Package 'ORIClust'

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Description A user-friendly R-based software package for gene clustering. Clusters are given by genes matched to prespecified profiles across various ordered treatment groups. It is particularly useful for analyzing data obtained from short time-course or dose-response microarray experiments.					
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ORIClust-package Order-restricted Information Criterion-based Clustering Algorithm

Description

ORIClust is a user-friendly R-based software package for gene clustering. Clusters are given by genes matched to prespecified profiles across various ordered treatment groups. It is particularly useful for analyzing data obtained from short time-course or dose-response microarray experiments.

Details

Package:	ORIClust
Type:	Package
Version:	1.0
Date:	2009-05-24
License:	GPL-2
LazyLoad:	yes

The main functions are ORICC1 and ORICC2, see the documentation files with examples.

Author(s)

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References

Liu, T., Lin, N., Shi, N. and Zhang, B. (2009), Information criterion-based clustering with orderrestricted candidate profiles in short time-course microarray experiments, BMC Bioinformatics, 10: 146.

Breast

Breast cancer cell line data

Description

This data set comes from a breast cancer cell line microarray study. The experiment was done as follows. First, the MCF-7 breast cancer cell line was treated with 17 beta-estradiol or ethanol (vehicle control). Then, samples were harvested at 1, 4, 12, 24, 36 and 48 hours after treatment. At each time point, M = 8 replicate arrays were prepared with each array consisting of G = 1901 genes.

complete.profile

Usage

Breast

Format

A matrix containing 1901 rows and 50 columns.

References

Lobenhofer, E., Bennett, L., Cable, P., Li, L., Bushel, P., and Afshari, C. (2002), *Regulation of DNA replication fork genes by 17 beta-estradiol*. Molec. Endocrin., **16**, 1215-1229.

complete.profile complete.profile

Description

Returns the log-maximum likelihood and the estimator of the mean when there is no inequality constraint.

Usage

```
complete.profile(data,x,n.rep)
```

Arguments

data	A vector containing the expressions of one gene.
x	A vector consisting of the average expression at time points $(1, 2,, T)$, where T is the total number of time points.
n.rep	A vector consisting of the number of replicate arrays at time points $(1, 2,, T)$, where T is the total number of time points.

Value

logelr	Log-maximum likelihood
mu	A vector containing the estimator of the mean

Author(s)

Tianqing Liu, Nan Lin, Ningzhong Shi and Baoxue Zhang Maintainer:Tianqing Liu <tianqingliu@gmail.com> cyclical.max.min cyclical.max.min

Description

Returns the log-maximum likelihood and the estimator of the mean under cyclical profile with maximum at max1 and minimum at min1 (max1 < min1).

Usage

cyclical.max.min(data,x,n.rep,max1,min1)

Arguments

data	A vector containing the expressions of one gene.
x	A vector consisting of the average expression at time points $(1, 2,, T)$, where T is the total number of time points.
n.rep	A vector consisting of the number of replicate arrays at time points $(1, 2,, T)$, where T is the total number of time points.
max1	Cyclical profile with maximum at max1.
min1	Cyclical profile with minimum at min1.

Value

logelr	Log-maximum likelihood
mu	A vector containing the estimator of the mean

Author(s)

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References

Liu, T., Lin, N., Shi, N. and Zhang, B. (2009), *Information criterion-based clustering with orderrestricted candidate profiles in short time-course microarray experiments.* BMC Bioinformatics, **10**: 146.

Robertson, T., Wright, F. T. and Dykstra, R. L. (1988). Order restricted statistical inference. New York: Wiley.

cyclical.min.max cyclical.min.max

Description

Returns the log-maximum likelihood and the estimator of the mean under cyclical profile with minimum at min1 and maximum at max1 (min1 < max1).

Usage

cyclical.min.max(data,x,n.rep,min1,max1)

Arguments

A vector containing the expressions of one gene.
A vector consisting of the average expression at time points $(1, 2,, T)$, where T is the total number of time points.
A vector consisting of the number of replicate arrays at time points $(1, 2,, T)$, where T is the total number of time points.
Cyclical profile with minimum at min1
cyclical profile with maximum at max1

Value

logelr	Log-maximum likelihood
mu	A vector containing the estimator of the mean

Author(s)

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References

Liu, T., Lin, N., Shi, N. and Zhang, B. (2009), *Information criterion-based clustering with orderrestricted candidate profiles in short time-course microarray experiments.* BMC Bioinformatics, **10**: 146.

