

# Package ‘CTD’

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**Title** CTD method for “connecting the dots” in weighted graphs

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**Description** 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

**License** MIT License

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**LazyData** true

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data.HMDBtoKEGG	<i>Convert HMDB IDs to KEGG compound IDs.</i>
-----------------	---

---

### 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)
```

---

data.LabeltoKEGG	<i>Convert compound names to KEGG compound IDs.</i>
------------------	---

---

### 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.

### Examples

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

---

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.

### Usage

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

**Arguments**

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

```
# Look at main_CTD.r script for full analysis script: https://github.com/BRL-BCM/CTD.
# 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 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
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)
```

---

graph.diffuseP1Movie    *Make a movie of the diffusion of 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.

**Usage**

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

**Arguments**

p1	- The probability being dispersed from the starting node, startNode.
startNode	- The first variable drawn in the adaptive permutation node sequence, from which p1 gets dispersed.
G	- A list of probabilities, with names of the list being the node names in the background knowledge graph.
visitedNodes	- A character vector of node names, storing the history of previous draws in the permutation sequence.
graphNumber	- 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.

**Examples**

```
# 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())
```

---

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.

## 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	<i>Metabolite set enrichment analysis (MSEA) using pathway knowledge curated by Metabolon</i>
----------------------	---

---

## 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.

## Examples

```
# 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, perms)
    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)
```

---

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

---

## 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, perms)
```

**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).  
 ptID2 - The identifier associated with patient 2's sample.

**Value**

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

**Examples**

```
# 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, perms)
      res[[k]]$ncd[ptID, ptID2] = tmp$NCD
      res[[k]]$ncd[ptID2, ptID] = tmp$NCD
    }
  }
}
```

---

mle.kraftMcMillan

*Apply the Kraft-McMillan 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, memory = FALSE)
```



**Arguments**

- G - A character vector of all node names in the background knowledge graph.
- k - The size of the node name subsets of G.

**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

**Examples**

```
G = list(A=0, B=0, C=0, D=0, E=0, F=0, G=0)
names(G) = tolower(names(G))
adj_mat = rbind(c(0,2,1,0,0,0,0), #A's neighbors
               c(2,0,1,0,0,0,0), #B's neighbors
               c(1,1,0,1,0,0,0), #C's neighbors
               c(0,0,1,0,2,1,0), #D's neighbors
               c(0,0,0,2,0,2,1), #E's neighbors
               c(0,0,0,1,2,0,1), #F's neighbors
               c(0,0,0,0,1,1,0)  #G's neighbors
               )
rownames(adj_mat) = names(G)
colnames(adj_mat) = names(G)
adjacency_matrix = list(adj_mat)
IA = mle.kraftMcMillian(G, 2)
# Power to find effects is
sum(2^-unlist(IA))
```

---

multiNode.getPermMovie

*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.getPermMovie(subset.nodes, ig, output_filepath, movie = TRUE,
                        zoomIn = FALSE)
```

**Arguments**

- 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 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 permutation starting with a perturbed variable.
zoomIn	- Boolean. Delete nodes outside of node subset's order 1 neighborhood?. Default is FALSE.

### Value

permutationByStartNode - a list object of node permutations. 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.getPermMovie(subset.nodes, ig, output_filepath = getwd(), movie=TRUE)
```

---

multiNode.getPerms	<i>Generate the "adaptive walk" node permutations, starting from a given perturbed variable</i>
--------------------	---

---

### Description

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

### Usage

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

### Arguments

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

**Value**

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

**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 permutations for graph
G = vector(mode="list", length=length(V(ig)$name))
names(G) = V(ig)$name
S = names(G)[1:3]
perms = multiNode.getPerms(S, G)
```

---

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, perms)
```

**Arguments**

S                      - A character vector of node names describing the node subset to be encoded.  
perms                  - The list of permutations 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, perms)
}
```

```
pathway.ListMaps_metabolon
```

*Get All Metabolites In Metabolon's Pathway Knowledgebase*

---

### Description

Get All Metabolites In Metabolon's Pathway Knowledgebase

### Usage

```
pathway.ListMaps_metabolon(Pathway.Knowledgebase.Path)
```

### Value

Pathway.Knowledgebase.Path - local path to extdata folder containing metabolon's pathway knowledgebase files.

### Examples

```
metabolon_knowledgebase_path = sprintf("%s/extdata", find.package("CTD"))
pwys = pathway.ListMaps_metabolon(metabolon_knowledgebase_path)
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)
```

---

plot.getPathwayIgraph *plot.getPathwayIgraph*


---

**Description**

plot.getPathwayIgraph

**Usage**

plot.getPathwayIgraph(input, Pathway.Name, pmap.path = "../extdata")

