Scalable and Adaptive Graph Querying with MapReduce

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SUMMARY We address the problem of processing graph pattern matching queries over a massive set of data graphs in this letter. As the number of data graphs is growing rapidly, it is often hard to process such queries with serial algorithms in a timely manner. We propose a distributed graph querying algorithm, which employs feature-based comparison and a filter-and-verify scheme working on the MapReduce framework. Moreover, we devise an efficient scheme that adaptively tunes a proper feature size at runtime by sampling data graphs. With various experiments, we show that the proposed method outperforms conventional algorithms in terms of scalability and efficiency.

key words: graph query, parallel processing, MapReduce, adaptive tuning

1. Introduction

Graphs are widely used to model complex structures such as protein-to-protein interactions and Web data [1]. Graph data are growing rapidly and massively, e.g., PubChem [2] now serves more than 40 million graphs that represent chemical compounds. The overall storage size of the graphs hits tens of terabytes [2]. A graph pattern matching query is to find all data graphs that contain a given query graph from a set of data graphs. A graph pattern matching query requires to process expensive subgraph isomorphism test, which is known to be NP-complete [3]. Thus, many studies have been reported in the literature to cut down the number of subgraph isomorphism tests [4], [5]. Although the algorithms are prominent, they are not scalable enough to handle a massive set of data graphs like PubChem since they all are serial, designed to work on a single machine.

Meanwhile, it is important to set a proper feature size in a filter-and-verify scheme since it dominantly determines the overall performance of query processing (refer to Sect. 6). However, it is hard to find a proper feature size before query processing since it depends on the statistics about both query graphs and data graphs. If the feature size is too large, the overhead of extracting and storing features increases sharply. On the other hand, many false positives are produced at filtering stage if the feature size is too small, involving many obsolete subgraph isomorphism tests.

To tackle the problems, we propose MR-Graph, a distributed graph querying algorithm based on MapReduce [6]. MR-Graph tunes the feature size at runtime. It determines the most appropriate feature size by sampling some portions of data graphs. MR-Graph processes multiple queries over a massive set of a data graph simultaneously. In addition, MR-Graph does not require any index building and the modification of MapReduce internals. MR-Graph is sophisticatedly tailored to work in harmony with MapReduce’s programming model.

2. Related Work

A major issue in the graph pattern matching problem is to reduce the number of pairwise subgraph isomorphism testing. Most approaches to the issue are based on a filter-and-verify scheme with index techniques [4], [5]. They extract substructures called features from data graphs and then filter irrelevant graphs before actual subgraph isomorphism testing. It may greatly reduce the number of actual subgraph isomorphism tests. However, all the algorithms are serial and thus not scalable enough to process a massive set of data graphs. MapReduce is a programming model that enables to process a massive volume of data in parallel with a large cluster of commodity PCs [6]. It provides a simple but powerful method that enables automatic parallelization and distribution of large-scale computation. Recently, Luo et al. [7] proposed an index-based method for parallel graph pattern matching on MapReduce. They build an edge index over data graphs, which maps each distinct edge to the list of data graphs that include the edge. However, their approach has a restricted assumption that every edge in a query graph must be uniquely identified by labels of its endpoints and itself. Thus it requires a subsequent verification phase to process general graph pattern matching queries. There are some studies about processing graphs with MapReduce. Lin et al. [8] introduce several design patterns suitable for iterative graph algorithms such as PageRank. Pregel [9] and Pegasus [10] are systems devised for processing large-scale graphs. However, they are different from our work in that they focused on processing a single large graph such as Web data or social network.

3. Preliminaries

We denote a graph by a tuple \( g = (V, E, L, l) \), where \( V \) is a set of vertices and \( E \) is a set of undirected edges such that \( E \subseteq V \times V \). \( L \) denotes a set of labels for both vertices and...
Given two graphs $g = (V, E, l)$ and $g' = (V', E', l')$, $g$ is subgraph isomorphic to $g'$, denoted by $g \subseteq^I g'$, if only if there exists an injective function $\phi : V \rightarrow V'$ such that

1. $\forall u \in V$, $\phi(u) \in V'$ and $l(u) = l'(\phi(u))$,
2. $\forall (u, v) \in E$, $(\phi(u), \phi(v)) \in E'$ and $l(u, v) = l'(\phi(u), \phi(v))$.

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**Problem statement** Let $D = \{g_1, g_2, \cdots, g_n\}$ be a set of data graphs and $Q = \{q_1, q_2, \cdots, q_m\}$ be a set of query graphs, where $m << n$. For each query $q \in Q$, we find all graphs $D_q$ such that $D_q = \{g | q \subseteq^I g, g_i \in D\}$.

Figure 1 shows an example which will be used throughout this paper. In the example, answers for two query graphs are $D_{q_3} = \{g_1, g_2, g_3\}$ and $D_{q_2} = \{g_1\}$, where $D_q$ means a set of answers for $q_i$.

