

# Package ‘EpiModel’

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**Title** Mathematical Modeling of Infectious Disease Dynamics

**Description** Tools for simulating mathematical models of infectious disease dynamics. Epidemic model classes include deterministic compartmental models, stochastic individual-contact models, and stochastic network models. Network models use the robust statistical methods of exponential-family random graph models (ERGMs) from the Statnet suite of software packages in R. Standard templates for epidemic modeling include SI, SIR, and SIS disease types. EpiModel features an API for extending these templates to address novel scientific research aims.

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**License** GPL-3

**URL** <http://epimodel.org/>, <http://github.com/statnet/EpiModel>

**BugReports** <https://github.com/statnet/EpiModel/issues>

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EpiModel-package	<i>Mathematical Modeling of Infectious Disease Dynamics</i>
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Description

Package:	EpiModel
Type:	Package
Version:	1.7.5
Date:	2019-12-29
License:	GPL-3
LazyLoad:	yes

Details

The EpiModel software package provides tools for building, solving, and visualizing mathematical models of infectious disease dynamics. These tools allow users to simulate epidemic models in multiple frameworks for both pedagogical purposes ("base models") and novel research purposes ("extension models").

Model Classes and Infectious Disease Types

EpiModel provides functionality for three classes of epidemic models:

- **Deterministic Compartmental Models:** these continuous-time models are solved using ordinary differential equations. EpiModel allows for easy specification of sensitivity analyses to compare multiple scenarios f the same model across different parameter values.
- **Stochastic Individual Contact Models:** a novel class of individual-based, microsimulation models that were developed to add random variation in all components of the transmission system, from infection to recovery to vital dynamics (arrivals and departures).
- **Stochastic Network Models:** with the underlying statistical framework of temporal exponential random graph models (ERGMs) recently developed in the **Statnet** suite of software

in R, network models over epidemics simulate edge (e.g., partnership) formation and dissolution stochastically according to a specified statistical model, with disease spread across that network.

EpiModel supports three infectious disease types to be run across all of the three classes.

- **Susceptible-Infectious (SI):** a two-state disease in which there is life-long infection without recovery. HIV/AIDS is one example, although for this case it is common to model infection stages as separate compartments.
- **Susceptible-Infectious-Recovered (SIR):** a three-stage disease in which one has life-long recovery with immunity after infection. Measles is one example, but modern models for the disease also require consideration of vaccination patterns in the population.
- **Susceptible-Infectious-Susceptible (SIS):** a two-stage disease in which one may transition back and forth from the susceptible to infected states throughout life. Examples include bacterial sexually transmitted diseases like gonorrhea.

These basic disease types may be extended in any arbitrarily complex way to simulate specific diseases for research questions.

## Model Parameterization and Simulation

EpiModel uses three model setup functions for each model class to input the necessary parameters, initial conditions, and control settings:

- `param.dcm`, `param.icm`, and `param.net` are used to input epidemic parameters for each of the three model classes. Parameters include the rate of contacts or acts between actors, the probability of transmission per contact, and recovery and demographic rates for models that include those transitions.
- `init.dcm`, `init.icm`, and `init.net` are used to input the initial conditions for each class. The main conditions are limited to the numbers or, if applicable, the specific agents in the population who are infected or recovered at the simulation outset.
- `control.dcm`, `control.icm`, and `control.net` are used to specify the remaining control settings for each simulation. The core controls for base model types include the disease type, number of time steps, and number of simulations. Controls are also used to input new model functions (for DCMs) and new model modules (for ICMs and network models) to allow the user to simulate fully original epidemic models in EpiModel. See the documentation for the specific control functions help pages.

With the models parameterized, the functions for simulating epidemic models are:

- `dcm` for deterministic compartmental models.
- `icm` for individual contact models.
- Network models are simulated in a three-step process:
  1. `netest` estimates the statistical model for the network structure itself (i.e., how partnerships form and dissolve over time given the parameterization of those processes). This function is a wrapper around the `ergm` and `stergm` functions in the `ergm` and `tergm` packages. The current statistical framework for model simulation is called "egocentric inference": target statistics summarizing these formation and dissolution processes collected from an egocentric sample of the population.

2. `netdx` runs diagnostics on the dynamic model fit by simulating the base network over time to ensure the model fits the targets for formation and dissolution.
3. `netsim` simulates the stochastic network epidemic models, with a given network model fit in `netest`. Here the function requires this model fit object along with the parameters, initial conditions, and control settings as defined above.

## References

The EpiModel website is at <http://epimodel.org/>, and the source code is at <http://github.com/statnet/EpiModel>. Bug reports and feature requests are welcome there.

Our primary methods paper on EpiModel is published in the **Journal of Statistical Software**. If you use EpiModel for any research or teaching purposes, please cite this reference:

Jenness SM, Goodreau SM and Morris M. EpiModel: An R Package for Mathematical Modeling of Infectious Disease over Networks. *Journal of Statistical Software*. 2018; 84(8): 1-47. doi: 10.18637/jss.v084.i08 (<http://doi.org/10.18637/jss.v084.i08>).

We have also developed an extension package specifically for modeling HIV and related sexually transmitted infections, called EpiModelHIV and available on Github at <http://github.com/statnet/EpiModelHIV>.

---

apportion\_lr

*Apportion Least-Remainder Method*


---

## Description

Apportions a vector of values given a specified frequency distribution of those values such that the length of the output is robust to rounding and other instabilities.

## Usage

```
apportion_lr(vector.length, values, proportions, shuffled = FALSE)
```

## Arguments

<code>vector.length</code>	Length for the output vector.
<code>values</code>	Values for the output vector.
<code>proportions</code>	Proportion distribution with one number for each value. This must sum to 1.
<code>shuffled</code>	If TRUE, randomly shuffle the order of the vector.

---

as.data.frame.dcm      *Extract Model Data for Deterministic Compartmental Models*


---

## Description

This function extracts a model run from an object of class dcm into a data frame using the generic as.data.frame function.

## Usage

```
## S3 method for class 'dcm'
as.data.frame(x, row.names = NULL, optional = FALSE, run, ...)
```

## Arguments

x	An EpiModel object of class <a href="#">dcm</a> .
row.names	See <a href="#">as.data.frame.default</a> .
optional	See <a href="#">as.data.frame.default</a> .
run	Run number for model; used for multiple-run sensitivity models. If not specified, will output data from all runs in a stacked data frame.
...	See <a href="#">as.data.frame.default</a> .

## Details

Model output from dcm simulations are available as a data frame with this helper function. The output data frame will include columns for time, the size of each compartment, the overall population size (the sum of compartment sizes), and the size of each flow.

For models with multiple runs (i.e., varying parameters - see example below), the default with the run parameter not specified will output all runs in a single stacked data frame.

## Examples

```
## Example 1: One-group SIS model with varying act.rate
param <- param.dcm(inf.prob = 0.2, act.rate = seq(0.05, 0.5, 0.05),
  rec.rate = 1/50)
init <- init.dcm(s.num = 500, i.num = 1)
control <- control.dcm(type = "SIS", nsteps = 10)
mod1 <- dcm(param, init, control)
as.data.frame(mod1)
as.data.frame(mod1, run = 1)
as.data.frame(mod1, run = 10)

## Example 2: Two-group SIR model with vital dynamics
param <- param.dcm(inf.prob = 0.2, inf.prob.g2 = 0.1,
  act.rate = 3, balance = "g1",
  rec.rate = 1/50, rec.rate.g2 = 1/50,
  a.rate = 1/100, a.rate.g2 = NA,
```

```

        ds.rate = 1/100, ds.rate.g2 = 1/100,
        di.rate = 1/90, di.rate.g2 = 1/90,
        dr.rate = 1/100, dr.rate.g2 = 1/100)
init <- init.dcm(s.num = 500, i.num = 1, r.num = 0,
               s.num.g2 = 500, i.num.g2 = 1, r.num.g2 = 0)
control <- control.dcm(type = "SIR", nsteps = 10)
mod2 <- dcm(param, init, control)
as.data.frame(mod2)

```

---

as.data.frame.icm      *Extract Model Data for Stochastic Models*


---

## Description

This function extracts model simulations for objects of classes `icm` and `netsim` into a data frame using the generic `as.data.frame` function.

## Usage

```

## S3 method for class 'icm'
as.data.frame(
  x,
  row.names = NULL,
  optional = FALSE,
  out = "vals",
  sim,
  qval,
  ...
)

## S3 method for class 'netsim'
as.data.frame(x, row.names = NULL, optional = FALSE, out = "vals", sim, ...)

```

## Arguments

<code>x</code>	An <code>EpiModel</code> object of class <code>icm</code> or <code>netsim</code> .
<code>row.names</code>	See <a href="#">as.data.frame.default</a> .
<code>optional</code>	See <a href="#">as.data.frame.default</a> .
<code>out</code>	Data output to data frame: "mean" for row means across simulations, "sd" for row standard deviations across simulations, "qnt" for row quantiles at the level specified in <code>qval</code> , or "vals" for values from individual simulations.
<code>sim</code>	If <code>out="vals"</code> , the simulation number to output. If not specified, then data from all simulations will be output.
<code>qval</code>	Quantile value required when <code>out="qnt"</code> .
<code>...</code>	See <a href="#">as.data.frame.default</a> .

## Details

These methods work for both `icm` and `netsim` class models. The available output includes time-specific means, standard deviations, quantiles, and simulation values (compartment and flow sizes) from these stochastic model classes. Means, standard deviations, and quantiles are calculated by taking the row summary (i.e., each row of data corresponds to a time step) across all simulations in the model output.

## Examples

```
## Stochastic ICM SIS model
param <- param.icm(inf.prob = 0.8, act.rate = 2, rec.rate = 0.1)
init <- init.icm(s.num = 500, i.num = 1)
control <- control.icm(type = "SIS", nsteps = 10,
                      nsims = 3, verbose = FALSE)
mod <- icm(param, init, control)

# Default output all simulation runs, default to all in stacked data.frame
as.data.frame(mod)
as.data.frame(mod, sim = 2)

# Time-specific means across simulations
as.data.frame(mod, out = "mean")

# Time-specific standard deviations across simulations
as.data.frame(mod, out = "sd")

# Time-specific quantile values across simulations
as.data.frame(mod, out = "qnt", qval = 0.25)
as.data.frame(mod, out = "qnt", qval = 0.75)

## Not run:
## Stochastic SI network model
nw <- network.initialize(n = 100, directed = FALSE)
formation <- ~edges
target.stats <- 50
coef.diss <- dissolution_coefs(dissolution = ~offset(edges), duration = 20)
est <- netest(nw, formation, target.stats, coef.diss, verbose = FALSE)

param <- param.net(inf.prob = 0.5)
init <- init.net(i.num = 10)
control <- control.net(type = "SI", nsteps = 10, nsims = 3, verbose = FALSE)
mod <- netsim(est, param, init, control)

# Same data extraction methods as with ICMs
as.data.frame(mod)
as.data.frame(mod, sim = 2)
as.data.frame(mod, out = "mean")
as.data.frame(mod, out = "sd")
as.data.frame(mod, out = "qnt", qval = 0.25)
as.data.frame(mod, out = "qnt", qval = 0.75)

## End(Not run)
```



---

as.data.frame.netdx     *Extract Timed Edgelists netdx Objects*

---

## Description

This function extracts timed edgelists for objects of class netdx into a data frame using the generic as.data.frame function.

## Usage

```
## S3 method for class 'netdx'  
as.data.frame(x, row.names = NULL, optional = FALSE, sim, ...)
```

## Arguments

x	An EpiModel object of class netdx.
row.names	See <a href="#">as.data.frame.default</a> .
optional	See <a href="#">as.data.frame.default</a> .
sim	The simulation number to output. If not specified, then data from all simulations will be output.
...	See <a href="#">as.data.frame.default</a> .

## Examples

```
# Initialize and parameterize the network model  
nw <- network.initialize(n = 100, directed = FALSE)  
formation <- ~edges  
target.stats <- 50  
coef.diss <- dissolution_coefs(dissolution = ~offset(edges), duration = 20)  
  
# Model estimation  
est <- netest(nw, formation, target.stats, coef.diss, verbose = FALSE)  
  
# Simulate the network with netdx  
dx <- netdx(est, nsims = 3, nsteps = 10, keep.tedgelist = TRUE, verbose = FALSE)  
  
# Extract data from the first simulation  
as.data.frame(dx, sim = 1)  
  
# Extract data from all simulations  
as.data.frame(dx)
```

---

as.network.transmat	<i>Converts transmat infection tree into a network object</i>
---------------------	---

---

### Description

Converts the edges of the infection tree described in the [transmat](#) object into a [network](#) object, copying in appropriate edge attributes for 'at', 'infDur', 'transProb', 'actRate', and 'finalProb' and constructing a vertex attribute for 'at'.

### Usage

```
## S3 method for class 'transmat'
as.network(x, ...)
```

### Arguments

x	an object of class transmat to be converted into a network object
...	unused

---

as.phylo.transmat	<i>Convert transmat infection tree into a phylo object</i>
-------------------	--

---

### Description

Converts the edgelist matrix in the transmat object into a phylo object by doing the required reordering and labeling.

### Usage

```
## S3 method for class 'transmat'
as.phylo(x, collapse.singles, vertex.exit.times, ...)
```

### Arguments

x	An object of class "transmat", the output from <a href="#">get_transmat</a> .
collapse.singles	logical, DEPRECATED
vertex.exit.times	optional numeric vector providing the time of departure of vertices, to be used to scale the lengths of branches reaching to the tips. Index position on vector corresponds to network id. NA indicates no departure, so branch will extend to the end of the tree.
...	further arguments (unused)

## Details

Converts a `transmat` object containing information about the history of a simulated infection into a `phylo` object representation suitable for plotting as a tree with `plot.phylo`. Each infection event becomes a 'node' (horizontal branch) in the resulting phylo tree, and each network vertex becomes a 'tip' of the tree. The infection events are labeled with the vertex id of the infector to make it possible to trace the path of infection.

The infection timing information is included to position the phylo-nodes, with the lines to the tips drawn to the max time value +1 (unless `vertex.exit.times` are passed in it effectively assumes all vertices are active/alive until the end of the simulation).

If the `transmat` contains multiple infection seeds (there are multiple trees with separate root nodes) it will return a list of class 'multiPhylo', each element of which is a `phylo` object. See [read.tree](#).

Note that in EpiModel versions  $\leq 1.2.4$ , the phylo tree was constructed differently, translating network vertices to both phylo-nodes and tips and requiring 'collapse.singles' to prune it to an appropriate branching structure.

## Examples

```
set.seed(10)
nw <- network.initialize(n = 100, directed = FALSE)
formation <- ~edges
target.stats <- 50
coef.diss <- dissolution_coefs(dissolution = ~offset(edges), duration = 20)

est1 <- netest(nw, formation, target.stats, coef.diss, verbose = FALSE)

param <- param.net(Inf.prob = 0.5)
init <- init.net(i.num = 1)
control <- control.net(type = "SI", nsteps = 40, nsims = 1, verbose = FALSE,
                      use.pids = FALSE)

mod1 <- netsim(est1, param, init, control)
tm <- get_transmat(mod1)
tmPhylo <- as.phylo.transmat(tm)
plot(tmPhylo, show.node.label = TRUE,
     root.edge = TRUE,
     cex = 0.5)
```

---

check\_bip\_degdist

*Check Degree Distribution for Bipartite Target Statistics*

---

## Description

Checks for consistency in the implied network statistics of a bipartite network in which the mode size and mode-specific degree distributions are specified.

**Usage**

```
check_bip_degdist(num.m1, num.m2, deg.dist.m1, deg.dist.m2)
```

**Arguments**

num.m1	Number of nodes in mode 1.
num.m2	Number of nodes in mode 2.
deg.dist.m1	Vector with fractional degree distribution for mode 1.
deg.dist.m2	Vector with fractional degree distribution for mode 2.

**Details**

This function outputs the number of nodes of degree 0 to m, where m is the length of a fractional degree distribution vector, given that vector and the size of the mode. This utility is used to check for balance in implied degree given that fractional distribution within bipartite network simulations, in which the degree-constrained counts must be equal across modes.

**See Also**

For a detailed explanation of this function, see the tutorial: [EpiModel Network Utility Functions](#).

**Examples**

```
# An imbalanced distribution
check_bip_degdist(num.m1 = 500, num.m2 = 500,
                  deg.dist.m2 = c(0.40, 0.55, 0.03, 0.02),
                  deg.dist.m1 = c(0.48, 0.41, 0.08, 0.03))

# A balanced distribution
check_bip_degdist(num.m1 = 500, num.m2 = 500,
                  deg.dist.m1 = c(0.40, 0.55, 0.04, 0.01),
                  deg.dist.m2 = c(0.48, 0.41, 0.08, 0.03))
```

---

color\_tea

---

*Creates a TEA Variable for Infection Status for ndtv Animations*


---

**Description**

Creates a new color-named temporally-extended attribute (TEA) variable in a networkDynamic object containing a disease status TEA in numeric format.

**Usage**

```
color_tea(
  nd,
  old.var = "testatus",
  old.sus = "s",
  old.inf = "i",
  old.rec = "r",
  new.var = "ndtvcol",
  new.sus,
  new.inf,
  new.rec,
  verbose = TRUE
)
```

**Arguments**

nd	An object of class networkDynamic.
old.var	Old TEA variable name.
old.sus	Status value for susceptible in old TEA variable.
old.inf	Status value for infected in old TEA variable.
old.rec	Status value for recovered in old TEA variable.
new.var	New TEA variable name to be stored in networkDynamic object.
new.sus	Status value for susceptible in new TEA variable.
new.inf	Status value for infected in new TEA variable.
new.rec	Status value for recovered in new TEA variable.
verbose	Print progress to console.

