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Title Network Reconstruction and Changepoint Detection

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Description Package EDISON (Estimation of Directed Interactions from Sequences Of Non-homogeneous gene expression) runs an MCMC simulation to reconstruct networks from time series data, using a non-homogeneous, time-varying dynamic Bayesian network. Networks segments and changepoints are inferred concurrently, and information sharing priors provide a reduction of the inference uncertainty.

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EDISON-package

Allows for network reconstruction and changepoint detection.

Description

This package runs an MCMC simulation to reconstruct networks from time series data, using a nonhomogeneous, time-varying dynamic Bayesian network. Networks segments and changepoints are inferred concurrently, and information sharing priors provide a reduction of the inference uncertainty.

Details

Package:	EDISON
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Author(s)

Frank Dondelinger, Sophie Lebre

Maintainer: Frank Dondelinger <fdondelinger.work@gmail.com>

References

Dondelinger et al. (2012), "Non-homogeneous dynamic Bayesian networks with Bayesian regularization for inferring gene regulatory networks with gradually time-varying structure", Machine Learning.

Husmeier et al. (2010), "Inter-time segment information sharing for non-homogeneous dynamic Bayesian networks", NIPS.

See Also

corpcor

Examples

```
# Generate random gene network and simulate data from it
dataset = simulateNetwork(l=25)
# Run MCMC simulation to infer networks and changepoint locations
result = EDISON.run(dataset$sim_data, num.iter=500)
# Calculate posterior probabilities of changepoints
cps = calculateCPProbabilities(result)
# Calculate marginal posterior probabilities of edges in the network
network = calculateEdgeProbabilities(result)
```

AcceptableMove Check if move is acceptable.

Description

This function takes as input a new network proposal and checks that the proposal does not exceed the maximum number of parents for a node, and that there are no self loops (if self loops have been disallowed).

Usage

```
AcceptableMove(proposal, qmax, self.loops, target, fixed.edges)
```

Arguments

proposal	The proposed network (K-by-q matrix with K segments and q parent sets).
qmax	Maximum number of parents allowed.
self.loops	Flag indicating whether self loops are allowed.
target	The current target node (only needed to find out which parent would be the self loop).
fixed.edges	Which edges in the network should be fixed for all segments (q-by-q matrix with entries 0 for fixed non-edge, 1 for fixed edge, -1 for non-fixed edge).

Value

Returns TRUE if the proposed move is acceptable, FALSE otherwise.

Author(s)

Frank Dondelinger

See Also

make_structure_move

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addProposalNetworkInfo

Add the proposed new network to the new.nets list.

Description

Updates the network.info data structure so that it stays consistent.

Usage

addProposalNetworkInfo(network.info, newS, E)

Arguments

network.info	Data structure containing the current network.
newS	Proposed new network for this target, a num.local.segs by num.parents matrix.
Е	The current vector of local segments for this target (only used to check for con- sistency with the network.info change points).

Value

Updated network.info data structure, with new network added to new.nets.

Author(s)

Frank Dondelinger

BinoHyperMove Makes a binomial hyperparameter move.

Description

This function proposes a move for one of the hyperparameters of the binomial prior, calculates the acceptance probability and accepts the move accordingly.

Usage

BinoHyperMove(network.info, node.sharing, GLOBvar)

network.info	The collected network information obtained using CollectNetworkInfo.
node.sharing	Which type of node sharing is used, either 'soft' or 'hard' sharing.
GLOBvar	Global variables of the MCMC.

Value

Returns a list with elements:

move	The move type (in this case, 2).
move.made	1 if the move was proposed, 0 otherwise.
network.info	The network information, including the new hyperparameters if the move was accepted.
accept	Whether the move was accepted or not.

Author(s)

Frank Dondelinger

References

For information about the binomial information sharing prior, see:

Husmeier et al. (2010), "Inter-time segment information sharing for non-homogeneous dynamic Bayesian networks", NIPS.

Dondelinger et al. (2012), "Non-homogeneous dynamic Bayesian networks with Bayesian regularization for inferring gene regulatory networks with gradually time-varying structure", Machine Learning.

See Also

BinoHyperMove

BinoHyperRatio Calculates the MH ratio of the binomial prior.

Description

This function calculates the ratio of the binomial information sharing prior with the proposed new hyperparameter values, and the binomial prior with the current hyperparameter values.

Usage

```
BinoHyperRatio(params.proposed, changed, node.sharing, network.info)
```

params.proposed		
	The new proposed hyperparameter values for the binomial prior.	
changed	Gives the index of the parameter that has changed.	
node.sharing	Type of information sharing among nodes: 'soft' or 'hard'.	
network.info	The network information as collected by CollectNetworkInfo.	

Value

This function returns a number greater than zero which represents the ratio of binomial priors.

Author(s)

Frank Dondelinger

References

For information about the binomial information sharing prior, see:

Husmeier et al. (2010), "Inter-time segment information sharing for non-homogeneous dynamic Bayesian networks", NIPS.

Dondelinger et al. (2012), "Non-homogeneous dynamic Bayesian networks with Bayesian regularization for inferring gene regulatory networks with gradually time-varying structure", Machine Learning.

See Also

BinoHyperMove

bp.computeAlpha Computes the acceptance ratio of two changepoint configurations.

Description

This function computes the acceptance ratio of two changepoint configurations with networks in a changepoint birth or death move.

Usage

```
bp.computeAlpha(birth, lNew, kminus, Ekl, Estar, Ekr, yL, PxL, yR, PxR, y2, Px2,
D, delta2, q, smax, v0, gamma0, prior_ratio = 1)
```

birth	1 for a changepoint birth move, -1 for a changepoint death move.
1New	Number of edges in the new segment.
kminus	Minimal number of changepoints between the two compared models (equal to s for a birth move, s-1 for a death move.
Ekl	Changepoint on the left of proposed changepoint.
Estar	Changepoint being inserted or deleted.
Ekr	Changepoint on the right of proposed changepoint.
уL	Response data (left).
PxL	Projection matrix (left).

уR	Response data (right).
PxR	Projection matrix (right).
y2	Response data (both).
Px2	Projection matrix (both).
D	Hyperparameters for the number of edges in each segment.
delta2	Hyperparameters for the empirical covariance (signal-to-noise ratio).
q	Total number of nodes in the network.
smax	Maximum number of changepoints.
v0	Hyperparameter.
gamma0	Hyperparameter.
prior_ratio	Ratio of network structure priors.

Author(s)

Sophie Lebre

References

For more information about the model, see:

Dondelinger et al. (2012), "Non-homogeneous dynamic Bayesian networks with Bayesian regularization for inferring gene regulatory networks with gradually time-varying structure", Machine Learning.

See Also

cp.birth, cp.death

buildXY

Builds response Y and predictor X.

Description

This function builds the response variables Y and predictor variables X from the input data.

Usage

buildXY(targetData, predData, GLOBvar)

targetData	Target input data.
predData	Predictor input data.
GLOBvar	Global variables of the MCMC simulation.

CalculateChanges

Value

A list with elements:

Х	Predictor variables.
Υ	Response variables.

Author(s)

Sophie Lebre

CalculateChanges Function to calculate the number of differences between adjaccent network segments.

Description

This function takes the current network structure, compares each segment to the next one, and calculates the number of changes. If soft information sharing among nodes is active, then this procedure is only done for the current target node.

Usage

CalculateChanges(network.info, node.sharing)

Arguments

network.info	The network information collected by function CollectNetworkInfo.
node.sharing	Specifies the type of information sharing among nodes: 'soft' or 'hard'

Value

Returns a vector with 4 elements: the number of coinciding edges, the number of edges in the previous segment that are absent in the next one, the number of edges in the next segment that are absent in the previous one and the number of coinciding non-edges.

Author(s)

Frank Dondelinger

calculateCPPGlobal Calculated the global changepoint probabilities.

Description

This function calculates the global probability of a changepoint at each measured timepoint, using the node-specific probabilities.

Usage

```
calculateCPPGlobal(prob.cps)
```

Arguments

prob.cps Node-specific changepoint probabilities, a NumNodes by NumTimepoints matrix.

Value

A matrix of length 1 by NumTimepoints, containing the global changepoint probabilities.

Author(s)

Frank Dondelinger

See Also

calculateCPProbabilities

calculateCPProbabilities

Calculate the changepoint probabilities.

