Random graph models for dynamic networks

Xiao Zhang\textsuperscript{1, a}, Cristopher Moore\textsuperscript{2}, and Mark E.J. Newman\textsuperscript{1, 2}

\textsuperscript{1} Department of Physics, University of Michigan, Ann Arbor, MI 48109, USA
\textsuperscript{2} Santa Fe Institute, 1399 Hyde Park Road, Santa Fe, NM 87501, USA

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Abstract. Recent theoretical work on the modeling of network structure has focused primarily on networks that are static and unchanging, but many real-world networks change their structure over time. There exist natural generalizations to the dynamic case of many static network models, including the classic random graph, the configuration model, and the stochastic block model, where one assumes that the appearance and disappearance of edges are governed by continuous-time Markov processes with rate parameters that can depend on properties of the nodes. Here we give an introduction to this class of models, showing for instance how one can compute their equilibrium properties. We also demonstrate their use in data analysis and statistical inference, giving efficient algorithms for fitting them to observed network data using the method of maximum likelihood. This allows us, for example, to estimate the time constants of network evolution or infer community structure from temporal network data using cues embedded both in the probabilities over time that node pairs are connected by edges and in the characteristic dynamics of edge appearance and disappearance. We illustrate these methods with a selection of applications, both to computer-generated test networks and real-world examples.

1 Introduction

Networked systems, such as social, technological, and biological networks, have been the subject of a vigorous research effort over the last decade \cite{1}, but most of this work has focused on static networks that do not change over time. In reality, almost all networks do in fact change, with nodes or edges appearing or disappearing as the system evolves, and a body of new work aimed at quantifying, modeling, and understanding such temporal or dynamic networks has recently emerged, driven in part by the increasing availability of relevant data \cite{2,3}.

Data on dynamic networks comes in a variety of forms, but the most common form, and the one we consider in this paper, is that of a set of \textit{snapshots} of network structure taken at successive times, usually (though not always) evenly spaced. Such sets are a special case of a more general “multiplex” network, meaning a set of different networks defined on the same set of nodes \cite{4,5}. Multiplex networks include many non-dynamic kinds, such as social networks with different types of interactions between the same set of actors. Our focus in this paper, however, is solely on dynamic networks. We also limit ourselves to networks defined on a fixed and unchanging set of nodes, so that only edges appear and disappear, not nodes.

In many early analyses of dynamic networks, researchers treated snapshots as independent measurements of network structure, analyzing each one separately using conventional static network methods \cite{3}. This, however, ignores the often strong correlations between snapshots, and thereby also ignores a potential rich source of information hidden in the data. In a friendship network, for instance, one expects to still be friends next week with most of the same people one is friends with this week. Moreover, it could be the case that two edges in a network are each present in half the snapshots, but that one of these edges flickers on and off rapidly, while the other varies more slowly. Treating snapshots independently would accurately measure the overall probability that such edges exist, but would be completely insensitive to their rate of appearance and disappearance.

A better approach is to employ methods where the fundamental unit of analysis is not an individual snapshot but the entire history of the network, and a number of researchers have followed this line of reasoning in recent years. Grindrod and coworkers \cite{6,7,8}, for instance, construct models in which the appearance and disappearance of edges in a network obeys a continuous-time Markov process, meaning that edges appear and disappear by making transitions from present to absent or \textit{vice versa} at certain rates. Crucially, these rates can differ from edge to edge, which can induce complex structure in the network. In \cite{6}, for example, the authors considered rates that depend on the “range” of an edge, i.e., its length in some latent space, while \cite{7,8} focus on local properties such as the current degree (the number of neighbors of a node) or transitivity (the number of common neighbors of a node pair).

\textsuperscript{a} e-mail: tomzhang@umich.edu
The models we study in this paper are a special case of models in this class, chosen so that they precisely generalize some of the best known static network models. In particular we do the following. (1) We spell out explicitly dynamic generalizations of the classic random graph, the configuration model, and the widely used stochastic block model, specifically its degree-corrected variant. The models we describe are contained within the larger class defined in [6–8] but are simpler than their more general cousins, making possible certain calculations that would be harder, or even impossible, in the general case. (2) We give calculations of the equilibrium properties of these models and demonstrate explicitly the sense in which they generalize the static models. (3) The bulk of our presentation is given over to the derivation and application of methods for fitting the models to observed network data, which allows us to infer large-scale structure, including (but not limited to) community structure, using maximum-likelihood techniques akin to those developed previously for the static case.

In addition to the work of [6–8], a number of other authors have considered dynamic generalizations of network models, including the random graph [9] and especially the stochastic block model [8,10–17]. The ordinary static version of the stochastic block model divides network nodes into groups or communities and then places edges between them with probabilities that depend on group membership. Dynamic variants of the model have been investigated in which nodes can change their community membership over time, which can cause edge probabilities also to change and hence edges to appear or disappear from one snapshot to the next. Versions of this idea include the dynamic mixed-membership model of Xing et al. [10] and the multi-group membership model studied by Yang et al. [11] and Kim and Leskovec [12]. In Matias and Miele [13] and Ghasemian et al. [14], group memberships can change but edges at successive times are independent conditioned on the groups. Xu [16] has studied a dynamic block model with edge dynamics controlled by a Markov process, which has some elements in common with our approach. Another model similar to ours (but without degree correction) was defined in [8], though in contrast to our approach the authors did not use maximum likelihood to jointly infer the dynamical parameters and the block memberships. Matias et al. [17] have considered “longitudinal” networks where contacts between nodes are governed by a Poisson process. And Perra et al. [18] and Liu et al. [19] have proposed “activity driven” network models, also employing Markov processes, to model heterogeneous activities in dynamic networks, while Ogura and Preciado [20] have considered spreading processes on networks with Markov processes on the edges.

A little further from our focus in this paper are the multilayer stochastic block model studied for instance in references [21,22]. As with dynamic models, these models generate a set of different networks or “layers” built upon the same set of nodes, but there is now no ordering of the layers or any assumption that adjacent layers are more similar than distant ones. Han et al. [21] have used such multilayer models to derive more consistent estimation of community structure for certain data sets than those derived from standard stochastic block models. Stanley et al. [22] studied a variant in which different layers (“strata” in their terminology) are generated from different underlying parameters. Finally, a number of authors have considered heuristic methods for link prediction in networks [23–25], which can be viewed as methods for estimating the future evolution of a network.

In the following section, we describe the models studied in this paper, and derive a range of properties and methods for their application.

2 Dynamic network models

Each of the models we study has a fixed number $n$ of nodes, plus edges between them that appear and disappear as the network evolves over time. Starting from some initial condition at time $t = 0$, our models generate continuous-time network histories, where edges appear and disappear at a sequence of real-valued times. In some data sets, events like these can be observed directly, for instance in a network of telephone calls where we are given the time and duration of each call. Here, however, we assume that the network is only observed at a set of $T$ further snapshots, evenly spaced at integer times $t = 1, \ldots, T$. Including the initial state there are, thus, a total of $T+1$ distinct snapshots. Note that the network is assumed to exist and to continue to evolve unobserved between the snapshots. Note also that since we assume the number of nodes to be fixed our models are not appropriate for networks that are rapidly growing or shrinking over time, such as the world wide web for example.

