

Package ‘GRENITS’

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Type Package

Title Gene Regulatory Network Inference Using Time Series

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Description The package offers four network inference statistical models using Dynamic Bayesian Networks and Gibbs Variable Selection: a linear interaction model, two linear interaction models with added experimental noise (Gaussian and Student distributed) for the case where replicates are available and a non-linear interaction model.

License GPL (>= 2)

LazyLoad yes

Depends R (>= 2.12.0), Rcpp (>= 0.8.6), RcppArmadillo (>= 0.2.8), ggplot2 (>= 0.9.0)

Imports graphics, grDevices, reshape2, stats, utils

Suggests network

LinkingTo Rcpp, RcppArmadillo

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<i>analyse.output</i>	<i>Analysis Plots</i>
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Description

Analyse output from network inference functions. Basic convergence and analysis plots.

Usage

```
analyse.output(output.folder, timeSeries = NULL)
```

Arguments

- output.folder Name of folder (including path) where chains are kept
- timeSeries Only used by NonLinearModel analysis. Data matrix containing gene expression time series. Where genes will be placed in rows and time points in columns.

Details

Read first two chains run and plot some basic convergence plots (ConvergencePlots.pdf), analysis plots (AnalysisPlots.pdf), as well as inferred network probabilities in two formats (NetworkProbability_List.txt and NetworkProbability_Matrix.txt).

Value

The output of the analysis will be four files (five if nonLinearNet). The contents of the two plot files change depending on the network inference function used.

ConvergencePlots.pdf

Basic convergence plots. The posterior means of each variable are compared.

AnalysisPlots.pdf

Heatmap plot of network link probabilities as well as marginal network uncertainty plot. A plot of the number of links predicted by the model for a given probability threshold. For ReplicatesNet_student, the posterior distribution of the degrees of freedom are also plotted. For NonLinearNet, the posterior of the smoothness parameter is plotted.

NetworkProbability_List.txt

Posterior probabilities for each network connection in list format, including posterior interaction strength for linear models. Can be imported with network analysis software such as cytoscape.

NetworkProbability_Matrix.txt

Posterior probabilities for each network connection in matrix format.

ProbNumParents.txt

Posterior probabilities for number of regulators for each gene.

InferredFunctionPlots.pdf

(Only for nonLinearNet) Posterior distribution of predicted functions. Data values are plotted as circles.

References

Morrissey, E.R., Juarez, M.A., Denby, K.J. and Burroughs, N.J. 2010. On reverse engineering of gene interaction networks using time course data with repeated measurements. Bioinformatics 2010; doi: 10.1093/bioinformatics/btq421

Morrissey, E.R., Juarez, M.A., Denby, K.J. and Burroughs, N.J. 2011 Inferring the time-invariant topology of a nonlinear sparse gene regulatory network using fully Bayesian spline autoregression Biostatistics 2011; doi: 10.1093/biostatistics/kxr009

See Also

[NonLinearNet](#), [LinearNet](#), [ReplicatesNet_student](#) , [ReplicatesNet_gauss](#).

Examples

```
# Load A. thaliana circadian clock ODE generated data
data(Athaliana_ODE)
# Folder where raw runs will be kept and analysed
output.folder <- paste(tempdir(), "/Example_LinearNet", sep="")
# Run network inference, place raw results in output.folder
LinearNet(output.folder, Athaliana_ODE)
# Analyse raw results, place analysis plots and files in output.folder
analyse.output(output.folder)
```

Athaliana_ODE*Gene expression time series generated with ODE model***Description**

This data set was generated using a circadian clock regulatory network ODE model. The data was simulated using COPASI, subsampled to produce hourly data and after this the log was calculated.

Usage

```
data(Athaliana_ODE)
```

Format

A matrix containing genes in rows and (1h) time points in columns.

References

Hoops, S., Sahle, S., Gauges, R., Lee, C., Pahle, J., Simus, N., Singhal, M., Xu, L., Mendes, P. and Kummer, U. (2006) COPASI a COmplex PATHway SImulator. Bioinformatics, 22, 3067-3074.

