

Package ‘xergm’

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Title Extensions of Exponential Random Graph Models

Description Extensions of Exponential Random Graph Models (ERGM): Temporal Exponential Random Graph Models (TERGM), Generalized Exponential Random Graph Models (GERGM), and Temporal Network Autocorrelation Models (TNAM).

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Imports stats4, utils, methods, statnet, statnet.common, network, sna, ergm (>= 3.2.4), texreg (>= 1.34), Matrix, parallel, boot, coda, stats, ROCR, igraph, vegan, lme4 (>= 1.0), Rcpp (>= 0.11.0), speedglm

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xergm-package	<i>Extensions of Exponential Random Graph Models (ERGM)</i>
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Description

Extensions of Exponential Random Graph Models (ERGM).

Details

The **xergm** package implements extensions of exponential random graph models, in particular bootstrapped temporal ERGMs (`btergm`) and generalized ERGMs (`gergm`). The `btergm` function estimates temporal exponential random graph models (TERGM) by bootstrapped pseudolikelihood and provides various goodness-of-fit and diagnostic measures and plots. The `tnam` function estimates (temporal) network autocorrelation models. GERGMs have not been implemented yet. To display citation information, type `citation("xergm")`.

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See Also

[btergm](#) [mtergm](#) [preprocess](#) [timecov](#) [simulate.btergm](#) [gof](#) [interpret](#) [btergm-class](#) [tnam](#) [tnam-terms](#)
[knecht](#) [chemnet](#)

adjust	<i>Adjust the dimensions of a matrix to the dimensions of another matrix</i>
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Description

Adjust the dimensions of a matrix to the dimensions of another matrix.

Usage

```
adjust(source, target, remove = TRUE, add = TRUE, value = NA,  
        returnlabels = FALSE)
```

Arguments

source	A matrix, network, list or data.frame object or a vector which should be adjusted.
target	A matrix, network, list or data.frame object or a vector to which the source object is compared with regard to its labels.
remove	Should rows and columns that are not present in the target object be removed?
add	Should rows and columns that are present in the target object but not in the source object be added to the source object?
value	The value to be inserted if a new row or column is added. By default, new cells are filled with NA values, but other sensible values may include $-\text{Inf}$ or \emptyset .
returnlabels	Return a list of added and removed row and column labels rather than the actual matrix, vector, or network object?

Details

An adjacency matrix (the source matrix) is compared to another adjacency matrix (the target matrix) by matching the row or column labels. If the target matrix contains rows/columns which are not present in the source matrix, new rows and columns with the corresponding labels and NA values in the cells are inserted into the source matrix. If the source matrix contains rows/columns which are not present in the target matrix, these rows and columns are removed from the source matrix. In addition to adjacency matrices, two-mode matrices, network objects (also with vertex attributes), and vectors are supported.

See Also

[xrgm-package handleMissings preprocess](#)

btergm

TERGM by bootstrapped pseudolikelihood or MCMC MLE

Description

TERGM by bootstrapped pseudolikelihood or MCMC MLE.

Usage

```
btergm(formula, R = 500, offset = FALSE, parallel = c("no",
  "multicore", "snow"), ncpus = 1, cl = NULL,
  verbose = TRUE, ...)
```

```
mtergm(formula, offset = FALSE, constraints = ~ .,
  estimate = c("MLE", "MPLE"), verbose = TRUE, ...)
```

Arguments

formula	Formula for the TERGM. Model construction works like in the ergm package with the same model terms etc. (for a list of terms, see <code>help("ergm-terms")</code>). The networks to be modeled on the left-hand side of the equation must be given either as a list of network objects with more recent networks last (i.e., chronological order) or as a list of matrices with more recent matrices at the end. <code>dyadcov</code> and <code>edgescov</code> terms accept time-independent covariates (as network or matrix objects) or time-varying covariates (as a list of networks or matrices with the same length as the list of networks to be modeled).
R	Number of bootstrap replications. The higher the number of replications, the more accurate but also the slower is the estimation.
offset	If <code>offset = TRUE</code> is set, a list of offset matrices (one for each time step) with structural zeros is handed over to the pseudolikelihood routine. The offset matrices contain structural zeros where either the dependent networks or any of the covariates have missing nodes (if <code>auto.adjust = TRUE</code> is used). All matrices and network objects are inflated to the dimensions of the largest object, and the offset matrices inform the estimation routine which dyads are constrained to be absent (that is, fixed at $-\infty$). If <code>offset = FALSE</code> is set (the default behavior) and <code>auto.adjust</code> is switched on, all nodes that are not present across all covariates and networks within a time step are removed completely from the respective object(s) before estimation begins. This is computationally more efficient than using an offset matrix.
parallel	Use multiple cores in a computer or nodes in a cluster to speed up bootstrapping computations. The default value "no" means parallel computing is switched off. If "multicore" is used, the <code>mclapply</code> function from the parallel package (formerly in the multicore package) is used for parallelization. This should run on any kind of system except MS Windows because it is based on forking. It is usually the fastest type of parallelization. If "snow" is used, the <code>parLapply</code> function from the parallel package (formerly in the snow package) is used for

parallelization. This should run on any kind of system including cluster systems and including MS Windows. It is slightly slower than the former alternative if the same number of cores is used. However, "snow" provides support for MPI clusters with a large amount of cores, which **multicore** does not offer (see also the `c1` argument). The backend for the bootstrapping procedure is the **boot** package.

ncpus	The number of CPU cores used for parallel computing (only if <code>parallel</code> is activated). If the number of cores should be detected automatically on the machine where the code is executed, one can set <code>ncpus = detectCores()</code> after loading the parallel package. On some HPC clusters, the number of available cores is saved as an environment variable; for example, if MOAB is used, the number of available cores can sometimes be accessed using <code>Sys.getenv("MOAB_PROCCOUNT")</code> , depending on the implementation.
c1	An optional parallel or snow cluster for use if <code>parallel = "snow"</code> . If not supplied, a PSOCK cluster is created temporarily on the local machine.
constraints	Constraints of the ERGM. See ergm for details.
estimate	Estimation procedure of the ERGM. MCMC MLE by default, but MPLE with uncorrected standard errors is possible. See ergm for details.
verbose	Print details about data preprocessing and estimation settings.
...	Further arguments to be handed over to subroutines.

Details

The `btergm` function computes temporal exponential random graph models (TERGM) by bootstrapped pseudolikelihood, as described in Desmarais and Cranmer (2012).

The `mtergm` function computes TERGMs by MCMC MLE (or MPLE with uncorrected standard errors) via blockdiagonal matrices and structural zeros. The `btergm` function is faster than the `mtergm` function.

References

- Cranmer, Skyler J., Tobias Heinrich and Bruce A. Desmarais (2014): Reciprocity and the Structural Determinants of the International Sanctions Network. *Social Networks* 36(1): 5–22. <http://dx.doi.org/10.1016/j.socnet.2013.01.001>.
- Desmarais, Bruce A. and Skyler J. Cranmer (2012): Statistical Mechanics of Networks: Estimation and Uncertainty. *Physica A* 391: 1865–1876. <http://dx.doi.org/10.1016/j.physa.2011.10.018>.
- Desmarais, Bruce A. and Skyler J. Cranmer (2010): Consistent Confidence Intervals for Maximum Pseudolikelihood Estimators. *Neural Information Processing Systems 2010 Workshop on Computational Social Science and the Wisdom of Crowds*.

See Also

[xergm-package](#) [simulate.btergm](#) [gof](#) [knecht](#) [btergm-class](#) [preprocess](#) [timecov](#)

Examples

```

# A simple toy example:

library("statnet")
set.seed(5)

networks <- list()
for(i in 1:10){          # create 10 random networks with 10 actors
  mat <- matrix(rbinom(100, 1, .25), nrow = 10, ncol = 10)
  diag(mat) <- 0        # loops are excluded
  nw <- network(mat)    # create network object
  networks[[i]] <- nw   # add network to the list
}

covariates <- list()
for (i in 1:10) {       # create 10 matrices as covariate
  mat <- matrix(rnorm(100), nrow = 10, ncol = 10)
  covariates[[i]] <- mat # add matrix to the list
}

fit <- btergm(networks ~ edges + istar(2) +
  edgescov(covariates), R = 100)

summary(fit)           # show estimation results

## Not run:
# The same example using MCMC MLE:

fit2 <- mtergm(networks ~ edges + istar(2) +
  edgescov(covariates))

summary(fit2)

## End(Not run)

# For an example with real data, see help("knecht").

# Examples for parallel processing:

# Some preliminaries:
# - "Forking" means running the code on multiple cores in the same
#   computer. It's fast but consumes a lot of memory because all
#   objects are copied for each node. It's also restricted to
#   cores within a physical computer, i.e. no distribution over a
#   network or cluster. Forking does not work on Windows systems.
# - "MPI" is a protocol for distributing computations over many
#   cores, often across multiple physical computers/nodes. MPI
#   is fast and can distribute the work across hundreds of nodes
#   (but remember that R can handle a maximum of 128 connections,
#   which includes file access and parallel connections). However,
#   it requires that the Rmpi package is installed and that an MPI

```

```

# server is running (e.g., OpenMPI).
# - "PSOCK" is a TCP-based protocol. It can also distribute the
# work to many cores across nodes (like MPI). The advantage of
# PSOCK is that it can as well make use of multiple nodes within
# the same node or desktop computer (as with forking) but without
# consuming too much additional memory. However, the drawback is
# that it is not as fast as MPI or forking.
# The following code provides examples for these three scenarios.

# btergm works with clusters via the parallel package. That is, the
# user can create a cluster object (of type "PSOCK", "MPI", or
# "FORK") and supply it to the 'cl' argument of the 'btergm'
# function. If no cluster object is provided, btergm will try to
# create a temporary PSOCK cluster (if parallel = "snow") or it
# will use forking (if parallel = "multicore").

## Not run:

# To use a PSOCK cluster without providing an explicit cluster
# object:
require("parallel")
fit <- btergm(networks ~ edges + istar(2) + edgecov(covariates),
  R = 100, parallel = "snow", ncpus = 25)

# Equivalently, a PSOCK cluster can be provided as follows:
require("parallel")
cores <- 25
cl <- makeCluster(cores, type = "PSOCK")
fit <- btergm(networks ~ edges + istar(2) + edgecov(covariates),
  R = 100, parallel = "snow", ncpus = cores, cl = cl)
stopCluster(cl)

# Forking (without supplying a cluster object) can be used as
# follows.
require("parallel")
cores <- 25
fit <- btergm(networks ~ edges + istar(2) + edgecov(covariates),
  R = 100, parallel = "multicore", ncpus = cores)
stopCluster(cl)

# Forking (by providing a cluster object) works as follows:
require("parallel")
cores <- 25
cl <- makeCluster(cores, type = "FORK")
fit <- btergm(networks ~ edges + istar(2) + edgecov(covariates),
  R = 100, parallel = "snow", ncpus = cores, cl = cl)
stopCluster(cl)

# To use MPI, a cluster object MUST be created beforehand. In
# this example, a MOAB HPC server is used. It stores the number of
# available cores as a system option:
require("parallel")
cores <- as.numeric(Sys.getenv("MOAB_PROCCOUNT"))

```

```

cl <- makeCluster(cores, type = "MPI")
fit <- btergm(networks ~ edges + istar(2) + edgecov(covariates),
  R = 100, parallel = "snow", ncpus = cores, cl = cl)
stopCluster(cl)

# In the following example, the Rmpi package is used to create a
# cluster. This may not work on all systems; consult your local
# support staff or the help files on your HPC server to find out how
# to create a cluster object on your system.

# snow/Rmpi start-up
if (!is.loaded("mpi_initialize")) {
  library("Rmpi")
}
library(snow);

mpirank <- mpi.comm.rank (0)
if (mpirank == 0) {
  invisible(makeMPIcluster())
} else {
  sink (file="/dev/null")
  invisible(slaveLoop (makeMPImaster()))
  mpi.finalize()
  q()
}
# End snow/Rmpi start-up

cl <- getMPIcluster()

fit <- btergm(networks ~ edges + istar(2) + edgecov(covariates),
  R = 100, parallel = "snow", ncpus = 25, cl = cl)

## End(Not run)

```

btergm-class

Class "btergm"

Description

btergm objects result from the estimation of a bootstrapped TERGM via the btergm function in the **xergm** package. btergm objects contain the coefficients, the bootstrapping samples of the coefficients, the number of replications, the number of observations, the number of time steps, the original formula, and the response, effects and weights objects that were fed into the glm call for estimating the model.

Usage

```
## S4 method for signature 'btergm'
```



```

summary(object, level = 0.95, ...)

## S4 method for signature 'btergm'
show(object)

## S4 method for signature 'btergm'
nobs(object)

## S4 method for signature 'btergm'
coef(object, ...)

## S4 method for signature 'btergm'
confint(object, parm, level = 0.95, ...)

btergm.se(object, print = FALSE)

btergm.timesteps(object)

```

Arguments

object	A btergm object.
level	The significance level for computation of the confidence intervals. The default is 0.95 (that is, an alpha value of 0.05). Other common values include 0.999, 0.99, 0.9, and 0.5.
parm	Parameters (specified by integer position or character string).
print	Should the formatted coefficient table be printed to the R console along with significance stars (print = TRUE), or should the plain coefficient matrix be returned (print = FALSE)?
...	Further arguments to be handed over to subroutines.

