Package 'RIbench'

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Title Benchmark Suite for Indirect Methods for RI Estimation

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Depends R (>= 3.3.0)

Imports stats, optparse, digest, data.table, graphics, grDevices,

RColorBrewer

Suggests knitr, rmarkdown

Description

The provided benchmark suite enables the automated evaluation and comparison of any existing and novel indirect method for reference interval ('RI') estimation in a systematic way.

Indirect methods take routine measurements of diagnostic tests, containing pathological and non-pathological samples as input and use sophisticated

statistical methods to derive a model describing the distribution of the nonpathological samples, which can then be

used to derive reference intervals. The benchmark suite contains 5,760 simu-

lated test sets with varying difficulty.

To include any indirect method, a custom wrapper function needs to be provided.

The package offers functions for generating the test sets, executing the indirect method and evaluating the results.

See ?RIbench or vignette(``RIbench_package") for a more comprehensive description of the features.

A detailed description and application is described in Ammer T., Schuetzenmeister A., Prokosch H.-U., Zierk J., Rank C.M., Rauh M. ``RIbench: A Proposed Benchmark for the Standardized Evaluation of Indirect Methods for Reference Interval Estimation". Clinical Chemistry (2022) <doi:10.1093/clinchem/hvac142>.

License GPL (>= 3)

VignetteBuilder knitr, rmarkdown

NeedsCompilation no

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R topics documented:

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RIbench-package

RIbench-package

RIbench: Benchmark Suite for the Standardized Evaluation of Indirect Methods for Reference Interval Estimation

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Description

RIbench enables the automated evaluation and comparison of any existing and novel indirect method in a systematic way. Indirect methods take routine measurements of diagnostic tests, containing pathological and non-pathological samples as input and use sophisticated statistical methods to derive a model describing the distribution of the non-pathological samples, which can then be used to derive reference intervals. The benchmark suite contains 5,760 simulated data sets with varying difficulty. To include any indirect method, a custom wrapper function needs to be provided. The package offers functions for generating the test sets <code>generateBiomarkerTestSets</code>, executing the indirect method <code>evaluateBiomarkerTestSets</code> and evaluating the results <code>evaluateAlgorithmResults</code>.

Details

Package: RIbench
Type: Package
Version: 1.0.2
Date: 2022-11-25
License: GPL (>=3)
LazyLoad: yes

Author(s)

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References

Ammer, T., Schuetzenmeister, A., Prokosch, HU., Zierk, J., Rank, C.M., Rauh, M. RIbench: A Proposed Benchmark for the Standardized Evaluation of Indirect Methods for Reference Interval Estimation. Clin Chem (2022) [Accepted, July 12].

4 as.rgb

addGrid	Add a grid to an existing plot.	
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Description

It is possible to use automatically determined grid lines (x=NULL, y=NULL) or specifying the number of cells x = 3, y = 4 as done by grid. Additionally, x- and y-locations of grid-lines can be specified, e.g. x = 1:10, y = seq(0,10,2).

Usage

```
addGrid(x = NULL, y = NULL, col = "lightgray", lwd = 1L, lty = 3L)
```

Arguments

X	(integer, numeric) single integer specifies number of cells, numeric vector specifies vertical grid-lines
У	(integer, numeric) single integer specifies number of cells, numeric vector specifies horizontal grid-lines
col	(character) color of grid-lines
lwd	(integer) line width of grid-lines
lty	(integer) line type of grid-lines

Author(s)

Andre Schuetzenmeister <andre.schuetzenmeister@roche.com>

as.rgb	Convert color-names or RGB-code to possibly semi-transparent RGB-
	code.

Description

Function takes the name of a color and converts it into the rgb space. Parameter "alpha" allows to specify the transparency within [0,1], 0 meaning completey transparent and 1 meaning completey opaque. If an RGB-code is provided and alpha != 1, the RGB-code of the transparency adapted color will be returned.

Usage

```
as.rgb(col = "black", alpha = 1)
```

BoxCox 5

Arguments

col (character) name of the color to be converted/transformed into RGB-space (code).

Only those colors can be used which are part of the set returned by function col-

ors(). Defaults to "black".

alpha (numeric) value specifying the transparency to be used, 0 = completely trans-

parent, 1 = opaque.

Value

RGB-code

Author(s)

Andre Schuetzenmeister <andre.schuetzenmeister@roche.com>

BoxCox

One-parameter Box-Cox transformation.

Description

One-parameter Box-Cox transformation.

Usage

```
BoxCox(x, lambda)
```

Arguments

x (numeric) data to be transformed

lambda (numeric) Box-Cox transformation parameter

Value

(numeric) vector with Box-Cox transformation of x

Author(s)

Andre Schuetzenmeister <andre.schuetzenmeister@roche.com>

6 computeDirect

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Function to simulate the direct method

Description

Function to simulate the direct method

Usage

```
computeDirect(
   N = 120,
   analyte,
   params,
   seed = 123,
   NIter = 10000,
   RIperc = c(0.025, 0.975)
)
```

Arguments

N	(integer) specifying the number of samples used as sample size for the direct method, default: 120
analyte	(character) specifying the biomarker that is currently simulated
params	(list) of parameters for non-pathological distribution (nonp_mu, nonp_sigma, nonp_lambda, and nonp_shift)
seed	(integer) specifying the seed used for the simulation, default: 123
NIter	(integer) specifying the number of times N samples should be drawn out of the simulated non-pathological distribution (default: $10,000$)
RIperc	(numeric) value specifying the percentiles, which define the reference interval

Value

(data frame) with the estimated reference intervals for NIter iterations

Author(s)

computePerfMeas 7

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Function for computing performance measurements

Description

Function for computing performance measurements

Usage

```
computePerfMeas(
  analyte,
  algo,
  resRIs,
  subTable,
  RIperc = c(0.025, 0.975),
  cutoffZ = 5
)
```

Arguments

analyte	(character) specifiyng current analyzed analyte
algo	(character) specifying used algorithm
resRIs	(data.frame) with all calculated reference intervals
subTable	(data.frame) containing all information about the simulated test sets
RIperc	(numeric) vector specifying the percentiles for the reference interval, default: $0.025 \ \mathrm{and} \ 0.975$
cutoffZ	(numeric) specifying if a cutoff should be used to classify results as implausible and exclude from analysis

Value

updated data frame with computed performance measures

Author(s)

8 computeRIs

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compute	PerfMe	asAll

Function for computing reference intervals for all markers

Description

Function for computing reference intervals for all markers

Usage

```
computePerfMeasAll(analytes, algo, risIn, tableTCs, cutoffZ = 5)
```

Arguments

analytes	(character) listing all analytes for which the result files should be parsed
algo	(character) specifying used algorithm
risIn	(list) with data frame of all calculated reference intervals
tableTCs	(data.frame) containing all information about the simulated test sets
cutoffZ	(integer) specifying if and if so which cutoff should be used to classify results

as implausible (default: 5)

