Privacy Preserving Subgraph Matching on Large Graphs in Cloud

Zhao Chang∗, Lei Zou†, Feifei Li‡

∗Peking University, China; †University of Utah, USA;
{changzhao, zoulei}@pku.edu.cn; {zchang, lifeifei}@cs.utah.edu

ABSTRACT
The wide presence of large graph data and the increasing popularity of storing data in the cloud drive the need for graph query processing on a remote cloud. But a fundamental challenge is to process user queries without compromising sensitive information. This work focuses on privacy preserving subgraph matching in a cloud server. The goal is to minimize the overhead on both cloud and client sides for subgraph matching, without compromising users’ sensitive information. To that end, we transform an original graph G into a privacy preserving graph Gk, which meets the requirement of an existing privacy model known as k-automorphism. By making use of the symmetry in a k-automorphic graph, a subgraph matching query can be efficiently answered using a graph Gk, a small subset of G. This approach saves both space and query cost in the cloud server. We also anonymize the query graphs to protect their label information using label generalization technique. To reduce the search space for a subgraph matching query, we propose a cost model to select the more effective label combinations. The effectiveness and efficiency of our method are demonstrated through extensive experimental results on real datasets.

1. INTRODUCTION
Owing to the rich semantic and structure information represented by a graph, structured graph data is used in numerous applications, such as social networks, web graphs, biological networks, transportation networks, knowledge base, and RDF graphs. Many emerging applications rely on large graphs to satisfy their query needs, such as Google’s knowledge graph and Facebook’s graph search. Thus, graph data management has attracted significant attention, and efficient graph query processing on large graph is an important subject of study. In this paper, we focus on subgraph matching query [25, 18, 12], which is a key component of numerous applications. For example, answering SPARQL query Q is equal to finding subgraph matches of Q on an RDF graph G [15]. Some graph databases also provide query language that is based on subgraph matching semantics, such as Cypher in Neo4j.

Meanwhile, increasingly, companies choose the “cloud” as their IT infrastructure platform. Using the cloud allows users to avoid expensive upfront infrastructure costs, and focus on projects that differentiate their businesses instead of on infrastructure [2]. Many public cloud services are available, such as Amazon cloud and Microsoft Azure. Some graph systems (such as GraphLab [14] and Neo4j [1]) offer SaaS (software-as-a-service)-style cloud services. In other words, they allow users to upload their graph data to their cloud platforms and provide cloud-based computing services over the outsourced graph data.

However, while utilizing cloud services for building graph applications is a cost-effective solution, the potential risk of compromising sensitive information is a serious problem.

One serious privacy leakage is the “identity disclosure” problem [20, 13]. Assume that an adversary can locate the target entity t as a vertex v of a social network graph G with a high probability. We say that the identity of t is disclosed. A naïve anonymization solution is to remove all identifiable personal information before publishing the network, such as names and social security numbers. However, even when a network is published without any identity information, it is still possible to locate the target with a high probability based on some structural information around the target [10].

For example, if an adversary knows the local graph structure (such as degree, 1-neighbor graph) around the target t, he/she may issue a subgraph query representing the local graph structure to find the matching position. If there are only few matches in graph G, the target will be identified with a high probability. Once it is determined that a vertex v corresponds to the target t, all sensitive attributes associated with v will be compromised. These are called structural attacks; see details in [13, 24, 10].

To address these threats, many privacy preserving graph data publishing techniques have been proposed [26, 6, 22]. A typical solution to protect graph privacy is based on the symmetry of the released graph data [26, 6, 22]. Specifically, given a graph G, we transform G into a k-automorphic graph Gk by introducing some noise edges, where each vertex has at least (k − 1) other symmetric vertices. It means there are no structural differences between v and each of its (k − 1) symmetric vertices. Thus it is impossible to distinguish v from the other (k − 1) symmetric vertices. In other words, the k-automorphism strategy [26] can defend against any structure-based attack.

These privacy preserving graph data publishing techniques seem like perfect solutions to our problem, but they compromise the utility of query results. In particular, they may return false positives regarding subgraph matching queries, due to the noise edges and vertices that were added. However, in a cloud setting, it is possible to offload most query processing costs to the cloud server and ask the clients to execute a simple and efficient filtering step to remove false positives and find the exact answers.

EXAMPLE 1. Consider a professional social network in Figure 1 that is modeled as a graph G. Each vertex in G represents an
entity, such as an individual (p), a company (c), or a school (s). Each edge in G represents a relation between two entities, such as "spouse" relation, "work at" relation and "graduate from" relation. Each entity has some attributes. For example, the attributes associated with individuals are "gender", "occupation" and so on. A user wants to find "two individuals satisfying (1) both of them graduated from the same university located in Illinois, and (2) one of them is working at a software company and the other one is working at an Internet company". The user issues a subgraph pattern matching query Q over G, as shown in Figure 1.

Sending the original graph G to the cloud will leak user privacy in G. Thus, we resort to the privacy preserving graph publishing techniques. For example, we can use the k-automorphism model [25]; the k-automorphic graph Gk for G and k = 2 is shown in Figure 2, where noise edges are shown by red dashed lines. Clearly, Gk protects the structure privacy of the graph G. To protect label privacy, we also use label anonymization where each vertex label in Gk is replaced by a label group (i.e., a generalized label). We use Label Correspondence Table (LCT) (in Figure 2) to represent the mapping between label groups and vertex labels. For example, vertex p1 in Gk has label groups C and E in Figure 2.

A straightforward method is to upload G2 to the cloud and perform graph queries over G2. Although the approach does not leak any sensitive information, the answers to query Q over graph G(L(Q,G2)) are different from those over G (L(Q,G)). For example, there are only two subgraph matches of Q over the original graph G. However, due to the noise edges introduced into the k-automorphic graph Gk and the label anonymization, there are eight matches over Gk. Nevertheless, when the cloud returns R(Q,Gk) with eight matches, the client can efficiently filter out false positives based on G to derive R(Q,G).

Note that the filtering step at the client side is much cheaper than asking the client to run the subgraph matching query Q over G (which is known to be expensive especially over large graphs). In other words, the cloud server did most of the work in this process, making the cloud service worthwhile and attractive for the client. We do assume that the client is the data owner who has access to G for the filtering step. For the general case, a query client (who is trusted and authenticated by the data owner) can ask for the data owner to execute the (very lightweight) filtering step.

However, the example above shows some major limitations of directly applying the existing graph data publishing techniques [25, 26, 22]. Firstly, most of these techniques that focus on structural attacks do not protect label privacy at the same time. But more importantly, to achieve higher privacy, they need to add a lot of noise edges and/or vertices (e.g., a large k value in the case of the k-automorphism model) to the original graph G. This results in a much larger graph (than the original graph) on the cloud side, which leads to much more expensive storage cost, much larger communication overhead, and much higher query costs for both the cloud server and the query client.

This work focuses on reducing these overheads without compromising either data and query graphs' privacy or the correctness of the final query results.

Solution overview. In order to achieve that, we first propose a basic solution. We transform the data graph G into an outsourced graph G2 using existing privacy-preserving graph publication techniques, such as the k-automorphism model [25]; and then upload G2 to the cloud. At query time, we propose a two-phase query evaluation strategy. First, given a query graph Q, we transform Q into an outsourced query graph Q'. In the cloud, we evaluate subgraph graph Q' over G2 to obtain query results R(Q', G2). Then, in the client side, we filter out false positives in R(Q', G2) based on G to find the final, correct results R(Q, G), i.e., subgraph matches of Q over the original graph G. To improve the query performance, we propose a solution that only uploads a small part of G to the cloud by leveraging the symmetry of the k-automorphic graph Gk. This method saves both space cost and query processing time in the cloud significantly. Although only a (small) part of G is uploaded to the cloud, our method still guarantees the correctness of the query results. Furthermore, we also propose a cost model based label combination/generalization strategy to reduce the search space for subgraph matching queries in the cloud.