**Details**

The ndtv package (<https://cran.r-project.org/package=ndtv>) produces animated visuals for dynamic networks with evolving edge structures and nodal attributes. Nodal attribute dynamics in ndtv movies require a temporally extended attribute (TEA) containing a standard R color for each node at each time step. By default, the EpiModel package uses TEAs to store disease status history in network model simulations run in netsim. But, that status TEA is in numeric format (0, 1, 2). The color\_tea function transforms those numeric values of that disease status TEA into a TEA with color values in order to visualize status changes in ndtv.

The convention in plot.netsim is to color the susceptible nodes as blue, infected nodes as red, and recovered nodes as green. Alternate colors may be specified using the new.sus, new.inf, and new.rec parameters, respectively.

Using the color\_tea function with a netsim object requires that TEAs for disease status be used and that the networkDynamic object be saved in the output: both tea.status and save.network must be set to TRUE in control.net.

**See Also**

netsim and the ndtv package documentation.

comp\_plot

*Plot Compartment Diagram for Epidemic Models***Description**

Plots a compartment flow diagram for deterministic compartmental models, stochastic individual contact models, and stochastic network models.

**Usage**

```
comp_plot(x, at, digits, ...)

## S3 method for class 'dcm'
comp_plot(x, at = 1, digits = 3, run = 1, ...)

## S3 method for class 'icm'
comp_plot(x, at = 1, digits = 3, ...)

## S3 method for class 'netsim'
comp_plot(x, at = 1, digits = 3, ...)
```

**Arguments**

x	An EpiModel object of class dcm, icm, or netsim.
at	Time step for model statistics.
digits	Number of significant digits to print.
...	Additional arguments passed to plot (not currently used).
run	Model run number, for dcm class models with multiple runs (sensitivity analyses).

**Details**

The comp\_plot function provides a visual summary of an epidemic model at a specific time step. The information contained in comp\_plot is the same as in the summary functions for a model, but presented graphically as a compartment flow diagram.

For dcm class plots, specify the model run number if the model contains multiple runs, as in a sensitivity analysis. For icm and netsim class plots, the run argument is not used; the plots show the means and standard deviations across simulations at the specified time step.

These plots are currently limited to one-group and one-mode models for each of the three model classes. That functionality may be expanded in future software releases.

## Examples

```
## Example 1: DCM SIR model with varying act.rate
param <- param.dcm(inf.prob = 0.2, act.rate = 5:7,
                  rec.rate = 1/3, a.rate = 1/90, ds.rate = 1/100,
                  di.rate = 1/35, dr.rate = 1/100)
init <- init.dcm(s.num = 1000, i.num = 1, r.num = 0)
control <- control.dcm(type = "SIR", nsteps = 25, verbose = FALSE)
mod1 <- dcm(param, init, control)
comp_plot(mod1, at = 25, run = 3)

## Example 2: ICM SIR model with 3 simulations
param <- param.icm(inf.prob = 0.2, act.rate = 3, rec.rate = 1/50,
                  a.rate = 1/100, ds.rate = 1/100,
                  di.rate = 1/90, dr.rate = 1/100)
init <- init.icm(s.num = 500, i.num = 1, r.num = 0)
control <- control.icm(type = "SIR", nsteps = 25,
                      nsims = 3, verbose = FALSE)
mod2 <- icm(param, init, control)
comp_plot(mod2, at = 25, digits = 1)
```

control.dcm

*Control Settings for Deterministic Compartmental Models*

## Description

Sets the controls for deterministic compartmental models simulated with [dcm](#).

## Usage

```
control.dcm(
  type,
  nsteps,
  dt = 1,
  odemethod = "rk4",
  dede = FALSE,
  new.mod = NULL,
  sens.param = TRUE,
  print.mod = FALSE,
  verbose = FALSE,
  ...
)
```

## Arguments

type	Disease type to be modeled, with the choice of "SI" for Susceptible-Infected diseases, "SIR" for Susceptible-Infected-Recovered diseases, and "SIS" for Susceptible-Infected-Susceptible diseases.
------	--

nsteps	Number of time steps to solve the model over or vector of times to solve the model over. If the number of time steps, then this must be a positive integer of length 1.
dt	Time unit for model solutions, with the default of 1. Model solutions for fractional time steps may be obtained by setting this to a number between 0 and 1.
odemethod	Ordinary differential equation (ODE) integration method, with the default of the "Runge-Kutta 4" method (see <a href="#">ode</a> for other options).
dede	If TRUE, use the delayed differential equation solver, which allows for time-lagged variables.
new.mod	If not running an base model type, a function with a new model to be simulated (see details).
sens.param	If TRUE, evaluate arguments in parameters with length greater than 1 as sensitivity analyses, with one model run per value of the parameter. If FALSE, one model will be run with parameters of arbitrary length.
print.mod	If TRUE, print the model form to the console.
verbose	If TRUE, print model progress to the console.
...	additional control settings passed to model.

### Details

`control.dcm` sets the required control settings for any deterministic compartmental models solved with the `dcm` function. Controls are required for both base model types and original models. For an overview of control settings for base DCM class models, consult the [Basic DCMs](#) tutorial. For all base models, the `type` argument is a necessary parameter and it has no default.

### New Model Functions

The form of the model function for base models may be displayed with the `print.mod` argument set to TRUE. In this case, the model will not be run. These model forms may be used as templates to write original model functions.

These new models may be input and solved with `dcm` using the `new.mod` argument, which requires as input a model function. Details and examples are found in the [Solving New DCMs](#) tutorial.

### See Also

Use `param.dcm` to specify model parameters and `init.dcm` to specify the initial conditions. Run the parameterized model with `dcm`.



control.icm

*Control Settings for Stochastic Individual Contact Models***Description**

Sets the controls for stochastic individual contact models simulated with `icm`.

**Usage**

```
control.icm(
  type,
  nsteps,
  nsims = 1,
  rec.rand = TRUE,
  a.rand = TRUE,
  d.rand = TRUE,
  initialize.FUN = initialize.icm,
  infection.FUN = infection.icm,
  recovery.FUN = recovery.icm,
  departures.FUN = departures.icm,
  arrivals.FUN = arrivals.icm,
  get_prev.FUN = get_prev.icm,
  verbose = FALSE,
  verbose.int = 0,
  skip.check = FALSE,
  ...
)
```

**Arguments**

type	Disease type to be modeled, with the choice of "SI" for Susceptible-Infected diseases, "SIR" for Susceptible-Infected-Recovered diseases, and "SIS" for Susceptible-Infected-Susceptible diseases.
nsteps	Number of time steps to solve the model over. This must be a positive integer.
nsims	Number of simulations to run.
rec.rand	If TRUE, use a stochastic recovery model, with the number of recovered at each time step a function of random draws from a binomial distribution with the probability equal to <code>rec.rate</code> . If FALSE, then a deterministic rounded count of the expectation implied by that rate.
a.rand	If TRUE, use a stochastic arrival model, with the number of arrivals at each time step a function of random draws from a binomial distribution with the probability equal to the governing arrival rates. If FALSE, then a deterministic rounded count of the expectation implied by those rates.
d.rand	If TRUE, use a stochastic departure model, with the number of departures at each time step a function of random draws from a binomial distribution with the probability equal to the governing departure rates. If FALSE, then a deterministic rounded count of the expectation implied by those rates.

initialize.FUN	Module to initialize the model at the outset, with the default function of <a href="#">initialize.icm</a> .
infection.FUN	Module to simulate disease infection, with the default function of <a href="#">infection.icm</a> .
recovery.FUN	Module to simulate disease recovery, with the default function of <a href="#">recovery.icm</a> .
departures.FUN	Module to simulate departures or exits, with the default function of <a href="#">departures.icm</a> .
arrivals.FUN	Module to simulate arrivals or entries, with the default function of <a href="#">arrivals.icm</a> .
get_prev.FUN	Module to calculate disease prevalence at each time step, with the default function of <a href="#">get_prev.icm</a> .
verbose	If TRUE, print model progress to the console.
verbose.int	Time step interval for printing progress to console, where 0 (the default) prints completion status of entire simulation and positive integer x prints progress after each x time steps.
skip.check	If TRUE, skips the default error checking for the structure and consistency of the parameter values, initial conditions, and control settings before running base epidemic models. Setting this to FALSE is recommended when running models with new modules specified.
...	Additional control settings passed to model.

## Details

`control.icm` sets the required control settings for any stochastic individual contact model solved with the [icm](#) function. Controls are required for both base model types and when passing original process modules. For an overview of control settings for base ICM class models, consult the [Basic ICMs](#) tutorial. For all base models, the `type` argument is a necessary parameter and it has no default.

## New Modules

Base ICM models use a set of module functions that specify how the individual agents in the population are subjected to infection, recovery, demographics, and other processes. Core modules are those listed in the `.FUN` arguments. For each module, there is a default function used in the simulation. The default infection module, for example, is contained in the [infection.icm](#) function.

For original models, one may substitute replacement module functions for any of the default functions. New modules may be added to the workflow at each time step by passing a module function via the `...` argument.

## See Also

Use [param.icm](#) to specify model parameters and [init.icm](#) to specify the initial conditions. Run the parameterized model with [icm](#).

---

control.net*Control Settings for Stochastic Network Models*

---

**Description**

Sets the controls for stochastic network models simulated with [netsim](#).

**Usage**

```
control.net(  
  type,  
  nsteps,  
  start = 1,  
  nsims = 1,  
  ncores = 1,  
  depend,  
  rec.rand = TRUE,  
  a.rand = TRUE,  
  d.rand = TRUE,  
  tea.status = TRUE,  
  attr.rules,  
  epi.by,  
  use.pids = TRUE,  
  pid.prefix,  
  initialize.FUN = initialize.net,  
  departures.FUN = departures.net,  
  arrivals.FUN = arrivals.net,  
  recovery.FUN = recovery.net,  
  edges_correct.FUN = edges_correct,  
  resim_nets.FUN = resim_nets,  
  infection.FUN = infection.net,  
  get_prev.FUN = get_prev.net,  
  verbose.FUN = verbose.net,  
  module.order = NULL,  
  set.control.ergm,  
  set.control.stergm,  
  save.nwstats = TRUE,  
  nwstats.formula = "formation",  
  delete.nodes = FALSE,  
  save.transmat = TRUE,  
  save.network = TRUE,  
  save.other,  
  verbose = TRUE,  
  verbose.int = 1,  
  skip.check = FALSE,  
  ...  
)
```

**Arguments**

type	Disease type to be modeled, with the choice of "SI" for Susceptible-Infected diseases, "SIR" for Susceptible-Infected-Recovered diseases, and "SIS" for Susceptible-Infected-Susceptible diseases.
nsteps	Number of time steps to simulate the model over. This must be a positive integer that is equal to the final step of a simulation. If simulation is restarted with start argument, this number must be at least one greater than that argument's value.
start	For dependent simulations, time point to start up simulation. For restarted simulations, this must be one greater than the final time step in the prior simulation and must be less than the value in nsteps.
nsims	The total number of disease simulations.
ncores	Number of processor cores to run multiple simulations on, using the foreach and doParallel implementations.
depend	If TRUE, resimulate the network at each time step. This occurs by default with two varieties of dependent models: if there are any vital dynamic parameters in the model (or if non-standard arrival or departures modules are passed into control.net), or if the network model formation formula includes the "status" attribute.
rec.rand	If TRUE, use a stochastic recovery model, with the number of recovered at each time step a function of random draws from a binomial distribution with the probability equal to rec.rate. If FALSE, then a deterministic rounded count of the expectation implied by that rate.
a.rand	If TRUE, use a stochastic arrival model, with the number of arrivals at each time step a function of random draws from a binomial distribution with the probability equal to the governing arrival rates. If FALSE, then a deterministic rounded count of the expectation implied by those rates.
d.rand	If TRUE, use a stochastic departure model, with the number of departures at each time step a function of random draws from a binomial distribution with the probability equal to the governing departure rates. If FALSE, then a deterministic rounded count of the expectation implied by those rates.
tea.status	If TRUE, use a temporally extended attribute (TEA) to store disease status. A TEA is needed for plotting static networks at different time steps and for animating dynamic networks with evolving status. TEAs are computationally inefficient for large simulations and should be toggled off in those cases. This argument automatically set to FALSE if delete.nodes=TRUE.
attr.rules	A list containing the rules for setting the attributes of incoming nodes, with one list element per attribute to be set (see details below).
epi.by	A character vector of length 1 containing a nodal attribute for which subgroup epidemic prevalences should be calculated. This nodal attribute must be contained in the network model formation formula, otherwise it is ignored.
use.pids	If TRUE, use persistent ids for vertices; otherwise, numeric ids will be recycled in models with vital dynamics. For one-mode simulations, this will be a random hexadecimal value; for bipartite simulations, it will be based on pid.prefix.
pid.prefix	For bipartite network simulations with vital dynamics, a character vector of length 2 containing the prefixes, with the default of c("F", "M").

initialize.FUN	Module to initialize the model at time 1, with the default function of <a href="#">initialize.net</a> .
departures.FUN	Module to simulate departure or exit, with the default function of <a href="#">departures.net</a> .
arrivals.FUN	Module to simulate arrivals or entries, with the default function of <a href="#">arrivals.net</a> .
recovery.FUN	Module to simulate disease recovery, with the default function of <a href="#">recovery.net</a> .
edges_correct.FUN	Module to adjust the edges coefficient in response to changes to the population size, with the default function of <a href="#">edges_correct</a> that preserves mean degree.
resim_nets.FUN	Module to resimulate the network at each time step, with the default function of <a href="#">resim_nets</a> .
infection.FUN	Module to simulate disease infection, with the default function of <a href="#">infection.net</a> .
get_prev.FUN	Module to calculate disease prevalence at each time step, with the default function of <a href="#">get_prev.net</a> .
verbose.FUN	Module to print simulation progress to screen, with the default function of <a href="#">verbose.net</a> .
module.order	A character vector of module names that lists modules the order in which they should be evaluated within each time step. If NULL, the modules will be evaluated as follows: first any new modules supplied through ... in the order in which they are listed, then the built-in modules in their order of the function listing. The initialize.FUN will always be run first and the verbose.FUN always last.
set.control.ergm	Control arguments passed to simulate.ergm. See the help file for <a href="#">netdx</a> for details and examples on specifying this parameter.
set.control.stergm	Control arguments passed to simulate.stergm. See the help file for <a href="#">netdx</a> for details and examples on specifying this parameter.
save.nwstats	If TRUE, save network statistics in a data frame. The statistics to be saved are specified in the nwstats.formula argument.
nwstats.formula	A right-hand sided ERGM formula that includes network statistics of interest, with the default to the formation formula terms.
delete.nodes	If TRUE, delete inactive nodes from the network after each time step, otherwise deactivate them but keep them in the network object. Deleting nodes increases computational efficiency in large network simulations.
save.transmat	If TRUE, save a transmission matrix for each simulation. This object contains one row for each transmission event (see <a href="#">discord_edgelist</a> ).
save.network	If TRUE, save a networkDynamic object containing full edge history for each simulation. If delete.nodes is set to TRUE, this will only contain a static network with the edge configuration at the final time step of each simulation.
save.other	A vector of elements on the dat master data list to save out after each simulation. One example for base models is the attribute list, "attr", at the final time step.
verbose	If TRUE, print model progress to the console.
verbose.int	Time step interval for printing progress to console, where 0 prints completion status of entire simulation and positive integer x prints progress after each x time steps. The default is to print progress after each time step.

skip.check	If TRUE, skips the default error checking for the structure and consistency of the parameter values, initial conditions, and control settings before running base epidemic models. Setting this to FALSE is recommended when running models with new modules specified.
...	Additional control settings passed to model.

## Details

control.net sets the required control settings for any network model solved with the [netsim](#) function. Controls are required for both base model types and when passing original process modules. For an overview of control settings for base models, consult the [Basic Network Models](#) tutorial. For all base models, the type argument is a necessary parameter and it has no default.

## The attr.rules Argument

The attr.rules parameter is used to specify the rules for how nodal attribute values for incoming nodes should be set. These rules are only necessary for models in which there are incoming nodes (i.e., arrivals) and there is also a nodal attribute in the network model formation formula set in [netest](#). There are three rules available for each attribute value:

- **"current"**: new nodes will be assigned this attribute in proportion to the distribution of that attribute among existing nodes at that current time step.
- **"t1"**: new nodes will be assigned this attribute in proportion to the distribution of that attribute among nodes at time 1 (that is, the proportions set in the original network for [netest](#)).
- **<Value>**: all new nodes will be assigned this specific value, with no variation.

For example, the rules list `attr.rules = list(race = "t1", sex = "current", status = "s")` specifies how the race, sex, and status attributes should be set for incoming nodes. By default, the rule is "current" for all attributes except status, in which case it is "s" (that is, all incoming nodes are susceptible).

## New Modules

Base network models use a set of module functions that specify how the individual nodes in the network are subjected to infection, recovery, demographics, and other processes. Core modules are those listed in the .FUN arguments. For each module, there is a default function used in the simulation. The default infection module, for example, is contained in the [infection.net](#) function.