Value

list with the calculated errors as data frame for each marker

Author(s)

Tatjana Ammer <tatjana.ammer@roche.com>

 ${\tt computeRIs}$

Function for computing reference intervals

Description

Function for computing reference intervals

Usage

```
computeRIs(
  analyte,
  algo,
  results,
  tableTCs,
  RIperc = c(0.025, 0.975),
  truncNormal = FALSE
)
```

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Arguments

algo (character) specifying used algorithm

results (list) with all calculated results as RWDRI objects

tableTCs (data frame) containing all information about the simulated test sets

RIperc (numeric) vector specifying the percentiles for the reference interval, default:

0.025 and 0.975

truncNormal (logical) specifying if a normal distribution truncated at zero shall be assumed

Value

data frame with computed reference intervals

Author(s)

Tatjana Ammer <tatjana.ammer@roche.com>

computeRIsAll	Function for computing reference intervals for all markers
computeRIsAll	Function for computing reference intervals for all markers

Description

Function for computing reference intervals for all markers

Usage

```
computeRIsAll(analytes, algo, resIn, tableTCs, truncNormal = FALSE)
```

Arguments

analvtes	(-1	\ 1:	. 11	£ 1 - 1 - 41 -	14 C1	should be parsed

algo (character) specifying used algorithm

resIn (list) with all calculated results for all markers as RWDRI objects tableTCs (data.frame) containing all information about the simulated test sets

truncNormal (logical) specifying if a normal distribution truncated at zero shall be assumed

Value

list with the calculated reference intervals as data frame for each marker

Author(s)

10 computeSubResults

computeRuntimeAll

Function to compute runtime statistics for all analytes

Description

Function to compute runtime statistics for all analytes

Usage

```
computeRuntimeAll(analytes, algo, risIn, tableTCs)
```

Arguments

analytes (character) listing all analytes for which the result files should be parsed (character) specifying used algorithm

risIn (list) with data frame of all calculated reference intervals and runtime tableTCs (data.frame) containing all information about the simulated test cases

Value

(list) wit runtime statistics per analyte and data frames with raw runtime overall and per analyte

Author(s)

Tatjana Ammer <tatjana.ammer@roche.com>

computeSubResults

Helper function to compute the subscores for the distribution types and the mentioned categories

Description

Helper function to compute the subscores for the distribution types and the mentioned categories

Usage

```
computeSubResults(
  errorDf,
  tableTCs,
  distCat,
  errorParam,
  catList,
  catLabels,
  perfCombination = "mean"
)
```

defineSubset 11

Arguments

errorDf	(data frame) containing the estimate reference intervals and all computed error measures
tableTCs	(data.frame) containing all information about the simulated test sets
distCat	(character) specifying the distribution category
errorParam	(character) specifiying for which error parameter the data frame should be generated
catList	(character) vector containing the categories to split the dataset
catLabels perfCombinatio	(character) vector containing the labels that will be used for the categories

(character) specifying if mean (default), median or sum should be computed

Value

(data frame) containing the computed subscores

Author(s)

Tatjana Ammer <tatjana.ammer@roche.com>

defineSubset	Function for defining a subset that is used for analyizing the computa-
	tion time and can be used for other subanalyses.
	·

Description

Function for defining a subset that is used for analyzing the computation time and can be used for other subanalyses.

Usage

```
defineSubset(tableTCs = NULL, N = 50, seed = 123)
```

Arguments

tableTCs (data frame) describing the pre-defined testcases

N (integer) describing the number of testcases per biomarker contained in the sub-

set (default: 50)

seed (integer) specifying the seed used for defining the subset, default: 123

Value

(data frame) describing the updated table with all test case definitions.

Author(s)

evaluate Algorithm Results

Convenience Function to generate all result plots and calculate the benchmark score

Description

Convenience Function to generate all result plots and calculate the benchmark score

Usage

```
evaluateAlgorithmResults(
  workingDir = "",
  algoNames = NULL,
  subset = "all",
  evalFolder = "Evaluation",
  withDirect = TRUE,
  withMean = TRUE,
  outline = TRUE,
  errorParam = c("zzDevAbs_Ov", "AbsPercError_Ov", "AbsError_Ov"),
  cutoffZ = 5,
  cols = NULL,
  ...
)
```

workingDir	(character) specifying the working directory: Plots will be stored in 'workingDir/evalFolder' and results will be used from 'workingDir/Results/algoName/biomarker'
algoNames	(character) vector specifying all algorithms that should be part of the evaluation
subset	(character, numeric, or data.frame) to specify for which subset the algorithm should be evaluated. character options: 'all' (default) for all test sets, a distribution type: 'normal', 'skewed', 'heavilySkewed', 'shifted'; a biomarker: 'Hb', 'Ca', 'FT4', 'AST', 'LACT', 'GGT', 'TSH', 'IgE', 'CRP', 'LDH'; 'Runtime' for runtime analysis subset; numeric option: number of test sets per biomarker, e.g. 10; data.frame: customized subset of table with test set specifications
evalFolder	(character) specifying the name of the ouptut directory, Plots will be stored in workingDir/evalFolder, default: 'Evaluation'
withDirect	(logical) indicating whether the direct method should be simulated for comparison (default:TRUE)
withMean	(logical) indicating whether the mean should be plotted as well (default: TRUE)
outline	(logical) indicating whether outliers should be drawn (TRUE, default), or not (FALSE)
errorParam	(character) specifying for which error parameter the data frame should be generated, choose between absolute z-score deviation ("zzDevAbs_Ov"), absolute percentage error ("AbsPercError_Ov"), and absolute error ("AbsError_Ov")

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cutoffZ	(integer) specifying if and if so which cutoff for the absolute z-score deviation should be used to classify results as implausible and exclude them from the overall benchmark score (default: 5)
cols	(character) vector specifying the colors used for the different algorithms
	additional arguments to be passed to the method, e.g. "truncNormal" (logical) vector specifying if a normal distribution truncated at zero shall be assumed, can be either TRUE/FALSE or a vector with TRUE/FALSE for each algorithm; "colDirect" (character) specifying the color used for the direct method, default: "grey" "ylab" (character) specifying the label for the y-axis

Value

(data frame) containing the computed benchmark results

Author(s)

Tatjana Ammer <tatjana.ammer@roche.com>

Examples

```
## Not run:
# Ensure that 'generateBiomarkerTestSets()' and 'evaluateBiomarkerTestSets() is called
# with the same workingDir and for all mentioned algorithms before calling this function.

# first example, evaluation for several algorithms
benchmarkScore <- evaluateAlgorithmResults(workingDir=tempdir(),
algoNames=c("Hoffmann", "TML", "kosmic", "TMC", "refineR"))
# The function will create several plots saved in workingDir/Evaluation.

# second example, evaluation for only one algorithm and a defined subset
benchmarkScore <- evaluateAlgorithmResults(workingDir = tempdir(),
algoNames = "refineR", subset = 'Ca')

# third example, saving the results in a different folder, and setting a different cutoff
# for the absolute z-score deviation
benchmarkScore <- evaluateAlgorithmResults(workingDir = tempdir(), algoNames = "refineR",
subset = 'Ca', cutoffZ = 4, evalFolder = "Eval_Test")

## End(Not run)</pre>
```

evaluateBiomarkerTestSets

Wrapper function to evaluate all test sets or a specified subset for a specified algorithm.

