A Bayesian Approach to Network Modularity

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We present an efficient, principled, and interpretable technique for inferring module assignments and for identifying the optimal number of modules in a given network. We show how several existing methods for finding modules can be described as variant, special, or limiting cases of our work, and how the method overcomes the resolution limit problem, accurately recovering the true number of modules. Our approach is based on Bayesian methods for model selection which have been used with success for almost a century, implemented using a variational technique developed only in the past decade. We apply the technique to synthetic and real networks and outline how the method naturally allows selection among competing models.

Large-scale networks describing complex interactions among a multitude of objects have found application in a wide array of fields, from biology to social science to information technology [1, 2]. In these applications one often wishes to model networks, suppressing the complexity of the full description while retaining relevant information about the structure of the interactions [3]. One such network model groups nodes into modules, or “communities,” with different densities of intra- and inter-connectivity for nodes in the same or different modules. We present here a computationally efficient Bayesian framework for inferring the number of modules, model parameters, and module assignments for such a model.

The problem of finding modules in networks (or “community detection”) has received much attention in the physics literature, wherein many approaches [4, 5] focus on optimizing an energy-based cost function with fixed parameters over possible assignments of nodes into modules. The particular cost functions vary, but most compare a given node partitioning to an implicit null model, 

\[ H = - \sum_{i>j} (J_L A_{ij} - J_G) \delta_{\sigma_i, \sigma_j} + \sum_{\mu=1}^K h_\mu \sum_{i=1}^N \delta_{\sigma_i, \mu}, \]

a Potts model Hamiltonian with unknown coupling constants \( J_G \equiv \ln(1-\vartheta_d)/(1-\vartheta_c), J_L \equiv \ln \vartheta_c/\vartheta_d + J_G, \) and

\[ p(A, \sigma|\pi, \vartheta, K) = p(A|\sigma, \vartheta) p(\sigma|\pi) \quad \text{(conditional dependence on } K \text{ has been suppressed below for brevity)} \]

where \( c_+ \equiv \sum_{i>j} A_{ij} \delta_{\sigma_i, \sigma_j} \) is the number of edges contained within communities, \( c_- \equiv \sum_{i>j} (1 - A_{ij}) \delta_{\sigma_i, \sigma_j} \) is the number of non-edges contained within communities, \( d_+ \equiv \sum_{i>j} A_{ij} (1 - \delta_{\sigma_i, \sigma_j}) \) is the number of edges between different communities, \( d_- \equiv \sum_{i>j} (1 - A_{ij}) (1 - \delta_{\sigma_i, \sigma_j}) \) is the number of non-edges between different communities, and \( n_\mu \equiv \sum_{i=1}^N \delta_{\sigma_i, \mu} \) is the occupation number of the \( \mu \)-th module. Defining \( \mathcal{H} \equiv -\ln p(A, \sigma|\pi, \vartheta) \) and regrouping terms by local and global counts, we recover (up to additive constants) a generalized version of [10]:

\[ \mathcal{H} = - \sum_{i>j} (J_L A_{ij} - J_G) \delta_{\sigma_i, \sigma_j} + \sum_{\mu=1}^K h_\mu \sum_{i=1}^N \delta_{\sigma_i, \mu}, \]
chemical potentials \( h_{\mu} \equiv -\ln \pi_{\mu} \). (Note that many previous methods omit a chemical potential term, implicitly assuming equally-sized groups.)

While previous approaches \cite{1} \cite{10} minimize related Hamiltonians as a function of \( \vec{\sigma} \), these methods require that the user specifies values for these unknown constants, which gives rise to the resolution limit problem \cite{8} \cite{9}. Our approach, however, uses a disorder-averaged calculation to infer distributions over these parameters, avoiding this issue. To do so, we take beta (\( B \)) and Dirichlet (\( D \)) distributions over \( \vec{\sigma} \) and \( \vec{\pi} \), respectively:

\[
p(\vec{\sigma}) p(\vec{\pi}) \equiv B(\vec{\sigma}_c; \vec{c}_+, \vec{c}_0) B(\vec{\pi}_d; \vec{d}_+, \vec{d}_0) D(\vec{\pi}; \vec{n}_0).
\]  

These conjugate prior distributions, are defined on the full range of \( \vec{\sigma} \) and \( \vec{\pi} \), respectively, and their functional forms are preserved when integrated against the model to obtain updated parameter distributions. Their hyperparameters \( \{\vec{c}_+, \vec{c}_0, \vec{d}_+, \vec{d}_0, \vec{n}_0\} \) act as pseudocounts that augment observed edge counts and occupation numbers.

