

Package ‘MXM’

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Type Package

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Description Many feature selection methods for a wide range of response variables, including minimal, statistically-equivalent and equally-predictive feature subsets. Bayesian network algorithms and related functions are also included. The package name 'MXM' stands for ``Mens eX Machina'', meaning ``Mind from the Machine'' in Latin. Reference: Feature Selection with the R Package MXM: Discovering Statistically Equivalent Feature Subsets, Lagani, V. and Athineou, G. and Farcomeni, A. and Tsagris, M. and Tsamardinos, I. (2017). Journal of Statistical Software, 80(7). <doi:10.18637/jss.v080.i07>.

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MXM-package	4
Ancestors and descendants of a node in a directed graph	6
Backward phase of MMPC	7
Backward selection regression	9
Backward selection regression for GLMM	10
Backward selection regression for GLMM using the eBIC	12
Backward selection regression using the eBIC	13
Backward selection with generalised linear regression models	15
Bayesian Network construction using a hybrid of MMPC and PC	16
Beta regression	19
BIC based forward selection	20
BIC based forward selection with generalised linear models	22
Calculation of the constant and slope for each subject over time	24
Certificate of exclusion from the selected variables set using SES or MMPC	25
Check Markov equivalence of two DAGs	26
Check whether a directed graph is acyclic	27
CondIditional independence tests	28
Conditional independence test for binary, categorical or ordinal data	35
Conditional independence test for case control data	39
Conditional independence test for longitudinal and clustered data using GEE	42
Conditional independence test for longitudinal and clustered data using GLMM	45
Conditional independence test for proportions/percentages	48
Conditional independence test for the static-longitudinal scenario	50
Conditional independence tests counting the number of times a possible collider d- separates two nodes	53
Conditional independence tests for continous univariate and multivariate data	55
Conditional independence tests for count data	58
Conditional independence tests for left censored data	61
Conditional independence tests for positive data	64
Conditional independence tests for sucess rates	67
Conditional independence tests for survival data	69
Conditional independence tests with and without permutation p-value	73
Constraint based feature selection algorithms	75
Constraint based feature selection algorithms for longitudinal and clustered data	81
Constraint based feature selection algorithms for multiple datasets	86
Correlation based conditonal independence tests	91
Cross-validation for ridge regression	94
Cross-Validation for SES and MMPC	96
Data simulation from a DAG	100
Drop all possible single terms from a model using the partial correlation	103
Effective sample size for G^2 test in BNs with case control data	104
Estimation of the percentage of Null p-values	105
Feature selection using SES and MMPC for classification with longitudinal data	106
Fit a mixture of beta distributions in p-values	109
Forward Backward Early Dropping selection regression	110
Forward Backward Early Dropping selection regression with GEE	113

Forward Backward Early Dropping selection regression with GLMM	116
Forward selection regression	118
Forward selection with generalised linear regression models	121
Forward selection with linear regression models	123
G-square conditional independence test for discrete data	124
Generalised linear mixed models based on temporal SES and MMPC outputs	127
Generalised ordinal regression	129
Generalised orthogonal matching pursuit (gOMP)	131
Generate random folds for cross-validation	132
IAMB backward selection phase	133
IAMB variable selection	135
Incremental BIC values and final regression model of the FBED algorithm	137
Interactive plot of an (un)directed graph	138
Lower limit of the confidence of an edge	139
mammpc.output-class	140
Many simple beta regressions	141
Many simple zero inflated Poisson regressions	142
Many Wald based tests for logistic and Poisson regressions with continuous predictors	144
Markov Blanket of a node in a directed graph	145
mases.output-class	146
MMPC solution paths for many combinations of hyper-parameters	147
MMPC.gee.output-class	149
MMPC.temporal.output-class	151
MMPCoutput-class	152
MXM-internal	153
Neighbours of nodes in an undirected graph	158
Network construction using the partial correlation based forward regression or FBED	159
Orientation rules for the PC algorithm	161
Partial correlation between two variables	163
Permutation based p-value for the Pearson correlation coefficient	164
Plot of longitudinal data	165
Regression models based on SES and MMPC outputs	167
Regression models based on SES.timeclass and MMPC.timeclass outputs	169
Regression models fitting	171
Ridge regression	173
Ridge regression coefficients plot	174
ROC and AUC	175
Search for triangles in an undirected graph	176
SES.gee.output-class	177
SES.temporal.output-class	178
SESoutput-class	179
Skeleton (local) around a node of the max-min hill-climbing (MMHC) algorithm	180
Skeleton of the max-min hill-climbing (MMHC) algorithm	182
Skeleton of the PC algorithm	185
Structural Hamming distance between two partially oriented DAGs	189
Symmetric conditional independence test with clustered data	190
Symmetric conditional independence test with mixed data	191
The max-min Markov blanket algorithm	193

Topological sort of a DAG	195
Total causal effect of a node on another node	196
Transformation of a DAG into an essential graph	197
Transitive closure of an adjacency matrix	199
Undirected path(s) between two nodes	200
Univariate regression based tests	201
Utilities for the skeleton of a (Bayesian) Network	204
Zero inflated Poisson regression	206

Index 209

MXM-package	<i>This is an R package that currently implements feature selection methods for identifying minimal, statistically-equivalent and equally-predictive feature subsets. In addition, two algorithms for constructing the skeleton of a Bayesian network are included.</i>
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Description

'MXM' stands for Mens eX Machina, meaning 'Mind from the Machine' in Latin. The package provides source code for the SES algorithm and for some appropriate statistical conditional independence tests. (Fisher and Spearman correlation, G-square test are some examples. Currently the response variable can be univariate or multivariate Euclidean, proportions within 0 and 1, compositional data without zeros and ones, binary, nominal or ordinal multinomial, count data (handling also overdispersed and with more zeros than expected), longitudinal, clustered data, survival and case-control. Robust versions are also available in some cases and a K-fold cross validation is offered. Bayesian network related algorithms and ridge regression are also included. Read the package's help pages for more details.

MMPC and SES can handle even thousands of variables and for some tests, even many sample sizes of tens of thousands. The user is best advised to check his variables in the beginning. For some regressions, logistic and Poisson for example, we have used C++ codes for speed reasons. Thus no check is done for a variable with zero variance for instance. Something like "colVars" (in "Rfast") could be used in the first place to remove variables with zero variance.

For more information the reader is addressed to Lagani V., Athineou G., Farcomeni A., Tsagris M. and Tsamardinos I. (2017). Feature Selection with the R Package MXM: Discovering Statistically Equivalent Feature Subsets. Journal of Statistical Software, 80(7), doi:10.18637/jss.v080.i07.

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Maintainer

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Disclaimer: Professor Tsamardinos is the creator of this package. Dr Lagani supervised Mr Athineou built it and Dr Tsagris is the maintainer.

Author(s)

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See Also

[SES](#), [MMPC](#), [censIndCR](#), [testIndFisher](#), [testIndLogistic](#), [gSquare](#), [testIndRQ](#)

Ancestors and descendants of a node in a directed graph

Returns and plots, if asked, the descendants or ancestors of one or all node(s) (or variable(s))

Description

Returns and plots, if asked, the descendants or ancestors of one or all node(s) (or variable(s))

Usage

```
findDescendants(G, node = NULL, graph = FALSE)
findAncestors(G, node = NULL, graph = FALSE)
```

Arguments

G	The graph matrix as produced from pc.or or any other algorithm which produces directed graphs.
node	A numerical value indicating the node (or variable) whose descendants are to be returned.
graph	A boolean variable. If TRUE the relevant graph will appear (if there are descendants).

Details

The functions searches for the descendants of some node. This is an S3 class output.

Value

isAnc	A matrix of the same dimensions as the original graph matrix with 0s and 1s. isAnc[i, j] = 1 indicates that the i-th node is an ancestor of the j-th node. If the argument "node" is NULL, only this matrix will be returned.
Ganc	A matrix of dimensions equal to the number of descendants of the node with 0s and 1s, if the argument "node" is not NULL.
anc	The descendants of the node if the argument "node" is not NULL.

Author(s)

Anna Roumpelaki

R implementation and documentation: Anna Roumpelaki <anna.roumpelaki@gmail.com>

See Also

[plotnetwork](#), [nei](#), [mb](#), [pc.or](#)

Examples

```
# simulate a dataset with continuous data
# simulate a dataset with continuous data
y = rdag(1000, 10, 0.3)
tru = y$G
x = y$x
mod = pc.con(x)
G = pc.or(mod)$G
plotnetwork(G)
findDescendants(G, 4, graph = FALSE)
findAncestors(G, 4, graph = FALSE)
findAncestors(G)
```

Backward phase of MMPC

Backward phase of MMPC

Description

Backward phase of MMPC.

Usage

```
mmpcbackphase(target, dataset, max_k = 3, threshold = 0.05, test = NULL,
wei = NULL, R = 1)
```

Arguments

target	The class variable. Provide either a string, an integer, a numeric value, a vector, a factor, an ordered factor or a Surv object. See also Details.
dataset	The data-set; provide either a data frame or a matrix (columns = variables , rows = samples). Alternatively, provide an ExpressionSet (in which case rows are samples and columns are features, see bioconductor for details).
max_k	The maximum conditioning set to use in the conditional independence test (see Details). Integer, default value is 3.
threshold	Threshold (suitable values in [0,1]) for assessing p-values significance. Default value is 0.05.
test	The conditional independence test to use. Type the test without " ", e.g. type testIndFisher, Not "testIndFisher". Default value is NULL. See also CondIndTests .
wei	A vector of weights to be used for weighted regression. The default value is NULL.

R The number of permutations, set to 1 by default (no permutations based test). There is a trick to avoid doing all permutations. As soon as the number of times the permuted test statistic is more than the observed test statistic is more than 50 (if threshold = 0.05 and R = 999), the p-value has exceeded the significance level (threshold value) and hence the predictor variable is not significant. There is no need to continue do the extra permutations, as a decision has already been made.

Details

For each of the selected variables (dataset) the function performs conditional independence tests where the conditioning sets are formed from the other variables. All possible combinations are tried until the variable becomes non significant. The maximum size of the conditioning set is equal to `max_k`. This is called in the [MMPC](#) when the backward phase is requested.

Value

A list including:

<code>met</code>	A numerical vector of size equal to the number of columns of the dataset.
<code>counter</code>	The number of tests performed.
<code>pvalue</code>	The maximum p-value for the association of each predictor variable.

Author(s)

Ioannis Tsamardinos, Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

Tsamardinos, Brown and Aliferis (2006). The max-min hill-climbing Bayesian network structure learning algorithm. *Machine learning*, 65(1), 31-78.

See Also

[MMPC](#), [mmhc.skel](#), [CondIndTests](#), [cv.mmmpc](#)

Examples

```
set.seed(123)
require("hash", quietly = TRUE)

#simulate a dataset with continuous data
dataset <- matrix(runif(1000 * 200, 1, 100), ncol = 200)

#define a simulated class variable
target <- 3 * dataset[, 10] + 2 * dataset[, 100] + 3 * dataset[, 20] + rnorm(1000, 0, 5)

# define some simulated equivalences
dataset[, 15] <- dataset[, 10] + rnorm(1000, 0, 2)
```

```

dataset[, 150] <- dataset[, 100] + rnorm(1000, 0, 2)
dataset[, 130] <- dataset[, 100] + rnorm(1000, 0, 2)

# MMPC algorithm
m1 <- MMPC(target, dataset, max_k = 3, threshold = 0.05, test="testIndFisher");
m2 <- MMPC(target, dataset, max_k = 3, threshold = 0.05, test="testIndFisher", backward = TRUE);
x <- dataset[, m1@selectedVars]
mmpcbackphase(target, x, test = testIndFisher)

```

Backward selection regression

Variable selection in regression models with backward selection

Description

Variable selection in regression models with backward selection

Usage

```
bs.reg(target, dataset, threshold = 0.05, wei = NULL, test = NULL, user_test = NULL)
```

Arguments

target	The class variable. Provide either a string, an integer, a numeric value, a vector, a factor, an ordered factor or a Surv object. See also Details. For the gamma regression this must a vector with strictly positive numbers (no zeros allowed).
dataset	The dataset; provide either a data frame or a matrix (columns = variables, rows = samples). In either case, only two cases are available, either all data are continuous, or categorical.
threshold	Threshold (suitable values in [0,1]) for assessing p-values significance. Default value is 0.05.
test	The regression model to use. Available options are most of the tests for SES and MMPC. The ones NOT available are "gSquare", "censIndER", "testIndMVreg", "testIndSpearman". If you want to use multinomial or ordinal logistic regression, make sure your target is factor. See also SES and CondIndTests for the tests.
wei	A vector of weights to be used for weighted regression. The default value is NULL.
user_test	A user-defined conditional independence test (provide a closure type object). Default value is NULL. If this is defined, the "test" argument is ignored.

Details

If the sample size is less than the number of variables a message will appear and no backward regression is performed.

Value

The output of the algorithm is S3 object including:

runtime	The run time of the algorithm. A numeric vector. The first element is the user time, the second element is the system time and the third element is the elapsed time.
info	A matrix with the non selected variables and their latest test statistics and p-values.
mat	A matrix with the selected variables and their latest statistics and logged p-values.
ci_test	The conditional independence test used.
final	The final regression model.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

See Also

[glm.fsreg](#), [lm.fsreg](#), [bic.fsreg](#), [bic.glm.fsreg](#), [CondIndTests](#), [MMPC](#), [SES](#)

Examples

```
set.seed(123)
dataset <- matrix( runif(1000 * 10, 1, 100), ncol = 10 )
target <- rnorm(1000)
a <- bs.reg(target, dataset, threshold = 0.05, test = "testIndRQ")
b <- bs.reg(target, dataset, threshold = 0.05, test = "testIndReg")
b2 <- bs.reg(target, dataset, threshold = 0.05, test = "testIndFisher")
```

Backward selection regression for GLMM

Backward selection regression for GLMM

Description

Backward selection regression for GLMM

Usage

```
glmm.bsreg(target, dataset, id, threshold = 0.05, wei = NULL, test = "testIndGLMMReg")
```

Arguments

target	The class variable. This can be a numerical vector with continuous data, binary or discrete valued data. It can also be a factor variable with two levels only.
dataset	The dataset; provide a numerical a matrix (columns = variables, rows = samples).
id	This is a numerical vector of the same size as target denoting the groups or the subjects.
threshold	Threshold (suitable values in [0,1]) for assessing p-values significance. Default value is 0.05.
wei	A vector of weights to be used for weighted regression. The default value is NULL.
test	This is for the type of regression to be used, "testIndGLMMReg", for Gaussian regression, "testIndGLMMLogistic" for logistic regression or "testIndGLMM-Pois" for Poisson regression.

Details

If the sample size is less than the number of variables a message will appear and no backward regression is performed.

Value

The output of the algorithm is S3 object including:

runtime	The run time of the algorithm. A numeric vector. The first element is the user time, the second element is the system time and the third element is the elapsed time.
info	A matrix with the non selected variables and their latest test statistics and logged p-values.
mat	A matrix with the selected variables and their latest statistics and p-values.
final	The final regression model.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

Eugene Demidenko (2013). Mixed Models: Theory and Applications with R, 2nd Edition. New Jersey: Wiley & Sons.

See Also

[fbed.glm.reg](#), [ebic.glm.bsreg](#), [MMPC.temporal](#)

Examples

```
## Not run:
require(lme4)
data(sleepstudy)
attach(sleepstudy)
x <- matrix(rnorm(180 * 20), ncol = 20) ## unrelated predictor variables
m1 <- glmm.bsreg(Reaction, x, Subject, wei = NULL, gam = NULL, type = "gaussian")
m2 <- MMPC.temporal(target = Reaction, group = Subject, dataset = x)

## End(Not run)
```

Backward selection regression for GLMM using the eBIC

Backward selection regression for GLMM using the eBIC

Description

Backward selection regression for GLMM using the eBIC

Usage

```
ebic.glmm.bsreg(target, dataset, id, wei = NULL, gam = NULL, test = "testIndGLMMReg")
```

Arguments

target	The class variable. This can be a numerical vector with continuous data, binary or discrete valued data. It can also be a factor variable with two levels only.
dataset	The dataset; provide a numerical a matrix (columns = variables, rows = samples).
id	This is a numerical vector of the same size as target denoting the groups or the subjects.
wei	A vector of weights to be used for weighted regression. The default value is NULL.
gam	In case the method is chosen to be "eBIC" one can also specify the <i>gamma</i> parameter. The default value is "NULL", so that the value is automatically calculated.
test	This is for the type of regression to be used, "testIndGLMMReg", for Gaussian regression, "testIndGLMMLogistic for logistic regression or "testIndGLMM-Pois" for Poisson regression.

Details

The algorithm is a variation of the usual forward selection. At every step, the most significant variable enters the selected variables set. In addition, only the significant variables stay and are further examined. The non significant ones are dropped. This goes until no variable can enter the set. The user has the option to redo this step 1 or more times (the argument K). In the end, a backward selection is performed to remove falsely selected variables.

Value

A list including:

runtime	The runtime required.
info	A matrix with the number of variables and the number of tests performed (or models fitted) at each round (value of K).
mat	A matrix with the selected variables and their eBIC.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

Borboudakis G. and Tsamardinos I. (2017). Forward-Backward Selection with Early Dropping. <https://arxiv.org/pdf/1705.10770.pdf>

Eugene Demidenko (2013). Mixed Models: Theory and Applications with R, 2nd Edition. New Jersey: Wiley & Sons.

See Also

[fbed.glm.reg](#), [glm.bsreg](#), [MMPC.temporal](#)

Examples

```
## Not run:
require(lme4)
data(sleepstudy)
attach(sleepstudy)
x <- matrix(rnorm(180 * 20), ncol = 20) ## unrelated predictor variables
m1 <- ebic.glm.bsreg(Reaction, x, Subject, wei = NULL, gam = NULL, type = "gaussian")
m2 <- MMPC.temporal(target = Reaction, group = Subject, dataset = x)

## End(Not run)
```

Backward selection regression using the eBIC

Backward selection regression using the eBIC

Description

Backward selection regression using the eBIC

Usage

```
ebic.bsreg(target, dataset, test = NULL, wei = NULL, gam = NULL)
```

Arguments

target	The class variable. Provide either a string, an integer, a numeric value, a vector, a factor, an ordered factor or a Surv object.
dataset	The dataset; provide either a data frame or a matrix (columns = variables, rows = samples).
test	The available tests: "testIndReg", "testIndPois", "testIndNB", "testIndLogistic", "testIndMMReg", "testIndBinom", "censIndCR", "censIndWR", "testIndBeta", "testIndZIP", "testIndGamma", "testIndNormLog" and "testIndTobit".
wei	A vector of weights to be used for weighted regression. The default value is NULL. It is not suggested when testIndMMReg is used.
gam	In case the method is chosen to be "eBIC" one can also specify the <i>gamma</i> parameter. The default value is "NULL", so that the value is automatically calculated.

Details

The algorithm is a variation of the usual forward selection. At every step, the most significant variable enters the selected variables set. In addition, only the significant variables stay and are further examined. The non significant ones are dropped. This goes until no variable can enter the set. The user has the option to redo this step 1 or more times (the argument K). In the end, a backward selection is performed to remove falsely selected variables.

Value

A list including:

runtime	The runtime required.
info	A matrix with the number of variables and the number of tests performed (or models fitted) at each round (value of K).
mat	A matrix with the selected variables and their eBIC.
back.rem	The variables removed in the backward phase.
back.n.tests	The number of models fitted in the backward phase.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

Borboudakis G. and Tsamardinos I. (2017). Forward-Backward Selection with Early Dropping. <https://arxiv.org/pdf/1705.10770.pdf>

See Also

[fs.reg](#), [bic.fsreg](#), [MMPC](#)

Examples

```
#simulate a dataset with continuous data
dataset <- matrix( runif(100 * 30, 1, 100), ncol = 30 )
#define a simulated class variable
target <- rt(100, 10)

a1 <- ebic.bsreg(target, dataset, test = "testIndReg")
target <- rpois(100, 10)
a2 <- ebic.bsreg(target, dataset, test = "testIndPois")
```

Backward selection with generalised linear regression models

Variable selection in generalised linear regression models with backward selection

Description

Variable selection in generalised linear regression models with backward selection

Usage

```
glm.bsreg(target, dataset, threshold = 0.05, wei = NULL, test = NULL)

gammabsreg(target, dataset, threshold = 0.05, wei = NULL)

normlog.bsreg(target, dataset, threshold = 0.05, wei = NULL)
```

Arguments

target	The class variable. Provide either an integer, a numeric value, or a factor. It can also be a matrix with two columns for the case of binomial regression. In this case, the first column is the number of successes and the second column is the number of trials. See also the Details.
dataset	The dataset; provide either a data frame or a matrix (columns = variables, rows = observations). In either case, only two cases are available, either all data are continuous, or categorical.
threshold	Threshold (suitable values in (0, 1)) for assessing p-values significance. Default value is 0.05.
wei	A vector of weights to be used for weighted regression. The default value is NULL. It is not suggested when robust is set to TRUE.
test	See CondIndTests for a list of the appropriate tests.

Details

This functions currently implements only linear, binomial, binary logistic and Poisson regression. If the sample size is less than the number of variables a message will appear and no backward regression is performed.

Value

The output of the algorithm is S3 object including:

<code>runtime</code>	The run time of the algorithm. A numeric vector. The first element is the user time, the second element is the system time and the third element is the elapsed time.
<code>info</code>	A matrix with the variables and their latest test statistics and p-values.
<code>mat</code>	A matrix with the selected variables and their latest test statistic and logged p-value.
<code>ci_test</code>	The conditional independence test used.
<code>final</code>	The final regression model.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

See Also

[fs.reg](#), [lm.fsreg](#), [bic.fsreg](#), [bic.glm.fsreg](#), [CondIndTests](#), [MMPC](#), [SES](#)

Examples

```
set.seed(123)

#simulate a dataset with continuous data
dataset <- matrix( runif(100 * 10, 1, 100), ncol = 10 )

#define a simulated class variable
target <- rpois(100, 10)
a <- glm.bsreg(target, dataset, threshold = 0.05)

target <- rbinom(100, 1, 0.6)
b <- glm.bsreg(target, dataset, threshold = 0.05)

target <- rgamma(100, 1, 2)
b1 <- gammabsreg(target, dataset, threshold = 0.05)
b2 <- normlog.bsreg(target, dataset, threshold = 0.05)
```

Bayesian Network construction using a hybrid of MMPC and PC

Bayesian Network construction using a hybrid of MMPC and PC

Description

Bayesian Network construction using a hybrid of MMPC and PC.

Usage

```
mmpc.or(x, max_k = 5, threshold = 0.01, test = "testIndFisher", backward = TRUE,
        symmetry = TRUE, ini.pvalue = NULL)
```

Arguments

<code>x</code>	A matrix with the variables. The user must know if they are continuous or if they are categorical. If you have a matrix with categorical data, i.e. 0, 1, 2, 3 where each number indicates a category, the minimum number for each variable must be 0. <code>data.frame</code> is also supported, as the dataset in this case is converted into a matrix.
<code>max_k</code>	The maximum conditioning set to use in the conditional independence test (see Details of SES or MMPC).
<code>threshold</code>	Threshold (suitable values in (0, 1)) for assessing p-values significance. Default value is 0.05.
<code>test</code>	The conditional independence test to use. Default value is "testIndFisher". This procedure allows for "testIndFisher", "testIndSPearman" for continuous variables and "gSquare" for categorical variables.
<code>backward</code>	If TRUE, the backward (or symmetry correction) phase will be implemented. This removes any falsely included variables in the parents and children set of the target variable. It calls the <code>link{mmpcbackphase}</code> for this purpose. For <code>perm.ses</code> and <code>wald.ses</code> this is not yet applicable.
<code>symmetry</code>	In order for an edge to be added, a statistical relationship must have been found from both directions. If you want this symmetry correction to take place, leave this boolean variable to TRUE. If you set it to FALSE, then if a relationship between Y and X is detected but not between X and Y, the edge is still added.
<code>ini.pvalue</code>	This is a list with the matrix of the univariate p-values. If you want to run <code>mmhc.skel</code> again, the univariate associations need not be calculated again.

Details

The MMPC is run on every variable. The backward phase (see Tsamardinos et al., 2006) can then take place. After all variables have been used, the matrix is checked for inconsistencies and they are corrected if you want. The "symmetry" argument. Do you want the edge to stay if it was discovered from both variables when they were considered as responses?

Value

A list including:

<code>ini.pvalue</code>	A matrix with the p-values of all pairwise univariate associations.
<code>kapa</code>	The maximum number of conditioning variables ever observed.
<code>ntests</code>	The number of tests MMPC (or SES) performed at each variable.
<code>info</code>	Some summary statistics about the edges, minimum, maximum, mean, median number of edges.

density	The number of edges divided by the total possible number of edges, that is $\#edges / n(n - 1)/2$, where n is the number of variables.
runtime	The run time of the skeleton phase of the algorithm. A numeric vector. The first element is the user time, the second element is the system time and the third element is the elapsed time.
runtime.or	The run time of the PC orientation rules. A numeric vector. The first element is the user time, the second element is the system time and the third element is the elapsed time.
Gini	The adjacency matrix. A value of 1 in $G[i, j]$ appears in $G[j, i]$ also, indicating that i and j have an edge between them.
G	The final adjacency matrix with the orientations. If $G[i, j] = 2$ then $G[j, i] = 3$. This means that there is an arrow from node i to node j . If $G[i, j] = G[j, i] = 0$; there is no edge between nodes i and j . If $G[i, j] = G[j, i] = 1$; there is an (undirected) edge between nodes i and j .
sepset	A list with the separating sets for every value of k .

Bear in mind that the values can be extracted with the \$ symbol, i.e. this is an S3 class output.

Author(s)

Michail Tsagris

R implementation and documentation: Giorgos Athineou <athineou@csd.uoc.gr> and Michail Tsagris <mtsagris@csd.uoc.gr>

References

Tsamardinos, Brown and Aliferis (2006). The max-min hill-climbing Bayesian network structure learning algorithm. *Machine learning*, 65(1), 31-78.

Spirtes P., Glymour C. and Scheines R. (2001). *Causation, Prediction, and Search*. The MIT Press, Cambridge, MA, USA, 3rd edition.

See Also

[pc.skel](#), [pc.or](#), [corfs.network](#)

Examples

```
y <- rdag2(1000, p = 20, nei = 3)
ind <- sample(1:20, 20)
x <- y$x[, ind]
a1 <- mmpc.or(x, max_k = 3, threshold = 0.01, test = "testIndFisher" )
b <- pc.skel( x, method = "comb.fast", alpha = 0.01 )
a2 <- pc.or(b)
```

Beta regression	<i>Beta regression</i>
-----------------	------------------------

Description

Beta regression.

Usage

```
beta.mod(target, dataset, wei = NULL, xnew= NULL)
```

Arguments

target	The target (dependent) variable. It must be a numerical vector with integers.
dataset	The independent variable(s). It can be a vector, a matrix or a dataframe with continuous only variables, a data frame with mixed or only categorical variables.
wei	A vector of weights to be used for weighted regression. The default value is NULL.
xnew	If you have new values for the predictor variables (dataset) whose target variable you want to predict insert them here. If you put the "dataset" or leave it NULL.

Details

The beta regression is fitted. The "beta.reg" is an internal wrapper function and is used for speed up purposes. It is not to be called directly by the user unless they know what they are doing.

Value

A list including:

be	The estimated coefficients of the model.
phi	The estimated precision parameter.
loglik	The log-likelihood of the regression model.
est	The estimated values if xnew is not NULL.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

Ferrari S.L.P. and Cribari-Neto F. (2004). Beta Regression for Modelling Rates and Proportions. Journal of Applied Statistics, 31(7): 799-815.

See Also

[beta.regs](#), [testIndBeta](#), [reg.fit](#), [ridge.reg](#)

Examples

```
y <- rbeta(500, 3, 5)
x <- matrix( rnorm(500 * 2), ncol = 2)
a1 <- beta.mod(y, x)
w <- runif(500)
a2 <- beta.mod(y, x, w)
```

BIC based forward selection

Variable selection in regression models with forward selection using BIC

Description

Variable selection in regression models with forward selection using BIC

Usage

```
bic.fsreg(target, dataset, test = NULL, wei = NULL, tol = 2, ncores = 1)
```

Arguments

target	The class variable. Provide either a string, an integer, a numeric value, a vector, a factor, an ordered factor or a Surv object. See also Details.
dataset	The dataset; provide either a data frame or a matrix (columns = variables, rows = samples). The data can be either euclidean, categorical or both.
test	The regression model to use. Available options are "testIndReg" for normal linear regression, "testIndBeta" for beta regression, "censIndCR" or "censIndWR" for Cox proportional hazards and Weibull regression respectively, "testIndLogistic" for binomial, multinomial or ordinal regression, "testIndPois" for poisson regression, "testIndNB" for negative binomial regression, "testIndZIP" for zero inflated poisson regression, "testIndRQ" for quantile regression, "testIndGamma" for gamma regression, "testIndNormLog" for linear regression with the log-link (non negative data), "testIndTobit" for Tobit regression. If you want to use multinomial or ordinal logistic regression, make sure your target is factor. See also SES and CondIndTests for the tests.
wei	A vector of weights to be used for weighted regression. The default value is NULL. It is not suggested when robust is set to TRUE.
tol	The difference bewtween two successive values of the stopping rule. By default this is is set to 2. If for example, the BIC difference between two successive models is less than 2, the process stops and the last variable, even though significant does not enter the model.

`ncores` How many cores to use. This plays an important role if you have tens of thousands of variables or really large sample sizes and tens of thousands of variables and a regression based test which requires numerical optimisation. In other cases it will not make a difference in the overall time (in fact it can be slower). The parallel computation is used in the first step of the algorithm, where univariate associations are examined, those take place in parallel. We have seen a reduction in time of 50% with 4 cores in comparison to 1 core. Note also, that the amount of reduction is not linear in the number of cores.

Details

If the current 'test' argument is defined as NULL or "auto" and the `user_test` argument is NULL then the algorithm automatically selects the best test based on the type of the data. Particularly:

- if target is a factor, the multinomial or the binary logistic regression is used. If the target has two values only, binary logistic regression will be used.
- if target is an ordered factor, the ordinal regression is used.
- if target is a numerical vector or a matrix with at least two columns (multivariate) linear regression is used.
- if target is discrete numerical (counts), the poisson regression conditional independence test is used. If there are only two values, the binary logistic regression is to be used.
- if target is a `Surv` object, the Survival conditional independence test (Cox regression) is used.

Value

The output of the algorithm is an S3 object including:

<code>runtime</code>	The run time of the algorithm. A numeric vector. The first element is the user time, the second element is the system time and the third element is the elapsed time.
<code>mat</code>	A matrix with the variables and their latest test statistics and p-values.
<code>info</code>	A matrix with the selected variables, and the BIC of the model with that and all the previous variables.
<code>ci_test</code>	The conditional independence test used.
<code>final</code>	The final regression model.

Author(s)

Michail Tsagris

R implementation and documentation: Giorgos Athineou <athineou@csd.uoc.gr> Michail Tsagris <mtsagris@csd.uoc.gr>

References

Tsamardinos I., Aliferis C. F. and Statnikov, A. (2003). Time and sample efficient discovery of Markov blankets and direct causal relations. In Proceedings of the 9th ACM SIGKDD international conference on Knowledge discovery and data mining (pp. 673-678).

See Also

[glm.fsreg](#), [lm.fsreg](#), [bic.glm.fsreg](#), [CondIndTests](#), [MMPC](#), [SES](#)

Examples

```
set.seed(123)
dataset <- matrix( runif(1000 * 20, 1, 100), ncol = 20 )
target <- 3 * dataset[, 10] + 2 * dataset[, 15] + 3 * dataset[, 20] + rnorm(1000, 0, 5)

a1 <- bic.fsreg(target, dataset, tol = 4, ncores = 1, test = "testIndReg" )
a3 <- MMPC(target, dataset, ncores = 1)
target <- round(target)
b1 <- bic.fsreg(target, dataset, tol = 2, ncores = 1, test = "testIndReg" )
```

BIC based forward selection with generalised linear models

Variable selection in generalised linear models with forward selection based on BIC

Description

Variable selection in generalised linear models with forward selection based on BIC

Usage

```
bic.glm.fsreg( target, dataset, wei = NULL, tol = 0, ncores = 1)
bic.mm.fsreg( target, dataset, wei = NULL, tol = 0, ncores = 1)
bic.gammafsreg(target, dataset, wei = NULL, tol = 0, ncores = 1)
bic.normlog.fsreg(target, dataset, wei = NULL, tol = 0, ncores = 1)
```

Arguments

target	The class variable. It can be either a vector with binary data (binomial regression), counts (poisson regression). If none of these is identified, linear regression is used.
dataset	The dataset; provide either a data frame or a matrix (columns = variables, rows = samples). These can be continuous and or categorical.
wei	A vector of weights to be used for weighted regression. The default value is NULL. It is not suggested when robust is set to TRUE.
tol	The difference between two successive values of BIC. By default this is set to 2. If for example, the BIC difference between two successive models is less than 2, the process stops and the last variable, even though significant does not enter the model.

`ncores` How many cores to use. This plays an important role if you have tens of thousands of variables or really large sample sizes and tens of thousands of variables and a regression based test which requires numerical optimisation. In other cases `cammb` it will not make a difference in the overall time (in fact it can be slower). The parallel computation is used in the first step of the algorithm, where univariate associations are examined, those take place in parallel. We have seen a reduction in time of 50% with 4 cores in comparison to 1 core. Note also, that the amount of reduction is not linear in the number of cores.

Details

Forward selection via the BIC is implemented. A variable which results in a reduction of BIC will be included, until the reduction is below a threshold set by the user (argument "tol").

Value

The output of the algorithm is S3 object including:

<code>runtime</code>	The run time of the algorithm. A numeric vector. The first element is the user time, the second element is the system time and the third element is the elapsed time.
<code>mat</code>	A matrix with the variables and their latest test statistics and p-values.
<code>info</code>	A matrix with the selected variables, and the BIC of the model with that and all the previous variables.
<code>ci_test</code>	The conditional independence test used.
<code>final</code>	The final regression model.

Author(s)

Michail Tsagris

R implementation and documentation: Giorgos Athineou <athineou@csd.uoc.gr> Michail Tsagris <mtsagris@csd.uoc.gr>

See Also

[fs.reg](#), [lm.fsreg](#), [bic.fsreg](#), [CondIndTests](#), [MMPC](#), [SES](#)

Examples

```
set.seed(123)
dataset <- matrix( runif(1000 * 50, 1, 100), ncol = 50 )
target <- 3 * dataset[, 10] + 2 * dataset[, 20] + 3 * dataset[, 30] + rnorm(1000, 0, 5)
a1 <- bic.glm.fsreg(target, dataset, tol = 2, ncores = 1 )
a2 <- bic.glm.fsreg( round(target), dataset, tol = 2, ncores = 1 )
y <- target ; me <- median(target) ; y[ y < me ] <- 0 ; y[ y >= me ] <- 1
a3 <- bic.glm.fsreg( y, dataset, tol = 2, ncores = 1 )
```

Calculation of the constant and slope for each subject over time
Calculation of the constant and slope for each subject over time

Description

Calculation of the constant and slope for each subject over time.

Usage

```
group.mvbetas(x, id, reps)
```

Arguments

x	The dataset; provide a numerical a matrix. This should contain longitudinal data. Each variable is a column, and rows are longitudinal data.
id	This is a numerical vector denoting the subjects. Its length must be equal to the number of rows of the x matrix.
reps	This is a numerical vector with the time points. Its length must be equal to the number of rows of the x matrix.

Details

This function is used internally in [SES](#) and [MMPC](#) and does calculations required bys the first step of the **Static-Longitudinal** scenario of Tsagris, Lagani and Tsamardinos (2018). The measurements of each subject are regressed againt time. So, for each subject, we get the constant and interecept over time and this is repated for very feature.

Value

A matrix. The first r (= length(unique(id)), the nubmer of subjects) rows contain the constants and the other r rows contain the slopes.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

Tsagris M., Lagani V., & Tsamardinos I. (2018). Feature selection for high-dimensional temporal data. *BMC bioinformatics*, 19(1), 17.

McCullagh, Peter, and John A. Nelder. *Generalized linear models*. CRC press, USA, 2nd edition, 1989.

See Also

[fbed.gEE.reg](#), [glmm.bsreg](#), [MMPC.temporal](#)

Examples

```
## assume these are longitudinal data, each column is a variable (or feature)
x <- matrix( rnorm(100 * 100), ncol = 100 )
id <- rep(1:20, each = 5) ## 20 subjects
reps <- rep( seq(4, 12, by = 2), 20) ## 5 time points for each subject
a <- group.mvbetas(x, id, reps)
dim(a) ## 5 200
## these are the regression coefficients of the first subject's values on the
## reps (which is assumed to be time in this example)
a[c(1, 21), 1]
coef( lm( x[id == 1, 1] ~ reps[1:5] ) )
```

Certificate of exclusion from the selected variables set using SES or MMPC
*Certificate of exclusion from the selected variables set using SES or
 MMPC*

Description

Information on why one ore more variables were not selected.

Usage

```
certificate.of.exclusion(xIndex, sesObject = NULL, mmpcObject = NULL)
```

Arguments

<code>xIndex</code>	A numerical vector with the indices of the predictor variables.
<code>sesObject</code>	If you ran SES, <code>wald.ses</code> or <code>perm.ses</code> , give the whole SES object here, otherwise leave it NULL.
<code>mmpcObject</code>	If you ran MMPC, <code>wald.mmpc</code> or <code>prm.mmpc</code> , give the whole SES object here, otherwise leave it NULL.

Value

A list with the conditioning variables (if any), the test statistic and the logarithm of the p-value. In case a variable has been selected a message appears.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

See Also[MMPC](#)**Examples**

```

set.seed(123)
require("hash", quietly = TRUE)
#simulate a dataset with continuous data
dataset <- matrix(runif(1000 * 200, 1, 100), ncol = 200)
#define a simulated class variable
target <- 3 * dataset[, 10] + 2 * dataset[, 100] + 3 * dataset[, 20] + rnorm(1000, 0, 5)
# define some simulated equivalences
dataset[, 15] <- dataset[, 10] + rnorm(1000, 0, 2)
dataset[, 150] <- dataset[, 100] + rnorm(1000, 0, 2)
dataset[, 130] <- dataset[, 100] + rnorm(1000, 0, 2)
# run the SES algorithm
mod1 <- SES(target, dataset, max_k = 5, threshold = 0.05, test = "testIndFisher",
hash = TRUE, hashObject = NULL);
mod2 <- MMPC(target, dataset, max_k = 5, threshold = 0.05, test = "testIndFisher",
hash = TRUE, hashObject = NULL);
certificate.of.exclusion(c(10,15,30,45,20), mod1)
certificate.of.exclusion(c(10,15,30,45,20), NULL, mod2)

```

Check Markov equivalence of two DAGs

Check Markov equivalence of two DAGs

Description

Check Markov equivalence of two DAGs.

Usage

```
equivdags(g1, g2)
```

Arguments

g1	The matrix of a DAG or a partially directed graph as produced from pc.or or any other algorithm.
g2	The matrix of a DAG or a partially directed graph as produced from pc.or or any other algorithm.

Details

Two DAGs are Markov equivalent if a) they have the same adjancencies (regardlsee of the mark, arrowhead, tail or nothing) and b) they have the same unshielded colliders.

Value

A list including:

`apofasi` A boolean variable, TRUE of FALSE.
`mes` A message specifying the result, the dimensions of the adjacency matrices do not match for example, or the number of adjancencies is not the same, they do not share the same unshilded colliders, or they are Markov equivalent.

Author(s)

Michail Tsagris

R implementation and documentation: Giorgos Athineou <athineou@csd.uoc.gr> and Michail Tsagris <mtsagris@csd.uoc.gr>

References

Tsamardinos, Brown and Aliferis (2006). The max-min hill-climbing Bayesian network structure learning algorithm. *Machine learning*, 65(1), 31-78.

See Also

[pc.or](#), [mmhc.skel](#), [pc.con](#)

Examples

```
y <- rdag(1000, 10, 0.3)
tru <- y$G
x <- y$x
mod <- pc.con(x)

eg <- dag2eg(y$G) ## make it essential graph first
est <- pc.or(mod)$G

equivdags(est, tru)
```

Check whether a directed graph is acyclic

Check whether a directed graph is acyclic

Description

Check whether a directed graph is acyclic.

Usage

```
is.dag(dag)
```

Arguments

`dag` A square matrix representing a directed graph which contains either 0, 1 or 0, 2, and 3. In the first case where $G[i, j] = 1$, means there is an arrow from node i to node j . In the second case $G[i, j] = 2$ and $G[j, i] = 3$ means that there is an arrow from node i to node j , where the 2 indicates the arrowhead and the 3 indicates the arrowtail.

Details

The topological sort is performed. If it cannot be performed, NAs are returned. Hence, the functions checks for NAs.

Value

A logical value, TRUE if the matrix represents a DAG and FALSE otherwise.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

Chickering, D.M. (1995). A transformational characterization of equivalent Bayesian network structures. Proceedings of the 11th Conference on Uncertainty in Artificial Intelligence, Montreal, Canada, 87-98.

See Also

[topological_sort](#), [dag2eg](#), [pc.or](#)

Examples

```
# simulate a dataset with continuous data
# simulate a dataset with continuous data
G <- rdag(100, 40, 0.3)$G
is.dag(G) ## TRUE
```

Description

Currently the **MXM** package supports numerous tests for different types of target (dependent) and predictor (independent) variables. The target variable can be of continuous, discrete, categorical and of survival type. As for the predictor variables, they can be continuous, categorical or mixed.

The **testIndFisher** and the **gSquare** tests have two things in common. They do not use a model implicitly (i.e. estimate some beta coefficients), even though there is an underlying assumed one. Secondly they are pure tests of independence (again, with assumptions required).

As for the other tests, they share one thing in common. For all of them, two parametric models must be fit. The null model containing the conditioning set of variables alone and the alternative model containing the conditioning set and the candidate variable. The significance of the new variable is assessed via a log-likelihood ratio test with the appropriate degrees of freedom. All of these tests which are available for SES and MMPC are summarized in the below table.

Target variable	Predictor variables	Available tests	Short explanation
Continuous	Continuous	testIndFisher	Partial correlation
Continuous	Continuous	testIndMMFisher	Robust partial correlation
Continuous	Continuous	testIndSpearman	Partial correlation
Continuous	Mixed	testIndMMReg	MM regression
Continuous	Mixed	testIndRQ	Median regression
Proportions	Continuous	testIndFisher	Partial correlation
Proportions	Continuous	testIndMMFisher	Robust partial correlation
Proportions	Continuous	testIndSpearman	Partial correlation
Proportions	Mixed	testIndReg	Linear regression
Proportions	Mixed	testIndMMReg	MM regression
Proportions	Mixed	testIndRQ	Median regression
Proportions	Mixed	testIndBeta	Beta regression
Strictly positive	Mixed	testIndIGreg	Inverse Gaussian regression
Strictly positive	Mixed	testIndGamma	Gamma regression
Non negative	Mixed	testIndNormLog	Gaussian regression with log link
Strictly Positive	Mixed	censIndWR	Weibull regression
Successes \& totals	Mixed	testIndBinom	Binomial regression
Discrete	Mixed	testIndPois	Poisson regression
Discrete	Mixed	testIndZIP	Zero Inflated Poisson regression
Discrete	Mixed	testIndNB	Negative binomial regression
Discrete	Mixed	testIndQPois	Quasi Poisson regression
Factor with two levels or binary	Mixed	testIndLogistic	Binary logistic regression
Factor with two levels or binary	Mixed	testIndQBinom	Quasi binomial regression
Factor with more than two levels (unordered)	Mixed	testIndMultinom	Multinomial logistic regression
Factor with more than two levels (ordered)	Mixed	testIndOrdinal	Ordinal logistic regression
Categorical	Categorical	gSquare	G-squared test of independence
Categorical	Categorical	testIndMultinom	Multinomial logistic regression
Categorical	Categorical	testIndOrdinal	Ordinal logistic regression

Survival	Mixed	censIndCR	Cox regression
Survival	Mixed	censIndWR	Weibull regression
Survival	Mixed	censIndER	Exponential regression
Left censored	Mixed	testIndTobit	Tobit regression
Case-control	Mixed	testIndClogit	Conditional logistic regression
Multivariate continuous	Mixed	testIndMVreg	Multivariate linear regression
Compositional data (no zeros)	Mixed	testIndMVreg after multivariate logit transformation	Multivariate linear regression
Longitudinal	Continuous	testIndLMMReg	Linear mixed models
Binary	Continuous	testIndLMMLogistic	Logistic mixed regression
Discrete	Continuous	testIndLMMPois	Poisson mixed regression

Details

These tests can be called by SES, MMPC, wald.mmpc or individually by the user. In all regression cases, there is an option for weights.

Log-likelihood ratio tests

1. **testIndFisher**. This is a standard test of independence when both the target and the set of predictor variables are continuous (continuous-continuous). When the joint multivariate normality of all the variables is assumed, we know that if a correlation is zero this means that the two variables are independent. Moving in this spirit, when the partial correlation between the target variable and the new predictor variable conditioning on a set of (predictor) variables is zero, then we have evidence to say they are independent as well. An easy way to calculate the partial correlation between the target and a predictor variable conditioning on some other variables is to regress the both the target and the new variable on the conditioning set. The correlation coefficient of the residuals produced by the two regressions equals the partial correlation coefficient. If the robust option is selected, the two aforementioned regression models are fitted using M estimators (Marona et al., 2006). If the target variable consists of proportions or percentages (within the (0, 1) interval), the logit transformation is applied beforehand.
2. **testIndSpearman**. This is a non-parametric alternative to **testIndFisher** test. It is a bit slower than its competitor, yet very fast and suggested when normality assumption breaks down or outliers are present. In fact, within SES, what happens is that the ranks of the target and of the dataset (predictor variables) are computed and the **testIndSpearman** is applied. This is faster than applying Fisher with M estimators as described above. If the target variable consists of proportions or percentages (within the (0, 1) interval), the logit transformation is applied beforehand.
3. **testIndReg**. In the case of target-predictors being continuous-mixed or continuous-categorical, the suggested test is via the standard linear regression. In this case, two linear regression models are fitted. One with the conditioning set only and one with the conditioning set plus the new variable. The significance of the new variable is assessed via the F test, which calculates the residual sum of squares of the two models. The reason for the F test is because the new variable may be categorical and in this case the t test cannot be used. It makes sense to say, that this test can be used instead of the **testIndFisher**, but it will be slower. If the robust option is

selected, the two models are fitted using M estimators (Marona et al. 2006). If the target variable consists of proportions or percentages (within the (0, 1) interval), the logit transformation is applied beforehand.

4. **testIndRQ**. An alternative to **testIndReg** for the case of continuous-mixed (or continuous-continuous) variables is the **testIndRQ**. Instead of fitting two linear regression models, which model the expected value, one can choose to model the median of the distribution (Koenker, 2005). The significance of the new variable is assessed via a rank based test calibrated with an F distribution (Gutenbrunner et al., 1993). The reason for this is that we performed simulation studies and saw that this type of test attains the type I error in contrast to the log-likelihood ratio test. The benefit of this regression is that it is robust, in contrast to the classical linear regression. If the target variable consists of proportions or percentages (within the (0, 1) interval), the logit transformation is applied beforehand.
5. **testIndBeta**. When the target is proportion (or percentage, i.e., between 0 and 1, not inclusive) the user can fit a regression model assuming a beta distribution. The predictor variables can be either continuous, categorical or mixed. The procedure is the same as in the **testIndReg** case.
6. **Alternatives to testIndBeta**. Instead of **testIndBeta** the user has the option to choose all the previous to that mentioned tests by transforming the target variable with the logit transformation. In this way, the support of the target becomes the whole of \mathbb{R}^d and then depending on the type of the predictors and whether a robust approach is required or not, there is a variety of alternative to beta regression tests.
7. **testIndIGreg**. When you have non negative data, i.e. the target variable takes positive values (including 0), a suggested regression is based on the the inverse gaussian distribution. The link function is not the inverse of the square root as expected, but the logarithm. This is to ensure that the fitted values will be always be non negative. The predictor variables can be either continuous, categorical or mixed. The significance between the two models is assessed via the log-likelihood ratio test. Alternatively, the user can use the Weibull regression (**censIndWR**), gamma regression (**testIndGamma**) or Gaussian regression with log link (**testIndNormLog**).
8. **testIndGamma**. This is an alternative to **testIndIGreg**.
9. **testIndNormLog**. This is a second alternative to **testIndIGreg**.
10. **testIndPois**. When the target is discrete, and in specific count data, the default test is via the Poisson regression. The predictor variables can be either continuous, categorical or mixed. The procedure is the same as in all the previously regression model based tests, i.e. the log-likelihood ratio test is used to assess the conditional independence of the variable of interest.
11. **testIndNB**. As an alternative to the Poisson regression, we have included the Negative binomial regression to capture cases of overdispersion. The predictor variables can be either continuous, categorical or mixed.
12. **testIndZIP**. When the number of zeros is more than expected under a Poisson model, the zero inflated poisson regression is to be employed. The predictor variables can be either continuous, categorical or mixed.
13. **testIndLogistic**. When the target is categorical with only two outcomes, success or failure for example, then a binary logistic regression is to be used. Whether regression or classification is the task of interest, this method is applicable. The advantage of this over a linear or quadratic discriminant analysis is that it allows for categorical predictor variables as well and for mixed types of predictors.

