Stochastic blockmodels and community structure in networks

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Stochastic blockmodels have been proposed as a tool for detecting community structure in networks as well as for generating synthetic networks for use as benchmarks. Most blockmodels, however, ignore variation in vertex degree, making them unsuitable for applications to real-world networks, which typically display broad degree distributions that can significantly distort the results. Here we demonstrate how the generalization of blockmodels to incorporate this missing element leads to an improved objective function for community detection in complex networks. We also propose a heuristic algorithm for community detection using this objective function or its non-degree-corrected counterpart and show that the degree-corrected version dramatically outperforms the uncorrected one in both real-world and synthetic networks.

I. INTRODUCTION

A stochastic blockmodel is a generative model for blocks, groups, or communities in networks. Stochastic blockmodels fall in the general class of random graph models and have a long tradition of study in the social sciences and computer science [1–5]. In the simplest stochastic blockmodel (many more complicated variants are possible), each of \( n \) vertices is assigned to one of \( K \) blocks, groups, or communities, and undirected edges are placed independently between vertex pairs with probabilities that are a function only of the group memberships of the vertices. If we denote by \( g_i \) the group to which vertex \( i \) belongs, then we can define a \( K \times K \) matrix \( \psi \) of probabilities such that the matrix element \( \psi_{g_i g_j} \) is the probability of an edge between vertices \( i \) and \( j \).

While simple to describe, this model can produce a wide variety of different network structures. For example, a diagonal probability matrix would produce networks with disconnected components, while the addition of small off-diagonal elements would generate conventional “community structure”—a set of communities with dense internal connections and sparse external ones. Other choices of probability matrix can generate core-periphery, hierarchical, or multipartite structures, among others. This versatility, combined with analytic tractability, has made the blockmodel a popular tool in a number of contexts. For instance, the planted partition model [6], which is equivalent to the model above with a specific parametrization of the matrix \( \psi \), is widely used as a theoretical testbed for graph partitioning and community detection algorithms [7, 8].

Another important application, and the one that is the primary focus of this paper, is the fitting of blockmodels to empirical network data as a way of discovering block structure, an approach referred to in the social networks literature as \textit{a posteriori} blockmodeling [4]. A number of ways of performing the fitting have been suggested, including some that make use of techniques from physics [9, 10]. \textit{A posteriori} blockmodeling can be thought of as a method for community structure detection in networks [8], though blockmodeling is considerably more general than traditional community detection methods, since it can detect many forms of structure in addition to simple communities of dense links. Moreover, it has the desirable property (not shared by most other approaches) of asymptotic consistency under certain conditions [11], meaning that if applied to networks that were themselves generated from the same blockmodel, the method can correctly recover the block structure.

Unfortunately, however, the simple blockmodel described above does not work well in many applications to real-world networks. The model is not flexible enough to generate networks with structure even moderately similar to that found in most empirical network data, meaning that \textit{a posteriori} fits to such data often give poor results [12]. Just as the fitting of a straight line to intrinsically curved data is likely to miss important features of the data, so a fit of the simple stochastic blockmodel to the structure of a complex network is likely to miss much and, as we will show, can in some cases give radically incorrect answers.

Attempts to overcome these problems by extending the blockmodel have focused particularly on the use of (more complicated) \( p^* \) or exponential random graph models, but while these are conceptually appealing, they quickly lose the analytic tractability of the original blockmodel as their complexity increases. Other recent attempts to extend blockmodels take the flavor of mixture models that allow vertices to participate in overlapping groups [13] or to have mixed membership [14, 15].

In this paper we adopt a different approach, considering a simple and apparently minor extension of the classic stochastic blockmodel to include heterogeneity in the degrees of vertices. Despite its innocuous appearance, this extension turns out to have substantial effects, as we will see. A number of previous authors have considered similar extensions of blockmodels. As early as 1987, Wang and Wong [16] proposed a stochastic blockmodel for directed simple graphs incorporating arbitrary expected in- and out-degrees, along with a selection of other features. Unfortunately, this model is not solvable for its parameter values in closed form which limits its usefulness for the types of calculations we consider. Several more recent
works have also explored blockmodels with various forms of degree heterogeneity [17–21], motivated largely by the recent focus on degree distributions in the networks literature. We note particularly the currently unpublished work of Patterson and Bader [21], who apply a variational Bayes approach to a model close, though not identical, to the one considered here.

In this paper we build upon the ideas of these authors but take a somewhat different tack, focusing on the question of why degree heterogeneity in blockmodels is a good idea. To study this question, we develop a degree-corrected blockmodel with closed-form parameter solutions, which allows us more directly to compare traditional and degree-corrected models. As we show, the incorporation of degree heterogeneity in the stochastic blockmodel results in a model that in practice performs much better, giving significantly improved fits to network data, while being only slightly more complex than the simple model described above. Although we here examine only the simplest version of this idea, the approaches we explore could in principle be incorporated into other blockmodels, such as the overlapping or mixed membership models.

In outline, the paper is as follows. We first review the ideas behind the ordinary stochastic blockmodel to understand why degree heterogeneity causes problems. Then we introduce a degree-corrected version of the model and demonstrate its use in a posteriori blockmodeling to infer group memberships in empirical network data, showing that the degree-corrected model outperforms the original model both on actual networks and on new synthetic benchmarks. The benchmarks introduced, which generalize previous benchmarks for community detection, may also be of independent interest.

II. STANDARD STOCHASTIC BLOCKMODEL

In this section we review briefly the use of the original, non-degree-corrected blockmodel, focusing on undirected networks since they are the most commonly studied.

