## Package 'GlobalAncova'

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Title Calculates a global test for differential gene expression between groups

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**Description** We give the following arguments in support of the

GlobalAncova approach: After appropriate normalisation, gene-expression-

data appear rather symmetrical and outliers

are no real problem, so least squares should be rather robust.

ANCOVA with interaction yields saturated data modelling e.g.

different means per group and gene. Covariate adjustment can

help to correct for possible selection bias. Variance

homogeneity and uncorrelated residuals cannot be expected.

Application of ordinary least squares gives unbiased, but no longer optimal estimates (Gauss-

Markov-Aitken). Therefore, using the classical F-test is inappropriate, due to

correlation. The test statistic however mirrors deviations

from the null hypothesis. In combination with a permutation

approach, empirical significance levels can be approximated.

Alternatively, an approximation yields asymptotic p-values.

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biocViews Microarray, OneChannel, Bioinformatics, Differential Expression, Pathways

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Depends methods, corpcor, globaltest

Imports annotate, AnnotationDbi

Suggests Biobase, annotate, GO.db, KEGG.db, golubEsets, hu6800.db,vsn, GSEABase, Rgraphviz

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colon.normal

Gene expression data

## Description

Normalized gene expression data of 12 patients with colorectal cancer. Samples are taken from inside the tumours. Additionally, from same patients samples are taken from normal tissue, see colon.normal. The expression matrix is only an exemplary subset of 1747 probe sets associated with cell proliferation.

## Usage

```
data(colon.normal)
```

## **Format**

```
The format is:
num [1:1747, 1:12] 8.74 10.53 8.48 12.69 8.55 ...
- attr(*, "dimnames")=List of 2
..$ : chr [1:1747] "200808_s_at" "215706_x_at" "217185_s_at" "202136_at" ...
..$: chr [1:12] "Co10.N.E.84.F.CEL" "Co14.N.E.89.F.CEL" "Co17.N.E.1037.F.CEL" "Co1.N.E.31.F.CEL" ...
```

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## References

Groene, J. et al., 2006, Transcriptional census of 36 microdissected colorectal cancers yields a gene signature to distinguish UICC II and III, *Int J Cancer* 119(8):1829–36.

## **Examples**

```
data(colon.normal)
#str(colon.normal)
```

colon.pheno

Covariate information for the colon data

## Description

Covariate data for the colon data example:

```
sex Sex of the patient.
```

age Age of the patient.

location Location of the tumour.

grade Histologic tumour grade.

UICC.stage UICC stage of colorectal carcinoma.

#### Usage

```
data(colon.pheno)
```

## **Format**

```
The format is:
```

```
'data.frame': 12 obs. of 5 variables:
$sex: Factor w/ 2 levels "0","1": 2 2 1 2 2 1 2 1 2 1 ...
$age: int 71 76 63 73 58 66 60 66 86 76 ...
$location: Factor w/ 2 levels "distal","proximal": 1 1 1 1 1 1 1 1 2 1 ...
$grade: Factor w/ 2 levels "2","3": 1 1 2 2 1 2 1 2 2 2 ...
$UICC.stage: Factor w/ 2 levels "2","3": 2 1 2 1 2 1 1 1 2 1 ...
```

#### References

Groene, J. et al., 2006, Transcriptional census of 36 microdissected colorectal cancers yields a gene signature to distinguish UICC II and III, *Int J Cancer* 119(8):1829–36.

## Examples

```
data(colon.pheno)
#str(colon.pheno)
```

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colon.tumour

Gene expression data

## **Description**

Normalized gene expression data of 12 patients with colorectal cancer. Samples are taken from inside the tumours. Additionally, from same patients samples are taken from normal tissue, see colon.normal. The expression matrix is only an exemplary subset of 1747 probe sets associated with cell proliferation.

## Usage

```
data(colon.tumour)
```

#### **Format**

```
The format is: num [1:1747, 1:12] 8.77 10.40 8.52 12.86 8.28 ... - attr(*, "dimnames")=List of 2 ... *: chr [1:1747] "200808_s_at" "215706_x_at" "217185_s_at" "202136_at" ... ... *: chr [1:12] "Co10.T.IT.83.F.CEL" "Co14.T.IT.88.F.CEL" "Co17.T.IT.563.F.CEL" "Co1.T.IT.30.F.CEL" ...
```

#### References

Groene, J. et al., 2006, Transcriptional census of 36 microdissected colorectal cancers yields a gene signature to distinguish UICC II and III, *Int J Cancer* 119(8):1829–36.

## **Examples**

```
data(colon.tumour)
#str(colon.tumour)
```

GlobalAncova

Global test for differential gene expression

## Description

Computation of a F-test for the association between expression values and clinical entities. In many cases a two way layout with gene and a dichotomous group as factors will be considered. However, adjustment for other covariates and the analysis of arbitrary clinical variables, interactions, gene co-expression, time series data and so on is also possible. The test is carried out by comparison of corresponding linear models via the extra sum of squares principle. Corresponding p-values, permutation p-values and/or asymptotic p-values are given.

