Package ‘Mfuzz’

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R topics documented:

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This function extracts genes forming the alpha cores of soft clusters

Usage

```r
acore(eset, cl, min.acore = 0.5)
```

Arguments

- `eset`: object of the class `ExpressionSet`.
- `cl`: An object of class `flcust` as produced by `mfuzz`.
- `min.acore`: minimum membership values of gene belonging to the cluster core.

Value

The function produces an list of alpha cores including genes and their membership values for the corresponding cluster.

Author(s)

Matthias E. Futschik (http://itb.biologie.hu-berlin.de/~futschik)

Examples

```r
if (interactive()){
  ### Data loading and pre-processing
  data(yeast) # data set includes 17 measurements
  yeastF <- filter.NA(yeast)
  yeastF <- fill.NA(yeastF)
  yeastF <- standardise(yeastF)

  ### Soft clustering and visualisation
  cl <- mfuzz(yeastF, c=20, m=1.25)
  acore.list <- acore(yeastF, cl=cl, min.acore=0.7)
}
```
**cselection**

Repeated soft clustering for detection of empty clusters for estimation of optimised number of clusters

---

**Description**

This function performs repeated soft clustering for a range of cluster numbers \( c \) and reports the number of empty clusters detected.

**Usage**

```r
cselection(eset, m, crange = seq(4, 32, 4), repeats = 5, visu = TRUE, ...)
```

**Arguments**

- `eset`: object of class `ExpressionSet`.
- `m`: value of fuzzy c-means parameter \( m \).
- `crange`: range of number of clusters \( c \).
- `repeats`: number of repeated clusterings.
- `visu`: If \( \text{visu} = \text{TRUE} \) plot of number of empty clusters is produced.
- `...`: additional arguments for underlying `mfuzz`.

**Details**

A soft cluster is considered as empty, if none of the genes has a corresponding membership value larger than 0.5

**Value**

A matrix with the number of empty clusters detected is generated.

**Note**

The `cselection` function may help to determine an accurate cluster number. However, it should be used with care, as the determination remains difficult especially for short time series and overlapping clusters. A better way is likely to perform clustering with a range of cluster numbers and subsequently assess their biological relevance e.g. by GO analyses.

**Author(s)**


**References**


L. Kumar and M. Futschik, Mfuzz: a software package for soft clustering of microarray data, Bioinformation, 2(1) 5-7, 2007
Examples

```r
if (interactive()){
  data(yeast)
  # Data pre-processing
  yeastF <- filter.NA(yeast)
  yeastF <- fill.NA(yeastF)
  yeastF <- standardise(yeastF)

  ### parameter selection
  # Empty clusters should not appear
  cl <- mfuzz(yeastF,c=20,m=1.25)
  mfuzz.plot(yeastF,cl=cl,mfrow=c(4,5))

  # Note: The following calculation might take some time
  tmp <- cselection(yeastF,m=1.25,crange=seq(5,40,5),repeats=5,visu=TRUE)
  # derivation of number of non-empty clusters (crosses) from diagonal
  # line indicate appearance of empty clusters

  # Empty clusters might appear
  cl <- mfuzz(yeastF,c=40,m=1.25)
  mfuzz.plot(yeastF,cl=cl,mfrow=c(4,5))
}
```

---

### Dmin

Calculation of minimum centroid distance for a range of cluster numbers for estimation of optimised number of clusters

---

**Description**

This function performs repeated soft clustering for a range of cluster numbers \( c \) and reports the minimum centroid distance.

**Usage**

```r
Dmin(eset,m,crange=seq(4,40,4),repeats=3,visu=TRUE)
```

**Arguments**

- `eset` object of class `ExpressionSet`
- `m` value of fuzzy \( c \)-means parameter \( m \)
- `crange` range of number of clusters \( c \)
- `repeats` number of repeated clusterings.
- `visu` If `visu=TRUE` plot of average minimum centroid distance is produced

**Details**

The minimum centroid distance is defined as the minimum distance between two cluster centers produced by the \( c \)-means clusterings.
**Value**

The average minimum centroid distance for the given range of cluster number is returned.

