

# Package ‘morse’

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**Type** Package

**Title** Modelling Reproduction and Survival Data in Ecotoxicology

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**URL** <https://cran.r-project.org/package=morse>

**BugReports** <https://gitlab.com/qonfluens/model/morse>

**Description** Advanced methods for a valuable quantitative environmental risk assessment using Bayesian inference of survival and reproduction Data. Among others, it facilitates Bayesian inference of the general unified threshold model of survival (GUTS). See our companion paper Baudrot and Charles (2021) <[doi:10.21105/joss.03200](https://doi.org/10.21105/joss.03200)>, as well as complementary details in Baudrot et al. (2018) <[doi:10.1021/acs.est.7b05464](https://doi.org/10.1021/acs.est.7b05464)> and Delignette-Muller et al. (2017) <[doi:10.1021/acs.est.6b05326](https://doi.org/10.1021/acs.est.6b05326)>.

**Depends** R (>= 3.5.0)

**SystemRequirements** JAGS (>= 4.0.0) (see <https://mcmc-jags.sourceforge.io>)

**Imports** coda, deSolve, dplyr, epitools, graphics, grDevices, ggplot2 (>= 2.1.0), grid, gridExtra, magrittr, methods, reshape2, rjags (>= 4.0), stats, tibble, tidyr, zoo

**License** MIT + file LICENSE

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**VignetteBuilder** knitr

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## R topics documented:

morse-package	3
cadmium1	5
cadmium2	6
chlordan	7
copper	8
dichromate	8
FOCUSprofile	9
is_exposure_constant	10
LCx	10
MFx	12
modelData	16
modelData.survDataCstExp	17
modelData.survDataVarExp	17
plot.LCx	18
plot.MFx	19
plot.reproData	21
plot.reproFitTT	22
plot.survDataCstExp	24
plot.survDataVarExp	25
plot.survFitCstExp	26
plot.survFitPredict	27
plot.survFitPredict_Nsurv	29
plot.survFitTKTD	30
plot.survFitTT	32
plot.survFitVarExp	34
plotDoseResponse	36
plotDoseResponse.reproData	36
plotDoseResponse.survDataCstExp	38
plot_prior_post	40
plot_prior_post.survFit	40
ppc	41
predict.survFit	45
predict_Nsurv_check	48
predict_ode	49

predict_ode.survFit . . . . .	50
print.msgTable . . . . .	51
print.reproFitTT . . . . .	52
print.survFitCstExp . . . . .	53
print.survFitTKTD . . . . .	54
print.survFitTT . . . . .	55
print.survFitVarExp . . . . .	56
priors_distribution . . . . .	57
priors_distribution.survFit . . . . .	57
priors_survData . . . . .	58
propiconazole . . . . .	59
propiconazole_pulse_exposure . . . . .	59
propiconazole_split . . . . .	60
reproData . . . . .	61
reproDataCheck . . . . .	62
reproFitTT . . . . .	63
summary.reproData . . . . .	65
summary.reproFitTT . . . . .	66
summary.survDataCstExp . . . . .	67
summary.survDataVarExp . . . . .	68
summary.survFit . . . . .	69
summary.survFitTKTD . . . . .	70
summary.survFitTT . . . . .	71
survData . . . . .	72
survData_join . . . . .	74
survFit . . . . .	75
survFitTKTD . . . . .	78
survFitTT . . . . .	80
survFitTT.survDataCstExp . . . . .	80
zinc . . . . .	82
<b>Index</b>	<b>83</b>

## Description

Provides tools for the analysis of survival/reproduction toxicity test data in quantitative environmental risk assessment. It can be used to explore/visualize experimental data, and to get estimates of  $LC_x$  ( $X\%$  Lethal Concentration) or,  $EC_x$  ( $X\%$  Effective Concentration) by fitting exposure-response curves. The  $LC_x$ ,  $EC_x$  and parameters of the curve are provided along with an indication of the uncertainty of the estimation. `morse` can also be used to get an estimation of the  $NEC$  (No Effect Concentration) by fitting a Toxicokinetic Toxicodynamic (TKTD) model (GUTS: General Unified Threshold model of Survival). Within the TKTD-GUTS approach,  $LC(x, t)$ ,  $EC(x, t)$  and  $MF(x, t)$  ( $x\%$  Multiplication Factors aka Lethal Profiles) can be explored in proportion  $x$  and time  $t$ .

## Details

Estimation procedures in `morse` can be used without a deep knowledge of their underlying probabilistic model or inference methods. Rather, they were designed to behave as well as possible without requiring a user to provide values for some obscure parameters. That said, `morse` models can also be used as a first step to tailor new models for more specific situations.

The package currently handles survival and reproduction data. Functions dedicated to survival (resp. reproduction) analysis start with a `surv` (resp. `repro`) prefix. `morse` provides a similar workflow in both cases:

1. create and validate a data set
2. explore a data set
3. plot a data set
4. fit a model on a data set and output the expected estimates
5. check goodness of fit with posterior predictive check plot (ppc)

More specifically, for survival data handles with TKTD 'GUTS' model, `morse` provides:

1. plot  $LC(x, t)$  and  $MF(x, t)$ .
2. compute goodness-of-fit measures (PPC percent, NRMSE and SPPE)

Those steps are presented in more details in the "Tutorial" vignette, while a more formal description of the estimation procedures are provided in the vignette called "Models in `morse` package". Please refer to these documents for further introduction to the use of `morse`.

This reference manual is a detailed description of the functions exposed in the package.

**Getting started** The package uses the `rjags` package (Plummer, 2013), an R interface to the JAGS library for Bayesian model estimation. Note that the `rjags` package does not include a copy of the JAGS library: you need to install it separately. For instructions on downloading JAGS, see the home page at <https://mcmc-jags.sourceforge.io>. Once done, simply follow the steps described in the tutorial vignette.

```
Package: morse
Type: Package
Version: 3.2.0
Date: 2018-11-15
License: GPL (>=2)
```

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## References

- Delignette-Muller, M.L., Ruiz P. and Veber P. (2017) *Robust fit of toxicokinetic-toxicodynamic models using prior knowledge contained in the design of survival toxicity tests*.
- Delignette-Muller, M.L., Lopes, C., Veber, P. and Charles, S. (2014) *Statistical handling of reproduction data for exposure-response modelling*.
- Forfait-Dubuc, C., Charles, S., Billoir, E. and Delignette-Muller, M.L. (2012) *Survival data analyses in ecotoxicology: critical effect concentrations, methods and models. What should we use?*
- Plummer, M. (2013) *JAGS Version 4.0.0 user manual*. [https://sourceforge.net/projects/mcmc-jags/files/Manuals/4.x/jags\\_user\\_manual.pdf/download](https://sourceforge.net/projects/mcmc-jags/files/Manuals/4.x/jags_user_manual.pdf/download)
- Baudrot, V., Preux, S., Ducrot, V., Pavé, A. and Charles, S. (2018) *New insights to compare and choose TKTD models for survival based on an inter-laboratory study for Lymnaea stagnalis exposed to Cd*.
- EFSA PPR Scientific Opinion (2018) *Scientific Opinion on the state of the art of Toxicokinetic/Toxicodynamic (TKTD) effect models for regulatory risk assessment of pesticides for aquatic organisms* <https://www.efsa.europa.eu/en/efsajournal/pub/5377>.

## See Also

[rjags](#), [ggplot2](#)

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cadmium1

*Reproduction and survival data sets for Daphnia magna exposed to cadmium during 21 days*

---

## Description

Reproduction and survival data sets of chronic laboratory toxicity tests with *Daphnia magna* freshwater invertebrate exposed to five concentrations of cadmium during 21 days. Five concentrations were tested, with four replicates per concentration. Each replicate contained 10 organisms. Reproduction and survival were monitored at 10 time points.

## Usage

```
data(cadmium1)
```

## Format

A data frame with 200 observations of the following five variables:

`replicate` A vector of class `numeric` with the replicate code (1 to 20).

`conc` A vector of class `numeric` with the cadmium concentrations in  $\mu\text{g.L}^{-1}$ .

`time` A vector of class `integer` with the time points (in days from the beginning of the experiment  $t = 0$ ).

`Nsurv` A vector of class `integer` with the number of alive individuals at each time point for each concentration and each replicate.

`Nrepro` A vector of class `integer` with the number of offspring at each time point for each concentration and each replicate.

## References

Billoir, E., Delhaye, H., Forfait, C., Clement, B., Triffault-Bouchet, G., Charles, S. and Delignette-Muller, M.L. (2012) Comparison of toxicity tests with different exposure time patterns: The added value of dynamic modelling in predictive ecotoxicology, *Ecotoxicology and Environmental Safety*, 75, 80-86.

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cadmium2	<i>Reproduction and survival data sets for Lymnaea stagnalis exposed to cadmium during 28 days</i>
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## Description

Reproduction and survival data sets of chronic laboratory toxicity tests with snails (*Lymnaea stagnalis*) exposed to six concentrations of cadmium during 28 days. Six concentrations were tested, with six replicates per concentration. Each replicate contained five organisms. Reproduction and survival were monitored at 17 time points.

## Usage

```
data(cadmium2)
```

## Format

A data frame with 612 observations of the following five variables:

`replicate` A vector of class `numeric` with the replicate code (1 to 36).

`conc` A vector of class `integer` with the cadmium concentrations in  $\mu\text{g}\cdot\text{L}^{-1}$ .

`time` A vector of class `integer` with the time points (in days from the beginning of the experiment  $t = 0$ ).

`Nsurv` A vector of class `integer` with the number of alive individuals at each time point for each concentration and each replicate.

`Nrepro` A vector of class `integer` with the number of clutches at each time point for each concentration and each replicate.

## References

Ducrot, V., Askem, C., Azam, D., Brettschneider, D., Brown, R., Charles, S., Coke, M., Collinet, M., Delignette-Muller, M.L., Forfait-Dubuc, C., Holbech, H., Hutchinson, T., Jach, A., Kinnberg, K.L., Lacoste, C., Le Page, G., Matthiessen, P., Oehlmann, J., Rice, L., Roberts, E., Ruppert, K., Davis, J.E., Veauvy, C., Weltje, L., Wortham, R. and Lagadic, L. (2014) Development and validation of an OECD reproductive toxicity test guideline with the pond snail *Lymnaea stagnalis* (Mollusca, Gastropoda), *Regulatory Toxicology and Pharmacology*, 70(3), 605-14.

Charles, S., Ducrot, V., Azam, D., Benstead, R., Brettschneider, D., De Schamphelaere, K., Filipe Goncalves, S., Green, J.W., Holbech, H., Hutchinson, T.H., Faber, D., Laranjeiro, F., Matthiessen, P., Norrgren, L., Oehlmann, J., Reategui-Zirena, E., Seeland-Fremer, A., Teigeler, M., Thome, J.P., Tobor Kaplon, M., Weltje, L., Lagadic, L. (2016) Optimizing the design of a reproduction toxicity

test with the pond snail *Lymnaea stagnalis*, *Regulatory Toxicology and Pharmacology*, vol. 81 pp.47-56.

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chlordan	<i>Reproduction and survival data sets for Daphnia magna exposed to chlordan during 21 days</i>
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### Description

Reproduction and survival data sets of chronic laboratory toxicity tests with *Daphnia magna* freshwater invertebrate exposed to six concentrations of one organochlorine insecticide (chlordan) during 21 days. Six concentrations were tested, with 10 replicates per concentration. Each replicate contained one organism. Reproduction and survival were monitored at 22 time points.

### Usage

```
data(chlordan)
```

### Format

A data frame with 1320 observations of the following five variables:

`replicate` A vector of class `numeric` with the replicate code (1 to 60).

`conc` A vector of class `numeric` with the chlordan concentrations in  $\mu\text{g.L}^{-1}$ .

`time` A vector of class `integer` with the time points (in days from the beginning of the experiment  $t = 0$ ).

`Nsurv` A vector of class `integer` with the number of alive individuals at each time point for each concentration and each replicate.

`Nrepro` A vector of class `integer` with the number of offspring at each time point for each concentration and each replicate.

### References

Manar, R., Bessi, H. and Vasseur, P. (2009) Reproductive effects and bioaccumulation of chlordan in *Daphnia magna*, *Environmental Toxicology and Chemistry*, 28, 2150-2159.

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copper	<i>Reproduction and survival data sets for Daphnia magna exposed to copper during 21 days</i>
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### Description

Reproduction and survival data sets of chronic laboratory toxicity tests with *Daphnia magna* freshwater invertebrate exposed to five concentrations of copper during 21 days. Five concentrations were tested, with three replicates per concentration. Each replicate contained 20 organisms. Reproduction and survival were monitored at 16 time points.

### Usage

```
data(copper)
```

### Format

A data frame with 240 observations of the following five variables:

`replicate` A vector of class `numeric` with the replicate code (1 to 15).

`conc` A vector of class `numeric` with the copper concentrations in  $\mu\text{g}\cdot\text{L}^{-1}$ .

`time` A vector of class `integer` with the time points (in days from the beginning of the experiment  $t = 0$ ).

`Nsurv` A vector of class `integer` with the number of alive individuals at each time point for each concentration and each replicate.

`Nrepro` A vector of class `integer` with the number of offspring at each time point for each concentration and each replicate.

### References

Billoir, E., Delignette-Muller, M.L., Pery, A.R.R. and Charles, S. (2008) A Bayesian Approach to Analyzing Ecotoxicological Data, *Environmental Science & Technology*, 42 (23), 8978-8984.

---

dichromate	<i>Survival data set for Daphnia magna exposed to dichromate during 21 days</i>
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### Description

Survival data set of chronic laboratory toxicity tests with *Daphnia magna* freshwater invertebrate exposed to six concentrations of one oxidizing agent (potassium dichromate) during 21 days. Six concentrations were tested with one replicate of 50 organisms per concentration. Survival is monitored at 10 time points.



**Usage**

```
data(dichromate)
```

**Format**

A data frame with 60 observations on the following four variables:

`replicate` A vector of class `numeric` with the replicate code (1).

`conc` A vector of class `numeric` with dichromate concentrations in  $mg.L^{-1}$ .

`time` A vector of class `integer` with the time points (in days from the beginning of the experiment  $t = 0$ ).

`Nsurv` A vector of class `integer` with the number of alive individuals at each time point for each concentration and each replicate.

**References**

Bedaux, J., Kooijman, SALM (1994) Statistical analysis of toxicity tests, based on hazard modeling, *Environmental and Ecological Statistics*, 1, 303-314.

---

FOCUSprofile

*A simulated exposure profile with 11641 time points.*

---

**Description**

Exposure profile of 11641 time points used for prediction.

**Usage**

```
data(FOCUSprofile)
```

**Format**

A data frame with 11641 observations on the following two variables:

`time` A vector of class `numeric`.

`conc` A vector of class `numeric` with exposure concentrations.and

`replicate` A vector of class `factor`.

---

`is_exposure_constant` *Test in a well-formed argument to function 'survData' if the concentration is constant and different from NA for each replicate (each time-serie)*

---

### Description

Test in a well-formed argument to function 'survData' if the concentration is constant and different from NA for each replicate (each time-serie)

### Usage

```
is_exposure_constant(x)
```

### Arguments

`x` an object of class `data.frame`

### Value

a boolean TRUE if concentration in replicate is constant, or FALSE if the concentration in at least one of the replicates is time-variable, and/or if NA occurs.

