Modeling valued networks with statnet

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1 Getting the software

If you have not already done so, please make sure that you have a reasonably new version of R, preferably the latest (3.0.0) (R Core Team, 2013). Then, download and install the latest versions of the Statnet (Handcock et~al., 2008; Goodreau et~al., 2008) packages, in particular ergm version 3.1 (Hunter et~al., 2008; Handcock et~al., 2013), ergm.count version 3.1 (Krivitsky, 2013), latentnet version 2.4-4 (Krivitsky and Handcock, 2013), and their dependencies. You can accomplish this by typing:

```
> install.packages("ergm.count")
> install.packages("latentnet")

> library(ergm.count)
> library(latentnet)

We will also need a temporary workaround for a bug:
> as.mcmc.default <- coda:::as.mcmc.default</pre>
```

> as.mcmc.list.default <- coda:::as.mcmc.list.default</pre>

2 network and edge attributes

network (Butts, 2008; Butts et~al., March 15, 2013) objects have three types of attributes:

network attributes attributes which pertain to the whole network and include such information as network size, directedness, and multiplicity;

vertex attributes attributes which pertain to the individual vertices in the network and include such information as vertex label, as well as group assignment or some other property of the individual being represented;

edge attributes attributes which pertain to edges in the network and include such information as edge value.

An edge attribute is only defined for edges that exist in the network. Thus, in a matter of speaking, to set an edge value, one first has to *create* an edge and then *set* its attribute.

As with network and vertex attributes, edge attributes that have been set can be listed with list.edge.attributes. Every network has at least one edge attribute: "na", which, if set to TRUE, marks an edge as missing.

2.1 Constructing valued networks

2.1.1 Sampson's Monks, pooled

The first dataset that we'll be using is the (in)famous Sampson's monks. Dataset samplk in package ergm contains three (binary) networks: samplk1, samplk2, and samplk3, containing the Monks' top-tree friendship nominations at each of the three survey time points. We are going to construct a valued network that pools these nominations.

Method 1: From a sociomatrix Suppose that a valued sociomatrix is available.

```
> data(samplk)
> ls()
[1] "as.mcmc.default"
                            "as.mcmc.list.default" "samplk1"
                            "samplk3"
[4] "samplk2"
> as.matrix(samplk1)[1:5,1:5]
            Ramuald Bonaventure Ambrose Berthold Peter
Ramuald
Bonaventure
                   0
                               0
                                        0
                                                 0
                                                       1
Ambrose
                   0
                                        0
                                                 0
                                                       0
                               1
Berthold
                   0
                               0
                                        1
                                                 0
                                                       1
                   1
                               1
                                        0
                                                 1
                                                       0
Peter
> # A sociomatrix totaling the nominations.
> samplk.tot.m<-as.matrix(samplk1)+as.matrix(samplk2)+as.matrix(samplk3)
> samplk.tot.m[1:5,1:5]
            Ramuald Bonaventure Ambrose Berthold Peter
Ramuald
                   0
Bonaventure
                   0
                               0
                                        1
                                                 0
                                                       3
Ambrose
                   0
                               3
                                        0
                                                 0
                                                       0
                                        3
                                                       3
Berthold
                   0
                               1
                                                 0
                               3
                                                 3
Peter
                                                       0
> # Create a network where the number of nominations becomes an attribute of an edge.
> samplk.tot <- as.network(samplk.tot.m, directed=TRUE, matrix.type="a",</pre>
                              ignore.eval=FALSE, names.eval="nominations" # Important!
> # Add vertex attributes.
> samplk.tot %v% "group" <- samplk1 %v% "group" # Groups identified by Sampson
> samplk.tot %v% "group"
 [1] "Waverers" "Loyal"
                            "Loyal"
                                        "Loyal"
                                                   "Loyal"
                                                               "Loyal"
 [7] "Waverers" "Turks"
                            "Turks"
                                        "Turks"
                                                   "Turks"
                                                               "Turks"
[13] "Turks"
                "Turks"
                            "Waverers" "Outcasts" "Outcasts" "Outcasts"
```

```
> samplk.tot %v% "vertex.names" <- samplk1 %v% "vertex.names" # Names
> # We can view the attribute as a sociomatrix.
> as.matrix(samplk.tot,attrname="nominations")[1:5,1:5]
            Ramuald Bonaventure Ambrose Berthold Peter
Ramuald
                              2
                                                0
                                      1
Bonaventure
                  0
                              0
                                       1
                                                0
                                                      3
                              3
                                                      0
Ambrose
                  0
                                      0
                                                0
Berthold
                  0
                              1
                                       3
                                                0
                                                      3
Peter
                  1
                              3
                                       0
                                                3
                                                      0
> # Also, note that samplk.tot now has an edge if i nominated j *at least once*.
> as.matrix(samplk.tot)[1:5,1:5]
            Ramuald Bonaventure Ambrose Berthold Peter
Ramuald
                                       1
                                                0
                              1
Bonaventure
                  0
                               0
                                                0
Ambrose
                              1
                                       0
                                                0
                  0
Berthold
                  0
                              1
                                      1
                                                0
                                                      1
Peter
                  1
                              1
                                     0
                                                1
                                                      0
Method 2: From an edgelist Now, suppose that instead we have an edgelist with values:
```

```
> samplk.tot.el <- as.matrix(samplk.tot, attrname="nominations",</pre>
                              matrix.type="edgelist")
> samplk.tot.el[1:5,]
     [,1] [,2] [,3]
[1,]
             1
[2,]
[3,]
             2
[4,]
                   3
        3
[5,]
> # and an initial empty network.
> samplk.tot2 <- samplk1 # Copy samplk1</pre>
> delete.edges(samplk.tot2, seq_along(samplk.tot2$mel)) # Empty it out
> samplk.tot2
 Network attributes:
  vertices = 18
  directed = TRUE
  hyper = FALSE
  loops = FALSE
  multiple = FALSE
  bipartite = FALSE
```

```
total edges= 0
    missing edges= 0
    non-missing edges= 0
 Vertex attribute names:
    cloisterville group vertex.names
> samplk.tot2[samplk.tot.el[,1:2], names.eval="nominations", add.edges=TRUE] <-
    samplk.tot.el[,3]
> as.matrix(samplk.tot2,attrname="nominations")[1:5,1:5]
            Ramuald Bonaventure Ambrose Berthold Peter
Ramuald
                   0
                               2
                                        1
Bonaventure
                   0
                               0
                                        1
                                                  0
                                                        3
Ambrose
                   0
                               3
                                        0
                                                  0
                                                        0
                                1
                                        3
                                                  0
                                                        3
Berthold
                   0
                                3
                                                  3
Peter
                   1
                                        0
                                                        0
```

In general, the construction net[i,j, names.eval="attrname", add.edges=TRUE] <-value can be used to modify individual edge values for attribute "attrname". This way, we can also add more than one edge attribute to a network.

