Efficient Maximal Motif-Clique Enumeration over Large Heterogeneous Information Networks

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ABSTRACT

In the heterogeneous information network (HIN), a motif-clique is a "complete graph" for a given motif (or a small connected graph) that could capture the desired relationship in the motif. The maximal motif-cliques of HINs have found various applications in community discovery, recommendation, and biological network analysis. The state-of-the-art algorithm for enumerating maximal motifcliques may have to explore all possible subgraphs of a maximal motif-clique and check whether a maximal motif-clique has been enumerated at each recursive step, which is very time-consuming. To improve the efficiency of enumeration, in this paper, we develop efficient algorithms for maximal motif-clique enumeration over large HINs. We first introduce an order-based framework to avoid duplicated enumeration, which results in lower time complexity compared to the existing algorithm. We then propose a pivot-based pruning strategy, which significantly reduces the search space. We further optimize the process of identifying the candidate sets and locating the subgraphs containing the maximal motif-cliques. Extensive experiments on five real-world HINs demonstrate that our proposed algorithm achieves high efficiency and is up to three orders of magnitude faster than the state-of-the-art algorithm.

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The source code, data, and/or other artifacts have been made available at https://github.com/EnderturtleOrz/VLDB2024-Mclique.

1 INTRODUCTION

Heterogeneous information networks (HINs) are networks with multiple typed objects and multiple typed links denoting different semantic relations. These graph data sources are prevalent in various domains, including bibliographic networks [70, 73], co-purchasing networks [71, 92], and knowledge graphs [31]. For example, Figure

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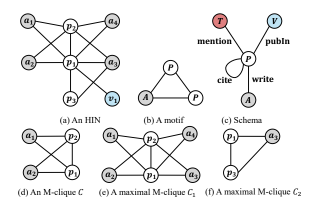


Figure 1: An example HIN of DBLP network.

1(a) illustrates an example HIN of the DBLP bibliographic network, consisting of four authors (i.e., a_1, \dots, a_4), three papers (i.e., p_1, p_2, p_3), and one venue (i.e., v_1).

In this paper, we study the efficient solutions for the problem of Maximal Motif-Clique Enumeration (or MMCE problem) over large HINs [38, 46]. As a fundamental graph mining topic, maximal clique enumeration (MCE) has gained tremendous attention [9, 14, 21-23, 29, 58, 75], but little has been paid to MCE on HINs. Recently, the MCE on HINs [38, 46] has been studied by using a motif to capture the desired relationship in it. A motif, a.k.a. higher-order structure or graphlet, is a small connected subgraph or pattern. As pointed out by [19, 39, 56, 61, 62, 66, 77], a motif is a fundamental building block of large and complex networks, and it enables "higher-order semantics" analysis for HINs. Figure 1(b) depicts a triangle motif, describing that an author writes two papers and one paper cites the another one. The "higher-order semantics" is a widely-used concept in graph mining [4, 51, 62, 89], indicating the complex multi-hop relationships captured by a motif, instead of an edge with direct connection. Given a motif, a motif-clique (M-clique) [38, 46] is a "complete graph" for capturing the higher-order semantics in the motif. Conceptually, the M-clique is a generalization of the classic clique for HINs, representing a user's defined patterns (motifs) rather than edges; as such, it is a "complete subgraph" according to the motif. For example, if a motif \mathcal{M} is an edge linked by two vertices of the same type, then the M-clique is a clique in a homogeneous network; if \mathcal{M} is an edge linked by two vertices of different types, then the M-clique actually is a biclique [1, 17] in a bipartite graph.

Given an HIN \mathcal{H} and a motif \mathcal{M} , a vertex set R is called an M-clique, if for any subset T of its vertices, that contains the same

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number of vertices and the same types of vertices as \mathcal{M} , the induced subgraph of T is subgraph isomorphic to \mathcal{M} . For example, Figure 1(d) shows an M-clique C, since both subsets $\{a_1,p_1,p_2\}$ and $\{a_2,p_1,p_2\}$ satisfy the above constrains. An M-clique is maximal if it is not a subgraph of any other M-clique. Figures 1(e) and (f) present two maximal M-cliques for the above motif in the HIN of Figure 1(a). Similar to the classic maximal clique enumeration (MCE) in homogeneous networks that has been applied to various applications like social network analysis [45, 59], financial network analysis [8], and biological network analysis [43], the MMCE on HINs has also found many real-world applications, to name a few:

- Community discovery [38]. In social networks (e.g., Facebook), the communities can be modeled as maximal M-cliques, by using a specific motif as a guide. Our experiments in Section 7.4 show that the maximal M-cliques are effective for revealing communities in DBLP network. For example, given a motif that represents two authors who have published the same papers, we can discover the maximal M-cliques [38], which can be viewed as close-knit communities, since all members have a close collaborative relationship.
- Bundle recommendation [12]. The MMCE solutions can be used to enhance the performance of bundle recommendation models, since a bundle can often be described by a motif. As shown in our case study in Section 7.4, the MMCE algorithms can improve the performance of SOTA bundle recommendation method BGCN [12], by first finding maximal M-cliques of motifs of bundles and then augmenting the networks of users, items, and bundles.
- Biological network analysis [46]. In a biological graph, the maximal M-cliques can disclose new side effects of a drug, and potential drugs for healing diseases. For example, in the HIN of diseases, genes, and drugs, discovering the maximal M-cliques under specified motifs (e.g., a drug node and a disease node linking to the same gene node) could be used to study medicine characteristics and find potential side effects of drugs [46].

While it is very useful, the MMCE problem is NP-hard [38]. Hu et al. attempted to solve it by proposing a recursive backtracking algorithm META [38] based on the Bron-Kerbosch (BK) algorithm [9], which is a classic MCE algorithm for homogeneous graphs. It first identifies all subgraphs to which the motif is isomorphic. Here, the sets of vertices of these subgraphs are also called the *motif* instances of the motif. Then, for each motif instance, it iteratively enumerates all the maximal M-cliques that contain it by following a node expansion process. However, META is very costly, because to enumerate all the maximal M-cliques containing a specific motif instance, it needs to explore all the possible M-cliques that contain it. Besides, it treats checking whether each maximal M-clique has been enumerated as a subset query problem, i.e., if the M-clique *R* includes a motif instance Γ that is already visited, then the expansion of R can be skipped. Although it can avoid outputting duplicated maximal M-cliques, it is still inefficient since adding each new vertex to an M-clique requires a subset query.

We notice that to avoid duplicated search and reduce the time complexity of MCE, the previous algorithms of the classic MCE problem often define a search order for the vertices [17, 24, 26, 29, 48, 54]. The main idea is that for each vertex v, only its neighbors with an order greater than v need to be included in the enumeration process, so the overall search space can be dramatically reduced. However, the search order above cannot be directly applied to MMCE problem either. This is because, unlike MCE, MMCE needs to iteratively find maximal M-cliques containing each motif instance,

rather than a single vertex, and defining an order for these motif instances is not straightforward. Besides, the BK algorithm can be significantly speeded up by exploiting the pivot principle [9, 29, 58, 75], whose key idea is that every maximal clique must contain either a vertex p or a non-neighbor of p. Thus, during the MCE expansion process, if a vertex p (known as a pivot vertex) is selected, then all its neighbors can be pruned in the current recursion, thus significantly reducing search space. Unfortunately, we demonstrate that the pivot principle cannot be directly applied to MMCE, because in each recursion, the neighbors of a pivot vertex may form a larger M-clique with the current M-clique, so it cannot be skipped.

Our technical contributions. To resolve the above issues, we first propose a new elegant order-based framework that can incorporate different kinds of vertex orders. Under the order-based framework, in the enumeration process, for each motif instance Γ , we only take into account the larger-order vertices. Here, a larger-order vertex is a vertex, if its order is higher than the maximum order of vertices that are with the same type in Γ . By exploiting the orders, the search space can be reduced dramatically.

We also design a novel pivot principle, called motif-pivot, which is very different from the previous pivot principle. The key idea of motif-pivot is that when a pivot vertex p is selected, we first identify a set I(p) of vertices, called M-clique precedence set, for reducing the search space. More precisely, in each recursion, any M-clique in $R \cup I(p)$ can be enlarged by p. Since $R \cup I(p)$ does not contain any solutions, the enumeration algorithm can safely prune the vertices in I(p) from the candidate set, which contains all the possible vertices that are in the maximal M-cliques. However, given a pivot vertex p, detecting the M-clique precedence set I(p) is an NP-hard problem, as we will prove later. To tackle this challenge, we propose a fast algorithm to quickly identify an approximate subset of I(p). As shown in our experiments later, this approximate set is very close to the original set, and it also can be quickly identified.

In addition, to further improve the efficiency, we introduce two non-trivial optimization techniques. The first one is to quickly identify the candidate set for each motif instance, and the second one is a graph reduction approach for eliminating edges and vertices that are not included in any maximal M-cliques.

By combining the above techniques together, we obtain a fast Pivot and Order-based MMCE Algorithm, also called POMA. We have performed extensive efficiency evaluation on five real-world large HINs. The results show that POMA achieves higher efficiency and scalability than the state-of-the-art algorithm META on all datasets, and is up to three orders of magnitude faster than META. We have also conducted two case studies on real-world HINs which show that enumerating maximal M-cliques could be useful in community discovery and bundle recommendation.

Contributions. In summary, our main contributions are:

- We propose an elegant order-based search framework to avoid duplicated computation;
- We develop a novel pivot principle to reduce the search space which significantly speedups the process of MMCE;
- We propose non-trivial optimization techniques to quickly identify the candidate set for each motif instance and locate the subgraphs containing the maximal M-cliques.
- We conduct experiments on five real-world large HINs to demonstrate the efficiency and scalability of our algorithm.

