Package 'OncoBayes2'

December 12, 2019

```
Type Package
Title Bayesian Logistic Regression for Oncology Dose-Escalation Trials
Description Bayesian logistic regression model with optional
      EXchangeability-NonEXchangeability parameter modelling for flexible
      borrowing from historical or concurrent data-sources. The safety model
      can guide dose-escalation decisions for adaptive oncology Phase I
      dose-escalation trials which involve an arbitrary number of
      drugs. Please refer to Neuenschwander et al. (2008)
      <doi:10.1002/sim.3230> and Neuenschwander et al. (2016)
      <doi:10.1080/19466315.2016.1174149> for details on the methodology.
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```

44

Index

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R topics documented:

bind_rows_0	3
blrm_exnex	3
blrm_formula_linear	0
blrm_trial	1
codata_combo2	5
dose_info_combo2	6
drug_info_combo2	7
example-combo2	8
example-combo2_trial	0
example-combo3	2
example-single-agent	5
example_model	7
hist_combo2	8
hist_combo3	9
hist_SA	0
nsamples.blrmfit	2
OncoBayes2	3
posterior_interval.blrmfit	4
posterior_linpred.blrmfit	5
posterior_predict.blrmfit	6
predictive_interval.blrmfit	8
prior_summary.blrmfit	9
summary.blrmfit	0
summary.blrm_trial	1
update.blrmfit	2
update.blrm_trial	3

bind_rows_0

bind_rows_0

Bind rows of multiple data frames with zero fill

Description

A version of bind_rows out of dplyr that fills non-common columns with zeroes instead of NA. Gives an error if any of the input data contains NA already.

Usage

```
bind_rows_0(...)
```

Arguments

... Data frames to combine, passed into bind_rows (see dplyr documentation)

Examples

```
## Setting up dummy sampling for fast execution of example
## Please use 4 chains and 100x more warmup & iter in practice
.user_mc_options <- options(OncoBayes2.MC.warmup=10, OncoBayes2.MC.iter=20, OncoBayes2.MC.chains=1)
library(tibble)

dose_info_A <- tibble(
    group_id = "hist_A",
    drug_A = 1
)

dose_info_B <- tibble(
    group_id = "hist_B",
    drug_B = 100 * (1:2)
)

bind_rows_0(dose_info_A, dose_info_B)

## Recover user set sampling defaults
options(.user_mc_options)</pre>
```

blrm_exnex

Bayesian Logistic Regression Model for N-compounds with EXNEX

Description

Bayesian Logistic Regression Model (BLRM) for N compounds using EXchangability and NonEXchangability (EXNEX) modeling.

Usage

```
blrm_exnex(formula, data, prior_EX_mu_mean_comp, prior_EX_mu_sd_comp,
  prior_EX_tau_mean_comp, prior_EX_tau_sd_comp, prior_EX_corr_eta_comp,
  prior_EX_mu_mean_inter, prior_EX_mu_sd_inter, prior_EX_tau_mean_inter,
  prior_EX_tau_sd_inter, prior_EX_corr_eta_inter, prior_is_EXNEX_inter,
 prior_is_EXNEX_comp, prior_EX_prob_comp, prior_EX_prob_inter,
  prior_NEX_mu_mean_comp, prior_NEX_mu_sd_comp, prior_NEX_mu_mean_inter,
  prior_NEX_mu_sd_inter, prior_tau_dist,
  iter = getOption("OncoBayes2.MC.iter", 2000),
 warmup = getOption("OncoBayes2.MC.warmup", 1000),
  thin = getOption("OncoBayes2.MC.thin", 1),
  init = getOption("OncoBayes2.MC.init", 0.5),
  chains = getOption("OncoBayes2.MC.chains", 4),
  cores = getOption("mc.cores", 1L),
  control = getOption("OncoBayes2.MC.control", list()),
  prior_PD = FALSE, verbose = FALSE)
## S3 method for class 'blrmfit'
print(x, ..., prob = 0.95, digits = 2)
```

Arguments

formula

the model formula describing the linear predictors of the model. The lhs of the formula is a two-column matrix which are the number of occured events and the number of times no event occured. The rhs of the formula defines the linear predictors for the marginal models for each drug component, then the interaction model and at last the grouping and optional stratum factors of the models. These elements of the formula are separated by a vertical bar. The marginal models must follow a intercept and slope form while the interaction model must not include an interaction term. See the examples below for an example instantiation.

data

optional data frame containing the variables of the model. If not found in data, the variables are taken from environment(formula).

```
prior_EX_mu_mean_comp, prior_EX_mu_sd_comp
```

Mean and sd for the prior on the mean parameters $\mu_i = (\mu_{\alpha i}, \mu_{\beta i})$ of each component. Two column matrix (intercept, log-slope) with one row per component.

```
prior_EX_tau_mean_comp, prior_EX_tau_sd_comp
```

Prior mean and sd for heterogeniety parameter $\tau_{si}=(\tau_{\alpha si},\tau_{\beta si})$ of each stratum. If no differential discounting is required (i.e. if there is only one stratum s=1), then it is a two-column matrix (intercept, log-slope) with one row per component. Otherwise it is a three-dimensional array whose first dimension indexes the strata, second dimension indexes the components, and third dimension of length two for (intercept, log-slope).

