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Tolerating the Community Detection Resolution Limit with Edge Weighting

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Communities of vertices within a giant network such as the World-Wide Web are likely to be vastly smaller than the network itself. However, Fortunato and Barthélemy have proved that modularity maximization algorithms for community detection may fail to resolve communities with fewer than $\sqrt{L/2}$ edges, where L is the number of edges in the entire network. This resolution limit leads modularity maximization algorithms to have notoriously poor accuracy on many real networks.

Fortunato and Barthélemy's argument can be extended to networks with weighted edges as well, and we derive this corollary argument. We conclude that weighted modularity algorithms may fail to resolve communities with less than $\sqrt{W\epsilon/2}$ total edge weight, where W is the total edge weight in the network and ϵ is the maximum weight of an inter-community edge. If ϵ is small, then small communities can be resolved.

Given a weighted or unweighted network, we describe how to derive new edge weights in order to achieve a low ϵ , we modify the "CNM" community detection algorithm to maximize weighted modularity, and show that the resulting algorithm has greatly improved accuracy. In experiments with an emerging community standard benchmark, we find that our simple CNM variant is competitive with the most accurate community detection methods yet proposed.

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I. INTRODUCTION

Maximizing the modularity of a network, as defined by Girvan and Newman [1], is perhaps the most popular and cited paradigm for detecting communities in networks. There are many algorithms for approximately maximizing modularity and its variants, such as [2-4]. Community assignments of good modularity feature groups of nodes that are more tightly connected than would be expected. We give the formal definition of modularity below. Recent literature, however, has begun to focus on paradigms other than modularity maximization. This is in part due to Clauset, Newman, and Moore [5], who now advocate a more general notion of "community" than that associated with modularity. The shift away from modularity maximization is also due to Fortunato and Barthèlemy [6], who prove that any community assignment produced by a modularity maximization algorithm will have predictable deficiencies in certain realistic situations. Specifically, they argue that any solution of maximum modularity will suffer from a resolution limit that prevents small communities from being detected in large networks. Furthermore, work by Dunbar [7] indicates that true human communities are generally smaller that 150 nodes. This size is far below the resolution limit inherent in many large networks, such as various social networking sites on the World Wide Web.

We agree with Clauset, Newman, and Moore's [5] idea that it is useful to consider more general definitions for "community"; however, we maintain that it is still important to detect traditional, tightly-connected communities of nodes. In this paper, we revisit the negative result of Fortunato and Barthèlemy and analyze it in a different light. We show that positive results are possible without

contradicting the resolution limit. The key is to apply carefully computed weights to the edges of the network.

With one exception, previous methods for tolerating this resolution limit require searching over an input parameter. For example, Li, et al. [8] address the resolution limit problem by defining a modularity alternative called modularity density. Given a fixed number of communities k, solving a k-means problem will maximize modularity density. Li, et al. generalize modularity density so that tuning a parameter λ favors either small communities (large λ) or large communities (small λ) [8]. Arenas, Fernandez, and Gomez also address the problem of resolution limits [9]. They provide the user with a parameter r that modifies the natural community sizes for modularity maximization algorithms. By tuning r, they influence the natural resolution limit. At certain values of r, small communities will be natural, and at other values of r, large communities will be natural. Our methods apply without specifying any target scale for natural communities, and resolve small and large communities simultaneously.

One solution that resolves communities at multiple scales with no tuning parameter is the HQcut algorithm of Ruan and Zhang [10]. This algorithm alternates between spectral methods and efficient local improvement. It uses a statistical test to determine whether to split each community. Ruan and Zhang argue that a subnetwork with modularity significantly greater than that expected of a random network with the same sequence of vertex degrees is likely to have sub-communities, and therefore should be split. As Fortunato points out in his recent survey [11], though, this stopping criterion is an ad-hoc construction.

Nevertheless, Ruan and Zhang present compelling ev-

idence that the accuracy of HQcut often exceeds that of competitors such as Newman's spectral method followed by Kernighan-Lin local improvement [12] and the simulated annealing method of Guimerà and Amaral [13]. The HQcut solution is not simply the solution of global maximum modularity, so it is not bound by the resolution limit. We obtained the authors' Matlab code for HQcut and we present comparisons with our approach below.