The fundamental idea behind all of the models we consider is that the connection between every pair of nodes obeys a continuous-time Markov process, edges appearing and disappearing with constant rates, though the rates can differ from one node pair to another depending on various latent properties of the nodes. By choosing this dependence appropriately we can model various kinds of dynamic network structure, including fluctuating densities, degree distributions, and community structure. (The assumption of a Markov process does however exclude networks with bursty or intermittent dynamics, for which our models would not be appropriate).

To make our discussion more concrete, consider a particular pair of nodes in the network. Let us define $\lambda$ to be the rate (in continuous time) at which an edge appears between these two nodes where previously there was none, and let us define $\mu$ to be the rate at which an existing edge disappears. If we denote by $p(t)$ and $p_0(t)$ respectively the probabilities that there is and is not an edge between our nodes at time $t$ then

$$p(t + dt) = p(t) + \lambda p_0(t) \, dt - \mu p_1(t) \, dt,$$  
$$p_0(t + dt) = p_0(t) - \lambda p_0(t) \, dt + \mu p_1(t) \, dt,$$
and hence $p_1$ satisfies the master equation

$$\frac{dp_1}{dt} = -dp_0 = \lambda p_0(t) - \mu p_1(t),$$

which has the solution

$$p_1(t) = \frac{\lambda}{\mu + \lambda} - c e^{-(\mu + \lambda)t},$$

where $c$ is an integration constant and we have made use of $p_0 = 1 - p_1$.

Now suppose that there is no edge between our two nodes at time $t = 0$, i.e., that $p_1(0) = 0$, which corresponds to the choice $c = \lambda/(\mu + \lambda)$. Then the probability of having an edge between our nodes at the next snapshot of the network, at time $t = 1$, is equal to $p_1(1)$, which takes the value

$$\alpha = \frac{\lambda}{\mu + \lambda} [1 - e^{-(\mu + \lambda)}].$$

This is the probability of appearance of an edge between one snapshot and the next. Similarly we can show that the probability of disappearance of an edge is

$$\beta = \frac{\mu}{\mu + \lambda} [1 - e^{-(\mu + \lambda)}],$$

and the equilibrium probability of an edge in the limit of long time is

$$p = \lim_{t \to \infty} p_1(t) = \frac{\lambda}{\mu + \lambda} = \frac{\alpha}{\alpha + \beta}.$$

It will be more convenient to define our models in terms of probabilities such as these, which can always be calculated if necessary from the continuous-time rates $\lambda$ and $\mu$.

### 2.1 Dynamic random graph

The random graph $G(n, p)$, famously studied by Erdős and Rényi in the 1950s and 60s [26,27], is perhaps the most fundamental of all network models. In this model edges are placed between node pairs independently with probability $p$ (or not with probability $1 - p$). In this section we define the first and simplest of our dynamic network models as a direct dynamic counterpart to the random graph.

The definition of the model is straightforward. Starting from some initial state at time $t = 0$, at every snapshot $t$ each node pair not connected by an edge in the previous snapshot gains an (undirected) edge with probability $\alpha$, or not with probability $1 - \alpha$. Similarly each existing edge disappears with probability $\beta$ or not with probability $1 - \beta$.

The net result after $T$ time-steps is a sequence of $T + 1$ snapshots which can be represented by a set of symmetric adjacency matrices $A(t)$ having elements $A_{ij}(t) = 1$ if nodes $i$ and $j$ are connected by an edge in snapshot $t$ and $A_{ij}(t) = 0$ otherwise.

In the limit of long time $T \to \infty$, the average probability of an edge between two nodes in this model is given by equation (7) to be $p = \alpha/(\alpha + \beta)$, the same for every node pair. Hence the stationary distribution of the model is simply the random graph $G(n, p)$. It is in this sense that the model is a dynamic generalization of the random graph.

This is a particularly simple example of the class of models we study – we will look at more complex ones shortly – but even so there are various reasons to be interested in a model of this kind. One could use it for instance to compute the time variation of network properties such as connectivity or component sizes, or the density of specific subgraphs – computations akin to the classic calculations of Erdős and Rényi and others for the static case [26,27]. Our primary interest in this paper, however, is in the use of this and other models as tools for understanding observed network data, using methods of statistical inference: we fit the model to the data by the method of maximum likelihood and the parameters of the fit tell us about our data in much the same way that fitting a straight line through a set of points can tell us about their slope.

Suppose that we have a set of $T + 1$ observed snapshots of some network, measured at uniform intervals over time. If we hypothesize that the data were in fact generated from our dynamic random graph model, then the probability, or likelihood, that we observe this particular set of snapshots, given the parameters $\alpha$, $\beta$ of the model, has the form

$$P(A(t)) = \prod_{i<j} P(A_{ij}(0)|\alpha, \beta) \prod_{t=1}^{T} P(A_{ij}(t)|\alpha, \beta, A_{ij}(t-1)).$$

Note that we have separate terms in this expression for the first snapshot and all succeeding snapshots. The first snapshot differs from the others because it has no preceding snapshots, so its probability is not conditioned on those before it. The probabilities of all later snapshots, on the other hand, depend on the previous state of the network. Because of the assumption that network evolution follows a Markov process, each snapshot only depends directly on the immediately preceding snapshot, hence the inclusion of $A_{ij}(t-1)$ in the second product.

The two probabilities $P(A_{ij}(0)|\alpha, \beta)$ and $P(A_{ij}(t)|\alpha, \beta, A_{ij}(t-1))$ are straightforward to write down. With the assumption that the first snapshot is drawn from the stationary distribution of the Markov process, the first probability, which represents the probability of observing $A_{ij}(0)$ given no information about the previous history of the network, is equal to the stationary probability of an edge or non-edge within the model, which as we have said is $p = \alpha/(\alpha + \beta)$ for an edge, or $1 - p$ for a non-edge. Hence