14. **testIndMultinom.** If the target has more than two outcomes, but it is of nominal type, there is no ordering of the outcomes, multinomial logistic regression will be employed. Again, this regression is suitable for classification purposes as well and it allows for categorical predictor variables.
15. **testIndOrdinal.** This is a special case of multinomial regression, in which case the outcomes have an ordering, such as **not satisfied, neutral, satisfied**. The appropriate method is ordinal logistic regression.
16. **testIndBinom.** When the target variable is a matrix of two columns, where the first one is the number of successes and the second one is the number of trials, binomial regression is to be used.
17. **gSquare.** If all variables, both the target and predictors are categorical the default test is the G-square test of independence. It is similar to the chi-squared test of independence, but instead of using the chi-squared metric between the observed and estimated frequencies in contingency tables, the Kullback-Leibler divergence of the observed from the estimated frequencies is used. The asymptotic distribution of the test statistic is a chi-squared distribution on some appropriate degrees of freedom. The target variable can be either ordered or unordered with two or more outcomes.
18. **Alternatives to gSquare.** An alternative to the **gSquare** test is the **testIndLogistic**. Depending on the nature of the target, binary, un-ordered multinomial or ordered multinomial the appropriate regression model is fitted.
19. **censIndCR.** For the case of time-to-event data, a Cox regression model is employed. The predictor variables can be either continuous, categorical or mixed. Again, the log-likelihood ratio test is used to assess the significance of the new variable.
20. **censIndWR.** A second model for the case of time-to-event data, a Weibull regression model is employed. The predictor variables can be either continuous, categorical or mixed. Again, the log-likelihood ratio test is used to assess the significance of the new variable. Unlike the semi-parametric Cox model, the Weibull model is fully parametric.
21. **censIndER.** A third model for the case of time-to-event data, an exponential regression model is employed. The predictor variables can be either continuous, categorical or mixed. Again, the log-likelihood ratio test is used to assess the significance of the new variable. This is a special case of the Weibull model.
22. **testIndClogit.** When the data come from a case-control study, the suitable test is via conditional logistic regression.
23. **testIndMVreg.** In the case of multivariate continuous targets, the suggested test is via a multivariate linear regression. The target variable can be compositional data as well. These are positive data, whose vectors sum to 1. They can sum to any constant, as long as it the same, but for convenience reasons we assume that they are normalised to sum to 1. In this case the additive log-ratio transformation (multivariate logit transformation) is applied beforehand.
24. **testIndGLMM.** In the case of a longitudinal or clustered targets (continuous, proportions, binary or counts), the suggested test is via a (generalised) linear mixed model.

Wald based tests

The available tests for `wald.ses` and `wald.mmpc` are listed below. Note, that only continuous predictors are allowed.

Target variable	Available tests	Short explanation	
Continuous	waldMMReg	MM regression	
Proportions	waldMMReg	MM regression after logit transformation	
Proportions	waldBeta	Beta regression	
Non negative	waldIGreg	Inverse Gaussian regression	
Strictly positive	waldGamma	Gamma regression	
Non negative	waldNormLog	Gaussian regression with log link	
Successes \& totals	testIndBinom	Binomial regression	
Discrete	waldPois	Poisson regression	
Discrete	waldSpeedPois	Poisson regression	
Discrete	waldZIP	Zero Inflated	Poisson regression
Discrete	waldNB	Negative binomial regression	
Factor with two levels or binary	waldLogistic	Logistic regression	
Factor with more than two levels (ordered)	waldOrdinal	Ordinal logistic regression	
Left censored	waldTobit	Tobit regression	
Case-control	Mixed	testIndClogit	Conditional logistic regression
Survival	waldCR	Cox regression	
Survival	waldWR	Weibull regression	
Survival	waldER	Exponential regression	

Permutation based tests

The available tests for perm.ses and perm.mmpc are listed below. Note, that only continuous predictors are allowed.

Target variable	Available tests	Short explanation	
Continuous	permFisher	Pearson correlation	
Continuous	permMMFisher	Robust Pearson correlation	
Continuous	permDcor	Distance correlation	
Continuous	permReg	Linear regression	
Proportions	permReg	Linear regression after logit transformation	
Proportions	permBeta	Beta regression	
Non negative	permIGreg	Inverse Gaussian regression	
Strictly positive	permGamma	Gamma regression	
Non negative	permNormLog	Gaussian regression with log link	
Non negative	permWR	Weibull regression	
Successes \& totals	permBinom	Binomial regression	
Discrete	permPois	Poisson regression	
Discrete	permZIP	Zero Inflated	Poisson regression
Discrete	permNB	Negative binomial regression	
Factor with two levels or binary	permLogistic	Binary logistic regression	

Factor with more than two levels (nominal)	permMultinom	Multinomial logistic regression
Factor with more than two levels (ordered)	permOrdinal	Ordinal logistic regression
Left censored	permTobit	Tobit regression
Survival	permCR	Cox regression
Survival	permWR	Weibull regression
Survival	permER	Exponential regression

Author(s)

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Conditional independence test for binary, categorical or ordinal data

Conditional independence test for binary, categorical or ordinal class variables

Description

The main task of this test is to provide a p-value PVALUE for the null hypothesis: feature 'X' is independent from 'TARGET' given a conditioning set CS. The pvalue is calculated by comparing a logistic model based on the conditioning set CS against a model whose regressor are both X and CS. The comparison is performed through a chi-square test with the appropriate degrees of freedom on the difference between the deviances of the two models.

Usage

```
testIndLogistic(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL)
```

```
testIndMultinom(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL)
```

```
testIndOrdinal(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL)
```

```
testIndQBinom(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL)
```

```
permLogistic(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL,
threshold = 0.05, R = 999)
```

```
permMultinom(target, dataset, xIndex, csIndex, wei = NULL,
```

```
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL,
threshold = 0.05, R = 999)
```

```
permOrdinal(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL,
threshold = 0.05, R = 999)
```

```
waldLogistic(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL)
```

```
waldQBinom(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL)
```

```
waldOrdinal(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL)
```

Arguments

target	A numeric vector containing the values of the target variable. For the "testInd-Logistic" this can either be a binary numerical variable or a factor variable. The factor variable can have two values (binary logistic regression), more than two values (multinomial logistic regression) or it can be an ordered factor with more than two values (ordinal regression). The last one is for example, factor(x, ordered = TRUE). The "waldBinary" is the Wald test version of the binary logistic regression. The "waldOrdinal" is the Wald test version of the ordinal regression.
dataset	A numeric matrix or data frame, in case of categorical predictors (factors), containing the variables for performing the test. Rows as samples and columns as features. In the cases of "waldBinary", "waldOrdinal" this is strictly a matrix.
xIndex	The index of the variable whose association with the target we want to test.
csIndex	The indices of the variables to condition on. If you have no variables set this equal to 0.
wei	A vector of weights to be used for weighted regression. The default value is NULL.
univariateModels	Fast alternative to the hash object for univariate test. List with vectors "pvalues" (p-values), "stats" (statistics) and "flags" (flag = TRUE if the test was succesful) representing the univariate association of each variable with the target. Default value is NULL.
hash	A boolean variable which indicates whether (TRUE) or not (FALSE) to use the hash-based implementation of the statistics of SES. Default value is FALSE. If TRUE you have to specify the stat_hash argument and the pvalue_hash argument.
stat_hash	A hash object (hash package required) which contains the cached generated statistics of a SES run in the current dataset, using the current test.
pvalue_hash	A hash object (hash package required) which contains the cached generated p-values of a SES run in the current dataset, using the current test.

threshold	Threshold (suitable values in [0,1]) for assessing p-values significance.
R	The number of permutations, set to 999 by default. There is a trick to avoid doing all permutations. As soon as the number of times the permuted test statistic is more than the observed test statistic is more than 50 (if threshold = 0.05 and R = 999), the p-value has exceeded the significance level (threshold value) and hence the predictor variable is not significant. There is no need to continue do the extra permutations, as a decision has already been made.

Details

The `testIndQBinom` does quasi binomial regression which is suitable for binary targets and proportions including 0 and 1.

If `hash = TRUE`, `testIndLogistic` requires the arguments `'stat_hash'` and `'pvalue_hash'` for the hash-based implementation of the statistic test. These hash Objects are produced or updated by each run of `SES` (if `hash == TRUE`) and they can be reused in order to speed up next runs of the current statistic test. If `"SESoutput"` is the output of a `SES` run, then these objects can be retrieved by `SESoutput@hashObject$stat_hash` and the `SESoutput@hashObject$pvalue_hash`.

Important: Use these arguments only with the same dataset that was used at initialization. For all the available conditional independence tests that are currently included on the package, please see `"?CondIndTests"`.

The log-likelihood ratio test used in `"testIndLogistic"` requires the fitting of two models. The Wald test used in `waldBinary` and `waldOrdinal` requires fitting of only one model, the full one. The significance of the variable is examined only. Only continuous (or binary) predictor variables are currently accepted in this test.

Value

A list including:

<code>pvalue</code>	A numeric value that represents the logarithm of the generated p-value.
<code>stat</code>	A numeric value that represents the generated statistic.
<code>stat_hash</code>	The current hash object used for the statistics. See argument <code>stat_hash</code> and details. If argument <code>hash = FALSE</code> this is <code>NULL</code> .
<code>pvalue_hash</code>	The current hash object used for the p-values. See argument <code>stat_hash</code> and details. If argument <code>hash = FALSE</code> this is <code>NULL</code> .

Note

This test uses the function `multinom` (package `nnet`) for multinomial logistic regression, the function `clm` (package `ordinal`) for ordinal logit regression and the function `glm` (package `stats`) for binomial regression.

Author(s)

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See Also

[SES](#), [gSquare](#), [CondIndTests](#)

Examples

```
#require(nnet)
#require(ordinal)

#simulate a dataset with categorical data
dataset_m <- matrix( sample(c(0, 1, 2), 50 * 100, replace = TRUE), ncol = 50)
#initialize categorical target
target_m <- dataset_m[, 50]
#remove target from the dataset
dataset_m <- dataset_m[, -50]

#run the conditional independence test for the nominal class variable
results_m <- testIndMultinom(target_m, dataset_m, xIndex = 44, csIndex = c(10, 20) )
results_m

#run the SES algorithm using the testIndLogistic conditional independence test
#for the nominal class variable
sesObject <- SES(as.factor(target_m), dataset_m, max_k = 3, threshold = 0.05,
test = "testIndMultinom");
#print summary of the SES output
summary(sesObject);
# plot the SES output
# plot(sesObject, mode = "all");

#####
#run the conditional independence test for the ordinal class variable
results_o <- testIndOrdinal( factor(target_m, ordered = TRUE), dataset_m,
xIndex = 44, csIndex = c(10, 20) )
results_o

#run the SES algorithm using the testIndLogistic conditional independence test
#for the ordinal class variable
sesObject <- SES(factor(target_m, ordered=TRUE), dataset_m, max_k = 3 ,
threshold = 0.05, test = "testIndOrdinal");
#print summary of the SES output
summary(sesObject);
# plot the SES output
# plot(sesObject, mode = "all");
```

```
#####

#simulate a dataset with binary data
dataset_b <- matrix(sample(c(0,1),50 * 60, replace = TRUE), ncol = 50)
#initialize binary target
target_b <- dataset_b[, 50]
#remove target from the dataset
dataset_b <- dataset_b[, -50]

#run the conditional independence test for the binary class variable
results_b <- testIndLogistic( target_b, dataset_b, xIndex = 44, csIndex = c(10, 20) )
results_b

#run the SES algorithm using the testIndLogistic conditional independence test
#for the binary class variable
sesObject <- SES(target_b, dataset_b, max_k = 3, threshold = 0.05,
test = "testIndLogistic");
#print summary of the SES output
summary(sesObject);
```

Conditional independence test for case control data

Conditional independence test based on conditional logistic regression for case control studies

Description

The main task of this test is to provide a p-value PVALUE for the null hypothesis: feature 'X' is independent from 'TARGET' given a conditioning set CS. The pvalue is calculated by comparing a conditional logistic regression model based on the conditioning set CS against a model whose regressors are both X and CS. The comparison is performed through a chi-square test with the appropriate degrees of freedom on the difference between the deviances of the two models. This is suitable for a case control design

Usage

```
testIndClogit(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL)
```

```
permClogit(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL,
threshold = 0.05, R = 999)
```

Arguments

target A matrix with two columns, the first one must be 0 and 1, standing for 0 = control and 1 = case. The second column is the id of the patients. A numerical variable, for example c(1,2,3,4,5,6,7,1,2,3,4,5,6,7).

dataset	A numeric matrix or a data.frame in case of categorical predictors (factors), containing the variables for performing the test. Rows as samples and columns as features.
xIndex	The index of the variable whose association with the target we want to test.
csIndex	The indices of the variables to condition on. If you have no variables set this equal to 0.
wei	A vector of weights to be used for weighted regression. The default value is NULL and it should stay NULL as weights are ignored at the conditional logistic regression. See the survival package for more information about conditional logistic regression.
univariateModels	Fast alternative to the hash object for univariate test. List with vectors "pvalues" (p-values), "stats" (statistics) and "flags" (flag = TRUE if the test was succesful) representing the univariate association of each variable with the target. Default value is NULL.
hash	A boolean variable which indicates whether (TRUE) or not (FALSE) to use the hash-based implementation of the statistics of SES. Default value is FALSE. If TRUE you have to specify the stat_hash argument and the pvalue_hash argument.
stat_hash	A hash object (hash package required) which contains the cached generated statistics of a SES run in the current dataset, using the current test.
pvalue_hash	A hash object (hash package required) which contains the cached generated p-values of a SES run in the current dataset, using the current test.
threshold	Threshold (suitable values in [0,1]) for assessing p-values significance.
R	The number of permutations, set to 999 by default. There is a trick to avoid doing all permutations. As soon as the number of times the permuted test statistic is more than the observed test statistic is more than 50 (if threshold = 0.05 and R = 999), the p-value has exceeded the significance level (threshold value) and hence the predictor variable is not significant. There is no need to continue do the extra permutations, as a decision has already been made.

Details

If hash = TRUE, testIndClogit requires the arguments 'stat_hash' and 'pvalue_hash' for the hash-based implementation of the statistic test. These hash Objects are produced or updated by each run of SES (if hash == TRUE) and they can be reused in order to speed up next runs of the current statistic test. If "SESoutput" is the output of a SES run, then these objects can be retrieved by SESoutput@hashObject\$stat_hash and the SESoutput@hashObject\$pvalue_hash.

Important: Use these arguments only with the same dataset that was used at initialization. For all the available conditional independence tests that are currently included on the package, please see "?CondIndTests".

This is for case control studies. The log-likelihood for a conditional logistic regression model equals the log-likelihood from a Cox model with a particular data structure. When a well tested Cox model routine is available many packages use this "trick" rather than writing a new software routine from scratch, and this is what the "clogit" function in the "survival" package does. In detail, a stratified

Cox model with each case/control group assigned to its own stratum, time set to a constant, status of 1=case 0=control, and using the exact partial likelihood has the same likelihood formula as a conditional logistic regression. The "clogit" routine creates the necessary dummy variable of times (all 1) and the strata, then calls the function "coxph".

Note that this function is a bit sensitive and prone to breaking down for some reason which I have not yet figured out. In case of singular design matrix it will work, it identifies the problem and handles it internally like most regression functions do. However, problems can still occur.

Value

A list including:

pvalue	A numeric value that represents the logarithm of the generated p-value due to the conditional logistic regression (see reference below).
stat	A numeric value that represents the generated statistic due to the conditional logistic regression (see reference below).
stat_hash	The current hash object used for the statistics. See argument stat_hash and details. If argument hash = FALSE this is NULL.
pvalue_hash	The current hash object used for the p-values. See argument stat_hash and details. If argument hash = FALSE this is NULL.

Author(s)

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References

Michell H. Gail, Jay H. Lubin and Lawrence V. Rubinstein (1980). Likelihood calculations for matched case-control studies and survival studies with tied death times. *Biometrika* 68:703-707.

See Also

[SES](#), [testIndLogistic](#), [censIndCR](#), [censIndWR](#)

Examples

```
#simulate a dataset with continuous data
dataset <- matrix(rnorm(300 * 40), nrow = 300 )
#the target feature is the last column of the dataset as a vector
case <- rbinom(300, 1, 0.6)
ina <- which(case == 1)
ina <- sample(ina, 100)
case[-ina] = 0
id <- rep(1:100, 3)
target <- cbind(case, id)

results <- testIndClogit(target, dataset, xIndex = 34, csIndex = 30)
```

results

```
#run the SES algorithm using the testIndClogit conditional independence test
a1 <- SES(target, dataset, max_k = 3, threshold = 0.05, test = "testIndClogit");
a2 <- MMPC(target, dataset, max_k = 3, threshold = 0.05, test = "testIndClogit");
# print summary of the SES output
summary(a1);
```

Conditional independence test for longitudinal and clustered data using GEE
*Linear mixed models conditional independence test for longitudinal
class variables*

Description

The main task of this test is to provide a p-value PVALUE for the null hypothesis: feature 'X' is independent from 'TARGET' given a conditioning set CS. The pvalue is calculated by comparing a linear model based on the conditioning set CS against a model with both X and CS. The comparison is performed through an F test the appropriate degrees of freedom on the difference between the deviances of the two models. This test accepts a longitudinal target and longitudinal, categorical, continuous or mixed data as predictor variables.

Usage

```
testIndGEEReg(target, reps = NULL, group, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL,
correl = "exchangeable", se = "jack")
```

```
testIndGEELogistic(target, reps = NULL, group, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL,
correl = "exchangeable", se = "jack")
```

```
testIndGEEPois(target, reps = NULL, group, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL,
correl = "exchangeable", se = "jack")
```

```
testIndGEEGamma(target, reps = NULL, group, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL,
correl = "exchangeable", se = "jack")
```

```
testIndGEENormLog(target, reps = NULL, group, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL,
correl = "exchangeable", se = "jack")
```

```
testIndGEEOrdinal(target, reps = NULL, group, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL,
correl = "exchangeable", se = "jack")
```

Arguments

target	A numeric vector containing the values of the target variable. If the values are proportions or percentages, i.e. strictly within 0 and 1 they are mapped into R using $\log(\text{target}/(1 - \text{target}))$. In both cases a linear mixed model is applied. It can also be a binary variable (binary logistic regression) or a discrete, counts (Poisson regression), thus fitting generalised linear mixed models.
reps	A numeric vector containing the time points of the subjects. It's length is equal to the length of the target variable. If you have clustered data, leave this NULL.
group	A numeric vector containing the subjects or groups. It must be of the same length as target.
dataset	A numeric matrix or data frame, in case of categorical predictors (factors), containing the variables for performing the test. Rows as samples and columns as features.
xIndex	The index of the variable whose association with the target we want to test.
csIndex	The indices of the variables to condition on. If you have no variables set this equal to 0.
wei	A vector of weights to be used for weighted regression. The default value is NULL. It is mentioned in the "geepack" that weights is not (yet) the weight as in sas proc genmod, and hence is not recommended to use.
univariateModels	Fast alternative to the hash object for univariate test. List with vectors "pvalues" (p-values), "stats" (statistics) and "flags" (flag = TRUE if the test was succesful) representing the univariate association of each variable with the target. Default value is NULL.
hash	A boolean variable which indicates whether (TRUE) or not (FALSE) to use the hash-based implementation of the statistics of SES. Default value is FALSE. If TRUE you have to specify the stat_hash argument and the pvalue_hash argument.
stat_hash	A hash object (hash package required) which contains the cached generated statistics of a SES run in the current dataset, using the current test.
pvalue_hash	A hash object (hash package required) which contains the cached generated p-values of a SES run in the current dataset, using the current test.
correl	The correlation structure. For the Gaussian, Logistic, Poisson and Gamma regression this can be either "exchangeable" (compound symmetry, suitable for clustered data) or "ar1" (AR(1) model, suitable for longitudinal data). For the ordinal logistic regression its only the "exchangeable" correlation sturcture. This is obsolete for "testIndGEEOrdinal", but is here for compatibility reasons.
se	The method for estimating standard errors. This is very important and crucial. The available options for Gaussian, Logistic, Poisson and Gamma regression are: a) 'san.se', the usual robust estimate. b) 'jack': if approximate jackknife variance estimate should be computed. c) 'j1s': if 1-step jackknife variance estimate should be computed and d) 'fij': logical indicating if fully iterated jackknife variance estimate should be computed. If you have many clusters (sets of

repeated measurements) "san.se" is fine as it is asymptotically correct, plus jackknife estimates will take longer. If you have a few clusters, then maybe it's better to use jackknife estimates.

The jackknife variance estimator was suggested by Paik (1988), which is quite suitable for cases when the number of subjects is small ($K < 30$), as in many biological studies. The simulation studies conducted by Ziegler et al. (2000) and Yan and Fine (2004) showed that the approximate jackknife estimates are in many cases in good agreement with the fully iterated ones. This is obsolete for "testIndGEEOrdinal", but is here for compatibility reasons.

Details

If `hash = TRUE`, `testIndGEE` requires the arguments `'stat_hash'` and `'pvalue_hash'` for the hash-based implementation of the statistic test. These hash Objects are produced or updated by each run of `SES` (if `hash == TRUE`) and they can be reused in order to speed up next runs of the current statistic test. If `"SESoutput"` is the output of a `SES.temp` run, then these objects can be retrieved by `SESoutput@hashObject$stat_hash` and the `SESoutput@hashObject$pvalue_hash`.

Important: Use these arguments only with the same dataset that was used at initialization. For all the available conditional independence tests that are currently included on the package, please see `"?CondIndTests"`.

This test is for longitudinal and clustered data. Bear in mind that the time effect, for the longitudinal data case, is linear. It could be of higher order as well, but this would be a hyper-parameter, increasing the complexity of the models to be tested.

Value

A list including:

<code>pvalue</code>	A numeric value that represents the logarithm of the generated p-value due to the (generalised) linear mixed model (see reference below).
<code>stat</code>	A numeric value that represents the generated statistic due to the (generalised) linear mixed model (see reference below).
<code>stat_hash</code>	The current hash object used for the statistics. See argument <code>stat_hash</code> and details. If argument <code>hash = FALSE</code> this is <code>NULL</code> .
<code>pvalue_hash</code>	The current hash object used for the p-values. See argument <code>stat_hash</code> and details. If argument <code>hash = FALSE</code> this is <code>NULL</code> .

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

- Liang K.Y. and Zeger S.L. (1986). Longitudinal data analysis using generalized linear models. *Biometrika*, 73(1): 13-22.
- Prentice R.L. and Zhao L.P. (1991). Estimating equations for parameters in means and covariances of multivariate discrete and continuous responses. *Biometrics*, 47(3): 825-839.

Heagerty P.J. and Zeger S.L. (1996) Marginal regression models for clustered ordinal measurements. *Journal of the American Statistical Association*, 91(435): 1024-1036.

Paik M.C. (1988). Repeated measurement analysis for nonnormal data in small samples. *Communications in Statistics-Simulation and Computation*, 17(4): 1155-1171.

Ziegler A., Kastner C., Brunner D. and Blettner M. (2000). Familial associations of lipid profiles: A generalised estimating equations approach. *Statistics in medicine*, 19(24): 3345-3357

Yan J. and Fine J. (2004). Estimating equations for association structures. *Statistics in medicine*, 23(6): 859-874

Eugene Demidenko (2013). *Mixed Models: Theory and Applications with R*, 2nd Edition. New Jersey: Wiley & Sons.

Tsagris, M., Lagani, V., & Tsamardinos, I. (2018). Feature selection for high-dimensional temporal data. *BMC bioinformatics*, 19(1), 17.

See Also

[SES.temporal](#), [MMPC.temporal](#), [CondIndTests](#)

Examples

```
y <- rnorm(150)
x <- matrix(rnorm(150 * 5), ncol = 5)
id <- sample(1:20, 150, replace = TRUE)
testIndGEEReg(y, group=id, dataset=x, xIndex=1, csIndex=3)
```

Conditional independence test for longitudinal and clustered data using GLMM
*Linear mixed models conditional independence test for longitudinal
class variables*

Description

The main task of this test is to provide a p-value PVALUE for the null hypothesis: feature 'X' is independent from 'TARGET' given a conditioning set CS. The pvalue is calculated by comparing a linear model based on the conditioning set CS against a model with both X and CS. The comparison is performed through an F test the appropriate degrees of freedom on the difference between the deviances of the two models. This test accepts a longitudinal target and longitudinal, categorical, continuous or mixed data as predictor variables.

Usage

```
testIndGLMMReg(target, reps = NULL, group, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL,
pvalue_hash=NULL, slopes = FALSE)
```

```
testIndGLMMLogistic(target, reps = NULL, group, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL,
```

```
pvalue_hash=NULL, slopes = FALSE)
```

```
testIndGLMMPois(target, reps = NULL, group, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL,
pvalue_hash=NULL, slopes = FALSE)
```

```
testIndLMM(target, reps = NULL, group, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL,
pvalue_hash=NULL, slopes = FALSE)
```

Arguments

target	A numeric vector containing the values of the target variable. If the values are proportions or percentages, i.e. strictly within 0 and 1 they are mapped into R using $\log(\text{target}/(1 - \text{target}))$. In both cases a linear mixed model is applied. It can also be a binary variable (binary logistic regression) or a discrete, counts (Poisson regression), thus fitting generalised linear mixed models.
reps	A numeric vector containing the time points of the subjects. It's length is equal to the length of the target variable. If you have clustered data, leave this NULL.
group	A numeric vector containing the subjects or groups. It must be of the same length as target.
dataset	A numeric matrix or data frame, in case of categorical predictors (factors), containing the variables for performing the test. Rows as samples and columns as features.
xIndex	The index of the variable whose association with the target we want to test.
csIndex	The indices of the variables to condition on. If you have no variables set this equal to 0.
wei	A vector of weights to be used for weighted regression. The default value is NULL.
univariateModels	Fast alternative to the hash object for univariate test. List with vectors "pvalues" (p-values), "stats" (statistics) and "flags" (flag = TRUE if the test was succesful) representing the univariate association of each variable with the target. Default value is NULL.
hash	A boolean variable which indicates whether (TRUE) or not (FALSE) to use the hash-based implementation of the statistics of SES. Default value is FALSE. If TRUE you have to specify the stat_hash argument and the pvalue_hash argument.
stat_hash	A hash object (hash package required) which contains the cached generated statistics of a SES run in the current dataset, using the current test.
pvalue_hash	A hash object (hash package required) which contains the cached generated p-values of a SES run in the current dataset, using the current test.
slopes	A boolean variable which indicates whether (TRUE) to or not (FALSE) random slopes in the time effect as well. By default random intercepts are considered.

Details

If `hash = TRUE`, `testIndGLMM` requires the arguments `'stat_hash'` and `'pvalue_hash'` for the hash-based implementation of the statistic test. These hash Objects are produced or updated by each run of `SES` (if `hash == TRUE`) and they can be reused in order to speed up next runs of the current statistic test. If `"SESoutput"` is the output of a `SES.temp` run, then these objects can be retrieved by `SESoutput@hashObject$stat_hash` and the `SESoutput@hashObject$pvalue_hash`.

Important: Use these arguments only with the same dataset that was used at initialization. For all the available conditional independence tests that are currently included on the package, please see `"?CondIndTests"`.

This test is for longitudinal and clustered data. Bear in mind that the time effect, for the longitudinal data case, is linear. It could be of higher order as well, but this would be a hyper-parameter, increasing the complexity of the models to be tested.

The `testIndLMM` is used for linear mixed models with no weights, no slopes and no reps. This is a random intercepts model only. The advantage of this function is that it is tens of times faster than `testIndGLMM`.

Value

A list including:

<code>pvalue</code>	A numeric value that represents the logarithm of the generated p-value due to the (generalised) linear mixed model (see reference below).
<code>stat</code>	A numeric value that represents the generated statistic due to the (generalised) linear mixed model (see reference below).
<code>stat_hash</code>	The current hash object used for the statistics. See argument <code>stat_hash</code> and details. If argument <code>hash = FALSE</code> this is <code>NULL</code> .
<code>pvalue_hash</code>	The current hash object used for the p-values. See argument <code>stat_hash</code> and details. If argument <code>hash = FALSE</code> this is <code>NULL</code> .

Author(s)

Vincenzo Lagani, Ioannis Tsamardinos, Michail Tsagris and Giorgos Athineou

R implementation and documentation: Giorgos Athineou <athineou@csd.gr>, Vincenzo Lagani <vlagani@csd.uoc.gr> and Michail Tsagris <mtsagris@csd.uoc.gr>

References

Eugene Demidenko (2013). *Mixed Models: Theory and Applications with R*, 2nd Edition. New Jersey: Wiley & Sons.

Jose Pinheiro Jose and Douglas Bates. *Mixed-effects models in S and S-PLUS*. Springer Science & Business Media, 2006.

See Also

[SES.temporal](#), [MMPC.temporal](#), [CondIndTests](#)

Examples

```

y <- rnorm(150)
x <- matrix(rnorm(150 * 5), ncol = 5)
id <- sample(1:20, 150, replace = TRUE)
testIndGLMMReg(y, group=id, dataset=x, xIndex=1, csIndex=3)
testIndLMM(y, group=id, dataset=x, xIndex=1, csIndex=3)

```

Conditional independence test for proportions/percentages

Beta regression conditional independence test for proportions/percentage class dependent variables and mixed predictors

Description

The main task of this test is to provide a p-value PVALUE for the null hypothesis: feature 'X' is independent from 'TARGET' given a conditioning set CS. The pvalue is calculated by comparing a Beta regression model based on the conditioning set CS against a model whose regressor are both X and CS. The comparison is performed through a chi-square test with the appropriate degrees of freedom on the difference between the deviances of the two models.

Usage

```

testIndBeta(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL)

```

```

permBeta(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL,
threshold = 0.05, R = 999)

```

```

waldBeta(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL)

```

Arguments

target	A numeric vector containing the values of the target variable. They must be percentages or proportions, i.e. within the (0, 1) interval. Currently 0 and/or 1 values are not allowed.
dataset	A numeric matrix or data frame, in case of categorical predictors (factors), containing the variables for performing the test. Rows as samples and columns as features. In the case of "waldBeta" this is strictly a matrix.
xIndex	The index of the variable whose association with the target we want to test.
csIndex	The indices of the variables to condition on. If you have no variables set this equal to 0.
wei	A vector of weights to be used for weighted regression. The default value is NULL.

univariateModels	Fast alternative to the hash object for univariate test. List with vectors "pvalues" (p-values), "stats" (statistics) and "flags" (flag = TRUE if the test was succesful) representing the univariate association of each variable with the target. Default value is NULL.
hash	A boolean variable which indicates whether (TRUE) or not (FALSE) to use the hash-based implementation of the statistics of SES. Default value is FALSE. If TRUE you have to specify the stat_hash argument and the pvalue_hash argument.
stat_hash	A hash object (hash package required) which contains the cached generated statistics of a SES run in the current dataset, using the current test.
pvalue_hash	A hash object (hash package required) which contains the cached generated p-values of a SES run in the current dataset, using the current test.
threshold	Threshold (suitable values in [0,1]) for assessing p-values significance.
R	The number of permutations, set to 999 by default. There is a trick to avoid doing all permutations. As soon as the number of times the permuted test statistic is more than the observed test statistic is more than 50 (if threshold = 0.05 and R = 999), the p-value has exceeded the significance level (threshold value) and hence the predictor variable is not significant. There is no need to continue do the extra permutations, as a decision has already been made.

Details

If hash = TRUE, testIndBeta requires the arguments 'stat_hash' and 'pvalue_hash' for the hash-based implementation of the statistic test. These hash Objects are produced or updated by each run of SES (if hash == TRUE) and they can be reused in order to speed up next runs of the current statistic test. If "SESoutput" is the output of a SES run, then these objects can be retrieved by SESoutput@hashObject\$stat_hash and the SESoutput@hashObject\$pvalue_hash.

Important: Use these arguments only with the same dataset that was used at initialization. For all the available conditional independence tests that are currently included on the package, please see "?CondIndTests". An alternative regression to this is "testIndReg" and "testIndRQ". In these two latter cases, the logit transformation is first applied to the target variable.

The log-likelihood ratio test used in "testIndBeta" requires the fitting of two models. The Wald test used in "waldBeta" requires fitting of only one model, the full one. The significance of the variable is examined only. Only continuous (or binary) predictor variables are currently accepted in this test.

Value

A list including:

pvalue	A numeric value that represents the logarithm of the generated p-value due to Beta regression (see reference below).
stat	A numeric value that represents the generated statistic due to Beta regression (see reference below).
stat_hash	The current hash object used for the statistics. See argument stat_hash and details. If argument hash = FALSE this is NULL.
pvalue_hash	The current hash object used for the logged p-values. See argument stat_hash and details. If argument hash = FALSE this is NULL.

Author(s)

Vincenzo Lagani, Ioannis Tsamardinos, Michail Tsagris and Giorgos Athineou

R implementation and documentation: Giorgos Athineou <athineou@csd.uoc.gr>, Vincenzo Lagani <vlagani@csd.uoc.gr> and Michail Tsagris <mtsagris@csd.uoc.gr>

References

Ferrari S.L.P. and Cribari-Neto F. (2004). Beta Regression for Modelling Rates and Proportions. Journal of Applied Statistics, 31(7): 799-815.

See Also

[SES](#), [testIndReg](#), [testIndRQ](#), [testIndFisher](#), [CondIndTests](#)

Examples

```
#simulate a dataset with continuous data
dataset <- matrix(runif(100 * 20, 1, 1000), ncol = 20 )
#the target feature is the last column of the dataset as a vector
target <- dataset[, 20]
dataset <- dataset[, -20]
target <- target / (max(target) + 2 )

results <- testIndBeta(target, dataset, xIndex = 14, csIndex = 9)
results

#run the SES algorithm using the testIndBeta conditional independence test
sesObject <- SES(target, dataset, max_k = 3, threshold = 0.05, test = "testIndBeta");
#print summary of the SES output
summary(sesObject);
```

Conditional independence test for the static-longitudinal scenario
Conditional independence test for the static-longitudinal scenario

Description

The main task of this test is to provide a p-value PVALUE for the null hypothesis: feature 'X' is independent from 'TARGET' given a conditioning set CS. The pvalue is calculated by comparing a logistic model based on the conditioning set CS against a model whose regressor are both X and CS. The comparison is performed through a chi-square test with the appropriate degrees of freedom on the difference between the deviances of the two models.

Usage

```
testIndTimeLogistic(target, dataset, xIndex, csIndex, wei = NULL,
  univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL)
```

```
testIndTimeMultinom(target, dataset, xIndex, csIndex, wei = NULL,
  univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL)
```

Arguments

target	A numeric vector containing the values of the target variable. For the "testInd-Logistic" this can either be a binary numerical variable or a factor variable. The factor variable can have two values (binary logistic regression), more than two values (multinomial logistic regression) or it can be an ordered factor with more than two values (ordinal regression). The last one is for example, factor(x, ordered = TRUE). The "waldBinary" is the Wald test version of the binary logistic regression. The "waldOrdinal" is the Wald test version of the ordinal regression.
dataset	A numeric matrix with the constants and slopes stack one upo the other. The first r rows are the constants and the rest of the rows contains the slopes. In some the matrix can be calculated using the group.mvbetas function.
xIndex	The index of the variable whose association with the target we want to test.
csIndex	The indices of the variables to condition on. If you have no variables set this equal to 0.
wei	A vector of weights to be used for weighted regression. The default value is NULL.
univariateModels	Fast alternative to the hash object for univariate test. List with vectors "pvalues" (p-values), "stats" (statistics) and "flags" (flag = TRUE if the test was succesful) representing the univariate association of each variable with the target. Default value is NULL.
hash	A boolean variable which indicates whether (TRUE) or not (FALSE) to use the hash-based implementation of the statistics of SES. Default value is FALSE. If TRUE you have to specify the stat_hash argument and the pvalue_hash argument.
stat_hash	A hash object (hash package required) which contains the cached generated statistics of a SES run in the current dataset, using the current test.
pvalue_hash	A hash object (hash package required) which contains the cached generated p-values of a SES run in the current dataset, using the current test.

Details

This conditional independence test is devised for the static-longitudinal scenario of Tsagris, Lagani and Tsamardinos (2018). The idea is that you have many features of longitudinal data for many subjects. For each subject you have calculated the coefficients of a simple linear regression over time and this is repeated for each feature. In the end, assuming p features, you have p constants and p slopes for each subject, each constant and slope refers to a feature for a subject.

Value

A list including:

pvalue	A numeric value that represents the logarithm of the generated p-value.
stat	A numeric value that represents the generated statistic.
stat_hash	The current hash object used for the statistics. See argument stat_hash and details. If argument hash = FALSE this is NULL.
pvalue_hash	The current hash object used for the p-values. See argument stat_hash and details. If argument hash = FALSE this is NULL.

Note

This test uses the function multinom (package nnet) for multinomial logistic regression, the function clm (package ordinal) for ordinal logit regression and the function glm (package stats) for binomial regression.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

Tsagris M., Lagani V., & Tsamardinos I. (2018). Feature selection for high-dimensional temporal data. *BMC bioinformatics*, 19(1), 17.

Vincenzo Lagani, George Kortas and Ioannis Tsamardinos (2013), Biomarker signature identification in "omics" with multiclass outcome. *Computational and Structural Biotechnology Journal*, 6(7):1-7.

McCullagh, Peter, and John A. Nelder. *Generalized linear models*. CRC press, USA, 2nd edition, 1989.

See Also

[SES](#), [gSquare](#), [CondIndTests](#)

Examples

```
## assume these are longitudinal data, each column is a variable (or feature)
x <- matrix( rnorm(400 * 100), ncol = 100 )
id <- rep(1:80, each = 5) ## 80 subjects
reps <- rep( seq(4, 12, by = 2), 80) ## 5 time points for each subject
dataset <- group.mvbetas(x, id, reps)
## these are the regression coefficients of the first subject's values on the
## reps (which is assumed to be time in this example)
target <- rbinom(80, 1, 0.5)
testIndTimeLogistic(target, dataset, xIndex = 1 ,csIndex = 0)
testIndTimeLogistic(target, dataset, xIndex = 1 ,csIndex = 2)
```

Conditional independence tests counting the number of times a possible collider d-separates two nodes
Many conditional independence tests counting the number of times a possible collider d-separates two nodes

Description

Many conditional independence tests counting the number of times a possible collider d-separates two nodes .

Usage

```
condis(ind1, ind2, cs1, cs2, Var, dat, type = "pearson", rob = FALSE, max_k = 2, R = 1 )
```

Arguments

ind1	The index of the one variable to be considered.
ind2	The index of the other variable to be considered.
cs1	The index or indices of the conditioning set of variable(s). These are the neighbours of node ind1.
cs2	The index or indices of the conditioning set of variable(s). These are the neighbours of node ind2.
Var	The index of the possible collider.
dat	A numerical matrix or a data.frame with numerical, binary, nominal and ordinal variables only.
type	This is either "pearson", "spearman", "cat", "distcor", "ci.mm" or "ci.fast".
rob	In case you want robust estimation of the Pearson correlation prior to applying the conditional independence test. This is activated only when type = "pearson".
max_k	The maximum number of conditioning variables to consider. It can be the case that each node ind1 and ind2 has 10 neighbours. We should try all possible combinations of the neighbours of ind1 and then of ind2. To reduce the computational cost we search the subsets with at most max_k variables.
R	This is used by most tests, except for type = "ci.mm" and type = "ci.fast".

Details

This is to be used in the conservative version of Rule 0 of the Pc algorithm. When one wants to know whether a variable is a possible collider, Ramsey, Spirtes and Zhang (2005) propose to perform the following action. For every unshielded triple (X, Y, Z) check all subsets of X's possible parents and of Z's possible parents. a) If Y is NOT in any such set conditional on which, X and Z are independent orient X - Y - Z as X -> Y <- Z. b) If Y is in ALL sets conditional on which, X and Z are independent, leave X - Y - Z as it is, i.e. a non-collider. c) Mark the triple X - Y - Z as "unfaithfull" otherwise. This modification leads to the so called conservative PC (CPC) algorithm.

54 Conditional independence tests counting the number of times a possible collider d-separates two nodes

A few years later, Colombo and Maathuis (2014) suggested a modification of the previous action, called the majority rule. If Y is less than 50% of the sets that render X and Z independent orient $X - Y - Z$ as $X \rightarrow Y \leftarrow Z$. If Y is found in exactly 50% of sets that render X and Z independent, this triple is marked as "ambiguous". This modification leads to the so-called majority rule PC (MPC) algorithm.

This function we have implemented here, does exactly this. It applies tests to many subsets and returns a matrix with two columns. The first one contains 0 or 1 and the second is the p-value. A value of 0 indicates absence of the possible collider from the set that produced that p-value, whereas a value of 1 indicates its presence in the set.

This way, we can measure the proportion of times the possible collider Y was in a subset that rendered X and Z independent.

Value

A matrix with two columns. The second one is the logarithm of the p-value. The first one contains 0s and 1s. The value of 0 means that the candidate collider was not in that set which produced the relevant p-value, whereas a value of 1 indicates that it was a member of that conditioning set.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

Ramsey, J., Zhang, J., Spirtes, P., 2006. Adjacency-faithfulness and conservative causal inference. Proceedings of the 22nd Annual Conference on Uncertainty in Artificial Intelligence (UAI 2006). <https://arxiv.org/ftp/arxiv/papers/1206/1206.6843.pdf>

Colombo, Diego, and Marloes H. Maathuis (2014). Order-independent constraint-based causal structure learning. *The Journal of Machine Learning Research* 15(1): 3741–3782.

See Also

[SES](#), [MMPC](#), [testIndLogistic](#)

Examples

```
x <- rdag2(1000, p = 10, nei = 5)
G <- x$G
dat <- x$x
cs1 <- which(G[6, ] > 0 | G[, 6] > 0)
cs2 <- which(G[7, ] > 0 | G[, 7] > 0)
cs1 <- setdiff( cs1, c(7, 3) )
cs2 <- setdiff( cs2, c(6, 3) )
condis(6, 7, cs1, cs2, 3, dat, type = "pearson", rob = FALSE, max_k = 3, R = 1 )
```

Conditional independence tests for continuous univariate and multivariate data
*Linear (and non-linear) regression conditional independence test for
continuous univariate and multivariate response variables*

Description

The main task of this test is to provide a p-value PVALUE for the null hypothesis: feature 'X' is independent from 'TARGET' given a conditioning set CS. The pvalue is calculated by comparing a linear regression model based on the conditioning set CS against a model whose regressor are both X and CS. The comparison is performed through an F test the appropriate degrees of freedom on the difference between the deviances of the two models.

Usage

```
testIndReg(target, dataset, xIndex, csIndex, wei = NULL,  
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL)
```

```
testIndRQ(target, dataset, xIndex, csIndex, wei = NULL,  
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL)
```

```
testIndMVreg(target, dataset, xIndex, csIndex, wei = NULL,  
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL)
```

```
testIndMMReg(target, dataset, xIndex, csIndex, wei = NULL,  
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL)
```

```
waldMMReg(target, dataset, xIndex, csIndex, wei = NULL,  
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL)
```

```
permReg(target, dataset, xIndex, csIndex, wei = NULL,  
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL,  
threshold = 0.05, R = 999)
```

```
permMMReg(target, dataset, xIndex, csIndex, wei = NULL,  
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL,  
threshold = 0.05, R = 999)
```

```
permRQ(target, dataset, xIndex, csIndex, wei = NULL,  
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL,  
threshold = 0.05, R = 999)
```

```
permMVreg(target, dataset, xIndex, csIndex, wei = NULL,  
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL,  
threshold = 0.05, R = 999)
```

Arguments

target	A numeric vector containing the values of the target variable. If the values are proportions or percentages, i.e. strictly within 0 and 1 they are mapped into R using $\log(\text{target}/(1 - \text{target}))$. In the case of <code>testIndMVreg</code> , the same takes place true. See details for more information.
dataset	A numeric matrix or data frame, in case of categorical predictors (factors), containing the variables for performing the test. Rows as samples and columns as features. In the cases of "waldIGreg" and "waldMMreg" this is strictly a matrix.
xIndex	The index of the variable whose association with the target we want to test.
csIndex	The indices of the variables to condition on. If you have no variables set this equal to 0.
wei	A vector of weights to be used for weighted regression. The default value is NULL. They are not take into account in the robust regression via MM estimation (<code>testIndMMReg</code> and <code>permMMreg</code>).
univariateModels	Fast alternative to the hash object for univariate test. List with vectors "pvalues" (p-values) and "stats" (statistics) representing the univariate association of each variable with the target. Default value is NULL.
hash	A boolean variable which indicates whether (TRUE) or not (FALSE) to use the hash-based implementation of the statistics of SES. Default value is FALSE. If TRUE you have to specify the <code>stat_hash</code> argument and the <code>pvalue_hash</code> argument.
stat_hash	A hash object (hash package required) which contains the cached generated statistics of a SES run in the current dataset, using the current test.
pvalue_hash	A hash object (hash package required) which contains the cached generated p-values of a SES run in the current dataset, using the current test.
threshold	Threshold (suitable values in [0,1]) for assessing p-values significance.
R	The number of permutations, set to 999 by default. There is a trick to avoid doing all permutations. As soon as the number of times the permuted test statistic is more than the observed test statistic is more than 50 (if <code>threshold = 0.05</code> and <code>R = 999</code>), the p-value has exceeded the significance level (threshold value) and hence the predictor variable is not significant. There is no need to continue do the extra permutations, as a decision has already been made.

Details

If `hash = TRUE`, all three tests require the arguments 'stat_hash' and 'pvalue_hash' for the hash-based implementation of the statistic test. These hash Objects are produced or updated by each run of SES (if `hash == TRUE`) and they can be reused in order to speed up next runs of the current statistic test. If "SESoutput" is the output of a SES run, then these objects can be retrieved by `SESoutput@hashObject$stat_hash` and the `SESoutput@hashObject$pvalue_hash`.

Important: Use these arguments only with the same dataset that was used at initialization.

`TestIndReg` offers linear regression.

`testIndMMReg` offers robust linear MM estimation regression.

TestIndRQ offers quantile (median) regression as a robust alternative to linear regression.

In both cases, if the dependent variable consists of proportions (values between 0 and 1) the logit transformation is applied and the tests are applied then.

testIndMVreg is for multivariate continuous response variables. Compositional data are positive multivariate data and each vector (observation) sums to the same constant, usually taken 1 for convenience. A check is performed and if such data are found, the additive log-ratio (multivariate logit) transformation (Aitchison, 1986) is applied beforehand. Zeros are not allowed.

For all the available conditional independence tests that are currently included on the package, please see "?CondIndTests".

The Wald test used in "waldMMReg" requires fitting of only one model, the full one. The significance of the variable is examined only. Only continuous (or binary) predictor variables are currently accepted in this test.

Value

A list including:

pvalue	A numeric value that represents the logarithm of the generated p-value due to linear regression (see reference below).
stat	A numeric value that represents the generated statistic due to linear regression (see reference below).
stat_hash	The current hash object used for the statistics. See argument stat_hash and details. If argument hash = FALSE this is NULL.
pvalue_hash	The current hash object used for the p-values. See argument stat_hash and details. If argument hash = FALSE this is NULL.

Author(s)

Vincenzo Lagani, Ioannis Tsamardinos, Michail Tsagris and Giorgos Athineou

R implementation and documentation: Giorgos Athineou <athineou@csd.gr>, Vincenzo Lagani <vlagani@csd.uoc.gr> and Michail Tsagris <mtsagris@csd.uoc.gr>

References

- Draper, N.R. and Smith H. (1988). Applied regression analysis. New York, Wiley, 3rd edition.
- Hampel F. R., Ronchetti E. M., Rousseeuw P. J., and Stahel W. A. (1986). Robust statistics: the approach based on influence functions. John Wiley & Sons.
- Koenker R.W. (2005). Quantile regression. New York, Cambridge University Press.
- Mardia, Kanti, John T. Kent and John M. Bibby. Multivariate analysis. Academic press, 1979.
- John Aitchison. The Statistical Analysis of Compositional Data, Chapman & Hall; reprinted in 2003, with additional material, by The Blackburn Press.

See Also

[testIndRQ](#), [testIndFisher](#), [testIndSpearman](#), [CondIndTests](#)

Examples

```

#simulate a dataset with continuous data
dataset <- matrix(runif(100 * 100, 1, 100), ncol = 100 )
#the target feature is the last column of the dataset as a vector
target <- dataset[, 100]
dataset <- dataset[, -100]

testIndReg(target, dataset, xIndex = 44, csIndex = 50)
testIndMMReg(target, dataset, xIndex = 44, csIndex = 50)
testIndRQ(target, dataset, xIndex = 44, csIndex = 50)
testIndIGreg(target, dataset, xIndex = 44, csIndex = 50)

#define class variable (here tha last column of the dataset)
#run the SES algorithm using the testIndReg conditional independence test
sesObject <- SES(target, dataset, max_k = 3, threshold = 0.05, test = "testIndReg");
sesObject2 <- SES(target, dataset, max_k = 3, threshold = 0.05, test = "testIndRQ");
#print summary of the SES output

summary(sesObject);
summary(sesObject2);

```

Conditional independence tests for count data

Regression conditional independence test for discrete (counts) class dependent variables

Description

The main task of this test is to provide a p-value PVALUE for the null hypothesis: feature 'X' is independent from 'TARGET' given a conditioning set CS. The pvalue is calculated by comparing a Poisson regression model based on the conditioning set CS against a model whose regressor are both X and CS. The comparison is performed through a chi-square test with the appropriate degrees of freedom on the difference between the deviances of the two models. The models supported here are poisson, zero inflated poisson and negative binomial.

