Online Search of Overlapping Communities

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ABSTRACT
A great deal of research has been conducted on modeling and discovering communities in complex networks. In most real life networks, including social networks and bio-chemical networks, an object often participates in multiple overlapping communities. In view of this, recent research has focused on mining overlapping community structures in complex networks. The algorithms essentially materialize a snapshot of the overlapping communities in the network. This approach has three drawbacks, however. First, the mining algorithm uses the same global criterion to decide whether a subgraph qualifies as a community. In other words, the criterion is fixed and predetermined. But in reality, communities for different vertices may have very different characteristics. Second, it is costly, time consuming, and often unnecessary to find communities for an entire network. Third, the approach does not support dynamically evolving networks. In this paper, we focus on online search of overlapping communities, that is, given a query vertex, we find meaningful overlapping communities the vertex belongs to in an online manner. In doing so, each search can use community criterion tailored for the vertex in the search. To support this approach, we introduce a novel model for overlapping communities, and we provide theoretical guidelines for tuning the model. We present several algorithms for online overlapping community search and we conduct comprehensive experiments to demonstrate the effectiveness of the model and the algorithms. We also suggest many potential applications of our model and algorithms.

1. INTRODUCTION
Most complex networks in nature and human society contain community structures that serve as functional building blocks for the networks. Furthermore, communities often overlap with each other, that is, a vertex in a network may belong to more than one community.

1.1 Overlapping communities
Overlapping community structures can be observed in many real networks, including social networks, biology networks, and semantic networks, and they are vital in revealing the internal structure of large networks [1]. The ubiquity of overlapping communities signifies the importance of finding overlapping communities in graphs.

Communities can be defined in many ways. In general, a community refers to a group of vertices that are densely connected to each other and sparsely connected to other vertices in the graph. The density of a community usually is measured by its average degree. In this paper, we define a community as a k-clique [1]. We will show the rational in Section 3. A k-clique is a complete graph of k vertices. Given a graph G, we can find all k-cliques in G. We say two k-cliques are adjacent if they share k − 1 vertices in G. We can then create a k-clique graph for G. In the k-clique graph, each vertex is a k-clique and an edge between two vertices means the two k-cliques they represent are adjacent in G. A community is defined as a connected component, called a k-clique component, in the k-clique graph. Clearly, the larger the k value, the denser the communities. By tuning k, we obtain communities of different density. Next, we use a toy social network to illustrate overlapping communities.

EXAMPLE 1 (OVERLAPPING COMMUNITIES). Consider the toy social graph shown in Figure 1. Suppose we need to find the communities containing Bob. When k = 3, we get two overlapping communities: abcde and bfghi (for simplicity, we use the first character of a person’s name to represent the person). When k = 4, we find two communities: abcde and bfghi, but bfghi is denser than abcde because Issac, who has fewer connections to the community, is excluded.

1.2 OCD vs OCS
There are two related problems: overlapping community detection (OCD), which finds overlapping communities in the entire network, and overlapping community search (OCS), which finds overlapping communities that a specific vertex belongs to.

The difference between OCD and OCS is clear. For a given graph, OCD takes k as input, and finds overlapping communities in the entire graph, using a batch process. As a result, all detected communities are of the density that is not less than the same threshold. On the other hand, OCS takes a vertex and a specific k as input, and finds overlapping communities containing the vertex, using an online process.

One common operation in community analysis is to retrieve the communities containing a vertex. To support online query answer-
ing by OCD, we need to detect and materialize all communities in an offline pre-computation stage and then index the communities. In contrast, with OCS, we just need to specify the vertex of interest in the query. Using OCD to support this common operation is less desirable for large graphs due to the following reasons.

- First, it is costly and time consuming to find communities for the entire graph by OCD. OCD in general is NP-hard, and many real life graphs are quite large, with millions or even billions of nodes. For example, the friendship network of Facebook [2] contains over 800 million nodes and 100 billion links. As a result, OCD is computationally prohibitive on real large graphs. In contrast, with OCS, we just need to find communities within the local neighborhoods of the vertex. In the experimental section, we will show that in most cases, our OCS solution only needs several milliseconds to find answers, independent of the entire graph size.

- Second, OCD uses a global criterion for community detection for all vertices in a network. The semantics of the discovered community in general heavily depends on the parameters. Example 1 has shown that for the same vertex, a different $k$ leads to communities of different density and different semantics. Results on real networks (the details are presented in the experiment section) further shows that under a given $k$, some vertices may have no corresponding valid communities. In other words, a fixed $k$ is not appropriate for all vertices. OCD does not support using the most appropriate $k$ for each individual vertex.

- Third, it is difficult for OCD to support dynamically evolving graphs. Graphs in real life are always evolving over time. OCD can only be used for offline analysis. Generally, we cannot afford to run OCD very frequently. As a result, a solution of OCD usually loses its freshness and effectiveness after a short period of time. In contrast, with OCS, we are free to issue queries for any vertex, which will not significantly hinder system performance.

In summary, OCD plus indexing suffers when the graph is dynamically evolving or users want to tune the density parameters for a better result. Because it is costly to rerun OCD and rebuild the index. Unfortunately, most real networks, such as Facebook friendship networks, are evolving and users in these networks usually expect to freely explore his or her neighborhood community. This motivates us to study online OCS.

### 1.3 Challenges

To understand the major challenges of OCS, we compare OCS with a related and well-studied problem: Community Search (CS). Given a vertex (or a set of vertices), community search finds the community (instead of a set of overlapping communities) that contains the vertex [3]. In the following analysis, we show that CS and OCS are fundamentally different, and methods of CS cannot be directly applied to solve the OCS problem. The challenges of OCS are summarized as the follows:

- Our first challenge lies in modeling the overlapping community search. CS and OCS have different semantics. For OCS, each community has a “functioning model”. For example, a person may be part of a “weekend hiking” community and a “software engineer community” at the same time. If we mix the two communities, then we incur a loss of semantics. Typical CS approaches cannot tell the difference. For example, one approach finds the subgraph that has the largest minimal degree as the community [3]. Clearly, if two subgraphs satisfy the criterion, then their union also satisfies the criterion. This shows that CS is not a good model for overlapping community. In comparison, OCS is more semantically aware in the sense that its goal is more about finding a functioning module that can better reveal the role of a vertex. Hence, we need to define the community more carefully and precisely to reflect this requirement. Specifically, we need to prescribe when two overlapping groups can be merged into one community, and when they are two separate communities.

- Our second challenge is that OCS is computationally harder than CS. We show that OCS in general is #P-Complete. Intuitively, in OCS, all communities need to be enumerated. In some extreme cases, when there are an exponential number of valid communities, such enumeration is quite costly. Even if there is only a small number of valid communities, finding them among an exponential number of candidates is still a great challenge. In contrast, in most CS [3], for a given query vertex, there is at most one community to detect.

- Our third challenge is to adopt approximate approaches so that we can scale to large graphs with millions of nodes. Due to its computation intractability, it is difficult to scale up to million-node real life graphs. Finding the approximate results without compromising too much the quality of the solution (particularly the semantics of the community) is a challenge.

### 1.4 Contributions

In this paper, we propose a novel model for OCS, the theory based on the model, and the algorithms that solve OCS. More specifically, we make the following contributions:

- We introduce a model for OCS that satisfies not only the traditional community semantics, but also the semantics of overlapping. Furthermore, the two types of semantics are consistent in our model.

- We propose several algorithms to solve OCS. These algorithms produce meaningful results, and can handle million-node graphs.

- We offer the theoretical guidelines for parameter selection. The guidelines take into consideration the structural properties of real networks, and ensure the discovery of meaningful overlapping communities in real life complex networks.

- We conduct extensive experiments on real networks to justify our algorithms and theories. We also show many applications of OCS model and algorithms.

## 1.5 Paper Organization

The rest of the paper is organized as follows. Section 2 discusses related work. Section 3 focuses on the model and the problem of overlapping community search. Section 4 describes a naive algorithm and an improved algorithm. We further propose a more efficient approximate algorithm. Experimental results in Section 5 show the performance and effectiveness of both the exact and the approximate algorithm. We show some potential applications for OCS model in Section 6. We conclude in Section 7.

