# Package 'crmPack'

# December 16, 2014

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crmPack-package

Object-oriented implementation of CRM designs

# Description

Object-oriented implementation of CRM designs

## Author(s)

Daniel Sabanes Bove <sabanesd@roche.com>

approximate

 $Approximate\ posterior\ with\ (log)\ normal\ distribution$ 

# Description

Approximate posterior with (log) normal distribution

## Usage

```
approximate(object, ...)
```

## Arguments

object the object ... unused

#### Value

the approximation model

```
approximate,Samples-method
```

Approximate posterior with (log) normal distribution

## **Description**

It is recommended to use set.seed before, in order to be able to reproduce the resulting approximating model exactly.

## Usage

```
## S4 method for signature 'Samples'
approximate(object, model, data, points = seq(from =
    min(data@doseGrid), to = max(data@doseGrid), length = 5L),
    refDose = median(points), logNormal = FALSE, verbose = TRUE, ...)
```

## **Arguments**

object	the Samples object
model	the Model object
data	the Data object
points	optional parameter, which gives the dose values at which the approximation should rely on (default: 5 values equally spaced from minimum to maximum of the dose grid)
refDose	the reference dose to be used (default: median of points)
logNormal	use the log-normal prior? (not default) otherwise, the normal prior for the logistic regression coefficients is used
verbose	be verbose (progress statements and plot)? (default)
•••	additional arguments for ${\tt Quantiles2LogisticNormal}, e.g.$ in order to control the approximation quality, etc.
	unused

# Value

the approximation Model

```
as.list,Data-method as.list method for the "Data" class
```

## **Description**

```
as.list method for the "Data" class
```

## Usage

```
## S4 method for signature 'Data' as.list(x, ...)
```

6 CohortSizeDLT-class

## **Arguments**

x the Data object we want to convert

... objects, possibly named.

#### Value

a list of all slots in x

CohortSize-class

The virtual class for cohort sizes

## Description

The virtual class for cohort sizes

#### See Also

CohortSizeMax, CohortSizeMin, CohortSizeRange, CohortSizeDLT, CohortSizeConst, CohortSizeParts

CohortSizeConst-class Constant cohort size

## Description

This class is used when the cohort size should be kept constant.

## Slots

size the constant integer size

CohortSizeDLT-class

Cohort size based on number of DLTs

# Description

Cohort size based on number of DLTs

#### **Slots**

DLTintervals a vector with the bounds of the relevant DLT intervals of length n cohortSize an integer vector of length n-1 with the cohort sizes in the DLTintervals

CohortSizeMax-class 7

CohortSizeMax-class Size based on maximum of multiple cohort size rules

## **Description**

This class can be used to combine multiple cohort size rules with the MAX operation.

#### **Details**

cohortSizeList contains all cohort size rules, which are again objects of class CohortSize. The maximum of these individual cohort sizes is taken to give the final cohort size.

#### **Slots**

cohortSizeList list of cohort size rules

CohortSizeMin-class Size based on minimum of multiple cohort size rules

#### **Description**

This class can be used to combine multiple cohort size rules with the MIN operation.

#### **Details**

cohortSizeList contains all cohort size rules, which are again objects of class CohortSize. The minimum of these individual cohort sizes is taken to give the final cohort size.

## Slots

cohortSizeList list of cohort size rules

CohortSizeParts-class Cohort size based on the parts

#### **Description**

This class is used when the cohort size should change for the second part of the dose escalation. Only works in conjunction with DataParts objects.

#### **Slots**

sizes the two sizes for part 1 and part 2

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CohortSizeRange-class Cohort size based on dose range

## Description

Cohort size based on dose range

## **Slots**

intervals a vector with the bounds of the relevant dose intervals of length n cohortSize an integer vector of length n-1 with the cohort sizes in the intervals

crmPackExample

Open the example pdf for crmPack

## Description

Calling this helper function should open the example.pdf document, residing in the doc subfolder of the package installation directory.

## Usage

```
crmPackExample()
```

#### Value

nothing

# Author(s)

Daniel Sabanes Bove <sabanesd@roche.com>

crmPackHelp

Open the browser with help pages for crmPack

## Description

This convenience function opens your browser with the help pages for crmPack.

## Usage

```
crmPackHelp()
```

#### Value

nothing

## Author(s)

Daniel Sabanes Bove <sabanesd@roche.com>

Data-class 9

Data-class	Class for the data input	

#### **Description**

Class for the data input

#### **Slots**

```
x the doses for the patients
y the vector of toxicity events (0 or 1 integers)

ID unique patient IDs (integer vector)
cohort the cohort indices (sorted values from 0, 1, 2, ...)
doseGrid the vector of all possible doses (sorted), i.e. the dose grid
nObs number of observations
nGrid number of gridpoints
xLevel the levels for the doses the patients have been given
```

DataDual-class

Class for the dual endpoint data input

#### **Description**

This is a subclass of Data, so contains all slots from Data, and in addition biomarker values.

## **Slots**

w the continuous vector of biomarker values

DataParts-class Class for the data with two study parts

## **Description**

This is a subclass of Data, so contains all slots from Data, and in addition information on the two study parts.

#### **Slots**

```
part integer vector; which part does each of the patients belong to?
nextPart integer; what is the part for the next cohort?
part1Ladder sorted numeric vector; what is the escalation ladder for part 1? This shall be a subset of the doseGrid.
```

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Design-class

Class for the CRM design

## Description

Class for the CRM design

#### **Slots**

```
model the model to be used, an object of class Model
nextBest how to find the next best dose, an object of class NextBest
stopping stopping rule(s) for the trial, an object of class Stopping
increments how to control increments between dose levels, an object of class Increments
cohortSize rules for the cohort sizes, an object of class CohortSize
data what is the dose grid, any previous data, etc., contained in an object of class Data
startingDose what is the starting dose? Must lie on the grid in data
```

dose

Compute the doses for a given probability, given model and samples

## Description

Compute the doses for a given probability, given model and samples

## Usage

```
dose(prob, model, samples, ...)
```

## Arguments

```
prob the probability
model the Model
samples the Samples
... unused
```

```
dose, numeric, Model, Samples-method
```

Compute the doses for a given probability, given model and samples

#### **Description**

Compute the doses for a given probability, given model and samples

#### Usage

```
## S4 method for signature 'numeric, Model, Samples'
dose(prob, model, samples, ...)
```

#### **Arguments**

prob the probability model the Model samples the Samples ... unused

DualEndpoint-class

Dual endpoint model

#### **Description**

todo: describe the model

#### **Slots**

mu For the probit toxicity model, mu contains the prior mean vector

Sigma For the probit toxicity model, contains the prior covariance matrix

- sigma2betaW For the biomarker model, contains the prior variance factor of the random walk prior. If it is not a single number, it can also contain a vector with elements a and b for the inverse-gamma prior on sigma2betaW.
- sigma2W Either a fixed value for the biomarker variance, or a vector with elements a and b for the inverse-gamma prior parameters.
- rho Either a fixed value for the correlation (between -1 and 1), or a vector with elements a and b for the Beta prior on the transformation kappa = (rho + 1) / 2, which is in (0, 1). For example, a=1, b=1 leads to a uniform prior on rho.
- useRW1 for specifying the random walk prior on the biomarker level: if TRUE, RW1 is used, otherwise RW2.
- useFixed a list with logical value for each of the three parameters sigma2betaW, sigma2W and rho indicating whether a fixed value is used or not.

extract

Extract something from an object and produce a data.frame

## Description

Extract something from an object and produce a data.frame

# Usage

```
extract(object, ...)
```

## Arguments

```
object the object ... unused
```

## Value

the data frame

```
extract, Samples-method
```

Extract certain parameter from Samples object

# Description

Extract certain parameter from Samples object

# Usage

```
## S4 method for signature 'Samples'
extract(object, parameter, ...)
```

# Arguments

object the Samples object

parameter the name of the parameter

... unused

## Value

the data frame suitable for use with ggmcmc

fitted,Samples-method

fitted, Samples-method Fit method for the Samples class

## **Description**

Fit method for the Samples class

## Usage

```
## S4 method for signature 'Samples'
fitted(object, model, data, points = data@doseGrid,
  quantiles = c(0.025, 0.975), middle = mean, ...)
```

#### **Arguments**

object the Samples object model the Model object data the Data object

points at which dose levels is the fit requested? default is the dose grid

quantiles the quantiles to be calculated (default: 0.025 and 0.975) middle the function for computing the middle point. Default: mean

... other arguments.

