## Exponential Random Graph models for Little Networks

#### 2 Abstract

Statistical models for social networks have enabled researchers to study complex social phenomena that give rise to observed patterns of relationships among social actors and to gain a rich understanding of the interdependent nature of social ties and actors. Much of this research has focused on social networks within medium to large social groups. To date, these advances in statistical models for social networks, and in particular, of Exponential-Family Random Graph Models (ERGMS), have rarely been applied to the study of small networks, despite small network data in teams, families, and personal networks being common in many fields. In this paper, we revisit the estimation of ERGMs for small networks and propose using exhaustive enumeration when possible. We developed an R package that implements the estimation of pooled ERGMs for small networks using Maximum Likelihood Estimation (MLE), called "ergmito". Based on the results of an extensive simulation study to assess the properties of the MLE estimator, we conclude that there are several benefits of direct MLE estimation compared to approximate methods and that this creates opportunities for valuable methodological innovations that can be applied to modeling social networks with ERGMs.

- 3 Keywords: exponential random graph models, small networks, exact
- 4 statistics, simulation study, teams

#### 5 1. Introduction

- Statistical models for social networks have enabled researchers to study com-
- 7 plex social phenomena that give rise to observed patterns of relationships among
- social actors, and to gain a rich understanding of the interdependent nature of
- 9 social ties and social actors [28, 41]. For example, this research has provided new

insights into the role that the attributes of social actors (e.g., their characteristics, beliefs, and decisions), and endogenous structural processes (e.g., social balance, and relationship reciprocity) play in shaping social networks across different populations and social settings, and how these social networks, in turn, influence individuals and groups.

Much of this research has focused on social networks within medium to 15 large social groups: networks ranging from dozens or hundreds of members (e.g., classrooms and organizations) to millions (e.g., online social networks). However, modern advances in statistical models for social networks have rarely 18 been applied to the study of small networks, despite small network data from teams, families, and personal (ego-centric) networks being common in many 20 fields that study social phenomena [3, 7, 8, 18]. The study of small networks 21 often uses descriptive statistics that summarize basic structural features of the network; for example, the density, degree distribution, or triad count. However, 23 researchers in these fields are often interested in testing hypotheses about why localized social structures, such as reciprocity, balance, and homophily, emerge 25 in these small groups. A key limitation to such work has been the availability of statistical models for networks that can flexibly test and control for the kind of dependencies inherent to network data. In this paper, we propose an approach 28 for applying one of the most widely used statistical models for social networks exponential random graph models, or ERGMs-to small graphs, to enable new 30 research on "little networks".

#### 2. Exponential-Family Random Graph Models

Exponential-family random graph models (ERGMs) are one of the most popular tools used by social scientists to understand social networks and test hypotheses about these networks [13, 19, 35, 42, 47, and others]. In this family of models, an observed graph y, comprised of a set of nodes (vertices) and ties (edges), is characterized by a set of sufficient statistics defined on the graph, s(y), and parameters  $\theta$ . In a model that also includes node characteristics X, this leads to the following equation:

Representation	Description
<b></b>	Mutual Ties (Reciprocity) $\sum_{i \neq j} y_{ij} y_{ji}$
	Transitive Triad (Balance) $\sum_{i \neq j \neq k} y_{ij} y_{jk} y_{ik}$
	Homophily $\sum_{i \neq j} y_{ij} 1 (x_i = x_j)$
$\bigcirc$	Attribute-receiver effect $\sum_{i \neq j} y_{ij} x_j$
	Four Cycle $\sum_{i \neq j \neq k \neq l} y_{ij} y_{jk} y_{kl} y_{li}$

Figure 1: Besides of the common edge count statistic (number of ties in a graph), ERGMs allow measuring other more complex structures that can be captured as sufficient statistics.

$$\Pr\left(Y = y \mid \theta, X\right) = \frac{\exp\left\{\theta^{\mathsf{t}} s\left(y, X\right)\right\}}{\kappa\left(\theta, X\right)}, \quad \forall y \in \mathcal{Y}$$
(1)

Where  $\kappa\left(\theta,X\right)=\sum_{y\in\mathcal{Y}}\exp\left\{\theta^{\mathbf{t}}s\left(y,X\right)\right\}$  is the normalizing constant, and  $\mathcal{Y}$  is the support of the model that is usually assumed to include all graphs of the 41 same type (e.g., directed or undirected) and size, that do not include self-ties. In the directed graph case, the size of  $\mathcal{Y}$  equals  $2^{n(n-1)}$  possible graphs. This makes the exact calculation of  $\kappa(\theta, X)$ , and therefore of (1), computationally expensive. A sophisticated array of parameters can be specified for ERGMs that reflect social and structural process of interest to social scientists, such 46 as social closure, connectivity, and other affiliation preferences. Figure 1 shows some examples of the structures (statistics) that can be estimated with ERGMs. While other methods for studying small graphs exist, e.g. non-parametric 49 tests like the Conditionally Uniform Graph tests (CUG tests in the social net-50 works literature [1, 12] and rewiring algorithms in the network science literature 51 [29, 30]), all in all, ERGMs have more flexibility because they can be used to 52 test complex hypotheses in a multivariate framework. As noted in [5], most of these non-parametric methods can be written in the form of (1), which means 54 that ERGMs can be viewed as a generalized version of many of these tests. 55

56

Although "small networks" is a topic mentioned several times in the liter-

ature on social network models [13, 41, 47], interest in larger social networks has dominated the field. Thus, ERGM methods have been developed to accommodate larger networks (although it is only very recent developments that have begun to scale well to "very large" networks of several thousand nodes or more [44]). One example of this is the calculation of the likelihood function: rather than being calculated using exhaustive enumeration (which we will refer to as "exact likelihood"), the most popular software packages used for estimating these models apply simulation-based estimation methods. As a consequence, current methods used to estimate ERGMs for medium to large networks do not translate well to small network data (i.e., 6 or fewer nodes in a directed network), and applications of these statistical network models to small networks are rare.

One major technical and theoretical issue in ERGM estimation generally, which is exacerbated with small networks, is the problem of non-existence of Maximum Likelihood Estimation (MLE). Non-existence of MLEs (or the convex-71 hull problem) occurs when the observed graph's statistics lie in a region on or 72 near the boundary of the support [2], and can be stressed when estimation depends on Monte Carlo Integration [15]. Small networks, which are more likely to be nearly empty or nearly full, have a smaller region of support, and 75 are more likely to be on or near the boundary of that support. For example, 76 if we are trying to estimate an ERGM in a network with only three nodes, in 77 the scenario where the graph is directed and does not allow for self-ties, the chances of obtaining a graph with either one or zero ties (i.e., empty or almost completely empty), or a graph with five or six ties (i.e., fully or almost fully connected) is about 20% using a uniform sampler.<sup>2</sup> 81

Because researchers studying small networks often have observed *samples* of small networks (e.g., multiple team, family, or personal/egocentric networks), a common work-around to the issue of non-existence of MLE is to combine the in-

82

 $<sup>^{1}\</sup>mathrm{This}$  is perhaps because, as put by [41], small networks are considered to be "uninteresting special cases"

<sup>&</sup>lt;sup>2</sup>For more on the discussion on existence, degeneracy, and instability see [24, 34, 36].

dependent small networks into a single larger block-diagonal graph. Estimation then proceeds by assuming that ties between blocks are impossible (i.e., treated as structural zeros in estimation). The major problems with this approach are that it can be complicated to fit, and difficult to extend. As an example of the former, the same set of constraints (the structural zeros) that allow for the model to be fit can also make the estimation procedure more difficult, and increase the 90 possibility of sampling problems during MCMC estimation. However, a more important challenge with the block-diagonal approach are difficulties with extension. A basic "complete pooling" model, which assumes a common data 93 generating process across all networks, is straightforward to define. However, relaxing that assumption to allow for variability across graphs (i.e., unpooled 95 or partially-pooled models) can be problematic; it would typically require the creation of block-wise node membership attributes, and complex interaction terms involving subgraph statistics and node membership variables. Moreover, extending this framework to not only allow for between-group variability, but to explicitly *predict* it (for example, as a function of additional group-level vari-100 ables), is not straightforward with this complete-pooling approach. 101

To overcome the challenges described above for fitting ERGMs to small 102 networks, we leverage the fact that in the case of small networks, the full likeli-103 hood function is tractable. This allows the direct estimation of model parame-104 ters without using Markov Chain Monte Carlo (MCMC) or other approximate 105 methods, avoiding some of the convergence issues associated with the convexhull problem [15]. It also makes it much easier to combine ERGMs with other 107 statistical techniques, opening the door for many possibilities of richer methods 108 to model and understand small-group network structure and dynamics. In this 109 paper, we describe how modern computational power allows for the complete 110 specification of the likelihood for small graphs, and how this specification al-111 lows us to use the standard tools of MLE, instead of approximate methods. We 112 present examples using these techniques; provide some initial results on empir-113 ical bias, type I error rates, and power based on a simulation study; illustrate 114 the flexibility of this method with an empirical application; and discuss future extensions these techniques make feasible.