II. RESOLUTION LIMITS

Fortunato and Barthélemy [6] define a *module* to be a set of vertices with positive modularity:

$$\frac{l_s}{L} - \left(\frac{d_s}{2L}\right)^2 > 0,\tag{1}$$

where l_s is the number of undirected edges (links) within the set, d_s is the sum of the degrees of the vertices within the set, and L is the number of undirected links in the entire network. These modules contain more edges than we would expect from a set of vertices with the same degrees, were edges to be assigned randomly (respecting the invariant vertex degrees). iLet us define such modules to be natural communities with respect to modularity maximization. We say that a natural community is minimal if it contains no other natural communities. We wish to resolve the minimal natural communities, and we will discuss this goal in Section VIII B.

In order to ensure that such modules are resolved in a global community assignment with maximum modularity, Fortunato and Barthélemy [6] argue that the following must hold:

$$l_s \ge \sqrt{\frac{L}{2}}. (2)$$

They back up this mathematical argument with empirical evidence. Even in a pathologically easy situation, in which the modules are cliques, and only one edge links any module to a neighboring module, the individual modules will not be resolved in any solution of maximum modularity. Instead, several cliques will be merged into one module. Experiments show that the numbers of links in the resulting modules closely track the $\sqrt{L/2}$ prediction.

Work by Dunbar [7] indicates that true human communities are generally limited to roughly 150 members, and this is corroborated by the recent work of Leskovec, Lang, Dasgupta, and Mahoney [14]. Such communities will have dramatically fewer than $\sqrt{L/2}$ edges in practice. Based on this argument, it would seem that there is little hope for the solutions of modularity maximizing algorithms to be applied in real situations in which $L \gg l_s$. Indeed, partially due to the resolution limit result, the general direction of research in community detection seems to have shifted away from modularity maximization in favor of machine learning techniques.

In this paper, we revisit the resolution limit in the context of edge weighting and derive more positive results.

III. RESOLUTION WITH EDGE WEIGHTS

The definition of a module in equation [1] can easily be generalized when edges have weights. Let w_s be the sum of the weights of all undirected edges connecting vertices within Set s. Let $d^w(v)$, the weighted degree, or strength of vertex v, be the sum of the weights of all edges incident on v. We define $d^w_s = \sum_{v \in s} d^w(v)$ to be the sum of the strengths of the vertices in Set s. Then Set s is a module if and only if:

$$\frac{w_s}{W} - \left(\frac{d_s^w}{2W}\right)^2 > 0. (3)$$

Following [6] step-by-step, when considering a module, we use w_s^{out} to denote the sum of the weights of the edges leaving Set s, and also note that $w_s^{\text{out}} = \alpha_s w_s$, where α_s is a convenience that enables us to rewrite the definition of a module in a useful way. We now have $d_s^w = 2w_s + w_s^{\text{out}} = (\alpha_s + 2)w_s$, and a new, equivalent, definition of a module:

$$\frac{w_s}{W} - \left(\frac{(\alpha_s + 2)w_s}{2W}\right)^2 > 0. \tag{4}$$

Manipulating the inequality, we obtain the relationship:

$$w_s < \frac{4W}{(\alpha_s + 2)^2}. (5)$$

Thus, sets representing communities must not have too much weight in order to be modules.

IV. THE MAXIMUM WEIGHTED MODULARITY

Fortunato and Barthélemy describe the most modular network possible. This yields both computed figures that can be corroborated by experimental evidence, and intuition that the resolution limit in community detection has a natural scale that is related to the total number of links in the network. We will use the same strategy for the weighted case.

First, we imagine a network in which every module is a clique. For a given number of nodes and number of cliques, the modularity will be maximized if each clique has the same size. Weighting does not change the argument of [6] that the modularity approaches 1.0 as the number of cliques goes to infinity. Now, following [6], we consider a slight relaxation of the simple case above: the most modular connected network. This will be our set of m cliques with at least m-1 edges to connect them. Without loss of generality, we consider the case

of m connecting edges — a ring of cliques, as studied by [15].

Departing for a moment from [6], we now consider an edge weighting for the network. With edge weights in the range [0,1], the optimal weighting would assign 1 to each intra-clique edge and 0 to each connecting edge. The weighted modularity of this weighted network would be equivalent to the unweighted modularity of the m independent cliques described above, and would tend to 1.

Relaxing this idealized condition, now assume that we have a weighting function that assigns ϵ to each connecting edge, and 1.0 to each intra-clique edge. We now analyze the resulting weighted modularity.

