

# Package ‘MultiDataSet’

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**Type** Package

**Title** Implementation of MultiDataSet and ResultSet

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**Description** Implementation of the BRGE's (Bioinformatic Research Group in Epidemiology from Center for Research in Environmental Epidemiology) MultiDataSet and ResultSet. MultiDataSet is designed for integrating multi omics data sets and ResultSet is a container for omics results. This package contains base classes for MEAL and rexposome packages.

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**LazyData** TRUE

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limma

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---

|          |   |
|----------|---|
| add_eset | <i>Method to add an eSet to MultiDataSet.</i> |
|----------|---|

---

### Description

This method adds or overwrites a slot of a `MultiDataSet` with the content of the given `eSet`.

### Usage

```
add_eset(
  object,
  set,
  dataset.type,
  dataset.name = NULL,
  sample.tables = NULL,
  feature.tables = NULL,
  warnings = TRUE,
  overwrite = FALSE,
```

```

    GRanges
  )

```

### Arguments

|                |  |
|----------------|--|
| object         | MultiDataSet that will be filled.  |
| set            | Object derived from eSet to be used to fill the slot.  |
| dataset.type   | Character with the type of data of the omic set (e.g. expression, methylation...)  |
| dataset.name   | Character with the specific name for this set (NULL by default). It is useful when there are several sets of the same type (e.g. multiple expression assays) |
| sample.tables  | Character with the names of the slots with sample data besides phenoData.  |
| feature.tables | Character with the names of the slots with feature data besides featureData.   |
| warnings       | Logical to indicate if warnings will be displayed.   |
| overwrite      | Logical to indicate if the set stored in the slot will be overwritten.   |
| GRanges        | GenomicRanges to be included in rowRanges slot.  |

### Value

A new MultiDataSet with a slot filled.

### See Also

[add\\_methy](#), [add\\_genexp](#), [add\\_rnaseq](#), [add\\_snps](#)

### Examples

```

multi <- createMultiDataSet()
eset <- new("ExpressionSet", exprs = matrix(runif(10), 5))
multi <- add_eset(multi, eset, "exampledata", GRanges = NA)

```

---

add\_genexp

*Method to add an expression microarray dataset to MultiDataSet.*

---

### Description

This method adds or overwrites the slot "expression" of an MultiDataSet with the content of the given ExpressionSet. The fData of the ExpressionSet must contain the columns chromosome, start and end.

### Usage

```
add_genexp(object, gexpSet, ...)
```

**Arguments**

object            MultiDataSet that will be filled.  
 gexpSet          ExpressionSet to be used to fill the slot.  
 ...              Arguments to be passed to add\_eset.

**Value**

A new MultiDataSet with the slot "expression" filled.

**Examples**

```
multi <- createMultiDataSet()
eset <- new("ExpressionSet", exprs = matrix(runif(4), 2))
fData(eset) <- data.frame(chromosome = c("chr1", "chr2"), start = c(12414, 1234321),
  end = c(121241, 124124114), stringsAsFactors = FALSE)
multi <- add_genexp(multi, eset)
```

---

add\_methy                      *Method to add a slot of methylation to MultiDataSet.*

---

**Description**

This method adds or overwrites the slot "methylation" of an MultiDataSet with the content of the given MethylationSet or RatioSet. The fData of the input object must contain the columns chromosome and position.

**Usage**

```
add_methy(object, methySet, ...)
```

**Arguments**

object            MultiDataSet that will be filled.  
 methySet          MethylationSet or RatioSet to be used to fill the slot.  
 ...              Further arguments to be passed to add\_eset.

**Value**

A new MultiDataSet with the slot "methylation" filled.

**Examples**

```
if (require(brgedata)){
  multi <- createMultiDataSet()
  multi <- add_methy(multi, brge_methy[1:100, ])
}
```

---

|            |   |
|------------|---|
| add_rnaseq | <i>Method to add an expression RNA seq dataset to MultiDataSet.</i> |
|------------|---|

---

### Description

This method adds or overwrites the slot "rnaseq" of an MultiDataSet with the content of the given ExpressionSet. The fData of the ExpressionSet must contain the columns chromosome, start and end.

