Package ‘clustDRM’

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Description Functions to identify the pattern of a dose-response curve. Then fit a set of appropriate models to it according to the identified pattern, followed by model averaging to estimate the effective dose.
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clustDRMapp

launch the shiny app for an easier use of the package

Description
launch the shiny app for an easier use of the package

Usage
clustDRMapp()

Details
The shiny app made for an easy use of the functionalities of the clustDRM package. It can be launched using command clustDRMapp(). It imports the data (in csv format), performs the clustering on it (for monotone patterns using E2 test, and for general patterns using ORICC2 and MCT). It also can estimate EDP for different dose-response curves using appropriate models. Plotting dose-response curves are also possible for any of these operations. A simulation tab will help to decide about the design of the study using a simulation study.

Author(s)
Vahid Nassiri and Yimer Wasihun.

clustDRMappSimple

launch the simpler version of the shiny app for an easier use of the package

Description
launch the simpler version of the shiny app for an easier use of the package

Usage
clustDRMappSimple()
Details

A simpler version of the shiny app made for an easy use of the functionalities of the clustDRM package. It can be launched using command clustDRMappSimple(). It imports the data (in csv format), performs the clustering on it (for monotone patterns using E2 test, and for general patterns using ORICC2 and MCT). It also can estimate EDp for different dose-response curves using appropriate models. Plotting dose-response curves are also possible for any of these operations.

Author(s)

Vahid Nassiri and Yimer Wasihun.

fitDRM

fitting dose-response model according to the identified pattern.

Description

Function to fit several dose-response candidate models according to the identified pattern, and combine their results using model selection and/or model averaging.

Usage

fitDRM(inputDataset, dose, response, ID, subsettingID = NULL, transform = c("none", "log", "sRoot", "qRoot", "boxcox"), addCovars = ~1, patternClusters, EDp = 0.5, addCovarsVar = TRUE, alpha = 0.05, na.rm = FALSE, imputationMethod = c("mean", "median"), nCores = 1)

Arguments

inputDataset a data frame containing the input dataset, it should at least include dose, response, and ID.

dose either a single string or a scalar, indicating the name of the dose column or its index.

response either a single string or a scalar, indicating the name of the response column or its index.

ID either a single string or a scalar, indicating the name of the ID column or its index.

subsettingID a vector of ID’s of the subjects, in case one wants to fit the models only to a subset of the data. Default is NULL, i.e., all the subjects in the inputDataset will be used.

transform single string indicating what kind of transform should be applied on the response data. It takes "none" (no transform, default), "log" (natural log), "sRoot" (square root), and "qRoot" (cubic root), and "boxcox" (Box-Cox transformation).

addCovars formula specifying extra linear covariate, e.g., ~x1+x2
patternClusters
a vector of the same length as the number of rows in inputData (number of subjects) indicating a pattern for each subject. Note that the keywords which are recognized are: "increasing", "decreasing", "flat", "complete", and "up down max at x" and "down up min at x", which x is one of the doses. The "flat" and "complete" patterns would not be considered.

EDp
scalar in (0,1), indicating with EDp should be computed, default is 0.5 (ED50).

addCovarsVar
logical variable (TRUE as default), indicating whether the variance of the extra covariates presented in addCovars (unless it is only intercept) should also be computed or not.

alpha
scalar in (0,1), level of significance with default alpha = 0.05.

na.rm
logical variable indicating whether missing values should be removed (TRUE) or not (FALSE, default).

imputationMethod
single string taking values from "mean" (default), and "median", which indicates how the missing values should be treated. "mean" would replace them with the mean of the observed ones, and "median" will use median of them for imputation.

nCores
scalar, indicating the number of cores should be used to perform LRT and MCT tests. Default is 1 which means sequential computation (no parallel computation).

Details
Note that the dose column of the inputDataset should be a numeric variable.