Details

Various generic methods are available for btergm objects: The coef and show methods return the coefficients; the summary method gives a model summary. The nobs method returns the number of observations. The confint method returns confidence intervals from the bootstrap replications of btergm objects, and the user can specify the confidence level. The method returns a matrix with three columns: the estimate, the lower bound, and the upper bound of the confidence interval for each model term.

The btergm.se function computes standard errors and p values for btergm objects. It returns a matrix with four columns: the estimate, the standard error, the z value, and the p value for each model term. If the argument print = TRUE is used, the matrix is printed to the R console as a formatted coefficient matrix with significance stars instead. Note that confidence intervals are the preferred way of interpretation for bootstrapped TERGMs; standard errors are only accurate if the bootstrapped data are normally distributed, which is not always the case. Various methods for checking for normality for each model term are available, for example quantile-quantile plots (e.g., qqnorm(x@bootsamp[, 1])) for the first model term in the btergm object called x).

The `btergm.timesteps` function extracts the number of time steps from a `btergm` object. The number of time steps is the number of networks being modeled on the left-hand side of the model formula.

Slots

`coef`: Object of class "numeric". The coefficients.

`bootsamp`: Object of class "matrix". The bootstrapping sample.

`R`: Object of class "numeric". Number of replications.

`nobs`: Object of class "numeric". Number of observations.

`time.steps`: Object of class "numeric". Number of time steps.

`formula`: Object of class "formula". The original model formula (without indices for the time steps).

`response`: Object of class "integer". The response variable.

`effects`: Object of class "data.frame". The effects that went into the `glm` call.

`weights`: Object of class "integer". The weights of the observations.

`auto.adjust`: Object of class "logical". Indicates whether automatic adjustment of dimensions was done before estimation.

`offset`: Object of class "logical". Indicates whether an offset matrix with structural zeros was used.

`directed`: Object of class "logical". Are the dependent networks directed?

`bipartite`: Object of class "logical". Are the dependent networks bipartite?

References

Cranmer, Skyler J., Tobias Heinrich and Bruce A. Desmarais (2014): Reciprocity and the Structural Determinants of the International Sanctions Network. *Social Networks* 36(1): 5–22. <http://dx.doi.org/10.1016/j.socnet.2013.01.001>.

Desmarais, Bruce A. and Skyler J. Cranmer (2012): Statistical Mechanics of Networks: Estimation and Uncertainty. *Physica A* 391: 1865–1876. <http://dx.doi.org/10.1016/j.physa.2011.10.018>.

Desmarais, Bruce A. and Skyler J. Cranmer (2010): Consistent Confidence Intervals for Maximum Pseudolikelihood Estimators. *Neural Information Processing Systems 2010 Workshop on Computational Social Science and the Wisdom of Crowds*.

See Also

[xergm-package](#) [btergm](#) [simulate.btergm](#) [gof](#) [knecht](#) [getformula](#) [interpret](#)

chemnet	<i>German Toxic Chemicals Policy Network in the 1980s (Volker Schneider)</i>
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Description

The chemnet dataset contains network and attribute data and for the 30 most influential political actors with regard to toxic chemicals regulation in Germany in 1983/1984. While the original dataset contains up to 47 actors, this dataset contains the "complete influence core" of mutually relevant actors. The data are cross-sectional. There are no missing data; the response rate was 100 percent. Volker Schneider (University of Konstanz) collected this dataset for his dissertation (Schneider 1988). The dataset was later re-used for a journal publication on information exchange in policy networks (Leifeld and Schneider 2012).

The chemnet dataset contains network relations on political/strategic and technical/scientific information exchange, influence attribution, and membership in policy committees/forums, as well as nodal attributes on the actor type and opinions about the six most salient issues related to the political process that was leading to a new chemicals law at the time being.

Usage

`data(chemnet)`

Format

`pol` is a directed 30 x 30 adjacency matrix indicating which row actor sends political/strategic information to which column actor. 1 indicates an information exchange tie, and 0 indicates the absence of a network tie.

`sci to` is a directed 30 x 30 adjacency matrix indicating which row actor sends technical/scientific information to which column actor. 1 indicates an information exchange tie, and 0 indicates the absence of a network tie. In contrast to political/strategic information exchange, two separate survey questions were asked about technical/scientific information exchange: sending information, and receiving information. The two matrices contain the same relation but one time from the sender's perspective and one time from the receiver's perspective. By combining the two matrices, one can create a "confirmed" technical/scientific information exchange relation. The `sci to` matrix contains ties from the sender's perspective.

`sci from` is a directed 30 x 30 adjacency matrix indicating which row actor receives technical/scientific information from which column actor. 1 indicates an information exchange tie, and 0 indicates the absence of a network tie. In contrast to political/strategic information exchange, two separate survey questions were asked about technical/scientific information exchange: sending information, and receiving information. The two matrices contain the same relation but one time from the sender's perspective and one time from the receiver's perspective. By combining the two matrices, one can create a "confirmed" technical/scientific information exchange relation. The `sci from` matrix contains ties from the receiver's perspective.

`infrep` is a directed 30 x 30 adjacency matrix indicating which row actor deems which column actor "particularly influential". 1 indicates such a tie, and 0 indicates the absence of an influence attribution tie.

committee is a 30 x 20 two-mode (bipartite) network matrix indicating which row actor is a member of which policy committee/forum (as indicated by the column labels). 1 indicates a membership tie, and 0 indicates non-membership.

types is a one-column data.frame where the type variable contains the actor type of each node. The following values are possible:

- gov (government actor, e.g., a federal ministry)
- ig (interest group)
- io (international organization)
- par (political party)
- sci (scientific organization)

intpos is a 30 x 6 matrix containing the interest positions of the 30 political actors on the six most salient political issues related to a pending new chemicals law. -1 indicates a negative stance, i.e., the actor rejects the proposal; 1 indicates a positive stance, i.e., the actor supports the proposal; and 0 indicates a neutral or absent opinion.

Source

The data were collected using paper-based questionnaires. The questionnaires were administered in personal interviews (PAPI). Further information, including the actual survey, data on additional actors, the full names of the policy committees/forums, and the full list of unabbreviated actor names can be found online at <http://hdl.handle.net/1902.1/17004> in the replication archive of Leifeld and Schneider (2012).

- Replication archive: <http://hdl.handle.net/1902.1/17004>
- AJPS publication: <http://dx.doi.org/10.1111/j.1540-5907.2011.00580.x>

The dataset is publicly available. Questions about the data or the original study should be directed to Volker Schneider (volker.schneider@uni-konstanz.de), the author of the original study and person who collected the data.

References

Leifeld, Philip and Volker Schneider (2012): Information Exchange in Policy Networks. *American Journal of Political Science* 53(3): 731–744. <http://dx.doi.org/10.1111/j.1540-5907.2011.00580.x>.

Schneider, Volker (1988): *Politiknetzwerke der Chemikalienkontrolle. Eine Analyse einer transnationalen Politikentwicklung*. Walter de Gruyter: Berlin/New York.

Schneider, Volker and Philip Leifeld (2009): Ueberzeugungssysteme, Diskursnetzwerke und politische Kommunikation: Ein zweiter Blick auf die deutsche Chemikalienkontrolle der 1980er Jahre. In: Volker Schneider, Frank Janning, Philip Leifeld and Thomas Malang (editors): *Politiknetzwerke. Modelle, Anwendungen und Visualisierungen*. Pages 139–158. Wiesbaden: VS Verlag fuer Sozialwissenschaften. http://dx.doi.org/10.1007%2F978-3-531-91883-9_6.

Examples

```

## Not run:
# Replication code for Leifeld and Schneider (2012), AJPS.
# Note that the estimates can only be reproduced approximately
# due to internal changes in the statnet package.

# preparatory steps
library("statnet")
library("xergm")
library("texreg")
seed <- 12345
set.seed(seed)
data("chemnet")

# create confirmed network relation
sci <- scito * t(scifrom) # equation 1 in the AJPS paper
prefsim <- dist(intpos, method = "euclidean") # equation 2
prefsim <- max(prefsim) - prefsim # equation 3
prefsim <- as.matrix(prefsim)
committee <- committee
diag(committee) <- 0 # the diagonal has no meaning
types <- types[, 1] # convert to vector

# create network objects and store attributes
nw.pol <- network(pol) # political/strategic information exchange
set.vertex.attribute(nw.pol, "orgtype", types)
set.vertex.attribute(nw.pol, "betweenness",
  betweenness(nw.pol)) # centrality

nw.sci <- network(sci) # technical/scientific information exchange
set.vertex.attribute(nw.sci, "orgtype", types)
set.vertex.attribute(nw.sci, "betweenness",
  betweenness(nw.sci)) # centrality

# ERGM: model 1 in the AJPS paper; only preference similarity
model1 <- ergm(nw.pol ~ edges + edgescov(prefsim),
  control = control.ergm(seed = seed))
summary(model1)

# ERGM: model 2 in the AJPS paper; complete model
model2 <- ergm(nw.pol ~
  edges +
  edgescov(prefsim) +
  mutual +
  nodemix("orgtype", base = -7) +
  nodeifactor("orgtype", base = -1) +
  nodeofactor("orgtype", base = -5) +
  edgescov(committee) +
  edgescov(nw.sci) +
  edgescov(infrep) +
  gwesp(0.1, fixed = TRUE) +
  gwesp(0.1, fixed = TRUE),

```

```

    control = control.ergm(seed = seed)
  )
summary(model2)

# ERGM: model 3 in the AJPS paper; only preference similarity
model3 <- ergm(nw.sci ~ edges + edgescov(prefsim),
  control = control.ergm(seed = seed))
summary(model3)

# ERGM: model 4 in the AJPS paper; complete model
model4 <- ergm(nw.sci ~
  edges +
  edgescov(prefsim) +
  mutual +
  nodemix("orgtype", base = -7) +
  nodeifactor("orgtype", base = -1) +
  nodeofactor("orgtype", base = -5) +
  edgescov(committee) +
  edgescov(nw.pol) +
  edgescov(infrep) +
  gwesp(0.1, fixed = TRUE) +
  gwesp(0.1, fixed = TRUE),
  control = control.ergm(seed = seed)
)
summary(model4)

# regression table using the texreg package
screenreg(list(model1, model2, model3, model4))

# goodness of fit using the xergm package
gof2 <- gof(model2, roc = FALSE, pr = FALSE)
gof2 # print gof output
plot(gof2) # visual inspection of GOF

gof4 <- gof(model4, roc = FALSE, pr = FALSE)
gof4
plot(gof4)

# MCMC diagnostics
pdf("diagnostics2.pdf")
mcmc.diagnostics(model2)
dev.off()

pdf("diagnostics4.pdf")
mcmc.diagnostics(model4)
dev.off()

## End(Not run)

```

Description

Extract the model formula from ergm or btergm objects.

Usage

```
getformula(x)

## S4 method for signature 'ergm'
getformula(x)

## S4 method for signature 'btergm'
getformula(x)
```

Arguments

x A model object, for example a btergm or an ergm object.

Details

Extract the model formula from ergm or btergm objects.

See Also

[gof](#)

gof-methods	<i>Conduct Goodness-of-Fit Diagnostics on ERGMs, TERGMs, SAOMs, and logit models</i>
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Description

Assess goodness of fit and degeneracy of btergm and other network models.

Usage

```
## S4 method for signature 'btergm'
gof(object, target = NULL,
     formula = getformula(object), nsim = 100, MCMC.interval = 1000,
     MCMC.burnin = 10000, parallel = c("no", "MPI", "SOCK"),
     ncpus = 1, cl = NULL, classicgof = TRUE, rocprgof = TRUE,
     checkdegeneracy = TRUE, statistics = c("dsp", "esp", "geodist",
     "degree", "idegree", "odegree", "kstar", "istar", "ostar"),
     pr.impute = "poly4", verbose = TRUE, ...)

## S4 method for signature 'ergm'
gof(object, target = NULL,
     formula = getformula(object), nsim = 100, MCMC.interval = 1000,
```

```

MCMC.burnin = 10000, parallel = c("no", "MPI", "SOCK"),
ncpus = 1, cl = NULL, classicgof = TRUE, rocprgof = TRUE,
checkdegeneracy = TRUE, statistics = c("dsp", "esp", "geodist",
"degree", "idegree", "odegree", "kstar", "istar", "ostar"),
pr.impute = "poly4", verbose = TRUE, ...)

## S4 method for signature 'sienaAlgorithm'
gof(object, siena.data, siena.effects,
predict.period = NULL, nsim = 50, parallel = c("no", "multicore",
"snow"), ncpus = 1, cl = NULL, target.na = NA,
target.na.method = "remove", target.structzero = 10,
classicgof = TRUE, rocprgof = TRUE, statistics = c("dsp", "esp",
"geodist", "degree", "idegree", "odegree", "kstar", "istar",
"ostar"), pr.impute = "poly4", ...)

## S4 method for signature 'sienaModel'
gof(object, siena.data, siena.effects,
predict.period = NULL, nsim = 50, parallel = c("no", "multicore",
"snow"), ncpus = 1, cl = NULL, target.na = NA,
target.na.method = "remove", target.structzero = 10,
classicgof = TRUE, rocprgof = TRUE, statistics = c("dsp", "esp",
"geodist", "degree", "idegree", "odegree", "kstar", "istar",
"ostar"), pr.impute = "poly4", ...)

## S4 method for signature 'network'
gof(object, covariates, coef, target = NULL,
nsim = 100, mcmc = FALSE, MCMC.interval = 1000,
MCMC.burnin = 10000, parallel = c("no", "MPI", "SOCK"),
ncpus = 1, cl = NULL, classicgof = TRUE, rocprgof = TRUE,
statistics = c("dsp", "esp", "geodist", "degree", "idegree",
"odegree", "kstar", "istar", "ostar"), pr.impute = "poly4",
verbose = TRUE, ...)

## S4 method for signature 'matrix'
gof(object, covariates, coef, target = NULL,
nsim = 100, mcmc = FALSE, MCMC.interval = 1000,
MCMC.burnin = 10000, parallel = c("no", "MPI", "SOCK"),
ncpus = 1, cl = NULL, classicgof = TRUE, rocprgof = TRUE,
statistics = c("dsp", "esp", "geodist", "degree", "idegree",
"odegree", "kstar", "istar", "ostar"), pr.impute = "poly4",
verbose = TRUE, ...)

