# Package 'CTD'

### November 10, 2019

<b>Title</b> CTD method for "connecting the dots" in weighted graphs
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<b>Description</b> An R package for pattern discovery in weighted graphs. Two use cases are achieved: 1) Given a weighted graph and a subset of its nodes; do the nodes show significant connectedness? 2) Given a weighted graph and two subsets of its nodes; do the subsets show significant similarity?
Depends R (>= 3.3.0), igraph, plotly, gplots, RColorBrewer
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### Description

A function that converts HMDB IDs to KEGG compounds.

### Usage

```
data.HMDBtoKEGG(hmdb.ids)
```

### **Arguments**

hmdb.ids - A character vector of HMDB IDs.

### Examples

```
kegg.ids = data.HMDBtoKEGG(hmdb.ids)
```

 ${\tt data.LabeltoKEGG}$ 

Convert compound names to KEGG compound IDs.

### Description

A function that converts HMDB IDs to KEGG compounds.

### Usage

```
data.LabeltoKEGG(compound.names)
```

### Arguments

compound.names - A character vector of metabolite (compound) names.

```
kegg.ids = data.HMDBtoKEGG(compound.names)
```

data.surrogateProfiles 3

```
data.surrogateProfiles
```

Surrogate profiles

#### Description

Fill in a data matrix rank, when your data is low n, high p. Fill in rank with surrogate profiles.

#### Usage

```
data.surrogateProfiles(data, sd = 1, useMnDiseaseProfile = FALSE,
  addHealthyControls = TRUE, ref_data = NULL)
```

#### **Arguments**

data

- Data matrix with observations as rows, features as columns.

sd

- The level of variability (standard deviation) around each feature's mean you want to add in surrogate profiles.

useMnDiseaseProfile

- Boolean. For disease cohorts not showing homogeneity, mean across disease profiles and generate disease surrogates around this mean.

addHealthyControls

- Boolean. Add healthy control profiles to data?

#### Value

data\_mx - Data matrix with added surrogate profiles.

graph.diffuseP1

Diffuse Probability P1 from a starting node.

#### **Description**

Recursively diffuse probability from a starting node based on the connectivity of the background knowledge graph, representing the likelihood that a variable will be most influenced by a perturbation in the starting node.

```
graph.diffuseP1(p1, startNode, G, visitedNodes, graphNumber = 1,
  verbose = FALSE)
```

The probability being dispersed from the starting node, startNode.
 The first variable drawn in the node ranking, from which p1 gets dispersed.
 A list of probabilities, with names of the list being the node names in the background knowledge graph.
 The history of previous draws in the node ranking sequence.
 If testing against multiple background knowledge graphs, this is the index associated with the adjacency matrix that codes for G. Default value is 1.
 If debugging or tracking a diffusion event, verbose=TRUE will activate print statements. Default is FALSE.

#### Value

G - A list of returned probabilities after the diffusion of probability has truncated, with names of the list being the node names in the background knowledge graph.

#### **Examples**

```
# Read in any network via its adjacency matrix
tmp=matrix(1, nrow=100, ncol=100)
for (i in 1:100) {
  for (j in 1:100) {
    tmp[i, j]=rnorm(1, mean=0, sd=1)
}
colnames(tmp)=sprintf("MolPheno%d", 1:100)
ig=graph.adjacency(tmp, mode="undirected", weighted=TRUE, add.colnames="name")
V(ig)$name=tolower(V(ig)$name)
adjacency_matrix=list(as.matrix(get.adjacency(ig, attr="weight"))) # Must have this declared as a GLOBAL var
# Set other tuning parameters
p0=0.1 # 10% of probability distributed uniformly
p1=0.9 # 90% of probability diffused based on edge weights in networks
thresholdDiff=0.01
G=vector(mode="list", length=length(V(ig)$name))
names(G)=V(ig)$name
G=lapply(G, function(i) i[[1]]=0)
startNode=names(G)[1]
visitedNodes=G[1]
probs_afterCurrDraw=graph.diffuseP1(p1, startNode, G, visitedNodes, 1, TRUE)
```

#### Description

Recursively diffuse probability from a starting node based on the connectivity of the background knowledge graph, representing the likelihood that a variable will be most influenced by a perturbation in the starting node.

graph.diffuseP1Movie 5

#### Usage

```
graph.diffuseP1Movie(p1, startNode, G, visitedNodes, ig,
  recursion_level = 1, output_dir = getwd())
```

#### Arguments

The probability being dispersed from the starting node, startNode.
 The first variable drawn in the node ranking, from which p1 gets dispersed.
 A list of probabilities, with names of the list being the node names in the background knowledge graph.
 A character vector of node names, storing the history of previous draws in the node ranking.
 If testing against multiple background knowledge graphs, this is the index associated with the adjacency matrix that codes for G. Default value is 1.

