CDatanet: An R package to simulate and estimate a Count Data Model with Social Interactions

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1 Introduction

There is a large and growing literature on peer effects in economics.¹ Recent contributions include among other, models for discrete data such as binary (e.g., Lee et al., 2014; Liu, 2019), ordered (e.g., Boucher et al., 2018) and multinomial (e.g., Guerra and Mohnen, 2017). However, there are hardly any models dealing with count data (e.g., data collected by asking respondents: "How many times do you smoke daily?").

I present a model in which the outcome takes count values (0, 1, 2, ...). In this model, individuals interact through a directed network and their peers' decision may influence their own decision. Moreover, they do not observe their peers' outcome and form rational belief about that.

I also present an easy-to-use R package—named **CDatanet**—for implementing the model. This note is a supplement for the package manual. I provide examples with simulated data to show how to use the package. The package is located on GitHub at github.com/ahoundetoungan/CDatanet. All the results of the note can be replicated following the code provided.

The rest of this note is organized as follow. Section 2 briefly introduces the model. Section 3 explains how to install the package. Sections 4 and 5 provide examples in which the network matrix is assumed exogenous and endogenous respectively. They present Monte Carlo experiments to assess the performance of the estimator of the model parameters. They also discuss how to control for the endogeneity of the network. Section 6 concludes.

2 Model

Let $\mathcal{V} = \{1, ..., n\}$ be a set of n players indexed by i and y_i the observed integer outcome of player i (e.g., the number of cigarettes smoked per day or per week). I denote by G the adjacency matrix such that $G_{ij} = \frac{1}{n_i}$ if i knows j and $G_{ij} = 0$ otherwise, where n_i the number of friends of i. Let also X be the matrix of explanatory variables.

As in Liu (2019), I assume that there is a latent variable y_i^* which determines the observed count variable y_i . This latent variable y_i^* and the observed y_i are linked as follow.

Let be $(a_q)_{q\in\mathbb{N}}$ a sequence given by $a_0=-\infty,\ a_1\in\mathbb{R}$ and $a_q=a_1+\gamma(q-1)$ for $q\in\mathbb{N}^*$ and $\gamma\in\mathbb{R}_+^*$. If $y_i^*\in(a_q,a_{q+1}]$ then $y_i=q$.

The first order conditions of the game equilibrium implies that

$$y_i^* = \lambda \sum_{i=1}^n \sum_{r=0}^\infty r G_{ij} p_{ir} + \boldsymbol{x}_i' \boldsymbol{\beta} + \varepsilon_i$$

¹For recent reviews, see Boucher and Fortin (2016), De Paula (2017) and Bramoullé et al. (2019).

where p_{ir} is the probability of $y_i = r$, λ is the peer effect and $\varepsilon_i \sim \mathcal{N}\left(0, \sigma_{\varepsilon}^2\right)$.

The game equilibrium is given by

$$p_{ir} = \Phi\left(\frac{\lambda \sum_{i=1}^{n} \sum_{r=0}^{\infty} rG_{ij}p_{ir} + \boldsymbol{x}_{i}'\boldsymbol{\beta} - a_{q}}{\sigma_{\varepsilon}}\right) - \Phi\left(\frac{\lambda \sum_{i=1}^{n} \sum_{r=0}^{\infty} rG_{ij}p_{ir} + \boldsymbol{x}_{i}'\boldsymbol{\beta} - a_{q+1}}{\sigma_{\varepsilon}}\right)$$

where Φ is the probability density function of $\mathcal{N}(0,1)$.

To estimate $\theta = (\lambda, \beta', \sigma_{\varepsilon})'$, I rely on the NPL method. I refer the reader to Houndetoungan (2020) for more details. The estimation method is implemented in the package **CDatanet**.

3 CDatanet Installation

The code source of the package is written in C++ with the package **Rcpp** (Eddelbuettel et al., 2020). **CDatanet** is currently available on GitHub. Its installation requires to prior install **devtools** package (Wickham et al., 2020). Moreover, Windows users should install Rtools compatible with their R version.

CDatanet package can be installed from this GitHub repos using the install_github function of devtools. All the dependencies will also be installed automatically.

```
library(devtools)
install_github("ahoundetoungan/CDatanet")
```

4 Examples with exogenous network

In this section, I present examples to show how to use the package. I simulate data following the model by assuming that the network matrix is exogenous. I then show how to estimate the model parameters using functions implemented in **CDatanet**. I also use Monte Carlo simulations to assess the performance of the estimator of the model parameters.