Locke, J.C.W., Kozma-Bognar, L., Gould, P.D., Feher, B., Kevei, E., Nagy, F., Turner, M.S., Hall, A. and Millar, A.J. (2006) Experimental validation of a predicted feedback loop in the multi-oscillator clock of *Arabidopsis thaliana*. Molecular Systems Biology

Athaliana_ODE_4NoiseReps*Gene expression time series generated with ODE model with added noise***Description**

This data set was generated using a circadian clock regulatory network ODE model after which noise was added. The data was simulated using COPASI, subsampled to produce hourly data and after this the log was caluclated. Further to this, four replicates were produced by adding student noise.

Usage

```
data(Athaliana_ODE_4NoiseReps)
```

Format

A matrix containing genes in rows and (1h) time points in columns. The four replicates are appended (columns)

References

Hoops, S., Sahle, S., Gauges, R., Lee, C., Pahle, J., Simus, N., Singhal, M., Xu, L., Mendes, P. and Kummer, U. (2006) COPASI a COmplex PATHway SImulator. Bioinformatics, 22, 3067-3074.

Locke, J.C.W., Kozma-Bognar, L., Gould, P.D., Feher, B., Kevei, E., Nagy, F., Turner, M.S., Hall, A. and Millar, A.J. (2006) Experimental validation of a predicted feedback loop in the multi-oscillator clock of *Arabidopsis thaliana*. Molecular Systems Biology

Description

Run Bayesian inference of linear interaction network. The function generates MCMC chains that can later be analysed.

Usage

```
LinearNet( resultsFolder,           timeSeries,   ParamVec = NULL,
           chains = 2, user.seeds = NULL, Regulators = NULL,
           fixMe = NULL)
```

Arguments

resultsFolder	Name of output folder. The folder will be created and the output of the run will be placed there.
timeSeries	Data matrix containing gene expression time series. Where genes will be placed in rows and time points in columns. Gene names may be included as row names.
ParamVec	A parameter vector created using "mcmc.defaultParams_Linear". If none is given, default parameters will be used. The vector contains parameters associated to the priors as well as MCMC run length. (See mcmc.defaultParams_Linear)
chains	Number of MCMC chains to run.
user.seeds	An optional vector with seeds to use for MCMC chains.
Regulators	An optional vector with the indices of which genes are regulators. If provided, all non-regulator genes will not be allowed to regulate.
fixMe	An optional matrix of size genes x genes, where columns represent regulators and rows regulated genes. The matrix informs the model of network connections known to be present/absent. For each position use either 0 (no regulation, fix off), 1 (known regulatory interaction, fix on) or NaN (no information, do not fix).

Value

For each chain run, a folder (chain1, chain2, ...) will be created and the output of the MCMC run will be placed there. The files will be B_mcmc (the coefficients of the linear regression), Gamma_mcmc (the indicator variables of Gibbs variable selection), Lambda_mcmc (the precision of each regression), Mu_mcmc (the intercept of each regression) and Rho_mcmc (the network connectivity parameter).

References

- Morrissey, E.R., Juarez, M.A., Denby, K.J. and Burroughs, N.J. 2010. On reverse engineering of gene interaction networks using time course data with repeated measurements. *Bioinformatics* 2010; doi: 10.1093/bioinformatics/btq421
- Morrissey, E.R., Juarez, M.A., Denby, K.J. and Burroughs, N.J. 2011 Inferring the time-invariant topology of a nonlinear sparse gene regulatory network using fully Bayesian spline autoregression. *Biostatistics* 2011; doi: 10.1093/biostatistics/kxr009

See Also

`mcmc.defaultParams_Linear`, `analyse.output`.

Examples

```
# Load A. thaliana circadian clock ODE generated data
data(Athaliana_ODE)
# Folder where raw runs will be kept and analysed
output.folder <- paste(tempdir(), "/Example_LinearNet", sep="")
# Run network inference, place raw results in output.folder
LinearNet(output.folder, Athaliana_ODE)
# Analyse raw results, place analysis plots and files in output.folder
analyse.output(output.folder)
```

`mcmc.defaultParams_gauss`

Default Parameters for Linear Model with Gaussian distributed replicates

Description

Create parameter vector with default parameters for `ReplicatesNet_gauss` function

Usage

`mcmc.defaultParams_gauss()`

Details

Use this function to generate a template parameter vector to use non-default parameters for the `ReplicatesNet_gauss` model.

