

Package ‘SummarizedBenchmark’

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Type Package

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Description This package defines the BenchDesign and SummarizedBenchmark classes for building, executing, and evaluating benchmark experiments of computational methods. The SummarizedBenchmark class extends the RangedSummarizedExperiment object, and is designed to provide infrastructure to store and compare the results of applying different methods to a shared data set. This class provides an integrated interface to store metadata such as method parameters and software versions as well as ground truths (when these are available) and evaluation metrics.

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<code>.modmethod</code>	<i>Modify BenchDesign Method</i>
-------------------------	----------------------------------

Description

Given a method defined in a BenchDesign, this helper function returns a modified method with new parameters defined as a list of quosures.

Usage

`.modmethod(m, q, .overwrite)`

Arguments

- | | |
|-------------------------|---|
| <code>m</code> | method |
| <code>q</code> | quosure list of new parameters |
| <code>.overwrite</code> | logical whether to overwrite parameters |

Value

modified method.

Author(s)

Patrick Kimes

addMethod

Add method to BenchDesign object

Description

Takes a [BenchDesign](#) object and the definition of a new method for benchmarking and returns the original [BenchDesign](#) with the new method included.

At a minimum, a method label (label =), and the workhorse function for the method (func =) must be specified for the new method.

Parameters for the method must be specified as a [quos](#) named list of parameter = value pairs mapping entries in the benchmarking data to the function parameters. For users familiar with the [ggplot2](#) package, this can be viewed similar to the aes = mapping of data to geometry parameters.

An optional secondary function, post, can be specified if the output of the workhorse function, func, needs to be further processed. As an example, post may be a simple "getter" function for accessing the column of interest from the large object returned by func.

Usage

```
addMethod(bd, label, func, params = rlang::quos(), post = NULL,
          meta = NULL)
```

Arguments

bd	BenchDesign object.
label	Character name for the method.
func	Primary function to be benchmarked.
params	Named quosure list created using quos of parameter = value pairs to be passed to func.
post	Optional post-processing function that takes results of func as input. Ignored if NULL. If multiple assays (metrics) should be generated for each method, this can be accomplished by specifying a named list of post-processing functions, one for each assay. (default = NULL)
meta	Optional metadata information for method to be included in colData of SummarizedBenchmark object generated using <code>link{buildBench}</code> . See Details for more information. Ignored if NULL. (default = NULL)

Details

The optional meta parameter accepts a named list of metadata tags to be included for the method in the resulting [SummarizedBenchmark](#) object. This can be useful for two primary cases. First, it can help keep analyses better organized by allowing the specification of additional information that should be stored with methods, e.g. a tag for "method type" or descriptive information on why the method was included in the comparison. Second, and more importantly, the meta parameter can be used to overwrite the package and version information that is automatically extracted from the function specified to func. This is particularly useful when the function passed to func is a

wrapper for a script in (or outside of) R, and the appropriate package and version information can't be directly pulled from `func`. In this case, the user can either manually specify the `"pkg_name"` and `"pkg_vers"` values to `meta` as a list, or specify a separate function that should be used to determine the package name and version. If a separate function should be used, it should be passed to `meta` as a list entry with the name `pkg_func` and first quoted using `quo`, e.g. `list(pkg_func = quo(p.adjust))`.

Value

Modified `BenchDesign` object with new method added.

Author(s)

Patrick Kimes

See Also

`modifyMethod`, `expandMethod`, `dropMethod`

Examples

```
## create example data set of p-values
df <- data.frame(pval = runif(100))

## example calculating qvalue from pvalues

## using standard call
qv <- qvalue::qvalue(p = df$pval)
qv <- qv$qvalue

## adding same method to BenchDesign
bench <- BenchDesign(data = df)
bench <- addMethod(bench,
  label = "qv",
  func = qvalue::qvalue,
  post = function(x) { x$qvalue },
  params = rlang::quos(p = pval))
```

`addPerformanceMetric` *Add performance metric to SummarizedBenchmark object*

Description

This is a function to define performance metrics for benchmarking methods. The function is saved into the `performanceMetrics` slot.

Usage

```
addPerformanceMetric(object, evalMetric, assay, evalFunction = NULL)
```

Arguments

object	A SummarizedBenchmark object.
evalMetric	A string with the name of the evaluation metric.
assay	A string with an assay name. Indicates the assay that should be given as input to this performance metric.
evalFunction	A function that calculates a performance metric. It should contain at least two arguments, query and truth, where query is the output vector of a method and truth is the vector of true values. If additional parameters are specified, they must contain default values. If NULL, the 'evalMetric' string must be the name of a predefined metric available through 'availableMetrics()\$function'.

Value

A [SummarizedBenchmark](#) object.

Author(s)

Alejandro Reyes

See Also

[availableMetrics](#), [performanceMetrics](#)

Examples

```
data( sb )
sb <- addPerformanceMetric(
  object=sb,
  assay="qvalue",
  evalMetric="TPR",
  evalFunction = function( query, truth, alpha=0.1 ){
    goodHits <- sum( (query < alpha) & truth == 1 )
    goodHits / sum(truth == 1)
  }
)
```

allSB

SummarizedBenchmark object of isoform quantification results

Description

This object is a [SummarizedBenchmark](#) object containing isoform quantifications from salmon, sailfish and kallisto from 4 mouse samples (2 hearts and 2 brains) part of the Mouse BodyMap. Its generation is described in one of the vignettes of this package.

Source

Mouse BodyMap (Li et al, 2014). SRA accession numbers SRR5273705, SRR5273689, SRR5273699 and SRR5273683.

Examples

```
data(quantSB)
```

```
assayNames<- , SummarizedBenchmark, character-method
```

Set assay names in SummarizedBenchmark object

Description

Modifies the assay names of a [SummarizedBenchmark](#) object.

Usage

```
## S4 replacement method for signature 'SummarizedBenchmark, character'
assayNames(x, ...) <- value
```

Arguments

<code>x</code>	A SummarizedBenchmark object.
<code>...</code>	Futher arguments, perhaps used by methods.
<code>value</code>	A character vector.

Value

Modified [SummarizedBenchmark](#) object.