Outline. We formally present the MMCE problem in Section 2. Section 3 analyzes the limitations of the state-of-the-art algorithm.

The overall POMA algorithm and two optimizations are presented in Section 4. The order-based and pivot-based of our POMA algorithm are detailed in Sections 5 and 6, respectively. We report the experimental results in Section 7. We review the related works in Section 8 and conclude in Section 9. For lack of space, all the proofs in this paper are included in the technical report [90].

2 PROBLEM DEFINITION

In this section, we formally present the definition of MMCE problem. Table 1 summarizes the notations frequently used in this paper.

Table 1: Notations and meanings.

Notation	Meaning
$\mathcal{H} = (\mathcal{V},$	An HIN with vertex set $\mathcal V$, edge set $\mathcal E$, vertex type
$\mathcal{E}, \psi, \phi)$	mapping function ψ , edge type mapping function ϕ
$\mathcal{M} = (\mathcal{V}_{\mathcal{M}},$	A motif with vertex set $V_{\mathcal{M}}$, edge set $\mathcal{E}_{\mathcal{M}}$, vertex type
$\mathcal{E}_{\mathcal{M}}, \psi, \phi)$	mapping function ψ , edge type mapping function ϕ
Γ	A motif instance of $\mathcal M$ in $\mathcal H$
Φ	All motif instances of $\mathcal M$ in $\mathcal H$
$\mathcal{N}(v)$	The set of neighbors of v in \mathcal{H} : $\mathcal{N}(v) = \{u \in \mathcal{V} v \in \mathcal{V}\}$
N(0)	$(u,v)\in\mathcal{E}$
$\mathcal{N}_{\mathcal{M}}(v)$	The set of neighbors of v in \mathcal{M} : $\mathcal{N}_{\mathcal{M}}(v) = \{u \in \mathcal{V}_{\mathcal{M}} v \in \mathcal{N}_{\mathcal{M}} v \in N$
$\mathcal{M}(v)$	$(u,v)\in\mathcal{E}_{\mathcal{M}}$
Δ	The number of motif instances of $\mathcal M$ in $\mathcal H$
\mathcal{D}^R_u	The set of vertices in R dominated by u
$\mathcal{H}[T]$	The subgraph of \mathcal{H} induced by vertices set T
h	The number of vertex types in $\mathcal M$
\mathcal{P}_R	The candidate pivot set of <i>R</i>

DEFINITION 1 (HIN [73]). An HIN is an undirected graph $^1\mathcal{H} = (\mathcal{V}, \mathcal{E}, \psi, \phi)$ with a vertex type mapping function $\psi : \mathcal{V} \to \mathcal{A}$, and an edge type mapping function $\phi : \mathcal{E} \to \mathcal{R}$, where \mathcal{A} is the set of vertex types, \mathcal{R} is the set of edge types, each vertex $v \in \mathcal{V}$ belongs to a vertex type $\psi(v) \in \mathcal{A}$, each edge $e \in \mathcal{E}$ belongs to an edge type $\phi(e) \in \mathcal{R}$, $|\mathcal{A}| \geq 1$, $|\mathcal{R}| \geq 1$, and $|\mathcal{A}| + |\mathcal{R}| > 2$.

An HIN often follows a schema, or a graph defined over vertex types \mathcal{A} and edge types \mathcal{R} , denoted by $T_G = (\mathcal{A}, \mathcal{R})$. The HIN schema describes all allowable edge types between vertex types. Figure 1(b) gives the HIN schema of DBLP network. We use $h = |\mathcal{A}|$ to denote the number of vertex types in \mathcal{A} , and use \mathcal{A}_i to denote the i-th vertex type in \mathcal{A} . Denoted by $\mathcal{N}(v) = \{u \in \mathcal{V} | (u, v) \in \mathcal{E}\}$, the set of neighbors of v in \mathcal{H} .

Given an HIN \mathcal{H} , a motif is a small connected HIN $\mathcal{M} = (\mathcal{V}_{\mathcal{M}}, \mathcal{E}_{\mathcal{M}}, \psi, \phi)$. Notice that a valid motif must adhere to the constraints imposed by \mathcal{H} 's schema, including adhering to vertex types defined by the schema and ensuring that edges follow the vertex relationships permitted by the schema. Denote by $\mathcal{N}_{\mathcal{M}}(v) = \{u \in \mathcal{V}_{\mathcal{M}} \mid (u,v) \in \mathcal{E}_{\mathcal{M}}\}$, a set of neighbors of v in \mathcal{M} . Since a motif is often small, it cannot capture the cohesive relationship among many vertices. Thus, people are more interested in motif-based subgraphs.

Definition 2 (Subgraph isomorphism [63]). A motif \mathcal{M} is subgraph isomorphic to an HIN \mathcal{H} , if there exists an injective mapping $\tau\colon \mathcal{V}_{\mathcal{M}} \to \mathcal{V}$, s.t., $\forall u \in \mathcal{V}_{\mathcal{M}}, \ \psi(u) = \psi(\tau(u))$ and $\forall u,v \in \mathcal{V}_{\mathcal{M}},$ if $(u,v) \in \mathcal{E}_{\mathcal{M}}$, then $(\tau(u),\tau(v)) \in \mathcal{E}$ and $\phi(u,v) = \phi(\tau(u),\tau(v))$, where $\tau(u)$ is the vertex to which u is mapped.

Clearly, \mathcal{M} is subgraph isomorphic to \mathcal{H} iff \mathcal{M} is isomorphic to a subgraph (not necessarily induced) of \mathcal{H} .

DEFINITION 3 (TYPE-MATCHED VERTEX SET [38]). Given an HIN \mathcal{H} and a motif \mathcal{M} , a vertex set $T \subseteq \mathcal{V}$ is a type-matched vertex set of \mathcal{M} , if T and \mathcal{M} have the same number of vertices (i.e., $|T| = |\mathcal{V}_{\mathcal{M}}|$), and there exists a bijection $\tau: T \to \mathcal{V}_{\mathcal{M}}$, s.t., $\forall u \in T, \psi(u) = \psi(\tau(u))$.

In the case without ambiguity, we will simply use "match-set" to denote "type-matched vertex set". If a motif \mathcal{M} is subgraph isomorphic to the induced subgraph of a match-set T, then T is called a motif instance of \mathcal{M} .

DEFINITION 4 (MOTIF-CLIQUE [38, 46, 80]). Given an HIN \mathcal{H} and a motif \mathcal{M} , a vertex set R is a motif-clique (M-clique), if for each match-set T in R, \mathcal{M} is subgraph isomorphic to the induced subgraph $\mathcal{H}[T]$.

An M-clique R is maximal if there does not exist any other vertex set R' in \mathcal{H} such that R' is an M-clique and $R \subseteq R'$. Since a maximal M-clique often contains an exponential number of small M-cliques, enumerating all M-cliques will be extremely costly. Thus, we focus on enumerating maximal M-cliques rather than all M-cliques. We now formally present the definition of MMCE problem [38, 46].

PROBLEM 1 (MMCE PROBLEM [38, 46]). Given an HINH and a motif \mathcal{M} , enumerate all the maximal M-cliques of \mathcal{M} in \mathcal{H} .

EXAMPLE 1. Consider the HIN and motif in Figure 1. There are five match-sets of \mathcal{M} , i.e., $T_1 = \{a_1, p_1, p_2\}$, $T_2 = \{a_2, p_1, p_2\}$, \cdots , $T_5 = \{a_3, p_1, p_2\}$. Clearly, all these match-sets are motif instances of \mathcal{M} , because for each match-set T_i , \mathcal{M} is subgraph isomorphic to $\mathcal{H}[T_i]$. Then, we can find that there are two maximal \mathcal{M} -cliques $R_1 = \{a_1, a_2, a_3, a_4, p_1, p_2\}$ and $R_2 = \{a_3, p_1, p_2\}$ in the HIN, as depicted in Figures 1 (d) and (e) respectively.

3 EXISTING APPROACH

In this section, we review existing algorithm META [38] for solving the MMCE problem, and then discuss its limitations.

3.1 The META algorithm

To solve the MMCE problem, Hu et al. [38] proposed a recursive backtracking algorithm META based on the classic BK algorithm [9], as shown in Algorithm 1. The main idea is to maintain three disjoint sets R, C, and X in the recursive enumeration procedure, where R is an M-clique, C is a set of candidates that can be added to R to form a larger M-clique, and X is a set of vertices that have already been explored from C. In each recursion, the three sets keep the invariance that $R \cup \{v\}$ is an M-clique if $v \in C \cup X$, and $R \cup \{v\}$ is not an M-clique if $v \notin C \cup X$. The algorithm recursively processes the vertices in C to expand the current M-clique R, until $C = \emptyset$.

In Algorithm 1, it first computes all the motif instances of $\mathcal M$ (line 1). Then, for each motif instance Γ , it invokes the procedure GetMMC to enumerate all the maximal M-cliques that contain it (lines 2-5). GetMMC recursively processes each vertex $u \in C$ to expand the current M-clique R. In each iteration, it updates the sets C and X using function Refine to keep the invariance that $R \cup \{u\}$ is also an M-clique when $u \in C' \cup X'$ (lines 15-16). Afterwards, it removes u from C and adds it into X (line 12). Finally, GetMMC stops when all the vertices in C are processed.

