## Package 'SPADEVizR'

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```
Type Package
```

**Title** Visualization and statistical analyses of cell clustering results generated by the SPADE clustering algorithm

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#### **Description**

Flow and mass cytometry are experimental techniques used for the characterization of cell phenotypes. Automatic gating algorithms can be used to automatically identified clusters of cells having similar phenotypes. The SPADE algorithm has been proposed as a new way to analysis and explore mass-cytometry data. This algorithm performs a density-based down-sampling combined with an agglomerative hierarchical clustering. XXX While SPADE offers new opportunities for identifying cell populations, complementary approaches are needed to improve the characterization of identified cell populations. SPADEVizR is an R package designed to better visualize and analyze SPADE clustering results. This package extends the original SPADE outputs with techniques such as parallel coordinates, heatmaps, multidimensional scaling, volcano plots or streamgraph representations. Moreover several statistical methods allow the identification of SPADE clusters with relevant biological behaviors.

#### License GPL-3

```
Imports data.table,
      flowCore,
      ggdendro,
      ggnetwork,
      ggplot2,
      ggrepel,
      gridExtra,
      gtable,
      gtools,
      igraph,
      MASS,
      network,
      plyr,
      reshape2
LazyData true
Suggests knitr
VignetteBuilder knitr
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```

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abundantClustersViewer

Visualization of abundant clusters

#### **Description**

Generates a scatter plot representation showing for each cluster its mean abundance and associated p-value.

## Usage

```
abundantClustersViewer(AC, show.cluster_sizes = TRUE,
    show.all_labels = FALSE)
```

#### **Arguments**

#### **Details**

By default, only significant abundant clusters are labeled. Labels for all clusters can be displayed by setting the 'show.all\_labels' parameter to TRUE.

#### Value

a 'ggplot' object

AC-class

Abundant Clusters (AC class) definition

#### **Description**

The 'AC' object is a S4 object containing the information related to the abundant clusters in a given biological condition. Moreover this object contains all parameters used in the statistical analysis.

#### **Details**

A cluster is considered as a significant abundant cluster if its associated p-value and mean are below the specific thresholds 'th.pvalue' and 'th.mean'.

The 'print()' and 'show()' can be used to display a summury of this object. Moreover all information about this object could be saved as a tab separated file using the 'export()' method. This object is returned by the 'identifyAC()' function.

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#### **Slots**

sample.names a character vector containing the samples used to compute the abundant clusters cluster.size a numeric vector containing the number of cells ( - sum of all samples - ) for each

use.percentages a logical specifying if computation was performed on percentage of cell abundance

method a character containing the name of the statistical test used to identify the abundant clusters method.adjust a character containing the name of the multiple correction method used (if any) th.mean a numeric value specifying the mean threshold

th.pvalue a numeric value specifying the p-value threshold

result a data frame containing for each cluster (first column): the mean (second column) and the standard deviation (third column) of the biological condition, the associated p-value (fourth column) and a logical (fifth column) specifying if the cluster is significantly abundant.

biplotViewer biplotViewer

#### **Description**

Generates a biplot representation with two markers

#### Usage

```
biplotViewer(SPADEResults, x.marker, y.marker, samples = NULL,
 clusters = NULL, sample.merge = FALSE, resample.ratio = NULL)
```

#### Arguments

**SPADEResults** a SPADEResults object (Results object is not accepted) a character indicating the marker name of the first dimension x.marker a character indicating the marker name of the second dimension y.marker samples named vector providing the correspondence between samples name (in rowname) and the logical value TRUE to use these samples (all samples by default) clusters a character vector containing the clusters names to be visualized (by default all

clusters will be used)

sample.merge

resample.ratio a numeric ratio (between 0 and 1) specifying the downsample ratio to show less

dots (or NULL)

#### **Details**

In such representation, each dot corresponds to a cell profile and dots are ploted in a 2-dimentional space corresponding to the selected markers. When too cells dots are displayed, it can require some seconds. In order to seep up the computation, it is possible to reduce the number of cells displayed (downsampling) using the 'resample.ratio' parameter.

#### Value

if return.gg is TRUE, the function returns a list of ggplot objects

boxplotViewer 5

boxplotViewer	Visualization of cluster enrichment profiles conditions	

## **Description**

Generate a boxplot representation displaying the cell abundance for each cluster. Clusters are gathered by given biological conditions.

#### Usage

```
boxplotViewer(Results, conditions, clusters = NULL, use.percentages = TRUE,
    show.legend = FALSE, show.violin = TRUE)
```

#### **Arguments**

Results a SPADEResults or Results object

conditions conditions a named vector providing the correspondence between a sample name

(in row names) and the condition of this sample or NA to exclude

clusters a character vector containing the clusters names to be visualized (by default all

clusters will be displayed)

use.percentages

a logical specifying if the visualization must be performed on percentage

show.legend a logical specifying if the legend must be displayed

show.violin a logical specifying if the count distribution must be displayed

#### Details

Cells clusters are colored based on theirs associated biological samples.

#### Value

```
a 'ggplot' object
```

CC-class	Correlated Clusters (CC class) definition
	, , , ,

#### **Description**

The 'CC' object is a S4 object containing coefficient of correlation associated between each cluster and a phenotypic variable. Moreover this object contains all parameters used in the statistical analysis.

#### **Details**

A cluster is considered as a significant correlated cluster if its associated p-value and correlation threshold are below the specific thresholds 'th.pvalue' and 'th.correlation'.

The 'print()' and 'show()' can be used to display a summury of this object. Moreover all information about this object could be saved as a tab separated file using the 'export()' method. This object is returned by the 'identifyCC()' function.

