

Package ‘tergm’

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Suggests lattice, snow

Description An integrated set of extensions to the `ergm` package to analyze and simulate network evolution based on exponential-family random graph models (ERGM). ``tergm`` is a part of the ``statnet`` suite of packages for network analysis.

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URL <http://www.statnet.org>

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tergm-package	<i>Fit, Simulate and Diagnose Dynamic Network Models derived from Exponential-Family Random Graph Models</i>
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Description

`tergm` is a collection of extensions to the `ergm` package to fit, diagnose, and simulate models for dynamic networks — networks that evolve over time — based on exponential-family random graph models (ERGMs). For a list of functions type `help(package='tergm')`

When publishing results obtained using this package, please cite the original authors as described in `citation(package="tergm")`.

All programs derived from this package must cite it.

Details

An exponential-family random graph model (ERGM) postulates an exponential family over the sample space of networks of interest, and `ergm` package implements a suite of tools for modeling single networks using ERGMs.

More recently, there has been a number of extensions of ERGMs to model evolution of networks, including the temporal ERGM (TERGM) of Hanneke et al. (2010) and the separable temporal ERGM (STERGM) of Krivitsky and Handcock (2013). The latter model allows familiar ERGM terms and statistics to be reused in a dynamic context, interpreted in terms of formation and dissolution of ties. Krivitsky (2012) suggested a method for fitting dynamic models when only a cross-sectional network is available, provided some temporal information for it is available as well.

This package aims to implement these and other ERGM-based models for network evolution. At this time, it implements, via the `stergm` function, the STERGMs, both a conditional MLE (CMLE) fitting to a series of networks and an Equilibrium Generalized Method of Moments Estimation

(EGMME) for fitting to a single network with temporal information. For further development, see the referenced papers.

For detailed information on how to download and install the software, go to the Statnet project website: statnet.org. A tutorial, support newsgroup, references and links to further resources are provided there.

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 coef.stergm

Extract Model Coefficients

Description

coef is a Method which extracts model coefficients from objects returned by the [stergm](#) function. coefficients is an *alias* for it.

Usage

```
## S3 method for class 'stergm'
coef(object, ...)
## S3 method for class 'stergm'
coefficients(object, ...)
```

Arguments

object	A stergm fit.
...	other arguments.

Value

Coefficients extracted from object in the form of a list with two elements: formation, a vector of formation coefficients and dissolution, a vector of dissolution coefficients.

See Also

[stergm](#) and [ergm](#)

 control.simulate.stergm

Auxiliary for Controlling Separable Temporal ERGM Simulation

Description

Auxiliary function as user interface for fine-tuning STERGM simulation.

Usage

```
control.simulate.stergm(MCMC.burnin.min=NULL, MCMC.burnin.max=NULL,
                        MCMC.burnin.pval=NULL, MCMC.burnin.add=NULL,
                        MCMC.burnin=NULL, MCMC.burnin.mul=NULL,
                        MCMC.prop.weights.form=NULL, MCMC.prop.args.form=NULL,
                        MCMC.prop.weights.diss=NULL, MCMC.prop.args.diss=NULL,
                        MCMC.init.maxedges=NULL,
```

```

MCMC.packagenames=NULL,
MCMC.init.maxchanges=NULL)

control.simulate.network(MCMC.burnin.min=1000, MCMC.burnin.max=100000,
MCMC.burnin.pval=0.5, MCMC.burnin.add=1,
MCMC.burnin=NULL, MCMC.burnin.mul=NULL,
MCMC.prop.weights.form="default",MCMC.prop.args.form=NULL,
MCMC.prop.weights.diss="default",MCMC.prop.args.diss=NULL,
MCMC.init.maxedges=20000,
MCMC.packagenames=c(),
MCMC.init.maxchanges=1000000)

```

Arguments

- `MCMC.burnin.min`, `MCMC.burnin.max`, `MCMC.burnin.pval`, `MCMC.burnin.add`
Number of Metropolis-Hastings steps per phase (formation and dissolution) per time step used in simulation. By default, this is determined adaptively by keeping track of increments in the Hamming distance between the transitioned-from network and the network being sampled (formation network or dissolution network). Once `MCMC.burnin.min` steps have elapsed, the increments are tested against 0, and when their average number becomes statistically indistinguishable from 0 (with the p-value being greater than `MCMC.burnin.pval`), or `MCMC.burnin.max` steps are proposed, whichever comes first, the simulation is stopped after an additional `MCMC.burnin.add` times the number of elapsed steps had been taken. (Stopping immediately would bias the sampling.)
To use a fixed number of steps, set both `MCMC.burnin.min` and `MCMC.burnin.max` to the desired number of steps.
- `MCMC.prop.weights.form`, `MCMC.prop.weights.diss`
Specifies the proposal distribution used in the MCMC Metropolis-Hastings algorithm for formation and dissolution, respectively. Possible choices are "TNT" or "random"; the "default". The TNT (tie / no tie) option puts roughly equal weight on selecting a dyad with or without a tie as a candidate for toggling, whereas the random option puts equal weight on all possible dyads, though the interpretation of random may change according to the constraints in place. When no constraints are in place, the default is TNT, which appears to improve Markov chain mixing particularly for networks with a low edge density, as is typical of many realistic social networks.
- `MCMC.prop.args.form`, `MCMC.prop.args.diss`
An alternative, direct way of specifying additional arguments to proposals.
- `MCMC.init.maxchanges`
Maximum number of toggles changes for which to allocate space.
- `MCMC.packagenames`
Names of packages in which to look for change statistic functions in addition to those autodetected. This argument should not be needed outside of very strange setups.
- `MCMC.init.maxedges`
Maximum number of edges expected in network.

MCMC.burnin, MCMC.burnin.mul

No longer used. See MCMC.burnin.min, MCMC.burnin.max, MCMC.burnin.pval, MCMC.burnin.pval, and MCMC.burnin.add.

Details

This function is only used within a call to the [simulate](#) function. See the usage section in [simulate.stergm](#) for details.

Value

A list with arguments as components.

See Also

[simulate.stergm](#), [simulate.formula.control.stergm](#) performs a similar function for [stergm](#).

control.stergm	<i>Auxiliary for Controlling Separable Temporal ERGM Fitting</i>
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Description

Auxiliary function as user interface for fine-tuning 'stergm' fitting.