Robertson, T., Wright, F. T. and Dykstra, R. L. (1988). Order restricted statistical inference. New York: Wiley.

decreasing

Description

Returns the log-maximum likelihood and the estimator of the mean under the monotone decreasing profile.

Usage

decreasing(data,x,n.rep)

Arguments

data	A vector containing the expressions of one gene.
x	A vector consisting of the average expression at time points $(1, 2,, T)$, where T is the total number of time points.
n.rep	A vector consisting of the number of replicate arrays at time points $(1, 2,, T)$, where T is the total number of time points.

Value

logelr	Log-maximum likelihood
mu	A vector containing the estimator of the mean

Author(s)

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References

Liu, T., Lin, N., Shi, N. and Zhang, B. (2009), *Information criterion-based clustering with orderrestricted candidate profiles in short time-course microarray experiments*. BMC Bioinformatics, **10**: 146.

Robertson, T., Wright, F. T. and Dykstra, R. L. (1988). Order restricted statistical inference. New York: Wiley.

down.up

Description

Returns the log-maximum likelihood and the estimator of the mean under down-up profile with minimum at h.

Usage

down.up(data,x,n.rep,h)

Arguments

data	A vector containing the expressions of one gene.
x	A vector consisting of the average expression at time points $(1, 2,, T)$, where T is the total number of time points.
n.rep	A vector consisting of the number of replicate arrays at time points $(1, 2,, T)$, where T is the total number of time points.
h	Down-up profile with minimum at h.

Value

logelr	Log-maximum likelihood
mu	A vector containing the estimator of the mean

Author(s)

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References

Liu, T., Lin, N., Shi, N. and Zhang, B. (2009), *Information criterion-based clustering with orderrestricted candidate profiles in short time-course microarray experiments*. BMC Bioinformatics, **10**: 146.

Robertson, T., Wright, F. T. and Dykstra, R. L. (1988). Order restricted statistical inference. New York: Wiley.

flat.pattern

Description

Returns the log-maximum likelihood and the estimator of the mean under the equality constraint that all means are equal.

Usage

flat.pattern(data,x,n.rep)

Arguments

data	A vector containing the expressions of one gene.
x	A vector consisting of the average expression at time points $(1, 2,, T)$, where T is the total number of time points.
n.rep	A vector consisting of the number of replicate arrays at time points $(1, 2,, T)$, where T is the total number of time points.

Value

logelr	Log-maximum likelihood
mu	A vector containing the estimator of the mean

Author(s)

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References

Liu, T., Lin, N., Shi, N. and Zhang, B. (2009), *Information criterion-based clustering with orderrestricted candidate profiles in short time-course microarray experiments.* BMC Bioinformatics, **10**: 146.

Robertson, T., Wright, F. T. and Dykstra, R. L. (1988). *Order restricted statistical inference*. New York: Wiley.

increasing

increasing

Description

Returns the log-maximum likelihood and the estimator of the mean under the monotone increasing profile.

Usage

increasing(data,x,n.rep)

Arguments

data	A vector containing the expressions of one gene.
x	A vector consisting of the average expression at time points $(1, 2,, T)$, where T is the total number of time points.
n.rep	A vector consisting of the number of replicate arrays at time points $(1, 2,, T)$, where T is the total number of time points.

Value

logelr	Log-maximum likelihood
mu	A vector containing the estimator of the mean

Author(s)

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References

Liu, T., Lin, N., Shi, N. and Zhang, B. (2009), *Information criterion-based clustering with orderrestricted candidate profiles in short time-course microarray experiments*. BMC Bioinformatics, **10**: 146.

Robertson, T., Wright, F. T. and Dykstra, R. L. (1988). Order restricted statistical inference. New York: Wiley.

isodecre

isodecre

Description

Isotonic regression of a with weights w under monotone decreasing profile.

Usage

isodecre(a, w)

Arguments

а	A vector consisting of the average expression at time points $(1, 2,, T)$, where T is the total number of time points.
W	The weights.

Value

is A vector containing the estimator of the me	ean
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Author(s)

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References

Liu, T., Lin, N., Shi, N. and Zhang, B. (2009), *Information criterion-based clustering with orderrestricted candidate profiles in short time-course microarray experiments*. BMC Bioinformatics, **10**: 146.