Contributions. To the best of our knowledge, this is the first work that supports privacy preserving subgraph matching queries over a large graph in the cloud while protecting privacy without undermining query results. In this paper, we consider the cloud server "honest-but-curious", which is consistent with most related works in the literature. In other words, the cloud server always offers correct computations without cheating. However, the cloud server is "curious" to learn the graph data, its index structure, and user queries so as to gain sensitive information if he can. Our main contributions are summarized as follows.

- We propose an effective strategy to provide exact subgraph matching query services in public cloud while preserving private information in the data graph and query graphs.
- To save both space cost and query processing cost, we only upload a small subset of the anonymized data graph to the cloud. We answer subgraph matching queries efficiently in the cloud by making use of the symmetry of the k-automorphic graph. These techniques contribute to saving cost while minimizing the overhead in the client side.
- In order to reduce the search space in answering subgraph matching queries, we design a novel cost model to select effective vertex label combinations for anonymizing labels in the data graph and query graphs.
- We study the effectiveness and efficiency of our method through extensive experiments over several large real graphs.
Table 1: Notations

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>G</td>
<td>The original data graph</td>
</tr>
<tr>
<td>G²</td>
<td>The data graph released by k-automorphism algorithm</td>
</tr>
<tr>
<td>G³</td>
<td>The outsourced data graph</td>
</tr>
<tr>
<td>Q</td>
<td>The original query graph</td>
</tr>
<tr>
<td>Qo</td>
<td>The outsourced query graph</td>
</tr>
<tr>
<td>Q</td>
<td>The outsourced query graph</td>
</tr>
<tr>
<td>S, k</td>
<td>The k-th star of G generated by query decomposition</td>
</tr>
<tr>
<td>R(Q,G)</td>
<td>The set of candidate matching results of Q over G</td>
</tr>
<tr>
<td>R(G)</td>
<td>The set of subgraph matches of Q on G</td>
</tr>
</tbody>
</table>

2. BACKGROUND

2.1 Preliminary

Traditional subgraph matching methods always adopt the vertex labeled graph model [23], where each vertex has a single label. But this model is not desirable to model complex graph data, such as social networks and RDF graphs. In this work, we adopt the attributed graph model, where each vertex has a rich data structure including vertex type, vertex attributes and vertex labels (i.e., attribute values). For simplicity, we only consider rich data structures on vertices and ignore those on edges, although handling the more general case is not more complicated. For example, we can introduce an imaginary vertex to represent an edge of interest and assign the rich data structure on the edge to the new vertex. Formally, our graph model is defined as follows. Table 1 lists some frequently-used notations in this paper.

Definition 1. Attributed Graph Model. An attributed graph is defined as \( G = (V(G), E(G), T, \Gamma, L) \), where (1) \( V(G) \) is a set of vertices; (2) \( E(G) \subseteq V(G) \times V(G) \) is a set of undirected edges; (3) \( T \) is a set of vertex types, where each vertex has and only has one vertex type; (4) \( \Gamma \) is a set of vertex attributes, where each vertex type has one or more vertex attributes and different vertex types have different vertex attributes; and (5) \( L \) is a set of vertex labels, where each vertex attribute has one or more vertex labels.

The vertex type, vertex attributes and vertex labels of vertex \( v \) are denoted as \( T(v), \Gamma(v), L(v) \), respectively. For any two different vertices \( v_1 \) and \( v_2 \), if and only if \( T(v_1) = T(v_2) \), then \( \Gamma(v_1) = \Gamma(v_2) \). For one vertex attribute \( A \in \Gamma(v) \), \( (A, \{a_1, \cdots, a_n\}) \) represents the vertex attribute values on attribute \( A \) which are the vertex labels of \( A \). Note that one vertex attribute may have multiple vertex labels (i.e., attribute values). For example, the “Company Type” of one company can be “Internet”, “Software”, and so on.

The data graph \( G \) and the query graphs \( Q \) in the running example follow the attributed graph model.

Definition 2. Subgraph Match. Given a data graph \( G = (V(G), E(G), T, \Gamma, L) \) and a query graph \( Q = (V(Q), E(Q), T, \Gamma, L) \), \( Q \) is subgraph isomorphic to \( G \), if and only if there exists at least one injective function \( \gamma : V(Q) \rightarrow V(G) \) such that

1) \( \forall q \in V(Q), \gamma(q) \in V(G) \Rightarrow L(q) \subseteq L(\gamma(q)) \); and
2) \( \forall q, q_\prime \in V(Q), \text{edge } \gamma(q)\gamma(q_\prime) \in E(Q) \Rightarrow \text{edge } \gamma(q)\gamma(q_\prime) \in E(G) \).

Figure 2: A k-automorphic graph and an anonymized query graph on the same professional social network graph for \( k = 2 \).

Table 2: A k-automorphic graph and an anonymized query graph on the same professional social network graph for \( k = 2 \).

2.2 K-Automorphism

As explained in Section 1 (Example 1 and its discussion), it is possible to apply existing privacy preserving graph publication techniques in constructing a baseline solution. However, they suffer from major efficiency limitations. Furthermore, most of these techniques that focus on structural attacks do not protect label privacy at the same time. Nevertheless, we can use these techniques as a preprocessing step (e.g., k-automorphism [26]) towards constructing more effective and efficient solutions. That said, we will briefly review k-automorphism [26] next.

The basic idea of k-automorphism is as follows. Given a graph \( G \) to be published, we transform it into \( G^2 \) by introducing more vertices and edges, where \( G^2 \) satisfies the k-automorphic graph model (Definition 3). It means that any vertex \( v \in G^2 \) cannot distinguish itself from each of its symmetric \( (k - 1) \) vertices in \( G^2 \). Therefore, \( G^2 \) is safe to publish. Here, it is assumed that the graph is unlabeled. Figure 3 shows an example of the algorithm in [26].

Definition 3. K-Automorphic Graph [26]. A k-automorphic graph \( G^k \) is defined as \( G^k = (V(G^k), E(G^k)) \), where \( V(G^k) \) can be divided into \( k \) blocks and each block has \( \lceil |V(G)|/k \rceil \) vertices. Any vertex \( v \) has \( k - 1 \) symmetric vertices \( v' \) in the other \( k - 1 \) blocks.

The k-automorphic graph is generated with the help of the k-automorphic function defined as follows.

Definition 4. K-Automorphic Function [26]. Given a vertex \( v \) in a k-automorphic graph \( G^k \), \( v \) and its corresponding \( k - 1 \) symmetric vertices form an alignment vertex instance (AVI).

"Symmetric" means swapping \( v \) and \( v' \) gives an isomorphic graph.
Go graph solution works as follows. We transform a VI in AVT is represented as a circularly-linked list. Specifically, symmetric vertices. For example, the first row of AVT contain-

Table (AVT) is given in Figure 4(a). Each row of AVT shows alignment on k blocks (adding edge (p_i, p_j) and edge (p_j, p_k) in Figure 3(b)), where B_1 and B_2 are isomorphic to each other. Lastly, the “edge copy” technique is used to deal with the crossing edges between different blocks (adding edge (p_2, s_2) that corresponds to edge (p_3, s_3) in Figure 3(c)). Readers can refer to [26] for further details. Furthermore, it is easy to show that any vertex in the k-automorphic graph G cannot distinguish itself from each of its (k−1) symmetric vertices. It means that any adversary cannot identify any target vertex with a probability higher than \( \frac{1}{k} \).