Description

This function calculates the marginal changepoint probabilities from the changepoint samples taken during the MCMC simulation.

Usage

calculateCPProbabilities(network.samples)

Arguments

network.samples

List of network and changepoint samples collected during the MCMC simulation by EDISON.run and runDBN.

Value

Returns a matrix of dimension NumNodes by NumTimePoints, where each entry contains the marginal posterior probability of a changepoint for that node at that timepoint.

Author(s)

Frank Dondelinger

Examples

```
# Generate random gene network and simulate data from it
dataset = simulateNetwork()
# Due MCMC simulation to infer networks and sharesensitt less
```

Run MCMC simulation to infer networks and changepoint locations result = EDISON.run(dataset\$sim_data, num.iter=500)

```
# Calculate posterior probabilities of changepoints
cps = calculateCPProbabilities(result)
```

calculateEdgeProbabilities

Calculate the edge probabilities.

Description

This function calculates the marginal posterior probabilities of the edges in the network segments, for each timepoint, and optionally calculates the same for specified changepoints.

Usage

```
calculateEdgeProbabilities(network.samples, cps = NULL)
```

Arguments

network.samples

	Network samples obtained from the MCMC simulation using EDISON.run and runDBN.
cps	Optionally specifies changepoints to allow for calculating the marginal posterior edge probabilities for specific segments.

Value

A list with elements:

probs.all	A list containing marginal edge posterior probabilities for each timepoint.
probs.segs	A list containing marginal edge posterior probabilities for each specified seg
	ment.

Author(s)

Frank Dondelinger

See Also

calculateEdgeProbabilitiesTimePoints, calculateEdgeProbabilitiesSegs

Examples

```
# Generate random gene network and simulate data from it
dataset = simulateNetwork(1=25)
```

Run MCMC simulation to infer networks and changepoint locations
result = EDISON.run(dataset\$sim_data, num.iter=500)

```
# Calculate marginal posterior probabilities of edges in the network
network = calculateEdgeProbabilities(result)
```

```
# Calculate marginal posterior probabilities of edges in the network,
# using the true changepoints
true.cps = c(2,dataset$epsilon)
network = calculateEdgeProbabilities(result, cps=true.cps)
```

calculateEdgeProbabilitiesSegs Calculate edge probabilities for fixed segments.

Description

This function calculates the marginal posterior probabilities for the edges in each network for the specified segments.

Usage

```
calculateEdgeProbabilitiesSegs(prob.networks, cps, numNodes)
```

prob.networks	List containing the marginal posterior probabilities for the edges of each network at each timepoint, from calculateEdgeProbabilitiesTimePoints.
cps	Changepoints defining the segments for which the edge probabilities should be calculated. Note that these are global changepoints that apply to the whole network.
numNodes	Number of nodes in the network.

Value

Returns a list of length equal to the number of segments, with each entry containing a matrix of size NumNodes by NumNodes which contains the marginal edge probabilities for that segment.

Author(s)

Frank Dondelinger

See Also

calculateEdgeProbabilities, calculateEdgeProbabilitiesTimePoints

calculateEdgeProbabilitiesTimePoints Calculate the edge posterior probabilities for each timepoint.

Description

This function calculates the marginal posterior edge probabilities of the network at each timepoint.

Usage

```
calculateEdgeProbabilitiesTimePoints(network.samples, cps, numNodes)
```

Arguments

network.samples

	Collection of network and changepoint samples of the MCMC simulation, as
	obtained by EDISON.run, runDBN.
cps	Changepoint vector.
numNodes	Number of nodes in the network.

Value

A list of length equal to the number of timepoints, where each entry contains a matrix of size NumNodes by NumNodes with the marginal posterior edge probabilities of the network at this timepoint.

Author(s)

Frank Dondelinger

See Also

calculateEdgeProbabilities, calculateEdgeProbabilitiesSegs

```
CalculateLikelihoodRatio
```

Calculates the ratio of two likelihoods in a structure move.

Description

This function calculates the ratio of the liklihoods in a network structure move. The returned value is the ratio for the modification of one edge in one segment.

Usage

```
CalculateLikelihoodRatio(gamma0, y, Pxlm, Pxl, v0, delta2, dir)
```

Arguments

gamma0	Hyperparameter.
У	Target data.
Pxlm	Projection matrix with modified edge.
Pxl	Original projection matrix.
v0	Hyperparameter.
delta2	Delta squared parameter (signal-to-noise).
dir	Direction of the change: $1 = Added$ an edge. $2 = Removed$ an edge. $0 = No$ change.

Value

Returns the likelihood ratio.

Author(s)

Frank Dondelinger

References

For more information about the hyperparameters and the functional form of the likelihood, see:

Dondelinger et al. (2012), "Non-homogeneous dynamic Bayesian networks with Bayesian regularization for inferring gene regulatory networks with gradually time-varying structure", Machine Learning.

See Also

CalculatePriorRatio

CalculatePriorRatio Calculates the network prior ratio.

Description

This function calculates the ratio of the network structure priors for a structure move.

Usage

CalculatePriorRatio(method, q, lambda, network.info)

Arguments

method	Indicates which prior to use: 'poisson' for the standard Poisson prior (no in- formation sharing), 'exp_soft' or 'exp_hard' for the exponential informa- tion sharing prior with soft or hard sharing among nodes and 'bino_soft' or 'bino_hard' for the binomial information sharing prior with soft or hard shar- ing among nodes.
q	Number of nodes in the network.
lambda	Vector of lambda hyperparameter values for each network (needed for the Poisson prior).
network.info	The network information collected using CollectNetworkInfo.

Value

Returns the ratio of the network structure priors for the proposed structure move.

Author(s)

Frank Dondelinger

References

For more information on the network structure priors, see:

Husmeier et al. (2010), "Inter-time segment information sharing for non-homogeneous dynamic Bayesian networks", NIPS.

Dondelinger et al. (2012), "Non-homogeneous dynamic Bayesian networks with Bayesian regularization for inferring gene regulatory networks with gradually time-varying structure", Machine Learning.

See Also

CalculateLikelihoodRatio

CollectNetworkInfo Collects all the network information in one list.

Description

This function collects information about the current network segments and hyperparameters for the information sharing priors.

Usage

CollectNetworkInfo(Sall, Eall, prior.params, posPhase, target, q, self.loops, k)

Arguments

Sall	Structure of all segments. A list of length q, where each element is a K_i by q matrix containing the parents for the current node in each of the K_i segments.
Eall	Positions of segment boundaries. A list of length q, where each element is a vector containing the segment boundaries for the current parent node.
prior.params	The hyperparameters of the information sharing prior (if applicable).
posPhase	The segment being changed.
target	The target parent node whose edge is being changed.
q	The total number of nodes in the network.
self.loops	Whether self-loops are allowed in the network.
k	The level-2 hyperparameter for the exponential prior.

Value

The function returns a list with the following elements:

nets	The structure of all segments, a list of length K where K is the total number of segments over all nodes.
segment	Identical to posPhase.
target.nets	Identical to Sall.
prior.params	Identical to prior.params.
self.loops	Identical to self.loops.
k	Identical to k.
new.nets	Dummy variable for holding the proposed network in a network structure move. Originally identical to variable nets.

Author(s)

Frank Dondelinger

computePx

Description

This function computes the projection matrix that is needed for calculation of the likelihood.

Usage

computePx(len, x, delta2)

Arguments

len	Delimiting breakpoints.
x	The observations of x in the corresponding state.
delta2	Signal-to-noise ratio hyperparameter.

Value

The projection matrix.

Author(s)

Sophie Lebre

References

For more information about the hyperparameters and the functional form of the likelihood, see:

Dondelinger et al. (2012), "Non-homogeneous dynamic Bayesian networks with Bayesian regularization for inferring gene regulatory networks with gradually time-varying structure", Machine Learning.

See Also

CalculateLikelihoodRatio

computeRho4

Description

This function calculates the frequency at which each of the different changepoint moves is proposed. For the poisson network structure prior, this ensures that the proposal frequency is equal to the prior probability.

Usage

computeRho4(k, kmin, kmax, c, lambda)

Arguments

k	The number of hidden states.
kmin	Minimum number of hidden states.
kmax	Maximum number of hidden states
с	Parameter.
lambda	Hyperparameter controlling the number of hidden states.

