# Package 'clusterCons'

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Maintainer Dr. T. Ian Simpson <ian.simpson@ed.ac.uk></ian.simpson@ed.ac.uk>
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Description Functions for calculation of robustness measures for clusters and cluster membership based on generating consensus matrices from bootstrapped clustering experiments in which a random proportion of rows of the data set are used in each individual clustering. This allows the user to prioritise clusters and the members of clusters based on their consistency in this regime. The functions allow the user to select several algorithms to use in the resampling scheme and with any of the parameters that the algorithm would normally take. See Simpson, T. I., Armstrong, J. D. & Jarman, A. P. (2010) <doi:10.1186 1471-2105-11-590=""> and Monti, S., Tamayo, P., Mesirov, J. &amp; Golub, T. (2003) <doi:10.1023 a:1023949509487=""></doi:10.1023></doi:10.1186>
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<b>Author</b> Dr. T. Ian Simpson [aut, cre, cph] ( <a href="https://orcid.org/0000-0003-0495-7187">https://orcid.org/0000-0003-0495-7187</a> )
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# Description

clusterCons is a package containing functions that generate robustness measures for clusters and cluster membership based on generating consensus matrices from bootstrapped clustering experiments in which a random proportion of rows of the data set are used in each individual clustering. This allows the user to prioritise clusters and the members of clusters based on their consistency in this regime. The functions allow the user to select several algorithms to use in the re-sampling scheme and with any of the parameters that the algorithm would normally take.

# **Details**

Package: clusterCons
Type: Package
Version: 1.0
Date: 2010-10-12
License: GPL
LazyLoad: yes

Depends: methods, cluster, lattice, RColor Brewer, grid, apcluster

Extends: cluster Suggests: latticeExtra clusterCons-package 3

The user should first prepare an entirely numeric data. frame in which the conditions to be clustered are the column names and the unique ids of the entities are the row names. Compatibility of the resulting data.fram can be checked by using the data\_check function.

#### Functions to run the consensus clustering and retrieve robustness information

cluscomp - generate consensus matrices from re-sampled clustering experiments with the option of multiple algorithms and parameters

clrob - calculate the robustness of the clusters from the consensus matrix memrob - calculate the cluster membership robustness from the consensus matrix

# Internal functions to call the individual clustering algorithms

agnes\_clmem - wrapper for the agnes function of package cluster diana\_clmem - wrapper for the diana function of package cluster hclust\_clmem - wrapper for the hclust function of package cluster kmeans\_clmem - wrapper for the kmeans function of package cluster pam\_clmem - wrapper for the pam function of package cluster apcluster\_clmem - wrapper for the apclusterK function of package apcluster

#### **Functions to calculate AUC related metrics**

auc - calculates the area under the curve for a series of clustering experiments with the same cluster number

aucs - calculates the areas under the curves of a series of clustering experiments over a range of cluster numbers

deltak - calculates the change in the area under the curve

#### Functions to check data and object validity

data\_check - check that the provided data.frame is formatted correctly expSetProcess - extracts the data set from an object of class expressionSet validConsMatrixObject - check the validity of a consmatrix object validMergeMatrixObject - check the validity of a mergematrix object validMemRobListObject - check the validity of a membership robustness list object validMemRobMatrixObject - check the validity of a membership robustness matrix object validAUCObject - check the validity of an "auc" class object validDkObject - check the validity of an "dk" class object

# Functions to plot out performance curves

aucplot - plot area under the curve (AUC) plots from consensus clustering results dkplot - plot change in AUC by cluster number (delta-K plot) expressionPlot - plot the original data partitioned by cluster membership membBoxPlot - plot a box and whisker plot of the membership robustness for each cluster

