Package 'birta'

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Title Bayesian Inference of Regulation of Transcriptional Activity

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Description Expression levels of mRNA molecules are regulated by different processes, comprising inhibition or activation by transcription factors and post-transcriptional degradation by microRNAs. birta (Bayesian Inference of Regulation of Transcriptional Activity) uses the regulatory networks of TFs and miRNAs together with mRNA and miRNA expression data to predict switches in regulatory activity between two conditions. A Bayesian network is used to model the regulatory structure and Markov-Chain-Monte-Carlo is applied to sample the activity states.

License GPL (>= 2)

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birta-package

Joint Bayesian Inference of miRNA and Transcription Factor Activities

Description

Expression levels of mRNA molecules are regulated by different processes, comprising inhibition or activation by transcription factors and post-transcriptional degradation by microRNAs. birta uses the regulatory networks of TFs and miRNAs together with mRNA and miRNA expression data to predict switches in regulatory activity between two conditions. A Bayesian network is used to model the regulatory structure and Markov-Chain-Monte-Carlo is applied to sample the activity states.

Details

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Author(s)

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References

B. Zacher, K. Abnaof, S. Gade, E. Younesi, A. Tresch, and H. Frohlich. Joint Bayesian Inference of Condition Specific miRNA and Transcription Factor Activities from Combined Gene and microRNA Expression Data. *submitted*, 2012.

birta

Main interface for Bayesian Inference of Regulation of Transcriptional Activity.

Description

The function estimates parameterization of the model and then executes MCMC sampling to infer activity states.

birta

Usage

birta(dat.mRNA, dat.miRNA, TFexpr, limmamRNA=NULL, limmamiRNA=NULL, limmaTF=NULL, nrep=NULL, fdr.

Arguments

dat.mRNA	mRNA expression data (ExpressionSet or matrix). IMPORTANT: Replicates must be ordered according to nrep!
dat.miRNA	miRNA expression data (ExpressionSet or matrix).
TFexpr	TF expression data (ExpressionSet or matrix).
limmamRNA	Output of limma analysis for mRNA data (list: pvalue.tab, lm.fit).
limmamiRNA	Output of limma analysis for miRNA data (list: pvalue.tab, lm.fit).
limmaTF	Output of limma analysis for TF expression data (list: pvalue.tab, lm.fit).
nrep	Vector containing four integers. Entry 1 and 2 are the number of miRNA measurement replicates for condition 1 and 2. Entry 3 and 4 are the number of mRNA measurement replicates for condition 1 and 2.
fdr.mRNA	False discovery rate (FDR) cutoff for significance of the logFC for mRNA data.
fdr.miRNA	False discovery rate (FDR) cutoff for significance of the logFC for miRNA data.
lfc.mRNA	Additional logFC cutoff for significance in mRNA data.
lfc.miRNA	Additional logFC cutoff for significance in miRNA data.
genesets	Combined TF / miRNA network. IMPORTANT: Names of TF entries must start with V\\$.
lambda	Regularization parameter for edge weights.
sample.weights	Should edge weights be adapted during sampling?
one.regulator.	-
	Sould weights of all edges for a regulator to be the same?
theta_TF	Expected fraction of active TFs.
theta_miRNA	Expected fraction of active miRNAs.
model	Type of model. One out of c("all-plug-in", "weight-plug-in", "no-plug-in").
niter	Number of MCMC iterations (AFTER burnin).
nburnin potential_swap:	Number of MCMC iterations UNTIL burnin is assumed to be finished.
	Pre-computed potential swaps (OPTIONAL, see get_potential_swaps).
run.pretest	Initialize miRNA and TF states via the result of a hypergeometric test in order to improve convergence (should be taken with care; advise: only use it in case of observed convergence problems!).
condition.spec:	
	Should inference on TF / miRNA activities be made only RELATIVE to a ref- erence condition or independently in both conditions?
thin	Thinning of Markov chain: only use every thin's sample for posterior computa- tion.
only_switches	Should only switches be performed?
weightSampleMea	
	Mean for normal distribution used for sampling the omegas.
weightSampleVa	Variance for normal distribution used for sampling the omegas.

