# DECONSTRUCTING DATALOG 

by

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#### Abstract

The deductive query language Datalog has found a wide array of uses, including static analysis (Smaragdakis and Bravenboer, 2010), business analytics (Aref et al., 2015), and distributed programming (Alvaro et al., 2010, 2011). Datalog is high-level and declarative, but simple and well-studied enough to admit efficient implementation strategies. For example, Whaley et al. found they could replace a hand-tuned C implementation of context-sensitive pointer analysis with a comparably-performing Datalog program that was 100x smaller (Whaley and Lam, 2004; Whaley et al., 2005).

However, Datalog's semantics are not stable under extensions. For instance, adding arithmetic operations breaks Datalog's termination guarantee. Despite this, nearly all practical implementations extend Datalog beyond its theoretical core to add niceties such as arithmetic, datatypes, aggregations, and so on. Moreover, pure Datalog cannot abstract over repeated code: one may express a static analysis over a particular program, but to express the same analysis over multiple programs, one must duplicate the analysis code for each program analyzed.

This thesis deconstructs Datalog from a categorical and type theoretic perspective to determine what makes it tick. Datalog's semantic guarantees are provided by brute syntactic restrictions, such as stratification and the absence of function symbols. In place of these, we find compositional semantic properties such as monotonicity, which we capture using types. We show that this permits integrating Datalog's features with those of typed functional languages, such as algebraic data types and higher order functions.

In particular, this thesis makes the following contributions: 1. We define and expound the semantics and metatheory of Datafun, a pure and total higher-order typed functional language capturing the essence of Datalog. Where Datalog has predicates defined by a restricted class of Horn clauses, Datafun has finite sets and set comprehensions; Datalog's bottom-up recursive queries become iterative fixed points; and Datalog's stratification condition becomes a matter of tracking monotonicity with types. 2. We show how to generalize seminaïve evaluation to handle higher-order functions. Seminaïve evaluation is a technique from the Datalog literature which improves the performance of Datalog's most distinctive feature: recursive queries. These are computed iteratively, and under a naïve evaluation strategy, each iteration recomputes all previous values. Seminaïve evaluation computes a safe approximation of the difference between iterations. This can asymptotically improve the performance of Datalog queries. Seminaïve evaluation is defined partly as a program transformation and partly as a modified iteration strategy, and takes advantage of the first-order nature of Datalog. We extend this transformation to handle higher-order programs written in Datafun. 3. In the process of generalizing seminaïve evaluation, we uncover a theory of incremental, monotone, higher-order computation, in which values change over time by growing larger, and programs respond incrementally to these increases.


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## Chapter 1

## Introduction

### 1.1 Monotone fixed points

A remarkable number of computational problems can be expressed as finding the least fixed point of a monotone map on a semilattice satisfying the ascending chain condition. The utility of Datalog is explained by the fact that it captures this pattern, albeit restricted to the semilattice of finite sets under union. To understand this pattern better, let's consider three examples of increasing complexity: (1) reachability in a graph; (2) single-source shortest paths; and (3) analyzing which variable assignments may reach a given line in a simple imperative program (called "reaching definitions").

Reachability Consider a graph and suppose we wish to find all nodes reachable from some designated start node. We proceed as follows: first, we put a check mark next to the start node; then, repeatedly, we pick a node and put a check next to it if any of its neighbors is checked. Once there are no nodes which we can mark this way - in particular, when there are no edges between checked and unchecked nodes - we are done; the reachable nodes are exactly the checked nodes.

Shortest paths Now suppose each edge $e$ in the graph has an associated non-negative length $\mathrm{d}_{e}$, and we wish to find the minimum distance to each reachable node. We use a small modification of the previous procedure: instead of a check mark, we annotate nodes $v$ with the length $\mathrm{d}_{v}$ of the shortest path to them we've discovered so far. Initially we mark the start node with 0 and every other node with $\infty$ (representing "no known path"). Then, whenever an edge provides a shorter path to a node, we update its annotation that is, for any edge $v \xrightarrow{e} u$ we may update $d_{u}:=\min \left(d_{u}, d_{v}+d_{e}\right)$. Once no shortening edges exist, the annotations $d_{v}$ cannot change, and we are done. ${ }^{1}$

Reaching definitions Finally, let's consider something seemingly completely different: statically analyzing a simple imperative program (figure 1.1). In particular, we wish to determine which assignments may reach a given program line; for example, the print on line 2 will receive only the value of $x$ assigned on line 1 , while the print on line 4 may receive the values assigned on both lines 1 and 5 .

[^0]```
\(x\) := 0
print \(x\)
while true do
    print x
    x := x + 1
```

FIGURE I.I Example program

We determine this by propagating information along the control flow graph of our program. At each line we maintain a set of assignments (line-number/variable pairs) that we know can reach that line. Each line collects the assignments from all lines that can transfer control to it - usually the immediately preceding line, but loops and conditionals complicate this. However, lines which assign to a variable add themselves to this set, and discard other assignments to the same variable from incoming lines.

Once there is no line whose corresponding assignment-set is changing, the analysis is finished. See figure 1.2 for a step-by-step diagram of this process.

How do these three examples fit into our proposed pattern: finding the least fixed point of a monotone map on a semilattice satisfying the ascending chain condition? Let's break down each point in turn:

Fixed points In each example, we maintained some state - check marks or distance annotations on nodes, sets of reaching assignments on lines - that changed over time, and we terminated when there was no action we could take - no node, edge, or line we could examine - which would change this state. In other words, we halted once our state was fixed under our transition function.

Monotone Although our state changed over time, it did not change in arbitrary ways: there was a direction to it. Nodes went from unchecked to checked; distances to nodes decreased; and sets of reaching assignments grew. We can formalize this by giving our states a partial order representing the direction they change as computation progresses and information increases - an order with respect to which our state increases monotonically over time.

For example, in graph reachability a node's state is a boolean flag; we'll regard it as true if the node is checked, false otherwise. Since nodes go from unchecked to checked but not vice-versa, we say that false $<$ true. In single-source shortest-path our distances change toward zero, so we order them inversely, $\infty<\ldots<3<2<1<0$. And in reaching definitions our assignment-sets change by gaining new elements, so we order them by the subset relation, $s \leqslant t \Longleftrightarrow s \subseteq t$.

Semilattices In each example, we had some way of combining information from multiple sources. In graph reachability, we marked a node if any of its neighboring nodes were marked; in shortest paths, when there were multiple edges/paths into a node, we took the minimum among these competing options; and in reaching definitions, when a line could receive control from multiple lines, we took the union of their reaching assignment sets.

Not coincidentally, these operations - boolean disjunction, minimum, and union respectively - are the least upper bound operators for the partial orders we imposed on

| reaching <br> assignments | 1 | $x:=0$ |
| :--- | :--- | :--- | | reaching |
| :--- |
| assignments |

STEP 1 The assignment on line 1 adds itself.
STEP 2 Information propagates to line 2.

| 1 | $x:=0$ | $(x, 1)$ |
| :--- | :--- | :--- |
| 2 | print $x$ | $(x, 1)$ |
| 3 | while true do | $(x, 1)$ |
| 4 | print $x$ |  |
| 5 | $x:=x+1$ |  |


| 1 | $\mathrm{x}:=0$ | $(\mathrm{x}, 1)$ |
| :--- | :--- | :--- |
| 2 | print x | $(\mathrm{x}, 1)$ |
| 3 | while true do | $(\mathrm{x}, 1)$ |
| 4 | print x | $(\mathrm{x}, 1)$ |
| 5 | $\mathrm{x}:=\mathrm{x}+1$ |  |

STEP 3 Information propagates to line 3.
STEP 4 Information propagates to line 4.

| $1 \mathrm{x}:=0$ | $(\mathrm{x}, 1)$ | 1 | $\mathrm{x}:=0$ | $(\mathrm{x}, 1)$ |
| :--- | :--- | :--- | :--- | :--- |
| 2 | print x | $(\mathrm{x}, 1)$ | 2 | print x |
| 3 | while true do | $(\mathrm{x}, 1)$ | 3 while true do | $(\mathrm{x}, 1)$ |
| 4 | $\mathrm{print} \mathrm{x})$ | $(\mathrm{x}, 5)$ |  |  |
| 5 | $\mathrm{x}:=\mathrm{x}+1$ | $(\mathrm{x}, 1)$ | 4 | print x |
| $(\mathrm{x}, 5)$ | 5 | $\mathrm{x}:=\mathrm{x}+1$ | $(\mathrm{x}, 5)$ |  |

STEP 5 The assignment on line 5 adds itself, discarding the assignment from line 1.

STEP 6 Information propagates from line 5 to line 3 because of the loop.

| 1 | $x:=0$ | $(x, 1)$ |
| :--- | :---: | :--- |
| 2 | print $x$ | $(x, 1)$ |
| 3 | while true do | $(x, 1) \quad(x, 5)$ |
| 4 | print $x$ | $(x, 1) \quad(x, 5)$ |
| 5 | $x:=x+1$ | $(x, 5)$ |

STEP 7 Information propagates to line 4.

| $1 \mathrm{x}:=0$ | $(x, 1)$ |  |
| :--- | :---: | :--- |
| 2 | print $x$ | $(x, 1)$ |
| 3 | while true do | $(x, 1)(x, 5)$ |
| 4 | print $x$ | $(x, 1)(x, 5)$ |
| 5 | $x:=x+1$ | $(x, 5)$ |

STEP 8 No changes-done!
our states, making those partial orders into join-semilattices. In general, we write $x \vee y$ for the least upper bound/semilattice join of $x$ and $y$. If our partial order represents the direction of increasing information, $x \vee y$ is a natural way to combine information: it includes all information from (is greater than) both $x$ and $y$, but does not jump to unnecessary conclusions - it is the least, most conservative, upper bound.

Ascending chain condition Finally, in each case there was a limit on how much information we could possibly learn, and thus how many transitions we could take. For instance, in graph reachability, there were finitely many nodes, and each node could only transition from unchecked to checked once. This argument can be formalized by showing our partial order on states obeys the ascending chain condition (ACC), which asserts that there are no infinite strictly ascending chains, $x_{0}<x_{1}<x_{2}<\ldots$; consequently, any process producing an increasing state-sequence must halt. We leave it as an exercise for the reader to convince themselves each of our state-posets satisfies this property. ${ }^{2}$

The essence of our computational pattern, then, is this: we follow rules which accumulate information monotonically until there is nothing left to learn (a fixed point). We usually need a way to combine information from multiple sources (our semilattice); and if we are to finish, we cannot keep learning forever (the ascending chain condition). As this thesis progresses, we will see more examples of this pattern and develop a more precise mathematical understanding of what it consists in. To do this, it will help to examine a language which is at once an instance of this pattern for us to deconstruct and a vehicle for expressing further instances of it: Datalog.

### 1.2 Datalog

Datalog may be seen either as a restricted logic programming language or an expressive database query language. To start with, we consider the former view, explaining Datalog in terms of deduction. Here is a simple Datalog program:

```
parent(alice, bob).
parent(bob, charlie).
grandparent (X, Z) \leftarrow parent (X, Y), parent(Y, Z).
```

We can see each line, or clause, of this program as an inference rule. The first two lines are axioms, or inference rules with no premises; the last line is a rule with two premises. In inference rule notation we might write this:

$$
\overline{\text { parent(alice, bob })} \quad \overline{\text { parent(bob, charlie })} \quad \frac{\operatorname{parent}(\mathrm{X}, \mathrm{Y}) \quad \operatorname{parent}(\mathrm{Y}, \mathrm{Z})}{\operatorname{grandparent}(\mathrm{X}, \mathrm{Z})}
$$

More formally, a Datalog program is a sequence of clauses terminated by periods. Each clause is an implication with one conclusion and an optional list of premises, written conclusion-first, "B $\leftarrow A_{1}, A_{2}, \ldots, A_{n}$."; or if there are no premises, simply "B.". The premises and conclusion

[^1]are atoms of first-order logic: a predicate applied to a sequence of terms, $\mathrm{P}\left(\mathrm{T}_{1}, \ldots, \mathrm{~T}_{n}\right)$, or a negation of the same, $\neg \mathrm{P}\left(\mathrm{T}_{1}, \ldots, \mathrm{~T}_{\mathrm{n}}\right)$. Moreover, the conclusion of a clause must be positive, not negated.

As is the convention when interpreting inference rules, in Datalog the variables in a clause (for which we use capital letters $X, Y, Z$ ) are considered to be universally quantified: the logical interpretation of our third line, for example, is $(\forall X, Y, Z) \operatorname{parent}(X, Y) \wedge \operatorname{parent}(Y, Z) \Longrightarrow$ grandparent $(X, Z)$.

The intended interpretation of a Datalog program is the set of all facts deducible from its clauses. We access this set via queries such as grandparent(alice, $X$ ), which asks for a list of all $X$ such that grandparent (alice, $X$ ) is deducible. In general we allow conjunctive queries: lists of conjoined atoms, for instance the unlikely query parent $(X, Y)$, parent $(Y, X)$, which asks for a pair of people each the parent of the other (or, in the case $X=Y$, a single person who is their own parent). Finally, a query without any variables, such as grandparent(charlie, alice), amounts to asking a yes-or-no question: is the query deducible or not?

Computing the set of all deducible facts fits neatly into our description of monotone fixed points: applying inference rules is a monotone process, adding but never removing knowledge; we desire the fixed point of this process, where everything that can be deduced has been; our semilattice is sets of atoms under union. The ascending chain condition, however, does not obviously hold - perhaps there are infinitely many deducible facts? To answer this question, we must consider what differentiates Datalog from other logic programming languages.

### 1.2.1 Termination and recursion

Thus far our description of Datalog has not distinguished it from its ancestor Prolog; this is because the difference lies not so much in their syntax as in their semantics. Without further restrictions, whether a proposition is deducible from a collection of clauses is in general undecidable. Prolog's solution is to specify its proof search strategy. This lets Prolog programmers reason about the execution of their programs, but it can mean that some logically sensible recursive programs fail to terminate. For example:

```
reachable}(\textrm{Y})\leftarrow\mathrm{ reachable (X), edge( }\textrm{X},\textrm{Y})
reachable(start).
```

These rules encode our graph reachability example: a node is reachable if one of its neighbors is, or if it is the start node. In Prolog, however, any query to reachable will loop.

This is because Prolog uses backward chaining (also called goal-directed or top-down) depth-first search: we start from a goal and reason backward, applying rules that might prove it. These rules are applied in the order they occur in the program, so to solve the query reachable(st-louis), Prolog will apply the first rule and try recursively to solve reachable ( X ) for unknown $X$. This in turn will apply the same rule, solving reachable $\left(X_{2}\right)$ for unknown $X_{2}$, which will solve reachable $\left(X_{3}\right)$ for unknown $X_{3}$, and so on and on interminably. ${ }^{3}$
${ }^{3}$ One natural approach to this problem is to keep backward chaining, but use a complete search strategy instead of depth-first search; this is the approach adopted by miniKanren (Friedman et al., 2005). This restores some declarativeness to logic programming; in particular, reordering rules can no longer cause unproductive infinite looping. However, introducing a "redundant" rule like reachable $(X) \leftarrow$ reachable $(X)$ will still cause proof search to continue indefinitely (although it won't prevent any proofs from being found). And while this example is

Datalog takes a different tack: rather than fix a proof search strategy, it imposes limitations that keep proof search decidable. In particular, ignoring for now the issue of negation, it imposes two restrictions which keep all relations finite:

1. Clauses are range-restricted: all variables in the conclusion of a clause must occur positively in its premises. For example, the premiseless clause "equal $(X, X)$." is disallowed; while logically sensible (it asserts equal is reflexive), it leaves the variable $X$ unconstrained, which would generate an infinite relation.
2. Programs are constructor-free: predicate arguments are either atomic terms or variables. This prevents the introduction of new terms that don't already appear in the program, as this could also result in an infinite relation; for example, the relation containing all digit-lists (the use of a 'constructor' is underlined and red):
```
digits(nil).
digits(cons(X,Xs))}\leftarrow\operatorname{digit}(X),\operatorname{digits}(Xs)
```

Range-restriction and constructor-freedom together ensure that relations are finite and thus enforce the ascending chain condition. This permits the most common Datalog implementation strategy, forward chaining. In backward chaining we start from a goal ("can we reach St. Louis?") and reason backward, applying rules that might prove it. In forward chaining, we start from what we know ("we can reach Chicago, and there's an edge from Chicago to St. Louis") and apply rules whose premises are satisfied.

The weakness of forward chaining is that it's undirected: it deduces everything it can! If all you want to know is whether you can reach St. Louis, this is wasteful. On the other hand, it's much easier to know when to stop: when there is no rule whose application yields a new fact. This is the operational justification for range-restriction and constructor-freedom: by ensuring all predicates are finite, we guarantee forward-chaining deduction terminates.

### 1.2.2 Stratified negation

Negation and deduction have an interesting relationship. Consider the following program:

$$
\text { undeducible() } \leftarrow \neg \text { undeducible(). }
$$

In classical logic, an implication $B \leftarrow A$ is equivalent to $B \vee \neg A$. Applying this, the above is equivalent to undeducible() $\vee \neg \neg$ undeducible(), and thus simply to undeducible(). Regarded as a rule of inference, however - as a strategy for deducing new facts from ones already known - this clause makes little sense: we cannot invoke it unless we have proved its conclusion is false!

To avoid this sort of gap between the logical meaning of a program and its interpretation as inference rules, Datalog allows only programs where uses of negation can be stratified: a recursively defined predicate (or mutually recursive group of predicates) cannot use its own negation in its definition.

[^2]This restriction also avoids the need to make arbitrary choices. For example, this is disallowed:

```
marry-rochester() \leftarrow\negmarry-st-john().
marry-st-john() \leftarrow\negmarry-rochester().
```

This is classically equivalent to marry-rochester () $\vee$ marry-st-john(). While sensible logically, this means answering simple yes-or-no queries requires making an arbitrary choice: if we query the propositions marry-rochester() and marry-st-john() we may consistently answer either Rochester or St John or both. The symmetry of our program makes answering either Rochester or St John unprincipled, and both is simply not deducible from the given rules.

Thus, unlike the preceding restrictions, stratified negation is not about finiteness or decidability; rather, it is motivated by interpreting a program as a set of rules for deduction and not merely a set of propositions. In other words, in Datalog truth is identified with deducibility.

Following this principle, we regard anything not deducible as false. This conforms with the programmer's expectation that anything not explicitly declared to be true is false. For example, returning to our graph reachability example:

```
reachable(Y) \leftarrow reachable (X), edge (X, Y).
reachable(start).
```

Regarded as mere propositions, these do not rule out the possibility that reachable $(X)$ is true for all vertices $X$, regardless of the edge relation (in other words, one model of these propositions makes reachable the entire vertex set). But this is clearly not the programmer's intent, which is to capture reachability: not every graph is completely connected!

The principle of regarding anything not deducible as false is known as negation as failure: to deduce $\neg$ reachable $(X)$ it suffices to attempt to deduce reachable $(X)$ and fail. ${ }^{4}$ Forwardchaining provides a natural implementation strategy for negation-as-failure: once we have deduced all facts of the form reachable ( X ), if a particular such fact was not deduced, for example reachable(a-centauri), we regard this as proof of its negation.

However, because a forward-chaining system must wait until all facts reachable (X) are deduced before handling negative queries $\neg$ reachable $(X)$, it cannot handle such negative queries in reachable's own definition. This is the operational justification for stratified negation: we must be able to stratify our Datalog program into layers, each of which may only use the negation of predicates defined in the preceding layers.

Returning to our computational pattern, if range-restriction and constructor-freedom are about establishing the ascending chain condition, stratification is about establishing monotonicity within each stratum - each recursively-defined relation or group of relations. These strata correspond to individual fixed point computations. But negation is non-monotone: as its input grows toward truth, its result decreases to falsehood. This makes applying a rule with a negated premise $\neg \mathrm{P}(\ldots)$ dangerous: if as our knowledge grows we learn that $\mathrm{P}(\ldots)$ holds after all, we must retract our conclusion because it is not deducible. But this violates the condition that our state - the set of deduced atoms - grows over time!
${ }^{4}$ The closely related closed-world assumption states that whatever is true is known to be true; conversely, anything not known to be true is false. If we take "known" to mean "deducible", then the closed-world assumption justifies negation as failure.

### 1.3 Datalog for static analysis

Datalog has been successfully applied in various domains: for business analytics (Aref et al., 2015), as a general purpose database query language (Hickey et al., 2012), in network protocols (Alvaro et al., 2010; Loo et al., 2009), and for implementing distributed systems algorithms (Alvaro et al., 2011; Conway et al., 2012). But probably Datalog's most significant real-world adoption has been as a tool for scalable static analysis.

Datalog makes defining simple static analyses almost trivial. For instance, the essence of reaching definitions analysis can be expressed as follows:

```
reaches \((\mathrm{L}, \mathrm{V}, \mathrm{L}) \leftarrow \operatorname{assigns}(\mathrm{L}, \mathrm{V})\).
reaches \(\left(\mathrm{L}_{\text {dest }}, \mathrm{V}, \mathrm{L}_{\text {src }}\right) \leftarrow \neg \operatorname{assigns}\left(\mathrm{L}_{\text {dest }}, \mathrm{V}\right)\), reaches \(\left(\mathrm{L}_{\text {prev }}, \mathrm{V}, \mathrm{L}_{\text {src }}\right)\), flows \(\left(\mathrm{L}_{\text {prev }}, \mathrm{L}_{\text {dest }}\right)\).
```

If flows $\left(\mathrm{L}_{1}, \mathrm{~L}_{2}\right)$ means that line $\mathrm{L}_{1}$ may transfer control to $\mathrm{L}_{2}$, and assigns $(\mathrm{L}, \mathrm{V})$ means that line $L$ assigns to variable $V$, the above defines reaches $\left(L_{\text {dest }}, V, L_{\text {src }}\right)$ to mean that the assignment to V at line $\mathrm{L}_{\text {src }}$ may reach $\mathrm{L}_{\text {dest }}$.

Datalog's fluency at defining static analyses is not limited to toy examples. For instance, it has been successfully commercialized by Semmle, a company which uses a custom in-house Datalog dialect \& engine (Avgustinov et al., 2016) to do static analysis on large codebases to find potential security vulnerabilities - in particular variant analysis, which searches for variants of previously discovered issues. Their system has been used on NASA's Curiosity Rover, ${ }^{5}$ at Microsoft, ${ }^{6}$ and at the Nasdaq stock exchange company, ${ }^{7}$ among others.

On the academic side, the Doop project implements a state-of-the-art points-to analysis for Java code entirely in Datalog (Bravenboer and Smaragdakis, 2009). In a testimonial of sorts for the application of Datalog to static analysis, Smaragdakis and Bravenboer (2010) give several reasons why they found Datalog to be useful when implementing a scalable points-to analysis, compared in particular with a conventional language like Java or C++:

1. The high-level, declarative nature of Datalog allowed them to experiment with implementation techniques and algorithmic tweaks without having to entirely rewrite their analyses; for instance, they needed to carefully choose how to index their relations for maximal performance, a choice that would in a conventional language involve rewriting your data-access code. In their words:
[W]e believe that our ability to efficiently optimize our implementation was largely due to the declarative specifications of analyses. Working at the Datalog level eliminated much of the artificial complexity of a points-to analysis implementation, allowing us to concentrate on indexing optimizations and on the algorithmic essence of each analysis. (p. 1)
2. Datalog easily handles highly mutually-recursive relation definitions, which are common in static analysis: for example, "the logic for computing a callgraph depends on having

[^3]points-to information for pointer expressions, which, in turn, requires a callgraph." (p. 3) They also observe that one of their crucial optimizations, performing exception analysis on-the-fly, "would be quite hard to consider in a non-declarative context" because it "results in highly recursive definitions of core relations" (p. 6).
3. Rather than hand-optimizing their code as might be necessary in a conventional language, in Datalog they could rely on the language implementation to do a good chunk of this work for them by applying decades of work on query optimisation:

> We relied on query optimization (i.e., intra-rule, as opposed to inter-rule, optimization) being performed automatically. This was crucial for performance and, although a straightforward optimization in the context of database relations, results in far more automation than programming in a mainstream high-level language. (p. 6)
4. As we've seen with our examples so far, Datalog is very concise:

Generally, the declarative nature of Doop often allows for very concise specifications of analyses. We show in an earlier publication the striking example of the logic for the Java cast checking-i.e., the answer to the question "can type A be cast to type B?" The Datalog rules are almost an exact transcription of the Java Language Specification. (p. 4)

We must also consider that static analysis blunts one of Datalog's primary drawbacks. As mentioned previously, one weakness of forward chaining is that it is undirected, deducing everything it can. If we only need the results of a narrow query, this can waste a lot of effort. Various optimizations and alternative evaluation strategies exist to address this problem, including magic sets (Bancilhon et al., 1986; Beeri and Ramakrishnan, 1987), which statically rewrites recursive relation definitions to produce only a subset of facts relevant to a given query (for instance, automatically transforming all-pairs graph reachability to single-source reachability); and tabling (Swift and Warren, 2012; Tekle and Liu, 2011), which hybridizes forward- and backward-chaining by careful use of memoization. In static analysis, however, the eagerness of forward-chaining is less problematic, as we generally wish to analyse the whole program exhaustively, whether for optimization or for bug-finding purposes.

### 1.4 What Datalog can't do

In §1.2 we discussed Datalog's semantics and the deliberate restrictions that make it tractable; in $\S 1.3$ we saw how this made Datalog an attractive language for writing static analyses. However, Datalog's restrictions are not without their drawbacks. In this section will consider four things which Datalog does not support and their use-cases: (1) functional abstraction, (2) semilattices other than set union, (3) arithmetic and aggregations, and (4) compound data.

### 1.4.1 Functional abstraction

Consider our graph-reachability example again:

```
reachable(start).
reachable (Y) \leftarrow reachable (X), edge (X, Y).
```

Suppose we wish to compute reachability over multiple different graphs. One way is to repeat ourselves:

```
reachable1(start1).
reachable1 (Y) \leftarrow reachable (X), edge1 (X, Y).
reachable2(start2).
reachable2(Y) \leftarrow reachable (X), edge2 (X, Y).
reachable3(start3).
reachable3(Y) \leftarrow reachable (X), edge3(X, Y).
```

This quickly becomes obnoxious if we wish to repeat something more complex than a twoliner like graph reachability. In an ordinary language, we would factor out this repeated code into a procedure or a function. Unfortunately, Datalog cannot do this: Datalog does not have functions, procedures, or modules, only relations; and relations are first-order, unable to manipulate or abstract over other relations. It is unclear how to lift this restriction without giving up the guarantees that make forward-chaining Datalog evaluation tractable.

### 1.4.2 Semilattices other than set union

We have shown how to define reachability and reaching-definitions in Datalog. What about our second example, single-source shortest paths? A naive approach at expressing this in Datalog might look like this:

```
shortest(start, 0).
shortest \(\left(\mathrm{Y}, \mathrm{D}_{3}\right) \leftarrow \operatorname{shortest}\left(\mathrm{X}, \mathrm{D}_{1}\right)\), edge \(\left(\mathrm{X}, \mathrm{Y}, \mathrm{D}_{2}\right), \mathrm{D}_{1}+\mathrm{D}_{2}=\mathrm{D}_{3}\).
```

The immediate issue here is the use of the infinite relation $D_{1}+D_{2}=D_{3}$; recall that infinite relations can cause a problem for a forward-chaining evaluator. However, in this case $\mathrm{D}_{1}$ and $D_{2}$ are supplied by the first two premises, and together these allow computing $D_{3}$. The deeper problem is that, despite its name, shortest $(\mathrm{X}, \mathrm{D})$ finds all distances D from start to X rather than only the shortest. Besides failing to capture our intent, if the graph has cycles, there may be infinitely many such distances, causing an infinite loop.

Datalog really only understands one semilattice - finite sets under union - and has no way to specify that multiple sources of information, like the distance $D$ in shortest $(X, D)$, should be combined using a different strategy. This puts any computation using a custom semilattice out of reach. This particularly impacts Datalog's application to static analysis, where custom semilattices are frequently used to represent carefully-chosen approximations of the full set of values an expressions or variables may take on - for instance, "flat" lattices for constant propagation, or numeric intervals (or for multiple variables, convex polyhedra) under convex hull and intersection.

### 1.4.3 Arithmetic, user-defined functions, and aggregation

Pure Datalog does not permit using arithmetic or other functions. Although it is not difficult to extend a forward-chaining Datalog evaluator with support for these, and consequently almost everybody does so, this sacrifices Datalog's termination guarantee, as we saw with shortest paths. Even more problematic is support for aggregations, without which arithmetic by itself is of limited use. We also saw the need for aggregation in our shortest paths example: after calculating the lengths of multiple paths to the same node, we need to aggregate them and keep only their minimum.

In this example, our desired aggregation arises from a semilattice - we are taking the least upper bound $\bigvee_{p \in s}$ length $(p)$ over some set $s$ of paths $p$; to add support for other semilattices to Datalog, we need the ability to compute these semilattice aggregations. Semilattice aggregations are well-behaved because they are monotone: the value of $\bigvee_{x \in s} f(x)$ grows monotonically with respect to both the set $s$ and the function $f$. Thus the ascending chain condition provides sufficient conditions for their termination when used as part of a fixed point computation. However, not all useful aggregations form semilattices; for instance summing or averaging do not. Aggregations like this may or may not be monotone depending upon the order involved. Thus adding aggregations without careful restrictions can invalidate Datalog's least-fixed-point semantics.

### 1.4.4 Compound data

Datalog's constructor-freedom prevents code that would directly manipulate compound data structures, like lists, trees, or even tuples. This restriction rules out some infinite relations, such as our "lists of digits" example from page 8 , but it also rules out some legitimate, finite use-cases as well. For instance, Smaragdakis and Bravenboer (2010) added a macro system to their Datalog dialect to make writing context-sensitive static analyses less of a chore.

In a context-sensitive analysis, each rule depends upon a context representing an approximation of the conditions the code is executing under; for example, what call-site the function being analysed was invoked from. A deep context requires multiple pieces of data to specify (e.g. the preceding two or three functions on the call stack). It would be convenient to bundle this information into a single piece of data, hiding the choice of exactly how deep the context is, a choice orthogonal to the essence of the analysis. Although logically and operationally unproblematic, this use of compound data is not possible in Datalog, barring a macro system or some more principled extension.

### 1.5 Our goal and strategy

We started by introducing the computational pattern of monotone fixed points. We've seen that Datalog can express and compute some, but not all, instances of this pattern. Datalog's limitations are both practical - it is not obvious how to extend Datalog with higher-order abstraction - and theoretical - real world Datalog engines often support aggregation, arithmetic, and compound data, but in doing so raise the question of these features' semantics.

The goal of this thesis is to design a language which improves on Datalog's ability to express monotone fixed point computation over semilattices by finding ways to lift Datalog's restrictions without sacrificing either its simple semantics or its practical implementation
strategies. Our approach will be to combine Datalog with typed functional programming, on the basis of three hypotheses:

1. We can gain functional abstraction by mixing Datalog with higher-order functional programming. From a traditional logic programming perspective, this is a strange move: functions are a special-case of relations, so the "natural" way to make a logic language higher-order is to allow higher-order relations.
Datalog's power, however, comes from carefully limiting, not expanding, how relations may be defined; precisely because of their generality, higher-order relations are more complex to implement than higher-order functions, especially if one wishes to keep a natural semantics (Charalambidis et al., 2013). By separating our facility for deduction (relations, queries, \& fixed points) from our facility for abstraction (functions), we hope to gain the best of both worlds.
2. We can capture the restrictions that make Datalog work by deconstructing it typetheoretically. Type theory studies compositional properties of programs: if we recast syntactic restrictions (stratification, constructor-freedom) in terms of the properties they ensure (monotonicity, ACC), we can design a type system to capture these properties by finding compositional ways to provide them. Guided by this type system and its semantics, we can add practical features to our language, such as semilattices, aggregation, arithmetic, and compound datatypes, without sacrificing these properties.

## Chapter 2

## The Datafun Language

### 2.1 Syntax sketch

The idea behind Datafun is to capture the essence of Datalog in a typed, higher-order, functional setting. Since the key restriction that makes Datalog's combination of recursion and negation tractable - stratification - requires tracking monotonicity, we locate Datafun's semantics in the category Poset of partial orders and monotone maps. Since Poset is cartesian closed, it can interpret the simply typed $\lambda$-calculus, giving us a notation for writing monotone and higher-order functions. This lets us abstract over Datalog rules, something impossible in Datalog itself! In this section we sketch the construction of Datafun hewing closely to this semantic intuition.

Datafun begins as the simply-typed $\lambda$-calculus with functions ( $\lambda X$. e and ef), sums ( $\mathrm{in}_{i} e$ and case $e$ of $\ldots$ ), and products ( $(e, f)$ and $\pi_{i} e$ ). To represent relations, we add a type of finite sets $\left\{A_{\varepsilon Q}\right\},{ }^{1}$ introduced with set literals $\left\{e_{0}, \ldots e_{n}\right\}$, and eliminated using Moggi's monadic bind syntax, for $\left(x \in e_{1}\right) e_{2}$, which binds $x$ in $e_{2}$ successively to each element of $e_{1}$ and takes the union; in other words, $\bigcup_{x \in e_{1}} e_{2}$. Since we are working in Poset, each type comes with a partial order on it; sets are ordered by inclusion, $x \leqslant y:\left\{A_{\varepsilon Q}\right\} \Longleftrightarrow x \subseteq y$.

As long as all primitives are monotone, every definable function is also monotone. This is necessary for defining fixed points, but may seem too restrictive. There are many useful non-monotone operations, such as equality tests $e=f$. For example, $\}=\{ \}$ is true, but if the first argument increases to $\{1\}$ it becomes false, a decrease (as we'll see later, in Datafun, false $<$ true).

How can we express non-monotone operations if all functions are monotone? We cut this Gordian knot using a discreteness type constructor, $\square A$. The elements of $\square A$ are the same as those of $A$, but the partial order on $\square A$ is discrete, $x \leqslant y: \square A \Longleftrightarrow x=y$. Monotonicity of a function $\square A \rightarrow B$ is vacuous: $x=y$ implies $f(x) \leqslant f(y)$ by reflexivity. In this way we represent ordinary, possibly non-monotone, functions $A \rightarrow B$ as monotone functions $\square A \rightarrow B$.

Semantically, $\square$ is a monoidal comonad or necessity modality, and so we base our syntax on Pfenning and Davies (2001)'s syntax for the necessity fragment of constructive S4 modal logic. This involves distinguishing two kinds of variable: discrete variables are in lowercase

[^4]| types | A, B | $:=$ | $\{\mathrm{A}\}\|1\| A \times B\|A+B\| A \rightarrow B \mid \square A$ |
| :---: | :---: | :---: | :---: |
| eqtypes | $\underset{\text { eq }}{\text { A, }}$, B | $:=$ |  |
| semilattices | L, M | $:=$ |  |
| finite eqtypes ${ }^{2}$ | A, ${ }_{\text {fin }}$ | $:=$ |  |
| fixtypes | $\underset{\mathrm{Flx}}{\mathrm{L}}$, M | : $=$ |  |
| terms | $e, f$ | := | ```X\|x|\lambdaX.e|ef|()|(e,f)| | | e in}\mp@subsup{i}{i}{e| {\mp@subsup{e}{i}{}\mp@subsup{}}{i}{}|}|\mathrm{ for (x ( ``` |
|  |  |  | [e] \| let $[\mathrm{x}]=\mathrm{e}$ in f |
|  |  |  | $e=f \mid$ empty ${ }^{\text {e }} \mathrm{e} \mid$ split $e$ |
|  |  |  | $\perp\|e \vee f\| f i x e$ |

monotone variables $X, Y, Z$ are abstract names discrete variables $x, y, z$ are abstract names

Figure 2.I Datafun syntax
( $x, y$, foo, bar), while monotone variables are capitals ( $X, Y, Z$ ). Discrete variables may be used without restrictions, but monotone variables may only be used in ways that respect the ordering on their types: they must be used monotonically. This is enforced by a kind of variable hygiene: we remove monotone variables from scope within non-monotone expressions. For example, we cannot compare two monotone variables for equality, $X=Y$, because equality comparison is non-monotone. To aid the reader, we highlight non-monotone expressions with a light blue background; monotone variables bound outside of a non-monotone expression like $e=f$ may not be used within it. Putting this all together, we construct the type $\square A$ with the non-monotone introduction form $[e]$ and eliminate it by pattern-matching, let $[x]=e$ in $f$, giving access to a discrete variable $x$.

Finally, to express Datalog's recursive queries, Datafun includes a fixed point combinator fix, which computes the least fixed point of a map $f$. To ensure termination, this map must be monotone and take a semilattice type with decidable equality and no infinite ascending chains, $\underset{\sim}{L \times 1} \times$. For our purposes, a semilattice is a partial order with a least element $\perp$ and a least upper bound operation $\vee$ ("join"). Finite sets (with the empty set as least element, and union as join) are an example, as are tuples of semilattices. As long as the semilattice has no infinite ascending chains, the chain of iterations $\perp \leqslant f(\perp) \leqslant f(f(\perp)) \leqslant \ldots$ is guaranteed to stabilize at the least fixed point $f^{i}(\perp)=f^{i+1}(\perp)$ for some finite $i$. Decidable equality ensures we can tell when this fixed point is reached. While not used by fixed point iteration itself, semilattice join is used to define $f$ in all our motivating examples (following the pattern from §1.1), so we don't bother introducing "posets with bottom" as a concept separate from semilattices. ${ }^{2}$

[^5]
### 2.2 Examples

For brevity and clarity, the examples that follow make use of some syntax sugar:

1. We mentioned earlier that Datafun's boolean type bool is ordered false $<$ true. This is because we encode booleans as sets of empty tuples, or the type $\{1\}$, so written because 1 is the "unit type" of empty tuples. We desugar true to the singleton $\{()\}$ and false to the empty set $\}$. In a loop over a boolean, for $(x \in e) f$, the variable $x$ contains no useful information; if for brevity we omit it, the condensed expression for (e) f may be thought of as a "one-sided" conditional:

$$
\text { for }(e) f= \begin{cases}f & \text { if } e \text { is true } \\ \perp & \text { if } e \text { is false }\end{cases}
$$

Compared with encoding booleans as a sum type $1+1$, our approach has the advantage that it can express the type of "monotone" predicates $\mathrm{P}: A \rightarrow$ bool such that $\mathrm{P}(x)$ may change from false to true as $x$ grows, but cannot revert from true to false.
2. We make use of pattern matching. Besides the usual sum/tuple patterns, we support box-patterns $[p]$ and equality-check patterns !e. Box patterns $[p]$ correspond to boxelimination let $[x]=e$ in $f$, and their effect is to make all of the variables bound by $p$ discrete. The equality-check pattern !e matches only a value equal to $e$ - this is particularly useful when combined with set comprehensions.
3. We make use of set comprehensions, which can be desugared into the monadic operator for (Wadler, 1992). We also support looping/comprehending over only those elements of a set which match a certain pattern.
4. We express fixed points as a binding form, fix $X$ is $e$, instead of explicitly supplying a lambda to the fix combinator, fix $[\lambda X . e]$.
5. We permit ourselves a top-level surface syntax similar to Haskell or SML. In particular, we allow curried function definitions, type aliases, and algebraic datatype definitions:

```
disjunction : bool }->\mathrm{ bool }->\mathrm{ bool
disjunction X Y = X V Y
type colorname = string
data color = BLACK | NAMED colorname | RGB int int int
```

We do not allow data-types to be defined recursively, so they can be easily desugared into sums of products in the standard way. Similarly, we allow ourselves n-ary tuples, which are easily desugared into nested binary tuples.

We summarize the desugaring rules we use in figure 2.2, excepting our top-level declarations and the desugaring of algebraic data types, which should be fairly familiar.

$$
\begin{aligned}
& \text { types A, B : }=\ldots \text { | bool } \\
& \text { terms } e, f:=\ldots \text { | true | false } \\
& \text { for (C) } e \mid\{e \mid C\} \\
& \text { if } p \leftarrow e \text { then } f_{1} \text { else } f_{2} \\
& \text { fix } X \text { is } e \\
& \text { discrete patterns } p \quad:=x\left|-|!e|\left(p_{1}, p_{2}\right)\right| i n_{i} p \\
& \text { loop clauses } C, D:=p \in e|e| C, D \mid \\
& \text { bool } \xrightarrow{\text { desugar }}\{1\} \\
& \text { for }() e \xrightarrow{\text { desugar }} e \\
& \text { false } \xrightarrow{\text { desugar }}\} \\
& \text { for (C, D) e } \xrightarrow{\text { desugar }} \text { for (C) for (D) } e \\
& \text { true } \xrightarrow{\text { desugar }}\{()\} \\
& \text { for }(e) f \xrightarrow{\text { desugar }} \text { for }\left(\_\in e\right) f \\
& \{e \mid C\} \xrightarrow{\text { desugar }} \text { for }(C)\{e\} \quad \text { for }(p \in e) f \xrightarrow{\text { desugar }} \text { for }(x \in e) \text { if } p \leftarrow x \text { then } f \text { else } \perp \\
& \text { fix } X \text { is e } \xrightarrow{\text { desugar }} \text { fix [ } \lambda X, e \text { ] } \\
& \lambda[p] . e \xrightarrow{\text { desugar }} \lambda Y \text {. let }[p]=Y \text { in } e \\
& \text { let }[(x, y)]=e \text { in } f \xrightarrow{\text { desugar }} \text { let }[z]=e \text { in let }[x]=\left[\pi_{1} z\right] \text { in let }[y]=\left[\pi_{2} z\right] \text { in } f \\
& \text { if }{ }_{-} \leftarrow e \text { then } f_{1} \text { else } f_{2} \xrightarrow{\text { desugar }} f_{1} \\
& \text { if } x \leftarrow e \text { then } f_{1} \text { else } \mathrm{f}_{2} \xrightarrow{\text { desugar }} \text { let }[\mathrm{x}]=[\mathrm{e}] \text { in } \mathrm{f}_{1} \\
& \text { if }!e_{1} \leftarrow e_{2} \text { then } f_{1} \text { else } f_{2} \xrightarrow{\text { desugar }} \text { case empty? }\left(e_{1}=e_{2}\right) \text { of } \text { in }_{1} \rightarrow f_{2} ; \text { in }_{2} \rightarrow f_{1} \\
& \text { (NB. empty? e yields in } \mathrm{n}_{1} \text { if e is empty, i.e. false.) } \\
& \text { if }\left(p_{1}, p_{2}\right) \leftarrow e \text { then } f_{1} \text { else } f_{2} \xrightarrow{\text { desugar }} \text { if } p_{1} \leftarrow \pi_{1} e \\
& \text { if } \mathrm{in}_{i} p \leftarrow e \text { then } f_{1} \text { else } f_{2} \xrightarrow{\text { desugar }} \text { case split }[e] \text { of } \\
& \operatorname{in}_{i} X \rightarrow \text { let }[y]=X \text { in (if } p \leftarrow y \text { then } f_{1} \text { else } f_{2} \text { ) } \\
& \operatorname{in}_{(3-i)} X \rightarrow f_{2}
\end{aligned}
$$

Fresh variables introduced by desugaring are colored pink.
FIGURE 2.2 Syntax sugar

### 2.2.1 Set operations and relational algebra

One of the main features of Datafun is that it permits manipulating relations as first class values. In this subsection we will show how a variety of standard operations on sets can be represented in Datafun. The first operation we consider is testing membership:

```
member: }\square\textrm{E
member [x] S = for ( }\textrm{y}\in\textrm{S})\textrm{x}=\textrm{y
```

This checks if $x$ is equal to any element $y \in S$. The argument $x$ is discrete because increasing $x$ might send it from being in the set to being outside the set: although $1 \leqslant 2$ and $1 \in\{1\}$, nonetheless $2 \notin\{1\}$. Notice that here we're taking advantage of encoding booleans as sets of empty tuples - unioning these sets implements logical or.

Next we turn to set union and intersection. Union is baked into Datafun as the semilattice join, $x \cup y=x \vee y$, while intersection is definable using member, by taking the union of every singleton $\{x\}$ such that $x$ is in both $s$ and $t$ :

$$
\begin{aligned}
& S \cap T=\text { for }(x \in S \text {, member }[x] T)\{x\}
\end{aligned}
$$

Using comprehensions, this could alternately be written as:

$$
S \cap T=\{x \mid x \in S, \text { member }[x] T\}
$$

From now on, we'll use comprehensions whenever possible. For example, comprehensions make cross product and relational composition look almost exactly like their mathematical definitions:

$$
\begin{aligned}
& S \times T=\{(x, y) \mid x \in S, y \in T\}
\end{aligned}
$$

$$
\begin{aligned}
& S \bullet T=\left\{(a, c) \mid\left(a, b_{1}\right) \in S,\left(b_{2}, c\right) \in T, b_{1}=b_{2}\right\}
\end{aligned}
$$

The definitions of functional programming stalwarts filter and map (or in relational algebra terms, select and project) are slightly complicated by the need to be explicit about (non-) monotonicity:

$$
\begin{aligned}
& \text { filter } F S=\{x \mid x \in S, F[x]\}
\end{aligned}
$$

$$
\begin{aligned}
& \text { map }[f] S=\{f[x] \mid x \in S\}
\end{aligned}
$$

Why is filter monotone in its function argument while map is not? Recall that functions are ordered pointwise while sets are ordered by inclusion, and observe that increasing the filtering function (making it true on more inputs) enlarges the result of filter, but the same does not hold for map:

$$
\begin{aligned}
& \text { filter }(\leqslant 0)\{0,1\}=\quad\{0\} \\
& \operatorname{map}(\leqslant 0)\{0,1\}=\{0,1\}=\text { filter }(\leqslant 1)\{0,1\} \\
& =\{\text { true }, \text { false }\} \nsubseteq\{\text { true }\}=\operatorname{map}(\leqslant 1)\{0,1\}
\end{aligned}
$$

We can also define set difference, although we must first detour into boolean negation:

```
\(\neg: \square\) bool \(\rightarrow\) bool
\(\neg[\mathrm{t}]=\) case empty? t of \(\mathrm{in}_{1}() \rightarrow\) true; \(\mathrm{in}_{2}() \rightarrow\) false
```



```
\(S \backslash[t]=\{x \mid x \in S, \neg[\) member \([x] t]\}\)
```

To implement boolean negation, we need the primitive operator empty? e, which produces a tag indicating whether its argument $e$ (a boolean, i.e. a set of empty tuples) is the empty set. This in turn lets us define set difference, the analogue in Datafun of negation in Datalog. Note that in both boolean negation and set difference the "negated" argument $t$ is boxed, because the operation is not monotone in $t$. This enforces stratification.

Finally, generalizing our Datalog graph reachability example in $\S 1.2$, we can define the transitive closure of a relation:

```
trans: \(\square\{\underset{\text { FiN }}{A} \times \underset{\text { fin }}{A}\} \rightarrow\{\underset{\text { fiN }}{A} \times \underset{\text { fiN }}{A}\}\)
trans [edge] \(=\) fix R is edge \(\vee\) (edge \(\bullet \mathrm{R})\)
```

This definition uses a least fixed point, just like the mathematical definition - a transitive closure is the least relation R including both the original relation edge and the composition of edge with $R$. However, a peculiar feature of this definition is that the argument type is $\square\left\{_{\text {FIN }} \times A_{\text {FN }}\right\}$; transitive closure takes a discrete relation. This is because we must use the relation within the fixed point, and Datafun treats fix as a discrete operator. This restriction is artificial - transitive closure is semantically a monotone operation - but its explanation will have to wait until §3.3.5.

### 2.2.2 Regular expression combinators

Datafun permits tightly integrating the higher-order functional and bottom-up logic programming styles. To illustrate the benefits of doing so, in this section we implement a regular expression matching library in combinator style. Like combinator parsers in functional languages, the code is very concise. However, support for the relational style ensures we can write naïve code without the exponential backtracking cliffs typical of parser combinators in functional languages.

For these examples we'll assume the existence of eqtypes string, char, and int, an addition operator + , and functions length and chars satisfying:

```
length : }\square\mathrm{ string }->\mathrm{ int
length [s] = the length of s
chars: }\square\mathrm{ string }->\mathrm{ {int }\times\mathrm{ char}
chars [s]={(i,c)| the ith character of s is c}
```

Note that by always boxing string arguments, we avoid committing ourselves to any particular partial ordering on string.

These assumed, we define the type of regular expression matchers:
type regex $=\square$ string $\rightarrow\{$ int $\times$ int $\}$

A regular expression takes a discrete string [s] and returns the set of all pairs $(i, j)$ such that the substring $s_{i}, \ldots, s_{j-1}$ matches the regular expression. For example, to find all matches for a single character $c$, we return the range $(i, i+1)$ whenever $(i, c) \in$ chars $[s]$ :

```
sym : }\square\mathrm{ char }->\mathrm{ regex
sym [c] [s]={(i,i+1)|(i,!c)\in chars[s]}
```

To find all matches for the empty regex, i.e. all empty substrings, including the one "beyond the last character":

```
nil : regex
nil [s]={(i,i)|(i,_) \in chars [s]}\vee{(length [s], length [s])}
```

Appending regexes $R_{1}, R_{2}$ amounts to relation composition, since we wish to find all substrings consisting of adjacent substrings $s_{i} \ldots s_{j-1}$ and $s_{j} \ldots s_{k-1}$ matching $R_{1}$ and $R_{2}$ respectively:

```
seq : regex }->\mathrm{ regex }->\mathrm{ regex
seq R R R R S = R R S \bullet R2 S
```

Similarly, regex alternation $r_{1} \mid r_{2}$ is accomplished by unioning all matches of each:

$$
\begin{aligned}
& \text { alt }: \text { regex } \rightarrow \text { regex } \rightarrow \text { regex } \\
& \text { alt } R_{1} R_{2} S=R_{1} S \vee R_{2} S
\end{aligned}
$$

The most interesting regular expression combinator is Kleene star. Thinking relationally, if we consider the set of pairs $(i, j)$ matching some regex $r$, then $r *$ matches its reflexive, transitive closure. This can be accomplished by combining nil and trans.

