Package: BayesianMCPMod (via r-universe)

March 7, 2025

Title Simulate, Evaluate, and Analyze Dose Finding Trials with Bayesian MCPMod

Version 1.1.0

Description Bayesian MCPMod (Fleischer et al. (2022) <doi:10.1002/pst.2193>) is an innovative method that improves

the traditional MCPMod by systematically incorporating historical data, such as previous placebo group data. This R package offers functions for simulating, analyzing, and evaluating Bayesian MCPMod trials with normally distributed endpoints. It enables the assessment of trial designs incorporating historical data across various true dose-response relationships and sample sizes. Robust mixture prior distributions, such as those derived with the Meta-Analytic-Predictive approach (Schmidli et al. (2014) <doi:10.1111/biom.12242>), can be specified for each dose group. Resulting mixture posterior distributions are used in the Bayesian Multiple Comparison Procedure and modeling steps. The modeling step also includes a weighted model averaging approach (Pinheiro et al. (2014) <doi:10.1002/sim.6052>). Estimated dose-response relationships can be bootstrapped and visualized.

License Apache License (>= 2)

URL https://boehringer-ingelheim.github.io/BayesianMCPMod/,

https://github.com/Boehringer-Ingelheim/BayesianMCPMod

BugReports https://github.com/Boehringer-Ingelheim/BayesianMCPMod/issues

Depends R (>= 4.2)

- **Imports** checkmate, DoseFinding (>= 1.1-1), future.apply, ggplot2, methods, nloptr, RBesT, stats
- Suggests clinDR, data.table, doFuture, doRNG, dplyr, future, kableExtra, knitr, MCPModPack, reactable, rmarkdown, spelling, testthat (>= 3.0.0), tibble, tidyr

VignetteBuilder knitr

Config/testthat/edition 3

Encoding UTF-8

Language en-US

RoxygenNote 7.3.2

NeedsCompilation no

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Repository CRAN

Date/Publication 2025-03-07 19:00:07 UTC

Config/pak/sysreqs cmake make libicu-dev

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assessDesign assessDesign

Description

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This function performs simulation based trial design evaluations for a set of specified dose-response models

assessDesign

Usage

```
assessDesign(
  n_patients,
  mods,
  prior_list,
  sd,
  n_{sim} = 1000,
  alpha_crit_val = 0.05,
  modeling = FALSE,
  simple = TRUE,
  avg_fit = TRUE,
  reestimate = FALSE,
  contr = NULL,
  dr_means = NULL,
  delta = NULL,
  evidence_level = NULL,
 med_selection = c("avgFit", "bestFit")
)
```

Arguments

n_patients	Vector specifying the planned number of patients per dose group. A minimum of 2 patients are required in each group.
mods	An object of class "Mods" as specified in the DoseFinding package.
prior_list	A prior_list object specifying the utilized prior for the different dose groups
sd	A positive value, specification of assumed sd
n_sim	Number of simulations to be performed
alpha_crit_val	(Un-adjusted) Critical value to be used for the MCP testing step. Passed to the getCritProb() function for the calculation of adjusted critical values (on the probability scale). Default is 0.05.
modeling	Boolean variable defining whether the Mod part of Bayesian MCP-Mod will be performed in the assessment. More heavy on resources. Default FALSE.
simple	Boolean variable defining whether simplified fit will be applied. Passed to the getModelFits function. Default FALSE.
avg_fit	Boolean variable, defining whether an average fit (based on generalized AIC weights) should be performed in addition to the individual models. Default TRUE.
reestimate	Boolean variable defining whether critical value should be calculated with re- estimated contrasts (see getCritProb function for more details). Default FALSE
contr	An object of class 'optContr' as created by the getContr() function. Allows specification of a fixed contrasts matrix. Default NULL
dr_means	A vector, allows specification of individual (not model based) assumed effects per dose group. Default NULL
delta	A numeric value for the threshold Delta for the MED assessment. If NULL, no MED assessment is performed. Default NULL.

evidence_level	A numeric value between 0 and 1 for the evidence level gamma for the MED
	assessment. Only required for Bayesian MED assessment, see ?getMED for
	details. Default NULL.
med_selection	A string, either "avgFit" or "bestFit", for the method of MED selection. Default "avgFit".