4.1 Data simulation

Given the adjacency matrix, the explanatory variables and θ , the function simCDnet can used to simulate data. I assume that there are M sub-networks. The number of individuals in each sub-network is randomly chosen between 100 and 1000. I assume two exogenous variables (plus the intercept) and I control for the contextual effect.

```
set.seed(2020)
library(CDatanet)
# Groups' size
       <- 5 # Number of sub-groups
       <- round(runif(M, 100, 1000))
nvec
print(nvec)
## [1] 682 455 657 529 222
       <- sum(nvec)
print(n)
## [1] 2545
# Parameters
                           # peer effect
      <-c(2, -1.9, 0.8) # own effects
gamma <- c(1.5, -1.2)
                           # contextual effects
sigma <- 1.5
                           # standard deviation of epsilon
```

```
theta <- c(lambda, beta, gamma, sigma)

# X
data <- data.frame(x1 = rnorm(n, 1, 1), x2 = rexp(n, 0.4))</pre>
```

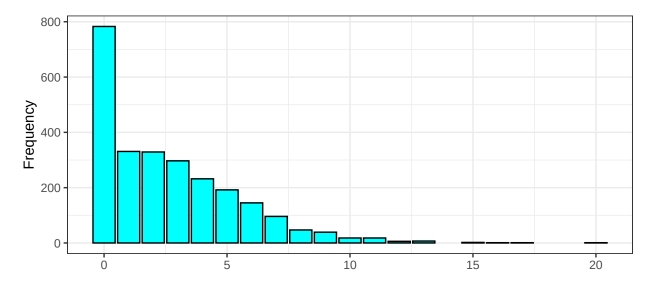
To simulate the network matrix, I assume that the number of friends of each individual is randomly chosen between 0 and 30.

```
# Network
Glist <- list()</pre>
for (m in 1:M) {
                <- nvec[m]
 nm
  Gm
               <- matrix(0, nm, nm)
               <- 30
 max_d
 for (i in 1:nm) {
               <- sample((1:nm)[-i], sample(0:max_d, 1))
    Gm[i, tmp] <- 1
  }
                <- rowSums(Gm); rs[rs == 0] <- 1
 rs
  Gm
                <- Gm/rs
  Glist[[m]]
                <- Gm
}
```

I can now simulate the count data. The output of simCDnet includes yst the vector of y_i^* , y the vector of y_i^* , yb the vector of $\sum_{r=0}^{\infty} rp_{ir}$, Gyb the vector of $\sum_{i=1}^{n} \sum_{r=0}^{\infty} rG_{ij}p_{ir}$ and iteration as the number of iterations performed to find the fixed point yb.

```
## Min. 1st Qu. Median Mean 3rd Qu. Max.
## 0.000 0.000 2.000 2.617 4.000 20.000
table(y)
```

```
## y
                 3
                         5
                             6
                                 7
                                     8
                                         9
                                             10
                                                 11
                                                     12
                                                         13
                                                             15
                                                                 16
                                                                     17
## 783 331 329 297 232 192 145 96
                                            18
                                    47
                                        39
                                                18
# Plot data histogram
library(ggplot2)
print(ggplot(data = data.frame(y = y), aes(x = y)) +
        geom_bar(color = "black", fill = "cyan") +
        theme_bw() + xlab("") + ylab("Frequency"))
```



