Efficient Maximum (α, β) -Quasi Biclique Computation on Bipartite Graphs*

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Abstract. A bipartite cohesive subgraph refers to a bipartite subgraph in which the vertices are highly interconnected. In this paper, we consider the (α,β) -quasi biclique model, which allows for connection proportions on two sides of vertices to be no less than α and β , respectively. Building upon this model, we propose the maximum (α,β) -quasi biclique search problem, which finds applications in various domains, including abnormal behavior detection, community group search, and biological gene expression detection. Due to the inefficiency of the baseline, we design effective preprocessing methods, upper bounding and reduction techniques, and advanced branching strategies, which collectively constitute our optimized algorithm called MVQB. Finally, through extensive experiments on real-world bipartite graphs, we validate the efficiency improvement of our proposed algorithm MVQB over the baseline, which demonstrates a speed improvement of up to six orders of magnitude.

Keywords: bipartite cohesive graphs \cdot maximum (α, β) -quasi biclique

1 Introduction

Bipartite graphs serve as a robust structure for modeling interactions between two distinct groups of entities [5,17]. This unique property enables these graphs to effectively represent and analyze complex relationships in many real-world scenarios, such as collaboration networks [13], customer-product networks [26], and user-page networks [4]. Furthermore, dense or cohesive subgraphs are particularly valuable in bipartite graphs as they often contain significant information relevant to addressing real-world challenges [10, 20, 29]. These subgraphs, characterized by a high degree of connectivity between their vertices, can reveal patterns and anomalies that are crucial for practical applications. For example, in fraud detection [29], the cohesive bipartite subgraphs help identify unusual groupings of interactions that may indicate fraudulent activity.

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To identify dense subgraphs within bipartite graphs, researchers have proposed a range of cohesive subgraph models. Some of the representatives include bicliques [20], (α, β) -bicores [10], and k-biplexs [24, 29]. These cohesive models serve diverse purposes and impose varying conditions on the connectivity patterns within the subgraphs. For instance, bicliques require complete interconnectivity between the two sides of vertices, while (α, β) -bicores and k-biplexes relax bicliques by requiring that each vertex connects to at least a constant number and disconnects from at most a *constant* number of vertices on the opposite side, respectively. We observe that research on models of bipartite cohesive subgraphs with more relaxed conditions is still insufficient based on the following two aspects. (1) Lack of density: Since these cohesiveness constants are unrelated to the size of cohesive subgraph, the identified cohesive subgraphs with unreasonable cohesiveness constants often exhibit low density, rendering them inadequate for effectively representing a cohesive structure in practice. (2) Incomplete parameter settings: Since bipartite graphs usually represent two types of entities, and these entities may not necessarily be of the same size, it is necessary to control the missing parameters separately for each side. In this paper, we focus on the (α, β) -quasi biclique model, which requires that the connection of each vertex in the subgraph reaches a certain proportion in relation to the size of vertices on the opposite side. Based on this cohesive bipartite model, we investigate the Maximum (α, β) -Quasi Biclique Search (MQBS) problem.

Challenges. The MQBS problem has been proven to be NP-hard [17]. Previous studies [17,28] then focus on either heuristic or approximation methods; however, these approaches cannot be modified to guarantee an exact solution for MQBS. To efficiently find the exact solution, a common approach is to adopt the branch-and-bound framework, similar to that used for other cohesive models mentioned earlier. However, unlike the biclique and k-biplex models, a notable difference of quasi-bicliques is that they are non-hereditary, i.e., an induced subgraph of an (α, β) -quasi biclique is not necessarily an (α, β) -quasi biclique. This implies that we cannot leverage the nice hereditary property to design useful pruning and reduction techniques in the branch-and-bound framework.

Our Solutions. We propose two exact algorithms based on the branch-and-bound framework for solving the MQBS problem. First, we provide an intuitive baseline method called ENUM. This method initially reduces the input graph into a bicore, then adopts a random vertex selection strategy to exhaustively explore all vertex combinations through a simple binary branching method. Although the methodology of ENUM is straightforward, it suffers from poor graph preprocessing, a lack of effective upper bounding and reduction rules, and a naive branching strategy. To address these limitations, we propose our advanced algorithm, termed MVQB. MVQB introduces a more powerful preprocessing method, which effectively utilizes the common neighbor information between two vertices on the same side and graph coloring techniques. Subsequently, we design a series of upper bounding methods and reduction rules, which are useful for pruning unnecessary branches and reducing the candidate sets during enumeration. Finally, to improve the performance of MVQB in branching, we devise an advanced

pivot-based branching strategy, which selects a pivot vertex and conducts specific branching operations based on this pivot. Compared to the simple branching in ENUM, the pivot-based branching effectively leverages the inherent characteristics of the (α, β) -quasi biclique model, thereby improving performance.

Contributions. The contributions of this paper can be summarized as follows.

- We propose exact solutions for the MQBS problem, i.e., a baseline approach called ENUM and an advanced vet efficient method named MVQB.
- We develop a series of novel techniques in MVQB, including refined graph preprocessing techniques, upper bounding and reduction rules, and a novel branching method. These techniques effectively mitigate the inherent limitations of the (α, β) -quasi biclique model, which lacks the hereditary property.
- We conduct extensive experiments between ENUM and MVQB across ten real-world datasets. The results demonstrate that 1) MVQB outperforms ENUM by up to six orders of magnitude in practical efficiency; and 2) the proposed advanced techniques designed for MVQB improve the algorithmic performance.