The total edge weight contained within the cliques is

$$\sum_{s=1}^{m} w_s = W - \epsilon m. \tag{6}$$

Each clique is a module by (3) provided that ϵ is sufficiently small. Summing the contributions of the modules, we find the weighted modularity of the network when broken into these cliques is:

$$Q = \sum_{s} \left[\frac{w_s}{W} - \left(\frac{2w_s + 2\epsilon}{2W} \right)^2 \right]. \tag{7}$$

Since all modules contain the same weight, for all s:

$$w_s = \frac{W - \epsilon m}{m} = \frac{W}{m} - \epsilon \tag{8}$$

The maximum modularity of any solution with m communities is:

$$Q_M(m, W) = m \left[\frac{W/m - \epsilon}{W} - \left(\frac{W/m}{W} \right)^2 \right] = 1 - \frac{\epsilon m}{W} - \frac{1}{m}$$
(9)

To quantify this maximum, we take the derivative with respect to m:

$$\frac{dQ_M}{dm}(m,W) = \frac{-\epsilon}{W} + \frac{1}{m^2} \tag{10}$$

Setting this to zero, we find the number of communities in the optimal solution:

$$m^* = \sqrt{\frac{W}{\epsilon}}. (11)$$

Substituting into (9), we find the maximum possible weighted modularity:

$$Q_M(W) = 1 - \frac{2}{\sqrt{W/\epsilon}}. (12)$$

The unweighted versions of equations [11] and [9] from [6] are, respectively, $m^* = \sqrt{L}$, and $Q_M(L) = 1 - \frac{2}{\sqrt{L}}$. In this unweighted case, the natural scale is clearly related to L. We don't expect to be able to find many more than \sqrt{L} modules in any solution of optimal unweighted modularity.

Our weighted case is similar, but the introduction of ϵ leads to some intriguing possibilities. If ϵ can be made small enough, for example, then there is no longer any limit to the number of modules we might expect in any solution of maximum weighted modularity.

V. THE WEIGHTED RESOLUTION LIMIT

In [6], Fortunato and Barthélemy prove that any module in which $l < \sqrt{L/2}$ may not be resolved by algorithms that maximize modularity. Their argument characterizes the condition under which two true modules linked to each other by any positive number of edges will contribute more to the global modularity as one unit rather than as two separate units. This result is corroborated by experiment. In a large real-world dataset such as the WWW, modules with $l \ll L$ will almost certainly exist.

Following the arguments of [6] directly, while considering edge weights, we now argue that any module s in which

$$w_s < \sqrt{\frac{W\epsilon}{2}} - \epsilon \tag{13}$$

may not be resolved. Consider a scenario in which two small modules are either merged or not. Suppose that the first module has intra-module edges of net weight w_1 , and the second has intra-module edges of net weight w_2 . iWe assume that inter-module edges between these two modules have weight ϵ , explicitly write the expressions for weighted modularity in both cases, and find their difference. The weighted modularity of the solution in which these two modules are resolved exceeds that in which they are merged, provided:

$$w < \frac{2W\epsilon/w}{(\frac{\epsilon}{2t} + \frac{\epsilon}{2t} + 2)(\frac{\epsilon}{w} + \frac{\epsilon}{w} + 2)}$$
 (14)

where w could be either w_1 or w_2 . Manipulation of this expression gives (13).

Two challenges remain: finding a method to set edge weights that achieve a small ϵ , and adapting modularity maximization algorithms to use weights. The second challenge is partially addressed by [16] and [4], but we take a different approach.

VI. EDGE WEIGHTING

There are myriad ways to identify local structure with local computations. Several approaches to community detection, such as [3, 17, 18], are based upon this idea. We use local computations to derive new edge weights. Our approach is to reward an edge for each short cycle connecting its endpoints. These suggest strong interconnections.

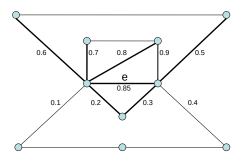


FIG. 1: (Color online) Edge neighborhood weighting

For a vertex v, let E(v) be the set of all undirected edges incident on v. We also define the following sets to express triangle and rectangle relationships between pairs of edges.

 $T_e = \{e' : \text{there exists a 3-cycle containing both } e \text{ and } e'\}$

 $R_e = \{e' : \text{there exists a 4-cycle containing both } e \text{ and } e'\}$

Note that e can be a member of T_e and R_e .

