

Toward Full Elasticity in Distributed Static Analysis: The Case of Callgraph Analysis

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ABSTRACT

In this paper we present the design and implementation of a distributed, whole-program static analysis framework that is designed to scale with the size of the input. Our approach is based on the actor programming model and is deployed in the cloud. Our reliance on a cloud cluster provides a degree of elasticity for CPU, memory, and storage resources. To demonstrate the potential of our technique, we show how a typical call graph analysis can be implemented in a distributed setting. The vision that motivates this work is that every large-scale software repository such as GitHub, BitBucket or Visual Studio Online will be able to perform static analysis on a large scale.

We experimentally validate our implementation of the distributed call graph analysis using a combination of both synthetic and real benchmarks. To show scalability, we demonstrate how the analysis presented in this paper is able to handle inputs that are almost 10 million lines of code (LOC) in size, without running out of memory. Our results show that the analysis scales well in terms of memory pressure independently of the input size, as we add more virtual machines (VMs). As the number of worker VMs increases, we observe that the analysis time generally improves as well. Lastly, we demonstrate that querying the results can be performed with a median latency of 15 ms.

CCS CONCEPTS

• **Theory of computation** → **Distributed algorithms**; • **Software and its engineering** → **Automated static analysis**; **Integrated and visual development environments**; • **Computer systems organization** → *Cloud computing*;

KEYWORDS

Development environments and tools, Parallel, distributed, and concurrent systems, Performance and scalability, Program analysis, Program comprehension and visualization

*Part of this work was done when the authors were visiting Microsoft.

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1 INTRODUCTION

In the last decade, we have seen a number of attempts to build increasingly more scalable whole program analysis tools. Advances in scalability have often come from improvements in underlying solvers such as SAT and Datalog solvers as well as sometimes improvements to the data representation in the analysis itself; we have seen much of this progress in the space of pointer analysis [3, 15, 16, 22, 25, 26, 42].

Limits of scalability: A typical whole-program analysis is designed to run on a single machine, primarily storing its data structures in memory. Despite the intentions of the analysis designer, this approach ultimately leads to scalability issues as the input program size increases, with even the most lightweight of analyses. Indeed, if the analysis is stateful, i.e. it needs to store data about the program as it progresses, typically, in memory, eventually this approach ceases to scale to very large inputs. Memory is frequently a bottleneck even if the processing time is tolerable, despite various memory compression techniques such as BDDs. We believe that the need to develop scalable program analyses is now greater than ever. This is because we see a shift toward developing large projects in centralized source repositories such as GitHub, which opens up opportunities for creating powerful and scalable *analysis backends* that go beyond what any developer's machine may be able to accomplish.

Distributed analysis: In this paper we explore an alternative approach to build distributed static analysis tools, designed to scale with the input size, with the goal of achieving full elasticity. In other words, no matter how big the input program is, given enough *computing resources*, i.e. machines to execute on, the analysis will complete in a reasonable time. Our analysis architecture assumes that the static analysis runs in the cloud, which gives us elasticity for CPU and memory resources, as well as storage. More specifically, in the context of large-scale code repositories, even code understanding and code browsing tasks are made challenging by the size of the code base. We have seen the emergence of scalable online code browsers such as Mozilla's LXR [28]. These tools often operate in batch mode, and thus have a hard time keeping up

with a rapidly changing code repository in real time, especially for repositories with many simultaneous contributors. We aim to show how a nimbler system can be designed, where analysis results are largely stored in memory, spread across multiple machines. This design results in more responsive queries to obtain analysis results.

1.1 Motivation: Static Analysis Backend

Imagine a large project hosted within a centralized source repository such as GitHub, BitBucket or Visual Studio Online¹. We see an emerging opportunity to perform server-side analysis in such a setting. Indeed, the backends of many such repositories consists of a large collection of machines, not all of which are fully utilized at any given time. During the downtime, some of the available cycles could be used to do static analysis of the code base. This can help developers with both program understanding tasks, such as code browsing as well as other static analysis applications, such as finding bugs.

The ever-changing code base: As is typical for large projects, multiple developers constantly update the code base, so it is imperative that the server-side analysis be both responsive to read-only user queries and propagate code updates fast. At the same time, within a large code base, many parts of the code, often entire directories remain unchanged for days or months at a time. Often, there is no reason to access these for analysis purposes. Therefore, to ensure that we do not run out of memory, it is important to have a system that is able to bring analysis nodes into memory on demand and persist them to disk (put them to sleep) when they are no longer needed.

1.2 Call Graph Computation

In this paper we advocate the use of the *actor model* as a building block of typical worklist-based analysis approaches. More specifically, we use this approach to implement a typical call graph construction algorithm. While the algorithm itself is quite well-known and is not a contribution of this paper, the way it is implemented in a distributed setting is. Also note that call graph information is used for interactive tasks such as autocomplete (or Intellisense), as shown in Figure 6. For tasks like this, both accuracy and responsiveness are important. Call graph construction is a fundamental step of most whole-program analysis techniques. However, most of the time, call graph analysis computation is a *batch* process: starting with one or more entry points such as `Main`, the call graph is iteratively updated until no more methods are discovered.

Interactive analysis: Our setting in this paper is a little different. Our goal is to answer *interactive* user queries quickly. Our queries are the kind that are most frequently posed in the context of code browsing and debugging, and are already supported on a syntactic level by many IDEs. Specifically, our analysis in this paper has been developed to provide semantic, analysis-backed answers for the following IDE-based tasks: (1) **Go to definition:** Given a symbol in the program, find its possible definitions²; (2) **Who calls me:** Given

¹Visual Studio Online: <https://www.visualstudio.com/team-services/>.

²Note that this process is challenging due to the presence of polymorphism, common in object-oriented languages. Given a call site, it is not always possible to determine which is the actual method implementation being invoked. This problem known as *call site devirtualization* is well-studied in the literature. Therefore, a static analysis can only approximate the target method definitions for a virtual method invocation.

a method definition, find all of its callers; (3) **Auto-complete:** Auto-completion, invoked when the developer presses a dot is one of the most common and well-studied tasks within an IDE [7, 19, 23, 29–31]. If the variable or expressions on the left-hand side of the dot is of a generic interface type, completion suggestions are not particularly useful or too general. It is therefore helpful to know which concrete type flow to a given abstract location.

We have architected our analysis backend to respond to REST calls [1] that correspond to the queries above. These queries constitute an important part of what is collectively known as *language services* and can be issued by both online IDEs, sophisticated code editors such as SublimeText, and full-fledged IDEs such as Eclipse and Visual Studio. Figure 6 shows examples of an IDE in action.

Soundness: Given the nature of such tasks that focus on *program understanding*, the goal is not to always be absolutely precise, but to be both useful to the end user and responsive. Our analysis judiciously cuts corners in the spirit of soundness [21]. As the analysis results are used in an advisory role in the context of program understanding in an interactive setting, complete soundness is not the goal. For instance, we do not attempt to model reflective constructs. While we focus on C# as the input language, our work should apply equally well to analyzing large projects in Java and other similar object-oriented languages. It is not, however, our goal to faithfully handle all the tricky language features such as reflection, runtime code generation, and pinvoke-based native calls.