For original models, one may substitute replacement module functions for any of the default functions. New modules may be added to the workflow at each time step by passing a module function via the ... argument. Consult the [New Network Models](#) tutorial. One may remove existing modules, such as `arrivals.FUN`, from the workflow by setting the parameter value for that argument to NULL.

## See Also

Use [param.net](#) to specify model parameters and [init.net](#) to specify the initial conditions. Run the parameterized model with [netsim](#).

## Description

Solves deterministic compartmental epidemic models for infectious disease.

## Usage

```
dcm(param, init, control)
```

## Arguments

param	Model parameters, as an object of class <code>param.dcm</code> .
init	Initial conditions, as an object of class <code>init.dcm</code> .
control	Control settings, as an object of class <code>control.dcm</code> .

## Details

The `dcm` function uses the ordinary differential equation solver in the `deSolve` package to model disease as a deterministic compartmental system. The parameterization for these models follows the standard approach in `EpiModel`, with epidemic parameters, initial conditions, and control settings. A description of solving DCMs with the `dcm` function may be found in the [Basic DCMs](#) tutorial.

The `dcm` function performs modeling of both base model types and original models with new structures. Base model types include one-group and two-group models with disease types for Susceptible-Infected (SI), Susceptible-Infected-Recovered (SIR), and Susceptible-Infected-Susceptible (SIS). New model types may be written and input into `dcm` following the steps outlined in the [Solving New DCMs](#) tutorial. Both base and original models require the `param`, `init`, and `control` inputs.

## Value

A list of class `dcm` with the following elements:

- **param:** the epidemic parameters passed into the model through `param`, with additional parameters added as necessary.
- **control:** the control settings passed into the model through `control`, with additional controls added as necessary.
- **epi:** a list of data frames, one for each epidemiological output from the model. Outputs for base models always include the size of each compartment, as well as flows in, out of, and between compartments.

## References

Soetaert K, Petzoldt T, Setzer W. Solving Differential Equations in R: Package `deSolve`. *Journal of Statistical Software*. 2010; 33(9): 1-25. <http://www.jstatsoft.org/v33/i09/>.

## See Also

Extract the model results with `as.data.frame.dcm`. Summarize the time-specific model results with `summary.dcm`. Plot the model results with `plot.dcm`. Plot a compartment flow diagram with `comp_plot`.

## Examples

```
## Example 1: SI Model (One-Group)
# Set parameters
param <- param.dcm(inf.prob = 0.2, act.rate = 0.25)
init <- init.dcm(s.num = 500, i.num = 1)
control <- control.dcm(type = "SI", nsteps = 500)
mod1 <- dcm(param, init, control)
mod1
plot(mod1)

## Example 2: SIR Model with Vital Dynamics (One-Group)
param <- param.dcm(inf.prob = 0.2, act.rate = 5,
                   rec.rate = 1/3, a.rate = 1/90, ds.rate = 1/100,
                   di.rate = 1/35, dr.rate = 1/100)
init <- init.dcm(s.num = 500, i.num = 1, r.num = 0)
control <- control.dcm(type = "SIR", nsteps = 500)
mod2 <- dcm(param, init, control)
mod2
plot(mod2)

## Example 3: SIS Model with act.rate Sensitivity Parameter
param <- param.dcm(inf.prob = 0.2, act.rate = seq(0.1, 0.5, 0.1),
                   rec.rate = 1/50)
init <- init.dcm(s.num = 500, i.num = 1)
control <- control.dcm(type = "SIS", nsteps = 500)
mod3 <- dcm(param, init, control)
mod3
plot(mod3)

## Example 4: SI Model with Vital Dynamics (Two-Group)
param <- param.dcm(inf.prob = 0.4, inf.prob.g2 = 0.1,
                   act.rate = 0.25, balance = "g1",
                   a.rate = 1/100, a.rate.g2 = NA,
                   ds.rate = 1/100, ds.rate.g2 = 1/100,
                   di.rate = 1/50, di.rate.g2 = 1/50)
init <- init.dcm(s.num = 500, i.num = 1,
                 s.num.g2 = 500, i.num.g2 = 0)
control <- control.dcm(type = "SI", nsteps = 500)
mod4 <- dcm(param, init, control)
mod4
plot(mod4)
```



---

dissolution\_coefs      *Dissolution Coefficients for Stochastic Network Models*


---

## Description

Calculates dissolution coefficients, given a dissolution model and average edge duration, to pass as offsets to an ERGM/STERGM model fit in `netest`.

## Usage

```
dissolution_coefs(dissolution, duration, d.rate = 0)
```

## Arguments

dissolution	Right-hand sided STERGM dissolution formula (see <a href="#">netest</a> ). See below for list of supported dissolution models.
duration	A vector of mean edge durations in arbitrary time units.
d.rate	Departure or exit rate from the population, as a single homogenous rate that applies to the entire population.

## Details

This function performs two calculations for dissolution coefficients used in a network model estimated with [netest](#):

1. **Transformation:** the mean duration of edges in a network are mathematically transformed to logit coefficients.
2. **Adjustment:** in a dynamic network simulation in an open population (in which there are departures), it is further necessary to adjust these coefficients for dynamic simulations; this upward adjustment accounts for departure as a competing risk to edge dissolution.

The current dissolution models supported by this function and in network model estimation in [netest](#) are as follows:

- `~offset(edges)`: a homogeneous dissolution model in which the edge duration is the same for all partnerships. This requires specifying one duration value.
- `~offset(edges) + offset(nodematch("<attr>"))`: a heterogeneous model in which the edge duration varies by whether the nodes in the dyad have similar values of a specified attribute. The duration vector should now contain two values: the first is the mean edge duration of non-matched dyads, and the second is the duration of the matched dyads.
- `~offset(edges) + offset(nodemix("<attr>"))`: a heterogenous model that extends the `nodematch` model to include non-binary attributes for homophily. The duration vector should first contain the base value, then the values for every other possible combination in the term.
- `~offset(edges) + offset(nodefactor("<attr>"))`: a heterogenous model in which the edge duration varies by a specified attribute. The duration vector should first contain the base value, then the values for every other value of that attribute in the term.

**Value**

A list of class `dissscoef` with the following elements:

- **dissolution:** right-hand sided STERGM dissolution formula passed in the function call.
- **duration:** mean edge durations passed into the function.
- **coef.crude:** mean durations transformed into logit coefficients.
- **coef.adj:** crude coefficients adjusted for the risk of departure on edge persistence, if the `d.rate` argument is supplied.
- **d.rate:** the departure rate.

**See Also**

The theory and details of this function are explained in detail in the [EpiModel Network Utility Functions](#) tutorial.

**Examples**

```
## Homogeneous dissolution model with no departures
dissolution_coefs(dissolution = ~offset(edges), duration = 25)

## Homogeneous dissolution model with departures
dissolution_coefs(dissolution = ~offset(edges), duration = 25,
                  d.rate = 0.001)

## Heterogeneous dissolution model in which same-race edges have
## shorter duration compared to mixed-race edges, with no departures
dissolution_coefs(dissolution = ~offset(edges) + offset(nodematch("race")),
                  duration = c(20, 10))

## Heterogeneous dissolution model in which same-race edges have
## shorter duration compared to mixed-race edges, with departures
dissolution_coefs(dissolution = ~offset(edges) + offset(nodematch("race")),
                  duration = c(20, 10), d.rate = 0.001)

## Not run:
## Extended example for differential homophily by age group
# Set up the network with nodes categorized into 5 age groups
nw <- network.initialize(1000, directed = FALSE)
age.grp <- sample(1:5, 1000, TRUE)
nw <- set.vertex.attribute(nw, "age.grp", age.grp)

# durations = non-matched, age.grp1 & age.grp1, age.grp2 & age.grp2, ...
# TERGM will include differential homophily by age group with nodematch term
# Target stats for the formation model are overall edges, and then the number
#   matched within age.grp 1, age.grp 2, ..., age.grp 5
form <- ~edges + nodematch("age.grp", diff = TRUE)
target.stats <- c(450, 100, 125, 40, 80, 100)

# Target stats for the dissolution model are duration of non-matched edges, then
#   duration of edges matched within age.grp 1, age.grp 2, ..., age.grp 5
```

```

durs <- c(60, 30, 80, 100, 125, 160)
diss <- dissolution_coefs(~offset(edges) +
                          offset(nodematch("age.grp", diff = TRUE)),
                          duration = durs)

# Fit the TERGM
fit <- netest(nw, form, target.stats, diss)

# Full diagnostics to evaluate model fit
dx <- netdx(fit, nsims = 10, ncores = 4, nsteps = 300)
print(dx)

# Simulate one long time series to examine timed edgelist
dx <- netdx(fit, nsims = 1, nsteps = 5000, keep.tedgelist = TRUE)

# Extract timed-edgelist
te <- as.data.frame(dx)
head(te)

# Limit to non-censored edges
te <- te[which(te$onset.censored == FALSE & te$terminus.censored == FALSE),
        c("head", "tail", "duration")]
head(te)

# Look up the age group of head and tail nodes
te$ag.head <- age.grp[te$head]
te$ag.tail <- age.grp[te$tail]
head(te)

# Recover average edge durations for age-group pairing
mean(te$duration[te$ag.head != te$ag.tail])
mean(te$duration[te$ag.head == 1 & te$ag.tail == 1])
mean(te$duration[te$ag.head == 2 & te$ag.tail == 2])
mean(te$duration[te$ag.head == 3 & te$ag.tail == 3])
mean(te$duration[te$ag.head == 4 & te$ag.tail == 4])
mean(te$duration[te$ag.head == 5 & te$ag.tail == 5])
durs

## End(Not run)

```

---

edgelist\_censor

---

*Table of Edge Censoring*


---

## Description

Outputs a table of the number and percent of edges that are left-censored, right-censored, both-censored, or uncensored for a `networkDynamic` object.

**Usage**

```
edgelist_censor(el)
```

**Arguments**

`el` Timed edgelist with start and end times extracted from a `networkDynamic` object using the `as.data.frame.networkDynamic` function.

**Details**

Given a STERGM simulation over a specified number of time steps, the edges within that simulation may be left-censored (started before the first step), right-censored (continued after the last step), right and left-censored, or uncensored. The amount of censoring will increase when the average edge duration approaches the length of the simulation.

**Examples**

```
# Initialize and parameterize network model
nw <- network.initialize(n = 100, directed = FALSE)
formation <- ~edges
target.stats <- 50
coef.diss <- dissolution_coefs(dissolution = ~offset(edges), duration = 20)

# Model estimation
est <- netest(nw, formation, target.stats, coef.diss, verbose = FALSE)

# Simulate the network and extract a timed edgelist
dx <- netdx(est, nsims = 1, nsteps = 100, keep.tedgelist = TRUE, verbose = FALSE)
el <- as.data.frame(dx)

# Calculate censoring
edgelist_censor(el)
```

**Description**

Runs a web browser-based GUI of deterministic compartmental models, stochastic individual contact models, and basic network models.

**Usage**

```
epiweb(class, ...)
```

**Arguments**

`class`            Model class, with options of "dcm", "icm" and "net".  
`...`            Additional arguments passed to `shiny::runApp`.

**Details**

epiweb runs a web-based GUI of one-group deterministic compartmental models, stochastic individual contact models, and stochastic network models with user input on model type, state sizes, and parameters. Model output may be plotted, summarized, and saved as raw data using the core EpiModel functionality for these model classes. These applications are built using the shiny package framework.

**References**

RStudio. shiny: Web Application Framework for R. R package version 1.0.5. 2015. <http://www.rstudio.com/shiny/>

**See Also**

[dcm](#), [icm](#), [netsim](#)

**Examples**

```
## Not run:
## Deterministic compartmental models
epiweb(class = "dcm")

## Stochastic individual contact models
epiweb(class = "icm")

## Stochastic network models
epiweb(class = "net")

## End(Not run)
```

---

geom\_bands

*ggplot2 geom for Quantile Bands*


---

**Description**

Plots quantile bands given a data.frame with stochastic model results from `icm` or `netsim`.

**Usage**

```
geom_bands(mapping, lower = 0.25, upper = 0.75, alpha = 0.25, ...)
```

**Arguments**

mapping	standard aesthetic mapping <code>aes()</code> input for <code>ggplot2</code> .
lower	Lower quantile for the time series.
upper	Upper quantile for the time series.
alpha	Transparency of the ribbon fill.
...	Additional arguments passed to <code>stat_summary</code> .

**Details**

This is a wrapper around `ggplot::stat_summary` with a ribbon geom as aesthetic output.

**Examples**

```
param <- param.icm(inf.prob = 0.2, act.rate = 0.25)
init <- init.icm(s.num = 500, i.num = 1)
control <- control.icm(type = "SI", nsteps = 250, nsims = 5)
mod1 <- icm(param, init, control)
df <- as.data.frame(mod1)
df.mean <- as.data.frame(mod1, out = "mean")

library(ggplot2)
ggplot() +
  geom_line(data = df, mapping = aes(time, i.num, group = sim), alpha = 0.25,
            lwd = 0.25, color = "firebrick") +
  geom_bands(data = df, mapping = aes(time, i.num),
            lower = 0.1, upper = 0.9, fill = "firebrick") +
  geom_line(data = df.mean, mapping = aes(time, i.num)) +
  theme_minimal()
```

---

get\_args

---

*Get Arguments from EpiModel Parameterization Functions*


---

**Description**

Returns a list of argument names and values for use for parameter processing functions.

**Usage**

```
get_args(formal.args, dot.args)
```

**Arguments**

formal.args	The output of <code>formals(sys.function())</code> .
dot.args	The output of <code>list(...)</code> .

get\_degree

*Get Individual Degree from Network or Edgelist***Description**

A fast method for querying the current degree of all individuals within a network.

**Usage**

```
get_degree(x)
```

**Arguments**

**x** Either an object of class `network` or `edgelist` generated from a network. If `x` is an `edgelist`, then it must contain an attribute for the total network size, `n`.

**Details**

Individual-level data on the current degree of nodes within a network is often useful for summary statistics and modeling complex interactions between degree. Given a `network` class object, `net`, one way to look up the current degree is to get a summary of the ERGM term, `sociality`, as in: `summary(net ~ sociality(base = 0))`. But that is computationally inefficient for a number of reasons. This function provide a fast method for generating the vector of degree using a query of the `edgelist`. It is even faster if the parameter `x` is already transformed as an `edgelist`.

**Examples**

```
nw <- network.initialize(500, directed = FALSE)

set.seed(1)
fit <- ergm(nw ~ edges, target.stats = 250)
sim <- simulate(fit)

# Slow ERGM-based method
ergm.method <- unname(summary(sim ~ sociality(base = 0)))
ergm.method

# Fast tabulate method with network object
deg.net <- get_degree(sim)
deg.net

# Even faster if network already transformed into an edgelist
el <- as.edgelist(sim)
deg.el <- get_degree(el)
deg.el

identical(as.integer(ergm.method), deg.net, deg.el)
```

---

get\_formula\_term\_attr *Outputs ERGM Formula Attributes into a Character Vector*

---

### Description

Given a formation formula for a network model, outputs it into a character vector of vertex attributes to be used in netsim simulations.

### Usage

```
get_formula_term_attr(form, nw)
```

### Arguments

form	an ergm model formula
nw	a network object

---

get_network	<i>Extract networkDynamic and network Objects from Network Simulations</i>
-------------	--

---

### Description

Extracts the networkDynamic object from a either a network epidemic model object generated with netsim or a network diagnostic simulation generated with netdx, with the option to collapse the extracted networkDynamic object down to a static network object.

### Usage

```
get_network(x, sim = 1, network = 1, collapse = FALSE, at)
```

### Arguments

x	An EpiModel object of class <a href="#">netsim</a> or <a href="#">netdx</a> .
sim	Simulation number of extracted network.
network	Network number, for netsim objects with multiple overlapping networks (advanced use, and not applicable to netdx objects).
collapse	If TRUE, collapse the networkDynamic object to a static network object at a specified time step.
at	If collapse is TRUE, the time step at which the extracted network should be collapsed.



## Details

This function requires that the networkDynamic is saved during the network simulation while running either `netsim` or `netdx`. For the former, that is specified with the `save.network` parameter in `control.net`. For the latter, that is specified with the `keep.tedgelist` parameter directly in `netdx`.

## Examples

```
# Set up network and TERGM formula
nw <- network.initialize(n = 100, bipartite = 50, directed = FALSE)
formation <- ~edges
target.stats <- 50
coef.diss <- dissolution_coefs(dissolution = ~offset(edges), duration = 20)

# Estimate the model
est <- netest(nw, formation, target.stats, coef.diss)

# Run diagnostics, saving the networkDynamic objects
dx <- netdx(est, nsteps = 10, nsims = 3, keep.tnetwork = TRUE, verbose = FALSE)

# Extract the network for simulation 2 from dx object
get_network(dx, sim = 2)

# Extract and collapse the network from simulation 1 at time step 5
get_network(dx, collapse = TRUE, at = 5)

# Parameterize the epidemic model, and simulate it
param <- param.net(inf.prob = 0.3, inf.prob.m2 = 0.15)
init <- init.net(i.num = 10, i.num.m2 = 10)
control <- control.net(type = "SI", nsteps = 10, nsims = 3, verbose = FALSE)
mod <- netsim(est, param, init, control)

# Extract the network for simulation 2 from mod object
get_network(mod, sim = 2)

## Extract and collapse the network from simulation 1 at time step 5
get_network(mod, collapse = TRUE, at = 5)
```

---

get\_nwstats

---

*Extract Network Statistics from netsim or netdx Object*


---

## Description

Extracts a data frame of network statistics from a network epidemic model simulated with `netsim` or a network diagnostics object simulated with `netdx`.

