

Package ‘sparsebn’

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Title Learning Sparse Bayesian Networks from High-Dimensional Data

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Description

Fast methods for learning sparse Bayesian networks from high-dimensional data using sparse regularization, as described in Aragam, Gu, and Zhou (2017) <arXiv:1703.04025>. Designed to handle mixed experimental and observational data with thousands of variables with either continuous or discrete observations.

Depends R (>= 3.2.3), sparsebnUtils (>= 0.0.5), ccdAlgorithm (>= 0.0.4), discretedAlgorithm (>= 0.0.5)

Suggests knitr, rmarkdown, mvtnorm, igraph, graph, testthat

URL <https://github.com/itsrainingdata/sparsebn>

BugReports <https://github.com/itsrainingdata/sparsebn/issues>

License GPL (>= 2)

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cytometryContinuous *The continuous cytometry network*

Description

Data and network for analyzing the flow cytometry experiment from [Sachs et al. \(2005\)](#) [1]. This dataset contains the raw measurements from these experiments.

Usage

```
data(cytometryContinuous)
```

Format

A **list** with three components:

- dag An **edgeList** containing the consensus network (11 nodes, 17 edges).
- data A **data.frame** with 11 variables and 7466 observations.
- ivn A **list** specifying which nodes are under intervention in each observation. Compatible with the input to **sparsebnData**.

Details

The dataset consists of $n = 7466$ observations of $p = 11$ continuous variables corresponding to different proteins and phospholipids in human immune system cells, and each observation indicates the measured level of each biomolecule in a single cell under different experimental interventions. Based on this data, a consensus network was reconstructed and validated, which is included as well.

References

[1] Sachs, Karen, et al. "[Causal protein-signaling networks derived from multiparameter single-cell data.](#)" *Science* 308.5721 (2005): 523-529.

Examples

```
# Create a valid sparsebnData object from the cytometry data
data(cytometryContinuous)
dat <- sparsebnData(cytometryContinuous$data, type = "c", ivn = cytometryContinuous$ivn)
```

cytometryDiscrete *The discrete cytometry network*

Description

Data and network for analyzing the flow cytometry experiment from [Sachs et al. \(2005\)](#) [1]. The data is a cleaned and discretized version of the raw data (see [cytometryContinuous](#) for details) from these experiments.

Usage

```
data(cytometryDiscrete)
```

Format

A [list](#) with three components:

- dag An [edgeList](#) containing the consensus network (11 nodes, 17 edges).
- data A [data.frame](#) with 11 variables and 5400 observations.
- ivn A [list](#) specifying which nodes are under intervention in each observation. Compatible with the input to [sparsebnData](#).

Details

After cleaning and pre-processing, the raw continuous measurements have been binned into one of three levels: low = 0, medium = 1, or high = 2. Due to the pre-processing, the discrete data contains fewer observations (n = 5400) compared to the raw, continuous data.

References

[1] Sachs, Karen, et al. "[Causal protein-signaling networks derived from multiparameter single-cell data.](#)" *Science* 308.5721 (2005): 523-529.

Examples

```
# Create a valid sparsebnData object from the cytometry data
data(cytometryDiscrete)
dat <- sparsebnData(cytometryDiscrete$data, type = "d", ivn = cytometryDiscrete$ivn)
```

estimate.covariance *Covariance estimation*

Description

Methods for inferring (i) Covariance matrices and (ii) Precision matrices for continuous, Gaussian data.

Usage

```
estimate.covariance(data, ...)
```

```
estimate.precision(data, ...)
```

Arguments

`data` data as [sparsebnData](#) object.
`...` (optional) additional parameters to [estimate.dag](#)

Details

For Gaussian data, the precision matrix corresponds to an undirected graphical model for the distribution. This undirected graph can be tied to the corresponding directed graphical model; see Sections 2.1 and 2.2 (equation (6)) of Aragam and Zhou (2015) for more details.

Value

Solution path as a plain [list](#). Each component is a [Matrix](#) corresponding to an estimate of the covariance or precision (inverse covariance) matrix for a given value of lambda.

Examples

```
data(cytometryContinuous)
dat <- sparsebnData(cytometryContinuous$data, type = "c", ivn = cytometryContinuous$ivn)
estimate.covariance(dat) # estimate covariance
estimate.precision(dat) # estimate precision
```

estimate.dag	<i>Estimate a DAG from data</i>
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Description

Estimate the structure of a DAG (Bayesian network) from data. Works with any combination of discrete / continuous and observational / experimental data.