### Examples

```
# (1) Load the survival data set and test if concentration in replicates is constant
data("propiconazole")
is_exposure_constant(propiconazole)
is_exposure_constant(survData(propiconazole))

# (1) Load the survival data set and test if concentration in replicates is constant
data("propiconazole_pulse_exposure")
is_exposure_constant(propiconazole_pulse_exposure)
```

**Description**

Predict median and 95% credible interval of the  $x\%$  Lethal Concentration.

The function `LCx`,  $x\%$  Lethal Concentration ( $LC_x$ ), is use to compute the dose required to kill  $x\%$  of the members of a tested population after a specified test duration (`time_LCx`) (default is the maximum time point of the experiment).

Mathematical definition of  $x\%$  Lethal Concentration at time  $t$ , denoted  $LC(x, t)$ , is:

$$S(LC(x, t), t) = S(0, t) * (1 - x/100),$$

where  $S(LC(x, t), t)$  is the survival probability at concentration  $LC(x, t)$  at time  $t$ , and  $S(0, t)$  is the survival probability at no concentration (i.e. concentration is 0) at time  $t$  which reflect the background mortality  $h_b$ :

$$S(0, t) = \exp(-hb * t).$$

In the function `LCx`, we use the median of  $S(0, t)$  to rescale the  $x\%$  Lethal Concentration at time  $t$ .

**Usage**

```
LCx(object, ...)
```

```
## S3 method for class 'survFit'
```

```
LCx(object, X, time_LCx = NULL, conc_range = NULL, npoints = 100, ...)
```

**Arguments**

<code>object</code>	An object of class <code>survFit</code>
<code>...</code>	Further arguments to be passed to generic methods
<code>X</code>	Percentage of individuals dying (e.g., 50 for $LC_{50}$ , 10 for $LC_{10}$ , ...)
<code>time_LCx</code>	A number giving the time at which $LC_x$ has to be estimated. If <code>NULL</code> , the latest time point of the experiment is used.
<code>conc_range</code>	A vector of length 2 with minimal and maximal value of the range of concentration. If <code>NULL</code> , the range is define between 0 and the highest tested concentration of the experiment.
<code>npoints</code>	Number of time point in <code>conc_range</code> between 0 and the maximal concentration. 100 by default.

**Details**

When class of object is `survFit`, see [LCx.survFit](#).

**Value**

returns an object of class `LCx`.

The function returns an object of class `LCx`, which is a list with the following information:

<code>X_prop</code>	Survival probability of individuals surviving considering the median of the background mortality (i.e. $S(0, t) * (1 - x/100)$ )
---------------------	--

X_prop_provided	Survival probability of individuals surviving as provided in arguments (i.e. $(100 - X)/100$ )
time_LCx	A number giving the time at which $LC_x$ has to be estimated as provided in arguments or if NULL, the latest time point of the experiment is used.
df_LCx	A data.frame with quantiles (median, 2.5% and 97.5%) of $LC_X$ at time time_LCx for $X\%$ of individuals
df_dose	A data.frame with four columns: concentration, and median q50 and 95% credible interval (qinf95 and qsup95) of the survival probability at time time_LCx

### Examples

```
# (1) Load the data
data("propiconazole")

# (2) Create an object of class 'survData'
dataset <- survData(propiconazole)

# (3) Run the survFit function with model_type SD (or IT)
out_SD <- survFit(dataset, model_type = "SD")

# (4) estimate LC50 at time 4
LCx(out_SD, X = 50, time_LCx = 4)
```

---

MF<sub>x</sub>

*Predict the Multiplication Factor leading to  $x\%$  of reduction in survival at a specific time.*

---

### Description

Generic method for MF<sub>x</sub>, a function denoted  $MF(x, t)$  for  $x\%$  Multiplication Factor at time  $t$ .

The function MF<sub>x</sub>,  $x\%$  Multiplication Factor at time  $t$ , ( $MF(x, t)$ ), is used to compute the multiplication factor applied to the concentration exposure profile in order to reduce by  $x\%$  (argument  $X$ ) the survival probability at a specified test duration  $t$  (argument time\_MF<sub>x</sub>) (default is the maximum time point of the experiment).

Mathematical definition of  $x\%$  Multiplication Factor at time  $t$  (at the end of a time series  $T = \{0, \dots, t\}$ ), denoted  $MF(x, t)$ , is given by:

$$S(MF(x, t) * C_w(\tau \in T), t) = S(C_w(\tau \in T), t) * (1 - x/100),$$

where  $C_w(\tau \in T)$  is the initial exposure profile without multiplication factor. And so the expression  $S(MF(x, t) * C_w(\tau \in T), t)$  is the survival probability after an exposure profile  $MF(x, t) * C_w(\tau \in T)$  at time  $t$ .

This is a method to replace function MFx used on survFit object when computing issues happen. MFx\_ode uses the deSolve library to improve robustness. However, time to compute may be longer.

The function MFx\_ode,  $x\%$  Multiplication Factor at time  $t$ , ( $MF(x, t)$ ), is used to compute the multiplication factor applied to the concentration exposure profile in order to reduce by  $x\%$  (argument X) the survival probability at a specified test duration  $t$  (argument time\_MF<sub>x</sub>) (default is the maximum time point of the experiment).

Mathematical definition of  $x\%$  Multiplication Factor at time  $t$  (at the end of a time series  $T = \{0, \dots, t\}$ ), denoted  $MF(x, t)$ , is given by:

$$S(MF(x, t) * C_w(\tau \in T), t) = S(C_w(\tau \in T), t) * (1 - x/100),$$

where  $C_w(\tau \in T)$  is the initial exposure profile without multiplication factor. And so the expression  $S(MF(x, t) * C_w(\tau \in T), t)$  is the survival probability after an exposure profile  $MF(x, t) * C_w(\tau \in T)$  at time  $t$ .

## Usage

```
MFx(object, ...)
```

```
## S3 method for class 'survFit'
```

```
MFx(
  object,
  data_predict,
  X = 50,
  time_MFx = NULL,
  MFx_range = c(0, 1000),
  mcmc_size = 1000,
  hb_value = TRUE,
  spaghetti = FALSE,
  accuracy = 0.01,
  quiet = FALSE,
  threshold_iter = 100,
  hb_valueFORCED = 0,
  ode = TRUE,
  interpolate_length = NULL,
  interpolate_method = "linear",
  ...
)
```

```
MFx_ode(object, ...)
```

```
## S3 method for class 'survFit'
```

```
MFx_ode(
  object,
  data_predict,
  X = 50,
  time_MFx = NULL,
  MFx_range = c(0, 1000),
  mcmc_size = 1000,
```

```

    hb_value = FALSE,
    spaghetti = FALSE,
    accuracy = 0.01,
    quiet = FALSE,
    threshold_iter = 100,
    interpolate_length = NULL,
    interpolate_method = "linear",
    ...
)

```

### Arguments

object	An object of class <code>survFit</code> .
...	Further arguments to be passed to generic methods
data_predict	A dataframe with two columns <code>time</code> and <code>conc</code> .
X	Percentage of survival change (e.g., 50 for survival decrease of 50% , or -50 for survival increase of 50%).The default is 50. Only time series computed during the adaptation using a binary search in $O(\log(n))$ are returned. However, if NULL, all time series computed from the vector <code>MFx_range</code> are returned.
time_MFx	A number giving the time at which $MF(x, t)$ has to be estimated. If NULL, the latest time point of the profile is used.
MFx_range	A vector from which lower and upper bound of the range of the multiplication factor MFx are generated. The default is a vector <code>c(0, 1000)</code> . If argument X is NULL, then all the time series generated with <code>MFx_range</code> are returned.
mcmc_size	Can be used to reduce the number of MCMC samples in order to speed up the computation. The default is 1000.
hb_value	If TRUE, the background mortality hb is taken into account from the posterior. If FALSE, parameter hb is set to 0. The default is TRUE.
spaghetti	If TRUE, return a set of survival curves using parameters drawn from the posterior distribution.
accuracy	Accuracy of the multiplication factor. The default is 0.01.
quiet	If FALSE, print the evolution of accuracy.
threshold_iter	Threshold number of iteration.
hb_valueFORCED	If <code>hb_value</code> is FALSE, it fix hb.
ode	IF <code>ode</code> is TRUE, also use <code>predict_ode</code> rather than <code>predict</code> . Default is TRUE.
interpolate_length	Length of the time sequence for which output is wanted.
interpolate_method	The interpolation method for concentration. See package <code>deSolve</code> for details. Default is <code>linear</code> .

### Details

When class of object is `survFit`, see [MFx.survFit](#).

**Value**

returns an object of class MFx

The function returns an object of class MFx, which is a list with the following information:

X_prop	Survival probability for X percent of reduction of the initial median survival probability at time time_MF <sub>x</sub> .
X_prop_provided	A number giving the proportion of reduction in survival.
time_MF <sub>x</sub>	A number giving the time at which $MF(x, t)$ has to be estimated as provided in arguments or if NULL, the latest time point of the profile is used.
df_MF <sub>x</sub>	A data.frame with quantiles (median, 2.5% and 97.5%) of $MF(x, t)$ at time $t$ , time_MF <sub>x</sub> , for $x\%$ of survival reduction.
df_dose	A data.frame with quantiles (median, 2.5% and 97.5%) of survival probability along the computed multiplication factor and at time time_MF <sub>x</sub> .
MF <sub>x</sub> _tested	A vector of all multiplication factors computed.
ls_predict	A list of all object of class survFitPredict obtained from computing survival probability for every profiles build from the vector of multiplication factors MF <sub>x</sub> _tested.

The function returns an object of class MFx, which is a list with the following information:

X_prop	Survival probability for X percent of reduction of the initial median survival probability at time time_MF <sub>x</sub> .
X_prop_provided	A number giving the proportion of reduction in survival.
time_MF <sub>x</sub>	A number giving the time at which $MF(x, t)$ has to be estimated as provided in arguments or if NULL, the latest time point of the profile is used.
df_MF <sub>x</sub>	A data.frame with quantiles (median, 2.5% and 97.5%) of $MF(x, t)$ at time $t$ , time_MF <sub>x</sub> , for $x\%$ of survival reduction.
df_dose	A data.frame with quantiles (median, 2.5% and 97.5%) of survival probability along the computed multiplication factor and at time time_MF <sub>x</sub> .
MF <sub>x</sub> _tested	A vector of all multiplication factors computed.
ls_predict	A list of all object of class survFitPredict obtained from computing survival probability for every profiles build from the vector of multiplication factors MF <sub>x</sub> _tested.

**Examples**

```
# (1) Load the data
data("propiconazole")

# (2) Create an object of class 'survData'
dataset <- survData(propiconazole)
```

```

# (3) Run the survFit function with model_type SD (or IT)
out_SD <- survFit(dataset, model_type = "SD")

# (4) data to predict
data_4prediction <- data.frame(time = 1:10, conc = c(0,0.5,3,3,0,0,0.5,3,1.5,0))

# (5) estimate MF(x=30, t=4), that is for 30% reduction of survival at time 4
MFx_SD_30.4 <- MFx(out_SD, data_predict = data_4prediction , X = 30, time_MFx = 4)

# (5bis) estimate MF(x,t) along the MF_range from 5 to 10 (50) (X = NULL)
MFx_SD_range <- MFx(out_SD, data_predict = data_4prediction ,
                    X = NULL, time_MFx = 4, MFx_range = seq(5, 10, length.out = 50))

# (1) Load the data
data("propiconazole")

# (2) Create an object of class 'survData'
dataset <- survData(propiconazole)

# (3) Run the survFit function with model_type SD (or IT)
out_SD <- survFit(dataset, model_type = "SD")

# (4) data to predict
data_4prediction <- data.frame(time = 1:10, conc = c(0,0.5,3,3,0,0,0.5,3,1.5,0))

# (5) estimate MF(x=30, t=4), that is for 30% reduction of survival at time 4
MFx_SD_30.4 <- MFx_ode(out_SD, data_predict = data_4prediction , X = 30, time_MFx = 4)

# (5bis) estimate MF(x,t) along the MF_range from 5 to 10 (50) (X = NULL)
MFx_SD_range <- MFx_ode(out_SD, data_predict = data_4prediction ,
                      X = NULL, time_MFx = 4, MFx_range = seq(5, 10, length.out = 50))

```

---

modelData

*Create a list giving data to use in Bayesian inference.*

---

### Description

Create a list giving data to use in Bayesian inference.

### Usage

```
modelData(x, ...)
```



**Arguments**

x                    An object of class survData  
 ...                  Further arguments to be passed to generic methods

**Value**

A list for parameterization of priors for Bayesian inference.

---

```
modelData.survDataCstExp
```

*Create a data set to analyse a survDataCstExp object.*

---

**Description**

Create a data set to analyse a survDataCstExp object.

**Usage**

```
## S3 method for class 'survDataCstExp'  
modelData(x, model_type = NULL)
```

**Arguments**

x                    An object of class survData  
 model\_type        TKTD GUTS model type ('SD' or 'IT')

**Value**

A list for parameterization of priors for Bayesian inference.

---

```
modelData.survDataVarExp
```

*Create a data set to analyse a survDataVarExp object.*

---

**Description**

Create a data set to analyse a survDataVarExp object.

**Usage**

```
## S3 method for class 'survDataVarExp'  
modelData(x, model_type = NULL, extend_time = 100, ...)
```

**Arguments**

x	An object of class survData
model_type	TKTD GUTS model type ('SD' or 'IT')
extend_time	Number of for each replicate used for linear interpolation (comprise between time to compute and fitting accuracy)
...	Further arguments to be passed to generic methods

**Value**

A list for parameterization of priors for Bayesian inference.

---

plot.LCx	<i>Plotting method for LCx objects</i>
----------	--

---

**Description**

This is the generic plot S3 method for the `LCx` class. It plots the survival probability as a function of concentration.

**Usage**

```
## S3 method for class 'LCx'
plot(
  x,
  xlab = "Concentration",
  ylab = "Survival probability \n median and 95 CI",
  main = NULL,
  subtitle = NULL,
  ...
)
```

**Arguments**

x	An object of class LCx.
xlab	A label for the $X$ -axis, by default Concentration.
ylab	A label for the $Y$ -axis, by default Survival probability median and 95 CI.
main	A main title for the plot.
subtitle	A subtitle for the plot
...	Further arguments to be passed to generic methods.

**Value**

a plot of class ggplot

**Examples**

```

# (1) Load the data
data("propiconazole")

# (2) Create an object of class 'survData'
dataset <- survData(propiconazole)

# (3) Run the survFit function with model_type SD (or IT)
out_SD <- survFit(dataset, model_type = "SD")

# (4) estimate LC50 at time 4
LCx_SD <- LCx(out_SD, X = 50, time_LCx = 4)

# (5) plot the object of class 'LCx'
plot(LCx_SD)

```

---

plot.MFx

*Plotting method for MFx objects*


---

**Description**

This is the generic plot S3 method for the MFx class. It plots the survival probability as a function of the multiplication factor applied or as a function of time.

**Usage**

```

## S3 method for class 'MFx'
plot(
  x,
  x_variable = "MFx",
  xlab = NULL,
  ylab = "Survival probability \n median and 95 CI",
  main = NULL,
  log_scale = FALSE,
  ncol = 3,
  ...
)

```

**Arguments**

x	An object of class MFx.
x_variable	A character to define the variable for the X-axis, either "MFx" or "Time". The default is "MFx".
xlab	A label for the X-axis, by default NULL and depend on the argument x_variable.