Description

Wrapper function to evaluate all test sets or a specified subset for a specified algorithm.

Usage

```
evaluateBiomarkerTestSets(
  workingDir = "",
  algoName = "refineR",
  algoFunction = "findRI",
  libs = "refineR",
  sourceFiles = NULL,
  params = NULL,
  requireDecimals = FALSE,
  requirePercentiles = FALSE,
  subset = "all",
  timeLimit = 14400,
  verbose = TRUE,
  showWarnings = FALSE,
  ...
)
```

Arguments

workingDir (character) specifying the working directory: Results will be stored in 'work-

ingDir/Results/algo/biomarker' and data will be used from 'workingDir/Data/biomarker'

algoName (character) specifying the name of the algorithm that is evaluated

algoFunction (character) specifying the name of the function needed for estimating RIs

libs (list) containing all libraries needed for executing the algorithm sourceFiles (list) containing all source files needed for executing the algorithm params (list) with additional parameters needed for calling algoFunction

requireDecimals

(logical) indicating whether the algorithm needs the number of decimal places

(TRUE) or not (FALSE, default)

requirePercentiles

(logical) indicating whether only percentiles and no model is estimated

subset (character, numeric, or data.frame) to specify for which subset the algorithm

should be executed. character options: 'all' (default) for all test sets; a distribution type: 'normal', 'skewed', 'heavilySkewed', 'shifted'; a biomarker: 'Hb', 'Ca', 'FT4', 'AST', 'LACT', 'GGT', 'TSH', 'IgE', 'CRP', 'LDH'; 'Runtime' for runtime analysis subset; numeric option: number of test sets per biomarker, e.g. 10; data.frame: customized subset of table with test set specifications

timeLimit (integer) specifying the maximum amount of time in seconds allowed to execute

one single estimation (default: 14400 sec (4h))

verbose (logical) indictaing if the progress counter should be shown (default: TRUE)

showWarnings (logical) indicating whether warnings from the call to the indirect method/algorithm

should be shown (default: FALSE)

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additional arguments to be passed to the method, e.g. specified in- and output directory ('inputDir', 'outputDir')

Value

(data frame) containing information about the test sets where the algorithm terminated the R session or failed to report a result

Author(s)

Tatjana Ammer <tatjana.ammer@roche.com>

Examples

```
## Not run:
# The evaluation of all test sets can take several hours depending on
# the computation time of the algorithm.
# Wrapper function for indirect method required, see vignette("RIbench_package")
# Ensure that 'generateBiomarkerTestSets()' is called with the same workingDir
     before calling this function.
# first generic example
evaluateBiomarkerTestSets(workingDir = tempdir(), algoName = 'myOwnAlgo',
algoFunction = 'estimateModel', libs = c('myOwnAlgo'),
sourceFiles = list("C:\\Temp\\MyAlgoWrapper.R"),
requireDecimals = FALSE, requirePercentiles = FALSE,
subset ='all', timeLimit = 14400)
# second example, evaluation for only 'Calcium' test sets.
progress <- evaluateBiomarkerTestSets(workingDir = tempdir(), algoName = 'myOwnAlgo',</pre>
algoFunction = 'estimateModel', libs = c('myOwnAlgo'), subset = "Ca")
# third example, evaluation for only a subset testsets that follow a skewed distribution.
progress <- evaluateBiomarkerTestSets(workingDir = tempdir(), algoName = 'myOwnAlgo',</pre>
algoFunction = 'estimateModel', libs = c('myOwnAlgo'), subset = "skewed")
# forth example, evaluation for a subset of 3 testsets per biomarker.
progress <- evaluateBiomarkerTestSets(workingDir = tempdir(), algoName = 'myOwnAlgo',</pre>
algoFunction = 'estimateModel', libs = c('myOwnAlgo'), subset = 3)
# fifth example, evaluation for a customized subset with all test sets that have
# a pathological fraction <= 30%.
testsets <- loadTestsetDefinition()</pre>
progress <- evaluateBiomarkerTestSets(workingDir = tempdir(), algoName = 'myOwnAlgo',</pre>
algoFunction = 'estimateModel', libs = c('myOwnAlgo'),
subset = testsets[testsets$fractionPathol <= 0.3,] )</pre>
```

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```
# sixth example, evaluation forwarding additional parameters to the 'algoFunction'
progress <- evaluateBiomarkerTestSets(workingDir = tempdir(), algoName = 'myOwnAlgo',
algoFunction = 'estimateModel', libs = c('myOwnAlgo'),
sourceFiles = list("Test_RIEst_2pBoxCox"), params = list("model='2pBoxCox'"))

# seventh example, evaluation for indirect method that requires the number of
# decimal points as input
evaluateBiomarkerTestSets(workingDir = tempdir(), algoName = 'myOwnAlgo',
algoFunction = 'estimateModelDec', libs = c('myOwnAlgo'),
sourceFiles = "C:\\Temp\\Test_RIEst_dec.R", requireDecimals = TRUE)

# eigth example, evaluation for indirect method that directly estimates the percentiles
evaluateBiomarkerTestSets(workingDir = tempdir(), algoName="myOwnAlgo",
algoFunction="estimateRIs", libs="myOwnAlgo",
sourceFiles = "C:\\Temp\\Test_RIEst_RIEst_R", requirePercentiles=TRUE)

## End(Not run)</pre>
```

formatNumber

Rounding method with trailing zeros.

Description

Rounding method with trailing zeros.

Usage

```
formatNumber(x, digits)
```

Arguments

```
x (numeric) value that is roundeddigits (integer) indicating the number of decimal places to be used
```

Value

Rounded value with trailing zeros

Author(s)

Christopher Rank <christopher.rank@roche.com>

generateBiomarkerTestSets

Convenience function to generate simulated data and save each test set as a separate file

Description

Convenience function to generate simulated data and save each test set as a separate file

Usage