## 2. RELATED WORKS

Our work is closely related to work on several topics, including overlapping community detection, dense subgraph discovery, and clique finding. We describe their similarity and differences in this section.

**Overlapping community detection.** Palla et al. [1] first noticed that most real networks have overlapping community structures and proposed a percolation based method to detect overlapping communities in real networks. They also defined a community as a $k$-clique component. But they did not generalize it to allow relaxation. They developed a software called CFinder [4] based
on the method to detect overlapping communities on biology networks. Their work was further extended with the EAGLE [5] algorithm that merges two cliques into one community if their Jaccard similarity score is large enough. Gregory et al. [6] transformed the problem of overlapping community detection (OCD) on graph $G$ into an equivalent non-overlapping community detection problem on another graph $G'$, which is produced by duplicating certain vertices or edges in $G$. They used betweenness as the heuristic to select the vertex or edge to be duplicated. They further improved its performance by using local betweenness in a way that can be quickly computed to approximate the betweenness [7]. Ahn et al. [8] reinvented a community as groups of links instead of vertices so that they can always produce a hierarchical dendrogram on vertex-overlapped networks. Baumes et al. [9] proposed a greedy solution to grow overlapping communities. The algorithm first ranks all nodes according to some criterion, such as PageRank. Then, each node is added in turn into existing communities whose first ranks all nodes according to some criterion, such as PageRank. 

Then, each node is added in turn into existing communities whose first ranks all nodes according to some criterion, such as PageRank. These vertices, called hubs, usually bridge different communities. So they proposed a linear algorithm, called SCAN, to find hubs and communities. Recently some efficient overlapping community detection algorithms were proposed, including SLPA [11], GCE [12] and OSLOM [13]. SLPA simulates the human communication behavior and realizes it into a speaker-listener label propagation algorithm. GCE identifies distinct cliques as seeds and expands these seeds by greedily optimizing a local fitness function. OSLOM finds overlapping communities by local optimization of a fitness function expressing the statistical significance of clusters with respect to random fluctuations. Compared to OCD, OSCs are more of a light-weight problem, and it is for online query answering. By definition, OCD inherently implies intractable computation complexity. For example, CFinder needs to enumerate all $k$-cliques in a graph and then combine them in terms of overlap between cliques. Both clique enumeration and pairwise computation of clique similarity are costly. In contrast, in OSCs, most resulting communities tend to occur around the local neighborhoods of the query vertex, allowing finding the answers by only exploring a part of the graph instead of processing the entire graph.

**Dense subgraph discovery.** In many graphs whose dense subgraphs have clear boundaries and can be separated from each other, communities are dense subgraphs. In this sense, our work is closely related to dense subgraph discovery. The problem of finding the densest subgraph is NP-hard, where the density of a graph $G(V, E)$ is usually measured by $\frac{|E|}{V(V-1)/2}$. Doron et al. [14] gave an approximate algorithm for finding the dense $k$-vertex subgraphs of a given graph, with approximation ratio $O(n^{\delta})$ for some $\delta < \frac{1}{2}$. For practical use, Gibson et al. [15] used recursive shingling to find dense subgraphs on massive graphs. Their algorithms are scalable to web-scale graph with 50M hosts and 11B edges but without guarantees for the density of the result. To handle Internet-sized graph data in the form of stream, Bahman et al. [16] proposed a greedy solution for graph streams, which takes $O(\log n + n)$ passes for any $\epsilon \geq 0$ yielding approximation factor of $2(1 + \epsilon)$. Some other methods propose different definitions of dense subgraphs. For example, DLS graph is proposed in [17] and accompanied with a triangle-counting based solution. Although dense subgraph discovery is related to community discovery, they are essentially different from each other. A community is far beyond a dense subgraph. As a community, the density within the subgraph is expected to be significantly larger than the outside world of the subgraph. Furthermore, our problem is a typical search problem, in which a query vertex needs to be specified.

**Clique finding.** $k$-cliques underlie our community definition. There is already many $k$-clique related research. Most of it concerns clique finding, i.e., finding a maximal clique or finding cliques with a size constraint. In general, the decision version of clique finding is NP-Complete [18]. Derényi [19] gave the threshold of edge-linking probability for the emergence of a giant $k$-clique component in an Erdös Renyi(ER) [20] random graph. To solve the problem practically, local search [21] is widely employed in many efficient solutions [22, 23, 24]. Along this direction, reactive local search (RLS) [22], dynamic local search (DLS) [23] and an improved DLS [24] have been proposed in turn. For disk-resident massive graphs, Cheng et al. [25] proposed an external-memory algorithm ExtMCE to reduce memory usage. They built a summary of small-sized graph, called H*-graph, to precisely encode the neighborhood information in the original graph. We only use $k$- clique to define community. In our problems, we not only need to find cliques around the query vertex, we also need to take care of clique adjacency. The second issue is not addressed in these clique-related works.

### 3. OVERLAPPING COMMUNITY

In the following, we first define the model as well as the problem (Section 3.1). Then, we justify the model by highlighting some of its important properties (Section 3.2). Finally, we discuss how to tune the parameters in the model to produce meaningful results (Section 3.3).

#### 3.1 Model and Problem Definition

We start with the definition of OSC problem. Then, we generalize it into the $(\alpha, \gamma, \gamma)$-OCS problem.

**3.1.1 OCS**

Network communities are usually modeled by $k$-cliques [1]. Given an original network $G$, we derive a $k$-clique graph from $G$. In the $k$-clique graph, each node represents a $k$-clique in $G$, and an edge between two nodes means that the $k$-cliques the two nodes represent are adjacent in $G$, where adjacency is defined as follows:

**DEFINITION 1 (CLIQUE ADJACENCY).** Two $k$-cliques are adjacent if they share $k - 1$ vertices.

We define OSCs based on the concept of $k$-clique components.

**DEFINITION 2 ($k$-CLIQUE COMPONENT).** Let $C$ be a connected component in the $k$-clique graph. A $k$-clique component is the union of all $k$-cliques represented by nodes in $C$.

Intuitively, given a vertex $v_0$, there may be multiple communities that contain $v_0$. All these communities naturally constitute the overlapping communities that contain $v_0$. Thus, the problem is finding all such communities.

**PROBLEM 1 (OVERLAPPING COMMUNITY SEARCH (OCS)).**

Given a graph $G(V, E)$, a query vertex $v_0$ and a positive integer $k$, we need to find all $k$-clique components containing $v_0$.

**Example 1 (OCS).** Consider the graph shown in Figure 1. Assume we want to find overlapping communities that contain node $b$ (i.e., $v_0 = b$). Let $k = 4$. We have three $k$-cliques containing $b$: $C_1 = abed, C_2 = abce$ and $C_3 = bfgh$. The clique graph is

![Clique graphs are sensitive to the parameters of OCS.](image)

(a) The clique graph for $(3, 1)$-OCS
(b) The clique graph for $(4, 8)$-OCS

Figure 2: Clique graphs are sensitive to the parameters of OCS.
shown in Figure 2(a). Hence, $C_1 \cup C_2$ and $C_3$ are returned as two overlapping communities containing the query vertex.

3.1.2 ($\alpha$, $\gamma$)-OCS

The definition of overlapping community given above is too restrictive: First, every pair of nodes in each $k$-clique by definition must be connected. Second, two $k$-cliques are considered adjacent iff they share as many as $k - 1$ nodes. In this section, we define a more general problem, $(\alpha$, $\gamma)$-OCS, by relaxing the two constraints.

First, we relax the adjacency requirement. In $(\alpha$, $\gamma)$-OCS, two $k$-cliques are considered adjacent if they share $\alpha$ nodes. This leads to the definition of $\alpha$-adjacency.

**Definition 3** ($\alpha$-adjacency). Two subgraphs $G_1$ and $G_2$ of graph $G(V, E)$ are $\alpha$-adjacent if they share at least $\alpha$ vertices.

Second, we relax the requirement of cliqueness. A $k$-clique is the densest graph among all $k$-node graphs. We generalize it to $\gamma$-quasi-clique by exposing parameter $\gamma$ to tune the density of the subgraph.

**Definition 4** ($\gamma$-quasi-$k$-clique [26]). A $\gamma$-quasi-$k$-clique of graph $G$ is a $k$-node subgraph of $G$ with at least $\lceil \frac{\gamma |V|}{k} \rceil$ edges (where $0 \leq \gamma \leq 1$).