## **Keywords**

cluster

#### See Also

cluster, lattice, apcluster

# **Examples**

```
#load data data(sim_profile);

#perform consensus clustering cmr <- cluscomp(sim_profile,algo=list('agnes','pam','kmeans'),clmin=2,clmax=7,rep=10,me
#see the consensus and merge matrices summary(cmr);

#fetch the cluster robustness for agnes consensus clustering with k=3 clrob(cmr$e1_agnes_k3);

#show the membership robustness for cluster 1 memrob(cmr$e1_agnes_k3)$cluster1

#show the same, but for the merge against the k=3 agnes clustering structure #note we provide the
reference matrix (which is the original cluster membership for agnes where k=3) clrob(cmr$merge_k3,cmr$e1_agnes_k3@rm
memrob(cmr$merge_k3,cmr$e1_agnes_k3@rm)$cluster1;

#calculate the AUCs acs <- aucs(cmr);

#plot the AUC curves aucplot(acs);

#calculate the delta-K curves dkplot(dks);

#plot the delta-K curves dkplot(dks);

#plot the expression profiles expressionPlot(sim_profile,cmr$e1_agnes_k3);

#plot the bwplot of membership robustness for the same membBoxPlot(memrob(cmr$e1_agnes_k3));
```

# Author(s)

Dr. T. Ian Simpson <ian.simpson@ed.ac.uk>

# References

Merged consensus clustering to assess and improve class discovery with microarray data. Simpson TI, Armstrong JD and Jarman AP. BMC Bioinformatics 2010, 11:590.

Consensus clustering: A resampling-based method for class discovery and visualization of gene expression microarray data. Monti, S., Tamayo, P., Mesirov, J. and Golub, T. Machine Learning, 52, July 2003.

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auc

Calculate area under the curve statistics

# **Description**

These functions calculate the area under the curve (AUC) for cumulative density functions of a consensus matrix. The function auc operates on an indvidual consensus matrix whereas aucs operates on an entire cluscomp analysis result as described below.

# Usage

```
auc(x)
aucs(x)
```

## **Arguments**

Х

For auc(x), provide a numeric square data matrix such as an individual consensus matrix. For aucs(x) provide a list of "consmatrix" class objects (see consmatrix-class for details) such as those produced directly by the cluscomp function

The functions will not allow any missing values (NAs).

# Value

```
auc(x) returns an individual AUC value.
```

aucs(x) returns a data.frame with the following variables.

```
k cluster number as a factora algorithm identifier as a factoraucs the AUC value
```

# Author(s)

```
Dr. T. Ian Simpson <ian.simpson@ed.ac.uk>
```

#### References

Merged consensus clustering to assess and improve class discovery with microarray data. Simpson TI, Armstrong JD and Jarman AP. BMC Bioinformatics 2010, 11:590.

#### See Also

```
consmatrix-class
```

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## **Examples**

```
#load up a test cluscomp result
data('testcmr');

#look at the result structure
summary(testcmr);

#calculate an individual AUC value for a consensus matrix
ac <- auc(testcmr$e1_kmeans_k2@cm);

#calculate all of the AUC values from the \code{cluscomp} result for algorithm 'kmeans'
kmeanscmr <- testcmr[grep('kmeans',names(testcmr))];
acs <- aucs(kmeanscmr);</pre>
```

auc-class

Class "auc"

# **Description**

Objects of class 'auc' contain a data.frame which have three variables k, a and auc as described in the aucs function description. This class simply holds the result from a call to aucs.

# **Objects from the Class**

Objects can be created by calls of the form new("auc", ...), although they are normally generated internally by the aucs function.

# Author(s)

Dr. T. Ian Simpson <ian.simpson@ed.ac.uk>

# References

Merged consensus clustering to assess and improve class discovery with microarray data. Simpson TI, Armstrong JD and Jarman AP. BMC Bioinformatics 2010, 11:590.

# See Also

Also see the aucs function.

```
showClass("auc")
```

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aucplot

Generate an area under the curve plot using lattice graphics

# **Description**

This function uses the lattice function xyplot to generate an AUC plot from a valid "auc" class object (see auc-class).

# Usage

```
aucplot(x)
```

## **Arguments**

Х

a valid "auc" class object (see auc-class), normally generated by the aucs function.

# Value

No return value, called for side effects

# Author(s)

```
Dr. T. Ian Simpson <ian.simpson@ed.ac.uk>
```

## References

Merged consensus clustering to assess and improve class discovery with microarray data. Simpson TI, Armstrong JD and Jarman AP. BMC Bioinformatics 2010, 11:590.

