Towards Efficient Large-Scale Interprocedural Program Static Analysis on Distributed Data-Parallel Computation

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Abstract—Static program analysis has been widely applied along the whole process of the program development for bug detection, code optimization, testing, etc. Although researchers have made significant work in static program analysis, it is still challenging to perform sophisticated interprocedural analysis on large-scale modern software. The underlying reason is that interprocedural analysis for large-scale modern software is highly computation- and memory-intensive, leading to poor efficiency and scalability. In this article, we introduce an efficient distributed and scalable solution for sophisticated static analysis. Specifically, we propose a data-parallel algorithm and a join-process-filter computation model for the CFL-reachability-based interprocedural analysis. Based on that, an efficient distributed static analysis engine called BigSpa is developed, which is composed of an offline batch static program analysis system and an online incremental static program analysis system. The BigSpa system has high generality and can support all kinds of static analysis tasks that can be expressed as CFL reachability problems. The performance of BigSpa is evaluated on real-world large-scale software datasets. Our experiments show that the offline batch system can exceed an order of magnitude compared with the most advanced analysis tools available on performance, and for incremental analysis with small batch updates on the same data sets, the online analysis system can achieve near real-time response, which is very fast and flexible.

Index Terms—Interprocedural static analysis, distributed systems, data-parallel computation

1 INTRODUCTION

In the entire process of software development, interprocedural static program analyses are widely used for tasks such as bug detection, code optimization, testing, debugging and so on.

Motivation. Nowadays, people apply the simple pattern-based or intraprocedural analysis to uncover various software problems in practice ([1], [2], [3]). However, due to the lack of considering adequately rich/complete semantics information, such simple analyses suffer from severely low accuracy (30-100 percent false positive rate [4]), leading to dead poor applicability [5]. For the sake of precise and thus useful analysis results, sophisticated interprocedural analyses are essential, e.g., interprocedural context-, path-, flow-, field-sensitive analysis. Different from the imprecise but efficient intraprocedural analysis, interprocedural analyses take into account more semantics information, resulting in high analysis complexity. For instance, in a context-sensitive interprocedural analysis, the number of calling contexts can easily exceed millions [6], making the analysis both computation- and memory-intensive. In addition, as the sizes of modern software grow, performing precise analysis on real-world modern software systems like Linux kernel, becomes more and more challenging. It is reported that in a context-sensitive interprocedural alias analysis for Linux kernel (more than 16M lines of code), there are more than 1 billion edges generated. None of the state-of-the-art analysis tools can accomplish the analysis task due to either analysis time or memory usage issues [7]. The poor scalability greatly hinders the interprocedural analysis being more widely adopted in industry.

From our point of view, the reason of poor scalability is two-fold. First, the majority of existing interprocedural analysis systems are usually based on sequential algorithms. They lack the support for highly efficient parallel and distributed computing. Second, most state-of-the-art algorithms are purely memory-based. The limited size of available memory severely impedes the applicability of static analyses in large-scale scenarios.

Researchers have been endeavouring to improve the analysis scalability by proposing approximations, abstractions, or various heuristics. However, approximations usually trade off analysis capability for scalability, rendering analysis less precise and useful. Even worse, the analysis with lower accuracy sometimes still failed to analyze large-scale systems code. In short, it is still too challenging to efficiently perform interprocedural analysis on large-scale codebases.

Approach. Previous work [7] introduces the idea of turning Big Code analysis into Big Data analytics. Unfortunately, as for the sophisticated and large-scale analysis workloads, it still severely suffers from slow convergence
In this work, we make the following contributions:

- We devise a data-parallel algorithm and a join-process-filter computation model for CFL-reachability based interprocedural analysis. Based on that, a distributed offline batch static program analysis system is implemented. To further improve performance, we propose both algorithm- and system-level optimizations, including computation closure to reduce iteration rounds, finer-grained partitioning for load balancing, and pre-shuffle to save network traffics. We also design a data structure with compression effect and manage to reduce the memory usage to one-third of the original.

- Further, to deal with mini-batch code updating scenarios, an online incremental static program analysis system supporting small batches of code update is designed and implemented. A triangle counting algorithm together with an improved data structure for online analysis is proposed to support the continuous updating of the graph topology. Considering the occurrence of large-scale computation triggered by a small batch update, we design a mechanism for the computation scale prediction and automatic computation mode switching.

- Based on the big data processing platform Spark, distributed file system HDFS and distributed in-memory database Redis, we implement a system and framework called BigSpa, integrating computation, storage and query for large-scale static program analysis. The system has high generality and can support all kinds of static analysis tasks that can be expressed as CFL reachability problems. We have made BigSpa publicly available at https://github.com/PasaLab/BigSpa.

- We empirically validate the BigSpa’s performance with extensive experiments over real-world large-scale software datasets. The experimental results show that compared with existing state-of-the-art analysis tools, our offline batch static program analysis system is one or more orders of magnitude faster. Also, for small batch updates, the online analysis system can make a near real-time response in seconds, which is very fast and flexible.

The rest of this article is organized as follows. Section 2 introduces the background of this article. Section 3 revisits the static analysis problem from the "Big Data" angle and proposes a scalable data-parallel solution. Section 4 introduces the design and optimization details of the offline batch program static analysis system. Section 5 further extends the previous section and designs an online incremental system. Section 6 introduces the design and usage details of the entire framework. The performance evaluation results are presented and analyzed in Section 7. We discuss related work in Section 8, and concludes the article in Section 9 finally.

## 2 Background

This section provides the necessary background about CFL-reachability based static analysis. We demonstrate how two typical analysis examples (i.e., pointer/alias analysis and dataflow analysis) are formulated as CFL reachability problems, and introduce the traditional worklist algorithm for CFL reachability.

### 2.1 CFL Reachability

CFL-reachability is initially proposed by Yannakakis [15] for Datalog query evaluation. Later, a large body of program analysis problems [8], [9], [16] are formulated as CFL reachability instances.

Informally, the CFL reachability computation is often guided by a context-free language \( \mathcal{L} \) over an alphabet \( \Sigma \). Given a graph \( G \), whose edges are labelled with members of \( \Sigma \). A vertex in \( G \) is \( \mathcal{L} \)-reachable from another vertex if and only if there exists a path between them, the word formed by concatenating labels along which is a member of \( \mathcal{L} \). The (all-pairs) CFL reachability problem is to determine all pairs of vertices \( v, w \) such that \( w \) is \( \mathcal{L} \)-reachable from \( v \).

### 2.2 Two Examples

We give two typical analysis examples as CFL reachability problems in the following.

**Alias Analysis.** Two variables are aliased if they point to the same memory location. Alias analysis is used to determine if two variables are aliases. Alias analysis can be formulated as CFL-reachability under the program expression graph representation of programs [16]. The analysis we implement is flow-insensitive in the sense that we do not care about the program control flow.