Usage

```
control.stergm(init.form=NULL,
               init.diss=NULL,
               init.method=NULL,
               force.main = FALSE,
               MCMC.prop.weights.form="default",MCMC.prop.args.form=NULL,
               MCMC.prop.weights.diss="default",MCMC.prop.args.diss=NULL,
               MCMC.init.maxedges=20000,
               MCMC.init.maxchanges=20000,
               MCMC.packagenames=c(),
               CMLE.MCMC.burnin = 10000,
               CMLE.MCMC.interval = 100,
               CMLE.control=NULL,
               CMLE.control.form=control.ergm(init=init.form,
               MCMC.burnin=CMLE.MCMC.burnin,
               MCMC.interval=CMLE.MCMC.interval,
               MCMC.prop.weights=MCMC.prop.weights.form,
               MCMC.prop.args=MCMC.prop.args.form,
               MCMC.init.maxedges=MCMC.init.maxedges,
               MCMC.packagenames=MCMC.packagenames,
               parallel=parallel,
               parallel.type=parallel.type,
               parallel.version.check=parallel.version.check,
```

```
    force.main=force.main),
CMLE.control.diss=control.ergm(init=init.diss,
  MCMC.burnin=CMLE.MCMC.burnin,
  MCMC.interval=CMLE.MCMC.interval,
  MCMC.prop.weights=MCMC.prop.weights.diss,
  MCMC.prop.args=MCMC.prop.args.diss,
  MCMC.init.maxedges=MCMC.init.maxedges,
  MCMC.packagenames=MCMC.packagenames,
  parallel=parallel,
  parallel.type=parallel.type,
  parallel.version.check=parallel.version.check,
  force.main=force.main),
CMLE.NA.impute=c(),
CMLE.term.check.override=FALSE,
EGMME.main.method=c("Gradient-Descent"),
EGMME.MCMC.burnin.min=1000,
EGMME.MCMC.burnin.max=100000,
EGMME.MCMC.burnin.pval=0.5,
EGMME.MCMC.burnin.add=1,
MCMC.burnin=NULL, MCMC.burnin.mul=NULL,
SAN.maxit=10,
SAN.control=control.san(coef=init.form,
  SAN.prop.weights=MCMC.prop.weights.form,
  SAN.prop.args=MCMC.prop.args.form,
  SAN.init.maxedges=MCMC.init.maxedges,
SAN.burnin=round(sqrt(EGMME.MCMC.burnin.min * EGMME.MCMC.burnin.max)),
  SAN.packagenames=MCMC.packagenames,
  parallel=parallel,
  parallel.type=parallel.type,
  parallel.version.check=parallel.version.check),
SA.restarts=10,
SA.burnin=1000,
SA.plot.progress=FALSE,
SA.max.plot.points=400,
SA.plot.stats=FALSE,
SA.init.gain=0.1,
SA.gain.decay=0.5,
SA.runlength=25,
SA.interval.mul=2,
SA.init.interval=500,
SA.min.interval=20,
SA.max.interval=500,
SA.phase1.minruns=4,
SA.phase1.tries=20,
SA.phase1.jitter=0.1,
SA.phase1.max.q=0.1,
SA.phase1.backoff.rat=1.05,
SA.phase2.levels.max=40,
```

```

SA.phase2.levels.min=4,
SA.phase2.max.mc.se=0.001,
SA.phase2.repeats=400,
SA.stepdown.maxn=200,
SA.stepdown.p=0.05,
SA.stop.p=0.1,
SA.stepdown.ct=5,
SA.phase2.backoff.rat=1.1,
SA.keep.oh=0.5,
SA.keep.min.runs=8,
SA.keep.min=0,
SA.phase2.jitter.mul=0.2,
SA.phase2.maxreljump=4,
SA.guard.mul = 4,
SA.robust = FALSE,
SA.refine=c("mean","linear","none"),
SA.se=TRUE,
SA.phase3.samplesize.runs=10,
SA.restart.on.err=TRUE,
seed=NULL,
parallel=0,
parallel.type=NULL,
parallel.version.check=TRUE)

```

Arguments

`init.form`, `init.diss`

numeric or NA vector equal in length to the number of parameters in the formation/dissolution model or NULL (the default); the initial values for the estimation and coefficient offset terms. If NULL is passed, all of the initial values are computed using the method specified by `control$init.method`. If a numeric vector is given, the elements of the vector are interpreted as follows:

- Elements corresponding to terms enclosed in `offset()` are used as the fixed offset coefficients.
- Elements that do not correspond to offset terms and are not NA are used as starting values in the estimation.
- Initial values for the elements that are NA are fit using the method specified by `control$init.method`.

Passing `control.ergm(init=coef(prev.fit))` can be used to “resume” an uncovered `ergm` run, but see [enformulate.curved](#).

`init.method`

Estimation method used to acquire initial values for estimation. Unused at this time.

`force.main`

Logical: If TRUE, then force MCMC-based estimation method, even if the exact MLE can be computed via maximum pseudolikelihood estimation.

`MCMC.prop.weights.form`, `MCMC.prop.weights.diss`

Specifies the method to allocate probabilities of being proposed to dyads in the formation/dissolution phase. Defaults to “default”, which picks a reasonable

- default for the specified constraint. Possible values include "TNT", "random", though not all values may be used with all possible constraints.
- MCMC.prop.args.form, MCMC.prop.args.diss
An alternative, direct way of specifying additional arguments to the proposal in the formation/dissolution phase.
- MCMC.init.maxedges
Maximum number of edges for which to allocate space.
- MCMC.init.maxchanges
Maximum number of changes in dynamic network simulation for which to allocate space.
- MCMC.packagenames
Names of packages in which to look for change statistic functions in addition to those autodetected. This argument should not be needed outside of very strange setups.
- CMLE.MCMC.burnin
Maximum number of Metropolis-Hastings steps per phase (formation and dissolution) per time step used in CMLE fitting.
- CMLE.MCMC.interval
Number of Metropolis-Hastings steps between successive draws when running MCMC MLE.
- CMLE.control A convenience argument for specifying both CMLE.control.form and CMLE.control.diss at once. See [control.ergm](#).
- CMLE.control.form, CMLE.control.diss
Control parameters used to fit the CMLE for the formation/dissolution ERGM. See [control.ergm](#).
- CMLE.NA.impute In STERGM CMLE, missing dyads in transitioned-to networks are accommodated using methods of Handcock and Gile (2009), but a similar approach to transitioned-from networks requires much more complex methods that are not, currently, implemented. CMLE.NA.impute controls how missing dyads in transitioned-from networks are be imputed. See argument imputers of [impute.network.list](#) for details.
By default, no imputation is performed, and the fitting stops with an error if any transitioned-from networks have missing dyads.
- CMLE.term.check.override
The method `stergm{stergm}` uses at this time to fit a series of more than two networks requires certain assumptions to be made about the ERGM terms being used, which are tested before a fit is attempted. This test sometimes fails despite the model being amenable to fitting, so setting this option to TRUE overrides the tests.
- EGMME.main.method
Estimation method used to find the Equilibrium Generalized Method of Moments estimator. Currently only "Gradient-Descent" is implemented.
- EGMME.MCMC.burnin.min, EGMME.MCMC.burnin.max, EGMME.MCMC.burnin.pval, EGMME.MCMC.burnin.add
Number of Metropolis-Hastings steps per phase (formation and dissolution) per time step used in EGMME fitting. By default, this is determined adaptively by

keeping track of increments in the Hamming distance between the transitioned-from network and the network being sampled (formation network or dissolution network). Once `EGMME.MCMC.burnin.min` steps have elapsed, the increments are tested against 0, and when their average number becomes statistically indistinguishable from 0 (with the p-value being greater than `EGMME.MCMC.burnin.pval`), or `EGMME.MCMC.burnin.max` steps are proposed, whichever comes first, the simulation is stopped after an additional `EGMME.MCMC.burnin.add` times the number of elapsed steps had been taken. (Stopping immediately would bias the sampling.)

To use a fixed number of steps, set both `EGMME.MCMC.burnin.min` and `EGMME.MCMC.burnin.max` to the desired number of steps.

