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Description It includes four methods: DCOL-based K-profiles clustering, non-linear network reconstruction, non-linear hierarchical clustering, and variable selection for generalized additive model. References: Tianwei Yu (2018) DOI:10.1002/sam.11381>; Haodong Liu and others (2016) DOI:10.1371/journal.pone.0158247>; Kai Wang and others (2015) DOI:10.1155/2015/918954>; Tianwei Yu and others (2010) DOI:10.1109/TCBB.2010.73>.	
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data.	gen	

Simulated Data Generation

Description

Generating gene matrix as a example of input.

Usage

```
data.gen(n.genes=100, n.samples=100, n.grps=10, aver.grp.size=10,
n.fun.types=6, epsilon=0.1, n.depend=0)
```

Arguments

n.genes the number of rows of the matrix.n.samples the number of columns of the matrix.

n.grps the number of hidden clusters.

aver.grp.size averge number of genes in a cluster. n.fun.types number of function types to use.

epsilon noise level.

n. depend data generation dependence structure. can be 0, 1, 2.

Details

The data generation scheme is described in detail in IEEE ACM Trans. Comput. Biol. Bioinform. 10(4):1080-85.

Value

return the data including gene and clustering.

data the gene matrix

grps the predicted clustering

Author(s)

Tianwei Yu<tyu8@emory.edu>

```
##generating a gene matrix with 100 genes, some in 5 clusters, and 100 samples per gene.
output<-data.gen(n.genes=100, n.samples=10, n.grps=5)
##get the gene matrix from the source of data.
matrix<-output$data
##get the hiden clusters from the source of data.
grps<-output$grp</pre>
```

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KPC

implementation of K-Profiles Clustering

Description

implementation of K-Profiles Clustering

Usage

```
KPC(dataset, nCluster, maxIter = 100, p.max = 0.2, p.min = 0.05)
```

Arguments

dataset the data matrix with genes in the row and samples in the column

nCluster the number of clusters K

maxIter the maximum number of iterations

p.max the starting p-value cutoff to exclude noise genesp.min the final p-value cutoff to exclude noise genes

Value

Return a list about gene cluster and the list of value p

cluster gene cluster
p.list a list of value p

Author(s)

Tianwei Yu <tianwei.yu@emory.edu>

References

http://www.hindawi.com/journals/bmri/aa/918954/

See Also

```
data.gen
```

```
## generating the data matrix & hiden clusters as a sample
input<-data.gen(n.genes=40, n.grps=4)
## now input includes data matrix and hiden clusters, so get the matrix as input.
input<-input$data
## set nCluster value to 4
kpc<-KPC(input,nCluster=4)</pre>
```

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```
##get the hiden cluster result from "KPC"
cluster<-kpc$cluster
##get the list of p
p<-kpc$p.list</pre>
```

nlhc

Non-Linear Hierarchical Clustering

Description

The non-linear hierarchical clustering based on DCOL

Usage

```
nlhc(array, hamil.method = "nn", concorde.path = NA,
use.normal.approx = FALSE, normalization = "standardize", combine.linear = TRUE,
use.traditional.hclust = FALSE, method.traditional.hclust = "average")
```

Arguments

the data matrix with no missing values array hamil.method the method to find the hamiltonian path. concorde.path If using the Concorde TSP Solver, the local directory of the solver use.normal.approx whether to use the normal approximation for the null hypothesis. normalization the normalization method for the array. combine.linear whether linear association should be found by correlation to combine with nonlinear association found by DCOL. use.traditional.hclust whether traditional agglomerative clustering should be used. method.traditional.hclust the method to pass on to hclust() if traditional method is chosen.

Details

Hamil.method: It is passed onto the function tsp of library TSP. To use linkern method, the user needs to install concord as instructed in TSP.

use.normal.approx: If TRUE, normal approximation is used for every feature, AND all covariances are assumed to be zero. If FALSE, generates permutation based null distribution - mean vector and a variance-covariance matrix.

normalization: There are three choices - "standardize" means removing the mean of each row and make the standard deviation one; "normal_score" means normal score transformation; "none" means do nothing. In that case we still assume some normalization has been done by the user such that each row has approximately mean 0 and sd 1.

combine.linear: The two pieces of information is combined at the start to initiate the distance matrix.

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Value

Returns a helust object same as the output of helust(). Reference: help(helust)

merge an n-1 by 2 matrix. Row i of merge describes the merging of clusters at step i

of the clustering. If an element j in the row is negative, then observation -j was merged at this stage. If j is positive then the merge was with the cluster formed

at the (earlier) stage j of the algorithm.

height a set of n-1 real values, the value of the criterion associated with the clusterig

method for the particular agglomeration

order a vector giving the permutation of the original observations suitable for plotting,

in the sense that a cluster plot using this ordering and matrix merge will not have

crossings of the branches.

labels labels for each of the objects being clustered

the call which produced the result

dist.method the distance that has been used to create d height.0 original calculation of merging height

Author(s)

Tianwei Yu <tianwei.yu@emory.edu>

References

http://www.ncbi.nlm.nih.gov/pubmed/24334400

See Also

