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Detecting Highly Overlapping Community Structure by Greedy Clique Expansion

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ABSTRACT
In complex networks it is common for each node to belong to several communities, implying a highly overlapping community structure. Recent advances in benchmarking indicate that the existing community assignment algorithms that are capable of detecting overlapping communities perform well only when the extent of community overlap is kept to modest levels. To overcome this limitation, we introduce a new community assignment algorithm called Greedy Clique Expansion (GCE). The algorithm identifies distinct cliques as seeds and expands these seeds by greedily optimizing a local fitness function. We perform extensive benchmarks on synthetic data to demonstrate that GCE’s good performance is robust across diverse graph topologies. Significantly, GCE is the only algorithm to perform well on these synthetic graphs, in which every node belongs to multiple communities. Furthermore, when put to the task of identifying functional modules in protein interaction data, and college dorm assignments in Facebook friendship data, we find that GCE performs competitively.

Categories and Subject Descriptors: H.2.8 Database Management: Database Applications – Data Mining
Keywords: Community Assignment, Overlapping, Local Clustering Algorithm, Complex Networks

1. INTRODUCTION

Community structure has been recognized in networks that come from a wide range of domains, such as social and biological networks. While concrete definitions of community vary by domain, a community may generally be described as a set of nodes with dense internal connections, exhibiting comparatively sparse connections to the rest of the network. Knowledge of community structure can reveal functional organization in networks, much as identifying organs in the body can reveal the role of various tissues. In recent years, numerous community assignment algorithms (CAAs) have been suggested, as computer scientists and physicists have taken on the problem of algorithmically finding communities (for an excellent recent review of the field, see Fortunato [1]).

Despite their proliferation, it is difficult to determine the performance of CAAs for two reasons. On the one hand, there is a lack of large empirical datasets where the a priori or ground truth communities are known; and on the other hand, most synthetic data—especially the most popular, the GN model [2]—is overly simplistic and unrealistic, lacking key topological features such as a heterogeneous degree distribution, varied community sizes, and triadic closure, while also requiring that every node belong to exactly one community. The lack of realistic benchmark graphs has led to a situation where researchers know that many algorithms perform well on simple networks, but are unaware how these perform on more complex empirical data.

This problem is so pronounced that in his comprehensive review of the field, Fortunato states with regard to benchmarking: “...the issue of testing algorithms has received very little attention in the literature on graph clustering. This is a serious limit of the field. Because of that, it is still impossible to state which method (or subset of methods) is the most reliable in applications...”

In the last year, Lancichinetti and Fortunato [3] have addressed this uncertainty by specifying a means of creating more realistic synthetic benchmark graphs, which have scale-free degree and community size distributions as well as overlapping communities. Using their specification (called LFR), they and others have subsequently discovered—with a level of subtlety previously unattained—under what topological conditions a wide range of CAAs perform well or poorly [4,5].

One surprising result revealed by this recent benchmarking is the poor performance of many CAAs when it comes to detecting moderately overlapping community structure. It is intuitive from our knowledge of real world domains that many complex networks will have communities that overlap, potentially to a high degree. Consider, for example, a social network site like Facebook. On average, a Facebook user has 130 “friends,” who typically belong to multiple distinct social groups [6]. These groups may correspond to ties formed in high-school, college, professional settings, and family. Figure 1 which depicts the ego-centric network of a Facebook user, demonstrates this tendency for a user to belong to multiple groups. The analysis of Marlow et al. [7] suggests that the groups apparent in this user’s ego-centric network correspond to acquaintances formed at different stages of life, and that most of these groups are dormant. Clearly, if this type of ego-centric network is typical of Facebook users, then any CAA that partitions nodes into non-overlapping communities (henceforth, non-overlapping CAA) will perform poorly: such CAAs can assign each node to only one of its many communities. Similarly, in complex networks of interactions between proteins, it has been claimed that many proteins belong to multiple communities, each of which in turn corresponds to some biological function [8,9]. Since 2005, the year in which Palla et al.
published a groundbreaking CAA capable of detecting overlapping communities, a number of algorithms have been developed that are able to assign nodes to more than one community \cite{9,5,16}. However, using LFR networks \cite{4,5} and other synthetic networks \cite{8}, recent work has indicated that many CAAs that are supposed to be capable of detecting overlapping community structure perform quite poorly when more than a minority of nodes belong to multiple communities.

The purpose of this paper is two-fold. On the one hand, in sections 2 and 3 we introduce a new algorithm, called Greedy Clique Expansion (GCE) 1 that is designed to perform well in domains with highly overlapping community structure. On the other hand, in section 4 we thoroughly benchmark this algorithm alongside several other leading CAAs that are designed to detect overlapping community structure. We run benchmarks on graphs with high levels of community overlap to clear up uncertainty of the performance of CAAs designed for this domain. None of these CAAs have been subjected to such benchmarks. Our results indicate that GCE is the only algorithm capable of accurately detecting communities when nodes belong to several communities.

