

# Package: smoothSurv (via r-universe)

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**Title** Survival Regression with Smoothed Error Distribution

**Depends** R (>= 3.0.0), survival

**Imports** graphics, stats

**Description** Contains, as a main contribution, a function to fit a regression model with possibly right, left or interval censored observations and with the error distribution expressed as a mixture of G-splines. Core part of the computation is done in compiled 'C++' written using the 'Scythe' Statistical Library Version 0.3.

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**License** GPL (>= 2)

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**NeedsCompilation** yes

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confint.smoothSurvReg *Confidence Intervals for Regression Parameters of 'smoothSurvReg' Model*

---

### Description

Computes confidence intervals for one or more regression related parameters (regression coefficients, scale parameter or regression coefficients in a model for scale) for a 'smoothSurvReg' model.

### Usage

```
## S3 method for class 'smoothSurvReg'
confint(object, parm, level = 0.95,
        method = c("pseudo-variance", "asymptotic"), ...)
```

### Arguments

object	Object of class smoothSurvReg.
parm	A specification of which parameters are to be given confidence intervals, either a vector of numbers or a vector of names. If missing, all parameters are considered.
level	The confidence level required.
method	Type of confidence intervals to be calculated. Option "pseudo-variance" provides confidence intervals derived from inverted minus second derivatives of the penalized log-likelihood (pseudo-variance matrix), option "asymptotic" provides confidence intervals derived from the asymptotic covariance matrix of the parameter estimates.
...	Argument included in the function parameters for the compatibility with the generic function.

**Value**

A matrix (or vector) with columns giving lower and upper confidence limits for each parameter. These will be labelled as  $(1 - \text{level})/2$  and  $1 - (1 - \text{level})/2$  in % (by default 2.5 % and 97.5 %).

**Author(s)**

Arnošt Komárek <arnost.komarek@mff.cuni.cz>

**See Also**

[smoothSurvReg](#)

---

estimTdiff	<i>Estimate expectation of survival times and their difference from the results given by survival regression function</i>
------------	---

---

**Description**

Estimate expectation of survival times and their difference from the results given by survival regression function

**Usage**

```
estimTdiff(x, ...)
## S3 method for class 'smoothSurvReg'
estimTdiff(x, cov1, cov2, logscale.cov1, logscale.cov2,
           time0 = 0, conf.level=0.95, ...)
```

**Arguments**

x	Object of an appropriate class.
cov1	Vector or matrix with covariates values for which the expectations of the first survival time are to be computed. It must be a matrix with as many columns as is the number of covariates (interactions included, intercept excluded) or the vector of length equal to the number of covariates (interactions included, intercept excluded). If matrix is supplied then is assumed that each row of this matrix gives one covariate combination for the first survival time. Intercept is not to be included in cov1. If cov1 is missing an expectation of a survivor time for the value of a covariate vector equal to zero is computed. If there is only intercept in the model, this parameter must be always missing.
cov2	Vector or matrix with covariate values for which the expectations of the second survival time are to be computed. It must be of same size as cov1.
logscale.cov1	Vector or matrix with covariate values for the expression of log-scale (if this depended on covariates) for the first survival time. It can be omitted in the case that log-scale was common for all observations.

logscale.cov2	Vector or matrix with covariate values for the expression of log-scale (if this depended on covariates) for the second survival time. It can be omitted in the case that log-scale was common for all observations.
time0	Starting time of the follow-up as used in the model. I.e. the model is assumed to be $\log(T - time0) = x'\beta + \sigma\varepsilon$
conf.level	confidence level of produced confidence intervals.
...	who knows

**Value**

A data.frame with columns named “ET1”, “sd.ET1”, “ET1.lower”, “ET1.upper”, “ET2”, “sd.ET2”, “ET2.lower”, “ET2.upper”, “diffT”, “sd.diffT”, “diffT.lower”, “diffT.upper” giving the estimates of expected values of the survival times for covariate values given in rows of cov1 and logscale.cov1, their standard errors, estimates of expected values of survival times for covariate values given in rows of cov2 and logscale.cov2, their standard errors and estimates of a difference of expected values of survival times for covariate values given in rows of cov1, logscale.cov1 and cov2, logscale.cov2, their standard errors and confidence intervals.

**Author(s)**

Arnošt Komárek <arnost.komarek@mff.cuni.cz>

**See Also**

[smoothSurvReg](#)

---

eval.Gspline

*Evaluate a G-spline in a grid of values*

---

**Description**

This function computes values of

$$f(x) = \sum_{j=1}^g c_j \varphi_{\mu_j, \sigma_j^2}(x)$$

in a grid of  $x$  values.

In above expression,  $\varphi_{\mu_j, \sigma_j^2}(x)$  denotes a density of  $N(\mu_j, \sigma_j^2)$ .

**Usage**

eval.Gspline(Gspline, grid)

**Arguments**

Gspline	A data frame with at least three columns named “Knot”, “SD basis” and “c coef.” which determine $\mu_1, \dots, \mu_g, \sigma_1, \dots, \sigma_g$ and $c_1, \dots, c_g$ . Data.frame with such properties can be found e.g. as spline component of the resulting object returned by functions <a href="#">smoothSurvReg</a> and <a href="#">minPenalty</a> .
grid	A numeric vector giving the grid of $x$ values at which the G-spline is to be evaluated.

**Value**

A data.frame with columns named “x” (grid) and “y” (G-spline values).

**Author(s)**

Arnošt Komárek <arnost.komarek@mff.cuni.cz>

**Examples**

```
spline <- minPenalty(knots=seq(-4.2, 4.2, by=0.3), sdspline=0.2, difforder=3)$spline
values <- eval.Gspline(spline, seq(-4.5, 4.5, by=0.05))
plot(values, type="l", bty="n", lwd=3)
```

---

extreme value

*Density of the Extreme Value Distribution of a Minimum.*

---

**Description**

Density function of the extreme value distribution of a minimum with location  $\alpha$  and scale  $\beta$  and the density of the standardized version (with zero mean and unit variance).

**Usage**

```
dextreme(x, alpha=0, beta=1)
dstextreme(x)
```

**Arguments**

x	Vector of quantiles.
alpha	Vector of location parameters.
beta	Vector of scale parameters.