Lemma 3.1 ([38]). Given an HIN $\mathcal H$ with n vertices and a motif $\mathcal M$, META costs $O(\gamma + \alpha n!)$ time, where γ and α denote the time cost of finding all motif instances of $\mathcal M$ in $\mathcal H$ and checking whether a vertex can enlarge the current M-clique, respectively.

 $^{^{\}rm 1}$ In this paper, we follow the [38] and focus on undirected HINs for simplicity, but our techniques can be readily extended to handle the directed HINs.

Algorithm 1: META $(\mathcal{H}, \mathcal{M})$ [38]

```
input : An HIN {\mathcal H} and a motif {\mathcal M}
    output: All the maximal M-cliques of \mathcal{M} in \mathcal{H}

Φ ← compute all motif instances of M in H;

2 foreach \Gamma \in \Phi do
          \Gamma' \leftarrow \{u | u \in V \setminus \Gamma \land \mathcal{N}(u) \cap \Gamma \neq \emptyset\};
          C \leftarrow \mathsf{Refine}(\Gamma, \Gamma');
         GetMMC(\Gamma, C, \emptyset);
 5
6 Function GetMMC(R, C, X):
          if C \cup X = \emptyset then report R as a maximal M-clique;
          foreach u \in C do
                C' \leftarrow \mathsf{Refine}(R \cup \{u\}, C);
 9
                X' \leftarrow \mathsf{Refine}(R \cup \{u\}, X);
10
                GetMMC(R \cup \{u\}, C', X');
11
               X \leftarrow X \cup \{u\};
12
13 Function Refine(R, C):
          C' \leftarrow \emptyset;
14
          foreach u \in C do
15
           if R \cup \{u\} is an M-clique then C' \leftarrow C' \cup \{u\};
16
          return C'
17
```

LEMMA 3.2 ([38]). Given an HIN \mathcal{H} and a motif \mathcal{M} , META costs $O(\Delta)$ space where Δ is the number of motif instances of \mathcal{M} in \mathcal{H} .

Note that verifying whether a vertex can be added to an existing M-clique to generate a larger one is also NP-hard [38]. To alleviate this issue, Hu et al. [38] proposed a pruning criterion by introducing the concept of dominance relationships among vertices.

DEFINITION 5 (DOMINANCE [38]). Given an HIN \mathcal{H} , an M-clique R and two vertices $u \in \mathcal{V} \setminus R$ and $v \in R$ with $\psi(u) = \psi(v)$, v is dominated by u if $(\mathcal{N}(v) \cap R) \subseteq (\mathcal{N}(u) \cap R)$.

Let \mathcal{D}_v^R denote the set of vertices in R dominated by v. As proved in [38], if $1 + |\mathcal{D}_v^R| > t$, where t denotes the number of vertices in M with type $\psi(v)$, we can directly append v from C to R without performing any additional checks.

In this case, for any match-set $T \subseteq R \cup \{u\}$ containing $u, T \cup \{v\} \setminus \{u\}$, i.e., replacing u with any $v \in \mathcal{D}_u^R$, is a motif instance, which means T is also a motif instance. Besides, Hu et al. presented an early stop strategy and set-trie structure to avoid redundant searches. For lack of space, we omit the details.

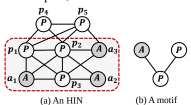


Figure 2: An example illustration the limitations of META.

3.2 Limitations of META

A major limitation of META is that to find a maximal M-clique, it has to explore all the possible non-maximal M-cliques contained by it, which is very costly. For motif instance, consider the HIN (shown in the shaded region), and motif in Figure 2. To enumerate all maximal M-cliques containing the motif instance $\Gamma = \{a_1, p_1, p_2\}$, META has to explore a total number of seven M-cliques, which respectively include the (2^3-1) subsets of $\{a_2, a_3, p_3\}$, but there is

only one maximal M-clique containing Γ . Meanwhile, when a new vertex is added into R to expand the current M-clique, META needs to check if this M-clique is already checked (i.e., before line 10 in Algorithm 1), which is also time-consuming. Our later experiments show that META is very inefficient for processing large HINs (e.g., on a DBLP dataset with two million edges, it cannot enumerate all the maximal M-cliques for 100 motifs within 30 days). Hence, there is much room for improvement.

Opportunities: The most of the existing MCE algorithms [17, 24, 26, 29, 48, 54] utilize a fixed total search order to ensure that each maximal clique is enumerated only once, which dramatically shrinks the search space and also reduces the time complexity. The key idea is that for each vertex v, only its neighbors with an order greater than v need to be included in the enumeration process. However, the search order above cannot be directly applied to the MMCE problem either. This is because, unlike MCE, MMCE needs to iteratively find maximal M-cliques containing each motif instance, rather than a single vertex. Hence, defining an order for these motif instances is not easy.

Besides, the pivot principle has been widely used to accelerate MCE [9, 29, 75]. The key idea is that any maximal clique of a graph either contains a vertex v or a non-neighbor of v. In other words, the clique containing only the neighbors of v must not be a maximal clique, because v can be added into it to form a larger one. Thus, if a vertex v in $C \cup X$, called a pivot vertex, is selected, then all neighbors of $v \in C$ can be skipped to expand the current clique R, thus significantly reducing the number of recursive calls. Unfortunately, the pivot principle cannot be directly applied to the MMCE problem, since the key property for maximal cliques above does not hold for maximal M-cliques — the maximal M-clique may not contain v but only contains its neighbor. Reconsider the HIN and motif in Figure 2, for example, the vertex set $\{a_1, p_1, p_2, p_4\}$ is an M-clique. It does not contain vertex p_5 but only contains one of its neighbor p_4 , since $\{a_1, p_4, p_5\}$ is not a motif instance of the motif.

Inspired by the discussions above, we develop a novel $\underline{\mathbf{P}}$ ivot and $\underline{\mathbf{O}}$ rder-based $\underline{\mathbf{M}}$ MCE $\underline{\mathbf{A}}$ lgorithm POMA in the following sections.

4 OUR POMA ALGORITHM

In this section, we first give an overview of our POMA algorithm, and then introduce the two optimization techniques.

4.1 Overview of POMA

Algorithm 2 gives an overview of POMA which sequentially performs the following five steps:

- (1) Removing vertices and edges that cannot be included in any M-cliques by reduceHIN from the HIN (see Section 4.2);
- (2) Computing all motif instances of motif \mathcal{M} in the HIN by using a subgraph isomorphism algorithm (e.g., VF3 [11]);
- (3) Identifying candidate sets by selectCand (see Section 4.3);
- (4) For each motif instance Γ, imposing an order constraint to reduce the redundant computation (see Section 5);
- (5) For each motif instance Γ, finding all the maximal M-cliques containing it recursively with motif-pivot principle (see Section 6);

Next, we present the algorithms for Steps (1) and (3) in Sections 4.2 and 4.3 respectively. The two key techniques of ordering strategy and pivot principle in Steps (4) and (5) will be discussed in Sections 5 and 6 respectively. In Step (2), any subgraph isomorphism algorithm can be used and we use the SOTA algorithm VF3 [11].

Algorithm 2: $POMA(\mathcal{H}, \mathcal{M})$

4.2 Graph reduction

We introduce two reduction rules to eliminate some vertices and edges that cannot be included in any M-cliques.

• **Rule 1.** For each vertex $v \in \mathcal{V}$, if there does not exist a vertex $v' \in \mathcal{V}_{\mathcal{M}}$ such that $\psi(v) = \psi(v')$, then we can remove v from \mathcal{H} directly, since it is not included in any M-clique.

Similarly, for each edge $e \in \mathcal{E}$, if there does not exist an edge $e' \in \mathcal{E}_{\mathcal{M}}$ such that $\phi(e) = \phi(e')$, then we can remove e from \mathcal{H} directly.

Next, we present two definitions before introducing Rule 2.

DEFINITION 6 (MOTIF ORBIT [51]). Given a motif \mathcal{M} , two vertices $u, v \in \mathcal{V}_{\mathcal{M}}$ are in the same orbit, if there is an automorphism mapping $\tau : \mathcal{V}_{\mathcal{M}} \to \mathcal{V}_{\mathcal{M}}$ with $\tau(u) = v$.

Note that if vertex u is in the same orbit as vertex v, and v is in the same orbit as vertex x, then u and x must be in the same orbit.

DEFINITION 7 (ORBIT TYPE). Given a motif M, a vertex type γ is said to be an orbit type, if M contains either a single vertex of type γ or all vertices of type γ belong to the same orbit.

Note that for any two vertices x, y of the same orbit type in the motif, we can assert that the neighbors of x and y have the same number of vertices of each type in \mathcal{M} . Otherwise, x and y cannot be on the same orbit, which contradicts our assumption.

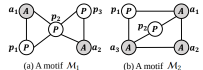


Figure 3: Illustration of the orbit type.

EXAMPLE 2. In the motif \mathcal{M}_1 of Figure 3, "author" is an orbit type, since there exists an automorphism mapping $\tau: \mathcal{V}_{\mathcal{M}_1} \to \mathcal{V}_{\mathcal{M}_1}$ with $\tau(a_1) = a_2, \tau(p_1) = p_3$, and $\tau(p_2) = p_2$. However, "paper" is not an orbit type, because p_2 has a larger degree than p_1 and p_3 .

• **Rule 2.** Given two vertices $v \in \mathcal{V}$, $x \in \mathcal{V}_{\mathcal{M}}$, where $\psi(v) = \psi(x)$ and $\psi(x)$ is an orbit type in the motif \mathcal{M} , if there exists a vertex type γ such that

$$|\{u \in \mathcal{N}(v) \mid \psi(u) = \gamma\}| < |y \in \mathcal{N}_{\mathcal{M}}(x) \mid \psi(y) = \gamma\}|, \qquad (1)$$

then we can remove v from $\mathcal H$ because it is not included in any maximal M-clique.