$$P(A_{ij}(0)|\alpha, \beta) = p A_{ij}(0) (1 - p)^{1 - A_{ij}(0)}.$$

The second probability is only a little more complicated, taking one of four values for edges that appear or not and
ones that disappear or not:
\[
P(A_{ij}(t) | \alpha, \beta, A_{ij}(t-1)) = \alpha^{1-A_{ij}(t-1)} A_{ij}(t) \\
\times (1 - \alpha)^{1-A_{ij}(t-1)} (1 - A_{ij}(t)) \\
\times \beta^{A_{ij}(t-1)} (1 - A_{ij}(t)) (1 - \beta)^{A_{ij}(t-1)} A_{ij}(t),
\]
(10)
Substituting (9) and (10) into equation (8) then gives us the full likelihood for our data. In fact, as is often the case, it is more convenient to work with the logarithm \( \mathcal{L} \) of the likelihood, which has its maximum in the same place. Taking the log of equation (8) and applying (9) and (10), we have
\[
\mathcal{L} = \log P(\{A(t)\} | \alpha, \beta) \\
= \sum_{ij} \{ A_{ij}(0) \log p + [1 - A_{ij}(0)] \log(1 - p) \\
+ \sum_{t=1}^{T} \left[ [1 - A_{ij}(t-1)] A_{ij}(t) \log \alpha \\
+ [1 - A_{ij}(t-1)][1 - A_{ij}(t)] \log(1 - \alpha) + A_{ij}(t-1) \\
\times [1 - A_{ij}(t)] \log \beta + A_{ij}(t-1) A_{ij}(t) \log(1 - \beta) \right] \}.
\]
(11)
Given the likelihood, we can estimate the parameters \( \alpha \) and \( \beta \) by maximizing, which gives
\[
\alpha = \frac{\sum_{ij} [A_{ij}(0) - p + \sum_{t=1}^{T} [1 - A_{ij}(t-1)] A_{ij}(t)]}{\sum_{ij} [A_{ij}(0) - p + \sum_{t=1}^{T} [1 - A_{ij}(t-1)]]},
\]
(12)
\[
\beta = \frac{\sum_{ij} [p - A_{ij}(0) + \sum_{t=1}^{T} A_{ij}(t-1) [1 - A_{ij}(t)]]}{\sum_{ij} [p - A_{ij}(0) + \sum_{t=1}^{T} A_{ij}(t-1)]}.
\]
(13)
Note that these expressions differ from the naive estimates of \( \alpha \) and \( \beta \) given by Laplace’s rule of succession [6], i.e., the number of times an edge appeared or disappeared divided by the number of times it could potentially have done so. The difference arises because the initial state of the network is chosen from the stationary distribution, and the probability \( p \) that \( A_{ij}(0) = 1 \) in this initial state itself depends on \( \alpha \) and \( \beta \). As \( T \to \infty \) the effect of the initial state becomes progressively diluted relative to the effect of the other snapshots and equations (12) and (13) converge to their naive values.

Because \( p \) appears on the right-hand side of (12) and (13), calculating the rates \( \alpha \) and \( \beta \) requires us to find self-consistent solutions to the equations. In fact, it is possible to eliminate the dependence on \( p \) on the right-hand side and derive explicit closed-form equations, but the expressions are somewhat complicated. In practice we have found it simpler just to solve equations (12) and (13) by iteration from a suitable initial condition.

What do these equations tell us? For a given data set, they give us an optimal estimate – better than the naive estimate – of the rate at which edges appear and disappear in our network. This gives us information about the correlation between adjacent snapshots. The combined values of \( \alpha \) and \( \beta \) also give us the maximum-likelihood estimate of the average density of the network, via the average probability \( p = \alpha / (\alpha + \beta) \) of an edge.

This model, however, while illustrative, is not, in practice, very useful. Like the static random graph which inspired it, it is too simple to capture most of the interesting structure in real networks, and in particular it generates networks with Poisson degree distributions, wholly unlike those of real-world networks, which typically have broad and strongly non-Poisson distributions. In the world of static network models, this latter shortcoming is remedied by the configuration model, a more sophisticated random graph that can accommodate arbitrary degree distributions [28, 29]. In the next section, we show how to define a dynamic equivalent of the configuration model.

2.2 Dynamic random graphs with arbitrary expected degrees

The configuration model is a model of a random graph with a given degree sequence [28, 29]. One fixes the degree \( d_i \) of each node \( i = 1, \ldots, n \), then places edges at random subject to the constraints imposed by the degrees. This can be achieved in practice by endowing each node \( i \) with \( d_i \) “half-edges” and choosing a matching of half-edges uniformly at random from the set of all possible matchings. In the limit \( n \to \infty \) the expected number of edges falling between nodes \( i \) and \( j \) in this model is \( d_i d_j / 2m \), where \( m = \frac{1}{2} \sum d_i \) is the total number of edges in the network, and the actual number of edges between each pair of nodes is Poisson distributed with this mean. There is nothing in this model to stop a pair of nodes having two or more edges connecting them – a so-called multiedge – and in general there will be some multiedges in networks generated using the configuration model. Self-loops – edges connecting a node to itself – can and do also appear. Although this is not realistic behavior for most real-world networks, versions of the configuration model that explicitly forbid multiedges and self-loops are much harder to work with than those that do not. Moreover, if the degree distribution has finite mean and variance, the expected number of multiedges and self-loops in the network is constant, independent of \( n \), so they have vanishing density as \( n \to \infty \). For these reasons, one normally puts up with the presence of a few multiedges and self-loops for the sake of simplicity.

A commonly studied variant of the configuration model, which is easier to treat in some ways, involves explicitly placing between each node pair a Poisson-distributed number of edges with mean \( d_i d_j / 2m \). In this variant, sometimes called the Chung–Lu model after two of the first authors to study it [30], the numbers of edges between node pairs are independent random variables, making analysis simpler. The price one pays for this simplicity is that the degrees of individual nodes are no longer fixed, themselves being Poisson-distributed (and asymptotically independent) with mean \( d_i \). Thus \( d_i \) in this case
represents not the actual degree but the expected degree of a node. (The random graph of Erdős and Rényi, with mean degree \( c \), is then the special case of this model where \( d_i = c \) for all \( i \).

In this section we define a dynamic analog of the Chung–Lu model in the sense of this paper: its edges have a dynamics chosen so that their stationary distribution is precisely that of the Chung–Lu model. A dynamic network model where the edge rates depend on the current degrees of their endpoints was proposed in [7]. In our model, by contrast, each node has a given average degree in the limit \( T \to \infty \). Since the Chung–Lu model can contain multiedges, we consider a process for adding and removing edges slightly different from that of the previous section, such that each pair of nodes can have any nonnegative number \( k \) of edges connecting it. Specifically, for each node pair we consider the Poisson process where edges are added at rate \( \lambda \), and each of the existing edges is removed independently at rate \( \mu \). Thus \( k \) is incremented at rate \( \lambda k \) and decremented at rate \( \mu k \).

Let \( p_k(t) \) denote the probability that a node pair has \( k \) edges at time \( t \). Then \( p_k \) satisfies the master equation
\[
\frac{dp_k}{dt} = \lambda p_{k-1}(t) + (k+1)\mu p_{k+1}(t) - (\lambda + k\mu)p_k(t).
\]
(14)

We can solve this equation by defining a generating function \( g(z, t) = \sum_{k=0}^{\infty} p_k(t) z^k \), multiplying both sides of (14) by \( z^k \), and summing over \( k \) to get
\[
\frac{\partial g}{\partial t} = (z - 1) \left[ \lambda g - \mu \frac{\partial g}{\partial z} \right].
\]
(15)
The general solution to this equation is
\[
g(z, t) = e^{(z-1)/(\mu)} f((z-1)e^{-\mu t}),
\]
(16)
where \( f(x) \) is any once-differentiable function of its argument satisfying \( f(0) = 1 \), the latter condition being necessary to fulfill the normalization requirement \( g(1, t) = \sum_k p_k(t) = 1 \) for all \( t \).

In the limit of long time we have \( g(z, t) \to e^{(z-1)/\mu} \), which is the generating function of a Poisson-distributed variable with mean \( \lambda / \mu \). Hence the number of edges between any pair of nodes in this model is Poisson-distributed in the stationary state. If we make the choice
\[
\lambda = \frac{\mu d_i d_j}{2m}
\]
(17)
for some set of values \( d_i \), with \( m = \frac{1}{2} \sum_i d_i \) as previously and any value of \( \mu \), then the mean number of edges between nodes \( i \) and \( j \) is \( \lambda / \mu = d_i d_j / 2m \). In other words, the stationary state of this model is precisely the Chung–Lu model with expected degrees \( d_i \).

