Package: csdR (via r-universe)

June 30, 2024

Title Differential gene co-expression

Version 1.11.0 **Date** 04-02-2021

Description This package contains functionality to run differential gene co-expression across two different conditions. The algorithm is inspired by Voigt et al. 2017 and finds Conserved, Specific and Differentiated genes (hence the name CSD). This package include efficient and variance calculation by bootstrapping and Welford's algorithm.

Depends R (>= 4.1.0)

Imports WGCNA, glue, RhpcBLASctl, matrixStats, Rcpp

License GPL-3 Encoding UTF-8

NeedsCompilation yes

BugReports https://github.com/AlmaasLab/csdR/issues

URL https://almaaslab.github.io/csdR,
 https://github.com/AlmaasLab/csdR

Roxygen list(markdown = TRUE)

RoxygenNote 7.1.2

LinkingTo Rcpp

biocViews DifferentialExpression, GraphAndNetwork, GeneExpression, Network

Suggests rmarkdown, knitr, testthat (>= 3.0.0), BiocStyle, magrittr, igraph, dplyr

Config/testthat/edition 3

VignetteBuilder knitr

RemoteUrl https://bioc.r-universe.dev **RemoteUrl** https://github.com/bioc/csdR

RemoteRef HEAD

RemoteSha 14407273b7752f26d52e3cc78db6603f3e7ca680

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data-	-expression	San	ıple	ex	pre	ess	ioi	n n	nat	ric	ces	fe	or	C_{λ}	SD)								

Description

Sample expression matrices of thyroid gland tissue for thyroid cancer patients and healthy individuals. These datasets were pre-processed by Gulla et al. (2019). Due to size requirements, only 1000 randomly selected genes are provided in the dataset. Number of samples are 399 and 504 in the healthy controls and the sick samples, respectively.

Usage

```
data(normal_expression)
data(sick_expression)
```

Format

Numeric matrices of normalized gene expression. Genes are in columns, whereas samples are in rows.

Source

For the expression matrix for healthy individuals, GenotypeTissue Expression (GTEx) V7. For the thyroid cancer patients, the data are obtained for the Thyroid Cancer project (THCA) from The Cancer Genome Atlas (TCGA).

References

Gulla, Almaas, Eivind, & Voigt, André (2019). An integrated systems biology approach to investigate transcriptomic data of thyroid carcinoma. NTNU. http://hdl.handle.net/11250/2621725

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partial_argsort

Extract indecies corresponding to the largest elements

Description

Extracts the indecies of the n largest elements of the input This procedure is equivalent to order(x, decreasing = TRUE)[1:n_elements], but is much faster and avoids the overhead of sorting discarded elements. This function is useful for extracting the rows in a data frame having the largest values in one of the columns.

Usage

```
partial_argsort(x, n_elements)
```

Arguments

x Numeric vector, the vector containing the numbers to sort.

n_elements Integer scalar, the number of indecies to return.

Value

Numeric vector, the indecies of the largest elements (in sorted order) in x.

Examples

```
x <- c(10L,5L,-2L,12L,15L)
max_indecies <- partial_argsort(x,3L)
max_indecies
x[max_indecies]
order(x)[1:3]
mtcars[partial_argsort(mtcars$hp,5L),]</pre>
```

run_cor_bootstrap

Run bootstrapping of Spearman correlations within a dataset

Description

This function provides the more low-level functionality of bootstrapping the Spearman correlations of the columns within a dataset. Only use this function if you want a low-level interface, else run_csd provides a more streamlined approach if you want to do a CSD analysis.

run_csd

Usage

```
run_cor_bootstrap(
    x,
    n_it = 20L,
    nThreads = 1L,
    verbose = TRUE,
    iterations_gap = 1L
)
```

Arguments

x Numeric matrix, the gene expression matrix to analyse. Genes are in columns,

samples are in rows.

n_it Integer, number of bootstrap iterations

nThreads Integer, number of threads to use for computations

verbose Logical, should progress be printed?

iterations_gap If output is verbose - number of iterations after issue a status message (Default=1

- displayed only if verbose=TRUE)

Value

A list with two fields

rho Numeric matrix constaining the bootstrapped mean of the Spearman correlation between each

var Numeric matrix constaining the bootstrapped variance of the Spearman correlation between each column

Examples

```
data("normal_expression")
cor_res <- run_cor_bootstrap(
    x = normal_expression,
    n_it = 100, nThreads = 2L
)</pre>
```

run_csd

Run CSD analysis

Description

This function implements the a CSD based on the one presented by Voigt et al. 2017. All pairs of genes are first compared within each condition by the Spearman correlation and the correlation and its variance are estimated by bootstrapping. Finally, the results for the two conditions are compared and C-, S- and D-values are computed and returned.

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Usage

```
run_csd(
    x_1,
    x_2,
    n_it = 20L,
    nThreads = 1L,
    verbose = TRUE,
    iterations_gap = 1L
)
```

Arguments

x_1	Numeric matrix, the gene expression matrix for the first condition. Genes are in columns, samples are in rows. The columns must be named with the name of the genes. Missing values are not allowed.										
x_2	Numeric matrix, the gene expression matrix for the second condition.										
n_it	Integer, number of bootstrap iterations										
nThreads	Integer, number of threads to use for computations										
verbose	Logical, should progress be printed?										
iterations_gap	If output is verbose - number of iterations after issue a status message (Default=1 - displayed only if verbose=TRUE)										

Details

The gene names in x_1 and x_2 do not need to be in the same order, but must be in the same namespace. Only genes present in both datasets will be considered for the analysis.

Value

A data.frame with the additional class attribute csd_res with the results of the CSD analysis. This frame has a row for each pair of genes and has the following columns:

Gene1 Character, the name of the first gene

Gene2 Character, the name of the second gene

rho1 Mean correlation of the two genes in the first condition

rho2 Mean correlation of the two genes in the second condition

var1 The estimated variance of rho1 determined by bootstrapping

var2 The estimated variance of rho2 determined by bootstrapping

cVal Numeric, the conserved score. A high value indicates that the co-expression of the two genes have the same sign in both conditions

sVal Numeric, the specific score. A high value indicates that the co-expression of the two genes have a high degree of co-expression in one condition, but not the other.

dVal Numeric, the differentiated score. A high value indicates that the co-expression of the two genes have a high degree of co-expression in both condition, but the sign of co-expression is different.

fun_csd

References

Voigt A, Nowick K and Almaas E 'A composite network of conserved and tissue specific gene interactions reveals possible genetic interactions in glioma' In: *PLOS Computational Biology* 13(9): e1005739. (doi: https://doi.org/10.1371/journal.pcbi.1005739)

Examples

```
data("sick_expression")
data("normal_expression")
cor_res <- run_csd(
    x_1 = sick_expression, x_2 = normal_expression,
    n_it = 100, nThreads = 2L
)
c_max <- max(cor_res$cVal)</pre>
```

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