Package ‘STATegRa’

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Function to compute a `bioDistclass` object from profile data and a mapping. For details of the process see the user’s guide, but briefly the process involves using the mapping to identify reference features appropriate to each surrogate feature (if any), aggregating the surrogate data into pseudo-data for each reference feature, and then calculating the correlation distance between the reference features according to the surrogate data.
bioDist

Usage

```r
bioDist(referenceFeatures=NULL, reference=NULL, mapping=NULL,
referenceData=NULL, surrogateData=NULL, filtering=NULL,
noMappingDist=NA, distance="spearman", aggregation="sum",
maxitems=NULL, selectionRule="maxFC", expfac=NULL,
name=NULL, ...)
```

Arguments

**referenceFeatures**
subset of features to be considered for the computation of the distances. If NULL then the features are first gathered from the features in referenceData. If referenceData is not provided then the list of features are gathered from mapping (bioMap class) and using the reference.

**reference**
A character indicating the variable that is being used as features to compute distance between

**mapping**
The mapping between feature types

**referenceData**
ExpressionSet object with the data from the reference features.

**surrogateData**
ExpressionSet object with the data from the surrogate features.

**filtering**
A filtering for the bioMap class. To be implemented.

**noMappingDist**
Distance value to be used when a reference feature do not map to any surrogate feature. If "max", maximum indirect distance among the rest of reference features is taken. If NA, distance weights are re-scaled so this surrogate association is not considered. If a number then the missing values are replaces with that value.

**distance**
Distance between features to be computed. Possible values are "pearson", "kendall", "spearman", "euclidean", "maximum", "manhattan", "canberra", "binary" and "minkowski". Default is "spearman".

**aggregation**
Action to perform when a reference feature maps to more than one surrogate feature. Options are "max", "sum", "mean" or "median" and the the values are aggregated according to the chosen statistic.

**maxitems**
The maximum number of surrogate features per reference feature to be used, selected according to "selectionRule" parameter. Default is 2.

**selectionRule**
Rule to select the surrogate features to be used (the number is determined by "maxitems"). It can be one of the following: (1) "maxcor" those presenting maximum correlation with corresponding main feature; in this case "referenceData" must be provided and the columns must overlap in at least 3 samples; (2) "maxmean": average across samples is computed and those features with higher mean are selected; case (3) is simmilar to (2) but considering other statistics: "maxmedian", "maxdiff", "maxFC", "sd", "ee".

**expfac**
Not in use yet.

**name**
Character that describes the nature of the bioDist class computed

... extra arguments passed to `dist`, eg "p=value" for the power used if calculating minkowski distance
**Value**

An object of class bioDistclass containing distances between the features in surrogateData.

**Author(s)**

David Gomez-Cabrero

**Examples**

```r
data(STATegRa_S1)
data(STATegRa_S2)
require(Biobase)

# Truncate data for brevity
Block1 <- Block1[1:100,]
Block2 <- Block2[1:100,]

## Create ExpressionSets
mRNA.ds <- createOmicsExpressionSet(Data=Block1,pData=ed,pDataDescr=c("classname"))
miRNA.ds <- createOmicsExpressionSet(Data=Block2,pData=ed,pDataDescr=c("classname"))

## Create the bioMap
map.gene.miRNA<-bioMap(name = "Symbol-miRNA",
                         metadata = list(type_v1="Gene",type_v2="miRNA",
                                         source_database="targetscan.Hs.eg.db",
                                         data_extraction="July2014"),
                         map=mapdata)

# Create Gene-gene distance computed through miRNA data
bioDistmiRNA<-bioDist(referenceFeatures = rownames(Block1),
                       reference = "Var1",
                       mapping = map.gene.miRNA,
                       surrogateData = miRNA.ds, ### miRNA data
                       referenceData = mRNA.ds, ### mRNA data
                       maxitems=2,
                       selectionRule="sd",
                       expfac=NULL,
                       aggregation = "sum",
                       distance = "spearman",
                       noMappingDist = 0,
                       filtering = NULL,
                       name = "mRNAbymiRNA")

# Create Gene-gene distance through mRNA data
bioDistmRNA<-new("bioDistclass",
                 name = "mRNAbymRNA",
                 distance = cor(t(exprs(mRNA.ds)),method="spearman"),
                 map.name = "id",
                 map.metadata = list(),
                 params = list())

          # Generation of the list of Surrogated distances.
```
bioDistclass

bioDistList <- list(bioDistmRNA, bioDistmiRNA)
sample.weights <- matrix(0, 4, 2)
sample.weights[, 1] <- c(0, 0.33, 0.67, 1)
sample.weights[, 2] <- c(1, 0.67, 0.33, 0)

##### Generation of the list of bioDistWclass objects.

bioDistWList <- bioDistW(referenceFeatures = rownames(Block1),
                        bioDistList = bioDistList,
                        weights = sample.weights)

##### Plot of distances.

bioDistWPlot(referenceFeatures = rownames(Block1),
              listDistW = bioDistWList,
              method.cor = "spearman")

##### Computing the matrix of features/distances associated.

fm <- bioDistFeature(Feature = rownames(Block1)[1],
                     listDistW = bioDistWList,
                     threshold.cor = 0.7)

bioDistFeaturePlot(data = fm)

---

bioDistclass

Description

Class to manage mappings between genomic features.

Usage

bioDistclass(name, distance, map.name, map.metadata, params)

Arguments

- name: Name assigned to the object
- distance: Matrix giving the distance between features
- map.name: Charactering giving the name of the bioMap object used to compute the distance
- map.metadata: List of parameters used to generate the mapping
- params: List of parameters used to generate the distance
bioDistFeature

Description

Function that computes for a given selected feature the closest features given a selected set of weighted distances.