2 Preliminaries

Let $G=(L\cup R,E)$ be an undirected, unweighted bipartite graph, where L and R are two disjoint sets of vertices, and $E\subseteq L\times R$ represents the set of edges. We denote the total number of vertices in G by |L|+|R|=n and the number of edges by |E|=m, respectively. Given a pair of vertex sets (X,Y) with $X\subseteq L$ and $Y\subseteq R$, we use $G[X\cup Y]=(X\cup Y,E_{X,Y})$ to define a subgraph of G induced by the vertex sets X and Y, where $E_{X,Y}=\{(x,y)\in E\mid x\in X,y\in Y\}$. For a vertex $u\in X$ (resp. $v\in Y$), the neighbor set and degree of u (resp. v) are denoted by $\Gamma_G(u,Y)$ and $\delta_G(u,Y)$ (resp. $\Gamma_G(v,X)$ and $\delta_G(v,X)$), respectively, where $\delta_G(u,Y)=|\Gamma_G(u,Y)|$ (resp. $\delta_G(v,X)=|\Gamma_G(v,X)|$). Particularly, when the context is clear, we use $\Gamma_G(u)$ and $\delta_G(u)$ to denote the neighbor set and degree of a vertex u, respectively. Besides, we define the common neighbors of two vertices u_1 and u_2 from the same side of vertices (i.e., both from L or R) as $\Delta(u_1,u_2)$, i.e., $\Delta(u_1,u_2)=\{w\mid (u_1,w)\in E \text{ and } (u_2,w)\in E\}$.

A classic bipartite subgraph model is called *biclique*, which has complete interconnectivity between the two sides of vertices. Obviously, the biclique model is the most cohesive subgraph model in bipartite graphs. To extend the practical applicability of bicliques, we consider a relaxed model of bicliques.

Definition 1 ((α, β) -Quasi Biclique). Given a bipartite graph $G = (L \cup R, E)$ and two real numbers α, β in [0, 1], an induced subgraph $G([X \cup Y])$ is an (α, β) -quasi biclique (QBC) if and only if every vertex $u \in X$ has a degree no less than $\alpha \cdot |Y|$ and every vertex $v \in Y$ has a degree no less than $\beta \cdot |X|$.

Compared to the biclique model, our proposed QBC model introduces two additional parameters α and β to relax the requirement of full connectivity. Note that when $\alpha=\beta=1$, the QBC model degenerates to the biclique model. Nonetheless, for general cases, QBC differs greatly from the biclique model in that

the former lacks the *hereditary property* inherent to the latter. More specifically, any induced subgraph of a biclique remains a biclique; however, this is not always the case for QBC. We next introduce the problem studied in this paper.

Problem 1. Given a bipartite graph $G = (L \cup R, E)$, two values $\alpha, \beta \in (0.5, 1]$, and thresholds θ_L, θ_R , the Maximum (α, β) -Quasi Biclique Search (MQBS) problem aims to find an (α, β) -quasi biclique $G([X \cup Y])$ with the largest cardinality, where $|X| \geq \theta_L$ and $|Y| \geq \theta_R$.

We note that in the above MQBS problem, we introduce two practical constraints, which are based on the followings. First, without imposing constraints on the parameters α and β , MQBS may yield a disconnected subgraph, which is less useful in practice. On the other hand, if the values of α and β lie within the range (0.5,1], we can can always ensure that the obtained (α,β) -quasi biclique is a connected subgraph, as proved in the below Lemma 1. Second, without imposing constraints on the size of the returned subgraph, MQBS will spend considerable time searching for solutions that are too small to be practically meaningful.

Lemma 1. Given two real numbers α, β in (0.5, 1], any two vertices in the same side of vertices of an (α, β) -quasi biclique have at least one common neighbor.

The omitted proofs can be found in Appendix of the technical report [1]. In this paper, we focus on providing a practically efficient exact algorithm for MQBS.

2.1 A Baseline Solution

A straightforward method to find the maximum (α, β) -quasi biclique is to enumerate the vertex sets and return the one with the largest cardinality. Such a baseline (advanced algorithm introduced in Section 3) follow the branch-and-bound framework, which recursively solves the problem via a process of branching. This process involves dividing the problem of finding the largest (α, β) -quasi biclique into multiple sub-problems. Each sub-problem, or branch, is defined by a unique pair of disjoint vertex sets, $(S_L \cup S_R, C_L \cup C_R)$. Here, the partial set $S_L \cup S_R$ contains vertices that form an (α, β) -quasi biclique, which must be included in any solution within this specific branch. Conversely, the candidate set $C_L \cup C_R$ consists of vertices that may potentially be added to $S_L \cup S_R$ to create a larger solution. The solution of a branch is clearly the largest (α, β) -quasi biclique within that branch. A given (α, β) -quasi biclique is considered part of a branch $(S_L \cup S_R, C_L \cup C_R)$ if it 1) includes all vertices in $S_L \cup S_R$ and 2) is a subgraph of $G[(S_L \cup S_R) \cup (C_L \cup C_R)]$. Thus, solving the branch $(\emptyset, L \cup R)$ is equivalent to finding the maximum (α, β) -quasi biclique in $G[L \cup R]$.