**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.

**Examples**

```
data(Miller2015)
# Input is supplied by R shiny app, but you can hard code parameters as a list object, too, to test functionality
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 patientSim.

**Examples**

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

---

plot.knnSim	<i>Visualize the confusion matrix using nearest neighbor as classification model.</i>
-------------	---

---

**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 patientSim. The names of this vector are patient IDs, and values are diagnostic labels.
diag	- The diagnosis associated with positive controls in your data.

**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	<i>View patient clusters using multi-dimensional scaling.</i>
-------------	---

---

**Description**

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

**Usage**

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

**Arguments**

- patientSim      - The patient similarity matrix.
- diagnoses      - A character vector of diagnostic labels associated with the rownames of patientSim.
- k                - The number of dimension you want to plot your data using multi-dimensional scaling.
- diag            - The diagnosis associated with positive controls in your data.

**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.mdsSim(patientSim, diagnoses, k=2, diag="diseased")
p
p = plot.mdsSim(patientSim, diagnoses, k=3, diag="diseased")
p
```

---

plot.pathwayMap	<i>Generate pathway map with patient perturbation data superimposed.</i>
-----------------	--

---

**Description**

Generate pathway map with patient perturbation data superimposed.

**Usage**

```
plot.pathwayMap(Pathway.Name, PatientID, patient.zscore, zscore.threshold, scalingFactor, outputFile)
```

**Arguments**

- Pathway.Name    - The name of the pathway map you want to plot patient data on.
- PatientID        - An identifier string associated with the patient.
- patient.zscore   - A named vector of metabolites with corresponding z-scores.
- zscore.threshold      - Plot all z-scores > or < this threshold.
- scalingFactor    - Integer associated with increase in node size.
- outputFilePath   - 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.Name = pathway.ListMaps_metabolon()
data(Miller2015)
Miller2015 = Miller2015[,grep("IEM", colnames(Miller2015))]
PatientID = colnames(Miller2015)[1]
patient.zscore = Miller2015[,1]
plot.pathwayMap(Pathway.Name[1], PatientID, patient.zscore, zscore.threshold, scalingFactor=1, outputFile)
```

---

singleNode.getPermMovie

*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

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

## Arguments

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 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 permutation starting with a perturbed variable.
zoomIn	- Boolean. Delete nodes outside of node subset's order 1 neighborhood?. Default is FALSE.

## Value

permutationByStartNode - a list object of node permutations. 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.getPermMovie(subset.nodes, ig, output_filepath = getwd(), movie=TRUE)
```

---

singleNode.getPermN	<i>Generate the "adaptive walk" node permutations, starting from a given perturbed variable</i>
---------------------	---

---

## Description

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

## Usage

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

## Arguments

n	- The index (out of a vector of node names) of the permutation 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 permutations for graph
perms = list()
for (n in 1:length(G)) {
  print(sprintf("Generating node permutation starting with node %s", names(G)[n]))
  perms[[n]] = singleNode.getPermN(n, G)
}
names(perms) = names(G)
```

---

`singleNode.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 memoryless network walker trying to find the variables in the encoded subset, given the background knowledge graph.

### Usage

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

### Arguments

<code>S</code>	- A character vector of node names describing the node subset to be encoded.
<code>perms</code>	- The list of permutations calculated over all possible nodes, starting with each node in subset of interest.
<code>num.misses</code>	- 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

### 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, perms)
}
```

---

`stats.entropyFunction` *Entropy of a bit-string*

---

### Description

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

### Usage

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

### Arguments

<code>x</code>	- A vector of 0's and 1's.
----------------	----------------------------

**Value**

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

**Examples**

```
stats.entropyFunction(c(1,0,0,0,1,0,0,0,0,0,0,0))
> 0.6193822
stats.entropyFunction(c(1,1,1,1,1,1,1,0,0,0,0,0))
> 1
stats.entropyFunction(c(1,1,1,1,1,1,1,1,1,1,1,1))
> 0
```

---

stats.fishersMethod	<i>Fisher's Combined P-value</i>
---------------------	----------------------------------

---

**Description**

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

**Usage**

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

**Arguments**

x - 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	<i>Metabolite set enrichment analysis (MSEA) using pathway knowledge curated by Metabolon</i>
-------------------------	---

---

**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_sampl
# Manually add the following text to 2nd line of .cls: #disease control
write.table(diag.ind, file="MSEA_Datasets/Miller2015_arg.cls", sep=" ", quote=FALSE, row.names = FALSE, col.

# 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="MSEA_Datasets/Miller2015.gct", 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("Pathway_GMTs/Metabolon.gmt")
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)]))
  print(sprintf("%s\t%s", paths.hsa[p], paste(pathCompIDs, collapse=" ")), quote=FALSE)
}
sink()
print("test")
abs_filename_dataset = "MSEA_Datasets/Miller2015.gct"
abs_filename_classes = "MSEA_Datasets/Miller2015_arg.cls"
pathway.data = stats.getMSEA_Metabolon(abs_filename_dataset, abs_filename_classes, pathway_knowledgebase = "
```

---

`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.

### Usage

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

### Arguments

<code>met.profile</code>	- 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.
<code>threshold</code>	- A cutoff to select metabolites with a zscore > threshold or < -1*threshold.
<code>type</code>	- Either "p-value" or "z-score".
<code>gene.profile</code>	- 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, threshold=3, "z-score", NULL)
```

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