#### Value

G - A list of returned probabilities after the diffusion of probability has truncated, with names of the list being the node names in the background knowledge graph.

```
# 7 node example graph illustrating diffusion of probability based on network connectivity
# from Thistlethwaite et al., 2019.
adj_mat = rbind(c(0,2,1,0,0,0,0), # A
                c(2,0,1,0,0,0,0), # B
                c(1,0,0,1,0,0,0), # C
                c(0,0,1,0,2,0,0), # D
                c(0,0,0,2,0,2,1), # E
                c(0,0,0,1,2,0,1), # F
                c(0,0,0,0,1,1,0) # G
rownames(adj_mat) = c("A", "B", "C", "D", "E", "F", "G")
colnames(adj_mat) = c("A", "B", "C", "D", "E", "F", "G")
adjacency_matrix = list(adj_mat)
ig = graph.adjacency(as.matrix(adj\_mat), \ mode="undirected", \ weighted=TRUE) \\
G=vector(mode="list", length=7)
G[1:length(G)] = 0
names(G) = c("A", "B", "C", "D", "E", "F", "G")
startNode = "A"
visitedNodes = startNode
# Diffuse 100% of probability from startNode "A"
p1 = 1.0
# Probability diffusion truncates at
thresholdDiff=0.01
coords = layout.fruchterman.reingold(ig)
V(ig)$x = coords[,1]
V(ig)$y = coords[,2]
# Global variable imgNum
imgNum=1
G_new = graph.diffuseP1Movie(p1, startNode, G, visitedNodes, ig, 1, getwd())
```

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graph.naivePruning	Pruning edges from disease differential network that also occur in reference-only network.

#### **Description**

Recursively diffuse probability from a starting node based on the connectivity of the background knowledge graph, representing the likelihood that a variable will be most influenced by a perturbation in the starting node.

### Usage

```
graph.naivePruning(ig_dis, ig_ref)
```

ig\_pruned=graph.naivePruning(ig\_dis, ig\_ref)

#### **Arguments**

ig_dis	- The igraph object associated with the disease+reference trained differential interaction network.
ig_ref	- The igraph object associated with the reference-only trained interaction network

#### Value

ig\_pruned - The pruned igraph object of the disease+reference differential interaction network, with reference edges subtracted.

### **Examples**

```
mle.blowoutSim

Module that best explains the patient similarity assigned between a set of patients.
```

#### **Description**

Module that best explains the patient similarity assigned between a set of patients.

### Usage

```
mle.blowoutSim(patientSim, data_mx, ptIDs, ig_pruned, kmx)
```

### Arguments

patientSim	- A similarity matrix, where row and columns are patient identifiers.
data_mx	- The matrix that gives the perturbation strength (z-scores) for all variables (columns) for each patient (rows).
ptIDs	- The identifier associated with patient 1's sample.
ig_pruned	- The igraph object associated with the pruned disease+reference differential interaction network.
kmx	- The maximum metabolite set size probed when assessing patient similarity.

#### Value

ptsim\_blowout - An igraph object showing the module blowout describing the similarity between patients in ptIDs.

#### **Examples**

```
require(CTD)
data(Thistlethwaite2019)
data_mx = as.matrix(data_mx)
data_mx = suppressWarnings(apply(data_mx, c(1,2), as.numeric))
data_mx = data_mx[,-c(1,2,3,4,5,6,7,8)]
# Load your background network, ig_pruned and your computed patientSim matrix
kmns.clust = kmeans(patientSim, centers=4)
table(kmns.clust$cluster)
ptIDs = names(kmns.clust$cluster[which(kmns.clust$cluster==1)])
ptsim_blowout = mle.blowoutSim(patientSim, data_mx, ptIDs, ig_pruned, kmx=15)
plot.igraph(ptsim_blowout, layout=layout.circle, edge.width=50*abs(E(ptsim_blowout)$weight))
```

mle.getEncodingLength Minimum encoding length (MLE)

#### **Description**

This function calculates the minimum encoding length associated with a subset of variables given a background knowledge graph.