```
prior_EX_corr_eta_comp
```

Prior LKJ correlation parameter for each component given as numeric vector. If missing, then a 1 is assumed corresponding to a marginal uniform prior of the correlation.

prior_EX_mu_mean_inter, prior_EX_mu_sd_inter

Prior mean and sd for population mean parameters $\mu_{\eta k}$ of each interaction parameter. Vector of length equal to the number of interactions.

prior_EX_tau_mean_inter, prior_EX_tau_sd_inter

Prior mean and sd for heterogeniety parameter $\tau_{\eta sk}$ of each stratum. Matrix with one column per interaction and one row per stratum.

prior_EX_corr_eta_inter

Prior LKJ correlation parameter for interaction given as numeric. If missing, then a 1 is assumed corresponding to a marginal uniform prior of the correlations.

prior_is_EXNEX_inter

Defines if non-exchangability is admitted for a given interaction parameter. Logical vector of length equal to the number of interactions. If missing FALSE is assumed for all interactions.

prior_is_EXNEX_comp

Defines if non-exchangability is admitted for a given component. Logical vector of length equal to the number of components. If missing TRUE is assumed for all components.

prior_EX_prob_comp

Prior probability p_{ij} for exchangability of each component per group. Matrix with one column per component and one row per group. Values must lie in [0-1] range.

prior_EX_prob_inter

Prior probability $p_{\eta kj}$ for exchangability of each interaction per group. Matrix with one column per interaction and one row per group. Values must lie in [0-1] range.

prior_NEX_mu_mean_comp, prior_NEX_mu_sd_comp

Prior mean m_{ij} and sd $s_{ij} = \text{diag}(S_{ij})$ of each component for non-exchangable case. Two column matrix (intercept, log-slope) with one row per component. If missing set to the same prior as given for the EX part. It is required that the specification be the same across groups j.

prior_NEX_mu_mean_inter, prior_NEX_mu_sd_inter

Prior mean $m_{\eta kj}$ and sd $s_{\eta kj}$ for each interaction parameter for non-exchangable case. Vector of length equal to the number of interactions. If missing set to the same prior as given for the EX part.

prior_tau_dist Defines the distribution used for heterogeniety parameters. Choices are 0=fixed to it's mean, 1=log-normal, 2=truncated normal.

iter number of iterations (including warmup).

warmup number of warmup iterations.
thin period of saving samples.

init positive number to specify uniform range on unconstrained space for random

initialization. See stan.

chains number of Markov chains.

cores number of cores for parallel sampling of chains.
control additional sampler parameters for NuTS algorithm

prior_PD	Logical flag (defaults to FALSE) indicating if to sample the prior predictive distribution instead of conditioning on the data.
verbose	$\label{logical} Logical flag \ (defaults \ to \ {\tt FALSE}) \ controlling \ if \ additional \ output \ like \ stan \ progress \ is \ reported.$
X	blrmfit object to print
	not used in this function
prob	central probability mass to report, i.e. the quantiles 0.5 -prob/2 and 0.5 +prob/2 are displayed. Multiple central widths can be specified.
digits	number of digits to show

Details

blrm_exnex is a flexible function for Bayesian meta-analytic modeling of binomial count data. In particular, it is designed to model counts of the number of observed dose limiting toxicities (DLTs) by dose, for guiding dose-escalation studies in Oncology. To accommodate dose escalation over more than one agent, the dose may consist of combinations of study drugs, with any number of treatment components.

In the simplest case, the aim is to model the probability π that a patient experiences a DLT, by complementing the binomial likelihood with a monotone logistic regression

$$\operatorname{logit} \pi(d) = \operatorname{log} \alpha + \beta t(d),$$

where $\beta > 0$. Most typically, d represents the dose, and t(d) is an appropriate transformation, such as $t(d) = \log(d/d^*)$. A joint prior on $\theta = (\log \alpha, \log \beta)$ completes the model and ensures monotonicity $\beta > 0$.

Many extensions are possible. The function supports general combination regimens, and also provides framework for Bayesian meta-analysis of dose-toxicity data from multiple historical and concurrent sources.

For an example of a single-agent trial refer to example-single-agent.

Value

The function returns a S3 object of type blrmfit.

Methods (by generic)

• print: print function.

Combination of two treatments

For a combination of two treatment components, the basic modeling framework is that the DLT rate $\pi(d_1,d_2)$ is comprised of (1) a "no-interaction" baseline model $\tilde{\pi}(d_1,d_2)$ driven by the single-agent toxicity of each component, and (2) optional interaction terms $\gamma(d_1,d_2)$ representing synergy or antagonism between the drugs. On the log-odds scale,

$$logit \pi(d_1, d_2) = logit \tilde{\pi}(d_1, d_2) + \eta \gamma(d_1, d_2).$$

The "no interaction" part $\tilde{\pi}(d_1, d_2)$ represents the probability of a DLT triggered by either treatment component acting *independently*. That is,

$$\tilde{\pi}(d_1, d_2) = 1 - (1 - \pi_1(d_1))(1 - \pi_2(d_2)).$$

In simple terms, P(no DLT for combination) = P(no DLT for drug 1) * P(no DLT from drug 2). To complete this part, the treatment components can then be modeled with monotone logistic regressions as before.

$$logit \pi_i(d_i) = log \alpha_i + \beta_i t(d_i)$$

where $t(d_i)$ is a monotone transformation of the doses, such as $t(d_i) = \log(d_i/d_i^*)$.

The inclusion of an interaction term $\gamma(d_1, d_2)$ allows DLT rates above or below the "no-interaction" rate. The magnitude of the interaction term may also be made dependent on the doses (or other covariates) through regression. As an example, one could let

$$\gamma(d_1, d_2) = \frac{d_1}{d_1^*} \frac{d_1}{d_2^*}.$$

The specific functional form is specified in the usual notation for a design matrix. The interaction model must respect the constraint that whenever any dose approaches zero, then the interaction term must vanish as well. Therefore, the interaction model must not include an intercept term which would violate this consistency requirement. A dual combination example can be found in example-combo2.

General combinations

The model is extended to general combination treatments consisting of N components by expressing the probability π on the logit scale as

$$\operatorname{logit} \pi(d_1, \dots, d_N) = \operatorname{logit} \left(1 - \prod_{i=1}^N (1 - \pi_i(d_i)) \right) + \sum_{k=1}^K \eta_k \, \gamma_k(d_1, \dots, d_N),$$

Multiple drug-drug interactions among the N components are now possible, and are represented through the K interaction terms γ_k .

Regression models can be again be specified for each π_i and γ_k , such as

$$\operatorname{logit} \pi_i(d_i) = \log \alpha_i + \beta_i t(d_i)$$

Interactions for some subset $I(k) \subset \{1, ..., N\}$ of the treatment components can be modeled with regression as well, for example on products of doses,

$$\gamma_k(d_1,\ldots,d_N) = \prod_{i\in I(k)} \frac{d_i}{d_i^*}.$$

For example, $I(k) = \{1, 2, 3\}$ results in the three-way interaction term

$$\frac{d_1}{d_1^*} \frac{d_2}{d_2^*} \frac{d_3}{d_3^*}$$

for drugs 1, 2, and 3.

For a triple combination example please refer to example-combo3.

Meta-analytic framework

Information on the toxicity of a drug may be available from multiple studies or sources. Furthermore, one may wish to stratify observations within a single study (for example into groups of patients corresponding to different geographic regions, or multiple dosing dose_info corresponding to different schedules).

blrm_exnex provides tools for robust Bayesian hierarchical modeling to jointly model data from multiple sources. An additional index $j=1,\ldots,J$ on the parameters and observations denotes the J groups. The resulting model allows the DLT rate to differ across the groups. The general N-component model becomes

$$\log \operatorname{it} \pi_j(d_1, \dots, d_N) = \operatorname{logit} \left(1 - \prod_{i=1}^N (1 - \pi_{ij}(d_i)) \right) + \sum_{k=1}^K \eta_{kj} \, \gamma_k(d_1, \dots, d_N),$$

for groups $j=1,\ldots,J$. The component toxicities π_{ij} and interaction terms γ_k are modelled, as before, through regression. For example, π_{ij} could be a logistic regression on $t(d_i) = \log(d_i/d_i^*)$ with intercept and log-slope θ_{ij} , and γ_k regressed with coefficient η_{kj} on a product $\prod_{i\in I(k)}(d_i/d_i^*)$ for some subset I(k) of components.

Thus, for $j=1,\ldots,J$, we now have group-specific parameters $\boldsymbol{\theta}_{ij}=(\log \alpha_{ij},\log \beta_{ij})$ and $\boldsymbol{\nu}_j=(\eta_{1j},\ldots,\eta_{Kj})$ for each component $i=1,\ldots,N$ and interaction $k=1,\ldots,K$.

The structure of the prior on $(\theta_{i1}, \dots, \theta_{iJ})$ and (ν_1, \dots, ν_J) determines how much information will be shared across groups j. Several modeling choices are available in the function.

• EX (Full exchangeability): One can assume the parameters are conditionally exchangeable given hyperparameters

$$\theta_{ij} \sim N(\mu_{\theta i}, \Sigma_{\theta i}),$$

independently across groups $j=1,\ldots,J$ and treatment components $i=1,\ldots,N$. The covariance matrix $\Sigma_{\theta i}$ captures the patterns of cross-group heterogeneity, and is parametrized with standard deviations $\tau_{\theta i}=(\tau_{\alpha i},\tau_{\beta i})$ and the correlation ρ_i . Similarly for the interactions, the fully-exchangeable model is

$$oldsymbol{
u}_j \sim \mathrm{N}ig(oldsymbol{\mu}_{oldsymbol{
u}}, oldsymbol{\Sigma}_{oldsymbol{
u}}ig)$$

for groups $j=1,\ldots,J$ and interactions $k=1,\ldots,K$, and the prior on the covariance matrix Σ_{ν} captures the amount of heterogeneity expected in the interaction terms a-priori. The covariance is again parametrized with standard deviations $(\tau_{\eta 1},\ldots,\tau_{\eta K})$ and its correlation matrix

• Differential discounting: For one or more of the groups $j=1,\ldots,J$, larger deviations of θ_{ij} may be expected from the mean μ_i , or of the interactions η_{kj} from the mean $\mu_{\eta,k}$. Such differential heterogeneity can be modeled by mapping the groups $j=1,\ldots,J$ to strata through $s_j \in \{1,\ldots,S\}$, and modifying the model specification to

$$oldsymbol{ heta}_{ij} \sim \mathrm{N}ig(oldsymbol{\mu}_{oldsymbol{ heta}i}, oldsymbol{\Sigma}_{oldsymbol{ heta}ij}ig),$$

where

$$\boldsymbol{\Sigma}_{\boldsymbol{\theta}ij} = \left(\begin{array}{cc} \tau_{\alpha s_j i}^2 & \rho_i \tau_{\alpha s_j i} \tau_{\beta s_j i} \\ \rho_i \tau_{\alpha s_j i} \tau_{\beta s_j i} & \tau_{\beta s_j i}^2 \end{array} \right).$$

For the interactions, the model becomes

$$\nu_j \sim \mathrm{N}(\mu_{\nu}, \Sigma_{\nu j}),$$

where the covariance matrix $\Sigma_{\nu j}$ is modelled as stratum specific standard deviations $(\tau_{\eta 1s_j},\ldots,\tau_{\eta Ks_j})$ and a stratum independent correlation matrix. Each stratum $s=1,\ldots,S$ then corresponds to its own set of standard deviations τ leading to different discounting per stratum. Independent priors are specified for the component parameters $\tau_{\alpha si}$ and $\tau_{\beta si}$ and for the interaction parameters $\tau_{\eta sk}$ for each stratum $s=1,\ldots,S$. Inference for strata s where the prior is centered on larger values of the τ parameters will exhibit less shrinkage towards the the means, $\mu_{\theta i}$ and μ_{ν} respectively.

• EXNEX (Partial exchangeability): Another mechansim for increasing robustness is to introduce mixture priors for the group-specific parameters, where one mixture component is shared across groups, and the other is group-specific. The result, known as an EXchangeable-NonEXchangeable (EXNEX) type prior, has a form

$$\boldsymbol{\theta}_{ij} \sim p_{\boldsymbol{\theta}ij} \, \mathrm{N}(\boldsymbol{\mu}_{\boldsymbol{\theta}i}, \boldsymbol{\Sigma}_{\boldsymbol{\theta}i}) + (1 - p_{\boldsymbol{\theta}ij}) \, \mathrm{N}(\boldsymbol{m}_{\boldsymbol{\theta}ij}, \boldsymbol{S}_{\boldsymbol{\theta}ij})$$

when applied to the treatment-component parameters, and

$$\nu_{kj} \sim p_{\nu_{kj}} \operatorname{N}(\mu_{\nu}, \Sigma_{\nu})_k + (1 - p_{\nu_{kj}}) \operatorname{N}(m_{\nu_{kj}}, s_{\nu_{kj}}^2)$$

when applied to the interaction parameters. The exchangeability weights $p_{\theta ij}$ and $p_{\nu_{kj}}$ are fixed constants in the interval [0,1] that control the degree to which inference for group j is informed by the exchangeable mixture components. Larger values for the weights correspond to greater exchange of information, while smaller values increase robustness in case of outlying observations in individual groups j.

References

Neuenschwander, B., Roychoudhury, S., & Schmidli, H. (2016). On the use of co-data in clinical trials. *Statistics in Biopharmaceutical Research*, 8(3), 345-354.

Neuenschwander, B., Wandel, S., Roychoudhury, S., & Bailey, S. (2016). Robust exchangeability designs for early phase clinical trials with multiple strata. *Pharmaceutical statistics*, 15(2), 123-134.

Neuenschwander, B., Branson, M., & Gsponer, T. (2008). Critical aspects of the Bayesian approach to phase I cancer trials. *Statistics in medicine*, 27(13), 2420-2439.

Neuenschwander, B., Matano, A., Tang, Z., Roychoudhury, S., Wandel, S. Bailey, Stuart. (2014). A Bayesian Industry Approach to Phase I Combination Trials in Oncology. In *Statistical methods in drug combination studies* (Vol. 69). CRC Press.