As mentioned in Section 2, an (α, β) -quasi biclique does not enjoy the hereditary property for efficient pruning during the enumeration. In other words, our baseline solution only examines the current vertex set based on the definition of (α, β) -quasi biclique and the size constraint associated with the problem. To speed up the whole enumeration process, we provide a necessary definition for graph preprocessing operations based on the following definition.

Algorithm 1: The baseline method: ENUM

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Input: A bipartite graph G = (L \cup R, E), two real numbers \alpha and \beta in (0.5, 1], and two thresholds \theta_L and \theta_R

Output: The maximum (\alpha, \beta)-quasi biclique

1 B^* \leftarrow \emptyset;

2 Reduce G to its (\lceil \alpha \cdot \theta_R \rceil, \lceil \beta \cdot \theta_L \rceil)-bicore by Lemma 2;

3 Enum_Rec(\emptyset, L \cup R);

4 return G[B^*];

Procedure: Enum_Rec(S_L \cup S_R, C_L \cup C_R)

5 if |S_L \cup S_R \cup C_L \cup C_R| \le |B^*| or |S_L \cup C_L| < \theta_L or |S_R \cup C_R| < \theta_R then

6 \lfloor \text{return};

7 if G[(S_L \cup C_L) \cup (S_R \cup C_R)] is an (\alpha, \beta)-quasi biclique then

8 \lfloor B^* = (S_L \cup C_L) \cup (S_R \cup C_R); return;

9 v \leftarrow \text{Randomly select a vertex from } C_L \cup C_R;

10 Enum_Rec(S_L \cup S_R \cup \{v\}, C_L \cup C_R \setminus \{v\});

11 Enum_Rec(S_L \cup S_R, C_L \cup C_R \setminus \{v\});
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Definition 2 $((\alpha, \beta)$ -Bicore [14]). Given a bipartite graph $G = (L \cup R, E)$ and two integers α and β , an (α, β) -bicore $G([X \cup Y])$ is the maximal induced subgraph of G in which all the vertices in $u \in X$ have degree at least α and all the vertices in $v \in Y$ have degree at least β .

Based on Definition 2, we can easily obtain the following graph preprocessing.

Lemma 2. An (α, β) -quasi biclique $G[X \cup Y]$ is in an $(\lceil \alpha | Y \rceil \rceil, \lceil \beta | X \rceil)$ -bicore.

We summarize our baseline method ENUM in Algorithm 1, where the description is deferred to Appendix of the technical report [1] due to the space limit.

Limitations of the Baseline. Although the idea of ENUM is quite straightforward, it still has several limitations. <u>First</u>, despite some preprocessing operations, the candidate set of ENUM remains large, which affects the subsequent exact search in the branch-and-bound framework. <u>Second</u>, ENUM does not apply any effective upper bound techniques or reduction rules specific to (α, β) -quasi bicliques, preventing it from pruning unnecessary branches and candidate vertices. <u>Third</u>, the branching strategy of ENUM, based on random vertex selection, is simple, leading to numerous search operations during the branching process.

3 Our Advanced Approach

To solve the limitations of ENUM, we introduce our advanced method, MVQB, for the MQBS problem in this section. In particular, we first present a more efficient graph preprocessing technique before conducting the branch-and-bound search in Section 3.1. Then, we introduce effective upper bounding techniques and reduction rules tailored for (α, β) -quasi bicliques in Section 3.2. Finally, Section 3.3 gives a more sophisticated branching strategy based on pivot vertices, along with our advanced algorithm MVQB equipped with all the proposed techniques.

3.1 A More Effective Graph Preprocessing Method

Effective graph preprocessing techniques can further reduce the search space that needs to be explored in the enumeration. However, ENUM only considers the degree of vertices during the graph preprocessing, while overlooking the significance of the relationships between vertex pairs. Thus, we propose a new preprocessing technique based on the number of common neighbors between vertex pairs.

Proposition 1. If there exist two vertices $u_1, u_2 \in L$ (resp. $u_1, u_2 \in R$) such that $\Delta(u_1, u_2) < 2 \lceil \alpha \cdot \theta_R \rceil - \theta_R - 1$ (resp. $2 \lceil \beta \cdot \theta_L \rceil - \theta_L - 1$), then vertices u_1 and u_2 cannot coexist in an (α, β) -quasi biclique with threshold θ_R (resp. θ_L).

By Proposition 1, we can determine whether two vertices on the same side of a bipartite graph can coexist in a solution by calculating the number of common neighbors between them. We next introduce our new preprocessing technique.

Graph Coloring-based Preprocessing Technique. Given a bipartite graph $G = (L \cup R, E)$ and two thresholds θ_L and θ_R , any two vertices u_1 and u_2 on the same side of G can coexist in the same (α, β) -quasi biclique if and only if they do not violate the conditions specified in Proposition 1. Based on this, we formalize the coexistence relationships between vertices on the two sides, L and R, of G by abstracting them into two simple undirected graphs, G_L and G_R , respectively. details as follows. If two vertices u_1 and u_2 on the same side of a bipartite graph can coexist within the same (α, β) -quasi biclique according to Proposition 1, an edge is generated between their corresponding vertices in the corresponding undirected graph, i.e., G_L or G_R . After constructing G_L and G_R , we apply a greedy graph coloring algorithm [23] to the two undirected graphs, assigning each vertex a unique color such that no two adjacent vertices share the same color. For a given vertex v, let C(v) represent the set of unique color numbers assigned to its adjacent vertices. Proposition 2 details our proposed graph coloring-based preprocessing technique.