```
generateBiomarkerTestSets(
  workingDir = "",
  subset = "all",
  rounding = TRUE,
  verbose = TRUE
)
```

Arguments

workingDir (character) specifying the working directory where 'workingDir/Data/biomarker'

folders will be generated containing the simulated data

subset (character, numeric, or data.frame) to specify for which subset the data should

be generated and the algorithms later applied to. character options: 'all' (default) for all test sets; a distribution type: 'normal', 'skewed', 'heavilySkewed', 'shifted'; a biomarker: 'Hb', 'Ca', 'FT4', 'AST', 'LACT', 'GGT', 'TSH', 'IgE', 'CRP', 'LDH'; 'Runtime' for runtime analysis subset; numeric option: number of test sets per biomarker, e.g. 10; data.frame: customized subset of table with

test set specification

rounding (logical) indicating whether decimal places stated in test set specification should

be applied (default, TRUE), if FALSE, data will be rounded to 5 decimal places

to mimic unrounded data

verbose (logical) indicating if the progress counter should be shown (default: TRUE)

Value

No return value, instead the data files are generated and saved in the workingDir

Author(s)

Examples

```
## Not run:
workingDir <- "C:\\Temp\\RIbench\\"
generateBiomarkerTestSets(workingDir = workingDir)
## End(Not run)

# example generating a subset of 2 test sets per biomarker
generateBiomarkerTestSets(workingDir = tempdir(), subset = 2)</pre>
```

generateBoxPlotOneAnalyte

Wrapper function to generate one boxplot for a specified analyte

Description

Wrapper function to generate one boxplot for a specified analyte

Usage

```
generateBoxPlotOneAnalyte(
 errorListAll,
  colList,
 nameList,
  catList,
  catLabels,
  a,
 errorParam,
 outline = TRUE,
 withMean = TRUE,
 withCats = TRUE,
 withDirect = TRUE,
  titlePart = NULL,
  outputDir,
  filenamePart = NULL,
 ylim1 = c(0, 100),
 ylim2 = c(100, 1000),
)
```

Arguments

errorListAll	(list) containing the overall benchmark results per algorithm
colList	(character) vector specifying the colors used for the different algorithms (should correspond to columns of benchmark results)
nameList	(character) vector specifying the names used in the legend (should correspond to columns of benchmark results), if NULL, colnames will be used
catList	(character) vector specifying the categories for which the boxes should be drawn
catLabels	(character) vector specifying the labels to the associated categories used for the x-axis
a	(character) specifying the analyte for which the boxplot should be generated
errorParam	(charcter) specifying for which error measure the plot should be generated
outline	(logical) indicating whether outliers should be drawn (TRUE, default), or not (FALSE) $$
withMean	(logical) indicating whether the mean should be plotted as well (default: TRUE)
withCats	(logical) set to TRUE if categories (e.g. pathological fraction) should be plotted (default: $FALSE$)
withDirect	(logical) indicating whether the box of the direct method should be elongated to facilitate comparison (default:TRUE)
titlePart	(character) specifying the latter part of the title
outputDir	(character) specifying a output directory
filenamePart	(character) specifying a filename for the plot
ylim1	(numeric) vector specifying the limits in y-direction for the first granular scale
ylim2	(numeric) vector specifying the limits in y-direction for the second less detailed scale
• • •	additional arguments passed forward to other functions

Value

No return value. Instead, a plot is generated.

Author(s)

```
{\tt generateBoxplotsDistTypes}
```

Wrapper function to generate all boxplots for the specified distribution types split by defined categories

Description

Wrapper function to generate all boxplots for the specified distribution types split by defined categories

Usage

```
generateBoxplotsDistTypes(
  errorListAll,
  colList,
 nameList,
  catList,
  catLabels,
  errorParam = "zzDevAbs_Ov",
 outline = TRUE,
 withMean = TRUE,
 withDirect = TRUE,
 withCats = TRUE,
  titlePart = NULL,
 outputDir = NULL,
  filenamePart = NULL,
 ylim1Vec = NULL,
 ylim2Vec = NULL,
 yticks1Vec = NULL,
 yticks2Vec = NULL,
)
```

errorListAll	(list) containing the overall benchmark results per algorithm
colList	(character) vector specifying the colors used for the different algorithms (should correspond to columns of benchmark results)
nameList	(character) vector specifying the names used in the legend (should correspond to columns of benchmark results), if NULL, colnames will be used
catList	(character) vector specifying the categories for which the boxes should be drawn
catLabels	(character) vector specifying the labels to the associated categories used for the x-axis
errorParam	(charcter) specifying for which error measure the plot should be generated

outline	(logical) indicating whether outliers should be drawn (TRUE, default), or not (FALSE)
withMean	(logical) indicating whether the mean should be plotted as well (default: TRUE)
withDirect	(logical) indicating whether the box of the direct method should be elongated to facilitate comparison (default:TRUE)
withCats	(logical) set to TRUE if categories (e.g. pathological fraction) should be plotted (default: FALSE)
titlePart	(character) specifying the latter part of the title
outputDir	(character) specifying a output directory
filenamePart	(character) specifying a filename for the plot
ylim1Vec	(numeric) vector specifying the limits in y-direction for the first granular scale
ylim2Vec	(numeric) vector specifying the limits in y-direction for the second less detailed scale
yticks1Vec	(numeric) vector specifying the ticks in y-direction for the first granular scale
yticks2Vec	(numeric) vector specifying the ticks in y-direction for the second less detailed scale
	additional arguments passed forward to other functions

Value

No return value. Instead, a plot is generated.

Author(s)

Tatjana Ammer <tatjana.ammer@roche.com>

```
{\tt generateBoxplotsMultipleCats}
```

Wrapper function to generate all boxplots for the specified analytes split by defined categories

Description

Wrapper function to generate all boxplots for the specified analytes split by defined categories

Usage

```
generateBoxplotsMultipleCats(
   analytes,
   errorListAll,
   colList,
   nameList,
   category = c("fractionPathol", "fractionPathol_cum", "N", "N_cum", "OvFreq",
        "OvFreq_cum"),
```