```
## Setting up dummy sampling for fast execution of example
## Please use 4 chains and 100x more warmup & iter in practice
.user_mc_options <- options(OncoBayes2.MC.warmup=10, OncoBayes2.MC.iter=20, OncoBayes2.MC.chains=1)
# fit an example model. See documentation for "combo3" example
example_model("combo3")</pre>
```

10 blrm_formula_linear

```
# print a summary of the prior
prior_summary(blrmfit, digits = 3)
# print a summary of the posterior (model parameters)
print(blrmfit)
# summary of posterior for DLT rate by dose for observed covariate levels
summ <- summary(blrmfit, interval_prob = c(0, 0.16, 0.33, 1))
print(cbind(hist_combo3, summ))
# summary of posterior for DLT rate by dose for new set of covariate levels
newdata <- expand.grid(</pre>
 stratum = "BID", group_id = "Combo",
 drug_A = 400, drug_B = 800, drug_C = c(320, 400, 600, 800),
 stringsAsFactors = FALSE
summ_pred <- summary(blrmfit, newdata = newdata, interval_prob = c(0, 0.16, 0.33, 1))
print(cbind(newdata, summ_pred))
# update the model after observing additional data
newdata$num_patients <- rep(3, nrow(newdata))</pre>
newdatanum_toxicities <- c(0, 1, 2, 2)
library(dplyr)
blrmfit_new <- update(blrmfit,</pre>
                      data = rbind(hist_combo3, newdata) %>%
                               arrange(stratum, group_id))
# updated posterior summary
summ_upd <- summary(blrmfit_new, newdata = newdata, interval_prob = c(0, 0.16, 0.33, 1))
print(cbind(newdata, summ_upd))
## Recover user set sampling defaults
options(.user_mc_options)
```

blrm_formula_linear

Build a BLRM formula with linear interaction term in logit-space

Description

blrm_formula_linear is a convenience function for generating a formula for blrm_trial and blrm_exnex.

Usage

```
blrm_formula_linear(ref_doses, max_interaction_level = 2)
```

Arguments

ref_doses Numeric vector of reference doses with names corresponding to drug names max_interaction_level

Highest interaction order to consider [1 - Inf]. Default: 2

Value

The function returns an object of class blrm_formula.

Examples

```
ref_doses <- c(drug_A=10, drug_B=20)
# can be used with blrm_trial
blrm_formula_linear(ref_doses)</pre>
```

blrm_trial

Dose-Escalation Trials guided by Bayesian Logistic Regression Model

Description

blrm_trial facilitates the conduct of dose escalation studies guided by Bayesian Logistic Regression Models (BLRM). While the blrm_exnex only fits the BLRM model to data, the blrm_trial function standardizes the specification of the entire trial design and provides various standardized functions for trial data accrual and derivation of model summaries needed for dose-escalation decisions.

Usage

```
blrm_trial(data, dose_info, drug_info, simplified_prior = FALSE,
    EXNEX_comp = TRUE, EX_prob_comp_hist = 1, EX_prob_comp_new = 0.8,
    EXNEX_inter = FALSE, EX_prob_inter = 0.8,
    formula_generator = blrm_formula_linear, interval_prob = c(0, 0.16,
    0.33, 1), interval_max_mass = c(prob_underdose = 1, prob_target = 1,
    prob_overdose = 0.25), ...)

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

Arguments

data dose-toxicity data available at design stage of trial dose_info specification of the dose levels as planned for the ongoing trial arms. drug_info specification of drugs used in trial arms. simplified_prior

logical (defaults to FALSE) indicating whether a simplified prior should be employed based on the reference_p_dlt values provided in drug_info. **Warning:** The simplified prior will change between releases. Please read instructions below in the respective section for the simplified prior.

EXNEX_comp logical (default to TRUE) indicating whether EXchangeable-NonEXchangeable

priors should be employed for all component parameters

EX_prob_comp_hist

prior weight ([0, 1], default to 1) on exchangeability for the component parameters in groups representing historical data

EX_prob_comp_new

prior weight ([0,1], default to 0.8) on exchangeability for the component parameters in groups representing new or concurrent data

EXNEX_inter logical (default to FALSE) indicating whether EXchangeable-NonEXchangeable priors should be employed for all interaction parameters

EX_prob_inter prior weight ([0,1], defaults to 0.8) on exchangeability for the interaction parameters

formula_generator

formula generation function (see for example blrm_formula_linear). The formula generator defines the employed interaction model.

interval_prob defines the interval probabilities reported in the standard outputs. Defaults to c(0,0.16,0.33,1).

interval_max_mass

named vector defining for each interval of the interval_prob vector a maxmimal admissable probability mass for a given dose level. Whenever the posterior probability mass in a given interval exceeds the threshold, then the Escalation With Overdose Control (EWOC) criterion is considered to be not fullfilled. Dose levels not fullfilling EWOC are ineligible for the next cohort of patients. The default restricts the overdose probability to less than 0.25.

Additional arguments are forwarded to blrm_exnex, i.e. for the purpose of prior specification.

x blrm_trial object to print

Details

blrm_trial constructs an object of class blrm_trial which stores the compelte information about the study design of a dose-escalation trial. The study design is defined through the data sets (see sections below for a definition of the columns):

data (historical data) The data argument defines available dose-toxicity data at the design stage of the trial. Together with the prior of model (without any data) this defines the prior used for the trial conduct.

dose_info Definition of the pre-specified dose levels explored in the ongoing trial arms. Thus, all dose-toxcitiy trial data added to the object is expected correspond to one of the dose levels in the pre-defined set of dose_info.

drug_info Determines the drugs used in the trial, their units, reference dose level and optionally defines the expected probability for a toxicity at the reference dose.

Once the blrm_trial object is setup the complete trial design is specified and the model is fitted to the given data. This allows evaluation of the pre-specified dose levels of the trial design wrt. to safety, i.e. whether the starting dose of the trial fullfills the escalate with overdose criterion (EWOC) condition.

The blrm_trial trial can also be constructed in a 2-step process which allows for a more convenient specification of the prior since meta data like number of drugs and the like can be used. See the example section for details.

After setup of the initial blrm_trial object additional data is added through the use of the update method which has a add_data argument intended to add data from the ongoing trial. The summary function finally allows to extract various model summaries. In particular, the EWOC criterion can be calculated for the pre-defined dose levels of the trial.

Value

The function returns an object of class blrm_trial.

Methods (by generic)

• print: print function.

Simplified prior

As a convenience for the user, a simplified prior can be specifed whenever the reference_p_dlt column is present in the drug_info data set. However, the user is **warned** that the simplified prior will change in future releases of the package and thus **we strongly discourage the use of the simplified prior for setting up trial designs**. The functionality is intended to provide the user a quick start and as a starting point. The actually instantiated prior can be seen as demonstrated below in the examples.

Input data

The data given to the data argument of blrm_trial is considered as the available at design stage of the trial. The collected input data thus does not necessarily need to have the same dose levels as the pre-specified dose_info for the ongoing trial(s). It's data columns must include, but are not limited to:

```
group_id study
```

stratum_id optional, only required for differential discounting of groups

num_patients number of patients

num_toxicities number of toxicities

drug_A Columns for the dose of each treatment component, with column names matching the drug_name values specified in the drug_info argument of blrm_trial

Drug info data

The drug information data-set defines drug properties. The fields included are:

drug_name name of drug which is also used as column name for the dose

dose ref reference dose

dose_unit units used for drug amounts

reference_p_dlt optional; if provided, allows setup of a simplified prior

Dose info data

The drug_info data-set pre-specifies the dose levels of the ongoing trial. Thus, all data added to the blrm_trial through the update command must be consistent with the pre-defined dose levels as no other than those pre-specified ones can be explored in an ongoing trial.

dose_id optional column which assigns a unique id to each group_id/dose combination. If not specified the column is internally generated.

```
group_id study
```

drug_A Columns for the dose of each treatment component, with column names matching the drug_name values specified in the drug_info argument of blrm_trial

References

Babb, J., Rogatko, A., & Zacks, S. (1998). Cancer phase I clinical trials: efficient dose escalation with overdose control. *Statistics in medicine*, 17(10), 1103-1120.

Neuenschwander, B., Roychoudhury, S., & Schmidli, H. (2016). On the use of co-data in clinical trials. *Statistics in Biopharmaceutical Research*, 8(3), 345-354.

See Also

Other blrm_trial combo2 example: dose_info_combo2, drug_info_combo2, example-combo2_trial

```
## Setting up dummy sampling for fast execution of example
## Please use 4 chains and 100x more warmup & iter in practice
.user_mc_options <- options(OncoBayes2.MC.warmup=10, OncoBayes2.MC.iter=20, OncoBayes2.MC.chains=1)
# construct initial blrm_trial object from built-in example datasets
combo2_trial_setup <- blrm_trial(</pre>
 data = hist_combo2,
 dose_info = dose_info_combo2,
 drug_info = drug_info_combo2,
 simplified_prior = TRUE
)
# extract blrm_call to see setup of the prior as passed to blrm_exnex
summary(combo2_trial_setup, "blrm_exnex_call")
# Warning: The simplified prior will change between releases!
# please refer to the combo2_trial example for a complete
# example. You can obtain this example with
# ?example-combo2_trial
# or by running
# example_model("combo2_trial")
## Recover user set sampling defaults
options(.user_mc_options)
```

codata_combo2

codata_combo2

Dataset: historical and concurrent data on a two-way combination

Description

One of two datasets from the application described in Neuenschwander et al (2016). In the study trial_AB, the risk of DLT was studied as a function of dose for two drugs, drug A and drug B. Historical information on the toxicity profiles of these two drugs is available from single agent trials trial_A and trial_B. Another study IIT was run concurrently to trial_AB, and studies the same combination. A second dataset hist_combo2 is available from this example, which includes only the data from the single agent studies, prior to the initiation of trial_AB and IIT.

