

Package ‘dhga’

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Title Differential Hub Gene Analysis

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Description Identification of hub genes in a gene co-expression network from gene expression data. The differential network analysis for two contrasting conditions leads to the identification of various types of hubs like Housekeeping, Unique to stress (Disease) and Unique to control (Normal) hub genes.

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Adjacency

Construction of Gene Co-expression Network

Description

The function computes the adjacency matrix using the soft threshold parameter to construct a Gene Co-expression Network

Usage

```
Adjacency(x, beta, threshold)
```

Arguments

| | |
|-----------|---|
| x | x is a data frame of gene expression values where rows represent genes and columns represent samples/time points under a particular condition/stress/trait. |
| beta | beta is a soft threshold parameter determined from the scale free property of biological networks (Gene Co-expression Network). |
| threshold | threshold is a scalar representing the threshold value for the adjacency matrix to construct the Gene Co-expression Network. |

Value

The function returns a list of outputs like adjacency matrix, Node-list and Edge-list for constructing a Gene Co-expression Network at certain threshold level

Author(s)

Samarendra Das and Baidya Nath Mandal

Examples

```
data(rice_normal)
beta=6
threshold=0.4
Adjacency(rice_normal,6,0.4)
```

DiffHub

Differential Hub status of the genes in a gene co-expression network

Description

The function returns hub status of each gene in a gene co-expression network under two contrasting conditions

Usage

```
DiffHub(x,y,m1,m2,s,beta,alpha, plot=TRUE)
```

Arguments

| | |
|-------|---|
| x | x is a data frame of gene expression values where rows represent genes and columns represent samples/time point under stress condition. |
| y | y is a data frame of gene expression values where rows represent genes and columns represent samples/time points under control condition. |
| m1 | m1 is a scalar representing sample size and is less than or equal to number of columns in x. |
| m2 | m2 is a scalar representing sample size and is less than or equal to number of columns in y. |
| s | s is a scalar representing number of times each of the m samples will be resampled. |
| beta | beta is a soft threshold parameter determined from the scale free property of biological networks (GCN). |
| alpha | alpha is a scalar representing statistical level of significance. Default is alpha=0.0001 |
| plot | plot is a character representing whether the hubplot can be drawn (TRUE) or not (FALSE). |

Value

The function returns a list with two components. First component returns the list of genes along with their hub status. Second component gives a table containing number of genes under different categories of hubs viz. housekeeping, unique to stress and unique to normal. The function also returns a venn plot of hub genes and unique hub genes under two conditions.

Author(s)

Samarendra Das and Baidya Nath Mandal

Examples

```
data(rice_salt)
data(rice_normal)
DiffHub(rice_salt,rice_normal,18,18,80, 6, alpha=0.0001, plot=TRUE)
```

| | |
|-----------------|---|
| hub.pval.cutoff | <i>Identification of hub genes in a Gene Co-expression Network based on gene connection significance values</i> |
|-----------------|---|

Description

The function returns list of top ranked hub genes under a particular condition/trait

Usage

```
hub.pval.cutoff(x, beta, m, s, n)
```

Arguments

| | |
|------|---|
| x | x is a data frame of gene expression values where rows represent genes and columns represent samples/time point under a particular condition/trait. |
| beta | beta is a soft threshold parameter determined from the scale free property of biological networks (Gene Co-expression Network). |
| m | m is a scalar representing sample size and is less than or equal to number of columns in x. |
| s | s is a scalar representing number of times each of the m samples will be resampled. |
| n | n is a scalar representing number of top ranked hub genes to be selected under that particular condition/trait |

Value

The function returns a vector containing list of top ranked hub genes for that particular condition/trait

Author(s)

Samarendra Das and Baidya Nath Mandal

Examples

```
data(rice_salt)
x=as.data.frame(rice_salt)
beta=6
m=18
s=80
n=20
hub.pval.cutoff(x, beta, m, s, n)
```

| | |
|---------|---|
| hub.wgs | <i>Identification of hub genes in a Gene Co-expression Network based on Weighted Gene Score</i> |
|---------|---|

Description

The function returns the list of hub genes in a Gene Co-expression Network based on Weighted Gene Score without considering statistical significance values.

Usage

```
hub.wgs(x, beta, n)
```

Arguments

| | |
|------|--|
| x | x is a data frame of gene expression values where rows represent genes and columns represent samples/time. |
| beta | beta is a soft threshold parameter determined from the scale free property of biological networks (Gene Co-expression Network). |
| n | n is a scalar representing number of genes to be declared as hub in the Gene Co-expression Network based on Weighted Gene Score. |

Value

The function returns a list of hub genes in the Gene Co-expression Network

Author(s)

Samarendra Das and Baidya Nath Mandal

Examples

```
data(rice_salt)
hub.wgs(rice_salt, beta=6, n=20)
```

| | |
|---------|--|
| HubPlot | <i>Venn plot of hub genes under two conditions namely stress and control</i> |
|---------|--|

Description

The function produces a venn diagram of number of hub genes and unique hub genes under two conditions

Usage

```
HubPlot(pvalue.stress, pvalue.control, alpha)
```

Arguments

`pvalue.stress` `pvalue.stress` is a vector of p-values for genes in the GCN under stress condition. `pvalue.stress` can be obtained from the output of `hub.stress` function.

`pvalue.control` `pvalue.control` is a vector of p-values for genes in the GCN under normal or control condition. `pvalue.control` can be obtained from the output of `hub.control` function.

