Package ‘PANR’

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R topics documented:

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assoScore

Association scores for gene pairs

Description
This function compute similarity scores to quantify associations between pairs of genes, given measured rich phenotypes.

Usage
assoScore(pheno, metric="cosine", upperTri=TRUE, transform=TRUE, verbose=TRUE, ...)

Arguments
- pheno: a numeric matrix of rich phenotypes with rows and columns specifying samples and genes, respectively.
- metric: a character value specifying the metric to compute similarity scores. Currently, 'cosine' and 'correlation' are supported (see details for more).
- upperTri: a logical value specifying whether (if TRUE) to take the upper triangular of the similarity matrix or not (if FALSE).
- transform: a logical value specifying whether to transform (if TRUE) the range of association scores from \([-1, 1]\) to \([0, 1]\) or not (if FALSE).
- verbose: a logical value to switch on (if TRUE) or off if FALSE detailed run-time message.
- ...: other arguments for function cor

Details
This function aims at quantifying the associations between genes of interest given certain phenotyping measurements (e.g. gene expressions by microarray, cell viabilities, morphological phenotypes, etc.). For the current version of the package, the user can either choose 'cosine' or 'correlation'. When the latter is chosen, additional arguments (e.g. 'method') for the function cor are allowed.

Value
This function will return either a vector (if upperTri=TRUE) or a matrix (if upperTri=FALSE) of association scores for given phenotypes.
BetaMixture-class

Author(s)

Xin Wang <xw264@cam.ac.uk>

References


See Also
cor, cosineSim

Examples

```
toydata<-matrix(rnorm(n=2000, mean=0, sd=4), nrow=100, ncol=20)
toyasso<-assoScore(t(toydata), "cosine", upperTri=FALSE, transform=FALSE)
# transform to [0, 1]
toyasso01<-assoScore(t(toydata), "cosine", upperTri=FALSE, transform=TRUE)
# transform to [0, 1] and return only the upper triangular
toyasso01upper<-assoScore(t(toydata), "cosine", upperTri=TRUE, transform=TRUE)
# use spearman correlation
toyassoSp<-assoScore(t(toydata), "correlation", upperTri=FALSE, transform=FALSE, method="spearman")
```

BetaMixture-class  An S4 class for beta mixture modelling of functional gene associations

Description

This S4 class includes methods to do beta-mixture modelling of functional gene associations given rich phenotyping screens.

Objects from the Class

Objects of class BetaMixture can be created from `new("BetaMixture", metric, order, association, model, pheno, partition)` (see the example below for details).

Slots

- `pheno`: a numeric matrix of rich phenotypes with rows and columns specifying genes and samples, respectively.
- `metric`: a character value specifying the metric to compute similarity scores. Currently, 'cosine' and 'correlation' are supported (see assoScore for more details).
- `order`: a numeric value specifying the order of the similarity score to be computed. Only 1 and 2 are supported for the current version. The first order (when `order=1`) similarities are used for quantification of the strength of functional associations between genes, whilst the second order (when `code=2`) ones are employed to compute the strength of modularity between genes.
- `association`: a numeric vector providing all association scores between genes. This can be useful when `pheno` is not available or the user has a different way to compute functional associations.
model: a character value specifying whether the original (if global) or extended (if stratified) model is used.

partition: a numeric of gene partition labels (e.g. `c(rep(1, 100), rep(2, 20), rep(3, 80))`) is a valid vector of partition labels for a vector of associations falling into three categories of interaction types 1, 2 and 3).

result: a list storing results from S4 methods of this class.

summary: a list of summary information for available results.

Methods

An overview of methods (More detailed introduction can be found in help for each specific function.):

- `permNULL` do permutations for input rich phenotyping screens (`'pheno'`).
- `fitNULL` fit the permuted association scores to a beta distribution.
- `fitBM` fit the functional association scores computed from input screens to a three-beta mixture model.
- `p2SNR` Translate p-values to Signal-to-Noise Ratios.
- `SNR2p` Translate Signal-to-Noise Ratios to p-values.
- `view` view the fitting results (a histogram of the original data and fitted probability density curves) for NULL and real data.
- `summarize` summarize results including input data and parameters, NULL fitting and beta mixture fitting.