This then defines our model: to generate a dynamic network with \( n \) nodes, we specify the expected degree \( d_i \) for each node and the parameter \( \mu \). We generate the initial state of the network from the Chung–Lu model with these expected degrees, and then generate future states by adding edges between each node pair \( i, j \) at rate \( \lambda_{ij} = \mu d_i d_j / 2m \) and removing existing edges at rate \( \mu \). We sample \( T \) snapshots of the resulting network at integer intervals \( t = 1, \ldots, T \) which, along with the initial state at \( t = 0 \), comprise the \( T + 1 \) total snapshots generated by the model. We represent these snapshots by adjacency matrices \( A(t) \).

One could use this model for various purposes, such as making calculations of expected structural properties, but our principal interest here is again in fitting the model to observed network data. As before we achieve this by maximizing a likelihood function, which has the same basic form as previously:
\[
P(\{A(t)\}|\{d_i\}, \mu) = \prod_{i<j} P(A_{ij}(0)|d_i, d_j, \mu) \prod_{t=1}^{T} \prod_{t=1}^{T} P(A_{ij}(t)|d_i, d_j, \mu, A_{ij}(t-1))
\]
(18)
The first probability on the right-hand side is straightforward to write down, since we know that the stationary distribution places a Poisson-distributed number of edges between nodes \( i \) and \( j \) with mean \( d_i d_j / 2m \). Thus
\[
P(A_{ij}(0)|d_i, d_j, \mu) = \frac{(d_i d_j / 2m)^{A_{ij}(0)}}{A_{ij}(0)!} e^{-d_i d_j / 2m},
\]
(19)
which is independent of \( \mu \).

The second probability \( P(A_{ij}(t)|d_i, d_j, \mu, A_{ij}(t-1)) \) is more involved, but the calculation is simplified by noting that even though the model can possess multiedges, the observed network data will normally have at most a single edge between any pair of nodes, so that the only allowed edge transitions are the appearance and disappearance of single edges.

Suppose that a given node pair is connected by zero edges at time \( t = 0 \). Then, setting \( t = 0 \) in equation (16), we find that \( f(x) = e^{-\lambda x / \mu} \), which implies that one timestep later at \( t = 1 \) we have
\[
g(z, 1) = e^{(z-1)(1-e^{-\mu}) / \mu} = e^{(z-1)\beta d_i d_j / 2m},
\]
(20)
where we have made use of equation (17) and for convenience defined the quantity
\[
\beta = 1 - e^{-\mu},
\]
(21)
which (by analogy with our use of the same symbol \( \beta \) in Sect. 2.1) is equal to the total probability that an existing edge disappears during a single unit of time, i.e., between two successive snapshots.

The probabilities \( p_{0|0} \) and \( p_{0|1} \) of a transition from zero edges to, respectively, zero or one edges in a single timestep are then equal to the probabilities \( p_{0}(1) \) and \( p_{1}(1) \) of having zero or one edges at \( t = 1 \). These are given by the zeroth and first coefficients in the expansion of \( g(z, 1) \) in powers of \( z \):
\[
p_{0|0} = e^{-\beta d_i d_j / 2m},
\]
(22)
\[
p_{0|1} = \beta \frac{d_i d_j}{2m} e^{-\beta d_i d_j / 2m},
\]
(23)
By a similar method we also have
\[ p_{1 \to 0} = \beta e^{-\beta d_i/2m}, \]  
\[ p_{1 \to 1} = (1 - \beta) e^{-\beta d_i/2m}, \]  
where we have ignored terms of second and higher order in \(1/m\) in (25). Equations (22)–(25) give the expressions for the probabilities of a transition from zero edges to zero edges, from zero edges to one edge, from one edge to zero edges, and from one edge to one edge respectively. This specifies the full transition matrix for a network that has single edges only and no multiedges. We can now write down the transition probability \(P(A_{ij}(t)|d_i, d_j, \mu, A_{ij}(t-1))\) as a function of \(\beta\):
\[
P(A_{ij}(t)|d_i, d_j, \beta, A_{ij}(t-1)) = (\beta d_i d_j/2m)^{[1-A_{ij}(t-1)]} \times \beta^{A_{ij}(t-1)[1-A_{ij}(t)]} (1 - \beta)^{A_{ij}(t-1)[1-A_{ij}(t)]}. \]  

Substituting this into equation (18) and taking logs, we get the following expression for the log-likelihood in our model:
\[
\mathcal{L} = \sum_{ij} \left( A_{ij}(0) + \sum_{t=1}^{T} [1 - A_{ij}(t-1)] A_{ij}(t) \right) \log \frac{d_i d_j}{2m} + 2(m^{0-1} + m^{1-0}) \log \beta + 2m^{1-1} \log (1 - \beta) - 2m(1+T\beta),
\]  

where we have ignored constant terms and
\[
m^{0-1} = \frac{1}{2} \sum_{i=1}^{T} \sum_{j}[1 - A_{ij}(t-1)] A_{ij}(t)
\]
is the total number of newly appearing edges in the observed data, and similarly
\[
m^{1-0} = \frac{1}{2} \sum_{i=1}^{T} \sum_{j} A_{ij}(t-1)[1 - A_{ij}(t)],
\]
\[
m^{1-1} = \frac{1}{2} \sum_{i=1}^{T} \sum_{j} A_{ij}(t-1) A_{ij}(t).
\]
Maximizing the log-likelihood with respect to the edge disappearance rate \(\beta\), we then find that the optimal value of \(\beta\) is the positive solution of the quadratic equation
\[
mT\beta^2 - (m + m^{0-1} + m^{1-0} + m^{1-1}) \beta + m^{0-1} + m^{1-0} = 0.
\]

Similarly, maximizing with respect to \(d_i\) and bearing in mind that \(m = \frac{1}{2} \sum_i d_i\), we find that \(d_i\) obeys
\[
\frac{2}{d_i} \sum_{j} \left[ A_{ij}(0) + \sum_{t=1}^{T} [1 - A_{ij}(t-1)] A_{ij}(t) \right] = \frac{1}{\sum_{j} d_j} \sum_{j} \left[ A_{ij}(0) + \sum_{t=1}^{T} [1 - A_{ij}(t-1)] A_{ij}(t) \right] - (1 + T\beta) = 0,
\]
which has the solution
\[
d_i = \frac{1}{1 + T\beta} \sum_j \left[ A_{ij}(0) + \sum_{t=1}^{T} [1 - A_{ij}(t-1)] A_{ij}(t) \right].
\]
The sum in this expression is the number of edges initially connected to node \(i\) plus the number that later appear. The divisor \(1 + T\beta\) is the effective number of independent measurements of an edge that we make during our \(T\) snapshots. If \(\beta = 0\), so that edges never appear or disappear, then in effect we only have one measurement of each edge -- the initial snapshot at \(t = 0\). Conversely, if \(\beta = 1\), so that every observed edge immediately disappears on the next snapshot, then all snapshots are independent and the number of independent measurements is \(T + 1\). Thus equation (31) measures the number of observed edges between node pairs divided by the number of independent observations of each node pair.