Author(s)

Alejandro Reyes

See Also

[SummarizedBenchmark](#)

Examples

```
data(sb)
assayNames(sb)[2] <- "log2FC"
```

availableMetrics	<i>List pre-defined metrics for SummarizedBenchmark objects</i>
------------------	---

Description

This function returns a data frame summarizing the default performance metrics provided in this package. The data.frame contains three columns, functions is the name of the performance metric, description is longer description of the performance metric and requiredTruth is logical depending on whether the performance metrics require ground truths.

Usage

```
availableMetrics()
```

Value

A data.frame summarizing the default performance metrics provided in this package.

Author(s)

Alejandro Reyes

Examples

```
availableMetrics()
```

BDDData	<i>Create a new BDDData object</i>
---------	------------------------------------

Description

Initializes a new BDDData object of benchmarking data.

Data sets are stored as BDDData objects within BenchDesign objects as well as SummarizedBenchmark objects. However, because data is directly specified to the BenchDesign constructor, there is usually no need to call the BDDData constructor to create completely new data objects.

The BDDData constructor is most useful for extracting the data sets contained in BenchDesign objects as well as SummarizedBenchmark objects. By default, the BDDData object stored in SummarizedBenchmark objects will be MD5 hashes rather than the complete original data set. [compareBDDData](#) can be used to compare various forms of BDDData, as shown in the examples below.

Usage

```
BDData(data)

## S4 method for signature 'ANY'
BDData(data)

## S4 method for signature 'BenchDesign'
BDData(data)

## S4 method for signature 'SummarizedBenchmark'
BDData(data)

## S4 method for signature 'BDData'
BDData(data)
```

Arguments

data a list object of data or MD5 hash string

Value

BDData object

Author(s)

Patrick Kimes

See Also

[BDData-class](#), [BenchDesign](#)

Examples

```
## construct from data.frame
datadf <- data.frame(x = 1:5, y = runif(5))
bdd_df <- BDData(datadf)
bdd_df

## construct from MD5 hash of data.frame
bdd_md5 <- BDData(digest::digest(datadf))
bdd_md5

## compare two BDData objects
compareBDData(bdd_df, bdd_md5)

## note that the data is the same, i.e. the MD5 hashes match, but the
## data types ("data" vs. "md5has") are different
```

BDData-class

BDData class

Description

Container for data in a BenchDesign object.

Slots

data a list or MD5 hash of the data.

type a character string indicating whether the data slot contains the 'data' or a 'md5hash' of the data.

Author(s)

Patrick Kimes

See Also

[BDData](#), [BenchDesign-class](#), [BDMethod-class](#), [BDMethodList-class](#)

BDData<-

Set data in BenchDesign object

Description

Adds, removes or replaces [BDData](#) in [BenchDesign](#) object. Data can be removed by setting the value to NULL.

Usage

```
BDData(x) <- value
```

```
## S4 replacement method for signature 'BenchDesign,BDDataOrNULL'
BDData(x) <- value
```

Arguments

x [BenchDesign](#) object.

value [BDData](#) or NULL.

Value

modified BenchDesign object

Author(s)

Patrick Kimes

See Also[BDDData](#)**Examples**

```
bd <- BenchDesign()
BDDData(bd) <- BDDData(data.frame(x1 = runif(5)))
bd
```

BDMMethod

*Create a new BDMethod object***Description**

Initializes a new BenchDesign method object for benchmarking.

New [BDMethod](#) objects are typically not directly constructed as they have limited use outside of [BenchDesign](#) objects. Instead, methods in a [BenchDesign](#) object are more commonly created, modified or removed using function calls on the [BenchDesign](#), e.g. using [addMethod](#) to add a new method object.

The constructor can also be used to access [BDMethod](#) objects stored in [BDMethodList](#) and [BenchDesign](#) objects.

Usage

```
BDMethod(x, params = rlang::quos(), post = NULL, meta = NULL, ...)
```

```
## S4 method for signature 'quosure'
```

```
BDMethod(x, params = rlang::quos(), post = NULL,
  meta = NULL, ...)
```

```
## S4 method for signature ``function`'
```

```
BDMethod(x, params = rlang::quos(), post = NULL,
  meta = NULL, ...)
```

```
## S4 method for signature 'BDMethodList'
```

```
BDMethod(x, i = 1)
```

```
## S4 method for signature 'BenchDesign'
```

```
BDMethod(x, i = 1)
```

Arguments

x	main method function or function quosure. Alternative, may be a BDMethodList or BenchDesign object from which the BDMethod should be extracted.
params	list of quosures specifying function parameters. (default = <code>rlang::quos()</code>)
post	list of functions to be applied to the output of x. (default = <code>NULL</code>)
meta	list of metadata. (default = <code>NULL</code>)
...	other parameters.
i	integer index or character name of BDMethod in BDMethodList or BenchDesign object.

Value

BDMethod object

Author(s)

Patrick Kimes

See Also

[BDMethod-class](#), [BenchDesign](#), [BDMethodList](#)

Examples

```
## create a simple BDMethod
bdm1 <- BDMethod(x = base::mean)

## create a more complex BDMethod
bdm2 <- BDMethod(x = function(x) { x^2 }, post = base::sqrt,
                 meta = list(note = "simple example"))

## construct a BenchDesign with the BDMethod objects
bd <- BenchDesign(method1 = bdm1, method2 = bdm2)

## access a BDMethod in the BenchDesign
BDMethod(bd, "method1")
```

BDMethod-class

BDMethod class

Description

Container for individual methods to be compared as part of a benchmark experiment defined in a [BenchDesign](#) object. In the SummarizedBenchmark framework, methods are defined by a unique combination of functions, parameters, and any relevant meta data.

New BDMethod objects can be created using the [BDMethod](#) constructor.

Slots

f a function to be benchmarked
fc a captured expression of the function **f**
params a list of quosures specifying function parameters
post a list of functions to be applied to the output of **f**
meta a list of meta data

Author(s)

Patrick Kimes

See Also

[BDMETHOD](#), [BenchDesign-class](#), [BDMETHODList-class](#), [BDDData-class](#)

 BDMETHOD<-

Set method in list or BenchDesign object

Description

Adds, replaces or removes a named [BDMETHOD](#) method in a [BDMETHODList](#) or [BenchDesign](#) object with a specified [BDMETHOD](#) object.