```

Arguments

object	A <code>btergm</code> , <code>ergm</code> , <code>sienaAlgorithm</code> , or <code>sienaModel</code> object (for the <code>btergm</code> , <code>ergm</code> , <code>sienaAlgorithm</code> , and <code>sienaModel</code> methods, respectively). Or a network object or matrix (for the <code>network</code> and <code>matrix</code> methods, respectively).
siena.data	An object of the class <code>siena</code> , which is usually created using the <code>sienaDataCreate</code>

	function in the <code>RSiena</code> package.
<code>siena.effects</code>	An object of the class <code>sienaEffects</code> , which is usually created using the <code>getEffects()</code> and the <code>includeEffects()</code> function in the <code>RSiena</code> package.
<code>predict.period</code>	Which time period should be predicted? By default, the last time period is predicted based on the last simulation of the second-last time period. The time period can be provided as a numeric, e.g., <code>predict.period = 4</code> for predicting the fourth network.
<code>target</code>	A network or list of networks to which the simulations are compared. If left empty, the original networks from the <code>btergm</code> object <code>x</code> are used as observed networks.
<code>formula</code>	A model formula from which networks are simulated for comparison. By default, the formula from the <code>btergm</code> object <code>x</code> is used. It is possible to hand over a formula with only a single response network and/or dyad or edge covariates or with lists of response networks and/or covariates. It is also possible to use indices like <code>networks[[4]]</code> or <code>networks[3:5]</code> inside the formula.
<code>nsim</code>	The number of networks to be simulated at each time step. Example: If there are six time steps in the <code>formula</code> and <code>nsim = 100</code> , a total of 600 new networks is simulated.
<code>MCMC.interval</code>	Internally, this package uses the simulation facilities of the ergm package to create new networks against which to compare the original network(s) for goodness-of-fit assessment. This argument sets the MCMC interval to be passed over to the simulation command. The default value is <code>1000</code> , which means that every 1000th simulation outcome from the MCMC sequence is used. There is no general rule of thumb on the selection of this parameter, but if the results look suspicious (e.g., when the model fit is perfect), increasing this value may be helpful.
<code>MCMC.burnin</code>	Internally, this package uses the simulation facilities of the ergm package to create new networks against which to compare the original network(s) for goodness-of-fit assessment. This argument sets the MCMC burnin to be passed over to the simulation command. The default value is <code>10000</code> . There is no general rule of thumb on the selection of this parameter, but if the results look suspicious (e.g., when the model fit is perfect), increasing this value may be helpful.
<code>parallel</code>	Use multiple cores in a computer or nodes in a cluster to speed up the simulations. The default value <code>"no"</code> means parallel computing is switched off. If <code>"multicore"</code> is used (only available for <code>sienaAlgorithm</code> and <code>sienaModel</code> objects), the <code>mclapply</code> function from the parallel package (formerly in the multicore package) is used for parallelization. This should run on any kind of system except MS Windows because it is based on forking. It is usually the fastest type of parallelization. If <code>"snow"</code> is used (only available for <code>sienaAlgorithm</code> and <code>sienaModel</code> objects), the <code>parLapply</code> function from the parallel package (formerly in the snow package) is used for parallelization. This should run on any kind of system including cluster systems and including MS Windows. It is slightly slower than the former alternative if the same number of cores is used. However, <code>"snow"</code> provides support for MPI clusters with a large amount of cores, which multicore does not offer (see also the <code>c1</code> argument). If <code>"MPI"</code> is used (only available for <code>btergm</code> and <code>ergm</code> objects), MPI parallelization as

implemented in the **ergm** package is used. And if "SOCK" is used (only available for **btergm** and **ergm** objects), a SOCK cluster as implemented in the **ergm** package (via the deprecated **snow** package) is used for parallelization. Note that "multicore" and "SOCK" will only work if all cores are on the same node. For example, if there are three nodes with eight cores each, a maximum of eight CPUs can be used.

<code>ncpus</code>	The number of CPU cores used for parallel simulations (only if <code>parallel</code> is activated). If the number of cores should be detected automatically on the machine where the code is executed, one can try the <code>detectCores()</code> function from the parallel package. On some HPC clusters, the number of available cores is saved as an environment variable; for example, if MOAB is used, the number of available cores can sometimes be accessed using <code>Sys.getenv("MOAB_PROCCOUNT")</code> , depending on the implementation. Note that the maximum number of connections in a single R session (i.e., to other cores or for opening files etc.) is 128, so fewer than 128 cores should be used at a time.
<code>cl</code>	An optional parallel or snow cluster for use if <code>parallel = "snow"</code> . If not supplied, a cluster on the local machine is created temporarily.
<code>target.na</code>	Which value was used for missing data in the dependent variable?
<code>target.na.method</code>	How should missing data be handled when comparing the simulations to the empirical (= observed) network? Two options are possible: remove drops nodes with missing ties both from the simulations (after running the simulations) and from the observed network before the comparison. <code>fillmode</code> replaces missing values by the mode of the network matrix (usually \emptyset).
<code>target.structzero</code>	Which value was used for structural zeroes (usually nodes which have dropped out of the network or have not yet joined the network) in the dependent variable? These nodes are removed from the observed network and the simulations before comparison.
<code>classicgof</code>	If <code>classicgof = TRUE</code> is set, the classic statnet-style goodness-of-fit comparison is conducted. This means that shared-partner statistics, the geodesic distance distribution and the degree distribution are compared between observed and simulated networks. The results can be plotted as boxplots or printed as tables. Note that the <code>classicgof</code> , <code>rocprgof</code> and <code>checkdegeneracy</code> arguments can be used together. In that case, the resulting btergm object will contain all three types of GOF/degeneracy assessment.
<code>rocprgof</code>	If <code>rocprgof = TRUE</code> is set, the coordinates of ROC and PR curves as well as the AUC measure are stored in the resulting btergm object. The results can be plotted as curves or printed as tables. Note that the <code>classicgof</code> , <code>rocprgof</code> and <code>checkdegeneracy</code> arguments can be used together. In that case, the resulting btergm object will contain all three types of GOF/degeneracy assessment.
<code>checkdegeneracy</code>	If <code>checkdegeneracy = TRUE</code> is set, the global statistics of the observed and simulated networks are compared for each observed time step separately. Frequent significant deviations indicate degeneracy. The results can be printed as tables. Note that the <code>classicgof</code> , <code>rocprgof</code> and <code>checkdegeneracy</code> arguments

	can be used together. In that case, the resulting <code>btergm</code> object will contain all three types of GOF/degeneracy assessment.
<code>statistics</code>	A character vector of auxiliary statistics used for comparison of observed and simulated networks. Valid values are "dsp", "esp", "geodist", "degree", "idegree", "odegree", "kstar", "istar", and "ostar". By default, all of these statistics are used.
<code>pr.impute</code>	In some cases, the first precision value of the precision-recall curve is undefined. The <code>pr.impute</code> argument serves to impute this missing value to ensure that the AUC-PR value is not severely biased. Possible values are "no" for no imputation, "one" for using a value of 1.0, "second" for using the next (= adjacent) precision value, "poly1" for fitting a straight line through the remaining curve to predict the first value, "poly2" for fitting a second-order polynomial curve etc. until "poly9" Warning: this is a pragmatic solution. Please double-check whether the imputation makes sense. This can be checked by plotting the resulting <code>btergm</code> object and using the <code>pr.poly</code> argument to plot the predicted curve on top of the actual PR curve.
<code>covariates</code>	A list of matrices or network objects that serve as covariates for the dependent network. The covariates in this list are automatically added to the formula as <code>edgecov</code> terms.
<code>coef</code>	A vector of coefficients.
<code>mcmc</code>	Should <code>statnet</code> 's MCMC methods be used for simulating new networks? If <code>mcmc = FALSE</code> , new networks are simulated based on predicted tie probabilities of the regression equation.
<code>verbose</code>	Print details?
<code>...</code>	Arbitrary further arguments are handed over to the <code>simulate.formula</code> function or the <code>siena07</code> function. For details, refer to the help page of these functions.

Details

The generic `gof` function provides goodness-of-fit measures and degeneracy checks for `btergm`, `ergm`, `SAOM`, and custom dyadic-independent models. Three different types of GOF/degeneracy assessment are possible with this function:

- (1) Classic `statnet`-type GOF assessment by comparing summary statistics of observed and simulated networks. The `gof` function has six built-in statistics: dyad-wise shared partners (`dsp`), edge-wise shared partners (`esp`), degree (for undirected networks only), indegree (for directed networks only), outdegree (for directed networks only), and geodesic distances. The comparison can be plotted using boxplots for the simulations and lines for the observed network(s) or printed using t-tests (testing whether simulated and observed networks are significantly different for all values in the distributions of the summary statistics).
- (2) An assessment of the classification performance using receiver operating characteristics (ROC) and precision-recall (PR) curves as well as the area under the curve (AUC) for the ROC curve.
- (3) For bootstrapped TERGMs: A degeneracy check by comparing the global statistics of simulated networks to those of the observed networks at each observed time step. If the global statistics differ significantly, this is indicated by small p values. If there are many significant results, this indicates degeneracy.

For all three types of GOF assessment, by default, in-sample predictive performance is assessed by comparing all observed networks to all simulations from the same networks (just like in the **ergm** package, but aggregated over several time steps). If an observed network or a list of observed networks is provided as the `target` argument, the simulations are compared to these networks instead. This is useful for out-of-sample prediction. If a formula is provided, the simulations are based on the networks and covariates specified in the formula. This is helpful in situations where complex out-of-sample predictions have to be evaluated. A usage scenario could be to simulate from a network at time t (provided through the `formula` argument) and compare to an observed network at time $t + 1$ (the `target` argument). This can be done, for example, to assess predictive performance between time steps of the original networks, or to check whether the model performs well with regard to a newly measured network given the old data from the previous time step.

Predictive fit can also be assessed for stochastic actor-oriented models (SAOM) as implemented in the **RSiena** package. After compiling the usual objects (model, data, effects), one of the time steps can be predicted based on the previous time step and the SAOM using the `sienaAlgorithm` (for **RSiena** \geq 1.1-227) or `sienaModel` (for **RSiena** $<$ 1.1-227) method of the `gof` function.

The `gof` methods for networks and matrices serve to assess the goodness of fit of a dyadic-independence model. To do this, the method requires a vector of coefficients (one coefficient for the intercept or edges term and one coefficient for each covariate), a list of covariates (in matrix or network shape), and a dependent network or matrix. This is useful for assessing the goodness of fit of QAP-adjusted logistic regression models (as implemented in the `netlogit` function in the **sna** package) or other dyadic-independence models, such as models fitted using `glm`. Note that this method only works with cross-sectional models and does not accept lists of networks as input data.

See also the [plot.btergm](#) help page for details on the plotting and printing options for GOF assessment.

See Also

[xergm-package](#) [btergm](#) [simulate.btergm](#) [simulate.formula](#) [plot.btergm](#) [gof](#)

handleMissings

Handle missing data in matrices.

Description

Handle missing data in matrices.

Usage

```
handleMissings(mat, na = NA, method = "remove", logical = FALSE)
```

Arguments

<code>mat</code>	A matrix object.
<code>na</code>	The value that missing data are coded as. Usually NA, sometimes 9 or 10.

method	What should be done with the missing data? If <code>method = "remove"</code> is set, the function determines how many missing entries are in each row and column and iteratively removes rows or columns with the largest amount of missing data until no missing data are left in the matrix. If <code>method = "fillmode"</code> is set, the modal value of the matrix is identified (usually 0 in network matrices) and missing cells are imputed by filling in this modal value. <code>method = "zero"</code> replaces NAs by 0s.
logical	Return a matrix with logical values indicating which cells should be removed? By default the manipulated matrix is returned.

Details

This function deals with missing data in matrices or network objects used for inferential network analysis. It can either remove missing rows and/or columns iteratively (rows and columns with more NA values first, then successively rows and columns with fewer NA entries) or replace missing values by the modal value of the matrix or by 0. The function can return either the manipulated matrix or a matrix with logical values indicating which of the cells should be removed.

See Also

[xergm-package adjust preprocess](#)

interpret

Interpretation functions for ergm and btergm objects

Description

Interpretation functions for ergm and btergm objects.

