# Package 'dynamichazard'

October 14, 2019

Type Package

Title Dynamic Hazard Models using State Space Models

Version 0.6.6

Description Contains functions that lets you fit dynamic hazard models using state space models. The first implemented model is described in Fahrmeir (1992) <doi:10.1080/01621459.1992.10475232> and Fahrmeir (1994)
 <doi:10.1093/biomet/81.2.317>. Extensions hereof are available where the Extended Kalman filter is replaced by an unscented Kalman filter and other options including particle filters. The implemented particle filters support more general state space models.

License GPL-2

LazyData TRUE

LinkingTo Rcpp, RcppArmadillo

**Imports** parallel, Rcpp (>= 0.12.6), boot

**Depends** R (>= 3.5.0), stats, graphics, utils, survival

RoxygenNote 6.1.1

**Suggests** testthat, knitr, rmarkdown, timereg, captioner, biglm, httr, mgcv, shiny, formatR, R.rsp, speedglm, dichromat, colorspace, plyr, gsl, mvtnorm, nloptr (>= 1.2.0)

VignetteBuilder knitr, R.rsp

BugReports https://github.com/boennecd/dynamichazard/issues

SystemRequirements C++11

URL https://github.com/boennecd/dynamichazard

**Encoding** UTF-8

NeedsCompilation yes

Author Benjamin Christoffersen [cre, aut], Alan Miller [cph], Anthony Williams [cph], Boost developers [cph], R-core [cph] Maintainer Benjamin Christoffersen <boennecd@gmail.com>

**Repository** CRAN

Date/Publication 2019-10-14 08:50:02 UTC

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ddFixed

Auxiliary Functions for Fixed Effects

# Description

Functions used in formula of ddhazard for time-invariant effects. ddFixed\_intercept is only used for the intercept.

# ddhazard

#### Usage

ddFixed(object)

```
ddFixed_intercept(random_intercept = FALSE)
```

#### Arguments

object expression that would be used in formula. E.g. x or poly(x,degree = 3).
random\_intercept

TRUE if a zero mean time-varying process should be included at as an additional term. Only relevant in stationary models. See the type argument in PF\_EM.

# Examples

```
# we can get a time-invariant effect of `x1` by
## Not run:
ddhazard(Surv(stop, event) ~ ddFixed(x1), data)
## End(Not run)
# all of the calls below will yield the same result with a time-invariant
# intercept:
## Not run:
ddhazard(Surv(stop, event) ~ ddFixed_intercept() + x1, data)
ddhazard(Surv(stop, event) ~ -1 + ddFixed_intercept() + x1, data)
```

## End(Not run)

ddhazard

Fitting Dynamic Hazard Models

#### Description

Function to fit dynamic hazard models using state space models.

# Usage

```
ddhazard(formula, data, model = "logit", by, max_T, id, a_0, Q_0,
    Q = Q_0, order = 1, weights, control = ddhazard_control(),
    verbose = F)
```

formula	coxph like formula with Surv(tstart,tstop,event) on the left hand site of
	~.
data	data.frame or environment containing the outcome and covariates.

model	"logit", "cloglog", or "exponential" for respectively the logistic link func- tion with discrete outcomes, the inverse cloglog link function with discrete out- comes, or for the continuous time model with piecewise constant exponentially distributed arrival times.
by	interval length of the bins in which parameters are fixed.
max_T	end of the last interval interval.
id	vector of ids for each row of the in the design matrix.
a_0	vector $a_0$ for the initial coefficient vector for the first iteration (optional). Default is estimates from static model (see static_glm).
Q_0	covariance matrix for the prior distribution.
Q	initial covariance matrix for the state equation.
order	order of the random walk.
weights	weights to use if e.g. a skewed sample is used.
control	list of control variables from ddhazard_control.
verbose	TRUE if you want status messages during execution.

# Details

This function can be used to estimate survival models where the regression parameters follows a given order random walk. The order is specified by the order argument. 1. and 2. order random walks is implemented. The regression parameters are updated at time by, 2by, ..., max\_T. See the vignette("ddhazard", "dynamichazard") for details.

All filter methods needs a state covariance matrix  $Q_0$  and state vector  $a_0$ . An estimate from a time-invariant model is used for  $a_0$  if it is not supplied (the same model you would get from static\_glm). A diagonal matrix with large entries is recommended for  $Q_0$ . What is large dependents on the data set and model. Further, a covariance matrix for the first iteration Q is needed. The Q and  $a_0$  are estimated with an EM-algorithm.

The model is specified through the model argument. The discrete outcome models are where outcomes are binned into the intervals. Be aware that there can be "loss" of information due to binning if outcomes are not discrete to start with. It is key for these models that the id argument is provided if individuals in the data set have time-varying covariates. The the exponential model use a piecewise constant exponential distribution for the arrival times where there is no "loss" information due to binning. Though, one of the assumptions of the model is not satisfied if outcomes are only observed in discrete time intervals.

It is recommended to see the Shiny app demo for this function by calling ddhazard\_app().

# Value

A list with class ddhazard. The list contains

formula	the passed formula.
call	the matched call.
state_vecs	2D matrix with the estimated state vectors (regression parameters) in each bin.
state_vars	3D array with smoothed variance estimates for each state vector.

#### ddhazard

lag_one_cov	3D array with lagged correlation matrix for each for each change in the state vector. Only present when the model is logit and the method is EKF.
n_risk	the number of observations in each interval.
times	the interval borders.
risk_set	the object from get_risk_obj if saved.
data	the data argument if saved.
weights	weights used in estimation if saved.
id	ids used to match rows in data to individuals.
order	order of the random walk.
F_	matrix which map from one state vector to the next.
method	method used in the E-step.
est_Q_0	TRUE if Q_0 was estimated in the EM-algorithm.
family	$Rcpp \ {\tt Module} \ with \ C++ \ functions \ used \ for \ estimation \ given \ the \ {\tt model} \ argument.$
discrete_hazard	_func
	the hazard function corresponding to the model argument.
terms	the terms object used.
has_fixed_inter	cept
	TRUE if the model has a time-invariant intercept.
xlev	a record of the levels of the factors used in fitting.

# References

Fahrmeir, Ludwig. *Dynamic modelling and penalized likelihood estimation for discrete time survival data*. Biometrika 81.2 (1994): 317-330.

Durbin, James, and Siem Jan Koopman. *Time series analysis by state space methods*. No. 38. Oxford University Press, 2012.

# See Also

plot, residuals, predict, static\_glm, ddhazard\_app, ddhazard\_boot

# Examples

```
# example with first order model
library(dynamichazard)
fit <- ddhazard(
  Surv(time, status == 2) ~ log(bili), pbc, id = pbc$id, max_T = 3600,
  Q_0 = diag(1, 2), Q = diag(1e-4, 2), by = 50,
  control = ddhazard_control(method = "GMA"))
  plot(fit)
# example with second order model
fit <- ddhazard(
  Surv(time, status == 2) ~ log(bili), pbc, id = pbc$id, max_T = 3600,
  Q_0 = diag(1, 4), Q = diag(1e-4, 2), by = 50,
```

```
control = ddhazard_control(method = "GMA"),
order = 2)
plot(fit)
```

ddhazard\_app ddhazard Demo

# Description

ddhazard\_app runs a shiny app with demonstration of models.

# Usage

```
ddhazard_app(quietly = F, ...)
```

# Arguments

quietly	TRUE if no messages should be printed when the app is run.
	starting values for the shiny app.

# Details

Runs a shiny app where you try different model specifications on simulated data.

# Examples

```
## Not run:
dynamichazard::ddhazard_app()
dynamichazard::ddhazard_app(seed = 1, more_options = TRUE)
```

## End(Not run)

ddhazard\_boot Bootstrap for ddhazard Object

#### Description

See the vignette vignette("Bootstrap\_illustration", "dynamichazard"). The do\_stratify\_with\_event may be useful when either cases or non-cases are very rare to ensure that the model estimation succeeds.

#### Usage

```
ddhazard_boot(ddhazard_fit, strata, unique_id, R = 100,
    do_stratify_with_event = F, do_sample_weights = F,
    LRs = ddhazard_fit$control$LR * 2^(0:(-4)), print_errors = F)
```

```
6
```

# Arguments

ddhazard_fit	returned object from a ddhazard call.	
strata	strata to sample within. These need to be on an individual by individual basis and not rows in the design matrix.	
unique_id	unique ids where entries match entries of strata.	
R	number of bootstrap estimates.	
do_stratify_with_event		
	TRUE if sampling should be by strata of whether the individual has an event. An interaction factor will be made if strata is provided.	
do_sample_weigh	ts	
	TRUE if weights should be sampled instead of individuals.	
LRs	learning rates in decreasing order which will be used to estimate the model.	
print_errors	TRUE if errors should be printed when estimations fails.	

#### Value

An object like from the **boot** function.