```
data.gen
```

```
## generating the data matrix & hiden clusters as a sample
input<-data.gen(n.genes=40, n.grps=4)
## now input includes data matrix and hiden clusters, so get the matrix as input.
input<-input$data

nlhc.data<-nlhc(input)
plot(nlhc.data)
##get the merge from the input.
merge<-nlhc.data$merge</pre>
```

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Non-Linear Network reconstruction from expression matrix

Description

Non-Linear Network reconstruction method

Usage

```
nlnet(input, min.fdr.cutoff=0.05,max.fdr.cutoff=0.2, conn.proportion=0.007,
gene.fdr.plot=FALSE, min.module.size=0, gene.community.method="multilevel",
use.normal.approx=FALSE, normalization="standardize", plot.method="communitygraph")
```

Arguments

input the data matrix with no missing values.

min.fdr.cutoff the minimun allowable value of the local false discovery cutoff in establishing

links between genes.

max.fdr.cutoff the maximum allowable value of the local false discovery cutoff in establishing

links between genes.

conn.proportion

the target proportion of connections between all pairs of genes, if allowed by the

fdr cutoff limits.

gene.fdr.plot whether plot a figure with estimated densities, distribution functions, and (local)

false discovery rates.

min.module.size

the min number of genes together as a module.

gene.community.method

the method for community detection.

use.normal.approx

whether to use the normal approximation for the null hypothesis.

normalization the normalization method for the array.

plot.method the method for graph and community ploting.

Details

gene.community.method: It provides three kinds of community detection method: "mutilevel", "label.propagation" and "leading.eigenvector".

use.normal.approx: If TRUE, normal approximation is used for every feature, AND all covariances are assumed to be zero. If FALSE, generates permutation based null distribution - mean vector and a variance-covariance matrix.

normalization: There are three choices: "standardize" means removing the mean of each row and make the standard deviation one; "normal_score" means normal score transformation; "none"

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means do nothing. In that case we still assume some normalization has been done by the user such that each row has approximately mean 0 and sd 1.

plot.method: It provides three kinds of ploting method: "none" means ploting no graph, "communitygraph" means ploting community with graph, "graph" means ploting graph, "membership" means ploting membership of the community

Value

it returns a graph and the community membership of the graph.

algorithm The algorithm name for community detection

graph An igraph object including edges: Numeric vector defining the edges, the first

edge points from the first element to the second, the second edge from the third

to the fourth, etc.

community Numeric vector, one value for each vertex, the membership vector of the com-

munity structure.

Author(s)

Haodong Liu < liuhaodong 0828@gmail.com>

References

https://www.ncbi.nlm.nih.gov/pubmed/27380516

See Also

```
data.gen
```

```
## generating the data matrix & hiden clusters as a sample
  input<-data.gen(n.genes=40, n.grps=4)
## now input includes data matrix and hiden clusters, so get the matrix as input.
input<-input$data
##change the ploting method
  result<-nlnet(input,plot.method="graph")
  ## get the result and see it values
  graph<-result$graph ##a igraph object.
  comm<-result$community ##community of the graph

