

Higher-Order, Data-Parallel Structured Deduction

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State-of-the-art Datalog engines include expressive features such as ADTs (structured heap values), stratified aggregation and negation, various primitive operations, and the opportunity for further extension using FFI. Current parallelization approaches for state-of-art Datalogs target shared-memory locking data-structures using conventional multi-threading, or use the map-reduce model for distributed computing. Furthermore, current state-of-art approaches cannot scale to formal systems which pervasively manipulate structured data due to their lack of indexing for structured data stored in the heap.

In this paper, we describe a new approach to data-parallel structured deduction that involves a key semantic extension of Datalog to permit first-class facts and higher-order relations via defunctionalization, an implementation approach that enables parallelism uniformly both across sets of disjoint facts and over individual facts with nested structure. We detail a core language, DL_S , whose key invariant (subfact closure) ensures that each subfact is materialized as a top-class fact. We extend DL_S to $SLOG$, a fully-featured language whose forms facilitate leveraging subfact closure to rapidly implement expressive, high-performance formal systems. We demonstrate $SLOG$ by building a family of control-flow analyses from abstract machines, systematically, along with several implementations of classical type systems (such as $STLC$ and LF). We performed experiments on EC2, Azure, and ALCF’s Theta at up to 1000 threads, showing orders-of-magnitude scalability improvements versus competing state-of-art systems.

CCS Concepts: • **Software and its engineering** → **Compilers; Parallel programming languages; Constraint and logic languages.**

Additional Key Words and Phrases: declarative programming, semantics engineering, data-parallel deduction

1 STRUCTURED DECLARATIVE REASONING

Effective programming languages permit their user to write high-performance code in a manner that is as close to the shape of her own thinking as possible. A long-standing dream of our field has been to develop especially high-level *declarative* languages that help bridge this gap between specification and implementation. Declarative programming permits a user to provide a set of high-level rules and declarations that offer the sought-after solution as a latent implication to be materialized automatically by the computer. The semantics of a declarative language does the heavy lifting in operationalizing this specification for a target computational substrate—one with its own low-level constraints and biases. Modern computers provide many threads of parallel computation, may be networked to further increase available parallelism, and are increasingly virtualized within “cloud” services. To enable scalable cloud-based reasoning, the future of high-performance declarative languages must refine their suitability on both sides of this gulf: becoming both more tailored to human-level reasoning and to modern, massively-parallel, multi-node machines.

Logic-programming languages that extend Datalog have seen repeated resurgences in interest since their inception, each coinciding with new advances in their design and implementation. For example, Bddbddb [80] suggested that binary decision diagrams (BDDs) could be used to compress relational data while permitting fast algebraic operations such as relational join, but required *a priori* knowledge of efficient BDD-variable orderings to enable its compression, which proved to be a significant constraint. LogicBlox and Soufflé [4, 40] have since turned research attention back to semi-naïve evaluation over extensional representations of relations, using compression techniques sparingly (i.e., compressed prefix trees) and focusing on the development of high-performance

shared-memory data structures. Soufflé represents the current state of the art at a low thread count, but struggles to scale well due to internal locking and its coarse-grained approach to parallelism. RadLog (i.e., BigDatalog) [69] has proposed scaling deduction to many-thread machines and clusters using Hadoop and the map-reduce paradigm for distributed programming. Unfortunately, map-reduce algorithms suffer from a (hierarchical) many-to-one collective communication bottleneck and are increasingly understood to be insufficient for leading high-performance parallel-computing environments [3, 64].

Most modern Datalogs are Turing-equivalent extensions, not simply finite-domain first-order HornSAT, offering stratified negation, abstract data-types (ADTs), ad hoc polymorphism, aggregation, and various operations on primitive values. Oracle’s Soufflé has added flexible pattern matching for ADTs, and Formulog [9] shows how these capabilities can be used to perform deductive inference of formulas; it seems likely future (extended) Datalogs will be used to implement symbolic execution and formal verification in a scalable, parallel manner.

In this paper, we introduce a new approach to simultaneously improve the expressiveness and data-parallelism of such deductive logic-programming languages. Our approach has three main parts: (1) a key semantic extension to Datalog, *subfacts* and a *subfact-closure property*, that is (2) implemented uniformly via ubiquitous fact interning, supported within relational algebra operations that are (3) designed from the ground-up to automatically balance their workload across available threads, using MPI to address the available data-parallelism directly. We show how our extension to Datalog permits deduction of structured facts, defunctionalization and higher-order relations, and more direct implementations of abstract machines (CEK, Krivine’s, CESK), rich program analyses (k -CFA, m -CFA), and type systems. We detail our implementation approach and evaluate it against the best current Datalog systems, showing improved scalability and performance.

We offer the following contributions to the literature:

- (1) An architecture for extending Datalog to structured recursive data and higher-order relations, uniform with respect to parallelism, allowing inference of tree-shaped facts which are indexed and data-parallel both horizontally (across facts) and vertically (over subfacts).
- (2) A formalism of our core language, relationship to Datalog, and equivalence of its model theoretic and fixed-point semantics, mechanized in Isabelle/HOL.
- (3) A high-performance implementation of our system, SLOG, with a compiler, REPL, and runtime written in Racket (10.6kloc), Python (2.5kloc), and C++ (8.5kloc).
- (4) An exploration of SLOG’s applications in the engineering of formal systems, including program analyses and type systems. We include a presentation of the systematic development of program analyses from corresponding abstract-machine interpreters—the abstracting abstract machines (AAM) methodology—where each intermediate step in the AAM process may also be written using SLOG.
- (5) An evaluation comparing SLOG’s performance against Soufflé and RadLog on EC2 and Azure, along with a strong-scaling study on the ALCF’s Theta supercomputer which shows promising strong scaling up to 800 threads. We observe improved scaling efficiency and performance at-scale, compared with both Soufflé and RadLog, and better single-thread performance vs. Soufflé when comparing SLOG subfacts to Soufflé ADTs.

2 SLOG: DECLARATIVE PARALLEL DEDUCTION OF STRUCTURED DATA

For Datalogs used in program analysis, manipulation of abstract syntax trees (ASTs) is among the most routine tasks. Normally, to provide such ASTs as input to a modern Datalog engine, one first requires an external flattening tool that walks the richly structured syntax tree and produces a stream of flat, first-order facts to be provided as an input database. For example, the Datalog-based

Java-analysis framework DOOP [12, 13], ported to Soufflé [4] in 2017, has a substantial preprocessor (written in Java) to be run on a target JAR to produce an input database of AST facts for analysis.

A key observation that initially motivated our work into this subject was that although this preparatory transformation is required to provide an AST as a database of first-order facts, the same work could not be done from within these Datalogs because it required generating unique identities (i.e., pointers to intern values) for inductively defined terms. In fact, any work generating ASTs as facts can not be done within Datalog itself but must be an extension to the language. Consider the pair of nested expressions that form an identity function:

$$\begin{array}{ccc}
 (\text{lam } "x" \text{ (ref } "x")}) & \xrightarrow{\text{flattens}} & \begin{array}{l} (= \text{ lam-id (lam } "x" \text{ ref-id)}) \\ (= \text{ ref-id (ref } "x")}) \end{array}
 \end{array}$$

Supplying unique intern values `lam-id` and `ref-id` as an extra column for those relations, and thus permitting them to be linked together, is the substance of this preparatory transformation. Our language, SLOG, proposes this interning behavior for facts be ubiquitous, accounted for at every iteration of relational algebra used to implement the underlying HornSAT fixed point.

In Soufflé, the language has more recently provided abstract data-type (ADT) declarations and struct/record types for heap-allocated values which can be built up into ASTs or other such structured data. These datatypes must be declared and can then be used as \$ expressions within rules; e.g., `$lam("x", $ref("x"))`. The downside of these ADTs in Soufflé is that they are not treated as facts for the purposes of triggering rules and are not indexed as facts, which would permit more efficient access patterns.

Our language, SLOG, respects a *subfact closure property*: every subfact is itself a first-class fact in the language and every top-level fact (and subfact) is a first-class value and has a unique identity (as an automatic column-0 value added to the relation). A clause (`foo x y`) in SLOG, always has an implied identity column and is interpreted the same as `(= _ (foo x y))` if it's missing (where underscore is a wildcard variable). A nested pair of linked facts like `(foo x (bar y) z)` is desugared as `(= _ (foo x id z))` and `(= id (bar y))`. Thus we can represent an identity function's AST in SLOG as the directly nested fact and subfact `(lam "x" (ref "x"))`; under the hood this will be equivalent to two flat facts with a 0-column id provided by an interning process that occurs at the discovery of each new SLOG fact.

- In SLOG, each structurally unique fact/subfact has a unique intern-id stored in its 0 column so it may be referenced as another fact's subfact and treated as a first-class value.
- In SLOG, all data is at once a first-class fact (able to trigger rule evaluation), a first-class value (able to be referenced by other facts/values), and a first-class thread of execution (treated uniformly by a data-parallel MPI backend that dynamically distributes the workload spatially within, and temporally across, fixed-point iterations).

With subfacts as first-class citizens of the language (see Section 3 for details), able to trigger rules, various useful idioms emerge in which a subfact triggers a response from another rule via an enclosing fact (see Section 3.1 for extensions and idioms). Using these straightforward syntactic extensions enables a wide range of deduction and reasoning systems (see Section 4 for a discussion of applications in program analyses and type systems). Because subfacts are first-class in SLOG, rules that use them will naturally force the compiler to include appropriate indices enabling efficient access patterns, and represent thread joins in the natural data-parallelism SLOG exposes. As a result, we are able to show a deep algorithmic improvement and parallelism over current state-of-art systems in the implementations of analyses we generate (see Section 6 for our evaluation with apples-to-apples comparisons against Soufflé and RadLog). In some experiments, SLOG finishes in 4–8 seconds with Soufflé taking 1–3 hours—attesting to the importance of subfact indices. In

others, we observe efficient strong-scaling on up to hundreds of threads, showing the value of our data-parallel backend.

3 STRUCTURALLY RECURSIVE DATALOG

The core semantic difference between SLOG and Datalog is to allow structurally recursive, first-class facts. This relatively minor semantic change enables both enhanced expressivity (naturally supporting a wide range of Turing-equivalent idioms, as we demonstrate in Section 4) and anticipates compilation to parallel relational algebra (which interns all facts and distributes facts via their intern key). In this section, we present the formal semantics of a language we call Structurally Recursive Datalog (henceforth DL_s), the core language extending Datalog to which SLOG programs compile. All of the definitions related to DL_s have been formalized, and all of the lemmas and theorems presented in this section have been formally proven in Isabelle.

Syntax. The syntax of DL_s is shown in Figure 1. As in Datalog, a DL_s program is a collection of Horn clauses. Each rule R contains a set of body clauses and a head clause, denoted by $Body(R)$ and $Head(R)$ respectively. DL_s (and SLOG) programs must also be well-scoped: variables appearing in a head clause must also be contained in the body.

We define a strict syntactic subset of DL_s , DL as the restriction of DL_s to clauses whose arguments are literals (i.e., $\langle Subcl \rangle_{DL} ::= \langle Var \rangle \mid \langle Lit \rangle$). This subset (and its semantics) corresponds to Datalog.

$\langle Prog \rangle ::= \langle Rule \rangle^*$
 $\langle Rule \rangle ::= \langle Clause \rangle \leftarrow \langle Clause \rangle^*$
 $\langle Clause \rangle ::= (tag \langle Subcl \rangle^*)$
 $\langle Subcl \rangle ::= (tag \langle Subcl \rangle^*) \mid \langle Var \rangle \mid \langle Lit \rangle$
 $\langle Lit \rangle ::= \langle Number \rangle \mid \langle String \rangle \mid \dots$

Fig. 1. Syntax of DL_s : *tag* is a relation name.

