

# Package ‘CoreGx’

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**Type** Package

**Title** Classes and Functions to Serve as the Basis for Other 'Gx' Packages

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**Description** A collection of functions and classes which serve as the foundation for our lab's suite of R packages, such as 'PharmacoGx' and 'RadioGx'. This package was created to abstract shared functionality from other lab package releases to increase ease of maintainability and reduce code repetition in current and future 'Gx' suite programs. Major features include a 'CoreSet' class, from which 'RadioSet' and 'PharmacoSet' are derived, along with get and set methods for each respective slot. Additional functions related to fitting and plotting dose response curves, quantifying statistical correlation and calculating area under the curve (AUC) or survival fraction (SF) are included. For more details please see the included documentation, as well as:

Smirnov, P., Safikhani, Z., El-Hachem, N., Wang, D., She, A., Olsen, C., Freeman, M., Selby, H., Gendoo, D., Grossman, P., Beck, A., Aerts, H., Lupien, M., Goldenberg, A. (2015) <[doi:10.1093/bioinformatics/btv723](https://doi.org/10.1093/bioinformatics/btv723)>. Manem, V., Labie, M., Smirnov, P., Kofia, V., Freeman, M., Koritzinksy, M., Abazeed, M., Haibe-Kains, B., Bratman, S. (2018) <[doi:10.1101/449793](https://doi.org/10.1101/449793)>.

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  'DataMapper-class.R' 'LongTable-accessors.R'  

  'LongTable-utils.R' 'LongTableDataMapper-class.R'  

  'LongTableDataMapper-accessors.R'  

  'TreatmentResponseExperiment-class.R' 'TREDataMapper-class.R'  

  'adaptiveMatthewCor.R' 'aggregate-methods.R'  

  'callingWaterfall.R' 'connectivityScore.R' 'cosinePerm.R'  

  'datasets.R' 'deprecated.R' 'endoaggregate-methods.R'  

  'globals.R' 'gwc.R' 'matthewCor.R' 'mergeAssays-method.R'  

  'methods-coerce.R' 'methods-dim.R' 'methods-dimnames.R'  

  'methods-drugSensitivitySig.R' 'methods-guessMapping.R'  

  'methods-metaConstruct.R' 'methods-subsetTo.R'  

  'optimizeCoreGx.R' 'updateObject-methods.R' 'utilities.R'  

  'utils-iteration.R' 'utils-messages.R' 'utils-optimization.R'  

  'utils-testing.R' 'utils-updateS4.R'
```

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  'methods-metaConstruct.R' 'methods-subsetTo.R'  

  'optimizeCoreGx.R' 'updateObject-methods.R' 'utilities.R'  

  'utils-iteration.R' 'utils-messages.R' 'utils-optimization.R'  

  'utils-testing.R' 'utils-updateS4.R'
```

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---

*Convenience function for converting R code to a call*

---

## Description

This is used to pass through unevaluated R expressions into subset and [, where they will be evaluated in the correct context.

## Usage

`.(....)`

## Arguments

`...` **pairlist** One or more R expressions to convert to calls.

## Value

**call** An R call object containing the quoted expression.

## Examples

```
.(sample_line1 == 'A2058')
```

---

`.assayToBumpyMatrix`     *Convert a LongTable assay into a BumpyMatrix object*

---

## Description

Convert a LongTable assay into a BumpyMatrix object

## Usage

`.assayToBumpyMatrix(LT, assay, rows, cols, sparse = TRUE)`

## Arguments

LT	LongTable with assay to convert into BumpyMatrix
assay	character(1) A valid assay name in LT, as returned by <code>assayNames(LT)</code> .
rows	character() The rownames associated with the assay rowKey
cols	character() The names associated with the assay colKey
sparse	logical(1) Should the BumpyMatrix be sparse (i.e., is the assay sparse).

## Value

BumpyMatrix containing the data from assay.

---

`.convertCSetMolecularProfilesToSE`  
*CSet molecularProfiles from ESets to SEs*

---

## Description

Converts all ExpressionSet objects within the molecularProfiles slot of a CoreSet to SummarizedExperiments

## Usage

`.convertCSetMolecularProfilesToSE(cSet)`

## Arguments

cSet	S4 A CoreSet containing molecular data in ExpressionSets
------	--

## Value

S4 A CoreSet containing molecular data in a SummarizedExperiments

---

.distancePointLine      *Calculate shortest distance between point and line*

---

## Description

This function calculates the shortest distance between a point and a line in 2D space.

## Usage

```
.distancePointLine(x, y, a = 1, b = 1, c = 0)
```

## Arguments

x	x-coordinate of point
y	y-coordinate of point
a	numeric(1) The coefficient in line equation $a * x + b * y + c = 0$ . Defaults to 1.
b	numeric(1) The coefficient in line equation $a * x + b * y + c = 0$ . Defaults to 1.
c	numeric(1) The intercept in line equation $a * x + b * y + c = 0$ . Defaults to 0.

## Value

numeric The shortest distance between a point and a line.

## Examples

```
.distancePointLine(0, 0, 1, -1, 1)
```

---

.distancePointSegment      *Calculate shortest distance between point and line segment*

---

## Description

This function calculates the shortest distance between a point and a line segment in 2D space.

## Usage

```
.distancePointSegment(x, y, x1, y1, x2, y2)
```

## Arguments

x	x-coordinate of point
y	y-coordinate of point
x1	x-coordinate of one endpoint of the line segment
y1	y-coordinate of line segment endpoint with x-coordinate x1
x2	x-coordinate of other endpoint of line segment
y2	y-coordinate of line segment endpoint with x-coordinate x2

**Value**

`numeric` The shortest distance between a point and a line segment

**Examples**

```
.distancePointSegment(0, 0, -1, 1, 1, -1)
```

---

`.fitCurve2`

*Curve fitting via stats::optim L-BFGS-B with fall-back grid/pattern search if convergence is not achieved.*

---

**Description**

Function to fit curve via stats::optim

**Usage**

```
.fitCurve2(
  par,
  x,
  y,
  fn,
  loss,
  lower = -Inf,
  upper = Inf,
  precision = 1e-04,
  density = c(2, 10, 5),
  step = 0.5/density,
  ...,
  loss_args = list(),
  span = 1,
  optim_only = FALSE,
  control = list(factr = 1e-08, ndeps = rep(1e-04, times = length(par)), trace = 0)
)
```

**Arguments**

<code>par</code>	<code>numeric</code> Vector of intial guesses for the parameters. For each index <code>i</code> of <code>par</code> , <code>par[i]</code> must be within the range ( <code>lower[i]</code> , <code>upper[i]</code> ). If only a single <code>upper</code> or <code>lower</code> value is present, that range is used for all parameters in <code>par</code> .
<code>x</code>	<code>numeric</code> Values to evaluate <code>fn</code> for.
<code>y</code>	<code>numeric</code> Target output values to optimze <code>fn</code> against.
<code>fn</code>	<code>function</code> A function to optimize. Any <code>fn</code> arguments passed via <code>...</code> will be treated as constant and removed from the optimization. It is assumed that the first argument is the <code>x</code> value to optimize over and any subsequent arguments are free parameters to be optimized. Transformed to be optim compatible via <code>make_optim_function</code> is the first arguement isn't already <code>par</code> .
<code>loss</code>	<code>character(1)</code> or <code>function</code> Either the name of one of the bundled loss functions (see details) or a custom loss function to compute for the output of <code>fn</code> over <code>x</code> .

lower	numeric(1) Lower bound for parameters. Parallel to par.
upper	numeric(1) Upper bound for parameters. Parallel to par.
precision	numeric smallest step size used in pattern search, once step size drops below this value, the search terminates.
density	numeric how many points in the dimension of each parameter should be evaluated (density of the grid)
step	initial step size for pattern search.
...	pairlist Fall through arguments to fn.
loss_args	list Additional argument to the loss function. These get passed to loss via do.call analagously to using ....
span	numeric Can be safely kept at 1, multiplicative ratio for initial step size in pattern search. Must be larger than precision.
optim_only	logical(1) Should the fall back methods when optim fails be skipped? Default is FALSE.
control	list List of control parameters to pass to optim. See ?optim for details.

## Details

TODO

## Value

numeric Vector of optimal parameters for fn fit against y on the values of x.

## Examples

```
## Not run:
# Four parameter hill curve equation
hillEqn <- function(x, Emin, Emax, EC50, lambda) {
  (Emin + Emax * (x / EC50)^lambda) / (1 + (x / EC50)^lambda)
}
# Make some dummy data
doses <- rev(1000 / (2^(1:20)))
lambda <- 1
Emin <- 1
Emax <- 0.1
EC50 <- median(doses)
response <- hillEqn(doses, Emin=Emin, lambda=lambda, Emax=Emax, EC50=EC50)
nresponse <- response + rnorm(length(response), sd=sd(response)*0.1) # add noise
# 3-parameter optimization
3par <- .fitCurve2(
  par=c(Emax, EC50, lambda),
  x=doses,
  y=nresponse,
  fn=hillEqn,
  Emin=Emin, # set this as constant in the function being optimized (via ...)
  loss=.normal_loss,
  loss_args=list(trunc=FALSE, n=1, scale=0.07),
  upper=c(1, max(doses), 6),
  lower=c(0, min(doses), 0)
)
# 2-parameter optimization
2par <- .fitCurve2(
```

```

par=c(Emax, EC50),
x=doses,
y=response,
fn=hillEqn,
Emin=Emin, # set this as constant in the function being optimized (via ...)
lambda=1,
loss=.normal_loss,
loss_args=list(trunc=FALSE, n=1, scale=0.07),
upper=c(1, max(doses)),
lower=c(0, min(doses))
)
## End(Not run)

```

---

**.intersectList**      *Intersect A List of More Than Two Vectors*

---

### Description

Utility to find the intersection between a list of more than two vectors or lists. This function extends the native intersect function to work on two or more arguments.

### Usage

```
.intersectList(...)
```

### Arguments

...      A list of or any number of vector like objects of the same mode, which could also be operated on by the native R set operations

### Value

A vector like object of the same mode as the first argument, containing only the intersection common to all arguments to the function

### Examples

```

list1 <- list('a', 'b', 'c')
list2 <- list('a', 'c')
list3 <- list('a', 'c', 'd')
listAll <- .intersectList(list1, list2, list3)
listAll

```

---

.longTableToSummarizedExperiment

*Convert LongTable to gDR Style SummarizedExperiment*

---

**Description**

Convert LongTable to gDR Style SummarizedExperiment

**Usage**

```
.longTableToSummarizedExperiment(LT, assay_names)
```

**Arguments**

LT	LongTable to convert to gDR SummarizedExperiment format.
assay_names	character() Names to rename the assays to. These are assumed to be in the same order as assayNames(LT).

**Value**

SummarizedExperiment object with all assay from LT as BumpyMatrixes.

---

.symSetDiffList

*Utility to find the symmetric set difference of a list of two or more vectors or lists*

---

**Description**

The function finds the symmetric set differences between all the arguments, defined as Union(args)-Intersection(args)

**Usage**

```
.symSetDiffList(...)
```

**Arguments**

...	A list of or any number of vector like objects of the same mode, which could also be operated on by the native R set operations
-----	---

**Value**

A vector like object of the same mode as the first argument, containing only the symmetric set difference

**Examples**

```
list1 <- list('a', 'b', 'c')
list2 <- list('a', 'c')
list3 <- list('a', 'c', 'd')
listAll <- .symSetDiffList(list1, list2, list3)
listAll
```

---

.unionList	<i>Utility to find the union between a list of more than two vectors or lists</i>
------------	---

---

### Description

This function extends the native union function to work on two or more arguments.

### Usage

```
.unionList(...)
```

### Arguments

...	A list of or any number of vector like objects of the same mode, which could also be operated on by the native R set operations
-----	---

### Value

A vector like object of the same mode as the first argument, containing all the elements of all arguments passed to the function

### Examples

```
list1 <- list('a', 'b')
list2 <- list('a', 'c')
list3 <- list('c', 'd')
listAll <- .unionList(list1, list2, list3)
listAll
```

---

aggregate, data.table-method	<i>Functional S4 API for aggregation over a data.table object.</i>
------------------------------	--

---

### Description

Compute a group-by operation over a data.table in a functional, pipe compatible format.

### Usage

```
## S4 method for signature 'data.table'
aggregate(
  x,
  by,
  ...,
  subset = TRUE,
  nthread = 1,
  progress = TRUE,
  BPPARAM = NULL,
  enlist = TRUE,
  moreArgs = list()
)
```

## Arguments

x	data.table to compute aggregation over.
by	character One or more valid column names in x to compute groups using.
...	call One or more aggregations to compute for each group by in x. If you name aggregation calls, that will be the column name of the value in the resulting data.table otherwise a default name will be parsed from the function name and its first argument, which is assumed to be the name of the column being aggregated over.
subset	call An R call to evaluate before performing an aggregate. This allows you to aggregate over a subset of columns in an assay but have it be assigned to the parent object. Default is TRUE, which includes all rows. Passed through as the i argument in [.data.table.
nthread	numeric(1) Number of threads to use for split-apply-combine parallelization. Uses BiocParallel::bplapply if nthread > 1 or you pass in BPPARAM. Does not modify data.table threads, so be sure to use setDTthreads for reasonable nested parallelism. See details for performance considerations.
progress	logical(1) Display a progress bar for parallelized computations? Only works if bpprogressbar<- is defined for the current BiocParallel back-end.
BPPARAM	BiocParallelParam object. Use to customized the the parallelization back-end of bplapply. Note, nthread over-rides any settings from BPPARAM as long as bpworkers<- is defined for that class.
enlist	logical(1) Default is TRUE. Set to FALSE to evaluate the first call in ... within data.table groups. See details for more information.
moreArgs	list() A named list where each item is an argument one of the calls in ... which is not a column in the table being aggregated. Use to further parameterize your calls. Please note that these are not added to your aggregate calls unless you specify the names in the call.

## Details

This S4 method override the default aggregate method for a data.frame and as such you need to call aggregate.data.frame directly to get the original S3 method for a data.table.

### Use of Non-Standard Evaluation:

Arguments in ... are substituted and wrapped in a list, which is passed through to the j argument of [.data.table internally. The function currently tries to build informative column names for unnamed arguments in ... by appending the name of each function call with the name of its first argument, which is assumed to be the column name being aggregated over. If an argument to ... is named, that will be the column name of its value in the resulting data.table.

### Enlisting:

The primary use case for enlist=FALSE is to allow computation of dependent aggregations, where the output from a previous aggregation is required in a subsequent one. For this case, wrap your call in {} and assign intermediate results to variables, returning the final results as a list where each list item will become a column in the final table with the corresponding name. Name inference is disabled for this case, since it is assumed you will name the returned list items appropriately. A major advantage over multiple calls to aggregate is that the overhead of parallelization is paid only once even for complex multi-step computations like fitting a model, capturing its parameters, and making predictions using it. It also allows capturing arbitrarily complex calls which can be

recomputed later using the `update`, `TreatmentResponseExperiment`-method. A potential disadvantage is increased RAM usage per thread due to storing intermediate values in variables, as well as any memory allocation overhead associate therewith.

### Value

`data.table` of aggregated results with an `aggregations` attribute capturing metadata about the last aggregation performed on the table.

---

### aggregate,LongTable-method

*Functional API for aggregation over a LongTable or inheriting class*

---

### Description

Compute a group-by operation over a `LongTable` object or it's inheriting classes.

### Usage

```
## S4 method for signature 'LongTable'
aggregate(
  x,
  assay,
  by,
  ...,
  subset = TRUE,
  nthread = 1,
  progress = TRUE,
  BPPARAM = NULL,
  enlist = TRUE,
  moreArgs = list()
)
```

### Arguments

<code>x</code>	<code>LongTable</code> or inheriting class to compute aggregation on.
<code>assay</code>	<code>character(1)</code> The assay to aggregate over.
<code>by</code>	<code>character</code> One or more valid column names in <code>x</code> to compute groups using.
<code>...</code>	<code>call</code> One or more aggregations to compute for each group by in <code>x</code> . If you name aggregation calls, that will be the column name of the value in the resulting <code>data.table</code> otherwise a default name will be parsed from the function name and its first argument, which is assumed to be the name of the column being aggregated over.
<code>subset</code>	<code>call</code> An R call to evaluate before performing an aggregate. This allows you to aggregate over a subset of columns in an assay but have it be assigned to the parent object. Default is <code>TRUE</code> , which includes all rows. Passed through as the <code>i</code> argument in <code>[.data.table</code> .
<code>nthread</code>	<code>numeric(1)</code> Number of threads to use for split-apply-combine parallelization. Uses <code>BiocParallel::bplapply</code> if <code>nthread &gt; 1</code> or you pass in <code>BPPARAM</code> . Does not modify <code>data.table</code> threads, so be sure to use <code>setDTthreads</code> for reasonable nested parallelism. See details for performance considerations.

progress	logical(1) Display a progress bar for parallelized computations? Only works if <code>bpprogressbar&lt;-</code> is defined for the current BiocParallel back-end.
BPPARAM	<code>BiocParallelParam</code> object. Use to customized the the parallelization back-end of <code>bplapply</code> . Note, <code>nthread</code> over-rides any settings from <code>BPPARAM</code> as long as <code>bpworkers&lt;-</code> is defined for that class.
enlist	logical(1) Default is TRUE. Set to FALSE to evaluate the first call in ... within <code>data.table</code> groups. See details for more information.
moreArgs	<code>list()</code> A named list where each item is an argument one of the calls in ... which is not a column in the table being aggregated. Use to further parameterize your calls. Please note that these are not added to your aggregate calls unless you specify the names in the call.

## Details

### Use of Non-Standard Evaluation:

Arguments in ... are substituted and wrapped in a list, which is passed through to the `j` argument of `[.data.table` internally. The function currently tries to build informative column names for unnamed arguments in ... by appending the name of each function call with the name of its first argument, which is assumed to be the column name being aggregated over. If an argument to ... is named, that will be the column name of its value in the resulting `data.table`.

### Enlisting:

The primary use case for `enlist=FALSE` is to allow computation of dependent aggregations, where the output from a previous aggregation is required in a subsequent one. For this case, wrap your call in `{` and assign intermediate results to variables, returning the final results as a list where each list item will become a column in the final table with the corresponding name. Name inference is disabled for this case, since it is assumed you will name the returned list items appropriately. A major advantage over multiple calls to `aggregate` is that the overhead of parallelization is paid only once even for complex multi-step computations like fitting a model, capturing its parameters, and making predictions using it. It also allows capturing arbitrarily complex calls which can be recomputed later using the `update`, `TreatmentResponseExperiment`-method. A potential disadvantage is increased RAM usage per thread due to storing intermediate values in variables, as well as any memory allocation overhead associated therewith.