There are three possible ways of using GlobalAncova. The general way is to define formulas for the full and reduced model, respectively, where the formula terms correspond to variables in

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model.dat. An alternative is to specify the full model and the name of the model terms that shall be tested regarding differential expression. In order to make this layout compatible with the function call in the first version of the package there is also a method where simply a group variable (and possibly covariate information) has to be given. This is maybe the easiest usage in cases where no 'special' effects like e.g. interactions are of interest.

## Usage

## **Arguments**

xx	Matrix of gene expression data, where columns correspond to samples and rows to genes. The data should be properly normalized beforehand (and log- or otherwise transformed). Missing values are not allowed. Gene and sample names can be included as the row and column names of xx.
formula.full	Model formula for the full model.
formula.red	Model formula for the reduced model (that does not contain the terms of interest.)
model.dat	Data frame that contains all the variable information for each sample.
group	Vector with the group membership information.
covars	Vector or matrix which contains the covariate information for each sample.
test.terms	Character vector that contains names of the terms of interest.
test.genes	Vector of gene names or a list where each element is a vector of gene names.
method	p-values can be calculated permutation-based ("permutation") or by means of an approximation for a mixture of chi-square distributions ("approx"). Both p-values are provided when specifying method = "both". With option "Fstat" only the global F-statistics are returned without p-values or further information.
perm	Number of permutations to be used for the permutation approach. The default is 10,000.
max.group.size	Maximum size of a gene set for which the asymptotic p-value is calculated. For bigger gene sets the permutation approach is used.
eps	Resolution of the asymptotic p-value.
acc	Accuracy parameter needed for the approximation. Higher values indicate higher

accuracy.

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#### Value

If test.genes = NULL a list with components

effect Name(s) of the tested effect(s)

ANOVA ANOVA table

test.result F-value, theoretical p-value, permutation-based and/or asymptotic p-value

terms Names of all model terms

If a collection of gene sets is provided in test.genes a matrix is returned whose columns show the number of genes, value of the F-statistic, theoretical p-value, permutation-based and/or asymptotic p-value for each of the gene sets.

#### Methods

- xx = "matrix", formula.full = "formula", formula.red = "formula", model.dat = "ANY", group = "missing", covars In this method, besides the expression matrix xx, model formulas for the full and reduced model and a data frame model.dat specifying corresponding model terms have to be given. Terms that are included in the full but not in the reduced model are those whose association with differential expression will be tested. The arguments group, covars and test.terms are "missing" since they are not needed for this method.
- xx = "matrix", formula.full = "formula", formula.red = "missing", model.dat = "ANY", group = "missing", covars = In this method, besides the expression matrix xx, a model formula for the full model and a data frame model.dat specifying corresponding model terms are required. The character argument test.terms names the terms of interest whose association with differential expression will be tested. The basic idea behind this method is that one can select single terms, possibly from the list of terms provided by previous GlobalAncova output, and test them without having to specify each time a model formula for the reduced model. The arguments formula.red, group and covars are "missing" since they are not needed for this method.
- xx = "matrix", formula.full = "missing", formula.red = "missing", model.dat = "missing", group = "ANY", covars = Besides the expression matrix xx a clinical variable group is required. Covariate adjustment is possible via the argument covars but more complex models have to be specified with the methods described above. This method emulates the function call in the first version of the package. The arguments formula.full, formula.red, model.dat and test.terms are "missing" since they are not needed for this method.

#### Note

This work was supported by the NGFN project 01 GR 0459, BMBF, Germany.

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with contributions from Sven Knueppel

## References

Mansmann, U. and Meister, R., 2005, Testing differential gene expression in functional groups, *Methods Inf Med* 44 (3).

#### See Also

```
Plot.genes, Plot.subjects, GlobalAncova.closed, GAGO, GlobalAncova.decomp
```

## **Examples**

```
data(vantVeer)
data(phenodata)
data(pathways)

GlobalAncova(xx = vantVeer, formula.full = ~metastases + ERstatus, formula.red = ~ERstatus, model.dat = phenodata,
GlobalAncova(xx = vantVeer, formula.full = ~metastases + ERstatus, test.terms = "metastases", model.dat = phenodata
GlobalAncova(xx = vantVeer, group = phenodata$metastases, covars = phenodata$ERstatus, test.genes=pathways[1], me
```

```
GlobalAncova gene set testing methods
```

Gene set testing of gene set databases using GlobalAncova

## **Description**

Three functions adapted from package **globaltest** to test gene sets from databases for association of the gene expression profile with a response variable. Three function are provided for KEGG, for Gene Ontology and for the Broad Institute's gene sets.