```
star : }\square\mathrm{ regex }->\mathrm{ regex
star [r] [s]= nil [s] V trans [r [s]]
```

Note that the argument r must be discrete because trans uses it to compute a fixed point. ${ }^{3}$

### 2.2.3 Regular expression combinators, take two

The combinators in the previous section found all matches within a given substring, but often we are not interested in all matches: we only want to know if a string can match starting at a particular location. We can easily refactor the combinators above to work in this style, which illustrates the benefits of tightly integrating functional and relational styles of programming - we can use functions to manage strict input/output divisions, and relations to manage nondeterminism and search.

$$
\text { type regex }=\square(\text { string } \times \text { int }) \rightarrow\{\text { int }\}
$$

Our new type of combinators takes a string and a starting position, and returns a set of ending positions. For example, sym [c] checks if $c$ occurs at the start position $\mathfrak{i}$, yielding $\{i+1\}$ if it does and the empty set otherwise, while nil simply returns the start position $i$.

[^6]```
sym : \(\square\) char \(\rightarrow\) regex
\(\operatorname{sym}[\mathrm{c}][(\mathrm{s}, \mathrm{i})]=\{i+1 \mid!(i, c) \in \operatorname{chars}[s]\}\)
nil : regex \(\rightarrow\) regex
nil \([(s, i)]=\{i\}\)
```

Appending regexes seq $R_{1} R_{2}$ simply applies $R_{2}$ starting from every ending position that $R_{1}$ can find:

```
seq : regex }->\mathrm{ regex }->\mathrm{ regex
seq R R R R [(s,i)]= for (j }\in\mp@subsup{R}{1}{}[(s,i)])\mp@subsup{R}{2}{}[(s,j)
```

Regex alternation alt is effectively unchanged:

```
alt : regex \(\rightarrow\) regex \(\rightarrow\) regex
alt \(R_{1} R_{2} X=R_{1} X \vee R_{2} X\)
```

Finally, Kleene star is implemented by recursively appending $r$ to a set $X$ of matches found so far:

```
star : \(\square\) regex \(\rightarrow\) regex
\(\operatorname{star}[r][(s, i)]=f \mathbf{f i x} X\) is \((\{i\} \vee\) for \((j \in X) r[(s, j)])\)
```

It's worth noting that this definition is effectively left-recursive - it takes the endpoints from the fixed point $x$, and then continues matching using the argument $r$. This should make clear that this is not just plain old functional programming - we are genuinely relying upon the fixed point semantics of Datafun.

### 2.2.4 CYK parsing

Parsing can be understood logically: a parse tree is a proof that a string belongs to a language, and parsing is proof search (Shieber et al., 1995). One of the simplest parsing algorithms is the Cocke-Younger-Kasami (CYK) algorithm for parsing grammars in Chomsky normal form; that is, where each production is either of the form $A \rightarrow B C$ or $A \rightarrow \vec{a}$, with $A, B, C$ ranging over nonterminals and $\vec{a}$ over strings of terminals. Fix a Chomsky-normal grammar G and a word $w=w_{0} w_{1} \ldots w_{n-1}$ to be parsed, and write $w_{i . . j}$ for the substring $w_{i} \ldots w_{j-1}$. Now, we introduce a family of predicates $A(i, j)$ (sometimes called facts), intended to represent the proposition that $w_{i . . j}$ is generated by the nonterminal $A$. Then, we can specify the CYK algorithm with the following two inference rules:

$$
\frac{(A \rightarrow B C) \in G \quad B(i, j) \quad C(j, k)}{A(i, k)} \quad \frac{(A \rightarrow \vec{a}) \in G \quad \vec{a}=w_{i . . j}}{A(i, j)}
$$

Then the whole word $w$ is generated by the start symbol $S$ if $S(0, n)$ is deducible.
In Datafun, this rule-based description of the algorithm can be transliterated almost directly into code. We begin by introducing a few basic types.

```
type symbol = string
data rule = STRING string | CONCAT symbol symbol
type grammar ={symbol }\times\mathrm{ rule }
type fact = symbol }\times\mathrm{ int }\times\mathrm{ int
```

The symbol type is a type synonym representing nonterminal names with strings. The rule type is the type of the right-hand-sides of productions in Chomsky normal form - either a string, or a pair of nonterminals. A grammar is just a set of productions - a set of pairs of nonterminals paired with their rules. The type fact represents the atomic facts deduced by the CYK inference system - they are triples of the rulename, the start position, and the end position.

```
length : \(\square\) string \(\rightarrow\) int
range : \(\square \mathrm{int} \rightarrow\) int \(\rightarrow\{\mathrm{int}\}\)
substring : \(\square\) (string \(\times\) int \(\times\) int) \(\rightarrow\) string
\((+):\) int \(\rightarrow\) int \(\rightarrow\) int
\((-):\) int \(\rightarrow \square\) int \(\rightarrow\) int
```

With these types in hand, we can write the CYK algorithm as a fixed point computation. In fact, it is convenient to break it into two pieces, by first defining the function whose fixed point we take. So we can write down the iter function, which represents one step of the fixed point iteration.

```
iter : \(\square\) string \(\rightarrow \square\) grammar \(\rightarrow\{\) fact \(\} \rightarrow\{\) fact \(\}\)
iter [text] [grammar] \(\mathrm{F}=\)
        \(\{(a, i, k) \mid(a\), concat \(b c) \in \operatorname{grammar},(!b, i, j) \in F,(!c,!j, k) \in F\}\)
    \(\cup\{(a, i, i+\) length \(s) \mid(a\), string \(s) \in\) grammar,
        \(i \in\) range [ 0 ] (length text - [length \(s]\) ),
        substring [text, \(i, i+\) length \(s]\}\)
```

We can then use iter to implement the parse function:

```
parse : \(\square\) string \(\rightarrow \square\) grammar \(\rightarrow\) \{symbol \(\}\)
parse [text] [grammar] \(=\)
    \(\{\mathbf{a} \mid(\mathrm{a},!0\), !(length text) \() \in \mathbf{f i x} \mathbf{X}\) is iter [text] [grammar] X\(\}\)
```

This finds all nonterminals in grammar that generate the entire string text.
This program is not expressible in Datalog, because Datalog provides no way to abstract over grammars. The rules of a grammar are easily represented as Datalog relations - but since Datalog is first-order, it cannot parameterize one relation by another; so there is no way in Datalog to express a generic parser. This demonstrates one of the key benefits of moving to a functional language like Datafun.

### 2.2.5 Dataflow analysis

In this section, we show how some simple dataflow analyses can be expressed in Datafun. We begin with the types in these programs.

```
type var = string
type label = int
data op = EQ | LE | ADD | SUB | MUL| DIV
data atom = VAR var | NUM int
data expr = ATOM atom | APPLY op atom atom
data statement = ASSICN var expr | IF expr label label
type program = {label }\times\mathrm{ statement }
```

We represent a program as a set of nodes. Each node has a label and contains a statement, either an assignment (ASSIGN) or a conditional jump (IF). Valid programs $p$ associate any label $l$ with at most one node $(l, s) \in p$. In what follows, we use a few trivial functions whose definitions we omit.

```
labels: program }->{\mathrm{ {label}
uses: }\square\mathrm{ statement }->{\mathrm{ var}
defines: }\square\mathrm{ statement }->\mathrm{ {var}
```

The labels function returns the set of labels in a program. The uses function returns the set of variables used by the expressions in a statement. The defines function returns the set of variables defined by a statement (i.e., at most one variable - the target of the assignment).

Given a program, we can recover its 1-step control flow graph with the flow function:

```
type flow \(=\{\) label \(\times\) label \(\}\)
flow : program \(\rightarrow\) flow
flow \(\mathrm{P}=\) for \(((i, s) \in P)\)
    if \(\operatorname{IF} \quad \mathrm{j} k \leftarrow \mathrm{~s}\)
    then \(\{(i, j),(i, k)\}\)
    else \(\{(i, i+1) \mid!(i+1) \in\) labels \(P\}\)
```

This says that if a program node $(i, s)$ is a conditional jump, $I F_{\sim} j k$, then it may flow to either $j$ or $k$; otherwise, it flows to the next program position $i+1$ if it exists.

Using this we can define liveness analysis, one of the classic backwards dataflow analyses. The function live takes a program P and its flow graph F and produces a set of label/variable pairs $(i, v)$ indicating that the variable $v$ is live at program point $i$.

```
live : \(\square\) program \(\rightarrow \square\) flow \(\rightarrow\{\) label \(\times\) var \(\}\)
live [program] [flow] \(=\)
    fix \(L\) is
    for \(((i, s) \in\) program \()\)
        ( \(\{i\} \times\) uses \(s)\)
    \(\cup\{(\mathbf{i}, v) \mid(!i, j) \in\) flow, \((!j, v) \in \mathrm{L}, \neg[\) member \([v]\) (defines \(s)]\}\)
```

At each label $i$, a variable $v$ is live if either of two conditions holds: (1) it is used by the current statement $s$, or (2) it is live at some label $j$ to which $i$ flows, and which does not define the variable $v$ for itself.

Next we give one of the classic forwards dataflow analyses: reaching definitions. This determines which program points the value assigned by a particular statement (a "definition") can reach.

```
reaching : \(\square\) program \(\rightarrow \square\) flow \(\rightarrow\{\) var \(\times\) label \(\times\) label \(\}\)
reaching [program] [flow] \(=\)
    fix \(R\) is
    for \(((k, s) \in\) program \()\)
        \(\{(v, k, k) \mid v \in\) defines \(s\}\)
    \(\cup\{(v, \mathfrak{i}, k) \mid(\mathfrak{j},!\mathrm{k}) \in\) flow, \((v, \mathfrak{i},!\mathfrak{j}) \in \mathrm{R}, \neg[\) member \([v]\) (defines \(s)]\}\)
```

The function reaching takes a program and its flow graph, and returns a set of tuples ( $v, i, j)$ indicating that the definition of $v$ at $i$ might reach the program point $j$. These tuples are
contexts $\Gamma:=\varepsilon \mid \Gamma, H$ hypotheses $H:=X: A \mid x: A$

$$
\begin{aligned}
\lceil\varepsilon\rceil & =\varepsilon \\
\lceil\Gamma, \mathrm{X}: А\rceil & =\lceil\Gamma\rceil \\
\lceil\Gamma, x: A\rceil & =\lceil\Gamma\rceil, x:: A
\end{aligned}
$$

| var | DVAR | LAM | APP |  |
| :---: | :---: | :---: | :---: | :---: |
| $X: A \in \Gamma$ | $x:: A \in \Gamma$ | $\Gamma, \mathrm{X}: \mathrm{A} \vdash \mathrm{e}: \mathrm{B}$ | $\Gamma \vdash \mathrm{e}: \mathrm{A} \rightarrow \mathrm{B}$ | $\Gamma \vdash \mathrm{f}: ~ A$ |
| $\overline{\Gamma \vdash X: A}$ | $\overline{\Gamma \vdash x: A}$ | $\Gamma \vdash \lambda X . e: A \rightarrow B$ | $\Gamma \vdash e f$ |  |


| UNIT | PAIR | PRJ | INJ |
| :--- | :--- | :--- | :--- |
| $\overline{\Gamma \vdash(): 1}$ | $\frac{\left(\Gamma \vdash e_{i}: A_{i}\right)_{i}}{\Gamma \vdash\left(e_{1}, e_{2}\right): A_{1} \times A_{2}}$ | $\frac{\Gamma \vdash e: A_{1} \times A_{2}}{\Gamma \vdash \pi_{i} e: A_{i}}$ | $\frac{\Gamma \vdash e: A_{i}}{\Gamma \vdash \mathrm{in} i_{i} e: A_{1}+A_{2}}$ |

$$
\begin{array}{ll}
\text { CASE } & \text { вох } \\
\frac{\Gamma \vdash e: A_{1}+A_{2}}{\Gamma \vdash \text { case } e \text { of }\left(\text { in }_{i} X_{i} \rightarrow \mathrm{f}_{\mathrm{i}}\right)_{i}: B} & \frac{\Gamma \Gamma\rceil \vdash e: A}{\Gamma \vdash[e]: \square A}
\end{array}
$$

Letbox

$$
\begin{array}{ll}
\text { вот } & \text { JOIN }
\end{array}
$$

$\Gamma \vdash e: \square A \quad \Gamma, x:: A \vdash f: B$
$\Gamma \vdash$ let $[x]=e$ in $f: B$

$$
\overline{\Gamma \vdash \perp: L} \quad \frac{\left(I \vdash e_{i}: L\right)_{i}}{\Gamma \vdash e_{1} \vee e_{2}: L}
$$

$$
\frac{\left(\lceil\Gamma\rceil \vdash e_{i}: \underset{E Q}{A}\right)_{i}}{\Gamma \vdash\left\{e_{i}\right\}_{i}:\{\underset{E Q}{A}\}}
$$

$$
\left.\begin{array}{rl}
\frac{\text { FOR }}{\Gamma \vdash e:\{\underset{E Q}{A}\}} \quad & \Gamma, x:: \underset{E Q}{A} \vdash f: L \\
\Gamma \vdash \text { for }(x \in e) f: L
\end{array}\right]
$$

EQ
EMPTY

$$
\frac{\left(\lceil\Gamma\rceil \vdash e_{i}: \underset{E Q}{A}\right)_{i}}{\Gamma \vdash e_{1}=e_{2}: \text { bool }}
$$

$$
\frac{\lceil\Gamma\rceil \vdash e:\{1\}}{\Gamma \vdash \text { empty? } e: 1+1}
$$

## FIGURE 2.3 Datafun typing rules

generated by two rules, corresponding to the two clauses in reaching's inner loop: (1) if $v$ is defined at $i$, then it reaches $i$, and (2) if the definition of $v$ at $i$ reaches $j$ and $j$ flows to $k$ then the definition also reaches k unless $\mathfrak{j}$ has an intervening definition of $v$.

### 2.3 Typing and denotational semantics

Our guiding intuition so far has been that Datafun is a language for writing monotone, higher-order functions. Here we substantiate that intuition by giving typing rules for core Datafun and showing how to interpret well-typed Datafun terms into Poset, the category of partially ordered sets and monotone maps.

### 2.3.1 Typing rules

The syntax of core Datafun is given in figure 2.1 and its typing rules in figure 2.3. Contexts are lists of hypotheses $H$; a hypothesis gives the type of either a monotone variable $X$ : $A$ or a discrete variable $x:: A$. The typing judgement $\Gamma \vdash e: A$ may be read as "assuming the variables in $\Gamma$ have their given types, the term $e$ has type $A$; moreover, $e$ is monotone with
respect to the monotone variables in $\Gamma^{\prime \prime}$.
The var and dvar rules say that both monotone hypotheses $X$ : A and discrete hypotheses $x:=A$ justify ascribing their variable the type $A$. The lam rule is the familiar rule for $\lambda$ abstraction. However, note that we introduce the argument variable $X$ : $A$ as a monotone hypothesis, not a discrete one. (This is the "right" choice because in Poset the exponential object is the poset of monotone functions.) The application rule APP is standard, as are the rules unit, pair, prj, inj. Case analysis case is also standard, noting only that as with LAM, the variables $X_{i}: A_{i}$ bound in each branch $f_{i}$ are monotone.
box says that [e] has type $\square A$ when $e$ has type $\mathcal{A}$ in the stripped context $\lceil\Gamma\rceil$. The stripping operation $\lceil\Gamma\rceil$ drops all monotone hypotheses from the context $\Gamma$, removing them from scope in $e$ and implementing the "variable hygiene" discussed in §2.1. This ensures we don't smuggle any information we must treat monotonically into a discretely-ordered $\square$ expression. The elimination rule letbox for (let $[x]=e$ in $f$ ) allows us to "cash in" a boxed expression $e: \square A$ by binding its result to a discrete variable $x: A$ in the body $f$.

At this point, our typing rules correspond to standard constructive S4 modal logic (Pfenning and Davies, 2001). We get to Datafun by adding a handful of domain-specific types and operations. First, split provides an operator split : $\square(A+B) \rightarrow \square A+\square B$ to distribute box across sum types. The other direction, $\square A+\square B \rightarrow \square(A+B)$, is already derivable, as is the isomorphism $\square A \times \square B \cong \square(A \times B)$. This is used implicitly by box pattern-matching - e.g., in the pattern $\left[\left(\mathrm{in}_{1} x, \mathrm{in}_{2} y\right)\right]$, the variables $x$ and $y$ are both discrete, which is information we propagate via these conversions. Semantically, all of these operations are the identity, as we shall see shortly. ${ }^{4}$

This leaves only the rules for manipulating sets and other semilattices. вот and Join tell us that $\perp$ and $\vee$ are valid at any semilattice type $L$, that is, at sets and products of semilattice types. The rule for set-elimination, FOR, is almost a monadic bind. However, we generalize it by allowing for $(x \in e) f$ to eliminate into any semilattice type, not just sets, denoting a "big semilattice join" rather than a "big union".

The set-introduction rule SET gives $\left\{e_{i}\right\}_{i \in I}$ type $\left\{\underset{\in Q}{ }\left\{\begin{array}{l}\text { A }\end{array}\right\}\right.$ when each of the $e_{i}$ has type $\underset{E Q}{A}$. Just as in box, each $e_{i}$ has to typecheck in a stripped context; constructing a set is a discrete operation, since $1 \leqslant 2$ but $\{1\} \nsubseteq\{2\}$.

Likewise discrete is equality comparison $e_{1}=e_{2}$, whose rule EQ is otherwise straightforward; and EMPTY, which requires more explanation. The idea is that empty? e determines whether $e:\{1\}$ is empty, returning $\mathrm{in}_{1}()$ if it is, and $\mathrm{in}_{2}()$ if it isn't. This lets us turn "booleans" (sets of units) into values we can case-analyse. This is, however, not monotone, because while booleans are ordered false < true, sum types are ordered disjointly; $\mathrm{in}_{1}()$ and $\mathrm{in}_{2}()$ are incomparable.

Finally, the rule FIX says that the fixed point fix combinator accepts a boxed function $\mathrm{f}: \square(\mathrm{LIX} \rightarrow \underset{f \mid x}{\mathrm{~L}})$ and returns a value of type $\underset{\mathrm{fx}}{\mathrm{L}}$. The restriction to "fixtypes" ensures $\mathrm{L}_{\mathrm{fIX}}^{\mathrm{L}}$ has no infinite ascending chains, guaranteeing the recursion will terminate. The restriction to boxed functions, treating fix as a non-monotone operator, is motivated not by our semantics but by our strategy for evaluating Datafun efficiently. This will be explained in detail in §3.3.5, but as a foreglimpse, to evaluate Datafun efficiently, we incrementalize monotone functions; incrementally maintaining fix is difficult, so we treat it as non-monotone.

4 An alternative to box pattern-matching and split, pursued in Arntzenius and Krishnaswami (2016), would be to give two rules for case, depending on whether or not the scrutinee can be typechecked in a stripped context.

### 2.3.2 The category Poset and its structures

An object of Poset is a pair $\left(A, \leqslant_{A}\right)$ consisting of a set $A$ and a reflexive, transitive, antisymmetric relation $\leqslant_{A} \subseteq A \times A$. For convenience, we usually denote these by a single letter $A$, leaving $\leqslant_{A}$ implicit. Following this convention, a morphism $f: A \rightarrow B$ is a function such that $x \leqslant_{A} y \Longrightarrow f(x) \leqslant_{B} f(y)$.

## Bicartesian structure

The bicartesian closed structure of Poset is largely the same as in Set. The product and sum sets are constructed the same way, and ordered pointwise:

$$
\begin{aligned}
&(a, b) \leqslant A \times B \\
&\left.i n_{i} x \leqslant_{A_{1}+A_{2}}, b_{j}\right) \Longleftrightarrow a \leqslant A a^{\prime} \wedge b \leqslant B b^{\prime}
\end{aligned} \Longleftrightarrow \mathfrak{i}=j \wedge x \leqslant_{A_{i}} y
$$

Projections $\pi_{i}$, injections $\mathrm{in}_{\mathrm{i}}$, tupling $\langle\mathrm{f}, \mathrm{g}\rangle$ and case-analysis $[\mathrm{f}, \mathrm{g}]$ are all the same as in Set, pausing only to note that all these operations preserve monotonicity, as we need.

The exponential $A \Rightarrow B$ consists of the monotone maps $f: A \rightarrow B$, again ordered pointwise:

$$
f \leqslant_{A \Rightarrow B} g \Longleftrightarrow\left(\forall x \leqslant_{A} y\right) f x \leqslant_{B} g y
$$

Currying $\lambda$ and evaluation are the same as in Set. Supposing $f: A \times B \rightarrow C$, then:

$$
\begin{array}{ll}
\lambda(f): A \rightarrow(B \Rightarrow C) & \text { eval }_{A, B}:(A \Rightarrow B) \times A \rightarrow B \\
\lambda(f)=x \mapsto y \mapsto f(x, y) & \text { eval }_{A, B}=(g, x) \mapsto g(x)
\end{array}
$$

Monotonicity here follows from the monotonicity of $f$ and $g$ and the pointwise ordering of $A \Rightarrow B$.

## The discreteness comonad

Given a poset $\left(A, \leqslant_{A}\right)$ we define the discreteness comonad $\square\left(A, \leqslant_{A}\right)$ as $\left(A, \leqslant \square_{A}\right)$, where $a \leqslant \square A a^{\prime} \Longleftrightarrow a=a^{\prime}$. That is, the discrete order preserves the underlying elements, but reduces the partial order to mere equality. This forms a rather boring comonad whose functorial action $\square(f)$, extraction $\varepsilon_{A}: \square A \rightarrow A$, and duplication $\delta_{A}: \square A \rightarrow \square \square A$ are all identities on the underlying sets:

$$
\square(\mathrm{f})=\mathrm{f} \quad \varepsilon_{\mathrm{A}}=\mathrm{a} \mapsto \mathrm{a} \quad \delta_{\mathrm{A}}=\mathrm{a} \mapsto \mathrm{a}
$$

This makes the functor and comonad laws trivial. Monotonicity holds in each case because all functions are monotone with respect to $\leqslant \square A$. It is also immediate that $\square$ is monoidal with respect to both products and sums. That is, $\square(A \times B) \cong \square A \times \square B$ and $\square(A+B) \cong \square A+\square B$. In both cases the isomorphism is witnessed by identity on the underlying elements. These lift to $n$-ary products and sums as well, which we write as dist ${ }_{\square}^{\times}: \prod_{i} \square A_{i} \rightarrow \square \prod_{i} A_{i}$ and dist $_{+}^{\square}: \square \sum_{i} A_{i} \rightarrow \sum_{i} \square A_{i}$.

## Sets and semilattices

Given a poset ( $A, \leqslant_{A}$ ) we define the finite powerset poset $P\left(A, \leqslant_{A}\right)$ as ( $\left.\mathcal{P}_{\text {fin }} A, \subseteq\right)$, that is, the finite subsets of $A$ ordered by inclusion. ${ }^{5}$ Finite sets admit a pair of useful morphisms:

$$
\begin{array}{ll}
\text { singleton : } \square A \rightarrow P A & \text { isEmpty : } \square P A \rightarrow 1+1 \\
\text { singleton }=a \mapsto\{a\} & \text { isEmpty }=X \mapsto \begin{cases}\operatorname{in}_{1}() & \text { when } X=\emptyset \\
\operatorname{in}_{2}() & \text { otherwise }\end{cases}
\end{array}
$$

The singleton function takes a value and makes a singleton set out of it. The domain must be discrete, as otherwise the map will not be monotone (sets are ordered by inclusion, and set membership relies on equality, not the partial order). Similarly, the emptiness test isEmpty also takes a discrete set-valued argument, because otherwise the boolean test would not be monotone.

Sets also form a semilattice, with the least element given by the empty set, and join given by union. For this and other semilattices $L \in$ Poset, in particular products of semilattices, we write join $\mathrm{n}_{n}^{\mathrm{L}}: \mathrm{L}^{n} \rightarrow \mathrm{~L}$ to denote the n -ary semilattice join (least upper bound). Moreover, if $f: A \times \square B \rightarrow L$, we define a morphism collect( $f$ ) : $A \times P B \rightarrow L$ as follows:

$$
\operatorname{collect}(f)=(a, X) \mapsto \bigvee_{b \in X} f(a, b)
$$

We will use this to interpret for-loops. However, it is worth noting that the discreteness of singleton means finite sets do not quite form a monad in Poset.

## Equality

Every object $A \in$ Poset admits an equality-test morphism eq:

$$
\begin{aligned}
& \text { eq }: \square A \times \square A \rightarrow \mathrm{P} 1 \\
& \text { eq }=(x, y) \mapsto \begin{cases}\{()\} & \text { if } x=y \\
\emptyset & \text { otherwise }\end{cases}
\end{aligned}
$$

The domain must be discrete, since $x=y$ and $y \leqslant z$ certainly doesn't imply $x=z$. Although in principle every object $A \in$ Poset admits eq, in practice our semantics only uses it when equality is decidable.

## Fixed points

Given a semilattice $L \in$ Poset without infinite ascending chains, we can define a family of fixed point morphisms fix : $\square(\mathrm{L} \Rightarrow \mathrm{L}) \rightarrow \mathrm{L}$ as follows: ${ }^{6}$

[^7]$$
\mathrm{fix}=\mathrm{f} \mapsto \bigvee_{\mathrm{n} \in \mathbb{N}} \mathrm{f}^{\mathrm{n}}(\perp)
$$

A routine inductive argument shows this must yield a least fixed point.

### 2.3.3 Interpretation of Datafun in Poset

Figure 2.4 shows how to interpret Datafun into Poset using the structures developed above. We interpret Datafun types and typing contexts as Poset-objects $\llbracket A \rrbracket, \llbracket \Gamma \rrbracket$ and well-typed Datafun terms (or more precisely, their typing derivations) $\Gamma \vdash e: A$ as Poset-morphisms $\llbracket \Gamma \rrbracket \rightarrow \llbracket A \rrbracket$. This follows the usual interpretation for constructive S4 (Alechina et al., 2001), with the addition of sets, semilattices, fixed points, and the ability to distribute $\square$ over sums. We give the interpretation in combinatory style; to increase readability, we freely use $n$-ary products to represent our typing context, to avoid the book-keeping of reassociating binary products.

Regarding notation, we write composition in diagrammatic or "pipeline" order with a semicolon, so $f ; g: A \rightarrow C$ means $f: A \rightarrow B$ followed by $g: B \rightarrow C$. If $f_{i}: A \rightarrow B_{i}$ we write $\left\langle f_{i}\right\rangle_{i}: A \rightarrow \prod_{i} B_{i}$ for the "tupling map" such that $\left\langle f_{i}\right\rangle_{i} ; \pi_{j}=f_{j}$. In particular, $\rangle$ is the map into the terminal object. Dually, if $g_{i}: A_{i} \rightarrow B$ we write $\left[g_{i}\right]_{i}: \sum_{i} A_{i} \rightarrow B$ for the "case-analysis map" such that $\mathrm{in}_{\mathrm{j}} ;\left[\mathrm{g}_{\mathrm{i}}\right]_{i}=\mathrm{g}_{\mathrm{j}}$.

### 2.4 Operational semantics

We consider the denotational semantics to be primary in Datafun; as with Datalog, any implementation technique is valid so long as it lines up with this semantics. To show such an implementation is possible, we present a simple call-by-value structural operational semantics in figure 2.5 and show that all well-typed terms terminate. In our operational semantics we:

1. Drop the distinction between discrete and monotone variables, writing both in lowercase $x, y, z$, and cease using a light blue background for non-monotone expressions.
2. Assume all equality tests and all semilattice operations ( $\perp, \vee$, for, and fix) are subscripted with their type.
3. Add iter expressions, which occur as intermediate forms in the evaluation of fix.

We use a small-step operational semantics with evaluation contexts E (Felleisen and Hieb, 1992) to enforce a call-by-value evaluation order; an evaluation context $E$ is an expression with a hole in it, written $O$, such that whatever is in the hole is next in line to be evaluated (if it is not a value already). To fill the hole in an evaluation context E with the expression $e$, we write $\mathrm{E}\{\mathrm{O} \mapsto \mathrm{e}\}$.

We define a relation $e \longmapsto e^{\prime}$ for expressions $e$ whose outermost structure is immediately reducible; we extend this relation to all expressions with the rule:

$$
\begin{aligned}
& \text { EVAL CONTEXT } \\
& \stackrel{e}{\mathrm{E}\{\mathrm{O} \mapsto \mathrm{e}\}} \mathrm{e}^{\prime} \longmapsto \mathrm{E}\left\{\mathrm{O} \mapsto e^{\prime}\right\}
\end{aligned}
$$

$$
\begin{array}{rlrl}
\llbracket 1 \rrbracket & =1 & \llbracket A \rightarrow B \rrbracket & =\llbracket \mathrm{A} \rrbracket \Rightarrow \llbracket \mathrm{~B} \rrbracket \\
\llbracket\{\mathrm{~A}\} \rrbracket & =\mathrm{P} \llbracket \mathrm{~A} \rrbracket & \llbracket \mathrm{~A} \times \mathrm{B} \rrbracket & =\llbracket \mathrm{A} \rrbracket \times \llbracket \mathrm{B} \rrbracket \\
\llbracket \square \mathrm{~A} \rrbracket & =\square \llbracket \mathrm{A} \rrbracket & \llbracket \mathrm{~A}+\mathrm{B} \rrbracket & =\llbracket \mathrm{A} \rrbracket+\llbracket \mathrm{B} \rrbracket \\
\llbracket \Gamma \rrbracket=\prod_{H \in \Gamma} \llbracket \mathrm{H} \rrbracket \\
\llbracket \mathrm{X}: \mathrm{A} \rrbracket=\llbracket \mathrm{A} \rrbracket & \llbracket x:: \mathrm{A} \rrbracket=\square \llbracket \mathrm{A} \rrbracket & \llbracket \Gamma \vdash \mathrm{~A} \rrbracket=\operatorname{Poset}(\llbracket \Gamma \rrbracket, \llbracket \mathrm{A} \rrbracket)
\end{array}
$$

TERM DENOTATIONS

$$
\begin{aligned}
& \llbracket \Gamma \vdash X: A \rrbracket=\pi_{X} \quad(\text { for } X: A \in \Gamma) \\
& \llbracket \Gamma \vdash x: A \rrbracket=\pi_{x} ; \varepsilon \quad(\text { for } x:: A \in \Gamma) \\
& \llbracket \Gamma \vdash \lambda X . e: A \rightarrow B \rrbracket=\lambda_{X} \llbracket \Gamma, X: A \vdash e: B \rrbracket \\
& \llbracket \Gamma \vdash e_{1} e_{2}: B \rrbracket=\left\langle\llbracket \Gamma \vdash e_{1}: A \rightarrow B \rrbracket, \llbracket \Gamma \vdash e_{2}: A \rrbracket\right\rangle ; \text { eval } \\
& \llbracket \Gamma \vdash\left(e_{1}, e_{2}\right): A_{1} \times A_{2} \rrbracket=\left\langle\llbracket \Gamma \vdash e_{1}: A_{1} \rrbracket, \llbracket \Gamma \vdash e_{2}: A_{2} \rrbracket\right\rangle \\
& \llbracket \Gamma \vdash \pi_{i} e: A_{i} \rrbracket=\llbracket \Gamma \vdash e: A_{1} \times A_{2} \rrbracket ; \pi_{i} \\
& \llbracket \Gamma \vdash \mathrm{in}_{\mathrm{i}} e: A_{1}+A_{2} \rrbracket=\llbracket \Gamma \vdash e: A_{i} \rrbracket ; \mathrm{in}_{\mathrm{i}} \\
& \llbracket \Gamma \vdash \text { case } e \text { of }\left(\mathrm{in}_{\mathrm{i}} \mathrm{X}_{\mathrm{i}} \rightarrow \mathrm{f}_{\mathrm{i}}\right)_{\mathrm{i}}: \mathrm{B} \rrbracket=\left\langle\mathrm{id}_{\llbracket \Gamma \rrbracket}, \llbracket \Gamma \vdash e: \mathrm{A}_{1}+\mathrm{A}_{2} \rrbracket\right\rangle \\
& \text {; } \text { dist }_{+}^{\times} \\
& ;\left[\llbracket \Gamma, X_{i}: A_{i} \vdash f_{i}: B \rrbracket\right]_{i \in\{1,2\}} \\
& \llbracket \Gamma \vdash[e]: \square A \rrbracket=\operatorname{box}_{\Gamma}(\llbracket\lceil\Gamma\rceil \vdash e: A \rrbracket) \\
& \llbracket \Gamma \vdash \text { let }[x]=e \text { in } f: B \rrbracket=\left\langle\mathrm{id}_{\llbracket \Gamma \rrbracket}, \llbracket \Gamma \vdash e: \square A \rrbracket\right\rangle ; \llbracket \Gamma, x:: A \vdash \mathrm{f}: \mathrm{B} \rrbracket \\
& \llbracket \Gamma \vdash \perp: \mathrm{L} \rrbracket=\langle \rangle ; \mathrm{join}_{0}^{\llbracket \mathrm{L} \rrbracket} \\
& \llbracket \Gamma \vdash e \vee f: L \rrbracket=\langle\llbracket \Gamma \vdash e: L \rrbracket, \llbracket \Gamma \vdash \mathrm{f}: \mathrm{L} \rrbracket\rangle ; \mathrm{join}_{2}^{\llbracket L \rrbracket} \\
& \llbracket \Gamma \vdash \text { empty? } e: 1+1 \rrbracket=\text { box }_{\Gamma}(\llbracket\lceil\Gamma\rceil \vdash e:\{1\} \rrbracket) \text {; isEmpty } \\
& \llbracket \Gamma \vdash \text { split } e: \square A+\square \mathrm{B} \rrbracket=\llbracket \Gamma \vdash e: \square(A+B) \rrbracket ; \text { dist }_{+}^{\square} \\
& \llbracket \Gamma \vdash e_{1}=e_{2}: \operatorname{bool} \rrbracket=\left\langle\operatorname{box}_{\Gamma}\left(\llbracket\lceil\Gamma\rceil \vdash e_{1}: \underset{E q}{A} \rrbracket\right), \operatorname{box}_{\Gamma}\left(\llbracket\lceil\Gamma\rceil \vdash e_{2}: A_{\varepsilon q}^{A} \rrbracket\right)\right\rangle ; \text { eq }
\end{aligned}
$$

$$
\begin{aligned}
& \llbracket \Gamma \vdash \text { for }(x \in e) f: L \rrbracket=\left\langle\operatorname{id}_{\llbracket \Gamma \rrbracket}, \llbracket \Gamma \vdash e:\left\{{ }_{\varepsilon Q}^{A}\right\rceil \rrbracket\right\rangle ; \operatorname{collect}(\llbracket \Gamma, x:: \underset{\in Q}{A} \vdash f: L \rrbracket) \\
& \text { dist }_{+}^{\times}: A \times\left(B_{1}+B_{2}\right) \rightarrow\left(A \times B_{1}\right)+\left(A \times B_{2}\right) \quad \text { box }_{\Gamma}: \llbracket\lceil\Gamma\rceil \vdash A \rrbracket \rightarrow \llbracket \Gamma \vdash \square A \rrbracket \\
& \operatorname{dist}_{+}^{\times}=\left\langle\pi_{2} ;\left[\lambda\left(\left\langle\pi_{2}, \pi_{1}\right\rangle ; \mathrm{in}_{i}\right)\right]_{i}, \pi_{1}\right\rangle ; \text { eval } \quad \operatorname{box}_{\Gamma}(f)=\left\langle\pi_{x} ; \delta\right\rangle_{x:: A \in \Gamma} ; \operatorname{dist}_{\square}^{\times} ; \square(f)
\end{aligned}
$$

## ADDITIONAL SYNTAX

$$
\begin{aligned}
& \operatorname{iter}_{\mathcal{A}}(v, e, f) \\
& \text { values } v, u \quad:=\lambda x . e|()|(v, u)\left|\operatorname{in}_{i} v\right|\left\{v_{i}\right\}_{i} \mid[v] \\
& \text { evaluation contexts } \mathrm{E} \quad:=\mathrm{O}|\mathrm{Ee}| v \mathrm{E}|(\mathrm{E}, e)|(v, \mathrm{E}) \mid \pi_{i} \mathrm{E} \\
& \mathrm{in}_{i} \mathrm{E} \mid \text { case } \mathrm{E} \text { of }\left(\mathrm{in}_{\mathrm{i}} x_{i} \rightarrow e_{i}\right)_{i} \\
& {[E] \mid \text { let }[x]=E \text { in } e} \\
& \{\vec{v}, \mathrm{E}, \vec{e}\}\left|\mathrm{E} \vee_{\mathrm{L}} \mathrm{e}\right| v \vee_{\mathrm{L}} \mathrm{E} \mid \text { for }_{\mathrm{L}}(x \in \mathrm{E}) \mathrm{e} \\
& \mathrm{E}={ }_{A} \mathrm{e}\left|v={ }_{\mathrm{A}} \mathrm{E}\right| \text { split } \mathrm{E} \mid \text { empty? } \mathrm{E} \\
& \operatorname{fix}_{\mathrm{FI} \mathrm{~L}} \mathrm{E}\left|\operatorname{iter}_{A}(v, \mathrm{E}, \mathrm{f})\right| \operatorname{iter}_{A}(v, u, \mathrm{E}) \\
& () \leqslant(): 1 \quad \frac{v_{1} \leqslant u_{1}: \underset{\varepsilon Q}{A} \quad v_{2} \leqslant u_{2}: \underset{\varepsilon Q}{B}}{\left(v_{1}, v_{2}\right) \leqslant\left(u_{1}, u_{2}\right)::_{\varepsilon Q}^{A} \times \underset{\varepsilon Q}{B}} \quad \frac{v \leqslant u: A_{i}}{\operatorname{in}_{i} v \leqslant \operatorname{in}_{i} u: A_{1}+A_{2}}
\end{aligned}
$$

## $\beta$ reductions

$$
\begin{array}{rlrl}
(\lambda x . e) v & \longmapsto e\{x \mapsto v\} & \text { let }[x]=[v] \text { in } e & \longmapsto e\{x \mapsto v\} \\
\pi_{i}\left(v_{1}, v_{2}\right) & \longmapsto v_{i} \\
\text { for }_{L}(x \in\{ \}) e & \longmapsto \perp_{L} \\
\text { for }_{L}(x \in\{\vec{u}, v\}) e & \longmapsto\left(\text { for }_{L}(x \in\{\vec{u}\}) e\right) \vee_{L}(e\{x \mapsto v\})
\end{array}
$$

## OTHER REDUCTIONS

$$
\begin{aligned}
& \perp_{1} \longmapsto() \quad() \vee_{1}() \longmapsto() \\
& \perp_{\{A\}} \longmapsto\{ \} \quad\{\vec{v}\} \vee_{\{A\}}\{\vec{u}\} \longmapsto\{\vec{v}, \vec{u}\} \\
& \perp_{L \times M} \longmapsto\left(\perp_{L}, \perp_{M}\right) \quad\left(v_{1}, v_{2}\right) \vee_{L \times M}\left(u_{1}, u_{2}\right) \longmapsto\left(v_{1} \vee_{L} u_{1}, v_{2} \vee_{M} u_{2}\right) \\
& \text { empty? }\left\} \longmapsto \operatorname{in}_{1}() \quad \text { empty? }\{v, \vec{u}\} \longmapsto \operatorname{in}_{2}()\right. \\
& \text { split }\left[\mathrm{in}_{i} \nu\right] \longmapsto \mathrm{in}_{\mathrm{i}}[v] \\
& v=\underset{\in R}{A} u \longmapsto \begin{cases}\text { true } & \text { if } v=u: \underset{\varepsilon \in}{A} \\
\text { false } & \text { otherwise }\end{cases} \\
& \underset{\mathrm{fix}}{ }[v] \longmapsto \operatorname{iter}_{\mathrm{F} \mid \mathrm{X}}\left(v, \perp_{\mathrm{F} \mid \mathrm{X}}, v \underset{\mathrm{FIX}}{ }\right)
\end{aligned}
$$

FIGURE 2.5 Operational semantics

In our rules for $e \longmapsto e^{\prime}$ where $e$ is an iter expression we make use of a decidable ordering test on values, $v \leqslant u:{ }_{\mathrm{EQ}}^{\mathrm{A}}$, and a corresponding equality test $v=u: \underset{\varepsilon}{\mathrm{E}}$. We define these using inference rules, but they are easily seen to be decidable by induction on $A_{\varepsilon}$.

Our implementation strategy for fix $f$ is straightforward: starting from $\perp$, iteratively apply $f$ until quiescence. We introduce the form iter ( $f, e_{1}, e_{2}$ ) to represent these intermediate iterative steps. The intention is that $e_{1}, e_{2}$ shall be successive iterations of $f$, with $e_{2}=f e_{1}$. The fixed point expression fix $f$, after evaluating $f$, steps to iter $(f, \perp, f e$ ), which kicks off the first two iterations. Once these have reduced to values, $\operatorname{iter}\left(f, u_{1}, u_{2}\right)$ tests $u_{1}=u_{2}$ to determine if a fixed point has been reached. If so, its value $u_{1}$ is returned; otherwise we step to iter $\left(f, u_{2}, f u_{2}\right)$ to evaluate the next iteration, and so on.

Observe that values don't step and evaluation is deterministic:
Lemma 1 (Values don't step). If $v$ is a value, there is no $e$ such that $v \longmapsto e$.
Proof. The left hand side of each reduction rule can never be a value. This is easily verified by inspection for the rules in figure 2.5; and for eval context we can see by the definition of evaluation contexts $E$ that filling a hole with a non-value always produces a non-value.

Lemma 2 (Determinism). If $e \longmapsto e_{1}^{\prime}$ and $e \longmapsto e_{2}^{\prime}$ then $e_{1}^{\prime}=e_{2}^{\prime}$; thus inductively, since values don't step, if $e \longmapsto^{*} v$ and $e \longmapsto^{*} u$ then $v=u$.

Proof. The left-hand sides of all reduction rules $e \longmapsto e^{\prime}$ are disjoint; there is no term to which two distinct reduction rules could apply. This applies inductively to eval context because decompositions of a term into an evaluation context and a reducible subterm are unique.

### 2.4.1 A logical relation for termination

To prove that all well-typed terms terminate according to our operational semantics, we use a logical relations argument. As a matter of notation, we will let $v, u, w$ range over values; $\mathrm{a}, \mathrm{b}, \mathrm{c}$ range over closed terms; and $\gamma, \sigma$ range over closing substitutions.

Our guiding intuition is that since we need an order structure in our denotational semantics to prove the definedness of fixed points, we likewise need an order structure on our syntax to prove the termination of fixed points. To this end we interpret each type $A$ as a partial preorder, $x \prec y: A$. A partial preorder is a relation which is transitive and partially reflexive, that is, $x \prec y \Longrightarrow x \prec x \wedge y \prec y$. While reflexivity may be glossed as "every element is related to itself", partial reflexivity glosses as "if an element is related to anything, it is related to itself"; in other words, unlike reflexivity, it permits some elements to be "outside the relation" and unrelated to anything, even themselves. Any partial preorder $x \prec y$ gives rise to a symmetric, transitive relation $x \equiv y \Longleftrightarrow x \prec y \wedge y \prec x .{ }^{7}$

In fact we define a mutually inductive collection of partial preorders: on values $v \prec u: A$, an extension to closed terms $\mathrm{a} \prec \mathrm{b}: A$, on closing substitutions $\gamma \prec \sigma: \Gamma$, on open terms

7 Symmetric, transitive relations are also known as partial equivalence relations (PERs). Moreover, letting $[x]_{\equiv}$ denote the equivalence class of $x$ (defined only when $x \prec x$ ), the relation $[x]_{\equiv} \leqslant[y]_{\equiv} \Longleftrightarrow x \prec y$ is a partial order over these equivalence classes; so our approach may also be considered to interpret types as PERs equipped with partial orders on their equivalence classes.
$e \prec f: \Gamma \vdash A$, and on open terms paired with closing substitutions $\gamma_{1}, e_{1} \prec \gamma_{2}, e_{2}: \Gamma, A$. The rules for the value-relation are:

$$
\begin{aligned}
& \overline{() \prec(): 1} \quad \frac{v_{1} \prec u_{1}: A \quad v_{2} \prec u_{2}: B}{\left(v_{1}, v_{2}\right) \prec\left(u_{1}, u_{2}\right): A \times B} \quad \frac{v \prec u: A_{i}}{\operatorname{in}_{i} v \prec \mathrm{in}_{i} u: A_{1}+A_{2}}
\end{aligned}
$$

Note that LR FN depends on the relation for open terms, making this definition mutually inductive. The second premise of LR SET may seem strange but it is necessary to ensure partial reflexivity. We extend this value-relation to closed terms:

$$
\mathrm{a} \prec \mathrm{~b}: A \Longleftrightarrow(\exists v, u) \mathrm{a} \longmapsto \longmapsto^{*} v \wedge \mathrm{~b} \longmapsto{ }^{*} u \wedge v \prec u: A
$$

Note that if $a, b$ are values, this definition coincides with the relation on values, since values do not step; this justifies using the same notation for the relation on values and closed terms. We extend this relation to closing substitutions pointwise, noting that discrete hypotheses are required to be equivalent:

$$
\gamma \prec \sigma: \Gamma \Longleftrightarrow\left((\forall X: A \in \Gamma) \gamma_{x} \prec \sigma_{x}: A\right) \wedge\left((\forall x:: A \in \Gamma) \gamma_{x} \equiv \sigma_{x}: A\right)
$$

Finally, we extend the relation to open terms, which involves an auxiliary relation on pairs of terms and closing substitutions:

$$
\begin{aligned}
e_{1} \prec e_{2}: \Gamma \vdash A & \Longleftrightarrow\left(\forall \gamma_{1} \prec \gamma_{2}: \Gamma\right) \gamma_{1}, e_{1} \prec \gamma_{2}, e_{2}: \Gamma, A \\
\gamma_{1}, e_{1} \prec \gamma_{2}, e_{2}: \Gamma, A & \Longleftrightarrow(\forall i=1,2) \gamma_{i}\left(e_{1}\right) \prec \gamma_{i}\left(e_{2}\right): A \wedge \gamma_{1}\left(e_{i}\right) \prec \gamma_{2}\left(e_{i}\right): A
\end{aligned}
$$

Note that $\gamma_{1}, e_{1} \prec \gamma_{2}, e_{2}: \Gamma, \mathcal{A}$ may be seen as a transitive square:


This ensures partial reflexivity; if we replace $e_{1}$ with $e_{2}$ or vice-versa this square collapses to one of its sides. If we had instead only required the diagonal, $\gamma_{1}\left(e_{1}\right) \prec \gamma_{2}\left(e_{2}\right): A$, we could not derive $\gamma_{1}\left(e_{1}\right) \prec \gamma_{2}\left(e_{1}\right)$ : A (or the same for $e_{2}$ ) as required by partial reflexivity.

Theorem 3 (Fundamental theorem). If $\Gamma \vdash e: A$ then $e \prec e: \Gamma \vdash A$.
Termination of well-typed programs follows as a corollary by unrolling definitions:
Theorem 4 (Termination). Every closed, well-typed program $\varepsilon \vdash \mathrm{a}: \mathcal{A}$ terminates.

Proof.

$$
\begin{aligned}
& \varepsilon \vdash a: A \\
\Longrightarrow & a \prec a: \varepsilon \vdash A \\
\Longrightarrow & \left(\forall \gamma_{1} \prec \gamma_{2}: \varepsilon\right) \gamma_{1}, a \prec \gamma_{2}, a: \Gamma, A \\
\Longrightarrow & (), a \prec(), a: \varepsilon, A \\
\Longrightarrow & a \prec a: A \\
\Longrightarrow & (\exists v, u) a \longmapsto^{*} v \wedge a \longmapsto^{*} u \wedge v \prec u: A \\
\Longrightarrow & (\exists v) a \longmapsto^{*} v
\end{aligned}
$$

Fundamental theorem
expand the definition
since ()$\prec(): \varepsilon$ vacuously expand the definition and simplify expand the definition simplify

The proof of the fundamental theorem itself proceeds by induction on $\Gamma \vdash e: A$. The key case is the fixed point rule, whose proof is a syntactic version of the proof of the existence of least fixed points in the denotational semantics. We give the proof of the fundamental theorem at the end of this section; to build up to it we must first develop several auxiliary definitions and lemmas.

### 2.4.2 Metatheory of the logical relation

First, any partial preorder over a set $S$ gives rise to a subset $\{x \in S \mid x \prec x\}$ over which $\prec$ is reflexive and thus a true preorder. It will be convenient to apply this point of view to our logical relations:

Definition 5 (Good terms). We define the following preordered sets of "good" terms:

$$
\begin{aligned}
\operatorname{Ok}_{v}(A) & =\{v \mid v \prec v: A\} \\
\operatorname{Ok}_{c}(A) & =\{\mathrm{a} \mid \mathrm{a} \prec \mathrm{a}: A\} \\
\operatorname{Ok}(\Gamma \vdash A) & =\{e \mid e \prec e: \Gamma \vdash A\}
\end{aligned}
$$

the good values
the good closed terms
the good open terms

We preorder these by the corresponding logical relation, so $v \leqslant u: \mathrm{Ok}_{v}(A) \Longleftrightarrow v \prec u: A$, etc.