2.1.2 Zachary's Karate club

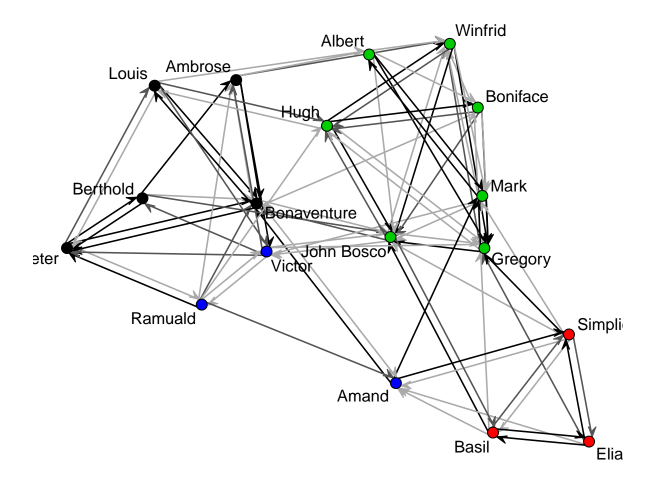
The other dataset we'll be using is almost as (in)famous Zachary's Karate Club dataset is a collapsed multiplex network that counts the number of social contexts in which each pair of individuals associated with the Karate Club in question interacted. A total of 8 contexts were considered, but as the contexts themselves were determined by the network process, this limit itself can be argued to be endogenous.

Over the course of the study, the club split into two factions one led by the instructor, "Mr. Hi" and the other led by the Club President, "John A.". Zachary also recorded the faction alignment of every regular attendee in the club. This dataset is included in the ergm.count package, as zach.

2.2 Visualizing a valued network

The network's plot method for networks can be used to plot a sociogram of a network. When plotting a valued network, we might want to do color the ties depending on their value. Function gray can be used to generate a gradient of colors, with gray(0) generating black and gray(1) generating white. This can then be passed to the edge.col argument of plot.network.

Sampson's Monks For the monks, let's do it as a matrix.

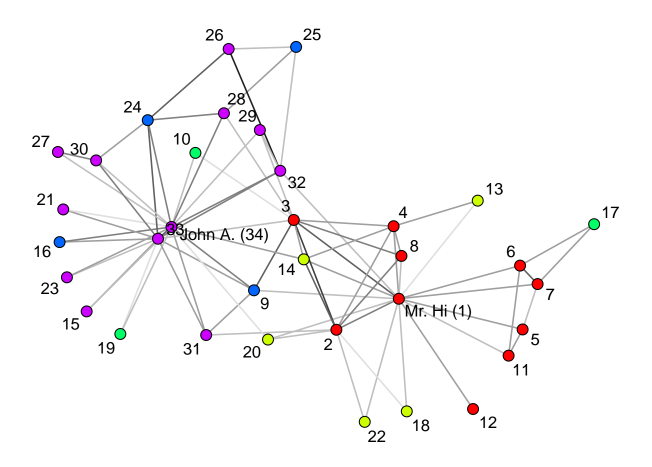


Edge color can also be passed as a vector of colors corresponding to edges. It's more efficient, but the ordering in the vector must correspond to network object's internal ordering, so it should be used with care.

Zachary's Karate Club In the following plot, we plot those strongly aligned with Mr. Hi as red, those with John A. with purple, those neutral as green, and those weakly aligned with

colors in between.

```
> data(zach)
> zach.ecol <- gray(1 - (zach %e% "contexts")/8)
> zach.vcol <- rainbow(5)[zach %v% "faction.id"+3]
> par(mar=rep(0,4))
> plot(zach, edge.col=zach.ecol, vertex.col=zach.vcol, displaylabels=TRUE)
```



3 Modeling dyad-dependent interaction counts with valued ERGMs using ergm.count

Many of the functions in package ergm, including ergm, simulate, and summary, have been extended to handle networks with valued relatins. They switch into this valued mode when passed the response argument, specifying the name of the edge attribute to use as the response variable. For example, a new valued term sum evaluates the sum of the values of all of the relations: $\sum_{(i,j)\in\mathbb{Y}} y_{i,j}$. So,

> summary(samplk.tot~sum)

produces an error (because no such term has been implemented for binary mode), while

> summary(samplk.tot~sum, response="nominations")

sum 168

gives the summary statistics. We will introduce more statistics shortly. First, we need to introduce the notion of valued ERGMs.

For a more in-depth discussion of the following, see (Krivitsky, 2012).

3.1 Valued ERGMs

Generally, a valued ERGM (for discrete variables) looks like this:

$$\mathrm{Pr}_{h,\boldsymbol{g}}(\boldsymbol{Y}=\boldsymbol{y};\boldsymbol{\theta}) = \frac{h(\boldsymbol{y})\exp(\boldsymbol{\theta}^{\top}\boldsymbol{g}(\boldsymbol{y}))}{c_{h,\boldsymbol{g}}(\boldsymbol{\theta})}, \ \boldsymbol{y} \in \mathcal{Y},$$

where the normalizing constant

$$c_{h,\boldsymbol{g}}(\boldsymbol{\theta}) = \sum_{\boldsymbol{y} \in \mathcal{Y}} h(\boldsymbol{y}) \exp\left(\boldsymbol{\theta}^{\top} \boldsymbol{g}(\boldsymbol{y})\right).$$

3.1.1 New concept: a reference distribution

With binary ERGMs, we only concern ourselves with presence and absence of ties among actors — who is connected with whom? If we want to model values as well, we need to think about who is connected with whom and how strong or intense these connections are. In particular, we need to think about how the values for connections we measure are distributed. The reference distribution (a reference measure, for the mathematically inclined) specifies the model for the data before we add any ERGM terms, and is the first step in modeling these values. The reference distribution is specified using a one-sided formula as a reference argument to an ergm or simulate call. Running

> help("ergm-references")

will list the choices implemented in the various packages, and are given as a one-sided formula.

Conceptually, it has two ingredients: the sample space and the baseline distribution (h(y)). An ERGM that "borrows" these from a distribution X for which we have a name is called an X-reference ERGM.

The sample space For binary ERGMs, the sample space \mathcal{Y} — the set of possible networks that can occur — is some subset of $2^{\mathbb{Y}}$, the set of all possible ways in which relationships among the actors may occur.

For the sample space of valued ERGMs, we need to define \mathbb{S} , the set of possible values each relationship may take. For example, for count data, that's $\mathbb{S} = \{0, 1, ..., s\}$ if the maximum count is s and $\{0, 1, ...\}$ if there is no a priori upper bound. Having specified that, \mathcal{Y} is defined as some subset of $\mathbb{S}^{\mathbb{Y}}$: the set of possible ways to assign to each relationship a value.

As with binary ERGMs, other constraints like degree distribution may be imposed on \mathcal{Y} .

h(y): The baseline distribution What difference does it make?

Suppose that we have a sample space with $\mathbb{S} = \{0, 1, 2, 3\}$ (e.g., number of monk-monk nominations) and let's have one ERGM term: the sum of values of all relations: $\sum_{(i,j)\in\mathbb{Y}} y_{i,j}$:

$$\operatorname{Pr}_{h,oldsymbol{g}}(oldsymbol{Y}=oldsymbol{y};oldsymbol{ heta}) \propto h(oldsymbol{y}) \exp\left(oldsymbol{ heta} \sum_{(i,j) \in \mathbb{Y}} oldsymbol{y}_{i,j}
ight).$$

There are two values for h(y) that might be familiar:

$$h(y) = 1$$
 (or any constant) $\implies Y_{i,j} \stackrel{\text{i.i.d}}{\sim} \text{Uniform or truncated geometric}$

$$h(\boldsymbol{y}) = \binom{m}{\boldsymbol{y}_{i,j}} = \frac{m!}{\boldsymbol{y}_{i,j}!(m-\boldsymbol{y}_{i,j})!} \implies \boldsymbol{Y}_{i,j} \overset{\text{i.i.d.}}{\sim} \text{Binomial}(m, \text{logit}^{-1}(\boldsymbol{\theta}))$$