2.3 Our Framework and Analysis Overview

In order to provide the correct cloud-based subgraph query ser-

vices while preserving privacy in data graph and query graphs, our solution works as follows. We transform G into an outsourced graph G^o and upload G^o to the cloud. We guarantee that G^o does not leak any private information in the original graph G. Specifically, we require that any adversary, who sees G^o, cannot identify any target vertex in graph G with a probability higher than \( \frac{1}{k} \). We adopt the k-automorphism model [26] to achieve this objective, by generating G^o based on G^o. At query time, given a query graph Q, we transform Q into an outsourced query graph Q^o. Leveraging the symmetry of the k-automorphic graph, the cloud server answers subgraph matching query Q^o based on G^o and returns R_m (a small subset of R(Q^o, G^o)), see Section 4.2.1) to the client. Based on R_m, the client can efficiently recover R(Q^o, G^o) and finally find the final results R(Q, G). We highlight several key requirements.

Firstly, we need to define what sensitive information is in da-

ta graph and query graphs. We do not consider the vertex types T and vertex attributes \( \Gamma \) as sensitive information, since they do not compromise users’ privacy. For any vertex v in G or Q, we consider vertex labels L(v) (i.e., attribute values) as private infor-

thus. The “label privacy” considered in this paper refers to the privacy of the attribute values of each vertex in the graph. For example, given an “individual” vertex in graph G in Figure 1, the values of gender and occupation are users’ privacy. Subgraph matching queries will reveal vertex labels in original data graph G, if we do not anonymize the vertex labels (i.e., attribute values) in query graphs. Thus, the vertex labels in query graph Q are also considered as sensitive information.

Secondly, we must reduce expenses for the cloud server; that is to say, we need to save query time and storage space in the cloud and reduce communication overhead between the client and the cloud. To protect data privacy, we have to introduce noise edges into the data graph and anonymize attribute values of vertices. Doing so will lead to larger search space in subgraph matching. Therefore, in Section 5, we propose a cost model to guide how to select a good graph anonymization strategy. We also study how to reduce the storage space and communication overhead by leveraging the symmetry of the anonymized graph in Section 4, which also helps reduce the cloud’s query processing cost.

Thirdly, we should minimize the overhead in the client side. Sub-

graph matching is an expensive task in terms of both time and space complexity. In our framework, the client only needs to filter out false positives using a hash index in linear time complexity (in terms of the number of candidate results generated by the cloud).

3. A BASELINE SOLUTION

We first describe a simple scheme to help illustrate the basic principle of our approach. For a graph G, we first generate the k-automorphic graph G^o. A baseline solution is to upload G^o to the cloud directly. Note that k-automorphism [26] assumes that the graph is unlabeled. To protect the vertex labels in data graph and query graphs, we resort to the generalization technique, which is also used in k-anonymity [17, 19]. Specifically, each vertex label in data graph and query graphs is represented by a generalized label. We refer to a generalized vertex label as a “label group” in the following discussion. Here, we assume that each label group contains not less than \( \theta \) distinct labels, where \( \theta \) is a user-specified parameter. For example, a label group “A” in Figure 2 includes \( \theta = 2 \) labels (Internet, Software).

The Label Correspondence Table (LCT) (in Figure 2) shows all label groups in the running example. Obviously, the label generalization strategy will affect the query performance. We assume that the label groups are given until Section 5, where we will propose a cost model to find a good label generalization strategy.

Using label generalization on a data graph G, we obtain a graph G^o whose vertices hide vertex labels by label groups. Next, using the k-automorphism algorithm in [26], we obtain a k-automorphic data graph G^o and the corresponding AVT. Each vertex v in G^o has

2The proof was given in Theorem 4.4 of [26].
4.1 Outsourced Graph

executing the very expensive subgraph matching query herself.

an outsourced graph of G is defined as Go

increases.

Theorem 1. For graph G and query graph Q, R(Q,G) \subseteq R(Q',G').

Finally, R(Q',G') are sent to the client and the client filters out
false positives in R(Q',G') based on G to obtain R(Q,G).

The baseline solution suffers from the following limitations. First,
we need to upload the k-automorphic graph Gk to the cloud. Since
the size of Gk can be significantly larger than |G|, this introduces
communication overhead and higher storage cost. Secondly,
a larger graph Gk naturally leads to a larger search space for subgraph
matching, which leads to higher query cost, especially as k increases.

Nevertheless, the baseline solution already saves the client from
executing the very expensive subgraph matching query herself.

4. THE OPTIMIZED METHOD

4.1 Outsourced Graph

According to Definition 3, Gk is a k-automorphic graph consisting
of k blocks. Each vertex v in Gk has (k−1) symmetric vertices in
the other (k−1) blocks. Intuitively, we only need to upload a block
of Gk together with k automorphic functions F, (i = 1, ..., k−1) to
the cloud, since the cloud can recover the whole Gk based on
Gk and the functions F. This is the motivation of defining the out-
sourced graph Gk (Definition 5), which is exactly the first block of
Gk together with the 1-hop neighbors in Gk.

Definition 5. Outsourced Graph. For a graph G(V(G), E(G),
T, Γ, L) and its k-automorphic graph Gk(V(Gk), E(Gk), T, Γ, L),
an outsourced graph of G is defined as Gk = (V(Gk), E(Gk), T, Γ, L),
where (1) Gk follows the attributed graph model; (2) V(Gk) is
the union of vertices in the first block of Gk, denoted as V(B1),

Figure 5: The outsourced data graph G' for G in Example 1.

(k−1) symmetric vertices F, (i = 1, ..., k−1). The k vertices, i.e., v together with F(v) for i = 1, ..., k−1, form a symmetric vertex group, which corresponds to a row in AVT (an AVI). For each symmetric vertex group, we ensure that all vertices in it have the same label groups, i.e., a vertex v's label groups are L(v) = L(F(v)) ∪ ... ∪ L(F(k−1)(v)).

Example 2. Given the original data graph G in Figure 1, the
k-automorphic graph Gk together with its label correspondence table (LCT) is given in Figure 2. A naïve solution is to upload Gk to the cloud directly.

Given a query graph Q, we anonymize Q by representing each vertex label using its corresponding label group. The anonymized query graph is denoted as Q , which is submitted to the cloud. The cloud server answers subgraph query Q k over the k-automorphic graph Gk. We know that R(Q,G) \subseteq R(Q',G'). Intuitively, we introduce more edges and vertices into G to form G', and Q' and G' s vertex labels are anonymized using the same LCT. Formally,

Theorem 1. For graph G and query graph Q, R(Q,G) \subseteq R(Q',G').

Finally, R(Q',G') are sent to the client and the client filters out
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Nevertheless, the baseline solution already saves the client from
executing the very expensive subgraph matching query herself.

4.2 Privacy Preserving Subgraph Query

Given a query Q at the client side, we generalize its vertex labels
to form Q'. Specifically, for each vertex label in Q, we replace it
with the corresponding label group according to the LCT (an example
is shown in Figure 2). Q' has the same size as Q, although each vertex
of Q' has a generalized label group. Q' ensures the privacy of the vertex label information in Q. It is a trade-off between the query
privacy and the query performance. Thus, label generalization
is an interesting challenge in its own and we assume for now
that this is done and present the details in Section 5.