**Note**

The minimum centroid distance can be used as cluster validity index. For an optimal cluster number, we may see a ‘drop’ of minimum centroid distance when plotted versus a range of cluster number and a slower decrease of the minimum centroid distance for higher cluster number. More information and some examples can be found in the study of Schwaemmle and Jensen (2010). However, it should be used with care, as the determination remains difficult especially for short time series and overlapping clusters. Alternatively, the function `cselection` can be used or functional enrichment analysis (e.g. using Gene Ontology) can help to adjust the cluster number.

**Author(s)**


**References**


L. Kumar and M. Futschik, Mfuzz: a software package for soft clustering of microarray data, Bioinformation, 2(1) 5-7, 2007

Schwaemmle and Jensen, Bioinformatics, Vol. 26 (22), 2841-2848, 2010

**Examples**

```r
if (interactive()){
  data(yeast)
  # Data pre-processing
  yeastF <- filter.NA(yeast)
  yeastF <- fill.NA(yeastF)
  yeastF <- standardise(yeastF)

  #### parameter selection
  # For fuzzifier m, we could use mestimate
  m1 <- mestimate(yeastF)
  m1 # 1.15

  # or the function partcoef (see example there)

  # For selection of c, either cselection (see example there)
  # or

  tmp <- Dmin(eset,m=m1,crange=seq(4,40,4),repeats=3,visu=TRUE)# Note: This calculation might take some time

  # It seems that the decrease for c ~ 20 - 25 24 and thus 20 might be
  # a suitable number of clusters
}
```
Replacement of missing values

Description

Methods for replacement of missing values. Missing values should be indicated by NA in the expression matrix.

Usage

fill.NA(eset, mode="mean", k=10)

Arguments

- **eset**: object of the class `ExpressionSet`.
- **mode**: method for replacement of missing values:
  - `mean` - missing values will be replaced by the mean expression value of the gene,
  - `median` - missing values will be replaced by the median expression value of the gene,
  - `knn` - missing values will be replaced by the averaging over the corresponding expression values of the k-nearest neighbours,
  - `knnw` - same replacement method as `knn`, but the expression values averaged are weighted by the distance to the corresponding neighbour
- **k**: Number of neighbours, if one of the `knn` method for replacement is chosen (`knn`, `knnw`).

Value

The function produces an object of the `ExpressionSet` class with missing values replaced.

Note

The replacement methods `knn` and `knnw` can computationally intensive for large gene expression data sets. It may be a good idea to run these methods as a ‘lunchtime’ or ‘overnight’ job.

Author(s)

Matthias E. Futschik ([http://itb.biologie.hu-berlin.de/~futschik](http://itb.biologie.hu-berlin.de/~futschik)) and Lokesh Kumar

Examples

```r
if (interactive()){  
data(yeast) # data set includes 17 measurements  
  yeastF <- filter.NA(yeast)  
  yeastF <- fill.NA(yeastF)  
}
```
**filter.NA**

*Filtering of genes based on number of non-available expression values.*

**Description**

This function can be used to exclude genes with a large number of expression values not available.

**Usage**

```r
filter.NA(eset, thres=0.25)
```

**Arguments**

- `eset` object of the class “ExpressionSet”.
- `thres` threshold for excluding genes. If the percentage of missing values (indicated by NA in the expression matrix) is larger than `thres`, the corresponding gene will be excluded.

**Value**

The function produces an object of the ExpressionSet class. It is the same as the input `eset` object, except for the genes excluded.

**Author(s)**

Matthias E. Futschik ([http://www.sysbiolab.eu](http://www.sysbiolab.eu))

**Examples**

```r
if (interactive()){  
data(yeast) # data set includes 17 measurements  
yeastF <- filter.NA(yeast) # genes are excluded if more than 4 measurements are missing  }
```

---

**filter.std**

*Filtering of genes based on their standard deviation.*

**Description**

This function can be used to exclude genes with low standard deviation.

**Usage**

```r
filter.std(eset, min.std, visu=TRUE)
```
**Arguments**

- **eset**: object of the class `ExpressionSet`.
- **min.std**: threshold for minimum standard deviation. If the standard deviation of a gene’s expression is smaller than min.std the corresponding gene will be excluded.
- **visu**: If visu is set to TRUE, the ordered standard deviations of genes’ expression values will be plotted.

**Value**

The function produces an object of the `ExpressionSet` class. It is the same as the input eset object, except for the genes excluded.

**Note**

As soft clustering is noise robust, pre-filtering can usually be avoided. However, if the number of genes with small expression changes is large, such pre-filtering may be necessary to reduce noise.