# See Also

ddhazard, plot

# Examples

```
## Not run:
library(dynamichazard)
set.seed(56219373)
fit <- ddhazard(
  Surv(time, status == 2) ~ log(bili), pbc, id = pbc$id, max_T = 3000,
  Q_0 = diag(1, 2), Q = diag(1e-4, 2), by = 100,
  control = ddhazard_control(method = "GMA"))
bt <- ddhazard_boot(fit, R = 999)
plot(fit, ddhazard_boot = bt, level = .9)
```

## End(Not run)

ddhazard\_control Auxiliary for Controlling Dynamic Hazard Models

#### Description

Auxiliary for additional settings with ddhazard.

# Usage

```
ddhazard_control(kappa = NULL, alpha = 1, beta = 0, NR_eps = NULL,
  LR = 1, n_max = 10^2, eps = 0.001, est_Q_0 = F, method = "EKF",
  save_risk_set = T, save_data = T, eps_fixed_parems = 1e-04,
  fixed_parems_start = NULL,
  n_threads = getOption("ddhazard_max_threads"), denom_term = 1e-05,
  fixed_terms_method = "E_step", Q_0_term_for_fixed_E_step = NULL,
  permu = if (!is.null(method)) method == "SMA" else F,
  posterior_version = "cholesky", GMA_max_rep = 25,
  GMA_NR_eps = 1e-04, est_a_0 = TRUE, ...)
```

kappa	hyper parameter $\kappa$ in the unscented Kalman Filter.
alpha	hyper parameter $\alpha$ in the unscented Kalman Filter.
beta	hyper parameter $\beta$ in the unscented Kalman Filter.
NR_eps	tolerance for the Extended Kalman filter. Default is NULL which means that no extra iteration is made in the correction step.
LR	learning rate.
n_max	maximum number of iteration in the EM-algorithm.
eps	tolerance parameter for the EM-algorithm
est_Q_0	TRUE if you want the EM-algorithm to estimate Q_0. Default is FALSE.
method	set to the method to use in the E-step. Either "EKF" for the Extended Kalman Filter, "UKF" for the Unscented Kalman Filter, "SMA" for the sequential posterior mode approximation method or "GMA" for the global mode approximation method. "EKF" is the default.
save_risk_set	TRUE if you want to save the list from get_risk_obj used to estimate the model. It may be needed for later calls to e.g., residuals, plot and logLike.
save_data	TRUE if you want to keep the data argument. It may be needed for later calls to e.g., residuals, plot and logLike.
eps_fixed_paren	nS
	tolerance used in the M-step of the Fisher's scoring algorithm for the fixed effects
fixed_parems_st	tart
	starting value for fixed terms.
n_threads	maximum number of threads to use.
<pre>denom_term fixed terms met</pre>	term added to denominators in either the EKF or UKF.
	the method used to estimate the fixed effects. Either 'M_step' or 'E_step' for estimation in the M-step or E-step respectively.
Q_0_term_for_fi	ixed_E_step
	the diagonal value of the initial covariance matrix, $Q_0$ , for the fixed effects if fixed effects are estimated in the E-step.

# get\_cloud\_means

permu	TRUE if the risk sets should be permutated before computation. This is TRUE
	by default for posterior mode approximation method and FALSE for all other
	methods.
posterior_versi	on
	the implementation version of the posterior approximation method. Either "woodbury" or "cholesky".
GMA_max_rep	maximum number of iterations in the correction step if method = 'GMA'.
GMA_NR_eps	tolerance for the convergence criteria for the relative change in the norm of the coefficients in the correction step if method = 'GMA'.
est_a_0	FALSE if the starting value of the state model should be fixed.
	additional undocumented arguments.

# Value

A list with components named as the arguments.

# See Also

ddhazard

get\_cloud\_means Compute Mean Estimates from Particle Cloud

#### Description

Computes the estimated means from a particle cloud.

#### Usage

```
get_cloud_means(object, ...)
## S3 method for class 'PF_EM'
get_cloud_means(object, ...)
## S3 method for class 'PF_clouds'
get_cloud_means(object, cov_index = NULL,
    type = c("smoothed_clouds", "forward_clouds", "backward_clouds"), ...)
```

# Arguments

object	object with class PF_EM or PF_clouds.
	named arguments to pass to the PF_clouds method.
cov_index	integer vector with indices of the random effect to include.
type	character with the type of cloud to compute means for.

# Value

A matrix which rows are time indices and columns are random effect indices.

get\_cloud\_quantiles Compute Quantile Estimates from Particle Cloud

# Description

Computes the estimated quantiles from a particle cloud.

#### Usage

```
get_cloud_quantiles(object, ...)
## S3 method for class 'PF_EM'
get_cloud_quantiles(object, ...)
## S3 method for class 'PF_clouds'
get_cloud_quantiles(object, cov_index = NULL,
    qlvls = c(0.05, 0.5, 0.95), type = c("smoothed_clouds",
    "forward_clouds", "backward_clouds"), ...)
```

# Arguments

object	object with class PF_EM or PF_clouds.
	named arguments to pass to the PF_clouds method.
cov_index	integer vector with indices of the random effect to include.
qlvls	numeric vector with values in $\left[0,1\right]$ with the quantiles to compute.
type	character with the type of cloud to compute quantiles for.

# Value

A 3 dimensional array where the first dimension is the quantiles, the second dimension is the random effect, and the third dimension is the time.

get_Q_0 Compute Time-Invariant Covariance Matrix	
--	--

# Description

Computes the invariant covariance matrix for a vector autoregression model.

# Usage

get\_Q\_0(Qmat, Fmat)

# get\_risk\_obj

#### Arguments

Qmat	covariance matrix in transition density.
Fmat	coefficients in transition density.

# Value

The invariant covariance matrix.

# Examples

```
Fmat <- matrix(c(.8, .4, .1, .5), 2, 2)
Qmat <- matrix(c( 1, .5, .5, 2), 2)
x1 <- get_Q_0(Qmat = Qmat, Fmat = Fmat)
x2 <- Qmat
for(i in 1:101)
    x2 <- tcrossprod(Fmat %*% x2, Fmat) + Qmat
stopifnot(isTRUE(all.equal(x1, x2)))</pre>
```

get\_risk\_obj Risk Set on an Equidistant Distant Grid

# Description

Get the risk set at each bin over an equidistant distant grid.

# Usage

```
get_risk_obj(Y, by, max_T, id, is_for_discrete_model = T,
    n_threads = 1, min_chunk = 5000)
```

by length of each bin.	
max_T last observed time.	
id vector with ids where entries match with outcomes Y. is_for_discrete_model	
TRUE if the model outcome is discrete. For example, a logit model is discr whereas what is is referred to as the exponential model in this package i dynamic model.	ete s a
n_threads set to a value greater than one to use mclapply to find the risk object.	
min_chunk minimum chunk size of ids to use when parallel version is used.	

# Value

a list with the following elements

risk_sets	list of lists with one for each bin. Each of the sub lists have indices that corre- sponds to the entries of Y that are at risk in the bin.
min_start	start time of the first bin.
I_len	length of each bin.
d	number of bins.
is_event_in	indices for which bin an observation $Y$ is an event. $\neg 1$ if the individual does not die in any of the bins.
is_for_discrete_model	
	value of is_for_discrete_model argument.

# Examples

```
# small toy example with time-varying covariates
dat <- data.frame(
    id = c(1, 1, 2, 2),
    tstart = c(0, 4, 0, 2),
    tstop = c(4, 6, 2, 4),
    event = c(0, 1, 0, 0))
with(dat, get_risk_obj(Surv(tstart, tstop, event), by = 1, max_T = 6, id = id))</pre>
```

# Description

Function used to get data. frame with weights for a static fit for survivals.

#### Usage

```
get_survival_case_weights_and_data(formula, data, by, max_T, id,
    init_weights, risk_obj, use_weights = T, is_for_discrete_model = T,
    c_outcome = "Y", c_weights = "weights", c_end_t = "t")
```

formula	coxph like formula with Surv(tstart,tstop,event) on the left hand site of
	~.
data	data.frame or environment containing the outcome and covariates.
by	interval length of the bins in which parameters are fixed.

max_T	end of the last interval interval.
id	vector of ids for each row of the in the design matrix.
init_weights	weights for the rows in data. Useful e.g., with skewed sampling.
risk_obj	a pre-computed result from a get_risk_obj. Will be used to skip some computations.
use_weights	TRUE if weights should be used. See details.
is_for_discrete	_model
	TRUE if the model is for a discrete hazard model is used like the logistic model.
c_outcome, c_wei	ights, c_end_t
	alternative names to use for the added columns described in the return section.
	Useful if you already have a column named Y, t or weights.

#### Details

This function is used to get the data.frame for e.g. a glm fit that is comparable to a ddhazard fit in the sense that it is a static version. For example, say that we bin our time periods into (0, 1], (1,2] and (2,3]. Next, consider an individual who dies at time 2.5. He should be a control in the the first two bins and should be a case in the last bin. Thus the rows in the final data frame for this individual is c(Y = 1, ..., weights = 1) and c(Y = 0, ..., weights = 2) where Y is the outcome, ... is the covariates and weights is the weights for the regression. Consider another individual who does not die and we observe him for all three periods. Thus, he will yield one row with c(Y = 0, ..., weights = 3).