The client then sends Q' to the cloud. The cloud first finds sub-
graph matches of all basic units of Q' over Gk, which will be
defined shortly. Using the symmetry of the k-automorphic graph, the
cloud can then obtain R(Q',G'), but without Gk since it only has
G', by joining these intermediate results. Lastly, R(Q',G') is sen-
to the client, who can obtain R(Q,G) by pruning the false
positives in R(Q',G'). Theorem 1 guarantees the correctness of
this framework.

4.2.1 Processing in the Cloud

A k-automorphic graph Gk consists of k symmetric blocks (B1,
..., Bk), while the outsourced graph Gk contains only one block
together with the first-hop neighbors of its boundary vertices. The challenge is how to find subgraph matches crossing multiple blocks
of Gk without accessing G', since only Gk resides in the cloud.

We adopt the query decomposition method. Given a query Q,' the cloud server decompose Q' into a set of stars (S_i), i = 1, ..., n,
where a star is a root vertex together with its adjacent edges and
neighbors in G'. Then, it finds R(S_i,G') for each star graph S_i,
for i = 1, ..., n. Leveraging the symmetry of G', we can then obtain
R(S_i,G'), based on R(S_i,G'). Finally, we obtain R(Q',G')
by joining the matching results for these stars.

Query Decomposition. We first discuss how to decompose Q'
to a set of stars and how to find the optimal query decomposition.

Consider a query Q' in Figure 2. The stars rooted at vertices
with Type P are shown in Figure 6, where a shaded circle denotes the center (aka root) of a star.

Intuitively, a query decomposition is a set of stars that collectively
cover the outsourced query graph Q'. Figure 6 shows the query
decomposition {S_1, S_2} over query Q'. The set of subgraph match-
es of a star graph S_i with respect to the outsourced graph Gk is
R(S_i,G'), or simply R(S_i) when the context is clear. To reduce
the number of intermediate results, we design a cost model to estimate
the number of matches, |R(S_i)|, for each star S_i. We will discuss the technical
details of the cost model in Section 5. Here, we assume

(3) E(G') is the subset of undirected edges from E(G') that connect
vertices within V(B_1) and vertices between V(B_1) and V(N_1).
Star Matching. decomposition with the minimum cost. Formally, problem. Although ILP is still an NP-hard problem, use an available ILP tool, e.g., the Gurobi ILP solver, to solve this query decomposition with the minimum cost with respect to Definition /*Each edge is contained in at least one star.*/

Qo query decomposition or not (xvi vertex label, and the other one is regarding the neighbourhood structure. Thus, there are two components in the index structure, as shown in Figure 7 which are constructed based on LCT (label correspondence table) and G* (see Figure 2 and Figure 5 respectively). They are the Vertex Bit Vector (VBV) Table and the Neighbour Label Bit Vector (LBV) Table, respectively. For convenience, the first block of G* is denoted as B1. Obviously, B1 is a subgraph of G*.

Each VBV corresponds to a label group, where the corresponding bit in the VBV for a vertex \( v \) in \( B1 \) is set to 1 iff \( v \) contains that label group. For example, \( p2 \) contains \( D \) but not \( E \) in \( B1 \). Thus the two corresponding bits in VBV are 1 and 0, respectively.

In LBV, for each vertex \( v \) in \( B1 \), the corresponding bit for a label group \( L \) is set to 1 if and only if \( L \) is contained in at least one label set for \( v \)'s neighbor vertices. For example, \( D \) is contained in the label set of \( p2 \) that is one of \( p1 \)'s neighbors. However, \( E \) is not contained in the label set of any of \( p1 \)'s neighbors. Thus the two corresponding bits for \( p1 \) in LBV are 1 and 0, respectively.

Algorithm 1 Star Matching Algorithm

Require: Input: \( G* \) (the outsourced data graph) and \( S* \) (the set of stars that contains star \( S1 \) with center \( v1 \) for \( 1 \leq i \leq n \)).

1. Initialize \( RS := \emptyset \).
2. for \( i := 1 \) to \( n \) do
3. Initialize \( RSi := \emptyset \).
4. Set \( a := \{ BS(VBVi, 1) \land VBVi(VBVi, 2) \land \cdots \land VBVi(VBVi, |VBVi|)\} \).
5. for each vertex \( v1 \) that corresponds to a non-zero bit in \( a \) do
6. if \( LBVi(v1) \land LBVi(v1) = LBVi(v1) \) then
7. Generate the set of matches of \( S1 \) with center \( v1 \), denoted as \( RSi \).
8. \( RS := RS \cup RSi \).
9. \( RS := RS \land RSi \).
10: Return \( RS \).

The Star Matching Algorithm is presented in Algorithm 1. Without loss of generality, assume that the center of a star \( Si \) is vertex \( vi \).

For each star \( Si \) for \( 1 \leq i \leq n \), we first find each Vertex Bit Vector that corresponds to every label group of \( v1; e.g., VBVi(C) = (1, 1) \) in our example. Recall \( L(v1) \) is the set of label groups for vertex \( v1 \).

We use \( L(v1) \) to denote the \( j \)-th label group of \( v1 \), for \( 1 \leq j \leq |L(v1)| \).

Hence, the first step is to find \( VBVi(L(v1), j) \) for \( 1 \leq j \leq |L(v1)| \). We perform a bitwise AND operation on these \( L(v1) \) bit vectors to obtain the result vector \( a \) (Line 4).

Each vertex \( v1 \) that corresponds to a non-zero bit in \( a \) is a candidate match of \( v1 \). If each label group of \( v1 \)'s neighbors can be found in the label sets of \( v1 \)'s neighbors (Line 6), some of \( v1 \)'s neighbors can be candidate matches of \( v1 \)'s neighbors. By enumerating candidate vertex combinations of \( v1 \) and its neighbor vertices, we generate matches of \( S1 \) and add them to the result set (Line 7-8).

Result Join. The next challenge is how to compute \( R(S*, G*) \) based on the star matching results \( R(S1, G*) \), for \( t = 1, \ldots, n \). A straightforward solution works as follows. First, using the \( k \)-automorphic function \( F_{j} \) (defined in Figure 4), we can derive \( R(S1, G*) \) and \( R(S2, G*) \).

Specifically, for each match \( M \) in \( R(S1, G*) \), using the automorphic function \( F_{j} \), \( F_{j}(M) \) derives another match \( M' \) in \( R(S1, G*) \). For example, \( (p1, c1, s1) \) is a match of \( S1 \) over \( G* \); thus, \( F_{j}(p1, c1) \), \( F_{j}(c1, s1) \) gives another match \( (p2, c2, s2) \). We get \( R(S2, G*) = R(S1, G*) \cup F_{j}(R(S1, G*)) \cup \cdots \cup F_{k}(R(S1, G*)) \). We can obtain
Algorithm 2 Result Join Algorithm

Require: Input: $R$ (the set of $R(S_a, G^a)$, $1 \leq a \leq n$) and Alignment Vertex Table ($AVT$).
Output: $R_m$.
1: Initialize $R_m := R(S_a, G^a)$, where $|R(S_a)|$ is minimum over all stars.
2: $R := := R = R_m$
3: while $|S| > 0$ do
4: Initialize a set $R_{next} := R(S_a, G^a)$ $(1 \leq i \leq a \leq n)$, where $S_i$ overlaps with the part of query graph that corresponds to current matches in $R_m$, and $|R(S_a)|$ is minimum over all such overlapping stars.
5: Initialize a set $R_{new} := \phi$.
6: for $m := 0$ to $k - 1$ do
7: $R_{next} := R_{next} \cup F_j(R_{next})$.
8: $R_{next} := R_{next} \cup F_j(R_{next})$.
9: $R := R := R_{next}$.
10: for each match $M$ in $R_m$ do
11: if $M$ contains duplicate vertices then
12: $R_m := R_m \setminus M$.
13: $R_m := R_m \setminus M$.
14: Return $R_m$.