Value
an object of class fittedDRM which is a list with the following objects: fittedModels the outcome of DoseFinding::fitMod for all the suitable models estAICNonmonotone: the computed AIC for the models fitted to the subjects with a non-monotone pattern estEDpNonmonotone: the computed EDp for the models fitted to the subjects with a non-monotone pattern estAICMonotone: the computed AIC for the models fitted to the subjects with a monotone pattern estEDpMonotone: the computed EDp for the models fitted to the subjects with a monotone pattern extraCovarsMonotone: if any extra covariates are added to the model their estimates and possibly standard errors (if addCovarsVar = TRUE) are given for subjects with monotone pattern. extraCovarsNonmonotone: if any extra covariates are added to the model their estimates and possibly standard errors (if addCovarsVar = TRUE) are given for subjects with non-monotone pattern.

Author(s)
Vahid Nassiri, and Yimer Wasihun

See Also
DoseFinding
Examples

```r
## generating data
set.seed(11)
doses2Use <- c(0, 5, 20)
numRep2Use <- c(6, 3, 3)
generatedData <- cbind(rep(1, sum(numRep2Use)),
MCPMod::genDFdata("logistic", c(5, 3, 10, 0.05), doses2Use,
numRep2Use, 1),
matrix(rnorm(1*sum(numRep2Use)), sum(numRep2Use), 1))
colnames(generatedData) <- c("ID", "dose", "response", "x1")
for (iGen in 2:15){
genData0 <- cbind(rep(iGen, sum(numRep2Use)),
MCPMod::genDFdata("logistic", c(5, 3, 10, 0.05), doses2Use,
numRep2Use, 1), matrix(rnorm(1*sum(numRep2Use)),
sum(numRep2Use), 1))
colnames(genData0) <- c("ID", "dose", "response", "x1")
generatedData <- rbind(generatedData, genData0)
}
## transforming it for clustering
toInput <- inputDataMaker(2, 3, 1, generatedData)
## general pattern clustering
generalPatternClust <- generalPatternClustering(
  inputData = toInput$inputData, colsData = toInput$colsData ,
  colID = toInput$colID, doseLevels = toInput$doseLevels,
  numReplications = toInput$numReplicates, na.rm = FALSE,
  imputationMethod = "mean", ORICC = "two", transform = "none",
  plotFormat = "eps", LRT = TRUE, MCT = TRUE,
  adjustMethod = "BH", nPermute = 100, useSeed = NULL,
  theLeastNumberOfMethods = 2, alpha = 0.05, nCores = 1)
## fitDRM
fittedModel <- fitDRM (inputDataset = generatedData, dose = 2,
  response = 3, ID = 1, subsettingID = NULL,
  transform = c("none"), addCovars = ~x1,
  patternClusters =
generalPatternClust$clusteringORICC2$Results$clusteringResultsORICC2,
  EDp = 0.5, addCovarsVar = TRUE, alpha = 0.05, na.rm = FALSE,
  imputationMethod = c("mean"), nCores = 1)
```

---

generalPatternClustering

*Clustering dose-response curves based on their pattern*

---

Description

function to cluster dose-response curves based on their pattern.
generalPatternClustering

Usage

generalPatternClustering(inputData, colsData, colID, doseLevels,
numReplications, na.rm = FALSE, imputationMethod = c("mean",
"median"), ORICC = c("two", "one", "both"), transform = c("none",
"log", "sRoot", "qRoot", "boxcox"), plotFormat = c("eps", "jpg"),
LRT = TRUE, MCT = FALSE, adjustMethod = c("BH", "holm", "hochberg",
"hommel", "bonferroni", "BY", "fdr", "none"), nPermute = 1000,
useSeed = NULL, theLeastNumberOfMethods = c(1, 2, 3, 4),
alpha = 0.05, nCores = 1)

Arguments

inputData data matrix which should include ID’s of the subjects, as well as the measurements (gene expressions, etc.) for all replications of different as columns.

colsData vector indicating the index of columns in the inputData which correspond to the measurement for different replications of different doses.

colID scalar indicating the index of column corresponding to data ID.

doseLevels vector with dose levels.

numReplications vector with the same length as doseLevels with number of replications for each dose.

na.rm logical variable indicating whether missing values should be removed (TRUE) or not (FALSE, default)

imputationMethod signle string taking values from "mean" (default), and "median", which indicates how the missing values should be treated. "mean" would replace them with the mean of the observed ones, and "median" will use median of them for imputation.