Usage

bioDistFeature(Feature, listDistW, threshold.cor)

Arguments

Feature Feature A selected as a reference.
listDistW A list of bioDistWclass objects. All the objects must contain the Feature A selected and all of them must contain the same set of features.
threshold.cor A threshold to select the features associated to Feature A

Value

Matrix with the associated features given the different weighted distances considered

Author(s)

David Gomez-Cabrero

Examples

data(STATegRa_S1)
data(STATegRa_S2)
require(Biobase)

# Truncate data for brevity
Block1 <- Block1[1:100,]
Block2 <- Block2[1:100,]

## Create ExpressionSets
mRNA.ds <- createOmicsExpressionSet(Data=Block1,pData=ed,pDataDescr=c("classname"))
miRNA.ds <- createOmicsExpressionSet(Data=Block2,pData=ed,pDataDescr=c("classname"))

## Create the bioMap
map.gene.miRNA<-bioMap(name = "Symbol-miRNA",
                        metadata = list(type_v1="Gene",type_v2="miRNA",
                                        source_database="targetscan.Hs.eg.db",
                                        data_extraction="July2014"),
                        map=mapdata)

# Create Gene-gene distance computed through miRNA data
bioDistmiRNA<-bioDist(referenceFeatures = rownames(Block1),
  reference = "Var1",
  mapping = map.gene.miRNA,
  surrogateData = miRNA.ds, ### miRNA data
  referenceData = mRNA.ds, ### mRNA data
  maxitems=2,
  selectionRule="sd",
  expfac=NULL,
  aggregation = "sum",
  distance = "spearman",
  noMappingDist = 0,
  filtering = NULL,
  name = "mRNAbymiRNA")

# Create Gene-gene distance through mRNA data
bioDistmRNA<-new("bioDistclass",
       name = "mRNAbymRNA",
       distance = cor(t(exprs(mRNA.ds)),method="spearman"),
       map.name = "id",
       map.metadata = list(),
       params = list())

###### Generation of the list of Surrogated distances.
bioDistList<-list(bioDistmRNA,bioDistmiRNA)
sample.weights<-matrix(0,4,2)
sample.weights[,1]<-c(0,0.33,0.67,1)
sample.weights[,2]<-c(1,0.67,0.33,0)

###### Generation of the list of bioDistWclass objects.
bioDistWList<-bioDistW(referenceFeatures = rownames(Block1),
  bioDistList = bioDistList,
  weights=sample.weights)

###### Plot of distances.
bioDistWPlot(referenceFeatures = rownames(Block1) ,
  listDistW = bioDistWList,
  method.cor="spearman")

###### Computing the matrix of features/distances associated.
fm<-bioDistFeature(Feature = rownames(Block1)[1] ,
  listDistW = bioDistWList,
  threshold.cor=0.7)
bioDistFeaturePlot(data=fm)
bioDistFeaturePlot

Description

Function that plots the results from a bioDistFeature analysis

Usage

bioDistFeaturePlot(data)

Arguments

data Matrix produced by bioDistFeature

Value

Generates a heatmap plot

Author(s)

David Gomez-Cabrero

Examples

data(STATegRa_S1)
data(STATegRa_S2)
require(Biobase)

# Truncate data for brevity
Block1 <- Block1[1:100,]
Block2 <- Block2[1:100,]

## Create ExpressionSets
mRNA.ds <- createOmicsExpressionSet(Data=Block1,pData=ed,pDataDescr=c("classname"))
miRNA.ds <- createOmicsExpressionSet(Data=Block2,pData=ed,pDataDescr=c("classname"))

## Create the bioMap
map.gene.miRNA<-bioMap(name = "Symbol-miRNA",
                        metadata = list(type_v1="Gene",type_v2="miRNA",
                                        source_database="targetscan.Hs.eg.db",
                                        data_extraction="July2014"),
                        map=mapdata)

# Create Gene-gene distance computed through miRNA data
bioDistmiRNA<-bioDist(referenceFeatures = rownames(Block1),
                       reference = "Var1",
                       mapping = map.gene.miRNA,
                       surrogateData = miRNA.ds, ### miRNA data
                       referenceData = mRNA.ds, ### mRNA data
                       maxitems=2,
                       selectionRule="sd",
                       expfac=NULL,
                       aggregation = "sum",
                       distance = "spearman",
                       noMappingDist = 0,
# Create Gene-gene distance through mRNA data
bioDistmRNA<-new("bioDistclass",
    name = "mRNAbymRNA",
    distance = cor(t(exprs(mRNA.ds)),method="spearman"),
    map.name = "id",
    map.metadata = list(),
    params = list())

##### Generation of the list of Surrogated distances.
bioDistList<-list(bioDistmRNA,bioDistmiRNA)
sample.weights<-matrix(0,4,2)
sample.weights[,1]<-c(0,0.33,0.67,1)
sample.weights[,2]<-c(1,0.67,0.33,0)

##### Generation of the list of bioDistWclass objects.
bioDistWList<-bioDistW(referenceFeatures = rownames(Block1),
    bioDistList = bioDistList,
    weights=sample.weights)

##### Plot of distances.
bioDistWPlot(referenceFeatures = rownames(Block1) ,
    listDistW = bioDistWList,
    method.cor="spearman")

##### Computing the matrix of features/distances associated.
fm<-bioDistFeature(Feature = rownames(Block1)[1] ,
    listDistW = bioDistWList,
    threshold.cor=0.7)
bioDistFeaturePlot(data=fm)

### Description

Function that computes weighted distances between a list of bioDistclass objects.

### Usage

bioDistW(referenceFeatures, bioDistList, weights)
Arguments

**referenceFeatures**

The set of features that weighted distance is computed between.

**bioDistList**

A list of bioDistclass objects. All the objects must contain the set of features selected.

**weights**

A matrix where the number of columns equals the number of elements included in the bioDistList list.

Value

Returns a list of bioDistWclass objects. Each element in the list returns the weighted distance associated to each row in the "weights" matrix.