The fitness function is of central importance to the GCE algorithm; it can be interpreted as steering the growth of a seed such that it expands into the desired community. Just as there is no universally correct concept of community that spans all domains, one cannot argue that any given fitness function will be appropriate for all types of network data.

Nevertheless, in our experiments, we found that the fitness function defined by Lancichinetti et al. \cite{14} provided good results on a wide range of synthetic and empirical data. They define the fitness of a community \( S \) in terms of \( S \)'s internal degree \( k_{in}^S \) and external degree \( k_{out}^S \). \( k_{in}^S \) is equal to twice the number of edges that both start and end in \( S \) (i.e., it is the sum of the internal degrees of the nodes in \( S \)), and \( k_{out}^S \) is the number of edges that have only one end in \( S \). In this notation, they define community fitness as

\[
F_S = \frac{k_{in}^S}{(k_{in}^S + k_{out}^S)^\alpha},
\]

where \( \alpha \) is a parameter that can be tuned. Lower values of \( \alpha \) result in larger communities being fitter. We found that \( \alpha \) values in the range 0.9 – 1.5 provided the best results, which is in line with the experience of Lancichinetti et al. We resume discussion of this parameter below.

Expanding a single seed. Assume, as above, that \( S \) is an induced subgraph of \( G \) that can be thought of as the “seed” or core of a community \( C \). In other words, \( S \) is embedded in some larger community \( C \), such that all of its nodes are part of \( C \), but not all nodes in \( C \) are included in \( S \). The task at hand is to expand \( S \) by adding nodes to it until it includes all nodes in \( C \). Previous work in community assignment suggests that—by utilizing a community fitness function such as eq. \( \{1\} \)—\( S \) can be efficiently expanded into \( C \).
through a technique of greedy local optimization. \cite{10, 14, 15, 17}

This technique can be varied, but can generally be summarized in the following steps:

1. For each node $v$ in the frontier of $S$ (e.g., the red nodes in fig.1), calculate $v$’s node fitness, i.e., how much the addition of $v$ to $S$ would raise or lower the community fitness of $S$.

2. Select the node with the largest fitness, $v_{\text{max}}$.

3. If $v_{\text{max}}$’s fitness is positive, then add it to $S$ and loop back to step 1. Otherwise, terminate and return $S$.

The works cited above vary this technique by either using different fitness functions or, for example, after each addition, removing any nodes in $S$ if their removal would improve $\text{F}(S)$. Also, they vary in their approach of finding starting seeds. In general, this approach of greedy local optimization scales well with the size and order of the network because it works locally; we discuss complexity in more detail in section \[3\].

While GCE shares this general strategy of expanding seeds via greedy local optimization, one key difference from previous algorithms is the choice of starting seed.

**Cliques as starting seeds.** Various approaches have been used to find the seeds of communities in the above-mentioned greedy algorithms. Lancichinetti et al. \cite{14}’s LFM algorithm keeps randomly selecting nodes that have not yet been assigned to any community, until every node belongs to at least one community. This method assumes that each node belongs to at least one community, and that, as soon as every node has been assigned to at least one community, there are no further communities that should be found. We take issue with both of these assumptions; we believe that the latter—by implicitly placing an upper bound on the number of communities that can be found at $|V|$—is responsible for the declining performance of LFM as each node tends to belong to more communities in the benchmarks that follow.

Baumes et al. \cite{15}’s Iterative Scan algorithm selects random edges as seeds, and keeps expanding seeds until the new seeds produce communities that are duplicates of previously-found communities. While this method of seed selection is less arbitrary than \cite{14}’s, it is inefficient because, if one wants to exhaustively search for seeds, the algorithm is unlikely to terminate before it has expanded a vast number of duplicate communities.

We use maximal cliques (i.e., fully-connected sets of nodes that are not completely contained in any larger set of fully-connected nodes), as seeds; we will henceforth refer to maximal cliques simply as cliques. This choice of seeds is motivated by the observation that, on the one hand, cliques are one of the characteristic structures contained within communities \cite{13}, while on the other hand—if one discards smaller cliques that are highly embedded in larger cliques—they are rare structures. We note that other CAAs exploit these properties of cliques \cite{9, 16, 19}, but none of them utilize cliques as seeds in the greedy expansion strategy mentioned above.