Value

Returns a single vector with the following elements (in this order):

- | | |
|-------------|--|
| (1) samples | Number of MCMC iterations to run |
| (2) burn.in | Number of initial iterations to discard as burn in |

(3) thin	Subsampling frequency
(4) c	Shape parameter 1 for Beta(c,d) prior on rho (connectivity parameter)
(5) d	Shape parameter 2 for Beta(c,d) prior on rho (connectivity parameter)
(6) sigma.s	Standard deviation parameter for N(0,sigma.s) prior on B (Regression coefficients)
(7) a	Shape parameter for Gamma(a,b) prior on lambda (Regression precision)
(8) b	Rate parameter for Gamma(a,b) prior on lambda (Regression precision)
(9) a_exp	Shape parameter for Gamma(a_exp,b_exp) prior on tau (Replicates precision)
(10) b_exp	Rate parameter for Gamma(a_exp,b_exp) prior on tau (Replicates precision)
(11) sigma.mu	Standard deviation parameter for N(0,sigma.mu) prior on mu (Regression intercept)
(12) fix.y.iter	Number of iterations for which sampled data Y is fixed

References

Morrissey, E.R., Juarez, M.A., Denby, K.J. and Burroughs, N.J. 2010. On reverse engineering of gene interaction networks using time course data with repeated measurements. Bioinformatics 2010; doi: 10.1093/bioinformatics/btq421

See Also

[plotPriors](#), [ReplicatesNet_gauss](#).

Examples

```
# Get default parameters
linearNet_Gauss.params <- mcmc.defaultParams_gauss()

# Change run length
linearNet_Gauss.params[1] <- 200000

# Change prior regression precision
linearNet_Gauss.params[7] <- 0.001
linearNet_Gauss.params[8] <- 0.001

# Plot to visualise changes
plotPriors(linearNet_Gauss.params)

## Use to run ReplicatesNet_gauss ...
```

mcmc.defaultParams_Linear*Default Parameters for Linear Model***Description**

Create parameter vector with default parameters for LinearNet function

Usage

```
mcmc.defaultParams_Linear()
```

Details

Use this function to generate a template parameter vector to use non-default parameters for the LinearNet model.

Value

Returns a single vector with the following elements (in this order):

- | | |
|--------------|--|
| (1) samples | Number of MCMC iterations to run. |
| (2) burn.in | Number of initial iterations to discard as burn in. |
| (3) thin | Subsampling frequency |
| (4) c | Shape parameter 1 for Beta(c,d) prior on rho (connectivity parameter) |
| (5) d | Shape parameter 2 for Beta(c,d) prior on rho (connectivity parameter) |
| (6) sigma.s | Standard deviation parameter for N(0,sigma.s) prior on B (Regression coefficients) |
| (7) a | Shape parameter for Gamma(a,b) prior on lambda (Regression precision) |
| (8) b | Rate parameter for Gamma(a,b) prior on lambda (Regression precision) |
| (9) sigma.mu | Standard deviation parameter for N(0,sigma.mu) prior on mu (Regression intercept) |

References

Morrissey, E.R., Juarez, M.A., Denby, K.J. and Burroughs, N.J. 2010. On reverse engineering of gene interaction networks using time course data with repeated measurements. Bioinformatics 2010; doi: 10.1093/bioinformatics/btq421

Morrissey, E.R., Juarez, M.A., Denby, K.J. and Burroughs, N.J. 2011 Inferring the time-invariant topology of a nonlinear sparse gene regulatory network using fully Bayesian spline autoregression Biostatistics 2011; doi: 10.1093/biostatistics/kxr009

See Also

[plotPriors](#), [LinearNet](#).

Examples

```
# Get default parameters
linearNet.params <- mcmc.defaultParams_Linear()

# Change run length
linearNet.params[1] <- 150000

# Change prior regression precision
linearNet.params[7] <- 0.001
linearNet.params[8] <- 0.001

# Plot to check changes
plotPriors(linearNet.params)

## Use to run LinearNet ...
```

mcmc.defaultParams_nonLinear

Default Parameters for non-Linear Model

Description

Create parameter vector with default parameters for NonLinearNet function

Usage

```
mcmc.defaultParams_nonLinear()
```

Details

Use this function to generate a template parameter vector to use non-default parameters for the NonLinearNet model.