For consistency with the degree-corrected case we will allow our networks to contain both multi-edges and self-edges, even though many real-world networks have no such edges. Like most random graph models for sparse networks the incorporation of multi-edges and self-edges makes computations easier without affecting the fundamental outcome significantly—typically their inclusion gives rise to corrections to the results that are of order $1/n$ and hence vanishing as the size $n$ of the network becomes large. For networks with multi-edges, the previously-defined probability $\psi_{rs}$ of an edge between vertices in groups $r$ and $s$ is replaced by the expected number of such edges, and the actual number of edges between any pair of vertices will be drawn from a Poisson distribution with this mean. In the limit of a large sparse graph, where the probability of an edge and the expected number of edges become equal, there is essentially no difference between the model described here and the standard blockmodel.

With this in mind, the model we study is now defined as follows. Let $G$ be an undirected multigraph on $n$ vertices, possibly including self-edges, and let $A_{ij}$ be an element of the adjacency matrix of the multigraph. Recall that the adjacency matrix for a multigraph is conventionally defined such that $A_{ij}$ is equal to the number of edges between vertices $i$ and $j$ when $i \neq j$, but the diagonal element $A_{ii}$ is equal to twice the number of self-edges from $i$ to itself (and hence is always an even number).

We let the number of edges between each pair of vertices (or between a vertex and itself in the case of self-edges) be independently Poisson distributed and define $\omega_{rs}$ to be the expected value of the adjacency matrix element $A_{ij}$ for vertices $i$ and $j$ lying in groups $r$ and $s$ respectively. Note that this implies that the expected number of self-edges at a vertex in group $r$ is $\frac{1}{2} \omega_{rr}$, because of the factor of two in the definition of the diagonal elements of the adjacency matrix. Now we can write the probability $P(G|\omega, g)$ of graph $G$ given the parameters and group assignments as

$$P(G|\omega, g) = \prod_{i<j} \frac{\omega_{g,ij} A_{ij}}{A_{ij}!} \exp(-\omega_{g,ij}) \times \prod_i \frac{1}{2} \omega_{g,ii} A_{ii}/2! \exp(-\frac{1}{2} \omega_{g,ii}). \quad (1)$$

Given that $A_{ij} = A_{ji}$ and $\omega_{rs} = \omega_{sr}$, Eq. (1) can after a small amount of manipulation be rewritten in the more convenient form

$$P(G|\omega, g) = \frac{1}{\prod_{i<j} A_{ij}! \prod_i 2 A_{ii}/2!} \times \prod_{rs} m_{rs}^{\omega_{rs}/2} \exp(-\frac{1}{2} n_r n_s \omega_{rs}), \quad (2)$$

where $n_r$ is the number of vertices in group $r$ and

$$m_{rs} = \sum_{ij} A_{ij} \delta_{g_i, r} \delta_{g_j, s}, \quad (3)$$

which is the total number of edges between groups $r$ and group $s$, or twice that number if $r = s$.

Our goal is to maximize this probability with respect to the unknown model parameters $\omega_{rs}$ and the group assignments of the vertices. In most cases, it will in fact be simpler to maximize the logarithm of the probability (whose maximum is in the same place). Neglecting constants and terms independent of the parameters and group assignments (i.e., independent of $\omega_{rs}$, $n_r$, and $m_{rs}$), the logarithm is given by

$$\log P(G|\omega, g) = \sum_{rs} (m_{rs} \log \omega_{rs} - n_r n_s \omega_{rs}). \quad (4)$$

We will maximize this expression in two stages, first with respect to the model parameters $\omega_{rs}$, then with
respect to the group assignments $g_i$. The maximum-likelihood values $\hat{\omega}_{rs}$ of the model parameters (where hatted variables indicate maximum-likelihood estimates) are found by simple differentiation to be

$$\hat{\omega}_{rs} = \frac{m_{rs}}{n_r n_s}, \quad (5)$$

and the value of Eq. (4) at this maximum is

$$\log P(G|\hat{\omega}, g) = \sum_{rs} m_{rs} \log \frac{m_{rs}}{n_r n_s} - 2m, \quad (6)$$

where $m = \frac{1}{2} \sum_{rs} m_{rs}$ is the total number of edges in the network. Dropping the constant final, we define the unnormalized log-likelihood for the group assignment $g$:

$$\mathcal{L}(G|g) = \sum_{rs} m_{rs} \log \frac{m_{rs}}{n_r n_s}. \quad (6)$$

The maximum of this quantity with respect to the group assignments now tells us the most likely set of assignments [11, 22]. In effect, Eq. (6) gives us an objective or quality function which is large for “good” group assignments, and small for “poor” ones. Many such objective functions have been defined elsewhere in the literature on community detection and graph partitioning, but Eq. (6) differs from most other choices in being derived from first principles, rather than heuristically motivated or simply proposed ad hoc.