Usage

```
codata_combo2
```

Format

A data frame with 20 rows and 5 variables:

```
group_id study
drug_A dose of Drug A
drug_B dose of Drug B
num_patients number of patients
num_toxicities number of DLTs
cohort_time cohort number of patients
```

References

Neuenschwander, B., Roychoudhury, S., & Schmidli, H. (2016). On the use of co-data in clinical trials. *Statistics in Biopharmaceutical Research*, 8(3), 345-354.

```
## Setting up dummy sampling for fast execution of example
## Please use 4 chains and 100x more warmup & iter in practice
.user_mc_options <- options(OncoBayes2.MC.warmup=10, OncoBayes2.MC.iter=20, OncoBayes2.MC.chains=1)
library(RBesT)
dref <- c(300, 960)

num_comp <- 2 # two investigational drugs
num_inter <- 1 # one drug-drug interaction needs to be modeled
num_groups <- nlevels(codata_combo2$group_id) # no stratification needed
num_strata <- 1 # no stratification needed
blrmfit <- blrm_exnex(</pre>
```

16 dose_info_combo2

```
cbind(num_toxicities, num_patients - num_toxicities) ~
     1 + I(log(drug_A / dref[1])) |
     1 + I(log(drug_B / dref[2])) |
     0 + I(drug_A/dref[1] *drug_B/dref[2]) |
     group_id,
 data = codata_combo2,
 prior_EX_mu_mean_comp = matrix(
    c(logit(0.1), 0, # hyper-mean of (intercept, log-slope) for drug A
      logit(0.1), 0), # hyper-mean of (intercept, log-slope) for drug B
      nrow = num_comp,
      ncol = 2,
      byrow = TRUE
     ),
 prior_EX_mu_sd_comp = matrix(
    c(3.33, 1, # hyper-sd of mean mu for (intercept, log-slope) for drug A
      3.33, 1), # hyper-sd of mean mu for (intercept, log-slope) for drug B
      nrow = num_comp,
      ncol = 2,
      byrow = TRUE
     ),
 prior_EX_tau_mean_comp = matrix(
    c(\log(0.25), \log(0.125),
      log(0.25), log(0.125)),
      nrow = num_comp,
      ncol = 2,
      byrow = TRUE
     ),
 prior_EX_tau_sd_comp = matrix(
    c(log(4) / 1.96, log(4) / 1.96,
      log(4) / 1.96, log(4) / 1.96),
      nrow = num_comp,
      ncol = 2,
      byrow = TRUE
     ),
 prior_EX_mu_mean_inter = 0,
 prior_EX_mu_sd_inter = 1.121,
 prior_EX_tau_mean_inter = matrix(log(0.125), nrow = num_inter, ncol = num_strata),
 prior_EX_tau_sd_inter = matrix(log(4) / 1.96, nrow = num_inter, ncol = num_strata),
 prior_is_EXNEX_comp = rep(FALSE, num_comp),
 prior_is_EXNEX_inter = rep(FALSE, num_inter),
 prior_EX_prob_comp = matrix(1, nrow = num_groups, ncol = num_comp),
 prior_EX_prob_inter = matrix(1, nrow = num_groups, ncol = num_inter),
 prior_tau_dist = 1
)
## Recover user set sampling defaults
options(.user_mc_options)
```

drug_info_combo2

Description

The data set defines all pre-defined dose-levels which can be explored in the dual-agent example trial

Usage

```
dose_info_combo2
```

Format

An object of class tbl_df (inherits from tbl, data.frame) with 33 rows and 4 columns.

Details

```
group_id studydrug_A drug A dose amountdrug_B drug B dose amountdose_id unique id of record
```

See Also

Other blrm_trial combo2 example: blrm_trial, drug_info_combo2, example-combo2_trial

drug_info_combo2

Dataset: drug information for a dual-agent combination study

Description

Data set describing the two drugs involved in the example for a dual-agent combination study.

Usage

```
drug_info_combo2
```

Format

A tibble with 2 rows (one per durg) and 4 columns:

```
drug_name name of drug
dose_ref reference dose
dose_unit units used for drug amounts
reference_p_dlt a-priori probability for a DLT at the reference dose
```

See Also

Other blrm_trial combo2 example: blrm_trial, dose_info_combo2, example-combo2_trial

18 example-combo2

example-combo2

Two-drug combination example

Description

Example using a combination of two experimental drugs.

Details

The following example is described in the reference Neuenschwander, B. et al (2016). The data are described in the help page for codata_combo2. In the study trial_AB, the risk of DLT was studied as a function of dose for two drugs, drug A and drug B. Historical information on the toxicity profiles of these two drugs was available from single agent trials trial_A and trial_B. Another study IIT was run concurrently to trial_AB, and studies the same combination.

The model described in Neuenschwander, et al (2016) is adapted as follows. For groups $j = 1, \ldots, 4$ representing each of the four sources of data mentioned above,

$$\operatorname{logit} \pi_{1j}(d_1) = \log \, \alpha_{1j} + \beta_{1j} \, \log \left(\frac{d_1}{d_1^*} \right),$$

and

$$\operatorname{logit} \pi_{2j}(d_2) = \log \, \alpha_{2j} + \beta_{2j} \, \log \left(\frac{d_2}{d_2^*}\right),$$

are logistic regressions for the single-agent toxicity of drugs A and B, respectively, when administered in group j. Conditional on the regression parameters $\theta_{1j} = (\log \alpha_{1j}, \log \beta_{1j})$ and $\theta_{2j} = (\log \alpha_{2j}, \log \beta_{2j})$, the toxicity $\pi_j(d_1, d_2)$ for the combination is modeled as the "no-interaction" DLT rate,

$$\tilde{\pi}_i(d_1, d_2) = 1 - (1 - \pi_{1i}(d_1))(1 - \pi_{2i}(d_2))$$

with a single interaction term added on the log odds scale,

$$\operatorname{logit} \pi_j(d_1, d_2) = \operatorname{logit} \tilde{\pi}_j(d_1, d_2) + \eta_j \frac{d_1}{d_1^*} \frac{d_2}{d_2^*}.$$

A hierarchical model across the four groups j allows dose-toxicity information to be shared through common hyperparameters.

For the component parameters θ_{ij} ,

$$\theta_{ij} \sim \text{BVN}(\boldsymbol{\mu}_i, \boldsymbol{\Sigma}_i).$$

For the mean, a further prior is specified as

$$\boldsymbol{\mu}_i = (\mu_{\alpha i}, \mu_{\beta i}) \sim \text{BVN}(\boldsymbol{m}_i, \boldsymbol{S}_i),$$

with $m_i = (\text{logit } 0.1, \log 1)$ and $S_i = \text{diag}(3.33^2, 1^2)$ for each i = 1, 2. For the standard deviations and correlation parameters in the covariance matrix,

$$oldsymbol{\Sigma}_i = \left(egin{array}{cc} au_{lpha i}^2 &
ho_i au_{lpha i} au_{eta i} \
ho_i au_{lpha i} au_{eta i} & au_{eta i}^2 \end{array}
ight),$$

example-combo2 19

```
the specified priors are \tau_{\alpha i} \sim \text{Log-Normal}(\log 0.25, ((\log 4)/1.96)^2),

\tau_{\beta i} \sim \text{Log-Normal}(\log 0.125, ((\log 4)/1.96)^2), \text{ and } \rho_i \sim \text{U}(-1, 1) \text{ for } i = 1, 2.
```

For the interaction parameters η_i in each group, the hierarchical model has

$$\eta_j \sim N(\mu_\eta, \tau_\eta^2),$$

for j = 1, ..., 4, with $\mu_{\eta} \sim N(0, 1.121^2)$ and $\tau_{\eta} \sim \text{Log-Normal}(\log 0.125, ((\log 4)/1.96)^2)$.

Below is the syntax for specifying this fully exchangeable model in blrm_exnex.

References

Neuenschwander, B., Roychoudhury, S., & Schmidli, H. (2016). On the use of co-data in clinical trials. *Statistics in Biopharmaceutical Research*, 8(3), 345-354.

```
## Setting up dummy sampling for fast execution of example
## Please use 4 chains and 100x more warmup & iter in practice
.user_mc_options <- options(OncoBayes2.MC.warmup=10, OncoBayes2.MC.iter=20, OncoBayes2.MC.chains=1)
library(RBesT)
dref <- c(300, 960)
num_comp <- 2 # two investigational drugs</pre>
num_inter <- 1 # one drug-drug interaction needs to be modeled
num_groups <- nlevels(codata_combo2$group_id) # no stratification needed</pre>
num_strata <- 1 # no stratification needed</pre>
blrmfit <- blrm_exnex(</pre>
  cbind(num_toxicities, num_patients - num_toxicities) \sim
      1 + I(log(drug_A / dref[1])) |
      1 + I(log(drug_B / dref[2])) |
      0 + I(drug_A/dref[1] *drug_B/dref[2]) |
      group_id,
  data = codata_combo2,
  prior_EX_mu_mean_comp = matrix(
     c(logit(0.1), 0, # hyper-mean of (intercept, log-slope) for drug A
       logit(0.1), 0), # hyper-mean of (intercept, log-slope) for drug B
       nrow = num_comp,
       ncol = 2,
       byrow = TRUE
      ),
  prior_EX_mu_sd_comp = matrix(
     c(3.33, 1, # hyper-sd of mean mu for (intercept, log-slope) for drug A
       3.33, 1), # hyper-sd of mean mu for (intercept, log-slope) for drug B
       nrow = num_comp,
       ncol = 2,
       byrow = TRUE
      ),
  prior_EX_tau_mean_comp = matrix(
     c(\log(0.25), \log(0.125),
```

```
log(0.25), log(0.125)),
       nrow = num_comp,
       ncol = 2,
       byrow = TRUE
     ),
 prior_EX_tau_sd_comp = matrix(
     c(\log(4) / 1.96, \log(4) / 1.96,
       log(4) / 1.96, log(4) / 1.96),
       nrow = num_comp,
       ncol = 2,
       byrow = TRUE
     ),
 prior_EX_mu_mean_inter = 0,
 prior_EX_mu_sd_inter = 1.121,
 prior_EX_tau_mean_inter = matrix(log(0.125), nrow = num_inter, ncol = num_strata),
 prior_EX_tau_sd_inter = matrix(log(4) / 1.96, nrow = num_inter, ncol = num_strata),
 prior_is_EXNEX_comp = rep(FALSE, num_comp),
 prior_is_EXNEX_inter = rep(FALSE, num_inter),
 prior_EX_prob_comp = matrix(1, nrow = num_groups, ncol = num_comp),
 prior_EX_prob_inter = matrix(1, nrow = num_groups, ncol = num_inter),
 prior_tau_dist = 1
)
## Recover user set sampling defaults
options(.user_mc_options)
```

example-combo2_trial Two-drug combination example using BLRM Trial

Description

Example using blrm_trial to guide the built-in two-drug combination study example.

Details

blrm_trial is used to collect and store all relevant design information for the example. Subsequent use of the update.blrm_trial command allows convenient model fitting via blrm_exnex. The summary.blrm_trial method allows exploration of the design and modeling results.

To run this example, use example_model("combo2_trial"). See example_model.