One of the key parameters of our algorithm, $k$, is the minimum number of nodes that a clique must contain if it is to be accepted as a seed. On the one hand, $k$ should be sufficiently large such that any clique of size $k$ or larger indicates the presence of a community; otherwise, one risks expanding a seed into a region with no community structure. For example, if $k = 3$, then triangles can be accepted as seeds. In some networks, one cannot assume that all triangles are embedded in a community. In such cases, if $k = 3$, then one risks accepting seeds that are not embedded in any community, and these seeds could expand into a region of the graph with no community structure. Such communities can be thought of as false positives. On the other hand, one should choose $k$ to be sufficiently small such that all of the communities that one wishes to detect contain at least one clique of size $k$. If $k$ is chosen to be too large, then those communities that ought to be detected, but contain no sufficiently large cliques, will not be detected. Such cases could be considered false negatives. We find that $k$ values of 3 or 4 will generally satisfy these constraints, and that one can decide between these two possibilities by considering whether one’s preferred semantic definition of community includes small structures.

This choice of seeds comes with an implicit requirement that any community that can be found by GCE must contain a clique with $k$ or more nodes. One might object that this assumption is too strict and will result in many false negatives. However, our results on the LFR benchmark graphs indicate otherwise. The LFR benchmarks provide a challenge in this respect because the process which creates communities does not favor the generation of cliques. In particular, creates edges such that the probability that any pair of nodes in a community is connected by an edge is independent of whether those two nodes share neighbors, much as in a classic Erdős-Rényi random graph \cite{1}. This generation technique does not lead to the high number of triangles and cliques that have been observed in empirical graphs. Empirical networks show a strong tendency for transitivity, i.e., for two neighbors of a given node to be connected to each other \cite{20, 21}, a process which leads to higher clustering coefficients and more cliques than one would expect to find in correspondingly sparse Erdős-Rényi graphs. Thus, the fact that GCE performs well on these synthetic graphs—despite the fact that one expects fewer cliques in these than in empirical data—indicates that this minimum clique size requirement does not cripple the sensitivity of GCE.

**Duplicates and community distance.** Overlapping CAAs must include some implicit or explicit strategy for dealing with near-duplicate communities (in contrast to the more numerous, non-overlapping CAAs, whose communities can share no nodes). This problem arises from the fact that many seeds can grow into near-duplicate versions of the same community. This is undesirable from the perspective of the network analyst because results that contain a large number of near-duplicate communities are hard to interpret and report statistics on.

To rid our results of near-duplicate communities, we must formally define what we mean by near-duplicate communities. Along the lines of Baumes et al. \cite{15}, we begin by defining a community distance measure. We choose a symmetric measure of community distance that can be thought of as the percent non-embedded. Given two communities $S$ and $S'$, this measure is defined as

$$
\delta_E(S, S') = 1 - \frac{|S \cap S'|}{\min(|S|, |S'|)},
$$

which can be interpreted as the proportion of the smaller community’s nodes that are not embedded in the larger community.

Given a set of communities $W$ and a community $S$, we can define the near-duplicates of $S$ to be all communities in $W$ that are within a distance $\epsilon$ of $S$, where $\epsilon$ is the minimum community distance parameter.

**Overview of GCE.** Now that we have covered the requisite concepts of community fitness, expanding a seed, choosing seeds, and near-duplicate seeds, we can outline the GCE algorithm. Given a graph $G$, a minimum clique size $k$, a minimum community distance $\epsilon$, and a scaling parameter $\alpha$, our algorithm:

1. Finds seeds by detecting all maximal cliques in $G$ with at
least \( k \) nodes.

2. Creates a candidate community \( C' \) by choosing the largest unexpanded seed and greedily expanding it with a community fitness function \( F \) until the addition of any node would lower fitness.

3. If \( C' \) is within \( \epsilon \) of any already accepted community \( C \), then \( C \) and \( C' \) are near-duplicates, so discard \( C' \). Otherwise, if no near-duplicates are found, accept \( C' \).

4. Continues to loop back to step 2 until no seeds remain.

We note that although GCE allows the user to specify the values of three parameters, two of these—\( k \) and \( \epsilon \)—allow for versatile default values. The value of \( k \) should usually be 4; if one is interested in very small communities (as in the case of the protein complexes of three parameters, two of these—\( k \) and \( \epsilon \)—allow for versatile default values. The value of \( k \) should usually be 4; if one is interested in very small communities (as in the case of the protein complexes presented in section 5), then \( k \) should be set to 3. We find 0.25 to be a good default value for \( \epsilon \)—if one finds too many near-duplicate communities in the output, then \( \epsilon \) should be increased.

The scaling parameter \( \alpha \) lends itself least to a versatile default value. For best results, one should first run GCE with \( \alpha \) set to 1.0, look at the results, and decide whether the communities found by GCE should be larger (using a lower \( \alpha \)) or smaller. However, in cases where the user knows little about the community structure, such tuning may be difficult. For this reason, in the synthetic and empirical benchmarks that follow, we report results where \( \alpha \) is set to 1.0, rather than tuning this parameter for best results.

The good benchmarking results indicate that even if one cannot tune \( \alpha \), GCE will often return good results with all parameters set to their default values.