Fixed-Point Semantics. The fixed-point semantics of a DL_s program P is given via the least fixed point of an *immediate consequence* operator $IC_P : DB \rightarrow DB$. Intuitively, this immediate consequence operator derives all of the immediate implications of the set of rules in P . A database db is a set of facts ($db \in DB = \mathcal{P}(Fact)$). A fact is a clause without variables:

$$Fact ::= (tag \ Val^*) \qquad Val ::= (tag \ Val^*) \mid Lit$$

In Datalog, *Vals* are restricted to a finite set of atoms ($Val_{DL} ::= Lit$). To define IC_P , we first define the immediate consequence of a rule $IC_R : DB \rightarrow DB$, which supplements the provided database with all the facts that can be derived directly from the rule given the available facts in the database:

$$IC_R(db) \triangleq db \cup \bigcup \{ unroll(Head(R)[\overrightarrow{v_i \setminus x_i}]) \mid \{ \overrightarrow{x_i \rightarrow v_i} \} \subseteq (Var \times Val) \wedge Body(R)[\overrightarrow{v_i \setminus x_i}] \subseteq db \}$$

The *unroll* function has the following definition:

$$\begin{aligned} unroll((tag \ item_1 \dots item_n)) &\triangleq \{ (tag \ item_1 \dots item_n) \} \cup \bigcup_{i \in 1 \dots n} unroll(item_i) \\ unroll(v)_{v \in Lit} &\triangleq \{ \} \end{aligned}$$

The purpose of the *unroll* function is to ensure that all nested facts are included in the database as well, a property we call *subfact-closure*. This property is crucial to the semantics of DL_s (and SLOG), because in DL_s , each nested fact is a fact in own right, and not merely a carrier of structured data. Later sections (starting in section 3.1) illustrate the importance of subfact closure by demonstrating how we utilize this behavior to construct idioms that make programming in SLOG more convenient.

The immediate consequence of a program is the union of the immediate consequence of each of its constituent rules, $IC_P(db) \triangleq db \cup \bigcup_{R \in P} IC_R(db)$. Observe that IC_P is monotonic over the

the lattice of databases whose bottom element is the empty database. Therefore, if IC_P has any fixed points, it also has a least fixed point [72]. Iterating to this least fixed point directly gives us a naïve, uncomputable fixed-point semantics for DL_s programs. Unlike pure Datalog, existence of a finite fixed point is not guaranteed in DL_s . This is indeed a reflection of the fact that DL_s is Turing-complete. The DL_s programs whose immediate consequence operators have no finite fixed points are non-terminating.

As discussed earlier, all SLOG databases must be subfact-closed (i.e. all subfacts are first-class facts). We can show that the least fixed point of the immediate consequence operator has the property that it is subfact-closed.

LEMMA 3.1. (*Formalized in Isabelle.*) *The least fixed point of IC_P is subfact-closed.*

It is worth pointing out that the fixed point semantics of Datalog is similar, the only difference being that the *unroll* function is not required, as Datalog clauses do not contain subclauses.

Model Theoretic Semantics. The model theoretic semantics of DL_s closely follows the model theoretic semantics of Datalog, as presented in, e.g., [18]. The *Herbrand universe* of a DL_s program is the set of all of the facts that can be constructed from the relation symbols appearing in the program. Because DL_s facts can be nested, the Herbrand universe of any nontrivial DL_s program is infinite. One could for example represent natural numbers in DL_s using the zero-arity relation *Zero* and the unary relation *Succ*. The Herbrand universe produced by just these two relations, one zero arity and one unary, is inductively infinite.

A *Herbrand Interpretation* of a DL_s program is any subset of its Herbrand universe that is subfact-closed. In other words, if I is a Herbrand Interpretation, then $I = \bigcup \{unroll(f) \mid f \in I\}$. For Datalog, the Herbrand Interpretation is defined similarly, with the difference that subfact-closure is not a requirement for Datalog, as Datalog facts do not contain subfacts.

Given a Herbrand Interpretation I of a DL_s program P , and a rule R in P , we say that R is true in I ($I \models R$) iff for every substitution of variables in R with facts in I , if all the body clauses with those substitutions are in I , so is the head clause of R with the same substitutions of variables.

$$I \models R \text{ iff } \forall \{\overrightarrow{x_i} \rightarrow \overrightarrow{v_i}\} . Body(R) [\overrightarrow{v_i} \setminus \overrightarrow{x_i}] \subseteq I \longrightarrow Head(R) [\overrightarrow{v_i} \setminus \overrightarrow{x_i}] \in I$$

If every rule in P is true in I , then I is a *Herbrand model* for P . The denotation of P is the intersection of all Herbrand models of P . We define $M(P)$ to be the set of all Herbrand models of P , and $D(P)$ to be the denotation of P . We then have $D(P) \triangleq \bigcap_{I \in M(P)} I$. It can be shown that such an

intersection is a Herbrand model itself:

LEMMA 3.2. *The intersection of a set of Herbrand models is also a Herbrand model.*

Unlike Datalog, nontrivial DL_s programs have Herbrand universes that are infinite. Thus, a DL_s program may have only infinite Herbrand models. If a DL_s program has no finite Herbrand models, its denotation is infinite and so no fixed-point may be finitely calculated using the fixed-point semantics. We now relate the operational semantics of DL_s to its model-theoretic semantics.

Equivalence of Model-Theoretic and Fixed-Point Semantics. To show that the model-theoretic and fixed-point semantics of DL_s compute the same Herbrand model, we need to show that the least fixed point of the immediate consequence operator is equal to the intersection of all the Herbrand models for any program. We start by proving the following lemmas (proved in Isabelle; proofs elided for space).

LEMMA 3.3. *Herbrand models of a DL_s program are fixed points of the immediate consequence operator.*

LEMMA 3.4. *Fixed points of the immediate consequence operator of a DL_s program that are subfact-closed are Herbrand models of the program.*

```

⟨toplvl-rule⟩ ::= ⟨rule⟩ | ⟨hclause⟩
⟨rule⟩        ::= [ ⟨hd-item⟩* <- ⟨bd-item⟩* ]
                | [ ⟨bd-item⟩* -> ⟨hd-item⟩* ]
⟨bd-item⟩     ::= ⟨rule⟩ | ⟨bclause⟩
⟨hd-item⟩     ::= ⟨rule⟩ | ⟨hclause⟩
⟨bclause⟩     ::= (⟨tag⟩ ⟨ibclause⟩*)
                | (= ⟨var⟩ (⟨tag⟩ ⟨ibclause⟩*))
⟨hclause⟩     ::= (⟨tag⟩ ⟨ihclause⟩*)
                | (= ⟨var⟩ (⟨tag⟩ ⟨ihclause⟩*))
⟨atom⟩        ::= ⟨var⟩ | ⟨lit⟩
⟨lit⟩         ::= ⟨string⟩ | ⟨number⟩
⟨ihclause⟩    ::= (⟨tag⟩ ⟨ihclause⟩*)
                | ?(⟨tag⟩ ⟨ibclause⟩*)
                | {⟨tag⟩ ⟨ibclause⟩*}
                | [⟨hlist-item⟩*]
                | ?[⟨blist-item⟩*]
                | ⟨atom⟩
⟨ibclause⟩    ::= (⟨tag⟩ ⟨ibclause⟩*)
                | !(⟨tag⟩ ⟨ihclause⟩*)
                | {⟨tag⟩ ⟨ibclause⟩*}
                | [⟨blist-item⟩*]
                | ![⟨hlist-item⟩*]
                | ⟨atom⟩
⟨hlist-item⟩  ::= ⟨ihclause⟩ | ⟨ihclause⟩ ...
⟨blist-item⟩  ::= ⟨ibclause⟩ | ⟨ibclause⟩ ...

```

Fig. 2. The syntax of SLOG. $\langle var \rangle$ is the set of variables, and $\langle tag \rangle$ is the set of relation names. A few syntactic forms, including disjunction, have been elided.

from the ? clauses, the rule may be written without square braces and an arrow to show direction.

Two more rules are needed to define a free-variables analysis. The second of these shows another extension: disjunction in the body of a rule is pulled to the top level and splits the rule into multiple rules. In this case, there is both a rule saying that a free variable in Ef is free in the application and a rule saying that a free variable in Ea is free in the application.

By proving that the Herbrand models and subfact-closed fixed points of the immediate consequence operator are the same, we conclude that the least fixed point of the immediate consequence operator IC_P (a subfact-closed database) is equal to the intersection of all its Herbrand models.

THEOREM 3.5. *The model theoretic semantics and fixed point semantics of DL_s are equivalent.*

Proof sketch: Form the lemma that all Herbrand models are fixed points of IC_P , we conclude that $D(P)$ is a superset of the intersection of all the fixed points. We know that the least fixed point of IC_P (which we'll call LFP_P) is a subset of the intersection of all the fixed points. We therefore have $LFP_P \subseteq D(P)$. From the fact the LFP_P is a Herbrand model, we conclude that $D(P) \subseteq LFP_P$. Putting these facts together, we conclude that $LFP_P = D(P)$.

3.1 Key extensions to the core language

With subfacts, a common idiom becomes for a subfact to appear in the body of a rule, while its surrounding fact and any associated values are meant to appear in the head. For these cases, we use a ? clause, an s-expression marked with a “?” at the front to indicate that although it may appear to be a head clause, it is actually a body clause and the rule does not fire without this fact present to trigger it. The following rule says that if a (**ref** x) AST exists, then x is a free variable with respect to it.

```
(free ?(ref x) x)
```

which desugars to the rule

```
[(= e-id (ref x)) -> (free e-id x)]
```

exposing that the ? clause is an implicit body clause. But if there are no body clauses apart

```

[(/= x y) (free Eb y)
 --> (free ?(lam x Eb) y)]
[(or (free Ef x) (free Ea x))
 --> (free ?(app Ef Ea) x)]

```

Another core mechanism in SLOG is to put head clauses in position where a body clause is expected. Especially because an inner clause can be *responded to* by a fact surrounding it, or by rules producing that fact, being able to emit a fact on-the-way to computing a larger rule is what permits natural-deduction-style rules through a kind of rule splitting, closely related to continuation-passing-style (CPS) conversion [5]. A **!** clause, under a **?** clause or otherwise in the position of a body clause, is a clause that will be deduced as the surrounding rule is evaluated, so long as any **?** clauses are satisfied and any subexpressions are ground (any clauses it depends on have been matched already). These **!** clauses are intermediate head clauses; technically the head clauses of subrules, which they are compiled into internally.

Consider the example in Figure 3, which lets us prove an arithmetic statement like $(\text{plus } (\text{plus } (\text{nat } 1) (\text{nat } 2)) (\text{nat } 1)) \Downarrow 4$. We can construe this rule in a few ways, as written. It could be that both the expression and value should be provided and are proved according to these rules, or it could be treated as a calculator, with the expression provided as input.

$$\begin{array}{c}
 (\text{interp } ?(\text{do-interp } (\text{nat } n)) \ n) \\
 \\
 [(\text{interp } !(\text{do-interp } e_0) \ v_0) \\
 (\text{interp } !(\text{do-interp } e_1) \ v_1) \\
 (+ \ v_0 \ v_1 \ v) \\
 \text{--> ;-----} \ [\text{plus}] \\
 (\text{interp } ?(\text{do-interp } (\text{plus } e_0 \ e_1)) \ v)]
 \end{array}
 \qquad
 \frac{
 \frac{}{(\text{nat } n) \Downarrow n} [\text{nat}]
 }{
 \frac{e_0 \Downarrow v_0 \quad e_1 \Downarrow v_1 \quad v = v_0 + v_1}{(\text{plus } e_0 \ e_1) \Downarrow v} [\text{plus}]
 }$$

Fig. 3. Natural-deduction-style reasoning with **!** clauses in SLOG.

