## **Relating Complexity and Precision in Control Flow Analysis**

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## Abstract

We analyze the computational complexity of *k*CFA, a hierarchy of control flow analyses that determine which functions may be applied at a given call-site. This hierarchy specifies related decision problems, quite apart from any algorithms that may implement their solutions. We identify a simple decision problem answered by this analysis and prove that in the 0CFA case, the problem is complete for polynomial time. The proof is based on a nonstandard, symmetric implementation of Boolean logic within multiplicative linear logic (MLL). We also identify a simpler version of 0CFA related to  $\eta$ -expansion, and prove that it is complete for logarithmic space, using arguments based on computing paths and permutations.

For any fixed k > 0, it is known that kCFA (and the analogous decision problem) can be computed in time exponential in the program size. For k = 1, we show that the decision problem is NP-hard, and sketch why this remains true for larger fixed values of k. The proof technique depends on using the *approximation* of CFA as an essentially nondeterministic computing mechanism, as distinct from the exactness of normalization. When k = n, so that the "depth" of the control flow analysis grows linearly in the program length, we show that the decision problem is complete for exponential time.

In addition, we sketch how the analysis presented here may be extended naturally to languages with control operators. All of the insights presented give clear examples of how straightforward observations about linearity, and linear logic, may in turn be used to give a greater understanding of functional programming and program analysis.

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### 1. Introduction

We investigate the *precision* of static, compile-time analysis, and the necessary analytic *tradeoff* with the computational *resources* that go into the analysis.

Control flow analysis provides a fundamental and ubiquitous static analysis of higher-order programs. Heintze and McAllester (1997b) point out that "in fact, some form of CFA is used in most forms of analyses for higher-order languages." Control flow analysis answers the following basic questions (Palsberg 1995):

- 1. For every application, which abstractions can be applied?
- 2. For every abstraction, to which arguments can it be applied?

These questions specify well-defined, significant *decision problems*, quite apart from any algorithm proposed to solve them. What is the inherent computational difficulty of solving these problems? What do they have to do with normalization (evaluation), as opposed to approximation? Where is the fulcrum located between exact computation and approximation to it, and how does that balance make this compile-time program analysis tractable?

To answer either of the enumerated questions above, we must be able to approximate the set of values to which any given subexpression may evaluate. This approximation can be formulated as a *flows to* relation between fragments of program text—the set of values a subexpression may evaluate to, then, is the set of appropriately defined (program text) values that flow into that subexpression. This analysis possibly includes "false positives" (values that *may* flow to a call site, but in fact do not), and thus a *flows to* analysis might relate every program fragment. We focus attention on an acceptably "least" analysis, that is, a solution to the constraints which has a minimum of false positives. More precision means fewer false positives.

To ensure tractability of any static analysis, there has to be an *approximation* of something, where information is deliberately *lost* in the service of providing what's left in a reasonable amount of time. A good example of what is lost during static analysis is that the information gathered for each occurrence of a bound variable is merged. When variable f occurs twice in function position with two different arguments, and a substitution of a function is made for f, a monovariant CFA will blur which copy of the function is applied to which argument.

Further refining the relatively coarse approximation of the above control flow analysis, a hierarchy of increasingly polyvariant analyses was developed by Olin Shivers in his Ph.D. thesis (Shivers 1991), the so-called kCFA analyses, of which the above analysis marks the base—0CFA. A 1CFA, for example, uses a *contour* to distinguish a more specific and dynamic calling context in the analysis of each program point; kCFA, then, distinguishes between k levels of such contexts. Moving up the hierarchy increases the precision of this analysis, by constructing more elaborate contours. However, this increased precision is accompanied by an empiri-

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cally observed increase in cost. As Shivers observed in his "Best of PLDI" retrospective on the kCFA work:

It did not take long to discover that the basic analysis, for any k > 0, was intractably slow for large programs. In the ensuing years, researchers have expended a great deal of effort deriving clever ways to tame the cost of the analysis.

**Technical contributions:** We identify a simple decision problem answered by control flow analysis and prove that in the 0CFA case, the problem is complete for polynomial time. The proof is based on a nonstandard, symmetric implementation of Boolean logic within multiplicative linear logic (MLL). We also identify a simpler version of 0CFA related to  $\eta$ -expansion, and prove that it is complete for logarithmic space, using arguments based on computing paths and permutations.

Moreover, we show that the decision problem for 1CFA is NP-hard. Given that there is a naive exponential algorithm, we conjecture that it is in fact NP-complete, where prudent guessing in the computation of the naive algorithm can answer specific questions about flows. The proof generalizes to an NP-hardness proof for all fixed k > 1, demonstrating that from a complexity point of view, the hierarchy is flat for all fixed k > 0.

Like all good proofs, this NP-hardness result has simple intuitions. 1CFA depends on *approximation*, where multiple values (here, closures) flow to the same program point, including false positives. The bottleneck of the naive algorithm is its handling of closures with many free variables. For example,  $\lambda w.wx_1x_2\cdots x_n$ has *n* free variables, with an exponential number of possible associated environments mapping these variables. Approximation allows us to bind each  $x_i$ , independently, to either closed  $\lambda$ -terms for "true" or "false". In turn, application to an *n*-ary Boolean function, as analyzed in CFA, must then evaluate the function on each of the possible environments. Asking whether "true" can flow out of the call site then becomes a way of asking if the Boolean function is satisfiable.

No such "pseudo parallelism" would be possible in an exact normalization—it is the existence of *approximation* that mashes these distinct closures together. Similar NP-hardness of kCFA for k > 1 results by suitably "padding" the construction for 1CFA so as to render the added precision useless. We remark also that the NP-hardness construction cannot be coded in 0CFA because in that case, there are no environments and hence no closures.

Despite being the fundamental analysis of higher-order programs, despite being the subject of investigation for over twentyfive years (Jones 1981), and the great deal of expended "effort deriving clever ways to tame the cost", there has remained a poverty of analytic knowledge on the complexity of control flow analysis and the kCFA hierarchy, the essence of how this hierarchy is computed, and where the sources of approximation occur that make the analysis work. This paper is intended to repair such lacunae in our understanding of program analysis.

## 2. Preliminaries

In this section, we describe the programming language that is the subject of our control flow analysis and provide the necessary mechanics of the graphical representation of programs employed in our algorithms. The language on which the control flow analysis will be performed is the untyped  $\lambda$ -calculus extended by a labelling scheme serving to index subexpressions of a program. Following Nielson et al. (1999), we define the following syntactic categories:



 $t \in \mathbf{Term}$  terms (or unlabeled expressions)

A countably infinite set of labels (Lab) is assumed and for simplicity we suppose the set of variable names are included in Lab.



Figure 1. Graphical coding of expressions.

The syntax of the language is given by the following grammar:<sup>1</sup>

$e ::= t^{\ell}$	expressions (or labeled terms)
$t ::= x \mid (e \ e) \mid (\lambda x.e)$	terms (or unlabeled expressions)

Expressions are represented graphically as follows: a graph consists of ternary apply (@), abstraction ( $\lambda$ ), and sharing nodes ( $\nabla$ ); unary weakening nodes ( $\odot$ ); and wires between these nodes. Each node has a distinguished *principal port*; ternary node's other *auxilary ports* are distinguished as the left and right ports. We call the principal port of an @-node the *function*, the left port the *continuation*, and the right port the *argument*. Likewise, the principal port of a  $\lambda$ -node is the *root*, the left port is the *body*, and the right port is the *parameter*. That is:



Wires are labeled to correspond with the labels on terms, i.e. the continuation wire for the graph of  $(e \ e)^{\ell}$  will be labeled  $\ell$  and the root wire for  $(\lambda x.e)^{\ell}$  will be labeled  $\ell$ . Parameter wires will be labeled with the bound variable's name.

Figure 1 gives the graphical coding of expressions. In the  $\lambda x.e$ case, the figure is drawn as though x appears twice in e, and thus the sharing node is used to duplicate the wire, one going to each occurrence. The wire between the  $\lambda$ -node and the sharing node is implicitly labeled x. The two wires attached to the auxiliary ports of the sharing node will be labeled with the distinct labels of each occurrence of x. If x occurred more than twice, more sharing nodes would be attached to fan out the binding the appropriate amount. If the bound variable occurred exactly once, the wire would connect the  $\lambda$  node directly to the variable occurrence; the label x and the label used at the occurrence of x would both refer to the same wire. If the bound variable x did not occur in e, the wire from the  $\lambda$ -node would attach to a weakening node. The "dangling wire" from the graph of the body of the function denotes a variable free in e. In the case of  $(e_0 \ e_1)$ , the figure is drawn with  $e_0$  and  $e_1$  both having a free variable in common, i.e. there is an x in  $\mathbf{fv}(e_0) \cap \mathbf{fv}(e_1)$ , and both  $e_0$  and  $e_1$  have another free variable not occuring in the other.