117

142

143

144

#### 3. ERGMitos: ERGMs for small networks

With modern computers, calculating the exact likelihood function of an 118 ERGM for a small network becomes computationally feasible. This has an im-119 portant implication: the process for estimating the parameters of an ERGM for 120 small networks can be done directly. Many innovative techniques have been de-121 veloped to handle models with intractable normalizing constants (e.g., Markov 122 Chain Monte Carlo [MCMC] based estimation methods, Bayesian techniques 123 such as the exchange sampler, etc.), and often these techniques work quite well. 124 Of course, no techniques are without tradeoffs; MCMC-based estimation can be 125 sensitive to starting values, and the quality of standard errors can depend on 126 the availability of analytic gradients [32]. Bayesian techniques like the exchange sampler [31] may be comparatively slow, which may be an issue when many 128 networks are to be analyzed. 129

Moreover, simulation-based methods may have particular susceptibilities to 130 the convex-hull problem. As stated by [15, p. 7], "[i]f the model used to simulate 131 the graphs is not close enough to produce realizations that cover the observed 132 values of the statistics, the MC-MLE will not exist even in cases where the MLE 133 does." For example, many common network models, such as triangle-based 134 models, can lead to bimodal distributions of graph statistics that simulation-135 based methods have difficulty with; even when the MLE falls between the modes [22]. Therefore, even though the non-existence issue is not completely avoided, 137 a method based on exact (non-simulation) inference may not only provide a 138 better solution (in general) by avoiding the additional uncertainty induced by 139 simulations and approximations, but it may also help to mitigate the problem 140 in cases where the MLE exists. 141

Of course, the statistical analysis of a single small network could be uninformative due to the small numbers of dyads, and a high restriction in the variability of possible subgraph statistics. Fortunately, research on small networks typically involves collecting data from *samples* of small groups (vs. the more typical 'case studies' of single larger networks), which allows for the development of models to analyze structural variation both within and across small networks. If we assume that the sample of networks comes from a population of networks (groups) that are governed by the same data generating process, we end up with the following likelihood, defining a completely-pooled model:

$$\Pr\left(Y_{1} = y_{1}, \dots, Y_{P} = y_{P} \mid \theta, X_{1}, \dots, X_{p}\right) = \prod_{p=1}^{P} \frac{\exp\left\{\theta^{\mathbf{t}} s\left(y_{p}, X_{p}\right)\right\}}{\kappa_{p}\left(\theta, X_{p}\right)} \quad (2)$$

Where P denotes the number of networks used in the model, and  $\kappa_p(\theta, X_p)$  is explicitly calculated, unlike existing approaches to ERGM estimation. We call this framework, which is a revisited version of ERGM in the case of small networks, ERGM*ito*. In general, this extension can be feasibly applied to small graphs containing at most 6 nodes if directed, or 8 if undirected.

Not to be confused with *pooled estimators* – i.e. aggregating various parameter estimates from independent model fits– pooled-data models have several benefits, including the ability to consider small networks that otherwise would be excluded from an analysis; e.g., because they are fully connected or empty graphs. Moreover, as we will emphasize later in section 5, as long as at least one network in the sample has values on the boundary for each type of sufficient statistic, the MLEs will generally exist [see 15].

One issue that may be of concern is the feasibility of the underlying assumptions when estimating pooled-data models with networks of different sizes. Because parameter estimates often encode network size, one may argue that pooling networks of different sizes into a single model may not be appropriate. However, there are several ways to control for size-induced heterogeneity; for example, including fixed or random effects at the graph level to account for size, or using approaches such as those described in [6, 25, 26]. In the cases presented in this paper, we focus on samples of networks that are of similar sizes (networks of size 4 and 5); thus, these issues are unlikely to be of great concern within a

small range of values, although we demonstrate how they can be accounted for 172 in our applied example (section 6). 173

In the following sections we illustrate and investigate the properties of estimating ERGMs for small networks using this approach. All simulations and 175 model fitting were conducted using the R package ergmito, which has been de-176 veloped to implement the methods described in this paper. 177

#### 4. Illustration with simulated data: fivenets 178

4.1. Data-generating-process and model fitting 179

Starting with a s simple example, we now look at a simulated data set that 180 was created using the data-generating-process of ERGMitos. This particular 181 dataset, which we call "fivenets", is included in the in the R package ergmito<sup>3</sup>. 182 The data set contains five small graphs with nodal attributes (we use gender in the following example), with the networks generated using the following speci-184 fication: 185

$$\Pr\left(Y = y \mid X, \theta\right) = \frac{\exp\left\{\theta_{edges}\left(\sum_{i, j} y_{i j}\right) + \theta_{same}\left(\sum_{i, j} y_{i j} \mathbf{1}\left(X_{i} = X_{j}\right)\right)\right\}}{\kappa\left(\theta, X\right)}$$

where  $\theta_{edges} = -2.0$  and  $\theta_{same} = 2.0$ . Using this equation we draw five networks of size four. The process of "homophily" is represented by a parameter that is defined as the number of ties in which ego and alter have the same gender,  $\theta_{same}$ . Before drawing the networks we randomly generated the node attribute 189 (gender) to each vertex as a Bernoulli with parameter 0.5. Figure 2 shows the 190 generated networks, including their nodal attributes. 191 Using the ergmito R package, we fit three different models to the data: (1) 192

a Bernoulli graph, which is a model that only includes the "edges" parameter, (2) a model with "gender homophily" as its only parameter, and finally (3) a 194

<sup>&</sup>lt;sup>3</sup>The R package is available to be downloaded at BLIND REVIEW.

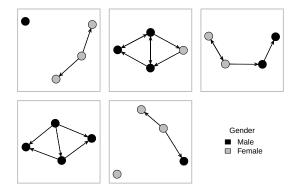


Figure 2: Fivenets data set. These graphs were randomly drawn from an ERGM distribution with two parameters: number of edges and gender homophily, with parameters equal to -2.0 and 2.0 respectively.

model including both "edges" and "gender homophily", which is the correct specification of the model. Some details regarding the computational aspects of the model fitting process are provided in Appendix A.

198

199

200

203

204

In general, while practitioners are accustomed to dealing with a single set of observed sufficient statistics, sometimes called "target" statistics, pooled models instead feature an array of such statistics. Table 1 displays the counts used in this model, from the Fivenets data.

Net id	edgecount	count of gender homophilic ties
1	2	2
2	7	5
3	4	3
4	5	5
5	2	1

Table 1: Observed sufficient for the *fivenets* dataset. In the case of pooled-data models, there is no one set of observed (target) sufficient statistics, but an array of such statistics. This table shows the *edgecount* and the *count of gender homophilic ties* in the *fivenets* dataset.

Table 2 shows the estimation results of the three different specifications of the model and, as expected, model (3) has the best overall fit to the data. Furthermore, since all three models were fitted using MLE, we can compare the edgecount and homophily models with the full model using Likelihood Ratio

### 206 tests [51].

	Homopholy	Edgecount	Full model
Edgecount		$-0.69^*$	-1.70**
		(0.27)	(0.54)
Homophily (on Gender)	-0.12		$1.59^*$
	(0.34)		(0.64)
LR-test statistic $(\chi^2)$	7.04**	13.72***	
AIC	85.06	78.38	73.34
BIC	87.15	80.48	77.53
Log Likelihood	-41.53	-38.19	-34.67
Num. networks	5	5	5

<sup>\*\*\*</sup>p < 0.001; \*\*p < 0.01; \*p < 0.05

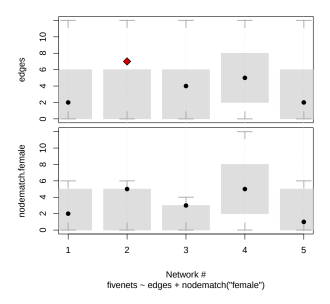
Table 2: Fitted ERGMitos using the fivenets dataset. Looking at AICs and LR-test statistics, the full model (last column of the table) is the one with the best fit to the observed data. More over, the 95% level CI of each covers the true parameters:  $\hat{\theta}_{edges} \in [-2.77, -0.64]$ ;  $\hat{\theta}_{Homophily} \in [0.33, 2.85]$ .

It is important to note that the ergm package can also be used to calculate exact likelihoods, and that this feature has been available for a long time. Some of the additional features and extensions provided in the ergmito package, which are illustrated in subsequent sections of the paper, are: a simple way of estimating pooled-data models, simulating small networks using exact likelihoods, evaluating goodness-of-fit at the graph level for pooled-data models, and including arbitrary effects like interaction effects and transformation of the canonical ERGM terms. The goodness of fit of this model is evaluated in the following section.

#### 4.2. Goodness-of-fit in ERGMitos

Researchers that apply ERGMs should be familiar with the graphical goodness-of-fit (GOF) diagnostics that are used to assess how well the estimated model can reproduce graphs that are similar to the observed graph on a range of local and global graph statistics [21]. In the case of ERGM*itos* applied to small networks, local graph statistics will be more relevant than global statistics to assess GOF. For example, the graph geodesic distribution (i.e., the distribution of

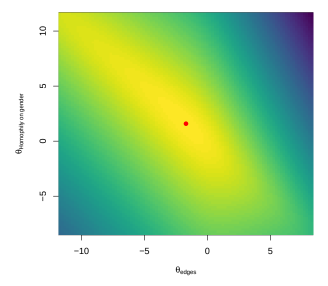
Figure 3: Goodness-of-fit in ERGMitos. This illustrates how the observed sufficient statistics of each one of the 5 networks (x-axis) locate in the overall estimated distribution based on the fitted ERGMito. The gray lines in each box show the minimum and maximum value that the sufficient statistics can take in each one of the 5 networks, whereas the dotted lines provide a 90% confidence interval. The dots are the observed statistics in each network.



shortest-path lengths) is often used to assess GOF for larger networks, but this is clearly less relevant in the case of small networks (like in our case, containing at most 6 nodes if directed, or 8 if undirected) because the shortest-path length between any two nodes typically lies between one and three steps. Therefore, we focus the GOF analysis on the parameters fit in the model as the minimum set of local graph statistics, as shown in Figure 3; and depending on the model complexity a more comprehensive set of local statistics may be needed. An important difference in our approach compared to traditional GOF assessments for ERGMs is that we are able to enumerate the full support of the model, and so instead of showing a boxplot we present a 90% exact confidence interval per-statistic per network, comparing the fitted model's distribution with the observed parameters.