## Usage

```
get_nwstats(x, sim, network = 1)
```

**Arguments**

x	An EpiModel object of class <code>netsim</code> or <code>netdx</code> .
sim	A vector of simulation numbers from the extracted object
network	Network number, for <code>netsim</code> objects with multiple overlapping networks (advanced use, and not applicable to <code>netdx</code> objects).

**Examples**

```
# Bipartite Bernoulli random graph TERGM
nw <- network.initialize(n = 100, bipartite = 50, directed = FALSE)
formation <- ~edges
target.stats <- 50
coef.diss <- dissolution_coefs(dissolution = ~offset(edges), duration = 20)
est <- netest(nw, formation, target.stats, coef.diss, verbose = FALSE)

dx <- netdx(est, nsim = 3, nsteps = 10, verbose = FALSE,
            nwstats.formula = ~edges + isolates)
get_nwstats(dx)
get_nwstats(dx, sim = 1)

# SI epidemic model
param <- param.net(inf.prob = 0.3, inf.prob.m2 = 0.15)
init <- init.net(i.num = 10, i.num.m2 = 10)
control <- control.net(type = "SI", nsteps = 10, nsims = 3,
                      nwstats.formula = ~edges + meandeg + degree(0:5),
                      verbose = FALSE)
mod <- netsim(est, param, init, control)

# Extract the network statistics from all or sets of simulations
get_nwstats(mod)
get_nwstats(mod, sim = 2)
get_nwstats(mod, sim = c(1, 3))

# On the fly summary stats
summary(get_nwstats(mod))
colMeans(get_nwstats(mod))
```

get\_sims

*Extract Network Simulations***Description**

Subsets the entire `netsim` object to a subset of simulations, essentially functioning like a reverse of `merge`.

**Usage**

```
get_sims(x, sims, var)
```

**Arguments**

<code>x</code>	An object of class <code>netsim</code> .
<code>sims</code>	A numeric vector of simulation numbers to retain in the output object, or "mean" which selects the one simulation with the value of the variable specified in <code>var</code> closest to the mean of <code>var</code> across all simulations.
<code>var</code>	A character vector of variables to retain from <code>x</code> if <code>sims</code> is a numeric vector, or a single variable name for selecting the average simulation from the set if <code>sims</code> = "mean".

**Examples**

```
# Network model estimation
nw <- network.initialize(n = 100, directed = FALSE)
formation <- ~edges
target.stats <- 50
coef.diss <- dissolution_coefs(dissolution = ~offset(edges), duration = 20)
est1 <- netest(nw, formation, target.stats, coef.diss, verbose = FALSE)

# Epidemic model
param <- param.net(inf.prob = 0.3)
init <- init.net(i.num = 10)
control <- control.net(type = "SI", nsteps = 10, nsims = 3, verbose.int = 0)
mod1 <- netsim(est1, param, init, control)

# Get sim 2
sim2 <- get_sims(mod1, sims = 2)

# Get sims 2 and 3 and keep only a subset of variables
sim2.small <- get_sims(mod1, sims = 2:3, var = c("i.num", "si.flow"))

# Extract the mean simulation for the variable i.num
sim.mean <- get_sims(mod1, sims = "mean", var = "i.num")
```

icm

*Stochastic Individual Contact Models***Description**

Simulates stochastic individual contact epidemic models for infectious disease.

**Usage**

```
icm(param, init, control)
```

**Arguments**

<code>param</code>	Model parameters, as an object of class <code>param.icm</code> .
<code>init</code>	Initial conditions, as an object of class <code>init.icm</code> .
<code>control</code>	Control settings, as an object of class <code>control.icm</code> .

## Details

Individual contact models are intended to be the stochastic microsimulation analogs to deterministic compartmental models. ICMs simulate disease spread on individual agents in discrete time as a function of processes with stochastic variation. The stochasticity is inherent in all transition processes: infection, recovery, and demographics. A detailed description of these models may be found in the [Basic ICMs](#) tutorial.

The `icm` function performs modeling of both the base model types and original models. Base model types include one-group and two-group models with disease types for Susceptible-Infected (SI), Susceptible-Infected-Recovered (SIR), and Susceptible-Infected-Susceptible (SIS). Original models may be built by writing new process modules that either take the place of existing modules (for example, disease recovery), or supplement the set of existing processes with a new one contained in an original module.

## Value

A list of class `icm` with the following elements:

- **param:** the epidemic parameters passed into the model through `param`, with additional parameters added as necessary.
- **control:** the control settings passed into the model through `control`, with additional controls added as necessary.
- **epi:** a list of data frames, one for each epidemiological output from the model. Outputs for base models always include the size of each compartment, as well as flows in, out of, and between compartments.

## See Also

Extract the model results with [as.data.frame.icm](#). Summarize the time-specific model results with [summary.icm](#). Plot the model results with [plot.icm](#). Plot a compartment flow diagram with [comp\\_plot](#).

## Examples

```
## Not run:
## Example 1: SI Model
param <- param.icm(inf.prob = 0.2, act.rate = 0.25)
init <- init.icm(s.num = 500, i.num = 1)
control <- control.icm(type = "SI", nsteps = 500, nsims = 10)
mod1 <- icm(param, init, control)
mod1
plot(mod1)

## Example 2: SIR Model
param <- param.icm(inf.prob = 0.2, act.rate = 0.25, rec.rate = 1/50)
init <- init.icm(s.num = 500, i.num = 1, r.num = 0)
control <- control.icm(type = "SIR", nsteps = 500, nsims = 10)
mod2 <- icm(param, init, control)
mod2
plot(mod2)
```

```
## Example 3: SIS Model
param <- param.icm(inf.prob = 0.2, act.rate = 0.25, rec.rate = 1/50)
init <- init.icm(s.num = 500, i.num = 1)
control <- control.icm(type = "SIS", nsteps = 500, nsims = 10)
mod3 <- icm(param, init, control)
mod3
plot(mod3)

## Example 4: SI Model with Vital Dynamics (Two-Group)
param <- param.icm(inf.prob = 0.4, inf.prob.g2 = 0.1,
  act.rate = 0.25, balance = "g1",
  a.rate = 1/100, a.rate.g2 = NA,
  ds.rate = 1/100, ds.rate.g2 = 1/100,
  di.rate = 1/50, di.rate.g2 = 1/50)
init <- init.icm(s.num = 500, i.num = 1,
  s.num.g2 = 500, i.num.g2 = 0)
control <- control.icm(type = "SI", nsteps = 500, nsims = 10)
mod4 <- icm(param, init, control)
mod4
plot(mod4)

## End(Not run)
```

init.dcm

*Initial Conditions for Deterministic Compartmental Models***Description**

Sets the initial conditions for deterministic compartmental models simulated with dcm.

**Usage**

```
init.dcm(s.num, i.num, r.num, s.num.g2, i.num.g2, r.num.g2, ...)
```

**Arguments**

s.num	Number of initial susceptible. For two-group models, this is the number of initial group 1 susceptible.
i.num	Number of initial infected. For two-group models, this is the number of initial group 1 infected.
r.num	Number of initial recovered. For two-group models, this is the number of initial group 1 recovered. This parameter is only used for the SIR model type.
s.num.g2	Number of initial susceptible in group 2. This parameter is only used for two-group models.
i.num.g2	Number of initial infected in group 2. This parameter is only used for two-group models.

r.num.g2	Number of initial recovered in group 2. This parameter is only used for two-group SIR models.
...	Additional initial conditions passed to model.

### Details

The initial conditions for a model solved with `dcm` should be input into the `init.dcm` function. This function handles initial conditions for both base model types and original models. For an overview of initial conditions for base DCM class models, consult the [Basic DCMs](#) tutorial.

Original models may use the parameter names listed as arguments here, a new set of names, or a combination of both. With new models, initial conditions must be input in the same order that the solved derivatives from the model are output. More details on this requirement are outlined in the [Solving New DCMs](#) tutorial.

### See Also

Use `param.dcm` to specify model parameters and `control.dcm` to specify the control settings. Run the parameterized model with `dcm`.

---

init.icm

---

*Initial Conditions for Stochastic Individual Contact Models*


---

### Description

Sets the initial conditions for stochastic individual contact models simulated with `icm`.

### Usage

```
init.icm(s.num, i.num, r.num, s.num.g2, i.num.g2, r.num.g2, ...)
```

### Arguments

s.num	Number of initial susceptible. For two-group models, this is the number of initial group 1 susceptible.
i.num	Number of initial infected. For two-group models, this is the number of initial group 1 infected.
r.num	Number of initial recovered. For two-group models, this is the number of initial group 1 recovered. This parameter is only used for the SIR model type.
s.num.g2	Number of initial susceptible in group 2. This parameter is only used for two-group models.
i.num.g2	Number of initial infected in group 2. This parameter is only used for two-group models.
r.num.g2	Number of initial recovered in group 2. This parameter is only used for two-group SIR models.
...	Additional initial conditions passed to model.

## Details

The initial conditions for a model solved with `icm` should be input into the `init.icm` function. This function handles initial conditions for both base models and original models using new modules. For an overview of initial conditions for base ICM class models, consult the [Basic ICMs](#) tutorial.

## See Also

Use `param.icm` to specify model parameters and `control.icm` to specify the control settings. Run the parameterized model with `icm`.

---

init.net

---

*Initial Conditions for Stochastic Network Models*


---

## Description

Sets the initial conditions for stochastic network models simulated with `netsim`.

## Usage

```
init.net(i.num, r.num, i.num.m2, r.num.m2, status.vector, infTime.vector, ...)
```

## Arguments

<code>i.num</code>	Number of initial infected. For bipartite models, this is the number of initial mode 1 infected.
<code>r.num</code>	Number of initial recovered. For bipartite models, this is the number of initial mode 1 recovered. This parameter is only used for the SIR model type.
<code>i.num.m2</code>	Number of initial infected in mode 2. This parameter is only used for bipartite models.
<code>r.num.m2</code>	Number of initial recovered in mode 2. This parameter is only used for bipartite SIR models.
<code>status.vector</code>	A vector of length equal to the size of the input network, containing the status of each node. Setting status here overrides any inputs passed in the <code>.num</code> arguments.
<code>infTime.vector</code>	A vector of length equal to the size of the input network, containing the (historical) time of infection for each of those nodes with a current status of "i". Can only be used if <code>status.vector</code> is used, and must contain NA values for any nodes whose status is not "i".
<code>...</code>	Additional initial conditions passed to model.

## Details

The initial conditions for a model solved with `netsim` should be input into the `init.net` function. This function handles initial conditions for both base models and new modules. For an overview of specifying initial conditions across a variety of base network models, consult the [Basic Network Models](#) tutorial.

**See Also**

Use [param.net](#) to specify model parameters and [control.net](#) to specify the control settings. Run the parameterized model with [netsim](#).

**Examples**

```
# Example of using status.vector and infTime.vector together
n <- 100
status <- sample(c("s", "i"), size = n, replace = TRUE, prob = c(0.8, 0.2))
infTime <- rep(NA, n)
infTime[which(status == "i")] <- -rgeom(sum(status == "i"), prob = 0.01) + 2

init.net(status.vector = status, infTime.vector = infTime)
```

---

InitErgmTerm.absdiffby

*Definition for absdiffby ERGM Term*


---

**Description**

This function defines and initialize the absdiffby ERGM term that allows for targeting age homophily by sex.

**Usage**

```
InitErgmTerm.absdiffby(nw, arglist, ...)
```

**Arguments**

nw	An object of class network.
arglist	A list of arguments as specified in the <code>ergm.userterms</code> package framework.
...	Additional data passed into the function as specified in the <code>ergm.userterms</code> package framework.

**Details**

This ERGM user term was written to allow for age-based homophily in partnership formation that is asymmetric by sex. The absdiff component targets age homophily while the by component allows that to be structured by a binary attribute such as "male", in order to enforce an offset in the average difference.

**Author(s)**

Samuel M. Jenness



---

InitErgmTerm.absdiffnodemix

*Definition for absdiffnodemix ERGM Term*


---

## Description

This function defines and initialize the absdiffnodemix ERGM term that allows for targeting age homophily by race.

## Usage

```
InitErgmTerm.absdiffnodemix(nw, arglist, ...)
```

## Arguments

nw	An object of class network.
arglist	A list of arguments as specified in the ergm.userterms package framework.
...	Additional data passed into the function as specified in the ergm.userterms package framework.

## Details

This ERGM user term was written to allow for age-based homophily in partnership formation that is heterogenous by race. The absdiff component allows targets the distribution of age mixing on that continuous variable, and the nodemix component differentiates this for black-black, black-white, and white-white couples.

## Author(s)

Steven M. Goodreau

---

is.transmat

*Extract Transmissions Matrix from Network Epidemic Model*


---

## Description

Extracts the matrix of transmission data for each transmission event that occurred within a network epidemic model.

## Usage

```
is.transmat(x)
```

```
get_transmat(x, sim = 1)
```

**Arguments**

**x**                      An EpiModel object of class `netsim`.  
**sim**                    Simulation number of extracted network.

**Value**

A data frame with the following columns

- **at**: the time step at which the transmission occurred.
- **sus**: the ID number of the susceptible (newly infected) node.
- **inf**: the ID number of the infecting node.
- **infDur**: the duration of the infecting node's disease at the time of the transmission.
- **transProb**: the probability of transmission per act.
- **actRate**: the rate of acts per unit time.
- **finalProb**: the final transmission probability for the transmission event.

**Examples**

```
## Simulate SI epidemic on bipartite Bernoulli random graph
nw <- network.initialize(n = 100, bipartite = 50, directed = FALSE)
formation <- ~edges
target.stats <- 50
coef.diss <- dissolution_coefs(dissolution = ~offset(edges), duration = 20)
est <- netest(nw, formation, target.stats, coef.diss, verbose = FALSE)
param <- param.net(inf.prob = 0.3, inf.prob.m2 = 0.15)
init <- init.net(i.num = 10, i.num.m2 = 10)
control <- control.net(type = "SI", nsteps = 10, nsims = 3, verbose = FALSE)
mod <- netsim(est, param, init, control)

## Extract the transmission matrix from simulation 2
get_transmat(mod, sim = 2)
```

---

merge.icm

---

*Merge Data across Stochastic Individual Contact Model Simulations*


---

**Description**

Merges epidemiological data from two independent simulations of stochastic individual contact models from icm.

**Usage**

```
## S3 method for class 'icm'
merge(x, y, ...)
```

**Arguments**

x	An EpiModel object of class <code>icm</code> .
y	Another EpiModel object of class <code>icm</code> , with the identical model parameterization as x.
...	Additional merge arguments (not used).

**Details**

This merge function combines the results of two independent simulations of `icm` class models, simulated under separate function calls. The model parameterization between the two calls must be exactly the same, except for the number of simulations in each call. This allows for manual parallelization of model simulations.

This merge function does not work the same as the default merge, which allows for a combined object where the structure differs between the input elements. Instead, the function checks that objects are identical in model parameterization in every respect (except number of simulations) and binds the results.

**Examples**

```
param <- param.icm(inf.prob = 0.2, act.rate = 0.8)
init <- init.icm(s.num = 1000, i.num = 100)
control <- control.icm(type = "SI", nsteps = 10,
                      nsims = 3, verbose = FALSE)
x <- icm(param, init, control)

control <- control.icm(type = "SI", nsteps = 10,
                      nsims = 1, verbose = FALSE)
y <- icm(param, init, control)

z <- merge(x, y)
x$epi
y$epi
z$epi
```

**Description**

Merges epidemiological data from two independent simulations of stochastic network models from `netsim`.

**Usage**

```
## S3 method for class 'netsim'
merge(
  x,
  y,
  keep.transmat = TRUE,
  keep.network = TRUE,
  keep.nwstats = TRUE,
  keep.other = TRUE,
  param.error = TRUE,
  ...
)
```

**Arguments**

x	An EpiModel object of class <code>netsim</code> .
y	Another EpiModel object of class <code>netsim</code> , with the identical model parameterization as x.
keep.transmat	If TRUE, keep the transmission matrices from the original x and y elements.
keep.network	If TRUE, keep the networkDynamic objects from the original x and y elements.
keep.nwstats	If TRUE, keep the network statistics (as set by the <code>nwstats.formula</code> parameter in <code>control.netsim</code> ) from the original x and y elements.
keep.other	If TRUE, keep the other simulation elements (as set by the <code>save.other</code> parameter in <code>control.netsim</code> ) from the original x and y elements.
param.error	If TRUE, if x and y have different params (in <code>param.net</code> ) or controls (passed in <code>control.net</code> ) an error will prevent the merge. Use FALSE to override that check.
...	Additional merge arguments (not currently used).

**Details**

This merge function combines the results of two independent simulations of `netsim` class models, simulated under separate function calls. The model parameterization between the two calls must be exactly the same, except for the number of simulations in each call. This allows for manual parallelization of model simulations.

This merge function does not work the same as the default merge, which allows for a combined object where the structure differs between the input elements. Instead, the function checks that objects are identical in model parameterization in every respect (except number of simulations) and binds the results.