## 3. 0CFA

Control flow analysis seeks to answer questions such as "what functions can be applied at a give application position?" Because both functions and applications may be copied, such questions become ambiguous. When one points to an application in the program text, running the program may duplicate this point several times, so which of the copies are we really asking about? Likewise, if a lambda term is said to flow into some application, which copy of the term is going to be applied? The answer 0CFA gives is: all of them—all copies of a term are identified. Later, we see how con-

<sup>&</sup>lt;sup>1</sup> Unlike Nielson et al. (1999), constants, binary operators, and recursiveand let-binding syntax are omitted. These language features add nothing interesting to the computational complexity of the analysis.



Figure 2. CFA virtual wire propagation rules.

textual information can be used to distinguish the copies and give more precise answers to these questions.

We follow Nielson et al. (1999) and say the result of 0CFA is an *abstract cache*  $\hat{C}$  associating abstract values with each labeled program point. More precisely:

$\widehat{v}$	$\in$	Val	=	$\mathcal{P}(\mathbf{Term})$	abstract values
ĉ	$\in$	Cache	=	$\mathbf{Lab}  ightarrow \widehat{\mathbf{Val}}$	abstract caches

An *abstract cache* maps a program label to an *abstract value*  $\hat{v}$ , a set of lambda expressions, which represents the set of (textual) values that may *flow into* that label's subexpression during evaluation. Similarly, the *abstract environment* maps a variable to an abstract value, which represents the set of (textual) values that may be bound to that variable during evaluation.

An acceptable control flow analysis for an expression e is written  $\widehat{C} \models e$ . Recalling again from Nielson et al. (1999), the acceptability relation is given by the greatest fixed point of the functional defined according to the following clauses:

$$\begin{array}{lll} \widehat{\mathsf{C}} & \models & x^{\ell} \operatorname{iff} \widehat{\mathsf{C}}(x) \subseteq \widehat{\mathsf{C}}(\ell) \\ \widehat{\mathsf{C}} & \models & (\lambda x.e)^{\ell} \operatorname{iff} (\lambda x.e) \in \widehat{\mathsf{C}}(\ell) \\ \widehat{\mathsf{C}} & \models & (t_{1}^{\ell_{1}} t_{2}^{\ell_{2}})^{\ell} \operatorname{iff} \widehat{\mathsf{C}} \models t_{1}^{\ell_{1}} \wedge \widehat{\mathsf{C}} \models t_{2}^{\ell_{2}} \wedge \forall (\lambda x.t_{0}^{\ell_{0}}) \in \widehat{\mathsf{C}}(\ell_{1}) : \\ & & \widehat{\mathsf{C}} \models t_{0}^{\ell_{0}} \wedge \widehat{\mathsf{C}}(\ell_{2}) \subseteq \widehat{\mathsf{C}}(x) \wedge \widehat{\mathsf{C}}(\ell_{0}) \subseteq \widehat{\mathsf{C}}(\ell) \end{array}$$

We now describe an algorithm for performing control flow analysis that is based on the graph coding of terms. The graphical formulation consists of generating a set of *virtual paths* for a program graph. Virtual paths describe an approximation of the real paths that will arise during program execution.

Figure 2 defines the virtual path propagation rules. The left hand rule states that a virtual wire is added from the continuation wire to the body wire and from the variable wire to the argument wire of each  $\beta$ -redex. The right hand rule states analogous wires are added to each *virtual*  $\beta$ -redex—an apply and lambda node connected by a virtual path. There is a *virtual path* between two wires  $\ell$  and  $\ell'$ , written  $\ell \rightsquigarrow \ell'$  in a CFA-graph iff: 1)  $\ell \equiv \ell'$ , 2) there is a virtual wire from  $\ell$  to  $\ell'$ , 3)  $\ell$  connects to an auxiliary port and  $\ell'$  connects to the principal port of a sharing node, or 4)  $\ell \rightsquigarrow \ell''$  and  $\ell'' \rightsquigarrow \ell'$ .

Some care must be taken to ensure leastness when propagating virtual wires. In particular, wires are added only when there is a virtual path between a *reachable* apply and a lambda. An apply node is reachable if it is on the spine of the program, i.e., if  $e = (\cdots ((e_0e_1)^{\ell_1}e_2)^{\ell_2}\cdots e_n)^{\ell_n}$  then the apply nodes with continuation wires labeled  $\ell_1, \ldots, \ell_n$  are reachable, or it is on the spine of an expression with a virtual path from a reachable apply node.

The graph-based analysis can now be performed in the following way: construct the CFA graph according to the rules in Figure 2, then define  $\widehat{C}(\ell)$  as  $\{(\lambda x.e)^{\ell'} \mid \ell \rightsquigarrow \ell'\}$ . It is easy to see that the algorithm constructs answers that satisfy the acceptability relation specifying the analysis. Moreover, this algorithm constructs least solutions according to the partial order  $\widehat{C} \sqsubseteq_e \widehat{C}'$  iff  $\forall \ell \in \mathbf{Lab}_e : \widehat{C}(\ell) \subseteq \widehat{C}'(\ell)$ , where  $\mathbf{Lab}_e$  denotes the set of labels restricted to those occurring in *e*, the program of interest.



Figure 3. Graph coding and CFA graph.

**Lemma 1.**  $\widehat{C}' \models e$  implies  $\widehat{C} \sqsubseteq_e \widehat{C}'$  for  $\widehat{C}$  constructed for e as described above.

We now consider an example of use of the algorithm. Consider the labeled program:

$$((\lambda f.((f^1f^2)^3(\lambda y.y^4)^5)^6)^7(\lambda x.x^8)^9)^{10}$$

Figure 3 shows the graph coding of the program and the corresponding CFA graph. The CFA graph is constructed by adding virtual wires  $10 \rightsquigarrow 6$  and  $f \rightsquigarrow 9$ , induced by the actual  $\beta$ -redex on wire 7. Adding the virtual path  $f \rightsquigarrow 9$  to the graph creates a virtual  $\beta$ -redex via the route  $1 \rightsquigarrow f$  (through the sharing node), and  $f \rightsquigarrow 9$  (through the virtual wire). This induces  $3 \rightsquigarrow 8$  and  $8 \rightsquigarrow 2$ . There is now a virtual  $\beta$ -redex via  $3 \rightsquigarrow 8 \rightsquigarrow 2 \rightsquigarrow f \rightsquigarrow 9$ , so wires  $6 \rightsquigarrow 8$  and  $8 \rightsquigarrow 5$  are added. This addition creates another virtual redex via  $3 \rightsquigarrow 8 \rightsquigarrow 2 \rightsquigarrow 5$ , which induces virtual wires  $6 \rightsquigarrow 4$  and  $4 \rightsquigarrow 5$ . No further wires can be added, so the CFA graph is complete. The resulting abstract cache gives:

$$\begin{array}{lll} \mathsf{C}(1) = \{\lambda x\} & \mathsf{C}(6) &= \{\lambda x, \lambda y\} \\ \widehat{\mathsf{C}}(2) = \{\lambda x\} & \widehat{\mathsf{C}}(7) &= \{\lambda f\} & \widehat{\mathsf{C}}(f) = \{\lambda x\} \\ \widehat{\mathsf{C}}(3) = \{\lambda x, \lambda y\} & \widehat{\mathsf{C}}(8) &= \{\lambda x, \lambda y\} & \widehat{\mathsf{C}}(x) = \{\lambda x, \lambda y\} \\ \widehat{\mathsf{C}}(4) = \{\lambda y\} & \widehat{\mathsf{C}}(9) &= \{\lambda x\} & \widehat{\mathsf{C}}(y) = \{\lambda y\} \\ \widehat{\mathsf{C}}(5) = \{\lambda y\} & \widehat{\mathsf{C}}(10) = \{\lambda x, \lambda y\} \end{array}$$

We now describe a natural decision problem answered by this control flow analysis. After describing the problem here, subsequent sections will consider variants of it for kCFA.

#### 3.1 The 0CFA decision problem

A decision problem—a question that can be answered with a yes or a no—makes the analysis insensitive to the output size of any control flow analysis. Typically, this analysis computes the answer to questions like "what functions can be applied at a particular call site?" or "what arguments can a particular function be applied to?", so a natural decision problem based on these questions are "is this particular function applied at this particular call site?" or "does this function get applied to this argument?", where the function is denoted by some lambda expression or application in the program text. These questions provide ways of answering the more general "to what values can a subexpression evaluate?"