An important advantage of the ERGMitos over "regular" ERGMs is that

Figure 4: Surface of the log-likelihood function of the pooled ERGM*ito* model. Lighter colors represent higher values while darker ones represent lower values. The red dot corresponds to the location of the MLE estimate of the model.



we can observe the surface of the log-likelihood over different combinations of parameters in a rather straightforward way. This, together with the GOF analysis should be a routine step done after every ERGM*ito* fit. Figure 4 shows the surface of the log-likelihood function around the solution parameters to the maximization problem.

The ability to calculate the surface of the exact likelihood function provides additional tools for assessing the quality of the estimated set of parameters. One good use of this diagnostic is to evaluate the roughness of the log-likelihood function, which in principle should give us an idea of the likelihood of the maximization process failing to reach a global maxima, or estimates being close to problematic (e.g., generating empty or fully connected graphs) areas of the parameter space.

#### 5. Simulation study

We conducted two sets of simulations where we compare the performance 249 of the Maximum Likelihood Estimator [MLE] with that of the Monte Carlo 250 MLE [MC-MLE] and Robbins-Monro Stochastic Approximation [RM] in terms 251 of bias, power, type I error rates, and overall computation time. In the first 252 set of simulations, we analyze empirical bias, empirical power, and overall com-253 putation time of each estimator in a scenario where the ERGM is defined by 254 edgecounts and transitive triads. For the second set of simulations, we look at 255 empirical type I error rates when ERGMs are mis-specified by including a tran-256 sitive triad term in the context of a data-generating-process that only includes 257 an edgecount statistic. 258

The code used to reproduce this entire section can be found at BLIND REVIEW.

#### 5.1. Empirical Bias and Power

261

272

273

274

275

276

Using the ERGMito R package, we generated 20,000 samples (datasets), 262 with each sample consisting of several small networks defined by the parameters 263 edges (edgecount) and ttriads (number of transitive triads). Each sample was generated using different combinations of parameters. While all come from an 265 ERGM model defined by edgecounts and number of transitive triads, for every 266 sample we specified: (1) population parameters for the ERGM, (2) the size of 267 the sample (i.e., the number of networks in the sample), and (3) the composition of the sample in terms of the combination of networks of size four and five. A 269 detailed description of each one of these three components used to draw the 270 samples follows: 271

1. **Population parameters**: First we drew two numbers from a piece-wise Uniform distribution with values in  $[-2, -.1] \cup [.1, 2]$ ,  $(\theta_{edges}, \theta_{ttriads})$ , which corresponded to the parameters associated to the statistics **edge-count** and **number of transitive triads**. This specifies the ERGM from which we will draw the networks from. This is akin to the approach taken

by [37], although we took a more conservative approach than their ranges of (-5,0) and (0, 5) for the parameters "edges" and "triangles" in order to increase the number of *irrelevant* draws (i.e., samples composed mostly of either empty of fully connected graphs, or networks with no transitive triads).

- 2. Number of networks per sample Then, we specified the number of networks to generate from the models defined in the previous step, using one of the following sample sizes {5, 10, 30, 50, 100, 150, 200, 300}. The 20,000 simulations were equally split across the various sample sizes (i.e., the simulation study was based on 2,500 samples comprised of 5 networks; 2,500 samples comprised of 10 networks, etc.)
- 3. Number of nodes per network Finally, the composition of each sample, in terms of the number of nodes that each network has, was uniformly-random selected from the pairs  $\{N,0\}, \{N-1,1\}, \ldots, \{1,N-1\}, \{0,N\},$  where the first number of each pair is the number of networks of size 4, and the second is the number of networks of size 5 in the sample. As an example, if the sample size selected in the previous step was 30, then the possible pairs to select from would be  $\{30,0\}, \{29,1\}, \ldots, \{1,29\}, \{0,30\},$  so that samples in which all networks were of size 4 (meaning we draw the pair  $\{30,0\}$ ) or size 5 (again, selecting the pair  $\{0,30\}$ ) were equally likely.

For each one of the 20,000 simulated datasets, we then estimated the model using MLE, as implemented in the *ergmito* R package, and MC-MLE and RM, as implemented in statnet's *ergm* R package [16, 23]. In the case of the latter two, the pooled estimation was done by fitting what is known in the literature as a block-diagonal model in which (a) networks are stacked together in a single adjacency matrix, and (b) the sampling space for the MCMC process is constrained to sample from graphs where ties are only possible within blocks. In the case of the MCMC estimator, we set the control parameters *interval* and samplesize to 2,048, with a burn-in of 2,048 x 16 = 32,768; all double the of the

current default values specified in the *ergm* package, so that we could increase the precision of our estimates. And in cases where the algorithm failed to return any estimates, we increased the control parameters *interval* and *samplesize* to 10,000.

#### 311 5.1.1. Analysis preface

332

333

334

335

After simulating the data and estimating the models, we found that there 312 were several cases in which the programs implementing the three algorithms 313 did not converge, and either returned estimates with a warning to the user, 314 or failed without returning a meaningful message to the user. First, the MLE 315 implementation in ergmito had zero failures, meaning that, even if the optimiza-316 tion failed to converge, the program provided the user with a meaningful report 317 in all cases. Second, while the MC-MLE implementation of the ergm package 318 did fail without returning any form of results in some cases (97 of the 20,000), in 319 each of these instances the program provided the user with a meaningful report 320 of what caused the error. Third, in the case of the Robbins-Monro algorithm 321 [RM], as implemented in the ergm package, we observed a high error rate: in about 25% of the samples, the ergm function failed during the estimation pro-323 cess, and returned an uninformative error message to the user ("NA/NaN/Inf324 in foreign function call (arg 13)"). This error rate should be interpreted with 325 some context; the implementation of the RM algorithm has received less atten-326 tion, and thus less optimization, that the MC-MLE method. While the PNet 327 [46] software provides a more mature implementation of the RM algorithm, we 328 chose to use statuet's implementation as it was better suited for the implemen-329 tation of our simulation study. Table 3 shows the number of errors as a function 330 of sample size (number of networks) for each estimation method. 331

Nearly all of the errors (cases in which the software failed and returned with an error) observed in RM, all but three occur on realizations of the data-generating-process that yielded uninteresting cases, where either of the observed sufficient statistics was on the boundary of their support, e.g. fully connected graphs or graphs with no triads.

	# of errors					
Sample size	MLE	MC-MLE	RM			
5	0	44	1,274			
10	0	21	1,058			
30	0	10	760			
50	0	3	668			
100	0	6	583			
150	0	3	507			
200	0	4	508			
300	0	6	460			
Total	0	97	5,818			

Table 3: Number of times the program failed to fit a model and returned with an error. This shows the overall error rate over the full set of 20,000 simulated samples. All but 3 errors of the RM implementation happened on cases where the sufficient statistics were on the boundary.

With respect to those cases in which the algorithm failed to converge (which 337 includes both software errors and the program reporting lack of convergence), 338 Figure 5 shows the distribution of the sufficient statistic split based on whether 339 the algorithm converged or failed to do so. As shown in the figure, when the algorithms did not converge it was typically due to sufficient statistics falling on the boundary of its support (convex-hull problem). This was especially 342 true for the case of the MLE implementation of ergmito, as all but one of the 343 convergence failures were on the boundary. While MLEs can be obtained in some of those cases (see appendix Appendix B and [15]), in general, estimating such models has no practical utility. We therefore focused our analysis on samples 346 of networks for which the aforementioned model is appropriate: all subsequent 347 analyses include only those data sets where the observed sufficient statistics, 348 edgecounts and number of transitive triads, were not on the boundary of its 349 support for at least one network in the sample. In other words, we included the sample if it: (a) had at least one graph that was not fully connected, and (b) 351 had at least one transitive triad in at least one network. Of the 20,000 simulated 352 data sets, 14,185 met the criteria. 353

Overall, practitioners should bear in mind that the cause of errors that arise during the estimation process can be based on the method or software, and when this is captured by the program it can be informative to both users and developers.

#### 358 5.1.2. Empirical Bias and Power

359

360

363

As shown in Figure 6, all three estimation methods behaved similarly in terms of empirical bias in the models studied here. As the size of the sample of networks in the dataset increased (i.e., when there were more networks within the sample), the empirical bias of all three, MLE, MC-MLE and RM, decreased, as expected.

Looking closer at the biases, we noticed that, while all methods show some kind of bias, MLE has (on average) the smallest. As showed in Table 4, at the 95% confidence level, all three methods tend to overestimate the *edges* parameter. On the other hand, with the exception of the RM method, both MLE and MC-MLE tend to underestimate the *transitive triads* parameter; yet, the RM method has the widest confidence interval for that parameter.

	MLE	MC-MLE	RM
edges	[0.27, 0.36]	[1.23, 1.65]	[0.55, 1.54]
ttriads	[-0.05, -0.03]	[-0.22, -0.16]	[-0.15, 0.48]

Table 4: Empirical bias. Each cell shows the 95% confidence interval of each methods' empirical bias.

