R documentation

of all in 'man'

October 8, 2018

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| | AutoPipe_tSNE | |

Description

This function is used to upload a table into R for further use in the AutoPipe

Usage

```
AutoPipe_tSNE(me,perplexity=30,max_iter=500)
```

Arguments

me The path of the expression table perplexity numeric; Perplexity parameter

max_iter integer; Number of iterations (default: 1000)

groups_men the data frame with the group clustering that the function Groups_Sup or top_supervised

(2. place on the list) returns with the data about each sample and its coresspond-

ing cluster.

2 Groups_Sup

| cons_clust | A function to plot do a Consensus clustering to validate the results |
|------------|--|
| | |

Description

this function calls the ConsensusClusterPlus function with thedaraset and plots a plot with the heatmaps of the clustering for each number of clusters from 2 to max_clust

Usage

```
cons_clust(data,max_clust,TOPgenes)
```

Arguments

data this is the data for the ConsensusClusterPlus

max_clust the max number of clusters that should be evaluated.

TOPgenes the number of the top genes to choose for the clustering

Value

plots a plot with all the heatmaps from the ConsensusClusterPlus for the number ofd clusters 2 to max_clust the same return value as the COnsensusClusterPlus

Examples

```
data(rna)
cons_clust(rna,5,TOPgenes=150)
```

Groups_Sup

cluster the samples

Description

This function clusters the samples into x clusters.

Usage

```
Groups_Sup(me_TOP, me, number_of_k,TRw)
```

Arguments

me_TOP the matrix with the n top genes, usually the from output of the function TopPAM the original expression matrix. (with genes in rows and samples in columns).

number_of_k the number of clusters

TRw threshold for the elemenation of the samples with a Silhouette width lower than

TRw. Default value is -1.

read_expression_file 3

Examples

```
## load data
library(org.Hs.eg.db)
data(rna)
me_x=rna
res<-AutoPipe::TopPAM(me_x,max_clusters = 8, TOP=1000)
me_TOP=res[[1]]
number_of_k=res[[3]]
File_genes=Groups_Sup(me_TOP, me=me_x, number_of_k,TRw=-1)
groups_men=File_genes[[2]]
me_x=File_genes[[1]]</pre>
```

Description

This function is used to upload a table into R for further use in the AutoPipe

Usage

```
read_expression_file(file, format="csv", sep";",gene_name="SYMBOL",Trans=F)
```

Arguments

file

| format | The format of the table "csv" or "txt" |
|-----------|---|
| sep | The seperator of the input table |
| gene_name | Genes are given in "SYMBOL" or "ENTREZID" |
| Trans | Need Matrix Transpose TRUE or FALSE |

The path of the expression table

Value

A data.frame with a gene expression matrix

4 rna

rna

rna egene expression of 48 meningiomas

Description

A dataset containing the gene expression data od 48 meningioma tumors

Usage

rna

Format

A data frame with 17288 rows and 48 variables:

```
BT_1008 sample BT_1008,
```

```
BT_700 sample BT_700,
BT_738 sample BT_738,
BT_751 sample BT_751,
BT_771 sample BT_771,
BT_797 sample BT_797,
BT_803 sample BT_803,
BT_808 sample BT_808,
BT_820 sample BT_820,
BT_837 sample BT_837,
BT_855 sample BT_855,
BT_862 sample BT_862,
BT_873 sample BT_873,
BT_882 sample BT_882,
BT_887 sample BT_887,
BT_900 sample BT_900,
BT_905 sample BT_905,
BT_907 sample BT_907,
BT_920 sample BT_920,
BT_944 sample BT_944,
BT_962 sample BT_962,
BT 963 sample BT 963,
BT_982 sample BT_982,
BT_990 sample BT_990, ...
```

Supervised_Cluster_Heatmap

Produce a Heatmap using a Supervised clustering Algorithm

Description

This function produces a plot with a Heatmap using a supervised clustering algorithm which the user choses. with a the mean Silhouette width plotted on the right top corner and the Silhouette width for each sample on top. On the right side of the plot the n highest and lowest scoring genes for each cluster will added. And next to them the coressponding pathways (see Details)

Usage

```
Supervised_Cluster_Heatmap(groups_men, gene_matrix, method="PAMR",TOP=1000,TOP_Cluster=150, show_sil=FALSE,show_clin=FALSE,genes_to_print=5, print_genes=FALSE,samples_data=NULL,colors="RdBu", GSE=FALSE,topPaths=5,db="c2",plot_mean_sil=FALSE,stats_clust =NULL,threshold=2)
```