An existing method can be removed by setting the value to NULL.

Usage

```

BDMETHOD(x, i) <- value

## S4 replacement method for signature 'BDMETHODList,character,BDMETHOD'
BDMETHOD(x, i) <- value

## S4 replacement method for signature 'BDMETHODList,character,`NULL`'
BDMETHOD(x, i) <- value

## S4 replacement method for signature 'BenchDesign,character,BDMETHOD'
BDMETHOD(x, i) <- value

## S4 replacement method for signature 'BenchDesign,character,`NULL`'
BDMETHOD(x, i) <- value

```

Arguments

x [BenchDesign](#) or [BDMETHODList](#) object.
i character name of method.
value [BDMETHOD](#) or NULL.

Value

modified BenchDesign object

Author(s)

Patrick Kimes

See Also

[BDMethod](#)

Examples

```
bd <- BenchDesign()
BDMethod(bd, "avg") <- BDMethod(x = base::mean)
bd
```

BDMethodList

Create a new BDMethodList object

Description

Initializes a new SimpleList of BenchDesign method ([BDMethod](#)) objects.

Similar to [BDMethod](#) objects, [BDMethodList](#) typically do not need to be directly constructed. Because the list of methods is only useful as part of a [BenchDesign](#) object, it is more common to simply manipulate the list of methods through calls to the corresponding [BenchDesign](#), e.g. [addMethod](#) to add a new method to the list.

The constructor can also be used to access the [BDMethodList](#) list of methods in a [BenchDesign](#) object.

Usage

```
BDMethodList(..., x = NULL)

## S4 method for signature 'ANY'
BDMethodList(..., x = NULL)

## S4 method for signature 'BenchDesign'
BDMethodList(..., x = NULL)

## S4 method for signature 'SummarizedBenchmark'
BDMethodList(..., x = NULL)
```

Arguments

... a named list of [BDMETHOD](#) objects

x a [BenchDesign](#) or [SummarizedBenchmark](#) object to extract the list of methods from. (default = NULL)

Value

BDMETHODList object

Author(s)

Patrick Kimes

See Also

[BDMETHODList-class](#), [BenchDesign](#), [BDMETHOD](#)

Examples

```
## construct an empty list
bdml <- BDMETHODList()

## construct a list with BDMETHOD objects
bdml <- BDMETHODList(m_method = BDMETHOD(base::mean),
                    s_method = BDMETHOD(function(x) { x^2 })))
bdml

## construct a BenchDesign with a BDMETHODList
bd <- BenchDesign(methods = bdml)

## access the BDMETHODList in the BenchDesign
BDMETHODList(bd)
```

BDMETHODList-class	<i>BDMETHODList class</i>
--------------------	---------------------------

Description

Extension of the SimpleList class to contain a list of BDMETHOD objects. The class serves as the primary container for the set of methods in the BenchDesign class.

New BDMETHODList objects can be created using the [BDMETHODList](#) constructor.

Author(s)

Patrick Kimes

See Also

[BDMETHODList](#), [BenchDesign-class](#), [BDMETHOD-class](#), [BDDData-class](#)

<code>BDMETHODList<-</code>	<i>Set method list in BenchDesign object</i>
--------------------------------	--

Description

Replaces the [BDMETHODList](#) list of methods in a [BenchDesign](#) object with a specified [BDMETHODList](#) object.

Usage

```
BDMETHODList(x) <- value

## S4 replacement method for signature 'BenchDesign,BDMETHODList'
BDMETHODList(x) <- value
```

Arguments

<code>x</code>	BenchDesign object.
<code>value</code>	BDMETHODList list of methods.

Value

modified [BenchDesign](#) object

Author(s)

Patrick Kimes

See Also

[BDMETHOD](#), [BDMETHODList](#)

Examples

```
bd <- BenchDesign()
BDMETHODList(bd) <- BDMETHODList(avg = BDMETHOD(x = base::mean))
bd
```

BenchDesign

*Create a new BenchDesign object***Description**

Initializes a new BenchDesign object of benchmarking methods and data.

The BenchDesign class serves as the core container for methods and data used for benchmarking in the SummarizedBenchmark package. The object can be initialized with a list of methods to be benchmarked, a default benchmarking data set, both or neither. Methods must be passed to the constructor as BDMethod or BDMethodList objects.

The constructor can also be used to access the BenchDesign stored in a SummarizedBenchmark object.

Usage

```
BenchDesign(..., methods = NULL, data = NULL)
```

```
## S4 method for signature 'ANY'
```

```
BenchDesign(..., methods = NULL, data = NULL)
```

```
## S4 method for signature 'SummarizedBenchmark'
```

```
BenchDesign(methods, data)
```

Arguments

...	named set of BDMethod objects and/or unnamed BenchDesign objects. Only the methods of any BenchDesign object will be used, and the data slot of the objects will be ignored.
methods	named set of BDMethod objects and/or unnamed BenchDesign objects as a list. (default = NULL)
data	optional data.frame or other list object to be used in the benchmark. (default = NULL)

Value

BenchDesign object.

Author(s)

Patrick Kimes

See Also

[BenchDesign-class](#), [BDMethod](#), [BDMethodList](#)

Examples

```
## with no input
bd <- BenchDesign()

## with data - data must be a named argument
datadf <- data.frame(pval = runif(20), x1 = rnorm(20))
bd <- BenchDesign(data = datadf)

## with two methods and data
method_bh <- BDMethod(stats::p.adjust, params = rlang::quos(p = pval, method = "BH"))
method_bf <- BDMethod(stats::p.adjust, params = rlang::quos(p = pval, method = "bonferroni"))
bd <- BenchDesign(bh = method_bh, bonf = method_bf,
                  data = datadf)

## with BDMethodList and data
bdml <- BDMethodList(bh = method_bh, bonf = method_bf)
bd <- BenchDesign(methods = bdml, data = datadf)
```

BenchDesign-class	<i>BenchDesign class</i>
-------------------	--------------------------

Description

Along with the SummarizedBenchmark class, one of the two main classes of the Summarized-Benchmark package. The BenchDesign class serves as a container for both the set of methods to be benchmarked and optionally the data to be used for benchmarking.