Equation (6) has an interesting information-theoretic interpretation. By adding and dividing by constant factors of the total number of vertices and edges the equation can be written in the alternative form

$$\mathcal{L}(G|g) = \frac{\sum m_{rs} \log \frac{m_{rs}}{2m}}{n_r n_s / n^2}, \quad (7)$$

where again we have neglected irrelevant constants. Now imagine, for a given set of group assignments, that we choose an edge uniformly at random from our network, and let $X$ be the group assignment at one (randomly selected) end of the edge and $Y$ be the group assignment at the other end of the edge. The probability distribution of the variables $X$ and $Y$ is then $P_K(X = r, Y = s) = P_K(r, s) = m_{rs} / 2m$, which appears twice in the above expression. The remaining terms in the denominator of the logarithm in (7) are equal to the expected value of the same probability in a network with the same group assignments but different edges, the edges now being placed completely at random without regard for the groups. Call this second distribution $P_1(r, s)$. Equation (7) can then be written

$$\mathcal{L}(G|g) = \sum_{rs} p_{K}(r, s) \log \frac{p_{K}(r, s)}{p_1(r, s)}, \quad (8)$$

which is the well-known Kullback–Leibler divergence between the probability distributions $p_K$ and $p_1$ [23].

The Kullback–Leibler divergence is not precisely a distance measure since it’s not symmetric in $p_K$ and $p_1$. However, if the logarithms are taken base 2 then it measures the expected number of extra bits required to encode $X$ and $Y$ if $p_1$ is mistakenly used as the distribution for $X$ and $Y$ instead of the assumed true distribution $p_K$. So intuitively it can be considered as measuring how far $p_K$ is from $p_1$. The most likely group assignments under the ordinary stochastic blockmodel are then those assignments that require the most information to describe starting from a model that does not have group structure.

This type of approach, in which one constructs an objective function that measures the difference between an observed quantity and the expected value of the same quantity under an appropriate null model, is common in work on community detection in networks. One widely used objective function is the so-called modularity:

$$Q(g) = \frac{1}{2m} \sum_{ij} [A_{ij} - P_{ij}] \delta(g_i, g_j), \quad (9)$$

where $A_{ij}$ is an element of the adjacency matrix and $P_{ij}$ is the expected value of the same element under some null model. The null model assumed in our blockmodel calculation is one in which $P_{ij}$ is constant. Making the same choice for the modularity would lead to

$$Q(g) = \sum_{r=1}^{K} [p_K(r, r) - p_1(r, r)]. \quad (10)$$

The modularity, however, is not normally used this way and for good reason. This null model, corresponding to a multigraph version of the Erdős–Rényi random graph, produces highly unrealistic networks, even for networks with no community structure. Specifically, it produces networks with Poisson degree distributions, in stark contrast to most real networks, which tend to have broad distributions of vertex degree. To avoid this problem, modularity is usually defined using a different null model that fixes the expected degree sequence to be the same as that of the observed network. Within this model $P_{ij} = k_i k_j / 2m$ where $k_i$ is the degree of vertex $i$. Then the probability distribution over the group assignments at the end of a randomly chosen edge becomes

$$p_{\text{degree}}(X = r, Y = s) = p_{\text{degree}}(r, s) = \frac{K_r}{2m} \frac{K_s}{2m}, \quad (11)$$

where

$$K_r = \sum_s m_{rs} = \sum_i k_i \delta(g_i, r) \quad (12)$$

is the total number of ends of edges, commonly called stubs, that emerge from vertices in group $r$, or equivalently the sum of the degrees of the vertices in group $r$. (Note that Eq. (12) correctly counts two stubs for edges that both start and end in group $r$.) Then the desired group assignments are given by the maximum of

$$Q(g) = \sum_{r=1}^{K} [p_K(r, r) - p_{\text{degree}}(r, r)]. \quad (13)$$

This choice of null model is found to give significantly better results than the original uniform model because
it allows for the fact that vertices with high degree are, all other things being equal, more likely to be connected
than those with low degree, simply because they have more edges. From an information-theoretic viewpoint, an
edge between two high-degree vertices is less surprising
than an edge between two low-degree vertices and we get
better results if we incorporate this observation in our
model.

Returning to the stochastic blockmodel, using \( p_i \) instead of \( p_{\text{degree}} \) in the objective function causes problems
similar to those that affect the modularity. Fits to the
model may incorrectly suggest that structure in the net-
work due merely to the degree sequence is a result in-
stead of group memberships. We will shortly see explicit
real-world cases in which such incorrect conclusions arise.
The solution to this problem, as with the modularity, is
to define a stochastic blockmodel that directly incorpo-
rates arbitrary heterogeneous degree distributions.

III. DEGREE-CORRECTED STOCHASTIC
BLOCKMODEL

In the degree-corrected blockmodel, the probability
distribution over undirected multigraphs with self-edges
(again denoted \( G \)) depends not only on the parameters
introduced previously but also on a new set of parameters \( \theta_i \)
controlling the expected degrees of vertices \( i \).

As before, we assume there are \( K \) groups, \( \omega_{rs} \) is a
\( K \times K \) symmetric matrix of parameters controlling edges
to between groups \( r \) and \( s \), and \( g_i \) is the group assignment
of vertex \( i \). As in the uncorrected blockmodel, let the
numbers of edges each be drawn from a Poisson distri-
bution, but now, following [20] and [21], let the expected
value of the adjacency matrix element \( A_{ij} \) be \( \theta_i \delta_{g_i, g_j} \).

