Package ‘DCA’

Type Package

Title Dynamic Correlation Analysis for High Dimensional Data

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Description Finding dominant latent signals that regulate dynamic correlation between many pairs of variables.

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## Dynamic Correlation Analysis for high dimensional data

### Description

Given a data matrix with variables in rows and samples in the columns, the method DCA finds dominant latent signals that regulate the dynamic correlation between many pairs of variables.

### Details

The subroutine dca() computes dynamic correlation signals from the data matrix. It can use PCA, SPCA, and kmeans clustering to find dominant signals. The subroutine find.xy() subsequently finds variable pairs that are associated with each latent signal. The subroutine plot.la() plots the dynamic correlation of two variables X and Y given the Z vector.

### Author(s)

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### References

https://ru.arxiv.org/pdf/1705.02479


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## dca

### Dynamic Correlation Analysis

### Description

The method finds a series of latent vectors, which serve as the LA scouting vectors for large numbers of variable pairs.

### Usage

```r
dca(array, top.pairs.prop = 0.95, max.pairs = 1e+06, n.fac = 10,
    sumabsv = sqrt(max.pairs)/10, normalization = "standardize", method = "PCA")
```
Arguments

array  The data matrix, with variables in the rows and samples in the columns.
top.pairs.prop  The method ranks all variable pairs from the most likely to have dynamic correlation relationship to the least likely. The top pairs are used for detection of latent signals. This parameter controls the percentage of pairs used in the computation.
max.pairs  The maximum number of pairs to use. When the data contains too many variables, such as tens of thousands of variables in a gene expression matrix, this parameter limits the maximum number of variable pairs to enter the calculation.
n.fac  The number of top latent factors to report. If the method "kmeans" is used, this parameter is used as the number of clusters.
sumabsv  The sumabsv parameter to be passed on to the SPC() method.
normalization  The way the data matrix is to be row-normalized. The method requires each row to have mean 0 and SD 1. There are two options, "standardize", or "normal score".
method  The method for finding the latent factors. Current choices are "PCA", "SPCA", and "kmeans".

Details

After finding the factors, the method attempts to rotate the factor using oblique rotation to achieve more interpretable results.

Value

The method returns a list.

fac  The original factors found. This is the PC, SPC, or cluster mean vector depending on the method chosen.
rotated  The factors after rotation.
ss.proj  The sum of squared attributed to each rotated factor.

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See Also

find.xy()

Examples

x<-la.simu.gen(n=100,p=200,n.grp=2, n.noise.gene=100, rho=0.5, pwr=0.5)
z<-dca(x$dat, n.fac=2)
cor(z[,2], x$z, method="spearman")
find.xy

Find variable pairs for a given set of LA scouting vectors.

Description

After finding the latent vectors, this function can be used to find the pairs of variables (rows of the original data matrix) that are associated with each latent vector in terms of dynamic correlation.

Usage

```r
find.xy(array, z, fdr.cut=0.05, normalization="standardize",
         center.z=FALSE, lac.percentile=0.8)
```

Arguments

- `array`: The data matrix with variables in the rows and samples in the columns.
- `z`: The matrix of latent variables. Each column is a latent vector.
- `fdr.cut`: The threshold of local fdr for the selection of variable pairs.
- `normalization`: The way the data matrix is to be row-normalized. The method requires each row to have mean 0 and SD 1. There are two options, "standardize", or "normal score".
- `center.z`: Whether to remove mean from each z vector.
- `lac.percentile`: The variable pairs that are considered potentially dynamically correlated will enter the computation. This is determined by the percentile of the LAC score among all variable pairs. If the percentile is higher than the provided threshold, then they are considered.

Value

A list is returned. Each numbered item is a matrix with three columns: row number 1, row number 2, and local fdr value. Only those selected by fdr threshold are returned.

Author(s)

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See Also

dca()

Examples

```r
x<-la.simu.gen(n=100, p=80, n.grp=2, n.noise.gene=100, rho=0.5, pwr=0.25)
z<-dca(x$dat, n.fac=2)
xy<-find.xy(x$dat, z[[2]], fdr.cut=0.01)
summary(xy)
xy[[1]][1:5,]
xy[[2]][1:5,]
```
Simulation of a data matrix with underlying dynamic correlation signal in the Liquid Association (LA) framework

Description

The simulation follows the LA framework, namely dynamic correlation is in the form of \( X \sim N(0,1), Y \sim N(0,1), Z \sim N(0,1), E(XY|Z) \) is a function of \( Z \).

Usage

```r
la.simu.gen(n, p, n.grp, n.noise.gene, rho, pwr)
```

Arguments

- **n**: Sample size (number of columns of the data matrix).
- **p**: The number of genes in each LA module, i.e. a group of genes regulated by the same latent dynamic correlation factor.
- **n.grp**: The number of LA modules to simulate.
- **n.noise.gene**: The number of pure noise genes to add to the matrix.
- **rho**: The standard deviation of the Gaussian noise to be added to the simulated data in the modules.
- **pwr**: The power for the transformation (see details)

Details

Between modules, the latent LA factor \( z \)'s are independent.

Within each module, 10 sub-modules are simulated. For each sub-module, we first generate a pair of \( X \) and \( Y \) vectors, which follows:

\[
X \sim N(0,1), Y \sim N(0,1) \quad u = (\text{pnorm}(z) - 0.5) \times 2 \quad E(XY|z) = \text{sign}(u) \times |u|^{pwr}
\]

Then white noise with SD of \( \rho \) is added to the hidden \( X, Y \) pair to generate pairs of observed \( X, Y \) vectors.

Value

A list is returned.

- **dat**: The data matrix.
- **z**: The true \( z \) vectors.

Author(s)

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See Also
dca()

Examples
   x<-la.simu.gen(n=100, p=200, n.grp=3, n.noise.gene=100, rho=0.5, pwr=1)
x$dat[1:5,1:5]
x$z[1:5,]

plot_la  Visualizing the dynamic correlation between a pair of genes given the AL scouting vector.

Description
   Given three vectors, x, y and z, the function produces a color scatter plot of x and y, colored by the grouping of z.

Usage
   plot_la(x, y, z, use.locfdr = FALSE, cols = c("red", "green", "blue"), cex = 0.5)

Arguments
   x   The x vector.
   y   The y vector.
   z   The z vector, conditioned on which X and Y have dynamic correlation.
   use.locfdr   Whether to use local fdr to group the z values. If TRUE, locfdr() is used and the fdr (posterior probability of belonging to the more extreme groups) threshold is 0.5. If FALSE, the z values are cut at the 0.33 and 0.67 quantiles into three groups.
   cols   The colors of the groups.
   cex   The point size to be passed to plot().

Details
   The locfdr approach is only to be used when it is clear z has heavy tails, and it is believed such points are driving the dynamic correlation.

Value
   A vector, each item is a string, recording the LA score and the correlation in each of the point groups as determined by z values.

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plot_la

Examples

```r
x <- la.simu.gen(n=100, p=20, n.grp=2, n.noise.gene=10, rho=0.25, pwr=1)
plot_la(x$dat[,1], x$dat[,2], x$z[,1], use.locfdr=FALSE)
```
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