## Value

`data.table` of aggregation results.

## See Also

`data.table::[.data.table`, `BiocParallel::bplapply`

---

aggregate2

*Functional API for `data.table` aggregation which allows capture of associated aggregate calls so they can be recomputed later.*

---

## Description

Functional API for `data.table` aggregation which allows capture of associated aggregate calls so they can be recomputed later.

## Usage

```
aggregate2(
  x,
  by,
  ...,
  nthread = 1,
  progress = interactive(),
  BPPARAM = NULL,
  enlist = TRUE,
  moreArgs = list()
)
```

## Arguments

x	data.table
by	character One or more valid column names in x to compute groups using.
...	call One or more aggregations to compute for each group by in x. If you name aggregation calls, that will be the column name of the value in the resulting data.table otherwise a default name will be parsed from the function name and its first argument, which is assumed to be the name of the column being aggregated over.
nthread	numeric(1) Number of threads to use for split-apply-combine parallelization. Uses BiocParallel::bplapply if nthread > 1 or you pass in BPPARAM. Does not modify data.table threads, so be sure to use setDTthreads for reasonable nested parallelism. See details for performance considerations.
progress	logical(1) Display a progress bar for parallelized computations? Only works if bpprogressbar<- is defined for the current BiocParallel back-end.
BPPARAM	BiocParallelParam object. Use to customized the the parallelization back-end of bplapply. Note, nthread over-rides any settings from BPPARAM as long as bpworkers<- is defined for that class.
enlist	logical(1) Default is TRUE. Set to FALSE to evaluate the first call in ... within data.table groups. See details for more information.
moreArgs	list() A named list where each item is an argument one of the calls in ... which is not a column in the table being aggregated. Use to further parameterize your calls. Please note that these are not added to your aggregate calls unless you specify the names in the call.

## Details

### Use of Non-Standard Evaluation:

Arguments in ... are substituted and wrapped in a list, which is passed through to the j argument of `[.data.table` internally. The function currently tries to build informative column names for unnamed arguments in ... by appending the name of each function call with the name of its first argument, which is assumed to be the column name being aggregated over. If an argument to ... is named, that will be the column name of its value in the resulting data.table.

### Enlisting:

The primary use case for `enlist=FALSE` is to allow computation of dependent aggregations, where the output from a previous aggregation is required in a subsequent one. For this case, wrap your call in `{` and assign intermediate results to variables, returning the final results as a list where each

list item will become a column in the final table with the corresponding name. Name inference is disabled for this case, since it is assumed you will name the returned list items appropriately. A major advantage over multiple calls to aggregate is that the overhead of parallelization is paid only once even for complex multi-step computations like fitting a model, capturing its parameters, and making predictions using it. It also allows capturing arbitrarily complex calls which can be recomputed later using the update, TreatmentResponseExperiment-method. A potential disadvantage is increased RAM usage per thread due to storing intermediate values in variables, as well as any memory allocation overhead associated therewith.

### Value

`data.table` of aggregation results.

### See Also

`data.table::[.data.table, BiocParallel::bplapply`

---

amcc

*Calculate an Adaptive Matthews Correlation Coefficient*

---

### Description

This function calculates an Adaptive Matthews Correlation Coefficient (AMCC) for two vectors of values of the same length. It assumes the entries in the two vectors are paired. The Adaptive Matthews Correlation Coefficient for two vectors of values is defined as the Maximum Matthews Coefficient over all possible binary splits of the ranks of the two vectors. In this way, it calculates the best possible agreement of a binary classifier on the two vectors of data. If the AMCC is low, then it is impossible to find any binary classification of the two vectors with a high degree of concordance.

### Usage

`amcc(x, y, step.prct = 0, min.cat = 3, nperm = 1000, nthread = 1, ...)`

### Arguments

<code>x, y</code>	Two paired vectors of values. Could be replicates of observations for the same experiments for example.
<code>step.prct</code>	Instead of testing all possible splits of the data, it is possible to test steps of a percentage size of the total number of ranks in x/y. If this variable is 0, function defaults to testing all possible splits.
<code>min.cat</code>	The minimum number of members per category. Classifications with less members fitting into both categories will not be considered.
<code>nperm</code>	The number of permutations to use for estimating significance. If 0, then no p-value is calculated.
<code>nthread</code>	Number of threads to parallelize over. Both the AMCC calculation and the permutation testing is done in parallel.
<code>...</code>	Additional arguments

### Value

Returns a list with two elements. `$amcc` contains the highest 'mcc' value over all the splits, the p value, as well as the rank at which the split was done.

## Examples

```
x <- c(1,2,3,4,5,6,7)
y <- c(1,3,5,4,2,7,6)
amcc(x,y, min.cat=2)
```

---

as

*Coerce a LongTable to a TreatmentResponseExperiment*

---

## Description

Coerce a LongTable into a data.table.

Currently only supports coercing to data.table or data.frame

Coerce a data.table with the proper configuration attributes back to a LongTable

## Arguments

from            A LongTableDataMapper to coerce.

## Value

The data in object, as the child-class TreatmentResponseExperiment.

A data.table with the data from a LongTable.

data.table containing the data from the LongTable, with the ‘longTableDataMapper’ attribute containing the metadata needed to reverse the coercing operation.

LongTable object configured with the longTableDataMapper

data.table with long format of data in from

data.frame with long format of data in from.

SummarizedExperiment with each assay as a BumpyMatrix

A TREDDataMapper object.

## See Also

[TreatmentResponseExperiment](#)

[BumpyMatrix::BumpyMatrix](#)

## Examples

```
data(clevelandSmall_cSet)
TRE <- as(treatmentResponse(clevelandSmall_cSet),
           "TreatmentResponseExperiment")
TRE

as(merckLongTable, 'data.table')

dataTable <- as(merckLongTable, 'data.table')
print(attr(dataTable, 'longTableDataMapper')) # Method doesn't work without this
as(dataTable, 'LongTable')
```

```
SE <- molecularProfilesSlot(clevelandSmall_cSet)[[1]]
as(SE, 'data.table')

SE <- molecularProfilesSlot(clevelandSmall_cSet)[[1]]
as(SE, 'data.frame')
```

**as.long.table***Coerce from data.table to LongTable***Description**

Coerce a data.table with the proper configuration attributes back to a LongTable

**Usage**

```
as.long.table(x)
```

**Arguments**

**x** A data.frame with the 'longTableDataMapper' attribute, containing three lists named assayCols, rowDataCols and colDataCols. This attribute is automatically created when coercing from a LongTable to a data.table.

**Value**

LongTable object configured with the longTableDataMapper

**Examples**

```
dataTable <- as(merckLongTable, 'data.table')
print(attr(dataTable, 'longTableDataMapper')) # Method doesn't work without this
as.long.table(dataTable)
```

**assay**, LongTableDataMapper, ANY-method*Extract the data for an assay from a LongTableDataMapper***Description**

Extract the data for an assay from a LongTableDataMapper

**Usage**

```
## S4 method for signature 'LongTableDataMapper,ANY'
assay(x, i, withDimnames = TRUE)
```

**Arguments**

x LongTableDataMapper The object to retrieve assay data from according to the assayMap slot.

i character(1) Name of an assay in the assayMap slot of x.

withDimnames logical(1) For compatibility with SummarizedExperiment::assay generic. Not used.

**Value**

`data.table` Data for the specified assay extracted from the `rawdata` slot of x.

`assay`, `TREDataMapper`, `ANY`-method

*Extract the data for an assay from a TREDataMapper*

**Description**

Extract the data for an assay from a `TREDataMapper`

**Usage**

```
## S4 method for signature 'TREDataMapper,ANY'
assay(x, i, withDimnames = TRUE)
```

**Arguments**

x `TREDataMapper` The object to retrieve assay data from according to the assayMap slot.

i character(1) Name of an assay in the assayMap slot of x.

withDimnames logical(1) For compatibility with `SummarizedExperiment::assay` generic. Not used.

**Value**

`data.table` Data for the specified assay extracted from the `rawdata` slot of x.

`assayCols`

*Generic to access the assay columns of a rectangular object.*

**Description**

Generic to access the assay columns of a rectangular object.

**Usage**

```
assayCols(object, ...)
```

**Arguments**

object	S4 An object to get assay ids from.
...	Allow new arguments to this generic.

**Value**

Depends on the implemented method.

**Examples**

```
print("Generics shouldn't need examples?")
```

---

assayIndex

*Retrieve and assayIndex*

---

**Description**

Retrieve and assayIndex

**Usage**

```
assayIndex(x, ...)
```

**Arguments**

x	An S4 object.
...	pairlist Allow definition of new parameters for implementations of this generic.

**Value**

An object representing the "assayIndex" of an S4 object.

**Examples**

```
print("Generics shouldn't need examples?")
```

---

assayKeys	<i>Retrieve a set of assayKeys</i>
-----------	------------------------------------

---

### Description

Retrieve a set of assayKeys

### Usage

```
assayKeys(x, ...)
```

### Arguments

x	An S4 object.
...	pairlist Allow definition of new parameters for implementations of this generic.

### Value

An object representing the "assayKeys" of an S4 object.

### Examples

```
print("Generics shouldn't need examples?")
```

---

assays,LongTableDataMapper-method	<i>Extract the data for all assays from a LongTableDataMapper</i>
-----------------------------------	---

---

### Description

Extract the data for all assays from a LongTableDataMapper

### Usage

```
## S4 method for signature 'LongTableDataMapper'
assays(x, withDimnames = TRUE)
```

### Arguments

x	LongTableDataMapper The object to retrieve assay data from according to the assayMap slot.
withDimNames	logical(1) For compatibility with SummarizedExperiment::assay generic. Not used.

### Value

list Data for all assays extracted from the rawdata slot of x as a list of data.tables, where the keys for each table are their id\_columns.

---

assays, TREDatMapper-method

*Extract the data for all assays from a TREDatMapper*

---

## Description

Extract the data for all assays from a TREDatMapper

## Usage

```
## S4 method for signature 'TREDatMapper'
assays(x, withDimnames = TRUE)
```

## Arguments

x	TREDatMapper The object to retrieve assay data from according to the assayMap slot.
withDimnames	logical(1) For compatibility with SummarizedExperiment::assay generic. Not used.

## Value

list Data for all assays extracted from the rawdata slot of x as a list of data.tables, where the keys for each table are their id\_columns.

---

assignment-immutable *Intercept assignment operations for "immutable" S3 objects.*

---

## Description

Prevents modification of objects labelled with the "immutable" S3-class by intercepting assignment during S3-method dispatch and returning an error.

## Usage

```
\method{subset}{immutable}(object, ...) <- value

## S3 replacement method for class 'immutable'
object[...] <- value

## S3 replacement method for class 'immutable'
object[[...]] <- value

## S3 replacement method for class 'immutable'
object$... <- value

## S3 replacement method for class 'immutable'
names(x) <- value
```

```
## S3 replacement method for class 'immutable'
dimnames(x) <- value

\method{colnames}{immutable}(x) <- value

\method{rownames}{immutable}(x) <- value
```

### Arguments

object, x	An R object inheriting from the "immutable" S3-class.
...	Catch subset arguments for various dimensions.
value	Not used.

### Value

None, throws an error.

### Examples

```
immutable_df <- immutable(data.frame(a=1:5, b=letters[1:5]))
# return immutable data.frame
immutable_df[1:4, ]
# return immutable vector
immutable_df$a
```

**buildComboProfiles** *Build an assay table with an S4 object.*

### Description

Build an assay table with an S4 object.

### Usage

```
buildComboProfiles(object, ...)
```

### Arguments

object	S4 An S4 object a list-like slot containing assays for the object.
...	Allow new arguments to be defined for this generic.

### Value

`data.table`.

### Examples

```
"This is a generic method!"
```

---

**buildComboProfiles,LongTable-method**

*Build an assay table with selected assay profiles for drug combinations*

---

**Description**

Build an assay table with selected assay profiles for drug combinations

**Usage**

```
## S4 method for signature 'LongTable'  
buildComboProfiles(object, profiles)
```

**Arguments**

object            LongTable or inheriting class containing curated drug combination data.  
profiles          character a vector of profile names, i.e., column names of assays.

**Value**

A data.table containing fields treatment1id, treatment1dose, treatment2id, treatment2dose, sampleid, which are used as keys to keep track of profiles, along with columns of selected profiles from their assays. Each \*\_1 is the monotheapy profile of treatment 1 in the combination, and the same rule applies to treatment 2.

**Examples**

```
## Not run:  
combo_profile_1 <- buildComboProfiles(tre, c("auc", "SCORE"))  
combo_profile_2 <- buildComboProfiles(tre, c("HS", "EC50", "E_inf", "ZIP"))  
  
## End(Not run)
```

---

**buildLongTable**

*Build a LongTable object*

---

**Description**

Build a LongTable object

**Usage**

```
buildLongTable(from, ...)
```

**Arguments**

from            What to build the LongTable from?  
...              pairlist Allow definition of new parameters for implementations of this generic.

**Value**

Depends on the implemented method

**Examples**

```
print("Generics shouldn't need examples?")
```

**buildLongTable,character-method**

*LongTable build method from character*

**Description**

**LongTable** Create a LongTable object from a single .csv file

**Usage**

```
## S4 method for signature 'character'
buildLongTable(from, rowDataCols, colDataCols, assayCols)
```

**Arguments**

<code>from</code>	character Path to the .csv file containing the data and metadata from which to build the LongTable.
<code>rowDataCols</code>	list List with two character vectors, the first specifying one or more columns to be used as cell identifiers (e.g., cell-line name columns) and the second containing any additional metadata columns related to the cell identifiers.
<code>colDataCols</code>	list List with two character vectors, the first specifying one or more columns to be used as column identifiers (e.g., drug name columns) and the second containing any additional metadata columns related to the column identifiers.
<code>assayCols</code>	list A named list of character vectors specifying how to parse assay columns into a list of data.tables. Each list data.table will be named for the name of corresponding list item and contain the columns specified in the character vector of column names in each list item.

**Value**

A LongTable object containing one or more assays, indexed by rowID and colID.

---

buildLongTable,data.frame-method  
*LongTable build method*

---

**Description**

Create a LongTable object from a single data.table or data.frame object.

**Usage**

```
## S4 method for signature 'data.frame'  
buildLongTable(from, rowDataCols, colDataCols, assayCols)
```

**Arguments**

from	character Path to the .csv file containing the data and metadata from which to build the LongTable.
rowDataCols	list List with two character vectors, the first specifying one or more columns to be used as cell identifiers (e.g., cell-line name columns) and the second containing any additional metadata columns related to the cell identifiers. If you wish to rename any of these columns, assign the new names to their respective character vectors.
colDataCols	list List with two character vectors, the first specifying one or more columns to be used as column identifiers (e.g., drug name columns) and the second containing any additional metadata columns related to the column identifiers. If you wish to rename any of these columns, assign the new names to their respective character vectors.
assayCols	list A named list of character vectors specifying how to parse assay columns into a list of data.tables. Each list data.table will be named for the name of corresponding list item and contain the columns specified in the character vector of column names in each list item. If there are no names for assayCols, the assays will be numbered by instead.

**Value**

A LongTable object containing one or more assays, indexed by rowID and colID.

---

buildLongTable,list-method  
*LongTable build method from list*

---

**Description**

Create a LongTable object from a list containing file paths, data.frames and data.tables.

**Usage**

```
## S4 method for signature 'list'  
buildLongTable(from, rowDataCols, colDataCols, assayCols)
```

**Arguments**

from	list A list containing any combination of character file paths, data.tables and data.frames which will be used to construct the LongTable.
rowDataCols	list List with two character vectors, the first specifying one or more columns to be used as cell identifiers (e.g., cell-line name columns) and the second containing any additional metadata columns related to the cell identifiers.
colDataCols	list List with two character vectors, the first specifying one or more columns to be used as column identifiers (e.g., drug name columns) and the second containing any additional metadata columns related to the column identifiers.
assayCols	list A named list of character vectors specifying how to parse assay columns into a list of data.tables. Each list data.table will be named for the name of corresponding list item and contain the columns specified in the character vector of column names in each list item.

**Value**

A LongTable object constructed with the data in `from`.

**Examples**

```
## Not run:
assayList <- assays(merckLongTable, withDimnames=TRUE)
rowDataCols <- list(rowIDs(merckLongTable), rowMeta(merckLongTable))
colDataCols <- list(colIDs(merckLongTable), colMeta(merckLongTable))
assayCols <- assayCols(merckLongTable)
longTable <- buildLongTable(from=assayList, rowDataCols, colDataCols, assayCols)

## End(Not run)
```

**c.immutable**

*Intercept concatenation for "immutable" class objects to return another "immutable" class object.*

**Description**

Ensures that `c` and `append` to an "immutable" class object return an immutable class object.

**Usage**

```
## S3 method for class 'immutable'
c(x, ...)
```

**Arguments**

x	An R object inheriting from the "immutable" S3-class
...	Objects to concatenate to x.

**Value**

`x` with one or more values appended to it.

---

 callingWaterfall *Drug sensitivity calling using waterfall plots*


---

### Description

1. Sensitivity calls were made using one of IC50, ActArea or Amax

### Usage

```
callingWaterfall(
  x,
  type = c("IC50", "AUC", "AMAX"),
  intermediate.fold = c(4, 1.2, 1.2),
  cor.min.linear = 0.95,
  name = "Drug",
  plot = FALSE
)
```

### Arguments

x	What type of object does this take in?
type	ic50: IC50 values in micro molar (positive values) actarea: Activity Area, that is area under the drug activity curve (positive values) amax: Activity at max concentration (positive values)
intermediate.fold	vector of fold changes used to define the intermediate sensitivities for ic50, actarea and amax respectively
cor.min.linear	numeric The minimum linear correlation to require?
name	character The name of the output to use in plot
plot	boolean Whether to plot the results

### Details

1. Sort log IC50s (or ActArea or Amax) of the samples to generate a “waterfall distribution”
2. Identify cutoff:
  - 3.1 If the waterfall distribution is non-linear (pearson cc to the linear fit  $\leq 0.95$ ), estimate the major inflection point of the log IC50 curve as the point on the curve with the maximal distance to a line drawn between the start and end points of the distribution.
  - 3.2 If the waterfall distribution appears linear (pearson cc to the linear fit  $> 0.95$ ), then use the median IC50 instead.
1. Samples within a 4-fold IC50 (or within a 1.2-fold ActArea or 20% Amax difference) difference centered around this inflection point are classified as being “intermediate”, samples with lower IC50s (or ActArea/Amax values) than this range are defined as sensitive, and those with IC50s (or ActArea/Amax) higher than this range are called “insensitive”.
2. Require at least x sensitive and x insensitive samples after applying these criteria (x=5 in our case).