Methods are organized as BDMethod objects and stored in as a list using the BDMethodList class. The BDData class is used to store benchmarking data, or in some cases, just the MD5 hash of the original data set. Any list object, including data.frame objects, can be specified for data. More details on the component classes are provided in the corresponding class documentation.

For details on how to create new BenchDesign objects, see the documentation for the [BenchDesign](#) constructor.

Slots

data a list containing the data to be used in the benchmark.

methods a BDMethodList list of BDMethod objects to be compared in the benchmark.

Author(s)

Patrick Kimes

See Also

[BenchDesign](#), [BDMethod-class](#), [BDMethodList-class](#), [BDData-class](#)

buildBench	<i>Execute BenchDesign</i>
------------	----------------------------

Description

Function to evaluate methods defined in a [BenchDesign](#) on a supplied data set to generate a [SummarizedBenchmark](#) of benchmarking results. In addition to the results of applying each method on the data, the returned [SummarizedBenchmark](#) also includes metadata for the methods in the colData of the returned object, metadata for the data in the rowData, and session information in the metadata.

Usage

```
buildBench(bd, data = NULL, truthCols = NULL, ftCols = NULL,
  sortIDs = FALSE, keepData = FALSE, catchErrors = TRUE,
  parallel = FALSE, BPPARAM = bpparam())
```

Arguments

bd	BenchDesign object.
data	Data set to be used for benchmarking, will take priority over data set specified to BenchDesign object. Ignored if NULL. (default = NULL)
truthCols	Character vector of column names in data set corresponding to ground truth values for each assay. If specified, column will be added to the groundTruth DataFrame of the returned SummarizedBenchmark object. If the BenchDesign includes only a single assay, the same name will be used for the assay. If the BenchDesign includes multiple assays, to map data set columns with assays, the vector must have names corresponding to the assay names specified to the post parameter at each addMethod call. (default = NULL)
ftCols	Vector of character names of columns in data set that should be included as feature data (row data) in the returned SummarizedBenchmark object. (default = NULL)
sortIDs	Whether the output of each method should be merged and sorted using IDs. See Details for more information. (default = FALSE)
keepData	Whether to store the data as part of the BenchDesign slot of the returned SummarizedBenchmark object. If FALSE, a MD5 hash of the data will be stored with the BenchDesign slot. (default = FALSE)
catchErrors	logical whether errors produced by methods during evaluation should be caught and printed as a message without stopping the entire build process. (default = TRUE)
parallel	Whether to use parallelization for evaluating each method. Parallel execution is performed using BiocParallel . Parameters for parallelization should be specified with register or through the BPPARAM parameter. (default = FALSE)
BPPARAM	Optional BiocParallelParam instance to be used when parallel is TRUE. If not specified, the default instance from the parameter registry is used.

Details

Parallelization is performed across methods. Therefore, there is currently no benefit to specifying more cores than the total number of methods in the [BenchDesign](#) object.

By default, errors thrown by individual methods in the [BenchDesign](#) are caught during evaluation and handled in a way that allows [buildBench](#) to continue running with the other methods. The error is printed as a message, and the corresponding column in the returned [SummarizedBenchmark](#) object is set to NA. Since many benchmarking experiments can be time and computationally intensive, having to rerun the entire analysis due to a single failed method can be frustrating. Default error catching was included to alleviate these frustrations. However, if this behavior is not desired, setting `catchErrors = FALSE` will turn off error handling.

If `sortIDs = TRUE`, each method must return a named vector or list. The names will be used to align the output of each method in the returned [SummarizedBenchmark](#). Missing values from each method will be set to NA. This can be useful if the different methods return overlapping, but not identical, results. If `truthCols` is also specified, and sorting by IDs is necessary, rather than specifying `sortIDs = TRUE`, specify the string name of a column in the data to use to sort the method output to match the order of `truthCols`.

When a method specified in the [BenchDesign](#) does not have a postprocessing function specified to `post =`, the trivial `base::identity` function is used as the default postprocessing function.

Value

[SummarizedBenchmark](#) object.

Author(s)

Patrick Kimes

See Also

[updateBench](#)

Examples

```
## with toy data.frame
df <- data.frame(pval = rnorm(100))
bench <- BenchDesign(data = df)

## add methods
bench <- addMethod(bench, label = "bonf", func = p.adjust,
  params = rlang::quos(p = pval, method = "bonferroni"))
bench <- addMethod(bench, label = "BH", func = p.adjust,
  params = rlang::quos(p = pval, method = "BH"))

## evaluate benchmark experiment
sb <- buildBench(bench)

## evaluate benchmark experiment w/ data sepecified
sb <- buildBench(bench, data = df)
```

compareBDData	<i>Compare BDData objects</i>
---------------	-------------------------------

Description

Simple comparison of two BDData objects based on comparing both type and data hash.

Usage

```
compareBDData(x, y)
```

Arguments

x	a BDData or BenchDesign object
y	a BDData or BenchDesign object

Value

list of two values giving agreement of "data" and "type".

Author(s)

Patrick Kimes

See Also

[compareBenchDesigns](#)

Examples

```
## compare data with same MD5 hash value
bdd1 <- BDData(data.frame(x = 1:10))
bdd1h <- hashBDData(bdd1)
compareBDData(bdd1, bdd1h)

## compare different data, both same type
bdd2 <- BDData(data.frame(x = 2:11))
bdd2h <- hashBDData(bdd2)
compareBDData(bdd1, bdd2)
compareBDData(bdd1h, bdd2h)

## compare completely different data
compareBDData(bdd1, bdd2h)
```

compareBDMethod	<i>Compare BDMethod objects</i>
-----------------	---------------------------------

Description

Simple comparison of two BDMethod objects based on meta data.

Usage

```
compareBDMethod(x, y)
```

Arguments

x	a BDMethod object
y	a BDMethod object

Value

logical value indicating whether the two objects produced the same meta data.