Lemma 6 (Closed term evaluation map). There exists a monotone map value ${ }_{A}: \mathrm{Ok}_{\mathrm{c}}(A) \rightarrow$ $\mathrm{Ok}_{v}(\mathcal{A})$ such that value $\mathrm{a}=v$ if and only if $\mathrm{a} \longmapsto{ }^{*} v$.

Proof. The definition of $a \prec a: A$ shows that every good closed term evaluates to some good value. Determinism shows this value is unique, so we can name it value a. And given $\mathrm{a} \prec \mathrm{b}: A$, applying its definition and this uniqueness shows that value $\mathrm{a} \prec$ value $\mathrm{b}: A$, showing monotonicity.

Lemma 7 (Closed term application map). If $a_{1} \prec a_{2}: A \rightarrow B$ and $b_{1} \prec b_{2}: A$ then $a_{1} b_{1} \prec a_{2} b_{2}$ : B. Equivalently, there exists a monotone map apply $A, B: \mathrm{Ok}_{C}(A \rightarrow B) \times$ $\mathrm{Ok}_{\mathrm{c}}(A) \rightarrow \mathrm{Ok}_{v}(B)$ such that apply $(\mathrm{a}, \mathrm{b})=$ value $(\mathrm{a} b)=v$ if and only if $a b \longmapsto^{*} v$.

Proof. Suppose $a_{1} \prec a_{2}: A \rightarrow B$ and $b_{1} \prec b_{2}: A$. Unrolling these assumptions, we have $e_{1}, e_{2}, u_{1}, u_{2}$ satisfying:

$$
\begin{array}{lll}
a_{1} \longmapsto{ }^{*} \lambda X . e_{1} & a_{2} \longmapsto{ }^{*} \lambda X . e_{2} & e_{1} \prec e_{2}:(X: A) \vdash B \\
b_{1} \longmapsto{ }^{*} u_{1} & b_{2} \longmapsto{ }^{*} u_{2} & u_{1} \prec u_{2}: A
\end{array}
$$

From $u_{1} \prec u_{2}$ : A we have $\left(X \mapsto u_{1}\right) \prec\left(X \mapsto u_{2}\right):(X: A)$, and applying this to $e_{1} \prec e_{2}$ : $(X: A) \vdash B$ we have a transitive square:

$$
\begin{array}{rll}
e_{1}\left\{X \mapsto u_{1}\right\} & \prec & e_{2}\left\{X \mapsto u_{1}\right\} \\
\curlywedge & & \curlywedge \\
e_{1}\left\{X \mapsto u_{2}\right\} & \prec & e_{2}\left\{X \mapsto u_{2}\right\}
\end{array}
$$

Taking the diagonal of this square, we have $e_{1}\left\{X \mapsto u_{1}\right\} \prec e_{2}\left\{X \mapsto u_{2}\right\}$ : B and therefore $v_{1}, v_{2}$ such that $e_{i}\left\{X \mapsto u_{i}\right\} \longmapsto{ }^{*} v_{i}$ and $v_{1} \prec v_{2}: B$. Thus for $i \in\{1,2\}$ we have:

$$
a_{i} b_{i} \longmapsto{ }^{*}\left(\lambda X . e_{i}\right) b_{i} \longmapsto{ }^{*}\left(\lambda X . e_{i}\right) u_{i} \longmapsto e_{i}\left\{X \mapsto u_{i}\right\} \longmapsto{ }^{*} v_{i}
$$

and $v_{1} \prec v_{2}: \mathrm{B}$ as desired.
Lemma 8 (Closed term pairing). If $a_{1} \prec a_{2}$ : $A$ and $b_{1} \prec b_{2}$ : $B$ then $\left(a_{1}, b_{1}\right) \prec\left(a_{2}, b_{2}\right)$ : $A \times B$.

Proof. Applying our assumptions' definitions we have $v_{1}, v_{2}, u_{1}, u_{2}$ such that:

$$
\begin{array}{lll}
\mathrm{a}_{1} \longmapsto{ }^{*} v_{1} & \mathrm{a}_{2} \longmapsto{ }^{*} v_{2} & v_{1} \prec v_{2}: A \\
\mathrm{~b}_{1} \longmapsto{ }^{*} \mathrm{u}_{1} & \mathrm{~b}_{2} \longmapsto{ }^{*} u_{2} & \mathrm{u}_{1} \prec \mathrm{u}_{2}: B
\end{array}
$$

From this we have:

$$
\left(a_{1}, b_{1}\right) \longmapsto{ }^{*}\left(v_{1}, b_{1}\right) \longmapsto{ }^{*}\left(v_{1}, u_{1}\right) \quad\left(a_{2}, b_{2}\right) \longmapsto{ }^{*}\left(v_{2}, b_{2}\right) \longmapsto \longmapsto^{*}\left(v_{2}, u_{2}\right)
$$

And $\left(v_{1}, u_{1}\right) \prec\left(v_{2}, u_{2}\right): A \times B$ because $v_{1} \prec v_{2}: A$ and $u_{1} \prec u_{2}: B$, which is what we wished to show.

Lemma 9 (Closure under stepping). $\prec$ is closed under $\longmapsto^{*}$; that is, if $a \prec b: A$ and $\left(a^{\prime} \longmapsto{ }^{*} a\right) \vee\left(a \longmapsto^{*} a^{\prime}\right)$ and $\left(b^{\prime} \longmapsto{ }^{*} b\right) \vee\left(b \longmapsto{ }^{*} b^{\prime}\right)$, then $a^{\prime} \prec b^{\prime}: A$.

Proof. $\mathrm{a} \longmapsto^{*} v$ and $\mathrm{b} \longmapsto^{*} u$ such that $v \prec u: A$ so by determinism $\mathrm{a}^{\prime} \longmapsto^{*} v$ and $b^{\prime} \longmapsto{ }^{*} u$, thus $a^{\prime} \prec b^{\prime}: A$.

Lemma 10 (First-order agreement on values). If $v \in \operatorname{Ok}_{v}(\underset{\varepsilon Q}{A})$ then $\varepsilon \vdash v: \underset{\varepsilon q}{A}$ and moreover


$$
v \prec u: \underset{E Q}{A} \Longleftrightarrow \llbracket v \rrbracket \leqslant \llbracket u \rrbracket: \llbracket \llbracket A_{\varepsilon Q}^{A} \rrbracket \Longleftrightarrow v \leqslant u: \underset{E Q}{A}
$$

(To be precise, by $\llbracket v \rrbracket: \llbracket \llbracket{ }_{\varepsilon q}^{A} \rrbracket$ we mean the map $\llbracket \varepsilon \vdash v: \underset{\varepsilon Q}{A} \rrbracket: \operatorname{Poset}(\llbracket \varepsilon \rrbracket, \llbracket \underset{\varepsilon q}{A} \rrbracket)$ applied to the empty environment () : $\llbracket \varepsilon \rrbracket$.)
 (e.g. for ${\underset{E Q}{1}}_{A_{1}} \times A_{E_{2}}$ we find that $v=\left(u_{1}, u_{2}\right)$ for some $u_{1}, u_{2} \in \operatorname{OK}\left(\underset{\mathrm{EQ}}{ }\left(A_{1}\right)\right)$ followed by applying our inductive hypotheses and the following typing rules (here specialized to $\Gamma=\varepsilon$ ):

$$
\begin{array}{llll}
\text { UNIT } & \begin{array}{l}
\text { PAIR } \\
\varepsilon \vdash(): 1
\end{array} & \frac{\left(\varepsilon \vdash e_{i}: A_{i}\right)_{i}}{\varepsilon \vdash\left(e_{1}, e_{2}\right): A_{1} \times A_{2}} & \frac{\text { INJ }}{\varepsilon \vdash i n_{i} e: A_{1}+A_{2}}
\end{array} \quad \frac{\text { SET }}{\varepsilon \vdash}
$$

As for equivalence of the orderings, observe that:

$$
\begin{aligned}
& \llbracket() \rrbracket=(): \llbracket 1 \rrbracket \\
& \llbracket \mathrm{in}_{\mathrm{i}} v \rrbracket=\mathrm{in}_{\mathrm{i}} \llbracket v \rrbracket: \llbracket \mathcal{A}_{\mathrm{E}}^{\mathrm{A}}+\underset{\mathrm{E}}{\mathrm{~B}} \rrbracket \\
& \llbracket(v, u) \rrbracket=(\llbracket v \rrbracket, \llbracket u \rrbracket): \llbracket A_{\varepsilon} \mathcal{A} \times \underset{\varepsilon_{Q}}{B} \rrbracket \\
& \llbracket\left\{v_{i}\right\}_{i} \rrbracket=\left\{\llbracket v_{i} \rrbracket\right\}_{i}: \llbracket\left\{\mathcal{A R}_{\varepsilon}\right\} \rrbracket
\end{aligned}
$$

and therefore:

$$
\begin{aligned}
& \llbracket() \rrbracket \leqslant \llbracket() \rrbracket: \llbracket 1 \rrbracket \Longleftrightarrow \top
\end{aligned}
$$

In each case, these coincide (after applying our inductive hypothesis) with the rules defining
 $(\forall j) u_{j} \prec u_{j}: A_{E Q} ;$ but this is satisfied by the assumption $\left\{u_{j}\right\}_{j} \in O k_{v}\left(\left\{\left\{_{\varepsilon Q}\right\}\right)\right.$. Thus inductively all three preorders coincide on good values of equality types.

Lemma 11 (Bottom is bottom). $\perp_{\mathrm{L}} \in \mathrm{Ok}_{\mathrm{C}}(\mathrm{L})$ and $\left(\forall \mathrm{a} \in \mathrm{Ok}_{\mathrm{C}}(\mathrm{L})\right) \perp_{\mathrm{L}} \prec \mathrm{a}: \mathrm{L}$.
Proof. By induction on L:
Case 1: Follows trivially from $\perp_{1} \longmapsto()$ and the relation at 1 .
Case $\mathrm{L}_{1} \times \mathrm{L}_{2}$ : We have $\mathrm{a} \longmapsto \longmapsto^{*}\left(v_{1}, v_{2}\right)$ for good $v_{i}$ and by IH we have $\perp_{\mathrm{L}_{i}} \prec \nu_{i}: \mathrm{L}_{\mathrm{i}}$. Thus by closed term pairing $\left(\perp_{L_{1}}, \perp_{\mathrm{L}_{2}}\right) \prec\left(v_{1}, v_{2}\right): \mathrm{L}_{1} \times \mathrm{L}_{2}$ and since $\perp_{\mathrm{L}_{1} \times \mathrm{L}_{2}} \longmapsto\left(\perp_{\mathrm{L}_{1}}, \perp_{\mathrm{L}_{2}}\right)$ by closure under stepping we have what we desire.
 from LR SET; the first premise is vacuous and the second follows from goodness of $v$.

Lemma 12 (Join is join). $a V_{L} b$ is the least upper bound of $a, b \in \operatorname{Ok}_{C}(L)$ with respect to the logical relation. That is, $a \prec a \vee_{L} b: L$ and $b \prec a \vee_{L} b: L$ and for any $c$ such that $a \prec c: L$ and $b \prec c:$ L we have $a V_{L} b \prec c: L$.

Proof. By closure under stepping it suffices to show the same for only good values. For this it suffices to show that $\llbracket v a l u e\left(v \vee_{\mathrm{L}} u\right) \rrbracket=\llbracket v \rrbracket \vee \llbracket u \rrbracket$ because by lemma 10 the semantic ordering and the logical relation ordering agree, so a least upper bound in one is a least upper bound in the other. We show this by induction on L:

Case 1: Follows from ()$\vee_{\mathrm{L}}() \longmapsto{ }^{*}()$ and the trivial order on 1.
Case $\mathrm{L} \times \mathrm{M}$ : Then we have

$$
\begin{array}{rlrl} 
& \llbracket \text { value }\left(\left(v_{1}, v_{2}\right) \vee_{\mathrm{L} \times \mathrm{M}}\left(u_{1}, u_{2}\right)\right) \rrbracket & \\
= & \llbracket\left(\text { value }\left(v_{1} \vee_{\mathrm{L}} u_{1}\right) \text {, value }\left(v_{2} \vee_{\mathrm{M}} u_{2}\right)\right) \rrbracket & & \text { calculation } \\
= & \left(\llbracket \text { value }\left(v_{1} \vee_{\mathrm{L}} u_{1}\right) \rrbracket, \llbracket \text { value }\left(v_{2} \vee_{\mathrm{M}} u_{2}\right) \rrbracket\right) & & \text { calculation } \\
= & \left(\llbracket v_{1} \rrbracket \vee \llbracket u_{1} \rrbracket, \llbracket v_{2} \rrbracket \vee \llbracket u_{2} \rrbracket\right) & & \text { inductive hy } \\
= & \left(\llbracket v_{1} \rrbracket, \llbracket v_{2} \rrbracket\right) \vee\left(\llbracket u_{1} \rrbracket, \llbracket u_{2} \rrbracket\right) & & \text { join in prod } \\
= & \llbracket\left(v_{1}, v_{2}\right) \rrbracket \vee \llbracket\left(u_{1}, u_{2}\right) \rrbracket & & \text { calculation }
\end{array}
$$ calculation inductive hypothesis join in product semilattice calculation

Case $\left\{\begin{array}{l}\text { feq } \\ \text { A }\end{array}\right\}$ : We have

$$
\begin{aligned}
\llbracket \text { value }\left(\{\vec{v}\} \vee_{\left\{\mathcal{A A}^{\mathrm{EQ}}\right\}}\{\vec{u}\}\right) \rrbracket & =\llbracket\{\vec{v}, \vec{u}\} \rrbracket & & \text { calculation } \\
& =\left\{\llbracket v_{i} \rrbracket\right\}_{i} \cup\left\{\llbracket u_{j} \rrbracket\right\}_{j} & & \text { calculation } \\
& =\llbracket\{\vec{v}\} \rrbracket \vee \llbracket\{\vec{u}\} \rrbracket \rrbracket & & \text { calculation }
\end{aligned}
$$

Lemma 13 (Discrete contexts make terms equivalent). If $e \prec f:\lceil\Gamma\rceil \vdash A$ and $\gamma_{1} \prec \gamma_{2}: \Gamma$ then $\gamma_{1}(e) \equiv \gamma_{2}(f): A$.

Proof. From $\gamma_{1} \prec \gamma_{2}: \Gamma$ we have $\gamma_{1} \equiv \gamma_{2}:\lceil\Gamma\rceil$ because $\lceil\Gamma\rceil$ restricts to only discrete hypotheses $x:: A \in \Gamma$ for which we know $\gamma_{1}(x) \equiv \gamma_{2}(x): A$. Thus applying $e \prec f:\lceil\Gamma\rceil \vdash A$ we have $\gamma_{1}(e) \equiv \gamma_{2}(f): A$ as desired.

### 2.4.3 Proof of the fundamental theorem

We now have the groundwork to prove the fundamental theorem, starting with the crucial case of fixed point expressions fix $e$.

Theorem 3 (Fundamental theorem). If $\Gamma \vdash e: A$ then $e \prec e: \Gamma \vdash A$.
Proof. Unrolling the definition of the logical relation, we may assume $\gamma_{1} \prec \gamma_{2}$ : $\Gamma$ and wish to show from this that $\gamma_{1}(e) \prec \gamma_{2}(e): A$. We do this by induction on $\Gamma \vdash e: A$.

Case $\frac{\Gamma \vdash e: \square(\underset{\mathrm{flx}}{\mathrm{L}} \rightarrow \underset{\mathrm{fx}}{\mathrm{L}})}{\Gamma \vdash f \mathrm{x} e \mathrm{e}: \mathrm{H}}$. We wish to show that

By our inductive hypothesis we have $\gamma_{1}(e) \prec \gamma_{2}(e): \square(\underset{f \mid x}{ } \rightarrow \underset{f \mid x}{\mathrm{f}})$; applying this we have $\nu_{1}, \nu_{2}$ such that $\gamma_{i}(e) \longmapsto \longmapsto^{*}\left[\nu_{i}\right]$ and $\nu_{1} \equiv v_{2}: \underset{\text { fix }}{\mathrm{L}} \rightarrow \underset{\mathrm{fX}}{\mathrm{L}}$. Thus for $i \in\{1,2\}$ :

Let $\boldsymbol{f}_{\mathfrak{i}}(u)=$ apply $\left(v_{i}, u\right)$ for brevity. Then applying lemmas 6 and 7 we have:

$$
\operatorname{iter}_{\mathrm{FIX}}\left(v_{i}, \perp_{\mathrm{FIX}}, v_{i} \perp_{\mathrm{LIX}}\right) \longmapsto{ }^{*} \operatorname{iter}_{\mathrm{FIX}}\left(v_{i}, \text { value } \perp_{\mathrm{LIX}}, f_{i}\left(\text { value } \perp_{\mathrm{LIX}}\right)\right)
$$

By bottom-is-bottom we have value $\perp_{\mathrm{t} \mid \mathrm{L}} \prec \mathrm{f}_{\mathfrak{i}}\left(\perp_{\mathrm{L}}\right)$ : $:$. To understand the way evaluation will proceed from here, consider the generalized situation $\operatorname{iter}_{\substack{\text { FIX }}}\left(v_{i}, u, f_{\mathfrak{i}}(u)\right)$ where $u \prec f_{i}(u):$ L. . This steps like so:

$$
\operatorname{iter}_{\mathrm{FI}}\left(v_{i}, u, f_{i}(u)\right) \longmapsto \longmapsto^{*} \begin{cases}u & \text { if } u=f_{i}(u): \underset{E Q}{A} \\ \operatorname{iter}_{\mathcal{E Q}}\left(v, f_{i}(u), f_{i}\left(f_{i}(u)\right)\right) & \text { otherwise }\end{cases}
$$

Starting with $u=\perp_{\text {Lix }}$, this calculates the sequence $u, f_{i}(u), f_{i}^{2}(u), f_{i}^{3}(u), \ldots$ until the first $k$ such that $f_{i}^{k}(u)=f_{i}^{k+1}(u): \underset{f \mid X}{L}$ and returns $f_{i}^{k}(u)$. Note that $f_{i}: \operatorname{Ok}_{C}(\underset{F X}{ }) \rightarrow \operatorname{Ok}_{V}\left(\mathcal{L E X}^{L}\right)$ is monotone by lemma 7 and therefore this sequence ascends in the logical relation:
 by partial reflexivity, we have inductively that $f_{1}^{j}(u) \equiv f_{2}^{j}(u):$ L. By lemma 10 these also hold in the semantic order, so the denotations both ascend $\llbracket f_{i}^{\mathfrak{j}}(u) \rrbracket \leqslant \llbracket f_{i}^{j+1}(u) \rrbracket: \llbracket \mathbb{L x}_{\ddagger x} \rrbracket$
 there must be some $k$ such that $\llbracket f_{1}^{k}(u) \rrbracket=\llbracket f_{1}^{k+1}(u) \rrbracket$; and as the sequences for $f_{1}, f_{2}$ coincide, $\llbracket f_{1}^{k}(u) \rrbracket=\llbracket f_{1}^{k+1}(u) \rrbracket=\llbracket f_{2}^{k}(u) \rrbracket=\llbracket f_{2}^{k+1}(u) \rrbracket$. Applying lemma 10 again this shows $f_{i}^{k}(u)=f_{i}^{k+1}(u): L$, and therefore $f_{i}^{k}(u)$ for the least such $k$ is the value we terminate with; since $f_{1}^{\mathrm{k}}(u) \equiv f_{2}^{\mathrm{k}}(u): \mathrm{L}_{\mathrm{f} \times \mathrm{x}}$, applying closure under stepping we are done.

Cases $\frac{X: A \in \Gamma}{\Gamma \vdash X: A} \quad \frac{x:: A \in \Gamma}{\Gamma \vdash x: A}$. Follows directly from $\gamma_{1} \prec \gamma_{2}: \Gamma$.

Case $\frac{\Gamma, X: A \vdash e: B}{\Gamma \vdash \lambda X . e: A \rightarrow B}$. What we wish to show is equivalent to:

$$
\begin{aligned}
& \gamma_{1}(\lambda X . e) \prec \gamma_{2}(\lambda X . e): A \rightarrow B \\
\Longleftrightarrow & \lambda X \cdot \gamma_{1}(e) \prec \lambda X . \gamma_{2}(e): A \rightarrow B \\
\Longleftrightarrow & \gamma_{1}(e) \prec \gamma_{2}(e):(X: A) \vdash B
\end{aligned}
$$

For the last it suffices to assume $a_{1} \prec a_{2}: A$ and show the transitive square at the logical relation for B:

$$
\begin{array}{rlcc}
e\left\{\gamma_{1}, X \mapsto a_{1}\right\} & & e\left\{\gamma_{2}, X \mapsto a_{1}\right\} \\
\lambda & & \lambda \\
e\left\{\gamma_{1}, X \mapsto a_{2}\right\} & \prec & e\left\{\gamma_{2}, X \mapsto a_{2}\right\}
\end{array}
$$

By our IH we have $\left(\forall \sigma \prec \sigma^{\prime}: \Gamma, X: A\right) \sigma(e) \prec \sigma^{\prime}(e): B$, so it suffices to show the transitive square at the logical relation for $\Gamma, X: A$ :

$$
\begin{array}{ccc}
\left(\gamma_{1}, \mathrm{X} \mapsto \mathrm{a}_{1}\right) & \prec & \left(\gamma_{2}, \mathrm{X} \mapsto \mathrm{a}_{1}\right) \\
\curlywedge & & \curlywedge \\
\left(\gamma_{1}, \mathrm{X} \mapsto \mathrm{a}_{2}\right) & \prec & \left(\gamma_{2}, \mathrm{X} \mapsto \mathrm{a}_{2}\right)
\end{array}
$$

This holds by $\gamma_{1} \prec \gamma_{2}$ : $\Gamma$ and $a_{1} \prec a_{2}$ : A and the definition of the logical relation for contexts.

Case $\frac{\Gamma \vdash e: A \rightarrow B \quad \Gamma \vdash f: A}{\Gamma \vdash e f: B}$. We wish to show $\gamma_{1}(e) \gamma_{1}(f) \prec \gamma_{2}(e) \gamma_{2}(f):$ B. By IH we have $\gamma_{1}(e) \prec \gamma_{2}(e): A \rightarrow B$ and $\gamma_{1}(f) \prec \gamma_{2}(f): A$. What we wish to show then follows from lemma 7.

Case $\overline{\Gamma \vdash(): 1}$. Trivial.

Case $\frac{\left(\Gamma \vdash e_{i}: A_{i}\right)_{i}}{\Gamma \vdash\left(e_{1}, e_{2}\right): A_{1} \times A_{2}}$. Apply our inductive hypotheses to get $\gamma_{i}\left(e_{j}\right) \longmapsto \longmapsto^{*} v_{i, j}$ with $v_{1, \mathrm{j}} \prec \nu_{2, \mathrm{j}}: A_{\mathrm{j}}$; this shows $\left(v_{1,1}, v_{1,2}\right) \prec\left(v_{2,1}, v_{2,2}\right): A_{1} \times A_{2}$ and since $\gamma_{i}\left(\left(e_{1}, e_{2}\right)\right)=$ $\left(\gamma_{i}\left(e_{1}\right), \gamma_{i}\left(e_{2}\right)\right) \longmapsto{ }^{*}\left(\nu_{i, 1}, \gamma_{i}\left(e_{2}\right)\right) \longmapsto{ }^{*}\left(\nu_{i, 1}, \nu_{i, 2}\right)$ by closure under stepping we are done.

Case $\frac{\Gamma \vdash e: A_{1} \times A_{2}}{\Gamma \vdash \pi_{i} e: A_{i}}$. By IH we have $\gamma_{j}(e) \longmapsto \longmapsto^{*}\left(v_{1, j}, \nu_{2, j}\right)$ with $v_{i, 1} \prec v_{i, 2}: A_{i}$; thus we have $\gamma_{j}\left(\pi_{i} e\right)=\pi_{i} \gamma_{j}(e) \longmapsto{ }^{*} \pi_{i}\left(v_{1, j}, \nu_{2, j}\right) \longmapsto \nu_{i, j}$ and we are done.

Case $\frac{\Gamma \vdash e: A_{i}}{\Gamma \vdash i n_{i} e: A_{1}+A_{2}}$. By IH we have $\gamma_{j}(e) \longmapsto \longmapsto^{*} v_{j}$ for some $v_{1} \prec v_{2}: A_{i}$. Applying the definition of the LR we have $\mathrm{in}_{i} \nu_{1} \prec \mathrm{in}_{i} \nu_{2}: A_{1}+A_{2}$ and since $\gamma_{\mathrm{j}}\left(\mathrm{in}_{i} e\right)=$ $\mathrm{in}_{\mathrm{i}} \gamma_{\mathrm{j}}(e) \longmapsto{ }^{*} \mathrm{in}_{\mathrm{i}} \nu_{j}$ by closure under stepping we are done.

Case $\frac{\Gamma \vdash e: A_{1}+A_{2} \quad\left(\Gamma, X_{i}: A_{i} \vdash f_{i}: B\right)_{i}}{\Gamma \vdash \text { case } e \text { of }\left(i n_{i} X_{i} \rightarrow f_{i}\right)_{i}: B}$. By IH for $e$ and the LR for $A_{1}+A_{2}$ we have $\gamma_{j}(e) \longmapsto \longmapsto^{*} \operatorname{in}_{i} v_{j}$ for some $i$ and $v_{1} \prec v_{2}$ : $A_{i}$. Using this and our IH for $f_{i}$ we have $f_{i}\left\{\gamma_{1}, X_{i} \mapsto \nu_{1}\right\} \prec f_{i}\left\{\gamma_{2}, X_{i} \mapsto \nu_{2}\right\}$ : B. Then by calculation:

$$
\begin{aligned}
\gamma_{j}\left(\text { case } e \text { of }\left(\mathrm{in}_{i} X_{i} \rightarrow f_{i}\right)_{i}\right) & \longmapsto{ }^{*} \text { case } \text { in }_{i} v_{j} \text { of }\left(\mathrm{in}_{i} X_{i} \rightarrow \gamma_{j}\left(f_{i}\right)\right)_{i} \\
& \longmapsto f_{i}\left\{\gamma_{j}, X_{i} \mapsto v_{j}\right\}
\end{aligned}
$$

and by closure under stepping we are done.
Case $\frac{\lceil\Gamma\rceil \vdash e: A}{\Gamma \vdash[e]: \square A}$. By our IH and lemma 13 we have for some $v_{i}$ that $\gamma_{i}(e) \longmapsto{ }^{*} v_{i}$ with $v_{1} \equiv v_{2}: A$. Thus we have $\gamma_{i}([e])=\left[\gamma_{i}(e)\right] \longmapsto^{*}\left[v_{i}\right]$ and by the definition of the LR for $\square A$ we are done.

Case $\frac{\Gamma \vdash e: \square A \quad \Gamma, x:: A \vdash f: B}{\Gamma \vdash \operatorname{let}[\mathrm{x}]=\mathrm{e} \text { in } \mathrm{f}: \mathrm{B}}$. By our IH for e and the logical relation for $\square \mathrm{A}$ we
have $\gamma_{i}(e) \longmapsto \longmapsto^{*}\left[\nu_{i}\right]$ for some $v_{1} \equiv v_{2}: A$. Then using this and applying our IH for f we have $f\left\{\gamma_{i}, x \mapsto v_{i}\right\} \longmapsto{ }^{*} u_{i}$ for some $u_{1} \prec u_{2}: B$. Then we have:

$$
\text { let } \begin{aligned}
{[x]=\gamma_{i}(e) \text { in } \gamma_{i}(f) } & \longmapsto{ }^{*} \text { let }[x]=\left[v_{i}\right] \text { in } \gamma_{i}(f) \\
& \longmapsto f\left\{\gamma_{i}, x \mapsto v_{i}\right\} \\
& \longmapsto \mathfrak{u}_{i}
\end{aligned}
$$

And by closure under stepping we are done.

Case $\overline{\Gamma \vdash \perp: L}$. By bottom is bottom.

Case $\frac{\left(\Gamma \vdash e_{i}: L\right)_{i}}{\Gamma \vdash e_{1} \vee e_{2}: L}$. Applying our IH we have $\gamma_{1}\left(e_{i}\right) \prec \gamma_{2}\left(e_{i}\right): L$; since join is join we know in particular that $\vee$ is a monotone operator on good closed terms and therefore $\gamma_{1}\left(e_{1}\right) \vee \gamma_{1}\left(e_{2}\right) \prec \gamma_{2}\left(e_{1}\right) \vee \gamma_{2}\left(e_{2}\right): L$ as desired.

Case $\frac{\left(\lceil\Gamma\rceil \vdash e_{i}:{\underset{\varepsilon Q}{ }}_{A}\right)_{i}}{\Gamma \vdash\left\{e_{i}\right\}_{i}:\left\{{\underset{\varepsilon Q}{ }\}}_{A}^{A}\right\}}$. Applying lemma 13 we have by our IH that $\gamma_{1}\left(e_{i}\right) \equiv \gamma_{2}\left(e_{i}\right): \underset{\mathrm{EQ}}{A}$ and thus for some $v_{i, j}$ we have $\gamma_{j}\left(e_{i}\right) \longmapsto{ }^{*} v_{i, j}$ with $v_{i, 1} \equiv v_{i, 2}: \underset{\varepsilon}{ }$. . Then by calculation we have $\gamma_{j}\left(\left\{e_{i}\right\}_{i}\right) \longmapsto{ }^{*}\left\{\nu_{i, j}\right\}_{i}$ and applying LR SET (using partial reflexivity for its second premise) we have $\left\{v_{i, 1}\right\}_{j} \prec\left\{v_{i, 2}\right\}_{i}:\left\{\mathcal{A R Q}^{\mathcal{R}}\right\}$ and by closure under stepping we are done.


$$
\text { for }\left(x \in \gamma_{1}(e)\right) \gamma_{1}(f) \prec \text { for }\left(x \in \gamma_{2}(e)\right) \gamma_{2}(f): L
$$

By our IH we have $\gamma_{1}(e) \prec \gamma_{2}(e):\left\{\right.$ A $\left._{\varepsilon Q}\right\}$. Applying the definition of this, let $v_{i, j} \in \operatorname{Ok}_{\mathrm{V}}(\underset{\varepsilon}{ }(\underset{Q}{A})$ and $a_{i, j}$ be defined by:

$$
\left\{v_{i, j}\right\}_{j}=\text { value }\left(\gamma_{i}(e)\right) \quad a_{i, j}=f\left\{\gamma_{i}, x \mapsto v_{i, j}\right\}
$$

From the logical relation we have $(\forall \mathfrak{j}, \exists \mathrm{k}) \nu_{1, \mathrm{j}} \prec \nu_{2, \mathrm{k}}$ : $\underset{\in \mathrm{E}}{ }$. From this, our IH for f , and the definition of the logical relation for contexts we have $(\forall i, \exists j) a_{1, i} \prec a_{2, j}: A$. Observe that:

$$
\text { for } \begin{aligned}
\left(x \in \gamma_{i}(e)\right) \gamma_{i}(f) & \longmapsto *^{*} \text { for }\left(x \in\left\{v_{i, j}\right\}_{j}\right) \gamma_{i}(f) \\
& \longmapsto{ }^{*} \perp_{L} \vee_{L} f\left\{\gamma_{i}, x \mapsto v_{i, 1}\right\} \vee_{L} f\left\{\gamma_{i}, x \mapsto v_{i, 2}\right\} \vee_{L} \ldots \\
& =\perp_{L} \vee_{L} a_{i, 1} \vee_{L} a_{i, 2} \vee_{L} \ldots
\end{aligned}
$$

Then by closure under stepping we wish to show:

$$
\left(\perp_{\mathrm{L}} \vee_{\mathrm{L}} a_{1,1} \vee_{\mathrm{L}} a_{1,2} \vee_{\mathrm{L}} \ldots\right) \prec\left(\perp_{\mathrm{L}} \vee_{\mathrm{L}} a_{2,1} \vee_{\mathrm{L}} a_{2,2} \vee_{\mathrm{L}} \ldots\right): L
$$

Applying bottom is bottom and join is join, this follows from $(\forall i, \exists j) a_{1, i} \prec a_{2, j}: A$, because then transitively

$$
a_{1, i} \prec a_{2, i} \prec\left(\perp_{L} \vee_{L} a_{2,1} \vee_{L} a_{2,2} \vee_{L} \ldots\right): L
$$

Showing that the latter is an upper bound of each $a_{1, i}$ and since $\left(\perp_{L} V_{L} a_{1,1} V_{L}\right.$ $a_{1,2} \vee_{L} \ldots$ ) is least among such upper bounds, we have what we desire.

Case $\frac{\left(\lceil\Gamma\rceil \vdash e_{i}: \underset{\varepsilon Q}{A}\right)_{i}}{\Gamma \vdash e_{1}=e_{2}: \text { bool }}$. By lemma 13 for some $v_{i, j}$ we have $\gamma_{j}\left(e_{i}\right) \longmapsto^{*} v_{i, j}$ such that $\nu_{i, 1} \equiv \nu_{i, 2}:$ A. It suffices to show that $\left(v_{1,1}=v_{2,1}\right) \equiv\left(v_{1,2}=v_{2,2}\right)$ : bool. Since pretty plainly true $\prec$ true : bool and false $\prec$ false : bool (by the same reasoning as in the case for $\left\{e_{i}\right\}_{i}$ ), and since $v_{1, j}=v_{2, \mathrm{j}}$ steps to either true or false depending on
 By lemma 10 we know that this decidable ordering test coincides with equivalence in the logical relation, thus this holds because $v_{i, 1} \equiv v_{i, 2}$ : A. .

Case $\frac{\lceil\Gamma\rceil \vdash e:\{1\}}{\Gamma \vdash e m p t y ? e: 1+1}$. Applying our IH and lemma 13 we have $\gamma_{1}(e) \equiv \gamma_{2}(e):\{1\}$. This means we have some $\vec{v}, \vec{u}$ such that $\gamma_{1}(e) \longmapsto \longmapsto^{*}\{\vec{v}\}$ and $\gamma_{2}(e) \longmapsto^{*}\{\vec{v}\}$ and $\{\vec{v}\} \equiv\{\vec{u}\}:\{1\}$. This implies that $\vec{v}$ is empty if and only if $\vec{u}$ is empty. In one case empty? $\gamma_{i}(e) \longmapsto{ }^{*} \mathrm{in}_{1}()$; in the other, $\mathrm{in}_{2}()$. So by closure under stepping it suffices to show $\mathrm{in}_{\mathfrak{j}}() \in \mathrm{Ok}_{\mathrm{v}}(1+1)$ for $\mathfrak{j} \in\{1,2\}$; which it is by unrolling the definitions involved.

Case $\frac{\Gamma \vdash e: \square\left(A_{1}+A_{2}\right)}{\Gamma \vdash \text { split } e: \square A_{1}+\square A_{2}}$. By our IH and the LR for $\square\left(A_{1}+A_{2}\right)$ we have for some $\mathfrak{i} \in$ $\{1,2\}$ and $v_{1} \equiv v_{2}: A_{i}$ that $\gamma_{j}(e) \longmapsto{ }^{*}\left[\mathrm{in}_{i} v_{j}\right]$. Then $\gamma_{j}($ split $e) \longmapsto{ }^{*}$ split $\left[\mathrm{in}_{i} v_{j}\right] \longmapsto$ $\mathrm{in}_{\mathrm{i}}\left[v_{j}\right]$, and using the definition of the LR we have $\mathrm{in}_{\mathrm{i}}\left[\nu_{1}\right] \prec \mathrm{in}_{\mathrm{i}}\left[\nu_{2}\right]: \square A_{1}+\square A_{2}$ and by closure under stepping we are done.

## Chapter 3

## Seminaïve Evaluation

In chapter 2 we presented Datafun's syntax and semantics. These semantics are straightforward to implement directly; implementing them efficiently is more difficult. Datalog has decades of well-studied implementation and optimization techniques. To explore whether these techniques can be transferred to Datafun, in this chapter we'll examine just one classic Datalog optimization, seminaïve evaluation, which makes practical Datalog and Datafun's defining feature: iterative fixed points.

In $\S 3.1$ we'll see how the direct approach to finding fixed points wastes time by rediscovering previously-known facts at each iteration, and how seminaïve evaluation fixes this by computing the differences or changes between iterations. We therefore see seminaïve evaluation as a matter of incremental computation, that is, efficiently responding to change. To apply this insight, in $\S 3.2$ we adapt prior work on the incremental lambda calculus (Cai et al., 2014) to construct a category of incrementalizable monotone maps capable of interpreting Datafun's semantics. Using this construction as a guide, our central contribution (§3.4) is a pair of static Datafun-to-Datafun translations which enable the seminaïve fixed-point-finding strategy. Finally, we prove these transformations correct using a logical relation (§3.5).

### 3.1 Seminaïve evaluation as incremental computation

Consider the following Datalog program:

```
path (X, Z)\leftarrow edge(X, Z).
path }(X,Z)\leftarrow\operatorname{edge}(X,Y), path(Y,Z)
```

Suppose edge denotes a linear graph, $\{(1,2),(2,3), \ldots,(n-1, n)\}$. Then path will denote reachability by a sequence of one or more edges, $\{(\mathfrak{i}, \mathfrak{j}) \mid 1 \leqslant \mathfrak{i}<\mathfrak{j} \leqslant n\}$, or the transitive closure of edge. How can we compute this? The simplest approach is to begin with nothing in the path relation and repeatedly apply its rules until nothing more is deducible. We can make this strategy explicit by time-indexing the path relation:

$$
\begin{aligned}
& \operatorname{path}_{i+1}(X, Z) \leftarrow \operatorname{edge}(X, Z) . \\
& \operatorname{path}_{i+1}(X, Z) \leftarrow \operatorname{edge}(X, Y), \operatorname{path}_{i}(Y, Z) .
\end{aligned}
$$

When we index a relation in this way, the indexed relation at time $i$ will contain exactly those facts deducible by applying the original rules at most $i$ times. For instance, since the rules for path append edges one at a time, we can show by induction that path ${ }_{i}$ contains exactly the
nonempty paths of $i$ or fewer edges. By omission path ${ }_{0}=\emptyset$ : there are no empty nonempty paths. Inductively assuming for some $\mathfrak{i} \geqslant 0$ that path ${ }_{i}$ contains the nonempty paths of at most $i$ edges, note that "a nonempty path of length at most $i+1$ " is the same as "an edge, optionally followed by a nonempty path of length at most $i$ ". The singleton edges are included by the first clause, and edges followed by paths by the second clause (applying our inductive hypothesis); so path ${ }_{i+1}$ contains exactly the paths of length at most $i+1$ as desired.

The first clause path ${ }_{i+1}(X, Z) \leftarrow$ edge $(X, Z)$ ensures path ${ }_{i+1}$ includes all 1-edge paths; path $_{i+1}(X, Z) \leftarrow$ edge $(X, Y)$, path ${ }_{i}(Y, Z)$ includes all paths formed by prepending an edge to a path from path ${ }_{i}$, i.e. (by our inductive hypothesis) paths of between 2 and $i+1$ edges. So path ${ }_{i+1}$ contains exactly the paths of length 1 to $i+1$, as desired.

Unfortunately, this strategy re-deduces each previously known fact on every subsequent iteration. For example, suppose path $h_{\mathfrak{i}}(\mathfrak{j}, k)$ holds. Then at step $\mathfrak{i}+1$ the second rule deduces $\operatorname{path}_{i+1}(\mathfrak{j}-1, k)$ from edge $(j-1, j) \wedge$ path $_{i}(j, k)$. But since path ${ }_{i+1}(j, k)$ holds (a path of length at most $i$ is also a path of length at most $i+1$ ), we perform the same deduction at time $\mathfrak{i}+2$, and again at $\mathfrak{i}+3, \mathfrak{i}+4$, etc.

For our linear graph, it's easy to calculate how much work these re-deductions waste. The longest path in a linear graph of $n$ nodes has $n-1$ edges, so we take $n$ steps to discover a fixed point path $h_{n-1}=$ path $_{n}$. Since step $i$ involves $\mid$ path ${ }_{i} \mid=\sum_{j=1}^{i} n-j \in \Theta(i \cdot n)$ deductions, we make $\Theta\left(n^{3}\right)$ deductions in total. There being only $\Theta\left(n^{2}\right)$ paths in the final result, this is terribly wasteful; hence we term this naïve evaluation.

Now let's move from Datalog to Datafun. ${ }^{1}$ The transitive closure of edge is the least fixed point of the monotone function step defined by:

$$
\text { step } s=\text { edge } \cup\{(x, z) \mid(x, y) \in \text { edge, }(y, z) \in s\}
$$

The naïve way to compute this is to simply iterate step, computing path ${ }_{i}=$ step $^{i} \emptyset$ inductively by letting:

$$
\text { path }_{0}=\emptyset \quad \text { path }_{i+1}=\text { step path }_{i}
$$

But as before, path ${ }_{i} \subseteq$ step path $_{i}$; each iteration re-computes the paths found by its predecessor. We'd rather not compute the entire set, step path ${ }_{i}$, but instead find a smaller subset of new paths, let's call them dpath ${ }_{i}$, such that path ${ }_{i} \cup$ dpath $_{i}=$ step path $_{i}$. The smallest such set is of course step path ${ }_{i} \backslash$ path $_{i}$, but we won't need this most-precise difference to prove our strategy correct, and the freedom to approximate can be useful for avoiding unnecessary work. Our iteration strategy then becomes:

$$
\begin{array}{rlrl}
\text { path }_{0} & =\emptyset & \text { path }_{i+1} & =\text { path }_{i} \cup \text { dpath }_{i} \\
\text { dpath }_{0}=? & \text { dpath }_{i+1}=?
\end{array}
$$

The base case is easily solved by letting dpath $_{0}=\operatorname{step} \emptyset$. This wastes no work since there are no previously-known paths to be rediscovered. In the inductive case, we need to compute dpath $_{i+1}$ from path $_{i}$ and dpath ${ }_{i}$. Let's imagine we have a function that does this, called step':

$$
\begin{aligned}
\text { path }_{0} & =\emptyset & \text { path }_{i+1} & =\text { path }_{i} \cup \text { dpath }_{\mathfrak{i}} \\
\text { dpath }_{0} & =\text { step } \emptyset & \text { dpath }_{i+1} & =\text { step' }^{\prime} \text { path }_{i} \text { dpath }_{i}
\end{aligned}
$$

[^8]What must step' satisfy to prove this iteration strategy correct? We wish to show inductively that step path ${ }_{i}=$ path $_{i+1}$ for all $i$. The base case is trivial. So assuming step path ${ }_{i}=$ path $_{i+1}$, let's look at what we wish to prove and simplify it:

$$
\begin{aligned}
{\text { step } \text { path }_{i+1}} & =\text { path }_{i+2} & \text { what we wish to show } \\
\text { step }\left(\text { path }_{i} \cup \text { dpath }_{i}\right) & =\text { path }_{i+1} \cup \text { dpath }_{i+1} & \text { apply definitions } \\
\text { step }\left(\text { path }_{\mathfrak{i}} \cup \text { dpath }_{i}\right) & =\text { step path }_{\mathfrak{i}} \cup \text { step' }^{\prime} \text { path }_{i} \text { dpath }_{i} & \text { apply IH and definitions }
\end{aligned}
$$

So it suffices for step' to have the following property:

$$
\text { step }(s \cup d s)=\text { step } s \cup \text { step }^{\prime} s d s
$$

Intuitively speaking, step' s ds captures how step's output changes in response to changing input: as $s$ grows to $s \cup$ ds, how does step $s$ grow to step ( $s \cup d s$ )? This makes sense: our iterations are the outputs of step applied to increasing inputs. To compute the changes between them, we want to know how step's output responds to growth in its input.

The next question is: can we find a step' with this property? We can: for instance, step' $s$ ds $=$ step $(s \cup d s) \backslash$ step $s$, or the even simpler step' $s$ ds $=$ step $(s \cup d s)$. While technically correct, these solutions are not efficient: if we plug them into our revised iteration strategy, we find ourselves repeatedly calling step on ever-growing inputs, returning us to naïve iteration. So let's examine the behavior of step $(s \cup d s)$ to see if we can find a better alternative:

```
    step \((s \cup d s)\)
\(=\) edge \(\cup\{(x, z) \mid(x, y) \in\) edge, \((y, z) \in s \cup d s\}\)
\(=\) edge \(\cup\{(x, z) \mid(x, y) \in\) edge, \((y, z) \in s\} \cup\{(x, z) \mid(x, y) \in\) edge, \((y, z) \in d s\}\)
\(=\) step \(s \cup\{(x, z) \mid(x, y) \in\) edge, \((y, z) \in d s\}\)
```

Thus, a satisfactory definition of step ${ }^{\prime}$ is:

$$
\text { step' }^{\prime} \mathrm{d} \text { ds }=\{(x, z) \mid(x, y) \in \text { edge, }(y, z) \in \mathrm{ds}\}
$$

Is this efficient? Plugging this into our iteration strategy:

$$
\begin{aligned}
\operatorname{path}_{0} & =\emptyset \\
\text { dpath }_{0} & =\text { step } \emptyset \\
& =\text { edge }
\end{aligned}
$$

$$
\operatorname{path}_{i+1}=\text { path }_{i} \cup \text { dpath }_{i}
$$

$$
\begin{aligned}
\text { dpath }_{i+1} & =\text { step }^{\prime} \text { path }_{i} \text { dpath }_{i} \\
& =\left\{(x, z) \mid(x, y) \in \text { edge, }(y, z) \in \text { dpath }_{i}\right\}
\end{aligned}
$$

Applying this to our original linear graph example, edge $=\{(0,1),(1,2), \ldots,(n-1, n)\}$, we find:

$$
\begin{aligned}
\operatorname{dpath}_{0} & =\{(\mathfrak{i}, \mathfrak{i}+1) \mid 0 \leqslant \mathfrak{i}<\mathfrak{i}+1 \leqslant \mathfrak{n}\} \\
\operatorname{dpath}_{1} & =\{(\mathfrak{i}, \mathfrak{i}+2) \mid 0 \leqslant \mathfrak{i}<\mathfrak{i}+2 \leqslant \mathfrak{n}\} \\
\operatorname{dpath}_{2} & =\{(\mathfrak{i}, \mathfrak{i}+3) \mid 0 \leqslant \mathfrak{i}<\mathfrak{i}+3 \leqslant \mathfrak{n}\} \\
& \vdots \\
\operatorname{dpath}_{\mathrm{k}} & =\{(\mathfrak{i}, \mathfrak{i}+\mathfrak{k}+1) \mid 0 \leqslant \mathfrak{i}<\mathfrak{i}+\mathfrak{k}+1 \leqslant \mathfrak{n}\}
\end{aligned}
$$

Thus dpath ${ }_{k}$ captures exactly the paths of length $k$, so each path is discovered exactly once: we have avoided redundant work by computing only the change between iterations of our step function.

The problem of seminaïve evaluation for Datafun reduces to automatically finding functions, like step', that efficiently compute the change in a function's output given a change to its input. This is a problem of incremental computation, and since Datafun is a functional language, we use an approach rooted in the incremental $\lambda$-calculus (Cai et al., 2014; Giarrusso, 2020; Giarrusso et al., 2019).

### 3.2 Change structures for Datafun

To solve the problem of computing how a function's output changes in response to its input, we must first make precise the notion of change for each type in our language. To do this, incremental $\lambda$-calculi associate every type $A$ with a change structure. In our case, noting that Datafun types denote posets, we define change structures as follows:

Definition 14. A change structure $A$ consists of a poset $V A$, a poset $\Delta A$, and a relation $R_{A} \subseteq$ $\Delta A \times V A \times V A$. For $d x: \Delta A$ and $x, y: V A$, we will write $(d x, x, y) \in R_{A}$ interchangeably as $\mathrm{d} x \triangleright x \hookrightarrow y: A$. This relation must satisfy three properties:

Functionality If $d x \triangleright x \hookrightarrow y: A$ and $d x \triangleright x \hookrightarrow z: A$ then $y=z$.
Soundness If $d x \triangleright x \hookrightarrow y: A$ then $x \leqslant A y$.
Zero changes If $x: V A$ there is some $d x: \Delta A$ such that $d x \triangleright x \hookrightarrow x: A$.
Some useful terminology and notation: We can think of elements $d x \in \Delta A$ as changes or "diffs" to values $x \in V A$. The relation $R_{A}$ tell us how changes affect values: we gloss $d x \triangleright x \hookrightarrow y: A$ as " $d x$ changes $x$ into $y$ ". We say that $d x$ is a valid change to $x$ if there is some $y$ such that $d x \triangleright x \hookrightarrow y: A$. When $d x \triangleright x \hookrightarrow x$ we call $d x$ a zero change to $x$; when we need to pick such a change we write $\mathcal{O}_{x}$. By the axiom of choice, the zero changes property is equivalent to the existence of such a 0 function.

Although we use multi-letter variable names prefixed with "d" for elements $d x, d y: \Delta A$ of delta posets, this is merely a naming convention; we could instead use single-letter variables like $p, q: \Delta A$ with the same meaning.