Equations (29) and (31) give us the maximum-likelihood estimates of the rate parameter \(\beta\) and the expected degrees of the nodes. We note two points. First, these equations have to be solved self-consistently, since the first equation depends on \(d_i\) via \(m = \frac{1}{2} \sum_i d_i\) and the second depends on \(\beta\). Second, neither \(\beta\) nor \(d_i\) are equal to their naive estimates from the data. One might imagine, for instance, that \(d_i\) would be given by the average of \(\sum_j A_{ij}(t)\) over all snapshots, but our results indicate that the maximum-likelihood estimate differs from this value. Both of these effects arise, as in the previous section, because of the information provided by the initial state. Because the initial state is drawn from the stationary distribution, which depends on the model parameters, we can make a better estimate of those parameters by taking it into account than not. On the other hand, the advantage of doing so dwindles as \(T\) becomes large and vanishes in the \(T \to \infty\) limit.

We could use these results, for example, to define in a principled fashion an equivalent of the “degree” for a node in a dynamic network. The actual degree of a node in such a network is a fluctuating quantity, but using our results one can define a single number \(d_i\) for each node that, like the degree in a static network, is a measure of the propensity of that node to connect to others. We give some examples in Section 3.

2.3 Dynamic block models

The stochastic block model is a random graph model of a network that incorporates modules or community structure -- groups of nodes with varying densities of within- and between-group edges. The standard stochastic block model, first proposed by Holland et al. in 1983 [31], is the community-structured equivalent of the random graph of Erdős and Rényi, but like the latter it has shortcomings as a model of real-world networks because the networks it generates have Poisson degree distributions. The degree-corrected stochastic block model [32] is a variant on the same idea that is analogous to the model of Chung.
and Lu [30], allowing us to choose any set of values for the expected degrees of nodes, while also generating a community-structured network.

A number of dynamic versions of block models have been proposed previously, as discussed in the introduction [8,10–17]. In this section we define a dynamic equivalent of the degree-corrected model and show how it can be used to infer community structure from dynamic network data using maximum likelihood. The standard (static) degree-corrected block model divides a network of $n$ nodes into $k$ nonoverlapping groups labeled by integers $1, \ldots, k$. Let us denote by $g_i$ the group to which node $i$ belongs. Then we place a Poisson-distributed number of edges between each node pair $i, j$ with mean equal to $\omega_{g_i}g_{\theta_i}\theta_j$, where $\theta_i$ is a degree-like parameter and $\omega_{g_i}$ is a further set of parameters which control the density of edges within and between each pair of groups. If the diagonal elements $\omega_{g_i}$ are greater than the off-diagonal ones, the model generates networks with conventional “assortative” community structure – dense in-group connections and sparser between-group ones – although other choices of $\omega_{g_i}$ are also possible and are observed in real-world situations.

This description does not completely fix the parameters of the model: they are arbitrary to within a multiplicative constant, since one can multiply all the $\theta_i$ in any group by a constant and divide the same constant out of $\omega_{g_i}$ without affecting the behavior of the model. This is why we refer to $\theta_i$ as a “degree-like parameter” – it plays a role similar to degree in the configuration model but is arbitrary to within a group-dependent multiplicative constant. Following [32], we remove this ambiguity by making a specific choice of normalization, that the sum of $\theta_i$ within any group should be 1:

$$\sum_i \theta_i \delta_{g_i, r} = 1, \quad (32)$$

where $\delta_{ij}$ is the Kronecker delta. This gives us $k$ constraints, one for each of the $k$ groups, and hence fixes all the remaining degrees of freedom.

To generalize this model to the dynamic case we again divide our $n$ nodes into $k$ groups and assign to each of them a degree-like parameter $\theta_i$ satisfying (32). We generate an initial state drawn from the static degree-corrected block model with these parameters. We then generate a history for the network by adding edges between each node pair $i, j$ at rate

$$\lambda_{ij} = \mu_{rs} \omega_{g_i}g_{\theta_i}\theta_j, \quad (33)$$

and removing existing edges independently at rate $\mu_{rs}$, where $r = g_i$ and $s = g_j$ are respectively the groups to which $i$ and $j$ belong. Note the similarity between equations (17) and (33), the primary differences being that the parameter $\mu_{rs}$ now depends on the group memberships and that the factor $1/2m$ has been replaced by the quantity $\omega_{g_i}$, which also depends on the group memberships. By the same argument as before, the number of edges between $i$ and $j$ in the stationary state is Poisson distributed with mean

$$\frac{\lambda_{ij}}{\mu_{rs}} = \omega_{r}g_{\theta_i}\theta_j, \quad (34)$$

which makes the stationary state of this model equivalent to the degree-corrected stochastic block model as desired. Also by the same argument as before, we can calculate the transition rates for edges to appear and disappear between one snapshot and the next, which are

$$p_{0 \rightarrow 0} = e^{-\beta_{r}g_{\omega_i}g_{\theta_i}\theta_j}, \quad (35)$$
$$p_{0 \rightarrow 1} = \beta_{r}g_{\omega_i}g_{\theta_i}\theta_j e^{-\beta_{r}g_{\omega_i}g_{\theta_i}\theta_j}, \quad (36)$$
$$p_{1 \rightarrow 0} = \beta_{r}g_{e^{-\beta_{r}g_{\omega_i}g_{\theta_i}\theta_j}}, \quad (37)$$
$$p_{1 \rightarrow 1} = (1 - \beta_{r})e^{-\beta_{r}g_{\omega_i}g_{\theta_i}\theta_j}, \quad (38)$$

Here

$$\beta_{rs} = 1 - e^{-\mu_{rs}} \quad (39)$$

is the total probability for an existing edge between nodes in groups $r$ and $s$ to disappear in the unit of time between successive snapshots. (Also as before we have in equation (38) discarded terms beyond leading order in the small quantities $\omega_{g_i}$).

By fitting this model to observed network data, we can determine the parameters $\beta_{rs}$, $\omega_{g_i}$, and $\theta_i$ along with the group assignment parameters $g_i$. The likelihood as a function of the four sets of parameters $\{\beta_{rs}\}$, $\{\omega_{g_i}\}$, $\{\theta_i\}$, and $\{g_i\}$ takes the form

$$P(\{A(t)\} | \{\beta_{rs}\}, \{\omega_{g_i}\}, \{\theta_i\}, \{g_i\}) = \prod_{i<j} P(A_{ij}(0) | \beta_{g_i,g_j}, \omega_{g_i,g_j}, \theta_i, \theta_j)$$
$$\times \prod_{t=1}^{T} P(A_{ij}(t) | \beta_{g_i,g_j}, \omega_{g_i,g_j}, \theta_i, \theta_j, A_{ij}(t - 1)). \quad (40)$$

The first probability on the right is straightforward, taking the value

$$P(A_{ij}(0) | \beta_{g_i,g_j}, \omega_{g_i,g_j}, \theta_i, \theta_j) = \frac{(\omega_{g_i,g_j}g_{\theta_i}\theta_j)^{A_{ij}(0)}}{A_{ij}(0)!} e^{-\omega_{g_i,g_j}g_{\theta_i}\theta_j} \quad (41)$$

by definition (which is independent of $\beta_{g_i,g_j}$), while the second can be expressed in terms of the transition probabilities, equations (35)–(38). The resulting expression for the log-likelihood is

See equation (42) next page.