**Value**

factor Containing the drug sensitivity status of each sample.

**Examples**

```
# Dummy example
1 + 1
```

**checkColumnCardinality**

*Search a data.frame for 1:cardinality relationships between a group of columns (your identifiers) and all other columns.*

**Description**

Search a data.frame for 1:cardinality relationships between a group of columns (your identifiers) and all other columns.

**Usage**

```
checkColumnCardinality(df, group, cardinality = 1, ...)
```

**Arguments**

df	A data.frame to search for 1:cardinality mappings with the columns in group.
group	A character vector of one or more column names to check the cardinality of other columns against.
cardinality	The cardinality of to search for (i.e., 1:cardinality) relationships with the combination of columns in group. Defaults to 1 (i.e., 1:1 mappings).
...	Fall through arguments to data.table::[:]. For developer use. One use case is setting verbose=TRUE to diagnose slow data.table operations.

**Value**

A character vector with the names of the columns with cardinality of 1:cardinality with the columns listed in group.

**Examples**

```
df <- rawdata(exampleDataMapper)
checkColumnCardinality(df, group='treatmentid')
```

---

checkCsetStructure     *A function to verify the structure of a CoreSet*

---

### Description

This function checks the structure of a PharamcoSet, ensuring that the correct annotations are in place and all the required slots are filled so that matching of samples and drugs can be properly done across different types of data and with other studies.

### Usage

```
checkCsetStructure(object, plotDist = FALSE, result.dir = tempdir())
```

### Arguments

object	A CoreSet to be verified
plotDist	Should the function also plot the distribution of molecular data?
result.dir	The path to the directory for saving the plots as a string. Defaults to this R sessions tempdir().

### Value

Prints out messages whenever describing the errors found in the structure of the cSet object passed in.

### Examples

```
checkCsetStructure(clevelandSmall_cSet)
```

---

clevelandSmall\_cSet     *Cleaveland\_mut RadioSet subsetted and cast as CoreSet*

---

### Description

This dataset is just a dummy object derived from the Cleveland\_mut RadioSet in the RadioGx R package. It's contents should not be interpreted and it is only present to test the functions in this package and provide examples

### Usage

```
data(clevelandSmall_cSet)
```

### Format

CoreSet object

### References

Lamb et al. The Connectivity Map: using gene-expression signatures to connect small molecules, genes, and disease. Science, 2006.

## colData, LongTableDataMapper-method

*Convenience method to subset the colData out of the rawData slot using the assigned colDataMap metadata.*

**Description**

Convenience method to subset the colData out of the rawData slot using the assigned colDataMap metadata.

**Usage**

```
## S4 method for signature 'LongTableDataMapper'
colData(x, key = TRUE)
```

**Arguments**

x	LongTableDataMapper object with valid data in the rawData and colDataMap slots.
key	logical(1) Should the table be keyed according to the id_columns of the colDataMap slot? This will sort the table in memory. Default is TRUE.

**Value**

data.table The colData as specified in the colDataMap slot.

## colData, TREDataMapper-method

*Convenience method to subset the colData out of the rawData slot using the assigned colDataMap metadata.*

**Description**

Convenience method to subset the colData out of the rawData slot using the assigned colDataMap metadata.

**Usage**

```
## S4 method for signature 'TREDataMapper'
colData(x, key = TRUE)
```

**Arguments**

x	TREDataMapper object with valid data in the rawData and colDataMap slots.
key	logical(1) Should the table be keyed according to the id_columns of the colDataMap slot? This will sort the table in memory. Default is TRUE.

**Value**

data.table The colData as specified in the colDataMap slot.

---

colIDs	<i>Generic to access the row identifiers for an object.</i>
--------	---

---

## Description

Generic to access the row identifiers for an object.

## Usage

```
colIDs(object, ...)
```

## Arguments

object	S4 An object to get column id columns from.
...	Allow new arguments to this generic

## Value

Depends on the implemented method.

## Examples

```
print("Generics shouldn't need examples?")
```

---

collect_fn_params	<i>Collects all function arguments other than the first into a single list parameter.</i>
-------------------	---

---

## Description

Useful for converting a regular function into a function amenable to optimization via `stats::optim`, which requires all free parameters be passed as a single vector `par`.

## Usage

```
collect_fn_params(fn)
```

## Arguments

fn	function A non-primitive function to refactor such that the first argument becomes the second argument and all other parameters must be passed as a vector to the first argument of the new function via the <code>par</code> parameter.
----	--

## Details

Takes a function of the form `f(x, ...)`, where `...` is any number of additional function parameters (but not literal `...!`) and parses it to a function of the form `f(par, x)` where `par` is a vector of values for `...` in the same order as the arguments appear in `fn`.

**Value**

function A new non-primitive function where the first argument is `par`, which takes a vector of parameters being optimized, and the second argument is the old first argument to `fn` (usually `x` since this is the independent variable to optimize the function over).

---

colMeta	<i>Generic to access the column identifiers for a rectangular object.</i>
---------	---

---

**Description**

Generic to access the column identifiers for a rectangular object.

**Usage**

```
colMeta(object, ...)
```

**Arguments**

object	S4 An object to get column metadata columns from.
...	Allow new arguments to this generic

**Value**

Depends on implemented method.

**Examples**

```
print("Generics shouldn't need examples?")
```

---

connectivityScore	<i>Function computing connectivity scores between two signatures</i>
-------------------	--

---

**Description**

A function for finding the connectivity between two signatures, using either the GSEA method based on the KS statistic, or the gwc method based on a weighted spearman statistic. The GSEA analysis is implemented in the piano package.

**Usage**

```
connectivityScore(
  x,
  y,
  method = c("fgsea", "gwc"),
  nperm = 10000,
  nthread = 1,
  gwc.method = c("spearman", "pearson"),
  ...
)
```

## Arguments

x	A matrix with the first gene signature. In the case of GSEA the vector of values per gene for GSEA in which we are looking for an enrichment. In the case of gwc, this should be a matrix, with the per gene responses in the first column, and the significance values in the second.
y	A matrix with the second signature. In the case of GSEA, this is the vector of up and down regulated genes we are looking for in our signature, with the direction being determined from the sign. In the case of gwc, this should be a matrix of identical size to x, once again with the per gene responses in the first column, and their significance in the second.
method	character string identifying which method to use, out of 'fgsea' and 'gwc'
nperm	numeric, how many permutations should be done to determine significance through permutation testing? The minimum is 100, default is 1e4.
nthread	numeric, how many cores to run parallel processing on.
gwc.method	character, should gwc use a weighted spearman or pearson statistic?
...	Additional arguments passed down to gsea and gwc functions

## Value

numeric a numeric vector with the score and the p-value associated with it

## References

F. Pozzi, T. Di Matteo, T. Aste, 'Exponential smoothing weighted correlations', The European Physical Journal B, Vol. 85, No 6, 2012. DOI: 10.1140/epjb/e2012-20697-x

Varemo, L., Nielsen, J. and Nookaew, I. (2013) Enriching the gene set analysis of genome-wide data by incorporating directionality of gene expression and combining statistical hypotheses and methods. Nucleic Acids Research. 41 (8), 4378-4391. doi: 10.1093/nar/gkt111

## Examples

```

xValue <- c(1,5,23,4,8,9,2,19,11,12,13)
xSig <- c(0.01, 0.001, .97, 0.01,0.01,0.28,0.7,0.01,0.01,0.01,0.01)
yValue <- c(1,5,10,4,8,19,22,19,11,12,13)
ySig <- c(0.01, 0.001, .97,0.01, 0.01,0.78,0.9,0.01,0.01,0.01,0.01)
xx <- cbind(xValue, xSig)
yy <- cbind(yValue, ySig)
rownames(xx) <- rownames(yy) <- c('1','2','3','4','5','6','7','8','9','10','11')
data.cor <- connectivityScore(xx,yy,method='gwc', gwc.method='spearman', nperm=300)

```

## Description

List of deprecated or defunct methods in the CoreGx R package.

## Details

### deprecated:

CoreSet: The CoreSet constructor is being updated to have a new API. This API is currently available via the CoreSet2 constructor. In Bioconductor 3.16, the old constructor will be renamed CoreSet2 and the new constructor will be renamed CoreSet.

### defunct:

buildLongTable: This function no longer works as building a LongTable or TreatmentResponseExperiment now uses a DataMapper and the metaConstruct method. See vignette("LongTable") for a detailed description of how to create a LongTable object.

---

CoreSet

*CoreSet constructor*

---

## Description

A constructor that simplifies the process of creating CoreSets, as well as creates empty objects for data not provided to the constructor. Only objects returned by this constructor are expected to work with the CoreSet methods.

## Usage

```
CoreSet(
  name,
  molecularProfiles = list(),
  sample = data.frame(),
  sensitivityInfo = data.frame(),
  sensitivityRaw = array(dim = c(0, 0, 0)),
  sensitivityProfiles = matrix(),
  sensitivityN = matrix(nrow = 0, ncol = 0),
  perturbationN = array(NA, dim = c(0, 0, 0)),
  curationSample = data.frame(),
  curationTissue = data.frame(),
  curationTreatment = data.frame(),
  treatment = data.frame(),
  datasetType = c("sensitivity", "perturbation", "both"),
  verify = TRUE,
  ...
)
```

## Arguments

name	A character string detailing the name of the dataset
molecularProfiles	A list of SummarizedExperiment objects containing molecular profiles for each molecular data type.
sample	A data.frame containing the annotations for all the sample profiled in the data set, across all data types. Must contain the mandatory sampleid column which uniquely identifies each sample in the object.

<code>sensitivityInfo</code>	A <code>data.frame</code> containing the information for the sensitivity experiments. Must contain a 'sampleid' column with unique identifiers to each sample, matching the <code>sample</code> object and a 'treatmentid' columns with unique identifiers for each treatment, matching the <code>treatment</code> object.
<code>sensitivityRaw</code>	A 3 Dimensional array containing the raw drug dose response data for the sensitivity experiments
<code>sensitivityProfiles</code>	<code>data.frame</code> containing drug sensitivity profile statistics such as IC50 and AUC
<code>sensitivityN, perturbationN</code>	A <code>data.frame</code> summarizing the available sensitivity/perturbation data
<code>curationSample, curationTissue, curationTreatment</code>	A <code>data.frame</code> mapping the names for samples, tissues and treatments used in the data set to universal identifiers used between different CoreSet objects
<code>treatment</code>	A <code>data.frame</code> containing annotations for all treatments profiled in the dataset. Must contain the mandatory <code>treatmentid</code> column which uniquely identifies each treatment in the object.
<code>datasetType</code>	A <code>character(1)</code> string of 'sensitivity', 'perturbation', or 'both' detailing what type of data can be found in the CoreSet, for proper processing of the data
<code>verify</code>	<code>logical(1)</code> Should the function verify the CoreSet and print out any errors it finds after construction?
<code>...</code>	Catch and parse any renamed constructor arguments.

## Details

### WARNING::

Parameters to this function have been renamed!

- `cell` is now `sample`
- `drug` is now `treatment`

## Value

An object of class `CoreSet`

## Examples

```
data(clevelandSmall_cSet)
clevelandSmall_cSet
```

## Description

Documentation for the various setters and getters which allow manipulation of data in the slots of a `CoreSet` object.

**Usage**

```
## S4 method for signature 'CoreSet'  
annotation(object)  
  
## S4 replacement method for signature 'CoreSet,list'  
annotation(object) <- value  
  
## S4 method for signature 'CoreSet'  
dateCreated(object)  
  
## S4 replacement method for signature 'CoreSet,character'  
dateCreated(object) <- value  
  
## S4 method for signature 'CoreSet'  
name(object)  
  
## S4 replacement method for signature 'CoreSet'  
name(object) <- value  
  
## S4 method for signature 'CoreSet'  
sampleInfo(object)  
  
## S4 replacement method for signature 'CoreSet,data.frame'  
sampleInfo(object) <- value  
  
## S4 method for signature 'CoreSet'  
sampleNames(object)  
  
## S4 replacement method for signature 'CoreSet,character'  
sampleNames(object) <- value  
  
## S4 method for signature 'CoreSet'  
treatmentInfo(object)  
  
## S4 replacement method for signature 'CoreSet,data.frame'  
treatmentInfo(object) <- value  
  
## S4 method for signature 'CoreSet'  
treatmentNames(object)  
  
## S4 replacement method for signature 'CoreSet,character'  
treatmentNames(object) <- value  
  
## S4 method for signature 'CoreSet'  
curation(object)  
  
## S4 replacement method for signature 'CoreSet,list'  
curation(object) <- value  
  
## S4 method for signature 'CoreSet'  
datasetType(object)
```

```
## S4 replacement method for signature 'CoreSet,character'
datasetType(object) <- value

## S4 method for signature 'CoreSet'
molecularProfiles(object, mDataType, assay)

## S4 replacement method for signature 'CoreSet,character,character,matrix'
molecularProfiles(object, mDataType, assay) <- value

## S4 replacement method for signature 'CoreSet,character,missing,matrix'
molecularProfiles(object, mDataType, assay) <- value

## S4 replacement method for signature 'CoreSet,missing,missing,list_OR_MAE'
molecularProfiles(object, mDataType, assay) <- value

## S4 method for signature 'CoreSet'
featureInfo(object, mDataType)

## S4 replacement method for signature 'CoreSet,character,data.frame'
featureInfo(object, mDataType) <- value

## S4 method for signature 'CoreSet,character'
phenoInfo(object, mDataType)

## S4 replacement method for signature 'CoreSet,character,data.frame'
phenoInfo(object, mDataType) <- value

## S4 method for signature 'CoreSet,character'
fNames(object, mDataType)

## S4 replacement method for signature 'CoreSet,character,character'
fNames(object, mDataType) <- value

## S4 method for signature 'CoreSet'
mDataNames(object)

## S4 replacement method for signature 'CoreSet'
mDataNames(object) <- value

## S4 method for signature 'CoreSet'
molecularProfilesSlot(object)

## S4 replacement method for signature 'CoreSet,list_OR_MAE'
molecularProfilesSlot(object) <- value

## S4 method for signature 'CoreSet'
sensitivityInfo(object, dimension, ...)

## S4 replacement method for signature 'CoreSet,data.frame'
sensitivityInfo(object, dimension, ...) <- value

## S4 method for signature 'CoreSet'
```

```

sensitivityMeasures(object)

## S4 replacement method for signature 'CoreSet,character'
sensitivityMeasures(object) <- value

## S4 method for signature 'CoreSet'
sensitivityProfiles(object)

## S4 replacement method for signature 'CoreSet,data.frame'
sensitivityProfiles(object) <- value

## S4 method for signature 'CoreSet'
sensitivityRaw(object)

## S4 replacement method for signature 'CoreSet,array'
sensitivityRaw(object) <- value

## S4 method for signature 'CoreSet'
treatmentResponse(object)

## S4 replacement method for signature 'CoreSet,list_OR_LongTable'
treatmentResponse(object) <- value

## S4 method for signature 'CoreSet'
sensNumber(object)

## S4 replacement method for signature 'CoreSet,matrix'
sensNumber(object) <- value

## S4 method for signature 'CoreSet'
pertNumber(object)

## S4 replacement method for signature 'CoreSet,array'
pertNumber(object) <- value

```

## Arguments

object	A CoreSet object.
value	See details.
mDataType	character(1) The name of a molecular datatype to access from the <code>molecularProfiles</code> of a CoreSet object.
assay	character(1) A valid assay name in the <code>SummarizedExperiment</code> of <code>@molecularProfiles</code> of a CoreSet object for data type <code>mDataType</code> .
dimension	See details.
...	See details.

## Details

### **@annotation:**

**annotation:** A list of CoreSet annotations with items: 'name', the name of the object; 'dateCreated', date the object was created; 'sessionInfo', the `sessionInfo()` when the object was created; 'call', the R constructor call; and 'version', the object version.

**annotation<-:** Setter method for the annotation slot. Arguments:

- value: a list of annotations to update the CoreSet with.

**@dateCreated:**

**dateCreated:** character(1) The date the CoreSet object was created, as returned by the date() function.

**dateCreated<-:** Update the 'dateCreated' item in the annotation slot of a CoreSet object. Arguments:

- value: A character(1) vector, as returned by the date() function.

**name:** character(1) The name of the CoreSet, retrieved from the @annotation slot.

**name<-:** Update the @annotation\$name value in a CoreSet object.

- value: character(1) The name of the CoreSet object.

**cellInfo:** data.frame Metadata for all sample in a CoreSet object.

**sampleInfo<-:** assign updated sample annotations to the CoreSet object. Arguments:

- value: a data.frame object.

**sampleNames:** character Retrieve the rownames of the data.frame in the sample slot from a CoreSet object.

**sampleNames<-:** assign new rownames to the sampleInfo data.frame for a CoreSet object. Arguments:

- value: character vector of rownames for the sampleInfo(object) data.frame.

**treatmentInfo:** data.frame Metadata for all treatments in a CoreSet object. Arguments:

- object: CoreSet An object to retrieve treatment metadata from.

**treatmentInfo<-:** CoreSet object with updated treatment metadata. object. Arguments:

- object: CoreSet An object to set treatment metadata for.
- value: data.frame A new table of treatment metadata for object.

**treatmentNames:** character Names for all treatments in a CoreSet object. Arguments:

- object: CoreSet An object to retrieve treatment names from.

**treatmentNames<-:** CoreSet Object with updates treatment names. object. Arguments:

- object: CoreSet An object to set treatment names from.
- value: character A character vector of updated treatment names.

**@curation:**

**curation:** A list of curated mappings between identifiers in the CoreSet object and the original data publication. Contains two data.frames, 'sample' with sample ids and 'tissue' with tissue ids.

**curation<-:** Update the curation slot of a CoreSet object. Arugments:

- value: A list of data.frames, one for each type of curated identifier. For a CoreSet object the slot should contain tissue and sample id data.frames.

**datasetType slot:**

**datasetType:** character(1) The type treatment response in the sensitivity slot. Valid values are 'sensitivity', 'perturbation' or 'both'.

