

# Package ‘sigaR’

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**Type** Package

**Title** Statistics for Integrative Genomics Analyses in R

**Version** 1.28.0

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**Description** Facilitates the joint analysis of high-throughput data from multiple molecular levels. Contains functions for manipulation of objects, various analysis types, and some visualization.

**License** GPL (>= 2)

**LazyLoad** yes

**URL** <http://www.few.vu.nl/~wvanwie>

**Depends** Biobase, CGHbase, methods, mvtnorm,

**Imports** corpcor (>= 1.6.2), graphics, igraph, limma, marray, MASS, penalized, quadprog, snowfall, stats

**Suggests** CGHcall

**biocViews** Microarray, DifferentialExpression, aCGH, GeneExpression, Pathways

**NeedsCompilation** no

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sigar-package

*statistics for integrative genomics analyses in R*

---

### Description

The package facilitates several types of integrative analysis of high-throughput data from various molecular levels. In addition, it includes functions for data management and visualization.

### Details

Package: sigar  
Type: Package  
Version: 1.0  
Date: 2011-04-15  
License: What license is it under?  
LazyLoad: yes

### Author(s)

Author: Wessel N. van Wieringen Maintainer: Wessel N. van Wieringen <w.vanwieringen@vumc.nl>

### References

- Van Wieringen, W.N., Van de Wiel, M.A. (2009), "Non-parametric testing for DNA copy number induced differential mRNA gene expression", *Biometrics*, 65(1), 19-29.
- Van Wieringen, W.N., Berkhof, J., Van de Wiel, M.A. (2010), "A random coefficients model for regional co-expression associated with DNA copy number", *Statistical Applications in Genetics and Molecular Biology*, Volume 9, Issue1, Article 25, 1-28.
- Van Wieringen, W.N., Van der Vaart, A.W. (2011), "Statistical analysis of the cancer cell's molecular entropy using high-throughput data", *Bioinformatics*, 27(4), 556-563.
- Van Wieringen, W.N., Unger, K., Leday, G.G.R., Krijgsman, O., De Menezes, R.X., Ylstra, B., Van de Wiel, M.A. (2012), "Matching of array CGH and gene expression microarray features for the purpose of integrative analysis", *submitted for publication*.

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.RCMloss-method

*Internal function*

---

### Description

Internal function.

### Note

Not to be called by the user.

---

`cghCall2cghSeg`*Genomic ordering of cghSeg-objects.*

---

**Description**

Transforms a `cghCall`-object to a `cghSeg`-object, by removing the slots present in the former but not in the latter.

**Usage**

```
cghCall2cghSeg(CNdata, verbose=TRUE)
```

**Arguments**

<code>CNdata</code>	Object of class <code>cghCall</code> .
<code>verbose</code>	Logical indicator: should intermediate output be printed on the screen?

**Value**

Object of class `cghSeg`.

**Author(s)**

Wessel N. van Wieringen: <w.vanwieringen@vumc.nl>

**References**

Van de Wiel, M.A., Kim, K.I., Vosse, S.J., Van Wieringen, W.N., Wilting, S.M., Ylstra, B. (2007), "CGHcall: an algorithm for calling aberrations for multiple array CGH tumor profiles", *Bioinformatics*, 23, 892-894.

**See Also**

`cghCall`, `cghSeg`.

**Examples**

```
# load data
data(pollackCN16)

# reduce the cghCall-object to a cghSeg-object
pollackCN16seg <- cghCall2cghSeg(pollackCN16)
```

---

cghCall2maximumSubset *Maximum subsetting cghCall-objects.*

---

### Description

Limit an `cghCall` object to a subset of its features, selecting those features with the most deviating copy number signal.

### Usage

```
cghCall2maximumSubset(CNdata, featuresAndWeights, chr, bpstart,  
bpend, ncpus = 1, verbose=TRUE)
```

### Arguments

CNdata	Object of class <code>cghCall</code> .
featuresAndWeights	Object of class <code>list</code> . Each list item is a matrix. The first column of this matrix contains the row numbers of features to be maintained in the <code>cghCall</code> -object. The second column contains the weights of each features, to be used in the calculation of the weighted average copy number signal.
chr	Column in the slot <code>featureData</code> of the <code>cghCall</code> -object specifying the chromosome information of the features.
bpstart	Column in the slot <code>featureData</code> of the <code>cghCall</code> -object specifying the start basepair information of the features.
bpend	Column in the slot <code>featureData</code> of the <code>cghCall</code> -object specifying the end basepair information of the features.
ncpus	Number of cpus to be used in computations.
verbose	Logical indicator: should intermediate output be printed on the screen?

### Details

Per entry of the `featuresAndWeights`-object and per sample the feature with the maximum absolute segmented DNA copy number signal is selected.

### Value

Object of class `cghCall`, restricted to the specified subset of features.

### Author(s)

Wessel N. van Wieringen: <w.vanwieringen@vumc.nl>

### References

Van Wieringen, W.N., Unger, K., Leday, G.G.R., Krijgsman, O., De Menezes, R.X., Ylstra, B., Van de Wiel, M.A. (2012), "Matching of array CGH and gene expression microarray features for the purpose of integrative analysis", *BMC Bioinformatics*, 13:80.

**See Also**[matchAnn2Ann](#)**Examples**

```
# load data
data(pollackCN16)

# extract genomic information from ExpressionSet-object
chr <- fData(pollackCN16)[,1]
bpstart <- fData(pollackCN16)[,2]
bpend <- fData(pollackCN16)[,3]

# find unique genomic locations
uniqInfo <- uniqGenomicInfo(chr, bpstart, bpend, verbose = FALSE)

# subset cghCall-object to features with unique genomic locations
pollackCN16 <- cghCall2maximumSubset(pollackCN16, uniqInfo, 1, 2, 3)
```

---

`cghCall2order`*Genomic ordering of cghCall-objects.*

---

**Description**

Orders the features within a `cghCall`-object in accordance with their genomic order.

**Usage**

```
cghCall2order(CNdata, chr, bpstart, verbose=TRUE)
```

**Arguments**

<code>CNdata</code>	Object of class <code>cghCall</code> .
<code>chr</code>	Column in the slot <code>featureData</code> of the <code>cghCall</code> -object specifying the chromosome information of the features.
<code>bpstart</code>	Column in the slot <code>featureData</code> of the <code>cghCall</code> -object specifying the start basepair information of the features.
<code>verbose</code>	Logical indicator: should intermediate output be printed on the screen?

**Value**

Object of class `cghCall`, now genomically ordered.

**Author(s)**

Wessel N. van Wieringen: <w.vanwieringen@vumc.nl>

**References**

Van de Wiel, M.A., Kim, K.I., Vosse, S.J., Van Wieringen, W.N., Wilting, S.M., Ylstra, B. (2007), "CGHcall: an algorithm for calling aberrations for multiple array CGH tumor profiles", *Bioinformatics*, 23, 892-894.

**See Also**

[cghCall](#).

**Examples**

```
# load data
data(pollackCN16)

# order the copy number data genomically
pollackCN16 <- cghCall2order(pollackCN16, 1, 2)
```

---

cghCall2subset	<i>Subsetting cghCall-objects.</i>
----------------	------------------------------------

---

**Description**

Limit an [cghCall](#) object to a subset of its features.

**Usage**

```
cghCall2subset(CNdata, featureSubset, verbose=TRUE)
```

**Arguments**

CNdata	Object of class <a href="#">cghCall</a> .
featureSubset	Object of class <code>numeric</code> , containing the row numbers of features to be maintained in the <a href="#">cghCall</a> -object.
verbose	Logical indicator: should intermediate output be printed on the screen?

**Value**

Object of class [cghCall](#), restricted to the specified subset of features.

**Author(s)**

Wessel N. van Wieringen: <w.vanwieringen@vumc.nl>

**References**

Van de Wiel, M.A., Kim, K.I., Vosse, S.J., Van Wieringen, W.N., Wilting, S.M., Ylstra, B. (2007), "CGHcall: an algorithm for calling aberrations for multiple array CGH tumor profiles", *Bioinformatics*, 23, 892-894.

**See Also**

[cghCall](#).

**Examples**

```
# load data
data(pollackCN16)

# order the copy number data genomically
pollackCN16 <- cghCall2subset(pollackCN16, c(1:50))
```

---

cghCall2weightedSubset

*Weighted subsetting cghCall-objects.*

---

### Description

Limit an `cghCall` object to a subset of its features, using weighted averaging of the copy number signal.

### Usage

```
cghCall2weightedSubset(CNdata, featuresAndWeights, chr, bpstart,  
bpend, ncpus = 1, verbose=TRUE)
```

### Arguments

CNdata	Object of class <code>cghCall</code> .
featuresAndWeights	Object of class <code>list</code> . Each list item is a matrix. The first column of this matrix contains the row numbers of features to be maintained in the <code>cghCall</code> -object. The second column contains the weights of each features, to be used in the calculation of the weighted average copy number signal.
chr	Column in the slot <code>featureData</code> of the <code>cghCall</code> -object specifying the chromosome information of the features.
bpstart	Column in the slot <code>featureData</code> of the <code>cghCall</code> -object specifying the start basepair information of the features.
bpend	Column in the slot <code>featureData</code> of the <code>cghCall</code> -object specifying the end basepair information of the features.
ncpus	Number of cpus to be used in computations.
verbose	Logical indicator: should intermediate output be printed on the screen?

### Value

Object of class `cghCall`, restricted to the specified subset of features.

### Warning

The `phenoData`, `experimentData`, and other slots of the `cghCall`-object are currently not passed on to the subsetted object.

### Note

This is a more intricate version of the `cghCall2subset` function. They exist parallel because this function is (much) slower than its counterpart.

### Author(s)

Wessel N. van Wieringen: <w.vanwieringen@vumc.nl>



## References

Van Wieringen, W.N., Unger, K., Leday, G.G.R., Krijgsman, O., De Menezes, R.X., Ylstra, B., Van de Wiel, M.A. (2012), "Matching of array CGH and gene expression microarray features for the purpose of integrative analysis", *BMC Bioinformatics*, 13:80.

## See Also

cghCall2subset

## Examples

```
# load data
data(pollackCN16)

# extract genomic information from ExpressionSet-object
chr <- fData(pollackCN16)[,1]
bpstart <- fData(pollackCN16)[,2]
bpend <- fData(pollackCN16)[,3]

# find unique genomic locations
uniqInfo <- uniqGenomicInfo(chr, bpstart, bpend, verbose = FALSE)

# subset cghCall-object to features with unique genomic locations
pollackCN16 <- cghCall2weightedSubset(pollackCN16, uniqInfo, 1, 2, 3)
```

---

cghSeg2order

*Genomic ordering of cghSeg-objects.*

---

## Description

Orders the features within a `cghSeg`-object in accordance with their genomic order.

## Usage

```
cghSeg2order(CNdata, chr, bpstart, verbose=TRUE)
```

## Arguments

CNdata	Object of class <code>cghSeg</code> .
chr	Column in the slot <code>featureData</code> of the <code>cghSeg</code> -object specifying the chromosome information of the features.
bpstart	Column in the slot <code>featureData</code> of the <code>cghSeg</code> -object specifying the start base-pair information of the features.
verbose	Logical indicator: should intermediate output be printed on the screen?

## Value

Object of class `cghSeg`, now genomically ordered.

## Author(s)

Wessel N. van Wieringen: <w.vanwieringen@vumc.nl>

## References

Van de Wiel, M.A., Kim, K.I., Vosse, S.J., Van Wieringen, W.N., Wilting, S.M. , Ylstra, B. (2007), "CGHcall: an algorithm for calling aberrations for multiple array CGH tumor profiles", *Bioinformatics*, 23, 892-894.