**Details**

Extreme value distribution of a minimum with the location  $\alpha$  and the scale  $\beta$  has a density

$$f(x) = \frac{1}{\beta} \exp \left[ \frac{x - \alpha}{\beta} - \exp \left( \frac{x - \alpha}{\beta} \right) \right]$$

the mean equal to  $\alpha - \beta e$ , where  $e$  is approximately 0.5772 and the variance equal to  $\beta^2 \frac{\pi}{6}$ . Its standardized version is obtained with  $\alpha = \frac{\sqrt{6}}{\pi} e$  and  $\beta = \frac{\sqrt{6}}{\pi}$

**Value**

The value of the density.

**Author(s)**

Arnošt Komárek <arnost.komarek@mff.cuni.cz>

**Examples**

```
dextreme(1, (sqrt(6)/pi)*0.5772, sqrt(6)/pi)
dstextreme(1)      ## approximately same result as on the previous row
```

---

fdensity.smoothSurvReg

*Density for Objects of Class 'smoothSurvReg'*

---

**Description**

Compute and plot density function for given combinations of covariates based on the fitted model.

**Usage**

```
## S3 method for class 'smoothSurvReg'
fdensity(x, cov, logscale.cov, time0 = 0, plot = TRUE,
         by, xlim, ylim, xlab = "t", ylab = "f(t)",
         type = "l", lty, main, sub, legend, bty = "n", cex.legend = 1, ...)
```

**Arguments**

x	Object of class smoothSurvReg.
cov	Vector or matrix with covariates values for which the survivor curve/cdf is to be computed and plotted. It must be a matrix with as many columns as is the number of covariates (interactions included) or the vector of length equal to the number of covariates (interactions included). Intercept is not to be included in cov. If cov is missing a survivor curve for the value of a covariate vector equal to zero is plotted. If there is only intercept in the model the survivor curve based on the fitted error distribution is always plotted.
logscale.cov	Vector or matrix with covariate values for the expression of log-scale (if this depended on covariates). It can be omitted in the case that log-scale was common for all observations.
time0	Starting time of the follow-up as used in the model. I.e. the model is assumed to be $\log(T - time0) = x'\beta + \sigma\varepsilon$
plot	If TRUE the plot is directly produced, otherwise only a data.frame with information used for later plotting is returned.
by	Step for a plotting grid. If missing it is automatically computed.

xlim, ylim	Arguments passed to the <a href="#">plot</a> function.
xlab, ylab	Arguments passed to the <a href="#">plot</a> function.
type, lty	Arguments passed to the <a href="#">plot</a> function.
main, sub	Arguments passed to the <a href="#">plot</a> function.
legend, bty	Argument passed to the <a href="#">plot</a> function.
cex.legend	argument passed to cex argument of the <a href="#">legend</a> function.
...	Arguments passed to the <a href="#">plot</a> function.

**Value**

A dataframe with columns named x and y where x gives the grid and y the values of the density function at that grid.

**Author(s)**

Arnošt Komárek <arnost.komarek@mff.cuni.cz>

**See Also**

[smoothSurvReg](#), [plot](#)

---

hazard.smoothSurvReg *Hazard Curves for Objects of Class 'smoothSurvReg'*

---

**Description**

Compute and plot hazard function for given combinations of covariates based on the fitted model.

**Usage**

```
## S3 method for class 'smoothSurvReg'
hazard(x, cov, logscale.cov, time0 = 0, plot = TRUE,
       by, xlim, ylim, xlab = "t", ylab = "h(t)",
       type = "l", lty, main, sub, legend, bty = "n", cex.legend = 1, ...)
```

**Arguments**

x	Object of class smoothSurvReg.
cov	Vector or matrix with covariates values for which the survivor curve/cdf is to be computed and plotted. It must be a matrix with as many columns as is the number of covariates (interactions included) or the vector of length equal to the number of covariates (interactions included). Intercept is not to be included in cov. If cov is missing a survivor curve for the value of a covariate vector equal to zero is plotted. If there is only intercept in the model the survivor curve based on the fitted error distribution is always plotted.

logscale.cov	Vector or matrix with covariate values for the expression of log-scale (if this depended on covariates). It can be omitted in the case that log-scale was common for all observations.
time0	Starting time of the follow-up as used in the model. I.e. the model is assumed to be $\log(T - \text{time0}) = x'\beta + \sigma\varepsilon$
plot	If TRUE the plot is directly produced, otherwise only a data.frame with information used for later plotting is returned.
by	Step for a plotting grid. If missing it is automatically computed.
xlim, ylim	Arguments passed to the <code>plot</code> function.
xlab, ylab	Arguments passed to the <code>plot</code> function.
type, lty	Arguments passed to the <code>plot</code> function.
main, sub	Arguments passed to the <code>plot</code> function.
legend, bty	Argument passed to the <code>plot</code> function.
cex.legend	argument passed to cex argument of the <code>legend</code> function.
...	Arguments passed to the <code>plot</code> function.

**Value**

A dataframe with columns named x and y where x gives the grid and y the values of the hazard function at that grid.

**Author(s)**

Arnošt Komárek <arnost.komarek@mff.cuni.cz>

**See Also**

[smoothSurvReg](#), [plot](#)

---

minPenalty

*Minimize the penalty term under the two (mean and variance) constraints*

---

**Description**

This function minimizes

$$\frac{1}{2} \sum_{j=m+1}^g \left( \Delta^m a_j \right)^2$$

with respect to  $a_1, \dots, a_g$  under the constraints

$$\sum_{j=1}^g c_j \mu_j = 0$$



and

$$\sum_{j=1}^g c_j (\mu_j^2 + \sigma_0^2) = 1,$$

where

$$c_j = \frac{\exp(a_j)}{\sum_{l=1}^g \exp(a_l)}$$

with one of  $a$ 's fixed to zero.

Note that the minimum is always zero. We are thus mainly interested in the point where the minimum is reached.