#### Value

data frame with dose, middle, lower and upper quantiles

Increments-class

The virtual class for controlling increments

# Description

The virtual class for controlling increments

#### See Also

IncrementsRelative, IncrementsRelativeDLT, IncrementsRelativeParts

IncrementsRelative-class

Increments control based on relative differences in intervals

#### **Description**

Increments control based on relative differences in intervals

#### **Slots**

intervals a vector with the bounds of the relevant intervals of length n increments a vector of length n-1 with the maximum allowable relative increments in the intervals

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IncrementsRelativeDLT-class

Increments control based on relative differences in terms of DLTs

#### **Description**

Increments control based on relative differences in terms of DLTs

#### **Slots**

DLTintervals a vector with the bounds of the relevant DLT intervals of length n increments a vector of length n-1 with the maximum allowable relative increments in the DLTintervals

IncrementsRelativeParts-class

Increments control based on relative differences in intervals, with special rules for part 1 and beginning of part 2

## **Description**

Note that this only works in conjunction with DataParts objects. If the part 2 will just be started in the next cohort, then the next maximum dose will be either dltStart (e.g. -1) shift of the last part 1 dose in case of a DLT in part 1, or cleanStart shift (e.g. 0) in case of no DLTs in part 1. If part 1 will still be on in the next cohort, then the next dose level will be the next higher dose level in the part1Ladder of the data object. If part 2 has been started before, the usual relative increment rules apply, see IncrementsRelative.

#### Slots

dltStart integer giving the dose level increment for starting part 2 in case of a DLT in part 1 cleanStart integer giving the dose level increment for starting part 2 in case of a DLT in part 1. If this is less or equal to 0, then the part 1 ladder will be used to find the maximum next dose. If this is larger than 0, then the relative increment rules will be applied to find the next maximum dose level.

initialize,Data-method

Initialization method for the "Data" class

## Description

This is the method for initializing a "Data" class object.

#### Usage

```
## S4 method for signature 'Data'
initialize(.Object, x = numeric(), y = integer(),
   ID = integer(), cohort = integer(), doseGrid = numeric(), ...)
```

#### **Arguments**

.Object	the Data we want to initialize
x	the doses for the patients
У	the vector of toxicity events (0 or 1 integers)
ID	unique patient IDs (integer vector)
cohort	the cohort indices (sorted values from 0, 1, 2,)
docoCrid	the vector of all possible doses

doseGrid the vector of all possible doses

data to include in the new object. Named arguments correspond to slots in the

class definition. Unnamed arguments must be objects from classes that this class

extends.

#### **Details**

Note that ID and cohort can be missing, then a warning will be issued and the variables will be filled with default IDs and best guesses, respectively.

```
initialize, DualEndpoint-method
```

Initialization method for the "DualEndpoint" class

#### **Description**

Initialization method for the "DualEndpoint" class

## Usage

```
## $4 method for signature 'DualEndpoint'
initialize(.Object, mu, Sigma, sigma2betaW, sigma2W,
   rho, smooth = c("RW1", "RW2"), ...)
```

## Arguments

. Object the DualEndpoint we want to initialize

mu see DualEndpoint
Sigma see DualEndpoint
sigma2betaW see DualEndpoint
sigma2W see DualEndpoint
rho see DualEndpoint

smooth either "RW1" (default) or "RW2", for specifying the random walk prior on the

biomarker level.

... data to include in the new object. Named arguments correspond to slots in the

class definition. Unnamed arguments must be objects from classes that this class

initialize,LogisticKadane-method

Initialization method for the "LogisticKadane" class

## **Description**

Initialization method for the "LogisticKadane" class

## Usage

```
## S4 method for signature 'LogisticKadane'
initialize(.Object, theta, xmin, xmax, ...)
```

## **Arguments**

. Object the LogisticKadane we want to initialize

theta the target toxicity probability

xmin the minimum of the dose range

xmax the maximum of the dose range

... data to include in the new object. Named arguments correspond to slots in the

class definition. Unnamed arguments must be objects from classes that this class

extends.

initialize,LogisticLogNormal-method

Initialization method for the "LogisticLogNormal" class

## **Description**

Initialization method for the "LogisticLogNormal" class

## Usage

```
## S4 method for signature 'LogisticLogNormal'
initialize(.Object, mean, cov, refDose, ...)
```

# **Arguments**

.Object the LogisticLogNormal we want to initialize

mean the prior mean vector
cov the prior covariance matrix

refDose the reference dose

... data to include in the new object. Named arguments correspond to slots in the

class definition. Unnamed arguments must be objects from classes that this class

```
initialize,LogisticNormal-method
```

Initialization method for the "LogisticNormal" class

#### **Description**

Initialization method for the "LogisticNormal" class

#### Usage

```
## S4 method for signature 'LogisticNormal'
initialize(.Object, mean, cov, refDose, ...)
```

#### **Arguments**

.Object the LogisticNormal we want to initialize

mean the prior mean vector cov the prior covariance matrix

refDose the reference dose

... data to include in the new object. Named arguments correspond to slots in the

class definition. Unnamed arguments must be objects from classes that this class

extends.

initialize, Logistic Normal Fixed Mixture-method

 $Initialization\ method\ for\ the\ "Logistic Normal Fixed Mixture"\ class$ 

## **Description**

Initialization method for the "LogisticNormalFixedMixture" class

# Usage

```
## S4 method for signature 'LogisticNormalFixedMixture'
initialize(.Object, components, weights,
  refDose, logNormal = FALSE, ...)
```

#### **Arguments**

.Object the LogisticNormalFixedMixture we want to initialize

components the specifications of the mixture components: a list with one list of mean and

cov for each bivariate (log) normal prior

weights the weights of the components, these must be positive and will be normalized to

sum to 1

refDose the reference dose

logNormal should a log normal prior be specified, such that the mean vectors and covariance

matrices are valid for the intercept and log slope? (not default)

... data to include in the new object. Named arguments correspond to slots in the

class definition. Unnamed arguments must be objects from classes that this class

```
initialize,LogisticNormalMixture-method
```

Initialization method for the "LogisticNormalMixture" class

#### **Description**

Initialization method for the "LogisticNormalMixture" class

## Usage

```
## $4 method for signature 'LogisticNormalMixture'
initialize(.Object, comp1, comp2, weightpar,
  refDose, ...)
```

## **Arguments**

.Object the LogisticNormalMixture we want to initialize

comp1 the specifications of the first component: a list with mean and cov for the first

bivariate normal prior

comp2 the specifications of the second component

weightpar the beta parameters for the weight of the first component

refDose the reference dose

... data to include in the new object. Named arguments correspond to slots in the

class definition. Unnamed arguments must be objects from classes that this class

extends.

initialize,McmcOptions-method

Initialization method for the "McmcOptions" class

#### **Description**

Initialization method for the "McmcOptions" class

## Usage

```
## S4 method for signature 'McmcOptions'
initialize(.Object, burnin = 10000L, step = 2L,
   samples = 10000L, ...)
```

## **Arguments**

.Object the McmcOptions we want to initialize

burnin number of burn-in iterations which are not saved (default: 10,000) step only every step-th iteration is saved after the burn-in (default: 2) samples number of resulting samples (by default 10,000 will result)

... data to include in the new object. Named arguments correspond to slots in the

class definition. Unnamed arguments must be objects from classes that this class

LogisticKadane-class 19

LogisticKadane-class Reparametrized logistic model

## **Description**

This is the logistic model in the parametrization of Kadane et al. (1980).