**Author(s)**

Matthias E. Futschik ([http://itb.biologie.hu-berlin.de/~futschik](http://itb.biologie.hu-berlin.de/~futschik))

**Examples**

```r
data(yeast)  # data set includes 17 measurements
yeastF <- filter.NA(yeast)  # filtering of genes based on missing values
yeastF <- filter.std(yeastF,min.std=0.3)  # filtering of genes based on standard deviation
```

**Description**

This function is a wrapper function for `kmeans` of the e1071 package. It performs hard clustering of genes based on their expression values using the k-means algorithm.

**Usage**

```r
kmeans2(eset,k,iter.max=100)
```

**Arguments**

- **eset**: object of the class `ExpressionSet`.
- **k**: number of clusters.
- **iter.max**: maximal number of iterations.

**Value**

An list of clustering components (see `kmeans`).

**Author(s)**

Matthias E. Futschik ([http://itb.biologie.hu-berlin.de/~futschik](http://itb.biologie.hu-berlin.de/~futschik))
kmeans2.plot

See Also

kmeans

Examples

```r
if (interactive()){
  data(yeast)
  # Data pre-processing
  yeastF <- filter.NA(yeast)
  yeastF <- fill.NA(yeastF)
  yeastF <- standardise(yeastF)

  # K-means clustering and visualisation
  kl <- kmeans2(yeastF,k=20)
  kmeans2.plot(yeastF,kl=kl,mfrow=c(2,2))
}
```

kmeans2.plot  
Plotting results for k-means clustering

Description

This function visualises the clusters produced by kmeans2.

Usage

```r
kmeans2.plot(eset,kl,mfrow=c(1,1))
```

Arguments

- `eset`  
  object of the class "ExpressionSet".
- `kl`  
  list produced by kmeans2.
- `mfrow`  
  determines splitting of graphic window.

Value

The function displays the temporal profiles of clusters detected by k-means.

Author(s)

Matthias E. Futschik [http://itb.biologie.hu-berlin.de/~futschik](http://itb.biologie.hu-berlin.de/~futschik)

Examples

```r
if (interactive()){
  data(yeast)
  # Data pre-processing
  yeastF <- filter.NA(yeast)
  yeastF <- fill.NA(yeastF)
  yeastF <- standardise(yeastF)

  # K-means clustering and visualisation
  kl <- kmeans2(yeastF,k=20)
```
Calculating of membership values for new data based on existing clustering

Description

Function that calculates the membership values of genes based on provided data and existing clustering

Usage

`membership(x, clusters, m)`

Arguments

- `x` expression vector or expression matrix
- `clusters` cluster centroids from existing clustering
- `m` fuzzification parameter

Value

Matrix of membership values for new genes

Note

This function calculates membership values for new data based on existing cluster centroids and fuzzification parameter. It can be useful, for instance, when comparing two time series, to assess whether the same gene in the different time series changes its cluster association.

Author(s)

Matthias E. Futschik ([http://www.sysbiolab.eu](http://www.sysbiolab.eu))

Examples

```r
if (interactive()){
  data(yeast)
  yeastF <- filter.NA(yeast)
  yeastF <- fill.NA(yeastF) # for illustration only; rather use knn method
  yeastF <- standardise(yeastF)

  cl <- mfuzz(yeastF,c=20,m=1.25)

  m <- 1.25
  clusters <- cl[[1]]
  x <- matrix(rnorm(2*17),nrow=2) # new expression matrix with two genes
  mem.tmp <- membership(x,clusters=clusters,m=m) #membership values
}
```
**mestimate**

Estimate for optimal fuzzifier \( m \)

**Description**

This function estimates an optimal setting of fuzzifier \( m \)

**Usage**

```r
mestimate(eset)
```

**Arguments**

- `eset` object of class “ExpressionSet”

**Details**

Schwaemmle and Jensen proposed a method to estimate \( m \), which was motivated by the evaluation of fuzzy clustering applied to randomized datasets. The estimated \( m \) should give the minimum fuzzifier value which prevents clustering of randomized data.

**Value**

Estimate for optimal fuzzifier.