## See Also

[cghSeg](#).

## Examples

```
# load data
data(pollackCN16)

# transform the cghCall-object to a cghSeg-object
pollackCN16 <- cghCall2cghSeg(pollackCN16)

# order the copy number data genomically
pollackCN16 <- cghSeg2order(pollackCN16, 1, 2)
```

---

cghSeg2subset	<i>Subsetting cghSeg-objects.</i>
---------------	-----------------------------------

---

## Description

Limit an [cghSeg](#) object to a subset of its features.

## Usage

```
cghSeg2subset(CNdata, featureSubset, verbose=TRUE)
```

## Arguments

CNdata	Object of class <a href="#">cghSeg</a> .
featureSubset	Object of class <code>numeric</code> , containing the row numbers of features to be maintained in the <a href="#">cghSeg</a> -object.
verbose	Logical indicator: should intermediate output be printed on the screen?

## Value

Object of class [cghSeg](#), restricted to the specified subset of features.

## Author(s)

Wessel N. van Wieringen: <w.vanwieringen@vumc.nl>

## References

Van de Wiel, M.A., Kim, K.I., Vosse, S.J., Van Wieringen, W.N., Wilting, S.M. , Ylstra, B. (2007), "CGHcall: an algorithm for calling aberrations for multiple array CGH tumor profiles", *Bioinformatics*, 23, 892-894.

**See Also**

[cghSeg](#).

**Examples**

```
# load data
data(pollackCN16)

# transform the cghCall-object to a cghSeg-object
pollackCN16 <- cghCall2cghSeg(pollackCN16)

# subset the copy number data
pollackCN16 <- cghSeg2subset(pollackCN16, c(1:50))
```

---

cghSeg2weightedSubset *Weighted subsetting cghSeg-objects.*

---

**Description**

Limit an [cghSeg](#) object to a subset of its features, using weighted averaging of the copy number signal.

**Usage**

```
cghSeg2weightedSubset(CNdata, featuresAndWeights, chr, bpstart,
  bpend, ncpus = 1, verbose=TRUE)
```

**Arguments**

CNdata	Object of class <a href="#">cghSeg</a> .
featuresAndWeights	Object of class <code>list</code> . Each list item is a <code>matrix</code> . The first column of this <code>matrix</code> contains the row numbers of features to be maintained in the <a href="#">cghSeg</a> -object. The second column contains the weights of each features, to be used in the calculation of the weighted average copy number signal.
chr	Column in the slot <code>featureData</code> of the <a href="#">cghSeg</a> -object specifying the chromosome information of the features.
bpstart	Column in the slot <code>featureData</code> of the <a href="#">cghSeg</a> -object specifying the start base-pair information of the features.
bpend	Column in the slot <code>featureData</code> of the <a href="#">cghSeg</a> -object specifying the end base-pair information of the features.
ncpus	Number of <code>cpus</code> to be used in computations.
verbose	Logical indicator: should intermediate output be printed on the screen?

**Value**

Object of class [cghSeg](#), restricted to the specified subset of features.

**Warning**

The phenoData, experimentData, and other slots of the cghSeg-object are currently not passed on to the subsetted object.

**Note**

This is a more intricate version of the cghSeg2subset function. They exists parallel because this function is (much) slower than its counterpart.

**Author(s)**

Wessel N. van Wieringen: <w.vanwieringen@vumc.nl>

**References**

Van Wieringen, W.N., Unger, K., Leday, G.G.R., Krijgsman, O., De Menezes, R.X., Ylstra, B., Van de Wiel, M.A. (2012), "Matching of array CGH and gene expression microarray features for the purpose of integrative analysis", *BMC Bioinformatics*, 13:80.

**See Also**

cghSeg2subset

**Examples**

```
# load data
data(pollackCN16)

# extract genomic information from ExpressionSet-object
chr <- fData(pollackCN16)[,1]
bpstart <- fData(pollackCN16)[,2]
bpend <- fData(pollackCN16)[,3]

# find unique genomic locations
uniqInfo <- uniqGenomicInfo(chr, bpstart, bpend, verbose = FALSE)

# transform the cghCall-object to a cghSeg-object
pollackCN16 <- cghCall2cghSeg(pollackCN16)

# subset cghSeg-object to features with unique genomic locations
pollackCN16 <- cghSeg2weightedSubset(pollackCN16, uniqInfo, 1, 2, 3)
```

---

cisEffectPlot

*DNA-mRNA plot*

---

**Description**

A variant on the boxplot, plotting the gene expression against the DNA copy number data. For each individual an open blue circle per call is plotted, all with their centerpoint at the height of the individual's expression level. The radius of the circles is proportional to the corresponding call probabilities. Call probabilities equal to zero reduce circles to dots. The red filled circles have a radius proportional to the estimated expected call probabilities, with their centerpoints at the estimated mean expression for the respective call.

**Usage**

```
cisEffectPlot(geneId, CNdata, GEdata, verbose=FALSE)
```

**Arguments**

geneId	Indicator of the gene to be plotted. Indicator refers to the row in the <a href="#">ExpressionSet</a> -object.
CNdata	Object of class <code>cghCall</code> , containing (among others) annotation and call probabilities. Features should be matched with those of the accompanying <code>ExpressionSet</code> -object (as may be done using the <code>matchAnn2Ann</code> -function).
GEdata	Object of class <code>ExpressionSet</code> . Features should be matched with those of the accompanying <code>cghCall</code> -object (as may be done using the <code>matchAnn2Ann</code> -function).
verbose	Logical indicator: should intermediate output be printed on the screen?

**Note**

This function is a rewritten version of the `intCNGEan.plot` function of the `intCNGEan`-package.

**Author(s)**

Wessel N. van Wieringen: <w.vanwieringen@vumc.nl>

**References**

Van Wieringen, W.N., Van de Wiel, M.A. (2009), "Non-parametric testing for DNA copy number induced differential mRNA gene expression", *Biometrics*, 65(1), 19-29.

**See Also**

`boxplot`, `cisEffectTune`, `cisEffectTest`, `matchAnn2Ann`

**Examples**

```
# load data
data(pollackCN16)
data(pollackGE16)

# plot DNA copy number vs. gene expression.
cisEffectPlot(225, pollackCN16, pollackGE16)
```

---

<code>cisEffectTable</code>	<i>Table of cis-effect test results</i>
-----------------------------	---

---

**Description**

Function to display the results of `cisEffectTest`-function in a table-format. Table may be restricted to a specified number of genes and sorted by relevant statistics.

**Usage**

```
cisEffectTable(testRes, number=10, sort.by=NULL)
```

**Arguments**

testRes	Object of class cisTest as produced by the <code>cisEffectTest</code> -function.
number	Number of genes whose results are to be included in the table.
sort.by	character indicating how the table is to sorted: NULL no sorting (genomic order), p.value, R2 or effect sort the table by the corresponding statistic.

**Author(s)**

Wessel N. van Wieringen: <w.vanwieringen@vumc.nl>

**References**

Van Wieringen, W.N., Van de Wiel, M.A. (2009), "Non-parametric testing for DNA copy number induced differential mRNA gene expression", *Biometrics*, 65(1), 19-29.

**See Also**

`cisEffectTest`

**Examples**

```
# load data
data(pollackCN16)
data(pollackGE16)

# test cis-effect of DNA copy number on gene expression levels
cisRes <- cisEffectTest(pollackCN16, pollackGE16, 1:nrow(pollackGE16), 1, nPerm=25)

# display top results
cisEffectTable(cisRes, number=10, sort.by="R2")
```

---

<code>cisEffectTest</code>	<i>Nonparametric testing for copy number induced differential gene expression.</i>
----------------------------	--

---

**Description**

A nonparametric test for the detection of copy number induced differential gene expression. The test incorporates the uncertainty of the calling of genomic aberrations: weighted version of well-known test statistics are used. An efficient permutation re-sampling procedure is used for p-value calculation. The test statistics may be "shrunk" to borrow information across neighboring genes that share the same copy number signature.

**Usage**

```
cisEffectTest(CNdata, GEdata, genes2test=NULL, GEchr,
analysisType="univariate", testStatistic="wcvm",
nPerm = 10000, lowCiThres = 0.1, verbose=TRUE)
```

**Arguments**

CNdata	Object of class <code>cghCall</code> , containing (among others) annotation and call probabilities. Features should be matched with those of the accompanying <code>ExpressionSet</code> -object (as may be done using the <code>matchAnn2Ann</code> -function).
GEdata	Object of class <code>ExpressionSet</code> . Features should be matched with those of the accompanying <code>cghCall</code> -object (as may be done using the <code>matchAnn2Ann</code> -function).
genes2test	Numeric indicator vector containing row number of genes for which the DNA copy number cis-effect should be tested. The function <code>cisEffectTune</code> yields an optimal selection.
GEchr	Column in the slot <code>featureData</code> of the <code>ExpressionSet</code> -object <code>GEdata</code> specifying the chromosome information of the features.
analysisType	Indicator to determine whether the test statistic should be "shrunk" within a region. Either "univariate" (no shrinkage) or "regional" (shrinkage).
testStatistic	Test statistic to be used, either "wcvm" or "wmw", the weighted Cramer-Von Mises and the weighted Mann-Whitney test statistic, respectively.
nPerm	Number of permutations used for the p-value calculation.
lowCiThres	A value between 0 and 1. Determines speed of efficient p-value calculation. Genes with a probability smaller than 0.001 of a p-value smaller than <code>eff.p.val.thres</code> are discarded from the permutation analysis and their p-value is set equal to 1. Should be chosen in accordance with the FDR-threshold for significance.
verbose	Logical indicator: should intermediate output be printed on the screen?

**Value**

Object of class `cisTest`.

**Note**

This function is a rewritten version of the `intCNGEan.test` function of the `intCNGEan`-package.

**Author(s)**

Wessel N. van Wieringen: <w.vanwieringen@vumc.nl>

**References**

Van Wieringen, W.N., Van de Wiel, M.A. (2009), "Non-parametric testing for DNA copy number induced differential mRNA gene expression", *Biometrics*, 65(1), 19-29.

**See Also**

`matchAnn2Ann`, `cisEffectTune`, `cisEffectTable`, `cisEffectPlot`

**Examples**

```
# load data
data(pollackCN16)
data(pollackGE16)

# test cis-effect of DNA copy number on gene expression levels
cisRes <- cisEffectTest(pollackCN16, pollackGE16, 1:nrow(pollackGE16), 1, nPerm=25)
```

---

cisEffectTune                      *Pre-test and tuning.*

---

### Description

Decides which test to perform: loss vs. no-loss (tumor suppressor) or no-gain vs gain (proto-onco). Followed by a tuning algorithm that enhances the overall power of the FDR procedure by excluding genes with either unbalanced (many samples having a high call probability of, say, a loss) or imprecise (many call probabilities close to 0.5) soft calls, which is likely to increase the probability of detection for genes with a more favorable call probability distribution.

### Usage

```
cisEffectTune(CNdata, GEdata, testStatistic, nGenes=250,
nPerm=250, minCallProbMass=0.10, verbose=TRUE)
```

### Arguments

CNdata	Object of class <code>cghCall</code> , containing (among others) annotation and call probabilities. Features should be matched with those of the accompanying <code>ExpressionSet</code> -object (as may be done using the <code>matchAnn2Ann</code> -function).
GEdata	Object of class <code>ExpressionSet</code> . Features should be matched with those of the accompanying <code>cghCall</code> -object (as may be done using the <code>matchAnn2Ann</code> -function).
testStatistic	Test statistic to be used, either "wcvm" or "wmw", the weighted Cramer-Von Mises and the weighted Mann-Whitney test statistic, respectively.
nGenes	Number of genes used for tuning.
nPerm	Number of permutation used for tuning.
minCallProbMass	A number inbetween 0 and 1. Genes with a marginal call probabilities in one of the classes smaller than <code>minCallProbMass</code> are discarded from further analysis. Effectively, this ensures identifiability of copy number effect on expression.
verbose	Boolean to suppress output, either FALSE and TRUE.