#### Usage

```
mle.getEncodingLength(bs, pvals, ptID, G)
```

#### **Arguments**

bs	- A list of bitstrings associated with a given patient's perturbed variables.
pvals	- The matrix that gives the perturbation strength significance for all variables (columns) for each patient (rows)
ptID	- The row name in data.pvals corresponding to the patient you specifically want encoding information for.
G	- A list of probabilities with list names being the node names of the background graph.

### Value

df - a data.frame object, for every bitstring provided in bs input parameter, a row is returned with the following data: the patientID; the bitstring evaluated where T denotes a hit and 0 denotes a miss; the subsetSize, or the number of hits in the bitstring; the individual p-values associated with the variable's perturbations, delimited by '/'; the combined p-value of all variables in the set using Fisher's method; Shannon's entropy, IS.null; the minimum encoding length IS.alt; and IS.null-IS.alt, the d.score.

mle.getMinPtDistance

#### **Examples**

```
# Identify the most significant subset per patient, given the background graph
data_mx.pvals = t(apply(data_mx, c(1,2), function(i) 2*pnorm(abs(i), lower.tail = FALSE)))
for (pt in 1:ncol(data_mx)) {
   ptID = colnames(data_mx)[pt]
   res = mle.getEncodingLength(ptBSbyK[[ptID]], data_mx.pvals, ptID, G)
   res = res[order(res[,"d.score"], decreasing=TRUE),]
   print(res)
}
```

mle.getMinPtDistance Metabolite set enrichment analysis (MSEA) using pathway knowledge curated by Metabolon

### **Description**

A function that returns the pathway enrichment score for all perturbed metabolites in a patient's full metabolomic profile.

### Usage

```
mle.getMinPtDistance(allSimMatrices)
```

#### **Arguments**

allSimMatrices - A list of all similarity matrices, across all k for a given graph, or across many graphs.

```
# Get patient distances
data_mx.pvals = apply(data_mx, c(1,2), function(i) 2*pnorm(abs(i), lower.tail = FALSE))
res = list()
t = list(ncd=matrix(NA, nrow=ncol(data_mx), ncol=ncol(data_mx)))
rownames(t$ncd) = colnames(data_mx)
colnames(t$ncd) = colnames(data_mx)
for (i in 1:kmx) {
  res[[i]] = t
for (pt in 1:ncol(data_mx)) {
  print(pt)
  ptID = colnames(data_mx)[pt]
  for (pt2 in pt:ncol(data_mx)) {
    ptID2 = colnames(data_mx)[pt2]
    tmp = mle.getPtSim(ptBSbyK[[ptID]], ptID, ptBSbyK[[ptID2]], ptID2, data_mx, ranks)
    for (k in 1:kmx) {
      res[[k]]$ncd[ptID, ptID2] = tmp$NCD[k]
      res[[k]]$ncd[ptID2, ptID] = tmp$NCD[k]
    }
  }
patientSimilarity = mle.getMinPtDistance(res)
```

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mle.getPtSim	Patient similarity using mutual information MLE metric of patients' most modular, perturbed subsets.
--------------	--

#### **Description**

This function calculates the universal distance between patients, using a mutual information metric, where self-information comes from the minimum encoding length of each patient's encoded modular perturbations in the background knowledge graph.

### Usage

```
mle.getPtSim(p1.optBS, ptID, p2.optBS, ptID2, data_mx, ranks)
```

#### **Arguments**

```
p1.optBS - The optimal bitstring associated with patient 1.

ptID - The identifier associated with patient 1's sample.

p2.optBS - The optimal bitstring associated with patient 2.

data_mx - The matrix that gives the perturbation strength (z-scores) for all variables (columns) for each patient (rows).

ptID - The identifier associated with patient 2's sample.
```

