in each simulation. It seems to that the different EKF methods and the UKF performs comparably. Another question is how often the various method got a given rank within a simulation in terms of their Brier score. We answer this question below (the rank are given as the first printed value such that one implies being the lowest Brier score in a given simulation):

Table 7: Number of times each set got a given rank in terms of Brier Score

	Static	EKF	EKF with extra correction	UKF
1	17	34	29	20
2	5	31	42	22
3	5	22	26	47
4	73	13	3	11

The EKF method does better with these specification in terms of getting the lowest mean out-sample Brier score and getting the lowest Brier score in most of the simulation.

Linear Time complexity

We will illustrate that the EKF and UKF have linear time complexity in the number of observation. This is particularly easy because the simulation function start of by simulating the coefficients as shown below (hence, variation will not be due to different coefficients vectors and only the number of series):

```
some_seed <- 69284
set.seed(some_seed)
res_1 <- test_sim_func_logit(100)

set.seed(some_seed)
res_2 <- test_sim_func_logit(1000) # different number of series
all.equal(res_1$betas, res_2$betas) # Coeffecients are equal</pre>
```

[1] TRUE

Next, we plot the computation time versus the number of simulation for the EKF and UKF method. Further, we print the linear regression slope for the log-log regression. The slope is close to one implying that the linear time complexity is linear in the number of observations

```
# Define function to record run time for a given number of series
run_time_func <- function(n, sim_args = default_args){</pre>
  set.seed(7851348) # Use the same seed
  sim_args$n_series <- n</pre>
  sims <- do.call(test_sim_func_logit, sim_args)</pre>
  time_EKF <- system.time(fit_EKF <- fit_funcs$dd(sims$res))</pre>
  time UKF <- system.time(</pre>
    fit_UKF <- ddhazard(</pre>
      formula = Surv(tstart, tstop, event) \sim x1 + x2 + x3 + x4 + x5,
      data = sims$res, max_T = T_max, by = 1, id = sims$res$id,
      Q_0 = diag(.1, n_beta + 1), Q = diag(.1, n_beta + 1),
      control = list(
        eps = 0.1,
        alpha = 1,
        beta = 0,
        method = "UKF")))
  # Check that both succed to fit
  if(is.na(fit_EKF) || is.na(fit_UKF))
    stop()
  list(time_EKF = time_EKF, time_UKF = time_UKF)
n_{for_{test}} <- 2^{(10:19)}
run_time <- sapply(n_for_test, run_time_func)</pre>
# Plot EKF and print log-log regression slope
ekf_time <- sapply(run_time["time_EKF", ], function(x) x[["user.self"]])</pre>
plot(n_for_test, ekf_time, type = "p", log = "xy",
     xlab = "Number of series", ylab = "Computation time for EKF")
```