### Value

A numeric-object with the genes selected for testing. Numbering corresponds to genes of the pre-tuned, but matched data set.

### Note

This function is a rewritten version of the `intCNGEan.tune` function of the `intCNGEan`-package.

### Author(s)

Wessel N. van Wieringen: <w.vanwieringen@vumc.nl>

### References

Van Wieringen, W.N., Van de Wiel, M.A. (2009), "Non-parametric testing for DNA copy number induced differential mRNA gene expression", *Biometrics*, 65(1), 19-29.



**See Also**

matchAnn2Ann, cisEffectTest

**Examples**

```
# load data
data(pollackCN16)
data(pollackGE16)

# select genes that are likely to have a significant genomic cis-effect on expression levels
genes2test <- cisEffectTune(pollackCN16, pollackGE16, "wmw", nGenes=50, nPerm=50)
```

---

cisTest-class	<i>Class "cisTest" for storing the results of the function cisEffectTest.</i>
---------------	---

---

**Description**

The class cisTest is the output of a call to `cisEffectTest`. It stores results from a hypothesis test.

**Slots**

**geneInfo:** Object of class "data.frame". E.g., annotation information of genes.

**geneId:** Object of class "numeric". Row number in `ExpressionSet`-object used in `{cisEffectTest}`, corresponding to a gene.

**comparison:** Object of class "numeric". Indicator of test performed, either "1" (loss vs. no-loss) or "2" (no-gain vs. gain).

**av.prob1:** Object of class "numeric". The estimated marginal call probability.

**av.prob2:** Object of class "numeric". The estimated marginal call probability.

**effectSize:** Object of class "numeric". Estimated genomic cis-effect on gene expression.

**R2:** Object of class "numeric". Percentage of explained variation in expression levels by .

**regId:** Object of class "numeric". Indicator for the region (NULL in the regional-analysis).

**beginReg:** Object of class "numeric". Row number in `ExpressionSet`-object corresponding to the first gene of the region (NULL in the regional-analysis).

**endReg:** Object of class "numeric". Row number in `ExpressionSet`-object corresponding to the last gene of the region (NULL in the regional-analysis).

**shrinkage:** Object of class "numeric". Amount of shrinkage applied in the regional analysis (NULL in the regional-analysis).

**p.value:** Object of class "numeric". P-value for the non-parametric test of the genomic cis-effect on expression levels.

**adjP.value:** Object of class "numeric". BH-multiple testing correct p-values.

**analysisType** Indicator whether the test statistic has been "shrunk" within a region. Either "univariate" (no shrinkage) or "regional" (shrinkage).

**testStatistic** Test statistic used, either "wcvm" or "wmw", the weighted Cramer-Von Mises and the weighted Mann-Whitney test statistic, respectively.

**nPerm** Number of permutations used for the p-value calculation.

**Methods**

**cisEffectTable** signature(object = "cisTest"): Prints the test results.

**Author(s)**

Wessel van Wieringen: <w.vanwieringen@vumc.nl>

**See Also**

[cisEffectTest](#)

**Examples**

```
showClass("cisTest")
```

---

CNGEheatmaps

*Parallel CN and GE heatmap plotting*

---

**Description**

Heatmaps of DNA copy number and gene expression data are plotted together.

**Usage**

```
CNGEheatmaps(CNdata, GEdata, location = "mode", colorbreaks = "equiquantiles")
```

**Arguments**

CNdata	Object of class <code>cghCall</code> , containing (among others) annotation and call probabilities. Features should be matched with those of the accompanying <code>ExpressionSet</code> -object (as may be done using the <code>matchCGHcall2ExpressionSet</code> -function).
GEdata	Object of class <code>ExpressionSet</code> . Features should be matched with those of the accompanying <code>cghCall</code> -object (as may be done using the <code>matchCGHcall2ExpressionSet</code> -function).
location	Parameter (median, mean, or mode) specifying how the center of the gene expression heatmap color-scheme is determined.
colorbreaks	Parameter specifying how the color distribution of the gene expression heatmap is determined, either equiquantiles or equidistant.

**Details**

The DNA copy number data heatmap is generated as follows. The DNA copy number data are used to determine the genomic segments exhibiting no difference in DNA copy number between the array elements that map to that segment. This resembles the dimension reduction technique employed in the `CGHregions`-package. Consequently, within a segment the DNA copy number for one sample is constant, but may vary between samples. Note that a region may comprise of a whole chromosome, but also of a focal amplification. It is the DNA copy number signature of the segments that is depicted in the heatmap of the DNA copy number data.

For the gene expression heatmap segments as constructed for the array CGH data are adopted. For each segment-sample combination the expression levels of the genes that map to that segment are averaged. Next, the gene expression data is also collapsed to the segment format. It is this collapsed and averaged expression data that is depicted in the corresponding heatmap.

**Author(s)**

Wessel N. van Wieringen: <w.vanwieringen@vumc.nl>

**References**

Van de Wiel, M.A., Van Wieringen, W.N. (2007), "CGHregions: dimension reduction for array CGH data with minimal information loss", *Cancer Informatics*, 2, 55-63.

Van Wieringen, W.N., Van de Wiel, M.A. (2009), "Non-parametric testing for DNA copy number induced differential mRNA gene expression", *Biometrics*, 65(1), 19-29.

**See Also**

cghCall, ExpressionSet, matchCGHcall2ExpressionSet, profilesPlot,

**Examples**

```
# load data
data(pollackCN16)
data(pollackGE16)

# plot heatmaps
CNHeatmaps(pollackCN16, pollackGE16, location = "mode", colorbreaks = "equiquantiles")
```

---

entropyTest

*One-sided two-sample test for entropy comparison*

---

**Description**

A one-sided two-sample test compares the entropy of a (high-dimensional) multivariate random variable between two groups. The test is one-sided: one group is a priori suspected to have a larger entropy. The null distribution is obtained via an efficient permutation resampling algorithm.

**Usage**

```
entropyTest(Y, id, nPerm = 1000, method = "normal", k0 = 1, k1 = 1,
center = TRUE, lowCiThres=0.10, ncpus=1, verbose=FALSE)
```

**Arguments**

Y	(High-dimensional) matrix. Rows are assumed to represent the samples, and columns represent the samples' genes or traits.
id	An indicator variable for the two groups to be compared. The groups should be coded as 0 and 1. There is an asymmetric interest in the groups: the group indicated by 1 is believed to exhibit a larger entropy.
nPerm	Number of permutations.
method	Distributional assumption under which entropy is to be estimated.
k0	k-nearest neighbor parameter for group comprising of samples indicated by a zero in the indicator variable id.
k1	k-nearest neighbor parameter for group comprising of samples indicated by a one in the indicator variable id.

center	Logical indicator: should the columns of Y be centered around zero?
lowCiThres	A value between 0 and 1. Determines speed of efficient p-value calculation. If the probability of a p-value being below lowCiThres is smaller than 0.001 (read: the test is unlikely to become significant), the permutation analysis is terminated and a p-value of 1.00 is reported.
ncpus	Number of cpus used for the permutations.
verbose	Logical indicator: should intermediate output be printed on the screen?

**Value**

Object of entTest-class.

**Author(s)**

Wessel N. van Wieringen: <w.vanwieringen@vumc.nl>

**References**

Van Wieringen, W.N., Van der Vaart, A.W. (2011), "Statistical analysis of the cancer cell's molecular entropy using high-throughput data", *Bioinformatics*, 27(4), 556-563.

Van Wieringen, W.N., Van de Wiel, M.A., Van der Vaart, A.W. (2008), "A test for partial differential expression", *Journal of the American Statistical Association*, 103(483), 1039-1049.

**See Also**

[hdEntropy](#)

**Examples**

```
# load data
data(pollackGE16)
Y <- exprs(pollackGE16)

# assign samples to groups
id <- sample(c(0,1), 41, replace=TRUE)

# perform testing and print test results
testRes <- entropyTest(t(Y), id, nPerm = 5, method="knn")
summary(testRes)
```

---

entTest-class

*Class "entTest" for storing the results of the function entropyTest.*

---

**Description**

The class entTest is the output of a call to [entropyTest](#). It stores results from a hypothesis test.

**Slots**

**statistic:** Object of class "numeric". Observed test statistic (i.e., estimated mutual information).

**p.value:** Object of class "numeric". P-value for the mutual information test.

**null.dist:** Object of class "numeric". The permutation null distribution for the test statistic.

**nperm:** Object of class "numeric". Number of permutation used for p-value calculation.

**remark:** Object of class "character". Tells whether the permutation algorithm was terminated prematurely or not.

**Methods**

**summary** signature(object = "entTest"): Prints the test results.

**Author(s)**

Wessel van Wieringen: <w.vanwieringen@vumc.nl>

**See Also**

entTest

**Examples**

```
showClass("entTest")
```

---

expandMatching2singleIDs

*Expand matching to single entries*

---

**Description**

In case a feature of platform 1 has been matched to multiple features of another platform, instead of averaging the data from these features, one may consider maintaining all features, each matched individually the feature of platform 1. This function modifies the results from the matching function `matchAnn2Ann` to facilitate this. The result can than directly be used in the subsetting functions `cghCall2weightedSubset` and `ExpressionSet2weightedSubset`.

**Usage**

```
expandMatching2singleIDs(matchedIDs)
```

**Arguments**

`matchedIDs` An object of class `list`, as returned by the `matchAnn2Ann`-function.

**Value**

An object of class `list`, similar to that returned by the `matchAnn2Ann`-function.

**Author(s)**

Wessel N. van Wieringen: <w.vanwieringen@vumc.nl>

## References

Van Wieringen, W.N., Unger, K., Leday, G.G.R., Krijgsman, O., De Menezes, R.X., Ylstra, B., Van de Wiel, M.A. (2012), "Matching of array CGH and gene expression microarray features for the purpose of integrative analysis", *BMC Bioinformatics*, 13:80.

## See Also

[matchAnn2Ann](#), [cghCall2weightedSubset](#), [ExpressionSet2weightedSubset](#).

## Examples

```
# load data
data(pollackCN16)
data(pollackGE16)

# extract genomic information from cghCall-object
chr1 <- fData(pollackCN16)[,1]
bpstart1 <- fData(pollackCN16)[,2]
bpend1 <- fData(pollackCN16)[,3]

# extract genomic information from ExpressionSet-object
chr2 <- fData(pollackGE16)[,1]
bpstart2 <- fData(pollackGE16)[,2]
bpend2 <- fData(pollackGE16)[,3]

# match features from both platforms
matchedFeatures <- matchAnn2Ann(chr1, bpstart1, bpend1, chr2,
bpstart2, bpend2, method = "distance", maxDist = 10000)

# expand
matchedFeatures <- expandMatching2singleIDs(matchedFeatures)
```

---

ExpressionSet2order     *Genomic ordering of ExpressionSet-objects.*

---

## Description

Orders the features within a [ExpressionSet](#)-object in accordance with their genomic order.

## Usage

```
ExpressionSet2order(GEdata, chr, bpstart, verbose=TRUE)
```

## Arguments

GEdata	Object of class <a href="#">ExpressionSet</a> .
chr	Column in the slot featureData of the <a href="#">ExpressionSet</a> -object specifying the chromosome information of the features.
bpstart	Column in the slot featureData of the <a href="#">ExpressionSet</a> -object specifying the start basepair information of the features.
verbose	Logical indicator: should intermediate output be printed on the screen?