<code>ylab</code>	A label for the <i>Y</i> -axis, by default Survival probability median and 95 CI.
<code>main</code>	A main title for the plot.
<code>log_scale</code>	If TRUE, the x-axis is log-scaled. Default is FALSE.
<code>ncol</code>	An interger for the number of columns when several panels are plotted.
<code>...</code>	Further arguments to be passed to generic methods.

**Value**

a plot of class `ggplot`

**Examples**

```
# (1) Load the data
data("propiconazole")

# (2) Create an object of class 'survData'
dataset <- survData(propiconazole)

# (3) Run the survFit function with model_type SD (or IT)
out_SD <- survFit(dataset, model_type = "SD")

# (4) data to predict
data_4prediction <- data.frame(time = 1:10, conc = c(0,0.5,3,3,0,0,0.5,3,1.5,0))

# (5) estimate MF for 30% reduction of survival at time 4
MFx_SD_30.4 <- MFx(out_SD, data_predict = data_4prediction , X = 30, time_MFx = 4)

# (6) plot the object of class 'MFx'
plot(MFx_SD_30.4)

# (6bis) plot with log-scale of x-axis
plot(MFx_SD_30.4, log_scale = TRUE)

# (6ter) plot with "Time" as the x-axis
plot(MFx_SD_30.4, x_variable = "Time")

# (7) plot when X = NULL and along a MFx_range from 5 to 10:
MFx_SD_range <- MFx(out_SD, data_predict = data_4prediction ,
  X = NULL, time_MFx = 4, MFx_range = seq(5, 10, length.out = 50))
plot(MFx_SD_range)
plot(MFx_SD_range, x_variable = "Time", ncol = 10)
```

---

plot.reproData      *Plotting method for reproData objects*

---

### Description

This is the generic plot S3 method for the reproData class. It plots the cumulated number of offspring as a function of time.

### Usage

```
## S3 method for class 'reproData'
plot(
  x,
  xlab,
  ylab = "Cumulated Number of offspring",
  main = NULL,
  concentration = NULL,
  style = "ggplot",
  pool.replicate = FALSE,
  addlegend = FALSE,
  remove.someLabels = FALSE,
  ...
)
```

### Arguments

x	an object of class reproData
xlab	label of the X-axis
ylab	label of the Y-axis, by default Cumulated Number of offspring
main	main title for the plot
concentration	a numeric value corresponding to some concentration in data. If concentration = NULL, draws a plot for each concentration
style	graphical backend, can be 'ggplot' or 'generic'
pool.replicate	if TRUE, the datapoints of each replicate are summed for a same concentration
addlegend	if TRUE, adds a default legend to the plot
remove.someLabels	if TRUE, removes 3/4 of X-axis labels in 'ggplot' style to avoid the label overlap
...	Further arguments to be passed to generic methods

### Value

a plot of class ggplot

**Note**

When `style = "generic"`, the function calls the generic function `plot`

When `style = "ggplot"`, the function return an object of class `gg` and `ggplot`, see function `ggplot`

**Examples**

```
# (1) Load the data
data(cadmium1)

# (2) Create an object of class 'reproData'
cadmium1 <- reproData(cadmium1)

# (3) Plot the reproduction data
plot(cadmium1)

# (4) Plot the reproduction data for a fixed concentration
plot(cadmium1, concentration = 4.36, style = "generic")
```

---

`plot.reproFitTT`      *Plotting method for reproFitTT objects*

---

**Description**

This is the generic `plot` S3 method for the `reproFitTT` class. It plots the concentration-effect fit under target time reproduction analysis.

**Usage**

```
## S3 method for class 'reproFitTT'
plot(
  x,
  xlab = "Concentration",
  ylab = "Nb of offspring per ind/day",
  main = NULL,
  fitcol = "orange",
  fitlty = 1,
  fitlwd = 1,
  spaghetti = FALSE,
  cicol = "orange",
  cilty = 2,
  cilwd = 1,
  ribcol = "grey70",
  addlegend = FALSE,
  log.scale = FALSE,
  style = "ggplot",
  ...
)
```

**Arguments**

x	an object of class reproFitTT
xlab	a label for the $X$ -axis, by default Concentration
ylab	a label for the $Y$ -axis, by default Nb of offspring per ind/day
main	main title for the plot
fitcol	color of the fitted curve
fitlty	line type of the fitted curve
fitlwd	width of the fitted curve
spaghetti	if TRUE, the credible interval is represented by multiple curves
cicol	color of the 95 % credible limits
cilty	line type of the 95 % credible limits
cilwd	width of the 95 % credible limits
ribcol	color of the ribbon between lower and upper credible limits. Transparent if NULL
addlegend	if TRUE, adds a default legend to the plot
log.scale	if TRUE, displays $X$ -axis in log-scale
style	graphical backend, can be 'ggplot' or 'generic'
...	Further arguments to be passed to generic methods

**Details**

The fitted curve represents the **estimated reproduction rate** at the target time as a function of the chemical compound concentration. The function plots 95% credible intervals for the estimated reproduction rate (by default the grey area around the fitted curve). Typically a good fit is expected to display a large overlap between the two types of intervals. If spaghetti = TRUE, the credible intervals are represented by two dotted lines limiting the credible band, and a spaghetti plot is added to this band. It consists of the representation of simulated curves using parameter values sampled in the posterior distribution (10% of the MCMC chains are randomly taken for this sample).

**Value**

a plot of class ggplot

**Note**

When style = "generic", the function calls the generic function [plot](#)

When style = "ggplot", the function return an object of class ggplot, see function [ggplot](#)

**Examples**

```
# (1) Load the data
data(cadmium1)

# (2) Create an object of class "reproData"
```

```

dataset <- reproData(cadmium1)

# (3) Run the reproFitTT function with the log-logistic gamma-Poisson model
out <- reproFitTT(dataset, stoc.part = "gammapoisson",
                  ecx = c(5, 10, 15, 20, 30, 50, 80), quiet = TRUE)

# (4) Plot the fitted curve with generic style
plot(out, xlab = expression("Concentration in" ~ mu~g.L^{-1}),
      fitcol = "blue", cicol = "lightblue",
      main = "Log-logistic response to concentration")

```

---

plot.survDataCstExp    *Plotting method for survData objects*

---

### Description

This is the generic plot S3 method for the survData class. It plots the number of survivors as a function of time.

### Usage

```

## S3 method for class 'survDataCstExp'
plot(
  x,
  xlab = "Time",
  ylab = "Number of survivors",
  main = NULL,
  concentration = NULL,
  style = "ggplot",
  pool.replicate = FALSE,
  addlegend = FALSE,
  remove.someLabels = FALSE,
  ...
)

```

### Arguments

x	an object of class survData
xlab	a label for the X-axis, by default Time
ylab	a label for the Y-axis, by default Number of survivors
main	main title for the plot
concentration	a numeric value corresponding to some concentration(s) in data. If concentration = NULL, draws a plot for each concentration



**style** graphical backend, can be 'generic' or 'ggplot'  
**pool.replicate** if TRUE, the datapoints of each replicate are summed for a same concentration  
**addlegend** if TRUE, adds a default legend to the plot  
**remove.someLabels**  
if TRUE, removes 3/4 of X-axis labels in 'ggplot' style to avoid label overlap  
**...** Further arguments to be passed to generic methods

**Value**

a plot of class ggplot

**Note**

When style = "ggplot" (default), the function calls function [ggplot](#) and returns an object of class ggplot.

**Examples**

```

# (1) Load the data
data(zinc)
zinc <- survData(zinc)

# (2) Plot survival data with a ggplot style
plot(zinc)

# (3) Plot the survival data for one specific concentration
plot(zinc, concentration = 0.66)

```

---

plot.survDataVarExp *Plotting method for survDataVarExp objects*

---

**Description**

This is the generic plot S3 method for the survDataVarC class. It plots the number of survivors as a function of time.

**Usage**

```

## S3 method for class 'survDataVarExp'
plot(
  x,
  xlab = "Time",
  ylab = "Number of survivors",
  main = NULL,
  one.plot = FALSE,
  facetting_level = NULL,
  ...
)

```

**Arguments**

x	an object of class survDataVarExp
xlab	a label for the <i>X</i> -axis, by default Time
ylab	a label for the <i>Y</i> -axis, by default Number of survivors
main	main title for the plot
one.plot	if TRUE, draws all the points in one plot instead of one per replicate
facetting_level	a vector of characters to rank replicates in the multi plot (i.e. one.plot == FALSE)
...	Further arguments to be passed to generic methods

**Value**

an object of class ggplot, see function [ggplot](#)

---

plot.survFitCstExp     *Plotting method for survFit objects*

---

**Description**

This is the generic plot S3 method for the survFit. It plots the fit obtained for each concentration of chemical compound in the original dataset.

**Usage**

```
## S3 method for class 'survFitCstExp'
plot(
  x,
  xlab = "Time",
  ylab = "Survival probability",
  main = NULL,
  concentration = NULL,
  spaghetti = FALSE,
  one.plot = FALSE,
  adddata = TRUE,
  addlegend = FALSE,
  style = "ggplot",
  ...
)
```

**Arguments**

<code>x</code>	An object of class <code>survFit</code> .
<code>xlab</code>	A label for the $X$ -axis, by default <code>Time</code> .
<code>ylab</code>	A label for the $Y$ -axis, by default <code>Survival probability</code> .
<code>main</code>	A main title for the plot.
<code>concentration</code>	A numeric value corresponding to some specific concentrations in data. If <code>concentration = NULL</code> , draws a plot for each concentration.
<code>spaghetti</code>	if <code>TRUE</code> , draws a set of survival curves using parameters drawn from the posterior distribution
<code>one.plot</code>	if <code>TRUE</code> , draws all the estimated curves in one plot instead of one plot per concentration.
<code>adddata</code>	if <code>TRUE</code> , adds the observed data to the plot with (frequentist binomial) confidence intervals
<code>addlegend</code>	if <code>TRUE</code> , adds a default legend to the plot.
<code>style</code>	graphical backend, can be <code>'generic'</code> or <code>'ggplot'</code>
<code>...</code>	Further arguments to be passed to generic methods.

**Details**

The fitted curves represent the **estimated survival probability** as a function of time for each concentration. The black dots depict the **observed survival probability** at each time point. Note that since our model does not take inter-replicate variability into consideration, replicates are systematically pooled in this plot. The function plots both 95% credible intervals for the estimated survival probability (by default the grey area around the fitted curve) and 95% binomial confidence intervals for the observed survival probability (as black error bars if `adddata = TRUE`). Both types of intervals are taken at the same level. Typically a good fit is expected to display a large overlap between the two types of intervals. If `spaghetti = TRUE`, the credible intervals are represented by two dotted lines limiting the credible band, and a spaghetti plot is added to this band. This spaghetti plot consists of the representation of simulated curves using parameter values sampled in the posterior distribution (2% of the MCMC chains are randomly taken for this sample).

**Value**

a plot of class `ggplot`

---

`plot.survFitPredict` *Plotting method for survFitPredict objects*

---

**Description**

This is the generic plot S3 method for the `survFitPredict`. It plots the predicted survival probability for each concentration of the chemical compound in the provided dataset.

**Usage**

```
## S3 method for class 'survFitPredict'
plot(
  x,
  xlab = "Time",
  ylab = "Survival probability",
  main = NULL,
  spaghetti = FALSE,
  one.plot = FALSE,
  mcmc_size = NULL,
  ...
)
```

**Arguments**

x	An object of class <code>survFitPredict</code> .
xlab	A label for the <i>X</i> -axis, by default <code>Time</code> .
ylab	A label for the <i>Y</i> -axis, by default <code>Survival probability</code> .
main	A main title for the plot.
spaghetti	If <code>TRUE</code> , draws a set of survival curves using parameters drawn from the posterior distribution
one.plot	if <code>TRUE</code> , draws all the estimated curves in one plot instead of one plot per concentration.
mcmc_size	A numerical value referring by default to the size of the mcmc in object <code>survFitPredict</code> . This option is specific to <code>survFitPredict</code> objects for which computing time may be long. <code>mcmc_size</code> can be used to reduce the number of mcmc samples in order to speed up the computation.
...	Further arguments to be passed to generic methods.

**Details**

The fitted curves represent the **predicted survival probability** as a function of time for each concentration. The function plots both the 95% credible band and the predicted survival probability over time. If `spaghetti = TRUE`, the credible intervals are represented by two dotted lines limiting the credible band, and a spaghetti plot is added to this band. This spaghetti plot consists of the representation of simulated curves using parameter values sampled in the posterior distribution (10% of the MCMC chains are randomly taken for this sample).

**Value**

a plot of class `ggplot`

**Examples**

```
# (1) Load the survival data
data("propiconazole_pulse_exposure")
```

```
# (2) Create an object of class "survData"
dataset <- survData(propiconazole_pulse_exposure)

# (3) Run the survFit function
out <- survFit(dataset , model_type = "SD")

# (4) Create a new data table for prediction
data_4prediction <- data.frame(time = 1:10, conc = c(0,5,5,5,0,0,5,5,5,5),
  replicate= rep("predict", 10))

# (5) Predict on a new dataset
predict_out <- predict(out, data_predict = data_4prediction, spaghetti = TRUE)

# (6) Plot the predicted curve
plot(predict_out)
plot(predict_out, spaghetti = TRUE)
```

---

plot.survFitPredict\_Nsurv

*Plotting method for survFitPredict\_Nsurv objects.*

---

## Description

This is the generic plot S3 method for the survFitPredict\_Nsurv. It plots the predicted survival probability for each concentration of the chemical compound in the provided dataset.

## Usage

```
## S3 method for class 'survFitPredict_Nsurv'
plot(
  x,
  xlab = "Time",
  ylab = "Number of survivors",
  main = NULL,
  spaghetti = FALSE,
  one.plot = FALSE,
  mcmc_size = NULL,
  ...
)
```

## Arguments

x	An object of class survFitPredict_Nsurv.
xlab	A label for the X-axis, by default Time.

ylab	A label for the Y-axis, by default Survival probability.
main	A main title for the plot.
spaghetti	If TRUE, draws a set of survival curves using parameters drawn from the posterior distribution
one.plot	if TRUE, draws all the estimated curves in one plot instead of one plot per concentration.
mcmc_size	A numerical value refering by default to the size of the mcmc in object survFitPredict. This option is specific to survFitPredict objects for which computing time may be long. mcmc_size can be used to reduce the number of mcmc samples in order to speed up the computation.
...	Further arguments to be passed to generic methods.

### Details

The fitted curves represent the **predicted survival probability** as a function of time for each concentration. The function plots both the 95% credible band and the predicted survival probability over time. If spaghetti = TRUE, the credible intervals are represented by two dotted lines limiting the credible band, and a spaghetti plot is added to this band. This spaghetti plot consists of the representation of simulated curves using parameter values sampled in the posterior distribution (10% of the MCMC chains are randomly taken for this sample).

### Value

a plot of class ggplot

---

plot.survFitTKTD      *Plotting method for survFitTKTD objects*

---

### Description

This is the generic plot S3 method for the survFitTKTD. It plots the fit obtained for each concentration of chemical compound in the original dataset.