#### **Slots**

sample.names a character vector containing the samples used to compute correlated clusters variable a numeric vector containing the expression values of the associated variable

cluster.size a numeric vector containing number of cells ( – sum of all samples – ) for each cluster

use.percentages a logical specifying if computation was performed on percentage of cell abundance

method a character containing the name of the statistical test used to identify the CC

method adjust a character containing the name of the multiple correction method used (if any)

th.correlation a numeric value specifying the correlation threshold (R)

th.pvalue a numeric value specifying the p-value threshold

result a data.frame containing for each cluster (first column): the coefficiant of correlation R (second column), the associated p-value (third column) and a logical (fourth column) specifying if the cluster is significantly correlated.

classifyEnrichmentProfiles

Classification of clusters based on their enrichment profiles

#### **Description**

Classifies clusters based on their enrichment profiles.

#### Usage

```
classifyEnrichmentProfiles(Results, method = "hierarchical_h",
  method.parameter = NULL)
```

#### **Arguments**

Results a Results or SPADEResults object

method a character specifying the clustering method among one of those: 'hierarchical\_h','hierarchical\_k','k-

means', 'eigencell', 'clique' (hierarchical\_h by default)

method.parameter

a numeric specifying the numeric value required by the selected method

#### **Details**

The classification is done on cell abundances of each clusters and could be performed using 5 methods:

• "hierarchical\_k" This method first compute the pearson correlation matrix and then use this matrix to performs a hierarchical classification. The hierarchical classification is cutted in order to return the desired number of classes. This number of classes must be provided as a numeric integer using the 'method.parameter' parameter. It is to note that negative correlations are considered as uncorrelated

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• "hierarchical\_h" (default method) This method works in the same way than 'hierarchical\_k' but the height where the hierarchical tree is specified. This height is a correlation threshold (a numeric double between 0 and 1 included, default is 0.7) provided using the 'method.parameter' parameter.

- "eigencell" This method compute the performs a eigen vector decomposition and then calculate the correlations between cluster expressions and these vectors. Clusters which correlate above a specific threshold with the same eigen vector are classified together. This correlation threshold (a numeric double between 0 and 1 included, default is 0.8) provided using the 'method.parameter' parameter.
- "clique" This method first compute the Pearson correlation matrix and then use this matrix to generate an undirected graph. In this graph, an edge is drawn between two nodes if the correlation coefficient in the adjacency matrix is above a specific threshold. This correlation threshold (a numeric double between 0 and 1 included, default is 0.7) provided using the 'method parameter' parameter. After building the graph, the method looking for the largest cliques wich are considered as classes of nodes. Cliques correspond to subgraph in which every two distinct vertices are adjacent.

#### Value

a S4 object of class 'EnrichmentProfiles'

classifyPhenoProfiles Classification of clusters based on their phenotype profiles

## **Description**

Classifies clusters based on their phenotype profiles.

#### Usage

```
classifyPhenoProfiles(Results, method = "hierarchical_h",
  method.parameter = NULL)
```

## **Arguments**

Results or SPADEResults object

method a character specifying the clustering method among one of those: "hierarchi-

cal\_h", "hierarchical\_k", "k-means", "eigencell", "clique"

method.parameter

a numeric specifying the numeric value required by the selected method

## **Details**

The classification is done on cell abundances of each clusters and could be performed using 5 methods:

• "hierarchical\_k" This method first compute the pearson correlation matrix and then use this matrix to performs a hierarchical classification. The hierarchical classification is cutted in order to return the desired number of classes. This number of classes must be provided as a numeric integer using the 'method.parameter' parameter. It is to note that negative correlations are considered as uncorrelated

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• "hierarchical\_h" (default method) This method works in the same way than 'hierarchical\_k' but the height where the hierarchical tree is specified. This height is a correlation threshold (a numeric double between 0 and 1 included, default is 0.7) provided using the 'method.parameter' parameter.

- "eigencell" This method compute the performs a eigen vector decomposition and then calculate the correlations between cluster expressions and these vectors. Clusters which correlate above a specific threshold with the same eigen vector are classified together. This correlation threshold (a numeric double between 0 and 1 included, default is 0.8) provided using the 'method.parameter' parameter.
- "clique" This method first compute the Pearson correlation matrix and then use this matrix to generate an undirected graph. In this graph, an edge is drawn between two nodes if the correlation coefficient in the adjacency matrix is above a specific threshold. This correlation threshold (a numeric double between 0 and 1 included, default is 0.7) provided using the 'method parameter' parameter. After building the graph, the method looking for the largest cliques wich are considered as classes of nodes. Cliques correspond to subgraph in which every two distinct vertices are adjacent.

#### Value

a S4 object of class 'PhenoProfiles'

|--|

## **Description**

Generates a parallel coordinate plot representation showing for each cluster the marker median expressions.

## Usage

```
clusterViewer(Results, clusters = NULL, samples = NULL, markers = NULL,
    show.mean = "both")
```

## Arguments

Results	a SPADEResults or Result object
clusters	a character vector containing the clusters names to be visualized (by default all clusters will be displayed)
samples	named vector providing the correspondence between samples name (in rowname) and the logical value TRUE to use these samples (all samples by default)
markers	a character vector specifying the markers to be displayed
show.mean	a character specifying if marker means expression should be displayed, possible value are among: "none", "only" or "both"

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#### **Details**

If 'Results' is an object of class 'SPADEResults' object then the quantile ranges will be displayed using a ribbon.

The 'samples' parameter required a named vector providing the correspondence between a sample name (in rowname) and the logical value TRUE to show this sample or FALSE otherwise.

The 'show.mean' parameter allows to visualize three kinds of information:

- "none" value will show marker median expressions for each selected samples;
- "only" value will show only the mean of median maker expressions for all selected samples (displayed as black dashed line);
- "both" value will show marker median expressions for each selected samples together with the mean of median maker expressions for all selected samples.