Based on the formulation, a program expression graph is constructed where each pointer expression (reference variable \( x \), dereference expression \( *x \), and address-of expression \( &x \)) corresponds to a graph vertex. Five kinds of three-
address statements listed below are considered to add respective edges into the graph

\[
\begin{array}{lll}
\text{Type} & \text{Stmt} & \text{Edge} \\
\text{memoryallocation} & x = \text{malloc()} & x \leftarrow m A \\
\text{assignment} & x = y & x \leftarrow y \\
\text{store} & \ast x = y & \ast x \leftarrow y \\
\text{load} & x = \ast y & x \leftarrow \ast y \\
\text{address - of} & x = \& y & x \leftarrow \& y \\
\end{array}
\]

As for a memory allocation, we add the edge from \( A \) to \( x \) labeled by \( m \). All other four types of statements would trigger the addition of an edge, accordingly. Moreover, we add dereference edge \((d)\) from each pointer variable \( x \) to \( \ast x \) and from \( \& x \) to \( x \)

\[
\begin{array}{ll}
\text{Type} & \text{Edge} \\
\text{dereference} & \ast x \leftarrow x \\
& x \leftarrow \ast x \leftarrow \& x. \\
\end{array}
\]

Given a program expression graph, the language adopted in the alias analysis is as follows:

\[
\begin{align*}
\text{Object flow} & : = m VF \\
\text{Value flow} & : = (a MA | MA | a)^* \\
\text{Memory alias} & : = \overline{d} VF d.
\end{align*}
\]

A non-terminal \( OF \) represents the points-to relation. One vertex \( x \) in the expression graph which is \( OF \)-reachable from \( o \) (i.e., there is a path from \( o \) to \( x \), along which the labels form a word of \( OF \)), means that a variable \( x \) points to an object \( o \) during execution. The definition of \( OF \) is straightforward: it must start with an alloc \((m)\) edge, followed by a \( VF \) path that propagates the object address to a variable. A \( VF \) path is either a sequence of simple assignment \((a)\) edges or a mix of assignments edges and \( MA \) (memory alias) paths. An \( MA \) path is represented by \( \overline{d} VF d \). Each edge has an inverse edge with a “bar” label. If \( o \rightarrow x \) occurs in the graph, \( x \rightarrow \overline{o} \) also exists. \( \overline{d} \) represents the inverse of a dereference and is essentially equivalent to an address-of. \( \overline{d} VF d \) represents the fact that if we take the address of a variable \( x \), propagate the address through a \( VF \) path to another variable \( y \), and then do a dereference on \( y \), the result is the same as the value in \( x \).

Fig. 1 shows a simple program and its expression graph. Solid and dashed edges are original edges in the graph and they are labeled \( m, a, \) or \( d \), respectively. Dotted edges are transitive edges added by BigSpa into the graph, as discussed shortly.

In Fig. 1, \( t \) points to \( A_3 \), since the \( m \) edge between them forms an \( OF \) path. There is a \( VF \) path from \( \& s \) to \( p \), which enables an \( MA \) path from \( s \) to \( \ast p \) due to the balanced parentheses \( d \) and \( \overline{d} \). This path then induces an additional \( VF \) path from \( \& s \) to \( q \), which, in turn, contributes to the forming of the \( VF \) path from \( n \) to \( r \), making \( \ast s \) and \( \ast r \) memory aliases. The dotted edges in Fig. 1 show these transitive edges.

**Dataflow Analysis.** Following Rep et al.’s interprocedural, finite, distributive, subset (IFDS) framework [8], we have also formulated a fully context-sensitive dataflow analysis as a grammar-guided reachability problem. Specifically, we use this dataflow analysis to track NULL value propagation. Under the IFDS framework, each dataflow fact corresponds to a vertex. The dataflow transfer function is represented as the relation mapping (edges) between vertices at different program points. Finally, an exploded supergraph is generated and the dataflow analysis is equivalent to the reachability computation on the graph. For more details, please refer to [8].

One slight difference between the IFDS framework and our formulation is that we achieve context sensitivity also by cloning intraprocedural graphs instead of using the summary-based approach in [8], which has been demonstrated [6] to fall short in answering many user queries.

### 2.3 Worklist Algorithm

Most of the state-of-the-art approaches [9], [16] implemented the CFL reachability in a search fashion. Specifically, a worklist is maintained to store the reachable vertices. Each vertex is associated with the path information, i.e., the sequence of labels along the path leading to the vertex. The process is iteratively performed, each time a vertex from the worklist is popped off and the newly reachable vertices are added into the worklist by computing the path information. However, this search-based implementation is inherently unsuitable for parallelism, especially in our case the traversal is dynamically decided [17], [18]. In addition, as each path is traversed separately from start to end and none of the intermediate path information is cached for reuse, duplicated traversals are caused.

In addition, we reviewed a dynamic programming based CFL reachability algorithm [19] which all the intermediate edges are physically added. This algorithm maintains a graph \( G \) and a list \( W \) containing new edges. Each time, (1) one edge \( e \) is popped from \( W \); (2) then it generates new edges \( e' \) based on edge \( e \) and all its adjacent edges in \( G \) if the labels on the edges match a production rule in CFL like the example shown above in Fig. 1; (3) it determines then if \( e' \) already exists in \( G \), if not, 4) add \( e' \) into both \( W \) and \( G \). It repeats the 4 steps until \( W \) becomes empty.

However, this dynamic programming based algorithm has significant drawbacks. First, the traditional algorithm is sequential, which is difficult to be directly parallelized due to a large number of read and write operations on shared variables (list \( W \) and graph \( G \)). The parallel implementation by means of locks suffers from unacceptable synchronization.
overheads and is impractical in a distributed environment due to the massive communication costs across networks. Second, the worklist algorithm needs to maintain a huge number of edges in memory. The limited memory size greatly hinders its applicability in practice.

In this work, we devise a data-parallel algorithm for CFL reachability, which is highly parallel and can leverage disks during computation.

3 A DISTRIBUTED SOLUTION

A naive parallel version of the dynamic programming algorithm [19] is to launch multiple threads to deal with multiple edges in W simultaneously. In this way, we have to achieve synchronization by aid of locks, resulting in massive synchronization overheads. Even worse, in a distributed environment, as the communications are required between tasks, a great deal of message passing occurs, which leads to prohibitive communication cost. As a matter of fact, each task only plays with a small portion of the shared data. Fully locking of the shared variables is not necessary. For the sake of low synchronization overheads, we are intended to treat the shared data at a finer granularity.

Algorithm 1. A Data-Parallel Worklist Algorithm

Data: The input graph G (a set of edges)
Result: Updated graph G with new edges
1 \(W \leftarrow G\)
2 repeat
3 \(P \leftarrow G \text{ join } W\)
4 \(E \leftarrow \text{map} (P)\)
5 \(W \leftarrow \text{distinct (E diff G)}\)
6 \(G \leftarrow W \text{ union G}\)
7 until \(W = \emptyset\)
8 return \(G\)
9 Function map \((P)\)
10 \(E \leftarrow \emptyset\)
11 foreach edge pair \(x \rightarrow y, y \rightarrow z\) in \(P\) do
12 if a production rule \(C ::= A \impliedby B\) exists then
13 \(e \leftarrow x \rightarrow z\)
14 \(E \leftarrow E \cup \{e\}\)
15 return \(E\)

Different from the task-parallel algorithm, we regard the CFL reachability as a big data problem which is composed of a large dataset and the simple computation on that dataset. With proper modeling of this problem, we can then obtain an efficient solution with good performance and scalability through cluster computing. As such, we propose a data-parallel algorithm based on Map/Reduce-style operations, shown as Algorithm 1. Specifically, the process is executed in an iterative manner. Each iteration first performs a join on \(G\) and \(W\) to produce all the possible edge pairs, e.g., \(A \rightarrow B, x \rightarrow y \rightarrow z\) (Line 3). Next, a map operation is followed to perform the label matching on each edge pair (Lines 11-14) resulting in a variety of new edges \(E\) (Line 4). As the same edge could be generated in different ways multiple times, duplicated edges thus exist in \(E\). In order to avoid redundant computation thereafter, we need to filter out all the duplicates so as to obtain the newly added edges \(W\) (Line 5). These new edges are merged to the graph \(G\) via union for the next iteration (Line 6). The iterative process continues until no new edges are produced, i.e., \(W\) is empty (Line 7).

The above data-parallel algorithm is distinct from the task-parallel one in several aspects. First, map/reduce operations intuitively partition the shared data into multiple small blocks. The data is processed in a divide-and-conquer manner, thus reducing synchronization overheads and achieving high parallelism. Second, we can choose the particular computation model according to the data locality to reduce the communication costs in the distributed environment. Finally, it is readily to utilize disks during computation by means of the mature distributed file systems.

4 OFFLINE BATCH STATIC PROGRAM ANALYSIS SYSTEM

We implemented the data-parallel algorithm in Section 3, and built an offline batch static program analysis system.