**Value**

Object of class [ExpressionSet](#), now genomically ordered.

**Author(s)**

Wessel N. van Wieringen: <w.vanwieringen@vumc.nl>

**See Also**

[ExpressionSet](#).

**Examples**

```
# load data
data(pollackGE16)

# order the copy number data genomically
pollackGE16 <- ExpressionSet2order(pollackGE16, 1, 2)
```

---

ExpressionSet2subset *Subsetting ExpressionSet-objects.*

---

**Description**

Limit an [ExpressionSet](#) object to a subset of its features.

**Usage**

```
ExpressionSet2subset(GEdata, featureSubset, verbose=TRUE)
```

**Arguments**

GEdata	Object of class <a href="#">ExpressionSet</a> .
featureSubset	Object of class numeric, containing the row numbers of features to be maintained in the <a href="#">ExpressionSet</a> -object.
verbose	Logical indicator: should intermediate output be printed on the screen?

**Value**

Object of class [ExpressionSet](#), restricted to the specified subset of features.

**Author(s)**

Wessel N. van Wieringen: <w.vanwieringen@vumc.nl>

**See Also**

[ExpressionSet](#).

**Examples**

```
# load data
data(pollackGE16)

# order the copy number data genomically
pollackGE16 <- ExpressionSet2subset(pollackGE16, c(1:50))
```

---

ExpressionSet2weightedSubset

*Weighted subsetting ExpressionSet-objects.*

---

**Description**

Limit an [ExpressionSet](#) object to a subset of its features, using weighted averaging of the expression signal.

**Usage**

```
ExpressionSet2weightedSubset(GEdata, featuresAndWeights, chr,
                             bpstart, bpend, ncpus = 1, verbose=TRUE)
```

**Arguments**

GEdata	Object of class <a href="#">ExpressionSet</a> .
featuresAndWeights	Object of class <code>list</code> . Each list item is a matrix. The first column of this matrix contains the row numbers of features to be maintained in the <a href="#">ExpressionSet</a> -object. The second column contains the weights of each features, to be used in the calculation of the weighted average gene expression signal.
chr	Column in the slot <code>featureData</code> of the <a href="#">ExpressionSet</a> -object specifying the chromosome information of the features.
bpstart	Column in the slot <code>featureData</code> of the <a href="#">ExpressionSet</a> -object specifying the start basepair information of the features.
bpend	Column in the slot <code>featureData</code> of the <a href="#">ExpressionSet</a> -object specifying the end basepair information of the features.
ncpus	Number of cpus to be used in computations.
verbose	Logical indicator: should intermediate output be printed on the screen?

**Details**

Annotation information of features with multiplicity larger than one is compressed as follows. It is assumed that all features map to the same chromosome, leaving no ambiguity. The start base pair of the "new" feature is the smallest start base pair of features from which it has been formed. The end base pair of the "new" feature is the largest end base pair of features from which it has been formed.

**Value**

Object of class [ExpressionSet](#), restricted to the specified subset of features.



**Warning**

The phenoData, experimentData, and other slots of the ExpressionSet-object are currently not passed on to the subsetted object.

**Note**

This is a more intricate version of the ExpressionSet2subset function. They exist parallel because this function is much slower than its counterpart.

**Author(s)**

Wessel N. van Wieringen: <w.vanwieringen@vumc.nl>

**References**

Van Wieringen, W.N., Unger, K., Leday, G.G.R., Krijgsman, O., De Menezes, R.X., Ylstra, B., Van de Wiel, M.A. (2012), "Matching of array CGH and gene expression microarray features for the purpose of integrative analysis", *BMC Bioinformatics*, 13:80.

**See Also**

ExpressionSet2subset

**Examples**

```
# load data
data(pollackGE16)

# extract genomic information from ExpressionSet-object
chr <- fData(pollackGE16)[,1]
bpstart <- fData(pollackGE16)[,2]
bpend <- fData(pollackGE16)[,3]

# find unique genomic locations
uniqInfo <- uniqGenomicInfo(chr, bpstart, bpend, verbose = FALSE)

# subset ExpressionSet-object to features with unique genomic locations
pollackGE16 <- ExpressionSet2weightedSubset(pollackGE16, uniqInfo, 1, 2, 3)
```

---

getSegFeatures

*Identical signature features selection from cghCall-object.*

---

**Description**

Given an example, selects features (contiguous to the example) with the same signature (as the example) across samples from an [cghCall](#)-object.

**Usage**

```
getSegFeatures(featureNo, CNdata, verbose=TRUE)
```

**Arguments**

featureNo	Row number of example feature.
CNdata	Object of class <a href="#">cghCall</a> .
verbose	Logical indicator: should intermediate output be printed on the screen?

**Value**

Object of class `numeric`, containing the row numbers of those contiguous features with the same segmented log<sub>2</sub>-ratio signatures as `featureNo` across samples.

**Author(s)**

Wessel N. van Wieringen: <w.vanwieringen@vumc.nl>

**References**

Van Wieringen, W.N., Berkhof, J., Van de Wiel, M.A. (2010), "A random coefficients model for regional co-expression associated with DNA copy number", *Statistical Applications in Genetics and Molecular Biology*, Volume 9, Issue1, Article 25, 1-28.

**See Also**

[cghCall](#), [RCMestimation](#).

**Examples**

```
# load data
data(pollackCN16)

# feature of interest
featureNo <- 7

# extract all features with identical copy number signature (over the samples)
getSegFeatures(featureNo, pollackCN16)
```

---

hdEntropy

*Entropy estimation.*

---

**Description**

The (differential) entropy of a high-dimensional multivariate random variable is estimated from a (high-dimensional matrix) under a normality or k-NN distributional assumption.

**Usage**

```
hdEntropy(Y, method = "normal", k = 1, center = TRUE, indKnn = TRUE)
```

**Arguments**

Y	(High-dimensional) matrix. Rows are assumed to represent the samples, and columns represent the samples' genes or traits.
method	Distributional assumption under which entropy is to be estimated.
k	k-nearest neighbor parameter.
center	Logical indicator: should the columns of Y be centered around zero?
indKnn	Logical indicator: should samples' individual contributions to the k-NN entropy be reported?

**Value**

The entropy estimate is returned as a numeric.

**Author(s)**

Wessel N. van Wieringen: <w.vanwieringen@vumc.nl>

**References**

Van Wieringen, W.N., Van der Vaart, A.W. (2011), "Statistical analysis of the cancer cell's molecular entropy using high-throughput data", *Bioinformatics*, 27(4), 556-563.

**See Also**

[entropyTest](#).

**Examples**

```
data(pollackGE16)
hdEntropy(t(exprs(pollackGE16)), method="knn")
```

---

hdMI

*Mutual information estimation.*

---

**Description**

The mutual information between two high-dimensional multivariate random variables is estimated from two (high-dimensional matrix) under a normality or k-NN distributional assumption.

**Usage**

```
hdMI(Y, X, method = "normal", k = 1, center = TRUE, rescale = TRUE)
```

**Arguments**

Y	(High-dimensional) matrix. Rows are assumed to represent the samples, and columns represent the samples' genes or traits.
X	(High-dimensional) matrix. Rows are assumed to represent the samples, and columns represent the samples' genes or traits. The number of rows of X must be identical to that of Y.
method	Distributional assumption under which mutual information is to be estimated.
k	k-nearest neighbor parameter.
center	Logical indicator: should the columns (traits) of Y and X be centered at zero? Applied only under the normality assumption.
rescale	Logical indicator: should Y and X be rescaled to have the same scale? Applied only under the k-NN assumption.

**Value**

The mutual information estimate is returned as a numeric.

**Author(s)**

Wessel N. van Wieringen: <w.vanwieringen@vumc.nl>

**References**

Van Wieringen, W.N., Van der Vaart, A.W. (2011), "Statistical analysis of the cancer cell's molecular entropy using high-throughput data", *Bioinformatics*, 27(4), 556-563.

**See Also**

[mutInfTest](#).

**Examples**

```
data(pollackCN16)
data(pollackGE16)
hdMI(t(exprs(pollackGE16)), t(copynumber(pollackCN16)), method="knn")
```

---

matchAnn2Ann

*Genomic location matching of two sets of features*

---

**Description**

Genomic location matching of two sets of features

**Usage**

```
matchAnn2Ann(chr1, bpstart1, bpend1, chr2, bpstart2, bpend2,
method = "distance", maxDist = 10000, minPerc = 0,
reference = 1, ncpus = 1, verbose=TRUE)
```

**Arguments**

chr1	Object of class <code>numeric</code> containing chromosome information of features from set 1.
bpstart1	Object of class <code>numeric</code> containing start base pair information of features from set 1. Of same length as chr1.
bpend1	Object of class <code>numeric</code> containing end base pair information of features from set 1. Of same length as chr1.
chr2	Object of class <code>numeric</code> containing chromosome information of features from set 2.
bpstart2	Object of class <code>numeric</code> containing start base pair information of features from set 2. Of same length as chr2.
bpend2	Object of class <code>numeric</code> containing end base pair information of features from set 2. Of same length as chr2.
method	Matching method to be applied, either "distance" or "overlap". See below for details.
maxDist	Maximum number of bases two features are allowed to be separated for a match. Only used in combination with method="distance".
minPerc	Minimum percentage of overlap between two features required for a match. Only used in combination with method="overlap".
reference	Platform that is taken as a reference in the calculation of the percentage, should equal 1 or two, referring to the platform.
ncpus	Number of cpus to be used in the computation.
verbose	Logical indicator: should intermediate output be printed on the screen?

**Details**

The features of set 1 (chr1, bpstart1, bpend1) are matched to the features of set 2 (chr2, bpstart2, bpend2). That is, for every feature in set 2, features in set 1 are sought.

In case method="distance", the midpoint of set 1 and set 2 features are calculated and for each feature of set 2 all features of set 1 with midpoints not further than maxDist are selected. If there are no features in set 1 satisfying this criterion, the feature of set 2 that could not be matched is discarded.

If method="overlap", each feature of set 1 is matched to the feature of set 2 on the basis of the percentage of overlap. All features of set 1 with a percentage exceeding minPerc are selected. In case no feature in set 1 had any overlap with the features from set 2, the feature of set 2 that could not be matched is discarded.

**Value**

An object of class `list`. Each list item is a three-column matrix with the matched features information. The first column contains feature numbers of set 1 in the order as supplied. The second column contains feature numbers of set 2 in the order as supplied. Each row thus has two entries. The first entry contains the feature number of set 1 that has been matched to second entry, representing the feature number of set 2. The third column contains either the percentage of overlap (method="overlap") or the distance between the the midpoints of the two features (method="distance").

**Warning**

Base pair information of features from both sets should be on the same scale!

Features with incomplete annotation information are removed before matching. For clarity, they are not included in the object with matched features.

**Author(s)**

Wessel N. van Wieringen: <w.vanwieringen@vumc.nl>

**References**

Van Wieringen, W.N., Unger, K., Leday, G.G.R., Krijgsman, O., De Menezes, R.X., Ylstra, B., Van de Wiel, M.A. (2012), "Matching of array CGH and gene expression microarray features for the purpose of integrative analysis", *BMC Bioinformatics*, 13:80.