#### **Details**

Let  $\rho_0 = p(x_{min})$  be the probability of a DLT and the minimum dose  $x_{min}$ , and let  $\gamma$  be the dose with target toxicity probability  $\theta$ , i.e.  $p(\gamma) = \theta$ . Then it can easily be shown that the logistic regression model has intercept

$$\frac{\gamma logit(\rho_0) - x_{min}logit(\theta)}{\gamma - x_{min}}$$

and slope

$$\frac{logit(theta) - logit(\rho_0)}{\gamma - x_{min}}$$

The prior is a uniform distribution for  $\gamma$  between  $x_{min}$  and  $x_{max}$ , and for  $\rho_0$  as well a uniform distribution between 0 and  $\theta$ .

The slots of this class, required for creating the model, are the target toxicity, as well as the minimum and maximum of the dose range. Note that these can be different from the minimum and maximum of the dose grid in the data later on.

#### **Slots**

theta the target toxicity probability  $\theta$  xmin the minimum of the dose range  $x_{min}$  xmax the maximum of the dose range  $x_{max}$ 

LogisticLogNormal-class

Standard logistic model with bivariate (log) normal prior

# Description

This is the usual logistic regression model with a bivariate normal prior on the intercept and log slope.

#### **Details**

The covariate is the natural logarithm of the dose x divided by the reference dose  $x^*$ :

$$logit[p(x)] = \alpha + \beta \cdot log(x/x^*)$$

where p(x) is the probability of observing a DLT for a given dose x.

The prior is

$$(\alpha, \log(\beta)) \sim Normal(\mu, \Sigma)$$

The slots of this class contain the mean vector and the covariance matrix of the bivariate normal distribution, as well as the reference dose.

#### **Slots**

```
mean the prior mean vector \mu cov the prior covariance matrix \Sigma refDose the reference dose x^*
```

LogisticNormal-class Standard logistic model with bivariate normal prior

## **Description**

This is the usual logistic regression model with a bivariate normal prior on the intercept and slope.

#### **Details**

The covariate is the natural logarithm of the dose x divided by the reference dose  $x^*$ :

$$logit[p(x)] = \alpha + \beta \cdot \log(x/x^*)$$

where p(x) is the probability of observing a DLT for a given dose x.

The prior is

$$(\alpha, \beta) \sim Normal(\mu, \Sigma)$$

The slots of this class contain the mean vector, the covariance and precision matrices of the bivariate normal distribution, as well as the reference dose.

#### **Slots**

```
mean the prior mean vector \mu cov the prior covariance matrix \Sigma prec the prior precision matrix \Sigma^{-1} refDose the reference dose x^*
```

LogisticNormalFixedMixture-class

Standard logistic model with fixed mixture of multiple bivariate (log) normal priors

## **Description**

This is standard logistic regression model with a mixture of multiple bivariate (log) normal priors on the intercept and slope parameters. The weights of the normal priors are fixed, hence no additional model parameters are introduced. This type of prior is often used to better approximate a given posterior distribution, or when the information is given in terms of a mixture.

#### **Details**

The covariate is the natural logarithm of the dose x divided by the reference dose  $x^*$ :

$$logit[p(x)] = \alpha + \beta \cdot \log(x/x^*)$$

where p(x) is the probability of observing a DLT for a given dose x.

The prior is

$$(\alpha, \beta) \sim \sum_{j=1}^{K} w_j Normal(\mu_j, \Sigma_j)$$

if a normal prior is used and

$$(\alpha, \log(\beta)) \sim \sum_{j=1}^{K} w_j Normal(\mu_j, \Sigma_j)$$

if a log normal prior is used.

The weight  $w_i$  of the components are fixed and sum to 1.

The (additional) slots of this class comprise two lists, containing the mean vector, the covariance and precision matrices of the two bivariate normal distributions each, the parameters of the beta prior for the first component weight, as well as the reference dose. Moreover, a slot specifies whether a log normal prior is used.

## **Slots**

components a list with one entry per component of the mixture. Each entry is a list with mean, cov and prec for the bivariate normal prior

weights the weights of the components, these must be positive and sum to 1

refDose the reference dose  $x^*$ 

logNormal is a log normal prior specified for each of the components?

LogisticNormalMixture-class

Standard logistic model with flexible mixture of two bivariate normal priors

## **Description**

This is standard logistic regression model with a mixture of two bivariate normal priors on the intercept and slope parameters. The weight of the two normal priors is a model parameter, hence it is a flexible mixture. This type of prior is often used with a mixture of a minimal informative and an informative component, in order to make the CRM more robust to data deviations from the informative component.

#### **Details**

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The covariate is the natural logarithm of the dose x divided by the reference dose  $x^*$ :

$$logit[p(x)] = \alpha + \beta \cdot \log(x/x^*)$$

where p(x) is the probability of observing a DLT for a given dose x.

The prior is

$$(\alpha, \beta) \sim w * Normal(\mu_1, \Sigma_1) + (1 - w) * Normal(\mu_2, \Sigma_2)$$

The weight w for the first component is assigned a beta prior B(a, b).

The slots of this class comprise two lists, containing the mean vector, the covariance and precision matrices of the two bivariate normal distributions each, the parameters of the beta prior for the first component weight, as well as the reference dose.

#### **Slots**

comp1 the specifications of the first component: a list with mean, cov and prec for the first bivariate normal prior

comp2 the specifications of the second component

weightpar the beta parameters for the weight of the first component

refDose the reference dose  $x^*$ 

logit

Shorthand for logit function

## **Description**

Shorthand for logit function

## Usage

logit(x)

#### **Arguments**

х

the function argument

#### Value

the logit(x)

maxDose 23

maxDose

Determine the maximum possible next dose

## **Description**

Determine the upper limit of the next dose based on the increments rule.

## Usage

```
maxDose(increments, data, ...)
```

## **Arguments**

increments The rule, an object of class Increments
data The data input, an object of class Data

... further arguments

#### **Details**

This function outputs the maximum possible next dose, based on the corresponding rule increments and the data.

#### Value

the maximum possible next dose

```
maxDose,IncrementsRelative,Data-method
```

Determine the maximum possible next dose based on relative increments

# Description

Determine the maximum possible next dose based on relative increments

## Usage

```
## S4 method for signature 'IncrementsRelative,Data'
maxDose(increments, data, ...)
```

## **Arguments**

increments The rule, an object of class Increments

data The data input, an object of class Data

... further arguments

 ${\tt maxDose}$ ,  ${\tt IncrementsRelativeDLT}$ ,  ${\tt Data-method}$ 

Determine the maximum possible next dose based on relative increments determined by DLTs so far

## **Description**

Determine the maximum possible next dose based on relative increments determined by DLTs so far

## Usage

```
## S4 method for signature 'IncrementsRelativeDLT,Data'
maxDose(increments, data, ...)
```

## **Arguments**

increments The rule, an object of class Increments
data The data input, an object of class Data

... further arguments

 $\verb|maxDose|, Increments Relative Parts|, Data Parts-method|$ 

Determine the maximum possible next dose based on relative increments and part 1 and 2

#### **Description**

Determine the maximum possible next dose based on relative increments and part 1 and 2

## Usage

```
## S4 method for signature 'IncrementsRelativeParts,DataParts'
maxDose(increments, data, ...)
```

#### **Arguments**

increments The rule, an object of class Increments
data The data input, an object of class Data
... further arguments

maxSize 25

maxSize

"MAX" combination of cohort size rules

# Description

This function combines cohort size rules by taking the maximum of all sizes.

## Usage

```
maxSize(...)
```

## **Arguments**

... Objects of class CohortSize

## Value

the combination as an object of class CohortSizeMax

#### See Also

minSize

maxSize,CohortSize-method

The method combining cohort size rules by taking maximum

## **Description**

The method combining cohort size rules by taking maximum

# Usage

```
## S4 method for signature 'CohortSize'
maxSize(...)
```

# **Arguments**

... Objects of class CohortSize

mcmc

Obtain posterior samples for all model parameters

## **Description**

Obtain posterior samples for all model parameters

#### Usage

```
mcmc(data, model, options, ...)
```

#### **Arguments**

data The data input, an object of class Data
model The model input, an object of class Model
options MCMC options, an object of class McmcOptions
... unused

#### **Details**

This is the function to actually run the MCMC machinery to produce posterior samples from all model parameters and required derived values. It is a generic function, so that customized versions may be conveniently defined for specific subclasses of Data, Model, and McmcOptions input.