4.1 Computation Model

We propose a join-process-filter computation model in our design illustrated as Fig. 2. The whole computation is processed iteratively. At the beginning of each iteration, all the edges in both the graph \(G\) and worklist \(W\) are partitioned according to their source and destination vertices. Edges with the same source or destination vertex (i.e., the key) are put together into the same partition corresponding to the rectangle with gray background in Fig. 2. For brevity, we only use two partitions for each dataset. Note that computations for different partitions are done in parallel.

Join. Once the partitioning is done, we next join all edge partitions resulting that two edges are gathered if the source vertex of one edge is identical to the destination of the other. As shown in Fig. 2, the edges with the yellow source are connected together with edges of yellow destination forming multiple pairs of adjacent edges, e.g., \((x \rightarrow y, y \rightarrow z)\).

Process. After joining, we obtain multiple partitions, each consisting of multiple pairs of adjacent edges. We next perform a map to process all the partitions in parallel. The computation required for each partition is simply to generate new edges by matching edge labels based on the grammar rules.

Filter. When the map is completed, edges are generated in each partition shown as \(E\). Duplicates exist in the sense that the same edge could be generated multiple times. We need to filter out all these duplicated edges. Our filtering consists of two phases: (1) global filtering which removes out from \(E\) all the edges already existing in \(G\) with a distributed cache system (For example, we can store the edges of \(G\) in a distributed database such as Redis [20] and query the
elements in $E$ from the database. Only those elements with empty query results will be retained. Redis is an in-memory key-value store. It is very fast by adopting many advanced system technologies, including HashMap-based memory storage model, simple data structure, multiplexing IO, non-blocking IO and so on.; (2) "local filtering" which gets rid of all the duplicated edges in $E$ itself via a distinct operation.

After the entire join-process-filter procedure, a set of new edges $E'$ is acquired. These new edges are repartitioned and merged into $G$. Simultaneously, we replace $E$ with $E'$ for the subsequent iteration. The computation terminates until no new edges are generated (i.e., $E'$ is empty).

### 4.2 Data Structure

An edge in the program graph can be defined using a triplet $(src, dst, label)$, which represents the source node, target node and label of the edge. However, this straightforward representation is not suitable for our computation model. First, simply using triplets to store edges is quite space-inefficient. A large amount of redundancy exists. Second, at each iteration, the process part in Section 4.1 involves the join operation and label matching among edges, which requires expensive searches among a large number of triplets. Therefore, we design a compressed data structure with index optimization.

We use a key-value form to represent the edge set in a partition, with a vertex as the key and the set of edges connected to the vertex as the value. The basis for compression and optimization with indexes is that the symbols in the CFL are finite, to the vertex as the value. The basis for compression and optimization with indexes is that the symbols in the CFL are finite, which means there are only a limited number of types of edges. In practice, we can classify edges from three aspects:

1. Given a vertex, there are two directions for adjacent edges: incoming and outgoing;
2. According to the calculation state of the edge, there are two states: active (the edges to be processed) and silent (the edges that have already been processed);
3. Assuming that there are a total of $L$ terminal and non-terminal symbols in the CFL, we have $L$ kinds of labels.

Hence, all the edges associated with a certain vertex can be classified into $2 \times 2 \times L = 4L$ kinds.

Fig. 3 shows an example of the structure, where the original edge set is compressed into two arrays (the indexes of arrays start from 1 in this figure). The array of neighbours contains the nodes adjacent to the intermediate node $m$. It is ensured that all the nodes of the same edge type are stored consecutively, and their starting position in the array is maintained respectively in the array of starting indexes. In this way, when searching for a particular type of edges connected to $m$, we are able to achieve an efficient random access by leveraging the index range information. For example, in Fig. 3 if we are looking for the edges of type $[(E \_	ext{in, outgoing}, active)]$, we simply look at the $(k - 1) \ast 4 + 2 + 1)$th element in the array of starting indexes. Given $k = 2$, the 7th element in the index array is 11 and the following (8th) one is 12, we know that there is only one outgoing and active edge connected to $m$ with label $L_2$, i.e., $(m, v_1, L_2)$.

Given that there are $4L$ types of edges, the length of the array of starting indexes is $4L$. The total storage size of the two arrays is $(4L + X)$. Therefore, when there are $X$ edges and $X \gg 4L$, the amount of data stored is reduced from $3X$ to $X$. When the number of edges is large, the storage amount is reduced to one third (ignoring the platform implementation difference).

The index-based data structure not only reduces storage overheads, but also improves computational efficiency: when matching edges according to grammar rules, we can directly find the index range of the corresponding label with time complexity $O(1)$, instead of traversing the entire edge set.

### 4.3 Optimizations

#### Load Balancing
Load balancing plays a crucial role in the performance of parallel computing. In our data-parallel algorithm, if the sizes of partitions vary significantly, it probably arrives at a status that the majority of threads finish the tasks quickly and become idle waiting for one thread, bringing about poor performance. In our computation model, for each vertex, all of its incoming and outgoing edges need to be visible to perform label matching and edge addition. For instance, when $y$ is processed, both $x \rightarrow y$ and $y \rightarrow z$ need to be accessed to add the edge from $x$ to $z$ (shown as Line 11 to 14 in Algorithm 1). There perhaps exist some hot vertices which are associated with a huge number of adjacent edges. As a consequence, partitions would be unbalanced.

In order to address load imbalance, we adopt a finer-grained partitioning scheme. To be clear, we define the number of Matchable Edge-pairs (MEP) as Formula (11), where $m$ is the intermediate vertex representing an edge-pair, and $E_m(direction, status, label)$ is the number of edges connected to $m$ of a given type.

$$MEP(m) = \sum_{C::A, B \in Production} \left( E_m(in, active, A) \cdot E_m(out, all, B) + E_m(in, silent, A) \cdot E_m(out, active, B) \right).$$
The MEPs are calculated immediately after the join process of each iteration. If the MEP of certain vertex is beyond some threshold, we split it into multiple small partitions. In practice, the threshold can be set to an empirical value or the average MEP of all the vertices. Since we conduct the edge generation and filtering separately in our join-process-filter model, splitting edge pairs of one vertex into multiple partitions does not affect the correctness of computation. The computation time cost on each node shown in Figs. 12 and 13 empirically validated the effectiveness of our load balancing strategy.

Pre-Shuffle. Pre-shuffle is intended to reduce the network traffic when a join is performed on two datasets of key-value pairs. The normal join shuffles all partitions from the two datasets over the network. The pre-shuffle avoids this network communication by partitioning two datasets with the same partitioner in advance and joining corresponding partitions directly in the same node without the network traffic. In BigSpa, the join of old edges and new edges triggers shuffle at each iteration. We maintain a global partitioner during the execution and assure that the old edges are always partitioned by the global partitioner. When new edges are generated, we partition it with the global partitioner and join it with old edges via pre-shuffle. The united dataset becoming the old edges in the next iteration inherits the global partitioner as its partitioner automatically. In this way, the network communication is only involved during the repartitioning of the new edges at each iteration. BigSpa avoids re-shuffling old edges which become larger and larger, greatly reducing network traffic.

Computation Closure. Computation closure is a design optimization to accelerate the convergence of edge addition for specific analysis workloads. We check the analysis grammar and find that certain edges labelled by terminal symbols are static during the iterations. We broadcast these edges to all the worker nodes so that these edges can be accessed locally. At each iteration, each node computes a local closure to generate as much more edges as possible with the help of the broadcasted static edges. Take the dataflow analysis as an example, there exists one production rule \( Ne \rightarrow ew \) in the grammar. The edges labelled by terminal \( e \) are static. In other words, no new \( e \) edges will be added during computation. Given a path in the input graph \( x \rightarrow y \rightarrow z \rightarrow w \), we should get two new edges \( x \rightarrow z \) and \( x \rightarrow w \). During the mini-batch update process, although the updated parts to achieve high efficiency in time and memory usage. To handle this scenario, we further design an online incremental static program analysis system.

