

Cactus Environment Machine

Shared Environment Call-by-Need

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Abstract. Existing machines for lazy evaluation use a *flat* representation of environments, storing the terms associated with free variables in an array. Combined with a heap, this structure supports the shared intermediate results required by lazy evaluation. We propose and describe an alternative approach that uses a *shared* environment to minimize the overhead of delayed computations. We show how a shared environment can act as both an environment and a mechanism for sharing results. To formalize this approach, we introduce a calculus that makes the shared environment explicit, as well as a machine to implement the calculus, the *Cactus Environment Machine*. A simple compiler implements the machine and is used to run experiments for assessing performance. The results show reasonable performance and suggest that incorporating this approach into real world compilers could give performance benefits in some scenarios.

1 Introduction

Call-by-need evaluation is a formalization of the idea that work should be delayed until needed, and performed only once. Existing implementations of call-by-need take care in *packaging* a delayed computation, or *thunk*, by building a closure with an array that contains the bindings of all free variables [24, 7]. The overhead induced by this operation is well known, and one reason existing implementations avoid thunks wherever possible [18]. The key insight of our Cactus Environment (\mathcal{CE}) Machine is that this overhead can be minimized by only recording a location in a shared environment.

As an example, consider the application $f e$. In existing call-by-need implementations, e.g., the STG machine[24], a closure with a flat environment will be constructed for e . Doing so incurs a time and memory cost proportional to the number of free variables of e .³ We minimize this packaging cost by recording a location in a shared environment, which requires only two machine words (and two instructions) for the thunk: one for the code pointer, and one for the environment pointer. One way to think about the approach is that it is *lazier* about lazy evaluation: in the case that e is unneeded, the work to package it in a thunk is entirely wasted. In the spirit of lazy evaluation, we attempt to minimize this potentially unnecessary work.

The main contributions of the paper are:

³ In some implementations, these are lambda lifted to be formal parameters, but the principle is the same.

- A big-step calculus and small-step abstract machine that formalize the notion of a shared environment for call-by-need evaluation using an explicitly shared environment (Section 4).
- A simple implementation of the abstract machine that compiles to x86 assembly with a preliminary evaluation that shows performance comparable to existing implementations (Sections 6 and 7).

Section 2 reviews relevant background material, and Section 3 discusses the current landscape of environment representations, highlighting the opportunity for combining shared environments with lazy evaluation. We then provide some intuition for why this combination is desirable, and formalize the connection between call-by-need evaluation and shared environments in a calculus (Section 4). Section 5 uses the calculus to derive a novel abstract machine, the \mathcal{CE} machine, explains how \mathcal{CE} uses the shared environment in a natural way to implement lazy evaluation, and gives its formal semantics. We then describe a straightforward implementation of \mathcal{CE} in Section 6, extended with machine literals and primitive operations, compiling directly to native code. We evaluate the implementation in Section 7, showing that it is capable of performing comparably to existing implementations, despite lacking several common optimizations, and discuss the results. We discuss related work, the limitations of our approach, and some ideas for future work in Section 8, and conclude the paper in Section 9.

2 Background and Motivation

This section provides relevant background for the \mathcal{CE} machine, outlining lambda calculus, evaluation strategies, and Curien’s calculus of closures.

2.1 Preliminaries

We begin with the simple lambda calculus [5]:

$$t ::= x \mid \lambda x.t \mid t t$$

where x is a variable, $\lambda x.t$ is an abstraction, and $t t$ is an application. We also use lambda calculus with deBruijn indices, which replace variables with a natural number indexing into the binding lambdas. This calculus is given by the syntax:

$$t ::= i \mid \lambda t \mid t t$$

where $i \in \mathbb{N}$. In both cases, we use the standard Barendregt syntax conventions, namely that applications are left associative and the bodies of abstractions extend as far as possible to the right [5]. A *value* in lambda calculus refers to an abstraction. We are concerned only with evaluation to weak head normal form (WHNF), which terminates on an abstraction without entering its body.

In mechanical evaluation of expressions, it would be too inefficient to perform explicit substitution. To solve this, the standard approach uses closures [20, 8, 24, 6]. Closures combine a term with an environment, which binds the free variables in the term to closures.

For a formal basis for closures, we use Curien’s calculus of closures [8], given in Figure 2. It is a formalization of closures with an environment represented as a list of closures, indexed by deBruijn indices. We will occasionally modify this calculus by replacing the deBruijn indices with variables for readability, in which case variables are looked up in the environment instead of indexed, e.g., $t[x = c, y = c']$ [5]. We also add superscript and subscript markers to denote unique syntax elements, e.g., $t', t_1 \in \text{Term}$.