Author(s)

David Gomez-Cabrero

Examples

data(STATegRa_S1)
data(STATegRa_S2)
require(Biobase)

# Truncate data for brevity
Block1 <- Block1[1:100,]
Block2 <- Block2[1:100,]

## Create ExpressionSets
mRNA.ds <- createOmicsExpressionSet(Data=Block1,pData=ed,pDataDescr=c("classname"))
miRNA.ds <- createOmicsExpressionSet(Data=Block2,pData=ed,pDataDescr=c("classname"))

## Create the bioMap
map.gene.miRNA<-bioMap(name = "Symbol-miRNA",
metadata = list(type_v1="Gene",type_v2="miRNA",
source_database="targetscan.Hs.eg.db",
data_extraction="July2014"),
map=mapdata)

# Create Gene-gene distance computed through miRNA data
bioDistmiRNA<-bioDist(referenceFeatures = rownames(Block1),
reference = "Var1",
mapping = map.gene.miRNA,
surrogateData = miRNA.ds, ### miRNA data
referenceData = mRNA.ds, ### mRNA data
maxitems=2,
selectionRule="sd",
expfac=NULL,
aggregation = "sum",
distance = "spearman",
noMappingDist = 0,
filtering = NULL,
# Create Gene-gene distance through mRNA data

```r
distmRNA <- new("bioDistClass",
    name = "mRNAbymRNA",
    distance = cor(t(exprs(mRNA.ds)),method="spearman"),
    map.name = "id",
    map.metadata = list(),
    params = list())
```

##### Generation of the list of Surrogated distances.

```r
distList <- list(distmRNA, distmiRNA)
sample.weights <- matrix(0,4,2)
sample.weights[,1] <- c(0, 0.33, 0.67, 1)
sample.weights[,2] <- c(1, 0.67, 0.33, 0)
```

##### Generation of the list of bioDistWclass objects.

```r
distWList <- bioDistW(referenceFeatures = rownames(Block1),
    bioDistList = distList,
    weights = sample.weights)
```

##### Plot of distances.

```r
distWPlot(referenceFeatures = rownames(Block1),
    listDistW = distWList,
    method.cor = "spearman")
```

##### Computing the matrix of features/distances associated.

```r
fm <- bioDistFeature(Feature = rownames(Block1)[1],
    listDistW = distWList,
    threshold.cor = 0.7)

bioDistFeaturePlot(data = fm)
```

### Description

Function that plots the "distance relation" between features computed through different surrogate features.

### Usage

```r
bioDistWPlot(referenceFeatures, listDistW, method.cor)
```
bioDistWPlot

Arguments

referenceFeatures
The set of features to be used.

listDistW
A list of bioDistWclass objects.

method.cor
Method to compute distances between the elements in the listDistW. The default is spearman correlation.

Value
Makes a plot with the projected distance between the listDistW objects.

Author(s)
David Gomez-Cabrero

Examples

data(STAteRa_S1)
data(STAteRa_S2)
require(Biobase)

# Truncate data for brevity
Block1 <- Block1[,1:100,]
Block2 <- Block2[,1:100,]

## Create ExpressionSets
mRNA.ds <- createOmicsExpressionSet(Data=Block1,pData=ed,pDataDescr=c("classname"))
miRNA.ds <- createOmicsExpressionSet(Data=Block2,pData=ed,pDataDescr=c("classname"))

## Create the bioMap
map.gene.miRNA<-bioMap(name = "Symbol-miRNA",
                     metadata = list(type_v1="Gene",type_v2="miRNA",
                                     source_database="targetscan.Hs.eg.db",
                                     data_extraction="July2014"),
                     map=mapdata)

# Create Gene-gene distance computed through miRNA data
bioDistmiRNA<-bioDist(referenceFeatures = rownames(Block1),
                     reference = "Var1",
                     mapping = map.gene.miRNA,
                     surrogateData = miRNA.ds, ### miRNA data
                     referenceData = mRNA.ds, ### mRNA data
                     maxitems=2,
                     selectionRule="sd",
                     expfac=NULL,
                     aggregation = "sum",
                     distance = "spearman",
                     noMappingDist = 0,
                     filtering = NULL,
                     name = "mRNAbymiRNA")
# Create Gene-gene distance through mRNA data

```r
bioDistmRNA <- new("bioDistclass",
    name = "mRNAbymRNA",
    distance = cor(t(exprs(mRNA.ds)), method = "spearman"),
    map.name = "id",
    map.metadata = list(),
    params = list())
```

##### Generation of the list of Surrogated distances.

```r
bioDistList <- list(bioDistmRNA, bioDistmiRNA)
```

```r
sample.weights <- matrix(0, 4, 2)
sample.weights[, 1] <- c(0, 0.33, 0.67, 1)
sample.weights[, 2] <- c(1, 0.67, 0.33, 0)
```

##### Generation of the list of bioDistWclass objects.

```r
bioDistWList <- bioDistW(referenceFeatures = rownames(Block1),
    bioDistList = bioDistList,
    weights = sample.weights)
```

##### Plot of distances.

```r
bioDistWPlot(referenceFeatures = rownames(Block1),
    listDistW = bioDistWList,
    method.cor = "spearman")
```

##### Computing the matrix of features/distances associated.

```r
fm <- bioDistFeature(Feature = rownames(Block1)[1],
    listDistW = bioDistWList,
    threshold.cor = 0.7)
```

```r
bioDistFeaturePlot(data = fm)
```

---

### bioMap

**Description**

Function to generate a bioMap object.

**Usage**

```r
bioMap(name, metadata, map)
```

**Arguments**

- `name` : Name to assign the object
metadata A list with information of the mapping. Elements expected in the list are: (1) "type_v1" and "type_v2", refer to the nature of the features mapped; a vocabulary we recommend is "gene", "mRNA", "miRNA", "proteins", etc. (2) "source_database", provides information on the source of the mapping; from a specific data-base e.g. "targetscan.Hs.eg.db" to a genomic location mapping. (3) "data_extraction" stores information on the data the mapping was generated or downloaded.

map A data.frame object storing the mapping. The data.frame may include an unlimited number of columns, however the first column must be named "Var1" and refer to the elements of "type_v1" and similarly for the second column ("Var2","type_v2").

Value
An object of class bioMap

Author(s)
David Gomez-Cabrero

Examples
```
data(STATegRa_S2)
map.gene.miRNA<-bioMap(name = "Symbol-miRNA",
    metadata = list(type_v1="Gene",type_v2="miRNA",
        source_database="targetscan.Hs.eg.db",
        data_extraction="July2014"),
    map=mapdata)
```

Description
Stores the results of any of the omicsPCA analyses.