**See Also**

matchCGHcall2ExpressionSet

**Examples**

```
# load data
data(pollackCN16)
data(pollackGE16)

# extract genomic information from cghCall-object
chr1 <- fData(pollackCN16)[,1]
bpstart1 <- fData(pollackCN16)[,2]
bpend1 <- fData(pollackCN16)[,3]

# extract genomic information from ExpressionSet-object
chr2 <- fData(pollackGE16)[,1]
bpstart2 <- fData(pollackGE16)[,2]
bpend2 <- fData(pollackGE16)[,3]

# match features from both platforms
matchedFeatures <- matchAnn2Ann(chr1, bpstart1, bpend1, chr2,
bpstart2, bpend2, method = "distance", maxDist = 10000)
```

---

matchCGHcall2ExpressionSet

*Genomic location matching of CN and GE data*

---

**Description**

Integrative CN-GE analysis requires the copy number data of all genes on the expression array to be available. `intCNGEan.match` matches the features of the copy number platform to the genes of the expression array. This is done using their genomic locations on the basis of either proximity or overlap.

**Usage**

```
matchCGHcall2ExpressionSet(CNdata, GEdata, CNchr, CNbpstart,
  CNbpend, GEchr, GEbpstart, GEbpend, method = "distance",
  reference=1, ncpus=1, verbose=TRUE)
```

**Arguments**

CNdata	Object of class <code>cghCall</code> , containing (among others) annotation and call probabilities.
GEdata	Object of class <code>ExpressionSet</code> .
CNchr	Column in the slot <code>featureData</code> of the <code>cghCall</code> -object specifying the chromosome information of the features.
CNbpstart	Column in the slot <code>featureData</code> of the <code>cghCall</code> -object specifying the start basepair information of the features.
CNbpend	Column in the slot <code>featureData</code> of the <code>cghCall</code> -object specifying the end basepair information of the features.
GEchr	Column in the slot <code>featureData</code> of the <code>ExpressionSet</code> -object specifying the chromosome information of the features.
GEbpstart	Column in the slot <code>featureData</code> of the <code>ExpressionSet</code> -object specifying the start basepair information of the features.
GEbpend	Column in the slot <code>featureData</code> of the <code>ExpressionSet</code> -object specifying the end basepair information of the features.
method	Matching method to be applied, either "distance", "overlap" or "overlapPlus". See below for details.
reference	Platform that is taken as a reference in the calculation of the percentage, should equal 1 or two, referring to the platform.
ncpus	Number of cpus to be used in the computation.
verbose	Logical indicator: should intermediate output be printed on the screen?

**Details**

Ideally full annotation information (chromosome number, start base pair, end base pair) for both copy number and gene expression data is available. In case only start base pair information is available, let `CNbpend` and `GEbpend` refer to the same columns as `CNbpstart` and `GEbpstart`. Base pair information of copy number and expression data should be on the same scale.

Matching occurs on the basis of genomic locations. In case `method="distance"`, the midpoint of CN and GE features are calculated and for each gene on the expression array the closest feature of the copy number platform is selected. If `method="overlap"`, each gene in the `ExpressionSet`-object is matched to the feature from the copy number platform with the maximum percentage of overlap. If the maximum percentage of overlap equals zero, the gene is not included in the matched objects. If `method="overlapPlus"`, the features are first matched by their percentage of overlap (as with the `method="overlap"`-option). For all non-matched GE features its closest two CN features (one down- and one upstream) are determined. If the copy number signature of these two CN features is identical, intrapolation seems reasonable, and the GE feature is matched to the closest of these two CN features. Hence, `method="overlapPlus"` makes use of the copy number data, consequently, matching may be different for different data sets.

**Value**

A two-column matrix with the matched features entries. The first column contains feature numbers of the cghCall-object. The second column contains feature numbers of the ExpressionSet-object. Each row thus has two entries. The first entry contains the feature number of the cghCall-object that has been matched to second entry, representing the feature number of the ExpressionSet-object.

**Warning**

Features with incomplete annotation information are removed before matching. For clarity, they are not included in the objects with matched features.

**Note**

The matching process implemented here is different from the one implemented in the (deprecated) ACEit-package (Van Wieringen et al., 2006).

**Author(s)**

Wessel N. van Wieringen: <w.vanwieringen@vumc.nl>

**References**

Van Wieringen, W.N., Belien, J.A.M., Vosse, S.J., Achame, E.M., Ylstra, B. (2006), "ACE-it: a tool for genome-wide integration of gene dosage and RNA expression data", *Bioinformatics*, 22(15), 1919-1920.

Van Wieringen, W.N., Unger, K., Leday, G.G.R., Krijgsman, O., De Menezes, R.X., Ylstra, B., Van de Wiel, M.A. (2012), "Matching of array CGH and gene expression microarray features for the purpose of integrative analysis", *BMC Bioinformatics*, 13:80.

**See Also**

[cghCall](#), [ExpressionSet](#)

**Examples**

```
# load data
data(pollackCN16)
data(pollackGE16)

# match features from both platforms
featureMatch <- matchCGHcall2ExpressionSet(pollackCN16, pollackGE16, 1, 2, 3, 1, 2, 3)
```

---

merge2cghCalls

*Merge two cghCall-objects into one cghCall-object*

---

**Description**

Merge two cghCall-objects into one cghCall-object.

**Usage**

```
merge2cghCalls(CNdata1, CNdata2, verbose=TRUE)
```



**Arguments**

CNdata1	Object of class <a href="#">cghCall</a> .
CNdata2	Object of class <a href="#">cghCall</a> .
verbose	Logical indicator: should intermediate output be printed on the screen?

**Details**

Data of the two objects is assumed to originate from the same samples, and are presented in the same order.

Only the experimental data and annotation information is inherited by the merged object.

**Value**

Object of class [cghCall](#), restricted to the specified subset of features.

**Author(s)**

Wessel N. van Wieringen: <w.vanwieringen@vumc.nl>

**References**

Van de Wiel, M.A., Kim, K.I., Vosse, S.J., Van Wieringen, W.N., Wilting, S.M., Ylstra, B. (2007), "CGHcall: an algorithm for calling aberrations for multiple array CGH tumor profiles", *Bioinformatics*, 23, 892-894.

Van Wieringen, W.N., Unger, K., Leday, G.G.R., Krijgsman, O., De Menezes, R.X., Ylstra, B., Van de Wiel, M.A. (2012), "Matching of array CGH and gene expression microarray features for the purpose of integrative analysis", *BMC Bioinformatics*, 13:80.

**See Also**

[cghCall](#).

**Examples**

```
# load data
data(pollackCN16)

# create two cghCall-objects
ids1 <- sample(1:dim(pollackCN16)[1], 10)
CNdata1 <- pollackCN16[ids1,]
CNdata2 <- pollackCN16[-ids1,]

# order the copy number data genomically
pollackCN16 <- merge2cghCalls(CNdata1, CNdata2)
```

---

merge2ExpressionSets *Merge two ExpressionSet-objects into one ExpressionSet-object*

---

### Description

Merge two ExpressionSet-objects into one ExpressionSet-object

### Usage

```
merge2ExpressionSets(GEdata1, GEdata2, verbose=TRUE)
```

### Arguments

GEdata1	Object of class <a href="#">ExpressionSet</a> .
GEdata2	Object of class <a href="#">ExpressionSet</a> .
verbose	Logical indicator: should intermediate output be printed on the screen?

### Details

Data of the two objects is assumed to originate from the same samples, and are presented in the same order.

Only the experimental data and annotation information is inherited by the merged object.

### Value

Object of class [ExpressionSet](#), restricted to the specified subset of features.

### Author(s)

Wessel N. van Wieringen: <w.vanwieringen@vumc.nl>

### References

Van de Wiel, M.A., Kim, K.I., Vosse, S.J., Van Wieringen, W.N., Wilting, S.M., Ylstra, B. (2007), "CGHcall: an algorithm for calling aberrations for multiple array CGH tumor profiles", *Bioinformatics*, 23, 892-894.

Van Wieringen, W.N., Unger, K., Leday, G.G.R., Krijgsman, O., De Menezes, R.X., Ylstra, B., Van de Wiel, M.A. (2012), "Matching of array CGH and gene expression microarray features for the purpose of integrative analysis", *BMC Bioinformatics*, 13:80.

### See Also

[ExpressionSet](#).

## Examples

```
# load data
data(pollackGE16)

# create two cghCall-objects
ids1 <- sample(1:dim(pollackGE16)[1], 10)
GEdata1 <- pollackGE16[ids1,]
GEdata2 <- pollackGE16[-ids1,]

# order the copy number data genomically
pollackGE16 <- merge2ExpressionSets(GEdata1, GEdata2)
```

---

miTest-class

*Class "miTest" for storing the results of the function mutInfTest.*

---

## Description

The class miTest is the output of a call to [mutInfTest](#). It stores results from a hypothesis test.

## Slots

**statistic:** Object of class "numeric". Observed test statistic (i.e., estimated mutual information).

**p.value:** Object of class "numeric". P-value for the mutual information test.

**null.dist:** Object of class "numeric". The permutation null distribution for the test statistic.

**nperm:** Object of class "numeric". Number of permutation used for p-value calculation.

**remark:** Object of class "character". Tells whether the permutation algorithm was terminated prematurely or not.

## Methods

**summary** signature(object = "miTest"): Prints the test results.

## Author(s)

Wessel van Wieringen: <w.vanwieringen@vumc.nl>

## See Also

[mutInfTest](#)

## Examples

```
showClass("miTest")
```

---

mutInfTest	<i>Test for mutual information</i>
------------	------------------------------------

---

### Description

A test evaluates the significance of the mutual information between two (high-dimensional) multivariate random variables. The null distribution is obtained via an efficient permutation resampling algorithm.

### Usage

```
mutInfTest(Y, X, nPerm = 1000, method = "normal", k = 1, center = TRUE,
rescale = TRUE, lowCiThres=0.10, ncpus=1, verbose=FALSE)
```

### Arguments

Y	(High-dimensional) matrix. Columns are assumed to represent the samples, and rows represent the samples' genes or traits.
X	(High-dimensional) matrix. Columns are assumed to represent the samples, and rows represent the samples' genes or traits. The number of columns of X must be identical to that of Y.
nPerm	Number of permutations.
method	Distributional assumption under which mutual information is to be estimated.
k	k-nearest neighbor parameter.
center	Logical indicator: should the rows of Y and X be centered at zero? Applied only under the normality assumption.
rescale	Logical indicator: should Y and X be rescaled to have the same scale? Applied only under the k-NN assumption.
lowCiThres	A value between 0 and 1. Determines speed of efficient p-value calculation. If the probability of a p-value being below lowCiThres is smaller than 0.001 (read: the test is unlikely to become significant), the permutation analysis is terminated and a p-value of 1.00 is reported.
ncpus	Number of cpus used for the permutations.
verbose	Logical indicator: should intermediate output be printed on the screen?

### Value

Object of miTest-class.

### Author(s)

Wessel N. van Wieringen: <w.vanwieringen@vumc.nl>

### References

Van Wieringen, W.N., Van der Vaart, A.W. (2011), "Statistical analysis of the cancer cell's molecular entropy using high-throughput data", *Bioinformatics*, 27(4), 556-563.

Van Wieringen, W.N., Van de Wiel, M.A., Van der Vaart, A.W. (2008), "A test for partial differential expression", *Journal of the American Statistical Association*, 103(483), 1039-1049.

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

```
# load data
data(pollackCN16)
data(pollackGE16)
Y <- t(exprs(pollackGE16))
X <- t(copynumber(pollackCN16))

# perform testing and print test results
testRes <- mutInfTest(Y, X, nPerm = 1000)
summary(testRes)
```

---

nBreakpoints	<i>Number of breakpoints</i>
--------------	------------------------------

---

**Description**

The number of samples with at least one breakpoint is calculated for each transcript.