Value

Returns success probabilities for the different assumed dose-response shapes, attributes also includes information around average success rate (across all assumed models) and prior Effective sample size

Examples

```
mods <- DoseFinding::Mods(linear</pre>
                                      = NULL,
                                      = c(0.5, 1.2),
                          emax
                          exponential = 2,
                          doses
                                      = c(0, 0.5, 2, 4, 8),
                          maxEff
                                      = 6)
sd <- 12
prior_list <- list(Ctrl = RBesT::mixnorm(comp1 = c(w = 1, m = 0, s = 12), sigma = 2),
                   DG_1 = RBesT::mixnorm(comp1 = c(w = 1, m = 1, s = 12), sigma = 2),
                   DG_2 = RBesT::mixnorm(comp1 = c(w = 1, m = 1.2, s = 11), sigma = 2) ,
                   DG_3 = RBesT::mixnorm(comp1 = c(w = 1, m = 1.3, s = 11), sigma = 2) ,
                   DG_4 = RBesT::mixnorm(comp1 = c(w = 1, m = 2, s = 13), sigma = 2))
n_patients <- c(40, 60, 60, 60, 60)
success_probabilities <- assessDesign(</pre>
 n_patients = n_patients,
             = mods,
 mods
 prior_list = prior_list,
              = sd,
 sd
 n_sim
              = 1e2) # speed up example run time
success_probabilities
if (interactive()) { # takes typically > 5 seconds
# with MED estimation without bootstrapping
# see ?getMED for details
success_probabilities <- assessDesign(</pre>
              = n_patients,
 n_patients
 mods
                = mods,
                = prior_list,
 prior_list
 sd
                 = sd,
 modeling
                 = TRUE,
                 = 10, # speed up example run time
 n_sim
 delta
                 = 7)
 success_probabilities
```

```
# with MED estimation with bootstrapping
success_probabilities <- assessDesign(</pre>
             = n_patients,
 n_patients
 mods
                = mods,
 prior_list = prior_list,
 sd
                = sd,
 modeling
                = TRUE,
 n_sim
                = 10, # speed up example run time
 delta
                = 7,
 evidence_level = 0.8)
 success_probabilities
}
```

getBootstrapQuantiles getBootstrapQuantiles

Description

A function for the calculation of bootstrapped model predictions. Samples from the posterior distribution are drawn (via the RBesT function rmix()) and for every sample the simplified fitting step (see getModelFits() function) and a prediction is performed. These fits are then used to identify the specified quantiles. This approach can be considered as the Bayesian equivalent of the frequentist bootstrap approach described in O'Quigley et al. (2017). Instead of drawing n bootstrap samples from the sampling distribution of the trial dose-response estimates, here the samples are directly taken from the posterior distribution.

Usage

```
getBootstrapQuantiles(model_fits, quantiles, n_samples = 1000, doses = NULL)
```

Arguments

model_fits	An object of class modelFits, i.e. information about fitted models & correspond- ing model coefficients as well as the posterior distribution that was the basis for the model fitting
quantiles	A vector of quantiles that should be evaluated
n_samples	Number of samples that should be drawn as basis for the bootstrapped quantiles
doses	A vector of doses for which a prediction should be performed. If NULL, the dose levels of the model_fits will be used. Default NULL.

Value

A data frame with columns for model, dose, and bootstrapped samples

References

O'Quigley J, Iasonos A, Bornkamp B. 2017. Handbook of Methods for Designing, Monitoring, and Analyzing Dose-Finding Trials (1st ed.). Chapman and Hall/CRC. doi:10.1201/9781315151984

Examples

```
posterior_list <- list(Ctrl = RBesT::mixnorm(comp1 = c(w = 1, m = 0, s = 1), sigma = 2),</pre>
                      DG_1 = RBesT::mixnorm(comp1 = c(w = 1, m = 3, s = 1.2), sigma = 2),
                     DG_2 = RBesT::mixnorm(comp1 = c(w = 1, m = 4, s = 1.5), sigma = 2) ,
                     DG_3 = RBesT::mixnorm(comp1 = c(w = 1, m = 6, s = 1.2), sigma = 2) ,
                     DG_4 = RBesT::mixnorm(comp1 = c(w = 1, m = 6.5, s = 1.1), sigma = 2))
models
               <- c("exponential", "linear")
dose_levels
               <- c(0, 1, 2, 4, 8)
model fits
               <- getModelFits(models
                                            = models.
                               posterior = posterior_list,
                               dose_levels = dose_levels,
                                simple
                                            = TRUE)
bs_quantiles <- getBootstrapQuantiles(model_fits = model_fits,</pre>
                                       quantiles = c(0.025, 0.5, 0.8, 0.975),
                                       n_samples = 10, # speeding up example run time
                                       doses
                                                  = c(0, 6, 8))
bs_quantiles
```