Robertson, T., Wright, F. T. and Dykstra, R. L. (1988). Order restricted statistical inference. New York: Wiley.

isoincre

isoincre

Description

Isotonic regression of a with weights w under monotone increasing profile.

Usage

isoincre(a, w)

Arguments

а	A vector consisting of the average expression at time points $(1, 2,, T)$, where T is the total number of time points.
W	The weights.

Value

is A vector containing the estimator of the me	ean
--	-----

Author(s)

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References

Liu, T., Lin, N., Shi, N. and Zhang, B. (2009), *Information criterion-based clustering with orderrestricted candidate profiles in short time-course microarray experiments*. BMC Bioinformatics, **10**: 146.

Robertson, T., Wright, F. T. and Dykstra, R. L. (1988). Order restricted statistical inference. New York: Wiley.

ORICC1

Description

One-stage ORICC is a computationally efficient information criterion-based clustering algorithm for selecting and clustering genes according to their time-course or dose-response profiles. This algorithm takes account of the ordering in time-course or dose-response experiments by embedding the order-restricted inference into a model selection framework. This algorithm mainly consist of two steps. In the first step, candidate profiles are defined in terms of inequalities among mean expression levels at different time points or doses levels. In the second step, genes are assigned to the best matched profiles determined by an information criterion for order-restricted inference.

Usage

Arguments

data	A matrix containing the gene expressions.
data.col	Column indices of the gene expression data.
id.col	Column index of the gene ID. Defaults to 1.
n.rep	A vector consisting of the number of replicate arrays at time points $(1, 2,, T)$, where T is the total number of time points.
n.top	The number of genes kept for the final clustering result. Genes are ranked based on expression variation across time or dose levels. Defaults to all genes ORICC1 selects
transform	Transformation of the original data:
	0=None, 1=natural log, 2=square root, 3=cubic root. Defaults to 0.
name.profile	A character string specifying the collection of candidate profiles. This option only supports monotone, up-down and down-up profiles specified as by "decreasing";
	<pre>naste("up down max at", i.sep=""):</pre>
	<pre>paste("down up min at", j, sep=" ");</pre>
	If name.profile="all", the 'decreasing', 'increasing' and all 'up-down' and 'down-up' profiles will be included.
	If name.profile=NULL, 'decreasing', 'increasing' and all 'up-down' and 'down- up' profiles will be absent. Defaults to NULL.
	One can also specify several up-down or down-up profiles together as follows.

profile1=paste("up down max at", c(2,4), sep=" ");

profile2=paste("down up min at", c(3,5), sep=" ");

name.profile=c(profile1,profile2);

then up-down profile with maxima at 2 and 4 as well as down-up profile with minima at 3 and 5 will be included.

cyclical.profile

A matrix with 2 columns. Each element of the matrix must be a number in the set $\{2,3,\ldots,T-1\}$. Each row of the matrix represents a cyclical profile with minima at the first entry of the row and maxima at the 2nd entry. As a result, two elements in the same row must be different. For example, if

cyclical.profile=matrix(c(2,3,4,3),2,2,byrow=T), then the cyclical profile with minima at 2 and maxima at 3 and the cyclical profile with minima at 4 and maxima at 3 will be included as candidate profiles.

If cyclical.profile=NULL, all cyclical profiles will be absent. Defaults to NULL.

complete.profile

	The complete.profile means a profile in which there is no defined inequality constraint.
	If the complete.profile is a candidate profile,
	<pre>complete.profile=1, otherwise,</pre>
	complete.profile=NULL. Defaults to NULL.
onefile	logical: if true (the default) multiple figures for different clusters are output in one file. If FALSE, each cluster is plotted in a seperate file. Defaults to TRUE.
plot.format	The format of the output file containing plots of gene clusters.Users can choose between 'eps' and 'jpg'. Defaults to 'eps'.

Details

The gene expression dataset should be in a tab-delimited txt file, in which the first two columns contain the gene names and their accession numbers or descriptions, and the remaining columns, in their orders, are the geneexpression data (contain multiple columns, i.e. data.col). The dataset is assumed to have been processed so that each row contains the expressions of only one gene.