The total weight of edges incident on the endpoints of edge e = (u, v) is

$$W_e = \sum_{e' \in E(u) \cup E(v)} w_{e'}.$$

We consider incident edges that reside on paths of at most three edges connecting the endpoints of e to be "good" with respect to e.

$$G_e = \sum_{e' \in E(u) \cup E(v) \cap (T_e \cup R_e)} w_{e'}.$$

Such edges add credence to the proposition that e is an intra-community edge. We define $neighborhood\ coherence$ of e as follows:

$$C(e) = \frac{G_e}{W_e}$$

For example, in Figure 1, the coherence is computed by summing the weights of the thickened edges and dividing by the total weight of edges incident on the endpoints of e: $C(e) = \frac{4.85}{5.35}$. Alternate definitions are possible, of course, but this weighting is intuitive and performs well in practice.

Arenas, Fernandes, and Gomez, by contrast, add self-loops to vertices according to their r parameter, thereby

"weighting" the nodes, and also adding more intracommunity edges to each module. Thus, they pack more edges into each module in order to satisfy Inequality [2].

We have considered generalizing C(e) to include cycles of length 5 and greater, but this would be a considerable computational expense, and we expect diminishing marginal benefit.

Now we give a simple iterative algorithm for computing edge weights:

- 1. Set $w_e = 1.0$ for each edge e in the network (or accept w_e as input if the edges are already weighted).
- 2. Compute C(e) for each e, set $w_e = C(e)$.
- 3. If any w_e 's changed within some tolerance, go to Step 2

This process will tend to siphon weight out of the intermodule edges (those with smaller C(e)), thus lowering ϵ . We find in practice that it terminates in a small number of iterations. Computing C(e) reduces to finding the triangles and 4-cycles in the graph. This can be done naively in $O(mn \log n)$ time on scale-free graphs. We use Cohen's data structures [19] that admit more efficient algorithms in practice. For WWW-scale graphs, it may be necessary for efficiency reasons to ignore edges incident on high-degree vertices. This would isolate these vertices. However, since such vertices often have special roles in real networks, they might require individual attention anyway.

We define Algorithm W(k) to be k iterations through the loop in Steps 2–3.

VII. WEIGHTED CLAUSET-NEWMAN-MOORE

Any modularity maximization algorithm could be made to leverage edge weights such as those computed in the previous section. Newman replaces individual weighted edges with sets of multiple edges, each with integral weight [16]. We modify the agglomerative algorithm of Clauset, Newman, and Moore (CNM) [2] to handle arbitrary weights directly.

The CNM algorithm efficiently computes the change in modularity (ΔQ) associated with all possible mergers of two existing communities. At the beginning, each vertex is in its own singleton community. Unweighted modularity is defined as follows:

$$Q = \frac{1}{2L} \sum_{vw} \left[A_{vw} - \frac{k_v k_w}{2L} \right] \delta(c_v, c_w)$$
$$= \sum_s (e_{ss} - a_s^2).$$

 A_{vw} is the adjacency matrix entry for directed edge (v, w), k_v is the degree of vertex v, e_{rs} is the fraction of edges that link vertices in community r to vertices in community s, and $a_s = \sum_r e_{rs}$ is the sum of the degrees

of all vertices in community s divided by the total degree. The function $\delta(c_v, c_w)$ equals 1 if v and w are in the same community, and 0 otherwise.

Since vertices i and j initially reside in their own singleton communities, e_{ij} is initially simply $\frac{A_{ij}}{2L}$. The first step in CNM is to initialize ΔQ for all possible mergers:

$$\Delta Q = \begin{cases} 1/(L) - 2k_i k_j/(2L)^2 & \text{if } i,j \text{ are connected} \\ 0 & \text{otherwise.} \end{cases}$$
 (15)

Algorithm	ϵ	m^*	S	Q_M	Q
CNM	N.A.	108	108	0.980	0.980
wCNM1	0.111	286	263	0.9930	0.9928
wCNM5	< 0.000001	1000	1000	0.9999	0.9986

TABLE I: These results from the ring of 1000 5-cliques illustrate gains made by considering weighting. Predicted (m^*) and algorithmically discovered (|S|) numbers of communities match well and indicate that careful weighting makes it possible to resolve all 1000 cliques as modules in a solution of maximal weighted modularity. Q_M is defined in (12), m^* is defined in (11), and ϵ is the weight of the heaviest edge between two communities.

