

Package ‘ORCME’

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Type Package

Title Order Restricted Clustering for Microarray Experiments

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Description Provides clustering of genes with similar dose response (or time course) profiles. It implements the method described by Lin et al. (2012).

Imports Iso

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`doseData`*Dose Data Example*

Description

Dose data; a vector of length 12 with 3 observations for each of 4 doses.

Usage

```
data(doseData)
```

Format

The format is: num [1:12] 1 1 1 2 2 2 3 3 ...

Examples

```
data(doseData)
```

```
doseData
```

`geneData`*Gene Expression Data Example*

Description

This dose-response microarray data contains 1000 genes and 4 doses (one control dose (zero dose) and three increasing dose) with 3 arrays at each dose level. Due to confidentiality, it is only part of the real data set.

Usage

```
data(geneData)
```

Format

A data frame with 1000 observations on the following 12 variables.

X1 Sample one with zero dose

X1.1 Sample two with zero dose

X1.2 Sample three with zero dose

X2 Sample one with second dose

X2.1 Sample two with second dose

- X2.2 Sample three with second dose
- X3 Sample one with third dose
 - X3.1 Sample two with third dose
 - X3.2 Sample three with third dose
- X4 Sample one with fourth dose
 - X4.1 Sample two with fourth dose
 - X4.2 Sample three with fourth dose

References

Testing for Trend in Dose-Response Microarray Experiments: a Comparison of Testing Procedures, Multiplicity, and Resampling-Based Inference, Lin et al. 2007, Stat. App. in Gen. & Mol. Bio., 6(1), article 26.

Examples

```
data(geneData)
head(geneData)
```

monotoneDirection	<i>The monotone means under increasing/decreasing trend</i>
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Description

The function calculates the likelihood for the increasing and decreasing trend in the dose response for all the given genes separately gene-by-gene. The trend with the higher likelihood is chosen and the isotonic regression is applied on the means.

Usage

```
monotoneDirection(geneData, doseData)
```

Arguments

geneData	gene expression matrix for all genes
doseData	indicates the dose levels

Value

A list with components

direction	the direction with the higher likelihood of increasing (indicated by "up") or decreasing (indicated by "dn") trend.
incData	isotonic means with respect to dose for those genes that were classified as following the increasing trend.
decData	isotonic means with respect to dose for those genes that were classified as following the decreasing trend.
obsincData	observed gene expression matrix for those genes that were classified as following the increasing trend.
obsdecData	observed gene expression matrix for those genes that were classified as following the decreasing trend.
arrayMean	isotonic means with respect to dose for all genes.

Author(s)

Adetayo Kasim, Martin Otava and Tobias Verbeke

References

Lin D., Shkedy Z., Yekutieli D., Amaratunga D., and Bijmens, L. (editors). (2012) Modeling Dose-response Microarray Data in Early Drug Development Experiments Using R. Springer.

Cheng, Y. and Church, G. M. (2000). Biclustering of expression data. In: Proceedings of the Eighth International Conference on Intelligent Systems for Molecular Biology, 1, 93-103.

See Also

[ORCME, plotIsomeans](#)

Examples

```
data(doseData)
data(geneData)

dirData <- monotoneDirection(geneData = geneData,doseData = doseData)

## direction of monotone trend
Direction <- dirData$direction
## Isotonic means for upward genes
incData <- as.data.frame(dirData$incData)
##Isotonic means for downward genes
decData <- as.data.frame(dirData$decData)
## observed data upward genes
obsIncData <- as.data.frame(dirData$obsincData)
## observed data for downward genes
obsDecData <- as.data.frame(dirData$obsdecData)
## isotonic means for all genes
isoMeans <- as.data.frame(dirData$arrayMean)
```

ORCME	<i>Order restricted clustering for dose-response trends in microarray experiments</i>
-------	---

Description

The function performs delta-clustering of a microarray data. It can be used for clustering of both the time-course or dose-response microarray data.

Usage

```
ORCME(DRdata, lambda, phi, robust=FALSE)
```

Arguments

DRdata	matrix of a microarray data with rows corresponding to genes and columns corresponding to time points or different doses
lambda	assumed proportion of coherence relative to the observed data, it ranges between 0 and 1. A lambda value of 1 considers the observed data as a cluster and lambda value of 0 finds every possible pattern within the data.
phi	minimum number of genes in a cluster
robust	logical variable that determines, if algorithm uses robust version based on median polish and absolute values, instead of mean square error. Default is FALSE.