Value

Vector containing the proposal frequencies for the different changepoint moves.

Author(s)

Sophie Lebre

References

For more information about the hyperparameters and the functional form of the likelihood, see:

Dondelinger et al. (2012), "Non-homogeneous dynamic Bayesian networks with Bayesian regularization for inferring gene regulatory networks with gradually time-varying structure", Machine Learning. convert_nets

Description

Converts from representing the network as a list of target nodes to representing it as a list of segments.

Usage

convert_nets(Ball, Eall)

Arguments

Ball	Input network: List of target nodes, where each element is a NumSegs by NumNodes matrix giving the parents for the target node in each segment.
Eall	Changepoints: List of target nodes, where each element contains a vector of changepoints.

Value

List with elements:

B_nets	List of segments, where each element contains a matrix of size NumNodes by
	NumNodes, representing the network for that segment.
segs	Vector containing the global segment boundaries.

Author(s)

Frank Dondelinger

cp.birth

Make changepoint birth move.

Description

This function makes a changepoint birth move, possibly adding a changepoint.

Usage

```
cp.birth(Eall, Sall, Ball, Sig2all, X, Y, D, GLOBvar, HYPERvar, target)
```

Arguments

Eall	Changepoints: List of target nodes, where each element contains a vector of changepoints.
Sall	Network structure: List of target nodes, where each element is a NumSegs by NumNodes matrix giving the parents for the target node in each segment. A binary matrix.
Ball	Network parameters: Similar to network structure, but with regression parameters included.
Sig2all	Sigma squared parameters.
Х	Response data.
Υ	Target data.
D	Hyperparameter.
GLOBvar	Global variables of the MCMC simulation.
HYPERvar	Hyperparameter variables.
target	Which target node the move is being proposed for.

Value

A list with elements:

E	New changepoint vector for target node.
Sall	Updated network structure.
Ball	Updated network structure with regression parameters.
Sig2all	Udated sigma squared.
prior.params	Updated vector of structure prior hyperparameters.
accept	Whether the move was accepted or not.
move	What type of move was made. In this case move=1 for a changepoint birth move.
alpha	The acceptance ratio of the move.
estar	The location of the new changepoint.
k	Hyperparameter.

Author(s)

Sophie Lebre Frank Dondelinger

References

For more information about the different changepoint moves, see:

Dondelinger et al. (2012), "Non-homogeneous dynamic Bayesian networks with Bayesian regularization for inferring gene regulatory networks with gradually time-varying structure", Machine Learning.

cp.death

See Also

cp.death, cp.shift

cp.death

Make changepoint death move.

Description

This function makes a changepoint death move, possibly removing a changepoint.

Usage

```
cp.death(Eall, Sall, Ball, Sig2all, X, Y, D, GLOBvar, HYPERvar, target)
```

Arguments

Eall	Changepoints: List of target nodes, where each element contains a vector of changepoints.
Sall	Network structure: List of target nodes, where each element is a NumSegs by NumNodes matrix giving the parents for the target node in each segment. A binary matrix.
Ball	Network parameters: Similar to network structure, but with regression parameters included.
Sig2all	Sigma squared parameters.
Х	Response data.
Υ	Target data.
D	Hyperparameter.
GLOBvar	Global variables of the MCMC simulation.
HYPERvar	Hyperparameter variables.
target	Which target node the move is being proposed for.

Value

A list with elements:

E	New changepoint vector for target node.
Sall	Updated network structure.
Ball	Updated network structure with regression parameters.
Sig2all	Updated sigma squared.
prior.params	Updated vector of structure prior hyperparameters.
accept	Whether the move was accepted or not.
move	What type of move was made. In this case move=2 for a changepoint death
	move.
alpha	The acceptance ratio of the move.
estar	The location of the removed changepoint.
k	Hyperparameter.

Author(s)

Sophie Lebre

Frank Dondelinger

References

For more information about the different changepoint moves, see:

Dondelinger et al. (2012), "Non-homogeneous dynamic Bayesian networks with Bayesian regularization for inferring gene regulatory networks with gradually time-varying structure", Machine Learning.

See Also

cp.birth, cp.shift

cp.shift

Makes a changepoint shift move.

Description

This function makes a changepoint shift move, possibly moving one of the changepoints.

Usage

cp.shift(Eall, Sall, Ball, Sig2all, X, Y, GLOBvar, HYPERvar, target)

Arguments

Eall	Changepoints: List of target nodes, where each element contains a vector of changepoints.
Sall	Network structure: List of target nodes, where each element is a NumSegs by NumNodes matrix giving the parents for the target node in each segment. A binary matrix.
Ball	Network parameters: Similar to network structure, but with regression parameters included.
Sig2all	Sigma squared parameters.
Х	Response data.
Υ	Target data.
GLOBvar	Global variables of the MCMC simulation.
HYPERvar	Hyperparameter variables.
target	Which target node the move is being proposed for.

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defaultOptions

Value

A list with elements:

E	New changepoint vector for target node.
Sall	Updated network structure.
Ball	Updated network structure with regression parameters.
Sig2all	Updated sigma squared.
prior.params	Updated vector of structure prior hyperparameters.
accept	Whether the move was accepted or not.
move	What type of move was made. In this case move=2 for a changepoint death move.
alpha	The acceptance ratio of the move.
estar	The location of the removed changepoint.
k	Hyperparameter.

Author(s)

Sophie Lebre

References

For more information about the different changepoint moves, see:

Dondelinger et al. (2012), "Non-homogeneous dynamic Bayesian networks with Bayesian regularization for inferring gene regulatory networks with gradually time-varying structure", Machine Learning.

See Also

cp.birth, cp.death

defaultOptions Set the default options for the MCMC simulation.

Description

This function creates a list with the default options of the MCMC simulation.

Usage

defaultOptions()

Value

A list of default options with elements:

lmax	Maximum number of parent nodes. Default=5.
m	Number of repeated measurements. Default=1 (no repeats).
dyn	Lag for the DBN model. Default = 1 when $X(t)$ depends on the previous measurement $X(t-1)$, but dyn can be chosen equal to 2, 3,
minPhase	Minimal length of a segment. Default=2.
maxCP	Maximal number of changepoints. Default=10.
maxTF	Maximal number of incoming edges for each node. Default=5.
alphaCP	Hyperparameter for the number of changepoints. Default=1.
betaCP	Hyperparameter for the number of changepoints. Default=0.5.
alphaTF	Hyperparameter for the number of incoming edges. Default=1.
betaTF	Hyperparameter for the number of incoming edges. Default=0.5.
burnin	Whether to include a burnin period. Default=F.
psrf.check	Whether to calculate the potential scale reduction factor (PSRF). Default=F.
pp.11	Proposal frequency for level-1 hyperparameter moves. Default=0.2.
pp.12	Proposal frequency for level-2 hyperparameter moves. Default=0.01.
save.by.node	Whether to save results separately for each target node. Default=F.
save.file	Whether to save the results to a file. Default=F.
hyper.fixed	Whether to keep the network structure prior hyperparameters fixed. Default=F.
cp.fixed	Whether to keep the changepoints fixed. Default=F.
hyper.init	Initial values for the network structure prior hyperparameters. Default=NULL.
cp.init	Initial values for the changepoint locations. Default=NULL.

Author(s)

Frank Dondelinger

Examples

```
# Set options to allow saving network and changepoint samples to file
options = defaultOptions()
options$save.file = TRUE
# NOT EXECUTED
# result.bino2 = EDISON.run(dataset$sim_data,
# information.sharing='bino_hard',
# num.iter=5000, output.file='bino2.results',
# options=options)
```

dinvgamma

Description

This function calculates the density of the inverse gamma distribution.

Usage

```
dinvgamma(x, shape, scale = 1, log = FALSE)
```

Arguments

х	Input.
shape	Shape parameter.
scale	Scale parameter (1/rate).
log	Whether to return the log density.

Value

Returns the density (or log density).

Author(s)

Frank Dondelinger

Examples

```
# Draw samples from inverse gamma distribution with shape parameter 1
# and scale parameter 1
samples = rinvgamma(100, shape=1, scale=1)
```

```
# Calculate density of samples
densities = dinvgamma(samples, shape=1, scale=1)
```

EDISON.run

Wrapper function for starting an MCMC simulation

Description

This function provides a wrapper for starting an MCMC simulation, using only the data and some basic options as input.