ORICC signle string taking value "two", "one", and "both", indicating which ORICC procedure should be used. "one" refers to one-stage ORICC only, "two" (default) refers to two-stage ORICC only, and "both" will perform both of them.

transform single string indicating what kind of transform should be applied on the response data. It takes "none" (no transform, default), "log" (natural log), "sRoot" (square root), and "qRoot" (cubic root), and "boxcox" (Box-Cox transformation).

plotFormat plotFormat string gets two values "eps" (default), and "jpg" indicating the format of the output plot.

LRT logical indicating whether a permutation-based likelihood ratio test should be applied (TRUE) on the subjects which their trend is identified as non-flat by ORICC1 or not (FALSE).

MCT logical indicating whether a multiple comparison test (with "UmbrellaWilliams" contrast matrix) should be applied (TRUE) on the subjects which their trend is identified as non-flat by ORICC1 or not (FALSE).

adjustMethod The method for multiplicity adjustment for p-values. The possible values for this argument are "BH", "holm", "hochberg", "hommel", "bonferroni", "BY", "fdr", "none" with "BH" (Benjamini-Hochberg) as default.
**generalPatternClustering**

- **nPermute**: scalar indicating number of permutations in LRT.
- **useSeed**: scalar, indicating the seed should be used to generate LRT permutations. The default is NULL.
- **theLeastNumberOfMethods**: scalar taking values from 1, 2, 3, and 4, indicating how many methods should approve a non-flat trend that it can be selected. Its value depends on how many tests are asked to be done, for the maximum happens when ORICC = "both" and both LRT and MCT are TRUE. For example, when this argument sets to 2 and ORICC = "two", LRT = TRUE, and MCT = TRUE, it means if two-stage ORICC identifies a non-flat pattern and at least one of the LRT and MCT also accepts (at the level of alpha), then that compound is selected as one with a non-flat pattern. Note that the comparison with alpha is done for adjusted p-values.
- **alpha**: the significance level to compare the adjusted p-value with it.
- **nCores**: nCores scalar, indicating the number of cores should be used to perform LRT and MCT tests. Default is 1 which means sequential computation (no parallel computation).

**Details**

This function first use ORIC1 or ORIC2 (or both) to identify the pattern of the dose-response curve for each subject. Once the pattern is identified, for non-flat ones, a permutation-based likelihood ratio test (for exactly the identified pattern, if LRT = TRUE), and a multiple comparisons test (to test H0: flat vs. H1: non-flat, if MCT = TRUE) will be performed to further filter the flat patterns.

**Value**

a list of the following objects:

- **selectedSubjects**: a data frame indicating the ID’s of the selected subjects in the first columns and the identified trend in the second column.
- **clusteringORICC1Results** and/or **clusteringORICC2Results**: a list with four elements providing the raw data as the outcome of the ORICC procedure (rawDataORICC1 and/or rawDataORICC2), the pattern identified by the ORICC procedure (clusteringResultsORICC1 and/or clusteringResultsORICC2), results of LRT (resultsLRT) and results of MCT (resultsMCT). Both of them provide the adjusted and unadjusted p-values, but for MCT the selected contrast will be provided as well.

**Author(s)**

Vahid Nassiri, and Yimer Wasihan.