Value

Returns a single vector with the following elements (in this order):

- (1) samples Number of MCMC iterations to run.
- (2) burn.in Number of initial iterations to discard as burn in.
- (3) thin Subsampling frequency
- (4) c Shape parameter 1 for Beta(c,d) prior on rho (connectivity parameter)
- (5) d Shape parameter 2 for Beta(c,d) prior on rho (connectivity parameter)
- (6) trunc Truncation parameter for InvertedPareto prior on tau (smoothness parameter)
- (7) tau0 Precision parameter for N(0, tau0^(-0.5)) prior on B (first two coefficients)
- (8) M Numer of knots used for each spline function
- (9) a Shape parameter for Gamma(a,b) prior on lambda (Regression precision)

- (10) b Rate parameter for Gamma(a,b) prior on lambda (Regression precision)
- (11) sigma.mu Standard deviation parameter for N(0,sigma.mu) prior on mu (Regression intercept)
- (12) a_pareto Pareto parameter for InvertedPareto prior on tau (smoothness parameter)

References

Morrissey, E.R., Juarez, M.A., Denby, K.J. and Burroughs, N.J. 2011 Inferring the time-invariant topology of a nonlinear sparse gene regulatory network using fully Bayesian spline autoregression Biostatistics 2011; doi: 10.1093/biostatistics/kxr009

See Also

[plotPrior](#), [NonLinearNet](#).

Examples

```
# Get default parameters
nonLinearNet.params <- mcmc.defaultParams_nonLinear()

# Change run length
nonLinearNet.params[1] <- 150000

# Change prior on smoothness parameter
nonLinearNet.params[6] <- 30000 # Change truncation
nonLinearNet.params[12] <- 3 # Concentrate more mass close to linear region

# Plot to check changes
plotPrior(nonLinearNet.params)

## Use to run LinearNet ...
```

mcmc.defaultParams_student

Default Parameters for Linear Model with Student distributed replicates

Description

Create parameter vector with default parameters for ReplicatesNet_student function

Usage

`mcmc.defaultParams_student()`

Details

Use this function to generate a template parameter vector to use non-default parameters for the ReplicatesNet_student model.

Value

Returns a single vector with the following elements (in this order):

(1) samples	Number of MCMC iterations to run
(2) burn.in	Number of initial iterations to discard as burn in
(3) thin	Subsampling frequency
(4) c	Shape parameter 1 for Beta(c,d) prior on rho (connectivity parameter)
(5) d	Shape parameter 2 for Beta(c,d) prior on rho (connectivity parameter)
(6) sigma.s	Standard deviation parameter for N(0,sigma.s) prior on B (Regression coefficients)
(7) a	Shape parameter for Gamma(a,b) prior on lambda (Regression precision)
(8) b	Rate parameter for Gamma(a,b) prior on lambda (Regression precision)
(9) a_exp	Shape parameter for Gamma(a_exp,b_exp) prior on tau (Replicates precision)
(10) b_exp	Rate parameter for Gamma(a_exp,b_exp) prior on tau (Replicates precision)
(11) a_deg	Shape parameter for Gamma(a_deg,b_deg) prior on nu (Student degrees of freedom)
(12) b_deg	Rate parameter for Gamma(a_deg,b_deg) prior on nu (Student degrees of freedom)
(13) sigma.mu	Standard deviation parameter for N(0,sigma.mu) prior on mu (Regression intercept)
(14) fix.y.iter	Number of iterations for which sampled data Y is fixed

References

Morrissey, E.R., Juarez, M.A., Denby, K.J. and Burroughs, N.J. 2010. On reverse engineering of gene interaction networks using time course data with repeated measurements. Bioinformatics 2010; doi: 10.1093/bioinformatics/btq421

See Also

[plotPriors](#), [ReplicatesNet_student](#).

Examples

```
# Get default parameters
linearNet_Student.params <- mcmc.defaultParams_student()

# Change run length
linearNet_Student.params[1] <- 200000

# Change prior regression precision
linearNet_Student.params[7] <- 0.001
linearNet_Student.params[8] <- 0.001

# Plot to visualise changes
```

```
plotPriors(linearNet_Student.params)

## Use to run ReplicatesNet_student ...
```

Description

Run Bayesian inference of non-linear interaction network. Non linear interactions are modelled using Penalised Splines. The function generates MCMC chains that can later be analysed.