Proposition 2. Given a bipartite graph $G = (L \cup R, E)$ and two threshold values θ_L and θ_R , we can safely remove a vertex $v \in L$ (resp. $v \in R$) if $C(v) < \theta_L - 1$ (resp. $C(v) < \theta_R - 1$) in the corresponding undirected graph G_L (resp. G_R).

We also present an example for the above technique in Appendix of [1].

3.2 Upper Bounding and Reduction Rules

Effective upper bounding and reduction rules can reduce the search space in the enumeration, thereby improving the practical performance of the branch-and-bound based algorithm. Generally speaking, for models with hereditary properties (such as bicliques), if the selected set in a particular branch no longer constitutes a feasible solution, we can easily prune that branch since any sub-branch containing the selected set cannot yield a feasible solution. However, due to the lack of hereditary property in the (α, β) -quasi biclique model, designing appropriate upper bounding and reduction rules is extremely challenging.

Nevertheless, in this section, we address this challenge by focusing on certain properties of (α, β) -quasi bicliques.

Upper Bounding Rules. The upper bound technique provides an estimation on the size of the optimum solution that can be obtained from a particular branch, and then uses this bound to determine whether the branch can be pruned or not. Next, we introduce an upper bound based on vertex degree, derived from the definition of (α, β) -quasi bicliques.

Lemma 3 (Degree-based Upper Bound). Given a branch $B = (S_L \cup S_R, C_L \cup C_R)$, the degree-based upper bound UB_d is shown as $UB_d = UB_{dl} + UB_{dr}$, where $UB_{dl} = \lfloor \min_{v \in S_L} \delta_G(v, S_R \cup C_R)/\alpha \rfloor$ and $UB_{dr} = \lfloor \min_{v \in S_R} \delta_G(v, S_L \cup C_L)/\beta \rfloor$.

With Lemma 3, we can safely prune the branch if this upper bound is less than the size of the recorded optimum solution. Besides, we also identify an important characteristic of (α, β) -quasi biclique that exhibits monotonicity, which enables the design of another efficient upper bound technique. This characteristic relates to the number of missing neighbors in the selected set of a branch. Specifically, within a given branch $B = (S_L \cup S_R, C_L \cup C_R)$, the number of neighbor vertices absent in S_R (resp. S_L) for a vertex $v \in S_L$ (resp. $v \in S_R$) demonstrates a non-decreasing characteristic during the branching process. Our second upper bounding technique is to calculate the upper bound of missing neighbors allowable for the vertices in the selected set of a branch, and then determine whether this bound has exceeded the current number of missing neighbors in the selected set. We summarize this upper bound technique in Lemma 4.

Lemma 4 (Missing Degree-based Upper Bound). Given a branch $B = (S_L \cup S_R, C_L \cup C_R)$, the missing degree-based upper bounds UB_{ml} and UB_{mr} on the left and right sides of B are shown as $UB_{ml} = \lfloor (1 - \alpha) \cdot UB_{dl} \rfloor$ and $UB_{mr} = \lfloor (1 - \beta) \cdot UB_{dr} \rfloor$.

Lemma 4 only defines the maximum number of missing neighbors allowed for a branch. To unleash its pruning power, we let $\nabla(S_L)$ be the maximum missing degree of vertices on the left side of the selected set, i.e., $\nabla(S_L) = |S_R| - \min_{u \in S_L} \delta_G(u, S_R)$. Note that a similar definition $\nabla(S_R)$ can be defined symmetrically on S_R . Now we can prune a branch if $\nabla(S_L) > UB_{ml}$ or $\nabla(S_R) > UB_{mr}$.

Reduction Rules. If there are vertices in the candidate set that cannot be added to the selected set to generate a larger solution, these vertices can be removed from the candidate set. For ease of presentation, we define the lower bound on the right side as $LB(B_R) = \left\lceil \frac{\nabla(S_L)}{1-\alpha} \right\rceil$. The lower bound on the left side can be defined symmetrically. Subsequently, we propose two novel reduction rules based on the information of degree and common neighbor, respectively.

Lemma 5 (Degree-based Reduction Rule). Given a branch $B = (S_L \cup S_R, C_L \cup C_R)$ and a vertex $u \in C_L$ (resp. $u \in C_R$), u can be removed from the candidate set if $\delta_G(u, S_R \cup C_R) < \alpha \cdot LB(B_R)$ (resp. $\delta_G(u, S_L \cup C_L) < \beta \cdot LB(B_L)$).

Lemma 6 (Common Neighbor-based Reduction Rule). For a branch $B = (S_L \cup S_R, C_L \cup C_R)$ and a vertex $u \in C_L$ (resp. $u \in C_R$), u can be removed from the candidate set if there exists a vertex $v \in S_L$ (resp. $v \in S_R$) such that $\Delta(u, v) < 2 \lceil \alpha \cdot LB(B_R) \rceil - LB(B_R) - 1$ (resp. $\Delta(u, v) < 2 \lceil \alpha \cdot LB(B_L) \rceil - LB(B_L) - 1$).