getBootstrapSamples getBootstrapSamples

Description

A function to return bootstrap samples from the fitted dose-response models. Samples from the posterior distribution are drawn (via the RBesT function rmix()) and for every sample the simplified fitting step (see getModelFits() function) and a prediction is performed. These samples are returned by this function. This approach can be considered as the Bayesian equivalent of the frequentist bootstrap approach described in O'Quigley et al. (2017). Instead of drawing n bootstrap samples from the sampling distribution of the trial dose-response estimates, here the samples are directly taken from the posterior distribution.

Usage

```
getBootstrapSamples(model_fits, n_samples = 1000, doses = NULL)
```

Arguments

model_fits	An object of class modelFits, i.e. information about fitted models & correspond- ing model coefficients as well as the posterior distribution that was the basis for the model fitting
n_samples	Number of samples that should be drawn
doses	A vector of doses for which a prediction should be performed

getContr

Value

A data frame with entries model, dose, and sample

getContr

References

O'Quigley J, Iasonos A, Bornkamp B. 2017. Handbook of Methods for Designing, Monitoring, and Analyzing Dose-Finding Trials (1st ed.). Chapman and Hall/CRC. doi:10.1201/9781315151984

Examples

```
posterior_list <- list(Ctrl = RBesT::mixnorm(comp1 = c(w = 1, m = 0, s = 1), sigma = 2),</pre>
                      DG_1 = RBesT::mixnorm(comp1 = c(w = 1, m = 3, s = 1.2), sigma = 2),
                     DG_2 = RBesT::mixnorm(comp1 = c(w = 1, m = 4, s = 1.5), sigma = 2) ,
                     DG_3 = RBesT::mixnorm(comp1 = c(w = 1, m = 6, s = 1.2), sigma = 2),
                     DG_4 = RBesT::mixnorm(comp1 = c(w = 1, m = 6.5, s = 1.1), sigma = 2))
models
               <- c("exponential", "linear")
dose_levels
               <- c(0, 1, 2, 4, 8)
model_fits
               <- getModelFits(models
                                            = models.
                                posterior = posterior_list,
                                dose_levels = dose_levels,
                                simple
                                            = TRUE)
bs_samples <- getBootstrapSamples(model_fits = model_fits,</pre>
                                  n_samples = 10, # speeding up example run time
                                   doses
                                              = c(0, 6, 8))
bs_samples
```

Description

getContr

This function calculates contrast vectors that are optimal for detecting certain alternatives via applying the function optContr() of the DoseFinding package. Hereby, 4 different options can be distinguished that are automatically executed based on the input that is provided

- Bayesian approach: If dose_weights and a prior_list are provided an optimized contrasts for the posterior sample size is calculated. In detail, in a first step the dose_weights (typically the number of patients per dose group) and the prior information is combined by calculating for each dose group a posterior effective sample. Based on this posterior effective sample sizes the allocation ratio is derived, which allows for a calculation on pseudo-optimal contrasts via regular MCPMod are calculated from the regular MCPMod for these specific weights
- 2. Frequentist approach: If only dose_weights are provided optimal contrast vectors are calculated from the regular MCPMod for these specific weights
- Bayesian approach + re-estimation: If only a sd_posterior (i.e. variability of the posterior distribution) is provided, pseudo-optimal contrasts based on these posterior weights will be calculated

4. Frequentist approach+re-estimation: If only a se_new_trial (i.e. the estimated variability per dose group of a new trial) is provided, optimal contrast vectors are calculated from the regular MCPMod for this specific vector of standard errors. For the actual evaluation this vector of standard errors is translated into a (diagonal) matrix of variances

Usage

```
getContr(
  mods,
  dose_levels,
  dose_weights = NULL,
  prior_list = NULL,
  sd_posterior = NULL,
  se_new_trial = NULL
)
```