What do they look like? Let's simulate!

```
> y <- network.initialize(2,directed=FALSE) # A network with one dyad!
> ## Discrete Uniform reference
> # 0 coefficient: discrete uniform
> sim.du3<-simulate(y~sum, coef=0, reference=~DiscUnif(0,3),
                    response="w",statsonly=TRUE,nsim=1000)
> # Negative coefficient: truncated geometric skewed to the right
> sim.trgeo.m1<-simulate(y~sum, coef=-1, reference=~DiscUnif(0,3),</pre>
                         response="w",statsonly=TRUE,nsim=1000)
> # Positive coefficient: truncated geometric skewed to the left
> sim.trgeo.p1<-simulate(y~sum, coef=+1, reference=~DiscUnif(0,3),</pre>
                        response="w",statsonly=TRUE,nsim=1000)
> # Plot them:
> par(mfrow=c(1,3))
> hist(sim.du3,breaks=diff(range(sim.du3))*4)
> hist(sim.trgeo.m1,breaks=diff(range(sim.trgeo.m1))*4)
> hist(sim.trgeo.p1,breaks=diff(range(sim.trgeo.p1))*4)
```

Histogram of sim.du3 Histogram of sim.trgeo.m1 Histogram of sim.trgeo.p1 250 500 200 500 -requency -requency 150 300 300 100 50 100 100

0.0

1.0

sim.trgeo.p1

2.0

3.0

0.0

1.0

sim.trgeo.m1

2.0

3.0

-requency

0.0

1.0

sim.du3

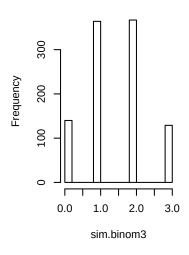
2.0

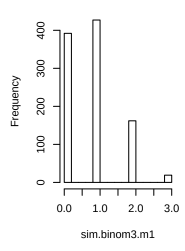
3.0

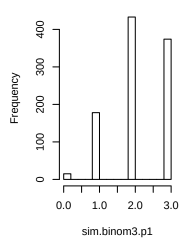
Histogram of sim.binom3

Histogram of sim.binom3.m

Histogram of sim.binom3.p







Now, suppose that we don't have an a priori upper bound on the counts — $\mathbb{S} = \{0, 1, \dots\}$ — then there are two familiar reference distributions:

$$h(y) = 1$$
 (or any constant) $\Longrightarrow Y_{i,j} \stackrel{\text{i.i.d.}}{\sim} \text{Geometric}(p = 1 - \exp(\theta))$

$$h(\boldsymbol{y}) = 1/\prod_{(i,j)\in\mathbb{Y}} \boldsymbol{y}_{i,j}! \implies \boldsymbol{Y}_{i,j} \overset{\text{i.i.d.}}{\sim} \text{Poisson}(\mu = \exp(\boldsymbol{\theta}))$$

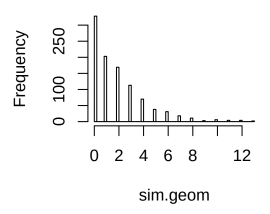
- > mean(sim.geom)
- [1] 1.955
- > mean(sim.pois)
- [1] 2.024

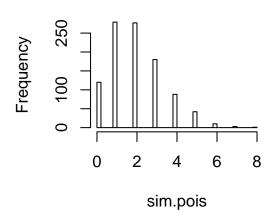
Similar means. But, what do they look like?

- > par(mfrow=c(1,2))
- > hist(sim.geom,breaks=diff(range(sim.geom))*4)
- > hist(sim.pois,breaks=diff(range(sim.pois))*4)

Histogram of sim.geom

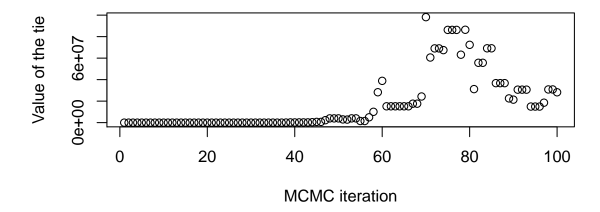
Histogram of sim.pois





Where did log(2) and log(2/3) come from? Later.

Warning: Parameter space What happens if we simulate from a geometric-reference ERGM with all coefficients set to 0?



Why does it do that? Because

$$\mathrm{Pr}_{h,\boldsymbol{g}}(\boldsymbol{Y}=\boldsymbol{y};\boldsymbol{\theta}) = \frac{\exp(\boldsymbol{\theta} \sum_{(i,j) \in \mathbb{Y}} \boldsymbol{y}_{i,j})}{c_{h,\boldsymbol{g}}(\boldsymbol{\theta})}$$

for $\theta \geq 0$, is not a valid distribution, because $c_{h,g}(\theta) = \infty$. Using reference="Geometric can be dangerous for this reason. This issue only arises with ERGMs that have an infinite sample space.

3.2 Initial values

Before we can start fitting these models, we need a place to start. Unfortunately, there is no general MPLE for these models that we have (yet), so there are two options:

0 This is the default. It can work reasonably well for some models, but it can also bog down for larger networks.

A closed-form submodel If the model fit has a simpler submodel that we can estimate exactly, we can use that as a starting point. That is, we find a coefficient for the submodel, and then start the main fit with all other coefficients set to 0.

Just as almost ever application of binary ERGMs uses the edges term as the baseline, almost every model we are likely to use will have the baseline intensity (sum) term. Given a network y with attribute "a" to be used as the response, the following should give adequate starting coefficients for sum for:

Bernoulli: Binary ERGM. Use MPLE.

Binomial(trials):

```
p <- sum(y %e% "a")/trials/network.dyadcount(y) init.sum <- \log(p/(1-p))
```

Poisson: m <- sum(y %e%)

```
m <- sum(y %e% "a")/network.dyadcount(y)
init.sum <- log(m)</pre>
```

Geometric:

```
m <- sum(y %e% "a")/network.dyadcount(y)
init.sum <- log(1-1/(m+1))</pre>
```

This will probably be automated in the next version.

3.2.1 Example: Sampson's Monks with Binomial(3) reference

Suppose our model of interest contains the term sum, and we are using the Binomial(3) reference distribution. Then, plugging in samplk.tot for y and "nominations" for "a",

```
> p <- sum(samplk.tot %e% "nominations")/3/network.dyadcount(samplk.tot)
> p
[1] 0.1830065
> samplk.sum.init <- log(p/(1-p)) # i.e., logit(p)
> samplk.sum.init
[1] -1.496109
```

3.2.2 Example: Zachary's Karate Club with Poisson reference

Suppose our model of interest contains the term sum, and we are using the Poisson reference distribution. Then, plugging in zach for y and "contexts" for "a",

```
> m <- sum(zach %e% "contexts")/network.dyadcount(zach)
> m

[1] 0.4117647

> zach.sum.init <- log(m)
> zach.sum.init

[1] -0.8873032
```

We will be using these in the following examples.

3.3 ERGM terms

3.3.1 GLM-style terms

Many of the familiar ERGM effects can be modeled using the very same terms, but applied a little differently.

Any dyad-independent binary ERGM statistic can be expressed as $\mathbf{g}_{k} = \sum_{(i,j) \in \mathbb{Y}} \mathbf{x}_{k,i,j} \mathbf{y}_{i,j}$ for some covariate matrix \mathbf{x}_{k} . If $\mathbf{y}_{i,j}$ is allowed to have values other than 0 and 1, then simply using such a term in a Poisson-reference ERGM creates the familiar log-linear effect. Similarly, in a Binomial-reference ERGM, such terms produce an effect on log-odds of a success.