4.2 Estimation

Using the simulated data, I estimate the count data model as well as the SART and SAR models. The output of these models has a summary class to summarize the results. In addition, the summary function computes marginal effects for the count data model.

```
Count data
     <- CDnetNPL(formula = y ~ x1 + x2, contextual = TRUE, Glist = Glist,
                  optimizer = "nlm",
                 data = data, npl.ctr = list(print = FALSE, maxit = 5e3))
summary(CD)
## Count data Model with Social Interactions
##
## Method: Nested pseudo-likelihood (NPL)
## Iteration: 12
##
## Network:
## Number of groups
## Sample size
                                2545
## Average number of friends:
                               14.98075
##
## Coefficients:
##
                Estimate Std. Error t value Pr(>|t|)
## lambda
                0.377313
                           0.075603
                                        4.99 6.02e-07 ***
                           0.155886
## (Intercept) 1.940142
                                       12.45
                                               <2e-16 ***
## x1
               -1.836252
                           0.037055
                                     -49.55
                                               <2e-16 ***
## x2
                0.804280
                           0.012901
                                       62.34
                                               <2e-16 ***
## G: x1
                1.587260
                           0.112376
                                       14.12
                                               <2e-16 ***
## G: x2
               -1.211202
                           0.074873
                                     -16.18
                                               <2e-16 ***
##
## Marginal Effects:
                Estimate Std. Error t value Pr(>|t|)
##
                           0.056942
                                        4.98 6.31e-07 ***
## lambda
                0.283665
## (Intercept) 1.458607
                           0.117014
                                       12.47
                                               <2e-16 ***
## x1
               -1.380502
                           0.033611
                                     -41.07
                                               <2e-16 ***
## x2
                0.604661
                           0.011963
                                       50.55
                                               <2e-16 ***
## G: x1
                1.193309
                           0.086218
                                       13.84
                                               <2e-16 ***
```

```
-0.910587 0.057630 -15.80 <2e-16 ***
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## sigma: 1.503368
## log pseudo-likelihood: -3650.167
# SART
SART <- SARTML(formula = y ~ x1 + x2, contextual = TRUE, Glist = Glist,
             optimizer = "nlm", data = data, print = FALSE)
summary(SART)
## SART Model
## Method: Maximum Likelihood (ML)
## Network:
## Number of groups
                      : 5
## Sample size
                       : 2545
## Average number of friends: 14.98075
## Coefficients:
             Estimate Std. Error t value Pr(>|t|)
##
## lambda
            ## (Intercept) 2.526843 0.150888 16.75 <2e-16 ***
           -1.929519 0.039198 -49.23 <2e-16 ***
## x1
            ## x2
## G: x1
            ## G: x2
            -1.157105 0.063618 -18.19 <2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## sigma: 1.630561
## log likelihood: -3811.928
SAR <- SARML(formula = y ~ x1 + x2, contextual = TRUE, Glist = Glist,
           optimizer = "nlm", data = data, print = FALSE)
summary(SAR)
## SAR Model
## Method: Maximum Likelihood (ML)
## Network:
## Number of groups
                      : 5
## Sample size
## Average number of friends: 14.98075
## Coefficients:
##
            Estimate Std. Error t value Pr(>|t|)
           ## lambda
## (Intercept) 2.532121 0.117490 21.55 <2e-16 ***
           -1.386272 0.027828 -49.82
## x1
                                     <2e-16 ***
## x2
            0.706970 0.010947 64.58 <2e-16 ***
## G: x1
            1.084204 0.085506 12.68 <2e-16 ***
```

Using the count data model, the peer effect is estimated at 0.38, which is close to the true value, 0.4. By replicating this experience several times, the average of the estimates is likely to be almost equal to the true value (see next section). Note that this parameter is not interpretable. Instead, one should interpret the marginal effect: increasing of the expected y_j of the peers by one implies an increasing of the expected y_i of the individual by 0.28. However, estimating the SART model using data simulated from the count data model underestimates the peer effect (0.27). The package does not implement the marginal effect under the SART model. This latter should be much lower. Lastly, the SAR model more underestimates the peer effect (0.16). In addition, this is directly the marginal effect because the SAR model does not assume a limited dependent variable.

4.3 Monte Carlo Simulations

In this section, I conduct Monte Carlo simulations to assess the performance of the estimator. I use the same settings as in Section 4.1 except that the number of individuals in each sub-network is set to 500. Let's build the following Monte Carlo function that simulates the data, estimates the three models and returns their estimators.

```
fMC <- function(s) {</pre>
  # Groups' size
  М
          <- 5
          \leftarrow rep(500, 5);
  nvec
          <- sum(nvec)
  # Parameters
  lambda \leftarrow 0.4
          <-c(2, -1.9, 0.8)
  beta
  gamma \leftarrow c(1.5, -1.2)
  sigma <- 1.5
  theta <- c(lambda, beta, gamma, sigma)
  # X
          \leftarrow data.frame(x1 = rnorm(n, 1, 1), x2 = rexp(n, 0.4))
  data
  # Network
  Glist <- list()</pre>
  for (m in 1:M) {
    nm
                   <- nvec[m]
                   <- matrix(0, nm, nm)
    Gm
                   <- 30
    max_d
    for (i in 1:nm) {
                   <- sample((1:nm)[-i], sample(0:max_d, 1))
      tmp
      Gm[i, tmp] <- 1
    }
                   <- rowSums(Gm); rs[rs == 0] <- 1
    rs
    Gm
                   <- Gm/rs
    Glist[[m]]
                   <- Gm
  }
  # y
```

```
<- simCDnet(formula = ~ x1 + x2 | x1 + x2, Glist = Glist, theta = theta,
                   data = data)
       <- ytmp$y
data$y <- y
# Models
       <- CDnetNPL(formula = y ~ x1 + x2, contextual = TRUE, Glist = Glist,
CD
                   optimizer = "nlm",
                   data = data, npl.ctr = list(print = FALSE, maxit = 5e3))
SART
       <- SARTML(formula = y ~ x1 + x2, contextual = TRUE, Glist = Glist,
                 optimizer = "nlm", data = data, print = FALSE)
       <- SARML(formula = y ~ x1 + x2, contextual = TRUE, Glist = Glist,
SAR
                optimizer = "nlm", data = data, print = FALSE)
c(CD$estimate, SART$estimate, SAR$estimate)
```