**Author(s)**

Matthias E. Futschik ([http://itb.biologie.hu-berlin.de/~futschik](http://itb.biologie.hu-berlin.de/~futschik))

**References**

Schwaemmle and Jensen, Bioinformatics, Vol. 26 (22), 2841-2848, 2010

**Examples**

```r
if (interactive()){
  data(yeast)
  # Data pre-processing
  yeastF <- filter.NA(yeast)
  yeastF <- fill.NA(yeastF)
  yeastF <- standardise(yeastF)

  #### parameter selection
  # For fuzzifier \( m \), we could use mestimate
  m1 <- mestimate(yeastF)
  m1 # 1.15
  cl <- mfuzz(yeastF,c=20,m=m1)
  mfuzz.plot(yeastF,cl=cl,mfrow=c(4,5))
}
```
mfuzz

Function for soft clustering based on fuzzy c-means.

Description

This function is a wrapper function for cmeans of the e1071 package. It performs soft clustering of genes based on their expression values using the fuzzy c-means algorithm.

Usage

mfuzz(eset, centers, m, ...)

Arguments

eset        object of the class “ExpressionSet”.
centers     number of clusters.
m          fuzzification parameter.
...         additional parameters for cmeans.

Details

This function is the core function for soft clustering. It groups genes based on the Euclidean distance and the c-means objective function which is a weighted square error function. Each gene is assigned a membership value between 0 and 1 for each cluster. Hence, genes can be assigned to different clusters in a gradual manner. This contrasts hard clustering where each gene can belongs to a single cluster.

Value

An object of class flcust (see cmeans) which is a list with components:
centers    the final cluster centers.
size       the number of data points in each cluster of the closest hard clustering.
cluster    a vector of integers containing the indices of the clusters where the data points are assigned to for the closest hard clustering, as obtained by assigning points to the (first) class with maximal membership.
iter       the number of iterations performed.
membership a matrix with the membership values of the data points to the clusters.
withinerror the value of the objective function.
call       the call used to create the object.

Note

Note that the clustering is based solely on the exprs matrix and no information is used from the phenoData. In particular, the ordering of samples (arrays) is the same as the ordering of the columns in the exprs matrix. Also, replicated arrays in the exprs matrix are treated as independent by the mfuzz function i.e. they should be averagered prior to clustering or placed into different distinct “ExpressionSet” objects.
Author(s)
Matthias E. Futschik (http://www.sysbiolab.eu)

References
L. Kumar and M. Futschik, Mfuzz: a software package for soft clustering of microarray data, Bioinformation, 2(1) 5-7, 2007

See Also
cmeans

Examples
if (interactive()){  
data(yeast)  
  # Data pre-processing  
  yeastF <- filter.NA(yeast)  
  yeastF <- fill.NA(yeastF) # for illustration only; rather use knn method  
  yeastF <- standardise(yeastF)  
  # Soft clustering and visualisation  
  cl <- mfuzz(yeastF,c=20,m=1.25)  
  mfuzz.plot(yeastF,cl=cl,mfrow=c(2,2))  
  # Plotting center of cluster 1  
  X11(); plot(cl[[1]][1,],type="l",ylab="Expression")  
  # Getting the membership values for the first 10 genes in cluster 1  
  cl[[4]][1:10,1]  
}
min.mem  Genes with membership values below min.mem will not be displayed.
time.labels  labels can be given for the time axis.
new.window  should a new window be opened for graphics.

Value
The function generates plots where the membership of genes is color-encoded.

Author(s)
Matthias E. Futschik (http://www.sysbiolab.eu/matthias)

Examples
if (interactive()){
data(yeast)
# Data pre-processing
yeastF <- filter.NA(yeast)
yeastF <- fill.NA(yeastF)
yeastF <- standardise(yeastF)

# Soft clustering and visualisation
cl <- mfuzz(yeastF,c=20,m=1.25)
mfuzz.plot(yeastF,cl=cl,mfrow=c(2,2))

# display of cluster cores with alpha = 0.5
mfuzz.plot(yeastF,cl=cl,mfrow=c(2,2),min.mem=0.5)

# display of cluster cores with alpha = 0.7
mfuzz.plot(yeastF,cl=cl,mfrow=c(2,2),min.mem=0.7)
}

mfuzz.plot2

Plotting results for soft clustering with additional options

Description
This function visualises the clusters produced by mfuzz. It is similar to mfuzz.plot, but offers more options for adjusting the plots.