We now give a generalized version of overlapping community search: $(\alpha$, $\gamma)$-OCS. Similar to the basic version, we also intend to find all overlapping communities containing a query vertex. But each community here is a $\gamma$-quasi-$k$-clique component and the clique adjacency is defined by $\alpha$ adjacency. Clearly, Problem 1 is simply $(k - 1, 1)$-OCS.

**Problem 2** ($(\alpha$, $\gamma$)-OCS). For a graph $G$, a query vertex $v_0$ and a positive integer $k$, the $(\alpha$, $\gamma$)-OCS problem finds all $\gamma$-quasi-$k$-clique components containing $v_0$.

**Example 3** ($(\alpha$, $\gamma$)-OCS). To continue with the previous example, consider (4,0.8)-OCS and $k = 5$ and $v_0 = \text{abcde}$. bgf is a two resulting communities. This can be directly obtained from the clique graph shown in Figure 2(b).

3.1.3 Complexity

We next analyze the complexity of overlapping community search.

**Theorem 1.** The $(\alpha$, $\gamma)$-OCS problem, as well as its special case $(k - 1, 1)$-OCS, are #P-Complete, if $k$ is part of the input.

**Proof.** We show that $(k - 1, 1)$-OCS is #P-Complete. Then, as a more general problem, $(\alpha$, $\gamma)$-OCS is at least #P-Complete. Given a query vertex $v_0$, the decision version of $(k - 1, 1)$-OCS is to decide whether there are any $k$-cliques that contain $v_0$. Since $k$ is not a constant, this decision problem is NP-complete, which can be reduced from the $k$-clique problem.

### 3.2 The Rationale

Is the model proposed in Section 3.1 meaningful? In this section, we justify it from the following three aspects: community density, overlapping-awareness and consistency.

#### 3.2.1 Community Density

To qualify as a community, a subgraph obviously needs to have enough density. In $(\alpha$, $\gamma)$-OCS, the parameters $\alpha$ and $\gamma$ together control the closeness among the members of a community. In this subsection, we establish a lower bound (Theorem 2) of the average degree for communities found by $(\alpha$, $\gamma)$-OCS. Theorem 2 is based on Lemma 1. As an example of Theorem 2, the average degree of a community of $(k - 1, 1)$-OCS is at least $k - 1$.

**Lemma 1.** For a $\gamma$-quasi-$k$-clique, the number of edges incident to an arbitrary set of $x$ vertices is at least $\max \{0, \gamma \binom{x}{2} - \binom{k - 1}{2} \}$.

**Proof.** The total number of edges of this clique is at least $\gamma \binom{x}{2}$.

The subgraph induced by the rest $\text{k} - x$ vertices has at most $\binom{x}{2}$ edges, which are not incident to the selected vertices. Therefore, the lemma holds.

**Theorem 2.** (Lower bound on average degree). The average degree of each community found by $(\alpha$, $\gamma)$-OCS has a lower bound

$$2 \max \{0, \min \{f(1), f(\alpha)\} \}$$

where $f(x) = \frac{(\gamma \binom{x}{2} - \binom{k - 1}{2})}{2}$. 

**Proof.** First, we show that for $x \in \{1, 2, ..., \alpha\}$, it holds that $f(x) \geq \min \{f(1), f(\alpha)\}$. We have $f'(x) = -\frac{1}{2} + \frac{(1 - 2\alpha)(k - 1)}{2x^2}$, which satisfies the following inequalities:

$$f'(x) \geq 0, \quad 0 < x \leq \sqrt{(1 - \gamma)k(k - 1)}$$

Thus, $f(x)$ takes its minimum value either at $x = 1$ or $x = \alpha$, which means $f(x) \geq \min \{f(1), f(\alpha)\}$.

Now, suppose $C$ is a community found by $(\alpha$, $\gamma)$-OCS, and $C$ consists of $t$ different cliques $C_1, C_2, ..., C_t$. Let $V_i = \bigcup_{1 \leq j \leq t} C_j$. Let $v_i = [V_i - V_{i - 1}]$. We have $|C| = \sum_{0 \leq i < t} v_i$. Let $e_i$ be the number of edges in $E[V_i] - E[V_{i - 1}]$, where $E[V_i]$ is the edges induced by $V_i$. The average degree of $C$ is $\frac{2|E(C)|}{|V(C)|}$, which equals

$$\frac{2|E(C)|}{|V(C)|} = \frac{2}{|C|} \sum_{0 \leq i < t} e_i \geq \frac{2}{|C|} \sum_{0 \leq i < t} v_i \times \frac{\gamma}{2}.$$

Using the minimal value of $\frac{\gamma}{2}$ to replace each $\frac{\gamma}{2}$, we get

$$\frac{|E(C)|}{|V(C)|} \geq \frac{2}{|C|} \sum_{0 \leq i < t} v_i \min \{\frac{e_i}{v_i} \} = 2 \min \left\{ \frac{\sum_{0 \leq i < t} e_i}{\sum_{0 \leq i < t} v_i} \right\}.$$

According to Lemma 1 and the fact that $f(x) \geq \min \{f(1), f(\alpha)\}$, we find that the average degree of a community found by $(\alpha$, $\gamma)$-OCS has a lower bound $2 \max \{0, \min \{f(1), f(\alpha)\} \}$.

#### 3.2.2 Overlapping Awareness

The model should be able to reveal the overlapping relationships of multiple communities. Note that this is not a trivial task. When two subgraphs overlap, it is hard to tell whether they are two components of a single community or two overlapping communities. In OCS, given two subgraphs $H_1, H_2$ that contain the query vertex, we have two choices: return $H_1 \cup H_2$ as a single community or return $H_1, H_2$ as two overlapping communities.

Previous models of communities were overlapping-unaware, and even overlapping-unfriendly. A widely adopted model [3] considers a subgraph whose minimal degree is larger than a given threshold as a valid community. Thus, the union of two communities is also a valid community. Hence, two communities can always be merged, even if they have very little overlap.

In $(\alpha$, $\gamma)$-OCS, we use $\alpha$ to decide whether two components have enough overlapping to be considered and merged as a single community. Specifically, if two $\gamma$-quasi-$k$-cliques share fewer than $\alpha$ vertices and they are not reachable in any clique component, they are considered as two overlapping communities instead of a single community.

#### 3.2.3 Consistency

Community search must be consistent, that is, if a community $C$ is considered a community for a query vertex $v_0$, then if we use any other vertex in $C$ as a query vertex, we should also obtain $C$ as its community. We establish the consistency of $(\alpha$, $\gamma)$-OCS in Theorem 3, which can be easily derived according to the fact that clique adjacency is an equivalence relationship.

**Theorem 3** (Consistency). In $(\alpha$, $\gamma)$-OCS, if $C$ is a community that contains query vertex $v_0$, then for any other vertex $v \in C$ as the query vertex, $C$ is also returned as its community.
3.3 Parameter Selection

The model for overlapping community contains 3 parameters: \( k, \alpha, \gamma \). The selection of these parameters is crucial to producing meaningful results. In this section, we will give the theoretic guideline for selecting appropriate parameters in real networks.

3.3.1 Selection of \( \gamma \) and \( k \)

Parameter \( \gamma \) controls the density of the community. The selection of \( k \) and \( \gamma \) is related, in the sense that the larger the \( k \), the more difficult it is to find communities with the high density. Hence, in order to generate meaningful results, \( \gamma \) should vary with \( k \). Next, we will reveal the relationship between \( \gamma \) and \( k \). We first give a simple relationship established to avoid a trivial case. Then, we establish a more complicated relationship between them on ER random graphs.

