Advanced Network Analysis

ERGMs for Valued Networks

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Readings

- Pavel N Krivitsky. Exponential-family random graph models for valued networks. *Electronic Journal of Statistics*, 6: 1110--1128, 2012.
- Bruce A. Desmarais and Skyler J. Cranmer. Statistical inference for valued-edge networks: The generalized exponential random graph model. *PloS One*, 7(1), 2012.

Overview

- Many networks of interest have valued rather than binary edges, e.g. trade network, friendships, romantic relationships.
- Can generalize ERGMs to modeling networks with count or rank-ordered valued edges.
- Similar properties and estimation approach.

Estimation

Install Packages:

```
rm(list=ls())
#install.packages("ergm.count")
library(statnet)
library(ergm.count)
library(networkdata)
```

Load the Data

We are going to use Gade et al's data. Our dependent variable--- the valued network--- records the number of collaborations between rebel groups.

```
data(gadeData)
# data characs
actors = sort(unique(c(gadeData$Var1, gadeData$Var2)))
gadeData<-sort(gadeData)
gadeData$coopActions<-round(gadeData$coopActions^2)
#These are the dyadic variables. They
#must be in matrix form.
dyadVars = names(gadeData)[c(3,5:8)]
n = length(actors) ; p = length(dyadVars)</pre>
```

```
# create empty arr object for all dyad vars
dyadArray = array(0,
    dim=c(n,n,p),
    dimnames=list(actors,actors,dyadVars)
)
# loop through and fill in
for(param in dyadVars){
    for(i in 1:nrow(gadeData)){
        a1 = gadeData$Var1[i]
        a2 = gadeData$Var2[i]
        val =gadeData[i,param]
        dyadArray[a1,a2,param] = val
    }
}
```

```
# These are node-level variables.
nodeVars = names(gadeData)[9:11]
nodeData = unique(gadeData[,c('Var1',nodeVars)])
rownames(nodeData) = nodeData$Var1
nodeData = nodeData[actors,c(-1)]
# The DV must be a network object
net = as.network(
    dyadArray[,,'coopActions'],
    directed=FALSE, loops=FALSE,
    matrix.type='adjacency',
    ignore.eval = FALSE,
    names.eval = "coopActions"
    )
```

Look at the first 10 rows and cols:

```
as.matrix(net[1:10,1:10])
```

```
101st 13th 1st AALS AARB AASB AASG ADE AF AFB
##
## 101st
             0
                      0
                            0
                                 0
                                                   0
                                                       0
## 13th
                      0
                                                       0
## 1st
                      0
## AALS
                                               0 0
                                                       0
                      0
## AARB
                      0
                            0
                                               0
             0
                                                       0
## AASB
             0
                      0
                                                       0
## AASG
                      0
                            0
                                               0 0
                                 0
                                      0
                                                       1
## ADF
                  0
                            0
                                               0 0
                                                       0
                      0
                                 0
                                           0
                      0
                                               0 0
## AF
                  0
                            0
                                 1
                                      0
                                           0
                                                       0
## AFB
                  0
                      1
                                           1
                            0
                                 0
                                      0
                                                0
                                                       0
                                                   0
```

```
# Set node attributes
for(param in nodeVars){
    network::set.vertex.attribute(net, param, nodeData[,param])
}

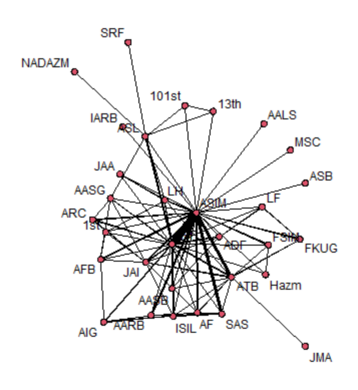
# Set network attributes:
set.network.attribute(net,'loc.dyad',dyadArray[,,'loc.dyad'])
set.network.attribute(net,'spons.dyad',dyadArray[,,'spons.dyad'])
# We can view the attribute as a sociomatrix.
as.matrix(net, attrname = "coopActions")[1:10, 1:10]
```