Usage

```
interpret(object, ...)

## S4 method for signature 'ergm'
interpret(object, formula = object$formula,
  coefficients = coef(object), network = eval(parse(text =
  deparse(formula[[2]]))), type = "tie", i, j, ...)

## S4 method for signature 'btergm'
interpret(object, formula = object@formula,
  coefficients = coef(object), network = eval(parse(text =
  deparse(formula[[2]]))), type = "tie", i, j,
  t = 1:length(network), ...)
```

Arguments

object	An <code>ergm</code> or <code>btergm</code> object.
formula	The formula to be used for computing probabilities. By default, the formula embedded in the model object is retrieved and used.
coefficients	The estimates on which probabilities should be based. By default, the coefficients from the model object are retrieved and used. Custom coefficients can be handed over, for example, in order to compare versions of the model where the reciprocity term is fixed at 0 versus versions of the model where the reciprocity term is left as in the empirical result. This is one of the examples described in Desmarais and Cranmer (2012).
network	The response network on which probabilities are based. Depending on whether the function is applied to an <code>ergm</code> or <code>btergm</code> object, this can be either a single network or a list of networks.
type	If <code>type = "tie"</code> is used, probabilities at the edge level are computed. For example, what is the probability of a specific node i to be connected to a specific node j given the rest of the network and given the model? If <code>type = "dyad"</code> is used, probabilities at the dyad level are computed. For example, what is the probability that node i is connected to node j but not vice-versa, or what is the probability that nodes i and j and mutually connected in a directed network? If <code>type = "node"</code> is used, probabilities at the node level are computed. For example, what is the probability that node i is connected to a set of three other j nodes given the rest of the network and the model?
i	A single (sender) node i or a set of (sender) nodes i . If <code>type = "node"</code> is used, this can be more than one node and should be provided as a vector. The i argument can be either provided as the index of the node in the sociomatrix (e.g., the fourth node would be $i = 4$) or the row name of the node in the sociomatrix (e.g., $i = \text{"Peter"}$). If more than one node is provided and <code>type = "node"</code> , there can be only one (receiver) node j . The i and j arguments are used to specify for which nodes probabilities should be computed. For example, what is the probability that $i = 4$ is connected to $i = 7$?
j	A single (receiver) node j or a set of (receiver) nodes j . If <code>type = "node"</code> is used, this can be more than one node and should be provided as a vector. The j argument can be either provided as the index of the node in the sociomatrix (e.g., the fourth node would be $j = 4$) or the row name of the node in the sociomatrix (e.g., $j = \text{"Mary"}$). If more than one node is provided and <code>type = "node"</code> , there can be only one (sender) node i . The i and j arguments are used to specify for which nodes probabilities should be computed. For example, what is the probability that $i = 4$ is connected to $i = 7$?
t	A vector of (numerical) time steps for which the probabilities should be computed. This only applies to <code>btergm</code> objects because <code>ergm</code> objects are by definition based on a single time step. By default, all available time steps are used. It is, for example, possible to compute probabilities only for a single time step by specifying, e.g., $t = 5$ in order to compute probabilities for the fifth response network.
...	Further arguments to be handed over to subroutines.

Details

The `interpret` function facilitates interpretation of ERGMs and TERGMs at the micro level, as described in Desmarais and Cranmer (2012). There are generic methods which work with `ergm` objects and `btergm` objects. The function can be used to interpret these models at the tie or edge level, dyad level, and block level.

For example, what is the probability that two specific nodes i (the sender) and node j (the receiver) are connected given the rest of the network and given the model? Or what is the probability that any two nodes are tied at $t = 2$ if they were tied (or disconnected) at $t = 1$ (i.e., what is the amount of tie stability)? These tie- or edge-level questions can be answered if the `type = "tie"` argument is used.

Another example: What is the probability that node i has a tie to node j but not vice-versa? Or that i and j maintain a reciprocal tie? Or that they are disconnected? How much more or less likely are i and j reciprocally connected if the `mutual` term in the model is fixed at 0 (compared to the model that includes the estimated parameter for reciprocity)? See example below. These dyad-level questions can be answered if the `type = "dyad"` argument is used.

Or what is the probability that a specific node i is connected to nodes j_1 and j_2 but not to j_3 and j_4 ? And how likely is any node i to be connected to exactly four j nodes? These node-level questions (focusing on the ties of node i or node j) can be answered by using the `type = "node"` argument.

References

Desmarais, Bruce A. and Skyler J. Cranmer (2012): Micro-Level Interpretation of Exponential Random Graph Models with Application to Estuary Networks. *The Policy Studies Journal* 40(3): 402–434.

See Also

[xergm-package btergm btergm.timesteps](#)

Examples

```
##### The following example is a TERGM adaptation of the #####
##### dyad-level example provided in figure 5(c) on page #####
##### 424 of Desmarais and Cranmer (2012) in the PSJ. At #####
##### each time step, it compares dyadic probabilities #####
##### (no tie, unidirectional tie, and reciprocal tie #####
##### probability) between a fitted model and a model #####
##### where the reciprocity effect is fixed at 0 based #####
##### on 20 randomly selected dyads per time step. The #####
##### results are visualized using a grouped bar plot. #####

## Not run:
# create toy dataset and fit a model
networks <- list()
for (i in 1:3) {           # create 3 random networks with 10 actors
  mat <- matrix(rbinom(100, 1, 0.25), nrow = 10, ncol = 10)
  diag(mat) <- 0          # loops are excluded
  nw <- network(mat)     # create network object
  networks[[i]] <- nw    # add network to the list
}
```

```

}
fit <- btergm(networks ~ edges + istar(2) + mutual, R = 200)

# extract coefficients and create null hypothesis vector
null <- coef(fit) # estimated coeffs
null[3] <- 0      # set mutual term = 0

# sample 20 dyads per time step and compute probability ratios
probabilities <- matrix(nrow = 9, ncol = length(networks))
# nrow = 9 because three probabilities + upper and lower CIs
colnames(probabilities) <- paste("t =", 1:length(networks))
for (t in 1:length(networks)) {
  d <- dim(as.matrix(networks[[t]])) # how many row and column nodes?
  size <- d[1] * d[2]                # size of the matrix
  nw <- matrix(1:size, nrow = d[1], ncol = d[2])
  nw <- nw[lower.tri(nw)]            # sample only from lower triangle b/c
  samp <- sample(nw, 20)             # dyadic probabilities are symmetric
  prob.est.00 <- numeric(0)
  prob.est.01 <- numeric(0)
  prob.est.11 <- numeric(0)
  prob.null.00 <- numeric(0)
  prob.null.01 <- numeric(0)
  prob.null.11 <- numeric(0)
  for (k in 1:20) {
    i <- arrayInd(samp[k], d)[1, 1] # recover 'i's and 'j's from sample
    j <- arrayInd(samp[k], d)[1, 2]
    # run interpretation function with estimated coeffs and mutual = 0:
    int.est <- interpret(fit, type = "dyad", i = i, j = j, t = t)
    int.null <- interpret(fit, coefficients = null, type = "dyad",
      i = i, j = j, t = t)
    prob.est.00 <- c(prob.est.00, int.est[[1]][1, 1])
    prob.est.11 <- c(prob.est.11, int.est[[1]][2, 2])
    mean.est.01 <- (int.est[[1]][1, 2] + int.est[[1]][2, 1]) / 2
    prob.est.01 <- c(prob.est.01, mean.est.01)
    prob.null.00 <- c(prob.null.00, int.null[[1]][1, 1])
    prob.null.11 <- c(prob.null.11, int.null[[1]][2, 2])
    mean.null.01 <- (int.null[[1]][1, 2] + int.null[[1]][2, 1]) / 2
    prob.null.01 <- c(prob.null.01, mean.null.01)
  }
  prob.ratio.00 <- prob.est.00 / prob.null.00 # ratio of est. and null hyp
  prob.ratio.01 <- prob.est.01 / prob.null.01
  prob.ratio.11 <- prob.est.11 / prob.null.11
  probabilities[1, t] <- mean(prob.ratio.00) # mean estimated 00 tie prob
  probabilities[2, t] <- mean(prob.ratio.01) # mean estimated 01 tie prob
  probabilities[3, t] <- mean(prob.ratio.11) # mean estimated 11 tie prob
  ci.00 <- t.test(prob.ratio.00, conf.level = 0.99)$conf.int
  ci.01 <- t.test(prob.ratio.01, conf.level = 0.99)$conf.int
  ci.11 <- t.test(prob.ratio.11, conf.level = 0.99)$conf.int
  probabilities[4, t] <- ci.00[1]           # lower 00 conf. interval
  probabilities[5, t] <- ci.01[1]           # lower 01 conf. interval
  probabilities[6, t] <- ci.11[1]           # lower 11 conf. interval
  probabilities[7, t] <- ci.00[2]           # upper 00 conf. interval
  probabilities[8, t] <- ci.01[2]           # upper 01 conf. interval
}

```



```

    probabilities[9, t] <- ci.11[2]          # upper 11 conf. interval
  }

# create barplots from probability ratios and CIs
require("gplots")
bp <- barplot2(probabilities[1:3, ], beside = TRUE, plot.ci = TRUE,
  ci.l = probabilities[4:6, ], ci.u = probabilities[7:9, ],
  col = c("tan", "tan2", "tan3"), ci.col = "grey40",
  xlab = "Dyadic tie values", ylab = "Estimated Prob./Null Prob.")
mtext(1, at = bp, text = c("(0,0)", "(0,1)", "(1,1)"), line = 0, cex = 0.5)

## End(Not run)

```

knecht	<i>Longitudinal classroom friendship network and behavior (Andrea Knecht)</i>
--------	---

Description

The Knecht dataset contains the friendship network of 26 pupils in a Dutch school class measured at four time points along with several demographic and behavioral covariates like age, sex, ethnicity, religion, delinquency, alcohol consumption, primary school co-attendance, and school advice. Some of these covariates are constant while others vary over time.

The full dataset (see Knecht 2006 and 2008) contains a large number of classrooms while the dataset presented here is an excerpt based on one single classroom. This excerpt was first used in a tutorial for the software **Siena** and the corresponding R package **RSiena** (Snijders, Steglich and van de Bunt 2010). The following description was largely copied from the original data description provided on the homepage of the **Siena** project (see below for the URL).

The data were collected between September 2003 and June 2004 by Andrea Knecht, supervised by Chris Baerveldt, at the Department of Sociology of the University of Utrecht (NL). The entire study is reported in Knecht (2008). The project was funded by the Netherlands Organisation for Scientific Research NWO, grant 401-01-554. The 26 students were followed over their first year at secondary school during which friendship networks as well as other data were assessed at four time points at intervals of three months. There were 17 girls and 9 boys in the class, aged 11–13 at the beginning of the school year. Network data were assessed by asking students to indicate up to twelve classmates which they considered good friends. Delinquency is defined as a rounded average over four types of minor delinquency (stealing, vandalism, graffiti, and fighting), measured in each of the four waves of data collection. The five-point scale ranged from ‘never’ to ‘more than 10 times’, and the distribution is highly skewed. In a range of 1–5, the mode was 1 at all four waves, the average rose over time from 1.4 to 2.0, and the value 5 was never observed.

Usage

```
data(knecht)
```

Format

Note: the data have to be transformed before they can be used with the `btergm` function from the `xergm` package (see examples below).

`friendship` is a list of adjacency matrices at four time points, containing friendship nominations of the column node by the row node. The following values are used: 0 = no, 1 = yes, `NA` = missing, 10 = not a member of the classroom (structural zero).

`demographics` is a data frame with 26 rows (the pupils) and four demographic variables about the pupils:

- sex (1 = girl, 2 = boy)
- age (in years)
- ethnicity (1 = Dutch, 2 = other, 0 = missing)
- religion (1 = Christian, 2 = non-religious, 3 = non-Christian religion, 0 = missing)

`primary` is a 26 x 26 matrix indicating whether two pupils attended the same primary school. 0 = no, 1 = yes.

`delinquency` is a data frame with 26 rows (the pupils) and four columns (the four time steps). It contains the rounded average of four items (stealing, vandalizing, fighting, graffiti). Categories: frequency over last three months, 1 = never, 2 = once, 3 = 2–4 times, 4 = 5–10 times, 5 = more than 10 times; 0 = missing.

`alcohol` is a data frame with 26 rows (the pupils) and 3 columns (waves 2, 3, and 4). It contains data on alcohol use (“How often did you drink alcohol with friends in the last three months?”). Categories: 1 = never, 2 = once, 3 = 2–4 times, 4 = 5–10 times, 5 = more than 10 times; 0 = missing.

`advice` is a data frame with one variable, “school advice”, the assessment given at the end of primary school about the school capabilities of the pupil (4 = low, 8 = high, 0 = missing)

Source

The data were gathered by Andrea Knecht, as part of her PhD research, building on methods developed by Chris Baerveldt, initiator and supervisor of the project. The project is funded by the Netherlands Organisation for Scientific Research NWO, grant 401-01-554, and is part of the research program “Dynamics of Networks and Behavior” with principle investigator Tom A. B. Snijders.

- Complete original data: <https://easy.dans.knaw.nl/ui/datasets/id/easy-dataset:48665>
- This excerpt in Siena format: <http://www.stats.ox.ac.uk/~snijders/siena/klas12b.zip>
- Siena dataset description: http://www.stats.ox.ac.uk/~snijders/siena/tutorial2010_data.htm

Permission to redistribute this dataset along with the `xergm` package was granted by Andrea Knecht on April 17, 2014. Questions about the data or the original study should be directed to her.