## Usage

#### **Arguments**

Matrix of gene expression data, where columns correspond to samples and rows хx to genes. Gene names have to be included as the row names of xx Arguments describing the tests to be performed are passed on to Global Ancova. Note that only the approximative version of GlobalAncova is used here and hence the parameter method is not available. Even though the number of permutations (perm) may be specified since very large gene sets (with more genes than max.group.size) are treated with the permutation test. id The identifier(s) of gene sets to be tested (character vector). If omitted, tests all gene sets in the database. annotation The name of the probe annotation package for the microarray that was used, or the name of the genome wide annotation package for the species (e.g. org. Hs.eg.db for human). If an organism package is given, the argument probe2entrez must be supplied. Use only if no probe annotation package is available. A mapping from probe probe2entrez identifiers to entrez gene ids. May be an environment, named list or named vector. multtest The method of multiple testing correction. Choose from: Benjamini and Hochberg FDR control (BH); Benjamini and Yekutieli FDR control (BY) or Holm familywise error control (holm). For GAGO also the focus level method is available. See focusLevel. If TRUE, sorts the results to increasing p-values. sort ontology The ontology or ontologies to be used. Default is to use all three ontologies. minsize The minimum number of probes that may be annotated to a gene set. Gene sets with fewer annotated probes are discarded. maxsize The maximum number of probes that may be annotated to a gene set. Gene sets with more annotated probes are discarded. focuslevel The focus level to be used for the focus level method. Either a vector of gene set ids, or a numerical level. In the latter case, findFocus is called with maxsize at the specified level to find a focus level. The Broad gene set collection, created by a call to getBroadSets. collection category The subcategory of the Broad collection to be tested. The default is to test all sets.

## **Details**

These are utility functions to make it easier to do gene set testing of gene sets available in gene set databases. The functions automatically retrieve the gene sets, preprocess and select them, perform global test, do multiple testing correction, and sort the results on the basis of their p-values. All functions require that annotate and the appropriate annotation packages are installed. GAKEGG additionally requires the KEGG.db package; GAGO requires the GO.db package; GABroad requires the user to download the XML file "msigdb\_v2.5.xml" from http://www.broad.mit.edu/gsea/downloads.jsp, and to preprocess that file using the getBroadSets function.

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#### Value

The function returns a data frame with raw and multiplicity-adjusted p-values for each gene set.

#### Note

Functions GAGO, GAKEGG and GABroad correspond to functions gtGO, gtKEGG and gtBroad in package **globaltest**. The difference is in the use of the GlobalAncova test instead of gt within the procedures.

#### Author(s)

Jelle Goeman: <j.j.goeman@lumc.nl>; Jan Oosting; Manuela Hummel

#### References

Goeman, J.J. and Mansmann, U., Multiple testing on the directed acyclic graph of Gene Ontology. Bioinformatics 2008; 24(4): 537-44.

#### See Also

gtGO, gtKEGG, gtBroad, GlobalAncova, gt,

## **Examples**

# see vignettes of packages GlobalAncova and globaltest and help of gtGO

GlobalAncova-methods Methods for Function GlobalAncova

#### **Description**

There are three possible ways of using GlobalAncova. The general way is to define formulas for the full and reduced model, respectively, where the formula terms correspond to variables in model.dat. An alternative is to specify the full model and the name of the model terms that shall be tested regarding differential expression. In order to make this layout compatible with the function call in the first version of the package there is also a method where simply a group variable (and possibly covariate information) has to be given. This is maybe the easiest usage in cases where no 'special' effects like e.g. interactions are of interest.

#### Methods

xx = "matrix", formula.full = "formula", formula.red = "formula", model.dat = "ANY", group = "missing", covars In this method, besides the expression matrix xx, model formulas for the full and reduced model and a data frame model.dat specifying corresponding model terms have to be given. Terms that are included in the full but not in the reduced model are those whose association with differential expression will be tested. The arguments group, covars and test.terms are "missing" since they are not needed for this method.

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xx = "matrix", formula.full = "formula", formula.red = "missing", model.dat = "ANY", group = "missing", covars =
In this method, besides the expression matrix xx, a model formula for the full model and a data
frame model.dat specifying corresponding model terms are required. The character argument
test.terms names the terms of interest whose association with differential expression will
be tested. The basic idea behind this method is that one can select single terms, possibly from
the list of terms provided by previous GlobalAncova output, and test them without having
to specify each time a model formula for the reduced model. The arguments formula.red,
group and covars are "missing" since they are not needed for this method.

xx = "matrix", formula.full = "missing", formula.red = "missing", model.dat = "missing", group = "ANY", covars = Besides the expression matrix xx a clinical variable group is required. Covariate adjustment is possible via the argument covars but more complex models have to be specified with the methods described above. This method emulates the function call in the first version of the package. The arguments formula.full, formula.red, model.dat and test.terms are "missing" since they are not needed for this method.

 $\begin{array}{ll} {\hbox{\sf GlobalAncova.closed}} & {\hbox{\it Closed testing procedure for testing several groups of genes using}} \\ {\hbox{\it GlobalAncova}} \end{array}$ 

## Description

Computation of a closed testing procedure for several groups of genes, e.g. pathways, as an alternative of correcting for multiple testing. Starting from the pathways of interest a family of null hypotheses is created that is closed under intersection. Each null hypothesis can be rejected at a given level if it is rejected along with all hypotheses included in it.