CNM also initializes $a_i = \frac{k_i}{2L}$ for each vertex i. Once the initializations are complete, the algorithm repeatedly selects the best merger, then updates the ΔQ and a_i values, until only one community remains. The solution is the community assignment with the largest value of Q encountered during this process. Clever data structures allow efficient update of the ΔQ values.

To modify CNM to work on weighted graphs, we need only change the initialization step. The update steps are identical. We simply define and compute the strength of each vertex $k_i^w = \sum_j w_{ij}$. The initialization becomes:

$$\Delta Q^w = \begin{cases} w_{ij}/(W) - 2k_i^w k_j^w/(2W)^2 & \text{if } i,j \text{ are connected} \\ 0 & \text{otherwise,} \end{cases}$$
(16)

and $a_i^w = \frac{k_i^w}{2W}$. With these initializations, normal CNM merging greedily maximizes weighted modularity Q^w . We refer to this algorithm as wCNM. Note that our definition of Q^w is equivalent to that of [4].

VIII. RESULTS

Given an undirected, weighted or unweighted network, we apply the Algorithm W(k) to set our edge weights, then run wCNM. We use wCNMk to denote this two-step process. Note that running wCNM0 is equivalent to running CNM.

We will consider two different datasets: the ring of cliques example discussed above, and the benchmark of [20], which is a generalization of the 128-node benchmark of Girvan and Newman [21].

A. The ring of cliques

Refer to Table I for the following discussion. Danon, Díaz-Guilera, Duch, and Arenas [15] considered m disconnected cliques as a pathological example of maximum modularity (which approaches 1.0 as the number of cliques increases). Fortunato and Barthèlemy [6] add single connections between cliques to form a ring. Our intuition is that the natural communities in such a graph are the cliques. However, the resolution limit argument of Fortunato and Barthèlemy indicates that this will not be the solution of maximum modularity if each clique has fewer than $\frac{\sqrt{L}}{2}$ edges. They confirm this via experiment, and we have reproduced their results for an instance with 1000 cliques of size five. Table I summarizes the performance of CNM and wCNM for this case. The m^* column contains the number of communities expected in a solution of maximum weighted modularity, as defined in 11. The first row shows the unweighted case, in which m^* is equivalent to that defined in [6]. CNM achieves this theoretical maximum by finding 108 communities, which is much smaller than the number of cliques.

If we run wCNM1, which performs one iteration of neighborhood coherence, we obtain the results in Row 2 of Table I. The value of ϵ we observe is 0.047, leading via (11) to an estimate of 286 resolved communities. The wCNM1 algorithm resolves 263. In a run with five iterations, labeled wCNM5, we both expect and find 1000 communities, resolving all of the natural communities and simultaneously observing our highest weighted modularity. Iterating further reduces ϵ without changing the community assignment.

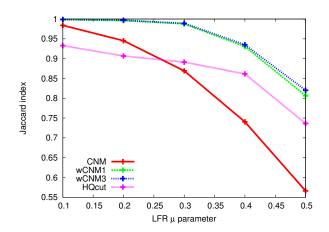
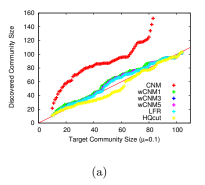
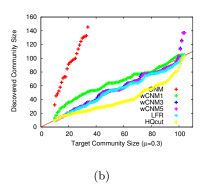


FIG. 2: (Color) Normalized Mutual Information study for the LFR benchmark.





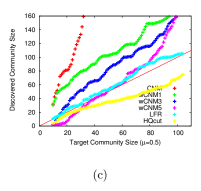


FIG. 3: (Color) Example distributions of community sizes are shown in these quantile-quantile plots. The line y = x represents a perfect match between discovered community sizes and the LFR power-law distribution.

B. The LFR Benchmark

Lancichinetti, Fortunato, and Radicchi [20] (LFR) give a generalization of the popular Girvan and Newman benchmark [21] (GN) for evaluating community detection algorithms. The latter consists of 128-vertex random graphs, each with 4 natural communities of size 32. The proportion of intra-community edges is parameterized. Many authors use GN to create plots of "mutual information," or agreement in node classification between algorithm-discovered communities and natural communities. The LFR benchmark is similar in spirit, but considerably more realistic. It allows the user to specify distributions both for the community sizes and the vertex degrees. Users also specify the average ratio (per vertex) of inter-community adjacencies to total adjacencies, called mixing parameter μ . At $\mu = 0.0$, all edges are intra-community.