Slots
- `InitialData` List of ExpressionSets, one for each set of omics data
- `Names` Character vector giving names for the input data
- `preprocessing` Character vector describing the preprocessing applied to the data
- `preproData` List of matrices containing data after preprocessing
- `caMethod` Character giving the component analysis method name
- `commonComps` Numeric giving the number of common components
- `distComps` Numeric vector giving the number of distinctive components for each block
- `scores` List of matrices of common and distinctive scores
**combiningMappings**

loadings  List of matrices of common and distinctive loadings
VAF  List of matrices indicating VAF (Variability Explained For) for each component in each block of data
others  List containing other miscellaneous information specific to different SCA methods

**Author(s)**

Patricia Sebastian Leon

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**combiningMappings**  *combiningMappings, combining several mappings for use in the omicsNPC function*

**Description**

This function combines several annotation so that measurements across different datasets are mapped to the same reference elements (e.g., genes). The annotations should all be either data frame / matrices, named vectors/lists, or bioMap objects. See the examples for further details

**Usage**

`combiningMappings(mappings, reference = NULL, retainAll = FALSE)`

**Arguments**

- `mappings`  List of annotations.
- `reference`  If the annotations are data frame, matrices or bioMap objects, the name of the column containing the reference elements
- `retainAll`  Logical, if set to TRUE measurements that have no counterparts in other datasets are retained

**Value**

A data frame encoding the mapping across several dataset

**Author(s)**

Vincenzo Lagani

**References**

Nestoras Karathanasis, Ioannis Tsamardinos and Vincenzo Lagani. omicsNPC: applying the Non-Parametric Combination methodology to the integrative analysis of heterogeneous omics data. Submitted to PlosONE.
Example 1
#Mapping with data frames
mRNA <- data.frame(gene = rep(c('G1', 'G2', 'G3'), each = 2), probeset = paste('p', 1:6, sep = ''));
methylation <- data.frame(gene = c(rep('G1', 3), rep('G2', 4)),
                         methy = paste('methy', 1:7, sep = ''));
miRNA <- data.frame(gene = c(rep('G1', 2), rep('G2', 1), rep('G3', 2)),
                     miR = c('miR1', 'miR2', 'miR1', 'miR1', 'miR2'));
mappings <- list(mRNA = mRNA, methylation = methylation, miRNA = miRNA);
combiningMappings(mappings = mappings, retainAll = TRUE)

Example 2
#Mapping with character vectors
mRNA <- rep(c('G1', 'G2', 'G3'), each = 2);
names(mRNA) = paste('p', 1:6, sep = '');
methylation <- c(rep('G1', 3), rep('G2', 4));
names(methylation) = paste('methy', 1:7, sep = '');
miRNA <- c(rep('G1', 2), rep('G2', 1), rep('G3', 2));
names(miRNA) = c('miR1', 'miR2', 'miR1', 'miR1', 'miR2');
mappings <- list(mRNA = mRNA, methylation = methylation, miRNA = miRNA);
combiningMappings(mappings = mappings, retainAll = TRUE)

createOmicsExpressionSet

createOmicsExpressionSet

Description
This function allows the user to create a ExpressionSet object from a matrix representing an omics dataset. It allows to include the experimental design and annotation in the ExpressionSet object.

Usage
createOmicsExpressionSet(Data, pData = NULL, pDataDescr = NULL,
                          feaData = NULL, feaDataDescr = NULL)

Arguments
Data Omics data
pData Data associated with the samples/phenotype
pDataDescr Description of the phenotypic data
feaData Data associated with the variables/features annotation
feaDataDescr Description of the feature annotation

Details
In Data matrix, samples has to be in columns and variables has to be in rows
getInitialData

Value

ExpressionSet with the data provided

Author(s)

Patricia Sebastian-Leon

Examples

data(STATegRa_S3)
B1 <- createOmicsExpressionSet(Data=Block1.PCA,pData=ed.PCA,
pDataDescr=c("classname"))
B2 <- createOmicsExpressionSet(Data=Block2.PCA,pData=ed.PCA,
pDataDescr=c("classname"))

getInitialData Retrieve initial data from caClass objects

Description

Generic function to retrieve the initial data used for by omicsCompAnalysis from a caClass-class object

Usage

getInitialData(x, block=NULL)

Arguments

x caClass-class object.
block Character indicating the block of data to be returned. It can be specified by the position of the block ("1" or "2") or the name assigned in the caClass-class object. If it is NULL both blocks are displayed.

Value

The requested data block or blocks

Author(s)

Patricia Sebastian-Leon

See Also

omicsCompAnalysis, caClass-class
Examples

data("STATegRa_S3")
B1 <- createOmicsExpressionSet(Data=Block1.PCA, pData=ed.PCA,
   pDataDescr=c("classname"))
B2 <- createOmicsExpressionSet(Data=Block2.PCA,
   pData=ed.PCA, pDataDescr=c("classname"))

# Omics components analysis
res <- omicsCompAnalysis(Input=list(B1, B2), Names=c("expr", "mirna"),
   method="DISCOSCA", Rcommon=2, Rspecific=c(2, 2),
   center=TRUE, scale=TRUE, weight=TRUE)
getInitialData(res)
getInitialData(res, block="expr")

getLoadings

Retrieve component analysis loadings

Description

Generic function to retrieve loadings (common and distinctive) found by `omicsCompAnalysis` on a `caClass-class` object.

Usage

getLoadings(x, part=NULL, block=NULL)

Arguments

x `caClass-class` object.

part Character indicating whether "common" or "distinctive" loadings should be displayed

block Character indicating the block of data for which the loadings will be given. It can be specified by the position of the block ("1" or "2") or the name assigned in the `caClass-class` object. If it is NULL both blocks are displayed.

Value

A list containing the requested information.