### Usage

```
add_rnaseq(object, rnaSet, ...)
```

### Arguments

|        |  |
|--------|--|
| object | MultiDataSet that will be filled.          |
| rnaSet | ExpressionSet to be used to fill the slot. |
| ...    | Arguments to be passed to add_eset.        |

### Value

A new MultiDataSet with the slot "rnaseq" filled.

### Examples

```
multi <- createMultiDataSet()  
eset <- new("ExpressionSet", exprs = matrix(runif(4), 2))  
fData(eset) <- data.frame(chromosome = c("chr1", "chr2"), start = c(12414, 1234321),  
  end = c(121241, 12122414), stringsAsFactors = FALSE)  
multi <- add_genexp(multi, eset)
```

---

|         |  |
|---------|--|
| add_rse | <i>Method to add a RangedSummarizedExperiment to MultiDataSet.</i> |
|---------|--|

---

### Description

This method adds or overwrites a slot of a MultiDataSet with the content of the given RangedSummarizedExperiment.

**Usage**

```
add_rse(
  object,
  set,
  dataset.type,
  dataset.name = NULL,
  sample.tables = NULL,
  feature.tables = NULL,
  warnings = TRUE,
  overwrite = FALSE
)
```

**Arguments**

|                             |  |
|-----------------------------|--|
| <code>object</code>         | MultiDataSet that will be filled.  |
| <code>set</code>            | Object derived from RangedSummarizedExperiment to be used to fill the slot.  |
| <code>dataset.type</code>   | Character with the type of data of the omic set (e.g. expression, methylation...)  |
| <code>dataset.name</code>   | Character with the specific name for this set (NULL by default). It is useful when there are several sets of the same type (e.g. multiple expression assays) |
| <code>sample.tables</code>  | Character with the names of the slots with sample data besides colData.  |
| <code>feature.tables</code> | Character with the names of the slots with feature data besides rowData.   |
| <code>warnings</code>       | Logical to indicate if warnings will be displayed.   |
| <code>overwrite</code>      | Logical to indicate if the set stored in the slot will be overwritten.   |

**Value**

A new MultiDataSet with a slot filled.

**Examples**

```
if (require(GenomicRanges) & require(SummarizedExperiment)){
  multi <- createMultiDataSet()
  counts <- matrix(runif(200 * 6, 1, 1e4), 200)
  rowRanges <- GRanges(rep(c("chr1", "chr2"), c(50, 150)),
    IRanges(floor(runif(200, 1e5, 1e6)), width=100),
    strand=sample(c("+", "-"), 200, TRUE),
    feature_id=sprintf("ID%03d", 1:200))
  colData <- DataFrame(Treatment=rep(c("ChIP", "Input"), 3),
    row.names=LETTERS[1:6], id = LETTERS[1:6])
  names(rowRanges) <- 1:200
  rse <- SummarizedExperiment(assays=SimpleList(counts=counts),
    rowRanges=rowRanges, colData=colData)
  multi <- add_rse(multi, rse, "rseEx")
}
```

---

|        |  |
|--------|--|
| add_se | <i>Method to add a SummarizedExperiment to MultiDataSet.</i> |
|--------|--|

---

## Description

This method adds or overwrites a slot of a `MultiDataSet` with the content of the given `SummarizedExperiment`.

## Usage

```
add_se(  
  object,  
  set,  
  dataset.type,  
  dataset.name = NULL,  
  sample.tables = NULL,  
  feature.tables = NULL,  
  warnings = TRUE,  
  overwrite = FALSE,  
  GRanges  
)
```

## Arguments

|                             |  |
|-----------------------------|--|
| <code>object</code>         | <code>MultiDataSet</code> that will be filled.   |
| <code>set</code>            | Object derived from <code>SummarizedExperiment</code> to be used to fill the slot.   |
| <code>dataset.type</code>   | Character with the type of data of the omic set (e.g. expression, methylation...)  |
| <code>dataset.name</code>   | Character with the specific name for this set (NULL by default). It is useful when there are several sets of the same type (e.g. multiple expression assays) |
| <code>sample.tables</code>  | Character with the names of the slots with sample data besides <code>colData</code> .  |
| <code>feature.tables</code> | Character with the names of the slots with feature data besides <code>rowData</code> .   |
| <code>warnings</code>       | Logical to indicate if warnings will be displayed.   |
| <code>overwrite</code>      | Logical to indicate if the set stored in the slot will be overwritten.   |
| <code>GRanges</code>        | <code>GenomicRanges</code> to be included in <code>rowRanges</code> slot.  |

## Value

A new `MultiDataSet` with a slot filled.