I run the Monte Carlo function fMC 1000 times and compute the average of each estimator. To make it faster, I run the replications in parallel using doParallel package (Corporation and Weston, 2019).

```
library(doParallel)
n.cores <- 32
replic <- 1000 #Number of replications
out.mc <- mclapply(1:replic, fMC, mc.cores = n.cores)
out.mc <- apply(t(do.call(cbind, out.mc)), 2, mean)
out.mc <- cbind(theta, out.mc[1:7], out.mc[8:14], out.mc[15:21])
colnames(out.mc) <- c("TrueValue", "CoutData", "SART", "SAR")
print(out.mc)</pre>
```

```
##
               TrueValue
                           CoutData
                                          SART
                                                      SAR
## lambda
                         0.4014868 0.2073720
                                               0.1379745
                     0.4
## (Intercept)
                     2.0 1.9967916
                                    2.7235037
                                                2.6711174
## x1
                    -1.9 -1.8990528 -1.9996477 -1.4179726
## x2
                     0.8
                         0.7998454
                                    0.8260788
                                               0.7005770
## G: x1
                                    1.3591351
                     1.5 1.4982137
                                               0.9242272
## G: x2
                    -1.2 -1.1994937 -1.0790096 -0.6846019
                     1.5
## sigma
                         1.4974801 1.6256067
                                                1.3795906
```

The estimator of the count data model seems unbiased while the SART and the SAR models underestimate the peer effect.

5 Examples with endogenous network

Peer effect estimation is generally based on the assumption of exogeneity of the adjacency matrix. This means that the probability of link formation is not correlated to the error term in the count data model. Such an assumption is strong as the link formation probability may depend on unobserved characteristics (e.g, gregariousness) that also influence the outcome. In this section, I simulate data with endogenous network (which violates the assumption). I then show that the peer effect is overestimated when one does not control for the endogeneity of the network. I also present a method to control for the endogeneity. Finally, I use Monte Carlo simulations to prove that this method performs well.

5.1 Data simulation

I assume three sub-networks. The number of individuals in each sub-network is randomly chosen between 100 and 500.

```
rm(list = ls())
set.seed(2020)
# Groups' size
       <- 3 # Number of sub-groups
       <- round(runif(M, 100, 500))
print(nvec)
## [1] 359 258 347
       <- sum(nvec)
print(n)
## [1] 964
# Parameters
lambda \leftarrow 0.4
                            # peer effect
       <-c(2, -1.9, 0.8)
                            # own effects
gamma <- c(1.5, -1.2) # contextual effects
sigma <- 1.5
                            # standard deviation of epsilon
theta <- c(lambda, beta, gamma, sigma)
```

The network matrix follows the dyadic linking model presented in Houndetoungan (2020). Let \mathbf{A} be the matrix of links such that $A_{ij} = 1$ if i knows j and $A_{ij} = 0$. In other words, $A_{ij} = 1$ if $G_{ij} = \frac{1}{n_i}$. It is easier to work with \mathbf{A} (binary data) in the network formation model. However, in the count data model, I use \mathbf{G} as row-normalized equivalent of \mathbf{A} .

The probability of link formation, P_{ij} depends on $\Delta x_{1ij} = |x_{1i} - x_{1j}|$ and $\Delta x_{2ij} = |x_{2i} - x_{2j}|$ (observed dyad-specific variables) as well as on unobserved individual-level attributes μ_i and μ_j . Typically, let a_{ij}^* defined by

$$a_{ij}^* = \Delta x_{ij}' \bar{\beta} + \mu_i + \mu_j + \varepsilon_{ij}^*,$$

where $\varepsilon_{ij}^* \sim logistic$ distribution. I assume that $A_{ij} = 1$ if $a_{ij}^* > 0$. In that case, P_{ij} is given by

 \leftarrow data.frame(x1 = rnorm(n, 1, 1), x2 = rexp(n, 0.4))

X

data

$$P_{ij} = \frac{\exp\left(\Delta x'_{ij}\bar{\beta} + \mu_i + \mu_j\right)}{1 + \exp\left(\Delta x'_{ij}\bar{\beta} + \mu_i + \mu_j\right)}.$$