There are three possible ways of using GlobalAncova. Also GlobalAncova. closed can be invoked with these three alternatives.

max.group.size = 2500, eps = 1e-16, acc = 50)

## Usage

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#### **Arguments**

Matrix of gene expression data, where columns correspond to samples and rows

to genes. The data should be properly normalized beforehand (and log- or otherwise transformed). Missing values are not allowed. Gene and sample names

can be included as the row and column names of xx.

test.genes A list of named pathways that shall be tested, each containing vectors of gene

names.

previous.test The output of a call to Global Ancova with specified option test.genes accord-

ing to the pathways of interest (optional).

level The global level of significance of the testing procedure.

formula.full Model formula for the full model.

formula.red Model formula for the reduced model (that does not contain the terms of inter-

est).

model.dat Data frame that contains all the variable information for each sample.

group Vector with the group membership information.

covars Vector or matrix which contains the covariate information for each sample.

test.terms Character vector that contains names of the terms of interest.

method Raw p-values can be calculated permutation-based ("permutation") or by means

of an approximation ("approx").

perm Number of permutations to be used for the permutation approach. The default

is 10,000.

max.group.size Maximum size of a gene set for which the asymptotic p-value is calculated. For

bigger gene sets the permutation approach is used.

eps Resolution of the asymptotic p-value.

acc Accuracy parameter needed for the approximation. Higher values indicate higher

accuracy.

## Value

#### A list with components

new.data Family of null hypotheses (vectors of genes to be tested simultaneously with

GlobalAncova).

test.results Test results for each pathway of interest and all hypotheses included in it.

significant Names of the significant pathways.

not.significant

Names of the non significant pathways.

#### Methods

xx = "matrix", test.genes="list", formula.full = "formula", formula.red = "formula", model.dat = "ANY", group = '
In this method, besides the expression matrix xx and the list of gene groups test.genes,
model formulas for the full and reduced model and a data frame model.dat specifying corresponding model terms have to be given. Terms that are included in the full but not in the

reduced model are those whose association with differential expression will be tested. The arguments group, covars and test.terms are "missing" since they are not needed for this method.

- xx = "matrix", test.genes="list", formula.full = "formula", formula.red = "missing", model.dat = "ANY", group = "
  In this method, besides the expression matrix xx and the list of gene groups test.genes, a
  model formula for the full model and a data frame model.dat specifying corresponding model
  terms are required. The character argument test.terms names the terms of interest whose
  association with differential expression will be tested. The arguments formula.red, group
  and covars are "missing" since they are not needed for this method.
- xx = "matrix", test.genes="list", formula.full = "missing", formula.red = "missing", model.dat = "missing", group =
  Besides the expression matrix xx and the list of gene groups test.genes a clinical variable
  group is required. Covariate adjustment is possible via the argument covars but more complex models have to be specified with the methods described above. This method emulates the
  function call in the first version of the package. The arguments formula.full, formula.red,
  model.dat and test.terms are "missing" since they are not needed for this method.

#### Note

This work was supported by the NGFN project 01 GR 0459, BMBF, Germany.

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#### References

Marcus, R., Peritz, E. and Gabriel, K.R., 1976, On closed testing procedures with special reference to ordered analysis of variance, *Biometrika* 63 (3): 655–660.

#### See Also

GlobalAncova, Plot.genes, Plot.subjects

GlobalAncova.closed-methods

Methods for Function GlobalAncova.closed

## **Description**

There are three possible ways of using GlobalAncova, use methods? GlobalAncova for getting more information. Also GlobalAncova.closed can be invoked with these three alternatives.

## Methods

- xx = "matrix", test.genes="list", formula.full = "formula", formula.red = "formula", model.dat = "ANY", group = "In this method, besides the expression matrix xx and the list of gene groups test.genes, model formulas for the full and reduced model and a data frame model.dat specifying corresponding model terms have to be given. Terms that are included in the full but not in the reduced model are those whose association with differential expression will be tested. The arguments group, covars and test.terms are "missing" since they are not needed for this method.
- xx = "matrix", test.genes="list", formula.full = "formula", formula.red = "missing", model.dat = "ANY", group = "
  In this method, besides the expression matrix xx and the list of gene groups test.genes, a
  model formula for the full model and a data frame model.dat specifying corresponding model
  terms are required. The character argument test.terms names the terms of interest whose
  association with differential expression will be tested. The arguments formula.red, group
  and covars are "missing" since they are not needed for this method.
- xx = "matrix", test.genes="list", formula.full = "missing", formula.red = "missing", model.dat = "missing", group = Besides the expression matrix xx and the list of gene groups test.genes a clinical variable group is required. Covariate adjustment is possible via the argument covars but more complex models have to be specified with the methods described above. This method emulates the function call in the first version of the package. The arguments formula.full, formula.red, model.dat and test.terms are "missing" since they are not needed for this method.

GlobalAncova decomp GlobalAncova with sequential and type III sum of squares decomposition and adjustment for global covariates

## **Description**

Computation of a F-test for the association between expression values and clinical entities. The test is carried out by comparison of corresponding linear models via the extra sum of squares principle. In models with various influencing factors extra sums of squares can be treated with sequential and type III decomposition. Adjustment for global covariates, e.g. gene expression values in normal tissue as compared to tumour tissue, can be applied. Given theoretical p-values may not be appropriate due to correlations and non-normality. The functions are hence seen more as a descriptive tool.

