# Package 'DRaWR'

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Title Discriminative Random Walk with Restart

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Author Charles Blatti [aut, cre]

Maintainer Charles Blatti <blatti@illinois.edu>

**Description** We present DRaWR, a network-based method for ranking genes or properties related to a given gene set. Such related genes or properties are identified from among the nodes of a large, heterogeneous network of biological information. Our method involves a random walk with restarts, performed on an initial network with multiple node and edge types, preserving more of the original, specific property information than current methods that operate on homogeneous networks. In this first stage of our algorithm, we find the properties that are the most relevant to the given gene set and extract a subnetwork of the original network, comprising only the relevant properties. We then rerank genes by their similarity to the given gene set, based on a second random walk with restarts, performed on the above subnetwork.

**Depends** R (>= 3.0)

License GPL-2

LazyData true

Imports Matrix, ROCR

RoxygenNote 5.0.1.9000 NeedsCompilation no

**Repository** CRAN

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DRaWR

#### Description

This function runs the DRaWR two stage random walk with restart method.

#### Usage

```
DRaWR(possetfile = "extdata/sample.setlist", unifile = "extdata/sample.uni",
    networkfile = "extdata/sample.edge", outdir = "output_",
    restarts = c(0.7), nfolds = 1, st2keep = 1, undirected = TRUE,
    unweighted = FALSE, normalize = "type", maxiters = 50, thresh = 1e-04,
    property_types = c("allen_brain_atlas", "chip_binding", "gene_ontology",
    "motif_u5", "pfam_domain", "T1", "T2"), writepreds = 0)
```

#### Arguments

possetfile	(string): location of file containing location of gene sets to test.
unifile	(string): location of file listing gene universe.
networkfile	(string): location of file containing network contents.
outdir	(string): prefix of location of file to write performance results (optionally prediction results).
restarts	(vector): vector of restart values to test. Default is $c(0.7)$ .
nfolds	(int): number of folds for cross validation, 1 is no cross-validation. Default is 4.
st2keep	(int): number of property nodes to keep in second stage for each property type. Default is 50.
undirected	(bool): boolean to make network undirected.
unweighted	(bool): boolean to make network unweighted.
normalize	(string): "type" or "none". Default is 'type'.
maxiters	(int): maximum number of allowable iterations. Default is 50.
thresh	(float): threshold for L1 norm convergence. Default is 0.001.
property_types	(vector): list of possible property types. Default is c("go_curated_evidence", "go_inferred_evidence", "pfam_domain").
writepreds	(boolean): write predictions out to a file. Default is FALSE

#### Examples

```
DRaWR(possetfile = system.file("extdata", "sample.setlist", package="DRaWR"),
unifile = system.file("extdata", "sample.uni", package="DRaWR"),
networkfile = system.file("extdata", "sample.edge", package="DRaWR"),
outdir = "output_", restarts = c(.7), nfolds = 1, st2keep = 1,
undirected = TRUE, unweighted = FALSE, normalize = "type", maxiters = 50,
thresh = 0.0001, property_types = c("T1", "T2"), writepreds = 0)
```

#### Description

RWR

This function runs a random walk with restart using two supported matrix representations.

#### Usage

RWR(boolSparceMat, transmat, restart, query, startvec, maxiters, thresh)

#### Arguments

boolSparceMat	(bool): Boolean to indicate sparce Matrix or list matrix
transmat	(sparce Matrix / list matrix): transition probabilites.
restart	(float): probability of restart.
query	(vector): probability of restarting at all nodes.
startvec	(vector): initial probability of being at any node.
maxiters	(int): maximum number of allowable iterations.
thresh	(float): threshold for L1 norm convergence.

### Value

list of 'iter':number of iterations, 'diff': L1 norm of difference, 'vec': converged probability distribution vector.

#### Examples

```
RWR(boolSparceMat=TRUE, transmat=transmat, restart=.3, query=c(rep(0.1,10),rep(0,5)),
startvec=rep(1/15,15), maxiters=10, thresh=0.001)
```

threeCol2listMat threeCol2listMat

#### Description

This function takes a three vectors of equal length (source nodes, target nodes, and edge weights) and return the adjacency matrix as a list of vectors.

#### Usage

```
threeCol2listMat(a = c("a", "b", "c", "c"), b = c("a", "b", "b", "b"),
v = c(1, 2, 3, 4))
```

#### Arguments

а	(vector): vector of source node names.
b	(vector): vector of target node names.
v	(vector): vector of edge weights names.

#### Value

list of vectors matrix representation.

#### Examples

```
threeCol2listMat(a = c("a","b","c","c"), b = c("a","b","b","b"), v = c(1,2,3,4))
```

|--|

#### Description

This function takes a three vectors of equal length (source nodes, target nodes, and edge weights) and return the adjacency matrix as a sparse Matrix.

#### Usage

```
threeCol2MaxMat(a = c("a", "b", "c", "c"), b = c("a", "b", "b", "b"),
 v = c(1, 2, 3, 4))
```

#### Arguments

а	(vector): vector of source node names.
b	(vector): vector of target node names.
v	(vector): vector of edge weights names.

#### Value

sparce Matrix.

#### Examples

threeCol2MaxMat(a = c("a","b","c","c"), b = c("a","b","b","b"), v = c(1,2,3,4))

transmat

## Description

A Matrix containing the normalized transition matrix from the test network

#### Usage

transmat

#### Format

a Matrix containing the normalized transition matrix from the test network.

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