Author(s)

Patricia Sebastian-Leon

See Also

`omicsCompAnalysis, caClass-class`
getMethodInfo

Examples

data("STATegRa_S3")
B1 <- createOmicsExpressionSet(Data=Block1.PCA, pData=ed.PCA, pDataDescr=c("classname"))
B2 <- createOmicsExpressionSet(Data=Block2.PCA, pData=ed.PCA, pDataDescr=c("classname"))
# Omics components analysis
res <- omicsCompAnalysis(Input=list(B1, B2), Names=c("expr", "mirna"), method="DISCOSCA", Rcommon=2, Rspecific=c(2, 2), center=TRUE, scale=TRUE, weight=TRUE)
getLoadings(res)
getLoadings(res, part="common", block="expr")
getLoadings(res, part="distinctive", block="expr")

getMethodInfo

Retrieve information about component analysis method

Description

Generic function to retrieve information about the method used by omicsCompAnalysis on a caClass-class object.

Usage

getMethodInfo(x, method=FALSE, comps=NULL, block=NULL)

Arguments

x caClass-class object.
method Logical indicating whether to return the method name.
comps Character indicating which component number to return ("common", "distinctive" or "all")
block Character indicating the block of data for which the component count will be given. It can be specified by the position of the block ("1" or "2") or the name assigned in the caClass-class object. If it is NULL both blocks are displayed.

Value

A list containing the requested information.

Author(s)

Patricia Sebastian-Leon

See Also

omicsCompAnalysis, caClass-class
getPreprocessing

Retrieve information about preprocessing

Description
Generic function to retrieve information about the preprocessing done by omicsCompAnalysis on a caClass-class object.

Usage
getPreprocessing(x, process=FALSE, preproData=FALSE, block=NULL)

Arguments
x caClass-class object.
process Logical indicating whether to return information about the processing done.
preproData Logical indicating whether to return the pre-processed data matrices.
block Character indicating the block of data to be returned. It can be specified by the position of the block ("1" or "2") or the name assigned in the caClass-class object. If it is NULL both blocks are displayed.

Value
If both process and preproData are specified, a list containing (otherwise the individual item):

process Character indicating the processing done
preproData Matrix (or list of matrices, depending on block) containing pre-processed data

Author(s)
Patricia Sebastian-Leon
getScores

See Also
omicsCompAnalysis, caClass-class

Examples

data("STATegRa_S3")
B1 <- createOmicsExpressionSet(Data=Block1.PCA, pData=ed.PCA,
pDataDescr=c("classname"))
B2 <- createOmicsExpressionSet(Data=Block2.PCA,
pData=ed.PCA, pDataDescr=c("classname"))
# Omics components analysis
res <- omicsCompAnalysis(Input=list(B1, B2), Names=c("expr", "mirna"),
                         method="DISCOSCA", Rcommon=2, Rspecific=c(2, 2),
                         center=TRUE, scale=TRUE, weight=TRUE)
getPreprocessing(res, process=TRUE)
getPreprocessing(res, preproData=TRUE, block="1")

getScores

Retrieve component analysis scores

Description
Generic function to retrieve scores (common and distinctive) found by omicsCompAnalysis on a caClass-class object.

Usage
getScores(x, part=NULL, block=NULL)

Arguments

x caClass-class object.

part Character indicating whether "common" or "distinctive" scores should be displayed

block Character indicating the block of data for which the scores will be given. It can be specified by the position of the block ("1" or "2") or the name assigned in the caClass-class object. If it is NULL both blocks are displayed.

Value
A list containing the requested information.

Author(s)
Patricia Sebastian-Leon
See Also

omicsCompAnalysis, caClass-class

Examples

data("STATegRa_S3")
B1 <- createOmicsExpressionSet(Data=Block1.PCA, pData=ed.PCA,
   pDataDescr=c("classname"))
B2 <- createOmicsExpressionSet(Data=Block2.PCA,
   pData=ed.PCA, pDataDescr=c("classname"))
# Omics components analysis
res <- omicsCompAnalysis(Input=list(B1, B2), Names=c("expr", "mirna"),
   method="DISCOSCA", Rcommon=2, Rspecific=c(2, 2),
   center=TRUE, scale=TRUE, weight=TRUE)
getScores(res)
getScores(res, part="common")
getScores(res, part="distinctive", block="expr")

getVAF

Retrieve information about VAF

Description

Generic function to retrieve VAF found by omicsCompAnalysis on a caClass-class object.

Usage

getVAF(x, part=NULL, block=NULL)

Arguments

x caClass-class object.
part Character indicating whether "common" or "distinctive" VAF should be displayed
block Character indicating the block of data for which the VAF will be given. It can be specified by the position of the block ("1" or "2") or the name assigned in the caClass-class object. If it is NULL both blocks are displayed.

Value

A list containing the requested information.

Author(s)

Patricia Sebastian-Leon

See Also

omicsCompAnalysis, caClass-class
Examples

data("STATegRa_S3")
B1 <- createOmicsExpressionSet(Data=Block1.PCA, pData=ed.PCA, pDataDescr="classname")
B2 <- createOmicsExpressionSet(Data=Block2.PCA, pData=ed.PCA, pDataDescr="classname")
# Omics components analysis
res <- omicsCompAnalysis(Input=list(B1, B2), Names=c("expr", "mirna"),
                           method="DISCOSCA", Rcommon=2, Rspecific=c(2, 2),
                           center=TRUE, scale=TRUE, weight=TRUE)
getVAF(res)
getVAF(res, part="common")
getVAF(res, part="distinctive", block="expr")

holistOmics

holistOmics an application of NPC on omics datasets

Description

This function is defunct. Use omicsNPC instead.