To motivate our three properties, it will help to consider an example of a change structure corresponding to an important Datafun type: finite sets $\left\{\begin{array}{l}\text { A }\end{array}\right\}$. Recall that our goal is to speed up fixed point computation. Since iterations toward a fixed point grow monotonically, in Datafun we only need increasing changes. Therefore, changes to sets are themselves sets, to be unioned in:

Functionality says that $d x \triangleright x \hookrightarrow y:\left\{\begin{array}{c}A \\ \text { eq }\end{array}\right\}$ must be a partial function from $(d x, x)$ to $y$. In this case, it's a total function: set union. Soundness requires that all changes are increasing, which is true since $x \subseteq x \cup d x$. Finally, zero changes holds since $x \cup \emptyset=x$; one can leave a set unchanged by adding nothing. ${ }^{2}$

[^9]We'll see more examples of change structures later, including ones where the validity relation is a partial rather than a total function, but first, let's revisit our transitive closure example from §3.1. Using change structures we can generalize the relation between step and step'. We call step' a derivative, because it tells us how step's output changes in respond to its input changing:

Definition 15. A derivative of a monotone map $f: A \rightarrow B$ between change structures $A, B$ is a monotone map $f^{\prime}: \square \mathrm{VA} \rightarrow \Delta \mathrm{A} \rightarrow \Delta \mathrm{B}$ satisfying the law (for all $x, y, d x$ ):

$$
d x \triangleright x \hookrightarrow y: A \Longrightarrow f^{\prime} x d x \triangleright f x \hookrightarrow f y: B
$$

We say $a$ derivative, not the derivative, because derivatives are not necessarily unique. This is because changes are not necessarily unique: for fixed $x, y$ there may be many $d x$ such that $\mathrm{d} x \triangleright x \hookrightarrow y$.

Applying this definition to our change structure for finite sets, we recover the relationship we needed between step and step' in $\S 3.1$ for seminaïve evaluation:

$$
\begin{aligned}
& d s \triangleright s \hookrightarrow t:\left\{\begin{array}{c}
A \\
\varepsilon Q
\end{array}\right\} \Longrightarrow \text { step's ds } \triangleright \text { step } s \hookrightarrow \text { step } t:\left\{\begin{array}{l}
\text { AR }
\end{array}\right\} \\
& \text { iff } \\
& \mathrm{s} \cup \mathrm{ds}=\mathrm{t} \Longrightarrow \text { step } \mathrm{s} \cup \text { step }^{\prime} \mathrm{s} \mathrm{~d} s=\text { step } \mathrm{t} \\
& \text { iff } \\
& \text { step }(s \cup d s)=\text { step } s \cup \text { step }^{\prime} s d s
\end{aligned}
$$

This generalization is useful because differentiable maps (that is, maps possessing a derivative in the above sense) compose; in fact, they form a category:

Definition 16. The category $\Delta$ Poset has as objects change structures $A, B$ and as morphisms differentiable monotone maps $\mathrm{f}: \mathrm{VA} \rightarrow \mathrm{VB}$, that is, maps having at least one derivative $\mathrm{f}^{\prime}: \square \mathrm{VA} \rightarrow \Delta \mathrm{A} \rightarrow \Delta \mathrm{B}$ (also monotone). Morphism composition and the identity morphism are both as in Poset.

Proof. Of course, for this to be a category we need to show that: (1) the identity map is differentiable; (2) the composition of two differentiable maps is differentiable. We also need associativity and identity of composition, but these follow from the same in Poset. The derivative for identity is trivial, while the derivative for composition follows the pattern of the chain rule from calculus:

$$
\text { id }^{\prime} x d x=d x \quad(g \circ f)^{\prime} x d x=g^{\prime}(f x)\left(f^{\prime} x d x\right)
$$

That id' is a derivative of id is straightforward, while for composition we need to pick maps $f^{\prime}, g^{\prime}$ which are derivatives of $f, g$ respectively; then, applying the definition of derivatives:

$$
\begin{aligned}
& d x \triangleright x \hookrightarrow y \\
\Longrightarrow & f^{\prime} x d x \triangleright f x \hookrightarrow f y \\
\Longrightarrow & g^{\prime}(f x)\left(f^{\prime} x d x\right) \triangleright g(f x) \hookrightarrow g(f y)
\end{aligned}
$$

[^10]We also need id' $x d x$ and $(g \circ f)^{\prime} x d x$ to be monotone in $d x$, which they are, for straightforward compositional reasons; in general, for the remainder of $\S 3.2$ and 3.3 , we omit showing that functions are monotone unless the argument is non-obvious.

In the next section we will sketch the most important structures in $\Delta$ Poset needed to support Datafun's semantics, providing a recipe for incrementalizing Datafun. Applied correctly, this will let us automatically find derivatives for functions used by fix expressions, allowing us to employ the seminaïve evaluation strategy for finding fixed points faster.

### 3.3 The structure of $\Delta$ Poset

We should note up front that ours is only one among many reasonable notions of change structure. For instance, Giarrusso (2020) defines both basic change structures (definition 12.1.1), consisting only of a delta-set and a relation, and the more elaborate change structures (definition 13.1.1) that have an update operator $\oplus: A \times \Delta A \rightarrow A$, a difference operator $\ominus: A \times A \rightarrow \Delta A$, and composition of changes $\odot: \Delta A \times \Delta A \rightarrow \Delta A$; while Alvarez-Picallo (2020) uses a definition based on monoid actions. We will compare these with our approach in more detail in $\S 5.2$, but the "big picture" difference is that we are pervasively concerned with monotone functions, increasing changes, and higher-order computation; most of our choices flow from one more more of these considerations.

Our eventual destination is a static transformation on Datafun source code which implements seminaïve evaluation (§3.4). This transformation, originally presented by Arntzenius and Krishnaswami (2020), predates the construction of $\Delta$ Poset presented here and is independent of it. The transformation itself is quite intricate; our aim in presenting $\Delta$ Poset is to break its core concepts down into small pieces, showing how this complexity arises and suggesting potential alternatives for future investigation. In service of this goal, we have chosen what seems the simplest definition of change structure that both supports the features of Datafun and provides useful intuition. Ultimately, however, we deploy a logical relations argument to prove the translation correct (§3.5). A reader who does not care for a categorical view and is prepared to jump "in the deep end," therefore, may skim this section or jump straight to the definition of the seminaïve transformation itself in §3.4.

Recall that the structures we needed to interpret Datafun into the category Poset in §2.3.2 and 2.3.3 were: products, sums, exponentials, a discreteness comonad to interpret $\square$, sets and semilattice objects, equality-test morphisms, and fixed points. In $\Delta$ Poset we will cover products, sums, exponentials, the discreteness comonad, and fixed points, as they are the most significant for understanding the broad structure of our approach; the other structures are straightforward (with the exception of collect(f), which we discuss as for in §3.4.6).

### 3.3.1 Products

Products and the terminal object in $\Delta$ Poset mirror those in Poset:

$$
\begin{array}{rlrl}
\mathrm{V} 1 & =1 & \mathrm{~V}(\mathrm{~A} \times \mathrm{B}) & =\mathrm{VA} \times \mathrm{VB} \\
\Delta 1 & =1 & \Delta(\mathrm{~A} \times \mathrm{B}) & =\Delta \mathrm{A} \times \Delta \mathrm{B} \\
& & \begin{array}{l}
\text { PRodUct change }
\end{array} \\
() \triangleright() \hookrightarrow(): 1 & \frac{\mathrm{da} \triangleright \mathrm{a} \hookrightarrow \mathrm{a}^{\prime}: A}{(\mathrm{da}, \mathrm{db}) \triangleright(\mathrm{a}, \mathrm{~b}) \hookrightarrow\left(\mathrm{a}^{\prime}, \mathrm{b}^{\prime}\right): \mathrm{A} \times \mathrm{B}}
\end{array}
$$

These satisfy functionality, soundness, and zero changes by invoking the corresponding properties at $A$ and $B$. For instance, picking zero changes $0_{a}, 0_{b}$ for $a, b$ respectively, Product change tells us ( $0_{a}, 0_{b}$ ) is a zero change to ( $a, b$ ).

Finally, the terminal map $\left\rangle\right.$, projection $\pi_{i}$, and the tupling $\langle f, g\rangle$ of $f: A \rightarrow B$ and $g: A \rightarrow C$ are all the same as in Poset (thus inheriting the necessary universal properties), with derivatives given by:

$$
\begin{aligned}
\rangle: A \rightarrow 1 & \left\rangle^{\prime} a d a\right. & =() \\
\pi_{i}: A_{1} \times A_{2} \rightarrow A_{i} & \pi_{i}^{\prime}\left(x_{1}, x_{2}\right)\left(d x_{1}, d x_{2}\right) & =d x_{i} \\
\langle f, g\rangle: A \rightarrow B \times C & \langle f, g\rangle^{\prime} a d a & =\left(f^{\prime} a d a, g^{\prime} a \operatorname{da}\right)
\end{aligned}
$$

The correctness of $\left\rangle^{\prime}\right.$ is trivial; correctness of $\pi_{i}^{\prime}$ follows by inversion of Product change; and $\langle f, g\rangle^{\prime}$ is correct by Product Change and correctness of $f^{\prime}, g^{\prime} .{ }^{3}$

### 3.3.2 Sums

Sums and the initial object also mirror those in Poset:

$$
\begin{array}{rc}
\mathrm{V} 0=0 & \mathrm{~V}(\mathrm{~A}+\mathrm{B})=\mathrm{VA}+\mathrm{VB} \\
\Delta 0=0 & \Delta(\mathrm{~A}+\mathrm{B})=\Delta \mathrm{A}+\Delta \mathrm{B} \\
& \text { sum CHANGE } \\
\mathrm{R}_{0}=\emptyset & \frac{\mathrm{dx} \triangleright \mathrm{x} \hookrightarrow \mathrm{y}: \mathrm{A}_{\mathrm{i}}}{\mathrm{in}_{\mathrm{i}} \mathrm{~d} x \triangleright \mathrm{in}_{i} \mathrm{x} \hookrightarrow \mathrm{in}_{i} y: A_{1}+\mathrm{A}_{2}}
\end{array}
$$

These satisfy functionality, soundness, and zero changes pretty straightforwardly using the corresponding properties at $A$ and $B$. For instance, $i n_{i} ~_{x}$ is a zero change to in $n_{i} \chi$.

The initial map [], injection $\mathrm{in}_{\mathrm{i}}$, and case-analysis [ $\mathrm{f}_{1}, \mathrm{f}_{2}$ ] (given $\mathrm{f}_{1}: A_{1} \rightarrow \mathrm{C}, \mathrm{f}_{2}: A_{2} \rightarrow$ C ) are the same as in Poset (inheriting its universal properties), with derivatives as follows:

$$
\begin{aligned}
& \text { []:0 } \quad[]^{\prime}=[] \quad \text { (the domain is empty) } \\
& \mathrm{in}_{\mathrm{i}}: \mathrm{A}_{\mathrm{i}} \rightarrow \mathrm{~A}_{1}+\mathrm{A}_{2} \\
& \mathrm{in}_{\mathrm{i}}^{\prime} x \mathrm{~d} x=\mathrm{in} \mathrm{n}_{\mathrm{i}} \mathrm{~d} x \\
& {[f, g]: A_{1}+A_{2} \rightarrow C} \\
& {\left[f_{1}, f_{2}\right]^{\prime}\left(\mathrm{in}_{i} x\right)\left(\mathrm{in}_{\mathrm{j}} \mathrm{dx}\right)= \begin{cases}\mathrm{f}_{\mathrm{i}}^{\prime} \times \mathrm{d} x & \text { if } \mathfrak{i}=\mathfrak{j} \\
\text { anything oftype } \Delta \mathrm{C} & \text { if } \mathfrak{i} \neq \mathfrak{j}\end{cases} }
\end{aligned}
$$

Correctness of []' is vacuous; correctness of in ${ }_{i}^{\prime}$ follows directly from sum change; but the definition of $\left[\mathrm{f}_{1}, \mathrm{f}_{2}\right]^{\prime}$ requires explanation. If we take the proposition that $\left[\mathrm{f}_{1}, \mathrm{f}_{2}\right]^{\prime}$ is a derivative of $\left[f_{1}, f_{2}\right]$ and apply the definition of $R_{A_{1}+A_{2}}$ (namely SUM CHANGE), we find that it simplifies to:

$$
d x \triangleright x \hookrightarrow y: A_{i} \Longrightarrow\left[f_{1}, f_{2}\right]^{\prime}\left(i n_{i} x\right)\left(i n_{i} d x\right) \triangleright f_{i} x \hookrightarrow f_{i} y: C
$$

This only constrains the behavior of $\left[f_{1}, f_{2}\right]^{\prime}\left(i n_{i} x\right)\left(i n_{j} d x\right)$ when $\mathfrak{i}=\mathfrak{j}$; and in this case, we have $f_{i}^{\prime} \times d x \triangleright f_{i} x \hookrightarrow f_{i} y: C$ as desired. Since the $i \neq j$ case is unconstrained, any

[^11]value of type $\Delta \mathrm{C}$ will suffice; all we need for differentiability is to show one exists, i.e. that $\Delta \mathrm{C}$ is inhabited. Fortunately, in this case we have an $x: V A_{i}$ and a differentiable function $f_{i}: A_{i} \rightarrow$. Applying zero changes at $A_{i}$ we can pick a zero-change $0_{x}$ (although it being a zero-change is unnecessary; all we need is an element of $\Delta A_{i}$ ) and take $f_{i}^{\prime} \times 0_{x}: \Delta C$. Or, we could use zero-changes at C instead and take $\mathrm{O}_{\left(\mathrm{f}_{i} x\right)}: \Delta \mathrm{C}$.

This $i \neq j$ case is related to the partiality of the validity relation: $\mathrm{in}_{1} \mathrm{dx}$ is never a valid change to $\mathrm{in}_{2} \chi$. This is hard to avoid given our definition of change structures: to differentiate $\mathrm{in}_{i}$ and $\left[\mathrm{f}_{1}, \mathrm{f}_{2}\right]$ we need $\Delta\left(A_{1}+A_{2}\right)$ to include both $\Delta A_{1}$ and $\Delta A_{2}$ somehow; and a change $\mathrm{d} x \in \Delta \mathrm{~A}_{1}$ has no natural meaning applied to a value $x \in V A_{2}$. Furthermore, the fact that the $\mathfrak{i} \neq \mathfrak{j}$ case is unconstrained - essentially "dead code" - means that if we had defined $\Delta$ Poset morphisms as maps equipped with a particular derivative (rather than merely differentiable) we would be unable to prove the uniqueness of $[f, g]^{\prime}$ required by the universal property for sums. ${ }^{4}$

This could, for instance, be addressed by changing the definition of $\Delta$ Poset to only require derivatives to be defined for valid changes. We don't do this because from a type-theoretic perspective, this requires a dependent or refinement type, while we want the types of our derivatives to be simple so our category corresponds closely with a static transformation on Datafun, a simply-typed language.

### 3.3.3 Exponentials

The values of the exponentials in $\Delta$ Poset capture differentiable, monotone maps:

$$
\begin{aligned}
V(A \Rightarrow B) & =\operatorname{differentiable~monotone~maps~} V A \rightarrow V B \text {, ordered pointwise } \\
& =(\Delta \operatorname{Poset}(A, B),\{(f, g):(\forall x) f x \leqslant g x\})
\end{aligned}
$$

We might expect changes $\Delta(A \Rightarrow B)$ to be given pointwise, as (not necessarily monotone) functions $V A \rightarrow \Delta \mathrm{~B}$ mapping each input to the change in the corresponding output:

$$
\Delta(A \Rightarrow B)=\square V A \Rightarrow \Delta B \quad \frac{(\forall x) \mathrm{df} x \triangleright \mathrm{f} x \hookrightarrow \mathrm{~g} x: \mathrm{B}}{\mathrm{df} \triangleright \mathrm{f} \hookrightarrow \mathrm{~g}: \mathrm{A} \Rightarrow \mathrm{~B}} \quad x \text { not an exponential }
$$

However, this choice makes it difficult to differentiate function application. The function application map eval : $(A \Rightarrow B) \times A \rightarrow B$ is, of course, given by eval $(f, x)=f x$. To differentiate this is to ask for some eval' $(f, x)(d f, d x)$ that captures how $f x$ changes as both $f$ and $x$ change simultaneously: supposing $d f \triangleright f \hookrightarrow g$ and $d x \triangleright x \hookrightarrow y$, how do we find a change $f x \hookrightarrow g y$ ?

Using a pointwise change $\mathrm{df}: V A \rightarrow \Delta B$, we can find $\mathrm{df} x \triangleright \mathrm{f} x \hookrightarrow \mathrm{~g} x$; and applying differentiability of $f$ we can find some $f^{\prime} x d x \triangleright f x \hookrightarrow f y$. The former handles a change to the function, the latter a change to the argument. These form two sides of a "square of changes":

[^12]

We need the diagonal of this square. One approach would be to use pointwise function changes but augment our definition of change structures to allow composing changes, and find the diagonal by composing sides. Unfortunately, this is more difficult than it appears: eval' is applied to $f, x, d f, d x$, but to compute $d f y$ or $g^{\prime} x d x$ we need either $y$ or $g$. This seems to require equipping change structures with an operator $\oplus_{\mathrm{A}}: \square \mathrm{VA} \times \Delta \mathrm{A} \rightarrow \mathrm{VA}$ that extends the validity relation $R_{A}$ from a partial to a total function (since eval' is defined for all inputs, not merely valid ones); then we can recover $y=x \oplus d x$ or $g=f \oplus d f$. But $\oplus_{A \rightarrow B}$ is difficult to construct, because we must guarantee that $f \oplus d f$ is a monotone function, no matter the value of $\mathrm{df}: \square \mathrm{VA} \rightarrow \Delta \mathrm{B}$; it is easy to come up with a df such that $\lambda x$. $\mathrm{f} x \oplus \mathrm{df} x$ (the natural definition of $f \oplus d f$ ) is non-monotone. ${ }^{5}$

Perhaps there is some way through these difficulties; fortunately, there is a simple approach that side-steps them entirely: following the original incremental $\lambda$-calculus (Cai et al., 2014) we require function changes to produce the diagonal directly. Since this diagonal depends on the change $d x$ to the argument, function changes $d f$ become two-argument functions:

$$
\Delta(A \Rightarrow B)=\square V A \Rightarrow(\Delta A \Rightarrow \Delta B) \quad \frac{(\forall \mathrm{dx} \triangleright x \hookrightarrow \mathrm{y}: \mathrm{A}) \mathrm{df} x \mathrm{dx} \triangleright \mathrm{f} x \hookrightarrow \mathrm{~g} y: \mathrm{B}}{\mathrm{df} \triangleright \mathrm{f} \hookrightarrow \mathrm{~g}: \mathrm{A} \rightarrow \mathrm{~B}}
$$

With this definition, function changes are exactly what is needed to incrementalize function application $f x$. A change to a function $d f \triangleright f \hookrightarrow g$ accepts a change in its argument $d x \triangleright x \hookrightarrow y$ and produces the the change in its output, $d f x d x \triangleright f x \hookrightarrow g y$. If we return to our square of changes, we find it now has a zig-zag shape, with the diagonal filled in but missing the vertical sides:


To recover the missing sides, we can apply df to zero-changes $0_{x}$ and $0_{y}$ instead of $d x$ :


[^13]Zero changes thus let us recover a pointwise change $\lambda x$. df $x \mathcal{O}_{\chi}$ from any $\mathrm{df} \triangleright \mathrm{f} \hookrightarrow \mathrm{g}$ : $A \Rightarrow B$.

Note also the mixture of monotonicity and non-monotonicity in $\square V A \Rightarrow \Delta A \Rightarrow \Delta B$. Since our functions are monotone (increasing inputs yield increasing outputs), we expect function changes to be monotone with respect to input changes $\Delta \mathrm{A}$ : a larger increase in the input yields a larger increase in the output. However, there's no reason to expect the change in the output to grow as the base point increases - hence the first argument is discrete, $\square \mathrm{V} A$.

Although this nicely solves the problem of differentiating eval, it is not immediately obvious that $R_{A \Rightarrow B}$ is functional, sound, and possesses zero-changes. ${ }^{6}$ The first two are quite similar, so we'll tackle them together: Suppose $d f \triangleright f \hookrightarrow g: A \Rightarrow B$ and likewise df $\triangleright \mathrm{f} \hookrightarrow \mathrm{h}$ and fix some $x \in V A$. For functionality, we wish to show $g x=h x$; for soundness, we wish to show $\mathrm{f} x \leqslant \mathrm{~g} x$. By zero changes at $A$ we can pick some $0_{x} \triangleright x \hookrightarrow x$. Inverting fn change we have df $x 0_{x} \triangleright f x \hookrightarrow g x: B$ and likewise $d f x 0_{x} \triangleright f x \hookrightarrow h x$. Then by functionality at $B$ we have $g x=h x$; and by soundness at $B$ we have $f x \leqslant g x$.

Showing zero changes is simple but illuminating. By definition, every $f: V(A \Rightarrow B)$ is differentiable, and a derivative $f^{\prime}$ of $f$ is exactly a zero change $f^{\prime} \triangleright f \hookrightarrow f: A \Rightarrow B$ :

$$
\begin{array}{rlrl} 
& \mathrm{df} \triangleright \mathrm{f} \hookrightarrow \mathrm{f}: \mathrm{A} \rightarrow \mathrm{~B} & \\
\Longleftrightarrow & (\forall \mathrm{~d} x \triangleright \mathrm{x} \hookrightarrow \mathrm{y}: \mathrm{A}) \mathrm{df} x \mathrm{~d} x \triangleright \mathrm{f} x \hookrightarrow \mathrm{f} y: \mathrm{B} & \text { FN CHANGE } \\
\Longleftrightarrow & \mathrm{df} \text { is a derivative of } \mathrm{f} & & \text { definition } 15
\end{array}
$$

This happens because we've defined function changes $d f \triangleright f \hookrightarrow g: A \Rightarrow B$ to tell us how function application responds to changes in both the function and its argument. If the function doesn't change, this reduces to how the function's output changes as its argument changes: exactly what a derivative does.

Finally, to make $A \Rightarrow B$ an exponential we need function application eval ${ }_{A, B}:(A \Rightarrow$ B) $\times A \rightarrow B$ (which we have already discussed), and for any $f: C \times A \rightarrow B$, its currying $\lambda f: C \rightarrow A \Rightarrow B$. These are defined as in Poset, which ensures their universal property holds; but since $V(A \Rightarrow B)$ contains only differentiable maps, besides eval and $\lambda f$ we also require ( $\lambda \mathrm{f} \mathrm{c}$ ) to be differentiable:
${ }^{6}$ We also promised in a footnote on page 45 to show that completeness was problematic at function types. In other words, supposing $f \leqslant g: A \Rightarrow B$, why can't we find some df $\triangleright f \hookrightarrow g$ ? Well, we would need df $x \mathrm{~d} x \triangleright \mathrm{f} x \hookrightarrow \mathrm{~g} y$ : B whenever $\mathrm{d} x \triangleright x \hookrightarrow y$. If we inductively suppose completeness at $B$, we could pick such a change, since by monotonicity we can show $f x \leqslant g y$. Of course, we need $d f$ to be defined over all $x, \mathrm{~d} x$, not merely valid ones, but this is nothing the axiom of choice can't handle. More problematic is that we need $d f x d x$ to be monotone with respect to $d x$. So merely picking changes is not enough; we have to pick them in a way that preserves monotonicity.

We conjecture this can be done by strengthening change structures to include (1) monotone completeness and (2) monotone change composition. Monotone completeness strengthens completeness by requiring an operator $y \ominus_{A} x$ defined for $y \geqslant x$ : VA such that $y \ominus_{A} x \triangleright x \hookrightarrow y: A$ and which is monotone in $y$ and anti-monotone in $x$, that is, $x^{\prime} \leqslant x \wedge y \leqslant y^{\prime} \Longrightarrow y \ominus x \leqslant y^{\prime} \ominus x^{\prime}$. Monotone change composition requires a monotone operator $\odot_{A}: \Delta A \times \Delta A \rightarrow \Delta A$ such that if $d x \triangleright x \hookrightarrow y$ and $d y \triangleright y \hookrightarrow z$ then $d y \odot d x \triangleright x \hookrightarrow z$. Then we can define $g \ominus_{A \Rightarrow B} f=\lambda x . \lambda d x . g^{\prime} x d x \odot_{B}\left(g x \ominus_{B} f x\right)$ and $d g \odot_{A \Rightarrow B} d f=\lambda x . \lambda d x . d g x d x \circ d f x 0_{x}$.

We have not done this because it considerably complicates the definition of change structures and does not help explain any features of the translation given in §3.4, but it might be an interesting direction for future work.

$$
\begin{array}{rlrl}
\text { eval }(f, x) & =f x & \lambda f c x & =f(c, x) \\
\text { eval }^{\prime}(f, a)(d f, d a) & =d f a d a & (\lambda f)^{\prime} c d c a d a & =f^{\prime}(c, a)(d c, d a) \\
(\lambda f c)^{\prime} a d a & =f^{\prime}(c, a)\left(O_{c}, d a\right)
\end{array}
$$

We've already seen how eval's correctness follows from Fn Change. Applying $R_{A \Rightarrow B}$ and $R_{C \times A}$, we find that $(\lambda f)^{\prime}$ is a derivative for $\lambda f$ when $f^{\prime}$ is a derivative for $f$ :
$(\lambda f)^{\prime}$ is a derivative of $\lambda f$
$\Longleftrightarrow\left(\forall \mathrm{dc} \triangleright \mathrm{c} \hookrightarrow \mathrm{c}^{\prime}: \mathrm{C}\right)(\lambda \mathrm{f})^{\prime} \mathrm{c} d \mathrm{c} \triangleright \lambda \mathrm{fc} \hookrightarrow \lambda \mathrm{f} \mathrm{c}^{\prime}: A \Rightarrow B$
$\Longleftrightarrow\left(\forall d c \triangleright c \hookrightarrow c^{\prime}: C\right)\left(\forall d a \triangleright a \hookrightarrow a^{\prime}: A\right)(\lambda f)^{\prime} c$ dc $a d a \triangleright \lambda f c a \hookrightarrow \lambda f c^{\prime} a^{\prime}: B$
$\Longleftrightarrow\left(\forall(d c, d a) \triangleright(c, a) \hookrightarrow\left(c^{\prime}, a^{\prime}\right): C \times A\right) f^{\prime}(c, a)(d c, d a) \triangleright f(c, a) \hookrightarrow f\left(c^{\prime}, a^{\prime}\right): B$
$\Longleftrightarrow f^{\prime}$ is a derivative of $f$
Finally, the correctness of $(\lambda f c)^{\prime}$ follows from that of $f^{\prime}$ by applying $\mathcal{O}_{c} \triangleright c \hookrightarrow c: C$ :
$f^{\prime}$ is a derivative of $f$ and $O_{c}$ is a zero change to $c$
$\Longrightarrow\left(\forall d a \triangleright a \hookrightarrow a^{\prime}: A\right) f^{\prime}(c, a)\left(O_{c}, d a\right) \triangleright f(c, a) \hookrightarrow f\left(c, a^{\prime}\right): B$
$\Longleftrightarrow\left(\forall d a \triangleright a \hookrightarrow a^{\prime}: A\right)(\lambda f c)^{\prime} a d a \triangleright(\lambda f c) a \hookrightarrow(\lambda f c) a^{\prime}: B$
$\Longleftrightarrow(\lambda f c)^{\prime}$ is a derivative of $(\lambda f c)$

### 3.3.4 Semilattice change structures and seminaïve fixed points

We've already been introduced to the finite powerset change structure, as our introductory example in §3.2. But to define it properly as a functor P : $\Delta$ Poset $\rightarrow \Delta$ Poset, inheriting from the corresponding P on Poset:

$$
V P A=P V A \quad \Delta P A=P V A \quad d x \triangleright x \hookrightarrow y: P A \Longleftrightarrow x \cup d x=y
$$

This finite powerset change structure forms the prototype for our change structures for semilattices in general, which we need to support various language features, most importantly fixed points. We saw in §3.1-3.2 that given a function $f: P A \rightarrow P A$ and a derivative for it $f^{\prime}: \square P A \rightarrow P A \rightarrow P A$ we can compute its fixed point seminaïvely as follows:

$$
\begin{aligned}
x_{0} & =\emptyset \\
d x_{0} & =\mathrm{f} \emptyset
\end{aligned}
$$

$$
\begin{aligned}
x_{i+1} & =x_{i} \cup d x_{i} \\
d x_{i+1} & =f^{\prime} x_{i} d x_{i}
\end{aligned}
$$

This takes advantage of the fact that the change to a set is another set, and we apply a change using set union/semilattice join. Following this pattern, we can endow any semilattice L : Poset with a similar change structure:

$$
\mathrm{VL}=\mathrm{L} \quad \Delta \mathrm{~L}=\mathrm{L} \quad \mathrm{~d} x \triangleright x \hookrightarrow \mathrm{y}: \mathrm{L} \Longleftrightarrow x \vee \mathrm{~d} x=y
$$

This satisfies functionality ( $V$ is a function), soundness ( $x \leqslant x \vee y$ ), and zero-changes ( $x \vee \perp=x$ ). Let's call this the semilattice change structure on $L$. By construction, the finite powerset change structure $P A$ is the semilattice change structure on $P V A$; and our seminaïve fixed point strategy generalizes to any semilattice change structure:

Definition 17 (semifix). Given a semilattice $L$ with no infinite ascending chains and monotone maps $f: L \rightarrow L$ and $f^{\prime}: \square L \rightarrow L \rightarrow L$, let semifix $\left(f, f^{\prime}\right)=V_{i} x_{i}$ be the limit of the ascending chain defined by:

$$
\begin{aligned}
x_{0} & =\perp & x_{i+1} & =x_{i} \vee d x_{i} \\
d x_{0} & =\mathrm{f} \perp & \mathrm{~d} x_{i+1} & =\mathrm{f}^{\prime} x_{i} d x_{i}
\end{aligned}
$$

Theorem 18. semifix $x_{L}\left(f, f^{\prime}\right)$ is the least fixed point of $f$ if $f^{\prime}$ is a derivative of $f$.
Proof. It suffices to show inductively that $x_{i+1}=f x_{i}$; from this it follows that $x_{i}=f^{i} \perp$, as in the naïve approach to computing a fixed point. We prove this with essentially the same argument used in §3.1 (page 44). The base case is $x_{1}=x_{0} \vee \mathrm{~d} x_{0}=\perp \vee \mathrm{f} \perp=\mathrm{f} \perp=\mathrm{f} \mathrm{x}_{0}$, and the inductive case is:

$$
\begin{aligned}
x_{\mathfrak{i}+2} & =x_{i+1} \vee d x_{i+1} & & \text { definition of } x_{i+2} \\
& =f x_{\mathfrak{i}} \vee f^{\prime} x_{\mathfrak{i}} d x_{\mathfrak{i}} & & \text { inductive hypothesis, definition of } d x_{i+1} \\
& =f\left(x_{i} \vee d x_{i}\right) & & f^{\prime} \text { is a derivative of } f \\
& =f x_{i+1} & & \text { definition of } x_{i+1}
\end{aligned}
$$

### 3.3.5 Fixed points and discreteness comonads

Theorem 18 shows we can speed up fixed points by exploiting the power of derivatives. It may seem as though this justifies a morphism fix : $(\mathrm{L} \Rightarrow \mathrm{L}) \rightarrow \mathrm{L}: \Delta$ Poset for any semilattice change structure L satisfying ACC. However, morphisms in $\Delta$ Poset must be differentiable: does fix have a derivative? Prior work (Alvarez-Picallo et al., 2019; Arntzenius, 2017) has answered this affirmatively. One solution is to find the fixed point of the function change:

$$
f x^{\prime} f d f=f i x(d f(f i x f))
$$

How and why this works is non-obvious; we refer the reader to Arntzenius (2017) for a full explanation. So there is indeed a morphism fix : $(\mathrm{L} \Rightarrow \mathrm{L}) \rightarrow \mathrm{L}: \Delta$ Poset. However, from the perspective of our original goal of speeding up fixed point computations, the derivative of this morphism presents two issues. First, it isn't actually incremental: computing fix' $f$ df using this derivative requires re-computing fix $f$ as an argument to $d f!^{7}$ Second, we would naturally like to compute fix ( $\mathrm{df}(\mathrm{fixf})$ ) seminaïvely, but we have no guarantee that ( $\mathrm{df}(\mathrm{fixf})$ ) is differentiable! This requires a higher-order derivative; a coherent theory of higher-order derivatives and higher-order change structures would be enormously interesting, but we leave it to future work.

Instead, we deliberately limit the scope of our approach to avoid the need to incrementally maintain fixed points. As we mentioned in §2.3.1, in Datafun fix is not treated as a monotone operator; correspondingly the morphisms we require to interpret it are not fix : (L $\Rightarrow \mathrm{L}) \rightarrow \mathrm{L}$

[^14]but rather fix : $\square(\mathrm{L} \Rightarrow \mathrm{L}) \rightarrow \mathrm{L}$. The idea here is that, just as $\square$ in Poset captures nonmonotonicity in an otherwise monotone world, in $\Delta$ Poset we can use it to capture nondifferentiability or non-incrementalizability in an otherwise differentiable world.

More concretely, since we only consider increasing changes and $\square A$ is ordered discretely, $x \leqslant y: \square A \Longleftrightarrow x=y$, the only possible "change" is to stay the same. We can thus extend the discreteness comonad $\square$ on Poset to a comonad $\square$ on $\Delta$ Poset by letting the space of changes be trivial:

$$
\mathrm{V} \square \mathrm{~A}=\square \mathrm{VA} \quad \Delta \square \mathrm{~A}=1 \quad() \triangleright \mathrm{a} \hookrightarrow \mathrm{a}: \square \mathrm{A}
$$

This straightforwardly satisfies functionality, soundness, and zero changes. Moreover, it inherits the monoidal comonad structure of $\square$ from Poset. Fixing some map $f: A \rightarrow B$, the derivatives of functorial action, extraction, duplication, and distribution are mostly trivial:

$$
\begin{aligned}
\square(\mathrm{f}) & : \square \mathrm{A} \rightarrow \square \mathrm{~B} \\
\varepsilon_{A} & : \square \mathrm{A} \rightarrow \mathrm{~A} \\
\delta_{A} & : \square \mathrm{A} \rightarrow \square \square \mathrm{~A} \\
\operatorname{dist}_{\square}^{\times} & : \prod_{i} \square A_{\mathrm{i}} \rightarrow \square \prod_{i} A_{i} \\
\operatorname{dist}_{+}^{\square} & : \square \sum_{i} A_{i} \rightarrow \sum_{i} \square A_{i}
\end{aligned}
$$

$$
\begin{aligned}
\square(\mathrm{f})^{\prime} \times() & =() \\
\varepsilon_{\mathcal{A}}^{\prime} \times() & \left.=0_{x} \quad \text { (see footnote }{ }^{8}\right) \\
\delta_{A}^{\prime} \times() & =() \\
\operatorname{dist}_{\square}^{\times \prime} \times \mathrm{d} x & =() \\
\operatorname{dist}_{+}^{\square^{\prime}}\left(\mathrm{in}_{\mathrm{i}} x\right)() & =\mathrm{in}_{\mathrm{i}}()
\end{aligned}
$$

Finally, observe that any monotone map $\mathrm{f}: \mathrm{V} \square \mathrm{A} \rightarrow \mathrm{VB}$ is trivially differentiable by letting $\mathrm{f}^{\prime} \mathrm{x}()=\mathcal{O}_{(\mathrm{f} x)}$, confirming our intuition that differentiable maps $\square A \rightarrow B$ should coincide with not-necessarily-differentiable maps $A \rightarrow B$.

### 3.4 The $\phi$ and $\delta$ transforms

Our goal in this chapter is to automatically speed up computation of fixed points in Datafun using seminaïve evaluation, replacing fix $f$ by semifix ( $f, f^{\prime}$ ) where $f^{\prime}$ is a derivative for $f$. The previous section suggests this is possible: $\Delta$ Poset appears capable of interpreting Datafun's semantics; its exponential objects consist of differentiable monotone maps; and its construction provides a recipe for calculating concrete derivatives witnessing this differentiability. In this section, we carry out this recipe (with one significant change), producing two static transformations, $\phi$ and $\delta$, defined in figure 3.2.

The "speed-up" transform $\phi$ replaces fix $f$ by semifix ( $f, f^{\prime}$ ) and decorates other expressions with the information we need to compute $f^{\prime}$. In particular, to find $f^{\prime}$ we need the "incrementalization" transform, $\delta$, which propagates changes through a program. For instance, the derivative of ( $\lambda$ X. e) depends on how e changes in response to changes in $X$. In general, $\delta e$ computes the change in $\phi e$ given changes to its free variables. The rules defining $\delta$ closely resemble both the incremental $\boldsymbol{\lambda}$-calculus' Derive operator (Cai et al., 2014) and the derivatives given in the previous section for morphisms in $\Delta$ Poset.

Unfortunately, when we consider terms with free variables, there is a gap between derivatives and changes: we cannot simply let $f^{\prime}=\delta f$, because we want $f^{\prime}$ to be the derivative of the

8 It is again important here that in $\Delta$ Poset our morphisms are merely differentiable, not equipped with derivatives, that is, that we do not distinguish morphisms by their derivative. Otherwise naturality of $\varepsilon$ would require that $(\varepsilon \circ \square f)^{\prime}=(f \circ \varepsilon)^{\prime}$, which requires $\varepsilon^{\prime}(f x)()=f^{\prime} x\left(\varepsilon^{\prime} x()\right)$ and thus $0_{(f x)}=f^{\prime} \times 0_{x}$, which is difficult to guarantee without unique zero-changes.

$$
\begin{aligned}
\Phi 1 & =1 \\
\Phi\{\mathcal{E Q}\} & =\left\{\Phi \text { A }_{\mathrm{EQ}}\right\} \quad \text { (see lemma 19) } \\
\Phi(\square \mathrm{A}) & =\square(\Phi A \times \Delta \Phi A) \\
\Phi(\mathrm{A} \times \mathrm{B}) & =\Phi A \times \Phi B \\
\Phi(\mathrm{~A}+\mathrm{B}) & =\Phi A+\Phi B \\
\Phi(\mathrm{~A} \rightarrow \mathrm{~B}) & =\Phi A \rightarrow \Phi B
\end{aligned}
$$

$$
\begin{aligned}
\Delta 1 & =1 \\
\Delta\left\{\mathrm{AQ}_{\mathrm{EQ}}\right\} & =\{\mathrm{A}\} \\
\Delta(\square \mathrm{A}) & =1 \\
\Delta(\mathrm{~A} \times \mathrm{B}) & =\Delta \mathrm{A} \times \Delta \mathrm{B} \\
\Delta(\mathrm{~A}+\mathrm{B}) & =\Delta \mathrm{A}+\Delta \mathrm{B} \\
\Delta(\mathrm{~A} \rightarrow \mathrm{~B}) & =\square \mathrm{A} \rightarrow \Delta \mathrm{~A} \rightarrow \Delta \mathrm{~B}
\end{aligned}
$$

## FIGURE 3.I $\Delta$ and $\Phi$ type transformations

function $f$, not the change to it. One way to solve this problem, suggested by the exponential in $\Delta$ Poset, would be to have $\phi$ decorate every function with its derivative. However, this is overkill: we only need derivatives for functions used in fixed point computations.

This leads to our significant change. Recall from §3.3.3 that a zero change to a function is a derivative for it: the problematic gap between derivatives and changes disappears if the function does not change. We ensure this by giving the fix the argument type $\square(\mathrm{L} \mid \mathrm{Lx} \rightarrow \underset{\text { fix }}{\mathrm{L}})$; as we saw in §3.3.5, the type $\square A$ represents values which do not change. Thus, rather than decorate every function with its derivative, the key strategy of the $\phi$ transformation is to decorate expressions of type $\square \boldsymbol{A}$ with their zero changes. In this way, we hijack the $\square$ comonad to track functions that are used inside fixed points and make their derivatives available where we need them: at fix expressions.

### 3.4.1 Typing $\phi$ and $\mathcal{\delta}$

In order to decorate expressions with extra information, $\phi$ also needs to decorate their types. In figure 3.1 we give a type translation $\Phi A$ capturing this. In particular, if $e: \square A$ then $\phi e$ will have type $\Phi(\square A)=\square(\Phi A \times \Delta \Phi A)$. The idea is that evaluating $\phi e$ will produce a pair $[(x, d x)]$ where $x: \Phi A$ is the sped-up result and $d x: \Delta \Phi A$ is a zero change to $x$. For example, if $e: \square(A \rightarrow B)$, then $\phi e$ will compute $\left[\left(f, f^{\prime}\right)\right]$, where $f^{\prime}$ is the derivative of $f$.

On types other than $\square A$, there is no information we need to add, so $\Phi$ simply distributes. In particular, source programs and sped-up programs agree on the shape of first-order data:

Lemma 19. $\Phi \underset{\varepsilon}{ } A_{\varepsilon}={\underset{\varepsilon}{e}}_{A}$ for all equality types $\underset{\varepsilon}{A}$.
Proof. Induct on $\underset{\varepsilon}{A}$ applying the equations in figure 3.1, recalling from figure 2.1 that the

It will also be important that, as in our semilattice change structures in $\Delta$ Poset, changes at a semilattice type L are drawn from the very same type:

Lemma 20. At each semilattice type L , we have $\Delta \mathrm{L}=\mathrm{L}$.
Proof. Induct on L applying the equations in figure 3.1, recalling from figure 2.1 that the grammar of semilattice types is $L::=1\left|L_{1} \times L_{2}\right|\left\{\begin{array}{c}\text { AR }\end{array}\right\}$.

As we'll see in $\S 3.4 .3$ and 3.4.4, $\phi$ and $\delta$ are mutually recursive. To make this work, $\delta e$ must find the change to $\phi e$ rather than $e$. So if $e: A$ then $\phi e: \Phi A$ and $\delta e: \Delta \Phi A$. However, so far we have neglected to say what $\phi$ and $\delta$ do to typing contexts. To understand this, it's helpful to look at what $\Phi$ and $\Delta \Phi$ do to functions and to $\square$. This is because expressions

$$
\begin{aligned}
\phi X & =X \\
\phi(\lambda X . e) & =\lambda X . \phi e \\
\phi\left(e_{i}\right)_{i} & =\left(\phi e_{i}\right)_{i} \\
\phi\left(\mathrm{in}_{i} e\right) & =\mathrm{in}_{i} \phi e \\
\phi \perp & =\perp \\
\phi\left(\left\{e_{i}\right\}_{i}\right) & =\left\{\phi e_{i}\right\}_{i} \\
\phi[e] & =[(\phi e, \delta e)] \\
\phi(e=f) & =(\phi e=\phi f) \\
\phi(f i x e) & =\text { semifix } \phi e
\end{aligned}
$$

$$
\begin{aligned}
& \phi x= x \\
& \phi(e f)= \phi e \phi f \\
& \phi\left(\pi_{i} e\right)= \pi_{i} \phi e \\
& \phi\left(\text { case } e \text { of }\left(i n_{i} X \rightarrow f_{i}\right)_{i}\right)= \text { case } \phi e \text { of }\left(i n_{i} X \rightarrow \phi f_{i}\right)_{i} \\
& \phi(e \vee f)= \phi e \vee \phi f \\
& \phi(\text { for }(x \in e) f)= \text { for }(x \in \phi e) \text { let }[d x]=[0 x] \text { in } \phi f \\
& \phi(\text { let }[x]=e \text { in } f)= \text { let }[(x, d x)]=\phi e \text { in } \phi f \\
& \phi(\text { empty? } e)= \text { empty? } \phi e \\
& \phi(\text { split } e) \stackrel{\star}{=} \text { case } \phi e \text { of } \\
&\left(\left[\left(i i_{i} x, i i_{i} d x\right)\right] \rightarrow i i_{i}[(x, d x)]\right)_{i} \\
&\left(\left[\left(i n_{i} x, i n_{j}\right)\right] \rightarrow i n_{i}[(x, \text { dummy } x)]\right)_{i \neq j}
\end{aligned}
$$

## DERIVATIVE TRANSLATION $\delta$

$$
\delta \perp=\delta\left\{e_{i}\right\}_{i}=\delta(e=f)=\delta(\text { fix } e)=\perp
$$

$$
\delta\left(\text { case } e \text { of }\left(\mathrm{in}_{i} X \rightarrow \mathrm{f}_{i}\right)_{i}\right) \stackrel{\star}{=} \text { case split }[\phi e], \delta e \text { of }
$$

$$
\begin{aligned}
& \left(\operatorname{in}_{i}[x], \text { in }_{i} D X \rightarrow \delta f_{i_{i}}\right)_{i} \\
& \left(\mathrm{in}_{i}[x], \text { in }{ }_{j} \rightarrow \text { let } D X=\text { dummy } x \text { in } \delta f_{i}\right)_{i \neq j} \\
\delta(\text { for }(x \in e) \text { f })= & (\text { for }(x \in \delta e) \text { let }[d x]=[0 x] \text { in } \phi f) \\
\vee & (\text { for }(x \in \phi e \vee \delta e) \text { let }[d x]=[0 x] \text { in } \delta f)
\end{aligned}
$$

Equations marked with a red star, $\stackrel{\star}{=}$, use pattern-matching syntax sugar we have not previously defined; see figure 3.3 for expansions.

FIGURE $3.2 \phi$ and $\delta$ term translations

$$
\begin{aligned}
& \delta X=D X \\
& \delta(\lambda X . e)=\lambda[x] . \lambda D X . \delta e \\
& \delta\left(e_{i}\right)_{i}=\left(\delta e_{i}\right)_{i} \\
& \delta\left(\mathrm{in}_{i} e\right)=\mathrm{in}_{\mathrm{i}} \delta e \\
& \delta[e]=() \\
& \delta(e m p t y ? e)=e m p t y ? \phi e \\
& \delta x=d x \\
& \delta(e f)=\delta e[\phi f] \delta f \\
& \delta\left(\pi_{i} e\right)=\pi_{i} \delta e \\
& \delta(e \vee f)=\delta e \vee \delta f \\
& \delta(\text { let }[x]=e \text { in } f)=\operatorname{let}[(x, d x)]=\phi e \text { in } \delta f \\
& \delta(\text { split } e) \stackrel{\star}{=} \text { case } \phi e \text { of }\left(\left[\left(i n_{i},{ }^{\prime}\right)\right] \rightarrow i n_{i}()\right)_{i}
\end{aligned}
$$

ADDITIONAL DESUGARINGS
$\phi($ split $e) \stackrel{\star}{=}$ case $\phi e$ of
$\left(\left[\left(\mathrm{in}_{i} x, \mathrm{in}_{i} \mathrm{~d} x\right)\right] \rightarrow \mathrm{in}_{\mathrm{i}}[(\mathrm{x}, \mathrm{d} x)]\right)_{i}$
$\left(\left[\left(\mathrm{in}_{\mathrm{i}} x, \mathrm{in} \mathrm{n}_{\mathrm{j}}\right)\right] \rightarrow \mathrm{in}_{\mathrm{i}}[(\mathrm{x} \text {, dummy } \mathrm{x})]\right)_{\mathrm{i} \neq \mathrm{j}}$
$=$ let $[z]=\phi e$ in
case split $\left[\pi_{1} z\right]$ of

$$
\left(\mathrm{in}_{\mathrm{i}} Y \rightarrow \text { let }[\mathrm{x}]=\mathrm{Y}\right. \text { in }
$$

case split $\left[\pi_{2} z\right]$ of
$\operatorname{in}_{i}$ DY $\rightarrow$ let $[d x]=D Y$ in in $_{i}[(x, d x)]$
$\mathrm{in}_{\mathrm{i}+1 \bmod 2}{ }_{-} \rightarrow \mathrm{in}_{i}[(x$, dummy $\left.x)]\right)_{i}$
$\delta($ split $e) \stackrel{\star}{=}$ case $\phi e$ of $\left.\left(\left[\left(\mathrm{in}_{i_{-}},\right)_{-}\right)\right] \rightarrow \mathrm{in}_{\mathrm{i}}()\right)_{\mathrm{i}}$

$$
=\text { let }[y]=\phi e \text { in case } \pi_{1} y \text { of }\left(\mathrm{in}_{i_{-}} \rightarrow \mathrm{in}_{i}()\right)_{i \in\{1,2\}}
$$

$\delta\left(\right.$ case $e$ of $\left.\left(i n_{i} X \rightarrow f_{i}\right)_{i}\right) \stackrel{\star}{=}$ case split [ $\left.\phi e\right], \delta e$ of

$$
\left(\mathrm{in}_{\mathfrak{i}}[x], \mathrm{in}_{\mathrm{i}} \mathrm{DX} \rightarrow \delta \mathrm{f}_{\mathrm{i}}\right)_{\mathrm{i}}
$$

$$
\left(\mathrm{in}_{\mathfrak{i}}[x], \mathrm{in}_{\mathrm{j}-} \rightarrow \text { let } \mathrm{DX}=\text { dummy } x \text { in } \delta \mathrm{f}_{\mathfrak{i}}\right)_{i \neq j}
$$

$=$ case split [ $\phi e]$ of
$\left(\mathrm{in}_{\mathrm{i}} \mathrm{Y} \rightarrow\right.$ let $[\mathrm{x}]=\mathrm{Y}$ in
$\left(\lambda D X . \delta f_{i}\right)\left(\right.$ case $\delta e$ of $\mathrm{in}_{\mathrm{i}} \mathrm{DX} \rightarrow \mathrm{DX}$
$\mathrm{in}_{\mathrm{i}+1 \bmod 2} \rightarrow$ dummy x$\left.)\right)_{\mathrm{i}}$
Fresh variables introduced by desugaring are colored pink.
FIGURE 3.3 Additional syntax sugar for $\phi$ and $\delta$ transformations
denote functions of their free variables. Moreover, in Datafun free variables come in two flavors, monotone and discrete, and discrete variables are semantically $\square$-ed.