This function use similar logic as the ddhazard for individuals with time varying covariates (see the vignette ("ddhazard", "dynamichazard") for details).

If use\_weights = FALSE then the two previously mentioned individuals will yield three rows each. The first individual will have c(Y = 0, t = 1, ..., weights = 1), c(Y = 0, t = 2, ..., weights = 1), c(Y = 1, t = 3, ..., weights = 1) while the latter will have three rows <math>c(Y = 0, t = 1, ..., weights = 1), c(Y = 0, t = 2, ..., weights = 1), c(Y = 0, t = 3, ..., weights = 1). This kind of data frame is useful if you want to make a fit with e.g. gam function in the mgcv package as described en Tutz et. al (2016).

#### Value

Returns a data.frame where the following is added (column names will differ if you specified them): column Y for the binary outcome, column weights for weights of each row and additional rows if applicable. A column t is added for the stop time of the bin if use\_weights = FALSE. An element Y with the used Surv object is added if is\_for\_discrete\_model = FALSE.

#### References

Tutz, Gerhard, and Matthias Schmid. *Nonparametric Modeling and Smooth Effects*. Modeling Discrete Time-to-Event Data. Springer International Publishing, 2016. 105-127.

#### See Also

ddhazard, static\_glm

# Examples

```
library(dynamichazard)
# small toy example with time-varying covariates
dat <- data.frame(</pre>
       = c( 1,
 id
                    1, 2,
                               2),
 tstart = c(0,
                    4, 0,
                              2),
 tstop = c(4,
                    6, 2,
                              6),
 event = c(0,
                    1, 0,
                              0),
       = c(1.09, 1.29, 0, -1.16))
 x1
get_survival_case_weights_and_data(
 Surv(tstart, tstop, event) ~ x1, dat, by = 1, id = dat$id)$X
get_survival_case_weights_and_data(
 Surv(tstart, tstop, event) ~ x1, dat, by = 1, id = dat$id,
 use_weights = FALSE)$X
```

hatvalues.ddhazard Hat Values for ddhazard Object

#### Description

Computes hat-"like" values from usual L2 penalized binary regression.

# Usage

## S3 method for class 'ddhazard'
hatvalues(model, ...)

#### Arguments

model	a fit from ddhazard
	not used.

# Details

Computes hat-"like" values in each interval for each individual at risk in the interval. See the vignette("ddhazard", "dynamichazard") vignette for details.

# Value

A list of matrices. Each matrix has three columns: the hat values, the row number of the original data point and the id the row belongs to.

# See Also

ddhazard

# Examples

hds

```
library(dynamichazard)
fit <- ddhazard(
  Surv(time, status == 2) ~ log(bili), pbc, id = pbc$id, max_T = 3000,
  Q_0 = diag(1, 2), Q = diag(1e-4, 2), by = 100,
  control = ddhazard_control(method = "GMA"))
hvs <- hatvalues(fit)
head(hvs[[1]])
head(hvs[[2]])</pre>
```

hds

Hard Drive Failures

#### Description

A data set containing hard drive failures data from Backblaze in the start-stop format used in the survival package.

#### Usage

hds

# Format

A data. frame with the following columns:

serial\_number Serial number for the hard disk which the row belongs to.

model hard disk model.

manufacturer manufacturer of the hard disk model.

tstart,tstop start and stop times on the SMART 9 attribute scale.

fails 1 if the hard disk fails at tstop.

size\_tb hard disk size in terabytes.

smart\_x the raw SMART attribute x value. E.g., smart\_12 is the power cycle count.

**smart\_x\_bin** 1 if the SMART attribute x value is non-zero.

...\_cumsum cumulative sum of the prefix ....

**n\_fails** number of failures in the original data. Hard disk should only fail once but this is not the case in the raw data.

**n\_records** number of records in the original source.

min\_date,max\_date first and last date in the original source.

min\_hours,max\_hours smallest and largest value of the SMART 9 attribute in the original source.

#### Details

Details about the the SMART attributes can be found on https://en.wikipedia.org/wiki/S.M. A.R.T.. As stated in the original source

"Reported stats for the same SMART stat can vary in meaning based on the drive manufacturer and the drive model. Make sure you are comparing apples-to-apples as drive manufacturers don't generally disclose what their specific numbers mean."

There are some notes on <a href="https://en.wikipedia.org/wiki/S.M.A.R.T">https://en.wikipedia.org/wiki/S.M.A.R.T</a>. regarding which attributes that have vendor specific raw value. Further,

"The values in the files are the values reported by the drives. Sometimes, those values are out of whack. For example, in a few cases the RAW value of SMART 9 (Drive life in hours) reported a value that would make a drive 10+ years old, which was not possible. In other words, it's a good idea to have bounds checks when you process the data."

See this github page for the processing steps https://github.com/boennecd/backblaze\_survival\_ analysis\_prep.

#### Source

Raw data from https://www.backblaze.com/b2/hard-drive-test-data.html. Data have been processed to get a start-stop data.frame format.

logLik.ddhazard Log Likelihood of Mean Path of ddhazard Object

#### Description

Computes the log likelihood of (a potentially new) data set given the estimated:

```
E_{\theta}(\alpha_1|y_{1:d}), E_{\theta}(\alpha_2|y_{1:d}), \dots, E_{\theta}(\alpha_d|y_{1:d})
```

of the ddhazard object. Note that this is not the log likelihood of the observed data given the outcome.

# Usage

## S3 method for class 'ddhazard'
logLik(object, data = NULL, id, ...)

#### Arguments

object	an object of class ddhazard.
data	new data to evaluate the likelihood for.
id	the individual identifiers as in ddhazard.
	unused.

# logLik.PF\_EM

#### Examples

```
library(dynamichazard)
fit <- ddhazard(
  Surv(time, status == 2) ~ log(bili), pbc, id = pbc$id, max_T = 3600,
  Q_0 = diag(1, 2), Q = diag(1e-4, 2), by = 50,
  control = ddhazard_control(method = "GMA"))
logLik(fit)</pre>
```

logLik.PF\_EM

Approximate Log-Likelihood from a Particle Filter

# Description

Computes the approximate log-likelihood using the forward filter clouds. See the vignette("Particle\_filtering", "dyna for details.

#### Usage

```
## S3 method for class 'PF_EM'
logLik(object, ...)
## S3 method for class 'PF_clouds'
logLik(object, df = NA_real_, nobs = NA_integer_,
```

...)

# Arguments

object	an object of class PF_clouds or PF_EM.
	unused.
df	degrees of freedom used in the model.
nobs	integer with number of individuals used to estimate the model.

# Value

The approximate log-likelihood value given the observed data and set of parameter used when simulating the clouds. An attribute " $P(y_t|y_{1:(t-1)})$ " has the  $P(y_t|y_{1:(t-1)})$  terms.

PF\_control

# Description

Auxiliary for additional settings with PF\_EM.

# Usage

```
PF_control(N_fw_n_bw = NULL, N_smooth = NULL, N_first = NULL,
eps = 0.01, forward_backward_ESS_threshold = NULL,
method = "AUX_normal_approx_w_cloud_mean", n_max = 25,
n_threads = getOption("ddhazard_max_threads"),
smoother = "Fearnhead_O_N", Q_tilde = NULL, est_a_0 = TRUE,
N_smooth_final = N_smooth, nu = 0L, covar_fac = -1,
ftol_rel = 1e-08, averaging_start = -1L, fix_seed = TRUE)
```

N_fw_n_bw	number of particles to use in forward and backward filter.
N_smooth	number of particles to use in particle smoother.
N_first	number of particles to use at time 0 and time $d + 1$ .
eps	convergence threshold in EM method.
forward_backwar	d_ESS_threshold
	required effective sample size to not re-sample in the particle filters.
method	method for forward, backward and smoothing filter.
n_max	maximum number of iterations of the EM algorithm.
n_threads	maximum number threads to use in the computations.
smoother	smoother to use.
Q_tilde	covariance matrix of additional error term to add to the proposal distributions. NULL implies no additional error term.
est_a_0	FALSE if the starting value of the state model should be fixed. Does not apply for type = "VAR".
N_smooth_final	number of particles to sample with replacement from the smoothed particle cloud with $N_{smooth}$ particles using the particles' weights. This causes additional sampling error but decreases the computation time in the M-step.
nu	integer with degrees of freedom to use in the (multivariate) t-distribution used as the proposal distribution. A (multivariate) normal distribution is used if it is zero.
covar_fac	factor to scale the covariance matrix with. Ignored if the values is less than or equal to zero.
ftol_rel	relative convergence tolerance of the mode objective in mode approximation.

#### PF\_control

averaging_start	
	index to start averaging. Values less then or equal to zero yields no averaging.
fix_seed	TRUE if the same seed should be used. E.g., in PF_EM the same seed will be used in each iteration of the E-step of the MCEM algorithm.