##		101st	13th	1st	AALS	AARB	AASB	AASG	ADF	ΑF	AFB
##	101st	0	1	0	0	0	0	0	0	0	0
##	13th	1	0	0	0	0	0	0	0	0	0
##	1st	0	0	0	0	0	0	0	0	0	1
##	AALS	0	0	0	0	0	0	0	0	0	0
##	AARB	0	0	0	0	0	0	0	0	2	0
##	AASB	0	0	0	0	0	0	0	1	0	0
##	AASG	0	0	0	0	0	0	0	0	0	1
##	ADF	0	0	0	0	0	1	0	0	0	0
##	AF	0	0	0	0	2	0	0	0	0	0
##	AFB	0	0	1	0	0	0	1	0	0	0

Make a Network Graph with Valued Edges:

```
plot(net, edge.col = "black", usecurve = TRUE,
    edge.curve = 0, edge.lwd=.25*dyadArray[,,"coopActions"],
    displaylabels = TRUE)
```

Make a Network Graph with Valued Edges:



Valued ERGMs

- The package ergm.count extends the ergm package to allow for modeling networks with valued edges. This is done by specifying the response argument with the name of the edge attribute to use as the response variable.
- New concept: a reference distribution
 - Need to think about how the values for connections we measure are distributed. The reference distribution specifies the model for the data before we add any ERGM terms.

```
help("ergm-references")
```

The Reference Distribution

Possible reference measures to represent baseline distributions

Reference measures currently available are:

Poisson

Poisson-reference ERGM: Specifies each dyad's baseline distribution to be Poisson with mean 1: h(y)=\(\int_{i,j}\) 1/y_{i,j,j}!, with the support of y_{i,j}\) being natural numbers (and 0). Using valued_ERGM_terms that are "generalized" from their binary counterparts, with form "sum" (see previous link for the list) produces Poisson regression. Using CMP induces a Conway-Maxwell-Poisson distribution that is Poisson when its coefficient is 0 and geometric when its coefficient is 1.

Three proposal functions are currently implemented, two of them designed to improve mixing for sparse networks. They can can be selected via the MCMC.prop.weights= control parameter. The sparse proposals work by proposing a jump to 0. Both of them take an optional proposal argument p0 (i.e., MCMC.prop.args=list(p0=...)) specifying the probability of such a jump. However, the way in which they implement it are different:

"random"

Select a dyad (i,j)at random, and draw the proposal $y_{i,j}^{(j,j)}$ (i,j) $^{(i,j)}$ (a Poisson distribution with mean slightly higher than the current value and conditional on not proposing the current value).

"0inflated"

As "random" but, with probability p0, propose a jump to 0 instead of a Poisson jump (if not already at 0). If p0 is not given, defaults to the "surplus" of 0s in the observed network, relative to Poisson.

"TNT" (the default)

As "Oinflated" but instead of selecting a dyad at random, select a tie with probability p0, and a random dyad otherwise, as with the binary TNT. Currently, p0 defaults to 0.2.

Geometric

Geometric-reference ERGM: Specifies each dyad's baseline distribution to be uniform on the natural numbers (and 0):
h(y)=1. In itself, this "distribution" is improper, but in the presence of gum, a geometric distribution is induced. Using CMP (in addition to gum) induces a Conway-Maxwell-Poisson distribution that is geometric when its coefficient is 0 and Poisson when its coefficient is -1.

Binomial(trials)

The Sample Space

- For binary ERGMs, the sample space (or support) \mathcal{Y} the set of possible networks that can occur is usually some subset of 2^N , 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 \mathcal{S} , the set of possible values each relationship may take. For example, for count data, that's $\mathcal{S} = \{0, 1, \ldots, s\}$ if the maximum count is s and $\{0, 1, \ldots\}$ if there is no a priori upper bound. Having specified that, \mathcal{Y} is defined as some subset of $\mathcal{S}^{\mathcal{Y}}$: the set of possible ways to assign to each relationship a value.