# See Also

```
consmatrix-class
```

```
#load up a test cluscomp result
data('testcmr');

#look at the result structure
summary(testcmr);

#calculate all of the AUC values from the \code{cluscomp} result for algorithm 'kmeans'
kmeanscmr <- testcmr[grep('kmeans',names(testcmr))];
acs <- aucs(kmeanscmr);

#plot the AUC curve
aucplot(acs);</pre>
```

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checks

Functions to check the integrity of various objects

# **Description**

These methods are mainly internal although the user may like to check their original data using data\_check before they perform consensus clustering experiments.

# Usage

```
data_check(x)
validConsMatrixObject(object)
validMemRobListObject(object)
validMemRobMatrixObject(object)
validMergeMatrixObject(object)
validAUCObject(object)
validDkObject(object)
```

# **Arguments**

x The data.frame object to be checked prior to using with the cluscomp function.

The object to be checked with the suitable function by type. These are used internally by several of the functions in the package.

#### Value

returns TRUE if check is passed or an error message if it is not

## Author(s)

```
Dr. T. Ian Simpson <i an.simpson@ed.ac.uk>
```

# References

Merged consensus clustering to assess and improve class discovery with microarray data. Simpson TI, Armstrong JD and Jarman AP. BMC Bioinformatics 2010, 11:590.

```
#load data
data(sim_profile);

#check if this can be used by cluscomp
data_check(sim_profile);

#perform a clusomp run
cmr <- cluscomp(sim_profile,clmin=2,clmax=2,rep=10);</pre>
```

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#check one of the consensus matrices
validConsMatrixObject(cmr\$e1\_kmeans\_k2)

clrob

Calculate the cluster robustness from consensus clustering results

# **Description**

This function calculates the cluster robustness from a consmatrix or mergematrix class object.

#### **Usage**

clrob(x,rm)

# **Arguments**

either a consmatrix or mergematrix object.

rm

Х

(optional) if a mergematrix object is passed then you must provide a reference clustering structure to calculate cluster robustness against. These structures are stored with every consmatrix object in the 'rm' slot. You would normally select a reference matrix for a cluster number matching that of the mergematrix (see example below).

# Value

Returns a data.frame of the cluster robustness values indexed by cluster number.

# Author(s)

Dr. T. Ian Simpson <ian.simpson@ed.ac.uk>

#### References

Merged consensus clustering to assess and improve class discovery with microarray data. Simpson TI, Armstrong JD and Jarman AP. BMC Bioinformatics 2010, 11:590.

# See Also

Also see cluscomp, consmatrix and mergematrix.

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# **Examples**

```
#load cmr (consensus clustering result produced by cluscomp)
data(testcmr);

#calculate the cluster robustness of the consensus matrix for pam where k=4
clrob(testcmr$e1_kmeans_k4);

#calculate the cluster robustness of the merge matrix in reference
#to the clustering structrure of pam where k=4
clrob(testcmr$merge_k4,testcmr$e1_kmeans_k4@rm);
```

cluscomp

Perform consensus clustering with the option of using multiple algorithms and parameters and merging

# Description

Calculates an NxN consensus matrix for each clustering experiment performed where each entry has a value between 0 (never observed) and 1 (always observed)

When running with more than one algorithm or with the same algorithm and multiple conditions a consensus matrix will be generated for each. These can optionally be merged into a mergematrix by cluster number by setting merge=1.

## Usage

```
cluscomp(
    x,
    diss=FALSE,
    algorithms = list("kmeans"),
    alparams = list(),
    alweights = list(),
    clmin = 2,
    clmax = 10,
    prop = 0.8,
    reps = 50,
    merge = 0
)
```

# Arguments

Х

data.frame of numerical data with conditions as the column names and unique ids as the row names. All variables must be numeric. Missing values(NAs) are not allowed. Optionally you can pass a distance matrix directly, in which case you must ensure that the distance matrix is a data.frame and that the row and column names match each other (as the distance matrix is a pair-wise distance calculation).

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diss	set to TRUE if you are providing a distance matrix, default is FALSE
algorithms	list of algorithm names which can be drawn from 'agnes', 'diana', 'pam', 'kmeans' or 'hclust'. The user can also write a simple wrapper for any other clustering method (see details)
alparams	list of algorithm paramter lists using the same specification as for the individual algorithm called (see details)
alweights	list of integer weights for each algorithm (only used when merging consensus results between algorithms)
clmin	integer for the smallest cluster number to consider
clmax	integer for the largest cluster number to consider
prop	numeric for the proportion of rows to sample during the process. Must be between $0 \ \mathrm{and} \ 1$
reps	integer for the number of iterations to perform per clustering
merge	an integer indicating whether you also want the merged matrices (1) or just the consensus ones (0), accepts only 1 or 0.