**Control Flow Problem (0CFA):** Given expressions e,  $(\lambda x.e_0)$ , and label  $\ell$ , is  $(\lambda x.e_0) \in \widehat{C}(\ell)$  in a least analysis of e?

The graphical analogue of this problem point to a  $\lambda$ -node and an application node, and ask if there a *virtual path* (describing a  $\beta$ redex from reductions to take place) from the function port of the apply to the root port of the  $\lambda$ ? Or likewise, point to two  $\lambda$ -nodes: is there a virtual path from the variable port of one to the root of the other? The graph-reduction characterization has the virtue of being free of a lot of notational clutter (like variable renaming).

We now prove that this decision problem is complete for polynomial time. A graph-based argument for containment in PTIME is straightforward:

#### Theorem 1. OCFA is contained in PTIME.

*Proof.* OCFA computes a binary relation over a fixed structure (the graph description of a program). The computation of the relation is monotone: it begins as empty and is added to incrementally. Because the structure is finite, a fixed point must be reached by this incremental computation. The binary relation can be at most polynomial in size, and each increment is computed in polynomial time.

We now turn to the more interesting hardness of the problem. As has been observed by Henglein and Mairson (1991); Mairson (2004); Neergaard and Mairson (2004), linearity is the key ingredient in understanding the lower-bound complexity of analyses. In looking at the control flow analysis algorithm of section 3, we may ask what is the source of approximation? The answer is that every copy of a function is identified, and every copy of an application site is identified. In other words, a program with copying engenders an approximate analysis. More concretely, notice that an application site has multiple virtual paths to a value only through virtual paths that pass through sharing nodes. If there are no sharing nodes-as in a linear term, a term where every bound variable occurs exactly once-it is simple to see that there is only a single virtual path. By the conservativity of the analysis, it follows immediately that if there is a single path from an apply node to a lambda node, then these nodes will fuse during reduction. Thus, linearity subverts approximation and renders control flow analysis synonymous with normalization. A lower bound on the complexity of the analysis, then, is the expressivity of the language it normalizes. Proving these lower bounds amounts to functional hacking within the constraints of linearity.

Recall the canonical PTIME-complete decision problem (Ladner 1975):

**Circuit Value Problem:** Given a Boolean circuit C of n inputs and one output, and truth values  $\vec{x} = x_1, \ldots, x_n$ , is  $\vec{x}$  accepted by C?

A problem is PTIME-hard if any instance of it can be compiled using only  $O(\ln |x|)$  space into an instance of the circuit value problem. We now show how to program circuits using only linear terms, proving 0CFA to be PTIME-hard.

As a calculational tool and *lingua franca* amongst functional programmers (and compilers), we use a linear fragment of Standard ML to illustrate our constructions. The Boolean values True and False are built out of the constants TT and FF:

```
- fun TT (x:'a,y:'a)= (x,y);
val TT = fn : 'a * 'a -> 'a * 'a
- fun FF (x:'a,y:'a)= (y,x);
val FF = fn : 'a * 'a -> 'a * 'a
- val True= (TT: ('a * 'a -> 'a * 'a),
FF: ('a * 'a -> 'a * 'a));
val True = (fn,fn) : ('a * 'a -> 'a * 'a)
* ('a * 'a -> 'a * 'a)
TT: ('a * 'a -> 'a * 'a);
val False = (ff,fn) : ('a * 'a -> 'a * 'a);
* ('a * 'a -> 'a * 'a);
```

This little hack will print out what Boolean we are talking about:

The way we compute And is to use the famous 1930s-era coding of conjunction (hacked by Alonzo Church?) in the first component, and the disjunction in the second component. That way, we are guaranteed that the junk in v and v' is symmetric: one is TT, and the other is FF. Then function composition can be used to erase the symmetric garbage—the slogan is, "symmetric garbage is self-annihilating."

```
- fun And (p,p') (q,q')=
    let val ((u,v),(u',v')) = (p (q,FF), p' (TT,q'))
    in (u,Compose (Compose (u',v),Compose (v',FF)))
    end;
val And = fn
    : ('a * ('b * 'b -> 'b * 'b) -> 'c * ('d -> 'e))
    * (('f * 'f -> 'f * 'f) * 'g
                          -> ('e -> 'h) * ('i * 'i -> 'd))
                          -> 'a * 'g -> 'c * ('i * 'i -> 'h)
- Show (And True False);
val it = (false,true) : bool * bool
```

Notice that since p' is the complement of p, and q' that of q, we know u' is the complement of u. Composing v, v' and FF is always the *identity function* TT, which can then be composed with u' without changing the value of u'.

The construction of the Or term is symmetric to the And term, the Not term is just an inversion<sup>2</sup>—all this an obvious consequence of deMorgan duality and the construction here of the Booleans. The Copy gate uses the fact that either p or p' will invert its argument, so that either ((TT,FF),(TT,FF)) or ((FF,TT),(FF,TT)) is returned:

By writing logic gates in continuation-passing style, for example:

- fun Andgate p q k= k (And p q)

we can then write circuits that look like straight-line code:

<sup>&</sup>lt;sup>2</sup> Both are omitted for space reasons.

The above code says: compute the and of e2 and e3, putting the result in register  $e7, \ldots$ , make two copies of register f, putting the values in registers e9 and  $e10, \ldots$ , compute the or of e11 and e12, putting the result in the output register.

Now for the decision problem: the pair (x,y) is either the pair (f,g) or (g,f), depending on the Boolean circuit computing output u. Thus, is f applied to a and g to b, or the other way around? Because the computation is entirely linear, the set of possible binders to x is—this must be emphasized—not {f, g}: it is precisely {f}, or {g}. The 0CFA "approximation" is in fact an exact normalization.

**Theorem 2.** OCFA is complete for PTIME.

### **4.** *k***CFA**

Increasing the precision of the coarse approximation given by the above control flow analysis, a hierarchy of more and more refined analyses was developed. These analyses are *context-sensitive* or *polyvariant* because they distinguish functions applied in distinct call sites to increase the precision of the analysis. A 1CFA analysis, for example, uses a *contour* to distinguish *one* level of dynamic calling context in the analysis of each program point.

Contours  $\delta$  are strings of labels of length at most k; they serve to record the last k dynamic call points and thus distinguish instances of variables and program points. Contour environments map variable names to contours. Contours describe the context in which an instance of a term evaluates, and contour environments, similarly describe the context under which a variable was bound.

For example, in the expression  $(e_0((\lambda x.e_1)e_2)^{\ell_1})^{\ell_2}$ , the subexpression  $e_1$  evaluates in contour  $\ell_2\ell_1$ . Likewise, x is bound in contour  $\ell_2\ell_1$ .

$$\begin{array}{rcl} \delta & \in & \Delta & = & \mathbf{Lab}^{\leq k} & \text{contour} \\ ce & \in & \mathbf{CEnv} & = & \mathbf{Var} \to \Delta & \text{contour environment} \end{array}$$

Abstract values are extended to pairs of (textual) program values and contour environments closing the term, i.e. abstract closures. The environment maps variable names to the contours in place at the definition point for the free variables. The abstract cache now maps a label and a contour to an abstract (closure) value. That is:

$$\begin{array}{rcl} v & \in & \widehat{\mathbf{Val}} & = & \mathcal{P}(\mathbf{Term} \times \mathbf{CEnv}) & \text{abstract values} \\ \widehat{\mathsf{C}} & \in & \widehat{\mathbf{Cache}} & = & (\mathbf{Lab} \times \Delta) \to \widehat{\mathbf{Val}} & \text{abstract caches} \end{array}$$

An acceptable k-level control flow analysis for an expression e is written  $\widehat{C} \models_{\delta}^{ce} e$ , which states that  $\widehat{C}$  is an acceptable analysis of e in the context of the current environment ce and current contour  $\delta$  (for the top level analysis of a program, these will both be empty).

Colloquially we understand these judgments as follows: when running the program, fragments of the original program text may be duplicated, such as when a function copies its argument:

$$((\lambda f. \cdots (fe_1)^{\ell_1} \cdots (fe_2)^{\ell_2} \cdots)(\lambda x.e))$$

Contours distinguish between these copies (for copies created via at most k application points, beyond this the distinction in copies is blurred), so when judging an analysis correct for a program

fragment, we use  $\delta$  to tell us which copy of the text is being analyzed, and *ce* to tell us which copies of the free variables in this copy of the program are being analyzed. So in the above example, we have seperate judgements  $\widehat{C} \models_{\ell_1}^{x \mapsto \ell_1} e$  and  $\widehat{C} \models_{\ell_2}^{x \mapsto \ell_2} e$  as a way of talking about seperate run-time instances of the expression *e* that occurs statically only once.