A simple relationship. In general, as a meaningful community, the quasi clique needs to be connected. For \( k \) vertices, even when there are \( \frac{k(k-1)}{2} - (k-1) \) edges among them, it is still possible that they are disconnected. It happens when the \( k \)-vertex graph consists of an isolated vertex and a \( (k-1) \)-vertex clique. To avoid the disconnected case, \( \gamma \) should be big enough. Hence, we need to ensure that:

\[
\left\lfloor \frac{k(k-1)}{2} \right\rfloor > \frac{k(k-1)}{2} - k + 1,
\]

By transformation, we have

\[
\gamma \geq \frac{k-2}{k} + \frac{2}{k(k-1)}
\]

(1)

Relationship on ER random graphs. Given a graph with \( N \) nodes and \( M \) edges, the density of the graph is given by \( \frac{2M}{N(N-1)} \). Next, we use the ER random graph model [20] as an example to reveal the relationship between \( \gamma \) and \( k \). The ER random graph is used to simulate real graphs, such as road networks. Consider an ER random graph \( G \in \mathcal{G}(n, p) \) that has \( n \) vertices and each pair of vertices is linked with probability \( p \), independent of other vertices and edges. Suppose we randomly pick \( k \) vertices from \( G \). Let \( X \) be the random variable that represents the number of edges among the selected vertices. Then, the probability that \( X = i \) is

\[
Pr(X = i) = \binom{m}{i} p^i (1-p)^{m-i}
\]

where \( m = k(k-1)/2 \). Let \( Y \) be the random variable that represents the density of the subgraph induced by the randomly selected \( k \) vertices. Clearly, we have \( Y = X/m \). Then, the probability that \( k \) selected vertices have density \( \gamma \) is a function of \( k \) and \( \gamma \), that is:

\[
Pr(Y = \gamma) = \binom{\left\lfloor \frac{k(k-1)}{2} \right\rfloor}{\left\lfloor \frac{k(k-1)}{2} \right\rfloor} p^{\left\lfloor \frac{k(k-1)}{2} \right\rfloor} (1-p)^{\left\lfloor \frac{k(k-1)}{2} \right\rfloor - \left\lfloor \frac{k(k-1)}{2} \right\rfloor}
\]

(3)

We give the simulation of \( Pr(Y = \gamma) \) in Figure 3(a) for \( p = 0.3 \). We can clearly see that when \( \gamma \) is fixed, \( Pr(Y = \gamma) \) is a monotonically decreasing function of \( k \). This suggests that we can find communities for small \( k \) with high probability. We justify this in the experimental section by showing that for most real networks communities exist mostly for 4 \( \leq k \leq 10 \).

When \( k \) is fixed, \( Pr(Y = \gamma) \) reaches its maximum when \( \gamma \) is close to the average density. To show this more clearly, we give \( Pr(Y = \gamma) \) as a function of \( \gamma \) for two fixed \( k \) in Figure 3(b). The simulation results imply that there are many subgraphs of average density in the networks. In general, a meaningful community should have a much larger density than the average density. Hence, we should select a large enough density to disfavor those trivial communities with average density.

More formally, once \( k \) is given, we may use a confidence value such as \( c = 0.05 \) to construct an inequality \( Pr(Y \geq \gamma) \leq c \) which is equivalent to

\[
\sum_{i\leq j\leq m} \binom{m}{i} p^i (1-p)^{m-i} \leq c
\]

, since \( Pr(X \geq i) = Pr(Y \geq \gamma) \) when \( i = \gamma m \). By solving above inequation, we find the appropriate \( i \) value and corresponding \( \gamma \). By tuning \( c \), we can control \( \gamma \). A smaller \( c \) generally leads to larger \( \gamma \).

3.3.2 Selection of \( \alpha \)

Parameter \( \alpha \), as we mentioned before, controls how strictly we consider two communities to be one community or components of the same community. Our major goal in selecting a meaningful \( \alpha \) is to avoid two trivial cases. The first trivial case occurs when the density of a \( \gamma \)-quasi-\( k \)-clique is contributed mainly by its \( \alpha \)-size subset. As a result, any other \( k - \alpha \) vertices outside of the clique combined with the subset will be a valid adjacent clique. This trivial case obviously leads to meaningless communities. To avoid this case, we need \( \binom{\alpha}{2} < \gamma \binom{k}{2} \). By simple transformation, we have

\[
\alpha \leq \left\lceil \sqrt{1/4 + k(k-1)\gamma - 1/2} \right\rceil
\]

(4)

The second trivial case occurs when two \( \gamma \)-quasi-\( k \)-cliques intersect at a \( \alpha \)-size subset that is quite sparse. In real networks, if the common part of two subgraphs is too sparse, the two subgraphs can rarely be regarded as a single closely-connected community. If the common part of two subgraphs is too sparse, the two subgraphs can rarely be regarded as a single closely-connected community. This case happens if all edges are located among the rest \( k - \alpha \) vertices. To avoid this, we set \( \gamma \binom{\alpha}{2} > \binom{k-\alpha}{2} \) so that the common part has at least one shared edge. By transformation, we have

\[
\alpha \geq \frac{k+1/2 - \sqrt{k^2 - \gamma k + 1/4}}{2}
\]

(5)

Combining the two cases, given \( k \) and \( \gamma \), we chose \( \alpha \) to be an integer between the lower and upper bounds. As an example, when \( \gamma = 1 \), we have \( 1 \leq \alpha \leq \left\lfloor \sqrt{1/4 + k(k-1) - 1/2} \right\rfloor \). Furthermore, when \( k = 4 \), we have \( 1 \leq \alpha \leq 3 \). Hence, usually we use \( \alpha = 3 \).

3.3.3 Practical selection of parameters

Based on the above analysis, we show that \( \alpha = k - 1 \), \( \gamma = 1 \) (or values close to 1) are typical settings of \( (\alpha, \gamma) \)-OCS. By simple transformation of Eq. 1, we have

\[
\gamma \geq 1 - \frac{2k-2}{k-1} = f(k)
\]

. When \( k > 3 \), \( f(k) \) converges to 1. It can be observed from the simulation of \( f(k) \) in Figure 4. Hence, \( \gamma = 1 \) (or values close to 1).
1) is a typical setting. Next, we show that \( \alpha = k - 1 \) is also a typical setting because only \( \alpha = k - 1 \) lies between the lower and upper bound of \( \alpha \) for any \( k \) and \( \frac{k-2}{k} + \frac{2}{k(k-1)} \leq \gamma \leq 1 \) (shown in Theorem 4).

![Figure 4: Simulation of \( f(k) \).](image)

**Theorem 4.** Let \( \alpha = k - \Delta \) and \( \Delta \) be a positive integer less than \( k \). Only when \( \Delta = 1 \), \( \alpha \) satisfies Eq. 4 and Eq. 5 for any \( k \geq 2 \) and \( \frac{k-2}{k} + \frac{2}{k(k-1)} \leq \gamma \leq 1 \).

**Proof.** First, consider \( k = 2 \). Since \( 0 < \alpha = k - \Delta < k \), we have \( 2 - \Delta > 0 \). Hence, only \( \Delta = 1 \) satisfies this inequality. Next, for any \( k \) and \( \frac{k-2}{k} + \frac{2}{k(k-1)} \leq \gamma \leq 1 \), we have

\[
\left[1 + \frac{1}{k} - \frac{4}{k(k-1)} \right] \geq \left[ \sqrt{\frac{9}{4}} + (k - 1)(k - 2) - 1/2 \right],
\]

which is no less than \( k - 1 \); and

\[
\left[1 + 2 - \sqrt{\gamma(k^2 - \gamma k + 1/4)} \right] \leq \left[ k - 1/2 - \sqrt{(k-1)(k-2) + 9/4} \right]
\]

which is no longer than \( k - 1 \). Thus only \( \alpha = k - 1 \), i.e., \( \Delta = 1 \), satisfies the constraints. \( \square \)

### 4. Algorithms

In this section, we will first propose two exact algorithms to solve OCS. The first is a straightforward solution. The second is a more efficient exact solution. To further improve the performance, we also propose a more efficient approximate algorithm.

#### 4.1 Exact algorithm

##### 4.1.1 Naive Algorithm

We give a naive algorithm according to the definition of OCS. The direct solution for OCS consists of three major steps, which are shown in Algorithm 1. In the first step, we find the \( \gamma \)-quasi-\( k \)-clique that contains \( v_0 \). Then for each newly discovered vertex, we find all \( \gamma \)-quasi-\( k \)-cliques that contain the vertex. We repeat the above finding procedure until no new vertex can be discovered. In the second step, we calculate the clique adjacency for all cliques found in the first step. Finally, we return all clique components as the resulting communities.