5 Process of Incremental Update

5.1 Process of Incremental Update

Synchronously, edges in the program analysis graph represent the reachability relationships between vertices, and there are two types of them: directly reachable (those formed by directly adding edges to the graph) and indirectly reachable (those generated by label matching based on the CFL syntax). When adding or deleting edges, the indirect reachability relationships generated by these edges should be changed accordingly. Therefore, an update on the graph requires iterative processing until the topology of the graph reaches a stable state.

In order to improve the performance, we use the minibatch update strategy to perform incremental static analysis, that is, multiple edges are added to the graph at the same time, and then the updated transitive closure is computed. During the mini-batch update process, although the order of adding and deleting edges may change, the graph topology of the final update graph is determined. Obviously, the same topology will lead to the same reachability relationship. This ensures the correctness of our mini-batch update method.

5.2 Method for Online Incremental Analysis

5.2.1 Triangle Count

As shown in Fig. 4, the difficulty of deleting edges is that an indirect edge could be generated on multiple paths. Without extra information, we cannot decide whether to delete that indirect edge or not. Simply storing all the generating paths will greatly increase the storage and calculation overheads (as discussed in Filter part in Section 4.1). To deal with the edge deletion correctly and efficiently, we propose a method based on the triangle count (short as \( \text{tri}_\text{count} \)). In this article, we define the triangle count of an edge \( e \) as the

5 Online Incremental Static Program Analysis System

In the previous section, we introduced how to process large scale static program analysis over the entire code dataset in the batch processing way. In fact, in the real world, software codebases can be updated day by day, especially for those popular open source software with large scale developer communities. In this scenario, when a program to be analyzed is changed or updated, re-analyzing the entire graph can be time-consuming and cumbersome. Instead, the system can perform local analysis on the updated parts to achieve high efficiency in time and memory usage. To handle this scenario, we further design an online incremental static program analysis system.
number of triangles with $e$ as one of their sides. Essentially, triangle count ($\text{tri}_\text{count}(x \rightarrow y)$) is the number of 1- or 2-hop paths from $x$ to $y$ in the reachability graph. Whenever an edge is deleted, other relevant edges remain if and only if their respective triangle counts are not decreased to be 0. The calculation of the triangle count is shown in Formula (12) (transitive closure is short as $\text{TC}$)

$$\text{tri}_\text{count}(v_1 \rightarrow v_3) = |\{v_2 \mid (v_1, v_2, A) \in \text{TC} \&\& (v_2, v_3, B) \in \text{TC} \&\& C := AB\}|.$$  

(12)

Fig. 5 describes the edge addition and deletion process based on the triangle count method, and the operations are quite similar. When a new indirect edge is newly generated, its $\text{tri}_\text{count}$ is initialized to 1. During the iterative calculation, each time an addition/deletion of an indirect edge is triggered, the $\text{tri}_\text{count}$ of this edge is increased/decreased by one. For example, if an edge $x \rightarrow A \rightarrow y$ is added, the system matches edge pairs based on labels and triggers the addition of an indirect edge $x \rightarrow C \rightarrow z$. When the edge $x \rightarrow A \rightarrow y$ is removed, the same label matching process is performed and we can find the same indirect edge $x \rightarrow C \rightarrow z$, which was generated by $x \rightarrow A \rightarrow y$. As one of its generating path is broken by the deletion, the triangle count of $x \rightarrow z$ should therefore be decreased by 1. An indirect edge is removed from the graph only when its $\text{tri}_\text{count}$ is reduced to zero. In this way, based on the edge-pair computation model, BigSpa can simplify the maintenance of provenance information via triangle counting. A theoretical proof for the correctness of using triangle counts to determine reachability can be found in the appendix of our technical report [21].

### 5.2.2 Online Incremental Computation Algorithm

Algorithm 2 shows the procedure of online incremental analysis. Compared with Algorithm 1, the main change is that a new variable $W_{\text{count}}$ is defined, which stores the new edges and the number of times they are added or deleted. It is used to update the triangle counts and generate the edge set $W$ to be calculated. During each iteration, $W_{\text{count}}$ is updated to ensure that it only covers the additions and deletions generated within the current round of computation, so that no triangles are missed or double-counted.

**Algorithm 2. Online Incremental Computation Algorithm**

Data: A batch of updates on graph $G$, Production set $P$
Result: TC of $G$

1. $W_{\text{count}} \leftarrow \text{count batch}$
2. $W \leftarrow W_{\text{count}} \text{ join } \text{TC}$
3. repeat
4. $P \leftarrow \text{TC join } W$
5. $W \leftarrow W_{\text{count}} \text{ diff } \text{TC}$
6. $W_{\text{count}} \leftarrow \text{map } (P)$
7. $W \leftarrow W_{\text{count}} \text{ diff } \text{TC}$
8. until $W = \emptyset$
9. $W_{\text{count}} \leftarrow W_{\text{count}} \text{ join } \text{TC}$

The computation model is thus modified as Fig. 6. In the online processing system, in order to get a lower response time for queries and updates, the distributed databases are utilized not only for filtering, but also for storing and updating the graph.

### 5.3 Data Structure

To record the triangle counts of each edge and reach the requirement of real-time response, we then improve the data structure introduced in Section 4.2. As shown in Fig. 7, according to their labels (assuming there are $L$ types of labels) and directions (inside and out), the adjacent edges of
each intermediate node are divided into $2L$ subsets. Note that the edge calculation state is not considered here, as all edges stored in the online system are silent edges (i.e., the reachable relationships that have already been calculated).

In general, iterative calculations caused by incremental updates are local, which means there is no need for the whole graph or all adjacent edges of a node to be involved in the calculation. With this fine-grained data structure, the system can access data on demand when updating the graph. Besides, there are two more advantages to storing the adjacent edges separately: first, it can avoid a single record in the database being too large; second, the system can directly locate the edges needed for calculation, reducing the cost for intermediate results.

5.4 Adaptive Computation Mode Switching Mechanism

The online incremental analysis system is designed to deal with small batches of update data, which, in most cases, trigger a small amount of computation. However, in some scenarios, small batch updates can also trigger large-scale calculations (as is the case where a large number of indirect edges are generated, mentioned in Section 4.3). Distributed platforms are suitable for the latter case, but they can be greatly slowed down by transmission between computing nodes in the former case. Considering that, we design an adaptive mode switching mechanism. Based on the prediction of computation scale, the system can automatically switch between single-machine multi-threaded computing mode and distributed computing mode.

In online incremental processing, after the join operations, a calculation scale prediction process is performed. Specifically, we calculate the sum of MEP (defined in Section 4.3) of all the intermediate nodes, which is the number of new edges to be generated and can indicate the scale of calculation. Hence, if the sum of MEP is below a threshold, the map process will be performed in the single-machine multi-threaded mode, otherwise the data to be processed is divided into different partitions and the distributed computing mode is activated. During each iteration, the system performs the prediction and switches between the two modes flexibly. In this way, we can both avoid computational failure due to insufficient single-machine resources and reduce the resource waste and timeout due to excessive overheads in distributed platforms.

6 SYSTEM DESIGN AND USAGE

6.1 System Architecture

The overall system architecture of BigSpa is shown in Fig. 8, which consists of four layers:

- Analysis Layer: The top layer of the program static analysis system includes the offline batch and online incremental analysis system described in Sections 4 and 5.
- API Layer: There are five types of operators to complete specific reading and calculation tasks for application usage.
- System Optimization Layer: This layer implements the various optimization measures, such as pre-shuffle, load balancing, computation closure and computation mode switching mechanism.
- Underlying Distributed Platform Layer: This layer includes a distributed data processing engine, a distributed file system, and a distributed database. In implementation, we choose Spark as the data processing platform, Hadoop HDFS as the distributed file system, and Redis as the distributed database.