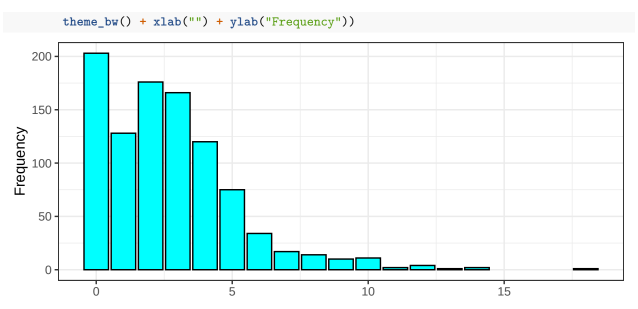
I also assume that μ_i is random. If i is from the s-th sub-network, then $\mu_i \sim \mathcal{N}\left(u_{\mu s}, \sigma_{\mu s}^2\right)$. The mean and the variance of μ_i vary across the sub-networks. This is a way to control for the sub-network heterogeneity as fixed effects in the link formation probability.

```
# Parameter for network model
betanet <-c(-2.8, -1.5)
                             # beta
Glist
         <- list()
                             # adjacency matrix row normalized
Network <- list()</pre>
                             # adjacency matrix row non-normalized
         <- matrix(0, 0, 2) # observed dyad-specific variables
dΧ
         <- list()
                             # unobserved individual-level attribute
mu
         <- runif(M, -1, 1) # mean of uu in each sub-network
uu
sigma2u <- runif(M, 0.5, 4) # variance of uu in each sub-network
# Network
for (m in 1:M) {
  nm
               <- nvec[m]
               <- rnorm(nm, uu[m], sqrt(sigma2u[m]))
  mum
```

```
Z1
                <- matrix(0, nm, nm)
  Z2
                <- matrix(0, nm, nm)
  for (i in 1:nm) {
    for (j in 1:nm) {
      Z1[i, j] <- abs(data$x1[i] - data$x1[j])</pre>
      Z2[i, j] \leftarrow abs(data$x2[i] - data$x2[j])
    }
  }
                <- 1*((Z1*betanet[1] + Z2*betanet[2] +
  Gm
                          kronecker(mum, t(mum), "+") + rlogis(nm^2)) > 0)
                <- 0
  diag(Gm)
  diag(Z1)
                <- NA
  diag(Z2)
                <- NA
                <- Z1[!is.na(Z1)]
  Z2
                <- Z2[!is.na(Z2)]
                <- rbind(dX, cbind(Z1, Z2))
  dX
  Network[[m]] <- Gm</pre>
                <- rowSums(Gm); rs[rs == 0] <- 1
                <- Gm/rs
  Glist[[m]]
                <- Gm
  mu[[m]]
                <- mum
}
               <- unlist(mu)
```

To simulate the dependent variable, I use μ , the vector of μ_i as additional explanatory variable in the count variable model. I set that $Cor(\mu_i, \varepsilon_i) = \rho$ and $Cor(\sum_{j=1}^n G_{ij}\mu_i, \varepsilon_i) = \bar{\rho}$. In that case, $\varepsilon_i = \rho \sigma_{\varepsilon} \tilde{\mu}_i + \bar{\rho} \sigma_{\varepsilon} \bar{\tilde{\mu}}_i + \tilde{\nu}_i$, where $\tilde{\mu}_i = \frac{\mu_i - u_{\mu s}}{\sigma_{\mu s}}$, $\bar{\tilde{\mu}}_i = \sum_{j=1}^n G_{ij}\tilde{\mu}_j$ is the average of $\tilde{\mu}_i$ among *i*'s friends and $\tilde{\nu}_i$ is the new error term following normal distribution of variance $\bar{\sigma}_{\varepsilon}^2 = (1 - \rho^2 - \bar{\rho}^2) \sigma_{\varepsilon}^2$.

```
<- (mu - rep(uu, nvec))/sqrt(rep(sigma2u, nvec))
data$tmu <- tmu
        <- 0.24
rho
rhobar
       <- 0.18
thetanet <- c(lambda, beta, sigma*rho, gamma, sigma*rhobar,
              sigma*(1 - rho - rhobar))
         <- simCDnet(formula = ~ x1 + x2 + tmu | x1 + x2 + tmu,
ytmp
                     Glist = Glist, theta = thetanet, data = data)
         <- ytmp$y
# Add y to data
data$y <- y
# Summarize y
summary(y)
##
      Min. 1st Qu. Median
                              Mean 3rd Qu.
                                              Max.
```



In real life, $\tilde{\mu}_i$ and $\bar{\tilde{\mu}}_i$ are not observed and they are included in the error term ε_i . This implies that the network is endogenous.

