Package: tightClust (via r-universe)

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Type Package

Title Tight Clustering

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tightClust-package Tight Clustering Package

Description

This package could perform tight clustering algorithm proposed by George C. Tseng and Wing H. Wong.

Details

Package: tightClust Type: Package Version: 1.0

Date: 2012-08-28 License: GPL (>=2)

Author(s)

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References

George C. Tseng and Wing H. Wong. (2005) Tight Clustering: A Resampling-based Approach for Identifying Stable and Tight Patterns in Data. Biometrics.61:10-16.

plot.tight.clust Plot tight cluster result

Description

A function to plot the heatmap of the tight cluster result.

Usage

```
## S3 method for class 'tight.clust'
plot(x, standardize.gene = TRUE, order.sample = FALSE, plot.noise=TRUE, ...)
```

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Arguments

x Return value of the tight.clust function

standardize.gene

If standardize each gene vector to mean 0 and sd 1.

order.sample It specifies whether to order samples (features) using the hierarchical clustering

method.

plot.noise It specifies whether to plot the remaining noise genes (objects).

... Arguments to image.

Author(s)

Chi Song <chs108@pitt.edu>

References

George C. Tseng and Wing H. Wong. (2005) Tight Clustering: A Resampling-based Approach for Identifying Stable and Tight Patterns in Data. Biometrics.61:10-16.

tclust.test.data

test data for tight clustering package

Description

Sample microarray data

Usage

```
data(tclust.test.data)
```

Format

The data is a list of 3 items:

GeneID ID of each gene

Annotation Annotation information of each gene

Data Data matirx of gene expression: each row represent one gene; each column represent one sample

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tight.clust	Tight Clustering	

Description

This function could perform the tight clustering algorithm.

Usage

```
tight.clust(x, target, k.min, alpha = 0.1, beta = 0.6,
top.can = 7, seq.num = 2, resamp.num = 10,
samp.p = 0.7, nstart = 1, remain.p = 0.1,
k.stop = 5, standardize.gene=TRUE, random.seed=NULL)
```

Arguments

	X	Input data, should be matrix. Each row represents a gene (object) to be clustered. Gene (object) names are usually represented in the row names and sample (feature) names are in the column names of the matrix.		
	target	The total number of clusters that the user aims to find.		
	k.min	The starting point of k0. See 'Details' for more information.		
	alpha	The threshold of comembership index. Default value is suggested to be used.		
	beta	The threshold of clusters stably found in consecutive $k0$. Default value is suggested to be used.		
	top.can	The number of top (size) candidate clusters for a specific $k\theta$. Default value is suggested to be used.		
	seq.num	The number of subsequent $k\theta$ that finds the tight cluster. Default value is suggested to be used.		
	resamp.num	Total number of resampling to obtain comembership matrix. Default value is suggested to be used.		
	samp.p	Percentage of subsamples selected. Default value is suggested to be used.		
	nstart	Number of different random inital for K-means. Default value is suggested to be used.		
	remain.p	Stop searching when the percentage of remaining points <= remain.p. Default value is suggested to be used.		
	k.stop	Stop decreasing k0 when k0<=k.stop. Default value is suggested to be used.		
standardize.gene				
		It specifies whether to standardize each gene vector to mean 0 and sd 1. Default value is suggested to be used.		
	random.seed	If random. seed is NULL no random seed will be set. If random. seed is a number, it will be used as the random seed. This parameters should be used to get the same result for different runs.		

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Details

Tight clustering method is a resampling-evaluated clustering method that aims to directly identify tight clusters in a high-dimensional complex data set and allow a set of scattered objects without being clustered. The method was originally developed for gene cluster analysis in microarray data but can be applied in any complex data. The most important parameter is k.min. A large k.min results in smaller and tighter clusters. Normally k.min>=target+5 is suggested. All other parameters do not affect the quality of final clustering results too much and are suggested to remain unchanged.

Value

Returned value is a "tight.clust" object (list). The first element is the original data matrix. The second element is a vector representing the cluster identity (-1: scattered gene set; 1: the first cluster; 2: the second cluster; ...). The third element is a vector of the size of each tight cluster.

Author(s)

Chi Song <chs108@pitt.edu>

References

George C. Tseng and Wing H. Wong. (2005) Tight Clustering: A Resampling-based Approach for Identifying Stable and Tight Patterns in Data. Biometrics.61:10-16.

Examples

```
## load the test dataset
data(tclust.test.data)
## find 10 tight clusters
ptm<-proc.time()</pre>
## k.min=25, tighter clusters will be found
## target=1 is used to save time, target=10 is recommended
tclust1<-tight.clust(tclust.test.data$Data, target=1, k.min=25, random.seed=12345)
proc.time()-ptm
## plot the heat map of cluster result
plot(tclust1)
## write the cluster result
write.tight.clust(tclust1)
ptm<-proc.time()</pre>
## k.min=10, looser clusters will be found
## target=1 is used to save time, target=5 is recommended
tclust2<-tight.clust(tclust.test.data$Data, target=1, k.min=10, random.seed=12345)
proc.time()-ptm
## plot the heat map of cluster result
plot(tclust2)
## write the cluster result
write.tight.clust(tclust2)
```

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write.tight.clust

Write tight cluster result

Description

A function to print the tight cluster result to a file or connection.

Usage

```
write.tight.clust(x, ...)
```

Arguments

x Return value of the tight.clust function

... Arguments to write.table.

Author(s)

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References

George C. Tseng and Wing H. Wong. (2005) Tight Clustering: A Resampling-based Approach for Identifying Stable and Tight Patterns in Data. Biometrics.61:10-16.

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