**datasetType<-:** Update the datasetType slot of a CoreSet object. Arguments:

- value: A character(1) vector with one of 'sensitivity', 'perturbation' or 'both'

**@molecularProfiles:**

**molecularProfiles:** matrix() Retrieve an assay in a SummarizedExperiment from the molecularProfiles slot of a CoreSet object with the specified mDataType. Valid mDataType arguments can be found with mDataNames(object). Exclude mDataType and assay to access the entire slot. Arguments:

- assay: Optional character(1) vector specifying an assay in the SummarizedExperiment of the molecularProfiles slot of the CoreSet object for the specified mDataType. If excluded, defaults to modifying the first assay in the SummarizedExperiment for the given mDataType.

**molecularProfiles<-:** Update an assay in a SummarizedExperiment from the molecularProfiles slot of a CoreSet object with the specified mDataType. Valid mDataType arguments can be found with mDataNames(object). Omit mDataType and assay to update the slot.

- assay: Optional character(1) vector specifying an assay in the SummarizedExperiment of the molecularProfiles slot of the CoreSet object for the specified mDataType. If excluded, defaults to modifying the first assay in the SummarizedExperiment for the given mDataType.
- value: A matrix of values to assign to the assay slot of the SummarizedExperiment for the selected mDataType. The rownames and column names must match the associated SummarizedExperiment.

**featureInfo:** Retrieve a DataFrame of feature metadata for the specified mDataType from the molecularProfiles slot of a CoreSet object. More specifically, retrieve the @rowData slot from the SummarizedExperiment from the @molecularProfiles of a CoreSet object with the name mDataType.

**featureInfo<-:** Update the featureInfo(object, mDataType) DataFrame with new feature metadata. Arguments:

- value: A data.frame or DataFrame with updated feature metadata for the specified molecular profile in the molecularProfiles slot of a CoreSet object.

**phenoInfo:** Return the @colData slot from the SummarizedExperiment of mDataType, containing sample-level metadata, from a CoreSet object.

**phenoInfo<-:** Update the @colData slot of the SummarizedExperiment of mDataType in the @molecularProfiles slot of a CoreSet object. This updates the sample-level metadata in-place.

- value: A data.frame or DataFrame object where rows are samples and columns are sample metadata.

**fNames:** character() The features names from the rowData slot of a SummarizedExperiment of mDataType within a CoreSet object.

**fNames:** Updates the rownames of the feature metadata (i.e., rowData) for a SummarizedExperiment of mDataType within a CoreSet object.

- value: character() A character vector of new features names for the rowData of the SummarizedExperiment of mDataType in the @molecularProfiles slot of a CoreSet object. Must be the same length as nrow(featureInfo(object, mDataType)), the number of rows in the feature metadata.

**mDataNames:** character Retrieve the names of the molecular data types available in the `molecularProfiles` slot of a CoreSet object. These are the options which can be used in the `mDataType` parameter of various `molecularProfiles` slot accessors methods.

**mDataNames:** Update the molecular data type names of the `molecularProfiles` slot of a CoreSet object. Arguments:

- `value`: character vector of molecular datatype names, with length equal to `length(molecularProfilesSlot(obj))`

**molecularProfilesSlot:** Return the contents of the `@molecularProfiles` slot of a CoreSet object. This will either be a list or `MultiAssayExperiment` or `SummarizedExperiments`.

**molecularProfilesSlot<-:** Update the contents of the `@molecularProfiles` slot of a CoreSet object. Arguments:

- `value`: A list or `MultiAssayExperiment` or `SummarizedExperiments`. The list and assays should be named for the molecular datatype in each `SummarizedExperiment`.

#### **@treatmentResponse:**

*Arguments::*

- `dimension`: Optional character(1) One of 'treatment', 'sample' or 'assay' to retrieve `rowData`, `colData` or the 'assay\_metadata' assay from the CoreSet `@sensitivity` `LongTable` object, respectively. Ignored with warning if `@treatmentResponse` is not a `LongTable` object.
- `...`: Additional arguments to the `rowData` or `colData` `LongTable` methods. Only used if the `sensitivity` slot contains a `LongTable` object instead of a list and the `dimension` argument is specified.

*Methods::*

**sensitivityInfo:** `DataFrame` or `data.frame` of sensitivity treatment combo by sample metadata for the CoreSet object. When the `dimension` parameter is used, it allows retrieval of the dimension specific metadata from the `LongTable` object in `@treatmentResponse` of a CoreSet object.

**sensitivityInfo<-:** Update the `@treatmentResponse` slot metadata for a CoreSet object. When used without the `dimension` argument it behaves similar to the old CoreSet implementation, where the `@treatmentResponse` slot contained a list with a `$info` `data.frame` item. When the `dimension` argument is used, more complicated assignments can occur where 'sample' modifies the `@sensitivity` `LongTable` `colData`, 'treatment' the `rowData` and 'assay' the 'assay\_metadata' assay. Arguments:

- `value`: A `data.frame` of treatment response experiment metadata, documenting experiment level metadata (mapping to treatments and samples). If the `@treatmentResponse` slot doesn't contain a `LongTable` and `dimension` is not specified, you can only modify existing columns as returned by `sensitivityInfo(object)`.

**sensitivityMeasures:** Get the 'sensitivityMeasures' available in a CoreSet object. Each measure represents some summary of sample sensitivity to a given treatment, such as `ic50`, `ec50`, `AUC`, `AAC`, etc. The results are returned as a character vector with all available metrics for the `PSet` object.

**sensitivityMeasures:** Update the sensitivity measure in a CoreSet object. These values are the column names of the 'profiles' assay and represent various computed sensitivity metrics such as `ic50`, `ec50`, `AUC`, `AAC`, etc.

- `value`: A character vector of new sensitivity measure names, the then length of the character vector must match the number of columns of the 'profiles' assay, excluding metadata and key columns.

**sensitivityProfiles**: Return the sensitivity profile summaries from the sensitivity slot. This data.frame cotanins vaarious sensitivity summary metrics, such as ic50, amax, EC50, aac, HS, etc as columns, with rows as treatment by sample experiments.

**sensitivityProfiles<-**: Update the sensitivity profile summaries the sensitivity slot. Arguments: - value: A data.frame the the same number of rows as as returned by `sensitivityProfiles(object)`, but potentially modified columns, such as the computation of additional summary metrics.

**sensitivityRaw**: Access the raw sensitiity measurements for a CoreSet object. A 3D array where rows are experiment\_ids, columns are doses and the third dimension is metric, either 'Dose' for the doses used or 'Viability' for the sample viability at that dose.

**sensitivityRaw<-**: Update the raw dose and viability data in a CoreSet.

- value: A 3D array object where rows are experiment\_ids, columns are replicates and pages are c('Dose', 'Viability'), with the corresponding dose or viability measurement for that experiment\_id and replicate.

**sensNumber**: Return a count of viability observations in a CoreSet object for each treatment-combo by sample combination.

**sensNumber<-**: Update the 'n' item, which holds a matrix with a count of treatment by sample-line experiment counts, in the list in `@treatmentResponse` slot of a CoreSet object. Will error when `@sensitivity` contains a `LongTable` object, since the counts are computed on the fly. Arguments:

- value: A matrix where rows are samples and columns are treatments, with a count of the number of experiments for each combination as the values.

**pertNumber**: array Summary of available perturbation experiments from in a CoreSet object. Returns a 3D array with the number of perturbation experiments per treatment and sample, and data type.

**pertNumber<-**: Update the `@perturbation$n` value in a CoreSet object, which stores a summary of the available perturbation experiments. Arguments:

- value: A new 3D array with the number of perturbation experiments per treatment and sample, and data type

## Value

Accessors: See details.

Setters: An updated CoreSet object, returned invisibly.

## Examples

```
data(clevelandSmall_cSet)

## @annotation

annotation(clevelandSmall_cSet)

annotation(clevelandSmall_cSet) <- annotation(clevelandSmall_cSet)

dateCreated(clevelandSmall_cSet)

## dateCreated
dateCreated(clevelandSmall_cSet) <- date()

name(clevelandSmall_cSet)
```

```
name(clevelandSmall_cSet) <- 'new_name'

sampleInfo(clevelandSmall_cSet) <- sampleInfo(clevelandSmall_cSet)

sampleNames(clevelandSmall_cSet)

sampleNames(clevelandSmall_cSet) <- sampleNames(clevelandSmall_cSet)

treatmentInfo(clevelandSmall_cSet)

treatmentInfo(clevelandSmall_cSet) <- treatmentInfo(clevelandSmall_cSet)

treatmentNames(clevelandSmall_cSet)

treatmentNames(clevelandSmall_cSet) <- treatmentNames(clevelandSmall_cSet)

## curation
curation(clevelandSmall_cSet)

curation(clevelandSmall_cSet) <- curation(clevelandSmall_cSet)

datasetType(clevelandSmall_cSet)

datasetType(clevelandSmall_cSet) <- 'both'

# No assay specified
molecularProfiles(clevelandSmall_cSet, 'rna') <- molecularProfiles(clevelandSmall_cSet, 'rna')

# Specific assay
molecularProfiles(clevelandSmall_cSet, 'rna', 'exprs') <-
  molecularProfiles(clevelandSmall_cSet, 'rna', 'exprs')

# Replace the whole slot
molecularProfiles(clevelandSmall_cSet) <- molecularProfiles(clevelandSmall_cSet)

featureInfo(clevelandSmall_cSet, 'rna')

featureInfo(clevelandSmall_cSet, 'rna') <- featureInfo(clevelandSmall_cSet, 'rna')

phenoInfo(clevelandSmall_cSet, 'rna')

phenoInfo(clevelandSmall_cSet, 'rna') <- phenoInfo(clevelandSmall_cSet, 'rna')

fNames(clevelandSmall_cSet, 'rna')

fNames(clevelandSmall_cSet, 'rna') <- fNames(clevelandSmall_cSet, 'rna')

mDataNames(clevelandSmall_cSet)

mDataNames(clevelandSmall_cSet) <- mDataNames(clevelandSmall_cSet)

molecularProfilesSlot(clevelandSmall_cSet)

molecularProfilesSlot(clevelandSmall_cSet) <- molecularProfilesSlot(clevelandSmall_cSet)

sensitivityInfo(clevelandSmall_cSet)
```

```

sensitivityInfo(clevelandSmall_cSet) <- sensitivityInfo(clevelandSmall_cSet)

sensitivityMeasures(clevelandSmall_cSet) <- sensitivityMeasures(clevelandSmall_cSet)

sensitivityMeasures(clevelandSmall_cSet) <- sensitivityMeasures(clevelandSmall_cSet)

sensitivityProfiles(clevelandSmall_cSet)

sensitivityProfiles(clevelandSmall_cSet) <- sensitivityProfiles(clevelandSmall_cSet)

head(sensitivityRaw(clevelandSmall_cSet))

sensitivityRaw(clevelandSmall_cSet) <- sensitivityRaw(clevelandSmall_cSet)

treatmentResponse(clevelandSmall_cSet)

treatmentResponse(clevelandSmall_cSet) <- treatmentResponse(clevelandSmall_cSet)

sensNumber(clevelandSmall_cSet)

sensNumber(clevelandSmall_cSet) <- sensNumber(clevelandSmall_cSet)

pertNumber(clevelandSmall_cSet)

pertNumber(clevelandSmall_cSet) <- pertNumber(clevelandSmall_cSet)

```

---

### CoreSet-class

*CoreSet - A generic data container for molecular profiles and treatment response data*

---

### Description

CoreSet - A generic data container for molecular profiles and treatment response data

### Details

The CoreSet (cSet) class was developed as a superclass for pSets in the PharmacoGx and RadioGx packages to contain the data generated in screens of cancer sample lines for their genetic profile and sensitivities to therapy (Pharmacological or Radiation). This class is meant to be a superclass which is contained within the PharmacoSet (pSet) and RadioSet (rSet) objects exported by PharmacoGx and RadioGx. The format of the data is similar for both pSets and rSets, allowing much of the code to be abstracted into the CoreSet super-class. However, the models involved with quantifying sampleular response to Pharmacological and Radiation therapy are widely different, and extension of the cSet class allows the packages to apply the correct model for the given data.

### Slots

annotation See Slots section.  
 molecularProfiles See Slots section.  
 sample See Slots section.  
 treatment See Slots section.

treatmentResponse See Slots section.  
perturbation See Slots section.  
curation See Slots section.  
datasetType See Slots section.

## Slots

- annotation: A list of annotation data about the CoreSet, including the \$name and the session information for how the object was created, detailing the exact versions of R and all the packages used.
- molecularProfiles: A list or MultiAssayExperiment containing CoreSet object.
- sample: A data.frame containing the annotations for all the samples profiled in the data set, across all molecular data types and treatment response experiments.
- treatment: A data.frame containing the annotations for all treatments in the dataset, including the mandatory 'treatmentid' column to uniquely identify each treatment.
- treatmentResponse: A list or LongTable containing all the data for the treatment response experiment, including \$info, a data.frame containing the experimental info, \$raw a 3D array containing raw data, \$profiles, a data.frame containing sensitivity profiles statistics, and \$n, a data.frame detailing the number of experiments for each sample-drug/radiationInfo pair
- perturbation: list containing \$n, a data.frame summarizing the available perturbation data. This slot is currently being deprecated.
- curation: list containing mappings for treatment, sample and tissue names used in the data set to universal identifiers used between different CoreSet objects.
- datasetType: character string of 'sensitivity', 'perturbation', or both detailing what type of data can be found in the CoreSet, for proper processing of the data

## See Also

[CoreSet-accessors](#)

---

CoreSet-utils

*Utility methods for a CoreSet object.*

---

## Description

Documentation for utility methods for a CoreSet object, such as set operations like subset and intersect. See @details for information on different types of methods and their implementations.

## Usage

```
## S4 method for signature 'CoreSet'  
subsetBySample(x, samples)  
  
## S4 method for signature 'CoreSet'  
subsetByTreatment(x, treatments)  
  
## S4 method for signature 'CoreSet'  
subsetByFeature(x, features, mDataTypes)
```

## Arguments

x	A CoreSet object.
samples	character() vector of sample names. Must be valid rownames from sampleInfo(x).
treatments	character() vector of treatment names. Must be valid rownames from treatmentInfo(x). This method does not work with CoreSet objects yet.
features	character() vector of feature names. Must be valid feature names for a given mDataType
mDataTypes	character() One or more molecular data types to to subset features by. Must be valid rownames for the selected SummarizedExperiment mDataTypes.

## Details

### subset methods:

**subsetBySample**: Subset a CoreSet object by sample identifier.

- value: a CoreSet object containing only samples.

### subset methods:

**subsetByTreatment**: Subset a CoreSet object by treatment identifier.

- value: a CoreSet object containing only treatments.

### subset methods:

**subsetByFeature**: Subset a CoreSet object by molecular feature identifier.

- value: a CoreSet object containing only features.

## Value

See details.

## Examples

```
data(clevelandSmall_cSet)

## subset methods

### subsetBySample
samples <- sampleInfo(clevelandSmall_cSet)$sampleid[seq_len(10)]
clevelandSmall_cSet_sub <- subsetBySample(clevelandSmall_cSet, samples)

## subset methods

### subsetByTreatment
#treatments <- treatmentInfo(clevelandSmall_cSet)$treatmentid[seq_len(10)]
#clevelandSmall_cSet_sub <- subsetByTreatment(clevelandSmall_cSet, treatments)

## subset methods

### subsetByFeature
features <- fNames(clevelandSmall_cSet, 'rna')[seq_len(5)]
clevelandSmall_cSet_sub <- subsetByFeature(clevelandSmall_cSet, features, 'rna')
```

---

**CoreSet2***Make a CoreSet with the updated class structure*

---

**Description**

New implementation of the CoreSet constructor to support MAE and TRE. This constructor will be swapped with the original CoreSet constructor as part of an overhaul of the CoreSet class structure.

**Usage**

```
CoreSet2(  
  name = "emptySet",  
  treatment = data.frame(),  
  sample = data.frame(),  
  molecularProfiles = MultiAssayExperiment(),  
  treatmentResponse = LongTable(),  
  datasetType = "sensitivity",  
  perturbation = list(n = array(dim = 3), info = "No perturbation data!"),  
  curation = list(sample = data.frame(), treatment = data.frame())  
)
```

**Arguments**

name	A character(1) vector with the CoreSet objects name.
treatment	A data.frame with treatment level metadata.
sample	A data.frame with sample level metadata for the union of samples in treatmentResponse and molecularProfiles.
molecularProfiles	A MultiAssayExperiment containing one SummarizedExperiment object for each molecular data type.
treatmentResponse	A LongTable or LongTableDataMapper object containing all treatment response data associated with the CoreSet object.
datasetType	A deprecated slot in a CoreSet object included for backwards compatibility. This may be removed in future releases.
perturbation	A deprecated slot in a CoreSet object included for backwards compatibility. This may be removed in future releases.
curation	A list(2) object with two items named treatment and sample with mappings from publication identifiers to standardized identifiers for both annotations, respectively.

**Value**

A CoreSet object storing standardized and curated treatment response and multiomic profile data associated with a given publication.

**Examples**

```
data(clevelandSmall_cSet)  
clevelandSmall_cSet
```

---

**cosinePerm***Cosine Permutations*

---

## Description

Computes the cosine similarity and significance using permutation test. This function uses random numbers, to ensure reproducibility please call `set.seed()` before running the function.

## Usage

```
cosinePerm(  
  x,  
  y,  
  nperm = 1000,  
  alternative = c("two.sided", "less", "greater"),  
  include.perm = FALSE,  
  nthread = 1,  
  ...  
)
```

## Arguments

<code>x</code>	factor is the factors for the first variable
<code>y</code>	factor is the factors for the second variable
<code>nperm</code>	integer is the number of permutations to compute the null distribution of MCC estimates
<code>alternative</code>	string indicates the alternative hypothesis and must be one of "two.sided", "greater" or "less". You can specify just the initial letter. "greater" corresponds to positive association, "less" to negative association. Options are 'two.sided', 'less', or 'greater'
<code>include.perm</code>	boolean indicates whether the estimates for the null distribution should be returned. Default set to 'FALSE'
<code>nthread</code>	integer is the number of threads to be used to perform the permutations in parallel
<code>...</code>	A list of fallthrough parameters

## Value

A list estimate of the cosine similarity, p-value and estimates after random permutations (null distribution) in `include.perm` is set to 'TRUE'

## Examples

```
x <- factor(c(1,2,1,2,1))  
y <- factor(c(2,2,1,1,1))  
cosinePerm(x, y)
```

---

DataMapper-accessors *Accessing and modifying data in a DataMapper object.*

---

## Description

Documentation for the various setters and getters which allow manipulation of data in the slots of a DataMapper object.