Usage

```
EDISON.run(input, output.file = "EDISON.output",
information.sharing = "poisson", num.iter = 10000, prior.params = NULL,
options = NULL, fixed.edges = NULL)
```

Arguments

input	Input data. Either a filename pointing to an R data file containing the results of simulateNetwork, or a NumTimePoints by NumNodes matrix.	
output.file	Where to save the output of the MCMC simulation.	
information.sharing		
	Which information sharing prior to use: 'poisson' for the Poisson prior (no information sharing), 'exp_hard' or 'exp_soft' for the exponential prior with hard or soft coupling among nodes, respectively, and 'bino_hard' or 'bino_soft' for the binomial prior with hard or soft coupling among nodes.	
num.iter	Number of iterations of the MCMC simulation.	
prior.params	Initial values of the hyperparameters of the information sharing priors.	
options	Settings for the MCMC simulation, as generated by defaultOptions.	
fixed.edges	Matrix of size NumNodes by NumNodes, with fixed.edges[i,j]==1 $ 0$ if the edge between nodes i and j is fixed, and -1 otherwise. Defaults to NULL (no edges fixed).	

Value

Returns the results of the MCMC simulation, similar to runDBN.

Author(s)

Sophie Lebre Frank Dondelinger

References

For details on the model and MCMC simulation, see:

Dondelinger et al. (2012), "Non-homogeneous dynamic Bayesian networks with Bayesian regularization for inferring gene regulatory networks with gradually time-varying structure", Machine Learning.

See Also

runDBN

Examples

```
# Generate random gene network and simulate data from it
dataset = simulateNetwork(1=25)
```

Run MCMC simulation to infer networks and changepoint locations

Uses default settings: Poisson prior and 1500 iterations

ExpHyperMove

```
result.poisson = EDISON.run(dataset$sim_data, num.iter=500)
# Use the binomial information sharing prior with hard node coupling, and
# run for 5000 iterations
# NOT EXECUTED
#result.bino = EDISON.run(dataset$sim_data,
                 information.sharing='bino_hard', num.iter=5000)
#
# Set options to allow saving network and changepoint samples to file
options = defaultOptions()
options$save.file = TRUE
# NOT EXECUTED
# result.bino2 = EDISON.run(dataset$sim_data,
#
                   information.sharing='bino_hard',
                   num.iter=5000, output.file='bino2.results',
#
#
                   options=options)
```

```
ExpHyperMove
```

Makes an exponential hyperparameter move.

Description

This function tries to make a level-1 or level-2 hyperparameter move for the exponential prior

Usage

ExpHyperMove(network.info, node.sharing, GLOBvar, hyper.proposals)

Arguments

network.info	The network information collected by CollectNetworkInfo.	
node.sharing	The type of information sharing among nodes: <code>'soft'</code> or <code>'hard'</code> .	
GLOBvar	Collection of global variables of the MCMC.	
hyper.proposals		
	Proposal width of the hyperparameter move.	

Value

Returns a list with elements:

move.made	1 if a level-1 hyperparameter move has been made, 0 otherwise.
network.info	Network information with updated hyperparameters if the move was accepted.
accept	Whether a level-1 hyperparameter move has been accepted or not.
move.made.k	1 if a level-2 hyperparameter move has been made, 0 otherwise.
accept.k	Whether a level-2 hyperparameter move has been accepted or not.
move	Type of move: 2 for a level-1 hyperparameter move, 3 for a level-2 hyperparam-
	eter move.

Author(s)

Frank Dondelinger

References

For information about the exponential information sharing prior, see:

Husmeier et al. (2010), "Inter-time segment information sharing for non-homogeneous dynamic Bayesian networks", NIPS.

Dondelinger et al. (2012), "Non-homogeneous dynamic Bayesian networks with Bayesian regularization for inferring gene regulatory networks with gradually time-varying structure", Machine Learning.

See Also

ExpHyperRatioTarget

ExpHyperRatioTarget Calculates the ratio of an exponential hyperparameter move.

Description

This function calculates the acceptance ratio of a level-1 hyperparameter move for a given target node.

Usage

```
ExpHyperRatioTarget(beta.proposed, beta.old, target.net, self.loops)
```

Arguments

beta.proposed	Proposed new hyperparameter value.
beta.old	Previous value of hyperparameter beta.
target.net	Network segments for the target node associated with this hyperparameter value.
self.loops	'TRUE' if self-loops are acceptable, 'FALSE' otherwise.

Value

Returns the ratio of the exponential prior with the previous hyperparameter value and the proposed new hyperparameter value.

Author(s)

Frank Dondelinger

fix_eigenvalues

References

For information about the exponential information sharing prior, see:

Husmeier et al. (2010), "Inter-time segment information sharing for non-homogeneous dynamic Bayesian networks", NIPS.

Dondelinger et al. (2012), "Non-homogeneous dynamic Bayesian networks with Bayesian regularization for inferring gene regulatory networks with gradually time-varying structure", Machine Learning.

See Also

ExpHyperMove

fix_eigenvalues Modify network to ensure stationarity.

Description

This function ensures that the eigenvalues of the network structure matrix are smaller or equal to 1, thereby ensuring stationarity of the regression. This is done by removing edges at random until the condition is satisfied.

Usage

fix_eigenvalues(network, q, gauss_weights)

Arguments

network	Original network structure, a matrix of size NumNodes by NumNodes.
q	Number of nodes.
gauss_weights	If TRUE, use Gaussian regression weight, if FALSE conserve original weights.

Value

Returns a network with fewer eigenvalues than the original network, but satisfying the stationarity condition.

Author(s)

Frank Dondelinger

See Also

generateNetwork

generateNetwork

Description

This function generates a random network with changepoints for structure changes, for simulating synthetic data.

Usage

```
generateNetwork(lambda_2 = 0.45, q = 10, min_phase_length = 1,
k_bar = 5, l = 10, lambda_3 = 2, spacing = 1, gauss_weights = TRUE,
same = FALSE, change_method = "sequential", fixed = FALSE, cps = NULL)
```