#### **Details**

cluscomp is an implementation of a consensus clustering methodology first proposed by Monti et al. (2003) in which the connectivity between any two members of a data matrix is tested by resampling statistics. The principle is that by only sampling a random proportion of rows in the data matrix and performing many clustering experiments we can capture information about the robustness of the clusters identified by the full unsampled clustering result.

For each re-sampling experiment run a zero square matrix is created with identical rows and columns matching the unique ids of the rows of the data matrix, this matrix is called the connectivity matrix. A second identically sized matrix is created to count the number of times that any pair of row ids are called in any one re-sampled clustering. This matrix is called the identity matrix. For each iteration within the experiment the rows sampled are recorded in the identity matrix and then the co-occurrence of all pairs are recorded in the connectivity matrix. These values are incremented for each iteration until finally a conensensus matrix is generated by dividing the connectivity matrix by the identity matrix.

The consensus matrix is the raw output from cluscomp implemented as a class consmatrix. If the user has specified to return a merged matrix in addition to the consensus matrices then for each clustering with the same k (cluster number value) an object of class mergematrix is also returned in the list which is identical to a consmatrix with the exception that the 'cm' slot is occupied by the merged matrix (a weighted average of all the consensus matrices for the cluster number matched consensus matrices) and there is no reference matrix slot (as there is no reference clustering for the merge). The user should instead call the memrob function using the merge matrix and providing a reference matrix from one of the cluster number matched consmatrix objects from which the merge was generated. This provides a way to quantify the difference between single and multi-algorithm resampling schemes.

#### Value

a list of objects of class consmatrix and (if merge specified) mergematrix. See consmatrix and mergematrix for details.

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## Author(s)

Dr. T. Ian Simpson <ian.simpson@ed.ac.uk>

#### References

Merged consensus clustering to assess and improve class discovery with microarray data. Simpson TI, Armstrong JD and Jarman AP. BMC Bioinformatics 2010, 11:590.

Consensus clustering: A resampling-based method for class discovery and visualization of gene expression microarray data. Monti, S., Tamayo, P., Mesirov, J. and Golub, T. Machine Learning, 52, July 2003.

#### See Also

cluster,clrob,memrob

```
#load test data
data(sim_profile);
#perform a group of re-sampling clustering experiments accepting default parameters
#for the clustering algorithms
cmr <- cluscomp(</pre>
sim_profile,
algorithms=list('kmeans','pam'),
merge=1,
clmin=2,
clmax=5,
reps=5
#display resulting matrices contained in the consensus result list
summary(cmr);
#display the cluster robusteness for the kmeans k=4 consensus matrix
clrob(cmr$e2_pam_k4);
#plot a heatmap of the consensus matrix, note you access the cluster matrix object
#through the cm slot
#heatmap(cmr$e2_pam_k4@cm);
#display the membership robustness for kmeans k=4 cluster 1
memrob(cmr$e2_pam_k4)$cluster1;
#merged consensus example
#data(testcmr);
#calculate the membership robustness for the merge matrix when cluster number k=4,
#in reference to the pam scaffold. (see memrob for more details).
```

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```
#mr <- memrob(testcmr$merge_k4,testcmr$e1_kmeans_k4@rm);
#show the membership robustness for cluster 1
#mr$cluster1;</pre>
```

consmatrix-class

Class "consmatrix"

# **Description**

Objects of class 'consmatrix' are created to hold the results of a consensus clustering experiment along with the necessary ancillary data to allow the subsequent downstream calculations such as cluster and membership robustness. In addition the object holds the original call made when running cluscomp.

# **Objects from the Class**

Objects can be created by calls of the form new("consmatrix", ...), but are normally created internally by the cluscomp function to store consensus matrices and their associated meta-data.

#### Slots