#### Details

The method argument can take the following values

- bootstrap\_filter for a bootstrap filter.
- PF\_normal\_approx\_w\_cloud\_mean for a particle filter where a Gaussian approximation is used using a Taylor approximation made at the mean for the current particle given the mean of the parent particles and/or mean of the child particles.
- AUX\_normal\_approx\_w\_cloud\_mean for an auxiliary particle filter version of PF\_normal\_approx\_w\_cloud\_mean.
- PF\_normal\_approx\_w\_particles for a filter similar to PF\_normal\_approx\_w\_cloud\_mean and differs by making a Taylor approximation at a mean given each sampled parent and/or child particle.
- AUX\_normal\_approx\_w\_particles for an auxiliary particle filter version of PF\_normal\_approx\_w\_particles.

The smoother argument can take the following values

- Fearnhead\_0\_N for the smoother in Fearnhead, Wyncoll, and Tawn (2010).
- Brier\_0\_N\_square for the smoother in Briers, Doucet, and Maskell (2010).

#### Value

A list with components named as the arguments.

#### References

Gordon, N. J., Salmond, D. J., and Smith, A. F. (1993) Novel approach to nonlinear/non-Gaussian Bayesian state estimation. *In IEE Proceedings F (Radar and Signal Processing)*, (Vol. 140, No. 2, pp. 107-113). IET Digital Library.

Pitt, M. K., and Shephard, N. (1999) Filtering via simulation: Auxiliary particle filters. *Journal of the American statistical association*, **94(446)**, 590-599.

Fearnhead, P., Wyncoll, D., and Tawn, J. (2010) A sequential smoothing algorithm with linear computational cost. *Biometrika*, **97**(**2**), 447-464.

Briers, M., Doucet, A., and Maskell, S. (2010) Smoothing algorithms for state-space models. *Annals of the Institute of Statistical Mathematics*, **62(1)**, 61.

#### See Also

PF\_EM

# Description

Method to estimate the hyper parameters with an EM algorithm.

# Usage

```
PF_EM(formula, data, model = "logit", by, max_T, id, a_0, Q_0, Q,
order = 1, control = PF_control(...), trace = 0, seed = NULL,
type = "RW", fixed = NULL, random = NULL, Fmat, fixed_effects, G,
theta, J, K, psi, phi, ...)
```

formula	coxph like formula with Surv(tstart,tstop,event) on the left hand site of $\sim$ .
data	data.frame or environment containing the outcome and covariates.
model	either 'logit' for binary outcomes with the logistic link function, 'cloglog' for binary outcomes with the inverse cloglog link function, or 'exponential' for piecewise constant exponential distributed arrival times.
by	interval length of the bins in which parameters are fixed.
max_T	end of the last interval interval.
id	vector of ids for each row of the in the design matrix.
a_0	vector $a_0$ for the initial coefficient vector for the first iteration (optional). Default is estimates from static model (see static_glm).
Q_0	covariance matrix for the prior distribution.
Q	initial covariance matrix for the state equation.
order	order of the random walk.
control	see PF_control.
trace	argument to get progress information. Zero will yield no info and larger integer values will yield incrementally more information.
seed	seed to set at the start of every EM iteration. See set.seed.
type	type of state model. Either "RW" for a [R]andom [W]alk or "VAR" for [V]ector [A]uto[R]egression.
fixed	two-sided formula to be used with random instead of formula. It is of the form Surv(tstart,tstop,event) ~ x or Surv(tstart,tstop,event) ~ $-1$ for no fixed effects.
random	one-sided formula to be used with fixed instead of formula. It is of the form $\sim z$ .
Fmat	starting value for F when type = "VAR". See 'Details' in $PF\_EM$ .

# Details

Estimates a state model of the form

$$\alpha_t = F\alpha_t + R\epsilon_t, \qquad \epsilon_t \sim N(0, Q)$$

where  $F \in \mathbb{R}^{p \times p}$  has full rank,  $\alpha_t \in \mathbb{R}^p$ ,  $\epsilon_t \in \mathbb{R}^r$ ,  $r \leq p$ , and  $R = (e_{l_1}, e_{l_2}, \ldots, e_{l_r})$  where  $e_k$  is column from the p dimensional identity matrix and  $l_1 < l_2 < \ldots < l_r$ . The time zero state is drawn from

$$\alpha_0 \sim N(a_0, Q_0)$$

with  $Q_0 \in \mathbb{R}^{p \times p}$ . The latent states,  $\alpha_t$ , are related to the output through the linear predictors

$$\eta_{it} = X_t (R^+ \alpha_t) + Z_t \beta$$

where  $X_t \in \mathbb{R}^{n_t \times r}$  and  $Z_t \mathbb{R}^{n_t \times c}$  are design matrices and the outcome for a individual *i* at time *t* is distributed according to an exponential family member given  $\eta_{it}$ .  $\beta$  are constant coefficients. See vignette("Particle\_filtering", "dynamichazard") for details.

# Value

An object of class PF\_EM.

# Warning

The function is still under development so the output and API may change.

#### See Also

PF\_forward\_filter to get a more precise estimate of the final log-likelihood.

See the examples at https://github.com/boennecd/dynamichazard/tree/master/examples.

