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Parallelizing Graph Structural Recursion with BSP

Chong Li†  Le-Duc Tung‡  Nhat-Tan Duong§
Soichiro Hidaka†‡  Zhenjiang Hu†‡

†National Institute of Informatics, Japan
‡Graduate University of Advanced Studies, Japan
§Hanoi University of Science and Technology, Vietnam

{chong, tung, hidaka, hu}@nii.ac.jp  dn.nhattan@gmail.com

Abstract

The ever-increasing size of data today creates a critical need for scalable systems that can process large data efficiently. Graph structure can scale naturally to large datasets, as it does not require expensive join operations that are often needed by relational database querying. However, a complex query on a large graph is still very expensive in computation. Moreover, optimizing algorithms of different queries is still needed to study case by case. Queries can be expressed by structural recursion like first-order logic extended with transitive closures. It gives us a new thinking that graph queries can be generalized in a structural-recursion way. Therefore, querying can be systematically evaluated in an efficient way by optimizing the structural recursion. In this paper, the structural recursion is first time implemented in parallel. A novel framework, based on Bulk Synchronous Parallelism (BSP), is thus proposed to evaluate large-graph queries. It provides a systematic way to deal with general graph queries. The implementation of the framework puts the structural recursion into practice and completes many unclear parts that cannot be covered by the theory. The performance evaluation shows that BSP can handle large-graph querying efficiently with a good scalability. The validation of our framework is an important step towards a systematic development of algorithms on large distributed graphs in which we can apply rules to automatically reasoning about programs.

Keywords: Structural recursion, Graph querying, Distributed computing, BSP

1 Introduction

The ever-increasing size of data today creates a critical need for scalable systems that can process large data efficiently. Graphs are useful to manage ad-hoc and evolving data like the World Wide Web [AJB99], social network [CSW05], biologic information [MV07], transportation system [APT04], etc. Moreover, graph structure can scale naturally to large datasets, as it does not require expensive join operations. However, a complex query on a large graph is still very expensive in computation. Efficient processing of large graphs becomes an open problem and more and more emergent in today’s big-data era.

Researchers proposed different algorithms for challenging graph computing problems such as shortest paths, clustering, page ranking, minimum cut, connected components, triangle counting, community detection, maximum flow, and so on. Many of the above algorithms were studied and implemented into single-computer-processing libraries like the Boost Graph Library (BGL) [SLL01], LEDA [Meh99], the Stanford GraphBase [Knu93], or JDSL [GHHT99]. However, such libraries were designed to handle the graphs that can be fitted into a single computer’s memory. The scale of problems is therefore limited by the memory of machine, since they cannot be directly used on distributed architecture for large distributed graphs. A parallel/distributed platform is desired for handling large-graph processing.
Inspired by the complexity theory of PRAM model [GMR94] of parallel computers, Leslie Valiant [Val90] introduced the Bulk Synchronous Parallel (BSP) model. The BSP parallel algorithms can be designed and measured by taking into account not only the classical balance between time and parallel space (hence the number of processors) but also communication and synchronization. A BSP computation is a sequence of so-called supersteps. Each superstep combines asynchronous local computation with point-to-point communications that are coordinated by a global synchronization to ensure coherence and determinism. BSP provides a clear and sequential view of the parallel system, that is very efficient for designing complex parallel algorithms. Inspired by BSP, Google [MAB+10] developed Pregel, a system for large-scale graph processing. Several open-source Pregel-like systems [SYK+10, Ave11, SW13] were after proposed too. These systems offer the possibility to design efficient parallel algorithms for large-graph processing. However, graph algorithms are still needed to be studied case by case. Implementing various algorithms is a very labour-intensive work. Moreover, there is no systematic guidance for designing efficient algorithms. Everything is still based on the experiences of developer. How to provide a systematic way to deal with large-graph queries is still an open problem.

Buneman et al. [BFS00] introduced a simple and powerful query language for graphs using structural recursion, which allows queries being expressed by first-order logic extended with transitive closures. It gives us a new thinking that graph queries can be generalized in a structural recursive way. Therefore, instead of studying graph algorithms case by case, handling the structural recursion can systematically optimize large-graph querying. However, Buneman’s system “is adequate for small input graphs, e.g., graphs with at most 1000 nodes and 10000 edges, but it does not scale to large graphs with 10000 or more nodes”. There is still a big gap between structural recursion and parallel system. Parallel implementation of structural recursion does not exist. Previous researches discussed only the possibility of parallelizing structural recursion in theory briefly. We do not know how the evaluation works in parallel precisely in practice. Moreover, those theoretical proposals were never experimented. Implementing structural recursion on a parallel platform becomes an essential challenge, that is an important step towards a systematic development of parallel algorithms on large graphs.

![Diagram of General Graph Query and Parallel Execution](image)

**Figure 1: Overview of our parallel framework**

In this paper, we attempt to bridge the gap between structural recursion and parallel system, by proposing a parallel framework in which general graph queries can be optimized systematically for large-graph processing. Figure 1 is an overview of our framework. The first step for bridging the gap is to study how a graph general query can be translated into parallelizable structural
recursion, and how such structural recursion can be evaluated efficiently in parallel.

This paper is organized as follows: Section 2 reviews the graph data model and the structural recursion proposed by Buneman et al. [BFS00]. Section 3 exploits the parallelizability of structural recursion. In Section 4, we show how to implement efficient parallel structural recursion evaluation by using Bulk-Synchronous ML (BSML), a variant of CAML programming language developed by Loulergue et al. [Lou00]. The scalability of our implementation are studied in Section 5. The related work is discussed in Section 6 and the paper is concluded by Section 7.

Our contributions in the paper are: 1) studying the parallelizability of structural recursion with semantics inference rules; 2) developing the first parallel implementation of structural recursion evaluation, where we discover several optimizations and have improved the performance of evaluation; 3) validating the scalability of parallel evaluation of structural recursion, which encourages future researches on efficient structural recursion; and 4) proposing a general framework that can systematically process large-graph queries in an efficient way.

2 Graph Structural Recursion

Graph structural recursion, proposed by Buneman et al. [BFS00], is powerful with its simplicity and expressibility. In this section, we will review the graph structural recursion and its graph data model. A method for translating select-where queries into structural recursion is also presented in this section. Reader can, if he/she already knows Buneman et al.’s work on [BFS00], skip this section and go directly to next section.

2.1 Graph Data Model

Graphs we deal with in this paper is rooted, directed, and edge-labelled with no order on outgoing edges. They are edge-labelled in the sense that all information is stored on labels of edges while labels of nodes serve only as a unique identifier without a particular meaning. Our graph data model has two prominent features, markers and \(\epsilon\)-edges. Nodes may be marked with input and output markers, which are used as an interface to connect them to other graphs. An \(\epsilon\)-edge represents a shortcut of two nodes, working like the \(\epsilon\)-transition in an automaton. We use \(\text{Label}\) to denote the set of labels and \(\mathcal{M}\) to denote the set of markers.