Arguments

mods	An object of class 'Mods' as created by the function 'DoseFinding::Mods()'
dose_levels	Vector containing the different dosage levels.
dose_weights	Vector specifying weights for the different doses. Please note that in case this information is provided together with a prior (i.e. Option 1) is planned these two inputs should be provided on the same scale (e.g. patient numbers). Default NULL
prior_list	A list of objects of class 'normMix' as created with 'RBesT::mixnorm()'. Only required as input for Option 1. Default NULL
sd_posterior	A vector of positive values with information about the variability of the posterior distribution, only required for Option 3. Default NULL
se_new_trial	A vector of positive values with information about the observed variability, only required for Option 4. Default NULL

Value

An object of class 'optContr' as provided by the function 'DoseFinding::optContr()'.

Examples

```
dose_levels <- c(0, 0.5, 2, 4, 8)
mods <- DoseFinding::Mods(</pre>
              = NULL,
  linear
              = c(0.5, 1.2),
  emax
  exponential = 2,
              = dose_levels,
  doses
  maxEff
              = 6)
sd_posterior <- c(2.8, 3, 2.5, 3.5, 4)
contr_mat <- getContr(</pre>
  mods
              = mods,
  dose_levels = dose_levels,
```

getCritProb

```
sd_posterior = sd_posterior)
```

getCritProb getCritProb

Description

This function calculates multiplicity adjusted critical values. The critical values are calculated in such a way that when using non-informative priors the actual error level for falsely declaring a significant trial in the Bayesian MCPMod is controlled (by the specified alpha level). Hereby optimal contrasts of the frequentist MCPMod are applied and two options can be distinguished

- 1. Frequentist approach: If only dose_weights are provided optimal contrast vectors are calculated from the regular MCPMod for these specific weights and the corresponding critical value for this set of contrasts is calculated via the critVal() function of the DoseFinding package.
- 2. Frequentist approach + re-estimation: If only a se_new_trial (i.e. the estimated variability per dose group of a new trial) is provided, optimal contrast vectors are calculated from the regular MCPMod for this specific vector of standard errors. Here as well the critical value for this set of contrasts is calculated via the critVal() function of the DoseFinding package.

Usage

```
getCritProb(
  mods,
  dose_levels,
  dose_weights = NULL,
  se_new_trial = NULL,
  alpha_crit_val = 0.025
)
```

Arguments

mods	An object of class "Mods" as specified in the DoseFinding package.
dose_levels	Vector containing the different dosage levels.
dose_weights	Vector specifying weights for the different doses, only required for Option i). Default NULL
<pre>se_new_trial</pre>	A vector of positive values, only required for Option ii). Default NULL
alpha_crit_val	Significance level. Default set to 0.025.

Value

Multiplicity adjusted critical value on the probability scale.

Examples

|--|--|

Description

This function calculates the effective sample size for every dose group via the RBesT function ess().

Usage

```
getESS(post_list)
```

Arguments

post_list A posterior list object, for which the effective sample size for each dose group should be calculated

Value

A vector of the effective sample sizes for each dose group

|--|--|--|

Description

This function provides information on the minimally efficacious dose (MED). The MED evaluation can either be based on the fitted model shapes (model_fits) or on bootstrapped quantiles (bs_quantiles).

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getMED

Usage

```
getMED(
   delta,
   evidence_level = 0.5,
   dose_levels = NULL,
   model_fits = NULL,
   bs_quantiles = NULL
)
```

Arguments

delta	A numeric value for the threshold Delta.
evidence_level	A numeric value between 0 and 1 for the evidence level gamma. Used for the bs_quantiles-based evaluation and not used for the model_fits-based evaluation. Default 0.5.
dose_levels	A vector of numerics containing the different dosage levels. Default NULL.
model_fits	An object of class modelFits as created with getModelFits(). Default NULL.
bs_quantiles	A dataframe created with getBootstrapQuantiles(). Default NULL.

Details

The function assumes that the 1st dose group is the control dose group.

The bootstrap approach allows for an MED based on decision rules of the form

$$\widehat{\text{MED}} = \arg\min_{d \in \{d_1, \dots, d_k\}} \left\{ \Pr\left(f(d, \hat{\theta}) - f(d_1, \hat{\theta}) > \Delta \right) > \gamma \right\}.$$

The model-shape approach takes the point estimate of the model into account.