1.3 Contributions

This paper makes the following contributions:

- We propose a distributed static analysis approach, based on the monotone framework, and show how to apply it to call graph construction for answering program understanding and code browsing queries.
- We describe how the analysis is implemented on top of the Orleans distributed programming platform and is deployed on legacy hardware in the cloud using Microsoft Azure.
- We experimentally demonstrate the scalability of our distributed call graph implementation using a range of synthetic and real benchmarks. The results show that our analysis scales well in terms of memory pressure independently of the input size, as we add more machines. Despite using stock hardware and incurring a non-trivial communication overhead, we scale to inputs containing 10 million LOC: for inputs of 1M LOC, the analysis requires at least 4 machines; for 10M LOC, 16 machines. While the communication overhead can become a bottleneck, we show that as the number of machines increases (up to 64), the analysis time generally drops. Depending on the setting, partial results can be queried before the analysis has finished. Lastly, we demonstrate that querying the results can be performed with an acceptable median latency of 15 ms.

2 OVERVIEW

Our goal is to have the analysis backend respond to queries quickly, independently of the input size. Of course, we also need to make sure that the backend does not run out of memory or timeout in some unpredictable way. Our requirements force us to rethink some of the typical assumptions of whole-program analysis.

```

1: while |MQ| > 0 do
2:   ⟨a, m⟩ := MQ.choose()
3:   v := UNPACK(m) ⊔ VALUE[a]
4:   if v ⊑ VALUE[a] then
5:     continue
6:   end if
7:   v' := TF[a](v)
8:
9:   if v ⊑ v' then
10:    U := DELTA(v, v')
11:    for each u in U do
12:      MQ := MQ ∪ PACK(a, u)
13:    end for
14:    VALUE[a] := v'
15:   end if
16: end while

```

Figure 1: Distributed worklist algorithm.

2.1 Analysis Design Principles

We use a *distributed actor model* [2] as the basis of our distributed static analysis engine. For a program written in an object-oriented language such as Java or C#, a natural fit is to have an actor per *method* within the program. We could choose to have an actor per *class* in a program, or any other well-defined program entity. These actors are responsible for receiving messages from other actors, processing them using local state (a representation of the method body, for instance), and sending information to other methods that depend on it. For example, for a call graph construction analysis, actors representing individual methods may send messages to actors for their callers and callees. Our analysis design adhere to the following distilled principles.

Minimal in-memory state per actor: We want to “pack” as many actors per machine as possible without creating undue memory pressure, leading to swapping, etc.

Design for lightweight serialization: We have designed our analysis so that the updates sent from one actor to another are generally small and easily serialized. There is minimal sharing among actors, as actor holds on to its local state and occasionally sends small updates to others. The same principle applies to persistent per-actor state as well. We only serialize the bare minimum to disk, before the actor is put to sleep. This can happen when the actor runtime decides to page an actor out due to memory pressure or lack of recent use.

State can be recomputed on demand: In a distributed setting, we have to face the reality that processes may die due to hardware and/or software faults. It is therefore imperative to be able to recover in case of state loss. While it is possible to commit local state to persistent store, we eschew the overhead of such an approach and instead choose to recompute per-node state on demand.

Locality optimizations to minimize communication: We attempt to place related actors together on the same machine. In the case of a call graph analysis, this often means that entire strongly connected components co-exist on the same physical box, which minimizes the number of messages that we actually need to dispatch across the network.

2.2 Distributed Worklist Algorithm

We now present a high-level view of a distributed analysis problem as a pair $\langle A, L \rangle$ where:

- A is a set of actors distributed in a network.
- $\langle L, \sqsubseteq, \sqcup \rangle$ is a complete semi-lattice of finite height³.

Each actor $a \in A$ has the following associated functions:

- $VALUE[a] = v \in L$ is the local state of actor a ;

³The finite height requirement can be avoided with the use of a widening operator.

- $TF[a] : L \mapsto L$ is the transfer function for the local computation performed within actor a . We assume all TF are monotone;

The following helper functions are for communicating state changes among actors:

- $DELTA(v, v')$ computes a set U of (global) updates required when switching from local state v to $v' \in L$;
- $PACK(a, u)$ is a function that given an update at actor $a \in A$ produces one or several messages to communicate to other actors.
- $UNPACK(m)$ is a function that unpacks a message and returns a value $v \in L$.

Figure 1 shows the pseudocode for a distributed worklist algorithm. The algorithm makes use of a global message queue, denoted as MQ ⁴. The queue is initialized with a set of starting messages that will depend on the actual analysis instance.

2.3 Termination and Non-Determinism

We would like to show that the presented distributed worklist algorithm terminates.

Let H denote the (finite⁵) height of semi-lattice L and let $N = |A|$. Consider iterations through the loop on line 1. Let’s consider two sets of sequences of iterations, I_1 are iterations that lead to a value increase on line 7 and I_2 are those that do not.

We can have at most $H \times N$ iterations in I_1 given the finite size of the lattice. For iterations in I_2 , the size of MQ decreases because at least one message is consumed but it does not generate other messages. We consider two possibilities:

- Starting from some iteration i , we only have iterations in I_2 . This, however, means that on every iteration the size of MQ decreases, until it eventually becomes empty.
- The other possibility is that we will have an infinite number of iterations in I_1 . This is clearly impossible because the size of I_1 is bounded by $H \times N$.

It is important to emphasize the difference between this distributed algorithm and a single-node worklist approach. If a message is in flight, we do not wish the program analysis to terminate. However, detecting the emptiness of MQ is not trivial, so in practice we must have an effective means for detecting termination. We make use of an *orchestrator* mechanism for termination detection, as described in Section 4.2.

While the algorithm in Figure 1 reaches a fixpoint independently of the arrival order of messages, it is natural to ask whether that is the only fixpoint that can be reached. Given that $TF[a]$ is monotone and L is of a finite height the uniqueness of least fixpoint is guaranteed [9, 18].

3 CALL GRAPH ANALYSIS

In this section we present an instantiation of the general framework described in the previous section for computing call graphs. Our analysis is a distributed interprocedural inclusion-based static analysis inspired by the Variable Type Analysis (VTA) presented in [36]. This flow-insensitive analysis computes the set of potential types

⁴Note that MQ is a *mathematical abstraction*: we do not actually use a global message queue in our implementation. Conceptually, we can think of a (local) worklist maintained on a per-actor basis. Termination is achieved when all the worklists are empty.

⁵Note that our approach can also terminate for an infinite height lattice with the use of a widening operator.

for each *object reference* (variable, field, etc.) by solving a system of inclusion constraints. Because it propagates type constraints from object allocation sites to their uses, this kind of analysis is sometimes referred to as *concrete type analysis*.