## Usage

```
## S4 method for signature 'DataMapper'  
rawdata(object)  
  
## S4 replacement method for signature 'DataMapper,ANY'  
rawdata(object) <- value
```

## Arguments

object	A DataMapper object to get or set data from.
value	A list-like object to assign to the rawdata slot. Should be a <code>data.frame</code> or <code>data.table</code> with the current implementation.

## Details

**rawdata:** Get the raw data slot from a DataMapper object. Returns a list-like containing one or more raw data inputs to the DataMapper object.

**rawdata:** Set the raw data slot from a DataMapper object. **value:** The list-like object to set for the rawdata slot. Note: this currently only supports `data.frame` or `data.table` objects.

## Value

Accessors: See details

Setters: An update DataMapper object, returned invisibly.

## See Also

Other DataMapper-accessors: [LongTableDataMapper-accessors](#), [TREDataMapper-accessors](#)

---

DataMapper-class *An S4 Class For Mapping from Raw Experimental Data to a Specific S4 Object*

---

## Description

This object will be used as a way to abstract away data preprocessing.

## Slots

- **rawdata:** A list-like object containing one or more pieces of raw data that will be processed and mapped to the slots of an S4 object.
- **metadata:** A List of object level metadata.

---

drop_fn_params	<i>Drop parameters from a function and replace them with constants inside the function body.</i>
----------------	--

---

**Description**

Drop parameters from a function and replace them with constants inside the function body.

**Usage**

```
drop_fn_params(fn, args)
```

**Arguments**

fn	function A non-primitive function to remove parameters from (via <code>base:::formals(fn)</code> ).
args	list A list where names are the function arguments (parameters) to remove and the values are the appropriate value to replace the parameter with in the function body.

**Value**

function A new non-primitive function with the parameters named in `args` deleted and their values fixed with the values from `args` in the function body.

---

drugSensitivitySig	<i>Compute the correlation between a molecular feature and treatment response</i>
--------------------	---

---

**Description**

Compute the correlation between a molecular feature and treatment response

**Usage**

```
drugSensitivitySig(object, ...)
```

**Arguments**

object	An object inheriting from the <code>CoreGx::CoreSet</code> class
...	Allow definition of new arguments to this generic

**Value**

A 3D array of genes x drugs x metric

---

endoaggregate	<i>Perform aggregation over an S4 object, but return an object of the same class.</i>
---------------	---

---

## Description

Perform aggregation over an S4 object, but return an object of the same class.

## Usage

```
endoaggregate(x, ...)
```

## Arguments

x	An S4 object to endomorphically aggregate over.
...	pairlist Allow definition of new parameters for implementations of this generic.

## Value

An object with the same class as x.

## Examples

```
print("Generics shouldn't need examples?")
```

---

endoaggregate,LongTable-method
<i>Functional API for endomorphic aggregation over a LongTable or inheriting class</i>

---

## Description

Compute a group-by operation over a LongTable object or its inheriting classes.

## Usage

```
## S4 method for signature 'LongTable'
endoaggregate(
  x,
  ...,
  assay,
  target = assay,
  by,
  subset = TRUE,
  nthread = 1,
  progress = TRUE,
  BPPARAM = NULL,
  enlist = TRUE,
  moreArgs = list()
)
```

## Arguments

x	LongTable or inheriting class to compute aggregation on.
...	call One or more aggregations to compute for each group by in x. If you name aggregation calls, that will be the column name of the value in the resulting data.table otherwise a default name will be parsed from the function name and its first argument, which is assumed to be the name of the column being aggregated over.
assay	character(1) The assay to aggregate over.
target	character(1) The assay to assign the results to. Defaults to assay.
by	character One or more valid column names in x to compute groups using.
subset	call An R call to evaluate before performing an aggregate. This allows you to aggregate over a subset of columns in an assay but have it be assigned to the parent object. Default is TRUE, which includes all rows. Passed through as the i argument in [.data.table.
nthread	numeric(1) Number of threads to use for split-apply-combine parallelization. Uses BiocParallel::bplapply if nthread > 1 or you pass in BPPARAM. Does not modify data.table threads, so be sure to use setDTthreads for reasonable nested parallelism. See details for performance considerations.
progress	logical(1) Display a progress bar for parallelized computations? Only works if bpprogressbar<- is defined for the current BiocParallel back-end.
BPPARAM	BiocParallelParam object. Use to customized the the parallelization back-end of bplapply. Note, nthread over-rides any settings from BPPARAM as long as bpworkers<- is defined for that class.
enlist	logical(1) Default is TRUE. Set to FALSE to evaluate the first call in ... within data.table groups. See details for more information.
moreArgs	list() A named list where each item is an argument one of the calls in ... which is not a column in the table being aggregated. Use to further parameterize your calls. Please note that these are not added to your aggregate calls unless you specify the names in the call.

## Details

### Use of Non-Standard Evaluation:

Arguments in ... are substituted and wrapped in a list, which is passed through to the j argument of [.data.table internally. The function currently tries to build informative column names for unnamed arguments in ... by appending the name of each function call with the name of its first argument, which is assumed to be the column name being aggregated over. If an argument to ... is named, that will be the column name of its value in the resulting data.table.

### Enlisting:

The primary use case for enlist=FALSE is to allow computation of dependent aggregations, where the output from a previous aggregation is required in a subsequent one. For this case, wrap your call in {} and assign intermediate results to variables, returning the final results as a list where each list item will become a column in the final table with the corresponding name. Name inference is disabled for this case, since it is assumed you will name the returned list items appropriately. A major advantage over multiple calls to aggregate is that the overhead of parallelization is paid only once even for complex multi-step computations like fitting a model, capturing its parameters, and making predictions using it. It also allows capturing arbitrarily complex calls which can be recomputed later using the update, TreatmentResponseExperiment-method. A potential disadvantage is increased RAM usage per thread due to storing intermediate values in variables, as well as any memory allocation overhead associated therewith.

**Value**

Object with the same class as x, with the aggregation results assigned to target, using strategy if target is an existing assay in x.

**See Also**

`data.table::[.data.table, BiocParallel::bplapply`

---

exampleDataMapper

*Example LongTableDataMapper*

---

**Description**

A dummy LongTableDataMapper object to be used in package examples.

**Usage**

`data(exampleDataMapper)`

**Format**

LongTableDataMapper object

---

getIntern

*Retrieve the specified item from object internal metadata.*

---

**Description**

Internal slot for storing metadata relevant to the internal operation of an S4 object.

**Usage**

`getIntern(object, x, ...)`

**Arguments**

object	S4 An object with an <code>@.intern</code> slot containing an environment.
x	character One or more symbol names to retrieve from the object <code>@.intern</code> environment.
...	Allow new parameters to be defined for this generic.

**Details**

Warning: This method is intended for developer use and can be ignored by users.

**Value**

Depends on the implemented method

**Examples**

```
print("Generics shouldn't need examples?")
```

---

`getIntern<-`*Set the internal structural metadata for an S4 class*

---

**Description**

Set the internal structural metadata for an S4 class

**Usage**

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

**Arguments**

`object` An R object to update internal structural metadata for.  
`value` An `immutable_list` object, being a class union between `list` and `immutable` S3 classes.

**Value**

Updates the object and returns invisibly.

**Examples**

```
print("Generics shouldn't need examples?")
```

---

`getIntern<-,LongTable,immutable_list-method`*Set the .intern slot of a LongTable*

---

**Description**

Set the `.intern` slot of a `LongTable`

**Usage**

```
## S4 replacement method for signature 'LongTable,immutable_list'  
getIntern(object) <- value
```

**Arguments**

`object` `LongTable`  
`value` An `immutable_list` object, being a class union between `list` and `immutable` S3 classes.

**Value**

Updates the object and returns invisibly.

---

guessMapping	<i>Generic for Guessing the Mapping Between Some Raw Data and an S4 Object</i>
--------------	--

---

## Description

Generic for Guessing the Mapping Between Some Raw Data and an S4 Object

## Usage

```
guessMapping(object, ...)
```

## Arguments

object	An S4 object containing so raw data to guess data to object slot mappings for.
...	Allow new arguments to be defined for this generic.

## Value

A list with mapping guesses as items.

## Examples

```
"Generics shouldn't need examples!"
```

---

guessMapping,LongTableDataMapper-method	<i>Guess which columns in raw experiment data map to which dimensions.</i>
---	--

---

## Description

Checks for columns which are uniquely identified by a group of identifiers. This should be used to help identify the columns required to uniquely identify the rows, columns, assays and metadata of a DataMapper class object.

## Usage

```
## S4 method for signature 'LongTableDataMapper'  
guessMapping(object, groups, subset, data = FALSE)
```

## Arguments

object	A LongTableDataMapper object.
groups	A list containing one or more vector of column names to group-by. The function uses these to determine 1:1 mappings between the combination of columns in each vector and unique values in the raw data columns.
subset	A logical vector indicating whether to subset out mapped columns after each grouping. Must be a single TRUE or FALSE or have the same length as groups, indicating whether to subset out mapped columns after each grouping. This will prevent mapping a column to two different groups.
data	A logical vector indicating whether you would like the data for mapped columns to be returned instead of their column names. Defaults to FALSE for easy use as signing mapped columns to a DataMapper object.

## Details

Any unmapped columns will be added to the end of the returned list in an item called unmapped. The function automatically guesses metadata by checking if any columns have only a single value. This is returned as an additional item in the list.

## Value

A list, where each item is named for the associated groups item the guess is for. The character vector in each item are columns which are uniquely identified by the identifiers from that group.

## Examples

```
guessMapping(exampleDataMapper, groups=list(rows='treatmentid', cols='sampleid'),
subset=FALSE)
```

---

gwc	<i>GWC Score</i>
-----	------------------

---

## Description

Calculate the gwc score between two vectors, using either a weighted spearman or pearson correlation

## Usage

```
gwc(
  x1,
  p1,
  x2,
  p2,
  method.cor = c("pearson", "spearman"),
  nperm = 10000,
  truncate.p = 1e-16,
  ...
)
```

**Arguments**

x1	numeric vector of effect sizes (e.g., fold change or t statistics) for the first experiment
p1	numeric vector of p-values for each corresponding effect size for the first experiment
x2	numeric effect size (e.g., fold change or t statistics) for the second experiment
p2	numeric vector of p-values for each corresponding effect size for the second experiment
method.cor	character string identifying if a pearson or spearman correlation should be used
nperm	numeric how many permutations should be done to determine
truncate.p	numeric Truncation value for extremely low p-values
...	Other passed down to internal functions

**Value**

numeric a vector of two values, the correlation and associated p-value.

**Examples**

```
data(clevelandSmall_cSet)
x <- molecularProfiles(clevelandSmall_cSet, 'rna')[,1]
y <- molecularProfiles(clevelandSmall_cSet, 'rna')[,2]
x_p <- rep(0.05, times=length(x))
y_p <- rep(0.05, times=length(y))
names(x_p) <- names(x)
names(y_p) <- names(y)
gwc(x, x_p, y, y_p, nperm=100)
```

**idCols**

*Generic to access the unique id columns in an S4 object used to*

**Description**

Generic to access the unique id columns in an S4 object used to

**Usage**

```
idCols(object, ...)
```

**Arguments**

object	An S4 object to get id columns from.
...	Allow new arguments to this generic.

**Value**

Depends on the implemented method

## Examples

```
print("Generics shouldn't need examples?")
```

---

immutable

*Constructor for "immutable" S3-class property*

---

## Description

This method should allow any S3 object in R to become immutable by intercepting `[<-`, `[[<-`, `$<-` and `c` during S3-method dispatch and returning an error.

Reverse with call to the `mutable` function.

## Usage

```
immutable(object)

is.immutable(object)

## S3 method for class 'immutable'
print(x, ...)

show.immutable(x)
```

## Arguments

<code>object, x</code>	Any R object which uses S3 method dispatch
<code>...</code>	Fallthrough arguments to <code>print.default</code> .

## Details

The motivation for this class was to create pseudo-private slots in an R S4 object by preventing mutation of those slots outside of the accessors written for the class. It should behave as expected for R object which operate with 'copy-on-modify' semantics, including most base R functions and S3 objects.

An environment was not suitable for this case due to the 'copy-by-reference' semantics, such that normal R assignment, which users assume makes a copy of the object, actually references the same environment in both the original and copy of the object.

**WARNING:** This implementation is unable to intercept modifications to a `data.table` via the `set*` group of methods. This is because these methods are not S3 generics and therefore no mechanism exists for hooking into them to extend their functionality. In general, this helper class will only work for objects with an S3 interface.

## Value

The object with "immutable" prepended to its class attribute.

`logical(1)` Does the object inherit from the "immutable" S3-class?

`None, invisible(NULL)`

**See Also**[assignment-immutable](#), [setOps-immutable](#)**Examples**

```
immutable_list <- immutable(as.list(1:5))
class(immutable_list)
# errors during assignment operations
tryCatch({ immutable_list$new <- 1 }, error=print)

immutable_list <- immutable(as.list(1:5))
is.immutable(immutable_list)
```

---

**is.items***Get the types of all items in a list*

---

**Description**

Get the types of all items in a list

**Usage**

```
is.items(list, ..., FUN = is)
```

**Arguments**

list	A list to get the types from
...	pairlist Additional arguments to FUN
FUN	function or character Either a function, or the name of a function which returns a single logical value. The default function uses <code>is</code> , specify the desired type in <code>...</code> . You can also use other type checking functions such as <code>is.character</code> , <code>is.numeric</code> , or <code>is.data.frame</code> .

**Value**

logical A vector indicating if the list item is the specified type.

**Examples**

```
list <- list(c(1,2,3), c('a','b','c'))
is.items(list, 'character')
```

---

<code>is_optim_compatible</code>	<i>Check whether a function signature is amenable to optimization via stats:::optim.</i>
----------------------------------	--

---

## Description

Functions compatible with `optim` have the parameter named `par` as their first formal argument where each value is a respective free parameter to be optimized.

## Usage

```
is_optim_compatible(fn)
```

## Arguments

<code>fn</code>	function A non-primitive function.
-----------------	------------------------------------

## Value

`logical(1)` TRUE if the first value of `formalArg(fn)` is "par", otherwise FALSE.

---

<code>lapply,MultiAssayExperiment-method</code>	<i>lapply lapply method for MultiAssayExperiment</i>
---	--

---

## Description

`lapply` `lapply` method for `MultiAssayExperiment`

## Usage

```
## S4 method for signature 'MultiAssayExperiment'
lapply(X, FUN, ...)
```

## Arguments

<code>X</code>	A <code>MultiAssayExperiment</code> object.
<code>FUN</code>	A function to be applied to each <code>SummarizedExperiment</code> in <code>X</code> in <code>X</code> .
<code>...</code>	Fall through parameters to <code>FUN</code>

## Value

A `MultiAssayExperiment` object, modified such that `experiments(X) <- endoapply(experiments(X), FUN, ...)`.

---

list\_OR\_LongTable-class

*A class union to allow multiple types in a CoreSet slot*

---

**Description**

A class union to allow multiple types in a CoreSet slot

---

## LongTable

*LongTable constructor method*

---

**Description**

LongTable constructor method

**Usage**

```
LongTable(
  rowData,
  rowIDs,
  colData,
  colIDs,
  assays,
  assayIDs,
  metadata = list(),
  keep.rownames = FALSE
)
```

**Arguments**

rowData	data.frame	A rectangular object coercible to a data.table.
rowIDs	character	A vector of rowData column names needed to uniquely identify each row in a LongTable.
colData	data.frame	A rectangular object coercible to a data.table.
colIDs	character	A vector of colData column names needed to uniquely identify each column in a LongTable.
assays	list	A list of rectangular objects, each coercible to a data.table. Must be named and item names must match the assayIDs list.
assayIDs	list	A list of character vectors specifying the columns needed to uniquely identify each row in an assay. Names must match the assays list.
metadata	list	A list of one or more metadata items associated with a LongTable experiment.
keep.rownames	logical(1) or character(1)	Should rownames be retained when coercing to data.table inside the constructor. Default is FALSE. If TRUE, adds a 'rn' column to each rectangular object that gets coerced from data.frame to data.table. If a string, that becomes the name of the rownames column.

**Value**

A LongTable object containing the data for a treatment response experiment and configured according to the rowIDs and colIDs arguments.

**Examples**

```
"See vignette('The LongTable Class', package='CoreGx')"
```

LongTable-accessors     *Accessing and modifying information in a LongTable*

**Description**

Documentation for the various setters and getters which allow manipulation of data in the slots of a LongTable object.

**Value**

Accessors: See details.

Setters: An updated LongTable object, returned invisibly.

**Examples**

```
data(merckLongTable)
```

LongTable-class     *LongTable class definition*

**Description**

Define a private constructor method to be used to build a LongTable object.

This is used as an alternative to R attributes for storing structural metadata of an S4 objects.

Add or replace an assay in a LongTable by name. Currently this function only works when the assay has all columns in row and column data tables (i.e., when assays is retured withDimnames=TRUE).

Select an assay from within a LongTable object.