where

$$m_{rs}(0) = \sum_{ij} A_{ij}(0) \delta_{g_i,g_j}\delta_{g_j}, \quad (43)$$

and

$$m_{rs}^{0 \rightarrow 1} = \sum_{ij} [1 - A_{ij}(t - 1)] A_{ij}(t) \delta_{g_i,g_j}\delta_{g_j}, \quad (44)$$
\[
\mathcal{L} = \sum_{ij} \left\{ A_{ij}(0) \log(\omega_{g_i,g_j} \vartheta \theta_j) - \omega_{g_i,g_j} \vartheta \theta_j + \sum_{t=1}^{T} \left[ (1 - A_{ij}(t-1)) A_{ij}(t) \log(\beta_{g_i,g_j} \omega_{g_i,g_j} \vartheta \theta_j) \right. \right.
\]
\[
\left. + A_{ij}(t-1) [1 - A_{ij}(t)] \log(\beta_{g_i,g_j} + A_{ij}(t-1) A_{ij}(t) \log(1 - \beta_{g_i,g_j}) - \beta_{g_i,g_j} \omega_{g_i,g_j} \vartheta \theta_j) \right\}
\]
\[
= \sum_{i,j} \sum_{rs} \delta_{g_i,r} \delta_{g_j,s} \left\{ A_{ij}(0) \log(\omega_{r,s} \vartheta \theta_j) - \omega_{r,s} \vartheta \theta_j + \sum_{t=1}^{T} \left[ (1 - A_{ij}(t-1)) A_{ij}(t) \log(\beta_{r,s} \omega_{r,s} \vartheta \theta_j) \right. \right.
\]
\[
\left. + A_{ij}(t-1) [1 - A_{ij}(t)] \log(\beta_{r,s} + A_{ij}(t-1) A_{ij}(t) \log(1 - \beta_{r,s}) - \beta_{r,s} \omega_{r,s} \vartheta \theta_j) \right\}
\]
\[
= \sum_{i,j} \left\{ A_{ij}(0) + \sum_{t=1}^{T} (1 - A_{ij}(t-1)) A_{ij}(t) \right\} \log(\theta_j) + \sum_{rs} \left\{ m_{rs}(0) \log \omega_{rs} + m_{rs}^{0-1} \log(\beta_{r,s}) + m_{rs}^{1-0} \log(1 - \beta_{r,s}) - (1 + T \beta_{r,s}) \omega_{rs} \right\},
\]

where we made use of equation (32) in the final equality. Hence the degrees are simply proportional to \( \theta_i \), with a constant of proportionality that can be easily calculated once we have the values of \( \omega_{rs} \) from equation (47).

This still leaves us to calculate the maximum-likelihood estimates of the group assignments \( g_i \). To do this, we substitute our estimates of the parameters back into the log-likelihood, equation (42), to get the so-called profile likelihood, which is then maximized over the group assignments \( g_i \). Note that there is no need to calculate the last term \( \sum_{rs} (1 + T \beta_{r,s}) \omega_{rs} \) in the likelihood since, by equation (47), it is equal to \( \sum_{rs} [m_{rs}(0) + m_{rs}^{0-1}] \), which is independent of the group assignments and hence has no effect on the position of the maximum.

Maximization of the profile likelihood over the values of \( g_i \) is harder than maximizing with respect to the other parameters, since the values of the \( g_i \) are discrete. We perform the maximization numerically, using a heuristic algorithm analogous to that used for the static block model in [32], which was in turn inspired by the classic Kernighan–Lin algorithm for graph partitioning [33]. Starting from a random group assignment, we move a single node to a different group, choosing from among all possible such moves the one that most increases (or least decreases) the profile likelihood. We repeat this process, making a chain of successive single-node moves, but with the important qualification that each node is moved only once. When all nodes have been moved once, we reexamine every state passed through during the process to find the one with the highest profile likelihood, then take that state as the starting point for a new repetition of the same algorithm. We continue repeating until no further improvement in the profile likelihood is found. As with many other optimization algorithms, the results can vary from one run to another because of the random initial condition, so one commonly performs several complete runs with different initial conditions, taking as the final answer the output of the run that gives the highest overall value of the profile likelihood.

An alternative way to fit our model would be to use an expectation-maximization (EM) algorithm in which the
model parameters are assigned their maximum-likelihood values but one computes an entire posterior distribution over divisions of the network into groups. The latter distribution, being a large object, is normally evaluated only approximately, either by Monte Carlo sampling or using a belief propagation algorithm [34] in which nodes pass each other estimates of their marginal probabilities of belonging to each group. A belief propagation algorithm was used previously for a different dynamic block model in [14], where each node sends messages both along “spatial” edges to its neighbors in each snapshot and along “temporal” edges to its past and future selves in adjacent snapshots. A similar approach could work in the present case, although our model differs from that of [14] in assuming unchanging group memberships but correlated edges where [14] makes the opposite assumption of time-varying group memberships but independent edges between snapshots.

3 Applications

In this section we give examples of fits of dynamic network data to the dynamic configuration model of Section 2.2 and the dynamic block model of Section 2.3.

3.1 Synthetic networks

Our first set of examples make use of synthetic data sets – computer-generated networks with known structure that we attempt to recover using the maximum-likelihood fit. We demonstrate this approach using the dynamic block model of Section 2.3 and the test networks we use are themselves generated using the same model. Our code for performing the fit, along with an example set of test networks, is available online[1].

We look in particular at the case where the expected degree parameters $d_i$ for all nodes are the same, equal to a constant $c$. For the tests reported here we use $c=16$. At the same time we vary the strength of the community structure, encapsulated in the parameters $\omega_{rs}$, according to

$$\omega_{rs} = \delta \omega_{rs}^{\text{planted}} + (1 - \delta) \omega_{rs}^{\text{random}}$$

(51)

Here $\omega_{rs}^{\text{planted}}$ is diagonal (all elements with $r \neq s$ are zero), $\omega_{rs}^{\text{random}}$ is a flat matrix (all elements are the same), and $\delta \in [0,1]$ is an interpolating parameter. Thus by varying $\delta$ we span the range from a uniform random graph with no community structure ($\delta = 0$) to a network in which all edges lie within communities and none between communities ($\delta = 1$), so that the communities are completely disconnected components.

We similarly vary the rate constants $\beta_{rs}$ according to a second parameter $\eta$, also lying in $[0,1]$, such that

$$\beta_{rs} = \eta \beta_{rs}^{\text{planted}} + (1 - \eta) \beta_{rs}^{\text{uniform}}.$$  

(52)

which interpolates between values that are the same for all groups and the heterogeneous choice $\beta_{rs}^{\text{planted}}$, which can be anything we choose. Note that while varying $\beta_{rs}$ does not change the expected degree or average density of edges in the network, it does change how rapidly edges appear and disappear. Thus $\eta$ controls the extent to which the dynamics of the network, as opposed to merely its average behavior, gives additional information about the community structure.

Once the parameters are fixed, we generate a set of networks, which in our tests have $n=500$ nodes divided into two groups of equal size. For each network we generate an initial state followed by up to five further snapshots. The initial state is generated from the stationary distribution (i.e., from a traditional degree-corrected block model) and the following snapshots are generated according to the prescription of Section 2.3.