<code>SAN.maxit</code>	When <code>target.stats</code> argument is passed to ergm , the maximum number of attempts to use san to obtain a network with statistics close to those specified.
<code>SAN.control</code>	SAN control parameters. See control.san
<code>SA.restarts</code>	Maximum number of times to restart a failed optimization process.
<code>SA.burnin</code>	Number of time steps to advance the starting network before beginning the optimization.
<code>SA.plot.progress</code> , <code>SA.plot.stats</code>	Logical: Plot information about the fit as it proceeds. If <code>SA.plot.progress==TRUE</code> , plot the trajectories of the parameters and target statistics as the optimization progresses. If <code>SA.plot.stats==TRUE</code> , plot a heatmap representing correlations of target statistics and a heatmap representing the estimated gradient. Do NOT use these with non-interactive plotting devices like pdf . (In fact, it will refuse to do that with a warning.)
<code>SA.max.plot.points</code>	If <code>SA.plot.progress==TRUE</code> , the maximum number of time points to be plotted. Defaults to 400. If more iterations elapse, they will be thinned to at most 400 before plotting.
<code>SA.init.gain</code>	Initial gain, the multiplier for the parameter update size. If the process initially goes crazy beyond recovery, lower this value.
<code>SA.gain.decay</code>	Gain decay factor.
<code>SA.runlength</code>	Number of parameter trials and updates per C run.
<code>SA.interval.mul</code>	The number of time steps between updates of the parameters is set to be this times the mean duration of extant ties.
<code>SA.init.interval</code>	Initial number of time steps between updates of the parameters.
<code>SA.min.interval</code> , <code>SA.max.interval</code>	Upper and lower bounds on the number of time steps between updates of the parameters.
<code>SA.phase1.tries</code>	Number of runs trying to find a reasonable parameter and network configuration.
<code>SA.phase1.jitter</code>	Initial jitter standard deviation of each parameter.

SA.phase1.max.q	Q-value (false discovery rate) that a gradient estimate must obtain before it is accepted (since sign is what is important).
SA.phase1.backoff.rat, SA.phase2.backoff.rat	If the run produces this relative increase in the approximate objective function, it will be backed off.
SA.phase1.minruns	Number of runs during Phase 1 for estimating the gradient, before every gradient update.
SA.phase2.levels.min, SA.phase2.levels.max	Range of gain levels (subphases) to go through.
SA.phase2.max.mc.se	Approximate precision of the estimates that must be attained before stopping.
SA.phase2.repeats, SA.stepdown.maxn, SA.stepdown.p, SA.stepdown.ct	A gain level may be repeated multiple times (up to SA.phase2.repeats) if the optimizer detects that the objective function is improving or the estimating equations are not centered around 0, so slowing down the parameters at that point is counterproductive. To detect this it looks at the the window controlled by SA.keep.oh, thinning objective function values to get SA.stepdown.maxn, and 1) fitting a GLS model for a linear trend, with AR(2) autocorrelation and 2) conducting an approximate Hotelling's T^2 test for equality of estimating equation values to 0. If there is no significance for either at SA.stepdown.p SA.stepdown.ct runs in a row, the gain level (subphase) is allowed to end. Otherwise, the process continues at the same gain level.
SA.stop.p	At the end of each gain level after the minimum, if the precision is sufficiently high, the relationship between the parameters and the targets is tested for evidence of local nonlinearity. This is the p-value used. If that test fails to reject, a Phase 3 run is made with the new parameter values, and the estimating equations are tested for difference from 0. If this test fails to reject, the optimization is finished. If either of these tests rejects, at SA.stop.p, optimization is continued for another gain level.
SA.keep.oh, SA.keep.min, SA.keep.min.runs	Parameters controlling how much of optimization history to keep for gradient and covariance estimation. A history record will be kept if it's at least one of the following: <ul style="list-style-type: none"> • Among the last SA.keep.oh (a fraction) of all runs. • Among the last SA.keep.min (a count) records. • From the last SA.keep.min.runs (a count) optimization runs.
SA.phase2.jitter.mul	Jitter standard deviation of each parameter is this value times its standard deviation without jitter.
SA.phase2.maxreljump	To keep the optimization from "running away" due to, say, a poor gradient estimate building on itself, if a magnitude of change (Mahalanobis distance) in parameters over the course of a run divided by average magnitude of change for

	recent runs exceeds this, the change is truncated to this amount times the average for recent runs.
SA.guard.mul	The multiplier for the range of parameter and statistics values to compute the guard width.
SA.robust	Whether to use robust linear regression (for gradients) and covariance estimation.
SA.refine	Method, if any, used to refine the point estimate at the end: "linear" for linear interpolation, "mean" for average, and "none" to use the last value.
SA.se	Logical: If TRUE (the default), get an MCMC sample of statistics at the final estimate and compute the covariance matrix (and hence standard errors) of the parameters. This sample is stored and can also be used by mcmc.diagnostics.stergm to assess convergence.
SA.phase3.samplesize.runs	This many optimization runs will be used to determine whether the optimization has converged and to estimate the standard errors.
SA.restart.on.err	Logical: if TRUE (the default) an error somewhere in the optimization process will cause it to restart with a smaller gain value. Otherwise, the process will stop. This is mainly used for debugging
seed	Seed value (integer) for the random number generator. See set.seed
parallel	Number of threads in which to run the sampling. Defaults to 0 (no parallelism). See the entry on parallel processing for details and troubleshooting.
parallel.type	API to use for parallel processing. Supported values are "MPI" and "SOCK". Defaults to using the snow package default.
parallel.version.check	Logical: If TRUE, check that the version of ergm running on the slave nodes is the same as that running on the master node.
MCMC.burnin, MCMC.burnin.mul	No longer used. See EGMME.MCMC.burnin.min , EGMME.MCMC.burnin.max , EGMME.MCMC.burnin.pval , EGMME.MCMC.burnin.pval , EGMME.MCMC.burnin.add and CMLE.MCMC.burnin and CMLE.MCMC.interval .

Details

This function is only used within a call to the [stergm](#) function. See the usage section in [stergm](#) for details.

Value

A list with arguments as components.

References

- Boer, P., Huisman, M., Snijders, T.A.B., and Zeggelink, E.P.H. (2003), StOCNET User's Manual. Version 1.4.
- Firth (1993), Bias Reduction in Maximum Likelihood Estimates. *Biometrika*, 80: 27-38.

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See Also

`stergm`. The `control.simulate.stergm` function performs a similar function for `simulate.stergm`.

ergm-constraints	<i>Formation and Dissolution Constraints for Exponential Family Random Graph Models</i>
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Description

This page describes the network sample space constraints that are included with the `tergm` package. For more information, and instructions for using constraints, see `ergm-constraints` and `ergm`.

Constraints implemented in the `tergm` package

- `atleast(nw)` *The Formation Constraint*: Preserve all ties in network `nw`. Only dyads that are not ties in `nw` may be changed.
- `atmost(nw)` *The Dissolution Constraint*: Prevent all nonties in network `nw`. Only dyads that have ties in `nw` may be changed.

References

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- Goodreau SM, Handcock MS, Hunter DR, Butts CT, Morris M (2008a). A **statnet** Tutorial. *Journal of Statistical Software*, 24(8). <http://www.jstatsoft.org/v24/i08/>.
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Description

Unlike ordinary [ergm-terms](#), which take only a single network as an argument, the terms documented here also take into account the "ages" of extant ties in the network: the time elapsed since their formation.

As implemented, many of these terms cannot be used to "drive" the process of network evolution, but they can be used as target statistics to infer the terms that do. More concretely, they may appear in `targets=` or `monitor=` formulas of `stergm`, `simulate.stergm`, or `summary` (with an ERGM formula), but they may not appear in their `formation=` and `dissolution=` formulas. These terms are marked with "(target-only)".