Viewed as functions of their free variables, $\delta e$ denotes the derivative of $\phi e$. And just as the derivative of a unary function $f x$ has two arguments, $d f x d x$, the derivative of an expression $e$ with $n$ variables $x_{1}, \ldots, x_{n}$ will have $2 n$ variables: the original $x_{1}, \ldots, x_{n}$ and their changes $d x_{1}, \ldots, d x_{n} .{ }^{9}$ However, this says nothing yet about monotonicity or discreteness. To make this precise, we'll use three context transformations, named according to the analogous type operators $\square$, $\Phi$, and $\Delta$ :

$$
\begin{array}{ll}
\square(X: A)=x:: A & \square(x:: A)=x:: A \\
\Phi(X: A)=X: \Phi A & \Phi(x:: A)=x: \Phi A, d x: \Delta \Phi A \\
\Delta(X: A)=D X: \Delta A & \Delta(x: A)=\varepsilon
\end{array}
$$

Otherwise all three operators distribute; e.g. $\square \varepsilon=\varepsilon$ and $\square\left(\Gamma_{1}, \Gamma_{2}\right)=\square \Gamma_{1}, \square \Gamma_{2}$. Intuitively, $\square \Gamma, \Phi \Gamma$, and $\Delta \Gamma$ mirror the effect of $\square$, $\Phi$, and $\Delta$ on the semantics of $\Gamma$ :

$$
\begin{array}{lll}
\llbracket \square \Gamma \rrbracket \cong \square \llbracket \Gamma \rrbracket & \llbracket \Phi(X: A) \rrbracket \cong \llbracket \Phi A \rrbracket & \llbracket \Delta(X: A) \rrbracket \cong \llbracket \Delta A \rrbracket \\
& \llbracket \Phi(x:: A) \rrbracket \cong \llbracket \Phi \square A \rrbracket & \llbracket \Delta(x:: A) \rrbracket \cong \llbracket \Delta \square A \rrbracket
\end{array}
$$

These defined, we can state the types of $\phi e$ and $\delta e$ :
Theorem 21 (Well-typedness of $\phi, \delta$ ). If $\Gamma \vdash e: A$, then $\phi e$ and $\delta e$ have the following types:

$$
\begin{gathered}
Ф Г \vdash \phi e: Ф А \\
\square Ф Г, \Delta \Phi Г \vdash \delta e: \Delta \Phi А
\end{gathered}
$$

Proof. By induction on the derivation of $\Gamma \vdash e: A$, although as we'll see shortly we will need weakening (theorem 22) in some places.

As expected, if we view expressions as functions of their free variables, and pretend $\Gamma$ is a type, these correspond to $\Phi(\Gamma \rightarrow \mathcal{A})$ and $\Delta \Phi(\Gamma \rightarrow \mathcal{A})$ respectively:

$$
\Phi(\Gamma \rightarrow \mathcal{A})=\Phi Г \rightarrow \Phi А \quad \Delta \Phi(\Gamma \rightarrow А)=\square Ф \Gamma \rightarrow \Delta \Phi \Gamma \rightarrow \Delta \Phi А
$$

To get the hang of these context and type transformations, suppose $x:: A, Y: B \vdash e: C$. Then theorem 21 tells us:

$$
\begin{gathered}
x:: \Phi A, \mathrm{dx}:: \Delta \Phi A, \mathrm{Y}: \Phi \mathrm{B} \vdash \phi e: \Phi \mathrm{C} \\
\mathrm{x}:: \Phi \overline{\mathrm{c}} \mathrm{dx}:: \Delta \Phi A, \mathrm{y}:: \Phi \mathrm{B}, \mathrm{DY}: \Delta \Phi \mathrm{B} \vdash \delta e: \Delta \Phi \mathrm{C}
\end{gathered}
$$

Along with the original program's variables, $\phi e$ requires zero change variables $d x$ for every discrete source variable $x$. Meanwhile, $\delta e$ requires changes for every source program variable (for discrete variables these will be zero changes), and moreover is discrete with respect to the source program variables (the "base points").

We now have enough information to tackle the definitions of $\phi$ and $\delta$ given in figure 3.2. In the remainder of this section, we'll examine the most interesting and important parts of these definitions in detail.

[^15]
### 3.4.2 Fixed points

The whole purpose of $\phi$ and $\delta$ is to speed up fixed points, so let's start there. In a fixed point expression fix $e$, we know $e: \square\left(L_{\text {fix }} \rightarrow \underset{\text { fix }}{L}\right)$. Consequently the type of $\phi e$ is

$$
\begin{aligned}
& \Phi(\square(\underset{\mathrm{FX}}{\mathrm{~L}} \rightarrow \underset{\mathrm{FX}}{\mathrm{~L}}))=\square(\Phi(\underset{\mathrm{fX}}{\mathrm{~L}} \rightarrow \underset{\mathrm{FX}}{\mathrm{~L}}) \times \Delta \Phi(\underset{\mathrm{FX}}{\mathrm{~L}} \rightarrow \underset{\mathrm{FXX}}{\mathrm{~L}})) \\
& =\square\left(\left(\Phi \Phi_{\mathrm{FX}}^{\mathrm{L}} \rightarrow \Phi \Phi_{\mathrm{FX}}^{\mathrm{L}}\right) \times\left(\square \Phi \Phi_{\mathrm{FX}}^{\mathrm{L}} \rightarrow \Delta \Phi \Phi_{\mathrm{FX}}^{\mathrm{L}} \rightarrow \Delta \Phi \Phi_{\mathrm{FX}}^{\mathrm{L}}\right)\right)
\end{aligned}
$$

$$
\begin{aligned}
& =\square\left((\mathrm{LEX} \rightarrow \underset{\mathrm{FX}}{\mathrm{~L}}) \times\left(\square \square_{\mathrm{FX}}^{\mathrm{L}} \rightarrow \underset{\mathrm{FX}}{\mathrm{~L}} \rightarrow \underset{\mathrm{FIX}}{\mathrm{~L}}\right) \quad \text { by lemma 20, } \Delta \mathrm{fix}_{\mathrm{L}}^{\mathrm{L}}=\underset{\mathrm{fix}}{\mathrm{~L}}\right.
\end{aligned}
$$

The behavior of $\phi e$ is to compute a boxed pair $\left[\left(f, f^{\prime}\right)\right]$, where $f: \underset{\mathrm{Hx}}{\mathrm{L}} \rightarrow \underset{\mathrm{fx}}{\mathrm{L}}$ is a sped-up function and $f^{\prime}: \square L_{\mathrm{FIX}} \rightarrow \underset{\mathrm{flx}}{\mathrm{L}} \rightarrow \underset{\mathrm{F} \mid \mathrm{x}}{\mathrm{L}}$ is its derivative. This is exactly what we need in order to call semifix. Therefore $\phi($ fix e) $=$ semifix $\phi e$. However, if we're going to use semifix in the output of $\phi$, we ought to give it a typing rule and semantics:

$$
\begin{array}{cc}
\Gamma \vdash e: \square\left(\left(\left(_{\mathrm{fx}}^{\mathrm{L}} \rightarrow \underset{\mathrm{fix}}{\mathrm{~L}}\right) \times(\square \mathrm{f} \underset{\mathrm{tix}}{\mathrm{~L}} \rightarrow \underset{\mathrm{fix}}{\mathrm{~L}} \rightarrow \underset{\mathrm{fix}}{\mathrm{~L}})\right.\right. & \text { semifix e』 } \gamma=\text { semifix }\left(\mathrm{f}, \mathrm{f}^{\prime}\right) \\
\Gamma \vdash \text { semifix } e: \underset{\mathrm{fx}}{\mathrm{~L}} & \text { where }\left(\mathrm{f}, \mathrm{f}^{\prime}\right)=\llbracket e \rrbracket \gamma
\end{array}
$$

As for $\delta$ (fix e), since e can't change (having $\square$ type), neither can fix e (or semifix $\phi e$ ). All we need is a zero change at type $\underset{f \times x}{ }$; by lemma $20, \perp$ suffices.

### 3.4.3 Variables, $\lambda$-abstraction, and application

At the core of a functional language are variables, $\lambda$-abstraction, and application. The $\phi$ translation leaves these alone, simply distributing over subexpressions. On variables, $\delta$ yields the corresponding change variables. On functions and application, $\delta$ is more interesting:

$$
\begin{aligned}
\Delta \Phi(A \rightarrow B) & =\square \Phi A \rightarrow \Delta \Phi A \rightarrow \Delta \Phi B \\
\delta(\lambda X . e) & =\lambda[x] . \lambda D X . \delta e \\
\delta(e f) & =\delta e[\phi f] \delta f
\end{aligned}
$$

The intuition behind $\delta(\lambda X . e)=\lambda[x] . \lambda D X . \delta e$ is that a function change takes two arguments, a base point $x$ and a change $D X$, and yields the change in the result of the function, $\delta e$. However, we are given an argument of type $\square \Phi A$, but consulting theorem 21 for the type of $\delta e$, we need a discrete variable $x:=\Phi A$, so we use pattern-matching to unbox our argument. The intuition behind $\delta(e f)=\delta e[\phi f] \delta f$ is much the same: $\delta e$ needs two arguments, the original input $\phi \mathrm{f}$ and its change $\delta \mathrm{f}$, to return the change in the function's output. Moreover, it's discrete in its first argument, so we need to box it, [ $\phi f]$.

One might ask why this type-checks, since $\phi e$ and $\delta e$ don't use the same typing context. We're even boxing $\phi \mathrm{f}$, hiding all monotone variables; consequently, it gets the context $\lceil\square \Phi \Gamma, \Delta \Phi\lceil \rceil$. However, $\square$ makes every variable discrete, and $\lceil-\rceil$ leaves discrete variables alone, so this provides at least $\square Ф \Gamma$, while the context $\phi \mathrm{f}$ needs is $\Phi \Gamma$. Thus really this is a question about the interaction of weakening and discreteness: can a discrete variable always substitute for a monotone one?

Indeed it may: making a variable discrete only increases the number of places it can be used, because while some typing rules discard monotone variables, they never discard discrete ones. We formalize this using a weakening relation $\Gamma \sqsubseteq \Delta$ (figure 3.4; note that H

| EMPTY | CONS | DROP <br> $\bar{\varepsilon} \sqsubseteq \varepsilon$ | $\frac{\Gamma \sqsubseteq \Delta}{\Gamma, \mathrm{H} \sqsubseteq \Delta, \mathrm{H}}$ |
| :--- | :--- | :--- | :--- |$\quad \frac{\text { DISC }}{\Gamma \sqsubseteq \Delta, \mathrm{H}} \quad$| $\Gamma, \mathrm{X}: \mathrm{A} \sqsubseteq \Delta, x: \mathrm{A}$ |
| :---: |

FIGURE 3.4 Weakening relation
for "hypothesis" ranges over all variable typings, monotone or discrete), which is standard except for the rule DISC, which says that a discrete hypothesis is weaker than a monotone one. We can then show that typing respects weakening:

Theorem 22 (Weakening). If $\Delta \sqsupseteq \Gamma$ and $\Gamma \vdash e: A$ then $\Delta \vdash e: A$.
Proof. By induction on the derivation of $\Gamma \vdash e: A$; see appendix A.2.
We use this without further note throughout the $\phi$ and $\delta$ transformations.

### 3.4.4 The discreteness comonad, $\square$

Our strategy hinges on decorating expressions of type $\square A$ with their zero changes, so the translations of $[e]$ and (let $[x]=e$ in $f$ ) are of particular interest. The most trivial of these is $\delta[e]=()$; this follows from $\Delta \Phi \square A=1$, since boxed values cannot change.

Next, consider $\phi[e]=[(\phi e, \delta e)]$. The intuition here is straightforward: $\phi$ needs to decorate $e$ with its zero change; since $e$ is discrete and cannot change, we use $\delta e$. However! In general, one cannot use $\delta$ inside the $\phi$ translation and expect the result to be well-typed; $\phi$ and $\delta$ require different typing contexts. To see this, let's apply theorem 21 to singleton contexts:

| $\Gamma$ (context of e) | $Ф \Gamma$ (context of фe) | $\square Ф Г, \triangle Ф Г$ (context of $\delta$ e) |
| :---: | :---: | :---: |
| X:A | X: ФА | $x:=Ф А, \mathrm{DX}: \triangle \Phi \mathcal{}$ |
|  | $x:=\Phi А, \mathrm{dx}$ :: $\Delta \Phi \mathcal{A}$ | $\mathrm{x}:=\Phi \mathcal{A}, \mathrm{d} x:=\Delta \Phi \mathcal{}$ |

Luckily, although $Ф\lceil$ and $\square Ф \Gamma, \Delta Ф \Gamma$ differ on monotone variables, they agree on discrete ones. And since $e$ is discrete, it has no free monotone variables, justifying the use of $\delta e$ in $\phi[e]=[(\phi e, \delta e)]$.

Next we come to (let $[x]=e$ in $f$ ), whose $\phi$ and $\delta$ translations are very similar:

$$
\left.\begin{array}{rl}
\phi(\text { let }[x] & =e \text { in } f)
\end{array}=\text { let }[(x, d x)]=\phi e \text { in } \phi f \text { let }[x]=e \text { in } f\right)=\text { let }[(x, d x)]=\phi e \text { in } \delta f \text { }
$$

Since $x$ is a discrete variable, both $\phi f$ and $\delta f$ need access to its zero change $d x$. Luckily, $\phi e: \square(\Phi A \times \Delta \Phi A)$ provides it, so we simply unpack it. We don't use $\delta e$ in $\delta$ f, but this is unsurprising when you consider that its type is $\Delta \Phi \square \mathcal{A}=1$.

$$
\begin{aligned}
\text { dummy }_{\{\mathcal{A A R}\}} & =\{ \} & \text { dummy }_{A \times B}(x, y) & =(\text { dummy } x \text {, dummy } y) \\
\text { dummy }_{1}() & =() & \text { dummy }_{A+B}\left(\text { in }_{i} x\right) & =\text { in }_{i}(\text { dummy } x) \\
\text { dummy }_{\square A}[x] & =[\text { dummy } x] & \text { dummy }_{A \rightarrow B} f & =\lambda x . \text { dummy }(f x)
\end{aligned}
$$

FIGURE 3.5 The function dummy $_{A}: A \rightarrow \Delta A$

### 3.4.5 Case analysis, split, and dummy

The derivative of case-analysis, $\delta$ (case $e$ of $\left.\left(\mathrm{in}_{i} X_{i} \rightarrow f_{i}\right)_{i}\right)$, is complex. Suppose $\phi e$ evaluates to $\mathrm{in}_{i} x$ and its change $\delta e$ evaluates to $\mathrm{in}_{\mathrm{j}} \mathrm{d} x$. Recall that sums are ordered disjointly (§2.3.2); the value $x$ can increase, but the tag $\mathrm{in}_{\mathrm{i}}$ must remain the same. Since $\delta e$ is a change to $\phi e$, the change structure on sums (§3.3.2) tells us that $i=j$ ! So the desired change $\delta$ (case e of ...) is given by $\delta f_{i}$ in a context supplying a discrete base point $x$ (the value $x$ ) and the change DX. To bind $x$ discretely, we need to use $[\phi e]: \square(\Phi A+\Phi B)$; to pattern-match on this, we need split to distribute the $\square$.

This handles the first two cases, $\left(i n_{i}[x], i n_{i} D X \rightarrow \delta f_{i}\right)_{i}$. Since we know the tags on $\phi e$ and $\delta e$ agree, these are the only possible cases. However, since the output of our translation is Datafun code, to appease the type-checker we must handle the impossible case that $\mathfrak{i} \neq \mathfrak{j}$. This case is dead code: it needs to typecheck, but is otherwise irrelevant. It suffices to generate a dummy change $\mathrm{dx}: \Delta \Phi A_{i}$ from our base point $x:=\Phi A_{i}$. We do this using a simple function dummy $_{A}: A \rightarrow \Delta A$ (figure 3.5).

We also need dummy in the definition of $\phi$ (split $e$ ). In effect split has type $\square(A+B) \rightarrow$ $\square A+\square B$. Observe that

$$
\begin{aligned}
& \Phi(\square(\mathrm{A}+\mathrm{B}))=\square((\Phi \mathrm{A}+\Phi \mathrm{B}) \times(\Delta \Phi \mathrm{A}+\Delta \Phi \mathrm{B})) \\
& \Phi(\square \mathrm{A}+\square \mathrm{B})=\square(\Phi \mathrm{A} \times \Delta \Phi \mathcal{A})+\square(\Phi \mathrm{B} \times \Delta \Phi \mathrm{B})
\end{aligned}
$$

So while $\phi e$ yields a boxed pair of tagged values, $\left[\left(\mathrm{in}_{\mathrm{i}} x, \mathrm{in}_{\mathrm{j}} \mathrm{d} x\right)\right]$, we need $\phi($ split $e)$ to yield a tagged boxed pair, $\mathrm{in}_{i}[(x, d x)]$. Again we use dummy to handle the impossible case $\mathfrak{i} \neq \mathrm{j}$.

Finally, observe that $\delta$ (split $e$ ) has type $\Delta \Phi(\square A+\square B)=\Delta \Phi \square A+\Delta \Phi \square B=1+1$. All it must do is return ( $\mathrm{in} \mathrm{n}_{\mathrm{i}}()$ ) with a tag that matches $\phi$ (split e) and $\phi e$; case-analysing фe suffices.

### 3.4.6 Semilattices and comprehensions

The translation $\phi(e \vee f)=\phi e \vee \phi f$ is straightforward, but $\delta(e \vee f)=\delta e \vee \delta f$ is not as simple as it seems. Restricting to sets, suppose that $d x$ changes $x$ into $x^{\prime}$ and $d y$ changes $y$ to $y^{\prime}$. In particular, suppose these changes are precise: that $d x=x^{\prime} \backslash x$ and $d y=y^{\prime} \backslash y$. Then the precise change from $x \cup y$ into $x^{\prime} \cup y^{\prime}$ is:

$$
\left(x^{\prime} \cup y^{\prime}\right) \backslash(x \cup y)=\left(x^{\prime} \backslash x \backslash y\right) \cup\left(y^{\prime} \backslash y \backslash x\right)=(d x \backslash y) \cup(d y \backslash x)
$$

This suggests letting $\delta(e \cup f)=(\delta e \backslash \phi f) \cup(\delta f \backslash \phi e)$. This is a valid derivative, but it involves recomputing $\phi e$ and $\phi f$, and our goal is to avoid recomputation. So instead, we overapproximate the derivative: $\delta e \cup \delta f$ might contain some unnecessary elements, but we expect it to be cheaper to include these than to recompute $\phi e$ and $\phi f$. This overapproximation
agrees with seminaïve evaluation in Datalog: Datalog implicitly unions the results of different rules for the same predicate (e.g. those for path in §3.1), and the seminaïve translations of these rules do not include negated premises to compute a more precise difference.

Now let's consider for $(x \in e)$ f. Its $\phi$-translation is straightforward, with one hitch: because $x:: \underset{\boxminus}{A}$ is a discrete variable, the inner loop $\phi \mathrm{f}$ needs access to its zero change $\mathrm{dx}:: \Delta{\underset{\varepsilon}{\varepsilon}}^{\mathrm{A}}$. Conveniently, at eqtypes (although not in general), the dummy function computes zero changes:

Lemma 23. If $x: \underset{⿷ \in}{A}$ then dummy $x \triangleright x \hookrightarrow x: \underset{E Q}{A}$.
Proof. By induction on $\underset{\varepsilon \in}{A}$, unfolding the definition of (dummy $\mathrm{d} x \triangleright x \hookrightarrow x: \underset{\varepsilon}{ }$ A) from $\S 3.2$ at each step. For example, when $\underset{\varepsilon}{A}=\left\{B_{\varepsilon}\right\}$, we need to show that $x \cup$ dummy $d x=x$, which is

For clarity, we write 0 rather than dummy when we use it to produce zero changes; we only call it dummy in dead code.

Finally, we come to $\delta($ for $(x \in e) f)$, the computational heart of the seminaïve transformation, as for is what enables embedding relational algebra (the right-hand-sides of Datalog clauses) into Datafun. Here there are two things to consider, corresponding to the two for-clauses generated by $\delta($ for $(x \in e)$ f). First, if the set $\phi e$ we're looping over gains new elements $x \in \delta e$, we need to compute $\phi f$ over these new elements. Second, if the inner loop $\phi \mathrm{f}$ changes, we need to add in its changes $\delta \mathrm{f}$ for every element, new or old, in the looped-over set, $\phi e \vee \delta e$. Just as in the $\phi$-translation, we use $0 /$ dummy to calculate zero changes to set elements.

### 3.4.7 Leftovers

The $\phi$ rules we haven't yet discussed simply distribute $\phi$ over subexpressions. The remaining $\delta$ rules mostly do the same, with a few exceptions. In the case of $\delta\left(\left\{e_{i}\right\}_{i}\right)=\delta(e=f)=\perp$, the sub-expressions are discrete and cannot change, so we produce a zero change $\perp$. This is also the case for $\delta$ (empty? e) = empty? $\phi e$, but as with $\delta$ (split $e$ ), the zero change here is at type $1+1$, so to get the tag right we must analyse $\phi e$.

### 3.5 Proving the seminaïve transformation correct

We have given two program transformations: $\phi e$, which optimizes $e$ by computing fixed points seminaïvely; and $\delta e$, which finds the change in $\phi e$ under a change in its free variables. To state the correctness of $\phi$ and $\delta$, we need to show that $\phi e$ preserves the meaning of $e$ and that $\delta e$ correctly updates $\phi e$ with respect to changes in its variable bindings. Since our transformations modify the types of higher-order expressions to include the extra information needed for seminaïve evaluation, we cannot directly prove that the semantics is preserved. Instead, we formalize the relationship between $e, \phi e$, and $\delta e$ using a logical relation, and use this relation to prove an adequacy theorem saying that the semantics is preserved for closed, first-order programs.

So, inductively on types $A$, letting $\mathrm{a}, \mathrm{b} \in \llbracket A \rrbracket, x, y \in \llbracket \Phi A \rrbracket$, and $d x \in \llbracket \Delta \Phi A \rrbracket$, we define a five place relation $d x \triangleright_{A} x \not z a \rightarrow y\{b$, meaning roughly " $x, y$ speed up $a, b$ respectively, and $d x$ changes $x$ into $y$ ". The full definition is in figure 3.6.

$$
\begin{aligned}
& \text { () } \triangleright_{1}() \nsucceq() \rightarrow() \nsucceq() \Longleftrightarrow \top \\
& \overrightarrow{\mathrm{dx}} \triangleright_{A_{1} \times \mathrm{A}_{2}} \vec{x} \nmid \overrightarrow{\mathrm{a}} \rightarrow \overrightarrow{\mathrm{y}} \not \overrightarrow{\mathrm{~b}} \Longleftrightarrow(\forall \mathrm{i}) \mathrm{d} x_{\mathrm{i}} \triangleright_{\mathrm{A}_{\mathrm{i}}} x_{\mathrm{i} \nless} \not \mathrm{a}_{\mathrm{i}} \rightarrow \mathrm{y}_{\mathrm{i}} \nless \mathrm{~b}_{\mathrm{i}}
\end{aligned}
$$

$$
\begin{aligned}
& \mathrm{df} \triangleright_{\mathrm{A} \rightarrow \mathrm{~B}} \mathrm{f} \not \mathrm{f}^{\prime} \rightarrow \mathrm{g} \not \mathrm{~g}^{\prime} \Longleftrightarrow\left(\forall \mathrm{d} x \triangleright_{\mathrm{A}} x \notin \mathrm{a} \rightarrow \mathrm{y} \nless \mathrm{~b}\right) \\
& \text { df } x d x \triangleright_{B} f x \notin f^{\prime} a \rightarrow g y \not g^{\prime} b \\
& \mathrm{dx} \triangleright_{\left\{\begin{array}{c}
\text { A }\}
\end{array}\right.} x \notin \mathrm{a} \rightarrow \mathrm{y} \dot{z} \mathrm{~b} \Longleftrightarrow(x, y, x \cup \mathrm{~d} x)=(\mathrm{a}, \mathrm{~b}, \mathrm{y}) \\
& \text { () } \triangleright_{\square A}(x, d x) \notin a \rightarrow(y, d y) \notin b \Longleftrightarrow(a, x, d x)=(b, y, d y) \wedge d x \triangleright_{A} x \notin a \rightarrow y \notin b
\end{aligned}
$$

## FIGURE 3.6 Definition of the logical relation

At product, sum, and function types this is essentially a more elaborate version of the change structures given in §3.2. At set types, changes are still a set of values added to the initial value, but we additionally insist that the "slow" $a, b$ and "speedy" $x, y$ are equal. This is because we have engineered the definitions of $\Phi$ and $\phi$ to preserve behavior on equality types. Finally, since $\square A$ represents values which cannot change, $d x$ is an uninformative empty tuple and the original and updated values are identical. However, the "speedy" values are now pairs of a value and its zero change. This ensures that at a boxed function type, we will always have a derivative (a zero change) available.

The logical relation is defined on simple values, and so before we can state the fundamental theorem, we have to extend it to contexts $\Gamma$ and substitutions, letting $\rho, \rho^{\prime} \in \llbracket \Gamma \rrbracket, \gamma, \gamma^{\prime} \in \llbracket \Phi \Gamma \rrbracket$, and $\mathrm{d} \gamma \in \llbracket \Delta Ф Г \rrbracket$ :

$$
\begin{aligned}
\mathrm{d} \gamma \triangleright_{\Gamma} \gamma \& \rho \rightarrow \gamma^{\prime} \& \rho^{\prime} & \Longleftrightarrow(\forall X: A \in \Gamma) d \gamma_{D x} \triangleright_{A} \gamma_{x} \& \rho_{X} \rightarrow \gamma_{X}^{\prime} \& \rho_{X}^{\prime} \\
& \wedge(\forall x:: A \in \Gamma)() \triangleright_{\square A}\left(\gamma_{x}, \gamma_{d x}\right) \& \rho_{x} \rightarrow\left(\gamma_{x}^{\prime}, \gamma_{d x}^{\prime}\right) \& \rho_{x}^{\prime}
\end{aligned}
$$

With that in place, we can state the fundamental theorem, showing that $\phi$ and $\delta$ generate expressions which satisfy this logical relation:

Theorem 24 (Fundamental property). If $\Gamma \vdash e: A$ and $d \gamma \triangleright_{\Gamma} \gamma z \rho \rightarrow \gamma^{\prime} \& \rho^{\prime}$ then

$$
\llbracket \delta e \rrbracket(\gamma, \mathrm{~d} \gamma) \triangleright_{\mathrm{A}} \llbracket \phi e \rrbracket \gamma z \llbracket e \rrbracket \rho \rightarrow \llbracket \phi e \rrbracket \gamma^{\prime} z \llbracket e \rrbracket \rho^{\prime}
$$

## Proof. See appendix A.2.

This theorem follows by a structural induction on typing derivations as usual, but requires a number of lemmas. By and large, these lemmas generalize or build on results stated earlier regarding the simpler change structures from §3.2. For example, we build on lemmas 19 and 23 to characterize the logical relation at equality types $\underset{\mathrm{EQ}}{A}$ and the behavior of dummy:

Lemma 25 (Equality changes). If $d x \triangleright_{\underset{E}{ }} x \notin a \rightarrow y \notin b$ then $x=a$ and $y=b$.

Lemma 26 (Dummy is zero at eqtypes). If $x \in \llbracket \underset{E Q}{A} \rrbracket$ then dummy $x \triangleright_{\underset{E}{A}}^{A} x \not x \rightarrow x \not x x$.
Proof. In each case, induct on $\underset{E Q}{A}$. See appendix A. 2 .

Lemma 25 tells us that at equality types, the sped-up version of a value is the value itself. This is used later to prove our adequacy theorem. Lemma 26 is an analogue of lemma 23, showing that dummy function computes zero changes at equality types. This is used in the proof of the fundamental theorem for for-loops, in whose $\phi$ and $\delta$ translations 0 is implemented by dummy.

Next, we generalize lemma 20 to characterize changes at semilattice type:
Lemma 27 (Semilattice changes). At any semilattice type L , we have $\Delta \mathrm{L}=\mathrm{L}$, and moreover $d x \triangleright_{\mathrm{L}} x \& a \rightarrow y$ b $b$ iff $x=a$ and $y=b=x \vee_{L} d x$

Proof. By induction on semilattice types L, applying lemma 25 (noting that every semilattice type is an equality type).

We require this lemma in the proofs of the fundamental theorem in all the cases involving semilattice types - namely $\perp$, $\vee$, for, and fix.

Since typing rules that involve discreteness (such as the $\square$ rules) manipulate the context, we need some lemmas regarding these manipulations. First, we show that all valid changes for a context with only discrete variables send substitutions to themselves, recalling that $\lceil\Gamma\rceil$ contains only the discrete variables from $\Gamma$.

Lemma 28 (Discrete contexts don’t change). If () $\triangleright_{\lceil\Gamma\rceil} \gamma \& \rho \rightarrow \gamma^{\prime}$ \& $\rho^{\prime}$ then $\gamma=\gamma^{\prime}$ and $\rho=\rho^{\prime}$.

Proof. All variables in the stripped contexts are discrete, and therefore the logical relation for discrete variables in contexts, which invokes the logical relation at $\square$ type, requires their corresponding components be equal.

We use this lemma in combination with the next, which says that any valid context change gives rise to a valid change on a stripped context:

Lemma 29 (Context stripping). If $\mathrm{d} \gamma \triangleright_{\Gamma} \gamma\left\{\rho \rightarrow \gamma^{\prime} \& \rho^{\prime}\right.$ then

$$
() \triangleright_{\Gamma \Gamma\rceil} \operatorname{strip}_{\Phi \Gamma}(\gamma) \& \operatorname{strip}_{\Gamma}(\rho) \rightarrow \operatorname{strip}_{\Phi \Gamma}\left(\gamma^{\prime}\right) \text { strip }\left(\rho^{\prime}\right)
$$

where strip ${ }_{\Gamma}=\left\langle\pi_{\chi}\right\rangle_{x:: А \in \Gamma}$ keeps only the discrete variables from a substitution.

Proof. Immediate from the definitions.
Jointly, these two lemmas ensure that a valid change to any context is an identity on the discrete part. We use these in all the cases of the fundamental theorem involving discrete expressions - equality $e_{1}=e_{2}$, set literals $\left\{e_{i}\right\}_{i}$, emptiness tests empty? $e$, and box introduction [e].

Combining all these lemmas to establish the fundamental theorem, adequacy follows immediately:

Proof. By theorem 24, noting the premise is trivial since the context is empty, we have $\llbracket \delta e \rrbracket \triangleright_{\substack{A \\ \varepsilon}} \llbracket \phi e \rrbracket z \llbracket e \rrbracket \rightarrow \llbracket \phi e \rrbracket$ 文 $\llbracket e \rrbracket$, which by lemma 25 implies $\llbracket \phi e \rrbracket=\llbracket e \rrbracket$.

## Chapter 4

## Implementation and Efficiency

The previous chapter was entirely theoretical, formalizing the intuition that seminaïve evaluation works by computing the changes between iterations toward a fixed point by, first, constructing a theory of changes for Datafun; and second, applying that theory to construct and prove correct a program transformation which implements this strategy. However, the purpose of seminaïve evaluation is not to push changes around, but to compute results faster. We have proven that our transformed program computes the same result, but not shown that it does so more efficiently. In this chapter we remedy this experimentally, observing that at least two further optimizations are necessary to achieve asymptotic performance improvements.

First, in $\S 4.1$ we apply the seminaïve program transformation by hand to our running example, transitive closure. In the process we uncover some obvious inefficiencies in the transformed code and demonstrate how to optimize them away. In $\S 4.2$ we discuss our implementation of a Datafun-to-Haskell compiler, which we use to demonstrate experimentally that seminaïve evaluation can produce asymptotic performance improvements when combined with these optimizations.

Second, in $\S 4.3$ we observe that even with these optimizations, there remain cases where we do asymptotically more work than necessary, not because of inefficiencies in the transformed program, but because of the imprecision of our derivatives. This results in overly large changes which accumulate across fixed point iterations. We implement a simple solution based on change minimization and test it experimentally.

### 4.1 Applying the seminaïve transformation to transitive closure

Let's try applying the seminaïve transform to a simple Datafun program: the transitive closure function trans from §2.2.1:
trans [edge] $=$ fix $R$ is edge $\cup($ edge $\bullet R)$
$S \bullet T=$ for $\left(\left(x, y_{1}\right) \in S\right)$ for $\left(\left(y_{2}, z\right) \in T\right)$ for $\left(y_{1}=y_{2}\right)\{(x, z)\}$
In the process we'll discover that besides $\phi$ itself we need a few simple optimisations to actually speed up our program: most importantly, we need to propagate $\perp$ expressions.

In our experience, performing $\phi$ and $\delta$ by hand is easiest done from the inside outwards. At the core of transitive closure is a relation composition, $(e \bullet p)$, and at the core of relation composition is a "one-sided conditional", for $\left(y_{1}=y_{2}\right)\{(x, z)\}$. Let's take a look at its $\phi$ and $\delta$ translations:

$$
\phi\left(\text { for }\left(y_{1}=y_{2}\right)\{(x, z)\}\right)
$$

$$
=\phi\left(\text { for }\left(() \in y_{1}=y_{2}\right)\{(x, z)\}\right) \quad \text { desugar }
$$

$$
=\text { for }\left(() \in y_{1}=y_{2}\right) \phi\{(x, z)\} \quad \text { apply } \phi \text { and omit unused let }
$$

$$
=\text { for }\left(y_{1}=y_{2}\right)\{(x, z)\} \quad \text { resugar }
$$

Frequently, as in this case, $\phi$ does nothing interesting. For brevity we'll skip such no-op translations. Now for the $\delta$ translation:

$$
\begin{array}{rll} 
& \delta\left(\text { for }\left(y_{1}=y_{2}\right)\{(x, z)\}\right) & \text { desugar } \\
= & \delta\left(\text { for }\left(() \in y_{1}=y_{2}\right)\{(x, z)\}\right) & \\
= & \text { for }\left(() \in \delta\left(y_{1}=y_{2}\right)\right) \phi\{(x, z)\} & \text { apply } \phi / \delta \text { to } y_{1}=y_{2} \text { and }\{(x, z)\} \\
& \cup \text { for }\left(() \in \phi\left(y_{1}=y_{2}\right) \cup \delta\left(y_{1}=y_{2}\right)\right) \delta\{(x, z)\} & \text { apply } \delta \text { and omit unused lets } \\
= & \text { for }(() \in \perp)\{(x, z)\} & \text { propagate } \perp
\end{array}
$$

Thus:

$$
\begin{equation*}
\delta\left(\text { for }\left(y_{1}=y_{2}\right)\{(x, z)\}\right)=\perp \tag{4.1}
\end{equation*}
$$

The core insight here is that neither $y_{1}=y_{2}$ nor $\{(x, z)\}$ can change. Propagating this information - for example, rewriting (for (...) $\perp$ ) to $\perp$ - can simplify derivatives and eliminate expensive for-loops.

Now let's pull out and examine for $\left(\left(y_{2}, z\right) \in t\right)$ for $\left(y_{1}=y_{2}\right)\{(x, z)\}$. The $\phi$ translation is again a no-op, and the $\delta$ translation is:

$$
\begin{array}{rll} 
& \delta\left(\text { for }\left(\left(y_{2}, z\right) \in t\right) \text { for }\left(y_{1}=y_{2}\right)\{(x, z)\}\right) & \\
= & \text { for }\left(\left(y_{2}, z\right) \in d t\right) \phi\left(\text { for }\left(y_{1}=y_{2}\right)\{(x, z)\}\right) & \text { apply } \delta \text { and omit unused lets } \\
& \cup \text { for }\left(\left(y_{2}, z\right) \in t \cup d t\right) \delta\left(\text { for }\left(y_{1}=y_{2}\right)\{(x, z)\}\right) & \\
= & \text { for }\left(\left(y_{2}, z\right) \in d t\right) \text { for }\left(y_{1}=y_{2}\right)\{(x, z)\} & \text { use equation 4.1, propagate } \perp
\end{array}
$$

Thus:

$$
\begin{equation*}
\delta\left(\text { for }\left(\left(y_{2}, z\right) \in t\right) \text { for }\left(y_{1}=y_{2}\right)\{(x, z)\}\right)=\text { for }\left(\left(y_{2}, z\right) \in d t\right) \text { for }\left(y_{1}=y_{2}\right)\{(x, z)\} \tag{4.2}
\end{equation*}
$$

Tackling the outermost for loop:

$$
\begin{array}{rlr} 
& \delta\left(\text { for }\left(\left(x, y_{1}\right) \in s\right) \text { for }\left(\left(y_{2}, z\right) \in t\right) \text { for }\left(y_{1}=y_{2}\right)\{(x, z)\}\right) & \\
= & \text { for }\left(\left(x, y_{1}\right) \in d s\right) \phi\left(\text { for }\left(\left(y_{2}, z\right) \in t\right) \text { for }\left(y_{1}=y_{2}\right)\{(x, z)\}\right) & \text { apply } \delta(\text { for } \ldots) \\
\cup & \text { for }\left(\left(x, y_{1}\right) \in s \cup d s\right) \delta\left(\text { for }\left(\left(y_{2}, z\right) \in t\right) \text { for }\left(y_{1}=y_{2}\right)\{(x, z)\}\right) & \\
= & \text { for }\left(\left(x, y_{1}\right) \in d s\right) \text { for }\left(\left(y_{2}, z\right) \in t\right) \text { for }\left(y_{1}=y_{2}\right)\{(x, z)\} & \text { use equation } 4.2 \\
\cup \text { for }\left(\left(x, y_{1}\right) \in s \cup d s\right) \text { for }\left(\left(y_{2}, z\right) \in d t\right) \text { for }\left(y_{1}=y_{2}\right)\{(x, z)\} & \\
= & (d s \bullet t) \cup((s \cup d s) \bullet d t) & \text { rewrite using } \bullet
\end{array}
$$

This, then, is the derivative of relation composition:

$$
\begin{equation*}
\delta(s \bullet t)=(d s \bullet t) \cup((s \cup d s) \bullet d t) \tag{4.3}
\end{equation*}
$$

Distributing composition over union, this is equivalent to $(d s \bullet t) \cup(s \bullet d t) \cup(d s \bullet d t)$, which is perhaps the derivative a human would give.

Let's use this to figure out $\phi$ (trans [e]). Working inside out, we start with the derivative of the loop body, $\delta(e \cup(e \bullet p))$ :

$$
\begin{aligned}
& \delta(e \cup(e \bullet p)) \\
= & \delta e \cup \delta(e \bullet p) \\
= & \delta e \cup(\delta e \bullet p) \cup((e \cup \delta e) \bullet d p \\
= & \perp \cup(\perp \bullet p) \cup((e \cup \perp) \bullet d p) \\
= & e \bullet d p
\end{aligned}
$$

$$
=\delta e \cup(\delta e \bullet p) \cup((e \cup \delta e) \bullet d p) \quad \text { use equation } 4.3
$$

$\delta e$ is a zero change; replace with $\perp$ propagate $\perp$

Note that the penultimate step here requires a new optimization: by definition $\delta e=d e$, but since $e$ is discrete we know de is a zero change, so we may safely replace it by $\perp$, as it will have the same effect. Thus:

$$
\begin{equation*}
\delta(e \cup(e \bullet p))=e \bullet d p \tag{4.4}
\end{equation*}
$$

Putting everything together, we have:

$$
\begin{array}{rlr} 
& \phi(f \text { fix } P \text { is } e \cup(e \bullet p) & \\
= & \phi(\text { fix }[\lambda p . e \cup(e \bullet p)]) & \text { desugaring } \\
= & \operatorname{semifix} \phi[\lambda p . e \cup(e \bullet p)] & \\
= & \operatorname{semifix}[(\phi(\lambda p . e \cup(e \bullet p)), \delta(\lambda p . e \cup(e \bullet p)))] & \\
= & \operatorname{semifix}[((\lambda p . e \cup(e \bullet p)),(\lambda[p] . \lambda d p . e \bullet d p))] & \text { use equation } 4.4
\end{array}
$$

Examining the recurrence produced by this use of semifix, we recover the seminaïve transitive closure algorithm from §3.1:

$$
\begin{array}{rlrl}
x_{0} & =\perp & x_{i+1} & =x_{i} \cup d x_{i} \\
d x_{0} & =(\lambda p . e \cup(e \bullet p)) \perp=e & d x_{i+1} & =(\lambda[p] . \lambda d x . e \bullet d p)\left[x_{i}\right] d x_{i}=e \bullet d x_{i}
\end{array}
$$

### 4.2 Implementation

To put the seminaïve transformation presented in chapter 3 to the test, we have implemented it as part of a compiler and runtime for a fragment of Datafun (omitting sum types and patternmatching), available at https://github.com/rntz/datafun/tree/popl20/v4-fastfix. In §4.2.1 we sketch the compiler's front-to-back structure, from Datafun source code through several intermediate stages to Haskell output. In $\S 4.2 .2$ we explore how an example Datafun program gets compiled, exhibit the small auxiliary library necessary to run the compiled outputs, and explain why we chose Haskell as a target language. Finally, in $\S 4.2 .3$ we perform some simple benchmarks to test for the expected efficiency gains.


Figure 4.I Stages of the Datafun compiler

### 4.2.1 The compiler structure

The compiler proceeds in several passes, shown as arrows in figure 4.1. It begins with fairly standard parsing and typechecking phases. For parsing, we use OCaml's Menhir library. ${ }^{1}$ The typechecker is bidirectional (Pierce and Turner, 2000) rather than performing full inference, for simplicity of implementation. After these, we have a choice of paths.

The simplest approach is to translate Datafun fairly directly into Haskell. Many of Datafun's features have direct parallels in Haskell, including sum, product, and function types; Datafun's sets are implemented using Haskell's Data. Set module; Datafun's semilattice types are implemented as a Haskell typeclass; and Datafun's fixed points can be implemented by naïve bottom-up iteration. (We discuss these implementation details in §4.2.2.)

One feature of Datafun that does not translate straightforwardly into Haskell is the discreteness comonad $\square$. However, its only purpose is to track (non-)monotonicity; at runtime, $\square A$ may as well be $A$. Thus before emitting Haskell we "drop boxes", eliminating $\square$ and its related term syntax by rewriting $[e] \rightsquigarrow e$ and let $[x]=e$ in $f \rightsquigarrow$ let $x=e$ in $f$ and dropping the distinction between discrete and monotone variables. This takes us to an intermediate language, unimaginatively dubbed "IL", supporting Datafun's computational features but lacking its modal typing, which we translate directly into Haskell.

Thus naïve compilation of Datafun code takes a straight path through figure 4.1:


For seminaïve evaluation, however, we must apply the $\phi$ transform from chapter 3 to the

[^16]typed syntax before dropping boxes, since the transform works precisely by annotating boxed terms with their zero-changes. As we have no further use for $\square$, the compiler pass implementing $\phi$ omits boxes from its output. Thus after applying the $\phi$ transform we can directly emit Haskell.

As we saw in $\S 4.1$, however, the $\phi$ transform produces code that can often be simplified by replacing certain expressions by $\perp$. Therefore we implement two optimization passes:

1. We propagate $\perp$ by applying the following rewrites to the IL:

$$
\begin{aligned}
e \vee \perp & \rightsquigarrow e \\
\text { for }(x \in e) \perp & \rightsquigarrow \perp \\
\text { let } x=\perp \text { in } e & \rightsquigarrow e\{x \mapsto \perp\} \\
(\perp, \perp) & \rightsquigarrow \perp
\end{aligned}
$$

$$
\begin{aligned}
\perp \vee e & \rightsquigarrow e \\
\text { for }(x \in \perp) e & \rightsquigarrow \perp \\
\text { let } x=e \text { in } \perp & \rightsquigarrow \perp \\
\pi_{i} \perp & \rightsquigarrow \perp
\end{aligned}
$$

The most important rewrite here is for $(x \in e) \perp \rightsquigarrow \perp$ : to evaluate the left hand side directly, we evaluate $e$ and iterate over the resulting set, which takes work proportional to its size; but evaluating the right hand side takes constant work. This is where the asymptotic speedups originate. The other rewrites are useful primarily because they enable this one.
2. To make $\perp$-propagation more effective, we first insert $\perp$ in place of semilattice-valued zero changes. To this end, the $\phi$ transform pass explicitly marks certain terms produced by $\delta$ which are guaranteed to be zero-changes, namely:

$$
\begin{aligned}
\delta x & =\mathrm{d} x \quad(\text { for discrete } x) \\
\delta[e] & =() \\
\delta\left\{e_{i}\right\}_{i} & =\delta\left(e_{1}=e_{2}\right)=\delta(\text { fix } e)=\delta \perp=\perp
\end{aligned}
$$

The insert $\perp$ pass replaces these marked expressions with $\perp$ when they have semilattice type, along with some compound terms guaranteed to produce zero-changes:

$$
\begin{array}{ll}
\text { let } x=e \text { in } f & \text { when } f \text { is a zero change } \\
x & \text { when } x \text { is let-bound to a zero change } \\
e_{1} e_{2} e_{3} & \text { when } e_{1}, e_{3} \text { are both zero changes }
\end{array}
$$

In case it is not clear, the last case corresponds to the derivative of function application, $\delta(e f)=\delta e[\phi f] \delta f$, when neither the function nor the argument are changing.

To test whether insert $\perp$ is actually useful, we also implement a do nothing pass for comparison, which simply ignores the zero-change annotations produced by the $\phi$ transform. Altogether, we have three new paths through the compiler. First, after parsing and typechecking we we can apply the seminaïve transform without further optimizations:

$$
\text { Typed syntax } \xrightarrow{\phi \text { transform }} \text { IL with zero } \xrightarrow{\text { do nothing }} \text { IL } \xrightarrow{\text { emit }} \text { Haskell }
$$

Second, we can optimize the output of the $\phi$ transform by propagating $\perp$ :


Or finally, we can replace semilattice zero changes with $\perp$ to make $\perp$ propagation more effective:

$$
\text { Typed syntax } \xrightarrow{\text { ¢ transform }} \text { IL with zero } \xrightarrow{\text { insert } \perp} \text { IL } \xrightarrow{\text { propagate } \perp} \text { IL } \xrightarrow{\text { emit }} \text { Haskell }
$$

### 4.2.2 Compiling transitive closure

To understand the compiler's behavior more concretely, let's consider what it does to a Datafun program implementing transitive closure:

```
trans: \(\square\left\{\mathcal{E Q}_{\varepsilon \in}^{A} \times \underset{E Q}{A}\right\} \rightarrow\{\underset{\in Q}{A} \times \underset{\varepsilon \in}{A}\}\)
trans [edge] \(=\) fix \(P\) is edge \(\vee\{(x, z) \mid(x, y) \in\) edge, \((!y, z) \in P\}\)
```

This code needs a few changes for the compiler to accept it. First, we must replace $\underset{\varepsilon}{ } \mathcal{E}_{\mathrm{\varepsilon}}$ with a specific, concrete type, as the compiler does not support polymorphism. Second, the compiler does not support pattern-matching, so we must replace box-patterns with let-unboxing, tuple-patterns with projections, and equality patterns !y with equality tests:

```
trans : \(\square\{\) string \(\times\) string \(\} \rightarrow\{\) string \(\times\) string \(\}\)
trans \(=\lambda \mathrm{E}\).
    let [edge] \(=\mathrm{E}\) in
    fix \(P\) is
        edge \(\vee\left\{\left(\pi_{1} a, \pi_{2} b\right) \mid a \in e d g e, b \in P, \pi_{2} a=\pi_{1} b\right\}\)
```

In the ASCII syntax the compiler accepts, this becomes:

```
@([\{str,str\}] -> \{str,str\}) -- type annotation
\e.
    let [edge] = e in
    fix \(p\) is
        edge or \(\{(\) pi1 \(a, ~ p i 2 b)\) for \(a\) in edge for \(b\) in \(p\) when pi2 \(a=p i 1 b\}\)
```

Passing this through the naïve compilation path, parse $\rightarrow$ typecheck $\rightarrow$ dropboxes $\rightarrow$ emit, produces closely corresponding Haskell code (also shown in figure 4.4):

```
\e_0 ->
    let edge_1 = e_0 in
    fix (\p_2 ->
        union edge_1
            (forIn edge_1 (\a_3 ->
            forIn p_2 (\b_4 ->
            guard (snd a_3 == fst b_4)
            (set [(fst a_3, snd b_4)])))))
```

This code has been prettified by removing unnecessary parentheses and adding line breaks and indentation. Besides minor syntactic details, the primary changes are:

1. Variable names have had unique numeric suffixes added; this is an artifact of the compiler internals.
2. Boxes have been dropped, so let [edge] = e becomes simply let edge_1 = e_0.
3. The set comprehension is translated into nested calls to forIn, guard, and set. This uglifies the code, replacing binding forms with higher-order functions applied to lambdas, but otherwise corresponds to desugaring the comprehension (figure 2.2):

$$
\begin{aligned}
& \quad\left\{\left(\pi_{1} a, \pi_{2} b\right) \mid a \in \text { edge, } b \in P, \pi_{2} a=\pi_{1} b\right\} \\
& \xrightarrow{\text { desugar }} \text { for }(a \in \text { edge }) \text { for }(b \in P) \text { for }\left(\_\in \pi_{2} a=\pi_{1} b\right)\left\{\left(\pi_{1} a, \pi_{2} b\right)\right\}
\end{aligned}
$$

4. In a similar way, other Datafun features such as fixed points fix $X$ is $e$ and semilattice join $e_{1} \vee e_{2}$ have been translated into calls to functions like fix and union.

The resulting Haskell code depends on functions (highlighted in pink) supplied by a small auxiliary library (figure 4.2). The primary reason we chose Haskell as a target language is because it simplifies this step of translating language features into calls to library functions: Datafun's concept of a "semilattice type" $L$, over which $\vee, \perp$, fix, and for-loops are parameterized, translates directly to a Haskell typeclass Semilat defined in this auxiliary library. This spares the Datafun compiler the work of determining at which types these primitives are actually invoked and generating the library code necessary for that subset of types.