Usage

```
NonLinearNet( resultsFolder,           timeSeries,   ParamVec = NULL,
              chains = 2, user.seeds = NULL, Regulators = NULL,
              fixMe = NULL)
```

Arguments

<code>resultsFolder</code>	Name of output folder. The folder will be created and the output of the run will be placed there.
<code>timeSeries</code>	Data matrix containing gene expression time series. Where genes will be placed in rows and time points in columns. Gene names may be included as row names.
<code>ParamVec</code>	A parameter vector created using "mcmc.defaultParams_nonLinear". If none is given, default parameters will be used. The vector contains parameters associated to the priors as well as MCMC run length. (See <code>mcmc.defaultParams_nonLinear</code>)
<code>chains</code>	Number of MCMC chains to run.
<code>user.seeds</code>	An optional vector with seeds to use for MCMC chains.
<code>Regulators</code>	An optional vector with the indices of which genes are regulators. If provided, all non-regulator genes will not be allowed to regulate.
<code>fixMe</code>	An optional matrix of size genes x genes, where columns represent regulators and rows regulated genes. The matrix informs the model of network connections known to be present/absent. For each position use either 0 (no regulation, fix off), 1 (known regulatory interaction, fix on) or NaN (no information, do not fix).

Value

For each chain run, a folder (chain1, chain2, ...) will be created and the output of the MCMC run will be placed there. The files will be Gamma_mcmc (the indicator variables of Gibbs variable selection), Lambda_mcmc (the precision of each regression), Mu_mcmc (the intercept of each regression), Rho_mcmc (the network connectivity parameter), Tau_mcmc (the "smoothness parameter"), all_f (posterior mean of all functions), all_f_sqr (posterior mean of the square of all functions) and Full_F_sqr (posterior mean of the square of the sum of all functions, for each regression). For the files all_f and all_f_sqr functions are placed in column-wise order. The file is filled by placing all interactions for each regression one after another.

References

Morrissey, E.R., Juarez, M.A., Denby, K.J. and Burroughs, N.J. 2011 Inferring the time-invariant topology of a nonlinear sparse gene regulatory network using fully Bayesian spline autoregression Biostatistics 2011; doi: 10.1093/biostatistics/kxr009

See Also

[mcmc.defaultParams_nonLinear](#), [analyse.output](#).

Examples

```
# Synthetic data
data(Athaliana_ODE)
# Reduced data set to 3 genes 20 TP for faster run
Athaliana_ODE.reduced <- Athaliana_ODE[c(1,3,5),1:20]
# Folder where raw runs will be kept and later analysed
output.folder <- paste(tempdir(), "/ExampleNonLinearNet", sep = "")
# Run network inference and place raw results in output.folder
NonLinearNet(output.folder , Athaliana_ODE.reduced)
# Analyse raw results, place analysis plots and files in output.folder
analyse.output(output.folder, Athaliana_ODE.reduced)
```

plotPriors

Plot prior using parameter vector

Description

Plot appropriate priors using parameters from vector

Usage

plotPriors(parameter.vec)

Arguments

parameter.vec MCMC parameter vector of the type generated by e.g. [mcmc.defaultParams_Linear](#)

Details

This function takes the parameter vector that will be used for network inference function and plots the priors associated with the parameters given.

References

Morrissey, E.R., Juarez, M.A., Denby, K.J. and Burroughs, N.J. 2010. On reverse engineering of gene interaction networks using time course data with repeated measurements. Bioinformatics 2010; doi: 10.1093/bioinformatics/btq421

Morrissey, E.R., Juarez, M.A., Denby, K.J. and Burroughs, N.J. 2011 Inferring the time-invariant topology of a nonlinear sparse gene regulatory network using fully Bayesian spline autoregression Biostatistics 2011; doi: 10.1093/biostatistics/kxr009

See Also

`mcmc.defaultParams_gauss`, `mcmc.defaultParams_Linear`, `mcmc.defaultParams_nonLinear`,
`mcmc.defaultParams_student`.