```
cm: Object of class "matrix" - the consensus matrix itself
```

rm: Object of class "data.frame" - the cluster membership of the full (i.e. not consensus) clustering result when the current algorith is called with the same algorithm parameters as the consensus clustering run. This is needed to be able to work with merge matrices that need a clustering structure on which to operate to produce cluster and membership robustness values.

a: Object of class "character" - the clustering algorithm name

k: Object of class "numeric" - the cluster number (k) used

call: Object of class "call" - the original parameters passed to cluscomp for provenance and reproducibility

#### Author(s)

Dr. T. Ian Simpson <ian.simpson@ed.ac.uk>

# References

Merged consensus clustering to assess and improve class discovery with microarray data. Simpson TI, Armstrong JD and Jarman AP. BMC Bioinformatics 2010, 11:590.

#### See Also

See Also cluscomp

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#### **Examples**

```
showClass("consmatrix");
#you can access the slots in useful ways
#load a cmr
data(testcmr);
#get a consensus clustering matrix via the 'cm' slot
cm <- testcmr$e1_kmeans_k4@cm;</pre>
#this can be used as a distance matrix, e.g. for a heatmap
heatmap(cm);
#or as a new distance matrix
dm <- data.frame(cm) #first convert to a data.frame</pre>
#make sure names are the same for rows and columns
names(dm) <- row.names(dm);</pre>
#you need to explicitly tell cluscomp that you are passing a distance matrix
cmr2 <- cluscomp(dm,diss=TRUE,clmin=2,clmax=4,rep=2);</pre>
#for merge consensus clustering you take advantage of the reference matrix (rm) slot
#cluster robustness for agnes with cluster number (k) = 3
clrob(testcmr$merge_k3, testcmr$e1_kmeans_k3@rm);
#membership robustness for cluster 1
memrob(testcmr$merge_k3,testcmr$e1_kmeans_k3@rm)$cluster1;
```

data

Data sets for the clusterCons package

# **Description**

These data sets are used by the examples in the package function descriptions and allow the user to explore the functionality of the package

#### Usage

```
data(golub);
data(sim_class);
data(sim_profile);
data(testcmr);
```

#### **Format**

golub: data.frame of gene expression values for 999 genes for 38 leukemia patients (1-27) ALL and (28-38) AML. sim\_class: data.frame of 200 simulated gene expression values for 30 conditions where there are 4 discrete classes of expression profile, for testing clustering with the transposed data (clustering by column). sim\_profile: data.frame of 120 simulated gene expression values for

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4 conditions where there are 4 discrete classes of expression profile, for testing general clustering (clustering by row). testcmr: list of consensus and merge matrix results from a cluscomp run (see consmatrix-class and mergematrix-class).

# Author(s)

Dr. T. Ian Simpson <ian.simpson@ed.ac.uk>

#### References

Merged consensus clustering to assess and improve class discovery with microarray data. Simpson TI, Armstrong JD and Jarman AP. BMC Bioinformatics 2010, 11:590.

Molecular classification of cancer: class discovery and class prediction by gene expression monitoring. Golub, TR and Slonim, DK and Tamayo, P and Huard, C and Gaasenbeek, M and Mesirov, JP and Coller, H and Loh, ML and Downing, JR and Caligiuri, MA and Bloomfield, CD and Lander, ES. Science 1999, 286:531-537

# **Examples**

```
#cluster by class
data(sim_class);
cutree(agnes(t(sim_class)),4);
#cluster by profile
data(sim_profile);
cutree(agnes(sim_profile),4);
```

deltak

Function to calculate the change in the area under the curve (AUC) across a range of cluster number values

# **Description**

This function takes an "auc" class object and calculates the difference in AUC value by cluster number (called delta-K). Peaks in delta-K coincide with the cluster numbers that are most robust and provide estimates for the optimal cluster number.

#### Usage

```
deltak(x)
```

# **Arguments**

x a valid "auc" class object, normally provided as a result from the aucs function.

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## Value

deltak(x) returns a data.frame with the following variables.

k cluster number as a factor
a algorithm identifier as a factor

dk the delta-K value

## Author(s)

Dr. T. Ian Simpson <ian.simpson@ed.ac.uk>

#### References

Merged consensus clustering to assess and improve class discovery with microarray data. Simpson TI, Armstrong JD and Jarman AP. BMC Bioinformatics 2010, 11:590.

## See Also

Also see the aucs function.

# **Examples**