The acceptability relation is given by the greatest fixed point of the functional defined according to the following clauses (again we are concerned only with least solutions):<sup>3</sup>

$$\begin{split} \widehat{\mathsf{C}} &\models_{\delta}^{ce} \quad x^{\ell} \text{ iff } \widehat{\mathsf{C}}(x, ce(x)) \subseteq \widehat{\mathsf{C}}(\ell, \delta) \\ \widehat{\mathsf{C}} &\models_{\delta}^{ce} \quad (\lambda x.e)^{\ell} \text{ iff } \langle (\lambda x.e), ce_0 \rangle \in \widehat{\mathsf{C}}(\ell, \delta) \\ & \text{where } ce_0 = ce |\mathbf{fv}(\lambda x.e_0) \\ \widehat{\mathsf{C}} &\models_{\delta}^{ce} \quad (t_1^{\ell_1} t_2^{\ell_2})^{\ell} \text{ iff } \widehat{\mathsf{C}} \models_{\delta}^{ce} t_1^{\ell_1} \wedge \widehat{\mathsf{C}} \models_{\delta}^{ce} t_2^{\ell_2} \wedge \\ & \forall \langle (\lambda x.t_0^{\ell_0}), ce_0 \rangle \in \widehat{\mathsf{C}}(\ell_1, \delta) : \widehat{\mathsf{C}} \models_{\delta_0}^{ce_0'} t_0^{\ell_0} \wedge \\ & \widehat{\mathsf{C}}(\ell_2, \delta) \subseteq \widehat{\mathsf{C}}(x, \delta_0) \wedge \widehat{\mathsf{C}}(\ell_0, \delta_0) \subseteq \widehat{\mathsf{C}}(\ell, \delta) \\ & \text{where } \delta_0 = [\delta, \ell]_k \\ & \text{and } ce_0' = ce_0[x \mapsto \delta_0] \end{split}$$

The notation  $[\delta, \ell]_k$  denotes the string obtained by appending  $\ell$  to the end of  $\delta$  and taking the rightmost k labels.

Returning to the example given in section 3, the analysis is able to distinguish each occurrence of f as distinct. The increased precision allows us to conclude that the program is *approximated* by the (singleton) set  $\{\lambda y. y\}$ .

Let us consider an acceptable least analysis for the program in Figure 3. We write  $\epsilon$  for the empty contour and [] for the empty contour environment. Since every  $\lambda$ -term in the program is closed, the contour environments in the results will always be empty and so we omit it from this table:

$$\begin{array}{ll} \widehat{\mathsf{C}}(1,10) = \{\lambda x\} & \widehat{\mathsf{C}}(7,\epsilon) = \{\lambda f\} \\ \widehat{\mathsf{C}}(2,10) = \{\lambda x\} & \widehat{\mathsf{C}}(8,3) = \{\lambda x\} & \widehat{\mathsf{C}}(f,10) = \{\lambda x\} \\ \widehat{\mathsf{C}}(3,10) = \{\lambda x\} & \widehat{\mathsf{C}}(8,6) = \{\lambda y\} & \widehat{\mathsf{C}}(x,3) = \{\lambda x\} \\ \widehat{\mathsf{C}}(5,10) = \{\lambda y\} & \widehat{\mathsf{C}}(9,\epsilon) = \{\lambda x\} & \widehat{\mathsf{C}}(x,6) = \{\lambda y\} \\ \widehat{\mathsf{C}}(6,10) = \{\lambda y\} & \widehat{\mathsf{C}}(10,\epsilon) = \{\lambda y\} \end{array}$$

And the following holds:

$$\widehat{\mathsf{C}}\models^{[]}_{\epsilon}((\lambda f.((f^{1}f^{2})^{3}(\lambda y.y^{4})^{5})^{6})^{7}(\lambda x.x^{8})^{9})^{10}$$

#### 4.1 The *k*CFA decision problem

As we did in subsection 3.1, we now formulate a decision problem naturally answered by the analysis and ask: What is the difficulty of computing within this hierarchy? What are the sources of approximation that render such analysis tractable?

**Control Flow Problem** (*k***CFA**): Given an expression *e*, an abstract closure  $\langle (\lambda x.e_0), ce_0 \rangle$ , and a label-contour pair  $(\ell, \delta)$  with  $|\delta| \leq k$ , is  $\langle (\lambda x.e_0), ce_0 \rangle \in \widehat{C}(\ell, \delta)$  in a least analysis of *e*?

The source of approximation in *k*CFA is the bounding of the length of contour strings. But suppose k is sufficiently large that  $\delta$  is never truncated during the analysis. What can be said about the precision of the result? If the contour is never truncated, the analysis is just normalization. The acceptability relation above can be read as specifying a non-standard interpreter, which is given an expression and constructs a table from which the normalized program can be retrieved.

<sup>&</sup>lt;sup>3</sup> To be precise, we take as our starting point *uniform* kCFA rather than a kCFA in which  $\widehat{\mathbf{Cache}} = (\mathbf{Lab} \times \mathbf{CEnv}) \rightarrow \widehat{\mathbf{Val}}$ . The differences are immaterial for our purposes.

Let's rewrite the specification to make this clear. Evaluation is paramaterized by an initially empty table and inclusion constraints are interpreted as destructive updates to the table.  $\mathcal{E}[t^{\ell}]_{\delta}^{ce}$  evaluates *t* and writes the result into the table  $\widehat{C}$  at location  $(\ell, \delta)$ .

$$\begin{split} \mathcal{E}[\![x^{\ell}]\!]_{\delta}^{ce} &= \widehat{\mathbb{C}}(\ell,\delta) \leftarrow \widehat{\mathbb{C}}(x,ce(x)) \\ \mathcal{E}[\![(\lambda x.e_0)^{\ell}]\!]_{\delta}^{ce} &= \widehat{\mathbb{C}}(\ell,\delta) \leftarrow \langle (\lambda x.e_0),ce_0 \rangle \\ & \text{where } ce_0 = ce|\mathbf{fv}(\lambda x.e_0) \\ \mathcal{E}[\![(t_1^{\ell_1}t_2^{\ell_2})^{\ell}]\!]_{\delta}^{ce} &= \mathcal{E}[\![t_1^{\ell_1}]\!]_{\delta}^{ce}; \mathcal{E}[\![t_2^{\ell_2}]\!]_{\delta}^{ce}; \\ & \text{let } \langle (\lambda x.t_0^{\ell_0}),ce_0 \rangle = \widehat{\mathbb{C}}(\ell_1,\delta) \text{ in } \\ \widehat{\mathbb{C}}(x,\delta\ell) \leftarrow \widehat{\mathbb{C}}(\ell_2,\delta); \\ \mathcal{E}[\![t_0^{\ell_0}]\!]_{\delta\ell}^{ce} = \widehat{\mathbb{C}}(\ell_0,\delta\ell) \end{aligned}$$

The contour environment plays much the same role as an environment in a typical interpreter, but rather than mapping a variable to its value, it maps a variable to a location in the table where its value is found. The contour  $\delta$  is a history of call sites in place at the current point of evaluation and serves to keep locations in the table distinct (a simple induction proof shows that  $(\ell, \delta)$  is unique).

In the application case, the operator and operand are evaluated, updating the table at positions  $(\ell_1, \delta)$  and  $(\ell_2, \delta)$ , respectively. The closure in  $(\ell_1, \delta)$  is retrieved from the table and the variable is "bound" by writing the value of the argument (found at position  $(\ell_2, \delta)$ ) into position  $(x, \delta \ell)$ . The body of the closure is evaluated in an extended environment that maps x to the location in the table where its value is stored. After evaluating the body, the table is updated to record the value of the application as being the value found in  $(\ell_0, \delta \ell)$ .

This evaluation function can be seen as a variant of the exact collecting semantics from which the analysis was originally abstracted<sup>4</sup>, in other words, if k is big enough to that it is not truncated, the analysis is simply normalization.

## **Lemma 2.** If e has an exact kCFA, then $\mathcal{E}[\![e]\!]_{\epsilon}^{[]}$ constructs it.

A lower bound on the hardness of the analysis, then, is the expressivity of the language that can be evaluated in the above interpreter with a  $\delta$  of length at most k.