Estimate a Valued ERGM

```
m0 <- ergm(net ~ sum +
    nodecov('averageId.node') +
    nodecov('size.node') +
    nodecov('spons_actor.node') +
    absdiff('averageId.node') +
    absdiff('size.node') +
    edgecov('loc.dyad') +
    edgecov('spons.dyad'),
    response = "coopActions", reference = ~Poisson)
mcmc.diagnostics(m0)</pre>
```

Diagnostics

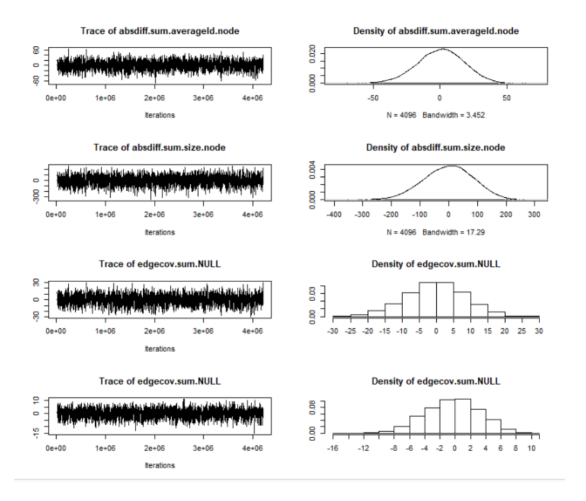


Table of Results

```
summary(m0)
## Iterations: 5 out of 20
##
## Monte Carlo MLE Results:
                              Estimate Std. Error MCMC % z value Pr(>|z|)
##
## sum
                              -8.10533
                                         1.05447
                                                      0 -7.687 < 1e-04 
                                                      0 11.459 < 1e-04 *
## nodecov.sum.averageId.node
                               0.55962
                                         0.04884
## nodecov.sum.size.node
                               0.07523
                                         0.00605
                                                      0 12.434 < 1e-04 *
                                         0.13749
                               0.73878
                                                      0 5.373 < 1e-04 *
## nodecov.sum.spons_actor.node
## absdiff.sum.averageId.node
                                                      0 -3.720 0.000199 *
                              -0.17780
                                         0.04779
## absdiff.sum.size.node
                              -0.08958
                                         0.01007
                                                      0 -8.897 < 1e-04 *
## edgecov.sum.NULL
                                                      0 3.997 < 1e-04 *
                               3.95933
                                         0.99062
## edgecov.sum.NULL.1
                               0.58640
                                         0.14944
                                                      0 3.924
                                                                < 1e-04 *
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
##
       Null Deviance:
                        0.0 on 465 degrees of freedom
   Residual Deviance: -692.7 on 457 degrees of freedom
##
##
## Note that the null model likelihood and deviance are defined to be 0. This
## (LRT, Analysis of Deviance, AIC, BIC, etc.) is only valid between models w
```

Interpretation

Let's calculate the expected number of links between two average rebel groups:

- Set all *x*s to their mean values
- ullet Calculate the expected number as e^{X^Teta}

```
#to get the mean values (in same order as in our model)
x_mean<-apply(data[,c(9,10,11,5,6,7,8)],2,mean)
exp(c(1,x_mean)%*%m0$coef) #expected number of collaborations</pre>
```

Your Turn

Calculate the expected number of links between two average rebel groups with no ideological differences:

Accounting for Network Dependencies

Individual Heterogeneity

Actors may have different overall propensities to interact. This has been modeled using using degeneracy-prone terms like k-star counts. With valued ERGMs, a more robust measure is:

$$m{g_{actor~cov.}}(m{y}) = \sum\limits_{i \in N} rac{1}{n-2} \sum\limits_{j,k \in \mathbb{Y}_{\mathrm{i}} \wedge j < k} (\sqrt{m{y}_{i,j} - \overline{\sqrt{m{y}}}}) (\sqrt{m{y}_{i,k} - \overline{\sqrt{m{y}}}})$$

This is essentially a measure of covariance between the squared values of edges incident (originating) from the same actor. Implemented with the term nodesqrtcovar.