**Algorithm 1 Naive OCS**

**Input:** \( G(V, E), v_0, \alpha, \gamma, k \);

**Output:** The overlapping communities containing \( v_0 \);

1: \( V_C \leftarrow \{v_0\} \);

2: for all unvisited vertex \( v \in V_C \) do

3: \( \text{find clique}(v) \); \( \triangleright \) find all cliques that contain \( v \);

4: Add all vertices of these cliques into \( V_C \);

5: end for

6: Calculate clique component

7: Calculate the adjacency matrix of candidate cliques;

8: Stage 3. Return all clique components as communities

#### 4.1.2 New algorithm framework

The naive algorithm is not smart yet. As illustrated in Example 4, many cliques enumerated in the naive algorithm will not belong to a valid community. Hence, enumerating these invalid cliques is wasteful. This can be avoided if we check the adjacency when a clique is enumerated. Following this idea, we propose an improved algorithm, shown in Algorithm 2. The algorithm runs iteratively. In each iteration, we find an unvisited clique containing the query vertex by next clique(). If such a clique exists, we find the clique component that the clique belongs to by expand().

**Algorithm 2 Improved OCS**

**Input:** \( G(V, E), v_0, \alpha, \gamma, k \);

**Output:** The overlapping communities containing \( v_0 \);

1: \( R \leftarrow \emptyset \);

2: while \( C \leftarrow \text{next clique}(v_0), C \neq \emptyset \) do

3: \( C \leftarrow \text{expand}(C) \); \( \triangleright \) find the clique component of \( C \);

4: \( R \leftarrow R \cup \{C\} \);

5: end while

6: Return \( R \);

Note that for any two adjacent cliques, their clique components are identical. To avoid redundant enumeration, we record the visit status of each clique in both next clique() and expand(), and only enumerate unvisited cliques. We illustrate this in Example 5. The new algorithm framework also allows us to further optimize it. We will present such optimizations in the following sections.

**Example 5 (Improved Algorithm).** Continue with the previous example but change \( v_0 = b \). The next clique() may first return \( b, f, g, h \). By calling expand() on this clique, we find the first community \( \{b, f, g, h\} \). The next call of next clique() may return abc. By calling expand() on this clique, we find abcde. Thus, \( \{a, b, c, d, e\} \) is the second community. Then any call of next clique() will return nothing and the algorithm will terminate since all cliques have been visited either in next clique() or expand().

##### 4.1.3 Implementation of next clique()

next clique() is responsible for enumerating each unvisited \( \gamma \)-quasi-\( k \)-clique containing vertex \( v_0 \). Finding all cliques is clearly \#P-complete. A naive solution needs to exhaustively enumerate all \( k \)-subsets with \( k \) containing \( v_0 \), and check whether it is a valid clique. The solution costs \( O(V^k k^2) \) time. Because there are \( O(V^k) \) \( k \)-size subsets and checking whether the subset is a valid clique needs \( O(k^2) \) time.

Clique enumeration is a well-known computationally hard problem. Even allowing approximation, clique finding is hard to approximate. Hence, brute-force enumeration seems to be inevitable to produce exact results. The major procedure is a depth-first search with backtracking. The procedure is shown in Algorithm 3. The search starts from \( U = \{v_0\} \). Then, the procedure iteratively adds a new vertex into \( U \) until a new valid clique is found (line 5, 6). The procedure will stop until all \( \gamma \)-quasi-\( k \)-cliques containing the query vertex are found.

The backtracking search can be improved from two aspects. First, we just need to select a new vertex from the neighbors of the current vertex (line 13). This is because as a valid parameter, \( \gamma \) needs to satisfy Eq. 1, which ensures the result is a connected subgraph. Second, we speedup the search by pruning the impossible enumeration (line 11). Let \( |E(U)| \) be the number of edges in the subgraph induced by \( U \). Thus the maximal number of edges that the resulting clique has is \( |E(U)| + (k - |U|)(|U| + |k - |U| - 1|) \delta \), which equals \( |E(U)| + \frac{(k-|U|)(|U|+|k-|U|-1|)}{2} \). When the maximal number of edges is less than \( \gamma \frac{k(k-1)}{2} \), that is

\[
g(U) = |E(U)| + \frac{(k - |U|)(k + |U| - 1)}{2} < \gamma \frac{k(k-1)}{2}
\]

(6)
The same clique from different paths. For example, for the clique

\[ |U| = k \]

we can certainly prune the current \( U \). The left side of the inequality is a function of \( U \). We use \( g(U) \) to denote it.

**Algorithm 3 DFS procedure of next_clique(0)**

1: \( U \leftarrow \{v_0\} \);
2: \( DFS(U, v_0) \);
3: procedure DFS(U, a)
4:   if \( |U| = k \) then
5:     \( \text{if } U \subset C \text{ and } U \text{ is unvisited } \) then
6:       \( U \leftarrow v \);
7:       \( \text{Return } U; \)
8:     else
9:       \( \text{end if } \)
10:    \( \text{end if } \)
11:   \( \text{if } g(U) < \gamma^{k(k-1)} \) then Return;
12: \( \text{end if } \)
13: for all \( (u, v) \in E, v \notin U \) do
14: \( DFS(U \cup \{v\}, v) \);
15: \( \text{end for } \)
16: \( \text{end procedure } \)

### 4.1.4 Implementation of \( expand() \)

Function \( expand(C) \) is used for finding the clique component containing \( C \). By any graph traverse on the clique graph, we can certainly find the connected component. But the traverse orders are influential on the performance. Here, we use DFS order.

The procedure of \( expand(C) \) is shown in Algorithm 4. The major framework is a DFS traverse on the clique graph. In the algorithm, \( A \) is a global variable storing the currently found vertices in the clique component. The key operation in the traverse is finding a subset \( S_2 \) from \( C \)'s neighborhoods to replace one of \( C \)'s subset \( S_1 \) so that the new combination \( C' \) is a valid clique. To keep the size of the clique as \( k \), we enforce \( |S_1| = |S_2| \) (line 8). We can also use Eq. 6 to terminate the search that will not lead to a valid clique (line 6).

The function \( Candidate(C - S_1) \) is used for finding a subset \( S_2 \) with the same size as \( S_1 \) from the local neighborhoods of \( C - S_1 \). Note that we do not need to enumerate \( S_2 \) from the entire \( V \). Because under the constraint of Eq. 1, the induced graph of \( (C - S_1) \cup S_2 \) is always connected. Hence, in the worst case, we just need to explore the \( |S_1| \)-hop-neighborhood of \( C - S_1 \), which contains all vertices that are at most \( |S_1| \) hops away from any vertex in \( C - S_1 \). When \( k \) is small, the exploration of \( |S_1| \)-hop-neighborhood will be effective.

**Algorithm 4 expand(0)**

Input: A \( \gamma \)-quasi-\( k \)-clique \( C \);
Output: The clique component of \( C \);
1: \( A \leftarrow C \);
2: \( \text{Expand_clique}(C) \);
3: return \( A \);
4: procedure Expand_clique(C)
5: for all \( S_1 \subset C, |S_1| \leq k - \alpha \) do
6: \( \text{if } C - S_1 < \gamma^{k(k-1)} \) then Continue
7: \( \text{end if } \)
8: for all \( S_2 \in Candidate(C - S_1), |S_2| = |S_2| \) do
9: \( C' \leftarrow (C - S_1) \cup S_2 \); \( \) \( \text{if } C' \text{ is unvisited } \) \( \text{and } C' \text{ is a } \gamma \)-quasi-\( k \)-clique \) then
10: \( A \leftarrow A \cup S_2 \);
11: \( \text{Expand_clique}(C') \); \( \) \( \text{end if } \)
12: \( \text{end for } \)
13: \( \text{end for } \)
14: \( \text{end procedure } \)