When the analysis is inexact, the evaluator must be modified to perform kCFA by truncating contours at length k. The relevant change to the evaluator is in the application case:

$$\begin{split} \mathcal{E}\llbracket (t_1^{\ell_1} t_2^{\ell_2})^{\ell} \rrbracket_{\delta}^{ce} &= \mathcal{E}\llbracket t_1^{\ell_1} \rrbracket_{\delta}^{ce}; \mathcal{E}\llbracket t_2^{\ell_2} \rrbracket_{\delta}^{ce}; \\ &\forall \langle \lambda x. t_0^{\ell_0}, ce_0 \rangle \in \widehat{\mathsf{C}}(\ell_1, \delta) : \\ &\widehat{\mathsf{C}}(x, \lceil \delta, \ell \rceil_k) \leftarrow \widehat{\mathsf{C}}(\ell_2, \delta); \\ &\mathcal{E}\llbracket t_0^{\ell_0} \rrbracket_{\lceil \delta, \ell \rceil_k}^{ce_0[x \mapsto \lceil \delta, \ell \rceil_k]}; \\ &\widehat{\mathsf{C}}(\ell, \delta) \leftarrow \widehat{\mathsf{C}}(\ell_0, \lceil \delta, \ell \rceil_k) \end{split}$$

Notice that truncation destroys the uniqueness of locations in the table, so in evaluating an application, any number of closures can flow into the operator position and *all* of them must be applied. Furthermore, the evaluator has to be iterated until a fixed point in the table  $\hat{C}$  is reached.

# **Lemma 3.** The least kCFA analysis of e is constructed by iterating $\mathcal{E}[\![e]\!]_{e}^{[]}$ until $\widehat{C}$ reaches a fixed point.

In this case, the table  $\widehat{C}$  is finite and has  $n^{k+1}$  entries. Each entry contains a set of *values* and the only values are closures

 $\langle (\lambda x.e_0), ce_0 \rangle$ ; the environment in a closure maps p free variables to any one of  $n^k$  contours. Because there are n possible  $\lambda x.e_0$  and  $n^{kp}$  such environments, there are sets of size at most  $n^{1+kp}$  in any table entry.

Observe that the above evaluation is *monotonic*: each table entries is initialzed to the empty set, and built up incrementally. Thus in *k*CFA, there can be at most  $n^{1+(k+1)p}$  updates to  $\widehat{C}$ , and  $\mathcal{E}[t^{\ell}]_{\delta}^{ce}$  then has at most  $n^{O(p)}$  program states during evaluation. Because  $p \leq n$ , we conclude with the well-known observation—see, for example, Nielson et al. (1999, page 193):

**Theorem 3.** *kCFA is contained in* EXPTIME.

#### 4.1.1 kCFA is NP-hard

Because CFA makes approximations, many closures can flow to a single program point and contour. In 1CFA, for example,  $\lambda w.wx_1x_2 \cdots x_n$  has *n* free variables, with an exponential number of possible associated environments mapping these variables to program points (contours of length 1). Approximation allows us to bind each  $x_i$ , independently, to either of the closed  $\lambda$ -terms for True or False that we saw in the PTIME-completeness proof for 0CFA. In turn, application to an *n*-ary Boolean function necessitates computation of all  $2^n$  such bindings in order to compute the flow out from the application site. The term True can only flow out if the Boolean function is satisfiable by some truth valuation.

$$\begin{aligned} &(\lambda f_1.(f_1 \operatorname{True})(f_1 \operatorname{False})) \\ &(\lambda x_1. \\ &(\lambda f_2.(f_2 \operatorname{True})(f_2 \operatorname{False})) \\ &(\lambda x_2. \\ &(\lambda f_3.(f_3 \operatorname{True})(f_3 \operatorname{False})) \\ &(\lambda x_3. \\ & \cdots \\ &(\lambda f_n.(f_n \operatorname{True})(f_n \operatorname{False})) \\ &(\lambda x_n. \\ &C[(\lambda v.\phi v)(\lambda w.w x_1 x_2 \cdots x_n)]) \cdots)))) \end{aligned}$$

For an appropriately chosen program point (label)  $\ell$ , the cache location  $\widehat{C}(v, \ell)$  will contain the set of all possible closures which are approximated to flow to v. This set is that of all closures

$$\langle (\lambda w.wx_1x_2\cdots x_n), ce \rangle$$

where *ce* ranges over all assignments of True and False to the free variables (or more precisely assignments of locations in the table containing True and False to the free variables). The Boolean function  $\phi$  is completely linear, as in the PTIME-completeness proof; the context *C* uses the Boolean output(s) as in the conclusion to that proof: mixing in some ML, the context is:

Again, a can only flow as an argument to f if True flows to (u, u'), leaving (f,g) unchanged, which can only happen if *some* closure  $\langle (\lambda w.wx_1x_2\cdots x_n), ce \rangle$  provides a satisfying truth valuation for  $\phi$ . We have as a consequence:

#### Theorem 4. 1CFA is NP-hard.

We observe that while the computation of the *entire* cache  $\widehat{C}$  requires exponential time, the existence of a *specific* flow in it may well be computable in NP. A nondeterministic polynomial might compute using the "collection semantics"  $\mathcal{E}[t^{\ell}]_{\delta}^{ce}$ , but rather than compute entire sets, *choose* the element of the set that bears witness to the flow.

<sup>&</sup>lt;sup>4</sup> However it varies in flavor—being a big step semantics rather than the structured operational semantics of Nielson and Nielson (1997). The motivation for SOS in Nielson and Nielson (1997) was to prove correctness of the analysis for non-terminating programs. Our evaluator only works for finite programs, but since we are investigating the complexity of the analysis, this is agreeable.

Increasing the precision to kCFA with k > 1 undermines the approximation that allows for construction of the NP-hardness proof. We use the following program transformation to render a kCFA of the original synonymous with a (k + 1)CFA of the transformed program. The NP-hardness of kCFA for k > 0 falls out by iterating the transformation k - 1 times on the 1CFA construction.

Transform expressions as follows, where  $\ell^*$  is a distinguished label and k a distinguished variable not appearing in the source expression:

$$\begin{aligned} \langle x^{\ell} \rangle &= x^{\ell} \\ \langle (e_1 e_2)^{\ell} \rangle &= (\langle e_1 \rangle \langle e_2 \rangle)^{\ell} \\ \langle (\lambda x. e)^{\ell} \rangle &= ((\lambda k. (\lambda x. \langle e \rangle)^{\ell^*}) (\lambda y. y)) \end{aligned}$$

The transformation works by nesting every  $\lambda$ -abstraction in a single application context (a *K*-redex), which consumes the added precision of a (k + 1)CFA. The following lemma states that top-level flows are analogous in the transformed program under a more precise analysis:

**Lemma 4.** If  $\widehat{C} \models_{\epsilon}^{[]} t^{\ell}$  is a least kCFA, and  $\widehat{C}' \models_{\epsilon}^{[]} \langle t^{\ell} \rangle$  is a least (k+1)CFA, then  $\langle (\lambda x.e), ce \rangle \in \widehat{C}(\ell)$  iff  $\langle (\lambda x.\langle e \rangle), ce' \rangle \in \widehat{C}'(\ell)$ , where ce and ce' are isomorphic upto placement of  $\star$ -labels.

For any fixed *k*, this constant-factor expansion can be computed by a logspace-computable reduction. From this, we conclude:

**Theorem 5.** *kCFA is* NP-*hard, for any* k > 0.

#### **4.1.2** *n***CFA is complete for** EXPTIME

We now examine the complexity of kCFA where k is allowed to vary linearly in the size of the program.

In the previous section, we saw 0CFA was sufficient to evaluate linear  $\lambda$ -terms. Following the construction of Neergaard and Mairson (2004), we can code a Turing machine transition function and machine IDs with linear terms. Suppose  $\phi$  is a linear transition function that takes a tuple  $\langle q, L, R \rangle$  consisting of the current machine state, the tape to the left of the head in reverse order, and the tape to the right of the head. Applying  $\phi$  returns a tuple  $\langle q', L', R' \rangle$ consisting of the new state and tape. Let I be the initial configuration, then  $\phi I$  simulates one step of the machine,  $\phi(\phi I)$  two steps, and  $\phi^n I$ , n steps.

If 0CFA is sufficient to evaluate  $\phi I$ , then what is 1CFA sufficient for? By introducing non-linearity using the Church numeral  $\overline{2}$ , we can iterate the transition function twice, as follows:

$$\overline{2}\phi I \equiv (\lambda s.\lambda z.(s(sz)^1)^2)\phi I$$

A contour of length 1 is sufficient to distinguish between the application of  $\phi$  in the calling context of 4 and that of 5, we are able to maintain an exact analysis. Scaling up, suppose we have:

$$\overline{2}(\overline{2}\phi)I \equiv (\lambda s.\lambda z.(s(sz)^3)^4)((\lambda s.\lambda z.(s(sz)^1)^2)\phi)I$$

A contour of length 2 is sufficient to distinguish between the application of  $\phi$  in the calling context 13, 14, 23 and 24. In general, *n*CFA is sufficient for an exact analysis of  $\overline{2}^n \phi I$ —all of the calling contexts in which  $\phi$  is applied are distinguished. Thus *n*CFA is synonymous with normalization and the program normalizes to  $\phi^{2^n} I$ . So when *k* is linear in the size of the program *n*, we can simulate a Turing machine for an exponential number of steps.