### Usage

```
minPenalty(knots = NULL, dist.range = c(-6, 6), by.knots = 0.3, sdspline = NULL,
  difforder = 3, init.c,
  maxiter = 200, rel.tolerance = 1e-10, toler.chol = 1e-15, toler.eigen = 1e-3,
  maxhalf = 10, debug = 0, info = TRUE)
```

### Arguments

knots	A vector of knots $\mu_1, \dots, \mu_g$ .
dist.range	Approximate minimal and maximal knot. If not given by knots the knots are determined as $c(\text{seq}(0, \text{dist.range}[2], \text{by} = \text{by.knots}), \text{seq}(0, \text{dist.range}[1], \text{by} = -\text{by.knots}))$ . The sequence of knots is sorted and multiple entries are removed.
by.knots	The distance between the two knots used when building a vector of knots if these are not given by knots.
sdspline	Standard deviation $\sigma_0^2$ of the basis G-spline (here it appears only in the variance constraint). If not given it is determined as $2/3$ times the maximal distance between the two knots. If $\text{sdspline} \geq 1$ it is changed to 0.9 to be able to satisfy the constraints.
difforder	The order of the finite difference used in the penalty term.
init.c	Optional vector of the initial values for the G-spline coefficients $c$ , all values must lie between 0 and 1 and must sum up to 1.
maxiter	Maximum number of Newton-Raphson iterations.
rel.tolerance	(Relative) tolerance to declare the convergence. For this function, the convergence is declared if absolute value of the penalty is lower than $\text{rel.tolerance}$ and if both constraints are satisfied up to $\text{rel.tolerance}$ .
toler.chol	Tolerance to declare Cholesky decomposition singular.
toler.eigen	Tolerance to declare an eigen value of a matrix to be zero.
maxhalf	Maximum number of step-halving steps if updated estimate leads to a decrease of the objective function.
debug	If non-zero print debugging information.
info	If TRUE information concerning the iteration process is printed during the computation to the standard output.

**Value**

A list with the components “spline”, “penalty”, “warning”, “fail”.

spline	A data frame with columns named “Knot”, “SD basis”, “c coef.” and “a coef.” which gives the optimal values of $c_1, \dots, c_g$ and $a_1, \dots, a_g$ in the latter two columns. This data.frame can be further worked out using the function <code>eval.Gspline</code> .
penalty	The value of the penalty term when declaring convergence.
warning	Possible warnings concerning the convergence.
fail	Failure indicator. It is zero if everything went OK.

**Author(s)**

Arnošt Komárek <arnost.komarek@mff.cuni.cz>

**Examples**

```
optimum <- minPenalty(knots=seq(-4.2, 4.2, by = 0.3), sdspline=0.2, difforder=3)
where <- optimum$spline
print(where)
show <- eval.Gspline(where, seq(-4.2, 4.2, by=0.05))
plot(show, type="l", bty="n", lwd=2)
```

---

piece

*Left Continuous Piecewise Constant Function with a Finite Support.*

---

**Description**

Function to evaluate a left continuous piecewise constant function with a finite support.

**Usage**

```
piece(x, breaks, values)
```

**Arguments**

x	Vector of values where the piecewise constant function should be evaluated.
breaks	Vector of sorted breakpoints of the piecewise constant function.
values	Values of the piecewise constant function. It takes the value <code>value[i]</code> on the interval <code>(breaks[i], breaks[i+1]]</code> . The function is assumed to be zero outside its range specified as <code>(breaks[1], breaks[length(breaks)])</code> . The length of the vector values must be equal to <code>length(breaks) - 1</code>

**Value**

The value of the piecewise constant function.

**Author(s)**

Arnošt Komárek <arnost.komarek@mff.cuni.cz>

**Examples**

```
my.breaks <- c(-2, 1.5, 4, 7)
my.values <- c(0.5, 0.9, -2)
grid <- seq(-3, 8, by = 0.25)
piece(grid, my.breaks, my.values)
```

---

plot.smoothSurvReg      *Plot Objects of Class 'smoothSurvReg'*

---

**Description**

Plot the fitted error distribution.

**Usage**

```
## S3 method for class 'smoothSurvReg'
plot(x, plot = TRUE, resid = TRUE, knots = TRUE,
     compare = TRUE, components = FALSE, standard = TRUE,
     by, toler.c = 1e-5,
     xlim, ylim,
     xlab = expression(epsilon), ylab = expression(paste("f(", epsilon, ")", sep = "")),
     type = "l", lty = 1, main, sub, bty = "n", ...)
```

**Arguments**

x	Object of class smoothSurvReg.
plot	If TRUE the plot is directly produced, otherwise only a data.frame with information used for later plotting is returned.
resid	If resid & plot residuals are added to the plot in the following way. Residuals based on exact observations are plotted by pch = 3 (+), residuals based on right censored observations using pch = 4 (x), residuals based on left censored observations using pch = 2 (triangle) and midpoints of residuals based on interval censored observations using pch = 5 (diamond).
knots	If knots & plot bullets indicating (transformed) knots corresponding to 'c' G-spline coefficients higher than toler.c are added to the x-axis of the plot.
compare	If compare & !components & standard & plot plots of the three parametric distributions (st. normal, st. logistic and st. minimum extreme value) are added to the plot for comparison.
components	If components & standard & plot dashed lines with weighted G-spline components are added to the plot.

standard	If standard data for plotting the fitted error distribution are created, otherwise data for plotting the distribution of $\alpha + \sigma\varepsilon$ are created. In the case that $\log(\sigma)$ depends on covariates and standard is FALSE, all covariates equal to zero are used to compute $\log(\sigma)$ .
by	Step for a plotting grid. If NULL it is automatically computed.
toler.c	Tolerance to indicate zero 'c' G-spline coefficients used if knots == TRUE.
xlim, ylim	Arguments passed to the <code>plot</code> function.
xlab, ylab	Arguments passed to the <code>plot</code> function.
type, lty	Arguments passed to the <code>plot</code> function.
main, sub	Arguments passed to the <code>plot</code> function.
bty	Argument passed to the <code>plot</code> function.
...	Arguments passed to the <code>plot</code> function.

**Value**

A dataframe with columns named x and y where x gives the grid and y the values of the density at that grid.