#### Examples

```
## Not run:
#####
# Fit model with lung data set from survival
# Warning: long-ish computation time
library(dynamichazard)
.lung <- lung[!is.na(lung$ph.ecog), ]</pre>
```

```
# standardize
.lung$age <- scale(.lung$age)</pre>
# fit
set.seed(43588155)
pf_fit <- PF_EM(</pre>
Surv(time, status == 2) ~ ddFixed(ph.ecog) + age,
data = .lung, by = 50, id = 1:nrow(.lung),
Q_0 = diag(1, 2), Q = diag(.5^2, 2),
max_T = 800,
control = PF_control(
    N_fw_n_bw = 500, N_first = 2500, N_smooth = 5000,
    n_max = 50, eps = .001, Q_tilde = diag(.2^2, 2), est_a_0 = FALSE,
    n_threads = max(parallel::detectCores(logical = FALSE), 1)))
# Plot state vector estimates
plot(pf_fit, cov_index = 1)
plot(pf_fit, cov_index = 2)
# Plot log-likelihood
plot(pf_fit$log_likes)
## End(Not run)
## Not run:
######
# example with fixed intercept
# prepare data
temp <- subset(pbc, id <= 312, select=c(id, sex, time, status, edema, age))</pre>
pbc2 <- tmerge(temp, temp, id=id, death = event(time, status))</pre>
pbc2 <- tmerge(pbc2, pbcseq, id=id, albumin = tdc(day, albumin),</pre>
              protime = tdc(day, protime), bili = tdc(day, bili))
pbc2 <- pbc2[, c("id", "tstart", "tstop", "death", "sex", "edema",</pre>
                 "age", "albumin", "protime", "bili")]
pbc2 <- within(pbc2, {</pre>
log_albumin <- log(albumin)</pre>
log_protime <- log(protime)</pre>
log_bili <- log(bili)</pre>
})
# standardize
for(c. in c("age", "log_albumin", "log_protime", "log_bili"))
pbc2[[c.]] <- drop(scale(pbc2[[c.]]))</pre>
# fit model with extended Kalman filter
ddfit <- ddhazard(</pre>
Surv(tstart, tstop, death == 2) ~ ddFixed_intercept() + ddFixed(age) +
   ddFixed(edema) + ddFixed(log_albumin) + ddFixed(log_protime) + log_bili,
 pbc2, Q_0 = 100, Q = 1e-2, by = 100, id = pbc2$id,
model = "exponential", max_T = 3600,
control = ddhazard_control(eps = 1e-5, NR_eps = 1e-4, n_max = 1e4))
summary(ddfit)
```

```
# fit model with particle filter
set.seed(88235076)
pf_fit <- PF_EM(</pre>
  Surv(tstart, tstop, death == 2) ~ ddFixed_intercept() + ddFixed(age) +
    ddFixed(edema) + ddFixed(log_albumin) + ddFixed(log_protime) + log_bili,
  pbc2, Q_0 = 2^2, Q = ddfit (* 100, # use estimate from before
  by = 100, id = pbc2$id,
  model = "exponential", max_T = 3600,
  control = PF_control(
   N_fw_n_bw = 500, N_smooth = 2500, N_first = 1000, eps = 1e-3,
   method = "AUX_normal_approx_w_cloud_mean", est_a_0 = FALSE,
   Q_tilde = as.matrix(.1^2),
   n_max = 25, # just take a few iterations as an example
   n_threads = max(parallel::detectCores(logical = FALSE), 1)))
# compare results
plot(ddfit)
plot(pf_fit)
sqrt(ddfit$Q * 100)
sqrt(pf_fit$Q)
rbind(ddfit$fixed_effects, pf_fit$fixed_effects)
## End(Not run)
## Not run:
#####
# simulation example with `random` and `fixed` argument and a restricted
# model
# g groups with k individuals in each
g <- 3L
k <- 400L
# matrices for state equation
p <- g + 1L
G <- matrix(0., p^2, 2L)
for(i in 1:p)
  G[i + (i - 1L) * p, 1L + (i == p)] <- 1L
theta <- c(.9, .8)
# coefficients in transition density
(F. <- matrix(as.vector(G %*% theta), 4L, 4L))</pre>
J \leq matrix(0., ncol = 2L, nrow = p)
J[-p, 1L] <- J[p, 2L] <- 1
psi <- c(log(c(.3, .1)))</pre>
K <- matrix(0., p * (p - 1L) / 2L, 2L)</pre>
j <- 0L
for(i in (p - 1L):1L){
  j <- j + i
 K[j, 2L] <- 1
}
K[K[, 2L] < 1, 1L] <- 1
```

```
phi <- \log(-(c(.8, .3) + 1) / (c(.8, .3) - 1))
V <- diag(exp(drop(J %*% psi)))</pre>
C \le diag(1, ncol(V))
C[lower.tri(C)] <- 2/(1 + exp(-drop(K %*% phi))) - 1
C[upper.tri(C)] <- t(C)[upper.tri(C)]</pre>
(Q <- V %*% C %*% V)
                          # covariance matrix in transition density
cov2cor(Q)
Q_0 <- get_Q_0(Q, F.) # time-invariant covariance matrix
beta <- c(rep(-6, g), 0) # all groups have the same long run mean intercept
# simulate state variables
set.seed(56219373)
n_periods <- 300L
alphas <- matrix(nrow = n_periods + 1L, ncol = p)</pre>
alphas[1L, ] <- rnorm(p) %*% chol(Q_0)
for(i in 1:n_periods + 1L)
  alphas[i, ] <- F. %*% alphas[i - 1L, ] + drop(rnorm(p) %*% chol(Q))</pre>
alphas <- t(t(alphas) + beta)</pre>
# plot state variables
matplot(alphas, type = "1", lty = 1)
# simulate individuals' outcome
n_{obs} <-g * k
df <- lapply(1:n_obs, function(i){</pre>
  # find the group
  grp <- (i - 1L) %/% (n_obs / g) + 1L
  # left-censoring
  tstart <- max(0L, sample.int((n_periods - 1L) * 2L, 1) - n_periods + 1L)</pre>
  # covariates
  x <- c(1, rnorm(1))</pre>
  # outcome (stop time and event indicator)
  osa <- NULL
  oso <- NULL
  osx <- NULL
  y <- FALSE
  for(tstop in (tstart + 1L):n_periods){
    sigmoid <- 1 / (1 + exp(- drop(x %*% alphas[tstop + 1L, c(grp, p)])))</pre>
    if(sigmoid > runif(1)){
      y <- TRUE
      break
    }
    if(.01 > runif(1L) && tstop < n_periods){</pre>
      # sample new covariate
      osa <- c(osa, tstart)
      tstart <- tstop
      oso <- c(oso, tstop)
```

```
osx <- c(osx, x[2])
     x[2] <- rnorm(1)
   }
 }
 cbind(
    tstart = c(osa, tstart), tstop = c(oso, tstop),
    x = c(osx, x[2]), y = c(rep(FALSE, length(osa)), y), grp = grp,
    id = i)
})
df <- data.frame(do.call(rbind, df))</pre>
df$grp <- factor(df$grp)</pre>
# fit model. Start with "cheap" iterations
fit <- PF_EM(</pre>
 fixed = Surv(tstart, tstop, y) ~ x, random = ~ grp + x - 1,
 data = df, model = "logit", by = 1L, max_T = max(df$tstop),
 Q_0 = diag(1.5<sup>2</sup>, p), id = df$id, type = "VAR",
 G = G, theta = c(.5, .5), J = J, psi = log(c(.1, .1)),
 K = K, phi = log(-(c(.4, 0) + 1) / (c(.4, 0) - 1)),
 control = PF_control(
   N_fw_n_bw = 100L, N_smooth = 100L, N_first = 500L,
   method = "AUX_normal_approx_w_cloud_mean",
   nu = 5L, # sample from multivariate t-distribution
   n_max = 100L, averaging_start = 50L,
   smoother = "Fearnhead_0_N", eps = 1e-4, covar_fac = 1.2,
   n_threads = 4L # depends on your cpu(s)
 ),
 trace = 1L)
plot(fit$log_likes) # log-likelihood approximation at each iterations
# take more iterations with more particles
cl <- fit$call</pre>
ctrl <- cl[["control"]]</pre>
ctrl[c("N_fw_n_bw", "N_smooth", "N_first", "n_max",
       "averaging_start")] <- list(500L, 2000L, 5000L, 200L, 30L)
cl[["control"]] <- ctrl</pre>
cl[c("phi", "psi", "theta")] <- list(fit$phi, fit$psi, fit$theta)</pre>
fit_extra <- eval(cl)</pre>
plot(fit_extra$log_likes) # log-likelihood approximation at each iteration
# check estimates
sqrt(diag(fit_extra$Q))
sqrt(diag(Q))
cov2cor(fit_extra$Q)
cov2cor(Q)
fit_extra$F
F.
# plot predicted state variables
for(i in 1:p){
 plot(fit_extra, cov_index = i)
```

```
abline(h = 0, lty = 2)
lines(1:nrow(alphas) - 1, alphas[, i] - beta[i], lty = 3)
}
## End(Not run)
```

PF\_forward\_filter Forward Particle Filter

#### Description

Functions to only use the forward particle filter. Useful for log-likelihood evaluation though there is an  $O(d^2)$  variance of the estimate where d is the number of time periods. The number of particles specified in the control argument has no effect.

The function does not alter the .Random.seed to make sure the same rng.kind is kept after the call. See PF\_EM for model details.

#### Usage

```
PF_forward_filter(x, N_fw, N_first, ...)
## S3 method for class 'PF_EM'
PF_forward_filter(x, N_fw, N_first, seed, ...)
## S3 method for class 'formula'
PF_forward_filter(x, N_fw, N_first, data,
    model = "logit", by, max_T, id, a_0, Q_0, Q, fixed_effects,
    control = PF_control(...), seed = NULL, trace = 0, G, theta, J, K,
    psi, phi, type = "RW", Fmat, ...)
## S3 method for class 'data.frame'
PF_forward_filter(x, N_fw, N_first, formula,
    model = "logit", by, max_T, id, a_0, Q_0, Q, fixed_effects,
    control = PF_control(...), seed = NULL, trace = 0, fixed = NULL,
    random = NULL, G, theta, J, K, psi, phi, type = "RW", Fmat,
    order = 1, ...)
```

#### Arguments

x	an PF_EM or formula object.
N_fw	number of particles.
N_first	number of time zero particles to draw.
	optional way to pass arguments to control.
seed	.GlobalEnv $\$ .Random.seed to set. Not seed as in set.seed function. Can be used with the .Random.seed returned by PF_EM.
data	data.frame or environment containing the outcome and covariates.

model	either 'logit' for binary outcomes with the logistic link function, 'cloglog' for binary outcomes with the inverse cloglog link function, or 'exponential' for piecewise constant exponential distributed arrival times.
by	interval length of the bins in which parameters are fixed.
max_T	end of the last interval interval.
id	vector of ids for each row of the in the design matrix.
a_0	vector $a_0$ for the initial coefficient vector for the first iteration (optional). Default is estimates from static model (see static_glm).
Q_0	covariance matrix for the prior distribution.
Q	initial covariance matrix for the state equation.
fixed_effects	values for the fixed parameters.
control	see PF_control.
trace	argument to get progress information. Zero will yield no info and larger integer values will yield incrementally more information.
G, theta, J, K, p	osi, phi
	parameters for a restricted type = "VAR" model. See the vignette mentioned in 'Details' of PF_EM and the examples linked to in 'See Also'.
type	type of state model. Either "RW" for a [R]andom [W]alk or "VAR" for [V]ector [A]uto[R]egression.
Fmat	starting value for $F$ when type = "VAR". See 'Details' in PF_EM.
formula	coxph like formula with $Surv(tstart,tstop,event)$ on the left hand site of ~.
fixed	two-sided formula to be used with random instead of formula. It is of the form Surv(tstart,tstop,event) ~ x or Surv(tstart,tstop,event) ~ $-1$ for no fixed effects.
random	one-sided formula to be used with fixed instead of formula. It is of the form $\sim z.$
order	order of the random walk.

#### Value

An object of class PF\_clouds.

# Methods (by class)

- PF\_EM: Forward particle filter using the estimates of an PF\_EM call.
- formula: Forward particle filter with formula input.
- data.frame: Forward particle filter with data.frame data input as x instead of data. Can be used with fixed and random argument.

# Warning

The function is still under development so the output and API may change.