The good news is that almost every dyad-independent ergm term has been reimplemented to allow this. It is invoked by specifying "form="sum" argument for one of the terms inherited from binary ERGMs, though this not required, as it's the default. Also, note that for valued ERGMs, the "intercept" term is sum, not edges.

```
> help("ergm-terms")
```

has the complete list.

Example: Sampson's Monks For example, we can fit the equivalent of logistic regression on the probability of nomination, with every ordered pair of monks observed 3 times. We will look at differential homophily on group. That is, $Y_{i,j} \stackrel{\text{ind}}{\sim} \text{Binomial}(3, \pi_{i,j})$ where

```
\begin{aligned} \text{logit}(\pmb{\pi}_{i,j}) &= \pmb{\beta}_1 + \pmb{\beta}_2 \mathbb{I} \left( i \text{ and } j \text{ are both in the Loyal Opposition} \right) \\ &+ \pmb{\beta}_3 \mathbb{I} \left( i \text{ and } j \text{ are both Outcasts} \right) + \pmb{\beta}_4 \mathbb{I} \left( i \text{ and } j \text{ are both Young Turks} \right) \\ &+ \pmb{\beta}_5 \mathbb{I} \left( i \text{ and } j \text{ are both Waverers} \right) \end{aligned}
```

To actually fit it, we need to specify the initial parameters. How many do we need?

Monte Carlo MLE Results:

```
Estimate Std. Error MCMC % p-value
                              -2.3253
                                          0.1346
                                                      0 <1e-04 ***
sum
nodematch.sum.group.Loyal
                               2.2553
                                          0.2940
                                                      0 <1e-04 ***
nodematch.sum.group.Outcasts
                               3.5661
                                          0.5886
                                                      0 <1e-04 ***
                               2.1999
                                                      0 <1e-04 ***
nodematch.sum.group.Turks
                                          0.2202
nodematch.sum.group.Waverers
                               1.0732
                                          0.5813
                                                      0 0.0658 .
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
     Null Deviance:
                      0.0 on 306 degrees of freedom
 Residual Deviance: -555.5 on 301 degrees of freedom
Note that the null model likelihood and deviance are defined to be 0.
AIC: -545.5
               BIC: -526.8
                              (Smaller is better.)
```

Based on this, we can say that the odds of a monk nominating another monk not in the same group during a given time step are $\exp(\beta_1) = \exp(-2.3253) = 0.0978$, that the odds of a Loyal Opposition monk nominating another Loyal Opposition monk are $\exp(\beta_2) = \exp(2.2553) = 9.5379$ times higher, etc..

Example: Zachary's Karate Club We will use a Poisson log-linear model for the number of contexts in which each pair of individuals interacted, as a function of whether this individual is a faction leader (Mr. Hi or John A.) That is, $Y_{i,j} \stackrel{\text{ind.}}{\sim} \text{Poisson}(\mu_{i,j})$ where

```
\log(\pmb{\mu}_{i.i}) = \pmb{\beta}_1 + \pmb{\beta}_2(\mathbb{I}\left(i \text{ is a faction leader}\right) + \mathbb{I}\left(j \text{ is a faction leader}\right))
```

We will do this by constructing a dummy variable, a vertex attribute "leader":

```
_____
Summary of model fit
          zach ~ sum + nodefactor("leader")
Formula:
Iterations: 20
Monte Carlo MLE Results:
                         Estimate Std. Error MCMC % p-value
                          -1.2204
                                     0.0799
                                                0 <1e-04 ***
                           1.4396
                                     0.1176
                                                0 <1e-04 ***
nodefactor.sum.leader.TRUE
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
    Null Deviance: 0.0 on 561 degrees of freedom
Residual Deviance: -349.9 on 559 degrees of freedom
Note that the null model likelihood and deviance are defined to be 0.
AIC: -345.9
             BIC: -337.2
                            (Smaller is better.)
```

Based on this, we can say that the expected number of contexts of interaction between two non-leaders is $\exp(\beta_1) = \exp(-1.2204) = 0.2951$, that the expected number of contexts of interaction between a leader and a non-leader is $\exp(\beta_2) = \exp(1.4396) = 4.2192$ times higher, and that the expected number of contexts of interaction between the two leaders is $\exp(2\beta_2) = \exp(2 \cdot 1.4396) = 17.8016$ times higher than that between two non-leaders. (Because the leaders were hostile to each other, this may not be a very good prediction.)

3.3.2 Sparsity and zero-modification

It is often the case that in networks of counts, the network is sparse, yet if two actors do interact, their interaction count is relatively high. This amounts to zero-inflation.

We can model this using the binary-ERGM-based terms with the term nonzero $(\boldsymbol{g}_k = \sum_{(i,j)\in\mathbb{Y}}\mathbb{I}\left(\boldsymbol{y}_{i,j}\neq 0\right))$ and GLM-style terms with argument form="nonzero": $\boldsymbol{g}_k = \sum_{(i,j)\in\mathbb{Y}}\boldsymbol{x}_{k,i,j}\mathbb{I}\left(\boldsymbol{y}_{i,j}\neq 0\right)$. For example,

```
> samplk.tot.nm.nz <-
    ergm(samplk.tot~sum + nonzero + nodematch("group",diff=TRUE,form="sum"),
        response="nominations", reference=~Binomial(3),
        control=control.ergm(init=c(samplk.sum.init,0, 0,0,0,0)))
> mcmc.diagnostics(samplk.tot.nm.nz)
> summary(samplk.tot.nm.nz)
```

```
_____
```

```
Summary of model fit
```

```
Formula: samplk.tot ~ sum + nonzero + nodematch("group", diff = TRUE,
    form = "sum")
```

Estimate Ctd Essess MCMC % -- --- lue

Iterations: 20

Monte Carlo MLE Results:

	Estimate	Sta. Error	MCMC %	p-value	
sum	-0.3334	0.1968	0	0.0912	
nonzero	-2.9730	0.3220	1	<1e-04	***
nodematch.sum.group.Loyal	1.2191	0.2220	0	<1e-04	***
${\tt nodematch.sum.group.Outcasts}$	1.9709	0.4837	0	<1e-04	***
nodematch.sum.group.Turks	1.1895	0.1779	0	<1e-04	***
${\tt nodematch.sum.group.Waverers}$	0.5998	0.4195	1	0.1538	

```
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

```
Null Deviance: 0.0 on 306 degrees of freedom Residual Deviance: -646.3 on 300 degrees of freedom
```

Note that the null model likelihood and deviance are defined to be 0.

