Package 'CNA2Subpathway'

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Title Identification of Dysregulated Subpathway Driven by Copy Number Alteration

Type Package

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Description Identification of dysregulated subpathway driven by Copy Number Alterations(CNA). We first integrated CNA and gene expression data to calculate CNADEscore for each gene in cancer. Secondly, we construct a subpathway network based on the bical function relationship between subpathways, and the weight of this network regulated by the CNADEscore. Finally, we use an eigenvector centrality measure to determine the nificance of a subpathway, and the statistical significance of these subpathways is assessed using a bootstrap-based randomization method.	e sig-
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CNA2Subpathway-package

The CNA2Subpathway package

Description

CNA2Subpathway is a R package for based on Copy Number Alteration to identify dysregulated subpathway in cancer.

Details

CNA2Subpathway

The main goals of the CNA2Subpathway package is to provide main functions and some datas.

The main functions includes: 1) getCNADEscore 2) getCNAlabel 3) IDSubpath and two visual functions: 1) plotDiscovermap 2) plotSubPathway.

For more details, please see browseVignettes("CNA2Subpathway")

envData

The variables in the environment include an copy number variation data, amplified genes, deleted genes, an example expression profile, A matrix of subpathway infomation, Biological functions data, Jaccard coefficient of the network point, Overlapping genes of the network point, A list of subpathway map infomation, A binary matrix of CNA profile, A matric of CNADEscor, Optimized subpathway, and the statistical significance p value and FDR for these optimal subpathways

Description

Identification of dysregulated subpathways driven by Copy Number aberrations(CNA). We firstly calculate the CNADEscore by copy number aberration and gene expression. Secondly, we constructing subpathways network that implicitly links subpathways through their common function and weight of the network based on the CNADEscore. Finally, we calculate an eigenvector centrality and the statistical significance of these subpathways is assessed using a bootstrap-based randomization method.

Format

An environment variable

Details

The environment variable includes the variable ampGene, delGene,CNAlabelresult,GEP,CNAP,GoInfo,SubPathwayInfo

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getCNADEscore	getCNADEscore	
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Description

The function getCNADEscore used calculate the CNA-DEscore. The CNA-DEscore is defined as differentially expressed genes driven by CNA. We used the Student's t-test to calculate the t scores with the CNA label, and standardized them to a normal distribution with a mean of 0 and an SD of 1.

Usage

```
getCNADEscore(inexpData, CNALabel)
```

Arguments

inexpData Gene Expression profile(GEP)

CNALabel The CNA label (the result of function getCNAlabel)

Value

A matrix of the Student's t-test result and CNA-DEscore

Examples

```
# obtain GEP data
GEP<-GetExampleData("GEP")
# obtain the result of getCNAlabel function
CNAlabelresult<-GetExampleData("CNAlabelresult")
# Calculate the CNA-DEscore
CNADEscore<-getCNADEscore(GEP,CNAlabelresult)
# view first the ten gene CNA-DEscore
head(CNADEscore[1:10,])</pre>
```

getCNAlabel getCNAlabel

Description

The function getCNAlabel used to discretize the CNA profile to a binary matrix.

Usage

```
getCNAlabel(CNAprofile, ampGene, delGene)
```

Arguments

CNAprofile Copy Number Alteration profile

ampGene A vector of strings,the gene symbol of amplified genes delGene A vector of strings,the gene symbol of deleted genes

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Value

A binary matrix

Examples

```
# obtain Copy Number Alteration profile data
CNAP<-GetExampleData("CNAP")
head(CNAP[1:10,1:10])
# obtion amplified genes
ampGene<-GetExampleData("ampGene")
# obtion deleted genes
delGene<-GetExampleData("delGene")
# Obtin the CNA label.
CNAlabelresult<-getCNAlabel(CNAP,ampGene,delGene)
# get the result of the getCNAlabel function
CNAlabelresult<-GetExampleData("CNAlabelresult")
# view first ten genes and ten sample of the label
head(CNAlabelresult[1:10,1:10])</pre>
```

GetExampleData

GetExampleData

Description

Get the example data of test package for litte trials

Usage

GetExampleData(exampleData)

Arguments

exampleData

A character, should be one of "ampGene", "delGene", "GEP", "CNAP", "SubPathwayInfo", "GoSubPconGene", "Jaccardscore", "SubPathwaymapdata", "CNAlabelresult", "CNAlabelresult",

"CNA2Subpathwayresult".

Value

data

IDSubPath

IDSubPath

Description

The function IDSubPath used to find significant dysregulate subpathways by the IDSCNA method. According our method, in this function, the user need input 6 variable. Four of six variable can obtain from our example data, those data from KEGG and GO, and the user can change at will also. The "CNADEscore" variable is result of function getCNADEscore. And the last variable is nperm, that representative number of disturbances, usually nperm = 1000 or bigger as statistically significant.