Author(s)

Xin Wang <xw264@cam.ac.uk>

References


See Also

`permNULL fitNULL fitBM view summarize`

Examples

```R
## Not run:
data(Bakal2007)
bm1 <- new("BetaMixture", pheno=Bakal2007, metric="cosine", model="global", order=1)
bm1 <- fitNULL(bm1, nPerm=10, thetaNULL=c(alphaNULL=4, betaNULL=4), sumMethod="median", permMethod="all", verbose=TRUE)
bm1 <- fitBM(bm1, para=list(zInit=NULL, thetaInit=c(alphaNeg=2, betaNeg=4, alphaNULL=bm1@result$fitNULL$thetaNULL[["alphaNULL"]], betaNULL=bm1@result$fitNULL$thetaNULL[["betaNULL"]], alphaPos=4, betaPos=2), gamma=NULL), ctrl=list(fitNULL=FALSE, tol=1e-1), verbose=TRUE, gradtol=1e-3)
view(bm1, "fitNULL")
```
buildPAN

view(bm1, "fitBM")
bm1

## End(Not run)

---

**buildPAN**

*Build an igraph or RedeR graph for PAN*

**Description**

The function builds a graph for the inferred PAN so that it can be visualize in igraph or RedeR.

**Usage**

```r
buildPAN(object, engine="igraph", para=list(nodeColor=NULL, nodeSize=NULL, edgeWidth=NULL, edgeColor=NULL, nodeSumCols=1, nodeSumMethod="none", hideNeg=TRUE), verbose=TRUE, ...)
```

**Arguments**

- `object` an object of S4 class `PAN`.
- `engine` a character value specifying which graphics engine to use: ‘igraph’ or ‘RedeR’.
- `para` a list of parameters specifying graph attributes (see details)
- `verbose` a logical value to switch on (if `TRUE`) or off if `FALSE` detailed run-time message.
- `...` all the other arguments accepted by the function `pvclust`.

**Details**

Here are the introductions for the detailed graph attributes that the user can specify:

- `'nodeColor'` and `'nodeSize'` - a vector of node colors or sizes. Please note that the order of color or size must be concordant with the gene ids in PANR:the argument `pheno` of the object of class `BetaMixture`.
- `'nodeSumCols'` and `'nodeSumMethod'` - these two arguments are used to scale the colors of nodes by phenotypes. The former argument is a numeric vector specifying the columns in slot `pheno` of the object of class `BetaMixture`; while the latter one is a character value giving the method to summarize these columns of phenotypes: either ‘mean’ or ‘median’.
- `'edgeColor'` and `'edgeWidth'` - a vector of edge colors or width.
- `'pValCutoff'` - the argument is only used when `what='module'` and for module searching based on `pvclust`. Only significant modules will be displayed.
- `'minSize'` and `'maxSize'` - two arguments controlling the size of modules which will be used to filtered out modules that are too small or too large.
- `'hideNeg'` - a logical value specifying whether or not to hide edges with negative associations.

**Value**

This function will return an object of class `PAN` with inferred gene modules (`modules$clusters`) and corresponding p-values (`modules$pval`) updated in slot 'modules'.
Author(s)

Xin Wang <xw264@cam.ac.uk>

References


Examples

data(bm, package="PANR")
pan<-new("PAN", bm=bm)
pan<-infer(pan, para=list(type="SNR", log=TRUE, sign=TRUE, cutoff=log(5)), filter=FALSE, verbose=TRUE)
data(Bakal2007Cluster, package="PANR")
pan<-buildPAN(pan, engine="igraph", para=list(nodeColor=nodeColor, hideNeg=TRUE), verbose=TRUE)

---

**cosineSim**

**Compute cosine similarities or distances between pairs of genes**

Description

This function compute cosine similarities or distances between pairs of genes, given measured rich phenotypes.

Usage

```r
cosineSim(x)
cosineDist(x)
```

Arguments

- `x` a numeric matrix of rich phenotypes with rows and columns specifying samples and genes, respectively.