Author(s)

Patrick Kimes

See Also

[compareBenchDesigns](#)

Examples

```
bdm1 <- BDMethod(stats::rnorm, params = rlang::quos(n = 100))
bdm2 <- BDMethod(stats::rt, params = rlang::quos(n = 100, df = 1))

compareBDMethod(bdm1, bdm2)
```

compareBenchDesigns	<i>Compare BenchDesign objects</i>
---------------------	------------------------------------

Description

Comparison of BenchDesign objects and BenchDesign method information stored in SummarizedBenchmark objects. Inputs can be either BenchDesign or SummarizedBenchmark objects. If SummarizedBenchmark objects are specified, the method metadata stored in the colData will be used for the comparison. If only a single SummarizedBenchmark object is specified, the colData information will be compared with the BenchDesign object in the BenchDesign slot of the object. To compare the BenchDesign slots of SummarizedBenchmark objects, the BenchDesigns should be extracted with BenchDesign(sb) and passed as inputs (see Examples).


```

                                post = sd))
bd2 <- addMethod(bd1, "chi_sd",
                func = stats::rchisq,
                params = rlang::quos(n = n, df = 1),
                post = sd)

compareBenchDesigns(bd1, bd2)

```

dropMethod

Remove method from BenchDesign object

Description

Takes a [BenchDesign](#) object and the name of an existing method and returns a reduced [BenchDesign](#) object with the method removed.

Usage

```
dropMethod(bd, label)
```

Arguments

bd	BenchDesign object.
label	Character name of method.

Value

Modified [BenchDesign](#) object with specified method dropped.

Author(s)

Patrick Kimes

See Also

[modifyMethod](#), [expandMethod](#), [addMethod](#)

Examples

```

## empty BenchDesign
bench <- BenchDesign()

## add methods
bench <- addMethod(bench, label = "bonf", func = p.adjust,
                  params = rlang::quos(p = pval, method = "bonferroni"))
bench <- addMethod(bench, label = "BH", func = p.adjust,
                  params = rlang::quos(p = pval, method = "BH"))
BDMethodList(bench)

```

```
## remove methods
bench <- dropMethod(bench, label = "bonf")
BDMethodList(bench)
```

estimateMetricsForAssay

Estimate performance metrics in SummarizedBenchmark object

Description

These functions estimate the performance metrics, either passed as arguments or added previously with the [addPerformanceMetric](#) function. The function will estimate the performance metric for each method.

Usage

```
estimateMetricsForAssay(object, assay, evalMetric = NULL,
  addColData = FALSE, evalFunction = NULL, tidy = FALSE, ...)

estimatePerformanceMetrics(object, addColData = FALSE, tidy = FALSE,
  rerun = TRUE, ...)
```

Arguments

object	A SummarizedBenchmark object.
assay	A string with an assay name. Indicates the assay that should be given as input to this performance metric.
evalMetric	A string with the name of the evaluation metric.
addColData	Logical (default: FALSE). If TRUE, the results are added to the colData slot of the SummarizedExperiment object and the object is returned. If FALSE, only a DataFrame with the results is returned.
evalFunction	A function that calculates a performance metric. It should contain at least two arguments, query and truth, where query is the output vector of a method and truth is the vector of ground true values. If additional parameters are specified, they must contain default values. If this parameter is passed, the metrics in the object are ignored and only this evaluation metric is estimated.
tidy	Logical (default: FALSE). If TRUE, a long formatted data.frame is returned.
...	Additional parameters passed to the performance functions.
rerun	Logical (default: TRUE). By default, all performance metrics are recalculated everytime that estimatePerformanceMetrics is called. If FALSE, performance metrics will only be calculated for newly added methods or modified methods.

Value

Either a [SummarizedBenchmark](#) object, a [DataFrame](#) or a [data.frame](#).

Functions

- `estimateMetricsForAssay`: Estimate performance metrics for a given assay
- `estimatePerformanceMetrics`: Estimate performance metrics for all assays

Author(s)

Alejandro Reyes

See Also

[availableMetrics](#), [performanceMetrics](#)

Examples

```
data( sb )
sb <- addPerformanceMetric(
  object=sb,
  assay="qvalue",
  evalMetric="TPR",
  evalFunction = function( query, truth, alpha=0.1 ){
    goodHits <- sum( (query < alpha) & truth == 1 )
    goodHits / sum(truth == 1)
  }
)

qvalueMetrics <- estimateMetricsForAssay( sb, assay="qvalue" )
allMetrics <- estimatePerformanceMetrics( sb )
allMetricsTidy <- estimatePerformanceMetrics( sb, tidy=TRUE )
```

expandMethod

Expand method in BenchDesign object

Description

Takes a [BenchDesign](#) object, the name of an existing method, and new parameter specifications, and returns a modified [BenchDesign](#) object with new methods added. The named method is "expanded" to multiple methods according to the specified set of parameters.

Usage

```
expandMethod(bd, label, params, onlyone = NULL, .replace = FALSE,
  .overwrite = FALSE)
```

Arguments

bd	BenchDesign object.
label	Character name of method to be expanded.
params	Named list of quosure lists specifying the label of the new methods to be added to the BenchDesign , and the set of parameters to overwrite in the original method definition for each new method. Alternatively, if <code>onlyone</code> is non-NULL, a single quosure list with <code>name = value</code> pairs specifying the label of the new methods and the values to use for overwriting the parameter specified in <code>onlyone</code> .
onlyone	Character name of a parameter to be modified. Only specify if just a single parameter should be replaced in the original method definition. Ignored if NULL. (default = NULL)
.replace	Logical whether original label method should be removed. (default = FALSE)
.overwrite	Logical whether to overwrite the existing list of parameters (TRUE) or to simply add the new parameters to the existing list (FALSE). (default = FALSE)

Value

Modified [BenchDesign](#) object with new methods with specified parameters added.

Author(s)

Patrick Kimes

See Also

[modifyMethod](#), [addMethod](#), [dropMethod](#)

Examples

```
## empty BenchDesign
bench <- BenchDesign()

## add basic 'padjust' method
bench <- addMethod(bench, label = "padjust",
  func = p.adjust,
  params = rlang::quos(p = pval, method = "none"))

## modify multiple parameters - params is a list of quosure lists
newparams <- list(bonf = rlang::quos(p = round(pval, 5), method = "bonferonni"),
  bh = rlang::quos(p = round(pval, 3), method = "BH"))
bench_exp <- expandMethod(bench, label = "padjust", params = newparams)
BDMethodList(bench_exp)

## only modify a single parameter - params is a quosure list
newparams <- rlang::quos(bonf = "bonferonni", BH = "BH")
bench_exp <- expandMethod(bench, label = "padjust", onlyone = "method", params = newparams)
BDMethodList(bench_exp)
```

groundTruths	<i>Get ground truths in SummarizedBenchmark object</i>
--------------	--

Description

Method to get groundTruths in SummarizedBenchmark object.