## Usage

GlobalAncova.decomp(xx, formula, model.dat = NULL, method = c("sequential", "type3", "all"), test.gene

#### **Arguments**

xx Matrix of gene expression data, where columns correspond to samples and rows

to genes. The data should be properly normalized beforehand (and log- or otherwise transformed). Missing values are not allowed. Gene and sample names

can be included as the row and column names of xx.

formula Model formula for the linear model.

model.dat	Data frame that contains all the variable information for each sample.
method	Whether sequential or type III decomposition or both should be calculated.
test.genes	Vector of gene names or a list where each element is a vector of gene names.
genewise	Shall the sequential decomposition be displayed for each single gene in a (small) gene set?
zz	Global covariate, i.e. matrix of same dimensions as xx.
zz.per.gene	If set to TRUE the adjustment for the global covariate is applied on a gene-wise basis.

#### Value

Depending on parameters test.genes, method and genewise ANOVA tables, or lists of ANOVA tables for each decomposition and/or gene set, or lists with components of ANOVA tables for each gene are returned.

#### Note

This work was supported by the NGFN project 01 GR 0459, BMBF, Germany.

## Author(s)

```
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Manuela Hummel <hummel@ibe.med.uni-muenchen.de>
Urlich Mansmann <mansmann@ibe.med.uni-muenchen.de>
```

## See Also

```
Plot.sequential, pair.compare, GlobalAncova
```

## **Examples**

```
data(vantVeer)
data(phenodata)
data(pathways)

# sequential or type III decomposition
GlobalAncova.decomp(xx = vantVeer, formula = ~ grade + metastases + ERstatus, model.dat = phenodata, method = "sequence GlobalAncova.decomp(xx = vantVeer, formula = ~ grade + metastases + ERstatus, model.dat = phenodata, method = "type:
# adjustment for global covariate
data(colon.tumour)
data(colon.normal)
data(colon.pheno)
```

GlobalAncova.decomp(xx = colon.tumour, formula = ~ UICC.stage + sex + location, model.dat = colon.pheno, method = "a

pair.compare 15

pair.compare	Pairwise comparisons of factor levels within GlobalAncova	

## Description

Pairwise comparisons of gene expression in different levels of a factor by GlobalAncova tests. The method uses the reduction in residual sum of squares obtained when two respective factor levels are set to the same level. Holm-adjusted permutation-based p-values are given.

## Usage

```
pair.compare(xx, formula, group, model.dat = NULL, test.genes = NULL, perm = 10000)
```

## Arguments

xx	Matrix of gene expression data, where columns correspond to samples and rows to genes. The data should be properly normalized beforehand (and log- or otherwise transformed). Missing values are not allowed. Gene and sample names can be included as the row and column names of xx.
formula	Model formula for the linear model.
group	Factor for which pairwise comparisons shall be calculated.
model.dat	Data frame that contains all the variable information for each sample.
test.genes	Vector of gene names or a list where each element is a vector of gene names.
perm	Number of permutations to be used for the permutation approach. The default is 10,000.

## Value

An ANOVA table, or list of ANOVA tables for each gene set, for the pairwise comparisons.

## Note

This work was supported by the NGFN project 01 GR 0459, BMBF, Germany.

## Author(s)

```
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Urlich Mansmann <mansmann@ibe.med.uni-muenchen.de>
```

## See Also

GlobalAncova, GlobalAncova.decomp

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## **Examples**

```
data(vantVeer)
data(phenodata)
data(pathways)

pair.compare(xx = vantVeer, formula = ~ grade, group = "grade", model.dat = phenodata, test.genes = pathways[1:3], pair.compare(xx = vantVeer, formula = ~ grade, group = "grade", model.dat = phenodata, test.genes = pathways[1:3], pair.compare(xx = vantVeer, formula = ~ grade, group = "grade", model.dat = phenodata, test.genes = pathways[1:3], pair.compare(xx = vantVeer, formula = ~ grade, group = "grade", model.dat = phenodata, test.genes = pathways[1:3], pair.compare(xx = vantVeer, formula = ~ grade, group = "grade")
```

pathways

Cancer related pathways

## Description

A list of nine cancer related pathways corresponding to the van t'Veer data. Each element contains a vector gene names corresponding to those in the data set.

#### Usage

data(pathways)

## Format

```
The format is:
List of 9
$ androgen_receptor_signaling: chr [1:72] "AW025529" "NM_001648" "NM_001753" "NM_003298" ...
$ apoptosis : chr [1:187] "AB033060" "NM_002341" "NM_002342" "AI769763" ...
$ cell_cycle_control : chr [1:31] "NM_001759" "NM_001760" "NM_001786" "NM_001789" ...
$ notch_delta_signalling : chr [1:34] "NM_002405" "AL133036" "NM_003260" "NM_004316" ...
$ p53_signalling : chr [1:33] "NM_002307" "NM_002392" "NM_003352" "NM_002745" ...
$ ras_signalling : chr [1:266] "D25274" "AI033397" "NM_003029" "NM_001626" ...
$ tgf_beta_signaling : chr [1:82] "NM_003036" "AI090812" "AI697699" "AI760298" ...
$ tight_junction_signaling : chr [1:326] "D25274" "AA604213" "AF018081" "NM_003005" ...
$ wnt_signaling : chr [1:176] "AB033058" "AB033087" "NM_003012" "NM_003014" ...
```

## Examples

```
data(pathways)
#str(pathways)
```

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phenodata

Covariate information for the van t'Veer data

## **Description**

Covariate data for the van t'Veer example:

Sample Sample number.

metastases Development of distant metastases within five years (0-no/1-yes).

grade Tumor grade (three ordere levels).