Subclauses, written with parentheses, are treated as top-level clauses whose id column value is unified at the position of the subclause. Another common use for a relation is as a function, or with a designated output column, deterministic or not, so SLOG also supports this type of access via **{}** inner clauses, which have their final-column value unified at the position of the curly-brace subclause. For example, the rule in Figure 3 could also have been written as below, with the clause **{+ v₀ v₁}** in place of variable *v*. This example illustrates that this syntax can also be used for built-in relations like **+**.

Putting this all together and adding SLOG’s built-in list syntax—currently implemented as linked-lists of SLOG facts in the natural way—we can implement rules for appending lists, a naturally direct-recursive task due to a linked list naturally having its first element at its front, so a second list can only be appended to the back of the first list, and the front element onto the front of that.

```

[ (interp ! (do-interp e0) v0)
  (interp ! (do-interp e1) v1)
  --> ;----- [plus]
  (interp ? (do-interp (plus e0 e1))
    {+ v0 v1}) ]

(append ? (do-append [] ls) ls)

[ (append ! (do-append lso lsi) ls')
  -->
  (append ? (do-append [x lso ...] lsi)
    [x ls' ...]) ]

; or ind. case could even be written:
(append ? (do-append [x lso ...] lsi)
  [x
    {append ! (do-append lso lsi)}
    ...])

```

4 APPLICATIONS

In this section, we will examine several related applications of SLOG: implementing reduction systems, natural deduction systems, AAM-based program analyses, and natural-deduction-style type systems.

We start with a λ -calculus interpreter. Let’s observe how β -reduction can be defined via capture-avoiding substitution. If a **do-subst** fact is emitted where a reference to variable *x* is being substituted with expression *E*, associate it in the **subst** relation with *E*:

```
(subst ?(do-subst (ref x) x E) E)
```

However, if x and y are distinct variables, the substitution yields expression $(\text{ref } x)$ unchanged:

```
[(/= x y) --> (subst ?(do-subst (ref x) y E) (ref x))]
```

Recall that $?$ -clauses are body clauses, so these rules could also have been written more verbosely:

```
[(= d (do-subst (ref x) x E)) --> (subst d E)]  
[(/= x y) (= d (do-subst (ref x) y E)) --> (subst d (ref x))]
```

At a lambda, where the formal parameter shadows the variable being substituted, its scope ends and substitution stops:

```
(subst ?(do-subst (lam x Ebody) x E) (lam x Ebody))
```

If the variable does not match and is not free in the (argument) expression E , substitution may continue under the lambda, triggered by a $!$ clause:

```
[(/= x y) ~(free E x)  
--> (subst ?(do-subst (lam x Ebody) y E)  
      (lam x {subst !(do-subst Ebody y E)}))]
```

Three further syntactic extensions are being used in this rule. First off, the negated $\sim(\text{free } E \ x)$ clause in the body requires that the compiler stratify computation of free , as normal when adding otherwise nonmonotonic rule dependance to Datalog programs.

Second, the process of rewriting the lambda body is triggered by the establishment of a **do-subst** fact via a $!$ clause. These $!$ clauses generate facts on-the-fly during rule evaluation, allowing other rules to hook-in by generating a fact to trigger them (using a $!$ clause) and expecting a response in the surrounding body clauses. These $!$ clauses are implemented by generating a subrule whose head clause is the intermediate $!$ clause and whose body contains all body clauses the $!$ clause depends upon, along with and any $?$ clauses in the rule (which are always required to trigger any rule). In this case, term-substitution rules respond to the **do-subst** request via the **subst** relation, as queried here by the $\{\}$ expression in $\{\text{subst } !(\text{do-subst } Ebody \ y \ E)\}$.

Third, this $\{\}$ syntax allows for looking up the final column of a relation by providing all but the final-column value. $(\text{foo } x \ \{\text{bar } y\})$ desugars into $(\text{and } (\text{foo } x \ z) \ (\text{bar } y \ z))$, allowing for the looked-up value to be unified with the position of the $\{\}$ expression in a natural way. Do note that the relation need not actually be functional and could just as easily associate multiple values with any input. $\{\}$ expressions and $!$ clauses are especially expressive when used together in this way for direct recursion.

If we were to desugar the $\{\}$ syntax, $?$ clause, and $!$ clause in this rule, we would obtain two rules. The rule below on the left emits a **do-subst** fact for the body of the lambda, if it qualifies for rewriting, and a **ruleXX-midpoint** fact saving pertinent details of the rule needed in its second half. Below on the right, the second half of the rule requires that the first half of the rule triggered and that the **subst** relation has responded with a rewritten lambda body for the **do-subst** fact do' .

```
[(/= x y) ~(free E x)  
(= do (do-subst (lam x Ebody) y E))      [(ruleXX-midpoint do do' x)  
-->                                     (subst do' Ebody')  
(= do' (do-subst Ebody y E))            -->  
(ruleXX-midpoint do do' x)]             (subst do (lam x Ebody'))]]
```

Finally, in the case of an application, substitution is performed down both subexpressions. If one $!$ clause were nested under the other, they would need to be ordered. In this case, the compiler will detect that both $!$ -clause facts can be emitted in parallel, so this rule will also split into two rules as in the rule above. The first rule will generate both $!$ -clause facts and the second rule will await a response for both. In this way, SLog's semantics for $!$ clauses assists to naturally enable exposure of parallelism in semantics it models.


```
(subst ?(do-subst (app Ef Ea) x E)
      (app {subst !(do-subst Ef x E)}
           {subst !(do-subst Ea x E)}))
```

subexpression to a lambda, substitutes the argument for the formal parameter, and reduces the body.

The compiler will detect in this case that the **do-subst** !-clause fact depends on variable *body*, and the **do-interp** ! clause in the head depends on *body*', but that the first **do-interp** !-clause fact only depends on the variable *fun* from the original ?-clause fact kicking off the rule. These three sequential ! clauses split the rule into four parts during compilation, just as a continuation-passing-style (CPS) transformation [5] would explicitly break a traditional functional implementation of this recursive, substitution-based interpreter into one function entry point and three continuation entry points. Unlike traditional CPS translation of functional programs however, ! clauses in SLOG will naturally emit multiple facts for parallel processing in a nonblocking manner when variable dependence allows for parallelism.

With a substitution function defined, we can define evaluation using a pair of relations: **interp** and **do-interp**. A lambda (**lam** *x* *body*) is already fully reduced. An application reduces its left-hand

```
; values
(interp ?(do-interp (lam x body))
      (lam x body))

; application
[ (interp !(do-interp fun) (lam x body))
  (subst !(do-subst body x arg) body')
  -->
  (interp ?(do-interp (app fun arg))
        {interp !(do-interp body')}) ]
```

4.1 Abstract Machines

Next, instead of using terms alone to represent intermediate points in evaluation, we may wish to explicitly represent facets of evaluation such as the environment, the stack, and the heap. Instead of representing environments through substitution, we may want to represent them explicitly in a higher-order way. As shown in Figure 4, with first-class facts and ad hoc polymorphic rules, we can use defunctionalization to implement first-class relations, providing a global **env-map** relation, we can read with an {**env-map** *env* *x*} expression (assuming ground variables *env* and *x*), along with a (**ext-env** *env* *x* *val*) facility for deriving an extended environment.

```
; environments (defunctionalized)
(env-map ?(ext-env env x val) x val)
[(/= x y) --> (env-map ?(ext-env env x _) y {env-map env y})]
```

Fig. 4. Defunctionalized environments; extension via (**ext-env** *env* *x* *v*), lookup via {**env-map** *env* *x*}.

On the left of Figure 5 shows an abstract machine for CBN evaluation, and on the right, an abstract machine for CBV evaluation. At the top, the rules for reference use {**env-map** *env* *x*} to access the value from the defunctionalized **env-map** relation. In the CBN version, we cannot count on the stored closure to be a lambda closure, so we continue interpretation, using another {} expression to drop-in the transitive reduction of the stored argument closure. Lambda closures are the base case which **interp** as themselves. Finally, application closures trigger a closure to evaluate *Ef* via a ! clause, !(**clo** *Ef* *env*), and the lambda closure that finally results has its body evaluated under its environment, extended with parameter mapped to argument. In the CBN interpreter, (**ext-env** *env*' *x* (**clo** *Ea* *env*)) puts the argument expression *Ea* in the environment, closed with the current environment. In the CBV interpreter, (**ext-env** *env*' *x* *Eav*) puts the argument value *Eav* in the environment (after first evaluating it). In both these interpreters, the **app**-handling rules use ! clauses to implicitly create handling rules and a chain of continuation facts so **interp** maybe be utilized in a direct-recursive manner. The ! syntax introduces a CPS-like transformation that provides a stack in the interpretation of SLOG rules for these CE interpreters to map their stack onto.

```

; ref
(interp ?(clo (ref x) env)
  {interp {env-map env x}})

; lam
(interp ?(clo (lam x Eb) env)
  (clo (lam x Eb) env))

; app
[ (interp !(clo Ef env)
  (clo (lam x Eb) env'))
  (= env' (ext-env env' x (clo Ea env)))
  (interp !(clo Eb env') v)
  -->
  (interp ?(clo (app Ef Ea) env) v) ]

; ref
(interp ?(clo (ref x) env)
  {env-map env x})

; lam
(interp ?(clo (lam x Eb) env)
  (clo (lam x Eb) env))

; app
[ (interp !(clo Ef env)
  (clo (lam x Eb) env'))
  (interp !(clo Ea env) Eav)
  (interp !(clo Eb (ext-env env' x Eav)) v)
  -->
  (interp ?(clo (app Ef Ea) env) v) ]

```

Fig. 5. Two CE (closure-creating) interpreters in SLOC; for CBN eval. (left) and CBV eval. (right).

```

; eval ref
(interp ?(cek (clo (ref x) env) k)
  {interp !(cek env-map env x k)})

; eval lam (apply)
(interp ?(cek (clo (lam x Eb) env)
  [aclo k ...])
  {interp !(cek (clo Eb
    (ext-env env x aclo))
    k)})

; eval app
(interp ?(cek (clo (app Ef Ea) env) k)
  {interp !(cek (clo Ef env)
    [(clo Ea env) k ...])})

; return / halt
(interp ?(cek (clo (lam x Eb) env) [])
  (clo (lam x Eb) env))

; eval ref
(interp ?(cek (clo (ref x) env) k)
  {interp !(cek env-map env x k)})

; eval lam (ret to ar-k)
(interp ?(cek (clo (lam x Eb) env)
  (ar-k aclo k))
  {interp !(cek aclo
    (fn-k (clo (lam x Eb) env)
    k)})})

; eval lam (ret to fn-k)
(interp ?(cek (= aclo (clo (lam _ _) _))
  (fn-k (clo (lam x Eb) env) k))
  {interp !(cek (clo Eb
    (ext-env env x aclo))
    k)})})

; eval app
(interp ?(cek (clo (app Ef Ea) env) k)
  {interp !(cek (clo Ef env)
    (ar-k (clo Ea env) k))})

; return to (halt-k)
(interp ?(cek (clo (lam x Eb) env) (halt-k))
  (clo (lam x Eb) env))

```

Fig. 6. Two CEK (stack-passing) interpreters in SLOC; for CBN eval. (left) and CBV eval. (right).