Usage
mfuzz.plot2(eset,cl,mfrow=c(1,1),colo,min.mem=0,time.labels,time.points,
ylim.set=c(0,0), xlab="Time",ylab="Expression changes",x11=TRUE,
ax.col="black", bg = "white",col.axis="black",col.lab="black",
col.main="black",col.sub="black",col="black",centre=FALSE,
centre.col="black",centre.lwd=2,
Xwidth=5,Xheight=5,single=FALSE,...)
Arguments

eset  object of the class ExpressionSet.
cl   object of class flclust.
mfrow determines splitting of graphic window. Use mfrow=NA if layout is used (see example).
colo color palette to be used for plotting. If the color argument remains empty, the default palette is used. If the colo = “fancy”, an alternative (fancier) palette will be used.
min.mem Genes with membership values below min.mem will not be displayed.
time.labels labels for ticks on x axis.
time.points numerical values for the ticks on x axis. These can be used if the measured time points are not equidistant.
ylim.set Vector of min. and max. y-value set for plotting. If ylim.set=c(0,0), min. and max. value will be determined automatically.
xlab label for x axis
ylab label for y axis
x11 If TRUE, a new window will be open for plotting.
ax.col Color of axis line.
bg Background color.
col.axis Color for axis annotation.
col.lab Color for axis labels.
col.main Color for main titles.
col.sub Color for sub-titles.
col Default plotting color.
centre If TRUE, a line for the cluster centre will be drawn.
centre.col Color of the line for the cluster centre
centre.lwd Width of the line for the cluster centre
Xwidth Width of window.
Xheight Height of window.
single Integer if a specific cluster is to be plotted, otherwise it should be set to FALSE.
... Additional, optional plotting arguments passed to plot.default and axes functions such as cex.lab,cex.main,cex.axis

Value

The function generates plots where the membership of genes is color-encoded.

Author(s)

Matthias E. Futschik (http://www.sysbiolab.eu/matthias)
Examples

```r
if (interactive()){
  data(yeast)
  # Data pre-processing
  yeastF <- filter.NA(yeast)
  yeastF <- fill.NA(yeastF)
  yeastF <- standardise(yeastF)

  # Soft clustering and visualisation
  cl <- mfuzz(yeastF,c=20,m=1.25)
  mfuzz.plot2(yeastF, cl=cl, mfrow=c(2,2)) # same output as mfuzz.plot
  mfuzz.plot2(yeastF, cl=cl, mfrow=c(2,2), centre=TRUE) # lines for cluster centres will be included

  # More fancy choice of colors
  mfuzz.plot2(yeastF, cl=cl, mfrow=c(2,2), colo="fancy",
  ax.col="red", bg = "black", col.axis="red", col.lab="white",
  col.main="green", col.sub="blue", col="blue", cex.main=1.3, cex.lab=1.1)

  ### Single cluster with colorbar (cluster # 3)
  X11(width=12)
  mat <- matrix(1:2, ncol=2, nrow=1, byrow=TRUE)
  l <- layout(mat, width=c(5,1))
  mfuzz.plot2(yeastF, cl=cl, mfrow=NA, colo="fancy", ax.col="red", bg = "black", col.axis="red", col.lab="white",
  col.main="green", col.sub="blue", col="blue", cex.main=2, single=3, x11=FALSE)
  mfuzzColorBar(col="fancy", main="Membership", cex.main=1)

  ### Single cluster with colorbar (cluster # 3)
  X11(width=14)
  mat <- matrix(1:2, ncol=2, nrow=1, byrow=TRUE)
  l <- layout(mat, width=c(5,1))
  mfuzz.plot2(yeastF, cl=cl, mfrow=NA, colo="fancy", ax.col="red", bg = "black", col.axis="red", col.lab="white",
  time.labels = c(paste(seq(0,160,10), "min")),
  col.main="green", col.sub="blue", col="blue", cex.main=2, single=3, x11=FALSE)
  mfuzzColorBar(col="fancy", main="Membership", cex.main=1)

}
```

```

mfuzzColorBar

Plots a colour bar

Description

This function produces a (separate) colour bar for graphs produced by mfuzz.plot

Usage

mfuzzColorBar(col, horizontal=FALSE,...)
```
Mfuzzgui

Arguments

col vector of colours used. If missing, the same vector as the default vector for mfuzz.plot is used. If col="fancy", an alternative color palette is used (see mfuzz.plot2).

horizontal If TRUE, a horizontal colour bar is generated, otherwise a vertical one will be produced.