See Also

Other blrm_trial combo2 example: blrm_trial, dose_info_combo2, drug_info_combo2

```
## Setting up dummy sampling for fast execution of example
## Please use 4 chains and 100x more warmup & iter in practice
.user_mc_options <- options(OncoBayes2.MC.warmup=10, OncoBayes2.MC.iter=20, OncoBayes2.MC.chains=1)</pre>
```

```
library(tibble)
library(dplyr)
library(tidyr)
library(RBesT)
# Combo2 example using blrm_trial functionality
# construct initial blrm_trial object from built-in example datasets
combo2_trial_setup <- blrm_trial(</pre>
  data = hist_combo2,
  dose_info = dose_info_combo2,
  drug_info = drug_info_combo2,
  simplified_prior = FALSE
)
dims <- summary(combo2_trial_setup, "dimensionality")</pre>
# Fit the initial model with the historical data and fully specified prior
combo2_trial_start <- update(</pre>
  combo2_trial_setup,
  prior_EX_mu_mean_comp = matrix(
    c(logit(0.1), 0, \# hyper-mean of (intercept, log-slope) for drug A
      logit(0.1), 0), # hyper-mean of (intercept, log-slope) for drug B
    nrow = dims$num_components,
   ncol = 2,
    byrow = TRUE
  ),
  prior_EX_mu_sd_comp = matrix(
   c(3.33, 1, # hyper-sd of mean mu for (intercept, log-slope) for drug A
      3.33, 1), # hyper-sd of mean mu for (intercept, log-slope) for drug B
    nrow = dims$num_components,
   ncol = 2,
   byrow = TRUE
  ),
  prior_EX_tau_mean_comp = matrix(
   c(\log(0.25), \log(0.125),
      log(0.25), log(0.125)),
    nrow = dims$num_components,
    ncol = 2,
   byrow = TRUE
  ),
  prior_EX_tau_sd_comp = matrix(
   c(\log(4) / 1.96, \log(4) / 1.96,
      log(4) / 1.96, log(4) / 1.96),
    nrow = dims$num_components,
   ncol = 2,
   byrow = TRUE
  ),
  prior_EX_mu_mean_inter = 0,
  prior_EX_mu_sd_inter = 1.121,
  prior_EX_tau_mean_inter = matrix(log(0.125),
                                    nrow = dims$num_interaction_terms,
```

22 example-combo3

```
ncol = dims$num_strata),
 prior_EX_tau_sd_inter = matrix(log(4) / 1.96,
                                 nrow = dims$num_interaction_terms,
                                 ncol = dims$num_strata),
 prior_is_EXNEX_comp = rep(FALSE, dims$num_components),
 prior_is_EXNEX_inter = rep(FALSE, dims$num_interaction_terms),
 prior_EX_prob_comp = matrix(1,
                              nrow = dims$num_groups,
                              ncol = dims$num_components),
 prior_EX_prob_inter = matrix(1,
                               nrow = dims$num_groups,
                               ncol = dims$num_interaction_terms),
 prior_tau_dist = 1
# print summary of prior specification
prior_summary(combo2_trial_start)
# summarize inference at observed dose levels
summary(combo2_trial_start, "data_prediction")
# summarize inference at specified dose levels
summary(combo2_trial_start, "dose_prediction")
# Update again with new data
# using update() with data argument supplied
# dem <- update(combo2_trial_start, data = codata_combo2)</pre>
# alternate way using update() with add_data argument for
# new observations only (those collected after the trial
# design stage).
new_data <- filter(codata_combo2, cohort_time > 0)
combo2_trial <- update(combo2_trial_start, add_data = new_data)</pre>
summary(combo2_trial, "data") # cohort_time is tracked
summary(combo2_trial, "data_prediction")
summary(combo2_trial, "dose_prediction")
rm(dims, new_data)
## Recover user set sampling defaults
options(.user_mc_options)
```

example-combo3 23

Description

Example using a combination of two experimental drugs, with EXNEX and differential discounting.

Details

This dataset involves a hypothetical dose-escalation study of combination therapy with three treatment components. From two previous studies HistAgent1 and HistAgent2, historical data is available on each of the treatments as single-agents, as well as two of the two-way combinations. However, due to a difference in treatment schedule between the Combo study and the historical studies, a stratification (through stratum) is made between the groups to allow differential discounting of the alternate-schedule data. The association is as below.

group_id (j):	stratum (s_j):
Combo (1)	BID (1)
HistAgent1 (2)	QD (2)
HistAgent2 (3)	QD (2)

For additional robustness, EXNEX priors are used for all group-level treatment component and interaction parameters, to limit the amount of borrowing in case of significant heterogeneity across groups.

The complete model is as follows. As a function of doses d_1, d_2, d_3 , the DLT rate in group j is, for j = 1, ..., 3,

$$\operatorname{logit} \pi_j(d_1,d_2,d_3) = \operatorname{logit} \Big(1 - \prod_{i=1}^3 (1 - \pi_{ij}(d_i)) \Big) + \eta_j^{(12)} \frac{d_1}{d_1^*} \frac{d_2}{d_2^*} + \eta_j^{(13)} \frac{d_1}{d_1^*} \frac{d_3}{d_3^*} + \eta_j^{(23)} \frac{d_2}{d_2^*} \frac{d_3}{d_3^*} + \eta_j^{(123)} \frac{d_1}{d_1^*} \frac{d_2}{d_2^*} \frac{d_3}{d_3^*} \Big) + \eta_j^{(12)} \frac{d_1}{d_2^*} \frac{d_2}{d_3^*} \frac{d_3}{d_3^*} + \eta_j^{(12)} \frac{d_1}{d_2^*} \frac{d_2}{d_3^*} \frac{d_3}{d_3^*} + \eta_j^{(12)} \frac{d_1}{d_2^*} \frac{d_2}{d_3^*} \frac{d_3}{d_3^*} \Big) + \eta_j^{(12)} \frac{d_1}{d_2^*} \frac{d_2}{d_3^*} \frac{d_3}{d_3^*} + \eta_j^{(12)} \frac{d_1}{d_2^*} \frac{d_2}{d_3^*} \frac{d_3}{d_3^*} + \eta_j^{(12)} \frac{d_1}{d_2^*} \frac{d_2}{d_3^*} \frac{d_3}{d_3^*} \Big) + \eta_j^{(12)} \frac{d_1}{d_2^*} \frac{d_2}{d_3^*} \frac{d_2}{d_3^*} \frac{d_3}{d_3^*} + \eta_j^{(12)} \frac{d_1}{d_3^*} \frac{d_2}{d_3^*} \frac{d_3}{d_3^*} \frac{d_3}{d_3^$$

In group j each treatment component i toxicity is modeled with logistic regression,

logit
$$\pi_{ij}(d_i) = \log \alpha_{ij} + \beta_{ij} \log \left(\frac{d_i}{d_i^*}\right)$$
.

The intercept and log-slope parameters $\theta_{ij} = (\log \alpha_{ij}, \log \beta_{ij})$ are are given an EXNEX prior

$$\boldsymbol{\theta}_{ij} \sim p_{ij} \text{BVN}(\boldsymbol{\mu}_i, \boldsymbol{\Sigma}_{ij}) + (1 - p_{ij}) \text{BVN}(\boldsymbol{m}_{ij}, \boldsymbol{S}_{ij}),$$

where the exchangeability weights are all $p_{ij}=0.9$. The NEX parameters are set to $m_{ij}=(\log i (1/3), \log 1)$, $S_{ij}=\operatorname{diag}(2^2,1^2)$ for all components i=1,2,3 and groups j=1,2,3, and the EX parameters are modeled hierarchically. The mean of the exchangeable part has the distribution

$$\boldsymbol{\mu}_i = (\mu_{\alpha i}, \mu_{\beta i}) \sim \text{BVN}(\boldsymbol{m}_i, \boldsymbol{S}_i),$$

with $m_i = (\text{logit}(1/3), \log 1)$ and $S_i = \text{diag}(2^2, 1^2)$ for each component i = 1, 2, 3. For differentially discounting data from each schedule (QD and BID), the covariance parameters for the exchangeable part

$$\Sigma_{ij} = \begin{pmatrix} \tau_{\alpha s_j i}^2 & \rho_i \tau_{\alpha s_j i} \tau_{\beta s_j i} \\ \rho_i \tau_{\alpha s_j i} \tau_{\beta s_j i} & \tau_{\beta s_j i}^2 \end{pmatrix}.$$

are allowed to vary across groups j depending on their mapping to strata s(j) as described above. For stratum s=1 (BID, which contains only the group j=1 (Combo)), the standard deviations are modeled as

$$\tau_{\alpha 1i} \sim \text{Log-Normal}(\log 0.25, (\log 4/1.96)^2)$$

24 example-combo3

$$\tau_{\beta 1i} \sim \text{Log-Normal}(\log 0.125, (\log 4/1.96)^2).$$

Whereas in stratum s=2 (QD, which contains the historical groups j=2,3 (HistData1, HistData2)), the standard deviations are

$$\tau_{\alpha 2i} \sim \text{Log-Normal}(\log 0.5, (\log 4/1.96)^2)$$

$$\tau_{\beta 2i} \sim \text{Log-Normal}(\log 0.25, (\log 4/1.96)^2).$$

For all interaction parameters $\eta_j^{(12)}$, $\eta_j^{(13)}$, $\eta_j^{(23)}$, and $\eta_j^{(123)}$ (j=1,2,3), the following prior is assumed:

$$\eta_{j}^{(\cdot)} \sim p_{\eta j}^{(\cdot)} \mathrm{N}(\mu_{\eta}^{(\cdot)}, {\tau_{\eta s_{j}}^{(\cdot)}}^{2}) + (1 - p_{\eta j}^{(\cdot)}) \mathrm{N}({m_{\eta j}^{(\cdot)}, s_{\eta j}^{(\cdot)}}^{2}).$$

The exchangeability weights are $p_{\eta j}^{(\cdot)}=0.9$ for all interaction parameters and all groups. Here, for each $\mu_{\eta}^{(12)},\mu_{\eta}^{(13)},\mu_{\eta}^{(23)}$, and $\mu_{\eta}^{(123)}$, we take

$$\mu_{\eta}^{(\cdot)} \sim N(0, 1/2),$$

and for each $au_{\eta s}^{(12)}, au_{\eta s}^{(13)}, au_{\eta s}^{(23)},$ and $au_{\eta s}^{(123)},$

$$\tau_{ns}^{(\cdot)} \sim \text{Log-Normal}(\log(0.25), (\log 2/1.96)^2),$$

for both strata s=1,2. Furthermore, $m_{\eta j}^{(\cdot)}=0$ and $s_{\eta j}^{(\cdot)^2}=1/2$, uniformly across all indices. Below is the syntax for specifying this model in blrm_exnex.