Usage

holistOmics(dataInput, dataTypes, comb.method = c("Fisher", "Liptak", "Tippett"),
            numPerm = 1000, numCores = 1, verbose = FALSE)

Arguments

dataInput List of ExpressionSet objects, one for each data modality.
dataTypes Character vector with possible values: 'RNA-seq', 'microarray'
comb.method Character vector with possible values: 'Fisher', 'Liptak', 'Tippett', if more than
              one is specified, all will be used.
numPerm Number of permutations
numCores Number of CPU cores to use
verbose Logical, if set to TRUE holistOmics prints out the step that it performs

Value

A data.frame

Author(s)

Nestoras Karathanasis

References

Pesarin, Fortunato, and Luigi Salmaso. Permutation tests for complex data: theory, applications
Examples

```r
# Load the data
data("TCGA_BRCA_Batch_93")
# Setting dataTypes, the first two ExpressionSets include RNAseq data,
# the third ExpressionSet includes Microarray data.
dataTypes <- c("RNAseq", "RNAseq", "Microarray")
# Setting methods to combine pvalues
comb.method = c("Fisher", "Liptak", "Tippett")
# Setting number of permutations
numPerm = 1000
# Setting number of cores
numCores = 1
# Setting holistOmics to print out the steps that it performs.
verbose = TRUE
# Run holistOmics analysis.
# The output is a data.frame of p-values.
# Each row corresponds to a gene name. Each column corresponds to a method
# used in the analysis.
## Not run: out <- holistOmics(dataInput = TCGA_BRCA_Data, dataTypes = dataTypes,
##                            comb.method = comb.method, numPerm = numPerm,
##                            numCores = numCores, verbose = verbose)
## End(Not run)
```

---

**modelSelection**

*Find optimal common and distinctive components*

**Description**

Estimate the optimal number of common and distinctive components according to given selection criteria.

**Usage**

```r
modelSelection(Input, Rmax, fac.sel, varthreshold=NULL, nvar=NULL, PCnum=NULL, center=FALSE, scale=FALSE, weight=FALSE, plot_common=FALSE, plot_dist=FALSE)
```

**Arguments**

- **Input** List of ExpressionSet objects, one for each block of data
- **Rmax** Maximum common components
- **fac.sel** PCA criteria for selection ("%accum", "single%", "rel.abs", "fixed.num")
- **varthreshold** Cumulative variance criteria for PCA selection. Threshold for "%accum" or "single%" criteria.
- **nvar** Relative variance criteria. Threshold for "rel.abs".
- **PCnum** Fixed number of components for "fixed.num".
- **center** Character (or FALSE) specifying which (if any) centering will be applied before analysis. Choices are "PERBLOCKS" (each block separately) or "ALL-BLOCKS" (all data together).
modelSelection

scale Character (or FALSE) specifying which (if any) scaling will be applied before analysis. Choices are "PERBLOCKS" (each block separately) or "ALL-BLOCKS" (all data together).

weight Logical indicating whether weighting is to be done. Choices are "BETWEEN-BLOCKS"

plot_common Logical indicating whether to plot the explained variances (SSQ) of each block and its estimation and the ratios

plot_dist Logical indicating whether to plot the explained variances (SSQ) and the accumulated variance for each block

Value

List containing:

common List with common components results

commonComps Optimal number of common components

ssqs Matrix of SSQ for each block and estimator

pssq ggplot object showing SSQ for each block and estimator

pratios ggplot object showing SSQ ratios between each block and estimator

dist List containing the results of distinct PCA for each input block; for each block PCAres and numComps is returned within a list

PCAres List containing results of PCA, with fields "eigen", "var.exp", "scores" and "loadings"

nomComps Number of components selected

Author(s)

Patricia Sebastian-Leon

See Also

omicsCompAnalysis

Examples

data(STATegRa_S3)
B1 <- createOmicsExpressionSet(Data=Block1.PCA,pData=ed.PCA,pDataDescr=c("classname"))
B2 <- createOmicsExpressionSet(Data=Block2.PCA,pData=ed.PCA,pDataDescr=c("classname"))
ms <- modelSelection(Input=list(B1, B2), Rmax=3, fac.sel="single\%", varthreshold=0.03, center=TRUE, scale=FALSE, scale=FALSE, weight=FALSE, plot_common=FALSE, plot_dist=FALSE)
ms
Components analysis for multiple objects

Description

This function performs a components analysis of object wise omics data to understand the mechanisms that underlay all the data blocks under study (common mechanisms) and the mechanisms underlying each of the data block independently (distinctive mechanisms). This analysis include both, the preprocessing of data and the components analysis by using three different methodologies.

Usage

```r
omicsCompAnalysis(Input, Names, method, Rcommon, Rspecific,
                   convThres=1e-10, maxIter=600, center=FALSE,
                   scale=FALSE, weight=FALSE)
```

Arguments

- **Input**: List of `ExpressionSet` objects, one for each block of data.
- **Names**: Character vector giving names for each `Input` object.
- **method**: Method to use for analysis (either "DISCOSCA", "JIVE", or "O2PLS").
- **Rcommon**: Number of common components between all blocks.
- **Rspecific**: Vector giving number of unique components for each input block.
- **convThres**: Stop criteria for convergence.
- **maxIter**: Maximum number of iterations.
- **center**: Character (or FALSE) specifying which (if any) centering will be applied before analysis. Choices are "PERBLOCKS" (each block separately) or "ALL-BLOCKS" (all data together).
- **scale**: Character (or FALSE) specifying which (if any) scaling will be applied before analysis. Choices are "PERBLOCKS" (each block separately) or "ALL-BLOCKS" (all data together).
- **weight**: Logical indicating whether weighting is to be done.

Value

An object of class `caClass-class`.

Author(s)

Patricia Sebastian Leon
Examples

```r
data("STATegRa_S3")
B1 <- createOmicsExpressionSet(Data=Block1.PCA,pData=ed.PCA,
pDataDescr=c("classname"))
B2 <- createOmicsExpressionSet(Data=Block2.PCA,
pData=ed.PCA,pDataDescr=c("classname"))
# Omics components analysis
discoRes <- omicsCompAnalysis(Input=list(B1,B2),Names=c("expr","mirna"),
method="DISCOSCA",Rcommon=2,Rspecific=c(2,2),
center=TRUE,scale=TRUE,weight=TRUE)
jiveRes <- omicsCompAnalysis(Input=list(B1,B2),Names=c("expr","mirna"),
method="JIVE",Rcommon=2,Rspecific=c(2,2),
center=TRUE,scale=TRUE,weight=TRUE)
o2plsRes <- omicsCompAnalysis(Input=list(B1,B2),Names=c("expr","mirna"),
method="O2PLS",Rcommon=2,Rspecific=c(2,2),
center=TRUE,scale=TRUE,weight=TRUE)
```

Description

This function applies the Non-Parametric Combination methodology on the integrative analysis of different omics data modalities. It retrieves genes associated to a given outcome, taking into account all omics data. First, each datatype is analyzed independently using the appropriate method. omicsNPC analyses continuous data (microarray) using limma, while count data (e.g., RNAseq) are first preprocessed with the "voom" function. The user can also specify their own function for computing deregulation/association. The p-values from the single dataset analysis are then combined employing Fisher, Liptak and Tippett combining functions. The Tippett function returns findings which are supported by at least one omics modality. The Liptak function returns findings which are supported by most modalities. The Fisher function has an intermediate behavior between those of Tippett and Liptak.