Value

This function will return either a numeric matrix of cosine similarities or an object of `dist`.

Author(s)

Xin Wang <xw264@cam.ac.uk>

References

Rich morphological phenotypes for gene overexpression and RNA interference screens

The data set we use here comes from quantitative morphological screening for 249 gene-overexpression or RNAi knock-down experiments. For each individual cell, 145 different geometric features were computed by imaging analysis, and are subsequently scored with NNs trained to discriminate seven reference TCs with distinctive morphologies. For each TC, NN z-scores were computed from all scored cells in this TC (more details in Bakal 2007).

Bakal2007: a matrix of NN z-scores with rows and columns corresponding to 249 TCs and seven reference TCs.

Bakal2007Cluster: unsupervised hierarchical clustering results by Bakal et al.
nodeColor: colors scaled according to the clustering results by Bakal et al.

bm1: an object of BetaMixture, which includes data and results for beta-mixture modelling on the association densities based on first-order cosine similarities of the phenotyping screens.

Usage

##see example for details

Author(s)

Xin Wang <xw264@cam.ac.uk>

References


Examples

data(Bakal2007)
dim(Bakal2007)
data(bm)
ob1

exportPAN

Export inferred PAN or module graphs to files

Description
Powered by function write.graph in package igraph, this function writes the inferred PAN or module graphs to files in a variety of formats that are supported by igraph.

Usage
exportPAN(object, file="pan", what="graph", moduleID=1, format="gml", verbose=TRUE, ...)

Arguments
- object: an object of S4 class PAN.
- file: the name of the file to write the graph or module (no need to put a filename extension).
- what: a character value specifying what to write: 'graph' or 'pvclustModule'.
- moduleID: a numeric or integer vector of modules to write (see details in sigModules).
- format: a character value specifying the format to write (see more details in write.graph).
- verbose: a logical value to switch on (if TRUE) or off if FALSE detailed run-time message.
- ...: not in use, but only for further extension.

Value
a numeric vector of ids for significant gene modules

Author(s)
Xin Wang <xw264@cam.ac.uk>

Examples
## Not run:
data(bm, package="PANR")
pan<-new("PAN", bm1=bm)
pan<-infer(pan, para=list(type="SNR", log=TRUE, sign=TRUE, cutoff=log(5)),
filter=FALSE, verbose=TRUE)
data(Bakal2007Cluster, package="PANR")
pan<-buildPAN(pan, engine="igraph", para=list(nodeColor=nodeColor,
hideNeg=TRUE), verbose=TRUE)
exportPAN(pan, file="pan", what="graph", format="ncol")
## End(Not run)
fitBM

**Fit a three-beta mixture model to densities of functional gene associations**

---

**Description**

The function fits a three-beta mixture model to densities of functional gene associations computed from rich phenotyping screens.

**Usage**

```r
fitBM(object, para=list(zInit=NULL, thetaInit=c(alphaNeg=2, betaNeg=4, alphaNULL=4, betaNULL=4, alphaPos=4, betaPos=2), gamma=NULL), ctrl=list(fitNULL=FALSE, tol=1e-3, maxIter=NULL), verbose=TRUE, ...)
```

**Arguments**

- `object`: an object of S4 class `BetaMixture`.
- `para`: a list of initial values for parameter estimation in fitting a three-beta mixture model (see 'details').
- `ctrl`: a list of control parameters for the mixture model fitting (see 'details').
- `verbose`: a logical value to switch on (if `TRUE`) or off (if `FALSE`) detailed run-time messages.
- `...`: other arguments of the function `nlm`.

**Details**

This function fits a beta-mixture model to functional gene associations using the Expectation-Maximization algorithm. The function allows various parameter settings to perform fitting by the original (if `model='global'`) or stratified (if `model='stratified'`) beta-mixture model (the model should be specified when creating a new object of `BetaMixture`).

The initial values of the beta distributions can be set by `thetaInit`, a numeric vector including the two shape parameters for the '-' (negative), 'x' (NULL) and '+' (positive) distributions. Please note that if `ctrl$NULL` is `TRUE`, meaning that the NULL distribution has already been fitted, then `para$alphaNULL` and `para$betaNULL` are supposed to be filled in the estimated NULL parameters by the function `fitNULL`.