#### Value

The posterior samples, an object of class Samples.

#### **Description**

The fast method for the LogisticNormal class

## Usage

```
## S4 method for signature 'Data,LogisticNormal,McmcOptions'
mcmc(data, model, options,
  verbose = FALSE, ...)
```

# Arguments

verbose shall messages be printed? (not default)
data The data input, an object of class Data
model The model input, an object of class Model
options MCMC options, an object of class McmcOptions

.. unused

## Description

Standard method which uses JAGS/BUGS

# Usage

```
## S4 method for signature 'Data,Model,McmcOptions'
mcmc(data, model, options,
   program = c("JAGS", "OpenBUGS", "WinBUGS"), verbose = FALSE, ...)
```

## Arguments

program	the program which shall be used: either "JAGS" (default), "OpenBUGS" or "WinBUGS"
verbose	shall messages be printed? (not default)
data	The data input, an object of class Data
model	The model input, an object of class Model
options	MCMC options, an object of class McmcOptions
	unused

McmcOptions-class

Class for the three canonical MCMC options

## Description

Class for the three canonical MCMC options

## **Slots**

iterations number of MCMC iterations burnin number of burn-in iterations which are not saved step only every step-th iteration is saved after the burn-in 28 minSize

MinimalInformative

Construct a minimally informative prior

#### **Description**

This function constructs a minimally informative prior, which is captured in a LogisticNormal object.

#### Usage

```
MinimalInformative(dosegrid, refDose, threshmin = 0.2, threshmax = 0.3, ...)
```

#### **Arguments**

dosegrid the dose grid

refDose the reference dose

threshmin Any toxicity probability above this threshold would be very unlikely (5%) at the minimum dose (default: 0.2)

threshmax Any toxicity probability below this threshold would be very unlikely (5%) at the maximum dose (default: 0.3)

... additional arguments for computations, see Quantiles2LogisticNormal

#### **Details**

Based on the proposal by Neuenschwander et al (2008, Statistics in Medicine), a minimally informative prior distribution is constructed. The required key input is the minimum ( $d_1$  in the notation of the Appendix A.1 of that paper) and the maximum value ( $d_J$ ) of the dose grid supplied to this function. Then threshmin is the probability threshold  $q_1$ , such that any probability of DLT larger than  $q_1$  has only 5% probability. Likewise, threshmax is the probability threshold  $q_J$ , such that any probability of DLT smaller than  $q_J$  has only 5% probability. Subsequently, for all doses supplied in the dosegrid argument, Beta distributions are set up, and Quantiles2LogisticNormal is used to transform the resulting quantiles into an approximating LogisticNormal model.

#### Value

see Quantiles2LogisticNormal

minSize

"MIN" combination of cohort size rules

#### **Description**

This function combines cohort size rules by taking the minimum of all sizes.

## Usage

```
minSize(...)
```

#### **Arguments**

... Objects of class CohortSize

#### Value

the combination as an object of class CohortSizeMin

#### See Also

maxSize

minSize, CohortSize-method

The method combining cohort size rules by taking minimum

## **Description**

The method combining cohort size rules by taking minimum

#### Usage

```
## S4 method for signature 'CohortSize'
minSize(...)
```

## **Arguments**

... Objects of class CohortSize

Model-class

Class for the model input

# Description

This is the general model class, from which all other specific models inherit.

## **Details**

The datamodel must obey the convention that the data input is called exactly as in the Data class. All prior distributions for parameters should be contained in the model function priormodel. The background is that this can be used to simulate from the prior distribution, before obtaining any data.

The dose function has as first argument prob, a scalar toxicity probability which is targeted. Additional arguments are model parameters. Then it computes, using model parameter(s) (samples), the resulting dose. Note that the model parameters are called exactly as in the model and must be included in the sample vector. The vectors of all samples for these parameters will then be supplied to the function. So your function must be able to process vectors of the model parameters, i.e. it must vectorize over them.

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The prob function has as first argument dose, which is a scalar dose. Additional arguments are model parameters. Then it computes, using model parameter(s) (samples), the resulting probability of toxicity at that dose. Again here, the function must vectorize over the model parameters.

If you work with multivariate parameters, then please assume that your the two functions receive either one parameter value as a row vector, or a samples matrix where the rows correspond to the sampling index, i.e. the layout is then nSamples x dimParameter.

Note that dose and prob are the inverse functions of each other.

#### **Slots**

datamodel a function representing the BUGS data model specification (see the details above) priormodel a function representing the BUGS prior specification (see the details above)

datanames The names of all Data slots that are used in the datamodel and/or priormodel definition. Note that you cannot specify more variables than those that are really used in the model!

modelspecs a function computing the list of the data model and prior model specifications that are required for fully specifying them (e.g. prior parameters, reference dose, etc.), based on the Data slots that are then required as arguments of this function. This will then be passed to BUGS for the computations.

dose a function computing the dose reaching a specific target probability, based on the model parameters and additional prior settings (see the details above)

prob a function computing the probability of toxicity for a specific dose, based on the model parameters and additional prior settings (see the details above)

init a function computing the list of starting values for parameters required to be initialized in the MCMC sampler, based on the Data slots that are then required as arguments of this function

sample names of all parameters from which you would like to save the MCMC samples. These must include the ones required by the dose and prob functions.

## See Also

LogisticNormal, LogisticLogNormal, LogisticKadane, DualEndpoint

nextBest Find the next best dose

# Description

Compute the recommended next best dose.

## Usage

```
nextBest(nextBest, doselimit, samples, model, data, ...)
```

#### **Arguments**

nextBest The rule, an object of class NextBest doselimit The maximum allowed next dose

samples the Samples object

model The model input, an object of class Model data The data input, an object of class Data

... possible additional arguments without method dispatch

#### **Details**

This function outputs the next best dose recommendation based on the corresponding rule nextBest, the posterior samples from the model and the underlying data.

#### Value

a list with the next best dose (element value) on the grid defined in data, and a plot depicting this recommendation (element plot)

nextBest, NextBestDualEndpoint, numeric, Samples, DualEndpoint, Data-method

Find the next best dose based on the dual endpoint model

## Description

Find the next best dose based on the dual endpoint model

## Usage

```
## S4 method for signature
## 'NextBestDualEndpoint, numeric, Samples, DualEndpoint, Data'
nextBest(nextBest,
  doselimit, samples, model, data, ...)
```

## **Arguments**

nextBest The rule, an object of class NextBest doselimit The maximum allowed next dose

samples the Samples object

model The model input, an object of class Model data The data input, an object of class Data

... possible additional arguments without method dispatch

nextBest, NextBestMTD, numeric, Samples, Model, Data-method

Find the next best dose based on the MTD rule

## **Description**

Find the next best dose based on the MTD rule

#### Usage

```
## $4 method for signature 'NextBestMTD, numeric, Samples, Model, Data'
nextBest(nextBest, doselimit,
   samples, model, data, ...)
```

#### **Arguments**

nextBest The rule, an object of class NextBest doselimit The maximum allowed next dose

samples the Samples object

model The model input, an object of class Model data The data input, an object of class Data

... possible additional arguments without method dispatch

nextBest, NextBestNCRM, numeric, Samples, Model, Data-method

Find the next best dose based on the NCRM method

## **Description**

Find the next best dose based on the NCRM method

#### Usage

```
## S4 method for signature 'NextBestNCRM, numeric, Samples, Model, Data'
nextBest(nextBest,
   doselimit, samples, model, data, ...)
```

# **Arguments**

nextBest The rule, an object of class NextBest doselimit The maximum allowed next dose

samples the Samples object

model The model input, an object of class Model data The data input, an object of class Data

... possible additional arguments without method dispatch

nextBest,NextBestNCRM,numeric,Samples,Model,DataParts-method

Find the next best dose based on the NCRM method when two parts

trial is used

## **Description**

Find the next best dose based on the NCRM method when two parts trial is used

#### Usage

```
## S4 method for signature 'NextBestNCRM,numeric,Samples,Model,DataParts'
nextBest(nextBest,
   doselimit, samples, model, data, ...)
```

NextBest-class 33

#### **Arguments**

nextBest The rule, an object of class NextBest doselimit The maximum allowed next dose

samples the Samples object

model The model input, an object of class Model data The data input, an object of class Data

... possible additional arguments without method dispatch

NextBest-class

The virtual class for finding next best dose

## **Description**

The virtual class for finding next best dose

#### See Also

NextBestMTD, NextBestNCRM, NextBestDualEndpoint

NextBestDualEndpoint-class

The class with the input for finding the next dose based on the dual endpoint model

# Description

The class with the input for finding the next dose based on the dual endpoint model

#### **Slots**

target the biomarker level, relative to the maximum, that needs to be reached. For example, 0.9 means that a dose with 90 of the maximum biomarker level is considered as having reached sufficient biomarker level.

overdose the overdose toxicity interval

maxOverdoseProb maximum overdose probability that is allowed

NextBestMTD-class

The class with the input for finding the next best MTD estimate

## **Description**

The class with the input for finding the next best MTD estimate

## **Slots**

target the target toxicity probability

derive the function which derives from the input, a vector of posterior MTD samples called mtdSamples, the final next best MTD estimate.