**ERstatus** Estrogen receptor status (pos-positive/neg-negative).

## Usage

```
data(phenodata)
```

#### **Format**

The format is:

```
'data.frame': 96 obs. of 4 variables:
$Sample: int 1 2 3 4 5 6 7 8 9 10 ...
$metastases: int 0 0 0 0 0 0 0 0 0 ...
$grade: int 2 1 3 3 3 2 1 3 3 2 ...
```

\$ERstatus: Factor w/ 2 levels "neg", "pos": 2 2 1 2 2 2 2 1 2 2 ...

## **Examples**

```
data(phenodata)
#str(phenodata)
```

Plot.all

Combined visualization of sequential decomposition and influence of single genes on the GlobalAncova statistic

## **Description**

Plot that combines Plot.genes and Plot.sequential into one graphic.

## Usage

```
Plot.all(xx, formula, model.dat = NULL, test.genes = NULL, name.geneset = "")
```

Plot.genes

## **Arguments**

xx Matrix of gene expression data, where columns correspond to samples and rows

to genes. The data should be properly normalized beforehand (and log- or otherwise transformed). Missing values are not allowed. Gene and sample names

can be included as the row and column names of xx.

formula Model formula for the linear model.

model.dat Data frame that contains all the variable information for each sample.

test.genes Vector of gene names or gene indices specifying a gene set.

name.geneset Name of the plotted geneset.

## Note

This work was supported by the NGFN project 01 GR 0459, BMBF, Germany.

## Author(s)

```
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Reinhard Meister <meister@tfh-berlin.de>
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Urlich Mansmann <mansmann@ibe.med.uni-muenchen.de>
```

## See Also

```
Plot.genes, Plot.sequential, GlobalAncova.decomp, GlobalAncova
```

## **Examples**

```
data(vantVeer)
data(phenodata)
data(pathways)
```

Plot.all(vantVeer, formula = ~ ERstatus + metastases + grade, model.dat = phenodata, test.genes = pathways[[3]], na

Plot.genes	Genes Plot for Global Ancova

## **Description**

Produces a plot to show the influence of individual genes on the test result produced by GlobalAncova.

There are three possible ways of using GlobalAncova. Also Plot . genes can be invoked with these three alternatives.

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## Usage

```
## S4 method for signature 'matrix,formula,formula,ANY,missing,missing,missing'
Plot.genes(xx, formula.full, formula.red, model.dat, group, covars = NULL,test.terms,test.genes, Color
## S4 method for signature 'matrix,formula,missing,ANY,missing,missing,character'
Plot.genes(xx, formula.full, formula.red, model.dat, group, covars = NULL,test.terms,test.genes, Color
## S4 method for signature 'matrix,missing,missing,missing,ANY,ANY,missing'
Plot.genes(xx,formula.full, formula.red, model.dat, group, covars = NULL,test.terms,test.genes, Colors
```

## Arguments

XX	Matrix of gene expression data, where columns correspond to samples and rows to genes. The data should be properly normalized beforehand (and log- or otherwise transformed). Missing values are not allowed. Gene and sample names can be included as the row and column names of xx.
formula.full	Model formula for the full model.
formula.red	Model formula for the reduced model (that does not contain the terms of interest.)
model.dat	Data frame that contains all the variable information for each sample.
group	Vector with the group membership information.
covars	Vector or matrix which contains the covariate information for each sample.
test.terms	Character vector that contains names of the terms of interest.
test.genes	Vector of gene names or gene indices specifying the gene set. If missing, the plot refers to all genes in xx.
Colorgroup	Character variable giving the group that specifies coloring. If the function is called using the argument group then this variable is assumed to be relevant for coloring.
legendpos	Position of the legend (a single keyword from the list '"bottomright"', '"bottom"', '"bottomleft"', '"left"', '"topleft"', '"top"', '"topright"', '"right"' and '"center"').
returnValues	Shall bar heights (gene-wise reduction in sum of squares) be returned?
bar.names	Vector of bar labels. If missing, gene names from test.genes or row names of xx are taken.
	Graphical parameters for specifying colors, titles etc.

#### Methods

xx = "matrix", formula.full = "formula", formula.red = "formula", model.dat = "ANY", group = "missing", covars In this method, besides the expression matrix xx, model formulas for the full and reduced model and a data frame model.dat specifying corresponding model terms have to be given. Terms that are included in the full but not in the reduced model are those whose association with differential expression will be tested. The arguments group, covars and test.terms are "missing" since they are not needed for this method.