#### Value

patientSim - a similarity matrix, where row and columns are patient identifiers.

```
# Get patient distances
data_mx.pvals = apply(data_mx, c(1,2), function(i) 2*pnorm(abs(i), lower.tail = FALSE))
res = list()
tt = list(ncd=matrix(NA, nrow=ncol(data_mx), ncol=ncol(data_mx)))
rownames(t$ncd) = colnames(data_mx)
colnames(t$ncd) = colnames(data_mx)
for (i in 1:kmx) {
  res[[i]] = tt
for (pt in 1:ncol(data_mx)) {
  print(pt)
  ptID = colnames(data_mx)[pt]
  for (pt2 in pt:ncol(data_mx)) {
    ptID2 = colnames(data_mx)[pt2]
    for (k in 1:kmx) {
    tmp = mle.getPtSim(ptBSbyK[[ptID]][k], ptID, ptBSbyK[[ptID2]][k], ptID2, data_mx, ranks)
      res[[k]]$ncd[ptID, ptID2] = tmp$NCD
      res[[k]]$ncd[ptID2, ptID] = tmp$NCD
    }
 }
}
```

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mle.kraftMcMillan	Apply the Kraft-McMillian Inequality using a specific encoding algorithm.
-------------------	---

#### **Description**

A power analysis of the encoding algorithm using to encode subsets of S in G.

#### Usage

```
mle.kraftMcMillan(G, k, multiNode = FALSE)
```

#### **Arguments**

G - A character vector of all node names in the background knowledge graph.

- The size of the node name subsets of G.

multiNode - Boolean, indicating whether to use the multi-node diffusion encoding algo-

rithm (TRUE) or the single-node diffusion encoding algorithm (FALSE). De-

fault is FALSE.

#### Value

IA - a list of bitlengths associated with all outcomes in the N choose K outcome space, with the names of the list elements the node names of the encoded nodes

multiNode.getNodeRanks

Generate the "adaptive walk" node rankings, starting from a given perturbed variable

#### **Description**

This function calculates the node rankings starting from a given perturbed variable in a subset of variables in the background knowledge graph.

### Usage

```
multiNode.getNodeRanks(S, G, num.misses = NULL)
```

#### **Arguments**

S

- A character vector of the node names for the subset of nodes you want to encode.

#### Value

ranks - A list of character vectors of node names in the order they were drawn by the probability diffusion algorithm, from each starting node in S.

### **Examples**

```
# Read in any network via its adjacency matrix
tmp = matrix(1, nrow=100, ncol=100)
for (i in 1:100) {
    for (j in 1:100) {
        tmp[i, j] = rnorm(1, mean=0, sd=1)
      }
}
colnames(tmp) = sprintf("Compound%d", 1:100)
ig = graph.adjacency(tmp, mode="undirected", weighted=TRUE, add.colnames="name")
V(ig)$name = tolower(V(ig)$name)
# Get node rankings for graph
G = vector(mode="list", length=length(V(ig)$name))
names(G) = V(ig)$name
S = names(G)[1:3]
ranks = multiNode.getNodeRanks(S, G)
```

multiNode.getNodeRanksMovie

Capture the movement of the adaptive walk of the diffusion probability method.

#### **Description**

Make a movie of the adaptive walk the diffusion probability method makes in search of a given patient's perturbed variables.

#### Usage

```
multiNode.getNodeRanksMovie(subset.nodes, ig, output_filepath,
  movie = TRUE, zoomIn = FALSE)
```

#### **Arguments**

 $\begin{tabular}{ll} subset.nodes & - The subset of variables, S, in a background graph, G. \\ ig & - The igraph object associated with the background knowledge graph. \\ output\_filepath & - The igraph object associated with the background knowledge graph. \\ output\_filepath & - The igraph object associated with the background knowledge graph. \\ \hline \end{tabular}$ 

- The local directory at which you want still images to be saved.

movie - If you want to make a movie, set to TRUE. This will produce a set of still

images that you can stream together to make a movie. Default is TRUE. Alternatively (movie=FALSE), you could use this function to get the node labels

returned for each node ranking starting with a perturbed variable.

- Boolean. Delete nodes outside of node subset's order 1 neighborhood?. De-

fault is FALSE.

#### Value

zoomIn

ranksByStartNode - a list object of node rankings Each element is based on a different startNode. Images are also generated in the output\_directory specified.