3.3 Our Advanced Branching Method and the Final Algorithm

To address the limitations of ENUM in the branching strategy, we present our advanced branching method in this part. We also present our advanced algorithm, named MVQB, that integrates all the previously mentioned techniques.

Pivot-based Branching Rules. In the baseline ENUM, the branching strategy involves randomly selecting a vertex from the candidate set as the branching vertex. This arbitrary branching behavior often results in many unnecessary branches, significantly impacting the running time of the branch-and-bound algorithm (see our experiments). Instead of randomly picking a vertex to branch on, our goal is to choose a vertex that is likely to lead to faster pruning of the search space. This can be achieved by identifying a vertex (the pivot one) that is well-connected to the current set of vertices that are already included in the left or right branch. By selecting a pivot vertex with specific properties, the algorithm can potentially reduce the number of redundant branches and prune parts of the search space more effectively, leading to a more efficient exploration. We introduce a sophisticated pivot-based branching rule as follows. We first define a methodology for finding a suitable pivot vertex.

Definition 3 (Pivot Vertex). For a branch $B = (S_L \cup S_R, C_L \cup C_R)$, a vertex w is said to be the *pivot vertex* if w satisfies the following conditions:

- (1) We first look for vertices that satisfy $\delta_G(w, S_R \cup C_R) < \alpha |S_R \cup C_R|$ when $w \in S_L$ or $\delta_G(w, S_L \cup C_L) < \beta |S_L \cup C_L|$ when $w \in S_R$;
- (2) If no vertice meets (1), then we look for vertices that satisfy $\delta_G(w, S_R \cup C_R) < \alpha |S_R \cup C_R| \ (w \in C_L) \text{ or } \delta_G(w, S_L \cup C_L) < \beta |S_L \cup C_L| \ (w \in C_R);$
- (3) If multiple vertices satisfy (1) or (2), then w is the one with the minimum degree among them.

Intuitively, a pivot vertex w that meets the criteria of Definition 3 can always be found. Moreover, depending on whether w resides in $S_L \cup S_R$ or $C_L \cup C_R$, two distinct branching cases are applicable.

Branching Case 1: $w \in S_L \cup S_R$. The purpose of this case is to incorporate a certain number of non-neighboring vertices of w into $S_L \cup S_R$, thus ensuring that any sub-branch containing the selected set cannot yield any valid solution. Without loss of generality, we assume that the pivot vertex $w \in S_L$, but the following description symmetrically applies to $w \in S_R$.

We first define the number of missing neighbors of w in the candidate set as $p = |C_R| - \delta_G(w, C_R)$. Then, we use q to denote the maximum number of non-neighbors of w that can be added to S_R from C_R :

$$q = |(1 - \alpha) \cdot |\delta_G(w, S_R \cup C_R)/\alpha|| - (|S_R| - \delta_G(w, S_R)). \tag{1}$$

The first half of Equation 1 represents the maximum number of non-neighbors that vertex w can additionally include in a valid (α, β) -quasi biclique, while the second half indicates the number of neighbors that w has already lost in the selected set. We remark that p > q since the current graph is not a qualified (α, β) -quasi biclique, otherwise we can terminate the branching operation directly. Upon defining p and q, it becomes apparent that any sub-branch with the selected set can contain at most q vertices in $C_R \setminus \Gamma_G(w, C_R) = \{v_1, v_2, \ldots, v_p\}$. Consequently, the branching strategy for Case 1 results in q + 1 sub-branches. (1) The first sub-branch removes v_1 from C_R ; (2) The i_{th} sub-branch removes v_i from C_R and includes $\{v_1, v_2, \ldots, v_{p}\}$ from C_R and includes $\{v_1, v_2, \ldots, v_{q}\}$ to S_R .

Branching Case 2: $w \in C_L \cup C_R$. The main objective of Case 2 is to move w to the selected set through a simple binary branching operation, thereby making one of the sub-branches satisfies the conditions of Case 1 again. The branching strategy of Case 2 generates the following two sub-branches.

(1) The first sub-branch includes the pivot vertex w to $S_L \cup S_R$, i.e., $B_1 = (S_L \cup S_R \cup \{w\}, C_L \cup C_R \setminus \{w\});$ (2) The second sub-branch removes w from $C_L \cup C_R$, i.e., $B_2 = (S_L \cup S_R, C_L \cup C_R \setminus \{w\}).$

Remark: Our branching method is different from the one for k-biplex in [30]. For k-biplex, we can easily find the upper bound of the non-neighbors (i.e., k), but the upper bound for quasi biclique (i.e., the following q) is not as obvious. To find a tighter q and minimize the search branches, we need to identify the pivot vertex which has the maximum number of non-neighbors that can be added.