4.2.2 Processing in the Client Side

According to Algorithm 2, the cloud obtains the set $R_m$ and transmits it to the client. There are two stages in the client side processing. The pseudocode is given in Algorithm 3.

First, we need to compute $R_{out}$ according to the filtering functions $F_j (j = 1, \ldots, k - 1)$ (Line 1-Line 4 in Algorithm 3). For each match $M$ in $R_m$, we compute $F_j (M)$ $(j = 1, \ldots, k - 1)$ and put them into $R_{out}$. Obviously, $R(Q^c, G^c) = R_m \cup R_{out}$ (Line 5). Note that this step can also be done by the cloud.

Secondly, the client computes the final result set $R(Q, G)$ by filtering out the false positives in $R(Q^c, G^c)$ (Line 6-Line 23). The filtering process also has two steps. In the first step, we remove some matches in $R(Q^c, G^c)$ that contain vertices or edges that do not exist in the original graph $G$ (Line 8-Line 20). Also note that we anonymize the vertex labels in the query graph by using vertex label groups. Thus in the second step, we need to filter out matches that contain vertices whose labels cannot match those of the corresponding vertices in the original query graph $Q$ (Line 21-Line 22).

It is straightforward to see that the time complexity of the client processing is linear with the number of matches in $R(Q^c, G^c)$. Furthermore, it is easy to design some hashing techniques to speed up the filtering processing.

5. COST MODEL

In order to reduce the search space of subgraph matching, we need a cost model for both label generalization and query decomposition. Different label combinations lead to different search s-
pances in subgraph matching. Since our subgraph matching algorithm is based on joining star matches, we require that the number of star matches (|R(S)|) should be as small as possible. Furthermore, the query decomposition method in Section 4.2.1 also relies on estimating |R(S)|. Thus, we propose a cost model for estimating |R(S)|.

5.1 Estimating |R(S)|

Given a star query $S$ with center $q$, it matches a star with the center $v$ in $B_1$ (i.e., the first block of $G^2$) if and only if the following two conditions hold. Therefore, we should consider the two factors while estimating $|R(S)|$:

- center $q$ should match $v$;
- each neighbor of $q$ should match one of $v$'s neighbors.

First Factor. The first factor measures the number of candidate matching vertices of the center of the star query $S$. Given an outsourced graph $G^2$, the vertices matching the center $q$ of $S$ only locate in the first block of $G^2$, i.e., block $B_1$. That is to say, we need to find the number of such candidate vertices $v$ in $B_1$, where each $v$ and $q$ share the same vertex type and each such candidate $v$ contains $q$'s vertex label group. $|V(B_1)|$ is the number of vertices in $B_1$. Given a center $q$, we should estimate the probability that a vertex $v$ in $B_1$ has the same vertex type with $q$ and $v$ contains $q$'s vertex label group.

Due to the symmetry in $G^2$, the first block $B_1$ has the same vertex label distribution with $G^2$. Thus, we use $G^2$ to derive the probability. Let $V(G^2,j)$, $V(G^2,(j,))$, and $V(G^2,(j,i))$ denote the set of vertices with the $j$-th vertex type, label $l_j$, (1-th label of $j$th vertex type), and label group $L_j$, respectively. Then, we define:

$$F_{G^2}(j) = \frac{|V(G^2,j)|}{|V(G^2)|}, \quad F_{G^2}(j,i) = \frac{|V(G^2,(j,i))|}{|V(G^2,j)|}, \quad F_{G^2}(j,i) = \frac{|V(G^2,(j,i))|}{|V(G^2)|}$$

for the graph $G^2$.

Intuitively, $F_{G^2}(j)$ estimates the probability of a vertex being the $j$th vertex type; $F_{G^2}(j,i)$ estimates the probability of a vertex that's the $j$th vertex type having an $i$th label; lastly, $F_{G^2}(j,i)$ estimates the probability a vertex that's the $j$th vertex type having an $i$th label group after the label generalization.

Similarly, we can also define $F_3(j)$, $F_3(j,i)$ and $F_3(j,i)$ for a star query graph $S$, replacing $G^2$ by star $S$ in the equation.

Without loss of generality, assume that for the $j$-th $(1 \leq j \leq t)$ vertex type, there are $\theta_1$ different labels ($l_1, l_2, \ldots, l_{\theta_1}$). These labels can be combined into $h_1$ label groups ($L_1, L_2, \ldots, L_{h_1}$). The $i$-th group $L_{ij}$ contains $\theta_i$ different labels, which are $l_{i_1}, l_{i_1+1}, \ldots, l_{i_n}$ for $1 \leq i \leq h_j$. Note that $(p_1, p_2, \ldots, p_{h_j})$ forms a permutation of $(1, 2, \ldots, h_j)$.

$F_{G^2}(j)F_3(j)$ is the probability that a vertex $v$ in $G^2$ has the same vertex type with the star center $q$. Consider each possible vertex type. For the $j$-th vertex type, $\sum_{i=1}^{b_j} F_3(j)F_3(j,i)$ denotes the probability that vertex $v$ and the query center has the same label group. Therefore, the first factor estimating the number of vertices that can match the center of the star query is as follows.

$$\sum_{i=1}^{b_j} F_3(j)F_3(j,i)$$

Second Factor. The second factor measures the search space of checking whether each of $q$’s neighbors can find its matching vertex in $v$’s neighbors. The estimation here is similar with that of the first factor. The difference is that the candidate matching vertex $v$ of the center $q$ of $S$ has been given. Thus, to match the first vertex of $q$’s neighbors, the number of candidate vertices that we should search is the degree of vertex $v$ rather than $\frac{|V(B_1)|}{\theta_1}$. Since there are several candidate vertices $v$, we use the average degree of vertices in $G^2$ to estimate the degree of vertex $v$. Here, we denote it as $D(G^2)$. Then, to match the second vertex of $v$’s neighbors, the potential search space is $D(G^2)$ – 1. The rest can be done in the same manner. Suppose the center of the star query $S$ has $D(S)$ neighbors. Thus, this part of the search space can be estimated as $D(G^2)\cdot(D(G^2)−1)\cdot\cdots\cdot(D(G^2)−D(S)+1)$. For the sake of simplicity, we can estimate it as $D(G^2)^{D(S)}$. As with the estimation of the first factor, we should also consider the probability of sharing the same vertex type and containing the corresponding vertex label group.

Thus, we can define the second factor that estimates the search space of matching the star center $q$’s neighbors as follows.

$$D(G^2)^{D(S)}\sum_{i=1}^{b_j} F_3(j)F_3(j,i)\sum_{i=1}^{b_j} F_3(j)F_3(j,i)$$

The Cost Model. Our cost model is given by Expression 4, which is simply the product of factor 1 and factor 2 from (2) and (3).