#### **Examples**

```
# Read in any network via its adjacency matrix
tmp = matrix(1, nrow=100, ncol=100)
for (i in 1:100) {
  for (j in 1:100) {
    tmp[i, j] = rnorm(1, mean=0, sd=1)
}
colnames(tmp) = sprintf("Compound%d", 1:100)
ig = graph.adjacency(tmp, mode="undirected", weighted=TRUE, add.colnames="name")
V(ig)$name = tolower(V(ig)$name)
# Set other tuning parameters
p0=0.1 # 10% of probability distributed uniformly
p1=0.9 # 90% of probability diffused based on edge weights in networks
thresholdDiff=0.01
G = vector(mode="list", length=length(V(ig)$name))
names(G) = V(ig) name
subset.nodes = names(G)[sample(1:length(G), 3)]
multiNode.getNodeRanksMovie(subset.nodes, ig, output_filepath = getwd(), movie=TRUE)
```

multiNode.getPtBSbyK Generate patient-specific bitstrings from adaptive network walk.

### **Description**

This function calculates the bitstrings (1 is a hit; 0 is a miss) associated with the adaptive network walk made by the diffusion algorithm trying to find the variables in the encoded subset, given the background knowledge graph.

#### Usage

```
multiNode.getPtBSbyK(S, ranks)
```

#### **Arguments**

S

- A character vector of node names describing the node subset to be encoded.

ranks

- The list of node ranks calculated over all possible nodes, starting with each node in subset of interest.

#### Value

pt.byK - a list of bitstrings, with the names of the list elements the node names of the encoded nodes

#### **Examples**

```
# Get bitstrings associated with each patient's top kmx variable subsets
kmx = 15
ptBSbyK = list()
for (pt in 1:ncol(data_mx)) {
   S = data_mx[order(abs(data_mx[,pt]), decreasing=TRUE),pt][1:kmx]
   ptBSbyK[[ptID]] = multiNode.getPtBSbyK(S, ranks)
}
```

```
pathway.ListMaps_metabolon
```

Get All Metabolites In Metabolon's Pathway Knowledgebase

### Description

Get All Metabolites In Metabolon's Pathway Knowledgebase

#### Usage

```
pathway.ListMaps_metabolon()
```

#### Value

pwys - List of pathway maps curated by Metabolon's Metabolync.

```
pwys = pathway.ListMaps_metabolon()
print(pwys)
```

pathway.ListMetabolites\_metabolon

Get All Metabolites In Metabolon's Pathway Knowledgebase

### Description

Get All Metabolites In Metabolon's Pathway Knowledgebase

#### Usage

```
pathway.ListMetabolites_metabolon()
```

#### Value

mets - a character vector of unique metabolites and enzymes found in at least 1 pathway in Metabolon's pathway knowledgebase.

### **Examples**

```
mets = pathway.ListMetabolites_metabolon()
print(mets)
```

 $\verb"plot.getPathwayIgraph" plot.getPathwayIgraph"$ 

#### **Description**

plot.getPathwayIgraph

#### Usage

```
plot.getPathwayIgraph(input, Pathway.Name)
```

#### **Arguments**

input - A list object of parameters (esp. from R shiny app). Required parameters are

ptIDs, diagClass and pathwayMapId.

Pathway. Name - The name of the pathway map for which you want the topological information.

#### Value

template.ig - Igraph object of selected pathway map.

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#### **Examples**

```
data(Miller2015)
# Input is supplied by R shiny app, but you can hard code parameters as a list object, too, to test functional:
input = list()
input$ptIDs = colnames(Miller2015)[4]
input$diagClass = "paa"
input$pathwayMapId = "All"
ig = plot.getPathwayIgraph(input, Miller2015)
# Returns a blank template for selected pathway.
plot.igraph(ig, edge.arrow.size = 0.01)
```

plot.hmSim

Generate heatmap plot of patient similarity matrix.

#### **Description**

This function plots a heatmap of a patient similarity matrix.

#### Usage

```
plot.hmSim(patientSim, path=getwd(), diagnoses=NULL)
```

#### **Arguments**

patientSim

- The patient similarity matrix.

path

- The filepath to a directory in which you want to store the .png file.

diagnoses

- A character vector of diagnostic labels associated with the rownames of pa-

tientSim.

#### **Examples**

```
plot.hmSim(patientSim, path=getwd(), diagnoses)
```

plot.knnSim

Visualize the confusion matrix using nearest neighbor as classification

model.

### Description

Visualize the confusion matrix using nearest neighbor as classification model.