5.2 Estimation

When the endogeneity of the network is not taken into account, the peer effect is overestimated. The reason is that the peer effect also captures other effects. The model considers any common shock on $\tilde{\mu}_i$ and $\tilde{\bar{\mu}}_i$ as peer effect. In the example below, the peer effect is estimated at 0.49 when I do not control for the endogeneity of the network.

```
# Count data model
CDexo <- CDnetNPL(formula = y ~ x1 + x2, contextual = TRUE, Glist = Glist,</pre>
                   optimizer = "optim", data = data, npl.ctr = list(print = FALSE))
summary(CDexo)
## Count data Model with Social Interactions
##
## Method: Nested pseudo-likelihood (NPL)
## Iteration:
               22
##
## Network:
## Number of groups
                                3
## Sample size
                                964
## Average number of friends:
                                27.30809
##
##
  Coefficients:
##
                Estimate Std. Error t value Pr(>|t|)
                                        12.92
## lambda
                0.487503
                            0.037720
                                                <2e-16 ***
                                        12.87
                                                <2e-16 ***
## (Intercept)
                1.825234
                            0.141787
## x1
               -1.947378
                            0.042677
                                       -45.63
                                                <2e-16 ***
## x2
                0.803085
                            0.015453
                                        51.97
                                                <2e-16 ***
## G: x1
                1.705766
                            0.076044
                                        22.43
                                                <2e-16 ***
## G: x2
               -1.201023
                            0.026497
                                       -45.33
                                                <2e-16 ***
##
## Marginal Effects:
##
                Estimate Std. Error t value Pr(>|t|)
```

```
## lambda
               0.410015
                          0.032023
                                    12.80
                                            <2e-16 ***
                          0.118774 12.92
## (Intercept) 1.535114
                                            <2e-16 ***
## x1
              -1.637843
                         0.042362 -38.66
                                            <2e-16 ***
## x2
               0.675435
                                   45.74
                          0.014767
                                            <2e-16 ***
## G: x1
               1.434635
                         0.067551
                                    21.24
                                            <2e-16 ***
## G: x2
              -1.010121
                          0.025133 -40.19
                                            <2e-16 ***
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## sigma: 0.9333558
## log pseudo-likelihood: -1139.806
```

To correct the endogeneity, $\tilde{\mu}_i$ and $\tilde{\bar{\mu}}_i$ should be included in the model as additional explanatory variables. However, these variables are not observed.

I use Markov Chain Monte Carlo (MCMC) to estimate the parameters of the dyadic linking model (these include the unobserved individual-level attributes μ_i). The estimation can be done using the function netformation.

I randomly choose some parameters and present their posterior distribution. It stands out that the MCMC converges quickly and the simulations are pretty close to the true values.

```
# plot simulations
par(mfrow = c(4,2), mar = c(2, 2, 2, 1.9))
plot(net\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\poste
                         main = bquote(beta[1]), ylab = "")
abline(h = betanet[1], col = "red")
plot(net\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\posterior\poste
                         main = bquote(beta[2]), ylab = "")
abline(h = betanet[2], col = "red")
plot(net$posterior$mu[,10], type = "l", col = "blue",
                         main = bquote(mu[10]), ylab = "")
abline(h = mu[10], col = "red")
plot(net$posterior$mu[,542], type = "1", col = "blue",
                         main = bquote(mu[542]), ylab = "")
abline(h = mu[542], col = "red")
plot(net$posterior$mu[,10], type = "l", col = "blue",
                         main = bquote(mu[10]), ylab = "")
abline(h = mu[10], col = "red")
plot(net$posterior$mu[,849], type = "1", col = "blue",
                         main = bquote(mu[849]), ylab = "")
abline(h = mu[849], col = "red")
plot(net$posterior$mu[,752], type = "1", col = "blue",
                         main = bquote(mu[752]), ylab = "")
abline(h = mu[752], col = "red")
```

```
plot(net$posterior$sigmamu2[,3], type = "l", col = "blue",
    main = bquote(sigma[mu][3]^2), ylab = "")
abline(h = sigma2u[3], col = "red")
```