**Graph representation** In MapReduce, all input data are represented by key-value pairs. To parallelize graph query processing with MapReduce, we first encode each graph as a single key-value pair. A graph $g$ is encoded as a pair of $(ID, CODE)$ where $ID$ is a unique identifier for each graph and $CODE$ is a serialized version of $g$. CODE is composed of the number of vertices, the number of edges, a list of vertex labels, and a list of edges. In the format, each edge is represented by a triple that consists of from-vertex $ID$, to-vertex $ID$, and edge label. For example, query graph $q_2$ in Fig. 1 (a) is encoded as $(q_2, \langle 3, 3, A, B, B, \emptyset, 1, b, 0, 2, b, 1, 2, b, b \rangle)$.

**Features in graph query processing** A feature is a loosely defined term for a substructure of a graph. There are various kinds of features such as path, subtree, and subgraph [4], [5]. In this letter, we use a subgraph as feature because it exhibits the best filtering power among them [5]. However, any other substructures can be used in MR-Graph without loss of generality.

### 4. Basic Algorithm

A MapReduce-based algorithm works in two stages: **map** and **reduce**. In MapReduce, input data are first partitioned into equal-sized blocks and each block is assigned to a mapper at map stage and mapped outputs are then stored into local disks. Finally, the outputs are shuffled and pulled to reducers for further processing at reduce stage. Algorithm 1 is the basic algorithm of MR-Graph. Each mapper processes queries over data graphs and each reducer groups mapped outputs by query graph’s ID.

| Algorithm 1: Basic MR-Graph |
|-----------------------------|
| **input:** $D$: a set of data graphs, $Q$: a set of query graphs  |
| **output:** $(ID_q, a$ list of $ID_q)$ pairs |
| init() \hspace{1cm} // initializing a mapper |
| $\overrightarrow{F}[] = \text{ExtractFeatures}(Q)$ |
| map (String $ID_q$, String $CODE_q$) \hspace{1cm} // mapper |
| $\overrightarrow{F}_q = \text{ExtractFeatures}(CODE_q)$ |
| foreach $(ID_q, CODE_q) \in Q$ do |
| if CheckContainment($\overrightarrow{F}[ID_q], \overrightarrow{F}_q$) then |
| if TestSubIso (CODE_q, CODE_q) then |
| emit $(ID_q, ID_q)$ |
| reduce (String $ID_q$, iterator $\overrightarrow{ID}_q$) \hspace{1cm} // reducer |
| answer = Enlist $(\overrightarrow{ID}_q)$ |
| emit $(ID_q, answer)$ |

At map stage, we extract features from a query graph and a data graph by different rules (line 2 and 4). Graph query processing is performed in a filter-and-verify scheme. For each data graph, MR-Graph tests whether the data graph contains all the features that compose a certain query graph in a query set $Q$ (line 6). If it is true, the algorithm pairs data graph with the query graph and tests subgraph isomorphism (line 7). At reduce stage, result data graphs are grouped by a query graph ID and emitted. MR-Graph greatly reduces the number of graphs that must be tested by subgraph isomorphism based on the filter-and-verify scheme.

When it comes to feature extraction, we extract $\min(k, |q|)$-sized features for a query graph $q$ where $|q|$ is the size of query graph $q$. Note that we do not use features whose sizes are less than $\min(k, |q|)$. We explain a rationale behind this. Suppose that a query graph $q$ has two features $f_s$ and $f_y$, where $f_s \subseteq^I f_y$ and the size of feature $f_y$, denoted by $|f_y|$, is set to $\min(k, |q|)$. It is straightforward that a data graph $g$ contains $f_s$ if it contains $f_y$. Thus, testing if $g$ contains $f_s$ is unnecessary when $g$ is confirmed to contain $f_y$. Meanwhile, a data graph is processed with a set of various sized query graphs. For a data graph $g$, we extract features whose sizes are up to $\min(k, |q|)$. In other words, the size of feature $f$ for a data graph $g$ must satisfy the condition, $1 \leq |f| \leq \min(k, |q|)$. The ExtractFeatures function in Algorithm 1 extracts features from both query graphs and data graphs with the feature size following the rules described thus far.

### 5. AdaptiveTune: Tuning the Feature Size at Runtime

Note that the feature size $k$ in Algorithm 1 is determined by a user before runtime. However, feature size decision is not easy since the best feature size varies according to the characteristics of query graphs and data graphs. To address the issue, we devise an adaptive feature size tuning technique. **AdaptiveTune** examines a set of trial values for $k$, denoted by $V_k$, in the first few mappers, called trial mappers, then determines a proper feature size. Note that the number of...
Algorithm 2: AdaptiveTune

input: $k_{init}$: initial feature size, $V_k$: trial values for $k$,
$\alpha$: number of trials for each feature size in $V_k$
output: $k$: the best appropriate feature size
1. The initial feature size $k_{init}$ is set into DCS.
2. If a mapper is a trial mapper, choose one value from $V_k$.
   Otherwise, use the feature size stored in DCS.
3. The trial mapper updates DCS.
4. DCS notifies its update to all running mappers.
5. For each running mapper,
   a. Estimate the remaining time left till the mapper ends,
      denoted by $T_{current}$, and the remaining time expected if
      the mapper uses the best feature size, denoted by $T_{best}$.
   b. If $T_{best} < T_{current}$, change the feature size with one in DCS.
   c. Process the remaining data graphs in the input split.