We can also implement the stack ourselves within our interpreter, thereby eliminating its need for our interpreter itself, by applying a stack-passing transformation. On the left, Figure 6 shows Krivine’s machine [43], a tail-recursive abstract machine for CBN evaluation, and on the right, a tail-recursive abstract machine for CBV evaluation. Each of these machines incrementally constructs and passes a stack. In the CBN stack-passing interpreter, each application reached pushes a closure for the argument expression onto the stack. When a lambda is reached, this continuation is handled by popping its latest closure, the argument value. In the CBV stack-passing interpreter, each application reached pushes an *ar-k* continuation frame on the stack to save the argument value and environment. When a lambda is reached, this continuation is handled by swapping it for a *fn-k* continuation that saves the function value while the argument expression is evaluated (before application). Finally, when a lambda is reached, the *fn-k* continuation is handled by applying the saved closure. Now that the stack is entirely maintained by the interpreter itself, you may note that all recursive uses of `{interp (cek ...)}` are in tail position for the result column of relation `interp`.

4.2 Abstracting Abstract Machines

The *abstracting abstract machines* (AAM) methodology [57, 79] proscribes a particular systematic application of abstract interpretation [19–21] on abstract-machine operational semantics like those we’ve just built in SLOG. AAM proposes key preparatory refactorings of an abstract machine, to remove direct sources of unboundedness through recursion, before more straightforward structural abstraction can be applied. In particular, there are two main sources of unboundedness in the CEK machines: environments and continuations. Environments contain closures which themselves contain environments; continuations are a stack of closures formed inductively in the CBV CEK machine and formed using SLOG’s list syntax in the CBN CEK machine to more closely follow the usual presentation of Krivine’s machine [43]. AAM proposes threading each such fundamental source of unboundedness through a store, added in a normal store-passing transformation of the interpreter that might be used to add direct mutation or other effects to the language. Environments will map variables to addresses in the store, not to closures directly, and the stack will be store allocated at least once per function application so the stack may not grow indefinitely without the store likewise growing without bound. These two changes will permit us to place a bound on the addresses allocated, and thereby finitize the machine’s state space as a whole.

Figure 7 shows the CBV CEK machine of Figure 6 modified in a few key ways, yielding a CESKT machine with control expression, environment, store, continuation, and timestamp/contour components:

- *abstract-machine states have been factored* into **eval**, **apply**, and **ret** configurations; an **eval** state has a control expression, environment (mapping variables to addresses), store (mapping addresses to closures and continuations), current continuation, and timestamp (tracking the size of the store, and thus the next address); an **apply** state has a closure being applied, argument value, store, continuation, and timestamp; and a **ret** state has a value being returned, a store, a continuation, and a timestamp;
- *state transitions have been written as small-step rules* that always terminate; previously, our CEK machines were written to take a big-step from **cek**-state to the final, denoted value as logged in the $(\text{interp } e \ v)$ relation, but in tail-recursive fashion, using $!$ clauses; Figure 7 has no explicit small-step relation, but simply says, for example, that the existence of a **ret** state permits us to deduce to existence of an **apply** state; if we were to want an explicit **step** relation, we could again give this rule a presentation with an implied body via a $?$ clause; for example:

```

; eval ref -> ret
[(eval (ref x) env sto k c)
 -->
 (ret {sto-map sto {env-map env x}} sto k c)]
; eval lam -> ret
[(eval (lam x Eb) env sto k c)
 -->
 (ret (clo (lam x Eb) env) sto k c)]
; eval app -> eval
[(eval (app Ef Ea) env sto k c)
 -->
 (eval Ef env sto (ar-k Ea env k) c)]
; ret to kaddr -> ret
[(ret vf sto (kaddr c') c)
 -->
 (ret vf sto {sto-map sto (kaddr c')} c)]
; ret to ar-k -> eval
[(ret vf sto (ar-k Ea env k) c)
 -->
 (eval Ea env sto (fn-k vf k) c)]
; ret to fn-k -> apply
[(ret va sto (fn-k vf k) c)
 -->
 (apply vf va sto k c)]
; apply -> eval
[(apply (clo (lam x Eb) env) va sto k c)
 -->
 (eval Eb
  (ext-env env x (addr c))
  (ext-sto (ext-sto sto (kaddr c) k)
    (addr c) va)
  k
  {+ 1 c})])

```

Fig. 7. A CESKT (control, environment, store, continuation, timestamp) interpreter in SLOG.

```
(step ?(ret va sto (fn-k vf k) c)
      (apply vf va sto k c))
```

- *states have been subjected to a store-passing transformation* which has added a store `sto` and timestamp (stored-value count) `c` to each state; environments now bind variables to addresses and the current store binds those addresses to values; we perform a variable lookup with `{sto-map sto {env-map env x}}`; at an `apply` state, we use the store count `c` to generate a fresh address (`addr c`) for the parameter `x`; we also store-allocate the current continuation at a continuation address (`kaddr c`), in preparation for modeling the stack finitely as well; when returning to a (`kaddr c`), the continuation is simply fetched from the store as in the fourth rule down (`ret to kaddr`).

From here it suffices to pick a finite set from which to draw addresses. To instantiate a monovariant control-flow analysis from this CESKT interpreter, it would be enough to use the variable name itself as the address or to generate an address (`addr x`). When the environment and store become finite, so does the number of possible states. Consider what happens, as the naturally relational `sto-map` relation encoding stores conflates multiple values at a single address for the same variable. Conflation in the store would lead naturally to nondeterminism in any `step` relation. When looking up a variable, two distinct `ret` states could result, leading to two distinct `apply` states after some further steps.

A (potentially) more precise, though (potentially) more costly analysis would be to specialize all control-flow points and store points by a finite history of recent or enclosing calls. Such a *k-call-sensitive* analysis can be instantiated using a specific instrumentation and allocation policy, as can many others [28]. It requires an instrumentation to track a history of *k* enclosing calls, and then an *abstract allocation policy* that specializes variables by this call history at binding time. Such context-sensitive techniques are a gambit that the distinction drawn between variable *x* when bound at one call-site vs another will prove meaningful—in that it may correlate with its distinct values. Increasing the polyvariance allows for greater precision while also increasing the upper-bound on analysis cost. In a well known paradox of programming analyses, greater precision sometimes goes hand-in-hand with lower cost in practice because values that are simpler and fewer are simpler to represent [82]. At the same time, we use the polyvariant entry point of each function, its body and abstract contour—(`kaddr Eb c'`)—to store allocate continuations as suggested by previous literature on selecting this address [31] so as to adapt to the value polyvariance chosen.

The per-state store by itself is a source of exponential blowup for any polyvariant control-flow analysis [56]. Instead, it is standard to use a global store and compute the least-upper-bound of all per-state stores. In SLOG this is as simple as using a single global (`store addr val`) relation instead of a defunctionalized (`sto-map sto addr val`) relation that approximates all per-state stores in one. The left side of Figure 8 shows a version of the CESKT machine with a global store and a tunable instrumentation that can be varied by changing the `tick` function rule. Currently, `tick` instantiates this to a 3-*k*-CFA: at each function application, the current call site (now saved in the `ar-k` and `fn-k` continuation frames to provide to the `apply` state) is saved in front of the current call history and the fourth-oldest call is dropped.

This is the classic *k*-CFA, except perhaps that the original *k*-CFA, formulated for CPS as it was, also tracked returns positively instead of reverting the timestamp as functions return like we do here. The original *k*-CFA used true higher-order environments, unlike equivalent analyses written for object oriented languages which implicitly had flat environments (objects) [58]. The corresponding CFA for functional languages is called *m*-CFA and is shown on the right side of Figure 8. *m*-CFA has only the latest call history as a flat context. Instead of having a per-variable address with a per-variable history tracked by a per-state environment, *m*-CFA stores a variable *x* at abstract contour *c* (i.e., abstract timestamp, instrumentation, 3-limited call-history) in the store

```

;; Eval states
[(eval (ref x) env k _)
-->
 (ret {store {env-map env x}} k)]
[(eval (lam x body) env k _)
-->
 (ret (clo (lam x body) env) k)]
[(eval (app ef ea) env k c)
-->
 (eval ef env
  (ar-k ea env (app ef ea) c k)
  c)]

;; Ret states
[(ret vf (ar-k ea env call c k))
-->
 (eval ea env (fn-k vf call c k) c)]
[(ret va (fn-k vf call c k))
-->
 (apply call vf va k c)]
[(ret v (kaddr e env))
 (store (kaddr e env) k)
-->
 (ret v k)]

;; Apply states
[(apply call (clo (lam x Eb) env) va k c)
-->
 (eval Eb env' (kaddr Eb env') c')
 (store (kaddr Eb env') k)
 (store (addr x c') va)
 (= env' (ext-env env x (addr x c'))))
 (= c' {tick !(do-tick call c)})]

;; tick (tuning for 3-k-CFA)
(tick ?(do-tick call [h0 h1 _])
 [call h0 h1])

;; Eval states
[(eval (ref x) k c)
-->
 (ret {store (addr x c)} k)]
[(eval (lam x body) k c)
-->
 (ret (clo (lam x body) c) k)]
[(eval (app ef ea) k c)
-->
 (eval ef (ar-k ea (app ef ea) c k) c)]

;; Ret states
[(ret vf (ar-k ea call c k))
-->
 (eval ea (fn-k vf call c k) c)]
[(ret va (fn-k vf call c k))
-->
 (apply call vf va k c)]
[(ret v (kaddr e c))
 (store (kaddr e c) k)
-->
 (ret v k)]

;; Apply states
[(apply call (clo (lam x Eb) _) va k c)
-->
 (eval Eb (kaddr Eb c') c')
 (store (kaddr Eb c') k)
 (store (addr x c') va)
 (= c' {tick !(do-tick call c)})]

;; Propagate free vars
[(free y (lam x body))
 (apply call (clo (lam x body) clam) _ _ c)
-->
 (store (addr y {tick !(do-tick call c)})
 {store (addr y clam)})]

```

Fig. 8. An AAM for global-store k -CFA (left) and m -CFA (right) in SLOG. These are evaluated in Section 6.

at the address (`addr x c`). This means at every update to the current flat context c , now taking the place of the environment, all free variables must be propagated into an address (`addr x c`).

4.3 Type Systems

Along with operational semantics and program analyses, SLOG naturally enables the realization of structural type systems based on constructive logics [61, 62]. These systems are often specified via an inductively-defined typing judgment, whose derivations may be represented in SLOG via (sub)facts and whose typing rules may be realized as rules in SLOG (providing their evaluation may be operationalized via SLOG's idioms). For example, the rules for the simply-typed λ -calculus (STLC) in Figure 9 define the judgment $\Gamma \vdash e : \tau$ —under typing environment Γ , e has been proven to have type τ . Proofs of this judgment are represented via SLOG facts of the form $(: (ck \Gamma e) \tau)$; each rule in the type system is then mirrored by a corresponding rule in SLOG.

