Package 'GeneExpressionSignature'

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Title Gene Expression Signature based Similarity Metric
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Description This package gives the implementations of the gene expression signature and its distance to each. Gene expression signature is represented as a list of genes whose expression is correlated with a biological state of interest. And its distance is defined using a nonparametric, rank based pattern-matching strategy based on the Kolmogorov-Smirnov statistic. Gene expression signature and its distance can be used to detect similarities among the signatures of drugs, diseases, and biological states of interest.
Depends R (>= 2.13), Biobase, PGSEA
Suggests apcluster, GEO query
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LazyLoad yes
biocViews Bioinformatics, GeneExpression
R topics documented:
GeneExpressionSignature getRLs RankMerging ScoreGSEA ScorePGSEA SignatureDistance
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exampleSet

sample data, a subset of the C-MAP

Description

sample data, a subset of the C-MAP as , which is a collection of 50 genome-wide transcriptional expression data from cultured human cells treated with 15 different small molecules.

Usage

data(exampleSet)

Format

A ExpressionSet: assay data represents the 50 genome-wide transcriptional expression data, phenotypic data describes 15 different small molecules corresponds to the expression data (assay data).

References

http://www.sciencemag.org/content/313/5795/1929.short Lamb et al., The Connectivity Map: Using Gene-Expression Signatures to Connect Small Molecules, Genes, and Disease, science 2006

Examples

data(exampleSet)

GeneExpressionSignature

Introduction to the GeneExpressionSignature Package

Description

The **GeneExpressionSignature** add-on is an implementation of computing distances among preprocessed gene-expression profiles of samples for R. The distances can be used to detect similarities among the signatures of drugs, diseases, and biological states of interest, and construct connectivity map.

Details

This package contains functions for the distances computation based on gene expression signature. First, list of genes is ranked according to their expression ratios to produce the Prototype Ranked List (PRL). Second, all the PRLs with the same state are aggregated by RankMerging functions. Finally, all the ranked lists are made as one input of the ScorePGSEA and ScoreGSEA functions to compute the pairwise distances.

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getRLs

Convert gene expression profiles to a ranked list (data preprocessing)

Description

Sorting the microarray probe-set identifiers according to the differential expression values with respect to the untreated hybridization to obtain a ranked list. Gene-expression profiles in are represented in a nonparametric fashion.

Usage

```
getRLs(control, treatment)
```

Arguments

control a matrix, including the vehicle control gene expression profiles corresponding

to the treatment gene expression profiles.

treatment a matrix, is composed of gene expression profiles.

Details

The genes on the array are rank-ordered according to their differential expression relative to the control. First, control and treatment values less than a primary threshold value (quartile) were set to that threshold value. Finally, probe sets were ranked in descending order of d, where d is the ratio of the corresponding treatment-to-control values. For probe sets where d=1, a lower threshold was applied to the original difference values and a new treatment to control ratio (d') calculated. These probe sets were then sub-sorted in descending order of d.

Value

A matrix is composed of ranked lists, a ranked list represents the corresponding gene expression profiles.

Examples

```
if (require(GEOquery)){
    #treatment gene-expression profiles
    GSM118720 <- getGEO(filename=system.file("extdata/GSM118720.soft",package=
    "GeneExpressionSignature"))
    #control gene-expression profiles
    GSM118721 <- getGEO(filename=system.file("extdata/GSM118721.soft",package=
    "GeneExpressionSignature"))
    #data ranking according to the different expression values
    control <- as.matrix(as.numeric(Table(GSM118721)[,2]))
    treatment <- as.matrix(as.numeric(Table(GSM118720)[,2]))
    ranked_list <-getRLs(control,treatment)
}</pre>
```

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RankMerging	Merging the ranker lists with the same labels of the biological states into a single list with the Iorio's method.

Description

Merging the assay data according to phenotypic data of the input ExpressionSet. Each group of the ranked lists with the same phenotypic data is aggregated into a single list, return it as an ExpressionSet object.

Usage

```
RankMerging(exprSet, MergingDistance = c("Spearman", "Kendall"), weighted = TRUE)
```

Arguments

exprSet an ExpressionSet object, each column of assay data represents a ranked list ob-

tained by preprocessing the corresponding gene expression profile, and phenotypic data represents the short description (characteristics of gene expression

profile, such as the drug type, the disease state) about the assay data.

MergingDistance

distance to be used which "measures" the similarity of ordered lists, the default

is "Spearman"

weighted there are tow rank merging approaches for two cases: if weighted=FALSE, all

ranked list with the same biological state are treated equally important, a simple but useful method average ranking technique is selected; otherwise, weighted=TRUE, each individual ranked lists has its own ranked weights, this takes the iterative

rank-aggregating algorithm, default is TRUE.

Details

The krubor function is used in the aggregating procedure. And the following methods are used in the implementation: a measure of the distance between two ranked lists (Spearman's Footrule), a method to merge two or more ranked lists the (Borda Merging Method), and a algorithm to obtain a single ranked list from a set of them in a hierarchical way (the Kruskal Algorithm). If choose Kendall as distance, the effectiveness of this function is certainly limited by the size of the merging problem.