Usage

```

testIndPois(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL)

testIndNB(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL)

testIndZIP(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL)

testIndQPois(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL)

```

```
permPois(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL,
threshold = 0.05, R = 999)
```

```
permNB(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL,
threshold = 0.05, R = 999)
```

```
permZIP(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL,
threshold = 0.05, R = 999)
```

```
waldPois(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL)
```

```
waldNB(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL)
```

```
waldZIP(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL)
```

Arguments

target	A numeric vector containing the values of the target variable.
dataset	A numeric matrix or data frame, in case of categorical predictors (factors), containing the variables for performing the test. Rows as samples and columns as features. In the cases of "waldPois", "waldNB" and "waldZIP" this is strictly a matrix.
xIndex	The index of the variable whose association with the target we want to test.
csIndex	The indices of the variables to condition on. If you have no variables set this equal to 0.
wei	A vector of weights to be used for weighted regression. The default value is NULL.
univariateModels	Fast alternative to the hash object for univariate test. List with vectors "pvalues" (p-values), "stats" (statistics) and "flags" (flag = TRUE if the test was succesful) representing the univariate association of each variable with the target. Default value is NULL.
hash	A boolean variable which indicates whether (TRUE) or not (FALSE) to use the hash-based implementation of the statistics of SES. Default value is FALSE. If TRUE you have to specify the stat_hash argument and the pvalue_hash argument.
stat_hash	A hash object (hash package required) which contains the cached generated statistics of a SES run in the current dataset, using the current test.
pvalue_hash	A hash object (hash package required) which contains the cached generated p-values of a SES run in the current dataset, using the current test.

threshold	Threshold (suitable values in [0,1]) for assessing p-values significance.
R	The number of permutations, set to 999 by default. There is a trick to avoid doing all permutations. As soon as the number of times the permuted test statistic is more than the observed test statistic is more than 50 (if threshold = 0.05 and R = 999), the p-value has exceeded the significance level (threshold value) and hence the predictor variable is not significant. There is no need to continue do the extra permutations, as a decision has already been made.

Details

If hash = TRUE, all three tests require the arguments 'stat_hash' and 'pvalue_hash' for the hash-based implementation of the statistic test. These hash Objects are produced or updated by each run of SES (if hash == TRUE) and they can be reused in order to speed up next runs of the current statistic test. If "SESoutput" is the output of a SES run, then these objects can be retrieved by SESoutput@hashObject\$stat_hash and the SESoutput@hashObject\$pvalue_hash.

Important: Use these arguments only with the same dataset that was used at initialization. For all the available conditional independence tests that are currently included on the package, please see "?CondIndTests".

If you have overdispersion, the variance is higher than the mean, a negative binomial is to be used. If you have more zeros than expected under a Poisson model, not overdispersion, then zero inflated Poisson is to be used. Bear in mind that if you have a small number of zeros, there is no reason to use this model. If for example you have count data but no, or 1 zeros, this will not work.

The log-likelihood ratio test used in "testIndPois", "testIndNB" and "testIndZIP" requires the fitting of two models. The Wald test used in "waldPois", "waldNB" and "waldZIP" requires fitting of only one model, the full one. The significance of the variable is examined only. Only continuous (or binary) predictor variables are currently accepted in this test.

The testIndQPois does quasi Poisson regression. The benefit of this regression is that it works for over and under dispersed data. Negative Binomial works for over dispersed data, but not for under dispersed data. In addition it is fast.

Value

A list including:

pvalue	A numeric value that represents the logarithm of the generated p-value due to the count data regression (see references below).
stat	A numeric value that represents the generated statistic due to Poisson regression(see reference below).
stat_hash	The current hash object used for the statistics. See argument stat_hash and details. If argument hash = FALSE this is NULL.
pvalue_hash	The current hash object used for the p-values. See argument stat_hash and details. If argument hash = FALSE this is NULL.

Author(s)

Vincenzo Lagani, Ioannis Tsamardinos, Michail Tsagris and Giorgos Athineou

R implementation and documentation: Giorgos Athineou <athineou@csd.uoc.gr>, Vincenzo Lagani <vlagani@csd.uoc.gr> and Michail Tsagris <mtsagris@csd.uoc.gr>

References

- McCullagh P. and Nelder J.A. (1989). Generalized linear models. CRC press, USA, 2nd edition.
- Lambert D. (1992). Zero-inflated Poisson regression, with an application to defects in manufacturing. *Technometrics*, 34(1):1-14.
- Joseph M.H. (2011). Negative Binomial Regression. Cambridge University Press, 2nd edition.

See Also

[testIndReg](#), [testIndNB](#), [testIndZIP](#), [gSquare](#), [CondIndTests](#)

Examples

```
#simulate a dataset with continuous data
dataset <- matrix(runif(400 * 50, 1, 50), ncol = 50 )
#the target feature is the last column of the dataset as a vector
target <- rpois(400, 10)
results <- testIndPois(target, dataset, xIndex = 24, csIndex = 10)
results

#run the SES algorithm using the testIndPois conditional independence test
sesObject <- SES(target, dataset, max_k = 3, threshold = 0.05, test = "testIndPois");
sesObject2 <- SES(target, dataset, max_k = 3, threshold = 0.05, test = "testIndNB");
summary(sesObject);
```

Conditional independence tests for left censored data

Conditional independence test for survival data

Description

The main task of this test is to provide a p-value PVALUE for the null hypothesis: feature 'X' is independent from 'TARGET' given a conditioning set CS. This test can be based on the Cox (semi-parametric) regression or on the Weibull (parametric) regression.

Usage

```
testIndTobit(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL)

waldTobit(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL)

permTobit(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL,
threshold = 0.05, R = 999)
```

Arguments

target	A Survival object (class Surv from package survival) with left censored data. This test works with responses that are left censored. See the examples below (the command <code>Surv</code> or the final example in the <code>survreg</code> documentation of the "survival" package) for more information on how to create the target variable.
dataset	A numeric matrix or data frame, in case of categorical predictors (factors), containing the variables for performing the test. Rows as samples and columns as features. In the cases of "waldTobit" and "permTobit" this is strictly a matrix.
xIndex	The index of the variable whose association with the target we want to test.
csIndex	The indices of the variables to condition on. If you have no variables set this equal to 0.
wei	A vector of weights to be used for weighted regression. The default value is NULL.
univariateModels	Fast alternative to the hash object for univariate test. List with vectors "pvalues" (p-values), "stats" (statistics) and "flags" (flag = TRUE if the test was succesful) representing the univariate association of each variable with the target. Default value is NULL.
hash	A boolean variable which indicates whether (TRUE) or not (FALSE) to use the hash-based implementation of the statistics of SES. Default value is FALSE. If TRUE you have to specify the stat_hash argument and the pvalue_hash argument.
stat_hash	A hash object (hash package required) which contains the cached generated statistics of a SES run in the current dataset, using the current test.
pvalue_hash	A hash object (hash package required) which contains the cached generated p-values of a SES run in the current dataset, using the current test.
threshold	Threshold (suitable values in [0,1]) for assessing p-values significance.
R	The number of permutations, set to 999 by default. There is a trick to avoid doing all permutations. As soon as the number of times the permuted test statistic is more than the observed test statistic is more than 50 (in this example case), the p-value has exceeded the signifiace level (threshold value) and hence the predictor variable is not significant. There is no need to continue do the extra permutations, as a decision has already been made.

Details

Tobit regression is performed. The implied model is Gaussian with left censored data.

If `hash = TRUE`, `censIndCR`, `censIndWR` and `censIndER` require the arguments 'stat_hash' and 'pvalue_hash' for the hash-based implementation of the statistic test. These hash Objects are produced or updated by each run of SES (if `hash == TRUE`) and they can be reused in order to speed up next runs of the current statistic test. If "SESoutput" is the output of a SES run, then these objects can be retrieved by `SESoutput@hashObject$stat_hash` and the `SESoutput@hashObject$pvalue_hash`.

Important: Use these arguments only with the same dataset that was used at initialization.

For all the available conditional independence tests that are currently included on the package, please see "?CondIndTests".

The log-likelihood ratio test used in "testIndTobit" requires the fitting of two models. The Wald test used in "waldTobit", requires fitting of only one model, the full one. The significance of the variable is examined only. Only continuous (or binary) predictor variables are currently accepted in this test.

Value

A list including:

pvalue	A numeric value that represents the logarithm of the generated p-value.
stat	A numeric value that represents the generated statistic.
stat_hash	The current hash object used for the statistics. See argument stat_hash and details. If argument hash = FALSE this is NULL.
pvalue_hash	The current hash object used for the p-values. See argument stat_hash and details. If argument hash = FALSE this is NULL.

Note

This test uses the functions "survreg" and Surv of the package survival and the function anova (analysis of variance) of the package stats.

Author(s)

R implementation and documentation: Michail Tsagris <mtsagris@yahoo.gr>

References

Tobin James (1958). Estimation of relationships for limited dependent variables. *Econometrica*. 26(1): 24-36.

See Also

[SES](#), [censIndWR](#), [testIndFisher](#), [gSquare](#), [testIndLogistic](#), [Surv](#), [anova](#), [CondIndTests](#)

Examples

```
require(survival, quietly = TRUE)

x <- matrix( rnorm(100 * 30), ncol = 30)
y <- x[, 1] - x[, 2] + rnorm(100, 5)
y[y < 0 ] <- 0
y <- Surv(y, y>0, type = 'left')

#run the censIndCR conditional independence test
testIndTobit(y, x, xIndex = 12, csIndex = c(5, 7, 4) )
waldTobit(y, x, xIndex = 12, csIndex = c(5, 7, 4) )
permTobit(y, x, xIndex = 12, csIndex = c(5, 7, 4), R = 499 )

#run the SES algorithm using the censIndCR conditional independence
#test for the survival class variable
a <- SES(y, x, max_k = 2, threshold = 0.05, test = "testIndTobit");
```

 Conditional independence tests for positive data

Regression conditional independence test for positive response variables.

Description

The main task of this test is to provide a p-value PVALUE for the null hypothesis: feature 'X' is independent from 'TARGET' given a conditioning set CS. The pvalue is calculated by comparing a Poisson regression model based on the conditioning set CS against a model whose regressor are both X and CS. The comparison is performed through a chi-square test with the appropriate degrees of freedom on the difference between the deviances of the two models. The models supported here are poisson, zero inflated poisson and negative binomial.

Usage

```
testIndGamma(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL)
```

```
testIndNormLog(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL)
```

```
testIndIGreg(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL)
```

```
permGamma(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL,
threshold = 0.05, R = 999)
```

```
permNormLog(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL,
threshold = 0.05, R = 999)
```

```
permIGreg(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL,
threshold = 0.05, R = 999)
```

```
waldGamma(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL)
```

```
waldNormLog(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL)
```

```
waldIGreg(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL)
```

Arguments

target	A numeric vector containing the values of the target variable. For the Gamma based tests, the values must be strictly greater than zero. For the NormLog case, zeros can be included.
dataset	A numeric matrix or data frame, in case of categorical predictors (factors), containing the variables for performing the test. Rows as samples and columns as features. In the cases of "waldPois", "waldNB" and "waldZIP" this is strictly a matrix.
xIndex	The index of the variable whose association with the target we want to test.
csIndex	The indices of the variables to condition on. If you have no variables set this equal to 0.
wei	A vector of weights to be used for weighted regression. The default value is NULL.
univariateModels	Fast alternative to the hash object for univariate test. List with vectors "pvalues" (p-values), "stats" (statistics) and "flags" (flag = TRUE if the test was succesful) representing the univariate association of each variable with the target. Default value is NULL.
hash	A boolean variable which indicates whether (TRUE) or not (FALSE) to use the hash-based implementation of the statistics of SES. Default value is FALSE. If TRUE you have to specify the stat_hash argument and the pvalue_hash argument.
stat_hash	A hash object (hash package required) which contains the cached generated statistics of a SES run in the current dataset, using the current test.
pvalue_hash	A hash object (hash package required) which contains the cached generated p-values of a SES run in the current dataset, using the current test.
threshold	Threshold (suitable values in [0,1]) for assessing p-values significance.
R	The number of permutations, set to 999 by default. There is a trick to avoid doing all permutations. As soon as the number of times the permuted test statistic is more than the observed test statistic is more than 50 (if threshold = 0.05 and R = 999), the p-value has exceeded the signifiacne level (threshold value) and hence the predictor variable is not significant. There is no need to continue do the extra permutations, as a decision has already been made.

Details

If hash = TRUE, all three tests require the arguments 'stat_hash' and 'pvalue_hash' for the hash-based implementation of the statistic test. These hash Objects are produced or updated by each run of SES (if hash = TRUE) and they can be reused in order to speed up next runs of the current statistic test. If "SESoutput" is the output of a SES run, then these objects can be retrieved by SESoutput@hashObject\$stat_hash and the SESoutput@hashObject\$pvalue_hash.

Important: Use these arguments only with the same dataset that was used at initialization. For all the available conditional independence tests that are currently included on the package, please see "?CondIndTests".

For the `testIndGamma` and `testIndNormLog` the F test is used and not the log-likelihood ratio test because both of these regression models have a nuisance parameter. The `testIndNormLog` can be seen as a non linear Gaussian model where the conditional mean is related with the covariate(s) via an exponential function.

`TestIndIGreg` fits an inverse gaussian distribution with a log link. The `testIndIGreg` has some problems due to problems in R's implementation of the inverse gaussian regression with a log link.

Value

A list including:

<code>pvalue</code>	A numeric value that represents the logarithm of the generated p-value due to the count data regression (see references below).
<code>stat</code>	A numeric value that represents the generated statistic due to Poisson regression(see reference below).
<code>stat_hash</code>	The current hash object used for the statistics. See argument <code>stat_hash</code> and details. If argument <code>hash = FALSE</code> this is NULL.
<code>pvalue_hash</code>	The current hash object used for the p-values. See argument <code>stat_hash</code> and details. If argument <code>hash = FALSE</code> this is NULL.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

McCullagh P., and Nelder J.A. (1989). Generalized linear models. CRC press, USA, 2nd edition.

See Also

[testIndReg](#), [testIndNB](#), [testIndZIP](#), [gSquare](#), [CondIndTests](#)

Examples

```
#simulate a dataset with continuous data
dataset <- matrix(rnorm(400 * 50, 1, 5), ncol = 50 )
#the target feature is the last column of the dataset as a vector
target <- rgamma(400, 1, 3)
testIndGamma(target, dataset, xIndex = 24, csIndex = 10)
testIndNormLog(target, dataset, xIndex = 24, csIndex = 10)
#run the SES algorithm using the testIndPois conditional independence test
sesObject <- SES(target, dataset, max_k = 3, threshold = 0.05, test = "testIndGamma");
sesObject2 <- SES(target, dataset, max_k = 3, threshold = 0.05, test = "testIndNormLog");
```

Conditional independence tests for success rates

*Binomial regression conditional independence test for success rates
(binomial)*

Description

The main task of this test is to provide a p-value PVALUE for the null hypothesis: feature 'X' is independent from 'TARGET' given a conditioning set CS. The pvalue is calculated by comparing a binomial logistic regression model based on the conditioning set CS against a model whose regressor are both X and CS. The comparison is performed through a chi-square test with the appropriate degrees of freedom on the difference between the deviances of the two models.

Usage

```
testIndBinom(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL)
```

```
permBinom(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL,
threshold = 0.05, R = 999)
```

```
waldBinom(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL)
```

Arguments

target	A matrix with two two columns, the first one is the number of successes, the cases (integer) and the second one is the totals (integers).
dataset	A numeric matrix or data frame, in case of categorical predictors (factors), containing the variables for performing the test. Rows as samples and columns as features. In the case of "waldBinom" this is strictly a matrix.
xIndex	The index of the variable whose association with the target we want to test.
csIndex	The indices of the variables to condition on. If you have no variables set this equal to 0.
wei	A vector of weights to be used for weighted regression. The default value is NULL and should stay as is, since the totals (second column of the target) is used as weights.
univariateModels	Fast alternative to the hash object for univariate test. List with vectors "pvalues" (p-values), "stats" (statistics) and "flags" (flag = TRUE if the test was succesful) representing the univariate association of each variable with the target. Default value is NULL.

hash	A boolean variable which indicates whether (TRUE) or not (FALSE) to use the hash-based implementation of the statistics of SES. Default value is FALSE. If TRUE you have to specify the stat_hash argument and the pvalue_hash argument.
stat_hash	A hash object (hash package required) which contains the cached generated statistics of a SES run in the current dataset, using the current test.
pvalue_hash	A hash object (hash package required) which contains the cached generated p-values of a SES run in the current dataset, using the current test.
threshold	Threshold (suitable values in [0,1]) for assessing p-values significance.
R	The number of permutations, set to 999 by default. There is a trick to avoid doing all permutations. As soon as the number of times the permuted test statistic is more than the observed test statistic is more than 50 (if threshold = 0.05 and R = 999), the p-value has exceeded the significance level (threshold value) and hence the predictor variable is not significant. There is no need to continue do the extra permutations, as a decision has already been made.

Details

If hash = TRUE, all three tests require the arguments 'stat_hash' and 'pvalue_hash' for the hash-based implementation of the statistic test. These hash Objects are produced or updated by each run of SES (if hash == TRUE) and they can be reused in order to speed up next runs of the current statistic test. If "SESoutput" is the output of a SES run, then these objects can be retrieved by SESoutput@hashObject\$stat_hash and the SESoutput@hashObject\$pvalue_hash.

Important: Use these arguments only with the same dataset that was used at initialization. For all the available conditional independence tests that are currently included on the package, please see "?CondIndTests".

If you have overdispersion, the variance is higher than the mean, a negative binomial is to be used. If you have more zeros than expected under a Poisson model, not overdispersion, then zero inflated Poisson is to be used. This is in fact a logistic regression where the target is the ratio of successes divided by the totals and the weights are the totals.

The log-likelihood ratio test used in "testIndBinom" requires the fitting of two models. The Wald test used in "waldBinom" requires fitting of only one model, the full one. The significance of the variable is examined only. Only continuous (or binary) predictor variables are currently accepted in this test.

Value

A list including:

pvalue	A numeric value that represents the logarithm of the generated p-value due to the count data regression (see references below).
stat	A numeric value that represents the generated statistic due to Poisson regression(see reference below).
stat_hash	The current hash object used for the statistics. See argument stat_hash and details. If argument hash = FALSE this is NULL.
pvalue_hash	The current hash object used for the p-values. See argument stat_hash and details. If argument hash = FALSE this is NULL.

Author(s)

Vincenzo Lagani, Ioannis Tsamardinos, Giorgos Athineou and Michail Tsagris.

R implementation and documentation: Giorgos Athineou <athineou@csd.uoc.gr>, Vincenzo Lagani <vlagani@csd.uoc.gr> and Michail Tsagris <mtsagris@csd.uoc.gr>

References

McCullagh P., and Nelder J.A. (1989). Generalized linear models. CRC press, USA, 2nd edition.

See Also

[testIndLogistic](#), [testIndBeta](#), [testIndReg](#), [CondIndTests](#)

Examples

```
#simulate a dataset with continuous data
dataset <- matrix(runif(400 * 50, 1, 50), ncol = 50 )
#the target feature is the last column of the dataset as a vector
y <- rbinom(400, 10, 0.6)
N <- sample(11:20, 400, replace = TRUE)
target <- cbind(y, N)
results <- testIndBinom(target, dataset, xIndex = 24, csIndex = 10)
results

#run the SES algorithm using the testIndPois conditional independence test
a1 <- SES(target, dataset, max_k = 3, threshold = 0.05, test = "testIndBinom");
a2 <- MMPC(target, dataset, max_k = 3, threshold = 0.05, test = "testIndBinom");
```

Conditional independence tests for survival data

Conditional independence test for survival data

Description

The main task of this test is to provide a p-value PVALUE for the null hypothesis: feature 'X' is independent from 'TARGET' given a conditioning set CS. This test can be based on the Cox (semi-parametric) regression or on the Weibull (parametric) regression.

Usage

```
censIndCR(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL)
```

```
censIndWR(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL)
```

```
censIndER(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL)
```

```
permCR(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL,
threshold = 0.05, R = 999)
```

```
permWR(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL,
threshold = 0.05, R = 999)
```

```
permER(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL,
threshold = 0.05, R = 999)
```

```
waldCR(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL)
```

```
waldWR(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL)
```

```
waldER(target, dataset, xIndex, csIndex, wei = NULL,
univariateModels = NULL, hash = FALSE, stat_hash = NULL, pvalue_hash = NULL)
```

Arguments

target	A Survival object (class <code>Surv</code> from package <code>survival</code>) containing the time to event data (time) and the status indicator vector (event). View Surv documentation for more information.
dataset	A numeric matrix or data frame, in case of categorical predictors (factors), containing the variables for performing the test. Rows as samples and columns as features. In the cases of "waldCR", "waldWR", "waldER", "waldCR", "permWR" and "permER" this is strictly a matrix.
xIndex	The index of the variable whose association with the target we want to test.
csIndex	The indices of the variables to condition on.
wei	A vector of weights to be used for weighted regression. The default value is <code>NULL</code> .
univariateModels	Fast alternative to the hash object for univariate test. List with vectors "pvalues" (p-values), "stats" (statistics) and "flags" (flag = <code>TRUE</code> if the test was succesful) representing the univariate association of each variable with the target. Default value is <code>NULL</code> .
hash	A boolean variable which indicates whether (<code>TRUE</code>) or not (<code>FALSE</code>) to use the hash-based implementation of the statistics of SES. Default value is <code>FALSE</code> . If <code>TRUE</code> you have to specify the <code>stat_hash</code> argument and the <code>pvalue_hash</code> argument.
stat_hash	A hash object (hash package required) which contains the cached generated statistics of a SES run in the current dataset, using the current test.

pvalue_hash	A hash object (hash package required) which contains the cached generated p-values of a SES run in the current dataset, using the current test.
threshold	Threshold (suitable values in [0,1]) for assessing p-values significance.
R	The number of permutations, set to 999 by default. There is a trick to avoid doing all permutations. As soon as the number of times the permuted test statistic is more than the observed test statistic is more than 50 (in this example case), the p-value has exceeded the significance level (threshold value) and hence the predictor variable is not significant. There is no need to continue do the extra permutations, as a decision has already been made.

Details

The censIndCR implies the Cox (semiparametric) regression, the censIndWR the Weibull (parametric) regression and the censIndER the exponential (parametric) regression, which is a special case of the Weibull regression (when shape parameter is 1). **Note:** When there are observations with zero values (time=0) the Weibull and Exponential regressions will not work. Only Cox regression will run.

If hash = TRUE, censIndCR, censIndWR and censIndER require the arguments 'stat_hash' and 'pvalue_hash' for the hash-based implementation of the statistic test. These hash Objects are produced or updated by each run of SES (if hash == TRUE) and they can be reused in order to speed up next runs of the current statistic test. If "SESoutput" is the output of a SES run, then these objects can be retrieved by SESoutput@hashObject\$stat_hash and the SESoutput@hashObject\$pvalue_hash.

Important: Use these arguments only with the same dataset that was used at initialization.

For all the available conditional independence tests that are currently included on the package, please see "?CondIndTests".

The log-likelihood ratio test used in "censIndCR", "censIndWR" and "censIndER" requires the fitting of two models. The Wald tests used in "waldCR", "waldWR" and "waldER" requires fitting of only one model, the full one. The significance of the variable is examined only. Only continuous (or binary) predictor variables are currently accepted in this test.

Value

A list including:

pvalue	A numeric value that represents the logarithm of the generated p-value.
stat	A numeric value that represents the generated statistic.
stat_hash	The current hash object used for the statistics. See argument stat_hash and details. If argument hash = FALSE this is NULL.
pvalue_hash	The current hash object used for the p-values. See argument stat_hash and details. If argument hash = FALSE this is NULL.

Note

This test uses the functions coxph and Surv of the package survival and the function anova (analysis of variance) of the package stats.

Author(s)

R implementation and documentation: Vincenzo Lagani <vlagani@csd.uoc.gr>, Giorgos Athineou <athineou@csd.uoc.gr>

References

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See Also

[SES](#), [censIndWR](#), [testIndFisher](#), [gSquare](#), [testIndLogistic](#), [Surv](#), [anova](#), [CondIndTests](#)

Examples

```
#create a survival simulated dataset
dataset <- matrix(runif(1000 * 20, 1, 100), nrow = 1000 , ncol = 20)
dataset <- as.data.frame(dataset);
timeToEvent <- numeric(1000)
event <- numeric(1000)
ca <- numeric(1000)
for(i in 1:1000) {
  timeToEvent[i] <- dataset[i, 1] + 0.5 * dataset[i, 10] + 2 * dataset[i, 15] + runif(1, 0, 3);
  event[i] <- sample( c(0, 1), 1)
  ca[i] <- runif(1, 0, timeToEvent[i] - 0.5)
  if(event[i] == 0) timeToEvent[i] = timeToEvent[i] - ca[i]
}

require(survival, quietly = TRUE)

#init the Surv object class feature
target <- Surv(time = timeToEvent, event = event)
#run the censIndCR conditional independence test
censIndCR( target, dataset, xIndex = 12, csIndex = c(5, 7, 4) )
# run the SESC algorithm
## Not run:
ses1 <- SES(target, dataset, max_k = 1, threshold = 0.05, test = "censIndCR");
ses2 <- SES(target, dataset, max_k = 1, threshold = 0.05, test = "censIndWR");

## End(Not run)
```

Conditional independence tests with and without permutation p-value
Conditional independence test for continuous class variables with and without permutation based p-value

Description

The main task of this test is to provide a permutation based p-value PVALUE for the null hypothesis: feature 'X' is independent from 'TARGET' given a conditioning set CS.

Usage

```
condi(ind1, ind2, cs, dat, type = "pearson", rob = FALSE, R = 1)
dist.condi(ind1, ind2, cs, dat, type = NULL, rob = FALSE, R = 499)
cat.ci(ind1, ind2, cs, dat, type, rob = FALSE, R = 1)
```

Arguments

ind1	The index of the one variable to be considered.
ind2	The index of the other variable to be considered.
cs	The index or indices of the conditioning set of variable(s). If you have no variables set this equal to 0.
dat	A matrix with the data. In the case of "cat.ci" the minimum must be 0, i.e. the data must be like 0, 1, 2 and NOT 1, 2, 3... There is a C++ code behind and the minimum must be 0.
type	Do you want the Pearson (type = "pearson") or the Spearman (type = "spearman") correlation to be used. For "dist.condi" this is an obsolete argument but it requires to exist when it is used in the PC algorithm. For "cat.ci" this should be a vector with the levels, the number of distinct (different) values of each categorical variable. Its length is equal to the number of variables used in the test (2 + the number of conditioning variables).
rob	If you choose type="pearson" then you can sapecify whether you want a robust version of it. For "dist.condi" and "cat.ci" this is an obsolete argument but it requires to exist when it is used in the PC algorithm.
R	If R = 1 then the asymptotic p-value is calculated. If R > 1 a permutation based p-value is returned. For the distance correlation based test, this is set to 499 by default and is used in the partial correlaiton test only.

Details

This test is currently designed for usage by the PC algorithm. The Fisher conditional independence test which is based on the Pearson or Spearman correlation coefficients is much faster than the distance based (partial) correlation test.

The distance correlation can handle non linear relationships as well. The p-value for the partial distance correlation is calculated via permutations and is slow.

Value

A vector including the test statistic, its associated p-value and the relevant degrees of freedom. In the case of a permutation based p-value, the returned test statistic is the observed test statistic divided by the relevant degrees of freedom (Pearson and Spearman correlation coefficients only). This is for the case of ties between many permutation based p-values. The PC algorithm choose a pair of variables based on the p-values. If they are equal it will use the test statistic.

Author(s)

Michail Tsagris

R implementation and documentation: Giorgos Athineou <athineou@csd.uoc.gr> and Michail Tsagris <mtsagris@csd.uoc.gr>

References

Hampel F. R., Ronchetti E. M., Rousseeuw P. J., and Stahel W. A. (1986). Robust statistics: the approach based on influence functions. John Wiley & Sons.

Lee Rodgers J., and Nicewander W.A. (1988). "Thirteen ways to look at the correlation coefficient". The American Statistician 42(1): 59-66.

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Szekely G.J. and Rizzo M.L. (2013). Energy statistics: A class of statistics based on distances. Journal of Statistical Planning and Inference 143(8): 1249–1272.

See Also

[testIndFisher](#), [testIndSpearman](#), [pc.skel](#), [gSquare](#), [CondIndTests](#)

Examples

```
#simulate a dataset with continuous data
dataset <- matrix(runif(500 * 5, 1, 100), nrow = 5 )
testIndFisher(dataset[, 1], dataset[, -1], xIndex = 1, csIndex = 2)
condi(ind1 = 1, ind2 = 2, cs = 3, dataset, R = 1)
condi(ind1 = 1, ind2 = 2, cs = 3, dataset, R = 999)
dist.condi(ind1 = 1, ind2 = 2, 0, dataset)
dist.condi(ind1 = 1, ind2 = 2, cs = 3, dataset, R = 99)
```

 Constraint based feature selection algorithms

SES: Feature selection algorithm for identifying multiple minimal, statistically-equivalent and equally-predictive feature signatures
MMPC: Feature selection algorithm for identifying minimal feature subsets

Description

SES algorithm follows a forward-backward filter approach for feature selection in order to provide minimal, highly-predictive, statistically-equivalent, multiple feature subsets of a high dimensional dataset. See also Details. MMPC algorithm follows the same approach without generating multiple feature subsets.

Usage

```
SES(target, dataset, max_k = 3, threshold = 0.05, test = NULL, ini = NULL,
wei = NULL, user_test = NULL, hash = FALSE, hashObject = NULL,
ncores = 1)
```

```
MMPC(target, dataset, max_k = 3, threshold = 0.05, test = NULL, ini = NULL,
wei = NULL, user_test = NULL, hash = FALSE, hashObject = NULL,
ncores = 1, backward = FALSE)
```

```
wald.ses(target, dataset, max_k = 3, threshold = 0.05, test = NULL, ini = NULL,
wei = NULL, user_test = NULL, hash = FALSE, hashObject = NULL,
ncores = 1)
```

```
wald.mmpc(target, dataset, max_k = 3, threshold = 0.05, test = NULL, ini = NULL,
wei = NULL, user_test = NULL, hash = FALSE, hashObject = NULL,
ncores = 1, backward = FALSE)
```

```
perm.ses(target, dataset, max_k = 3, threshold = 0.05, test = NULL, ini = NULL,
wei = NULL, user_test = NULL, hash=FALSE, hashObject = NULL,
R = 999, ncores = 1)
```

```
perm.mmpc(target, dataset, max_k = 3, threshold = 0.05, test = NULL, ini = NULL,
wei = NULL, user_test = NULL, hash=FALSE, hashObject = NULL,
R = 999, ncores = 1, backward = FALSE)
```

Arguments

target	The class variable. Provide either a string, an integer, a numeric value, a vector, a factor, an ordered factor or a Surv object. See also Details.
dataset	The data-set; provide either a data frame or a matrix (columns = variables , rows = samples).

max_k	The maximum conditioning set to use in the conditional independence test (see Details). Integer, default value is 3.
threshold	Threshold (suitable values in [0,1]) for assessing p-values significance. Default value is 0.05.
test	The conditional independence test to use. Default value is NULL. See also CondIndTests .
ini	This is supposed to be a list. After running SES or MMPC with some hyper-parameters you might want to run SES again with different hyper-parameters. To avoid calculating the univariate associations (first step of SES and of MMPC) again, you can extract them from the first run of SES and plug them here. This can speed up the second run (and subsequent runs of course) by 50%. See the details and the argument "univ" in the output values.
wei	A vector of weights to be used for weighted regression. The default value is NULL.
user_test	A user-defined conditional independence test (provide a closure type object). Default value is NULL. If this is defined, the "test" argument is ignored.
hash	A boolean variable which indicates whether (TRUE) or not (FALSE) to store the statistics calculated during SES execution in a hash-type object. Default value is FALSE. If TRUE a hashObject is produced.
hashObject	A List with the hash objects generated in a previous run of SES or MMPC. Each time SES runs with "hash=TRUE" it produces a list of hashObjects that can be re-used in order to speed up next runs of SES or MMPC. Important: the generated hashObjects should be used only when the same dataset is re-analyzed, possibly with different values of max_k and threshold.
R	The number of permutations, set to 999 by default. There is a trick to avoid doing all permutations. As soon as the number of times the permuted test statistic is more than the observed test statistic is more than 50 (if threshold = 0.05 and R = 999), the p-value has exceeded the significance level (threshold value) and hence the predictor variable is not significant. There is no need to continue do the extra permutations, as a decision has already been made.
ncores	How many cores to use. This plays an important role if you have tens of thousands of variables or really large sample sizes and tens of thousands of variables and a regression based test which requires numerical optimisation. In other cases it will not make a difference in the overall time (in fact it can be slower). The parallel computation is used in the first step of the algorithm, where univariate associations are examined, those take place in parallel. We have seen a reduction in time of 50% with 4 cores in comparison to 1 core. Note also, that the amount of reduction is not linear in the number of cores.
backward	If TRUE, the backward (or symmetry correction) phase will be implemented. This removes any falsely included variables in the parents and children set of the target variable. It call the <code>link{mmpcbackphase}</code> for this purpose. For <code>perm.ses</code> and <code>wald.ses</code> this is not yet applicable.

Details

The SES function implements the Statistically Equivalent Signature (SES) algorithm as presented in "Tsamardinos, Lagani and Pappas, HSCBB 2012" <http://www.mensxmachina.org/publications/discovering-multiple-equivalent-biomarker-signatures/>

The MMPC function implements the MMPC algorithm as presented in "Tsamardinos, Brown and Aliferis. The max-min hill-climbing Bayesian network structure learning algorithm" http://www.dsl-lab.org/supplements/mmhc_paper/paper_online.pdf

The output value "univ" along with the output value "hashObject" can speed up the computations of subsequent runs of SES and MMPC. The first run with a specific pair of hyper-parameters (threshold and max_k) the univariate associations tests and the conditional independence tests (test statistic and natural logarithm of their corresponding p-values) are stored and returned. In the next run(s) with different pair(s) of hyper-parameters you can use this information to save time. With a few thousands of variables you will see the difference, which can be up to 50%. For the non robust correlation based tests, the difference may not be significant though, because the unconditional correlation coefficients are calculated very efficiently.

The max_k option: the maximum size of the conditioning set to use in the conditioning independence test. Larger values provide more accurate results, at the cost of higher computational times. When the sample size is small (e.g., < 50 observations) the max_k parameter should be ≤ 5 , otherwise the conditional independence test may not be able to provide reliable results.

If the dataset (predictor variables) contains missing (NA) values, they will automatically be replaced by the current variable (column) mean value with an appropriate warning to the user after the execution.

If the target is a single integer value or a string, it has to correspond to the column number or to the name of the target feature in the dataset. In any other case the target is a variable that is not contained in the dataset.

If the current 'test' argument is defined as NULL or "auto" and the user_test argument is NULL then the algorithm automatically selects the best test based on the type of the data. Particularly:

- if the target is a factor, the multinomial or the binary logistic regression is used. If the target has two values only, binary logistic regression will be used.
- if target is an ordered factor, ordinal regression is used in the logistic test. Hence, if you want to use multinomial or ordinal logistic regression, make sure your target is factor.
- if target is a numerical vector and the dataset is a matrix or a data.frame with continuous variables, the Fisher conditional independence test is used. If the dataset is a data.frame and there are categorical variables, linear regression is used.
- if target is discrete numerical (counts), the Poisson regression conditional independence test is used. If there are only two values, the binary logistic regression is to be used.
- if target is a Surv object, a Survival conditional independence test is used.
- if target is a matrix with at least 2 columns, the multivariate linear regression is used.
- if target is a 2 column matrix whose columns are the number of successes and the number of trials (first and second column respectively) the testIndBinom should be used.

Conditional independence test functions to be passed through the user_test argument should have the same signature of the included test. See [testIndFisher](#) for an example.

For all the available conditional independence tests that are currently included on the package, please see [CondIndTests](#).

If two or more p-values are below the machine epsilon (`.Machine$double.eps` which is equal to $2.220446e-16$), all of them are set to 0. To make the comparison or the ordering feasible we use the logarithm of the p-value. The max-min heuristic though, requires comparison and an ordering of the p-values. Hence, all conditional independence tests calculate the logarithm of the p-value.

If there are missing values in the dataset (predictor variables) columnwise imputation takes place. The median is used for the continuous variables and the mode for categorical variables. It is a naive and not so clever method. For this reason the user is encouraged to make sure his data contain no missing values.

If you have percentages, in the (0, 1) interval, they are automatically mapped into R by using the logit transformation. If you set the test to `testIndBeta`, beta regression is used. If you have compositional data, positive multivariate data where each vector sums to 1, with NO zeros, they are also mapped into the Euclidean space using the additive log-ratio (multivariate logit) transformation (Aitchison, 1986).

If you use `testIndSpearman` (argument "test"), the ranks of the data calculated and those are used in the calculations. This speeds up the whole procedure.

As a rule of thumb you can try this. If for example you have counts and want to see which model fits best, there are two ways. Calculate the mean and the variance. If they are similar, use the Poisson instead of the negative binomial as it is much faster. If you are not convinced, you can either use the negative binomial or do the following simulation study.

```
# x <- matrix(rnorm(n * 1000), ncol = 1000) # a <- Rfast::univglms(y, x) # hist(a[, 2]) ## histogram of the p-values
```

If the histogram shows a uniform distribution, use the Poisson regression. If the histogram is not uniform, then repeat the simulation but with a negative binomial distribution. If the histogram is again not flat, then another model is necessary. If the data come from a Poisson or negative binomial, the histogram with a negative binomial regression will be flat. If the data come from a negative binomial, the histogram with a Poisson will not be uniform.

On the same page, if you have many zeros, try `Rfast::zip.mle` and see whether there are grounds to facilitate the use of a zero inflated Poisson model. Otherwise, do a simulation study like before.

Value

The output of the algorithm is an object of the class 'SESoutput' for SES or 'MMPCoutput' for MMPC including:

<code>selectedVars</code>	The selected variables, i.e., the signature of the target variable.
<code>selectedVarsOrder</code>	The order of the selected variables according to increasing pvalues.
<code>queues</code>	A list containing a list (queue) of equivalent features for each variable included in <code>selectedVars</code> . An equivalent signature can be built by selecting a single feature from each queue. Featured only in SES.
<code>signatures</code>	A matrix reporting all equivalent signatures (one signature for each row). Featured only in SES.
<code>hashObject</code>	The hashObject caching the statistic calculated in the current run.

<code>pvalues</code>	For each feature included in the dataset, this vector reports the strength of its association with the target in the context of all other variables. Particularly, this vector reports the max p-values found when the association of each variable with the target is tested against different conditional sets. Lower values indicate higher association.
<code>stats</code>	The statistics corresponding to "pvalues" (higher values indicates higher association).
<code>univ</code>	This is a list with the univariate associations. The test statistics and their corresponding logged p-values, along with their flag (1 if the test was performed and 0 otherwise). This list is very important for subsequent runs of SES with different hyper-parameters. After running SES with some hyper-parameters you might want to run SES again with different hyper-parameters. To avoid calculating the univariate associations (first step of SES or MMPC) again, you can take this list from the first run of SES and plug it in the argument "ini" in the next run(s) of SES or MMPC. This can speed up the second run (and subsequent runs of course) by 50%. See the argument "univ" in the output values.
<code>max_k</code>	The <code>max_k</code> option used in the current run.
<code>threshold</code>	The <code>threshold</code> option used in the current run.
<code>n.tests</code>	If you have set <code>hash = TRUE</code> , then the number of tests performed by SES or MMPC will be returned. If you have not set this to <code>TRUE</code> , the number of univariate associations will be returned. So be careful with this number.
<code>runtime</code>	The run time of the algorithm. A numeric vector. The first element is the user time, the second element is the system time and the third element is the elapsed time.
<code>test</code>	The character name of the statistic test used.

Generic Functions implemented for `SESoutput` Object:

```
plot(object=SESoutput, mode="all")
```

Plots the generated pvalues (using `barplot`) of the current `SESoutput` object in comparison to the threshold.

Argument `mode` can be either "all" or "partial" for the first 500 pvalues of the object.

Note

The packages required by the SES and MMPC algorithm operations are:

hash : for the hash-based implementation

quantreg: for the quantile (median) regression

MASS: for negative binomial regression

nnet : also `require(stats)` and `require(MASS)` for the `testIndLogistic` test

and **ordinal** : also `require(stats)` and `require(MASS)` for the `testIndLogistic` test

survival : for the `censIndCR`, `censIndWR` and the `censIndER` tests

doParallel: for parallel computations

lme4: for (generalised) linear mixed models

Rfast: for many fast functions.

Author(s)

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References

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Brown, L. E., Tsamardinos, I., & Aliferis, C. F. (2004). A novel algorithm for scalable and accurate Bayesian network learning. Medinfo, 711-715.

Tsamardinos, Brown and Aliferis (2006). The max-min hill-climbing Bayesian network structure learning algorithm. Machine learning, 65(1), 31-78.

See Also

[CondIndTests](#), [cv.ses](#)

Examples

```
set.seed(123)
require("hash", quietly = TRUE)

#simulate a dataset with continuous data
dataset <- matrix(runif(1000 * 200, 1, 100), ncol = 200)

#define a simulated class variable
target <- 3 * dataset[, 10] + 2 * dataset[, 100] + 3 * dataset[, 20] + rnorm(1000, 0, 5)

# define some simulated equivalences
dataset[, 15] <- dataset[, 10] + rnorm(1000, 0, 2)
dataset[, 150] <- dataset[, 100] + rnorm(1000, 0, 2)
dataset[, 130] <- dataset[, 100] + rnorm(1000, 0, 2)

# run the SES algorithm
sesObject <- SES(target , dataset, max_k = 5, threshold = 0.05, test = "testIndFisher",
hash = TRUE, hashObject = NULL);

# print summary of the SES output
summary(sesObject);
# plot the SES output
# plot(sesObject, mode = "all");
# get the queues with the equivalences for each selected variable
sesObject@queues
#get the generated signatures
```

```

sesObject@signatures;

# re-run the SES algorithm with the same or different configuration
# under the hash-based implementation of retrieving the statistics
# in the SAME dataset (!important)
hashObj <- sesObject@hashObject;
sesObject2 <- SES(target, dataset, max_k = 2, threshold = 0.01, test = "testIndFisher",
hash = TRUE, hashObject = hashObj);

sesObject3 <- SES(target, dataset, max_k = 2, threshold = 0.01, test = "testIndFisher",
ini = sesObject@univ, hash = TRUE, hashObject = hashObj);

# retrieve the results: summary, plot, sesObject2@...)
summary(sesObject2)
# get the run time
sesObject@runtime;
sesObject2@runtime;
sesObject3@runtime;

# MMPC algorithm
mmpcObject <- MMPC(target, dataset, max_k = 3, threshold = 0.05, test="testIndFisher");
mmpcObject@selectedVars
mmpcObject@runtime

```

Constraint based feature selection algorithms for longitudinal and clustered data
SES.temporal/SES.gee: Feature selection algorithm for identifying multiple minimal, statistically-equivalent and equally-predictive feature signatures
MMPC.temporal/MMPC.gee: Feature selection algorithm for identifying minimal feature subsets

Description

SES.temporal algorithm follows a forward-backward filter approach for feature selection in order to provide minimal, highly-predictive, statistically-equivalent, multiple feature subsets of a high dimensional dataset. See also Details. MMPC.temporal algorithm follows the same approach without generating multiple feature subsets. They are both adapted to longitudinal target variables.

Usage

```

SES.temporal(target, reps = NULL, group, dataset, max_k = 3, threshold = 0.05,
test = NULL, ini = NULL, wei = NULL, user_test = NULL, hash = FALSE,
hashObject = NULL, slopes = FALSE, ncores = 1)

```

```

MMPC.temporal(target, reps = NULL, group, dataset, max_k = 3, threshold = 0.05,
test = NULL, ini = NULL, wei = NULL, user_test = NULL, hash = FALSE,
hashObject = NULL, slopes = FALSE, ncores = 1)

```

```
MMPC.gee(target, reps = NULL, group, dataset, max_k = 3, threshold = 0.05,
test = NULL, ini = NULL, wei = NULL, user_test = NULL, hash = FALSE,
hashObject = NULL, correl = "exchangeable", se = "jack", ncores = 1)
```

```
SES.gee(target, reps = NULL, group, dataset, max_k = 3, threshold = 0.05,
test = NULL, ini = NULL, wei = NULL, user_test = NULL, hash = FALSE,
hashObject = NULL, correl = "exchangeable", se = "jack", ncores = 1)
```

Arguments

target	The class variable. Provide a vector with continuous (normal), binary (binomial) or discrete (Poisson) data.
reps	A numeric vector containing the time points of the subjects. It's length is equal to the length of the target variable. If you have clustered data, leave this NULL.
group	A numeric vector containing the subjects or groups. It must be of the same length as target.
dataset	The dataset; provide either a data frame or a matrix (columns = variables , rows = samples). Currently, only continuous datasets are supported.
max_k	The maximum conditioning set to use in the conditional independence test (see Details). Integer, default value is 3.
threshold	Threshold (suitable values in [0, 1]) for assessing p-values significance. Default value is 0.05.
test	The conditional independence test to use. Default value is NULL. Currently, the only available conditional independence test are the testIndGLMMLogistic , testIndGLMMPois and testIndGLMMReg which fit linear mixed models.
ini	This is a supposed to be a list. After running SES or MMPC with some hyper-parameters you might want to run SES again with different hyper-parameters. To avoid calculating the univariate associations (first step of SES and of MPPC) again, you can extract them from the first run of SES and plug them here. This can speed up the second run (and subsequent runs of course) by 50%. See the details and the argument "univ" in the output values.
wei	A vector of weights to be used for weighted regression. The default value is NULL.
user_test	A user-defined conditional independence test (provide a closure type object). Default value is NULL. If this is defined, the "test" argument is ignored.
hash	A boolean variable which indicates whether (TRUE) or not (FALSE) to store the statistics calculated during SES execution in a hash-type object. Default value is FALSE. If TRUE a hashObject is produced.
hashObject	A List with the hash objects generated in a previous run of SES.temporal. Each time SES runs with "hash=TRUE" it produces a list of hashObjects that can be re-used in order to speed up next runs of SES. Important: the generated hashObjects should be used only when the same dataset is re-analyzed, possibly with different values of max_k and threshold.
slopes	Should random slopes for the ime effect be fitted as well? Default value is FALSE.

correl	The correlation structure. For the Gaussian, Logistic, Poisson and Gamma regression this can be either "exchangeable" (compound symmetry, suitable for clustered data) or "ar1" (AR(1) model, suitable for longitudinal data). For the ordinal logistic regression its only the "exchangeable" correlation structure. This is obsolete for "testIndGEEOrdinal", but is here for compatibility reasons.
se	<p>The method for estimating standard errors. This is very important and crucial. The available options for Gaussian, Logistic, Poisson and Gamma regression are: a) 'san.se', the usual robust estimate. b) 'jack': if approximate jackknife variance estimate should be computed. c) 'j1s': if 1-step jackknife variance estimate should be computed and d) 'fj': logical indicating if fully iterated jackknife variance estimate should be computed. If you have many clusters (sets of repeated measurements) "san.se" is fine as it is asymptotically correct, plus jackknife estimates will take longer. If you have a few clusters, then maybe it's better to use jackknife estimates.</p> <p>The jackknife variance estimator was suggested by Paik (1988), which is quite suitable for cases when the number of subjects is small ($K < 30$), as in many biological studies. The simulation studies conducted by Ziegler et al. (2000) and Yan and Fine (2004) showed that the approximate jackknife estimates are in many cases in good agreement with the fully iterated ones. This is obsolete for "testIndGEEOrdinal", but is here for compatibility reasons.</p>
ncores	How many cores to use. This plays an important role if you have tens of thousands of variables or really large sample sizes and tens of thousands of variables and a regression based test which requires numerical optimisation. In other cases it will not make a difference in the overall time (in fact it can be slower). The parallel computation is used in the first step of the algorithm, where univariate associations are examined, those take place in parallel. We have seen a reduction in time of 50% with 4 cores in comparison to 1 core. Note also, that the amount of reduction is definitely not linear in the number of cores.

Details

The SES.temporal function implements the Statistically Equivalent Signature (SES) algorithm as presented in "Tsamardinos, Lagani and Pappas, HSCBB 2012" adapted to longitudinal data. The citation for this is "Tsagris, Lagani and tsamardinos, (2018)". These functions presented here are for the **temporal-logntudinal scenario**.

<http://www.mensxmachina.org/publications/discovering-multiple-equivalent-biomarker-signatures/>

The MMPC function implements the MMPC algorithm as presented in "Tsamardinos, Brown and Aliferis. The max-min hill-climbing Bayesian network structure learning algorithm" adapted to longitudinal data. http://www.dsl-lab.org/supplements/mmhc_paper/paper_online.pdf

The output value "univ" along with the output value "hashObject" can speed up the computations of subsequent runs of SES and MMPC. The first run with a specific pair of hyper-parameters (threshold and max_k) the univariate associations tests and the conditional independence tests (test statistic and logarithm of their corresponding p-values) are stored and returned. In the next run(s) with different pair(s) of hyper-parameters you can use this information to save time. With a few thousands of variables you will see the difference, which can be up to 50%.

The max_k option: the maximum size of the conditioning set to use in the conditioning independence test. Larger values provide more accurate results, at the cost of higher computational times.

When the sample size is small (e.g., < 50 observations) the `max_k` parameter should be ≤ 5 , otherwise the conditional independence test may not be able to provide reliable results.

If the dataset contains missing (NA) values, they will automatically be replaced by the current variable (column) mean value with an appropriate warning to the user after the execution.

If the target is a single integer value or a string, it has to correspond to the column number or to the name of the target feature in the dataset. In any other case the target is a variable that is not contained in the dataset.

If the current 'test' argument is defined as NULL or "auto" and the `user_test` argument is NULL then the algorithm automatically selects only available, which is `testIndGLMMReg`.

Conditional independence test functions to be passed through the `user_test` argument should have the same signature of the included test. See "`?testIndFisher`" for an example.

For all the available conditional independence tests that are currently included on the package, please see "`?CondIndTests`".