`zInit` is a matrix of posterior probabilities for gene associations following the three mixture components.

The hyper-parameters for the dirichlet priors for the mixture components can also be set by `para.gamma`, which is a numeric vector with rows and columns corresponding to association partitions and the three beta mixture components.

The other arguments to control the fitting algorithm are `tol` and `maxIter`, which are convergence tolerance and the maximal iterations.

Since the estimation of shape parameters of beta distributions are realized by the function `nlm` numerically, additional arguments for `nlm` are allowed by `...`.

**Value**

This function will return an updated object of class `BetaMixture`. 
fitNULL

Fit the NULL component of a three-beta mixture model for functional gene associations

Description

The function performs permutations to the input rich phenotyping screens, and subsequently fit a beta distribution to the densities.

Usage

```r
fitNULL(object, nPerm=20, thetaNULL=c(alphaNULL=4, betaNULL=4),
sumMethod="median", permMethod="keepRep", verbose=TRUE, ...)```

Arguments

- `object` an object of S4 class BetaMixture.
- `nPerm` a positive numeric or integer value specifying the number of permutations.
- `thetaNULL` a list of numeric values giving the initial values for estimating the two shape parameters (see `dbeta` for more details).
- `sumMethod` a character value specifying how to summarize estimated parameters from multiple permutations. The current version only supports 'median' and 'mean'.
infer

Infer a posterior association network

Description

The function infers a posterior association network from beta-mixture modelling of functional associations computed from rich phenotyping screens.

Usage

infer(object, para=list(type='SNR', log=TRUE, sign=TRUE, cutoff=0), filter=FALSE, verbose=TRUE, ...)
Arguments

object an object of S4 class PAN.
para a list of parameters to perform inference (see details).
filter a logical value specifying whether or not to filter out genes without any significant association with all the other genes.
verbose a logical value to switch on (if TRUE) or off if FALSE detailed run-time message.
... not in use, only for further extension.

Details

This function employs different edge weights to infer a posterior association network (see edgeWeight for more details). Multiple parameters are provided for the user to specify the network:

'type' - a character value giving the type of edge weights: signal-to-noise ratio ('SNR'), posterior probability ratio ('PPR') or posterior probability ('PP').
'log' - a logical value specifying whether or not to compute logarithms for edge weights.
'sign' - a logical value specifying whether a signed graph should be inferred. It is only valid when type='SNR'.
'cutoff' - a numeric value giving the threshold to tell the significance of an edge.

Value

This function will return an object of class PAN with inferred PAN updated in slot 'graph'.

Author(s)

Xin Wang <xw264@cam.ac.uk>

References


See Also

edgeWeight

Examples

data(bm, package="PANR")
pan<-new("PAN", bm1=bm1)
pan<-infer(pan, para=list(type="SNR", log=TRUE, sign=TRUE, cutoff=log(5)),
filter=FALSE, verbose=TRUE)
Description

The function translate p-values to Signal-to-Noise Ratios based on the fitted mixture model.

Usage

p2SNR(object, pval, ...)

Arguments

object  
an object of S4 class BetaMixture.
pval  
a numeric or integer specifying the p-value to translate
...
  
not in use, only for further extension.

Value

The function will return a data frame including p-values, lower and upper quantiles and corresponding Signal-to-Noise Ratios.

Author(s)

Xin Wang <xw264@cam.ac.uk>

References


See Also

p2SNR

Examples

## Not run:
data(Bakal2007)
bm1<-new("BetaMixture", pheno=Bakal2007, model="global", order=1)
bm1<-fitNULL(bm1, nPerm=10, thetaNULL=c(alphaNULL=4, betaNULL=4), sumMethod="median", permMethod="all", verbose=TRUE)
bm1<-fitBM(bm1, para=list(zInit=NULL, thetaInit=c(alphaNeg=2, betaNeg=4, alphaNULL=bm1@result$fitNULL$thetaNULL["alphaNULL"], betaNULL=bm1@result$fitNULL$thetaNULL["betaNULL"]), alphaPos=4, betaPos=2), gamma=NULL), ctrl=list(fitNULL=FALSE, tol=1e-1, verbose=TRUE, gradtol=1e-3)
p2SNR(bm1, pval=0.01)
## End(Not run)
An S4 class for inferring a posterior association network

Description

This S4 class includes methods to infer posterior association networks and enriched modules of functional gene interactions from rich phenotyping screens.

Objects from the Class