## Examples

```
multi <- createMultiDataSet()  
se <- SummarizedExperiment::SummarizedExperiment(matrix(runif(10), 5))  
multi <- add_se(multi, se, "exampledata", GRanges = NA)
```

---

|          |  |
|----------|--|
| add_snps | <i>Method to add a slot of SNPs to MultiDataSet.</i> |
|----------|--|

---

### Description

This method adds or overwrites the slot "snps" of an MultiDataSet with the content of the given SnpSet. The fData of the SnpSet must contain the columns chromosome and position.

### Usage

```
add_snps(object, snpSet, ...)
```

### Arguments

|        |                                     |
|--------|-------------------------------------|
| object | MultiDataSet that will be filled.   |
| snpSet | SnpSet to be used to fill the slot. |
| ...    | Arguments to be passed to add_eset. |

### Value

A new MultiDataSet with the slot "snps" filled.

### Examples

```
multi <- createMultiDataSet()
geno <- matrix(c(3,1,2,1), ncol = 2)
colnames(geno) <- c("VAL0156", "VAL0372")
rownames(geno) <- c("rs3115860", "SNP1-1628854")
map <- AnnotatedDataFrame(data.frame(chromosome = c("chr1", "chr2"), position = c(12414, 1234321),
  stringsAsFactors = FALSE))
rownames(map) <- rownames(geno)
snpSet <- new("SnpSet", call = geno, featureData = map)
pheno <- data.frame(id = c("VAL0156", "VAL0372"))
rownames(pheno) <- c("VAL0156", "VAL0372")
pData(snpSet) <- pheno
multi <- add_snps(multi, snpSet)
```

---

|           |  |
|-----------|--|
| add_table | <i>Method to add a matrix to MultiDataSet.</i> |
|-----------|--|

---

### Description

This method adds or overwrites a slot of a MultiDataSet with the content of the given matrix.



**Usage**

```
add_table(  
  object,  
  set,  
  dataset.type,  
  dataset.name = NULL,  
  warnings = TRUE,  
  overwrite = FALSE  
)
```

**Arguments**

|              |   |
|--------------|---|
| object       | MultiDataSet that will be filled.   |
| set          | matrix used to fill the slot.   |
| dataset.type | Character with the type of data   |
| dataset.name | Character with the specific name for this set (NULL by default). It is useful when there are several sets of the same type. |
| warnings     | Logical to indicate if warnings will be displayed.  |
| overwrite    | Logical to indicate if the set stored in the slot will be overwritten.  |

**Value**

A new MultiDataSet with a slot filled.

**Examples**

```
multi <- createMultiDataSet()  
mat <- matrix(runif(12), nrow = 3)  
colnames(mat) <- paste0("S", 1:4)  
rownames(mat) <- paste0("F", 1:3)  
multi <- add_table(multi, mat, "exampledata")
```

---

chrNumToChar

*Convert chr numbers to chr strings*

---

**Description**

Given a vector of number representing the chromosomes, convert them to string (e.g 1 to chr1). 23 is consider chrX, 24 is chrY, 25 is chrXY (probes shared between chromosomes X and Y) and 26 is chrMT.

**Usage**

```
chrNumToChar(vector)
```

**Arguments**

vector            The vector with the chromosome numbers

**Value**

A vector with the chromosomes in string format.

**Examples**

```
chromosomes <- c(1, 3, 4, 23, 15)
stringChrs <- chrNumToChar(chromosomes)
stringChrs
```

---

|           |   |
|-----------|---|
| commonIds | <i>Get the name of the ids common to all datasets</i> |
|-----------|---|

---

**Description**

Get the name of the ids common to all datasets

**Usage**

```
commonIds(object)
```

**Arguments**

object            MultiDataSet that will be filtered.

**Value**

Character vector with the common ids.