```
AIC: -634.3 BIC: -612 (Smaller is better.)
```

fits a zero-modified Binomial model, with a coefficient on the number of non-zero relations -2.973 is negative and highly significant, indicating that there is an excess of zeros in the data relative to the binomial distribution, and given the rest of the model.

3.3.3 Dispersion

Similarly, even if we may use Poisson as a starting distribution, the counts might be overdispersed or underdispersed relative to it. For now, ergm offers two ways to do so:

Conway-Maxwell-Poisson

- Implemented by adding a CMP term to a Poisson- or geometric-reference ERGM.
- Effectively replaces the "1/y!" part of a Poisson density with " $1/(y!)^{\theta_{\text{CMP}}}$ ".
- + Produces a continuum between a geometric distribution and a Bernoulli distribution.
- + Can represent both over- and under-dispersion.
- Has the parameter space problem; also, has some theoretical issues.

Fractional moments

- Implemented by adding a sum(pow=1/2) term to a Poisson-reference ERGM.
- Adds a statistic of the form $\sum_{(i,j)\in\mathbb{Y}} y_{i,j}^{1/2}$ to the model.
- + More stable.
- + For Poisson-like data, $\sqrt{y_{i,j}}$ is a variance-stabilizing transformation, so it could be interpreted as modeling (along with sum the first two moments of $\sqrt{y_{i,j}}$.
- Not well-understood.
- In extreme cases, creates a bimodal shape in the counts.

3.3.4 Mutuality

ergm binary mutuality statistic has the form $g_{\leftrightarrow} = \sum_{(i,j) \in \mathbb{Y}} y_{i,j} y_{j,i}$. It turns out that directly plugging counts into that statistic is a bad idea. mutuality(form) is a valued ERGM term, permitting the following generalizations:

"geometric": $\sum_{(i,j)\in\mathbb{Y}}\sqrt{y_{i,j}y_{j,i}}$ — can be viewed as uncentered covariance of variance-stabilized counts

"min": $\sum_{(i,j) \in \mathbb{Y}} \min oldsymbol{y}_{i,j}, oldsymbol{y}_{j,i}$ — easiest to interpret

"nabsdiff":
$$\sum_{(i,j)\in\mathbb{Y}} -|oldsymbol{y}_{i,j}-oldsymbol{y}_{j,i}|$$

Figure 1 visualizes their effects.

3.3.5 Individual heterogeneity

Different actors may have different overall propensities to interact. This has been modeled using random effects (as in the p_2 model and using degeneracy-prone terms like k-star counts.

ergm implements a number of statistics to model it, but the one that seems to work best so far seems to be

$$\boldsymbol{g}_{\text{actor cov.}}(\boldsymbol{y}) = \sum_{i \in N} \frac{1}{n-2} \sum_{j,k \in \mathbb{Y}_i \wedge j < k} (\sqrt{\boldsymbol{y}_{i,j}} - \overline{\sqrt{\boldsymbol{y}}}) (\sqrt{\boldsymbol{y}_{i,k}} - \overline{\sqrt{\boldsymbol{y}}}),$$

essentially a measure of covariance between the $\sqrt{y_{i,j}}$ s incident on the same actor. The term nodesqrtcovar implements it.

3.3.6 Triadic closure

Finally, to generalize the notion of triadic closure, ergm implements very flexible transitiveweights(twopath, combine, affect) and similar cyclicalweights statistics. The transitive weight statistic has the following general form:

$$oldsymbol{g}_{oldsymbol{v}}(oldsymbol{y}) = \sum_{(i,j) \in \mathbb{Y}} v_{ ext{affect}}\left(oldsymbol{y}_{i,j}, v_{ ext{combine}}\left(v_{2 ext{-path}}(oldsymbol{y}_{i,k}, oldsymbol{y}_{k,j})_{k \in N \setminus \{i,j\}}
ight)
ight),$$

and can be "customized" by varying the three functions

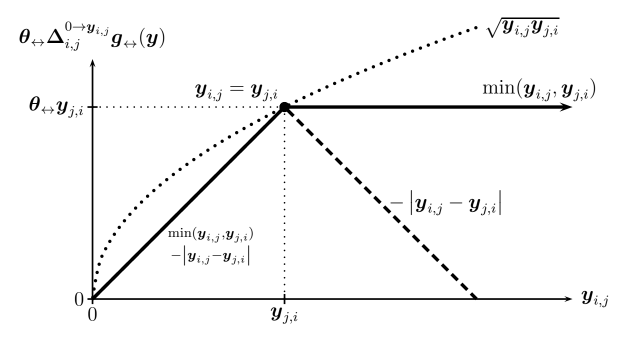


Figure 1: Effect of several mutuality forms on the probability of $Y_{i,j}$ having a certain value given a particular $y_{j,i}$.

 $v_{2\text{-path}}$ Given $y_{i,k}$ and $y_{k,j}$, what is the strength of the two-path they form?

"min" the minimum of their values — conservative

"geomean" their geometric mean — more able to detect effects, but more likely to cause "degeneracy"

 v_{combine} Given the strengths of the two-paths $y_{i\to k\to j}$ for all $k\neq i,j$, what is the combined strength of these two-paths between i and j?

"max" the strength of the strongest path — conservative; analogous to transitiveties
"sum" the sum of path strength — more able to detect effects, but more likely to cause
"degeneracy"; analogous to triangles

 $v_{\mathbf{affect}}$ Given the combined strength of the two-paths between i and j, how should they affect $Y_{i,j}$?

"min" conservative

"geomean" more able to detect effects, but more likely to cause "degeneracy"

These effects are analogous to mutuality.

3.4 Examples

3.4.1 Sampson's Monks

Suppose that we want to fit a model with a zero-modified Binomial baseline, mutuality, transitive (hierarchical) triads, and cyclical (antihierarchical) triads, to this dataset. Such a

```
model has 5 parameters, so
> theta0 <- c(samplk.sum.init, 0, 0, 0, 0)
> samplk.tot.ergm <-</pre>
    ergm(samplk.tot ~ sum + nonzero + mutual("min") +
        transitiveweights("min", "max", "min") +
         cyclicalweights("min","max","min"),
        reference="Binomial(3), response="nominations",
         control=control.ergm(init=theta0))
> mcmc.diagnostics(samplk.tot.ergm)
> summary(samplk.tot.ergm)
  _____
Summary of model fit
samplk.tot ~ sum + nonzero + mutual("min") + transitiveweights("min",
    "max", "min") + cyclicalweights("min", "max", "min")
Iterations: 20
Monte Carlo MLE Results:
                             Estimate Std. Error MCMC % p-value
                             -0.04967 0.18854
                                                     2 0.7924
sum
                             -3.45575
                                         0.31520
                                                     4 <1e-04 ***
nonzero
                                                     3 <1e-04 ***
mutual.min
                              1.38499
                                         0.24720
transitiveweights.min.max.min 0.21051
                                         0.16318
                                                     0 0.1980
cyclicalweights.min.max.min
                                                     1 0.0884 .
                             -0.24760
                                         0.14486
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
    Null Deviance:
                      0 on 306 degrees of freedom
 Residual Deviance: -603 on 301 degrees of freedom
Note that the null model likelihood and deviance are defined to be 0.
AIC: -593
            BIC: -574.4
                           (Smaller is better.)
```

What does it tell us? The negative coefficient on nonzero suggests zero-inflation, there is strong evidence of mutuality, and the positive coefficient on transitive weights and negative on the cyclical weights suggests hierarchy, but they are not significant.

3.4.2 Zachary's Karate Club

Now, let's try using Poisson to model the Zachary Karate Club data: a zero-modified Poisson, with potentially different levels of activity for the faction leaders, heterogeneity in actor

activity level overall, and an effect of difference in faction membership, a model that looks like this:

```
      sum
      nonzero

      231.00000
      78.00000

      nodefactor.sum.leader.TRUE
      absdiff.sum.faction.id.1

      90.00000
      74.00000

      absdiff.sum.faction.id.2
      absdiff.sum.faction.id.3

      12.00000
      11.00000

      absdiff.sum.faction.id.4
      nodesqrtcovar.centered

      10.00000
      16.17248
```

The parameter vector will have length 8, so

```
> theta0 <- c(zach.sum.init, rep(0,7))</pre>
```

A few other notes:

- We can make sampling from zero-inflated Poisson more efficient by telling ergm that that's what we expect by setting MCMC.prop.weights="0inflated" control parameter. The current version has a bug that requires the degree of inflation to be set by the user, though this will be automatic in the next release, so MCMC.prop.args=list(p0=0.5) is also needed.
- Informally, a Poisson random variable contains more "information" than a Bernoulli random variable, which means that large changes in likelihood are not necessarily symptomatic of a problem. Thus, it often helps to set MCMLE.trustregion, which is normally 20 to something higher.

Now, for the fit and the diagnostics:

```
Summary of model fit
```

Formula: zach ~ sum + nonzero + nodefactor("leader") + absdiffcat("faction.id") + nodesqrtcovar(TRUE)

Iterations: 20

Monte Carlo MLE Results:

	Estimate	Std. Error	MCMC %	p-value	
sum	0.99512	0.09250	7	< 1e-04	***
nonzero	-3.85145	0.27512	11	< 1e-04	***
nodefactor.sum.leader.TRUE	0.20062	0.06633	1	0.00261	**
absdiff.sum.faction.id.1	-0.10837	0.07810	0	0.16582	
absdiff.sum.faction.id.2	-0.66743	0.19232	1	0.00056	***
absdiff.sum.faction.id.3	-0.88254	0.21637	1	< 1e-04	***
absdiff.sum.faction.id.4	-1.16067	0.23913	1	< 1e-04	***
nodesqrtcovar.centered	1.81829	0.22266	1	< 1e-04	***

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Null Deviance: 0.0 on 561 degrees of freedom Residual Deviance: -805.4 on 553 degrees of freedom

Note that the null model likelihood and deviance are defined to be 0.

```
AIC: -789.4 BIC: -754.7 (Smaller is better.)
```

What does it tell us? The negative coefficient on nonzero suggests zero-inflation, the faction leaders clearly have more activity than others, and the more ideologically separated two individuals are, the less they interact. Over and above that, there is some additional heterogeneity in how active individuals are: if i has a lot of interaction with j, it is likely that i has more with j'. Could this mean a form of preferential attachment?

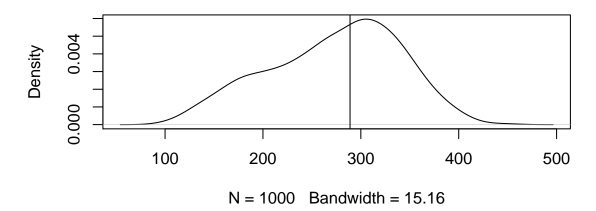
We can try seeing whether there is some friend of a friend effect above and beyond that. This can be done by fitting a model with transitivity and seeing whether the coefficient is significant, or we can perform a simulation test. In the following

- simulate unpacks the zach.pois ERGM fit, extracting the formula, the coefficient, and the rest of the information.
- nsim says how many networks to generate.
- statsonly=TRUE says that we only want to see the simulated statistics, not the networks.
- monitor="transitiveweights("geomean", "sum", "geomean") says that in addition to the statistics used in the fit, we want simulate to keep track of the transitive weights statistic.

We do not need to worry about degeneracy in this case, because we are not actually using that statistic in the model, only "monitoring" it.