Then graph \( G \) has probability

\[
P(G|\theta, \omega, g) = \frac{1}{\prod_{i\neq j} A_{ij}^{A_{ij}} \exp(-\theta_i \omega_{g_i, g_j})} \times \prod_i \frac{(\frac{1}{2} \theta_i^2 \omega_{g_i, g_i})^{A_{ii}/2}}{(A_{ii}/2)!} \exp(-\frac{1}{2} \theta_i^2 \omega_{g_i, g_i}).
\]

(14)

The \( \theta \) parameters are arbitrary to within a multiplicative
constant which is absorbed into the \( \omega \) parameters. Their
normalization can be fixed by imposing the constraint

\[
\sum_i \theta_i \delta_{g_i, r} = 1
\]

(15)

for all groups \( r \), which makes \( \theta_i \) equal to the probability
that an edge connected to the community to which \( i \) be-
longs lands on \( i \) itself. With this constraint, the proba-
bility \( P(G|\theta, \omega, g) \) can be simplified to the more convenient
form

\[
P(G|\theta, \omega, g) = \frac{1}{\prod_{i\neq j} A_{ij}^{A_{ij}} \prod_i 2^{A_{ii}/2}(A_{ii}/2)!} \times \prod_i \theta_i^{k_i} \prod_i \omega_{rs}^{m_{rs}/2} \exp(-\frac{1}{2} \omega_{rs}),
\]

(16)

with \( k_i \) being the degree of vertex \( i \) as previously and \( m_{rs} \)
defined as in Eq. (3). As before, rather than maximizing
this probability, it is more convenient to maximize its
logarithm, which, ignoring constants, is

\[
\log P(G|\theta, \omega, g) = 2 \sum_i k_i \log \theta_i + \sum_{r,s} (m_{rs} \log \omega_{rs} - \omega_{rs}).
\]

(17)

Allowing for the constraint (15), the maximum-likelihood
values of the parameters \( \theta_i \) and \( \omega_{rs} \) are then given by

\[
\hat{\theta}_i = \frac{k_i}{\kappa_{g_i}}, \quad \omega_{rs} = m_{rs},
\]

(18)

where \( \kappa_r \) is the sum of the degrees in group \( r \) as be-
fore (see Eq. (12)). This maximum-likelihood parameter
estimate has the appealing property of preserving the
expected numbers of edges between groups and the ex-
pected degree sequence of the network. To see this, let \( \langle x \rangle \) be the average of \( x \) in the ensemble of graphs with
parameters (18). Then the expected number of edges
between groups \( r \) and \( s \) is

\[
\sum_{ij} \langle A_{ij} \rangle \delta_{g_i, r} \delta_{g_j, s} = \sum_{ij} k_i k_j \frac{m_{g_i, g_j}}{\kappa_{g_i} \kappa_{g_j}} \delta_{g_i, r} \delta_{g_j, s} = m_{rs},
\]

(19)

where we have made use of Eq. (12). Similarly, the aver-
age degree of vertex \( i \) in the ensemble is

\[
\sum_j \langle A_{ij} \rangle = \sum_j \hat{\theta}_j \hat{\omega}_{g_i, j} = \frac{k_i}{\kappa_{g_i}} \sum_j \frac{k_j}{\kappa_{g_j}} m_{g_i, g_j}
\]

\[= \frac{k_i}{\kappa_{g_i}} \sum_j \frac{k_j}{\kappa_{g_j}} m_{g_i, r} \delta_{g_j, r}
\]

\[= \frac{k_i}{\kappa_{g_i}} \kappa_r m_{g_i, r} = k_i.
\]

(20)

Traditional blockmodels, by contrast, preserve only the
expected value of the matrix \( m_{rs} \) and not the expected
degree—every vertex in group \( r \) in the traditional block-
model has the same expected degree \( \sum_j m_{g_j, g_j}/(n_{r} n_{r}) = \kappa_{r}/n_{r} \).

Substituting Eq. (18) into Eq. (17), the maximum of
\log \( P(G|\theta, \omega, g) \) for the degree-corrected blockmodel is

\[
\log \begin{bmatrix} P(G|\theta, \omega, g) \end{bmatrix} = 2 \sum_i k_i \log \frac{k_i}{\kappa_{g_i}} + \sum_{r,s} m_{rs} \log m_{rs} - 2m.
\]

(21)

where as before \( m \) is the total number of edges in the net-
work. The first term in this expression can be rewritten
as

\[ 2 \sum_{i} k_{i} \log \kappa_{g_{i}} = 2 \sum_{i} k_{i} \log k_{i} - 2 \sum_{i} \sum_{r} k_{i} \delta_{g_{i},r} \log \kappa_{r} \]
\[ = 2 \sum_{i} k_{i} \log k_{i} - \sum_{r} \kappa_{r} \log \kappa_{r} - \sum_{s} \kappa_{s} \log \kappa_{s} \]
\[ = 2 \sum_{i} k_{i} \log k_{i} - \sum_{rs} m_{rs} \log \kappa_{r} \kappa_{s}, \quad (22) \]

where we have again made use of Eq. (12). Substituting back into Eq. (21) and dropping overall constants then gives us an unnormalized log-likelihood function of

\[ \mathcal{L}(G|g) = \sum_{rs} m_{rs} \log \frac{m_{rs}}{\kappa_{r} \kappa_{s}}. \quad \text{(23)} \]

Notice that the only difference between this degree-corrected log-likelihood and the uncorrected log-likelihood of Eq. (6) is the replacement of the number \( n_{r} \) of vertices in each group by the number \( \kappa_{r} \) of stubs. Minor though this replacement may seem, however, it has a big effect, as we will shortly see.