#### Examples

```
## Not run:
# head-and-neck cancer study data. See Efron, B. (1988) doi:10.2307/2288857
is_censored <- c(</pre>
 6, 27, 34, 36, 42, 46, 48:51, 51 + c(15, 30:28, 33, 35:37, 39, 40, 42:45))
head_neck_cancer <- data.frame(</pre>
 id = 1:96,
 stop = c(
   1, 2, 2, rep(3, 6), 4, 4, rep(5, 8),
    rep(6, 7), 7, 8, 8, 8, 9, 9, 10, 10, 10, 11, 14, 14, 14, 15, 18, 18, 20,
    20, 37, 37, 38, 41, 45, 47, 47,
   2, 2, 3, rep(4, 4), rep(5, 5), rep(6, 5),
    7, 7, 7, 9, 10, 11, 12, 15, 16, 18, 18, 18, 21,
    21, 24, 25, 27, 36, 41, 44, 52, 54, 59, 59, 63, 67, 71, 76),
 event = !(1:96 %in% is_censored),
 group = factor(c(rep(1, 45 + 6), rep(2, 45))))
# fit model
set.seed(61364778)
ctrl <- PF_control(</pre>
 N_fw_n_bw = 500, N_smooth = 2500, N_first = 2000,
 n_max = 1, # set to one as an example
 n_threads = max(parallel::detectCores(logical = FALSE), 1),
 eps = .001, Q_tilde = as.matrix(.3<sup>2</sup>), est_a_0 = FALSE)
pf_fit <- suppressWarnings(</pre>
 PF_EM(
    survival::Surv(stop, event) ~ ddFixed(group),
    data = head_neck_cancer, by = 1, Q_0 = 1, Q = 0.1^2, control = ctrl,
   max_T = 30))
# the log-likelihood in the final iteration
(end_log_like <- logLik(pf_fit))</pre>
# gives the same
fw_ps <- PF_forward_filter(</pre>
 survival::Surv(stop, event) ~ ddFixed(group), N_fw = 500, N_first = 2000,
 data = head_neck_cancer, by = 1, Q_0 = 1, Q = 0.1^2,
 a_0 = pf_fit$a_0, fixed_effects = -0.5370051,
 control = ctrl, max_T = 30, seed = pf_fit$seed)
all.equal(c(end_log_like), c(logLik(fw_ps)))
# will differ since we use different number of particles
fw_ps <- PF_forward_filter(</pre>
 survival::Surv(stop, event) ~ ddFixed(group), N_fw = 1000, N_first = 3000,
 data = head_neck_cancer, by = 1, Q_0 = 1, Q = 0.1^2,
 a_0 = pf_fit$a_0, fixed_effects = -0.5370051,
  control = ctrl, max_T = 30, seed = pf_fit$seed)
all.equal(c(end_log_like), c(logLik(fw_ps)))
# will differ since we use the final estimates
fw_ps <- PF_forward_filter(pf_fit, N_fw = 500, N_first = 2000)</pre>
all.equal(c(end_log_like), c(logLik(fw_ps)))
```

## End(Not run)

PF\_get\_score\_n\_hess Approximate Observed Information Matrix and Score Vector

#### Description

Returns a list of functions to approximate the observed information matrix and score vector.

# Usage

```
PF_get_score_n_hess(object, debug = FALSE, use_0_n_sq = FALSE)
```

#### Arguments

object	object of class PF_EM.
debug	TRUE if debug information should be printed to the console.
use_0_n_sq	TRUE if the method from Poyiadjis et al. (2011) should be used.

#### Details

The score vector and observed information matrix are computed with the (forward) particle filter. This comes at an  $O(d^2)$  variance where d is the number of periods. Thus, the approximation may be poor for long series. The score vector can be used to perform stochastic gradient descent.

If use\_0\_n\_sq is TRUE then the method in Poyiadjis et al. (2011) is used. This may only have a variance which is linear in the number of time periods. However, the present implementation is  $O(N^2)$  where N is the number of particles. The method uses a particle filter as in Section 3.1 of Lin et al. (2005). There is no need to call run\_particle\_filter unless one wants a new approximation of the log-likelihood as a separate filter is run with get\_get\_score\_n\_hess when use\_0\_n\_sq is TRUE.

#### Value

A list with the following functions as elements

run\_particle\_filter

function to run particle filter as with PF\_forward\_filter.

set\_parameters function to set the parameters in the model. The first argument is a vectorized version of F matrix and Q matrix. The second argument is the fixed effect coefficients.

```
set_n_particles
```

sets the number of particles to use in run\_particle\_filter and get\_get\_score\_n\_hess when use\_0\_n\_sq is TRUE.

get\_get\_score\_n\_hess

approximate the observed information matrix and score vector. The argument toggles whether or not to approximate the observed information matrix. The last particle cloud from run\_particle\_filter is used when use\_0\_n\_sq is FALSE.

#### Warning

The function is still under development so the output and API may change.

#### References

Cappe, O. and Moulines, E. (2005) Recursive Computation of the Score and Observed Information Matrix in Hidden Markov Models. *IEEE/SP 13th Workshop on Statistical Signal Processing*.

Cappe, O., Moulines, E. and Ryden, T. (2005) Inference in Hidden Markov Models (Springer Series in Statistics). Springer-Verlag.

Doucet, A., and Tadić, V. B. (2003) Parameter Estimation in General State-Space Models Using Particle Methods. *Annals of the Institute of Statistical Mathematics*, **55**(2), 409–422.

Lin, M. T., Zhang, J. L., Cheng, Q. and Chen, R. (2005) Independent Particle Filters. *Journal of the American Statistical Association*, **100(472)**, 1412-1421.

Poyiadjis, G., Doucet, A. and Singh, S. S. (2011) Particle Approximations of the Score and Observed Information Matrix in State Space Models with Application to Parameter Estimation. *Biometrika*, **98(1)**, 65–80.

# See Also

See the examples at https://github.com/boennecd/dynamichazard/tree/master/examples.

#### Examples

```
## Not run:
library(dynamichazard)
.lung <- lung[!is.na(lung$ph.ecog), ]</pre>
# standardize
.lung$age <- scale(.lung$age)</pre>
# fit model
set.seed(43588155)
pf_fit <- PF_EM(</pre>
 fixed = Surv(time, status == 2) ~ ph.ecog + age,
 random = ~ 1, model = "exponential",
 data = .lung, by = 50, id = 1:nrow(.lung),
 Q_0 = as.matrix(1), Q = as.matrix(.5^2), type = "VAR",
 max_T = 800, Fmat = as.matrix(.5),
 control = PF_control(
   N_fw_n_bw = 250, N_first = 2000, N_smooth = 500, covar_fac = 1.1,
   nu = 6, n_max = 1000L, eps = 1e-4, averaging_start = 200L,
   n_threads = max(parallel::detectCores(logical = FALSE), 1)))
# compute score and observed information matrix
comp_obj <- PF_get_score_n_hess(pf_fit)</pre>
comp_obj$set_n_particles(N_fw = 10000L, N_first = 10000L)
comp_obj$run_particle_filter()
(01 <- comp_obj$get_get_score_n_hess())</pre>
# O(N^2) method with lower variance as a function of time
comp_obj <- PF_get_score_n_hess(pf_fit, use_0_n_sq = TRUE)</pre>
```

```
comp_obj$set_n_particles(N_fw = 2500L, N_first = 2500L)
(o2 <- comp_obj$get_get_score_n_hess())
# approximations may have large variance
o3 <- replicate(10L, {
    runif(1)
    pf_fit$seed <- .Random.seed
    comp_obj <- PF_get_score_n_hess(pf_fit)
    comp_obj$set_n_particles(N_fw = 10000L, N_first = 10000L)
    comp_obj$set_get_score_n_hess()
}, simplify = FALSE)
sapply(o3, function(x) x$score)
sapply(o3, function(x) sqrt(diag(solve(x$obs_info))))
## End(Not run)
```

plot.ddhazard Plots for ddhazard Object

# Description

Plot of estimated state space variables from a ddhazard fit.

# Usage

```
## S3 method for class 'ddhazard'
plot(x, xlab = "Time", ylab = "Hazard",
   type = "cov", plot_type = "l", cov_index, ylim, col = "black",
   add = F, do_alter_mfcol = T, level = 0.95, ddhazard_boot, ...)
```

Х	result of ddhazard call.	
xlab, ylab, ylim, col		
	arguments to override defaults set in the function.	
type	type of plot. Currently, only "cov" is available for plot of the state space parameters.	
plot_type	the type argument passed to plot.	
cov_index	the index (indices) of the state space parameter(s) to plot.	
add	FALSE if you want to make a new plot.	
do_alter_mfcol	TRUE if the function should alter $par(mfcol)$ in case that $cov_index$ has more than one element.	
level	level (fraction) for confidence bounds.	
ddhazard_boot	object from a ddhazard_boot call which confidence bounds will be based on and where bootstrap samples will be printed with a transparent color.	
	arguments passed to plot.default or lines depending on the value of add.	