Value

The function returns a list containing the following entries:

miRNAstates1	Probability of each miRNA to be active in condition 1 (only for condition spe- cific sampling).
miRNAstates2	Probability of each miRNA to be active in condition 2 (only for condition spe- cific sampling).
miRNAActivityS	witch
	Probability of each miRNA switching its activities (non-condition specific sampling).
TFstates1	Probability of each TF to be active in condition 1 (only for condition specific sampling).
TFstates2	Probability of each TF to be active in condition 2 (only for condition specific sampling).
miRNAactivitySv	witch
	Probability of each TF switching its activities (non-condition specific sampling).
log_lik_trace	Log-likelihood trace of MCMC sampling.
TFomega	Weights of TF-target graph. (effect of a TF on its targets)
miRNAomega	Weights of miRNA-target graph. (effect of a miRNA on its targets)
genesetsTF	TF-target network. This might be different from the network submitted to the function, due to incompatibilities of network and experimental measurements. Check your warnings and command line output!
genesetsmiRNA	miRNA-target network. This might be different from the network submitted to the function, due to incompatibilities of network and experimental measurements. Check your warnings and command line output!
mRNAexpr	mRNA expression data. This might be different from the matrix submitted to the function, due to incompatibilities of network and experimental measurements. Check your warnings and command line output!
miRNAexpr	miRNA expression data. This might be different from the matrix submitted to the function, due to incompatibilities of network and experimental measurements. Check your warnings and command line output!
TFexpr	TF expression data (only if they are specifically included). This might be dif- ferent from the matrix submitted to the function, due to incompatibilities of network and experimental measurements. Check your warnings and command line output!

Author(s)

Holger Frohlich, Benedikt Zacher

Examples

```
data(humanSim)
design = model.matrix(~0+factor(c(rep("control", 5), rep("treated", 5))))
colnames(design) = c("control", "treated")
contrasts = "treated - control"
limmamRNA = limmaAnalysis(sim$dat.mRNA, design, contrasts)
limmamiRNA = limmaAnalysis(sim$dat.miRNA, design, contrasts)
sim_result = birta(sim$dat.mRNA, sim$dat.miRNA, limmamRNA=limmamRNA,
limmamiRNA=limmamiRNA, nrep=c(5,5,5,5), genesets=genesets,
```

birta-methods

```
model="all-plug-in", niter=50000, nburnin=10000,
sample.weights=FALSE, potential_swaps=potential_swaps)
```

birta-methods Ma

Methods for Function birta in Package birta

Description

Generic methods for function birta in package **birta**. The expression data can be stored in a matrix or an ExpressionSet.

Methods

The following signatures make sure, that the arguments, storing the expression data are in the correct format. signature(dat.mRNA = "ExpressionSet", dat.miRNA = "ExpressionSet", TFexpr = "ExpressionSet"gnatu

```
signature(dat.mRNA = "ExpressionSet", dat.miRNA = "missing", TFexpr = "ExpressionSet")
signature(dat.mRNA = "ExpressionSet", dat.miRNA = "missing", TFexpr = "missing")
```

```
signature(dat.mRNA = "matrix", dat.miRNA = "matrix", TFexpr = "matrix")
signature(dat.mRNA = "matrix", dat.miRNA = "matrix", TFexpr = "missing")
signature(dat.mRNA = "matrix", dat.miRNA = "missing", TFexpr = "matrix")
signature(dat.mRNA = "matrix", dat.miRNA = "missing", TFexpr = "missing")
```

EColiNetwork Example TF-target graph from Regulon DB.

Description

This list contains the TF-target graph used in the vignette.

Usage

EColiNetwork

Format

A list containing the target gene sets of 160 TFs.

Source

This TF-target graph was taken from (R. Castelo and A. Roverato, 2009). It is a pre-filtered version of RegulonDB 6.1.

References

R. Castelo and A. Roverato. Reverse engineering molecular regulatory networks from microarray data with qp-graphs. *J Comput Biol*, 16(2):213227, Feb 2009.

EColi0xygen

Description

This data set gives expression values for three experiments of the E. Coli K12 strain under aerobic and three experiments under aerobic growth. It is used in the vignette to illustrate application of birta to TFs only.

Usage

EColi0xygen

Format

ExpressionSet

Source

The original data comes from (Covert et al., 2004) The normalized data set used here is taken from the qpgraph package by R. Castelo and A. Roverato.