Empirical power levels, calculated as the proportion of times that the method 370 reported a significant effect at the 5% level in the same direction as the data-37 generating-process parameter, is depicted in Figure 7. For each method, a 372 single bar in the figure shows the empirical power level for the corresponding 373 combination of sample size (x-axis), parameter (columns), and effect size (rows). 374 There are three main findings to highlight: first, as expected, power increases 375 as both sample size and effect size increase; second, both MLE and MC-MLE 376 behave very similarly with no statistically significant differences across sample 377 and effect size; and third, compared to MLE, RM had a statistically significant 378 smaller power level at various sample and effect sizes combinations, with the largest differences observed on transitive triads Although there may be some

inherent properties of each method that may benefit MLEs, this again may be due to less emphasis on the implementation of RM in the ergm package. 382 Finally, as an anecdotal observation, it is interesting to see that, in the case of effect sizes of magnitude [0.5, 1.0), the discovery rate for the ttriads parameter 384 reaches nearly 0.75 for sample sizes between 30 to 50 networks, which is a rather 385 common sample size in the study of small networks such as teams, families, and 386 sometimes ego-networks. 387

Figure 8 shows the effect of the composition of the sample in each dataset, in terms of the proportion of networks of size five (vs. size four), and the number of networks per dataset. In this case, we observe no meaningful patterns that would indicate the dataset composition is related to power.

380

390

391

392

394

395

397

398

One remarkable difference between the three estimation methods featured by the simulations is the overall computing time needed to fit the models. While the computation of exact likelihoods and gradients is still very computationally intensive, the total time needed to obtain MLEs is still significantly less than what is needed to by the other two methods. As shown in Figure 9, MLE 396 can be orders of magnitude faster than MC-MLE and RM. Therefore, while all three estimators show very similar properties in terms of power and bias, practitioners will benefit by using MLE when modeling small networks because 399 it may substantially reduce computation time. 400

Nevertheless, while MLE is generally faster than the other two methods, 401 there are some scenarios in which the speed gains may not be as dramatic as those shown here. The biggest computational bottleneck that the MLE esti-403 mation faces is the calculation of the full support of the sufficient statistics. In 404 the case of structure-only statistics, ergmito, and actually ergm, computes the 405 full distribution very quickly, but, as the model starts to become more complex, 406 such calculation becomes more and more expensive. Yet, once the full enu-407 meration of the support of sufficient statistics is done, finding MLEs becomes trivial, making the implementation of other statistical tools such as bootstrap 409 or forward/backward model selection feasible to implement. Bootstrapping of 410 ERGMs is illustrated in section 6.

#### 5.2. Type I error rates

412

Using the same procedure described in subsection 5.1, we simulated 35,000 413 datasets comprised of Bernoulli networks (i.e., an ERGM model only defined 414 by the edgecounts sufficient statistic). In this case, we drew different sets of sample sizes: for each of {5, 10, 15, 20, 30, 50, 100} we generated 5,000 datasets 416 using the Bernoulli model with edgecount parameter uniformly distributed in 417 the range  $[-2, -.1] \cup [.1, 2]$ . We then estimated the models using MLE, MC-418 MLE, and RM and calculated the type I error rates using a misspecified model; 419 that is, fitting ERGMs that included a transitive triads count statistic. As with 420 the previous simulations, we only analyze datasets that either had at least one 421 not fully connected graph and had at least one transitivite triad in at least one 422 network. Fortunately, as Table 5 shows, most of the cases did. 423 Table 5 shows the type I error rates per sample size for each of the three methods. In general, MLE report lower error rates compared to MC-MLE and RM, when models were fit to datasets with sample sizes of 20 or fewer networks, 426 the MLE had a better performance than MC-MLE as it reported smaller type I error rates that were much closer to the nominal 5% level. Datasets with 30

the MLE had a better performance than MC-MLE as it reported smaller type I error rates that were much closer to the nominal 5% level. Datasets with 30 or more networks had no significantly different type I error rates between the two methods. Compared to RM, the simulation study shows MLE has a better performance when estimating pooled-data models with 10 or less networks.

No significant difference is observed when dealing with samples of 15 or more networks.

# 434 6. Extended Application: The role of gender-homophily on the formation of small teams

In this final section, we apply the ERGM*itos* framework to a set of observed social networks in an experimental setting. The data was generated as part of a study that examined the emergence of social networks in small teams.

The analytic sample consists of 31 small mixed-gender teams that include either four (17 teams) or five members (14 teams). Individuals recruited for the

-		P(Type I error)			$\chi^2$ (vs	MLE)
Sample size	N. Sims.	MLE	MC-MLE	RM	MC-MLE	RM
5	4,325	0.066	0.086	0.086	11.36 ***	11.36 ***
10	4,677	0.063	0.078	0.073	8.44 ***	3.73 *
15	4,818	0.060	0.072	0.063	5.50 **	0.41
20	4,889	0.054	0.065	0.061	5.30 **	2.05
30	4,946	0.053	0.059	0.055	1.60	0.07
50	4,987	0.053	0.055	0.047	0.16	1.67
100	4,999	0.054	0.054	0.050	0.00	0.81

Table 5: Empirical Type I error rates. The  $\chi^2$  statistic is from a 2-sample test for equality of proportions, and the significance levels are given by \*\*\* p < 0.01, \*\*\* p < 0.05, and \* p < 0.10.

study were University students, participating for research credit or compensa-441 tion, who were assigned to the teams with two conditioning factors: (1) they 442 did not know the other teammates, and (2) there must be at least one team member who identified as male, and one who identified as female. On average, 444 55% of each team's members were female, with no statistically significant differ-445 ence between the teams (test of equal proportions) nor within the teams when 446 compared to a null of 0.5 (exact binomial test). Each team met face-to-face in 447 a laboratory setting to complete about one hour of group tasks. Immediately after the completion of the group tasks, the team networks were measured using 449 name generators administered in an online survey (that was completed in the 450 lab). Advice seeking was one relationship measured, via the question "Who did 451 you go to for advice, information, or help to complete the group tasks?", and 452 participants could select as many or as few teammates as they liked. These data 453 were used to generate directed graphs that represent the advice-seeking network 454 in each team, where  $y_{ij} = 1$  if i identified j as someone they sought advice from. 455 One research question of interest in the field of team science is what is the 456 role of gender and gender-based homophily (i.e., the preference for individuals to 457 form social ties with teammates who match them on gender) in the formation of team networks. Using the ergmito R package to model the team advice networks 459 and test hypotheses about gender and network dynamics, we illustrate how 460 exact calculation of ERGM likelihoods can be leverage to go beyond traditional 461

ERGM analysis. Overall, the analysis consists of two parts: (1) building a baseline model that only includes structural features of the graph, and (2) using that model to test if gender-homophily is a prevalent feature of the data, while also controlling for other gender-based terms in a multivariate fashion.

In the structural-terms-only model, we fitted five different models based on the following terms:

468

469

470

471

472

473

474

475

- Edge count (edges): This accounts for the overall density of the graph and is usually compared to that of a constant term in regression analyses. This is calculated as  $\sum_{ij} y_{ij}$ .
- Number of transitive triads (ttriads): This statistic, also known as balanced triangles or transitive triples, captures the phenomenon of social clustering and balance; where "the friend of my friend is my friend". In this context it indicates that "the advisor of my advisor is my advisor". This term is calculated as follows:  $\sum_i \sum_{j < k} y_{ij} y_{jk} y_{ik}$ .

To illustrate the flexibility of estimating ERGMs with the ergmito R package, 476 we generated three additional terms to be included in the models using the edges 477 and ttriads terms. First, we included two interaction effects, one per term, with 478 an indicator variable which equals to one if the corresponding network was of size 479 five, and zero if it was size four. We also added an offset term as that proposed 480 by [25] which has the nice property of being size-invariant; i.e., it preserves the 481 mean degree as the network size increases. All of these additional terms allowed 482 us to control for differences as a function of the network size. A valuable benefit 483 of these additional statistics is that users can add interaction effects or variable 484 transformations to the models; a feature that, currently, is not easily achieved in 485 other available frameworks (see for example [17, 20]). Just like we showed earlier in Table 1, Table 6 shows an example of the target statistics used in the models 487 for 6 of the 31 networks (i.e., the array of observed sufficient statistics). With 488 these five statistics we estimated five different models, including a bootstrapped 489 version of the one with the best overall fit. Table 7 shows the results. 490

(1)	(2)	(3)	(4)	(5)	(6)
Size $(n)$	edges	ttriads	$\begin{array}{c} \text{edges} \times \\ 1 \left( n = 5 \right) \end{array}$	$ttriads \times 1 (n = 5)$	$\frac{\text{edges} \times}{\log 1/n}$
4	10	14	0	0	-13.86
4	6	2	0	0	-8.32
4	4	0	0	0	-5.55
5	6	1	6	1	-9.66
5	8	8	8	8	-12.88
5	6	2	6	2	-9.66
			. 25 more rows		

Table 6: Example of observed sufficient statistics for the team advice networks. Pooled-data ERGMs have multiple observed sufficient statistics (also known as target statistics). Furthermore, as shown here, we can manipulate common statistics as *edges* (2) and *ttriads* (3) to include, e.g. interaction effects (4) and (5), or more complex transformations, e.g. (6).