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```
xx = "matrix", formula.full = "formula", formula.red = "missing", model.dat = "ANY", group = "missing", covars = In this method, besides the expression matrix xx, a model formula for the full model and a data frame model.dat specifying corresponding model terms are required. The character argument test.terms names the terms of interest whose association with differential expression will be tested. The arguments formula.red, group and covars are "missing" since they are not needed for this method.
```

xx = "matrix", formula.full = "missing", formula.red = "missing", model.dat = "missing", group = "ANY", covars = Besides the expression matrix xx a clinical variable group is required. Covariate adjustment is possible via the argument covars but more complex models have to be specified with the methods described above. This method emulates the function call in the first version of the package. The arguments formula.full, formula.red, model.dat and test.terms are '"missing" since they are not needed for this method.

#### Note

This work was supported by the NGFN project 01 GR 0459, BMBF, Germany.

#### Author(s)

```
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Ulrich Mansmann <mansmann@ibe.med.uni-muenchen.de>
Manuela Hummel <hummel@ibe.med.uni-muenchen.de>
```

## See Also

```
GlobalAncova, Plot. subjects, Plot. sequential
```

## **Examples**

```
data(phenodata)
data(pathways)

Plot.genes(xx = vantVeer, formula.full = ~metastases + ERstatus, formula.red = ~ERstatus, model.dat = phenodata, te
Plot.genes(xx = vantVeer, formula.full = ~metastases + ERstatus, test.terms = "metastases", model.dat = phenodata,
Plot.genes(xx = vantVeer, group = phenodata$metastases, covars = phenodata$ERstatus, test.genes = pathways[[3]])
```

Plot.genes-methods

data(vantVeer)

Methods for Function Plot.genes

## **Description**

There are three possible ways of using GlobalAncova, use methods? GlobalAncova for getting more information. Also Plot. genes can be invoked with these three alternatives.

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#### Methods

xx = "matrix", formula.full = "formula", formula.red = "formula", model.dat = "ANY", group = "missing", covars In this method, besides the expression matrix xx, model formulas for the full and reduced model and a data frame model.dat specifying corresponding model terms have to be given. Terms that are included in the full but not in the reduced model are those whose association with differential expression will be tested. The arguments group, covars and test.terms are "missing" since they are not needed for this method.

- xx = "matrix", formula.full = "formula", formula.red = "missing", model.dat = "ANY", group = "missing", covars = In this method, besides the expression matrix xx, a model formula for the full model and a data frame model.dat specifying corresponding model terms are required. The character argument test.terms names the terms of interest whose association with differential expression will be tested. The arguments formula.red, group and covars are "missing" since they are not needed for this method.
- xx = "matrix", formula.full = "missing", formula.red = "missing", model.dat = "missing", group = "ANY", covars = Besides the expression matrix xx a clinical variable group is required. Covariate adjustment is possible via the argument covars but more complex models have to be specified with the methods described above. This method emulates the function call in the first version of the package. The arguments formula.full, formula.red, model.dat and test.terms are "missing" since they are not needed for this method.

Plot.sequential Visualization of sequential decomposition

## **Description**

Plot to show the sum of squares decomposition for each gene into parts according to all variables.

## Usage

Plot.sequential(xx, formula, model.dat = NULL, test.genes = NULL, name.geneset = "")

## **Arguments**

Matrix of gene expression data, where columns correspond to samples and rows

to genes. The data should be properly normalized beforehand (and log- or otherwise transformed). Missing values are not allowed. Gene and sample names

can be included as the row and column names of xx.

formula Model formula for the linear model.

model.dat Data frame that contains all the variable information for each sample.

test.genes Vector of gene names or gene indices specifying a gene set.

name.geneset Name of the plotted geneset.

#### Note

This work was supported by the NGFN project 01 GR 0459, BMBF, Germany.

Plot.subjects

#### Author(s)

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## See Also

```
GlobalAncova.decomp, Plot.genes, GlobalAncova
```

## **Examples**

```
data(vantVeer)
data(phenodata)
data(pathways)
```

 $Plot.sequential (vant Veer, formula = $^{\circ}$ ER status + metastases + grade, model.dat = phenodata, test.genes = pathways [[] and [] and [] are the properties of the proper$ 

Plot.subjects

Subjects Plot for GlobalAncova

## Description

Produces a plot to show the influence of the samples on the test result produced by GlobalAncova.

There are three possible ways of using GlobalAncova. Also Plot. subjects can be invoked with these three alternatives.

## Usage