Usage

```
groundTruths(object, ...)

## S4 method for signature 'SummarizedBenchmark'
groundTruths(object, ...)
```

Arguments

object	a SummarizedBenchmark object.
...	further arguments, perhaps used by methods.

Value

modified BenchDesign object

Author(s)

Alejandro Reyes

groundTruths<-	<i>Set ground truths in SummarizedBenchmark object</i>
----------------	--

Description

Method to set groundTruths in SummarizedBenchmark object.

Usage

```
groundTruths(object, ...) <- value

## S4 replacement method for signature 'SummarizedBenchmark'
groundTruths(object, ...) <- value
```

Arguments

object	a SummarizedBenchmark object.
...	further arguments, perhaps used by methods.
value	replacement set of ground truths.

Value

modified BenchDesign object

Author(s)

Alejandro Reyes, Patrick Kimes

hashBDData	<i>Hash data in BDData object</i>
------------	-----------------------------------

Description

Replaces data stored in a [BDData](#) object with the MD5 hash of the data. If the data was already a MD5 hash, the original object is returned unchanged. The method can be called directly on [BenchDesign](#) objects to hash the underlying data as well.

Usage

```
hashBDData(object)

## S4 method for signature 'BDData'
hashBDData(object)

## S4 method for signature 'BenchDesign'
hashBDData(object)
```

Arguments

object a BDData or BenchDesign object

Value

an object of the same class as object with data converted to a MD5 hash.

Author(s)

Patrick Kimes

```
mcols<- , SummarizedBenchmark-method
```

Set meta data columns in SummarizedBenchmark object

Description

Modifies the mcols slot of a [SummarizedBenchmark](#) object.

Usage

```
## S4 replacement method for signature 'SummarizedBenchmark'
mcols(x, ...) <- value
```

Arguments

x	A SummarizedBenchmark object.
...	Futher arguments, perhaps used by methods.
value	A DataFrame of meta data.

Value

Modified [SummarizedBenchmark](#) object.

Author(s)

Alejandro Reyes

See Also

[SummarizedBenchmark](#)

modifyMethod	<i>Modify method in BenchDesign object</i>
--------------	--

Description

Takes a [BenchDesign](#) object, the name of an existing method, and new parameter specifications, and returns a modified [BenchDesign](#) object with the specified changes.

Usage

```
modifyMethod(bd, label, params, .overwrite = FALSE)
```

Arguments

<code>bd</code>	BenchDesign object.
<code>label</code>	Character name of method to be modified.
<code>params</code>	Named quosure list created using quos of parameter = value pairs to replace in the method definition. The <code>post</code> , and <code>meta</code> parameters of the method can be modified using the special keywords, <code>bd.post</code> , and <code>bd.meta</code> (the prefix denoting that these values should modify BenchDesign parameters). All other named parameters will be added to the list of parameters to be passed to <code>func</code> .
<code>.overwrite</code>	Logical whether to overwrite the complete existing list of parameters to be passed to <code>func</code> (TRUE), or to simply add the new parameters to the existing list and only replace overlapping parameters (FALSE). (default = FALSE)

Value

Modified [BenchDesign](#) object with single method parameters modified.

Author(s)

Patrick Kimes

See Also

[addMethod](#), [expandMethod](#), [dropMethod](#)

Examples

```
## empty BenchDesign
bench <- BenchDesign()

## add method
bench <- addMethod(bench, label = "qv",
  func = qvalue::qvalue,
  post = function(x) { x$qvalue },
  meta = list(note = "storey's q-value"),
  params = rlang::quos(p = pval))

## modify method 'meta' property of 'qv' method
bench <- modifyMethod(bench, label = "qv",
  params = rlang::quos(bd.meta =
    list(note = "Storey's q-value"))))

## verify that method has been updated
printMethod(bench, "qv")
```

performanceMetrics	<i>Get performance metrics in SummarizedBenchmark object</i>
--------------------	--

Description

Given a [SummarizedBenchmark](#) object, returns a list of lists of performance metrics that have been defined for each assay. Optionally, if assay = is specified, performance metrics for only the specified subset of specified assays are returned.

Usage

```
performanceMetrics(object, ...)  
  
## S4 method for signature 'SummarizedBenchmark'  
performanceMetrics(object, assay = NULL)
```

Arguments

object	a SummarizedBenchmark object.
...	further arguments, perhaps used by methods.
assay	a character string indicating an assay name.

Value

A SimpleList with one element for each assay. Each element of the list contains a list of performance metric functions.

Author(s)

Alejandro Reyes

See Also

[addPerformanceMetric](#), [estimatePerformanceMetrics](#)

Examples

```
data(sb)  
performanceMetrics(sb)  
performanceMetrics(sb, assay = "qvalue")
```

```
performanceMetrics<- Set performance metrics in SummarizedBenchmark object
```

Description

Replaces the list of performance metrics in a [SummarizedBenchmark](#) object with a new list of performance metric lists.

Usage

```
performanceMetrics(object, ...) <- value
```

```
## S4 replacement method for signature 'SummarizedBenchmark, SimpleList'  
performanceMetrics(object) <- value
```

Arguments

object	a SummarizedBenchmark object.
...	further arguments, perhaps used by methods.
value	a SimpleList of the same length as the number of assays.

Value

Silently, the newly specified SimpleList of performance metric lists.

Author(s)

Alejandro Reyes

See Also

[addPerformanceMetric](#), [estimatePerformanceMetrics](#), [performanceMetrics](#)

Examples

```
data(sb)  
performanceMetrics(sb)  
performanceMetrics(sb) <- SimpleList(qvalue = list(), logFC = list())
```

plotMethodsOverlap *Plot UpSetR for SummarizedBenchmark object*

Description

This function looks for an assay, called by default 'qvalue', and given an alpha threshold, it binarizes the assay matrix depending on whether its values are below the alpha threshold. Then it uses the function [upset](#) to plot the overlaps. The plot is only generated if at least 2 methods have observations that pass the alpha threshold.