**Examples**

```
multi <- createMultiDataSet()
eset <- new("ExpressionSet", exprs = matrix(runif(9), ncol = 3))
fData(eset) <- data.frame(chromosome = c("chr1", "chr1", "chr1"),
                          start = c(1, 5, 10), end = c(4, 6, 14),
                          stringsAsFactors = FALSE)
sampleNames(eset) <- c("S1", "S2", "S3")
pData(eset) <- data.frame(id = c("S1", "S2", "S3"))
rownames(pData(eset)) <- c("S1", "S2", "S3")
multi <- add_genexp(multi, eset, dataset.name = "g1")
eset <- new("ExpressionSet", exprs = matrix(runif(8), ncol = 2))
fData(eset) <- data.frame(chromosome = c("chr1", "chr1", "chr1", "chr1"),
                          start = c(1, 14, 25, 104), end = c(11, 16, 28, 115),
                          stringsAsFactors = FALSE)
sampleNames(eset) <- c("S1", "G2")
pData(eset) <- data.frame(id = c("S1", "G2"))
```

```
rownames(pData(eset)) <- c("S1", "G2")

multi <- add_genexp(multi, eset, dataset.name="g2")
commonIds(multi)
```

---

|               |   |
|---------------|---|
| commonSamples | <i>Method to select samples that are present in all datasets.</i> |
|---------------|---|

---

## Description

This method subsets the datasets to only contain the samples that are in all datasets. All sets will have the samples in the same order, taking into account that there can be duplicates.

## Usage

```
commonSamples(object, unify.names = FALSE)
```

## Arguments

|             |   |
|-------------|---|
| object      | MultiDataSet that will be filtered.                               |
| unify.names | Logical indicating if sample names of the sets should be unified. |

## Details

If unify.names is TRUE, the sample names of the sets will be unified using the id column of phenodata. This option is only possible when there are no duplicated ids.

## Value

A new MultiDataSet with only the common samples.

## Examples

```
multi <- createMultiDataSet()
eset <- new("ExpressionSet", exprs = matrix(runif(9), ncol = 3))
fData(eset) <- data.frame(chromosome = c("chr1", "chr1", "chr1"),
  start = c(1, 5, 10), end = c(4, 6, 14),
  stringsAsFactors = FALSE)
sampleNames(eset) <- c("S1", "S2", "S3")
pData(eset) <- data.frame(id = c("S1", "S2", "S3"))
rownames(pData(eset)) <- c("S1", "S2", "S3")
multi <- add_genexp(multi, eset, dataset.name = "g1")
eset <- new("ExpressionSet", exprs = matrix(runif(8), ncol = 2))
fData(eset) <- data.frame(chromosome = c("chr1", "chr1", "chr1", "chr1"),
  start = c(1, 14, 25, 104), end = c(11, 16, 28, 115),
  stringsAsFactors = FALSE)
sampleNames(eset) <- c("S1", "G2")
pData(eset) <- data.frame(id = c("S1", "G2"))
rownames(pData(eset)) <- c("S1", "G2")
```

```
multi <- add_genexp(multi, eset, dataset.name="g2")
commonSamples(multi)
```

---

getAssociation                    *Method to extrat feature result from a ResultSet*

---

### Description

Homologous methods from limma, getAssociation returns a data.frame with the logFC and PValue per featrue for the selcted coef and for given result (rid).

### Usage

```
getAssociation(object, rid = 1, coef = 2, contrast = NULL, fNamees = NULL, ...)
```

### Arguments

|          |   |
|----------|---|
| object   | A <a href="#">ResultSet</a> object.   |
| rid      | The name or index of the result to be extracted.  |
| coef     | (default 2) Index of the coefficient to be extracted.   |
| contrast | (default 1) When code corresponds to a multicategorical variable, contastr selects the comparison.  |
| fNames   | (default c("chromosome", "start", "end", "genesymbol")) Corresponds to the columns selected from fData that will be incorporated into the resulting data.frame. |
| ...      | Further arguments passed to <a href="#">topTable</a>  |

### Value

A data.frame with the result of the association study, including P-Value and Fold Change.

### Examples

```
data(rset)
getAssociation(rset, rid=1, fNamees = c("chromosome", "position"))
```

---

|               |  |
|---------------|--|
| lambdaClayton | <i>Lambda Calculation for a vector of P-Values</i> |
|---------------|--|

---

**Description**

Implementation of Clayton's lambda score for a vector of P-Values

**Usage**

```
lambdaClayton(x, trim = 0.5)
```

**Arguments**

|      |                   |
|------|-------------------|
| x    | Vector of P-Value |
| trim | (default 0.5)     |

**Value**

A lambda value, indicating the inflation/deflation of the analysis.