#### Usage

```
plot.knnSim(patientSim, diagnoses, diag)
```

#### **Arguments**

patientSim

- The patient similarity matrix.

diagnoses

- A character vector of diagnostic labels associated with the rownames of pa-

tientSim. The names of this vector are patient IDs, and values are diagnostic

labels.

diag

- The diagnosis associated with positive controls in your data.

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#### Value

p - a plotly scatter plot colored by provided diagnostic labels.

#### **Examples**

```
\# if you have diagnostic labels associated with the colnames(data_mx), send them using diagnoses parameter p = plot.knnSim(patientSim, diagnoses, diag="diseased") p
```

plot.mdsSim

View patient clusters using multi-dimensional scaling.

#### **Description**

This function plots the provided patient similarity matrix in a lower dimensional space using multidimensional scaling, which is well suited for similarity metrics.

### Usage

```
plot.mdsSim(patientSim, diagnoses, k, diag)
```

### Arguments

- The patient similarity matrix.

- A character vector of diagnostic labels associated with the rownames of patientSim.

- The number of dimension you want to plot your data using multi-dimensional scaling.

- The diagnosis associated with positive controls in your data.

#### Value

p - a plotly scatter plot colored by provided diagnostic labels.

```
# if you have diagnostic labels associated with the colnames(data_mx), send them using diagnoses parameter
p = plot.mdsSim(patientSim, diagnoses, k=2, diag="diseased")
p
p = plot.mdsSim(patientSim, diagnoses, k=3, diag="diseased")
p
```

plot.pathwayMap 17

plot.pathwayMap	Generate pathway map with patient perturbation data superimposed.
• • •	

#### **Description**

Generate pathway map with patient perturbation data superimposed.

### Usage

```
plot.pathwayMap(Pathway, ptID, pt.zscore, zscore.threshold, scale, out.path, SVG=TRUE)
```

#### **Arguments**

- The name of the pathway map you want to plot patient data on.

ptID - An identifier string associated with the patient.

pt.zscore - A named vector of metabolites with corresponding z-scores.

zscore.threshold

- Plot all z-scores > or < this threshold.

scale - Integer associated with increase in node size.

out.path - The directory in which you want to store image files.

SVG - Save as SVG or PNG? If SVG is TRUE, then an SVG image is saved. If

FALSE, a PNG is saved.

#### **Examples**

```
Pathway = pathway.ListMaps_metabolon()
data(Miller2015)
Miller2015 = Miller2015[,grep("IEM", colnames(Miller2015))]
ptID = colnames(Miller2015)[1]
pt.zscore = Miller2015[,1]
plot.pathwayMap(Pathway[1], ptID, pt.zscore, zscore.threshold, scale=1, out.path=getwd(), SVG=TRUE)
```

```
singleNode.getNodeRanksMovie
```

Capture the movement of the fixed, single-node walk of the diffusion probability method.

#### **Description**

Make a movie of the fixed, single-node walk the diffusion probability method makes in search of a given patient's perturbed variables.

```
singleNode.getNodeRanksMovie(subset.nodes, ig, output_filepath,
  movie = TRUE, zoomIn = FALSE)
```

subset.nodes

- The subset of variables, S, in a background graph, G.

- The igraph object associated with the background knowledge graph.

output\_filepath

- The local directory at which you want still images to be saved.

movie

- If you want to make a movie, set to TRUE. This will produce a set of still images that you can stream together to make a movie. Default is TRUE. Alternatively (movie=FALSE), you could use this function to get the node labels returned for each node ranking starting with a perturbed variable.

zoomIn

- Boolean. Delete nodes outside of node subset's order 1 neighborhood?. Default is FALSE.

#### Value

ranksByStartNode - a list object of node rankings Each element is based on a different startNode. Images are also generated in the output\_directory specified.

#### **Examples**

```
# Read in any network via its adjacency matrix
tmp = matrix(1, nrow=100, ncol=100)
for (i in 1:100) {
  for (j in 1:100) {
    tmp[i, j] = rnorm(1, mean=0, sd=1)
  }
}
colnames(tmp) = sprintf("Compound%d", 1:100)
ig = graph.adjacency(tmp, mode="undirected", weighted=TRUE, add.colnames="name")
V(ig)name = tolower(V(ig)name)
adjacency_matrix = list(tmp) # MUST BE GLOBAL VARIABLE
# Set other tuning parameters
p0=0.1 # 10% of probability distributed uniformly
p1=0.9 # 90% of probability diffused based on edge weights in networks
thresholdDiff=0.01
G = vector(mode="list", length=length(V(ig)$name))
names(G) = V(ig) name
subset.nodes = names(G)[sample(1:length(G), 3)]
singleNode.getNodeRanksMovie(subset.nodes, ig, output_filepath = getwd(), movie=TRUE)
```

singleNode.getNodeRanksN

Generate the fixed, single-node diffusion node rankings, starting from a given perturbed variable.