#### Value

a list of ggplot objects

computeClique

Internal - Clique percolation classification

## Description

This function is used internally to classify clusters enrichment profiles or phenotype profiles using a clique percolation algorithm.

## Usage

```
computeClique(data, clique.correlation.th = 0.7)
```

## **Arguments**

```
data a numeric matrix with all clusters in rownames clique.correlation.th a numeric value indicating the correlation coefficient threshold
```

#### **Details**

xxx

## Value

a dataframe containing for each cluster, its name and class

compute Eigen Cell Clusters

Internal - Eigen vector classification

## Description

This function is used internally to classify clusters enrichment profiles or phenotype profiles using eigen vector decomposition.

## Usage

```
computeEigenCellClusters(data, eigencell.correlation.th = 0.8)
```

## **Arguments**

```
data a numeric matrix with all clusters in rownames eigencell.correlation.th a numeric value indicating the correlation coefficient threshold
```

#### **Details**

XXX

#### Value

a dataframe containing for each cluster, its name and class

```
computeHierarchicalClustering
```

Internal - Hierarchical classification

## Description

This function is used internally to classify clusters enrichment profiles or phenotype profiles using a hierarchical algorithm.

## Usage

```
computeHierarchicalClustering(data, class.number = NULL,
    hierarchical.correlation.th = 0.8)
```

## **Arguments**

```
data a matrix with all clusters in rownames
class.number a numeric specifying the number of classes
hierarchical.correlation.th
a numeric value specifying the cut height
```

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#### **Details**

This function compute the Pearson correlation matrix associated to the provided matrix. It is to note that negative correlations are considered as uncorrelated. This correlation matrix is used to performs a hierarchical classification. If 'class.number' is NULL, classification will be determined base on the cut height correlation threshold.

## Value

a dataframe containing for each cluster, its name and class

computeKmeans

Internal - Kmeans classification

## **Description**

This function is used internally to classify clusters enrichment profilies or phenotype profiles using a k-means algorithm.

## Usage

```
computeKmeans(data, k = NULL)
```

## **Arguments**

data a numeric matrix with cluster names in rownames k a numeric specifying the desired number of classes

#### **Details**

XXX

## Value

a dataframe containing for each cluster, its name and class

computePhenoTable

Internal - Generatate marker expression scores describing phenotypes

## Description

This function is used internally to generate a melted numeric matrix of discrete expression scores for each marker of each cluster.

## Usage

```
computePhenoTable(SPADEResults, num = 5)
```

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## **Arguments**

SPADEResults a SPADEResults object

num a numeric value specifying the number of markers expression categories

## **Details**

NA values are removed

#### Value

a numeric matrix of expression scores

 ${\tt compute} \\ {\tt Quantile}$ 

Internal - Compute quantile with FCS flowset marker by marker

## Description

This function is used internally to compute the maker range quantiles.

## Usage

```
computeQuantile(flowset, probs = c(0.05, 0.95))
```

## Arguments

flowset a flowCore flowset

probs a numeric vector of 2 values specifying the quantiles to compute

## **Details**

This function performs the exact calculation of quantiles with all cells but needs more memory than 'computeQuantile.approximation'.

## Value

a numeric matrix containing the quantiles of each marker

 ${\tt computeQuantile.approximation}$ 

Internal - Compute quantile with FCS flowset sample by sample

## Description

This function is used internally to provide the mean of quantiles from each sample to seed up computation.

## Usage

```
computeQuantile.approximation(flowset, probs = c(0.05, 0.95))
```

## **Arguments**

flowset a flowCore flowset

probs a numeric vector of 2 values specifying the quantiles to compute

#### **Details**

This function performs an approximate calculation of quantiles using less memory than compute-Quantile.

#### Value

a numeric matrix containing the quantiles of each marker

correlatedClustersViewer

Visualization of correlated clusters

## **Description**

Generate a scatter plot representation showing for each cluster

## Usage

```
correlatedClustersViewer(CC, show.cluster.sizes = TRUE,
    show.all_labels = FALSE)
```

## **Arguments**

```
CC an object of class 'CC' (object returned by the 'computeCC()' function)
show.cluster.sizes
a logical specifying if dot sizes are proportional to cell counts
show.all_labels
a logical specifying if all cluster label must be show or just significant cluster
```

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## **Details**

By default, only significant correlated clusters are labeled. Labels for all clusters can be displayed by setting the 'all.label' parameter to TRUE.

## Value

```
a 'ggplot' object
```

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## Description

Generate a two dimensional vizualisation showing the number of cells (sum of selected samples) of each cluster.

## Usage

```
countViewer(Results, samples = NULL, clusters = NULL, min.cells = 0,
    sort = TRUE, show.samples = TRUE)
```

## Arguments

Results	a SPADEResults or Results object
samples	a named vector providing the correspondence between samples name (in rownames) and the logical value TRUE to use these samples (all samples by default)
clusters	a character vector containing the clusters names to be visualized (by default all clusters will be displayed)
min.cells	a numeric specifying the minimum number of cell (sum of all selected samples) to display a cluster
sort	a logical specifying if clusters will be to be sorted (descending) based on the sum of all selected samples for each cluster
show.samples	a logical specifying if the number of cells for all selected samples will be displayed

#### **Details**

XXX

## Value

```
a 'ggplot' object
```

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DEC-class

Differentially Enriched Clusters (DEC class) definition

#### **Description**

The 'DEC' object is a S4 object containing the information related to the differentially enriched clusters between two given biological conditions. Moreover this object contains all parameters used in the statistical analysis.