AdaptiveTune that works with 4 mappers. Suppose that $V_k = \{k_1, k_2, k_3\}$ and $\alpha = 1$, then the number of trial mappers is 3. T1, T2, and T3 are trial mappers and they use one of $V_k$, while T4 uses the feature size stored in DCS, $k_{init}$. After finishing its input split, T2 updates DCS with $\{k_2, \ldots\}$ at time $t_1$. DCS notifies this tuple to the other mappers. Then the mappers determine whether they keep or change their feature size, and process remaining input splits with the feature size. Finally, T1 and T4 change the current feature size to $k_2$.

6. Experiments

Experimental setup We performed our experiments with Hadoop version 1.0.3 working on a 9-node cluster. Each data node is equipped with an Intel i5-2500 3.3 GHz processor, 8 GB memory, and a 7200 RPM HDD, running on CentOS 6.2. All nodes are connected via a Gigabit switching hub. The number of task slots is 32, which indicates how many tasks can run simultaneously on Hadoop. The total number of map and reduce tasks were set to be 8 times (256) and 2 (64) times as many as the number of task slots. We tested two datasets introduced in [11], i.e., PubChem for a real-world dataset and synthetic.E30.D3.L50 for a synthetic dataset. In addition, we varied the number of data graphs or query graphs. Table 1 presents the statistics of our datasets.

| Dataset | D/Q | | | | size |
|---------|-----|-----|-----|-----|
| Real    | D   | 10M | 23.98 | 25.76 | 78 | 2.31 GB |
|         | Q   | 1000 | 14.26 | 14.00 | 11 | 131 KB |
| Synthetic | D   | 10M | 14.23 | 30.53 | 47 | 2.62 GB |
|         | Q   | 1000 | 10.25 | 12.00 | 47 | 120 KB |

Table 1 Statistics of graph datasets.

We compared our method with a non-filtering approach, denoted by Basic, and Luo et al.’s approach, denoted by Luo [7]. All algorithms are written in C++ and the verification phase was attached into Luo for fair comparison (please refer to Sect. 2). In our system, we used VF2 [12] for subgraph isomorphism test, gSpan [13] for feature extraction, and ZooKeeper [14] for distributed coordination.

Result analysis We first tested how a feature size $k$ affects the overall performance of graph pattern matching. Figure 3 shows that the execution time, depicted by bars, is the fastest
when $k$ is set to 4 whereas the average number of candidates, depicted by lines, is the least when $k$ is 6 for the real dataset. The reason is that the cost of feature extraction and feature comparison offsets the benefit of fine filtering effect. The best value for $k$ on the synthetic dataset is 2 as shown in Fig. 4. Note that we omitted some results which exceeded 24 hours in our figures. The results also confirmed that the best $k$ can be different according to the characteristics of given data and query graph sets. Figures 5 and 6 show how a feature size impacted on the overall performance. As the feature size increases, both the total storage size of feature sets and the average feature extraction per data graph increase exponentially. This causes not only space overhead to store the features, but a lot of I/Os during query processing on Hadoop.

We now evaluate AdaptiveTune. Figures 7 and 8 show the experimental results. In the figures, the ‘Preset’ value is a fixed initial feature size whereas the ‘Best’ value is the best feature size computed after finishing all graph queries. In both cases, AdaptiveTune (AT) shows good execution time even compared to the best feature size. With the real dataset, AdaptiveTune was successful to reduce the execution time 52.5%.

We also compared our algorithm with conventional algorithms. We used 1 million real dataset rather than 10 million graphs in Table 1, since Luo exhibited limitation in scalability. As shown in Fig. 9, our algorithm is always better than the others in terms of execution time per query except in the case of processing a single graph query. As the number of query graphs becomes larger, MR-Graph outperforms two other algorithms more. We also noticed that the performance of Luo was even poorer than our Basic algorithm. The reason is that Luo suffered from frequent accesses to the storage of index files and a skewness problem; value distribution in the indexes is highly skewed [15]. Finally, we evaluate scalability of MR-Graph. As shown in Fig. 10, MR-Graph is scalable to the size of a cluster. Specifically, MR-Graph records 25.3 times faster than Basic in execution time when we use 8 data nodes.

7. Conclusions

We proposed an adaptive and parallel algorithm, MR-
Graph, which efficiently processes multiple query graphs over a massive set of data graphs based on MapReduce. AdaptiveTune improves query processing as it adaptively tunes a feature size at runtime, keeping the system in a balance between the effectiveness of graph filtering and the cost of feature extraction. Our experiments show that AdaptiveTune performs with very little overhead in execution time.

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