When implementing a type system in SLOG, it is crucial to consider some important differences between SLOG and natural deduction per-se. First, equivalence in SLOG is intensional, via fact interning (as in type theories such as Coq's Calculus of Inductive Constructions). For example, while our type checker for STLC decides $\Gamma \vdash e : \tau$ via structural recursion on e , there are infinitely

$\text{STLC Terms } e ::= \begin{array}{l} (\lambda(x:\tau) e) \\ (e_0 e_1) \\ x \end{array}$		$\text{STLC Types } \tau \in T ::= \begin{array}{l} \tau \rightarrow \tau \\ \text{nat} \\ \dots \end{array}$	
<div style="border: 1px solid black; padding: 2px; display: inline-block;">$\Gamma \vdash e : \tau$</div>			
T-VAR	$\frac{x:T \in \Gamma}{\Gamma \vdash x:T}$	$[\text{-->;} \text{----- T-Var} \\ (: ?(\text{ck } \Gamma \text{ (ref } x)) \{ \text{env-map } \Gamma \ x \})]$	
T-ABS	$\frac{\Gamma, x:T_1 \vdash e : T_2}{(\lambda (x:T_1) e) : T_1 \rightarrow T_2}$	$[(: !(\text{ck } (\text{ext-env } \Gamma \ x \ T_1) \ e) \ T_2) \\ \text{-->;} \text{----- T-Abs} \\ (: ?(\text{ck } \Gamma \ (\lambda \ x \ T_1 \ e)) \ (-> \ T_1 \ T_2))]$	
T-APP	$\frac{\Gamma \vdash e_0 : T_0 \rightarrow T_1 \quad e_1 : T_0}{\Gamma \vdash (e_0 \ e_1) : T_1}$	$[(: !(\text{ck } \Gamma \ e_0) \ (-> \ T_0 \ T_1)) \\ (: !(\text{ck } \Gamma \ e_1) \ T_0) \\ \text{-->;} \text{----- T-App} \\ (: ?(\text{ck } \Gamma \ (\text{app } e_0 \ e_1)) \ T_1)]$	

Fig. 9. Syntax (top) and semantics (left) of STLC; equivalent SLOG (right).

many $\Gamma' \supseteq \Gamma$ for which $\Gamma' \vdash e : \tau$ also holds—materializing these (infinite) Γ' would result in nontermination.

Here, we focus on the presentation of algorithmic (i.e., syntax-directed) type checking procedures. The decidability of these systems follows immediately from their structurally-recursive nature, a property inherited by their SLOG counterparts. We expect enumerating terms in other theories which enjoy strong normalization will readily follow. We anticipate SLOG’s declarative style may also be a natural fit for type synthesis, by bounding (potentially infinite) rewritings using a decreasing “fuel” parameter. However, we leave this, along with explorations of other (bidirectional, substructural, etc...) type systems in SLOG to future work.

Simply-typed λ -calculus. We begin with the simply-typed λ -calculus [8, 61]. The syntax of STLC terms and types is shown at the top of Figure 9. Our presentation roughly follows Chapter 9 of Benjamin Pierce’s *Types and Programming Languages* [61]; we use Scheme-style syntax to more closely mirror the SLOG presentation. STLC extends the untyped λ -calculus: λ -abstractions are annotated with types, and variable typing defers to a typing environment which assigns types to type variables and is subsequently extended at callsites. STLC defines a notion of “simple” types including (always) arrow types between simple types and (sometimes) base types (e.g., nat), depending on the presentation.

The key creative challenge in translating a type system into SLOG lies in the operationalization of its control flow. For example, consider the T-VAR rule on the left side of Figure 9: the rule is parametric over the typing environment Γ , however a terminating analysis necessarily inspects only a finite number of typing environments. The solution is interpret the rules in a demand-driven way, using subfacts to defer type checking until necessary. Operationally, each type rule is predicated upon a message $?(\text{ck } \Gamma \ e)$, which triggers type synthesis for e under the typing environment Γ . Using this technique, we may invoke the analysis by including a fact $(\text{ck } \Gamma \ e)$. Using stratified negation, we may treat $(: (\text{ck } \Gamma \ e) \ \tau)$ as a decision procedure, like so:

```
[(success Γ e τ) <-- (typecheck Γ e τ) ( : ! (ck Γ e) τ )]
[(failure Γ e τ) <-- ~(success Γ e τ)]
```

Here, the inclusion of $(\text{typecheck } \Gamma \ e \ \tau)$ forces the materialization of $!(\text{ck } \Gamma \ e)$, and subsequently the enumeration of the type of each of its subexpressions. Once the resulting type is

materialized, we force its unification with τ (implicitly using SLOG's intensional notion of equality) and, when successful, generate (**success** $\Gamma \vdash \tau$). The resulting SLOG implementation necessarily terminates because the analysis enumerates at most a finite number of (**ck** $\Gamma \vdash e$) facts (because of the structural recursion on e done by \vdash), which in-turn forces materialization of a finite number of (**ext-env** $\Gamma \times \tau$) facts, each of which forces a bounded materialization of **env-map**.

Natural Deduction and Per Martin-Löf's Type Theory. The well-known Curry-Howard isomorphism relates terms in pure functional languages to proofs in an appropriate constructive logic [22]. For STLC, the Curry-Howard isomorphism tells us that we may read our type checker as a decision procedure for intuitionistic propositional logic. As an alternative (but equivalent) perspective, we now consider how SLOG may represent proofs in Per Martin-Löf's intuitionistic type theory (ITT) [53].

By design, ITT cleanly separates propositions from their associated derivations (proof objects). In SLOG, we may represent derivations as structured facts, obtaining the nested structure of derivations via SLOG's subfacts. Adopting this perspective, checking a natural-deduction proof in ITT involves propagating assumptions to their usages (akin to the propagation achieved using maps in STLC).

$$\wedge I \quad \frac{A \text{ true} \quad B \text{ true}}{A \wedge B \text{ true}} \quad \left| \begin{array}{l} [(\text{true} \mid (\text{ck } A\text{-pf } A)) \\ (\text{true} \mid (\text{ck } B\text{-pf } B)) \\ \text{-->} \\ (\text{true} \mid ?(\text{ck } (\wedge I \text{ A-pf } B\text{-pf } (\wedge I \text{ A } B)) (\wedge I \text{ A } B)))] \end{array} \right.$$

Fig. 10. The introduction rule for \wedge in ITT.

Figure 10 shows the introduction form for \wedge (left) and its corresponding SLOG transliteration (right). Checking a derivation of $\wedge I$ forces the checking of each sub-derivation, and (upon success) populates the **true** relation with the appropriate derivation.

Implication in ITT is managed by introducing, and then discharging, assumptions: $A \supset B$ holds whenever B may be proven by assuming A . Figure 11 details the implication rule in ITT (left) and SLOG (right): the introduced hypothesis (named u in $\supset I$) is subsequently discharged to produce an assumption-free proof of $A \supset B \text{ true}$. In SLOG, the rule for $\supset I$ introduces an assumption by forcing the materialization of (**assuming** $A \text{ pf-B}$)—other rules then “push down” the assumption $A \text{ true}$ to their eventual uses (top right of Figure 11), performing transitive closure of assumptions to their usages in an on-demand fashion.

$$\supset I \quad \frac{\begin{array}{c} \overline{A \text{ true}}^u \\ \vdots \\ B \text{ true} \end{array}}{A \supset B \text{ true}} \quad \left| \begin{array}{l} ;; \text{ Propagate assumptions} \\ (\text{true} \mid ?(\text{ck } (\text{assuming } P (\text{assumption } P)) P) P) \\ \dots \\ [(\text{true} \mid (\text{ck } (\text{assuming } A \text{ pf-B } B)) \\ \text{-->} \\ (\text{true} \mid ?(\text{ck } (\supset I (\supset A B) \text{ pf-B } (\supset A B)))] \end{array} \right.$$

Fig. 11. Implication in ITT and SLOG

First-Order Dependent Types: λLF . The Edinburgh Logical Framework (λLF) is a dependently-typed λ -calculus [36]. It is a first-order dependent type system, in the sense that it stratifies its objects into kinds, types (families), and terms (values)—kinds may quantify over types, but not over other kinds. The syntax of λLF is detailed at the bottom of Figure 12—it extends STLC with kinds, which are either $*$ or (type families) $\Pi x : T.K$, where T is a simple type (of kind $*$). λLF generalizes the arrow type to the dependent product: $\Pi x : T.T$. System λLF enjoys several decidability properties

<i>λLF Contexts</i>	$\Gamma ::= \emptyset$ $\Gamma, x : T$ $\Gamma, x :: K$	<i>λLF Types</i>	$\tau \in T ::= X$ $\Pi x : T. T$ $(T e)$
<i>λLF Kinds</i>	$K ::= \Pi x : T. K \mid *$		
TA-Abs	$\frac{\Gamma \vdash S :: * \quad \Gamma, x : S \vdash t : T}{\Gamma \vdash (\lambda(x : S)t) : \Pi x : S. T}$		<pre> [(: ! (ck-k Γ S) (*)) (: ! (ck-t (ext-env Γ x S) t) T) -->;----- (: ? (ck-t Γ (λ x S t)) (Π x S T))] </pre>
TA-App	$\frac{\Gamma \vdash t_1 : \Pi x : S_1. T \quad \Gamma \vdash t_2 : S_2 \quad \Gamma \vdash S_1 \equiv S_2}{\Gamma \vdash (t_1 t_2) : [x \mapsto t_2] T}$		<pre> [(: ! (ck-t Γ t1) (Π x S1 T)) (: ! (ck-t Γ t2) S2) (true ! (== Γ S1 S2)) -->;----- (: ? (ck-t Γ (app t1 t2)) {subst ! (do-subst T x t2)}))] </pre>
KA-App	$\frac{\Gamma \vdash S :: \Pi x : T_1. K \quad \Gamma \vdash t : T_2 \quad \Gamma \vdash T_1 \equiv T_2}{\Gamma \vdash (S t) : [x \mapsto t] K}$		<pre> [(: ! (ch-k Γ S) (Π x T1 K)) (: ! (ck-t Γ t) T2) (true ! (== Γ T1 T2)) -->;----- (: ? (ch-t Γ (type-app S t)) {subst ! (do-subst K x t)}))] </pre>
	<pre> [(->wh ! (do->wh t1) t1') -->;----- (->wh ? (do->wh (app t1 t2)) (app t1' t2))] </pre>		<pre> (->wh ? (do->wh (app (λ x T1 t1) t2)) {subst ! (do-subst t1 x t2)}) </pre>

Fig. 12. *λLF*: Syntax (top), selected rules (left) and SLOG (right).

which make it particularly amenable to implementation in SLOG. The first is strong normalization, which implies that reduction sequences for well-typed terms in our implementation will be finite. The second is *λLF*'s focus on canonical forms and hereditary substitution [37]. In *λLF*, terms are canonicalized to weak-head normal form (WHNF); this choice enables inductive reasoning on these canonical forms, and this methodology forms the basis for Twelf [60].

The *judgments-as-types* principle interprets type checking for *λLF* as proving theorems in intuitionistic predicate logic; Using this principle, we may define traditional constructive connectives (such as \wedge , \vee , and \exists) via type families and their associated rules. For example, including in Γ a binding $\wedge \mapsto \Pi P : \text{prop}. \Pi Q : \text{prop}. *$ allows using the constructor \wedge , though \wedge must be instantiated with a suitable P and Q , which must necessarily be of some sort (e.g., *prop*) also bound in Γ .

We have performed a transliteration of *λLF* as formalized in Chapter 2 of *Advanced Topics in Types and Programming Languages (ATAPL)* [62]. Our transliteration (from pages 57–58) consists of roughly 150 lines of SLOG code. Figure 12 details several of the key rules. TA-Abs introduces a Π type, generalizing the T-Abs rule in Figure 9. The TA-App applies a term t_1 , of a dependent product type $\Pi x : S_1. T$, whenever the input t_2 shares an equivalent type, S_2 . The notion of equality here is worth mentioning: \equiv demands reduction of its arguments to WHNF—*λLF* is constructed to identify terms under WHNF, thus ensuring \equiv will terminate as long as the term is typeable. Reduction to WHNF is readily implemented in SLOG; two exemplary rules detailed at the bottom of Figure 12 outline the key invariant in WHNF: reduce down the leftmost spine, eliminating β -redexes via application. Equality checks are demanded by the TA-App and KA-App rules, and force normalization of their arguments to WHNF before comparison of canonical forms, generating a witness in **true** before triggering the head of the rule.