#### **Details**

A cluster is considered as a differentially enriched cluster if its associated p-value and fold-change are below the specific thresholds 'th.pvalue' and 'th.fc'.

The 'print()' and 'show()' can be used to display a summury of this object. Moreover all information about this object could be saved as a tab separated file using the 'export()' method. This object is returned by the 'identifyDEC()' function.

#### **Slots**

sample.cond1 a character specifying the names of the samples of the first biological condition
sample.cond2 a character specifying the names of the samples of the second biological condition
cluster.size a numeric vector containing number of cells ( - sum of all samples - ) for each
cluster

use.percentages a logical specifying if computation was performed on percentage of cell abundance

method a character containing the name of the statistical test used to identify the DEC

method.adjust a character containing the name of the multiple correction method used (if any)

method.paired a logical indicating if the statistical test have been performed in a paired manner

th.fc a numeric value specifying the fold-change threshold

th.pvalue a numeric value specifying the p-value threshold

result a data.frame containing for each cluster (first column): the fold-change (second column) and the standard deviation (third column) for the first biological condition, the fold-change (fourth column) and the standard deviation (fifth column) for the second biological condition, the associated p-value (sixth column) and a logical (seventh column) specifying if the cluster is significantly differentially enriched.

distogramViewer

Visualization of marker co-expressions

#### Description

Generate a distogram representation showing the marker co-expressions.

#### Usage

```
distogramViewer(Results, clusters = NULL, samples = NULL, markers = NULL)
```

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#### **Arguments**

Results a SPADEResults or Results object

clusters a character vector containing the clusters names to be use (by default all clusters

will be used)

samples a character vector specifying the samples to use

markers a character vector specifying the markers to be displayed

#### **Details**

A pearson correlation matrix is calculated between selected markers using selected clusters and samples. High positive correlated markers are shown by a green tile at their perpendicular intersection. In the same way, absence of correlation are shown by black tiles and negative correlation by red tiles.

#### Value

a list of 'ggplot' objects

EnrichmentProfiles-class

Classification of clusters based on their enrichment profiles (EnrichmentProfiles class) definition

#### **Description**

The 'EnrichmentProfiles' is a S4 object containing the information related to the cluster classification based on theirs enrichment profiles.

This object contains all information about the classification method and parameters used.

#### **Details**

Five methods are available to classify cellular clusters: 'hierarchical\_k', 'hierarchical\_h', 'kmeans', 'eigencell' and 'clique'. Each method can parameterized using the 'method.parameter' parameter.

The 'print()' and 'show()' can be used to display a summary of this object. Moreover all information about this object could be saved as a tab separated file using the 'export()' method. This object is returned by the 'classifyEnrichmentProfiles()' function.

#### **Slots**

method a character specifying the method used to classify cluster

method.parameter a numeric parameters associated with the chosen method

cluster.size a numeric vector with the number of cell in each cluster (sum of all samples)

 ${\tt cluster.number}\ \ a\ numeric\ providing\ the\ number\ of\ cluster$ 

class.number a numeric providing the number of classes

classes a two column dataframe with the cluster in first column and corresponding class in the second column

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exclude.markers

Internal - Removing of cell markers to exclude from a matrix

#### **Description**

This function is used internally to remove one or several cell markers.

#### Usage

```
exclude.markers(data, exclude, colnames.FCS = NULL)
```

## **Arguments**

data a numeric matrix or flowset

exclude a character vector containing the cell markers to be excluded (case intensive)

colnames.FCS a character vector containing column names if data is a FCS flowset

#### **Details**

If the data parameter is a dataframe the colnames.FCS parameter is ignored but if the data parameter is a flowset, the colnames.FCS parameter is required.

#### Value

a numeric matrix without the cell markers to exclude

export

Exportation of SPADEVizR objects

#### **Description**

Exports a SPADEVizR object into a tab separated file.

## Usage

```
export(object, filename = "export.txt")
## S4 method for signature 'Results'
export(object, filename = "export.txt")
## S4 method for signature 'AC'
export(object, filename = "export.txt")
## S4 method for signature 'DEC'
export(object, filename = "export.txt")
## S4 method for signature 'CC'
export(object, filename = "export.txt")
```

filter.medians

```
## S4 method for signature 'PhenoProfiles'
export(object, filename = "export.txt")

## S4 method for signature 'EnrichmentProfiles'
export(object, filename = "export.txt")
```

## **Arguments**

object a SPADEVizR object

filename a character indicating the location of output file

## Value

none

filter.medians

Internal - filter medians to exclude from a matrix

## Description

This function is used internally to remove raw or transform medians from SPADE matrix. CVS medians are always removed.

## Usage

```
filter.medians(data, use.raw.medians = FALSE)
```

## **Arguments**

```
data a SPADE matrix
use.raw.medians
a logical specifying if "transformed" or "raw" medians will be use (FALSE by default)
```

## Value

a numeric matrix without the cell markers to exclude

generateReport 19

generateReport	Generate a report including SPADEVizR plots.	