### Usage

```
## S3 method for class 'survFitTKTD'
plot(
  x,
  xlab = "Time",
  ylab = "Survival probablity",
  main = NULL,
  concentration = NULL,
  spaghetti = FALSE,
  one.plot = FALSE,
  adddata = FALSE,
  addlegend = FALSE,
```

```

    style = "ggplot",
    ...
  )

```

### Arguments

<code>x</code>	An object of class <code>survFitTKTD</code> .
<code>xlab</code>	A label for the $X$ -axis, by default <code>Time</code> .
<code>ylab</code>	A label for the $Y$ -axis, by default <code>Survival probability</code> .
<code>main</code>	A main title for the plot.
<code>concentration</code>	A numeric value corresponding to some specific concentration in data. If <code>concentration = NULL</code> , draws a plot for each concentration.
<code>spaghetti</code>	if <code>TRUE</code> , draws a set of survival curves using parameters drawn from the posterior distribution
<code>one.plot</code>	if <code>TRUE</code> , draws all the estimated curves in one plot instead of one plot per concentration.
<code>adddata</code>	if <code>TRUE</code> , adds the observed data to the plot with (frequentist binomial) confidence intervals
<code>addlegend</code>	if <code>TRUE</code> , adds a default legend to the plot.
<code>style</code>	graphical backend, can be <code>'generic'</code> or <code>'ggplot'</code>
<code>...</code>	Further arguments to be passed to generic methods.

### Details

The fitted curves represent the **estimated survival probability** as a function of time for each concentration. When `adddata = TRUE` the black dots depict the **observed survival probability** at each time point. Note that since our model does not take inter-replicate variability into consideration, replicates are systematically pooled in this plot. The function plots both 95% credible intervals for the estimated survival probability (by default the grey area around the fitted curve) and 95% binomial confidence intervals for the observed survival probability (as black error bars if `adddata = TRUE`). Both types of intervals are taken at the same level. Typically a good fit is expected to display a large overlap between the two types of intervals. If `spaghetti = TRUE`, the credible intervals are represented by two dotted lines limiting the credible band, and a spaghetti plot is added to this band. This spaghetti plot consists of the representation of simulated curves using parameter values sampled in the posterior distribution (2% of the MCMC chains are randomly taken for this sample).

### Value

a plot of class `ggplot`

### Examples

```

# (1) Load the survival data
data(propiconazole)

# (2) Create an object of class "survData"

```

```

dataset <- survData(propiconazole)

# (3) Run the survFitTKTD function ('SD' model only)
out <- survFitTKTD(dataset)

# (4) Plot the fitted curves in one plot
plot(out)

# (5) Plot one fitted curve per concentration with credible limits as
# spaghetti, data and confidence intervals
# and with a ggplot style
plot(out, spaghetti = TRUE , adddata = TRUE, one.plot = FALSE,
      style = "ggplot")

# (6) Plot fitted curve for one specific concentration
plot(out, concentration = 36, style = "ggplot")

```

---

plot.survFitTT

*Plotting method for survFitTT objects*


---

### Description

This is the generic plot S3 method for the survFitTT class. It plots concentration-response fit under target time survival analysis.

### Usage

```

## S3 method for class 'survFitTT'
plot(
  x,
  xlab = "Concentration",
  ylab = "Survival probability",
  main = NULL,
  fitcol = "orange",
  fitlty = 1,
  fitlwd = 1,
  spaghetti = FALSE,
  cicol = "orange",
  cilty = 2,
  cilwd = 1,
  ribcol = "grey70",
  adddata = FALSE,
  addlegend = FALSE,
  log.scale = FALSE,
  style = "ggplot",
  ...
)

```



### Arguments

x	an object of class survFitTT
xlab	a label for the <i>X</i> -axis, default is Concentration
ylab	a label for the <i>Y</i> -axis, default is Survival probability
main	main title for the plot
fitcol	color of the fitted curve
fitlty	line type of the fitted curve
fitlwd	width of the fitted curve
spaghetti	if TRUE, the credible interval is represented by multiple curves
cicol	color of the 95 % credible interval limits
cilty	line type for the 95 % credible interval limits
cilwd	width of the 95 % credible interval limits
ribcol	color of the ribbon between lower and upper credible limits. Transparent if NULL
adddata	if TRUE, adds the observed data with confidence intervals to the plot
addlegend	if TRUE, adds a default legend to the plot
log.scale	if TRUE, displays <i>X</i> -axis in log-scale
style	graphical backend, can be 'generic' or 'ggplot'
...	Further arguments to be passed to generic methods

### Details

The fitted curve represents the **estimated survival probability** at the target time as a function of the concentration of chemical compound; When `adddata = TRUE` the black dots depict the **observed survival probability** at each tested concentration. Note that since our model does not take inter-replicate variability into consideration, replicates are systematically pooled in this plot. The function plots both 95% credible intervals for the estimated survival probability (by default the grey area around the fitted curve) and 95% binomial confidence intervals for the observed survival probability (as black segments if `adddata = TRUE`). Both types of intervals are taken at the same level. Typically a good fit is expected to display a large overlap between the two intervals. If `spaghetti = TRUE`, the credible intervals are represented by two dotted lines limiting the credible band, and a spaghetti plot is added to this band. This spaghetti plot consists of the representation of simulated curves using parameter values sampled in the posterior distribution (10% of the MCMC chains are randomly taken for this sample).

### Value

a plot of class `ggplot`

### Note

When `style = "ggplot"`, the function calls function `ggplot` and returns an object of class `ggplot`.

**Examples**

```

# (1) Load the data
data(cadmium1)

# (2) Create an object of class "survData"
dat <- survData(cadmium1)

# (3) Run the survFitTT function with the log-logistic
#       binomial model
out <- survFitTT(dat, lcx = c(5, 10, 15, 20, 30, 50, 80),
                 quiet = TRUE)

# (4) Plot the fitted curve
plot(out, log.scale = TRUE, adddata = TRUE)

# (5) Plot the fitted curve with ggplot style
plot(out, xlab = expression("Concentration in" ~ mu~g.L^{-1}),
      fitcol = "blue", adddata = TRUE, cicol = "blue",
      style = "ggplot")

```

---

plot.survFitVarExp      *Plotting method for survFit objects*

---

**Description**

This is the generic plot S3 method for the survFit. It plots the fit obtained for each concentration profile in the original dataset.

**Usage**

```

## S3 method for class 'survFitVarExp'
plot(
  x,
  xlab = "Time",
  ylab = "Survival probability",
  main = NULL,
  spaghetti = FALSE,
  one.plot = FALSE,
  adddata = TRUE,
  mcmc_size = NULL,
  scales = "fixed",
  addConfInt = TRUE,
  ...
)

```

**Arguments**

<code>x</code>	An object of class <code>survFit</code> .
<code>xlab</code>	A label for the $X$ -axis, by default <code>Time</code> .
<code>ylab</code>	A label for the $Y$ -axis, by default <code>Survival probability</code> .
<code>main</code>	A main title for the plot.
<code>spaghetti</code>	if TRUE, draws a set of survival curves using parameters drawn from the posterior distribution
<code>one.plot</code>	if TRUE, draws all the estimated curves in one plot instead of one plot per concentration.
<code>adddata</code>	if TRUE, adds the observed data to the plot.
<code>mcmc_size</code>	A numerical value referring by default to the size of the mcmc in object <code>survFit</code> . This option is specific to <code>survFitVarExp</code> objects for which computing time may be long. <code>mcmc_size</code> can be used to reduce the number of mcmc samples in order to speed up the computation.
<code>scales</code>	Shape the scale of axis. Default is <code>"fixed"</code> , but can be <code>"free"</code> , or free in only one dimension <code>"free_x"</code> , <code>"free_y"</code> . (See <code>ggplot2</code> documentation for more details.)
<code>addConfInt</code>	If TRUE, add a 95% confidence interval on observed data from a binomial test
<code>...</code>	Further arguments to be passed to generic methods.

**Details**

The fitted curves represent the **estimated survival probability** as a function of time for each concentration profile. The black dots depict the **observed survival probability** at each time point. Note that since our model does not take inter-replicate variability into consideration, replicates are systematically pooled in this plot. The function plots both 95% binomial credible intervals for the estimated survival probability (by default the grey area around the fitted curve) and 95% binomial confidence intervals for the observed survival probability (as black segments if `adddata = TRUE`). Both types of intervals are taken at the same level. Typically a good fit is expected to display a large overlap between the two types of intervals. If `spaghetti = TRUE`, the credible intervals are represented by two dotted lines limiting the credible band, and a spaghetti plot is added to this band. This spaghetti plot consists of the representation of simulated curves using parameter values sampled in the posterior distribution (10% of the MCMC chains are randomly taken for this sample).

**Value**

a plot of class `ggplot`

**Examples**

```
# (1) Load the survival data
data("propiconazole_pulse_exposure")

# (2) Create an object of class "survData"
dataset <- survData(propiconazole_pulse_exposure)
```

```

# (3) Run the survFit function
out <- survFit(dataset , model_type = "SD")

# (4) Summary look the estimated values (parameters)
summary(out)

# (5) Plot the fitted curve
plot(out, adddata = FALSE)

# (6) Plot the fitted curve with ggplot style and CI as spaghetti
plot(out, spaghetti = TRUE)

```

---

plotDoseResponse      *Plot dose-response from raw data*

---

### Description

Plots the response of the effect as a function of the concentration at a given target time.

### Usage

```
plotDoseResponse(x, ...)
```

### Arguments

x                      an object used to select a method plotDoseRespons  
...                      Further arguments to be passed to generic methods

### Value

a plot of class ggplot

---

plotDoseResponse.reproData  
*Plot dose-response from reproData objects*

---

### Description

This is the generic plotDoseResponse S3 method for the reproData class. It plots the number of offspring per individual-days as a function of concentration at a given target time.

**Usage**

```
## S3 method for class 'reproData'
plotDoseResponse(
  x,
  xlab = "Concentration",
  ylab = "Nb of offspring per ind.day",
  main = NULL,
  ylim = NULL,
  target.time = NULL,
  style = "ggplot",
  log.scale = FALSE,
  remove.someLabels = FALSE,
  axis = TRUE,
  addlegend = TRUE,
  ...
)
```

**Arguments**

<code>x</code>	an object of class <code>reproData</code>
<code>xlab</code>	a label for the <i>X</i> -axis, by default <code>Concentration</code>
<code>ylab</code>	a label for the <i>Y</i> -axis, by default <code>Nb of offspring per ind.day</code>
<code>main</code>	main title for the plot
<code>ylim</code>	<i>Y</i> -axis limits
<code>target.time</code>	a numeric value corresponding to some observed time points in data
<code>style</code>	graphical backend, can be <code>'ggplot'</code> or <code>'generic'</code>
<code>log.scale</code>	if <code>TRUE</code> , displays <i>X</i> -axis in log-scale
<code>remove.someLabels</code>	if <code>TRUE</code> , removes 75% of <i>X</i> -axis labels in <code>'ggplot'</code> style to avoid the label overlap
<code>axis</code>	if <code>TRUE</code> displays ticks and label axis
<code>addlegend</code>	if <code>TRUE</code> , adds a default legend to the plot
<code>...</code>	Further arguments to be passed to generic methods

**Details**

The function plots the observed values of the reproduction rate (number of reproduction outputs per individual-day) at a given time point as a function of concentration. The 95 % Poisson confidence interval is added to each reproduction rate. It is calculated using function `pois.exact` from package `epi.tools`. As replicates are not pooled in this plot, overlapped points are shifted on the *x*-axis to help the visualization of replicates.

**Value**

a plot of class `ggplot`

**Note**

When `style = "generic"`, the function calls the generic function `plot`

When `style = "ggplot"`, the function return an object of class `ggplot`, see function `ggplot`

**See Also**

`pois.exact`

**Examples**

```
# (1) Load the data
data(zinc)

# (2) Create an object of class 'reproData'
zinc_rpr <- reproData(zinc)

# (3) Plot dose-response
plotDoseResponse(zinc_rpr)

# (4) Plot dose-response with a generic style
plotDoseResponse(zinc_rpr, style = "generic")
```

---

`plotDoseResponse.survDataCstExp`

*Plot dose-response from survData objects*

---

**Description**

This is the generic `plotDoseResponse` S3 method for the `survData` class. It plots the survival probability as a function of concentration at a given target time.

**Usage**

```
## S3 method for class 'survDataCstExp'
plotDoseResponse(
  x,
  xlab = "Concentration",
  ylab = "Survival probability",
  main = NULL,
  target.time = NULL,
  style = "ggplot",
  log.scale = FALSE,
  remove.someLabels = FALSE,
  addlegend = TRUE,
  ...
)
```

## Arguments

<code>x</code>	an object of class <code>survData</code>
<code>xlab</code>	a label for the $X$ -axis, by default <code>Concentration</code>
<code>ylab</code>	a label for the $Y$ -axis, by default <code>Survival probability</code>
<code>main</code>	main title for the plot
<code>target.time</code>	a numeric value corresponding to some observed time in data
<code>style</code>	graphical backend, can be <code>'ggplot'</code> or <code>'generic'</code>
<code>log.scale</code>	if <code>TRUE</code> , displays $X$ -axis in log-scale
<code>remove.someLabels</code>	if <code>TRUE</code> , removes 75% of $X$ -axis labels in <code>'ggplot'</code> style to avoid the label overlap
<code>addlegend</code>	if <code>TRUE</code> , adds a default legend to the plot
<code>...</code>	Further arguments to be passed to generic methods

## Details

The function plots the observed values of the survival probability at a given time point as a function of concentration. The 95 % binomial confidence interval is added to each survival probability. It is calculated using function `binom.test` from package `stats`. Replicates are systematically pooled in this plot.

## Value

a plot of class `ggplot`

## Note

When `style = "generic"`, the function calls the generic function `plot`

When `style = "ggplot"`, the function return an object of class `ggplot`, see function `ggplot`

## See Also

[binom.test](#)

## Examples

```
library(ggplot2)

# (1) Load the data
data(zinc)

# (2) Create an object of class 'survData'
zinc <- survData(zinc)

# (3) Plot dose-response
plotDoseResponse(zinc)
```

```
# (4) Plot dose-response with a generic style
plotDoseResponse(zinc, style = "generic")
```

---

plot_prior_post	<i>Generic method to plot priors and posteriors.</i>
-----------------	--

---

### Description

Plot priors and posteriors of a survFit object

### Usage

```
plot_prior_post(x, ...)
```

### Arguments

x	an object used to select a method plot_prior_post
...	Further arguments to be passed to generic methods

### Value

an object of class plot\_prior\_post

---

plot_prior_post.survFit	<i>Plot posteriors vs priors</i>
-------------------------	----------------------------------

---

### Description

Plot posteriors vs priors of a survFit object

### Usage

```
## S3 method for class 'survFit'
plot_prior_post(x, size_sample = 1000, EFSA_name = FALSE, ...)
```

### Arguments

x	an object of class survFit used to select a method plot_prior_post
size_sample	Size of the random generation of the distribution. Default is 1e3.
EFSA_name	If TRUE, replace the current terminology by the one used in the recent EFSA PPR Scientific Opinion (2018).
...	Further arguments to be passed to generic methods



**Value**

a plot of class ggplot

**References**

EFSA PPR Scientific Opinion (2018) *Scientific Opinion on the state of the art of Toxicokinetic/Toxicodynamic (TKTD) effect models for regulatory risk assessment of pesticides for aquatic organisms* <https://www.efsa.europa.eu/en/efsajournal/pub/5377>

---

 ppc

---

*Posterior predictive check plot*


---

**Description**

Plots posterior predictive check for reproFitTT, survFitTT, survFitTKTD, survFitCstExp and survFitVarExp objects.

This is the generic ppc S3 method for the reproFitTT class. It plots the predicted values with 95% credible intervals versus the observed values.