In this framework the problem of module detection can be stated as: given an adjacency matrix \( \mathbf{A} \), determine the most probable number of modules (i.e. occupied spin states) \( K^* = \text{argmax}_K p(K|\mathbf{A}) \) and infer posterior distributions over the model parameters (i.e. coupling constants and chemical potentials) \( p(\vec{\pi}, \vec{\sigma}|\mathbf{A}) \) and the latent module assignments (i.e. spin states) \( p(\vec{\sigma}|\mathbf{A}) \). In the absence of a priori belief about the number of modules, we demand that \( p(K) \) is sufficiently weak that maximizing \( p(K|A) \propto p(\mathbf{A}|K)p(K) \) is equivalent to maximizing \( p(\mathbf{A}|K) \), referred to as the evidence. This approach to model selection \cite{12} proposed by the statistical physicist Jeffreys in 1935 \cite{13} balances model fidelity and complexity to determine, in this context, the number of modules.

A more physically intuitive interpretation of the evidence is as the disorder-averaged partition function of a spin-glass, calculated by marginalizing over the possible quenched values of the parameters \( \vec{\sigma} \) and \( \vec{\pi} \) as well as the spin configurations \( \vec{\sigma} \):

\[
Z = p(\mathbf{A}|K) = \sum_{\vec{\sigma}} \int d\vec{\sigma} \int d\vec{\pi} p(\mathbf{A}, \vec{\sigma}|\vec{\pi}, \vec{\sigma}) p(\vec{\sigma}) p(\vec{\pi})
\]

\[
= \sum_{\vec{\sigma}} \int d\vec{\sigma} \int d\vec{\pi} e^{-\mathcal{H}} p(\vec{\sigma}) p(\vec{\pi}).
\]  

While the \( \vec{\sigma} \) and \( \vec{\pi} \) integrals in Eqn. 1 can be performed analytically, the remaining sum over module assignments \( \vec{\sigma} \) scales as \( K^N \) and becomes computationally intractable for networks of even modest sizes. To accommodate large-scale networks we use a variational approach that is well-known to the statistical physics community \cite{14} and has recently found application in the statistics and machine learning literature, commonly termed variational Bayes (VB) \cite{15}. We proceed by taking the negative log-

arithm of \( Z \) and using Gibbs’s inequality:

\[
-\ln Z = -\ln \sum_{\vec{\sigma}} \int d\vec{\sigma} \int d\vec{\pi} q(\vec{\sigma}, \vec{\pi}, \vec{\sigma}) \frac{p(\mathbf{A}, \vec{\sigma}, \vec{\pi}, \vec{\sigma})}{q(\vec{\sigma}, \vec{\pi}, \vec{\sigma})} K_0 \leq -\sum_{\vec{\sigma}} \int d\vec{\sigma} \int d\vec{\pi} q(\vec{\sigma}, \vec{\pi}, \vec{\sigma}) \ln \frac{p(\mathbf{A}, \vec{\sigma}, \vec{\pi}, \vec{\sigma})}{q(\vec{\sigma}, \vec{\pi}, \vec{\sigma})} K_0
\]

That is, we first multiply and divide by an arbitrary approximating distribution \( q(\vec{\sigma}, \vec{\pi}, \vec{\sigma}) \) and then upper-bound the log of the expectation by the expectation of the log. We define the quantity to be minimized – the expression in Eqn. 7 – as the variational free energy \( F(q; \mathbf{A}) \), a functional of \( q(\vec{\sigma}, \vec{\pi}, \vec{\sigma}) \). (Note that the negative log of \( q(\vec{\sigma}, \vec{\pi}, \vec{\sigma}) \) plays the role of a test Hamiltonian in variational approaches in statistical mechanics.)