## use different community detection method
  #nlnet(input,gene.community.method="label.propagation")

## change the fdr pro to control connections of genes
## adjust the modularity size
#nlnet(input,conn.proportion=0.005,min.module.size=10)</pre>
```

8 nvsd

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Nonlinear Variable Selection based on DCOL

Description

This is a nonlinear variable selection procedure for generalized additive models. It's based on DCOL, using forward stagewise selection. In addition, a cross-validation is conducted to tune the stopping alpha level and finalize the variable selection.

Usage

```
nvsd(X, y, fold = 10, step.size = 0.01, stop.alpha = 0.05, stop.var.count = 20,
max.model.var.count = 10, roughening.method = "DCOL", do.plot = F, pred.method = "MARS")
```

Arguments

X	The predictor matrix. Each row is a gene (predictor), each column is a sample. Notice the dimensionality is different than most other packages, where each column is a predictor. This is to conform to other functions in this package that handles gene expression type of data.
у	The numerical outcome vector.
fold	The fold of cross-validation.
step.size	The step size of the roughening process.

stop.alpha The alpha level (significance of the current selected predictor) to stop the itera-

tions.

stop.var.count The maximum number of predictors to select in the forward stagewise selection.

Once this number is reached, the iteration stops.

max.model.var.count

The maximum number of predictors to select. Notice this can be smaller than the stop.var.count. Stop.var.count can be set more liniently, and this parameter controls the final maximum model size.

roughening.method

The method for roughening. The choices are "DCOL" or "spline".

do.plot Whether to plot the points change in each step.

pred.method The prediction method for the cross validation variable selection. As forward

stagewise procedure doesn't do prediction, a method has to be borrowed from

existing packages. The choices include "MARS", "RF", and "SVM".

Details

Please refer to the reference for details.

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Value

A list object is returned. The components include the following.

selected.pred The selected predictors (row number).

all.pred The selected predictors by the forward stagewise selection. The \$selected.pred

is a subset of this.

Author(s)

Tianwei Yu<tianwei.yu@emory.edu>

References

https://arxiv.org/abs/1601.05285

See Also

stage.forward

Examples

```
X<-matrix(rnorm(2000),ncol=20)
y<-sin(X[,1])+X[,2]^2+X[,3]
nvsd(t(X),y,stop.alpha=0.001,step.size=0.05)</pre>
```

stage.forward

Nonlinear Forward stagewise regression using DCOL

Description

The subroutine conducts forward stagewise regression using DCOL. Either DCOL roughening or spline roughening is conducted.

Usage

```
stage.forward(X, y, step.size = 0.01, stop.alpha = 0.01,
stop.var.count = 20, roughening.method = "DCOL", tol = 1e-08,
spline.df = 5, dcol.sel.only = FALSE, do.plot = F)
```

Arguments

Χ	The predictor matrix. Each row is a gene (predictor), each column is a sample.
	Notice the dimensionality is different than most other packages, where each
	column is a predictor. This is to conform to other functions in this package that
	handles gene expression type of data.

y The numerical outcome vector.

step.size The step size of the roughening process.

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stop.alpha The alpha level (significance of the current selected predictor) to stop the itera-

tions.

stop.var.count The maximum number of predictors to select. Once this number is reached, the

iteration stops.

roughening.method

The method for roughening. The choices are "DCOL" or "spline".

tol The tolerance level of sum of squared changes in the residuals.

spline.df The degree of freedom for the spline.

dcol.sel.only TRUE or FALSE. If FALSE, the selection of predictors will consider both linear

and nonlinear association significance.

do.plot Whether to plot the points change in each step.

Details

Please refer to the reference manuscript for details.

Value

A list object is returned. The components include the following.

found.pred The selected predictors (row number).

ssx.rec The magnitude of variance explained using the current predictor at each step.

\$sel.rec The selected predictor at each step.

\$p.rec The p-value of the association between the current residual and the selected

predictor at each step.

Author(s)

Tianwei Yu<tianwei.yu@emory.edu>

References

https://arxiv.org/abs/1601.05285

See Also

nvsd

```
X<-matrix(rnorm(2000),ncol=20)
y<-sin(X[,1])+X[,2]^2+X[,3]
stage.forward(t(X),y,stop.alpha=0.001,step.size=0.05)</pre>
```

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