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Usage

```
IDSubPath(CNADEscore, nperm, Subpathway, Go, Jaccard, Go_SubPath_gene)
```

Arguments

CNADEscore The CNA-DEscore of genes.

nperm input Number of disturbance(>=1000).

Subpathway Subpathway information from SuupathwayMiner.

Go Biological functions data from Gene Ontology.

Jaccard Jaccard score shared by a sub-pathway with biological function. Used to opti-

mization.

Go_SubPath_gene

Genes symble shared by a sub-pathway with biological function. Used to opti-

mization.

Value

Optimized subpathway, and the statistical significance p value and FDR for these optimal subpathways

Examples

```
# load depend package
require(igraph)
# obtain subpathway data
SubPathwayInfo<-GetExampleData("SubPathwayInfo")</pre>
# obtain biological functions data
GoInfo<-GetExampleData("GoInfo")</pre>
# obtain some genes shared by a sub-pathway with biological function
GoSubPconGene<-GetExampleData("GoSubPconGene")</pre>
# obtain Jaccardscore shared by a sub-pathway with biological function
Jaccardscore<-GetExampleData("Jaccardscore")</pre>
# obtain the result of getDEGscore function
CNADEscore<-GetExampleData("CNADEscore")</pre>
# run the IDSubPath function
#CNA2Subpathwayresult<-IDSubPath(CNADEscore,nperm=100,
                       Subpathway=SubPathwayInfo,Go=GoInfo,
                       Jaccard=Jaccardscore,Go_SubPath_gene=GoSubPconGene)
# get the result of the IDSCNA function #only show
{\tt CNA2Subpathwayresult<-GetExampleData("CNA2Subpathwayresult")}
# View the result top 10 subpathways
head(CNA2Subpathwayresult[1:10,])
```

plotDiscovermap plotDiscovermap

Description

we use Copy Number Alteration data to polt discover.

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Usage

```
plotDiscovermap(genes, CNAmatrix)
```

Arguments

A vector of strings. The user intresting and happened CNA genes symbol. genes CNAmatrix

The copy number alteration matrix. It takes the from of a binary matrix. Only 0

ang 1 are allowed as values.

Value

Groupwise test of the discover result

Examples

```
# load depend package
require(discover)
# input gene
genes <- c("AKT3","AKT1","CALML5","PIK3CA","PIK3R1","CALM1","CALM3","CALML3")</pre>
# get the result of the getCNAlabel function
CNAlabelresult<-GetExampleData("CNAlabelresult")</pre>
# plot discover
discover<-plotDiscovermap(genes,CNAlabelresult)</pre>
plot(discover)
```

plotSubPathway

plotSubPathway

Description

The function plotsubpathway can visualization of a subpathway network map.

Usage

```
plotSubPathway(
  subpathwayID,
  subpathwaydata,
  layout = NULL,
  margin = 0,
  vertex.label.cex = 0.8,
  vertex.label.font = 0.8,
  vertex.label.dist = 1,
  vertex.size = 12,
  edge.arrow.width = 3,
  edge.label.cex = 0.6,
  vertex.label.color = "black",
  vertex.color = "#F08080",
  vertex.frame.color = "dimgray",
  edge.color = "grey70",
  edge.label.color = "dimgray",
  sub = NULL,
  main = NULL
)
```

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Arguments

subpathwayID subpathway id number subpathwaydata The subpathwaymap data

layout A matrix of x-y coordinates with two dims. Determine the placement of the

nodes for drawing a graph.

margin A numeric. The value is usually between -0.5 and 0.5, which is able to zoom in

or out a subpathway graph. The default is 0.

vertex.label.cex

A numeric vector of node label size.

vertex.label.font

A numeric vector of label font.

vertex.label.dist

A numeric vector of label dist.

vertex.size A numeric vector of Node size. See plot.igraph.

edge.arrow.width

Edge arrow width. The default is 3.

edge.label.cex Edge label size.

vertex.label.color

A vector of node label colors. The default is black.

vertex.color A vector of node colors. The default is the KEGG node color.

vertex.frame.color

A vector of node frame color. The default is dimgray.

edge.color A vector of edge color. The default is dimgray

edge.label.color

A vector of edge label color. The default is dimgray.

sub A character string of subtitle.

main A character string of main title.

Value

a subpathway map

Examples

```
# load depend package
require(igraph)
# obtain subpathwaymap data
SubPathwaymapdata<-GetExampleData("SubPathwaymapdata")
# plot network graph of the subpathway 04915_13
plotSubPathway("04915_13",SubPathwaymapdata)</pre>
```

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