Formally, a graph \(G\), sometimes denoted by \(G_{(V,E,I,O)}\), is a quadruple \((V,E,I,O)\), where \(V\) is a set of nodes, \(E \subseteq V \times (\text{Label} \cup \{\epsilon\}) \times V\) is a set of edges, \(I \subseteq \mathcal{M} \times V\) is a set of pairs of an input marker and the corresponding input node, and \(O \subseteq V \times \mathcal{M}\) is a set of pairs of output nodes and associated output markers. For each marker &\(x\) \(\in \mathcal{M}\), there is at most one node \(v\) such that \((&x, v) \in I\). The node \(v\) is called an input node with marker &\(x\) and is denoted by \(I(\&x)\). Unlike input markers, more than one node can be marked with an identical output marker. They are called output nodes. Intuitively, input nodes are root nodes of the graph (we allow a graph to have multiple root nodes, and for singly rooted graphs, we often use default marker & to indicate the root), while an output node can be seen as a “context-hole” of graphs where an input node with the same marker will be plugged later. We write \(\text{inMarker}(G)\) to denote the set of input markers and \(\text{outMarker}(G)\) to denote the set of output markers in a graph \(G\). In addition, we write \(\text{label}(\zeta)\) to denote the label of the edge \(\zeta\).

For instance, the graph in Figure 2 is denoted by \((V,E,I,O)\) where,

\[
V = \{0, 1, 2, 3, 4, \ldots, 40, 41\}
\]

\[
E = \{(41, \text{Paper}, 40), (40, \text{pubven}, 33),
(40, \text{reference}, 35), (35, \text{Paper}, 39),
(41, \text{Paper}, 39), (39, \text{title}, 30), \ldots\}
\]

\[
I = \{(&, 41)\}
\]

\[
O = \{\}\.
\]

To describe arbitrary (directed, edge-labelled, and rooted) graphs, we use here the following nine graph constructors proposed by Buneman et al. [BFS00]:
Figure 2: Graph of a digital library with 3 papers

\[
G := \begin{cases}
\emptyset & \text{(single node graph)}
| \\
a : G & \text{(an edge pointing to a graph)}
| \\
G_1 \cup G_2 & \text{(graph union)}
| \\
&x := G & \text{(label the root node with an input marker)}
| \\
&y & \text{(a node graph with an output marker)}
| \\
() & \text{(empty graph)}
| \\
G_1 \oplus G_2 & \text{(disjoint graph union)}
| \\
G_1 @ G_2 & \text{(append of two graphs)}
| \\
cycle(G) & \text{(graph with cycles)}
\end{cases}
\]

As showing in Figure 3, \(\emptyset\) constructs a root-only graph, \(\{a : G\}\) constructs a graph by adding an edge with label \(a \in Label \cup \{\epsilon\}\) pointing to the root of graph \(G\), and \(G_1 \cup G_2\) adds two \(\epsilon\)-edges from the new root to the roots of \(G_1\) and \(G_2\). Also, \&\(x := G\) associates an input marker, \&\(x\), to the root node of \(G\), \&\(y\) constructs a graph with a single node marked with one output marker \&\(y\), and () constructs an empty graph that has neither a node nor an edge. Further, \(G_1 \oplus G_2\) constructs a graph by using a componentwise \((V, E, I)\) union. Operator \(\cup\) differs from \(\oplus\) in that \(\cup\) unifies input nodes while \(\oplus\) does not. Operator \(\oplus\) requires input markers of operands to be disjoint, while \(\cup\) requires them to be identical. \(G_1 @ G_2\) composes two graphs vertically by connecting the output nodes of \(G_1\) with the corresponding input nodes of \(G_2\) with \(\epsilon\)-edges, and cycle\((G)\) connects the output nodes with the input nodes of \(G\) to form cycles. Newly created nodes have unique identifiers.

Figure 3: Graph Constructors

For example, the graph showing in Figure 2, that respects DBLP citation dataset schema [TZY+08], can be constructed by using the above graph constructors:

\[
&dl \ @
\]
\[
cycle((
&dl := \{\text{Paper:}\&p1, \text{Paper:}\&p2, \text{Paper:}\&p3\},
&p1 := \{\text{title:} \text{"SGL"}, \text{year:} \{\text{Int:}2011\},
\text{authors:} \text{"C. Li and G. Hains"}\},
\ldots
\}}
\]

\[
\]
Here `&dl@` which is used to declare that `&dl@` is the root input marker (by the convention root input marker is normally named `&`) and remove other input markers (e.g. `&p1`, `&p2` and `&p3` in our example) from the graph.

### 2.2 Graph Structural Recursion

Recursion is widely used by functional programming language for traversing dataset. However, different from list and tree structures, graph structure is much more general and complex, since it can include cyclic structure. A recursion without restriction might loop infinitely on such structure. That’s why we use a restricted form of recursion – structural recursion – to deal with the general structure. The restrictions of structural recursion are to ensure the termination of recursion.

A function \( f \) on graphs is called a structural recursion if it is defined by the following equations:

\[
\begin{align*}
  f(\{\}) & = \{\} \\
  f(\{l : g\}) & = e \circ f(g) \\
  f(g_1 \cup g_2) & = f(g_1) \cup f(g_2),
\end{align*}
\]

where the expression \( e \) may contain references to variables \( l \) and \( g \) (but no recursive calls to \( f \)). Since the first and the third equations are common in all structural recursions, we write the structural recursion simply as

\[
  f(db) = \text{rec}(\lambda(l,g).e(db))
\]

where \( e \) could be either graph variable (denoted by \( g \)), conditional expression (\text{if} \ l = l \text{then} \ e \text{else} \ e \), where \( l \) is either actual label or label variable \( l \), graph constructor (presented in Section 2.1) or structural recursion expression.

For example, for retrieving all papers’ conferences (pubven) in the graph of Figure 2, we can write the query in the structural recursion form as follows:

\[
\begin{align*}
Q1 := \\
  \text{rec}(\lambda(l,v).f(v)) \\
  \quad \text{if} \ l = \text{Paper} \text{then} \\
  \quad \text{rec}(\lambda(l,c).f(v)) \\
  \quad \quad \text{if} \ l = \text{pubven} \text{then} \\
  \quad \quad \{\text{Conference} : c\} \\
  \quad \quad \text{else} \ \{\} \\
  \quad \text{else} \ \{\} \\
  \quad \} \\
\end{align*}
\]

This query firstly retrieves all subgraphs \( G_1 \) that are under a \textbf{Paper} edge (\{\text{Paper} : \}) from the whole graph \( db \). Thus, each \( G_1 \) contains all information of a paper. After that, it retrieves subgraphs \( G_2 \) that are under a \textbf{pubven} edge (\{\text{pubven} : \}) from each \( G_1 \). So each \( G_2 \) is the conference information of a paper. At the end, we add a \textbf{Conference} edge before the actual information of conference by replacing the \textbf{pubven} edge. Figure 4 shows the result of the query.
2.3 Translating Queries into Structural Recursion

Writing SQL-like queries is often easier than directly writing in the structural recursion form. For example, Q1 that is used to retrieve all papers’ conference can be written in a select...where... surface syntax with pattern matching:

\[
\text{Q2 := select \{Conference : \$c\} where \{\_*.Paper.pubven : \$c\} in \$db}
\]

For any regular path pattern, according to [Suc02], it can be translated into structural recursion by expressing first the regular expression as an automaton, and associating a function with each state. For example, the regular expression \_*.Paper.pubven in Q2 can be represented by the automaton in Figure 5.