By combining the aforementioned two rules, when a motif \mathcal{M} is specified, we can locate the maximal M-cliques of \mathcal{M} in a small subgraph of \mathcal{H} by iteratively removing vertices and edges that violate the above rules from \mathcal{H} . We refer to this graph reduction algorithm as reduceHIN and provide the details in the technical report [90].

4.3 Fast candidate set calculation

To compute new candidates, we can use the dominance relationship between two vertices to significantly boost efficiency. Nonetheless, when $R = \Gamma$, the number of vertices that can be dominated by vertices in R is quite limited. Consequently, the process of determining the candidate set C for a motif instance Γ remains computationally intensive. Recall that each vertex u in the candidate set satisfies that $\Gamma \cup \{u\}$ forms an M-clique. To solve this computational bottleneck, we propose a novel algorithm that effectively computes the candidates set for all motif instances in polynomial time. Before introducing our algorithm, we give the following definition.

DEFINITION 8 (MOTIF INSTANCE NEIGHBOR). Given an HIN \mathcal{H} , a motif \mathcal{M} , and two motif instances Γ_1 , Γ_2 of \mathcal{M} , Γ_1 and Γ_2 are motif instance neighbor if $|\Gamma_1 \setminus \Gamma_2| = 1$, and $\Gamma_1 \cup \Gamma_2$ is also an \mathcal{M} -clique of \mathcal{M} . Let $\mathcal{S}[\Gamma]$ denote the set of motif instance neighbors of Γ .

Lemma 4.1 shows how to identify the candidate set.

LEMMA 4.1. Given a motif instance Γ of \mathcal{M} , the candidate set C of Γ can be identified by gathering the different vertices from all motif instance neighbors of Γ using the following equation:

$$C = \bigcup_{\Gamma_i \in \mathcal{S}[\Gamma]} \{ \Gamma_i \backslash \Gamma \} \tag{2}$$

According to Lemma 4.1, we can directly compute the candidate set for a motif instance Γ by identifying its motif instance neighbors. The following three steps outline the process: (1) compute a motif instances set $\mathcal{T}[\Gamma]$, s.t., $\forall T \in \mathcal{T}[\Gamma]$, $|T \setminus \Gamma| = 1$; (2) for each motif instance $T \in \mathcal{T}[\Gamma]$, denoting $u = T \setminus \Gamma$, if all match-sets in $\Gamma \cup \{u\}$ containing u are also motif instances of M, T is a motif instance neighbor of Γ ; (3) the candidate set for Γ is computed based on Lemma 4.1. The detailed steps are illustrated in Algorithm 3.

$\textbf{Algorithm 3:} \ \texttt{selectCand}(\mathcal{H}, \Phi)$

```
input : An HIN \mathcal{H} and a motif instance set \Phi of motif \mathcal{M} output: The candidate sets for all motif instances in \Phi

1 C \leftarrow \emptyset;  // The candidate sets

2 foreach \Gamma \in \Phi do

3  \mathcal{T}[\Gamma] \leftarrow \{T|(|T\backslash\Gamma| = 1) \land T \in \Phi\};

4  foreach T \in \mathcal{T}[\Gamma] do

5  u \leftarrow T\backslash\Gamma;

6  u \leftarrow T\backslash\Gamma;

9  u \leftarrow T\backslash\Gamma;

10 u \leftarrow T\backslash\Gamma are within u \leftarrow T if all match-sets in u \leftarrow T are within u \leftarrow T instance neighbors

10 u \leftarrow T\backslash\Gamma;

11 u \leftarrow T\backslash\Gamma;

12 u \leftarrow T\backslash\Gamma;

13 u \leftarrow T\backslash\Gamma;

14 u \leftarrow T\backslash\Gamma;

15 u \leftarrow T\backslash\Gamma;

16 u \leftarrow T\backslash\Gamma;

17 u \leftarrow T\backslash\Gamma;

18 u \leftarrow T\backslash\Gamma;

19 u \leftarrow T\backslash\Gamma;

10 u \leftarrow T\backslash\Gamma;

11 u \leftarrow T\backslash\Gamma;

12 u \leftarrow T\backslash\Gamma;

13 u \leftarrow T\backslash\Gamma;

14 u \leftarrow T\backslash\Gamma;

15 u \leftarrow T\backslash\Gamma;

16 u \leftarrow T\backslash\Gamma;

17 u \leftarrow T\backslash\Gamma;

18 u \leftarrow T\backslash\Gamma;

19 u \leftarrow T\backslash\Gamma;

10 u \leftarrow T\backslash\Gamma;
```

We first initialize $C=\emptyset$ and use it to save the candidate sets (line 1). Then, we compute the candidate set for each motif instance Γ in Φ one by one. Specifically, for each motif instance Γ , we compute a motif instance set $\mathcal{T}[\Gamma]$ (line 3). Next, for each motif instance $T\in\mathcal{T}[\Gamma]$, we denote $u=\{T\setminus\Gamma\}$, and then, we calculate all matchesets, denoted as \mathcal{Y} , that are formed by adding u into Γ . If all these match-sets in \mathcal{Y} are also motif instances of \mathcal{M} , T is a motif instance neighbor of Γ (lines 5-8). Afterward, we can compute the candidate set of Γ (line 9). Finally, after iterating all motif instances in Φ , we stop and return C (line 10).

EXAMPLE 3. Reconsider the HIN (depicted in the shaded region), and motif in Figure 2, taking $\Gamma = \{a_1, p_1, p_2\}$ as an example. There

are four motif instances that only have a one-vertex difference from Γ , namely $\Gamma_1 = \{a_1, p_1, p_3\}$, $\Gamma_2 = \{a_1, p_2, p_3\}$, $\Gamma_3 = \{a_2, p_1, p_2\}$, and $\Gamma_4 = \{a_3, p_1, p_2\}$, that differ from Γ by only one vertex. Therefore, we have $\mathcal{T}[\Gamma] = \{\Gamma_1, \Gamma_2, \Gamma_3, \Gamma_4\}$. For the motif instance $\Gamma_1 \in \mathcal{T}$, we can compute $\mathcal{Y} = \{\{a_1, p_1, p_3\}, \{a_1, p_2, p_3\}\}$. Notably, all match-sets in \mathcal{Y} are also motif instances of \mathcal{M} (i.e., Γ_1 , and Γ_2). Hence, Γ_1 is a motif instance neighbor of Γ . After scanning all motif instances in $\mathcal{T}[\Gamma]$, we can identify all the motif instance neighbors of Γ , represented as $\mathcal{S}[\Gamma] = \{\Gamma_1, \Gamma_2, \Gamma_3, \Gamma_4\}$. Therefore, we can compute the candidate set for Γ , $C[\Gamma] = \{a_2, a_3, p_3\}$.

Theorem 4.2. Algorithm 3 correctly computes the candidate sets for all motif instances of \mathcal{M} .

PROOF SKETCH. For any $Y \in \mathcal{T}[\Gamma]$, let $u = (Y \setminus \Gamma)$. Then, all match-sets in $\Gamma \cup \{u\}$ are also motif instances of \mathcal{M} , since $\Gamma \cup \{u\}$ is an M-clique. Hence, any $Y \in \mathcal{T}[\Gamma]$ and all motif instance neighbors of Γ can be computed by Algorithm 3.

LEMMA 4.3. The total time cost of selectCand is $O(|V| \cdot \Delta)$, where Δ is the number of motif instances of M in \mathcal{H} .

PROOF SKETCH. In the worst case, for each motif instance $\Gamma \in \Phi$, we need $O(|\mathcal{V}| \cdot |\mathcal{V}_{\mathcal{M}}|)$ time to compute its motif instance neighbor, where $|\mathcal{V}_{\mathcal{M}}|$ is the number of vertices in the motif, which can be considered as a constant in practice.

In addition, we summarize the time, and space complexities of each step of META and POMA, and the limitations of POMA in our technical report [90].

5 AN ORDER-BASED SEARCH FRAMEWORK

In this section, we introduce a novel ordering technology that can be used to avoid the duplicate enumeration of maximal M-cliques.

5.1 Our order-based search framework

The main idea of the order-based search framework in the MCE problem is that for each vertex v, only its neighbors with an order greater than v need to be included in the enumeration process, so the overall search space can be dramatically reduced, where degree order [48, 81, 87] and degeneracy order [13, 17, 24] are commonly used orders. While order-based techniques for improving the performance of MCE have been well studied, their utilization for accelerating the MMCE process remains unexplored. Unfortunately, those vertex order techniques cannot be directly applied to MMCE, because MMCE needs to iteratively enumerate maximal M-cliques containing each motif instance, rather than a single vertex. Hence, it is desired to design a new order-based search framework for the MMCE problem.

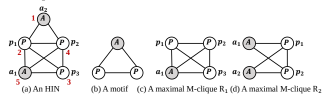


Figure 4: Illustrating the limitations of existing orders.

• A failed attempt. The key challenge in designing an orderbased search framework lies in determining how to compare the order of vertices in a motif instance with that of a vertex in its candidate set. The initial idea is to use either the maximum or minimum order value of all the vertices within a motif instance to denote its order. However, both methods have inherent limitations. We illustrate the limitations via Figure 4, where the order of each vertex is displayed below it in red, and there are four motif instances in the graph, i.e., $\Gamma_1 = \{a_1, p_1, p_2\}$, $\Gamma_2 = \{a_1, p_2, p_3\}$, $\Gamma_3 = \{a_1, p_1, p_3\}$, and $\Gamma_4 = \{a_2, p_1, p_2\}$.