The results, Table 7, indicate that transitive triads (ttriads) were more preva-491 lent than expected by chance; which is common in positive affiliation and col-492 laboration networks. Parameter estimates for the ttriads term were also robust with significant and positive effects across the different model specifications. 494 Second, we found that controlling for size of the network mattered. The results 495 of models (3) and (4) show that allowing networks of size 5 to have different 496 parameters associated with number of edges or transitive triads (with networks 497 of size 4 as a reference), significantly improved model fit relative to model (1). Yet, as shown in model (5), these interaction effects were not jointly significant. 499 Regarding model (2), which includes the offset edges  $\times \log 1/n$ , we see that the 500 edges parameter flips from negative 0.72, to positive 0.73, which should be in-501 terpreted in the context of this offset change. For example, in the case of the 502 Bernoulli model, the probability of an individual tie for a network of size 4 would be  $logit^{-1}(-log 4 + 0.73) \approx logit^{-1}(-0.66) \approx 0.34$ , i.e. less than 0.5 which is 504 the expected value under the null. 505 Of the five models, model (3) had the best overall fit, the lowest AIC and 506 BIC, and so it was retained as the structural baseline model for the subsequent 507 analyses. To finalize this first stage of analysis, we calculated the standard errors of model (3) using bootstrap [see 49]; with the results reported in column (3b).

	(1)	(2)	(3)	(4)	(5)	(3b)
edges	-0.72***	0.73***	-0.53***	-0.85***	-0.56*	-0.53***
	(0.13)	(0.13)	(0.15)	(0.14)	(0.23)	(0.12)
ttriad	0.29***	0.33***	0.36***	0.50***	0.38***	0.36***
	(0.05)	(0.05)	(0.06)	(0.07)	(0.11)	(0.05)
$edges \times 1 (n = 5)$			-0.53***		-0.49	-0.53***
			(0.12)		(0.28)	(0.12)
$ttriad \times 1 (n = 5)$				-0.22***	-0.02	
				(0.05)	(0.12)	
offset						
$edges \times \log(1/n)$		Yes				
AIC	651.38	641.02	637.28	640.40	639.26	637.28
BIC	659.74	649.39	649.83	652.95	655.99	649.83
Log Likelihood	-323.69	-318.51	-315.64	-317.20	-315.63	-315.64
Num. networks	31	31	31	31	31	31
Time (seconds)	0.55	0.99	0.74	0.76	0.74	10.12
N replicates						1000
N Used replicates						1000

 $<sup>^{***}</sup>p < 0.001, \, ^{**}p < 0.01, \, ^{*}p < 0.05$ 

516

517

518

519

520

521

522

523

Table 7: Structural models. Model (2) includes Krivitsky et al (2011) offset term. Besides of the common GOF statistics, the table includes the number of networks used, elapsed time to fit the model, and, in the case of Model (3b) which is a bootstrapped version of model (3), number of replicates fitted and included in the bootstrap variance estimate.

This final model had no meaningful changes in standard errors compared to (3); although they were slightly smaller compared to MLEs in (3). Additionally, the elapsed time for this bootstrapping process was negligible: remarkably, we fit 1,000 ERGMs in about 10 seconds, which further highlights how speed and model specification-flexibility are key features of fitting ERGMs using Maximum Likelihood.

The second phase of model specification, which uses model (3) as baseline, focused on evaluating the role of gender and gender-homophily in the advice networks, using the following terms:

- Gender homophily: This term equals to the number of ties in which ego and alter are matched on gender. This was calculated as:  $\sum_{ij} y_{ij} \mathbf{1} (X_i = X_j)$ , where  $X_i$  is one if i is a female, and zero otherwise.
- Female-sender effect: This term, also known as attribute-activity effect, captures the propensity of females to send ties. Is is calculated as:

 $\sum_{ij} y_{ij} X_i$ .

524

525

542

543

546

547

548

551

552

• Female-receiver effect: This term captures the propensity of females to receive ties. It is calculated as:  $\sum_{ij} y_{ij} X_j$ .

Ttaking advantage of the flexibility that the *ergmito* package, and ultimately, 527 using exact likelihoods provides, we also explored modifying the model by means 528 of transformations and offset terms. First, with the purpose of improving the 529 predictive capability of our model, we included the square root of the count of gender-homophilic ties. Other transformations such as interactions with other 531 terms, or centering around a given constant (for example, some population 532 average) could also be implemented. Second, while not the case in our data, 533 we illustrate a hypothetical scenario where the teams had to have at least 5 534 ties, and we constrained the support of the sufficient statistics to only include networks with five or more ties. We did this by using an offset parameter that 536 equaled  $-\infty$  if the network had four or less ties, and zero otherwise. Figure 10 537 illustrates the differences between the Cumulative Distribution Function (CDF) 538 associated with edges statistic (the probability of observing up to given number 539 of ties, x-axis) calculated from a model with (red line) and without (blue line) the constrained space. 541

Using offset terms to constraint the support of the model is not a new thing. The ergm package features this capability as well, in addition to specialized algorithms to constrain samplings space. Users can also set offsets to  $-\infty$  to forbid some configurations, yet, in the case of ergmito combining offset terms with the capability of mixing-transforming variables in the model provides the user with greater flexibility. As we did before, an example of six of the 31 networks is shown in Table 8.

Like in the first round of ERGMs, the standard errors of the final best model were re-calculated using bootstrap. Table 9 shows the results.

As illustrated in Table 9, in our first three specifications we found no evidence that gender-homophily was a prevalent feature of the advice networks, as the baseline (1), its constrained version (2), and the baseline including a transformed

(1) n	(2) Homophily (gender)	(3) Receiver (female)	(4) Sender (female)	(5) Homophily <sup><math>1/2</math></sup>
4	3	5	6	1.73
4	1	4	3	1.00
4	3	4	3	1.73
5	2	2	4	1.41
5	4	7	5	2.00
5	3	4	3	1.73
		25 more r	rows	

Table 8: Example of observed sufficient statistics for the team advice networks (bis). For the second set of ERGMs, we included gender-based effects: homophily (2), receiver (3), and sender (4). Variable (5) is the square root of variable (2).

version of gender-homophily (3) failed to reject the null  $\theta_{\text{Homophily}} = 0$ . Of the other gender-based effects, only the female-sender effect, model (4), was 555 significant. With a coefficient equal to 0.46, the model indicates that, compared 556 to males, females tended to nominate more of their team members as people 557 they sought advice from. Furthermore, we found that the term Sender (female) 558 was a confounder of gender-homophily, with the latter changing from -0.03 in 559 model (1), to -0.12 when the female-sender effect is included. Overall, these final models indicate that the team networks are best explained by preferences 561 for balanced advice-seeking triads, and a tendency for females to seek advice 562 from more of their teammates, compared to males. 563

With Model (4), Table 9, having the best fit overall (smallest AIC and BIC), we re-calculated its standard errors using bootstrapp, model (4b) with the elapsed time, again, remarkably short (~84 seconds to fit a thousand models). While model (4) took roughly five seconds to be fitted, most of the computation time lies on calculating the support of the space of sufficient statistics. Once the support of the sufficient statistics has been calculated, the optimization takes only a fraction of the time, which is why the bootstrap version of model (4) took about 0.09 seconds per repetition, and not 5.27 as the user may have expected. Details on the computational resources used for this section and the simulation studies are shown in Appendix C.

565

566

567

568

570

571

572

573

574

As part of the ad hoc diagnostics, Figure 11 shows the distribution of the suf-

	(1)	(2)	(3)	(4)	(5)	(4b)
edges	-0.52**	-0.91***	-0.54**	-0.72***	-0.48*	-0.72***
-	(0.17)	(0.23)	(0.18)	(0.19)	(0.19)	(0.17)
ttriads	0.36***	0.46***	0.37***	0.36***	0.36***	0.36***
	(0.06)	(0.06)	(0.06)	(0.06)	(0.06)	(0.05)
Homophily (gender)	-0.03	-0.01	-0.20	-0.12	-0.01	-0.12
	(0.20)	(0.21)	(0.46)	(0.20)	(0.20)	(0.20)
$edges \times 1 (n = 5)$	-0.53***	-0.47**	-0.52***	-0.53***	-0.53***	-0.53***
	(0.12)	(0.16)	(0.13)	(0.13)	(0.12)	(0.13)
$(Homophily)^{1/2}$			0.54			
			(1.32)			
Sender (female)				$0.46^{*}$		0.46*
				(0.18)		(0.18)
Receiver (female)					-0.08	
					(0.18)	
Constraint (offset)						
edge > 4		Yes				
AIC	639.26	569.93	641.08	634.68	641.07	634.68
BIC	655.99	586.66	661.99	655.59	661.98	655.59
Log Likelihood	-315.63	-280.96	-315.54	-312.34	-315.53	-312.34
Num. networks	31	28	31	31	31	31
Time (seconds)	2.26	2.32	2.28	5.10	5.19	83.97
N replicates						1000
N Used replicates						1000
***. < 0.001 **. < 0.0	01 * .005					

<sup>\*\*\*</sup>p < 0.001, \*\*p < 0.01, \*p < 0.05

Table 9: Testing for gender homopholy. Models (1) through (3) include either an interaction term or a transformation of the term *Homphily (gender)*. Models (4) and (5) include female sender and receiver effects, while model (4b) is a bootstrapped version of model (4). Model (2) constraints the sample space by setting an offset restricting the support to networks with at least 5 edges. Furthermore, since three of the 31 teams had less than five ties, these were excluded from the analysis, hence (2) includes 28 of the 31 available networks.

ficient statistic under the fitted model, 95% exact confidence intervals (CI), versus the observed set of sufficient statistics. With the exception of two networks one that is a full graph and another that only has one tie—the CIs generated by model (4) are able to cover all other network and term combinations.

We further discuss our results in the following section.