Arguments

qNumber of nodes.min_phase_lengtbMinimum segment length.k_barMaximum number of changepoints. If fixed=TRUE, this is equal to the number of changepoints.1Length of the time series.lambda_3Average number of structure changes between two segments (parameter for a Poisson distribution).spacing1 if segments are equally spaced, 0 if they are spaced randomly (subject to the constraints of min_phase_length).gauss_weights1 if edge weights in the network are drawn from N(0, 1), 0 if they are fixed to be 1.same1 if all segments have the same network structure (no changes), 0 otherwise.change_method'sequential' if the changes happen sequentially (i.e. changes at segment i are applied to segment i-1), 'hierarchical' if the changes happen with respect to a hypernetwork (i.e. changes at segment i are applied to segment 0).fixedT if the changepoint locations are fixed, F if they should be sampled.cpsChangepoint locations (if they are fixed).	lambda_2	Average number of parents for each node in the network (parameter for a Poisson distribution).
Minimum segment length.k_barMaximum number of changepoints. If fixed=TRUE, this is equal to the number of changepoints.1Length of the time series.lambda_3Average number of structure changes between two segments (parameter for a Poisson distribution).spacing1 if segments are equally spaced, 0 if they are spaced randomly (subject to the constraints of min_phase_length).gauss_weights1 if edge weights in the network are drawn from N(0, 1), 0 if they are fixed to be 1.same1 if all segments have the same network structure (no changes), 0 otherwise.change_method'sequential' if the changes happen sequentially (i.e. changes at segment i are applied to segment i-1), 'hierarchical' if the changes happen with respect to a hypernetwork (i.e. changes at segment i are applied to segment 0).fixedT if the changepoint locations are fixed, F if they should be sampled.	q	Number of nodes.
k_barMaximum number of changepoints. If fixed=TRUE, this is equal to the number of changepoints.1Length of the time series.lambda_3Average number of structure changes between two segments (parameter for a Poisson distribution).spacing1 if segments are equally spaced, 0 if they are spaced randomly (subject to the constraints of min_phase_length).gauss_weights1 if edge weights in the network are drawn from N(0, 1), 0 if they are fixed to be 1.same1 if all segments have the same network structure (no changes), 0 otherwise.'sequential' if the changes happen sequentially (i.e. changes at segment i are applied to segment i-1), 'hierarchical' if the changes happen with respect to a hypernetwork (i.e. changes at segment i are applied to segment 0).fixedT if the changepoint locations are fixed, F if they should be sampled.	min_phase_lengt	th
 of changepoints. 1 Length of the time series. 1 ambda_3 Average number of structure changes between two segments (parameter for a Poisson distribution). spacing 1 if segments are equally spaced, 0 if they are spaced randomly (subject to the constraints of min_phase_length). gauss_weights 1 if edge weights in the network are drawn from N(0, 1), 0 if they are fixed to be 1. same 1 if all segments have the same network structure (no changes), 0 otherwise. 'sequential' if the changes happen sequentially (i.e. changes at segment i are applied to segment i-1), 'hierarchical' if the changes happen with respect to a hypernetwork (i.e. changes at segment i are applied to segment 0). fixed T if the changepoint locations are fixed, F if they should be sampled. 		Minimum segment length.
 lambda_3 Average number of structure changes between two segments (parameter for a Poisson distribution). spacing 1 if segments are equally spaced, Ø if they are spaced randomly (subject to the constraints of min_phase_length). gauss_weights 1 if edge weights in the network are drawn from N(0, 1), Ø if they are fixed to be 1. same 1 if all segments have the same network structure (no changes), Ø otherwise. 'sequential' if the changes happen sequentially (i.e. changes at segment i are applied to segment i-1), 'hierarchical' if the changes happen with respect to a hypernetwork (i.e. changes at segment i are applied to segment 0). fixed T if the changepoint locations are fixed, F if they should be sampled. 	k_bar	
 Poisson distribution). spacing 1 if segments are equally spaced, Ø if they are spaced randomly (subject to the constraints of min_phase_length). gauss_weights 1 if edge weights in the network are drawn from N(0, 1), Ø if they are fixed to be 1. same 1 if all segments have the same network structure (no changes), Ø otherwise. change_method 'sequential' if the changes happen sequentially (i.e. changes at segment i are applied to segment i-1), 'hierarchical' if the changes happen with respect to a hypernetwork (i.e. changes at segment i are applied to segment 0). fixed T if the changepoint locations are fixed, F if they should be sampled. 	1	Length of the time series.
 constraints of min_phase_length). gauss_weights 1 if edge weights in the network are drawn from N(0, 1), Ø if they are fixed to be 1. same 1 if all segments have the same network structure (no changes), Ø otherwise. change_method 'sequential' if the changes happen sequentially (i.e. changes at segment i are applied to segment i-1), 'hierarchical' if the changes happen with respect to a hypernetwork (i.e. changes at segment i are applied to segment 0). fixed T if the changepoint locations are fixed, F if they should be sampled. 	lambda_3	
 be 1. same 1 if all segments have the same network structure (no changes), Ø otherwise. 'sequential' if the changes happen sequentially (i.e. changes at segment i are applied to segment i-1), 'hierarchical' if the changes happen with respect to a hypernetwork (i.e. changes at segment i are applied to segment 0). fixed T if the changepoint locations are fixed, F if they should be sampled. 	spacing	
 change_method 'sequential' if the changes happen sequentially (i.e. changes at segment i are applied to segment i-1), 'hierarchical' if the changes happen with respect to a hypernetwork (i.e. changes at segment i are applied to segment 0). fixed T if the changepoint locations are fixed, F if they should be sampled. 	gauss_weights	
applied to segment i-1), 'hierarchical' if the changes happen with respect to a hypernetwork (i.e. changes at segment i are applied to segment 0).fixedT if the changepoint locations are fixed, F if they should be sampled.	same	1 if all segments have the same network structure (no changes), Ø otherwise.
	change_method	applied to segment i-1), 'hierarchical' if the changes happen with respect to
cps Changepoint locations (if they are fixed).	fixed	T if the changepoint locations are fixed, F if they should be sampled.
	cps	Changepoint locations (if they are fixed).

Value

A list with the following elements:

network	The network, a list of length NumSegs, where each element is a NumNodes by
	NumNodes matrix.
epsilon	The vector of changepoint locations.
k	The number of changepoint.
changes	The number of changes among segments.

HyperparameterMove

Author(s)

Frank Dondelinger

See Also

simulateNetwork

Examples

```
# Generate random network with default parameters
network = generateNetwork()
# Simulate data using generated network
dataset = simulateNetwork(net=network)
# Generate random network with 4 changepoints and 15 nodes,
# with changepoints distributed over a timeseries of length 50
network = generateNetwork(1=50, q=15, fixed=TRUE, k_bar=4)
# Simulate data of length 50 using generated network
dataset = simulateNetwork(net=network)
```

HyperparameterMove *Make a hyperparameter move.*

Description

This function makes a hyperparameter move for the information sharing prior selected (or no move if no information sharing prior is selected).

Usage

HyperparameterMove(method, network.info, GLOBvar, hyper.proposals)

Arguments

method	The information sharing method used: 'poisson' for the Poisson prior (no information sharing), 'exp_soft' and 'exp_hard' for the exponential information sharing prior with soft or hard information sharing among nodes, respectively, 'bino_soft' and 'bino_hard' for the binomial information sharing prior with soft or hard information sharing among nodes, respectively.
network.info	Network information collected using CollectNetworkInfo.
GLOBvar	Global variables used during the MCMC.
hyper.proposals	
	Proposal width for hyperparameters

Proposal width for hyperparameters.

Value

List summing up the result of the hypermove. Contains at least:

move.made	Whether a hyperparameter move has been made.
network.info	The network information, possibly updated if the hyperparameter move was made and accepted.

May contain further elements depending on the type of information sharing prior used. See the prior-specific functions ExpHyperMove and BinoHyperMove for details.

Author(s)

Frank Dondelinger

References

For information about the information sharing priors, see:

Husmeier et al. (2010), "Inter-time segment information sharing for non-homogeneous dynamic Bayesian networks", NIPS.

Dondelinger et al. (2012), "Non-homogeneous dynamic Bayesian networks with Bayesian regularization for inferring gene regulatory networks with gradually time-varying structure", Machine Learning.

HyperParms

Sets up initial values of hyperparameters.

Description

This function initialises the variable HYPERvar with values for the various hyperparameters in the model.

Usage

```
HyperParms(options)
```

Arguments

options MCMC settings, possibly from defaultOptions.

Value

Settings for the HYPERvar variable:

cD	Proportion of changepoint moves proposed.
alphaD	Prior settings for the number of changepoints.
betaD	Prior settings for the number of changepoints.

с	Ratio of changepoint birth/death moves proposed.
v0	Prior settings for the sigma squared parameters.
gamma0	Prior settings for the sigma squared parameters.
alphad2	Prior settings for the signal-to-noise ratio delta squared.
betad2	Prior settings for the signal-to-noise ratio delta squared.
alphalbd	Prior settings for the number of transcription factors.
betalbd	Prior settings for the number of transcription factors.

Author(s)

Sophie Lebre Frank Dondelinger

init

Initialise the MCMC simulation.

Description

This function intialises the parameters and variables needed for the MCMC simulation.

Usage

init(X, Y, sinit, GLOBvar, HYPERvar, options)

Arguments

Input response data.
Input target data.
Initial changepoints.
Global variables used during the MCMC simulation.
Hyperparameter variables.
MCMC simulation options as obtained e.g. by defaultOptions.

Value

List with elements:

counters	Matrices for counting the number of moves made and accepted.
initState	Initial state of the variables of the MCMC simulation.
listStock	Variables for recording the network, changepoint and hyperparameter samples.

Author(s)

Sophie Lebre Frank Dondelinger

See Also

sampleParms

main

Main function of the MCMC simulation.

Description

This function executes the main loop of the MCMC simulation, making the different moves and recording samples.

Usage

main(X, Y, initiation, GLOBvar, HYPERvar)

Arguments

Х	Input response data.
Υ	Input target data.
initiation	Initialisation of the MCMC simulation, as obtained by function init.
GLOBvar	Global variables of the MCMC simulation.
HYPERvar	Hyperparameter variables.

Value

Returns a list with the following elements:

counters	List containing the different move counters for the number of times moves have been proposed and accepted.
listStock	List containing the recorded samples for the networks, changepoints and hyper- parameters

Author(s)

Sophie Lebre Frank Dondelinger

References

For more information about the MCMC simulations, see:

Dondelinger et al. (2012), "Non-homogeneous dynamic Bayesian networks with Bayesian regularization for inferring gene regulatory networks with gradually time-varying structure", Machine Learning.