... additional parameter passed to maColorBar (see also example in mfuzz.plot2)

Author(s)

Matthias E. Futschik (http://itb.biologie.hu-berlin.de/~futschik)

See Also

maColorBar

Examples

if (interactive()){
  X11(w=1.5,h=5);
  par(mar=c(1,1,1,5))
  mfuzzColorBar()

  mfuzzColorBar(col="fancy",main="Membership value")
}

Mfuzzgui

Graphical user interface for Mfuzz package

Description

The function Mfuzzgui provides a graphical user interface for clustering of microarray data and visualisation of results. It is based on the functions of the Mfuzz package.

Usage

Mfuzzgui()

Details

The function Mfuzzgui launches a graphical user interface for the Mfuzz package. It is based on Tk widgets using the R TclTk interface by Peter Dalgaard. It also employs some pre-made widgets from the tkWidgets Bioconductor-package by Jianhua Zhang for the selection of objects/files to be loaded.

Mfuzzgui provides a convenient interface to most functions of the Mfuzz package without restriction of flexibility. An exception is the batch processes such as partcoeff and cselection routines which are used for parameter selection in fuzzy c-means clustering of microarray data. These routines are not included in Mfuzzgui. To select various parameters, the underlying Mfuzz routines may be applied.

Usage of Mfuzzgui does not require an pre-built exprSet object but can be used with tab-delimited text files containing the gene expression data. Note, however, that the clustering is based
on the ordering of samples (arrays) as of the columns in the expression matrix of the exprSet object or in the uploaded table, respectively. Also, replicated arrays in the expression matrix (or table) are treated as independent by the mfuzz function and, thus, should be averaged prior to clustering.

For a overview of the functionality of Mfuzzgui, please refer to the package vignette. For a description of the underlying functions, please refer to the Mfuzz package.

Value

Mfuzzgui returns a tcclObj object.

Note

The newest versions of Mfuzzgui can be found at the Mfuzz webpage (http://itb.biologie.hu-berlin.de/~futschik/software/R/Mfuzz).

Author(s)

Matthias E. Futschik (http://itb.biologie.hu-berlin.de/~futschik) and Lokesh Kumar

References


See Also

mfuzz

overlap

Calculation of the overlap of soft clusters

Description

This function calculates the overlap of clusters produced by mfuzz.

Usage

overlap(cl)

Arguments

cl object of class flclust

Value

The function generates a matrix of the normalised overlap of soft clusters. The overlap indicates the extent of “shared” genes between clusters. For a mathematical definiton of the overlap, see the vignette of the package or the reference below.
Author(s)
Matthias E. Futschik (http://itb.biologie.hu-berlin.de/~futschik)

References

Examples
if (interactive()){
data(yeast)
# Data pre-processing
yeastF <- filter.NA(yeast)
yeastF <- fill.NA(yeastF)
yeastF <- standardise(yeastF)

# Soft clustering and visualisation
cl <- mfuzz(yeastF,c=20,m=1.25)
mfuzz.plot(yeastF,cl=cl,mfrow=c(4,5))

# Calculation of cluster overlap and visualisation
O <- overlap(cl)
X11()
Ptmp <- overlap.plot(cl,over=O,thres=0.05)
}

overlap.plot

Description
This function visualises the cluster overlap produced by overlap.

Usage
overlap.plot(cl,overlap,thres=0.1, scale=TRUE, magni=30, P=NULL)

Arguments
cl object of class “flclust”
overlap matrix of cluster overlap produced by overlap
thres threshold for visualisation. Cluster overlaps below the threshold will not be visualised.
scale Scale parameter for principal component analysis by prcomp
magni Factor for increase the line width for cluster overlap.
P Projection matrix produced by principal component analysis.
partcoef

Calculation of the partition coefficient matrix for soft clustering

Description

This function calculates partition coefficient for clusters within a range of cluster parameters. It can be used to determine the parameters which lead to uniform clustering.

Usage

partcoef(eset, crange=seq(4,32,4), mrange=seq(1.05,2,0.1), ...)