### 4.1.5 Duplication detection

In both \( next_clique() \) and \( expand() \) functions, we may meet the same clique from different paths. For example, for the clique

\[ a - b - c \text{ or } c - a - b \]

we may meet it by \( a - b - c \) or \( c - a - b \). Hence, we need to tell whether a clique has ever been visited. For this purpose, we use a hash table to store visited cliques, allowing in constant time to query a visited clique. We use the following hash function:

\[ h(C) = \left( \sum_{v_i \in C} \text{id}(v_i) \times a^{\text{id}(v_i)} \right) \mod b \]

where \( a, b \) are two large primes and \( \text{id}(v_i) \) is the id of \( v_i \). Hashing each \( k \)-size clique takes \( O(k) \) time. By DFS, such complexity can be further reduced to \( O(1) \). In DFS, two successively visited cliques have only two different vertices. Thus, we can calculate the hash value in an incremental way. Suppose, \( C_{i+1} = (C_i - \{v\}) \cup \{u\} \), we have

\[ h(C_{i+1}) = (h(C_i) - (\text{id}(v) \times a^{\text{id}(v)}) + \text{id}(u) \times a^{\text{id}(u)}) \mod b \]

### 4.1.6 Optimization on \((k-1,1)-OCS\)

In the exact solutions for the general OCS problems, we can only detect duplication after the enumeration of a clique. For a \( k \)-size subset, we may enumerate it \( O((k-1)) \) times. Next we show that on \((k-1,1)-OCS\), we can avoid such redundant enumerations. We only discuss it for \( next_clique(v_0) \). \( expand() \) can be optimized with the same technique. When we search for a \( \gamma \)-quasi-clique containing \( v_0 \), we need to search along a path (line 13-15 in Algorithm 3) to ensure that we find a connected subgraph. However, on \((k-1,1)-OCS\), we are searching for a standard clique (a complete subgraph). It is order independent to find such cliques. That is, for these cliques we can use any DFS order to enumerate the same clique. As a result, on \((k-1,1)-OCS\), we can pose an arbitrary linear order on \( V \). Then, we search for a clique according to the linear order. That is only adding a vertex with a superior order into the partial solution. In this way, redundant enumerations of a \( k \)-clique can be avoided.

### 4.2 Approximate Algorithm

Next, we propose an efficient approximate algorithm for \((\alpha, \gamma)\)-OCS. The algorithm shares the same framework as Algorithm 2 with \( next_clique() \) and \( expand() \) further approximated.

#### 4.2.1 Approximation in next_clique()

The \( next_clique() \) function may enumerate an exponential number of \( \gamma \)-quasi-\( k \)-cliques. To reduce the enumeration space, we only enumerate an unvisited clique which contains at least one new vertex that does not belong to any communities already found. This heuristic can be achieved by a slight change of the DFS procedure (Algorithm 3). Let \( v_0, v_1, \ldots, v_k \) be the vertex sequence (staring from the root \( v_0 \)) in the DFS tree. We require that the second vertex \( v_1 \) is a new vertex. In this way, we ensure that the new clique contains at least a new vertex. Note that we enforce that the vertex sequence contains a new vertex as early as possible and \( v_1 \) is the earliest vertex added into the sequence. Hence, \( v_1 \) is required to be a new vertex.

#### 4.2.2 Approximation in expand()

We first study the traverse orders in \( expand() \) function. We show in Example 6 that to meet all vertices in a community, the best case only requires a linear number cliques to be visited, while in the worst case, an exponential number may be visited.

**Example 6 (Influence of Traverse Orders).** Consider a complete graph with 100 vertices and \( k = 4 \). For \((k-1,1)-OCS\) on any query vertex, the graph itself is the unique answer. To find the exact result, we need to visit \( (\binom{100}{2}) \) approximately millions of \( k \)-cliques. Alternatively, if we always select a clique that is adjacent to previously visited cliques and contains at least one unvisited vertex, after visiting 97 cliques, all vertices in the result are met.

The above example implies that we may find a community quite early before we exhaustively enumerate all \( \gamma \)-quasi-\( k \)-cliques in the
community. Let \( C \) be a community of \((\alpha, \gamma)\)-OCS on graph \( G \), in general we expect to find a shortest clique sequence \( C_1, C_2, \ldots, C_m \) such that each \( C_i \) is adjacent to at least one of its preceding cliques, and \( \cup_{1 \leq i \leq m} C_i = C \). However, in general, there exists an exponential number of possible sequences, it is hard to select the shortest sequence to visit.

Here, we will present a heuristic to find a short clique sequence. We still use DFS order. Consider \( \text{expand}(C_i) \). Let \( C_{i-1} \) be two successively visited cliques; we always select \( C_{i+1} \) such that

\[
C_{i+1} = \bigcup_{j < i} C_j \neq \emptyset
\]  

(9)

In other words, \( C_{i+1} \) is the clique containing at least one unvisited vertex. The traverse stops at \( C_m \), to which no other unvisited cliques satisfying Equation 9 are adjacent. In this way, the complete clique sequence \( C_1, \ldots, C_m \) is found. We return the union of them as the result of \( \text{expand}(C_i) \). This heuristic can be implemented by simply adding a condition, \( S_i - A \neq \emptyset \), in line 10 of Algorithm 4.

4.2.3 Analysis

Consider a \( k \)-clique component \( C \). Compared to the exact algorithm, which explores \( O(|C|) \) cliques, the approximate \( \text{expand}() \) algorithm only explores exactly \( O(|C|) \) cliques. Because the heuristic defined in Eq. 9 ensures finding a new vertex each time a clique is visited. \( O(|C|) \) cliques will be visited and many wasteful enumerations are avoided. In this way, we reduce the exponential complexity to linear complexity. However, this heuristic may produce inexact results, which are illustrated in Example 7. We will show in the experimental section that this heuristic works quite well on real graphs, producing accurate results.

**Example 7 (Bad case).** Consider the \((2, 1)\)-OCS on the graph shown in Figure 1 with \( k = 3 \) and \( v_0 = f \). The right community is \( bfgi \). If cliques \( bhf, bgh \) were first visited, the search will stop since no new clique can be visited. As a result, \( i \) will be missed. If we first visit \( bh \) and \( bfg \), \( gi \) is qualified to be visited and then we can find the right community.

Due to the NP-hardness to approximate of clique finding, it is hard to give a theoretic guarantee of the approximation quality. However, we find that the approximate communities obtained by the approximation of \( \text{next\_clique}() \) and \( \text{expand}() \) are finer than the exact ones. More formally, overlapping communities can be considered as a collection of subsets \( P = \{ C_1, C_2, \ldots, C_k \} \). Then, for each community \( C_i \) in the approximate result, \( \exists C' \) such that \( C_i \subseteq C' \) and \( C' \) is an exact community of \( v_0 \). It clearly holds since the two approximations are equivalent to strengthen the constraint of a subgraph as a community. Since the community is finer, its size is smaller than or equal to the exact one. We have mentioned that the approximate algorithm only explores \( O(|C|) \) cliques. Therefore, this property implies that the approximate algorithm is more efficient.

5. EXPERIMENTS

In this section, we present the experimental study and show the effectiveness and efficiency of our models and algorithms.

5.1 Experiment Setup

We ran all experiments on a PC with Intel Core2 at 2.13GHz, 4G memory running 64-bit Windows 7. We implemented all algorithms in C++. We compared the improved exact algorithm and the approximate algorithm. We also compared the OCS model and the OCD model.

We used four real networks to test our solution. The basic statistics of these networks are shown in Table 1. DBLP is a scientific collaboration network extracted from a recent snapshot of the DBLP dataset. Each vertex in the graph represents an author and each edge indicates a coauthoring relationship. Livejournal is the friendship network of Livejournal, a social networking and blogging site. Google represents the web graph that was released in 2002 by Google. In the graph, nodes represent web pages and directed edges represent hyperlinks between them. We ignore the direction of the edges. WordNet is a semantic network in which each vertex represents a specific sense in language and each edge represents a certain semantic relationship between senses. WordNet has been widely used in a variety of real applications, such as word sense disambiguation, automatic text classification and automatic text summarization.

**Figure 5**: Case study on DBLP. OCS can find communities with different research topics. By tuning \( k \), OCS can find communities with different closeness. This case study justifies the effectiveness of the OCS model.

5.2 Case Study

We justify the model we use by two case studies: the scientific collaboration network and WordNet.