Triadic Closure

```
transitiveweights(twopath, combine, affect)
```

- twopath---given $oldsymbol{y}_{i,j}$ and $oldsymbol{y}_{k,j}$, how to compute the value for the two-path?
 - "min"---the minimum of their values
 - "geomean---geometric mean
- combine---given the strength of the two-paths $m{y}_{i->k->j}$ for all $k \neq i,j,$ how to combine the values?
 - "max"--- the strength of the strongest path
 - "sum"---the sum of path strength
- affect ---given the combined strength pf the two-paths between i and j, how should they affect $Y_{i,j}$?
 - ∘ "min"
 - ∘ "geomean"

Example

```
m1 <- ergm(net ~ sum +
    nodecov('averageId.node') +
    nodecov('size.node') +
    nodecov('spons_actor.node') +
    absdiff('averageId.node') +
    absdiff('size.node') +
    edgecov('loc.dyad') +
    edgecov('spons.dyad')+
    transitiveweights("min","max","min"),
    response = "coopActions", reference = ~Poisson)
par(mfrow = c(3,2))
mcmc.diagnostics(m1)</pre>
```

Diagnostics

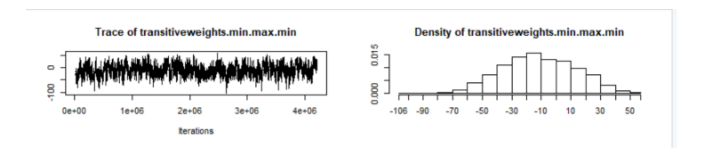


Table of Results

```
summary(m1)
## Iterations: 6 out of 20
##
## Monte Carlo MLE Results:
                                 Estimate Std. Error MCMC % z value Pr(>|z|)
##
## sum
                                -7.417775
                                            1.061091
                                                             -6.991
                                                                     < 1e-04
## nodecov.sum.averageId.node
                                 0.550254
                                           0.049272
                                                             11.168 < 1e-04
## nodecov.sum.size.node
                                 0.067013
                                            0.005905
                                                             11.348
                                                                     < 1e-04
                                                          0
## nodecov.sum.spons_actor.node
                                 0.827648
                                            0.132379
                                                          0 6.252
                                                                     < 1e-04
## absdiff.sum.averageId.node
                                                             -3.586 0.000336
                                -0.160610
                                           0.044788
                                                          0
## absdiff.sum.size.node
                                -0.077496
                                            0.009579
                                                          0
                                                             -8.090
                                                                     < 1e-04
                                                              3.762 0.000168
## edgecov.sum.NULL
                                 3.782666
                                            1.005441
                                                          0
## edgecov.sum.NULL.1
                                            0.128019
                                                              4.089
                                                                     < 1e-04
                                 0.523423
                                                          0
## transitiveweights.min.max.min -0.635663
                                                          0 -13.858
                                            0.045871
                                                                     < 1e-04
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
       Null Deviance: 0.0 on 465 degrees of freedom
##
   Residual Deviance: -790.2 on 456 degrees of freedom
##
##
```

Note that the null model likelihood and deviance are defined to be 26.74 his

Simulating Networks

Simulating Networks

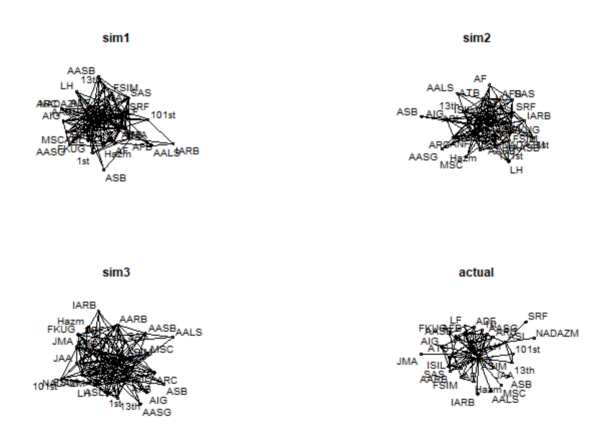
We can use the estimates from our model to simulate a network (just like with ERGMs). If the simulated networks look similar to the observed network, then our model has a good fit.

```
# Simulate from model fit:
simNets <- simulate(m1, nsim = 3)

# Define a plotting function:
plotSimNet = function(net, label){
    set.seed(6886)
    plot(net, edge.col = "black", usecurve = TRUE,
        edge.curve = 0, edge.lwd=.1*dyadArray[,,"coopActions"],
        displaylabels = TRUE)
    title(label) }</pre>
```

```
par(mfrow = c(2, 2))
# add actual network to list of sim nets
# for comparison
simNets[[4]] = net
labels = c(paste0("sim",1:3), 'actual')
lapply(1:length(simNets), function(i){
    plotSimNet(simNets[[i]], labels[i]) })
```