**Author(s)**

Arnošt Komárek <arnost.komarek@mff.cuni.cz>

**See Also**

[smoothSurvReg](#), [plot](#)

---

print.estimTdiff      *Print for Objects of Class 'estimTdiff'*

---

**Description**

Print a summary information of the estimates and tests for expected values of survival times based on a regression.

**Usage**

```
## S3 method for class 'estimTdiff'
print(x, digits = min(options())$digits, 4), ...)
```

**Arguments**

x	Object of class estimTdiff.
digits	Controls the number of digits to print when printing numeric values. It is a suggestion only. Valid values are 1...22.
...	Further arguments passed to or from other methods.

**Value**

No return value, called to print the object.

**Author(s)**

Arnošt Komárek <arnost.komarek@mff.cuni.cz>

**See Also**

[smoothSurvReg](#), [print](#)

---

print.smoothSurvReg    *Summary and Print for Objects of Class 'smoothSurvReg'*

---

**Description**

Print a summary information of the fitted model.

For **regression coefficients** the following information is given:

'Value'	- estimate of the coefficient
'Std.Error'	- estimated standard error based on the pseudo-variance estimate (3.1) in Komárek, Lesaffre and Hilton (2005)
'Std.Error2'	- estimated standard error based on the asymptotic variance estimate (3.2) in Komárek, Lesaffre and Hilton (2005)
'Z'	- the Wald statistic obtained as 'Value' divided by 'Std.Error'
'Z2'	- the Wald statistic obtained as 'Value' divided by 'Std.Error2'
'p'	- the two-sided P-value based on normality of the statistic 'Z'
'p2'	- the two-sided P-value based on normality of the statistic 'Z2'

Further, we print:

'Lambda'	- the optimal value of the smoothing hyperparameter divided by the sample size, i.e., $\lambda/n$ in the notation of Komárek, Lesaffre and Hilton (2005)
'Log(Lambda)'	- logarithm of the above
'df'	- effective degrees of freedom of the model, see Section 2.2.3 of Komárek, Lesaffre and Hilton (2005)
'AIC'	- Akaike's information criterion of the model, see Section 2.2.3 of Komárek, Lesaffre and Hilton (2005)

With argument **spline** set to TRUE, analogous table like that for the regression coefficients is printed also for the weights of the penalized Gaussian mixture (G-spline).

**Usage**

```
## S3 method for class 'smoothSurvReg'  
print(x, spline, digits = min(options())$digits, 4), ...)  
## S3 method for class 'smoothSurvReg'  
summary(object, spline, digits = min(options())$digits, 4), ...)
```

**Arguments**

x	Object of class smoothSurvReg.
object	Object of class smoothSurvReg.
spline	TRUE/FALSE. If TRUE an information on fitted G-spline is printed.
digits	Controls the number of digits to print when printing numeric values. It is a suggestion only. Valid values are 1...22.
...	Further arguments passed to or from other methods.

**Value**

No return value, called to print the object.

**Author(s)**

Arnošt Komárek <arnost.komarek@mff.cuni.cz>

**References**

Komárek, A., Lesaffre, E., and Hilton, J. F. (2005). Accelerated failure time model for arbitrarily censored data with smoothed error distribution. *Journal of Computational and Graphical Statistics*, **14**, 726–745.

Lesaffre, E., Komárek, A., and Declerck, D. (2005). An overview of methods for interval-censored data with an emphasis on applications in dentistry. *Statistical Methods in Medical Research*, **14**, 539–552.

**See Also**

[smoothSurvReg](#), [print](#), [summary](#)

---

residuals.smoothSurvReg

*Residuals for Objects of Class 'smoothSurvReg'*

---

**Description**

Compute residuals for the fitted model.

**Usage**

```
## S3 method for class 'smoothSurvReg'  
residuals(object, ...)
```

**Arguments**

object            Object of class smoothSurvReg.  
...                Argument included in the function parameters for the compatibility with the generic function.

**Value**

A dataframe with columns named res, res2 and censor where the column res2 is included only if there are any interval censored observations. Column res contains all residuals, column res2 is applicable only for interval censored observations. Column censor gives the type of censoring (0 for right censoring, 1 for exact observations, 2 for left censoring and 3 for interval censoring).

**Author(s)**

Arnošt Komárek <arnost.komarek@mff.cuni.cz>

**See Also**

[smoothSurvReg](#)

---

smoothSurvReg

*Regression for a Survival Model with Smoothed Error Distribution*

---

**Description**

Regression for a survival model. These are all time-transformed location models, with the most useful case being the accelerated failure models that use a log transformation. Error distribution is assumed to be a mixture of G-splines. Parameters are estimated by the penalized maximum likelihood method.

**Usage**

```
smoothSurvReg(formula = formula(data), logscale = ~1,  
  data = parent.frame(), subset, na.action = na.fail,  
  init.beta, init.logscale, init.c, init.dist = "best",  
  update.init = TRUE, aic = TRUE, lambda = exp(2:(-9)),  
  model = FALSE, control = smoothSurvReg.control(), ...)
```

