# dRWRcontact

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dRWRcontact Function to estimate RWR-based contact strength between samples

from an input gene-sample data matrix, an input graph and its pre-

computed affinity matrix

# **Description**

dRWRcontact is supposed to estimate sample relationships (ie. contact strength between samples) from an input gene-sample matrix, an input graph and its affinity matrix pre-computed according to random walk restart (RWR) of the input graph. It includes: 1) RWR-smoothed columns of input gene-sample matrix based on the pre-computed affinity matrix; 2) calculation of contact strength (inner products of RWR-smooth columns of input gene-sample matrix); 3) estimation of the contact signficance by a randomalisation procedure. Parallel computing is also supported for Linux or Mac operating systems.

### Usage

```
dRWRcontact(data, g, Amatrix, permutation = c("random", "degree"),
num.permutation = 10, p.adjust.method = c("BH", "BY", "bonferroni",
"holm", "hochberg", "hommel"), adjp.cutoff = 0.05, parallel = TRUE,
multicores = NULL, verbose = T)
```

## **Arguments**

data an input gene-sample data matrix used for seeds. Each value in input gene-

sample matrix does not necessarily have to be binary (non-zeros will be used as

a weight, but should be non-negative for easy interpretation).

g an object of class "igraph" or "graphNEL"

Amatrix an affinity matrix pre-computed from the input graph. Notes: columns for start-

ing nodes walking from, and rows for ending nodes walking to

permutation how to do permutation. It can be 'degree' for degree-preserving permutation,

'random' for permutation purely in random

num.permutation

the number of permutations used to for generating the distribution of contact

strength under randomalisation

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p.adjust.method

the method used to adjust p-values. It can be one of "BH", "BY", "bonferroni", "holm", "hochberg" and "hommel". The first two methods "BH" (widely used) and "BY" control the false discovery rate (FDR: the expected proportion of false discoveries amongst the rejected hypotheses); the last four methods "bonferroni", "holm", "hochberg" and "hommel" are designed to give strong control of the family-wise error rate (FWER). Notes: FDR is a less stringent condition than FWER

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adjp.cutoff the cutoff of adjusted pvalue to construct the contact graph

parallel logical to indicate whether parallel computation with multicores is used. By de-

fault, it sets to true, but not necessarily does so. Partly because parallel backends available will be system-specific (now only Linux or Mac OS). Also, it will depend on whether these two packages "foreach" and "doMC" have been installed. It can be installed via: source("http://bioconductor.org/biocLite.R"); biocLite(c("foreach", "doMC")). If not yet installed, this option will be dis-

abled

multicores an integer to specify how many cores will be registered as the multicore parallel

backend to the 'foreach' package. If NULL, it will use a half of cores available in a user's computer. This option only works when parallel computation is enabled

verbose logical to indicate whether the messages will be displayed in the screen. By

default, it sets to true for display

#### Value

an object of class "dContact", a list with following components:

- ratio: a symmetric matrix storing ratio (the observed against the expected) between pairwise samples
- zscore: a symmetric matrix storing zscore between pairwise samples
- pval: a symmetric matrix storing pvalue between pairwise samples
- adjpval: a symmetric matrix storing adjusted pvalue between pairwise samples
- cgraph: the constructed contact graph (as a 'igraph' object) under the cutoff of adjusted value
- call: the call that produced this result

## Note

none

#### See Also

dRWR, dCheckParallel

## **Examples**

```
# 1) generate a random graph according to the ER model
g <- erdos.renyi.game(100, 1/100)

# 2) produce the induced subgraph only based on the nodes in query
subg <- dNetInduce(g, V(g), knn=0)
V(subg)$name <- 1:vcount(subg)

# 3) pre-compute affinity matrix from the input graph</pre>
```

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```
Amatrix <- dRWR(g=subg, parallel=FALSE)

# 4) estimate RWR-based sample relationships

# define sets of seeds as data

# each seed with equal weight (i.e. all non-zero entries are 1)
aSeeds <- c(1,0,1,0,1)
bSeeds <- c(0,0,1,0,1)
data <- data.frame(aSeeds,bSeeds)
rownames(data) <- 1:5

# calcualte their two contacts
dContact <- dRWRcontact(data=data, g=subg, Amatrix=Amatrix, parallel=FALSE)
dContact
```