Objects of class PAN can be created from `new("PAN", bm1,bm2)` (see the example below for details).

Slots

- `bm1`: an object of S4 class BetaMixture, which models the first-order similarities between genes (see `BetaMixture`).
- `bm2`: an object of S4 class BetaMixture, which models the second-order similarities between genes (modularity).
- `edgeWt`: a weighted adjacency matrix computed from the posterior probabilities for gene associations to belong to mixture components (see `edgeWeight`).
- `engine`: the graphics visualization engine for PAN.
- `graph`: a weighted adjacency matrix with edge weights satisfying certain constraints specified by the user (see `infer`).
- `modules`: a list summarizing inferred enriched functional gene modules (see `pvclustModule`.
- `iPAN`: an igraph object for storing the inferred PAN.
- `legend`: a list of legends for built PAN graph.
- `summary`: a list of summary information for available results.

Methods

An overview of methods (More detailed introduction can be found in help for each specific function.):

- `edgeWeight` compute edge weights by signal-to-noise ratio, posterior odd or posterior probabilities (more details in `edgeWeight`).
- `infer` infer a posterior association network given the beta-mixture model(s) fitted to first- and/or second-order similarities (more details in `infer`).
- `pvclustModule` search significantly enriched functional gene modules by hierarchical clustering with bootstrap resampling based on the package pvclust (more details in `pvclustModule`).
- `exportPAN` export the inferred PAN or modules to file(s) in a variety of formats (more details in `exportPAN`).
- `sigModules` retrieve significant gene modules that satisfy the given p-value cutoff and module size range (more details in `sigModules`).
- `viewNestedModules` view a nested structure for gene modules searched by hierarchical clustering (more details in `viewNestedModules`).
- `viewPAN` view the inferred PAN or modules in igraph or RedeR (more details in `viewPAN`).
buildPAN  build a PAN graph for visualization in igraph or RedeR (more details in viewPAN).
viewLegend  View the legends for the graph built for PAN.
summarize  summarize results including input data and parameters, inferred graph and modules.

Author(s)
Xin Wang <xw264@cam.ac.uk>

References

See Also

edgeWeight infer pvclustModule exportPAN sigModules viewPAN viewNestedModules summarize

Examples

## Not run:
data(bm, package="PANR")
##create an object of 'PAN'
pan<-new("PAN", bm=bm)
##infer a PAN
pan<-infer(pan, para=list(type="SNR", log=TRUE, sign=TRUE, cutoff=log(5)),
filter=FALSE, verbose=TRUE)
##build a PAN graph for RedeR, hide negative edges
##using colors scaled based on the clustering results from Bakal et al. 2007
data(Bakal2007Cluster)
pan<-buildPAN(pan, engine="RedeR", para=list(nodeColor=nodeColor, hideNeg=TRUE))
##view PAN in RedeR
library(RedeR)
viewPAN(pan, what="graph")
##print a summary of results
summarize(pan, "ALL")

## End(Not run)

## Search enriched functional gene modules by pvclust

pvclustModule  

Search enriched functional gene modules by pvclust

Description
The function employs the R package pvclust to search significant functional gene modules using hierarchical clustering with bootstrap resampling.

Usage

pvclustModule(object, nboot=1000, metric="cosine", hclustMethod="average",
filter=TRUE, verbose=TRUE, ...)
Arguments

object: an object of S4 class PAN.

nboot: a numeric value giving the number of bootstraps for pvclust.

metric: a character value specifying which distance metric to use for the hierarchical clustering: 'correlation', 'cosine', 'abscor' or those allowed by the argument 'method' in dist.

hclustMethod: the agglomerative method used in hierarchical clustering: 'average', 'ward', 'single', 'complete', 'mcquitty', 'median' or 'centroid' (see the argument method in hclust for more details).

filter: a logical value specifying whether or not to filter out screening data of genes without significant associations with all the other genes.

verbose: a logical value to switch on (if TRUE) or off if FALSE detailed run-time message.

... all the other arguments accepted by the function pvclust.

Details

This function performs hierarchical clustering with bootstrap resampling to quantify significance of gene clusters (modules) based on the package pvclust.

Value

This function will return an object of class PAN with inferred gene modules (modules$clusters) and corresponding p-values (modules$pval) updated in slot 'modules'.

Author(s)

Xin Wang <xw264@cam.ac.uk>

References


Examples