**Author(s)**

Juan R. Gonzalez

**Examples**

```
lambdaClayton(runif(30))
```

---

|         |   |
|---------|---|
| mae2mds | <i>Convert a MultiAssayExperiment to a MultiDataSet</i> |
|---------|---|

---

**Description**

This function creates a MultiDataSet using the data of a MultiAssayExperiment.

**Usage**

```
mae2mds(MAE, warnings = TRUE)
```

**Arguments**

|          |  |
|----------|--|
| MAE      | a MultiAssayExperiment                             |
| warnings | Logical to indicate if warnings will be displayed. |

**Value**

MultiDataSet with the of the incoming MultiAssayExperiment.

---

|         |   |
|---------|---|
| mds2mae | <i>Convert a MultiDataSet to a MultiAssayExperiment</i> |
|---------|---|

---

### Description

This function creates a MultiAssayExperiment using the data of a MultiDataSet.

### Usage

```
mds2mae(MDS)
```

### Arguments

MDS            a MultiDataSet

### Value

MultiAssayExperiment with the of the incoming MultiDataSet.

---

|              |   |
|--------------|---|
| MultiDataSet | <i>MultiDataSet: Implementation of the BRGE's basic classes</i> |
|--------------|---|

---

### Description

Implementation of the BRGE's (Bioinformatic Research Group in Epidemiology from Center for Research in Environmental Epidemiology) MultiDataSet and MethylationSet. MultiDataSet is designed for integrating multi omics data sets and MethylationSet to contain normalized methylation data. MultiDataSet for integrating multi omics data sets

### See Also

[MultiDataSet](#)

---

MultiDataSet-class      *MultiDataSet instances*

---

### Description

The class MultiDataSet is a superior class to store multiple datasets in form of triplets (assayData-phenodata-featureData). The datasets can be eSet or SummarizedExperiment derived or matrices.

### Usage

```
## S4 method for signature 'MultiDataSet,eSet'
add_eset(
  object,
  set,
  dataset.type,
  dataset.name = NULL,
  sample.tables = "protocolData",
  feature.tables = NULL,
  warnings = TRUE,
  overwrite = FALSE,
  GRanges
)

## S4 method for signature 'MultiDataSet,ExpressionSet'
add_genexp(object, gexpSet, ...)

## S4 method for signature 'MultiDataSet,ExpressionSet'
add_rnaseq(object, rnaSet, ...)

## S4 method for signature 'MultiDataSet,GenomicRatioSet'
add_methy(object, methySet, ...)

## S4 method for signature 'MultiDataSet,RangedSummarizedExperiment'
add_rse(
  object,
  set,
  dataset.type,
  dataset.name = NULL,
  sample.tables = NULL,
  feature.tables = "elementMetadata",
  warnings = TRUE,
  overwrite = FALSE
)

## S4 method for signature 'MultiDataSet,SummarizedExperiment'
add_se(
  object,
```

```
    set,
    dataset.type,
    dataset.name = NULL,
    sample.tables = NULL,
    feature.tables = "elementMetadata",
    warnings = TRUE,
    overwrite = FALSE,
    GRanges
  )

## S4 method for signature 'MultiDataSet,SnpSet'
add_snps(object, snpSet, ...)

## S4 method for signature 'MultiDataSet,matrix'
add_table(
  object,
  set,
  dataset.type,
  dataset.name = NULL,
  warnings = TRUE,
  overwrite = FALSE
)

## S4 method for signature 'MultiDataSet'
as.list(x)

## S4 method for signature 'MultiDataSet'
commonIds(object)

## S4 method for signature 'MultiDataSet'
commonSamples(object, unify.names = FALSE)

createMultiDataSet()

## S4 method for signature 'MultiDataSet'
dims(x)

## S4 method for signature 'MultiDataSet'
w_iclusterplus(object, commonSamples = TRUE, ...)

## S4 method for signature 'MultiDataSet'
length(x)

## S4 method for signature 'MultiDataSet'
w_mcia(object, ...)

## S4 method for signature 'MultiDataSet'
names(x)
```



```

## S4 method for signature 'MultiDataSet'
ncol(x)

## S4 method for signature 'MultiDataSet'
nrow(x)

## S4 method for signature 'MultiDataSet'
rowRangesElements(object)

## S4 method for signature 'MultiDataSet'
sampleNames(object)

## S4 method for signature 'MultiDataSet'
assayData(object)

## S4 method for signature 'MultiDataSet'
fData(object)

## S4 method for signature 'MultiDataSet'
featureData(object)

## S4 method for signature 'MultiDataSet'
pData(object)

## S4 method for signature 'MultiDataSet'
phenoData(object)

## S4 method for signature 'MultiDataSet'
rowRanges(x)

## S4 method for signature 'MultiDataSet,ANY,ANY'
x[[i]]

## S4 method for signature 'MultiDataSet,ANY,ANY,ANY'
x[i, j, k, ..., drop = FALSE]

## S4 method for signature 'MultiDataSet'
subset(x, feat, phe, warnings = TRUE, keep = TRUE)