3. OPTIMIZATIONS

We begin this section by underscoring the point that one cannot satisfactorily characterize the average complexity of GCE purely in terms of \( |V| \) or \( |E| \); rather, the complexity depends on subtler local characteristics of \( G \) that are difficult to specify rigorously. Despite this, we can clearly discuss several important parts of the algorithm, and the heuristics and optimizations we have used to improve their performance.

First, we discuss finding the maximal cliques that form seeds, and consider the complexity of greedily expanding each seed. The remainder of this section is devoted to various problems related to detecting near-duplicate communities. We consider the cost of deciding whether a candidate community is sufficiently distinct to accept, and two heuristics to discard potentially indistinct seeds early. While our solutions to these problems are somewhat trivial, we describe them in detail both for replications and because the computational savings that they afford are so significant.

We note that GCE makes extensive use of set operations, such as set insertion and deletion and testing for set membership. In our implementation, we use the C++ standard template library to provide these operations.

Finding Maximal Cliques. Although finding all of the cliques in a graph is generally computationally expensive, cliques can be found quickly in graphs that are sufficiently sparse. To this end, our implementation makes use of the Bron-Kerbosch [22] clique enumeration algorithm to efficiently find the maximal cliques that form seeds. In the large synthetic and empirical networks that we analyze in section 4 and section 5, the computation required for finding cliques was a small part of the overall run time, compared to the computation required to expand seeds and check for near duplicates. To further support the claim that finding cliques in sparse graphs is scalable, we point out that Schmidt et al. [23] have recently introduced a parallel variant of the Bron-Kerbosch algorithm, which they demonstrate can achieve a linear parallel speed-up even when using 2048 processors.

Greedy Expansion. Greedy seed expansion requires that the frontier \( f(S) \) of each initial seed be identified, and that \( f(S) \) be updated as \( S \) is expanded.

The initial frontiers may be identified by calculating, for each edge, the symmetric difference of the sets of seeds for the endpoints of that edge. Identifying the initial frontiers therefore has complexity \( O(|E| \times M) \), where \( M \) is the number of cliques to be expanded.

As each seed is expanded by adding the fittest node from the frontier \( v_{\text{max}} \), its frontier becomes:

\[
 f(S \cup v_{\text{max}}) = (f(S) \cup N(v_{\text{max}}), S) - \{v_{\text{max}}\}
\]

where \( N(v_{\text{max}}) \) is the set of neighbors of \( v_{\text{max}} \). This requires at most \( \theta \) insertions into \( f(S) \), where \( \theta \) is the maximum degree in \( G \).

We note that the fitness \( F_{S \cup v_{\text{max}}} \) depends only on the internal and external degree of \( v_{\text{max}} \) and the total internal and external degrees of the vertices already in \( S \). To facilitate fast identification of \( v_{\text{max}} \), \( k^I_{\text{max}} \) and \( k^O_{\text{max}} \) are stored, along with the internal and external degrees of each \( v \) in \( f(S) \). These stored values are updated each time \( f(S) \) is updated, noting that \( k^I_{\text{max}} \) and \( k^O_{\text{max}} \) each change only by the internal and external degree of \( v_{\text{max}} \) and that the frontier degrees need only be updated for \( v \in N(v_{\text{max}}) \).

Identifying Near-duplicate Communities. To identify if a candidate community, \( C \), is a near-duplicate of an accepted community, it is necessary to calculate the overlap between community pairs. Finding the intersection between sets \( C_1 \) and \( C_2 \) has complexity \( 2(|C_1| + |C_2|) - 1 \), assuming sorted sets. In a naive implementation, after each seed expansion has completed, this must be carried out \( O(\zeta) \) times, with \( \zeta \) the number of accepted communities, resulting in at least \( O(\zeta^2) \) set intersections in total. Instead, we maintain, for each node \( v \), the set \( c(v) \) of accepted communities it belongs to. In a first pass, we identify those communities that have a non-empty overlap with the candidate community as \( \bigcup_{c \in C} c(v) \). The full intersection is then calculated only on the communities with non-empty overlap. At the cost of extra storage, this results in significant time savings as the number of accepted communities grows large.

Clique Coverage Heuristic (CCH). While developing this algorithm, we observed that many empirical datasets exhibited the following phenomenon: given some maximal clique, there exist numerous smaller cliques that are almost, but not fully, subgraphs of it. Shen et al. [16] have also observed this property; in fact, this is the key property exploited by k-clique percolation. [9] (Remember, whenever we use the term "clique" here and throughout our paper, we mean "maximal clique.") Typically, these cliques are within \( \epsilon \) distance of the larger clique and thus are near-duplicates of it.