3.1 Program Representation

Propagation graphs: At the method level, the inclusion-based analysis is implemented using a data structure we call a *propagation graph* (PG) [36]. A PG is a directed graph used to “push” type information to follow data flow in the program, as described by analysis rules. Our analysis naturally lands itself to incrementality, although we do not evaluate this experimentally in this paper. A typical change in the program would require often minimal recomputation within the modified code fragment as well as propagation of that information to its “neighbors”. Propagation graphs support incremental updates since the propagation of information is triggered when a new type reaches a node.

Terminology: More formally, let $PG = \langle R, E \rangle$ where R denotes a set of nodes representing *abstract locations* in the method (such as variables and fields) and E refers to a set of edges between them. An edge $e = (v_1, v_2) \in E$ connects nodes in the PG to model the potential flow of type information from v_1 to v_2 . Essentially, an edge represents a rule stating that $\text{Types}(v_2) \supseteq \text{Types}(v_1)$ (e.g. $v_2 = v_1$). To model interprocedural interaction, the PG also includes nodes representing method invocations (inv_{loc}) and return values (rv). Finally, $I \subseteq R$ denotes the set of invocations. Let T be the set of all possible types, $dType$ contains declared types (compile-time types) for abstract locations and $Types$ denotes concrete types inferred by our analysis.

3.2 Analysis Phases

In the actor model, the choice of granularity is key for performance. We decided to use one actor per method, although other design decisions such as one actor per class are also possible. Each method-level actor contains a PG that captures type information that propagates through the method. The analysis starts by analyzing an initial set of root methods M_0 . We describe both intra- and interprocedural processing below.

3.2.1 Intraprocedural Analysis.

This phase is the responsible of computing the local state of an actor representing a method.

Instantiating the problem: The lattice L for our analysis consists of a mapping from abstract locations to sets of possible types and is defined as

$$L = \langle \text{Types} : R \mapsto 2^T, \sqsubseteq_{type}, \sqcup_{type} \rangle$$

with \sqsubseteq_{type} defined as

$$l_1 \sqsubseteq_{type} l_2 \text{ iff } l_1.\text{Types}(r) \subseteq l_2.\text{Types}(r), \forall r \in R$$

and \sqcup_{type} defined as $l_1 \sqcup_{type} l_2 = l_3$ where

$$l_3.\text{Types}(r) = l_1.\text{Types}(r) \cup l_2.\text{Types}(r), \forall r \in R.$$

Analysis rules that compute $TF[a]$ are summarized in Figure 2 and cover the typical statement types such as loads, stores, allocations, etc. Object dereferences (i.e., $v.f$) are represented by using the

$$\begin{aligned} v_1 = v_2 &\implies \text{Types}(v_1) \supseteq \text{Types}(v_2) \\ v_1 = v_2.f &\implies \text{Types}(v_1) \supseteq \text{Types}(dType(v_2).f) \\ v_1.f = v_2 &\implies \text{Types}(dType(v_1).f) \supseteq \text{Types}(v_2) \\ v = \text{new } C() &\implies C \in \text{Types}(v) \\ \text{return } v &\implies \text{Types}(rv) \supseteq \text{Types}(v) \\ loc : v = v_0.m(v_1 \dots v_n) &\implies \text{Types}(inv_{loc}) \supseteq \bigcup_{j=0..n} \text{Types}(v_j) \end{aligned}$$

Figure 2: VTA analysis rules.

name of the class defining the field. That is, the analysis is field-sensitive but not object-sensitive. In the case of invocations there is an inclusion relation to model the flow of all the arguments to the invocation abstract location $inv_{loc} \in I \subseteq R$. Note that the left-hand side v of the invocation is not updated by the rule since it depends on the result of the invoked method. This will be handled by interprocedural analysis.

Notice that $TF[a]$ is monotonic because the propagation of types never removes a type and L satisfies the finite-height condition because it is a finite lattice.

3.2.2 Interprocedural Analysis. Once the intraprocedural phase finishes, relevant updates must be communicated to the corresponding methods (callees and callers). As mentioned, the analysis considers invocations using the set $I \subseteq R$. To handle callers’ updates, we need to extend the lattice to include the caller’s information for the current method. This has the form $\langle m, lhs \rangle$, where $m \in A$ denotes the caller’s name and $lhs \in R$ represents the left-hand side of the invocation made by the caller. The extended lattice is shown below.

$$\begin{aligned} L &= \langle \text{Types} : R \mapsto 2^T \times \text{Callers} : 2^{A \times R}, \sqsubseteq, \sqcup \rangle \\ l_1 \sqsubseteq l_2 &\text{ iff } l_1 \sqsubseteq_{type} l_2 \wedge \\ & l_1.\text{Callers} \subseteq l_2.\text{Callers} \\ l_1 \sqcup l_2 &= (ts, cs) \text{ where} \\ & ts = l_1 \sqcup_{type} l_2 \wedge \\ & cs = l_1.\text{Callers} \cup l_2.\text{Callers} \end{aligned}$$

A message m has the form $\langle kind, d, data \rangle$, where $kind \in \{\text{callMsg}, \text{retMsg}\}$ is the kind of message, $d \in A$ is the destination actor and $data$ is a tuple.

Instantiating DELTA: In Figure 3a we show the definition of the *DELTA* operation described in Section 2. It computes the set of invocations that were affected by the propagation. An invocation is affected if the set of types flowing to any of its parameters grew. Additionally, we also must consider changes in types that the return value may correspond to, since they need to be communicated to the callers.

Instantiating PACK: Figure 3b shows a definition of *PACK*. This function is in charge of converting local updates to messages that can be serialized and sent to other actors. For each invocation, the analysis uses the computed type information of the receiver argument to resolve potential callees.

Then, it builds a caller message including the potential types for each argument. Those types will be added to the set of types of the parameters on the caller actor. In case of an update in return value it builds a message to inform the caller about changes to the return value’s types. This message includes the (original) caller’s left-hand side, so that the caller can update its types.

$$\begin{aligned} \text{let } \text{diff}(v, v', r) &:= v'.\text{Types}(r) - v.\text{Types}(r) \\ \text{let } \text{Inv}(v, v') &:= \{inv \mid inv \in I \wedge \text{diff}(v, v', inv) \neq \emptyset\} \\ \text{let } \text{Rv}(v, v') &:= \begin{cases} \{rv\} & \text{if } \text{diff}(v, v', rv) \neq \emptyset \\ \emptyset & \text{otherwise} \end{cases} \\ \text{DELTA}(v, v') &\stackrel{\text{def}}{=} \text{Inv}(v, v') \cup \text{Rv}(v, v') \end{aligned}$$

(a) Definition of $\text{DELTA}(v, v')$

$$\begin{aligned} \text{let } \text{callees}(inv) &:= \{C.m \mid C \in I.\text{Types}(\overline{\text{args}(inv)_0})\} \\ \text{let } \text{callMsg}(a, inv) &:= \langle a, \text{lhs}(inv), I.\text{Types}(\overline{\text{args}(inv)}) \rangle \\ \text{let } \text{callMsgs}(a, inv) &:= \{\langle \text{callMsg}, d, \text{callMsg}(a, inv) \rangle \mid d \in \text{callees}(inv)\} \\ \text{let } \text{returnMsg}(a, c) &:= \langle a, \text{lhs}(c), I.\text{Types}(rv) \rangle \\ \text{let } \text{retMsgs}(a) &:= \{\langle \text{retMsg}, \text{method}(c), \text{returnMsg}(a, c) \rangle \mid c \in I.\text{Callers}\} \\ \text{PACK}(a, u) &\stackrel{\text{def}}{=} \begin{cases} \text{callMsgs}(a, u) & \text{if } u \in I \\ \text{retMsgs}(a) & \text{if } u = rv \end{cases} \end{aligned}$$