**See Also**

ORIClust ORIClust
Examples

```r
## generating data
set.seed(11)
doses2Use <- c(0, 5, 20)
numRep2Use <- c(3, 3, 3)
generatedData <- cbind(rep(1, sum(numRep2Use)),
MCPMod::genDFdata("logistic", c(5, 3, 10, 0.05), doses2Use,
numRep2Use, 1),
matrix(rnorm(1 * sum(numRep2Use)), sum(numRep2Use), 1))
colnames(generatedData) <- c("ID", "dose", "response", "x1")
for (iGen in 2:15){
genData0 <- cbind(rep(iGen, sum(numRep2Use)),
MCPMod::genDFdata("logistic", c(5, 3, 10, 0.05), doses2Use,
numRep2Use, 1),
matrix(rnorm(1 * sum(numRep2Use)), sum(numRep2Use), 1))
colnames(genData0) <- c("ID", "dose", "response", "x1")
generatedData <- rbind(generatedData, genData0)
}
## transforming it for clustering
toInput <- inputDataMaker(2, 3, 1, generatedData)

## general pattern clustering
generalPatternClust <- generalPatternClustering(inputData = toInput$inputData,
colsData = toInput$colsData, colID = toInput$colID,
doseLevels = toInput$doseLevels, numReplications = toInput$numReplicates,
na.rm = FALSE, imputationMethod = "mean", ORICC = "two", transform = "none", plotFormat = "eps",
LRT = TRUE, MCT = TRUE,
adjustMethod = "BH", nPermute = 100, useSeed = NULL,
theLeastNumberOfMethods = 2, alpha = 0.05, nCores = 1)
```

inputDataMaker  

Creating suitable `inputData` for clustering of the dose-response curve patterns

Description

function to create needed information as the input of the functions to cluster dose-response curve patterns.

Usage

```r
inputDataMaker(dose, response, ID, inputDataset)
```

Arguments

dose  
either a single string or a scalar, indicating the name of the dose column or its index.
inputDataMaker

response
either a single string or a scalar, indicating the name of the response column or its index.

ID
either a single string or a scalar, indicating the name of the ID column or its index.

inputDataset
a data frame containing the input dataset, it should at least include dose, response, and ID

Details
Note that the output of this function can be feed into the function for clustering dose-response curve patterns.

Value
a list with the following elements:
inputDataset: includes the ID (first column), and the response for all doses with their replications for each subject as rows. doseLevels: unique dose levels numReplications: number of replications per each unique dose level. colsData: the index of columns with responses. colID: the index of ID column.

Author(s)
Vahid Nassiri, and Yimer Wasihun

Examples

## generating data
set.seed(11)
doses2Use <- c(0, 5, 20)
umRep2Use <- c(3, 3, 3)
generatedData <- cbind(rep(1, sum(numRep2Use)),
MCPMod::genDFdata("logistic", c(5, 3, 10, 0.05),
doses2Use, numRep2Use, 1),
matrix(rnorm(1 * sum(numRep2Use)), sum(numRep2Use), 1))
colnames(generatedData) <- c("ID", "dose", "response", "x1")
for (iGen in 2:15){
genData0 <- cbind(rep(iGen, sum(numRep2Use)),
MCPMod::genDFdata("logistic", c(5, 3, 10, 0.05),
doses2Use, numRep2Use, 1),
matrix(rnorm(1 * sum(numRep2Use)), sum(numRep2Use), 1))
colnames(genData0) <- c("ID", "dose", "response", "x1")
generatedData <- rbind(generatedData, genData0)
}
## transforming it for clustering
toInput <- inputDataMaker(2, 3, 1, generatedData)
monotonePatternClustering

clustering dose-response curves based on their pattern when it is known to be monotone. function to cluster dose-response curves based on their pattern.

Description

clustering dose-response curves based on their pattern when it is known to be monotone. function to cluster dose-response curves based on their pattern.

Usage

monotonePatternClustering(inputData, colsData, colID, doseLevels, numReplications, transform = c("none", "log", "sRoot", "qRoot", "boxcox"), BHorBY = TRUE, SAM = FALSE, testType = c("E2", "Williams", "Marcus", "M", "ModifM"), adjustType = c("BH", "BY"), FDRvalue = c(0.05, 0.05), nPermute = c(1000, 1000), fudgeSAM = c("pooled", "none"), useSeed = c(NULL, NULL), theLeastNumberOfTests = 5, na.rm = FALSE, imputationMethod = c("mean", "median"))

Arguments

inputData  data matrix which should include ID's of the subjects, as well as the measurements (gene expressions, etc.) for all replications of different as columns.

colsData  vector indicating the index of columns in the inputData which correspond to the measurement for different replications of different doses.

colID  scalar indicating the index of column corresponding to data ID.

doseLevels  vector with dose levels.

numReplications  vector wit the same length as doseLevels with number of replications for each dose.

transform  single string indicating what kind of transform should be applied on the response data. It takes "none" (no transform, dafault), "log" (natural log), "sRoot" (square root), and "qRoot" (cubic root), and "boxcox" (Box-Cox transformation).