**Usage**

```
## S4 method for signature 'LongTable'
rowIDs(object, data = FALSE, key = FALSE)

## S4 method for signature 'LongTable'
rowMeta(object, data = FALSE, key = FALSE)

## S4 method for signature 'LongTable'
colIDs(object, data = FALSE, key = FALSE)
```

```
## S4 method for signature 'LongTable'
colMeta(object, data = FALSE, key = FALSE)

## S4 method for signature 'LongTable'
idCols(object)

## S4 method for signature 'LongTable'
assayIndex(x)

## S4 method for signature 'LongTable'
assayKeys(x, i)

## S4 method for signature 'LongTable'
assayCols(object, i)

## S4 method for signature 'LongTable,character'
getIntern(object, x)

## S4 method for signature 'LongTable,missing'
getIntern(object, x)

## S4 method for signature 'LongTable'
rowData(x, key = FALSE, use.names = FALSE, ...)

## S4 replacement method for signature 'LongTable'
rowData(x, ...) <- value

## S4 method for signature 'LongTable'
colData(x, key = FALSE, dimnames = FALSE, ...)

## S4 replacement method for signature 'LongTable,ANY'
colData(x, ...) <- value

## S4 method for signature 'LongTable'
assays(
  x,
  withDimnames = TRUE,
  metadata = withDimnames,
  key = !withDimnames,
  ...
)

## S4 replacement method for signature 'LongTable,list'
assays(x, withDimnames = TRUE, ...) <- value

## S4 method for signature 'LongTable,ANY'
assay(
  x,
  i,
  withDimnames = TRUE,
  summarize = withDimnames,
  metadata = !summarize,
```

```

key = !(summarize || withDimnames),
...
)

## S4 replacement method for signature 'LongTable,ANY'
assay(x, i) <- value

## S4 method for signature 'LongTable'
assayNames(x)

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

## S4 method for signature 'LongTable'
dim(x)

## S4 method for signature 'LongTable'
colnames(x)

## S4 method for signature 'LongTable'
rownames(x)

## S4 method for signature 'LongTable'
dimnames(x)

```

### Arguments

object	LongTable
data	logical Should the colData for the metadata columns be returned instead of the column names? Default is FALSE.
key	logical Should the key columns also be returned? Defaults to !withDimnames. This is incompatible with summarize=TRUE, which will drop the key columns regardless of the value of this argument.
x	The LongTable object to retrieve the dimnames for.
i	character(1) name or integer index of the desired assay.
use.names	logical This parameter is just here to stop matching the positional argument to use.names from the rowData generic. It doesn't do anything at this time and can be ignored.
...	For developer use only! Pass raw=TRUE to return the slot for modification by reference.
value	A data.frame or data.table to update the assay data with. This must at minimum contain the row and column data identifier columns to allow correctly mapping the assay keys. We recommend modifying the results returned by assay(longTable, 'assayName', withDimnames=TRUE). For convenience, both the [[ and \$ LongTable accessors return an assay with the dimnames.
withDimnames	logical(1) Should the dimension names be returned joined to the assay. This retrieves both the row and column identifiers and returns them joined to the assay. For
metadata	logical(1) Should all of the metadata also be joined to the assay. This is useful when modifying assays as the resulting list has all the information needed to recreate the LongTable object. Defaults to withDimnames.

summarize	logical(1) If the assays is a summary where some of idCols(x) are not in assayKeys(x, i), then those missing columns are dropped. Defaults to FALSE. When metadata is TRUE, only metadata columns with 1:1 cardinality with the assay keys for i.
‘x’	A LongTable or inheriting class.
‘i’	An optional valid assay name or index in x.

### Value

LongTable object containing the assay data from a treatment response experiment
A character vector of rowData column names if data is FALSE, otherwise a data.table with the data from the rowData id columns.
A character vector of rowData column names if data is FALSE, otherwise a data.table with the data from the rowData metadata columns.
A character vector of colData column names if data is FALSE, otherwise a data.table with the data from the colData id columns.
A character vector of colData column names if data is FALSE, otherwise a data.table with the data from the colData metadata columns.
character A character vector containing the unique rowIDs and colIDs in a LongTable object.
A mutable copy of the "assayIndex" for x
A mutable copy of the "assyKeys" for x
A list of character vectors containing the value column names for each assay if i is missing, otherwise a character vector of value column names for the selected assay.
immutable value of x if length(x) == 1 else named list of values for all symbols in x.
An immutable list.
A data.table containing rowID, row identifiers, and row metadata.
A copy of the LongTable object with the rowData slot updated.
A data.table containing row identifiers and metadata.
A copy of the LongTable object with the colData slot updated.
A list of data.table objects, one per assay in the object.
A copy of the LongTable with the assays modified.
LongTable With updated assays slot.
character Names of the assays contained in the LongTable.
numeric Vector of object dimensions.
character Vector of column names.
character Vector of row names.
list List with two character vectors, one for row and one for column names.

### Methods (by generic)

- `rowMeta(LongTable)`: Get the names of the non-id columns from rowData.
- `colIDs(LongTable)`: Get the names of the columns in colData required to uniquely identify each row.
- `colMeta(LongTable)`: Get the names of the non-id columns in the colData data.table.

- `idCols(LongTable)`: Get the names of all id columns.
- `assayIndex(LongTable)`: Get the assayIndex item from the objects internal metadata.
- `assayKeys(LongTable)`: Get the assayKeys item from the objects internal metadata.
- `assayCols(LongTable)`: Get a list of column names for each assay in the object.
- `getIntern(object = LongTable, x = character)`: Access structural metadata present within a LongTable object. This is mostly for developer use.
- `getIntern(object = LongTable, x = missing)`: Access all structural metadata present within a LongTable object. This is primarily for developer use.
- `rowData(LongTable)`: Get the row level annotations for a LongTable object.
- `rowData(LongTable) <- value`: Update the row annotations for a LongTable object. Currently requires that all columns in `rowIDs(longTable)` be present in value.
- `colData(LongTable)`: Get the column level annotations for a LongTable object.
- `colData(x = LongTable) <- value`: Update the colData of a LongTable object. Currently requires that all of the `colIDs(longTable)` be in the value object.
- `assays(LongTable)`: Get a list containing all the assays in a LongTable.
- `assays(x = LongTable) <- value`: Update the assays in a LongTable object. The `rowIDs` and `colIDs` must be present in all assays to allow successfully remapping the keys. We recommend modifying the list returned by `assays(longTable, withDimnames=TRUE)` and the reassigning to the LongTable.
- `assay(x = LongTable, i = ANY)`: Retrieve an assay `data.table` object from the assays slot of a LongTable object.
- `assay(x = LongTable, i = ANY) <- value`:
- `assayNames(LongTable)`: Return the names of the assays contained in a LongTable
- `x[[i]`: Get an assay from a LongTable object. This method returns the row and column annotations by default to make assignment and aggregate operations easier.
- `dim(LongTable)`: Get the number of row annotations by the number of column annotations from a LongTable object. Please note that row x columns does not necessarily equal the number of rows in an assay, since it is not required for each assay to have every row or column present.
- `colnames(LongTable)`: Retrieve the pseudo-colnames of a LongTable object, these are constructed by pasting together the `colIDs(longTable)` and can be used in the subset method for regex based queries.
- `rownames(LongTable)`: Retrieve the pseudo-rownames of a LongTable object, these are constructed by pasting together the `rowIDs(longTable)` and can be used in the subset method for regex based queries.
- `dimnames(LongTable)`: Get the pseudo-dimnames for a LongTable object. See `colnames` and `rownames` for more information.

## Slots

`rowData` See Slots section.  
`colData` See Slots section.  
`assays` See Slots section.  
`metadata` See Slots section.  
`.intern` See Slots section.

### Slots

- *rowData*: A `data.table` containing the metadata associated with the row dimension of a `LongTable`.
- *colData*: A `data.table` containing the metadata associated with the column dimension of a `LongTable`.
- *assays*: A list of `data.tables`, one for each assay in a `LongTable`.
- *metadata*: An optional list of additional metadata for a `LongTable` which doesn't map to one of the dimensions.
- *.intern*: An immutable list that holds internal structural metadata about a `LongTable` object, such as which columns are required to key the object.

### Examples

```
rowIDs(merckLongTable)

rowMeta(merckLongTable)

colIDs(merckLongTable)

colMeta(merckLongTable)

idCols(merckLongTable)

assayIndex(nci_TRE_small)

assayKeys(nci_TRE_small)
assayKeys(nci_TRE_small, "sensitivity")
assayKeys(nci_TRE_small, 1)

assayCols(merckLongTable)

getIntern(merckLongTable, 'rowIDs')
getIntern(merckLongTable, c('colIDs', 'colMeta'))

getIntern(merckLongTable)

rowData(merckLongTable)

rowData(merckLongTable) <- rowData(merckLongTable)

colData(merckLongTable)

# Get the keys as well, mostly for internal use
colData(merckLongTable, key=TRUE)

colData(merckLongTable) <- colData(merckLongTable)

assays(merckLongTable)

assays(merckLongTable) <- assays(merckLongTable, withDimnames=TRUE)

# Default annotations, just the key columns
assay(merckLongTable, 'sensitivity')
assay(merckLongTable, 1)
```

```

# With identifiers joined
assay(merckLongTable, 'sensitivity', withDimnames=TRUE)

# With identifiers and metadata
assay(merckLongTable, 'profiles', withDimnames=TRUE, metadata=TRUE)

assay(merckLongTable, 'sensitivity') <-
  assay(merckLongTable, 'sensitivity', withDimnames=TRUE)
assay(merckLongTable, 'sensitivity') <- merckLongTable$sensitivity

assayNames(merckLongTable)
names(merckLongTable)

merckLongTable[['sensitivity']]

dim(merckLongTable)

dim(merckLongTable)

head(colnames(merckLongTable))

head(rownames(merckLongTable))

lapply(dimnames(merckLongTable), head)

```

---

<b>LongTableDataMapper</b>	<i>Constructor for the LongTableDataMapper class, which maps from one or more raw experimental data files to the slots of a LongTable object.</i>
----------------------------	---

---

## Description

Constructor for the LongTableDataMapper class, which maps from one or more raw experimental data files to the slots of a LongTable object.

## Usage

```

LongTableDataMapper(
  rawdata = data.frame(),
  rowDataMap = list(character(), character()),
  colDataMap = list(character(), character()),
  assayMap = list(list(character(), character())),
  metadataMap = list(character())
)

```

## Arguments

<b>rawdata</b>	A <code>data.frame</code> of raw data from a treatment response experiment. This will be coerced to a <code>data.table</code> internally. We recommend using joins to aggregate your raw data if it is not present in a single file.
----------------	--

rowDataMap	A list-like object containing two character vectors. The first is column names in rawdata needed to uniquely identify each row, the second is additional columns which map to rows, but are not required to uniquely identify them. Rows should be drugs.
colDataMap	A list-like object containing two character vectors. The first is column names in rawdata needed to uniquely identify each column, the second is additional columns which map to rows, but are not required to uniquely identify them. Columns should be samples.
assayMap	A list-like where each item is a list with two character vectors defining an assay, the first containing the identifier columns in rawdata needed to uniquely identify each row an assay, and the second the rawdata columns to be mapped to that assay. The names of assayMap will be the names of the assays in the LongTable that is created when calling metaConstruct on this DataMapper object. If the character vectors have names, the value columns will be renamed accordingly.
metadataMap	A list-like where each item is a character vector of rawdata column names to assign to the @metadata of the LongTable, where the name of that assay is the name of the list item. If names are omitted, assays will be numbered by their index in the list.

## Details

The guessMapping method can be used to test hypotheses about the cardinality of one or more sets of identifier columns. This is helpful to determine the id columns for rowDataMap and colDataMap, as well as identify columns mapping to assays or metadata.

To attach metadata not associated with rawdata, please use the metadata assignment method on your LongTableDataMapper. This metadata will be merged with any metadata from metadataMap and added to the LongTable which this object ultimately constructs.

## Value

A LongTable object, with columns mapped to it's slots according to the various maps in the LongTableDataMapper object.

## See Also

[guessMapping](#)

## Examples

```
data(exampleDataMapper)  
exampleDataMapper
```

---

## LongTableDataMapper-accessors

*Accessing and modifying data in a LongTableDataMapper object.*

---

## Description

Documentation for the various setters and getters which allow manipulation of data in the slots of a LongTableDataMapper object.

## Usage

```
## S4 replacement method for signature 'LongTableDataMapper,list'
rawdata(object) <- value

## S4 method for signature 'LongTableDataMapper'
rowDataMap(object)

## S4 replacement method for signature 'LongTableDataMapper,list_OR_List'
rowDataMap(object) <- value

## S4 method for signature 'LongTableDataMapper'
colDataMap(object)

## S4 replacement method for signature 'LongTableDataMapper,list_OR_List'
colDataMap(object) <- value

## S4 method for signature 'LongTableDataMapper'
assayMap(object)

## S4 replacement method for signature 'LongTableDataMapper,list_OR_List'
assayMap(object) <- value

## S4 method for signature 'LongTableDataMapper'
metadataMap(object)

## S4 replacement method for signature 'LongTableDataMapper,list_OR_List'
metadataMap(object) <- value
```

## Arguments

object	A LongTableDataMapper object to get or set data from.
value	See details.

## Details

**rawdata:** Get the raw data slot from a LongTableDataMapper object. Returns a list-like containing one or more raw data inputs to the LongTableDataMapper object.

**rawdata:** Set the raw data slot from a LongTableDataMapper object. **value:** The list-like object to set for the rawdata slot. Note: this currently only supports data.frame or data.table objects.

**rowDataMap:** list of two character vectors, the first are the columns required to uniquely identify each row of a LongTableDataMapper and the second any additional row-level metadata. If the character vectors have names, the resulting columns are automatically renamed to the item name of the specified column.

**rowDataMap<-:** Update the @rowDataMap slot of a LongTableDataMapper object, returning an invisible NULL. Arguments:

- **value:** A list or List where the first item is the names of the identifier columns – columns needed to uniquely identify each row in rowData – and the second item is the metadata associated with those the identifier columns, but not required to uniquely identify rows in the object rowData.

**colDataMap**: list of two character vectors, the first are the columns required to uniquely identify each row of a LongTableDataMapper and the second any additional col-level metadata. If the character vectors have names, the resulting columns are automatically renamed to the item name of the specified column.

**colDataMap<-**: Update the @colDataMap slot of a LongTableDataMapper object, returning an invisible NULL. Arguments:

- value: A list or List where the first item is the names of the identifier columns – columns needed to uniquely identify each row in colData – and the second item is the metadata associated with those the identifier columns, but not required to uniquely identify rows in the object rowData.

**assayMap**: A list of character vectors. The name of each list item will be the assay in a LongTableDataMapper object that the columns in the character vector will be assigned to. Column renaming occurs automatically when the character vectors have names (from the value to the name).

**assayMap<-**: Updates the @assayMap slot of a LongTableDataMapper object, returning an invisible NULL. Arguments:

- value: A list of character vectors, where the name of each list item is the name of an assay and the values of each character vector specify the columns mapping to the assay in the S4 object the LongTableDataMapper constructs.

**metadataMap**: A list of character vectors. Each item is an element of the constructed objects @metadata slot.

**metadataMap<-**: Updates LongTableDataMapper object in-place, then returns an invisible(NULL). Arguments:

- value: A list of character vectors. The name of each list item is the name of the item in the @metadata slot of the LongTableDataMapper object created when metaConstruct is called on the DataMapper, and a character vector specifies the columns of @rawdata to assign to each item.

## Value

Accessors: See details

Setters: An update LongTableDataMapper object, returned invisibly.

## See Also

Other DataMapper-accessors: [DataMapper-accessors](#), [TREDataMapper-accessors](#)

## Examples

```
rowDataMap(exampleDataMapper)

rowDataMap(exampleDataMapper) <- list(c('treatmentid'), c())

colDataMap(exampleDataMapper)

colDataMap(exampleDataMapper) <- list(c('sampleid'), c())

assayMap(exampleDataMapper)
```

```
assayMap(exampleDataMapper) <- list(sensitivity=c(viability1='viability'))

metadataMap(exampleDataMapper)

metadataMap(exampleDataMapper) <- list(object_metadata=c('metadata'))
```

---

## LongTableDataMapper-class

*A Class for Mapping Between Raw Data and an LongTable Object*

---

### Description

A Class for Mapping Between Raw Data and an LongTable Object

### Usage

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

### Arguments

object            A LongTableDataMapper to display in the console.

### Value

invisible Prints to console.

### Functions

- `show(LongTableDataMapper)`: Show method for LongTableDataMapper. Determines how the object is displayed in the console.

### Slots

`rawdata` See Slots section.  
`rowDataMap` See Slots section.  
`colDataMap` See Slots section.  
`assayMap` See Slots section.  
`metadataMap` See Slots section.

### Slots

- `rowDataMap`: A list-like object containing two character vectors. The first is column names in `rawdata` needed to uniquely identify each row, the second is additional columns which map to rows, but are not required to uniquely identify them. Rows should be drugs.
- `colDataMap`: A list-like object containing two character vectors. The first is column names in `rawdata` needed to uniquely identify each column, the second is additional columns which map to rows, but are not required to uniquely identify them. Columns should be samples.

- assayMap: A list-like where each item is a list with two elements specifying an assay, the first being the identifier columns in `rawdata` needed to uniquely identify each row an assay, and the second a list of `rawdata` columns to be mapped to that assay. The names of `assayMap` will be the names of the assays in the `LongTable` that is created when calling `metaConstruct` on this `DataMapper` object.
- metadataMap: A list-like where each item is a character vector of `rawdata` column names to assign to the `@metadata` of the `LongTable`, where the name of that assay is the name of the list item. If names are omitted, assays will be numbered by their index in the list.
- `rawdata`: A list-like object containing one or more pieces of raw data that will be processed and mapped to the slots of an S4 object.
- `metadata`: A List of object level metadata.

## Examples

```
show(exampleDataMapper)
```

---

`make_optim_function`     *Takes a non-primitive R function and refactors it to be compatible with optimization via `stats:::optim`.*

---

## Description

Takes a non-primitive R function and refactors it to be compatible with optimization via `stats:::optim`.

## Usage

```
make_optim_function(fn, ...)
```

## Arguments

<code>fn</code>	function A non-primitive function
<code>...</code>	Arguments to <code>fn</code> to fix for before building the function to be optimized. Useful for reducing the number of free parameters in an optimization if there are insufficient degrees of freedom.

## See Also

[drop\\_fn\\_params](#), [collect\\_fn\\_params](#)

---

**mcc***Compute a Mathews Correlation Coefficient*

---

## Description

The function computes a Matthews correlation coefficient for two factors provided to the function. It assumes each factor is a factor of class labels, and the entries are paired in order of the vectors.

## Usage

```
mcc(  
  x,  
  y,  
  nperm = 1000,  
  nthread = 1,  
  alternative = c("two.sided", "less", "greater"),  
  ...  
)
```

## Arguments

<code>x, y</code>	factor of the same length with the same number of levels
<code>nperm</code>	numeric number of permutations for significance estimation. If 0, no permutation testing is done
<code>nthread</code>	numeric can parallelize permutation testing using BiocParallel's <code>bplapply</code>
<code>alternative</code>	indicates the alternative hypothesis and must be one of "two.sided", "greater" or "less". You can specify just the initial letter. "greater" corresponds to positive association, "less" to negative association.
<code>...</code>	list Additional arguments

## Details

Please note: we recommend you call `set.seed()` before using this function to ensure the reproducibility of your results. Write down the seed number or save it in a script if you intend to use the results in a publication.

## Value

A list with the MCC as the `$estimate`, and p value as `$p.value`

## Examples

```
x <- factor(c(1,2,1,2,3,1))  
y <- factor(c(2,1,1,1,2,2))  
mcc(x,y)
```

---

merckLongTable *Merck Drug Combination Data LongTable*

---

## Description

This is a LongTable object created from some drug combination data provided to our lab by Merck.