6.2 BigSpa Usage

Fig. 9 demonstrates how to use BigSpa to carry out various analyses. Analogous to the declarative program analysis [15], [22], [23], we separated the computation back-end from the client analysis implementations. As stated earlier, BigSpa as a general analysis computation engine supports multiple analyses [10]. In order for users to implement a particular sophisticated interprocedural analysis, a front-end is required. As shown in Fig. 9, we implemented two front-ends, one for dataflow and the other for pointer analysis (both field-insensitive and field-sensitive) in our experiments. Users can develop their own front-ends for specific client analyses and then leave the computation to BigSpa. In the following, we discuss the two tasks of which each front-end is comprised, namely generating the program graph and specifying the analysis grammar.

Generating Graphs. In order for BigSpa to perform an interprocedural analysis, users are first required to generate a specialized program graph tailored for the analysis. In our graph generation, we first perform intra-graph generation and call graph generation. We then perform the function inlining to produce the final input graph.

For intra-graph generation, we implemented each separate intra-graph generator for each particular client analysis based on different infrastructures. Specifically, for programs written in C/C++, each intra-graph generator is implemented as an analysis pass based on LLVM [24] which takes the LLVM IR as input and produces an intra-graph for each

![Fig. 8. System structure of BigSpa.](image-url)
function according to the particular client analysis. For Java subjects, our frontend is implemented based on the compiler infrastructure Soot [25].

During call graph generation, we launched an existing lightweight pointer/alias analysis to help track the precise call relations. Specifically, for C/C++ programs, we used an inclusion-based context-insensitive pointer analysis with support for function pointers (available in LLVM). To analyze Java, we adopted the context-insensitive Spark points-to analysis [26] in Soot to efficiently construct the (sound but relatively imprecise) call graph. **Specifying Grammar:** Given the graph generated, the front-end needs to specify a particular grammar guiding the edge addition for the analysis. For example, pointer/alias analysis could take advantage of the grammar similar to the rules 8 - 10 in Section 2.2.

For the purpose of performance, our model is devised to access at most two adjacent edges for label matching. Such simple computation logic makes our design easier and leads to a dynamic programming style algorithm which imposes lower complexity. To this end, each production rule in the grammar needs to be normalized so that it has no more than two terms on its right-hand-side (RHS). The transformation can be done by introducing new non-terminal symbols in linear time, and causes a linear increase in the size of the grammar [19]. The normalized grammars for field-insensitive pointer/alias analysis and dataflow analysis can be found in the appendix of our technical report [21].

### 7 Evaluation

This section presents an empirical evaluation of BigSpa. We focus on three research questions:

- **Q1:** What is the performance of BigSpa?
- **Q2:** How does BigSpa compare to the state-of-the-art single-machine analysis system?
- **Q3:** How does BigSpa compare to other state-of-the-art distributed systems for analysis workloads?

#### Hardware and Software Environment

Unless otherwise specified, experiments presented in this article were conducted in a cluster with 17 nodes (1 master node + 16 worker nodes). Each node was equipped with two Intel Xeon E5620 CPUs (2.4 GHz/12 MB Cache, 12 physical cores and 24 logical cores in total), 16 GB available memory and 3 TB HDD RAID0 disk storage. All nodes were connected via 1 Gbps Ethernet. Our system was implemented with Apache Spark 2.2.0 and used Apache Hadoop 2.7.2 to provide HDFS and Yarn. Redis 3.2.0 was employed as the distributed database. We compiled and ran all the programs with JDK 1.8.0 and Scala 2.11.8 on the RHEL 7 64-bit OS with Linux kernel v3.10.0.

#### 7.1 Evaluation of Offline Batch System

We first evaluate the performance of the offline batch system.

##### 7.1.1 Performance Analysis

**Performance and Scalability.** Table 3 reports the performance data of BigSpa. The columns \#V(M), \#EB(M), \#EA(M), \#Ites, PT(mins), CT(mins), and TT(mins) correspond to the number...
of vertices, number of edges before computation, number of edges after computation, the number of iterations, the preprocessing time, the computation time and the total time, respectively. All the numbers of vertices and edges reported are in millions and the time costs are in minutes. The preprocessing time is mainly taken by loading graphs and grammars into the distributed file system (HDFS) and the cache system (Redis), while the computation time shows the time cost for edge addition.

BigSpa managed to finish most of the analyses within 20 minutes in our experiments. Even for the biggest graph with more than a billion edges, BigSpa only took around half an hour to complete the computation. In the field-sensitive pointer/alias analysis, as each of the grammars contains thousands of symbols, the intermediate results take up extremely large amount of memory even with our compressed data structure. We increased the available memory of each node to 30 GB and completed the analysis of HDFS and Hadoop with 25 nodes and 32 nodes respectively. It took BigSpa tens of minutes to finish the analyses. By contrast, the cutting-edge single machine systems like Graspan [7] were unable to process neither of the above two workloads, because they greatly exceed the processing capability of one single machine.

Table 4 shows the proportion of time spent on updating database, shuffling data between machines and Java’s garbage collection on the representative subjects. For pointer/alias analysis, the computation time dominated the overall time cost. The communication was mainly spent on updating Redis, as the new edges generated in each iteration were written into the database. As for dataflow analysis, the preprocessing time became comparable with the computation time. The preprocessing time was mainly spent on broadcasting edges which is required by the closure optimization strategy discussed in Section 4.3. This also greatly reduced the number of edges that need to be written into Redis. GC took much more time when analyzing large or heavily-skewed graphs, since the computation process generates more intermediate objects.

Fig. 10 plots the analysis time of BigSpa running on a cluster with varying number of nodes. We present here the results of two analyses which can be finished in reasonable time even on a single node. The detailed data for all subject analyses can be checked shortly in Table 6. We can read from the trend that BigSpa scales well (near linear) with the number of nodes available.

CPU and Memory Usage. Fig. 11 illustrates the CPU and memory usage of a node when we ran BigSpa in a cluster for PostgreSQL. The memory occupied by each machine was composed of three parts: (1) Spark RDD memory, which stores the edge set and the intermediate results; (2) Redis memory, which stores the transitive closure for the filter process; (3) other memory reserved by Hadoop and Yarn, etc.

For pointer/alias analysis, the CPU utilization is relatively low and the memory consumption increases as BigSpa iterates. This is consistent with Fig. 15 in the sense that the map stage contributes little to the overall cost. It further confirms that the pointer/alias analysis is network-communication

---

**TABLE 3**

The Performance of BigSpa in Batch Processing

<table>
<thead>
<tr>
<th>Subj.</th>
<th>#V(M)</th>
<th>#EB(M)</th>
<th>#EA(M)</th>
<th>#Ite</th>
<th>PT(min)</th>
<th>CT(min)</th>
<th>TT(min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Linux</td>
<td>52.88</td>
<td>249.50</td>
<td>1.13B</td>
<td>73</td>
<td>2.51</td>
<td>27.3</td>
<td>29.81</td>
</tr>
<tr>
<td>PSQL</td>
<td>5.20</td>
<td>14.97</td>
<td>862.18</td>
<td>70</td>
<td>0.75</td>
<td>11.6</td>
<td>12.35</td>
</tr>
<tr>
<td>httpd</td>
<td>1.72</td>
<td>1.19</td>
<td>904.34</td>
<td>65</td>
<td>0.63</td>
<td>12.05</td>
<td>12.68</td>
</tr>
<tr>
<td>HDFS</td>
<td>5.34</td>
<td>19.10</td>
<td>1.80B</td>
<td>7</td>
<td>1.15</td>
<td>25.46</td>
<td>26.61</td>
</tr>
<tr>
<td>Hadoop</td>
<td>21.88</td>
<td>77.17</td>
<td>99.15</td>
<td>9</td>
<td>1.65</td>
<td>3.25</td>
<td>4.9</td>
</tr>
</tbody>
</table>

**TABLE 4**

Proportion of Time Spent on Communications and Garbage Collection

<table>
<thead>
<tr>
<th>Subj.</th>
<th>Redis ratio (%)</th>
<th>Shuffle ratio (%)</th>
<th>GC-ratio (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Pointer/Alias Analysis</td>
<td>Flow-scan Analysis</td>
<td></td>
</tr>
<tr>
<td>Linux</td>
<td>18.63</td>
<td>4.15</td>
<td>9.41</td>
</tr>
<tr>
<td>PSQL</td>
<td>35.12</td>
<td>2.71</td>
<td>5.07</td>
</tr>
<tr>
<td>httpd</td>
<td>35.17</td>
<td>1.99</td>
<td>7.62</td>
</tr>
<tr>
<td>HDFS</td>
<td>31.43</td>
<td>4.80</td>
<td>37.94</td>
</tr>
<tr>
<td>Hadoop</td>
<td>2.85</td>
<td>7.15</td>
<td>12.93</td>
</tr>
</tbody>
</table>

(a) Pointer/Alias Analysis

(b) Dataflow Analysis

Fig. 10. Scalability analysis.