![Figure 5: Automaton for Q2](image)

The initial state is \(z_1\). If we meet a Paper-edge in \(z_1\) then it is changed to the state \(z_3\) otherwise to the state \(z_4\). The states \(z_2\) and \(z_4\) are similar to \(z_1\), will be changed to \(z_3\) if meet a Paper-edge otherwise to \(z_4\). The \(z_3\) is the state where we met a Paper-edge and waiting for a pubven-edge. Therefore its state will be remained when meet a Paper-edge but changes to \(z_2\) iff meet pubven. The state \(z_2\) is terminal state.

Based on the translated automaton, Q2 can be written like Q1 with mutually-recursive function. Such mutually-recursive function can be translated into single-recursive function through tupling, where tuples of graphs are represented by disjoint union of graphs and projections of the tuples as the idiom \&z@ where \&z is the component that are extracted by the tuples. Therefore, we obtain:
Q3 :=
&z1
@ rec(\ { $L , $c ) .
i f $L = Paper
then (&z4 :=&z3 , & z3 :=&z3 , & z2 :=&z3 , & z1 :=&z3 )
else i f $L = pubven
then (&z4 :=&z4 , & z3 :=&z4 , & z2 :=&z4 , & z1 :=&z4 )
else (&z4 :=&z4 , & z3 :=&z4 , & z2 :=&z4 , & z1 :=&z4 )
( $db )

Now, we are able to write queries in a simple select . . . where . . . syntax with regular path pattern. We know also how to translate a select-where query into structural recursion. In addition, Hidaka et al. [HHI+12] proposed an optimization for rewriting structural recursion queries based on markers analysis.

3 Parallelizability of Structural Recursion

We exploit, in this section, the parallelizability of structural recursion. The bulk semantics, which is one of solutions to evaluate structural recursion, is studied here by using inference rules. Select-where query is also studied to guarantee the parallelizability for more general case. At the end of this section, we show how to decompose a graph into a distributed one.

3.1 Bulk Semantics

There are several solutions to evaluate structural recursion by avoiding that recursive functions run into infinite loops on cyclic graph. For example, we can memorize all recursive calls then avoid entering infinite loops of recursion evaluation on graph. We can also do an inductive definition based on a constructor expression for the given data graph. However, such a definition can only be done under some technical restrictions.

Here we present a parallelizable solution called bulk semantics, which is also mentioned briefly in [BFS00], but has never been implemented or experimented in parallel environment. The method applies the recursive function individually on all edges of the input graph. Therefore, all the possible results of the recursive functions are evaluated on each edge by using its label and subgraph as inputs of the function. We simply need to reconnect the above results according to the structural recursion. Moreover, each function will be applied only as many times as edges in the graph, and infinite loops are avoided.

Let $G[a_1 : G_1]$ denotes that graph $G$ is composed of $a_1 : G_1$, and $⟨e , G⟩ → G′$ denotes that applying expression $e$ to graph $G$ gets graph $G′$. Recursive evaluation of structural recursion expression $\text{rec}(\lambda(\$l, \$g).e)$ applied to input graph $G$ can be summarized by the following inference rules:

$$\frac{⟨e , G⟩ → G′}{\text{rec}(\lambda(\$l, \$g).e , G[a_1 : G_1]) → G′}$$

As defined in Section 2.2, the operation $@$ is associative. When the evaluation of expression $e$ depends only on $a_1$ but not on $G_i$, we are able to apply body $e$ independently on every pair $(a_i, G_i)$ in $G$ where $a_i$ is the label of the edge and $G_i$ is the graph that the edge is pointing to. Once the the edges of $G$ were evaluated with body $e$, then we can join the evaluated edges with $e$-edges (as in the $@$ constructor). Therefore, we obtain fully parallelizable bulk semantics for evaluating structural recursion:

$$\frac{⟨e , G⟩ → G′}{\text{rec}(\lambda(\$l, \$g).e , G[a_1 : G_1]) → G′}$$
However, it also shows that there is a constraint for the body \( e \): when applying the bulk semantics, we do not have full information of the input graph \( G \) any more, we shall use only the information of the actual edge \( a \).

### 3.2 Decomposed Query

Indeed, the current expression of structural recursion allows the body \( e \) uses the subgraph \( g \). But what will happen to bulk semantics? Here is an example for retrieving all \( \text{Paper} \) published in 2011 from the graph of Figure 2. Since \( \text{Paper} \) is not used at the end of expression but in the middle, we need to split the regular expression \( \rightarrow \text{Paper.year.Int.}2011 \) to at least two parts (at most five parts in our case: \( \rightarrow \text{Paper.year}. \text{Int} \) and 2011) in order to construct the select-where query:

\[
\begin{align*}
\text{select} & \quad \{ \text{Paper2011}: p \} \\
\text{where} & \quad \{ \_, \text{Paper}: p \} \quad \text{in} \quad db, \\
& \quad \{ \text{year}. \text{Int}: 2011 \} \quad \text{in} \quad p
\end{align*}
\]

In this query, the final result \( (p) \) depends not only on the actual edge (where label \( p = \text{Paper} \)), but also on its subgraph where it needs to satisfy the expression \( \text{year.Int} : 2011 \).