This is the generic ppc S3 method for the survFitCstExp class. It plots the predicted values along with 95% credible intervals versus the observed values for survFit objects.

This is the generic ppc S3 method for the survFitPredict\_Nsurv class. It plots the predicted values along with 95% credible intervals versus the observed values for survFitPredict\_Nsurv objects.

This is the generic ppc S3 method for the survFitTKTD class. It plots the predicted values along with 95% credible intervals versus the observed values for survFitTKTD objects.

This is the generic ppc S3 method for the survFitTT class. It plots the predicted values with 95 % credible intervals versus the observed values for survFitTT objects.

This is the generic ppc S3 method for the survFitVarExp class. It plots the predicted values along with 95% credible intervals versus the observed values for survFit objects.

**Usage**

```
ppc(x, ...)

## S3 method for class 'reproFitTT'
ppc(
  x,
  style = "ggplot",
  xlab = "Observed Cumul. Nbr. of offspring",
  ylab = "Predicted Cumul. Nbr. of offspring",
  main = NULL,
  ...
)

## S3 method for class 'survFitCstExp'
```

```

ppc(x, style = "ggplot", main = NULL, ...)

## S3 method for class 'survFitPredict_Nsurv'
ppc(
  x,
  xlab = "Observed nb of survivors",
  ylab = "Predicted nb of survivors",
  main = NULL,
  ...
)

## S3 method for class 'survFitTKTD'
ppc(x, style = "ggplot", main = NULL, ...)

## S3 method for class 'survFitTT'
ppc(x, style = "ggplot", main = NULL, ...)

## S3 method for class 'survFitVarExp'
ppc(
  x,
  xlab = "Observed nb of survivors",
  ylab = "Predicted nb of survivors",
  main = NULL,
  ...
)

```

### Arguments

x	An object of class <code>survFitVarExp</code>
...	Further arguments to be passed to generic methods
style	graphical backend, can be 'generic' or 'ggplot'
xlab	A label for the $X$ -axis, by default Observed nb of survivors.
ylab	A label for the $Y$ -axis, by default Predicted nb of survivors.
main	A main title for the plot.

### Details

Depending on the class of the object `x` see their links. for class `reproFitTT`: [ppc.reproFitTT](#) ; for class `survFitTT`: [ppc.survFitTT](#) ; for class `survFitTKTD`: [ppc.survFitTKTD](#) ; for class `survFitCstExp`: [ppc.survFitCstExp](#) and for class `survFitVarExp`: [ppc.survFitVarExp](#).

The coordinates of black points are the observed values of the cumulated number of reproduction outputs for a given concentration ( $X$ -scale) and the corresponding predicted values ( $Y$ -scale). 95% prediction intervals are added to each predicted value, colored in green if this interval contains the observed value and in red in the other case. As replicates are not pooled in this plot, overlapped points are shifted on the  $X$ -axis to help the visualization of replicates. The bisecting line ( $y = x$ ) is added to the plot in order to see if each prediction interval contains each observed value. As replicates are shifted on the  $X$ -axis, this line may be represented by steps.

The black points show the observed number of survivors (pooled replicates, on  $X$ -axis) against the corresponding predicted number ( $Y$ -axis). Predictions come along with 95% prediction intervals, which are depicted in green when they contain the observed value and in red otherwise. Samples with equal observed value are shifted on the  $X$ -axis. For that reason, the bisecting line ( $y = x$ ), is represented by steps when observed values are low. That way we ensure green intervals do intersect the bisecting line.

For `survFitPredict_Nsurv` object, PPC is based on times series simulated for each replicate. In addition, the black points show the observed number of survivors (on  $X$ -axis) against the corresponding predicted number ( $Y$ -axis). Predictions come along with 95% prediction intervals, which are depicted in green when they contain the observed value and in red otherwise.

The black points show the observed number of survivors (pooled replicates, on  $X$ -axis) against the corresponding predicted number ( $Y$ -axis). Predictions come along with 95% prediction intervals, which are depicted in green when they contain the observed value and in red otherwise. Samples with equal observed value are shifted on the  $X$ -axis. For that reason, the bisecting line ( $y = x$ ), is represented by steps when observed values are low. That way we ensure green intervals do intersect the bisecting line.

The coordinates of black points are the observed values of the number of survivors (pooled replicates) for a given concentration ( $X$ -axis) and the corresponding predicted values ( $Y$ -axis). 95% prediction intervals are added to each predicted value, colored in green if this interval contains the observed value and in red otherwise. The bisecting line ( $y = x$ ) is added to the plot in order to see if each prediction interval contains each observed value. As replicates are shifted on the  $x$ -axis, this line is represented by steps.

The black points show the observed number of survivors (on  $X$ -axis) against the corresponding predicted number ( $Y$ -axis). Predictions come along with 95% prediction intervals, which are depicted in green when they contain the observed value and in red otherwise.

## Value

a plot of class `ggplot`  
a plot of class `ggplot`  
a plot of class `ggplot`  
a plot of class `ggplot`  
a plot of class `ggplot`  
a plot of class `ggplot`  
a plot of class `ggplot`

## Examples

```
# (1) Load the data
data(cadmium1)

# (2) Create an object of class "reproData"
dataset <- reproData(cadmium1)
```

```
# (3) Run the reproFitTT function with the log-logistic gamma-Poisson model
out <- reproFitTT(dataset, stoc.part = "gammapoisson",
ecx = c(5, 10, 15, 20, 30, 50, 80), quiet = TRUE)
```

```
# (4) Plot observed versus predicted values
ppc(out)
```

```
# (1) Load the data
data(propiconazole)
```

```
# (2) Create an object of class "survData"
dataset <- survData(propiconazole)
```

```
# (3) Run the survFitTKTD function with the TKTD model ('SD' or 'IT')
out <- survFit(dataset, model_type = "SD")
```

```
# (4) Plot observed versus predicted values
ppc(out)
```

```
# (1) Load the data
data(propiconazole)
```

```
# (2) Create an object of class "survData"
dat <- survData(propiconazole)
```

```
# (3) Run the survFitTKTD function with the TKTD model ('SD' only)
out <- survFitTKTD(dat)
```

```
# (4) Plot observed versus predicted values
ppc(out)
```

```
# (1) Load the data
data(cadmium1)
```

```
# (2) Create an object of class "survData"
dat <- survData(cadmium1)
```

```
# (3) Run the survFitTT function with the log-logistic binomial model
out <- survFitTT(dat, lcx = c(5, 10, 15, 20, 30, 50, 80),
quiet = TRUE)
```

```
# (4) Plot observed versus predicted values
ppc(out)
```

```
# (1) Load the data
data(propiconazole_pulse_exposure)

# (2) Create an object of class "survData"
dat <- survData(propiconazole_pulse_exposure)

# (3) Run the survFitTKTD function with the TKTD model ('SD' or 'IT')
out <- survFit(dat, model_type = "SD")

# (4) Plot observed versus predicted values
ppc(out)
```

---

predict.survFit

*Predict method for survFit objects*

---

## Description

This is the generic predict S3 method for the survFit class. It provides simulation for "SD" or "IT" models under constant or time-variable exposure.

It provides the simulated number of survivors for "SD" or "IT" models under constant or time-variable exposure.

It provides the simulated number of survivors for "SD" or "IT" models under constant or time-variable exposure.

This is a method to replace function predict\_Nsurv used on survFit object when computing issues happen. predict\_nsurv\_ode uses the deSolve library to improve robustness. However, time to compute may be longer.

## Usage

```
## S3 method for class 'survFit'
predict(
  object,
  data_predict = NULL,
  spaghetti = FALSE,
  mcmc_size = NULL,
  hb_value = TRUE,
  ratio_no.NA = 0.95,
  hb_valueFORCED = NA,
  extend_time = 100,
  ...
)
```

```

predict_Nsurv(object, ...)

## S3 method for class 'survFit'
predict_Nsurv(
  object,
  data_predict = NULL,
  spaghetti = FALSE,
  mcmc_size = NULL,
  hb_value = TRUE,
  hb_valueFORCED = NA,
  extend_time = 100,
  ...
)

predict_Nsurv_ode(
  object,
  data_predict,
  spaghetti,
  mcmc_size,
  hb_value,
  hb_valueFORCED,
  extend_time,
  interpolate_length,
  interpolate_method,
  ...
)

```

### Arguments

object	An object of class <code>survFit</code> .
data_predict	A dataframe with three columns <code>time</code> , <code>conc</code> and <code>replicate</code> used for prediction. If <code>NULL</code> , prediction is based on <code>x</code> object of class <code>survFit</code> used for fitting.
spaghetti	If <code>TRUE</code> , return a set of survival curves using parameters drawn from the posterior distribution.
mcmc_size	Can be used to reduce the number of <code>mcmc</code> samples in order to speed up the computation. <code>mcmc_size</code> is the number of selected iterations for one chain. Default is 1000. If all MCMC is wanted, set argument to <code>NULL</code> .
hb_value	If <code>TRUE</code> , the background mortality <code>hb</code> is taken into account from the posterior. If <code>FALSE</code> , parameter <code>hb</code> is set to 0. The default is <code>TRUE</code> .
ratio_no.NA	A numeric between 0 and 1 standing for the proportion of non-NA values required to compute quantile. The default is 0.95.
hb_valueFORCED	If <code>hb_value</code> is <code>FALSE</code> , it fix <code>hb</code> .
extend_time	Length of time points interpolated with variable exposure profiles.
...	Further arguments to be passed to generic methods
interpolate_length	Length of the time sequence for which output is wanted.

interpolate\_method

The interpolation method for concentration. See package deSolve for details. Default is linear.

### Value

a list of data.frame with the quantiles of outputs in df\_quantiles or all the MCMC chains df\_spaghetti

an object of class predict\_Nsurv.

The function returns an object of class survFitPredict\_Nsurv, which is a list with the two following data.frame:

df\_quantile      A data.frame with 10 columns, time, conc, replicate, Nsurv (observed number of survivors) and other columns with median and 95% credible interval of the number of survivors computed with 2 different way refers as check and valid: Nsurv\_q50\_check, Nsurv\_qinf95\_check, Nsurv\_qsup95\_check, Nsurv\_q50\_valid, Nsurv\_qinf95\_valid, Nsurv\_qsup95\_valid. The \_check refers to the number of survivors at time  $t$  predicted using the observed number of survivors at time  $t - 1$ , while the \_valid refers to the number of survivors predicted at time  $t$  based on the predicted number of survivors at time  $t - 1$ .

df\_spaghetti      NULL if arguement spaghetti = FALSE. With spaghetti = TRUE, it returns a dataframe with all simulations based on MCMC parameters from a survFit object.

an object of class predict\_Nsurv\_ode.

### Examples

```
# (1) Load the survival data
data("propiconazole_pulse_exposure")

# (2) Create an object of class "survData"
dataset <- survData(propiconazole_pulse_exposure)

# (3) Run the survFit function
out <- survFit(dataset , model_type = "SD")

# (4) Create a new data table for prediction
data_4prediction <- data.frame(time = 1:10,
                               conc = c(0,5,30,30,0,0,5,30,15,0),
                               replicate= rep("predict", 10))

# (5) Predict on a new dataset
predict_out <- predict(object = out, data_predict = data_4prediction, spaghetti = TRUE)
```

```

# (1) Load the survival data
data("propiconazole_pulse_exposure")

# (2) Create an object of class "survData"
dataset <- survData(propiconazole_pulse_exposure)

# (3) Run the survFit function
out <- survFit(dataset , model_type = "SD")

# (4) Create a new data table for prediction
data_4prediction <- data.frame(time = 1:10,
                               conc = c(0,5,30,30,0,0,5,30,15,0),
                               replicate= rep("predict", 10),
                               Nsurv = c(20,20,17,16,15,15,15,14,13,12))

# (5) Predict Nsurv on a new data set
predict_out <- predict_Nsurv(object = out, data_predict = data_4prediction, spaghetti = TRUE)

```

---

predict\_Nsurv\_check    *Checking goodness-of-fit method for survFitPredict and survFitPredict\_Nsurv objects*

---

## Description

It returns measures of goodness-of-fit for predictions.

Provide various criteria for assessment of the model performance: (i) percentage of observation within the 95% credible interval of the Posterior Prediction Check (PPC), the Normalised Root Mean Square Error (NRMSE) and the Survival Probability Prediction Error (SPPE) as recommended by the recent Scientific Opinion from EFSA (2018).

## Usage

```

predict_Nsurv_check(object, ...)

## S3 method for class 'survFitPredict_Nsurv'
predict_Nsurv_check(object, ...)

```

## Arguments

object	an object of class survFitPredict_Nsurv
...	Further arguments to be passed to generic methods



**Value**

return a list of data.frame.

The function return a list with three items:

PPC	The criterion, in percent, compares the predicted median numbers of survivors associated to their uncertainty limits with the observed numbers of survivors. Based on experience, PPC resulting in less than 50% of the observations within the uncertainty limits indicate poor model performance. A fit of 100% may hide too large uncertainties of prediction (so covering all data).
PPC_global	percentage of PPC for the whole data set by gathering replicates.
NRMSE	The criterion, in percent, is based on the classical root-mean-square error (RMSE), used to aggregate the magnitudes of the errors in predictions for various time-points into a single measure of predictive power. In order to provide a criterion expressed as a percentage, NRMSE is the normalised RMSE by the mean of the observations.
NRMSE_global	NRMSE for the whole data set by gathering replicates.
SPPE	The SPPE indicator, in percent, is negative (between 0 and -100%) for an underestimation of effects, and positive (between 0 and 100) for an overestimation of effects. An SPPE value of 0 means an exact prediction of the observed survival probability at the end of the exposure profile.

@references EFSA PPR Scientific Opinion (2018) *Scientific Opinion on the state of the art of Toxicokinetic/Toxicodynamic (TKTD) effect models for regulatory risk assessment of pesticides for aquatic organisms* <https://www.efsa.europa.eu/en/efsajournal/pub/5377>

---

predict\_ode

*Predict method for survFit objects*

---

**Description**

This is a method to replace function predict used on survFit object when computing issues happen. predict\_ode uses the deSolve library to improve robustness. However, time to compute may be longer.

**Usage**

```
predict_ode(object, ...)
```

**Arguments**

object	an object used to select a method ppc
...	Further arguments to be passed to generic methods

**Value**

an object of class predict\_ode

---

predict\_ode.survFit *Predict method for survFit objects*

---

## Description

This is the generic predict S3 method for the survFit class. It provides predicted survival rate for "SD" or "IT" models under constant or time-variable exposure.