The LFR benchmark construction process begins by sampling vertex degrees and creating a graph with the selected degree distribution. It then samples community sizes. A vertex of degree k should have about $(1-\mu)k$ neighbors from the same community. Therefore, it is assigned to a community with at least $(1-\mu)k+1$ vertices. LFR assigns vertices to communities via an interated random process enforcing this constraint, then rewires until the average μ meets the desired value. We have a special interest in the LFR benchmark because it generates graphs with both small and large natural communities.

For several different values of μ , we used the C code from Fortunato's web site (cited in [20]) to generate 30 instances each of LFR benchmark graphs, each with 5000 vertices and average degree 8. The community sizes were selected from the power-law distribution $f(k) \sim k^{-1.5}$, with $k \in [10, 105]$. The degree distribution was $f(k) \sim k^{-2}$, with $k \in [2, 50]$. We specified an average degree of 8, which is roughly comparable to that of the WWW.

Figure 2 contains the mutual information plot for our experiments with LFR. Our metric for comparison is the

Jaccard index [22]:

$$J(A,B) = \frac{|A \cap B|}{|A \cup B|}$$

where A is the set of intra-community edges in the LFR ground truth, and B is the set of intra-community edges in an algorithm solution. As predicted by the resolution limit argument, CNM, an unweighted modularity maximization algorithm, is not able to resolve most of the natural communities. However, even with these more realistic data, wCNM achieves greater accuracy than the sophisticated HQcut algorithm. This is notable, considering the reputation for poor accuracy recently associated with agglomerative algorithms such as CNM and its variants [23]. The accuracy of our CNM variant, on the other hand, is competitive.

Iterating the neighborhood coherence weighting provides diminishing marginal returns. However, as we show below, such iteration does add value.

In addition to the mutual information, we wish to compare the distributions of the sizes of communities discovered by CNM and its weighted variants to the original distributions used in LFR generation. It is a challenge to fit empirical data to heavy-tailed power-law distributions. However, the discrete power-law distribution of community sizes used by LFR is not heavy-tailed.

LFR uses the following precise sampling process to determine ground truth community sizes:

- 1. Compute $\frac{1}{k}^{\tau}$, the probability that a community will have size k.
- 2. For all $a \le k \le b$, where a and b bound the community sizes, compute the empirical cumulative distribution function for k: $p_k = \sum_{k'=a}^k (\frac{1}{k'})^{\tau}$.
- 3. For a uniform random variate $x \in [0,1]$, find the minimum k' such that $p_{k'} \ge x$.

This process continues until the sum of the community sizes exceeds the number of vertices, and the final community is truncated.

We approach the problem of testing goodness-of-fit of sets of algorithm-generated community sizes by generating visualizations and performing hypothesis tests. In both cases, we compare the empirical distributions of community sizes with the untruncated discrete power-law distribution that underlies the LFR distribution.

For visualization, we generate quantile-quantile plots using the R language [24] and its quantile() function with interpolation type 8. This is the recommendation of Hyndman and Fan [25]. Figure 3 shows three such plots: one LFR instance each of μ values 0.1, 0.3, and 0.5. With the moderate community coherence of $\mu=0.3$, the wCNM variants track the target distribution closely, show a drastic improvement over CNM, and appear to dominate HQcut. This latter claim is corroborated by the hypothesis tests described below. At $\mu=0.5$, the advantage over CNM is still clear, but neither wCNM nor HQcut track the target distribution closely.

To augment our results with statistical evidence, we use the classical Kolmogorov-Smirnov (K–S) test as described, for example, in [26]. Our null hypothesis is that the algorithm-generated community sizes follow a discrete power-law with $\tau=1.5$. We computed critical values for each sample size between 10 and 290. The former sometimes occurs in CNM output because of the resolution limit, and the latter sometimes occurs in HQ-cut output as its stopping criterion encourages splitting communities with high modularity. The average number of target communities in our LFR instances is roughly 150. For each sample size, the critical value is the 95th percentile of computed K–S statistic values. We used 100,000 trials per sample size.

After computing critical values, we evaluated the K–S statistic for each of our trials at each value of μ . If we reject the null hypothesis then we have 95% confidence that the algorithm results do not follow the discrete power-law distribution, Table II summarizes our results for all instances, broken down by algorithm type and μ value.

