Package ‘lionessR’

May 20, 2024

Title  Modeling networks for individual samples using LIONESS
Version  1.18.0
Description  LIONESS, or Linear Interpolation to Obtain Network Estimates for Single Samples, can be used to reconstruct single-sample networks (https://arxiv.org/abs/1505.06440). This code implements the LIONESS equation in the lioness function in R to reconstruct single-sample networks. The default network reconstruction method we use is based on Pearson correlation. However, lionessR can run on any network reconstruction algorithms that returns a complete, weighted adjacency matrix. lionessR works for both unipartite and bipartite networks.
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LazyData  TRUE
Imports  stats, SummarizedExperiment, S4Vectors
Depends  R (>= 3.6.0)
Suggests  knitr, rmarkdown, igraph, reshape2, limma,
biocViews  Network, NetworkInference, GeneExpression
BugReports  https://github.com/mararie/lionessR/issues
URL  https://github.com/mararie/lionessR
Encoding  UTF-8
RoxygenNote  6.0.1
VignetteBuilder  knitr
git_url  https://git.bioconductor.org/packages/lionessR
git_branch  RELEASE_3_19
git_last_commit  d3a6951
git_last_commit_date  2024-04-30
Repository  Bioconductor 3.19
Date/Publication  2024-05-20
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Description

This function uses the LIONESS equation to estimate single-sample networks. The input supports numeric matrix or a SummerizedExperiment object.

Usage

lioness(x, f = netFun)

Arguments

x Numeric matrix with samples in columns or SummerizedExperiment object
f Network reconstruction function. Defaults to Pearson correlation.

Value

SummerizedExperiment object for single-sample association network. The rowData contains the information about the regulators and their targets, while the colData contains the information about the samples. The edge weights of sample specific networks can be accessed through the lioness assay of the object.

Examples

```r
exp <- matrix(sample(1000,1000)/1000, 100, 10)
genes <- paste("gene", c(1:nrow(exp)), sep="_")
samples <- paste("sample", c(1:ncol(exp)), sep="_")

rowData <- S4Vectors::DataFrame(row.names = genes, gene = genes)
colData <- S4Vectors::DataFrame(col.names = samples, sample = samples)

se <- SummarizedExperiment::SummarizedExperiment(assays = list(counts = exp),
    colData = colData, rowData = rowData)

lionessResults <- lioness(se, netFun)
```
### Description

This is the network reconstruction function that will be used to build aggregate networks.

### Usage

```r
netFun(x)
```

### Arguments

- **x**: Numeric matrix with samples in columns.

### Value

Numeric matrix of association network.

### Examples

```r
exp <- matrix(sample(1000, 1000)/1000, 100, 10)
netFun(exp)
```

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### OSdata

**OS data**

### Description

Pre-processed gene expression data from high-grade osteosarcoma biopsies and sample characteristics were downloaded from GEO (GSE42352). We converted nuIDs to gene symbols using the annotation platform GPL10295. For genes with duplicate gene symbols, we selected the gene with the highest variance. Finally, we subsetted the data to the 53 patients for which 5 year metastasis information was available. Finally, we extracted genes with top 10,000 variance to reduce the size of the dataset.

### Usage

```r
data(OSdata)
```

### Format

- **"exp"**: Data frame containing expression data for 10000 genes and 53 samples.
- **"targets"**: Data frame containing information on whether patients developed metastases within 5 years or not, 53 samples and 2 columns.
Index

* datasets
  OSdata, 3

* lioness
  lioness, 2

* netFun
  netFun, 3

exp (OSdata), 3
lioness, 2
netFun, 3
OSdata, 3
targets (OSdata), 3