Examples

```
# Get default parameters
nonLinearNet.params <- mcmc.defaultParams_nonLinear()

# Change run length
nonLinearNet.params[1] <- 150000

# Change prior on smoothness parameter
nonLinearNet.params[6] <- 30000 # Change truncation
nonLinearNet.params[12] <- 3 # Concentrate more mass close to linear region

# Plot to check changes
plotPriors(nonLinearNet.params)
```

read.chain

*Read MCMC Chains***Description**

Read MCMC chains for further analysis.

Usage

```
read.chain(output.folder, chainNumber)
```

Arguments

<code>output.folder</code>	Name of folder (including path) where chains are kept
<code>chainNumber</code>	Which of the chains will be read

Details

Read chains produced by NonLinearNet, LinearNet, ReplicatesNet_student and ReplicatesNet_gauss for further analysis.

Value

Returns a list of vectors/matrices with the value of the variables at each MCMC iteration.

References

Morrissey, E.R., Juarez, M.A., Denby, K.J. and Burroughs, N.J. 2010. On reverse engineering of gene interaction networks using time course data with repeated measurements. Bioinformatics 2010; doi: 10.1093/bioinformatics/btq421

Morrissey, E.R., Juarez, M.A., Denby, K.J. and Burroughs, N.J. 2011 Inferring the time-invariant topology of a nonlinear sparse gene regulatory network using fully Bayesian spline autoregression Biostatistics 2011; doi: 10.1093/biostatistics/kxr009

See Also

[NonLinearNet](#), [LinearNet](#), [ReplicatesNet_student](#), [ReplicatesNet_gauss](#).

Examples

```

#####
## Run inference using one chain
#####
# Load A. thaliana circadian clock ODE generated data
data(Athaliana_ODE)
# Folder where raw runs will be kept and analysed
output.folder <- paste(tempdir(), "/Example_LinearNet", sep="")
# Run network inference, place raw results in output.folder
# Run just one chain for example purpose
LinearNet(output.folder, Athaliana_ODE, chains = 1)

#####
## Read chain
#####
chain1 <- read.chain(output.folder, 1)

```

Description

Run Bayesian inference of linear interaction network on data with replicates. The replicates are assumed to follow a Gaussian distribution. The function generates MCMC chains that can later be analysed.

Usage

Arguments

<code>resultsFolder</code>	Name of output folder. The folder will be created and the output of the run will be placed there.
<code>timeSeries</code>	Data matrix containing gene expression time series. Where genes will be placed in rows and time points in columns. Each times series must be placed one after another, so that the first n columns correspond to time series replicate one, the next n columns to time series replicate two, etc. Gene names may be included as row names.
<code>numReps</code>	Number of replicate time series included in <code>timeSeries</code> matrix.
<code>ParamVec</code>	A parameter vector created using "mcmc.defaultParams_gauss". If none is given, default parameters will be used. The vector contains parameters associated to the priors as well as MCMC run length. (See <code>mcmc.defaultParams_gauss</code>)
<code>chains</code>	Number of MCMC chains to run.
<code>user.seeds</code>	An optional vector with seeds to use for MCMC chains.
<code>Regulators</code>	An optional vector with the indices of which genes are regulators. If provided, all non-regulator genes will not be allowed to regulate.
<code>fixMe</code>	An optional matrix of size genes x genes, where columns represent regulators and rows regulated genes. The matrix informs the model of network connections known to be present/absent. For each position use either 0 (no regulation, fix off), 1 (known regulatory interaction, fix on) or NaN (no information, do not fix).

Details

The order in which the replicates are placed do not affect the output. In other words swapping timepoint 2 replicate 1 and timepoint 2 replicate 2 makes no difference. For the cases where a measurement is not available, an NaN may be used.

Value

For each chain run, a folder (chain1, chain2, ...) will be created and the output of the MCMC run will be placed there. The files will be `B_mcmc` (the coefficients of the linear regression), `Gamma_mcmc` (the indicator variables of Gibbs variable selection), `Lambda_mcmc` (the precision of each regression), `Mu_mcmc` (the intercept of each regression), `Rho_mcmc` (the network connectivity parameter), `DataMean_standarised` (a times series with the mean of the replicates), `Lambda_exp_mcmc` (the precision of the replicate noise) and `Y_mean` (the mean posterior value of the inferred "true mRNA").