References

M. W. Covert, E. M. Knight, J. L. Reed, M. J. Herrgard, and B. O. Palsson. Integrating high-throughput and computational data elucidates bacterial networks. *Nature*, 429(6987):9296, May 2004.

genesets

TF-target and miRNA-target networks for simulated example.

Description

For miRNAs we looked at target predictions in human via miRanda (Betel et al., 2008), miRBase (Griffths-Jones et al., 2008) and miRDB (Wang and Naqa, 2008). Experimentally confirmed targets were retrieved from Tarbase (Papadopoulos et al., 2009). As trustworthy miRNA targets we considered those, which were either experimentally confirmed (i.e. listed in Tarbase) or predicted by at least two of the other three methods. In conclusion we arrived at a network with 583 miRNAs regulating between 1 and 1628 genes (median: 302).\ A TF-target gene network was compiled by computing TF binding affinities to promoter sequences of all human genes according to the TRAP model (Roideret al., 2007) via the author's R implementation. Upstream sequences of genes were retrieved here from the ENSEMBL database via biomaRt (Haider et al., 2009). We assumed that promoter sequences were located in the range 0 - 2Kbp upstream to the transcription start site of a gene. As trustworthy TF targets we considered those, for which a Holm corrected affinity p-value smaller than 0.001 was reported. In conclusion we arrived at a network with 272 TFs regulating between 1 and 16517 genes (median: 20).

Usage

genesets

Format

A list containing a TF- and miRNA-target graph.

Source

The networks were generated using the TRAP model on TRANSFAC matrices and miRNA-target annotations form different databases.

References

D. Betel, M. Wilson, A. Gabow, D. S. Marks, and C. Sander. The microrna.org resource: targets and expression. *Nucleic Acids Res*, 36(Database issue):D149-D153, Jan 2008.

S. Griffiths-Jones, H. K. Saini, S. van Dongen, and A. J. Enright. miRBase: tools for microRNA genomics. *Nucleic Acids Res*, 36(Database issue):D154-D158, Jan. 2008.

X. Wang and I. M. E. Naqa. Prediction of both conserved and nonconserved microrna targets in animals. *Bioinformatics*, 24(3):325-332, Feb 2008.

G. L. Papadopoulos, M. Reczko, V. A. Simossis, P. Sethupathy, and A. G. Hatzigeorgiou. The database of experimentally supported targets: a functional update of tarbase. *Nucleic Acids Res*, 37(Database issue):D155-D158, Jan 2009.

get_potential_swaps Calculate swap partner for TF-/miRNA-target graph.

Description

Calculates for TF-/miRNA-target graph all potential swap partner.

Usage

get_potential_swaps(genesetsTF=NULL, genesetsmiRNA=NULL, perc.overlap.cutoff = 0.8, integer.id=TR

Arguments

genesetsTF	Each entry corresponds to a TF and contains its target genes.
genesetsmiRNA	Each entry corresponds to a miRNA and contains its target genes.
perc.overlap.cu	utoff
	Percentage cutoff of minimal overlap between two miRNAs or TFs to be possi-
	ble swap partner.
integer.id	If TRUE, the swap partner are not outputted as characters, but as integer indices.
verbose	print process or not.

Value

The function returns a list, with the following entries:

T_potential_swaps

Contains for each TF all potential swap partner.

S_potential_swaps

Contains for each miRNA all potential swap partner.

Author(s)

Benedikt Zacher (zacher@lmb.uni-muenchen.de)

Examples

```
genesetsTF = list("V$1"=c("A", "B", "C", "D", "E"), "V$2"=c("A", "B", "C", "D"), "V$3"=c("A", "B", "C"))
genesetsmiRNA =list("miR-1"=c("C", "E", "D", "G", "H"), "miR-2"=c("C", "E", "D", "G"), "miR-3"=c("C", "E"))
get_potential_swaps(genesetsTF, genesetsmiRNA, integer.id=FALSE, perc.overlap.cutoff=0.7)
```

limmaAnalysis Perform a limma analysis on expression data.

Description

Needed for the parameterization of the model as well as the Fisher (pre-)test.

Usage

```
limmaAnalysis(dat, design, contrasts)
```

Arguments

dat	A matrix or ExpressionSet containing the expression values.
design	A design matrix.
contrasts	Contrast for the linear model.