```
#load a test cluscomp result set
data(testcmr)

#calculate all of the AUC values from the \code{cluscomp} result for algorithm 'kmeans'
kmeanscmr <- testcmr[grep('kmeans',names(testcmr))];
acs <- aucs(kmeanscmr);

#calculate the delta-K values
dks <- deltak(acs);</pre>
```

dk-class

Class "dk"

# Description

Objects of class 'dk' contain a data.frame which have three variables k, a and deltak as described in the deltak function description. This class simply holds the result from a call to deltak.

## **Objects from the Class**

Objects can be created by calls of the form new("dk", ...), although they are normally generated internally by the deltak function.

# Author(s)

```
Dr. T. Ian Simpson <ian.simpson@ed.ac.uk>
```

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## References

Merged consensus clustering to assess and improve class discovery with microarray data. Simpson TI, Armstrong JD and Jarman AP. BMC Bioinformatics 2010, 11:590.

# See Also

Also see the aucs function.

# **Examples**

```
showClass("dk")
```

dkplot

Generate a delta-K plot from area under the curve (AUC) values across multiple cluster numbers.

# **Description**

This function uses the lattice function xyplot to generate an delta-K plot from a valid "dk" class object (see dk-class).

# Usage

dkplot(x)

# Arguments

Х

a valid "dk" class object (see dk-class), normally generated by the deltak function.

# Value

No return value, called for side effects

## Author(s)

Dr. T. Ian Simpson <ian.simpson@ed.ac.uk>

# References

Merged consensus clustering to assess and improve class discovery with microarray data. Simpson TI, Armstrong JD and Jarman AP. BMC Bioinformatics 2010, 11:590.

# See Also

```
consmatrix-class
```

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# **Examples**

```
#load up a test cluscomp result
data('testcmr');

#look at the result structure
summary(testcmr);

#calculate all of the AUC values from the \code{cluscomp} result for algorithm 'kmeans'
kmeanscmr <- testcmr[grep('kmeans',names(testcmr))];
acs <- aucs(kmeanscmr);

#calculate all of the delta-K values
dks <- deltak(acs);

#plot the delta-K curve
dkplot(dks);</pre>
```

expressionPlot

Generate a profile plot for the data partitioned by cluster membership.

# **Description**

This function uses the lattice function xyplot to generate a profile plot of the data values grouped by cluster in a multi-panel plot. The function takes as input the original data.frame() and a valid "consmatrix" class object (see consmatrix-class) by which to segregate the data.

# Usage

```
expressionPlot(x,cm);
```

# **Arguments**

x the original data.frame() object used in the clustering.
cm a valid "consmatrix" class object generated by the cluscomp function.

#### Value

No return value, called for side effects

# Author(s)

```
Dr. T. Ian Simpson <ian.simpson@ed.ac.uk>
```

# References

Merged consensus clustering to assess and improve class discovery with microarray data. Simpson TI, Armstrong JD and Jarman AP. BMC Bioinformatics 2010, 11:590.

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## See Also

```
consmatrix-class
```

# **Examples**

```
#load up the data set
data(sim_profile);
#load up an example cluscomp result with this data
data('testcmr');
#plot the expression profiles
expressionPlot(sim_profile,testcmr$e1_kmeans_k4);
```

expSetProcess

Internal function to extract the data from an expressionSet class object from the affy package for use with cluscomp

# **Description**

This is a convenience function that is used internally to allow the user to pass an expressionSet object from the microarray processing package 'affy' directly to the cluscomp function.

# Usage

```
expSetProcess(x)
```

# **Arguments**

Χ

An object of class expressionSet from the Bioconductor package 'affy'.

# Value

when called directly, returns a suitably labeled data.frame() object of the expressionSet expression values.

# Author(s)

```
Dr. T. Ian Simpson <ian.simpson@ed.ac.uk>
```

#### References

Merged consensus clustering to assess and improve class discovery with microarray data. Simpson TI, Armstrong JD and Jarman AP. BMC Bioinformatics 2010, 11:590.

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membBoxPlot	Generate a box and whisker plot of membership robustness for all clusters

# Description

This function uses the lattice function bwplot to generate a box and whisker plot of membership robustness from the result of a call to the memrob function.

# Usage