By decomposing the above query and translating them into two automata (Figure 6 and Figure 7), we can obtain the following embedding structural recursion:

\[
\begin{align*}
& \text{kz1} \at \text{rec}(\{ (sL, sP) \}. \\
& \text{if} \quad sL = \text{Paper} \\
& \text{then} \quad (kz3 := \& kz2) \quad \text{U} \quad \text{kz1} \at \text{rec}(\{ (sL, sfv1) \}. \\
& \text{if} \quad sL = \text{year} \\
& \text{then} \quad \{ kz1 := \& kz12, \& kz12 := \& kz15, \& kz13 := \& kz15 \} \\
& \text{if} \quad sL = \text{Int} \\
& \text{then} \quad \{ kz1 := \& kz15, \& kz12 := \& kz13, \& kz13 := \& s15 \} \\
& \text{if} \quad sL = 2011 \\
& \text{then} \quad \{ kz1 := \& kz15, \& kz12 := \& s15, \& kz13 := \& s14 \quad \text{U} \quad \{ \text{Paper2011}: p \} \} \\
\text{else} \quad \{ kz1 := \& kz15, \& kz12 := \& s15, \& kz13 := \& s15 \} \\
\} \\
\text{(p)}), \\
& \text{kz2} := \& kz2 \quad \text{U} \quad \text{kz1} \at \text{rec}(\{ (sL, sfv1) \}. \\
& \text{if} \quad sL = \text{year} \\
& \text{then} \quad \{ kz1 := \& kz12, \& kz12 := \& s15, \& kz13 := \& s15 \} \\
& \text{if} \quad sL = \text{Int} \\
& \text{then} \quad \{ kz1 := \& s15, \& kz12 := \& s13, \& kz13 := \& s15 \} \\
& \text{if} \quad sL = 2011 \\
& \text{then} \quad \{ kz1 := \& s15, \& kz12 := \& s15, \& kz13 := \& s14 \quad \text{U} \quad \{ \text{Paper2011}: p \} \} \\
\text{else} \quad \{ kz1 := \& s15, \& kz12 := \& s15, \& kz13 := \& s15 \} \\
\} \\
\text{(p)}), \\
& \text{else} \quad \{ \text{kz3} := \& kz3, \& kz2 := \& kz3, \& kz1 := \& kz3 \}
\}
\end{align*}
\]

The inference rules of embedding structural recursion thus become:

\[
\begin{align*}
& \frac{(e, G) \circ (e, G) \circ \cdots \rightarrow G'[G_0 \circ G_0 \cdots] \quad \text{G}_0[G_0 \circ G_0] \cdots}{}
\end{align*}
\]

\[
\begin{align*}
& \frac{\{ e, G[a_1 : G_1] \} \circ \{ e, G'[a_2 : G_2] \} \circ \cdots \rightarrow G'[G'_0 \circ G'_0] \cdots}{}
\end{align*}
\]

\[
\begin{align*}
& \frac{\langle \text{rec}(\lambda (sL, sP). e), G[a_1 : G_1] \rangle \rightarrow G'}{}
\end{align*}
\]
where $e_g$ is the inner structural recursion $\text{rec}(\lambda(l, g).e_{inner})$ using the same bulk semantics inference rules, $e_a$ is partial expression of $e$ excluding the inner structural recursion, and $\circ$ is the relation between $e_g$ and $e_a$. Therefore, an embedding structural recursion is still parallelizable by using divide-and-conquer approach. However, information of graph need to be replicated recursively that will increase heavily communication cost in a distribute-memory environment.

There are also other cases of non-decomposed query. Since our select-where queries are using regular expression or its composition, we can always split queries down to decomposed ones in order to maintain the parallelizability of the querying. To have as less as possible the decomposed parts of a query $Q(g)$ or $Q(l, g)$, where $g$ is an input graph variable and $l$, if needed, is an input label variable, in order to have better performance, we here borrow six forms from [Suc02]. These forms are originally to restrict select-where queries on tree structure. A decomposed select-where subquery must be in one of these forms:

1. $g$;
2. $\{\}$;
3. $\{a : Q_1(l, g)\}$ where $a$ is either $l$ or a label constant;
4. $Q_1(l, g) \cup Q_2(l, g)$;
5. $\text{select } Q_1(g_1) \text{ where } \{R : g_1\} \text{ in } g$;
6. $\text{select } Q_1(l_1, g_1) \text{ where } \{R : l_1 : g_1\} \text{ in } g, P(l_1)$;

where $P(l_1)$ is a unary predicate, $R$ is regular expression, $Q_1$ and $Q_2$ are themselves restricted select-where queries with variable marked accordingly. Therefore, if a query $\text{select } E \text{ where } P \text{ in } db$ can be translated into one single structural recursion union with an input marker, which is used to indicate the initial state (e.g. $&z1$ in the structural recursion query on Page 8), then $P$ may introduce only one graph variable $g$ and, if needed, a label variable $l$ occurring right before $g$.

Let $f(G)$ be a function for querying the graph $G$, $f$ has decomposability iff:

- $\forall G_1, G_2 | f(G_1@G_2) = f(G_1)@f(G_2)$, and
In this subsection, we will study how to decompose a graph data into distributed one by extending the basic graph data using some special markers. Therefore, a very large graph data could be split into several partitions and distributed to computer cluster. These special markers are called cross links. Similar to [Suc02], cross links are used to split nodes in different partitions. A cross link is an edge $u \to v$ labelled with $\epsilon$ for which $u$ and $v$ are stored on different partitions. Therefore, a node $a$ can be split into two nodes $a$ and $a'$ with a cross link $a \leftarrow a'$.

In practice, we use input markers and output markers with the same name to describe cross links instead of creating real $\epsilon$-edges. For example, Figure 8 is a distributed graph\(^1\) with 3 partitions that contains the same information of the graph of Figure 2. The nodes 38 and 39 of Figure 2 are split here to create cross links that bridge partitions with input and output markers. Splitting the node under Paper is not mandatory, any node of the graph can be split to different partitions by using cross link.

Figure 8: Distributed Graph with 3 Partitions

\[ \forall G_1, G_2 | f(G_1 \oplus G_2) = f(G_1) \oplus f(G_2). \]

It provides the possibility to evaluate queries in parallel by decomposing the database.

### 3.3 Distributed Graph

In this subsection, we will study how to decompose a graph data into distributed one by extending the basic graph data using some special markers. Therefore, a very large graph data could be split into several partitions and distributed to computer cluster. These special markers are called cross links. Similar to [Suc02], cross links are used to split nodes in different partitions. A cross link is an edge $u \to v$ labelled with $\epsilon$ for which $u$ and $v$ are stored on different partitions. Therefore, a node $a$ can be split into two nodes $a$ and $a'$ with a cross link $a \leftarrow a'$.

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In general, a distributed graph $G$ can be expressed as:

\[ &z \circ \text{cycle}_{z}(G_1 \oplus \cdots \oplus G_p) \]

where $z$ is a set of input markers $z = \{ &z_1, \ldots, &z_n \}$ whose size is the number of cross links plus one, $&z$ is the root input marker, and $G_1, \ldots, G_p$ are the subgraphs on $p$ partitions.