```
> # Simulate from model fit:
> zach.sim <-
    simulate(zach.pois, monitor=~transitiveweights("geomean", "sum", "geomean"),
             nsim = 1000, statsonly=TRUE,
             control=control.simulate.ergm(
               MCMC.prop.weights="0inflated",
               MCMC.prop.args=list(p0=0.5) # Should not be necessary in the next version.
               ))
> # What have we simulated?
> colnames(zach.sim)
[1] "sum"
[2] "nonzero"
[3] "nodefactor.sum.leader.TRUE"
[4] "absdiff.sum.faction.id.1"
[5] "absdiff.sum.faction.id.2"
[6] "absdiff.sum.faction.id.3"
[7] "absdiff.sum.faction.id.4"
[8] "nodesqrtcovar.centered"
[9] "transitiveweights.geomean.sum.geomean"
> # How high is the transitiveweights statistic in the observed network?
> zach.obs <- summary(zach ~ transitiveweights("geomean","sum","geomean"),</pre>
                       response="contexts")
> zach.obs
transitiveweights.geomean.sum.geomean
                              288.9793
Let's plot the density of the simulated values of transitive weights statistic:
> par(mar=c(5, 4, 4, 2) + 0.1)
> # 9th col. = transitiveweights
> plot(density(zach.sim[,9]))
> abline(v=zach.obs)
```

density.default(x = zach.sim[, 9])



- > # Where does the observed value lie in the simulated?
- > # This is a p-value for the Monte-Carlo test:
- > min(mean(zach.sim[,9]>zach.obs), mean(zach.sim[,9]<zach.obs))*2</pre>

[1] 0.916

Looks like individual heterogeneity and faction alignment account for appearance of triadic effects. (Notably, the factions themselves may be endogenous, if social influence is a factor. Untangling selection from influence is hard enough when dynamic network data are available. We cannot do it here.)

3.5 Other notes

- Missing (NA) edges are handled automatically for valued ERGMs as well.
- ergm has an argument eval.loglik, which is TRUE by default. For valued ERGMs, it's quite a bit less efficient than for binary, at least for now. So, unless you need the AICs or BICs to compare models, and especially if your networks are not small, pass eval.loglik=FALSE.
- Writing user terms is possible, but the API is a little different from that of binary change statistics API.

3.6 Medium-term Roadmap

- A public release for rank ERGM implementation.
- Automatic starting values for valued ERGMs.
- More efficient sampling for valued ERGMs.

- Goodness of fit diagnostics.
- Other reference distributions, including ranks and continuous data. (Continuous uniform is already implemented, but not well-understood.)

4 Latent space models with non-binary response with latentnet

latentnet, as the name suggests, is designed to fit latent space models, but it can fit other dyad-independent network models as well. Let

Y be the random network being modeled;

y be the observed network;

 \boldsymbol{x} is a $p \times n \times n$ array of dyadic covariates, with

 $\boldsymbol{x}_{\cdot,i,j}$ being a *p*-vector of covariates for dyad (i,j);

 β be the p-vector of covariate coefficients;

Z be the $n \times d$ array of latent positions, with

 Z_i being the d-vector position of actor i;

 $\boldsymbol{\delta}$ be the *n*-vector of sender effects, with

 δ_i being the sender effect of δ ; and

 γ being a *n*-vector of receiver effects, with

 γ_i being the receiver effect of γ .

For brevity, let $\theta = (\beta, \mathbf{Z}, \delta, \gamma)$ and let $\theta_{i,j} = (\beta, \mathbf{Z}_i, \mathbf{Z}_j, \delta_i, \gamma_j)$. Generally, a latent space model that can be fit by latentnet has the following form:

$$\Pr(\mathbf{Y} = \mathbf{y} | \boldsymbol{\theta}, \mathbf{x}) = \prod_{(i,j) \in \mathbb{Y}} \Pr(\mathbf{Y}_{i,j} = \mathbf{y}_{i,j} | \boldsymbol{\theta}_{i,j}, \mathbf{x}_{\cdot,i,j}), \tag{1}$$

$$\Pr(\mathbf{Y}_{i,j} = \mathbf{y}_{i,j} | \boldsymbol{\theta}_{i,j}, \mathbf{x}_{\cdot,i,j}) = f(\mathbf{y}_{i,j} | \boldsymbol{\mu}_{i,j}), \tag{2}$$

$$\boldsymbol{\mu}_{i,j} = g^{-1}(\boldsymbol{\eta}_{i,j}),\tag{3}$$

$$\eta_{i,j} = \boldsymbol{x}_{\cdot,i,j}^{\top} \boldsymbol{\beta} + d(\boldsymbol{Z}_i, \boldsymbol{Z}_j) + \boldsymbol{\delta}_i + \boldsymbol{\gamma}_j,$$
(4)

for some latent position effect $d(\cdot, \cdot)$. Effects supported by latentnet are Euclidean, having $d(\mathbf{Z}_i, \mathbf{Z}_j) = -|\mathbf{Z}_i - \mathbf{Z}_j|$ and bilinear, having $d(\mathbf{Z}_i, \mathbf{Z}_j) = \mathbf{Z}_i^{\top} \mathbf{Z}_j$. Latent space positions \mathbf{Z} can further be modeled as a Gaussian mixture, and $\boldsymbol{\delta}$ and $\boldsymbol{\gamma}$ are likewise modeled as random effects.

In words, the values of individual relations are assumed to be independent (1), and each potential relation has a value modeled by a GLM, having some distribution with density f,

parametrized by its expected value (2), which is, in turn, a function, via a GLM link function g, of the linear predictor $\eta_{i,j}$ (3), which is a linear function combining all the parameters (4).

A latent space model for a binary network has

$$f(\mathbf{y}_{i,j}|\mathbf{\mu}_{i,j}) = (\mathbf{\mu}_{i,j})^{\mathbf{y}_{i,j}} (1 - \mathbf{\mu}_{i,j})^{1 - \mathbf{y}_{i,j}}$$
$$g(\mathbf{\mu}_{i,j}) = \text{logit}(\mathbf{\mu}_{i,j}),$$

a Bernoulli model with a logit link: a logistic regression model.

This is easily extended to any GLM. In particular,

Poisson log-linear model:

$$f(\boldsymbol{y}_{i,j}|\boldsymbol{\mu}_{i,j}) = \exp\left(-\boldsymbol{\mu}_{i,j}\right) \boldsymbol{\mu}_{i,j}^{\boldsymbol{y}_{i,j}}/\boldsymbol{y}_{i,j}!$$
$$g(\boldsymbol{\mu}_{i,j}) = \log(\boldsymbol{\mu}_{i,j})$$

Binomial logit with m trials:

$$f(\boldsymbol{y}_{i,j}|\boldsymbol{\mu}_{i,j}) = {m \choose \boldsymbol{y}_{i,j}} (\boldsymbol{\mu}_{i,j}/m)^{\boldsymbol{y}_{i,j}} (1 - \boldsymbol{\mu}_{i,j}/m)^{m-\boldsymbol{y}_{i,j}}$$
$$g(\boldsymbol{\mu}_{i,j}) = \operatorname{logit}(\boldsymbol{\mu}_{i,j})$$

Normal linear with variance σ^2 :

$$f(\boldsymbol{y}_{i,j}|\boldsymbol{\mu}_{i,j}) = \frac{1}{\sqrt{2\pi\sigma^2}} \exp\left(-\frac{(\boldsymbol{y}_{i,j} - \boldsymbol{\mu}_{i,j})^2}{2\sigma^2}\right)$$
$$g(\boldsymbol{\mu}_{i,j}) = \boldsymbol{\mu}_{i,j}$$

These are currently supported by latentnet. Unfortunately, σ^2 cannot be estimated by latentnet at this time.

4.1 A very quick overview of latentnet

The workhorse function ergmm (for Exponential Random Graph Mixed Model) has a syntax very similar to ergm: the model is specified as network ~ term1 + term2 + ... and some of the dyad-independent ergm terms still work. (We are working on replicating the full set of ergm terms.) For a full list of terms implemented, see ? terms.ergmm. Note that ergmm, like glm and unlike ergm, sets the first covariate $x_{1,i,j} \equiv 1$ for an intercept term, the (binary) ERGM equivalent of an edge count term.

For example,

> samplk.nm.l <- ergmm(samplk.tot~nodematch("group",diff=TRUE),tofit="mle", verbose=TRUE)

sets $\boldsymbol{x}_{1,i,j} \equiv 1$, $\boldsymbol{x}_{2,i,j} \equiv \mathbb{I}$ ($i \in \text{Loyal} \land j \in \text{Loyal}$), $\boldsymbol{x}_{3,i,j} \equiv \mathbb{I}$ ($i \in \text{Outcasts} \land j \in \text{Outcasts}$), etc., to produce a model of the form

 $logit(Pr(Y_{i,j} = 1)) = \beta_1 + \beta_2 \mathbb{I}(i \text{ and } j \text{ are both Loyal Opposition}) + \beta_3 \mathbb{I}(i \text{ and } j \text{ are both Outcasts}) + \cdots,$ equivalent to a ergm specification of

```
> samplk.nm.e <- ergm(samplk.tot~edges+nodematch("group",diff=TRUE))
  In fact, the produce the same coefficients and standard errors:
> summary(samplk.nm.l, point.est="mle", se=TRUE)
_____
Summary of model fit
_____
Formula:
         samplk.tot ~ nodematch("group", diff = TRUE)
Attribute: edges
Model:
         Bernoulli
MCMC sample of size 4000, draws are 10 iterations apart, after burnin of 10000 iterations.
Covariate coefficients MLE:
                      Estimate Std. Error z value Pr(>|z|)
                      edges
                      nodematch.group.Loyal
                                               0.98488
nodematch.group.Outcasts 17.05930 900.30202 0.0189
nodematch.group.Turks
                      nodematch.group.Waverers 1.66208 0.83596 1.9882
                                               0.04678 *
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
> summary(samplk.nm.e)
Summary of model fit
_____
         samplk.tot ~ edges + nodematch("group", diff = TRUE)
Formula:
Iterations: 20
Monte Carlo MLE Results:
                      Estimate Std. Error MCMC % p-value
edges
                      -1.6621 0.1793 NA <1e-04 ***
nodematch.group.Loyal
                       2.0675
                                 0.4904
                                         NA <1e-04 ***
                                         NA 0.9859
nodematch.group.Outcasts 18.3450 1038.5246
                               0.3858 NA <1e-04 ***
0.8360 NA 0.0477 *
nodematch.group.Turks
                     2.5784
nodematch.group.Waverers 1.6621
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
    Null Deviance: 424.2 on 306 degrees of freedom
 Residual Deviance: 289.1 on 301 degrees of freedom
AIC: 299.1
           BIC: 317.7
```

(Smaller is better.)

latentnet fits latent space models using terms euclidean and bilinear, which take two main arguments: d for dimension and G for number of clusters to use. So, for a classic example of the Sampson's monks fit, i.e.,

$$logit(Pr(\boldsymbol{Y}_{i,j}=1)) = \boldsymbol{\beta}_1 + |\boldsymbol{Z}_i - \boldsymbol{Z}_j|,$$

with Z_i modeled as 3 spherical Gaussian clusters with 2 dimensions,

> samplk.d2G3<-ergmm(samplk.tot~euclidean(d=2,G=3), verbose=TRUE)

Random actor-specific effects can also be used, with terms rsender, receiver, and rsociality, respectively: a receiver effects model of the form

$$logit(Pr(\boldsymbol{Y}_{i,j}=1)) = \boldsymbol{\beta}_1 + |\boldsymbol{Z}_i - \boldsymbol{Z}_j| + \boldsymbol{\gamma}_j,$$

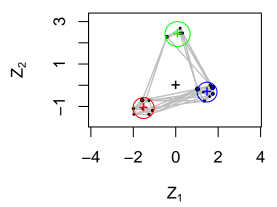
can be fit with

- > samplk.d2G3r<-ergmm(samplk.tot~euclidean(d=2,G=3)+rreceiver, verbose=TRUE)
- > mcmc.diagnostics(samplk.d2G3r)

The results can be visualized using

- > par(mfrow=c(1,2))
- > # Extract a clustering
- > Z.K.ref <- summary(samplk.d2G3,point.est="pmean")\$pmean\$Z.K
- > # Plot one model, saving positions, using Z.K.ref to set reference clustering.
- > Z.ref <- plot(samplk.d2G3, pie=TRUE, Z.K.ref=Z.K.ref)
- > # Plot the other model, using Z.ref and Z.K.ref to ensure similar
- > # orientation and coloring.
- > plot(samplk.d2G3r, rand.eff="receiver", pie=TRUE, Z.ref=Z.ref, Z.K.ref=Z.K.ref)

MKL Latent Positions of samplk.dKL Latent Positions of samplk.d samplk.tot ~ euclidean(d = 2, G = 3) samplk.tot ~ euclidean(d = 2, G = 3) + rreceiver



All ergmm terms also take additional arguments, specifying prior distributions. Their defaults are generally sensible, so we will not discuss them here.

Table 1: GLM families implemented by latentnet

Family	Link	family=	<pre>fam.par=list()</pre>
Bernoulli	logit	"Bernoulli"	
binomial	logit	"binomial"	$\mathtt{trials} = m$
Poisson	\log	"Poisson"	
Normal	linear	"normal"	$\mathtt{var} = \sigma^2$

4.2 Specifying non-binary models in ergmm

Fitting non-binary models using ergmm requires three additional arguments:

response An edge attribute in the network whose values are to be used as the response.

family A string specifying the family and the link to be used.

fam.par A named list of parameters required by some families.

The families listed above are specified using parameter values listed in Table 1. Also, see