```
membBoxPlot(x)
```

# **Arguments**

v

the result of a call to the memrob function.

#### Value

No return value, called for side effects

# Author(s)

```
Dr. T. Ian Simpson <ian.simpson@ed.ac.uk>
```

#### References

Merged consensus clustering to assess and improve class discovery with microarray data. Simpson TI, Armstrong JD and Jarman AP. BMC Bioinformatics 2010, 11:590.

# See Also

```
memroblist-class,memrob
```

```
#load up a test cluscomp result
data('testcmr');

#calculate the membershpi robustness for one of the clustering results
mr <- memrob(testcmr$e1_kmeans_k5);

#plot the bwplot
membBoxPlot(mr);</pre>
```

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memrob	Calculate the membership robustness from consensus clustering results

# **Description**

This function calculates the membership robustness from a consmatrix or mergematrix class object.

# Usage

```
memrob(x,rm)
```

# Arguments

x either a consmatrix or mergematrix object.

rm (optional) if a mergematrix object is passed then you must provide a reference

clustering structure to calculate cluster robustness against. These structures are stored with every consmatrix object in the 'rm' slot. You would normally select a reference matrix for a cluster number matching that of the mergematrix (see

example below).

## Value

Returns a list of memroblist class objects, one for each cluster, and the full membership robustness matrix as a memrobmatrix class object.

# Author(s)

```
Dr. T. Ian Simpson <ian.simpson@ed.ac.uk>
```

# References

Merged consensus clustering to assess and improve class discovery with microarray data. Simpson TI, Armstrong JD and Jarman AP. BMC Bioinformatics 2010, 11:590.

# See Also

Also see cluscomp, consmatrix and mergematrix.

```
#load cmr (consensus clustering result produced by cluscomp)
data(testcmr);
#calculate the cluster robustness of the consensus matrix for pam where k=4
mr1 <- memrob(testcmr$e1_kmeans_k4);</pre>
```

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```
#show the membership robustness of cluster 1
mr1$cluster1;

#calculate the cluster robustness of the merge matrix in reference
#to the clustering structure of pam where k=4
mr2 <- memrob(testcmr$merge_k4,testcmr$e1_kmeans_k4@rm);

#plot a heatmap of the full membership robustness matrix
heatmap(mr2$resultmatrix@mrm)</pre>
```

memroblist-class

Class "memroblist"

## **Description**

Objects of class 'memroblist' are created to hold the membership robustness scores for the features (e.g. genes) of a cluster.

# **Objects from the Class**

Objects can be created by calls of the form new("memroblist", ...), although these objects are normally created internally by the memrob function.

## **Slots**

```
mrl: Object of class "data.frame" - the membership robustness list itself
```

# Author(s)

```
Dr. T. Ian Simpson <ian.simpson@ed.ac.uk>
```

#### References

Merged consensus clustering to assess and improve class discovery with microarray data. Simpson TI, Armstrong JD and Jarman AP. BMC Bioinformatics 2010, 11:590.

# See Also

Also see the memrob function/

```
showClass("memroblist")

#load a cmr
data(testcmr);

#calculate the membership robustness for agnes, k=4
mr <- memrob(testcmr$e2_agnes_k4);</pre>
```

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```
#get a membership robustness list
mrl <- mr$cluster1;</pre>
```

memrobmatrix-class

Class "memrobmatrix"

# **Description**

Objects of class 'memrobmatrix' hold the full membership robustness matrix generated from analysis of a consensus matrix. This includes the calculations of membership robustness for all features (e.g. genes) for each cluster. This can be useful as it allows you to see what conribution a particular feature (e.g. gene) is making to other clusters. This could resonably be thought of as a measure similar to 'fuzziness' i.e. partial cluster membership. If the value of the membership robustness for a feature is similar in many clusters then that is additional evidence that the feature is not easily placed in any cluster.

# **Objects from the Class**

Objects can be created by calls of the form new("memrobmatrix", ...), although they are usually generated internally by the memrob function.

#### **Slots**

mrm: Object of class "matrix" - this is the full membership robustness matrix itself and therefore has the same dimensions as the original data object used in the clustering

## Author(s)

```
Dr. T. Ian Simpson <ian.simpson@ed.ac.uk>
```

## References

Merged consensus clustering to assess and improve class discovery with microarray data. Simpson TI, Armstrong JD and Jarman AP. BMC Bioinformatics 2010, 11:590.

#### See Also

Also see the memrob function.