Value

A matrix with rows for MED reached, MED, and MED index in the vector of dose levels and columns for the dose-response shapes.

Examples

```
posterior_list <- list(Ctrl = RBesT::mixnorm(comp1 = c(w = 1, m = 0, s = 1), sigma = 2),</pre>
                     DG_1 = RBesT::mixnorm(comp1 = c(w = 1, m = 3, s = 1.2), sigma = 2),
                     DG_2 = RBesT::mixnorm(comp1 = c(w = 1, m = 4, s = 1.5), sigma = 2) ,
                     DG_3 = RBesT::mixnorm(comp1 = c(w = 1, m = 6, s = 1.2), sigma = 2) ,
                    DG_4 = RBesT::mixnorm(comp1 = c(w = 1, m = 6.5, s = 1.1), sigma = 2))
              <- c("exponential", "linear")
models
               <- c(0, 1, 2, 4, 8)
dose_levels
model_fits
               <- getModelFits(models
                                           = models,
                               posterior = posterior_list,
                               dose_levels = dose_levels,
                                           = TRUE)
                               simple
```

MED based on the model_fit:

getModelFits getModelFits

Description

Fits dose-response curves for the specified dose-response models, based on the posterior distributions. For the simplified fit, multivariate normal distributions will be approximated and reduced by one-dimensional normal distributions. For the default case, the Nelder-Mead algorithm is used. In detail, for both approaches the mean vector θ^Y and the covariance Σ of the (mixture) posterior distributions and the corresponding posterior weights $\tilde{\omega}_l$ for $l \in 1, ..., L$ are used as basis For the full fit a GLS estimator is used to minimize the following expression for the respective dose-response models m

$$\hat{\theta}_m = \arg\min_{\theta_m} \sum_{l=1}^{L} \tilde{\omega}_l (\theta_{l_i}^Y - f(dose_i, \hat{\theta}_m))' \Sigma_l^{-1} (\theta_{l_i}^Y - f(dose_i, \hat{\theta}_m))$$

Therefore the function nloptr of the nloptr package is utilized. In the simplified case L = 1, as the dimension of the posterior is reduced to 1 first. The generalized AIC values are calculated via the formula

$$gAIC_{m} = \sum_{l=1}^{L} \tilde{\omega}_{l} \sum_{i=0}^{K} \frac{1}{\sum_{l_{i,i}}} (\theta_{l_{i}}^{Y} - f(dose_{i}, \hat{\theta}_{m}))^{2} + 2p$$

where p denotes the number of estimated parameters and K the number of active dose levels. Here as well for the simplified case the formula reduces to one summand as L = 1. Corresponding gAIC based weights for model M are calculated as outlined in Schorning et al. (2016)

$$\Omega_I(M) = \frac{\exp(-0.5gAIC_M)}{\sum_{m=1}^{Q} \exp(-0.5gAIC_m)}$$

where Q denotes the number of models included in the averaging procedure.

Usage

```
getModelFits(models, dose_levels, posterior, avg_fit = TRUE, simple = FALSE)
```

getModelFits

Arguments

models	List (or vector) of model names for which a fit will be performed.
dose_levels	A vector containing the different dosage levels.
posterior	A getPosterior object, containing the (multivariate) posterior distribution per dosage level.
avg_fit	Boolean variable, defining whether an average fit (based on generalized AIC weights) should be performed in addition to the individual models. Default TRUE.
simple	Boolean variable, defining whether simplified fit will be applied. Default FALSE.

Value

An object of class modelFits. A list containing information about the fitted model coefficients, the prediction per dose group as well as maximum effect and generalized AIC (and corresponding weight) per model.

References

Schorning K, Bornkamp B, Bretz F, Dette H. 2016. Model selection versus model averaging in dose finding studies. Stat Med; 35; 4021-4040.