```
catList = NULL,
catLabels = NULL,
errorParam = "zzDevAbs_Ov",
outline = TRUE,
withMean = TRUE,
withDirect = TRUE,
withCats = TRUE,
titlePart = NULL,
outputDir = NULL,
filenamePart = NULL,
ylim1Vec = NULL,
ylim2Vec = NULL,
yticks1Vec = NULL,
yticks2Vec = NULL,
...
)
```

analytes	(character) vector specifying for which analytes the plots should be generated
errorListAll	(named list) containing the overall benchmark results per algorithm (names of list elements should be the names of the algorithms)
colList	(character) vector specifying the colors used for the different algorithms (should correspond to columns of benchmark results)
nameList	(character) vector specifying the names used in the legend (should correspond to columns of benchmark results), if NULL, colnames will be used
category	(character) defining the category used for creating the subsets. All defined subfeatures are used for the categorization. Choose from "fractionPathol" (default), "N", or "OvFreq", individual or cumulative ("_cum"); if category is set this will be used to define catList and catLabels
catList	(character) vector specifying the categories for which the boxes should be drawn
catLabels	(character) vector specifying the labels to the associated categories used for the x-axis
errorParam	(charcter) specifying for which error measure the plot should be generated
outline	(logical) indicating whether outliers should be drawn (TRUE, default), or not (FALSE)
withMean	(logical) indicating whether the mean should be plotted as well (default: TRUE)
withDirect	(logical) indicating whether the box of the direct method should be elongated to facilitate comparison (default:TRUE)
withCats	(logical) set to TRUE if categories (e.g. pathological fraction) should be plotted (default: FALSE)
titlePart	(character) specifying the latter part of the title
outputDir	(character) specifying an output directory
filenamePart	(character) specifying a filename for the plot

generateDataFiles 23

ylim1Vec	(numeric) vector specifying the limits in y-direction for the first granular scale
ylim2Vec	(numeric) vector specifying the limits in y-direction for the second less detailed scale
yticks1Vec	(numeric) vector specifying the ticks in y-direction for the first granular scale
yticks2Vec	(numeric) vector specifying the ticks in y-direction for the second less detailed scale
	additional arguments passed forward to other functions

Value

No return value. Instead, a plot is generated.

Author(s)

Tatjana Ammer <tatjana.ammer@roche.com>

generateDataFiles	Generate simulated data with one start seed for each biomarker and save each test set as separate file
generateDataFiles	v

Description

Generate simulated data with one start seed for each biomarker and save each test set as separate file

Usage

```
generateDataFiles(
  tableTCs = NULL,
  outputDir = NULL,
  rounding = TRUE,
  verbose = TRUE
)
```

tableTCs	(data.frame) containing all information about the simulated test cases
outputDir	(character) specifying the output directory where the data files should be written to
rounding	(logical) indicating whether decimal places stated in tableTCs should be applied (default, TRUE), if FALSE, data will be rounded to 5 decimal places to mimic unrounded data
verbose	(logical) indicating if the progress counter should be shown (default: TRUE)

Value

No return value, instead the data files are generated

Author(s)

Tatjana Ammer <tatjana.ammer@roche.com>

generateMD5

Generate an MD5 hash sum for any R object.

Description

Generate an MD5 hash sum for any R object.

Usage

generateMD5(x)

Arguments

х

(object) any R object.

Value

(character) MD5 hash sum of the input object.

Author(s)

Christopher Rank < christopher.rank@roche.com>

 ${\tt generateScatterplotsAll}$

Wrapper function to generate scatterplots for the specified analytes

Description

Wrapper function to generate scatterplots for the specified analytes

Usage

```
generateScatterplotsAll(
  analytes,
  errorListAll,
  colList = NULL,
  nameList,
  tableTCs,
  errorParam = "zzDevAbs",
 withColorCat = NULL,
  titlePart = NULL,
  outputDir = NULL,
  filenamePart = NULL,
 ylim = NULL,
 xlim = NULL,
 xlab = NULL,
 ylab = NULL,
)
```

analytes	(character) vector specifying for which analytes the scatterplot should be generated
errorListAll	(list) containing the overall benchmark results per algorithm
colList	(character) vector specifying the colors used for the different algorithms (should correspond to columns of benchmark results)
nameList	(character) vector specifying the names used in the legend (should correspond to columns of benchmark results), if NULL, colnames will be used
tableTCs	(data frame) containing all test case information
errorParam	(character) specifying for which error measure the plot should be generated
withColorCat	(character) indicating if plot should be colored according to the pathological fraction ("fractionPathol"), sample size ("N"), or "overlapPatholLeft", "overlapPatholRight"
titlePart	(character) specifying the latter part of the title
outputDir	(character) specifying a output directory
filenamePart	(character) specifying a filename for the plot
ylim	(numeric) vector specifying the limits in y-direction for the first granular scale
xlim	(numeric) vector specifying the limits in y-direction for the second less detailed scale
xlab	(character) specifying x-axis label
ylab	(character) specifying y-axis label
	additional arguments passed forward to other functions

26 getBenchmarkResults

Value

No return value. Instead, a plot is generated.

Author(s)

Tatjana Ammer <tatjana.ammer@roche.com>

getBenchmarkResults

Computing benchmark table with the mean overall results.

Description

Computing benchmark table with the mean overall results.

Usage

```
getBenchmarkResults(
  errorList,
  nameVec,
  tableTCs,
  errorParam = "zzDevAbsCutoff_Ov",
  cutoffZ = 5,
  catList = c("fractionPathol <= 0.20 & N <= 5000",
        "fractionPathol <= 0.20 & N > 5000", "fractionPathol > 0.20 & N <= 5000",
        "fractionPathol > 0.20 & N > 5000"),
    catLabels = c("lowPlowN", "lowPhighN", "highPlowN", "highPhighN"),
    perfCombination = c("mean", "median", "sum")
)
```

errorList	(list) containing the the computed errors for the different (indirect) methods/algorithms	
nameVec	(character) vector specifying the names of the different (indirect) methods/algorithms	
tableTCs	(data.frame) containing all information about the simulated test sets	
errorParam	(character) specifying for which error parameter the data frame should be generated	
cutoffZ	(integer) specifying if and if so which cutoff for the absolute z-score deviation should be used to classify results as implausible and exclude them from the overall benchmark score (default: 5)	
catList	(character) vector containing the categories to split the dataset	
catLabels	(character) vector containing the labels that will be used for the categories	
perfCombination		
	(character) specifying which measure should be used to compute the overall benchmark score; choose from "mean" (default), "median", or "sum"	

getRI 27

Value

(data frame) containing the computed benchmark results

Author(s)

Tatjana Ammer <tatjana.ammer@roche.com>

getRI Method to calculate reference intervals (percentiles) for objects of class 'RWDRI'

Description

Method to calculate reference intervals (percentiles) for objects of class 'RWDRI'

Usage

```
getRI(
    x,
    RIperc = c(0.025, 0.975),
    CIprop = 0.95,
    pointEst = c("fullDataEst", "medianBS", "meanBS"),
    truncNormal = FALSE,
    Scale = c("original", "transformed")
)
```

Arguments

X	(object) of class 'RWDRI'
RIperc	(numeric) value specifying the percentiles, which define the reference interval
CIprop	(numeric) value specifying the central region for estimation of confidence intervals
pointEst	(character) specifying the point estimate determination: (1) using the full dataset ("fullDataEst"), (2) calculating the median from all bootstrap samples ("medianBS"), (2) works only if NBootstrap > 0 (3) calculating the mean from all bootstrap samples ("meanBS"), (3) works only if NBootstrap > 0
truncNormal	(logical) specifying if a normal distribution truncated at zero shall be assumed
Scale	(character) specifying if percentiles are calculated on the original scale ("Or") or the transformed scale ("Tr")

Value

(data.frame) with columns for percentile, point estimate and confidence intervals.

Author(s)

Christopher Rank christopher.rank@roche.com, Tatjana Ammer tatjana.ammer@roche.com>

28 getRuntime

getRIsAllwithoutModel Function for retrieving reference intervals if directly computed

Description

Function for retrieving reference intervals if directly computed

Usage