```
showClass("memrobmatrix")
#load cmr
data(testcmr);
#calculate membership robustness
mr <- memrob(testcmr$e1_kmeans_k3)</pre>
```

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```
#get the full membership robustness matrix (matrix itself held in slot 'mrm')
mrm <- mr$resultmatrix@mrm;</pre>
```

mergematrix-class

Class "mergematrix"

## **Description**

Objects of class 'mergematrix' hold the merge matrix in the same way that a consmatrix object holds a consensus matrix. As merge matrices only make sense in the context of the consensus clustering results that were used to generate them we do not store the meta-data for any one consensus clustering parameter set as we do for a 'consmatrix' object. All we need to identify the 'mergematrix' is the cluster number.

# **Objects from the Class**

Objects can be created by calls of the form new("mergematrix", ...), although they are normally generated by the cluscomp function when merge is specified.

#### Slots

```
cm: Object of class "matrix" - the merge matrix itselfk: Object of class "numeric" - the cluster number (k) value for which the merge was calculateda: Object of class "character" - always takes the value of 'merge' to identify it as a merge matrix
```

# Author(s)

```
Dr. T. Ian Simpson <ian.simpson@ed.ac.uk>
```

#### References

Merged consensus clustering to assess and improve class discovery with microarray data. Simpson TI, Armstrong JD and Jarman AP. BMC Bioinformatics 2010, 11:590.

# See Also

Also see the cluscomp function.

```
showClass("mergematrix")
#load the cmr
data(testcmr);
#get a merge matrix object
```

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```
mm <- testcmr$merge_k4;
#plot a heatmap of the merge matrix
heatmap(mm@cm);</pre>
```

wrappers

Functions to wrap command calls to clustering functions

#### **Description**

These are primarily internal functions called by cluscomp to execute clustering runs and are unlikely to be used directly. The wrappers are detailed in the algorithm.R file of the clusterCons package and the user can add their own wrappers to this to extend the number of algorithms supported. These five wrappers allow the user to specify the conditions under which the corresponding clustering algorithms are run and follow exactly the same specifications as the corresponding cluster functions (see agnes, pam, hclust, diana and kmeans).

# Usage

```
agnes_clmem(x, clnum, params = list())
pam_clmem(x, clnum, params = list())
hclust_clmem(x, clnum, params = list())
diana_clmem(x, clnum, params = list())
kmeans_clmem(x, clnum, params = list())
apcluster_clmem(x,clnum,params = list())
```

# **Arguments**

A data.frame of numerical values to be clustered which must pass the data\_check function. This function simply checks that there are no missing values, that all of the data is numeric and that row.names and column.names are unique. This is essential to ensure that individual rows (e.g. genes) and columns (e.g. experi-

mental conditions) can be identified consistently.

clnum The number of specified clusters. When using the cluscomp function, this will

be over-ridden by the cluster range specified using the parameters clmin and

clmax (see cluscomp for details).

params A list of key, value pairs specifying the parameters to pass to the clustering

algorithm. These follow the exact specification of the original functions in the

cluster package (see agnes, pam, hclust, diana and kmeans).

#### Value

cm

Returns a data.frame with row.names matching that of the data.

cluster membership identifier specifying the cluster into which the row has been classified

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## Author(s)

Dr. T. Ian Simpson <ian.simpson@ed.ac.uk>

## References

Merged consensus clustering to assess and improve class discovery with microarray data. Simpson TI, Armstrong JD and Jarman AP. BMC Bioinformatics 2010, 11:590.

# See Also

```
cluster, agnes, pam, hclust, diana, kmeans and apclusterK
```

```
#load some data
data(sim_profile);

#run a basic agnes clustering with 3 clusters
cm <- agnes_clmem(sim_profile,3);

#pass some more complex parameters
agnes_params = list(metric='manhattan',method='single');
cm <- agnes_clmem(sim_profile, 3,params=agnes_params);</pre>
```

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