We also observed that when expanded, such sets of near-duplicate cliques were likely to grow into the same or very similar regions of the graph, leaving us with many near-duplicate communities that would later need to be removed (as specified in subsection “Dupli cate and community distance” above). We could thus skip the expansion of these near duplicate seeds without significantly affecting our results and enjoy a significant savings in memory and computation time. We developed a simple heuristic to quickly prune such near-duplicate cliques from the our initial set of seeds. This method is designed solely as a heuristic computational speedup, and has
been found to effectively discard large numbers of near-duplicate cliques while not significantly altering benchmark results.

The Clique Coverage Heuristic (CCH) is as follows: We order the maximal cliques, largest first. Each clique is then either accepted or rejected. A clique is accepted if less than a certain proportion of its nodes are contained in at least two of the larger cliques that have already been accepted. We found that even values of \( \phi \) close to 0 resulted in the large numbers of near-duplicate cliques being rejected from consideration. Simultaneously, it appears that in each of the clique-dense areas of the graph that are likely to be embedded in a community, at least some of the original cliques are always preserved. Thus, each of these areas remains “seeded” with at least one clique, but most of the smaller, near-duplicate versions of this clique are removed. In our evaluation we choose a value of 0.25 for \( \phi \) – meaning that a clique will only be discarded if at least 75 percent of its nodes have already been covered twice by other, larger, accepted cliques.

Due to the potentially vast number of near-duplicate cliques found in complex networks, the impact of CCH can be large. For example, we ran GCE on the Oklahoma State Facebook subnetwork, which contains 17425 nodes and 829528 edges (the source of this data set is Traud et al. [24], which is described in section 5). On this data, CCH reduced the number of cliques from over 46 million to around 5000. To address concerns that pruning seeds using CCH may adversely affect the accuracy of GCE, we refer the reader to fig. 5 which includes one line displaying the results of GCE with the CCH, and one line for GCE without the CCH. The results are almost identical.

Abandoning suspiciously overlapping seeds. Let us call any seed that is undergoing expansion that is within some distance \( \Delta \) of an already accepted expanded community ‘suspicious’. We call it suspicious because for an appropriate value of \( \Delta \), such expanding seeds frequently expand to within \( \epsilon \) (i.e. become near-duplicates) of an already accepted community. In these cases, the computation required to expand the suspicious seed into a near-duplicate community is essentially wasted. One simple optimization, which we employ in all of the evaluations below, is to discard these suspicious seeds. In practice, we have found that the benchmark results of our implementation are not very sensitive to a range of \( \Delta \) values, and we use the value of 0.6 for both \( \Delta \) and \( \epsilon \).

Performance characteristics. We now briefly describe two experiments to reveal both a strength and a weakness of GCE’s runtime performance (we postpone discussion of accuracy, as this is the topic of the next section). These experiments are based on the above-mentioned LFR graphs, which are synthetic graphs whose specification will be explained in detail in section 4. The parameters used to construct the graphs used in these experiments are listed in the first two columns of table 1.

While the details of these graphs may not be clear for the reader until the next section, the essential point is that in fig. 3 we run GCE on a series of graphs that are identical in many ways—sharing the same degree distribution and community size distribution—but which have ever-more nodes and communities. In this figure we observe that as the size of the graph increases along the x-axis, the runtime of GCE scales favorably when compared to the runtime of other overlapping CAAs (which are also introduced in the next section).

In fig. 4 we again observe the runtime of GCE on a series of graphs that are nearly identical; however, in these graphs we do not vary the number of nodes (keeping it fixed at 5000), but rather we vary the degrees of the nodes. As the average degree increases along the x-axis, the runtime of GCE increases quite rapidly when compared with the quickest algorithm run on this experiment, COPRA (which is also introduced in the next section). We observe that even on these relatively small graphs, the runtime of GCE is quite sensitive to the average degree. In this figure, we also see the effect of the CCH optimization.

4. SYNTHETIC BENCHMARKS

In the following section, we pursue two objectives. First, we perform the central experiment of this paper: we benchmark many CAAs on the synthetic graphs in which nodes belong to up to five communities. Benchmarks with such a high degree of community overlap are uncharted territory. The results of the benchmarks are surprising, indicating that many algorithms that are specifically designed for detecting overlapping community structure perform poorly when more than a fraction of nodes belong to more than one community. While one other algorithm performs well up to the point where nodes belong to an average of 1.9 communities, GCE returns good results even when every node belongs to four communities.

Unfortunately, because benchmarks with such high levels of communities have not yet been carried out, we had to choose many of the parameters for creating the benchmark graphs. The secondary purpose of this section is to show that GCE also performs well on benchmarks graphs whose parameters have been defined by other,
less interested parties. To this end, we replicate a set of partitioning benchmarks performed by [4] in a recent review of several CAAs. We demonstrate that GCE performs competitively against the best known CAAs. This is significant, because these other algorithms are specialized for graph partitioning, while GCE is designed to handle the more general case of overlapping communities.