#### 7. Discussion

579

In this paper we revisit and extend Exponential Family Random Graph Models (ERGM) for the case of small networks. Given the interest in testing hypotheses about small networks in the literature, but limited application of statistical models to small network data [13, 19, 35, 42, 47, and others], we

shed new light to ERGMs for small networks, which we call ERGMitos. An appealing feature of ERGMitos is that it allows direct use of the full likeli-586 hood, with all that that entails, rather than costly approximations via MCMC methods. Overall, this approach provides a couple of important benefits for 588 small network data: (1) it increases the chances of obtaining estimates in the 589 case that the observed sufficient statistics are near the boundary of its support 590 (the convex-hull problem); as suggested by our simulation studies, (2) it has the 591 potential to improve power and reduce type I error rates, compared to MC-MLE 592 and the Robbins-Monro stochastic approximation; and (3) ERGMitos can be 593 significantly faster to estimate (orders of magnitude faster), which in turn makes 594 methods like bootstrap or other computationally intensive algorithms immedi-595 ately available to be used with ERGMs, which to date has been unthinkable.

598

601

604

607

611

614

Another major benefit of using the full likelihood directly, when feasible, is that it gives researchers tremendous flexibility in terms of constructing and estimating new models. In terms of estimation, for example, the ability to easily 599 calculate the likelihood allows researchers to make use of standard tools and 600 techniques for ML estimation and MCMC estimation. This is important, because current techniques for intractable models (e.g., auxiliary variable MCMC, 602 MC-MLE, and noise-contrastive estimation), while effective, are not necessarily 603 straightforward to implement for non-experts. This places a high barrier to entry for researchers who would like to develop statistical models that go beyond 605 the standard packages. As a simple example, take recent work on multilevel network models [40]. In that work, constructing a multilevel Bayesian model of a sample of networks, while conceptually straightforward, required the de-608 velopment of custom code and algorithms to implement. By contrast, having 609 the full likelihood available in R means that the same models studied in that 610 paper (assuming small-N networks) can be constructed and estimated as easily as any other non-intractable model for which we can calculate the likelihood, using the full range of traditional tools and algorithms for ML and Bayesian 613 estimation. This frees researchers from focusing only on models that are implementable in packages like *statnet*, and allows greater freedom to think about ways that models for graphs can be modified and incorporated into other statistical models. In addition, being able to estimate gradients opens the possibility
of estimating models using modern Bayesian algorithms like Hamiltonian Monte
Carlo (HMC) and stochastic gradient langevin dynamics (SGLD), which may
offer advantages in terms of speed or scalability, respectively.

The development and evaluation of ERGM itos in this simulation study also 621 brings up topics for future work. One is the evaluation of model goodness-of-622 fit, and identifying statistics that are most important to evaluate with small networks, and that are reasonable to expect in a model that would suggest 624 "good fit". Because ERGMitos enable a rather simple way of conducting 625 simulation studies (relative to traditional ERGMs), this will facilitate this work 626 in future. Another topic to explore in future work is the value of ERGM itos for 627 estimating ERGMs for very large networks, by drawing samples of local network structures from a large graph. There is ongoing work extending ERGMs to very 629 large networks [43, 44], and ERGMitos could be a valuable approach for fitting 630 these pooled models to a large sample (e.g., in the order of the thousands) 631 of small local network structures drawn from a large network. Although this 632 is an exciting extension to explore, this must proceed cautiously; while some 633 matters as the "projectivity problem" [39] are solvable [25, 26, 38], size-invariant 634 ERGMs (models in which the parameter estimates are scalable across graph 635 sizes), is an area of research very much under development. 636

In sum, ERGM*itos* provide a promising extension to the ERGM framework for the analysis of small social networks. In addition to all the theoretical benefits that using exact likelihoods carry with it [15], features related to computational efficiency and flexibility open the door to new *and* old statistical tools that have been unreachable in the ERGM framework. Ultimately, as a fundamental building block of larger social systems, a richer understanding of the local social processes that give rise to the formation of networks in small social groups is key for our understanding of larger social structures that these constitute.

Figure 5: Distribution of the average sufficient statistics per sample. Since samples can contain networks of sizes four and five, we have re-scaled the sufficient statistics counts by each network size's corresponding maximum value so these range from zero to one. Most of the cases in which methods failed to converge happened in scenarios where either all the graphs in the sample were fully connected or there was no transitive triad; exactly the cases that we excluded for the reminder of the analysis.

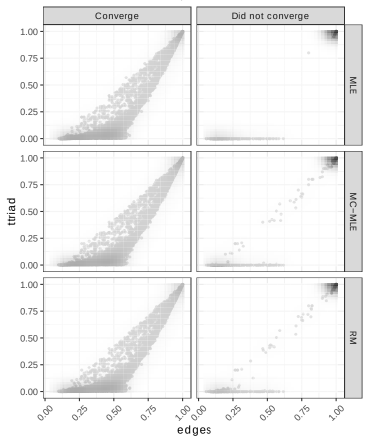


Figure 6: Empirical distribution of the bias per model parameter, for MC-MLE and MLE estimation methods. In general we see that the parameter estimates' bias is centered around zero and both MC-MLE (ERGM) and MLE (ERGMito) have about the same bias in our simulation study.

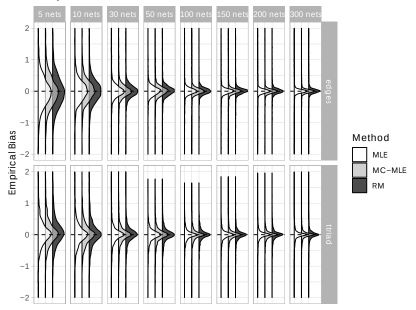


Figure 7: Empirical power by dataset size and effect size (the later considering only magnitude), for ERGM and ERGMito estimation methods. Power increases for both MC-MLE (ERGM) and MLE (ERGMito) with increases in the size of the dataset and effect size. There are indistinguishable differences in power between the two estimation methods.

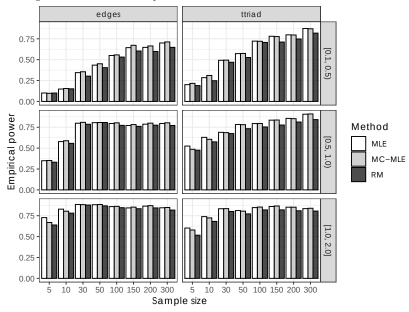


Figure 8: Empirical power by proportion of networks of size five in the sample (color coded) and sample size (rows).

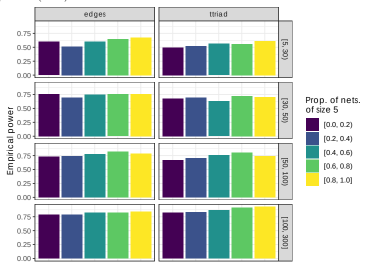
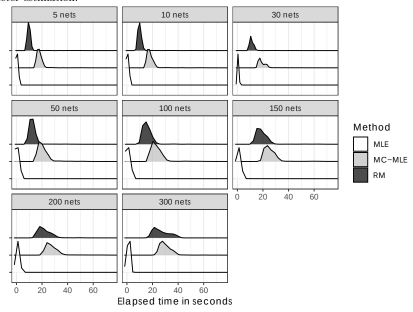


Figure 9: Distribution of elapsed time (in seconds) for the estimation process for MC-MLE (ERGM) versus MLE (using  $\mathrm{ERGM}ito$ ). Overall, the MLE implementation is orders of magnitude faster compared to the time required by the MC-MLE implementation to do the parameter estimation.



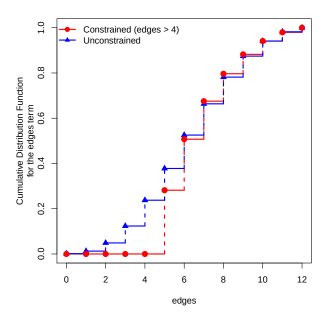


Figure 10: Marginalized Cumulative Distribution Function (CDF) for the *edges* sufficient statistic for a network of size 4 (up to 12 ties). The blue line shows the CDF for the edges term according to model (1) in Table 9, which was fitted without constraining the support of the sufficient statistics, while the red line shows the CDF of model (2), also in Table 9, which constrained the support to networks with at least 5 ties. Once again, since we can fully enumerate the support, both CDFs are exact, and thus, not simulated.

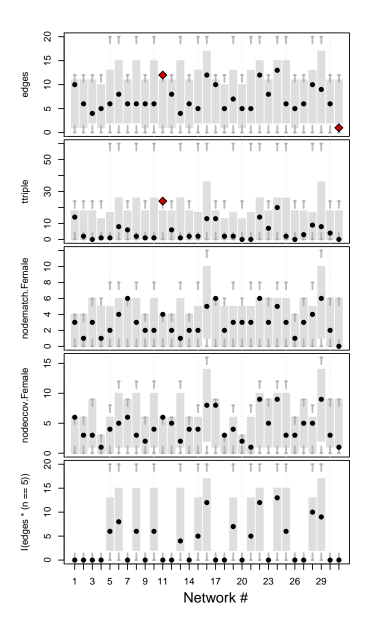


Figure 11: Distribution of the sufficient statistics under the ERGM specified by the parameters from model (4) in Table 9. Each bar represents the exact 95% confidence interval for the corresponding network+term combination, while the black dots show the location of the corresponding observed statistic. Red diamonds mark the observed statistics that fall out of the 95% confidence interval. Of the 31 networks in the sample, it is only in two networks that the CIs don't cover the observed statistic, one that is fully connected and another that only has one tie.