Value

A plot is generated based on a principal component analysis of the cluster centers. The overlap is visualised by lines with variable width indicating the strength of the overlap. Additionally, the matrix of principal components is returned. This matrix can be re-used for other projections to compare the overlap and global cluster structure of different clusterings.

Author(s)

Matthias E. Futschik (http://itb.biologie.hu-berlin.de/~futschik)

See Also

prcomp

Examples

if (interactive()){
  data(yeast)
  # Data pre-processing
  yeastF <- filter.NA(yeast)
  yeastF <- fill.NA(yeastF)
  yeastF <- standardise(yeastF)
  # Soft clustering
  cl <- mfuzz(yeastF,c=20,m=1.25)
  X11();mfuzz.plot(yeastF,cl=cl,mfrow=c(4,5))
  O <- overlap(cl)
  X11();Ptmp <- overlap.plot(cl,over=O,thres=0.05)
  # Alternative clustering
  cl <- mfuzz(yeastF,c=10,m=1.25)
  X11();mfuzz.plot(yeastF,cl=cl,mfrow=c(3,4))
  O <- overlap(cl)
  X11();overlap.plot(cl,over=0,P=Ptmp,thres=0.05)
  # visualisation based on principal components from previous projection
}
partcoef

Arguments

eset object of class “ExpressionSet”.

range range of number of clusters c.
mrange range of clustering parameter m.

... additional arguments for underlying mfuzz.

Details

Introduced by Bezdek (1981), the partition coefficient F is defined as the sum of squares of values of the partition matrix divided by the number of values. It is maximal if the partition is hard and reaches a minimum for \( U = 1/c \) when every gene is equally assigned to every cluster.

It is well-known that the partition coefficient tends to decrease monotonically with increasing n. To reduce this tendency we defined a normalized partition coefficient where the partition for uniform partitions are subtracted from the actual partition coefficients (Futschik and Kasabov, 2002).

Value

The function generates the matrix of partition coefficients for a range of c and m values. It also produces a matrix of normalised partition coefficients as well as a matrix with partition coefficient for uniform partitions.

Author(s)

Matthias E. Futschik (http://itb.biologie.hu-berlin.de/~futschik)

References

1. J.C. Bezdek, Pattern recognition with fuzzy objective function algorithms, Plenum, 1981

Examples

if (interactive()){
data(yeast)

# Data pre-processing
yeastF <- filter.NA(yeast)
yeastF <- fill.NA(yeastF)
yeastF <- standardise(yeastF)

#### parameter selection
yeastFR <- randomise(yeastF)
cl <- mfuzz(yeastFR,c=20,m=1.1)
mfuzz.plot(yeastFR,cl=cl,mfrow=c(4,5)) # shows cluster structures (non-uniform partition)

tmp <- partcoef(yeastFR) # This might take some time.
F <- tmp[[1]];F.n <- tmp[[2]];F.min <- tmp[[3]]

# Which clustering parameters result in a uniform partition?
F > 1.01 * F.min

c1 <- mfuzz(yeastFR,c=20,m=1.25) # produces uniform partition
standardise

mfuzz.plot(yeastFR, cl=c1, mfrow=c(4,5))
# uniform coloring of temporal profiles indicates uniform partition

randomise  

**Randomisation of data**

**Description**

This function randomise the time order for each gene separately.

**Usage**

randomise(eset)

**Arguments**

- eset: object of the class `ExpressionSet`.

**Value**

The function produces an object of the `ExpressionSet` class with randomised expression data.

**Author(s)**

Matthias E. Futschik (http://itb.biologie.hu-berlin.de/~futschik)

**Examples**

data(yeast) # data set includes 17 measurements
yeastR <- randomise(yeast)

standardise  

**Standardization of expression data for clustering.**

**Description**

Standardisation of the expression values of every gene/transcript/protein is carried out, so that the average expression value for each gene/transcript/protein is zero and the standard deviation of its expression profile is one.

**Usage**

standardise(eset)

**Arguments**

- eset: object of the class `ExpressionSet`.

**Value**

The function produces an object of the `ExpressionSet` class with standardised expression values.
Note

Mfuzz assumes that the given expression data are preprocessed (including the normalisation). The function standardise does not replace the normalisation step. Note the difference: Normalisation is carried out to make different samples comparable, while standardisation (in Mfuzz) is carried out to make transcripts (genes) comparable.