We now apply the fitting method of Section 2.3 to these networks to test whether it is able to successfully recover the community structure planted in them. Success, or lack of it, is quantified using the normalized mutual information [35,36], an information-theoretic metric that measures the agreement between two sets of group assignments. A normalized mutual information of 1 indicates exact recovery of the planted groups while 0 indicates complete failure – zero correlation between recovered and planted values.

Figure 1 shows the results of our tests. In panel (a) we fix $\eta = 0$, so that $\beta_{rs}$ is uniform and block structure is indicated only by the relative abundance of edges within and between groups. We use a value of $\beta^{\text{uniform}} = 0.4$, meaning that 40% of extant edges disappear at each timestep. The different curves in the figure show the normalized mutual information as a function of the parameter $\delta$ which measures the strength of the community structure, for different numbers of snapshots from $T=0$ to $T=5$. As we can see, our ability to recover the planted structure diminishes, and eventually fails completely, as the structure becomes weaker, but this effect is partly offset (as we might expect) by increasing the number of snapshots – the more snapshots we use the better we are able to infer the community structure. For larger numbers of snapshots, the algorithm is able to surpass the known “detectability threshold” below which community detection is impossible for single, static networks [34], which is indicated by the vertical dashed line in the plot. In other words the algorithm is able to integrate information about the network over time in order to better determine the shape of the communities.

In Figure 1b we set $\eta = 1$, so that $\beta_{rs} = \beta_{rs}^{\text{planted}}$, choosing the value of $\beta_{rs}^{\text{planted}}$ to be $\beta^{\text{in}} = 0.3$ along the diagonal and $\beta_{out} = 0.5$ off the diagonal, meaning that within-group edges are somewhat more persistent – more likely to be conserved from one snapshot to the next – than between-group edges. This behavior provides another signal of community structure, in addition to the differing time-averaged edge probabilities, which the algorithm can in principle use to determine group memberships. And indeed the results of Figure 1b reflect this, showing that the

[1] Code for fitting the dynamic degree-corrected block model to network data is available online for download at http://www.umich.edu/~mejn/dynamic.zip.
algorithm is able to determine group memberships even well below the detectability threshold, but only when \( T \) is large. If \( T \) is small, then it becomes difficult to determine the values of \( \beta_{in} \) from the data, and hence difficult to determine group membership for small \( \delta \). This point is discussed further below.

In Figure 1c we fix \( \delta = 0 \) and vary \( \eta \) between zero and one using values \( \beta_{uniform} = 0.4 \) as previously, and \( \beta_{in} = 0, \beta_{out} = 0.8 \). With \( \delta = 0 \) there is now no signal whatsoever of community structure present in the positions of the edges. The only clue to the group assignments lies in the rate of appearance and disappearance of edges within and between groups. As we would expect, the algorithm is unable to identify the communities at all when \( T = 0 \) or \( \eta = 0 \), but as \( \eta \) grows for \( T \geq 1 \) the algorithm assigns a larger and larger fraction of nodes to the correct groups, with better performance for larger values of \( T \). These results suggest the existence of a new detectability threshold as a function of \( \eta \), with location tending to zero as \( T \rightarrow \infty \). (A threshold like this was observed, for instance, by Ghasemian et al. [14] in their model, discussed in Sect. 2.3, which has a transition as a function of both the strength of community structure and the relevant rate parameters.)

### 3.2 Real-world examples

We have also tested our models against a number of empirical data sets representing the structure of real-world dynamic networks. We give three examples representing networks drawn from technological and social domains, finding in each case that our dynamic models and their associated algorithms perform better than static methods.

#### 3.2.1 Internet graph

Our first example is a network representation of the structure of the Internet at the level of autonomous systems (ASes), the fundamental units of global packet routing used by the Internet’s Border Gateway Protocol. The structure of the Internet changes constantly and is well documented: a number of ongoing projects collect snapshots of the structure at regular intervals and make them available for research. Here we use data from the CAIDA AS Relationships Dataset\(^2\), focusing on four snapshots of the network’s structure taken at three-month intervals during 2015. The spacing of the snapshots is chosen with an eye to the rate of growth of the network. The Internet has grown steadily in size over the several decades of its existence, and it is still growing today, but this growth is not captured by our models. To ensure better fits, therefore, we first restrict our data to the set of nodes that are present in all of our snapshots, and second choose snapshots that span a relatively short total time. Thus our four snapshots were chosen to be sufficiently far apart in time that the network sees significant change between one snapshot and the next, but close enough that the size of the network does not change greatly.

We fit our Internet data to the dynamic version of the configuration model described in Section 2.2, which

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\(^2\) The CAIDA AS relationships dataset. [http://www.caida.org/data/as-relationships](http://www.caida.org/data/as-relationships).
gives us a way to determine the parameter $\beta$ that controls the rate of appearance and disappearance of edges as well as the effective degrees $d_i$ of nodes $i$ in the network. For the rate parameter we find a maximum likelihood value of $\beta = 0.0896$, which indicates a fairly slow rate of turnover of the edges in the network. Recall that $\beta$ is the average probability that an edge will vanish from one snapshot to the next, so this value of $\beta$ implies that over 90% of edges remain intact between snapshots. As discussed in Section 2.2, one could make a naive estimate of the rate at which edges vanish simply by counting the number that do, but that estimate would be less accurate than the maximum-likelihood one.

Our fit also gives us estimates of the degree parameters $d_i$ from equation (31). Again, we could make naive estimates of the degrees, for instance by assuming snapshots to be independent and averaging the raw degrees of their nodes across snapshots. Figure 2 shows a comparison between the frequency distribution of estimated degrees for the Internet derived from the two methods. As the figure shows, there is in this case relatively little difference between the results from the two methods for most of the degree distribution, although there is some deviation for very high and low degrees. The dynamic method appears to produce a somewhat smoother distribution in these regions, suggesting that departures from the smooth distribution in the naive estimate may be due to transient effects.

The similarity between the maximum likelihood and naive estimates in Figure 2 is not entirely surprising, since the naive average is a correct estimator of the $d_i$ in the limit of a large number of snapshots – it will tend to the correct answer eventually. Even so, it is less than ideal. For instance, while it may give a correct estimate of the degrees, the estimate of the error on the values it gives would still be wrong. By assuming the snapshots to be independent, we effectively assume that we have more measurements of the network than we really do and hence underestimate the variance. For instance, if we observe that the naive degree of a node is unchanging for many snapshots in a row, we may conclude that the average of those values has a very small statistical error, because the fluctuations are small. This, however, would be erroneous if the small fluctuations are actually just a result of the fact that the network is only changing rather slowly.

And error estimates are not the only thing that will be affected by improperly using a naive degree estimate. The values of the degrees themselves can also be affected if the snapshots are strongly correlated, which they are in this case because of the small value of $\beta$. Strongly correlated snapshots will tend to give a node the same or similar degree on successive snapshots, but equation (31) implies that in this case our estimate of $d_i$ should actually decrease over time (as $T$ becomes larger in the denominator while the numerator remains constant). A naive estimate on the other hand would remain unchanged. At first sight the decrease in the maximum-likelihood estimate may appear counterintuitive, but it has a simple physical interpretation: for a node that truly has a constant value of $d_i$, we would expect additional edges to appear occasionally, at a rate dependent on that value. If we do not see any edges appearing, therefore, it implies our initial estimate of the degree was too high and we should revise it downward.