#### Details

Creates a plot of state variables or adds state variables to a plot with indices cov\_index. Pointwise 1.96 std. confidence intervals are provided with the smoothed co-variance matrices from the fit.

# Examples

```
library(dynamichazard)
fit <- ddhazard(
  Surv(time, status == 2) ~ log(bili), pbc, id = pbc$id, max_T = 3600,
  Q_0 = diag(1, 2), Q = diag(1e-4, 2), by = 50,
  control = ddhazard_control(method = "GMA"))
plot(fit)
plot(fit, cov_index = 2)</pre>
```

plot.ddhazard\_space\_errors

State Space Error Plot

# Description

Plot function for state space errors from ddhazard fit.

# Usage

```
## S3 method for class 'ddhazard_space_errors'
plot(x, mod, cov_index = NA,
   t_index = NA, p_cex = par()$cex * 0.2, pch = 16,
   ylab = "Std. state space error", x_tick_loc = NA, x_tick_mark = NA,
   xlab = "Time", ...)
```

х	result of residuals with a 'type' argument which yields state space errors.
mod	the ddhazard result used in the residuals call.
cov_index	the indices of state vector errors to plot. Default is to use all.
t_index	the bin indices to plot. Default is to use all bins.
p_cex	cex argument for the points
pch, ylab, xlab	arguments to override defaults set in the function.
<pre>x_tick_loc, x_t</pre>	ick_mark
	at and labels arguments passed to axis.
	arguments passed to plot.default.

plot.ddsurvcurve Create and plot survival curves

# Description

The function creates a predicted survival curve for a new observation using a estimated ddhazard model from ddhazard. The predicted curve is based on the predicted mean path of the state vector. Thus, the survival curve will not be a "mean" curve due to the non-linear relation between the probability of an event and the state vector.

#### Usage

```
## S3 method for class 'ddsurvcurve'
plot(x, y, xlab = "Time", ylab = "Survival",
    ylim, xaxs = "i", yaxs = "i", ...)
## S3 method for class 'ddsurvcurve'
lines(x, col = "Black", lty = 1,
    lwd = par()$lwd, ...)
```

```
ddsurvcurve(object, new_data, tstart = "", tstop = "")
```

х	a ddsurvcurve object.
У	not used.
xlab	xlab passed to plot.
ylab	ylab passed to plot.
ylim	ylim passed to plot.
xaxs	xaxs passed to plot.
yaxs	yaxs passed to plot.
	not used.
col	col passed to lines.
lty	lty passed to lines.
lwd	lwd passed to lines.
object	a ddhazard object.
new_data	a data.frame with the new data for the observation who the survival curve should be for. It can have more rows if tstart and tstop is supplied. The rows need to be consecutive and non-overlapping time intervals.
tstart	name of the start time column in new_data. It must be on the same time scale as the tstart used in the Surv(tstart,tstop,event) in the formula passed to ddhazard.
tstop	same as tstart for the stop argument.

ddsurvcurve returns an object of class ddsurvcurve. It elements are the predicted discrete survival curve, time points for the survival curve, point of the first time period, the call, the discrete probabilities of an event in each interval conditional on survival up to that point, and the name of the distribution family. It should be seen as a plug-in estimate.

# Methods (by generic)

- plot: method for plotting survival curve.
- lines: Method for adding survival curve to a plot.

#### plot.ddsurvcurve

Returns the same as lines.ddsurvcurve.

#### lines.ddsurvcurve

Either returns the objects used in the call to segments for discrete time hazard models, or the time points and survival function used to draw the survival curve.

#### See Also

ddhazard, and predict.ddhazard.

#### Examples

```
#####
# example with continuous time model
# setup data set. See `vignette("timedep", "survival")`
library(dynamichazard)
temp <- subset(pbc, id <= 312, select=c(id:sex, stage))</pre>
pbc2 <- tmerge(temp, temp, id=id, death = event(time, status))</pre>
pbc2 <- tmerge(pbc2, pbcseq, id = id, bili = tdc(day, bili))</pre>
# fit model
f1 <- ddhazard(
  Surv(tstart, tstop, death == 2) ~ ddFixed(log(bili)), pbc2, id = pbc2$id,
  max_T = 3600, Q_0 = 1, Q = 1e-2, by = 100, model = "exponential",
  control = ddhazard_control(method = "EKF", eps = 1e-4, n_max = 1000,
                              fixed_terms_method = "M_step"))
# predict with default which is all covariates set to zero
ddcurve <- ddsurvcurve(f1)</pre>
par(mar = c(4.5, 4, .5, .5))
plot(ddcurve, col = "DarkBlue", lwd = 2)
# compare with cox model
f2 <- coxph(Surv(tstart, tstop, death == 2) ~ log(bili), data = pbc2)</pre>
nw <- data.frame(bili = 1, tstart = 0, tstop = 3000)</pre>
lines(survfit(f2, newdata = nw))
```

#### plot.ddsurvcurve

```
# same as above but with bili = 3
nw <- data.frame(bili = 3)</pre>
lines(ddsurvcurve(f1, new_data = nw), col = "DarkBlue")
lines(survfit(f2, newdata = nw))
# change to time-varying slope
f3 <- ddhazard(
 Surv(tstart, tstop, death == 2) ~ log(bili), pbc2, id = pbc2$id,
 max_T = 3600, Q_0 = diag(1, 2), Q = diag(1e-2, 2), by = 100, model = "exponential",
 control = ddhazard_control(method = "EKF", eps = 1e-4, n_max = 1000))
# example with time-varying coefficient
nw <- data.frame(</pre>
 bili = c(2.1, 1.9, 3.3, 3.9, 3.8, 3.6, 4, 4.9, 4.2, 5.7, 10.2),
 tstart = c(0L, 225L, 407L, 750L, 1122L, 1479L, 1849L, 2193L, 2564L, 2913L,
             3284L),
 tstop = c(225L, 407L, 750L, 1122L, 1479L, 1849L, 2193L, 2564L, 2913L,
            3284L, 3600L))
ddcurve <- ddsurvcurve(f3, new_data = nw, tstart = "tstart", tstop = "tstop")</pre>
lines(ddcurve, "darkorange", lwd = 2)
# can condition on survival up to some time
ddcurve <- ddsurvcurve(f3, new_data = nw[-(1:5), ], tstart = "tstart",</pre>
                       tstop = "tstop")
lines(ddcurve, lty = 2, lwd = 2)
#####
# example with discrete time model
# head-and-neck cancer study data. See Efron, B. (1988) doi:10.2307/2288857
is_censored <- c(</pre>
 6, 27, 34, 36, 42, 46, 48:51, 51 + c(15, 30:28, 33, 35:37, 39, 40, 42:45))
head_neck_cancer <- data.frame(</pre>
 id = 1:96,
 stop = c(
   1, 2, 2, rep(3, 6), 4, 4, rep(5, 8),
   rep(6, 7), 7, 8, 8, 8, 9, 9, 10, 10, 10, 11, 14, 14, 14, 15, 18, 18, 20,
   20, 37, 37, 38, 41, 45, 47, 47,
   2, 2, 3, rep(4, 4), rep(5, 5), rep(6, 5),
   7, 7, 7, 9, 10, 11, 12, 15, 16, 18, 18, 18, 21,
   21, 24, 25, 27, 36, 41, 44, 52, 54, 59, 59, 63, 67, 71, 76),
 event = !(1:96 %in% is_censored),
 group = factor(c(rep(1, 45 + 6), rep(2, 45))))
# fit model
h1 <- ddhazard(
 Surv(stop, event) ~ group, head_neck_cancer, by = 1, max_T = 45,
 Q_0 = diag(2^2, 2), Q = diag(.01^2, 2), control = ddhazard_control(
   method = "GMA", eps = 1e-4, n_max = 200))
# plot predicted survival curve. Notice the steps since the model is discrete
nw <- data.frame(group = factor(1, levels = 1:2), tstart = 0, tstop = 30)</pre>
ddcurve <- ddsurvcurve(h1, new_data = nw, tstart = "tstart",</pre>
                       tstop = "tstop")
```

plot.PF\_clouds Plot of Clouds from a PF\_clouds Object

# Description

Plots mean curve along with quantiles through time for the forward, backward or smoothed clouds.

#### Usage

```
## S3 method for class 'PF_clouds'
plot(x, y, type = c("smoothed_clouds",
    "forward_clouds", "backward_clouds"), ylim, add = FALSE,
qlvls = c(0.05, 0.5, 0.95), pch = 4, lty = 1, col, ..., cov_index,
qtype = c("points", "lines"))
```

х	an object of class PF_clouds.
У	unused.
type	parameter to specify which cloud to plot.
ylim	ylim passed to matplot.
add	TRUE if a new plot should not be made.
qlvls	vector of quantile levels to be plotted.
pch	pch argument for the quantile points.
lty	lty argument for the mean curves.
col	col argument to matplot and matpoints or matlines.
	unused.
cov_index	indices of the state vector to plot. All are plotted if this argument is omitted.
qtype	character specifying how to show quantiles. Either "points" for crosses or "lines" for dashed lines.