If two or more p-values are below the machine epsilon (`.Machine$double.eps` which is equal to $2.220446e-16$), all of them are set to 0. To make the comparison or the ordering feasible we use the logarithm of the p-value. The max-min heuristic though, requires comparison and an ordering of the p-values. Hence, all conditional independence tests calculate the logarithm of the p-value.

If there are missing values in the dataset (predictor variables) columnwise imputation takes place. The median is used for the continuous variables and the mode for categorical variables. It is a naive and not so clever method. For this reason the user is encouraged to make sure his data contain no missing values.

If you have percentages, in the (0, 1) interval, they are automatically mapped into R by using the logit transformation and a linear mixed model is fitted. If you have binary data, logistic mixed regression is applied and if you have discrete data (counts), Poisson mixed regression is applied.

Value

The output of the algorithm is an object of the class 'SES.temporal.output' for SES.temporal or 'MMPC.temporal.output' for MMPC.temporal including:

<code>selectedVars</code>	The selected variables, i.e., the signature of the target variable.
<code>selectedVarsOrder</code>	The order of the selected variables according to increasing p-values.
<code>queues</code>	A list containing a list (queue) of equivalent features for each variable included in <code>selectedVars</code> . An equivalent signature can be built by selecting a single feature from each queue. Featured only in SES.
<code>signatures</code>	A matrix reporting all equivalent signatures (one signature for each row). Featured only in SES.
<code>hashObject</code>	The <code>hashObject</code> caching the statistic calculated in the current run.
<code>pvalues</code>	For each feature included in the dataset, this vector reports the strength of its association with the target in the context of all other variables. Particularly, this vector reports the max p-values found when the association of each variable with the target is tested against different conditional sets. Lower values indicate higher association.

<code>stats</code>	The statistics corresponding to "pvalues" (higher values indicates higher association).
<code>univ</code>	This is a list with the univariate associations. The test statistics and their corresponding logge p-values. This list is very important for subsequent runs of SES with different hyper-parameters. After running SES with some hyper-parameters you might want to run SES again with different hyper-parameters. To avoid calculating the univariate associations (first step of SES or MMPC) again, you can take this list from the first run of SES and plug it in the argument "ini" in the next run(s) of SES or MMPC. This can speed up the second run (and subsequent runs of course) by 50%. See the argument "univ" in the output values.
<code>max_k</code>	The <code>max_k</code> option used in the current run.
<code>threshold</code>	The <code>threshold</code> option used in the current run.
<code>slope</code>	Whether random slopes for the time effects were used or not, TRUE or FALSE.
<code>runtime</code>	The run time of the algorithm. A numeric vector. The first element is the user time, the second element is the system time and the third element is the elapsed time.

Generic Functions implemented for SESoutput Object:

```
plot(object=SES.temporal.output, mode="all")
```

Plots the generated pvalues (using barplot) of the current SESoutput object in comparison to the threshold. Argument mode can be either "all" or "partial" for the first 500 pvalues of the object.

Author(s)

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See Also

[CondIndTests](#), [testIndGLMMReg](#)

Examples

```
## Not run:
require(lme4)
data(sleepstudy)
attach(sleepstudy)
x <- matrix(rnorm(180 * 100),ncol = 100) ## unrelated predictor variables
m1 <- SES.temporal(Reaction, Days, Subject, x)
m2 <- MMPC.temporal(Reaction, Days, Subject, x)

## End(Not run)
```

Constraint based feature selection algorithms for multiple datasets

ma.ses: Feature selection algorithm for identifying multiple minimal, statistically-equivalent and equally-predictive feature signatures with multiple datasets
ma.mmpc: Feature selection algorithm for identifying minimal feature subsets with multiple datasets

Description

SES algorithm follows a forward-backward filter approach for feature selection in order to provide minimal, highly-predictive, statistically-equivalent, multiple feature subsets of two or more high dimensional datasets. See also Details. MMPC algorithm follows the same approach without generating multiple feature subsets.

Usage

```
ma.ses(target, dataset, ina, statistic = FALSE, max_k = 3, threshold = 0.05,
test = NULL, ini = NULL, user_test = NULL, hash = FALSE, hashObject = NULL,
ncores = 1)
```

```
ma.mmpc(target, dataset, ina, statistic = FALSE, max_k = 3, threshold = 0.05,
test = NULL, ini = NULL, user_test = NULL, hash = FALSE, hashObject = NULL,
ncores = 1, backward = FALSE)
```

Arguments

target	The class variable. Provide either a string, an integer, a numeric value, a vector, a factor, an ordered factor or a Surv object. See also Details.
dataset	The data-set; provide either a data frame or a matrix (columns = variables , rows = samples). Alternatively, provide an ExpressionSet (in which case rows are samples and columns are features, see bioconductor for details).
ina	A numerical vector indicating the dataset. The numbers must be 1, 2, 3,...
statistic	A boolean variable indicating whether the test statistics (TRUE) or the p-values should be combined (FALSE). See the details about this.
max_k	The maximum conditioning set to use in the conditional independence test (see Details). Integer, default value is 3.
threshold	Threshold (suitable values in [0,1]) for assessing p-values significance. Default value is 0.05.
test	The conditional independence test to use. Default value is NULL. See also CondIndTests .
ini	This is a supposed to be a list. After running SES or MMPC with some hyper-parameters you might want to run SES again with different hyper-parameters. To avoid calculating the univariate associations (first step of SES and of MPPC) again, you can extract them from the first run of SES and plug them here. This can speed up the second run (and subsequent runs of course) by 50%. See the details and the argument "univ" in the output values.
user_test	A user-defined conditional independence test (provide a closure type object). Default value is NULL. If this is defined, the "test" argument is ignored.
hash	A boolean variable which indicates whether (TRUE) or not (FALSE) to store the statistics calculated during SES execution in a hash-type object. Default value is FALSE. If TRUE a hashObject is produced.
hashObject	A List with the hash objects generated in a previous run of SES or MMPC. Each time SES runs with "hash=TRUE" it produces a list of hashObjects that can be re-used in order to speed up next runs of SES or MMPC. Important: the generated hashObjects should be used only when the same dataset is re-analyzed, possibly with different values of max_k and threshold.
ncores	How many cores to use. This plays an important role if you have tens of thousands of variables or really large sample sizes and tens of thousands of variables and a regression based test which requires numerical optimisation. In other cases it will not make a difference in the overall time (in fact it can be slower). The

parallel computation is used in the first step of the algorithm, where univariate associations are examined, those take place in parallel. We have seen a reduction in time of 50% with 4 cores in comparison to 1 core. Note also, that the amount of reduction is not linear in the number of cores.

`backward` If TRUE, the backward (or symmetry correction) phase will be implemented. This removes any falsely included variables in the parents and children set of the target variable. This is an experimental stage, so do not trust too much or even better do not use.

Details

This is more at an experimental stage at the present.

The SES function implements the Statistically Equivalent Signature (SES) algorithm as presented in "Tsamardinos, Lagani and Pappas, HSCBB 2012" <http://www.mensxmachina.org/publications/discovering-multiple-equivalent-biomarker-signatures/>

The MMPC function implements the MMPC algorithm as presented in "Tsamardinos, Brown and Aliferis. The max-min hill-climbing Bayesian network structure learning algorithm" http://www.dsl-lab.org/supplements/mmhc_paper/paper_online.pdf

The output value "univ" along with the output value "hashObject" can speed up the computations of subsequent runs of SES and MMPC. The first run with a specific pair of hyper-parameters (threshold and max_k) the univariate associations tests and the conditional independence tests (test statistic and logarithm of their corresponding p-values) are stored and returned. In the next run(s) with different pair(s) of hyper-parameters you can use this information to save time. With a few thousands of variables you will see the difference, which can be up to 50%. For the non robust correlation based tests, the difference may not be significant though, because the unconditional correlation coefficients are calculated very efficiently.

The max_k option: the maximum size of the conditioning set to use in the conditioning independence test. Larger values provide more accurate results, at the cost of higher computational times. When the sample size is small (e.g., < 50 observations) the max_k parameter should be ≤ 5 , otherwise the conditional independence test may not be able to provide reliable results.

If the dataset (predictor variables) contains missing (NA) values, they will automatically be replaced by the current variable (column) mean value with an appropriate warning to the user after the execution.

Conditional independence test functions to be passed through the user_test argument should have the same signature of the included test. See `testIndFisher` for an example.

For all the available conditional independence tests that are currently included on the package, please see `CondIndTests`.

If two or more p-values are below the machine epsilon (`.Machine$double.eps` which is equal to $2.220446e-16$), all of them are set to 0. To make the comparison or the ordering feasible we use the logarithm of the p-value. The max-min heuristic though, requires comparison and an ordering of the p-values. Hence, all conditional independence tests calculate the logarithm of the p-value.

If there are missing values in the dataset (predictor variables) columnwise imputation takes place. The median is used for the continuous variables and the mode for categorical variables. It is a naive and not so clever method. For this reason the user is encouraged to make sure his data contain no missing values.

If you have percentages, in the (0, 1) interval, they are automatically mapped into R by using the logit transformation.

If you use `testIndSpearman` (argument "test"), the ranks of the data calculated and those are used in the calculations. This speeds up the whole procedure.

Currently only the `testIndFisher` and `testIndSpearman` tests are supported for use in the algorithm.

If the argument **statistic** is set to FALSE, the p-values from the hypothesis test of each dataset are combined via Fisher's meta-analytic approach, that is $T = -2 \sum_{i=1}^k \log p_i$ and $T \chi_{2k}^2$. If **statistic** is TRUE, the test statistics are combined as $T = \frac{\sum_{i=1}^k t_i / se(t_i)}{\sum_{i=1}^k 1/se(t_i)}$ and $T N(0, 1)$.

Value

The output of the algorithm is an object of the class 'SESoutput' for SES or 'MMPCoutput' for MMPC including:

<code>selectedVars</code>	The selected variables, i.e., the signature of the target variable.
<code>selectedVarsOrder</code>	The order of the selected variables according to increasing pvalues.
<code>queues</code>	A list containing a list (queue) of equivalent features for each variable included in <code>selectedVars</code> . An equivalent signature can be built by selecting a single feature from each queue. Featured only in SES.
<code>signatures</code>	A matrix reporting all equivalent signatures (one signature for each row). Featured only in SES.
<code>hashObject</code>	The <code>hashObject</code> caching the statistic calculated in the current run.
<code>pvalues</code>	For each feature included in the dataset, this vector reports the strength of its association with the target in the context of all other variables. Particularly, this vector reports the max p-values (on a logarithmic scale) found when the association of each variable with the target is tested against different conditional sets. Lower values indicate higher association.
<code>stats</code>	The statistics corresponding to "pvalues" (higher values indicates higher association).
<code>univ</code>	This is a list with the univariate associations. The test statistics and their corresponding logged p-values, along with their flag (1 if the test was performed and 0 otherwise). This list is very important for subsequent runs of SES with different hyper-parameters. After running SES with some hyper-parameters you might want to run SES again with different hyper-parameters. To avoid calculating the univariate associations (first step of SES or MMPC) again, you can take this list from the first run of SES and plug it in the argument "ini" in the next run(s) of SES or MMPC. This can speed up the second run (and subsequent runs of course) by 50%. See the argument "univ" in the output values.
<code>max_k</code>	The <code>max_k</code> option used in the current run.
<code>threshold</code>	The <code>threshold</code> option used in the current run.
<code>runtime</code>	The run time of the algorithm. A numeric vector. The first element is the user time, the second element is the system time and the third element is the elapsed time.

`test` The character name of the statistic test used.

Generic Function implemented for SESoutput Object:

```
plot(object=SESoutput, mode="all")
```

Plots the generated pvalues (using barplot) of the current SESoutput object in comparison to the threshold.

Argument mode can be either "all" or "partial" for the first 500 pvalues of the object.

Author(s)

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Tsamardinos, Brown and Aliferis (2006). The max-min hill-climbing Bayesian network structure learning algorithm. *Machine learning*, 65(1), 31-78.

See Also

[SES](#), [CondIndTests](#), [cv.ses](#)

Examples

```
set.seed(123)
require(hash)

#simulate a dataset with continuous data
dataset <- matrix(runif(1000 * 100, 1, 100), ncol = 100)

#define a simulated class variable
target <- 3 * dataset[, 10] + 2 * dataset[, 20] + 3 * dataset[, 30] + rnorm(1000, 0, 5)

#define some simulated equivalences
dataset[, 15] <- dataset[, 10] + rnorm(1000, 0, 2)
dataset[, 10] <- dataset[, 10] + rnorm(1000, 0, 2)
dataset[, 25] <- dataset[, 20] + rnorm(1000, 0, 2)
dataset[, 23] <- dataset[, 20] + rnorm(1000, 0, 2)

require("hash", quietly = TRUE)
#run the SES algorithm
a1 <- SES(target, dataset, max_k = 5, threshold = 0.05, test = "testIndFisher",
```

```

hash = TRUE, hashObject = NULL)

ina <- rbinom(1000, 2, 0.5) + 1
a2 <- ma.ses(target , dataset, ina = ina, max_k = 5, threshold = 0.05, test = "testIndFisher",
hash = TRUE, hashObject = NULL)
a3 <- ma.mmpc(target , dataset, ina = ina, max_k = 5, threshold = 0.05, test = "testIndFisher",
hash = TRUE, hashObject = NULL)

#get the generated signatures
a1@signatures
a2@signatures
a3@selectedVars

```

Correlation based conditonal independence tests

Fisher and Spearman conditional independence test for continuous class variables

Description

The main task of this test is to provide a p-value PVALUE for the null hypothesis: feature 'X' is independent from 'TARGET' given a conditioning set CS.

Usage

```

testIndFisher(target, dataset, xIndex, csIndex, wei = NULL, statistic = FALSE,
univariateModels = NULL, hash = FALSE, stat_hash = NULL,
pvalue_hash = NULL)

testIndMMFisher(target, dataset, xIndex, csIndex, wei = NULL, statistic = FALSE,
univariateModels = NULL, hash = FALSE, stat_hash = NULL,
pvalue_hash = NULL)

testIndSpearman(target, dataset, xIndex, csIndex, wei = NULL, statistic = FALSE,
univariateModels = NULL, hash = FALSE, stat_hash = NULL,
pvalue_hash = NULL)

permFisher(target, dataset, xIndex, csIndex, wei = NULL, statistic = FALSE,
univariateModels = NULL, hash = FALSE, stat_hash = NULL,
pvalue_hash = NULL, threshold = 0.05, R = 999)

permMMFisher(target, dataset, xIndex, csIndex, wei = NULL, statistic = FALSE,
univariateModels = NULL, hash = FALSE, stat_hash = NULL,
pvalue_hash = NULL, threshold = 0.05, R = 999)

permDcor(target, dataset, xIndex, csIndex, wei = NULL, statistic = FALSE,
univariateModels = NULL, hash = FALSE, stat_hash = NULL,
pvalue_hash = NULL, threshold = 0.05, R = 499)

```

Arguments

target	A numeric vector containing the values of the target variable. If the values are proportions or percentages, i.e. strictly within 0 and 1 they are mapped into R using $\log(\text{target}/(1 - \text{target}))$. This can also be a list of vectors as well. In this case, the metanalytic approach is used.
dataset	A numeric matrix containing the variables for performing the test. Rows as samples and columns as features.
xIndex	The index of the variable whose association with the target we want to test.
csIndex	The indices of the variables to condition on. If you have no variables set this equal to 0.
wei	A vector of weights to be used for weighted regression. The default value is NULL. This is not used with "permDcor".
statistic	A boolean variable indicating whether the test statistics (TRUE) or the p-values should be combined (FALSE). See the details about this. For the permFisher test this is not taken into account.
univariateModels	Fast alternative to the hash object for univariate test. List with vectors "pvalues" (p-values), "stats" (statistics) and "flags" (flag = TRUE if the test was succesful) representing the univariate association of each variable with the target. Default value is NULL.
hash	A boolean variable which indicates whether (TRUE) or not (FALSE) to use the hash-based implementation of the statistics of SES. Default value is FALSE. If TRUE you have to specify the stat_hash argument and the pvalue_hash argument.
stat_hash	A hash object (hash package required) which contains the cached generated statistics of a SES run in the current dataset, using the current test.
pvalue_hash	A hash object (hash package required) which contains the cached generated p-values of a SES run in the current dataset, using the current test.
threshold	Threshold (suitable values in [0,1]) for assessing p-values significance. Default value is 0.05. This is actually obsolete here, but has to be in order tyo have a concise list of input arguments across the same family of functions.
R	The number of permutations to use. The default value is 999. For the "permDcor" this is set to 499.

Details

If hash = TRUE, testIndFisher requires the arguments 'stat_hash' and 'pvalue_hash' for the hash-based implementation of the statistic test. These hash Objects are produced or updated by each run of SES (if hash == TRUE) and they can be reused in order to speed up next runs of the current statistic test. If "SESoutput" is the output of a SES run, then these objects can be retrieved by SESoutput@hashObject\$stat_hash and the SESoutput@hashObject\$pvalue_hash.

Important: Use these arguments only with the same dataset that was used at initialization.

For all the available conditional independence tests that are currently included on the package, please see "?CondIndTests".

Note that if the `testIndReg` is used instead the results will not be the same, unless the sample size is very large. This is because the Fisher test uses the t distribution stemming from the Fisher's z transform and not the t distribution of the correlation coefficient.

BE CAREFUL with `testIndSpearman`. The Pearson's correlation coefficient is actually calculated. So, you must have transformed the data into their ranks before plugging them here. The reason for this is to speed up the computation time, as this test can be used in SES, MMPC and `mmhc.skel`. The variance of the Fisher transformed Spearman's correlation is $\frac{1.06}{n-3}$ and the variance of the Fisher transformed Pearson's correlation coefficient is $\frac{1}{n-3}$.

When performing the above tests with multiple datasets, the test statistic and the p-values are combined in a meta-analytic way. Is up to the user to decide whether to use the fixed effects model approach and combine the test statistics (statistic = TRUE), or combine the p-values as Fisher suggested (statistic = FALSE).

The argument `R` is useful only for the `permFisher` and `permDcor` tests. The `permDcor` test uses the distance correlation instead of the usual Pearson or Spearman correlations.

`TestIndMMFisher` does a robust estimation of the correlation via MM regression.

Value

A list including:

<code>pvalue</code>	A numeric value that represents the logarithm of the generated p-value due to Fisher's method (see reference below).
<code>stat</code>	A numeric value that represents the generated statistic due to Fisher's method (see reference below).
<code>stat_hash</code>	The current hash object used for the statistics. See argument <code>stat_hash</code> and details. If argument <code>hash = FALSE</code> this is NULL.
<code>pvalue_hash</code>	The current hash object used for the p-values. See argument <code>stat_hash</code> and details. If argument <code>hash = FALSE</code> this is NULL.

Author(s)

Vincenzo Lagani and Ioannis Tsamardinos

R implementation and documentation: Giorgos Athineou <athineou@csd.uoc.gr> Vincenzo Lagani <vlagani@csd.uoc.gr>.

References

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Lee Rodgers J., and Nicewander W.A. (1988). "Thirteen ways to look at the correlation coefficient." The American Statistician 42(1): 59–66.

Shevlyakov G. and Smirnov P. (2011). Robust Estimation of the Correlation Coefficient: An Attempt of Survey. Austrian Journal of Statistics, 40(1 & 2): 147–156.

Szekely G.J. and Rizzo, M.L. (2014). Partial distance correlation with methods for dissimilarities. The Annals of Statistics, 42(6): 2382–2412.

Szekely G.J. and Rizzo M.L. (2013). Energy statistics: A class of statistics based on distances. Journal of Statistical Planning and Inference 143(8): 1249–1272.

See Also

[testIndSpearman](#), [testIndReg](#), [SES](#), [testIndLogistic](#), [gSquare](#), [CondIndTests](#)

Examples

```
#simulate a dataset with continuous data
dataset <- matrix(runif(1000 * 200, 1, 1000), nrow = 1000 )
#the target feature is the last column of the dataset as a vector
target <- dataset[, 200]
res1 <- testIndFisher(target, dataset, xIndex = 44, csIndex = 100)
res2 <- testIndSpearman(target, dataset, xIndex = 44, csIndex = 100)
res3 <- permFisher(target, dataset, xIndex = 44, csIndex = 100, R = 999)
res4 <- permDcor(target, dataset, xIndex = 44, csIndex = 100, R = 99)

#define class variable (here the last column of the dataset)
dataset <- dataset[, -200];
#run the SES algorithm using the testIndFisher conditional independence test
sesObject <- SES(target, dataset, max_k = 3, threshold = 0.05, test = "testIndFisher");

#print summary of the SES output
summary(sesObject);
```

Cross-validation for ridge regression

Cross validation for the ridge regression

Description

Cross validation for the ridge regression is performed using the TT estimate of bias (Tibshirani and Tibshirani, 2009). There is an option for the GCV criterion which is automatic.

Usage

```
ridgereg.cv( target, dataset, K = 10, lambda = seq(0, 2, by = 0.1), auto = FALSE,
  seed = FALSE, ncores = 1, mat = NULL )
```

Arguments

target	A numeric vector containing the values of the target variable. If the values are proportions or percentages, i.e. strictly within 0 and 1 they are mapped into R using $\log(\text{target}/(1 - \text{target}))$.
dataset	A numeric matrix containing the variables. Rows are samples and columns are features.
K	The number of folds. Set to 10 by default.
lambda	A vector with the a grid of values of λ to be used.
auto	A boolean variable. If it is TRUE the GCV criterion will provide an automatic answer for the best λ . Otherwise k-fold cross validation is performed.
seed	A boolean variable. If it is TRUE the results will always be the same.
ncores	The number of cores to use. If it is more than 1 parallel computing is performed.
mat	If the user has its own matrix with the folds, he can put it here. It must be a matrix with K columns, each column is a fold and it contains the positions of the data, i.e. numbers, not the data. For example the first column is $c(1,10,4,25,30)$, the second is $c(21, 23,2, 19, 9)$ and so on.

Details

The `lm.ridge` command in MASS library is a wrapper for this function. If you want a fast choice of λ , then specify `auto = TRUE` and the λ which minimizes the generalised cross-validation criterion will be returned. Otherwise a k-fold cross validation is performed and the estimated performance is bias corrected as suggested by Tibshirani and Tibshirani (2009).

Value

A list including:

mspe	If <code>auto</code> is FALSE the values of the mean prediction error for each value of λ .
lambda	If <code>auto</code> is FALSE the λ which minimizes the MSPE.
performance	If <code>auto</code> is FALSE the minimum bias corrected MSPE along with the estimate of bias.
runtime	The run time of the algorithm. A numeric vector. The first element is the user time, the second element is the system time and the third element is the elapsed time.

Note

The values can be extracted with the `$` symbol, i.e. this is an S3 class output.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

Hoerl A.E. and R.W. Kennard (1970). Ridge regression: Biased estimation for nonorthogonal problems. *Technometrics*, 12(1):55-67.

Brown P. J. (1994). *Measurement, Regression and Calibration*. Oxford Science Publications.

Tibshirani R.J., and Tibshirani R. (2009). A bias correction for the minimum error rate in cross-validation. *The Annals of Applied Statistics* 3(2): 822-829.

See Also

[ridge.reg](#)

Examples

```
#simulate a dataset with continuous data
dataset <- matrix(runif(200 * 50, 1, 100), nrow = 200 )
#the target feature is the last column of the dataset as a vector
target <- dataset[, 50]
a1 <- ridgereg.cv(target, dataset, auto = TRUE)
a2 <- ridgereg.cv( target, dataset, K = 10, lambda = seq(0, 1, by = 0.1) )
```

Cross-Validation for SES and MMPC

Cross-Validation for SES and MMPC

Description

The function performs a k-fold cross-validation for identifying the best values for the SES and MMPC 'max_k' and 'threshold' hyper-parameters.

Usage

```
cv.ses(target, dataset, wei = NULL, kfolds = 10, folds = NULL,
        alphas = c(0.1, 0.05, 0.01), max_ks = c(3, 2), task = NULL,
        metric = NULL, modeler = NULL, ses_test = NULL, ncores = 1)

cv.mmpc(target, dataset, wei = NULL, kfolds = 10, folds = NULL,
        alphas = c(0.1, 0.05, 0.01), max_ks = c(3, 2), task = NULL,
        metric = NULL, modeler = NULL, mmpc_test = NULL, ncores = 1)

cv.waldses(target, dataset, wei = NULL, kfolds = 10, folds = NULL,
        alphas = c(0.1, 0.05, 0.01), max_ks = c(3, 2), task = NULL,
```

```
metric = NULL, modeler = NULL, ses_test = NULL, ncores = 1)

cv.waldmmmpc(target, dataset, wei = NULL, kfolds = 10, folds = NULL,
alphas = c(0.1, 0.05, 0.01), max_ks = c(3, 2), task = NULL,
metric = NULL, modeler = NULL, mmmpc_test = NULL, ncores = 1)

cv.permses(target, dataset, wei = NULL, kfolds = 10, folds = NULL,
alphas = c(0.1, 0.05, 0.01), max_ks = c(3, 2), task = NULL,
metric = NULL, modeler = NULL, ses_test = NULL, R = 999, ncores = 1)

cv.permmmmpc(target, dataset, wei = NULL, kfolds = 10, folds = NULL,
alphas = c(0.1, 0.05, 0.01), max_ks = c(3, 2), task = NULL,
metric = NULL, modeler = NULL, mmmpc_test = NULL, R = 999, ncores = 1)
```

Arguments

target	The target or class variable as in SES and MMPC. The difference is that it cannot accept a single numeric value, an integer indicating the column in the dataset.
dataset	The dataset object as in SES and MMPC.
wei	A vector of weights to be used for weighted regression. The default value is NULL.
kfolds	The number of the folds in the k-fold Cross Validation (integer).
folds	The folds of the data to use (a list generated by the function generateCVRRuns TunePareto). If NULL the folds are created internally with the same function.
alphas	A vector of SES or MMPC thresholds hyper parameters used in CV.
max_ks	A vector of SES or MMPC max_ks parameters used in CV.
task	A character ("C", "R" or "S"). It can be "C" for classification (logistic, multinomial or ordinal regression), "R" for regression (robust and non robust linear regression, median regression, (zero inflated) poisson and negative binomial regression, beta regression), "S" for survival regression (Cox, Weibull or exponential regression).
metric	A metric function provided by the user. If NULL the following functions will be used: auc.mxm, mse.mxm, ci.mxm for classification, regression and survival analysis tasks, respectively. See details for more. If you know what you have put it here to avoid the function choosing something else. Note that you put these words as they are, without "".
modeler	A modeling function provided by the user. If NULL the following functions will be used: glm.mxm, lm.mxm, coxph.mxm for classification, regression and survival analysis tasks, respectively. See details for more. If you know what you have put it here to avoid the function choosing something else. Note that you put these words as they are, without "".
ses_test	A function object that defines the conditional independence test used in the SES function (see also SES help page). If NULL, "testIndFisher", "testIndLogistic" and "censIndCR" are used for classification, regression and survival analysis tasks, respectively. If you know what you have put it here to avoid the function choosing something else. Not all tests can be included here. "testIndClogit",

	"testIndMVreg", "testIndIG", "testIndGamma", "testIndZIP" and "testIndTobit" are not available at the moment.
mmpc_test	A function object that defines the conditional independence test used in the MMPC function (see also SES help page). If NULL, "testIndFisher", "testIndLogistic" and "censIndCR" are used for classification, regression and survival analysis tasks, respectively.
R	The number of permutations, set to 999 by default. There is a trick to avoid doing all permutations. As soon as the number of times the permuted test statistic is more than the observed test statistic is more than 50 (if threshold = 0.05 and R = 999), the p-value has exceeded the significance level (threshold value) and hence the predictor variable is not significant. There is no need to continue do the extra permutations, as a decision has already been made.
ncores	This argument is valid only if you have a multi-threaded machine.

Details

Input for metric functions: predictions: A vector of predictions to be tested. test_target: target variable actual values to be compared with the predictions.

The output of a metric function is a single numeric value. **Higher values indicate better performance.** Metric based on error measures should be modified accordingly (e.g., multiplying the error for -1)

The metric functions that are currently supported are:

- auc.mxm: "area under the receiver operator characteristic curve" metric for binary logistic regression.
- acc.mxm: accuracy for binary logistic regression.
- fscore.mxm: F score for binary logistic regression.
- euclid_sens.spec.mxm: Euclidean norm of 1 - sensitivity and 1 - specificity for binary logistic regression.
- acc_multinom.mxm: accuracy for multinomial logistic regression.
- mse.mxm: $-1 * (\text{mean squared error})$, for robust and non robust linear regression and median (quantile) regression.
- ci.mxm: 1 - concordance index as provided in the rcorr.cens function from the surviva package. This is to be used with the Cox proportional hazards model only.
- ciwr.mxm concordance index as provided in the rcorr.cens function from the survival package. This is to be used with the Weibull regression model only.
- poisdev.mxm: Poisson regression deviance.
- nbdev.mxm: Negative binomial regression deviance.
- binomdev.mxm: Negative binomial regression deviance.
- ord_mae.mxm: Ordinal regression mean absolute error.

Usage: metric(predictions, test_target)

Input of modelling functions: train_target: target variable used in the training procedure. sign_data: training set. sign_test: test set.

Modelling functions provide a single vector of predictions obtained by applying the model fit on `sign_data` and `train_target` on the `sign_test`

The modelling functions that are currently supported are:

- `glm.mxm`: fits a glm for a binomial family (Classification task).
- `lm.mxm`: fits a linear model (stats) for the regression task.
- `coxph.mxm`: fits a cox proportional hazards regression model for the survival task.
- `weibreg.mxm`: fits a Weibull regression model for the survival task.
- `rq.mxm`: fits a quantile (median) regression model for the regression task.
- `lmrob.mxm`: fits a robust linear model for the regression task.
- `pois.mxm`: fits a poisson regression model for the regression task.
- `nb.mxm`: fits a negative binomial regression model for the regression task.
- `multinom.mxm`: fits a multinomial regression model for the regression task.
- `ordinal.mxm`: fits an ordinal regression model for the regression task.
- `beta.mxm`: fits a beta regression model for the regression task. The predicted values are transformed into R using the logit transformation. This is so that the "mse.mxm" metric function can be used. In addition, this way the performance can be compared with the regression scenario, where the logit is applied and then a regression model is employed.

Usage: `modeler(train_target, sign_data, sign_test)`

The procedure will be more automated in the future and more functions will be added. The multithreaded functions have been tested and no error has been detected. However, if you spot any suspicious results please let us know.

Value

A list including:

`cv_results_all` A list with predictions, performances and signatures for each fold and each SES or MMPC configuration (e.g `cv_results_all[[3]]$performances[1]` indicates the performance of the 1st fold with the 3d configuration of SES or MMPC). In the case of the multi-threaded functions (`cvses.par` and `cvmmmpc.par`) this is a list with a matrix. The rows correspond to the folds and the columns to the configurations (pairs of `threshold` and `max_k`).

`best_performance`

A numeric value that represents the best average performance.

`best_configuration`

A list that corresponds to the best configuration of SES or MMPC including `id`, `threshold` (named 'a') and `max_k`.

Bear in mind that the values can be extracted with the `$` symbol, i.e. this is an S3 class output.

Author(s)

R implementation and documentation: Giorgos Athineou <athineou@csd.uoc.gr> and Vincenzo Lagani <vlagani@csd.uoc.gr>

See Also

[SES](#), [CondIndTests](#), [testIndFisher](#), [testIndLogistic](#), [gSquare](#), [censIndCR](#)

Examples

```
set.seed(1234)

# simulate a dataset with continuous data
dataset <- matrix( rnorm(200 * 50), ncol = 50 )
# the target feature is the last column of the dataset as a vector
target <- dataset[, 50]
dataset <- dataset[, -50]

# get 50 percent of the dataset as a train set
train_set <- dataset[1:100, ]
train_target <- target[1:100]

require(hash)
# run a 10 fold CV for the regression task
best_model = cv.ses(target = train_target, dataset = train_set, kfolds = 5, task = "R")

# get the results
best_model$best_configuration
best_model$best_performance

# summary elements of the process. Press tab after each $ to view all the elements and
# choose the one you are interesting in.
# best_model$cv_results_all[[...]]$...
#i.e.
# mse value for the 1st configuration of SES of the 5 fold
abs(best_model$cv_results_all[[1]]$performances[5])

best_a <- best_model$best_configuration$a
best_max_k <- best_model$best_configuration$max_k
```

Data simulation from a DAG

Simulation of data from DAG (directed acyclic graph)

Description

The main task of this test is to provide a p-value PVALUE for the null hypothesis: feature 'X' is independent from 'TARGET' given a conditioning set CS. This test is based on the log likelihood ratio test.

Usage

```
rdag(n, p, s, a = 0, m, A = NULL, seed = FALSE)
rdag2(n, A = NULL, p, nei, low = 0.1, up = 1)
rmdag(n, A = NULL, p, nei, low = 0.1, up = 1)
```

Arguments

n	A number indicating the sample size.
p	A number indicating the number of nodes (or vertices, or variables).
nei	The average number of neighbours.
s	A number in (0, 1). This defines somehow the sparseness of the model. It is the probability that a node has an edge.
a	A number in (0, 1). The defines the percentage of outliers to be included in the simulated data. If $a = 0$, no outliers are generated.
m	A vector equal to the number of nodes. This is the mean vector of the normal distribution from which the data are to be generated. This is used only when $a > 0$ so as to define the mean vector of the multivariate normal from which the outliers will be generated.
A	If you already have an adjacency matrix in mind, plug it in here, otherwise, leave it NULL.
seed	If seed is TRUE, the simulated data will always be the same.
low	Every child will be a function of some parents. The beta coefficients of the parents will be drawn uniformly from two numbers, low and up. See details for more information on this.
up	Every child will be a function of some parents. The beta coefficients of the parents will be drawn uniformly from two numbers, low and up. See details for more information on this.

Details

In the case where no adjacency matrix is given, an $p \times p$ matrix with zeros everywhere is created. Every element below the diagonal is replaced by random values from a Bernoulli distribution with probability of success equal to s . This is the matrix B . Every value of 1 is replaced by a uniform value in $(0, 1)$. This final matrix is called A . The data are generated from a multivariate normal distribution with a zero mean vector and covariance matrix equal to $(\mathbf{I}_p - A)^{-1} (\mathbf{I}_p - A)$, where \mathbf{I}_p is the $p \times p$ identity matrix. If a is greater than zero, the outliers are generated from a multivariate normal with the same covariance matrix and mean vector the one specified by the user, the argument "m". The flexibility of the outliers is that you can specify outliers in some variables only or in all of them. For example, $m = c(0,0,5)$ introduces outliers in the third variable only, whereas $m = c(5,5,5)$ introduces outliers in all variables. The user is free to decide on the type of outliers to include in the data.

For the "rdag2", this is a different way of simulating data from DAGs. The first variable is normally generated. Every other variable can be a function of some previous ones. Suppose now that the i -th variable is a child of 4 previous variables. We need for coefficients b_j to multiply the 4 variables and then generate the i -th variable from a normal with mean $\sum_{j=1}^4 b_j X_j$ and variance 1. The b_j will be either positive or negative values with equal probability. Their absolute values ranges between "low" and "up". The code is accessible and you can see in detail what is going on. In addition, every generated data, are standardised to avoid numerical overflow.

The "rmdag" generates data from a BN with continuous, ordinal and binary data in proportions 50%, 25% and 25% respectively on average. This was used in the experiments run by Tsagris et al. (2017). If you want to generate data and then use them in the "pcalg" package with the function

"ci.fast2" or "ci.mm2" you should transform the resulting data into a matrix. The factor variables must become numeric starting from 0. See the examples for more on this.

Value

A list including:

nout	The number of outliers.
G	The adjacency matrix used. For the "rdag" if $G[i, j] = 2$, then $G[j, i] = 3$ and this means that there is an arrow from j to i . For the "rdag2" and "rmdag" the entries are either $G[i, j] = G[j, i] = 0$ (no edge) or $G[i, j] = 1$ and $G[j, i] = 0$ (indicating $i \rightarrow j$).
A	The matrix with the with the uniform values in the interval 0.1, 1. This is returned only by "rdag".
x	The simulated data.

Author(s)

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

Tsagris M., Borboudakis G., Lagani V. and Tsamardinos I. (2018). Constraint-based Causal Discovery with Mixed Data. *International Journal of Data Science and Analytics*.

<https://link.springer.com/content/pdf/10.1007>

<http://nugget.unisa.edu.au/CD2017/papersonly/constraint-based-causal-r1.pdf>

Spirtes P., Glymour C. and Scheines R. (2001). *Causation, Prediction, and Search*. The MIT Press, Cambridge, MA, USA, 3rd edition.

Colombo, Diego, and Marloes H. Maathuis (2014). Order-independent constraint-based causal structure learning. *The Journal of Machine Learning Research* 15(1): 3741–3782.

See Also

[pc.skel](#), [pc.or](#), [ci.mm](#), [mmhc.skel](#)

Examples

```
y <- rdag(100, 20, 0.2)
x <- y$x
tru <- y$G

mod <- pc.con(x)
b <- pc.or(mod)
plotnetwork(tru)
dev.new()
plotnetwork(b$G)

y <- rmdag(1000, p = 10, nei = 3)
x <- y$x
```

```
## make every factor numerical starting from 0.
for (i in 1:dim(x)[2]) {
  if ( is.factor(x[, i]) ) x[, i] <- as.numeric(x[, i]) - 1
}
```

Drop all possible single terms from a model using the partial correlation
Drop all possible single terms from a model using the partial correlation

Description

Drop all possible single terms from a model using the partial correlation.

Usage

```
cor.drop1(y, x, logged = FALSE)
```

Arguments

y	A numerical vector with the response variable.
x	A numerical matrix or a data.frame with the predictor variables. If it is a matrix it is internally transformed into a data.frame form, hence the user is advised to supply a data.frame in order to save some time. If the number of columns (variables) is higher than the number of rows (observations) the function will simply not work.
logged	If you want the p-values be returned leave this FALSE. If it is TRUE their logarithm is returned.

Details

This uses R's command `drop1` and modifies it so as to calculate the p-value using Fisher's conditional independence test.

Value

A matrix with two columns, the test statistic values and its associated p-value.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

See Also

[glm.bsreg](#), [fbed.reg](#), [mmpcbackphase](#)

Examples

```
y <- rnorm(1000)
x <- matrix( rnorm(1000 * 20), ncol = 20)
cor.drop1(y, x)
```

Effective sample size for G^2 test in BNs with case control data

Effective sample size for G^2 test in BNs with case control data

Description

Effective sample size for G^2 test in BNs with case control data.

Usage

```
Ness(propNt, N, K = 10000)
```

Arguments

propNt	A numerical vector with the proportions (distribution) of the (single) selection variable.
N	The sample size of the data.
K	The number of repetitions to be used for estimating the effective sample size.

Details

When dealing with case control data, spurious correlations or relationships arise. To deal with this one way is to adjust the sample size used in the G^2 test statistic. This function does exactly this, estimates the effective sample size as per the Borboudakis and Tsamardinos (2012) suggestion. The idea is that after learning the skeleton with the usual G^2 test, one should go to the edges and perform a conditional G^2

Value

The estimated effective sample size.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

Borboudakis G. and Tsamardinos I. (2015). Bayesian Network Learning with Discrete Case-Control Data. 31st Conference on Uncertainty in Artificial Intelligence (UAI), 151-160.

See Also

[SES](#), [MMPC](#), [testIndLogistic](#)

Examples

```
Ness(c(0.3, 0.7), N = 1000, K = 10000)
```

Estimation of the percentage of Null p-values

Estimation of the percentage of Null p-values

Description

Estimation of the percentage of Null p-values.

Usage

```
pi0est(p, lambda = seq(0.05, 0.95, by = 0.01), dof = 3)
```

Arguments

p	A vector of p-values.
lambda	A vector of values of the tuning parameter lambda.
dof	Number of degrees of freedom to use when estimating pi_0 with smoothing splines.

Details

The estimated proportion of null p-values is estimated the algorithm by Storey and Tibshirani (2003).

Value

The estimated proportion of non significant (null) p-values. In the paper Storey and Tibshirani mention that the estimate of pi0 is with lambda=1, but in their R code they use the highest value of lambda and thus we do the same here.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

Storey J.D. and Tibshirani R. (2003). Statistical significance for genome-wide experiments. Proceedings of the National Academy of Sciences, 100: 9440-9445.

See Also

[conf.edge.lower](#), [bn.skel.utils](#), [mmhc.skel](#), [pc.skel](#), [corfs.network](#), [local.mmhc.skel](#)

Examples

```
## simulate a dataset with continuous data
y <- rdag2(500, p = 25, nei = 3)
ind <- sample(1:25, 25)
x <- y$x[, ind]
mod <- pc.skel( x, method = "comb.fast", alpha = 0.01 )
pval <- exp(mod$pvalue)
pval <- lower.tri(pval)
pi0est(pval)
```

Feature selection using SES and MMPC for classification with longitudinal data
Feature selection using SES and MMPC for classification with longitudinal data

Description

SES algorithm follows a forward-backward filter approach for feature selection in order to provide minimal, highly-predictive, statistically-equivalent, multiple feature subsets of a high dimensional dataset. See also Details. MMPC algorithm follows the same approach without generating multiple feature subsets.

Usage

```
SES.timeclass(target, reps, id, dataset, max_k = 3, threshold = 0.05,
ini = NULL, wei = NULL, hash = FALSE, hashObject = NULL, ncores = 1)
```

```
MMPC.timeclass(target, reps, id, dataset, max_k = 3, threshold = 0.05,
ini = NULL, wei = NULL, hash = FALSE, hashObject = NULL, ncores = 1)
```

Arguments

target	The class variable. Provide a vector or a factor with discrete numbers indicating the class. Its length is equal to the number of rows of the dataset.
reps	A numeric vector containing the time points of the subjects. Its length is equal to the number of rows of the dataset.
id	A numeric vector containing the subjects or groups. Its length is equal to the number of rows of the dataset.
dataset	The dataset; provide a matrix. Currently, only continuous datasets are supported. The dataset contains longitudinal data, where each column is a variable. The repeated measurements are the samples.
max_k	The maximum conditioning set to use in the conditional independence test (see Details). Integer, default value is 3.

threshold	Threshold (suitable values in [0, 1]) for assessing p-values significance. Default value is 0.05.
ini	This is supposed to be a list. After running SES or MMPC with some hyper-parameters you might want to run SES again with different hyper-parameters. To avoid calculating the univariate associations (first step of SES and of MPPC) again, you can extract them from the first run of SES and plug them here. This can speed up the second run (and subsequent runs of course) by 50%. See the details and the argument "univ" in the output values.
wei	A vector of weights to be used for weighted regression. The default value is NULL.
hash	A boolean variable which indicates whether (TRUE) or not (FALSE) to store the statistics calculated during SES execution in a hash-type object. Default value is FALSE. If TRUE a hashObject is produced.
hashObject	A List with the hash objects generated in a previous run of SES.temporal. Each time SES runs with "hash=TRUE" it produces a list of hashObjects that can be re-used in order to speed up next runs of SES. Important: the generated hashObjects should be used only when the same dataset is re-analyzed, possibly with different values of max_k and threshold.
ncores	How many cores to use. This plays an important role if you have tens of thousands of variables or really large sample sizes and tens of thousands of variables and a regression based test which requires numerical optimisation. In other cases it will not make a difference in the overall time (in fact it can be slower). The parallel computation is used in the first step of the algorithm, where univariate associations are examined, those take place in parallel. We have seen a reduction in time of 50% with 4 cores in comparison to 1 core. Note also, that the amount of reduction is definitely not linear in the number of cores.

Details

This is SES and MMPC used in the static-longitudinal scenario of Tsagris, Lagani and Tsamardinos (2018). The idea is that you have many features of longitudinal data for many subjects. For each subject you have calculated the coefficients of a simple linear regression over time and this is repeated for each feature. In the end, assuming p features, you have p constants and p slopes for each subject, each constant and slope refers to a feature for a subject. Hence, each new feature consists of two vectors, the constants and the slopes and the feature selection takes place there.

Value

The output of the algorithm is an object of the class 'SESoutput' for SES or 'MMPCoutput' for MMPC including:

selectedVars	The selected variables, i.e., the signature of the target variable.
selectedVarsOrder	The order of the selected variables according to increasing pvalues.
queues	A list containing a list (queue) of equivalent features for each variable included in selectedVars. An equivalent signature can be built by selecting a single feature from each queue. Featured only in SES.

signatures	A matrix reporting all equivalent signatures (one signature for each row). Featured only in SES.
hashObject	The hashObject caching the statistic calculated in the current run.
pvalues	For each feature included in the dataset, this vector reports the strength of its association with the target in the context of all other variables. Particularly, this vector reports the max p-values found when the association of each variable with the target is tested against different conditional sets. Lower values indicate higher association.
stats	The statistics corresponding to "pvalues" (higher values indicates higher association).
univ	This is a list with the univariate associations. The test statistics and their corresponding logged p-values, along with their flag (1 if the test was performed and 0 otherwise). This list is very important for subsequent runs of SES with different hyper-parameters. After running SES with some hyper-parameters you might want to run SES again with different hyper-parameters. To avoid calculating the univariate associations (first step of SES or MMPC) again, you can take this list from the first run of SES and plug it in the argument "ini" in the next run(s) of SES or MMPC. This can speed up the second run (and subsequent runs of course) by 50%. See the argument "univ" in the output values.
max_k	The max_k option used in the current run.
threshold	The threshold option used in the current run.
n.tests	If you have set hash = TRUE, then the number of tests performed by SES or MMPC will be returned. If you have not set this to TRUE, the number of univariate associations will be returned. So be careful with this number.
runtime	The run time of the algorithm. A numeric vector. The first element is the user time, the second element is the system time and the third element is the elapsed time.
test	The character name of the statistic test used.

Generic Functions implemented for SESoutput Object:

`plot(object=SESoutput, mode="all")`

Plots the generated pvalues (using barplot) of the current SESoutput object in comparison to the threshold.

Argument mode can be either "all" or "partial" for the first 500 pvalues of the object.

Author(s)

Ioannis Tsamardinos, Vincenzo Lagani

R implementation and documentation: Giorgos Athineou <athineou@csd.uoc.gr> Vincenzo Lagani <vlagani@csd.uoc.gr>

References

Tsagris M., Lagani V., & Tsamardinos I. (2018). Feature selection for high-dimensional temporal data. *BMC bioinformatics*, 19(1), 17.

Vincenzo Lagani, George Kortas and Ioannis Tsamardinos (2013), Biomarker signature identification in "omics" with multiclass outcome. *Computational and Structural Biotechnology Journal*, 6(7):1-7.

McCullagh, Peter, and John A. Nelder. *Generalized linear models*. CRC press, USA, 2nd edition, 1989.

See Also

[mmpc.timeclass.model](#)

Examples

```
## assume these are longitudinal data, each column is a variable (or feature)
dataset <- matrix( rnorm(400 * 100), ncol = 100 )
id <- rep(1:80, each = 5) ## 80 subjects
reps <- rep( seq(4, 12, by = 2), 80) ## 5 time points for each subject
## dataset contains are the regression coefficients of each subject's values on the
## reps (which is assumed to be time in this example)
target <- rep(0:1, each = 200)
a <- MMPC.timeclass(target, reps, id, dataset)
```

Fit a mixture of beta distributions in p-values

Fit a mixture of beta distributions in p-values

Description

Fit a mixture of beta distributions in p-values.

Usage

```
pval.mixbeta(p)
```

Arguments

`p` A vector of p-values.

Details

The p-values are assumed to follow a mixture of two beta distributions. The null p-values follow $Be(1, 1)$ and the non-null p-values follow $Be(xi, 1)$. In the first step, the proportion of true null values using Storey and Tibshirani (2003) is calculated and then MLE is adopted to obtain xi . For more information on this see Triantafillou (2014).

Value

A vector with the estimated π_0 and ξ values.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

Triantafillou S., Tsamardinos I. and Roumpelaki A. (2014). Learning neighborhoods of high confidence in constraint-based causal discovery. In European Workshop on Probabilistic Graphical Models, pp. 487-502.

Storey J.D. and Tibshirani R. (2003). Statistical significance for genome-wide experiments. Proceedings of the National Academy of Sciences, 100: 9440-9445.

See Also

[pc.skel](#), [mmhc.skel](#), [corfs.network](#), [local.mmhc.skel](#), [conf.edge.lower](#)

Examples

```
## simulate a dataset with continuous data
y <- rdag2(500, p = 25, nei = 3)
ind <- sample(1:25, 25)
x <- y$x[, ind]
mod <- pc.skel( x, method = "comb.fast", alpha = 0.01 )
pval <- as.vector( mod$pvalue[lower.tri(mod$pvalue)] )
pval.mixbeta(pval)
```

Forward Backward Early Dropping selection regression

Forward Backward Early Dropping selection regression

Description

Forward Backward Early Dropping selection regression.

Usage

```
fbed.reg(target, dataset, ini = NULL, test = NULL, threshold = 0.05, wei = NULL,
K = 0, method = "LR", gam = NULL, backward = TRUE)
```

Arguments

target	The class variable. Provide either a string, an integer, a numeric value, a vector, a factor, an ordered factor or a Surv object.
dataset	The dataset; provide either a data frame or a matrix (columns = variables, rows = samples).
ini	If you already have the test statistics and the p-values of the univariate associations (the first step of FBED) supply them as a list with the names "stat" and "pvalue" respectively. If you have used the EBIC this list contains the eBIC of the univariate associations. Note, that the "gam" argument must be the same though.
test	The available tests: "testIndReg", "testIndPois", "testIndNB", "testIndLogistic", "testIndMMReg", "testIndBinom", "censIndCR", "censIndWR", "testIndBeta", "testIndZIP", "testIndGamma", "testIndNormLog", "testIndTobit", "testIndQPois", "testIndQBinom", "testIndFisher", "testIndMultinom", "testIndOrdinal", "testIndClogit". Note that not all of them work with eBIC.
threshold	Threshold (suitable values in [0,1]) for assessing p-values significance. Default value is 0.05.
wei	A vector of weights to be used for weighted regression. The default value is NULL. It is not suggested when robust is set to TRUE. If you want to use the "testIndBinom", then supply the successes in the y and the trials here.
K	How many times should the process be repeated? The default value is 0.
method	Do you want the likelihood ratio test to be performed ("LR" is the default value) or perform the selection using the "eBIC" criterion (BIC is a special case).
gam	In case the method is chosen to be "eBIC" one can also specify the <i>gamma</i> parameter. The default value is "NULL", so that the value is automatically calculated.
backward	After the Forward Early Dropping phase, the algorithm proceeds with the usual Backward Selection phase. The default value is set to TRUE. It is advised to perform this step as maybe some variables are false positives, they were wrongly selected. The backward phase using likelihood ratio test and eBIC are two different functions and can be called directly by the user. SO, if you want for example to perform a backward regression with a different threshold value, just use these two functions separately.