The maximum-likelihood estimator can, on the other hand, also have problems of its own if the model we are fitting is not a perfect description of the data. In the case of the Internet we see two possible sources of disagreement between data and model. First, even though the number of nodes in the network is held fixed, the number of edges is observed to grow over time – the network is becoming more dense. This effect is not included in our model, which assumes constant expected density. Second, we see some evidence that the removal of edges is not uniform as our model assumes, but that edges connected to high-degree nodes disappear at a higher rate than those connected to low-degree ones. Both of these behaviors could potentially affect our results. (It is interesting to ask whether and how the model could be generalized to include them, though we leave pursuit of that question for future work.)

3.2.2 Friendship network

Our second example focuses on a set of social networks from a study by Michell and West of friendship patterns and behaviors among school students in the UK [37]. High-school students at a school in the west of Scotland were polled about their friendship patterns, each student being allowed to name up to twelve friends, and they were also asked about their drinking, smoking, and drug use habits. The entire exercise was conducted a total of three times, at yearly intervals, with the same group of students. The study looked at all students in the school, but the most
detailed data were collected for a subset of 50 girls within the larger population and it is on this subset that we focus here.

The researchers were interested in the extent to which substance use behaviors correlated with friendship patterns. They found that although there was no single factor that would completely explain the friendships of the students, the network of friendships did display homophily according to substance use, meaning that students with similar use patterns were more likely to be friends [38,39].

In our analysis we divide the students into three groups: those who do not drink, smoke, or take drugs on a regular basis; those who exhibit one of these three behaviors; and those who exhibit two or more. We then ask whether it is possible to detect this division into groups based on network structure alone, without any knowledge of student behaviors. We find that when using the dynamic version of the degree-corrected block model described in Section 2.3 it is indeed possible to determine the groups, and to do so with better accuracy than can be achieved by standard static methods. Specifically, we compare results from our dynamic block model to those from the static degree-corrected block model fitted to an aggregate network formed from the union of the three snapshots. The aggregated network is allowed to contain multiedges if two students are connected in more than one snapshot.

Figure 3 shows three pictures of the overall aggregate network of friendships. Each picture is laid out identically, but with different coloring. In panel (a) the three colors represent the ground truth, with green, yellow, and red denoting students who engaged in zero, one, or two or more of the behaviors studied respectively. Panel (b) shows the communities found in the network by fitting to the dynamic block model. Though not perfect, this fit places 64% of the nodes in their correct groups. A random coloring, for comparison, would get only 33% right (since each node would have a $1/3$ probability of assignment to the correct group). Panel (c) shows the results from the standard static algorithm applied to the aggregated network. This fit places only 52% of the nodes in their correct groups.

3.2.3 Proximity network

Our third example is another social network, a network of physical proximity between students in a high school in France [40]. The data were collected using electronic proximity detectors worn by the participants, which recorded the presence and identity of other detectors in their vicinity at intervals of 20 s. The data were collected over five consecutive days, but on the last day only a half day’s worth of data were collected, which we discard, leaving four full days to work with. We construct one snapshot for each day and consider there to be an edge between two participants in a snapshot if three or more contacts between them were recorded during the relevant day. Requiring a minimum number of contacts in this way helps to remove spurious signals from the data, as discussed in [41].

We also restrict our study to those nodes that are present in all snapshots.

Fig. 3. Communities within the friendship network of UK high-school students described in the text. (a) Node colors and shapes indicate ground-truth data on substance use, divided into students who used no substances (green circles), one (yellow squares), or two or more substances (red triangles). (b) Colors and shapes indicate group assignments inferred by fitting the network to the dynamic block model of this paper using all three snapshots. (c) Colors and shapes indicate the group assignments inferred by fitting an aggregate of the three snapshots to the static degree-corrected stochastic block model.

The students in the study were divided among three subject specialties: mathematics/physics, physics/chemistry, and biology. Each specialty was further divided into three classes, so there are a total of nine classes in the data. We attempt to recover these classes from the network data alone, without other information, using both the dynamic model of this paper and a traditional static degree-corrected block model applied to the aggregated network. In this case both methods do well, which is perhaps unsurprising, given that the edges within each group are significantly denser than those between groups. Figure 4 shows the results for the dynamic model in panel (a) and the static model in panel (b). As we can see, both models achieve good classification of the nodes into their classes, although the dynamic model performs slightly better. The error rate – the fraction of incorrectly labeled nodes – is 4.1% for the dynamic model of panel (a) and 5.7% for the static model of panel (b).

The primary benefit of the dynamic model in this case, however, lies not in its ability to recover the communities but in what it reveals about the dynamics of the network. In addition to the communities themselves, the dynamic model also returns values for the rate parameters that can reveal features of the data not seen in the simple static fit.
Fig. 4. Student proximity network. The nine groups of nodes in each panel represent the nine classes and the colors and shapes represent the community structure found using (a) the dynamic model of this paper applied to the four snapshots and (b) the standard static degree-corrected block model applied to the aggregate of the snapshots. Note that classes in the same row belong to the same subject specialty, where PC stands for physics/chemistry, MP stands for math/physics and BIO stands for biology. Classes within the same subject specialty tend to have more inter-class edges than classes in different specialties.

to the aggregate network. Of particular interest in this case are the parameters $\beta_{rs}$, which measure the relative rates at which edges change within and between groups. Our fit gives estimates of

$$
\beta_{rs} \approx \begin{cases} 
0.51 \pm 0.05 & \text{within classes,} \\
0.75 \pm 0.19 & \text{different classes but same specialty,} \\
0.94 \pm 0.19 & \text{different specialties,}
\end{cases}
$$

(53)

where the errors indicate the standard deviation among classes.

In other words, connections are not only more likely between participants in the same class or specialty, but they are also more persistent, in some cases by a wide margin – only about 6% of connections persist from one snapshot to the next between individuals in the different specialties for example, but almost 50% persist within classes.

4 Conclusions

In this paper we have introduced dynamic generalizations of some of the best-known static network models, including the Erdős-Rényi random graph, the configuration model, and the degree-corrected stochastic block model. We have also derived and implemented algorithms for fitting these models to network data that allow us to infer maximum-likelihood estimates of rates of change, node degrees, and community structure. We have tested the performance of our dynamic versions of the configuration model and the block model on synthetic benchmark networks as well as on a selection of data sets representing real-world dynamic networks.

There are a number of directions in which this work could be extended. First, we have focused exclusively on edge dynamics here, but there are also networks in which nodes appear and disappear and it would be a natural generalization to study the dynamics of nodes also, or of both edges and nodes together. We could also allow node properties, such as expected degrees or community memberships, to change over time, as some other authors have done. Second, the assumption of an equilibrium continuous-time Markov processes for the edge dynamics is a particularly simple one, which could be relaxed to encompass more complicated situations, such as those in which the distribution from which snapshots are drawn is non-stationary or out of equilibrium. Third, in our community detection calculations we assume we know the number of communities the network contains, but in many cases we do not have this information. Methods have been developed for determining the number of communities in static networks and it is an interesting question whether those methods can be extended to the dynamic case as well.

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Author contribution statement

X.Z., C.M., and M.E.J.N. designed the research, conducted the research, and prepared the manuscript.
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