#### References

655

656

657

- [1] Brigham S Anderson, Carter Butts, and Kathleen Carley. "The interaction of size and density with graph-level indices". In: Social Networks 21.3 (July 1999), pp. 239-267. DOI: 10.1016/S0378-8733(99)00011-8. URL: http://www.sciencedirect.com/science/article/pii/S0378873399000118%20https://linkinghub.elsevier.com/retrieve/pii/S0378873399000118.
- O. Barndorff-Nielsen. Information and Exponential Families in Statistical Theory. 2014. ISBN: 9781118857281. DOI: 10.1002/9781118857281.
  - [3] E. Bott and E.B. Spillius. Family and Social Network. Families & Marriage. Routledge, 2001. ISBN: 9780415264174. URL: https://books.google.com/books?id=XVwMqHb56TsC.
- Carter T. Butts. sna: Tools for Social Network Analysis. R package version 2.5. 2019. URL: https://CRAN.R-project.org/package=sna.
- Carter T. Butts. "Social network analysis: A methodological introduction". In: Asian Journal Of Social Psychology 11.1 (Mar. 2008), pp. 13–41. ISSN: 1367-2223. DOI: 10.1111/j.1467-839X.2007.00241.x. URL: http://doi.wiley.com/10.1111/j.1467-839X.2007.00241.x.
- Carter T. Butts and Zack W. Almquist. "A Flexible Parameterization for Baseline Mean Degree in Multiple-Network ERGMs". In: Journal of Mathematical Sociology 39.3 (2015), pp. 163–167. DOI: 10.1080/0022250X.
   2014.967851.
- Dorothy R. Carter et al. "Social network approaches to leadership: An integrative conceptual review." In: Journal of Applied Psychology 100.3 (May 2015), pp. 597–622. ISSN: 1939-1854. DOI: 10.1037/a0038922. URL: http://doi.apa.org/getdoi.cfm?doi=10.1037/a0038922.
- [8] N. Crossley et al. Social Network Analysis for Ego-Nets: Social Network
  Analysis for Actor-Centred Networks. SAGE Publications, 2015. ISBN:
  9781473927339. URL: https://books.google.com/books?id=v0IdCAAAQBAJ.
- [9] Leonardo Dagum and Ramesh Menon. "OpenMP: an industry standard API for shared-memory programming". In: *IEEE computational science* and engineering 5.1 (1998), pp. 46–55.
- Dirk Eddelbuettel and Romain François. "Rcpp: Seamless R and C++
  Integration". In: Journal of Statistical Software 40.8 (2011), pp. 1–18. DOI:
  10.18637/jss.v040.i08. URL: http://www.jstatsoft.org/v40/i08/.
- Dirk Eddelbuettel and Conrad Sanderson. "RcppArmadillo: Accelerating R with high-performance C++ linear algebra". In: Computational Statistics and Data Analysis 71 (Mar. 2014), pp. 1054–1063. URL: http://dx.doi.org/10.1016/j.csda.2013.02.005.

- Katherine Faust. "A puzzle concerning triads in social networks: Graph constraints and the triad census". In: Social Networks 32.3 (2010), pp. 221–233. DOI: 10.1016/j.socnet.2010.03.004. URL: http://dx.doi.org/10.1016/j.socnet.2010.03.004.
- O Frank and David Strauss. "Markov graphs". In: Journal of the American Statistical Association 81.395 (1986), pp. 832–842. DOI: 10.2307/2289017.
   URL: http://amstat.tandfonline.com/doi/abs/10.1080/01621459.
   1986.10478342.
- Jeff Gill and Gary King. "What to Do When Your Hessian is Not Invertible: Alternatives to Model Respecification in Nonlinear Estimation".
   In: Sociological Methods & Research 33.1 (2004), pp. 54–87. DOI: 10.1177/0049124103262681. URL: https://doi.org/10.1177/0049124103262681.
- Mark S. Handcock. "Assessing Degeneracy in Statistical Models of Social Networks". In: Working Paper No. 39 76.39 (2003), pp. 33-50. ISSN: 1936900X. DOI: 10.1.1.81.5086. URL: http://citeseerx.ist.psu.edu/viewdoc/summary?doi=10.1.1.81.5086.
- Mark S. Handcock et al. ergm: Fit, Simulate and Diagnose Exponential-Family Models for Networks. R package version 3.9.4. The Statnet Project (http://www.statnet.org). 2018. URL: https://CRAN.R-project.org/package=ergm.
- Mark S. Handcock et al. ergm.userterms: User-specified Terms for the statnet Suite of Packages. R package version 3.10.0. The Statnet Project (https://statnet.org). 2019. URL: https://CRAN.R-project.org/package=ergm.userterms.
- Top [18] Kaisa Henttonen. "Exploring social networks on the team level-A review of the empirical literature". In: Journal of Engineering and Technology Management 27.1 (2010), pp. 74–109. ISSN: 0923-4748. DOI: https://doi.org/10.1016/j.jengtecman.2010.03.005.
- Paul W. Holland and Samuel Leinhardt. "An exponential family of probability distributions for directed graphs". In: *Journal of the American Statistical Association* 76.373 (1981), pp. 33–50. DOI: 10.2307/2287037.
- [20] David R. Hunter, Steven M. Goodreau, and Mark S. Handcock. "ergm.userterms:
   A Template Package for Extending statnet". In: *Journal of Statistical Software* 52.2 (2013), pp. 1–25.
- 719 [21] David R Hunter, Steven M Goodreau, and Mark S Handcock. "Goodness of Fit of Social Network Models". In: *Journal of the American Statistical*721 *Association* 103.481 (2008), pp. 248–258. DOI: 10.1198/016214507000000446.
  722 eprint: https://doi.org/10.1198/016214507000000446. URL: https:
  723 //doi.org/10.1198/016214507000000446.

- David R. Hunter, Pavel N. Krivitsky, and Michael Schweinberger. "Computational Statistical Methods for Social Network Models". In: Journal of Computational and Graphical Statistics 21.4 (2012). PMID: 23828720, pp. 856–882. DOI: 10.1080/10618600.2012.732921. eprint: https://doi.org/10.1080/10618600.2012.732921. URL: https://doi.org/10.1080/10618600.2012.732921.
- [23] David R. Hunter et al. "ergm: A Package to Fit, Simulate and Diagnose Exponential-Family Models for Networks". In: *Journal of Statistical Software* 24.3 (2008), pp. 1–29.
- 733 [24] Martin Jacobsen. "Existence and Unicity of MLEs in Discrete Exponential Family Distributions". In: Scandinavian Journal of Statistics 16.4 (1989), pp. 335–349. ISSN: 03036898, 14679469. URL: http://www.jstor.org/stable/4616145.
- Pavel N. Krivitsky, Mark S. Handcock, and Martina Morris. "Adjusting for network size and composition effects in exponential-family random graph models". In: Statistical Methodology 8.4 (July 2011), pp. 319-339.

  ISSN: 15723127. DOI: 10.1016/j.stamet.2011.01.005. URL: https:
  //linkinghub.elsevier.com/retrieve/pii/S1572312711000086.
- Pavel N. Krivitsky and Eric D. Kolaczyk. "On the Question of Effective Sample Size in Network Modeling: An Asymptotic Inquiry". In: Statistical Science 30.2 (May 2015), pp. 184–198. ISSN: 0883-4237. DOI: 10.1214/14-STS502. URL: http://projecteuclid.org/euclid.ss/1433341477.
- Philip Leifeld. "texreg: Conversion of Statistical Model Output in R to LATEX and HTML Tables". In: Journal of Statistical Software 55.8 (2013), p. 24. URL: http://www.jstatsoft.org/v55/i08/.
- D. Lusher, J. Koskinen, and G. Robins. Exponential Random Graph Models for Social Networks: Theory, Methods, and Applications. Structural Analysis in the Social Sciences. Cambridge University Press, 2012. ISBN: 9781139851039.
- 753 [29] Ron Milo et al. "Superfamilies of Evolved and Designed Networks". In:

  Science 303.5663 (Mar. 2004), pp. 1538-1542. DOI: 10.1126/science.

  1089167. URL: http://www.sciencemag.org/cgi/doi/10.1126/

  science.1089167.
- R Milo et al. "On the uniform generation of random graphs with prescribed degree sequences". In: *Arxiv preprint condmat0312028* cond-mat/0 (2004), pp. 1–4. arXiv: 0312028 [cond-mat]. URL: http://arxiv.org/abs/cond-mat/0312028.
- J. Møller et al. "An efficient Markov chain Monte Carlo method for distributions with intractable normalising constants". In: *Biometrika* 93.2 (June 2006), pp. 451-458. ISSN: 0006-3444. DOI: 10.1093/biomet/93.2.451. eprint: http://oup.prod.sis.lan/biomet/article-pdf/93/2/451/590012/932451.pdf. URL: https://doi.org/10.1093/biomet/93.2.451.

- 767 [32] Jaewoo Park and Murali Haran. "Bayesian Inference in the Presence of Intractable Normalizing Functions". In: Journal of the American Statistical Association 113.523 (2018), pp. 1372–1390. DOI: 10.1080/01621459. 2018. 1448824. eprint: https://doi.org/10.1080/01621459.2018.1448824. URL: https://doi.org/10.1080/01621459.2018.1448824.
- 772 [33] R Core Team. R: A Language and Environment for Statistical Comput-773 ing. R Foundation for Statistical Computing. Vienna, Austria, 2018. URL: 774 https://www.R-project.org/.
- Alessandro Rinaldo, Stephen E. Fienberg, and Yi Zhou. "On the geometry of discrete exponential families with application to exponential random graph models". In: *Electronic Journal of Statistics* 3 (2009), pp. 446–484.