Details

The algorithm is a variation of the usual forward selection. At every step, the most significant variable enters the selected variables set. In addition, only the significant variables stay and are further examined. The non significant ones are dropped. This goes until no variable can enter the set. The user has the option to re-do this step 1 or more times (the argument K). In the end, a backward selection is performed to remove falsely selected variables. Note that you may have specified, for example, K=10, but the maximum value FBED used can be 4 for example.

The "testIndQPois" and "testIndQBinom" do not work for method = "eBIC" as there is no BIC associated with quasi Poisson and quasi binomial regression models.

If you specify a range of values of K it returns the results of `fbed.reg` for this range of values of K. For example, instead of running `fbed.reg` with `K=0`, `K=1`, `K=2` and so on, you can run `fbed.reg` with `K=0:2` and the selected variables found at `K=2`, `K=1` and `K=0` are returned. Note also, that you may specify maximum value of K to be 10, but the maximum value FBED used was 4 (for example).

Value

If K is a single number a list including:

<code>univ</code>	If you have used the log-likelihood ratio test this list contains the test statistics and the associated p-values of the univariate associations tests. If you have used the EBIC this list contains the eBIC of the univariate associations. Note, that the "gam" argument must be the same though.
<code>res</code>	A matrix with the selected variables, their test statistic and the associated logged p-value.
<code>info</code>	A matrix with the number of variables and the number of tests performed (or models fitted) at each round (value of K). This refers to the forward phase only.
<code>runtime</code>	The runtime required.
<code>back.rem</code>	The variables removed in the backward phase.
<code>back.n.tests</code>	The number of models fitted in the backward phase.

If K is a sequence of numbers a list including:

<code>univ</code>	If you have used the log-likelihood ratio test this list contains the test statistics and the associated p-values of the univariate associations tests. If you have used the EBIC this list contains the eBIC of the univariate associations.
<code>res</code>	The results of <code>fbed.reg</code> with the maximum value of K.
<code>mod</code>	A list where each element refers to a K value. If you run FBED with method = "LR" the selected variables, their test statistic and their p-value is returned. If you run it with method = "eBIC", the selected variables and the eBIC of the model with those variables are returned.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

Borboudakis G. and Tsamardinos I. (2017). Forward-Backward Selection with Early Dropping. <https://arxiv.org/pdf/1705.10770.pdf>

See Also

[fs.reg](#), [ebic.bsreg](#), [bic.fsreg](#), [MMPC](#)

Examples

```
#simulate a dataset with continuous data
dataset <- matrix( runif(100 * 50, 1, 100), ncol = 50 )
#define a simulated class variable
target <- rt(100, 10)

a1 <- fbbed.reg(target, dataset, test = "testIndReg")
a2 <- fbbed.reg(target, dataset, test = "testIndFisher")
y <- rpois(100, 10)
a2 <- fbbed.reg(y, dataset, test = "testIndPois")
a3 <- MMPC(y, dataset)
```

Forward Backward Early Dropping selection regression with GEE

Forward Backward Early Dropping selection regression with GEE

Description

Forward Backward Early Dropping selection regression with GEE.

Usage

```
fbbed.greg(target, dataset, id, reps = NULL, ini = NULL, threshold = 0.05,
wei = NULL, K = 0, test = "testIndGEEReg", correl = "exchangeable", se = "jack")
```

Arguments

target	The class variable. This can be a numerical vector with continuous data, binary, discrete or an ordered factor variable. It can also be a factor variable with two levels only. Pay attention to this. If you have ordinal data, then supply an ordered factor variable.
dataset	The set of candidate predictor variables provide a numerical matrix or a data.frame in case of categorical variables (columns = variables, rows = samples). In the case of ordinal regression, this can only be a numerical matrix, i.e. only continuous predictor variables.
id	This is a numerical vector of the same size as target denoting the groups or the subjects.
reps	This is a numerical vector of the same size as target denoting the groups or the subjects. If you have longitudinal data and you know the time points, then supply them here.
ini	If you already have the test statistics and the logged p-values of the univariate associations (the first step of FBED) supply them as a list with the names "stat" and "pvalue" respectively.
threshold	Threshold (suitable values in [0,1]) for asmmmsing p-values significance. Default value is 0.05.

<code>wei</code>	A vector of weights to be used for weighted regression. The default value is NULL. It is mentioned in the "geepack" that weights is not (yet) the weight as in sas proc genmod, and hence is not recommended to use.
<code>K</code>	How many times should the process be repeated? The default value is 0.
<code>test</code>	This is for the type of regression to be used, "testIndGEEReg", for Gaussian regression, "testIndGEEGamma" for Gamma regression, "testIndGEELogistic" for logistic regression, "testIndGEEOrdinal" for ordinal regression, "testIndGEENorm-Log" for linear regression with a log link, or "testIndGEEPois" for Poisson regression.
<code>correl</code>	The correlation structure. For the Gaussian, Logistic, Poisson and Gamma regression this can be either "exchangeable" (compound symmetry, suitable for clustered data) or "ar1" (AR(1) model, suitable for longitudinal data). For the ordinal logistic regression its only the "exchangeable" correlation structure.
<code>se</code>	<p>The method for estimating standard errors. This is very important and crucial. The available options for Gaussian, Logistic, Poisson and Gamma regression are: a) 'san.se', the usual robust estimate. b) 'jack': if approximate jackknife variance estimate should be computed. c) 'jls': if 1-step jackknife variance estimate should be computed and d) 'fij': logical indicating if fully iterated jackknife variance estimate should be computed. If you have many clusters (sets of repeated measurements) "san.se" is fine as it is asymptotically correct, plus jackknife estimates will take longer. If you have a few clusters, then maybe it's better to use jackknife estimates.</p> <p>The jackknife variance estimator was suggested by Paik (1988), which is quite suitable for cases when the number of subjects is small ($K < 30$), as in many biological studies. The simulation studies conducted by Ziegler et al. (2000) and Yan and Fine (2004) showed that the approximate jackknife estimates are in many cases in good agreement with the fully iterated ones.</p>

Details

The algorithm is a variation of the usual forward selection. At every step, the most significant variable enters the selected variables set. In addition, only the significant variables stay and are further examined. The non significant ones are dropped. This goes until no variable can enter the set. The user has the option to redo this step 1 or more times (the argument K). In the end, a backward selection is performed to remove falsely selected variables.

Since GEE are likelihood free, all significance tests take place using the Wald test, hence we decided not to have a backward phase. This algorithm is suitable for both clustered and longitudinal (temporal) data.

If you specify a range of values of K it returns the results of `fbed.reg` for this range of values of K. For example, instead of running `fbed.reg` with $K=0$, $K=1$, $K=2$ and so on, you can run `fbed.reg` with $K=0:2$ and the selected variables found at $K=2$, $K=1$ and $K=0$ are returned. Note also, that you may specify maximum value of K to be 10, but the maximum value FBED used was 4 (for example).

Value

If K is a single number a list including:

res	A matrix with the selected variables, their test statistic and the associated p-value.
info	A matrix with the number of variables and the number of tests performed (or models fitted) at each round (value of K).
runtime	The runtime required.

If K is a sequence of numbers a list including:

univ	If you have used the log-likelihood ratio test this list contains the test statistics and the associated logged p-values of the univariate associations tests. If you have used the EBIC this list contains the eBIC of the univariate associations.
res	The results of <code>fbed.gEE.reg</code> with the maximum value of K.
mod	A list where each element refers to a K value. If you run FBEd with method = "LR" the selected variables, their test statistic and their p-value is returned. If you run it with method = "eBIC", the selected variables and the eBIC of the model with those variables are returned.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

- Borboudakis G. and Tsamardinos I. (2017). Forward-Backward Selection with Early Dropping. <https://arxiv.org/pdf/1705.10770.pdf>
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- Ziegler A., Kastner C., Brunner D. and Blettner M. (2000). Familial associations of lipid profiles: A generalised estimating equations approach. *Statistics in medicine*, 19(24): 3345-3357
- Yan J. and Fine J. (2004). Estimating equations for association structures. *Statistics in medicine*, 23(6): 859-874
- Eugene Demidenko (2013). *Mixed Models: Theory and Applications with R*, 2nd Edition. New Jersey: Wiley & Sons.

See Also

[fbed.glm.reg](#), [glm.bsreg](#), [MMPC.temporal](#), [fbed.reg](#)

Examples

```
## Not run:
require(lme4)
data(sleepstudy)
attach(sleepstudy)
x <- matrix(rnorm(180 * 20), ncol = 20) ## unrelated predictor variables
m1 <- fbed.gee.reg(Reaction, x, Subject)
m2 <- fbed.glmm.reg(Reaction, x, Subject, backward = FALSE)

## End(Not run)
```

Forward Backward Early Dropping selection regression with GLMM

Forward Backward Early Dropping selection regression with GLMM

Description

Forward Backward Early Dropping selection regression with GLMM.

Usage

```
fbed.glmm.reg(target, dataset, id, reps = NULL, ini = NULL, threshold = 0.05,
wei = NULL, K = 0, method = "LR", gam = NULL, backward = TRUE, test = "testIndGLMMReg")
```

Arguments

target	The class variable. This can be a numerical vector with continuous data, binary or discrete valued data. It can also be a factor variable with two levels only.
dataset	The dataset; provide a numerical a matrix (columns = variables, rows = samples).
id	This is a numerical vector of the same size as target denoting the groups or the subjects.
reps	This is a numerical vector of the same size as target denoting the groups or the subjects. If you have longitudinal data and you know the time points, then supply them here.
ini	If you already have the test statistics and the p-values of the univariate associations (the first step of FBED) supply them as a list with the names "stat" and "pvalue" respectively in the case of method = "LR". In the case of method = "eBIC" this is a list with an element called "bic".
threshold	Threshold (suitable values in [0,1]) for asmmmsing p-values significance. Default value is 0.05.
wei	A vector of weights to be used for weighted regression. The default value is NULL.
K	How many times should the process be repated? The default value is 0.

method	Do you want the likelihood ratio test to be performed ("LR" is the default value) or perform the selection using the "eBIC" criterion (BIC is a special case).
gam	In case the method is chosen to be "eBIC" one can also specify the <i>gamma</i> parameter. The default value is "NULL", so that the value is automatically calculated.
backward	After the Forward Early Dropping phase, the algorithm proceeds with the usual Backward Selection phase. The default value is set to TRUE. It is advised to perform this step as maybe some variables are false positives, they were wrongly selected. The backward phase using likelihood ratio test and eBIC are two different functions and can be called directly by the user. SO, if you want for example to perform a backward regression with a different threshold value, just use these two functions separately.
test	This is for the type of regression to be used, "testIndGLMMReg", for Gaussian regression, "testIndGLMMLogistic" for logistic regression or "testIndGLMM-Pois" for Poisson regression.

Details

The algorithm is a variation of the usual forward selection. At every step, the most significant variable enters the selected variables set. In addition, only the significant variables stay and are further examined. The non significant ones are dropped. This goes until no variable can enter the set. The user has the option to redo this step 1 or more times (the argument K). In the end, a backward selection is performed to remove falsely selected variables.

Bear in mind that for the "gaussian" case, the forward phase takes place using the F test (Wald statistic calibrated against an F distribution). The backward phase though takes place using the log-likelihood ratio test.

If you specify a range of values of K it returns the results of `fbed.reg` for this range of values of K. For example, instead of running `fbed.reg` with `K=0`, `K=1`, `K=2` and so on, you can run `fbed.reg` with `K=0:2` and the selected variables found at `K=2`, `K=1` and `K=0` are returned. Note also, that you may specify maximum value of K to be 10, but the maximum value FBED used was 4 (for example).

Value

If K is a single number a list including:

res	A matrix with the selected variables, their test statistic and the associated logged p-value.
info	A matrix with the number of variables and the number of tests performed (or models fitted) at each round (value of K). This refers to the forward phase only.
runtime	The runtime required.
back.rem	The variables removed in the backward phase.
back.n. tests	The number of models fitted in the backward phase.

If K is a sequence of numbers a list including:

univ	If you have used the log-likelihood ratio test this list contains the test statistics and the associated p-values of the univariate associations tests. If you have used the EBIC this list contains the eBIC of the univariate associations.
res	The results of <code>fbed.glm.reg</code> with the maximum value of K.
mod	A list where each element refers to a K value. If you run FBED with method = "LR" the selected variables, their test statistic and their p-value is returned. If you run it with method = "eBIC", the selected variables and the eBIC of the model with those variables are returned.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

Borboudakis G. and Tsamardinos I. (2017). Forward-Backward Selection with Early Dropping. <https://arxiv.org/pdf/1705.10770.pdf>

Eugene Demidenko (2013). Mixed Models: Theory and Applications with R, 2nd Edition. New Jersey: Wiley & Sons.

See Also

[fbed.gee.reg](#), [glm.bsreg](#), [MMPC.temporal](#), [fbed.reg](#)

Examples

```
## Not run:
require(lme4)
data(sleepstudy)
attach(sleepstudy)
x <- matrix(rnorm(180 * 20), ncol = 20) ## unrelated predictor variables
m1 <- fbed.glm.reg(Reaction, x, Subject)
m2 <- MMPC.temporal(target = Reaction, group = Subject, dataset = x)

## End(Not run)
```

Forward selection regression

Variable selection in regression models with forward selection

Description

Variable selection in regression models with forward selection

Usage

```
fs.reg(target, dataset, ini = NULL, threshold = 0.05, wei = NULL, test = NULL,
user_test = NULL, stopping = "BIC", tol = 2, ncores = 1)
```

Arguments

target	The class variable. Provide either a string, an integer, a numeric value, a vector, a factor, an ordered factor or a Surv object. See also Details.
dataset	The dataset; provide either a data frame or a matrix (columns = variables, rows = samples). In either case, only two cases are available, either all data are continuous, or categorical.
ini	If you have a set of variables already start with this one. Otherwise leave it NULL.
threshold	Threshold (suitable values in [0,1]) for asmmmsing p-values significance. Default value is 0.05.
wei	A vector of weights to be used for weighted regression. The default value is NULL.
test	The regression model to use. Available options are most of the tests for SES and MMPC. The ones NOT available are "gSquare", "censIndER", "testIndMVreg", "testIndClogit" and "testIndSpearman". If you have continuous predictor variables in matrix form, you can put "testIndFisher" and this is pretty fast. Instead of calculating partial F-tests, which requires linear regression models to be fit, it calculates partial correlation coefficients and this is much more efficient.
user_test	A user-defined conditional independence test (provide a closure type object). Default value is NULL. If this is defined, the "test" argument is ignored.
stopping	The stopping rule. The BIC is always used for all methods. If you have linear regression though you can change this to "adjrsq" and in this case the adjusted R squared is used.
tol	The difference between two successive values of the stopping rule. By default this is set to 2. If for example, the BIC difference between two successive models is less than 2, the process stops and the last variable, even though significant does not enter the model.
ncores	How many cores to use. This plays an important role if you have tens of thousands of variables or really large sample sizes and tens of thousands of variables and a regression based test which requires numerical optimisation. In other cases it will not make a difference in the overall time (in fact it can be slower). The parallel computation is used in the first step of the algorithm, where univariate associations are examined, those take place in parallel. We have seen a reduction in time of 50% with 4 cores in comparison to 1 core. Note also, that the amount of reduction is not linear in the number of cores.

Details

If the current 'test' argument is defined as NULL or "auto" and the user_test argument is NULL then the algorithm automatically selects the best test based on the type of the data. Particularly:

- if target is a factor, the multinomial or the binary logistic regression is used. If the target has two values only, binary logistic regression will be used.
- if target is an ordered factor, the ordered logit regression is used. Hence, if you want to use multinomial or ordinal logistic regression, make sure your target is factor.
- if target is a numerical vector and the dataset is not a matrix, but a data.frame linear regression is used. If however, the dataset is a matrix, the correlation based forward selection is used. That is, instead of partial F-tests, we do partial correlation tests.
- if target is discrete numerical (counts), the poisson regression conditional independence test is used. If there are only two values, the binary logistic regression is to be used.
- if target is a Surv object, the Survival conditional independence test is used.

Value

In the case of `test="testIndMMReg"` and `class(dataset) = matrix`, just one matrix is returned with the index of the selected variable(s), the p-value, the test statistic and the BIC value of each model. For all other cases, the output of the algorithm is S3 object including:

<code>runtime</code>	The run time of the algorithm. A numeric vector. The first element is the user time, the second element is the system time and the third element is the elapsed time.
<code>mat</code>	A matrix with the variables and their latest test statistics and p-values.
<code>info</code>	A matrix with the selected variables, their logged p-values and test statistics. Each row corresponds to a model which contains the variables up to that line. The BIC in the last column is the BIC of that model.
<code>ci_test</code>	The conditional independence test used.
<code>final</code>	The final regression model.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

See Also

[glm.fsreg](#), [lm.fsreg](#), [bic.fsreg](#), [bic.glm.fsreg](#), [CondIndTests](#), [MMPC](#), [SES](#)

Examples

```
set.seed(123)

#simulate a dataset with continuous data
dataset <- matrix( runif(1000 * 20, 1, 100), ncol = 20 )

#define a simulated class variable
target <- rt(1000, 10)

a0 <- fs.reg(target, dataset, threshold = 0.05, stopping = "BIC", tol = 2)
```

```

a1 <- fs.reg(target, dataset, threshold = 0.05, test = "testIndRQ", stopping = "BIC",
tol = 2)

y <- survival::Surv(rexp(1000), rep(1,1000) )
a2 <- fs.reg(y, dataset, threshold = 0.05, test = "censIndWR", stopping = "BIC", tol = 2)
a3 <- MMPC(target, dataset)

target <- factor( rbinom(1000, 1, 0.6) )
b2 <- fs.reg(target, dataset, threshold = 0.05, test = NULL, stopping = "BIC", tol = 2)

```

Forward selection with generalised linear regression models

Variable selection in generalised linear regression models with forward selection

Description

Variable selection in generalised linear regression models with forward selection

Usage

```

glm.fsreg(target, dataset, ini = NULL, threshold = 0.05, wei = NULL, tol = 2,
ncores = 1)

gammafsreg(target, dataset, ini = NULL, threshold = 0.05, wei = NULL, tol = 2,
ncores = 1)

normlog.fsreg(target, dataset, ini = NULL, threshold = 0.05, wei = NULL, tol = 2,
ncores = 1)

```

Arguments

target	A numerical vector or a factor variable with two levels. The Gamma regression requires strictly positive numbers, whereas the normlog requires positive numbers, zero included.
dataset	The dataset; provide either a data frame or a matrix (columns = variables, rows = samples). In either case, only two cases are available, either all data are continuous, or categorical.
ini	If you have a set of variables already start with this one. Otherwise leave it NULL.
threshold	Threshold (suitable values in [0,1]) for assessing the p-values significance. Default value is 0.05.
wei	A vector of weights to be used for weighted regression. The default value is NULL. It is not suggested when robust is set to TRUE.
tol	The difference between two successive values of the stopping rule. By default this is set to 2. If for example, the BIC difference between two successive models is less than 2, the process stops and the last variable, even though significant does not enter the model.

ncores How many cores to use. This plays an important role if you have tens of thousands of variables or really large sample sizes and tens of thousands of variables and a regression based test which requires numerical optimisation. In other cases it will not make a difference in the overall time (in fact it can be slower). The parallel computation is used in the first step of the algorithm, where univariate associations are examined, those take place in parallel. We have seen a reduction in time of 50% with 4 cores in comparison to 1 core. Note also, that the amount of reduction is not linear in the number of cores.

Value

The output of the algorithm is S3 object including:

runtime	The run time of the algorithm. A numeric vector. The first element is the user time, the second element is the system time and the third element is the elapsed time.
mat	A matrix with the variables and their latest test statistics and p-values. If you have logistic or Poisson regression with continuous predictor variables in a matrix form and no weights, this will not appear. In this case, a C++ code is called and the output is less.
info	A matrix with the selected variables, their logged p-values and test statistics. Each row corresponds to a model which contains the variables up to that line. The BIC in the last column is the BIC of that model.
ci_test	The conditional independence test used.
final	The final regression model.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

See Also

[fs.reg](#), [lm.fsreg](#), [bic.fsreg](#), [bic.glm.fsreg](#), [CondIndTests](#), [MMPC](#), [SES](#)

Examples

```
set.seed(123)

#simulate a dataset with continuous data
dataset <- matrix( runif(1000 * 50, 1, 100), ncol = 50 )

#define a simulated class variable
target <- rpois(1000, 10)

a <- glm.fsreg(target, dataset, threshold = 0.05, tol = 2, ncores = 1 )
b <- MMPC(target, dataset, max_k = 3, threshold = 0.05, test = "testIndPois")
```

 Forward selection with linear regression models

Variable selection in linear regression models with forward selection

Description

Variable selection in linear regression models with forward selection

Usage

```
lm.fsreg(target, dataset, ini = NULL, threshold = 0.05, wei = NULL, stopping = "BIC",
  tol = 2, ncores = 1)
```

Arguments

target	The class variable. Provide either a string, an integer, a numeric value, a vector, a factor, an ordered factor or a Surv object. See also Details.
dataset	The dataset; provide either a data frame or a matrix (columns = variables, rows = samples). In either case, only two cases are available, either all data are continuous, or categorical.
ini	If you have a set of variables already start with this one. Currently this can only be a matrix with continuous variables. In such cases, the dataset must also contain continuous variables only. Otherwise leave it NULL.
threshold	Threshold (suitable values in [0,1]) for assessing the p-values significance. Default value is 0.05.
wei	A vector of weights to be used for weighted regression. The default value is NULL. It is not suggested when robust is set to TRUE.
stopping	The stopping rule. The BIC ("BIC") or the adjusted R^2 ("adjrsq") can be used.
tol	The difference between two successive values of the stopping rule. By default this is set to 2. If for example, the BIC difference between two successive models is less than 2, the process stops and the last variable, even though significant does not enter the model. If the adjusted R^2 is used, the tol should be something like 0.01 or 0.02.
ncores	How many cores to use. This plays an important role if you have tens of thousands of variables or really large sample sizes and tens of thousands of variables and a regression based test which requires numerical optimisation. In other cases it will not make a difference in the overall time (in fact it can be slower). The parallel computation is used in the first step of the algorithm, where univariate associations are examined, those take place in parallel. We have seen a reduction in time of 50% with 4 cores in comparison to 1 core. Note also, that the amount of reduction is not linear in the number of cores.

Details

Only linear regression (robust and non robust) is supported from this function.

Value

The output of the algorithm is S3 object including:

runtime	The run time of the algorithm. A numeric vector. The first element is the user time, the second element is the system time and the third element is the elapsed time.
mat	A matrix with the variables and their latest test statistics and p-values.
info	A matrix with the selected variables, their logged p-values and test statistics. Each row corresponds to a model which contains the variables up to that line. The BIC in the last column is the BIC of that model.
models	The regression models, one at each step.
ci_test	The conditional independence test used, "testIndReg".
final	The final regression model.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

See Also

[fs.reg](#), [lm.fsreg](#), [bic.fsreg](#), [bic.glm.fsreg](#), [CondIndTests](#), [MMPC](#), [SES](#)

Examples

```
set.seed(123)

#simulate a dataset with continuous data
dataset <- matrix( runif(1000 * 30, 1, 100), ncol = 30 )

#define a simulated class variable
target <- 3 * dataset[, 10] + 2 * dataset[, 20] + rnorm(1000, 0, 5)
a1 <- lm.fsreg(target, dataset, threshold = 0.05, stopping = "BIC", tol = 2)
```

G-square conditional independence test for discrete data

G-square conditional independence test for discrete data

Description

The main task of this test is to provide a p-value PVALUE for the null hypothesis: feature 'X' is independent from 'TARGET' given a conditioning set CS. This test is based on the log likelihood ratio test.

Usage

```
gSquare(target, dataset, xIndex, csIndex, wei = NULL,
        univariateModels = NULL, hash = FALSE, stat_hash = NULL,
        pvalue_hash = NULL)
```

```
permgSquare(target, dataset, xIndex, csIndex, wei = NULL,
            univariateModels = NULL, hash = FALSE, stat_hash = NULL,
            pvalue_hash = NULL, threshold = 0.05, R = 999)
```

Arguments

target	A numeric vector containing the values of the target variable.
dataset	A numeric matrix containing the variables for performing the test. Rows as samples and columns as features.
xIndex	The index of the variable whose association with the target we want to test.
csIndex	The indices of the variables to condition on.
wei	This argument is not used in this test.
univariateModels	Fast alternative to the hash object for univariate tests. List with vectors "pvalues" (p-values), "stats" (statistics) and "flags" (flag = TRUE if the test was succesful) representing the univariate association of each variable with the target. Default value is NULL.
hash	A boolean variable which indicates whether (TRUE) or not (FALSE) to use the hash-based implementation of the statistics of SES. Default value is FALSE. If TRUE you have to specify the stat_hash argument and the pvalue_hash argument.
stat_hash	A hash object (hash package required) which contains the cached generated statistics of a SES run in the current dataset, using the current test.
pvalue_hash	A hash object (hash package required) which contains the cached generated p-values of a SES run in the current dataset, using the current test.
threshold	Threshold (suitable values in [0,1]) for assessing p-values significance. Default value is 0.05. This is actually obsolete here, but has to be in order to have a concise list of input arguments across the same family of functions.
R	The number of permutations to use. The default value is 999.

Details

If the number of samples is at least 5 times the number of the parameters to be estimated, the test is performed, otherwise, independence is not rejected (see Tsmardinis et al., 2006, pg. 43)

If hash = TRUE, testIndLogistic requires the arguments 'stat_hash' and 'pvalue_hash' for the hash-based implementation of the statistical test. These hash Objects are produced or updated by each run of SES (if hash == TRUE) and they can be reused in order to speed up next runs of the current statistic test. If "SESoutput" is the output of a SES run, then these objects can be retrieved by SESoutput@hashObject\$stat_hash and the SESoutput@hashObject\$pvalue_hash.

Important: Use these arguments only with the same dataset that was used at initialization.

For all the available conditional independence tests that are currently included on the package, please see "`?CondIndTests`".

Value

A list including:

<code>pvalue</code>	A numeric value that represents the logarithm of the generated p-value of the G^2 test (see reference below).
<code>stat</code>	A numeric value that represents the generated statistic of the G^2 test (see reference below).
<code>stat_hash</code>	The current hash object used for the statistics. See argument <code>stat_hash</code> and details. If argument <code>hash = FALSE</code> this is NULL.
<code>pvalue_hash</code>	The current hash object used for the p-values. See argument <code>stat_hash</code> and details. If argument <code>hash = FALSE</code> this is NULL.

Author(s)

R implementation and documentation: Giorgos Athineou <athineou@csd.uoc.gr>

References

Tsamardinos, Ioannis, Laura E. Brown, and Constantin F. Aliferis. The max-min hill-climbing Bayesian network structure learning algorithm. *Machine learning*, 2006 65(1): 31–78.

See Also

[SES](#), [testIndFisher](#), [testIndLogistic](#), [censIndCR](#), [CondIndTests](#)

Examples

```
#simulate a dataset with binary data
dataset <- matrix(rbinom(500 * 51, 1, 0.6), ncol = 51)
#initialize binary target
target <- dataset[, 51]
#remove target from the dataset
dataset <- dataset[, -51]

#run the gSquare conditional independence test for the binary class variable
results <- gSquare(target, dataset, xIndex = 44, csIndex = c(10,20) )
results

#run SES algorithm using the gSquare conditional independence test for the binary class variable
sesObject <- SES(target, dataset, max_k = 3, threshold = 0.05, test = "gSquare");
target <- as.factor(target)
sesObject2 <- SES(target, dataset, max_k = 3, threshold = 0.05, test = "testIndLogistic");
#print summary of the SES output
summary(sesObject);
summary(sesObject2);
```

Generalised linear mixed models based on temporal SES and MMPC outputs
*Generalised linear mixed model(s) based obtained from temporal SES
or MMPC*

Description

One or more regression models obtained from SES or MMPC, are returned.

Usage

```
mmpc.temporal.model(target, dataset, reps = NULL, group, slopes = FALSE, wei = NULL,
mmpctemporal.Object, test = NULL)
```

```
ses.temporal.model(target, dataset, reps = NULL, group, slopes = FALSE, wei = NULL,
sestemporal.Object, nsigmat = 1, test = NULL)
```

```
mmpc.gee.model(target, dataset, reps = NULL, group, correl = "exchangeable",
se = "jack", wei = NULL, mmpcgee.Object, test = NULL)
```

```
ses.gee.model(target, dataset, reps = NULL, group, correl = "exchangeable",
se = "jack", wei = NULL, sesgee.Object, nsigmat = 1, test = NULL)
```

Arguments

target	The class variable. Provide a vector with continuous (normal), binary (binomial) or discrete (Poisson) data.
dataset	The dataset; provide either a data frame or a matrix (columns = variables, rows = observations). Currently, only continuous datasets are supported.
reps	A numeric vector containing the time points of the subjects. Its length is equal to the length of the target variable. If you have clustered data, leave this NULL.
group	A numeric vector containing the subjects or groups. It must be of the same length as target.
slopes	Should random slopes for the ime effect be fitted as well? Default value is FALSE.
correl	The correlation structure. For the Gaussian, Logistic, Poisson and Gamma regression this can be either "exchangeable" (compound symmetry, suitable for clustered data) or "ar1" (AR(1) model, suitable for longitudinal data). For the ordinal logistic regression its only the "exchangeable" correlation sturcture. This is obsolete for "testIndGEEOrdinal", but is here for compatibility reasons.
se	The method for estimating standard errors. This is very important and crucial. The available options for Gaussian, Logistic, Poisson and Gamma regression are: a) 'san.se', the usual robust estimate. b) 'jack': if approximate jackknife variance estimate should be computed. c) 'j1s': if 1-step jackknife variance estimate should be computed and d) 'fij': logical indicating if fully iterated jackknife variance estimate should be computed. If you have many clusters (sets of

repeated measurements) "san.se" is fine as it is asymptotically correct, plus jackknife estimates will take longer. If you have a few clusters, then maybe it's better to use jackknife estimates.

The jackknife variance estimator was suggested by Paik (1988), which is quite suitable for cases when the number of subjects is small ($K < 30$), as in many biological studies. The simulation studies conducted by Ziegler et al. (2000) and Yan and Fine (2004) showed that the approximate jackknife estimates are in many cases in good agreement with the fully iterated ones. This is obsolete for "testIndGEEOrdinal", but is here for compatibility reasons.

<code>wei</code>	A vector of weights to be used for weighted regression. The default value is NULL.
<code>mmpctemporal.Object</code>	An object with the results of an MMPC.temporal run.
<code>sestemporal.Object</code>	An object with the results of a SES.temporal run.
<code>mmpcgee.Object</code>	An object with the results of an MMPC.gee run.
<code>sesgee.Object</code>	An object with the results of an SES.gee run.
<code>nsignat</code>	How many signatures to use. If <code>nsignat = 1</code> (default value) the first set of variables will be used for the model. If you want more, then specify the number of signatures you want. If you want the models based on all signatures, specify "all". If you put a number which is higher than the number of signatures, all models will be returned.
<code>test</code>	The conditional independence test to use. Default value is NULL. Currently, the only available conditional independence tests are the testIndGLMMReg , testIndGLMMLogistic , testIndGLMMPois and testIndLMM which fit linear mixed models.

Details

This command is useful if you want to see all models and check for example their fitting ability, MSE in linear models for example.

Value

A list including:

<code>mod</code>	Depending on the number of signatures requested, one or models will be returned.
<code>signature</code>	A matrix (or just one vector if one signature only) with the variables of each signature, along with the BIC of the corresponding regression model.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

- Tsagris, M., Lagani, V., & Tsamardinos, I. (2018). Feature selection for high-dimensional temporal data. *BMC bioinformatics*, 19(1), 17.
- I. Tsamardinos, M. Tsagris and V. Lagani (2015). Feature selection for longitudinal data. Proceedings of the 10th conference of the Hellenic Society for Computational Biology & Bioinformatics (HSCBB15)
- Eugene Demidenko (2013). *Mixed Models: Theory and Applications with R*, 2nd Edition. New Jersey: Wiley & Sons.
- Pinheiro J. and D. Bates. *Mixed-effects models in S and S-PLUS*. Springer Science & Business Media, 2006.

See Also

[SES](#), [MMPC](#), [cv.ses](#), [cv.mmpc](#)

Examples

```
## Not run:
require(lme4)
data(sleepstudy)
attach(sleepstudy)
x <- matrix(rnorm(180 * 100),ncol = 100) ## unrelated predictor variables
m1 <- SES.temporal(Reaction, Days, Subject, x)
m2 <- MMPC.temporal(Reaction, Days, Subject, x)
mod <- mmpc.temporal.model(target, dataset = x, reps = Days, group = Subject, slopes = FALSE,
mmpctemporal.Object = m2)

## End(Not run)
```

Generalised ordinal regression
Generalised ordinal regression

Description

Generalised ordinal regression.

Usage

```
ordinal.reg(formula, data)
```

Arguments

formula	An object of class "formula" (or one that can be coerced to that class): a symbolic description of the model to be fitted. This is the usual formula use by many regression models in R and other packages. Type "glm" or "formula" in R for more information.
data	A data.frame object carrying the relevant data.

Details

Generalised ordinal regression is fitted. This means the instead of having the same coefficient for each predictor variable, they are allowed to vary. The usual, proportional odds, ordinal regression specifies that the lines do not cross. This one does not need the proportional odds assumption. The proportional odds assumption does not always hold in practice and is a rather restrictive model. Be careful though, you may end up with negative probabilities. We do a tick to fix them, but in that case, you may have not found the optimal model. This is a problematic case unfortunately. Williams (2006) explains in a very nice way how one can fit this model by using many logistic regressions in an incremental way. The number of logistic regression models to be fit is the number of categories of the response variables - 1.

Value

A list including:

<code>mess</code>	If you hit negative probabilities, the message "problematic region" will be printed. Otherwise, this is NULL.
<code>be</code>	The regression coefficients.
<code>devi</code>	The deviance of the model.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

Williams, R. (2006). Generalized ordered logit/partial proportional odds models for ordinal dependent variables. *Stata Journal*, 6(1), 58-82

See Also

[pc.skel](#), [ridge.reg](#)

Examples

```
y <- factor( rbinom(100, 3, 0.5) )
x <- matrix( rnorm(100*3), ncol = 3)
ordinal.reg(y ~ x, data = data.frame(x) )
ordinal.reg(y ~ 1, data = data.frame(x) )
```

Generalised orthogonal matching pursuit (gOMP)
Generalised orthogonal matching pursuit (gOMP)

Description

Generalised orthogonal matching pursuit.

Usage

```
gomp(target, dataset, tol = qchisq(0.95, 1) + log(dim(dataset)[1]),
      test = "testIndFisher" )
```

Arguments

target	The response variable, a numeric vector, a matrix or a Surv object.
dataset	A matrix with continuous data, where the rows denote the samples and the columns are the variables.
tol	The tolerance value to terminate the algorithm. This is the change in the criterion value between two successive steps. The default value is the 95% quantile of the χ^2 distribution with 1 degree of freedom plus the logarithm of the sample size. For test = "testIndFisher" the BIC is already calculated, so better make it qchisq(0.95, 1).
test	This denotes the parametric model to be used each time. It depends upon the nature of the target variable. The possible values are "testIndFisher" (or "testIndReg" for the same purpose), "testIndLogistic", "testIndPois", "testIndQPois", "testIndQbinom", "testIndNormLog", "testIndMVreg", "testIndNB", "testIndBeta", "testIndGamma", "testIndMMReg", "testIndRQ", "testIndTobit", "censIndCR" and "censIndWR".

Value

A list including:

runtime	The runtime of the algorithm
res	A matrix with two columns. The selected variable(s) and the criterion value at every step.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@yahoo.gr>.

References

Pati Y. C., Rezaifar R. & Krishnaprasad P. S. (1993). Orthogonal matching pursuit: Recursive function approximation with applications to wavelet decomposition. In Signals, Systems and Computers. 1993 Conference Record of The Twenty-Seventh Asilomar Conference on. IEEE.

Mazin Abdulrasool Hameed (2012). Comparative analysis of orthogonal matching pursuit and least angle regression. MSc thesis, Michigan State University. <https://www.google.gr/url?sa=t&rct=j&q=&esrc=s&source=web&>

Razavi S. A. Ollila E., & Koivunen V. (2012). Robust greedy algorithms for compressed sensing. In Signal Processing Conference (EUSIPCO), 2012 Proceedings of the 20th European. IEEE.

Lozano A., Swirszcz G., & Abe N. (2011). Group orthogonal matching pursuit for logistic regression. In Proceedings of the Fourteenth International Conference on Artificial Intelligence and Statistics.

See Also

[cor.fbed](#), [cor.fsreg](#), [correls](#), [fs.reg](#)

Examples

```
x <- matrix( rnorm(100 * 100), ncol = 100)
y <- rnorm(100)
b <- MXM::gomp(y, x, test = "testIndFisher")
```

Generate random folds for cross-validation

Generate random folds for cross-validation

Description

Random folds for use in a cross validation are generated. There is the option for stratified splitting as well.

Usage

```
generatefolds(target, nfolds = 10, stratified = TRUE, seed = FALSE)
```

Arguments

target	A vector with some data, either continuous or categorical.
nfolds	The number of folds to produce.
stratified	A boolean variable specifying whether stratified random (TRUE) or simple random (FALSE) sampling is to be used when producing the folds.
seed	A boolean variable. If set to TRUE, the folds will always be the same.

Details

I was inspired by the sam command in the package **TunePareto** in order to do the stratified version.

Value

A list with `nfolds` elements where each element is a fold containing the indices of the data.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

See Also

[cv.ses](#)

Examples

```
a <- generatefolds(iris[, 5], nfolds = 5, stratified = TRUE)
table(iris[a[[1]], 5) ## 10 values from each group
```

IAMB backward selection phase

IAMB backward selection phase

Description

IAMB backward selection phase.

Usage

```
iamb.bs(target, dataset, threshold = 0.05, wei = NULL, test = NULL, user_test = NULL)
```

Arguments

<code>target</code>	The class variable. Provide either a string, an integer, a numeric value, a vector, a factor, an ordered factor or a <code>Surv</code> object.
<code>dataset</code>	The dataset; provide either a data frame or a matrix (columns = variables, rows = observations). In either case, only two cases are available, either all data are continuous, or categorical.
<code>threshold</code>	Threshold (suitable values in (0,1)) for assessing p-values significance. Default value is 0.05.
<code>test</code>	The regression model to use. Available options are most of the tests for SES and MMPC. The ones NOT available are "gSquare", "censIndER", "testIndMVreg", "testIndClogit", "testIndSpearman" and "testIndFisher".
<code>wei</code>	A vector of weights to be used for weighted regression. The default value is <code>NULL</code> . It is not suggested when <code>robust</code> is set to <code>TRUE</code> .
<code>user_test</code>	A user-defined conditional independence test (provide a closure type object). Default value is <code>NULL</code> . If this is defined, the "test" argument is ignored.

Details

IAMB stands for Incremental Association Markov Blanket. The algorithm comprises of a forward selection and a modified backward selection process. This functions does the modified backward selection process. In the usual backward selection, among the non significant variabls, the one with the maximum p-value is dropped. So, one variable is removed at every step. In the IAMB backward phase, at aevery step, all non significant variables are removed. This makes it a lot faster.

Value

The output of the algorithm is a list of an S3 object including:

<code>runtime</code>	The run time of the algorithm. A numeric vector. The first element is the user time, the second element is the system time and the third element is the elapsed time.
<code>ci_test</code>	The conditional independence test used.
<code>vars</code>	The selected variables.
<code>mat</code>	A matrix with the selected variables and their latest test statistic and p-value. If no variable is selected this is NULL.
<code>final</code>	The final regression model.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

Tsamardinos, I., Aliferis, C. F., Statnikov, A. R., & Statnikov, E. (2003). Algorithms for Large Scale Markov Blanket Discovery. In FLAIRS conference, pp. 376-380.

See Also

[glm.fsreg](#), [lm.fsreg](#), [bic.fsreg](#), [bic.glm.fsreg](#), [CondIndTests](#), [MMPC](#), [SES](#)

Examples

```
set.seed(123)
dataset <- matrix( runif(1000 * 10, 1, 100), ncol = 10 )
target <- rnorm(1000)

a1 <- iamb.bs(target, dataset, threshold = 0.05, test = "testIndRQ")
a2 <- bs.reg(target, dataset, threshold = 0.05, test = "testIndRQ")
```

 IAMB variable selection

IAMB variable selection

Description

IAMB variable selection.

Usage

```
iamb(target, dataset, threshold = 0.05, wei = NULL, test = NULL, user_test = NULL,
      stopping = "BIC", tol = 2, ncores = 1, back = "iambbs")
```

Arguments

target	The class variable. Provide either a string, an integer, a numeric value, a vector, a factor, an ordered factor or a Surv object.
dataset	The dataset; provide either a data frame or a matrix (columns = variables, rows = observations). In either case, only two cases are available, either all data are continuous, or categorical.
threshold	Threshold (suitable values in (0,1)) for assessing p-values significance. Default value is 0.05.
wei	A vector of weights to be used for weighted regression. The default value is NULL. It is not suggested when robust is set to TRUE.
test	The regression model to use. Available options are most of the tests for SES and MMPC. The ones NOT available are "gSquare", "censIndER", "testIndMVreg", "testIndClogit", "testIndSpearman" and "testIndFisher" and "testIndIGreg".
user_test	A user-defined conditional independence test (provide a closure type object). Default value is NULL. If this is defined, the "test" argument is ignored.
stopping	The stopping rule. The BIC is always used for all methods. If you have linear regression though you can change this to "adjrsq" and in this case the adjusted R squared is used.
tol	The difference between two successive values of the stopping rule. By default this is set to 2. If for example, the BIC difference between two successive models is less than 2, the process stops and the last variable, even though significant does not enter the model.
ncores	How many cores to use. This plays an important role if you have tens of thousands of variables or really large sample sizes and tens of thousands of variables and a regression based test which requires numerical optimisation. In other cases it will not make a difference in the overall time (in fact it can be slower). The parallel computation is used in the first step of the algorithm, where univariate associations are examined, those take place in parallel. We have seen a reduction in time of 50% with 4 cores in comparison to 1 core. Note also, that the amount of reduction is not linear in the number of cores.

back The backward phase. If this "iambbs" (default value) the IAMB backward phase is performed and hence the IAMB algorithm is completed. If "bs", a simple backward selection phase is performed. This way, the IAMB algorithm is slightly more general.

Details

IAMB stands for Incremental Association Markov Blanket. The algorithm comprises of a forward selection and a modified backward selection process. This functions does the modified backward selection process. In the usual backward selection, among the non significant variabls, the one with the maximum p-value is dropped. So, one variable is removed at every step. In the IAMB backward phase, at aevery step, all non significant variables are removed. This makes it a lot faster.

Value

The output of the algorithm is a list of an S3 object including:

vars	A vector with the selcted variables.
mod	The output of the backward phase. In the case of no backward procedure this is the output of the forward phase.
mess	If the forward regression returned at most one variable, no backward procedure takes place and a message appears informing the user about this.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

Tsamardinos, I., Aliferis, C. F., Statnikov, A. R., & Statnikov, E. (2003). Algorithms for Large Scale Markov Blanket Discovery. In FLAIRS conference, pp. 376-380.

See Also

[glm.fsreg](#), [lm.fsreg](#), [bic.fsreg](#), [bic.glm.fsreg](#), [CondIndTests](#), [MMPC](#), [SES](#)

Examples

```
set.seed(123)
dataset <- matrix( runif(100 * 100, 1, 100), ncol = 100 )
target <- rnorm(100)

target = rpois(100, 10)
a1 <- iamb(target, dataset, threshold = 0.05, stopping = "BIC", tol = 0, back = "iambbs")
a2 <- iamb(target, dataset, threshold = 0.05, stopping = "BIC", tol = 0, back = "bs")
```

Incremental BIC values and final regression model of the FBED algorithm
Incremental BIC values and final regression model of the FBED algorithm

Description

Incremental BIC values and final regression model of the FBED algorithm.

Usage

```
fbedreg.bic(target, dataset, wei = NULL, fbedreg.object, test = NULL, graph = TRUE)
```

Arguments

target	A numeric vector containing the values of the target variable. It can also discrete data, binary data (as factor), nominal or ordinal data (as factor). In contrast to SES, no position of the target variable in the dataset is accepted. The target must be a numerical vector.
dataset	A numeric matrix or data.frame containing the variables. Rows are samples and columns are features. If you have categorical variables, this should be a data frame.
wei	A vector of weights to be used for weighted regression. The default value is NULL. We suggest not to use weights if you choose "testIndMMReg" as weights are already being used there.
fbedreg.object	An object with the results of an FBED run.
test	If you know the test used in SES put it here, otherwise leave it NULL. It will take this information from the SES object.
graph	If you want a graphical representation of the drop in the BIC values set this to TRUE.

Details

This function takes the output of the FBED ([fbed.reg](#)) and fits successive models calculating the BIC for each of them. A graph can also be returned.

Value

A list including:

res	A matrix with the selected variables, their test statistic and p-value (taken from the fbedreg.object) along with the BIC. Each row contains a BIC, that is the BIC of the model with the variables up to that row.
mod	The final model with all selected variables.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

Borboudakis G. and Tsamardinos I. (2017). Forward-Backward Selection with Early Dropping. <https://arxiv.org/pdf/1705.10770.pdf>

See Also

[reg.fit](#), [mmpc.model](#), [SES](#), [MMPC](#), [cv.ses](#), [cv.mmpc](#)

Examples

```
dataset <- matrix( runif(100 * 100, 1, 100), ncol = 100 )
#define a simulated class variable
target <- rt(100, 10)
a <- fbed.reg(target, dataset, K = 10, test = "testIndFisher", method = "eBIC")
fbedreg.bic(target, dataset, fbedreg.object = a, test = "testIndFisher")
```

Interactive plot of an (un)directed graph

Interactive plot of an (un)directed graph

Description

Interactive plot of an (un)directed graph.

Usage

```
plotnetwork(G, titlos)
```

Arguments

G	The adjacency matrix as produced from mmhc.skel , pc.skel , pc.con or any other algorithm. This can correspond to an undirected, partially directed or a completely directed graph.
titlos	A character argument specifying the title of the graph, for example "PC network".

Details

This visualises the directed graph.

Value

The plot of the directed graph. This is interactive, in the sense that the user can "play" with it. Move the nodes, zoom it, stretch it etc.

Author(s)

Michail Tsagris

R implementation and documentation: Giorgos Athineou <athineou@csd.uoc.gr> and Michail Tsagris <mtsagris@csd.uoc.gr>

See Also

[mmhc.skel](#), [nei](#), [pc.skel](#), [mb](#)

Examples

```
# simulate a dataset with continuous data
dataset <- matrix(runif(1000 * 20, 1, 100), nrow = 1000 )
a <- mmhc.skel(dataset, max_k = 3, threshold = 0.05, test = "testIndFisher",
nc = 1)
plotnetwork(a$G)
plotnetwork(a$G, titlos = "DAG skeleton")
```

Lower limit of the confidence of an edge

Lower limit of the confidence of an edge

Description

Lower limit of the confidence of an edge.

Usage

```
conf.edge.lower(p)
```

Arguments

p A numerical vector with the proportion of times an edge was found in the bootstrapped PC algorithm or the confidence of the edge returned by [bn.skel.utils2](#).

Details

After having performed PC algorithm many times in the bootstrap samples (using [pc.skel.boot](#) for example) you get a symmetric matrix with the proportion of times an edge was discovered. Take the lower (or upper) triangular elements of that matrix and pass them as input in this function. This will tell you the minimum proportion required to be confident that an edge is truly significant.

Value

The estimated cutoff limit above which an edge can be deemed significant.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

Scutari M. and Nagarajan R. (2013). Identifying significant edges in graphical models of molecular networks. *Artificial Intelligence in Medicine*, 57: 207-217.

See Also

[pc.skel.boot](#)

Examples

```
y <- rdag2(200, p = 40, nei = 3)
x <- y$x
g <- pc.skel.boot(x, R = 199)$Gboot
a <- g[ lower.tri(g) ]
conf.edge.lower(a)
```

mammpc.output-class *Class "mammpc.output"*

Description

mammpc.output object class.

Objects from the Class

Objects can be created by calls of the form `new("mammpc.output", ...)`.

Slots

selectedVars: Object of class "numeric"
selectedVarsOrder: Object of class "numeric"
hashObject: Object of class "list"
pvalues: Object of class "numeric"
stats: Object of class "numeric"
univ: Object of class "list"
max_k: Object of class "numeric"
threshold: Object of class "numeric"
test: Object of class "character"
runtime: Object of class "proc_time"

Methods

plot `plot(x = "mammpc.output", mode = "all")`: Generic function for plotting the generated pvalues of the MMPCoutput object. Argument `mode = "all"` for plotting all the pvalues or `mode="partial"` for partial plotting the first 500 pvalues

Author(s)

Giorgos Athineou <athineou@csd.uoc.gr>

See Also

[ma.mmmpc](#), [ma.ses](#)

Examples

```
showClass("mammpc.output")
```

Many simple beta regressions

Many simple beta regressions.