```

### Arguments

|               |  |
|---------------|--|
| object        | MultiDataSet   |
| set           | Object derived from eSet to be used to fill the slot.                                    |
| dataset.type  | Character with the type of data of the omic set (e.g. expression, methylation...)        |
| dataset.name  | Character with the specific name for this set (NULL by default). It is useful when there |
| sample.tables | Character with the names of the slots with sample data besides phenoData.                |

|                |   |
|----------------|---|
| feature.tables | Character with the names of the slots with feature data besides featureData.                    |
| warnings       | Logical to indicate if warnings will be displayed.  |
| overwrite      | Logical to indicate if the set stored in the slot will be overwritten.                          |
| GRanges        | GenomicRanges to be included in rowRanges slot.   |
| gexpSet        | ExpressionSet to be used to fill the slot.  |
| ...            | Further arguments passed to add_rse or add_se   |
| rnaSet         | ExpressionSet to be used to fill the slot.  |
| methySet       | GenomicRatioSet to be used to fill the slot.  |
| snpSet         | SnpSet to be used to fill the slot.   |
| x              | MultiDataSet  |
| unify.names    | Logical indicating if sample names of the sets should be unified.                               |
| commonSamples  | Logical to indicate if common samples are selected  |
| i              | Character corresponding to selected sample names. They should match the id column of phenoData. |
| j              | Character with the name of the selected tables.   |
| k              | GenomicRange used to filter the features.   |
| drop           | If TRUE, sets with no samples or features will be discarded                                     |
| feat           | Logical expression indicating features to keep  |
| phe            | Logical expression indicating the phenotype of the samples to keep                              |
| keep           | If FALSE, sets where the expression cannot be evaluated will be discarded.                      |

### Details

The names of the three lists (assayData, phenoData and featureData) must be the same.

### Value

MultiDataSet  
MultiDataSet

### Methods (by generic)

- `add_eset`: Method to add an eSet to MultiDataSet.
- `add_genexp`: Method to add a slot of expression to MultiDataSet.
- `add_rnaseq`: Method to add a slot of (RNASeq) expression to MultiDataSet.
- `add_methy`: Method to add a slot of methylation to MultiDataSet from a GenomicRatioSet.
- `add_rse`: Method to add a RangedSummarizedExperiment to MultiDataSet.
- `add_se`: Method to add a SummarizedExperiment to MultiDataSet.
- `add_snps`: Method to add a slot of SNPs to MultiDataSet.
- `add_table`: Method to add a matrix to MultiDataSet.
- `as.list`: Returns a list with the first matrix of each dataset.

- `commonIds`: Get the name of the ids common to all datasets
- `commonSamples`: Get a MultiDataSet only with the samples present in all the tables
- `dims`: Returns the dimensions of the sets
- `w_iclusterplus`: Apply iClusterPlus clustering method to a MultiDataSet object
- `length`: Returns the number of sets into the object.
- `w_mcia`: Apply mcia integration method to a MultiDataSet object
- `names`: Get the names of the slots.
- `ncol`: Get number of samples of each set
- `nrow`: Get number of features of each set
- `rowRangesElements`: Get the name of the datasets that have rowRanges
- `sampleNames`: Get sample names
- `assayData`: Retrieve all assay data blocks.
- `fData`: Retrieve information on features.
- `featureData`: Retrieve information on features.
- `pData`: Retrieve information on experimental phenotypes
- `phenoData`: Retrieve information on experimental phenotypes
- `rowRanges`: Retrieve information on feature ranges.
- `[]`: Get a set from a slot
- `[:`: Subset a MultiDataSet
- `subset`: Filter a subset using feature ids or phenotypes

### Slots

`assayData` List of assayData elements.