As before, we can interpret the optimization of the objective function (23) through the lens of information theory. Adding and multiplying by constant factors allows us to write the log-likelihood in the form

\[ \mathcal{L}(G|g) = \sum_{rs} \frac{m_{rs}}{2m} \log \frac{m_{rs}/2m}{(\kappa_{r}/2m)(\kappa_{s}/2m)}. \quad \text{(24)} \]

which is the Kullback–Leibler divergence between \( p_{K} \) and \( p_{\text{degree}} \). Alternatively, noting that \( p_{\text{degree}} \) is the product of the marginal distributions \( \sum_{s} p_{K}(r,s) \) and \( \sum_{r} p_{K}(r,s) \), this particular form of divergence can also be thought of as the mutual information of the random variables representing the group labels at either end of a randomly chosen edge. Loosely speaking, the best fit to the degree-corrected stochastic blockmodel gives the group assignment that is most surprising compared to the null model with given expected degree sequence, whereas the ordinary stochastic blockmodel gives the group assignment that is most surprising compared to the Erdős–Rényi random graph.

Information-theoretic quantities have been proposed previously as possible objective functions for community detection or clustering. Dhillon et al. [24], for instance, used mutual information as an objective function for clustering bipartite graphs, as part of an approach they call “information-theoretic co-clustering.” Equation (23) is also somewhat reminiscent of an objective function of Reichardt et al. [18] which, if translated into our terminology and adapted to undirected networks, is equivalent to the total variation distance between \( p_{K} \) and \( p_{\text{degree}} \), variation distance being an alternative measure of the distance between two probability distributions. While the variation distance and the Kullback–Leibler divergence are related, both falling in the class of so-called \( f \)-divergences, the optimization of variation distance does not, to our knowledge, correspond to maximizing the likelihood of any generative model, and there are significant benefits to the connection with generative models. In particular, one can easily create networks from the ensemble of our model and in addition the connection to generative processes means that a posteriori blockmodeling fits into standard frameworks for statistical inference, which are well studied and understood in other contexts.

Equation (23) could also be used as a measure of assortative mixing among discrete vertex characteristics in networks [25, 26]. In a network such as a social network, where connections between individuals can depend on characteristics such as nationality, race, or gender, our objective function could be used, for instance, to quantify which of several such characteristics is more predictive of network structure.

A useful property of the objective function in Eq. (23) when used for a posteriori blockmodeling is that it is possible to quickly compute the change in the log-likelihood when a single vertex switches groups. When a vertex changes groups from \( r \) to \( s \) only \( \kappa_{r}, \kappa_{s}, m_{rs}, \) and \( m_{st} \) (for any \( t \)) can change (with \( m_{rs} \) symmetric). This means that many terms cancel out of the difference of log-likelihoods and can be ignored in the computations.

Consider moving vertex \( i \) from community \( r \) to community \( s \). Let \( k_{it} \) be the number of edges from vertex \( i \) to vertices in group \( t \) excluding self-edges, and let \( u_{i} \) be the number of self-edges for vertex \( i \). These quantities are the same for all possible moves of vertex \( i \). Define \( a(x) = 2x \log x \) and \( b(x) = x \log x \) where \( a(0) = 0 \) and \( b(0) = 0 \). Then the change in the log-likelihood can be written:

\[ \Delta \mathcal{L} = \sum_{t \neq r,s} \left[ a(m_{rt} + k_{it}) - a(m_{rt}) + a(m_{st} + k_{it}) - a(m_{st}) \right] + a(m_{rs} + k_{ir} - k_{is}) - a(m_{rs}) \]
\[ + b(m_{rr} - 2(k_{ir} + u_{i})) - b(m_{rr}) + b(m_{ss} + 2(k_{is} + u_{i})) - b(m_{ss}) - a(\kappa_{r} - k_{i}) + a(\kappa_{r}) - a(\kappa_{s} + k_{i}) + a(\kappa_{s}). \quad \text{(25)} \]

This quantity can be evaluated in time \( O(K + \langle k \rangle) \) on average and finding the \( s \) that gives the maximum \( \Delta \mathcal{L} \) for
given \(i\) and \(r\) can thus be done in time \(O(K(K + \langle k \rangle))\). Because these computations can be done quickly for a reasonable number of communities, local vertex switching algorithms, such as single-vertex Monte Carlo, can be implemented easily. Monte Carlo, however, is slow, and we have found competitive results using a local heuristic algorithm similar in spirit to the Kernighan–Lin algorithm used in minimum-cut graph partitioning [27].

Briefly, in this algorithm we divide the network into some initial set of \(K\) communities at random. Then we repeatedly move a vertex from one group to another, selecting at each step the move that will most increase the objective function—or least decrease it if no increase is possible—subject to the restriction that each vertex may be moved only once. When all vertices have been moved, we inspect the states through which the system passed from start to end of the procedure, select the one with the highest objective score, and use this state as the starting point for a new iteration of the same procedure. When a complete such iteration passes without any increase in the objective function, the algorithm ends. As with many deterministic algorithms, we have found it helpful to run the calculation with several different random initial conditions and take the best result over all runs.

IV. RESULTS

We have tested the performance of the degree-corrected and uncorrected blockmodels in applications both to real-world networks with known community assignments and to a range of synthetic (i.e., computer-generated) networks. We evaluate performance by quantitative comparison of the community assignments found by the algorithms and the known assignments. As a metric for comparison we use the normalized mutual information, which is defined as follows [7]. Let \(n_{rs}\) be the number of vertices in community \(r\) in the inferred group assignment and in community \(s\) in the true assignment. Then define \(p(X = r, Y = s) = n_{rs}/n\) to be the joint probability that a randomly selected vertex is in \(r\) in the inferred assignment and \(s\) in the true assignment. Using this joint probability over the random variables \(X\) and \(Y\), the normalized mutual information is

\[
NMI(X, Y) = \frac{2 \cdot MI(X, Y)}{H(X) + H(Y)},
\]

where \(MI(X, Y)\) is the mutual information and \(H(Z)\) is the entropy of random variable \(Z\). The normalized mutual information measures the similarity of the two community assignments and takes a value of one if the assignments are identical and zero if they are uncorrelated. A discussion of this and other measures can be found in Ref. [28].