**Arguments**

formula	A formula expression as for other regression models. See the documentation for <a href="#">lm</a> and <a href="#">formula</a> for details. Use <a href="#">Surv</a> on the left hand side of the formula.
logscale	A formula expression to determine a possible dependence of the log-scale on covariates.
data	Optional data frame in which to interpret the variables occurring in the formula.
subset	Subset of the observations to be used in the fit.
na.action	Function to be used to handle any NAs in the data. It's default value is <code>na.fail</code> . It is not recommended to change it in the case when <code>logscale</code> depends on covariates.
init.beta	Optional vector of the initial values of the regression parameter $\beta$ (intercept and regression itself).
init.logscale	Optional value of the initial value of the parameters that determines the log-scale parameter $\log(\sigma)$ .
init.c	Optional vector of the initial values for the G-spline coefficients <code>c</code> , all values must lie between 0 and 1 and must sum up to 1.
init.dist	A character string specifying the distribution used by <a href="#">survreg</a> to find the initial values for parameters (if not given by the user). It is assumed to name "best" or an element from <a href="#">survreg.distributions</a> . These include "weibull", "exponential", "gaussian", "logistic", "lognormal" and "loglogistic". If "best" is specified one of "lognormal", "weibull" and "loglogistic" giving the highest likelihood is used.
update.init	If TRUE, the initial values are updated during the grid search for the lambda parameter giving the optimal AIC. Otherwise, fits with all lambdas during the grid search start with same initials determine at the beginning either from the values of <code>init.beta</code> , <code>init.scale</code> , <code>init.c</code> or from the initial <a href="#">survreg</a> fit as determined by the parameter <code>init.dist</code> .
aic	If TRUE the optimal value of the tuning parameter $\lambda$ is determined via a grid search through the values specified by the parameter <code>lambda</code> . If FALSE, only the model with $\lambda = \text{lambda}[1]$ is fitted.
lambda	A grid of values of the tuning parameter $\lambda$ searched for the optimal value if <code>aic = TRUE</code> .
model	If TRUE, the model frame is returned.
control	A list of control values, in the format produced by <a href="#">smoothSurvReg.control</a> .
...	Other arguments which will be passed to <a href="#">smoothSurvReg.control</a> . See its help page for more options to control the fit and for the possibility to fix some values and not to estimate them.

**Details**

Read the papers referred below.



There is a slight difference in the definition of the penalty used by the R function compared to what is written in the paper. The penalized log-likelihood given in the paper has a form

$$\ell_P(\theta) = \ell(\theta) - \frac{\lambda}{2} \sum_{j=m+1}^g (\Delta^m a_j)^2,$$

while the penalized log-likelihood used in the R function multiplies the tuning parameter  $\lambda$  given by lambda by a sample size  $n$  to keep default values more or less useful for samples of different sizes. So that the penalized log-likelihood which is maximized by the R function has the form

$$\ell_P(\theta) = \ell(\theta) - \frac{\lambda \cdot n}{2} \sum_{j=m+1}^g (\Delta^m a_j)^2.$$

### Value

An object of class smoothSurvReg is returned. See [smoothSurvReg.object](#) for details.

### Author(s)

Arnošt Komárek <arnost.komarek@mff.cuni.cz>

### References

Komárek, A., Lesaffre, E., and Hilton, J. F. (2005). Accelerated failure time model for arbitrarily censored data with smoothed error distribution. *Journal of Computational and Graphical Statistics*, **14**, 726–745.

Lesaffre, E., Komárek, A., and Declerck, D. (2005). An overview of methods for interval-censored data with an emphasis on applications in dentistry. *Statistical Methods in Medical Research*, **14**, 539–552.

### Examples

```
##### EXAMPLE 1: Common scale
##### =====
### We generate interval censored data and fit a model with few artificial covariates
set.seed(221913282)
x1 <- rbinom(50, 1, 0.4)                ## binary covariate
x2 <- rnorm(50, 180, 10)                ## continuous covariate
y1 <- 0.5*x1 - 0.01*x2 + 0.005 *x1*x2 + 1.5*rnorm(50, 0, 1)  ## generate log(T), left limit
t1 <- exp(y1)                           ## left limit of the survival time
t2 <- t1 + rgamma(50, 1, 1)              ## right limit of the survival time
surv <- Surv(t1, t2, type = "interval2") ## survival object

## Fit the model with an interaction
fit1 <- smoothSurvReg(surv ~ x1 * x2, logscale = ~1, info = FALSE, lambda = exp(2:(-1)))

## Print the summary information
summary(fit1, spline = TRUE)

## Plot the fitted error distribution
```

```

plot(fit1)

## Plot the fitted error distribution with its components
plot(fit1, components = TRUE)

## Plot the cumulative distribution function corresponding to the error density
survfit(fit1, cdf = TRUE)

## Plot survivor curves for persons with (x1, x2) = (0, 180) and (1, 180)
cov <- matrix(c(0, 180, 0, 1, 180, 180), ncol = 3, byrow = TRUE)
survfit(fit1, cov = cov)

## Plot hazard curves for persons with (x1, x2) = (0, 180) and (1, 180)
cov <- matrix(c(0, 180, 0, 1, 180, 180), ncol = 3, byrow = TRUE)
hazard(fit1, cov = cov)

## Plot densities for persons with (x1, x2) = (0, 180) and (1, 180)
cov <- matrix(c(0, 180, 0, 1, 180, 180), ncol = 3, byrow = TRUE)
fdensity(fit1, cov = cov)

## Compute estimates expectations of survival times for persons with
## (x1, x2) = (0, 180), (1, 180), (0, 190), (1, 190), (0, 200), (1, 200)
## and estimates of a difference of these expectations:
## T(0, 180) - T(1, 180), T(0, 190) - T(1, 190), T(0, 200) - T(1, 200),
cov1 <- matrix(c(0, 180, 0, 0, 190, 0, 0, 200, 0), ncol = 3, byrow = TRUE)
cov2 <- matrix(c(1, 180, 180, 1, 190, 190, 1, 200, 200), ncol = 3, byrow = TRUE)
print(estimTdiff(fit1, cov1 = cov1, cov2 = cov2))

##### EXAMPLE 2: Scale depends on covariates
##### =====
### We generate interval censored data and fit a model with few artificial covariates
set.seed(221913282)
x1 <- rbinom(50, 1, 0.4) ## binary covariate
x2 <- rnorm(50, 180, 10) ## continuous covariate
x3 <- runif(50, 0, 1) ## covariate for the scale parameter
logscale <- 1 + x3
scale <- exp(logscale)
y1 <- 0.5*x1 - 0.01*x2 + 0.005 *x1*x2 + scale*rnorm(50, 0, 1) ## generate log(T), left limit
t1 <- exp(y1) ## left limit of the survival time
t2 <- t1 + rgamma(50, 1, 1) ## right limit of the survival time
surv <- Surv(t1, t2, type = "interval2") ## survival object

## Fit the model with an interaction
fit2 <- smoothSurvReg(surv ~ x1 * x2, logscale = ~x3, info = FALSE, lambda = exp(2:(-1)))

## Print the summary information
summary(fit2, spline = TRUE)

## Plot the fitted error distribution
plot(fit2)

## Plot the fitted error distribution with its components