#### **Theorem 6.** *nCFA is complete for* EXPTIME.

It should be remarked that in this case, the contours are large enough that the computation is essentially by *normalization*, without using the power of any approximation. Every location of the cache  $\hat{C}$  contains at most one value. Researchers have noted that computing a more precise analysis is often cheaper than performing a less precise one. A less precise analysis "yields coarser approximations, and thus induces more merging. More merging leads to more propagation, which in turn leads to more reevaluation" (Wright and Jagannathan 1998). Harnessing the computational power of this reevaluation is precisely what makes the NP-hardness construction work and relegates lower bounds using exact analyses to limit cases such as k = n; an analysis that is exact can only be polynomial in  $n^k$ .

On the other hand, these limiting cases shed analytic light on the nature of kCFA. Even when the polyvariance of the analysis is taken to an extreme as in k = n, the expressivity of the analysis is still limited to EXPTIME. The intractible feature of kCFA is the approximation of closures, not the degree of polyvariance. Polyvariant static analyses such as polymorphic splitting and poly-kCFA have since developed without the use of closures and enjoyed tractability.

### **5.** LOGSPACE and $\eta$ -expansion

In this section, we identify a restricted class of functional programs whose 0CFA decision problem may be simpler—namely, complete for LOGSPACE. Consider programs that are simply typed, and where a variable in the function position or the argument position of an application is fully  $\eta$ -expanded. This case—especially, but not only when the programs are linear—strongly resembles multiplicative linear logic with *atomic* axioms. This distinction is highlighted in the discussion below.

We remark that  $\eta$ -expansion changes control flow analysis. If 0CFA infers that a call site  $\chi$  may call a function  $\phi$  in a program  $\Pi$ , and we ask the same question of the residual  $\chi$  and  $\phi$  in the  $\eta$ -expanded version of  $\Pi$ , the answer may vary.

#### 5.1 MLL and (linear) functional programming

The sequent rules of MLL are:

$$\operatorname{Ax} \frac{}{A,A^{\perp}} \quad \operatorname{Cut} \frac{\Gamma, A \quad A^{\perp}, \Delta}{\Gamma, \Delta} \quad \otimes \frac{\Gamma, A, B}{\Gamma, A \otimes B} \quad \otimes \frac{\Gamma, A \quad \Delta, B}{\Gamma, \Delta, A \otimes B}$$

These rules have an easy functional programming interpretation as the types of a linear programming language (eg. linear ML), following the intuitions of the Curry-Howard correspondence (Girard et al. 1989, Chapter 3).<sup>5</sup>

The AXIOM rule says that a variable can be viewed simultaneously as a continuation  $(A^{\perp})$  or as an expression (A)—one man's ceiling is another man's floor. Thus we say "input of type A" and "output of type  $A^{\perp}$ " interchangably, along with similar dualisms. We also regard  $(A^{\perp})^{\perp}$  synonymous with A: for example, Int is an integer, and Int<sup> $\perp$ </sup> is a request (need) for an integer, and if you need to need an integer—(Int<sup> $\perp$ </sup>)<sup> $\perp$ </sup>—then you have an integer.

The CUT rule says that if you have two computations, one with an output of type A, another with an input of type A, you can plug them together.

The  $\otimes$ -rule is about pairing: it says that if you have separate computations producing outputs of types A and B respectively, you can combine the computations to produce a paired output of type  $A \otimes B$ . Alternatively, given two computations with A an output in one, and B an input (equivalently, continuation  $B^{\perp}$  an output) in the other, they get paired as a *call site* "waiting" for a function which produces an *output* of type B with an *input* of type A. Thus  $\otimes$  is both cons and function call (@).

The  $\mathcal{B}$ -rule is the linear unpairing of this  $\otimes$ -formation. When a computation uses inputs of types A and B, these can be combined

<sup>&</sup>lt;sup>5</sup> For a more detailed discussion of the correspondence between linear ML and MLL, see Mairson (2004).



Figure 4. Expansion algorithm.

as a single input pair, e.g., let (x,y)=p in... Alternatively, when a computation has an input of type A (output of continuation of type  $A^{\perp}$ ) and an output of type B, these can be combined to construct a function which inputs a call site pair, and unpairs them appropriately. Thus  $\mathfrak{B}$  is both unpairing and  $\lambda$ .

## 5.2 Atomic versus non-atomic axioms: PTIME versus LOGSPACE

The above AXIOM rule does not make clear whether the formula A is an atomic type variable or a more complex type formula. When a *linear* program only has atomic formulas in the "axiom" position, then we can evaluate (normalize) it in logarithmic space. When the program is not linear, we can similarly compute a 0CFA analysis in LOGSPACE. Moreover, these problems are complete for LOGSPACE.

MLL proofs with non-atomic axioms can be easily converted to ones with atomic axioms using the following transformation, analogous to  $\eta$ -expansion:

$$\frac{\overline{\alpha, \alpha^{\perp}} \quad \overline{\beta, \beta^{\perp}}}{\alpha \otimes \beta, \alpha^{\perp} \otimes \beta^{\perp}} \qquad \Rightarrow \qquad \frac{\overline{\alpha, \alpha^{\perp}} \quad \overline{\beta, \beta^{\perp}}}{\alpha \otimes \beta, \alpha^{\perp} \otimes \beta^{\perp}}$$

This transformation can increase the size of the proof. For example, in the circuit examples of the previous section (which are evidence for PTIME-completeness),  $\eta$ -expansion causes an exponential increase in the number of proof rules used.<sup>6</sup> A LOGSPACE evaluation is then polynomial-time and -space in the original circuit description.

The program transformation corresponding to the above proof expansion is a version of  $\eta$ -expansion: see Figure 4. The left hand expansion rule is simply  $\eta$ , dualized in the unusual right hand rule. The right rule is written with the @ above the  $\lambda$  only to emphasis its duality with the left rule. Although not shown in the graphs, but implied by the term rewriting rules, an axiom may pass through any number of sharing nodes.

## 5.3 Normalization and 0CFA for linear programs in LOGSPACE

A normalized *linear* program has no redexes. From the type of the program, one can reconstruct—in a totally syntax-directed way—what the structure of the term is. It is only the position of the *axioms* that is not revealed. For example, both TT and FF from the above circuit example have type 'a \* 'a -> 'a \* 'a.<sup>7</sup> From this type, we can see that the term is a  $\lambda$ -abstraction, the parameter is unpaired—and then, are the two components of type a repaired as before, or "twisted"? To twist or not to twist is what distinguishes TT from FF.

The geometry of interaction (GoI)—the semantics of linear logic—and the notion of paths provide a way to calculate normal forms, and may be viewed as the logician's way of talking about static program analysis.<sup>8</sup> To understand how this analysis works, we need to have a graphical picture of what a linear functional program looks like.

Without loss of generality, such a program has a type  $\phi$ . Nodes in its graphical picture are either  $\lambda$  or linear unpairing ( $\otimes$  in MLL), or application/call site or linear pairing ( $\otimes$  in MLL). We draw the graphical picture so that axioms are on top, and cuts (redexes, either  $\beta$ -redexes or pair-unpair redexes) are on the bottom.



Because the axioms all have atomic type, the graph has the following nice property:

**Lemma 5.** Begin at an axiom  $\alpha$  and "descend" to a cut-link, saving in an (initially empty) stack whether nodes are encountered on their left or right auxiliary port. Once a cut is reached, "ascend" the accompanying structure, popping the stack and continuing left or right as specified by the stack token. Then (1) the stack empties exactly when the next axiom  $\alpha'$  is reached, and (2) if the k-th node from the start traversed is a  $\otimes$ , the k-th node from the end traversed is a  $\otimes$ , and vice versa.

The path traced in the Lemma, using the stack, is geometry of interaction (GoI), also known as static analysis. The correspondence between the *k*-th node from the start and end of the traversal is precisely that between a *call site* ( $\otimes$ ) and a *called function* ( $\otimes$ ), or between a cons ( $\otimes$ ) and a linear unpairing ( $\otimes$ ).

A sketch of the "four finger" normalization algorithm: The stack height may be polynomial, but we do not need the stack! Put fingers  $\alpha$ ,  $\beta$  on the axiom where the path begins, and iterate over all possible choices of another two fingers  $\alpha'$ ,  $\beta'$  at another axiom. Now move  $\beta$  and  $\beta'$  towards the cut link, where if  $\beta$  encounters a node on the left (right), then  $\beta'$  must move left (right) also. If  $\alpha'$ ,  $\beta'$  were correctly placed initially, then when  $\beta$  arrives at the cut link, it must be met by  $\beta'$ . If  $\beta'$  isn't there, or got stuck somehow, then  $\alpha'$ ,  $\beta'$  were incorrectly placed, and we iterate to another placement and try again.