NextBestNCRM-class

The class with the input for finding the next dose in target interval

## **Description**

Note that to avoid numerical problems, the dose selection algorithm has been implemented as follows: First admissible doses are found, which are those with probability to fall in overdose category being below maxOverdoseProb. Next, within the admissible doses, the maximum probability to fall in the target category is calculated. If that is above 5% (i.e., it is not just numerical error), then the corresponding dose is the next recommended dose. Otherwise, the highest admissible dose is the next recommended dose.

## **Slots**

```
target the target toxicity interval

overdose the overdose toxicity interval

maxOverdoseProb maximum overdose probability that is allowed
```

or-Stopping-Stopping The method combining two atomic stopping rules

# Description

The method combining two atomic stopping rules

#### Usage

```
## S4 method for signature 'Stopping,Stopping'
e1 | e2
```

## **Arguments**

e1 First Stopping object e2 Second Stopping object

## Value

The StoppingAny object

```
or-Stopping-StoppingAny
```

The method combining a stopping list and an atomic

# Description

The method combining a stopping list and an atomic

## Usage

```
## S4 method for signature 'StoppingAny,Stopping'
e1 | e2
```

## **Arguments**

```
e1 StoppingAny object
e2 Stopping object
```

## Value

The modified StoppingAny object

```
or-StoppingAny-Stopping
```

The method combining an atomic and a stopping list

# Description

The method combining an atomic and a stopping list

# Usage

```
## S4 method for signature 'Stopping,StoppingAny'
e1 | e2
```

# Arguments

```
e1 Stopping object
e2 StoppingAny object
```

## Value

The modified StoppingAny object

```
plot, Data, missing-method
```

Plot method for the "Data" class

## **Description**

Plot method for the "Data" class

## Usage

```
## S4 method for signature 'Data,missing'
plot(x, y, ...)
```

## **Arguments**

x the Data object we want to plot

y the y coordinates of points in the plot, *optional* if x is an appropriate structure.

Arguments to be passed to methods, such as graphical parameters (see par).

Many methods will accept the following arguments:

type what type of plot should be drawn. Possible types are

- "p" for points,
- "1" for lines,
- "b" for **b**oth,
- "c" for the lines part alone of "b",
- "o" for both 'overplotted',
- "h" for 'histogram' like (or 'high-density') vertical lines,
- "s" for stair steps,
- "S" for other steps, see 'Details' below,
- "n" for no plotting.

All other types give a warning or an error; using, e.g., type = "punkte" being equivalent to type = "p" for S compatibility. Note that some methods, e.g. plot.factor, do not accept this.

```
main an overall title for the plot: see title. sub a sub title for the plot: see title. xlab a title for the x axis: see title. ylab a title for the y axis: see title. asp the y/x aspect ratio, see plot.window.
```

#### Value

the ggplot object

```
plot, DataDual, missing-method
```

Plot method for the "DataDual" class

# **Description**

Plot method for the "DataDual" class

# Usage

```
## S4 method for signature 'DataDual,missing' plot(x, y, ...)
```

#### **Arguments**

x the DataDual object we want to plot

y the y coordinates of points in the plot, *optional* if x is an appropriate structure.

Arguments to be passed to methods, such as graphical parameters (see par).

Many methods will accept the following arguments:

type what type of plot should be drawn. Possible types are

- "p" for **p**oints,
- "1" for lines,
- "b" for **b**oth,
- "c" for the lines part alone of "b",
- "o" for both 'overplotted',
- "h" for 'histogram' like (or 'high-density') vertical lines,
- "s" for stair steps,
- "S" for other steps, see 'Details' below,
- "n" for no plotting.

All other types give a warning or an error; using, e.g., type = "punkte" being equivalent to type = "p" for S compatibility. Note that some methods, e.g. plot.factor, do not accept this.

```
main an overall title for the plot: see title. sub a sub title for the plot: see title. xlab a title for the x axis: see title. ylab a title for the y axis: see title. asp the y/x aspect ratio, see plot.window.
```

## Value

```
the ggplot object
```

```
plot, Samples, DualEndpoint-method
```

Plot method for the "Samples" object, when we have the dual endpoint model

#### **Description**

Plot method for the "Samples" object, when we have the dual endpoint model

#### Usage

```
## S4 method for signature 'Samples,DualEndpoint'
plot(x, y, data, extrapolate = TRUE, ...)
```

#### **Arguments**

```
x the Samples object
y the DualEndpoint object
data the DataDual object
extrapolate should the biomarker fit be extrapolated to the whole dose grid? (default)
... Arguments to be passed to methods, such as graphical parameters (see par).
Many methods will accept the following arguments:
type what type of plot should be drawn. Possible types are
```

- "p" for **p**oints,
- "1" for lines,
- "b" for **b**oth,
- "c" for the lines part alone of "b",
- "o" for both 'overplotted',
- "h" for 'histogram' like (or 'high-density') vertical lines,
- "s" for stair steps,
- "S" for other steps, see 'Details' below,
- "n" for no plotting.

All other types give a warning or an error; using, e.g., type = "punkte" being equivalent to type = "p" for S compatibility. Note that some methods, e.g. plot.factor, do not accept this.

```
main an overall title for the plot: see title. sub a sub title for the plot: see title. xlab a title for the x axis: see title. ylab a title for the y axis: see title. asp the y/x aspect ratio, see plot.window.
```

#### Value

the ggplot object

```
plot, Samples, Model-method
```

Plot method for the "Samples" and "Model" object

# **Description**

Plot method for the "Samples" and "Model" object

# Usage

```
## S4 method for signature 'Samples,Model'
plot(x, y, data, ..., xlab = "Dose level",
  ylab = "Probability of DLT [%]")
```

# Arguments

```
x the Samples object

y the Model object

data the Data object

xlab the x axis label

ylab the y axis label

... Arguments to be passed to methods, such as graphical parameters (see par).

Many methods will accept the following arguments:
```

type what type of plot should be drawn. Possible types are

- "p" for points,
- "1" for lines,
- "b" for **b**oth,
- "c" for the lines part alone of "b",
- "o" for both 'overplotted',
- "h" for 'histogram' like (or 'high-density') vertical lines,
- "s" for stair steps,
- "S" for other steps, see 'Details' below,
- "n" for no plotting.

All other types give a warning or an error; using, e.g., type = "punkte" being equivalent to type = "p" for S compatibility. Note that some methods, e.g. plot.factor, do not accept this.

```
main an overall title for the plot: see title. sub a sub title for the plot: see title. xlab a title for the x axis: see title. ylab a title for the y axis: see title. asp the y/x aspect ratio, see plot.window.
```

# Value

the ggplot object

```
{\it plot}, {\it Simulations}, {\it missing-method} \\ {\it Plot simulations}
```

# **Description**

Summarize the simulations with plots

#### Usage

```
## S4 method for signature 'Simulations,missing'
plot(x, y, type = c("trajectory",
   "dosesTried"), ...)
```

#### **Arguments**

the Simulations object we want to plot from
 type the type of plots you want to obtain.
 the y coordinates of points in the plot, *optional* if x is an appropriate structure.
 Arguments to be passed to methods, such as graphical parameters (see par).
 Many methods will accept the following arguments:

type what type of plot should be drawn. Possible types are

- "p" for points,
- "1" for lines,
- "b" for **b**oth,
- "c" for the lines part alone of "b",
- "o" for both 'overplotted',
- "h" for 'histogram' like (or 'high-density') vertical lines,
- "s" for stair steps,
- "S" for other steps, see 'Details' below,
- "n" for no plotting.