References

Neuenschwander, B., Roychoudhury, S., & Schmidli, H. (2016). On the use of co-data in clinical trials. *Statistics in Biopharmaceutical Research*, 8(3), 345-354.

example-single-agent 25

```
+ I(drug_A/dref[1] * drug_B/dref[2])
                       + I(drug_A/dref[1] * drug_C/dref[3])
                       + I(drug_B/dref[2] * drug_C/dref[3])
                       + I(drug_A/dref[1] * drug_B/dref[2] * drug_C/dref[3]) |
                       stratum/group_id,
                       data=hist_combo3,
                  prior_EX_mu_mean_comp=matrix(c(logit(1/3), 0), nrow=num_comp, ncol=2, TRUE),
                       prior_EX_mu_sd_comp=matrix(c(2, 1), nrow=num_comp, ncol=2, TRUE),
        prior_EX_tau_mean_comp=abind(matrix(log( c(0.25, 0.125)), nrow=num_comp, ncol=2, TRUE),
                                matrix(log(2*c(0.25, 0.125)), nrow=num\_comp, ncol=2, TRUE),
                                         along=0),
                  prior_EX_tau_sd_comp=abind(matrix(log(4)/1.96, nrow=num_comp, ncol=2, TRUE),
                                         matrix(log(4)/1.96, nrow=num_comp, ncol=2, TRUE),
                                                   along=0),
                       prior_EX_mu_mean_inter=rep(0, num_inter),
                       prior_EX_mu_sd_inter=rep(sqrt(2)/2, num_inter),
                  \label{local_prior_EX_tau_mean_inter} prior_{EX\_tau\_mean\_inter=matrix} (log(0.25) \ , nrow=num\_strata, ncol=num\_inter),
                  prior_EX_tau_sd_inter=matrix(log(2)/1.96, nrow=num_strata, ncol=num_inter),
                       prior_EX_prob_comp=matrix(0.9, nrow=num_groups, ncol=num_comp),
                       prior_EX_prob_inter=matrix(0.9, nrow=num_groups, ncol=num_inter),
                     ## by default EXNEX is on for components and off for all interactions
                       prior_tau_dist=1,
                       prior_PD=FALSE
## Recover user set sampling defaults
options(.user_mc_options)
```

example-single-agent Single Agent Example

Description

Example using a single experimental drug.

Details

The single agent example is described in the reference Neuenschwander, B. et al (2008). The data are described in the help page for hist_SA. In this case, the data come from only one study, with the treatment being only single agent. Hence the model specified does not involve a hierarchical prior for the intercept and log-slope parameters. The model described in Neuenschwander, et al (2008) is adapted as follows:

$$\log i \pi(d) = \log \alpha + \beta \log \left(\frac{d}{d^*}\right),$$

where $d^* = 250$, and the prior for $\theta = (\log \alpha, \log \beta)$ is

$$\theta \sim N(m, S)$$
,

and $m = (\text{logit } 0.5, \log 1)$ and $S = \text{diag}(2^2, 1^2)$ are constants.

26 example-single-agent

In the blrm_exnex framework, in which the prior must be specified as a hierarchical model $\theta \sim N(\mu, \Sigma)$ with additional priors on μ and Σ , the simple prior distribution above is accomplished by fixing the diagonal elements τ_{α}^2 and τ_{β}^2 of Σ to zero, and taking

$$\mu \sim N(m, S)$$
.

The arguments prior_tau_dist and prior_EX_tau_mean_comp as specified below ensure that the τ 's are fixed at zero.

References

Neuenschwander, B., Branson, M., & Gsponer, T. (2008). Critical aspects of the Bayesian approach to phase I cancer trials. *Statistics in medicine*, 27(13), 2420-2439.

```
## Setting up dummy sampling for fast execution of example
## Please use 4 chains and 100x more warmup & iter in practice
. user\_mc\_options <- options (OncoBayes2.MC.warmup=10, OncoBayes2.MC.iter=20, OncoBayes2.MC.chains=1) \\
library(RBesT)
## Example from Neuenschwander, B., et al. (2009). Stats in Medicine
num_comp <- 1 # one investigational drug</pre>
num_inter <- 0 # no drug-drug interactions need to be modeled</pre>
num_groups <- nlevels(hist_SA$group_id) # no stratification needed</pre>
num_strata <- 1 # no stratification needed</pre>
dref <- 50
## Since there is no prior information the hierarchical model
## is not used in this example by setting tau to (almost) 0.
blrmfit <- blrm_exnex(</pre>
  cbind(num_toxicities, num_patients - num_toxicities) ~
    1 + log(drug_A / dref) |
    0 |
    group_id,
  data = hist_SA,
  prior_EX_mu_mean_comp = matrix(
    c(logit(1/2), # mean of intercept on logit scale
      log(1)), # mean of log-slope on logit scale
    nrow = num_comp,
    ncol = 2
  prior_EX_mu_sd_comp = matrix(
    c(2, # sd of intercept
      1), # sd of log-slope
    nrow = num_comp,
    ncol = 2
  ),
```

example_model 27

```
## Here we take tau as known and as zero.
 ## This disables the hierarchical prior which is
 ## not required in this example as we analyze a
 ## single trial.
 prior_EX_tau_mean_comp = matrix(
   c(0, 0),
   nrow = num_comp,
   ncol = 2
 ),
 prior_EX_tau_sd_comp = matrix(
   c(1, 1),
   nrow = num_comp,
   ncol = 2
 prior_EX_prob_comp = matrix(1, nrow = num_comp, ncol = 1),
 prior_tau_dist = 0,
 prior_PD = FALSE
)
## Recover user set sampling defaults
options(.user_mc_options)
```

example_model

Runs example models

Description

Runs example models

Usage

```
example_model(topic, envir = parent.frame(), silent = FALSE)
```

Arguments

topic example to run

envir environment which the example is loaded into. Defaults to the caller environ-

ment.

silent logical controlling if execution is run silently (defaults to FALSE)

Value

When topic is not specified a list of all possible topics is return. Whenever a valid topic is specified, the function inserts the example into the environment given and returns (invisibly) the updated environment.

28 hist_combo2

Examples

```
## Setting up dummy sampling for fast execution of example
## Please use 4 chains and 100x more warmup & iter in practice
.user_mc_options <- options(OncoBayes2.MC.warmup=10, OncoBayes2.MC.iter=20, OncoBayes2.MC.chains=1)

## get a list of available examples
example_model()

## run 3 component example
example_model("combo3")

## Recover user set sampling defaults
options(.user_mc_options)</pre>
```

hist_combo2

Dataset: historical data on two single-agents to inform a combination study

Description

One of two datasets from the application described in Neuenschwander et al (2016). The risk of DLT is to be studied as a function of dose for two drugs, drug A and drug B. Historical information on the toxicity profiles of these two drugs is available from single agent trials trial_A and trial_B. A second dataset codata_combo2 is available from this application, which includes additional dosetoxicity data from trial_AB and IIT of the combination of Drugs A and B.

Usage

hist_combo2

Format

A tibble with 11 rows and 5 variables:

```
group_id study
drug_A dose of Drug A
drug_B dose of Drug B
num_patients number of patients
num_toxicities number of DLTs
cohort_time cohort number of patients
```

References

Neuenschwander, B., Roychoudhury, S., & Schmidli, H. (2016). On the use of co-data in clinical trials. *Statistics in Biopharmaceutical Research*, 8(3), 345-354.

hist_combo3 29

hist_combo3

Dataset: historical and concurrent data on a three-way combination

Description

This dataset involves a hypothetical dose-escalation study of combination therapy with three treatment components. From two previous studies <code>HistAgent1</code> and <code>HistAgent2</code>, historical data is available on each of the treatments as single-agents, as well as two of the two-way combinations. However, due to a difference in treatment schedule between the Combo study and the historical studies, a stratification (through <code>stratum</code>) is made between the groups to allow differential discounting of the alternate-dose data.

Usage

hist_combo3

Format

A data frame with 18 rows and 7 variables:

```
group_id study
drug_A dose of Drug A
drug_B dose of Drug B
drug_C dose of Drug C
num_patients number of patients
num_toxicities number of DLTs
stratum stratum for group_id's used for differential discounting
```

30 hist_SA

```
1 + I(log(drug_B/dref[2])) |
                        1 + I(log(drug_C/dref[3])) |
                    + I(drug_A/dref[1] * drug_B/dref[2])
                    + I(drug_A/dref[1] * drug_C/dref[3])
                    + I(drug_B/dref[2] * drug_C/dref[3])
                    + I(drug_A/dref[1] * drug_B/dref[2] * drug_C/dref[3]) |
                    stratum/group_id,
                    data=hist_combo3,
                prior_EX_mu_mean_comp=matrix(c(logit(1/3), 0), nrow=num_comp, ncol=2, TRUE),
                    prior_EX_mu_sd_comp=matrix(c(2, 1), nrow=num_comp, ncol=2, TRUE),
       matrix(log(2*c(0.25, 0.125)), nrow=num_comp, ncol=2, TRUE),
                                    along=0),
                prior_EX_tau_sd_comp=abind(matrix(log(4)/1.96, nrow=num_comp, ncol=2, TRUE),
                                     matrix(log(4)/1.96, nrow=num_comp, ncol=2, TRUE),
                                              along=0),
                    prior_EX_mu_mean_inter=rep(0, num_inter),
                    prior_EX_mu_sd_inter=rep(sqrt(2)/2, num_inter),
                prior_EX_tau_mean_inter=matrix(log(0.25) , nrow=num_strata, ncol=num_inter),
                prior_EX_tau_sd_inter=matrix(log(2)/1.96, nrow=num_strata, ncol=num_inter),
                    prior_EX_prob_comp=matrix(0.9, nrow=num_groups, ncol=num_comp),
                    prior_EX_prob_inter=matrix(0.9, nrow=num_groups, ncol=num_inter),
                  ## by default EXNEX is on for components and off for all interactions
                    prior_tau_dist=1,
                    prior_PD=FALSE
## Recover user set sampling defaults
options(.user_mc_options)
```

hist_SA

Single-agent example

Description

Example data from the application in Neuenschwander, et. al. 2008, from an "open-label, multicenter, non-comparative, dose-escalation cancer trial to characterize the safety, tolerability, and pharmacokinetic profile of a drug and to determine its MTD."