All terms listed here are binary.

Terms to represent network statistics included in the `tergm` package

`degrange.mean.age(from, to=+Inf, byarg=NULL, emptyval=0)` (**target-only**) *Average age of ties incident on nodes having degree in a given range:* The `from` and `to` arguments are vectors of distinct integers or `+Inf`, for `to`. If one of the vectors has length 1, it is recycled to the length of the other. Otherwise, they must have the same length. This term adds one network statistic to the model for each element of `from` (or `to`); the i th such statistic equals the average, among all ties incident on nodes with degree greater than or equal to `from[i]` but strictly less than `to[i]`, of the amount of time elapsed since the tie's formation. The optional argument `by` is a character string giving the name of an attribute in the network's vertex attribute list. If specified, then separate degree statistics are calculated for nodes having each separate value of the attribute.

Because this average is undefined for a network that does not have any actors with degree in the specified range, the argument `emptyval` can be used to specify the value returned if this is the case. This is, technically, an arbitrary value, but it should not have a substantial effect unless a non-negligible fraction of networks at the parameter configuration of interest has no actors with specified degree.

`degree.mean.age(d, by=NULL, emptyval=0)` (**target-only**) *Average age of ties incident on nodes having a given degree:* The `d` argument is a vector of distinct integers. This term adds one network statistic to the model for each element in `d`; the i th such statistic equals the average, among all ties incident on nodes with degree exactly `d[i]`, of the amount of time elapsed since the tie's formation. The optional argument `by` is a character string giving the name of an attribute in the network's vertex attribute list. If specified, then separate degree statistics are calculated for nodes having each separate value of the attribute.

Because this average is undefined for a network that does not have any actors with degree `d[i]`, the argument `emptyval` can be used to specify the value returned if this is the case. This is, technically, an arbitrary value, but it should not have a substantial effect unless a non-negligible fraction of networks at the parameter configuration of interest has no actors with specified degree.

- `edges.ageinterval(from, to=+Inf)` (**dissolution- and target-only**) *Number of edges with age falling into a specified range:* This term counts the number of edges in the network for which the time elapsed since formation is greater than or equal to `from` but strictly less than `to`. In other words, it is in the semiopen interval $[from, to)$. `from` and `to` may be scalars, vectors of the same length, or one of them must have length one, in which case it is recycled.
- When used in the dissolution formula of a STERGM, it can be used to model a non-Markovian dissolution process, controlling the hazard function in the interval directly.
- `edge.ages` (**target-only**) *Sum of ages of extant ties:* This term adds one statistic equaling sum, over all ties present in the network, of the amount of time elapsed since formation.
- Unlike `mean.age`, this statistic is well-defined on an empty network. However, if used as a target, it appears to produce highly biased dissolution parameter estimates if the goal is to get an intended average duration.
- `edgecov.ages(x, attrname=NULL)` (**target-only**) *Weighted sum of ages of extant ties:* This term adds one statistic equaling sum, over all ties present in the network, of the amount of time elapsed since formation, multiplied by a dyadic covariate. See the help for the `edgecov` term for details for specifying the covariate.
- "Weights" can be negative.
- Unlike `edgecov.mean.age`, this statistic is well-defined on an empty network. However, if used as a target, it appears to produce highly biased dissolution parameter estimates if the goal is to get an intended average duration.
- `edgecov.mean.age(x, attrname=NULL, emptyval=0)` (**target-only**) *Weighted average age of an extant tie:* This term adds one statistic equaling the average, over all ties present in the network, of the amount of time elapsed since formation, weighted by a (nonnegative) dyadic covariate. See the help for the `edgecov` term for details for specifying the covariate.
- The behavior when there are negative weights is undefined.
- Because this average is undefined for an empty network (or a network all of whose extant edges have been weighted 0), the argument `emptyval` can be used to specify the value returned if this is the case. This is, technically, an arbitrary value, but it should not have a substantial effect unless a non-negligible fraction of networks at the parameter configuration of interest is empty and/or if only a few dyads have nonzero weights.
- `mean.age(emptyval=0)` (**target-only**) *Average age of an extant tie:* This term adds one statistic equaling the average, over all ties present in the network, of the amount of time elapsed since formation.
- Because this average is undefined for an empty network, the argument `emptyval` can be used to specify the value returned if it is. This is, technically, an arbitrary value, but it should not have a substantial effect unless a non-negligible fraction of networks at the parameter configuration of interest is empty.

References

- Handcock M. S., Hunter D. R., Butts C. T., Goodreau S. G., Krivitsky P. N. and Morris M. (2012). `_Fit, Simulate and Diagnose Exponential-Family Models for Networks_`. Version 3.1. Project home page at <URL: <http://www.statnet.org>>, <URL: CRAN.R-project.org/package=ergm>.
- Krivitsky, P.N. (2012). Modeling of Dynamic Networks based on Egocentric Data with Durational Information. *Pennsylvania State University Department of Statistics Technical Report*, 2012(2012-01). <http://stat.psu.edu/Research/2012-technical-reports>

- Krivitsky, P.N. (2012). Modeling Tie Duration in ERGM-Based Dynamic Network Models. *Pennsylvania State University Department of Statistics Technical Report*, 2012(2012-02). <http://stat.psu.edu/Research/2012-technical-reports>

See Also

[ergm-terms](#) (from the [ergm](#) package), [ergm](#), [network](#), [%v%](#), [%n%](#)

gof.stergm

Goodness-of-fit methods for STERGM CMLE and CMPLE fits

Description

For now, these are simple wrappers around [gof.ergm](#), [print.gofobject](#), [summary.gofobject](#), and [plot.gofobject](#), respectively, to run goodness-of-fit for formation and dissolution models separately. This may change in the future.

Usage

```
## S3 method for class 'stergm'
gof(object, ...)
## S3 method for class 'gof.stergm'
print(x, ...)
## S3 method for class 'gof.stergm'
summary(object, ...)
## S3 method for class 'gof.stergm'
plot(x, ..., main="Goodness-of-fit diagnostics")
```

Arguments

object, x	For gof.stergm , stergm conditional MLE (CMLE) or conditional MPLE (CMPLE) fit. For the others, a gof.stergm object returned by gof.stergm .
main	Gives the title of the goodness-of-fit plots, which will have "Formation:" and "Dissolution:" prepended to it.
...	Additional arguments passed through to the respective functions in the ergm package.

Value

For [gof.stergm](#), an object of class [gof.stergm](#), which is simply a list with two named elements: formation and dissolution, each of them a [gofobject](#) returned by [gof.ergm](#).

For the others, nothing.

See Also

[stergm](#), [ergm](#), [simulate.stergm](#), [print.gofobject](#), [plot.gofobject](#), [summary.gofobject](#), [mcmc.diagnostics.ergm](#)

Examples

```
data(samplk)

# Fit a transition from Time 1 to Time 2
samplk12 <- stergm(list(samplk1, samplk2),
  formation=~edges+mutual+transitiveties+cyclicalties,
  dissolution=~edges+mutual+transitiveties+cyclicalties,
  estimate="CMLE")

samplk12.gof <- gof(samplk12)

samplk12.gof

summary(samplk12.gof)

plot(samplk12.gof)

plot(samplk12.gof, plotlogodds=TRUE)
```

impute.network.list *Impute missing dyads in a series of networks*

Description

This function takes a list of networks with missing dyads and returns a list of networks with missing dyads imputed according to a list of imputation directives.