Usage

```
plotMethodsOverlap(object, assay = "qvalue", alpha = 0.1, ...)
```

Arguments

object	A SummarizedBenchmark object.
assay	The name of an assay.
alpha	An alpha value.
...	Further arguments passed to upset

Value

An upseR plot.

Author(s)

Alejandro Reyes

See Also

[plotROC](#), [estimatePerformanceMetrics](#)

Examples

```
data(sb)
## Not run:
plotMethodsOverlap(sb)

## End(Not run)
```

plotROC	<i>Plot ROC curve for SummarizedBenchmark object</i>
---------	--

Description

This function inputs a [SummarizedBenchmark](#) object, looks for an assay called 'qvalue' and plots receiver operating characteristic curves for each of the methods to benchmark.

Usage

```
plotROC(object, assay = "qvalue")
```

Arguments

object	A SummarizedBenchmark object.
assay	An assay name.

Value

A ggplot object.

Author(s)

Alejandro Reyes

See Also

[plotMethodsOverlap](#), [estimatePerformanceMetrics](#)

Examples

```
data(sb)
## Not run:
plotROC(sb)

## End(Not run)
```

printMethod	<i>Pretty print methods in a BenchDesign object</i>
-------------	---

Description

Print out details about a method included in the BenchDesign. The printMethods function is just a wrapper to call printMethod on all methods in the BenchDesign.

Usage

```
printMethod(bd, n = NULL)

printMethods(bd)
```

Arguments

bd	BenchDesign object.
n	name of a method in the BenchDesign to show.

Value

Brief description is returned to console.

Author(s)

Patrick Kimes

See Also

[BDMethod-class](#), [BenchDesign-class](#)

Examples

```
## create empty BenchDesign
bench <- BenchDesign()

## currently no methods
printMethods(bench)

## add method
bench <- addMethod(bench, label = "method_a", p.adjust)
bench <- addMethod(bench, label = "method_b", qvalue::qvalue)

## show a single method
printMethod(bench, "method_a")

## show all methods
printMethods(bench)
```

sb	<i>SummarizedBenchmark example</i>
----	------------------------------------

Description

This object contains the example data from the iCOBRA package reformatted as a SummarizedBenchmark object. It consists of differential expression results from DESeq2 edgeR and limma-voom.

Source

Example data from the iCOBRA package.

Examples

```
data(sb)
```

show,BDData-method	<i>Show BDData object</i>
--------------------	---------------------------

Description

Show BDData object

Usage

```
## S4 method for signature 'BDData'  
show(object)
```

Arguments

object	BDData object to show
--------	-----------------------

Value

Print description of BDData object to console

show,BDMethod-method *Show BDMethod object*

Description

Show BDMethod object

Usage

```
## S4 method for signature 'BDMethod'
show(object)
```

Arguments

object BDMethod object to show

Value

Print description of BDMethod object to console

show,BDMethodList-method
 Show BDMethodList object

Description

Show BDMethodList object

Usage

```
## S4 method for signature 'BDMethodList'
show(object)
```

Arguments

object BDMethodList object to show

Value

Print description of BDMethodList object to console

show,BenchDesign-method	<i>Show BenchDesign object</i>
-------------------------	--------------------------------

Description

Show BenchDesign object

Usage

```
## S4 method for signature 'BenchDesign'
show(object)
```

Arguments

object BenchDesign object to show

Value

Print description of BenchDesign object to console

SummarizedBenchmark	<i>Create a new SummarizedBenchmark object</i>
---------------------	--

Description

Function to construct SummarizedBenchmark objects.

Usage

```
SummarizedBenchmark(assays, colData, ftData = NULL, groundTruth = NULL,
  performanceMetrics = NULL, BenchDesign = NULL, ...)
```

Arguments

- | | |
|---------|--|
| assays | A list containing outputs of the methods to be benchmark. Each element of the list must contain a matrix or data.frame of n x m, n being the number of features tested (e.g. genes) and m being the number of methods in the benchmark. Each element of the list must contain a single assay (the outputs of the methods). For example, for a benchmark of differential expression methods, one assay could contain the q-values from the different methods and another assay could be the estimated log fold changes. |
| colData | A DataFrame describing the annotation of the methods. These could include version of the software or the parameters used to run them. |

ftData	A DataFrame object describing the rows. This parameter is equivalent to the parameter rowData of a SummarizedExperiment.
groundTruth	If present, a DataFrame containing the ground truths. If provided, the number of columns must be the same as the number of assays (NA's are accepted). The names of the columns should have the same names as the assays.
performanceMetrics	A SimpleList of the same length as the number of assays. Each element of the list must be a list of functions. Each function must contain the parameters 'query' and 'truth'.
BenchDesign	A BenchDesign containing the code used to construct the object. (default = NULL)
...	Additional parameters passed to SummarizedExperiment .

Value

A [SummarizedBenchmark](#) object.

Author(s)

Alejandro Reyes

Examples

```
## loading the example data from iCOBRA
library(iCOBRA)
data(cobradata_example)

## a bit of data wrangling and reformatting
assays <- list(
  qvalue=cobradata_example@padj,
  logFC=cobradata_example@score )
assays[["qvalue"]]$DESeq2 <- p.adjust(cobradata_example@pval$DESeq2, method="BH")
groundTruth <- DataFrame( cobradata_example@truth[,c("status", "logFC")] )
colnames(groundTruth) <- names( assays )
colData <- DataFrame( method=colnames(assays[[1]]) )
groundTruth <- groundTruth[rownames(assays[[1]]),]

## constructing a SummarizedBenchmark object
sb <- SummarizedBenchmark(
  assays=assays, colData=colData,
  groundTruth=groundTruth )
colData(sb)$label <- rownames(colData(sb))
```

SummarizedBenchmark-class

SummarizedBenchmark class

Description

Extension of the [RangedSummarizedExperiment](#) to store the output of different methods intended for the same purpose in a given dataset. For example, a differential expression analysis could be done using **limma**-voom, **edgeR** and **DESeq2**. The SummarizedBenchmark class provides a framework that is useful to store, benchmark and compare results.