**Benchmarking procedure and terminology.** To benchmark the performance of a CAA, we perform the following steps: first, we create a synthetic graph which, by construction, contains communities planted in it. We will refer to these communities as the ground truth communities. Next, we run a CAA on this graph; we call the communities returned by the CAA the found communities. Finally, we use some metric to compare the similarity of the ground truth communities to the found communities.

To construct synthetic graphs, we use the LFR specification, which allows one to create graphs with realistic properties such as scale-free degree and community size distributions. [3]. To measure the similarity of ground truth communities and found communities, we use normalized mutual information (NMI), an information-theoretic similarity measure. This measure is normalized such that the NMI of two sets of communities is 1 if they are identical, and 0 if they are totally independent of each other. Danon et al. [25] first applied NMI to the problem of evaluating the similarity of two sets of communities, but defined the measure only for partitions. In our benchmarks, we employ a variant of NMI introduced by Lancichinetti et al. [14] that is defined for covers, in which nodes may belong to multiple communities.

**Synthetic graph description & parameters.** To construct a LFR graph, one must specify ten parameters, which are listed in table 1. Note that the parameter \( O_n \) refers to the number of nodes in the graph that are overlapping (i.e., belong to more than one community), and \( O_m \) dictates how many communities each of the overlapping nodes belongs to.

<table>
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<th>Description</th>
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<th>fig. 4</th>
<th>fig. 5</th>
<th>fig. 6</th>
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<td>( N ) number of nodes</td>
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Table 1: Parameters for LFR synthetic benchmark graphs for various figures. Bolded values indicate the variable on the x-axis of each plot.

Benchmarks on graphs with overlapping communities. As we mentioned in the introduction, very little work has been done on benchmarking CAAs on graphs with overlapping community structure. The work that includes such benchmarks is limited to graphs with only moderate levels of community overlap, i.e., where some fraction of nodes belong to two communities, and the rest belong to one [4][5][8].

Our purpose here is to examine how CAAs perform on graphs with a higher degree of community overlap. We specify five graphs; in the first, each node belongs to one community, and in each successive graph all nodes belong to one more community, so that in the fifth graph, all nodes belong to five communities. In order for nodes to have enough edges for membership in an increasing number of communities, in each successive graph nodes are assigned a higher average degree. In the first graph, the average node degree is \( k = 18 \), and for each extra community that a node belongs to, the average degree increases by 18, so that by the fifth graph, \( k = 90 \). Other parameter values remain constant across all five graphs and are listed in the third column of table 1. In all synthetic and empirical benchmarks, we kept GCE’s parameters fixed at \( k = 4, \alpha = 1.0, \) and \( \epsilon = 0.6 \), with the exception that in the PPI benchmark, we set \( k = 3 \) because the domain contains very small communities.

In fig. 5 (A) and (B), we see the results of GCE alongside five other CAAs that were designed specifically to detect overlapping communities, and two that find non-overlapping communities. The five other overlapping CAAs that we benchmarked are CFinder, which employs a technique of k-clique percolation [3]; LFM, which uses a local greedy optimization strategy very similar to GCE, but selects seeds randomly [14]; COPRA, which utilizes a label propagation technique [6]; abchampions, which finds all regions of the graph with a certain difference between internal density and external sparsity [11]; and Iterative Scan [15], which we have described in section 2. All implementations we used were from the authors. Just as GCE’s parameters were fixed at default values, we left the parameters of the other algorithms set to their defaults with the following exceptions: CFinder, where we set \( k = 4 \), which returned the best results overall; COPRA, where we set \( v = 3 \), Iterative Scan, where we set the initial cluster size to 2, as recommended; and abchampions, where we set \( c_{min} = 5 \) and \( c_{max} = 100 \), as recommended by an author. For each point in the plot, we ran all CAAs on graphs with such a high degree of overlap: none of the existing algorithms perform well even when overlap is kept at moderate levels. The robust performance of GCE as the number of communities to which each node belongs increases is unprecedented. To more closely examine the limitations of the other three algorithms, in fig. 5 (B) and (D) we run additional benchmarks on graphs where some fraction of nodes belongs to two communities and the rest belong to one. In these graphs, the average degree \( k \) steadily rises from 18 (when each node belongs to one community) up to 36 (when every node belongs to two communities).

With regard to CFinder and COPRA, our results here mirror the results of [4][9], who benchmarked these algorithms on similar LFR-generated graphs. They also found that these algorithms could no longer accurately assign nodes to communities if all nodes belonged to two communities. We benchmark LFM for the first time on such overlapping synthetic graphs. It is interesting to note that although LFM and GCE use the same fitness function and a similar greedy heuristic, their results vary so greatly, uses the same fitness parameters; error bars represent the standard deviation of NMI over the 20 runs, which also holds for the benchmarks in fig. 6.