## Usage

```
data(merckLongTable)
```

## Format

LongTable object

## References

TODO:: Include a reference

---

mergeAssays *Merge assays with an S4 object.*

---

## Description

Merge assays with an S4 object.

## Usage

```
mergeAssays(object, ...)
```

## Arguments

object	S4 An S4 object a list-like slot containing assays for the object.
...	Allow new arguments to be defined for this generic.

## Value

A modified version of object.

## Examples

```
"This is a generic method!"
```

---

**mergeAssays,LongTable-method**

*Endomorphically merge assays within a LongTable or inheriting class*

---

**Description**

Endomorphically merge assays within a LongTable or inheriting class

**Usage**

```
## S4 method for signature 'LongTable'  
mergeAssays(object, x, y, target = x, ..., metadata = FALSE)
```

**Arguments**

object	A LongTable or inheriting class.
x	character(1) A valid assay name in object.
y	character(1) A valid assay name in object.
target	character(1) Name of the assay to assign the result to. Can be a new or existing assay. Defaults to x.
...	Fallthrough arguments to merge.data.table to specify the join type. Use this to specify which columns to merge on. If excluded, defaults to by=assayKeys(object, y).
metadata	logical A logical vector indicating whether to attach metadata to either assay before the merge occurs. If only one value is passed that value is used for both assays. Defaults to FALSE.

**Value**

A copy of object with assays x and y merged and assigned to target.

**Author(s)**

Christopher Eeles

**See Also**

[merge.data.table](#)

---

metaConstruct	<i>Generic for preprocessing complex data before being used in the constructor of an S4 container object.</i>
---------------	---

---

## Description

This method is intended to abstract away complex constructor arguments and data preprocessing steps needed to transform raw data, such as that produced in a treatment-response or next-gen sequencing experiment, and automate building of the appropriate S4 container object. This is intended to allow mapping between different experimental designs, in the form of an S4 configuration object, and various S4 class containers in the Bioconductor community and beyond.

## Usage

```
metaConstruct(mapper, ...)

## S4 method for signature 'LongTableDataMapper'
metaConstruct(mapper)

## S4 method for signature 'TREDataMapper'
metaConstruct(mapper)
```

## Arguments

mapper	An TREDataMapper object abstracting arguments to an the TreatmentResponseExperiment constructor.
...	Allow new arguments to be defined for this generic.

## Value

An S4 object for which the class corresponds to the type of the build configuration object passed to this method.  
A LongTable object, as specified in the mapper.  
A TreatmentResponseExperiment object, as specified in the mapper.

## Examples

```
data(exampleDataMapper)
rowDataMap(exampleDataMapper) <- list(c('treatmentid'), c())
colDataMap(exampleDataMapper) <- list(c('sampleid'), c())
assayMap(exampleDataMapper) <- list(sensitivity=list(c("treatmentid", "sampleid"), c('viability')))
metadataMap(exampleDataMapper) <- list(experiment_metadata=c('metadata'))
longTable <- metaConstruct(exampleDataMapper)
longTable

data(exampleDataMapper)
exampleDataMapper <- as(exampleDataMapper, "TREDataMapper")
rowDataMap(exampleDataMapper) <- list(c('treatmentid'), c())
colDataMap(exampleDataMapper) <- list(c('sampleid'), c())
assayMap(exampleDataMapper) <- list(sensitivity=list(c("treatmentid", "sampleid"), c('viability')))
metadataMap(exampleDataMapper) <- list(experiment_metadata=c('metadata'))
tre <- metaConstruct(exampleDataMapper)
```

---

```
tre
```

---



---

```
metadata,LongTable-method
```

*Getter method for the metadata slot of a LongTable object*

---

## Description

Getter method for the metadata slot of a LongTable object

## Usage

```
## S4 method for signature 'LongTable'  
metadata(x)
```

## Arguments

x The LongTable object from which to retrieve the metadata list.

## Value

list The contents of the metadata slot of the LongTable object.

---

```
metadata<-,LongTable-method
```

*Setter method for the metadata slot of a LongTable object*

---

## Description

Setter method for the metadata slot of a LongTable object

## Usage

```
## S4 replacement method for signature 'LongTable'  
metadata(x) <- value
```

## Arguments

x LongTable The LongTable to update

value list A list of new metadata associated with a LongTable object.

## Value

LongTable A copy of the LongTable object with the value in the metadata slot.

---

mutable	<i>Remove the "immutable" S3-class from an R object, allowing it to be modified normally again.</i>
---------	---

---

## Description

Remove the "immutable" S3-class from an R object, allowing it to be modified normally again.

## Usage

```
mutable(object)
```

## Arguments

object            An R object inheriting from the "immutable" class.

## Value

The object with the "immutable" class stripped from it.

## Examples

```
immut_list <- immutable(list())
mutable(immut_list)
```

---

nci_TRE_small	<i>NCI-ALMANAC Drug Combination Data TreatmentResponseExperiment Subset</i>
---------------	---

---

## Description

This is a TreatmentResponseExperiment object containing a subset of NCI-ALMANAC drug combination screening data, with 2347 unique treatment combinations on 10 cancer cell lines selected.

## Usage

```
data(nci_TRE_small)
```

## Format

TreatmentResponseExperiment object

## References

Susan L. Holbeck, Richard Camalier, James A. Crowell, Jeevan Prasaad Govindharajulu, Melinda Hollingshead, Lawrence W. Anderson, Eric Polley, Larry Rubinstein, Apurva Srivastava, Deborah Wilsker, Jerry M. Collins, James H. Doroshow; The National Cancer Institute ALMANAC: A Comprehensive Screening Resource for the Detection of Anticancer Drug Pairs with Enhanced Therapeutic Activity. *Cancer Res* 1 July 2017; 77 (13): 3564–3576. <https://doi.org/10.1158/0008-5472.CAN-17-0489>

---

optimizeCoreGx	<i>A helper method to find the best multithreading configuration for your computer</i>
----------------	--

---

## Description

A helper method to find the best multithreading configuration for your computer

## Usage

```
optimizeCoreGx(sample_data, set = FALSE, report = !set)
```

## Arguments

sample_data	TreatmentResponseExperiment
set	logical(1) Should the function modify your R environment with the predicted optimal settings? This changes the global state of your R session!
report	logical(1) Should a data.frame of results be returned by number of threads and operation be returned? Defaults to !set.

## Value

If set=TRUE, modifies data.table threads via setDTthreads(), otherwise displays a message indicating the optimal number of threads. If report=TRUE, also returns a data.frame of the benchmark results.

## Examples

```
data(merckLongTable)
optimizeCoreGx(merckLongTable)
```

---

printSlot	<i>Helper function to print slot information</i>
-----------	--

---

## Description

Helper function to print slot information

## Usage

```
printSlot(slotName, slotData)
```

## Arguments

slotName	character The name of the slot to print.
slotData	data.table The data to print.

---

**reindex***Generic method for resetting indexing in an S4 object*

---

## Description

This method allows integer indexes used to maintain referential integrity internal to an S4 object to be reset. This is useful particularly after subsetting an object, as certain indexes may no longer be present in the object data. Reindexing removes gaps integer indexes and ensures that the smallest contiguous integer values are used in an objects indexes.

## Usage

```
reindex(object, ...)
```

## Arguments

object	S4 An object to redo indexing for
...	pairlist Allow definition of new parameters to this generic.

## Value

Depends on the implemented method

## Examples

```
print("Generics shouldn't need examples?")
```

---

**reindex,LongTable-method***Redo indexing for a LongTable object to remove any gaps in integer indexes*

---

## Description

After subsetting a LongTable, it is possible that values of rowKey or colKey could no longer be present in the object. As a result there the indexes will no longer be contiguous integers. This method will calcualte a new set of rowKey and colKey values such that integer indexes are the smallest set of contiguous integers possible for the data.

## Usage

```
## S4 method for signature 'LongTable'  
reindex(object)
```

## Arguments

object	The LongTable object to recalculalte indexes (rowKey and colKey values) for.
--------	--

**Value**

A copy of the LongTable with all keys as the smallest set of contiguous integers possible given the current data.

## rowData, LongTableDataMapper-method

*Convenience method to subset the rowData out of the rawData slot using the assigned rowDataMap metadata.*

**Description**

Convenience method to subset the rowData out of the rawData slot using the assigned rowDataMap metadata.

**Usage**

```
## S4 method for signature 'LongTableDataMapper'
rowData(x, key = TRUE)
```

**Arguments**

x	LongTableDataMapper object with valid data in the rawData and colDataMap slots.
key	logical(1) Should the table be keyed according to the id_columns of the rowDataMap slot? This will sort the table in memory. Default is TRUE.

**Value**

data.table The rowData as specified in the rowDataMap slot.

## rowData, TREDatMapper-method

*Convenience method to subset the rowData out of the rawData slot using the assigned rowDataMap metadata.*

**Description**

Convenience method to subset the rowData out of the rawData slot using the assigned rowDataMap metadata.

**Usage**

```
## S4 method for signature 'TREDatMapper'
rowData(x, key = TRUE)
```

**Arguments**

x	TREDatMapper object with valid data in the rawData and colDataMap slots.
key	logical(1) Should the table be keyed according to the id_columns of the rowDataMap slot? This will sort the table in memory. Default is TRUE.

**Value**

`data.table` The `rowData` as specified in the `rowDataMap` slot.

---

rowIDs

*Generic to access the row identifiers from*

---

**Description**

Generic to access the row identifiers from

**Usage**

`rowIDs(object, ...)`

**Arguments**

`object` S4 An object to get row id columns from.  
`...` Allow new arguments to this generic.

**Value**

Depends on the implemented method.

**Examples**

```
print("Generics shouldn't need examples?")
```

---

rowMeta

*Generic to access the row identifiers from*

---

**Description**

Generic to access the row identifiers from

**Usage**

`rowMeta(object, ...)`

**Arguments**

`object` S4 An object to get row metadata columns from.  
`...` Allow new arguments to this generic.

**Value**

Depends on the implemented method.

**Examples**

```
print("Generics shouldn't need examples?")
```

---

sensitivityInfo	<i>Generic function to get the annotations for a treatment response experiment from an S4 class</i>
-----------------	---

---

## Description

Generic function to get the annotations for a treatment response experiment from an S4 class

## Usage

```
sensitivityInfo(object, ...)
```

## Arguments

object	An S4 object to get treatment response experiment annotations from.
...	Allow new arguments to be defined for this generic.

## Value

Depends on the implemented method

## Examples

```
print("Generics shouldn't need examples?")
```

---

sensitivityInfo<-	<i>sensitivityInfo&lt;- Generic Method</i>
-------------------	--

---

## Description

Generic function to get the annotations for a treatment response experiment from an S4 class.

## Usage

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

## Arguments

object	An S4 object to set treatment response experiment annotations for.
...	Allow new arguments to be defined for this generic.
value	The new treatment response experiment annotations.

## Value

Depends on the implemented method

## Examples

```
print("Generics shouldn't need examples?")
```

---

```
sensitivityMeasures    sensitivityMeasures Generic
```

---

### Description

Get the names of the sensitivity summary metrics available in an S4 object.

### Usage

```
sensitivityMeasures(object, ...)
```

### Arguments

object	An S4 object to retrieve the names of sensitivity summary measurements for.
...	Fallthrough arguments for defining new methods

### Value

Depends on the implemented method

### Examples

```
sensitivityMeasures(clevelandSmall_cSet)
```

---

```
sensitivityMeasures<- sensitivityMeasures<- Generic
```

---

### Description

Set the names of the sensitivity summary metrics available in an S4 object.

### Usage

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

### Arguments

object	An S4 object to update.
...	Allow new methods to be defined for this generic.
value	A set of names for sensitivity measures to use to update the object with.

### Value

Depends on the implemented method

### Examples

```
print("Generics shouldn't need examples?")
```

---

`sensitivityProfiles`    *sensitivityProfiles Generic*

---

### Description

A generic for sensitivityProfiles getter method

### Usage

```
sensitivityProfiles(object, ...)
```

### Arguments

<code>object</code>	The S4 object to retrieve sensitivity profile summaries from.
<code>...</code>	<code>pairlist</code> Allow defining new arguments for this generic.

### Value

Depends on the implemented method

### Examples

```
print("Generics shouldn't need examples?")
```

---

`sensitivityProfiles<-`    *sensitivityProfiles<- Generic*

---

### Description

A generic for the sensitivityProfiles replacement method

### Usage

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

### Arguments

<code>object</code>	An S4 object to update the sensitivity profile summaries for.
<code>...</code>	Fallthrough arguments for defining new methods
<code>value</code>	An object with the new sensitivity profiles. If a matrix object is passed in, converted to data.frame before assignment

### Value

Updated CoreSet

---

`sensitivityRaw`*sensitivityRaw Generic Method*

---

**Description**

Generic function to get the raw data array for a treatment response experiment from an S4 class.

**Usage**

```
sensitivityRaw(object, ...)
```

**Arguments**

object	An S4 object to extract the raw sensitivity experiment data from.
...	pairlist Allow new parameters to be defined for this generic.

**Value**

Depends on the implemented method

**Examples**

```
print("Generics shouldn't need examples?")
```

---

`sensitivityRaw<-`*sensitivityRaw<- Generic*

---

**Description**

Generic function to set the raw data array for a treatment response experiment in an S4 class.

**Usage**

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

**Arguments**

object	An S4 object to extract the raw sensitivity data from.
...	pairlist Allow new parameters to be defined for this generic.
value	An object containing dose and viability metrics to update the object with.

**Value**

Depends on the implemented method

---

**sensitivitySlotToLongTable**  
*sensitivitySlotToLongTable Generic*

---

### Description

Convert the sensitivity slot in an object inheriting from a CoreSet from a list to a LongTable.

### Usage

```
sensitivitySlotToLongTable(object, ...)
```

### Arguments

object	CoreSet Object inheriting from CoreSet.
...	Allow new arguments to be defined on this generic.

### Value

A LongTable object containing the data in the sensitivity slot.

### Examples

```
print("Generics shouldn't need examples?")
```

---

**setOps-immutable**      *Subset an immutable object, returning another immutable object.*

---

### Description

Subset an immutable object, returning another immutable object.

### Usage

```
subset.immutable(x, ...)

## S3 method for class 'immutable'
x[...]

## S3 method for class 'immutable'
x[[...]]

## S3 method for class 'immutable'
x$...
```

### Arguments

x	An R object inheriting from the "immutable" S3-class.
...	Catch any additional parameters. Lets objects with arbitrary dimensions be made immutable.

**Value**

An immutable subset of x.

**Examples**

```
immut_mat <- immutable(matrix(1:100, 10, 10))
immut_mat[1:5, 1:5]
```

---

show,CoreSet-method *Show a CoreSet*

---

**Description**

Show a CoreSet

**Usage**

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

**Arguments**

object           CoreSet object to show via cat.

**Value**

Prints the CoreSet object to the output stream, and returns invisible NULL.

**See Also**

[cat](#)

**Examples**

```
show(clevelandSmall_cSet)
```

---

show,LongTable-method *Show method for the LongTable class*

---

**Description**

Show method for the LongTable class

**Usage**

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

**Arguments**

object            A LongTable object to print the results for.

**Value**

invisible Prints to console.

**Examples**

```
show(merckLongTable)
```

showSigAnnot

*Get the annotations for a Signature class object, as returned by drugSensitivitysig or radSensitivitySig functions available in PharmacoGx and RadioGx, respectively.*

**Description**

Get the annotations for a Signature class object, as returned by drugSensitivitysig or radSensitivitySig functions available in PharmacoGx and RadioGx, respectively.

**Usage**

```
showSigAnnot(object, ...)
```

**Arguments**

object            A Signature class object  
 ...              Allow definition of new arguments to this generic

**Value**

NULL Prints the signature annotations to console

**Examples**

```
print("Generics shouldn't need examples?")
```

---

**subset,LongTable-method**

*Subset method for a LongTable object.*

---

**Description**

Allows use of the colData and rowData data.table objects to query based on rowID and colID, which is then used to subset all assay data.tables stored in the assays slot. This function is endomorphic, it always returns a LongTable object.

**Usage**

```
## S4 method for signature 'LongTable'  
subset(x, i, j, assays = assayNames(x), reindex = TRUE)
```

**Arguments**

x	LongTable	The object to subset.
i	character, numeric, logical or call	Character: pass in a character vector of rownames for the LongTable object or a valid regex query which will be evaluated against the rownames. Numeric or Logical: vector of indices or a logical vector to subset the rows of a LongTable. Call: Accepts valid query statements to the data.table i parameter, this can be used to make complex queries using the data.table API for the rowData data.table.
j	character, numeric, logical or call	Character: pass in a character vector of colnames for the LongTable object or a valid regex query which will be evaluated against the colnames. Numeric or Logical: vector of indices or a logical vector to subset the columns of a LongTable. Call: Accepts valid query statements to the data.table i parameter, this can be used to make complex queries using the data.table API for the colData data.table.
assays	character, numeric or logical	Optional list of assay names to subset. Can be used to subset the assays list further, returning only the selected items in the new LongTable.
reindex	logical(1)	Should index values be reset such that they are the smallest possible set of consecutive integers. Modifies the "rowKey", "colKey", and all assayKey columns. Initial benchmarks indicate reindex=FALSE saves ~20% of both execution time and memory allocation. The cost of reindexing decreases the smaller your subset gets.

**Value**

LongTable A new LongTable object subset based on the specified parameters.