**Lemma 6.** Any path from axiom  $\alpha$  to axiom  $\alpha'$  traced by the stack algorithm of the previous lemma is also traversed by the "four finger" normalization algorithm.

<sup>&</sup>lt;sup>6</sup> It is linear in the formulas used, whose length increases exponentially (not so if the formulas are represented by directed acyclic graphs).

<sup>&</sup>lt;sup>7</sup> The linear logic equivalent is  $(\alpha^{\perp} \otimes \alpha^{\perp}) \otimes (\alpha \otimes \alpha)$ . The  $\lambda$  is represented by the outer  $\otimes$ , the unpairing by the inner  $\otimes$ , and the consing by the  $\otimes$ .

<sup>&</sup>lt;sup>8</sup> See Mairson (2002) for an introduction to context semantics and normalization by static analysis in the geometry of interaction.

Normalization by static analysis is synonymous with traversing these paths. Because these fingers can be stored in logarithmic space, we conclude (Terui 2002; Mairson 2006):

## **Theorem 7.** Normalization of linear, simply-typed, and fully $\eta$ -expanded functional programs is contained in LOGSPACE.

That 0CFA is then contained in LOGSPACE is a casual byproduct of this theorem, due to the following observation: if application site  $\chi$  calls function  $\phi$ , then the  $\otimes$  and  $\otimes$  (synonymously, @ and  $\lambda$ ) denoting call site and function are in distinct trees connected by a CUT link. As a consequence the 0CFA computation is a subcase of the four-finger algorithm: traverse the two paths from the nodes to the cut link, checking that the paths are isomorphic, as described above. The full 0CFA calculation then iterates over all such pairs of nodes.

**Corollary 1.** *OCFA of linear, simply-typed, and fully*  $\eta$ *-expanded functional programs is contained in* LOGSPACE.

#### 5.4 OCFA in LOGSPACE

Now let us remove the linearity constraint, while continuing to insist on full  $\eta$ -expansion as described above, and simple typing. The normalization problem is no longer contained in LOGSPACE, but rather nonelementary recursive, (Statman 1979; Mairson 1992; Asperti and Mairson 1998). However, 0CFA remains contained in LOGSPACE, because it is now an *approximation*. This result follows from the following observation:

**Lemma 7.** Suppose  $(\lambda x.e_0)^{l'}$  and  $(t_1^l e_2)$  occur in a simply typed, fully  $\eta$ -expanded program and  $\lambda x.e_0 \in \widehat{C}(l)$ . Then the corresponding  $\otimes$  and  $\otimes$  occur in adjacent trees connected at their roots by a CUT-link and on dual, isomorphic paths modulo placement of sharing nodes.

Here "modulo placement" means: follow the paths to the cut then we encounter  $\otimes$  (resp.,  $\otimes$ ) on one path when we encounter  $\otimes$  (resp.,  $\otimes$ ) on the other, on the same (left, right) auxiliary ports. We thus *ignore* traversal of sharing nodes on each path in judging whether the paths are isomorphic. (Without sharing nodes, the  $\otimes$ and  $\otimes$  would annihilate—i.e., a  $\beta$ -redex—during normalization.)

**Theorem 8.** *OCFA of a simply-typed, fully*  $\eta$ *-expanded program is contained in* LOGSPACE.

Observe that 0CFA defines an *approximate* form of normalization which is suggested by simply *ignoring* where sharing occurs. Thus we may define the *set* of  $\lambda$ -terms to which that a term might evaluate. Call this *0CFA-normalization*.

**Theorem 9.** For fully  $\eta$ -expanded, simply-typed terms, OCFAnormalization can be computed in nondeterministic LOGSPACE.

**Conjecture 1.** For fully  $\eta$ -expanded, simply-typed terms, OCFAnormalization is complete for nondeterministic LOGSPACE.

The proof of the above conjecture likely depends on a coding of arbitrary directed graphs and the consideration of commensurate path problems.

**Conjecture 2.** An algorithm for OCFA normalization can be realized by optimal reduction, where sharing nodes always duplicate, and never annihilate.

## 5.5 LOGSPACE-hardness of normalization and 0CFA: linear, simply-typed, fully η-expanded programs

That the normalization and 0CFA problem for this class of programs is as hard as any LOGSPACE problem follows from the LOGSPACE-hardness of the *permutation problem*: given a permutation  $\pi$  on  $1, \ldots, n$  and integer  $1 \le i \le n$ , are 1 and *i* on the same cycle in  $\pi$ ? That is, is there a *k* where  $1 \le k \le n$  and  $\pi^k(1) = i$ ? Briefly, the LOGSPACE-hardness of the permutation problem is as follows. Given an arbitrary LOGSPACE Turing machine M and an input x to it, visualize a graph where the nodes are machine IDs, with directed edges connecting successive configurations. Assume that M always accepts or rejects in unique configurations. Then the graph has two connected components: the "accept" component, and the "reject" component. Each component is a directed tree with edges pointing towards the root (final configuration). Take an Euler tour around each component (like tracing the fingers on your hand) to derive two *cycles*, and thus a *permutation* on machine IDs. Each cycle is polynomial size, because the configurations only take logarithmic space. The equivalent permutation problem is then: does the initial configuration and the accept configuration sit on the same cycle?

The following linear ML code describes the "target" code of a transformation of an instance of the permutation problem. For a permutation on n letters, we take here an example where n = 3. Begin with a vector of length n set to False, and a permutation on n letters:

- val V= (False,False,False); val V = ((fn,fn),(fn,fn),(fn,fn)) : (('a \* 'a -> 'a \* 'a) \* ('a \* 'a -> 'a \* 'a)) \* (('a \* 'a -> 'a \* 'a) \* ('a \* 'a -> 'a \* 'a)) \* (('a \* 'a -> 'a \* 'a) \* ('a \* 'a -> 'a \* 'a))

Denote as  $\nu$  the type of vector V.

- fun Perm (P,Q,R)= (Q,R,P); val Perm = fn :  $\nu \rightarrow \nu$ 

The function Insert *linearly* inserts True in the first vector component, using all input exactly once:

```
- fun Insert ((p,p'),Q,R)= ((TT,Compose(p,p')),Q,R); val Insert = fn : \nu -> \nu
```

The function Select *linearly* selects the third vector component:

```
- fun Select (P,Q,(r,r'))=
    (Compose (r,Compose (Compose P, Compose Q)),r');
val Select = fn
    : ν -> (('a * 'a -> 'a * 'a) * ('a * 'a -> 'a * 'a))
```

Because Perm and Insert have the same flat type, they can be composed iteratively in ML without changing the type. (This clearly is *not* true in our coding of circuits, where the size of the type increases with the circuit. A careful coding limits the type size to be polynomial in the circuit size, regardless of circuit depth.)

**Lemma 8.** Let  $\pi$  be coded as permutation Perm. Define Foo to be Compose(Insert,Perm) composed with itself n times. Then 1 and i are on the same cycle of  $\pi$  iff Select (Foo V) normalizes to True.

Because 0CFA of a linear program is identical with normalization, we conclude:

**Theorem 10.** *OCFA of a simply-typed, fully*  $\eta$ *-expanded program is complete for* LOGSPACE.

As a consequence of the hardness construction having a constant type, we may conclude the 0CFA of any bounded type program ( $\eta$ -expanded or not) is LOGSPACE-hard:

**Theorem 11.** *OCFA of a bounded, simply-typed program is* LOGSPACE-*hard.* 

### 6. Languages with first-class control

Shivers (2004) argues that "CPS provide[s] a uniform representation of control structure," allowing "this machinery to be employed to reason about context, as well," and that "without CPS, seperate



Figure 5. Graph coding of call/cc and example CFA graph.

contextual analyses and transforms must be also implemented redundantly," in his view. Although our formulation of kCFA is a "direct-style" formulation, a graph representation enjoys the same benefits of a CPS representation, namely that control structures are made explicit—in a graph a continuation is simply a wire. Control constructs such as call/cc can be expressed directly (Lawall and Mairson 2000) and our graphical formulation of control flow analysis carries over without modification.