Description

Many simple beta regressions.

Usage

```
beta.regs(target, dataset, wei = NULL, check = FALSE, logged = FALSE, ncores = 1)
```

```
perm.betaregs(target, dataset, wei = NULL, check = FALSE, logged = FALSE,
threshold = 0.05, R = 999, ncores = 1)
```

```
wald.betaregs(target, dataset, wei = NULL, check = FALSE, logged = FALSE, ncores = 1)
```

Arguments

<code>target</code>	The target (dependent) variable. It must be a numerical vector with integers.
<code>dataset</code>	The independent variable(s). It can be a vector, a matrix or a dataframe with continuous only variables, a data frame with mixed or only categorical variables.
<code>wei</code>	A vector of weights to be used for weighted regression. The default value is NULL.
<code>check</code>	A boolean variable indicating whether to check for variables with identical values. The default is FALSE.
<code>logged</code>	A boolean variable; it will return the logarithm of the p-value if set to TRUE.
<code>threshold</code>	Threshold (suitable values in [0,1]) for assessing p-values significance. Default value is 0.05.

R	The number of permutations, set to 999 by default. There is a trick to avoid doing all permutations. As soon as the number of times the permuted test statistic is more than the observed test statistic is more than 50 (in this example case), the p-value has exceeded the significance level (threshold value) and hence the predictor variable is not significant. There is no need to continue do the extra permutations, as a decision has already been made.
ncores	The number of cores to use. The default value is 1.

Details

Many simple beta regressions are fitted.

Value

A matrix with the test statistic values, their relevant (logged) p-values and the BIC values.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

Ferrari S.L.P. and Cribari-Neto F. (2004). Beta Regression for Modelling Rates and Proportions. Journal of Applied Statistics, 31(7): 799-815.

See Also

[beta.mod](#), [testIndBeta](#), [reg.fit](#), [ridge.reg](#)

Examples

```
y <- rbeta(60, 5, 3)
x <- matrix( rnorm(60 * 10), ncol = 10)
a1 <- beta.regs(y, x)
a2 <- perm.betaregs(y, x[, 1:5])
```

Many simple zero inflated Poisson regressions

Many simple zero inflated Poisson regressions.

Description

Many simple zero inflated Poisson regressions.

Usage

```
zip.regs(target, dataset, wei = NULL, check = FALSE, logged = FALSE, ncores = 1)

perm.zipregs(target, dataset, wei = NULL, check = FALSE, logged = FALSE, R = 999,
threshold = 0.05, ncores = 1)

wald.zipregs(target, dataset, wei = NULL, check = FALSE, logged = FALSE, ncores = 1)
```

Arguments

target	The target (dependent) variable. It must be a numerical vector with integers.
dataset	The independent variable(s). It can be a vector, a matrix or a dataframe with continuous only variables, a data frame with mixed or only categorical variables.
wei	A vector of weights to be used for weighted regression. The default value is NULL.
check	A boolean variable indicating whether to check for variables with identical values. The default is FALSE.
logged	A boolean variable; it will return the logarithm of the pvalue if set to TRUE.
R	The number of permutations, set to 999 by default. There is a trick to avoid doing all permutations. As soon as the number of times the permuted test statistic is more than the observed test statistic is more than 50 (in this example case), the p-value has exceeded the significance level (threshold value) and hence the predictor variable is not significant. There is no need to continue do the extra permutations, as a decision has already been made.
threshold	Threshold (suitable values in [0,1]) for assessing p-values significance. Default value is 0.05.
ncores	The number of cores to use. The default value is 1.

Details

Many simple zero inflated Poisson regressions are fitted. The permutations based approach may cause some errors sometimes, and this is due to the nature of the distribution and its maximisation process. "nlm" and "optim" are used internally.

Value

A matrix with the test statistic values, their relevant (logged) p-values and the BIC values.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

Lambert D. (1992). Zero-inflated Poisson regression, with an application to defects in manufacturing. *Technometrics*, 34(1):1-14.

See Also

[zip.mod](#), [testIndZIP](#), [reg.fit](#), [ridge.reg](#)

Examples

```
y <- rpois(50, 3)
x <- matrix( rnorm(50 * 10), ncol = 10)
y[1:10] <- 0
a1 <- zip.regs(y, x)
a2 <- perm.zipregs(y, x[, 1:3])
```

Many Wald based tests for logistic and Poisson regressions with continuous predictors
Many Wald based tests for logistic and Poisson regressions with continuous predictors

Description

Many Wald based tests for logistic and Poisson regressions with continuous predictors.

Usage

```
wald.logisticregs(y, x, tol = 1e-09, wei = NULL, check = FALSE, logged = FALSE,
ncores = 1)
wald.poissonregs(y, x, tol = 1e-09, wei = NULL, check = FALSE, logged = FALSE,
ncores = 1)
```

Arguments

<code>y</code>	A vector with either 0s and 1 (logistic regression) or discrete data, counts (Poisson regression).
<code>x</code>	A data.frame, the predictor variables. If you have no categorical variables, the function will still work but it's better to use the score.glm s because it is faster.
<code>tol</code>	The tolerance value to stop the Newton-Raphson iterations. It is set to 1e-09 by default.
<code>wei</code>	A vector of weights to be used for weighted regression. The default value is NULL.
<code>check</code>	A boolean variable indicating whether to check for variables with identical values. The default is FALSE.
<code>logged</code>	A boolean variable; it will return the logarithm of the pvalue if set to TRUE.
<code>ncores</code>	How many to cores to useq the default value is 1.

Details

Instead of using R built-in function [glm](#) we implemented the newton-Raphson in order to avoid unnecessary calculations. The functions are much faster.

Value

A matrix with three columns, the test statistic, its associated (logged) p-value and the BIC of each model.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@yahoo.gr> and Manos Papadakis <papadakm95@gmail.com>.

References

Draper, N.R. and Smith H. (1988). Applied regression analysis. New York, Wiley, 3rd edition.

McCullagh, Peter, and John A. Nelder. Generalized linear models. CRC press, USA, 2nd edition, 1989.

See Also

[univregs](#), [perm.univregs](#)

Examples

```
## 200 variables, hence 200 univariate regressions are to be fitted
x <- matrix( rnorm(100 * 1000), ncol = 1000 )
y <- rpois(100, 4)
a <- wald.poissonregs(y, x)
b <- univregs(y, x, test = testIndPois)
cor(exp(a[, 2]), exp(b$pvalue) )
```

Markov Blanket of a node in a directed graph

Returns the Markov blanket of a node (or variable)

Description

Returns the Markov blanket of a node (or variable).

Usage

```
mb(G, node)
```

Arguments

G	The graph matrix as produced from pc.or or any other algorithm which produces directed graphs.
node	A vector with one or more numbers indicating the selected node(s) (or variable(s)).

Details

This is a way to see the network for some given nodes. It is useful if you have many nodes and the whole network is a bit difficult to see clearly. Bear in mind that the values can be extracted with the \$ symbol, i.e. this is an S3 class output.

Value

parents	The parents of the node of interest.
children	The children of the node of interest.
spouses	The spouses of the node of interest. These are the other parents of the children of the node of interest.
relatives	Nodes which are connected with the node of interest, but it is not known whether they are parents or children. The edge between them is undirected.
markov.blanket	The Markov blanket of the node of interest. The collection of all the previous.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

See Also

[plotnetwork](#), [nei](#), [pc.or](#)

Examples

```
# simulate a dataset with continuous data
# simulate a dataset with continuous data
y <- rdag(1000, 10, 0.3)
tru <- y$G
x <- y$x
mod <- pc.con(x)
G <- pc.or(mod)$G
plotnetwork(G)
dev.new()
mb(G, 8)
```

mases.output-class *Class "mases.output"*

Description

Meta analytic SES output object class.

Objects from the Class

Objects can be created by calls of the form `new("mases.output", ...)`.

Slots

selectedVars: Object of class "numeric"
selectedVarsOrder: Object of class "numeric"
queues: Object of class "list"
signatures: Object of class "matrix"
hashObject: Object of class "list"
pvalues: Object of class "numeric"
stats: Object of class "numeric"
univ: Object of class "list"
max_k: Object of class "numeric"
threshold: Object of class "numeric"
runtime: Object of class "proc_time"
test: Object of class "character"

Methods

plot plot(x = "mases.output", mode = "all"): Generic function for plotting the generated pvalues of the mases.output object. Argument mode = "all" for plotting all the pvalues or mode="partial" for partial plotting the first 500 pvalues

Author(s)

Giorgos Athineou <athineou@csd.uoc.gr>

See Also

[ma.ses](#), [ma.mmmpc](#)

Examples

```
showClass("mases.output")
```

MMPC solution paths for many combinations of hyper-parameters

MMPC solution paths for many combinations of hyper-parameters

Description

MMPC solution paths for many combinations of hyper-parameters.

Usage

```
mmpc.path(target, dataset, wei = NULL, max_ks = NULL, alphas = NULL, test = NULL,
user_test = NULL, ncores = 1)
```

```
waldmmpc.path(target, dataset, wei = NULL, max_ks = NULL, alphas = NULL, test = NULL,
user_test = NULL, ncores = 1)
```

Arguments

target	The class variable. Provide either a string, an integer, a numeric value, a vector, a factor, an ordered factor or a Surv object. See also Details.
dataset	The data-set; provide either a data frame or a matrix (columns = variables , rows = samples). Alternatively, provide an ExpressionSet (in which case rows are samples and columns are features, see bioconductor for details).
wei	A vector of weights to be used for weighted regression. The default value is NULL.
max_ks	A vector of possible max_k values. Can be a number as well, but this does not really make sense to do. If nothing is given, the values max_k = (4,3,2) are used by default.
alphas	A vector of possible threshold values. Can be a number as well, but this does not really make sense to do. If nothing is given, the values (0.1, 0.05, 0.01) are used by default.
test	The conditional independence test to use. Default value is NULL. See also CondIndTests .
user_test	A user-defined conditional independence test (provide a closure type object). Default value is NULL. If this is defined, the "test" argument is ignored.
ncores	How many cores to use. This plays an important role if you have tens of thousands of variables or really large sample sizes and tens of thousands of variables and a regression based test which requires numerical optimisation. In other cases it will not make a difference in the overall time (in fact it can be slower). The parallel computation is used in the first step of the algorithm, where univariate associations are examined, those take place in parallel. We have seen a reduction in time of 50% with 4 cores in comparison to 1 core. Note also, that the amount of reduction is not linear in the number of cores. This argument is used only in the first run of MMPC and for the univariate associations only and the results are stored (hashed). In the enxt runs of MMPC the results are used (cached) and so the process is faster.

Details

For different combinations of the hyper-parameters, max_k and the significance level (threshold or alpha) the MMPC algorithm is run.

Value

The output of the algorithm is an object of the class 'SESoutput' for SES or 'MMPCoutput' for MMPC including:

bic	A matrix with the BIC values of the final fitted model based on the selected variables identified by each configuration, combination of the hyper-parameters.
size	A matrix with the length of the selected variables identified by each configuration of MMPC.
variables	A list containing the variables from each configuration of MMPC
runtime	The run time of the algorithm. A numeric vector. The first element is the user time, the second element is the system time and the third element is the elapsed time.

Author(s)

Ioannis Tsamardinos, Vincenzo Lagani

R implementation and documentation: Giorgos Athineou <athineou@csd.uoc.gr> Vincenzo Lagani <vlagani@csd.uoc.gr>

References

I. Tsamardinos, V. Lagani and D. Pappas (2012). Discovering multiple, equivalent biomarker signatures. In proceedings of the 7th conference of the Hellenic Society for Computational Biology & Bioinformatics-HSCBB12.

Tsamardinos, Brown and Aliferis (2006). The max-min hill-climbing Bayesian network structure learning algorithm. Machine learning, 65(1): 31-78.

See Also

[CondIndTests](#), [cv.ses](#)

Examples

```
set.seed(123)
require(hash)
# simulate a dataset with continuous data
dataset <- matrix(runif(1000 * 101, 1, 100), nrow = 1000 )
#the target feature is the last column of the dataset as a vector
target <- dataset[, 101]
dataset <- dataset[, -101]

a <- mmpc.path(target, dataset, max_ks = NULL, alphas = NULL, test = NULL,
user_test = NULL, ncores = 1)
```

MMPC.gee.output-class *Class* "MMPC.gee.output"

Description

MMPC.temporal output object class.

Objects from the Class

Objects can be created by calls of the form `new("MMPC.gee.output", ...)`.

Slots

`selectedVars`: Object of class "numeric"
`selectedVarsOrder`: Object of class "numeric"
`hashObject`: Object of class "list"
`pvalues`: Object of class "numeric"
`stats`: Object of class "numeric"
`univ`: Object of class "list"
`max_k`: Object of class "numeric"
`threshold`: Object of class "numeric"
`runtime`: Object of class "proc_time"
`test`: Object of class "character"
`correl`: Object of class "character"
`se`: Object of class "character"

Methods

plot `plot(x = "MMPC.gee.output", mode = "all")`: Generic function for plotting the generated pvalues of the `MMPC.temporal.output` object. Argument `mode = "all"` for plotting all the pvalues or `mode="partial"` for partial plotting the first 500 pvalues

Author(s)

Giorgos Athineou and Michail Tsagris <mtsagris@csd.uoc.gr>

See Also

[MMPC.gee](#), [SES.gee](#)

Examples

```
showClass("MMPC.gee.output")
```

```
MMPC.temporal.output-class
      Class "MMPC.temporal.output"
```

Description

MMPC.temporal output object class.

Objects from the Class

Objects can be created by calls of the form `new("MMPC.temporal.output", ...)`.

Slots

selectedVars: Object of class "numeric"
selectedVarsOrder: Object of class "numeric"
hashObject: Object of class "list"
pvalues: Object of class "numeric"
stats: Object of class "numeric"
univ: Object of class "list"
max_k: Object of class "numeric"
threshold: Object of class "numeric"
runtime: Object of class "proc_time"
test: Object of class "character"
slope: Object of class "logical"

Methods

plot `plot(x = "MMPC.temporal.output", mode = "all")`: Generic function for plotting the generated pvalues of the MMPC.temporal.output object. Argument `mode = "all"` for plotting all the pvalues or `mode="partial"` for partial plotting the first 500 pvalues

Author(s)

Giorgos Athineou <athineou@csd.uoc.gr>

See Also

[MMPC.temporal](#), [SES.temporal](#)

Examples

```
showClass("MMPC.temporal.output")
```

MMPCoutput-class	Class "MMPCoutput"
------------------	--------------------

Description

MMPC output object class.

Objects from the Class

Objects can be created by calls of the form `new("MMPCoutput", ...)`.

Slots

selectedVars: Object of class "numeric"
selectedVarsOrder: Object of class "numeric"
hashObject: Object of class "list"
pvalues: Object of class "numeric"
stats: Object of class "numeric"
univ: Object of class "list"
max_k: Object of class "numeric"
threshold: Object of class "numeric"
n.tests: Object of class "numeric"
runtime: Object of class "proc_time"
test: Object of class "character"

Methods

plot `plot(x = "MMPCoutput", mode = "all")`: Generic function for plotting the generated pvalues of the MMPCoutput object. Argument `mode = "all"` for plotting all the pvalues or `mode="partial"` for partial plotting the first 500 pvalues

Author(s)

Giorgos Athineou <athineou@csd.uoc.gr>

See Also

[MMPC](#), [SES](#)

Examples

```
showClass("MMPCoutput")
```

Description

Internal functions of Package **MXM**

Details

These functions are only for internal usage of the MXM package - NOT to be called by the user.

Functions

- InternalSES(...)
- InternalMMPC(...)
- Internalmasess(...)
- Internalmammpc(...)
- IdentifyEquivalence(...)
- IdentifyEquivalence.ma(...)
- apply_ideq(...)
- apply_ideq.ma(...)
- compare_p_values(...)
- identifyTheEquivalent(...)
- identifyTheEquivalent.ma(...)
- max_min_assoc(...)
- max_min_assoc.ma(...)
- min_assoc(...)
- min_assoc.ma(...)
- univariateScore(...)
- univariateScore.ma(...)
- condi.perm(...)
- InternalSES.temporal(...)
- InternalMMPC.temporal(...)
- IdentifyEquivalence.temporal(...)
- apply_ideq.temporal(...)
- identifyTheEquivalent.temporal(...)
- max_min_assoc.temporal(...)
- min_assoc.temporal(...)
- univariateScore.temporal(...)

- `is.sepset(...)`
- `lm.fsreg_2(...)`
- `glm.fsreg_2(...)`
- `dag_to_eg(...)`
- `nchoosek(...)`
- `R0(...)`
- `R1(...)`
- `R2(...)`
- `R3(...)`
- `is.sepset(...)`
- `regbeta(...)`
- `regbetawei(...)`
- `betamle.wei(...)`
- `regzip(...)`
- `regzipawei(...)`
- `zipmle.wei(...)`
- `zipwei(...)`
- `bic.betafsreg(...)`
- `bic.zipfsreg(...)`
- `beta.fsreg(...)`
- `zip.fsreg(...)`
- `beta.bsreg(...)`
- `zip.bsreg(...)`
- `iamb.betabs(...)`
- `iamb.zipbs(...)`
- `iamb.glmbs(...)`
- `internaliamb.binombs(...)`
- `internaliamb.poisbs(...)`
- `internaliamb.lmbs(...)`
- `InternalMMPC.temporal(...)`
- `InternalSES.temporal(...)`
- `InternalSES(...)`
- `InternalMMPC(...)`
- `univariateScore(...)`
- `perm.univariateScore(...)`
- `max_min_assoc(...)`
- `min_assoc(...)`

- `nchoosek(...)`
- `Internalmasers(...)`
- `compare_p_values(...)`
- `perm.Internalmmpc`
- `wald.Internalmmpc`
- `wald.Internalses`
- `perm.IdentifyEquivalence(...)`
- `perm.identifyTheEquivalent(...)`
- `perm.apply_ideq(...)`
- `IdentifyEquivalence(...)`
- `apply_ideq(...)`
- `identifyTheEquivalent(...)`
- `cvses.par(...)`
- `cvmmpc.par(...)`
- `cvwaldses.par(...)`
- `cvwaldmmpc.par(...)`
- `cvpermses.par(...)`
- `cvpermmmpc.par(...)`
- `wald.univariateScore(...)`
- `univariateScore.ma(...)`
- `IdentifyEquivalence.ma(...)`
- `identifyTheEquivalent.ma(...)`
- `apply_ideq.ma(...)`
- `max_min_assoc.ma(...)`
- `min_assoc.ma(...)`
- `fs.reg_2(...)`
- `gammafsreg_2(...)`
- `beta.fsreg_2(...)`
- `zip.fsreg_2(...)`
- `internaliamb.zipbs(...)`
- `internaliamb.betabs(...)`
- `iamb.gammabs(...)`
- `iamb.normlogbs(...)`
- `internaliamb.gammabs(...)`
- `internaliamb.normlogbs(...)`
- `bic.tobit.fsreg(...)`
- `iamb.tobitbs(...)`

- `internaliamb.tobitbs(...)`
- `fbed.beta(...)`
- `fbed.cr(...)`
- `fbed.glm(...)`
- `fbed.lm(...)`
- `fbed.mmreg(...)`
- `fbed.multinom(...)`
- `fbed.nb(...)`
- `fbed.ordinal(...)`
- `fbed.tobit(...)`
- `fbed.wr(...)`
- `fbed.zip(...)`
- `ebic.fbed.beta(...)`
- `ebic.fbed.cr(...)`
- `ebic.fbed.glm(...)`
- `ebic.fbed.lm(...)`
- `ebic.fbed.mmreg(...)`
- `ebic.fbed.multinom(...)`
- `ebic.fbed.nb(...)`
- `ebic.fbed.ordinal(...)`
- `ebic.fbed.tobit(...)`
- `ebic.fbed.wr(...)`
- `ebic.fbed.zip(...)`
- `ebic.beta.bsreg(...)`
- `ebic.cr.bsreg(...)`
- `ebic.glm.bsreg(...)`
- `ebic.lm.bsreg(...)`
- `ebic.mm.bsreg(...)`
- `ebic.multinom.bsreg(...)`
- `ebic.ordinal.bsreg(...)`
- `ebic.tobit.bsreg(...)`
- `ebic.wr.bsreg(...)`
- `ebic.zip.bsreg(...)`
- `cat_condis(...)`
- `pearson_condis(...)`
- `pearson_condis.rob(...)`
- `disctor_condis(...)`

- `comb_condis(...)`
- `fbed.glmm(...)`
- `fbed.lmm(...)`
- `ebic.fbed.glmm(...)`
- `ebic.fbed.lmm(...)`
- `lmm.bsreg(...)`
- `clogit.fsreg(...)`
- `clogit.fsreg_2(...)`
- `bic.clogit.fsreg(...)`
- `fbed.glm2(...)`
- `quasibinom.bsreg(...)`
- `quasipois.bsreg(...)`
- `quasibinom.fsreg(...)`
- `quasipois.fsreg(...)`
- `quasibinom.fsreg_2(...)`
- `quasipois.fsreg_2(...)`
- `fbed.geeglm(...)`
- `fbed.geeglm.reps(...)`
- `fbed.ordgee(...)`
- `internaliamb.mmbs(...)`
- `iamb.gammabs(...)`
- `iamb.normlogbs(...)`
- `internaliamb.gammabs(...)`
- `internaliamb.normlogbs(...)`
- `bic.wr.fsreg(...)`
- `wr.fsreg(...)`
- `wr.fsreg_2(...)`
- `ebicScore(...)`
- `fbed.glmm.reps(...)`
- `fbed.lmm.reps(...)`
- `ebic.fbed.lmm.reps(...)`
- `ebic.fbed.glmm.reps(...)`
- `ebic.glmm.reps.bsreg(...)`
- `fbed.ordgee.reps(...)`
- `fbed.geelm(...)`
- `fbed.geelm.reps(...)`
- `univariateScore.gee(...)`

- `InternalMMPC.gee(..)`
- `InternalSES.gee(..)`
- `max_min_assoc.gee(...)`
- `min_assoc.gee(...)`
- `IdentifyEquivalence.gee(...)`
- `identifyTheEquivalent.gee(...)`
- `univariateScore.timeclass(...)`
- `InternalMMPC.timeclass(...)`
- `gomp2(...)`
- `mm.bsreg(...)`
- `kfbed.reg(...)`
- `kfbed.glmm.reg(...)`
- `kfbed.gee.reg(...)`

Neighbours of nodes in an undirected graph

Returns the node(s) and their neighbour(s), if there are any.

Description

Returns the node(s) and their neighbour(s) of one or more nodes (if there are any).

Usage

```
nei(G, node)
```

Arguments

G	The adjacency matrix of an undirected graph as produced by <code>mmhc.skel</code> , <code>pc.skel</code> or any other algorithm.
node	A vector with one or more numbers indicating the selected node(s) (or variable(s)).

Details

This is a way to see the network for some given nodes. It is useful if you have many nodes and the whole network is a bit difficult to see clearly.

Value

A list object called "geit" containing the neighbours of the node(s). If there are no neighbours a message appears and no plot is presented. If the "graph" argument is set to TRUE and there are neighbours, a plot will appear.

Bear in mind that the values can be extracted with the \$ symbol, i.e. this is an S3 class output.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

See Also

[mmhc.skel](#), [SES](#), [MMPC](#)

Examples

```
# simulate a dataset with continuous data
set.seed(1234)
dataset <- matrix(runif(1000 * 20, 1, 100), nrow = 1000 )
G <- pc.con(dataset)$G
plotnetwork(G)
dev.new()
nei( G, c(3, 4) )
nei( G, c(1, 3) )
```

Network construction using the partial correlation based forward regression or FBED
Network construction using the partial correlation based forward regression of FBED

Description

Network construction using the partial correlation based forward regression or FBED.

Usage

```
corfs.network(x, threshold = 0.05, tolb = 2, tolr = 0.02, stopping = "BIC",
symmetry = TRUE, nc = 1)
```

```
corfbed.network(x, threshold = 0.05, symmetry = TRUE, nc = 1)
```

Arguments

x	A matrix with continuous data.
threshold	Threshold (suitable values in (0, 1)) for assessing p-values significance. Default value is 0.05.
tolb	The difference in the BIC bewtween two successive values. By default this is set to 2. If for example, the BIC difference between two successive models is less than 2, the process stops and the last variable, even though significant does not enter the model.

tolr	The difference in the adjusted R^2 between two successive values. By default this is set to 0.02. If for example, the difference between the adjusted R^2 of two successive models is less than 0.02, the process stops and the last variable, even though significant does not enter the model.
stopping	The stopping rule. The "BIC" is the default value, but can change to "ar2" and in this case the adjusted R^2 is used. If you want both of these criteria to be satisfied, type "BICR2".
symmetry	In order for an edge to be added, a statistical relationship must have been found from both directions. If you want this symmetry correction to take place, leave this boolean variable to TRUE. If you set it to FALSE, then if a relationship between Y and X is detected but not between X and Y, the edge is still added.
nc	How many cores to use. This plays an important role if you have many variables, say thousands or so. You can try with $nc = 1$ and with $nc = 4$ for example to see the differences. If you have a multicore machine, this is a must option. There was an extra argument for plotting the skeleton but it does not work with the current visualisation packages, hence we removed the argument. Use plotnetwork to plot the skeleton.

Details

In the MMHC algorithm (see [mmhc.skel](#)), the MMPC or SES algorithms are run for every variable. Hence, one can use forward regression for each variable and this is what we are doing here. Partial correlation forward regression is very efficient, since only correlations are being calculated.

Value

A list including:

runtime	The run time of the algorithm. A numeric vector. The first element is the user time, the second element is the system time and the third element is the elapsed time.
density	The number of edges divided by the total possible number of edges, that is $\#edges / n(n - 1)/2$, where n is the number of variables.
info	Some summary statistics about the edges, minimum, maximum, mean, median number of edges.
ntests	The number of tests MMPC (or SES) performed at each variable.
G	The adjacency matrix. A value of 1 in $G[i, j]$ appears in $G[j, i]$ also, indicating that i and j have an edge between them.

Bear in mind that the values can be extracted with the \$ symbol, i.e. this is an S3 class output.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

- Sanford Weisberg (2014). Applied Linear Regression. Hoboken NJ: John Wiley, 4th edition.
- Draper N.R. and Smith H. (1988). Applied regression analysis. New York, Wiley, 3rd edition.
- Tsamardinos, Brown and Aliferis (2006). The max-min hill-climbing Bayesian network structure learning algorithm. Machine learning, 65(1), 31-78.
- Borboudakis G. and Tsamardinos I. (2017). Forward-Backward Selection with Early Dropping. <https://arxiv.org/pdf/1705.10770.pdf>

See Also

[mmhc.skel](#), [pc.skel](#)

Examples

```
# simulate a dataset with continuous data
dataset <- matrix(runif(1000 * 50, 1, 100), nrow = 1000 )
a1 <- mmhc.skel(dataset, max_k = 3, threshold = 0.05, test = "testIndFisher",
nc = 1)
a2 <- corfs.network(dataset, threshold = 0.05, tolb = 2, tolr = 0.02, stopping = "BIC",
symmetry = TRUE, nc = 1)
a1$runtime
a2$runtime
```

Orientation rules for the PC algorithm

The orientations part of the PC algorithm.

Description

The function takes the outcome of the PC algorithm, as produced by [pc.skel](#) or [pc.con](#) and performs the 4 orientation rules. A graph is also possible to visualize.

Usage

```
pc.or(mod)
```

Arguments

`mod` An object with the results of the PC algorithm, as produced by [pc.skel](#) or [pc.con](#). There was an extra argument for plotting the skeleton but it does not work with the current visualisation packages, hence we removed the argument. Use [plotnetwork](#) to plot the skeleton.

Details

After having calculated the skeleton of the PC algorithm one may want to perform orientations, leading to causal relationships. The rules as stated in Spirtes, Glymour and Scheines (2001) are

1. **Rule 1.** For each triple of vertices X, Y, Z such that the pair X, Y and the pair Y, Z are each adjacent in C but the pair X, Z are not adjacent in C , orient $X - Y - Z$ as $X \rightarrow Y \leftarrow Z$ if and only if Y is not in $\text{Sepset}(X, Z)$.
2. **Rule 2.** If $A \rightarrow B$, B and C are adjacent, A and C are not adjacent, and there is no arrowhead at B , then orient $B - C$ as $B \rightarrow C$.
3. **Rule 3.** If there is a directed path from A to B , and an edge between A and B , then orient $A - B$ as $A \rightarrow B$.
4. **Rule 4.** This was added by Zhang (2008). If $A \rightarrow B \leftarrow C$, $A - D - C$, A and C are not adjacent, and $D - B$, then orient $D - B$ as $D \rightarrow B$.

The first rule is applied once. Rules 2-4 are applied repeatedly until no more edges can be oriented. If however the argument "prve.cycles" is set to TRUE, then at any rule a cycle is detected, the rule is cancelled. Also, when applying Rules 1-3 we try to avoid the recreation of new v-structures ($X \rightarrow Y \leftarrow Z$).

Value

A list including:

Gini	The initial adjacency matrix, no orientations. This is the matrix produced by pc.skel or pc.con .
G	The final adjacency matrix with the orientations. If $G[i, j] = 2$ then $G[j, i] = 3$. This means that there is an arrow from node i to node j . If $G[i, j] = G[j, i] = 0$; there is no edge between nodes i and j . If $G[i, j] = G[j, i] = 1$; there is an (undirected) edge between nodes i and j .
runtime	The run time of the algorithm. A numeric vector. The first element is the user time, the second element is the system time and the third element is the elapsed time.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

Spirtes P., Glymour C. and Scheines R. (2001). Causation, Prediction, and Search. The MIT Press, Cambridge, MA, USA, 3rd edition.

Zhang, Jiji. (2008). On the completeness of orientation rules for causal discovery in the presence of latent confounders and selection bias. *Artificial Intelligence* 172(16): 1873–1896.

See Also

[pc.con](#), [pc.skel](#), [mmhc.skel](#), [is.dag](#), [mb](#)

Examples

```
# simulate a dataset with continuous data
y <- rdag2(2000, p = 20, nei = 3)
ind <- sample(1:20, 20)
tru <- y$G[ind, ind]
x <- y$x[, ind]
mod <- pc.con(x)
mod$runtime

plotnetwork(tru)

b <- pc.or(mod)
plotnetwork(b$G)

plotnetwork( dag2eg(tru) ) ## essential graph
plotnetwork(b$G)
```

Partial correlation between two variables
Partial correlation

Description

Partial correlation between two variables when a correlation matrix is given.

Usage

```
partialcor(R, indx, indy, indz)
```

Arguments

R	A correlation matrix.
indx	The index of the first variable whose conditional correlation is to estimated.
indy	The index of the second variable whose conditional correlation is to estimated.
indz	The index of the conditioning variables.

Details

Given a correlation matrix the function will calculate the partial correlation between variables `indx` and `indy` conditioning on variable(s) `indz`.

Value

A number, the partial correlation coefficient.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

See Also

[testIndFisher](#), [testIndSpearman](#), [permcors](#), [pc.con](#)

Examples

```
r <- cor( iris[, 1:4] )
partialcor(r, 1, 2, 0)
r[1, 2] ## the same as above
```

```
y = as.vector( iris[, 1] )
x = as.vector( iris[, 2] )
z = as.vector( iris[, 3] )
e1 = resid( lm(y ~ z) )
e2 = resid( lm(x ~ z) )
cor(e1, e2)
partialcor(r, 1, 2, 3)
```

Permutation based p-value for the Pearson correlation coefficient

Permutation based p-value for the Pearson correlation coefficient

Description

The main task of this test is to provide a p-value PVALUE for the null hypothesis: feature 'X' is independent from 'TARGET' given a conditioning set CS.

Usage

```
permcors(x1, x2, R = 999)
permcorsels(y, x, R = 999)
```

Arguments

x1	A numerical vector.
x2	A numerical vector of the same size as x1.
y	A vector whose length is equal to the number of rows of x.
x	This is a matrix with many variables.
R	The number of permutations to be conducted; set to 999 by default.

Details

This is a computational non parametric (permutation based) correlation coefficient test and is advised to be used when a small sample size is available. If you want to use the Spearman correlation instead, simply provide the ranks of x or of y and x.

Value

For the case of "permcov" a vector consisting of two values, the Pearson correlation and the permutation based p-value. For the "permcovrels" a vector with three values, the Pearson correlation, the test statistic value and the permutation based p-value.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

Legendre Pierre (2000). Comparison of permutation methods for the partial correlation and partial Mantel tests. *Journal of Statistical Computation and Simulation* 67(1):37-73.

See Also

[pc.skel](#), [testIndSpearman](#), [testIndFisher](#), [SES](#), [CondIndTests](#)

Examples

```
MXM::permcov(iris[, 1], iris[, 2], R = 9999)
x <- matrix(rnorm(50 * 2000), ncol = 2000)
a <- permcovrels(iris[1:50, 1], x)
```

Plot of longitudinal data

Plot of longitudinal data

Description

Plot of longitudinal data.

Usage

```
tc.plot(target, tp, id, type = "l", ylab = "Values", xlab = "Time points",
        col = 2, lwd = 1, lty = 2, pch = 1, main = "Spaghetti plot")
```

Arguments

target	A numerical vector with the longitudinal data.
tp	The time points. It can either be a vector with length either equal to the number of time points or equal to the length of the target.
id	A numerical vector specifying the subjects. It can either be a vector with length either equal to the number of subjects or equal to the length of the target.
type	This is a graphical parameter. You can have lines "l" everywhere or lines with points at each time point "p".
ylab	This is a graphical parameter. The label on the y axis.
xlab	This is a graphical parameter. The label on the x axis.
col	This is a graphical parameter. The color of the lines.
lwd	This is a graphical parameter. The thickness of the lines.
lty	This is a graphical parameter. The type of line, e.g. dashed, dotted, etc.
pch	This is a graphical parameter. If the type is "b", then you can specify if you want different signs, for example circles, crosses, diamonds etc.
main	This is a graphical parameter. The title of the graph.

Details

The data must be longitudinal (the same subject measured multiple times at different time points) and for one variable only. For the graphical parameters see [plot](#) or [par](#).

Value

A plot with the longitudinal data over time.

Author(s)

Michail Tsagris

R implementation and documentation: Giorgos Athineou <athineou@csd.uoc.gr> Michail Tsagris <mtsagris@csd.uoc.gr>

References

Tsamardinos I., Aliferis C. F. and Statnikov, A. (2003). Time and sample efficient discovery of Markov blankets and direct causal relations. In Proceedings of the 9th ACM SIGKDD international conference on Knowledge discovery and data mining (pp. 673-678).

See Also

[testIndGLMMReg](#), [SES.temporal](#)

Examples

```

y <- rnorm(70)
Days <- rep(1:7, each = 10)
id <- rep(1:10, 7)
tc.plot(y, Days, id)
tc.plot(y, Days, id, type = "b")

```

Regression models based on SES and MMPC outputs

Regression model(s) obtained from SES or MMPC

Description

One or more regression models obtained from SES or MMPC, are returned.

Usage

```
ses.model(target, dataset, wei = NULL, sesObject, nsignal = 1, test = NULL)
```

```
mmpc.model(target, dataset, wei = NULL, mmpcObject, test = NULL)
```

```
waldses.model(target, dataset, wei = NULL, wald.sesObject, nsignal = 1, test = NULL)
```

```
waldmmpc.model(target, dataset, wei = NULL, wald.mmpcObject, test = NULL)
```

Arguments

target	A numeric vector containing the values of the target variable. If the values are proportions or percentages, i.e. strictly within 0 and 1 they are mapped into R using $\log(\text{target}/(1 - \text{target}))$. It can also discrete data, binary data (as factor), nominal or ordinal data (as factor). In contrast to SES, no position of the target variable in the dataset is accepted. The target must be a numerical vector.
dataset	A numeric matrix or data.frame containing the variables. Rows are samples and columns are features. If you have categorical variables, this should be a data frame.
wei	A vector of weights to be used for weighted regression. The default value is NULL.
sesObject	An object with the results of a SES run.
mmpcObject	An object with the results of an MMPC run.
wald.sesObject	An object with the results of a wald.ses run.
wald.mmpcObject	An object with the results of an wald.mmpc run.

<code>nsignat</code>	How many signatures to use. If <code>nsignat = 1</code> (default value) the first set of variables will be used for the model. If you want more, then specify the number of signatures you want. If you want the models based on all signatures, specify "all". If you put a number which is higher than the number of signatures, all models will be returned.
<code>test</code>	If you know the test used in SES put it here, otherwise leave it NULL. It will take this information from the SEs object. If you used a robust version of a test (wherever possible), robust model(s) will be created.

Details

This command is useful if you want to see all models and check for example their fitting ability, MSE in linear models for example.

Value

A list including:

<code>mod</code>	Depending on the number of signatures requested, one or models will be returned.
<code>signature</code>	A matrix (or just one vector if one signature only) with the variables of each signature, along with the BIC of the corresponding regression model.

Author(s)

Michail Tsagris

R implementation and documentation: Giorgos Athineou <athineou@csd.uoc.gr> and Michail Tsagris <mtsagris@csd.uoc.gr>

References

- Aitchison J. (1986). *The Statistical Analysis of Compositional Data*, Chapman & Hall; reprinted in 2003, with additional material, by The Blackburn Press.
- Cox D.R. (1972). Regression models and life-tables. *J. R. Stat. Soc.*, 34, 187-220.
- Draper, N.R. and Smith H. (1988). *Applied regression analysis*. New York, Wiley, 3rd edition.
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- Gutenbrunner C., Jureckova J., Koenker R. and Portnoy S. (1993). Tests of Linear Hypothesis based on Regression Rank Scores, *Journal of NonParametric Statistics* 2, 307-331.
- Joseph M.H. (2011). *Negative Binomial Regression*. Cambridge University Press, 2nd edition.
- Koenker R.W. (2005). *Quantile Regression*, Cambridge University Press.
- Lagani V., Kortas G. and Tsamardinos I. (2013). Biomarker signature identification in "omics" with multiclass outcome. *Computational and Structural Biotechnology Journal*, 6(7): 1-7.
- Lagani V. and Tsamardinos I. (2010). Structure-based variable selection for survival data. *Bioinformatics Journal* 16(15): 1887-1894.

Lambert, Diane (1992). Zero-inflated Poisson regression, with an application to defects in manufacturing. *Technometrics* 34(1): 1-14.

Mardia K.V., Kent J.T. and Bibby J.M. (1979). *Multivariate Analysis*, Academic Press, New York, USA.

Maronna R.D. Yohai M.V. (2006). *Robust Statistics, Theory and Methods*. Wiley.

McCullagh P., and Nelder J.A. (1989). *Generalized linear models*. CRC press, USA, 2nd edition.

See Also

[SES](#), [MMPC](#), [cv.ses](#), [cv.mmpc](#)

Examples

```
# simulate a dataset with continuous data
dataset <- matrix(runif(1000 * 101, 1, 100), nrow = 1000 )
#the target feature is the last column of the dataset as a vector
target <- dataset[, 101]
dataset <- dataset[, -101]
sesObject <- SES(target , dataset , max_k=3 , threshold = 0.05);
ses.model(target, dataset, sesObject = sesObject, nsigmat = 1, test = NULL)
ses.model(target, dataset, sesObject = sesObject, nsigmat = 40, test = NULL)
mmpcObject <- MMPC(target , dataset , max_k=3 , threshold = 0.05);
mmpc.model(target, dataset, mmpcObject = mmpcObject, test = NULL)
```

Regression models based on *SES.timeclass* and *MMPC.timeclass* outputs

Regression model(s) obtained from SES.timeclass or MMPC.timeclass

Description

One or more regression models obtained from *SES.timeclass* or *MMPC.timeclass*, are returned.

Usage

```
mmpc.timeclass.model(target, dataset, id, reps, wei = NULL, mmpctimeclass.Object)
ses.timeclass.model(target, dataset, id, reps, wei = NULL, sestimeclass.Object,
nsigmat = 1)
```

Arguments

target	The class variable. Provide a vector or a factor with discrete numbers indicating the class. Its length is equal to the number of rows of the dataset.
dataset	The dataset; provide a matrix. Currently, only continuous datasets are supported. The dataset contains longitudinal data, where each column is a variable. The repeated measurements are the samples.
id	A numeric vector containing the subjects or groups. Its length is equal to the number of rows of the dataset.

<code>reps</code>	A numeric vector containing the time points of the subjects. Its length is equal to the number of rows of the dataset.
<code>wei</code>	A vector of weights to be used for weighted regression. The default value is <code>NULL</code> .
<code>mmptimeclass.Object</code>	An object with the results of an <i>MMPC.timeclass</i> run.
<code>sestimeclass.Object</code>	An object with the results of a <i>SES.timeclass</i> run.
<code>nsignat</code>	How many signatures to use. If <code>nsignat = 1</code> (default value) the first set of variables will be used for the model. If you want more, then specify the number of signatures you want. If you want the models based on all signatures, specify "all". If you put a number which is higher than the number of signatures, all models will be returned.

Details

This command is useful if you want to see all models and check for example their fitting ability.

Value

A list including:

<code>mod</code>	Depending on the number of signatures requested, one or models will be returned.
<code>signature</code>	A matrix (or just one vector if one signature only) with the variables of each signature, along with the BIC of the corresponding regression model.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

Tsagris, M., Lagani, V., & Tsamardinos, I. (2018). Feature selection for high-dimensional temporal data. *BMC bioinformatics*, 19(1), 17.

McCullagh P., and Nelder J.A. (1989). *Generalized linear models*. CRC press, USA, 2nd edition.

See Also

[MMPC.timeclass](#)

Examples

```
## assume these are longitudinal data, each column is a variable (or feature)
dataset <- matrix( rnorm(400 * 100), ncol = 100 )
id <- rep(1:80, each = 5) ## 80 subjects
reps <- rep( seq(4, 12, by = 2), 80) ## 5 time points for each subject
## dataset contains are the regression coefficients of each subject's values on the
```

```
## reps (which is assumed to be time in this example)
target <- rep(0:1, each = 200)
a <- MMPC.timeclass(target, reps, id, dataset)
mmpc.timeclass.model(target, dataset, id, reps, mmpctimeclass.Object = a)
```

Regression models fitting

Regression modelling

Description

Generic regression modelling function.

Usage

```
reg.fit(y, dataset, event = NULL, reps = NULL, group = NULL, slopes = FALSE,
reml = FALSE, model = NULL, wei = NULL, xnew = NULL)
```

Arguments

y	The target (dependent) variable. It can be a numerical variable, factor, ordinal factor, percentages, matrix, or time to event. If the values are proportions or percentages, i.e. strictly within 0 and 1 they are mapped into R using $\log(\text{target}/(1 - \text{target}))$. If they are compositional data the additive log-ratio (multivariate logit) transformation is applied beforehand.
dataset	The independent variable(s). It can be a vector, a matrix or a dataframe with continuous only variables, a data frame with mixed or only categorical variables.
event	This is NULL unless you have time to event data (survival regression).
reps	This is NULL unless you have time measurements (longitudinal data).
group	This is NULL unless you have grouped (or clustered) data or longitudinal data (is the latter case the argument reps is required).
slopes	This is for the longitudinal data only, TRUE or FALSE. Should random slopes be added or not?
reml	This is for the longitudinal or grouped data only, TRUE or FALSE. If TRUE, REML will be used, otherwise ML will be used.
model	The type of model you want to use. It can be specified by the user or left NULL, if other correct arguments have been passed. Possible values (apart from NULL) are: "gaussian" (default), "binary", "binomial", "multinomial", "poisson", "ordinal", "gamma", "normlog", "tobit", "cox", "weibull", "exponential", "zip", "beta", "median", "negbin", "longitudinal", "grouped", "qpois" and "qbinom". The "zip" means that the zero part is constant, the variables are not associated with the excessive zeros. The value "grouped" refers to grouped data, but this does not have to be given if the argument "group" is given, but not the argument "reps". The "binomial" is when you have the number of successes and also the number of trials. "MM" stands for robust regression using MM estimation.

<code>wei</code>	A vector of weights to be used for weighted regression. The default value is NULL.
<code>xnew</code>	If you have new data whose target values you want to predict put it here, otherwise leave it blank.

Details

This is a generic regression function, which offers prediction as well. It is important that you pass the arguments with their names, for example if you have time to event data, write "event = ..." and not just put your event variable. This will avoid confusion. For the mixed models you need to specify the relevant arguments, "slopes", "reps", "reml" and "group"

Value

A list including:

<code>mod</code>	The fitted model.
<code>pred</code>	If you have new data the predicted values of the target (dependent) variable.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

Almost the same as in [CondIndTests](#).

See Also

[fbedreg.bic](#), [mmpc.model](#), [ridge.reg](#)

Examples

```
#simulate a dataset with continuous data
dataset <- matrix(runif(100 * 10, 1, 100), nrow = 100 )
#the target feature is the last column of the dataset as a vector
target <- dataset[, 1]
dataset <- dataset[, -1]
a <- reg.fit(target, dataset)
```

Ridge regression *Ridge regression*

Description

Regularisation via ridge regression is performed.

Usage

```
ridge.reg(target, dataset, lambda, B = 1, newdata = NULL)
```

Arguments

target	A numeric vector containing the values of the target variable. If the values are proportions or percentages, i.e. strictly within 0 and 1 they are mapped into R using $\log(\text{target}/(1 - \text{target}))$.
dataset	A numeric matrix containing the variables. Rows are samples and columns are features.
lambda	The value of the regularisation parameter λ .
B	Number of bootstraps. If $B = 1$ no bootstrap is performed and no standard error for the regression coefficients is returned.
newdata	If you have new data and want to predict the value of the target put them here, otherwise, leave it NULL.

Details

There is also the `lm.ridge` command in MASS library if you are interested in ridge regression.

Value

A list including:

beta	The regression coefficients if no bootstrap is performed. If bootstrap is performed their standard error appears as well.
seb	The standard error of the regression coefficients. If bootstrap is performed their bootstrap estimated standard error appears.
est	The fitted values if no new data are available. If you have used new data these will be the predicted target values.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

- Hoerl A.E. and R.W. Kennard (1970). Ridge regression: Biased estimation for nonorthogonal problems. *Technometrics*, 12(1): 55-67.
- Brown P. J. (1994). *Measurement, Regression and Calibration*. Oxford Science Publications.

See Also

[ridgereg.cv](#)

Examples

```
#simulate a dataset with continuous data
dataset <- matrix(runif(100 * 50, 1, 100), nrow = 100 )
#the target feature is the last column of the dataset as a vector
target <- dataset[, 10]
dataset <- dataset[, -10]
a0 <- ridge.reg(target, dataset, lambda = 0, B = 1, newdata = NULL)
a1 <- ridge.reg(target, dataset, lambda = 0.5, B = 1, newdata = NULL)
a2 <- ridge.reg(target, dataset, lambda = 0.5, B = 100, newdata = NULL)
```

Ridge regression coefficients plot
Ridge regression

Description

A plot of the regularised parameters is shown.

Usage

```
ridge.plot(target, dataset, lambda = seq(0, 5, by = 0.1) )
```

Arguments

target	A numeric vector containing the values of the target variable. If the values are proportions or percentages, i.e. strictly within 0 and 1 they are mapped into R using $\log(\text{target}/(1 - \text{target}))$. In any case, they must be continuous only.
dataset	A numeric matrix containing the continuous variables. Rows are samples and columns are features.
lambda	A grid of values of the regularisation parameter λ .

Details

For every value of λ the coefficients are obtained. They are plotted versus the λ values.

Value

A plot with the values of the coefficients as a function of λ .

Author(s)

Michail Tsagris

R implementation and documentation: Giorgos Athineou <athineou@csd.uoc.gr>, Vincenzo Lagani <vlagani@csd.uoc.gr> and Michail Tsagris <mtsagris@csd.uoc.gr>

References

Hoerl A.E. and R.W. Kennard (1970). Ridge regression: Biased estimation for nonorthogonal problems. *Technometrics*, 12(1): 55-67.

Brown P. J. (1994). *Measurement, Regression and Calibration*. Oxford Science Publications.

See Also

[ridge.reg](#), [ridgereg.cv](#)

Examples

```
#simulate a dataset with continuous data
dataset <- matrix(runif(300 * 20, 1, 20), nrow = 300 )
#the target feature is the last column of the dataset as a vector
target <- dataset[, 20]
dataset <- dataset[, -20]
ridge.plot(target, dataset)
```

ROC and AUC

ROC and AUC

Description

Receiver operating curve and area under the curve.

Usage

```
auc(group, preds, roc = FALSE, cutoffs = NULL)
```

Arguments

group	A numerical vector with the predicted values of each group as 0 and 1.
preds	The predicted values of each group.
roc	If you want the ROC to appear set it to TRUE.
cutoffs	If you provide a vector with decreasing numbers from 1 to 0 that will be used for the ROC, otherwise, the values from 1 to 0 with a step equal to -0.01 will be used.

Details

The area under the curve is returned. The user has the option of getting the receiver operating curve as well.

Value

A list including:

cutoffs	The cutoff values.
sensitivity	The sensitivity values for each cutoff value.
specificity	The specificity value for each cutoff value.
youden	The pair of 1- specificity and sensitivity where the Youden's J appears on the graph and the Youden index which is defined as the maximum value of sensitivity - specificity + 1.
auc	The area under the curve, plus a circle with the point where Youden's J is located. If "roc" is set to FALSE, this is the only item in the list to be returned.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

See Also

[SES](#), [MMPC](#), [testIndLogistic](#)

Examples

```
# simulate a dataset with continuous data
g <- rbinom(150, 1, 0.6)
f <- rnorm(150)
auc(g, f, roc = FALSE)
```

Search for triangles in an undirected graph

Search for triangles in an undirected graph

Description

Search for triangles in an undirected graph.