**Usage**

```
nBreakpoints(featuresAndWeights, CNdata)
```

**Arguments**

featuresAndWeights	Object of class <code>list</code> . Each list item is a matrix. The first column of this matrix contains the row numbers of features to be maintained in subsetting of the <code>cghCall</code> -object. The second column contains the weights of each features, to be used in the calculation of the weighted average copy number signal.
CNdata	Object of class <code>cghCall</code>

**Details**

For each item of the object `featuresAndWeights` the segmented data from the `cghCall`-object is used to determine whether a sample exhibits a breakpoint for this transcript.

**Value**

Object of class `numeric` containing the number of samples with at least one breakpoint. It is of the same length as the `featuresAndWeights`-object.

**Author(s)**

Wessel N. van Wieringen: <w.vanwieringen@vumc.nl>

## References

Van Wieringen, W.N., Unger, K., Leday, G.G.R., Krijgsman, O., De Menezes, R.X., Ylstra, B., Van de Wiel, M.A. (2012), "Matching of array CGH and gene expression microarray features for the purpose of integrative analysis", *submitted for publication*.

## See Also

[matchAnn2Ann](#).

## Examples

```
# load data
data(pollackCN16)
data(pollackGE16)

# extract genomic information from cghCall-object
chr1 <- fData(pollackCN16)[,1]
bpstart1 <- fData(pollackCN16)[,2]
bpend1 <- fData(pollackCN16)[,3]

# extract genomic information from ExpressionSet-object
chr2 <- fData(pollackGE16)[,1]
bpstart2 <- fData(pollackGE16)[,2]
bpend2 <- fData(pollackGE16)[,3]

# match features from both platforms
matchedIDs <- matchAnn2Ann(chr1, bpstart1, bpend1, chr2, bpstart2,
bpend2, method = "distance", maxDist = 10000)

# extract ids for object subsetting
matchedIDsCN <- lapply(matchedIDs, function(Z){ return(Z[, -1, drop=FALSE]) })

# calculate the number of breakpoints
nBreakpoints(matchedIDsCN, pollackCN16)
```

---

pathway1sample

*Penalized estimation of a pathway's regulatory network from DNA copy number and gene expression data (one-sample).*

---

## Description

The regulatory relationships between DNA copy number and gene expression within a pathway are modeled by a simultaneous-equations model. Parameters of this model are fitted by minimizing of a penalized least squares criterion. The employed penalty is that of the lasso, encouraging sparsity.

## Usage

```
pathway1sample(Y, X, lambda1 = 1, constr = TRUE, startCis=numeric(),
startTrans=matrix(), verbose = FALSE)
```

**Arguments**

Y	matrix. Rows are assumed to represent the samples, and columns represent the samples' gene expression levels.
X	matrix. Rows are assumed to represent the samples, and columns represent the samples' genes or traits. The number of rows and columns of X must be identical to that of Y.
lambda1	numeric or matrix. The lasso parameter. In case lambda1 is of class numeric and its length equals one, the same penalty parameter is applied to all <i>trans</i> -effects. In case lambda1 is of class matrix its column and row dimension equal the number of columns of Y. A possibly different penalty parameter is applied to each <i>trans</i> -effect.
constr	logical. Should the <i>cis</i> -effect (the direct effect of a column of X on column of Y) be positive?
startCis	numeric. Starting values for the <i>cis</i> -effect.
startTrans	matrix. Starting values for the <i>trans</i> -effect.
verbose	logical. Should intermediate output be printed on the screen?

**Details**

The model is fitted equation-by-equation. This is warranted by the assumption of independent errors. The expression levels of one gene is regressed on its own DNA copy number data and the expression levels of all other genes in the pathway.

**Value**

Object of class pathwayFit.

**Author(s)**

Wessel N. van Wieringen: <w.vanwieringen@vumc.nl>

**References**

Van Wieringen, W.N., Van de Wiel, M.A. (2012), "Modeling the *cis*- and *trans*-effect of DNA copy number aberrations on gene expression levels in a pathway", *submitted for publication*.

**See Also**

See also pathwayFit and pathway2sample.

**Examples**

```
# set number of genes (p) and samples (n)
p <- 10
n <- 1000

# sample cis-effects
beta <- abs(rnorm(p))

# sample trans-effects
Theta <- matrix(sample(c(-1,1), p^2, replace=TRUE, prob=c(0.2, 0.8)), ncol=p) *
matrix(runif(p^2), ncol=p) / 4
```

```

diag(Theta) <- 1

# sample error variances
Sigma <- diag(rchisq(p, df=1)/5 + 0.5)

# sample DNA copy number data
X <- matrix(runif(n*p, min=-2, max=2), ncol=p)

# sample gene expression data
Y <- t(apply(X, 1, function(Y, beta){ Y * beta }, beta=beta)) %% t(solve(Theta)) +
rmvnorm(n, sigma=solve(Theta) %% Sigma %% t(solve(Theta)))

# fit model
pFit <- pathway1sample(Y, X, lambda1=1, verbose=TRUE)

# compare fit to "truth" for cis-effects
plot(pFit@Cis ~ beta, pch=20)

# compare fit to "truth" for trans-effects
penFits <- c(pFit@Trans[upper.tri(Theta)], pFit@Trans[lower.tri(Theta)])
truth <- c(Theta[upper.tri(Theta)], Theta[lower.tri(Theta)])
plot(penFits ~ truth, pch=20)

```

---

pathway2sample

*Penalized estimation of a pathway's regulatory network from DNA copy number and gene expression data (two-sample).*

---

## Description

The regulatory relationships between DNA copy number and gene expression within a pathway are modeled by a simultaneous-equations model. Parameters of this model are fitted by minimizing of a penalized least squares criterion. The employed penalty is a combination of the lasso and the fused lasso. This combination encourages within-sample sparsity (lasso), and limits the between-sample differences (fused lasso).

## Usage

```

pathway2sample(Y, X, id, lambda1 = 1, lambdaF = 1, method = "FL",
  constr = TRUE, startCis = numeric(), startTrans1 =
    matrix(), startTrans2 = matrix(), epsilon = 0, verbose = FALSE)

```

## Arguments

- |    |  |
|----|--|
| Y  | Object of class <code>matrix</code> . Rows are assumed to represent the samples, and columns represent the samples' gene expression levels.  |
| X  | Object of class <code>matrix</code> . Rows are assumed to represent the samples, and columns represent the samples' genes or traits. The number of rows and columns of X must be identical to that of Y. |
| id | An indicator variable of class <code>numeric</code> for the two groups to be compared. The groups should be coded as 0 and 1.  |



lambda1	Either a numeric- or matrix-object. The lasso parameter. In case lambda1 is of class numeric its length is one, and the same penalty parameter is applied to all <i>trans</i> -effects. In case lambda1 is of class matrix its column and row dimension equal the number of columns of Y. A possibly different penalty parameter is applied to each <i>trans</i> -effect.
lambdaF	Either a numeric- or matrix-object. The fused lasso parameter. In case lambdaF is of class numeric and of length one, the same penalty parameter is applied to all differential <i>trans</i> -effects. In case lambdaF is of class matrix its column and row dimension equal the number of columns of Y. A possibly different penalty parameter is applied to each differential <i>trans</i> -effect.
method	A character-object. Indicates which penalty to employ (see details).
constr	logical. Should the <i>cis</i> -effect (the direct effect of a column of X on column of Y) be positive?
startCis	numeric. Starting values for the <i>cis</i> -effect.
startTrans1	matrix. Starting values for the <i>trans</i> -effect of group 1 (coded as 0).
startTrans2	matrix. Starting values for the <i>trans</i> -effect of group 2 (coded as 1).
epsilon	A numeric. Non-negative positive in the low-dimensional case. epsilon is to assume a positive value in the high-dimensional case.
verbose	logical. Should intermediate output be printed on the screen?

## Details

The model is fitted equation-by-equation. This is warranted by the assumption of independent errors. The expression levels of one gene is regressed on its own DNA copy number data and the expression levels of all other genes in the pathway.

The method-option indicates which penalty is combined with the least squares loss function. In case `methode = FL`, this the fused lasso penalty (as described in Van Wieringen, W.N., Van de Wiel, M.A., 2012):

$$\lambda_1 \|\Theta^{(a)}\|_1 + \lambda_1 \|\Theta^{(b)}\|_1 + \lambda_F \|\Theta^{(a)} - \Theta^{(b)}\|_1.$$

When `methode = FLs`, this penalty is simplified to:

$$\lambda_1 \|\Theta^{(a)} + \Theta^{(b)}\|_1 + \lambda_F \|\Theta^{(a)} - \Theta^{(b)}\|_1.$$

The use of this penalty may be motivated as follows. The two samples used to share a common network architecture. One expects only a relatively limited number of edges to have changed. Hence, the majority of edges will have the same sign, resulting in equality of the two penalties. An other motivation for this second penalty arises from the the observation that it is computationally faster. And, as

$$\lambda_1 \|\Theta^{(a)}\|_1 + \lambda_1 \|\Theta^{(b)}\|_1 \geq \lambda_1 \|\Theta^{(a)} + \Theta^{(b)}\|_1,$$

it penalizes less. As such, the resulting FLs penalized estimates may be used as starting values for fitting the model with the FL penalty.

## Value

Object of class `pathwayFit`.

## Author(s)

Wessel N. van Wieringen: <w.vanwieringen@vumc.nl>

## References

Van Wieringen, W.N., Van de Wiel, M.A. (2012), "Modeling the *cis*- and *trans*-effect of DNA copy number aberrations on gene expression levels in a pathway", *submitted for publication*.

## See Also

See also pathwayFit and pathway1sample.

## Examples

```
# set number of genes (p) and samples (n)
p <- 10
n <- 1000

# sample cis-effects
beta <- abs(rnorm(p))

# sample trans-effects for first sample
Theta1 <- matrix(sample(c(-1,1), p^2, replace=TRUE, prob=c(0.2, 0.8)), ncol=p) *
matrix(runif(p^2), ncol=p) / 4
diag(Theta1) <- 1

# sample trans-effects for second sample
idDiff <- sample(which(Theta1 != 1), 10)
Theta2 <- Theta1
Theta2[idDiff] <- -Theta1[idDiff]

# sample error variances
Sigma <- diag(rchisq(p, df=1)/5 + 0.5)

# sample DNA copy number data of sample 1
X1 <- matrix(runif(n*p, min=-2, max=2), ncol=p)

# sample gene expression data
Y1 <- t(apply(X1, 1, function(Y, beta){ Y * beta }, beta=beta)) %*% t(solve(Theta1)) +
rmvnorm(n, sigma=solve(Theta1) %*% Sigma %*% t(solve(Theta1)))

# sample DNA copy number data of sample 1
X2 <- matrix(runif(n*p, min=-2, max=2), ncol=p)

# sample gene expression data
Y2 <- t(apply(X2, 1, function(Y, beta){ Y * beta }, beta=beta)) %*% t(solve(Theta2)) +
rmvnorm(n, sigma=solve(Theta2) %*% Sigma %*% t(solve(Theta2)))

# construct id-vector
id <- c(rep(0, n), rep(1, n))

# fit model
pFit <- pathway2sample(Y=rbind(Y1, Y2), X=rbind(X1, X2), id=id, lambda1=0, lambdaF=0.01)

# compare fit to "truth" for cis-effects
plot(pFit@Cis ~ beta, pch=20)

# compare fit to "truth" for differential trans-effects
penFits1 <- c(pFit@Trans1[upper.tri(Theta1)], pFit@Trans1[lower.tri(Theta1)])
penFits2 <- c(pFit@Trans2[upper.tri(Theta2)], pFit@Trans2[lower.tri(Theta2)])
```

```
truth1 <- c(Theta1[upper.tri(Theta1)], Theta1[lower.tri(Theta1)])
truth2 <- c(Theta2[upper.tri(Theta2)], Theta2[lower.tri(Theta2)])
plot(penFits1 - penFits2, truth1 - truth2, pch=20)
cor(penFits1 - penFits2, truth1 - truth2, m="s")
```

---

pathwayFit-class      *Class "pathwayFit" for storing the results of the function pathway1sample or pathway2sample.*

---

## Description

The class `pathwayFit` is the output of a call to `pathway1sample` and `pathway2sample`. It stores results from fitting a simultaneous-equations model from DNA copy number and gene expression data.