### **Description**

This function calculates the node rankings starting from a given perturbed variable in a subset of variables in the background knowledge graph.

```
singleNode.getNodeRanksN(n, G, S = NULL, misses.thresh = NULL)
```

n	- The index (out of a vector of node names) of the node ranking you want to calculate.
G	- A list of probabilities with list names being the node names of the background graph.
S	- A character vector of node names in the subset you want the network walker to find.
misses.thresh	- The number of "misses" the network walker will tolerate before switched to fixed length codes for remaining nodes to be found.

#### Value

current\_node\_set - A character vector of node names in the order they were drawn by the probability diffusion algorithm.

#### **Examples**

```
# Get node rankings for graph
ranks = list()
for (n in 1:length(G)) {
   print(sprintf("Generating node rankings starting with node %s", names(G)[n]))
   ranks[[n]] = singleNode.getNodeRanksN(n, G)
}
names(ranks) = names(G)
```

singleNode.getPtBSbyK Generate patient-specific bitstrings from the fixed, single-node network walker.

#### **Description**

This function calculates the bitstrings (1 is a hit; 0 is a miss) associated with the fixed, single-node network walker trying to find the variables in the encoded subset, given the background knowledge graph.

#### Usage

```
singleNode.getPtBSbyK(S, ranks, num.misses = NULL)
```

#### Arguments

S	- A character vector of node names describing the node subset to be encoded.
ranks	- The list of node ranks calculated over all possible nodes, starting with each node in subset of interest.
num.misses	- The number of misses tolerated by the network walker before path truncation occurs.

### Value

pt.byK - a list of bitstrings, with the names of the list elements the node names of the encoded nodes

20 stats.fishersMethod

#### **Examples**

```
# Load in your profiling data (rows are compounds, columns are samples)
data_mx = read.table("your_profiling_data.txt", sep="\t", header=TRUE)
# Get bitstrings associated with each patient's top kmx variable subsets
kmx = 15
ptBSbyK = list()
for (pt in 1:ncol(data_mx)) {
   S = data_mx[order(abs(data_mx[,pt]), decreasing=TRUE),pt][1:kmx]
   ptBSbyK[[ptID]] = singleNode.getPtBSbyK(S, ranks)
}
```

stats.entropyFunction Entropy of a bit-string

#### **Description**

The entropy of a bitstring (ex: 1010111000) is calculated.

### Usage

```
stats.entropyFunction(bitString)
```

#### **Arguments**

Χ

- A vector of 0's and 1's.

#### Value

e - a floating point percentage, between 0 and 1.

#### **Examples**

```
 \begin{array}{l} stats.entropyFunction(c(1,0,0,0,1,0,0,0,0,0,0,0,0,0)) \\ > 0.6193822 \\ stats.entropyFunction(c(1,1,1,1,1,1,1,1,0,0,0,0,0,0,0,0,0)) \\ > 1 \\ stats.entropyFunction(c(1,1,1,1,1,1,1,1,1,1,1,1,1,1,1,1)) \\ > 0 \\ \end{array}
```

stats.fishersMethod

Fisher's Combined P-value

### Description

Fisher's combined p-value, used to combine the results of individual statistical tests into an overall hypothesis.

```
stats.fishersMethod(x)
```

Х

- A vector of p-values (floating point numbers).

#### Value

a floating point number, a combined p-value using Fisher's method.

#### **Examples**

```
stats.fishersMethod(c(0.2,0.1,0.3)) > 0.1152162
```

```
stats.getMSEA_Metabolon
```

Metabolite set enrichment analysis (MSEA) using pathway knowledge curated by Metabolon

#### **Description**

A function that returns the pathway enrichment score for all perturbed metabolites in a patient's full metabolomic profile.