- (1) If the maximum order is adopted, some maximal M-cliques will be missed. Assume that we first detect all maximal M-cliques containing Γ_1 , with $C = \{a_2, p_3\}$ and the maximum vertex order of Γ_1 is 5. As all vertices in C have orders less than 5, R_1 cannot be obtained. Similarly, in the process of detecting the maximal M-cliques containing other motif instances, such as Γ_2 , Γ_3 , and Γ_4 , R_1 remains undiscovered.
- (2) Using the minimum order will lead to redundant computations. When we detect maximal M-cliques containing Γ₁, and Γ₁ has an order of 2, the candidate set *C* is {p₃}. As a result, R₁ will be enumerated. Similarly, when detecting maximal M-cliques containing Γ₃, R₁ will also be enumerated. Hence, the advantages of the order-based search strategy will not be fully utilized.
- A new search framework. To ensure each maximal M-clique is enumerated only once, we introduce a novel order-based search framework. The main idea of our new search framework is that for each motif instance Γ of \mathcal{M} , we only consider larger-order vertices in the candidate set of Γ . Here, a larger-order vertex is a vertex, if its order is higher than the maximum order of the vertices in Λ_{Γ} with the same type. Let $\Lambda_{\Gamma}[\cdot]$ denote the maximum order of the vertices in each vertex type in Γ :

$$\Lambda_{\Gamma}[\gamma] = \max\{\lambda(u)|u \in \Gamma \land \psi(u) = \gamma\},\tag{3}$$

where γ is a vertex type in Γ . Consequently, no result will be missed and each maximal M-clique is enumerated only once, which will be proved in Section 5.3. Based on the above analysis, we develop a new order-divide algorithm using any given vertex order, called Order-Divide, in Algorithm 4.

Algorithm 4: Order-Divide (C, Γ, λ)

input: A candidate set C, a motif instance Γ , and a vertex order λ **output**: A new candidate set C' and not set X

- $1 \ \Lambda_{\Gamma}[\cdot] \leftarrow \emptyset;$
- 2 foreach $v \in \Gamma$ do
- 3 $\Lambda_{\Gamma}[\psi(v)] \leftarrow \max\{\lambda(u)|u \in \Gamma \land \psi(u) = \psi(v)\};$
- 4 $C' \leftarrow \{u | u \in C \land \lambda(u) > \Lambda_{\Gamma}[\psi(u)]\};$
- 5 $X \leftarrow \{u | u \in C \land \lambda(u) < \Lambda_{\Gamma}[\psi(u)]\};$
- 6 **return** C' and X;

Specifically, we first compute the maximum order of the vertices for each vertex type in a motif instance Γ (lines 2-3). Next, we divide the candidate set C into the new candidate set C' and the not set X by only considering the larger-order vertices in C (lines 4-5). Finally, the new sets C' and X are returned (line 6).

Example 4. Reconsider the HIN and the motif instances in Figure 4 with $\Gamma_1=\{a_1,p_1,p_2\},\ \Gamma_2=\{a_1,p_2,p_3\},\ \Gamma_3=\{a_1,p_1,p_3\},\ and\ \Gamma_4=\{a_2,p_1,p_2\}.$ Taking $\Gamma_3=\{a_1,p_1,p_3\}$ as an example, we have $\Lambda_{\Gamma_3}[A]=5$ and $\Lambda_{\Gamma_3}[P]=3$, where 5 represents the order of a_1 (i.e., $\lambda(a_1)=5$) and 3 represents the maximum value of $\lambda(p_1)$ and $\lambda(p_3)$. By running Algorithm 4, we get $C=\{a_2,p_2\},\ C'=\{p_2\},\ and\ X=\{a_2\}$ for Γ_3 . The detailed information is provided in Table 2.

Table 2: Illustrating the Order-Divide.

Motif instances	$\Lambda_{\Gamma_{i}}[\cdot]$	С	C'	X
$\Gamma_1 = \{a_1, p_1, p_2\}$	[5, 4]	$\{a_2, p_3\}$	Ø	$\{a_2, p_3\}$
$\Gamma_2 = \{a_1, p_2, p_3\}$	[5, 4]	$\{a_2, p_1\}$	Ø	$\{a_2, p_1\}$
$\Gamma_3 = \{a_1, p_1, p_3\}$	[5, 3]	$\{a_2, p_2\}$	$\{p_2\}$	$\{a_{2}\}$
$\Gamma_4 = \{a_2, p_1, p_2\}$	[1, 4]	$\{a_1\}$	$\{a_1\}$	Ø

5.2 Ordering heuristics

As shown in Algorithm 4, different vertex orders may have significant effect on the efficiency of MMCE. However, finding the optimal order is an NP-hard problem [26], so in the literature, some heuristic orders have been proposed, such as degree order and degeneracy order, which are described as follows:

- **Degree order.** The degree order [48, 81, 87] is the simplest one, where the vertices are arranged in ascending order based on their degrees. By using the degree order, the size of the largest candidate set is no larger than the maximum degree.
- **Degeneracy order.** Another simple yet efficient ordering is to use the degeneracy order [24, 29], which is widely used in MCE algorithms. Such an order can be obtained by repeatedly peeling a vertex of the minimum degree in the remaining subgraph, where the vertex removing order is exactly the degeneracy order. Such a peeling procedure can be done in O(m+n), where m and n denote the number of edges and vertices in the graph, respectively. In the HIN, we can obtain a degeneracy order by ignoring the vertex and edge types and running classic core decomposition algorithm [3]. Since the degeneracy order is theoretically better than the degree order [29], so we employ it in this paper.

5.3 Theoretical correctness analysis for POMA

Lemma 5.1. For each motif instance Γ of \mathcal{M} , Algorithm 2 can enumerate all the maximal M-cliques containing it without missing any results.

PROOF SKETCH. Suppose there exists a maximal M-clique R containing Γ that cannot be enumerated. We can construct a new motif instance Γ' by selecting vertices of each type with the minimum order from R, and then R can be enumerated by the recursive enumeration starting from Γ' , which contradicts our assumption. \square

Lemma 5.2. Algorithm 2 ensures that each maximal M-clique will only be enumerated exactly once.

PROOF SKETCH. Proof this by contradiction, supposing a maximal M-clique R can be enumerated twice by two motif instances Γ_1 and Γ_2 . However, whether Λ_{Γ_1} is dominated by Λ_{Γ_2} or not, R cannot be enumerated twice.

Lemma 5.3. The total time cost of POMA is $O(\gamma + \alpha \Delta 2^{\beta})$, where $\beta = h \times \delta$, δ is the degeneracy number of the input HIN, h is the number of vertex types in the input motif, Δ is the number of motif instance of $\mathcal M$ in $\mathcal H$, and the other variables have the same meanings as those in Lemma 3.1.

PROOF SKETCH. In the worst-case scenario, each motif instance Γ necessitates the consideration of up to 2^{β} branches in Algorithm 2. For each branch, we need $O(\alpha)$ time to check whether a vertex can expand to the M-clique.

Discussions. Although our ordering framework shares some general purpose with those of classic MCE algorithms [29, 58, 75], it significantly differs from them in three aspects:

- Order definitions: In MCE algorithms, the order is defined based on comparing vertices, while in POMA, the order is defined based on comparing both vertices and motif instances.
- Order-based pruning strategies: In MCE, when enumerating
 the maximal cliques containing a vertex v, only the neighbours of v with larger orders than v need to be considered.
 In contrast, in POMA, we find maximal M-cliques containing
 each motif instance, rather than a single vertex, so we only
 consider "larger-order" vertices. Here, a larger-order vertex
 is a vertex whose order is higher than the maximum order
 of the vertices with the same type in this motif instance.
- Correctness proofs. In MCE problem, the correctness is evident as each maximal clique can be enumerated starting from the vertex with the smallest order in this clique. However, for MMCE problem, we need to theoretically prove Lemma 5.1 and 5.2 to confirm the correctness.

LEMMA 5.4. Given an HIN \mathcal{H} and a motif \mathcal{M} , POMA costs $O(\Delta)$ space, where Δ is the number of motif instance of \mathcal{M} in \mathcal{H} .

PROOF SKETCH. In the process of enumerating all maximal M-cliques, all the motif instances would be stored in the memory. \Box

6 PIVOT-BASED ENUMERATION TECHNIQUES

As discussed in Section 3, the classical pivot principle cannot be directly applied to the MMCE problem. In this section, we present a novel pivot principle for accelerating the MMCE process.

6.1 The motif-pivot principle

Recall that in the MCE on homogeneous graphs, we can just use the neighbors of the pivot to find the vertices that can be pruned. However, in our MMCE, pruning neighbors from the candidate set is not correct. To figure out the vertices to prune, we introduce the concept of M-clique precedence.

DEFINITION 9 (M-CLIQUE PRECEDENCE \prec). Given an HIN \mathcal{H} , a motif \mathcal{M} , an M-clique R, a candidate set C of vertices, and two vertices $u, v \in C$, v has M-clique precedence over u, denoted by $v \prec u$, if the following conditions are met:

- R ∪ {u} is an M-clique due to u ∈ C, and R ∪ {u, v} is also an M-clique;
- if there exists an edge $(x, y) \in \mathcal{E}_{\mathcal{M}}$ satisfying $\psi(x) = \psi(u) \land \psi(y) = \psi(v)$, we have $(u, v) \in \mathcal{E}$.