Figure 5 (A) and (C) suggests why no benchmarks have been carried out on graphs with such a high degree of overlap: none of the existing algorithms perform well even when overlap is kept at moderate levels. The robust performance of GCE as the number of communities to which each node belongs increases is unprecedented. To more closely examine the limitations of the other three algorithms, in fig. 5 (B) and (D) we run additional benchmarks on graphs where some fraction of nodes belongs to two communities and the rest belong to one. In these graphs, the average degree \( k \) steadily rises from 18 (when each node belongs to one community) up to 36 (when every node belongs to two communities).

With regard to CFinder and COPRA, our results here mirror the results of [4][9], who benchmarked these algorithms on similar LFR-generated graphs. They also found that these algorithms could no longer accurately assign nodes to communities if all nodes belonged to two communities. We benchmark LFM for the first time on such overlapping synthetic graphs. It is interesting to note that although LFM and GCE use the same fitness function and a similar greedy heuristic, their results vary so greatly, uses the same fitness parameters; error bars represent the standard deviation of NMI over the 20 runs, which also holds for the benchmarks in fig. 6.
function as GCE. We speculate that LFM’s performance drops because this seed selection strategy causes it to prematurely give up on trying to expand new regions of the graph that have unidentified communities.

**Benchmarks on graphs with disjoint communities.** Most of the benchmarking of CAAs has been on graphs with non-overlapping communities. In particular, Lancichinetti and Fortunato [4] have benchmarked a wide variety of CAAs on a particular set of LFR graphs. Gregory [5] have recently followed suit and benchmarked more algorithms on this set of graphs, so we continue in this vein and benchmark GCE and a number of other CAAs to see how they perform on this emerging standard. There are four graph specifications that are included in this standard set. The graphs in this quartet are either small or large ($N = 1000$ or $N = 5000$), and have either small or large communities (ranging between 10 – 50 nodes or 20 – 100 nodes). The results are displayed in fig. 6.

It is informative to compare the accuracy of GCE as displayed in fig. 6 with the results of various disjoint CAAs in the recent comparative benchmarking of Lancichinetti and Fortunato [4]. We note that COPRA and CFinder have been previously benchmarked on graphs with the same specification and returned the same results, indicating that we have accurately replicated this benchmark and that it is reasonable to compare our results with theirs. With regard to this matter of comparison, we also point out that in their review, Lancichinetti and Fortunato also used the overlapping version of NMI, so that we can directly compare the results from fig. 6 to their results. The comparison indicates that GCE’s accuracy on the task of partitioning graphs with non-overlapping community structure is among the best, even when compared to non-overlapping CAAs, which specialize in this task.

More specifically, GCE clearly outperforms the classic divisive GN algorithm of Newman and Girvan [26], a similar divisive algorithm by Radicchi et al. [27], the EM method of Newman and Leicht [28], the Markov clustering algorithm (MCL) of Van Dongen [29], an information theoretic approach by Rosvall and Bergstrom [30], and a spectral algorithm by Donetti and Munoz [31]. Against other algorithms, results were mixed. GCE performed better than a method based on modularity optimization via simulated annealing by Guimera and Amaral [32] in all cases except where the graph size was small and the community size large. GCE performs similarly to the modularity maximizing algorithm of Blondel et al. [33], which was among the CAAs that Lancichinetti and Fortunato identified as a top performer, and slightly worse than the other two top performers: another information theoretic algorithm from Rosvall and Bergstrom [30], and a Potts model approach by Ronhovde and Nussinov [34].

**5. EMPIRICAL BENCHMARKS**

In this section we strive to demonstrate GCE’s ability to identify meaningful communities in the context of non-trivial empirical networks, for which ground-truths are available. We agree with [11] that small the social networks which are typically used as empirical benchmarks for CAAs, such as Zachary’s Karate club, provide insufficient validation of a CAA. While we were unable to find an ideal, large-scale graph in which the ground-truth is completely known, we find two reasonable data sets for this purpose: a protein-protein interaction (PPI) network which includes a set of proteins with known complexes, and a collegiate Facebook network in...
Figure 6: NMI of GCE and other overlapping CAAs on all benchmarks used by Lancichinetti and Fortunato [4] and Gregory [5].

Table 2: NMI of various algorithms on PPI data, along with number of communities $|C|$ and average comm. per node.

| Algorithm     | NMI   | $|C|$ | Avg |
|---------------|-------|------|-----|
| GCE           | 0.55021 | 119  | 0.861 |
| abchampions [13] | 0.52983 | 63   | 0.485 |
| Clique percolation [36] | 0.522748 | 114  | 0.744 |
| MCL [29]     | 0.414983 | 298  | 1.000 |
| Blondel      | 0.328344 | 212  | 1.000 |
| Iterative Scan [15] | 0.301171 | 230  | 6.975 |
| COPRA [5]    | 0.299514 | 513  | 1.101 |
| LFM [14]     | 0.270601 | 58   | 0.646 |

which the dorm assignments are known.