(b) Definition of $\text{PACK}(a, u)$. $I.\text{Types}(\overline{\text{args}})$ is the lifting of $I.\text{Types}$ to the list of arguments, it returns a lists of set of types. Given $inv = \langle v = v_0.m(v_1 \dots v_n) \rangle$, $\text{args}(inv) = [v_0, v_1, \dots, v_n]$, $\text{lhs}(inv) = v$. For a caller $c = (m, \text{lhs}) \in I.\text{Callers}$, $\text{method}(c) = m$, the caller's name and $\text{lhs}(c) = \text{lhs}$, the left-hand side of the original invocation made by the caller.

$$\begin{aligned} \text{let } l_1.\text{Types}(r) &= \begin{cases} \text{argTypes}(m)_i & \text{if } r = p_i \\ \emptyset & \text{otherwise} \end{cases} \\ \text{let } l_1.\text{Callers} &= \{ \langle \text{sender}(m), \text{lhs}(m) \rangle \} \\ \text{let } l_2.\text{Types}(r) &= \begin{cases} \text{retTypes}(m) & \text{if } r = \text{lhs}(m) \\ \emptyset & \text{otherwise} \end{cases} \\ \text{UNPACK}(m) &\stackrel{\text{def}}{=} \begin{cases} l_1 & \text{if } \text{kind}(m) = \text{callMsg} \\ l_2 & \text{if } \text{kind}(m) = \text{retMsg} \end{cases} \end{aligned}$$

(c) Definition of $\text{UNPACK}(m)$. For a message $m = \langle \text{callMsg}, d, \langle a, \text{lhs}, [ts_0, ts_1, \dots, ts_n] \rangle \rangle$ $\text{argTypes}(m)_i = ts_i$, the set of potential types for the i^{th} argument p_i . $\text{lhs}(m) = \text{lhs}$, $\text{sender}(m) = a$. For a return message $m' = \langle \text{retMsg}, d, \langle a, \text{lhs}, ts \rangle \rangle$, $\text{retTypes}(m') = ts$ is the set of potential types of the method's return value.

Figure 3: Defining DELTA , UNPACK , and PACK .

Instantiating UNPACK : Function UNPACK in Figure 3c is responsible for processing messages received by an actor. This function converts a message into a value in the lattice of the local analysis that will be then joined into the local state. A message can be either a *call message* (i.e., an invocation made by a caller) or a *return message* (i.e., to inform a change in the callee's return value). For call messages we produce an element that incorporates the types for each call argument into the method parameters. We also update the set of callers. For return messages we need to update the left-hand side of the invocation with the potential types of the return value.

Example 1 This example illustrates the advantage of using concrete types as opposed to declared types to obtain more precision. Consider the small program in Figure 4a. In Figure 4b we show the propagation graphs for both methods. As the analysis starts, only the left-hand sides of allocations (lines 2 and 11) contain types.

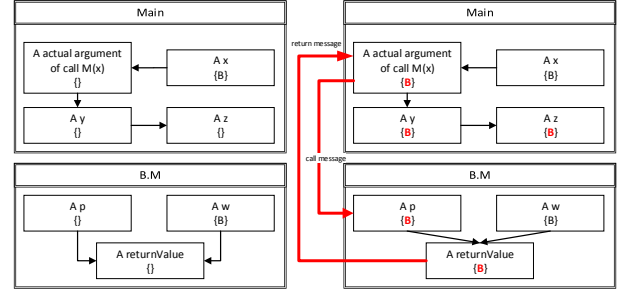
During propagation, type B flows from variable x into an invocation of M as an argument. This triggers a message to the actor for method B.M. The flow through parameter p and w makes the return

```

1 public static void Main() {
2   A x = new B(); // allocation
3   A y = x.M(x);
4   A z = y;
5 }
6 public class A {
7   public abstract A M(A p);
8 }
9 public class B : A {
10  public override A M(A p) {
11    A w = new B(); // allocation
12    return (p != null) ? p : w;
13  }
14 }

```

(a) Code example for interprocedural propagation.



(b) PGs for methods Main and B.M before (left) and after (right) the propagation for the code in Figure 4a.

Figure 4: Code and propagation graph for Example 1.

value of B.M to contain type B. This in turn triggers a return message that adds B to the types of y. This propagates to z. Concrete type analysis produces results that are more accurate for y, z, etc. than what we can obtain from their declared types. ■

Type approximation: In the interprocedural stage, our analysis sends information about concrete parameter types to its callees. However, when it comes to complex, nested objects, this information is potentially insufficient, as it only concerns one level of the object hierarchy. Consider the following example:

```

void Main {
  A x = new B();
  x.f = new B();
  y = M(x)
}

void M(A p) {
  A z = p.f;
  return z;
}

```

Function PACK will create a message that propagates the type of x into M and UNPACK will discover the type of p to be B. However, no information is given for the type of p.f, potentially leading to unsoundness. Instead of sending information about nested fields, which leads to increased message sizes, we opted to use the type of p.f given by a distributed version of the Rapid Type Analysis [5] that runs simultaneously on each method-actor; when RTA provides no useful information, we fall back on declared types. We did not observe imprecision caused by this over-approximation.

3.3 Other Uses of the Analysis Framework

The distributed algorithm in Figure 1 can be instantiated for other program analyses that follow the same design principle. For instance, consider an inclusion-based analysis like Andersen's points-to [4]. A possible instantiation may be as follows: (1) Each actor represents a method; (2) The transfer function implements Andersen's inclusion rules locally and, in case there is a change in an

argument of a method invocation, produces an update message to be sent to the potential callees.

Similarly, by just replacing the inclusion rules with unification rules in the transfer function, we can turn it into a unification based points-to analysis like Steensgaard's [35]. Context-sensitivity can be achieved by representing different *context* \times *method* combinations with different actors.

It is worth noticing that our analysis has similar characteristics as standard dataflow analyses, but an ordering on how information flows between the actors cannot be assumed. We envision future work where our distributed back-end would be combined with a natural front-end for this kind of analysis that uses Datalog, as previously proposed for single-machine analysis [17]. However, as we describe in Section 1.2, our evaluation in Section 5 focuses on quickly answering interactive questions related to call graph resolution in the context of a IDE.

4 IMPLEMENTATION

We implemented a prototype of our distributed approach⁶ to analyze large-scale projects written in C#. This prototype relies on Roslyn [27], a compiler framework for analyzing C# code and the Orleans framework [6], an implementation of a distributed actor model that can be deployed in the cloud. Although other deployment options such AWS are possible, for this paper we used Azure as a platform for running our experiments.