A. Empirical networks

We have tested our algorithms on real-world networks ranging in size from tens to tens of thousands of vertices. In networks with highly homogeneous degree distributions we find little difference in performance between the degree-corrected and uncorrected blockmodels, which is expected since for networks with uniform degrees the two models have the same likelihood up to an additive constant. Our primary concern, therefore, is with networks that have heterogeneous degree distributions, and we here give two examples that show the effects of heterogeneity clearly.

The first example, widely studied in the field, is the “karate club” network of Zachary [29]. This is a social network representing friendship patterns between the 34 members of a karate club at a US university. The club in question is known to have split into two different factions as a result of an internal dispute, and the members of each faction are known. It has been demonstrated that the factions can be extracted from a knowledge of the complete network by many community detection methods.

Applying our inference algorithms to this network,
corrected and uncorrected blockmodels with \( K = 2 \), we find the results shown in Fig. 1. As pointed out also by other authors [11, 30], the non-degree-corrected blockmodel fails to split the network into the known factions (indicated by the dashed line in the figure), instead splitting it into a group composed of high-degree vertices and another of low. The degree-corrected model, on the other hand, splits the vertices according to the known communities, except for the misidentification of one vertex on the boundary of the two groups. (The same vertex is also misplaced by a number of other commonly used community detection algorithms.)

The failure of the uncorrected model in this context is precisely because it does not take the degree sequence into account. The \textit{a priori} probability of an edge between two vertices varies as the product of their degrees, a variation that can be fit by the uncorrected blockmodel if we divide the network into high- and low-degree groups. Given that we have only one set of groups to assign, however, we are obliged to choose between this fit and the true community structure. In the present case it turns out that the division into high and low degrees gives the higher likelihood and so it is this division that the algorithm returns. In the degree-corrected blockmodel, by contrast, the variation of edge probability with degree is already included in the functional form of the likelihood, which frees up the block structure for fitting to the true communities.

Moreover it is apparent that this behavior is not limited to the case \( K = 2 \). For \( K = 3 \), the ordinary stochastic blockmodel will, for sufficiently heterogeneous degrees, be biased towards splitting into three groups by degree—high, medium, and low—and similarly for higher values of \( K \). It is of course possible that the true community structure itself corresponds entirely or mainly to groups of high and low degree, but we only want our model to find this structure if it is still statistically surprising once we know about the degree sequence, and this is precisely what the corrected model does.

As a second real-world example we show in Fig. 2 an application to a network of political blogs assembled by Adamic and Glance [31]. This network is composed of blogs (i.e., personal or group web diaries) about US politics and the web links between them, as captured on a single day in 2005. The blogs have known political leanings and were labeled by Adamic and Glance as either liberal or conservative in the data set. We consider the network in undirected form and examine only the largest connected component, which has 1222 vertices. Figure 2 shows that, as with the karate club, the uncorrected stochastic blockmodel splits the vertices into high- and low-degree groups, while the degree-corrected model finds a split more aligned with the political division of the network. While not matching the known labeling exactly, the split generated by the degree-corrected model has a normalized mutual information of 0.72 with the labeling of Adamic and Glance, compared with 0.0001 for the uncorrected model.

(To make sure that these results were not due to a failure of the heuristic optimization scheme, we also checked that the group assignments found by the heuristic have a higher objective score than the known group assignments, and that using the known assignments as the initial condition for the optimization recovers the same group assignments as found with random initial conditions.)

B. Generation of synthetic networks

We turn now to synthetic networks. The networks we use are themselves generated from the degree-corrected
stochastic blockmodel, which is ideally designed to play exactly this role. (Indeed, though it is not the primary focus of this article, we believe that the blockmodel may in general be of use as a source of flexible and challenging benchmark networks for testing the performance of community detection strategies.)

In order to generate networks we must first choose the values of $g$, $\omega$, and $\theta$. The group assignments $g$ can be chosen in any way we please, and we can also choose freely the values for the expected degrees of all vertices, which fixes the $\theta$ variables according to Eq. (18). Choosing the values of $\omega_{rs}$ involves a little more work. In principle, any set of nonnegative values is acceptable provided it is symmetric in $r$ and $s$ and satisfies $\sum_s \omega_{rs} = \kappa_r$, with $\kappa_r$ as in Eq. (12). However, because we wish to be able to vary the level of community structure in our networks we choose $\omega_{rs}$ in the present case to have the particular form

$$\omega_{rs} = \lambda \omega_{rs}^{\text{planted}} + (1 - \lambda) \omega_{rs}^{\text{random}}. \quad (27)$$

This form allows us to interpolate linearly between the values $\omega_{rs}^{\text{planted}}$ and $\omega_{rs}^{\text{random}}$ using the parameter $\lambda$. The $\omega_{rs}^{\text{random}}$ represents a fully random network with no group structure; it is defined to be the expected value of $\omega_{rs}$ in a random graph with fixed expected degrees [32], which is simply

$$\omega_{rs}^{\text{random}} = \frac{\kappa_r \kappa_s}{2m}. \quad (28)$$

The value of $\omega_{rs}^{\text{planted}}$ by contrast is chosen to create group structure. A simple example with four groups is:

$$\omega_{\text{planted}}^{\text{planted}} = \begin{pmatrix} \kappa_1 & 0 & 0 & 0 \\ 0 & \kappa_2 & 0 & 0 \\ 0 & 0 & \kappa_3 & 0 \\ 0 & 0 & 0 & \kappa_4 \end{pmatrix}. \quad (29)$$