Observed vs. Simulated Networks



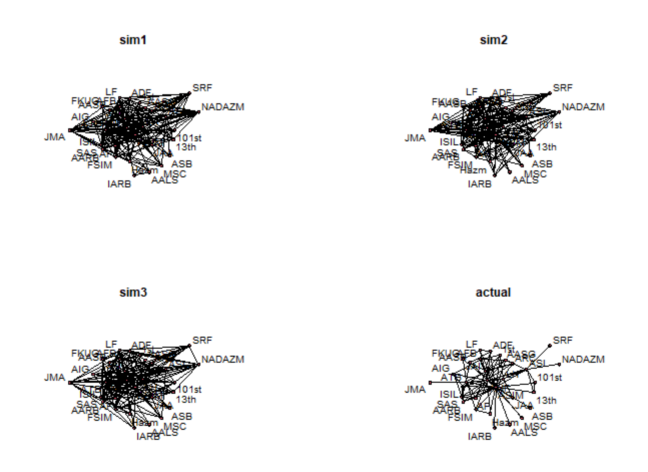
With Fixed Coordinates:

```
#Plot the original network to get the layout:
set.seed(6886)
p<-plot(net, edge.col = "black", usecurve = TRUE,
        edge.curve = 0, edge.lwd=.25*dyadArray[,,"coopActions"],
        displaylabels = TRUE)</pre>
```



```
# Define a plotting function:
plotSimNet = function(net, label){
    set.seed(6886)
    plot(net, edge.col = "black", usecurve = TRUE,
    edge.curve = 0, edge.lwd=.1*dyadArray[,,"coopActions"],
    displaylabels = TRUE, coord=p)
```

Observed vs. Simulated Networks (Fixed Coord.)



Assessing Model Fit

In the above exercise, we compared our network to only 3 simulated networks. Ideally, we would like to compare it to more than 3. Since it's difficult to look at thousands of simulated networks on a graph, a way to compare our network to thousands of such simulated networks is by summarizing the characteristics of these simulated networks, such as the sum of edges or various other measures.

Notice that in the code below that, in addition to network statistics included in model m1, we can also summarize statistics that were not explicitly included in m1, such as nodesquarecovar. This is because our simulated networks may still exhibit vertex heterogeneity as a function of the modeled network properties (e.g. triangles), i.e. more triangles may also lead to more k-stars.

Also notice that I specified output="stats". Since I only care about network summaries, I am telling the function to NOT save the actual simulated networks, but only their summary statistics. This saves memory space.

```
# Simulate from model fit:
simNets1000 <- simulate(m1, monitor = ~nodesqrtcovar(TRUE),
    nsim = 1000, output = "stats")
```

Results of the Simulation

colnames(simNets1000)

```
## [1] "sum" "nodecov.sum.averageId.node"
## [5] "absdiff.sum.averageId.node" "absdiff.sum.size.node"
## [9] "transitiveweights.min.max.min" "nodesqrtcovar.centered"
```

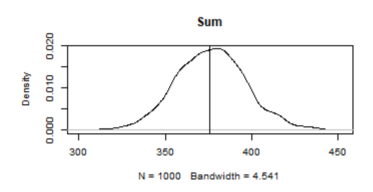
"node

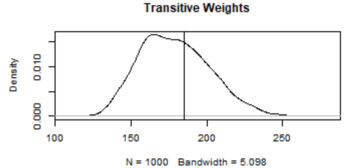
"edge

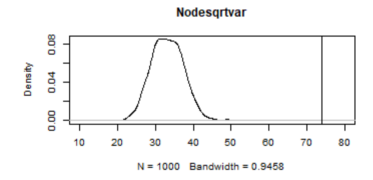
Plot the Summary of the Simulations

```
#How prevalent are k-stars in the observed network?
obsNet<-summary(net~sum+transitiveweights("min", "max", "min")+nodesqr1
                response = "coopActions")
par(mfrow = c(2, 2))
#1st col.=sum
plot(density(simNets1000[,1]), main="")
abline(v = obsNet[1])
title("Sum")
# 9th col. = transitiveweights
plot(density(simNets1000[,9]), main="")
abline(v = obsNet[2])
title("Transitive Weights")
# 10th col. = nodesartcovar
plot(density(simNets1000\lceil,10\rceil), main="", xlim=c(10,80))
abline(v = obsNet)
title("Nodesqrtvar")
```