Using simulations from the posterior distribution, I can construct good estimators for $\tilde{\mu}_i$ and $\tilde{\mu}_i$ and use them as additional supplementary explanatory variables. In the next example, I use the simulation with the highest posterior density to estimate $\tilde{\mu}_i$ and $\tilde{\bar{\mu}}_i$.

```
<- which.max(net$posterior$log.density)
t
print(t)
            <- net$posterior$mu[t,]
muest
            <- net$posterior$uu[t,]
uuest
sigma2uest <- net$posterior$sigmamu2[t,]</pre>
            <- (muest - rep(uuest, nvec))/sqrt(rep(sigma2uest, nvec))
tmuest
data$tmuest <- tmuest
            <- CDnetNPL(formula = y ~ x1 + x2 + tmuest, contextual = TRUE,
CDendo
                         Glist = Glist, optimizer = "optim", data = data,
                         npl.ctr = list(print = FALSE))
summary(CDendo)
```

The coefficients of $\tilde{\mu}_i$ and $\tilde{\bar{\mu}}_i$ are significant. This confirms that the network is endogenous. Moreover, the peer effect is estimated at 0.40, which is much better than the previous estimate.

The standard errors computed in this example are not valid because they assume that $\tilde{\mu}_i$ and $\bar{\tilde{\mu}}_i$ are observed. To correct the standard error, I replicate simulations of $\tilde{\mu}_i$ and $\bar{\tilde{\mu}}_i$ with replacement from the posterior distribution in order to take into account the variance of the MCMC.

```
<- function(s) {
fendo
  t
            <- sample(3001:8000, 1)
  datat
            <- data
            <- net$posterior$mu[t,]
  mils
            <- rep(net$posterior$uu[t,], nvec)
  uus
            <- rep(net$posterior$sigmamu2[t,], nvec)
  SIIS
            <- (mus - uus)/sqrt(sus)
  datat$tmu <- tmu
  CDnet
           <- CDnetNPL(formula = y ~ x1 + x2 + tmu, contextual = TRUE,
                       Glist = Glist, optimizer = "optim", data = datat,
                       npl.ctr = list(print = FALSE))
  summary(CDnet, Glist = Glist, data = datat)
}
n.cores
           <- 32
           <- 1000 #Number of replications
replic
          <- mclapply(1:replic, fendo, mc.cores = n.cores)</pre>
# The output of out.endo is a list of objects from "summary.CDnetNPL" class
# Let's set the class of out.endo as "summary.CDnetNPLs"
class(out.endo) <- "summary.CDnetNPLs"</pre>
# I can now summarize the results using print
# Object from "summary.CDnetNPL" has a print method
print(out.endo)
## Count data Model with Social Interactions
```

```
## Replication: 1000
##
##
  Coefficients:
                Estimate Std. Error t value Pr(>|t|)
##
## lambda
                0.402133
                            0.040005
                                        10.05
                                                <2e-16 ***
                                        12.75
## (Intercept)
                1.864045
                            0.146209
                                                <2e-16 ***
                                       -47.07
## x1
               -1.935345
                            0.041119
                                                <2e-16 ***
## x2
                0.806629
                            0.014940
                                        53.99
                                                <2e-16 ***
## tmu
                0.337809
                            0.037415
                                         9.03
                                                <2e-16 ***
## G: x1
                1.611186
                            0.078309
                                        20.57
                                                <2e-16 ***
## G: x2
                -1.212639
                            0.026483
                                       -45.79
                                                <2e-16 ***
                0.255580
## G: tmu
                            0.088880
                                         2.88
                                               0.00403
##
## Marginal Effects:
##
                Estimate Std. Error t value Pr(>|t|)
## lambda
                0.338501
                            0.033852
                                        10.00
                                                <2e-16 ***
  (Intercept)
                                        12.74
##
                1.569115
                            0.123193
                                                <2e-16 ***
               -1.629105
                            0.040575
                                       -40.15
                                                <2e-16 ***
## x2
                                        47.32
                0.678992
                            0.014350
                                                <2e-16 ***
## tmu
                0.284351
                            0.031994
                                         8.89
                                                <2e-16 ***
## G: x1
                1.356236
                            0.068747
                                        19.73
                                                <2e-16 ***
## G: x2
                -1.020757
                            0.025080
                                       -40.70
                                                <2e-16 ***
                                               0.00396 **
## G: tmu
                0.215120
                            0.074660
                                         2.88
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## sigma: 0.8792079
## log pseudo-likelihood:
         min
                  mean
                              max
## -1104.369 -1094.693 -1085.772
```

The standard errors are slightly greater than the previous estimate.

5.3 Monte Carlo Simulations

One simulation is not sufficient to confirm that the method used to correct the endogeneity performs well. In this section, I use Monte Carlo simulations to assess the performance of this method. I compare the estimates when the network is assumed exogenous to these when I control for the endogeneity.