5.2.1 Scientific collaboration network

We run both the improved exact algorithm and the approximate algorithm on the scientific collaboration network extracted from DBLP. The two algorithms produce almost the same results for most query vertices. Here, we use ‘Jiawei Han’ as the query vertex. Jiawei Han is a renowned computer scientist specializing in data mining and database. When \( k = 6 \), the exact result and the approximate result only differ in one community, where only two authors are missing in the approximate result. This justifies the effectiveness of the approximate algorithm. Hence, in the following evaluation, we will only use results produced by the approximate solution.

\[ ^2 \text{Available at http://dblp.uni-trier.de/xml/} \]

\[ ^3 \text{This and Google are available at http://snap.stanford.edu/data.} \]
We show in Figure 5(a) part of the overlapping communities that Jiawei Han resides in. It only takes 18ms for the approximate solution to get the results, which justifies the efficiency of the approximate solution. Next, we focus on the effectiveness of the approach. We find that our overlapping community search model is meaningful in many aspects.

- First, it successfully unveils multiple research interests of an author. For example, in Jiawei Han’s case, the community \( C_1 \) represents the topic of multimedia data mining and the community \( C_2 \) is about stream data mining. Each community has a clear boundary to be distinguished from others. We also can see that those authors with multiple research interests occur in different communities. For example, Jian Pei simultaneously occurs in \( C_2 \) and \( C_3 \), reflecting his research interest in both stream data mining and information network mining.

- Second, our model is flexible. By tuning parameter \( k \), our model can discover communities with different closeness. For example, when \( k = 9 \), we discover more closely-connected communities of ‘Jiawei Han’ as shown in Figure 5(b). From the result, we can see that community \( C_1 \) of \( k = 6 \) is broken up into two more-closely connected overlapping communities \( C_1, C_2 \) of \( k = 9 \). The two communities respectively reflect Jiawei’s more specific research interest. One is about multimedia data mining and the other is about association rule mining on large relational databases. When \( k \) becomes larger, many periphery members of communities who have relatively fewer collaborations with members in the community will be excluded from the community. For example, we find that 59 authors in the result when \( k = 6 \) are excluded from the result when \( k = 9 \).

5.2.2 WordNet

In WordNet, we use the word ’vessel’ as the query vertex and we set \( k = 3 \). The result in Figure 6 shows that ’vessel’ occurs in multiple communities, and each community represents a certain sense of ’vessel’. The community that contains ‘bowl’, ‘dish’ corresponds to tablewares. The community consisting of ‘barrel’ and ‘tube’ represents the meaning of bucket. Another two communities represent the meaning of ships. The difference between these two communities is that the one consisting of ‘ship’ and ‘galley’ is generally used for describing ships with large tonnage and the other community are usually used for describing small boats. This example shows that our OCS model is helpful for finding different senses of a word, which is a fundamental task in natural language understanding.

![Figure 6: Case study on Wordnet. OCS can find different senses of a word.](image)

### Table 1: Datasets, performance of OCD and OCS

<table>
<thead>
<tr>
<th>Dataset</th>
<th>#Vertices</th>
<th>#Edges</th>
<th>average degree</th>
<th>Performance of OCD and OCS</th>
</tr>
</thead>
<tbody>
<tr>
<td>WordNet</td>
<td>82670</td>
<td>153445</td>
<td>5.23</td>
<td>LA</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>51s</td>
</tr>
<tr>
<td>DBLP</td>
<td>560851</td>
<td>1816613</td>
<td>6.48</td>
<td>6183s</td>
</tr>
<tr>
<td>Google</td>
<td>916427</td>
<td>422051</td>
<td>9.43</td>
<td>22873s</td>
</tr>
<tr>
<td>Livejournal</td>
<td>4847571</td>
<td>42851237</td>
<td>17.7</td>
<td>&gt;= 24h</td>
</tr>
</tbody>
</table>

5.3 Performance

We first compare the performance of the exact algorithm and the approximate algorithm. Then, we compare the performance of OCS and OCD.

**OCS: exact vs approximate methods.** We tested the performance for both exact algorithms and approximate algorithms. For each \( k \), we randomly selected 100 valid query vertices (with degree at least \( k - 1 \)). We counted the average query answering time as well as the variance for these vertices under different \( k \). Note that the exact solution has exponential enumeration cost. Hence, when the running time exceeds 60s, we terminated the exact algorithm and recorded the entire query response time as 60s. Clearly, this favors exact solutions. However, approximate solution still shows significant advantage over exact solutions as the following results indicate.

The results on three real networks are given in Figure 7. We can see that in general approximate algorithm is more than two orders of magnitude faster than the exact one. Note that the performance priority of the approximation algorithm over the exact one is more striking than observed since we limit the running time of the exact algorithm to 60s. For Livejournal and Google, significant performance difference can be consistently observed over different \( k \). When \( k = 4 \), most valid query vertices have non-trivial result and the approximate algorithm shows a much clearer advantage over the exact algorithm. On DBLP, an increase in speed by three orders of magnitude is achieved for \( k = 4 \). For all the tested networks, the approximate solution’s performance is quite stable compared to the exact solution, especially on DBLP. Even for the biggest two datasets, Livejournal and Google, both with approximately a million nodes and tens of millions edges, the approximation algorithm returns results within less than 100ms for almost all \( k \). This sufficiently shows that the approximate solution can be used as online service for large real graphs.

![Figure 7: DBLP](image)

(a) DBLP

![Figure 7: Livejournal](image)

(b) Livejournal

![Figure 7: Google](image)

(c) Google

**OCS vs OCD.** For comparisons, we also give the running time of the state-of-the-art OCD algorithmss LA [9] and OSLOM2 [13] in Table 1. We use the –fast option to get the fastest results for OSLOM2. The time of OSLOM2 on Livejournal is not available, because the program hung up unexpectedly on this graph. We find that even on the smallest graph WordNet, two OCD algorithms still need 51s or 1913s, respectively. For a fair comparison, we also give the amortized running time of OCD, that is, running time of OCD - running time of the exact algorithm. Then, we compare the performance of OCD and OCS.
suitable for these applications. Is not a strict requirement and our approximate algorithm is quite community has vague meanings, the accuracy of the community more than 79%.

7.4 Quality
In this subsection, we show the quality of the solution provided by the approximate algorithm. Overlapping communities express a certain equivalence relationship on $V$. Any two vertices in $C_i$ are equivalent to each other under this equivalence relationship. Thus, we may use the overlapping ratio of the equivalence relationship to quantify the similarities between exact and inexact results. Let $P_i = \{C_1, C_2, ..., C_k\}$ be the approximate result, and let $C_i'$ be the exact community that includes $C_i$ as a subset. Such $C_i'$ certainly exists and is unique since $P_i$ is finer than the exact communities $P_2$. Then, we define the similarity between $P_i$ and $P_2$ as

$$sim(P_i, P_2) = \max_{1 \leq i \leq k} \sqrt{\frac{|C_i|}{|C_i'|}} \frac{\left|\left|C_i'\right| - 1\right|}{\left|\left|C_i\right| - 1\right|} \tag{10}$$

Clearly, we have $0 \leq sim(P_i, P_2) \leq 1$.

Figure 8: Accuracy of approximate solutions. Communities produced by approximate solutions are quite close to that produced by exact solutions.

We ran the approximate algorithm with 100 randomly selected valid query vertex on each network under different $k$. For each query vertex, we calculated the similarity value defined in Equation 10. We summarized the average and variance of the similarity value for 100 random query vertices. The results are shown in Figure 8. It is clear that for each tested network and each $k$, more than 70% accuracy can be achieved. In some special cases, for example, on DBLP and Google, almost 90% accuracy can be achieved.

We give the detailed accuracy results of DBLP in Table 2. In the table, for each $k$, 100 randomly selected valid vertices were queried. $\# has\ community$ denotes the number of vertices among them that have valid communities. $\# equivalent$ is the number of vertices for which the approximate algorithm produces the exact results. And $ratio = \frac{\# has\ community}{\# equivalent}$. We can see that for any $k$, more than 79% of query vertices have the exact results. When $k$ increases, the accuracy increases. In many real applications where community has vague meanings, the accuracy of the community is not a strict requirement and our approximate algorithm is quite suitable for these applications.