Value

Returns a list containing the following entries:

pvalue.tab	Containing the result of the topTable function from limma.
lm.fit	Linear fit to the model.
design	The design used.
contrast	The contrasts used.

Author(s)

Benedikt Zacher (zacher@lmb.uni-muenchen.de), Holger Frohlich

References

G. K. Smyth. Limma : Linear Models for Microarray Data. Bioinformatics, (2005):397-420.

See Also

lmFit, topTable

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limmaAnalysis-methods

Examples

```
data(humanSim)
design = model.matrix(~0+factor(c(rep("control", 5), rep("treated", 5))))
colnames(design) = c("control", "treated")
contrasts = "treated - control"
limmamRNA = limmaAnalysis(sim$dat.mRNA, design, contrasts)
```

limmaAnalysis-methods Methods for Function limmaAnalysis in Package birta

Description

Generic methods for function limmaAnalysis in package **birta**. The expression data can be stored in a matrix or an ExpressionSet.

Methods

```
signature(dat = "ExpressionSet", design = "matrix", contrasts = "character") Generic
function for objects of class ExpressionSet.
```

```
signature(dat = "matrix", design = "matrix", contrasts = "character") Generic func-
tion for objects of class matrix.
```

plotConvergence *Plotting the likelihood along MCMC sampling.*

Description

Plots the log likelihood along MCMC sampling.

Usage

```
plotConvergence(res, nburnin=NULL, title="")
```

Arguments

res	The result from birta.run (a list).
nburnin	Number of iterations used for the burn in.
title	Optional title of the plot.

Author(s)

Benedikt Zacher <zacher@lmb.uni-muenchen.de>

See Also

birta

Examples

```
data(humanSim)
data(humanSim)
design = model.matrix(~0+factor(c(rep("control", 5), rep("treated", 5))))
colnames(design) = c("control", "treated")
contrasts = "treated - control"
limmamRNA = limmaAnalysis(sim$dat.mRNA, design, contrasts)
limmamiRNA = limmaAnalysis(sim$dat.miRNA, design, contrasts)
sim_result = birta(sim$dat.mRNA, sim$dat.miRNA, limmamRNA=limmamRNA,
limmamiRNA=limmamiRNA, nrep=c(5,5,5,5), genesets=genesets,
model="all-plug-in", niter=50000, nburnin=10000,
sample.weights=FALSE, potential_swaps=potential_swaps)
plotConvergence(sim_result, nburnin=10000, title="simulation")
```

potential_swaps Potential swap moves for TF-target and miRNA-target networks (see genestes data set).

Description

Potential swap moves for TF-target and miRNA-target networks (see genestes data set) used for the simulation in the vignette.

Usage

potential_swaps

Format

A list containing potential swap moves for the TF- and miRNA-target graph used in the simulation.

sim

Simulated expression data for mRNAs and miRNAs.

Description

This data set contains simulated expression values of miRNAs and mRNAs, together with the associated TF- and miRNA-target networks.

Usage

sim

Format

A list containing the entries dat.mRNA (mRNA expression), dat.miRNA (miRNA expression), miRNAstates (activity states of miRNAs), TFstates (activity states of TFs, TRANSFAC), gene-setsmiRNA (miRNA-target graph, targetScan), genesetsTFs (TF-taget graph).

sim

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TFexpr

Source

Simulation (see reference for details).

References

B. Zacher, K. Abnaof, S. Gade, E. Younesi, A. Tresch, and H. Frohlich. Joint Bayesian Inference of Condition Specific miRNA and Transcription Factor Activities from Combined Gene and microRNA Expression Data. *submitted*, 2012.

TFexpr	Transcription factor	expression	values j	for	the	aerobic-anaerobic
	growth experiment.					

Description

This data set gives expression values for the 160 TF of the TF-target graph EColiNetwork used in the vignette.

Usage

TFexpr

Format

ExpressionSet. Rownames in the assayData correspond to entries in TF-target graph.

Source

See EColiOxygen and EColiNetwork (see reference for details).

References

M. W. Covert, E. M. Knight, J. L. Reed, M. J. Herrgard, and B. O. Palsson. Integrating high-throughput and computational data elucidates bacterial networks. *Nature*, 429(6987):92-96, May 2004.

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