```
getRIsAllwithoutModel(analytes, algo, resIn, tableTCs)
```

Arguments

analytes (character) listing all markers for which the result files should be parsed

algo (character) specifying used algorithm

resIn (list) with all calculated results for all markers as RWDRI objects tableTCs (data.frame) containing all information about the simulated test sets

Value

list with the calculated reference intervals as data frame for each marker

Author(s)

Tatjana Ammer <tatjana.ammer@roche.com>

getRuntime Helper function to compute runtime statistics

Description

Helper function to compute runtime statistics

Usage

```
getRuntime(x, analyte)
```

Arguments

```
x (data.frame) with one column specifying the Runtime
analyte (character) specifying current analyzed marker
```

Value

(data.frame) containing runtime statistics (min, mean, median, max)

getSubset 29

Author(s)

Tatjana Ammer <tatjana.ammer@roche.com>

getSubset

Function to group the data according to a specified feature.

Description

The feature can either be the pathological fraction, the sample size or the overlap (category) individually or cumulative (_cum). For an individualized categorisation see getSubsetForDefinedCats.

Usage

```
getSubset(
   subsetDef,
   distType = FALSE,
   tableTCs,
   errorList,
   category = c("fractionPathol", "fractionPathol_cum", "N", "N_cum", "OvFreq",
        "OvFreq_cum"),
   restrict = NULL
)
```

Arguments

subsetDef	(character) listing either the analytes or distribution types for which the result files should be parsed
distType	(logical) indicating if parameter subsetDef refers to analytes (FALSE, default) or distribution types (TRUE)
tableTCs	(data.frame) containing all information about the simulated test sets
errorList	(list) containing for each method the table with the computed error measurements
category	(character) defining the category used for creating the subsets. All defined subfeatures are used for the categorization. Choose from "fractionPathol" (default), "N", or "OvFreq", individual or cumulative ("_cum")
restrict	(character) indicating whether test sets should be filtered according to specified restriction, default NULL, e.g. fractionPathol <= 0.30

Value

(list) containing the performance measurements grouped according to specified subset definition and categories

Author(s)

```
{\tt getSubsetForDefinedCats}
```

Function to group the data according to a specified feature.

Description

Function to group the data according to a specified feature.

Usage

```
getSubsetForDefinedCats(
   subsetDef,
   distType = FALSE,
   tableTCs,
   errorList,
   catList = NULL,
   catLabels = NULL,
   restrict = NULL
)
```

Arguments

subsetDef	(character) listing either the analytes or distribution types for which the result files should be parsed
distType	(logical) indicating if 'subsetDef' refers to analytes (FALSE, default) or distribution types (TRUE) $$
tableTCs	(data.frame) containing all information about the simulated test sets
errorList	(list) containing the table with the computed error measurements
catList	(list) containing the categories to split the dataset
catLabels	(list) containing the labels that will be used for the categories
restrict	(character) indicating whether testcases should be filtered according to specified restriction, default NULL, e.g. fractionPathol <= 0.30

Value

(list) containing the performance measurements grouped according to specified subset definition and categories

Author(s)

invBoxCox 31

invBoxCox

Inverse of the one-parameter Box-Cox transformation.

Description

Inverse of the one-parameter Box-Cox transformation.

Usage

```
invBoxCox(x, lambda)
```

Arguments

x (numeric) data to be transformed

lambda (numeric) Box-Cox transformation parameter

Value

(numeric) vector with inverse Box-Cox transformation of x

Author(s)

Andre Schuetzenmeister <andre.schuetzenmeister@roche.com>

loadTestsetDefinition Convenience function to load the table with the information about the pre-defined test sets

Description

Convenience function to load the table with the information about the pre-defined test sets

Usage

```
loadTestsetDefinition()
```

Value

(data frame) containing the pre-defined parameter combinations to generate the simulations

Author(s)

32 mergeAnalytes

Examples

```
testsets <- loadTestsetDefinition()
str(testsets)</pre>
```

mergeAnalytes

Function to combine analytes for defined categories

Description

The feature can either be the pathological fraction, the sample size or the overlap (category) individually or cumulative (_cum). For a individualized categorisation see getSubsetForDefinedCats.

Usage

```
mergeAnalytes(
  tableTCs,
  errorList,
  catList = NULL,
  catLabels = NULL,
  distTypes = TRUE
)
```

Arguments

tableTCs	(data.frame) containing all information about the simulated test sets
errorList	(list) containing for each method the table with the computed error measurements
catList	(list) containing the categories to split the dataset
catLabels	(list) containing the labels that will be used for the categories
distTypes	(logical) indicating if 'catList' refers to analytes (FALSE, default) or distribution types (TRUE)

Value

(list) containing the merged performance measurements grouped according to specified category

Author(s)

mergeSummaryErrors 33

mergeSummaryErrors

Helper function to combine all computed summary errors

Description

Helper function to combine all computed summary errors

Usage

```
mergeSummaryErrors(
  errorList,
  nameVec,
  errorParam = "MedianAbsPercErrorOV",
  cutoffZ = FALSE
)
```

Arguments

errorList (list) of the error lists for the different methods for which the summary errors

should be combined

nameVec (character) vector specifying the names of the methods

errorParam (character) specifying for which error parameter the data frame should be gen-

erated

cutoffZ (logical) indicating if a cutoff was set, needed for CRP case

Value

(data frame) containing the summary errors per analyte per method

Author(s)

Tatjana Ammer <tatjana.ammer@roche.com>

plotBarplot

Plot method for generating a barplot out of the benchmark results

Description

Plot method for generating a barplot out of the benchmark results

34 plotBarplot

Usage