Value

The matrix of classification into clusters: each row represents one gene and columns found clusters. The matrix consist of the Booleans values, in each row there is only one of them TRUE which means that the gene was classified into the respective cluster.

Author(s)

Adetayo Kasim, Martin Otava and Tobias Verbeke

References

Lin D., Shkedy Z., Yekutieli D., Amaratunga D., and Bijmens, L. (editors). (2012) Modeling Dose-response Microarray Data in EarlyDrug Development Experiments Using R. Springer.

Cheng, Y. and Church, G. M. (2000). Biclustering of expression data. In: Proceedings of the Eighth International Conference on Intelligent Systems for Molecular Biology, 1, 93-103.

See Also

[monotoneDirection](#), [plotIsomeans](#)

Examples

```

data(doseData)
data(geneData)

dirData <- monotoneDirection(geneData = geneData,doseData = doseData)
incData <- as.data.frame(dirData$incData)

print(orcme <- ORCME(DRdata=incData,lambda=0.15,phi=2))
orcmeRobust <- ORCME(DRdata=incData,lambda=0.15,phi=2, robust=TRUE)

# number of genes within cluster
colSums(orcme)
colSums(orcmeRobust)

```

plotCluster

Plotting the gene specific profiles for one given cluster of genes

Description

The function is plotting the profiles of the genes that belongs to the same cluster. It is not providing the clustering itself, just plotting the results of clustering from input. Optionally, the function can center the profiles around the gene-specific means.

Usage

```

plotCluster(DRdata, doseData, ORCMEoutput, clusterID,
zeroMean=FALSE, xlabel, ylabel, main="")

```

Arguments

DRdata	the microarray data with rows corresponding to genes and columns corresponding to time points or different doses
doseData	indicates the dose levels
ORCMEoutput	the matrix of classification into clusters: each row represents one gene and columns found clusters. The matrix consist of the Booleans values, in each row there is only one of them TRUE which means that the gene was classified into the respective gene
clusterID	id of the cluster to be plotted
zeroMean	if TRUE, it centers the gene profiles around the gene-specific means, default is FALSE
xlabel	a title for the x axis
ylabel	a title for the y axis
main	an overall title for the plot

Value

Plot of the gene specific profiles dependent one the dose level (or time point) that are classified into the given cluster.

Author(s)

Adetayo Kasim, Martin Otava and Tobias Verbeke

References

Lin D., Shkedy Z., Yekutieli D., Amaratunga D., and Bijmens, L. (editors). (2012) Modeling Dose-response Microarray Data in Early Drug Development Experiments Using R. Springer.

Cheng, Y. and Church, G. M. (2000). Biclustering of expression data. In: Proceedings of the Eighth International Conference on Intelligent Systems for Molecular Biology, 1, 93-103.

See Also

[ORCME](#), [plotIsomeans](#)

Examples

```
data(doseData)
data(geneData)

dirData <- monotoneDirection(geneData = geneData,doseData = doseData)
incData <- as.data.frame(dirData$incData)
ORCMEoutput <- ORCME(DRdata=incData,lambda=0.15,phi=2)

plotCluster(DRdata=incData,doseData=doseData, ORCMEoutput=ORCMEoutput,
clusterID=4,zeroMean=FALSE, xlabel="Dose",ylabel="Gene Expression")
```

plotIsomeans	<i>Plot of the observed gene expression and the isotonic means with respect to dose</i>
--------------	---

Description

The function is plotting the observed data points of the gene expression and isotonic means with respect to dose for one particular gene.

Usage

```
plotIsomeans(monoData, obsData, doseData, geneIndex)
```

Arguments

monoData	isotonic means with respect to dose for all genes
obsData	observed gene expression for all genes
doseData	indicates the dose levels
geneIndex	index of the gene to be plotted

Value

Plot of the data points and the isotonic means for each dose with the isotonic regression curve.

Author(s)

Adetayo Kasim, Martin Otava and Tobias Verbeke

References

Lin D., Shkedy Z., Yekutieli D., Amaratunga D., and Bijmens, L. (editors). (2012) Modeling Dose-response Microarray Data in Early Drug Development Experiments Using R. Springer.

Cheng, Y. and Church, G. M. (2000). Biclustering of expression data. In: Proceedings of the Eighth International Conference on Intelligent Systems for Molecular Biology, 1, 93-103.