Examples

```
posterior_list <- list(Ctrl = RBesT::mixnorm(comp1 = c(w = 1, m = 0, s = 1), sigma = 2),</pre>
                      DG_1 = RBesT::mixnorm(comp1 = c(w = 1, m = 3, s = 1.2), sigma = 2),
                     DG_2 = RBesT::mixnorm(comp1 = c(w = 1, m = 4, s = 1.5), sigma = 2) ,
                     DG_3 = RBesT::mixnorm(comp1 = c(w = 1, m = 6, s = 1.2), sigma = 2) ,
                     DG_4 = RBesT::mixnorm(comp1 = c(w = 1, m = 6.5, s = 1.1), sigma = 2))
models
               <- c("emax", "exponential", "sigEmax", "linear")
               <- c(0, 1, 2, 4, 8)
dose_levels
fit
           <- getModelFits(models
                                       = models,
                           posterior = posterior_list,
                           dose_levels = dose_levels)
fit
fit_simple <- getModelFits(models</pre>
                                        = models,
                           posterior = posterior_list,
                           dose_levels = dose_levels,
                           simple
                                       = TRUE)
fit_simple
```

getPosterior

Description

Either the patient level data or both mu_hat as well as S_hat must to be provided. If patient level data is provided mu_hat and S_hat are calculated within the function using a linear model. This function calculates the posterior distribution. Depending on the input for S_hat this step is either performed for every dose group independently via the RBesT function postmix() or the mvpostmix() function of the DoseFinding package is utilized. In the latter case conjugate posterior mixture of multivariate normals are calculated (DeGroot 1970, Bernardo and Smith 1994)

Usage

```
getPosterior(
   prior_list,
   data = NULL,
   mu_hat = NULL,
   S_hat = NULL,
   calc_ess = FALSE
)
```

Arguments

prior_list	a prior list with information about the prior to be used for every dose group
data	dataframe containing the information of dose and response. Default NULL Also a simulateData object can be provided.
mu_hat	vector of estimated mean values (per dose group).
S_hat	Either a vector or a covariance matrix specifying the (estimated) variability can be specified. The length of the vector (resp. the dimension of the matrix) needs to match the number of dose groups. Please note that for a vector input the numbers should reflect the standard error per dose group (i.e. square root of variance), while for a matrix input the variance-covariance matrix should be provided.
calc_ess	boolean variable, indicating whether effective sample size should be calculated. Default FALSE

Details

Kindly note that one can sample from the posterior_list with lapply(posterior_list, RBesT::rmix, n = 10).

Value

posterior_list, a posterior list object is returned with information about (mixture) posterior distribution per dose group (more detailed information about the conjugate posterior in case of covariance input for S_hat is provided in the attributes)

References

BERNARDO, Jl. M., and Smith, AFM (1994). Bayesian Theory. 81.

Examples

performBayesianMCP performBayesianMCP

Description

Performs Bayesian MCP Test step, as described in Fleischer et al. (2022). Tests for a dose-response effect using a model-based multiple contrast test based on the (provided) posterior distribution. In particular for every dose-response candidate the posterior probability is calculated that the contrast is bigger than 0 (based on the posterior distribution of the dose groups). In order to obtain significant test decision we consider the maximum of the posterior probabilities across the different models. This maximum is compared with a (multiplicity adjusted) critical value (on the probability scale).

Usage

```
performBayesianMCP(posterior_list, contr, crit_prob_adj)
```

Arguments

posterior_list	An object derived with getPosterior with information about the (mixture) posterior distribution per dose group
contr	An object of class 'optContr' as created by the getContr() function. It contains the contrast matrix to be used for the testing step.
crit_prob_adj	A getCritProb object, specifying the critical value to be used for the testing (on the probability scale)

Value

Bayesian MCP test result, with information about p-values for the individual dose-response shapes and overall significance

References

Fleischer F, Bossert S, Deng Q, Loley C, Gierse J. 2022. Bayesian MCPMod. Pharmaceutical Statistics. 21(3): 654-670. doi:10.1002/pst.2193