`phenoData` List of AnnotatedDataFrame containing the phenoData of each assayData.

`featureData` List of AnnotatedDataFrame containing the featureData of each assayData.

`rowRanges` List of GenomicRanges containing the rowRanges of each assayData.

`extraData` List of other slots of the original object.

`return_method` List of functions used to create the original object.

### See Also

[add\\_eset](#), [add\\_rse](#)

### Examples

```
createMultiDataSet()
```

---

|     |   |
|-----|---|
| opt | <i>Method to get the options sued to create the ResultSet</i> |
|-----|---|

---

**Description**

Method that returns a list with the options used to create the ResultSet.

**Usage**

```
opt(object)
```

**Arguments**

object            A [ResultSet](#) object.

**Value**

A list with the options used to create the ResultSet.

**Examples**

```
data(rset)
opt(rset)
```

---

|         |  |
|---------|--|
| qq_plot | <i>Function to draw a QQ Plot from a vector of numbers</i> |
|---------|--|

---

**Description**

Function to draw a QQ Plot from a vector of numbers

**Usage**

```
qq_plot(values, show.lambda = TRUE)
```

**Arguments**

values            Numeric vector of P.Values  
show.lambda      (default: TRUE) If TRUE shows lambda score for the given model.

**Value**

An object obtained from [ggplot](#).

**Examples**

```
data(rset)
rst <- getAssociation(rset, rid = 1, fName = NULL)
qq_plot(rst$P.Value)
```

---

**ResultSet***Class ResultSet*

---

**Description**

Class `ResultSet` used to encapsulate results from MEAL and omicrexposome.

**Usage**

```
## S4 method for signature 'ResultSet'
fData(object)

## S4 method for signature 'ResultSet'
getAssociation(object, rid = 1, coef = 2, contrast = NULL, fName = NULL, ...)

## S4 method for signature 'ResultSet'
length(x)

## S4 method for signature 'ResultSet'
names(x)

## S4 method for signature 'ResultSet'
opt(object)

## S4 method for signature 'ResultSet,ANY'
plot(
  x,
  y,
  rid = 1,
  coef = 2,
  contrast = NULL,
  type,
  tFC = 2,
  tPV = -log10(0.001),
  show.labels = TRUE,
  show.effect = FALSE,
  show.lambda = TRUE,
  fName = c("chromosome", "start"),
  subset,
  highlight,
  ...
)

## S4 method for signature 'ResultSet'
varLabels(object)

create_resultset(fOrigin, lResults, fData, lOptions = list())
```

**Arguments**

|             |  |
|-------------|--|
| object      | A ResultSet object.  |
| rid         | Name or index of the internal result to be used  |
| coef        | Coefficient to be returned, usually 2  |
| contrast    | Numeric matrix with the contrasts used to perform the analyses                                     |
| fNames      | Character vector with the names of the fData columns that will be added to the results data.frame. |
| ...         | Further arguments passed to <a href="#">topTable</a>   |
| x           | A ResultSet object.  |
| y           | -  |
| type        | Type of plot to be drawn   |
| tFC         | Threshold for log FC of effect   |
| tPV         | Threshold for P-Value  |
| show.labels | (default TRUE) If set to TRUE, features are labelled.  |
| show.effect | (default: TRUE). Used in volcano plot. If TRUE, effect is shown as FC instead of logFC.            |
| show.lambda | (default: TRUE) If TRUE shows lambda score for the given model.                                    |
| subset      | GenomicRanges used to zoom a region in Manhattan plot  |
| highlight   | GenomicRanges used to highlight a region in Manhattan plot   |
| fOrigin     | Character with the function used to run the analysis.  |
| lResults    | List with the results  |
| fData       | List with the feature data.  |
| lOptions    | List with additional options   |

**Value**

An object of class ResultSet

**Methods (by generic)**

- `fData`: Returns data.frame with feature's data.
- `getAssociation`: Getter to obtain the raw data.frame from association and integration analysis.
- `length`: Returns the amount of analyses stored in the ResultSet.
- `names`: Returns the names of the omics data used to create the ResultSet.
- `opt`: Returns a list with the options used to create the ResultSet
- `plot`: Allows to plot a series of plots (QQ plot, Manhattan plot and Volcano plot) depending on the results stored in the ResultSet.
- `varLabels`: Returns the names of the variables of the models used in a ResultSet.