We next choose a factorized approximating distribution \( q(\vec{\sigma}, \vec{\pi}, \vec{\sigma}) = q_{\vec{\sigma}}(\vec{\sigma}) q_{\vec{\pi}}(\vec{\pi}) q_{\vec{\sigma}}(\vec{\sigma}) \) with \( q_{\vec{\sigma}}(\vec{\sigma}) = D(\vec{\sigma}; \vec{n}) \) and \( q_{\vec{\pi}}(\vec{\pi}) = D(\vec{\pi}; \vec{n}) \); as in mean field theory, we factorize \( q_{\vec{\sigma}}(\vec{\sigma}) \) as \( q(\sigma_i = \mu) = Q_{i\mu} \), an \( N \)-by-\( K \) matrix which gives the probability that the \( i \)-th node belongs to the \( \mu \)-th module. Evaluating \( F(q; \mathbf{A}) \) with this functional form for \( q(\vec{\sigma}, \vec{\pi}, \vec{\sigma}) \) gives a function of the variational parameters \( \{\vec{c}_+, \vec{c}_-, \vec{d}_+, \vec{d}_-\} \) and matrix elements \( Q_{i\mu} \) which can subsequently be minimized by taking the appropriate derivatives.

We summarize the resulting iterative algorithm, which provably converges to a local minimum of \( F(q; \mathbf{A}) \) and provides controlled approximations to the evidence \( p(\mathbf{A}|K) \) as well as the posteriors \( p(\vec{\pi}, \vec{\sigma}|\mathbf{A}) \) and \( p(\vec{\sigma}|\mathbf{A}) \).

Initialization.—Initialize the \( N \)-by-\( K \) matrix \( \mathbf{Q} = \mathbf{Q}_0 \) and set pseudocounts \( \vec{c}_+ = \vec{c}_{+0}, \vec{c}_- = \vec{c}_{-0}, \vec{d}_+ = \vec{d}_{+0}, \vec{d}_- = \vec{d}_{-0}, \) and \( \vec{n}_0 = \vec{n}_{0\mu} \).

Main Loop.—Until convergence in \( F(q; \mathbf{A}) \):

(i) Update the expected value of the coupling constants and chemical potentials

\[
\langle J_L \rangle = \psi(\vec{c}_+ - \psi(\vec{c}_-) - \psi(\vec{d}_+ + \psi(\vec{d}_-))
\]

\[
\langle J_G \rangle = \psi(\vec{d}_+ - \psi(\vec{d}_-) + \psi(\vec{c}_+ + \psi(\vec{c}_-))
\]

\[
\langle h_\mu \rangle = \psi \left( \sum_{\mu} \tilde{n}_\mu \right) - \psi(\vec{n}_0),
\]  

where \( \psi(x) \) is the digamma function;

(ii) Update the variational distribution over each spin \( \sigma_i \)

\[
Q_{i\mu} \propto \exp \left\{ \sum_{\mu \neq i} \left[ \langle J_L \rangle A_{ij} - \langle J_G \rangle \right] Q_{j\mu} - \langle h_\mu \rangle \right\}
\]

normalized such that \( \sum_{\mu} Q_{i\mu} = 1 \), for all \( i \);

(iii) Update the variational distribution over parameters from the expected counts and pseudocounts

\[
\tilde{n}_\mu = \langle n_\mu \rangle + \tilde{n}_{\mu 0} = \sum_{i=1}^N Q_{i\mu} + \tilde{n}_{\mu 0}
\]
\begin{equation}
\hat{c}_+ = \langle c_+ \rangle + \hat{c}_{+0} = \frac{1}{2} Tr(Q^T A Q) + \hat{c}_{+0}
\end{equation}
\begin{equation}
\hat{c}_- = \langle c_- \rangle + \hat{c}_{-0} = \frac{1}{2} Tr(Q^T (\hat{u} \langle n \rangle^T - Q)) - \langle c_+ \rangle + \hat{c}_{-0}
\end{equation}
\begin{equation}
\hat{d}_+ = \langle d_+ \rangle + \hat{d}_{+0} = M - \langle c_+ \rangle + \hat{d}_{+0}
\end{equation}
\begin{equation}
\hat{d}_- = \langle d_- \rangle + \hat{d}_{-0} = C - M - (c_-) + \hat{d}_{-0},
\end{equation}

where $C = N(N - 1)/2$, $M = \sum_{i>j} A_{ij}$, and $\hat{u}$ is a $N$-by-$1$ vector of $1$'s.

(iv) Calculate the updated optimized free energy

\begin{equation}
F(q; A) = -\ln \frac{Z_c Z_d Z_{\bar{\varphi}}}{Z_c Z_d Z_{\bar{\varphi}}} + \sum_{\mu=1}^{K} \sum_{i=1}^{N} Q_{i\mu} \ln Q_{i\mu},
\end{equation}

where $Z_{\bar{\varphi}} = B(\hat{u})$ is the beta function with a vector-valued argument, the partition function for the Dirichlet distribution $q_{\varphi}(\hat{\varphi})$ (likewise for $q_{\varphi}(\hat{\varphi})$, $Q_{i\mu}$).