```
plotBarplot(
  benchmarkRes,
  perDistType = FALSE,
  colList,
  nameList = NULL,
  withLabels = FALSE,
  withHorizLines = FALSE,
  title = NULL,
  xlim = NULL,
  xlab = "Mean of Absolute Z-Score Deviations",
  outputDir = NULL,
  filename = NULL,
  ...
)
```

Arguments

benchmarkRes	(data frame) containing the overall benchmark results
perDistType	(logical) indicating if one overall plot should be generated or if it should be separated by the distribution type
colList	$(character)\ vector\ specifying\ the\ colors\ used\ for\ the\ different\ algorithms\ (should\ correspond\ to\ columns\ of\ benchmark\ results)$
nameList	(character) vector specifying the names used in the legend (should correspond to columns of benchmark results), if NULL, colnames will be used
withLabels	(logical) indicating whether the corresponding values should be plotted as well (default: $FALSE$)
withHorizLines	(logical) indicating whether horizontal lines should be plotted for a better visual separation of the different categories (default:FALSE)
title	(character) specifying plot title
xlim	(numeric) vector specifying the limits in x-direction
xlab	(character) specifying the x-axis label
outputDir	(character) specifying a output directory
filename	(character) specifying a filename for the plot
	additional arguments passed forward to other functions

Value

No return value. Instead, a plot is generated.

Author(s)

plotBoxplot 35

plotBoxplot

Plot method for generating a boxplot of the benchmark results

Description

Plot method for generating a boxplot of the benchmark results

Usage

```
plotBoxplot(
  errorList,
  colList,
  nameList,
  outline = TRUE,
  withMean = TRUE,
  withCats = FALSE,
  withDirect = TRUE,
  title = "",
  outputDir = NULL,
  filename = NULL,
  ylim1 = c(0, 100),
  ylim2 = c(100, 1000),
  ...
)
```

errorList	containing the overall benchmark results
colList	(character) vector specifying the colors used for the different algorithms (should correspond to columns of benchmark results)
nameList	(character) vector specifying the names used in the legend (should correspond to columns of benchmark results), if NULL, colnames will be used
outline	(logical) indicating whether outliers should be drawn (TRUE, default), or not (FALSE)
withMean	(logical) indicating whether the mean should be plotted as well (default: TRUE)
withCats	(logical) set to TRUE if categories (e.g. pathological fraction) should be plotted (default: FALSE)
withDirect	(logical) indicating whether the box of the direct method should be elongated to facilitate comparison (default:TRUE)
title	(character) specifying plot title
outputDir	(character) specifying a output directory
filename	(character) specifying a filename for the plot
ylim1	(numeric) vector specifying the limits in y-direction for the first granular scale
ylim2	(numeric) vector specifying the limits in y-direction for the second less detailed scale
	additional arguments passed forward to other functions

36 plotScatterplot

Value

No return value. Instead, a plot is generated.

Author(s)

Tatjana Ammer <tatjana.ammer@roche.com>

plotScatterplot

Plot method for generating a scatterplot

Description

Plot method for generating a scatterplot

Usage

```
plotScatterplot(
  errorList,
  colList,
  nameList,
  withColor = NULL,
  cats = NULL,
  title = "",
  outputDir = NULL,
  filename = NULL,
  xlim = NULL,
  ylim = NULL,
  ylab = NULL,
  ...
)
```

errorList	(data frame) containing the overall benchmark results
colList	(character) vector specifying the colors used for the different algorithms (should correspond to columns of benchmark results)
nameList	(character) vector specifying the names used in the legend (should correspond to columns of benchmark results), if NULL, colnames will be used
withColor	(character) indicating if plot should be colored according to pathological fraction, sample size or pathological overlap left / right
cats	(character) specifying the category labels
title	(character) specifying plot title
outputDir	(character) specifying a output directory
filename	(character) specifying a filename for the plot

print.RWDRI 37

xlim	(numeric) vector specifying the limits in y-direction for the first granular scale
ylim	(numeric) vector specifying the limits in y-direction for the second less detailed scale
xlab	(character) specifying x-axis label
ylab	(character) specifying y-axis label
	additional arguments passed forward to other functions

Value

No return value. Instead, a plot is generated.

Author(s)

Tatjana Ammer <tatjana.ammer@roche.com>

print.RWDRI Standard print method for objects of class 'RWDRI'

Description

Standard print method for objects of class 'RWDRI'

Usage

```
## S3 method for class 'RWDRI'
print(
    x,
    RIperc = c(0.025, 0.975),
    CIprop = 0.95,
    pointEst = c("fullDataEst", "medianBS", "meanBS"),
    truncNormal = FALSE,
    ...
)
```

х	(object) of class 'RWDRI'
RIperc	(numeric) value specifying the percentiles, which define the reference interval
CIprop	(numeric) value specifying the central region for estimation of confidence intervals
pointEst	(character) specifying the point estimate determination: (1) using the full dataset ("fullDataEst"), (2) calculating the median from the bootstrap samples ("medianBS"), (2) works only if NBootstrap > 0 (3) calculating the mean from the bootstrap samples ("meanBS"), (3) works only if NBootstrap > 0
truncNormal	(logical) specifying if a normal distribution truncated at zero shall be assumed additional arguments passed forward to other functions.

38 readResultFiles

Value

No return value. Instead, a summary is printed.

Author(s)

Christopher Rank <christopher.rank@roche.com>

progressInd

Function for setting up the progress indicator.

Description

Function for setting up the progress indicator.

Usage

```
progressInd(value, maxValue, nCharMsg = 0)
```

Arguments

value (integer) indicating the current number
maxValue (integer) indicating the maximum number

nCharMsg (integer) indicating the number of characters the message already has

Value

(character) returing generated progress message

Author(s)

Tatjana Ammer <tatjana.ammer@roche.com>

readResultFiles

Function for reading in the result files for one marker

Description

Function for reading in the result files for one marker

Usage

```
readResultFiles(analyte, algo, path = NULL, tableTCs = NULL)
```

readResultFilesAll 39

Arguments

analyte (character) specifying analyte

algo (character) specifying used algorithm

path (character) specifying path to Results directories

tableTCs (data frame) containing all information about the simulated test sets

Value

list with caluclated results as RWDRI objects

Author(s)

Tatjana Ammer <tatjana.ammer@roche.com>

readResultFilesAll Function for reading all results files.

Description

Function for reading all results files.

Usage

```
readResultFilesAll(analytes, algo, baseDir = NULL, inputDir = NULL, tableTCs)
```

Arguments

analytes (character) listing all analytes for which the result files should be parsed

algo (character) specifying used algorithm

baseDir (character) specifying the baseDir: Results will be used from baseDir/Results/algo/marker

if baseDir is set, inputDir will be ignored; if baseDir is NULL, the current work-

ing directory will be used

inputDir (character) specifying path directly to Results directories

tableTCs (data frame) containing all information about the simulated test sets

Value

list with all caluclated results as RWDRI objects for each marker

Author(s)

read Results And Compute Errors

Function to read the result files and compute performance measures to create customized plots afterwards

Description

Function to read the result files and compute performance measures to create customized plots afterwards

Usage