4.1 Orleans and the Actor Model

Orleans [6] is a framework designed to simplify the development of distributed applications. It is based on the abstraction of virtual *actors*. In Orleans terminology, these actors are called *grains*. Orleans solves a number of the complex distributed systems problems, such as deciding where – on which machine – to allocate a given actor, sending messages across machines, etc., largely liberating developers from dealing with those concerns. At the same time, the Orleans runtime is designed to enable applications that have high degrees of responsiveness and scalability. Grains are the basic building blocks of Orleans applications and are the units of isolation and distribution. Every grain has a unique global identity that allows the underlying runtime to dispatch messages between actors. An actor encapsulates both behavior and mutable local state. State updates across grains can be initiated using messages.

The runtime decides which physical machine (*silo* in Orleans terminology) a given grain should execute on, given concerns such as memory pressure, amount of communication between individual grains, etc. This mechanism is designed to optimize for communication locality because even within the same cluster the amount of cross-machine messages are considerably smaller than the amount of local messages, within the same machine.

We follow a specific strategy in organizing grains at runtime. This strategy is driven by the input structure. The input consists of an MSBuild *solution*, a `.sln` file that can be opened in Visual Studio. Each solution consists of a set of *project files*, `*.csproj`, which may depend on each other. Roslyn allows us to enumerate all project files within a solution, source files within a project, classes within

a file, methods within a class, etc. Furthermore, Roslyn can use its built-in C# compiler to compile sources on the fly. We define grains for solutions, projects and methods. We did not find it necessary to provide grains for classes and other higher-level code artifacts such as namespaces.

A solution grain is a singleton responsible for maintaining the list of projects and providing functionality to find methods within projects; A project grain contains the source code of all files for that project and provides functionality to compute the information required by method grains (e.g., to build propagation graphs by parsing the method code) as well as type resolution (e.g., method lookup, subtyping queries, etc). Finally, a method grain is responsible for computing the local type propagation and resolving caller/callees queries; it stores type information for abstract locations within the method.

The solution grain reads the `*.sln` file from cloud storage; in our implementation we used Azure Files, but other forms of input that support file-like APIs such as GitHub or Dropbox are also possible. Project grains read `*.csproj` files and also proceed to compile the sources contained in the project to get a `Roslyn Compilation` object. This information is only contained in the project grain to minimize duplication. To obtain information about the rest of the project, method grains can consult the project grain. We use caching to reduce the number of messages between method and project grains.

Example 2 To illustrate persistent state for a typical method grain, consider the example in Figure 4a. The state of both methods is as follows.

Method Main:

```
Callers = {}
Types = {(x,{B}), (y,{B}), (z,{B}), (3,{B})}
```

Method B.M:

```
Callers = {(A.Main, y)}
Types = {(p,{B}), (w,{B}), (returnValue,{B})}
```

This minimal state is easily serialized to disk if the grains are ever deactivated by the Orleans runtime. Orleans deactivates grains when they aren't used for a long time, however, this never happened in our experiments. ■

4.2 Distributed Analysis Challenges

Implementing a distributed system like ours is fraught with some fundamental challenges.

Reentrancy: Since the callgraph can have cycles, a grain can start a propagation which will in turn eventually propagate to the original method. However, since Orleans uses turn-based concurrency this will create a deadlock. Even without recursion it is possible for a method grain that is currently being processed to receive another message (i.e. a return message).

Termination: In a distributed setting, detecting when we achieve termination is not so easy. This is in part because even if all the local worklists are empty, we may have messages in flight or those that have been delayed.

A naïve implementation is not going to work well because of reentrancy issues: we can block waiting for a message that waits for our response. In our implementation, we use *orchestrators* to

⁶Source code and benchmarks available on GitHub at: <https://github.com/too4words/Call-Graph-Builder-DotNet>.

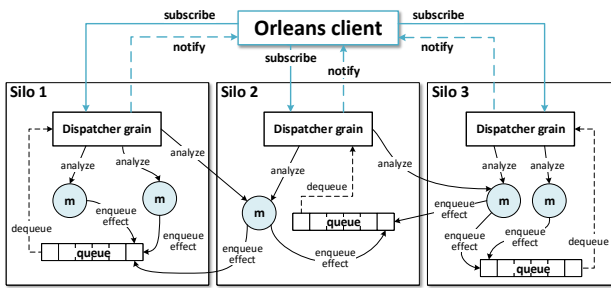


Figure 5: The multi-queue approach, illustrated. Method grains are circles shown in light blue. Solid and dashed arrows represent standard invocations and callbacks respectively. Each silo has each own dispatcher grain.

establish some degree of centralized control over the propagation process. Grains communicate with an orchestrator exclusively, instead of communicating with each other peer-to-peer. This avoids the issue of reentrancy by construction; only the orchestrator can send messages to grains via a single *message queue*. The orchestrator keeps track of the outstanding tasks and can therefore detect both termination and prevent reentrant calls from taking place.

The key disadvantage of this design is that it is possible to have a great deal of *contention* for access to the orchestrator. We observed this in practice, suggesting a different variant of this idea. We use a *collection* of queues distributed across the distributed system. Each method grain is a potential producer of *effects* to be processed by other method grains. To avoid reentrancy, this information is not sent directly to the target method grain but it is enqueued in one of the queues in a round robin fashion. The information is then consumed by *dispatchers grains* that pull the data from the queues and deliver it to the corresponding method grains; this is illustrated in Figure 5.

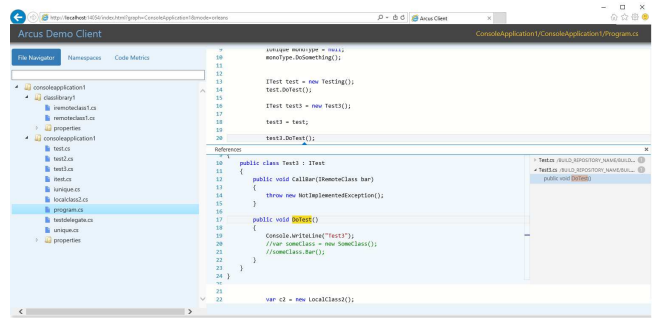
Using this mechanism we avoid both reentrancy, bottlenecks and single points of failure. The drawback is that detecting termination is more complex. For that, we use timers to determine when a dispatcher becomes idle (i.e., inactive longer than a predetermined threshold), at which point we notify the client. The analysis finishes when the client is sure that all dispatchers are idle⁷. In practice, we set the number of queues to be four times higher than the number of worker VMs (for example, 128 queues for 32 worker VMs) and set the termination threshold to 10 seconds.