## Usage

```
## S3 method for class 'survFit'
predict_ode(
  object,
  data_predict = NULL,
  spaghetti = FALSE,
  mcmc_size = 1000,
  hb_value = TRUE,
  interpolate_length = 100,
  interpolate_method = "linear",
  hb_valueFORCED = NA,
  ...
)
```

## Arguments

object	An object of class survFit.
data_predict	A dataframe with three columns time, conc and replicate used for prediction. If NULL, prediction is based on x object of class survFit used for fitting.
spaghetti	If TRUE, return a set of survival curves using parameters drawn from the posterior distribution.
mcmc_size	Can be used to reduce the number of mcmc samples in order to speed up the computation. mcmc_size is the number of selected iterations for one chain. Default is 1000. If all MCMC is wanted, set argument to NULL.
hb_value	If TRUE, the background mortality hb is taken into account from the posterior. If FALSE, parameter hb is set to a fixed value. The default is TRUE.
interpolate_length	Length of the time sequence for which output is wanted.
interpolate_method	The interpolation method for concentration. See package deSolve for details. Default is linear.
hb_valueFORCED	If hb_value is FALSE, it fix hb.
...	Further arguments to be passed to generic methods

**Value**

a list of data.frame with the quantiles of outputs in df\_quantiles or all the MCMC chains df\_spaghetti

**Examples**

```
# (1) Load the survival data
data("propiconazole_pulse_exposure")

# (2) Create an object of class "survData"
dataset <- survData(propiconazole_pulse_exposure)

# (3) Run the survFit function
out <- survFit(dataset , model_type = "SD")

# (4) Create a new data table for prediction
data_4prediction <- data.frame(time = 1:10,
                              conc = c(0,5,30,30,0,0,5,30,15,0),
                              replicate= rep("predict", 10))

# (5) Predict on a new data set
predict_out <- predict_ode(object = out, data_predict = data_4prediction,
                          mcmc_size = 1000, spaghetti = TRUE)
```

---

print.msgTable	<i>Print msgTables objects</i>
----------------	--------------------------------

---

**Description**

Print in the REPL the msgTables

**Usage**

```
## S3 method for class 'msgTable'
print(x, ...)
```

**Arguments**

x                    an object of class msgTables  
...                   Further arguments to be passed to generic methods

**Value**

Print in the REPL the msgTables

---

print.reproFitTT	<i>Print of reproFitTT object</i>
------------------	-----------------------------------

---

### Description

This is the generic print S3 method for the reproFitTT class. It prints the underlying JAGS model and some information on the Bayesian inference procedure.

### Usage

```
## S3 method for class 'reproFitTT'  
print(x, ...)
```

### Arguments

x	An object of class reproFitTT
...	Further arguments to be passed to generic methods

### Value

print the model text and the Jags Computing information

### Examples

```
# (1) Load the data  
data(cadmium1)  
  
# (2) Create an object of class 'reproData'  
cadmium1 <- reproData(cadmium1)  
  
# (3) Run the reproFitTT function with the log-logistic  
# model  
out <- reproFitTT(cadmium1, ecx = c(5, 10, 15, 20, 30, 50, 80),  
quiet = TRUE)  
  
# (4) Print the reproFitTT object  
print(out)
```

---

print.survFitCstExp    *Print of survFit object*

---

### Description

This is the generic print S3 method for the survFitCstExp class. It prints the underlying JAGS model and some information on the Bayesian inference procedure.

### Usage

```
## S3 method for class 'survFitCstExp'  
print(x, ...)
```

### Arguments

x	An object of class survFitCstExp
...	Further arguments to be passed to generic methods.

### Value

print the model text and the Jags Computing information

### Examples

```
# (1) Load the data  
data(propiconazole)  
  
# (2) Create an object of class 'survData'  
dat <- survData(propiconazole)  
  
# (3) Run the survFit function with TKTD model 'SD' or 'IT'  
out <- survFit(dat, quiet = TRUE, model_type="SD")  
  
# (4) Print the survFit object  
print(out)
```

---

print.survFitTKTD      *Print of survFitTKTD object*

---

### Description

This is the generic print S3 method for the survFitTKTD class. It prints the underlying JAGS model and some information on the Bayesian inference procedure.

### Usage

```
## S3 method for class 'survFitTKTD'  
print(x, ...)
```

### Arguments

x	An object of class survFitTKTD
...	Further arguments to be passed to generic methods.

### Value

print the model text and the Jags Computing information

### Examples

```
# (1) Load the data  
data(propiconazole)  
  
# (2) Create an object of class 'survData'  
dat <- survData(propiconazole)  
  
# (3) Run the survFitTKTD function  
out <- survFitTKTD(dat, quiet = TRUE)  
  
# (4) Print the survFitTKTD object  
print(out)
```

---

print.survFitTT	<i>Print of survFitTT object</i>
-----------------	----------------------------------

---

### Description

This is the generic print S3 method for the survFitTT class. It prints the underlying JAGS model and some information on the Bayesian inference procedure.

### Usage

```
## S3 method for class 'survFitTT'  
print(x, ...)
```

### Arguments

x	An object of class survFitTT
...	Further arguments to be passed to generic methods

### Value

print the model text and the Jags Computing information

### Examples

```
# (1) Load the data  
data(cadmium1)  
  
# (2) Create an object of class 'survData'  
cadmium1 <- survData(cadmium1)  
  
# (3) Run the survFitTT function with the log-logistic  
# binomial model  
out <- survFitTT(cadmium1, lcx = c(5, 10, 15, 20, 30, 50, 80),  
                 quiet = TRUE)  
  
# (4) Print the survFitTT object  
print(out)
```

---

print.survFitVarExp    *Print of survFitVarExp object*

---

### Description

This is the generic print S3 method for the survFitVarExp class. It prints the underlying JAGS model and some information on the Bayesian inference procedure.

### Usage

```
## S3 method for class 'survFitVarExp'  
print(x, ...)
```

### Arguments

x	An object of class survFitVarExp
...	Further arguments to be passed to generic methods.

### Value

print the model text and the Jags Computing information

### Examples

```
# (1) Load the data  
data(propiconazole_pulse_exposure)  
  
# (2) Create a survData object  
dataset <- survData(propiconazole_pulse_exposure)  
  
# (3) Run the survFit function with TKTD model 'SD' or 'IT'  
out <- survFit(dataset, model_type="SD")  
  
# (4) Print the survFit object  
print(out)
```



---

priors\_distribution    *Density distribution of priors.*

---

**Description**

Return a data.frame with prior density distributions of parameters used in object.

**Usage**

```
priors_distribution(object, ...)
```

**Arguments**

object	An object used to select a method
...	Further arguments to be passed to generic methods

**Details**

When the object is of class survFit, see [priors\\_distribution.survFit](#)

**Value**

an object of class priors\_distribution

---

priors\_distribution.survFit  
*Density distribution of priors from a survFit object.*

---

**Description**

Return a data.frame with priors distribution of parameters used in object.

**Usage**

```
## S3 method for class 'survFit'
priors_distribution(object, size_sample = 1000, EFSA_name = FALSE, ...)
```

**Arguments**

object	An object of class survFit.
size_sample	Size of the random generation of the distribution. Default is 1e3.
EFSA_name	If TRUE, replace the current terminology by the one used in the recent EFSA PPR Scientific Opinion (2018).
...	Further arguments to be passed to generic methods.

**Value**

a data.frame with prio distribution.

**References**

EFSA PPR Scientific Opinion (2018) *Scientific Opinion on the state of the art of Toxicokinetic/Toxicodynamic (TKTD) effect models for regulatory risk assessment of pesticides for aquatic organisms* <https://www.efsa.europa.eu/en/efsajournal/pub/5377>.

---

priors_survData	<i>Create a list of scalars giving priors to use in Bayesian inference.</i>
-----------------	---

---

**Description**

Create a list of scalars giving priors to use in Bayesian inference.

**Usage**

```
priors_survData(x, model_type = NULL)
```

**Arguments**

x	An object of class survData
model_type	TKTD model type ('SD' or 'IT')

**Value**

A list for parameterization of priors for Bayesian inference with JAGS.

**Examples**

```
# (1) Load the data
data(cadmium1)

# (2) Create a survData object
dat <- survData(cadmium1)

# (3) Create priors for SD model_type
priors_survData(dat, model_type = "SD")

# (4) Create priors for IT model_type
priors_survData(dat, model_type = "IT")
```

---

propiconazole	<i>Survival data set for Gammarus pulex exposed to propiconazole during four days</i>
---------------	---

---

### Description

Survival data set of chronic laboratory toxicity tests with *Gammarus pulex* freshwater invertebrate exposed to eight concentrations of one fungicide (propiconazole) during four days. Eight concentrations were tested with two replicates of 10 organisms per concentration. Survival is monitored at five time points.

### Usage

```
data(propiconazole)
```

### Format

A dataframe with 75 observations on the following four variables:

`replicate` A vector of class `factor` with the replicate code (SC for the control and A1 to G2 for other profiles).

`conc` A vector of class `numeric` with propiconazole concentrations in  $\mu\text{mol.L}^{-1}$ .

`time` A vector of class `integer` with the time points (in days from the beginning of the experiment  $t = 0$ ).

`Nsurv` A vector of class `integer` with the number of alive individuals at each time point for each concentration and each replicate.

### References

Nyman, A.-M., Schirmer, K., Ashauer, R., (2012) Toxicokinetic-toxicodynamic modelling of survival of *Gammarus pulex* in multiple pulse exposures to propiconazole: model assumptions, calibration data requirements and predictive power, *Ecotoxicology*, (21), 1828-1840.

---

propiconazole_pulse_exposure	<i>Survival data set for Gammarus pulex exposed to propiconazole during 10 days with time-variable exposure concentration (non-standard pulsed toxicity experiments)</i>
------------------------------	--

---

### Description

Survival data set of laboratory toxicity tests with *Gammarus pulex* freshwater invertebrates exposed to several profiles of concentrations (time-variable concentration for each time series) of one fungicide (propiconazole) during 10 days.

**Usage**

```
data(propiconazole_pulse_exposure)
```

**Format**

A data frame with 74 observations on the following four variables:

replicate A vector of class factor with the replicate code (varControl, varA, varB and varC).

conc A vector of class numeric with propiconazole concentrations in  $\mu\text{mol.L}^{-1}$ .

time A vector of class integer with the time points (in days from the beginning of the experiment  $t = 0$ ).

Nsurv A vector of class integer with the number of alive individuals at each time point for each concentration and each replicate.

**References**

Nyman, A.-M., Schirmer, K., Ashauer, R., (2012) Toxicokinetic-toxicodynamic modelling of survival of *Gammarus pulex* in multiple pulse exposures to propiconazole: model assumptions, calibration data requirements and predictive power, *Ecotoxicology*, (21), 1828-1840.

---

propiconazole\_split     *Survival data set for Gammarus pulex exposed to propiconazole during four days*

---

**Description**

Survival data set of chronic laboratory toxicity tests with *Gammarus pulex* freshwater invertebrate exposed to eight concentrations of one fungicide (propiconazole) during four days. Eight concentrations were tested with two replicates of 10 organisms per concentration. Survival is monitored at five time points.

**Usage**

```
data(propiconazole_split)
```

**Format**

A dataframe with 75 observations on the following four variables:

replicate A vector of class factor with the replicate code (SC for the control and A1 to G2 for other profiles).

conc A vector of class numeric with propiconazole concentrations in  $\mu\text{mol.L}^{-1}$ .

time A vector of class integer with the time points (in days from the beginning of the experiment  $t = 0$ ).

Nsurv A vector of class integer with the number of alive individuals at each time point for each concentration and each replicate.

## References

Nyman, A.-M., Schirmer, K., Ashauer, R., (2012) Toxicokinetic-toxicodynamic modelling of survival of *Gammarus pulex* in multiple pulse exposures to propiconazole: model assumptions, calibration data requirements and predictive power, *Ecotoxicology*, (21), 1828-1840.

---

reproData

*Creates a dataset for reproduction toxicity analysis*

---

## Description

This function creates a reproData object from experimental data provided as a data.frame. The resulting object can then be used for plotting and model fitting. The reproData class is a sub-class of survData, meaning that all functions and method available for survival analysis can be used with reproData objects.

## Usage

```
reproData(x)
```

## Arguments

x a dataframe as expected by survData containing one additional Nrepro column of class integer with positive values only. This column should provide the number of offspring produced since the last observation.

## Details

The x argument contains the experimental data, and should have the same structure than the argument of survData, plus a single additional column providing the total number of offspring observed since the last time point. The function fails if x does not meet the expected requirements. Please run [reproDataCheck](#) to ensure x is well-formed.

Note that experimental data with time-variable exposure are not supported.

## Value

An object of class reproData.

## Examples

```
# (1) Load reproduction dataset
data(cadmium1)

# (2) Create an object of class "reproData"
dat <- reproData(cadmium1)
class(dat)
```

---

reproDataCheck	<i>Checks if an object can be used to perform reproduction toxicity data analysis</i>
----------------	---

---

### Description

The reproDataCheck function can be used to check if an object containing data from a reproduction toxicity assay meets the expectations of the function [reproData](#).

### Usage

```
reproDataCheck(data, diagnosis.plot = TRUE)
```

### Arguments

data            any object  
diagnosis.plot if TRUE, produces a diagnosis plot

### Details

Since in morse' reproduction data sets are a special case of survival data sets, reproDataCheck performs the same verifications than [survDataCheck](#) plus additional ones that are specific to reproduction data.

### Value

The function returns a data.frame similar to the one returned by [survDataCheck](#), except that it may contain the following additional error ids:

- NreproInteger: column Nrepro contains values of class other than integer
- Nrepro0T0: Nrepro is not 0 at time 0 for each concentration and each replicate
- Nsurvt0Nreprotp1P: at a given time  $T$ , the number of alive individuals is null and the number of collected offspring is not null for the same replicate and the same concentration at time  $T+1$

### Note

If an error of type dataframeExpected or missingColumn is detected, the function reproDataCheck is stopped. When no error is detected the reproDataCheck function returns an empty dataframe.

### See Also

[reproData](#)

**Examples**

```
# Run the check data function
data(copper)
reproDataCheck(copper)

# Now we insert an error in the data set, by setting a non-zero number of
# offspring at some time, although there is no surviving individual in the
# replicate from the previous time point.
copper[148, "Nrepro"] <- as.integer(1)
reproDataCheck(copper)
```

---

reproFitTT	<i>Fits a Bayesian concentration-effect model for target-time reproduction analysis</i>
------------	---

---

**Description**

This function estimates the parameters of a concentration-effect model for target-time reproduction analysis using Bayesian inference. In this model the endpoint is the cumulated number of reproduction outputs over time, with potential mortality all along the experiment.

**Usage**

```
reproFitTT(
  data,
  stoc.part = "bestfit",
  target.time = NULL,
  ecx = c(5, 10, 20, 50),
  n.chains = 3,
  quiet = FALSE
)
```

**Arguments**

<code>data</code>	an object of class <code>reproData</code>
<code>stoc.part</code>	stochastic part of the model. Possible values are "bestfit", "poisson" and "gammapoisson"
<code>target.time</code>	defines the target time point at which to analyse the repro data. By default the last time point
<code>ecx</code>	desired values of $x$ (in percent) for which to compute $EC_x$
<code>n.chains</code>	number of MCMC chains. The minimum required number of chains is 2
<code>quiet</code>	if TRUE, does not print messages and progress bars from JAGS

## Details

Because some individuals may die during the observation period, the reproduction rate alone is not sufficient to account for the observed number of offspring at a given time point. In addition, we need the time individuals have stayed alive during this observation period. The `reproFitTT` function estimates the number of individual-days in an experiment between its start and the target time. This covariable is then used to estimate a relation between the chemical compound concentration and the reproduction rate *per individual-day*.

The `reproFitTT` function fits two models, one where inter-individual variability is neglected ("Poisson" model) and one where it is taken into account ("gamma-Poisson" model). When setting `stoc.part` to "bestfit", a model comparison procedure is used to choose between both. More details are presented in the vignette accompanying the package.