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- Knecht, Andrea (2008): *Friendship Selection and Friends' Influence. Dynamics of Networks and Actor Attributes in Early Adolescence*. PhD Dissertation, University of Utrecht. <http://igitur-archive.library.uu.nl/dissertations/2008-0125-200537/>.
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Examples

```
## Not run:
# for more details, see the package vignette -- run vignette("xergm")

require("texreg")
require("sna")
require("xergm")
require("RSiena")
data("knecht")

# what do the networks look like? plot them!
par(mfrow = c(2, 2), mar = c(0, 0, 1, 0))
for (i in 1:length(friendship)) {
  plot(network(friendship[[i]]), main = paste("t =", i),
        usearrows = FALSE, edge.col = "grey50")
}

# =====
# Preprocess data and estimate TERGMs using the btergm function
# =====

# first, estimate a temporally pooled ERGM

# the Knecht network has the same dimensions at all time steps; "10"
# values indicate composition change; NA values are missings
```

```

# step 1: make sure the network matrices have node labels
for (i in 1:length(friendship)) {
  rownames(friendship[[i]]) <- 1:nrow(friendship[[i]])
  colnames(friendship[[i]]) <- 1:ncol(friendship[[i]])
}
rownames(primary) <- rownames(friendship[[1]])
colnames(primary) <- colnames(friendship[[1]])

# step 2: preprocess dep. NW, remove struct. zeros, impute NA with 0:
# get rid of missing data in the friendship networks (the "dependent
# variable") and adjust the dimensions temporally

dep <- preprocess(friendship, primary, demographics$sex, lag = FALSE,
  covariate = FALSE, na = NA, na.method = "fillmode",
  structzero = 10, structzero.method = "remove")
  # this has to be repeated for the covariates if they have NAs

length(dep)          # make sure there are still four time steps
sapply(friendship, dim) # network dimensions: 26-26-26-26
sapply(dep, dim)      # after the adjustment: 26-26-25-25
rownames(dep[[3]])    # node 21 (an NA value) was removed

# step 3: preprocess covariates; adjust them to the "dep" dimensions

primary.cov <- preprocess(primary, dep, demographics$sex,
  lag = FALSE, covariate = TRUE)
sex.cov <- preprocess(demographics$sex, primary.cov, dep,
  lag = FALSE, covariate = TRUE)

sapply(primary.cov, dim) # 26-26-25-25
sapply(sex.cov, length) # 26-26-25-25

# step 4: add nodal covariates to the networks
for (i in 1:length(dep)) {
  dep[[i]] <- network(dep[[i]])
  odegsqrt <- sqrt(degree(dep[[i]], cmode = "outdegree"))
  idegsqrt <- sqrt(degree(dep[[i]], cmode = "indegree"))
  dep[[i]] <- set.vertex.attribute(dep[[i]], "odegsqrt", odegsqrt)
  dep[[i]] <- set.vertex.attribute(dep[[i]], "idegsqrt", idegsqrt)
  dep[[i]] <- set.vertex.attribute(dep[[i]], "sex",
    sex.cov[[i]])
}

# step 5: estimate the TERGM without any lagged terms
model1 <- btergm(dep ~ edges + mutual + ttriple + transitivities +
  ctriple + nodeicov("idegsqrt") + nodeicov("odegsqrt") +
  nodecov("odegsqrt") + nodeofactor("sex") + nodeifactor("sex") +
  nodematch("sex") + edgecov(primary.cov), R = 100)

summary(model1, level = 0.95) # look at the results of the estimation

# step 5: plot goodness of fit (see ?gof.btergm and ?plot.btergm)
gof1 <- gof(model1, nsim = 25)

```

```

plot(gof1) # display boxplot diagram

# =====
# Add cross-temporal dynamics
# =====

# we now include cross-temporal dynamics

# step 1: preprocess data; this time with a lag
# dep contains t = 2 to t = 4
# lag contains t = 1 to t = 3
# mem indicates edge stability between 1-2, 2-3, and 3-4
dep <- preprocess(friendship, primary, demographics$sex, lag = TRUE,
  covariate = FALSE, na = NA, na.method = "fillmode",
  structzero = 10, structzero.method = "remove")
lag <- preprocess(friendship, primary, demographics$sex, lag = TRUE,
  covariate = TRUE, na = NA, na.method = "fillmode",
  structzero = 10, structzero.method = "remove")
mem <- preprocess(friendship, primary, demographics$sex, lag = TRUE,
  covariate = TRUE, memory = "stability", na = NA,
  na.method = "fillmode", structzero = 10,
  structzero.method = "remove")
primary.cov <- preprocess(primary, dep, demographics$sex,
  lag = FALSE, covariate = TRUE)
sex.cov <- preprocess(demographics$sex, primary.cov, dep,
  lag = FALSE, covariate = TRUE)

length(dep)      # now there are only three time steps
sapply(dep, dim) # after the adjustment: 26-25-25
sapply(lag, dim) # 26-25-25
sapply(mem, dim) # 26-25-25 (stability between t and t - 1)

# delayed reciprocity: are ties from t - 1 reciprocated at t?
delrecip <- lapply(friendship, t) # transpose friendship matrices
delrecip <- preprocess(delrecip, primary, friendship, lag = TRUE,
  covariate = TRUE, na = NA, na.method = "fillmode",
  structzero = 10, structzero.method = "remove")
sapply(delrecip, dim)

# step 3: add nodal covariates to the networks
for (i in 1:length(dep)) {
  dep[[i]] <- network(dep[[i]])
  odegsqrt <- sqrt(degree(dep[[i]], cmode = "outdegree"))
  idegsqrt <- sqrt(degree(dep[[i]], cmode = "indegree"))
  dep[[i]] <- set.vertex.attribute(dep[[i]], "odegsqrt", odegsqrt)
  dep[[i]] <- set.vertex.attribute(dep[[i]], "idegsqrt", idegsqrt)
  dep[[i]] <- set.vertex.attribute(dep[[i]], "sex", sex.cov[[i]])
}

# step 4: estimate the TERGM with cross-temporal model terms
model2 <- btergm(dep ~ edges + mutual + ttriple + transitivities +
  ctriple + nodeicov("idegsqrt") + nodeicov("odegsqrt") +

```

```

nodeocov("odegsqrt") + nodeofactor("sex") + nodeifactor("sex") +
nodematch("sex") + edgecov(primary.cov) + edgecov(delrecip) +
edgecov(mem), R = 100)

# look at the results (first using tables, then visually)
summary(model2)
screenreg(list(model1, model2)) # compare the two models directly
plotreg(model2, custom.model.names = "Model 2", custom.coef.names =
  c("Edges", "Reciprocity", "Transitive triples",
    "Transitive ties", "Cyclic triples", "Indegree popularity",
    "Outdegree popularity", "Outdegree activity", "Ego = male",
    "Alter = male", "Both nodes = male", "Same primary school",
    "Delayed reciprocity", "Memory term (edge stability)",
    omit.coef = "Edges", file="coefs.pdf")

# step 5: plot goodness of fit (see ?gof.btergm and ?plot.btergm)
gof2 <- gof(model2, nsim = 25)
plot(gof2) # display boxplot diagram; model fit is much better now!

# =====
# assess out-of-sample predictive performance of the model
# =====

# we also want to try to predict the network at t = 4 based on a model
# estimated for t = 1 through t = 3
model3 <- btergm(dep[1:2] ~ edges + mutual + ttriple + transitivities +
  ctriple + nodeicov("idegsqrt") + nodeicov("odegsqrt") +
  nodeocov("odegsqrt") + nodeofactor("sex") + nodeifactor("sex") +
  nodematch("sex") + edgecov(primary.cov[1:2]) + edgecov(delrecip[1:2]) +
  edgecov(mem[1:2]), R = 100)

screenreg(list(model1, model2, model3)) # similar results as before

# simulate 100 networks from t = 3 and compare to t = 4
gof3 <- gof(model3, nsim = 100, target = dep[[3]], formula = dep[[3]] ~
  edges + mutual + ttriple + transitivities + ctriple +
  nodeicov("idegsqrt") + nodeicov("odegsqrt") +
  nodeocov("odegsqrt") + nodeofactor("sex") + nodeifactor("sex") +
  nodematch("sex") + edgecov(primary.cov[[3]]) +
  edgecov(delrecip[[3]]) + edgecov(mem[[3]]))

# display goodness of fit
plot(gof3, roc = FALSE, pr = FALSE) # predictive fit (boxplots)
gof3 # display goodness of fit tables
pdf("rocpr.pdf")
plot(gof3, boxplot = FALSE, pr = FALSE, roc = TRUE,
  roc.random = TRUE, pr.random = FALSE, ylab = "TPR/PPV",
  xlab = "FPR/TPR", roc.main = "ROC and PR curves") # ROC curve
plot(gof3, boxplot = FALSE, roc = FALSE, pr = TRUE,
  pr.random = TRUE, rocpr.add = TRUE) # add PR curve
dev.off()

```

```

# =====
# assess predictive goodness of fit of a SAOM (estimated using RSiena)
# =====

# determine composition change
comp <- rep(list(c(1, 4)), 26) # actors are present from t=1 to t=4
comp[[21]] <- c(1, 2) # actor 21 drops out after the second time step
changes <- sienaCompositionChange(comp)

# prepare networks and covariate model terms
siena.nets <- sienaNet(array(c(friendship[[1]], friendship[[2]],
  friendship[[3]], friendship[[4]]), dim = c(26, 26, 4)))
primaryCov <- coDyadCovar(primary)
sexCov <- coCovar(demographics$sex)
ageCov <- coCovar(demographics$age)
ethnicityCov <- coCovar(demographics$ethnicity)
religionCov <- coCovar(demographics$religion)

mymodel <- sienaModelCreate(useStdInits = FALSE, projname = "myproject")

mydata <- sienaDataCreate(siena.nets, primaryCov, sexCov, ageCov,
  ethnicityCov, religionCov, changes)

# add effects
myeff <- getEffects(mydata)
myeff <- includeEffects(myeff, transTrip)
myeff <- includeEffects(myeff, transTies)
myeff <- includeEffects(myeff, cycle3)
myeff <- includeEffects(myeff, outPopSqrt)
myeff <- includeEffects(myeff, egoX, interaction1 = "sexCov")
myeff <- includeEffects(myeff, altX, interaction1 = "sexCov")
myeff <- includeEffects(myeff, sameX, interaction1 = "sexCov")
myeff <- includeEffects(myeff, X, interaction1 = "primaryCov")
myeff <- includeEffects(myeff, inPopSqrt)
myeff <- includeEffects(myeff, outActSqrt)

# estimate the SIENA model
ans <- siena07(mymodel, data = mydata, effects = myeff, batch = TRUE,
  verbose = FALSE)
ans # display results

# finally, assess predictive performance of the model using the xergm
# package; this should take up to 10 minutes on a recent PC
siena.gof <- gof(
  mymodel, # hand over the SIENA model
  siena.data = mydata, # ... and the SIENA data
  siena.effects = myeff, # ... and the SIENA effects
  nsim = 3, # number of estimation replications
  # (should be 100, but is very slow!)
  target.na = NA, # how are NA values denoted?
  target.na.method = "remove", # remove NAs in comparison network
  target.structzero = 10, # how are structural zeros denoted?
  parallel = "no", # adjust this for parallel processing

```

```

    ncpus = 1          # number of CPU cores
  )

siena.gof # display goodness of fit results

# plot the goodness of fit (see ?gof.btergmgoF for details)
plot(siena.gof, roc = FALSE, pr = FALSE) # display boxplot diagram
plot(siena.gof, boxplot = FALSE, pr = FALSE) # display ROC curve
plot(siena.gof, boxplot = FALSE, roc = FALSE) # display PR curve

# compare ROC and PR curves for TERGM and SAOM
plot(siena.gof, boxplot = FALSE, pr = FALSE, roc.col = "red1",
     ylab = "", xlab = "", roc.main = "SAOM versus TERGM prediction")
plot(siena.gof, boxplot = FALSE, roc = FALSE, pr.col = "steelblue1",
     rocpr.add = TRUE)
plot(gof3, boxplot = FALSE, pr = FALSE, roc.col = "red4",
     rocpr.add = TRUE)
plot(gof3, boxplot = FALSE, roc = FALSE, pr.col = "steelblue4",
     rocpr.add = TRUE)
color <- c("red1", "red4", "steelblue1", "steelblue4")
label <- c("SAOM ROC", "TERGM ROC", "SAOM PR", "TERGM PR")
legend("bottom", legend = label, col = color, lty = 1, lwd = 3)

# compare area under the curve
ROC <- c(SAOM = siena.gof$auc.roc, TERGM = gof3$auc.roc)
PR <- c(SAOM = siena.gof$auc.pr, TERGM = gof3$auc.pr)
barplot(cbind(ROC, PR), beside = TRUE, col = color)
legend("topright", legend = label, pch = 15, col = color)

## End(Not run)

```

plot.btergmgoF *Plot or print btergmgoF objects*

Description

Plot or print formatted goodness-of-fit statistics and degeneracy checks from btergmgoF objects.