# plot.PF\_EM

# Value

List with quantile levels and mean curve.

plot.PF\_EM

# Plot for a PF\_EM Object

# Description

Short hand to call plot.PF\_clouds.

# Usage

## S3 method for class 'PF\_EM'
plot(x, y, ...)

# Arguments

х	an object of class PF_EM.
У	unused.
	arguments to plot.PF_clouds.

#### Value

See plot.PF\_clouds

predict.ddhazard Predict Method for ddhazard Object

# Description

Predict method for ddhazard.

# Usage

```
## S3 method for class 'ddhazard'
predict(object, new_data, type = c("response",
    "term"), tstart = "start", tstop = "stop", use_parallel, sds = F,
    max_threads, ...)
```

#### Arguments

object	result of a ddhazard call.
new_data	new data to base predictions on.
type	either "response" for predicted probability of an event or "term" for predicted terms in the linear predictor.
tstart	name of the start time column in new_data. It must be on the same time scale as the tstart used in the Surv(tstart,tstop,event) in the formula passed to ddhazard.
tstop	same as tstart for the stop argument.
use_parallel	not longer supported.
sds	TRUE if point wise standard deviation should be computed. Convenient if you use functions like ns and you only want one term per term in the right hand site of the formula used in ddhazard.
max_threads	not longer supported.
	not used.

#### Details

The function check if there are columns in new\_data which names match tstart and tstop. If matched, then the bins are found which the start time to the stop time are in. If tstart and tstop are not matched then all the bins used in the estimation method will be used.

#### Term

The result with type = "term" is a lists of list each having length equal to nrow(new\_data). The lists are

- terms It's elements are matrices where the first dimension is time and the second dimension is the terms.
- sds similar to terms for the point-wise confidence intervals using the smoothed co-variance matrices. Only added if sds = TRUE.

fixed\_terms contains the fixed (non-time-varying) effect.

- varcov similar to sds but differs by containing the whole covariance matrix for the terms. It is a 3D array where the third dimension is time. Only added if sds = TRUE.
- start numeric vector with start time for each time-varying term.

tstop numeric vector with stop time for each time-varying term.

#### Response

The result with type = "response" is a list with the elements below. If tstart and tstop are matched in columns in new\_data, then the probability will be for having an event between tstart and tstop conditional on no events before tstart.

fits fitted probability of an event.

istart numeric vector with start time for each element in fits.

istop numeric vector with stop time for each element in fits.

# print.ddhazard\_boot

#### Examples

```
fit <- ddhazard(
  Surv(time, status == 2) ~ log(bili), pbc, id = pbc$id, max_T = 3600,
  Q_0 = diag(1, 2), Q = diag(1e-4, 2), by = 50,
  control = ddhazard_control(method = "GMA"))
  predict(fit, type = "response", new_data =
    data.frame(time = 0, status = 2, bili = 3))
  predict(fit, type = "term", new_data =
    data.frame(time = 0, status = 2, bili = 3))
# probability of an event between time 0 and 2000 with bili = 3
  predict(fit, type = "response", new_data =
        data.frame(time = 0, status = 2, bili = 3, tstart = 0, tstop = 2000),
        tstart = "tstart", tstop = "tstop")
```

print.ddhazard\_boot Summary Statistics for a ddhazard\_boot Object

# Description

Arguments have the same effects as for an object from a boot call. See print.

#### Usage

```
## S3 method for class 'ddhazard_boot'
print(x, digits = getOption("digits"),
    index = 1L:ncol(boot.out$t), ...)
```

#### Arguments

х	returned object from a ddhazard_boot call.
digits	the number of digits to be printed in the summary statistics.
index	indices indicating for which elements of the bootstrap output summary statistics are required.
	not used.

#### See Also

ddhazard\_boot

```
print.summary.ddhazard
```

Summarizing Dynamic Hazard Models Fits

# Description

The sd printed for time-varying effects are point-wise standard deviations from the smoothed covariance matrices.

#### Usage

```
## S3 method for class 'summary.ddhazard'
print(x, digits = getOption("digits"), ...)
## S3 method for class 'ddhazard'
summary(object,
   var_indices = 1:ncol(object$state_vecs), max_print = 10, ...)
```

#### Arguments

Х	object returned from summary.ddhazard.
digits	number of digits to print.
	not used.
object	object returned from ddhazard.
var_indices	variable indices to print for time-varying effects.
max_print	maximum number of time points to print coefficients at.

residuals.ddhazard Residuals Method for ddhazard Object

#### Description

Residuals method for the result of a ddhazard call.

# Usage

```
## S3 method for class 'ddhazard'
residuals(object, type = c("std_space_error",
    "space_error", "pearson", "raw"), data = NULL, ...)
```

#### Arguments

object	result of ddhazard call.
type	type of residuals. Four possible values: "std_space_error", "space_error", "pearson" and "raw". See the sections below for details.
data	data.frame with data for the Pearson or raw residuals. This is only needed if the data set is not saved with the object. Must be the same data set used in the initial call to ddhazard.
	not used.

#### Pearson and raw residuals

Is the result of a call with a type argument of either "pearson" or "raw" for Pearson residuals or raw residuals. Returns a list with class "ddhazard\_residual" with the following elements.

residuals list of residuals for each bin. Each element of the list contains a 2D array where the rows corresponds to the passed data and columns are the residuals (residuals), estimated probability of death (p\_est), outcome (Y) and row number in the initial data set (row\_num). The data rows will only have a residuals in a given risk list if they are at risk in that risk set.

type the type of residual.

#### State space errors

Is the result of a call with a type argument of either "std\_space\_error" or "space\_error". The former is for standardized residuals while the latter is non-standardized. Returns a list with class. "ddhazard\_space\_errors" with the following elements:

- residuals 2D array with either standardized or non-standardized state space errors. The row are bins and the columns are the parameters in the regression.
- standardize TRUE if standardized state space errors.

Covariances 3D array with the smoothed co-variance matrix for each set of the state space errors.

#### Examples

```
library(dynamichazard)
fit <- ddhazard(
  Surv(time, status == 2) ~ log(bili), pbc, id = pbc$id, max_T = 3600,
  Q_0 = diag(1, 2), Q = diag(1e-4, 2), by = 50,
  control = ddhazard_control(method = "GMA"))
resids <- residuals(fit, type = "pearson")$residuals
head(resids[[1]])
head(resids[[2]])</pre>
```

static\_glm

#### Description

Method to fit a static model corresponding to a ddhazard fit. The method uses weights to ease the memory requirements. See get\_survival\_case\_weights\_and\_data for details on weights.

The parallelglm\_quick and parallelglm\_QR methods are similar to two methods used in bam function in the mgcv package (see the `use.chol` argument or Wood et al. 2015). parallelglm\_QR is more stable but slower. See Golub (2013) section 5.3 for a comparison of the Cholesky decomposition method and the QR method.

# Usage

```
static_glm(formula, data, by, max_T, ..., id, family = "logit",
model = F, weights, risk_obj = NULL, speedglm = F,
only_coef = FALSE, mf, method_use = c("glm", "speedglm",
"parallelglm_quick", "parallelglm_QR"),
n_threads = getOption("ddhazard_max_threads"))
```

formula	coxph like formula with Surv(tstart,tstop,event) on the left hand site of ~.
data	data.frame or environment containing the outcome and covariates.
by	interval length of the bins in which parameters are fixed.
max_T	end of the last interval interval.
	arguments passed to glm or speedglm. If only_coef = TRUE then the arguments are passed to glm.control if glm is used.
id	vector of ids for each row of the in the design matrix.
family	"logit", "cloglog", or "exponential" for a static equivalent model of ddhazard.
model	TRUE if you want to save the design matrix used in glm.
weights	weights to use if e.g. a skewed sample is used.
risk_obj	a pre-computed result from a get_risk_obj. Will be used to skip some compu- tations.
speedglm	depreciated.
only_coef	TRUE if only coefficients should be returned. This will only call the speedglm::speedglm.wfit or glm.fit which will be faster.
mf	model matrix for regression. Needed when only_coef = TRUE
method_use	<pre>method to use for estimation. glm uses glm.fit, speedglm::speedglm uses speedglm::speedglm.wfit and parallelglm_quick and parallelglm_QR uses a parallel C++ estimation method.</pre>
n_threads	number of threads to use when method_use is "parallelglm".

static\_glm

# Value

The returned list from the glm call or just coefficients depending on the value of only\_coef.

#### References

Wood, S.N., Goude, Y. & Shaw S. (2015) Generalized additive models for large datasets. Journal of the Royal Statistical Society, Series C 64(1): 139-155.

Golub, G. H., & Van Loan, C. F. (2013). Matrix computations (4th ed.). JHU Press.

# Examples

```
library(dynamichazard)
fit <- static_glm(
   Surv(time, status == 2) ~ log(bili), pbc, id = pbc$id, max_T = 3600,
   by = 50)
fit$coefficients</pre>
```

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