Let I(u) denote the set of vertices in C that have M-clique precedence over u, i.e., $I(u) = \{v \in C | v < u, u \in C\}$. In each recursion of MMCE, the pivot vertex p should be able to enlarge the M-cliques formed in $R \cup I(p)$, which will be further explained in Lemma 6.1. To ensure this, we introduce the concept of candidate pivot set.

DEFINITION 10 (CANDIDATE PIVOT SET). Given an HINH, a motif \mathcal{M} , and an M-clique R, we define \mathcal{P}_R as a candidate pivot set of R if, for every vertex v in \mathcal{P}_R , the condition $1 + |\mathcal{D}_v^R| > t$ holds, where \mathcal{D}_v^R denotes the set of vertices in R dominated by v, and t represents the number of vertices in \mathcal{M} with type $\psi(v)$.

It is noted that \mathcal{P}_R can be empty, indicating that we cannot prune any search space in this case. However, in practice, the case when

 $\mathcal{P}_R = \emptyset$ is not common, and as shown in our experiments, the pivot technique can significantly avoid redundant computation. Based on the definitions above, we give the following lemma:

Lemma 6.1. For any M-clique R and a vertex u in \mathcal{P}_R , any maximal M-clique $R' \supseteq R$ must include either u or one of the vertices that are not M-clique precedence over u, as otherwise, R' could be enlarged by adding u.

PROOF SKETCH. We assume that a maximal M-clique R' does not contain vertex u or any vertices that are not M-clique precedence over u. In other words, R' satisfies $R \subseteq R' \subseteq R \cup I(u)$. Then, $R' \cup \{u\}$ is also an M-clique, since we each match-set T in $R' \cup \{u\}$ containing u is also a motif instance.

Next, based on Lemma 6.1, we formally introduce our motif-pivot principle. Given a maximal M-clique, there are only two cases: it either contains the pivot vertex p or one of the vertices not in $\mathcal{I}(p)$. If an M-clique does not contain p, then it can only be enlarged by vertices from $C\setminus I(p)$. In this situation, the algorithm can safely prune vertices in I(p) from C.

Efficiently approximating I(p)

Based on the discussion above, for each pivot vertex p, we need to compute its M-clique precedence set I(p). However, as stated in [38], determining whether a vertex can be incorporated into an existing M-clique to form a larger one is an NP-hard problem. Consequently, computing I(p) is also an NP-hard problem, and it is not feasible to directly obtain. Luckily, we can quickly find a subset of I(p) via the dominance relationship. However, this only allows a limited number of vertices to be included in $\mathcal{I}(p)$. To better approximate I(p), we propose additional conditions to include more vertices in I(p) in the following lemma.

LEMMA 6.2 (APPROXIMATING I(p)). Given an HIN \mathcal{H} , a motif \mathcal{M} , an M-clique R and a pivot vertex p, a vertex v in C that satisfies at least one of the following conditions is M-clique precedence over p:

- Condition 1. 1 + |D_v^{R∪{p}}| > t.
 Condition 2. ψ(p) = ψ(v), and there is only one vertex in \mathcal{M} with type $\psi(p)$.
- Condition 3. Let $L = \{x | x \in R' \land \psi(x) = \psi(v)\}$, and $\left(\bigcup_{x\in L} (\mathcal{N}(x)\cap R')\right)\subseteq (\mathcal{N}(v)\cap R'), \text{ where } R'=R\cup\{p\};$

PROOF SKETCH. The Condition 1 is guaranteed by Lemma 6.4 in [38]. For Condition 2, since all new match-sets will contain v but not p, and $R \cup \{v\}$ is an M-clique, $R \cup \{p, v\}$ is an M-clique. For Condition 3, whether the match-set T in $R' \cup \{v\}$ includes v but not p, or both p and v, T is a motif instance.

The above lemma can be easily incorporated into our pivot principle, as discussed in Section 6.1. Specifically, in each recursive call, for the current M-clique R, the algorithm first selects a pivot vertex p from the \mathcal{P}_R . To compute I(p), we applied Lemma 6.2 to quickly detect an approximated subset of $\mathcal{P}(p)$ without subgraph isomorphism checking. The following example illustrates that Lemma 6.2 can better approximate I(p) than the dominance relationship.

Example 5. Consider the HIN and motif in Figure 2. Assume that $R = \{a_1, p_1, p_2\}, C = \{a_2, a_3, p_3, p_4, p_5\}, and p = a_2$. Here, we need to compute I(p). However, if we only rely on the dominance relationship (i.e., Condition 1 of Lemma 6.2), none of the vertices in C satisfy this

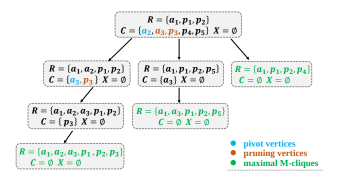


Figure 5: An enumeration tree for the HIN in Figure 2.

condition. According to Condition 2, a_3 can be appended into I(p), since there is only one author in the motif. In addition, based on Condition 3, p_3 can also be added to I(p), as the neighbors of p_1 and p_2 are also neighbors of p_3 .

To compute $\Gamma(p)$ exactly, we need to check whether a vertex vcan expand an M-clique via VF3, which invokes a subgraph isomorphism algorithm. There is no constant factor approximation algorithm for it [63]. Our approximation algorithm can quickly detect a subset of $\Gamma(p)$, which costs $O(n \cdot d_{max})$ time to check whether a vertex can be added into $\Gamma(p)$, while the exact algorithm requires O(n!) time in the worst case. Besides, the approximate I(u) can also be used to efficiently update C and X. Specifically, in Algorithm 1, after adding vertex u into R, we can compute the I(u) based on Lemma 6.2. Then, the vertices in $I(u) \cap C$ and $I(u) \cap X$ can safely be added to C' and X', respectively, without any extra checks.

6.3 A pivot-based enumeration algorithm

In this subsection, we develop a pivot-based enumeration algorithm, called MP-MMC, that fully integrates the proposed motif-pivot principle, as shown in Algorithm 5. Specifically, we first compute the candidate pivot set \mathcal{P}_R of R, and then select a pivot vertex from \mathcal{P}_R (lines 3-4). Note that if \mathcal{P}_R has multiple vertices, we select the one with the maximum degree as the pivot vertex. Next, we compute the M-clique precedence set I(p) based on the Lemma 6.2 (lines 5-8). Then, we iterate over the vertices in $C \setminus I(p)$ to expand the current M-clique R (lines 9-13). Example 6 further illustrates how MP-MMC reduces redundant computation.

```
Algorithm 5: MP-MMC (R, C, X)
   input: An HIN \mathcal{H} and a motif \mathcal{M}
   output: All maximal M-cliques of \mathcal{M} in \mathcal{H}
1 if C \cup X = \emptyset then report R as a maximal M-clique;
_{2} \mathcal{P}_{R} ← compute the candidate pivot set of R;
p ← select a pivot vertex from \mathcal{P}_R;
A R' \leftarrow R \cup \{p\};
5 foreach u \in C do
         if \exists (x, y) \in \mathcal{E}_{\mathcal{M}}, \psi(x) = \psi(p), \psi(y) = \psi(u) then
           if (p, u) \notin \mathcal{E} then continue;
        if Lemma 6.2 is satisfied then I(p) \leftarrow I(p) \cup \{u\};
9 foreach u \in C \setminus I(p) do
         C' \leftarrow \mathsf{Refine}(R \cup u, C);
11
         X' \leftarrow \mathsf{Refine}(R \cup u, X);
         MP-MMC(R \cup u, C', X');
12
        X \leftarrow X \cup \{u\};
```

EXAMPLE 6. Given an HIN $\mathcal H$ and a motif $\mathcal M$ in Figure 2, assume that $\Gamma=\{a_1,p_1,p_2\}$. The initial parameters of MP-MMC are $R=\{a_1,p_2,p_2\}$, $C=\{a_2,a_3,p_3,p_4,p_5\}$, and $X=\emptyset$ respectively. Assume that the pivot vertex of MP-MMC is a_2 (marked in blue in Figure 5). Then, $\{a_3,p_3\}$ (marked in orange in Figure 5) can be pruned from C, as all the vertices are M-clique precedence over a_2 . By our pivot technique, only vertices in $\{a_2,p_4,p_5\}$ are used to expand the current M-clique R. When R is expanded with the vertex a_2 , the candidate set is updated to $\{a_3,p_3\}$, which will be checked in the next recursive call. In the next recursion, suppose the selected pivot vertex is a_3 . Then, p_3 can be pruned from C. Thus, the algorithm only picks a_3 to expand R. The procedure continues until all vertices in C of the top recursion have been processed. The enumeration tree is shown in Figure 5.

textbfDiscussions. Our motif-pivot principle is different from the pivot strategies of existing MCE methods [9, 29, 58, 75] in two aspects:

- Pivot-based pruning strategies: In MCE algorithms, when a pivot p is selected, the neighbours of p can be pruned from C. However, in MMCE, such a pruning set no longer exists since a maximal M-clique may not contain the pivot p, but only contain its neighbours. Thus, we introduce the M-clique precedence relationship, then use the M-clique precedence set of p, I(p), as the pruning set, and finally propose an efficient algorithm to quickly approximate I(p).
- *Pivot selection processes*. In existing MCE algorithms, the pivot vertex is directly selected from the candidate set or not set. However, in MMCE problem, for an M-clique R, we can only select the pivot vertex from the candidate pivot set of R. In Lemma 6.1, we theoretically prove that when a pivot vertex p is selected, the M-cliques in $R \cup I(p)$ can be enlarged by p.