## Value

The function returns an object of class `reproFitTT` which is a list of the following objects:

<code>DIC</code>	DIC value of the selected model
<code>estim.ECx</code>	a table of the estimated 5, 10, 20 and 50 % effective concentrations (by default) and their 95 % credible intervals
<code>estim.par</code>	a table of the estimated parameters as medians and 95 % credible intervals
<code>mcmc</code>	an object of class <code>mcmc.list</code> with the posterior distribution
<code>model</code>	a JAGS model object
<code>warnings</code>	a <code>data.frame</code> with warning messages
<code>model.label</code>	a character string, "P" if the Poisson model is used, "GP" if the gamma-Poisson is used
<code>parameters</code>	a list of the parameter names used in the model
<code>n.chains</code>	an integer value corresponding to the number of chains used for the MCMC computation
<code>n.iter</code>	a list of two indices indicating the beginning and the end of monitored iterations
<code>n.thin</code>	a numerical value corresponding to the thinning interval
<code>jags.data</code>	a list of the data passed to the jags model
<code>transformed.data</code>	the <code>survData</code> object passed to the function
<code>dataTT</code>	the dataset with which the parameters are estimated

## Examples

```
# (1) Load the data
data(cadmium1)

# (2) Create an object of class "reproData"
dataset <- reproData(cadmium1)
```



```
# (3) Run the reproFitTT function with the log-logistic gamma-Poisson model
out <- reproFitTT(dataset, stoc.part = "gammapoisson",
                  ecx = c(5, 10, 15, 20, 30, 50, 80), quiet = TRUE)
```

---

summary.reproData      *Summary of reproData object*

---

### Description

This is the generic summary S3 method for the reproData class. It provides information about the structure of the data set and the experimental design.

### Usage

```
## S3 method for class 'reproData'
summary(object, quiet = FALSE, ...)
```

### Arguments

object	an object of class reproData
quiet	if TRUE, does not print
...	Further arguments to be passed to generic methods

### Value

The function returns a list with the same information than [summary.survDataCstExp](#) plus an additional one:

NboffTimeConc    nb of offspring for all concentrations and time points

### Examples

```
# (1) Load the data
data(cadmium1)

# (2) Create a reproData object
cadmium1 <- reproData(cadmium1)

# (3) Summarize the data set
summary(cadmium1)
```

---

summary.reproFitTT      *Summary of reproFitTT object*

---

### Description

This is the generic summary S3 method for the reproFitTT class. It shows the quantiles of priors and posteriors on parameters and the quantiles of the posterior on the ECx estimates.

### Usage

```
## S3 method for class 'reproFitTT'  
summary(object, quiet = FALSE, ...)
```

### Arguments

object	an object of class reproFitTT
quiet	when TRUE, does not print
...	Further arguments to be passed to generic methods

### Value

The function returns a list with the following information:

Qpriors	quantiles of the model priors
Qposteriors	quantiles of the model posteriors
QECx	quantiles of ECx estimates

### Examples

```
# (1) Load the data  
data(cadmium1)  
  
# (2) Create a reproData object  
cadmium1 <- reproData(cadmium1)  
  
# (3) Run the reproFitTT function with the log-logistic  
# model  
out <- reproFitTT(cadmium1, ecx = c(5, 10, 15, 20, 30, 50, 80),  
quiet = TRUE)  
  
# (4) summarize the reproFitTT object  
summary(out)
```

---

`summary.survDataCstExp`*Summary of survDataCstExp object*

---

## Description

The generic summary S3 method for the survDataCstExp class provides information about the structure of the data set and the experimental design.

## Usage

```
## S3 method for class 'survDataCstExp'  
summary(object, quiet = FALSE, ...)
```

## Arguments

<code>object</code>	an object of class survDataCstExp
<code>quiet</code>	when TRUE, does not print
<code>...</code>	Further arguments to be passed to generic methods

## Value

The function returns a list with the following information:

`NbrepTimeConc` nb of replicates for all concentrations and time points

`NbsurvTimeConc` nb of survivors. for all concentrations and time points

## Examples

```
# (1) Load the data  
data(cadmium1)  
  
# (2) Create a survDataCstExp object  
dat <- survData(cadmium1)  
  
# (3) Summarize the data set  
summary(dat)
```

---

`summary.survDataVarExp`*Summary of survDataVarExp object*

---

## Description

The generic summary S3 method for the survDataVarExp class provides information about the structure of the data set and the experimental design.

## Usage

```
## S3 method for class 'survDataVarExp'  
summary(object, quiet = FALSE, ...)
```

## Arguments

<code>object</code>	an object of class survDataVarExp
<code>quiet</code>	when TRUE, does not print
<code>...</code>	Further arguments to be passed to generic methods

## Value

The function returns a list with the following information:

<code>OccRepTime</code>	Occurence of replicates for all time points
<code>NbsurvTimeRep</code>	nb of survivors. for all replicates and time points
<code>ConcTimeRep</code>	Concentration for all replicates and time points

## Examples

```
# (1) Load the data  
data(propiconazole_pulse_exposure)  
  
# (2) Create a survDataVarExp object  
out <- survData(propiconazole_pulse_exposure)  
  
# (3) Summarize the data set  
summary(out)
```

---

summary.survFit      *Summary of survFit object*

---

### Description

This is the generic summary S3 method for the survFit class. It shows the quantiles of priors and posteriors on parameters.

### Usage

```
## S3 method for class 'survFit'  
summary(object, quiet = FALSE, EFSA_name = FALSE, ...)
```

### Arguments

object	An object of class survFit.
quiet	When TRUE, does not print.
EFSA_name	If TRUE, the current terminology by the one used in the recent EFSA PPR Scientific Opinion (2018).
...	Further arguments to be passed to generic methods.

### Value

The function returns a list with the following information:

Qpriors	quantiles of the model priors
Qposteriors	quantiles of the model posteriors

### References

EFSA PPR Scientific Opinion (2018) *Scientific Opinion on the state of the art of Toxicokinetic/Toxicodynamic (TKTD) effect models for regulatory risk assessment of pesticides for aquatic organisms* <https://www.efsa.europa.eu/en/efsajournal/pub/5377>.

### Examples

```
# (1) Load the data  
data(propiconazole)  
  
# (2) Create a survData object  
dat <- survData(propiconazole)  
  
# (3) Run the survFit function  
out <- survFit(dat, model_type = "SD")  
  
# (4) summarize the survFit object  
summary(out)
```

---

summary.survFitTKTD *Summary of survFitTKTD object*

---

### Description

This is the generic summary S3 method for the survFitTKTD class. It shows the quantiles of priors and posteriors on parameters.

### Usage

```
## S3 method for class 'survFitTKTD'  
summary(object, quiet = FALSE, ...)
```

### Arguments

object	an object of class survFitTKTD
quiet	when TRUE, does not print
...	Further arguments to be passed to generic methods.

### Value

The function returns a list with the following information:

Qpriors	quantiles of the model priors
Qposteriors	quantiles of the model posteriors

### Examples

```
# (1) Load the data  
data(propiconazole)  
  
# (2) Create a survData object  
dat <- survData(propiconazole)  
  
# (3) Run the survFitTKTD function  
out <- survFitTKTD(dat)  
  
# (4) summarize the survFitTKTD object  
summary(out)
```

---

summary.survFitTT      *Summary of survFitTT object*

---

### Description

This is the generic summary S3 method for the survFitTT class. It shows the quantiles of priors and posteriors on parameters and the quantiles of the posteriors on the LCx estimates.

### Usage

```
## S3 method for class 'survFitTT'  
summary(object, quiet = FALSE, ...)
```

### Arguments

object	an object of class survFitTT
quiet	when TRUE, does not print
...	Further arguments to be passed to generic methods

### Value

The function returns a list with the following information:

Qpriors	quantiles of the model priors
Qposteriors	quantiles of the model posteriors
QLCx	quantiles of LCx estimates

### Examples

```
# (1) Load the data  
data(cadmium1)  
  
# (2) Create a survData object  
cadmium1 <- survData(cadmium1)  
  
# (3) Run the survFitTT function with the log-logistic  
# binomial model  
out <- survFitTT(cadmium1, lcx = c(5, 10, 15, 20, 30, 50, 80),  
                  quiet = TRUE)  
  
# (4) summarize the survFitTT object  
summary(out)
```

---

survData	<i>Creates a data set for survival analysis</i>
----------	---

---

## Description

This function creates a `survData` object from experimental data provided as a `data.frame`. The resulting object can then be used for plotting and model fitting. It can also be used to generate *individual-time* estimates.

The `survDataCheck` function can be used to check if an object containing survival data is formatted according to the expectations of the `survData` function.

## Usage

```
survData(x)
```

```
survDataCheck(data, diagnosis.plot = FALSE)
```

## Arguments

<code>x</code>	a <code>data.frame</code> containing the following four columns: <ul style="list-style-type: none"><li>• <code>replicate</code>: a vector of any class numeric, character or factor for replicate identification. A given replicate value should identify the same group of individuals followed in time</li><li>• <code>conc</code>: a vector of class numeric with tested concentrations (positive values, may contain NAs)</li><li>• <code>time</code>: a vector of class integer with time points, minimal value must be 0</li><li>• <code>Nsurv</code>: a vector of class integer providing the number of alive individuals at each time point for each concentration and each replicate (may contain NAs)</li></ul>
<code>data</code>	any object
<code>diagnosis.plot</code>	if TRUE, the function may produce diagnosis plots

## Details

Survival data sets can be under either constant or time-variable exposure profile. The resulting object, in addition to its `survData` class, inherits the class `survDataCstExp` or `survDataVarExp` respectively.

The `x` argument describes experimental results from a survival toxicity test. Each line of the `data.frame` corresponds to one experimental measurement, that is a number of alive individuals at a given concentration at a given time point and in a given replicate. Note that either the concentration or the number of alive individuals may be missing. The data set is inferred to be under constant exposure if the concentration is constant for each replicate and systematically available. The function `survData` fails if `x` does not meet the expected requirements. Please run [survDataCheck](#) to ensure `x` is well-formed.



**Value**

A dataframe of class `survData` and column replicate as factor.

The function returns a dataframe of class `msgTable` and `data.frame` with two columns: `id` and `msg` of character strings. When no error is detected the object is empty. Here is the list of possible error ids with their meaning:

<code>dataframeExpected</code>	an object of class <code>data.frame</code> is expected
<code>missingColumn</code>	at least one expected column heading is missing
<code>firstTime0</code>	the first time point for some (concentration, replicate) couples is not 0
<code>concNumeric</code>	column <code>conc</code> contains a value of class other than numeric
<code>timeNumeric</code>	column <code>time</code> contains a value of class other than numeric
<code>NsurvInteger</code>	column <code>Nsurv</code> contains a value of class other than integer
<code>tablePositive</code>	some data are negative
<code>Nsurv0T0</code>	<code>Nsurv</code> is 0 at time 0 for some (concentration, replicate)
<code>duplicateID</code>	there are two identical (replicate, time) couples
<code>NsurvIncrease</code>	<code>Nsurv</code> increases at some time point of some (concentration, replicate)
<code>maxTimeDiffer</code>	maximum time for concentration is lower than maximum time for survival

**Note**

If an error of type `dataframeExpected` or `missingColumn` is detected, the function `survDataCheck` is stopped before looking for other errors.

**See Also**

[survDataCheck](#)

[survData](#)

**Examples**

```
# (1) Load the survival data set
data(zinc)

# (2) Create an objet of class 'survData'
dat <- survData(zinc)
class(dat)

# Run the check data function
data(zinc)
survDataCheck(zinc)

# Now we insert an error in the dataset, by artificially increasing the
# number of survivors at a given time point, in such a way that the number
# of individuals increases in the corresponding replicate
zinc[25, "Nsurv"] <- as.integer(20)
survDataCheck(zinc, diagnosis.plot = TRUE)
```

---

survData_join	<i>Joins a concentration with a survival data set into an argument for 'survData' when the concentration varies over time</i>
---------------	---

---

## Description

This function joins two data sets, one for exposure measurements, the other for survival measurements, into a single dataframe that can be used with the survData function.

## Usage

```
survData_join(x, y)
```

## Arguments

- |   |   |
|---|---|
| x | a data.frame containing the following three columns: <ul style="list-style-type: none"><li>• replicate: a vector of class integer or factor for replicate identification</li><li>• time: a vector of class integer with time points, min value must be 0</li><li>• Nsurv: a vector of class integer providing the number of alive individuals at some or all time points for each replicate</li></ul> |
| y | a data.frame containing the following three columns: <ul style="list-style-type: none"><li>• replicate: a vector of class integer or factor for replicate identification</li><li>• time: a vector of class integer with time points, min value must be 0</li><li>• conc: a vector of class numeric providing the concentration at some or all time points for each replicate</li></ul>                |

## Value

a dataframe suitable for 'survData'

## Examples

```
# (1) Load the two survival data sets
data(propiconazole_pulse_exposure)
exposure <- propiconazole_pulse_exposure[,c("replicate", "time", "conc")]
survival <- propiconazole_pulse_exposure[,c("replicate", "time", "Nsurv")]

# (2) Create an objet of class 'survData'
dat_join <- survData(survData_join(exposure, survival))
class(dat_join)
```

---

`survFit`*Fits a TKTD model for survival analysis using Bayesian inference*

---

**Description**

This function estimates the parameters of a TKTD model ('SD' or 'IT') for survival analysis using Bayesian inference. In this model, the survival rate of individuals is modeled as a function of the chemical compound concentration with a mechanistic description of the effects on survival over time.

**Usage**

```
survFit(  
  data,  
  model_type,  
  quiet,  
  n.chains,  
  n.adapt,  
  n.iter,  
  n.warmup,  
  thin.interval,  
  limit.sampling,  
  dic.compute,  
  dic.type,  
  hb_value,  
  hb_valueFIXED,  
  ...  
)  
  
## S3 method for class 'survDataCstExp'  
survFit(  
  data,  
  model_type = NULL,  
  quiet = FALSE,  
  n.chains = 3,  
  n.adapt = 3000,  
  n.iter = NULL,  
  n.warmup = NULL,  
  thin.interval = NULL,  
  limit.sampling = TRUE,  
  dic.compute = FALSE,  
  dic.type = "pD",  
  hb_value = TRUE,  
  hb_valueFIXED = NA,  
  ...  
)
```

```
## S3 method for class 'survDataVarExp'
survFit(
  data,
  model_type = NULL,
  quiet = FALSE,
  n.chains = 3,
  n.adapt = 1000,
  n.iter = NULL,
  n.warmup = NULL,
  thin.interval = NULL,
  limit.sampling = TRUE,
  dic.compute = FALSE,
  dic.type = "pD",
  hb_value = TRUE,
  hb_valueFIXED = NA,
  extend_time = 100,
  ...
)
```

### Arguments

<code>data</code>	An object of class <code>survDataCstExp</code> or <code>survDataVarExp</code> .
<code>model_type</code>	Can be "SD" or "IT" to choose between "Stochastic Death" or "Individual Tolerance" models (resp.). See the modeling vignette for details.
<code>quiet</code>	If FALSE, prints logs and progress bar from JAGS.
<code>n.chains</code>	A positive integer specifying the number of MCMC chains. The minimum required number of chains is 2.
<code>n.adapt</code>	A positive integer specifying the number of iterations for adaptation. If <code>n.adapt = 0</code> then no adaptation takes place.
<code>n.iter</code>	A positive integer specifying the number of iterations to monitor for each chain.
<code>n.warmup</code>	A positive integer specifying the number of warmup (aka burnin) iterations per chain.
<code>thin.interval</code>	A positive integer specifying the period to monitor.
<code>limit.sampling</code>	if FALSE (default is TRUE), there is no limit to the number of iterations in MCMC imposed by the <code>raftery.diag</code> test.
<code>dic.compute</code>	if TRUE (default is FALSE), it generates penalized deviance samples to compute the Deviance Information Criterion (DIC) with the <code>rjags</code> package
<code>dic.type</code>	type of penalty to use. A string identifying the type of penalty: <code>pD</code> or <code>popt</code> (see function <a href="#">dic.samples</a> )
<code>hb_value</code>	If TRUE, the background mortality <code>hb</code> is taken into account. If FALSE, parameter <code>hb</code> is set to 0. The default is TRUE.
<code>hb_valueFIXED</code>	If <code>hb_value</code> is FALSE, then <code>hb_valueFIXED</code> is the value to fix <code>hb</code> . If <code>hb_value</code> is FALSE and <code>hb_valueFIXED</code> is NA, then <code>hb</code> is fixed to 0.
<code>...</code>	Further arguments to be passed to generic methods
<code>extend_time</code>	Number of for each replicate used for linear interpolation (comprise between time to compute and fitting accuracy)

## Details

The function `survFit` returns the parameter estimates of Toxicokinetic-toxicodynamic (TKTD) models SD for 'Stochastic Death' or IT for 'Individual Tolerance'. TKTD models, and particularly the General Unified Threshold model of Survival (GUTS), provide a consistent process-based framework to analyse both time and concentration dependent datasets. In GUTS-SD, all organisms are assumed to have the same internal concentration threshold (denoted  $z$ ), and, once exceeded, the instantaneous probability to die increases linearly with the internal concentration. In GUTS-IT, the threshold concentration is distributed among all the organisms, and once exceeded in one individual, this individual dies immediately.