Usage

```
triangles.search(G)
```

Arguments

G The adjacency matrix of an undirected graph. $G[i, j] = G[j, i] = 1$ means there is an edge between modes i and j . Zero values indicate the absence of edges.

Details

The functions searches for triangles, that is for tripletes of nodes (or variables) for which X-Y, Y-Z and X-Z.

Value

A matrix with thre columns. If there are no triangles, the matrix is empty. If there is at least one triangle, then each row contains three numbers, one for each node. See the examples below.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

See Also

[plotnetwork](#), [pc.skel](#), [mmhc.skel](#)

Examples

```
set.seed(123)
x <- rdag2(1000, p = 30, nei = 5)$x
a <- pc.skel(x, alpha = 0.05)
plotnetwork(a$G)
triangles.search(a$G)
```

SES.gee.output-class *Class* "SES.gee.output"

Description

SES.gee output object class.

Objects from the Class

Objects can be created by calls of the form `new("SES.temporal.output", ...)`.

Slots

`selectedVars`: Object of class "numeric"
`selectedVarsOrder`: Object of class "numeric"
`queues`: Object of class "list"
`signatures`: Object of class "matrix"
`hashObject`: Object of class "list"
`pvalues`: Object of class "numeric"
`stats`: Object of class "numeric"

univ: Object of class "list"
max_k: Object of class "numeric"
threshold: Object of class "numeric"
runtime: Object of class "proc_time"
test: Object of class "character"
correl: Object of class "character"
se: Object of class "character"

Methods

plot plot(x = "SES.gee.output", mode = "all"): Generic function for plotting the generated pvalues of the SES.temporal.output object. Argument mode = "all" for plotting all the pvalues or mode="partial" for partial plotting the first 500 pvalues

Author(s)

Giorgos Athineou <athineou@csd.uoc.gr>

See Also

[MMPC.gee](#), [SES.gee](#)

Examples

```
showClass("SES.temporal.output")
```

```
SES.temporal.output-class  
  Class "SES.temporal.output"
```

Description

SES.temporal output object class.

Objects from the Class

Objects can be created by calls of the form new("SES.temporal.output", ...).

Slots

selectedVars: Object of class "numeric"
selectedVarsOrder: Object of class "numeric"
queues: Object of class "list"
signatures: Object of class "matrix"
hashObject: Object of class "list"

pvalues: Object of class "numeric"
stats: Object of class "numeric"
univ: Object of class "list"
max_k: Object of class "numeric"
threshold: Object of class "numeric"
runtime: Object of class "proc_time"
test: Object of class "character"
slope: Object of class "logical"

Methods

plot plot(x = "SES.temporal.output", mode = "all"): Generic function for plotting the generated pvalues of the SES.temporal.output object. Argument mode = "all" for plotting all the pvalues or mode="partial" for partial plotting the first 500 pvalues

Author(s)

Giorgos Athineou <athineou@csd.uoc.gr>

See Also

[MMPC.temporal](#), [SES.temporal](#)

Examples

```
showClass("SES.temporal.output")
```

SESoutput-class	<i>Class "SESoutput"</i>
-----------------	--------------------------

Description

SES output object class.

Objects from the Class

Objects can be created by calls of the form `new("SESoutput", ...)`.

Slots

selectedVars: Object of class "numeric"
selectedVarsOrder: Object of class "numeric"
queues: Object of class "list"
signatures: Object of class "matrix"
hashObject: Object of class "list"
pvalues: Object of class "numeric"
stats: Object of class "numeric"
univ: Object of class "list"
max_k: Object of class "numeric"
threshold: Object of class "numeric"
n.tests: Object of class "numeric"
runtime: Object of class "proc_time"
test: Object of class "character"

Methods

plot plot(x = "SESoutput", mode = "all"): Generic function for plotting the generated pvalues of the SESoutput object. Argument mode = "all" for plotting all the pvalues or mode="partial" for partial plotting the first 500 pvalues

Author(s)

Giorgos Athineou <athineou@csd.uoc.gr>

See Also

[SES](#)

Examples

```
showClass("SESoutput")
```

Skeleton (local) around a node of the max-min hill-climbing (MMHC) algorithm
Skeleton (local) around a node of the MMHC algorithm

Description

The local skeleton of a Bayesian network around a node produced by MMHC. No orientations are involved.

Usage

```
local.mmhc.skel(dataset, node, max_k = 3, threshold = 0.05, test = "testIndFisher")
```

Arguments

dataset	A matrix with the variables. The user must know if they are continuous or if they are categorical. If you have a matrix with categorical data, i.e. 0, 1, 2, 3 where each number indicates a category, the minimum number for each variable must be 0. data.frame is also supported, as the dataset in this case is converted into a matrix.
node	A number between 1 and the number of columns of the dataset. The local network (edges only) will be built around this node. At first the parents and children of this node are identified and then their parents and children. No inconsistencies correction whatsoever is attempted. A variable detected by the node, but the node was not detected by that variable.
max_k	The maximum conditioning set to use in the conditional independence test (see Details of SES or MMPC).
threshold	Threshold (suitable values in (0, 1)) for assessing p-values significance. Default value is 0.05.
test	The conditional independence test to use. Default value is "testIndFisher". This procedure allows for "testIndFisher", "testIndSPearman" for continuous variables and "gSquare" for categorical variables. Or in general, if the dataset is a data.frame with different types of data, leave this NULL. See also link{MMPC} for the automatic choice of tests.

Details

The MMPC is run on the user specific variable. The backward phase (see Tsamardinos et al., 2006) takes place automatically. Then, the MMPC is run on the parents and children of that variable. If the node variable is not detected by a variable, this variable is not removed though.

Value

A list including:

runtime	The run time of the algorithm. A numeric vector. The first element is the user time, the second element is the system time and the third element is the elapsed time.
ntests	The number of tests MMPC (or SES) performed at each variable.
res	A list with the parents and children of each variable. The first element is the parents and children of the node variable.
Gloc	The local adjacency matrix. A value of 1 in $G[i, j]$ may not appear appear in $G[j, i]$ also, indicating that variable j was discovered as a possible parent or child of node i , but not the converse. The usual MMHC (mmhc.skel) removes the edge between them as this is an inconsistency.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

Tsamardinos, Brown and Aliferis (2006). The max-min hill-climbing Bayesian network structure learning algorithm. *Machine learning*, 65(1), 31-78.

See Also

[mmhc.skel](#), [pc.skel](#), [pc.or](#), [corfs.network](#)

Examples

```
# simulate a dataset with continuous data
dataset <- matrix(runif(1000 * 30, 1, 100), nrow = 1000 )
a1 <- mmhc.skel(dataset, max_k = 3, threshold = 0.05, test = "testIndFisher")
a2 <- local.mmhc.skel(dataset, 4)
a1$runtime
a2$runtime

dataset <- rdag2(1000, p = 20, nei = 4)$x
a1 <- mmhc.skel(dataset, max_k = 3, threshold = 0.05, test = "testIndFisher")
a2 <- local.mmhc.skel(dataset, 5)
a1$runtime
a2$runtime
```

Skeleton of the max-min hill-climbing (MMHC) algorithm

The skeleton of a Bayesian network as produced by MMHC

Description

The skeleton of a Bayesian network produced by MMHC. No orientations are involved.

Usage

```
mmhc.skel(dataset, max_k = 3, threshold = 0.05, test = "testIndFisher", type = "MMPC",
backward = TRUE, symmetry = TRUE, nc = 1, ini.pvalue = NULL, hash = FALSE)
```

Arguments

dataset	A matrix with the variables. The user must know if they are continuous or if they are categorical. If you have a matrix with categorical data, i.e. 0, 1, 2, 3 where each number indicates a category, the minimum number for each variable must be 0. data.frame is also supported, as the dataset in this case is converted into a matrix.
max_k	The maximum conditioning set to use in the conditional independence test (see Details of SES or MMPC).
threshold	Threshold (suitable values in (0, 1)) for assessing p-values significance. Default value is 0.05.
test	The conditional independence test to use. Default value is "testIndFisher". This procedure allows for "testIndFisher", "testIndSPearman" for continuous variables and "gSquare" for categorical variables. In case the dataset is a dta.frame with mixed types of data leave this NULL and an appropriate test will be selected. See <code>link{MMPC}</code> for the automatic choice of tests.
type	The type of variable selection to take place for each variable (or node). The default (and standard) in "MMPC". You can also choose to run it via "SES" and thus allow for multiple signatures of variables to be connected to a variable.
backward	If TRUE, the backward (or symmetry correction) phase will be implemented. This removes any falsely included variables in the parents and children set of the target variable. It call the <code>link{mmpcbackphase}</code> for this purpose. For <code>perm.ses</code> and <code>wald.ses</code> this is not yet applicable.
symmetry	In order for an edge to be added, a statistical relationship must have been found from both directions. If you want this symmetry correction to take place, leave this boolean variable to TRUE. If you set it to FALSE, then if a relationship between Y and X is detected but not between X and Y, the edge is still added.
nc	How many cores to use. This plays an important role if you have many variables, say thousands or so. You can try with <code>nc = 1</code> and with <code>nc = 4</code> for example to see the differences. If you have a multicore machine, this is a must option. There was an extra argument for plotting the skeleton but it does not work with the current visualisation packages, hence we removed the argument. Use <code>plotnetwork</code> to plot the skeleton.
ini.pvalue	This is a list with the matrix of the univariate p-values. If you want to run <code>mmhc.skel</code> again, the univariate associations need not be calculated again.
hash	A boolean variable which indicates whether (TRUE) or not (FALSE) to store the statistics calculated during MMPC or SES execution in a hash-type object. Default value is FALSE. If TRUE a hashObject is produced. You should put it to TRUE is you want an accurate count of the the number of tests performed. The only reason I put FLASE is because I want it to pass the CRAN check.

Details

The MMPC is run on every variable. The backward phase (see Tsamardinos et al., 2006) takes place automatically. After all variables have been used, the matrix is checked for inconsistencies and they are corrected.

A trick mentioned in that paper to make the procedure faster is the following. In the k -th variable, the algorithm checks how many previously scanned variables have an edge with this variable and keeps them (it discards the other variables with no edge) along with the next (unscanned) variables. This trick reduces time, but can lead to different results. For example, if the i -th variable is removed, the k -th node might not remove an edge between the j -th variable, simply because the i -th variable that could separate them is missing.

The user is given this option via the argument "fast", which can be either TRUE or FALSE. Parallel computation is also available.

Value

A list including:

runtime	The run time of the algorithm. A numeric vector. The first element is the user time, the second element is the system time and the third element is the elapsed time.
density	The number of edges divided by the total possible number of edges, that is $\#edges / n(n - 1)/2$, where n is the number of variables.
info	Some summary statistics about the edges, minimum, maximum, mean, median number of edges.
ms	If you run "MMPC" for each variable this is NULL. If you run "SES" is a vector denoting which variables had more than one signature, i.e. more than one set of variables associated with them.
ntests	The number of tests MMPC (or SES) performed at each variable.
ini.pvalue	A matrix with the p-values of all pairwise univariate associations.
pvalue	A matrix with the final p-values. These are the maximum p-values calculated during the process. When the process finishes, the matrix is not symmetric. It becomes symmetric though by keeping the maximum between any two off-diagonal elements. These p-values now can be used to sort the strength of the edges. If you know the true adjacency matrix you can use them and create a ROC curve (see <code>link{bn.skel.utils}</code> for more information).
G	The adjacency matrix. A value of 1 in $G[i, j]$ appears in $G[j, i]$ also, indicating that i and j have an edge between them.

Bear in mind that the values can be extracted with the $\$$ symbol, i.e. this is an S3 class output.

Author(s)

Michail Tsagris

R implementation and documentation: Giorgos Athineou <athineou@csd.uoc.gr> and Michail Tsagris <mtsagris@csd.uoc.gr>

References

Tsamardinos, Brown and Aliferis (2006). The max-min hill-climbing Bayesian network structure learning algorithm. *Machine learning*, 65(1), 31-78.

Brown L. E., Tsamardinos I., and Aliferis C. F. (2004). A novel algorithm for scalable and accurate Bayesian network learning. *Medinfo*, 711-715.

See Also

[pc.skel](#), [pc.or](#), [corfs.network](#), [bn.skel.utils](#)

Examples

```
# simulate a dataset with continuous data
y <- rdag2(500, p = 20, nei = 3)
x <- y$x
a <- mmhc.skel(x, max_k = 5, threshold = 0.01, test = "testIndFisher" )
b <- pc.skel( x, alpha = 0.01 )
a$runtime ##
b$runtime ##
```

Skeleton of the PC algorithm

The skeleton of a Bayesian network produced by the PC algorithm

Description

The skeleton of a Bayesian network produced by the PC algorithm. No orientations are involved. The `pc.con` is for continuous data only and it calls the same C++ as `pc.skel`. Hence, you are advised to use `pc.skel`.

Usage

```
pc.skel(dataset, method = "pearson", alpha = 0.01, rob = FALSE, R = 1, stat = NULL,
ini.pvalue = NULL)
```

```
pc.con(dataset, method = "pearson", alpha = 0.01)
```

```
pc.skel.boot(dataset, method = "pearson", alpha = 0.01, R = 199, ncores = 1)
```

```
glmm.pc.skel(dataset, id, method = "comb.mm", alpha = 0.01, stat = NULL,
ini.pvalue = NULL)
```

Arguments

`dataset` A matrix with the variables. The user must know if they are continuous or if they are categorical. **If you have categorical data though, the user must transform the data.frame into a matrix. In addition, the numerical matrix must have values starting from 0. For example, 0, 1, 2, instead of "A", "B" and "C".** In the case of mixed variables, continuous, binary and ordinal this must a data.frame and the non continuous variables must be ordered factors, even the binary variables.

For the `pc.con`, `pc.skel.boot` and `glmm.pc.skel`, `dataset` can only be a matrix.

method	If you have continuous data, you can choose either "pearson", "spearman" or "distcor". The latter uses the distance correlation and should not be used with lots of observations as it is by default really slow. If you have categorical data, this must be "cat". If you have a mix of continuous, binary and ordinal data (we will expand the available dataset in the future) then choose "comb.fast" or "comb.mm". These two methods perform the symmetric test for mixed data (Tsagris et al., 2017). See details for more information on this. For the mixed models PC algorithm, this is the same argument, but currently only "comb.mm" is accepted.
id	This is to be used in the <code>glimm.pc.skel</code> only. It is a vector for identifying the grouped data, the correlated observations, the subjects.
alpha	The significance level (suitable values in (0, 1)) for assessing the p-values. Default value is 0.01.
rob	This is for robust estimation of the Pearson correlation coefficient. Default value is FALSE.
R	The number of permutations to be conducted. This is taken into consideration for the "pc.skel" only. The Pearson correlation coefficient is calculated and the p-value is assessed via permutations. There was an extra argument for plotting the skeleton but it does not work with the current visualisation packages, hence we removed the argument. Use <code>plotnetwork</code> to plot the skeleton. In the <code>pc.skel.boot</code> this is the number of bootstrap resamples to draw. The PC algorithm is performed in each bootstrap sample. In the end, the adjacency matrix on the observed data is returned, along with another adjacency matrix produced by the bootstrap. The latter one contains values from 0 to 1 indicating the proportion of times an edge between two nodes was present.
ncores	The number of cores to use. By default this is set to 1.
stat	If you have the initial test statistics (univariate associations) values supply them here.
ini.pvalue	If you have the initial p-values of the univariate associations supply them here.

Details

The PC algorithm as proposed by Spirtes et al. (2000) is implemented. The variables must be either continuous or categorical, only. The skeleton of the PC algorithm is order independent, since we are using the third heuristic (Spirtes et al., 2000, pg. 90). At every step of the algorithm use the pairs which are least statistically associated. The conditioning set consists of variables which are most statistically associated with each either of the pair of variables.

For example, for the pair (X, Y) there can be two conditioning sets for example (Z1, Z2) and (W1, W2). All p-values and test statistics and degrees of freedom have been computed at the first step of the algorithm. Take the p-values between (Z1, Z2) and (X, Y) and between (Z1, Z2) and (X, Y). The conditioning set with the minimum p-value is used first. If the minimum p-values are the same, use the second lowest p-value. In the event of 2 or more p-values being the same (with permutations for example), the test statistic divided by the degrees of freedom is used as a means of choosing which conditioning set is to be used first. If two or more p-values are below the machine epsilon (`.Machine$double.eps` which is equal to $2.220446e-16$), all of them are set to 0. To make the comparison or the ordering feasible we use the logarithm of the p-value. Hence, the logarithm of the p-values is always calculated and used.

In the case of the G^2 test of independence (for categorical data) we have incorporated a rule of thumb. If the number of samples is at least 5 times the number of the parameters to be estimated, the test is performed, otherwise, independence is not rejected (see Tsamardinos et al., 2006).

The "comb.fast" and "comb.mm" methods are used with mixed variables, continuous, binary and ordinal. The "comb.mm" performs two log-likelihood ratio tests. For every pair of variables each of the two variables is treated as response and the suitable regression model is fitted. Then, two likelihood ratio tests are performed and the 2 p-values are combined in a meta-analytic way. In the case of "comb.fast" one regression model is fitted, the easiest (between the two) to implement. The ordering of the "easiness" is as follows: linear regression > logistic regression > ordinal regression.

The "pc.con" is a faster implementation of the PC algorithm but for continuous data only, without the robust option, unlike pc.skel which is more general and even for the continuous datasets slower. pc.con accepts only "pearson" and "spearman" as correlations.

If there are missing values they are placed by their median in case of continuous data and by their mode (most frequent value) if they are categorical.

The "lmm.pc.skel" is designed for clustered or grouped data. It uses linear mixed models and works in the same way as the PC with mixed data. For each variable, a random intercepts model is fitted and the significance of the other variable is assessed. The two p-values are meta-analytically combined.

For all cases, except for method = c("pearson", "spearman", "cat") with $R = 1$, or $R > 1$, we return the maximum logged p-value of the conditional independence tests between the pairs. This can be used to order the strength of association between pairs of variables. In addition, one can use it to estimate the AUC. See the example.

Value

A list including:

stat	The test statistics of the univariate associations.
ini.pvalue	The initial univariate associations p-values.
pvalue	The logarithm of the maximum p-values from every conditional independence test. Note also, that if there is a log p-value smaller than the log of the threshold value that means there is an edge. This can be used to estimate the FDR (Tsamardinos and Brown, 2008). See details for more information. This is also an estimate of the strength of the association between pairs of variables. At the moment this is not possible for method = c("pearson", "spearman", "cat") with $R = 1$, or $R > 1$ because these cases are handled in C++. We will add those cases in the future.
info	Some summary statistics about the edges, minimum, maximum, mean, median number of edges.
runtime	The run time of the algorithm. A numeric vector. The first element is the user time, the second element is the system time and the third element is the elapsed time.
kappa	The maximum value of k, the maximum cardinality of the conditioning set at which the algorithm stopped.
density	The number of edges divided by the total possible number of edges, that is $\#edges / n(n - 1)/2$, where n is the number of variables.

info	Some summary statistics about the edges, minimum, maximum, mean, median number of edges.
G	The adjacency matrix. A value of 1 in G[i, j] appears in G[j, i] also, indicating that i and j have an edge between them.
sepset	A list with the separating sets for every value of k.
title	The name of the dataset.

Bear in mind that the values can be extracted with the \$ symbol, i.e. this is an S3 class output.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

Spirtes P., Glymour C. and Scheines R. (2001). Causation, Prediction, and Search. The MIT Press, Cambridge, MA, USA, 3rd edition.

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Tsagris M., Borboudakis G., Lagani V. and Tsamardinos I. (2018). Constraint-based Causal Discovery with Mixed Data. International Journal of Data Science and Analytics.

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Sedgewick, A. J., Ramsey, J. D., Spirtes, P., Glymour, C., & Benos, P. V. (2017). Mixed Graphical Models for Causal Analysis of Multi-modal Variables. arXiv preprint arXiv:1704.02621.

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See Also

[bn.skel.utils](#), [mmhc.skel](#), [corfs.network](#), [local.mmhc.skel](#)

Examples

```
# simulate a dataset with continuous data
y <- rdag2(500, p = 20, nei = 3)
ind <- sample(1:20, 20)
x <- y$x[, ind]
a <- mmhc.skel(x, max_k = 3, threshold = 0.05, test = "testIndFisher" )
b <- pc.skel( x, method = "pearson", alpha = 0.05 )
a$runtime ##
b$runtime ##
```

Structural Hamming distance between two partially oriented DAGs
Structural Hamming distance between two partially oriented DAGs

Description

Structural Hamming distance between two partially oriented DAGs.

Usage

```
shd(est, true)
```

Arguments

`est` The first (partially oriented) DAG. This could also be the estimated DAG.
`true` The second (partially oriented) DAG. This could also be the equivalence class of the true DAG.

Details

The structural Hamming distance as proposed by Tsamardinos et al. (2006) is calculated and returned. The cases are listed below

True	Estimated	Penalty
-		1
	-	1
->		1
	<-	1
->	-	1
-	<-	1
->	<-	1

Value

A list including

`mat` A table with the agreements and disagreements between the two DAGs.
`shd` The structural Hamming distance.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

Tsamardinos, Brown and Aliferis (2006). The max-min hill-climbing Bayesian network structure learning algorithm. *Machine learning*, 65(1), 31-78.

See Also

[pc.skel](#), [pc.or](#), [mmhc.skel](#), [plotnetwork](#)

Examples

```
y <- rdag(1000, 20, 0.2)
tru <- y$G
mod <- pc.skel(y$x)
a <- pc.or(mod)
shd( a$G, dag2eg(tru) )
```

Symmetric conditional independence test with clustered data

Symmetric conditional independence test with clustered data

Description

Symmetric conditional independence test with clustered data.

Usage

```
glm.ci.mm(ind1, ind2, cs = NULL, dat, id)
```

Arguments

<code>ind1</code>	The index of the one variable to be considered.
<code>ind2</code>	The index of the other variable to be considered.
<code>cs</code>	The index or indices of the conditioning set of variable(s). If you have no variables set this equal to 0.
<code>dat</code>	A numerical matrix with data.
<code>id</code>	This is a numerical vector denoting the groups or the subjects.

Details

Two linear random intercept models are fitted, one for each variable and the p-value of the hypothesis test that the other variable is significant is calculated. These two p-values are combined in a meta-analytic way.

Value

A vector including the test statistic, it's associated p-value and the relevant degrees of freedom.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

Tsagris M., Borboudakis G., Lagani V. and Tsamardinos I. (2018). Constraint-based Causal Discovery with Mixed Data. International Journal of Data Science and Analytics.

<https://link.springer.com/content/pdf/10.1007>

<http://nugget.unisa.edu.au/CD2017/papersonly/constraint-based-causal-r1.pdf>

Eugene Demidenko (2013). Mixed Models: Theory and Applications with R, 2nd Edition. New Jersey: Wiley & Sons.

See Also

[pc.skel](#), [condi](#), [testIndGLMMReg](#)

Examples

```
## we generate two independent vectors of clustered data
s1 <- matrix(1.5, 4, 4)
diag(s1) <- 2.5
s2 <- matrix(1.5, 4, 4)
diag(s2) <- 2
x1 <- MASS::mvrnorm(10, rnorm(4), s1)
x1 <- as.vector( t(x1) )
x2 <- MASS::mvrnorm(10, rnorm(4), s2)
x2 <- as.vector( t(x2) )
id <- rep(1:10, each = 4)
glmm.ci.mm(1, 2, dat = cbind(x1,x2), id = id)
```

Symmetric conditional independence test with mixed data

Symmetric conditional independence test with mixed data

Description

Symmetric conditional independence test with mixed data.

Usage

```
ci.mm(ind1, ind2, cs = NULL, dat, type, rob = FALSE, R = 1)
ci.fast(ind1, ind2, cs = NULL, dat, type, rob = FALSE, R = 1)
ci.mm2(ind1, ind2, cs = NULL, suffStat)
ci.fast2(ind1, ind2, cs = NULL, suffStat)
```

Arguments

ind1	The index of the one variable to be considered.
ind2	The index of the other variable to be considered.
cs	The index or indices of the conditioning set of variable(s). If you have no variables set this equal to 0.
dat	A data.frame with numerical, binary, nominal and ordinal variables only.
type	This is obsolete basically, but we need it here, so that the functions <code>ci.mm</code> and <code>ci.fast</code> have the same signatures as in <code>cat.ci</code> , <code>condi</code> and <code>dist.condi</code> .
rob	This is obsolete basically, but we need it here, so that the functions <code>ci.mm</code> and <code>ci.fast</code> have the same signatures as in <code>cat.ci</code> , <code>condi</code> and <code>dist.condi</code> .
R	This is obsolete basically, but we need it here, so that the functions <code>ci.mm</code> and <code>ci.fast</code> have the same signatures as in <code>cat.ci</code> , <code>condi</code> and <code>dist.condi</code> .
suffStat	This is a list with only the dataset and the name must be "dataset". A data.frame with numerical, binary, nominal and ordinal variables only.

Details

The functions "ci.mm" and "ci.fast" are general functions to be used anywhere. The functions "ci.mm2" and "ci.fast2" are designed to be accepted by the command "pc" in the package "pcalg". The functions "ci.mm2" and "ci.fast2" can be fed in the "pc" function of the "pcalg" package in order to produce a PDAG with mixed data using the PC algorithm. For more information see the relevant paper in the references.

The "ci.mm" and "ci.fast" work with linear, logistic, multinomial and ordinal regression, whereas the "ci.mm2" and "ci.fast2" work with linear, logistic and ordinal regression only.

Value

A vector including the test statistic, it's associated p-value and the relevant degrees of freedom.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

- Tsagris M., Borboudakis G., Lagani V. and Tsamardinos I. (2018). Constraint-based Causal Discovery with Mixed Data. *International Journal of Data Science and Analytics*.
<https://link.springer.com/content/pdf/10.1007>
<http://nugget.unisa.edu.au/CD2017/papersonly/constraint-based-causal-r1.pdf>
- Spirtes P., Glymour C. and Scheines R. (2001). *Causation, Prediction, and Search*. The MIT Press, Cambridge, MA, USA, 3rd edition.
- Sedgewick, A. J., Ramsey, J. D., Spirtes, P., Glymour, C., & Benos, P. V. (2017). *Mixed Graphical Models for Causal Analysis of Multi-modal Variables*. arXiv preprint arXiv:1704.02621.

See Also

[pc.skel](#), [mmhc.skel](#), [cat.ci](#), [condi](#)

Examples

```
ci.mm(1, 2, dat = iris)
ci.mm(1, 5, dat = iris)
ci.fast(1, 5, dat = iris)
x <- iris
x[, 5] <- as.numeric(x[, 5]) ## Caution:: this will be treated as ordered variable.
x[, 5] <- factor(x[, 5], ordered = TRUE)
## ci.mm2 and ci.fast2 do not perform multinomial regression.
ci.mm2(1, 5, suffStat = list(dataset = x) )
ci.fast2(1, 5, suffStat = list(dataset = x) )
```

The max-min Markov blanket algorithm

Max-min Markov blanket algorithm

Description

The MMMB algorithm follows a forward-backward filter approach for feature selection in order to provide a minimal, highly-predictive, feature subset of a high dimensional dataset. See also Details.

Usage

```
mmmb(target , dataset , max_k = 3 , threshold = 0.05 , test = "testIndFisher",
user_test = NULL, ncores = 1)
```

Arguments

<code>target</code>	The class variable. Provide either a string, an integer, a numeric value, a vector, a factor, an ordered factor or a Surv object. See also Details.
<code>dataset</code>	The dataset; provide either a data frame or a matrix (columns = variables, rows = samples). In either case, only two cases are available, either all data are continuous, or categorical.
<code>max_k</code>	The maximum conditioning set to use in the conditional independence test (see Details). Integer, default value is 3.
<code>threshold</code>	Threshold (suitable values in [0,1]) for assessing the p-values. Default value is 0.05.
<code>test</code>	The conditional independence test to use. Default value is "testIndFisher". See also link{CondIndTests} .
<code>user_test</code>	A user-defined conditional independence test (provide a closure type object). Default value is NULL. If this is defined, the "test" argument is ignored.

`ncores` How many cores to use. This plays an important role if you have tens of thousands of variables or really large sample sizes and tens of thousands of variables and a regression based test which requires numerical optimisation. In other cases it will not make a difference in the overall time (in fact it can be slower). The parallel computation is used in the first step of the algorithm, where univariate associations are examined, those take place in parallel. We have seen a reduction in time of 50% with 4 cores in comparison to 1 core. Note also, that the amount of reduction is not linear in the number of cores.

Details

The idea is to run the MMPC algorithm at first and identify the parents and children (PC) of the target variable. As a second step, the MMPC algorithm is run on the discovered variables to return PC. The parents of the children of the target are the spouses of the target. Every variable in PC_i is checked to see if it is a spouse of the target. If yes, it is included in the Markov Blanket of the target, otherwise it is thrown. If the data are continuous, the Fisher correlation test is used or the Spearman correlation (more robust). If the data are categorical, the G^2 test is used.

Value

The output of the algorithm is S3 object including:

`mb` The Markov Blanket of the target variable. The parents and children of the target variable, along with the spouses of the target, which are the parents of the children of the target variable.

`runtime` The run time of the algorithm. A numeric vector. The first element is the user time, the second element is the system time and the third element is the elapsed time.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

Tsamardinos I., Aliferis C. F. and Statnikov, A. (2003). Time and sample efficient discovery of Markov blankets and direct causal relations. In Proceedings of the 9th ACM SIGKDD international conference on Knowledge discovery and data mining, pp. 673-678.

See Also

[CondIndTests](#), [MMPC](#), [SES](#)

Examples

```
set.seed(123)
require(hash)

#simulate a dataset with continuous data
```

```
dataset <- matrix( runif(1000 * 50, 1, 100), ncol = 50 )
#define a simulated class variable
target <- 3 * dataset[, 10] + 2 * dataset[, 50] + 3 * dataset[, 20] + rnorm(1000, 0, 5)
aa <- mmb(target , dataset , max_k = 3 , threshold = 0.05, test= "testIndFisher",
ncores = 1,)
ab <- SES(target, dataset, test="testIndFisher")
```

Topological sort of a DAG

Topological sort of a DAG

Description

Topological sort of a DAG.

Usage

```
topological_sort(dag)
```

Arguments

`dag` A square matrix representing a directed graph which contains 0s and 1s. If $G[i, j] = 1$ it means there is an arrow from node i to node j . When there is no edge between nodes i and j if $G[i, j] = 0$.

Details

The function is an R translation from an old matlab code.

Value

A vector with numbers indicating the sorting. If the matrix does not correspond to a DAG, NA will be returned.

Author(s)

Ioannis Tsamardinos and Michail Tsagris

R implementation and documentation: Ioannis Tsamardinos <tsamard@csd.uoc.gr> and Michail Tsagris <mtsagris@csd.uoc.gr>

References

Chickering, D.M. (1995). A transformational characterization of equivalent Bayesian network structures. Proceedings of the 11th Conference on Uncertainty in Artificial Intelligence, Montreal, Canada, 87-98.

See Also

[plotnetwork](#), [nei](#), [pc.or](#)

Examples

```
# simulate a dataset with continuous data
# simulate a dataset with continuous data
G = rdag(1000, 10, 0.3)$G
G[G == 2] <- 1
G[G == 3] <- 0
topological_sort(G)
```

Total causal effect of a node on another node

Total causal effect of a node on another node

Description

Total causal effect of a node on another node.

Usage

```
ida(x, y, G, dataset)
```

Arguments

x	A number between 1 and the number of variables (nodes) indicating the node whose total causal effect we want to estimate. This is the independent variable. See details for more on this.
y	A number between 1 and the number of variables (nodes) indicating the node who is the dependent variable. The goal is to estimate the total causal effect of x on y.
G	A square matrix representing a (partially) directed graph which contains 0s and 1s. If $G[i, j] = 2$ it means there is an arrow from node i to node j. If $G[i, j] = 1$, there is an undirected edge between nodes i and j and there is no edge between nodes i and j if $G[i, j] = 0$.
dataset	The dataset. This is a numerical matrix with data.

Details

The total causal effect defined in Pearl's do-calculus is $E(Y|do(X=z+1)) - E(Y|do(X=z))$. As Pearl described it, he used linear regression, hence this function works for continuous data which are assumed to be Gaussian.

We estimate a set of possible total causal effects using linear regression. If y is a parent, or a descendant, of x in G, the estimated causal effect of x on y is zero. If y is not a parent of x, we take the regression coefficient of x in the regression $lm(y \sim x + pa(x))$, where $pa(x)$ denotes the parents of x. This is repeated using all parents of x (including the empty set) and all possible parents values of x and their combinations.

One restriction to bear in mind. If a collider is created that combination of nodes is not used in the regression.

Value

A list including:

tc	A matrix with 4 elements. The first column is the estimated beta coefficient, the second is its standard error, its t-value and the p-value for testing whether this is equal to 0.
mess	If the x node has no parents a message about this appears. Otherwise this is NULL.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

M.H. Maathuis, M. Kalisch and P. Buehlmann (2009). Estimating high-dimensional intervention effects from observational data. *Annals of Statistics* 37, 3133-3164.

Pearl (2005). *Causality. Models, reasoning and inference*. Cambridge University Press, New York.

See Also

[plotnetwork](#), [nei](#), [pc.or](#)

Examples

```
dataset <- rdag2(1000, p=20, nei = 3)$x
mod <- pc.skel(dataset, alpha = 0.01)
G <- pc.or(mod)$G
ida(10, 15, G, dataset)
```

Transformation of a DAG into an essential graph

Transforms a DAG into an essential graph

Description

Transforms a DAG into an essential graph.

Usage

```
dag2eg(dag, type = NULL)
```

Arguments

dag	The graph matrix as produced from pc.or or any other algorithm which produces directed graphs. A DAG in general.
type	This can be either NULL or 1 or 2. type = 1 means that the matrix contains 0, 1, 2, 3 where $G[i, j] = g[j, i] = 0$, means there is no edge between nodes i and j, $G[i, j] = g[j, i] = 1$, there is an edge between nodes i and j and $G[i, j] = 2$ and $G[j, i] = 3$ means that there is an arrow from node i to node j. If type 2, the matrix contains 0 for no edge and 1 for a directed edge. In this case, $G[i,j]=1$ and $G[j,i]=0$ means that there is an arrow from node i to node j. If you are not sure of what you have, just leave it NULL, the function will check to which case your matrix belongs.

Details

The function is an R translation from an old matlab code.

Value

The matrix of the essential graph.

Author(s)

Ioannis Tsamardinos and Michail Tsagris

R implementation and documentation: Ioannis Tsamardinos <tsamard@csd.uoc.gr> and Michail Tsagris <mtsagris@csd.uoc.gr>

References

Chickering, D.M. (1995). A transformational characterization of equivalent Bayesian network structures. Proceedings of the 11th Conference on Uncertainty in Artificial Intelligence, Montreal, Canada, 87-98.

See Also

[plotnetwork](#), [is.dag](#), [topological_sort](#)

Examples

```
# simulate a dataset with continuous data
# simulate a dataset with continuous data
y = rdag(1000, 10, 0.3)
tru = y$G
eg = dag2eg(tru)
par( mfrow = c(1, 2) )
plotnetwork(tru)
plotnetwork(eg)
```

Transitive closure of an adjacency matrix

Returns the transitive closure of an adjacency matrix

Description

Returns the transitive closure of an adjacency matrix.

Usage

```
transitiveClosure(amat)
```

Arguments

amat The adjacency matrix of a graph.

Details

A function that computes the transitive closure of a graph. The transitive closure $C(G)$ of a graph is a graph which contains an edge between nodes u and v whenever there is a directed path from u to v (Skiena 1990, p. 203). <http://mathworld.wolfram.com/TransitiveClosure.html>

Value

closure The transitive closure of the adjacency matrix representing a graph.

Author(s)

Anna Roumpelaki

R implementation and documentation: Anna Roumpelaki <anna.roumpelaki@gmail.com>

References

Skiena S. (1990). Implementing Discrete Mathematics: Combinatorics and Graph Theory with Mathematica. Reading, MA: Addison-Wesley

Examples

```
# example adjacency matrix
# simulate a dataset with continuous data
dataset <- matrix( runif(1000 * 10, 1, 100), nrow = 1000 )
test <- pc.con( dataset, method = "pearson", alpha = 0.05 )$G
transitiveClosure(test)
```

Undirected path(s) between two nodes

Undirected path(s) between two nodes

Description

Undirected path(s) between two nodes.

Usage

```
undir.path(G, y, x)
```

Arguments

G	An adjacency matrix where $G[i,j] = G[j, i] = 1$ means there is an edge between nodes i and j . If $G[i, j] = G[j, i] = 0$ there is no edge between them.
y	A numerical value indicating the first node, it has to be a number between 1 and the maximum number of variables.
x	A numerical value indicating the second node, it has to be a number between 1 and the maximum number of variables. The order of the nodes does not make a difference.

Details

The algorithm finds all the nodes between the two nodes. It finds all paths between the two chosen nodes.

Value

A vector with the two nodes and all nodes between them in the case of connecting nodes. Otherwise, a matrix with the neighbours of each node.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

Tsamardinos, Brown and Aliferis (2006). The max-min hill-climbing Bayesian network structure learning algorithm. *Machine learning*, 65(1), 31-78.

See Also

[SES](#), [MMPC](#), [pc.skel](#)

Examples

```
# simulate a dataset with continuous data
set.seed(1234)
dataset <- matrix(runif(1000 * 20, 1, 100), nrow = 1000 )
G <- pc.con(dataset)$G
plotnetwork(G)
undir.path(G, 3, 4)
undir.path(G, 1, 3)
```

Univariate regression based tests

Univariate regression based tests

Description

Univariate regression based tests.

Usage

```
univregs(target, dataset, targetID = -1, test = NULL, user_test = NULL, wei = NULL,
ncores = 1)
```

```
ebic.univregs(target, dataset, targetID = -1, test = NULL, user_test = NULL,
wei = NULL, ncores = 1, gam = NULL)
```

```
wald.univregs(target, dataset, targetID = - 1, test = NULL, user_test = NULL,
wei = NULL, ncores = 1)
```

```
perm.univregs(target, dataset, targetID = -1, test = NULL, user_test = NULL,
wei = NULL, threshold = 0.05, R = 999, ncores = 1)
```

```
score.univregs(target, dataset, test)
```

```
rint.regs(target, dataset, targetID = -1, id, reps = NULL, tol = 1e-08)
```

```
glmm.gee.univregs(target, reps = NULL, id, dataset, targetID = -1, test, wei = NULL,
slopes = FALSE, correl = "exchangeable", se = "jack", ncores = 1)
```

Arguments

target	The target (dependent) variable. It must be a numerical vector.
dataset	The indendent variable(s). For the "univregs" this can be a matrix or a dataframe with continuous only variables, a data frame with mixed or only categorical variables. For the "wald.univregs", "perm.univregs", "score.univregs" and "rint.regs" this can only by a numerical matrix.
targetID	This is by default -1. If the target is not a variable but is included in the dataset, you can specify the column of dataset playing the role of the target.

test	<p>For the "univregs" this can only be one of the following: testIndFisher, testIndSpearman, gSquare, testIndBeta, testIndReg, testIndLogistic, testIndMultinom, testIndOrdinal, testIndPois, testIndZIP, testIndNB, testIndClogit, testIndBinom, testIndIGreg, censIndCR, censIndWR, censIndWR, testIndGamma or testIndNormLog. For the testIndSpearman the user must supply the ranked target and ranked dataset. The reason for this, is because this function is called internally by SES and MMPC, the ranks have already been applied and there is no reason to re-rank the data.</p> <p>For the "wald.univregs" this can only be one of the following: waldBeta, waldCR, waldWR, waldER, waldClogit, waldLogistic, waldPois, waldNB, waldBinom, waldZIP, waldMMReg, waldIGreg, waldOrdinal, waldGamma or waldNormLog.</p> <p>For the "perm.univregs" this can only be one of the following: permFisher, permReg, permRQ, permBeta, permCR, permWR, permER, permClogit, permLogistic, permPois, permNB, permBinom, permgSquare, permZIP, permMVreg, permIGreg, permGamma or permNormLog.</p> <p>For the "score.univregs" this can only be one of the following: testIndBeta, testIndLogistic, testIndPois, testIndNB, testIndGamma or censIndWR. Bear in mind that for the score test in testIndLogistic the target must be either binary or nominal multinomial. Ordinal regression is not supported.</p> <p>Note that in all cases you must give the name of the test, without " ".</p>
user_test	A user-defined conditional independence test (provide a closure type object). Default value is NULL. If this is defined, the "test" argument is ignored.
wei	A vector of weights to be used for weighted regression. The default value is NULL.
gam	This is for ebic.univregs only and it refers to the <i>gamma</i> value of the eBIC. If it is NULL, the default value is calculated and if gam is zero, the usual BIC is returned. see Chen and Chen (2008) for more information.
threshold	Threshold (suitable values in [0,1]) for assessing p-values significance. Default value is 0.05. The reason for this is to speed up the computations. The permutation p-value is calculated as the proportion of times the permuted test statistic is higher than the observed test statistic. When running the permutations, if the proportion is more than 0.05, the process stops. A decision must be made fast, and if the non rejection decision has been made, there is no need to perform the rest permutations; the decision cannot change.
R	This is for the permutations based regression models. It is the number of permutations to apply. Note, that not all the number of permutations need be performed. If the number of times the test statistic is greater than the observed test statistic is more than $\text{threshold} * R$, the iterations stop, as a decision has already been made.
ncores	How many cores to use. This plays an important role if you have tens of thousands of variables or really large sample sizes and tens of thousands of variables and a regression based test which requires numerical optimisation. In other cases it will not make a difference in the overall time (in fact it can be slower). The parallel computation is used in the first step of the algorithm, where univariate associations are examined, those take place in parallel. We have seen a reduction

	in time of 50% with 4 cores in comparison to 1 core. Note also, that the amount of reduction is not linear in the number of cores.
id	A numerical vector of the same length as target with integer valued numbers, such as 1, 2, 3,... (zeros, negative values and factors are not allowed) specifying the clusters or subjects. This argument is for the rint.regs (see details for more information).
reps	If you have measurements over time (lognitudinal data) you can put the time here (the length must be equal to the length of the target) or set it equal to NULL. (see details for more information).
tol	The tolerance value to terminate the Newton-Raphson algorithm in the random effects models.
slopes	Should random slopes for the ime effect be fitted as well? Default value is FALSE.
correl	The correlation structure. For the Gaussian, Logistic, Poisson and Gamma regression this can be either "exchangeable" (compound symmetry, suitable for clustered data) or "ar1" (AR(1) model, suitable for longitudinal data). For the ordinal logistic regression its only the "exchangeable" correlation sturcture. This is obsolete for "testIndGEEOrdinal", but is here for compatibility reasons.
se	<p>The method for estimating standard errors. This is very important and crucial. The available options for Gaussian, Logistic, Poisson and Gamma regression are: a) 'san.se', the usual robust estimate. b) 'jack': if approximate jackknife variance estimate should be computed. c) 'j1s': if 1-step jackknife variance estimate should be computed and d) 'fij': logical indicating if fully iterated jackknife variance estimate should be computed. If you have many clusters (sets of repeated measurements) "san.se" is fine as it is astmpotically correct, plus jackknife estimates will take longer. If you have a few clusters, then maybe it's better to use jackknife estimates.</p> <p>The jackknife variance estimator was suggested by Paik (1988), which is quite suitable for cases when the number of subjects is small ($K < 30$), as in many biological studies. The simulation studies conducted by Ziegler et al. (2000) and Yan and Fine (2004) showed that the approximate jackknife estimates are in many cases in good agreement with the fully iterated ones. This is obsolete for "testIndGEEOrdinal", but is here for compatibility reasons.</p>

Details

This function is more as a help function for SES and MMPC, but it can also be called directly by the user. In some, one should specify the regression model to use and the function will perform all simple regressions, i.e. all regression models between the target and each of the variables in the dataset.

For the score.univregs, score based tests are used which are extremely fast.

For the rint.regs, we perform linear mixed models (weights are not allowed) with random intercepts only (no ranodm slopes). This function works for clustered or longitudinal data. The covariance structure we impose is compound symmetry, hence for longitudinal data, this may not be the best option, yet it will work.

Value

In the case of "ebic.univregs" a list with one element

ebic The eBIC of every model.

For all other cases a list including:

stat The value of the test statistic.

pvalue The logarithm of the p-value of the test.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

Chen J. and Chen Z. (2008). Extended Bayesian information criteria for model selection with large model spaces. *Biometrika*, 95(3): 759-771.

See Also

[SES](#), [MMPC](#), [CondIndTests](#), [reg.fit](#), [ridge.reg](#)

Examples

```
y <- rpois(50, 15)
x <- matrix( rnorm(50 * 10), ncol = 10)
a1 <- univregs(y, x, test = testIndPois)
a2 <- perm.univregs(y, x, test = permPois)
a3 <- wald.univregs(y, x, test = waldPois)
```

Utilities for the skeleton of a (Bayesian) Network

Utilities for the skeleton of a (Bayesian) Network

Description

Utilities for the skeleton of a (Bayesian) Network

Usage

```
bn.skel.utils(mod, G = NULL, roc = TRUE, alpha = 0.01)
bn.skel.utils2(mod, G = NULL, roc = TRUE, alpha = 0.01)
```

Arguments

<code>mod</code>	An object as returned by <code>pc.skel</code> , <code>glmm.pc.skel</code> or <code>mmhc.skel</code> .
<code>G</code>	The true adjacency matrix with 1 indicating an edge and zero its absence. Symmetric or not is not important. If this is not available, leave it NULL.
<code>roc</code>	Do you want a graph with the ROC curve be returned? Default value is TRUE.
<code>alpha</code>	The significance level (suitable values in (0, 1)) for assessing the p-values. Default value is 0.01.

Details

Given the true adjacency matrix one can evaluate the estimated adjacency matrix, skeleton, of the Pc or the MMHC algorithm.

The `bn.skels.utils` give you the area under the curve, false discovery rate and sorting of the edges based on their p-values.

The `bn.skel.utils2` estimates the confidence of each edge. The estimated proportion of null p-values is estimated the algorithm by Storey and Tibshirani (2003).

Value

For the "`bn.skel.utils`" a list including:

<code>fdr</code>	The false discovery rate as estimated using the Benjamini-Hochberg correction.
<code>area</code>	This is a list with the elements of the <code>auc</code> function. The area under the curve, the sensitivity and specificity for a range of values, the Youden index, etc.
<code>sig.pvalues</code>	A matrix with the row and column of each significant p-value sorted in ascending order. As we move down the matrix, the p-values increase and hence the strength of the associations decreases.

For the "`bn.skel.utils2`" a list including:

<code>area</code>	This is a list with the elements of the <code>auc</code> function. The area under the curve, the sensitivity and specificity for a range of values, the Youden index, etc.
<code>pxy</code>	A matrix with the row and column of the confidence of each p-value sorted in ascending order. As we move down the matrix, the confidences decrease.
<code>lower</code>	The lower confidence limit of an edge as estimated by <code>conf.edge.lower</code> .

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

Tsamardinos I. and Brown L.E. Bounding the False Discovery Rate in Local Bayesian Network Learning. AAAI, 2008.

Triantafillou S., Tsamardinos I. and Roumpelaki A. (2014). Learning neighborhoods of high confidence in constraint-based causal discovery. In European Workshop on Probabilistic Graphical Models, pp. 487-502.

Storey J.D. and Tibshirani R. (2003). Statistical significance for genome-wide experiments. Proceedings of the National Academy of Sciences, 100: 9440-9445.

Benjamini Y. and Hochberg Y. (1995). Controlling the false discovery rate: a practical and powerful approach to multiple testing. Journal of the Royal Statistical Society Series B, 57(1), 289-300.

Spirtes P., Glymour C. and Scheines R. (2001). Causation, Prediction, and Search. The MIT Press, Cambridge, MA, USA, 3rd edition.

See Also

[pc.skel](#), [mmhc.skel](#), [corfs.network](#), [local.mmhc.skel](#), [conf.edge.lower](#)

Examples

```
## simulate a dataset with continuous data
y <- rdag2(500, p = 25, nei = 3)
ind <- sample(1:25, 25)
x <- y$x[, ind]
mod <- pc.skel( x, method = "comb.fast", alpha = 0.01 )
G <- y$G[ind, ind]
G <- G + t(G)
bn.skel.utils(mod, G, roc = FALSE, alpha = 0.01)
bn.skel.utils2(mod, G, roc = FALSE, alpha = 0.01)
```

Zero inflated Poisson regression

Zero inflated Poisson regression

Description

Zero inflated Poisson regression.

Usage

```
zip.mod(target, dataset, wei = NULL, xnew = NULL)
zip.reg(target, dataset, wei = NULL, lgy = NULL)
```

Arguments

target	The target (dependent) variable. It must be a numerical vector with integers.
dataset	The independent variable(s). It can be a vector, a matrix or a dataframe with continuous only variables, a data frame with mixed or only categorical variables.
wei	A vector of weights to be used for weighted regression. The default value is NULL.
xnew	If you have new values for the predictor variables (dataset) whose target variable you want to predict insert them here. If you put the "dataset" or leave it NULL it will calculate the regression fitted values.
lgy	If you have already calculated the constant term of the ZIP regression plug it here. This is the sum of the logarithm of the factorial of the values.

Details

The zero inflated Poisson regression as suggested by Lambert (1992) is fitted. Unless you have a sufficient number of zeros, there is no reason to use this model. The "zip.reg" is an internal wrapper function and is used for speed up purposes. It is not to be called directly by the user unless they know what they are doing.