With this choice, all edges will be placed within communities when $\lambda = 1$ and none between communities. When $\lambda = 0$, on the other hand, all edges will be placed randomly, conditioned on the degree sequence, and for intermediate values of $\lambda$ we interpolate between these two extremes in a controlled fashion. (This model is similar to the benchmark network ensemble previously proposed by Lancichinetti [33]—roughly speaking it is the “canonical ensemble” version of the “microcanonical” model in [33].)

More complicated choices of $\omega_{rs}^{\text{planted}}$ are also possible. Examples include the core-periphery structure

$$\omega_{\text{planted}}^{\text{planted}} = \begin{pmatrix} \kappa_1 - \kappa_2 & \kappa_2 & 0 \\ \kappa_2 & -\kappa_2 & 0 \\ 0 & 0 & \kappa_3 \end{pmatrix}, \quad (30)$$

where $\kappa_1 \geq \kappa_2$. In the case where $\kappa_1 \approx \kappa_2$ this choice also generates approximately bipartite networks, where most edges run between the two groups and few lie inside. Another possibility is a hierarchical structure of the form

$$\omega_{\text{planted}}^{\text{planted}} = \begin{pmatrix} \kappa_1 - A & A & 0 \\ A & \kappa_2 - A & 0 \\ 0 & 0 & \kappa_3 \end{pmatrix}, \quad (30)$$

where $A \leq \min(\kappa_1, \kappa_2)$.

In mixed models such as these, each edge in effect has a probability $\lambda$ of being chosen from the planted structure and $1 - \lambda$ of being chosen from the null model. Among the edges attached to a given vertex, the expected fraction drawn from the planted structure is $\lambda$ and the remainder are drawn randomly.

Once we have chosen our values for $g$, $\theta$, and $\omega$, the network generation itself is a straightforward implementation of the blockmodel: we first draw a Poisson-distributed number of edges for each pair of groups $r, s$ with mean $\omega_{rs}$ (or $\frac{1}{2} \omega_{rs}$ when $r = s$), then we assign each end of an edge to a vertex in the appropriate group with probability $\theta_l$.

**C. Performance on synthetic networks**

There are two primary considerations in comparing the degree-corrected and uncorrected blockmodels on our synthetic benchmark networks. The first is how close the group assignments found in our calculations are to the planted group assignments. The second is the performance of the heuristic optimization algorithm. It is possible that the maximum-likelihood group assignment may be close to the true group assignment but that our heuristic is unable to find it. And if the heuristic performs better in general for either the corrected or uncorrected blockmodel it may make comparisons between the models unreliable: we want to claim that the degree-corrected model gives better results than the uncorrected version because it has a better objective function for heterogeneous networks and not because we used a biased optimization algorithm.

To shed light on these questions we take the following approach. For both the degree-corrected model and the uncorrected model we perform tests with random initial conditions and with initial conditions equal to the known planted group structure. The latter (planted) initializations tell us whether the planted group assignment, or something close to it, is a local optimum of the respective objective function—if it is, our heuristic should find that optimum most of the time and return a final assignment similar to the planted one. This should be true for essentially any reasonable heuristic, even a biased one, since the heuristic will be making only minimal changes to the group assignments (or none at all).

For small values of $\lambda$ we expect that the planted assignment is not near a local maximum, but for large $\lambda$ we would hope that it is. Thus, if we discover in the process of running our heuristic that it is not, it strongly suggests we have made a poor choice of objective function (and this conclusion should hold even if the heuristic is biased).

The results of such tests on our synthetic networks are shown in Fig. 3. We plot the normalized mutual information as a function of $\lambda$ for various choices of planted structure. Each data point represents an average over 30 networks of size $n = 1000$ for both the degree-corrected
and uncorrected objective functions. In the case of random initializations, ten initializations were performed for each network and we take the best result among the ten.

The left panel in the figure shows results for networks with two communities and just two possible values of the expected degree, 10 and 30. Each of the 1000 vertices was assigned to one of the four possible combinations of degree and community with equal probability, and the planted structure was chosen diagonal, as in Eq. (28).

The green points in the figure indicate the performance of the degree-corrected blockmodel, while the black points are for the uncorrected model. Solid squares and open circles show performance starting from the planted community structure and random assignments respectively. Bearing in mind that \( \lambda = 0 \) corresponds to zero planted structure (in which case neither algorithm should find any significant result) and that a normalized mutual information approaching 1 indicates successful detection of the planted structure, we can see from the figure that the degree-corrected blockmodel significantly out-performs the uncorrected one in this simple test. As \( \lambda \) increases from zero, the mutual information for all algorithms rises, but the corrected model starts to detect some signatures of the planted structure almost immediately and for \( \lambda = \frac{1}{2} \) returns a normalized mutual information above 0.7 for both initial conditions. The uncorrected model, by contrast, finds no planted structure at \( \lambda = \frac{1}{2} \) for either initialization—including when the algorithm is initialized to the known correct answer. The reason for this poor performance is precisely because of the variation in degrees: for values of \( \lambda \) up to around 0.6 the uncorrected model fits these networks better if vertices are assigned to groups according to their degree than if they are assigned according to the planted structure, and hence the best-fit group structure has no correlation with the planted structure.