Examples

```
mods <- DoseFinding::Mods(linear</pre>
                                      = NULL,
                          emax
                                      = c(0.5, 1.2),
                          exponential = 2,
                                      = c(0, 0.5, 2, 4, 8))
                          doses
dose_levels <- c(0, 0.5, 2, 4, 8)
sd_posterior <- c(2.8,3,2.5,3.5,4)</pre>
contr_mat <- getContr(</pre>
 mods
              = mods,
 dose_levels = dose_levels,
 sd_posterior = sd_posterior)
critVal <- getCritProb(</pre>
 mods
                 = mods,
 dose_weights = c(50, 50, 50, 50, 50), #reflecting the planned sample size
 dose_levels = dose_levels,
 alpha_crit_val = 0.05)
prior_list <- list(Ctrl = RBesT::mixnorm(comp1 = c(w = 1, m = 0, s = 5), sigma = 2),</pre>
                   DG_1 = RBesT::mixnorm(comp1 = c(w = 1, m = 1, s = 12), sigma = 2),
                   DG_2 = RBesT::mixnorm(comp1 = c(w = 1, m = 1.2, s = 11), sigma = 2) ,
                   DG_3 = RBesT::mixnorm(comp1 = c(w = 1, m = 1.3, s = 11), sigma = 2),
                   DG_4 = RBesT::mixnorm(comp1 = c(w = 1, m = 2, s = 13), sigma = 2))
mu <- c(0, 1, 1.5, 2, 2.5)
S_hat <- c(5, 4, 6, 7, 8)
posterior_list <- getPosterior(</pre>
 prior_list = prior_list,
 mu_hat
          = mu,
 S_hat
            = S_hat,
 calc_ess = TRUE)
performBayesianMCP(posterior_list = posterior_list,
                   contr = contr_mat,
                   crit_prob_adj = critVal)
```

performBayesianMCPMod performBayesianMCPMod

Description

Performs Bayesian MCP Test step and modeling in a combined fashion. See performBayesian-MCP() function for MCP Test step and getModelFits() for the modeling step

Usage

```
performBayesianMCPMod(
   posterior_list,
   contr,
   crit_prob_adj,
   simple = FALSE,
   avg_fit = TRUE,
   delta = NULL,
   evidence_level = NULL,
   med_selection = c("avgFit", "bestFit"),
   n_samples = 1000
)
```

Arguments

posterior_list	An object of class 'postList' as created by getPosterior() containing information about the (mixture) posterior distribution per dose group
contr	An object of class 'optContr' as created by the getContr() function. It contains the contrast matrix to be used for the testing step.
crit_prob_adj	A getCritProb object, specifying the critical value to be used for the testing (on the probability scale).
simple	Boolean variable, defining whether simplified fit will be applied. Passed to the getModelFits() function. Default FALSE.
avg_fit	Boolean variable, defining whether an average fit (based on generalized AIC weights) should be performed in addition to the individual models. Default TRUE.
delta	A numeric value for the threshold Delta for the MED assessment. If NULL, no MED assessment is performed. Default NULL.
evidence_level	A numeric value between 0 and 1 for the evidence level gamma for the MED assessment. Only required for Bayesian MED assessment, see ?getMED for details. Default NULL.
med_selection	A string, either "avgFit" or "bestFit" based on the lowest gAIC, for the method of MED selection. Default "avgFit".
n_samples	A numerical for the number of bootstrapped samples in case the Bayesian MED assessment is performed. Default 1e3.

Value

Bayesian MCP test result as well as modeling result.

Examples

```
mods <- DoseFinding::Mods(linear</pre>
                                      = NULL,
                                      = c(0.5, 1.2),
                          emax
                          exponential = 2,
                                      = c(0, 0.5, 2, 4, 8))
                          doses
dose_levels <- c(0, 0.5, 2, 4, 8)
sd_posterior <- c(2.8, 3, 2.5, 3.5, 4)
contr_mat <- getContr(</pre>
  mods
              = mods,
  dose_levels = dose_levels,
  sd_posterior = sd_posterior)
critVal <- getCritProb(</pre>
  mods
                 = mods,
  dose_weights = c(50, 50, 50, 50, 50), #reflecting the planned sample size
                 = dose_levels,
  dose_levels
  alpha_crit_val = 0.6) # unreasonable alpha for this example, rather choose 0.05
prior_list <- list(Ctrl = RBesT::mixnorm(comp1 = c(w = 1, m = 0, s = 5), sigma = 2),</pre>
                   DG_1 = RBesT::mixnorm(comp1 = c(w = 1, m = 1, s = 12), sigma = 2),
                   DG_2 = RBesT::mixnorm(comp1 = c(w = 1, m = 1.2, s = 11), sigma = 2) ,
                   DG_3 = RBesT::mixnorm(comp1 = c(w = 1, m = 1.3, s = 11), sigma = 2),
                   DG_4 = RBesT::mixnorm(comp1 = c(w = 1, m = 2, s = 13), sigma = 2))
mu <- c(0, 1, 1.5, 2, 2.5)
S_hat <- c(5, 4, 6, 7, 8)
posterior_list <- getPosterior(</pre>
  prior_list = prior_list,
  mu_hat
             = mu,
  S_hat
             = S_hat,
  calc_ess = TRUE)
performBayesianMCPMod(posterior_list = posterior_list,
                      contr = contr_mat,
                      crit_prob_adj = critVal,
                      simple
                                     = FALSE,
                      delta
                                     = 1)
```