**Examples**

```
# Network model
nw <- network.initialize(n = 100, directed = FALSE)
coef.diss <- dissolution_coefs(dissolution = ~offset(edges), duration = 10)
est <- netest(nw, formation = ~edges, target.stats = 25,
  coef.diss = coef.diss, verbose = FALSE)
```

```

# Epidemic models
param <- param.net(inf.prob = 1)
init <- init.net(i.num = 1)
control <- control.net(type = "SI", nsteps = 20, nsims = 2,
                      save.nwstats = TRUE,
                      nwstats.formula = ~edges + degree(0),
                      verbose = FALSE)
x <- netsim(est, param, init, control)
y <- netsim(est, param, init, control)

# Merging
z <- merge(x, y)
x$epi
y$epi
z$epi

```

modules.icm

*Modules for Stochastic Individual Contact Models*

## Description

Stochastic individual contact models of infectious disease simulate epidemics in which contacts between individuals are instantaneous events in discrete time. They are intended to be the stochastic microsimulation analogs to deterministic compartmental models.

The `icm` function handles both the simulation tasks. Within this function are a series of modules that initialize the simulation and then simulate new infections, recoveries, and vital dynamics at each time step. A module also handles the basic bookkeeping calculations for disease prevalence.

Writing original ICMs will require modifying the existing modules or adding new modules to the workflow in `icm`. The existing modules may be used as a template for replacement or new modules.

This help page presents a brief overview of the module functions in the order in which they are used within `icm`, in order to help guide users in writing their own module functions. These module functions are not shown on the help index since they are not called directly by the end-user. To understand these functions in more detail, review the separate help pages listed below.

## Initialization Module

This function sets up agent attributes, like disease status, on the network at the starting time step of disease simulation,  $t_1$ . For multiple-simulation function calls, these are reset at the beginning of each simulation.

- `initialize.icm`: sets which agents are initially infected, through the initial conditions passed in `init.icm`.

### Disease Status Modification Modules

The main disease simulation occurs at each time step given the current state of the population at that step. Infection of agents is simulated as a function of disease parameters and population composition. Recovery of agents is likewise simulated with respect to infected nodes. These functions also analyze the flows for summary measures such as disease incidence.

- `infection.icm`: randomly draws an edgelist given the parameters, subsets the list for discordant pairs, and simulates transmission on those discordant pairs through a series of draws from a binomial distribution.
- `recovery.icm`: simulates recovery from infection either to a lifelong immune state (for SIR models) or back to the susceptible state (for SIS models), as a function of the recovery rate specified in the `rec.rate` parameter. The recovery rate may vary for two-group models.

### Demographic Modules

Vital dynamics such as arrival and departure processes are simulated at each time step to update entries into and exits from the population. These are used in open-population ICMs.

- `departures.icm`: randomly simulates departures or exits for agents given the departure rate specified in the disease-state and group-specific departure parameters in `param.icm`. This involves deactivating agents from the population, but their historical data is preserved in the simulation.
- `arrivals.icm`: randomly simulates new arrivals into the population given the current population size and the arrival rate parameters. This involves adding new agents into the population.

### Bookkeeping Module

Simulations require bookkeeping at each time step to calculate the summary epidemiological statistics used in the model output analysis.

- `get_prev.icm`: calculates the number in each disease state (susceptible, infected, recovered) at each time step for those active agents in the population.

### Description

Stochastic network models of infectious disease in EpiModel require statistical modeling of networks, simulation of those networks forward through time, and simulation of epidemic dynamics on top of those evolving networks. The `netsim` function handles both the network and epidemic simulation tasks. Within this function are a series of modules that initialize the simulation and then simulate new infections, recoveries, and demographics on the network. Modules also handle the resimulation of the network and some bookkeeping calculations for disease prevalence.

Writing original network models that expand upon our "base" model set will require modifying the existing modules or adding new modules to the workflow in `netsim`. The existing modules may be used as a template for replacement or new modules.

This help page provides an orientation to these module functions, in the order in which they are used within `netsim`, to help guide users in writing their own functions. These module functions are not shown on the help index since they are not called directly by the end-user. To understand these functions in more detail, review the separate help pages listed below.

## Initialization Module

This function sets up nodal attributes, like disease status, on the network at the starting time step of disease simulation,  $t_1$ . For multiple-simulation function calls, these are reset at the beginning of each individual simulation.

- `initialize.net`: sets up the master data structure used in the simulation, initializes which nodes are infected (via the initial conditions passed in `init.net`), and simulates a first time step of the networks given the network model fit from `netest`.

## Disease Status Modification Modules

The main disease simulation occurs at each time step given the current state of the network at that step. Infection of nodes is simulated as a function of attributes of the nodes and the edges. Recovery of nodes is likewise simulated as a function of nodal attributes of those infected nodes. These functions also calculate summary flow measures such as disease incidence.

- `infection.net`: simulates disease transmission given an edgelist of discordant partnerships by calculating the relevant transmission and act rates for each edge, and then updating the nodal attributes and summary statistics.
- `recovery.net`: simulates recovery from infection either to a lifelong immune state (for SIR models) or back to the susceptible state (for SIS models), as a function of the recovery rate parameters specified in `param.net`.

## Demographic Modules

Demographics such as arrival and departure processes are simulated at each time step to update entries into and exits from the network. These are used in dependent network models, in which the network is resimulated at each time step to account for the nodal changes affecting the edges.

- `departures.net`: randomly simulates departure for nodes given their disease status (susceptible, infected, recovered), and their mode-specific departure rates specified in `param.net`. departures involve deactivating nodes, which are then deleted from the network if `delete.nodes=TRUE` is set in `control.net`.
- `arrivals.net`: randomly simulates new arrivals into the network given the current population size and the arrival rate specified in the `a.rate` parameters. This involves adding new nodes into the network.

## Network Resimulation Module

In dependent network models, the network object is resimulated at each time step to account for changes in the size of the network (changed through entries and exits), and the disease status of the nodes.

- `edges_correct`: adjusts the edges coefficient of a network model to account for changes in the population size due to entries and exits. The default behavior is to preserve the mean degree (average number of edges per person) in response to change population sizes.
- `resim_nets`: resimulates the network object one time step forward given the set of formation and dissolution coefficients estimated in `netest`. This function also deletes the inactive nodes if the `delete.nodes` control is set to `TRUE`.

## Bookkeeping Module

Network simulations require bookkeeping at each time step to calculate the summary epidemiological statistics used in the model output analysis.

- `get_prev.net`: calculates the number in each disease state (susceptible, infected, recovered) at each time step for those active nodes in the network. If the `epi.by` control is used, it calculates these statistics by a set of specified nodal attributes.
- `verbose.net`: summarizes the current state of the simulation and prints this to the console.

---

mutate_epi	<i>Add New Epidemiology Variables</i>
------------	---------------------------------------

---

## Description

Inspired by `dplyr::mutate`, `mutate_epi` adds new variables to the epidemiological and related variables within simulated model objects of any class in `EpiModel`.

## Usage

```
mutate_epi(x, ...)
```

## Arguments

<code>x</code>	An <code>EpiModel</code> object of class <code>dcm</code> , <code>icm</code> , or <code>netsim</code> .
<code>...</code>	Name-value pairs of expressions (see examples below).

## Examples

```
# DCM example
param <- param.dcm(inf.prob = 0.2, act.rate = 0.25)
init <- init.dcm(s.num = 500, i.num = 1)
control <- control.dcm(type = "SI", nsteps = 500)
mod1 <- dcm(param, init, control)
mod1 <- mutate_epi(mod1, prev = i.num/num)
plot(mod1, y = "prev")

# Network model example
nw <- network.initialize(n = 100, bipartite = 50, directed = FALSE)
formation <- ~edges
target.stats <- 50
```



```

coef.diss <- dissolution_coefs(dissolution = ~offset(edges), duration = 20)
est1 <- netest(nw, formation, target.stats, coef.diss, verbose = FALSE)

param <- param.net(inf.prob = 0.3, inf.prob.m2 = 0.15)
init <- init.net(i.num = 1, i.num.m2 = 0)
control <- control.net(type = "SI", nsteps = 10, nsims = 3,
                      verbose = FALSE)
mod1 <- netsim(est1, param, init, control)
mod1

# Add the prevalences to the dataset
mod1 <- mutate_epi(mod1, i.prev = i.num / num,
                  i.prev.m2 = i.num.m2 / num.m2)
plot(mod1, y = c("i.prev", "i.prev.m2"), qnts = 0.5, legend = TRUE)

# Add incidence rate per 100 person years (assume time step = 1 week)
mod1 <- mutate_epi(mod1, ir100 = 5200*(si.flow + si.flow.m2) /
                  (s.num + s.num.m2))

as.data.frame(mod1)
as.data.frame(mod1, out = "mean")

```

---

netdx

---

*Dynamic Network Model Diagnostics*


---

## Description

Runs dynamic diagnostics on an ERGM/STERGM estimated through netest.

## Usage

```

netdx(
  x,
  nsims = 1,
  dynamic = TRUE,
  nsteps,
  nwstats.formula = "formation",
  set.control.ergm,
  set.control.stergm,
  sequential = TRUE,
  keep.tedgelist = FALSE,
  keep.tnetwork = FALSE,
  verbose = TRUE,
  ncores = 1,
  skip.dissolution = FALSE
)

```

## Arguments

<code>x</code>	An EpiModel object of class <code>netest</code> .
<code>nsims</code>	Number of simulations to run.
<code>dynamic</code>	If TRUE, runs dynamic diagnostics. If FALSE and the <code>netest</code> object was fit with the Edges Dissolution approximation method, simulates from the static ERGM fit.
<code>nsteps</code>	Number of time steps per simulation (dynamic simulations only).
<code>nwstats.formula</code>	A right-hand sided ERGM formula with the network statistics of interest. The default is the formation formula of the network model contained in <code>x</code> .
<code>set.control.ergm</code>	Control arguments passed to <code>simulate.ergm</code> (see details).
<code>set.control.stergm</code>	Control arguments passed to <code>simulate.stergm</code> (see details).
<code>sequential</code>	For static diagnostics (dynamic=FALSE): if FALSE, each of the <code>nsims</code> simulated Markov chains begins at the initial network; If TRUE, the end of one simulation is used as the start of the next.
<code>keep.tedgelist</code>	If TRUE, keep the timed edgelist generated from the dynamic simulations. Returned in the form of a list of matrices, with one entry per simulation. Accessible at <code>\$edgelist</code> .
<code>keep.tnetwork</code>	If TRUE, keep the full <code>networkDynamic</code> objects from the dynamic simulations. Returned in the form of a list of <code>nD</code> objects, with one entry per simulation. Accessible at <code>\$network</code> .
<code>verbose</code>	Print progress to the console.
<code>ncores</code>	Number of processor cores to run multiple simulations on, using the <code>foreach</code> and <code>doParallel</code> implementations.
<code>skip.dissolution</code>	If TRUE, skip over the calculations of duration and dissolution stats in <code>netdx</code> .

## Details

The `netdx` function handles dynamic network diagnostics for network models fit with the `netest` function. Given the fitted model, `netdx` simulates a specified number of dynamic networks for a specified number of time steps per simulation. The network statistics in `nwstats.formula` are saved for each time step. Summary statistics for the formation model terms, as well as dissolution model and relational duration statistics, are then calculated and can be accessed when printing or plotting the `netdx` object.

## Control Arguments

Models fit with the full STERGM method in `netest` (setting `edapprox` argument to FALSE) require only a call to `simulate.stergm`. Control parameters for those simulations may be set using `set.control.stergm` in `netdx`. The parameters should be input through the `control.simulate.stergm()` function, with the available parameters listed in the [control.simulate.stergm](#) help page in the `tergm` package.

Models fit with the ERGM method with the edges dissolution approximation (setting `edapprox` to `TRUE`) require a call first to `simulate.ergm` for simulating an initial network, and second to `simulate.network` for simulating that static network forward through time. Control parameters may be set for both processes in `netdx`. For the first, the parameters should be input through the `control.simulate.ergm()` function, with the available parameters listed in the [control.simulate.ergm](#) help page in the `ergm` package. For the second, parameters should be input through the `control.simulate.network()` function, with the available parameters listed in the [control.simulate.network](#) help page in the `tergm` package. An example is shown below.

### See Also

Plot these model diagnostics with [plot.netdx](#).

### Examples

```
## Not run:
# Network initialization and model parameterization
nw <- network.initialize(100, directed = FALSE)
formation <- ~edges
target.stats <- 50
coef.diss <- dissolution_coefs(dissolution = ~offset(edges), duration = 25)

# Estimate the model
est <- netest(nw, formation, target.stats, coef.diss, verbose = FALSE)

# Static diagnostics on the ERGM fit
dx1 <- netdx(est, nsims = 1e4, dynamic = FALSE,
             nwstats.formula = ~edges + meandeg + concurrent)
dx1
plot(dx1, method = "b", stats = c("edges", "concurrent"))

# Dynamic diagnostics on the STERGM approximation
dx2 <- netdx(est, nsims = 5, nsteps = 500,
             nwstats.formula = ~edges + meandeg + concurrent,
             set.control.ergm = control.simulate.ergm(MCMC.burnin = 1e6))
dx2
plot(dx2, stats = c("edges", "meandeg"), plots.joined = FALSE)
plot(dx2, type = "duration")
plot(dx2, type = "dissolution", qnts.col = "orange2")
plot(dx2, type = "dissolution", method = "b", col = "bisque")

## End(Not run)
```

### Description

Estimates statistical network models using the exponential random graph modeling (ERGM) framework with extensions for dynamic/temporal models (STERGM).

**Usage**

```
netest(
  nw,
  formation,
  target.stats,
  coef.diss,
  constraints,
  coef.form = NULL,
  edapprox = TRUE,
  set.control.ergm,
  set.control.stergm,
  verbose = FALSE
)
```

**Arguments**

<code>nw</code>	An object of class <code>network</code> .
<code>formation</code>	Right-hand sided STERGM formation formula in the form <code>~edges + ...</code> , where <code>...</code> are additional network statistics.
<code>target.stats</code>	Vector of target statistics for the formation model, with one number for each network statistic in the model.
<code>coef.diss</code>	An object of class <code>disscoef</code> output from the <a href="#">dissolution_coefs</a> function.
<code>constraints</code>	Right-hand sided formula specifying constraints for the modeled network, in the form <code>~...</code> , where <code>...</code> are constraint terms. By default, no constraints are set.
<code>coef.form</code>	Vector of coefficients for the offset terms in the formation formula.
<code>edapprox</code>	If <code>TRUE</code> , use the indirect edges dissolution approximation method for the dynamic model fit, otherwise use the more time-intensive full STERGM estimation (see details).
<code>set.control.ergm</code>	Control arguments passed to <code>simulate.ergm</code> (see details).
<code>set.control.stergm</code>	Control arguments passed to <code>simulate.stergm</code> (see details).
<code>verbose</code>	Print model fitting progress to console.

**Details**

`netest` is a wrapper function for the `ergm` and `stergm` functions that estimate static and dynamic network models, respectively. Network model estimation is the first step in simulating a stochastic network epidemic model in `EpiModel`. The output from `netest` is a necessary input for running the epidemic simulations in [netsim](#). With a fitted network model, one should always first proceed to model diagnostics, available through the [netdx](#) function, to check model fit. A detailed description of fitting these models, along with examples, may be found in the [Basic Network Models](#) tutorial.

## Edges Dissolution Approximation

The edges dissolution approximation method is described in Carnegie et al. This approximation requires that the dissolution coefficients are known, that the formation model is being fit to cross-sectional data conditional on those dissolution coefficients, and that the terms in the dissolution model are a subset of those in the formation model. Under certain additional conditions, the formation coefficients of a STERGM model are approximately equal to the coefficients of that same model fit to the observed cross-sectional data as an ERGM, minus the corresponding coefficients in the dissolution model. The approximation thus estimates this ERGM (which is typically much faster than estimating a STERGM) and subtracts the dissolution coefficients.

The conditions under which this approximation best hold are when there are few relational changes from one time step to another; i.e. when either average relational durations are long, or density is low, or both. Conveniently, these are the same conditions under which STERGM estimation is slowest. Note that the same approximation is also used to obtain starting values for the STERGM estimate when the latter is being conducted. The estimation does not allow for calculation of standard errors, p-values, or likelihood for the formation model; thus, this approach is of most use when the main goal of estimation is to drive dynamic network simulations rather than to conduct inference on the formation model. The user is strongly encouraged to examine the behavior of the resulting simulations to confirm that the approximation is adequate for their purposes. For an example, see the vignette for the package `tergm`.

## Control Arguments

The `ergm` and `stergm` functions allow control settings for the model fitting process. When fitting a STERGM directly (setting `edapprox` to `FALSE`), control parameters may be passed to the `stergm` function with the `set.control.stergm` argument in `netest`. The controls should be input through the `control.stergm()` function, with the available parameters listed in the [control.stergm](#) help page in the `tergm` package.

When fitting a STERGM indirectly (setting `edapprox` to `TRUE`), control settings may be passed to the `ergm` function using `set.control.ergm` in `netest`. The controls should be input through the `control.ergm()` function, with the available parameters listed in the [control.ergm](#) help page in the `ergm` package. An example is below.

## References

- Krivitsky PN, Handcock MS. "A separable model for dynamic networks." *JRSS(B)*. 2014; 76.1:29-46.
- Carnegie NB, Krivitsky PN, Hunter DR, Goodreau SM. An approximation method for improving dynamic network model fitting. *Journal of Computational and Graphical Statistics*. 2014; 24(2): 502-519.
- Jenness SM, Goodreau SM and Morris M. EpiModel: An R Package for Mathematical Modeling of Infectious Disease over Networks. *Journal of Statistical Software*. 2018; 84(8): 1-47.