See Also

runDBN

make_structure_move *Makes a structure move*.

Description

This function makes a network structure move.

Usage

make_structure_move(x, y, S, B, Sig2, q, qmax, network.info, method, Mphase, E, fixed.edges, HYPERvar)

Arguments

х	Response data.
У	Target data.
S	Network structure for the current target node, a NumSegs by NumNodes matrix.
В	Same as S, but including the regression parameters.
Sig2	Sigma squared parameters.
q	Number of nodes.
qmax	Maximum number of parents.
network.info	Network information, as collected by CollectNetworkInfo.
method	Information sharing method: Either 'poisson', 'exp_hard', 'exp_soft', 'bino_hard', 'bino_soft'.
Mphase	Segment boundary positions.
E	Changepoint vector.
fixed.edges	Matrix of size NumNodes by NumNodes, with fixed.edges[i,j]==1 0 if the edge between nodes i and j is fixed, and -1 otherwise. Defaults to NULL (no edges fixed).
HYPERvar	Hyperparameter variables.

Value

Returns a list containing the following elements:

newS	Updated network structure.
newB	Updated network structure with regression parameters.
move	Type of move being made: 1 for network structure moves.
accept	1 if the move has been accepted, 0 otherwise.

Author(s)

Frank Dondelinger

References

For more information about the MCMC moves, see:

Dondelinger et al. (2012), "Non-homogeneous dynamic Bayesian networks with Bayesian regularization for inferring gene regulatory networks with gradually time-varying structure", Machine Learning.

NetworkProbBino Calculates the prior probability of the network segments under the binomial prior.

Description

This function calculates the (log) probability of the network segments using the binomial information sharing prior.

Usage

```
NetworkProbBino(network.info, node.sharing = "soft")
```

Arguments

network.info	Network information collected by function CollectNetworkInfo.
node.sharing	Coupling of hyperparameters among nodes: 'hard' or 'soft'.

Value

Returns the log prior probability of the network segments under the binomial prior.

Author(s)

Frank Dondelinger

References

For information about the binomial information sharing prior, see:

Husmeier et al. (2010), "Inter-time segment information sharing for non-homogeneous dynamic Bayesian networks", NIPS.

Dondelinger et al. (2012), "Non-homogeneous dynamic Bayesian networks with Bayesian regularization for inferring gene regulatory networks with gradually time-varying structure", Machine Learning.

See Also

NetworkRatioBino, CalculatePriorRatio

Description

This function calculates the log prior probability of the network structure. It uses the exponential information sharing prior.

Usage

```
NetworkProbExp(network.info)
```

Arguments

network.info Network information collected using the function CollectNetworkInfo

Value

Returns the log prior probability of the network segments.

Author(s)

Frank Dondelinger

References

For information about the exponential information sharing prior, see:

Husmeier et al. (2010), "Inter-time segment information sharing for non-homogeneous dynamic Bayesian networks", NIPS.

Dondelinger et al. (2012), "Non-homogeneous dynamic Bayesian networks with Bayesian regularization for inferring gene regulatory networks with gradually time-varying structure", Machine Learning.

See Also

NetworkRatioExp, CalculatePriorRatio

NetworkRatioBino

Description

This function calculates the ratio of binomial prior probabilities of two networks.

Usage

```
NetworkRatioBino(network.info, node.sharing)
```

Arguments

network.info	Network information collected by function CollectNetworkInfo. Note that
	network.info\$new.nets has to be set.
node.sharing	Type of coupling of hyperparameters among nodes: 'hard' or 'soft'.

Value

Returns the ratio of [prior of new network]/[prior of old network].

Author(s)

For information about the binomial information sharing prior, see:

Husmeier et al. (2010), "Inter-time segment information sharing for non-homogeneous dynamic Bayesian networks", NIPS.

Dondelinger et al. (2012), "Non-homogeneous dynamic Bayesian networks with Bayesian regularization for inferring gene regulatory networks with gradually time-varying structure", Machine Learning.

See Also

NetworkProbBino, CalculatePriorRatio

NetworkRatioExp Calculates the ratio of exponential network prior probabilities.

Description

This function calculates the ratio of exponential network information sharing prior probabilities.

Usage

```
NetworkRatioExp(network.info)
```

output

Arguments

network.info Network information collected using the function CollectNetworkInfo. Note that network.info\$new.nets has to be set.

Value

Returns the ratio [prior of new network]/[prior of old network].

Author(s)

Frank Dondelinger

References

For information about the exponential information sharing prior, see:

Husmeier et al. (2010), "Inter-time segment information sharing for non-homogeneous dynamic Bayesian networks", NIPS.

Dondelinger et al. (2012), "Non-homogeneous dynamic Bayesian networks with Bayesian regularization for inferring gene regulatory networks with gradually time-varying structure", Machine Learning.

See Also

NetworkProbExp, CalculatePriorRatio

output

Collects and saves output.

Description

This function collects the network, changepoint and hyperparameter samples taken from the MCMC simulation, and saves them to a file if appropriate.

Usage

```
output(counters, listStock, GLOBvar, HYPERvar, OUTvar)
```

List of counters for the number of moves that have been proposed and accepted.
Network, changepoint and hyperparameter samples.
Global variables of the MCMC simulation.
Hyperparameter variables.
Output variables, including the output file.

Value

Returns a list with an element for each target node which is also a list. Each sublist containts the elements:

cp_samples	Changepoint samples, a NumSamples by MaxNumChangePoints matrix.
edge_samples	Network samples (with regression parameters), a NumSamples by (NumSegs * NumNodes) matrix.
target	The target node for this subnetwork.
hyper_samples	Information sharing prior hyperparameter samples, a NumSamples by NumHyperParams matrix.
sampled	Sampled iterations.
counters	Counters for the number of proposed and accepted moves.

Author(s)

Frank Dondelinger

phase.update

Make a network structure or hyperparameter move.

Description

This function makes a network structure or information sharing hyperparameter move.

Usage

phase.update(Eall, Sall, Ball, Sig2all, X, Y, GLOBvar, HYPERvar, target)

Arguments

Eall	List of changepoints with one entry for each target node. Each entry has length equal to the number of changepoints for that target node.
Sall	Network structure: List of length equal to the number of target nodes, where each list entry is a NumSegs by NumNodes matrix.
Ball	Network structure with regression coefficients: Same as Sall, but with regression coefficients as matrix entries.
Sig2all	Sigma squared.
Х	Input response data.
Υ	Input target data.
GLOBvar	Global variables used during the MCMC simulation.
HYPERvar	Hyperparameter variables.
target	Current target node.

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PriorRatioPoisson

Value

Returns a list with the following elements:

E	Changepoints for the current target node.
Sall	Network structure (possibly updated).
Ball	Network structure regression coefficients (possibly updated).
Sig2all	Sigma squared.
prior.params	Information sharing prior hyperparameters (possibly updated).
k	Level-2 exponential prior hyperparameter (possibly updated).
move	Move type: 4 for a network structure move, 5 hyperparameter move.
move	Structure Move type: 1 for a network structure move, 2 for a level-1 hyperparameter move, 3 for a level-2 hyperparameter move.
accept	1 if the move has been accepted, 0 otherwise.

Author(s)

Sophie Lebre

Frank Dondelinger

References

For more information on network structure moves and information sharing priors, see:

Dondelinger et al. (2012), "Non-homogeneous dynamic Bayesian networks with Bayesian regularization for inferring gene regulatory networks with gradually time-varying structure", Machine Learning.

See Also

make_structure_move

PriorRatioPoisson Calculate network prior ratio with Poisson prior.

Description

This function calculates the ratio of the Poisson prior for two networks.

Usage

PriorRatioPoisson(network.info, q, lambda)

Arguments

network.info	Network information collected using CollectNetworkInfo. Note that one need	
	to set network.info\$new.nets.	
q	Number of nodes in the network.	
lambda	Vector of lambda hyperparameters for each network.	

Value

Returns the ratio [prior of new network]/[prior of old network].

Author(s)

Frank Dondelinger

References

For more information on the network structure priors, see:

Dondelinger et al. (2012), "Non-homogeneous dynamic Bayesian networks with Bayesian regularization for inferring gene regulatory networks with gradually time-varying structure", Machine Learning.