For example, $\&dl \circ \text{cycle}(db_1 \oplus db_2 \oplus db_3)$ is equal to the graph of Figure 2, where $db_1$, $db_2$ and $db_3$ are the 3 partitions of the distributed graph shown in Figure 8:

\[ \begin{align*}
&db_1 := \\
&\&dl \circ \text{cycle}((
&\&dl := \{ \text{Paper} : &p1, \text{Paper} : &p2, \text{Paper} : &p3 \}, \\
&\&p1 := \{ \text{title} : \{ \text{String} : "SGL" \}, \text{year} : \{ \text{Int} : 2011 \}, \\
&\&authors : \{ \text{String} : "C. Li and G. Hains" \}, \\
&\&pubven : \{ \text{String} : "HPCS" \}).
\end{align*} \]

\(^1\)The ID of nodes is local for each partition. For the full graph, it is needed to add partition ID to each node for distinguishing the nodes that have same node ID but at different partitions.
Therefore, for a decomposed function $f(G)$, with which we know $[Suc02]$ $f(\text{cycle}(G)) = \text{cycle}(f(G))$, can be applied on a distributed graph $G$ (e.g. the one in Figure 2), then we obtain

$$f(kz_1)@\text{cycle}_z(f(G_1) + \cdots + f(G_p))$$

Bulk semantics of structural recursion on distributed graph work also well in parallel:

$$\left(\left(\left(e, G[a_1^1 : G_1]\right) + \cdots \right) + \left(\left(e, G[a_p^1 : G_p]\right) + \cdots \right)\right) \rightarrow G'[G_1^0 + \cdots + G_p^0]$$

We are now able to store very large graph in a distributed environment by using cross links. We will describe the implementation of the parallel framework for structural recursion in next section.

## 4 Structural Recursion over BSP

GRoundTram$^2$ [HHI+11], which is a system to build bidirectional transformation between two models (graphs), uses the basic graph data model presented in Section 2. In this section, we will use BSML to extend the sequential structural recursion of GRoundTram to parallel one. The parallel bulk semantics is first time implemented in practice. Thanks to the clarity of BSP model, the performance of our parallel implementation can be understood easily.

### 4.1 Bulk-Synchronous Parallelism

#### 4.1.1 BSP Model

The Bulk-Synchronous Parallel (BSP) model is a parallel programming model introduced by Leslie Valiant [Val90]. It offers a high degree of abstraction like PRAM models [FW78] and yet allows portable and predictable performance on a wide variety of multi-processor architectures [SHM97]. The major difference between BSP and PRAM is that the local computations of BSP are asynchronous, and the cost of inter-processor communications in BSP is not neglected.

A BSP computer contains:

- a homogeneous set of uniform processor-memory pairs;
- a communication network allowing inter-processor delivery of messages;
- and a global synchronization unit which executes collective requests for a synchronization barrier.
A wide range of actual architectures can be simulated by a BSP computer, including shared-memory machines (via BS-PRAM \cite{Tis96}). Moreover, the synchronization unit is very rarely a hardware but rather a software event \cite{HS98}. Supercomputers and clusters of PCs can be modelled as BSP computers.

A BSP program is executed as a sequence of supersteps (Figure 10), each one is divided into (at most) three successive and logically disjoint phases:

1. In the first phase, each processor uses its local data (only) to perform sequential computations and to request data transfers to/from other nodes.
2. In the second phase, the network delivers the requested data transfers.
3. And in the third phase, a global synchronization barrier occurs, making the transferred data available for the next superstep.

The performance of the BSP machine is characterised by 4 parameters:

- the local processing speed \( r \);
- the number of processor-memory pairs \( p \);
- the time \( L \) required for a global synchronization (barrier);
- and the time \( g \) for collectively delivering a 1-relation communication phase where every processor receives/sends at most one word.

The network can deliver an \( h \)-relation (every processor receives/sends at most \( h \) words) in time \( g \times h \). To accurately estimate the execution time of a BSP program these 4 parameters could be easily benchmarked \cite{Bis04}.

\footnote{Source codes of GRoundTram are available online at \url{http://www.biglab.org/download.html}.}
The execution time (cost) of a superstep $s$ is the sum of the maximum local processing time, the data delivery and the global synchronisation times. It is expressed by the following formula:

$$Cost(s) = \max_{0 \leq i \leq p} \{w_i^s\} + \max_{0 \leq i \leq p} \{h_i^s \times g\} + L$$

where $w_i^s$ is the local processing time on processor $i$ during superstep $s$, and $h_i^s = \max\{h_i^{s+}, h_i^{s-}\}$ where $h_i^{s+}$ (resp. $h_i^{s-}$) is the number of words transmitted (resp. received) by processor $i$ during superstep $s$.

The total cost of a BSP program composed of $S$ supersteps is $\sum_S Cost(s)$. It is, therefore, the sum of three terms:

$$W + H \times g + S \times L$$

where $W = \sum_S \max_i \{w_i^s\}$ and $H = \sum_S \max_i \{h_i^s\}$.

4.1.2 BSML Programming

BSML$^3$ [LGB05] developed at Université d’Orléans and Université Paris-12 (now called Université Paris-Est Créteil or UPEC) is a library for OCaml$^4$ implementing partially the Bulk Synchronous Parallel ML language [Gav05]. There is in BSML an abstract polymorphic type $\alpha$ par which represents the type of $p$-wide parallel vectors of values of type $\alpha$, one per process. It is very different from usual SPMD programming where messages and processes are explicit, and programs may be non-deterministic or may contain deadlocks. In fact a large subset of BSML parallel programs are purely functional. The core BSML library is based on the following primitives:

- $\text{mkpar} : \langle \text{int} \rightarrow \alpha \rangle \rightarrow \alpha$ par
- $\text{proj} : \alpha$ par $\rightarrow \langle \text{int} \rightarrow \alpha \rangle$
- $\text{apply} : \langle \alpha \rightarrow \beta \rangle$ par $\rightarrow \alpha$ par $\rightarrow \beta$ par
- $\text{put} : \langle \text{int} \rightarrow \alpha \rangle$ par $\rightarrow \langle \text{int} \rightarrow \alpha \rangle$ par

The semantics of BSML primitives is described by the use of parallel values. Parallel value $<x_0, x_1, \ldots, x_{p-1}>$ represents a set of local values of a given type, such that $x_i$ is stored on processor $i$ and $p$ is the number of processors.

In BSML, $\text{mkpar}$ is the parallel constructor: $\text{mkpar}$ $f$ computes the value $<f\,0, f\,1, \ldots, f\,(p-1)>$. $\text{proj}$ is the parallel destructor: $\text{proj}$ $<x_0, x_1, \ldots, x_{p-1}>$ computes a function $f$ such that $(f\,i) = x_i$. $\text{apply}$ is the asynchronous parallel transformer: $\text{apply}$ $<f_0, f_1, \ldots, f_{p-1}> <x_0, x_1, \ldots, x_{p-1}>$ computes $<f_0 x_0, f_1 x_1, \ldots, f_{p-1} x_{p-1}>$.

Finally, $\text{put}$ is the synchronous (communicating) parallel transformer: $\text{put}$ $<g_0, g_1, \ldots, g_{p-1}>$ computes a parallel vector of functions that contains the transported messages that were specified by the $g_i$. The input local functions are used to specify the outgoing messages thus: $g_i$ $j$ is the value that processor $i$ wishes to send to processor $j$. The result of applying $\text{put}$ is a parallel vector of functions dual to the $g_i$: they specify which value was received from a given distant processor.