Usage

hist_SA

Format

A data frame with 5 rows and 4 variables:

```
group_id study
drug_A dose
```

hist_SA 31

```
num_patients number of patients
num_toxicities number of events
```

References

Neuenschwander, B., Branson, M., & Gsponer, T. (2008). Critical aspects of the Bayesian approach to phase I cancer trials. *Statistics in medicine*, 27(13), 2420-2439.

```
## Setting up dummy sampling for fast execution of example
## Please use 4 chains and 100x more warmup & iter in practice
.user_mc_options <- options(OncoBayes2.MC.warmup=10, OncoBayes2.MC.iter=20, OncoBayes2.MC.chains=1)
library(RBesT)
## Example from Neuenschwander, B., et al. (2009). Stats in Medicine
num_comp <- 1 # one investigational drug</pre>
num_inter <- 0 # no drug-drug interactions need to be modeled</pre>
num_groups <- nlevels(hist_SA$group_id) # no stratification needed</pre>
num_strata <- 1 # no stratification needed</pre>
dref <- 50
## Since there is no prior information the hierarchical model
## is not used in this example by setting tau to (almost) 0.
blrmfit <- blrm_exnex(</pre>
 cbind(num_toxicities, num_patients - num_toxicities) ~
    1 + \log(\text{drug\_A / dref}) |
   0 |
    group_id,
 data = hist_SA,
 prior_EX_mu_mean_comp = matrix(
    c(logit(1/2), # mean of intercept on logit scale
                # mean of log-slope on logit scale
      log(1)),
   nrow = num_comp,
   ncol = 2
 ),
 prior_EX_mu_sd_comp = matrix(
    c(2, # sd of intercept
     1), # sd of log-slope
   nrow = num_comp,
   ncol = 2
 ## Here we take tau as known and as zero.
 ## This disables the hierarchical prior which is
 ## not required in this example as we analyze a
 ## single trial.
 prior_EX_tau_mean_comp = matrix(
   c(0, 0),
```

32 nsamples.blrmfit

```
nrow = num_comp,
    ncol = 2
),
prior_EX_tau_sd_comp = matrix(
    c(1, 1),
    nrow = num_comp,
    ncol = 2
),
prior_EX_prob_comp = matrix(1, nrow = num_comp, ncol = 1),
prior_tau_dist = 0,
    prior_PD = FALSE
)
## Recover user set sampling defaults
options(.user_mc_options)
```

nsamples.blrmfit

Return the number of posterior samples

Description

Return the number of posterior samples

Usage

```
## S3 method for class 'blrmfit'
nsamples(object, ...)
```

Arguments

```
object fitted model object
... not used in this function
```

```
## Setting up dummy sampling for fast execution of example
## Please use 4 chains and 100x more warmup & iter in practice
.user_mc_options <- options(OncoBayes2.MC.warmup=10, OncoBayes2.MC.iter=20, OncoBayes2.MC.chains=1)

## run single-agent analysis which defines blrmfit model object
example_model("single_agent")

nsamples(blrmfit)

## Recover user set sampling defaults
options(.user_mc_options)</pre>
```

OncoBayes2 33

Description

Bayesian logistic regression model with optional EXchangeability-NonEXchangeability parameter modelling for flexible borrowing from historical or concurrent data-sources. The safety model can guide dose-escalation decisions for adaptive Oncology phase I dose-escalation trials which involve an arbitrary number of drugs.

Global Options

Option	Default	Description
OncoBayes2.MC.warmup	1000	MCMC warmup iterations
OncoBayes2.MC.iter	2000	total MCMC iterations
OncoBayes2.MC.chains	4	MCMC chains
OncoBayes2.MC.thin	1	MCMC thinning
OncoBayes2.MC.control	list(adapt_delta=0.99,	sets control argument for Stan call
	stepsize=0.1)	
OncoBayes2.abbreviate.min	0	Minimal length of variable names
		when abbreviating variable names.
		The default 0 disables abbreviation.

References

Neuenschwander, B., Roychoudhury, S., & Schmidli, H. (2016). On the use of co-data in clinical trials. *Statistics in Biopharmaceutical Research*, 8(3), 345-354.

Neuenschwander, B., Wandel, S., Roychoudhury, S., & Bailey, S. (2016). Robust exchangeability designs for early phase clinical trials with multiple strata. *Pharmaceutical statistics*, 15(2), 123-134.

Neuenschwander, B., Branson, M., & Gsponer, T. (2008). Critical aspects of the Bayesian approach to phase I cancer trials. *Statistics in medicine*, 27(13), 2420-2439.

Neuenschwander, B., Matano, A., Tang, Z., Roychoudhury, S., Wandel, S. Bailey, Stuart. (2014). A Bayesian Industry Approach to Phase I Combination Trials in Oncology. In *Statistical methods in drug combination studies* (Vol. 69). CRC Press.

Stan Development Team (2019). RStan: the R interface to Stan. R package version 2.19.2. https://mc-stan.org

```
posterior_interval.blrmfit

Posterior intervals
```

Description

Posterior intervals of all model parameters.

Usage

```
## S3 method for class 'blrmfit'
posterior_interval(object, prob = 0.95, ...)
```

Arguments

object fitted model object

prob central probability mass to report, i.e. the quantiles 0.5-prob/2 and 0.5+prob/2 are displayed. Multiple central widths can be specified.

... not used in this function

Details

Reports the quantiles of posterior parameters which correspond to the central probability mass specified. The output includes the posterior of the hyper-parameters and the posterior of each group estimate.

Value

Matrix of two columns for the central probability interval prob for all parameters of the model.

```
## Setting up dummy sampling for fast execution of example
## Please use 4 chains and 100x more warmup & iter in practice
.user_mc_options <- options(OncoBayes2.MC.warmup=10, OncoBayes2.MC.iter=20, OncoBayes2.MC.chains=1)
example_model("single_agent")

posterior_interval(blrmfit)

## Recover user set sampling defaults
options(.user_mc_options)</pre>
```

Description

Calculates the posterior of the linear predictor.

Usage

```
## S3 method for class 'blrmfit'
posterior_linpred(object, transform = FALSE, newdata,
    draws, ...)
```

Arguments

object fitted model object

transform logical (defaults to FALSE) indicating if the linear predictor on the logit link scale is transformed with inv_logit to the 0-1 response scale.

newdata optional data frame specifying for what to predict; if missing, then the data of the input model object is used

draws number of returned posterior draws; by default the entire posterior is returned

... not used in this function

Details

Simulates the posterior of the linear predictor of the model object for the specified data set.

Value

Matrix of dimensions draws by nrow(newdata) where row correspond to a draw of the posterior and each column corresponds to a row in newdata. The columns are labelled with the row.names of newdata.

Group and strata definitions

The groups and strata as defined when running the blrm_exnex analysis cannot be changed at a later stage. As a result no evaluations can be performed for groups which have not been present in the data set used for running the analysis. However, it is admissible to code the group (and/or stratum) column as a factor which contains empty levels. These groups are thus not contained in the fitting data set and they are assigned by default to the first stratum. In addition priors must be setup for these groups (and/or strata). These empty group (and/or strata) levels are then allowed in subsequent evaluations. This enables the evaluation of the hierarchical model in terms of representing a prior for future groups.

Examples

```
## Setting up dummy sampling for fast execution of example
## Please use 4 chains and 100x more warmup & iter in practice
.user_mc_options <- options(OncoBayes2.MC.warmup=10, OncoBayes2.MC.iter=20, OncoBayes2.MC.chains=1)
## run single-agent analysis which defines blrmfit model object
example_model("single_agent")
## obtain posterior of linear prediction on 0-1 scale
post_prob_dlt <- posterior_linpred(blrmfit, TRUE, newdata=hist_SA)</pre>
## name columns to obtain nice bayesplot labels
colnames(post_prob_dlt) <- hist_SA$drug_A</pre>
library(bayesplot)
library(ggplot2)
mcmc_intervals(post_prob_dlt, prob=0.5, prob_outer=0.95) +
   coord_flip() +
   vline_at(c(0.16, 0.33), linetype=2) +
   ylab("Dose [mg]") +
   ggtitle("Posterior Probability of a DLT") +
   scale_x_continuous(breaks=c(0.1,0.16,0.33, 0.5, 0.75))
## Recover user set sampling defaults
options(.user_mc_options)
```

```
posterior_predict.blrmfit
```

Posterior of predictive

Description

Simulation of the predictive distribution.

Usage

```
## S3 method for class 'blrmfit'
posterior_predict(object, newdata, draws, ...)
```

Arguments

object fitted model object

newdata optional data frame specifying for what to predict; if missing, then the data of the input model object is used

draws number of returned posterior draws; by default the entire posterior is returned not used in this function

Details

Simulates the posterior predictive of the model object for the specified data set.

Value

Matrix of dimensions draws by nrow(newdata) where row correspond to a draw of the posterior and each column corresponds to a row in newdata. The columns are labelled with the row.names of newdata.