BHorBY  logical indicating whether monotonicity tests (specified in argument testType) using BH or BY modifications should be performed. Default is TRUE.

SAM  logical indicating whether a SAM procedure should be performed. Default is FALSE.

testType  string a subset of c("E2", "Williams", "Marcus", "M", "ModifM"), indicating the monotonicity tests which should be applied.

adjustType  method of adjustment for multi-plicity in case BHorBY = TRUE. It takes values "BH" abd "BY" with "BH" as the default.
monotonePatternClustering

FDRvalue a numerical vector of length 2 indicating the FDR values for BHOrBY and SAM, the default is 0.05 for both.

nPermuted a numerical vector of length 2 indicating number of permutation for BHOrBY and SAM, the default for both is 1000.

fudgeSAM single string takes value from ("pooled", "none") specified the fudge factor in SAM test statistic. The default is "pooled".

useSeed a vector of length two specifying the seed value for BHOrBY and SAM, the default is NULL for both.

theLeastNumberOfTests A scalar indicating the minimum number of tests which should approve a monotone trend to consider a trend monotone. The default is 5, i.e., all of the tests should agree on the monotonicity.

na.rm logical variable indicating whether missing values should be removed (TRUE) or not (FALSE, default)

imputationMethod single string taking values from "mean" (default), and "median", which indicates how the missing values should be treated. "mean" would replace them with the mean of the observed ones, and "median" will use median of them for imputation.

Value

a list with the following objects:

selectedSubjects: provides the ID and identified patterns for the subjects which are selected based on the results of various tests and theLeastNumberOfTests.

subjectsPatterns: a vector of the same length as the number of subjects in the input dataset which indicates the identified patterns for all subjects (including flat ones).

resultsBH: a list with the results of selected tests (if BHOrBY = TRUE, NULL otherwise).

resultsSAM: a list with results of SAM procedure (if SAM = TRUE, NULL otherwise).

selectedSubjectsBH: a data frame of all of the subjects with then number of tests select them based on adjusted BH or BY methods.

selectedSubjectsSAM: a data frame of all of the subjects with then number of tests select them based on SAM procedure

Author(s)

Vahid Nassiri, and Yimer Wasihun.

See Also

IsoGene IsoGene ORCME
Examples

```r
## generating data, a sample of size 20
set.seed(11)
doses2Use <- c(0, 5, 20)
numRep2Use <- c(3, 3, 3)
generatedData <- cbind(rep(1, sum(numRep2Use)),
  MCPMod::genDFdata("logistic", c(5, 3, 10, 0.05),
  doses2Use, numRep2Use, 1),
  matrix(rnorm(1 * sum(numRep2Use)), sum(numRep2Use), 1))
colnames(generatedData) <- c("ID", "dose", "response", "x1")
for (iGen in 2:20){
genData0 <- cbind(rep(iGen, sum(numRep2Use)),
  MCPMod::genDFdata("logistic", c(5, 3, 10, 0.05),
  doses2Use, numRep2Use, 1),
  matrix(rnorm(1 * sum(numRep2Use)), sum(numRep2Use), 1))
colnames(genData0) <- c("ID", "dose", "response", "x1")
generatedData <- rbind(generatedData, genData0)
}
## transforming it for clustering
toInput <- inputDataMaker(2, 3, 1, generatedData)
## monotone pattern clustering
monotonePatternClust <- monotonePatternClustering (inputData =
toInput$inputData, colsData = toInput$colsData ,
colID = toInput$colID, doseLevels = toInput$doseLevels,
numReplications = toInput$numReplicates,
BHorBY = TRUE, SAM = FALSE, testType = c("E2"),
adjustType = "BH", FDRvalue = c(0.05, 0.05),
nPermute = c(100, 100), fudgeSAM = "pooled",
useSeed = c(NULL, NULL), theLeastNumberOfTests = 1,
na.rm = FALSE, imputationMethod = "mean")
```

plotDoseResponseData  plot dose-response curves

Description

function to plot dose-response curves with the possibility of adding lines indicating average response per dose levels. Also, provided a pattern for the dose-response curve, it can estimate the expected mean values per dose level for the given pattern and add them to the plot.