## See Also

Use [netdx](#) to diagnose the fitted network model, and [netsim](#) to simulate epidemic spread over a simulated dynamic network consistent with the model fit.

## Examples

```
# Initialize a network of 100 nodes
nw <- network.initialize(n = 100, directed = FALSE)

# Set formation formula
formation <- ~edges + concurrent

# Set target statistics for formation
target.stats <- c(50, 25)

# Obtain the offset coefficients
coef.diss <- dissolution_coefs(dissolution = ~offset(edges), duration = 10)

# Estimate the STERGM using the edges dissolution approximation
est <- netest(nw, formation, target.stats, coef.diss,
              set.control.ergm = control.ergm(MCMC.burnin = 1e5,
                                              MCMC.interval = 1000))

est

# To estimate the STERGM directly, use edapprox = FALSE
# est2 <- netest(nw, formation, target.stats, coef.diss, edapprox = FALSE)
```

---

netsim

*Stochastic Network Models*

---

## Description

Simulates stochastic network epidemic models for infectious disease.

## Usage

```
netsim(x, param, init, control)
```

## Arguments

x	Fitted network model object, as an object of class <code>netest</code> . Alternatively, if restarting a previous simulation, may be an object of class <code>netsim</code> .
param	Model parameters, as an object of class <code>param.net</code> .
init	Initial conditions, as an object of class <code>init.net</code> .
control	Control settings, as an object of class <code>control.net</code> .

## Details

Stochastic network models explicitly represent phenomena within and across edges (pairs of nodes that remain connected) over time. This enables edges to have duration, allowing for repeated transmission-related acts within the same dyad, specification of edge formation and dissolution rates, control over the temporal sequencing of multiple edges, and specification of network-level

features. A detailed description of these models, along with examples, is found in the [Basic Network Models](#) tutorial.

The `netsim` function performs modeling of both the base model types and original models. Base model types include one-mode and bipartite models with disease types for Susceptible-Infected (SI), Susceptible-Infected-Recovered (SIR), and Susceptible-Infected-Susceptible (SIS).

Original models may be parameterized by writing new process modules that either take the place of existing modules (for example, disease recovery), or supplement the set of existing processes with a new one contained in a new module. This functionality is documented in the [Solving New Network Models](#) tutorial. The list of modules within `netsim` available for modification is listed in [modules.net](#).

## Value

A list of class `netsim` with the following elements:

- **param:** the epidemic parameters passed into the model through `param`, with additional parameters added as necessary.
- **control:** the control settings passed into the model through `control`, with additional controls added as necessary.
- **epi:** a list of data frames, one for each epidemiological output from the model. Outputs for base models always include the size of each compartment, as well as flows in, out of, and between compartments.
- **stats:** a list containing two sublists, `nwstats` for any network statistics saved in the simulation, and `transmat` for the transmission matrix saved in the simulation. See [control.net](#) and the Tutorial for further details.
- **network:** a list of `networkDynamic` objects (or `network` objects if `delete.nodes` was set to `TRUE`), one for each model simulation.

## References

Jenness SM, Goodreau SM and Morris M. EpiModel: An R Package for Mathematical Modeling of Infectious Disease over Networks. *Journal of Statistical Software*. 2018; 84(8): 1-47.

## See Also

Extract the model results with [as.data.frame.netsim](#). Summarize the time-specific model results with [summary.netsim](#). Plot the model results with [plot.netsim](#).

## Examples

```
## Not run:
## Example 1: Independent SI Model
# Network model estimation
nw <- network.initialize(n = 100, bipartite = 50, directed = FALSE)
formation <- ~edges
target.stats <- 50
coef.diss <- dissolution_coefs(dissolution = ~offset(edges), duration = 20)
est1 <- netest(nw, formation, target.stats, coef.diss, verbose = FALSE)
```

```

# Epidemic model
param <- param.net(inf.prob = 0.3, inf.prob.m2 = 0.15)
init <- init.net(i.num = 10, i.num.m2 = 10)
control <- control.net(type = "SI", nsteps = 100, nsims = 5, verbose.int = 0)
mod1 <- netsim(est1, param, init, control)

# Print, plot, and summarize the results
mod1
plot(mod1)
summary(mod1, at = 50)

## Example 2: Dependent SIR Model
# Recalculate dissolution coefficient with departure rate
coef.diss <- dissolution_coefs(dissolution = ~offset(edges), duration = 20,
                              d.rate = 0.0021)

# Reestimate the model with new coefficient
est2 <- netest(nw, formation, target.stats, coef.diss, verbose = FALSE)

# Reset parameters to include demographic rates
param <- param.net(inf.prob = 0.3, inf.prob.m2 = 0.15,
                  rec.rate = 0.02, rec.rate.m2 = 0.02,
                  a.rate = 0.002, a.rate.m2 = NA,
                  ds.rate = 0.001, ds.rate.m2 = 0.001,
                  di.rate = 0.001, di.rate.m2 = 0.001,
                  dr.rate = 0.001, dr.rate.m2 = 0.001)
init <- init.net(i.num = 10, i.num.m2 = 10,
                r.num = 0, r.num.m2 = 0)
control <- control.net(type = "SIR", nsteps = 100, nsims = 5)

# Simulate the model with new network fit
mod2 <- netsim(est2, param, init, control)

# Print, plot, and summarize the results
mod2
plot(mod2)
summary(mod2, at = 100)

## End(Not run)

```

## Description

Sets the epidemic parameters for deterministic compartmental models simulated with dcm.



**Usage**

```

param.dcm(
  inf.prob,
  inter.eff,
  inter.start,
  act.rate,
  rec.rate,
  a.rate,
  ds.rate,
  di.rate,
  dr.rate,
  inf.prob.g2,
  act.rate.g2,
  rec.rate.g2,
  a.rate.g2,
  ds.rate.g2,
  di.rate.g2,
  dr.rate.g2,
  balance,
  ...
)

```

**Arguments**

<code>inf.prob</code>	Probability of infection per transmissible act between a susceptible and an infected person. In two-group models, this is the probability of infection for the group 1 members.
<code>inter.eff</code>	Efficacy of an intervention which affects the per-act probability of infection. Efficacy is defined as 1 - the relative hazard of infection given exposure to the intervention, compared to no exposure.
<code>inter.start</code>	Time step at which the intervention starts, between 1 and the number of time steps specified in the model. This will default to 1 if the <code>inter.eff</code> is defined but this parameter is not.
<code>act.rate</code>	Average number of transmissible acts per person per unit time. For two-group models, this is the number of acts per group 1 persons per unit time; a balance between the acts in groups 1 and 2 is necessary, and set using the <code>balance</code> parameter (see details).
<code>rec.rate</code>	Average rate of recovery with immunity (in SIR models) or re-susceptibility (in SIS models). The recovery rate is the reciprocal of the disease duration. For two-group models, this is the recovery rate for group 1 persons only. This parameter is only used for SIR and SIS models.
<code>a.rate</code>	Arrival or entry rate. For one-group models, the arrival rate is the rate of new arrivals per person per unit time. For two-group models, the arrival rate may be parameterized as a rate per group 1 person time (with group 1 persons representing females), and with the <code>a.rate.g2</code> rate set as described below.
<code>ds.rate</code>	Departure or exit rate for susceptible. For two-group models, it is the rate for the group 1 susceptible only.

di.rate	Departure or exit rate for infected. For two-group models, it is the rate for the group 1 infected only.
dr.rate	Departure or exit rate for recovered. For two-group models, it is the rate for the group 1 recovered only. This parameter is only used for SIR models.
inf.prob.g2	Probability of infection per transmissible act between a susceptible group 2 person and an infected group 1 person. It is the probability of infection to group 2 members.
act.rate.g2	Average number of transmissible acts per group 2 person per unit time; a balance between the acts in groups 1 and 2 is necessary, and set using the balance parameter (see details).
rec.rate.g2	Average rate of recovery with immunity (in SIR models) or re-susceptibility (in SIS models) for group 2 persons. This parameter is only used for two-group SIR and SIS models.
a.rate.g2	Arrival or entry rate for group 2. This may either be specified numerically as the rate of new arrivals per group 2 persons per unit time, or as NA in which case the group 1 rate, a.rate, governs the group 2 rate. The latter is used when, for example, the first group is conceptualized as female, and the female population size determines the arrival rate. Such arrivals are evenly allocated between the two groups.
ds.rate.g2	Departure or exit rate for group 2 susceptible.
di.rate.g2	Departure or exit rate for group 2 infected.
dr.rate.g2	Departure or exit rate for group 2 recovered. This parameter is only used for SIR model types.
balance	For two-group models, balance the act.rate to the rate set for group 1 (with balance="g1") or group 2 (with balance="g2"). See details.
...	Additional arguments passed to model.

### Details

param.dcm sets the epidemic parameters for deterministic compartmental models solved with the `dcm` function. The models may use the base types, for which these parameters are used, or original model specifications for which these parameters may be used (but not necessarily). A detailed description of DCM parameterization for base models is found in the **Basic DCMs** tutorial.

For base models, the model specification will be selected as a function of the model parameters entered here and the control settings in `control.dcm`. One-group and two-group models are available, where the former assumes a homogenous mixing in the population and the latter assumes a purely heterogenous mixing between two distinct partitions in the population (e.g., men and women). Specifying any group two parameters (those with a .g2) implies the simulation of a two-group model. All the parameters for a desired model type must be specified, even if they are zero.

### Act Balancing

In two-group models, a balance between the number of acts for group 1 members and those for group 2 members must be maintained. With purely heterogenous mixing, the product of one group size and act rate must equal the product of the other group size and act rate:  $N_1\alpha_1 = N_2\alpha_2$ , where  $N_i$  is the group size and  $\alpha_i$  the group-specific act rates at time  $t$ . The balance parameter here

specifies which group's act rate should control the others with respect to balancing. See the [Basic DCMs](#) tutorial for further details.

### Sensitivity Analyses

dcm has been designed to easily run DCM sensitivity analyses, where a series of models varying one or more of the model parameters is run. This is possible by setting any parameter as a vector of length greater than one. See both the example below and the [Basic DCMs](#) tutorial.

### New Model Types

To build original model specifications outside of the base models, start by consulting the [Solving New DCMs with EpiModel](#) tutorial. Briefly, an original model may use either the existing model parameters named here, an original set of parameters, or a combination of both. The `...` argument allows the user to pass an arbitrary set of new model parameters into `param.dcm`. Whereas there are strict checks for base models that the model parameters are valid, parameter validity is the user's responsibility with these original models.

### See Also

Use [init.dcm](#) to specify the initial conditions and [control.dcm](#) to specify the control settings. Run the parameterized model with [dcm](#).

---

param.icm

*Epidemic Parameters for Stochastic Individual Contact Models*

---

### Description

Sets the epidemic parameters for stochastic individual contact models simulated with `icm`.

### Usage

```
param.icm(
  inf.prob,
  inter.eff,
  inter.start,
  act.rate,
  rec.rate,
  a.rate,
  ds.rate,
  di.rate,
  dr.rate,
  inf.prob.g2,
  act.rate.g2,
  rec.rate.g2,
  a.rate.g2,
  ds.rate.g2,
  di.rate.g2,
```

```

    dr.rate.g2,
    balance,
    ...
)

```

### Arguments

<code>inf.prob</code>	Probability of infection per transmissible act between a susceptible and an infected person. In two-group models, this is the probability of infection for the group 1 members.
<code>inter.eff</code>	Efficacy of an intervention which affects the per-act probability of infection. Efficacy is defined as 1 - the relative hazard of infection given exposure to the intervention, compared to no exposure.
<code>inter.start</code>	Time step at which the intervention starts, between 1 and the number of time steps specified in the model. This will default to 1 if the <code>inter.eff</code> is defined but this parameter is not.
<code>act.rate</code>	Average number of transmissible acts per person per unit time. For two-group models, this is the number of acts per group 1 persons per unit time; a balance between the acts in groups 1 and 2 is necessary, and set using the <code>balance</code> parameter (see details).
<code>rec.rate</code>	Average rate of recovery with immunity (in SIR models) or re-susceptibility (in SIS models). The recovery rate is the reciprocal of the disease duration. For two-group models, this is the recovery rate for group 1 persons only. This parameter is only used for SIR and SIS models.
<code>a.rate</code>	Arrival or entry rate. For one-group models, the arrival rate is the rate of new arrivals per person per unit time. For two-group models, the arrival rate may be parameterized as a rate per group 1 person time (with group 1 persons representing females), and with the <code>a.rate.g2</code> rate set as described below.
<code>ds.rate</code>	Departure or exit rate for susceptible. For two-group models, it is the rate for the group 1 susceptible only.
<code>di.rate</code>	Departure or exit rate for infected. For two-group models, it is the rate for the group 1 infected only.
<code>dr.rate</code>	Departure or exit rate for recovered. For two-group models, it is the rate for the group 1 recovered only. This parameter is only used for SIR models.
<code>inf.prob.g2</code>	Probability of infection per transmissible act between a susceptible group 2 person and an infected group 1 person. It is the probability of infection to group 2 members.
<code>act.rate.g2</code>	Average number of transmissible acts per group 2 person per unit time; a balance between the acts in groups 1 and 2 is necessary, and set using the <code>balance</code> parameter (see details).
<code>rec.rate.g2</code>	Average rate of recovery with immunity (in SIR models) or re-susceptibility (in SIS models) for group 2 persons. This parameter is only used for two-group SIR and SIS models.
<code>a.rate.g2</code>	Arrival or entry rate for group 2. This may either be specified numerically as the rate of new arrivals per group 2 persons per unit time, or as NA in which case

	the group 1 rate, <code>a.rate</code> , governs the group 2 rate. The latter is used when, for example, the first group is conceptualized as female, and the female population size determines the arrival rate. Such arrivals are evenly allocated between the two groups.
<code>ds.rate.g2</code>	Departure or exit rate for group 2 susceptible.
<code>di.rate.g2</code>	Departure or exit rate for group 2 infected.
<code>dr.rate.g2</code>	Departure or exit rate for group 2 recovered. This parameter is only used for SIR model types.
<code>balance</code>	For two-group models, balance the <code>act.rate</code> to the rate set for group 1 (with <code>balance="g1"</code> ) or group 2 (with <code>balance="g2"</code> ). See details.
<code>...</code>	Additional arguments passed to model.

### Details

`param.icm` sets the epidemic parameters for the stochastic individual contact models simulated with the `icm` function. Models may use the base types, for which these parameters are used, or new process modules which may use these parameters (but not necessarily). A detailed description of ICM parameterization for base models is found in the [Basic ICMs](#) tutorial.

For base models, the model specification will be chosen as a result of the model parameters entered here and the control settings in `control.icm`. One-group and two-group models are available, where the former assumes a homogenous mixing in the population and the latter assumes a purely heterogenous mixing between two distinct partitions in the population (e.g., men and women). Specifying any group two parameters (those with a `.g2`) implies the simulation of a two-group model. All the parameters for a desired model type must be specified, even if they are zero.

### Act Balancing

In two-group models, a balance between the number of acts for group 1 members and those for group 2 members must be maintained. With purely heterogenous mixing, the product of one group size and act rate must equal the product of the other group size and act rate:  $N_1\alpha_1 = N_2\alpha_2$ , where  $N_i$  is the group size and  $\alpha_i$  the group-specific act rates at time  $t$ . The `balance` parameter here specifies which group's act rate should control the others with respect to balancing. See the [Basic DCMs](#) tutorial.

### New Modules

To build original models outside of the base models, new process modules may be constructed to replace the existing modules or to supplement the existing set. These are passed into the control settings in `control.icm`. New modules may use either the existing model parameters named here, an original set of parameters, or a combination of both. The `...` allows the user to pass an arbitrary set of original model parameters into `param.icm`. Whereas there are strict checks with default modules for parameter validity, these checks are the user's responsibility with new modules.

### See Also

Use `init.icm` to specify the initial conditions and `control.icm` to specify the control settings. Run the parameterized model with `icm`.

**Description**

Sets the epidemic parameters for stochastic network models simulated with [netsim](#).