All other types give a warning or an error; using, e.g., type = "punkte" being equivalent to type = "p" for S compatibility. Note that some methods, e.g. plot.factor, do not accept this.

```
main an overall title for the plot: see title. sub a sub title for the plot: see title. xlab a title for the x axis: see title. ylab a title for the y axis: see title. asp the y/x aspect ratio, see plot.window.
```

# **Details**

This plot method can be applied to Simulations objects in order to summarize them graphically. Possible types of plots at the moment are:

**trajectory** Summary of the trajectory of the simulated trials **dosesTried** Average proportions of the doses tested in patients

You can specify one or both of these in the type argument.

#### Value

A single ggplot2 object if a single plot is asked for, otherwise a gridExtra{gTree} object.

```
{\it plot}, {\it Simulations-summary, missing-method} \\ {\it Plot summaries of the simulations}
```

#### **Description**

Graphical display of the simulation summary

# Usage

```
## S4 method for signature 'Simulations-summary,missing'
plot(x, y, type = c("nObs",
   "doseSelected", "propDLTs", "nAboveTarget", "meanFit"), ...)
```

#### **Arguments**

x the Simulations-summary object we want to plot from

type the types of plots you want to obtain.

y the y coordinates of points in the plot, *optional* if x is an appropriate structure.

Arguments to be passed to methods, such as graphical parameters (see par).

Many methods will accept the following arguments:

type what type of plot should be drawn. Possible types are

- "p" for **p**oints,
- "1" for lines,
- "b" for **b**oth.
- "c" for the lines part alone of "b",
- "o" for both 'overplotted',
- "h" for 'histogram' like (or 'high-density') vertical lines,
- "s" for stair steps,
- "S" for other steps, see 'Details' below,
- "n" for no plotting.

All other types give a warning or an error; using, e.g., type = "punkte" being equivalent to type = "p" for S compatibility. Note that some methods, e.g. plot.factor, do not accept this.

```
main an overall title for the plot: see title. sub a sub title for the plot: see title. xlab a title for the x axis: see title. ylab a title for the y axis: see title. asp the y/x aspect ratio, see plot.window.
```

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#### **Details**

This plot method can be applied to Simulations objects in order to summarize them graphically. Possible types of plots at the moment are:

**nObs** Distribution of the number of patients in the simulated trials

**doseSelected** Distribution of the final selected doses in the trials. Note that this can include zero entries, meaning that the trial was stopped because all doses in the dose grid appeared too toxic.

**propDLTs** Distribution of the proportion of patients with DLTs in the trials

**nAboveTarget** Distribution of the number of patients treated at doses which are above the target toxicity interval (as specified by the truth and target arguments to summary, Simulations-method)

**meanFit** Plot showing the average fitted dose-toxicity curve across the trials, together with 95% credible intervals, and comparison with the assumed truth (as specified by the truth argument to summary, Simulations-method)

You can specify any subset of these in the type argument.

#### Value

A single ggplot2 object if a single plot is asked for, otherwise a gridExtra{gTree} object.

prob

Compute the probability for a given dose, given model and samples

## **Description**

Compute the probability for a given dose, given model and samples

# Usage

```
prob(dose, model, samples, ...)
```

# Arguments

```
dose the dose
model the Model
samples the Samples
... unused
```

```
prob, numeric, Model, Samples-method
```

Compute the probability for a given dose, given model and samples

# **Description**

Compute the probability for a given dose, given model and samples

# Usage

```
## S4 method for signature 'numeric, Model, Samples'
prob(dose, model, samples, ...)
```

#### **Arguments**

```
dose
                  the dose
model
                  the Model
                  the Samples
samples
                  unused
```

Quantiles2LogisticNormal

Convert prior quantiles (lower, median, upper) to logistic (log) normal model

# **Description**

This function uses generalised simulated annealing to optimise a LogisticNormal model to be as close as possible to the given prior quantiles.

# Usage

```
Quantiles2LogisticNormal(dosegrid, refDose, lower, median, upper,
 level = 0.95, logNormal = FALSE, parstart = NULL, parlower = c(-10,
  -10, 0, 0, -0.95), parupper = c(10, 10, 10, 10, 0.95), verbose = TRUE,
 control = list(threshold.stop = 0.01, maxit = 50000, temperature = 50000,
 max.time = 600))
```

# **Arguments**

dosegrid the dose grid refDose the reference dose lower the lower quantiles median the medians

upper the upper quantiles

level the credible level of the (lower, upper) intervals (default: 0.95) 44 sampleSize

logNormal	use the log-normal prior? (not default) otherwise, the normal prior for the logistic regression coefficients is used
parstart	starting values for the parameters. By default, these are determined from the medians supplied.
parlower	lower bounds on the parameters (intercept alpha and the slope beta, the corresponding standard deviations and the correlation.)
parupper	upper bounds on the parameters
verbose	be verbose? (default)
control	additional options for the optimisation routine, see GenSA for more details

#### Value

a list with the best approximating model (LogisticNormal or LogisticLogNormal), the resulting quantiles, the required quantiles and the distance to the required quantiles, as well as the final parameters (which could be used for running the algorithm a second time)

Samples-class

Class for the MCMC output

# Description

Class for the MCMC output

#### **Slots**

data a list where each entry contains the samples of a (vector-valued) parameter in a vector/matrix in the format (number of samples) x (dimension of the parameter).

options the  ${\tt McmcOptions}$  which have been used

 ${\tt sampleSize}$ 

Compute the number of samples for a given MCMC options triple

# **Description**

Compute the number of samples for a given MCMC options triple

# Usage

sampleSize(mcmcOptions)

# Arguments

mcmcOptions the McmcOptions object

#### Value

the resulting sample size

```
show, Simulations-summary-method
```

Show the summary of the simulations

#### **Description**

Show the summary of the simulations

#### Usage

```
## S4 method for signature 'Simulations-summary'
show(object)
```

#### **Arguments**

object

the Simulations-summary object we want to print

#### Value

invisibly returns a data frame of the results with one row and appropriate column names

```
simulate, Design-method
```

Simulate outcomes from a CRM design

#### **Description**

Simulate outcomes from a CRM design

# Usage

```
## S4 method for signature 'Design'
simulate(object, truth, args = NULL,
  firstSeparate = FALSE, nsim = 1L, mcmcOptions = new("McmcOptions"),
  seed = NULL, parallel = FALSE, ...)
```

# **Arguments**

object the Design object we want to simulate data from

truth a function which takes as input a dose (vector) and returns the true probability

(vector) for toxicity. Additional arguments can be supplied in args.

args data frame with arguments for the truth function. The column names corre-

spond to the argument names, the rows to the values of the arguments. The rows are appropriately recycled in the nsim simulations. In order to produce outcomes from the posterior predictive distribution, e.g, pass an object that contains the data observed so far, truth contains the prob function from the

model in object, and args contains posterior samples from the model.

firstSeparate enroll the first patient separately from the rest of the cohort? (not default) If yes,

the cohort will be closed if a DLT occurs in this patient.

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nsim the number of simulations (default: 1)

mcmcOptions object of class McmcOptions, giving the MCMC options for each evaluation in

the trial. By default, the standard options are used

seed an object specifying if and how the random number generator should be initial-

ized ("seeded"). Either NULL (default) or an integer that will be used in a call to set.seed before simulating the response vectors. If set, the value is saved as the seed slot of the returned object. The default, NULL will not change the

random generator state, and .Random. seed will be saved.

parallel should the simulation runs be parallelized across the clusters of the computer?