Fig. 11. CPU utilization and memory consumption for PostgreSQL.
intensive in BigSpa. For dataflow analysis, the memory consumption is steady since the number of new edges generated in the dataflow analysis is relatively small. Besides, in dataflow analysis Redis occupies only a small amount of memory. This is because only a few new edges are generated in the dataflow analysis and the Redis overhead allocated to each machine is minimal. The CPU utilization varies a lot, which has a strong correlation with the number of new edges added at each iteration. We only show the data for PostgreSQL here. Other subjects have the similar trends.

**Load Balancing.** We measured the execution time of map phase (i.e., computation) at each node in the cluster. The MEP threshold for node splitting was set to 1E6 in our experiments. Fig. 12 presents the data for pointer/alias analysis on Linux kernel. The results for other analyses show similar trend. X axis denotes the computing node ID in the cluster, while Y axis shows the execution time in seconds. The dashed line corresponds to the average time (56.21 seconds) across the whole cluster. We computed the standard deviation which is as low as 1.4 percent (0.81 / 56.21). Fig. 13 provides box-plots showing the distribution of the mean CPU usage of all cores of a node. It can be seen that during each iteration, the 24 CPU cores on the same node have no obvious load difference and the maximum CPU usage difference is about 10 percent. These results empirically validate that our load balancing strategy works well on both node level and thread level.

**Optimizations.** In this part, we evaluate the efficiency of the optimizations described in Section 4.3. Different optimizations work for different scenarios: the pre-shuffle and load balancing optimizations are tailored for pointer/alias analysis, while the closure optimization is for dataflow analysis.

Fig. 14 illustrates the effects of pre-shuffle and load balancing on the total analysis time. The optimizations can significantly lower the end-to-end time cost in general because the pre-shuffle strategy greatly reduces data transfer overhead, especially when a large number of new edges are generated. Apart from this, the node splitting operation can prevent too many computing tasks from being assigned to a single node. The improvement brought by load balancing is particularly obvious on some extremely skewed graphs like HDFS. Without this, it leads to much longer processing time or even failures. The only exception is pointer/alias analysis on Hadoop, where the optimizations introduce an additional time cost (about 30 seconds). Since the scale of graphs for this benchmark is small, the optimizations can only make a relatively small improvement but still introduce a little extra cost.

Table 5 lists the effect of closure optimization on dataflow analysis. Although the dataflow analysis generates a relatively small number of edges compared with pointer/alias analysis, it requires much more iterations and thus takes much longer computation time. The optimization of computation closure can greatly reduce the number of iterations needed, especially on the Linux dataset, where the number of iterations is reduced from 6,636 to 112 (a reduction of 98.31 percent).

**Time Cost Distribution.** We monitored the time cost distribution and number of edges added at each iteration of BigSpa for each analysis shown as Fig. 15. Within each iteration, the time cost comes from five stages, namely join, map, filter, distinct, and union corresponding to each stage in our computation model (Fig. 2).

For pointer/alias analysis (Fig. 15a), the time cost at each iteration ascends to a peak and then descends. After the peak, there is a long tail to converge. In addition, as shown in the plots, there exists a high correlation between the time cost and the number of edges added at each iteration. The trend of the join time verifies the efficiency of the pre-shuffle optimization discussed in Section 4.3. As BigSpa iterates, the total number of edges increases and the time spent on join should also increase. The pre-shuffle optimization makes BigSpa only shuffle newly added edges at each iteration, thus stabilizing the time cost at the join stage. Note that although the join time cost does not increase as the computation proceeds, it still dominates the total time cost in pointer/alias analysis.

The dataflow analysis (Fig. 15b) has the similar trend to the pointer/alias analysis but with the peaks appearing at the beginning. As the dataflow analysis exploits the computation closure optimization, a large number of new edges are quickly added in the first few iterations. Note that there

### Table 5

<table>
<thead>
<tr>
<th>Analysis</th>
<th>Subject</th>
<th>#It Before</th>
<th>#It After</th>
<th>Runtime Before(mins)</th>
<th>Runtime After(mins)</th>
</tr>
</thead>
<tbody>
<tr>
<td>dataflow</td>
<td>Linux</td>
<td>6,636</td>
<td>112</td>
<td>933.31</td>
<td>15.31</td>
</tr>
<tr>
<td></td>
<td>PSQL</td>
<td>721</td>
<td>59</td>
<td>58.93</td>
<td>9.98</td>
</tr>
<tr>
<td></td>
<td>httpd</td>
<td>181</td>
<td>23</td>
<td>8.55</td>
<td>3.42</td>
</tr>
</tbody>
</table>
is a peak of computation time in iteration 50. This is because a lot of edges are generated in this iteration but most of them are duplicated and as a result, the total number of edges does not change significantly.

7.1.2 Comparison With Graspan [7]

We compared BigSpa against a single-machine disk-based interprocedural analysis system, called Graspan [7]. BigSpa was experimented with 1, 2, 4, 8, 16 nodes successively. As a comparison, we ran C++ version Graspan on one single node equipped with the same CPU as described before. In our experiments, Graspan was configured to use 24 threads, 4 initial partitions and 16 GB memory. Since it is an out-of-core system, we conducted Graspan experiments on both HDD RAID0 disks and SSD RAID0 disks. Table 6 shows the detailed running time in minutes and COSN metric.

Because Graspan’s IO time accounts for only a small proportion of the total cost, and most of its disk reads and writes are sequential, using SSD can only bring little acceleration. With 16 nodes, BigSpa runs one or more orders of magnitude faster than Graspan in most cases. However, for certain analyses especially with large graphs, the performance of BigSpa running on a few (1 or 2) nodes is disappointing. The underlying reason is that BigSpa based on Apache Spark utilizes memory as higher priority for efficient computation. Unfortunately, once the memory consumption exceeds certain limit, heavy Garbage Collection (GC), frequent disk I/O and data serialization/de-serialization on massive data have to be involved resulting in unacceptable performance. The experiment running on a single node reported that, the maximum I/O could reach 200 MB per second. The GC time even occupies more than 70 percent of the overall execution time at late iterations. That explains why a super-linear speedup appeared when analyzing large-scale pointer analysis graphs. As more computer nodes become available, memory pressure is relieved. GC time hence drops dramatically, achieving superlinear speedup.

Note that BigSpa performs significantly well on dataflow analysis compared against Graspan. BigSpa running on one node still outperforms Graspan a lot. The performance advantage mainly gives the credit to the following two design optimizations. First, BigSpa adopts computation closure optimization to reduce the number of iterations needed. Second, duplication removal and edge union are implemented in a more efficient way in BigSpa than Graspan. Graspan applies the merge operation to filter out duplicated edges and to unite edges. Even though just a few edges are newly generated, Graspan still needs to reconsider a great number of old edges. BigSpa exploits the Redis database to conduct the filter operation and adopts pre-shuffle technique to join the new edges with the old ones efficiently. The costs of both operations are just related to the number of newly added edges.