```r
## Not run:
data(bm, package="PANR")
pan<-new("PAN", bm1=bm1)
pan<-infer(pan, para=list(type="SNR", log=TRUE, sign=TRUE, cutoff=log(5)),
filter=FALSE, verbose=TRUE)
data(Bakal2007Cluster, package="PANR")
pan<-buildPAN(pan, engine="igraph", para=list(nodeColor=nodeColor,
hideNeg=TRUE), verbose=TRUE)
##need pvclust to search modules
library(pvclust)
pan<-pvclustModule(pan, nboot=1000, metric="cosine",
hclustMethod="average", filter=TRUE, verbose=TRUE, r=c(5:12/7))
## End(Not run)
```
sigModules

Retrieve ids for significant gene modules searched by pvclust

Description

The function retrieve ids for significant gene modules that satisfy the given p-value cutoff and module size range.

Usage

```
sigModules(object, pValCutoff=0.01, minSize=3, maxSize=100, sortby="size", decreasing=FALSE, ...)```

Arguments

- **object**: an object of S4 class PAN.
- **pValCutoff**: a numeric value specifying the p-value cutoff to tell the significance of a gene module.
- **minSize**: a numeric or integer value giving the minimal size of gene modules.
- **maxSize**: a numeric or integer value giving the maximal size of gene modules.
- **sortby**: a character value specifying how to sort the list of gene modules: by ‘size’ (module size) or ‘pval’ (pvclust p-value).
- **decreasing**: a logical value specifying whether or not the sorting will be conducted decreasingly.
- **...**: not in use, but only for further extension.

Details

This function facilitates the user to retrieve significant gene modules found by pvclust and obtain their ids, which can be subsequently used for visualization (see `viewPAN` for details).

Value

a numeric vector of ids for significant gene modules

Author(s)

Xin Wang <xw264@cam.ac.uk>

References


Examples

```r
## Not run:
data(bm, package="PANR")
pan<-new("PAN", bm1=bm1)
pan<-infer(pan, para=list(type="SNR", log=TRUE, sign=TRUE, cutoff=log(5)), filter=FALSE, verbose=TRUE)
data(Bakal2007Cluster, package="PANR")
pan<-buildPAN(pan, engine="igraph", para=list(nodeColor=nodeColor, hideNeg=TRUE), verbose=TRUE)
##need pvclust to search modules
library(pvclust)
pan<-pvclustModule(pan, nboot=1000, metric="cosine", hclustMethod="average", filter=TRUE, verbose=TRUE, r=c(5:12/7))
inds<sigModules(pan, pValCutoff=0.01, minSize=5, maxSize=100, sortby="size", decreasing=FALSE)
pan@modules$clusters[inds]
## End(Not run)
```

---

**SNR2p**

*Translate p-values to Signal-to-Noise Ratios*

**Description**

The function translate Signal-to-Noise Ratios to p-values based on the fitted mixture model.

**Usage**

```r
SNR2p(object, SNR, ...)
```

**Arguments**

- `object` an object of S4 class `BetaMixture`.
- `SNR` a numeric or integer specifying the SNR to translate
- `...` not in use, only for further extension.

**Value**

The function will return a data frame including SNRs, lower and upper quantiles and corresponding p-values.