<table>
<thead>
<tr>
<th>$k$</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
</tr>
</thead>
<tbody>
<tr>
<td>$# equivalent$</td>
<td>66</td>
<td>69</td>
<td>58</td>
<td>43</td>
<td>37</td>
<td>29</td>
</tr>
<tr>
<td>$# has\ community$</td>
<td>84</td>
<td>83</td>
<td>65</td>
<td>49</td>
<td>40</td>
<td>30</td>
</tr>
<tr>
<td>ratio</td>
<td>79%</td>
<td>83%</td>
<td>89%</td>
<td>88%</td>
<td>93%</td>
<td>97%</td>
</tr>
</tbody>
</table>

Table 2: Accuracy on DBLP. For most query vertices, the approximate solution produces exact results.

5.5 Effectiveness
To show the effectiveness of the OCS model, we computed the proportion of valid query vertices that have communities. Let $x$ denote the number of valid query vertices. We ran the approximate algorithm on 100 randomly selected valid query vertices. Let $x$ denote the number of vertices with communities. Then, $rx\%$ is an estimation of the proportion of vertices that have valid communities in the entire graph. The results are shown in Figure 9.

From both figures, we can see that for each tested real network, our OCS model can recover meaningful communities for a significant number of vertices. On both DBLP and Google, when $k = 4$, among valid query vertices more than 90% of them have communities. For all the tested networks, the ratios of valid vertices and those with communities decrease as $k$ increases. This can be naturally explained, since the constraint on the community is more restricted for larger $k$.

5.6 Influence of $k$
The value $k$ is the most important parameter of OCS. In this subsection, we study the influence of $k$ on the number of communities. We are interested in the number of communities as well as the size of communities.

We first give our analysis about the influence of $k$ on community size and community number. In general, it is expected that the community size monotonically decreases with $k$. When $k$ is small, the constraint for a group of vertices forming a community is weak. Consequently, it is easy to find large communities. When $k$ grows, the community will become smaller. The influence of $k$ on community number is a little complicated. In general, there are two consequences caused by the growth of $k$:

- (1) The constraint on the closeness of a community becomes stronger. As a result, many communities found under a small $k$ will not be a valid community any more. Consequently, the number of communities will reduce when $k$ grows.
- (2) The condition for two communities merged into one single community becomes stronger. As a result, large communities will be broken into small communities. Consequently, the number of communities will increase with $k$.

Hence, it is quite possible that there is a critical point on which two forces are balanced and maximal number of communities can be achieved. To verify this conjecture, we studied communities of ‘Jiawei Han’ in DBLP. We summarized the community number and community size of ‘Jiawei Han’ as the function of $k$.

The results are shown in Figure 10 and they verify our conjecture. From Figure 10(a), we can clearly see that the community size monotonically decreases with $k$. When $k = 5$ and larger, the community size is almost stable. These communities in general are core-coauthoring team members of ‘Jiawei Han’. From Figure 10(b), we can clearly see that $k = 5$ is the critical point on which maximal number of communities are found. Before this, most communities are broken into small communities. After this, valid communities quickly vanish due to the strong constraint on the closeness of a community.

5.7 Influence of $\alpha$ and $\gamma$
In this subsection, we study the influence of $\alpha$ and $\gamma$. We show our results on WordNet. Similar results can be obtained on other
networks. Since the graph is sparse, we only tested our approach on \( k = 3 \) and \( k = 4 \). According to Eq. 1, we have \( \gamma \geq \frac{3}{4} \) for \( k = 3, 4 \). So we chose some representative \( \gamma \) satisfying these constraints. Note that for \( k = 3 \), only \( \gamma = 1 \) is meaningful. Hence, we only show the result for \( \gamma = 1 \) when \( k = 3 \). If \( k = 4 \), when \( \gamma \leq \frac{3}{4} \), the quasi-\( k \)-clique has at least four edges; when \( \gamma < 1 \), the quasi-\( k \)-clique has at least five edges. We used \( \gamma = 0.8 \) and \( \gamma = 0.9 \) as two representative values of these two ranges, respectively. For each selected \( k \) and \( \gamma \), we randomly selected 10 vertices as \( v_0 \) and recorded their average run time in Table 3. When the query time of a vertex exceeds 60s, we considered it as 60s.

From the results, we found that, for the same \( k \) and \( \alpha \), generally a smaller \( \gamma \) costs more run time. This is because higher \( \gamma \) means more strict constraint, which reduces the search space. For the same \( k \) and \( \gamma \), the run time increases monotonically with the growth of \( \alpha \). The reason is similar: a smaller \( \alpha \) leads to a bigger search space.

6. APPLICATIONS

In this section, we propose two typical applications of OCS: diversity based social network analysis and name disambiguation in social networks. Our model and solution enable us to measure the diversity of social ties for a person, which is critical for the diversity based analysis in a quantitative manner. The OCS model and our solution also provide new insights to solve name disambiguation in social networks, which is well-known as a hard problem.

6.1 Diversity-based Social Network Analysis

In social networks, an individual’s social ties are diverse if he or she maintains connections to different communities or groups. Individuals with diverse social ties are more competitive than others [27, 28], because they serve as an intermediary with a position bridging different groups in a social network, and consequently own more opportunities than others in the network. Diversity leading to competition advantage has been widely acknowledged in real life, but has been rarely verified in a quantified manner. Computing overlapping communities is obviously a critical step to quantify the diversity of an individual. Due to the computational hardness of overlapping community detection, previous diversity measures use attributes to derive overlapping communities. For example, Shi et al [29] uses the conferences at which an author has ever published a paper to obtain overlapping communities for DBLP scientific collaboration network.

Now, we can directly use our solution under the OCS model to calculate the diversity of an individual. Intuitively, the diversity of a person can be measured by the number of overlapping communities found by our approach. Next, we show some diversity related analysis on DBLP with diversity calculated by our OCS solution.

Diversity distribution. The first class of fundamental questions are: what is the distribution of diversity? Can we find people with really large diversity? To answer these questions, we summarize the diversity distributions for \( k = 3, 4, 5 \). The results are shown in Figure 11. We find that most persons have diversity 1, indicating that they tend to coauthor with authors in a local research community. These authors are mostly fresh hands in an area. However, there are some authors with large diversity. Some authors have diversity at about 20. These authors are mostly distinguished scientists in computer science. There also some ambiguous names shared by more than one author. We will discuss this problem in the next application.

Diversity leading to competitive advantage. Given the OCS model and the solution, we now have a chance to verify the widely-established conjecture: diversity of social ties leads to competitive advantage in a social network. In a social network, a person’s competitive power in general is positively correlated to his degree. Hence, to verify this conjecture, we summarize the correlation between degree and diversity. The result is shown in Figure 12. We find positive correlations between degree and diversity, which verifies the conjecture. This finding strongly suggests that diversity is one of the important driving forces for an author to become an academic star. This implication opens opportunity for some other interesting real applications, such as predicting rising stars.

6.2 Name Disambiguation

Name disambiguation is a typical problem in social network data management [30, 31]. For example, in DBLP there are at least 40 authors with the same name ‘Wei Wang’. In general, name disambiguation is a challenging problem when person identity information is missing in the data set. The OCS model and its solution provide new insight to solve this problem. They are at least helpful in two aspects: First, they can be used to identify ambiguous names. Second, the overlapping communities themselves are candidate entities with the same name.

Table 4 presents the ambiguous names that have significant number of entities in the DBLP dataset. We found that most of these
names have a large number of communities. Their community numbers in general can be considered as exceptions when compared to the community number distribution observed in Figure 11. For comparisons, we also give the community number of renowned scientists in the database community. It is clear that for a real person, even as famous scientists with many papers and abundant coauthoring relationships, their community numbers are smaller than the community number of ambiguous names. These facts strongly suggest that the large number of overlapping communities is a good indicator of ambiguous names referring to multiple entities. Besides name disambiguation in social networks, OCS is also helpful for sense disambiguation in semantic networks, as shown in the case study on WordNet.

7. CONCLUSION
Most real networks have overlapping community structures. In this paper we propose a novel overlappingness-aware community search problem: overlapping community search. Compared to overlapping community detection, our model is much lightweight and supports online query answering. We reveal the rationale behind the model and provide theoretical guidelines for tuning the model. We devise several exact algorithms and an efficient approximate algorithm to find meaningful overlapping communities. We conduct extensive experiments to show that both the exact algorithms and approximate algorithms are effective to discover meaningful overlapping communities in real networks, and the approximate solution is quite efficient, supporting online (within several ms) query answering on million-node graphs.

8. REFERENCES