Lawall and Mairson (2000) derive graph representations of programs with control operators such as call/cc by first translating programs into continuation passing style (CPS). They observed that when edges in the CPS graphs carrying answer values (of type  $\perp$ ) are eliminated, the original (direct-style) graph is regained, modulo placement of boxes and croissants that control sharing. Composing the two transformations results in a direct-style graph coding for languages with call/cc (hereafter,  $\lambda_{\mathcal{K}}$ ). The approach applies equally well to languages such as Filinski's symmetric  $\lambda$ -calculus (1989), Parigot's  $\lambda_{\mu}$  calculus (1992), and most any language expressible in CPS.<sup>9</sup>

The left side of Figure 5 shows the graph coding of call/cc. Examining this graph, we can read of an interpretation of call/cc, namely: call/cc is a function that when applied, copies the current continuation ( $\triangle$ ) and applies the given function f to a function ( $\lambda v \dots$ ) that when applied abandons the continuation at that point ( $\odot$ ) and gives its argument v to a copy of the continuation where call/cc was applied. If f never applies the function it is given, then control returns "normally" and the value f returns is given to the other copy of the continuation where call/cc was applied.

The right side of Figure 5 gives the CFA graph for the program:

$$(\operatorname{call/cc} (\lambda k.(\lambda x.\overline{1})(k\overline{2})))^{l}$$

From the CFA graph we see that  $\widehat{C}(l) = \{\overline{1}, \overline{2}\}$ , reflecting the fact that the program will return  $\overline{1}$  under a call-by-name reduction strategy and  $\overline{2}$  under call-by-value. Thus, the analysis is indifferent to the reduction strategy. Note that whereas before, approximation was introduced through non-linearity of bound variables, approximation can now be introduced via non-linear use of continuations, as seen in the example. In the same way that 0CFA considers all occurrences of a bound variable "the same", 0CFA considers all continuations obtained with each instance of call/cc "the same".

Note that we can ask new kinds of interesting questions in this analysis. For example, in Figure 5, we can compute which continuations are potentially *discarded*, by computing which continuations flow into the weakening node of the call/cc term. (The answer is the continuation  $((\lambda x.\overline{1})[])$ .) Likewise, it is possible to ask

which continuations are potentially *copied*, by computing which continuations flow into the principal port of the sharing node in the call/cc term (in this case, the top-level empty continuation []). Because continuations are used linearly in call/cc-free programs, the questions were uninteresting before—the answer is always *none*.

Our proofs for the PTIME-completeness of 0CFA for the untyped  $\lambda$ -calculus—and likewise for the results on kCFA—carry over without modification languages such as  $\lambda_{\mathcal{K}}$ ,  $\lambda_{\mu}$  and the symmetric  $\lambda$ -calculus. In other words, first-class control operators such as call/cc increase the expressivity of the language, but add nothing to the computational complexity of control flow analysis. In the case of simply-typed, fully  $\eta$ -expanded programs, the same can be said. A suitable notion of "simply-typed" programs is needed, such as that provided by Griffin (1990) for  $\lambda_{\mathcal{K}}$ . The type-based expansion algorithm of Figure 4 applies without modification and Lemma 7 holds, allowing 0CFA for this class of programs to be done in LOGSPACE. Linear logic provides a foundation for (classical)  $\lambda$ -calculi with control; related logical insights allow control flow analysis in this setting.

#### 7. Related work

The PTIME-completeness of 0CFA is most closely related to the PTIME-completeness of simply typing in the  $\lambda$ -caclculus (Mairson 2004). Both results use linearity to subvert the approximation of the analysis, and since both analyses rely on the same source of approximation, it is no suprise that they share the same lower bound on complexity.

ML typing can be viewed as a bounded running of a program (reducing all let-redexes) followed by a simple typing of the residual. The residual program can be exponentially larger, leading to EXPTIME-completeness results by using polymorphism to iterate a linear TM transition function (Mairson 1990). The EXPTIME-completeness of nCFA can be viewed in a similar light. Contours of length proportional to the program size provide a bounded "running" of the program by exact analysis of the non-linearity introduced by iterative doubling of the transition function.

The story is the same for k-rank bounded intesection typing—a program is run by computing k successive minimal complete developments and the residual is simply typed. The resulting hardness of typing is elementary in k (Neergaard and Mairson 2004), and thus the complexity class of each fixed k is seperated. On the other hand, for (k > 0)CFA, the complexity in the "hierarchy" remains the same as k grows. There should be a natural way of developing an alternative control flow hierarchy that relies on complete developments for its notion of bounded running that will be strictly more expressive than the kCFA examined in this paper. The result is likely to be similiar in spirit to that of Mossin (1997b), although Mossin's analysis is simply evaluation by virtue of its exactness. To remain useful, some information must be purposeless lost in order to compute an answer in less time than it takes to run the program.

It also seems likely that the linear logic based investigation into CFA presented here can be coupled with that of Neergaard and Mairson (2004) to provide the foundation for complexity results for the control flow analysis of rank-2 bounded intersection typed programs (Banerjee and Jensen 2003).

Static program analysis has been recast as various kinds of graph reachability problems, and parenthesis languages have been used to describe paths in these graphs; see Reps (2000) for example. Words in these languages are the *contexts* of the context semantics presentation (Mairson 2003) of the geometry of interaction (Girard 1989). The undecidability of decision problems for these specialized parenthesis languages corresponds naturally to versions of the halting problem.

<sup>&</sup>lt;sup>9</sup> Languages such as  $\lambda_{\xi}$ , which contains the "delimited control" operators *shift* and *reset* (Danvy and Filinski 1990), are not immediately amenable to this approach since the direct-style transformation requires all calls to functions or continuations be in tail position. Adapting this approach to such languages constitutes an active area of research for us.

The graph coding of terms in our development is based on the technology of *sharing graphs* in the untyped case, and *proof nets* in the typed case (Lafont 1995). The graph codings, CFA graphs, and virtual wire propogation rules share a strong resemblance to the "pre-flow" graphs, flow graphs, and graph "closing rules", respectively, of Mossin (1997a). Casting the analysis in this light leads to insights from linear logic and optimal reduction. For example, as Mossin (1997a, page 78) notes, the CFA virtual paths computed by 0CFA are an approximation of the actual runtime paths and correspond exactly to the "well-balanced paths" of Asperti and Laneve (1995) as an approximation to "legal paths" (Lévy 1978) and results on proof normalization in linear logic (Mairson and Terui 2003) informed the novel CFA algorithms presented here.

Other research has shown a correspondence between 0CFA and certain type systems (Palsberg and O'Keefe 1995; Heintze 1995) and a further connection has been made between intersection typing and kCFA (Mossin 1997a; Palsberg and Pavlopoulou 1998). Work has also been done on relating the various flavors of control flow analysis, such as 0CFA, kCFA, polymorphic splitting, and uniform kCFA (Nielson and Nielson 1997). Moreover, control flow analysis can be computed under a number of different guises such as set-based analysis (Heintze 1994), closure analysis (Sestoft 1988, 1989), abstract interpretation (Shivers 1991; Tang and Jouvelot 1994), and type and effect systems (Faxén 1995; Heintze 1995; Faxén 1997; Banerjee 1997).

We believe a useful taxonomy of these and other static analyses can be derived by investigating their computational complexity. In the preface to their textbook, Nielson et al. (1999) make an analogy between their approach to the study of program analysis and that of complexity theory, whereby seemingly unrelated problems are shown to be the same through notions such as logspace reduction. We believe there is more than an analogy here—results on the complexity of static analyses are a useful way of understanding when two seemingly different program analyses are in fact computing the same thing.

### 8. Conclusions and perspective

The most obvious thing you can say about program analysis is this: if you want to know what a program is going to do, run it and find out. If you don't have time to run the program, run it for a while. If you don't have time to run it for a while, make a crude approximation. If you want to make sure you have all the answers in such an approximation, you'll end up computing "false positives". A common approximation idea is to merge whatever is known for all occurrences of a variable. Approximations are refined by gently distinguishing these variable occurrences.

Type inference follows this blueprint. Simple typing is the approximation: all occurrences of a variable have to have the same type. ML let-polymorphism computes a *finite development* before simple typing. Rank-bounded intersection systems iterate several minimal complete developments before simple typing (with  $\wedge$ -idempotency) or a linearity check (without  $\wedge$ -idempotency): see (Neergaard and Mairson 2004).

This is also the story of OCFA: a truncated evaluation that merges all occurrences of a bound variable. The occurrences are then distinguished in kCFA (k > 0) by *contours* that annotate application sites, and an *environment* whose values are closures—and it is the free variables of these closures that are the bottleneck of kCFA. The NP-hardness lower bound, and likely NP-completeness given the known exponential upper bound, comes from computing with approximations rather than computing via exact normalization.

What other kind of static analysis could there possibly be? We would like to relate kCFA in a more precise way to the geometry of interaction, a complete static program analysis based on linear

logic. It should be possible to construct an exact correspondence when k is unbounded. A further extension would be a generalization of kCFA to the precision of iterated exponentials, combining its technique with finite developments à la ML. Finally, we hope to use the techniques described here to provide a useful taxonomy of other program analyses, viewed from the perspective of the computational resources needed to realize them.

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