See Also

 ${\tt Signature Distance}$

Examples

#load the sample expressionSet
data(exampleSet)

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Merging each group of the ranked lists in the exampleSet with the same phenotypic data into a single PRL
MergingSet=RankMerging(exampleSet, "Spearman", weighted=TRUE)

ScoreGSEA Compute pairwise distances between samples with method in package GSEA

Description

Compute pairwise distances between sample according to their (Prototype Ranked List) PRL, a N x N distance matrix is generated by calling this function, N is the length of PRL.

Usage

ScoreGSEA(MergingSet, SignatureLength, ScoringDistance = c("avg", "max"), p.value = F)

Arguments

MergingSet

an ExpressionSet object. The assay data represents the PRLs of the samples, each column represents one PRL. The number of sample of this argument must be greater than 1, otherwise, this function is not meaningful.

SignatureLength

the length of "gene signature". In order to compute pairwise distances among samples, genes lists are ranked according to the gene expression ratio (fold change). And the "gene signature" includes the most up-regulated genes (near the top of the list) and the most down-regulated genes (near the bottom of the list).

ScoringDistance

the distance measurements between PRLs: the Average Enrichment Score Distance (avg), and the Maximum Enrichment Score Distance (max).

p.value

logical, if TRUE return a matrix of p.values of the distance matrix, default FALSE

Details

Once the PRL obtained for each sample, the distances between samples are calculated base on gene signature, including the expression of genes that seemed to consistently vary in response to the across different experimental conditions (e.g., different cell lines and different dosages). We take two distance measurements between PRLs: the Average Enrichment-Score Distance Davg=(TESx,y+TESy,x)/2, and the Maximum Enrichment-Score Distance Dmax=Min(TESx,y,TESy,x)/2. The avg is more stringent than max, where max is more sensitive to weak similarities, with lower precision but large recall.

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Value

an distance-matrix, the max distance is more sensitive to weak similarities, providing a lower precision but a larger recall.

If p.value is set to TRUE, then a list is returned that consists of the distance matrix as well as their p.values, otherwise, without p.vlues in the result.

See Also

ScorePGSEA, SignatureDistance

ds=ScoreGSEA(MergingSet,250,"avg")

Examples

```
# load the sample expressionSet
data(exampleSet)

# Merging each group of the ranked lists in the exampleSet with the same phenotypic data into a single PRL
MergingSet=RankMerging(exampleSet, "Spearman")

# get the distance matrix
```

ScorePGSEA

Compute pairwise distances between samples with method in package PGSEA

Description

Compute pairwise distances between sample according to their (Prototype Ranked List) PRL, get a N x N distance matrix is generated by calling this function, N is the length of PRL.

Usage

```
ScorePGSEA(MergingSet, SignatureLength, ScoringDistance = c("avg", "max"), p.value = F)
```

Arguments

MergingSet

an ExpressionSet object. The assay data represents the PRLs of the samples, each column represents one PRL. The number of sample must be greater than 1, oherwise, this function is not meaningful.

SignatureLength

the length of "gene signature". In order to compute pairwise distances among samples, genes lists are ranked according to the gene expression ratio (fold change). And the "gene signature" includes the most up-regulated genes (near the top of the list) and the most down-regulated genes (near the bottom of the list).

ScoringDistance

the distance measurements between PRLs: the Average Enrichment Score Distance (avg), or the Maximum Enrichment Score Distance (max).

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p.value

logical, if TRUE return a matrix of p.values of the distance matrix, default FALSE

Details

This function has the same function with ScoreGSEA, just with different methods.

See Also

ScoreGSEA, SignatureDistance

Examples

```
# load the sample expressionSet
data(exampleSet)

# Merging each group of the ranked lists in the exampleSet with the same phenotypic data into a single PRL
MergingSet=RankMerging(exampleSet, "Spearman")

# get the distance matrix
ds=ScorePGSEA(MergingSet, 250, ScoringDistance="avg")
```

SignatureDistance

Compute pairwise distances comprehensively.

Description

This function integrated the function for rank merging and distance scoring, we can do the rank merging and distance scoring simply with it.

Usage

SignatureDistance(exprSet, SignatureLength, MergingDistance = c("Spearman", "Kendall"), ScoringMethod

Arguments

exprSet

an ExpressionSet object, each column of assay data represents a ranked list obtained by preprocessing the corresponding gene expression profile, and phenotypic data represents the short description (characteristics of gene expression profile, such as the drug type, the disease state) about the assay data.

SignatureLength

the length of "gene signature". In order to compute pairwise distances among samples, genes lists are ranked according to the gene expression ratio (fold change). And the "gene signature" includes the most up-regulated genes (near the top of the list) and the most down-regulated genes (near the bottom of the list).

MergingDistance

distance to be used which "measures" the similarity of ordered lists, Spearman or Kendall

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ScoringMethod method to be used to perform distance scoring, GSEA or PGSEA ScoringDistance

the distance measurements between PRLs: the Average Enrichment Score Dis-

tance (avg), and the Maximum Enrichment Score Distance (max).

weighted there are tow rank merging approaches for two cases: if weighted=FALSE, all

ranked list with the same biological state are treated equally important, a simple but useful method average ranking technique is selected; otherwise, weighted=TRUE, each individual ranked lists has its own ranked weights, this takes the iterative

rank-aggregating algorithm, default is TRUE.

... additional arguments can be passed to the internal procedures

See Also

RankMerging, ScoreGSEA, ScorePGSEA

Examples

#load the sample expressionSet
data(exampleSet)

#distance scoring

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