It does assume independence among the label distributions for the neighbors of a vertex. This may not always hold in practice. But our experimental results show that our assumption is acceptable in three real large graphs and our cost model is very effective. We also note that $|V_j(G^2,(j,i))| \leq |1 + \delta(k)| \cdot \sum_{i=1}^{h_j} |V(G^2,(j,i))|$ for some constant $0 \leq \delta(k) \leq k − 1$, i.e., the number of vertices from the $j$th vertex type having the $i$th label group is at most a constant factor of the total number of vertices with $j$th vertex type having a label that was generalized into this label group. Immediately, this implies that $F_3((j,i)) \leq |1 + \delta(k)| \cdot \sum_{i=1}^{h_j} F_3((j,i))$. Intuitively, each vertex $u$ in $G^2$ has $(k − 1)$ symmetric vertices. To ensure that they have the same vertex groups, we require that $u$ should have a union of all its symmetric vertices’ label groups. In the worst case, $F_3((j,i))$ will increase by a factor of $(k − 1)$. In fact, $\delta(k)$ can be much less than $(k − 1)$, and given $G$ and the corresponding $\kappa$-automorphic graph $G^2$, we can give a much tighter bound on the parameter $\delta(k)$.
will be 0. In practice, δ(k) is far less than 1 when k is small, as demonstrated in our experiments.

\[ |RS| \leq \frac{|V(G^s)|}{k} \sum_{j=1}^{l} \left( \frac{\sum_{i=1}^{l} F_{G^s}(j)}{|S|} \right) \left( \frac{\sum_{i=1}^{l} F_{G^s}(j)}{|S|} \right)^{D_s(S_{avg})} \]

(4)

5.2 Label Combination

The cost model based query decomposition has been studied in Section 4.2.1. Hence, we only need to show how to do label generalization (label combination) based on the cost model above. Our solution works as follows. Let us consider the j-th vertex type and its \( \theta \) different vertices (\( \{l_1, l_2, \ldots, l_{\theta}\} \)). Assume that \( P = \langle p_1, p_2, \ldots, p_{\theta}\rangle \) is a permutation of \( \{1, 2, \ldots, \theta\}\). We divide P sequentially into \( \theta \) groups, where each group has \( \theta \) vertex labels. Each permutation corresponds to one possible label combination. The goal is to find the optimal permutation, which leads to the minimum cost in star matching.

In the derivation of Expression 4, we only estimate the search space of a single star query \( S \). However, to perform the label combination of the \( k \)-automorphic graph \( G^s \), we need to estimate the search space of star queries in the average case. The average cost can be estimated by exploring different query patterns. Suppose \( S_{all} \) is the set of all possible star queries. Similar to the definitions in Equation 1 in Section 5.1, we can define

\[ F_{S_{avg}}(j) = \frac{\sum_{s \in S_{all}} |s| \cdot |s|}{|S_{all}|} \cdot F_{S_{avg}}(j, i) = \frac{\sum_{s \in S_{all}} |s| \cdot |s|}{|S_{all}|} \cdot F_{S_{avg}}(j, i) = \frac{\sum_{s \in S_{all}} |s| \cdot |s|}{|S_{all}|} \]

for star queries. These values rely on all possible star queries rather than any one single star query. Similarly, let \( D_s(S_{avg}) = \frac{\sum_{s \in S_{all}} D_s(s)}{|S_{all}|} \).

Now similar to the derivation of Expression 4, we can estimate the search space of star queries in the average case, denoted as \( |RS(S_{avg})| \), in Expression 5 (the derivation follows the same steps as that in Expression 4). According to Expression 5, the component that concerns the label combination is given in Expression 6.

\[ |RS(S_{avg})| \leq \frac{\sum_{s \in S_{all}} |s| \cdot |s|}{|S_{all}|} \cdot \left( \frac{\sum_{s \in S_{all}} |s| \cdot |s|}{|S_{all}|} \right)^{D_s(S_{avg})+1} \]

(5)

Therefore, we define the label combination cost as follows.

**Definition 7. Label Combination Cost.** Given the \( j \)-th vertex type and its \( \theta \) different vertex labels (\( \{l_1, l_2, \ldots, l_{\theta}\} \)), we can form a list \( \langle F_{G^s}(j, 1), F_{S_{avg}}(j, 1) \rangle, \langle F_{G^s}(j, 2), F_{S_{avg}}(j, 2) \rangle, \ldots, \langle F_{G^s}(j, \theta), F_{S_{avg}}(j, \theta) \rangle \rangle \), where \( \sum_{\theta=1}^{\theta} F_{G^s}(j, i) = 1 \) and \( \sum_{\theta=1}^{\theta} F_{S_{avg}}(j, i) \).

**Figure 9:** An example of label anonymization algorithm.

Thus we need to choose an effective permutation \( P \) to decrease \( \text{cost}(P) \) as much as possible. To achieve that, we propose a heuristic solution. It works in an iterative manner. Initially, we generate a random label combination. Then, in each iteration, we try to swap two labels in two label groups, respectively. If the swap leads to smaller cost (see Expression 6), we keep the swap; otherwise, we ignore that. For example, we keep the swap between \( l_{1,2} \) and \( l_{1,4} \) (see Figure 9) in the first iteration. We consider all possible swaps sequentially. Since we cannot find a swap that leads to smaller cost, the algorithm stops. Our experimental results show that for each of the three datasets in Section 6 and each vertex type, we generally need no more than 10 such iterations before the results converge.

An example of label anonymization algorithm for \( \theta = 2 \) (two labels in each label group) is represented in Figure 9, using our running example.

6. EXPERIMENTAL RESULTS

6.1 Datasets and Setup

We evaluated our method on three real datasets in our experiments. Statistics on these graphs are given in Table 2.

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<th>Dataset</th>
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**Table 2:** Real graph datasets.
In this subsection, we evaluate the performance of generating $G^k$ and $G^o$. The default value of $\theta$ (the number of labels contained in each label group) is 2 in all the experiments. The main results of these experiments are as follows.

UK-2002. UK-2002 is obtained from a crawl of the .uk domain performed by UbiCrawler in 2002. We find that the frequencies of different vertex labels on these graphs all (roughly) obey Zipf’s law of different skewness.

SETUP. We compare four methods EFF, BAS, FSIM and RAN, where EFF is our method with all optimizations discussed.

1. EFF applies both the “cost model based” label generalization and the graph size reduction (i.e., uploading $G^k$).
2. BAS applies the same cost model based label generalization approach with EFF; but BAS uploads the entire $G^o$ directly.
3. FSIM and RAN adopt two different label generalization approaches. FSIM combines vertex labels with similar frequencies in the data graph into the same label groups, while RAN randomly combines vertex labels into label groups. However, both FSIM and RAN adopt the graph size reduction approach proposed in this paper, i.e., they upload $G^k$ instead of $G^o$. We use a Windows 7 PC with 3.0 GHz Intel Core 2 Duo CPU and 8 GB main memory and four CPU cores. All methods are implemented in C++. Some additional experimental results are presented in Appendix B.

6.2 Cost of Generating $G^k$ and $G^o$

In this section, we evaluate the performance of generating $G^k$ and $G^o$. The default value of $\theta$ (the number of labels contained in each label group) is 2 in all the experiments. The main results of these experiments are as follows.

UK

Figure 10: Time cost in generating $G^2$.

(a) Web-NotreDame

(b) DBpedia

Figure 11: Number of noise edges in $G^2$.

(a) Web-NotreDame

(b) DBpedia

Figure 13: Index cost of our algorithm.

(a) Index size.

(b) Index construction time.

The cost model based label anonymization approach EFF is as efficient as simple strategies like RAN and FSIM when generating $G^k$, as shown in Figure 10. The cost on Web-NotreDame slightly increases when $k$ goes from 2 to 6, but the running time on DBpedia decreases as $k$ becomes larger. The reason is that there are a few high-degree vertices on DBpedia. We use the BFS strategy in graph alignment for generating $G^k$, when $k$ is small, such as 2, we have to do more explorations on these high-degree vertices. The experimental results on UK-2002 are similar with those on DBpedia; see Appendix B.