```

```

plot(fit2, components = TRUE)

## Plot survivor curves for persons with (x1, x2) = (0, 180) and (1, 180)
## x3 = 0.8 and 0.9
cov <- matrix(c(0, 180, 0, 1, 180, 180), ncol = 3, byrow = TRUE)
logscale.cov <- c(0.8, 0.9)
survfit(fit2, cov = cov, logscale.cov = logscale.cov)

## Plot hazard curves for persons with (x1, x2) = (0, 180) and (1, 180)
## x3 = 0.8 and 0.9
cov <- matrix(c(0, 180, 0, 1, 180, 180), ncol = 3, byrow = TRUE)
logscale.cov <- c(0.8, 0.9)
hazard(fit2, cov = cov, logscale.cov=c(0.8, 0.9))

## Plot densities for persons with (x1, x2) = (0, 180) and (1, 180)
## x3 = 0.8 and 0.9
cov <- matrix(c(0, 180, 0, 1, 180, 180), ncol = 3, byrow = TRUE)
logscale.cov <- c(0.8, 0.9)
fdensity(fit2, cov = cov, logscale.cov = logscale.cov)

## More involved examples can be found in script files
## used to perform analyses and draw pictures
## presented in above mentioned references.
## These scripts and some additional files can be found as *.tar.gz files
## in the /inst/doc directory of this package.
##

```

---

smoothSurvReg.control *More Options for 'smoothSurvReg'*

---

## Description

This function checks and sets the fitting options for smoothSurvReg. Its arguments can be used instead of ... in a call to smoothSurvReg.

## Usage

```

smoothSurvReg.control(est.c = TRUE, est.scale = TRUE,
  maxiter = 200, firstiter = 0, rel.tolerance = 5e-5,
  toler.chol = 1e-15, toler.eigen = 1e-3,
  maxhalf = 10, debug = 0, info = TRUE, lambda.use = 1.0, sdspline = NULL,
  difforder = 3, dist.range = c(-6, 6), by.knots = 0.3,
  knots = NULL, nsplines = NULL, last.three = NULL)

```

## Arguments

**est.c** If TRUE the G-spline coefficients are estimated. Otherwise, they are fixed to the values given by `init.c` parameter of [smoothSurvReg](#).

<code>est.scale</code>	If TRUE the scale parameter $\sigma$ is estimated. Otherwise, it is fixed to the value given by <code>init.scale</code> parameter of <code>smoothSurvReg</code> .
<code>maxiter</code>	Maximum number of Newton-Raphson iterations.
<code>firstiter</code>	The index of the first iteration. This option comes from older versions of this function.
<code>rel.tolerance</code>	(Relative) tolerance to declare the convergence. In this version of the function, the convergence is declared if the relative difference between two consecutive values of the penalized log-likelihood are smaller than <code>rel.tolerance</code> .
<code>toler.chol</code>	Tolerance to declare Cholesky decomposition singular.
<code>toler.eigen</code>	Tolerance to declare an eigen value of a matrix to be zero.
<code>maxhalf</code>	Maximum number of step-halving steps if updated estimate leads to a decrease of the objective function.
<code>debug</code>	If non-zero print debugging information.
<code>info</code>	If TRUE information concerning the iteration process is printed during the computation to the standard output.
<code>lambda.use</code>	The value of the tuning (penalty) parameter $\lambda$ used in a current fit by the <code>smoothSurvReg.fit</code> function. Value of this option is not interesting for the user. The parameter <code>lambda</code> of the function <code>smoothSurvReg</code> is more important for the user.
<code>sdspline</code>	Standard deviation of the basis G-spline. If not given it is determined as 2/3 times the maximal distance between the two knots. If <code>est.c = TRUE</code> and <code>sdspline &gt;= 1</code> it is changed to 0.9 to be able to satisfy the constraints imposed to the fitted error distribution.
<code>difforder</code>	The order of the finite difference used in the penalty term.
<code>dist.range</code>	Approximate minimal and maximal knot. If not given by <code>knots</code> the knots are determined as <code>c(seq(0, dist.range[2], by = by.knots), seq(0, dist.range[1], by = -by.knots))</code> . The sequence of knots is sorted and multiple entries are removed.
<code>by.knots</code>	The distance between the two knots used when building a vector of knots if these are not given by <code>knots</code> . This option is ignored if <code>nsplines</code> is not NULL.
<code>knots</code>	A vector of knots.
<code>nsplines</code>	This option is ignored at this moment. It is used to give the number of G-splines to the function <code>smoothSurvReg.fit</code> .
<code>last.three</code>	A vector of length 3 with indeces of reference knots. The 'a' coefficient of the knot[ <code>last.three[1]</code> ] is then equal to zero, 'a' coefficients with indeces <code>last.three[2:3]</code> are expressed as a function of remaining 'a' coefficients such that resulting error distribution has zero mean and unit variance. If <code>maxiter &gt; 0</code> <code>last.three</code> is determined after the convergence is reached. If <code>maxiter == 0</code> <code>last.three</code> is used to compute variance matrices.

**Value**

A list with the same elements as the input except `dist.range` and `by.knots` is returned.

**Author(s)**

Arnošt Komárek <arnost.komarek@mff.cuni.cz>

---

smoothSurvReg.fit      *Work Function to Fit the Model Using 'smoothSurvReg'*

---

## Description

Fit the survival regression model with smoothed error distribution. This function is not to be called by the user.

## Usage

```
smoothSurvReg.fit(x, z, y, offset = NULL, correctlik, init, controlvals, common.logscale)
```

## Arguments

x	A covariate matrix.
z	A covariate matrix for log(scale).
y	A two- or three- column matrix with response. The last column indicate the type of censoring. See <a href="#">Surv</a> for details.
offset	A vector with possible offset term.
correctlik	Correction term to the likelihood due to the log transformation of the response.
init	A list with components beta, scale, ccoef giving the initial values for the fitting process.
controlvals	A list returned by <a href="#">smoothSurvReg.control</a> .
common.logscale	Indicator (TRUE/FALSE) indicating whether the log-scale is same for all observations or whether it depends on covariates.