The execution of $\text{mkpar}$ is purely local and so is the execution of the $\text{apply}$ primitive. The execution of $\text{proj}$ uses an all-to-all communication and the execution of $\text{put}$ is a general BSP communication (any processor-processor relation can be implemented with it). Recently, Li and Hains [LH14] proposed a new function named $\text{gps}$ to simplify the $\text{put}$ function. This new function separates the meta data and actual data, and provides a sequential view to program parallel communication.

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$^3$Bulk Synchronous Parallel ML. http://traclifo.univ-orleans.fr/BSML/

$^4$Objective Caml. http://caml.inria.fr/
4.2 Parallelizing Input and Output

Our parallel implementation takes an input graph and a structural recursion query, and produces an output graph. Different from GRoundTram, the input and output graphs are both distributed using cross links presented in Section 3.3 and stored in different files. Therefore, large graphs can be efficiently loaded and saved in parallel by different processor with parallel I/O.

Naturally, a large input graph is always split in advance by database administrator and distributed on different nodes of a parallel machine. In our implementation, a distributed input graph with \( p \) partitions (like Figure 8 which is distributed into 3 partitions) is stored in \( p \) files. Therefore, we have distributed input graph

\[
G_1 \oplus G_2 \oplus \cdots \oplus G_p
\]

where \( G_i (i = 1 \ldots p) \) is a local subgraph in partition \( i \).

We want each partition is handled by a compute node (i.e. processor-memory pairs in BSP model). So BSML function \( \text{mkpar} \) is used to load the distributed graph into system:

\[
\text{mkpar} \left( \text{fun \ } \text{pid} \rightarrow \text{load\_subgraph } G_{\text{pid}+1} \right)
\]

Therefore, partitions \( G_1, G_2, \ldots, G_p \) are mapped to processors \( \text{pid} = 0, \text{pid} = 1, \ldots, \text{pid} = p - 1 \):

\[
< G_1, G_2, \ldots, G_p >
\]

In fact, the node IDs in each partition are used locally. But after loading the graph, node ID will be combined with partition ID for distinguishing nodes that have the same local ID from different partitions. This step does not produce any communication cost, and the computation costs \( \max_{0 \leq i \leq p} \{ \text{load\_subgraph } G_i \} \).

The other input argument is a structural recursion query \( Q \). Here we study only decomposed queries. If the input query is a more complex one, we shall decompose it by using the technique presented in Section 3.2. Since the bulk semantics will apply the structural recursion on each edge, we also need to replicate \( Q \) in order to map it to each processor:

\[
\text{mkpar} \left( \text{fun \ } \text{pid} \rightarrow Q \right)
\]

Then we obtain

\[
< Q, Q, \ldots, Q >
\]

Since the size of the input query \( Q \) is tiny comparing to the input graph, communication cost and computation cost here for distributing \( Q \) can be neglected.

We keep the output graph also distributed, because 1) the result could be very big, for example if we want to retrieve the whole database; 2) communication cost can be reduced, since we do not need to move nodes and edges from one partition to another partition. Figure 11 is the distributed output graph created by applying \( Q_3 \) on the distributed input graph of Figure 8. We will explain how to evaluate \( Q_3 \) in the following sections. Therefore, the final result will be in the form of

\[
< G'_1, G'_2, \ldots, G'_p >
\]

A non-distributed final result could also be constructed with these \( G'_i (i = 1 \ldots p) \):

\[
G'_1 \oplus G'_2 \oplus \cdots \oplus G'_p
\]
4.3 Parallelizing Bulk Evaluation

The form of our input structural recursion is like $\&z_1@\text{rec}(\lambda(l, g).e)$, where $e$ is the recursion body, and $\&z_1$ is the root input marker. Since the bulk semantics focus only on the structural recursion evaluation, before evaluating the query, we split it into 2 parts:

$$rc = \text{rec}(\lambda(l, g).e)$$

and

$$mk = \&z_1@rc.$$ 

The bulk evaluation only take care of $rc$. $mk$ will only be used later by the epsilon elimination step presented in next subsection. This optimization technique is coherent to Buneman et al.’s sequential implementation [BFS00].

According to the bulk semantics, applying $Q_{rc}$ to $(G_1 \oplus \cdots \oplus G_p)$ is equal to applying $Q_{rc}$ to each $G_i(i = 1 \ldots p)$ individual then union the evaluated subgraphs by using $\oplus$. Therefore, we can evaluate our distributed input graph in parallel by using the BSML function:

$$\text{apply} <Q_{rc}, Q_{rc}, \ldots, Q_{rc} > <G_1, G_2, \ldots, G_p >$$

This step is equal to

$$Q_{rc}(G_1) \oplus Q_{rc}(G_2) \oplus \cdots \oplus Q_{rc}(G_p)$$

We obtain, after this step, the bulk-evaluated distributed graph

$$< G'_1, G'_2, \ldots, G'_p > .$$

This step costs $\max_{0 \leq i \leq p}\{Q_{rc}(G_i)\}$ for computation and 0 for communication.

For example, Figure 12 is the result subgraphs of partition 1 of the distributed graph in Figure 2 after applying Q3 using bulk semantics. The subgraphs that have no input marker are not included in Figure 12. Because those nodes can never be reached.

4.4 Optimizing Epsilon Elimination

After bulk evaluation, the result distributed graph contains many edges and nodes that we do not need (cf. Figure 12). For obtaining a clean clear final result (like the one in Figure 11), we need to 1) glue nodes by removing the epsilon-edges, and 2) remove the nodes that cannot be reached by the root input marker described with $mk$.

The easiest-to-design way to obtain the final result is to gather all evaluated subgraphs, then rebuild epsilon-edge of cross-links by matching the input and output markers, and at the end fetch
the reachable part of the large assembled graph. However, the intermediate result after bulk semantics evaluation is very big (c.f. Figure 12), often bigger than the original graph. Gathering and centralizing such big data is very communication-intensive and not realistic.

Since the size of final result could be very big too, our implementation keeps the final result in a distributive way. To avoid to create unnecessary communication cost, the result edges shall stay in the same partition as where the original edges were. We propose here a recursive function \textit{reachability} (Figure 13) to compute all reachable elements (nodes, edges, input markers and output markers) from source sub-graph. This function takes 4 input arguments: 1. local source sub-graph, 2. map of (node \rightarrow outgoing edges) according to the local source sub-graph, 3. global reachable output markers accumulator (initialized with root input marker), 4. reachable graph elements accumulator (initialized to empty); and produces a quadruple of nodes, edges, input markers and output markers to create the reachable subgraph(s) of each partition. The function is processed as follows:

1. locally compute the reachable elements of local graph according to the markers from marker accumulator (line 2), the result is a tuple of graph elements accumulator and output marker accumulator;

2. separate the graph elements accumulator and the marker accumulator (lines 3 - 4);

3. gather all cumulated reachable markers and broadcast to all processors (line 5);

4. check if there are new found output markers that we did not compute (line 6);

5. if there is no more new untested marker, then we stop the recursion, otherwise we make the recursive call (lines 7 - 8).