References

Morrissey, E.R., Juarez, M.A., Denby, K.J. and Burroughs, N.J. 2010. On reverse engineering of gene interaction networks using time course data with repeated measurements. Bioinformatics 2010; doi: 10.1093/bioinformatics/btq421

See Also

[mcmc.defaultParams_gauss](#), [analyse.output](#).

Examples

```
# Synthetic data
data(Athaliana_ODE_4NoiseReps)
# Folder where raw runs will be kept and later analysed
output.folder <- paste(tempdir(), "/Example_ReplicatesNet_Gauss", sep="")
# Run network inference and place raw results in output.folder
ReplicatesNet_gauss(output.folder, Athaliana_ODE_4NoiseReps, numReps = 4)
# Analyse raw results, place analysis plots and files in output.folder
analyse.output(output.folder)
```

ReplicatesNet_student *Dynamic Bayesian Network Inference Using Linear Interactions and Student Experimental Noise*

Description

Run Bayesian inference of linear interaction network on data with replicates. The replicates are assumed to follow a Student distribution. The function generates MCMC chains that can later be analysed.

Usage

```
ReplicatesNet_student(    resultsFolder,    timeSeries,
                         ParamVec = NULL,    chains = 2, user.seeds = NULL,
                         Regulators = NULL, fixMe = NULL)
```

Arguments

<code>resultsFolder</code>	Name of output folder. The folder will be created and the output of the run will be placed there.
<code>timeSeries</code>	Data matrix containing gene expression time series. Where genes will be placed in rows and time points in columns. Each times series must be placed one after another, so that the first n columns correspond to time series replicate one, the next n columns to time series replicate two, etc. Gene names may be included as row names.
<code>numReps</code>	Number of replicate time series included in <code>timeSeries</code> matrix.
<code>ParamVec</code>	A parameter vector created using "mcmc.defaultParams_student". If none is given, default parameters will be used. The vector contains parameters associated to the priors as well as MCMC run length. (See <code>mcmc.defaultParams_student</code>)
<code>chains</code>	Number of MCMC chains to run.
<code>user.seeds</code>	An optional vector with seeds to use for MCMC chains.
<code>Regulators</code>	An optional vector with the indices of which genes are regulators. If provided, all non-regulator genes will not be allowed to regulate.

<code>fixMe</code>	An optional matrix of size genes x genes, where columns represent regulators and rows regulated genes. The matrix informs the model of network connections known to be present/absent. For each position use either 0 (no regulation, fix off), 1 (known regulatory interaction, fix on) or NaN (no information, do not fix).
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Details

The order in which the replicates are placed do not affect the output. In other words swapping timepoint 2 replicate 1 and timepoint 2 replicate 2 makes no difference. For the cases where a measurement is not available, an NaN may be used.

Value

For each chain run, a folder (chain1, chain2, ...) will be created and the output of the MCMC run will be placed there. The files will be `B_mcmc` (the coefficients of the linear regression), `Gamma_mcmc` (the indicator variables of Gibbs variable selection), `Lambda_mcmc` (the precision of each regression), `Mu_mcmc` (the intercept of each regression), `Rho_mcmc` (the network connectivity parameter), `DataMean_standarised` (a times series with the mean of the replicates), `Lambda_exp_mcmc` (the precision of the replicate noise), `Y_mean` (the mean posterior value of the inferred "true mRNA"), `DegFreedom_mcmc` (degrees of freedom of the student distributed replicate noise) and `acceptanceRatio` (acceptance ratio of the Metropolis-Hastings degrees of freedom update).

References

Morrissey, E.R., Juarez, M.A., Denby, K.J. and Burroughs, N.J. 2010. On reverse engineering of gene interaction networks using time course data with repeated measurements. Bioinformatics 2010; doi: 10.1093/bioinformatics/btq421

See Also

[mcmc.defaultParams_student](#), [analyse.output](#).

Examples

```
# Synthetic data
data(Athaliana_ODE_4NoiseReps)
# Folder where raw runs will be kept and later analysed
output.folder <- paste(tempdir(), "/Example_ReplicatesNet_Student", sep="")
# Run network inference and place raw results in output.folder
ReplicatesNet_student(output.folder, Athaliana_ODE_4NoiseReps, numReps = 4)
# Analyse raw results, place analysis plots and files in output.folder
analyse.output(output.folder)
```

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