  ISSN: 1935-7524. DOI: 10.1214/08-EJS350. URL: https://projecteuclid.org/euclid.ejs/1243343761.
- Garry Robins et al. "An introduction to exponential random graph (p\*) models for social networks". In: Social Networks 29.2 (2007), pp. 173–191.

  DOI: 10.1016/j.socnet.2006.08.002.
- 783 [36] Michael Schweinberger. "Instability, Sensitivity, and Degeneracy of Dis784 crete Exponential Families". In: Journal of the American Statistical As785 sociation 106.496 (Dec. 2011), pp. 1361–1370. ISSN: 0162-1459. DOI: 10.
  786 1198/jasa.2011.tm10747. URL: http://www.tandfonline.com/doi/
  787 abs/10.1198/jasa.2011.tm10747.
- Michael Schweinberger and Mark S. Handcock. "Local dependence in random graph models: characterization, properties and statistical inference".

  In: Journal of the Royal Statistical Society: Series B (Statistical Methodology) 77.3 (June 2015), pp. 647–676. ISSN: 13697412. DOI: 10.1111/rssb. 12081. URL: http://doi.wiley.com/10.1111/rssb.12081.
- 793 [38] Michael Schweinberger, Pavel N. Krivitsky, and Carter T. Butts. A note 794 on the role of projectivity in likelihood-based inference for random graph 795 models. 2017. arXiv: 1707.00211 [math.ST].
- Cosma Rohilla Shalizi and Alessandro Rinaldo. "Consistency under sampling of exponential random graph models". In: *Ann. Statist.* 41.2 (Apr. 2013), pp. 508–535. DOI: 10.1214/12-AOS1044. URL: https://doi.org/10.1214/12-AOS1044.
- Andrew J Slaughter and Laura M Koehly. "Multilevel models for social networks: hierarchical Bayesian approaches to exponential random graph modeling". In: *Social Networks* 44 (2016), pp. 334–345. DOI: 10.1016/j. socnet.2015.11.002.
- 804 [41] Tom A. B. Snijders. "Statistical Models for Social Networks". In: *Annual Review of Sociology* 37.1 (2011), pp. 131–153. DOI: 10.1146/annurev. soc.012809.102709.

- 807 [42] Tom A B Snijders et al. "New specifications for exponential random graph models". In: Sociological Methodology 36.1 (Dec. 2006), pp. 99-153. ISSN: 0081-1750. DOI: 10.1111/j.1467-9531.2006.00176.x. URL: http:
  810 //www.jstor.org/stable/25046693%20http://smx.sagepub.com/
  811 lookup/doi/10.1111/j.1467-9531.2006.00176.x.
- Alex D. Stivala et al. "Snowball sampling for estimating exponential random graph models for large networks". In: Social Networks 47 (2016), pp. 167–188. ISSN: 0378-8733. DOI: https://doi.org/10.1016/j.socnet.2015.11.003. URL: http://www.sciencedirect.com/science/article/pii/S0378873315000878.
- Alex Stivala, Garry Robins, and Alessandro Lomi. "Exponential random graph model parameter estimation for very large directed networks". In:

  PLOS ONE 15.1 (Jan. 2020). Ed. by Inés P. Mariño, e0227804. ISSN: 1932-6203. DOI: 10.1371/journal.pone.0227804. URL: https://dx.plos.org/10.1371/journal.pone.0227804.
- W. N. Venables and B. D. Ripley. Modern Applied Statistics with S. Fourth. ISBN 0-387-95457-0. New York: Springer, 2002. URL: http://www.stats.ox.ac.uk/pub/MASS4.
- Peng Wang, Garry Robins, and Philippa Pattison. "PNet: A program for the simulation and estimation of exponential random graph models". In:

  \*\*University of Melbourne\*\* (2006).
- Stanley Wasserman and Philippa Pattison. "Logit models and logistic regressions for social networks: I. An introduction to Markov graphs and p\*".

  In: Psychometrika 61.3 (1996), pp. 401–425. DOI: 10.1007/BF02294547.
- Hadley Wickham. ggplot2: Elegant Graphics for Data Analysis. Springer-Verlag New York, 2016. ISBN: 978-3-319-24277-4. URL: http://ggplot2.
- Jeffrey M Wooldridge. Econometric Analysis of Cross Section and Panel
   Data. 2nd. Cambridge: MIT Press, 2010, p. 1064. ISBN: 978-0-262-23258-6.
- Andy B Yoo, Morris A Jette, and Mark Grondona. "SLURM: Simple Linux Utility for Resource Management". In: Job Scheduling Strategies for Parallel Processing. Ed. by Dror Feitelson, Larry Rudolph, and Uwe Schwiegelshohn. Berlin, Heidelberg: Springer Berlin Heidelberg, 2003, pp. 44–60. ISBN: 978-3-540-39727-4. DOI: 10.1007/10968987\_3. URL: http://link.springer.com/10.1007/10968987%7B%5C\_%7D3.
- Achim Zeileis and Torsten Hothorn. "Diagnostic Checking in Regression Relationships". In: *R News* 2.3 (2002), pp. 7–10. URL: https://CRAN.R-project.org/doc/Rnews/.

#### 45 Appendix A. MLE

The estimation process of ERGM*itos* (as a pooled-data of small networks)
is done entirely on R using the *ergmito* R package. While a significant amount
of the implementation of the methods described here was done using Rcpp [10],
a core component of the package is based on statnet's *ergm* R package, and in
particular, in the function ergm.allstats which does exhaustive enumeration
of statistics in a compact way. In general, the estimation process for any list of
networks is as follows:

- 1. Analyze the model to be estimated: Extract the networks from the left-hand-side as specified in the ergm package, and calculate the exact statistics using the ergm.allstats function.
- 2. With the full enumeration of statistics, build the joint likelihood function of the model in a compact form (i.e., using the weights instead of the full enumeration of the support of the model). This improves speed when it comes to evaluating the log-likelihood function.
- 3. Because we are dealing with exact statistics, it is also possible to calculate the exact gradient function. We compute the gradient as follows:

$$\sum_{p} \nabla l_{p}(\theta) = s \left( y_{p}, X_{p} \right)^{\mathbf{t}} - \frac{Q_{p}^{\mathbf{t}} \left( W_{p}^{\mathbf{t}} \circ \exp \left\{ Q_{p} \theta \right\} \right)}{\kappa_{p}}$$
(A.1)

Where s(y, X) is a vector of observed sufficient statistics (usually called target statistics), Q is a matrix of sufficient statistics, in particular, the isomorphic sufficient statistics associated with the model, and W is a vector of frequency weights.

These first three steps carry the most part of the computing time.

4. Finally, the joint log-likelihood is maximized using the BFGS algorith implemented in the the optim function in the stats package.

The final set of estimates is analyzed separately by another program included in the package. The next section describes the evaluation steps followed by ERGM*ito*.

#### Appendix B. Evaluation of estimates

After the optimization procedure finalizes, the *ergmito* package performs a series of tests checking the quality of the estimates. In particular, we conduct the following evaluations after every call to the main optimization function:

- 1. In the case that the observed sufficient statistics lied on the boundary of its support, the parameter estimate is set to be equal to the corresponding  $\pm \infty$ .
- 2. If all parameters turn out to be ±∞ after this check, the function will send
  a warning message to the user and the function returns without computing
  the variance-covariance matrix. In general, the entries of the Hessian that
  involve a parameter estimate that diverge will be set to zero, which in
  turns results in zero variance-covariance for those entries when computing
  the Moore-Penrose generalized inverse.
  - 3. If, on the other hand, a fraction of the parameters were switched to  $\pm \infty$ , the function recalculates the Hessian and the log-likelihood using the value  $\operatorname{sign}(\hat{\theta}_i) \times 10^5$ . This is done instead of using  $\infty$ , because in most cases using infinite will result in the function being undefined. Again, the function will warn users about this issue. Nevertheless, this is mostly an implementation issue that we are already working on since the limiting values for the log-likelihood, gradient, and hessian are well defined in these cases (see [15]).
    - 4. In the case that the Hessian matrix is non-invertible (not positive-semi-definite [p.s.d.]), we use the Moore-Penrose generalized inverse algorithm as implemented in the R package MASS [45]. For more on the interpretation of variance-covariance matrices when the Hessian is not p.s.d., see [14].

The possible return codes are:

- on optim converged, no issues reported.
  - optim converged, but the Hessian is not p.s.d.

- 901 10 optim did not converged, but the estimates look OK.
- 902 11 optim did not converged, and the Hessian is not p.s.d.
- $\mathbf{20}$  A subset of the parameters estimates was replaced with +/-Inf.
- 21 A subset of the parameters estimates was replaced with +/-Inf, and the
  Hessian matrix is not p.s.d.
- 30 All parameters went to +/-Inf suggesting that the MLE may not exists.

#### Appendix C. Computation details

All the simulations presented in this paper were executed in a large High-Performance-Computing cluster. In general, we use *Slurm* [50] over job arrays with 400 processors. Running the 20,000 simulations took about two hours on the cluster.

In all other cases, i.e. not needing a large computing cluster, model fitting
was done using a laptop computer with Ubuntu 18.04 LTS, 8GB of RAM, a
quad-core processor Intel® Core i5-7200U CPU @ 2.50GHz, and using R version
3.6.3. The number of cores is relevant as the current implementation of the
ergmito R package uses RcppArmadillo R package [11] which can be compiled
using OpenMP [9], meaning that matrix algebra is multi-threaded.