5 IMPLEMENTATION

We have implemented SLOG in a combination of Racket (the compiler, roughly 10,600 lines), C++ (the runtime system and parallel RA backend; roughly 8,500 lines), Python (a REPL and daemon; roughly 2,500 lines) and Slog (60 lines for list-splicing support). In this section, we describe relevant particulars.

5.1 Compiler

Our Racket-based compiler translates Slog source code to C++ code that links against our parallel RA backend. Our compiler is structured in the nanopass style, composed of multiple passes which consume and produce various forms of increasingly-low-level intermediate representations (IRs) [41]. After parsing, *organize-pass* performs various simplifications, such as canonicalizing the direction of \rightarrow and splitting disjunctive body clauses and conjunctive head clauses into multiple rules. This pass also eliminates various syntactic niceties of the language, including $!$ and $?$ clauses, nested rules, list syntax, and splicing variables. When a program includes splicing variables, the compiler concatenates a small splicing support library. Finally, this pass performs static unification, syntactically identifying clauses and variables that are statically constrained to be equal.

SLOG’s distribution paradigm is built upon binary joins. Thus, after organization, *partitioning-pass* breaks down bodies with multiple clauses into sequences of binary joins. Partitioning represents an algorithmic challenge, as there may be many ways to partition a set of clauses into sequences of binary joins. For example, a rule such as $[H \leftarrow B_1 B_2 \dots B_n]$ may be converted into an $(n - 1)$ -length sequence of binary joins (first joining B_1 and B_2 to form an intermediate relation which is subsequently joined with B_3) or a tree of binary joins (joining each B_i and B_{i+1} into intermediate relations which are then joined). Unfortunately, optimal partitioning is undecidable in general, and our compiler relies upon a set of heuristics and practical optimizations we’ve found to work well in practice, along with enabling the user to manually suggest a partitioning by using a $--$ syntax operator. Our early implementations preferred to form trees of joins—based on the intuition that this would extract more parallel work—however, we found this often resulted in materializing large numbers of unnecessary facts (essentially forming large Cartesian products to later be filtered in subsequent joins). We have come to see effective partitioning as a fundamental part of writing high-performance code in SLOG, similar to Soufflé’s Sideways Information Passing Strategy [73].

After partitioning, *split-selections-pass* performs index selection by inspecting each rule in the program and calculating a set of indices. There are two important differences between SLOG and a typical shared-memory Datalog implementation. First, because of our distribution methodology, it is impossible to organize indices in a trie-shaped representation that allows overlap-based index compression [71]. Second, fact interning is slightly tricky in the presence of multiple indices: we must be careful to avoid a fact being assigned two distinct intern keys during the same iteration in separate indices. Our solution is to designate a special *canonical index* which is used for intern key origination, along with a set of special administrative rules which replicate, for each relation, the intern key to every non-canonical index.

The last two passes are strongly-connected component (SCC) construction and incrementalization. Datalog programs are typically stratified into a plan of SCCs for evaluation. This aids efficiency (a single-node implementation may ignore considering rules unnecessary to the current SCC, our distributed implementation evaluates SCCs using task-level parallelism) and is also semantically relevant in the presence of stratified negation (to ensure all negated relations are computed strictly-before negation runs). Finally, incrementalization (i.e., semi-naïve evaluation) transforms the program to use a worklist-based evaluation strategy, every relation appearing in a rule body is split into two versions—**delta** and **total**. Rules are rewritten to add facts to **delta**,

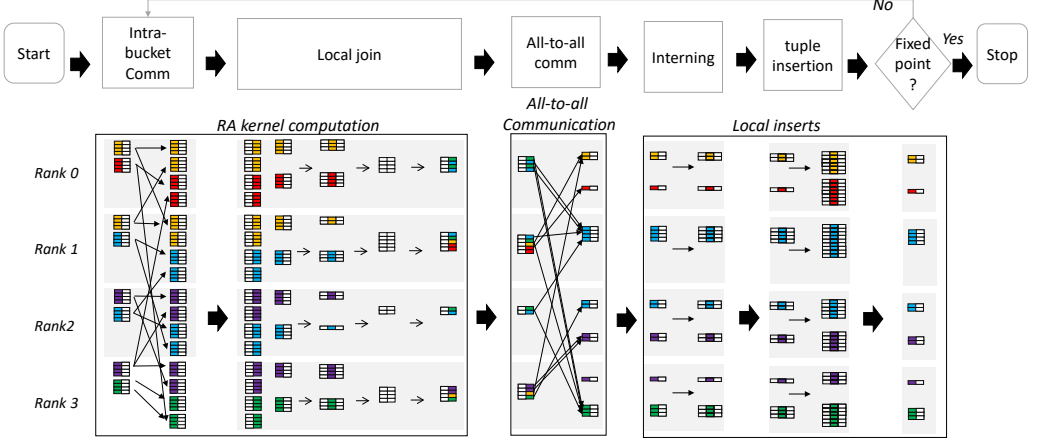


Fig. 13. An illustration of the main phases of our parallel RA backend.

while bodies are triggered by new entries in **delta**; our backend merges **delta** into **total** at the end of each iteration.

5.2 Backend

Our parallel relational-algebra backend supports fixed-point iterations and is designed for large-scale multi-node HPC clusters. Based on the bulk-synchronous-processing protocol and built using the MPI-everywhere model [26, 84], the parallel RA framework addresses the problem of partitioning and balancing workloads across processes by using a two-layered distribution hash-table [46]. In order to materialize newly generated facts within each iteration, and thus facilitate *iterated* RA (in a fixed-point loop), an all-to-all data exchange phase is used at every iteration. Figure 13 shows a schematic diagram of all the phases (including the interning phase) in the context of an incrementalized TC computation. There are three primary phases in the backend: (1) RA kernel computation, (2) all-to-all communication and (3) local insertion.

Figure 13 shows a schematic diagram of all the phases (including the interning phase) in the context of an incrementalized TC computation. There are three primary phases of our system: local RA computation, all-to-all data exchange, and materialization in appropriate indices. Workload (relations) are partitioned across processes using the the double hashing approach, which also adds an extra intra-bucket communication phase, to co-locate matching tuples. During the local computation phase, RA kernels (comprising of join, projection, union and others) are executed in parallel across all processes. Output tuples generated from local computation may each belong to an arbitrary bucket in the output relation, so an MPI all-to-all communication phase shuffles the output of all joins to their managing processes (preparing them for any subsequent rules or iterations). Once new tuples are received, we perform the interning phase, that first checks if the received fact was already populated before, if not then a new intern id is created and associated with the fact. The fact is then inserted into newt, and following the semi-naive evaluation approach, the newly generated facts forms the input for the following iteration of the fixed point. This process continues till the fixed-point is reached and no new facts can be discovered.

RA kernel computation. The two-layered distributed approach, with local hash-based joins and hash-based distribution of relations, is a foundational method to distribute RA-kernel (primarily join) operations over many nodes in a networked cluster computer. This algorithm involves partitioning

relations by their join-column values so that they can be efficiently distributed to participating processes [78]. The main insight behind this approach is that for each tuple in the outer relation, all relevant tuples in the inner relation must be hashed to the same MPI process or node, permitting joins to be performed locally on each process.

A major challenge with parallel workload partitioning is to ensure that every process gets to work on similar sized workloads that gives a load balanced system. A major challenge to enforce this load balance is to deal with inherently imbalanced data coming from key-skewed relations. To ensure uniform load across processes, we have built on previous approaches [46, 47] by developing strategies that mitigate load-imbalance in a dynamic manner. The approach [46] uses a two-layered distributed hash-table to partition tuples over a fixed set of *buckets*, and, within each bucket, to a dynamic set of *subbuckets* which may vary across buckets. Each tuple is assigned to a bucket based on a hash of its key-column values, but within each bucket tuples are hashed on non-join-column values, assigning them to a local subbucket, then mapped to an MPI process. Within subbuckets, tuples are stored in B-trees, organized by key-column values. Our scheme permits buckets that have more tuples to be split across multiple processes, but requires some additional communication among subbuckets for any particular bucket. We have developed and evaluated a dynamic refinement strategy, to decide how many subbuckets to allocate per-bucket. To distribute subbuckets to managing processes, we use a round-robin mapping scheme which we found to be significantly more effective than hashing.

A join operation can only be performed for two *co-located relations*: two relations each keyed on their respective join columns that share a bucket decomposition (but not necessarily a subbucket decomposition for each bucket). This ensures that the join operation may be performed separately on each bucket as all matching tuples will share a logical bucket; it does not, however, ensure that all two matching tuples will share the same subbucket as tuples are assigned to subbuckets (within a bucket) based on the values of non-join columns, separately for each relation. The first step in a join operation is therefore an *intra-bucket communication* (see Figure 13) phase within each bucket so that every subbucket receives all tuples for the outer relation across all subbuckets (while the inner relation only needs tuples belonging to the local subbucket). Following this, a *local join* operation (with any necessary projection and renaming) is performed in every subbucket.

All-to-all communication. To enable iterated parallel RA (in a fixed-point loop), processes must engage in a non-uniform all-to-all inter-process shuffle of generated tuples to their position in an output index. This data exchange is performed to *materialize* the output tuples generated from the local compute phase (where RA kernels are executed) to their appropriate processes (based on their bucket-subbucket assignment). Materializing a tuple in an output relation (resulting from an RA operation) involves hashing on its join and non-join columns to find its bucket and sub-bucket (respectively), and then transmitting it to the process that maintains that bucket/sub-bucket. As tuples generated from the local compute phase may each belong to an arbitrary bucket/sub-bucket in the output relation, an all-to-all communication phase is required to shuffle the output tuples to their managing processes. Given variability in the number of tuples generated across processes, and in their destination processes (due to inherent imbalance), the communication phase in our framework is *non-uniform* in nature. The output tuples may each belong to an arbitrary bucket in the output relation, an MPI *all-to-all* communication phase shuffles the output of all joins to their managing processes (preparing them for any subsequent iteration).

The overall scalability of the RA backend relies on the scalability of the all-to-all inter-process data exchange phase. However, all-to-all is notoriously difficult to scale [45, 67, 74]—largely because of the *quadratic* nature of its workload. We address this scaling issue by adopting recent advancements [24] that optimizes non-uniform all-to-all communication by extending the log-time Bruck algorithm [14,

74, 77] for non-uniform all-to-all workloads. Traditional algorithms to implement non-uniform all-to-all communication takes linear iterations as every process must send and receive from every other process. Bruck algorithm however, sends more amount of data in logarithmic steps, and therefore significantly improves overall performance of all-to-all data exchange. Using the Bruck implementation of non-uniform all-to-all algorithm was instrumental in the overall scalability of the backend.