## Slots

**Cis:** Object of class "numeric". Vector of estimated *cis*-effect.

**Trans:** Object of class "matrix". Matrix containing the *trans*-effects (one-sample only).

**Trans1:** Object of class "matrix". Matrix containing the *trans*-effects of the first sample (two-sample only).

**Trans2:** Object of class "matrix". Matrix containing the *trans*-effects of the second sample (two-sample only).

**Sigma:** Object of class "numeric". Vector of estimated residual variances.

**lambda1:** Object of class "matrix". Lasso parameter(s) employed.

**lambdaF:** Object of class "matrix". Fused lasso parameter(s) employed.

**constr:** Object of class "logical". Indicator for parameter constraints on *cis*-effect.

**epsilon:** Object of class "numeric". Constant used for the stabilization of estimation in a high-dimensional context.

**method:** Object of class "character". Indicator for method used in model fitting.

## Methods

**pathwayPlot** signature(object = "pathwayFit"): Plots the `pathwayFit`-object.

## Author(s)

Wessel N. van Wieringen: <w.vanwieringen@vumc.nl>

## See Also

See also `pathway1sample` and `pathway2sample`.

## Examples

```
showClass("pathwayFit")
```

---

pathwayPlot	<i>Plot of the pathway topology as reconstructed from DNA copy number and gene expression data (one-sample only).</i>
-------------	---

---

### Description

Plotting the topology of a pathway's regulatory network as reconstructed from DNA copy number and gene expression data by the `pathway1sample`-function.

### Usage

```
pathwayPlot(pFit, directed = TRUE, tWidth = 1, cWidth = 1, geWidth=10,
cnWidth=10, circleDist = 1.5, gNames = NULL, main = "", remove = FALSE)
```

### Arguments

<code>pFit</code>	Object of class <code>pathwayFit</code> as returned by the function <code>pathway1sample</code> .
<code>directed</code>	A logical indicating whether to plot directed or undirected <i>trans</i> -effects.
<code>tWidth</code>	A numeric that scales the width of the <i>trans</i> -effect edges.
<code>cWidth</code>	A numeric that scales the width of the <i>cis</i> -effect edges.
<code>geWidth</code>	A numeric that scales the width of the gene expression nodes.
<code>cnWidth</code>	A numeric that scales the width of the DNA copy number nodes.
<code>circleDist</code>	A numeric that scales the inner and outer circle.
<code>gNames</code>	A character containing the gene names to written inside the nodes.
<code>main</code>	The character to be plotted as plot title.
<code>remove</code>	A logical indicating whether to remove genes without <i>trans</i> -effects.

### Author(s)

Wessel N. van Wieringen: <w.vanwieringen@vumc.nl>

### References

Van Wieringen, W.N., Van de Wiel, M.A. (2012), "Modeling the *cis*- and *trans*-effect of DNA copy number aberrations on gene expression levels in a pathway", *submitted for publication*.

### See Also

See also `pathway1sample`.

### Examples

```
# set number of genes (p) and samples (n)
p <- 10
n <- 1000

# sample cis-effects
beta <- abs(rnorm(p))

# sample trans-effects
```

```
Theta <- matrix(sample(c(-1,1), p^2, replace=TRUE, prob=c(0.2, 0.8)), ncol=p) *
matrix(runif(p^2), ncol=p) / 4
diag(Theta) <- 1

# sample error variances
Sigma <- diag(rchisq(p, df=1)/5 + 0.5)

# sample DNA copy number data
X <- matrix(runif(n*p, min=-2, max=2), ncol=p)

# sample gene expression data
Y <- t(apply(X, 1, function(Y, beta){ Y * beta }, beta=beta)) %*% t(solve(Theta)) +
rmvnorm(n, sigma=solve(Theta) %*% Sigma %*% t(solve(Theta)))

# fit model
pFit <- pathway1sample(Y, X, lambda1=500)

# plot pathway topology
pathwayPlot(pFit, tWidth=5, cWidth=5)
```

---

pollackCN16

*Breast cancer data (copy number)*

---

## Description

Copy number data of chromosome 16 the breast cancer data set. Called using `CGHcall` with default settings, contains 240 features and 41 samples.

## Usage

```
data(pollackCN16)
```

## Format

An object of class `cghCall`.

## Source

Pollack, J. R., Sorlie, T., Perou, C. M., Rees, C. A., Jeffrey, S. S., Lonning, P. E., Tibshirani, R., Botstein, D., Borresen- Dale, A. L., Brown, P. O. (2002), "Microarray analysis reveals a major direct role of DNA copy number alteration in the transcriptional program of human breast tumors", *PNAS*, 99, 12963-12968.

## References

Van de Wiel, M. A., Kim, K. I., Vosse, S. J., Van Wieringen, W. N., Wilting, S. M., Ylstra, B. (2007), "CGHcall: Calling aberrations for array CGH tumor profiles", *Bioinformatics*, 23, 892-894.

## Examples

```
data(pollackCN16)
```

---

pollackGE16

*Breast cancer data (gene expression)*

---

### Description

Gene expression data of chromosome 16 of the breast cancer data set; contains 240 features and 41 samples.

### Usage

```
data(pollackGE16)
```

### Format

An object of class [ExpressionSet](#).

### Source

Pollack, J.R., Sorlie, T., Perou, C.M., Rees, C.A., Jeffrey, S.S., Lonning, P.E., Tibshirani, R., Botstein, D., Borresen-Dale, A.L., Brown, P.O. (2002), "Microarray analysis reveals a major direct role of DNA copy number alteration in the transcriptional program of human breast tumors", *PNAS*, 99, 12963-12968.

### Examples

```
data(pollackGE16)
```

---

profilesPlot

*CN-GE profiles plot*

---

### Description

Plots a sample's copy number and gene expression data side-by-side. This visualizes the relation between CN and GE within an individual sample.

### Usage

```
profilesPlot(CNdata, GEdata, sampleNo, chr = 0, verbose=TRUE)
```

### Arguments

CNdata	Object of class <a href="#">cghCall</a> , containing (among others) annotation and call probabilities. Features should be matched with those of the accompanying <a href="#">ExpressionSet</a> -object (as may be done using the <a href="#">matchCGHcall2ExpressionSet</a> -function).
GEdata	Object of class <a href="#">ExpressionSet</a> . Features should be matched with those of the accompanying <a href="#">cghCall</a> -object (as may be done using the <a href="#">matchCGHcall2ExpressionSet</a> -function).
sampleNo	Sample number of sample to be plotted. Corresponds to the order in which samples appear the CNdata- and GEdata-objects.
chr	Chromosome number for which the profiles are to be plotted. Default chr=0 for whole genome plotting.
verbose	Logical indicator: should intermediate output be printed on the screen?

**Details**

The blue lines in the gene expression profile plot are the median expressions of genes that map to the same copy number segment.

**Author(s)**

Wessel N. van Wieringen: <w.vanwieringen@vumc.nl>

**See Also**

[cghCall](#), [ExpressionSet](#)

**Examples**

```
# load data
data(pollackCN16)
data(pollackGE16)

# plot CN and GE profiles alongside
profilesPlot(pollackCN16, pollackGE16, 23, 16)
```

---

RCMestimation

*Fitting of the random coefficients model.*

---

**Description**

The parameters of the random coefficients model are estimated by means of the maximum likelihood method. The implemented maximum likelihood procedure has been optimized with respect to computational efficiency and memory usage.

**Usage**

```
RCMestimation(Y, X, R, hypothesis = "H2", shrinkType = "none",
  estType = "normal", corType = "unif", maxNoIt = 100,
  minSuccDist = 0.005, verbose = FALSE)
```

**Arguments**

Y	The matrix containing the (e.g., expression) data (number of columns equal to number of features, number of rows equal to number of samples).
X	The design matrix (number of rows equal to number of samples, number of columns equal to number of covariates).
R	The linear constraint matrix (number of columns equal to the number of covariates).
hypothesis	The hypothesis under which the model is fitted: H0 (H0 : R beta = 0 & tau2 = 0), H1 (H1 : R beta >= 0 & tau2 = 0), H2 (H2 : R beta >= 0 & tau2 >= 0).
shrinkType	The type of shrinkage to be applied to the error variances: none (shrinkage parameter is set equal to zero: no shrinkage), opt (shrinkage parameter is chosen to minimize the mean squared error criterion) or full (shrinkage parameter is set equal to one).

estType	Type of estimation, either normal (non-robust) or robust.
corType	Correlation structure to be used, either unif or ar1.
maxNoIt	Maximum number of iterations in the ML procedure.
minSuccDist	Minimum distance between estimates of two successive iterations to be achieved.
verbose	Logical indicator: should intermediate output be printed on the screen?

### Details

Details on the type of random coefficients model that is actually fitted are specified in the reference below.

### Value

Object of class `rcmFit`.

### Note

In case a covariate for the intercept is included in the design matrix  $X$  we strongly recommend the center, per feature, the data around zero.

### Author(s)

Wessel N. van Wieringen: <w.vanwieringen@vumc.nl>

### References

Van Wieringen, W.N., Berkhof, J., Van de Wiel, M.A. (2010), "A random coefficients model for regional co-expression associated with DNA copy number", *Statistical Applications in Genetics and Molecular Biology*, Volume 9, Issue1, Article 25, 1-28.

### See Also

[RCMrandom](#), [RCMtest](#), `rcmTest`.

### Examples

```
# load data
data(pollackCN16)
data(pollackGE16)

# select features belonging to a region
ids <- getSegFeatures(20, pollackCN16)

# extract segmented log2 ratios of the region
X <- t(segmented(pollackCN16)[ids[1], , drop=FALSE])

# extract segmented log2 ratios of the region
Y <- exprs(pollackGE16)[ids,]

# center the expression data (row-wise)
Y <- t(Y - apply(Y, 1, mean))

# specify the linear constraint matrix
R <- matrix(1, nrow=1)
```



```
# fit the random coefficients model to the random data
RCMresults <- RCMestimation(Y, X, R)
```

---

rcmFit-class                      *Class "rcmFit" for storing the results of the function RCMestimation.*

---

## Description

The class `rcmFit` is the output of a call to [RCMestimation](#). It stores results from fitting a random coefficients model.

## Slots

**betas:** Object of class "numeric". Vector of estimated global regression coefficients for each of the covariates in the design matrix.

**tau2s:** Object of class "numeric". Vector of estimated regression coefficient variances for each of the covariates in the design matrix X.

**sigma2s:** Object of class "numeric". Vector of estimated error variances for all genes.

**rho:** Object of class "numeric". Estimated correlation parameter between the error of two contiguous features.

**av.sigma2s:** Object of class "numeric". Average of the unshrunk estimated error variances.

**shrinkage:** Object of class "numeric". Applied shrinkage parameters in fitting the model.

**loglik:** Object of class "numeric". The log-likelihood of the fitted model.

**corType:** Object of class "character". Correlation structure of the error used.

**X:** Object of class "matrix". The design matrix.

## Methods

**.RCMloss** signature(object = "rcmFit"): Calculates the log-likelihood associated with the fitted model.

**RCMrandom** signature(object = "rcmFit"): Samples from the distribution induced by the fitted model.

**summary** signature(object = "rcmFit"): Prints the estimation result.

## Author(s)

Wessel N. van Wieringen: <w.vanwieringen@vumc.nl>

## See Also

[RCMestimation](#), [RCMrandom](#).