Our Advanced Algorithm: MVQB. After establishing the advanced pivot-based branching rule, we are ready to present our algorithm, namely MVQB, which integrates all the proposed techniques, in Algorithm 2. Basically, MVQB majorly consists of two parts: graph preprocessing (Lines 2-4) and recursive branch-andbound search (Line 5). Unlike ENUM, which only reduces the input graph into a bicore based on the given thresholds, MVQB incorporates our advanced preprocessing technique in Line 3. To maximize the effects of graph preprocessing. MVQB also iteratively applies the bicore reduction and the graph coloring-based preprocessing technique until no more vertices can be removed (Lines 2-4). After the graph preprocessing stage, MVQB invokes the procedure MVQB_Rec to exhaustively search for the largest (α, β) -quasi biclique. Specifically, MVQB_Rec can be divided into three sub-stages, which are pruning (Lines 7-11), graph reduction (Line 12), and branching (Lines 13-20), respectively. Pruning conditions include: 1) the size of the current graph is smaller than the size of the recorded optimal solution, or the number of vertices on one side is less than the given threshold (Line 7); 2) the entire branch already forms a valid (α, β) -quasi biclique (Lines 9-10); 3) the degree-based upper bound in the current branch is no greater than the current size lower bound $|B^*|$, or the missing degree on any side in the selected set exceeds the missing degree-based upper bound (Line 11). Once a branch passes these pruning tests, MVQB_Rec applies the reduction rules proposed in Lemma 5 and Lemma 6 to further narrow down the candidate vertex set (Line 12). Following this, MVQB_Rec enters its final sub-stage, i.e., the branch-