## Value

A list with components regres, spline, loglik, aic, degree.smooth, var, var2, dCdC, iter, estimated, warning, fail, H, I, G, U.

## Note

WARNING: Most users will call the higher level routine smoothSurvReg. Consequently, this function has very few error checks on its input arguments.

## Author(s)

Arnošt Komárek <arnost.komarek@mff.cuni.cz>

---

smoothSurvReg.object    *Smoothed Survival Regression Object*

---

### Description

This class of objects is returned by the smoothSurvReg class of functions to represent a fitted smoothed survival regression model.

Objects of this class have methods for the functions print, summary, plot, residuals, survfit.

### COMPONENTS I

The following components must be included in a legitimate smoothSurvReg object.

**fail** Indicator of the failure of the fitting procedure. Possible values are 0 for no problems, 3 if the iteration process was stopped because of non-positive definite minus Hessian, 4 if the iteration process was stopped because too many halving steps were performed, 5 if it was not possible to find the three reference knots (it was not then possible to perform optimization with respect to the full parameter vector), 6 if the maximal number of iterations was performed without reaching a convergence. The fail component is increased by 10 if the final minus Hessian of the penalized log-likelihood was not positive definite. The fail component is further increased by 20 if the computed effective degrees of freedom were non-positive. The fail component is further increased by 40 if there are negative estimates of standard errors for some regression parameters. The fail component is 99 or higher if the fitting procedure failed at all and there is no fit produced.

### COMPONENTS II

The following components must be included in a legitimate smoothSurvReg object if fail is lower than 99.

**regres** Estimates of the regression parameters  $\alpha, \beta, \sigma$  if these have been estimated with their standard errors stored in a data frame with colnames “Value”, “Std.Error”, “Std.Error2” and rownames derived from the names of the design matrix with “(Intercept)” for the intercept, “Scale” for the scale and “Log(scale)” for the log-scale. If the log-scale depends on covariates then rows named “LScale.(Intercept)”, “LScale.cov1” etc. give estimates of regression parameters for log-scale. The two standard errors are computed using either var or var2 described below.

**spline** Description of the fitted error density. A data frame with colnames “Knot”, “SD basis”, “c coef.”, “Std.Error.c”, “Std.Error2.c”, “a coef.”, “Std.Error.a” and “Std.Error2.a” and rownames knot[1], ..., knot[g] where  $g$  stands for the number of basis G-splines. The column “Knot” contains the knots in ascending order, “SD basis” the standard deviation of an appropriate basis G-spline, “c coef.” estimates of the G-spline coefficients and “Std.Error.c” and “Std.Error2.c” the estimates of their standard errors based either on var or var2. The column “a coef.” contains the estimates of transformed  $c$  coefficients where

$$c_j = \frac{\exp(a_j)}{\sum_{i=1}^g \exp(a_i)}, j = 1, \dots, g.$$

If the error distribution is estimated, one of the  $a$  coefficients is set to zero and two other  $a$ 's are expressed as a function of the remaining  $a$  coefficients (to avoid equality constraints concerning the mean and the variance of the error distribution). The standard error for these three  $a$  coefficients is then not available (it is equal to NA). Standard error is set to NaN if a diagonal element of the appropriate covariance matrix was negative.

**loglik** Maximized penalized log-likelihood, log-likelihood and the penalty term. A data frame with one row and three columns named "Log Likelihood", "Penalty" and "Penalized Log Likelihood".

**aic** Akaike's information criterion of the fitted model computed as a maximized value of the penalized log-likelihood minus the effective degrees of freedom.

**degree.smooth** Effective degrees of freedom, number of parameters and related information. A data frame with one row and columns named "Lambda", "Log(Lambda)", "df", "Number of parameters", "Mean param.", "Scale param.", "Spline param." where "Lambda" gives the value of the tuning parameter used in the final (optimal) fit, "df" the effective degrees of freedom, "Number of parameters" the real number of parameters and "Mean param.", "Scale param." and "Spline param." its decomposition. Note that if G-spline coefficients are estimated "Spline param." is equal to the number of basis G-spline with non-zero coefficients minus three.

**var** The estimate of the covariance matrix of the estimates based on the Bayesian approximation. It is equal to the inverse of the converged minus Hessian of the penalized log-likelihood. Note that there are no columns and rows corresponding to the three transformed G-spline coefficients since these are functions of the remaining transformed G-spline coefficients (to avoid equality constraints).

**var2** The estimate of the covariance matrix of the estimates based on the asymptotic theory for penalized models. It is equal to  $H^{-1} I H^{-1}$  where H is converged minus Hessian of the penalized log-likelihood and I is converged minus Hessian of the log-likelihood component of the penalized log-likelihood.

**dCdD** A matrix with derivatives of  $c$  spline coefficients with respect to  $d$  spline coefficients (these are  $a$  coefficients with three of them omitted). This matrix can be used later to compute estimates and standard errors of functions of original parameters using a Delta method. For closer definition of  $d$  coefficients see an enclosed document.

**iter** Used number of iterations to fit the model with the optimal  $\lambda$ .

**estimated** Indicator of what has really been estimated and not fixed. A four-component vector with component names "(Intercept)", "Scale", "ccoef", "common.logscale". The first component is TRUE if the intercept was included in the regression model. The second component is TRUE if the scale parameter was not fixed, the third component is TRUE if the G-spline coefficients were not fixed. The fourth component is TRUE if the log-scale does not depend on covariates.

**warning** A data frame with one column called "warnings" and three rows called "Convergence", "Final minus Hessian" and "df" containing a string information corresponding to the value of the fail component of the object. It contains a string "OK" if there are no problems with the appropriate part of the fitting process.