Author(s)

Matthias E. Futschik (http://www.sysbiolab.eu)

Examples

```r
if (interactive()){
  data(yeast)
  # Data pre-processing
  yeastF <- filter.NA(yeast)
  yeastF <- fill.NA(yeastF)
  yeastF <- standardise(yeastF)

  # Soft clustering and visualisation
  cl <- mfuzz(yeastF,c=20,m=1.25)
  mfuzz.plot(yeastF,cl=cl,mfrow=c(4,5))
}
```

---

**standardise2**

*Standardization in regards to selected time-point*

**Description**

Standardisation of the expression values of every gene is performed, so that the expression values at a chosen time point are zero and the standard deviation of expression profiles of individual genes/transcripts/proteins is one.

**Usage**

```r
standardise2(eset, timepoint=1)
```

**Arguments**

- `eset`: object of the class `ExpressionSet`.
- `timepoint`: integer: which time point should have expression values of zero.

**Value**

The function produces an object of the `ExpressionSet` class with standardised expression values.

**Note**

Mfuzz assumes that the given expression data are preprocessed (including the normalisation). The function `standardise2` does not replace the normalisation step. Note the difference: Normalisation is carried out to make different samples comparable, while standardisation (in Mfuzz) is carried out to make transcripts (genes) comparable.
table2eset

Conversion of table to Expression set object.

Description

A expression matrix stored as a table (in a defined format) is read and converted to Expression Set object.

Usage

table2eset(filename)

Arguments

filename name of file to be scanned in

Details

The expression matrix stored as table in the file has to follow some conventions in order to be able to be converted to an Expression Set object: The first row of the file contains sample labels and optionally, the second column can contains the time points. If the second row is used for the input the time, the first field in the second row must contain “Time”. Similarly, the first column contains unique gene IDs and optionally second row can contain gene names. If the second row contains gene names, the second field in the first row must contain “Gene.Name”. The rest of the file contains expression data. As example, two tables with expression data are provided. These examples can be viewed by inputing data(yeast.table) and data(yeast.table2) in the R console.

Value

An Expression Set object is generated.

Author(s)

Matthias E. Futschik (http://www.sysbiolab.eu)
top.count

Determines the number for which each gene has highest membership value in all cluster

Description

This function calculates the number, for which each gene appears to have the top membership score in the partition matrix of clusters produced by mfuzz.

Usage

top.count(cl)

Arguments

cl object of class “flclust”

Value

The function generates a vector containing a count for each gene, which is just the number of times that particular gene has acquired the top membership score.

Author(s)

Lokesh Kumar and Matthias E. Futschik (http://itb.biologie.hu-berlin.de/~futschik)

Examples

if (interactive()){  
data(yeast)  
  # Data pre-processing  
  yeastF <- filter.NA(yeast)  
  yeastF <- fill.NA(yeastF)  
  yeastF <- standardise(yeastF)  
  # Soft clustering and visualisation  
  cl <- mfuzz(yeastF,c=20,m=1.25)  
  top.count(cl)  
}

eyeast

Gene expression data of the yeast cell cycle

Description

The data contains gene expression measurements for 3000 randomly chosen genes of the yeast mutant cdc28 as performed and described by Cho et al. For details, see the reference.

Usage

data(yeast)
Format

An object of class “ExpressionSet”.

Source

The data was downloaded from Yeast Cell Cylce Analysis Project webservice and converted to an ExpressionSet object.

References


---

**yeast.table**

*Gene expression data of the yeast cell cycle as table*

Description

The data serves as an example for the format required for uploading tables with expression data into Mfuzzgui. The first row contains the names of the samples, the second row contains the measured time points. Note that “TIME” has to placed in the first field of the second row.

The first column contains unique identifiers for genes; optionally the second row can contain gene names if “GENE.NAMES” is in the second field in the first row.

An example for an table without optional fields is the dataset yeast.table2.

The exemplary tables can be found in the data sub-folder of the Mfuzzgui package.

References


See Also

yeast.table2

---

**yeast.table2**

*Gene expression data of the yeast cell cycle as table*

Description

The data serves as an example for the format required to upload tables with expression data into Mfuzzgui. The first row contains the names of the samples and the first column contains unique identifiers for genes. To input measurement time and gene names, refer to yeast.table.

The exemplary tables can be found in the data sub-folder of the Mfuzzgui package.

References

See Also

yeast.table
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