Usage

```

## S3 method for class 'btergmgoF'
plot(x, boxplot = TRUE, boxplot.mfrow = TRUE,
     boxplot.dsp = TRUE, boxplot.esp = TRUE, boxplot.geodist = TRUE,
     boxplot.degree = TRUE, boxplot.idegree = TRUE,
     boxplot.odegree = TRUE, boxplot.kstar = TRUE, boxplot.istar = TRUE,
     boxplot.ostar = TRUE, boxplot.dsp.max = NULL,
     boxplot.esp.max = NULL, boxplot.geodist.max = NULL,
     boxplot.degree.max = NULL, boxplot.idegree.max = NULL,

```



```

boxplot.odegree.max = NULL, boxplot.kstar.max = NULL,
boxplot.istar.max = NULL, boxplot.ostar.max = NULL,
boxplot.transform = function(x) x, boxplot.border = "darkgray",
boxplot.mean.col = "black", boxplot.median.col = "black",
boxplot.lwd = 0.8, boxplot.outline = FALSE, boxplot.ylab = "Frequency",
boxplot.main = NULL, boxplot.ylim = NULL, roc = TRUE, pr = TRUE,
rocpr.add = FALSE, rocpr.avg = c("none", "horizontal", "vertical",
"threshold"), rocpr.spread = c("boxplot", "stderror", "stddev"),
rocpr.lwd = 3, roc.main = NULL, roc.random = FALSE, roc.col = "#bd0017",
roc.random.col = "#bd001744", pr.main = NULL, pr.random = FALSE,
pr.col = "#5886be", pr.random.col = "#5886be44", pr.poly = 0, ...)

## S3 method for class 'btergmof'
print(x, classicgof = TRUE,
      rocprgof = TRUE, degeneracy = TRUE, ...)

## S3 method for class 'btergmof'
summary(object, classicgof = TRUE,
        rocprgof = TRUE, degeneracy = TRUE, ...)

```

Arguments

x	A btergmof object created by calling the gof.btergm method.
boxplot	Create a plot where simulated statistics are summarized as boxplots and observed networks as lines (= classic statnet-style GOF plots)?
boxplot.mfrow	Should the classic GOF plots come out separately (boxplot.mfrow = FALSE), or should all statistics be aligned in a single diagram (boxplot.mfrow = TRUE)? Returning the plots separately can be helpful if the output is redirected to a multipage PDF or TIFF file.
boxplot.dsp	Plot a GOF diagram for dyad-wise shared partners?
boxplot.esp	Plot a GOF diagram for edge-wise shared partners?
boxplot.geodist	Plot a GOF diagram for geodesic distances?
boxplot.degree	Plot a GOF diagram for the degree distribution?
boxplot.idegree	Plot a GOF diagram for the indegree distribution?
boxplot.odegree	Plot a GOF diagram for the outdegree distribution?
boxplot.kstar	Plot a GOF diagram for the kstar distribution?
boxplot.istar	Plot a GOF diagram for the instar distribution?
boxplot.ostar	Plot a GOF diagram for the outstar distribution?
boxplot.dsp.max	Upper bound of dyad-wise shared partners to report.
boxplot.esp.max	Upper bound of edge-wise shared partners to report.

boxplot.geodist.max	Upper bound of geodesic distances to report (excluding Inf values).
boxplot.degree.max	Upper bound of the degree statistic to report.
boxplot.idegree.max	Upper bound of the indegree statistic to report.
boxplot.odegree.max	Upper bound of the outdegree statistic to report.
boxplot.kstar.max	Upper bound of the kstar statistic to report.
boxplot.istar.max	Upper bound of the istar statistic to report.
boxplot.ostar.max	Upper bound of the ostar statistic to report.
boxplot.transform	A function which transforms the y values used for the boxplots. For example, if some of the values become very large and make the output illegible, <code>boxplot.transform = function(x) x^0.1</code> or a similar transformation of the values can be used. Note that logarithmic transformations often produce infinite values because $\log(0) = -\text{Inf}$, so one should rather use something like <code>boxplot.transform = function(x) log1p</code> to avoid infinite values.
boxplot.border	Color of the borders of the boxplots.
boxplot.mean.col	Color of the mean of the observed network statistic.
boxplot.median.col	Color of the median of the observed network statistic.
boxplot.lwd	Scaling factor for the line width of the boxplots and of the line for the observed network statistics.
boxplot.outline	Print outliers in the boxplots?
boxplot.ylab	Label of the y-axis of a GOF plot.
boxplot.main	Main title of a GOF plot. Note: The same main title is used for all resulting diagrams. To obtain diagrams with varying main titles, plot each statistic separately using a new call of the <code>plot.btergm</code> method.
boxplot.ylim	Vertical limit of the boxplots. Only the maximum value must be provided because the minimum is always 0. Providing a vertical limit may be sensible if multiple models have to be compared side by side.
roc	Should the ROC curve(s) be plotted?
pr	Should the precision-recall curve(s) be plotted?
rocpr.add	Add the ROC and/or PR curve to an existing plot?
rocpr.avg	Averaging pattern for the ROC and PR curve(s) if multiple target time steps were used. Allowed values are "none" (plot all curves separately), "horizontal" (horizontal averaging), "vertical" (vertical averaging), and "threshold" (threshold (= cutoff) averaging). Note that while threshold averaging is always feasible, vertical and horizontal averaging are not well-defined if the graph cannot be

represented as a function $x \rightarrow y$ and $y \rightarrow x$, respectively. More information can be obtained from the help pages of the **ROCR** package, the functions of which are employed here.

rocpr.spread	When multiple target time steps are used and curve averaging is enabled, the variation around the average curve can be visualized as standard error bars ("stderror"), standard deviation bars ("stddev"), or by using box plots ("boxplot"). Note that the function plotCI, which is used internally by the ROCR package to draw error bars, might raise a warning if the spread of the curves at certain positions is 0. More details can be found in the documentation of the ROCR package, the functions of which are employed here.
rocpr.lwd	Line width of the ROC and PR curve(s).
roc.main	Main title of the ROC plot.
roc.random	Draw an additional ROC curve for random graphs with the same tie probability (as a reference value)?
roc.col	Color of the ROC curve(s). By default, a dark red color is used.
roc.random.col	Color of the random graph ROC curve.
pr.main	Main title of the ROC plot.
pr.random	Draw an additional PR curve for random graphs with the same tie probability (as a reference value)?
pr.col	Color of the precision-recall curve(s). By default, a blue color is used.
pr.random.col	Color of the random graph PR curve.
pr.poly	If a value of 0 is set, nothing special happens. If a value of 1 is set, a straight line is fitted through the PR curve and displayed. Values between 2 and 9 fit higher-order polynomial curves through the PR curve and display the resulting curve. This argument allows to check whether the imputation of the first precision value in the PR curve yielded a reasonable result (in case the value had to be imputed).
classicgof	Print tables with t-tests comparing observed and simulated networks, as in statnet?
rocprgof	Print tables with the area under the curve (AUC) for the PR and ROC curves at each time step of the observed network(s)?
degeneracy	Print tables with t-tests comparing the global statistics of observed and simulated networks for each observed time step? This is essentially a degeneracy check.
object	A btergmof object created by calling the gof.btergm method.
...	Further arguments. These arguments are passed on to the plot command. This facilitates customization of the plots.

Details

These methods plot or print nicely formatted goodness-of-fit statistics from btergmof objects to the R console. The typical workflow is to estimate a TERGM using the [btergm](#) function and save it as an object, then use the [gof](#) function to produce a btergmof object, and finally use the functions and methods described here to show the output of this resulting object to assess the goodness of fit.

The `plot` method plots (1) boxplots of network statistics and draws the observed statistics as lines (classic `statnet`-like GOF boxplots) and (2) receiver operating characteristics (ROC) and precision recall (PR) curves.

The `print` and `summary` methods show (1) observed versus simulated network statistics (classic `statnet`-like GOF tables), (2) the area under the ROC curve (AUC) for each observed network, and (3) degeneracy checks by comparing global statistics of simulated versus observed networks.

See Also

[xergm-package btergm simulate.btergm gof](#)

```
preprocess
```

Preprocess lists of network matrices for use with btergm

Description

Remove NAs and structural zeroes, create lagged covariates, and create memory terms.

Usage

```
preprocess(object, ..., lag = FALSE, covariate = FALSE,
           memory = c("no", "autoregression", "stability",
                     "innovation", "loss"), na = NA, na.method = "fillmode",
           structzero = -9, structzero.method = "remove",
           verbose = FALSE)
```

Arguments

- | | |
|---------------------|--|
| <code>object</code> | The object of interest that should be manipulated. The object can be a matrix, vector, <code>data.frame</code> , or a list of several objects of this kind. Usually, this is a list of network matrices (the dependent variable for the <code>btergm</code> function) or a list of covariates that needs to be adjusted given the missing observations at previous or future time steps. The name of the object is not allowed to match any of the argument names of this function (e.g., handing over an object with the name <code>memory</code> will cause an error message). |
| <code>...</code> | One or multiple other objects, separated by commas. These are used for comparison. If data are missing in these objects but not in the object of interest (given as the <code>object</code> argument), the missing data are also removed from the object of interest. The names of these objects are not allowed to match any of the argument names of this function (e.g., handing over an object with the name <code>memory</code> will cause an error message). |
| <code>lag</code> | Take into account missing or unobserved data at the previous or next time step? Use this argument in conjunction with the <code>covariate</code> argument. If <code>lag = TRUE</code> and <code>covariate = TRUE</code> , the first item in the list is adjusted to the dimensions of the second item, the second item is adjusted to the dimensions of the third item, and so forth, and the last item is omitted because it cannot be used as a covariate |

anyway. If `lag = TRUE` and `covariate = FALSE`, the first item is omitted, the second item is adjusted to the dimensions of the first item, the third item is adjusted to the dimensions of the second item, and so forth. If `lag = FALSE` (the `covariate` argument does not matter in this case), the first item is adjusted to the dimensions of the first item, the second item is adjusted to the dimensions of the second item etc. In all cases, missing data are first removed cross-sectionally across objects.

<code>covariate</code>	Is the object of interest a covariate or a dependent variable/network? This governs whether forward- or backward-looking missing data adjustment is used (when <code>lag = TRUE</code> is active). See the entry for <code>lag</code> for more details.
<code>memory</code>	Create a memory term which can be used as an edge covariate. Memory terms describe the temporal stability of the network. Several types of memory terms can be created. The default value <code>memory = "no"</code> will not create a memory term. <code>memory = "stability"</code> creates a term that captures dyadic stability between one time step and the next. <code>memory = "autoregression"</code> creates a lagged dependent network and is equivalent to <code>memory = "no"</code> . The result is a positive autoregression term, which is the lagged dependent network. This captures the stability of edges (but not of non-edges). <code>memory = "innovation"</code> measures the extent to which new edges are introduced over time ("edge innovation"). <code>memory = "loss"</code> captures the extent to which previously existing edges are no longer present at each current time step. The memory argument only works when <code>lag = TRUE</code> and <code>covariate = TRUE</code> are also set.
<code>na</code>	A vector of values that are identified as missing data. These values can be removed or replaced (see the handleMissings function).
<code>na.method</code>	What should be done with missing data? Valid values are <code>remove</code> and <code>fillmode</code> . See the handleMissings function for details.
<code>structzero</code>	A vector of values that are identified as structural zeroes. These values can be removed or replaced (see the handleMissings function).
<code>structzero.method</code>	What should be done with structural zeroes? Valid values are <code>remove</code> and <code>fillmode</code> . See the handleMissings function for details.
<code>verbose</code>	Report the amount of missing data that were removed or replaced in each object?

Details

For use with the `btergm` function, lists of network matrices or covariates can be preprocessed using the `preprocess` function. The first object that is provided as an argument is adjusted to all remaining objects that are handed over via the `...` argument. The `preprocess` function can deal with constant or time-varying matrix, vector and `data.frame` objects. The user can specify whether the object of interest is a dependent network or a covariate and whether NA and structural zero removal should take into account missing data at previous or following time steps (`lag = TRUE`).

The preprocessing procedure consists of four steps: (1) remove or impute missing data and/or structural zeroes in each object at each time step; (2) cross-sectional adjustment of matrix dimensions (e.g., if the object of interest has more observations than another object at the current time step, the observations are dropped from the object of interest); (3) backward-looking adjustment of matrix dimensions (if the object of interest is a dependent variable/network and the `lag = TRUE` argument

is used); and (4) forward-looking adjustment of matrix dimensions (if the object of interest is a covariate and the `lag = TRUE` argument is used). The last two steps can be helpful for building lagged or delayed covariates which have to take into account missing data at other time steps as well.

For example, if 26 nodes are present during the first time step, 25 during the second, 23 during the third, and 25 during the fourth time step, the arguments `covariate = TRUE` and `lag = TRUE` return a list of three objects with 25, 23 and 25 objects, respectively, because the covariate is assumed to lag one step behind the other objects provided after the first object.

If the memory argument is specified (see below for details on the allowed values), the function will not return a lagged network, but will attempt to create a memory term. A memory term is a dyadic covariate which captures the temporal process.

