Package 'MCL'

July 21, 2025

Type Package
Title Markov Cluster Algorithm
Version 1.0
Date 2015-03-11
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Description Contains the Markov cluster algorithm (MCL) for identifying clusters in networks and graphs. The algorithm simulates random walks on a (n x n) matrix as the adjacency matrix of a graph. It alternates an expansion step and an inflation step until an equilibrium state is reached.
License GPL (>= 2)
Depends R ($>= 2.10.0$)
Imports expm
Encoding UTF-8
NeedsCompilation no
Repository CRAN
Date/Publication 2015-03-11 09:56:32
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MCL-package

Markov Cluster Algorithm

Description

Contains the Markov cluster algorithm (MCL) by van Dongen (2000) for identifying clusters in networks and graphs. The algorithm simulates random walks on a (n x n) matrix as the adjacency matrix of a graph. It alternates an expansion step and an inflation step until an equilibrium state is reached.

Details

Package: MCL
Type: Package
Version: 1.0
Date: 2015-03-10

License: GPL-2 | GPL-3 [expanded from: GPL (= 2)]

The Markov Cluster Algorithm (MCL) is a method to identify clusters in undirected network graphs. It is suitable for high-dimensional data (e.g. gene expression data).

The original MCL uses the adjacency matrix of a graph (propsed by van Dongen (2000)). The function mcl in this package allows in addition the input of a $(n \times n)$ matrix.

Note

We thank Moritz Hanke for his help in realizing this package.

Author(s)

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References

van Dongen, S.M. (2000) Graph Clustering by Flow Simulation. Ph.D. thesis, University of Utrecht. Utrecht University Repository: http://dspace.library.uu.nl/handle/1874/848

Examples

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```
0,0,0,0,0,0,0,0,0,0,0,0), \ byrow=TRUE, \ nrow=9) ### Run MCL mcl(x = adjacency, addLoops = TRUE)
```

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Markov Cluster Algorithm

Description

Perform the Markov Cluster Algorithm on an adjacency or (n x n) matrix.

Usage

Arguments

X	an adjacency or (n x n) matrix
addLoops	logical; if TRUE, self-loops with weight 1 are added to each vertex of \boldsymbol{x} (see Details).
expansion	numeric value > 1 for the expansion parameter
inflation	numeric value > 0 for the inflation power coefficient
allow1	logical; if TRUE, vertices are allowed to form their own cluster. If FALSE, clusters of size 1 are interpreted as background noise and grouped in one cluster.
max.iter	an interger, the maximum number of iterations for the MCL
ESM	logical whether the equilibrium state matrix should be returned (default is FALSE)

Details

The adjacency or correlation matrix x is clustered by the Markov Cluster algorithm. The algorithm is controlled by the expansion parameter and the inflation power coefficient (for further details, see reference below). Adding self-loops is necessary, if either x contains at least one vertex of degree 0 or x represents a directed, non-bipartite graph adjacency matrix (i.e. the upper or lower matrix of x contains only zeros).

Value

```
K the number of clusters

n.iterations the number of iterations

Cluster a vector of integers indicating the cluster to which each vertex is allocated Equilibrium.state.matrix

a matrix; rows contain clusters
```

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Note

If an error occurs, mcl returns the number of the last iteration. If an error occurs at iteration 1, there might be a problem with the matrix x. If an error occurs at iteration max.iter, x could not be transformed into an equilibrium state matrix.

Author(s)

Martin L. Jäger

References

van Dongen, S.M. (2000) Graph Clustering by Flow Simulation. Ph.D. thesis, University of Utrecht. Utrecht University Repository: http://dspace.library.uu.nl/handle/1874/848

Examples

```
### Generate adjacency matrix of undirected graph
0,1,0,0,0,0,1,1,1,0,0,0,0,0,0,0,1,0,0,
                    0,1,1,0,0,0,0,0,0,1,0,1,0,0,0,0,0,0,1,1,
                    byrow=TRUE, nrow=9)
### Plot graph (requires package igraph)
# library(igraph)
# gu <- graph.adjacency( adjacency, mode="undirected" )</pre>
# plot( gu )
### Run MCL
mcl(x = adjacency, addLoops=TRUE, ESM = TRUE)
### Allow clusters of size 1
mcl(x = adjacency, addLoops = TRUE, allow1 = TRUE)
### Error: Small inflation coefficient prevents that an
          equilibrium state matrix is reached within 100 iterations
mcl(x = adjacency, addLoops=TRUE, inflation = 1.01, max.iter = 100)
### Generate adjacency matrix of directed graph
dgr \leftarrow matrix(0, nrow = 10, ncol = 10)
dgr[2:3,1] <- 1; dgr[3:4,2] <- 1; dgr[5:6,4] <- 1
dgr[6:7,5] <- 1; dgr[8:9,7] <- 1; dgr[10,8:9] <- 1
### Plot graph (requires package igraph)
# library( igraph )
# gd <- graph.adjacency( dgr )</pre>
# plot( gd )
### Directed graphs require self-loops!
mcl(x = dgr, addLoops = TRUE)
```

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