Arguments

| groups_men | the data frame with the group clustering that the function Groups_Sup or top_supervised (2. place on the list) returns with the data about each sample and its coressponding cluster. |
|----------------|---|
| gene_matrix | the matrix of n selected genes that the function Groups_Sup returns |
| method | the method to cluster of Clustering. The default is "PAMR" which uses the pamr library. other methods are SAM and our own "EXReg" (see details) |
| TOP | the number of the top genes to take. the default value is 1000. |
| TOP_Cluster | a numeric variable for the number of genes to include in the clusters. Default is 150. |
| show_sil | a logical value that indicates if the function should show the Silhouette width for each sample. Default is FALSE. |
| show_clin | a logical value if TRUE the function will plot the clinical data provided by the user. Default value is FALSE. |
| genes_to_print | the number of genes to print for each cluster. this function adds on the right side. of the heatmap the n highest expressed genes and the n lowest expressed genes for each cluster. Default value is 5. |
| print_genes | a logical value indicating if or not to plot the TOP genes for each cluster.Default value is FALSE. |
| samples_data | the clinical data provided by the user to plot under the heatmap. it will be plotted only if show_clin is TRUE. Default value is NULL. see details for format. |
| colors | the colors for the Heatmap. The function RColorBrewer palletes. |
| GSE | a logical variable that indicates wether to plot thr Gene Set Enrichment Analysis next to the heatmap. Default value is FALSE. |
| topPaths | a numerical value that says how many pathways the Gene Set Enrichment plots should contain fo each cluster. Default value is 5. |
| db | a value for the database for the GSE to be used. Default value is "c1". the paramater can one of the values: "c1","c2","c3",c4","c5","c6","c7","h". See the broad institue GSE GSE webpage for further information in each dataset. |
| plot_mean_sil | A logical value. if TRUE the function plots the mean of the Silhouette width for each cluster number or gap statistic. |
| stats_clust | A vector with the mean Silhouette widths or gap statistic for the number of clusters. The first value should be for 2 Clusters. 2nd is for 3 clusters and so on. |
| threshold | the threshhold for the pam analysis default is 2. |

Details

sample data should be a data.frame with the sample names as rownames and the clinical triats as columns. each trait must be a numeric variable.

TopPAM 7

Examples

TopPAM

Compute Top genes

Description

This function computes the n=TOP genes and the the best number of clusters

Usage

```
TopPAM(me, max_clusters=15,TOP=1000)
```

Arguments

me a matrix with genes in rows and samples in columns

max_clusters max. number of clusters to check
TOP the number of genes to take.

B integer, number of Monte Carlo ("bootstrap") samples.

Gap_stat A logical value that indicates wether or not to calculate the Silhouette width and

Gap statistic.

Details

we use the clusGap algorithm from the package cluster to calculate the Gap statistic.

Value

a list of 1. A matrix with the top genes 2. A list of means of the Silhouette width for each number of clusters. 3. The optimal number of clusters. 4. gap_st the gap statistic of the clustering 5. best number of clusters according to the gap statistic.

8 top_supervised

Examples

```
##load the org.Hs.eg Library
library(org.Hs.eg.db)
#' ## load data
data(rna)
me_x=rna
res<-AutoPipe::TopPAM(me_x,max_clusters = 8, TOP=1000)
me_TOP=res[[1]]
number_of_k=res[[3]]</pre>
```

top_supervised

A Function for Assisting Supervised Clustering

Description

when perfoming a supervised clustering the user should run this function in order to get the best results.

Usage

```
top_supervised(me,TOP=1000,cluster_which,TRw=-1)
```

Arguments

me the matrix of the gene exporessions, the olums should be the samples and the

colnames the sample names the rownames should be the genes . at best the

ENTEREZID

TOP the top genes to choose, default is 1000.

cluster_which a dataframe with the supervised clustering arrangment of the samples. the

dataframe should have the sample names in the first column and the clustering

in the secound column.

TRw the threshhold for excluding samples with silhouette width < TRw

Value

a list. the first place is the expression matrix, the secound is the silhouette for each sample.

Examples

```
library(org.Hs.eg.db)
data(rna)
cluster_which<-cbind(colnames(rna),c(rep(1,times=24),rep(2,times=24)))
me_x=rna
## calculate best number of clusters and
res<-top_supervised(me_x,TOP = 1000,cluster_which)
me_TOP=res[[1]]</pre>
```

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UnSuperClassifier

Unsupervised Clustering

Description

A function for unsupervised Clustering of the data

Usage

```
UnSuperClassifier(data,clinical_data=NULL,thr=2,TOP_Cluster=150)
```

Arguments

data the data for the clustering. Data should be in the following format: samples in

columns and the genes in the rows (colnames and rownames accordingly). The

rownames should be Entrez ID in order to plot a gene set enrichment analysis.

clinical_data the clinical data provided by the user to plot under the heatmap. it will be plotted

only if show_clin is TRUE. Default value is NULL. see details for format.

thr The threshold for the PAMR algorithm default is 2.

TOP_Cluster numeric; Number of genes in each cluster.

Details

sample data should be a data frame with the sample names as rownames and the clinical triats as columns. each trait must be a numeric variable. @return the function is an autated Pipeline for clustering it plot cluster analysis for the geneset

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