(not default)

... additional optional arguments.

#### Value

an object of class Simulations

Simulations-class

Class for the simulations output

# **Description**

This class captures the trial simulations.

## **Details**

Here also the random generator state before starting the simulation is saved, in order to be able to reproduce the outcome. For this just use set.seed with the seed as argument before running simulate, Design-method.

#### Slots

data list of produced Data objects

doses the vector of final dose recommendations

fit list with the final fits

stopReasons list of stopping reasons for each simulation run

seed random generator state before starting the simulation

Simulations-summary-class

Class for the summary of simulations output

#### **Description**

Class for the summary of simulations output

#### **Slots**

target target toxicity interval
targetDoseInterval corresponding target dose interval
nsim number of simulations
propDLTs proportions of DLTs in the trials
meanToxRisk mean toxicity risks for the patients
doseSelected doses selected as MTD
toxAtDosesSelected true toxicity at doses selected
propAtTarget Proportion of trials selecting target MTD
doseMostSelected dose most often selected as MTD
obsToxRateAtDoseMostSelected observed toxicity rate at dose most often selected
fitAtDoseMostSelected fitted toxicity rate at dose most often selected
meanFit list with the average, lower (2.5 quantiles of the mean fitted toxicity at each dose level
nObs number of patients overall
nAboveTarget number of patients treated above target tox interval
doseGrid the dose grid that has been used

size

Determine the size of the next cohort

# Description

This function determines the size of the next cohort.

# Usage

```
size(cohortSize, dose, data, ...)
```

# **Arguments**

cohortSize The rule, an object of class CohortSize
dose the next dose
data The data input, an object of class Data
... additional arguments

#### Value

the size as integer value

# Description

Constant cohort size

# Usage

```
## S4 method for signature 'CohortSizeConst, ANY, Data'
size(cohortSize, dose, data, ...)
```

# **Arguments**

cohortSize The rule, an object of class CohortSize

dose the next dose

data The data input, an object of class Data

... additional arguments

size,CohortSizeDLT,ANY,Data-method

Determine the cohort size based on the number of DLTs so far

# Description

Determine the cohort size based on the number of DLTs so far

# Usage

```
## S4 method for signature 'CohortSizeDLT,ANY,Data'
size(cohortSize, dose, data, ...)
```

# Arguments

cohortSize The rule, an object of class CohortSize

dose the next dose

data The data input, an object of class Data

```
size,CohortSizeMax,ANY,Data-method
```

Size based on maximum of multiple cohort size rules

# Description

Size based on maximum of multiple cohort size rules

# Usage

```
## S4 method for signature 'CohortSizeMax,ANY,Data'
size(cohortSize, dose, data, ...)
```

# Arguments

cohortSize The rule, an object of class CohortSize

dose the next dose

data The data input, an object of class Data

... additional arguments

size,CohortSizeMin,ANY,Data-method

Size based on minimum of multiple cohort size rules

# **Description**

Size based on minimum of multiple cohort size rules

# Usage

```
## S4 method for signature 'CohortSizeMin,ANY,Data'
size(cohortSize, dose, data, ...)
```

# Arguments

cohortSize The rule, an object of class CohortSize

dose the next dose

data The data input, an object of class Data

# Description

Cohort size based on the parts

# Usage

```
## S4 method for signature 'CohortSizeParts,ANY,DataParts'
size(cohortSize, dose, data, ...)
```

#### **Arguments**

cohortSize The rule, an object of class CohortSize

dose the next dose

data The data input, an object of class Data

... additional arguments

 $\verb+size+, \verb+CohortSize+, \verb+ANY+, \verb+Data-method+ \\$ 

Determine the cohort size based on the range into which the next dose falls into

# **Description**

Determine the cohort size based on the range into which the next dose falls into

# Usage

```
## S4 method for signature 'CohortSizeRange,ANY,Data'
size(cohortSize, dose, data, ...)
```

# Arguments

cohortSize The rule, an object of class CohortSize

dose the next dose

data The data input, an object of class Data

Stopping-class 51

Stopping-class	The virtual class for stopping rules	

#### **Description**

The virtual class for stopping rules

#### See Also

Stopping List, Stopping Max Patients, Stopping Cohorts Near Dose, Stopping Patients Near Dose, Stopping Min Cohorts, Stopping Min Patients, Stopping Target Prob Stopping MTD distribution, Stopping Target Biomarker

StoppingAll-class Stop based on fullfillment of all multiple stopping rules

# **Description**

This class can be used to combine multiple stopping rules with an AND operator.

#### **Details**

stopList contains all stopping rules, which are again objects of class Stopping. All stopping rules must be fulfilled in order that the result of this rule is to stop.

# **Slots**

stopList list of stopping rules of the stopping rules into a single result

StoppingAny-class Stop based on fullfillment of any stopping rule

# **Description**

This class can be used to combine multiple stopping rules with an OR operator.

## **Details**

stopList contains all stopping rules, which are again objects of class Stopping. Any of these rules must be fulfilled in order that the result of this rule is to stop.

# **Slots**

stopList list of stopping rules of the stopping rules into a single result

StoppingCohortsNearDose-class

Stop based on number of cohorts near to next best dose

# Description

Stop based on number of cohorts near to next best dose

# **Slots**

nCohorts number of required cohorts
percentage percentage (between 0 and 100) within the next best dose the cohorts must lie

StoppingList-class

Stop based on multiple stopping rules

# Description

This class can be used to combine multiple stopping rules.

#### **Details**

stopList contains all stopping rules, which are again objects of class Stopping, and the summary is a function taking a logical vector of the size of stopList and returning a single logical value. For example, if the function all is given as summary function, then this means that all stopping rules must be fulfilled in order that the result of this rule is to stop.

#### **Slots**

stopList list of stopping rules

summary the summary function to combine the results of the stopping rules into a single result

StoppingMaxPatients-class

Stop based on maximum number of patients

# Description

Stop based on maximum number of patients

## **Slots**

nPatients maximum allowed number of patients

StoppingMinCohorts-class

Stop based on minimum number of cohorts

# **Description**

Stop based on minimum number of cohorts

# Slots

nCohorts minimum required number of cohorts

StoppingMinPatients-class

Stop based on minimum number of patients

# Description

Stop based on minimum number of patients

# **Slots**

nPatients minimum required number of patients

StoppingMTDdistribution-class

Stop based on MTD distribution

# Description

Has 90% probability above a threshold of 50% of the current MTD been reached? This class is used for this question.

# Slots

```
target the target toxicity probability (e.g. 0.33) defining the MTD thresh the threshold relative to the MTD (e.g. 0.5) prob required probability (e.g. 0.9)
```

StoppingPatientsNearDose-class

Stop based on number of patients near to next best dose

# Description

Stop based on number of patients near to next best dose

#### **Slots**

nPatients number of required patients
percentage percentage (between 0 and 100) within the next best dose the patients must lie

StoppingTargetBiomarker-class

Stop based on probability of target biomarker

# Description

Stop based on probability of target biomarker

# Slots

target the biomarker level, relative to the maximum, that needs to be reached prob required target probability for reaching sufficient precision

 ${\tt StoppingTargetProb-class}$ 

Stop based on probability of target tox interval

# **Description**

Stop based on probability of target tox interval

# **Slots**

target the target toxicity interval prob required target toxicity probability for reaching sufficient precision

stopTrial 55

	Stop the trial?	stopTrial
--	-----------------	-----------

# **Description**

This function returns whether to stop the trial.