In the \textit{reachability} function, \textit{parfun} : (\alpha \rightarrow \beta) \rightarrow \alpha \ par \rightarrow \beta \ par is equal to \texttt{fun } f \ a \rightarrow \texttt{apply} (\texttt{mkpar}(\texttt{fun} \ pid \rightarrow f))\ a; \textit{parfun2} : (\alpha \rightarrow \beta \rightarrow \gamma) \rightarrow \alpha \ par \rightarrow \beta \ par \rightarrow \gamma \ par is equal to \texttt{fun } f\ a\ b\ \rightarrow \texttt{apply} (\texttt{apply} (\texttt{mkpar}(\texttt{fun} \ pid \rightarrow f))\ a)\ b; \textit{parfun4} : (\alpha \rightarrow \beta \rightarrow \gamma \rightarrow \delta \rightarrow \epsilon) \rightarrow \alpha \ par \rightarrow \beta \ par \rightarrow \gamma \ par \rightarrow \delta \ par \rightarrow \epsilon \ par is same as \textit{parfun} and \textit{parfun2} but with 4 input parameters instead; and \textit{fold.direct} : (\alpha \rightarrow \alpha \rightarrow \alpha) \rightarrow \alpha \ par \rightarrow \alpha \ par is a collective communication function based on \texttt{put} that takes a binary function \oplus and a parallel
let rec reachability = fun sourceGraphs outgoEdgeMap markersCumul graphEltsCumul ->
  let locCumulReached = parfun4 localReachable sourceGraphs outgoEdgeMap markersCumul graphEltsCumul in
  let reachedGraphElts = parfun (fun (g, m) -> g) locCumulReached in
  let reachedMarkers = fold direct SetofMarker.union reachedMarkers in
  if (proj stopCond 0) then reachedGraphElts else reachability sourceGraphs outgoEdgeMap reachedMarkers reachedGraphElts

Figure 13: BSML code of compute function

value <v_0, v_1, ..., v_{p-1}>, then produces a parallel value <s_0, s_1, ..., s_{p-1}> where s_0 = ... = s_{p-1} = v_0 \oplus ... \oplus v_{p-1} and p is the number of processors. Communication cost of fold_direct is (p-1) \times n \times g + l, where n is the average size of values v_i, g is the cost that one processor transfers a word to another processor, and l is the cost of synchronizing all processors. The initial values of markersCumul is a set with one element which is the root input marker (left side of mk, e.g. \&z1 of \&z1@rc); and the value of sourceGraphs is the bulk-evaluated distributed graph <G_1', G_2', ..., G_p'> but in quadruple form (nodes, edges, input markers and output markers).

Removing ε-edges is the most expensive computation. We need to compute transitive closure for all ε-edges, and the transitive closure computation is very costly. The ε-edge removing is processed by adding new edges that glue nodes between ε-edges to skip the ε-closure. Removing ε-edges does not require information for neighbour partition, Our implementation uses simply apply to perform this step in parallel.

Sequential GRoundTram first perform the ε-edges elimination then the reachability. However, after our experiments and performance analysis, the optimal way, both for sequential and parallel computing, is to eliminate the unreachable parts first, then perform the epsilon elimination only on the reachable parts. Therefore, in our implementation, after the bulk evaluation, we firstly compute the reachability and remove the unreachable parts, then remove the ε-edges in parallel for each cleaned partition.

5 Evaluation

The implementation of structural recursion evaluation querying graphs in parallel is experimented and will be discussed in the section.

5.1 Environment

The parallel machine used for the experiments is a computer cluster at the International Institute for Computational Science and Engineering, HUST. This supercomputer consists of 1 rack with 15 compute nodes, where we use 5 of compute nodes to evaluate the efficiency and scalability of our parallel implementation. Each compute node is configured with:

- 4 Intel Xeon E5-2679 CPUs @ 2.6GHz (16 cores per node)
- 32 GB RAM @ 1600 MHz and 300 GB HDD
- FDR 56Gbs Infiniband
- CentOS 6.5 64-bit

The following software are also deployed on the cluster, and loaded into the environment of each compute node:

\[^{6}\text{Hanoi University of Science and Technology}\]
• OpenMPI version 1.8
• Objective Caml (OCaml) v 4.0.2
• Bulk Synchronous Parallel ML (BSML) library v 0.4+beta2
The BSML library, which is based on OCaml, supports both TCP and MPI in parallel mode. Here, BSML uses OpenMPI for its communication.

5.2 Datasets
Datasets to be queried shall be a rooted, directed, and edge-labelled graph distributed into partitions and connected via cross links. All information is stored on label of edges, while label of nodes has no particular meaning but serve only as a unique identifier. This distributed graph can be in any shape including cyclic graph and tree. Since in bulk semantics a graph is evaluated edge by edge individually, the shape of graph does not have impact to the performance of evaluation, but it could indeed affect in some case the performance of epsilon-closure computation.

We created a generator to produce distributed random graphs. Our generated distributed random graph is cyclic, and each edge is labelled with a randomly chosen letter which is one of 26 Latin letters between “a” to “z”. Three random datasets were generated (Table 1). They were split into 2, 4, 8 and 16 partitions in equilibrium.

<table>
<thead>
<tr>
<th>Partitions</th>
<th>Dataset 1</th>
<th>Dataset 2</th>
<th>Dataset 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>(Total)</td>
<td>Nodes</td>
<td>Edges</td>
<td>Nodes</td>
</tr>
<tr>
<td>160 000</td>
<td>198 499</td>
<td>128 000</td>
<td>129 810</td>
</tr>
<tr>
<td>80 000</td>
<td>88 250</td>
<td>64 000</td>
<td>64 950</td>
</tr>
<tr>
<td>40 000</td>
<td>44 125</td>
<td>32 000</td>
<td>32 475</td>
</tr>
<tr>
<td>20 000</td>
<td>22 063</td>
<td>16 000</td>
<td>16 238</td>
</tr>
<tr>
<td>10 000</td>
<td>11 032</td>
<td>8 000</td>
<td>8 119</td>
</tr>
</tbody>
</table>

5.3 Queries
Queries can be written in select... where... form with regular path patterns then translated into structural recursion as explained in Section 2.3. Here, we write queries directly in structural recursion form for having a clearer understanding. All queries described below are used for retrieving information from our distributed random graphs described in previous section. Evaluation of the queries is discussed in next section.