Both Figure 3 and Table II expose a phenomenon we call fracturing. We refer to the communities defined by LFR as target communities. There is no guarantee that target communities will be minimal natural communities. In fact, the subgraph induced by a target community is itself a random graph, and therefore we expect these to contain minimal natural communities occasionally. Modularity-based algorithms such as CNM, wCNM, and HQcut will find these smaller communities when they exist. In Table II, note that wCNM5 fails more K-S tests than does wCNM3 with increasing μ . As we add more iterations to the edge weighting scheme described in Section VI, we enable wCNM to resolve smaller communities. The most plausible explanation for the increased K-S failure rate of wCNM5, holding μ constant, is that we detect smaller communities whose sizes were not drawn from the LFR power-law.

Figure 3 (b) corroborates this observation. Note that for wCNM5, the quantile of target community size 10 corresponds to that of discovered community size less than

5. Figure 3 (c) shows that HQcut also finds communities smaller than size 10.

Algorithms such as wCNM and HQcut ascribe hierarchical community structure to a graph based on modularity. Some members of a large collection of random graphs, such as the LFR target communities, will have statistically significant sub-communities. Lang [27] uses an information theoretic metric to distinguish random graphs from those with community substructure. We conjecture that Lang's method will judge some LFR target communities to be non-random. Modularity-based algorithms will find substructure in these cases.

	LFR μ					
Algorithm	0.1	0.2	0.3	0.4	0.5	
CNM	0/29	0/30	0/30	0/30	0/29	
wCNM1	17/29	0/30	0/30	0/30	0/29	
wCNM3	28/29	29/30	29/30	23/30	0/29	
wCNM5	28/29	30/30	14/30	0/30	0/29	
HQcut	12/29	5/30	2/30	2/30	0/29	

TABLE II: This table shows Kolmogorov-Smirnov (K–S) results for experiments with 5000-vertex LFR instances (the number of passed tests divided by the number of instances). The critical values for the test were derived empirically by computing the K–S statistic for 100,000 samples, for each possible sample size between 10 and 290 communities. The hypothesis test results presented are at the 95% confidence level

We have not included formal running-time comparisons since Ruan and Zhang's publically available HQcut implementation is in Matlab and our implementation of wCNM is in C++. For anecdotal purposes, the wCNM runs on our 5000-vertex LFR instances took roughly 10s on a 3Ghz workstation, even with several iterations of weighting. The HQcut instances took 5-10 minutes on the same machine, though there were instances that took many hours. We killed such instances, and that is why we sometimes present fewer than 30 instances of HQcut results per μ .

IX. STATISTICAL CONFIDENCE

In recent work, several authors address the issue of statistical significance in detected communities. Bianconi, et al. [28–30] assess the relevance of a given community partition in explaining the topology of a given network. Bianconi [29] defines the entropy Σ_c of the ensemble of networks that are consistent with the given degree distribution and community assignments. This work leads to a z-score that quantifies the significance of the original community assignment compared to many random permutations of that assignment. Ref [29] also derives empirical confidence intervals for testing the null hypotheses that the given community structure is not significant. Unfortunately, these entropy computations rely on math-

LFR μ	child test pass rate	parent test pass rate	LFR μ	child test pass rate	parent test pass rate	
0.1	21/77 = 0.272	37/37 = 1.000	0.1	167/176 = 0.948	83/84 = 0.988	
0.2	4/41 = .097	20/20 = 1.000	0.2	157/167 = 0.940	83/84 = 0.988	
0.3	2/24 = .083	9/9 = 1.000	0.3	150/162 = 0.925	78/82 = 0.951	
0.4	2/18 = .111	8/8 = 1.000	0.4	141/167 = 0.844	67/79 = 0.848	
0.5	9/18 = .500	6/6 = 1.000	0.5	102/138 = 0.739	50/67 = 0.746	
(a) CNM			(b) wCNM3			

TABLE III: The results of parent–child tests on LFR instances. CNM under-resolves, so all dendrogram parents have substructure

ematical techniques that constrain the input to networks with fewer than \sqrt{N} communities. This is a regime we largely avoid by tolerating the resolution limit, so the method of Bianconi, et al. does not apply in our case.