Algorithm 2: Our advanced algorithm: MVQB

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Input: A bipartite graph G = (L \cup R, E), two real numbers \alpha and \beta in
               (0.5,1], and two thresholds \theta_L and \theta_R
    Output: The maximum (\alpha, \beta)-quasi biclique
 \mathbf{1} \ B^* \leftarrow \emptyset;
 2 while true do
         Remove vertices that (1) do not belong to (\lceil \alpha \cdot \theta_R \rceil, \lceil \beta \cdot \theta_L \rceil)-bicore by
           Lemma 2; (2) meet the conditions of Proposition 2;
         if no vertices are removed then break;
 5 MVQB_Rec(\emptyset, L \cup R);
 6 return G[B^*];
    Procedure: MVQB_Rec(S_L \cup S_R, C_L \cup C_R)
 7 if S_L \cup S_R \cup C_L \cup C_R \le |B^*| or |S_L \cup C_L| < \theta_L or |S_R \cup C_R| < \theta_R then
    return;
 9 if G[(S_L \cup C_L) \cup (S_R \cup C_R)] is an (\alpha, \beta)-quasi biclique then
     B^* = (S_L \cup C_L) \cup (S_R \cup C_R); return;
11 if UB_d \leq |B^*| or \nabla(S_L) > UB_{ml} or \nabla(S_R) > UB_{mr} then return;
12 Refine C_L \cup C_R by the reduction rules from Lemma 5 and Lemma 6;
13 w \leftarrow Find a pivot vertex according to Definition 3;
14 if w \in S_L \cup S_R then
         Assume w \in S_L; p = |C_R| - \delta_G(w, C_R);
15
         q = \lfloor (1 - \alpha) \cdot \lfloor \min_{v \in S_L} \delta(v, S_R \cup C_R) / \alpha \rfloor \rfloor - (|S_R| - \delta_G(w, S_R));
17
         Create q + 1 sub-branches by Branching Case 1;
18 else
         // Branching Case 2
         MVQB_Rec(S_L \cup S_R \cup \{w\}, C_L \cup C_R \setminus \{w\});
19
         MVQB_Rec(S_L \cup S_R, C_L \cup C_R \setminus \{w\});
20
```

ing stage (Lines 13-20). Unlike the random branching vertex selection strategy of the baseline ENUM, MVQB_Rec uses our proposed pivot-based branching method. That is, MVQB_Rec selects a pivot vertex w according to Definition 3 (Line 13), and divides into two different branching scenarios based on whether w is in the selected set S (Lines 14-17) or the candidate set C (Lines 18-20). If $w \in S$, for ease of description, we assume $w \in S_L$. MVQB_Rec then calculates the number of missing neighbors p regarding w in the candidate set (Line 15) and the maximum number of non-neighbors q that can still be added to the selected set (Line 16). Subsequently, we generate q+1 sub-branches following Case 1 of the proposed pivot-based branching strategy (Line 17). On the other hand, if $w \in C$, a simple binary branching is required as in our Branching Case 2 (Lines 19-20).

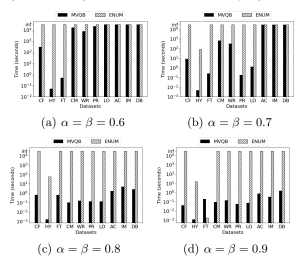
4 Experiments

We conduct extensive experimental studies by comparing the baseline algorithm Enum (Section 2.1) and our advanced algorithm MVQB (Section 3). All algorithms

Abbreviation Dataset |L||R|density θ^* m d_L d_R Cfat 200 200 1.534 3.84×10^{-3} 16 16 3 CF 5.70×10^{-1} ΗY Hywikibooks 242 1,714 2,379 683 17 1,508 1.14×10^{-2} FilmTrust 2.071 35,494 FT 244 1.044 46 2.00×10^{-2} $_{\mathrm{CM}}$ 16,726 22,015 58,595 CondMat 106 18 3.50×10^{-3} Writer WR 89,356 46,213 144,340 42 246 3.10×10^{-3} 30 Producer 48,833 138,844 207,268 512 3.20×10^{-3} Location LO 172,091 53,407 293,687 28 12,189 3.00×10^{-3} Actor AC 127,823 383,640 1,470,404 294 646 1.00×10^{-3} **IMDB** IM 428,439 896,308 3,782,463 1.334 1.590 26 2.00×10^{-4} Dblp DB 1.425.813 4,000,150 8,649,016 119

Table 1: Statistics of the datasets

Fig. 1: Running time in seconds when varying α and β



are written in C++, compiled using g++ with -O3, and executed on a Linux server equipped with an Intel(R) Core(TM) i5-10500 CPU. The running time of each experiment is limited to 30,000 seconds. Our codes can be found in [1]. **Datasets.** Our experiments select 10 real-world datasets from the KONECT Project⁴. Table 1 describes the statistics of the datasets, where the *density* is defined as $m/(|L| \cdot |R|)$, and d_L and d_R quantify the maximum degrees of vertices in vertex sets L and R, respectively. Moreover, the default threshold θ^* is set for different graphs. Especially, we set the $\alpha = \beta = 0.8$ as the default ratio.

4.1 Against the Baseline

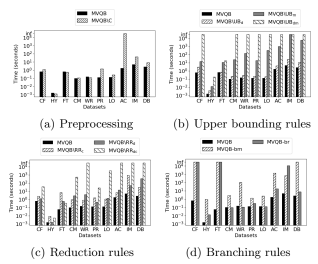
Exp-1: Efficiency Evaluation of Varying Values of α and β . In this experiment, we compare MVQB with ENUM by varying values of α and β , where the results are shown in Figure 1. Overall, we find that MVQB demonstrates superior practical performance compared to ENUM across all different values of α and β , with this advantage being more obvious at larger values of α and β (e.g., 0.8 and 0.9). The advantage of MVQB over ENUM is on the order of five magnitudes. This advantage is primarily due to MVQB's superior graph preprocessing techniques,

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Table 2: R	.ummme	иние	111	SECORGS	wnen	valiving	uniesi	IOIUS -

Algorithm		Е	NUM		MVQB			
Threshold	$\theta^* - 1$	θ^*	$\theta^* + 1$	$\theta^* + 2$	$\theta^* - 1$	θ^*	$\theta^* + 1$	$\theta^* + 2$
CF	OOT	TOO	OOT	OOT	1.19	0.65	0.08	0.07
HY	OOT	57.30	0.01	0.01	0.06	0.01	0.01	0.01
FT	OOT	TOO	OOT	1027.00	0.75	0.66	0.09	0.04
CM	OOT	TOO	OOT	OOT	14.80	0.10	0.05	0.01
WR	OOT	TOO	OOT	OOT	8.75	0.15	0.07	0.04
PR	OOT	TOO	OOT	0.03	1.98	0.13	0.06	0.05
LO	OOT	TOO	OOT	0.04	0.30	0.13	0.07	0.07
AC	OOT	TOO	OOT	OOT	29.40	1.73	0.93	0.91
IM	OOT	OOT	OOT	OOT	4.01	3.56	0.46	0.31
DB	ООТ	OOT	OOT	OOT	236.00	2.58	1.97	1.16

Fig. 2: Ablation studies of the proposed techniques in our algorithm MVQB



upper bounds, reduction rules, and branching strategies. Both algorithms perform undesirable on certain datasets for α and β values of 0.6 and 0.7, as smaller values of α and β indicate an increased difficulty in solving MQBS.

Exp-2: Efficiency Evaluation of Varying Threshold Values. We next compare the two algorithms across different values of thresholds in Table 2. Similarly, we observe that our proposed advanced method, MVQB, significantly outperforms the baseline algorithm, ENUM, across all different values of thresholds, achieving a lead of nearly six magnitudes in some cases. For instance, in the CM dataset with the threshold $\theta = \theta^* + 2$, ENUM fails to find the optimum solution within the time limit, while our MVQB completes the task in only 0.01 seconds. Furthermore, as the threshold values increase, both algorithms show clear improvements in running time, which can be due to the improved graph preprocessing and pruning effects provided by higher threshold values.

4.2 Ablation Studies

To further assess the effectiveness of our proposed advanced techniques adopted in MVQB, we conduct the following ablation studies.

Exp-3: Efficiency Evaluation of Preprocessing Technique. In this experiment, we validate the contribution of the graph coloring-based preprocessing technique to the efficiency improvement of MVQB (and defer the detailed preprocessing information, i.e., the size of the remaining graph after preprocessing, to Appendix of [1]). Our algorithms used in this experiment are MVQB and MVQB\C, where the latter is a variant of Algorithm 2 without the graph coloring-based preprocessing technique. The results are presented in Figure 3(a). We find that the total running time of MVQB is less than that of MVQB\C across most datasets. In particular, on the AC dataset, MVQB shows an efficiency advantage over MVQB\C that exceeds four magnitudes. This clearly demonstrates the effectiveness of our advanced preprocessing technique.