# Usage

```
stopTrial(stopping, dose, samples, model, data, ...)
```

#### **Arguments**

```
stopping The rule, an object of class Stopping dose the recommended next best dose samples the Samples object
```

model The model input, an object of class Model data The data input, an object of class Data

... additional arguments

#### Value

logical value: TRUE if the trial can be stopped, FALSE otherwise. It should have an attribute message which gives the reason for the decision.

```
stopTrial, StoppingAll, ANY, ANY, ANY, ANY-method

Stop based on fulfillment of all multiple stopping rules
```

# **Description**

Stop based on fulfillment of all multiple stopping rules

# Usage

```
## S4 method for signature 'StoppingAll,ANY,ANY,ANY,ANY'
stopTrial(stopping, dose, samples,
   model, data, ...)
```

# Arguments

stopping	The rule, an object of class Stopping
dose	the recommended next best dose
samples	the Samples object

model The model input, an object of class Model data The data input, an object of class Data

```
stopTrial, StoppingAny, ANY, ANY, ANY, ANY-method

Stop based on fulfillment of any stopping rule
```

#### **Description**

Stop based on fulfillment of any stopping rule

# Usage

```
## S4 method for signature 'StoppingAny,ANY,ANY,ANY',
stopTrial(stopping, dose, samples,
   model, data, ...)
```

# **Arguments**

stopping The rule, an object of class Stopping dose the recommended next best dose

samples the Samples object

model The model input, an object of class Model data The data input, an object of class Data

... additional arguments

stopTrial,StoppingCohortsNearDose,numeric,ANY,ANY,Data-method

Stop based on number of cohorts near to next best dose

# **Description**

Stop based on number of cohorts near to next best dose

# Usage

```
## S4 method for signature 'StoppingCohortsNearDose,numeric,ANY,ANY,Data'
stopTrial(stopping,
   dose, samples, model, data, ...)
```

# **Arguments**

stopping The rule, an object of class Stopping dose the recommended next best dose

samples the Samples object

model The model input, an object of class Model data The data input, an object of class Data

```
stopTrial,StoppingList,ANY,ANY,ANY,ANY-method

Stop based on multiple stopping rules
```

#### **Description**

Stop based on multiple stopping rules

# Usage

```
## S4 method for signature 'StoppingList, ANY, ANY, ANY, ANY'
stopTrial(stopping, dose, samples,
   model, data, ...)
```

# **Arguments**

stopping The rule, an object of class Stopping dose the recommended next best dose

samples the Samples object

model The model input, an object of class Model data The data input, an object of class Data

... additional arguments

```
stopTrial, StoppingMaxPatients, ANY, ANY, ANY, Data-method

Stop based on maximum number of patients
```

# **Description**

Stop based on maximum number of patients

# Usage

```
## S4 method for signature 'StoppingMaxPatients, ANY, ANY, ANY, Data'
stopTrial(stopping, dose,
   samples, model, data, ...)
```

# **Arguments**

stopping The rule, an object of class Stopping dose the recommended next best dose samples the Samples object

model The model input, an object of class Model data The data input, an object of class Data

stopTrial, StoppingMinCohorts, ANY, ANY, ANY, Data-method

Stop based on minimum number of cohorts

#### **Description**

Stop based on minimum number of cohorts

# Usage

```
## S4 method for signature 'StoppingMinCohorts, ANY, ANY, Data'
stopTrial(stopping, dose,
   samples, model, data, ...)
```

# **Arguments**

stopping The rule, an object of class Stopping dose the recommended next best dose

samples the Samples object

model The model input, an object of class Model data The data input, an object of class Data

... additional arguments

stopTrial, StoppingMinPatients, ANY, ANY, ANY, Data-method

Stop based on minimum number of patients

# Description

Stop based on minimum number of patients

# Usage

```
## S4 method for signature 'StoppingMinPatients, ANY, ANY, ANY, Data'
stopTrial(stopping, dose,
   samples, model, data, ...)
```

# **Arguments**

stopping The rule, an object of class Stopping dose the recommended next best dose

samples the Samples object

model The model input, an object of class Model data The data input, an object of class Data

 $stop Trial, Stopping \texttt{MTD} distribution, numeric, Samples, \texttt{Model}, \texttt{ANY-method} \\ Stop\ based\ on\ MTD\ distribution$ 

# Description

Stop based on MTD distribution

# Usage

```
## S4 method for signature 'StoppingMTDdistribution,numeric,Samples,Model,ANY'
stopTrial(stopping,
  dose, samples, model, data, ...)
```

# Arguments

stopping The rule, an object of class Stopping dose the recommended next best dose

samples the Samples object

model The model input, an object of class Model data The data input, an object of class Data

... additional arguments

stopTrial,StoppingPatientsNearDose,numeric,ANY,ANY,Data-method

Stop based on number of patients near to next best dose

## **Description**

Stop based on number of patients near to next best dose

#### Usage

```
## S4 method for signature 'StoppingPatientsNearDose,numeric,ANY,ANY,Data'
stopTrial(stopping,
  dose, samples, model, data, ...)
```

# **Arguments**

stopping The rule, an object of class Stopping dose the recommended next best dose

samples the Samples object

model The model input, an object of class Model data The data input, an object of class Data

#### **Description**

Stop based on probability of targeting biomarker

#### Usage

```
## S4 method for signature
## 'StoppingTargetBiomarker,numeric,Samples,DualEndpoint,ANY'
stopTrial(stopping,
  dose, samples, model, data, ...)
```

#### **Arguments**

stopping The rule, an object of class Stopping dose the recommended next best dose

samples the Samples object

model The model input, an object of class Model data The data input, an object of class Data

... additional arguments

# **Description**

Stop based on probability of target tox interval

#### Usage

```
## S4 method for signature 'StoppingTargetProb,numeric,Samples,Model,ANY'
stopTrial(stopping,
  dose, samples, model, data, ...)
```

#### **Arguments**

stopping The rule, an object of class Stopping dose the recommended next best dose

samples the Samples object

model The model input, an object of class Model data The data input, an object of class Data

```
summary, Simulations-method
```

Summarize the simulations, relative to a given truth

# **Description**

Summarize the simulations, relative to a given truth

# Usage

```
## S4 method for signature 'Simulations'
summary(object, truth, target = c(0.2, 0.35), \ldots)
```

#### **Arguments**

the Simulations object we want to summarize object a function which takes as input a dose (vector) and returns the true probability truth (vector) for toxicity. Additional arguments can be supplied via .... the target toxicity interval (default: 20-35%) used for the computations target

additional arguments affecting the summary produced. . . .

#### Value

an object of class Simulations-summary

update, Data-method

Update method for the "Data" class

# **Description**

Add new data to the Data object

# Usage

```
## S4 method for signature 'Data'
update(object, x, y, ID = (if (length(object@ID))
 max(object@ID) else OL) + seq_along(y), ...)
```

#### **Arguments**

```
the old Data object
object
                   the dose level (one level only!)
Х
                   the DLT vector (0/1 vector), for all patients in this cohort
У
ID
                   the patient IDs
```

Additional arguments to the call, or arguments with changed values. Use name = NULL . . . to remove the argument name.

#### Value

the new Data object

```
update, DataParts-method
```

Update method for the "DataParts" class

# Description

Add new data to the DataParts object

# Usage

```
## S4 method for signature 'DataParts'
update(object, x, y, ID = (if (length(object@ID))
  max(object@ID) else OL) + seq_along(y), ...)
```

# Arguments

```
the old DataParts object

x the dose level (one level only!)

y the DLT vector (0/1 vector), for all patients in this cohort

ID the patient IDs

... Additional arguments to the call, or arguments with changed values. Use name = NULL to remove the argument name.
```

# Value

the new DataParts object

```
&, Stopping, Stopping-method
```

The method combining two atomic stopping rules

# **Description**

The method combining two atomic stopping rules

# Usage

```
## S4 method for signature 'Stopping,Stopping'
e1 & e2
```

# Arguments

```
e1 First Stopping object
e2 Second Stopping object
```

#### Value

The StoppingAll object

```
&, Stopping, StoppingAll-method
```

The method combining an atomic and a stopping list

# Description

The method combining an atomic and a stopping list

# Usage

```
## S4 method for signature 'Stopping,StoppingAll'
e1 & e2
```

# **Arguments**

```
e1 Stopping object
e2 StoppingAll object
```

# Value

The modified StoppingAll object

```
&, StoppingAll, Stopping-method
```

The method combining a stopping list and an atomic

# Description

The method combining a stopping list and an atomic

# Usage

```
## S4 method for signature 'StoppingAll,Stopping'
e1 & e2
```

# **Arguments**

```
e1 StoppingAll object
e2 Stopping object
```

# Value

The modified StoppingAll object

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