We have also tested our blockmodels against synthetic networks with two other types of structure, one the core-periphery or approximately bipartite structure of Eq. (29) and the other the hierarchical structure of Eq. (30). In these examples we use a more realistic degree distribution that approximately follows a power law with a minimum expected degree of 10 and an exponent of \(-2.5\). For the core-periphery networks we randomly assign vertices to one of the two groups, while for the hierarchical networks we fix 500 vertices to be in the first of the three groups, put the rest randomly in the other two, and set \( A = \frac{1}{4} \min(\kappa_1, \kappa_2) \). (It has been suggested that choosing non-equal sizes for groups in this way presents a harder challenge for structure detection algorithms [30, 34].)

The performance of our blockmodels on these two classes of networks is shown in the middle (core-periphery) and right (hierarchical) panels of Fig. 3. Again we see that the normalized mutual information increases with increasing \( \lambda \) for all algorithms but that the degree-corrected blockmodel performs significantly better than the uncorrected model. The degree-corrected model with planted assignments consistently does the best among the four options as we would expect, and the degree-corrected model with random initializations performs respectably in all cases, although it’s entirely possible that better performance could be obtained with a better optimization strategy. The performance of the uncorrected model with random initializations, on the other hand, is quite poor [35]. But perhaps the most telling comparison is the one between the degree-corrected model with random initial assignments and the uncorrected model with the planted assignment. This comparison tilts the playing field heavily in favor of the uncorrected model and yet, as Fig. 3 shows, the degree-corrected model still performs about as well as, and in

![Fig. 3: The average normalized mutual information as a function of \( \lambda \) for the three synthetic tests described in the text. Filled squares and transparent circles indicate tests initialized with planted and random assignments respectively. Green points denote results for the degree-corrected blockmodel and black for the ordinary uncorrected model. The left, middle, and right panels show respectively the results for the two-group two-degree networks, core-periphery networks, and hierarchical networks. The error bars indicate the standard error on the mean computed from 30 networks per data point.](image-url)
some cases better than, the uncorrected model.

V. CONCLUSIONS

In this paper, we have studied how one can incorporate heterogeneous vertex degrees into stochastic blockmodels in a simple way, improving the performance of the models for statistical inference of group structure. The resulting degree-corrected blockmodels can also be used as generative models for creating benchmark networks, retaining the generality and tractability of other blockmodels while producing degree sequences closer to those of real networks.

We have found the performance of the degree-corrected model for inference of group structure to be quantitatively better on both synthetic and real-world test networks than the uncorrected model. In networks with substantial degree heterogeneity, the uncorrected model prefers to split networks into groups of high and low degree, and this preference can prevent it from finding the true group memberships. The degree-corrected model correctly ignores divisions based solely on degree and hence is more sensitive to underlying structure.

It seems likely that other more sophisticated blockmodels, such as the recently proposed overlapping and mixed membership models, would benefit from incorporating degree sequences also. In applications to online social network data, for example, where overlapping groups are common, there is frequently substantial degree heterogeneity and hence potentially significant benefits to using a degree-corrected model.

The degree-corrected blockmodel is not without its faults. For instance, the model as described can produce an unrealistic number of zero-degree vertices, and is also unable to model some degree sequences, such as those in which certain values of the degree are entirely forbidden. As a model of real-world networks, it may also fail to accurately represent higher-order network structure such as overrepresented network motifs or degree correlations. From a statistical point of view, it is also somewhat unsatisfactory that the number of parameters in the model scales with the size of the network, which for example prevents fits to a network of one size being used to generate synthetic networks of another size.

But perhaps the chief current drawback of the model is that the number \( K \) of blocks or groups in the network is assumed given. In most structure detection problems the number of groups is not known and a complete calculation will therefore require not only the algorithms described in this paper but also a method for estimating \( K \). Some previously suggested approaches to this problem include cross-validation [14], minimum description length methods using two-part or universal codes [30], maximization of a marginal likelihood [10], and nonparametric Bayesian methods. The marginal likelihood for our degree-corrected blockmodel can be computed explicitly if one assumes conjugate priors on the parameters—Dirichlet for \( \theta \) and gamma for \( \omega \)—but then one must also choose the parameters of those priors, called hyperparameters in the statistical literature. In principle one wants to choose values of the hyperparameters that provide asymptotic consistency—the blockmodel should return the correct number of groups when applied to a network generated from the same blockmodel, at least in certain limits. At present, however, it is not known how to make this choice. An alternative possibility is to note that the blockmodel used here is equivalent to a model that generates an ensemble of matrices with integer entries, implying potential connections to the large statistical literature on contingency table analysis that could be helpful in determining the number of groups in a principled fashion. We leave these questions for future work.

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[12] Technically this is only true if the number \( K \) of groups is moderate. If \( K \) can become arbitrarily large then it is possible to represent any network trivially as a stochastic blockmodel by making each vertex its own group and setting the independent probability of each edge to one.
or zero depending on whether an edge exists between the corresponding vertices in the empirical data.

[35] This algorithm’s performance may not be quite as bad as the figure seems to indicate because for high $\lambda$ the mutual information values are roughly bimodal, with most values being close to either 0 or 1. The mean shown is thus roughly the percentage of times that the uncorrected model successfully finds the planted structure. The same does not apply for the other curves, where the error represents relatively minor variation about the mean.