See Also

CalculatePriorRatio

proposalTuning *Tune the proposal width for betas.*

Description

This function adjusts the proposal width for the beta hyperparameter(s) of the exponential information sharing prior, so that the acceptance rate is close to 0.25.

Usage

proposalTuning(acceptRate, hyper.proposals)

Arguments

acceptRate Current acceptance rate. hyper.proposals Current proposal width.

Value

Returns the new proposal width.

proposeContinuous

Author(s)

Frank Dondelinger

proposeContinuous Propose a new real hyperparameter value.

Description

This function proposes a new real values hyperparameter for the information sharing prior.

Usage

```
proposeContinuous(orig_beta, proposal_range, limit = 30)
```

Arguments

orig_beta	Current value of the hyperparameter.
proposal_range	Range for the new value.
limit	Hard limit on the range.

Value

Returns a new uniformly random value within proposal_range of orig_beta and limited by limit.

Author(s)

Frank Dondelinger

See Also

ProposeDiscrete

```
# Previous parameter value
param = runif(1, 0, 1)
```

```
# Propose new value within range [0, 1], with proposal width 0.1
new.param = proposeContinuous(param, 0.1, 1)
```

ProposeDiscrete

Description

This function proposes a new discrete parameter, based on the previous value, within the given proposal range, making sure that the maximum range is not exceeded.

Usage

ProposeDiscrete(params.old, proposal.range, max.range)

Arguments

params.old	Old parameter value (an integer).
proposal.range	Range for new proposal (an integer).
max.range	Maximum value for new proposal (an integer).

Value

Returns the new proposed parameter, which will be an integer in the range [0, max.range], and within at most proposal.range of params.old.

Author(s)

Frank Dondelinger

See Also

proposeContinuous

```
# Previous parameter value
param = rpois(1, 5)
# Propose new value within range [0, 10], with proposal width 2
new.param = ProposeDiscrete(param, 2, 10)
```

Description

This function calculates the potential scale reduction factor of parameters or hyperparameters over several MCMC simulations (or one simulation split up). This can serve as a convergence diagnostic.

Usage

psrf(parameters)

Arguments

parameters A list of MCMC trajectories, where each trajectory is a matrix with NumParams rows and NumIterations columns, where NumParams is the number of parameters and NumIterations is the number of samples.

Value

A vectors of length NumParams, containing the PSRF values for each parameter.

Author(s)

Sophie Lebre Frank Dondelinger

References

Gelman and Rubin (1992) Inference from iterative simulation using multiple sequences, Statistical Science.

See Also

psrf_check, psrf_check_hyper

```
# Generate 5 'runs' of random samples from Gaussian N(0,1)
samples = list()
for(run in 1:5) {
   samples[[run]] = matrix(rnorm(1000), 1, 1000)
}
# Check potential scale reduction factor
# (Will be very close to 1 due to the samples being from
# the same distribution)
psrf.val = psrf(samples)
```

```
# Now use slightly different Gaussian distributions for each 'run'.
for(run in 1:5) {
    mean = runif(1, 0, 2)
    samples[[run]] = matrix(rnorm(1000, mean, 1), 1, 1000)
}
# Check potential scale reduction factor
# (Should be > 1.1)
psrf.val = psrf(samples)
```

psrf_check

Check the potential scale reduction factors for all parameters (edges).

Description

This function treats the edges of the network as parameters, calculates their potential scale reduction factors and returns the highest value.

Usage

psrf_check(params, q, k_max, num_it)

Arguments

params	Matrix of parameters.
q	Number of nodes.
k_max	Number of segments.
num_it	Number of iterations/samples.

Value

Returns the highest PSRF value.

Author(s)

Frank Dondelinger

References

Gelman and Rubin (1992) Inference from iterative simulation using multiple sequences, Statistical Science.

See Also

psrf, psrf_check_hyper

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psrf_check_hyper Checks the potential scale reduction factor for the hyperparameters.

Description

This function checks the potential scale reduction factors for the hyperparameters of the information sharing priors.

Usage

psrf_check_hyper(params, num_it)

Arguments

params	Matrix of hyperparameters.
num_it	Number of iterations/samples.

Value

Returns the maximum PSRF value.

Author(s)

Frank Dondelinger

References

Gelman and Rubin (1992) Inference from iterative simulation using multiple sequences, Statistical Science.

See Also

psrf, psrf_check

readDataTS Read target data.

Description

This function reads in the target data.

Usage

readDataTS(data, posI, t0, tf, m, n)

Arguments

data	Input data matrix to read
posI	Position of interest.
t0	First timepoint.
tf	Last timepoint.
m	Number of repetitions.
n	Number of timepoints.

Value

Returns the target data.

Author(s)

Sophie Lebre

See Also

buildXY

rinvgamma

Samples from the inverse gamma distribution.

Description

This function samples from the inverse gamma distribution.

Usage

rinvgamma(n, shape, scale)

Arguments

n	Number of values to sample.
shape	Shape parameter.
scale	Scale parameter (1/rate).

Value

Random sample from the inverse gamma distribution.

Author(s)

Frank Dondelinger

runDBN

See Also

dinvgamma

Examples

```
# Draw samples from inverse gamma distribution with shape parameter 1
# and scale parameter 1
samples = rinvgamma(100, shape=1, scale=1)
# Calculate density of samples
```

densities = dinvgamma(samples, shape=1, scale=1)

runDBN

Setup and run the MCMC simulation.

Description

This function initialises the variabes for the MCMC simulation, runs the simulation and returns the output.

Usage

```
runDBN(targetdata, preddata = NULL, q, n, multipleVar = TRUE,
minPhase = 2, niter = 20000, scaling = TRUE, method = "poisson",
prior.params = NULL, self.loops = TRUE, k = 15, options = NULL,
outputFile = ".", fixed.edges = NULL)
```

targetdata	Target input data: A matrix of dimensions NumNodes by NumTimePoints.
preddata	Optional: Input response data, if different from the target data.
q	Number of nodes.
n	Number of timepoints.
multipleVar	TRUE when a specific variance is estimated for each segment, FALSE otherwise.
minPhase	Minimal segment length.
niter	Number of MCMC iterations.
scaling	If TRUE, scale the input data to mean 0 and standard deviation 1, else leave it unchanged.
method	Network structure prior to use: 'poisson' for a sparse Poisson prior (no infor- mation sharing), 'exp_hard' or 'exp_soft' for the exponential information sharing prior with hard or soft node coupling, 'bino_hard' or 'bino_soft' with hard or soft node coupling.
prior.params	Initial hyperparameters for the information sharing prior.

self.loops	If TRUE, allow self-loops in the network, if FALSE, disallow self-loops.
k	Initial value for the level-2 hyperparameter of the exponential information sharing prior.
options	MCMC options as obtained e.g. by the function defaultOptions.
outputFile	File where the output of the MCMC simulation should be saved.
fixed.edges	Matrix of size NumNodes by NumNodes, with fixed.edges[i,j]==1 0 if the edge between nodes i and j is fixed, and -1 otherwise. Defaults to NULL (no edges fixed).

Value

A list containing the results of the MCMC simulation: network samples, changepoint samples and hyperparameter samples. For details, see output.

Author(s)

Sophie Lebre Frank Dondelinger

References

For more information about the MCMC simulations, see:

Dondelinger et al. (2012), "Non-homogeneous dynamic Bayesian networks with Bayesian regularization for inferring gene regulatory networks with gradually time-varying structure", Machine Learning.

See Also

output

sampleBinit

Sample initial regression coefficients.

Description

This function samples the initial regression coefficients for the networks.

Usage

```
sampleBinit(Si, sig2, delta2, X, q)
```

Si	Network structure.
sig2	Sigma squared.
delta2	Signal-to-noise ratio hyperparameter.
Х	Input data.
q	Number of nodes.

sampleBxy

Value

Returns a vector of regression coefficients.

Author(s)

Sophie Lebre

References

For details of the regression model, see:

Dondelinger et al. (2012), "Non-homogeneous dynamic Bayesian networks with Bayesian regularization for inferring gene regulatory networks with gradually time-varying structure", Machine Learning.

sampleBxy

Sample regression coefficients.

Description

This function samples the regression coefficients given the current state of the MCMC simulation.