This Semilat typeclass has two core methods, (<:) and unions: ${ }^{2}$

```
class Semilat a where
    (<:) :: a -> a -> Bool
    unions :: [a] -> a
    ... etc ...
```

The ( $<$ :) operator computes the type's order relation $\leqslant,{ }^{3}$ and is used to determine when fixed point iteration has stabilized: seminaïve iteration stabilizes once $d x_{i} \leqslant x_{i}$. (This is an optimization over checking equality $x_{i}=x_{i+1}$.) The unions method computes the semilattice join/least upper bound of its list of arguments. Using this we define binary union $/ \checkmark$ and nullary empty/ $\perp$ wrappers. We provide instances for empty and binary products, booleans, and sets, the Haskell types which our Datafun semilattice types get translated into.

The other functions provided by the runtime library are set, used to construct literal sets; forIn, which implements for-loops over sets; guard, which implements for-loops over booleans (i.e. one-sided conditionals for $\left(e_{1}\right) e_{2}$, also seen as boolean "guard" conditions in set comprehensions); and fix/semifix, which implement naïve and seminaïve iteration respectively. ${ }^{2}$

### 4.2.3 Benchmarking seminaïve evaluation

To test whether the $\phi$ translation can produce the asymptotic performance gains we claim, we benchmark two example Datafun programs:

[^17]```
module Runtime (Set, Semilat (..), set, guard, forIn,
    fix, semifix, semifixMinimized) where
import qualified Data.Set as Set
import Data.Set (Set)
class Semilat a where
    (<:) :: a -> a -> Bool
    unions :: [a] -> a
    empty :: a
    empty = unions []
    union :: a -> a -> a
    union x y = unions [x,y]
    diff :: a -> a -> a -- Law: union a (diff da a) = union a da
    diff dx x = dx -- always lawful but not always efficient
instance Semilat () where
    () <: () = True
    unions _ = ()
instance (Semilat a, Semilat b) => Semilat (a,b) where
    (a,x) <: (b,y) = a <: b && x < : y
    unions ts = (unions lefts, unions rights) where (lefts, rights) = unzip ts
    diff (da,db) (a,b) = (diff da a, diff db b)
instance Semilat Bool where
    x <: y = not x || y
    unions = or
instance Ord a => Semilat (Set a) where
    (<:) = Set.isSubsetOf
    unions = Set.unions
    diff = Set.difference
set :: Ord a => [a] -> Set a
set = Set.fromList
guard :: Semilat a => Bool -> a -> a
guard c x = if c then x else empty
forIn :: Semilat b => Set a -> (a -> b) -> b
forIn set f = unions [f x | x <- Set.toList set]
fix :: Semilat a => (a -> a) -> a
fix f = loop empty
    where loop x = if x' <: x then x else loop x'
        where x' = f x
semifix, semifixMinimized :: Semilat a => ((a -> a), (a -> a -> a)) -> a
semifix (f, df) = loop empty (f empty)
    where loop x dx = if dx <: x then x else loop (union x dx) (df x dx)
semifixMinimized (f, df) = loop empty (f empty)
    where loop x dx = if dx <: x then x else
                    let x' = union x dx in loop x' (df x dx `diff` x')
```

Figure 4.2 Datafun's runtime library

1. Finding the transitive closure of a linear graph using the trans function from §4.2.2 (first introduced in §2.2.1). As discussed in §3.1, transitive closure has a well understood asymptotic speed-up under seminaïve evaluation. This means that if we've failed to capture the essence of seminaïve evaluation, it should be highly visible.
2. Finding all matches of the regular expression / $a * /$ in the string $a^{n}$, using the regex combinators from §2.2.2. Finding all matches for /a*/amounts to finding the reflexive, transitive closure of the matches of $/ a /$, and on $a^{n}$ these form a linear graph. Thus this is a close analogue of our first example, but written in a higher-order style, as we represent regular expressions as functions and regex constructors as function combinators. We chose this example to test whether our extension of seminaïve evaluation properly handles Datafun's distinctive feature: higher-order programming.

We send each program through the four compiler paths described in §4.2.1: the direct/naïve translation (naïve); the $\phi$ transform alone (seminaïve raw); $\phi$ with $\perp$ propagation (seminaïve simplified); and $\phi$ with $\perp$ insertion and propagation (seminaïve optimized). We exhibit the four translations of trans in figures 4.4-4.7. We graph the benchmark results in figure 4.3, separately showing the running times as well as the speedup factor over naïve evaluation (on a logarithmic plot). The results support two conclusions:

1. The $\phi$ transformation combined with optimizations enables asymptotic speedups: seminaïve optimized is dramatically faster than naïve, and the speedup factor increases with the input size. ${ }^{4}$ Moreover, the measured times are similar for transitive closure and regex search across all optimization levels, suggesting that higher-order code does not pose a particular problem for our optimizations.
2. These asymptotic improvements depend on $\perp$ propagation: seminaïve raw yields only a small constant-factor speedup over naïve, roughly $20 \%$. However, the measured times for seminaïve simplified and seminaïve optimized are effectively identical, so $\perp$ insertion appears to be irrelevant, even (somewhat surprisingly) in higher-order code.

As alluded to in the previous section, these asymptotic speedups come from avoiding wasteful loops (for $\left.\left(x \in e_{1}\right) e_{2}\right)$ where $e_{1}$ grows as our input grows but $e_{2}$ always produces $\perp$. The $\phi / \delta$ translations alone do not accomplish this: both $\phi($ for $(x \in e) \ldots$ ) and $\delta($ for $(x \in e) \ldots$ ) produce loops that iterate over at least every $x \in \phi e$. Consulting our logical relation (figure 3.6) at set type, we see that $e$ and $\phi e$ compute identical sets, therefore the number of iterations never shrinks. For instance, in the seminaïve raw translation of trans (figure 4.5) the derivative passed to semifix contains the following wasteful loop:

```
forIn (union p_2 dp_2) ( \(\mathrm{Db}_{-} 4\)->
    let db_4 = ((), ()) in
    if (snd a_3 == fst b_4) then set [] else
    guard False
    (union (set [(fst a_3, snd b_4)]) (set [])))
```

4 Although faster than naïve evaluation, seminaïve optimized is still asymptotically quite slow in these benchmarks. On transitive closure, for example, doubling the graph size from 160 to 320 nodes yields a slowdown factor of $\frac{.407}{.054} \approx 7.54$ ! However, since there are quadratically many paths and we find all of them, the best possible runtime is $\mathrm{O}\left(\mathrm{n}^{2}\right)$. Moreover, our nested-loop relational joins are roughly a factor of n slower than optimal, so we expect $\mathrm{O}\left(\mathrm{n}^{3}\right)$ behavior, which predicts a slowdown of $2^{3}=8$, reasonably close to 7.54 .


|  | GRAPH SIZE / STRING LENGTH |  |  |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | 120 | 140 | 160 | 180 | 200 | 220 | 240 | 260 | 280 | 300 | 320 |
| REGEX SEARCH, NAÏVE | 1.665 | 2.940 | 4.924 | 7.872 | 12.251 | 18.789 | 27.828 | 48.393 | 69.337 | 96.371 | 131.300 |
| transitive Closure, naïve | 1.568 | 2.843 | 4.797 | 7.756 | 11.944 | 18.521 | 29.415 | 47.446 | 67.845 | 95.142 | 128.403 |
| REGEX SEARCH, SEMINAÏVE RAW | 1.317 | 2.410 | 4.047 | 6.568 | 9.909 | 14.840 | 21.636 | 39.629 | 57.017 | 80.622 | 109.707 |
| TRANSITIVE CLOSURE, SEMINAÏVE RAW | 1.275 | 2.351 | 4.040 | 6.429 | 9.880 | 14.656 | 22.886 | 39.007 | 56.686 | 79.837 | 109.552 |
| REGEX SEARCH, SEMINAÏVESIMPLIFIED | 0.024 | 0.037 | 0.055 | 0.079 | 0.107 | 0.141 | 0.182 | 0.228 | 0.279 | 0.347 | 0.407 |
| TRANSITIVE CLOSURE, SEMINAÏVESIMPLIFIED | 0.022 | 0.035 | 0.054 | 0.077 | 0.103 | 0.138 | 0.187 | 0.220 | 0.271 | 0.333 | 0.395 |
| REGEX SEARCH, SEMINAÏVE OPTIMIZED | 0.023 | 0.036 | 0.057 | 0.079 | 0.113 | 0.142 | 0.181 | 0.227 | 0.283 | 0.355 | 0.416 |
| TRANSITIVE CLOSURE, SEMINAÏVE OPTIMIZED | 0.022 | 0.035 | 0.054 | 0.081 | 0.102 | 0.137 | 0.174 | 0.216 | 0.272 | 0.336 | 0.407 |

FIGURE 4.3 Naïve vs seminaïve evaluation of transitive closure and regex matching in Datafun

```
\e_0 ->
    let edge_1 = e_0 in
    fix (\p_2 ->
        union edge_1
            (forIn edge_1 (\a_3 ->
                forIn p_2 (\b_4 ->
                guard (snd a_3 == fst b_4)
                (set [(fst a_3, snd b_4)])))))
```

FIGURE 4.4 Naïve translation of transitive closure

```
\e_0 ->
    let (edge_1, dedge_1) = e_0 in
    semifix
        ((\p_2 -> union edge_1
                            (forIn edge_1 (\a_3 ->
                            forIn p_2 (\b_4 ->
                                guard (snd a_3 == fst b_4)
                                (set [(fst a_3, snd b_4)]))))),
            (\p_2 -> \dp_2 ->
            union dedge_1
                (union
                                (forIn dedge_1 (\a_3 ->
                            let da_3 = ((), ()) in
                            forIn p_2 (\b_4 ->
                        guard (snd a_3 == fst b_4)
                            (set [(fst a_3, snd b_4)]))))
                                (forIn (union edge_1 dedge_1) (\a_3 ->
                        let da_3 = ((), ()) in
                    union
                            (forIn dp_2 (\b_4 ->
                            let db_4 = ((), ()) in
                            guard (snd a_3 == fst b_4)
                            (set [(fst a_3, snd b_4)])))
                                (forIn (union p_2 dp_2) (\b_4 ->
                            let db_4 = ((), ()) in
                            if (snd a_3 == fst b_4) then set [] else
                    guard False
                    (union (set [(fst a_3, snd b_4)]) (set [])))))))))
```

Figure 4.5 Raw seminaïve translation of transitive closure

```
\e_0 ->
    let (edge_1, dedge_1) = e_0 in
    semifix
        ((\p_2 -> union edge_1
                            (forIn edge_1 (\a_3 ->
                            forIn p_2 (\b_4 ->
                                guard (snd a_3 == fst b_4)
                            (set [(fst a_3, snd b_4)]))))),
        (\p_2 -> \dp_2 ->
        union dedge_1
            (union
                (forIn dedge_1 (\a_3 ->
                    let da_3 = ((), ()) in
                forIn p_2 (\b_4 ->
                guard (snd a_3 == fst b_4)
                (set [((fst a_3), (snd b_4))]))))
                (forIn (union edge_1 dedge_1) (\a_3 ->
                    let da_3 = ((), ()) in
                    forIn dp_2 (\b_4 ->
                    let db_4 = ((), ()) in
                    guard (snd a_3 == fst b_4)
                    (set [(fst a_3, snd b_4)])))))))
```

FIGURE 4.6 Seminaïve translation of transitive closure with $\perp$ propagation

```
\e_0 ->
    let (edge_1, dedge_1) = e_0 in
    semifix
        ((\p_2 -> union edge_1
                            (forIn edge_1 (\a_3 ->
                            forIn p_2 (\b_4 ->
                            guard (snd a_3 == fst b_4)
                            (set [(fst a_3, snd b_4)]))))),
        (\p_2 -> \dp_2 ->
        forIn edge_1 (\a_3 ->
        let da_3 = ((), ()) in
        forIn dp_2 (\b_4 ->
        let db_4 = ((), ()) in
        guard (snd a_3 == fst b_4)
        (set [(fst a_3, snd b_4)])))))
```

FIGURE 4.7 Seminaïve translation of transitive closure with $\perp$ insertion and propagation

Each branch of the if-statement computes an empty set, the first branch explicitly and the else-branch via (guard False ...). The $\perp$ propagation pass recognizes this and rewrites the entire loop to $\perp$, removing it from the simplified and optimized translations (figures 4.6 and 4.7).

### 4.3 Change minimization

Before concluding that we have captured the essence of seminaive evaluation, let's try a small twist on our running example: let's add self-loops to every node in our linear graph, producing the graph $(V, E)$ with $V=\{1, \ldots, n\}$ and $E=\{(i, j) \mid j \in\{i, i+1\}\}$. This makes our reachability relation reflexive, changing the transitive closure from the less-than relation $\{(\mathfrak{i}, \mathfrak{j}) \mid 1 \leqslant \mathfrak{i}<\mathfrak{j} \leqslant n\}$ to the less-than-or-equal-to relation $\{(i, j) \mid 1 \leqslant \mathfrak{i} \leqslant \mathfrak{j} \leqslant \mathfrak{n}\}$. This produces exactly $n$ new paths, namely $\{(\mathfrak{i}, \mathfrak{i}) \mid 1 \leqslant \mathfrak{i} \leqslant n\}$; since we already had quadratically many paths, ideally this won't affect our performance much.

Unfortunately, adding these self-loops produces an asymptotic slowdown, even with our seminaïve transformation and all optimizations applied (à la seminaïve optimized):


What's going on here? Recall from §4.1, page 67 (and confirmed by figure 4.7) that the seminaïve iteration strategy Datafun uses for transitive closure is:

$$
\begin{aligned}
x_{0} & =\emptyset & x_{i+1} & =x_{i} \cup \mathrm{~d} x_{i} \\
\mathrm{~d} x_{0} & =\text { edges } & \mathrm{d} x_{i+1} & =\text { edges } \bullet d x_{i}
\end{aligned}
$$

The key computation step here is $\mathrm{d} x_{i+1}=$ edges $\bullet \mathrm{d} x_{\mathrm{i}}$. In other words, we prepend edges out of each "frontier" $\mathrm{d} x_{i}$ to get the next frontier $\mathrm{d} x_{i+1}$. Ideally, each frontier consists of pairs $(x, y)$ newly discovered to be reachable; by accumulating them into $x_{i}=\bigcup_{j<i} d x_{j}$ we find all such pairs. In a linear graph without self-loops, as we saw in §3.1, this strategy discovers each reachable pair exactly once, because $d x_{i}$ captures paths of length exactly $i$, and each reachable pair corresponds to a unique path. But if our edge relation is reflexive, any path from $x$ to $y$ can be adjoined to a self-loop to find a longer path from $x$ to $y$; thus $d x_{i} \subseteq d x_{i+1}$. In turn this means that $d x_{i}=x_{i+1}$; by adding self-loops we've regressed to naïve evaluation!

Taking a logical perspective, at step $i$, naïve evaluation finds all derivations of depth $d \leqslant i$, while the "seminaïve" strategy we've presented so far finds only derivations of depth $d=i$. This is a clear improvement, but sometimes the same fact may be derived at multiple depths - as in our loopy linear graph, where derivation depth is path length. We care only about whether a fact has a derivation, so anything after the first (shallowest) derivation is redundant.

From an incremental computation perspective, this is a problem of unnecessarily large changes. Our seminaïve strategy looks for "new" ways to derive a tuple $(x, y)$, based on whatever was "newly" derived in the previous step. But our notion of "new" is a bit lax, because our derivatives are allowed to be imprecise. Our strategy for finding a fixed point of a function $f:\left\{A,\left\{\begin{array}{c}\text { FN }\end{array}\right\} \rightarrow\right.$ is:

$$
\begin{aligned}
x_{0} & =\emptyset & x_{i+1} & =x_{i} \cup d x_{i} \\
d x_{0} & =\mathrm{f} \emptyset & d x_{i+1} & =\mathrm{f}^{\prime} x_{i} d x_{i}
\end{aligned}
$$

For sets, the derivative property guarantees that $f x_{i} \cup f^{\prime} x_{i} d x_{i}=f x_{i+1}$, but not that $f^{\prime} x_{i} d x_{i}=f x_{i+1} \backslash f x_{i}$. This is exploited in, among others, the derivative rule $\delta\left(e_{1} \cup e_{2}\right)=\delta e_{1} \cup \delta e_{2}$. If $\delta e_{1}$ and $e_{2}$ intersect (or $\delta e_{2}$ and $e_{1}$ intersect), this generates an overly large change.

A more precise derivative would be $\delta\left(e_{1} \cup e_{2}\right)=\left(\delta e_{1} \backslash e_{2}\right) \cup\left(\delta e_{2} \backslash e_{1}\right)$. However, this does more work, not less: it does not avoid computing "old" elements $x \in e_{1} \cup e_{2}$, but rather discards them after-the-fact. Indeed, discovering something twice because it has two different derivations seems in general unavoidable; in graph reachability, for instance, how can we know two different paths lead to the same destination except by following them?

So if computing these overly large changes actually takes less work, where does the asymptotic slowdown originate? It happens because rediscovering a reachable pair ( $x, y$ ) at iteration $i$ causes redundant work in all subsequent iterations, because it is included in $\mathrm{d} x_{\mathrm{i}}$ (treated as "new") and used to compute $\mathrm{d} x_{\mathrm{i}+1}=\mathrm{f}^{\prime} x_{\mathrm{i}} \mathrm{d} x_{\mathrm{i}}$. Consequently, $\mathrm{d} x_{i+1}$ will include re-derivations of anything the presence of $(x, y)$ makes "newly" deducible; and so on in $d x_{i+2}, d x_{i+3}$, etc.

While we may not be able to avoid all rediscovery, we can avoid these unnecessary changes accumulating across iterations - and the resulting asymptotic wastefulness - by minimizing our changes. Let's change our strategy for finding the "new" frontier $\mathrm{d} \mathrm{x}_{i+1}$ to remove anything that's already in $x_{i+1}$ :

$$
\begin{aligned}
\text { for transitive closure } & d x_{i+1} & =\left(\text { edges } \bullet d x_{i}\right) \backslash x_{i+1} \\
\text { or more generally } & d x_{i+1} & =\left(f^{\prime} x_{i} d x_{i}\right) \backslash x_{i+1}
\end{aligned}
$$

This ensures each $d x_{i}$ is minimal, disjoint from $x_{i}$ and thus all prior $d x_{j}$ for $j<i$. (We don't need to do anything to minimize $d x_{0}$ since $x_{0}=\perp$.) This fixes our asymptotic slowdown:


If we examine all four options - with and without self-loops, with and without minimizing $\mathrm{d} x_{i}$ - and compare their slowdown factors, taking a loopless graph without change minimization


|  | GRAPH SIZE VS SECONDS TO EVALUATE |  |  |  |  |  |  |  |  |
| :--- | ---: | ---: | ---: | ---: | ---: | ---: | ---: | ---: | ---: |
|  | 120 | 140 | 160 | 180 | 200 | 220 | 240 | 260 | 280 |
| NOT MINIMIZING, LOOPY | 3.310 | 5.909 | 9.881 | 16.032 | 23.860 | 35.866 | 55.938 | 81.851 | 117.876 |
| MINIMIZING, LOOPY | 0.048 | 0.079 | 0.113 | 0.162 | 0.213 | 0.288 | 0.359 | 0.449 | 0.562 |
| MINIMIZING, LOOPLESS | 0.027 | 0.040 | 0.060 | 0.082 | 0.115 | 0.149 | 0.191 | 0.242 | 0.294 |
| NOT MINIMIZING, LOOPLESS | 0.024 | 0.036 | 0.055 | 0.077 | 0.104 | 0.135 | 0.177 | 0.222 | 0.274 |
|  |  |  |  |  |  |  |  |  |  |
| NOT MINIMIZING, LOOPY | 171.830 | 251.552 | 349.668 | 461.830 | 598.083 | 745.815 |  |  |  |
| MINIMIZING, LOOPY | 0.684 | 0.816 | 0.959 | 1.153 | 1.367 | 1.545 |  |  |  |
| MINIMIZING, LOOPLESS | 0.355 | 0.428 | 0.504 | 0.595 | 0.704 | 0.809 |  |  |  |
| NOTMINIMIZING, LOOPLESS | 0.332 | 0.401 | 0.490 | 0.558 | 0.672 | 0.779 |  |  |  |

FIGURE 4.8 The effect of self-loops and change minimization on seminaïve transitive closure on a line graph
as our baseline (figure 4.8), we find that (a) loopy graphs without change minimization are asymptotically slow; (b) on loopless graphs, minimizing changes has low constant overhead (versus not minimizing); and (c) moving from a loopless to a loopy graph causes a roughly 2 x slowdown when minimizing changes, because we must remove rediscovered paths on every iteration.

Two questions remain: (1) why is change minimization correct? and (2) how can we generalize it from finite sets to all semilattice types? As it happens, the answer to the first question also illuminates the second. Change minimization is correct because it preserves validity of changes: if $d x \triangleright x \hookrightarrow y:\left\{\mathcal{A}_{\varepsilon}\right\}$, in other words $x \cup d x=y$, then we also have $\mathrm{d} x \backslash x \triangleright x \hookrightarrow y:\{\underset{\varepsilon}{ }\}$, because $x \cup(\mathrm{~d} x \backslash x)=x \cup \mathrm{~d} x=y$. Thus minimizing changes preserves the inductive invariant that $\mathrm{d} \mathrm{x}_{\mathrm{i}} \triangleright \mathrm{x}_{\mathrm{i}} \hookrightarrow \mathrm{f} \mathrm{x}_{\mathrm{i}}$ which guarantees semifix finds a least fixed point.

This condition tells us how to generalize our approach: for each lattice type $L$, we require a change minimization operator $(\backslash)_{L}: L \rightarrow L \rightarrow L$ such that if $d x \triangleright x \hookrightarrow y: L$ then $\mathrm{d} x \backslash_{\mathrm{L}} \mathrm{x} \triangleright \mathrm{x} \hookrightarrow \mathrm{y}: \mathrm{L} .{ }^{5}$ In our runtime library (figure 4.2), we accomplish this by adding a diff method to our Semilat typeclass:

```
class Semilat a where
    diff :: a -> a -> a
    diff dx x = x
```

This method admits a default implementation, $d x \_{L} x=d x$, which trivially satisfies our correctness condition. Although generic, this is inefficient, for it degenerates to the original non-minimizing implementation of semifix:

$$
d x_{i+1}=\left(f^{\prime} x_{i} d x_{i}\right) \backslash_{L} x_{i+1}=f^{\prime} x_{i} d x_{i}
$$

Instead, we want $d x \_{L} x$ to be as small as possible (whatever that means for our runtime representation of $L$ ) to reduce the work done by any operators applied after change minimization. So we provide more practical implementations for finite sets and product types:

$$
d x \backslash_{\left\{\left\{_{\varepsilon}\right\}\right.} x=d x \backslash x \quad(d x, d y) \backslash_{L \times M}(x, y)=\left(\left(d x \backslash_{L} x\right),\left(d y \backslash_{M} y\right)\right)
$$

In our Haskell runtime this becomes:

```
instance Ord a => Semilat (Set a) where
    diff = Set.difference
instance (Semilat a, Semilat b) => Semilat (a,b) where
    diff (da,db) (a,b) = (diff da a, diff db b)
```

Unfortunately, for some semilattices the degenerate default is the only valid change minimization operator. This includes any totally-ordered semilattice, such as $\mathbb{N}_{\text {min }}^{\infty}$, the naturals extended with positive infinity under minimum, which is useful for shortest path computations. This inability to meaningfully minimize changes is concerning for the efficiency of computations which use these semilattices; an important direction for future work would be to characterize the worst-case efficiency of seminaïve evaluation over different classes of semilattice.

[^18]
## Chapter 5

## Related Work

### 5.1 Logic, higher-order abstraction, and semilattices

Datafun's value proposition is to extend bottom up logic programming à la Datalog with two additional features: higher-order functional abstraction and support for semilattices other than finite sets. There is a good deal of work on combining logic programming with one or the other of these features, and at least one other system - Flix - which, like Datafun, proposes to combine all three. We'll first briefly consider the systems which feature one or the other and then move on to a more detailed comparison with Flix.

Higher-order extensional logic programming There are many possible approaches to combining the power of logic programming and of higher-order abstractions. The most direct approach would be to directly extend logic programming with support for higher-order relations. Unfortunately, this quickly entangles one with thorny problems of decidability and efficiency. Nonetheless, a line of work starting with Wadge (1991) has explored this approach. Kountouriotis et al. (2005) investigate extending Datalog's bottom-up approach with higherorder relations, an approach very close to Datafun's; however, while they present a prototype implementation, they remark that it can be impractically slow for significantly higher-order programs because it needs to synthesize many large relations. Later approaches move from a bottom-up to a top-down approach, becoming less Datalog and more Prolog (Charalambidis et al., 2013). By contrast, Datafun attempts to reduce implementation difficulties by narrowing the scope of higher-order computation to functions, leaving relations first-order and decidable - higher-order functional programming and decidable first-order relational programming being both well-trodden areas.

Higher-order functional + top-down logic programming Of course, Datafun is not the first language to attempt to add higher-order features to logic programming by combining it with functional programming: Mercury (Somogyi et al., 1994) integrates higher-order functional programming into a logical paradigm, while Curry (Antoy and Hanus, 2010) cleanly combines the lazy functional language Haskell with top-down logic programming. The main difference is that Mercury and Curry's logical features are inspired by Prolog, employing top-down search and unification, while Datafun is inspired by Datalog, uses bottom-up enumeration, and imposes deliberate restrictions to ensure termination and avoid Turing-completeness.

Embedding database queries in functional languages Datalog has sometimes been described as "relational algebra plus fixed points", and there is a long line of work on embedding database query languages into general-purpose languages, including pioneering efforts such as Machiavelli (Ohori et al., 1989) and Kleisli (Wong, 2000), as well as more recent systems such as Ferry (Grust et al., 2009) and LINQ in C\# (Cheney et al., 2013). The focus of this work has been on embedding query languages based on relational algebra into general purpose languages, with an emphasis on statically compiling higher-order queries into the first-order queries supported by existing database systems (Cheney et al. (2014) is a representative example). Datafun's approach is different: rather than embed Datalog into a general purpose language, Datafun is also a "little language", albeit one that happens to be a higher-order functional language. We have not attempted to embed Datafun into an existing language, as this would greatly complicate the context-management operations needed to ensure monotonicity.

Logic with semilattices We know of two systems which extend Datalog with support for semilattices without incorporating higher-order abstraction. IncA $_{\mathrm{L}}$ (Szabó et al., 2018) is an incremental Datalog engine with support for custom semilattices, aimed at static analysis. Bloom ${ }^{\text {L }}$ (Conway et al., 2012) is a Datalog-inspired language aimed at distributed computing, where monotonicity is used to ensure eventual consistency (Alvaro et al., 2011) and custom semilattices are used to extend the range of types to which monotonicity analysis can be fruitfully applied. Although both of these are clearly related to Datafun's goals, neither includes the crucial ingredient of higher-order abstraction that leads to most of Datafun's unique capacities, as well as its unique design and implementation challenges.

### 5.1.1 Flix

Flix (Madsen et al., 2016) and Datafun both augment Datalog with higher-order functional programming and semilattice types, but they go about this merger of logic and functional programming in different ways. Datafun embeds Datalog's semantics into a functional language by adding first-class support for finite sets and monotone fixed points, ensuring monotonicity via a custom type system. Flix, however, is really two languages in a trenchcoat - one logical and Datalog-inspired, the other functional. These language halves interoperate closely: the logic fragment can use semilattices whose merge functions are defined in the functional fragment, and with a recent extension (Madsen and Lhoták, 2020) the functional fragment can manipulate first-class values representing groups of logic-fragment rules ("Datalog constraints") which can be composed and evaluated at run-time.

Our main contribution compared to Flix is to demonstrate that this separation between functional and logic layers is unnecessary; they can be smoothly integrated by paying close attention to semantics, and standard Datalog optimizations such as seminaïve evaluation can be generalized to operate on functional languages, with the unexpected side-benefit of revealing a theory of higher-order incremental monotone computation. The practical flip side of this theoretical contribution is that Flix, because it separates functional from logic, can reuse existing techniques for implementing and optimizing Datalog without needing to reinvent them in a higher-order setting.

Flix and Datafun also have different approaches to monotonicity: Flix does not check monotonicity via types, but monotonicity is nonetheless important for the interoperation of the
logical and functional fragments; functions invoked in certain positions in the logic fragment must be monotone to ensure that a fixed point exists. To this end, a verification toolchain has been developed for FLIx which employs SMT solving and symbolic execution to check properties, including monotonicity but also safety and soundness of static analyses (Madsen and Lhoták, 2018).

The advantage of checking these properties with such high-powered machinery is that it better supports user-defined posets and semilattices: in Datafun, adding a new datatype with a custom ordering is a matter for the language designer, but a Flix programmer may do it for themself, if they can convince the verification machinery. The disadvantage is that these verification techniques have more "black-box" behavior; they are less predictable and reliable (in terms of resource usage, error message quality, and code accepted) than a compositional type system. Nonetheless, for certain applications such as static analysis, the use of user-defined semilattice types is highly desirable. We believe a fruitful direction for future research would be to hybridize Datafun's monotonicity type system with lightweight verification techniques. In our ideal language, types with custom orderings would be defined in a modular, encapsulated fashion. Verification would take place inside the module, to ensure the interface it exposes has the properties (such as monotonicity) it claims, but a compositional type-system would handle code external to the module, which acts as a client of this verified interface.

### 5.2 Incremental computation

In chapter 3 we presented seminaive evaluation in Datalog and Datafun as a matter of incrementalizing the inner loop of our fixed points to avoid recomputation. Our approach was to only compute the changes between iterations, to which end we used a static transformation to push changes through expressions in our language. However, this is only one of many approaches to incremental computation. In this section we'll examine other approaches and how they relate to ours.

The core problem of incremental computation is to handle change while minimising work. Most approaches to incremental computation are built around one of two insights into how to do this: dependency tracking or finite differencing.

Dependency tracking The simplest way to avoid doing unnecessary work is to avoid reexecuting code if the data it depends on hasn't changed, and re-use its previous result instead. Generally these dependencies are represented by some sort of dependency graph. One of the oldest and most familiar applications of this idea is the build system Make and its many relatives. Perhaps even more ubiquitously, this strategy is used by spreadsheet software such as Microsoft Excel, where cells' contents may be (re)computed using the content of other cells, and so on recursively (Mokhov et al., 2020). A long line of work starting with self-adjusting computation (Acar, 2005; Acar et al., 2002) and descendant systems such as Adapton (Hammer et al., 2014) apply dependency tracking to general-purpose programming languages by tracing execution to construct a dynamic dependency graph.

It would be an intriguing line of future work to examine whether a dependency tracking approach, probably building on prior work such as SAC or Adapton, would be useful for computing Datafun's monotone fixed points. However, Datafun was inspired by Datalog, and
dependency tracking is not the approach taken by Datalog's seminaïve evaluation. Instead, it uses finite differencing.

Finite differencing Dependency tracking approaches incrementalization with a yes-or-no mindset: did our dependencies change? Suppose instead we ask the question: how did our dependencies change? By analyzing this difference we may be able to compute the resulting difference to our output. For constant-time operations over atomic data, like adding two 64-bit numbers, this may not be any faster than simple re-execution. But for bulk operations over collection types, where changes can be much smaller than the original data, differencing is a more natural approach.

For example, you could incrementally sum a list of changing numbers using a dependency graph - ideally a balanced tree, so updates take $O(\log n)$ steps. But with direct access to the difference, when an element changes you can simply add the difference to the previous sum. This is $\mathrm{O}(1)$, easily extended to handle new or removed elements, and doesn't require balancing or rebalancing a tree.

However, this example works only because addition on the integers commutes and associates. ${ }^{1}$ This is what allows us to combine a sum $\sum_{i} x_{i}$ with a difference $d x$ to the $j^{\text {th }}$ element and find the updated sum:

$$
d x+\sum_{i} x_{i}=\sum_{i} \begin{cases}x_{i}+d x & \text { if } \mathfrak{i}=\mathfrak{j} \\ x_{i} & \text { otherwise }\end{cases}
$$

Unlike the fairly generic dependency-graph strategy, efficient derivatives depend crucially on the structure of the modification to the input. Foundational work by Paige and Koenig (1982) takes this literally, differentiating set-valued expressions with respect to lines of program code modifying a depended-on variable. More recent work represents these modifications (the "differences" of "finite differencing") as values of a datatype equipped with some kind of algebraic structure. For instance, DBToaster incrementally maintains SQL queries using rings (Koch, 2010, 2013); Differential Dataflow uses groups (McSherry et al., 2013); recent work on incremental Datalog uses monoid actions (Alvarez-Picallo et al., 2019); and Datafun uses semilattices.

Since finite differencing is advantageous for operations over large collections, but requires algebraic insight that makes it non-obvious how to apply it to arbitrary programs, it should be no surprise that many of the examples just cited come from the database research community, which often deals with structured operations on bulk data. However, this approach has recently crossed over into the PL research community by way of the incremental $\lambda$-calculus, and it is from this line of work that Datafun takes inspiration.

### 5.2.1 The incremental $\lambda$-calculus

The incremental $\boldsymbol{\lambda}$-calculus was introduced by Cai, Giarrusso, Rendel, and Ostermann (2014) and further developed by Giarrusso, Régis-Gianas, and Schuster (2019) and in Giarrusso's PhD thesis (2020). We briefly discussed our adaptation of it in $\S 3.2$, but here we give a slightly
${ }^{1}$ More precisely, associativity and commutativity suffice because our aggregation (summation) is the same as the operator that applies our differences (addition). If we were taking the maximum of the list instead of its sum, this strategy would not work, even though maximum itself commutes and associates.
fuller comparison (though still only a summary) between our formulation, the original, and its further developments.

## Change structures

The insight of the incremental $\lambda$-calculus that is it is possible to extend the differencing approach to higher-order computation by finding an appropriate notion of change for functions. To this end, the incremental $\lambda$-calculus associates each type with a change structure capturing how values of that type may change, and how to represent these changes. The precise formal definition of a change structure differs between the various versions of the incremental $\lambda$-calculus. A good starting point is the notion of a basic change structure introduced in Giarrusso's PhD thesis (2020, chapter 12, definition 12.1.1): a basic change structure on a set $S$ consists of a change set $\Delta S$ and a validity relation $d x \triangleright x_{1} \hookrightarrow x_{2}: S$ indicating that $d x: \Delta S$ is a valid change from the base points $x_{1}: S$ to $x_{2}: S$.

For example, we can endow the naturals $\mathbb{N}$ with a basic change structure by letting changes be integer differences $\Delta \mathbb{N}=\mathbb{Z}$ and letting $d x \triangleright x_{1} \hookrightarrow x_{2}: \mathbb{N} \Longleftrightarrow x_{1}+d x=x_{2}$. To handle higher-order computation, Giarrusso endows function types $A \rightarrow B$ with a basic change structure as follows (definition 12.1.8): let $\Delta(A \rightarrow B)=A \rightarrow \Delta A \rightarrow \Delta B$ and define the validity relation by saying that a valid function change $d f \triangleright f_{1} \hookrightarrow f_{2}: A \rightarrow B$ is one which takes an argument $x: A$ and valid change to it $d x \triangleright x_{1} \hookrightarrow x_{2}: A$ to a valid output change df $x_{1} d x \triangleright f_{1} x_{1} \hookrightarrow f_{2} x_{2}$ : B, that is, a change between the original function applied to its original argument $f_{1} x_{1}$ and the updated function applied to an updated argument $f_{2} x_{2}$.

Giarrusso goes on to define more elaborate "full" change structures (definition 13.1.1) which additionally possesses operators $x \oplus d x$ for updating a base point by a change, $y \ominus x$ for finding a change between two points, 0 for finding a zero change from a point to itself, and $d x \odot d y$ for composing two changes. ${ }^{2}$

Datafun's change structures (definition 14) lie somewhere between Giarrusso's basic and "full" change structures, modified to handle monotonicity: in Datafun S (or VS in our notation) and $\Delta S$ are not sets but posets. For example, our change structures for functions almost coincide, except that because of Datafun's monotonicity typing we must let $\Delta(A \rightarrow B)=$ $\square \mathrm{A} \rightarrow \Delta \mathrm{A} \rightarrow \Delta \mathrm{B}$, using $\square$ to allow function changes to be non-monotone with respect to the base point. The root divergence here is one of goals: in Datafun we are not trying to respond to arbitrary changes to our whole program's input, but only to incrementalize the inner loops of fixed points to calculate them more efficiently. Because these fixed points are monotone, in Datafun we need only handle increasing change (enforced by our soundness condition). The price of this simplification is that we must pay careful attention to the interaction of incrementalization with Datafun's monotonicity-checking modal type system in our program transformation and its proof of correctness.

Because of the limited way Datafun uses incremental computation, we only need some of the operators from Giarrusso's full change structures, and only at certain types. We use 0

[^19]explicitly in the $\phi$ and $\delta$ translations of loops, for $(x \in e)$ f, to supply the zero change for the elements $x$; but since these elements are always of first-order type $\underset{E R}{A}$, we only need to compute zero changes for first order values. We implicitly use $\oplus$ in restricted form in the implementation of seminaïve fixed points, to combine the value $x_{i}$ of the $i^{\text {th }}$ iteration with its corresponding change $d x_{i}$. However, fixed points are always at first-order lattice types $\underset{f 1 \times}{\mathrm{L}}$, and because of the way our change types are constructed $\oplus$ at these types is simply $\vee$. Even more subtly, we use $\ominus$ in the same place, to find the kick-off change between the first iteration $\mathrm{x}_{1}=\mathrm{f} \perp$ and the zeroth $\mathrm{x}_{0}=\perp$; but, again because of the way these change types are constructed, $\mathrm{f} \perp \ominus \perp$ is simply $\mathrm{f} \perp$. These simplifications are fortuitous, because the interaction of the fully general versions of these operators with monotonicity typing presents several difficulties. ${ }^{3}$

## The derivative translation

Although change structures allow us to specify the notion of a derivative that takes input changes to output changes, they do not by themselves tell us how to find such a derivative. The incremental $\lambda$-calculus and Datafun both accomplish this by static differentiation: we give source-to-source translations from a program to its derivative, essentially by decomposing a program into primitive operations which we know how to differentiate and recombining these using an analogue of the chain rule.

Cai et al. (2014) call this translation $\mathcal{D e r i v e}$, while Datafun calls it $\delta$. Datafun's approach was directly inspired by Cai et al., and where their features overlap the two translations nearly coincide. For example, the derivative of function application is:

$$
\begin{aligned}
\operatorname{Derive}\left(e_{1} e_{2}\right) & =\operatorname{Derive}\left(e_{1}\right) e_{2} \operatorname{Derive}\left(e_{2}\right) \\
\delta\left(e_{1} e_{2}\right) & =\delta e_{1}\left[\phi e_{2}\right] \delta e_{2}
\end{aligned}
$$

Besides notation there are two differences here, which are indicative of the differences from the incremental $\lambda$-calculus more generally: (1) Datafun is concerned with monotonicity, and so since the function changes may be non-monotone in their first argument we need to box it; and (2) besides $\delta$, Datafun has another term translation $\phi$ which speeds up execution by executing fixed points seminaïvely, and for technical reasons these translations must be mutually-recursive; wherever $\mathcal{D e r i v e}$ uses a part of the original term it is given, $\delta$ instead uses its $\phi$-translation.
${ }^{3}$ For starters, in general $x \ominus y$ may not exist unless $x \geqslant y$ since in Datafun we only support increasing changes. For another example, it is difficult to define the $\oplus$ operator internally in a way that respects monotonicity typing at higher type. Recall that in Datafun all functions are monotone and $\Delta(A \rightarrow B)=\square A \rightarrow \Delta A \rightarrow \Delta B$. The natural definition of $\oplus_{A \rightarrow B}$ would seem to be:

$$
\begin{aligned}
\oplus_{A \rightarrow B} & :(A \rightarrow B) \times(\square A \rightarrow \Delta A \rightarrow \Delta B) \rightarrow(A \rightarrow B) \\
(f \oplus d f) & =\lambda X . f X \oplus d f[X](O X)
\end{aligned}
$$

[^20]While Datafun adds complications in the form of monotonicity typing and the seminaïve transformation, later work on the incremental $\lambda$-calculus adds complications of its own, extending it to handle the untyped $\lambda$-calculus and therefore nontermination; to prove correctness, they use a step-indexed logical relation (Giarrusso et al., 2019). They also address the problem of caching intermediate results, but in order to explain this problem and its relevance to Datafun, it will help to briefly revisit the idea of dependency tracking.

## Dependency tracking as a change structure

We started this section by comparing dependency tracking to finite differencing, observing informally that finite differencing generalizes dependency tracking by asking not merely "did our dependencies change?" but "how did our dependencies change?" This insight can be formalized using the incremental $\lambda$-calculus's change structures, as there is a simple generic change structure which captures the question "did it change?". Allowing ourselves to dip into pseudo-Haskell for a moment, consider the parameterized type Update A defined by:

```
data Update A = OLD | NEW A
```

Any type $A$ may be endowed with a basic change structure by letting $\Delta A=$ Update $A$ and letting $\mathrm{dx} \triangleright \mathrm{x} \hookrightarrow \mathrm{y}: A$ be defined by:

$$
\text { OLD } \triangleright x \hookrightarrow x: A \quad \text { NEW } y \triangleright x \hookrightarrow y: A
$$

This change structure has the wonderful property of trivializing differentiation; one valid derivative of $f: A \rightarrow B$ is simply:

$$
\begin{aligned}
& \mathrm{f}^{\prime} \times O L D=O L D \\
& \mathrm{f}^{\prime} \times(\text { NEW } y)=\operatorname{NEW}(\mathrm{f} y)
\end{aligned}
$$

The only complication is handling multi-argument functions: since Update $(A \times B) \neq$ Update $A \times$ Update $B$, a function taking multiple arguments $g: A \rightarrow B \rightarrow C$ needs a slightly more interesting derivative:

$$
\begin{aligned}
& \mathrm{g}^{\prime}: A \rightarrow \text { Update } \mathrm{A} \rightarrow \mathrm{~B} \rightarrow \text { Update } \mathrm{B} \rightarrow \text { Update } \mathrm{C} \\
& \mathrm{~g}^{\prime} \text { a OLD b OLD }=\text { OLD } \\
& g^{\prime} \text { _ (NEW a) b OLD }=\operatorname{NEW}(g a b) \\
& g^{\prime} a \operatorname{OLD} \quad-(N E W b)=\operatorname{NEW}(g a b) \\
& g^{\prime} \quad(\text { NEW } a)-(\text { NEW } b)=\operatorname{NEW}(g a b)
\end{aligned}
$$

The general strategy is to rerun the original function if any of its arguments change, reusing the previous value for arguments that did not change. This is precisely the strategy behind dependency tracking.

## Caching intermediate results

We have observed that dependency tracking's re-execution strategy can be seen as a special case of finite differencing. However, one thing all dependency tracking systems do is store intermediate results between runs so they can reuse them if they don't need to be recomputed
because their dependencies haven't changed. Thus far our presentation of the incremental $\lambda$ calculus (or indeed of Datafun) has not mentioned caching intermediate results. This presents an issue; although our goal is to translate input changes into output changes, in general computing the output difference may require both the input difference and the original input. For example, in the incremental $\lambda$-calculus the derivative of a function $f: A \rightarrow B$ has type $A \rightarrow \Delta A \rightarrow \Delta B$, taking the original argument $A$ as well as its change $\Delta A$. Where does this original argument come from?

By default, both Datafun and the incremental $\lambda$-calculus as originally introduced in Cai et al. (2014) recompute these arguments. For example, recall the derivatives of function application in each system:

$$
\begin{aligned}
\operatorname{Derive}\left(e_{1} e_{2}\right) & =\operatorname{Derive}\left(e_{1}\right) e_{2} \operatorname{Derive}\left(e_{2}\right) \\
\delta\left(e_{1} e_{2}\right) & =\delta e_{1}\left[\phi e_{2}\right] \delta e_{2}
\end{aligned}
$$

These expressions recalculate the original argument $e_{2}$ (or in the case of Datafun, its spedup version $\phi e_{2}$ ). On its own, recomputation is a losing strategy: the whole point of finite differencing is to compute the change more efficiently than by recomputing the output! Luckily, sometimes we do not need this original input. A simple example is summing a list of changing numbers: the change to the sum is simply the sum of the changes to each element; the original list is not necessary to compute the change. Or, for a natural example in the context of Datafun, fix a binary relation edge and consider two different functions, consing and appending, defined as follows (recall that $R \bullet S$ stands for relation composition):

$$
\begin{aligned}
\text { consing path } & =\text { edge } \cup(\text { edge } \bullet \text { path }) \\
\text { appending path } & =\text { edge } \cup(\text { path } \bullet \text { path })
\end{aligned}
$$

The fixed point of either function computes the transitive closure of edge: consing by extending paths one edge at a time, and appending by appending paths together. Now let's take a look at these functions' derivatives: ${ }^{4}$