**H** Converged minus Hessian of the penalized log-likelihood.

**I** Converged minus Hessian of the log-likelihood component of the penalized log-likelihood.  $I = H - G$ .

**G** Converged minus Hessian of the penalty term of the penalized log-likelihood.  $G = H - I$ .

- U** Converged score vector based on the penalized log-likelihood.
- na.action** The `na.action` attribute, if any, that was returned by the `na.action` routine.
- terms** The `terms` object used.
- formula** A symbolic description of the model to be fit.
- call** The matched call.
- init.dist** A string indicating the error distribution of the untransformed response to find the initial values. Possible values are “lognormal”, “loglogistic”, “weibull”.
- model** If requested, the model frame used.
- x** The model matrix used.
- y** The response matrix used (two columns if there were no interval censored observations, three columns if there were some interval censored observations). The last column indicates the death status.
- z** The model matrix used for the expression of log-scale.
- init.spline** A data frame describing the initial error density. It has columns named “Knot”, “SD basis”, “c coef.” and rows named “knot[1]”, ..., “knot[g]”.
- init.regres** Initial estimates of the regression parameters. A data frame with one column named “Value” and rows named as in the `regres` component of the `smoothSurvReg` object.
- adjust** Adjusted intercept and scale. A data frame with a column named “Value” and rows named “(Intercept)” and “Scale”. “(Intercept)” gives the overall intercept taking into account the mean of the fitted error distribution, “Scale” gives the overall scale taking into account the variance of the fitted error distribution. If the error distribution is standardized (always when G-spline coefficients are estimated) then the “(Intercept)” is equal to the “(Intercept)” from the `regres` component and “Scale” is equal to the “Scale” of either `regres` or `init.regres` component. NA’s appear in this data frame in the case that log-scale depends on covariates.
- error.dist** A data frame with columns named “Mean”, “Var” and “SD” and a row named “Error distribution: ” giving the mean, variance and the standard deviation of the fitted error distribution. These are equal to 0, 1 and 1 if the G-spline coefficients were estimated.
- searched** Information concerning the searched values of the tuning parameter  $\lambda$  when looking for the best AIC. A data frame with columns named “Lambda”, “Log(Lambda)”, “AIC”, “df”, “PenalLogLik”, “LogLik”, “nOfParm”, “fail”.

**Author(s)**

Arnošt Komárek <arnost.komarek@mff.cuni.cz>

---

standardized logistic *Density of Standardized Logistic Distribution.*

---

**Description**

Density function of the logistic distribution with zero mean and unit variance.



**Usage**

```
dstlogis(x)
```

**Arguments**

x                    Vector of quantiles.

**Details**

```
dstlogis(x) = dlogis(x, 0, sqrt(3)/pi)
```

**Value**

The value of the density.

**Author(s)**

Arnošt Komárek <arnost.komarek@mff.cuni.cz>

**See Also**

[dlogis](#) for the logistic distribution.

**Examples**

```
dstlogis(0)
dstlogis(seq(-3, 3, 0.2))
```

---

std.data

*Standardization of the Data*

---

**Description**

Chosen columns of a `data.frame` are standardized by using a sample mean and sample standard deviation.

**Usage**

```
std.data(datain, cols)
```

**Arguments**

datain                Input data frame.  
cols                   A character vector with names of the columns to be standardized.

### Details

For each chosen column the sample mean and the sample standard deviation is computed and then used to standardize the column. Missing values are ignored when computing the mean and the standard deviation.

### Value

A `data.frame` with same variables as the input `data.frame`. Chosen columns are standardized.

### Author(s)

Arnošt Komárek <arnost.komarek@mff.cuni.cz>

### See Also

[scale](#)

### Examples

```
variable1 <- rnorm(30)
variable2 <- rbinom(30, 1, 0.4)
variable3 <- runif(30)
data.example <- data.frame(variable1, variable2, variable3)
## We standardize only the first and the third column.
data.std <- std.data(data.example, c("variable1", "variable3"))
print(data.std)
print(c(mean(data.std$variable1), sd(data.std$variable1)))
print(c(mean(data.std$variable3), sd(data.std$variable3)))
```

---

survfit.smoothSurvReg *Survivor Curves for Objects of Class 'smoothSurvReg'*

---

### Description

Compute and plot survivor function/cumulative distribution function for given combinations of co-variates based on the fitted model.

### Usage

```
## S3 method for class 'smoothSurvReg'
survfit(formula, cov, logscale.cov, time0 = 0, plot = TRUE, cdf = FALSE,
  by, xlim, ylim = c(0, 1), xlab = "t", ylab,
  type = "l", lty, main, sub, legend, bty = "n", cex.legend = 1, ...)
```

**Arguments**

<code>formula</code>	Object of class <code>smoothSurvReg</code> .
<code>cov</code>	Vector or matrix with covariates values for which the survivor curve/cdf is to be computed and plotted. It must be a matrix with as many columns as is the number of covariates (interactions included) or the vector of length equal to the number of covariates (interactions included). Intercept is not to be included in <code>cov</code> . If <code>cov</code> is missing a survivor curve for the value of a covariate vector equal to zero is plotted. If there is only intercept in the model the survivor curve based on the fitted error distribution is always plotted.
<code>logscale.cov</code>	Vector or matrix with covariate values for the expression of log-scale (if this depended on covariates). It can be omitted in the case that log-scale was common for all observations.
<code>time0</code>	Starting time of the follow-up as used in the model. I.e. the model is assumed to be $\log(T - time0) = x'\beta + \sigma\varepsilon$
<code>plot</code>	If TRUE the plot is directly produced, otherwise only a <code>data.frame</code> with information used for later plotting is returned.
<code>cdf</code>	If TRUE cumulative distribution function is plotted instead of the survivor function.
<code>by</code>	Step for a plotting grid. If NULL it is automatically computed.
<code>xlim, ylim</code>	Arguments passed to the <code>plot</code> function.
<code>xlab, ylab</code>	Arguments passed to the <code>plot</code> function.
<code>type, lty</code>	Arguments passed to the <code>plot</code> function.
<code>main, sub</code>	Arguments passed to the <code>plot</code> function.
<code>legend, bty</code>	Argument passed to the <code>plot</code> function.
<code>cex.legend</code>	argument passed to <code>cex</code> argument of the <code>legend</code> function.
<code>...</code>	Arguments passed to the <code>plot</code> function.

**Value**

A dataframe with columns named `x` and `y` where `x` gives the grid and `y` the values of the survivor/cum. distribution function at that grid.

**Author(s)**

Arnošt Komárek <arnost.komarek@mff.cuni.cz>

**See Also**

[smoothSurvReg](#), [plot](#)

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