Plot the Summary of the Simulations







Improve Model Specofication

```
m2 <- ergm(net ~ sum +
    nodecov('averageId.node') +
    nodecov('size.node') +
    nodecov('spons_actor.node') +
    absdiff('averageId.node') +
    absdiff('size.node') +
    edgecov('loc.dyad') +
    edgecov('spons.dyad')+
    transitiveweights("min","max","min")+
    nodesqrtcovar(TRUE),
    response = "coopActions", reference = ~Poisson)</pre>
```

Improve Model Specification

constraints.

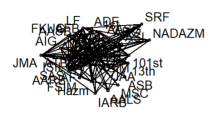
```
## Iterations: 12 out of 20
##
## Monte Carlo MLE Results:
                                 Estimate Std. Error MCMC % z value Pr(>|z|)
##
                                                          0 -4.274
## sum
                                -4.726287
                                            1.105867
                                                                     < 1e-04
## nodecov.sum.averageId.node
                                 0.296854
                                            0.056879
                                                          0 5.219
                                                                     < 1e-04
## nodecov.sum.size.node
                                 0.070925
                                           0.006855
                                                          0
                                                             10.346
                                                                     < 1e-04
## nodecov.sum.spons_actor.node
                                 0.230775
                                            0.140995
                                                             1.637
                                                                     0.10168
                                                          0
## absdiff.sum.averageId.node
                                -0.256984
                                                             -5.413
                                                                     < 1e-04
                                            0.047476
## absdiff.sum.size.node
                                -0.103376
                                            0.008632
                                                          0 -11.975
                                                                     < 1e-04
                                                                     0.00153
## edgecov.sum.NULL
                                 3.253555
                                            1.026670
                                                              3.169
                                                          0
## edgecov.sum.NULL.1
                                 0.241622
                                            0.159609
                                                             1.514
                                                                     0.13007
                                                          0
                                                             -6.702
## transitiveweights.min.max.min -0.690133
                                            0.102974
                                                                     < 1e-04
                                                          0
## nodesqrtcovar.centered
                                                              9.845
                                                                     < 1e-04
                                 2.627587
                                            0.266903
                                                          0
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
       Null Deviance: 0.0 on 465 degrees of freedom
##
   Residual Deviance: -932.9 on 455
                                      degrees of freedom
##
##
## Note that the null model likelihood and deviance are defined to be 0. This
## (LRT, Analysis of Deviance, AIC, BIC, etc.) is only valid between models w
```

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Your Turn

Follow the steps we did for m1 to evaluate the fit of model m2. Start by simulating a small number of networks, plot them and compare them to the observed network. Then simulate 1000 networks and compare them to the observed network on the key network statistics. Does m2 have a better fit?

sim1 sim2

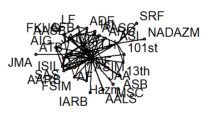




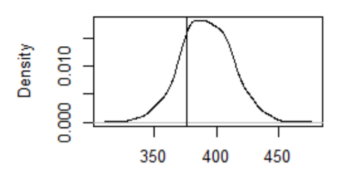
sim3



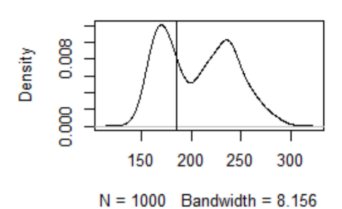
actual



Sum

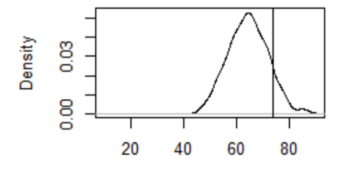


Transitive Weights



N = 1000 Bandwidth = 4.672

Nodesqrtvar



N = 1000 Bandwidth = 1.701