`alpha` `alpha` is a scalar representing statistical level of significance. Default is `alpha=0.0001`

Value

The function produces a venn plot of hub genes and unique hub genes under two conditions

Author(s)

Samarendra Das and Baidya Nath Mandal

Examples

```
data(rice_salt)
pval.stres <- pvalue.hub(rice_salt, beta=6, m=18, s=80, plot=FALSE)
pvalue.stress <- pval.stres[, 2]

data(rice_normal)
pval.control <- pvalue.hub(rice_normal, beta=6, m=18, s=80, plot=FALSE)
pvalue.control <- pval.control[, 2]
HubPlot(pvalue.stress, pvalue.control, alpha=0.0001)
```

HubStatus

Hub status of the genes in a gene co-expression network

Description

The function returns hub status of each gene in a gene co-expression network (GCN)

Usage

```
HubStatus(pvalue.stress, pvalue.control, alpha)
```

Arguments

- `pvalue.stress` `pvalue.stress` is a vector of p-values for genes in the GCN under stress condition. `pvalue.stress` can be obtained from the output of `hub.stress` function.
- `pvalue.control` `pvalue.control` is a vector of p-values for genes in the GCN under normal or control condition. `pvalue.control` can be obtained from the output of `hub.control` function.
- `alpha` `alpha` is a scalar representing statistical level of significance. Default is `alpha=0.0001`

Value

The function returns a list with two components. First component returns the list of genes along with their hub status. Second component gives a table containing number of genes under different categories of hubs viz. housekeeping, unique to stress and unique to normal

Author(s)

Samarendra Das and Baidya Nath Mandal

Examples

```
data(rice_salt)
pval.stres <- pvalue.hub(rice_salt, beta=6, m=18, s=80, plot=FALSE)
p1 <- pval.stres[, 2]

data(rice_normal)
pval.control <- pvalue.hub(rice_normal, beta=6, m=18, s=80, plot=FALSE)
p2 <- pval.control[, 2]
HubStatus(p1,p2,alpha=0.0001)
```

| | |
|------------|--|
| pvalue.hub | <i>Computation of gene connection sigificance values in a Gene Co-expression network</i> |
|------------|--|

Description

The function computes ths statistical significance values of gene connections in a GCN for a particular condition/trait/stress

Usage

```
pvalue.hub(x, beta, m, s, plot=TRUE)
```

Arguments

| | |
|------|---|
| x | x is a data frame of gene expression values where rows represent genes and columns represent samples/time point under a particular condition/trait/stress. |
| beta | beta is a soft threshold parameter determined from the scale free property of biological networks (Gene Co-expression network). |
| m | m is a scalar representing sample size and is less than or equal to number of columns in x. |
| s | s is a scalar representing number of times each of the m samples will be resampled. |
| plot | plot is a character must either take logical value TRUE/FALSE representing whether the plot of the pvalues of gene connections of all genes in Gene Co-expression Network needs to be constructed or not. |

Value

The function returns a vector of p-values for all the genes in the GCN

Author(s)

Samarendra Das and Baidya Nath Mandal

Examples

```
data(rice_salt)
x=as.data.frame(rice_salt)
beta=6
m=18
s=80
pvalue.hub(x, beta, m, s, plot=TRUE)
```

rice_normal

The gene expression data of rice under control or normal condition

Description

rice_normal data has the gene expression values of 200 genes over 20 samples. The row names of the data are the names of the probes.

Usage

```
data(rice_normal)
```

Format

A data frame with 200 rows and 20 columns.

Details

rice_normal data has the gene expression values of 200 genes identified by Wang et al. (2013). After loading the data, the dataset is available in R in the name of rice_normal.

Source

Wang J, Chen L, Wang Y, Zhang J, Liang Y, Xu D (2013) A Computational systems biology study for understanding salt tolerance mechanism in Rice. PLoS one 8(6): e64929. doi:10.1371/journal.pone.0064929.

Examples

```
data(rice_normal)
```

rice_salt

The gene expression data of rice under salinity stress condition

Description

rice_salt data has the gene expression values of 200 genes over 20 samples. The row names of the data are the names of the probes.

Usage

```
data("rice_salt")
```

Format

A data frame with 200 rows and 20 columns.

Details

rice_salt data has the gene expression values of 200 genes identified by Wang et al. (2013). After loading the data, the dataset is available in R in the name of rice_salt object.

Source

Wang J, Chen L, Wang Y, Zhang J, Liang Y, Xu D (2013) A Computational systems biology study for understanding salt tolerance mechanism in Rice. PLoS one 8(6): e64929. doi:10.1371/journal.pone.0064929.

Examples

```
data(rice_salt)
```

| | |
|-------------------|--|
| WeightedGeneScore | <i>Computation of weighted gene score of genes in Gene Co-expression Network</i> |
|-------------------|--|

Description

The function computes the weighted gene scores of all genes in a Gene Co-expression Network

Usage

```
WeightedGeneScore(x, beta, plot=TRUE)
```

Arguments

| | |
|------|---|
| x | x is a data frame of gene expression values where rows represent genes and columns represent samples/time points under a particular condition/stress/trait. |
| beta | beta is a soft threshold parameter determined from the scale free property of biological networks (Gene Co-expression Network). |
| plot | plot is a character must either take logical value TRUE/FALSE representing whether the plot of the gene connection degrees of all genes in Gene Co-expression Network needs to be constructed or not. |

Value

The function returns a vector of weighted gene scores of genes (nodes) in the Gene Co-expression Network along with the gene connectivity plot.

Author(s)

Samarendra Das and Baidya Nath Mandal

Examples

```
data(rice_normal)
beta=6
WeightedGeneScore(rice_normal, beta, plot=FALSE)
```

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