#### **Description**

Generate a customizable PDF report based on SPADEVizR vizualisation features. Available plots are :

- "count" (included by default):Display an representation showing the number of cells for each cluster
- "tree" (included by default):Display a tree representation showing combined SPADE trees
- "pheno" (included by default):Display an heatmap representation
- "boxplot":Display a boxplot representation. This plot required to provide the 'conditions' parameter
- "kinetics":Display a kinetic representation for each cluster. This plot required to provide the 'assignments' parameter
- "stream" (included by default):Display a streamgraphViewer representation showing the evolution of cells abundance
- "cluster" (included by default):Display a parallel coordinate representation showing for each cluster the marker median expression
- "MDSclusters" (included by default):Display the cluster similarities using MDS
- "MDSsamples":Display the samples similarities using MDS. This plot required to provide the 'assignment' parameter
- "disto" (included by default): Display a distogram representation showing the marker co-expressions
- "kinetics\_cluster":Display a kinetic representation and a parallel coordinate juxtaposed (are arranged one on the side of the other) for each cluster
- "boxplot\_cluster":Display a boxplot representation and a parallel coordinate juxtaposed (are arranged one on the side of the other) for each cluster

#### Usage

```
generateReport(Results, PDFfile = "report.pdf", plot.names = c("count",
   "pheno", "cluster", "tree", "disto", "stream", "MDSclusters"),
   clusters = NULL, markers = NULL, assignments = NULL,
   conditions = NULL, stat.objects = list(), profile.objects = list(),
   width = 29.7, height = 21)
```

#### **Arguments**

Results	a 'SPADEResults' or 'Result' object
PDFfile	a character specifying the output path
plot.names	a character vector specifying the names (see details) and the order of the desired plots
clusters	a character vector of clusters to include in the report (all will be included by default)
markers	a character vector of markers to include in the report (all will be included by default)

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assignments	a 2 column data.frame with the samples names in row names providing firstly the time-points (numeric) and secondly the individuals (character) of the experiment $\frac{1}{2}$	
conditions	conditions a named vector providing the correspondence between a sample name (in row names) and the condition of this sample or NA to exclude	
stat.objects	a vector of stat.object to be displayed in the report (object of class 'DEC', 'AC' or 'CC')	
profile.objects		
	a vector of profile.objects to be displayed in the report (object of class 'Pheno-Profiles' or 'EnrichmentProfiles')	
width	a numeric specifying the plot width	
height	a numeric specifying the plot height	

ggheatmap

Internal - Create a list of elements allowing to build a heatmap

## Description

This function is used internally to build the element needed for an heatmap

## Usage

```
ggheatmap(matrix, dendrogram.type = "rectangle", num = 5,
  clustering.markers = NULL)
```

## Arguments

```
matrix
                  a numeric matrix containing the markers expression categories
dendrogram.type
                  a character specifying the look of dendrograms ("rectangle" or "triangle", "rect-
                  angle" by default)
                  a numeric value specifying the number of markers expression categories
num
clustering.markers
                  a character vector of clustering markers
```

## Value

```
a list of 3 plots (top dendrogram, right dendrogram, heatmap)
```

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ggheatmap.plot

Internal - Generate an heatmap by assembling elements

## **Description**

This function displays the heatmap elements build by 'ggheatmap()'

## Usage

```
ggheatmap.plot(list, col.width = 0.15, row.width = 0.15)
```

## **Arguments**

list the list of ggplot object provided by ggheatmap

col.width size of horizontal dendrogram row.width size of vertical dendrogram

#### Value

a ggplot2 axis

g\_axis

Internal - Extraction of ggplot axes

## Description

This function is used internally to extract axes from a 'ggplot' objet.

## Usage

```
g_axis(gplot, x.axis = !y.axis, y.axis = !x.axis)
```

## **Arguments**

gplot a 'ggplot' plot

x.axis a logical value specifying if the x-axis must be extracty.axis a logical value specifying if the y-axis must be extract

## **Details**

It is to note that 'x' and 'y' are mutuality excluded (both cannot be both TRUE) with priority to 'x'.

## Value

```
a 'ggplot' axis object
```

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g\_dendro

Internal - Build a dendrograms plot

## Description

This function is used internally to generate a 'ggplot' dendrogram.

## Usage

```
g_dendro(dist, row = !col, col = !row)
```

## Arguments

dist a numeric matrix containing distances between objects

row a logical value specifying if the horizontal dendrogram must be computed col a logical value specifying if the vertical dendrogram must be computed

## **Details**

It is to note that 'row' and 'col' are mutuality excluded (both cannot be both TRUE) with priority to row.

#### Value

a 'ggplot' dendrogram object

g\_legend

Internal - Extraction of ggplot legend

## Description

This function is used internally to extract the legend from a 'ggplot' object.

## Usage

```
g_{legend(gplot)}
```

## **Arguments**

gplot

a 'ggplot' plot

## Value

a 'ggplot' legend object

identifyAC 23

identifyAC	Identification of the Abundant Clusters	

## **Description**

This function is used to identify the abundant clusters. That is to say clusters that have cell abundance statistically greater than a specific threshold.

## Usage

```
identifyAC(Results, condition, use.percentages = TRUE, method = "t.test",
  method.adjust = NULL, th.pvalue = 0.05, th.mean = 0)
```

## **Arguments**

Results	a 'Results' or 'SPADEResults' object
condition	a named vector providing the correspondence between a sample name (in row names) and the logical value TRUE to test abundance for this sample or FALSE otherwise
use.percentage	S
	a logical specifying if the computations should be performed on percentage
method	a character specifying the statistical method used to identify the abundant clusters. The parameter can take the values "t.test" or "wilcox test"

method.adjust a character specifying if the p-values should be corrected using multiple cor-

rection methods among: "holm", "hochberg", "hommel", "bonferroni", "BH",

"BY" and "fdr" (from 'stats::p.adjust' method)

th.pvalue a numeric specifying the p-value threshold

th.mean a numeric specifying the abundance mean threshold

## Value

a S4 object of class 'AC'

identifyCC	Identification of the correlation of SPADE cluster with a phenotype

## **Description**

This function is used to identify correlated clusters. That is to say clusters that correlate with a phenotypic variable.