## Examples

```
showClass("rcmFit")
```

---

RCMrandom

*Random data from the random coefficients model.*

---

### Description

The significance of hypotheses regarding parameters of the random coefficients model is assessed by means of the parametric bootstrap. Hereto random data from the fitted model under the null hypothesis of interest are drawn. This function provides.

### Usage

```
RCMrandom(object)
```

### Arguments

object            Object of class `rcmFit`.

### Details

Details on the type of random coefficients model from which data are drawn are specified in the reference below.

### Value

A matrix of dimension (number of genes) times (number of samples).

### Author(s)

Wessel N. van Wieringen: <w.vanwieringen@vumc.nl>

### References

Van Wieringen, W.N., Berkhof, J., Van de Wiel, M.A. (2010), "A random coefficients model for regional co-expression associated with DNA copy number", *Statistical Applications in Genetics and Molecular Biology*, Volume 9, Issue1, Article 25, 1-28.

### See Also

[RCMestimation](#), [rcmFit](#).

### Examples

```
# load data
data(pollackCN16)
data(pollackGE16)

# select features belonging to a region
ids <- getSegFeatures(20, pollackCN16)

# extract segmented log2 ratios of the region
X <- t(segmented(pollackCN16)[ids[1], , drop=FALSE])

# extract segmented log2 ratios of the region
```

```

Y <- exprs(pollackGE16)[ids,]

# center the expression data (row-wise)
Y <- t(Y - apply(Y, 1, mean))

# specify the linear constraint matrix
R <- matrix(1, nrow=1)

# fit the random coefficients model to the random data
RCMresults <- RCMestimation(Y, X, R)

# draw random data
Yrandom <- RCMrandom(RCMresults)

```

---

RCMrandom-method      *Methods for Function RCMrandom*

---

### Description

Methods for function RCMrandom

### Methods

signature(object = "rcmFit") Draws random data of same dimension as data on which the rcmFit-object was fitted.

---

RCMtest      *Hypothesis testing within the random coefficient model.*

---

### Description

Function that evaluates various hypothesis within the random coefficients model via bootstrap re-sampling.

### Usage

```

RCMtest(Y, X, R, testType = "I", nBoot = 100, lowCiThres = 0.1,
shrinkType = "none", estType = "normal", corType = "unif",
maxNoIt = 100, minSuccDist = 0.005, returnNullDist = FALSE,
ncpus=1, verbose = FALSE)

```

### Arguments

Y	The matrix containing the (e.g., expression) data (number of columns equal to number of features, number of rows equal to number of samples).
X	The design matrix (number of rows equal to number of samples, number of columns equal to number of covariates).
R	The linear constraint matrix (number of columns equal to the number of covariates).

testType	The hypothesis to be tested: I ( $H_0 : R \beta = 0 \ \& \ \tau^2 = 0$ ) vs. ( $H_2 : R \beta \geq 0 \ \vee \ \tau^2 \geq 0$ ), II ( $H_0 : R \beta = 0 \ \& \ \tau^2 = 0$ ) vs. ( $H_1 : R \beta \geq 0 \ \& \ \tau^2 = 0$ ), III ( $H_1 : R \beta \geq 0 \ \& \ \tau^2 = 0$ ) vs. ( $H_2 : R \beta \geq 0 \ \& \ \tau^2 \geq 0$ ).
nBoot	Number of bootstraps.
lowCiThres	A value between 0 and 1. Determines speed of efficient p-value calculation. If the probability of a p-value being below lowCiThres is smaller than 0.001 (read: the test is unlikely to become significant), bootstrapping is terminated and a p-value of 1.00 is reported.
shrinkType	The type of shrinkage to be applied to the error variances: none (shrinkage parameter is set equal to zero: no shrinkage), opt (shrinkage parameter is chosen to minimize the mean squared error criterion) or full (shrinkage parameter is set equal to one).
estType	Type of estimation, either normal (non-robust) or robust.
corType	Correlation structure to be used, either unif or ar1.
maxNoIt	Maximum number of iterations in the ML procedure.
minSuccDist	Minimum distance between estimates of two successive iterations to be achieved.
returnNullDist	Logical indicator: should the null distribution be returned?
ncpus	Number of cpus used for the bootstrap.
verbose	Logical indicator: should intermediate output be printed on the screen?

### Details

Details on the type of random coefficients model that is actually fitted are specified in the reference below.

### Value

Object of class rcmTest.

### Warning

In case a covariate for the intercept is included in the design matrix  $X$  we strongly recommend the center, per feature, the data around zero.

### Author(s)

Wessel N. van Wieringen: <w.vanwieringen@vumc.nl>

### References

Van Wieringen, W.N., Berkhof, J., Van de Wiel, M.A. (2010), "A random coefficients model for regional co-expression associated with DNA copy number", *Statistical Applications in Genetics and Molecular Biology*, Volume 9, Issue1, Article 25, 1-28.

Van Wieringen, W.N., Van de Wiel, M.A., Van der Vaart, A.W. (2008), "A test for partial differential expression", *Journal of the American Statistical Association*, 103(483), 1039-1049.

### See Also

[RCMestimation](#), [RCMrandom](#), [rcmTest](#).

**Examples**

```

# load data
data(pollackCN16)
data(pollackGE16)

# select features belonging to a region
ids <- getSegFeatures(20, pollackCN16)

# extract segmented log2 ratios of the region
X <- t(segmented(pollackCN16)[ids[1], , drop=FALSE])

# extract segmented log2 ratios of the region
Y <- exprs(pollackGE16)[ids,]

# center the expression data (row-wise)
Y <- t(Y - apply(Y, 1, mean))

# specify the linear constraint matrix
R <- matrix(1, nrow=1)

# fit the random coefficients model to the random data
RCMresults <- RCMestimation(Y, X, R)

# test for significance of effect of X on Y
RCMtestResults <- RCMtest(Y, X, R, nBoot=2)
summary(RCMtestResults)

```

rcmTest-class

*Class "rcmTest" for storing the results of the function RCMtest.***Description**

The class `rcmTest` is the output of a call to `RCMtest`. It stores results from a hypothesis test.

**Slots**

**statistic:** Object of class "numeric". Observed test statistic (i.e., estimated mutual information).

**p.value:** Object of class "numeric". P-value for the mutual information test.

**betas:** Object of class "numeric". Vector of estimated global regression coefficients for each of the covariates in the design matrix.

**tau2s:** Object of class "numeric". Vector of estimated regression coefficient variances for each of the covariates in the design matrix.

**sigma2s:** Object of class "numeric". Vector of estimated error variances for all features.

**rho:** Object of class "numeric". Estimated correlation parameter between the error of two contiguous features.

**av.sigma2s:** Object of class "numeric". Average of the unshrunk estimated error variances.

**shrinkage:** Object of class "numeric". Type of shrinkage applied in the estimation.

**loglik:** Object of class "numeric". The log-likelihood of the fitted model.

**nBoot:** Object of class "numeric". Number of bootstraps used for p-value calculation.

`corType`: Object of class "character". Correlation structure used in the fitted model.  
`null.dist`: Object of class "numeric". The permutation null distribution for the test statistic.  
`remark`: Object of class "character". Tells whether the bootstrapping was terminated prematurely or not.

### Methods

**summary** signature(object = "rcmTest"): Prints the test results.

### Author(s)

Wessel van Wieringen: <w.vanwieringen@vumc.nl>

### See Also

[RCMtest](#)

### Examples

```
showClass("rcmTest")
```

---

splitMatchingAtBreakpoints  
*Split matching at breakpoints*

---

### Description

In case a feature of platform 1 has been matched to multiple features of another platform, instead of averaging the data from these features, one may consider splitting the data at breakpoints within genes. This function modifies the results from the matching function `matchAnn2Ann` to facilitate this. The result can then directly be used in the subsetting functions `cghCall2weightedSubset` and `ExpressionSet2weightedSubset`.

### Usage

```
splitMatchingAtBreakpoints(matchedIDs, CNdata)
```

### Arguments

`matchedIDs`      An object of class `list`, as returned by the `matchAnn2Ann`-function.  
`CNdata`            Object of class `cghCall`.

### Value

An object of class `list`, similar to that returned by the `matchAnn2Ann`-function.

### Author(s)

Wessel N. van Wieringen: <w.vanwieringen@vumc.nl>

## References

Van Wieringen, W.N., Unger, K., Leday, G.G.R., Krijgsman, O., De Menezes, R.X., Ylstra, B., Van de Wiel, M.A. (2012), "Matching of array CGH and gene expression microarray features for the purpose of integrative analysis", *BMC Bioinformatics*, 13:80.

## See Also

[matchAnn2Ann](#), [cghCall2weightedSubset](#), [ExpressionSet2weightedSubset](#).

## Examples

```
# load data
data(pollackCN16)
data(pollackGE16)

# extract genomic information from cghCall-object
chr1 <- fData(pollackCN16)[,1]
bpstart1 <- fData(pollackCN16)[,2]
bpend1 <- fData(pollackCN16)[,3]

# extract genomic information from ExpressionSet-object
chr2 <- fData(pollackGE16)[,1]
bpstart2 <- fData(pollackGE16)[,2]
bpend2 <- fData(pollackGE16)[,3]

# match features from both platforms
matchedFeatures <- matchAnn2Ann(chr1, bpstart1, bpend1, chr2,
bpstart2, bpend2, method = "distance", maxDist = 10000)

# expand
matchedFeatures <- splitMatchingAtBreakpoints(matchedFeatures, pollackCN16)
```

---

summary-method

*Methods for Function summary*

---

## Description

Methods for function summary

## Methods

```
signature(object = "ANY") Regular.
signature(object = "entTest") Print output.
signature(object = "miTest") Print output.
signature(object = "rcmFit") Print output.
signature(object = "rcmTest") Print output.
```

---

uniqGenomicInfo	<i>Unique genomic location information</i>
-----------------	--

---

**Description**

Finds unique genomic location information.

**Usage**

```
uniqGenomicInfo(chr, bpstart, bpend, verbose = FALSE)
```

**Arguments**

chr	Object of class <code>numeric</code> containing chromosome information of features.
bpstart	Object of class <code>numeric</code> containing start base pair information of features. Of same length as chr.
bpend	Object of class <code>numeric</code> containing end base pair information of features. Of same length as chr.
verbose	Logical indicator: should intermediate output be printed on the screen?

**Value**

An object of class `list`. Each list item is a four-column matrix with the matched features information. The first column contains feature numbers of features with identical genomic location. The second, third and fourth column contain the chromosome, start and end base pair information of the features (should be the same for each feature).

**Author(s)**

Wessel N. van Wieringen: <w.vanwieringen@vumc.nl>

**References**

Van Wieringen, W.N., Unger, K., Leday, G.G.R., Krijgsman, O., De Menezes, R.X., Ylstra, B., Van de Wiel, M.A. (2012), "Matching of array CGH and gene expression microarray features for the purpose of integrative analysis", *BMC Bioinformatics*, 13:80.

**See Also**

`ExpressionSet2weightedSubset`, `cghCall2weightedSubset`

**Examples**

```
# load data
data(pollackGE16)

# extract genomic information from ExpressionSet-object
chr <- fData(pollackGE16)[,1]
bpstart <- fData(pollackGE16)[,2]
bpend <- fData(pollackGE16)[,3]

# find unique genomic locations
uniqInfo <- uniqGenomicInfo(chr, bpstart, bpend, verbose = FALSE)
```



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