In the experiments for field-sensitive pointer/alias analysis, BigSpa needed at most 32 nodes to finish processing the HDFS and Hadoop data sets in dozens of minutes (the detailed performance data can be found in Table 3). By contrast, Graspan failed to run on both HDFS and Hadoop data sets because the scale of both data sets greatly exceeded the processing capability of one single machine. Although the process of parallelization makes BigSpa introduce additional overhead for a single machine, it provides a more scalable solution for inter-procedural analysis. Users can utilize BigSpa to perform large-scale sophisticated analysis tasks simply by scaling the system to some extent. Moreover, with additional resources and the optimizations enabled, BigSpa managed to process the heavily skewed graphs and generated millions of edges within a few iterations, demonstrating the advantages of distributed computing.

7.1.3 Comparisons With Other Distributed Systems

We also conducted the comparisons with other distributed systems on static analysis workloads, namely BigDatalog [22], and distributed call graph analysis [30].

Comparison With BigDatalog. BigDatalog [22] is a distributed datalog-based [15] computation system which supports efficient static analysis. We compared the performance of BigDatalog and BigSpa for both pointer/alias and dataflow analyses on Linux kernel. All experiments done for BigSpa and BigDatalog were conducted in the same cluster with the same Spark installation.
As for dataflow analysis (Fig. 16a), BigSpa outperforms BigDatalog significantly. The number of iterations BigSpa needs is much smaller than that of BigDatalog. A large amount of time is taken by BigDatalog for convergence. With regard to the pointer/alias analysis, BigSpa outperforms BigDatalog with all different number of nodes. Note that Fig. 16b actually shows the performance results for a simplified pointer/alias analysis. Due to implementation issues, BigDatalog cannot support fully-precise pointer analysis. Runtime errors occurred when compiling rules of multiple consecutive joins, e.g., $M := \emptyset \cup V \cup d$. Reluctantly, we chose a simplified pointer analysis by removing certain rules for the sake of fair comparisons.

**Comparison With Distributed Call Graph Analysis.** We also compared with another distributed actor model-based analysis tool [30]. We exert ourselves to deploy BigSpa on Microsoft Azure cluster, and implemented a front-end for the inclusion-based Variable Type Analysis [31] to generate the propagation graph, which was fed into BigSpa to compute the CFL-reachability results. Hence, the time cost spent by BigSpa comes from two parts, frontend for propagation graph generation and backend for reachability computation. We conducted all the comparison experiments on a Microsoft Azure A3 cluster with 1 master and 4 workers, each equipped with 4 CPU cores and 7 GB memory. Table 7 shows performance results for the real programs chosen by [30]. Column “frontend” and “backend” indicate the time cost by generating propagation graph and computing reachability.

By comparing the total time spent, BigSpa outperforms Distributed Call Graph Analysis [30] especially on large graphs (e.g., more than 4 times faster on ILSpy).

### 7.2 Evaluation of Online Incremental System

In this subsection, we evaluate the performance of the online incremental static analysis system.

#### 7.2.1 Performance Analysis

We chose the largest Linux and the smallest httpd as the subjects for this experiment. We produced the program graphs for pointer/alias analysis and data flow analysis, and generated updating batches in two ways: (1) randomly divide them into multiple batches of edges, each with 1,000 edges; (2) add and delete nodes based on commits from the realistic version control repositories. We refer to these two experiments as uniform update and commit-based update in the following. For the commit-based update, we collected tens of thousands of commit records from the GitHub repositories of Linux and httpd respectively and remove the commits that do not involve code changes. Some of these commits only modify a few lines of code, while some others can cause thousands of edges to be modified. The details about the commits can be found in Table 8.

The mini-batches corresponding to both scenarios were then input into the incremental system. To evaluate the effectiveness of our system under real-time scenarios, we monitored the time taken for each update, and the tasks with processing time exceeding 60 seconds (can be configured by users) would be interrupted in our experiments. As can be seen from Tables 9 and 10, our system has significantly different performance on different tasks. The Response Time column lists the average response time and several upper quantiles. For example, “0.52” in the first row of Table 9 means that 90 percent of the updates were completed in 0.52 second.

The system shows good performance for pointer/alias analysis of Linux dataset, and for dataflow analysis of Linux and httpd dataset. In these three tasks, thousands of edges are processed per second on average and all the tasks can be finished within 15 seconds. However, in other tasks, the throughput is significantly reduced and timeout interruptions occur.

By monitoring the execution process of the tasks and analyzing the structure of the input graphs, we found that

1. The tasks with calculation time at millisecond level generally have fewer edges to update, and are calculated in the single-machine multi-thread mode.
2. In those tasks completed in seconds, the number of updated edges is larger, but still less than 100,000. In this case the system can still finish the calculation by switching to a distributed computing mode.
3. Overtime tasks often have super-large-scale updates and “hot vertices”.
4. In general, the system has higher throughput and lower processing latency in the commit-based experiment. This is because in large projects, most of the commits only trigger small-scale code updates.

In the above experiments, the system shows poor performance when there are a lot of “hot vertices” and tightly connected vertex clusters in the input mini-batches, as is the case of pointer/alias analysis on httpd. Updating these mini-batches is similar to updating a full graph ($O(|N|^2)$),

### TABLE 8

<table>
<thead>
<tr>
<th>Analysis</th>
<th>Subject</th>
<th># Commits</th>
<th>Proportion of commits with different number of modified edges (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>0.100</td>
</tr>
<tr>
<td>Pointer/Alias</td>
<td>Linux</td>
<td>35,235</td>
<td>57.09</td>
</tr>
<tr>
<td>Dataflow</td>
<td>Linux</td>
<td>35,235</td>
<td>57.09</td>
</tr>
</tbody>
</table>

* The SHA ids of these commits can be found at https://github.com/PasaLab/BigSpa/tree/master/data/online_commits
In contrast, on the Linux dataset, which is the largest one but has a more uniform distribution of edges, the incremental system performs excellently in both kinds of analyses. Therefore, it can be concluded that the online system is suitable for analyzing program graphs with uniform distribution of the transmission edges, and the performance would be affected when performing incremental analysis of program graphs with skew-distributed transmission edges.

From the application scenario, it is not realistic to lower the time cost of precise analysis of large-scale codebases to second level. Because of that, a timeout interruption mechanism is employed in our implementation to return only part of analysis results in time. If the interruption happens, the users will be informed by the output message which can be regarded as a signal. In this situation, they can use the offline batch analysis system for supplementary calculations to get the complete and accurate analysis results.

7.2.2 Comparison With Serial Incremental Analysis

For comparison, in this subsection, the same batches were input into a serial incremental analysis system in a serial manner.

The results of the comparison experiment are illustrated in Fig. 17. The horizontal axis indicates the software and analysis type, and the vertical axis indicates the average processing time of one batch in seconds. As we can see, the average response time of our mini-batch incremental system is obviously lower (less than one-twentieth) than that of the serial system. That is because there are many iterations in the process of incremental analysis, and serial iteration over multiple edges would result in a huge waste of computing resources. Besides, in the serial system, the amount of data to be processed per round is relatively small, therefore the memory can hardly be fully utilized. The above results show that the method of distributed mini-batch processing can effectively handle the incremental updates of program static analysis. Moreover, in practice, it is more common to update the codebases in mini-batches, which means the mini-batch update method is more suitable for real-world application scenarios.

8 RELATED WORK

Static Bug Detection. Static analysis is widely used to detect various software defects and security vulnerabilities [1], [2], [3]. Engler et al. [3] take advantage of the simple pattern-based analysis to discover a variety of bugs in Linux kernel. In addition, a series of commercial tools, e.g., Coverity [32], CodeSonar [33], KlocWork [34], are also developed and widely used in industries. Most of them are simply based on patterns or intra-procedural analysis which has low computation complexity, achieving good scalability. However, pattern-based or intra-procedural analysis suffers from low accuracy due to the lack of considering rich/complete semantics information. Some empirical studies show that the false positive rates reach 30-100 percent [4], leading to pretty poor applicability [5]. Differently, this work focuses on the sophisticated inter-procedural analysis. By developing a scalable inter-procedural analysis engine, we are able to perform precise sophisticated analysis on large-scale modern software.