#### Usage

```
stats.getMSEA_Metabolon(abs_filename_dataset, abs_filename_classes,
  pathway_knowledgebase = "Metabolon", output_dir = getwd(),
  expt_name = "msea_results")
```

#### Arguments

```
abs_filename_dataset
```

- Relative or absolute path to relevant .gct file. A .gct file contains profiling data, rows are compounds and columns are sample IDs.

abs\_filename\_classes

- Relative or absolute path to relevant .cls file. A .cls file contains a mapping of class labels to columns in the .gct file.

output\_dir

- The path associated with the folder in which MSEA results will be saved.

expt\_name

- A name to be associated with the experiment you are analyzing. This name will be used in filestems of results rendered in output\_dir.

#### pathway.knowledgebase

- The filename of the .gmt file associated with the pathway knowledge desired. Currently only "Metabolon" is offered, though "KEGG", "WikiPathways", "SM-PDB" and/or "Reactome" can be added in future versions.

#### **Examples**

```
data(Miller2015)
Miller2015 = Miller2015[,grep("IEM", colnames(Miller2015))]
# Generate a .cls file for your data.
diagnoses = gsub("[[:digit:]]", "", colnames(Miller2015))
diag.ind = diagnoses
diag.ind[which(diag.ind!="Argininemia")] = 0
diag.ind[which(diag.ind=="Argininemia")] = 1
diag.ind = as.numeric(diag.ind)
# Manually add the following text to 1st line of .cls,
  where num_samples is the length of diag.ind: #num_samples 1 2
# Manually add the following text to 2nd line of .cls: #disease control
write.table(diag.ind, file=system.file("extdata/MSEA_Datasets/Miller2015_arg.cls", package="CTD"),
            sep=" ", quote=FALSE, row.names = FALSE, col.names = FALSE)
# Create a .gct file.
data_mx = Miller2015
data_mx = data_mx[, order(diags.ind)]
data_mx = cbind(rep(NA, nrow(data_mx)), data_mx)
colnames(data_mx)[1] = "DESCRIPTION"
write.table(data_mx, file=system.file("extdata/MSEA_Datasets/Miller2015.gct", package="CTD"),
            sep="\t", quote=FALSE, row.names = TRUE)
# Generate a .gmt file.
population = names(met.profile)
paths.hsa = list.dirs(path="../inst/extdata", full.names = FALSE)
paths.hsa = paths.hsa[-which(paths.hsa %in% c("", "RData", "allPathways"))]
sink(system.file("extdata/Pathway_GMTs/Metabolon.gmt", package="CTD"))
for (p in 1:length(paths.hsa)) {
  load(sprintf("../inst/extdata/RData/%s.RData", paths.hsa[p]))
  pathway.compounds = V(ig)$label[which(V(ig)$shape=="circle")]
 pathCompIDs = unique(tolower(pathway.compounds[which(pathway.compounds %in% population)]))
                         %s", paths.hsa[p], paste(pathCompIDs, collapse="
                                                                             ")), quote=FALSE)
 print(sprintf("%s
}
sink()
print("test")
abs_filename_dataset = system.file("extdata/MSEA_Datasets/Miller2015.gct", package="CTD")
abs_filename_classes = system.file("extdata/MSEA_Datasets/Miller2015_arg.cls", package="CTD")
pathway.data = stats.getMSEA_Metabolon(abs_filename_dataset, abs_filename_classes, pathway_knowledgebase = '
                                       output_dir = getwd(), expt_name="msea_results")
```

stats.getORA\_Metabolon

Metabolite set enrichment analysis (MSEA) (using a hypergeometric test) using pathway knowledge curated by Metabolon

#### **Description**

A function that returns the pathway enrichment score for all perturbed metabolites in a patient's full metabolomic profile.

```
stats.getORA_Metabolon(met.profile, threshold = 3, type = "zscore",
   gene.profile = NULL)
```

met.profile - A character vector of a patient's metabolomic profile, including KEGG IDs and

the associated z-score or p-value describing the level of the metabolite compared

to controls.

threshold - A cutoff to select metabolites with a zscore > threshold or < -1\*threshold.

type - Either "p-value" or "z-score".

gene.profile - Default set to NULL, meaning the default enrichment analysis only considers

metabolites. However, if you have gene data, too, set this parameter to a character vector of the gene names with found variants in the patient's record. Gene

IDs must be converted to Entrez Identifiers.

### **Examples**

pathway.data = stats.getORA\_Metabolon(met.profile, threhold=3, "z-score", NULL)

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```