**Slots**

fun\_origin Character containing the function that creates the object.

results List containing the results of the association/integration.

fData List containing the feature-data of the original objects.

options list of options used to create the ResultSet.

**Examples**

```
create_resultset("hello", list(), list(), list())
```

---

rowRangesElements      *Get the name of the datasets that have rowRanges*

---

**Description**

Get the name of the datasets that have rowRanges

**Usage**

```
rowRangesElements(object)
```

**Arguments**

object                  MultiDataSet

**Value**

Character vector with the slots that have rowRanges.

**Examples**

```
multi <- createMultiDataSet()
eset <- new("ExpressionSet", exprs = matrix(runif(10), 5))
eset2 <- new("ExpressionSet", exprs = matrix(runif(8), ncol = 2))
fData(eset2) <- data.frame(chromosome = c("chr1", "chr1", "chr1", "chr1"),
                          start = c(1, 14, 25, 104), end = c(11, 16, 28, 115),
                          stringsAsFactors = FALSE)
multi <- add_eset(multi, eset, "exampledata", GRanges = NA)
multi <- add_genexp(multi, eset2)
rowRangesElements(multi)
```

---

|      |                          |
|------|--------------------------|
| rset | <i>Example ResultSet</i> |
|------|--------------------------|

---

**Description**

Example ResultSet used in the functions examples and in the tests. The script used to generate it can be found in inst/scripts.

**Usage**

```
rset
```

**Format**

```
ResultSet
```

---

|              |  |
|--------------|--|
| volcano_plot | <i>Function to draw a Volcano Plot</i> |
|--------------|--|

---

**Description**

Function that takes two numeric vectors (P-Value and fold change) and draws a volcano plot using ggplot2

**Usage**

```
volcano_plot(  
  pval,  
  fc,  
  names,  
  size = 2,  
  tFC = 2,  
  tPV = -log10(0.001),  
  show.labels = TRUE,  
  show.effect = FALSE  
)
```

**Arguments**

|       |   |
|-------|---|
| pval  | numeric vector of P.Values  |
| fc    | numeric vector of fold change   |
| names | character vector with the feature's names.                              |
| size  | (default 2) Size of the labels in case they are placed.                 |
| tFC   | (default 2) fold change threshold. It can be set to NULL to not filter. |



|             |   |
|-------------|---|
| tPV         | (default $-\log_{10}(0.001)$ ) P-Value threshold. It can be set to NULL to not filter.                |
| show.labels | (default TRUE) If set to TRUE, features are labelled.   |
| show.effect | (default FALSE) If set to TRUE, the X-axis will should $2^{\log_{2}FC}$ instead to the default logFC. |

**Value**

A ggplot object

**Examples**

```
data(rset)
w1 <- getAssociation(rset, rid = 1, fName = NULL)
volcano_plot(w1$P.Value, w1$logFC, rownames(w1))
```

---

|                |  |
|----------------|--|
| w_iclusterplus | <i>Apply iClusterPlus clustering method to a MultiDataSet object</i> |
|----------------|--|

---

**Description**

Method [iClusterPlus](#) is applied on a [MultiDataSet](#) object after getting the common samples along all the contained datasets.

**Usage**

```
w_iclusterplus(object, commonSamples = TRUE, ...)
```

**Arguments**

|               |   |
|---------------|---|
| object        | MultiDataSet  |
| commonSamples | Logical to indicate if common samples are selected        |
| ...           | Arguments passed to function <a href="#">iClusterPlus</a> |

**Value**

A list of results from [iClusterPlus](#)

**Note**

Argument type for [iClusterPlus](#) is filled within the method.

---

`w_mcia`*Apply mcia integration method to a MultiDataSet object*

---

**Description**

Method `mcia` is applied on a `MultiDataSet` object after getting the common samples along all the contained datasets.

**Usage**

```
w_mcia(object, ...)
```

**Arguments**

|                     |  |
|---------------------|--|
| <code>object</code> | <code>MultiDataSet</code>                      |
| <code>...</code>    | Arguments passed to function <code>mcia</code> |

**Value**

A list of results from `mcia`

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