Slots

performanceMetrics A [SimpleList](#) of the same length as the number of [assays](#) containing performance functions to be compared with the ground truths.

BenchDesign A [BenchDesign](#) originally used to generate the results in the object.

Author(s)

Alejandro Reyes

Examples

```
## loading the example data from iCOBRA
library(iCOBRA)
data(cobradata_example)

## a bit of data wrangling and reformatting
assays <- list(
  qvalue=cobradata_example@padj,
  logFC=cobradata_example@score )
assays[["qvalue"]]$DESeq2 <- p.adjust(cobradata_example@pval$DESeq2, method="BH")
groundTruth <- DataFrame( cobradata_example@truth[,c("status", "logFC")] )
colnames(groundTruth) <- names( assays )
colData <- DataFrame( method=colnames(assays[[1]]) )
groundTruth <- groundTruth[rownames(assays[[1]]),]

## constructing a SummarizedBenchmark object
sb <- SummarizedBenchmark(
  assays=assays, colData=colData,
  groundTruth=groundTruth )
```

tdat

Example data.frame containing results for 50 two-sample t-tests.

Description

Example data.frame containing results for 50 two-sample t-tests.

Format

a data.frame that contains the results of 50 simulated two-sample t-tests, with each row corresponding to an independent test. The data.frame includes the following 5 columns: 1. H = binary 0/1 whether data for the test was simulated under the null (0) or alternative (1) 2. test_statistic = test-statistics of the t-test 3. effect_size = mean difference between the two sample groups 4. pval = p-value of the t-test 5. SE = standard error of the t-test

Examples

```
data(tdat)
```

tidyBDMMethod

Tidy BDMMethod Data

Description

A helper function to extract information for a single or multiple BDMMethod object or a list of BDMMethod objects.

Usage

```
tidyBDMMethod(obj, dat = NULL, eval = FALSE, label = FALSE)
```

```
## S4 method for signature 'BDMMethod'
tidyBDMMethod(obj, dat, eval)
```

```
## S4 method for signature 'list'
tidyBDMMethod(obj, dat = NULL, eval = FALSE,
  label = FALSE)
```

```
## S4 method for signature 'SimpleList'
tidyBDMMethod(obj, dat = NULL, eval = FALSE,
  label = FALSE)
```

```
## S4 method for signature 'BenchDesign'
tidyBDMMethod(obj, dat = NULL, eval = FALSE,
  label = FALSE)
```

Arguments

obj	BDMethod object, list/List of BDMethod objects (e.g. a BDMethodList), or a BenchDesign object
dat	optional data object to use when evaluating any unevaluated expressions in bdmmeta data. (default = NULL)
eval	logical whether to evaluate any quosures in the meta slot of the BDMethod objects. (default = FALSE)
label	logical whether to add a "label" column to the resulting table containing the names of the methods if obj was specified as a named list. (default = FALSE)

Details

If any quosures are specified to the "meta" slot of a BDMethod object, the quosure is converted to a text string using `rlang::quo_text`.

Value

A named vector of meta data if only a single BDMethod object specified, else a tibble of meta data for the specified list of methods.

Author(s)

Patrick Kimes

tidyUpMetrics

Tidy up performance metrics in SummarizedBenchmark object

Description

This function takes as input a SummarizedBenchmark object, extracts the estimated performance metrics and reformats them into a long-formatted data frame.

Usage

```
tidyUpMetrics(object)
```

Arguments

object A [SummarizedBenchmark](#) object.

Value

A tidy data.frame

Author(s)

Alejandro Reyes

See Also[estimatePerformanceMetrics](#)**Examples**

```
data( "sb", package="SummarizedBenchmark" )
sb <- estimateMetricsForAssay( sb, assay="qvalue", evalMetric="rejections",
  evalFunction=function( query, truth, alpha=0.1 ){
    sum( query < alpha )
  },
  addColData=TRUE )
tidyUpMetrics( sb )
```

updateBench	<i>Check/Update SummarizedBenchmark</i>
-------------	---

Description

Function to update or check status of [SummarizedBenchmark](#) results.

If only a [SummarizedBenchmark](#) object is specified, the function will check whether ‘func’, ‘param’, ‘meta’, ‘post’ or the ‘pkg_vers’ of the methods in the [BenchDesign](#) stored with the [SummarizedBenchmark](#) do not match values stored in the colData. By default, no methods will be executed to update results. To actually execute updates, set dryrun = FALSE.

If a [BenchDesign](#) object is specified in addition to a [SummarizedBenchmark](#) object, the function will check which methods in the new [BenchDesign](#) need to be executed to update the [SummarizedBenchmark](#) results. Again, by default, no methods will be executed unless dryrun = FALSE is specified.

Unless reuseParams = FALSE is specified, the parameters of the last execution session stored in the the colData of the [SummarizedBenchmark](#) object will be used.

Usage

```
updateBench(sb, bd = NULL, dryrun = TRUE, version = FALSE,
  keepAll = TRUE, reuseParams = TRUE, ...)
```

Arguments

sb	a SummarizedBenchmark object
bd	a BenchDesign object
dryrun	logical whether to just print description of what would be updated rather than actually running any methods. (default = TRUE)
version	logical whether to re-run methods with only package version differences. (default = FALSE)
keepAll	logical whether to keep methods run in original SummarizedBenchmark but not in new BenchDesign . Only used if bd is not NULL. (default = TRUE)

reuseParams logical whether to reuse parameters from buildBench call used to create [SummarizedBenchmark](#) object (if available). Directly specified [buildBench](#) parameters still take precedence. (default = TRUE)

... optional parameters to pass to [buildBench](#).

Value

SumamrizedBenchmark object.

Author(s)

Patrick Kimes

See Also

[buildBench](#)

Examples

```
## load example SummarizedBenchmark object
data(allSB)
sb <- allSB[[1]]

## check if results are out of date
updateBench(sb)

## modify BenchDesign
bd <- BenchDesign(sb)
bd <- dropMethod(bd, "kallisto-default")

## check if results need to be updated with new BenchDesign
updateBench(sb, bd)
```

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