See Also

[ORCME](#), [monotoneDirection](#)

Examples

```
data(doseData)
data(geneData)

dirData <- monotoneDirection(geneData = geneData,doseData = doseData)
incData <- as.data.frame(dirData$incData)
obsIncData <- as.data.frame(dirData$obsincData)

## gene-specific profile plot
plotIsomeans(monoData=incData,obsData=obsIncData,doseData=
doseData, geneIndex=10)
```

plotLambda	<i>Plot the variety of the properties dependent on the proportion of heterogeneity in observed data set</i>
------------	---

Description

This function provides the plots of the dependency of the variety of properties on the proportion of heterogeneity in observed data set. It is not using the clustering as simple input, but it is also computing additional properties. The function can plot within cluster sum of squares, number of cluster, penalized within cluster sum of squares, Calsanzik and Harabax index and Hartigan index.

Usage

```
plotLambda(lambdaChoiceOutput,output)
```

Arguments

lambdaChoiceOutput	the output of the function resampleORCME
output	the variable that determines which output would be plotted, the values are "wss" for the cluster sum of squares, "ncluster" for the number of cluster, "pwss" for the penalized within cluster sum of squares, "ch" for the Calsanzik and Harabax index and "h" for the Hartigan index

Value

A plot of one of the properties mentioned above dependent on the proportion of heterogeneity. The confidence intervals are plotted instead of the point estimates.

Author(s)

Adetayo Kasim, Martin Otava and Tobias Verbeke

References

Lin D., Shkedy Z., Yekutieli D., Amaratunga D., and Bijnens, L. (editors). (2012) Modeling Dose-response Microarray Data in Early Drug Development Experiments Using R. Springer.

Cheng, Y. and Church, G. M. (2000). Biclustering of expression data. In: Proceedings of the Eighth International Conference on Intelligent Systems for Molecular Biology, 1, 93-103.

See Also

[ORCME](#), [resampleORCME](#)

Examples

```

data(doseData)
data(geneData)

dirData <- monotoneDirection(geneData = geneData, doseData = doseData)
incData <- as.data.frame(dirData$incData)

lambdaVector <- c(0.05, 0.50, 0.95)

lambdaChoiceOutput <- resampleORCME(clusteringData=incData, lambdaVector=lambdaVector)
plotLambda(lambdaChoiceOutput, output="wss")
plotLambda(lambdaChoiceOutput, output="ncluster")
plotLambda(lambdaChoiceOutput, output="pwss")
plotLambda(lambdaChoiceOutput, output="ch")
plotLambda(lambdaChoiceOutput, output="h")

```

resampleORCME

Estimation of the proportion of the heterogeneity in the observed data for clustering

Description

The function is computing within cluster sum of squares for given proportion of heterogeneity. Minimal number of genes per cluster is fixed as 2. The sum of squares is computed through resampling the 100 data sets with 100 genes randomly sampled with replacement from the reduced expression data.

Usage

```
resampleORCME(clusteringData, lambdaVector, robust=FALSE)
```

Arguments

<code>clusteringData</code>	the microarray data with rows corresponding to genes and columns corresponding to time points or different doses
<code>lambdaVector</code>	vector of assumed proportions of of heterogeneity of the observed data, it ranges between 0 and 1. A lambda value of 1 considers the observed data as a cluster and lambda value of 0 finds every possible pattern within the data
<code>robust</code>	logical variable that determines, if algorithm uses robust version based on median polish and absolute values, instead of mean square error. Default is FALSE.

Value

A list of matrices that represent one of the 100 iterations. Every matrix consist of the columns

lambda	vector of the proportions of heterogeneity given as input
WSS	within clusters sum of squares for given proportion of heterogeneity
TSS	total clusters sum of squares for given proportions of heterogeneity
nc	number of clusters as a function for given proportions of heterogeneity

Author(s)

Adetayo Kasim, Martin Otava and Tobias Verbeke

References

Lin D., Shkedy Z., Yekutieli D., Amaratunga D., and Bijmens, L. (editors). (2012) Modeling Dose-response Microarray Data in Early Drug Development Experiments Using R. Springer.

Cheng, Y. and Church, G. M. (2000). Biclustering of expression data. In: Proceedings of the Eighth International Conference on Intelligent Systems for Molecular Biology, 1, 93-103.

See Also

[ORCME](#), [plotLambda](#)

Examples

```
data(doseData)
data(geneData)

dirData <- monotoneDirection(geneData = geneData, doseData = doseData)
incData <- as.data.frame(dirData$incData)

lambdaVector <- c(0.05, 0.50, 0.95)

resampleORCME(clusteringData=incData, lambdaVector=lambdaVector, robust=FALSE)
```

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