• **Query 4**: Replace all vowels (a, e, i, o, u) by 1, 2, 3, 4, 5 respectively, and keep the consonants (other than vowels) as they are.

```ocaml
let zI @
rec($l, $g) =
  if $l = a then zI := \{1; zI\}
  else if $l = e then zI := \{2; zI\}
  else if $l = i then zI := \{3; zI\}
  else if $l = o then zI := \{4; zI\}
  else if $l = u then zI := \{5; zI\}
  else zI := \{$l; zI\}
  ) ($db)
```

• **Query 5**: Select all $x$ and $y$ that are reachable by an $i$ or a $j$ no matter after how many steps, but in between of the path we could not have $a$ or $b$. If there are more than one path from an $i$ to an $x$ and at least one of these paths does not contain $a$ or $b$ then this $x$ is reachable.
\( \text{rec}( (l, g). \quad \text{if } l = i \text{ or } l = j \quad \text{then } (z1 := z2, &z2 := &z2) \quad \text{else if } l = a \text{ or } l = b \quad \text{then } (z1 := z1, &z2 := &z1) \quad \text{else if } l = x \text{ or } l = y \quad \text{then } (z1 := z1, &z2 := \{ l; &z2 \}) \quad \text{else } (z1 := z1, &z2 := &z2) ) (db) \)

This query can be considered as an automaton with 3 states: initial state where we did not reach any \(i\) or \(j\); active state after reaching an \(i\) or \(j\) where it can be back to initial state if touch a \(a\) or \(b\); and the final state when reach \(x\) or \(y\). However, the final state has the same conditions as the active state: 1) back to the initial state if touch a \(a\) or \(b\) 2) maintain the state otherwise. That’s why we merged these two states into one.

### 5.4 Experiments

We experimented both Query 4 and Query 5 on the computer cluster described in Section 5.1. First, we fix the size of querying distributed graph and vary the number of processors. The size of subgraph in each partition will be decreased when the number of processors increases for keeping the size of the whole distributed graph consistent. After that, we fix the number of processors and vary the size of distributed graph. In this step, when the size of distributed graph increases, the size of each partition’s sug-graph increases too. Meanwhile, the number of cross links increases in the same time, since more nodes are spilt into different partitions. During our experiments, each processor processes one partition data. For each experiment, the execution times of bulk evaluation, reachability and epsilon elimination are measured separately.

![Figure 14: Execution time of experiments](image)

Query 4 is equivalent to an automaton with only 1 state. During bulk evaluation, each edge compare its label, then either relabel it if it is a vowel, or keep the consonant otherwise. Therefore, no edge will be deleted by transforming to \(\epsilon\)-edge. Epsilon elimination is very fast.
for this query. Since no edge is deleted, there is no long and deep chained $\epsilon$-edges but just only 1-step $\epsilon$-edges created during bulk evaluation. The execution time of reachability computation, which is used to clean the unreachable parts, is very short too for this query. As we know, this “automaton” has only 1 state, the number of candidate edges were therefore not multiplied. We need just to pass one and only one time for each edge and the number of edges is just same as the original graph. Figure 14 (a) (b) and (c) show the execution time of Query 4 by varying the number of processors (partitions) and the size of querying graph.

Figure 15: Automaton equivalent to Query 5

Query 5 fetches only the reachable $x$-edges and $y$-edges according to the conditions, all other edges will be contracted. The reachability computation is doubled because there is two states so each edge becomes two edges depending on states of automaton (Figure 15). In fact, during bulk evaluation, all edges are multiplied according to the number of state that the equivalent automaton has. Therefore a complex query will be very expensive. The epsilon elimination here is extremely expensive, because the transitive closure computation is very expensive by its nature. Furthermore, there are many long chained $\epsilon$-edges and the number of edges were already multiplied. That’s why, the sequential implementation of structural recursion evaluation proposed by Hidaka et al. [HHI+11] cannot handle epsilon elimination for large graphs. Our parallel implementation processes first the reachability computation then the epsilon elimination after. It optimized significantly the performance. Moreover, the parallel reachability computation is also optimized by using accumulators and collective communication via MPI. That’s why we can obtain a linear speedup, and it is even faster than other implementation in sequential without using parallelism features. Figure 14 (d) shows the execution time of Query 5 in varying the size of graph.

The results of experiments confirm the theoretical proposals and assumptions. Our implementation have linear speedup and sometimes super linear speedup for some complex queries since the reachability computation and epsilon elimination are optimized for both sequential computing and parallel computing. The validation of our implementation is an important step towards a systematic development of algorithms for large graph querying. More research can therefore be established based on this gratified results.

6 Related Work

Evaluating regular path queries on distributed, rooted, edge-labelled directed graphs were studied by Dan Suciu [Suc02]. Theoretically, algorithms in [Suc02] are efficient in terms of the total number of computation steps and the total number of data transferred during computation ($O(n^2)$, where $n$ is the number of cross-links between different sites). Later, Shoaran et. al [ST09] proposed an iterative approach with the same data communication complexity. Tung et. al [TNVH13] showed that $O(n^2)$ for data communication is not practical to big datasets such as Youtube, DBLP, and designed an efficient library using MapReduce framework. Queries in [TNVH13] are regular path queries.

Pregel [MAB+10] is a distributed programming framework to deal with very big graphs. It has focused on providing users with a natural API for programming graph algorithms, while
invisibly managing details of distribution such as message passing and fault tolerance. It is also inspired by the Bulk Synchronous Parallel model [Val90], which provides its synchronous superstep model of computation and communication. Pregel’s concepts have been cloned by several open source projects such as: Apache Hama [SYK10], Giraph [Apa13], Signal/Collect [SBC10]. Since Pregel is a vertex-centric model where vertices are first-class citizens, it is non-trivial to apply it to queries on edge-labelled graphs.

Pig [GNC09] and Hive [TSJ09] are two popular high-level dataflow systems on top of MapReduce to analyze large datasets in the spirit of SQL. Programs are compiled into sequences of Map-Reduce jobs and executed in Hadoop environment. However, they are not designed mainly to support scalable processing of graph-structured data.

7 Conclusion

In this paper, we have studied the parallelizability of graph structural recursion, and shown that BSP model is a good practical model in order to parallelize a class of structural recursions on graphs that are decomposable structural recursions. This is an important step towards a systematic development of algorithms on large distributed graphs where we can apply rules to automatically reasoning about programs. We have proposed a parallel programming framework based on the BSP model to efficiently compute decomposable structural recursions, in which the core component is an extension of functions in the GroundTram by using BSML library. We also discovered that, although, in theory, transitive closure is computed before removing inaccessible part, swapping those computations is much more efficient to deal with large graphs in practice. Experiments with random graphs showed that our solution is quite efficient and scalable to large graphs.

In the future, we will extend our solution to deal with structural recursions whose $e$ function refers to the variable $g$. It is interesting, but non-trivial to solve because such structural recursions are now not decomposable. Furthermore, porting the solution to Pregel-like systems in order to process big graphs is also a challenge, because Pregel-like systems are vertex-centric model, while ours is edge-centric model.

References