```
readResultsAndComputeErrors(
  workingDir = getwd(),
  algoName = NULL,
  subset = "all",
  cutoffZ = 5,
  ...
)
```

Arguments

workingDir	(character) specifying the working directory: Plots will be stored in workingDir/evalFolder and results will be used from workingDir/Results/algoName/biomarker;
algoName	(character) vector specifying one algorithm for which the performance measures should be evaluated
subset	(character, numeric, or data.frame) to specify for which subset the algorithm should be executed. character options: 'all' (default) for all test sets, a distribution type: 'normal', 'skewed', 'heavilySkewed', 'shifted'; a biomarker: 'Hb', 'Ca', 'FT4', 'AST', 'LACT', 'GGT', 'TSH', 'IgE', 'CRP', 'LDH'; 'runtime' for runtime analysis subset; numeric option: number of test sets per biomarker, e.g. 10; data.frame: customized subset of table with test set specifications
cutoffZ	(integer) specifying if and if so which cutoff for the absolute z-score deviation should be used to classify results as implausible and exclude them from the overall benchmark score (default: 5)
•••	additional arguments to be passed to the method truncNormal (logical) specifying if a normal distribution truncated at zero shall be assumed

Value

(list) with (data frame) and a (list) with the computed performance measures

Author(s)

restrictSet 41

restrictSet	Function to get error subsets for defined category and restriction.

Description

Function to get error subsets for defined category and restriction.

Usage

```
restrictSet(overallCat, tableTCs, errorList, distType = TRUE, restrict = NULL)
```

Arguments

overallCat	(list) containing the categories to split the dataset
tableTCs	(data.frame) containing all information about the simulated test sets
errorList	(list) containing for each method the table with the computed error measurements
distType	(logical) indicating if 'overallCat' refers to analytes (FALSE, default) or distribution types (TRUE) $$
restrict	(character) indicating whether testcases should be filtered according to specified restriction, default NULL, e.g. fractionPathol <= 0.30

Value

(list) containing the merged performance measurements grouped according to specified category

Author(s)

Tatjana Ammer <tatjana.ammer@roche.com>

runDirectMethod	Convenience function to simulate the direct method for the specified subset
	swe ser

Description

Convenience function to simulate the direct method for the specified subset

Usage

```
runDirectMethod(tableTCs = NULL, N = 120, cutoffZ = 5)
```

runTC_usingRscript

Arguments

tableTCs (data frame) containing the pre-defined parameter combinations to generate the

simulations

N (integer) specifying the number of samples used as sample size for the direct

method, default: 120

cutoffZ (numeric) specifying if a cutoff should be used to classify results as implausible

and exclude from analysis

Value

(data frame) with computed performance measures

Author(s)

Tatjana Ammer <tatjana.ammer@roche.com>

Examples

```
# example to run direct method only for test sets for hemoglobin (Hb)
testsets <- loadTestsetDefinition()
directRes <- runDirectMethod(tableTCs = testsets[testsets$Analyte =="Hb",], N = 120, cutoffZ = 5)</pre>
```

runTC_usingRscript

Function for running test sets per algorithm per marker with calling Rscript for each test set

Description

Function for running test sets per algorithm per marker with calling Rscript for each test set

Usage

```
runTC_usingRscript(
  biomarker = NULL,
  algoName = "myOwnAlgo",
  algoFunction = "estimateModel",
  sourceFiles = NULL,
  libs = NULL,
  params = NULL,
  decimals = FALSE,
  ris = FALSE,
  RIperc = c(0.025, 0.975),
  tableTCs = NULL,
```

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```
outputDir = NULL,
inputDir = NULL,
timeLimit = 14400,
subsetDef = "all",
verbose = TRUE,
showWarnings = FALSE,
...
)
```

Arguments

biomarker	(character) specifying the biomarker for which the algorithm should calculate RIs
algoName	(character) specifying the name of the algorithm that is evaluated
algoFunction	(character) specifying the name of the function needed for estimating RIs
sourceFiles	(list) containing all source files needed for executing the algorithm
libs	(list) containing all libraries needed for executing the algorithm
params	(list) with additional parameters needed for calling algoFunction
decimals	(logical) indicating whether the algorithm needs the number of decimal places (TRUE) or not (FALSE, default)
ris	(logical) indicating whether only percentiles and no model is estimated
RIperc	(numeric) value specifying the percentiles, which define the reference interval
tableTCs	(data.frame) with the information about the simulated test sets
outputDir	$(character)\ specifying\ the\ output Dir:\ Results\ will\ be\ stored\ in\ output Dir/Results/algo/biomarker$
inputDir	(character) specifying the inputDir: Data files should be stored in inputDir/Data/biomarker
timeLimit	(integer) specifying the maximum amount of time in seconds allowed to execute one single estimation (default: 14400 sec (4h))
subsetDef	(character) describing the specified subset of all test sets the algorithm is applied to, used for naming the progress file
verbose	(logical) indictaing if the progress counter should be shown (default: TRUE)
showWarnings	(logical) indicating whether warnings from the call to the indirect method/algorithm should be shown (default: FALSE)
	additional arguments to be passed to the method

Value

(data frame) containing information about the test sets where the algorithm terminated the R session or failed to report a result

Author(s)

44 setupDirStructure

setupDirStructure	Convenience function to set up the directory structure used for storing data and results.

Description

Convenience function to set up the directory structure used for storing data and results.

Usage

```
setupDirStructure(
  outputDir = NULL,
  onlyData = FALSE,
  onlyResults = FALSE,
  tableTCs = NULL,
  algoName = NULL
)
```

Arguments

outputDir	(character) specifying the base output directory. From here, Data/biomarker and Result/algoName/biomarker directories are generated
onlyData	(logical) if set to TRUE, only the biomarker subdirectories are generated, name of output directory is used as it is (default:FALSE)
onlyResults	(logical) if set to TRUE, only the algoName/biomarker subdirectories are generated, name of output directory is used as it is (default:FALSE)
tableTCs	(data frame) containing the pre-defined parameter combinations to generate the simulations
algoName	(character) specifying the name of the algorithm used for creating the subdirectory

Value

No return value. Instead, the directory structure is set up.

Author(s)

writeResFile 45

writeResFile	Helper function to write result file when time out occured or R session terminated

Description

Helper function to write result file when time out occured or R session terminated

Usage

```
writeResFile(
  algoName,
  biomarker,
  N = 0,
  error = NULL,
  runtime = NULL,
  filename = NULL,
  outputDir = NULL)
```

Arguments

algoName	(character) specifying the name of the algorithm that is evaluated
biomarker	(character) specifying the biomarker for which the algorithm should calculate RIs
N	(numeric) specifying the number of input data points
error	(character) specifying the type of error (e.g. timeout, RSessionTerminated)
runtime	(numeric) specifying the computation time up until the error occured
filename	(character) specifying the filename for which the algorithm failed
outputDir	(character) specifying the outputDir: Data files should be stored in output-Dir/Data/biomarker and Results will be stored in outputDir/Results/algo/biomarker

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