See Also

[xergm-package handleMissings adjust timecov](#)

Examples

```
## Not run:
# This example illustrates the usefulness of the preprocess function.

# first network: nodes a to j present
mat1 <- rbinom(100, 1, 0.1)
mat1 <- matrix(mat1, nrow = 10) # has 10 nodes
rownames(mat1) <- letters[1:10]
colnames(mat1) <- letters[1:10]

# second network: nodes c to n present
mat2 <- rbinom(144, 1, 0.1)
mat2 <- matrix(mat2, nrow = 12) # has 12 nodes
rownames(mat2) <- letters[3:14]
colnames(mat2) <- letters[3:14]

# third network: nodes a and d to k present
mat3 <- rbinom(81, 1, 0.1)
mat3 <- matrix(mat3, nrow = 9) # has 9 nodes
rownames(mat3) <- letters[c(1, 4:11)]
colnames(mat3) <- letters[c(1, 4:11)]

# fourth network: same as second matrix
mat4 <- mat2

networks <- list(mat1, mat2, mat3, mat4)

# btergm without cross-temporal dependencies:
model.1 <- btergm(networks ~ edges + mutual)
summary(model.1)

# When cross-temporal dependencies are specified, the dimensions
# of the matrices do not match. This would cause a problem for btergm:

\dontrun{
```

```

btergm(networks[2:4] ~ edges + mutual + edgecov(networks[1:3])) # ERROR!
}

# This is because the first network in the dependent network and the
# first network in the lagged covariate are expected to have the same
# dimensions (and also at the second and third time step, of course).

# Therefore, missing nodes in the covariate (here: {k, l, m, n} at t=1,
# {a} at t=2, and {c, l, m, n} at t=3) must be removed from the
# dependent network at t=2, t=3 and t=4 as well:

dep <- preprocess(networks, lag = TRUE, covariate = FALSE)

# This reduces the size of dep from 12 to 8 at t=2, from 9 to 8 at
# t=3, and from 12 to 8 at t=4, and it removes the first network from
# the list. Moreover, some nodes are present in the lagged covariate
# but not in the dependent network (that is, at the next time step).
# Therefore, node sets {a, b}, {c, l, m, n}, and {a} must be removed
# from the lagged covariate at t=1, t=2, and t=3, respectively, to make
# the dimensions compatible:

lag <- preprocess(networks, lag = TRUE, covariate = TRUE)

# To compare the dimensions of the original versus preprocessed
# dependent networks and covariates, try the following code:

cbind(
  "original_dep" = lapply(networks[2:4], nrow),
  "original_lag" = lapply(networks[1:3], nrow),
  "new_dep" = lapply(dep, nrow),
  "new_lag" = lapply(lag, nrow)
)

# The dependent networks were reduced from 12, 9 and 12 to 8, 8 and
# 8 nodes, and the lagged networks were reduced from 10, 12 and 9 to
# 8, 8 and 8 nodes, respectively. The lagged node sets are now
# compatible. To see this:

cbind(rownames(dep[[1]]), rownames(lag[[1]]))
cbind(rownames(dep[[2]]), rownames(lag[[2]]))
cbind(rownames(dep[[3]]), rownames(lag[[3]]))

# Note, however, that the composition still changes within each list
# across some of the time steps:

cbind(rownames(dep[[1]]), rownames(dep[[2]]), rownames(dep[[3]]))
cbind(rownames(lag[[1]]), rownames(lag[[2]]), rownames(lag[[3]]))

# We can now use the btergm function on the preprocessed lists:

model.2 <- btergm(dep ~ edges + mutual + edgecov(lag))
summary(model.2)

```

```

# The model can now be estimated because the current and lagged networks
# have the same node sets at each time step. The disadvantage of this
# approach is that some observations are lost. The advantage, however,
# is that cross-temporal theories can be tested.

# However, since the node sets still differ across time steps, ROC and
# PR curves cannot be estimated. This is true because a simulation from
# nodes {c ... j} cannot be compared to a target network with nodes
# {d ... k}. Therefore, the following command would compare the wrong
# sets of nodes to estimate prediction performance:

\dontrun{
gof.2 <- gof(model.2, classicgof = FALSE, rocprgof = TRUE) # PROBLEM!
}

# To solve this problem, the most obvious approach is to estimate the
# model at earlier time steps and compute the out-of-sample predictive
# performance only for the last network:

model.3 <- btergm(dep[1:2] ~ edges + mutual + edgescov(lag[1:2]))
gof.3 <- gof(model.3, target = dep[[3]], formula = dep[[3]] ~ edges +
  mutual + edgescov(lag[[3]]), classicgof = FALSE, rocprgof = TRUE)

# This models time steps 2 and 3 as a function of the lagged network
# at time steps 1 and 2, uses the resulting coefficients to predict
# the network at time step 4, and compares network 4 to simulations
# based on the coefficients from the previous time steps and the
# lagged network at the third time step. As the matrices within the
# third list item have identical node sets, predictive performance
# could be computed. The resulting ROC and PR curves can be plotted
# as follows:

plot(gof.3, boxplot = FALSE, pr = FALSE, roc.random = TRUE,
  ylab = "TPR/PPV", xlab = "FPR/TPR", roc.main = "ROC and PR")
plot(gof.3, boxplot = FALSE, roc = FALSE, pr.random = TRUE,
  rocpr.add = TRUE)
legend("right", legend = c("ROC", "ROC random graph", "PR",
  "PR random graph"), col = c("#bd0017", "#bd001744", "#5886be",
  "#5886be44"), lty = 1, lwd = 3)

# For another example with real-world data, see vignette("knecht")

## End(Not run)

```

simulate.btergm

Simulate new networks from btergm objects

Description

Simulate new networks from btergm objects.

Usage

```
## S3 method for class 'btergm'
simulate(object, nsim = 1, seed = NULL,
         index = NULL, formula = getformula(object),
         coef = object@coef, verbose = TRUE, ...)
```

Arguments

object	A btergm object, resulting from a call of the btergm function.
nsim	The number of networks to be simulated. Note that for values greater than one, a <code>network.list</code> object is returned, which can be indexed just like a <code>list</code> object, for example <code>mynetworks[[1]]</code> for the first simulated network in the object <code>mynetworks</code> .
seed	Random number integer seed. See set.seed .
formula	A model formula from which the new network(s) should be simulated. By default, the formula is taken from the btergm object.
index	Index of the network from which the new network(s) should be simulated. The index refers to the list of response networks on the left-hand side of the model formula. Note that more recent networks are located at the end of the list. By default, the first (= oldest) network is used.
coef	A vector of parameter estimates. By default, the coefficients are extracted from the given btergm object.
verbose	Print additional details while running the simulations?
...	Arbitrary further arguments are handed over to the simulate.formula function. For details, refer to the help page of the simulate.formula function.

Details

The `simulate.btergm` function is a wrapper for the `simulate.formula` function in the **ergm** package (see `help("simulate.formula")`). It can be used to simulate new networks from a btergm object. The `index` argument specifies from which of the original networks the new network(s) should be simulated. For example, if `object` is an estimation based on cosponsorship networks from the 99th to the 107th Congress (as in Desmarais and Cranmer 2012), and the cosponsorship network in the 108th Congress should be predicted using the `simulate.btergm` function, then the argument `index = 9` should be passed to the function because the network should be based on the 9th network in the list (that is, the latest network, which is the cosponsorship network for the 107th Congress). Note that all relevant objects (the networks and the covariates) must be present in the workspace (as was the case during the estimation of the model).

References

Desmarais, Bruce A. and Skyler J. Cranmer (2012): Statistical Mechanics of Networks: Estimation and Uncertainty. *Physica A* 391: 1865–1876.

See Also

[xergm-package btergm gof](#)

timecov	<i>Create interaction terms between covariates and (transformations of) time for TERGMs</i>
---------	---

Description

Create interaction terms between covariates and (transformations of) time for TERGMs.

Usage

```
timecov(covariate, minimum = 1, maximum = length(covariate),
        transform = function(t) 1 + (0 * t) + (0 * t^2),
        onlytime = FALSE)
```

Arguments

covariate	A list of matrices or network objects (a time-varying dyadic covariate). Note that nodal covariates can be manually converted into dyadic covariates to use them with this function.
minimum	The first matrix or network in the covariate list which should be non-zero. All matrices or networks before this time step are filled with zeros.
maximum	The last matrix or network in the covariate list which should be non-zero. All matrices or networks after this time step are filled with zeros.
transform	A function that transforms the time axis. By default, all time steps are weighted equally. For example, if there are five consecutive matrices, all of them are equally important, i.e., time is represented by weights of 1-1-1-1-1. A linearly increasing importance of time can be achieved by using a multiplication with time, for example $\text{function}(t) = 2 * t$ leads to a multiplication of all values in the covariates by the values 2-4-6-8-10, respectively by time step. Quadratic or higher-order polynomials are possible using more complex transformations of time.
onlytime	If <code>onlytime = TRUE</code> is set, the function returns a list of matrices where all entries within a matrix are identical and reflect the transformation of time. This is useful when a function of time per se should be included in the model as a covariate, e.g., when the tie formation probability increases or decreases over time. If <code>onlytime = FALSE</code> is set, the aforementioned list of matrices is interacted with the covariate list, i.e., each value in the matrix at time step t is multiplied by the transformation of time at that time step.

Details

The `timecov` function takes a list of matrices or networks (a varying dyadic covariate) and creates an interaction term with time. For example, if the rules of network formation are expected to change after some time steps (say, after the third out of six time steps), one can model the effect of the covariate for the first three time points and for the remaining three time points using separate model terms. To achieve this, the covariate matrix at each time step is multiplied by zeros or ones,

depending on whether the time step should be incorporated in that model term. In this situation, one would create two dyadic covariates, one where the first three time periods are present and the remaining three are set to zero, and the other one where the first three time periods are set to zero and the remaining ones are present.

Another usage scenario is that time per se may have a polynomial effect (of any shape) on the probability of forming a tie. In this case, the `timecov` function can be used to create a covariate list of matrices where the entries of the matrix correspond to polynomial functions of time. For example, if tie formation becomes increasingly likely (as expressed by a linear relationship), a list of matrices with linearly increasing entries over time can be created. Such time effects can also be interacted with other covariates. For example, the further time progresses, the more (or less) the property captured by the covariate becomes important for tie formation.

See Also

[xergm-package btergm preprocess](#)

tnam	<i>Fit (temporal) network autocorrelation models</i>
------	--

Description

Fit (temporal) network autocorrelation models.

Usage

```
tnam(formula, family = gaussian, re.node = FALSE,
      re.time = FALSE, time.linear = FALSE, time.quadratic = FALSE,
      center.y = FALSE, na.action = na.omit, ...)
```

```
tnamdata(formula, center.y = FALSE)
```

Arguments

- | | |
|---------|--|
| formula | A formula where the left-hand side specifies either a vector containing the outcome variable (for a cross-sectional model) or a list of such vectors (for modeling the outcome at multiple time steps) or a data frame with one time step per column (also for longitudinal models of behavior). The right-hand side of the formula consists of tnam-specific model terms like <code>netlag</code> , <code>structsim</code> and other terms which are described on the help page of tnam-terms . |
| family | The link function for fitting the generalized linear model or the mixed effects model, for example <code>gaussian</code> or <code>binomial</code> . The options are the same as in the glm and glmer functions. For details on the family argument, see the family help page. |
| re.node | If multiple time steps are present: should a random effect for the nodes be added to the model? This results in the estimation of a mixed effects model. |

<code>re.time</code>	If multiple time steps are present: should a random effect for the time steps be added to the model? This results in the estimation of a mixed effects model.
<code>time.linear</code>	If multiple time steps are present: should a linear effect for time be added to the model? This can be estimated in the standard GLM framework.
<code>time.quadratic</code>	If multiple time steps are present: should a squared effect for time be added to the model? This can be estimated in the standard GLM framework.
<code>center.y</code>	Center the dependent variable by subtracting the mean from the actual value within each time step?
<code>na.action</code>	How should missing values be treated? By default, they are omitted. See the na.omit help page for details.
<code>...</code>	Further arguments that should be passed to the <code>glm</code> , <code>lmer</code> , or <code>glmer</code> function, which is used under the hood for estimating the model.

Details

The `tnam` function serves to estimate temporal or cross-sectional network autocorrelation models. Model terms such as spatial lags, temporal lags, spatio-temporal lags, centrality etc. can be specified in the formula argument. Details on the model terms can be found on the [tnam-terms](#) help page.

The `tnamdata` function accepts a formula (like in the `tnam` function) and returns a data frame with the response variable and the covariates for estimation with any estimation function. `tnam` first calls `tnamdata` internally and then hands over the resulting data structure to a `glm`, `lmer`, or `nlmer` call. If models such as tobit, multinomial or multilevel models should be estimated, one can leave out the estimation step and feed the results of `tnamdata` manually into any type of model.