I build the following Monte Carlo function which simulates data based on endogenous network, and estimates the model by assuming that the network is exogenous and then endogenous.

```
fMCendo <- function(s) {</pre>
  # parameters
  М
            \leftarrow rep(500, 3)
  nvec
            <- sum(nvec)
  n
  lambda
            <- 0.4
                                  # peer effect
            <-c(2, -1.9, 0.8)
  beta
                                  # own effects
  gamma
            <-c(1.5, -1.2)
                                  # contextual effects
  sigma
            <- 1.5
                                  # standard deviation of epsilon
  theta
            <- c(lambda, beta, gamma, sigma)
            \leftarrow data.frame(x1 = rnorm(n, 1, 1), x2 = rexp(n, 0.4))
  data
```

```
# Network
betanet <-c(-2.8, -1.5)
                             # beta
         <- list()
Glist
                             # adjacency matrix row normalized
Network <- list()</pre>
                             # adjacency matrix row non-normalized
         <- matrix(0, 0, 2) # observed dyad-specific variables</pre>
mu
         <- list()
                             # unobserved individual-level attribute
         <- runif(M, -1, 1) # mean of uu in each sub-network
uu
sigma2u <- runif(M, 0.5, 4) # variance of uu in each sub-network
for (m in 1:M) {
               <- nvec[m]
 nm
               <- rnorm(nm, uu[m], sqrt(sigma2u[m]))
 mum
               <- matrix(0, nm, nm)
 Z1
               <- matrix(0, nm, nm)
 for (i in 1:nm) {
    for (j in 1:nm) {
      Z1[i, j] <- abs(data$x1[i] - data$x1[j])</pre>
     Z2[i, j] \leftarrow abs(data$x2[i] - data$x2[j])
   }
 }
               <- 1*((Z1*betanet[1] + Z2*betanet[2] +
                        kronecker(mum, t(mum), "+") + rlogis(nm^2)) > 0)
 diag(Gm)
 diag(Z1)
               <- NA
               <- NA
 diag(Z2)
               <- Z1[!is.na(Z1)]
 Z2
               <- Z2[!is.na(Z2)]
               <- rbind(dX, cbind(Z1, Z2))
 dΧ
 Network[[m]] <- Gm</pre>
               <- rowSums(Gm); rs[rs == 0] <- 1
               <- Gm/rs
 Glist[[m]]
              <- Gm
 mu[[m]]
               <- mum
}
mu
               <- unlist(mu)
# Data
         <- (mu - rep(uu, nvec))/sqrt(rep(sigma2u, nvec))
data$tmu <- tmu
         <- 0.24
rho
         <- 0.18
thetanet <- c(lambda, beta, sigma*rho, gamma, sigma*rhobar,
              sigma*(1 - rho - rhobar))
         \leftarrow simCDnet(formula = \sim x1 + x2 + tmu | x1 + x2 + tmu,
ytmp
                     Glist = Glist, theta = thetanet, data = data)
         <- ytmp$y
data$y
         <- y
```

```
# Count data model
CDexo <- CDnetNPL(formula = y ~ x1 + x2, contextual = TRUE, Glist = Glist,</pre>
                  optimizer = "optim", data = data, npl.ctr = list(print = FALSE))
# Dyadic linking model
      <- netformation(network = Network, formula = ~ dX, fixed.effects = TRUE,
                       mcmc.ctr = list(burnin = 1000, iteration = 2000), print = FALSE)
# Endogeneity
            <- which.max(net$posterior$log.density)
            <- net$posterior$mu[t,]
muest
            <- net$posterior$uu[t,]</pre>
uuest
sigma2uest <- net$posterior$sigmamu2[t,]</pre>
            <- (muest - rep(uuest, nvec))/sqrt(rep(sigma2uest, nvec))
tmuest
data$tmuest <- tmuest</pre>
CDendo
            <- CDnetNPL(formula = y ~ x1 + x2 + tmuest, contextual = TRUE,
                         Glist = Glist, optimizer = "optim", data = data,
                         npl.ctr = list(print = FALSE))
c(CDexo$estimate, CDendo$estimate)
```

I now run the function fMCendo 1000 times and compute the average of both estimators.

The results confirm that the method used to correct the endogeneity performs well. The average of the estimates is – whereas the true value is –. In contrast, the peer effect is overestimated when the network is assumed exogenous. The average of the estimates is –.

6 Conclusion

This paper provides technical details on the package **CDatanet**. It shows with simple and practical examples how to use the package. It provides comparison of the model with the SART and the SAR models. It also shows with Monte Carlo simulations how the model performs when the network matrix is exogenous and endogenous. Simulation results prove that the peer effect is overestimated when the network is endogenous. Lastly, it presents a way to control for the endogeneity of the network.

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