Usage

```r
omicsNPC(dataInput, dataMapping, dataTypes = rep('continuous', length(dataInput)),
combMethods = c("Fisher", "Liptak", "Tippett"), numPerms = 1000,
numCores = 1, verbose = FALSE, functionGeneratingIndex = NULL,
outcomeName = NULL, allCombinations = FALSE, dataWeights = rep(1, length(dataInput))/length(dataInput),
returnPermPvalues = FALSE, ...)
```

Arguments

dataInput
  List of ExpressionSet objects, one for each data modality.
dataMapping
  A data frame describing how to map measurements across datasets. See details for more information.
dataTypes Character vector with possible values: 'continuous', 'count'. Alternatively, a list of functions for assessing deregulation / association with an outcome

combMethods Character vector with possible values: 'Fisher', 'Liptak', 'Tippett'. If more than one is specified, all will be used.

numPerms Number of permutations

numCores Number of CPU cores to use

verbose Logical, if set to TRUE omicsNPC prints out the step that it performs

functionGeneratingIndex Function generating the indices for randomly permuting the samples

outcomeName Name of the outcome of interest / experimental factor, as reported in the design matrices. If NULL, the last column of the design matrices is assumed to be the outcome of interest.

allCombinations Logical, if TRUE all combinations of omics datasets are considered

dataWeights A vector specifying the weight to give to each dataset. Note that sum(dataWeights) should be 1.

returnPermPvalues Logical, should the p-values computed at each permutation being returned?

... Additional arguments to be passed to the user-defined functions

Value

A list containing: stats0 Partial deregulation / association statistics pvalues0 The partial p-values computed on each dataset pvaluesNPC The p-values computed through NPC. permPvalues The p-values computed at each permutation

Author(s)

Nestoras Karathanasis, Vincenzo Lagani

References


Examples

# Load the data
data("TCGA_BRCA_Batch_93")
# Setting dataTypes, the first two ExpressionSets include RNAseq data,
# the third ExpressionSet includes Microarray data.
dataTypes <- c("count", "count", "continuous")
# Setting methods to combine pvalues
combMethods = c("Fisher", "Liptak", "Tippett")
# Setting number of permutations
numPerms = 1000
# Setting number of cores
umCores = 1
# Setting omicsNPC to print out the steps that it performs.
verbose = TRUE
# Run omicsNPC analysis.
# The output contains a data.frame of p-values, where each row corresponds to a gene,
# and each column corresponds to a method used in the analysis.

## Not run: out <- omicsNPC(dataInput = TCGA_BRCA_Data, dataTypes = dataTypes,
combMethods = combMethods, numPerms = numPerms,
numCores = numCores, verbose = verbose)
## End(Not run)

---

**plotRes**

**Plot component analysis results**

**Description**

Plot scatterplots of scores or loadings, for common and distinctive parts as well as combined plots.

**Usage**

```r
plotRes(object, comps = c(1, 2), what = "both", type = "individual", combined = FALSE,
        block = NULL, color = NULL, shape = NULL, labels = NULL,
        title = NULL, xlabel = NULL, ylabel = NULL, background = TRUE,
        palette = NULL, pointSize = 4, labelSize = NULL,
        axisSize = NULL, titleSize = NULL, sizeValues = c(2, 4),
        shapeValues = c(17, 0))
```

**Arguments**

- **object**: *caClass-class* containing component analysis results
- **comps**: If combined = FALSE, it indicates the x and y components of the type and block chosen. If combined = TRUE, it indicates the component to plot for the first block of information and the component for the second block of information to plot together. By default the components are set to c(1, 2) if combined = FALSE and to c(1, 1) if combined = TRUE.
- **what**: Either "scores", "loadings" or "both"
- **type**: Either "common", "individual" or "both"
- **combined**: Logical indicating whether to make a simple plot of two components from one block, or components from different blocks
- **block**: Which block to plot, either "1" or "2" or the name of the block.
- **color**: Character specifying a pData column from the original data to use to color points
- **shape**: Character specifying a pData column to select point shape
- **labels**: Character specifying a pData column from which to take point labels
- **title**: Main title
- **xlabel**: x-axis name
**ylabel**

y-axis name

**background**

Logical specifying whether to make a grey background

**palette**

Vector giving the color palette for the plot

**pointSize**

Size of plot points

**labelSize**

Size of point labels if not NULL

**axisSize**

Size of axis text

**titleSize**

Size of title text

**sizeValues**

Vector containing sizes for scores and loadings

**shapeValues**

Vector indicating the shapes for scores and loadings

**Value**

`ggplot` object

**Author(s)**

Patricia Sebastian-Leon

**Examples**

data("STATegRa_S3")

B1 <- createOmicsExpressionSet(Data=Block1.PCA,pData=ed.PCA,
     pDataDescr=c("classname"))

B2 <- createOmicsExpressionSet(Data=Block2.PCA,
     pData=ed.PCA,pDataDescr=c("classname"))

# Omics components analysis
discoRes <- omicsCompAnalysis(Input=list(B1,B2),Names=c("expr","mirna"),
     method="DISCOSCA",Rcommon=2,Rspecific=c(2,2),
     center=TRUE,scale=TRUE,weight=TRUE)

jiveRes <- omicsCompAnalysis(Input=list(B1,B2),Names=c("expr","mirna"),
     method="JIVE",Rcommon=2,Rspecific=c(2,2),
     center=TRUE,scale=TRUE,weight=TRUE)

o2plsRes <- omicsCompAnalysis(Input=list(B1,B2),Names=c("expr","mirna"),
     method="O2PLS",Rcommon=2,Rspecific=c(2,2),
     center=TRUE,scale=TRUE,weight=TRUE)