The number of noise edges $|E(G^k)| - |E(G)|$ does not depend on the label anonymization approach. We find that the three approaches introduce nearly the same number of noise edges (as shown in Figure 11). The number of noise edges roughly grows linearly when $k$ goes from 2 to 6. The experimental results on UK-2002 are also similar with those on the other two datasets; see Appendix B.

We also report the number of edges in $G^k$ and $G^o$ (i.e., space and communication cost) in Figure 12. Since the three approaches introduce nearly the same number of noise edges, we only report the results generated by EFF method. As shown in Figure 12, the edge number of $G^k$ is much less than that of $G^o$. Especially, when $k$ is small, $|E(G^k)|$ is close to $|E(G)|$. It shows that our method not only ensures the data privacy but also saves the online query cost, since online subgraph matching is performed on $G^k$ rather than $G^o$.

Although the three label anonymization algorithms have similar efficiency in generating $G^o$, they do generate different $G^o$’s. EFF produces $G^o$’s that perform much better than the graphs produced by the other two methods in terms of the query performance. We will report these results shortly.

6.3 Performance in the Cloud

We first report the cost of index construction for our method. Figure 13 shows the space and time cost of constructing the index structure, which is discussed in Section 4.2.1. Both space and time costs of our index construction decrease while $k$ increases from 2 to 6. The reason lies in the fact that the size of our index (see Figure 7) depends on the vertex number of $G^k$ (i.e., $|V(G^k)|$). Obviously, $|V(G^k)|$ decreases with the increase of $k$, since $G^k$ roughly contains only the first block of vertices in $G^o$ and each block in $G^k$ has about $\frac{|V(G^o)|}{k}$ vertices. Thus, larger $k$ leads to less index size and less construction cost.

Then we pay attention to the cost of subgraph matching in the cloud. We generate query graphs by randomly extracting connected subgraphs from the data graph $G$, ensuring that $|E(Q)|$ meets a user-specified parameter value $N$. Specifically, we randomly locate the first edge $e$ from the data graph $G$ and set $E(Q) = e$. We then expand the current query graph $Q$ through a random walk over $G$ iteratively until it reaches $N$ edges. We used 100 queries and report the average. We compared our method with the baseline BAS and the other two label generalization techniques. First, we report the query performance for different query graph sizes $|E(Q)|$. Figure

Table: Number of edges in $G^k$ and $G^o$ using EFF.

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Table: Number of noise edges in $G^k$.

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Table: Number of noise edges in $G^o$.

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Table: Number of noise edges in $G^o$.

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<td>2</td>
<td>4</td>
<td>8</td>
<td>16</td>
<td>32</td>
</tr>
<tr>
<td>[G(G)]</td>
<td>2</td>
<td>4</td>
<td>8</td>
<td>16</td>
<td>32</td>
</tr>
<tr>
<td>[G(G)]</td>
<td>2</td>
<td>4</td>
<td>8</td>
<td>16</td>
<td>32</td>
</tr>
<tr>
<td>DBpedia</td>
<td>2</td>
<td>4</td>
<td>8</td>
<td>16</td>
<td>32</td>
</tr>
<tr>
<td>[G(G)]</td>
<td>2</td>
<td>4</td>
<td>8</td>
<td>16</td>
<td>32</td>
</tr>
<tr>
<td>UK-2002</td>
<td>2</td>
<td>4</td>
<td>8</td>
<td>16</td>
<td>32</td>
</tr>
</tbody>
</table>
Furthermore, BAS always performs the worst, since experimental results on UK-2002 are also similar with those on other cost model based label combination. Thus, the effectiveness of client model based label combination.

We find similar results on DBpedia, as shown in Figure 15. The experimental results on UK-2002 are also similar with those on other datasets; see Appendix B. When the query size becomes larger, EFF outperforms the other three approaches by at least one order of magnitude. Furthermore, BAS always performs the worst, since the search space on $G^*$ is much larger than that on $G^s$.

The running time of subgraph matching increases with $k$ from 2 to 6, as shown in Figure 14 and Figure 17. This is because that $|E(G^s)|$ increases with $k$ from 2 to 6, since we must insert more noise edges into $G^s$ when $k$ becomes larger. Nevertheless, EFF is always the best method. Furthermore, compared with other approaches, the advantage of EFF becomes even more significant for larger values of $k$.

We also find that both the query decomposition algorithm and the star matching algorithm run very fast. Even when $|E(Q)|$ is as large as 12, the time cost of query decomposition algorithm is less than 1 ms. As for the time cost of star matching algorithm, Figure 18 shows the experimental results which are also very efficient. We also report the size of the set $RS$ generated by our star matching algorithm in Figure 19. $|RS|$ has a large influence on the time cost of result join algorithm. These results show the importance of optimizing the result join algorithm.

### 6.4 Processing Cost in the Client Side

The last step of all methods involves processing in the client side. Figure 20 and Figure 21 illustrate this overhead on the first two real graphs. The goal of client side processing is to filter out all false positive matches due to the noise edges and labels in $G^s$.

First of all, note that for all methods, client processing cost is much less than cloud processing cost; it is orders of magnitude less. As Figure 20 and Figure 21 show, the overhead in the client side scales very well in terms of query size and $k$. EFF still outperforms both RAN and FSIM, but it is slightly worse than BAS in the client side. The reason is that EFF generates fewer intermediate results than RAN and FSIM, due to its effective label combination algorithm. However, following our processing framework, EFF, RAN, and FSIM produce $R_n$ (i.e., a small subset of $R(G^s, G^s)$, see Section 4.2.1) in the cloud, and the client needs to find $R(G^s, G^s)$ based on $R_n$ and our client side algorithm as shown in Section 4.2.2. In contrast, BAS obtains $R(G^s, G^s)$ from the cloud directly (since the cloud has $G^s$ instead of $G^s$ as that in our method).

However, note that client processing time is a tiny fraction of the entire processing time. Furthermore, compared with our method like EFF, BAS has much more expensive communication cost, since $|R(G^s, G^s)|$ is always much larger than $|R_n|$ (i.e., the size of a small subset of $R(G^s, G^s)$, see Section 4.2.1). For example, the average size of matching results for $|E(Q)| = 6, k = 4$ on UK-2002 in BAS is 12,224 bytes, while EFF only transmits 3,056 bytes.