Usage

```
estimate.dag(data, lambdas = NULL, lambdas.length = 20,
  whitelist = NULL, blacklist = NULL, error.tol = 1e-04,
  max.iters = NULL, edge.threshold = NULL, concavity = 2,
  weight.scale = 1, convLb = 0.01, upperbound = 100,
  adaptive = FALSE, verbose = FALSE)
```

Arguments

data	Data as sparsebnData .
lambdas	(optional) Numeric vector containing a grid of lambda values (i.e. regularization parameters) to use in the solution path. If missing, a default grid of values will be used based on a decreasing log-scale (see also generate.lambdas).
lambdas.length	Integer number of values to include in the solution path. If lambdas has also been specified, this value will be ignored.
whitelist	A two-column matrix of edges that are guaranteed to be in each estimate (a "white list"). Each row in this matrix corresponds to an edge that is to be whitelisted. These edges can be specified by node name (as a character matrix), or by index (as a numeric matrix).
blacklist	A two-column matrix of edges that are guaranteed to be absent from each estimate (a "black list"). See argument "whitelist" above for more details.
error.tol	Error tolerance for the algorithm, used to test for convergence.
max.iters	Maximum number of iterations for each internal sweep.
edge.threshold	Threshold parameter used to terminate the algorithm whenever the number of edges in the current estimate has > edge.threshold edges. NOTE: This is not the same as alpha in ccdr.run .
concavity	(CCDr only) Value of concavity parameter. If $\gamma > 0$, then the MCP will be used with gamma as the concavity parameter. If $\gamma < 0$, then the L1 penalty will be used and this value is otherwise ignored.
weight.scale	(CD only) A positive number to scale weight matrix.
convLb	(CD only) Small positive number used in Hessian approximation.
upperbound	(CD only) A large positive value used to truncate the adaptive weights. A -1 value indicates that there is no truncation.
adaptive	(CD only) TRUE / FALSE, if TRUE the adaptive algorithm will be run.
verbose	TRUE / FALSE whether or not to print out progress and summary reports.

Details

For details on the underlying methods, see `ccdr.run` and `cd.run`.

Value

A `sparsebnPath` object.

Examples

```
# Estimate a DAG from the cytometry data
data(cytometryContinuous)
dat <- sparsebnData(cytometryContinuous$data, type = "c", ivn = cytometryContinuous$ivn)
estimate.dag(dat)
```

pathfinder

The pathfinder network

Description

Simulated data and network for the pathfinder network from the [Bayesian network repository](#). Pathfinder is an expert system developed by [Heckerman et. al \(1992\)](#) [1] to assist with the diagnosis of lymph-node diseases.

Usage

```
data(pathfinder)
```

Format

A `list` with four components:

- `dag` An `edgeList` containing the pathfinder network (109 nodes, 195 edges).
- `data` A `data.frame` with 109 variables and 1000 observations.
- `ivn` A `list` specifying which nodes are under intervention in each observation; since this dataset is purely observational, this is just `NULL`. Compatible with the input to `sparsebnData`.
- `cov` Covariance matrix used to generate the data.

Details

This is a benchmark network used to test algorithms for learning Bayesian networks. The data is simulated from a Gaussian SEM assuming unit edge weights and unit variances for all nodes.

References

[1] Heckerman, David E., and Bharat N. Nathwani. "An evaluation of the diagnostic accuracy of Pathfinder." *Computers and Biomedical Research* 25.1 (1992): 56-74.

Examples

```
### Create a valid sparsebnData object from the simulated pathfinder data
data(pathfinder)
dat <- sparsebnData(pathfinder$data, type = "c")

### Code to reproduce this dataset by randomly generating edge weights
coefs <- runif(n = num.edges(pathfinder$dag), min = 0.5, max = 2) # coefficients
vars <- rep(1, num.nodes(pathfinder$dag)) # variances
params <- c(coefs, vars) # parameter vector
pathfinder.data <- generate_mvn_data(graph = pathfinder$dag,
                                   params = params,
                                   n = 1000)
```

plotDAG

Plot a DAG

Description

Using some sensible defaults for large graphs, plot a DAG object. Uses [igraph](#) package by default.

Usage

```
plotDAG(x, ...)
```

Arguments

x An [edgeList](#), [sparsebnFit](#), or [sparsebnPath](#) object.

... Additional arguments to [plot](#).

Details

This method is not intended for customization. For more control over the output, use [plot](#) and see [setPlotPackage](#) for plotting only and/or [setGraphPackage](#) for even more control. These methods grants the user the full feature set of the selected package.

sparsebn

sparsebn: Learning Sparse Bayesian Networks from High-Dimensional Data.

Description

Methods for learning sparse Bayesian networks and other graphical models from observational and experimental data via sparse regularization. Includes algorithms for both continuous and discrete data.

Details

The main methods for learning graphical models in [sparsebn](#) are:

- [estimate.dag](#) for directed acyclic graphs.
- [estimate.precision](#) for undirected graphs.
- [estimate.covariance](#) for covariance matrices.

The workhorse behind [sparsebn](#) is the [sparsebnUtils](#) package, which provides various S3 classes and methods for representing and manipulating graphs. For more details on this package and the functionality it provides, see [?sparsebnUtils](#).

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