**Protein-protein interactions and protein complexes.** Protein complexes tend to correspond to groups of proteins with many interactions, and can thus be detected by CAAs. We use a set of known protein complexes as an approximate ground truth.

To construct the PPI network, we used the interaction data found in the Combined-AP/MS network [35], which contains 1622 proteins and 9070 interactions. For the ground truth communities, we used the complexes listed in the CYC dataset of known complexes [36], selecting only those complexes that were also in the PPI network. Because many of these complexes were simply edges or triangles that are not recognizable as network communities, we removed all complexes with fewer than four proteins from the ground truth. Consequently, we use the value of 3 as the minimum clique size for all clique based algorithms. Note also, we use SCP [36] instead of CFinder here, as the latter fails to terminate on this dataset. The resulting ground truth contains 880 proteins; 136 of these belong to more than one complex. As the ground truth contains only a modest amount of overlap, we compare GCE against some disjoint CAA algorithms as well as some overlapping CAA algorithms. In table 2, we show the NMI achieved by each algorithm, along with the number of communities found.

We see that GCE’s found communities have the highest NMI with the ground truth, followed closely by abchampions and the clique percolation method. However, one might object that because we cannot assume that our ground-truth set of complexes is complete (due to the possibility of undiscovered complexes), the NMI measure is imperfect.

For this reason, in fig. 7 we look at these PPI results in more detail using the $F_1$-score, a summary of the classification metrics recall and precision. To evaluate a CAA performance, we performed the following procedure. For each known complex in the ground truth, we selected the found community with the highest $F_1$-score. An $F_1$-score of 1.0 indicates a perfectly recovered protein complex. We plotted all best $F_1$-scores in a histogram.

The histograms show that of all the overlapping CAAs, GCE has the most perfectly-recovered communities and the fewest poor matches. MCL, a non-overlapping CAA, returned slightly better results. Due to space constraints, we display the $F_1$-scores in fig. 7. We also looked at the second-best $F_1$-score to verify that the algorithms are finding each complex once, and only once; this is in-
We have introduced an algorithm, Greedy Clique Expansion, which combines the graph structure based approach of clique-finding methods with the greedy expansion strategy found in other algorithms. We demonstrate that GCE can accurately recover communities on synthetic networks in which every node belongs to four communities. We found that no other algorithm performed nearly as well on synthetic graphs in which every node belong to two or more communities. To determine whether these good results are robust, we performed further comparative benchmarks on a range of LFR graphs with non-overlapping communities, and found that GCE performed competitively. To complete our evaluation, we used GCE to recover biological ground truth communities from a reference protein-protein interaction network, and to infer non-network attributes from a social graph. Compared to other overlapping CAAs, GCE gave the best results.

6. CONCLUSION

We have introduced an algorithm, Greedy Clique Expansion, which combines the graph structure based approach of clique-finding methods with the greedy expansion strategy found in other algorithms. We demonstrate that GCE can accurately recover communities on synthetic networks in which every node belongs to four communities. We found that no other algorithm performed nearly as well on synthetic graphs in which every node belong to two or more communities. To determine whether these good results are robust, we performed further comparative benchmarks on a range of LFR graphs with non-overlapping communities, and found that GCE performed competitively. To complete our evaluation, we used GCE to recover biological ground truth communities from a reference protein-protein interaction network, and to infer non-network attributes from a social graph. Compared to other overlapping CAAs, GCE gave the best results.

Further Work. The community structure found in some networks may exist at multiple scales due to hierarchical organization of the system represented by the network. Ideally, a CAA would detect community structure at all scales. We are currently working with a modified version of GCE that expands all seeds in parallel, merging them as they become near-duplicates. By expanding all seeds to encompass the entire graph, and merging them along the way, the algorithm produces a dendrogram similar in some respects to the dendrograms produced by classic agglomerative algorithms such as that of Girvan and Newman [2], but which allows overlap, and in which leaves represent nodes rather than seeds. Communities could be extracted from this dendrogram by performing an analysis of stability. This variant of GCE might have the advantage of not only detecting hierarchy, but also of decreased sensitivity to the $\alpha$ parameter, because communities are selected based on stability rather than on the first local maximum of fitness.

We considered only a simple, greedy expansion heuristic. Future work should investigate using more sophisticated local heuristics that cleverly explore the most promising sections of the search space. Furthermore, due to the local nature of GCE, a parallel implementation should be straightforward to implement and would increase its scalability.

Finally, we would like better benchmarking abilities. On the one hand, we need a utility that uses synthetic graphs to systematically explore under what topological conditions the performance of a CAA breaks down. On the other hand, we need better empirical networks with ground truth communities to ensure that CAAs do not merely perform well on synthetic data.

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References