Local inserts. After all-to-all data exchange, every process receives a new set of facts that must be materialized to be used as input in the subsequent iteration of the fixed-point loop. Local insert is a two-step process involving interning and inserting newly generated facts in the appropriate version of a relation (delta and total). Interning assigns a unique 64-bit key to every fact, and in order to scale this process, it must be performed in an embarrassingly parallel manner without the need for any synchronization among processes. This is done by reserving the first 16 bits of the key for unique buckets ids, and the remaining 48-bits for facts. Since, a canonical-index (master index) bucket is never split across a process, reserving 16 bits for bucket ids ensures that globally unique intern keys can be created concurrently across processes. The fact-id component of the intern key is created by a bump pointer, which ensures that locally all facts receive a unique key. After interning, facts are added to their appropriate versions of the relations (delta or total), and a check is performed to see if a fixed-point has been reached. This check is performed by a global operation that checks the size of all relations across all processes, and if all sizes remains unchanged across two iterations, then this indicates that fixed-point has been attained and the program terminates, otherwise a new iteration is initiated.

6 EVALUATION

We aimed to measure and evaluate SLOG’s improved indexing and data parallelism, using three sets of performance benchmarks (PBs):

- PB1** (Section 6.1) How does SLOG compare against other systems designed for performance and parallelism on traditional Datalog workloads (without ADTs): Soufflé and RadLog?
- PB2** (Section 6.2) How do SLOG subfacts perform against Soufflé ADTs in the context of the m -CFA and k -CFA benchmarks developed in Section 4.
- PB3** (Section 6.3) How well can SLOG scale to many threads on a supercomputer?

We evaluated **PB1** and **PB2** by running a set of experiments on large cloud machines from Amazon AWS and Microsoft Azure. For **PB1**, we ran a set of strong scaling experiments of transitive closure on large graphs, picking transitive closure as an exemplary problem to measure end-to-end throughput of deductive inference at scale. For **PB2**, we measure the performance of the implementation of our k and m -CFA analyses from Section 4 compared to an equivalent implementation in Soufflé using abstract datatypes (ADTs). We answer **PB3** by running experiments on the Theta supercomputer at Argonne National Supercomputing Lab, scaling a control-flow analysis for the λ -calculus to 1000 threads on Argonne’s Theta.

6.1 Transitive Closure

We sought to compare SLOG’s full-system throughput on vanilla Datalog against two comparable production systems: Soufflé and Radlog. Soufflé is engineered to achieve the best-known performance on unified-memory architectures, and supports parallelism via OpenMP. Radlog is a Hadoop-based successor to the BigDatalog deductive inference system, which uses Apache Spark to perform distributed joins at scale [32]. We originally sought to compare SLOG directly against BigDatalog, but found it does not support recent versions of either Spark or Java (being built to

Table 1. Single-node TC Experiments

Name	Graph Properties		System	Time (s) at Process Count			
	Edges	TC		15	30	60	120
FB-MEDIA	206k	96,652,228	SLOG	62	40	21	18
			Soufflé	35	33	34	37
			Radlog	254	295	340	164
RING10000	10k	100,020,001	SLOG	363	218	177	115
			Soufflé	149	143	140	141
			Radlog	464	646	852	1292
SUITESPARSE	412k	3,354,219,810	SLOG	–	1,593	908	671
			Soufflé	1,417	1,349	1,306	1,282
			Radlog	–	–	–	–

target Java 1.5). Under direction of BigDatalog’s authors, we instead used Radlog, which is currently under active development and runs on current versions of Apache Spark.

We performed comparisons on an Standard_M128s instance rented from Microsoft Azure [55]. The node used in our experiments has 64 physical cores (128 threads) running Intel Xeon processors with a base clock speed of 2.7GHz and 2,048GB of RAM. To directly compare SLOG, Soufflé, and Radlog, we ran each on the same machine using 15, 30, 60, and 120 threads. We ran SLOG using OpenMPI version 4.1.1 and controlled core counts via `mpirun`. We compiled Soufflé from its Git repository, using Soufflé’s compiled mode to compile each benchmark separately to use the requisite number of threads before execution. Radlog runs natively on Apache Spark, which subsequently runs on Hadoop. To achieve a fair comparison against Soufflé and SLOG, we ran Radlog using Apache Spark configured in local mode; Spark’s local mode circumvents the network stack and runs the application directly in the JVM. We used three large graphs shown the first column of Table 1: FB-MEDIA is media-related pages on Facebook, RING10000 is ring graph of 10,000 nodes, and SUITESPARSE is from the UF Sparse Matrix Collection [23]. We configured Radlog according the directions on its website, experimenting with a variety of partitions (used for shuffling data between phases) to achieve the best performance we could. Ultimately, we used three times as many partitions as available threads, except for RING10000, for which we found higher partition counts caused significantly lower performance.

Table 1 details the results of our single-node performance comparisons in seconds for each thread count, where each datapoint represents the best of three runs (lower is better). Experiments were cut off after 30 minutes. In every case, we found that SLOG produced the best performance overall at 120 threads, even compared to Soufflé’s best time. However, as expected, our results indicate that Soufflé outperforms SLOG at lower core counts (below 60). Soufflé implements joins with tight loops in C++, and (coupled with its superior single-node datastructures) this allows Soufflé to achieve better performance than either SLOG or Radlog at lower core counts. We found that Radlog did not scale nearly as well as either Soufflé or SLOG. We expected this would be the case: both SLOG and Soufflé compile to C++. By comparison, Radlog’s Spark-based architecture incurs significant sequential overhead due to the fact that it is implemented on top of the JVM and pays a per-iteration penalty by using Hadoop’s aggregation and shuffling phase. SLOG also incurs sequential overhead compared to Soufflé due to its distributing results after every iteration, though results indicate that our MPI-based implementation helps ameliorate this compared to Radlog.

Table 2. Control-Flow Analysis Experimental Results: Slog vs. Soufflé

	Term Sz.	Iters	Cf. Pts	Sto. Sz.	8 Processes		64 Processes	
					Slog	Soufflé	Slog	Soufflé
3- <i>k</i> -CFA	8	1,193	98,114	23,413	00:01	01:07	0:02	00:15
	9	1,312	371,010	79,861	00:02	14:47	0:03	02:56
	10	1,431	1,441,090	291,317	00:06	☹	0:05	45:49
	11	1,550	5,678,402	1,107,957	00:27	☹	0:16	☹
	12	1,669	22,541,634	4,315,125	02:14	☹	1:07	☹
	13	1,788	89,822,530	17,022,965	12:17	☹	5:08	☹
4- <i>k</i> -CFA	9	1,363	311,790	65,397	00:01	14:38	00:03	02:08
	10	1,482	1,197,038	229,621	00:05	☹	00:05	40:30
	11	1,601	4,687,854	853,493	00:20	☹	00:13	☹
	12	1,720	18,550,766	3,281,909	01:40	☹	00:53	☹
	13	1,839	73,801,710	12,859,381	08:44	☹	03:58	☹
	14	1,958	294,404,078	50,892,789	60:53	☹	35:46	☹
5- <i>k</i> -CFA	9	1,429	203,674	50,677	00:02	05:30	0:03	001:15
	10	1,548	756,890	167,285	00:04	65:20	0:04	015:08
	11	1,667	2,911,898	597,493	00:13	☹	0:08	196:06
	12	1,786	11,416,218	2,245,109	00:56	☹	0:27	☹
	13	1,905	45,202,074	8,687,605	04:38	☹	2:00	☹
	14	2,024	179,882,650	34,158,581	25:14	☹	9:58	☹
10- <i>m</i> -CFA	50	6,120	21,005	656,847	00:02	00:02	00:10	00:01
	100	11,670	42,855	2,781,447	00:07	00:09	00:20	00:04
	200	22,770	86,555	11,440,647	00:26	00:56	00:42	00:23
	400	44,970	173,955	46,399,047	01:44	06:26	01:38	01:56
	800	89,370	348,755	186,875,847	07:35	45:22	04:21	09:33
	1600	178,170	698,355	750,069,447	32:56	☹	14:36	62:35
12- <i>m</i> -CFA	25	3,559	17,105	385,279	00:01	<0:01	0:06	<0:01
	50	6,434	36,280	1,885,179	00:04	000:03	0:11	00:03
	100	12,184	74,630	8,284,354	00:16	000:24	0:23	00:10
	200	23,684	151,330	34,680,204	01:10	002:37	0:53	00:55
	400	46,684	304,730	141,861,904	05:04	018:39	2:23	04:12
	800	92,684	611,530	573,785,304	22:46	2:38:22	7:28	24:58
15- <i>m</i> -CFA	12	2,211	14,461	136,740	00:01	<0:01	0:04	<0:01
	24	3,591	35,857	1,443,058	00:03	00:02	0:06	00:01
	48	6,351	78,649	8,292,250	00:14	00:15	0:14	00:07
	96	11,871	164,233	38,931,946	01:08	01:41	0:36	00:36
	192	22,911	335,401	167,976,586	05:15	12:10	1:49	02:51
	384	44,991	677,737	697,126,858	24:10	92:35	6:45	16:30

6.2 AAMs and CFAs

Next, we sought to benchmark the analyses described in Section 4, at scale, versus an equivalent implementation using ADTs in Soufflé (we ignore Radlog in this comparison due to its lack of

support for ADTs). We developed a SLOG analysis for each of six different polyvariance choices: three k -CFA ($k=3,4,5$) and three m -CFA ($m=10,12,15$) implementations. We then systematically derived six different Soufflé-based variants. We tested each of these on six different term sizes, drawn from a family of worst-case terms identified in David Van Horn’s thesis [38]. We then benchmark both SLOG and Soufflé on each of these instances and report on their results, scalability, and broad trends which we observed. Our Soufflé code is an exact port of the SLOG code we used (see Figure 7), except that $\$$ -ADT values are used in place of subfacts and the analysis was designed in the first place to avoid the need for these subfacts to trigger rule evaluation as they can in SLOG.

Experimental Setup. The experiments described in this subsection were run on a `c6a.metal` instance rented from Amazon Web Services (AWS), including 192 hardware threads (when run using the `.metal` instance types) and 384 GiB of RAM. Because both SLOG and Soufflé are designed to enable parallelism, we ran each experiment at two distinct scales: 8 and 64 processes (threads). SLOG was invoked using `mpirun`, and Soufflé’s compiled mode was used to produce a binary which was subsequently run and timed using `GNU time`. We did not systematically measure memory usage; recent microbenchmarks for TC report 3 – 5 \times memory blowup versus Soufflé. We record and report the best of three runs for each experiment (imposing a four hour cutoff). To avoid an unfair comparison to Soufflé with respect to on-disc ADT materialization (which may explode due to linearization of linked data), our Soufflé implementation does not output control-flow points or store directly—instead we measure and report their size using the `sum aggregate` (built in to Soufflé).

Results. We report our results in Table 2. Each of six distinct analysis choices is shown along the left side. Along rows of the table, we show experiments for a specific combination of analysis, precision, and term size. We detail the total number of iterations taken by the SLOG backend, along with control-flow points, store size, and runtime at both 8 and 64 threads for SLOG and Soufflé. Times are reported in minutes / seconds form; several runs of Soufflé took under 1 second (which we mark with `<0:01`), and `⌚` indicates that the run timed out after four hours.

Inspecting our results, we observed several broad trends. First, as problem size increases, SLOG’s runtime grows less-rapidly than Soufflé’s. This point may be observed by inspecting runtimes for a specific set of experiments. For example, 10- m -CFA with term size 200 took SLOG 26 seconds, while Soufflé’s run took 56 seconds. Doubling the term size to 400 takes 104 seconds in SLOG, but 398 seconds in Soufflé—a slowdown of 4 \times in SLOG, compared to a slowdown of 7 \times in Soufflé. A similar trend happens in many other experiments, e.g., 15 minutes to over three hours for Soufflé (13 \times slowdown) vs. 2 to 4 seconds (2 \times slowdown) in SLOG’s runtime on 5- k -CFA. Inspecting the output of Soufflé’s compiled C++ code for each experiment helped us identify the source of the slowdown. For example, the rule for returning a value to a continuation address `$KAddr(e, env)`, in Figure 8, must join a return state using this address with an entry in the continuation store for this address. Because Soufflé does not index subfacts, a scan-then-filter approach is used. In this case, as the subfact is an exact match, it could be optimized but in cases where a single variable in a subfact matches another, Soufflé’s scan-and-filter implementation cannot be avoided.