**Usage**

```
param.net(
  inf.prob,
  inter.eff,
  inter.start,
  act.rate,
  rec.rate,
  a.rate,
  ds.rate,
  di.rate,
  dr.rate,
  inf.prob.m2,
  rec.rate.m2,
  a.rate.m2,
  ds.rate.m2,
  di.rate.m2,
  dr.rate.m2,
  ...
)
```

**Arguments**

inf.prob	Probability of infection per transmissible act between a susceptible and an infected person. In bipartite models, this is the probability of infection to the mode 1 nodes. This may also be a vector of probabilities, with each element corresponding to the probability in that time step of infection (see Time-Varying Parameters below).
inter.eff	Efficacy of an intervention which affects the per-act probability of infection. Efficacy is defined as 1 - the relative hazard of infection given exposure to the intervention, compared to no exposure.
inter.start	Time step at which the intervention starts, between 1 and the number of time steps specified in the model. This will default to 1 if the inter.eff is defined but this parameter is not.
act.rate	Average number of transmissible acts <i>per partnership</i> per unit time (see act.rate Parameter below). This may also be a vector of rates, with each element corresponding to the rate in in that time step of infection (see Time-Varying Parameters below).

rec.rate	Average rate of recovery with immunity (in SIR models) or re-susceptibility (in SIS models). The recovery rate is the reciprocal of the disease duration. For bipartite models, this is the recovery rate for mode 1 persons only. This parameter is only used for SIR and SIS models. This may also be a vector of rates, with each element corresponding to the rate in that time step of infection (see Time-Varying Parameters below).
a.rate	Arrival or entry rate. For one-mode models, the arrival rate is the rate of new arrivals per person per unit time. For bipartite models, the arrival rate may be parameterized as a rate per mode 1 person time (with mode 1 persons representing females), and with the a.rate.g2 rate set as described below.
ds.rate	Departure or exit rate for susceptible. For bipartite models, it is the rate for the mode 1 susceptible only.
di.rate	Departure or exit rate for infected. For bipartite models, it is the rate for the mode 1 infected only.
dr.rate	Departure or exit rate for recovered. For bipartite models, it is the rate for the mode 1 recovered only. This parameter is only used for SIR models.
inf.prob.m2	Probability of transmission given a transmissible act between a susceptible mode 2 person and an infected mode 1 person. It is the probability of transmission to mode 2 members.
rec.rate.m2	Average rate of recovery with immunity (in SIR models) or re-susceptibility (in SIS models) for mode 2 persons. This parameter is only used for bipartite SIR and SIS models.
a.rate.m2	Arrival or entry rate for mode 2. This may either be specified numerically as the rate of new arrivals per mode 2 persons per unit time, or as NA in which case the mode 1 rate, a.rate, governs the mode 2 rate. The latter is used when, for example, the first mode is conceptualized as female, and the female population size determines the arrival rate. Such arrivals are evenly allocated between the two modes.
ds.rate.m2	Departure or exit rate for mode 2 susceptible.
di.rate.m2	Departure or exit rate for mode 2 infected.
dr.rate.m2	Departure or exit rate for mode 2 recovered. This parameter is only used for SIR model types.
...	Additional arguments passed to model.

## Details

param.net sets the epidemic parameters for the stochastic network models simulated with the [netsim](#) function. Models may use the base types, for which these parameters are used, or new process modules which may use these parameters (but not necessarily). A detailed description of network model parameterization for base models is found in the [Basic Network Models](#) tutorial.

For base models, the model specification will be chosen as a result of the model parameters entered here and the control settings in [control.net](#). One-mode and two-mode models are available, where the latter assumes a heterogeneous mixing between two distinct partitions in the population (e.g., men and women). Specifying any bipartite parameters (those with a .m2) implies the simulation of a bipartite model. All the parameters for a desired model type must be specified, even if they are zero.

### The `act.rate` Parameter

A key difference between these network models and DCM/ICM classes is the treatment of transmission events. With DCM and ICM, contacts or partnerships are mathematically instantaneous events: they have no duration in time, and thus no changes may occur within them over time. In contrast, network models allow for partnership durations defined by the dynamic network model, summarized in the model dissolution coefficients calculated in [dissolution\\_coefs](#). Therefore, the `act.rate` parameter has a different interpretation here, where it is the number of transmissible acts *per partnership* per unit time.

### Time-Varying Parameters

The `inf.prob`, `act.rate`, `rec.rate` arguments (and their `.m2` companions) may be specified as time-varying parameters by passing in a vector of probabilities or rates, respectively. The value in each position on the vector then corresponds to the probability or rate at that discrete time step for the infected partner. For example, an `inf.prob` of `c(0.5, 0.5, 0.1)` would simulate a 0.5 transmission probability for the first two time steps of a person's infection, followed by a 0.1 for the third time step. If the infected person has not recovered or exited the population by the fourth time step, the third element in the vector will carry forward until one of those events occurs or the simulation ends. For further examples, see the NME tutorial, [Time-Varying Biology & Behavior](#).

### New Modules

To build original models outside of the base models, new process modules may be constructed to replace the existing modules or to supplement the existing set. These are passed into the control settings in [control.net](#). New modules may use either the existing model parameters named here, an original set of parameters, or a combination of both. The `...` allows the user to pass an arbitrary set of original model parameters into `param.net`. Whereas there are strict checks with default modules for parameter validity, these checks are the user's responsibility with new modules.

### See Also

Use [init.net](#) to specify the initial conditions and [control.net](#) to specify the control settings. Run the parameterized model with [netsim](#).

---

plot.dcm

*Plot Data from a Deterministic Compartmental Epidemic Model*

---

### Description

Plots epidemiological data from a deterministic compartment epidemic model solved with `dcm`.

### Usage

```
## S3 method for class 'dcm'
plot(
  x,
  y,
```



```

    popfrac = FALSE,
    run,
    col,
    lwd,
    lty,
    alpha = 0.9,
    legend,
    leg.name,
    leg.cex = 0.8,
    axs = "r",
    grid = FALSE,
    add = FALSE,
    ...
)

```

### Arguments

x	An EpiModel object of class dcm.
y	Output compartments or flows from dcm object to plot.
popfrac	If TRUE, plot prevalence of values rather than numbers (see details).
run	Run number to plot, for models with multiple runs (default is run 1).
col	Color for lines, either specified as a single color in a standard R color format, or alternatively as a color palette from <a href="#">RColorBrewer</a> (see details).
lwd	Line width for output lines.
lty	Line type for output lines.
alpha	Transparency level for lines, where 0 = transparent and 1 = opaque (see <a href="#">transco</a> ).
legend	Type of legend to plot. Values are "n" for no legend, "full" for full legend, and "lim" for limited legend (see details).
leg.name	Character string to use for legend, with the default determined automatically based on the y input.
leg.cex	Legend scale size.
axs	Plot axis type (see <a href="#">par</a> for details), with default of "r".
grid	If TRUE, a grid is added to the background of plot (see <a href="#">grid</a> for details), with default of nx by ny.
add	If TRUE, new plot window is not called and lines are added to existing plot window.
...	Additional arguments to pass to main plot window (see <a href="#">plot.default</a> ).

### Details

This function plots epidemiological outcomes from a deterministic compartmental model solved with [dcm](#). Depending on the number of model runs (sensitivity analyses) and number of groups, the default plot is the fractional proportion of each compartment in the model over time. The specific compartments or flows to plot may be set using the y parameter, and in multiple run models the specific run may also be specified.

### The popfrac Argument

Compartment prevalences are the size of a compartment over some denominator. To plot the raw numbers from any compartment, use `popfrac=FALSE`; this is the default. The `popfrac` parameter calculates and plots the denominators of all specified compartments using these rules: 1) for one-group models, the prevalence of any compartment is the compartment size divided by the total population size; 2) for two-group models, the prevalence of any compartment is the compartment size divided by the group size.

### Color Palettes

Since `dcm` supports multiple run sensitivity models, plotting the results of such models uses a complex color scheme for distinguishing runs. This is accomplished using the `RColorBrewer` color palettes, which include a range of linked colors using named palettes. For `plot.dcm`, one may either specify a brewer color palette listed in [brewer.pal.info](http://www.cookbook-engineering.com/brewer.pal.info), or, alternatively, a vector of standard R colors (named, hexadecimal, or positive integers; see [col2rgb](http://www.cookbook-engineering.com/col2rgb)).

### Plot Legends

There are three automatic legend types available, and the legend is added by default for plots. To turn off the legend, use `legend="n"`. To plot a legend with values for every line in a sensitivity analysis, use `legend="full"`. With models with many runs, this may be visually overwhelming. In those cases, use `legend="lim"` to plot a legend limited to the highest and lowest values of the varying parameter in the model. In cases where the default legend names are not helpful, one may override those names with the `leg.name` argument.

### See Also

[dcm](#), [brewer.pal.info](#)

### Examples

```
# Deterministic SIR model with varying act rate
param <- param.dcm(inf.prob = 0.2, act.rate = 1:10,
                  rec.rate = 1/3, a.rate = 0.011, ds.rate = 0.01,
                  di.rate = 0.03, dr.rate = 0.01)
init <- init.dcm(s.num = 1000, i.num = 1, r.num = 0)
control <- control.dcm(type = "SIR", nsteps = 100, dt = 0.25)
mod <- dcm(param, init, control)

# Plot disease prevalence by default
plot(mod)

# Plot prevalence of susceptibles
plot(mod, y = "s.num", popfrac = TRUE, col = "Greys")

# Plot number of susceptibles
plot(mod, y = "s.num", popfrac = FALSE, col = "Greys", grid = TRUE)

# Plot multiple runs of multiple compartments together
plot(mod, y = c("s.num", "i.num"),
```

```

run = 5, xlim = c(0, 50), grid = TRUE)
plot(mod, y = c("s.num", "i.num"),
run = 10, lty = 2, legend = "n", add = TRUE)

```

plot.icm

*Plot Data from a Stochastic Individual Contact Epidemic Model***Description**

Plots epidemiological data from a stochastic individual contact model simulated with icm.

**Usage**

```

## S3 method for class 'icm'
plot(
  x,
  y,
  popfrac = FALSE,
  sim.lines = FALSE,
  sims,
  sim.col,
  sim.lwd,
  sim.alpha,
  mean.line = TRUE,
  mean.smooth = TRUE,
  mean.col,
  mean.lwd = 2,
  mean.lty = 1,
  qnts = 0.5,
  qnts.col,
  qnts.alpha,
  qnts.smooth = TRUE,
  legend,
  leg.cex = 0.8,
  axs = "r",
  grid = FALSE,
  add = FALSE,
  ...
)

```

**Arguments**

x	An EpiModel model object of class netsim.
y	Output compartments or flows from netsim object to plot.
popfrac	If TRUE, plot prevalence of values rather than numbers (see details).

<code>sim.lines</code>	If TRUE, plot individual simulation lines. Default is to plot lines for one-group models but not for two-group models.
<code>sims</code>	If <code>type="epi"</code> or <code>"formation"</code> , a vector of simulation numbers to plot. If <code>type="network"</code> , a single simulation number for network plot, or else <code>"min"</code> to plot the simulation number with the lowest disease prevalence, <code>"max"</code> for the simulation with the highest disease prevalence, or <code>"mean"</code> for the simulation with the prevalence closest to the mean across simulations at the specified time step.
<code>sim.col</code>	Vector of any standard R color format for simulation lines.
<code>sim.lwd</code>	Line width for simulation lines.
<code>sim.alpha</code>	Transparency level for simulation lines, where 0 = transparent and 1 = opaque (see <a href="#">transco</a> ).
<code>mean.line</code>	If TRUE, plot mean of simulations across time.
<code>mean.smooth</code>	If TRUE, use a lowess smoother on the mean line.
<code>mean.col</code>	Vector of any standard R color format for mean lines.
<code>mean.lwd</code>	Line width for mean lines.
<code>mean.lty</code>	Line type for mean lines.
<code>qnts</code>	If numeric, plot polygon of simulation quantiles based on the range implied by the argument (see details). If FALSE, suppress polygon from plot.
<code>qnts.col</code>	Vector of any standard R color format for polygons.
<code>qnts.alpha</code>	Transparency level for quantile polygons, where 0 = transparent and 1 = opaque (see <a href="#">transco</a> ).
<code>qnts.smooth</code>	If TRUE, use a lowess smoother on quantile polygons.
<code>legend</code>	If TRUE, plot default legend.
<code>leg.cex</code>	Legend scale size.
<code>axs</code>	Plot axis type (see <a href="#">par</a> for details), with default to <code>"r"</code> .
<code>grid</code>	If TRUE, a grid is added to the background of plot (see <a href="#">grid</a> for details), with default of <code>nx</code> by <code>ny</code> .
<code>add</code>	If TRUE, new plot window is not called and lines are added to existing plot window.
<code>...</code>	additional arguments to pass.

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

```
## Example 1: Plotting multiple compartment values from SIR model
param <- param.icm(inf.prob = 0.5, act.rate = 0.5, rec.rate = 0.02)
init <- init.icm(s.num = 500, i.num = 1, r.num = 0)
control <- control.icm(type = "SIR", nsteps = 100,
                      nsims = 3, verbose = FALSE)
mod <- icm(param, init, control)
```

```

plot(mod, grid = TRUE)

## Example 2: Plot only infected with specific output from SI model
param <- param.icm(inf.prob = 0.25, act.rate = 0.25)
init <- init.icm(s.num = 500, i.num = 10)
control <- control.icm(type = "SI", nsteps = 100,
                      nsims = 3, verbose = FALSE)
mod2 <- icm(param, init, control)

# Plot prevalence
plot(mod2, y = "i.num", mean.line = FALSE, sim.lines = TRUE)

# Plot incidence
par(mfrow = c(1, 2))
plot(mod2, y = "si.flow", mean.smooth = TRUE, grid = TRUE)
plot(mod2, y = "si.flow", qnts.smooth = FALSE, qnts = 1)

```

---

plot.netdx

---

*Plot Dynamic Network Model Diagnostics*


---

## Description

Plots dynamic network model diagnostics calculated in netdx.

## Usage

```

## S3 method for class 'netdx'
plot(
  x,
  type = "formation",
  method = "l",
  sims,
  stats,
  sim.lines,
  sim.col,
  sim.lwd,
  mean.line = TRUE,
  mean.smooth = TRUE,
  mean.col,
  mean.lwd = 2,
  mean.lty = 1,
  qnts = 0.5,
  qnts.col,
  qnts.alpha,
  qnts.smooth = TRUE,
  targ.line = TRUE,
  targ.col,

```

```

    targ.lwd = 2,
    targ.lty = 2,
    plots.joined,
    legend,
    grid = FALSE,
    ...
)

```

## Arguments

<code>x</code>	An EpiModel object of class <code>netdx</code> .
<code>type</code>	Plot type, with options of "formation" for network model formation statistics, "duration" for dissolution model statistics for average edge duration, or "dissolution" for dissolution model statistics for proportion of ties dissolved per time step.
<code>method</code>	Plot method, with options of "l" for line plots and "b" for boxplots.
<code>sims</code>	If <code>type="epi"</code> or "formation", a vector of simulation numbers to plot. If <code>type="network"</code> , a single simulation number for network plot, or else "min" to plot the simulation number with the lowest disease prevalence, "max" for the simulation with the highest disease prevalence, or "mean" for the simulation with the prevalence closest to the mean across simulations at the specified time step.
<code>stats</code>	Network statistics to plot, among those specified in the call to <code>netdx</code> , with the default to plot all statistics contained in the object.
<code>sim.lines</code>	If TRUE, plot individual simulation lines. Default is to plot lines for one-group models but not for two-group models.
<code>sim.col</code>	Vector of any standard R color format for simulation lines.
<code>sim.lwd</code>	Line width for simulation lines.
<code>mean.line</code>	If TRUE, plot mean of simulations across time.
<code>mean.smooth</code>	If TRUE, use a lowess smoother on the mean line.
<code>mean.col</code>	Vector of any standard R color format for mean lines.
<code>mean.lwd</code>	Line width for mean lines.
<code>mean.lty</code>	Line type for mean lines.
<code>qnts</code>	If numeric, plot polygon of simulation quantiles based on the range implied by the argument (see details). If FALSE, suppress polygon from plot.
<code>qnts.col</code>	Vector of any standard R color format for polygons.
<code>qnts.alpha</code>	Transparency level for quantile polygons, where 0 = transparent and 1 = opaque (see <a href="#">transco</a> ).
<code>qnts.smooth</code>	If TRUE, use a lowess smoother on quantile polygons.
<code>targ.line</code>	If TRUE, plot target or expected value line for the statistic of interest.
<code>targ.col</code>	Vector of standard R colors for target statistic lines, with default colors based on RColorBrewer color palettes.
<code>targ.lwd</code>	Line width for the line showing the target statistic values.

targ.lty	Line type for the line showing the target statistic values.
plots.joined	If TRUE and type="formation", combine all target statistics in one plot, versus one plot per target statistic if FALSE.
legend	If TRUE, plot default legend.
grid	If TRUE, a grid is added to the background of plot (see <a href="#">grid</a> for details), with default of nx by ny.
...	additional arguments to pass.

## Details

The plot function for netdx objects will generate plots of two types of model diagnostic statistics that run as part of the diagnostic tools within that function. The `formation` plot shows the summary statistics requested in `nwstats.formula`, where the default includes those statistics in the network model formation formula specified in the original call to `netest`.

The duration plot shows the average age of existing edges at each time step, up until the maximum time step requested. This is calculated with the `edgelist_meanage` function. The age is used as an estimator of the average duration of edges in the equilibrium state.

The dissolution plot shows the proportion of the extant ties that are dissolved at each time step, up until the maximum time step requested. Typically, the proportion of ties that are dissolved is the reciprocal of the mean relational duration. This plot thus contains similar information to that in the duration plot, but should reach its expected value more quickly, since it is not subject to censoring.

The `plots.joined` argument will control whether the statistics in the formation plot are joined in one plot or plotted separately. The default is based on the number of network statistics requested. The layout of the separate plots within the larger plot window is also based on the number of statistics.