7 EXPERIMENTS

We now present the experimental results. Section 7.1 discusses the setup. We discuss the efficiency results in Sections 7.2 and 7.3. We present the two case studies in Section 7.4.

7.1 Setup

Datasets. We use five real HINs: $Instacart^2$ [38], $WordNet^3$ [72], $DBLP^4$ [31], $DBpedia^5$ [88], and $Freebase^6$ [88]. Their statistics, including the numbers of vertices, edges, vertex types, edge types, and the degeneracy values δ , are reported in Table 3. Instacart is a co-purchasing network, where each product has a type showing its category (e.g., "personal care" and "beverages"), and each edge between two products means they have been purchased together more than 200 times. WordNet is a large lexical database, where nouns, verbs, adjectives and adverbs are grouped into sets of cognitive synonyms (synsets), each expressing a distinct concept. DBLP includes publication records in computer science areas, and the vertex types are authors, papers, venues and topics. DBpedia contains the data extracted from wikipedia infoboxes using the mapping-based extraction. Freebase contains all the entities and relations in the music domain.

Table 3: Datasets used in our experiments.

Dataset	Vertices	Edges	Vertex types	Edge types	δ	Motifs
Instacart	49,688	12,770	21	237	30	100
WordNet	76,853	240,798	5	25	10	100
DBLP	881,039	2,247,195	4	7	14	100
DBpedia	8,970,120	71,403,844	414	79,397	52	100
Freebase	347,463,729	1,110,001,528	10, 801	620,307	168	100

Queries. For each dataset, to generate a motif, we perform a random walk on the data graph to obtain a connected subgraph, following the approach in [7, 72]. For evaluation purposes, we generate five motif sets for each dataset, which contain 100 motifs with 3, 4, 5, 6, and 7 vertices respectively, in line with the range of motif sizes (from 3 to 7) typically encountered in various real-world applications [34, 56, 84, 89]. The default motif size is 4. The structures of motifs used in our paper are shown in our technical report [90]. For example, a motif with two users liking the same genre of movie or item shows value for recommendation purposes, and it can be found in both DBpedia and Freebase [57, 79, 91]. We implement all the algorithms in C++ with STL used and run experiments on a machine having an Intel(R) Xeon(R) Gold 6338R 2.0GHz CPU and 512GB of memory, with Ubuntu installed. If an algorithm cannot finish in 30 days, we mark its runtime as INF.

7.2 Overall efficiency results

In this section, we compare the efficiency of POMA and META.

- 1. Effect of motif sizes. Figure 6 compares the efficiency of these two algorithms by varying motif sizes. Clearly, POMA is up to three orders of magnitude faster than META, because it not only ensures that a maximal M-clique will not be enumerated multiple times but also dramatically reduces the search space, while META includes numerous redundant computation. Besides, as motif size becomes larger, the running time of all algorithms generally increases, since a larger motif means checking subgraph isomorphism is more time-consuming. POMA, on average, only requires a few tens of seconds to enumerate all maximal M-cliques for motifs with three vertices. In addition, on the largest two datasets, META could not complete its run for any motif within one month.
- **2. Scalability test.** For each HIN, we first randomly select 20%, 40%, 60%, 80%, and 100% of its edges and then obtain five sub-HINs induced by these edges, respectively. We then report the average efficiency of POMA and META on these sub-HINs in Figure 7. The time costs of both POMA and META scale almost exponentially with the number of vertices in the graph, but the growth rate of the curve of POMA is smaller, so POMA exhibits better scalability and META.
- 3. Comparing the search space of META and POMA. To measure the size of search space, we count the total number of branches that need to be enumerated by META and POMA (i.e., the number of nodes in the recursive tree), we use "N/A" to denote that the algorithm could not be finished within one month. Figure 8 shows the results on five datasets. Clearly, POMA is more effective for reducing the search space than META, thus achieving higher efficiency.
- 4. Comparing the space usage of META and POMA. We examine the memory usage of META and POMA on all datasets with motif size 4, reporting their results in Figure 9. The META and POMA have the same space complexity $O(\Delta)$, where Δ denotes the number of motif instances. Nevertheless, in practice, POMA usually takes slightly more memory than META, since it adapts a string hash operation in the selectCand step. This is an optimization technique at the

²https://www.instacart.com/datasets/grocery-shopping-2017

³https://wordnet.princeton.edu/

⁴https://www.aminer.cn/citation

⁵https://wiki.dbpedia.org/Datasets

⁶http://freebase-easy.cs.uni-freiburg.de/dump/

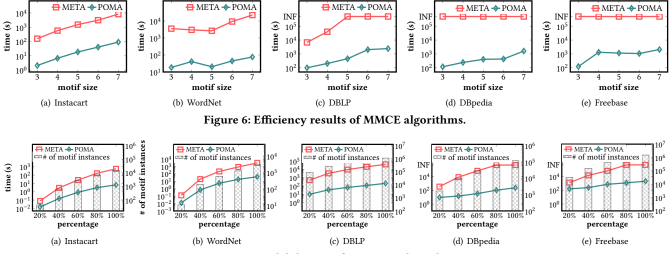


Figure 7: Scalability test for MMCE algorithms.

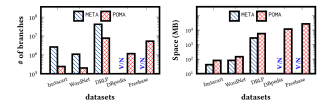


Figure 8: Search space.

Figure 9: Memory usage.

implementation level, which will increase the constant size of the space required by POMA, but will not increase its space complexity.

7.3 Detailed analysis of POMA

We extensively evaluate and analyze POMA from different angles.

- 1. Time cost of different steps in POMA. Recall that POMA sequentially performs the following five steps: (1) reducing the original HIN (reduceHIN), (2) computing all the motif instances (VF3 [11]), (3) identifying the candidate sets selectCand, (4) dividing the candidate sets and not vertex sets Order-Divide, and (5) performing the recursive search (MP-MMC). Figure 10 shows the time cost of these five steps (assuming the graph has been loaded into memory) on five datasets. We see that MP-MMC is the most computationally expensive step on all datasets. On DBpedia and Freebase datasets, the time cost of reduceHIN is significantly larger than that on other datasets. This is because these two datasets contain a much larger number of vertex types, allowing more vertices to be pruned since each motif includes a limited number of diverse vertex types. Besides, running the VF3 algorithm and Order-Divide only account for a relatively small portion of the total time during the execution of POMA, so we should focus on optimizing other steps.
- 2. Ablation study. To evaluate the effect of four key steps, we design four variants by adding reduceHIN, selectCand, ordering framework, and motif-pivot into META, which are denoted by META+R, META+C, META+O, and META+P respectively. We then run META and these variants on all datasets and report the efficiency results in Figure 11. We can see that POMA significantly outperforms the baseline META and the other variants. Moreover, these four variants are faster than META, further substantiating the efficacy of

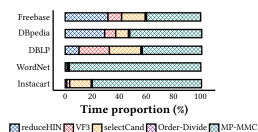
our four proposed techniques, especially on larger HINs. We also design other four variants by removing the four key steps above from POMA, and report their results in our technical report [90].

- **3. Effect of graph reduction technique.** We present the average numbers of vertices and edges before and after the graph reduction for different motifs across all datasets. Clearly, our graph reduction method can substantially reduce the size of the original HIN. The details are shown in our technical report [90].
- 4. Effect of different vertex ordering strategies. In this experiment, we consider two widely used vertex orders: degree order and degeneracy order. Clearly, both orders significantly outperform the case without vertex ordering, and the degeneracy order usually achieves better performance, and results are shown in [90].
- **5. Efficiency of the** I(p) **approximation algorithm.** To show the efficiency of the I(p) approximation algorithm, we denote the POMA that does not employ the algorithm proposed in Section 6.2 as POMA*. Then, we report the sum running time and the number of pruning vertices of POMA and POMA* on all datasets in Table 4. It is noted that if the algorithm cannot finish within three days, we mark its running time as $\geq 259,200$ s. Clearly, POMA significantly outperforms POMA* on larger HINs, as it obviates the need for subgraph isomorphism checking. More specifically, on the two largest datasets (DBpedia and Freebase), POMA* cannot finish within three days. Besides, from a theoretical perspective, POMA can only prune a subset of vertices that are pruned by POMA*. However, in practice, we observe that POMA can prune nearly the same number of vertices as POMA*, which underscores the practical effectiveness of POMA.

Dataset	Instacart	WordNet	DBLP	DBpedia	Freebase
POMA*	1,804.5s	1,877.1s	6,262.3s	≥ 259,200s	≥ 259,200s
POMA	136.2s	827.1s	4,117.9s	4,866.3s	12,097.8s
Speedup	13.2 ×	14.4 ×	1.5 ×	≥ 53 ×	≥ 21.4 ×
POMA*	36.5	117.0	6.37	l –	
POMA	35.6	117.0	6.37	161.8	50.0
ratio	97.5 %	100.0 %	100.0 %	_	_

Table 4: Comparing different pivot techniques.

6. The parallel version of POMA. Recall that in POMA, we need to enumerate the maximal M-cliques containing each motif instance, so the enumeration for all the motif instances can be performed



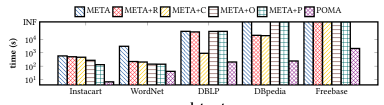


Figure 11: Efficiency comparison of MMCE algorithms.

Figure 10: Proportion of the time cost of each step in POMA.

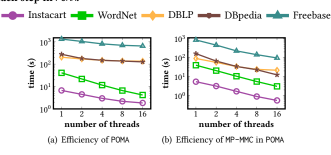


Figure 12: Effect of the number of threads.