```
> ? families.ergmm
```

for more information.

We have our network of nomination counts, with a maximum of three "successes", so we might model it using a Binomial distribution, i.e.,

$$\Pr(\boldsymbol{Y}_{i,j} = \boldsymbol{y}_{i,j} | \boldsymbol{\eta}_{i,j}) = \binom{3}{\boldsymbol{y}_{i,j}} (\operatorname{logit}^{\text{-}1}(\boldsymbol{\eta}_{i,j}))^{\boldsymbol{y}_{i,j}} (\operatorname{logit}^{\text{-}1}(\boldsymbol{\eta}_{i,j}))^{3-\boldsymbol{y}_{i,j}},$$

with $\eta_{i,j}$ being the same as in the binary case.

> plot(samplk.d2G3,pie=TRUE, Z.ref=Z.ref, Z.K.ref=Z.K.ref)

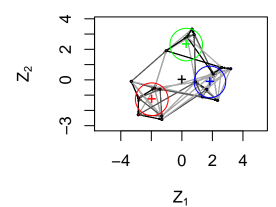
> plot(samplk.ct.d2G3,pie=TRUE, Z.ref=Z.ref, Z.K.ref=Z.K.ref, edge.col=samplk.ecol)

MKL Latent Positions of samplk.cKL Latent Positions of samplk.ct

samplk.tot ~ euclidean(d = 2, G = 3)

 Z_1

samplk.tot ~ euclidean(d = 2, G = 3) (nominations



Now, consider a latent cluster random effects model for the number of contexts of interactions, which takes into account that faction leaders are likely to have greater propensity to interact than non-leaders: $Y_{i,j} \stackrel{\text{ind.}}{\sim} \text{Poisson}(\mu_{i,j})$ with

$$\log \boldsymbol{\mu}_{i,j} = \boldsymbol{\eta}_{i,j} = \boldsymbol{\beta}_1 + \boldsymbol{\beta}_2(\mathbb{I}\left(i \text{ is a faction leader}\right) + \mathbb{I}\left(j \text{ is a faction leader}\right)) + |\boldsymbol{Z}_i - \boldsymbol{Z}_j| + \boldsymbol{\delta}_i + \boldsymbol{\delta}_j.$$

Some ergmm are missing that (for now), but the "leader" dummy variable can be used with ergmm's sendercov, receivercov, and/or socialitycov.

The term socialitycov(a) sets $x_{k,i,j} \equiv a_i + a_j$. Its ergm name is nodecov. Now, let's fit the model:

- > mcmc.diagnostics(zach.d2G2S)
- > summary(zach.d2G2S)

Summary of model fit

Formula: zach ~ socialitycov("leader") + euclidean(d = 2, G = 2) + rsociality

Attribute: contexts Model: Poisson

MCMC sample of size 4000, draws are 10 iterations apart, after burnin of 10000 iterations. Covariate coefficients posterior means:

Estimate 2.5% 97.5% Quantile of 0 edges 0.85346 0.18999 1.5176 0.00325 **

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Sociality effect variance: 0.7768003.

Overall BIC: 1149.352 Likelihood BIC: 801.9636

Latent space/clustering BIC: 267.818

Sociality effect BIC: 79.57072

Covariate coefficients MKL:

Estimate

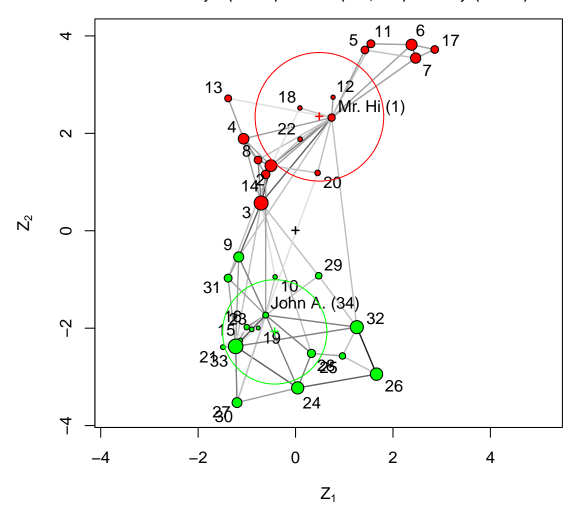
edges 0.3029993 socialitycov.leader.TRUE 2.0621726

> par(mar=c(5, 4, 4, 2) + 0.1)

> plot(zach.d2G2S, rand.eff="sociality", edge.col=zach.ecol, labels=TRUE)

MKL Latent Positions of zach.d2G2S

zach ~ socialitycov("leader") + euclidean(d = 2, G = 2) + rsociality (contexts)



Another useful application of socialitycov and others is when an undirected network of counts is a product of collapsing an affiliation network of actors to events to produce a network of actors only, counting the number of shared events for each pair of actors. If there is no particular pattern to the interaction, then the expected number of shared events between i and j would be proportional to the total number of events of i and to the number of events of j. If a_i is the total number of events associated with actor i, then using a vertex attribute equaling to $\log(a_i)$ in a socialitycov can be a good baseline model:

$$\log \boldsymbol{\mu}_{i,j} = \boldsymbol{\beta}_1 + \boldsymbol{\beta}_2(\log(a_i) + \log(a_j)) \Leftrightarrow \boldsymbol{\mu}_{i,j} = \exp(\boldsymbol{\beta}_1) \, a_i^{\boldsymbol{\beta}_2} a_j^{\boldsymbol{\beta}_2},$$

which produces proportionality when $\beta_2 \approx 1$. For an application of this, see Krivitsky et˜al. (2009).

4.3 Medium-term Roadmap

- Transition ergmm to use dyad-independent ergm terms.
- Enable latentnet to fit heterogeneous trial counts.
- Enable latentnet to fit normal distribution variance.

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