```
    consing' path dpath \(=\) edge \(\bullet\) dpath
appending' path dpath \(=(\) path \(\bullet\) dpath \() \cup(\) dpath \(\bullet\) path \() \cup(\) dpath \(\bullet\) dpath \()\)
```

Observe that consing', unlike appending ${ }^{\prime}$, does not need its first argument path, representing the original argument value.

Following the incremental $\lambda$-calculus we call functions whose derivatives do not depend on their original input, like consing or the sum of a list, self-maintainable. Because the transformation in Cai et al. (2014) does not cache intermediate results, it is really only suitable for programs composed primarily of self-maintainable functions, where the recomputation of these unused original arguments can be optimized out. To handle non-self-maintainable

4 To obtain these derivatives, let $\delta$ edge $=\varnothing$ since we assume the edge-set is fixed and apply the rules:

$$
\begin{aligned}
\delta(R \cup S) & =\delta R \cup \delta S \\
\delta(R \bullet S) & =(R \bullet \delta S) \cup(\delta R \bullet S) \cup(\delta R \bullet \delta S)
\end{aligned}
$$

behavior, follow-up work by Giarrusso et al. (2019) extends the derivative translation to cache intermediate results.

Datafun takes a simpler approach. We cache intermediate results in exactly one place: the implementation of semifix. Recall that, to compute the fixed point of a function $f$, semifix computes the sequences $x_{i}, \mathrm{~d} x_{i}$ defined by:

$$
\begin{aligned}
x_{0} & =\perp & x_{i+1} & =x_{i} \vee d x_{i} \\
\mathrm{~d} x_{0} & =\mathrm{f} \perp & \mathrm{~d} x_{i+1} & =\mathrm{f}^{\prime} x_{\mathrm{i}} \mathrm{~d} x_{\mathrm{i}}
\end{aligned}
$$

Since $x_{i+1}$ and $d x_{i+1}$ depend only on their immediate predecessors, we need exactly two pieces of state to produce this sequence: the previous iteration $x_{i}$ and its change $d x_{i}$. This is our only cache; unless $f$ is self-maintainable, any intermediate values it requires will be recomputed by $d x_{i+1}=f^{\prime} x_{i} d x_{i}$. Serendipitously, in practice most step functions are either self-maintainable or do not compute expensive intermediate results.

For instance, consider implementing transitive closure as the fixed point of either consing or appending. We have already observed that consing is self-maintainable. ${ }^{5}$ Even though appending is not, however, it does not require extensive recomputation:

$$
\mathrm{d} x_{i+1}=\text { appending }^{\prime} x_{i} \mathrm{~d} x_{i}=\left(x_{i} \bullet d x_{i}\right) \cup\left(d x_{i} \bullet x_{i}\right) \cup\left(\mathrm{d} x_{i} \bullet d x_{i}\right)
$$

All intermediate results in this expression (for example, $x_{i} \bullet d x_{i}$ ) depend upon $d x_{i}$; there is no work that could be saved by caching them, because the cache would be invalidated immediately. We conjecture that this holds so often for the programs we have examined because these fixed point step functions are essentially unions of (possibly many-way) relational joins. (Indeed, in Datalog this is baked into the language syntax!) It's not too hard to see that the derivative of a union of joins is itself a union of joins, and each component join will depend on at least one changing relation. So the question reduces to whether relational joins can be incrementalized efficiently without caching intermediate results. In the case of binary joins, at least, the answer is yes. Pursuing this conjecture further we leave to future work.

### 5.2.2 The monoidal approach to change

The incremental $\lambda$-calculus has its notion of a change structure; Datafun has another; but these do not exhaust the space of possibilities. A line of work starting with Alvarez-Picallo et al. (2019), and most thoroughly expounded in Mario Alvarez-Picallo's thesis (2020) has explored representing change structures using monoid actions, which they call change actions. A change action on a set $A$ consists of a monoid ( $\Delta A,+_{A}, O_{A}$ ) and a monoid action $\oplus_{A}: A \times \Delta A \rightarrow A$. As before we interpret $\oplus$ as applying a change to a base value. The monoid action law $x \oplus(d x+d y)=(x \oplus d x) \oplus d y$ says that + composes changes (corresponding to $\odot$ in the incremental $\lambda$-calculus); and the other law $x \oplus 0=x$ makes 0 a zero change to any value. The primary differences from the incremental $\lambda$-calculus are:

1. The lack of a $\ominus$ operator, thus allowing for incomplete change structures, where it may not be possible to find a change from one value to any other.

[^21]2. The requirement that there exists a single value $0: \Delta A$ which acts as a universal zero change, rather than an operator $0: A \rightarrow \Delta A$ that finds a zero change to a particular value.
3. Change actions lack a validity relation: every change must be valid for every base point

Tantalizingly, Alvarez-Picallo et al. (2019) explicitly use this notion of change action to give a differentiation/incrementalization transformation for Datalog programs. Might this monoidal approach work in Datafun as well? At first it seems as though it might: many of Datafun's change structures, in particular those on semilattice types, fit into this structure: for instance,
 $0=\varnothing$.

However, points (2) and (3) above produce problems when considering the change structure for functions. The incremental $\lambda$-calculus and Datafun both take function changes to be generalizations of function derivatives, such that the zero-change to a function is its derivative. But this is incompatible with requiring a universal zero-change $0: \Delta(A \rightarrow B)$; there is no "universal derivative". For this reason both Alvarez-Picallo et al. (2019) and Alvarez-Picallo (2020) use pointwise function changes $\Delta(A \rightarrow B)=A \rightarrow \Delta B$, letting $(f \oplus d f) x=f x \oplus d f x$. This fundamental divergence from the incremental $\lambda$-calculus would require redesigning the $\phi / \delta$ transformations - neither the 2019 paper nor Alvarez-Picallo's thesis give an explicit derivative transformation for a higher-order language.

However, there is a deeper issue: in a monotonicity-aware context like Datafun, we must choose whether the pointwise change functions are required to be monotone, $\Delta(A \rightarrow B)=$ $A \rightarrow \Delta B$, or allowed to be non-monotone, $\Delta(A \rightarrow B)=\square A \rightarrow \Delta B$. Both choices have serious problems:

Non-monotone changes If we allow function changes to be non-monotone we have an immediate problem: the updated function $f \oplus d f=x \mapsto f x \oplus d f x$ is not guaranteed to be monotone. The only way to repair this without requiring that the change itself be monotone would seem to be to re-introduce the incremental $\lambda$-calculus's notion of validity, and say that $d f$ is only a valid change to $f$ if $f \oplus d f$ remains monotone. This would require a considerable elaboration of the theory of change actions.

Monotone changes Unfortunately, insisting that pointwise function changes be monotone makes it impossible to differentiate some perfectly reasonable functions. For example, take the integers $\mathbb{Z}$ equipped with the natural change action $\Delta \mathbb{Z}=\mathbb{Z}, \oplus_{\mathbb{Z}}=+_{\mathbb{Z}}=$ ,$+ 0_{\mathbb{Z}}=0$, and consider the function expression $\lambda y . \max (x, y): \mathbb{Z} \rightarrow \mathbb{Z}$. Now suppose $x$ increases from 0 to 1 ; how does this function change in response? In other words, what is the change between $\max (0, \ldots)$ and $\max (1, \ldots)$ ? Tabulating its values for $y=0,1,2, \ldots$ we can clearly see it is not monotone:

$$
\begin{array}{r|llll}
y & 0 & 1 & 2 & \ldots \\
\max (0, y) & 0 & 1 & 2 & \ldots \\
\max (1, y) & 1 & 1 & 2 & \cdots \\
\max (1, y)-\max (0, y) & 1 & 0 & 0 & \cdots
\end{array}
$$

Thus, forcing function changes to be monotone with respect to the base point is a very limiting approach.

It is possible that by combining change actions with a validity relation, thus allowing the monoid action to be partial, one could achieve a synthesis of the change action approach and Datafun's higher-order monotonicity. We leave this to future work.

## Chapter 6

## Looking Back and Forward

To the extent that we have had in this dissertation a thesis, a singular statement we have aimed to demonstrate, it is that we can seamlessly integrate Datalog's features into a typed higher-order functional language by deconstructing them semantically. Or, as we said in the introduction (page 13):

> The goal of this thesis is to design a language which improves on Datalog's ability to express monotone fixed point computation over semilattices by finding ways to lift Datalog's restrictions without sacrificing either its simple semantics or its practical implementation strategies.

Unfortunately, we cannot claim to have definitively proven our thesis: we have made a start towards this goal, but much remains to be done.

The examples in chapter 2 show Datafun can at least express Datalog-style queries. Moreover in §1.4 we listed four things Datalog's restrictions do not permit: (1) functional abstraction, (2) semilattices other than set union, (3) arithmetic, user-defined functions, and aggregation, and (4) compound data. Of these, Datafun makes functional abstraction, arithmetic, user-defined functions, and compound data straightforward. It does not include semilattices other than sets (and products of sets), nor aggregations other than semilattice aggregation (for-loops). However, its design lays a clear foundation for such extensions: new semilattices can be added as semilattice types, and aggregations can be added as primitive higher-order functions.

In addition to lifting Datalog's restrictions, we also wished to preserve two desirable qualities: simple semantics and practical implementation strategies. On each count, we have achieved only qualified success.

Datafun possesses a simple denotational semantics that captures Datalog's ability to manipulate finite relations: Datalog's recursively-defined relations become Datafun's bottomup fixed points, and Datalog's stratification is enforced by Datafun's monotonicity types. However, least fixed points also require an ascending chain condition; here Datafun and Datalog diverge. Datalog makes a clear distinction between relations and the terms they range over, and enforces constructor-freedom: programs may not construct new terms not present in the source program. This ensures a finite universe of terms, and thus an ascending chain condition for relations over this universe.

By contrast, in Datafun relations and terms are simply types of values; this adds flexibility but requires a different way to guarantee the ascending chain condition. In theory we require
the type at which we take a fixed point to satisfy the ACC; for set types, this requires the element type be finite. In practice we have hand-waved this condition away - for instance, our regular expression combinators from $\S 2.2 .3$ use fix at type $\{i n t\}$, representing sets of indices into a string. Although valid indices into a particular string form a finite set, the integer type int is infinite. Semantically, this is a serious flaw in the foundations of our approach. Practically, it is on par with existing approaches: real Datalog engines routinely permit constructors or arithmetic, putting the onus on the programmer to avoid infinite relations and non-termination.

As for practical implementation strategies, we have constructed a Datafun implementation supporting seminaïve evaluation, a central Datalog technique without which recursive relations are impractical to compute; but this required a significant novel development of theory, and represents only one of many techniques necessary for an efficient implementation. It is plausible, but hardly certain, that other standard techniques - in particular query planning and optimization (necessary to replace nested for-loops with efficient relational joins) and demand transforms such as magic sets (which make queries over large recursively defined relations practical by computing only a smaller relevant subset of the relation) - could be extended to Datafun; this remains future work.

### 6.1 Directions forward

In this section we sketch some directions for future work on Datafun to address the shortcomings identified in the previous section.

The ascending chain condition As we have discussed, Datafun's semantics for fix require ACC; our motivating examples satisfy this in principle, but Datafun's simple type system cannot capture this. For instance, we might represent the nodes of a graph as integers; but while the graph may be finite, the integers are infinite. To remedy this, we must either (a) accept nontermination and adjust our semantics or (b) reject nontermination and adjust our language.

Accepting nontermination is simpler, but it removes potential optimizations by invalidating many program equalities, for instance loop interchange:

$$
\text { for }\left(x \in e_{1}, y \in e_{2}\right) e_{3}=\text { for }\left(y \in e_{2}, x \in e_{1}\right) e_{3} \quad\left(w h e n x, y \text { not used in } e_{1}, e_{2}\right)
$$

This equation fails if $e_{1}$ is nonterminating but $e_{2}=\emptyset$ or vice-versa. Perhaps some adjustment to how Datafun expresses queries like this, such as ditching monadic set-comprehensions in favor of applicative ones (see query planning and optimization below), might resolve this.

Rejecting nontermination requires creatively re-thinking how we guarantee ACC. We might, for instance, capture reasoning about the finiteness of sets at the type level, mechanizing our hand-waving about taking a fixed point over finite sets of nodes rather than of integers. Or, we might try to capture the spirit of Datalog's no-constructor restriction by singling out a class of "uncreative" functions that do not create new data; perhaps by treating "creating data" as an effectful operation, ${ }^{1}$ or perhaps by exploiting parametricity to guarantee all data in our output came from our input.

[^22]Query planning and optimization To implement Datafun efficiently, we need to be able to identify relational joins. Identifying joins is also a necessary first step if we wish to apply standard query planning and optimization techniques. Datafun expresses joins as nested loops; for instance, relational composition (a simple equijoin):