Value

The results are displayed in a graphical form. The graphics can be stored in a JPG or EPS format. Both the raw gene expression values and the estimated mean expressions are output to external files 'cluster of raw data.txt' and 'cluster of fitted mean data.txt', respectively.

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References

Liu, T., Lin, N., Shi, N. and Zhang, B. (2009), *Information criterion-based clustering with orderrestricted candidate profiles in short time-course microarray experiments*. BMC Bioinformatics, **10**: 146.

Examples

ORICC2

Two-stage ORICC

Description

It is a computationally efficient two-stage algorithm by adding a pre-screening stage. It first screens out genes that show no significant changes over time, and then applies the one-stage algorithm to a much smaller set of remained genes.

Usage

Arguments

data	A matrix containing the gene expressions.
data.col	Column indices of the gene expression data.
id.col	Column index of the gene ID. Defaults to 1.
n.rep	A vector consisting of the number of replicate arrays at time points $(1, 2,, T)$, where T is the total number of time points.
n.top	The number of genes kept for the final clustering result. Genes are ranked based on expression variation across time or dose levels. Defaults to all genes ORICC2 selects
transform	Transformation of the original data:
	0=None, 1=natural log, 2=square root, 3=cubic root. Defaults to 0.
name.profile	A character string specifying the collection of candidate profiles. This option only supports monotone, up-down and down-up profiles specified as by "decreasing";
	"increasing".
	<pre>paste("up down max at",i,sep=" ");</pre>

paste("down up min at", j, sep=" ");

If name.profile="all", the 'decreasing', 'increasing' and all 'up-down' and 'down-up' profiles will be included.

If name.profile=NULL, 'decreasing', 'increasing' and all 'up-down' and 'down-up' profiles will be absent. Defaults to NULL.

One can also specify several up-down or down-up profiles together as follows.

profile1=paste("up down max at", c(2,4), sep=" ");

profile2=paste("down up min at", c(3,5), sep=" ");

name.profile=c(profile1,profile2);

then up-down profile with maxima at 2 and 4 as well as down-up profile with minima at 3 and 5 will be included.

cyclical.profile

A matrix with 2 columns. Each element of the matrix must be a number in the set $\{2,3,\ldots,T-1\}$. Each row of the matrix represents a cyclical profile with minima at the first entry of the row and maxima at the 2nd entry. As a result, two elements in the same row must be different. For example, if

cyclical.profile=matrix(c(2,3,4,3),2,2,byrow=T), then the cyclical profile with minima at 2 and maxima at 3 and the cyclical profile with minima at 4 and maxima at 3 will be included as candidate profiles.

If cyclical.profile=NULL, all cyclical profiles will be absent. Defaults to NULL.

- onefile logical: if true (the default) multiple figures for different clusters are output in one file. If FALSE, each cluster is plotted in a seperate file. Defaults to TRUE.
- plot.format The format of the output file containing plots of gene clusters.Users can choose between 'eps' and 'jpg'. Defaults to 'eps'.

Details

The gene expression dataset should be in a tab-delimited txt file, in which the first two columns contain the gene names and their accession numbers or descriptions, and the remaining columns, in their orders, are the geneexpression data (contain multiple columns, i.e. data.col). The dataset is assumed to have been processed so that each row contains the expressions of only one gene.

Value

The results are displayed in a graphical form. The graphics can be stored in a JPG or EPS format. Both the raw gene expression values and the estimated mean expressions are output to external files 'cluster of raw data.txt' and 'cluster of fitted mean data.txt', respectively.

Author(s)

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References

Liu, T., Lin, N., Shi, N. and Zhang, B. (2009), *Information criterion-based clustering with orderrestricted candidate profiles in short time-course microarray experiments*. BMC Bioinformatics, **10**: 146.

Examples

Description

Returns the log-maximum likelihood and the estimator of the mean under up-down profile with maximum at h.

Usage

up.down(data,x,n.rep,h)

Arguments

data	A vector containing the expressions of one gene.
x	A vector consisting of the average expression at time points $(1, 2,, T)$, where T is the total number of time points.
n.rep	A vector consisting of the number of replicate arrays at time points $(1, 2,, T)$, where T is the total number of time points.
h	Up-down profile with maximum at h.

Value

logelr	Log-maximum likelihood
mu	A vector containing the estimator of the mean

Author(s)

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up.down

References

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