# Scatterplot of scores variables associated to common components

# DISCO-SCA
plotRes(object=discoRes,comps=c(1,2),what="scores",type="common",
     combined=FALSE,block=NULL,color="classname",shape=NULL,labels=NULL,
     background=TRUE,palette=NULL,pointSize=4,labelSize=NULL,
     axisSize=NULL,titleSize=NULL)

# JIVE
plotRes(object=jiveRes,comps=c(1,2),what="scores",type="common",
     combined=FALSE,block=NULL,color="classname",shape=NULL,labels=NULL,
     background=TRUE,palette=NULL,pointSize=4,labelSize=NULL,
     axisSize=NULL,titleSize=NULL)
# O2PLS
# Scatterplot of scores variables associated to common components
# Associated to first block
p1 <- plotRes(object=o2plsRes, comps=c(1,2), what="scores", type="common",
              combined=FALSE, block="expr", color="classname", shape=NULL,
              labels=NULL, background=TRUE, palette=NULL, pointSize=4,
              labelSize=NULL, axisSize=NULL, titleSize=NULL)
# Associated to second block
p2 <- plotRes(object=o2plsRes, comps=c(1,2), what="scores", type="common",
              combined=FALSE, block="mirna", color="classname", shape=NULL,
              labels=NULL, background=TRUE, palette=NULL, pointSize=4,
              labelSize=NULL, axisSize=NULL, titleSize=NULL)

# Combined plot of scores variables associated to common components
plotRes(object=o2plsRes, comps=c(1,1), what="scores", type="common",
        combined=TRUE, block=NULL, color="classname", shape=NULL,
        labels=NULL, background=TRUE, palette=NULL, pointSize=4,
        labelSize=NULL, axisSize=NULL, titleSize=NULL)

# Loadings plot for individual components
# Separately for each block
p1 <- plotRes(object=discoRes, comps=c(1,2), what="loadings", type="individual",
              combined=FALSE, block="expr", color="classname", shape=NULL,
              labels=NULL, background=TRUE, palette=NULL, pointSize=4,
              labelSize=NULL, axisSize=NULL, titleSize=NULL)
p2 <- plotRes(object=discoRes, comps=c(1,2), what="loadings", type="individual",
              combined=FALSE, block="mirna", color="classname", shape=NULL,
              labels=NULL, background=TRUE, palette=NULL, pointSize=4,
              labelSize=NULL, axisSize=NULL, titleSize=NULL)

# Biplot: scores + loadings
plotRes(object=discoRes, comps=c(1,2), what="both", type="common",
        combined=FALSE, block="expr", color="classname", shape=NULL,
        labels=NULL, background=TRUE, palette=NULL, pointSize=4,
        labelSize=NULL, axisSize=NULL, titleSize=NULL)

---

plotVAF  

Plot VAF (Variance Explained For) from Component Analysis

Description

This function visualises the VAF results from component analysis. The input is a caClass-class
object from omicsCompAnalysis. VAF cannot be calculated if mode "O2PLS" was used. The plots
for modes "DISCOSCA" and "JIVE" are different since DISCO-SCA distinctive components have
some VAF in the other block. This VAF can be interpreted as an error in the rotation.

Usage

plotVAF(object, mainTitle="")
Arguments

- **object**: `caClass-class` object containing component analysis results
- **mainTitle**: Plot title

Value

- ggplot object

Author(s)

Patricia Sebastian-Leon

Examples

```r
data("STATegRa_S3")
require(ggplot2)
B1 <- createOmicsExpressionSet(Data=Block1.PCA,pData=ed.PCA,
pDataDescr=c("classname"))
B2 <- createOmicsExpressionSet(Data=Block2.PCA,
pData=ed.PCA,pDataDescr=c("classname"))
# Omics components analysis
discoRes <- omicsCompAnalysis(Input=list(B1,B2),Names=c("expr","mirna"),
    method="DISCOSCA",Rcommon=2,Rspecific=c(2,2),
    center=TRUE,scale=TRUE,weight=TRUE)
jiveRes <- omicsCompAnalysis(Input=list(B1,B2),Names=c("expr","mirna"),
    method="JIVE",Rcommon=2,Rspecific=c(2,2),
    center=TRUE,scale=TRUE,weight=TRUE)

# DISCO-SCA plotVAF
plotVAF(discoRes)

# JIVE plotVAF
plotVAF(jiveRes)
```

Description

STATegRa is a package for the integrative analysis of multi-omic data-sets.

For full information, see the user’s guide.

See Also

- `STATegRaUsersGuide`
Defunct functions in STATegRa

Description
These functions are defunct and no longer available.

Details
- holistOmics: replaced by omicsNPC

STATegRaUsersGuide

Description
Finds the location of the STATegRa User's Guide and optionally opens it.

Usage
STATegRaUsersGuide(view = TRUE)

Arguments
  view  Whether to open a browser

Value
The path to the documentation

Author(s)
David Gomez-Cabrero

Examples
STATegRaUsersGuide(view=FALSE)
Data were downloaded from TCGA data portal, https://tcga-data.nci.nih.gov/tcga/. We downloaded sixteen tumour samples and the sixteen matching normal, for Breast invasive carcinoma, BRCA, batch 93. Herein, three types of data modalities are included, RNAseq (TCGA_BRCA_Data$RNAseq), RNAseqV2 (TCGA_BRCA_Data$RNAseqV2) and Expression-Genes (TCGA_BRCA_Data$Microarray). The Data Level was set to Level 3. For each data type, we pooled all data to one matrix, where rows corresponded to genes and columns to samples. Only the first 100 genes are included.
Format

One list, which contains three ExpressionSet objects.

Author(s)

Nestoras Karathanasis, Vincenzo Lagani

Source

See https://tcga-data.nci.nih.gov/tcga/.

Examples

# load data
data(TCGA_BRCA_Batch_93)