Usage

```
impute.network.list(nwl, imputers = c(), nwl.prepend = list(), nwl.append = list())
```

Arguments

nwl	A list of network objects or a network.list object.
imputers	A character vector giving one or more methods to impute missing dyads. Currently implemented methods are as follows: <ul style="list-style-type: none">"next" Impute the state of the same dyad in the next network in the list (or later, if that one is also missing). This imputation method is likely to lead to an underestimation of the formation and dissolution rates. The last network in the list cannot be imputed this way."previous" Impute the state of the same dyad in the previous network in the list (or earlier, if that one is also missing). The first network in the list cannot be imputed this way.

"majority" Impute the missing dyad with the value of the majority among the non-missing dyads in that time step's network. A network that has exactly the same number of ties as non-missing non-ties cannot be imputed this way.

"0" Assume missing dyads are all non-ties.

"1" Assume missing dyads are all ties.

If `length(imputers)>1` the specified imputation methods will be applied in succession. For example, `imputers=c("next", "previous", "majority", "0")` would first try to impute a missing dyad with the next time step's value. If it, and all of the later values for that dyad are missing, it will try to impute it with the previous time step's value. If it, and all of the earlier values for that dyad are missing as well, it will try to impute it with the value of the majority of non-missing dyads for that time step. If there is an exact tie, it will impute 0.

<code>nw1.prepend</code>	An optional list of networks to treat as preceding those in <code>nw1</code> . They will not be imputed or returned, but they can be useful for imputing dyads in the first network in <code>nw1</code> , when using "previous" imputer.
<code>nw1.append</code>	An optional list of networks to treat as following those in <code>nw1</code> . They will not be imputed or returned, but they can be useful for imputing dyads in the last network in <code>nw1</code> , when using "next" imputer.

Value

A list of networks with missing dyads imputed.

See Also

[network](#), [is.na](#)

<code>logLik.stergm</code>	A logLik method for stergm .
----------------------------	--

Description

Functions to return the log-likelihood associated with a [stergm](#) CMLE fit, evaluating it if necessary. See [logLik.ergm](#) documentation for details and caveats.

Usage

```
## S3 method for class 'stergm'
logLik(object,
        add=FALSE,
        force.reeval=FALSE,
        eval.loglik=add || force.reeval,
        control=control.logLik.stergm(),
        ...)
```

```
## S3 method for class 'stergm'
logLikNull(object, control=control.logLik.stergm(),
           ...)
```

Arguments

object	A stergm fit, returned by stergm , for estimate="CMLE".
add	Logical: If TRUE, instead of returning the log-likelihood, return object with log-likelihood value set.
force.reeval	Logical: If TRUE, reestimate the log-likelihood even if object already has an estimate.
eval.loglik	Logical: If TRUE, evaluate the log-likelihood if not set on object.
control	A list of control parameters for algorithm tuning. Constructed using control.logLik.ergm .
...	Other arguments to the likelihood functions.

Details

If the log-likelihood was not computed for object, produces an error unless `eval.loglik=TRUE`

Value

For `logLik.stergm`, `add=FALSE` (the default), a [logLik](#) object. If `add=TRUE` (the default), an [ergm](#) object or a [stergm](#) object with the log-likelihood set. For `logLikNull.stergm`, a [logLik](#) object.

References

Hunter, D. R. and Handcock, M. S. (2006) *Inference in curved exponential family models for networks*, Journal of Computational and Graphical Statistics.

See Also

[logLik](#), [ergm.bridge.llr](#), [ergm.bridge.dindstart.llk](#)

mcmc.diagnostics

Conduct MCMC diagnostics on an ergm or stergm fit

Description

This function prints diagnostic information and creates simple diagnostic plots for the MCMC sampled statistics produced from a [stergm](#) fit.

Usage

```
## S3 method for class 'stergm'
mcmc.diagnostics(object,
                  center=TRUE,
                  curved=TRUE,
                  vars.per.page=3,
                  ...)
```

Arguments

object	A stergm object. See documentation for stergm .
center	Logical: If TRUE, ; center the samples on the observed statistics.
curved	Logical: If TRUE, summarize the curved statistics (evaluated at the MLE of any non-linear parameters), rather than the raw components of the curved statistics.
vars.per.page	Number of rows (one variable per row) per plotting page. Ignored if latticeExtra package is not installed.
...	Additional arguments, to be passed to plotting functions.

Details

The plots produced are a trace of the sampled output and a density estimate for each variable in the chain. The diagnostics printed include correlations and convergence diagnostics.

In fact, an object contains the matrix of statistics from the MCMC run as component `$sample`. This matrix is actually an object of class `mcmc` and can be used directly in the coda package to assess MCMC convergence. *Hence all MCMC diagnostic methods available in coda are available directly.* See the examples and <http://www.mrc-bsu.cam.ac.uk/bugs/classic/coda04/readme.shtml>.

More information can be found by looking at the documentation of [stergm](#).

Value

`mcmc.diagnostics.ergm` returns some degeneracy information, if it is included in the original object. The function is mainly used for its side effect, which is to produce plots and summary output based on those plots.

References

Raftery, A.E. and Lewis, S.M. (1992). One long run with diagnostics: Implementation strategies for Markov chain Monte Carlo. *Statistical Science*, 7, 493-497.

Raftery, A.E. and Lewis, S.M. (1995). The number of iterations, convergence diagnostics and generic Metropolis algorithms. In *Practical Markov Chain Monte Carlo* (W.R. Gilks, D.J. Spiegelhalter and S. Richardson, eds.). London, U.K.: Chapman and Hall.

This function is based on the coda package It is based on the the R function `raftery.diag` in coda. `raftery.diag`, in turn, is based on the FORTRAN program `gibbsit` written by Steven Lewis which is available from the Statlib archive.

See Also

[ergm](#), [stergm](#), network package, coda package, [summary.ergm](#)

print.stergm

Separable Temporal Exponential Random Graph Models

Description

[print.stergm](#) and [summary.stergm](#) are the method used to print and summarize objects created by the [stergm](#) function.

Usage

```
## S3 method for class 'stergm'  
print(x, digits = max(3, getOption("digits") - 3), ...)  
## S3 method for class 'stergm'  
summary(object, ...)
```

Arguments

x, object	A stergm object. See documentation for stergm .
digits	Significant digits for coefficients
...	Additional arguments, to be passed to lower-level functions in the future.

Details

Automatically called when an object of class [stergm](#) is printed.

Value

The value returned is the [stergm](#) object itself.

See Also

[network](#), [stergm](#)

simulate.stergm	<i>Draw from the distribution of an Separable Temporal Exponential Family Random Graph Model</i>
-----------------	--

Description

`simulate` is used to draw from separable temporal exponential family random network models in their natural parameterizations. See `stergm` for more information on these models.