4.3 Deployment Details

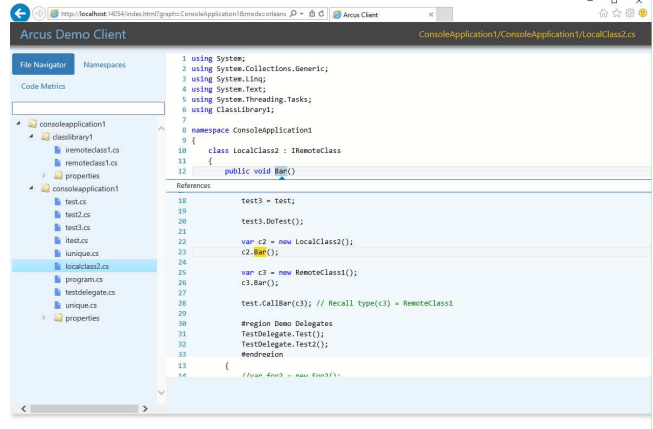
Our analysis is deployed in Azure as illustrated in Figure 7. On the left, there is the analysis client such as an IDE or a code editor like SublimeText. The cluster we used consists on one *front-end VM* and a number of worker VMs. The client used REST requests to communicate to the front-end VM. The job of the front-end VM is to (1) accept and process external analysis client requests; (2) dispatch jobs to the worker VMs and process the results; and (3) provide a Web UI with analysis results and statistics.

Interactive deployment within an IDE: In Figure 6 we show two screen-shots of an experimental IDE prototype that uses the

⁷We have a mechanism to detect when an idle dispatcher becomes active again.



(a) Visualizing callees: call site on line 20 invokes function `DoTest`.



(b) Visualizing callers: method `Bar` is called on line 23.

Figure 6: An experimental online IDE that uses analysis for resolving references for callees and callers.

API exposed by our analysis to resolve callers/callees queries. We should point out that the precision achieved by our analysis is enough for the autocomplete task.

5 EVALUATION

We aim to answer the following three research questions.

- RQ1:** Is our analysis capable of handling arbitrary amounts of input (i.e., more lines of code, files, projects, etc.) by increasing the number of worker VMs, without running out of memory?
- RQ2:** While the communication overhead can become significant, as more worker VMs are added, does an increase in the number of worker VMs significantly increase the overall analysis times?
- RQ3:** Is the analysis query latency small enough to allow for interactive use⁸?

The focus of our analysis is on being used in an interactive setting. Given the low latency times we can use our analysis interactively as a replacement of source code browsers such as `http://source.roslyn.io`. This browser provides code search and basic navigation facilities but lacks more advanced features like actual callers/callees inspection/navigation that we can provide

⁸Generally, query latencies of 10 to 20 ms are considered to be acceptable.

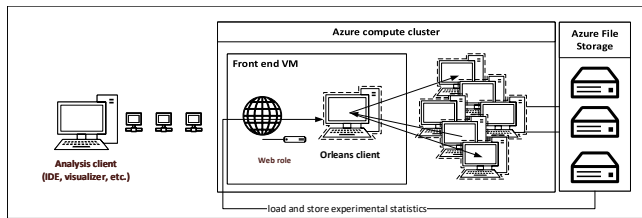


Figure 7: Azure-based deployment of our analysis. Actual work happens within worker VMs. The analysis client interacts with the cluster via a front-end VM.

with our analysis. At the same time, we are not as concerned about the completion time for the analysis as a whole as we are about its memory requirements on legacy VMs. Even if it takes longer to process, our goal is to engineer an always-on system that responds to messages sent to the cloud to service user requests, in the context of code browsing, and other tasks listed in Section 1.2. This work was performed in collaboration with the Roslyn team, and while we have not performed user studies, we believe latency numbers (most queries took under 20 ms) to be more than acceptable for interactive use.

5.1 Experimental Setup

All the experiments presented in this paper were executed in the cloud, on a commercially available Azure cluster. We could also have used an AWS cluster, as our dependency on Azure is small. The Azure cluster we used for the experiments consists on one front-end VM and up to 64 worker role VMs. The front-end VM is an Azure VM with 14 GB of RAM (this is an A4\ExtraLarge VM in Azure parlance⁹). Each worker role is an Azure VM with 7 GB of RAM (called A3\Large in Azure). For benchmarking purposes, we run our analysis with configurations that include 1, 2, 4, 8, 16, 32, and 64 worker VMs. To collect numbers for this paper, we used a custom-written experimental controller as our analysis client throughout this section; this setup is illustrated in Figure 7. The controller is scripted to issue commands to analyze the next `.sln` file, collect timings, etc.

We heavily instrumented our analysis to collect a set of relevant metrics. We instrumented our analysis code to measure the analysis elapsed time. We introduced wrappers around our grains (solution, project, and method grains) to distinguish between local messages (within the same VM) and network messages. Using Orleans-provided statistics, we measured the maximum memory consumption per VM. Lastly, we also have added instrumentation to measure query response times. While these measurements are collected at the level of an individual grain, we generally wanted to report aggregates. To collect these, we post grain-level statistics to a special auxiliary grain.

5.2 Benchmarks

For our inputs, we have used two categories of benchmarks, *synthetic* benchmarks we have generated specifically to test the scalability of our call graph analysis and a set of 3 real applications

written in C# that push our analysis implementation to be as complete as possible, in terms of handling tricky language features such as delegate, lambdas, etc. and see the impact of dealing with polymorphic method invocations. In all cases, we start with a solution file (`.sln`) which references several project files (`.csproj`), each of which in turn references a number of C# source files (`.cs`).

Benchmark	LOC	Projects	Classes	Methods
X1,000	9,196	10	10	1,000
X10,000	92,157	50	50	10,000
X100,000	904,854	100	100	100,000
X1,000,000	9,005,368	100	100	1,000,000

Figure 8: Information about synthetic benchmarks.

Synthetic benchmarks: We designed a set of synthetic benchmarks to test the scalability of our analysis approach. These are solution files generated to have the requisite number of methods (for the experiments, we ranged that number between 1,000 and 1,000,000).

The Figure 8 summarizes some statistics about the synthetic projects we have used for this evaluation. Synthetic benchmarks were generated to have the requisite number of methods, organized in classes and projects according to a maximum predefined number. Each method invokes between 1–11 other methods, with the only requirement that all methods be reachable. While synthetic programs measure the input size in a controlled way (e.g., LOCs, methods, invocations), the real benchmarks measure the overall complexity (e.g., polymorphism, complex program constructs).

Real-world benchmarks: We have selected several large open-source projects from GitHub for our analysis. A summary of information about these programs is shown in Figure 9. We tried to focus on projects that are under active development. To illustrate, one of our benchmarks, Azure Powershell is one of the most popular projects written in C# on GitHub. According to the project statistics, over a period of one month, 51 authors have pushed 280 commits to the main branch and 369 commits to all branches. There have been 342,796 additions and 195,366 deletions. We picked solution `ResourceManager.ForRefactoringOnly.sln` from Azure Powershell because it is the only one that contains all the projects. Generally, discovering good root methods to serve as starting points for the call graph analysis is not trivial. Because there is no natural `Main` method in several of these projects, we have decided to use as entry points the included *unit tests*, *event handlers*, and other public methods within the project to increase the number of methods our analysis reaches¹⁰.