Usage

sampleBxy(xi, y, Sig2, delta2)

Arguments

xi	Response data.
У	Target data.
Sig2	Sigma squared.
delta2	Signal-to-noise hyperparameter.

Value

The regression parameters.

Author(s)

Sophie Lebre

References

For details of the regression model, see:

Dondelinger et al. (2012), "Non-homogeneous dynamic Bayesian networks with Bayesian regularization for inferring gene regulatory networks with gradually time-varying structure", Machine Learning. sampleDelta2

Description

This function samples the signal-to-noise hyperparameter delta squared.

Usage

sampleDelta2(pos, x, q, B, S, sig2, alphad2, betad2)

Arguments

pos	The current segment.
x	Data,
q	Number of nodes.
В	Regression coefficients.
S	Network structure.
sig2	Sigma squared.
alphad2	Gamma prior hyperparameter.
betad2	Gamma prior hyperparameter.

Value

New sample of delta squared.

Author(s)

Sophie Lebre

References

For details of the sampling, see:

Dondelinger et al. (2012), "Non-homogeneous dynamic Bayesian networks with Bayesian regularization for inferring gene regulatory networks with gradually time-varying structure", Machine Learning.

sampleK

Description

This function samples the initial number of changepoints from a sparse Poisson prior.

Usage

sampleK(mini, maxi, lambda, nb)

Arguments

mini	Minimum value.
maxi	Maximum value.
lambda	Parameter of the Poisson distribution.
nb	Number of values to sample.

Value

The sampled number of changepoints.

Author(s)

Sophie Lebre

References

For more information on the prior choice and sampling, see:

Dondelinger et al. (2012), "Non-homogeneous dynamic Bayesian networks with Bayesian regularization for inferring gene regulatory networks with gradually time-varying structure", Machine Learning.

sampleParms

Sample initial parameters for the MCMC simulation.

Description

This function samples the initial hyperparameters and parameters that are needed for the MCMC simulation.

Usage

```
sampleParms(X, GLOBvar, HYPERvar, s_init = NULL, options)
```

Arguments

Input data.
Global variables of the MCMC simulation.
Hyperparameter variables.
Initial number of changepoints.
MCMC options, as given by e.g. default0ptions.

Value

Returns a list with elements:

E	The initial changepoint vector.
S	The intial networks structure.
В	The initial regression parameters.
Sig2	The inital sigma squared variances.
betas	The intial hyperparameters for the exponential information sharing prior.
hyper_params	The initial hyperparameters for the binomial information sharing prior.

Author(s)

Sophie Lebre Frank Dondelinger

References

For more information about the parameters and hyperparameters, see:

Dondelinger et al. (2012), "Non-homogeneous dynamic Bayesian networks with Bayesian regularization for inferring gene regulatory networks with gradually time-varying structure", Machine Learning.

See Also

init

sampleSig2 Sample initial sigma squared.

Description

This function samples the initial values for the sigma squared variance from the inverse gamma prior.

Usage

sampleSig2(y, Px, v0, gamma0)

simulateNetwork

Arguments

У	Input data.
Px	Projection matrix.
v0	Inverse gamma prior hyperparameter.
gamma0	Inverse gamma prior hyperparameter.

Value

The sampled sigma squared values.

Author(s)

Sophie Lebre

References

For more information about the model, see:

Dondelinger et al. (2012), "Non-homogeneous dynamic Bayesian networks with Bayesian regularization for inferring gene regulatory networks with gradually time-varying structure", Machine Learning.

simulateNetwork *Generate network and simulate data*.

Description

This function generates a random network with structure changepoints (or takes one as input) and simulated data from it using a regression model.

Usage

```
simulateNetwork(1 = 100, min_phase_length = 10, k_bar = 10, q = 10,
lambda_2 = 0.45, noise = 0.25, net = NULL, lambda_3 = 2,
spacing = 0, gauss_weights = FALSE, same = FALSE,
changes = "sequential", fixed = FALSE, cps = NULL, saveFile = NULL)
```

1	Length of the time series.
min_phase_length	
	Minimum segment length.
k_bar	Maximum number of changepoints.
q	Number of nodes.
lambda_2	Average number of parents for each node in the network (parameter for a Poisson distribution).

noise	Standard deviation of the Gaussian observation noise. Can be constant, or seg- ment specific (in which case the number of changepoints needs to be fixed and the noise needs to be a vector of the same length).
net	Input network, can be NULL if a new network should be generated.
lambda_3	Average number of structure changes between two segments (parameter for a Poisson distribution).
spacing	1 if segments are equally spaced, 0 if they are spaced randomly (subject to the constraints of min_phase_length).
gauss_weights	1 if edge weights in the network are drawn from $N(0, 1)$, 0 if they are fixed to be 1.
same	1 if the networks should all be the same (no changes), 0 otherwise.
changes	'sequential' if the changes happen sequentially (i.e. changes at segment i are applied to segment i-1), 'hierarchical' if the changes happen with respect to a hypernetwork (i.e. changes at segment i are applied to segment 0).
fixed	T if the changepoint locations are fixed, F if they should be sampled.
cps	Changepoint locations (if they are fixed).
saveFile	If not NULL, indicates the filename for saving the output in R data format.

Value

A list with elements:

sim_data	A matrix of length NumNodes by NumTimepoints containing the simulated data from the regression model.
epsilon	Changepoint vector.
k	Number of changepoints.
network	The network, a list of length NumSegs, where each element is a NumNodes by NumNodes matrix.
noise	The standard deviation of the applied Gaussian noise.

Author(s)

Frank Dondelinger

See Also

generateNetwork

```
# Generate random network and simulate data with default parameters
dataset = simulateNetwork()
```

```
# Generate random network and simulate data with an average of
# 1 change per node among network segments
dataset = simulateNetwork(lambda_3=1)
```

updateSigMulti

```
# Generate random network and simulate data with an average of
# 1 change per node among network segments and standard deviation
# of the Gaussian observation noise 0.5
dataset = simulateNetwork(lambda_3=1, noise=0.5)
# Generate random network with default parameters
network = generateNetwork()
# Simulate data using generated network
dataset = simulateNetwork(net=network)
# Generate random network with 4 changepoints and 15 nodes,
# with changepoints distributed over a timeseries of length 50
network = generateNetwork(l=50, q=15, fixed=TRUE, k_bar=4)
# Simulate data of length 50 using generated network
dataset = simulateNetwork(net=network)
```

updateSigMulti Update sigma squared variances.

Description

This function samples new values for the sigma squared variances, given the current network structure. A multivariate distribution is assumed.

Usage

```
updateSigMulti(phase, X, Y, E, Sall, Ball, Sig2, Mphase, alphad2, betad2, v0, gamma0)
```

phase	Current segment.
Х	Input response data.
Υ	Input target data.
E	Changepoints.
Sall	Network structure.
Ball	Regression coefficients.
Sig2	Current sigma squared values.
Mphase	Segment positions.
alphad2	Hyperparameter for gamma prior.
betad2	Hyperparameter for gamma prior.
v0	Hyperparameter for inverse gamma prior.
gamma0	Hyperparameter for inverse gamma prior.

The new samples sigma squared values.

Author(s)

Sophie Lebre

References

For more information about the model, see:

Dondelinger et al. (2012), "Non-homogeneous dynamic Bayesian networks with Bayesian regularization for inferring gene regulatory networks with gradually time-varying structure", Machine Learning.

See Also

updateSigSolo

updateSigSolo Sample new values for sigma squared.

Description

This function samples new values for the sigma squared variances, given the current network structure. A univariate distribution is assumed.

Usage

updateSigSolo(X, Y, E, Sall, Ball, Sig2, Mphase, alphad2, betad2, v0, gamma0)

Input response data.
Input target data.
Changepoints.
Network structure.
Regression coefficients.
Current sigma squared.
Segment position.
Gamma prior hyperparameter.
Gamma prior hyperparameter.
Inverse gamma prior hyperparameter.
Inverse gamma prior hyperparameter.

updateSigSolo

Value

Returns the new samples sigma squared values.

Author(s)

Sophie Lebre

References

For more information about the model, see:

Dondelinger et al. (2012), "Non-homogeneous dynamic Bayesian networks with Bayesian regularization for inferring gene regulatory networks with gradually time-varying structure", Machine Learning.

See Also

updateSigMulti

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