Usage

```
## S3 method for class 'stergm'
simulate(object, nsim=1, seed=NULL,
  coef.form = object$formation.fit$coef,
  coef.diss = object$dissolution.fit$coef,
  constraints = object$constraints,
  monitor = object$targets,
  time.slices=1, time.start=NULL, time.burnin=0, time.interval=1,
  control=control.simulate.stergm(),
  statsonly=NULL,
  output=c("networkDynamic", "stats", "changes", "final"),
  nw.start = NULL,
  stats.form = FALSE,
  stats.diss = FALSE,
  verbose=FALSE,
  ...)
## S3 method for class 'network'
simulate(object, nsim=1, seed=NULL,
  formation, dissolution,
  coef.form, coef.diss,
  constraints = ~.,
  monitor = NULL,
  time.slices=1, time.start=NULL, time.burnin=0, time.interval=1, time.offset=1,
  control=control.simulate.network(),
  statsonly=NULL,
  output=c("networkDynamic", "stats", "changes", "final"),
  stats.form = FALSE,
  stats.diss = FALSE,
  verbose=FALSE,
  ...)
## S3 method for class 'networkDynamic'
simulate(object, nsim=1, seed=NULL,
  formation = attr(object, "formation"),
  dissolution = attr(object, "dissolution"),
  coef.form = attr(object, "coef.form"),
  coef.diss = attr(object, "coef.diss"),
  constraints = NVL(attr(object, "constraints"), ~.),
```

```

monitor = attr(object, "monitor"),
time.slices=1, time.start=NULL, time.burnin=0, time.interval=1, time.offset=1,
control=control.simulate.network(),
statonly=NULL,
output=c("networkDynamic", "stats", "changes"),
stats.form = FALSE,
stats.diss = FALSE,
verbose=FALSE,
...)

```

Arguments

object	an R object of type <code>stergm</code> giving a model fit or of type <code>network</code> giving the initial network. simulate.network understands the <code>lasttoggle</code> "API".
formation, dissolution	One-sided <code>ergm</code> -style formulas for the formation and dissolution models, respectively.
nsim	Number of replications (separate chains of networks) of the process to run and return. The <code>networkDynamic</code> method only supports <code>nsim=1</code> .
seed	Random number integer seed. See <code>set.seed</code> .
coef.form	Parameters for the model from which the post-formation network is drawn.
coef.diss	As <code>coef.form</code> , but for the post-dissolution network.
constraints	A one-sided formula specifying one or more constraints on the support of the distribution of the networks being modeled, using syntax similar to the <code>formula</code> argument. Multiple constraints may be given, separated by "+" operators. Together with the model terms in the formula and the reference measure, the constraints define the distribution of networks being modeled. It is also possible to specify a proposal function directly by passing a string with the function's name. In that case, arguments to the proposal should be specified through the <code>prop.args</code> argument to <code>control.ergm</code> . The default is <code>~.</code> , for an unconstrained model. See the ERGM constraints documentation for the constraints implemented in the <code>ergm</code> package. Other packages may add their own constraints. For STERGMs in particular, the constraints apply to the post-formation and the post-dissolution network, rather than the final network. This means, for example, that if the degree of all vertices is constrained to be less than or equal to three, and a vertex begins a time step with three edges, then, even if one of its edges is dissolved during its time step, it won't be able to form another edge until the next time step. This behavior may change in the future. Note that not all possible combinations of constraints are supported.
monitor	Either a one-sided formula specifying one or more terms whose value is to be monitored, or a string containing "formation" or "dissolution", to monitor their respective terms, or "all" to monitor distinct terms from both.
time.slices	Number of time slices (or statistics) to return from each replication of the dynamic process. See below for return types. Defaults to 1, which, if <code>time.burnin==0</code> and <code>time.interval==1</code> (the defaults), advances the process one time step.

<code>time.start</code>	An optional argument specifying the time point at which the simulation is to start. See Details for further information.
<code>time.burnin</code>	Number of time steps to discard before starting to collect network statistics. Actual network will only be returned if <code>time.burnin==0</code> .
<code>time.interval</code>	Number of time steps between successive recordings of network statistics. Actual network will only be returned if <code>time.interval==1</code> .
<code>time.offset</code>	Argument specifying the offset between the point when the state of the network is sampled (<code>time.start</code>) and the beginning of the spell that should be recorded for the newly simulated network state.
<code>control</code>	A list of control parameters for algorithm tuning. Constructed using <code>control.simulate.stergm</code> or <code>control.simulate.network</code> .
<code>statonly</code>	Deprecated in favor of <code>output</code> .
<code>output</code>	A character vector specifying output type: one of "networkDynamic" (the default), "stats", and "changes", with partial matching allowed. See Value section for details.
<code>nw.start</code>	A specification for the starting network to be used by <code>simulate.stergm</code> , optional for EGMME fits, but required for CMLE and CMPLC fits: a numeric index <code>i</code> : use <code>i</code> th time-point's network, where the first network in the series used to fit the model is defined to be at the first time point; strings "first" or "last" : the first or last time point used in fitting the model; or a network object : specify the network directly. <code>networkDynamics</code> cannot be used as starting networks for <code>simulate.stergm</code> at this time. (They can be used as starting networks for <code>simulate.networkDynamic</code> , of course.)
<code>stats.form</code> , <code>stats.diss</code>	Logical: Whether to return formation/dissolution model statistics. This is not the recommended method: use <code>monitor</code> argument instead.
<code>verbose</code>	Logical: If TRUE, extra information is printed as the Markov chain progresses.
<code>...</code>	Further arguments passed to or used by methods.

Details

The dynamic process is run forward and the results are returned. For the method for `networkDynamic`, the simulation is resumed from the last generated time point of `object`, by default with the same model and parameters.

The starting network for the `stergm` object method (`simulate.stergm`) is determined by the `nw.start` argument.

The time index of the start of the simulation is determined as follows:

- If `time.start` is specified, it is used as the initial time index of the simulation.
- If `time.start` is not specified (is NULL), then if the object carries a time stamp from which to start or resume the simulation, either in the form of a "time" network attribute (for the `network` method — see the `lasttoggle` "API") or in the form of an `net.obs.period` network attribute (for the `networkDynamic` method), this attribute will be used. (If specified, `time.start` will override it with a warning.)

- Otherwise, the simulation starts at 0.

Value

Depends on the output argument:

"stats"	<p>If <code>stats.form==FALSE</code> and <code>stats.diss==FALSE</code>, an <code>mcmc</code> matrix with monitored statistics, and if either of them is <code>TRUE</code>, a list containing elements <code>stats</code> for statistics specified in the <code>monitor</code> argument, and <code>stats.form</code> and <code>stats.diss</code> for the respective formation and dissolution statistics.</p> <p>If <code>stats.form==FALSE</code> and <code>stats.diss==FALSE</code> and no monitored statistics are specified, an empty list is returned, with a warning.</p> <p>When <code>nsim>1</code>, an <code>mcmc.list</code> (or list of them) of the statistics is returned instead.</p>
"networkDynamic"	<p>A <code>networkDynamic</code> object representing the simulated process, with ties present in the initial network having onset <code>-Inf</code> and ties present at the end of the simulation having terminus <code>+Inf</code>. The method for <code>networkDynamic</code> returns the initial <code>networkDynamic</code> with simulated changes applied to it. The <code>net.obs.period</code> network attribute is updated (or added if not existing) to reflect the time period that was simulated.</p> <p>Additionally, attributes (<code>attr</code>, not network attributes) are attached as follows:</p> <p><code>formation, dissolution, monitor</code>: Formation, dissolution, and monitoring formulas used in the simulation, respectively.</p> <p><code>stats, stats.form, stats.diss</code>: Network statistics as above.</p> <p><code>coef.form, coef.diss</code>: Coefficients used in the simulation.</p> <p><code>changes</code>: A four-column matrix summarizing the changes in the "changes" output. (This may be removed in the future.)</p> <p>When <code>nsim>1</code>, a <code>network.list</code> of these <code>networkDynamics</code> is returned.</p>
"changes"	<p>An integer matrix with four columns (<code>time</code>, <code>tail</code>, <code>head</code>, and <code>to</code>), giving the timestamped changes relative to the current network. <code>to</code> is 1 if a tie was formed and 0 if a tie was dissolved. The convention for <code>time</code> is that it gives the time point during which the change is effective. For example, a row <code>c(5,2,3,1)</code> indicates that between time 4 and 5, a tie from node 2 to node 3 was formed, so that it was absent at time point 4 and present at time point 5; while a row <code>c(5,2,3,0)</code> indicates that in that time, that tie was dissolved, so that it was present at time point 4 and absent at time point 5.</p> <p>Additionally, same attributes (<code>attr</code>, not network attributes) as with <code>output=="networkDynamic"</code> are attached.</p> <p>When <code>nsim>1</code>, a list of these change matrices is returned.</p>
"final"	<p>A <code>network</code> object representing the last network in the series generated. This is not implemented in the method for <code>networkDynamic</code>.</p> <p><code>lasttoggle</code> attributes are also included.</p> <p>Additionally, attributes (<code>attr</code>, not network attributes) are attached as follows:</p> <p>formation, dissolution, monitor: Formation, dissolution, and monitoring formulas used in the simulation, respectively.</p> <p>stats, stats.form, stats.diss: Network statistics as above.</p>

coef.form, coef.diss: Coefficients used in the simulation.

changes A four-column matrix summarizing the changes in the "changes" output. (This may be removed in the future.)

When `nsim>1`, a `network.list` of these `networks` is returned.

Examples

```
logit<-function(p)log(p/(1-p))
coef.form.f<-function(coef.diss,density) -log(((1+exp(coef.diss))/(density/(1-density))))-1)

# Construct a network with 20 nodes and 20 edges
n<-20
target.stats<-edges<-20
g0<-network.initialize(n,dir=TRUE)
g1<-san(g0~edges,target.stats=target.stats,verbose=TRUE)

S<-10

# To get an average duration of 10...
duration<-10
coef.diss<-logit(1-1/duration)

# To get an average of 20 edges...
dyads<-network.dyadcount(g1)
density<-edges/dyads
coef.form<-coef.form.f(coef.diss,density)

# ... coefficients.
print(coef.form)
print(coef.diss)

# Simulate a networkDynamic
dynsim<-simulate(g1,formation=~edges,dissolution=~edges,
  coef.form=coef.form,coef.diss=coef.diss,
  time.slices=S,verbose=TRUE)

# "Resume" the simulation.
dynsim2<-simulate(dynsim,time.slices=S,verbose=TRUE)
```

Description

`stergm` is used for finding Separable Temporal ERGMs' (STERGMs) Conditional MLE (CMLE) (Krivitsky and Handcock, 2010) and Equilibrium Generalized Method of Moments Estimator (EGMME) (Krivitsky, 2009).

Usage

```
stergm (nw,
        formation,
        dissolution,
        constraints = ~.,
        estimate,
        times=NULL,
        offset.coef.form=NULL,
        offset.coef.diss=NULL,
        targets=NULL,
        target.stats=NULL,
        eval.loglik=TRUE,
        control=control.stergm(),
        verbose=FALSE,
        ...)
```

Arguments

- nw** A [network](#) object (for EGMME); or [networkDynamic](#) object, a [network.list](#) object, or a [list](#) containing networks (for CMLE and CPMLE). `simulate.network` understands the [lasttoggle](#) "API".
- formation, dissolution** One-sided [ergm](#)-style formulas for the formation and dissolution models, respectively.
- constraints** A one-sided formula specifying one or more constraints on the support of the distribution of the networks being modeled, using syntax similar to the `formula` argument. Multiple constraints may be given, separated by "+" operators. Together with the model terms in the formula and the reference measure, the constraints define the distribution of networks being modeled.
- It is also possible to specify a proposal function directly by passing a string with the function's name. In that case, arguments to the proposal should be specified through the `prop.args` argument to [control.ergm](#).
- The default is `~.`, for an unconstrained model.
- See the [ERGM constraints](#) documentation for the constraints implemented in the [ergm](#) package. Other packages may add their own constraints.
- For STERGMs in particular, the constraints apply to the post-formation and the post-dissolution network, rather than the final network. This means, for example, that if the degree of all vertices is constrained to be less than or equal to three, and a vertex begins a time step with three edges, then, even if one of its edges is dissolved during its time step, it won't be able to form another edge until the next time step. This behavior may change in the future.
- Note that not all possible combinations of constraints are supported.
- estimate** One of "EGMME" for Equilibrium Generalized Method of Moments Estimation, based on a single network with some temporal information and making an assumption that it is a product of a STERGM process running to its stationary (equilibrium) distribution; "CMLE" for Conditional Maximum Likelihood

	Estimation, modeling a transition between two networks, or "CMPLE" for Conditional Maximum PseudoLikelihood Estimation, using MPLE instead of MLE. CMPLE is extremely inaccurate at this time.
times	For CMLE and CMPLE estimation, times or indexes at which the networks whose transition is to be modeled are observed. Default to <code>c(0,1)</code> if <code>nw</code> is a <code>networkDynamic</code> and to <code>c(1:2)</code> (first two elements) if <code>nw</code> is a <code>network.list</code> or a <code>list</code> . Unused for EGMME.
offset.coef.form	Numeric vector to specify offset formation parameters.
offset.coef.diss	Numeric vector to specify offset dissolution parameters.
targets	One-sided <code>ergm</code> -style formula specifying statistics whose moments are used for the EGMME. Unused for CMLE and CMPLE.
target.stats	A vector specifying the values of the <code>targets</code> statistics that EGMME will try to match. Defaults to the statistics of <code>nw</code> . Unused for CMLE and CMPLE.
eval.loglik	Whether or not to calculate the log-likelihood of a CMLE STERGM fit. See <code>ergm</code> for details.
control	A list of control parameters for algorithm tuning. Constructed using <code>control.stergm</code> .
verbose	logical or integer; if TRUE or positive, the program will print out progress information. Higher values result in more output.
...	Additional arguments, to be passed to lower-level functions.

Value

`stergm` returns an object of class `stergm` that is a list consisting of the following elements:

formation, dissolution	Formation and dissolution formulas, respectively.
targets	The targets formula.
target.stats	The target statistics.
estimate	The type of estimate.
opt.history	A matrix containing the full trace of the EGMME optimization process: coefficients tried and target statistics simulated.
sample	An <code>mcmc</code> object containing target statistics sampled at the estimate.
covar	The full estimated variance-covariance matrix of the parameter estimates for EGMME. (Note that although the CMLE formation parameter estimates are independent of the dissolution parameter estimates due to the separability assumption, this is not necessarily the case for EGMME.)
formation.fit, dissolution.fit	For CMLE and CMPLE, <code>ergm</code> objects from fitting formation and dissolution, respectively. For EGMME, stripped down <code>ergm</code> -like lists.
network	For <code>estimate=="EGMME"</code> , the original network; for <code>estimate=="CMLE"</code> or <code>estimate=="CMPLE"</code> , a <code>network.list</code> (a discrete series of networks) to which the model was fit.
control	The control parameters used to fit the model.