Value

A list including:

be	The estimated coefficients of the model.
prop	The estimated proportion of zeros.
loglik	The log-likelihood of the regression model.
est	The estimated values if "xnew" is not NULL.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@csd.uoc.gr>

References

Lambert D. (1992). Zero-inflated Poisson regression, with an application to defects in manufacturing. *Technometrics*, 34(1):1-14.

See Also

[testIndZIP](#), [zip.regs](#), [reg.fit](#), [ridge.reg](#)

Examples

```
y <- rpois(100, 2)
x <- matrix( rnorm(100 * 2), ncol = 2)
a1 <- glm(y ~ x, poisson)
a2 <- zip.mod(y, x)
summary(a1)
logLik(a1)
a2 ## a ZIP is not really necessary
y[1:20] <- 0
a1 <- glm(y ~ x, poisson)
a2 <- zip.mod(y, x)
summary(a1)
logLik(a1)
a2 ## a ZIP is probably more necessary
```

Index

- *Topic **Ancestors**
 - Ancestors and descendants of a node in a directed graph, [6](#)
- *Topic **Area under the curve**
 - Conditional independence tests counting the number of times a possible collider d-separates two nodes, [53](#)
 - Drop all possible single terms from a model using the partial correlation, [103](#)
 - Effective sample size for G^2 test in BNs with case control data, [104](#)
 - ROC and AUC, [175](#)
- *Topic **Backward regression**
 - Backward selection regression, [9](#)
 - IAMB backward selection phase, [133](#)
 - IAMB variable selection, [135](#)
- *Topic **Beta regression**
 - Conditional independence test for proportions/percentages, [48](#)
- *Topic **Binary logistic regression**
 - Conditional independence test for binary, categorical or ordinal data, [35](#)
 - Conditional independence test for the static-longitudinal scenario, [50](#)
- *Topic **Binomial logistic regression**
 - Conditional independence tests for success rates, [67](#)
- *Topic **Case-control studies**
 - Conditional independence test for case control data, [39](#)
- *Topic **Conditional Independence Test**
 - Correlation based conditional independence tests, [91](#)
- *Topic **Conditional independence tests**
 - Conditional independence tests, [28](#)
- *Topic **Conditional independence test**
 - Conditional independence test for binary, categorical or ordinal data, [35](#)
 - Conditional independence test for case control data, [39](#)
 - Conditional independence test for longitudinal and clustered data using GLMM, [45](#)
 - Conditional independence test for proportions/percentages, [48](#)
 - Conditional independence test for the static-longitudinal scenario, [50](#)
 - Conditional independence tests for continuous univariate and multivariate data, [55](#)
 - Conditional independence tests for count data, [58](#)
 - Conditional independence tests for left censored data, [61](#)
 - Conditional independence tests for success rates, [67](#)
 - Conditional independence tests for survival data, [69](#)
 - G-square conditional independence test for discrete data, [124](#)
- *Topic **Conditional logistic regression**
 - Conditional independence test for case control data, [39](#)
- *Topic **Conditioning set**
 - Partial correlation between two variables, [163](#)
- *Topic **Cox regression**
 - Conditional independence tests for survival data, [69](#)
- *Topic **Cross validation**

- Cross-Validation for SES and MMPC, 96
- *Topic **DAGs**
 - Check Markov equivalence of two DAGs, 26
- *Topic **DAG**
 - Topological sort of a DAG, 195
- *Topic **Descendants**
 - Ancestors and descendants of a node in a directed graph, 6
- *Topic **Directed acyclic graph**
 - Check whether a directed graph is acyclic, 27
- *Topic **Directed acyclic graph**
 - Data simulation from a DAG, 100
- *Topic **Essential graph**
 - Transformation of a DAG into an essential graph, 197
- *Topic **Feature Selection**
 - Backward phase of MMPC, 7
 - Constraint based feature selection algorithms, 75
 - Constraint based feature selection algorithms for multiple datasets, 86
 - Feature selection using SES and MMPC for classification with longitudinal data, 106
 - MMPC solution paths for many combinations of hyper-parameters, 147
 - MXM-package, 4
- *Topic **Fisher's Test**
 - Correlation based conditional independence tests, 91
- *Topic **Fisher's test**
 - Conditional independence tests with and without permutation p-value, 73
- *Topic **Gamma regression**
 - Conditional independence tests for positive data, 64
- *Topic **Generation of folds**
 - Generate random folds for cross-validation, 132
- *Topic **Graphical visualisation**
 - Plot of longitudinal data, 165
- *Topic **Internal MXM Functions**
 - MXM-internal, 153
- *Topic **Linear mixed model**
 - Conditional independence test for longitudinal and clustered data using GLMM, 45
- *Topic **Linear regression test**
 - Conditional independence tests for continuous univariate and multivariate data, 55
- *Topic **Log likelihood ratio**
 - Conditional independence tests for left censored data, 61
 - Conditional independence tests for survival data, 69
 - G-square conditional independence test for discrete data, 124
- *Topic **Log link**
 - Conditional independence tests for positive data, 64
- *Topic **Longitudinal data**
 - Constraint based feature selection algorithms for longitudinal and clustered data, 81
 - Plot of longitudinal data, 165
- *Topic **MMHC algorithm**
 - Bayesian Network construction using a hybrid of MMPC and PC, 16
 - Skeleton (local) around a node of the max-min hill-climbing (MMHC) algorithm, 180
 - Skeleton of the max-min hill-climbing (MMHC) algorithm, 182
- *Topic **MMPC output**
 - MMPCoutput-class, 152
- *Topic **MMPC.temporal output**
 - MMPC.temporal.output-class, 151
- *Topic **MMPC**
 - Cross-Validation for SES and MMPC, 96
- *Topic **Markov Blanket**
 - Backward selection regression, 9
 - Backward selection with generalised linear regression models, 15
 - BIC based forward selection, 20
 - BIC based forward selection with

- generalised linear models, [22](#)
 - Forward selection regression, [118](#)
 - Forward selection with generalised linear regression models, [121](#)
 - Forward selection with linear regression models, [123](#)
 - IAMB backward selection phase, [133](#)
 - IAMB variable selection, [135](#)
 - The max-min Markov blanket algorithm, [193](#)
- *Topic **Markov equivalence**
 - Check Markov equivalence of two DAGs, [26](#)
- *Topic **Mixed models**
 - Constraint based feature selection algorithms for longitudinal and clustered data, [81](#)
- *Topic **Multinomial logistic regression**
 - Conditional independence test for binary, categorical or ordinal data, [35](#)
 - Conditional independence test for the static-longitudinal scenario, [50](#)
- *Topic **Multiple Feature Signatures**
 - Backward phase of MMPC, [7](#)
 - Constraint based feature selection algorithms, [75](#)
 - Constraint based feature selection algorithms for multiple datasets, [86](#)
 - Feature selection using SES and MMPC for classification with longitudinal data, [106](#)
 - MMPC solution paths for many combinations of hyper-parameters, [147](#)
 - MXM-package, [4](#)
- *Topic **Negative binomial regression**
 - Conditional independence tests for count data, [58](#)
- *Topic **Neighbour nodes**
 - Markov Blanket of a node in a directed graph, [145](#)
 - Neighbours of nodes in an undirected graph, [158](#)
- *Topic **Network construction**
 - Network construction using the partial correlation based forward regression or FBED, [159](#)
- *Topic **Network plot**
 - Interactive plot of an (un)directed graph, [138](#)
 - Markov Blanket of a node in a directed graph, [145](#)
 - Neighbours of nodes in an undirected graph, [158](#)
- *Topic **Ordinal logistic regression**
 - Conditional independence test for binary, categorical or ordinal data, [35](#)
 - Conditional independence test for the static-longitudinal scenario, [50](#)
- *Topic **PC algorithm**
 - Orientation rules for the PC algorithm, [161](#)
- *Topic **Partial correlation**
 - Partial correlation between two variables, [163](#)
- *Topic **Permutation test**
 - Permutation based p-value for the Pearson correlation coefficient, [164](#)
- *Topic **Poisson regressions**
 - Many Wald based tests for logistic and Poisson regressions with continuous predictors, [144](#)
- *Topic **Poisson regression**
 - Conditional independence tests for count data, [58](#)
- *Topic **Regression modelling**
 - Generalised linear mixed models based on temporal SES and MMPC outputs, [127](#)
 - Regression models based on SES and MMPC outputs, [167](#)
 - Regression models based on SES.timeclass and MMPC.timeclass outputs, [169](#)
- *Topic **Regression models**
 - Conditional independence tests, [28](#)
 - Many simple beta regressions, [141](#)
 - Many simple zero inflated Poisson regressions, [142](#)

- Univariate regression based tests, [201](#)
- *Topic **Regression model**
 - Beta regression, [19](#)
 - Generalised ordinal regression, [129](#)
 - Regression models fitting, [171](#)
 - Zero inflated Poisson regression, [206](#)
- *Topic **Ridge regression**
 - Cross-validation for ridge regression, [94](#)
 - Ridge regression, [173](#)
 - Ridge regression coefficients plot, [174](#)
- *Topic **SES output**
 - SESoutput-class, [179](#)
- *Topic **SES.temporal output**
 - SES.gee.output-class, [177](#)
 - SES.temporal.output-class, [178](#)
- *Topic **SES**
 - Backward phase of MMPC, [7](#)
 - Constraint based feature selection algorithms, [75](#)
 - Constraint based feature selection algorithms for longitudinal and clustered data, [81](#)
 - Constraint based feature selection algorithms for multiple datasets, [86](#)
 - Cross-Validation for SES and MMPC, [96](#)
 - Feature selection using SES and MMPC for classification with longitudinal data, [106](#)
 - MMPC solution paths for many combinations of hyper-parameters, [147](#)
 - MXM-package, [4](#)
- *Topic **Structurall Haming distance**
 - Structural Hamming distance between two partially oriented DAGs, [189](#)
- *Topic **Survival**
 - Conditional independence tests for left censored data, [61](#)
 - Conditional independence tests for survival data, [69](#)
- *Topic **Tobit regression**
 - Conditional independence tests for left censored data, [61](#)
- *Topic **Topological sort**
 - Topological sort of a DAG, [195](#)
- *Topic **Undirected path**
 - Undirected path(s) between two nodes, [200](#)
- *Topic **Variable Selection**
 - Backward phase of MMPC, [7](#)
 - Backward selection regression, [9](#)
 - Backward selection with generalised linear regression models, [15](#)
 - BIC based forward selection, [20](#)
 - BIC based forward selection with generalised linear models, [22](#)
 - Constraint based feature selection algorithms, [75](#)
 - Constraint based feature selection algorithms for longitudinal and clustered data, [81](#)
 - Constraint based feature selection algorithms for multiple datasets, [86](#)
 - Feature selection using SES and MMPC for classification with longitudinal data, [106](#)
 - Forward selection regression, [118](#)
 - Forward selection with generalised linear regression models, [121](#)
 - Forward selection with linear regression models, [123](#)
 - IAMB backward selection phase, [133](#)
 - IAMB variable selection, [135](#)
 - MMPC solution paths for many combinations of hyper-parameters, [147](#)
 - The max-min Markov blanket algorithm, [193](#)
- *Topic **Wald test**
 - Many Wald based tests for logistic and Poisson regressions with continuous predictors, [144](#)
- *Topic **Weibull regression**
 - Conditional independence tests for survival data, [69](#)
- *Topic **Zero inflated poisson regression**

- Conditional independence tests
 - for count data, [58](#)
- *Topic **beta distribution**
 - Beta regression, [19](#)
 - Many simple beta regressions, [141](#)
- *Topic **check for cycles**
 - Check whether a directed graph is acyclic, [27](#)
- *Topic **conditional independence test**
 - Conditional independence tests with and without permutation p-value, [73](#)
 - Permutation based p-value for the Pearson correlation coefficient, [164](#)
- *Topic **directed graph**
 - Interactive plot of an (un)directed graph, [138](#)
- *Topic **distance between DAGs**
 - Structural Hamming distance between two partially oriented DAGs, [189](#)
- *Topic **equivalence class**
 - Transformation of a DAG into an essential graph, [197](#)
- *Topic **exponential regression**
 - Conditional independence tests for survival data, [69](#)
- *Topic **forward regression**
 - Network construction using the partial correlation based forward regression or FBED, [159](#)
- *Topic **interactive graph**
 - Interactive plot of an (un)directed graph, [138](#)
- *Topic **list(list(G^2))**
 - G-square conditional independence test for discrete data, [124](#)
- *Topic **logistic regressions**
 - Many Wald based tests for logistic and Poisson regressions with continuous predictors, [144](#)
- *Topic **meta analytic ses output**
 - mases.output-class, [146](#)
- *Topic **meta-analytic MMPC output**
 - mammpc.output-class, [140](#)
- *Topic **ordinal regression**
 - Generalised ordinal regression, [129](#)
- *Topic **parallel**
 - Cross-Validation for SES and MMPC, [96](#)
- *Topic **pc algorithm**
 - Skeleton of the PC algorithm, [185](#)
- *Topic **permutation based p-value**
 - Conditional independence tests with and without permutation p-value, [73](#)
- *Topic **receiver operating curve**
 - Conditional independence tests counting the number of times a possible collider d-separates two nodes, [53](#)
 - Drop all possible single terms from a model using the partial correlation, [103](#)
 - Effective sample size for G^2 test in BNs with case control data, [104](#)
 - ROC and AUC, [175](#)
- *Topic **simulation of random values**
 - Data simulation from a DAG, [100](#)
- *Topic **test**
 - G-square conditional independence test for discrete data, [124](#)
- *Topic **zero inflated Poisson**
 - Many simple zero inflated Poisson regressions, [142](#)
 - Zero inflated Poisson regression, [206](#)
- *Topic
 - G-square conditional independence test for discrete data, [124](#)
- acc.mxm (Cross-Validation for SES and MMPC), [96](#)
- acc_multinom.mxm (Cross-Validation for SES and MMPC), [96](#)
- Ancestors and descendants of a node in a directed graph, [6](#)
- anova, [63](#), [72](#)
- apply_ideq (MXM-internal), [153](#)
- auc, [205](#)
- auc (ROC and AUC), [175](#)
- auc.mxm (Cross-Validation for SES and MMPC), [96](#)
- Backward phase of MMPC, [7](#)

- Backward selection regression, [9](#)
- Backward selection regression for GLMM, [10](#)
- Backward selection regression for GLMM using the eBIC, [12](#)
- Backward selection regression using the eBIC, [13](#)
- Backward selection with generalised linear regression models, [15](#)
- Bayesian Network construction using a hybrid of MMPC and PC, [16](#)
- Beta regression, [19](#)
- beta.bsreg (MXM-internal), [153](#)
- beta.fsreg (MXM-internal), [153](#)
- beta.fsreg_2 (MXM-internal), [153](#)
- beta.mod, [142](#)
- beta.mod (Beta regression), [19](#)
- beta.mxm (Cross-Validation for SES and MMPC), [96](#)
- beta.regs, [20](#)
- beta.regs (Many simple beta regressions), [141](#)
- betamle.wei (MXM-internal), [153](#)
- BIC based forward selection, [20](#)
- BIC based forward selection with generalised linear models, [22](#)
- bic.betafsreg (MXM-internal), [153](#)
- bic.clogit.fsreg (MXM-internal), [153](#)
- bic.fsreg, [10](#), [14](#), [16](#), [23](#), [112](#), [120](#), [122](#), [124](#), [134](#), [136](#)
- bic.fsreg (BIC based forward selection), [20](#)
- bic.gammasfsreg (BIC based forward selection with generalised linear models), [22](#)
- bic.glm.fsreg, [10](#), [16](#), [22](#), [120](#), [122](#), [124](#), [134](#), [136](#)
- bic.glm.fsreg (BIC based forward selection with generalised linear models), [22](#)
- bic.mm.fsreg (BIC based forward selection with generalised linear models), [22](#)
- bic.normlog.fsreg (BIC based forward selection with generalised linear models), [22](#)
- bic.tobit.fsreg (MXM-internal), [153](#)
- bic.wr.fsreg (MXM-internal), [153](#)
- bic.zipfsreg (MXM-internal), [153](#)
- bn.skel.utils, [106](#), [185](#), [188](#)
- bn.skel.utils (Utilities for the skeleton of a (Bayesian) Network), [204](#)
- bn.skel.utils2, [139](#)
- bn.skel.utils2 (Utilities for the skeleton of a (Bayesian) Network), [204](#)
- bs.reg (Backward selection regression), [9](#)
- Calculation of the constant and slope for each subject over time, [24](#)
- cat.ci, [192](#), [193](#)
- cat.ci (Conditional independence tests with and without permutation p-value), [73](#)
- cat_condis (MXM-internal), [153](#)
- censIndCR, [6](#), [41](#), [100](#), [126](#)
- censIndCR (Conditional independence tests for survival data), [69](#)
- censIndER (Conditional independence tests for survival data), [69](#)
- censIndWR, [41](#), [63](#), [72](#)
- censIndWR (Conditional independence tests for survival data), [69](#)
- Certificate of exclusion from the selected variables set using SES or MMPC, [25](#)
- certificate.of.exclusion (Certificate of exclusion from the selected variables set using SES or MMPC), [25](#)
- Check Markov equivalence of two DAGs, [26](#)
- Check whether a directed graph is acyclic, [27](#)
- ci.fast (Symmetric conditional independence test with mixed data), [191](#)
- ci.fast2 (Symmetric conditional independence test with mixed data), [191](#)
- ci.mm, [102](#)
- ci.mm (Symmetric conditional independence test with mixed data), [191](#)

- ci.mm2 (Symmetric conditional independence test with mixed data), [191](#)
- ci.mxm (Cross-Validation for SES and MMPC), [96](#)
- ciwr.mxm (Cross-Validation for SES and MMPC), [96](#)
- clogit.fsreg (MXM-internal), [153](#)
- clogit.fsreg_2 (MXM-internal), [153](#)
- comb_condis (MXM-internal), [153](#)
- compare_p_values (MXM-internal), [153](#)
- condi, [191–193](#)
- condi (Conditional independence tests with and without permutation p-value), [73](#)
- condi.perm (MXM-internal), [153](#)
- CondInditonal independence tests, [28](#)
- CondIndTests, [7–10, 15, 16, 20, 22, 23, 38, 45, 47, 50, 52, 57, 61, 63, 66, 69, 72, 74, 76, 78, 80, 86–88, 90, 94, 100, 120, 122, 124, 126, 134, 136, 148, 149, 165, 172, 194, 204](#)
- CondIndTests (CondInditonal independence tests), [28](#)
- condis (Conditional independence tests counting the number of times a possible collider d-separates two nodes), [53](#)
- Conditional independence test for binary, categorical or ordinal data, [35](#)
- Conditional independence test for case control data, [39](#)
- Conditional independence test for longitudinal and clustered data using GEE, [42](#)
- Conditional independence test for longitudinal and clustered data using GLMM, [45](#)
- Conditional independence test for proportions/percentages, [48](#)
- Conditional independence test for the static-longitudinal scenario, [50](#)
- Conditional independence tests counting the number of times a possible collider d-separates two nodes, [53](#)
- Conditional independence tests for continous univariate and multivariate data, [55](#)
- Conditional independence tests for count data, [58](#)
- Conditional independence tests for left censored data, [61](#)
- Conditional independence tests for positive data, [64](#)
- Conditional independence tests for sucess rates, [67](#)
- Conditional independence tests for survival data, [69](#)
- Conditional independence tests with and without permutation p-value, [73](#)
- conf.edge.lower, [106, 110, 205, 206](#)
- conf.edge.lower (Lower limit of the confidence of an edge), [139](#)
- Constraint based feature selection algorithms, [75](#)
- Constraint based feature selection algorithms for longitudinal and clustered data, [81](#)
- Constraint based feature selection algorithms for multiple datasets, [86](#)
- cor.drop1 (Drop all possible single terms from a model using the partial correlation), [103](#)
- cor.fbed, [132](#)
- cor.fsreg, [132](#)
- corfbed.network (Network construction using the partial correlation based forward regression or FBED), [159](#)
- corfs.network, [18, 106, 110, 182, 185, 188, 206](#)
- corfs.network (Network construction using the partial correlation based forward regression or FBED), [159](#)
- Correlation based conditonal independence tests, [91](#)
- correls, [132](#)
- coxph.mxm (Cross-Validation for SES and MMPC), [96](#)
- Cross-validation for ridge regression,

- 94
- Cross-Validation for SES and MMPC, 96
- cv.mmpc, 8, 129, 138, 169
- cv.mmpc (Cross-Validation for SES and MMPC), 96
- cv.permmmpc (Cross-Validation for SES and MMPC), 96
- cv.permses (Cross-Validation for SES and MMPC), 96
- cv.ses, 80, 90, 129, 133, 138, 149, 169
- cv.ses (Cross-Validation for SES and MMPC), 96
- cv.waldmmpc (Cross-Validation for SES and MMPC), 96
- cv.waldses (Cross-Validation for SES and MMPC), 96
- cvmmpc.par (MXM-internal), 153
- cvpermmpc.par (MXM-internal), 153
- cvpermses.par (MXM-internal), 153
- cvses.par (MXM-internal), 153
- cvwaldmmpc.par (MXM-internal), 153
- cvwaldses.par (MXM-internal), 153
- dag2eg, 28
- dag2eg (Transformation of a DAG into an essential graph), 197
- dag_to_eg (MXM-internal), 153
- Data simulation from a DAG, 100
- disctor_condis (MXM-internal), 153
- dist.condi, 192
- dist.condi (Conditional independence tests with and without permutation p-value), 73
- Drop all possible single terms from a model using the partial correlation, 103
- drop1, 103
- ebic.beta.bsreg (MXM-internal), 153
- ebic.bsreg, 112
- ebic.bsreg (Backward selection regression using the eBIC), 13
- ebic.cr.bsreg (MXM-internal), 153
- ebic.fbed.beta (MXM-internal), 153
- ebic.fbed.cr (MXM-internal), 153
- ebic.fbed.glm (MXM-internal), 153
- ebic.fbed.glmm (MXM-internal), 153
- ebic.fbed.lm (MXM-internal), 153
- ebic.fbed.lmm (MXM-internal), 153
- ebic.fbed.mmreg (MXM-internal), 153
- ebic.fbed.multinom (MXM-internal), 153
- ebic.fbed.nb (MXM-internal), 153
- ebic.fbed.ordinal (MXM-internal), 153
- ebic.fbed.tobit (MXM-internal), 153
- ebic.fbed.wr (MXM-internal), 153
- ebic.fbed.zip (MXM-internal), 153
- ebic.glm.bsreg (MXM-internal), 153
- ebic.glmm.bsreg, 11
- ebic.glmm.bsreg (Backward selection regression for GLMM using the eBIC), 12
- ebic.glmm.reps.bsreg (MXM-internal), 153
- ebic.lm.bsreg (MXM-internal), 153
- ebic.mm.bsreg (MXM-internal), 153
- ebic.multinom.bsreg (MXM-internal), 153
- ebic.ordinal.bsreg (MXM-internal), 153
- ebic.tobit.bsreg (MXM-internal), 153
- ebic.univregs (Univariate regression based tests), 201
- ebic.wr.bsreg (MXM-internal), 153
- ebic.zip.bsreg (MXM-internal), 153
- ebicScore (MXM-internal), 153
- Effective sample size for G^2 test in BNs with case control data, 104
- equivdags (Check Markov equivalence of two DAGs), 26
- Estimation of the percentage of Null p-values, 105
- euclid_sens.spec.mxm (Cross-Validation for SES and MMPC), 96
- fbed.beta (MXM-internal), 153
- fbed.cr (MXM-internal), 153
- fbed.gee.reg, 25, 118
- fbed.gee.reg (Forward Backward Early Dropping selection regression with GEE), 113
- fbed.geeglm (MXM-internal), 153
- fbed.geelm (MXM-internal), 153
- fbed.glm (MXM-internal), 153
- fbed.glm2 (MXM-internal), 153
- fbed.glmm (MXM-internal), 153
- fbed.glmm.reg, 11, 13, 115
- fbed.glmm.reg (Forward Backward Early Dropping selection regression with GLMM), 116
- fbed.lm (MXM-internal), 153
- fbed.lmm (MXM-internal), 153

- fbed.mmreg (MXM-internal), 153
- fbed.multinom (MXM-internal), 153
- fbed.nb (MXM-internal), 153
- fbed.ordgee (MXM-internal), 153
- fbed.ordinal (MXM-internal), 153
- fbed.reg, 103, 115, 118, 137
- fbed.reg (Forward Backward Early Dropping selection regression), 110
- fbed.tobit (MXM-internal), 153
- fbed.wr (MXM-internal), 153
- fbed.zip (MXM-internal), 153
- fbedreg.bic, 172
- fbedreg.bic (Incremental BIC values and final regression model of the FBED algorithm), 137
- Feature selection using SES and MMPC for classification with longitudinal data, 106
- findAncestors (Ancestors and descendants of a node in a directed graph), 6
- findDescendants (Ancestors and descendants of a node in a directed graph), 6
- Fit a mixture of beta distributions in p-values, 109
- Forward Backward Early Dropping selection regression, 110
- Forward Backward Early Dropping selection regression with GEE, 113
- Forward Backward Early Dropping selection regression with GLMM, 116
- Forward selection regression, 118
- Forward selection with generalised linear regression models, 121
- Forward selection with linear regression models, 123
- fs.reg, 14, 16, 23, 112, 122, 124, 132
- fs.reg (Forward selection regression), 118
- fs.reg_2 (MXM-internal), 153
- fscore.mxm (Cross-Validation for SES and MMPC), 96
- G-square conditional independence test for discrete data, 124
- gammabsreg (Backward selection with generalised linear regression models), 15
- gammafsreg (Forward selection with generalised linear regression models), 121
- gammafsreg_2 (MXM-internal), 153
- Generalised linear mixed models based on temporal SES and MMPC outputs, 127
- Generalised ordinal regression, 129
- Generalised orthogonal matching pursuit (gOMP), 131
- Generate random folds for cross-validation, 132
- generatefolds (Generate random folds for cross-validation), 132
- glm, 144
- glm.bsreg, 103
- glm.bsreg (Backward selection with generalised linear regression models), 15
- glm.fsreg, 10, 22, 120, 134, 136
- glm.fsreg (Forward selection with generalised linear regression models), 121
- glm.fsreg_2 (MXM-internal), 153
- glm.mxm (Cross-Validation for SES and MMPC), 96
- glmm.bsreg, 13, 25, 115, 118
- glmm.bsreg (Backward selection regression for GLMM), 10
- glmm.ci.mm (Symmetric conditional independence test with clustered data), 190
- glmm.gee.univregs (Univariate regression based tests), 201
- glmm.pc.skel (Skeleton of the PC algorithm), 185
- gomp (Generalised orthogonal matching pursuit (gOMP)), 131
- gomp2 (MXM-internal), 153
- group.mvbetas, 51
- group.mvbetas (Calculation of the constant and slope for each subject over time), 24
- gSquare, 6, 38, 52, 61, 63, 66, 72, 74, 94, 100
- gSquare (G-square conditional

- independence test for discrete data), 124
- iamb (IAMB variable selection), 135
- IAMB backward selection phase, 133
- IAMB variable selection, 135
- iamb.betabs (MXM-internal), 153
- iamb.bs (IAMB backward selection phase), 133
- iamb.gammabs (MXM-internal), 153
- iamb.glmb (MXM-internal), 153
- iamb.normlogbs (MXM-internal), 153
- iamb.tobitbs (MXM-internal), 153
- iamb.zipbs (MXM-internal), 153
- ida (Total causal effect of a node on another node), 196
- IdentifyEquivalence (MXM-internal), 153
- identifyTheEquivalent (MXM-internal), 153
- Incremental BIC values and final regression model of the FBED algorithm, 137
- Interactive plot of an (un)directed graph, 138
- internaliamb.betabs (MXM-internal), 153
- internaliamb.binombs (MXM-internal), 153
- internaliamb.gammabs (MXM-internal), 153
- internaliamb.lmb (MXM-internal), 153
- internaliamb.mmb (MXM-internal), 153
- internaliamb.normlogbs (MXM-internal), 153
- internaliamb.poisbs (MXM-internal), 153
- internaliamb.tobitbs (MXM-internal), 153
- internaliamb.zipbs (MXM-internal), 153
- Internalmampc (MXM-internal), 153
- Internalmases (MXM-internal), 153
- InternalMMPC (MXM-internal), 153
- InternalSES (MXM-internal), 153
- is.dag, 162, 198
- is.dag (Check whether a directed graph is acyclic), 27
- is.sepset (MXM-internal), 153
- kfbed.gEE.reg (MXM-internal), 153
- kfbed.glm.reg (MXM-internal), 153
- kfbed.reg (MXM-internal), 153
- lm.fsreg, 10, 16, 22, 23, 120, 122, 124, 134, 136
- lm.fsreg (Forward selection with linear regression models), 123
- lm.fsreg_2 (MXM-internal), 153
- lm.mxm (Cross-Validation for SES and MMPC), 96
- lmm.bsreg (MXM-internal), 153
- lmrob.mxm (Cross-Validation for SES and MMPC), 96
- local.mmhc.skel, 106, 110, 188, 206
- local.mmhc.skel (Skeleton (local) around a node of the max-min hill-climbing (MMHC) algorithm), 180
- Lower limit of the confidence of an edge, 139
- ma.mmhc, 141, 147
- ma.mmhc (Constraint based feature selection algorithms for multiple datasets), 86
- ma.ses, 141, 147
- ma.ses (Constraint based feature selection algorithms for multiple datasets), 86
- mampc.output (mampc.output-class), 140
- mampc.output-class, 140
- mampc.output-method (mampc.output-class), 140
- Many simple beta regressions, 141
- Many simple zero inflated Poisson regressions, 142
- Many Wald based tests for logistic and Poisson regressions with continuous predictors, 144
- Markov Blanket of a node in a directed graph, 145
- mases.output (mases.output-class), 146
- mases.output-class, 146
- mases.output-method (mases.output-class), 146
- max_min_assoc (MXM-internal), 153
- mb, 7, 139, 162
- mb (Markov Blanket of a node in a directed graph), 145
- min_assoc (MXM-internal), 153
- mm.bsreg (MXM-internal), 153
- mmhc.skel, 8, 27, 102, 106, 110, 138, 139, 158–162, 177, 181, 182, 188, 190, 193, 206

- mmhc.skel (Skeleton of the max-min hill-climbing (MMHC) algorithm), 182
- mmmb (The max-min Markov blanket algorithm), 193
- MMPC, 6, 8, 10, 14, 16, 22–24, 26, 54, 105, 112, 120, 122, 124, 129, 134, 136, 138, 152, 159, 169, 176, 194, 200, 204
- MMPC (Constraint based feature selection algorithms), 75
- MMPC solution paths for many combinations of hyper-parameters, 147
- MMPC.gee, 150, 178
- MMPC.gee (Constraint based feature selection algorithms for longitudinal and clustered data), 81
- mmpc.gee.model (Generalised linear mixed models based on temporal SES and MMPC outputs), 127
- MMPC.gee.output (MMPC.gee.output-class), 149
- MMPC.gee.output-class, 149
- MMPC.gee.output-method (MMPC.gee.output-class), 149
- mmpc.model, 138, 172
- mmpc.model (Regression models based on SES and MMPC outputs), 167
- mmpc.or (Bayesian Network construction using a hybrid of MMPC and PC), 16
- mmpc.path (MMPC solution paths for many combinations of hyper-parameters), 147
- MMPC.temporal, 11, 13, 25, 45, 47, 115, 118, 151, 179
- MMPC.temporal (Constraint based feature selection algorithms for longitudinal and clustered data), 81
- mmpc.temporal.model (Generalised linear mixed models based on temporal SES and MMPC outputs), 127
- MMPC.temporal.output (MMPC.temporal.output-class), 151
- MMPC.temporal.output-class, 151
- MMPC.temporal.output-method (MMPC.temporal.output-class), 151
- MMPC.timeclass, 170
- MMPC.timeclass (Feature selection using SES and MMPC for classification with longitudinal data), 106
- mmpc.timeclass.model, 109
- mmpc.timeclass.model (Regression models based on SES.timeclass and MMPC.timeclass outputs), 169
- mmpcbackphase, 103
- mmpcbackphase (Backward phase of MMPC), 7
- MMPCoutput (MMPCoutput-class), 152
- MMPCoutput-class, 152
- MMPCoutput-method (MMPCoutput-class), 152
- mse.mxm (Cross-Validation for SES and MMPC), 96
- multinom.mxm (Cross-Validation for SES and MMPC), 96
- MXM-internal, 153
- MXM-package, 4
- nb.mxm (Cross-Validation for SES and MMPC), 96
- nbdev.mxm (Cross-Validation for SES and MMPC), 96
- nchoosek (MXM-internal), 153
- nei, 7, 139, 146, 195, 197
- nei (Neighbours of nodes in an undirected graph), 158
- Neighbours of nodes in an undirected graph, 158
- Ness (Effective sample size for G^2 test in BNs with case control data), 104
- Network construction using the partial correlation based forward regression or FBED, 159
- normlog.bsreg (Backward selection with generalised linear regression models), 15
- normlog.fsreg (Forward selection with generalised linear regression

- models), [121](#)
- ord_mae.mxm (Cross-Validation for SES and MMPC), [96](#)
- ordinal.mxm (Cross-Validation for SES and MMPC), [96](#)
- ordinal.reg (Generalised ordinal regression), [129](#)
- Orientation rules for the PC algorithm, [161](#)
- par, [166](#)
- Partial correlation between two variables, [163](#)
- partialcor (Partial correlation between two variables), [163](#)
- pc.con, [27](#), [138](#), [161](#), [162](#), [164](#)
- pc.con (Skeleton of the PC algorithm), [185](#)
- pc.or, [6](#), [7](#), [18](#), [26–28](#), [102](#), [145](#), [146](#), [182](#), [185](#), [190](#), [195](#), [197](#), [198](#)
- pc.or (Orientation rules for the PC algorithm), [161](#)
- pc.skel, [18](#), [74](#), [102](#), [106](#), [110](#), [130](#), [138](#), [139](#), [158](#), [161](#), [162](#), [165](#), [177](#), [182](#), [185](#), [190](#), [191](#), [193](#), [200](#), [206](#)
- pc.skel (Skeleton of the PC algorithm), [185](#)
- pc.skel.boot, [139](#), [140](#)
- pearson_condis (MXM-internal), [153](#)
- perm.apply_ideq (MXM-internal), [153](#)
- perm.betaregs (Many simple beta regressions), [141](#)
- perm.IdentifyEquivalence (MXM-internal), [153](#)
- perm.identifyTheEquivalent (MXM-internal), [153](#)
- perm.Internalmmpc (MXM-internal), [153](#)
- perm.mmpc (Constraint based feature selection algorithms), [75](#)
- perm.ses (Constraint based feature selection algorithms), [75](#)
- perm.univariateScore (MXM-internal), [153](#)
- perm.univregs, [145](#)
- perm.univregs (Univariate regression based tests), [201](#)
- perm.zipregs (Many simple zero inflated Poisson regressions), [142](#)
- permBeta (Conditional independence test for proportions/percentages), [48](#)
- permBinom (Conditional independence tests for success rates), [67](#)
- permClogit (Conditional independence test for case control data), [39](#)
- permcors, [164](#)
- permcors (Permutation based p-value for the Pearson correlation coefficient), [164](#)
- permcors (Permutation based p-value for the Pearson correlation coefficient), [164](#)
- permCR (Conditional independence tests for survival data), [69](#)
- permDcor (Correlation based conditional independence tests), [91](#)
- permER (Conditional independence tests for survival data), [69](#)
- permFisher (Correlation based conditional independence tests), [91](#)
- permGamma (Conditional independence tests for positive data), [64](#)
- permGsquare (G-square conditional independence test for discrete data), [124](#)
- permIGreg (Conditional independence tests for positive data), [64](#)
- permLogistic (Conditional independence test for binary, categorical or ordinal data), [35](#)
- permMMFisher (Correlation based conditional independence tests), [91](#)
- permMMReg (Conditional independence tests for continuous univariate and multivariate data), [55](#)
- permMultinom (Conditional independence test for binary, categorical or ordinal data), [35](#)
- permMVreg (Conditional independence tests for continuous univariate and multivariate data), [55](#)
- permNB (Conditional independence tests for count data), [58](#)
- permNormLog (Conditional independence

- tests for positive data), 64
- permOrdinal (Conditional independence test for binary, categorical or ordinal data), 35
- permPois (Conditional independence tests for count data), 58
- permReg (Conditional independence tests for continuous univariate and multivariate data), 55
- permRQ (Conditional independence tests for continuous univariate and multivariate data), 55
- permTobit (Conditional independence tests for left censored data), 61
- Permutation based p-value for the Pearson correlation coefficient, 164
- permWR (Conditional independence tests for survival data), 69
- permZIP (Conditional independence tests for count data), 58
- pi0est (Estimation of the percentage of Null p-values), 105
- plot, 166
- Plot of longitudinal data, 165
- plot,mammpc.output,ANY-method (mammpc.output-class), 140
- plot,mammpc.output-method (mammpc.output-class), 140
- plot,mases.output,ANY-method (mases.output-class), 146
- plot,mases.output-method (mases.output-class), 146
- plot,MMPC.gee.output,ANY-method (MMPC.gee.output-class), 149
- plot,MMPC.gee.output-method (MMPC.gee.output-class), 149
- plot,MMPC.temporal.output,ANY-method (MMPC.temporal.output-class), 151
- plot,MMPC.temporal.output-method (MMPC.temporal.output-class), 151
- plot,MMPCoutput,ANY-method (MMPCoutput-class), 152
- plot,MMPCoutput-method (MMPCoutput-class), 152
- plot,SES.gee.output,ANY-method (SES.gee.output-class), 177
- plot,SES.gee.output-method (SES.gee.output-class), 177
- plot,SES.temporal.output,ANY-method (SES.temporal.output-class), 178
- plot,SES.temporal.output-method (SES.temporal.output-class), 178
- plot,SESoutput,ANY-method (SESoutput-class), 179
- plot,SESoutput-method (SESoutput-class), 179
- plotnetwork, 7, 146, 160, 161, 177, 183, 186, 190, 195, 197, 198
- plotnetwork (Interactive plot of an (un)directed graph), 138
- pois.mxm (Cross-Validation for SES and MMPC), 96
- poisdev.mxm (Cross-Validation for SES and MMPC), 96
- proc_time-class (MXM-internal), 153
- pval.mixbeta (Fit a mixture of beta distributions in p-values), 109
- quasibinom.bsreg (MXM-internal), 153
- quasibinom.fsreg (MXM-internal), 153
- quasibinom.fsreg_2 (MXM-internal), 153
- quasipois.bsreg (MXM-internal), 153
- quasipois.fsreg (MXM-internal), 153
- quasipois.fsreg_2 (MXM-internal), 153
- R0 (MXM-internal), 153
- R1 (MXM-internal), 153
- R2 (MXM-internal), 153
- R3 (MXM-internal), 153
- rdag (Data simulation from a DAG), 100
- rdag2 (Data simulation from a DAG), 100
- reg.fit, 20, 138, 142, 144, 204, 207
- reg.fit (Regression models fitting), 171
- regbeta (MXM-internal), 153
- regbetawei (MXM-internal), 153
- Regression models based on SES and MMPC outputs, 167
- Regression models based on SES.timeclass and MMPC.timeclass outputs, 169
- Regression models fitting, 171

- regzip (MXM-internal), [153](#)
- regzipwei (MXM-internal), [153](#)
- Ridge regression, [173](#)
- Ridge regression coefficients plot, [174](#)
- ridge.plot (Ridge regression coefficients plot), [174](#)
- ridge.reg, [20](#), [96](#), [130](#), [142](#), [144](#), [172](#), [175](#), [204](#), [207](#)
- ridge.reg (Ridge regression), [173](#)
- ridgereg.cv, [174](#), [175](#)
- ridgereg.cv (Cross-validation for ridge regression), [94](#)
- rint.regs (Univariate regression based tests), [201](#)
- rmdag (Data simulation from a DAG), [100](#)
- ROC and AUC, [175](#)
- rq.mxm (Cross-Validation for SES and MMPC), [96](#)
- score.glm, [144](#)
- score.univregs (Univariate regression based tests), [201](#)
- Search for triangles in an undirected graph, [176](#)
- SES, [6](#), [9](#), [10](#), [16](#), [20](#), [22–24](#), [38](#), [41](#), [50](#), [52](#), [54](#), [63](#), [72](#), [90](#), [94](#), [100](#), [105](#), [120](#), [122](#), [124](#), [126](#), [129](#), [134](#), [136](#), [138](#), [152](#), [159](#), [165](#), [169](#), [176](#), [180](#), [194](#), [200](#), [204](#)
- SES (Constraint based feature selection algorithms), [75](#)
- SES.gee, [150](#), [178](#)
- SES.gee (Constraint based feature selection algorithms for longitudinal and clustered data), [81](#)
- ses.gee.model (Generalised linear mixed models based on temporal SES and MMPC outputs), [127](#)
- SES.gee.output (SES.gee.output-class), [177](#)
- SES.gee.output-class, [177](#)
- SES.gee.output-method (SES.gee.output-class), [177](#)
- ses.model (Regression models based on SES and MMPC outputs), [167](#)
- SES.temporal, [45](#), [47](#), [151](#), [166](#), [179](#)
- SES.temporal (Constraint based feature selection algorithms for longitudinal and clustered data), [81](#)
- ses.temporal.model (Generalised linear mixed models based on temporal SES and MMPC outputs), [127](#)
- SES.temporal.output (SES.temporal.output-class), [178](#)
- SES.temporal.output-class, [178](#)
- SES.temporal.output-method (SES.temporal.output-class), [178](#)
- SES.timeclass (Feature selection using SES and MMPC for classification with longitudinal data), [106](#)
- ses.timeclass.model (Regression models based on SES.timeclass and MMPC.timeclass outputs), [169](#)
- SESoutput (SESoutput-class), [179](#)
- SESoutput-class, [179](#)
- SESoutput-method (SESoutput-class), [179](#)
- shd (Structural Hamming distance between two partially oriented DAGs), [189](#)
- Skeleton (local) around a node of the max-min hill-climbing (MMHC) algorithm, [180](#)
- Skeleton of the max-min hill-climbing (MMHC) algorithm, [182](#)
- Skeleton of the PC algorithm, [185](#)
- Structural Hamming distance between two partially oriented DAGs, [189](#)
- Surv, [62](#), [63](#), [70](#), [72](#)
- survreg, [62](#)
- Symmetric conditional independence test with clustered data, [190](#)
- Symmetric conditional independence test with mixed data, [191](#)
- tc.plot (Plot of longitudinal data), [165](#)
- testIndBeta, [20](#), [69](#), [78](#), [142](#)
- testIndBeta (Conditional independence test for proportions/percentages), [48](#)
- testIndBinom (Conditional independence tests for success rates), [67](#)

- testIndClogit (Conditional independence test for case control data), [39](#)
- testIndFisher, [6](#), [50](#), [57](#), [63](#), [72](#), [74](#), [77](#), [88](#), [100](#), [126](#), [164](#), [165](#)
- testIndFisher (Correlation based conditional independence tests), [91](#)
- testIndGamma (Conditional independence tests for positive data), [64](#)
- testIndGEEGamma (Conditional independence test for longitudinal and clustered data using GEE), [42](#)
- testIndGEELogistic (Conditional independence test for longitudinal and clustered data using GEE), [42](#)
- testIndGEENormLog (Conditional independence test for longitudinal and clustered data using GEE), [42](#)
- testIndGEEOrdinal (Conditional independence test for longitudinal and clustered data using GEE), [42](#)
- testIndGEEPois (Conditional independence test for longitudinal and clustered data using GEE), [42](#)
- testIndGEEReg (Conditional independence test for longitudinal and clustered data using GEE), [42](#)
- testIndGLMMLogistic, [82](#), [128](#)
- testIndGLMMLogistic (Conditional independence test for longitudinal and clustered data using GLMM), [45](#)
- testIndGLMMPois, [82](#), [128](#)
- testIndGLMMPois (Conditional independence test for longitudinal and clustered data using GLMM), [45](#)
- testIndGLMMReg, [82](#), [84](#), [86](#), [128](#), [166](#), [191](#)
- testIndGLMMReg (Conditional independence test for longitudinal and clustered data using GLMM), [45](#)
- testIndIGreg (Conditional independence tests for positive data), [64](#)
- testIndLMM, [128](#)
- testIndLMM (Conditional independence test for longitudinal and clustered data using GLMM), [45](#)
- testIndLogistic, [6](#), [41](#), [54](#), [63](#), [69](#), [72](#), [94](#), [100](#), [105](#), [126](#), [176](#)
- testIndLogistic (Conditional independence test for binary, categorical or ordinal data), [35](#)
- testIndMMFisher (Correlation based conditional independence tests), [91](#)
- testIndMMReg (Conditional independence tests for continuous univariate and multivariate data), [55](#)
- testIndMultinom (Conditional independence test for binary, categorical or ordinal data), [35](#)
- testIndMVreg (Conditional independence tests for continuous univariate and multivariate data), [55](#)
- testIndNB, [61](#), [66](#)
- testIndNB (Conditional independence tests for count data), [58](#)
- testIndNormLog (Conditional independence tests for positive data), [64](#)
- testIndOrdinal (Conditional independence test for binary, categorical or ordinal data), [35](#)
- testIndPois (Conditional independence tests for count data), [58](#)
- testIndQBinom (Conditional independence test for binary, categorical or ordinal data), [35](#)
- testIndQPois (Conditional independence tests for count data), [58](#)
- testIndReg, [50](#), [61](#), [66](#), [69](#), [93](#), [94](#)
- testIndReg (Conditional independence tests for continuous univariate and multivariate data), [55](#)

- testIndRQ, [6](#), [50](#), [57](#)
- testIndRQ (Conditional independence tests for continous univariate and multivariate data), [55](#)
- testIndSpearman, [57](#), [74](#), [94](#), [164](#), [165](#)
- testIndSpearman (Correlation based conditional independence tests), [91](#)
- testIndTimeLogistic (Conditional independence test for the static-longitudinal scenario), [50](#)
- testIndTimeMultinom (Conditional independence test for the static-longitudinal scenario), [50](#)
- testIndTobit (Conditional independence tests for left censored data), [61](#)
- testIndZIP, [61](#), [66](#), [144](#), [207](#)
- testIndZIP (Conditional independence tests for count data), [58](#)
- The max-min Markov blanket algorithm, [193](#)
- Topological sort of a DAG, [195](#)
- topological_sort, [28](#), [198](#)
- topological_sort (Topological sort of a DAG), [195](#)
- Total causal effect of a node on another node, [196](#)
- Transformation of a DAG into an essential graph, [197](#)
- Transitive closure of an adjacency matrix, [199](#)
- transitiveClosure (Transitive closure of an adjacency matrix), [199](#)
- triangles.search (Search for triangles in an undirected graph), [176](#)

- undir.path (Undirected path(s) between two nodes), [200](#)
- Undirected path(s) between two nodes, [200](#)
- Univariate regression based tests, [201](#)
- univariateScore (MXM-internal), [153](#)
- univregs, [145](#)
- univregs (Univariate regression based tests), [201](#)

- Utilities for the skeleton of a (Bayesian) Network, [204](#)

- wald.betaregs (Many simple beta regressions), [141](#)
- wald.Internalmmpc (MXM-internal), [153](#)
- wald.Internalses (MXM-internal), [153](#)
- wald.logisticregs (Many Wald based tests for logistic and Poisson regressions with continuous predictors), [144](#)
- wald.mmpc (Constraint based feature selection algorithms), [75](#)
- wald.poissonregs (Many Wald based tests for logistic and Poisson regressions with continuous predictors), [144](#)
- wald.ses (Constraint based feature selection algorithms), [75](#)
- wald.univariateScore (MXM-internal), [153](#)
- wald.univregs (Univariate regression based tests), [201](#)
- wald.zipregs (Many simple zero inflated Poisson regressions), [142](#)

- waldBeta (Conditional independence test for proportions/percentages), [48](#)
- waldBinom (Conditional independence tests for sucess rates), [67](#)
- waldCR (Conditional independence tests for survival data), [69](#)
- waldER (Conditional independence tests for survival data), [69](#)
- waldGamma (Conditional independence tests for positive data), [64](#)
- waldIGreg (Conditional independence tests for positive data), [64](#)
- waldLogistic (Conditional independence test for binary, categorical or ordinal data), [35](#)
- waldmmpc.model (Regression models based on SES and MMPC outputs), [167](#)
- waldmmpc.path (MMPC solution paths for many combinations of hyper-parameters), [147](#)
- waldMMReg (Conditional independence tests for continous univariate

and multivariate data), [55](#)
waldNB (Conditional independence tests for count data), [58](#)
waldNormLog (Conditional independence tests for positive data), [64](#)
waldOrdinal (Conditional independence test for binary, categorical or ordinal data), [35](#)
waldPois (Conditional independence tests for count data), [58](#)
waldQBinom (Conditional independence test for binary, categorical or ordinal data), [35](#)
waldses.model (Regression models based on SES and MMPC outputs), [167](#)
waldTobit (Conditional independence tests for left censored data), [61](#)
waldWR (Conditional independence tests for survival data), [69](#)
waldZIP (Conditional independence tests for count data), [58](#)
weibreg.mxm (Cross-Validation for SES and MMPC), [96](#)
wr.fsreg (MXM-internal), [153](#)
wr.fsreg_2 (MXM-internal), [153](#)

Zero inflated Poisson regression, [206](#)
zip.bsreg (MXM-internal), [153](#)
zip.fsreg (MXM-internal), [153](#)
zip.fsreg_2 (MXM-internal), [153](#)
zip.mod, [144](#)
zip.mod (Zero inflated Poisson regression), [206](#)
zip.reg (Zero inflated Poisson regression), [206](#)
zip.regs, [207](#)
zip.regs (Many simple zero inflated Poisson regressions), [142](#)
zipmle.wei (MXM-internal), [153](#)
zipwei (MXM-internal), [153](#)