**Author(s)**

Xin Wang `<xw264@cam.ac.uk>`

**References**

summarize

See Also

SNR2p

Examples

```r
## Not run:
data(Bakal2007)
bm1<-new("BetaMixture", pheno=Bakal2007, model="global", order=1)
bm1<-fitNULL(bm1, nPerm=10, thetaNULL=c(alphaNULL=4, betaNULL=4),
         sumMethod="median", permMethod="all", verbose=TRUE)
bm1<-fitBM(bm1, para=list(zInit=NULL, thetaInit=c(alphaNeg=2, betaNeg=4,
         alphaNULL=bm1@result$fitNULL$thetaNULL["alphaNULL"],
         betaNULL=bm1@result$fitNULL$thetaNULL["betaNULL"],
         alphaPos=4, betaPos=2, gamma=NULL),
         ctrl=list(fitNULL=FALSE, tol=1e-1, verbose=TRUE, gradtol=1e-3))
SNR2p(bm1, SNR=10)
## End(Not run)
```

summarize

`summarize` is a function that helps print a summary of an object of S4 class `BetaMixture` or `PAN`.

**Description**

The function helps print a summary of an object of S4 class `BetaMixture` or `PAN`.

**Usage**

```r
summarize(object, what='ALL', ...)
```

**Arguments**

- `object`: an object of S4 class `BetaMixture` or `PAN`.
- `what`: a character value specifying what to print (see details).
  - `'input'`: prints a summary of input parameters.
  - `'fitNULL'`: prints a summary of fitting results for the NULL distribution.
  - `'fitBM'`: prints a summary of fitting results for the beta-mixture model.
  - `'ALL'`: prints all above messages.
- `...`: not in use, only for further extension.

**Details**

This function prints a summary of an object of `BetaMixture` or `PAN`. The function is also called by S4 method `show`, which prints only a short message about the input parameters and data.

For an object of class `BetaMixture`:

- If `what='input'`, the function prints a summary of input parameters.
- If `what='fitNULL'`, the function prints a summary of fitting results for the NULL distribution.
- If `what='fitBM'`, the function prints a summary of fitting results for the beta-mixture model.
- If `what='ALL'`, all above messages will be printed.

For an object of class `PAN`:

- If `what='input'`, the function prints a summary of input objects of class `BetaMixture`.
- If `what='graph'`, the function prints a summary of inferred posterior association network.
- If `what='module'`, the function prints a summary of functional gene modules.
- If `what='ALL'`, all above messages will be printed.
Author(s)

Xin Wang <xw264@cam.ac.uk>

References


Examples

data(bm)
summarize(bm1, what='ALL')

Description

This function facilitate the user to view and check the fitting results of NULL and beta-mixture distributions.

Usage

view(object, what="fitBM", ...)

Arguments

object an object of S4 class BetaMixture.
what a character value specifying to show the fitting results of NULL (if what='fitNULL') or beta-mixture model (if what='fitBM').
... not in use, only for further extension.

Details

The function help the user to view and check the fitting of NULL and three beta-mixture model to permuted and real phenotyping screens, respectively. For either fitting, a histogram of association scores and the fitted beta distribution(s) will be plotted. For beta-mixture fitting, the integrated probability density function for the mixed distribution will also be plotted.

Author(s)

Xin Wang <xw264@cam.ac.uk>

References

viewLegend

See Also

fitNULL fitBM

Examples

data(bm)
view(bm1, what="fitNULL")
view(bm1, what="fitBM")

viewLegend

View the legends for the graph built for PAN

Description

This function helps the user to view legends for the built PAN graph.

Usage

viewLegend(object, what="nodeColor", ...)

Arguments

object an object of S4 class PAN.
what a character value specifying which legend to view: ‘nodeColor’, ‘nodeSize’ or ‘edgeWidth’.
... not in use, but only for further extension.

Author(s)

Xin Wang <xw264@cam.ac.uk>

Examples

## Not run:
data(bm, package="PANR")
pan<-new("PAN", bm1=bm)
pan<-infer(pan, para=list(type="SNR", log=TRUE, sign=TRUE, cutoff=log(5)),
filter=FALSE, verbose=TRUE)
data(Bakal2007Cluster, package="PANR")
pan<-buildPAN(pan, engine="igraph", para=list(nodeColor=nodeColor, hideNeg=TRUE), verbose=TRUE)
viewLegend(pan, "edgeWidth")

## End(Not run)
### Description

The function displays the nested enriched functional gene modules found by pvclust in a powerful graphic visualization software RedeR.