plot.modelFits plot.modelFits

Description

Plot function based on the ggplot2 package. Providing visualizations for each model and a average Fit. Black lines show the fitted dose response models and an AIC based average model. Dots indicate the posterior median and vertical lines show corresponding credible intervals (i.e. the variability of the posterior distribution of the respective dose group). To assess the uncertainty of the model fit one can in addition visualize credible bands (default coloring as orange shaded areas). The calculation of these bands is performed via the getBootstrapQuantiles() function. The default setting is that these credible bands are not calculated.

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plot.modelFits

Usage

```
## S3 method for class 'modelFits'
plot(
    x,
    gAIC = TRUE,
    cr_intv = TRUE,
    alpha_CrI = 0.05,
    cr_bands = FALSE,
    alpha_CrB = c(0.05, 0.2),
    n_bs_smpl = 1000,
    acc_color = "orange",
    plot_res = 100,
    ...
)
```

Arguments

х	An object of type modelFits
gAIC	Logical value indicating whether gAIC values are shown in the plot. Default TRUE
cr_intv	Logical value indicating whether credible intervals are included in the plot. Default TRUE
alpha_CrI	Numerical value of the width of the credible intervals. Default is set to 0.05 (i.e 95% CI are shown).
cr_bands	Logical value indicating whether bootstrapped based credible bands are shown in the plot. Default FALSE
alpha_CrB	Numerical vector of the width of the credible bands. Default is set to 0.05 and 0.5 (i.e 95% CB and 50% CB are shown).
n_bs_smpl	Number of bootstrap samples being used. Default 1000.
acc_color	Color of the credible bands. Default "orange".
plot_res	Number of plotted doses within the range of the dose levels, i.e., the resolution of the plot. Default 100.
	optional parameter to be passed to plot().

Value

A ggplot2 object

Examples

predict.modelFits predict.modelFits

Description

This function performs model predictions based on the provided model and dose specifications

Usage

S3 method for class 'modelFits'
predict(object, doses = NULL, ...)

Arguments

object	A modelFits object containing information about the fitted model coefficients
doses	A vector specifying the doses for which a prediction should be done
	Currently without function

Value

a list with the model predictions for the specified models and doses

Examples

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simulateData

predict(fit, doses = c(0, 1, 3, 4, 6, 8))

simulateData

Description

Function to simulate patient level data for a normally distributed endpoint

simulateData

Usage

```
simulateData(
  n_patients,
  dose_levels,
  sd,
  mods,
  n_sim = 1000,
  true_model = NULL,
  dr_means = NULL
)
```

Arguments

n_patients	Vector containing number of patients as a numerical value per dose-group.
dose_levels	Vector containing the different dosage levels.
sd	Standard deviation on patient level.
mods	An object of class "Mods" as specified in the DoseFinding package.
n_sim	Number of simulations to be performed, Default is 1000
true_model	Default value is NULL. Assumed true underlying model. Provided via a String. e.g. "emax". In case of NULL, all dose-response models, included in the mods input parameter will be used.
dr_means	a vector, with information about assumed effects per dose group. Default NULL.

Value

A list object, containing patient level simulated data for all assumed true models. Also providing information about simulation iteration, patient number as well as dosage levels.

Examples

```
models <- DoseFinding::Mods(linear</pre>
                                     = NULL,
                                   = NULL,
                           linlog
                                      = c(0.5, 1.2),
                           emax
                           exponential = 2,
                           doses
                                      = c(0, 0.5, 2,4, 8),
                           maxEff
                                       = 6)
dose_levels <- c(0, 0.5, 2,4, 8)
sd <- 12
n_patients <- c(40, 60, 60, 60, 60)
sim_data <- simulateData(n_patients = n_patients,</pre>
                        dose_levels = dose_levels,
                        sd
                                = sd,
                        su,
moαs = models,
n_sim = 100°
```

head(sim_data)

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