This return rule and its compiled C++ code is shown in Figure 14. Note that it uses two nested for loops to iterate over the entire `ref` relation, then iterate over the entire `kont_map` relation, and finally check if there is a match for all possible combinations. In this way, Soufflé’s lack of indices for structured values leaves it no other option but to materialize (in time) the entire Cartesian product as it checks for matches—rather than efficiently looking up relevant tuples in an appropriate index as would normally be the case for top-level relations.

For a fixed problem size, we found that Soufflé and SLOG both scaled fairly well. Soufflé consistently performed well on small input sizes; additional processes did not incur slowdowns, and Soufflé’s

```

// ret(av, k) :- ret(av, $KAddr(e, env)), kont_map($KAddr(e, env), k).
//      env0[0] ---^ env2[0]-^ ^--- env2[1]      env3[1] -----^

if(!(rel_13_delta_ret->empty()) && !(rel_18_kont_map->empty())) {
  for(const auto& env0 : *rel_13_delta_ret) {
    RamDomain const ref = env0[1];
    if (ref == 0) continue;
    const RamDomain *env1 = recordTable.unpack(ref,2);
    {
      if( (ramBitCast<RamDomain>(env1[0]) == ramBitCast<RamDomain>(RamSigned(3)))) {
        RamDomain const ref = env1[1];
        if (ref == 0) continue;
        const RamDomain *env2 = recordTable.unpack(ref,2);
        {
          for(const auto& env3 : *rel_18_kont_map) {
            // On this line we've omitted bitcasts and Tuple ctors:
            if( !(rel_19_delta_kont_map->contains({{ env2[0], env2[1] }, env3[1] })))
              && !(rel_12_ret->contains({{env0[0], env3[1]} }))) {
              // Omitted: null checks and insertion of {{ env0[0], env3[1] }} into ret
            }
          }
        }
      }
    }
  }
}

```

Fig. 14. Example C++ code generated by Soufflé.

efficiency was generally reasonable (roughly 50%) when algorithmic scalability did not incur slowdowns. For example, in 3- k -CFA ($n=8$), Soufflé took 67 seconds at 8 processes, and 15 seconds at 64 processes. SLOG’s parallelism doesn’t outweigh communication overhead on smaller problems, particularly on problems with high iteration count and low per-iteration work. As problem size increases, our SLOG implementations show healthy scalability; efficiency grows as problem size grows (e.g., 24:10 to 6:45 on 15- m -CFA/384, 22:46 to 7:28 on 12- m -CFA/800).

We found that extracting optimal efficiency from SLOG was facilitated by increasing per-iteration work and avoiding long sequences of sequential work with comparatively lower throughput. Because each iteration represents a synchronization point, greater numbers of iterations will decrease opportunities for parallelism. As an example of this principle, our 10- m -CFA uses a flat `ctx` fact to represent the context; a previous version used a linked list 10 elements deep, however this design achieved poorer scaling efficiency due to these 10-iteration-long sequences of work necessary to extend the instrumentation at each call-site. In our experiments scaling efficiency improved as polyvariance increased; e.g., improving by 2× for 10- m -CFA, but 3.5× for 15- m -CFA. We believe this is because of the relatively higher per-iteration work available with increased polyvariance.

6.3 Multi-node Scaling Experiments

In recent years, several cloud providers have launched MPI-capable HPC nodes for their cloud services and significantly upgraded their network interconnects [1, 16]. Alongside related advances in cloud architectures and MPI implementations, this move signals the ability for massively parallel analytics to be deployed as a service in the near future. In this spirit, we also conduct some preliminary strong-scaling experiments on the Theta supercomputer. For this we used a fully monomorphized m -CFA (distinct from the m -CFA used in the previous subsection)—experiments we had ready to go when our allocation on Theta became possible. The Theta Supercomputer [59] at the Argonne Leadership Computing Facility (ALCF) of the Argonne National Laboratory is a Cray machine with a peak performance of 11.69 petaflops. It is based on the second-generation Intel Xeon Phi processor and is made up of 281,088 compute cores. It has an aggregate of 843.264 TiB of DDR4 RAM, 70.272 TiB of MCDRAM and 10 PiB of online disk storage. The supercomputer has a Dragonfly network topology uses the Lustre filesystem.

We ran three sets of m -CFA worst-case experiments with 110 terms and $m = 6, 7$ and 8 . The results of the three sets of experiments, referred to as `dvh-110-6`, `dvh-110-7` and `dvh-110-8` are plotted in Figure 15. For all three set of experiments we observe improvement in performance with increase in process counts, until maximum efficiency is attained, after which performance degrades with increasing process counts, due to communication overhead and workload starvation. In general, for a given workload (problem size), we observe a range of processes that exhibit healthy scalability. `dvh-110-6` shows a near 100% scaling efficiency ($2\times$ speedup while increasing the process count from 100 to 200), performance however drops when the number of processes is increased to 300. Similarly, `dvh-110-7` shows a 75% scaling efficiency ($3\times$ speedup when the process count is increased from 100 to 400), and `dvh-110-8` shows a 71% scaling efficiency ($1.42\times$ speedup going 400 to 800).

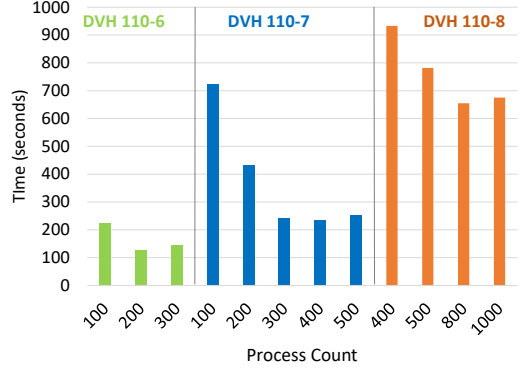


Fig. 15. Strong-scaling experiments on Theta.

7 RELATED AND FUTURE WORK

Distributed Datalog. There have been significant implementation efforts to scale Datalog-like languages to large clusters of machines. For example, BigDatalog [69], Distributed Socialite [68], Myria [35], and Radlog [33] all run on Apache Spark clusters (servers networked together via commodity switches within a datacenter). Extending Spark’s architecture with recursive queries (and aggregates), these frameworks scale to large datasets typical of Spark queries. SLOG differs from these systems in two primary ways. First, compared to SLOG’s MPI-based implementation, Apache Spark’s framework-imposed overhead is increasingly understood to be a bottleneck in scalable data analytics applications, with several authors noting order-of-magnitude improvements when switching from Spark to MPI [3, 44, 64]. Second, none of the aforementioned systems support first-class subfacts; for example, while Radlog can compute the length of the shortest path from a specific point, it cannot materialize the path per-se. Recently, Radlog’s authors have created DCDatalog, a parallel Datalog which targets shared-memory SMP architectures and demonstrate a $10\times$ runtime speedup compared to Soufflé on a machine with four eight-core processors and 256GB of RAM. Unfortunately, DCDatalog is not open-source, and we have not been able to obtain a copy for evaluation; we believe it is difficult to interpret DCDatalog’s results compared to SLOG and Soufflé, as their paper notes “Soufflé does not allow aggregates in recursion, and thus it must use a stratified query that results in very poor performance” for several evaluation queries. Last, Nexus (also closed-source) has claimed a significant performance boost (up to $4\times$) compared to BigDatalog by using Apache Flink, a data-flow processing language [39].

Datalog Extensions. Noting the first-order nature of vanilla Datalog—and often inspired by Datalog’s efficient semi-naïve evaluation strategy—there has been extensive work in extending Datalog with additional expressive power [6, 7, 9, 50, 52]. Flix augments Datalog with lattices [50, 52], but is not specifically focused on efficient compilation; recently, Ascent is a macro-based implementation of Datalog in Rust which includes lattices and shows orders-of-magnitude runtime improvements versus Flix [65]. Similarly, Datafun is a pure functional language which computes fixed points of monotone maps on semilattices [6, 7]. Compared to SLOG, Datafun’s evaluation strategy is top-down and based on the λ -calculus; the authors have recently studied semi-naïve

evaluation of Datafun upon work on the incremental λ -calculus [17, 27]. SLOG's primary difference from this work is that it is based on DL_s rather than the λ -calculus; because of this, semi-naïve evaluation for functions in SLOG (using defunctionalization) requires no extra logic.

Datalog + Constraints. An increasingly-popular semantic extension to Datalog is adding first-class constraints [9, 51, 75, 76]. These constraints typically allow interfacing with an SMT solver, potentially in a loop with subsequent analysis [9]. Formulog includes ADTs and first-order functions over ADTs, allowing Turing-equivalent to build formulas of arbitrary size to be checked by Z3 [9]; we anticipate SLOG will perform well compared to Formulog when subfact-indexing is of concern, though by Amdahl's law this effect will be smaller in code whose computation is dominated by calls to Z3. Similarly, Rosette efficiently compiles solver-aided queries to efficient implementations using host language constructs and a symbolic virtual machine (SVM) [75, 76]. SLOG is largely orthogonal to these systems, which focus on shared-memory implementations and are not primarily concerned with parallelization. We have transliterated proof-of-concept examples from both of these projects into SLOG, but it is currently impossible to call Z3 from SLOG as doing so would require all facts be resident on a single node. Semantically, SLOG is more directly comparable to constrained HornSAT or existential fixed-point logic, which have attracted recent interest for their application to program verification [10, 11, 25, 34]. DL_s can express constrained HornSAT problems as long as a decision procedure for the background logic is available; we plan to study usage of SLOG for CHCs in subsequent work.

Parallel Program Analyses. Given the algorithmic complexity intrinsic to large-scale program analyses, there has been significant interest in its parallelization [2, 4, 12, 15, 70, 83] or implementation using special-purpose datastructures [42, 49, 54, 63, 81]. There are a variety of fundamental approaches to scalability; for example, summarization-based analyses (such as Saturn [2, 83]) are attractive due to the task-level parallelism they expose. Much work in scaling program analysis has focused on context-insensitive analyses—wherein task-level parallelism is more directly exploitable. The goals of SLOG are most closely related to current efforts on scaling rich, whole-program context-sensitive analyses using deductive inference [4, 40, 66].

Parallel Relational Algebra. SLOG's backend builds upon recent successes in balanced, parallel relational algebra (BPRA) and follow-up work on compilation of vanilla Datalog to parallel relational algebra kernels [29, 30, 48]. However, that work focuses mainly on the low-level implementation of relational algebra kernels rather than a unified programming language, compiler, and runtime.

8 CONCLUSION

In this work we extended Datalog with subfacts, explicating a core language, DL_s , which supports subfacts as first-class facts, values, and threads. This straightforward semantic extension enables several key innovations, including the expression of higher-order functions via defunctionalization, subfact indexing, and a rich connection to constructive logic via the proofs-as-programs interpretation. We have implemented these ideas in SLOG, a fully featured language for data-parallel structured deduction, using a new MPI-based relational algebra backend, and we demonstrate its application to operational semantics, abstract interpreters, and formal systems broadly. Our experiments show that SLOG is competitive with, or (when programming with structured heap data) orders-of-magnitude faster than state-of-the-art systems, showing improved data-parallelism and scaling up to hundreds of cores on large unified machines and high-performance clusters.

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