### Usage

```r
viewNestedModules(object, pValCutoff=0.01, minSize=3, maxSize=100, verbose=TRUE, ...)
```

### Arguments

- **object**: an object of S4 class PAN.
- **pValCutoff**: a numeric value specifying the p-value cutoff to tell the significance of a gene module.
- **minSize**: a numeric or integer value giving the minimal size of gene modules.
- **maxSize**: a numeric or integer value giving the maximal size of gene modules.
- **verbose**: a logical value to switch on (if TRUE) or off if FALSE detailed run-time message.
- **...**: not in use, but only for further extension.

### Details

This function presents the searched enriched functional modules in RedeR - a bioconductor package for network visualization.

Please note that the user is expected to run `buildPAN` to build a graph and search modules using `pvclustModule` prior to visualize using this function.

Please also note that if ‘RedeR’ is selected as the graphics engine, it is suggested to manually organise the sizes and positions of containers (for nesting gene modules) run a dynamic layout to obtain the best structure for the network.

### Author(s)

Xin Wang <xw264@cam.ac.uk>

### References


### See Also

`addGraph, nestNodes, viewPAN, buildPAN`
viewPAN

Show posterior association networks or modules in ‘igraph’ or ‘RedeR’

Description

The function display inferred posterior association networks or enriched functional gene modules in igraph or a powerful graphical visualization software RedeR.

Usage

viewPAN(object, what="graph", moduleID=1, layout="layout.fruchterman.reingold", verbose=TRUE, ...)

Arguments

object an object of S4 class PAN.
what a character value specifying which to show: ‘graph’ or ‘pvclustModule’.
moduleID a numeric or integer vector of modules to view (see details in sigModules). This argument will be applied only when what='pvclustModule'.
layout a character value specifying the layout method (see details in layout). This argument will be applied only when engine='igraph' when build a graph for PAN using buildPAN.
verbose a logical value to switch on (if TRUE) or off if FALSE detailed run-time message.
... not in use, but only for further extension.

Details

This function presents the inferred posterior association network or enriched functional modules in igraph or RedeR depending on the graphics engine used when building the graph for PAN (details in buildPAN).

Please note that when viewing a dense PAN in ‘igraph’, it could be very messy. Multiple modules can be viewed at the same time in ‘RedeR’ powered by its feature of ‘containers’, which are used to group gene modules. When view multiple modules in ‘igraph’,

Examples

## Not run:
data(bm, package="PANR")
pan<-new("PAN", bm1=bm1)
pan<-infer(pan, para=list(type="SNR", log=TRUE, sign=TRUE, cutoff=log(5)),
filter=FALSE, verbose=TRUE)
data(Bakal2007Cluster, package="PANR")
pan<-buildPAN(pan, engine="igraph", para=list(nodeColor=nodeColor,
hideNeg=TRUE), verbose=TRUE)
##need pvclust to search modules
library(pvclust)
pan<-pvclustModule(pan, nboot=10000, metric="cosine2",
hclustMethod="average", filter=TRUE, verbose=TRUE, r=c(5:12/7))
viewNestedModules(pan, pValCutoff=0.05, minSize=5, maxSize=100)

## End(Not run)
Author(s)

Xin Wang <xw264@cam.ac.uk>

References


See Also

buildPAN, layout

Examples

```r
## Not run:
data(bm, package="PANR")
pan<-new("PAN", bm1=bm1)
pan<-infer(pan, para=list(type="SNR", log=TRUE, sign=TRUE, cutoff=log(5)),
filter=FALSE, verbose=TRUE)
data(Bakal2007Cluster, package="PANR")
pan<-buildPAN(pan, engine="igraph", para=list(nodeColor=nodeColor,
hideNeg=TRUE), verbose=TRUE)
##view inferred PAN
viewPAN(pan, what='graph', layout="layout.fruchterman.reingold")

## End(Not run)
```
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