As this provably converges to a local optimum, VB is best implemented with multiple randomly-chosen initializations of $Q_0$ to find the global minimum of $F(q; A)$.

Convergence of the above algorithm provides the approximate posterior distributions $q_{\varphi}(\hat{\varphi})$, $Q_{i\mu}$, and simultaneously returns $K^*$, the number of non-empty modules that maximizes the evidence. As such, one needs only to specify a maximum number of allowed modules and run VB; the probability of occupation for extraneous modules converges to zero as the algorithm runs and the most probable number of occupied modules remains.

This is significantly more accurate than other approximate methods, such as Bayesian Information Criterion (BIC) [16] and Integrated Classification Likelihood (ICL) [17, 18], and is less computationally expensive than empirical methods such as cross-validation (CV) [19, 20] in which one must perform the associated procedure after fitting the model for each considered value of $K$.

Specifically, BIC and ICL are suggested for single-peaked likelihood functions well-approximated by Laplace integration and studied in the large-$N$ limit. For a SBM the first assumption of a single-peaked function is invalidated by the underlying symmetries of the latent variables, i.e. nodes are distinguishable and modules indistinguishable. See Fig. 3 for comparison of our method with the Girvan-Newman modularity [5] in the resolution limit test [3, 9], where VB consistently identifies the correct number of modules. (Note that VB is both accurate and fast: it performs competitively in the “four groups” test [21] and scales as $O(MK)$. Runtime for the main loop in MATLAB on a 2GHz laptop is $\approx 6$ minutes for $N = 10^6$ nodes with average degree 16 and $K = 4$.)

Furthermore, we note that previous methods in which parameter inference is performed by optimizing a likelihood function via Expectation Maximization (EM) [11, 18] are also special cases of the framework presented here. EM is a limiting case of VB in which one collapses the distributions over parameters to point-estimates at the mode of each distribution; however EM is prone to overfitting and cannot be used to determine the appropriate number of modules, as the likelihood of observed data increases with the number of modules in the model. As such, VB performs at least as well as EM while simultaneously providing complexity control [22, 23].

In addition to validating the method on synthetic networks, we apply VB to the 2000 NCAA American football schedule shown in Fig. 2 [24]. Each of the 115 nodes represents an individual team and each of the 613 edges represents a game played between the nodes joined. The algorithm correctly identifies the presence of the 12 conferences which comprise the schedule, where teams tend to play more games within than between conferences, making most modules assortative. Of the 115 teams, 105 teams are assigned to their corresponding conferences, with the majority of exceptions belonging to the frequently-misclassified independent teams [25] – the only disassortative group in the network. We emphasize that, unlike other methods in which the number of conferences must be asserted, VB determines 12 as the most probable number of conferences automatically.

Posing module detection as inference of a latent variable within a probabilistic model has a number of advantages. It clarifies what precisely is to be optimized and suggests a principled and efficient procedure for how to perform this optimization. Inferring distributions over model parameters reveals the natural scale of a given modular network, avoiding resolution limit problems.

FIG. 1: Results for the resolution limit test suggested in [9] and [8]. Shapes and colors correspond to the inferred modules. (Left) Our method, variational Bayes, in which all 15 modules are correctly identified (each clique is assigned a unique color/shape). (Right) GN modularity optimization, where failure due to the resolution limit is observed – neighboring cliques are incorrectly grouped together. (Bottom) The results of this test implemented for a range of true number of modules, $K_{\text{true}}$, the number of 4-node cliques in the ring-like graph. Note that our method correctly infers the number of communities $K_{\text{VB}}$ over the entire range of $K_{\text{true}}$, while GN modularity initially finds the correct number of communities but fails for $K_{\text{true}} \geq 15$ as shown analytically in [9].
This method allows us to view a number of approaches to the problem by physicists, applied mathematicians, social scientists, and computer scientists as related sub-parts of a larger problem. In short, it suggests how a number of seemingly-disparate methods may be re-cast and united. A second advantage of this work is its generalization to other models, including those designed to reveal structural features other than modularity. Finally, use of the evidence allows model selection not only among nested models, e.g. models differing only in the number of parameters, but even among models of different parametric families. The last strikes us as a natural area for progress in the statistical study of real-world networks.

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