Usage

```r
plotDoseResponseData(inputDataset, dose, response, ID, subjectID,
  xlab = "Dose", ylab = "Response", addMean = TRUE,
  drcPattern = NULL)
```
Arguments

inputDataset a data frame containing the input dataset, it should at least include dose, response, and ID
dose either a single string or a scalar, indicating the name of the dose column or its index.
response either a single string or a scalar, indicating the name of the response column or its index.
ID either a single string or a scalar, indicating the name of the ID column or its index.
subjectID single input as the same type as given ID column with the ID of the subject to plot.
xlab single string with default "dose", the label on x axis.
ylab single string with default "response", the label on y axis.
addMean logical variable indicating whether mean values (connecting with lines) should be plotted or not.
drcPattern single string showing the identified pattern using clustering algorithms. The default is NULL. In such case, no extra line will be added to the plot regarding the estimated means via the identified pattern.

Details

with addMean = TRUE, a line will be added to the plot, connecting the averaged response per dose level. But when a pattern is provided for the dose-response curve via drcPattern, then a line will be added to the data with the means estimated assuming the identified pattern. If both addMean = TRUE and drcPattern != NULL, then two lines will be added to the plot. The line in purplish-colored with cross signs as points is the averaged response value per dose level, and the bluish-colored line with circled cross signs as points represents the estimated mean based on the pattern.

Value

make a plot.

Author(s)

Vahid Nassiri and Yimer Wasihun.

Examples

```r
## generating data, a sample of size 20
set.seed(11)
doses2Use <- c(0, 5, 20)
numRep2Use <- c(3, 3, 3)
generatedData <- cbind(rep(1,sum(numRep2Use)),
MCPMod::genDFdata("logistic",c(5, 3, 10, 0.05), doses2Use,
numRep2Use, 1),
matrix(rnorm(1*sum(numRep2Use)), sum(numRep2Use), 1))
colnames(generatedData) <- c("ID", "dose", "response", "x1")
```

for (iGen in 2:15) {
    genData0 <- cbind(rep(iGen, sum(numRep2Use)),
        MCPMod::genDFdata("logistic", c(5, 3, 10, 0.05),
        doses2Use, numRep2Use, 1),
    matrix(rnorm(1*sum(numRep2Use)), sum(numRep2Use), 1))
    colnames(genData0) <- c("ID", "dose", "response", "x1")
    generatedData <- rbind(generatedData, genData0)
}
## plotting dose response relation
plotDoseResponseData(generatedData, 2, 3, 1, 2)
## transforming it for clustering
plotDoseResponseData(generatedData, 2, 3, 1, 2, 
    addMean = FALSE,
    drcPattern = "increasing")

plotSimulDRM

Description

A function to make a heatmap of the simulation results for the given measure.

Usage

plotSimulDRM(simulDRMobj, quantity2Plot = c("mean", "bias", "mse",
"variance", "relativeBias", "absBias", "absRelativeBias"))

Arguments

- simulDRMobj: output of simulEvalDRM function
- quantity2Plot: single string, the measure which should be plotted. Available choices are: c("mean", "bias", "mse", "variance", "relativeBias", "absBias", "absRelativeBias")

Value

A heatmap

Author(s)

Vahid Nassiri and Yimer Wasihun.