[RQ1]: Scales with input size: To answer RQ1, we measured the memory consumption of each VM and computed the average and maximum memory consumption across all VMs. Figure 10 shows the *average* memory consumption for each benchmark during the run, for each experimental configuration, i.e. number of worker VMs used. As can be observed from the chart, the memory consumption decreases steadily as the number of worker VMs increases. Recall that worker VMs come equipped with 7 GB of memory, so these memory consumption numbers are nowhere near that limit.

⁹Up-to-date VM specifications are available at: <https://azure.microsoft.com/en-us/documentation/articles/virtual-workerVMs-size-specs/>.

¹⁰Note that we do not analyze libraries provided as DLLs; our analysis implementation works at the source level only.

Benchmark	URL	LOC	Projects	Classes	Methods	Main	Test	Event handlers	Public	Total	Reachable methods
Azure-PW	https://github.com/Azure/azure-powershell	416,833	60	2,618	23,617	0	997	1	18,747	18,759	23,663
ShareX	https://github.com/ShareX/ShareX	110,038	11	827	10,177	2	0	1,122	6,257	7,377	10,411
ILSpy	https://github.com/icsharpcode/ILSpy	300,426	14	2,606	25,098	1	0	119	14,343	14,498	21,944

Figure 9: Summary of information about real-world projects from GitHub. The number of reachable methods include also library methods invoked by the application methods. Note that some application methods might not be reachable.

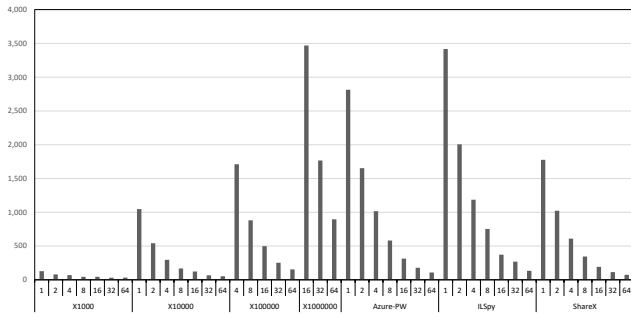


Figure 10: Average memory consumption in MB, for each benchmark as a function of the number of worker VMs. We see a steady decrease across the board.

Looking at Figure 10, we can see peaks of about 3.2 GB for a single worker VM while analyzing X1,000,000¹¹.

These experiments naturally highlight the notion of analysis *elasticity*. While we run the analysis with different number of VMs set for the sake of measurement, in reality, more machines would be added (or removed) due to memory pressure (or lack thereof) or to respond to how full analysis processing queues get. We can similarly choose to increase (or decrease) the number of queues and dispatchers involved in effect propagation. It is the job of the Orleans runtime to redistribute the grains to update the system with the new configuration.

RQ1: capable of handling input size?

The memory consumption per worker VMs steadily decreases as the number of worker VMs increases.

[RQ2]: Scales with the # of worker VMs: To answer RQ2, we proceeded to measure the total elapsed analysis time for each benchmark on all the configurations. Figure 11 shows the elapsed analysis time *normalized* by the number of methods in the input¹². Note that the real-world benchmarks shown on the right-hand side of the chart, despite containing fewer methods, require more time than the synthetic benchmarks with 100,000 methods. This is simply because of the analysis time that goes into analyzing more complex method bodies. Real-world benchmarks allocate more objects per method, involving more type propagation time, and perform more virtual invocations, adding to the method resolution time, while the synthetic benchmarks only perform static invocations and allocate

¹¹Note also that for that benchmark, we needed to use at least 16 worker VMs to fit all the methods into (their shared) memory. We needed at least 4 worker VMs for X100,000.

¹²Wall clock times range between less than 1 minute (64 VMs) to about 5 minutes (1 VM) in ShareX and 9 to 20 minutes in ILSpy. For other benchmarks elapsed time is typically less than 5 minutes for 16 VMs, except X1,000,000 that takes about 1 hour (40 minutes in 64 VMs).

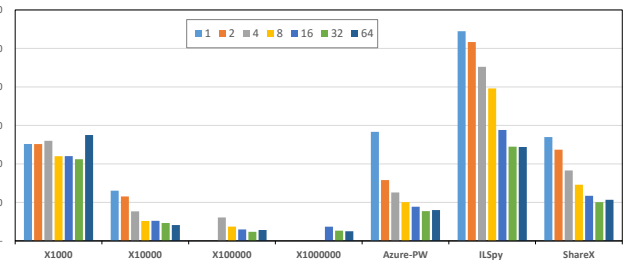
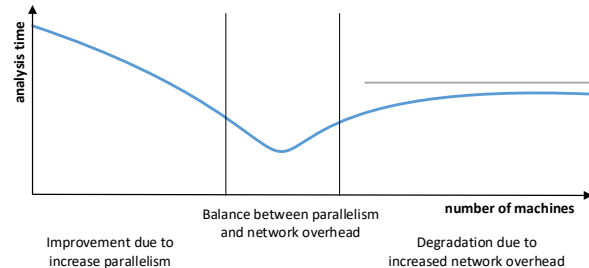


Figure 11: Elapsed analysis time in ms, as a function of the number of worker VMs per test, normalized by the number of reachable methods. The number of worker VMs is indicated in color in the legend above the figure.

relatively few objects. As the number of worker VMs increases, we see a consistent drop in the normalized analysis times. However, this effect generally diminishes after 16 VMs. This has to do with the tension between more parallel processing power of more machines and the increase in the network overhead, as shown below.



It is instructive to focus on the average number of (unprocessed) messages in the analysis queues. If the queues are *too full*, adding more machines will increase the number of queues, reducing the size of each one. More machines will increase the parallelism because of more dispatchers to process the messages in the new queues. As we add more resources, however, when the queues become *mostly empty*, their associated dispatchers will be mostly idle. So the cluster as a whole will have more computing resources than needed. Additionally, if more machines are added, the probability of sending a message to a grain on the same machine as the sender will be reduced, leading to more network overhead. So after reaching a certain cut-off point, adding more machines is not only not helping the analysis, but starts to degrade its performance.

RQ2: does adding more worker VMs increase analysis time?

Normalized analysis time generally *decreases*, as the number of worker VMs increases, up to a point, where the law of diminishing returns kicks in.

[RQ3]: Fast enough for interactive queries: One of the goals of our approach is to enable interactive queries submitted by an

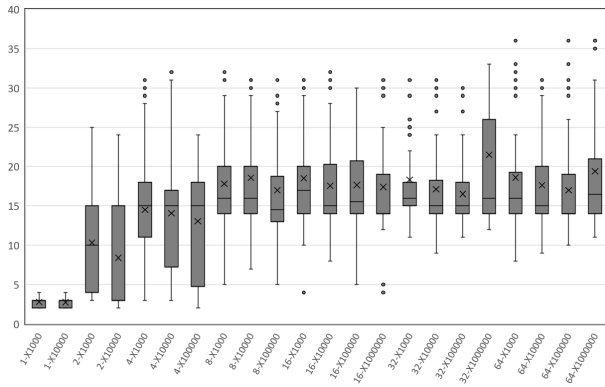


Figure 12: Mean and median query time (ms) for each worker VMs and synthetic test.

analysis client such as an IDE or a sophisticated code editor. In such a setting, responsiveness of such queries is paramount [31]. The user is unlikely to be happy with an IDE that takes several seconds to populate a list of auto-complete suggestions. We want to make sure that as the query times remain tolerable (under 20 ms) even as the size of input increases and the number of VMs goes up.