When class of object is `survDataCstExp`, see [survFit.survDataCstExp](#); and for a `survDataVarExp`, see [survFit.survDataVarExp](#).

## Value

an object of class `survFit`

The function returns an object of class `survFitCstExp`, which is a list with the following information:

<code>estim.par</code>	a table of the estimated parameters as medians and 95% credible intervals
<code>mcmc</code>	an object of class <code>mcmc.list</code> with the posterior distribution
<code>model</code>	a JAGS model object
<code>dic</code>	return the Deviance Information Criterion (DIC) if <code>dic.compute</code> is TRUE
<code>warnings</code>	a table with warning messages
<code>parameters</code>	a list of parameter names used in the model
<code>n.chains</code>	an integer value corresponding to the number of chains used for the MCMC computation
<code>mcmcInfo</code>	a table with the number of iterations, chains, adaptation, warmup and the thinning interval.
<code>jags.data</code>	a list of the data passed to the JAGS model
<code>model_type</code>	the type of TKTD model used: SD or IT

The function returns an object of class `survFitVarExp`, which is a list with the following information:

<code>estim.par</code>	a table of the estimated parameters as medians and 95% credible intervals
<code>mcmc</code>	an object of class <code>mcmc.list</code> with the posterior distribution
<code>model</code>	a JAGS model object
<code>dic</code>	return the Deviance Information Criterion (DIC) if <code>dic.compute</code> is TRUE
<code>warnings</code>	a table with warning messages
<code>parameters</code>	a list of parameter names used in the model
<code>n.chains</code>	an integer value corresponding to the number of chains used for the MCMC computation
<code>mcmcInfo</code>	a table with the number of iterations, chains, adaptation, warmup and the thinning interval
<code>jags.data</code>	a list of the data passed to the JAGS model
<code>model_type</code>	the type of TKTD model used: SD or IT

## References

Jager, T., Albert, C., Preuss, T. G. and Ashauer, R. (2011) General unified threshold model of survival-a toxicokinetic-toxicodynamic framework for ecotoxicology, *Environmental Science and Technology*, 45, 2529-2540. 303-314.

## Examples

```
# Example with time-variable exposure profile#
# (1) Load the survival data
data(propiconazole)
# (2) Create an object of class "survData"
dataset <- survData(propiconazole)

# (3) Run the survFit function with TKTD model 'SD' or 'IT'
out <- survFit(dataset , model_type = "SD")
# (4) Summarize look the estimated parameters
summary(out)
# (5) Plot the fitted curve
plot(out, adddata = TRUE)
# (6) Plot the fitted curve with ggplot style and CI as spaghetti
plot(out, spaghetti = TRUE , adddata = TRUE)

# Example with time-variable exposure profile
# (1) Load the survival data
data("propiconazole_pulse_exposure")
# (2) Create an object of class "survData"
dataset <- survData(propiconazole_pulse_exposure)

# (3) Run the survFit function with TKTD model 'SD' or 'IT'
out <- survFit(dataset , model_type = "SD")
# (4) Summarize look the estimated parameters
summary(out)
# (5) Plot the fitted curve
plot(out, adddata = FALSE)
# (6) Plot the fitted curve with ggplot style and CI as spaghetti
plot(out, spaghetti = TRUE)
```

## Description

This function estimates the parameters of a TKTD model for survival analysis using Bayesian inference. In this model, the survival rate of individuals is modeled as a function of the chemical compound concentration with a mechanistic description of the effects on survival over time.

## Usage

```
survFitTKTD(data, n.chains = 3, quiet = FALSE)
```

## Arguments

data	An object of class <code>survData</code> .
n.chains	Number of MCMC chains. The minimum required number of chains is 2.
quiet	If FALSE, prints logs and progress bar from JAGS.

## Value

The function returns an object of class `survFitTKTD`, which is a list with the following information:

estim.par	a table of the estimated parameters as medians and 95% credible intervals
mcmc	an object of class <code>mcmc.list</code> with the posterior distribution
warnings	a table with warning messages
model	a JAGS model object
parameters	a list of parameter names used in the model
n.chains	an integer value corresponding to the number of chains used for the MCMC computation
n.iter	a list of two indices indicating the beginning and the end of monitored iterations
n.thin	a numerical value corresponding to the thinning interval
jags.data	a list of data passed to the JAGS model

## References

Delignette-Muller ML, Ruiz P and Veber P (2017). *Robust fit of toxicokinetic-toxicodynamic models using prior knowledge contained in the design of survival toxicity tests*.

Bedaux, J., Kooijman, SALM (1994) Statistical analysis of toxicity tests, based on hazard modeling, *Environmental and Ecological Statistics*, 1, 303-314.

## Examples

```
# (1) Load the survival data
data(propiconazole)

# (2) Create an object of class "survData"
dataset <- survData(propiconazole)
```

```

# (3) Run the survFitTKTD function
out <- survFitTKTD(dataset)

# (4) Summarize look the estimated parameters
summary(out)

# (5) Plot the fitted curve
plot(out, adddata = TRUE)

# (6) Plot the fitted curve with ggplot style and CI as spaghetti
plot(out, spaghetti = TRUE , adddata = TRUE,
      style = "ggplot")

```

---

survFitTT	<i>Fits a Bayesian concentration-response model for target-time survival analysis</i>
-----------	---

---

### Description

Fits a Bayesian concentration-response model for target-time survival analysis

### Usage

```
survFitTT(data, ...)
```

### Arguments

data	an object used to select a method 'survFitTT'
...	Further arguments to be passed to generic methods

### Value

an object of class survFitTT

---

survFitTT.survDataCstExp	<i>Fits a Bayesian concentration-response model for target-time survival analysis</i>
--------------------------	---

---

### Description

This function estimates the parameters of an concentration-response model for target-time survival analysis using Bayesian inference. In this model, the survival rate of individuals at a given time point (called target time) is modeled as a function of the chemical compound concentration. The actual number of surviving individuals is then modeled as a stochastic function of the survival rate. Details of the model are presented in the vignette accompanying the package.



**Usage**

```
## S3 method for class 'survDataCstExp'
survFitTT(
  data,
  target.time = NULL,
  lcx = c(5, 10, 20, 50),
  n.chains = 3,
  quiet = FALSE,
  ...
)
```

**Arguments**

data	an object of class <code>survData</code>
target.time	the chosen endpoint to evaluate the effect of the chemical compound concentration, by default the last time point available for all concentrations
lcx	desired values of $x$ (in percent) for which to compute $LC_x$ .
n.chains	number of MCMC chains, the minimum required number of chains is 2
quiet	if TRUE, does not print messages and progress bars from JAGS
...	Further arguments to be passed to generic methods

**Details**

The function returns parameter estimates of the concentration-response model and estimates of the so-called  $LC_x$ , that is the concentration of chemical compound required to get an  $(1 - x/100)$  survival rate.

**Value**

The function returns an object of class `survFitTT`, which is a list with the following information:

estim.LCx	a table of the estimated $LC_x$ along with their 95% credible intervals
estim.par	a table of the estimated parameters (medians) and 95% credible intervals
det.part	the name of the deterministic part of the used model
mcmc	an object of class <code>mcmc.list</code> with the posterior distribution
warnings	a table with warning messages
model	a JAGS model object
parameters	a list of parameter names used in the model
n.chains	an integer value corresponding to the number of chains used for the MCMC computation
n.iter	a list of two indices indicating the beginning and the end of monitored iterations
n.thin	a numerical value corresponding to the thinning interval
jags.data	a list of the data passed to the JAGS model
transformed.data	the <code>survData</code> object passed to the function
dataTT	the dataset with which the parameters are estimated

**Examples**

```
# (1) Load the data
data(cadmium1)

# (2) Create an object of class "survData"
dat <- survData(cadmium1)

# (3) Run the survFitTT function with the log-logistic
#       binomial model
out <- survFitTT(dat, lcx = c(5, 10, 15, 20, 30, 50, 80),
                 quiet = TRUE)
```

---

zinc	<i>Reproduction and survival data sets for Daphnia magna exposed to zinc during 21 days</i>
------	---

---

**Description**

Reproduction and survival data sets of a chronic laboratory toxicity tests with *Daphnia magna* freshwater invertebrate exposed to four concentrations of zinc during 21 days. Four concentrations were tested with three replicates per concentration. Each replicate contained 20 organisms. Reproduction and survival were monitored at 15 time points.

**Usage**

```
data(zinc)
```

**Format**

A data frame with 180 observations on the following five variables:

`replicate` A vector of class `numeric` with the replicate code (1 to 12).

`conc` A vector of class `numeric` with zinc concentrations in  $mg.L^{-1}$ .

`time` A vector of class `integer` with the time points (in days from the beginning of the experiment  $t = 0$ ).

`Nsurv` A vector of class `integer` with the number of alive individuals at each time point for each concentration and each replicate.

`Nrepro` A vector of class `integer` with the number of offspring at each time point for each concentration and each replicate.

**References**

Billoir, E., Delignette-Muller, M.L., Pery, A.R.R. and Charles S. (2008) A Bayesian Approach to Analyzing Ecotoxicological Data, *Environmental Science & Technology*, 42 (23), 8978-8984.

# Index

- \* **check**
  - reproDataCheck, 62
- \* **data**
  - cadmium1, 5
  - cadmium2, 6
  - chlordan, 7
  - copper, 8
  - dichromate, 8
  - FOCUSprofile, 9
  - propiconazole, 59
  - propiconazole\_pulse\_exposure, 59
  - propiconazole\_split, 60
  - zinc, 82
- \* **estimation**
  - reproFitTT, 63
  - survFit, 75
  - survFitTKTD, 78
  - survFitTT.survDataCstExp, 80
- \* **plot**
  - plot.LCx, 18
  - plot.MFx, 19
  - plot.reproData, 21
  - plot.reproFitTT, 22
  - plot.survDataCstExp, 24
  - plot.survDataVarExp, 25
  - plot.survFitCstExp, 26
  - plot.survFitPredict, 27
  - plot.survFitPredict\_Nsurv, 29
  - plot.survFitTKTD, 30
  - plot.survFitTT, 32
  - plot.survFitVarExp, 34
  - plotDoseResponse.reproData, 36
  - plotDoseResponse.survDataCstExp, 38
- \* **print**
  - print.reproFitTT, 52
  - print.survFitCstExp, 53
  - print.survFitTKTD, 54
  - print.survFitTT, 55
  - print.survFitVarExp, 56
- \* **set**
  - cadmium1, 5
  - cadmium2, 6
  - chlordan, 7
  - copper, 8
  - dichromate, 8
  - FOCUSprofile, 9
  - propiconazole, 59
  - propiconazole\_pulse\_exposure, 59
  - propiconazole\_split, 60
  - zinc, 82
- \* **summary**
  - summary.reproData, 65
  - summary.reproFitTT, 66
  - summary.survDataCstExp, 67
  - summary.survDataVarExp, 68
  - summary.survFit, 69
  - summary.survFitTKTD, 70
  - summary.survFitTT, 71
- \* **transformation**
  - reproData, 61
  - survData, 72
- binom.test, 39
- cadmium1, 5
- cadmium2, 6
- chlordan, 7
- copper, 8
- dic.samples, 76
- dichromate, 8
- FOCUSprofile, 9
- ggplot, 22, 23, 25, 26, 33, 38, 39
- ggplot2, 5
- is\_exposure\_constant, 10

LCx, 10  
LCx.survFit, 11

MFx, 12  
MFx.survFit, 14  
MFx\_ode (MFx), 12  
modelData, 16  
modelData.survDataCstExp, 17  
modelData.survDataVarExp, 17  
morse (morse-package), 3  
morse-package, 3

plot, 22, 23, 38, 39  
plot.LCx, 18  
plot.MFx, 19  
plot.reproData, 21  
plot.reproFitTT, 22  
plot.survDataCstExp, 24  
plot.survDataVarExp, 25  
plot.survFitCstExp, 26  
plot.survFitPredict, 27  
plot.survFitPredict\_Nsurv, 29  
plot.survFitTKTD, 30  
plot.survFitTT, 32  
plot.survFitVarExp, 34  
plot\_prior\_post, 40  
plot\_prior\_post.survFit, 40  
plotDoseResponse, 36  
plotDoseResponse.reproData, 36  
plotDoseResponse.survDataCstExp, 38  
pois.exact, 37, 38  
ppc, 41  
ppc.reproFitTT, 42  
ppc.survFitCstExp, 42  
ppc.survFitTKTD, 42  
ppc.survFitTT, 42  
ppc.survFitVarExp, 42  
predict.survFit, 45  
predict\_Nsurv (predict.survFit), 45  
predict\_Nsurv\_check, 48  
predict\_Nsurv\_ode (predict.survFit), 45  
predict\_ode, 49  
predict\_ode.survFit, 50  
print.msgTable, 51  
print.reproFitTT, 52  
print.survFitCstExp, 53  
print.survFitTKTD, 54  
print.survFitTT, 55  
print.survFitVarExp, 56

priors\_distribution, 57  
priors\_distribution.survFit, 57, 57  
priors\_survData, 58  
propiconazole, 59  
propiconazole\_pulse\_exposure, 59  
propiconazole\_split, 60

reproData, 61, 62  
reproDataCheck, 61, 62  
reproFitTT, 63  
rjags, 5

summary.reproData, 65  
summary.reproFitTT, 66  
summary.survDataCstExp, 65, 67  
summary.survDataVarExp, 68  
summary.survFit, 69  
summary.survFitTKTD, 70  
summary.survFitTT, 71  
survData, 72, 73  
survData\_join, 74  
survDataCheck, 62, 72, 73  
survDataCheck (survData), 72  
survFit, 75  
survFit.survDataCstExp, 77  
survFit.survDataVarExp, 77  
survFitTKTD, 78  
survFitTT, 80  
survFitTT.survDataCstExp, 80

zinc, 82