CFL Reachability-Based Static Analysis. CFL-reachability is originally proposed by Yannakakis [15] for Datalog query evaluation. Reps [10] later adopt CFL-reachability to formulate a series of sophisticated static analysis. A variety of static analyses [8], [9], [16], [35] can be viewed as the instances of CFL-reachability problem, e.g., program slicing, inter-procedural dataflow analysis, pointer and alias analysis, shape analysis, specification inference, and information flow analysis. The worst case time complexity for solving CFL-reachability problems is $O(n^3)$, which is called cubic bottleneck [36]. As a result, it is really tough to perform highly precise analysis on large-scale software. Researchers have proposed various optimizations to enhance the analysis efficiency, including compositional analysis [37], sparse representation-based analysis [6], and demand-driven analysis [9] etc. The above work is mainly targeted at the optimized sequential and memory-based algorithm whose scalability is greatly limited.

Incremental Program Analysis. For efficient analysis of frequently changing codebases, researchers have also proposed many incremental analysis approaches. Reviser [38] checks the modified version of a code to find the changes of its control flow graph, and then propagates and updates the analysis results. Souter [39] and Ismail [40] employ specialized algorithms for incremental call graph reconstructions in IDE. IncA [41] is a domain-specific language for the definition of efficient incremental program analyses, which functions in a declarative way over AST representations of programs. Lu et al. [42] combine points-to analysis with graph reachability. They provide a trace-based incremental mechanism that precisely identifies and recomputes affected paths after a program changes. The existing algorithms usually suffer from the lack of good scalability and generality, while our incremental analysis system can incrementalize analysis of very large codebases with distributed computing.

### Table 9: Performance of Online Incremental System in the Uniform Update Experiments

<table>
<thead>
<tr>
<th>Analysis Subject</th>
<th>Function Time (s)</th>
<th>Task Interruption Rate (%)</th>
<th>Throughput (Speedup)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Average</td>
<td>95%</td>
<td>99%</td>
</tr>
<tr>
<td>Points/Aisle</td>
<td>Linux</td>
<td>0.31</td>
<td>0.34</td>
</tr>
<tr>
<td></td>
<td>BigTree</td>
<td>0.45</td>
<td>0.47</td>
</tr>
<tr>
<td>Dataflow</td>
<td>Linux</td>
<td>0.26</td>
<td>0.29</td>
</tr>
<tr>
<td></td>
<td>BigTree</td>
<td>0.30</td>
<td>0.33</td>
</tr>
</tbody>
</table>

### Table 10: Performance of Online Incremental System in the Commit-Based Update Experiments

<table>
<thead>
<tr>
<th>Analysis Subject</th>
<th>Function Time (s)</th>
<th>Task Interruption Rate (%)</th>
<th>Throughput (Speedup)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Average</td>
<td>95%</td>
<td>99%</td>
</tr>
<tr>
<td>Points/Aisle</td>
<td>Linux</td>
<td>0.58</td>
<td>0.63</td>
</tr>
<tr>
<td></td>
<td>BigTree</td>
<td>0.70</td>
<td>0.74</td>
</tr>
<tr>
<td>Dataflow</td>
<td>Linux</td>
<td>0.30</td>
<td>0.33</td>
</tr>
<tr>
<td></td>
<td>BigTree</td>
<td>0.33</td>
<td>0.36</td>
</tr>
</tbody>
</table>

Fig. 17. Comparison with a serial incremental analysis implementation.
Parallel and Distributed Static Analysis. In order to improve the efficiency of program analysis, researchers have developed parallel algorithms. For example, Mendez-Lojo et al. [43] propose a parallel points-to analysis based on constraint graph rewriting. Su et al. [44] develop a CFL-based parallel pointer analysis algorithm where they leverage data sharing and query scheduling to avoid redundant graph traversals. Graspan [7] is a single-machine disk-based inter-procedural analysis system using edge-pair centric computation model. It solves the scalability problem to some extent, but still suffers from the limited scalability due to the resource limitation of a single machine. [45] and [30] propose actor model-based parallel algorithms for dataflow analysis and call-graph analysis respectively. Alberghouthi et al. [46] parallelize the top-down interprocedural analysis based on MapReduce paradigm. Different from BigSpa, they focus on the demand-driven analysis and only implement a parallel version running on a multi-core machine. Google also implement the distributed static analyses to analyze their codebase [47], but only relatively simple (intra-procedural) analyses are considered due to performance limitations. In addition, Facebook propose their tool INFER [48] based on bi-abduction to check the memory safety properties of C code. Blaß et al. [49] present a parallel approach for fixpoint-based dataflow analysis on a GPU. Although the method performs efficiently, it requires extra extractors and can hardly scale out. IfA [50] uses Pointer Assignment Graphs (PAGs) to parallelize incremental Anderson’s pointer analysis.

Although the above studies solve the scalability problem to some extent, they still have several drawbacks. First, they usually focus on certain particular analysis and are not applicable to the general program analysis. Second, some actor model based algorithms are not scalable enough due to the big synchronization overheads. Finally, most existing parallel and distributed algorithms entirely rely on memory for computation. As stated in [51], the memory would be the major bottleneck for scaling analysis to large programs. We have also proposed the parallel offline batch interprocedural static program analytic algorithms in the conference version [52] of this article.

9 CONCLUSION AND FUTURE WORK

In this article, we propose an efficient, distributed interprocedural static analysis engine called BigSpa. We improve the idea of transforming the static analysis problem into a big data processing problem and devise a particular data-parallel algorithm. Based on that, we develop the scalable BigSpa system that can handle both offline batch and online incremental static analysis of programs. Specific optimizations for data structure and processing flow are also designed to ensure high overall efficiency. By utilizing BigSpa, we readily accomplished a series of precise static analyses for large-scale modern software.

In the future, we plan to eliminate redundancy in data storage. Also, we want to explore how to set proper thresholds in the system by using machine learning techniques.

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REFERENCES


Rong Gu received the PhD degree in computer science from Nanjing University, Nanjing, China, in December 2016. He is an associate researcher with the State Key Laboratory for Novel Software Technology, Nanjing University, China. His research interests include distributed storage, parallel computing, and distributed machine learning.

Zhiquiang Zuo received the PhD degree from the National University of Singapore, Singapore, in 2015. He is currently an assistant professor with the State Key Laboratory of Novel Software Technology, Nanjing University, China. His research interests span programming languages, software engineering, and systems.

Xi Jiang received the BS degree from the Nanjing University of Science and Technology, Nanjing, China, in 2016. He is currently working toward the MS degree at Nanjing University, Nanjing, China. His research interests include distributed storage system and parallel algorithms.

Han Yin received the BS degree from Nanjing University, Nanjing, China. He is currently working toward the MS degree at Nanjing University, Nanjing, China. His research interests include distributed storage system and parallel algorithms.
Zhaokang Wang received the BS degree from Nanjing University, Nanjing, China, in 2013. He is currently working toward the PhD degree at Nanjing University, Nanjing, China. His research interests include distributed graph algorithms and graph processing systems.

Xuandong Li received the MS and PhD degrees from Nanjing University, Nanjing, China, in 1991 and 1994, respectively. He is a full professor with the State Key Laboratory of Novel Software Technology, Nanjing University, China. His research interests include formal support for program design and analysis, software testing and verification.

Linzhang Wang received the PhD degree from Nanjing University, Nanjing, China, in 2005. He is currently a full professor with the State Key Laboratory of Novel Software Technology, Nanjing University, China. His research interests include software engineering and software security.

Yihua Huang received the PhD degrees in computer science from Nanjing University, Nanjing, China. He is a professor with Computer Science Department and State Key Laboratory for Novel Software Technology, Nanjing University, China. His main research interests include parallel and distributed computing and big data parallel processing.

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