<table>
<thead>
<tr>
<th>Dataset</th>
<th>$k = 3$</th>
<th>$k = 5$</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$</td>
<td>E(Q)</td>
</tr>
<tr>
<td>Web-NotreDame</td>
<td>EFF 9</td>
<td>31</td>
</tr>
<tr>
<td></td>
<td>RAN 19</td>
<td>69</td>
</tr>
<tr>
<td></td>
<td>FSIM 25</td>
<td>87</td>
</tr>
<tr>
<td></td>
<td>EFF 7</td>
<td>19</td>
</tr>
<tr>
<td></td>
<td>RAN 11</td>
<td>36</td>
</tr>
<tr>
<td></td>
<td>FSIM 14</td>
<td>50</td>
</tr>
<tr>
<td>DBpedia</td>
<td>EFF 7</td>
<td>24</td>
</tr>
<tr>
<td></td>
<td>RAN 11</td>
<td>38</td>
</tr>
<tr>
<td></td>
<td>FSIM 15</td>
<td>52</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Star Matching Time (ms)</th>
<th>$k = 3$</th>
<th>$k = 5$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Web-NotreDame</td>
<td>EFF 456</td>
<td>1262</td>
</tr>
<tr>
<td></td>
<td>FSIM 591</td>
<td>1533</td>
</tr>
<tr>
<td></td>
<td>RAN 552</td>
<td>1364</td>
</tr>
<tr>
<td>DBpedia</td>
<td>EFF 460</td>
<td>1258</td>
</tr>
<tr>
<td></td>
<td>FSIM 592</td>
<td>1534</td>
</tr>
<tr>
<td></td>
<td>RAN 552</td>
<td>1364</td>
</tr>
<tr>
<td>UK-2002</td>
<td>EFF 456</td>
<td>1262</td>
</tr>
<tr>
<td></td>
<td>FSIM 591</td>
<td>1533</td>
</tr>
<tr>
<td></td>
<td>RAN 552</td>
<td>1364</td>
</tr>
</tbody>
</table>

| $|RS|$ (MB) | $k = 3$ | $k = 5$ |
|------------|---------|---------|
| Web-NotreDame | 10      | 20      | RAN 25 | 50      |
|               | FSIM 30 | 60      | EFF 40 | 80      |
| DBpedia       | 15      | 30      | RAN 40 | 80      |
|               | FSIM 60 | 120    | EFF 80 | 160     |
| UK-2002       | 20      | 40      | RAN 50 | 100     |
|               | FSIM 100 | 200    | EFF 120 | 240     |
other competing methods significantly. In particular, EFF outperforms baseline label anonymization with graph size reduction approach, the client side. Figure 22 shows that our method EFF, the cost model for answering subgraph matching queries over encrypted graphs in the cloud [3], for a database of graphs. The method pre-builds a feature-based index for each data graph. After filtering in the cloud, each candidate supergraph is verified by checking subgraph isomorphism in the client. However, the method does not apply to our case where the goal is to find subgraph matches on a single large graph (instead of many small graphs).

### 8. CONCLUSION

In this paper, we present an efficient framework for privacy preserving subgraph matching on a large graph in the cloud. Our design protects both structural and label privacy in a graph, without losing data utility. By exploring a number of optimization techniques and leveraging an effective cost model, our method achieves superior query performance compared to the baseline method. Extensive experiments on large real graphs confirm the effectiveness and efficiency of our approach. Extending our framework to handling other important graph queries will lead to a number of interesting and important future works.

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Appendix

A. Proof of Theorems.

Theorem 1. Given a data graph $G$ and a query graph $Q$, $R(Q, G) \subseteq R(G', G^b)$.

Proof. For any match $M \in R(Q, G)$, we assume that $g : V(Q) \rightarrow V(G)$ is the injective function. We can let $g' : V(Q') \rightarrow V(G')$ have the same mapping relation with $g$, since $G$ is a subgraph of $G'$. Thus we have $\forall q \in V(Q) \Rightarrow g'(q) \in V(G')$ and $\forall q, q' \in E(Q')$. If $g(q, q') \in E(G)$, since we use the same label grouping strategy on $G$ and $Q$, $\forall q \in V(Q') \Rightarrow L(g(q)) \subseteq L(g'(q))$. According to Definition 2, $M \in R(G'; G^b)$, represented as $(g'(q_1), \ldots, g'(q_n))$.

Theorem 2. Given an outsourced query graph $Q^o$, finding the query decomposition with the minimum cost with respect to Definition 6 is an NP-hard problem.

Proof. Finding a minimum vertex cover is a classical NP-hard problem. When each $|R(S_x)| = 1$ ($i = 1, \ldots, n$), finding the optimal query decomposition is equivalent to finding a minimum vertex cover. Thus finding a minimum vertex cover is a subproblem of finding the optimal query decomposition.

Theorem 3. Given $Q^o$ with $m$ vertices $v_q$ ($a = 1, \ldots, m$), $M^o$ with $m$ vertices $v_{q^o}$ ($a = 1, \ldots, m$) is a subgraph match of $Q^o$ over graph $G^b$, where $v_{q^o}$ matches $v_q$. Consider a vertex $v_q$ in $Q^o$. If vertex $v_{q^o}$ matching $v_q$ is not in $B_1$ (the first block of $G^b$), we can find another match $M$ that contains a vertex $v_{q^o}$ matching $v_q$, where $v_{q^o} \in B_i$ and $v_{q^o} = F_j(v_{q^o})$ under some automorphic function $F_j$. Furthermore, $M^o = F_j(M)$.

Proof. Since $M^o$ is a subgraph of $G^b$, we have $v_{q^o} \in G^b$. Without loss of generality, assume that $v^i_{q^o} \in P_i$ ($1 < i \leq k$). We set $v_{q^o} = F_{k+1-i}(v_{q^o})$, i.e., $j = i - 1$ and $v_{q^o} = F_j(v_{q^o})$. Thus $v_{q^o} \in P_i$.

Here, we prove that $M = F_{k+1-i}(M^o)$ is another match of $Q^o$ over $G^b$. Thus $M^o = F_j(M)$.

For the match $M^o$, we assume that $g^o : V(Q^o) \rightarrow V(G^b)$ is the injective function. Here, we prove that $g^o = F_{k+1-i}(g^o)$ is the injective function of $M$. Obviously, $\forall q \in V(Q^o) \Rightarrow g^o(q) \in V(G^b) \Rightarrow F_{k+1-i}(g^o(q)) \in V(G^b) \Rightarrow g(q) \in V(G)$. In a similar way, $\forall q \in V(Q^o) \Rightarrow g^o(q) \in V(G^b)$, and $g(q) \in V(G)$. According to Definition 3, $\forall q \in V(Q^o) \Rightarrow L(g(q)) \subseteq L(g^o(q)) = L(g^o(q))$. According to Definition 2, $M \in R(G'; G^b)$, represented as $(g(q_1), \ldots, g(q_n))$.

B. Additional Experimental Results

Figure 23 shows the time cost in generating $G^b$ on UK-2002. Figure 24 shows the number of noise edges in $G^b$ on UK-2002. Figure 25 and Figure 26 show the query time in the cloud on UK-2002. Figure 27 shows the filtering time in the client side on UK-2002. As these figures show, the experiment results on UK-2002 are similar with those on other datasets. Figure 28, Figure 29 and Figure 30 show the query time in the cloud on the three datasets, respectively. Figure 31 shows the time cost of our star matching algorithm on the three datasets. Figure 32 shows the size of the set $RS$ generated by our star matching algorithm on the three datasets. Figure 33 shows the network overhead on the three datasets when transferring the candidate matching results from the cloud to the client side. Figure 34 shows the end-to-end running time on the three datasets. These results show similar trends as those in the experiment section and further confirm the superiority of our method.

$F_j(M)$ is a mapping graph of $M$ under an automorphic function $F_j$, which is defined in Definition 4.
Figure 23: Time cost in generating $G^k$ on UK-2002.

Figure 24: Number of noise edges in $G^k$ on UK-2002.

Figure 25: Query vs. $|E(Q)|$ on UK-2002.

Figure 26: Query time vs. $k$ on UK-2002.

Figure 27: Client processing time on UK-2002.

Figure 31: Star matching time.

Figure 32: The size of result set generated by star matching algorithm ($|RS|$).

Figure 33: Network overhead.

Figure 34: Overall running time.
Figure 28: Query time vs. $|E(Q)|$ on Web-NotreDame.

Figure 29: Query time vs. $|E(Q)|$ on DBpedia.

Figure 30: Query time vs. $|E(Q)|$ on UK-2002.