See the method `print.stergm` for details on how an `stergm` object is printed. Note that the method `summary.stergm` returns a summary of the relevant parts of the `stergm` object in concise summary format.

Model Terms

See `ergm` and `ergm-terms` for details. At this time, only linear ERGM terms are allowed.

References

- Krivitsky PN, Handcock MS (2010). A Separable Model for Dynamic Networks. <http://arxiv.org/abs/1011.1937>
- Krivitsky, P.N. (2012). Modeling of Dynamic Networks based on Egocentric Data with Durational Information. *Pennsylvania State University Department of Statistics Technical Report*, 2012(2012-01). <http://stat.psu.edu/Research/2012-technical-reports>

See Also

`ergm`, `network`, `%v%`, `%n%`, `ergm-terms`

Examples

```
# EGMME Example
par(ask=FALSE)
n<-30
g0<-network.initialize(n,dir=FALSE)

#           edges, degree(1), mean.age
target.stats<-c(  n*1/2,    n*0.6,      20)

dynfit<-stergm(g0,formation = ~edges+degree(1), dissolution = ~edges,
               targets = ~edges+degree(1)+mean.age,
               target.stats=target.stats, estimate="EGMME",
               control=control.stergm(SA.plot.progress=TRUE))

par(ask=TRUE)
mcmc.diagnostics(dynfit)
summary(dynfit)

# CMLE Example
data(samplk)

# Fit a transition from Time 1 to Time 2
samplk12 <- stergm(list(samplk1, samplk2),
                   formation=~edges+mutual+transitiveties+cyclicalities,
                   dissolution=~edges+mutual+transitiveties+cyclicalities,
                   estimate="CMLE")

mcmc.diagnostics(samplk12)
summary(samplk12)
```

```
# Fit a transition from Time 1 to Time 2 and from Time 2 to Time 3 jointly
samplk123 <- stergm(list(samplk1, samplk2, samplk3),
  formation=~edges+mutual+transitiveties+cyclicalities,
  dissolution=~edges+mutual+transitiveties+cyclicalities,
  estimate="CMLE")

mcmc.diagnostics(samplk123)
summary(samplk123)
```

```
summary.statistics.networkDynamic
```

Calculation of networkDynamic statistics.

Description

A method for [summary.statistics](#) to calculate the specified statistics for an observed [networkDynamic](#). See [ergm-terms](#) for more information on the statistics that may be specified.

Usage

```
## S3 method for class 'networkDynamic'
summary.statistics(object, at, ..., basis = NULL)
```

Arguments

object	An R formula object with a networkDynamic as its LHS. (See summary.statistics for more details.)
at	A vector of time points at which to calculate the statistics.
basis	An optional networkDynamic object relative to which the statistics should be calculated.
...	Further arguments passed to or used by methods.

Value

A matrix with `length(at)` rows, one for each time point, containing the statistics measured on the network.

See Also

[ergm](#), [networkDynamic](#), [ergm-terms](#), [summary.statistics.network](#)

tergm.godfather *A function to apply a given series of changes a network.*

Description

Gives the network a series of timed proposals it can't refuse. Returns the statistics of the network, and, optionally, the final network.

Usage

```
tergm.godfather(formula, changes = NULL, toggles = changes[, -4, drop = FALSE],
  start = NULL, end = NULL, end.network = FALSE, stats.start = FALSE, verbose = FALSE,
  control = control.tergm.godfather())
```

Arguments

formula	An summary.formula -style formula, with either a network or a networkDynamic as the LHS and statistics to be computed on the RHS. If LHS is a networkDynamic , it will be used to derive the changes to the network whose statistics are wanted. Otherwise, either changes or toggles must be specified, and the LHS network will be used as the starting network.
changes	A matrix with four columns: time, tail, head, and new value, describing the changes to be made. Can only be used if LHS of formula is not a networkDynamic .
toggles	A matrix with three columns: time, tail, and head, giving the dyads which had changed. Can only be used if LHS of formula is not a networkDynamic .
start	Time from which to start applying changes. Note that the first set of changes will take effect at start+1. Defaults to the time point 1 before the earliest change passed.
end	Time from which to finish applying changes. Defaults to the last time point at which a change occurs.
end.network	Whether to return a network that results. Defaults to FALSE.
stats.start	Whether to return the network statistics at start (before any changes are applied) as the first row of the statistics matrix. Defaults to FALSE, to produce output similar to that of simulate for STERGMs when output="stats", where initial network's statistics are not returned.
verbose	Whether to print progress messages.
control	A control list generated by control.tergm.godfather . So far, there is only one argument: <code>GF.init.maxedges.mul</code> , to control how much space is allocated for the edgelist of the final network. It is used adaptively, so should not be greater than 10.

Value

If `end.network==FALSE` (the default), an `mcmc` object with the requested network statistics associated with the network series produced by applying the specified changes. Its `mcmc` attributes encode the timing information: so `start(out)` gives the time point associated with the first row returned, and `end(out)` out the last. The "thinning interval" is always 1.

If `end.network==TRUE`, return a `network` object with `lasttoggle` "extension", representing the final network, with a matrix of statistics described in the previous paragraph attached to it as an `attr`-style attribute "stats".

See Also

`simulate.stergm`, `simulate.network`, `simulate.networkDynamic`

Examples

```
g1 <- network.initialize(10, dir=FALSE)
g1[1,2] <- 1
g1[3,4] <- 1
g1 %n% "time" <- 0
g1 %n% "lasttoggle" <- -1-rgeom(network.dyadcount(g1),1/4)

dc <- matrix(rnorm(100),10,10); dc <- dc+t(dc)

# Simulate a network, tracking its statistics.
simnet <- simulate(g1, formation=~edges, dissolution=~edges, coef.form=-1, coef.diss=1,
                  time.slices=50, monitor=~degree(1)+mean.age+degree.mean.age(1)+
                  edge.ages+edgecov(dc)+edgecov.ages(dc),
                  output="networkDynamic")

sim.stats <- attr(simnet, "stats")

print(head(sim.stats))
sim.stats <- as.matrix(sim.stats)

# Replay the simulation using a networkDynamic, monitoring a potentially different set of
# statistics (but same in this case).
gf1.stats <- tergm.godfather(simnet~degree(1)+mean.age+degree.mean.age(1)+
                             edge.ages+edgecov(dc)+edgecov.ages(dc),
                             start=0, end=50)

print(head(gf1.stats))
gf1.stats <- as.matrix(gf1.stats)

# Replay the simulation using the initial network + list of changes.

gf2.stats <- tergm.godfather(g1~degree(1)+mean.age+degree.mean.age(1)+
                             edge.ages+edgecov(dc)+edgecov.ages(dc),
                             start=0, end=50, changes=attr(simnet,"changes"))

print(head(gf2.stats))
```



```
gf2.stats <- as.matrix(gf2.stats)

# We can also compare them to the network statistics summarized.
summ.stats <- summary(simnet~degree(1)+mean.age+degree.mean.age(1)+
                      edge.ages+edgescov(dc)+edgescov.ages(dc), at=1:50)

print(head(summ.stats))

tol <- sqrt(.Machine$double.eps)
# If they aren't all identical, we are in trouble.
stopifnot(all.equal(sim.stats,gf1.stats),
          all.equal(sim.stats,gf2.stats),
          all.equal(sim.stats,summ.stats))
```

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