## Usage

```
identifyCC(Results, variable, use.percentages = TRUE, method = "pearson",
  method.adjust = NULL, th.pvalue = 0.05, th.correlation = 0.75)
```

24 identifyDEC

#### **Arguments**

Results a 'Results' or 'SPADEResults' object

variable a numerical named vector providing the correspondence between a sample name

(in rownames) and the specific phenotype or NA to ignore a sample

use.percentages

a logical specifying if the computations should be performed on percentage

method a character indicating the correlation method to use: "pearson", "spearman"

method.adjust a character specifying if the p-values should be corrected using multiple correction methods among: "holm", "hochberg", "hommel", "bonferroni", "BH",

rection methods among: noim, nochberg, nommer, bonierroni

"BY" and "fdr" (from 'stats::p.adjust' method)

th.pvalue a numeric specifying the p-value threshold

th.correlation a numeric specifying the absolute value of the correlation coefficient threshold

#### Value

a S4 object of class 'CC'

identifyDEC

Identification of the Differentially Enriched Clusters

#### **Description**

This function is used to identify differentially enriched clusters. That is to say clusters that are differentially abundant between two biologicals conditions.

## Usage

```
identifyDEC(Results, conditions, use.percentages = TRUE, method = "t.test",
  method.adjust = NULL, method.paired = FALSE, th.pvalue = 0.05,
  th.fc = 1)
```

#### **Arguments**

Results a 'Results' or 'SPADEResults' object

conditions a named vector providing the correspondence between a sample name (in row

names) and the condition of this sample: NA to exclude a sample from tests, 1

or 2 to attribute this sample, respectively to the first or second condition

use.percentages

a logical specifying if the computations should be performed on percentage

method a character specifying the name of the statistical test to use "t.test" or "wilcox.test"

method.adjust a character specifying if the p-values should be corrected using multiple cor-

rection methods among: "holm", "hochberg", "hommel", "bonferroni", "BH",

"BY" and "fdr" (from 'stats::p.adjust' method)

method.paired a logical indicating if the statistical test must be performed in a paired manner

th.pvalue a numeric specifying the p-value threshold th.fc a numeric specifying the fold-change threshold

#### Value

a S4 object of class 'DEC'

importSPADEResults 25

importResults	Import clustering results generated by other algorithms	

## **Description**

The 'importX()' function imports cell clustering results from two dataframes ('cells.count' and 'marker.expressions'). This function returns a 'Result' object.

## Usage

```
importResults(cells.count, marker.expressions)
```

## **Arguments**

cells.count a dataframe of cells abundances with clusters in row and samples in column marker.expressions

a dataframe containing median marker expression values for each cluster of each sample. In additions of markers, the 2 two first columns are are dedicated to "cluster" and "sample"

#### **Details**

The 'cells.count' dataframe must be formated as following:

cluster	sample1	sample1
cluster1	749	5421
cluster2	450	412

The 'marker.expressions' dataframe must be formated as following:

sample	cluster	marker1	marker2
sample1	cluster1	0.2	0.3
sample1	cluster2	0.1	0.3
sample2	cluster1	0.5	2.3
sample2	cluster2	1	1.3

## Value

a S4 object of class 'Results'

importSPADEResults	Import clustering results generated by SPADE

## Description

The 'importSPADEResults()' function imports SPADE cell clustering results from a specified path. This function returns a 'SPADEResult' object.

26 kinetics Viewer

This function import the expression matrix and count matrix as well as the SPADE tree. This function apply an hyperbolic sine transformation to imported FCS data and compute the maker range quantiles.

## Usage

```
importSPADEResults(path, dictionary = data.frame(),
  exclude.markers = c("cell_length", "FileNum", "density", "time"),
  probs = c(0.05, 0.95), use.raw.medians = FALSE,
  quantile.approximation = FALSE)
```

#### **Arguments**

path a character specify the path of SPADE results folder

dictionary a two column dataframe providing the correspondence between the original

marker names (first column) and the real marker names (second column)

exclude.markers

a character vector of markers to exclude (case insensitive)

probs a vector of probabilities with 2 values in [0,1] to compute maker range quantiles.

First is the lower bound and second is the upper bound.

use.raw.medians

a logical specifying if "transformed" or "raw" medians will be use in the cluster

expression matrix (FALSE by default)

quantile.approximation

a logical specifying if maker range quantiles are computed using all cells (FALSE),

or is the means of the quantile of each samples (TRUE)

#### **Details**

The computation of maker range quantiles can be approximated using 'quantile.approximation' parameter which is more efficient in term of loading time and memory usage.

#### Value

a S4 object of class 'SPADEResults'

kineticsViewer

Visualization of cluster enrichment profiles kinetics

#### **Description**

Generates a kinetics plot representation showing for each cluster its enrichment profiles at each time-point of each individual.

## Usage

```
kineticsViewer(Results, assignments, clusters = NULL,
  use.percentages = TRUE)
```

load.flowSet 27

#### **Arguments**

Results a SPADEResults or Results object

assignments a 2 column data frame with the samples names in row names providing firstly the

time-points (numeric) and secondly the individuals (character) of the experiment

clusters a character vector containing the clusters names to be visualized (by default all

clusters will be displayed)

use.percentages

a logical specifying if the visualization should be performed on percentage

#### **Details**

XXX

#### Value

a 'ggplot' object

load.flowSet

Load FCS files object into a 'SPADEResult' object

## **Description**

This function loads the FCS files to the 'flowset' slot of the 'SPADEResult' object.

## Usage

```
load.flowSet(SPADEResult = NULL, fcs.files, dictionary, exclude.markers)
```

## **Arguments**

SPADEResult a SPADEResult object (optional)

fcs.files a character vector containing the absolute path of the original FCS files

dictionary a two column data frame providing the correspondence between the original

marker names (first column) and the real marker names (second column)

exclude.markers

a character vector of markers to exclude (case insensitive)

## **Details**

If a 'SPADEResult' object is provided, others parameters ('fcs.files', 'dictionary', 'exclude.markers') will be ignored.