## See Also

[netdx](#)

## Examples

```
## Not run:
# Network initialization and model parameterization
nw <- network.initialize(100, directed = FALSE)
nw <- set.vertex.attribute(nw, "sex", rbinom(100, 1, 0.5))
formation <- ~edges + nodematch("sex")
target.stats <- c(50, 40)
coef.diss <- dissolution_coefs(dissolution = ~offset(edges), duration = 50)

# Estimate the model
est <- netest(nw, formation, target.stats, coef.diss, verbose = FALSE)

# Static diagnostics
dx1 <- netdx(est, nsims = 1e4, dynamic = FALSE,
             nwstats.formula = ~edges + meandeg + concurrent +
                               nodefactor("sex", base = 0) +
                               nodematch("sex"))
```

```

dx1

# Plot diagnostics
plot(dx1)
plot(dx1, stats = c("edges", "concurrent"), mean.col = "black",
      sim.lines = TRUE, plots.joined = FALSE)
plot(dx1, stats = "edges", method = "b",
      col = "seagreen3", grid = TRUE)

# Dynamic diagnostics
dx2 <- netdx(est, nsims = 10, nsteps = 500,
             nwstats.formula = ~edges + meandeg + concurrent +
                               nodefactor("sex", base = 0) +
                               nodematch("sex"))
dx2

# Formation statistics plots, joined and separate
plot(dx2, grid = TRUE)
plot(dx2, type = "formation", plots.joined = TRUE)
plot(dx2, type = "formation", sims = 1, plots.joined = TRUE,
      qnts = FALSE, sim.lines = TRUE, mean.line = FALSE)
plot(dx2, type = "formation", plots.joined = FALSE,
      stats = c("edges", "concurrent"), grid = TRUE)

plot(dx2, method = "b", col = "bisque", grid = TRUE)
plot(dx2, method = "b", stats = "meandeg", col = "dodgerblue")

# Duration statistics plot
plot(dx2, type = "duration", mean.col = "black", grid = TRUE)
plot(dx2, type = "duration", sims = 10, mean.line = FALSE, sim.line = TRUE,
      sim.col = "steelblue", sim.lwd = 3, targ.lty = 1, targ.lwd = 0.5)

# Dissolution statistics plot
plot(dx2, type = "dissolution", mean.col = "black", grid = TRUE)
plot(dx2, type = "dissolution", method = "b", col = "pink1")

## End(Not run)

```

---

plot.netsim

---

*Plot Data from a Stochastic Network Epidemic Model*


---

## Description

Plots epidemiological and network data from a stochastic network model simulated with netsim.

## Usage

```

## S3 method for class 'netsim'
plot(

```



```

x,
type = "epi",
y,
popfrac = FALSE,
sim.lines = FALSE,
sims,
sim.col,
sim.lwd,
sim.alpha,
mean.line = TRUE,
mean.smooth = TRUE,
mean.col,
mean.lwd = 2,
mean.lty = 1,
qnts = 0.5,
qnts.col,
qnts.alpha,
qnts.smooth = TRUE,
legend,
leg.cex = 0.8,
axs = "r",
grid = FALSE,
add = FALSE,
network = 1,
at = 1,
col.status = FALSE,
shp.bip = NULL,
stats,
targ.line = TRUE,
targ.col,
targ.lwd = 2,
targ.lty = 2,
plots.joined,
...
)

```

### Arguments

x	An EpiModel model object of class netsim.
type	Type of plot: "epi" for epidemic model results, "network" for a static network plot (plot.network), or "formation" for network formation statistics.
y	Output compartments or flows from netsim object to plot.
popfrac	If TRUE, plot prevalence of values rather than numbers (see details).
sim.lines	If TRUE, plot individual simulation lines. Default is to plot lines for one-group models but not for two-group models.
sims	If type="epi" or "formation", a vector of simulation numbers to plot. If type="network", a single simulation number for network plot, or else "min"

	to plot the simulation number with the lowest disease prevalence, "max" for the simulation with the highest disease prevalence, or "mean" for the simulation with the prevalence closest to the mean across simulations at the specified time step.
sim.col	Vector of any standard R color format for simulation lines.
sim.lwd	Line width for simulation lines.
sim.alpha	Transparency level for simulation lines, where 0 = transparent and 1 = opaque (see <a href="#">transco</a> ).
mean.line	If TRUE, plot mean of simulations across time.
mean.smooth	If TRUE, use a lowess smoother on the mean line.
mean.col	Vector of any standard R color format for mean lines.
mean.lwd	Line width for mean lines.
mean.lty	Line type for mean lines.
qnts	If numeric, plot polygon of simulation quantiles based on the range implied by the argument (see details). If FALSE, suppress polygon from plot.
qnts.col	Vector of any standard R color format for polygons.
qnts.alpha	Transparency level for quantile polygons, where 0 = transparent and 1 = opaque (see <a href="#">transco</a> ).
qnts.smooth	If TRUE, use a lowess smoother on quantile polygons.
legend	If TRUE, plot default legend.
leg.cex	Legend scale size.
axs	Plot axis type (see <a href="#">par</a> for details), with default to "r".
grid	If TRUE, a grid is added to the background of plot (see <a href="#">grid</a> for details), with default of nx by ny.
add	If TRUE, new plot window is not called and lines are added to existing plot window.
network	Network number, for simulations with multiple networks representing the population.
at	If type="network", time step for network graph.
col.status	If TRUE and type="network", automatic disease status colors (blue = susceptible, red = infected, green = recovered).
shp.bip	If type="network" and a bipartite simulation, shapes for the mode 2 vertices, with acceptable inputs of "triangle" and "square". Mode 1 vertices will be circles.
stats	If type="formation", network statistics to plot, among those specified in <code>nwstats.formula</code> of <a href="#">control.net</a> , with the default to plot all statistics.
targ.line	If TRUE, plot target or expected value line for the statistic of interest.
targ.col	Vector of standard R colors for target statistic lines, with default colors based on RColorBrewer color palettes.
targ.lwd	Line width for the line showing the target statistic values.
targ.lty	Line type for the line showing the target statistic values.
plots.joined	If TRUE and type="formation", combine all target statistics in one plot, versus one plot per target statistic if FALSE.
...	additional arguments to pass.

## Details

This plot function can produce three types of plots with a stochastic network model simulated through `netsim`:

1. `type="epi"`: epidemic model results (e.g., disease prevalence and incidence) may be plotted.
2. `type="network"`: a static network plot will be generated. A static network plot of a dynamic network is a cross-sectional extraction of that dynamic network at a specific time point. This plotting function wraps the `plot.network` function in the `network` package. Consult the help page for `plot.network` for all of the plotting parameters. In addition, four plotting parameters specific to `netsim` plots are available: `sim`, `at`, `col.status`, and `shp.bip`.
3. `type="formation"`: summary network statistics related to the network model formation are plotted. These plots are similar to the formation plots for `netdx` objects. When running a `netsim` simulation, one must specify there that `save.nwstats=TRUE`; the plot here will then show the network statistics requested explicitly in `nwstats.formula`, or will use the formation formula set in `netest` otherwise.

When `type="epi"`, this plotting function will extract the epidemiological output from a model object of class `netsim` and plot the time series data of disease prevalence and other results. The summary statistics that the function calculates and plots are individual simulation lines, means of the individual simulation lines, and quantiles of those individual simulation lines. The mean line, toggled on with `mean.line=TRUE` is calculated as the row mean across simulations at each time step.

Compartment prevalences are the size of a compartment over some denominator. To plot the raw numbers from any compartment, use `popfrac=FALSE`; this is the default for any plots of flows. The `popfrac` parameter calculates and plots the denominators of all specified compartments using these rules: 1) for one-group models, the prevalence of any compartment is the compartment size divided by the total population size; 2) for two-group models, the prevalence of any compartment is the compartment size divided by the group population size. For any prevalences that are not automatically calculated, the `mutate_epi` may be used to add new variables to the `netsim` object to plot or analyze.

The quantiles show the range of outcome values within a certain specified quantile range. By default, the interquartile range is shown: that is the middle 50% of the data. This is specified by `qnts=0.5`. To show the middle 95% of the data, specify `qnts=0.95`. To toggle off the polygons where they are plotted by default, specify `qnts=FALSE`.

When `type="network"`, this function will plot cross sections of the simulated networks at specified time steps. Because it is only possible to plot one time step from one simulation at a time, it is necessary to enter these in the `at` and `sims` parameters. To aide in visualizing representative and extreme simulations at specific time steps, the `sims` parameter may be set to `"mean"` to plot the simulation in which the disease prevalence is closest to the average across all simulations, `"min"` to plot the simulation in which the prevalence is lowest, and `"max"` to plot the simulation in which the prevalence is highest.

## See Also

`plot.network`, `mutate_epi`

## Examples

```
## Independent SI Model
# Initialize network and set network model parameters
nw <- network.initialize(n = 100, bipartite = 50, directed = FALSE)
formation <- ~edges
target.stats <- 50
coef.diss <- dissolution_coefs(dissolution = ~offset(edges), duration = 20)

# Estimate the network model
est <- netest(nw, formation, target.stats, coef.diss, verbose = FALSE)

# Simulate the epidemic model
param <- param.net(inf.prob = 0.3, inf.prob.m2 = 0.15)
init <- init.net(i.num = 10, i.num.m2 = 10)
control <- control.net(type = "SI", nsteps = 20, nsims = 3,
                      verbose = FALSE, save.nwstats = TRUE,
                      nwstats.formula = ~edges + meandeg + concurrent)
mod <- netsim(est, param, init, control)

# Plot epidemic trajectory
plot(mod)
plot(mod, type = "epi", grid = TRUE)
plot(mod, type = "epi", popfrac = TRUE)
plot(mod, type = "epi", y = "si.flow", qnts = 1, ylim = c(0, 4))

# Plot static networks
par(mar = c(0,0,0,0))
plot(mod, type = "network")

# Automatic coloring of infected nodes as red
par(mfrow = c(1, 2), mar = c(0, 0, 2, 0))
plot(mod, type = "network", main = "Min Prev | Time 50",
     col.status = TRUE, at = 20, sims = "min")
plot(mod, type = "network", main = "Max Prev | Time 50",
     col.status = TRUE, at = 20, sims = "max")

# Automatic shape by mode number (circle = mode 1)
par(mar = c(0,0,0,0))
plot(mod, type = "network", at = 20, col.status = TRUE, shp.bip = "square")
plot(mod, type = "network", at = 20, col.status = TRUE, shp.bip = "triangle")

# Plot formation statistics
par(mfrow = c(1,1), mar = c(3,3,1,1), mgp = c(2,1,0))
plot(mod, type = "formation", grid = TRUE)
plot(mod, type = "formation", plots.joined = FALSE)
plot(mod, type = "formation", sims = 2:3)
plot(mod, type = "formation", plots.joined = FALSE,
     stats = c("edges", "concurrent"))
plot(mod, type = "formation", stats = "meandeg",
     mean.lwd = 1, qnts.col = "seagreen", mean.col = "black")
```

---

plot.transmat	<i>Plot transmat infection tree in one of several styles</i>
---------------	--

---

### Description

Plots the infection tree described in a [transmat](#) object in one of several styles: phylogenetic tree, an un-rooted network, a hierarchical tree, or a transmissionTimeline.

### Usage

```
## S3 method for class 'transmat'
plot(x, style = c("phylo", "network", "transmissionTimeline"), ...)
```

### Arguments

x	A <a href="#">transmat</a> object to be plotted
style	Character name of plot style. One of "phylo", "network", or "transmissionTimeline"
...	Additional plot arguments to be passed to lower-level plot functions (plot.network, plot.phylo, etc)

### Details

The phylo plot requires the ape package. The ndtv::transmissionTimeline requires that the ndtv package is installed. All of the options are essentially wrappers to other plot calls with some appropriate preset arguments.

### See Also

[plot.network](#), [plot.phylo](#)

---

summary.dcm	<i>Summary Model Statistics</i>
-------------	---------------------------------

---

### Description

Extracts and prints model statistics solved with dcm.

### Usage

```
## S3 method for class 'dcm'
summary(object, at, run = 1, digits = 3, ...)
```

Arguments

object	An EpiModel object of class dcm.
at	Time step for model statistics.
run	Model run number, for dcm class models with multiple runs (sensitivity analyses).
digits	Number of significant digits to print.
...	Additional summary function arguments (not used).

Details

Summary statistics for the main epidemiological outcomes (state and transition size and prevalence) from an dcm model. Time-specific summary measures are provided, so it is necessary to input a time of interest. For multiple-run models (sensitivity analyses), input a model run number. See examples below.

See Also

[dcm](#)

Examples

```
## Deterministic SIR model with varying act.rate
param <- param.dcm(inf.prob = 0.2, act.rate = 2:4, rec.rate = 1/3,
                  a.rate = 0.011, ds.rate = 0.01,
                  di.rate = 0.03, dr.rate = 0.01)
init <- init.dcm(s.num = 1000, i.num = 1, r.num = 0)
control <- control.dcm(type = "SIR", nsteps = 50)
mod <- dcm(param, init, control)
summary(mod, at = 25, run = 1)
summary(mod, at = 25, run = 3)
summary(mod, at = 26, run = 3)
```

---

summary.icm	<i>Summary Model Statistics</i>
-------------	---------------------------------

---

Description

Extracts and prints model statistics simulated with icm.

Usage

```
## S3 method for class 'icm'
summary(object, at, digits = 3, ...)
```

**Arguments**

object	An EpiModel object of class icm.
at	Time step for model statistics.
digits	Number of significant digits to print.
...	Additional summary function arguments.

**Details**

Summary statistics for the main epidemiological outcomes (state and transition size and prevalence) from an icm model. Time-specific summary measures are provided, so it is necessary to input a time of interest.

**See Also**

[icm](#)

**Examples**

```
## Stochastic ICM SI model with 3 simulations
param <- param.icm(inf.prob = 0.2, act.rate = 1)
init <- init.icm(s.num = 500, i.num = 1)
control <- control.icm(type = "SI", nsteps = 50,
                      nsims = 5, verbose = FALSE)
mod <- icm(param, init, control)
summary(mod, at = 25)
summary(mod, at = 50)
```

---

summary.netsim	<i>Summary Model Statistics</i>
----------------	---------------------------------

---

**Description**

Extracts and prints model statistics simulated with netsim.

**Usage**

```
## S3 method for class 'netsim'
summary(object, at, digits = 3, ...)
```

**Arguments**

object	An EpiModel object of class netsim.
at	Time step for model statistics.
digits	Number of significant digits to print.
...	Additional summary function arguments.

## Details

Summary statistics for the main epidemiological outcomes (state and transition size and prevalence) from an `netsim` model. Time-specific summary measures are provided, so it is necessary to input a time of interest.

## See Also

[netsim](#)

## Examples

```
## Not run:
## Independent SI Model
# Initialize network and set network model parameters
nw <- network.initialize(n = 100, bipartite = 50, directed = FALSE)
formation <- ~edges
target.stats <- 50
coef.diss <- dissolution_coefs(dissolution = ~offset(edges), duration = 20)

# Estimate the ERGM models (see help for netest)
est1 <- netest(nw, formation, target.stats, coef.diss, verbose = FALSE)

# Parameters, initial conditions, and controls for model
param <- param.net(inf.prob = 0.3, inf.prob.m2 = 0.15)
init <- init.net(i.num = 10, i.num.m2 = 10)
control <- control.net(type = "SI", nsteps = 100, nsims = 5, verbose.int = 0)

# Run the model simulation
mod <- netsim(est1, param, init, control)

summary(mod, at = 1)
summary(mod, at = 50)
summary(mod, at = 100)

## End(Not run)
```

---

| truncate\_sim |

---

| *Truncate Simulation Time Series* |

---

## Description

Left-truncates a simulation epidemiological summary statistics and network statistics at a specified time step.

## Usage

```
truncate_sim(x, at)
```



**Arguments**

`x`                      Object of class `netsim` or `icm`.  
`at`                      Time step at which to left-truncate the time series.

**Details**

This function would be used when running a follow-up simulation from time steps `b` to `c` after a burnin period from time `a` to `b`, where the final time window of interest for data analysis is `b` to `c` only.

**Examples**

```
param <- param.icm(inf.prob = 0.2, act.rate = 0.25)
init <- init.icm(s.num = 500, i.num = 1)
control <- control.icm(type = "SI", nsteps = 200, nsims = 1)
mod1 <- icm(param, init, control)
df <- as.data.frame(mod1)
print(df)
plot(mod1)
mod1$control$nsteps

mod2 <- truncate_sim(mod1, at = 150)
df2 <- as.data.frame(mod2)
print(df2)
plot(mod2)
mod2$control$nsteps
```

---

update_dissolution	<i>Adjust Dissolution Component of Network Model Fit</i>
--------------------	--

---

**Description**

Adjusts the dissolution component of an dynamic ERGM fit using the `netest` function with the edges dissolution approximation method.

**Usage**

```
update_dissolution(old.netest, new.coef.diss)
```

**Arguments**

`old.netest`            An object of class `netest`, from the [netest](#) function.  
`new.coef.diss`        An object of class `disscoef`, from the [dissolution\\_coefs](#) function.

## Details

Fitting an ERGM is a computationally intensive process when the model includes dyadic dependent terms. With the edges dissolution approximation method of Carnegie et al, the coefficients for a temporal ERGM are approximated by fitting a static ERGM and adjusting the formation coefficients to account for edge dissolution. This function provides a very efficient method to adjust the coefficients of that model when one wants to use a different dissolution model; a typical use case may be to fit several different models with different average edge durations as targets. The example below exhibits that case.

## Examples

```
## Not run:
nw <- network.initialize(1000, directed = FALSE)

# Two dissolutions: an average duration of 300 versus 200
diss.300 <- dissolution_coefs(~offset(edges), 300, 0.001)
diss.200 <- dissolution_coefs(~offset(edges), 200, 0.001)

# Fit the two reference models
est300 <- netest(nw = nw,
                 formation = ~edges,
                 target.stats = c(500),
                 coef.diss = diss.300)

est200 <- netest(nw = nw,
                 formation = ~edges,
                 target.stats = c(500),
                 coef.diss = diss.200)

# Alternatively, update the 300 model with the 200 coefficients
est200.compare <- update_dissolution(est300, diss.200)

identical(est200$coef.form, est200.compare$coef.form)

## End(Not run)
```

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