See Also

[xergm-package tnam-terms preprocess knecht](#)

Examples

```
# The following example models delinquency among adolescents at
# multiple time steps as a function of (1) their nodal attributes
# like sex or religion, (2) their peers' delinquency levels, (3)
# their own and their peers' past delinquency behavior, and (4)
# their structural position in the network. See ?knecht for
# details on the dataset.

library("statnet")
library("xergm")
data("knecht")

# prepare the dependent variable y
delinquency <- as.data.frame(delinquency)
rownames(delinquency) <- letters

# replace structural zeros (denoted as 10) and add node labels
friendship[[3]][friendship[[3]] == 10] <- NA
friendship[[4]][friendship[[4]] == 10] <- NA
for (i in 1:length(friendship)) {
```

```

    rownames(friendship[[i]]) <- letters
    colnames(friendship[[i]]) <- letters
  }

  # prepare the sex variable (can be a list of vectors)
  sex <- as.numeric(demographics$sex)
  names(sex) <- letters
  sex <- list(sex, sex, sex, sex)

  # prepare the religion variable (can also be a data frame)
  rel <- as.numeric(demographics$religion)
  rel <- data.frame(rel, rel, rel, rel)
  rownames(rel) <- letters

  # Estimate the model. The first term is the sex of the respondent,
  # the second term is the religion of the respondent, the third
  # term is the previous delinquency behavior of the respondent,
  # the fourth term is the delinquency behavior of direct friends,
  # the fifth term is the delinquency behavior of indirect friends
  # at a path distance of 2, the sixth effect is the past delinquency
  # of direct friends, the seventh term indicates whether the
  # respondent has any contacts at all, and the last term captures
  # the effect of the betweenness centrality of the respondent on
  # his or her behavior. Apparently, previous behavior, being an
  # isolate, and religion seem to have an effect on delinquency in
  # this dataset. There is also a slight positive trend over time,
  # and direct friends exert a minor effect (not significant).
  # Note that a linear model may not be the best specification for
  # modeling the ordered categorical delinquency variable, but it
  # suffice here for illustration purposes.

  model1 <- tnam(
    delinquency ~
      covariate(sex, coefname = "sex") +
      covariate(rel, coefname = "religion") +
      covariate(delinquency, lag = 1, exponent = 1) +
      netlag(delinquency, friendship) +
      netlag(delinquency, friendship, pathdist = 2, decay = 1) +
      netlag(delinquency, friendship, lag = 1) +
      degreedummy(friendship, deg = 0, reverse = TRUE) +
      centrality(friendship, type = "betweenness"),
    re.node = TRUE, time.linear = TRUE
  )
  summary(model1)

  # for nice table output, use the texreg package
  library("texreg")
  screenreg(model1)

```

Description

The function `tnam` is used to fit (temporal) network autocorrelation models.

The function `tnamdata` can be used alternatively to create a data frame containing all the data ready for estimation. This may be useful when a non-standard model should be estimated, like a tobit model or a model with zero inflation, for example.

Both functions accept a formula containing several model terms. The model terms are themselves functions which can be called separately. For example, one model term is called `netlag`. This model term can be part of the formula handed over to the `tnam` function, or `netlag` can be called directly in order to create a single variable.

This help page describes the different model terms available in (temporal) network autocorrelation models. See the `tnam` help page for details on the model.

Usage

```
attribsim(y, attribute, match = FALSE, lag = 0,
          normalization = c("no", "row", "column"), center = FALSE,
          coefname = NULL)

centrality(networks, type = c("indegree", "outdegree", "freeman",
                              "betweenness", "flow", "closeness", "eigenvector",
                              "information", "load", "bonpow"), directed = TRUE, lag = 0,
          rescale = FALSE, center = FALSE, coefname = NULL, ...)

cliquelag(y, networks, k.min = 2, k.max = Inf, directed = TRUE,
          lag = 0, normalization = c("no", "row", "column"),
          center = FALSE, coefname = NULL)

clustering(networks, directed = TRUE, lag = 0, center = FALSE,
          coefname = NULL, ...)

covariate(y, lag = 0, exponent = 1, center = FALSE,
          coefname = NULL)

degreedummy(networks, deg = 0, type = c("indegree", "outdegree",
                                       "freeman"), reverse = FALSE, directed = TRUE, lag = 0,
          center = FALSE, coefname = NULL, ...)

interact(x, y, lag = 0, center = FALSE, coefname = NULL)

netlag(y, networks, lag = 0, pathdist = 1, decay = pathdist^-1,
        normalization = c("no", "row", "column", "complete"),
        reciprocal = FALSE, center = FALSE, coefname = NULL, ...)

structsim(y, networks, lag = 0, method = c("euclidean",
                                           "minkowski", "jaccard", "binary", "hamming"), center = FALSE,
          coefname = NULL, ...)
```

```
weightlag(y, networks, lag = 0, normalization = c("no", "row",
"column"), center = FALSE, coefname = NULL)
```

Arguments

attribute	A vector, list of vectors or data frame with the same dimensions as <i>y</i> . Based on this attribute, the similarity between nodes <i>i</i> and <i>j</i> will be calculated, and the resulting similarity matrix is used to weight the <i>y</i> variable.
center	Should the model term be centered? That is, should the mean of the variable be subtracted from the actual value at each time step?
coefname	An additional name that is used as part of the coefficient label for easier identification in the summary output of the model.
decay	For each value in <i>pathdist</i> , the decay argument specifies the relative importance. By default, a geometric decay is used, that is, the behavior of nodes at path distance 2 is counted only half as much as the behavior of adjacent nodes. Alternatively, if both are equally important, it is possible to write <i>pathdist</i> = <i>c</i> (1, 2) and <i>decay</i> = <i>c</i> (1, 1).
deg	The degree (e.g., <i>deg</i> = 2) or degree range (e.g., <i>deg</i> = 1:3).
directed	Is the input matrix or network a directed network?
exponent	The exponent of a covariate. For example, <i>exponent</i> = 2 creates a squared variable. This may be helpful for modeling non-linear effects or for modeling a quadratic behavior shape.
k.max	Maximal clique size.
k.min	Minimal clique size.
lag	The temporal lag. The default value 0 means there is no lag. A value of 1 would specify a single-period lag, that is, current behavior is modeled conditional on previous influence. A value of 2 would specify a two-period lag, that is, current behavior is modeled conditional on pre-previous influence, etc.
match	If <i>match</i> = FALSE, a similarity matrix is computed by subtracting node <i>j</i> 's attribute value from node <i>i</i> 's attribute value, standardizing the resulting distance between 0 and 1, and converting it into a similarity by subtracting it from 1. This similarity matrix is used as a weight matrix to compute a spatial lag. If <i>match</i> = TRUE is specified, the weight matrix contains values of 1 whenever node <i>i</i> and <i>j</i> have the same attribute value and 0 otherwise.
method	The distance function used for computing structural similarity. Possible values are "euclidean", "minkowski", "jaccard", "binary", and "hamming".
networks	The network(s) for computing the peer influence, also known as the weight matrix. This can be a matrix or a network object (for a single time step) or a list of matrices or network objects (for multiple time steps).
normalization	Possible values: "no" for switching off normalization, "row" for row normalization of the weight matrix, "column" for column normalization of the weight matrix, and "complete" for complete normalization. If "no" is selected, this corresponds to the total similarity or the sum of all influences of tied alters.

If "row" is selected, this corresponds to the average alter effect. If i is the row node and j is the column node, row normalization computes the peer influence of j on i as a fraction of i 's overall number of outgoing ties (i.e., i 's row sum or outdegree centrality). The theoretical intuition is that other nodes' influences on i are not cumulative; i rather perceives the average influence of his or her peers. Note that row normalization does not necessarily entail that the values are standardized between 0 and 1.

If "column" is selected, this corresponds to the peer influence of node j on node i as a fraction of the number of incoming ties j has (i.e., j 's column sum or indegree centrality). This captures the theoretical effect that j may distribute his or her influence among many nodes, in which case j 's influence on i is relatively weak. Thus the "exerted influence of j on i decreases with the number of actors j influences" (Leenders 2002). Note that column normalization does not necessarily entail that the values are standardized between 0 and 1.

If "complete" is selected, this captures the peer influence of node j on node i as a fraction of all nodes' cumulative outcome values (including non-tied dyads; except i 's own outcome value). In other words, complete normalization corresponds to the actual exposure of i to the influence of his or her tied alters j over the the exposure i could receive if i were tied to all other nodes in the network. If a decay ≤ 1 is used and if only direct friends are considered, this effectively standardizes the influence scores between 0 and 1.

pathdist	An integer or a vector of integers. For example, if pathdist = 1 is used, this computes the sum of the behavior of adjacent nodes. If pathdist = 2 is specified, this computes the effect of indirect paths of length 2 ("friends of friends"). If pathdist = 1:2 is set, both directly connected nodes' behavior and the behavior of nodes at a path distance of 2 from the focal node are counted. Arbitrary (sets of) path distances can be used. See also the decay argument.
reciprocal	If reciprocal = TRUE is specified, only the behavior of nodes to which a reciprocal relation exists is counted (that is, a link in both directions).
rescale	Should the centrality index be rescaled between 0 and 1?
reverse	Reverse the selection of degrees. For example, when deg = 0 and reverse = FALSE are specified, resulting values of 1 indicate that a node has no connections, whereas the combination deg = 0 and reverse = TRUE results in the value 1 representing nodes which have a degree of at least 1.
type	The type of centrality measure. Possible values are "indegree", "outdegree", "freeman", "betweenness", "flow", "closeness", "eigenvector", "information", "load", and "bonpow".
x	A variable that should be interacted with y . Either a vector or a list of vectors or another model term (this is the preferred way).
y	The outcome or behavior variable. Either a vector (for a single time step) or a list of vectors with named elements in each vector (for multiple time steps) or a data frame with row names where each column is one time step (for multiple time steps).
...	Additional arguments to be handed over to subroutines.

Model terms for tnam

- attribsim** *Spatial lag based on attribute similarity* The `attribsim` model term computes a similarity matrix based on the `attribute` argument and uses this similarity matrix to construct a spatial lag by multiplying the similarity matrix and the outcome vector `y`. The intuition behind this model term is that node `i`'s behavior may be influenced by node `j`'s behavior if nodes `i` and `j` are similar on another dimension. For example, if `i` and `j` both smoke while `k` does not smoke, `j`'s alcohol consumption may affect `i`'s alcohol consumption to a larger extent than node `k`'s alcohol consumption. In this example, the `y` outcome variable is alcohol consumption and the `attribute` argument is smoking. If `match = FALSE`, the absolute similarity between `i` and `j` is computed by subtracting `j`'s attribute value from `i`'s attribute value and taking the absolute value to construct the similarity matrix. If `match = TRUE`, the function computes a matrix containing values of 1 if `i` and `j` have the same attribute value and 0 otherwise. A scenario where the `attribsim` model term makes sense is degree assortativity: if `i` and `j` have the same degree centrality, they may be inclined to learn from each other's behavior, even in the absence of a direct connection between them.
- centrality** *Node centrality* The `centrality` model term computes a centrality index for the nodes in a network or matrix. This can capture important structural effects because being central often implies certain constraints or opportunities more peripheral nodes do not have. For example, central nodes in a network of employees might be able to perform better.
- cliquelag** *Spatial lag of k-clique co-members* The `cliquelag` model term computes a clique co-membership matrix and multiplies this matrix with the outcome variable. The intuition behind this is that in some settings individuals may be influenced to a particularly strong extent by peers in the same cliques. A clique is defined as a maximal connected subgraph of size `k`. For example, a deviant behavior of a person may be conditioned by the deviant behavior of the person's friends – but only if these friends are tied to each other as well so that a clique among these persons exists. A minimal and a maximal `k` may be defined, where `k` is the size of the cliques. In the clique co-membership matrix, all cliques with `k.min <= k <= k.max` are included.
- clustering** *Local clustering coefficient, or transitivity* The `clustering` model term computes the local clustering coefficient, which is also known as transitivity. This index has high values if the direct neighborhood of a node is densely interconnected. For example, if one's friends are friends with each other, this may have repercussions on ego's behavior.
- covariate** *Exogenous nodal covariate* The `covariate` model term adds an exogenous nodal covariate to the model. For example, when performance of employees is modeled, a covariate could be seniority of these employees. It is possible to add lagged covariates to model the effect of past nodal attributes on current behavior. Similarly, this model term can be used to add autoregressive terms, that is, the effect of previous behavior on current behavior.
- degreedummy** *Dummy variable for degree centrality values* The `degreedummy` model term controls for specific degree centralities or ranges of degree centrality. For example, do nodes with a degree of 0 (isolates) show different behavior than nodes who are connected? Or do nodes with a degree centrality larger than three exert different behavior?
- interact** *Interactions between other model terms* The `interact` model term adds an interaction effect between two other model terms by multiplying the result vectors of these two model terms. When using interaction terms, centering the result is recommended. Note that only the interaction term is created; the main effects must be introduced to the model using the other model terms.

netlag *Spatial network lag* The netlag model term captures the autocorrelation inherent in networks. For example, when political actors are members of a policy network, their success of achieving policy outcomes is not independent from each other. Most likely, being connected to policy winners increases the success rate. In many settings, indirect effects may be important as well: how does the behavior of my friends' friends affect my own behavior? In some contexts, spatio-temporal lags are useful: how does the past behavior of my friends affect my current behavior? The netlag model term is designed for binary networks because things like indirect effects, restriction to reciprocal dyads, decay of indirect relations etc. is possible. For weighted networks, the weightlag term is recommended. If no other arguments are specified and loops are absent and a binary matrix is used, both model terms produce the same results.

structsim *Structural similarity* The structsim model term computes the structural similarity with other nodes in the network and multiplies this similarity matrix with the outcome variable. The intuition is that behavior is sometimes affected by comparison with structurally similar nodes. For example, a worker may be impressed by the performance of other workers who are embedded in the same team or who report to the same bosses. As with the other model terms, temporal lags are possible.

weightlag *Weighted spatial lag* The weightlag model term captures spatial autocorrelation in weighted networks. For example, the GDP per capita of a country may be affected by the GDP of proximate other countries or by the GDP of trade partners. In these cases, indirect contacts etc. do not make any sense, therefore the distinction between the weightlag and the netlag model term. The weight matrix is multiplied by the outcome variable, possibly after row or column normalization.

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See Also

[xergm-package](#) [tnam](#) [tnamdata](#) [preprocess](#) [knecht](#)

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