#### Value

```
a S4 'flowSet' object
```

28 names

MDSViewer	Visualization of SPADE cluster or sample similarities using MDS

## Description

Generate a Multidimensional Scaling (MDS) representation showing the similarities between SPADE results based on theirs enrichment.

## Usage

```
MDSViewer(Results, use.percentages = TRUE, assignments, clusters = NULL,
   space = "clusters", dist.method = "euclidean")
```

## **Arguments**

Results	a SPADEResults or Results object
use.percentage	S
	a logical specifying if the visualization should be performed on percentage
assignments	a 2 column data.frame with the samples names in row names providing firstly the time-points (numeric) and secondly the individuals (character) of the experiment
clusters	a character vector containing the clusters names to be visualized (by default all clusters will be displayed)
space	a character specifying the space ("clusters" or "samples", "cluster" by default)
dist.method	a character string containing the name of the distance measure to use

## **Details**

The 'space' parameter specifying if the cluster or sample similarities will be determined using MDS. When the 'space' parameter is equal to 'sample', the 'assignments' parameter is required.

## Value

a list of ggplot objects

names	Definition of class names	
-------	---------------------------	--

## Description

Provides the name of each SPADEVizR object

PhenoProfiles-class 29

#### Usage

```
## S4 method for signature 'Results'
names(x)

## S4 method for signature 'SPADEResults'
names(x)

## S4 method for signature 'AC'
names(x)

## S4 method for signature 'DEC'
names(x)

## S4 method for signature 'CC'
names(x)

## S4 method for signature 'PhenoProfiles'
names(x)

## S4 method for signature 'EnrichmentProfiles'
names(x)
```

#### **Arguments**

Χ

a SPADEVizR object

## Value

a character providing the name of the object

PhenoProfiles-class

Classification of clusters based on their phenotype profiles (Pheno-Profiles class) definition

## **Description**

The 'PhenoProfiles' is a S4 object containing the information related to the cluster classification based on theirs marker expressions.

This object contains all information about the classification method and parameters used.

#### **Details**

Five methods are available to classify cellular clusters: 'hierarchical\_k', 'hierarchical\_h', 'kmeans', 'eigencell' and 'clique'. Each method can parameterized using the 'method.parameter' parameter.

The 'print()' and 'show()' can be used to display a summary of this object. Moreover all information about this object could be saved as a tab separated file using the 'export()' method. This object is returned by the 'classifyPhenoProfiles()' function.

30 phenoViewer

#### **Slots**

cluster.number a numeric providing the number of cluster

method a character specifying the method used to classify cluster

method.parameter a named list of parameters used by the classification method

cluster.size a numeric vector containing the number of cells associated with each cluster (– sum of all samples –)

class.number a numeric value specifying the number of clusters

classes a two column dataframe with the cluster in first column and corresponding classe in the second column

phenoViewer

Visualization of all clusters phenotypes as an heatmap

## **Description**

Generates an heatmap representation showing for all clusters the marker median expressions.

## Usage

```
phenoViewer(SPADEResults, num = 5)
```

#### **Arguments**

SPADEResults a SPADEResults object (Results object is not accepted)

num a numeric value specifying the number of markers expression categories to use

#### **Details**

For each marker, median expressions are discretized in severals categories corresponding to the heat intensities. This number of categories is provided using 'num' parameter.

By default the 'pheno.table' parameter is null and will be automatically compute using the 'computePhenoTable' function.

## Value

a list of ggplot objects

plot 31

plot

Graphical representation for some SPADEVizR objects

## Description

This function generates a graphical representation for 'AC', 'DEC', 'CC', 'PhenoProfiles' and 'EnrichmentProfiles' objects.

## Usage

```
plot(x, y = NULL, ...)
## S4 method for signature 'DEC,missing'
plot(x, y = NULL, ...)
## S4 method for signature 'AC,missing'
plot(x, y = NULL, ...)
## S4 method for signature 'CC,missing'
plot(x, y = NULL, ...)
## S4 method for signature 'PhenoProfiles,missing'
plot(x, y = NULL, ...)
## S4 method for signature 'EnrichmentProfiles,missing'
plot(x, y = NULL, ...)
```

## **Arguments**

```
x a 'AC', 'DEC', 'CC', 'PhenoProfiles' or 'EnrichmentProfiles' object
y a supplementary parameter transmited to 'AC', 'DEC' or 'CC' object
... some supplementaries parameters transmited to 'AC', 'DEC' or 'CC' object
```

#### Value

```
a 'ggplot' object
```

print

Textual previews for all SPADEVizR objects

## Description

Prints a previews for a SPADEVizR object.

32 profiles Viewer

#### Usage

```
## S4 method for signature 'Results'
print(x)

## S4 method for signature 'SPADEResults'
print(x)

## S4 method for signature 'AC'
print(x)

## S4 method for signature 'DEC'
print(x)

## S4 method for signature 'CC'
print(x)

## S4 method for signature 'PhenoProfiles'
print(x)

## S4 method for signature 'EnrichmentProfiles'
print(x)
```

#### **Arguments**

Χ

a SPADEVizR object

## Value

none

profilesViewer

profilesViewer

## Description

Generate a graph representation of PhenoProfiles classes

## Usage

```
profilesViewer(profile.object)
```

## **Arguments**

profile.object a PhenoProfiles object or an EnrichmentProfiles object

## **Details**

XXX

#### Value

```
a 'ggplot' object
```

rename.markers 33

rename.markers

Internal - Renaming cell markers

#### **Description**

This function is used internally to rename the cell markers based on a dictionary.