	Instacart	WordNet	DBLP	DBpedia	Freebase
γ	0.15 s	0.19 s	46.13 s	19.48 s	6.52 s
α	3.25 us	6.34 us	3.24 us	1.03 us	0.17 us

Table 5: Actual running time of γ and α .

in parallel. Specifically, we have implemented a parallel version of POMA by paralleling the MP-MCC step, denoted by POMA-Par, in which each thread is scheduled for enumerating the maximal Mcliques containing a specific motif instance. Figure 12(a) shows the time cost of the five steps of POMA and POMA-Par, and Figure 12(b) only shows the running time of MP-MMC step, by varying the number of threads from 1 to 16 across all datasets. Clearly, with an increasing number of threads, the overall runtime of POMA-Par exhibits a reduction, and if MP-MCC takes up a large portion of the whole running time, POMA-Par shows a strong parallel scalability. For example, on WordNet dataset, using 16 threads allows POMA-Par to achieve self-speedups of 13 times.

7. Statistical of the hyper-parameters. We mainly use five parameters, i.e., γ , α , Δ , δ , and β . For γ and α , they denote the time costs of computing all motif instances and checking whether a vertex can expand to M-clique respectively. Table 5 reports their actual running time costs. For Δ , δ , and β , they represent the average number of motif instances in the HIN, the degeneracy value of the graph, and the degeneracy multiplying the number of vertex types, respectively. The values of δ are included in Table 3. We report the values of Δ in Figure 7, which range from a few to two million. We observe that the running time of META and POMA on all datasets is proportional to the motif instance numbers since we need to enumerate the maximal M-cliques for each motif instance, which is aligned with our complexity analysis in Section 5.3.

7.4 Case studies

In this section, we present two case studies on real-world HINs.

Table 6: Community quality analysis on Instacart.

Method	Diameter		Simil	arity	
			Household	Breakfast	Frozen
R-com POMA	3.45 2.01	0.15 0.45	0.22 0.53	0.22 0.49	0.19 0.42

(1) Community detection. As the SOTA method of HIN community detection, R-com [42] takes input from a set of relational constraints which can be regarded as a motif, and finds communities with multiple vertex types. In this experiment, we compare the quality of communities detected by POMA and R-com on Instacart and DBLP datasets. The Specifically, we first randomly select 20 motifs with each having 4 vertices. Then, for each motif, we run POMA to detect the corresponding maximal M-cliques, and R-com to get the communities to adhere to the relation constraints in the motif. Finally, we compare the quality of communities in terms of community member similarity and closeness of communities, which are used for measuring community quality [30, 40, 42]. For the former one, we compute the Jaccard similarity for different types of vertices by following the method in [42]; for the latter one, we compute the diameter by following the method in [31, 40]. As shown in Table 6, we observe that the communities of POMA achieve both higher similarity values and smaller diameters than those of R-com on Instacart dataset. The results on DBLP dataset are shown in our technical report [90]. Thus, POMA is able to detect communities with higher quality.

Table 7: Statistics of datasets for bundle recommendation.

Dataset		# of vertice	# of e	dges	
Dataset	#User (U)	#Item (I)	#Bundle (B)	#U-I	#U-B
Netease	18,528	123,628	22,864	1,128,065	302,303
Youshu	8,039	32,770	4,771	138,515	51,377

Table 8: The Recall@K values on BGCN and BGCN-M.

Dataset	Method	Recall@10	Recall@20	Recall@30	Recall@40
Netease	BGCN	0.0369	0.0642	0.0845	0.1013
	BGCN-M	0.0391	0.0650	0.0863	0.1033
Youshu	BGCN	0.1596	0.2410	0.2984	0.3416
	BGCN-M	0.1600	0.2463	0.3018	0.3453

Table 9: The NDCG@K values on BGCN and BGCN-M.

Dataset	Method	NDCG@10	NDCG@20	NDCG@30	NDCG@40
Netease	BGCN	0.0202	0.0274	0.0321	0.0356
	BGCN-M	0.0209	0.0278	0.0327	0.0362
Youshu	BGCN	0.0934	0.1165	0.1303	0.1398
	BGCN-M	0.0955	0.1198	0.1331	0.1427

(2) Bundle recommendation. Our proposed algorithm POMA can be used to enhance the performance of the SOTA bundle recommendation method BGCN, which aims to recommend a bundle of items to a single user [10, 12, 37]. BGCN [12] trains a Graph Neural Networks (GNN)-based recommendation model using three networks, including the user-bundle interaction network, user-item interaction network, and bundle-item affiliation network, and then predict whether a user will interact with a certain bundle. However, due to exposure bias and item diversity imbalance [15, 83], the bundle-item affiliation network cannot capture the precise relationships between bundles and items, which may cause some items to be overlooked and not considered for the bundle, leading to sparse connections between bundles and items.

To alleviate the sparsity issue above, we propose to detect the maximal M-cliques from the network of items, and then use these maximal M-cliques to argument the bundle-item affiliation network, thereby enhancing the performance of BGCN. Specifically, we first build a co-purchasing network between items, where each edge between two items means they have been purchased together more than 5 times. Then, we use POMA to enumerate maximal M-cliques by using motifs formed by items in the bundles. Next, in the bundle-item affiliation network, we link items in the same maximal M-cliques to their respective bundles. Finally, we run BGCN on the augmented networks, and denote this approach by BGCN-M.

We have compared BGCN and BGCN-M on two real-world datasets, i.e., Netease and Youshu, whose statistics are shown in Table 7. The former one was collected [10] from the largest music platform ⁷ in China, which enables users to bundle songs with a specific theme, while the latter one was collected by [18] from Youshu platform, a Chinese book review site 8, where each bundle is a list of books selected by some users. We use Recall@K and NDCG@K [37, 83] to judge the performance of the ranking list, where Recall@K measures the ratio of test bundles that have been contained by the top-K ranking list, while NDCG@K complements Recall by assigning higher scores to the hits at higher positions of the list. As shown in Tables 8 and 9, BGCN-M significantly outperforms BGCN in terms of Recall@K and NDCG@K. This is because it uses maximal M-cliques to augment the bundle-item affiliation network, which alleviates the sparsity issue by more precisely capturing the relationships between bundles and items.

8 RELATED WORKS

This section mainly reviews the existing works of maximal clique enumeration (MCE) in both homogeneous and heterogeneous graphs.

• MCE in homogeneous graphs. As a fundamental problem for graph data analysis, the MCE problem has garnered plenty of research attention. The well-known solutions to the MCE problem are the classic BK algorithm [9] and its pivot-based variants [9, 29, 58, 75]. The BK algorithm [9], proposed by Bron and Kerbosch in 1973, enumerates maximal cliques in a recursive backtracking manner. Tomita et al. [75] showed that the time overhead to the MCE problem is optimal in the worst case using their pivot technique. Eppstein et al. [29] further derived a tighter worst-case time complexity for MCE based on the degeneracy order [52]. Naudé et al. [58] revised the pivot algorithm by refining the pivot selection process. Besides, the I/O efficient algorithms [21, 22], distributed algorithms [86], parallel [27, 67] algorithms, and output-sensitive

algorithms [14, 23, 55] have also been developed for the MCE problem. In addition, many variants of the MCE problem have been formulated for different types of homogeneous graphs, such as temporal [78], signed [20, 49], uncertain networks [24, 50], and efficient solutions have also been studied.

• MCE in heterogeneous graphs. Generally, the heterogeneous graphs can be classified as bipartite graphs and other general HINs [30]. On bipartite graphs, the problem of maximal bi-clique enumeration (MBCE) has received much attention. David Eppstein [28] developed a linear MBCE algorithm for any graph of bounded arboricity. In [65], the MBCE problem is solved by exhaustively listing subsets of vertices in one layer, subsequently identifying vertices in the other layer as their common neighbors, and then identifying the bicliques. Li et al. [47] applied efficient algorithms for mining closed patterns to the MBCE problem. Zhang et al. [93] proposed an algorithm iMBEA by fusing backtracking with a branch-and-bound framework. Recently, several approaches [1, 17] have employed the pivot principle to speed up the MBCE, and a special case of the MBCE problem, called maximum biclique search, has attracted much attention [16, 54, 68]. Besides, MCE on multi-partite graphs has also been studied [60]. For general HINs, Hu et al. [38] introduced the concept of motif-clique and proposed the META algorithm for the MMCE problem. However, META cannot efficiently process large HINs, calling for a faster solution.

In addition, many other works also find maximal solutions, such as maximal frequent itemset [5, 6, 33, 35, 36, 76], maximal independent set [32, 44, 53, 74, 85], maximal k-plex [2, 25, 41, 82], etc.

9 CONCLUSIONS

In this paper, we investigate the problem of efficient maximal motifclique enumeration (MMCE) over large HINs. The existing MMCE algorithm, following the classic BK algorithm, which explores all the possible subgraphs of a maximal motif-clique (M-clique) and checks whether each maximal M-clique has been enumerated at each recursive step, is very time-consuming. To improve the efficiency of MMCE, we introduce an order-based framework and propose a pivot-based pruning strategy. We further propose a series of optimization techniques. Our experimental results on five real large HINs show that our algorithm achieves up to three orders of magnitude faster than the state-of-the-art algorithm. In the future, we will develop a more generic definition of M-clique, that can incorporate various kinds of graph isomorphism, such as subgraph monomorphism [64], supergraph isomorphism [69], and graph homomorphism [63]. We will also study how to efficiently enumerate maximal M-cliques efficiently on large dynamic HINs since many real-world HINs are evolving over time.

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⁷https://music.163.com

⁸http://www.yousuu.com

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