```
## S4 method for signature 'matrix,formula,formula,ANY,missing,missing,missing'
Plot.subjects(xx, formula.full, formula.red, model.dat, group,covars = NULL, test.terms,test.genes, Co
## S4 method for signature 'matrix,formula,missing,ANY,missing,missing,character'
Plot.subjects(xx, formula.full,formula.red, model.dat, group,covars = NULL, test.terms,test.genes, Co]
## S4 method for signature 'matrix,missing,missing,missing,ANY,ANY,missing'
```

Plot.subjects(xx, formula.full, formula.red, model.dat, group, covars = NULL, test.terms, test.genes, Co

## Arguments

xx	Matrix of gene expression data, where columns correspond to samples and rows to genes. The data should be properly normalized beforehand (and log- or otherwise transformed). Missing values are not allowed. Gene and sample names can be included as the row and column names of xx.
formula.full	Model formula for the full model.
formula.red	Model formula for the reduced model (that does not contain the terms of interest.)

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model.dat Data frame that contains all the variable information for each sample.

group Vector with the group membership information.

covars Vector or matrix which contains the covariate information for each sample.

test. terms Character vector that contains names of the terms of interest.

test genes Vector of gene names or gene indices specifying the gene set. If missing, the

plot refers to all genes in xx.

Colorgroup Character variable giving the group that specifies coloring. If the function is

called using the argument group then this variable is assumed to be relevant for

coloring.

sort Should the samples be ordered by colorgroup?

legendpos Position of the legend (a single keyword from the list '"bottomright"', '"bot-

tom"', '"bottomleft"', '"left"', '"topleft"', '"top"', '"topright"', '"right"' and

'"center"').

returnValues Shall bar heights (subject-wise reduction in sum of squares) be returned?

bar.names Vector of bar labels. If missing, column names of xx are taken.

... Graphical parameters for specifying colors, titles etc.

#### Methods

xx = "matrix", formula.full = "formula", formula.red = "formula", model.dat = "ANY", group = "missing", covars
In this method, besides the expression matrix xx, model formulas for the full and reduced
model and a data frame model.dat specifying corresponding model terms have to be given.
Terms that are included in the full but not in the reduced model are those whose association
with differential expression will be tested. The arguments group, covars and test.terms
are "missing" since they are not needed for this method.

xx = "matrix", formula.full = "formula", formula.red = "missing", model.dat = "ANY", group = "missing", covars = In this method, besides the expression matrix xx, a model formula for the full model and a data frame model.dat specifying corresponding model terms are required. The character argument test.terms names the terms of interest whose association with differential expression will be tested. The arguments formula.red, group and covars are "missing" since they are not needed for this method.

xx = "matrix", formula.full = "missing", formula.red = "missing", model.dat = "missing", group = "ANY", covars = Besides the expression matrix xx a clinical variable group is required. Covariate adjustment is possible via the argument covars but more complex models have to be specified with the methods described above. This method emulates the function call in the first version of the package. The arguments formula.full, formula.red, model.dat and test.terms are "missing" since they are not needed for this method.

#### Note

This work was supported by the NGFN project 01 GR 0459, BMBF, Germany.

#### Author(s)

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#### See Also

GlobalAncova, Plot.genes, Plot.sequential

## **Examples**

data(vantVeer)

```
data(phenodata)
data(pathways)

Plot.subjects(xx = vantVeer, formula.full = ~metastases + ERstatus, formula.red = ~ERstatus, model.dat = phenodata
Plot.subjects(xx = vantVeer, formula.full = ~metastases + ERstatus, test.terms = "metastases", model.dat = phenodata
Plot.subjects(xx = vantVeer, group = phenodata$metastases, covars = phenodata$ERstatus, test.genes = pathways[[3]]
```

Plot.subjects-methods Methods for Function Plot.subjects

## **Description**

There are three possible ways of using GlobalAncova, use methods? GlobalAncova for getting more information. Also Plot. subjects can be invoked with these three alternatives.

#### Methods

- xx = "matrix", formula.full = "formula", formula.red = "formula", model.dat = "ANY", group = "missing", covars
  In this method, besides the expression matrix xx, model formulas for the full and reduced
  model and a data frame model.dat specifying corresponding model terms have to be given.
  Terms that are included in the full but not in the reduced model are those whose association
  with differential expression will be tested. The arguments group, covars and test.terms
  are "missing" since they are not needed for this method.
- xx = "matrix", formula.full = "formula", formula.red = "missing", model.dat = "ANY", group = "missing", covars = In this method, besides the expression matrix xx, a model formula for the full model and a data frame model.dat specifying corresponding model terms are required. The character argument test.terms names the terms of interest whose association with differential expression will be tested. The arguments formula.red, group and covars are "missing" since they are not needed for this method.
- xx = "matrix", formula.full = "missing", formula.red = "missing", model.dat = "missing", group = "ANY", covars = Besides the expression matrix xx a clinical variable group is required. Covariate adjustment is possible via the argument covars but more complex models have to be specified with the methods described above. This method emulates the function call in the first version of the package. The arguments formula.full, formula.red, model.dat and test.terms are "missing" since they are not needed for this method.

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vantVeer

Gene expression data

## Description

Normalized gene expression data for the van t'Veer example: A subset of 96 samples without BRCA1 or BRCA2 mutations and 1113 genes associated with nine cancer related pathways (see also ?pathways) was chosen.

## Usage

```
data(vantVeer)
```

## **Format**

```
The format is:
```

```
num [1:1113, 1:96] 0.13 0.936 -0.087 0.118 0.168 -0.081 0.023 -0.086 -0.154 0.025 ... - attr(*, "dimnames")=List of 2 ...$ : chr [1:1113] "AW025529" "NM_001648" "NM_001753" "NM_003298" ... ... : chr [1:96] "1" "2" "3" "4" ...
```

## **Examples**

```
data(vantVeer)
#str(vantVeer)
```

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