#### Usage

```
rename.markers(header, dictionary)
```

#### **Arguments**

header a character vector containing the original maker names

dictionary a character vector containing a correspondence between the original and the new

marker names

#### **Details**

Dictionary is a data.frame used to rename the marker names. The first column must correspond to the original marker names, the second column must correspond to the new marker names.

#### Value

a character vector containing the renamed marker names

Results-class

Results class definition

#### **Description**

The Results object is a S4 object containing cell clustering results obtained from various automatic gating algorithms.

This object mainly stores the count matrix (i.e. the number of cells associated with each cluster of each sample) and the cell cluster phenotypes (i.e. the marker median expressions for each cluster). It is to note that the Results object is a super class of the SPADEResult object.

#### **Details**

The 'cells.count' dataframe stores the number of cells associated with each cluster of each sample. This dataframe has in row the clusters and in column the samples.

The 'marker expressions' dataframe stores the marker median expressions for each cluster. This dataframe has in the first the sample names, in the second column the cluster names, and the maker median expressions in the others columns.

The 'print()' and 'show()' can be used to display a summary of this object. Moreover all information about this object could be saved as a tab separated file using the 'export()' method. This object is returned by the 'importX()' function.

34 show

#### **Slots**

cells.count a dataframe containing the number of cells for each cluster of each sample
marker.expressions a numerical dataframe containing marker median expressions for each cluster of each sample
sample.names a character vector containing the sample names
marker.names a character vector containing the markers names
cluster.number a numeric specifying the number of cell clusters

show

Textual previews for SPADEVizR objects

## **Description**

Show a previews for a SPADEVizR object.

#### Usage

```
## S4 method for signature 'Results'
show(object)

## S4 method for signature 'SPADEResults'
show(object)

## S4 method for signature 'AC'
show(object)

## S4 method for signature 'DEC'
show(object)

## S4 method for signature 'CC'
show(object)

## S4 method for signature 'PhenoProfiles'
show(object)

## S4 method for signature 'EnrichmentProfiles'
show(object)
```

## **Arguments**

object a SPADEVizR object

#### Value

none

SPADEResults-class 35

SPADEResults-class

SPADEResults class definition

#### **Description**

The 'SPADEResults' object is a S4 object containing cell clustering results obtained from SPADE.

This object inherits from the 'Result' object and stores the count matrix (i.e. the number of cells associated with each cluster of each sample) and the cell cluster phenotypes (i.e. the marker median expressions for each cluster). In addition to the 'Result' object, the 'SPADEResults' object contains information about SPADE clustering results, such as the SPADE tree, the clustering makers and the FCS files.

#### **Details**

The 'print()' and 'show()' can be used to display a summury of this object. Moreover all information about this object could be saved as a tab separated file using the 'export()' method. This object is returned by the 'importSPADEResult()' function.

#### **Slots**

use.raw.medians a logical specifying if the marker expressions correspond to the raw or transformed data

dictionary a two column data.frame providing the correspondence between the original marker names (first column) and the real marker names (second column)

marker.clustering a logical vector specifying marker that have been used during the clustering precedure

flowset a flowSet object containing the imported SPADE FCS file

fcs.files a character vector containing the absolute path of the original FCS files

quantiles a numeric data.frame containing the quantiles for each each markers of cluster

graph a igraph object containing the SPADE tree

graph.layout a numeric matrix containing the layout of the SPADE tree

streamgraphViewer

Visualization of cluster abundance dynamics

#### **Description**

Generate a streamgraph representation showing the dynamic evolution of the number of cells in clusters across samples.

## Usage

```
streamgraphViewer(Results, order = NULL, clusters = NULL,
  use.relative = FALSE)
```

36 tree Viewer

#### **Arguments**

Results a SPADEResults or Results object

order a named vector providing the correspondence between a sample name (in row-

names) and an integer ordering samples (NA to exclude a sample)

clusters a character vector containing the clusters names to be vizualised (by default all

clusters will be displayed)

use.relative a logical specifying if the visualization should be performed on relative abun-

dance

#### **Details**

XXX

#### Value

a 'ggplot' object

treeViewer

Visualization of combined SPADE trees

## **Description**

Generates a tree representation showing combined SPADE trees.

## Usage

```
treeViewer(SPADEResults, samples = NULL, stat.object = NULL,
    marker = NULL)
```

## Arguments

SPADEResults a SPADEResults object (Results object is not accepted)

samples a named vector providing the correspondence between samples name (in row-

name) and the logical value TRUE to use these samples (all samples by default)

stat.object an AC, DEC or CC object to highligth identified significant clusters in the

SPADE tree

marker a character specifying the marker name to display

#### **Details**

The size of tree nodes are related to the number of cells in each cluster. If the 'stat.object' parameter is provided node outlines are colored according to clusters signifiance. If the 'marker' parameter is provided, the nodes are colored according to mean expression for the selected marker using selected samples.

#### Value

a list of ggplot objects

unload.flowSet 37

unload.flowSet

Unload 'flowSet' object from a 'SPADEResult' object

#### **Description**

This function unloads the 'flowSet' object in a 'SPADEResult' object.

#### Usage

```
unload.flowSet(SPADEResult)
```

#### **Arguments**

```
SPADEResult a SPADEResult object (optional)
```

#### Value

The new 'SPADEResult' object

volcanoViewer

Visualization of differentially enriched clusters

## Description

Generates a Volcano plot representation showing for each cluster

## Usage

```
volcanoViewer(DEC = NULL, fc.log2 = TRUE, show.cluster.sizes = TRUE,
    show.all_labels = FALSE)
```

## **Arguments**

#### **Details**

By default, only significant differentially enriched clusters are labeled. Labels for all clusters can be displayed by setting the 'all.label' parameter to TRUE.

#### Value

```
a 'ggplot' object
```

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