Lancichinetti, et al. [31] seek to put a confidence score, called "c-score," on individual communities. This supports a hypothesis test per community, and we ran this test on the communities generated by our algorithm (wCNM), the Ruan and Zhang algorithm (HQcut), and the ground truth communities (generated by LFR). Our algorithm did consistently better than HQcut, and significantly better in some important cases. For example, with LFR graphs that have $\mu = 0.2$ (fairly well-defined communities), 91% of the communities wCNM found are judged to be significant by the c-score test, compared to only 66% of the communities found by HQcut. Roughly 95% of the LFR ground truth communities passed the test. However, it is problematic to present such aggregate results as evidence for statistical significance in the whole community assignment due to lack of independence. Each c-score computation operates on vertices outside of the community being evaluated.

Below, we present a double hypothesis test as our main evidence for statistical confidence in our wCNM solutions. We use the term "confidence" rather than "significance" because we provide evidence, but compute no p-values.

The wCNM algorithm returns a dendrogram containing hierarchical community structure. Each dendrogram node represents a subgraph. Some dendrogram nodes represent subgraphs compiled by a sequence of merges with positive change in weighted modularity. This set of subgraphs partitions the vertex set and determines the wCNM communities. The parents of these nodes in the community hierarchy are subgraphs that are not communities, but have at least one community as an immediate child. We tested not only the communities, but the parents as well. This will provide some evidence that the detected communities are not simply random fluctuations in a larger, random subgraph.

After the dendrogram has been computed, we evaluate each wCNM community using a significance test very similar to that of Ruan and Zhang [10]. The only exception is that Ruan and Zhang rely on an unknown distribution to compute their z-score, while we find empirical 95th percentile confidence bounds. For each detected

community we generate 20 randomly rewired instances and use wCNM to compute a weighted modularity for each. Like Ruan and Zhang, we rewire using the switching algorithm abstracted in [32] and discovered independently many times. After sorting these 20 numbers, the second and nineteenth determine an empirical 95% confidence region. If the weighted modularity of the original subgraph is within this range, we do not reject the null hypothesis that the subgraph is plausibly random. We call this the *child test* and consider it passed if the subgraph is found to be plausibly random. A parent test is performed on a parent subgraph and passed if the null hypothesis is rejected.

If a community passes the child test, it is plausible that it has no substructure other than random fluctuations. By construction, the HQcut algorithm is designed to return community partitions with 100% pass rate on both child and parent tests. Their method is top-down, and it splits a subgraph only if the parent test is passed. It stops only if the child test is passed. Despite this apparent statistical significance, Table 3 shows that HQcut consistently over-resolves the problem into communities that are too small, and visualization of the subgraphs themselves confirms that LFR ground truth communities are split into multiple pieces.

Unlike HQcut, the wCNM algorithm makes no decisions based on statistical tests. Our approach finds solutions that are closer to the LFR ground truth. Figure III contains the results of our parent–child tests on CNM and wCNM solutions. Since CNM under-resolves, the parent subgraphs are large and contain substructure. The wCNM parent–child pass rates are better balanced, and they provide some evidence that through $\mu=0.3$, we find communities that typically (1) do not have substructure other than random fluctuations, and (2) don't occur as random fluctuations in their parent subgraphs.

A community detection algorithm should not find structure in a purely random graph. Agglomerative algorithms such as CNM and wCNM will find such structure. However, our parent—child test can be used to mine the resulting dendrograms and disqualify these community assignments. Therefore, the combination of wCNM and the parent—child dendrogram test guards against the possibility of claiming significant community structure in random graphs.

X. CONCLUSIONS

Our weighted analogue to Fortunato and Barthèlemy's resolution argument shows that greater resolution in community detection is possible. We demonstrate this empirically with a simple adaptation of the CNM heuristic. Our algorithm is able to resolve communities of widely varying sizes in test data.

The CNM and wCNM agglomerative heuristics construct dendrograms of hierarchical community structure, and therefore contain more information than the flat community structure. Small modules within a community are represented in the dendrogram as well as the super-graph (parent) that contains a given community.

We mine the dendrograms produced by CNM and wCNM for this information, and use our parent—child test to check whether most of the children (communities) do not contain substructure, while most of the parents do. This two-sided test helps build confidence that the communities we find are not simply random fluctations in some larger subgraph. Our confidence is limited, however, because we do not tackle the intractable problem of examining all possible super-graphs of a community. Rather, we use the dendrogram parents to constrain the choice of larger subgraphs.

The edge weighting we describe is just one of many possible alternatives, and wCNM is just one of many potential weighted modularity algorithms. The primary contribution of this paper is to spread awareness that resolution limits may in fact be tolerated while retaining the advantages of modularity maximization and the efficient algorithms for this computation.

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