**Examples**

```
# Character  
subset(merckLongTable, 'ABT-888', 'CAOV3')  
# Numeric  
subset(merckLongTable, 1, c(1, 2))  
# Logical
```

```
subset(merckLongTable, , colData(merckLongTable)$sampleid == 'A2058')
# Call
subset(merckLongTable, drug1id == 'Dasatinib' & drug2id != '5-FU',
       sampleid == 'A2058')
```

**subsetTo**

*Subset a CoreSet object based on various parameters, such as cell lines, molecular features*

**Description**

Subset a CoreSet object based on various parameters, such as cell lines, molecular features

**Usage**

```
subsetTo(object, ...)
```

**Arguments**

object	An object inheriting from the CoreGx::CoreSet class
...	Allow definition of new arguments to this generic

**Value**

A subsetted version of the original object

**Examples**

```
"Generics shouldn't need examples!"
```

**summarizeMolecularProfiles**

*Summarize molecular profile data such that there is a single entry for each sample line/treatment combination*

**Description**

Summarize molecular profile data such that there is a single entry for each sample line/treatment combination

**Usage**

```
summarizeMolecularProfiles(object, ...)
```

**Arguments**

object	An S4 object to summarize the molecular profiles for.
...	Allow definition of new arguments to this generic

**Value**

Depends on the implemented method

**Examples**

```
print("Generics shouldn't need examples?")
```

---

summarizeSensitivityProfiles

*Summarize across replicates for a sensitivity dose-response experiment*

---

**Description**

Summarize across replicates for a sensitivity dose-response experiment

**Usage**

```
summarizeSensitivityProfiles(object, ...)
```

**Arguments**

object	An S4 object to summarize sensitivity profiles for.
...	Allow definition of new arguments to this generic

**Value**

Depends on the implemented method

**Examples**

```
print("Generics shouldn't need examples?")
```

---

TreatmentResponseExperiment

*TreatmentResponseExperiment constructor method*

---

**Description**

Builds a TreatmentResponseExperiment object from rectangular objects. The `rowData` argument should contain row level metadata, while the `colData` argument should contain column level metadata, for the experimental assays in the `assays` list. The `rowIDs` and `colIDs` lists are used to configure the internal keys mapping rows or columns to rows in the assays. Each list should contain at minimum one character vector, specifying which columns in `rowData` or `colData` are required to uniquely identify each row. An optional second character vector can be included, specifying any metadata columns for either dimension. These should contain information about each row but NOT be required to uniquely identify a row in the `colData` or `rowData` objects. Additional metadata can be attached to a TreatmentResponseExperiment by passing a list to the `metadata` argument.

**Usage**

```
TreatmentResponseExperiment(
  rowData,
  rowIDs,
  colData,
  colIDs,
  assays,
  assayIDs,
  metadata = list(),
  keep.rownames = FALSE
)
```

**Arguments**

rowData	<code>data.table</code> , <code>data.frame</code> , <code>matrix</code> A table like object coercible to a <code>data.table</code> containing the a unique <code>rowID</code> column which is used to key assays, as well as additional row metadata to subset on.
rowIDs	<code>character</code> , <code>integer</code> A vector specifying the names or integer indexes of the row data identifier columns. These columns will be pasted together to make up the <code>rownames</code> of the <code>TreatmentResponseExperiment</code> object.
colData	<code>data.table</code> , <code>data.frame</code> , <code>matrix</code> A table like object coercible to a <code>data.table</code> containing the a unique <code>colID</code> column which is used to key assays, as well as additional column metadata to subset on.
colIDs	<code>character</code> , <code>integer</code> A vector specifying the names or integer indexes of the column data identifier columns. These columns will be pasted together to make up the <code>colnames</code> of the <code>TreatmentResponseExperiment</code> object.
assays	A <code>list</code> containing one or more objects coercible to a <code>data.table</code> , and keyed by <code>rowIDs</code> and <code>colIDs</code> corresponding to the <code>rowID</code> and <code>colID</code> columns in <code>colData</code> and <code>rowData</code> .
assayIDs	<code>list</code> A list of <code>character</code> vectors specifying the columns needed to uniquely identify each row in an assay. Names must match the <code>assays</code> list.
metadata	A <code>list</code> of metadata associated with the <code>TreatmentResponseExperiment</code> object being constructed
keep.rownames	<code>logical</code> , <code>character</code> Logical: whether <code>rownames</code> should be added as a column if coercing to a <code>data.table</code> , default is <code>FALSE</code> . If <code>TRUE</code> , <code>rownames</code> are added to the column ' <code>rn</code> '. Character: specify a custom column name to store the <code>rownames</code> in.

**Details**

For now this class is simply a wrapper around a `LongTable` class. In the future we plan to refactor `CoreGx` such that the `LongTable` class is in a separate package. We can then specialize the implementation of `TreatmentResponseExperiment` to better capture the biomedical nature of this object.

**Value**

A `TreatmentResponseExperiment` object containing the data for a treatment response experiment configured according to the `rowIDs` and `colIDs` arguments.

---

**TreatmentResponseExperiment-class**

*TreatmentResponseExperiment class definition*

---

**Description**

Define a private constructor method to be used to build a TreatmentResponseExperiment object.

**Value**

TreatmentResponseExperiment object containing the assay data from a treatment response experiment

**Slots**

`rowData` See Slots section.

`colData` See Slots section.

`assays` See Slots section.

`metadata` See Slots section.

`.intern` See Slots section.

**Slots**

- `rowData`: A `data.table` containing the metadata associated with the row dimension of a TreatmentResponseExperiment.
- `colData`: A `data.table` containing the metadata associated with the column dimension of a TreatmentResponseExperiment.
- `assays`: A list of `data.tables`, one for each assay in a TreatmentResponseExperiment.
- `metadata`: An optional list of additional metadata for a TreatmentResponseExperiment which doesn't map to one of the dimensions.
- `.intern`: An environment that holds internal structural metadata about a TreatmentResponseExperiment object, such as which columns are required to key the object. An environment has been used to allow locking items, which can prevent accidental modification of a property required for the class to work.

---

**TREDataMapper**

*Constructor for the TREDataMapper class, which maps from one or more raw experimental data files to the slots of a LongTable object.*

---

**Description**

Constructor for the TREDataMapper class, which maps from one or more raw experimental data files to the slots of a LongTable object.

## Usage

```
TREDataMapper(
  rawdata = data.frame(),
  rawDataMap = list(character(), character()),
  colDataMap = list(character(), character()),
  assayMap = list(list(character(), character())),
  metadataMap = list(character())
)
```

## Arguments

rawdata	A <code>data.frame</code> of raw data from a treatment response experiment. This will be coerced to a <code>data.table</code> internally. We recommend using joins to aggregate your raw data if it is not present in a single file.
rowDataMap	A list-like object containing two character vectors. The first is column names in <code>rawdata</code> needed to uniquely identify each row, the second is additional columns which map to rows, but are not required to uniquely identify them. Rows should be treatments.
colDataMap	A list-like object containing two character vectors. The first is column names in <code>rawdata</code> needed to uniquely identify each column, the second is additional columns which map to rows, but are not required to uniquely identify them. Columns should be samples.
assayMap	A list-like where each item is a <code>list</code> with two character vectors defining an assay, the first containing the identifier columns in <code>rawdata</code> needed to uniquely identify each row an assay, and the second the <code>rawdata</code> columns to be mapped to that assay. The names of <code>assayMap</code> will be the names of the assays in the <code>TreatmentResponseExperiment</code> that is created when calling <code>metaConstruct</code> on this <code>DataMapper</code> object. If the character vectors have names, the value columns will be renamed accordingly.
metadataMap	A list-like where each item is a character vector of <code>rawdata</code> column names to assign to the <code>@metadata</code> of the <code>LongTable</code> , where the name of that assay is the name of the list item. If names are omitted, assays will be numbered by their index in the list.

## Details

The `guessMapping` method can be used to test hypotheses about the cardinality of one or more sets of identifier columns. This is helpful to determine the `id` columns for `rowDataMap` and `colDataMap`, as well as identify columns mapping to assays or metadata.

To attach metadata not associated with `rawdata`, please use the `metadata` assignment method on your `TREDataMapper`. This metadata will be merge with any metadata from `metadataMap` and added to the `LongTable` which this object ultimately constructs.

## Value

A `TREDataMapper` object, with columns mapped to it's slots according to the various maps in the `LongTableDataMapper` object.

## See Also

[guessMapping](#)

---

**TREDataMapper-accessors**

*Accessing and modifying data in a TREDataMapper object.*

---

**Description**

Documentation for the various setters and getters which allow manipulation of data in the slots of a TREDataMapper object.

**Usage**

```
## S4 replacement method for signature 'TREDataMapper, list'
rawdata(object) <- value

## S4 method for signature 'TREDataMapper'
rowDataMap(object)

## S4 replacement method for signature 'TREDataMapper, list_OR_List'
rowDataMap(object) <- value

## S4 method for signature 'TREDataMapper'
colDataMap(object)

## S4 replacement method for signature 'TREDataMapper, list_OR_List'
colDataMap(object) <- value

## S4 method for signature 'TREDataMapper'
assayMap(object)

## S4 replacement method for signature 'TREDataMapper, list_OR_List'
assayMap(object) <- value

## S4 method for signature 'TREDataMapper'
metadataMap(object)

## S4 replacement method for signature 'TREDataMapper, list_OR_List'
metadataMap(object) <- value
```

**Arguments**

object	A TREDataMapper object to get or set data from.
value	See details.

**Details**

**rawdata:** Get the raw data slot from a TREDataMapper object. Returns a list-like containing one or more raw data inputs to the TREDataMapper object.

**rawdata:** Set the raw data slot from a TREDataMapper object. **value:** The list-like object to set for the rawdata slot. Note: this currently only supports data.frame or data.table objects.

**rowDataMap**: list of two character vectors, the first are the columns required to uniquely identify each row of a TREDataMapper and the second any additional row-level metadata. If the character vectors have names, the resulting columns are automatically renamed to the item name of the specified column.

**rowDataMap<-**: Update the @rowDataMap slot of a TREDataMapper object, returning an invisible NULL. Arguments:

- value: A list or List where the first item is the names of the identifier columns – columns needed to uniquely identify each row in rowData – and the second item is the metadata associated with those the identifier columns, but not required to uniquely identify rows in the object rowData.

**colDataMap**: list of two character vectors, the first are the columns required to uniquely identify each row of a TREDataMapper and the second any additional col-level metadata. If the character vectors have names, the resulting columns are automatically renamed to the item name of the specified column.

**colDataMap<-**: Update the @colDataMap slot of a TREDataMapper object, returning an invisible NULL. Arguments:

- value: A list or List where the first item is the names of the identifier columns – columns needed to uniquely identify each row in colData – and the second item is the metadata associated with those the identifier columns, but not required to uniquely identify rows in the object rowData.

**assayMap**: A list of character vectors. The name of each list item will be the assay in a LongTableDataMapper object that the columns in the character vector will be assigned to. Column renaming occurs automatically when the character vectors have names (from the value to the name).

**assayMap<-**: Updates the @assayMap slot of a TREDataMapper object, returning an invisible NULL. Arguments:

- value: A list of character vectors, where the name of each list item is the name of an assay and the values of each character vector specify the columns mapping to the assay in the S4 object the TREDataMapper constructs.

**metadataMap**: A list of character vectors. Each item is an element of the constructed objects @metadata slot.

**metadataMap<-**: Updates TREDataMapper object in-place, then returns an invisible(NULL). Arguments:

- value: A list of character vectors. The name of each list item is the name of the item in the @metadata slot of the TREDataMapper object created when metaConstruct is called on the DataMapper, and a character vector specifies the columns of @rawdata to assign to each item.

## Value

Accessors: See details

Setters: An update TREDataMapper object, returned invisibly.

## See Also

Other DataMapper-accessors: [DataMapper-accessors](#), [LongTableDataMapper-accessors](#)

## Examples

```

rowDataMap(exampleDataMapper)

rowDataMap(exampleDataMapper) <- list(c('treatmentid'), c())

colDataMap(exampleDataMapper)

colDataMap(exampleDataMapper) <- list(c('sampleid'), c())

assayMap(exampleDataMapper)

assayMap(exampleDataMapper) <- list(sensitivity=c(viability1='viability'))

metadataMap(exampleDataMapper)

metadataMap(exampleDataMapper) <- list(object_metadata=c('metadata'))

```

---

TREDataMapper-class    *A Class for Mapping Between Raw Data and an TreatmentResponseExperiment Object*

---

## Description

A Class for Mapping Between Raw Data and an TreatmentResponseExperiment Object

## Slots

**rawdata** See Slots section.  
**rowDataMap** See Slots section.  
**colDataMap** See Slots section.  
**assayMap** See Slots section.  
**metadataMap** See Slots section.

## Slots

- **rowDataMap**: A list-like object containing two character vectors. The first is column names in `rawdata` needed to uniquely identify each row, the second is additional columns which map to rows, but are not required to uniquely identify them. Rows should be drugs.
- **colDataMap**: A list-like object containing two character vectors. The first is column names in `rawdata` needed to uniquely identify each column, the second is additional columns which map to rows, but are not required to uniquely identify them. Columns should be samples.
- **assayMap**: A list-like where each item is a list with two elements specifying an assay, the first being the identifier columns in `rawdata` needed to uniquely identify each row an assay, and the second a list of `rawdata` columns to be mapped to that assay. The names of `assayMap` will be the names of the assays in the `LongTable` that is created when calling `metaConstruct` on this `DataMapper` object.
- **metadataMap**: A list-like where each item is a character vector of `rawdata` column names to assign to the `@metadata` of the `LongTable`, where the name of that assay is the name of the list item. If names are omitted, assays will be numbered by their index in the list.

- `rawdata`: A list-like object containing one or more pieces of raw data that will be processed and mapped to the slots of an S4 object.
- `metadata`: A List of object level metadata.

## updateObject,CoreSet-method

*Update the CoreSet class after changes in it struture or API***Description**

Update the CoreSet class after changes in it struture or API

**Usage**

```
## S4 method for signature 'CoreSet'
updateObject(object, verify = FALSE, verbose = FALSE)
```

**Arguments**

<code>object</code>	A CoreSet object to update the class structure for.
<code>verify</code>	A logical(1) indicating is validObject should be called after updating the object. Defaults to TRUE, only set FALSE for debugging.
<code>verbose</code>	TRUE or FALSE, indicating whether information about the update should be reported

**Value**

CoreSet with update class structure.

## updateObject,LongTable-method

*Update the LongTable class after changes in it struture or API***Description**

Update the LongTable class after changes in it struture or API

**Usage**

```
## S4 method for signature 'LongTable'
updateObject(object, verify = FALSE, verbose = FALSE)
```

**Arguments**

<code>object</code>	A LongTable object to update the class structure for.
<code>verify</code>	A logical(1) indicating is validObject should be called after updating the object. Defaults to TRUE, only set FALSE for debugging.
<code>verbose</code>	TRUE or FALSE, indicating whether information about the update should be reported

**Value**

LongTable with update class structure.

---

updateSampleId	<i>Update the sample ids in a cSet object</i>
----------------	---

---

**Description**

Update the sample ids in a cSet object

**Usage**

```
updateSampleId(object, new.ids = vector("character"))
```

**Arguments**

object	The object for which the sample ids will be updated
new.ids	The new ids to assign to the object

**Value**

CoreSet The modified CoreSet object

**Examples**

```
updateSampleId(clevelandSmall_cSet, sampleNames(clevelandSmall_cSet))
```

---

updateTreatmentId	<i>Update the treatment ids in a cSet object</i>
-------------------	--

---

**Description**

Update the treatment ids in a cSet object

**Usage**

```
updateTreatmentId(object, new.ids = vector("character"))
```

**Arguments**

object	The object for which the treatment ids will be updated
new.ids	The new ids to assign to the object

**Value**

CoreSet The modified CoreSet object

**Examples**

```
updateTreatmentId(clevelandSmall_cSet, treatmentNames(clevelandSmall_cSet))
```

---

[,LongTable,ANY,ANY,ANY-method  
*[ LongTable Method*

---

## Description

Single bracket subsetting for a LongTable object. See subset for more details.

## Usage

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

x[i, j, assays = assayNames(x), ..., drop = FALSE]
```

## Arguments

x	LongTable The object to subset.
i	character, numeric, logical or call Character: pass in a character vector of drug names, which will subset the object on all row id columns matching the vector. This parameter also supports valid R regex query strings which will match on the colnames of x. For convenience, * is converted to .* automatically. Colon can be to denote a specific part of the colnames string to query. Numeric or Logical: these select based on the rowKey from the <code>rowData</code> method for the LongTable. Call: Accepts valid query statements to the <code>data.table</code> i parameter as a call object. We have provided the function <code>.()</code> to conveniently convert raw R statements into a call for use in this function.
j	character, numeric, logical or call Character: pass in a character vector of drug names, which will subset the object on all drug id columns matching the vector. This parameter also supports regex queries. Colon can be to denote a specific part of the colnames string to query. Numeric or Logical: these select based on the rowID from the <code>rowData</code> method for the LongTable. Call: Accepts valid query statements to the <code>data.table</code> i parameter as a call object. We have provided the function <code>.()</code> to conveniently convert raw R statements into a call for use in this function.
assays	character Names of assays which should be kept in the LongTable after subsetting.
...	Included to ensure drop can only be set by name.
drop	logical Included for compatibility with the '[' primitive, it defaults to FALSE and changing it does nothing.

## Details

This function is endomorphic, it always returns a LongTable object.

## Value

A LongTable containing only the data specified in the function parameters.

## Examples

```
# Character
merckLongTable['ABT-888', 'CAOV3']
# Numeric
merckLongTable[1, c(1, 2)]
# Logical
merckLongTable[, colData(merckLongTable)$sampleid == 'A2058']
# Call
merckLongTable[
  .(drug1id == 'Dasatinib' & drug2id != '5-FU'),
  .(sampleid == 'A2058'),
]
```

---

```
[[<,LongTable,ANY,ANY-method
  [[<- Method for LongTable Class
```

---

## Description

Just a wrapper around `assay<-` for convenience. See `?'assay<-,LongTable,character-method'`.

## Usage

```
## S4 replacement method for signature 'LongTable,ANY,ANY'
x[[i]] <- value
```

## Arguments

<code>x</code>	A <code>LongTable</code> to update.
<code>i</code>	The name of the assay to update, must be in <code>assayNames(object)</code> .
<code>value</code>	A <code>data.frame</code>

## Value

A `LongTable` object with the assay `i` updated using `value`.

## Examples

```
merckLongTable[['sensitivity']] <- merckLongTable[['sensitivity']]
```

---

`$,LongTable-method`     *Select an assay from a LongTable object*

---

### Description

Select an assay from a LongTable object

### Usage

```
## S4 method for signature 'LongTable'
x$name
```

### Arguments

<code>x</code>	A LongTable object to retrieve an assay from
<code>name</code>	character The name of the assay to get.

### Value

`data.frame` The assay object.

### Examples

```
merckLongTable$sensitivity
```

---

`$<-,LongTable-method`     *Update an assay from a LongTable object*

---

### Description

Update an assay from a LongTable object

### Usage

```
## S4 replacement method for signature 'LongTable'
x$name <- value
```

### Arguments

<code>x</code>	A LongTable to update an assay for.
<code>name</code>	character(1) The name of the assay to update
<code>value</code>	A <code>data.frame</code> or <code>data.table</code> to update the assay with.

### Value

Updates the assay name in `x` with `value`, returning an invisible `NULL`.

### Examples

```
merckLongTable$sensitivity <- merckLongTable$sensitivity
```

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