Group and strata definitions

The groups and strata as defined when running the blrm_exnex analysis cannot be changed at a later stage. As a result no evaluations can be performed for groups which have not been present in the data set used for running the analysis. However, it is admissible to code the group (and/or stratum) column as a factor which contains empty levels. These groups are thus not contained in the fitting data set and they are assigned by default to the first stratum. In addition priors must be setup for these groups (and/or strata). These empty group (and/or strata) levels are then allowed in subsequent evaluations. This enables the evaluation of the hierarchical model in terms of representing a prior for future groups.

```
## Setting up dummy sampling for fast execution of example
## Please use 4 chains and 100x more warmup & iter in practice
.user_mc_options <- options(OncoBayes2.MC.warmup=10, OncoBayes2.MC.iter=20, OncoBayes2.MC.chains=1)
example_model("single_agent")
post_pred <- posterior_predict(blrmfit)</pre>
## turn DLT counts into DLT rates
post_pred_rate <- sweep(post_pred, 2, hist_SA$num_patients, "/")</pre>
library(bayesplot)
library(ggplot2)
## compare posterior predictive of the model for the response rates
## with observed data
with(hist_SA,
   ppc_intervals(num_toxicities / num_patients, post_pred_rate, x=drug_A, prob_outer=0.95)) +
   xlab("Dose [mg]")
## Recover user set sampling defaults
options(.user_mc_options)
```

Description

Posterior predictive intervals of the model.

Usage

```
## S3 method for class 'blrmfit'
predictive_interval(object, prob = 0.95, newdata, ...)
```

Arguments

object fitted model object

prob central probability mass to report, i.e. the quantiles 0.5-prob/2 and 0.5+prob/2

are displayed. Multiple central widths can be specified.

newdata optional data frame specifying for what to predict; if missing, then the data of

the input model object is used

... not used in this function

Details

Reports for each row of the input data set the predictive interval according to the fitted model.

Value

Matrix with as many rows as the input data set and two columns which contain the lower and upper quantile corresponding to the central probability mass prob for the number of responses of the predictive distribution.

```
## Setting up dummy sampling for fast execution of example
## Please use 4 chains and 100x more warmup & iter in practice
.user_mc_options <- options(OncoBayes2.MC.warmup=10, OncoBayes2.MC.iter=20, OncoBayes2.MC.chains=1)
example_model("single_agent")
predictive_interval(blrmfit)
## Recover user set sampling defaults
options(.user_mc_options)</pre>
```

prior_summary.blrmfit 39

```
prior_summary.blrmfit Summarise model prior
```

Description

Extracts a summary of the prior in a structured data format.

Usage

```
## S3 method for class 'blrmfit'
prior_summary(object, digits = 2, ...)
```

Arguments

```
object blrmfit (blrm_trial) object as returned from blrm_exnex (blrm_trial) analysis
digits number of digits to show
... ignored by the function
```

Details

The summary of the prior creates a structured representation of the specified prior from a blrm_exnex (blrm_trial) analysis.

Value

Returns an analysis specific list, which has it's own print function. The returned list contains arrays which represent the prior in a structured format.

```
## Setting up dummy sampling for fast execution of example
## Please use 4 chains and 100x more warmup & iter in practice
.user_mc_options <- options(OncoBayes2.MC.warmup=10, OncoBayes2.MC.iter=20, OncoBayes2.MC.chains=1)
## run combo2 analysis which defines blrmfit model object
example_model("combo2")

prior_summary(blrmfit)

prior_sum <- prior_summary(blrmfit)

names(prior_sum)

## the entries of the prior list are labelled arrays
dimnames(prior_sum$EX_mu_log_beta)

## Recover user set sampling defaults
options(.user_mc_options)</pre>
```

40 summary.blrmfit

Description

Provides model summaries for blrm_exnex and blrm_trial analyses.

Usage

```
## S3 method for class 'blrmfit'
summary(object, newdata, transform = TRUE,
    prob = 0.95, interval_prob, ...)
```

Arguments

object fitted model object

newdata optional data frame specifying for what to predict; if missing, then the data of

the input model object is used

transform logical (defaults to FALSE) indicating if the linear predictor on the logit link scale

is transformed with inv_logit to the 0-1 response scale.

prob central probability mass to report, i.e. the quantiles 0.5-prob/2 and 0.5+prob/2

are displayed. Multiple central widths can be specified.

interval_prob optional vector of sorted quantiles for which the interval probabilities are calcu-

lated

... not used in this function

Details

The calculated posterior summaries are returned as a data.frame and contain optional interval probabilities for the specified vector of sorted quantiles. These summaries are calculated on the response scale by default and can be obtained on the link scale when setting transform=FALSE.

Value

Returns a data.frame of the key summaries of the posterior mean, standard deviation, central probability interval, median and optional interval probabilities. Each row of the data.frame corresponds to the respective input data which is by default the same data set as used for the blrm_exnex analysis or the data specified in the newdata argument.

```
## Setting up dummy sampling for fast execution of example
## Please use 4 chains and 100x more warmup & iter in practice
.user_mc_options <- options(OncoBayes2.MC.warmup=10, OncoBayes2.MC.iter=20, OncoBayes2.MC.chains=1)
example_model("single_agent")</pre>
```

summary.blrm_trial 41

```
## obtain underdosing (0-0.16), target dosing (0.16-0.33) and
## overdosing (0.33-1) probabilities
summary(blrmfit, interval_prob=c(0,0.16,0.33,1))
## Recover user set sampling defaults
options(.user_mc_options)
```

summary.blrm_trial

Summarise trial

Description

Provides model summaries for blrm_trial analyses.

Usage

```
## $3 method for class 'blrm_trial'
summary(object, summarize = c("blrmfit",
   "blrm_exnex_call", "dose_info", "dose_prediction", "data",
   "data_prediction", "dimensionality"), ...)
```

Arguments

object

blrm_trial object

summarize

one of the following options:

- blrmfit: summary of the underlying blrmfit object with further arguments
- blrm_exnex_call: blrm_exnex call used to create the blrmfit object
- dose_info: dose_info that were defined
- dose_prediction prediction for the defined dose_info
- data: data that were observed
- data_prediction: prediction for the observed data
- dimensionality: numeric vector with entries "num_components", "num_interaction_terms", "num_groups", "num_strata"

.. further arguments for summary.blrmfit

```
## Setting up dummy sampling for fast execution of example
## Please use 4 chains and 100x more warmup & iter in practice
.user_mc_options <- options(OncoBayes2.MC.warmup=10, OncoBayes2.MC.iter=20, OncoBayes2.MC.chains=1)
# construct initial blrm_trial object from built-in example datasets
combo2_trial_setup <- blrm_trial(
    data = hist_combo2,</pre>
```

42 update.blrmfit

```
dose_info = dose_info_combo2,
  drug_info = drug_info_combo2,
  simplified_prior = TRUE
)

# extract blrm_call to see setup of the prior as passed to blrm_exnex
  summary(combo2_trial_setup, "blrm_exnex_call")

## Recover user set sampling defaults
  options(.user_mc_options)
```

update.blrmfit

Update data of a BLRM analysis

Description

Adds data rows to a blrm_exnex or blrm_trial analysis object.

Usage

```
## S3 method for class 'blrmfit'
update(object, ..., add_data)
```

Arguments

object blrmfit analysis object

... passed to default update command

The data in add_data will be combined with data in object using bind_rows. The indices for groups and stratums (if defined) are matched between add_data

and the data of the analysis object.

Note that the add_data argument must be named explicitly as demonstrated in

the example.

add_data additional data added to analysis data of object

```
## Setting up dummy sampling for fast execution of example
## Please use 4 chains and 100x more warmup & iter in practice
.user_mc_options <- options(OncoBayes2.MC.warmup=10, OncoBayes2.MC.iter=20, OncoBayes2.MC.chains=1)
example_model("single_agent")
library(tibble)
new_cohort <- tibble(group_id="trial_A", drug_A=50, num_patients=4, num_toxicities=1)
## this would fail, since add_data argument must be named
## new_blrmfit <- update(blrmfit, new_cohort)</pre>
```

update.blrm_trial 43

```
new_blrmfit <- update(blrmfit, add_data=new_cohort)
## Recover user set sampling defaults
options(.user_mc_options)</pre>
```

update.blrm_trial

Update data and/or prior of a BLRM trial

Description

* Adds data rows to a blrm_trial object (add_data argument) * Replaces data of a blrm_trial object (data argument) * Sets the prior of a blrm_trial object (... argument will be passed to blrm_exnex)

Usage

```
## S3 method for class 'blrm_trial'
update(object, ...)
```

Arguments

object blrm_trial object
... passed to default update command of blrm_exnex

```
## Setting up dummy sampling for fast execution of example
## Please use 4 chains and 100x more warmup & iter in practice
.user_mc_options <- options(OncoBayes2.MC.warmup=10, OncoBayes2.MC.iter=20, OncoBayes2.MC.chains=1)

# the combo2_trial example demonstrates the use of add_data of
# update.blrmfit
example_model("combo2_trial")

## Recover user set sampling defaults
options(.user_mc_options)</pre>
```

Index

```
*Topic datasets
                                                posterior_predict
    codata_combo2, 15
                                                        (posterior_predict.blrmfit), 36
    dose_info_combo2, 16
                                                posterior_predict.blrmfit, 36
    drug_info_combo2, 17
                                                predictive_interval
    hist_combo2, 28
                                                        (predictive_interval.blrmfit),
    hist_combo3, 29
    hist_SA, 30
                                                predictive_interval.blrmfit, 38
                                                print.blrm_trial(blrm_trial), 11
bind_rows_0, 3
                                                print.blrmfit(blrm_exnex), 3
blrm_exnex, 3, 20, 39, 40, 42
                                                prior_summary(prior_summary.blrmfit),
blrm_formula_linear, 10
blrm_trial, 11, 17, 20, 39-43
                                                prior_summary.blrmfit, 39
codata_combo2, 15
                                                stan, 5
                                                summary.blrm_trial, 20, 41
dose_info_combo2, 14, 16, 17, 20
                                                summary.blrmfit, 40
drug_info_combo2, 14, 17, 17, 20
                                                update.blrm_trial, 20, 43
example-combo2, 18
                                                update.blrmfit, 42
example-combo2_trial, 20
example-combo3, 22
example-single-agent, 25
example_model, 20, 27
hist_combo2, 28
hist_combo3, 29
hist_SA, 30
nsamples (nsamples.blrmfit), 32
nsamples.blrmfit, 32
OncoBayes2, 33
OncoBayes2-package (OncoBayes2), 33
posterior_interval
        (posterior_interval.blrmfit),
        34
posterior_interval.blrmfit, 34
posterior_linpred
        (posterior_linpred.blrmfit), 35
posterior_linpred.blrmfit, 35
```