Examples

## generating data, a sample of size 20
set.seed(11)
doses2Use <- c(0, 5, 20)
numRep2Use <- c(3, 3, 3)
generatedData <- cbind(rep(1, sum(numRep2Use)),
        MCPMod::genDFdata("logistic", c(5, 3, 10, 0.05),
        doses2Use, numRep2Use, 1),
    matrix(rnorm(1*sum(numRep2Use)), sum(numRep2Use), 1))
    colnames(genData0) <- c("ID", "dose", "response", "x1")
    generatedData <- rbind(generatedData, genData0)
}
MCPMod::genDFdata("logistic",c(5, 3, 10, 0.05), doses2Use,
numRep2Use, 1),
matrix(rnorm(1*sum(numRep2Use)), sum(numRep2Use), 1))
colnames(generatedData) <- c("ID", "dose", "response", "x1")
for (iGen in 2:20){
genData0 <- cbind(rep(iGen,sum(numRep2Use)),
MCPMod::genDFdata("logistic",c(5, 3, 10, 0.05), doses2Use,
numRep2Use, 1),
matrix(rnorm(1*sum(numRep2Use)), sum(numRep2Use), 1))
colnames(genData0) <- c("ID", "dose", "response", "x1")
generatedData <- rbind(generatedData, genData0)
}
simRes <- simulEvalDRM (pilotData =
generatedData[generatedData$ID == 2, c(2,3)],
doseLevels = c(0, 4, 20),
numReplications = c(6, 3, 3), numSim = 10,
standardDeviation = 1, EDp = 0.5,
funcList = c("linlog", "emax", "sigEmax", "logistic"))
# plot the simulated results
plotSimulDRM(simRes, quantity2Plot = "mse")
**Value**

- `estsEDp`: a list of length of `funcList` providing the estimated EDp from models fitted to data generated from each model in `funcList`.
- `realEDp`: a vector of length `funcList`, the EDp’s computed based on the estimated parameters from different models fitted to `pilotData`.
- `bestModel`: a list of length `funcList`, a frequency table of best selected model for data generated from each model in `funcList`.
- `meanEDp`: a matrix showing mean of estimated EDp’s averaged over `numSim` replications.
- `biasEDp`: a matrix showing bias of estimated EDp’s averaged over `numSim` replications.
- `mseEDp`: a matrix showing MSE of estimated EDp’s averaged over `numSim` replications.
- `varEDp`: a matrix showing variance of estimated EDp’s averaged over `numSim` replications.
- `relativeBiasEDp`: a matrix showing relative bias of estimated EDp’s averaged over `numSim` replications.
- `absBiasEDp`: a matrix showing absolute bias of estimated EDp’s averaged over `numSim` replications.
- `absRelativeBiasEDp`: a matrix showing absolute bias of estimated EDp’s averaged over `numSim` replications.
- `averagedAIC`: a matrix showing AIC’s of different models averaged over `numSim` replications.
- `quantity2Plot`: which if needed will be passed to `plotSimulDRM` method.

The output of `simulEvalDRM` can be passed to the function `plotSimulDRM` to plot a heatmap for the desired the quantity of interest. Possible quantities are ("mean", "bias", "mse", "variance", "relativeBias", "absBias", "absRelativeBias")

**Author(s)**

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**Examples**

```r
## generating data, a sample of size 20
set.seed(11)
doses2Use <- c(0, 5, 20)
numRep2Use <- c(3, 3, 3)

generatedData <- cbind(rep(1,sum(numRep2Use)),
MCPMod::genDFdata("logistic",c(5, 3, 10, 0.05), doses2Use,
numRep2Use, 1),
matrix(rnorm(1*sum(numRep2Use)), sum(numRep2Use), 1))
colnames(generatedData) <- c("ID", "dose", "response", "x1")
for (iGen in 2:20){
genData0 <- cbind(rep(iGen,sum(numRep2Use)),
MCPMod::genDFdata("logistic",c(5, 3, 10, 0.05), doses2Use,
numRep2Use, 1),
matrix(rnorm(1*sum(numRep2Use)), sum(numRep2Use), 1))
colnames(genData0) <- c("ID", "dose", "response", "x1")
generatedData <- rbind(generatedData, genData0)
}
simRes <- simulEvalDRM (pilotData =
```
simulEvalDRM

generateData[generateData$ID == 2, c(2,3)],
doseLevels = c(0, 4, 20),
numReplications = c(6, 3, 3), numSim = 10,
standardDeviation = 1, EDp = 0.5,
funcList = c("linlog", "emax", "sigeMax", "logistic")
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