To evaluate query performance, we automatically generated sequences of 100 random queries, by repeating the following process. We would first pick a random method name from the list of all methods. Then we would (1) Request the solution grain for the corresponding method grain; (2) Select a random invocation from method and request the set of potential callees. In Figure 12 we show the mean and median query times (the latency of the two steps above) for each benchmark and worker VM configuration. Approximately 70% of queries took under 20 ms, 97% under 35 ms, 99.5% under 60 ms. Proper system warm-up may reduce the outliers.

RQ3: is response latency small enough?

The query median response time is consistently between 10 and 20 ms. Increasing the number of worker VMs and the input size does not negatively affect the query response times.

6 RELATED WORK

There exists a wealth of related work on traditional static analysis algorithms such as call graph construction [13, 14, 37]. A comparison of analysis precision is presented in Lhoták *et al.* [20]. As mentioned, our implementation is inspired in VTA [36]. While we have seen dedicated attempts to scale up important analyses such as points-to in the literature, we are unaware of projects that aim to bring call graph analysis to the cloud.

Many projects focus on speeding up the analysis through parallel computation (usually on one machine). Instead, we largely focus on handling memory pressure when analyzing large programs. There are two orthogonal ways to do that: 1) make compositional analysis using specs/summaries (like [8, 38]), abstractions, compact representations [40], demand-driven [34], and other techniques to scale-up; 2) partition analysis memory among several machines. Our analysis focuses on the latter by presenting an approach designed to run on a standard cluster. The engineering challenges are quite different, including state partitioning, decentralized control, number/size of messages sent, termination and network latency.

Scaling Points-to analysis: Hardekopf *et al.* [16] show how to scale up a points-to analysis using a staged approach. Their flow-sensitive algorithm is based on a sparse representation of program code created by a staged, flow-insensitive pointer analysis. They can analyze 1.9M LOC programs in under 14 minutes. The focus (as alleged by the authors) is in obtaining speedups, not in reducing memory pressure. In fact, their largest benchmark required a machine with 100 GB of memory, which is generally beyond the reach of most people. In contrast, we aim at analyzing large programs in clusters of low-cost hardware. Yu *et al.* [42] propose a method for analyzing pointers in a program level by level in terms of their points-to levels. This strategy enhances the scalability of a context- and flow-sensitive pointer analysis and can handle some programs with over a million lines of C code in minutes. The approach is neither parallel non-distributed, the focus is on speedups but some memory is saved by the use of BDDs. Mendez-Lojo *et al.* [26] propose a parallel analysis algorithm for inclusion-based points-to and show a speed up of up to 3× on an 8-core machine on code bases with size varying from 53K LOC to 0.5M LOC. Our focus is on bringing our approach to the cloud using legacy machines and going beyond multicore, to ultimately support code bases of arbitrary size, not being limited by the size of main memory. Voung *et al.* [38] propose a technique that uses the notion of a *relative lockset*, which allows functions to be summarized independent of the calling context. This, in turn, allows them to perform a modular, bottom-up analysis that is easy to parallelize. They have analyzed 4.5 million lines of C code in 5 hours, and, after applying some filters, found several dozen races. Knowing which methods to group together ahead of time would help our actor-machine allocation as well.

Frameworks: Albarghouthi *et al.* [3] present a generic framework to distribute top-down algorithms using a map-reduce strategy. Their focus is in obtaining speed ups in analysis elapsed times; they admit that a limiting scaling factor is memory consumption. McPeak *et al.* [25] propose a multicore analysis that allows them to handle millions LOC in several hours on an 8-core machine. In contrast, our approach focuses on a distributed analysis within a cloud cluster on often less powerful hardware. Boa (Dyer *et al.* [10–12]) is a domain-specific language for mining large code repositories like GitHub. However, while it uses a distributed backend, Boa is not a static analysis. Xie *et al.* [41] propose a bottom-up analysis that benefits from parallel processing on a multicore cluster. They rely on a central scheduler/server, while we use several orchestrators. They use method summaries while we flow the data from one method to another. Finally, we do not rely on a centralized DB, we use grains, which can be persisted or recomputed on-the-fly as needed. Rodriguez *et al.* [32] use an actor model approach in Scala to solve interprocedural distributive subset dataflow problems and evaluate it on an 8-core machine. Our work shares the idea of using actors for analysis but they focused on speed-ups, not memory pressure. Their approach leverages on the use of one computer to implement a global counter to monitor the size of a (virtual) global worklist. In contrast, we run in a cloud setting and must deal with network latency and serialization due to distribution. Pregel [24] is a system for large-scale graph processing that uses an asynchronous message passing model similar to actors, but execution on vertices happens in lockstep; the approach is illustrated for

algorithms such as PageRank and shortest path computation. Grasp [39], is a single-machine, disk-based parallel graph processing system for interprocedural static analyses. Grasp offers two major performance and scalability benefits: (1) the core computation of the analysis is automatically parallelized and (2) out-of-core disk support is exploited if the graph is too big to fit in memory. Our approach focuses on a cloud-based computation, in contrast. Tricorder [33] is a cloud-based tool from Google, designed for scaling program analysis. However, it is meant for simple, intraprocedural analyses, not distributed whole-program analyses.

7 CONCLUSIONS

As modern development is increasingly moving to large online cloud-backed repositories such as GitHub, and Visual Studio Online, is natural to wonder what kind of analysis can be performed on large bodies of code. In this paper, we explore an analysis architecture in which static analysis is executed on a *distributed cluster* composed of legacy VMs available from a commercial cloud provider.

We present a static analysis approach based on the actor model and designed for *elasticity*, i.e. to scale gracefully with the size of the input. To demonstrate the potential of our analysis, we show how a typical call graph analysis can be implemented and deployed in Microsoft Azure. Our call graph analysis implementation is able to handle inputs that are almost 10 million LOC in size. Our results show that our analysis scales well in terms of memory pressure independent of the input size, as we add more VMs. Despite using stock hardware and incurring a non-trivial communication overhead, our processing time for some of the benchmarks of close to 1 million LOC can be about 5 minutes, excluding compilation time. As the number of analysis VMs increases, we show that the analysis time does not suffer. Lastly, we demonstrate that querying the results can be performed with a median latency of 15 ms.

As future work we plan to investigate the performance of other instances of our distributed framework and understand the impact of changing the granularity of actors (e.g., from basic blocks to modules). We want to combine distributed processing with incremental analysis: we are ultimately interested in deploying an Azure-based distributed incremental analysis that can respond quickly to frequent updates in the code repository. We plan to incorporate the analysis into an IDE and to also perform user studies.

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