```
\(S \bullet T=\) for \(((a, b) \in S)\) for \(((!b, c) \in T)\{(a, c)\}\)
```

Implementing this as it is written, using a nested loop, has time-complexity $\mathrm{O}(|\mathrm{S}| \cdot|\mathrm{T}|)$. An on-the-fly hash-join, building an index on the $\underset{\in Q}{B}$ column of either $S$ or $T$, takes $O(|S|+|T|+|S \bullet T|)$. Unfortunately, nested for-loop expressions are in general not reducible to joins, because the inner loop can loop over a function of the outer loop variable: (for $(x \in S)$ for $(y \in f(x)) \ldots$ ). This does not happen in Datalog and makes query planning significantly harder: in database parlance, we've expressed joins (easy) as subqueries (hard). We could address this in the compiler by heuristically identifying nested loops which can be implemented as joins; or, we could force our comprehensions to be applicative rather than monadic, banning the problematic nesting entirely.

Moreover, there remains the question of what to do once we have identified joins. We could again attempt to lift standard database techniques to apply to a higher-order setting; or, we could attempt to sidestep this work by compiling Datafun into an existing Datalog dialect by using some variety of defunctionalisation.

Aggregations and the Boom hierarchy Aggregations can be added to Datafun as primitive higher-order functions and pose no semantic issues so long as their types properly capture their (non-)monotonicity. For instance, consider counting and summation:

$$
\begin{aligned}
\text { size } & :\left\{\begin{array}{l}
A \\
A
\end{array}\right\} \rightarrow \text { int } \\
\text { size } s & =|s|
\end{aligned}
$$

$$
\begin{aligned}
\text { sum } & :(\square A \rightarrow \mathrm{int}) \rightarrow \square\left\{\left\{_{\mathrm{ER}}\right\} \rightarrow \mathrm{int}\right. \\
\operatorname{sum} \mathrm{fs} & =\sum_{x \in s} \mathrm{f}(\mathrm{x})
\end{aligned}
$$

Throwing in each aggregation we need as a primitive function is, however, somewhat ad-hoc; we might unlock a richer approach if we pay attention to the semantics of aggregations. Many aggregations arise from free functors into categories of algebraic structures. For instance, bags (finite multisets) form the free commutative monoid; and given a map from a bag's elements into a commutative monoid $(M,+, 0)$, we can aggregate the bag's contents into $M$ :

$$
\begin{aligned}
\text { bagsum } & :(A \rightarrow M) \rightarrow \text { Bag } A \rightarrow M \\
\text { bagsum } f b & =\sum_{x \in b} f(x)
\end{aligned}
$$

Modulo monotonicity, sum arises from bagsum by letting $M=(\mathbb{Z},+, 0)$. Many common aggregations and some of the most common data structures used in programming arise in a

$$
\text { anoint }: A \rightarrow \text { Bless (Holy } A) \quad \text { disregard : Holy } A \rightarrow A
$$

such that map disregard (anoint $x$ ) = pure $x$. The idea is that Holy $A$ represents the finite subset of values of $A$ which have been anointed. Since functions passed to fix are pure, they cannot anoint new values; so we may treat Holy $A$ as a finite type, and thus (for instance) use fix at the type \{Holy int $\}$ even though we may not at \{int\}. Alas, actually programming with anoint/disregard is an exercise in boilerplate.
similar way from three free functors in particular: lists as the free monoid, bags as the free commutative monoid, and finite sets as the free idempotent commutative monoid (equivalent to a semilattice). These are three layers of the "Boom hierarchy" (Backhouse and Hoogendijk, 1993). ${ }^{2}$ Each free functor forms half of an adjunction between Set and the respective category of algebraic structures (Mon, CMon, and Semilat). Our semantics for Datafun is built over the adjunctions between Set, Poset, and Semilat; can we extend this to cover the other adjunctions of the Boom hierarchy, and uncover a language for expressing queries over these three data structures and their corresponding varieties of aggregation?

### 6.2 Lessons and surprises

Research is a process of discovery: sometimes you find what you expected, sometimes you do not; sometimes you find more questions. This section presents some insights gained and questions raised in the course of writing this dissertation which, while not entirely novel, at least surprised this author.

Change minimization and precise differences The need to minimize changes to avoid junk piling up during seminaïve fixed point iteration and eliminating any asymptotic speed-up is obvious in retrospect (see §4.3), but was overlooked by Arntzenius and Krishnaswami (2020), their reviewers, and the examiners of the initial version of this dissertation. The author only realized it when an undergraduate student pointed out an instance of it in the Q \& A for a talk given at POPL 2020.

The root of this oversight is an early decision, when applying the incremental $\lambda$-calculus to Datafun, to not compute precise changes - in particular, to let $\delta(e \cup f)=\delta e \cup \delta f$ instead of the more precise ( $\delta e \backslash f) \cup(\delta f \backslash e)$, because the former, simpler expression does less work and avoids recomputing $e$ and $f$. In light of the need for change minimization, this decision is questionable. But because our efforts in chapter 3 focused almost entirely on proving correctness, and because our initial benchmark in chapter 4 (a linear graph) has at most one path between two nodes and thus fails to trigger this issue, we failed to notice this inefficiency. Change minimization addresses this issue, but the question remains: might it be better to just compute precise changes in the first place?

Expressiveness versus tractability There is a tradeoff in programming language design between expressiveness and tractability: the more powerful the language, the more complex it becomes to reason about programs in that language. Datalog's power comes from its limitations: it restricts logic programming to the bare minimum - finite relations - and in return gains a rich theory of implementation and optimisation techniques. Datafun is an attempt to loosen Datalog's restrictions; in hindsight it's clear we might run into issues of tractability.

For instance, we have already seen that by allowing monadic set-comprehensions, a natural choice from a functional programming standpoint and one made without careful consideration on the part of this author, we complicated the task of query planning, ignoring
${ }^{2}$ The other layer of the Boom hierarchy is trees, specifically binary trees with data only at the leaves: the free magma. Few useful aggregations form magmas, and although trees are ubiquitous in computing, few are of this particular form.
hard-won wisdom from the field of databases incorporated implicitly into the design of Datalog.

More generally, by deeply integrating functional and logic programming, Datafun extends Datalog's expressivity - but is it worth the price? There are more conservative ways to address Datalog's shortcomings which may require less reinvention of existing techniques; for instance, using code generation or staged programming to allow modular code re-use; or the approach taken by Flix: two interlocking but separate logic and functional languages. The only examples we have given that deeply intertwine the functional and logic features of Datafun are the regular expression combinators from §2.2.2-2.2.3, which represent regular expressions as functions producing relations and regular expression constructions as higherorder functions (combinators). This is cute, but as justifications for a dissertation's worth of work go it is fairly thin. Higher-order programming sometimes unlocks unusually powerful or concise solutions to existing problems; can we find other motivating examples that take advantage of Datafun's unique feature set?

The diversity and unity of incremental computation The problem of seminaïve evaluation is an instance of incremental computation - one instance among many. As we explored in §5.2, there appear to be as many approaches to incremental computation as there are applications of it, including a zoo of incremental build systems, several varieties of selfadjusting computation, and multiple approaches to incremental maintenance in database systems. Even user interfaces involve incremental computation: for performance reasons, web UI libraries like React avoid re-rendering UI components whose state has not changed. ${ }^{3}$

These diverse systems nonetheless share a small set of core techniques: tracking dependencies so we know what needs to be updated, caching intermediate values to re-use them when they don't change, and propagating changes - perhaps as all-or-nothing updates, perhaps as diffs. This latter apparent difference (between what we called dependency tracking and finite differencing approaches) can, however, be resolved by seeing all-or-nothing updates as a degenerate kind of diff, as we showed in §5.2.1. The theory of change structures originated by Cai et al. (2014) and further developed by Giarrusso (2020), Alvarez-Picallo (2020), and in the present work - hints at the beginnings of a unified theory of incremental computation. However, all these works use slightly but crucially differing notions of change structure; perhaps incremental computation is destined never to be unified at all.

### 6.3 Successes summarized

So far in this chapter we have largely explored how this effort falls short or could be improved on in future work. However, chapters 2 and 3 respectively represent significant successes for this semantic-deconstruction approach; if we were to distill this dissertation into their key ideas, it would be these:

Model monotonicity with modal types. Datalog can be summarized as relational algebra plus stratified recursive queries. Modulo implementation subtleties, relational algebra embeds straightforwardly in a functional language via finite sets and set comprehensions. We

[^23]have shown that stratified recursive queries also embed nicely, so long as we locate our semantics in Poset to capture compositional reasoning about monotonicity. The main difficulty is the interaction of monotone and non-monotone functions; this arises from the discreteness comonad $\square$, and can be handled with a simple modal type system.

To find fixed points faster, incrementalize! Finding a fixed point by iteration involves repeatedly changing a function's input to match its changing output. Doing this naïvely is asymptotically inefficient; to do it efficiently, we must efficiently propagate changes. This is not only the essence of seminaïve evaluation in Datalog, but an instance of a greater problem of automatic incremental computation. Prior work on the incremental $\lambda$-calculus shows that incremental computation can be achieved in higher-order languages; we have extended it to Datafun and shown that by modifying it to consider only increasing changes, it gives rise to seminaïve evaluation.

## Appendix A

## Proofs omitted from main text

We state these lemmas and theorems in dependency order, so that nothing is used before it has been proven. This is not always the order in which they are stated in the text.

## A. 1 Datafun

Theorems and lemmas from chapter 2.

Lemma 31. If $X: A \in \Gamma$ or $x:: A \in \Gamma$ and $\Gamma \sqsubseteq \Delta$ then $X: A \in \Delta$ or $x:: A \in \Delta$.
Proof. Recall that although we write them differently we regard $X$ and $x$ as the same variable. Let H stand for the hypothesis in our assumption, either $\mathrm{X}: \mathrm{A}$ or $\mathrm{x}:: \mathrm{A}$ respectively. Then by induction on the derivation of $\Gamma \sqsubseteq \Delta$ :

Case $\varepsilon \sqsubseteq \varepsilon$. By contradiction, since $\mathrm{H} \in \varepsilon$ is impossible.
Case $\frac{\Gamma^{\prime} \sqsubseteq \Delta^{\prime}}{\Gamma^{\prime}, \mathrm{H}^{\prime} \sqsubseteq \Delta^{\prime}, \mathrm{H}^{\prime}}$. If $\mathrm{H}=\mathrm{H}^{\prime}$ we are done. Otherwise, $\mathrm{H} \in \Gamma^{\prime}$, so apply the IH .
Case $\frac{\Gamma \sqsubseteq \Delta^{\prime}}{\Gamma \sqsubseteq \Delta^{\prime}, \mathrm{H}}$. Apply the IH.
Case $\frac{\Gamma^{\prime} \sqsubseteq \Delta^{\prime}}{\Gamma^{\prime}, X: A \sqsubseteq \Delta^{\prime}, x: A}$. If $H=X: A$ we are done. Otherwise, $H \in \Gamma^{\prime}$, so apply the IH.

Lemma 32. If $\Gamma \sqsubseteq \Delta$ then $\lceil\Gamma\rceil \sqsubseteq\lceil\Delta\rceil$.
Proof. By induction on $\Gamma \sqsubseteq \Delta$ :
Case $\varepsilon \sqsubseteq \varepsilon$. Immediate.
Case $\frac{\Gamma^{\prime} \sqsubseteq \Delta^{\prime}}{\Gamma^{\prime}, \mathrm{H} \sqsubseteq \Delta^{\prime}, \mathrm{H}}$.

Either H is discrete $x:: \mathrm{A}$, in which case $\left\lceil\Gamma^{\prime}, \mathrm{H}\right\rceil=\left\lceil\Gamma^{\prime}\right\rceil, \mathrm{H} \sqsubseteq\left\lceil\Delta^{\prime}\right\rceil, \mathrm{H}=\left\lceil\Delta^{\prime}, \mathrm{H}\right\rceil$ by cons and our inductive hypothesis; or H is monotone $X: A$, in which case $\left\lceil\Gamma^{\prime}, H\right\rceil=$ $\left\lceil\Gamma^{\prime}\right\rceil \sqsubseteq\left\lceil\Delta^{\prime}\right\rceil=\left\lceil\Delta^{\prime}, \mathrm{H}\right\rceil$ by our inductive hypothesis alone.

Case $\frac{\Gamma \sqsubseteq \Delta^{\prime}}{\Gamma \sqsubseteq \Delta^{\prime}, \mathrm{H}}$.
Then $\lceil\Gamma\rceil \sqsubseteq\left\lceil\Delta^{\prime}\right\rceil$ by our inductive hypothesis. Depending upon whether H is monotone or discrete, we have either $\left\lceil\Delta^{\prime}, \mathrm{H}\right\rceil=\left\lceil\Delta^{\prime}\right\rceil$ (in which case our inductive hypothesis suffices) or $\left\lceil\Delta^{\prime}, \mathrm{H}\right\rceil=\left\lceil\Delta^{\prime}\right\rceil, \mathrm{H}$, in which case by cons and transitivity $\lceil\Gamma\rceil \sqsubseteq\left\lceil\Delta^{\prime}\right\rceil, \mathrm{H}$.

Case $\frac{\Gamma^{\prime} \sqsubseteq \Delta^{\prime}}{\Gamma^{\prime}, X: A \sqsubseteq \Delta^{\prime}, x:: A}$.
Then $\left\lceil\Gamma^{\prime}, \mathrm{X}: A\right\rceil=\left\lceil\Gamma^{\prime}\right\rceil$ while $\left\lceil\Delta^{\prime}, x:: A\right\rceil=\left\lceil\Delta^{\prime}\right\rceil, x:: A$. By our $I H$, we have $\left\lceil\Gamma^{\prime}\right\rceil \sqsubseteq$ $\left\lceil\Delta^{\prime}\right\rceil$, and therefore by drop we have $\left\lceil\Gamma^{\prime}\right\rceil \sqsubseteq\left\lceil\Delta^{\prime}\right\rceil, x$ : $A$ as desired.

## A. 2 Seminaïve evaluation

Theorems and lemmas from chapter 3.

Lemma 19. $\Phi \underset{\varepsilon}{\infty} \underset{\varepsilon}{A}=\underset{\varepsilon}{A}$ for all equality types $\underset{\varepsilon}{A}$.
Proof. Induct on $\underset{\varepsilon}{\mathcal{A}}$ applying the equations in figure 3.1, recalling from figure 2.1 that the grammar of equality types is $\underset{\varepsilon Q}{A}:=1|\underset{\varepsilon Q}{A} \times \underset{\varepsilon Q}{\mathcal{A}}| \underset{\varepsilon Q}{A}+\underset{\varepsilon Q}{B} \mid\{\underset{\varepsilon Q}{A}\}$.

Lemma 20. At each semilattice type L , we have $\Delta \mathrm{L}=\mathrm{L}$.
Proof. Induct on L applying the equations in figure 3.1, recalling from figure 2.1 that the grammar of semilattice types is $L::=1\left|L_{1} \times L_{2}\right|\{A\}$.

Theorem 22 (Weakening). If $\Delta \sqsupseteq \Gamma$ and $\Gamma \vdash e: A$ then $\Delta \vdash e: A$.
Proof. By induction on the derivation of $\Gamma \vdash e: A$.
Cases $\frac{X: A \in \Gamma}{\Gamma \vdash X: A} \quad \frac{x:: A \in \Gamma}{\Gamma \vdash x: A} . \quad$ By lemma 31.

Cases $\overline{\Gamma \vdash(): 1} \overline{\Gamma \vdash \perp: L} . \quad$ Trivial.

Cases where the premises have the same context as the conclusion, namely:

$$
\begin{aligned}
& \frac{\Gamma \vdash e: A \rightarrow B \quad \Gamma \vdash f: A}{\Gamma \vdash e f: B} \quad \frac{\left(\Gamma \vdash e_{i}: A_{i}\right)_{i}}{\Gamma \vdash\left(e_{1}, e_{2}\right): A_{1} \times A_{2}} \quad \frac{\Gamma \vdash e: A_{1} \times A_{2}}{\Gamma \vdash \pi_{i} e: A_{i}}
\end{aligned}
$$

Apply the same typing rule to our inductive hypotheses.

Cases where the premises add hypotheses to the context, namely:

$$
\begin{gathered}
\frac{\Gamma, X: A \vdash e: B}{\Gamma \vdash \lambda X . e: A \rightarrow B} \quad \frac{\Gamma \vdash e: \square A \quad \Gamma, x: A \vdash f: B}{\Gamma \vdash \text { let }[x]=e \text { in } f: B} \\
\frac{\Gamma \vdash e: A_{1}+A_{2} \quad\left(\Gamma, X_{i}: A_{i} \vdash f_{i}: B\right)_{i}}{\Gamma \vdash \text { case } e \text { of }\left(\operatorname{in}_{i} X_{i} \rightarrow f_{i}\right)_{i}: B} \quad \frac{\Gamma \vdash e:\{A\} \quad \Gamma, x:: A \vdash f: L}{\Gamma \vdash \text { for }(x \in e) f: L}
\end{gathered}
$$

Apply the inductive hypotheses, using cons when necessary to show that the modified contexts also satisfy our precondition, for example, $\Delta, X: A \sqsupseteq \Gamma, X: A$.

Case where the premises strip the context, namely:

Then $\lceil\Delta\rceil \sqsupseteq\lceil\Gamma\rceil$ by lemma 32, so we apply the inductive hypotheses.

Theorem 21 (Well-typedness of $\phi, \delta$ ). If $\Gamma \vdash e: A$, then $\phi e$ and $\delta e$ have the following types:

$$
\begin{gathered}
Ф Г \vdash \phi e: Ф А \\
Ф Г, \Delta Ф Г \vdash \delta e: \Delta \Phi А
\end{gathered}
$$

Proof. By induction on the derivation of $\Gamma \vdash e: A$, although as we'll see shortly we will need weakening (theorem 22) in some places.

Proof. By induction on $\underset{\mathrm{ER}}{\mathrm{A}}$, applying the definition from figure 3.6:
Case 1. Trivial.

Case $\underset{\varepsilon}{\mathrm{A}} \times \underset{\mathrm{E}}{\mathrm{A}} \times$. Then our assumption is equivalent to

$$
\left(d x_{1}, d x_{2}\right) \triangleright_{\underset{E}{A} \times B_{E Q}}\left(x_{1}, x_{2}\right) \notin\left(a_{1}, a_{2}\right) \rightarrow\left(y_{1}, y_{2}\right) \nLeftarrow\left(b_{1}, b_{2}\right)
$$

 which by our inductive hypotheses show $x_{1}=a_{1}, y_{1}=b_{1}$ and $x_{2}=a_{2}, y_{2}=b_{2}$, which suffices.

Case $\underset{\varepsilon Q 1}{A_{1}}+\underset{\varepsilon_{Q}}{A_{2}}$. Then for some $i \in\{1,2\}$ our assumption is equivalent to
and by unfolding this we have $d x \triangleright_{\substack{A_{i}}} x \nmid a \rightarrow y\{b$, which by our inductive hypothesis implies $x=a$ and $y=b$, which suffices.

Case $\{\underset{\varepsilon \mathrm{q}}{ } \mathrm{A}\}$. Then our assumption unfolds to $(x, y, x \cup d x)=(a, b, y)$, which suffices.

 (figures 3.5 and 3.6).

## Case 1. Trivial.

Case $\underset{E Q}{A} \times \underset{E Q}{\text { B }}$. Letting $x=(y, z)$, we have dummy $x=$ dummy $(y, z)=$ (dummy $y$, dummy $z$ ). By our inductive hypotheses, we have dummy $y \triangleright_{\underset{E}{A}} y\{y \rightarrow y$ y $y$ and likewise for $z$. By definition this shows that
as desired.
Case $\underset{\varepsilon}{ }{\underset{\varepsilon}{1}}^{A_{1}}+{\underset{\varepsilon}{2}}^{A_{2}}$. Without loss of generality we have $x=i_{i} y$ for some $i \in\{1,2\}$. Applying the definition of dummy we have dummy $x=\mathrm{in}_{\mathrm{i}}$ (dummy y ). By our inductive hypothesis we have dummy $y \triangleright_{\mathcal{A}_{i}} y\{y \rightarrow y\{y$, which suffices to show

$$
\mathrm{in}_{i}(\text { dummy } y) \triangleright_{A_{i}} i n_{i} y z i n_{i} y \rightarrow i n_{i} y\left\langle i n_{i} y\right.
$$

as desired.
Case $\left\{\begin{array}{l}\mathrm{A}\} \\ \mathrm{E}\end{array}\right]$. Unfolding our theorem's definition, we need to show that $(x, x, x \cup$ dummy $x)=$


Lemma 28 (Discrete contexts don’t change). If () $\triangleright_{\lceil\Gamma\rceil} \gamma \& \rho \rightarrow \gamma^{\prime} \& \rho^{\prime}$ then $\gamma=\gamma^{\prime}$ and $\rho=\rho^{\prime}$.

Proof. All variables in the stripped contexts are discrete, and therefore the logical relation for discrete variables in contexts, which invokes the logical relation at $\square$ type, requires their corresponding components be equal.

Lemma 29 (Context stripping). If $\mathrm{d} \gamma \triangleright_{\Gamma} \gamma \& \rho \rightarrow \gamma^{\prime} \& \rho^{\prime}$ then

$$
() \triangleright_{\Gamma \Gamma\rceil} \operatorname{strip}_{\Phi \Gamma}(\gamma) \& \operatorname{strip}_{\Gamma}(\rho) \rightarrow \operatorname{strip}_{\Phi \Gamma}\left(\gamma^{\prime}\right) ぬ \operatorname{strip}_{\Gamma}\left(\rho^{\prime}\right)
$$

where strip ${ }_{\Gamma}=\left\langle\pi_{x}\right\rangle_{x:: A \in \Gamma}$ keeps only the discrete variables from a substitution.
Proof. Immediate from the definitions.

Lemma 33 (Applying box). Given $\lceil\rceil \vdash e: A$ and $\gamma: \llbracket\ulcorner\rrbracket$,

$$
\llbracket[e] \rrbracket \gamma=\operatorname{box}_{\Gamma}(\llbracket e \rrbracket)(\gamma)=\llbracket e \rrbracket\left(\operatorname{strip}_{\Gamma}(\gamma)\right)
$$

Proof. Recall that the box comonad $\square$ 's functorial action, duplication map $\delta_{\mathrm{A}}$ :$\square A \rightarrow$
 and distribution dist ${ }_{\square}^{\times}: \prod_{i} \square A_{i} \rightarrow \square \prod_{i} A_{i}$ are all no-ops. Then:

$$
\begin{aligned}
\llbracket[e \rrbracket \rrbracket \gamma & =\operatorname{box}_{\Gamma}(\llbracket e \rrbracket)(\gamma) & & \text { definition of } \llbracket[e \rrbracket \rrbracket \\
& =\square \llbracket \rrbracket \rrbracket\left(\operatorname{dist}_{\square}^{\times}\left(\delta_{\mathcal{A}}\left(\gamma_{x}\right)_{x:: A \in \Gamma}\right)\right) & & \text { definition of box } \\
& =\llbracket e \rrbracket\left(\gamma_{x}\right)_{x:: A \in \Gamma} & & \text { no-ops } \\
& =\llbracket e \rrbracket\left(\operatorname{strip}_{\Gamma}(\gamma)\right) & & \text { definition of strip }
\end{aligned}
$$

Lemma 34 (Correctness of semifix). If $g^{\prime} \triangleright_{\substack{\mathrm{Lx} \rightarrow \mathrm{F}, \mathrm{L}}} \mathrm{g} \downarrow \mathrm{f} \rightarrow \mathrm{g} \downarrow \mathrm{f}$, then semifix $\left(g, g^{\prime}\right)=$ fix $f$. Proof. First, let's expand our assumption:

If we apply lemma 27 and simplify slightly, this is equivalent to:

$$
\begin{equation*}
(\forall x, d x: \underset{f \in x}{L}) g x=f x \text { and } g(x \vee d x)=f(x \vee d x)=g x \vee g^{\prime} x d x \tag{A.1}
\end{equation*}
$$

This in particular implies that $f=g$.
Now, recall that fix $f$ is defined as the limit $\bigvee_{i} f^{i} \perp$ of the iterations of $f$, while semifix $\left(g, g^{\prime}\right)$ is the limit $\bigvee_{i} x_{i}$ of the sequence $x_{i}$ given by:

$$
\begin{aligned}
x_{0} & =\perp & x_{i+1} & =x_{i} \vee d x_{i} \\
d x_{0} & =g \perp & d x_{i+1} & =g^{\prime} x_{i} d x_{i}
\end{aligned}
$$

Thus it suffices to show that $x_{i}=f^{i} \perp$ ，which we will show inductively，along with $x_{i} \vee d x_{i}=$ $\mathrm{f} \mathrm{x}_{\mathrm{i}}$ ．To establish the base case， $\mathrm{x}_{0}=\perp=\mathrm{f}^{0} \perp$ by definition and $\mathrm{x}_{0} \vee \mathrm{~d} x_{0}=\perp \vee \mathrm{g} \perp=\mathrm{f} \perp$ because $g=f$ ．Inductively assuming that $x_{i}=f^{i} \perp$ and $x_{i} \vee d x_{i}=f x_{i}$ ，we have that $x_{i+1}=x_{i} \vee d x_{i}=f x_{i}=f\left(f^{i} \perp\right)=f^{i+1} \perp$ as desired，and finally：

$$
\begin{aligned}
x_{i+1} \vee d x_{i+1} & =\left(x_{i} \vee d x_{i}\right) \vee g^{\prime} x_{i} d x_{i} & & \text { expanding definitions } \\
& =f x_{i} \vee g^{\prime} x_{i} d x_{i} & & \text { inductive hypothesis } \\
& =g x_{i} \vee g^{\prime} x_{i} d x_{i} & & \text { because } f=g \\
& =f\left(x_{i} \vee d x_{i}\right) & & \text { by equation A.1 } \\
& =f x_{i+1} & & \text { definition of } x_{i+1}
\end{aligned}
$$

Theorem 24 （Fundamental property）．If $\Gamma \vdash e: A$ and $d \gamma \triangleright_{\Gamma} \gamma$ 多 $\rho \rightarrow \gamma^{\prime}$ 名 $\rho^{\prime}$ then

$$
\llbracket \delta e \rrbracket(\gamma, \mathrm{~d} \gamma) \triangleright_{\mathrm{A}} \llbracket \phi e \rrbracket \gamma \text { 夕 } \llbracket e \rrbracket \rho \rightarrow \llbracket \phi e \rrbracket \gamma^{\prime} \text { 夕 } \llbracket e \rrbracket \rho^{\prime}
$$

Proof．By induction on the derivation of $\Gamma \vdash e: A$ ．We will refer to the other premise $\mathrm{d} \gamma \triangleright_{\Gamma} \gamma z \rho \rightarrow \gamma^{\prime} \& \rho^{\prime}$ as simply＂the assumption＂．

Case $\frac{X: A \in \Gamma}{\Gamma \vdash X: A}$ ．We wish to show：

$$
\begin{aligned}
& \llbracket \delta X \rrbracket(\gamma, \mathrm{~d} \gamma) \triangleright_{\mathrm{A}} \llbracket \phi X \rrbracket \gamma \text { 々 } \llbracket \mathrm{X} \rrbracket \rho \rightarrow \llbracket \phi X \rrbracket \gamma^{\prime} \text { 々 } \llbracket \mathrm{X} \rrbracket \rho^{\prime} \\
& \Longleftrightarrow \llbracket \mathrm{DX} \rrbracket(\gamma, \mathrm{~d} \gamma) \triangleright_{\mathrm{A}} \llbracket \mathrm{X} \rrbracket \gamma \text { 多 } \llbracket \mathrm{X} \rrbracket \rho \rightarrow \llbracket X \rrbracket \gamma^{\prime} \text { z } \llbracket \mathrm{X} \rrbracket \rho^{\prime} \\
& \Longleftrightarrow d \gamma_{D X} \triangleright_{A} \gamma_{X} \not \rho_{X} \rightarrow \gamma_{X}^{\prime} \& \rho_{X}^{\prime}
\end{aligned}
$$

which follows from the definition of our assumption．

Case $\frac{x:: A \in \Gamma}{\Gamma \vdash x: A}$ ．We wish to show：

$$
\begin{aligned}
& \llbracket \delta x \rrbracket(\gamma, d \gamma) \triangleright_{A} \llbracket \phi x \rrbracket \gamma z \llbracket x \rrbracket \rho \rightarrow \llbracket \phi x \rrbracket \gamma^{\prime} z \llbracket x \rrbracket \rho^{\prime} \\
& \Longleftrightarrow \llbracket d x \rrbracket(\gamma, d \gamma) \triangleright_{A} \llbracket x \rrbracket \gamma z \llbracket x \rrbracket \rho \rightarrow \llbracket x \rrbracket \gamma^{\prime} z \llbracket x \rrbracket \rho^{\prime} \\
& \Longleftrightarrow \gamma_{\mathrm{dx}} \triangleright_{\mathrm{A}} \gamma_{\mathrm{x}} \not \rho_{x} \rightarrow \gamma_{x}^{\prime} \not \rho_{x}^{\prime}
\end{aligned}
$$

If we apply our assumption we get：

$$
\begin{aligned}
& d \gamma \triangleright_{\Gamma} \gamma \& \rho \rightarrow \gamma^{\prime} \& \rho^{\prime} \\
\Longrightarrow & () \triangleright_{\square \mathrm{A}}\left(\gamma_{x}, \gamma_{\mathrm{dx}}\right) \& \rho_{x} \rightarrow\left(\gamma_{x}^{\prime}, \gamma_{\mathrm{d} x}^{\prime}\right) \& \rho_{x}^{\prime} \\
\Longrightarrow & \gamma_{\mathrm{dx}} \triangleright_{\mathrm{A}} \gamma_{x} \& \rho_{x} \rightarrow \gamma_{x}^{\prime} \& \rho_{x}^{\prime}
\end{aligned}
$$

as desired．

Case $\frac{\Gamma, X: A \vdash e: B}{\Gamma \vdash \lambda X . e: A \rightarrow B}$ Recall that $\phi(\lambda X . e)=\lambda X . \phi e$ and $\delta(\lambda X . e)=\lambda[x] . \lambda D X . \delta e$. We wish to show

$$
\llbracket \lambda[x] . \lambda D X . \delta e \rrbracket(\gamma, d \gamma) \triangleright_{A \rightarrow B} \llbracket \lambda X . \phi e \rrbracket \gamma z \llbracket \lambda X . e \rrbracket \rho \rightarrow \llbracket \lambda X . \phi e \rrbracket \gamma^{\prime} z \llbracket \lambda X . e \rrbracket \rho^{\prime}
$$

Applying the definition of the logical relation at $A \rightarrow B$, it suffices to assume (a) $\mathrm{d} x \triangleright_{\mathrm{A}} x \not z \mathrm{a} \rightarrow \mathrm{y} z \mathrm{~b}$ and prove
$\llbracket \lambda[x] . \lambda D X . \delta e \rrbracket(\gamma, d \gamma) x d x \triangleright_{B} \llbracket \lambda X . \phi e \rrbracket \gamma x \notin \llbracket \lambda X . e \rrbracket \rho a \rightarrow \llbracket \lambda X . \phi e \rrbracket \gamma^{\prime} y \sharp \llbracket \lambda X . e \rrbracket \rho^{\prime} b$ which, by calculation, is:

$$
\llbracket \delta e \rrbracket \sigma \triangleright_{B} \llbracket \phi e \rrbracket(\gamma, X \mapsto x) z \llbracket e \rrbracket(\rho, X \mapsto a) \rightarrow \llbracket \phi e \rrbracket(\gamma, X \mapsto y) z \llbracket e \rrbracket(\rho, X \mapsto b)
$$

where $\sigma=(\gamma, d \gamma, x \mapsto x, D X \mapsto d x)$. This follows from our inductive hypothesis if we can show that:

$$
(\mathrm{d} \gamma, \mathrm{DX} \mapsto \mathrm{dx}) \triangleright_{\Gamma, X: A}(\gamma, X \mapsto x) z(\rho, X \mapsto a) \rightarrow\left(\gamma^{\prime}, X \mapsto y\right) z\left(\rho^{\prime}, X \mapsto b\right)
$$

and this follows from our assumption and (a).

Case $\frac{\Gamma \vdash e: A \rightarrow B \quad \Gamma \vdash f: A}{\Gamma \vdash e f: B}$. Recall that $\phi(e f)=\phi e \phi f$ and $\delta(e f)=\delta e[\phi f] \delta f$. Thus we wish to show:
$\llbracket \delta e[\phi f] \delta f \rrbracket(\gamma, \mathrm{~d} \gamma) \triangleright_{\mathrm{B}} \llbracket \phi e \phi f \rrbracket \gamma z \llbracket e \mathrm{f} \rrbracket \rho \rightarrow \llbracket \phi e \phi f \rrbracket \gamma^{\prime} z \llbracket e f \rrbracket \rho^{\prime}$
Let:

$$
\begin{aligned}
\mathrm{d} x & =\llbracket \delta e \rrbracket(\gamma, \mathrm{~d} \gamma) & & \mathrm{d} y & =\llbracket \delta f \rrbracket(\gamma, \mathrm{~d} \gamma) & \\
x & =\llbracket \phi e \rrbracket \gamma & \mathrm{a} & =\llbracket e \rrbracket \rho & \mathrm{y} & =\llbracket \phi f \rrbracket \gamma \\
x^{\prime} & =\llbracket \phi e \rrbracket \gamma^{\prime} & \mathrm{a}^{\prime} & =\llbracket e \rrbracket \rho^{\prime} & y^{\prime} & =\llbracket \phi f \rrbracket \gamma^{\prime}
\end{aligned}
$$

By our IH for f we have $\mathrm{dy} \triangleright_{\mathrm{A}} \mathrm{y}\left\{\mathrm{b} \rightarrow \mathrm{y}^{\prime} \dot{\mathrm{b}} \mathrm{b}^{\prime}\right.$. By this and our IH for e we have $d x y d y \triangleright_{B} x y$ 多 $b \rightarrow x^{\prime} y^{\prime} \& a^{\prime} b^{\prime}$. By calculation, this is equal to what we wish to show.

Case $\overline{\Gamma \vdash(): 1}$. We wish to show:

$$
\begin{aligned}
& \llbracket() \rrbracket(\gamma, \mathrm{d} \gamma) \triangleright_{1} \llbracket() \rrbracket \gamma z \llbracket() \rrbracket \rho \rightarrow \llbracket() \rrbracket \gamma^{\prime} \& \llbracket() \rrbracket \rho^{\prime} \\
\Longleftrightarrow & () \triangleright_{1}() z() \rightarrow() z() \\
\Longleftrightarrow & T
\end{aligned}
$$

Case $\frac{\left(\Gamma \vdash e_{i}: A_{i}\right)_{i}}{\Gamma \vdash\left(e_{1}, e_{2}\right): A_{1} \times A_{2}}$ : Recall that $\phi\left(\left(e_{1}, e_{2}\right)\right)=\left(\phi e_{1}, \phi e_{2}\right)$ and $\delta\left(\left(e_{1}, e_{2}\right)\right)=\left(\delta e_{1}, \delta e_{2}\right)$. Thus we wish to show:
$\llbracket\left(\delta e_{1}, \delta e_{2}\right) \rrbracket(\gamma, \mathrm{d} \gamma) \triangleright_{A_{1} \times A_{2}} \llbracket\left(\phi e_{1}, \phi e_{2}\right) \rrbracket \gamma \psi \llbracket\left(e_{1}, e_{2}\right) \rrbracket \rho \rightarrow \llbracket\left(\phi e_{1}, \phi e_{2}\right) \rrbracket \gamma^{\prime} \sharp \llbracket\left(e_{1}, e_{2}\right) \rrbracket \rho^{\prime}$
Since in general $\llbracket\left(f_{1}, f_{2}\right) \rrbracket \sigma=\left(\llbracket f_{1} \rrbracket \sigma, \llbracket f_{2} \rrbracket \sigma\right)$, applying the definition of the LR at $A_{1} \times A_{2}$ this is equivalent to:

$$
(\forall i) \llbracket \delta e_{i} \rrbracket(\gamma, \mathrm{~d} \gamma) \triangleright_{A_{i}} \llbracket \phi e_{i} \rrbracket \gamma z \llbracket e_{i} \rrbracket \rho \rightarrow \llbracket \phi e_{i} \rrbracket \gamma^{\prime} z \llbracket e_{i} \rrbracket \rho^{\prime}
$$

which holds by our IH.

Case $\frac{\Gamma \vdash e: A_{1} \times A_{2}}{\Gamma \vdash \pi_{i} e: A_{i}}$. Recall that $\phi\left(\pi_{i} e\right)=\pi_{i} \phi e$ and $\delta\left(\pi_{i} e\right)=\pi_{i} \delta e$ and observe that $\llbracket \pi_{i} f \rrbracket \sigma=\pi_{\mathfrak{i}}(\llbracket f \rrbracket \sigma)$. Applying this, what we wish to show is

$$
\pi_{\mathrm{i}}(\llbracket \delta e \rrbracket(\gamma, \mathrm{~d} \gamma)) \triangleright_{\mathrm{A}_{\mathrm{i}}} \pi_{\mathrm{i}}(\llbracket \phi e \rrbracket \gamma) \& \pi_{\mathrm{i}}(\llbracket e \rrbracket \rho) \rightarrow \pi_{\mathrm{i}}\left(\llbracket \phi e \rrbracket \gamma^{\prime}\right) \& \pi_{\mathrm{i}}\left(\llbracket e \rrbracket \rho^{\prime}\right)
$$

which is a direct consequence of our IH.

Case $\frac{\Gamma \vdash e: A_{i}}{\Gamma \vdash i n_{i} e: A_{1}+A_{2}}$. Recall that $\phi\left(\mathrm{in}_{i} e\right)=\mathrm{in}_{i} \phi e$ and $\delta\left(\mathrm{in}_{i} e\right)=\mathrm{in} \delta e$. Observe that in general $\llbracket \mathrm{in}_{\mathrm{i}} \mathrm{f} \rrbracket \sigma=\mathrm{in}_{\mathrm{i}}(\llbracket \mathrm{f} \rrbracket \sigma)$ and therefore what we wish to show is equivalent to:

$$
\mathrm{in}_{\mathrm{i}}(\llbracket \delta e \rrbracket(\gamma, \mathrm{~d} \gamma)) \triangleright_{\mathrm{A}_{1}+\mathrm{A}_{2}} \mathrm{in}_{\mathrm{i}}(\llbracket \phi e \rrbracket \gamma)\left\langle\mathrm { in } _ { \mathrm { i } } ( \llbracket e \rrbracket \rho ) \rightarrow \mathrm { in } _ { \mathrm { i } } ( \llbracket \phi e \rrbracket \gamma ^ { \prime } ) \left\langle\mathrm{in}_{\mathrm{i}}\left(\llbracket e \rrbracket \rho^{\prime}\right)\right.\right.
$$

which is by definition equivalent to our inductive hypothesis.

Case $\underset{\Gamma \vdash \perp: \mathrm{L}}{ }$. Recall that $\phi \perp=\perp$ and $\delta \perp=\perp$ and $\llbracket \perp \rrbracket \sigma=\perp$. Thus it STS $\perp \triangleright_{\mathrm{L}}$ $\perp$ \& $\rightarrow \perp$ \& $\perp$, which holds by lemma 27.

Case $\frac{\left(\Gamma \vdash e_{i}: L\right)_{i}}{\Gamma \vdash e_{1} \vee e_{2}: L}$. Recall that $\phi\left(e_{1} \vee e_{2}\right)=\phi e_{1} \vee \phi e_{2}$. We wish to show: $\llbracket \delta e_{1} \vee \delta e_{2} \rrbracket(\gamma, d \gamma) \triangleright_{\mathrm{L}} \llbracket \phi e_{1} \vee \phi e_{2} \rrbracket \gamma \sharp \llbracket e_{1} \vee e_{2} \rrbracket \rho \rightarrow \llbracket \phi e_{1} \vee \phi e_{2} \rrbracket \gamma^{\prime} \sharp \llbracket e_{1} \vee e_{2} \rrbracket \rho^{\prime}$ Let:

$$
\mathrm{d} x_{i}=\llbracket \delta e_{i} \rrbracket(\gamma, \mathrm{~d} \gamma) \quad x_{i}=\llbracket \phi e_{i} \rrbracket \gamma \quad x_{i}^{\prime}=\llbracket \phi e_{i} \rrbracket \gamma^{\prime} \quad a_{i}=\llbracket e_{i} \rrbracket \rho \quad a_{i}^{\prime}=\llbracket e_{i} \rrbracket \rho^{\prime}
$$

Since $\llbracket f \vee g \rrbracket \sigma=\llbracket f \rrbracket \sigma \vee \llbracket g \rrbracket \sigma$, what we wish to show is equivalent to:

$$
d x_{1} \vee d x_{2} \triangleright_{L} x_{1} \vee x_{2} \vDash a_{1} \vee a_{2} \rightarrow x_{1}^{\prime} \vee x_{2}^{\prime} \& a_{1}^{\prime} \vee a_{2}^{\prime}
$$

Applying lemma 27 this is equivalent to:

$$
x_{1} \vee x_{2}=a_{1} \vee a_{2} \quad x_{1}^{\prime} \vee x_{2}^{\prime}=a_{1}^{\prime} \vee a_{2}^{\prime}=x_{1} \vee x_{2} \vee d x_{1} \vee d x_{2}
$$

By our IH and assumption and lemma 27 we have:

$$
x_{i}=a_{i} \quad x_{i}^{\prime}=a_{i}^{\prime}=x_{i} \vee d x_{i}
$$

which suffices by associativity and commutativity of $\vee$.
 $\perp$, by lemma 27 what we want to show is equivalent to:

$$
\llbracket\left\{\phi e_{i}\right\}_{i} \rrbracket \gamma=\llbracket\left\{e_{i}\right\}_{i} \rrbracket \rho=\llbracket\left\{\phi e_{i}\right\}_{i} \rrbracket \gamma^{\prime}=\llbracket\left\{e_{i}\right\}_{i} \rrbracket \rho^{\prime}
$$

Note that $\llbracket\left\{\mathrm{f}_{i}\right\}_{i} \rrbracket \sigma=\bigvee_{i}\left\{\operatorname{box}_{\Gamma}\left(\llbracket \mathrm{f}_{i} \rrbracket\right)(\sigma)\right\}=\left\{\llbracket \mathrm{f}_{i} \rrbracket\right.$ (strip $\left.\left.\sigma\right)\right\}_{i}$. Also observe that by lemmas 28 and 29 and our assumption, we have

$$
\begin{align*}
& () \triangleright_{\lceil\Gamma]} \text { strip } \gamma \& \text { strip } \rho \rightarrow \text { strip } \gamma^{\prime} \& \text { strip } \rho^{\prime}  \tag{A.2}\\
& \text { strip } \gamma=\text { strip } \gamma^{\prime} \quad \text { and } \quad \text { strip } \rho=\text { strip } \rho^{\prime} \tag{A.3}
\end{align*}
$$

So applying equation A. 3 it suffices to show that $\left\{\llbracket \phi e_{i} \rrbracket\right.$ (strip $\left.\left.\gamma\right)\right\}_{i}=\left\{\llbracket e_{i} \rrbracket\right.$ (strip $\left.\left.\rho\right)\right\}_{i}$, for which it suffices to show $\llbracket \phi e_{i} \rrbracket$ (strip $\left.\gamma\right)=\llbracket e_{i} \rrbracket$ (strip $\rho$ ). This holds by our IH for $e_{i}$ and equation A. 2 and lemma 25.

Case $\frac{\left.(\Gamma \Gamma\rceil \vdash e_{i}: \underset{\varepsilon}{2}\right)_{i}}{\Gamma \vdash e_{1}=e_{2}: \text { bool }}$. Recall that $\phi\left(e_{1}=e_{2}\right)=\phi e_{1}=\phi e_{2}$ and $\delta\left(e_{1}=e_{2}\right)=\perp$. Thus what we wish to show is:

$$
\llbracket \perp \rrbracket(\gamma, \mathrm{d} \gamma) \triangleright_{\text {bool }} \llbracket \phi e_{1}=\phi e_{2} \rrbracket \gamma \text { 名 } \llbracket e_{1}=e_{2} \rrbracket \rho \rightarrow \llbracket \phi e_{1}=\phi e_{2} \rrbracket \gamma^{\prime} z \llbracket e_{1}=e_{2} \rrbracket \rho^{\prime}
$$

Observing that $\llbracket \perp \rrbracket(\gamma, \mathrm{d} \gamma)=\emptyset$ and applying the definition of the logic relation for bool $=\{1\}$, this is equivalent to:

$$
\llbracket \phi e_{1}=\phi e_{2} \rrbracket \gamma=\llbracket \phi e_{1}=\phi e_{2} \rrbracket \gamma^{\prime}=\llbracket e_{1}=e_{2} \rrbracket \rho=\llbracket e_{1}=e_{2} \rrbracket \rho^{\prime}
$$

Observe by calculation that

$$
\llbracket \mathrm{f}_{1}=\mathrm{f}_{2} \rrbracket \sigma= \begin{cases}\{()\} & \text { if } \llbracket \mathrm{f}_{1} \rrbracket(\text { strip } \sigma)=\llbracket \mathrm{f}_{2} \rrbracket(\text { strip } \sigma) \\ \emptyset & \text { otherwise }\end{cases}
$$

Thus it suffices to show $\llbracket \phi e_{i} \rrbracket($ strip $\gamma)=\llbracket \phi e_{i} \rrbracket\left(\right.$ strip $\left.\gamma^{\prime}\right)=\llbracket e_{i} \rrbracket($ strip $\rho)=$ $\llbracket e_{i} \rrbracket$ (strip $\left.\rho^{\prime}\right)$. By our assumption and lemmas 28 and 29 we know strip $\gamma=$ strip $\gamma^{\prime}$ and strip $\rho=$ strip $\rho^{\prime}$ and ()$\triangleright_{\lceil\Gamma\rceil}$ strip $\gamma$ \& strip $\rho \rightarrow$ strip $\gamma^{\prime}$ \& strip $\rho^{\prime}$. By the first two equalities it now suffices to show $\llbracket \phi e_{i} \rrbracket$ (strip $\gamma$ ) $=\llbracket e_{i} \rrbracket$ (strip $\rho$ ). Applying our IH to the remaining third proposition we have

$$
\llbracket \delta e_{i} \rrbracket(\text { strip } \gamma) \triangleright_{\underset{\varepsilon}{A}} \llbracket \phi e_{i} \rrbracket(\text { strip } \gamma) \& \llbracket e_{i} \rrbracket(\text { strip } \rho) \rightarrow \llbracket \phi e_{i} \rrbracket\left(\text { strip } \gamma^{\prime}\right) z \llbracket e_{i} \rrbracket\left(\text { strip } \rho^{\prime}\right)
$$

which by lemma 25 implies $\llbracket \phi e_{i} \rrbracket($ strip $\gamma)=\llbracket e_{i} \rrbracket($ strip $\rho)$ as desired.

Case $\frac{\lceil\rceil \vdash e:\{1\}}{\Gamma \vdash e m p t y ? ~ e: 1+1}$. Recall that $\phi($ empty? $e)=\delta(e m p t y ? ~ e)=$ empty? $\phi e$. Note that by lemmas 28 and 29 we have strip $\gamma=$ strip $\gamma^{\prime}$ and strip $\rho=$ strip $\rho^{\prime}$ and () $\triangleright_{\lceil\Gamma 7}$ strip $\gamma$ \& strip $\rho \rightarrow$ strip $\gamma^{\prime}$ \& strip $\rho^{\prime}$; applying this to our inductive hypothesis and invoking lemma 25 on the result, we have

$$
\llbracket e \rrbracket(\text { strip } \rho)=\llbracket e \rrbracket\left(\text { strip } \rho^{\prime}\right)=\llbracket \phi e \rrbracket(\text { strip } \gamma)=\llbracket \phi e \rrbracket\left(\text { strip } \gamma^{\prime}\right)
$$

Thus all are equal to the same value. Now, observe by calculation that

$$
\llbracket e m p t y ? \mathrm{f} \rrbracket \sigma= \begin{cases}\mathrm{in} \mathrm{in}_{\mathrm{i}}( & \text { if } \llbracket \mathrm{f} \rrbracket(\text { strip } \sigma)=\emptyset \\ \mathrm{in}_{2}() & \text { otherwise }\end{cases}
$$

Thus, there is some $i$ such that $\llbracket e m p t y ? ~ e \rrbracket \rho=\llbracket e m p t y ? ~ e \rrbracket \rho^{\prime}=\llbracket e m p t y ? \phi e \rrbracket \gamma=$【empty? $\phi e \rrbracket \gamma^{\prime}=\mathrm{in}_{\mathrm{i}}()$ and we have $\mathrm{in}_{\mathrm{i}}() \triangleright_{1+1} \mathrm{in}_{\mathrm{i}}() \& \mathrm{in}_{\mathfrak{i}}() \rightarrow \mathrm{in}_{\mathrm{i}}() \& \mathrm{in} \mathrm{n}_{\mathrm{i}}()$ as desired.

Case $\frac{\Gamma \vdash e: A_{1}+A_{2} \quad\left(\Gamma, X_{i}: A_{i} \vdash f_{i}: B\right)_{i}}{\Gamma \vdash \text { case e of }\left(i n_{i} X \rightarrow f_{i}\right)_{i}: B}$. Recall that

$$
\begin{aligned}
\phi\left(\text { case } e \text { of }\left(i n_{i} X \rightarrow f_{i}\right)_{i}\right)= & \text { case } \phi e \text { of }\left(i n_{i} X \rightarrow \phi f_{i}\right)_{i} \\
\delta\left(\text { case } e \text { of }\left(i n_{i} X \rightarrow f_{i}\right)_{i}\right)= & \text { case split }[\phi e] \text { of } \\
& \left(i n_{i} Y \rightarrow\right. \\
& \text { let }[x]=Y \text { in } \\
& \left(\lambda D X . \delta f_{i}\right) \\
& \left(\text { case } \delta e \text { of } i n_{i} D X \rightarrow D X\right. \\
& \left.\left.i n_{i+1 \bmod 2} \rightarrow \text { dummy } x\right)\right)_{i}
\end{aligned}
$$

Let $\rho_{1}=\rho, \rho_{2}=\rho^{\prime}, \gamma_{1}=\gamma, \gamma_{2}=\gamma^{\prime}$. By our inductive hypothesis for $e$ there must be some $k \in\{1,2\}$ and some $d x, x_{i}, a_{i}$ such that $d x \triangleright_{A_{k}} x_{1} \& a_{1} \rightarrow x_{2} \& a_{2}$ and:

$$
\llbracket e \rrbracket \rho_{i}=\mathrm{in}_{k} \mathrm{a}_{\mathrm{i}} \quad \llbracket \phi e \rrbracket \gamma_{i}=\mathrm{in}_{k} x_{\mathrm{i}} \quad \llbracket \delta e \rrbracket(\gamma, \mathrm{~d} \gamma)=\mathrm{in}_{\mathrm{k}} \mathrm{~d} x
$$

Applying this and our inductive hypothesis for $f_{k}$ it will suffice to show that:

$$
\begin{align*}
\llbracket \text { case } e \text { of }\left(\mathrm{in}_{i} X \rightarrow \mathrm{f}_{\mathrm{i}}\right)_{i} \rrbracket \rho_{i} & =\llbracket \mathrm{f}_{\mathrm{k}} \rrbracket\left(\rho_{i}, \mathrm{X} \mapsto \mathrm{a}_{\mathrm{i}}\right)  \tag{A.4}\\
\llbracket \phi\left(\text { case } e \text { of }\left(\mathrm{in}_{i} \mathrm{X} \rightarrow \mathrm{f}_{\mathrm{i}}\right)_{\mathrm{i}}\right) \rrbracket \gamma_{i} & =\llbracket \phi \mathrm{f}_{\mathrm{k}} \rrbracket\left(\gamma_{i}, \mathrm{X} \mapsto \mathrm{x}_{\mathrm{i}}\right)  \tag{A.5}\\
\llbracket \delta\left(\text { case } e \text { of }\left(\mathrm{in} \mathrm{n}_{i} X \rightarrow \mathrm{f}_{\mathrm{i}}\right)_{\mathrm{i}}\right) \rrbracket(\gamma, \mathrm{d} \gamma) & =\llbracket \delta \mathrm{f}_{\mathrm{k}} \rrbracket\left(\gamma, \mathrm{~d} \gamma, \mathrm{x} \mapsto \mathrm{x}_{1}, \mathrm{DX} \mapsto \mathrm{dx}\right) \tag{A.6}
\end{align*}
$$

Showing this is a matter of calculation. Equations A. 4 and A. 5 are fairly straightforward to calculate, so we only show equation A. 6 in detail. Before starting, it will be useful to give some abbreviations for subterms of $\delta\left(\right.$ case $e$ of $\left.\left(i n_{i} X \rightarrow f_{i}\right)_{i}\right)$ :

$$
\begin{aligned}
h_{i} & =\text { case } \delta e \text { of } \text { in }_{i} D X \rightarrow D X ; \text { in }_{i+1 \bmod 2-} \rightarrow \text { dummy } x \\
g_{i} & =\text { let }[x]=Y \text { in }\left(\lambda D X . \delta f_{i}\right) h_{i}
\end{aligned}
$$

It will also be useful to note the type of $\phi e$ as it occurs in the sub-expression split [ $\phi e$ ] being immediately analyzed by $\delta\left(\right.$ case $e$ of $\left.\left(\mathrm{in}_{i} X \rightarrow f_{i}\right)_{i}\right)$; it has been weakened to the type $\square Ф Г \vdash$ фe: $\Phi A_{1}+\Phi A_{2}$.

$$
\begin{aligned}
& \llbracket \delta\left(\text { case } e \text { of }\left(\mathrm{in}_{i} X \rightarrow \mathrm{f}_{\mathrm{i}}\right)_{\mathrm{i}}\right) \rrbracket(\gamma, \mathrm{d} \gamma) \\
& =\left[\llbracket g_{i} \rrbracket\right]_{i}\left(\operatorname{dist}_{+}^{\times}((\gamma, \mathrm{d} \gamma), \llbracket \text { split }[\phi e] \rrbracket(\gamma, \mathrm{d} \gamma))\right) \\
& =\left[\llbracket g_{i} \rrbracket\right]_{i}\left(\operatorname{dist}_{+}^{\times}\left((\gamma, \mathrm{d} \gamma), \operatorname{dist}_{+}^{\square}(\llbracket[\phi e] \rrbracket(\gamma, \mathrm{d} \gamma))\right)\right) \\
& =\left[\llbracket g_{i} \rrbracket\right]_{i}\left(\operatorname{dist}_{+}^{\times}((\gamma, \mathrm{d} \gamma), \llbracket[\phi e] \rrbracket(\gamma, \mathrm{d} \gamma))\right) \quad \operatorname{dist}_{+}^{\square} \text { is the identity } \\
& =\left[\llbracket g_{i} \rrbracket\right]_{i}\left(\operatorname{dist}_{+}^{\times}\left((\gamma, \mathrm{d} \gamma), \llbracket \phi e \rrbracket\left(\operatorname{strip}_{\square \Phi \Gamma}(\gamma, \mathrm{d} \gamma)\right)\right)\right) \quad \text { lemma } 33 \\
& =\left[\llbracket g_{i} \rrbracket\right]_{i}\left(\operatorname{dist}_{+}^{\times}((\gamma, \mathrm{d} \gamma), \llbracket e \rrbracket \gamma)\right) \\
& =\left[\left[g_{i} \rrbracket_{i}\left(\operatorname{dist}_{+}^{\times}\left((\gamma, \mathrm{d} \gamma), \mathrm{in}_{\mathrm{k}} \mathrm{x}_{1}\right)\right)\right.\right. \\
& =\left[\llbracket g_{i} \rrbracket\right]_{i}\left(\mathrm{in}_{k}\left(\gamma, \mathrm{~d} \gamma, \mathrm{Y} \mapsto \mathrm{x}_{1}\right)\right) \\
& =\llbracket g_{k} \rrbracket\left(\gamma, \mathrm{~d} \gamma, \mathrm{Y} \mapsto \mathrm{x}_{1}\right) \\
& =\llbracket \text { let }[x]=Y \text { in }\left(\lambda D X . \delta f_{k}\right) h_{k} \rrbracket\left(\gamma, d \gamma, Y \mapsto x_{1}\right) \\
& =\llbracket\left(\lambda D X . \delta f_{k}\right) h_{k} \rrbracket\left(\gamma, d \gamma, x \mapsto x_{1}\right) \\
& =\llbracket\left(\lambda D X . \delta f_{k}\right) \rrbracket\left(\gamma, d \gamma, x \mapsto x_{1}\right)\left(\llbracket h_{k} \rrbracket\left(\gamma, d \gamma, x \mapsto x_{1}\right)\right) \\
& =\left(\mathrm{dx} \mapsto \llbracket \delta \mathrm{f}_{\mathrm{k}} \rrbracket\left(\gamma, \mathrm{~d} \gamma, \mathrm{x} \mapsto \mathrm{x}_{1}, \mathrm{DX} \mapsto \mathrm{dx}\right)\right) \\
& \left(\llbracket h_{k} \rrbracket\left(\gamma, d \gamma, x \mapsto x_{1}\right)\right) \\
& =\llbracket \delta f_{k} \rrbracket\left(\gamma, \mathrm{~d} \gamma, \mathrm{x} \mapsto \mathrm{x}_{1}, \mathrm{DX} \mapsto\left(\llbracket h_{\mathrm{k}} \rrbracket\left(\gamma, \mathrm{~d} \gamma, \mathrm{x} \mapsto \mathrm{x}_{1}\right)\right)\right)
\end{aligned}
$$

And therefore it suffices to show that $\llbracket h_{k} \rrbracket\left(\gamma, d \gamma, x \mapsto x_{1}\right)=d x$. Without loss of generality, assume $k=1$ :

$$
\begin{aligned}
& \llbracket h_{1} \rrbracket\left(\gamma, \mathrm{~d} \gamma, \mathrm{x} \mapsto \mathrm{x}_{1}\right) \\
= & \llbracket \text { case } \delta e \text { of in }{ }_{1} \mathrm{DX} \rightarrow \mathrm{DX} ; \mathrm{in}_{2}-\rightarrow \text { dummy } x \rrbracket\left(\gamma, \mathrm{~d} \gamma, x \mapsto \mathrm{x}_{1}\right) \\
= & {[\llbracket \mathrm{DX} \rrbracket, \llbracket \text { dummy } \mathrm{x} \rrbracket]\left(\text { dist }_{+}^{x}\left(\left(\gamma, \mathrm{~d} \gamma, \mathrm{x} \mapsto \mathrm{x}_{1}\right), \llbracket \delta \mathrm{d} \rrbracket\left(\gamma, \mathrm{~d} \gamma, x \mapsto \mathrm{x}_{1}\right)\right)\right) } \\
= & {[\llbracket \mathrm{DX} \rrbracket, \llbracket \text { dummy } x \rrbracket]\left(\left(\gamma, \mathrm{d} \gamma, \mathrm{x} \mapsto \mathrm{x}_{1}\right),\left(\mathrm{in}_{1} \mathrm{dx}\right)\right) } \\
= & {[\llbracket \mathrm{DX} \rrbracket \llbracket \text { dummy } x \rrbracket]\left(\mathrm{in}_{1}\left(\gamma, \mathrm{~d} \gamma, x \mapsto \mathrm{x}_{1}, \mathrm{DX} \mapsto \mathrm{~d} x\right)\right) } \\
= & \llbracket \mathrm{DX} \rrbracket\left(\gamma, \mathrm{~d} \gamma, x \mapsto \mathrm{x}_{1}, \mathrm{DX} \mapsto \mathrm{~d} x\right) \\
= & \mathrm{d} x
\end{aligned}
$$

Which is what we wished to show.

Case $\frac{\lceil\Gamma\rceil \vdash e: A}{\Gamma \vdash[e]: \square \mathcal{A}}, \phi[e]=[(\phi e, \delta e)], \delta[e]=()$.
For brevity, let

$$
\gamma_{s}=\operatorname{strip}_{\Phi \Gamma}(\gamma) \quad \gamma_{s}^{\prime}=\operatorname{strip}_{\Gamma}\left(\gamma^{\prime}\right) \quad \rho_{s}=\operatorname{strip}_{\Phi \Gamma}(\rho) \quad \rho_{s}^{\prime}=\operatorname{strip}_{\Gamma}\left(\rho^{\prime}\right)
$$

By applying lemma 29 to our assumption, we have

$$
\begin{equation*}
\text { () } \triangleright_{\Gamma} \gamma_{s} \& \rho_{s} \rightarrow \gamma_{s}^{\prime} \& \rho_{s}^{\prime} \tag{A.7}
\end{equation*}
$$

By applying lemma 28 we further know

$$
\begin{equation*}
\gamma_{s}=\gamma_{\mathrm{s}}^{\prime} \quad \text { and } \quad \rho_{\mathrm{s}}=\rho_{\mathrm{s}}^{\prime} \tag{A.8}
\end{equation*}
$$

We wish to show:

$$
\llbracket() \rrbracket(\gamma, \mathrm{d} \gamma) \triangleright \square \mathrm{A} \llbracket[(\phi e, \delta e)] \rrbracket \gamma z \llbracket[e] \rrbracket \rho \rightarrow \llbracket[(\phi e, \delta e)] \rrbracket \gamma^{\prime} z \llbracket[e] \rrbracket \rho^{\prime}
$$

Applying lemma 33 and further simplifying, this is equivalent to:

$$
() \triangleright_{\square A}\left(\llbracket \phi e \rrbracket \gamma_{s}, \llbracket \delta e \rrbracket \gamma_{s}\right) z \llbracket e \rrbracket \rho_{s} \rightarrow\left(\llbracket \phi e \rrbracket \gamma_{s}, \llbracket \delta e \rrbracket \gamma_{s}\right) z \llbracket e \rrbracket \rho_{s}
$$

Applying the definition of the logical relation at $\square A$, this requires that $\llbracket e \rrbracket \rho_{s}=\llbracket e \rrbracket \rho_{s}^{\prime}$ and $\llbracket \phi e \rrbracket \gamma_{s}=\llbracket \phi e \rrbracket \gamma_{s}^{\prime}$ and $\llbracket \delta e \rrbracket \gamma_{s}=\llbracket \delta e \rrbracket \gamma_{s}^{\prime}$, which hold by equation A.8, and that:

$$
\llbracket \delta e \rrbracket \gamma_{\mathrm{s}} \triangleright_{\mathrm{A}} \llbracket \phi e \rrbracket \gamma_{\mathrm{s}} z \llbracket e \rrbracket \rho_{\mathrm{s}} \rightarrow \llbracket \phi e \rrbracket \gamma_{\mathrm{s}}^{\prime} z \llbracket e \rrbracket \rho_{\mathrm{s}}^{\prime}
$$

which follows from our inductive hypothesis applied to equation A.7.

Case $\frac{\Gamma \vdash e: \square A \quad \Gamma, x: A \vdash f: B}{\Gamma \vdash \operatorname{let}[\mathrm{x}]=\mathrm{e} \text { in } \mathrm{f}: \mathrm{B}}$. Observe that

$$
\begin{aligned}
& \phi(\text { let }[x]=e \text { in } f) \\
&=\text { let }[(x, d x)]=\phi e \text { in } \phi f \\
& \delta(\text { let }[x]=e \text { in } f)=\text { let }[(x, d x)]=\phi e \text { in } \delta f
\end{aligned}
$$

Further observe that

$$
\llbracket \text { let }[(x, d x)]=\phi e \text { in } \phi f \rrbracket \gamma=\llbracket \phi f \rrbracket(\gamma, x, d x)
$$

and likewise for $\delta \mathrm{f}$ in place of $\phi \mathrm{f}$ and/or $\gamma^{\prime}$ in place of $\gamma$. For brevity, let

$$
\begin{aligned}
x, \mathrm{~d} x & =\llbracket \phi e \rrbracket \gamma \\
\mathrm{a} & =\llbracket e \rrbracket \rho
\end{aligned}
$$

$$
\begin{aligned}
x^{\prime}, \mathrm{d} x^{\prime} & =\llbracket \phi e \rrbracket \gamma^{\prime} \\
\mathrm{a}^{\prime} & =\llbracket e \rrbracket \rho^{\prime}
\end{aligned}
$$

Using this observation and these abbreviations, we have

$$
\begin{aligned}
\llbracket \text { let }[x]=e \text { in } f \rrbracket \rho & =\llbracket f \rrbracket(\rho, a) \\
\llbracket \text { let }[(x, d x)]=\phi e \text { in } \phi f \rrbracket \gamma & =\llbracket \phi f \rrbracket(\gamma, x, d x) \\
\llbracket \text { let }[(x, d x)]=\phi e \text { in } \delta f \rrbracket \gamma & =\llbracket \delta f \rrbracket(\gamma, x, d x)
\end{aligned}
$$

and likewise for $\gamma^{\prime}, \rho^{\prime}$. Then what we wish to show is that

$$
\llbracket \delta f \rrbracket(\gamma, x, d x) \triangleright_{\mathrm{B}} \llbracket \phi f \rrbracket(\gamma, x, d x) z \llbracket f \rrbracket(\rho, a) \rightarrow \llbracket \phi f \rrbracket\left(\gamma^{\prime}, x^{\prime}, \mathrm{d} x^{\prime}\right) z \llbracket f \rrbracket\left(\rho^{\prime}, a^{\prime}\right)
$$

By our inductive hypothesis for $f$, it suffices to show that

$$
\mathrm{d} \gamma \triangleright_{\Gamma, x:: A}(\gamma, x, \mathrm{dx}) \notin(\rho, a) \rightarrow\left(\gamma^{\prime}, x^{\prime}, \mathrm{d} x^{\prime}\right) \nless\left(\rho^{\prime}, a^{\prime}\right)
$$

Since our assumption tells us that $\mathrm{d} \gamma \triangleright_{\Gamma} \gamma \& \rho \rightarrow \gamma^{\prime} \& \rho^{\prime}$, it suffices to show that () $\triangleright_{\square A}(x, d x) \& a \rightarrow\left(x^{\prime}, d x^{\prime}\right) \& a^{\prime}$, which follows directly from our inductive hypothesis for $e$.

Case $\frac{\Gamma \vdash e:\left\{\begin{array}{c}A \\ A_{\varepsilon}\end{array}\right\} \quad \Gamma, x:: \underset{\varepsilon \in}{A} \vdash \mathrm{f}: \mathrm{L}}{\Gamma \vdash \text { for }(x \in e) \mathrm{f}: \mathrm{L}}$. Recall that:

$$
\begin{aligned}
\phi(\text { for }(x \in e) f) & =\text { for }(x \in \phi e) \text { let }[d x]=[0 x] \text { in } \phi f \\
\delta(\text { for }(x \in e) f) & =(\text { for }(x \in \delta e) \text { let }[d x]=[0 x] \text { in } \phi f) \\
& \vee(\text { for }(x \in \phi e \vee \delta e) \text { let }[d x]=[0 x] \text { in } \delta f)
\end{aligned}
$$

This case of the proof is quite complex; it will help to have a few abbreviations. First, we will be considering various definitons which differ only in whether they use the primed or un-primed versions of $\gamma, \rho$, so it will help to refer to these by subscript: $\gamma_{1}=\gamma$ and $\gamma_{2}=\gamma^{\prime}$ and $\rho_{1}=\rho$ and $\rho_{2}=\rho^{\prime}$.
With this in mind, apply lemma 27 to our inductive hypothesis for e :

$$
\begin{align*}
& \llbracket \delta e \rrbracket\left(\gamma_{1}, \mathrm{~d} \gamma\right) \triangleright_{\left\{A_{\mathrm{EQ}}\right\}} \llbracket \phi e \rrbracket \gamma_{1} z \llbracket e \rrbracket \rho_{1} \rightarrow \llbracket \phi e \rrbracket \gamma_{2} z \llbracket e \rrbracket \rho_{2}  \tag{A.9}\\
\Longleftrightarrow & \llbracket \phi e \rrbracket \gamma_{1}=\llbracket e \rrbracket \rho_{1} \text { and } \llbracket \phi e \rrbracket \gamma_{2}=\llbracket e \rrbracket \rho_{2}=\llbracket \phi e \rrbracket \gamma_{1} \vee \llbracket \delta e \rrbracket\left(\gamma_{1}, \mathrm{~d} \gamma\right) \tag{A.10}
\end{align*}
$$

Moving to our inductive hypothesis for $f$, for any $x \in \llbracket{ }_{\varepsilon \in} \rrbracket \rrbracket$, let:

$$
\rho_{i}^{x}=\left(\rho_{i}, x \mapsto x\right) \quad \gamma_{i}^{x}=\left(\gamma_{i}, x \mapsto x, \mathrm{~d} x \mapsto \text { dummy } x\right) \quad \mathrm{d} \gamma^{x}=\left(\gamma_{1}^{x}, \mathrm{~d} \gamma\right)
$$

 and our inductive hypothesis for f , applying lemma 27 :

$$
\begin{align*}
& \llbracket \delta f \rrbracket \mathrm{~d} \gamma^{\mathrm{x}} \triangleright_{\mathrm{L}} \llbracket \phi \mathrm{f} \rrbracket \gamma_{1}^{\mathrm{x}} \downarrow \llbracket \mathrm{f} \rrbracket \rho_{1}^{\mathrm{x}} \rightarrow \llbracket \phi \mathrm{f} \rrbracket \gamma_{2}^{\chi} \downarrow \llbracket \mathrm{f} \rrbracket \rho_{2}^{x}  \tag{A.11}\\
\Longleftrightarrow & \llbracket \phi \mathrm{f} \rrbracket \gamma_{1}^{\chi}=\llbracket \mathrm{f} \rrbracket \rho_{1}^{\mathrm{x}} \text { and } \llbracket \phi f \rrbracket \gamma_{2}^{x}=\llbracket \mathrm{f} \rrbracket \rho_{2}^{x}=\llbracket \phi \mathrm{f} \rrbracket \gamma_{1}^{\chi} \vee \llbracket \delta \mathrm{f} \rrbracket \mathrm{~d} \gamma^{x} \tag{A.12}
\end{align*}
$$

We summarize equations A. 10 and A. 12 as follows, introducing variables $s_{i}$, ds, $F_{i}$, dF:

$$
\begin{align*}
s_{i} & =\llbracket e \rrbracket \rho_{i}=\llbracket \phi e \rrbracket \gamma_{i} & \mathrm{~F}_{i}(\mathrm{x}) & =\llbracket \mathrm{f} \rrbracket \rho_{i}^{x}=\llbracket \phi f \rrbracket \gamma_{i}^{x}  \tag{A.13}\\
\mathrm{ds} & =\llbracket \delta e \rrbracket\left(\gamma_{1}, \mathrm{~d} \gamma\right) & \mathrm{dF}(\mathrm{x}) & =\llbracket \delta \mathrm{f} \rrbracket \mathrm{~d} \gamma^{\mathrm{x}} \\
\mathrm{~s}_{1} \cup \mathrm{ds} & =\mathrm{s}_{2} & \mathrm{~F}_{1}(\mathrm{x}) \vee \mathrm{dF}(\mathrm{x}) & =\mathrm{F}_{2}(\mathrm{x}) \tag{A.14}
\end{align*}
$$

Now let's give abbreviations to the denotations about which we are trying to prove something:

$$
\begin{aligned}
l_{i} & =\llbracket \text { for }(x \in e) f \rrbracket \rho_{i} \\
\mathrm{~m}_{\mathrm{i}} & =\llbracket \phi(\text { for }(x \in e) f) \rrbracket \gamma_{i} \\
\mathrm{dm} & =\llbracket \delta(\text { for }(x \in e) \mathrm{f}) \rrbracket\left(\gamma_{1}, \mathrm{~d} \gamma\right)
\end{aligned}
$$

Then what we wish to show is that $\mathrm{dm} \triangleright_{\mathrm{L}} \mathfrak{m}_{1} \& l_{1} \rightarrow \mathrm{~m}_{2} \& l_{2}$, or equivalently by applying lemma 27, that $m_{1}=l_{1}$ and $m_{2}=l_{2}=m_{1} \vee d m$. We will do this by showing that

$$
\begin{align*}
l_{i}=m_{i} & =\bigvee_{x \in s_{i}} F_{i}(x)  \tag{A.16}\\
\mathrm{dm} & =\left(\bigvee_{x \in \mathrm{ds}} F_{1}(x)\right) \vee\left(\bigvee_{x \in s_{1} \cup \mathrm{ds}} \mathrm{dF}(x)\right) \tag{А.17}
\end{align*}
$$

from which our result follows because:

$$
\begin{array}{rlr}
m_{2} & =\bigvee_{x \in s_{2}} F_{2}(x) & \\
& =\bigvee_{x \in s_{1} \cup d s}\left(F_{1}(x) \vee d F(x)\right) & \text { equation A. } 15 \\
& =\left(\bigvee_{x \in s_{1}} F_{1}(x)\right) \vee\left(\bigvee_{x \in d s} F_{1}(x)\right) \vee\left(\bigvee_{x \in s_{1} \cup d s} d F(x)\right) & \text { reassociate } \\
& =m_{1} \vee d m &
\end{array}
$$

So it suffices to show equations A. 16 and A.17. This is mostly a matter of pushing through denotations. For instance, starting with $l_{i}$ :

$$
\begin{align*}
l_{i} & =\llbracket \text { for }(x \in e) f \rrbracket \rho_{i} & & \\
& =\operatorname{collect}(\llbracket f \rrbracket)\left(\rho_{i}, \llbracket e \rrbracket \rho_{i}\right) & & \text { definition of } \llbracket \text { for... } \rrbracket \\
& =\bigvee_{x \in \llbracket e \rrbracket \rho_{i}} \llbracket f \rrbracket\left(\rho_{i}, x \mapsto x\right) & & \text { definition of collect } \\
& =\bigvee_{x \in s_{i}} F_{i}(x) & & \text { equation A.13 }
\end{align*}
$$

And $m_{i}$ ：

$$
\begin{aligned}
& m_{i}=\llbracket \text { for }(x \in \phi e) \text { let }[d x]=[0 x] \text { in } \phi f \rrbracket \gamma_{i} \\
& =\bigvee_{x \in \llbracket \phi e \rrbracket \gamma_{i}} \llbracket \text { let }[d x]=[0 x] \text { in } \phi f \rrbracket\left(\gamma_{i}, x \mapsto x\right) \\
& =\bigvee_{x \in s_{i}} \llbracket \phi \ddagger \rrbracket\left(\gamma_{i}, x \mapsto x, \mathrm{~d} x \mapsto \llbracket[0 x] \rrbracket\left(\gamma_{i}, x \mapsto x\right)\right) \quad \text { equation A. } 13 \\
& =\bigvee_{x \in s_{i}} \llbracket \phi f \rrbracket\left(\gamma_{i}, x \mapsto x, \mathrm{~d} x \mapsto \llbracket 0 x \rrbracket\left(\text { strip }\left(\gamma_{i}, x \mapsto x\right)\right)\right) \quad \text { lemma } 33 \\
& =\bigvee_{x \in s_{i}} \llbracket \phi f \rrbracket\left(\gamma_{i}, x \mapsto x, d x \mapsto \text { dummy } x\right) \quad \text { pushing definitions } \\
& =\bigvee_{x \in s_{i}} F_{i}(x)
\end{aligned}
$$

And finally， dm ．By expanding in the same manner as directly above we have that：

$$
\begin{gathered}
\llbracket \text { for }(x \in \delta e) \text { let }[d x]=[0 x] \text { in } \phi f \rrbracket\left(\gamma_{1}, d \gamma\right)=\bigvee_{x \in d s} F_{1}(x) \\
\llbracket \text { for }(x \in \phi e \vee \delta e) \text { let }[d x]=[0 x] \text { in } \delta f \rrbracket\left(\gamma_{1}, d \gamma\right)=\bigvee_{x \in s_{1} \cup d s} d F(x)
\end{gathered}
$$

And therefore：

$$
\begin{aligned}
\mathrm{dm} & =\llbracket \delta(\text { for }(x \in e) f) \rrbracket\left(\gamma_{1}, \mathrm{~d} \gamma\right) \\
& =\llbracket \text { for }(x \in \delta e) \text { let }[\mathrm{d} x]=[0 x] \text { in } \phi f \rrbracket\left(\gamma_{1}, \mathrm{~d} \gamma\right) \\
& \vee \llbracket \text { for }(x \in \phi e \vee \delta e) \text { let }[d x]=[0 x] \text { in } \delta f \rrbracket\left(\gamma_{1}, \mathrm{~d} \gamma\right) \\
& =\left(\bigvee_{x \in \mathrm{ds}} F_{1}(x)\right) \vee\left(\bigvee_{x \in s_{1} \cup \mathrm{ds}} \mathrm{dF}(x)\right)
\end{aligned}
$$

Which is what we wished to show．

$$
\text { Case } \frac{\Gamma \vdash e: \square(\underset{\mathrm{HX}}{\mathrm{~L}} \rightarrow \underset{\mathrm{HX}}{\mathrm{~L}})}{\Gamma \vdash f i x e: \text { Recall that }}
$$

$$
\begin{aligned}
& \phi(\text { fix } e)=\text { semifix } \phi e \\
& \delta(f i x e)=\perp
\end{aligned}
$$

For brevity，let

$$
\begin{aligned}
f & =\llbracket e \rrbracket \rho & f^{\prime} & =\llbracket e \rrbracket \rho^{\prime} \\
\left(g_{1}, g_{2}\right) & =\llbracket \phi e \rrbracket \gamma & \left(g_{1}^{\prime}, g_{2}^{\prime}\right) & =\llbracket \phi e \rrbracket \gamma^{\prime}
\end{aligned}
$$

What we wish to show is：

$$
\begin{aligned}
& \llbracket \perp \rrbracket(\gamma, \mathrm{d} \gamma) \triangleright_{\mathrm{E} \mid} \llbracket \text { semifix } \phi e \rrbracket \gamma \text { 文 } \llbracket \mathrm{fix} e \rrbracket \rho \rightarrow \llbracket \text { semifix } \phi e \rrbracket \gamma^{\prime} \text { 々 } \llbracket \mathrm{fix} e \rrbracket \rho^{\prime} \\
& \Longleftrightarrow \perp \triangleright_{\mathrm{L} \mid} \text { semifix }(\llbracket \phi e \rrbracket \gamma)\left\langle\text { fix } ( \llbracket e \rrbracket \rho ) \rightarrow \operatorname { s e m i f i x } ( \llbracket \phi e \rrbracket \gamma ^ { \prime } ) \left\langle\text { fix }\left(\llbracket e \rrbracket \rho^{\prime}\right)\right.\right. \\
& \Longleftrightarrow \perp \triangleright_{\text {FIX }} \text { semifix }\left(g_{1}, g_{2}\right) \gtreqless \text { fix } f \rightarrow \operatorname{semifix}\left(g_{1}^{\prime}, g_{2}^{\prime}\right) \gtreqless \text { fix } f^{\prime} \\
& \Longleftrightarrow \operatorname{semifix}\left(g_{1}, g_{2}\right)=\text { fix } f=\operatorname{semifix}\left(g_{1}^{\prime}, g_{2}^{\prime}\right)=\text { fix } f^{\prime} \quad \text { (applying lemma 27) }
\end{aligned}
$$

 and expanding gives us:

$$
\begin{align*}
& f=f^{\prime} \text { and } g_{1}=g_{1}^{\prime} \text { and } g_{2}=g_{2}^{\prime}  \tag{A.18}\\
& \mathrm{g}_{2} \triangleright_{\mathrm{fIX} \rightarrow \mathrm{FX}}^{\mathrm{L}} \mathrm{~g}_{1} \text { \& } \mathrm{f} \rightarrow \mathrm{~g}_{1}^{\prime} \text { \& } \mathrm{f}^{\prime} \tag{A.19}
\end{align*}
$$

Applying equation A. 18 reduces our goal to showing semifix $\left(g_{1}, g_{2}\right)=$ fix $f$, which holds by lemma 34 applied to equations A. 18 and A.19.

$$
\begin{aligned}
& \text { Case } \frac{\Gamma \vdash e: \square\left(A_{1}+A_{2}\right)}{\Gamma \vdash \text { split e }: \square A_{1}+\square A_{2}} \text {. Recall that } \\
& \phi(\text { split } \mathbf{e})=\text { let }[z]=\phi \mathbf{e} \text { in } \\
& \text { case split }\left[\pi_{1} z\right] \text { of } \\
& \left(\mathrm{in}_{\mathrm{i}} \mathrm{Y} \rightarrow \text { let }[\mathrm{x}]=\mathrm{Y}\right. \text { in } \\
& \text { case split }\left[\pi_{2} z\right] \text { of } \\
& \mathrm{in}_{\mathrm{i}} \mathrm{DY} \rightarrow \text { let }[\mathrm{d} x]=\mathrm{DY} \text { in } \mathrm{in}_{\mathrm{i}}[(\mathrm{x}, \mathrm{~d} x)] \\
& \left.\mathrm{in}_{\mathrm{i}+1 \bmod 2}{ }_{-} \rightarrow \mathrm{in}_{\mathrm{i}}[(\mathrm{x}, \text { dummy } \mathrm{x})]\right)_{i} \\
& \delta(\text { split } e)=\text { let }[y]=\phi e \text { in } \\
& \text { case } \pi_{1} y \text { of }\left(\mathrm{in}_{\mathrm{i}_{-}} \rightarrow \mathrm{in} \mathrm{i}_{\mathrm{i}}()\right)_{\mathrm{i} \in\{1,2\}}
\end{aligned}
$$

By unpacking our inductive hypothesis there must be some $k \in\{1,2\}$ and some $d x, x, a$ such that $d x \triangleright_{A_{k}} x \nmid a \rightarrow x \notin a$ and:

$$
\llbracket e \rrbracket \rho=\llbracket e \rrbracket \rho^{\prime}=\mathrm{in}_{\mathrm{k}} \mathrm{a} \quad \llbracket \phi e \rrbracket \gamma=\llbracket \phi e \rrbracket \gamma^{\prime}=\left(\mathrm{in}_{\mathrm{k}} \mathrm{x}, \mathrm{in}_{\mathrm{k}} \mathrm{~d} x\right) \quad \llbracket \delta e \rrbracket(\gamma, \mathrm{~d} \gamma)=()
$$

Applying the definition of what we wish to show, it therefore suffices to show that:

$$
\begin{align*}
\llbracket \text { split } e \rrbracket \rho=\llbracket \text { split } e \rrbracket \rho^{\prime} & =\mathrm{in}_{\mathrm{k}} \mathrm{a}  \tag{A.20}\\
\llbracket \phi(\text { split } e) \rrbracket \gamma=\llbracket \phi(\text { split } e) \rrbracket \gamma^{\prime} & =\mathrm{in}_{k}(\mathrm{x}, \mathrm{~d} x)  \tag{A.21}\\
\llbracket \delta(\text { split } e) \rrbracket(\gamma, \mathrm{d} \gamma) & =\mathrm{in}_{\mathrm{k}}() \tag{A.22}
\end{align*}
$$

Showing this is a matter of calculation. Equation A. 20 is immediate upon noting that dist ${ }_{+}^{\square}=\mathrm{id}$ and therefore $\llbracket$ split $e \rrbracket \rho=\llbracket e \rrbracket \rho$. The other two are no more difficult but considerably more tedious, so we omit the details of the calculations.

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[^0]:    ${ }^{1}$ The classic algorithm for single-source shortest paths, Dijkstra's algorithm, can be seen as a version of the algorithm we've described, but with a crucial optimization: it prioritizes which edges to consider next in a way that guarantees it never needs to revisit a node.

[^1]:    ${ }^{2}$ Although for simplicity's sake we have presented our partial orders as ranging over booleans, distances, and finite sets respectively, our states are actually maps from nodes or lines into booleans, distances, or finite sets. Fortunately, if the domain is finite, maps into a poset satisfying ACC also satisfy ACC.

[^2]:    contrived, the problem in general is fundamental: without further limitations on clauses, proof search is only semi-decidable, so while complete search strategies can guarantee finding all proofs, they cannot guarantee they'll halt after doing so. This is particularly important for the handling of negation-as-failure; see §1.2.2.

[^3]:    5 https://web.archive.org/web/20210727023523/https://semmle.com/case-studies/ semmle-nasa-landing-curiosity-safely-mars
    6 https://web.archive.org/web/20211130053831/https://msrc-blog.microsoft.com/2018/08/16/ vulnerability-hunting-with-semmle-ql-part-1/

    7 https://web.archive.org/web/20210624221802/https://semmle.com/case-studies/ semmle-nasdaq-improving-roi-and-reducing-time-market

[^4]:    ${ }^{1}$ To implement set types, their elements must support decidable equality. In our core calculus, we use a subgrammar of "eqtypes", and in our implementation (which compiles to Haskell) we use typeclass constraints to pick out such types.

[^5]:    ${ }^{2}$ We are sweeping a few technical details under the rug here. First, for reasons which will not be explained until $\S 3.3 .5$ we treat fix as if it were a non-monotone operator. Second, observe that the finite set type $\left\{\begin{array}{l}\text { A }\}\end{array}\right\}$ will possess infinite ascending chains if $\underset{\in \in}{A}$ has infinitely many inhabitants. Thus we need to distinguish a class of finite eqtypes A. Although their grammars in figure 2.1 are identical, their intent is different. For example, if we extended Datafun with integers, they would form an eqtype, but not a finite one.

[^6]:    ${ }^{3}$ Technically the inclusion order on sets of integer pairs does not satisfy the ascending chain condition, so this use of trans is not well-typed. However, since the positions in a particular string form a finite set, semantically there is no issue.

[^7]:    ${ }^{5}$ Note that the subset ordering completely ignores the element ordering $\leqslant_{A}$. There are orderings on $\mathcal{P}_{\text {fin }} A$ which are not so forgetful; for instance, the free semilattice FA consists of finite sets ordered by $s \leqslant_{\text {FA }} \mathrm{t} \Longleftrightarrow$ $(\forall x \in s, \exists y \in t) x \leqslant A y$ and quotiented by antisymmetry (this may also be seen as the semilattice of finitelygenerated downward closed sets under union, or of finite antichains). Our $P A$ is isomorphic to $F \square A$. However, Datalog's semantics use only the inclusion order, as do all of our motivating examples; so for simplicity we have stuck to it.
    ${ }^{6}$ In fact, fix is monotone and could be regarded as a map $(\mathrm{L} \Rightarrow \mathrm{L}) \rightarrow \mathrm{L}$, but because the typing rule for fix boxes its argument, we do the same here.

[^8]:    ${ }^{1}$ In this section we do not bother distinguishing monotone variables $X$ or discrete expressions $e$, as it muddies our examples to no benefit.

[^9]:    ${ }^{2}$ Indeed, sets have not only zero changes but all increasing changes: for any $x \leqslant y$ there is a $d x$ such that

[^10]:    $\mathrm{d} x \triangleright x \hookrightarrow \mathrm{y}:\{\underset{\in Q}{ } \underset{\varepsilon}{ }\}$; for instance one may let $\mathrm{d} x=\mathrm{y} \backslash x$, or indeed just y . We call this property completeness, as it is the converse of soundness. However, while our change structure for sets is complete, we will later observe that completeness is troublesome at function types, so we do not insist on it in general.

[^11]:    ${ }^{3}$ Note that had we chosen to let $\Delta(A \times B)=\Delta A+\Delta B$, representing a change to a tuple by a change to only one of its components, this would not allow us to differentiate tupling $\langle f, g\rangle$, since a change to the input may cause both components of the output to change simultaneously.

[^12]:    ${ }^{4}$ We could avoid partiality by defining $\mathrm{in}_{\mathrm{i}} \mathrm{dx} \triangleright \mathrm{in}_{\mathrm{j}} \mathrm{x} \hookrightarrow \mathrm{i} \mathrm{n}_{\mathrm{j}} x: A_{1}+A_{2}$ for $\mathfrak{i} \neq \mathfrak{j}$; that is, treating currently
    "invalid" changes as zero-changes. This unfortunately doesn't extend to the function case, which as we'll see shortly also needs a partial validity relation. Moreover, it doesn't ensure uniqueness of $\left[f_{1}, f_{2}\right]^{\prime}$ : although it requires $\left[f_{1}, f_{2}\right]^{\prime}\left(i_{i} x\right)\left(i n_{j} d x\right)$ to be a zero change to $f_{i} x$ when $\mathfrak{i} \neq \mathfrak{j}$, there may be multiple such zero changes.

[^13]:    ${ }^{5}$ We further discuss the issue of monotonicity and pointwise changes in §5.2.2.

[^14]:    7 The need to recompute fix f could likely be solved by caching intermediate values, which we discuss further in §5.2.1. Somewhat unusually, in this case we want to cache the previous output of an operation rather than its previous input.

[^15]:    9 For notational convenience we assume that source programs contain no variables starting with the letter $d$.

[^16]:    1 http://gallium.inria.fr/~fpottier/menhir/

[^17]:    2 The diff method on Semilat and the semifixMinimized function that uses it will be explained in §4.3.
    ${ }^{3}$ We did not use Haskell's built-in Ord typeclass for this because it is intended for total orders, not partial ones.

[^18]:    ${ }^{5}$ Given Datafun's monotonicity typing, the reader may wonder whether and with respect to what $(\backslash)_{\text {L }}$ must be monotonic. In practice, on finite sets $d x \backslash x$ is monotone in $d x$ but not $x$, and this pattern will hold for our other semilattice types as well. However, monotonicity in either argument is not required for correctness.

[^19]:    ${ }^{2}$ In the original presentation by Cai et al. (2014) these operators (except $\odot$, which is not present) are taken as primary rather than as additions to a basic change structure, and the validity relation $d x \triangleright x_{1} \hookrightarrow x_{2}: S$ is reduced to a set of valid changes $\Delta_{S} x_{1} \subseteq \Delta S$. Datafun's approach to seminaïve computation was originally inspired by this paper, but we moved to a validity-relation approach because of the difficulties of defining these operators in a monotonicity-aware setting; thus the validity relation approach to change structures appears to have been indendently invented in both Datafun and in Giarrusso's work.

[^20]:    However, this passes the monotone variable $X$ to the function change $d f$, which takes it as a discrete argument it does not type-check! Indeed, it is not hard to come up with functions $f, d f$ such that the result of $f \oplus d f$ as defined above is not a monotone function.

    These difficulties might be overcome with further careful work; for example, it should be possible to prove that $f \oplus d f$ is monotone so long as $d f$ is a valid change to $f$. This would make $\oplus$, like semifix, a "trusted primitive" whose implementation cannot be expressed in Datafun itself.

[^21]:    ${ }^{5}$ In fact, it is an exemplar of a large class of self-maintainable functions: join-distributive maps.

[^22]:    ${ }^{1}$ Here is a simple system along these lines: introduce a monad Bless and type constructor Holy with the methods:

[^23]:    ${ }^{3}$ https://web.archive.org/web/20220223155533/https://reactjs.org/; see also https://web.archive. org/web/20220907152744/https://blog.janestreet.com/incrementality-and-the-web/ for a discussion of incrementality in web UI libraries.