Exp-4: Efficiency Evaluation of Various Upper Bounds. We also implement multiple versions to thoroughly validate the effectiveness of our proposed upper bounding rules, namely MVQB, MVQB\ UB_d , MVQB\ UB_m , and MVQB\ UB_{dm} . Specifically, (1) MVQB\ UB_d denotes Algorithm 2 without using the degree-based upper bound (Lemma 3), (2) MVQB\ UB_m refers to Algorithm 2 excluding the missing degree-based upper bound (Lemma 4), and (3) MVQB\ UB_{dm} indicates that Algorithm 2 is implemented without using either of the upper bounding rules. The results are shown in Figure 3(b). Overall, our proposed two upper bounding techniques significantly enhance the runtime efficiency of MVQB. Without using either upper bound, namely MVQB\ UB_{dm} , eight out of ten datasets fail to find the optimum solution within the time limit. Furthermore, experimental results clearly suggest that the missing degree-based upper bound performs substantially better than the degree-based upper bound, as $MVQB \setminus UB_d$ outperforms $MVQB \setminus UB_m$ in the vast majority of datasets. Nevertheless, the degree-based upper bound still further improves the effectiveness of the missing degree-based upper bound across nearly all datasets.

Exp-5: Efficiency Evaluation of Various Reduction Rules. We implement several variants of our algorithm to assess the effectiveness of the proposed reduction rules, which include MVQB, MVQB\ RR_d , MVQB\ RR_c , and MVQB\ RR_{dc} . Specifically, (1) MVQB\ RR_d represents Algorithm 2 without the degree-based reduction rule (Lemma 5), (2) MVQB \R_c represents Algorithm 2 omitting the common neighbor-based reduction rule (Lemma 6), and (3) MVQB\ RR_{dc} is the implementation of Algorithm 2 without both reduction rules. As shown in Figure 3(c), the two reduction rules we propose make a significant contribution to the runtime efficiency of MVQB, since the performance of MVQB is superior to that of MVQB \RR_d and MVQB \R_c across all datasets. More specifically, our common neighbor-based reduction technique MVQB \RR_c yields a relatively greater performance improvement for the MVQB algorithm compared to the degree-based reduction rule. For instance, in the IM dataset, the running time of MVQB\ RR_d is 86 seconds, while the running time of MVQB\ RR_c reaches 1,000 seconds. This clearly demonstrates that the common neighbor reduction rule can greatly reduce the search space of the enumeration in certain scenarios.

Exp-6: Efficiency Evaluation of Varying Branching Rules. To validate the effectiveness of the proposed pivot-based branching rule, we use the following

three algorithms with different branching strategies, which are MVQB, MVQB-bm, and MVQB-br, respectively. Specifically, (1) MVQB-bm represents the method where Algorithm 2 branches by selecting the vertex with the *minimum degree* in the candidate set, and (2) MVQB-br is the approach in which Algorithm 2 uses the same *random* branching vertex selection approach as the baseline ENUM. The performance of these variants is illustrated in Figure 3(d). From the figure, it is clear that our proposed pivot-based branching method outperforms the other two branching strategies in the vast majority of cases. For instance, in the CH and FT datasets, MVQB-bm and MVQB-br fail to find the optimum solution, whereas our pivot-based branching method identifies the optimum solution in 0.6 second. These results clearly demonstrate the superiority of our advanced branching method.

5 Related Work

Quasi Biclique. James et al. [2] introduced the definition of γ -quasi bicliques, which requires the number of edges is at least a certain proportion γ of the total number of edges possible between its two vertex sets. Building on this concept, Dmitry et al. [12] developed a Mixed Integer Programming approach to search for the largest one. Mishara et al. [21] considered the vertex connectivity, introducing the concept of ϵ -quasi bicliques for studying cases where the degree of connectivity on one side does not exceed a certain missing proportion ϵ on the other side. On this basis, Liu et al. [15] designed a scalable algorithm using Giraph. However, this model, due to imposing constraints only on vertices on one side, could generate unbalanced subgraphs. To address this issue, Liu et al. [17] proposed the concept of δ -quasi bicliques, requiring the connectivity ratio between each vertex in one vertex set and those in the other set to not fall below δ . They proved that finding the largest δ -quasi biclique is an NPhard problem and introduced a heuristic algorithm to tackle this issue. Later, Wang [28] provided an approximation algorithm while violating δ by a bounded ratio. Chang et al. [5] extended this model to weighted bipartite graphs and utilized integer programming to solve the problem.

Other Cohesive Models in Bipartite Graphs. Eppstein et al. [11] proved that enumerating maximal bicliques cannot be done in polynomial time, which has inspired researchers to design more efficient algorithms [3,6]. Furthermore, there are studies that tackle the maximum biclique search problem, characterized by the vertex variant [9] and the edge variant [20, 22, 25, 27], depending on the definition of "maximum" in terms of the total number of vertices and edges, respectively. For the (α, β) -cores, Ding et al. [10] introduced an index structure to study the online version, and Liu et al. [14] reduced the space complexity to O(m). Liu et al. [16] studied the problem in a distributed fashion, while Luo et al. [19] considered the dynamic version of the problem. Zou [32] introduced the k-bitruss model, which requires each edge to be part of at least k "butterflies". Another related model is called the k-biplex, where each vertex in a k-biplex is allowed to disconnect at most k vertices in the other partition of vertex set.

Based on the k-biplex, the problems of maximum k-biplex search [18, 29] and maximal k-biplex enumeration [7, 8, 30, 31] are extensively studied.

6 Conclusion

In this paper, we elaborated on the (α, β) -quasi biclique model and explore the problem of finding the maximum (α, β) -quasi biclique. We proposed an exact algorithm MVQB for the problem, which incorporates efficient preprocessing methods, effective upper bounding and reduction techniques, and advanced branching strategies. Finally, the experimental comparisons on real-world datasets confirm the superior performance of MVQB over the baseline method by up to 6 orders of magnitude faster, achieving significant improvement. In the future, we plan to design the parallel version of our proposed MVQB algorithm.

Acknowledgments

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