2.2 Evaluation Strategies

There are three standard evaluation strategies for lambda calculus: call-by-value, call-by-need, and call-by-name. Call-by-value evaluates every argument to a value, whereas call-by-need and call-by-name only evaluate an argument if it is needed. If an argument is needed more than once, call-by-name re-computes the value, where call-by-need memoizes the value, so it is computed at most once. Thus, call-by-need attempts to embody the best of both worlds—never repeat work (call-by-value), and never perform unnecessary work (call-by-name). These are intuitively *good* properties to have, and we illustrate the correctness of such an intuition with the following example, modified from [10]:

$$\overbrace{c_m(c_m(\cdots(c_m \text{ id } \text{ id})\cdots)\text{ id})}^m \text{ true id bottom}$$

where $c_n = \lambda s. \lambda z. s (\overbrace{s \cdots (s z)}^n) \cdots$, $\text{true} = \lambda t. \lambda f. t$, $\text{id} = \lambda x. x$, and $\text{bottom} = (\lambda x. x x) \lambda x. x x$. Call-by-value never terminates, call-by-name takes exponential time, and call-by-need takes only polynomial time [10]. Of course, this is a contrived example, but it illustrates desirable properties of call-by-need.

In practice, however there are significant issues with call-by-need evaluation. We focus on the following: *Delaying a computation is slower than performing it immediately*. This issue is well known [18, 24], and has become part of the motivation for *strictness analysis* [23, 32], which transforms non-strict evaluation to strict when possible.

2.3 Existing Call-by-Need Machines

Diehl et al. [11] review the call-by-need literature in detail. Here we summarize the most relevant points.

The best known machine for lazy evaluation is the Spineless Tagless G-Machine (STG machine), which underlies the Glasgow Haskell Compiler (GHC). STG uses flat environments that can be allocated on the stack, the heap, or some combination [24].

Two other influential lazy evaluation machines relevant to the \mathcal{CE} machine are the call-by-need Krivine machines [14, 19, 28], and the three-instruction machine (TIM) [13]. Krivine machines started as an approach to call-by-name evaluation, and were later extended to call-by-need [19, 28, 10, 14]. The \mathcal{CE} modifies the lazy Krivine machine to capture the environment sharing given by the cactus environment. The TIM is an implementation of call-by-need and call-by-name [13]. It involves, as the name suggests, three machine instructions, TAKE, PUSH, and ENTER. In Section 6, we follow Sestoft [28] and re-appropriate these instructions for the \mathcal{CE} machine.

There has also been recent interest in *heapless* abstract machines for lazy evaluation. Danvy et al. [9] and Garcia et al. [27] independently derived similar machines from the call-by-need lambda calculus [4]. These are interesting approaches, but it is not yet clear how these machines could be implemented efficiently.

3 Environment Representations

As mentioned in Section 2, environments bind free variables to closures. There is significant flexibility in how they can be represented. In this section we review this design space in the context of existing work, both for call by value and call by need.⁴

There are two common approaches to environment representation: *flat* environments and *shared* environments (also known as linked environments) [2, 29]. A flat environment is one in which each closure has its own record of the terms its free variables are bound to. A shared environment is one in which parts of that record can be shared among multiple closures [2, 29]. For example, consider the following term:

$$(\lambda x. (\lambda y. t) (\lambda z. t_1)) t_2$$

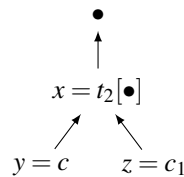
Assuming the term t has both x and y as free variables, we must evaluate it in the environment binding both x and y . Similarly, assuming t_1 contains both z and x as free variables, we must evaluate it in an environment containing bindings for both x and z . Thus, we can represent the closures for evaluating t and t_1 as

$$t[x = t_2[\bullet], y = c]$$

and

$$t_1[x = t_2[\bullet], z = c_1]$$

respectively. These are examples of *flat* environments, where each closure comes with its own record of all of its free variables. Because of the nested scope of the given term, x is bound to the same closure in the two environments. Thus, we can also create a shared, linked environment, represented by the following diagram:



Now each of the environments is represented by a linked list, with the binding of x shared between them. This is an example of a *shared* environment [2]. This shared, linked structure dates back to the first machine for evaluating expressions: Landin's SECD machine [20].

⁴ Some work refers to this space as *closure* representation rather than *environment* representation [29, 2]. Because the term part of the closure is simply a code pointer and the interesting design choices are in the environment, we refer to the topic as environment representation.

The drawbacks and advantages of each approach are well known. With a flat environment, variable lookup can be performed with a simple offset [24, 1]. On the other hand, significant duplication can occur, as we will discuss in Section 3.1. With a shared environment, that duplication is removed, but at the cost of possible link traversal upon dereference.

As with most topics in compilers and abstract machines, the design space is actually more complex. For example, Appel and Jim show a wide range of hybrids [2] between the two, and Appel and Shao [29] show an optimized hybrid that aims to achieve the benefits of both approaches. And as shown in the next section, choice of evaluation strategy further complicates the picture.

3.1 Existing Call-by-Need Environments

Existing call by need machines use flat environments with a heap of closures [24, 13, 18, 7]. These environments may contain some combination of primitive values and pointers into the heap (p below). The pointers and heap implement the memoization of results required for call by need. Returning to the earlier example, $(\lambda x.(\lambda y.t)(\lambda z.t_1))t_2$, we can view a simplified execution state for this approach when entering t as follows:

Closure

$$t[x = p, y = p_1]$$

Heap

$$\begin{aligned} p &\mapsto t_2[\bullet] \\ p_1 &\mapsto \lambda z.t_1[x = p] \end{aligned}$$

If $t_2[\bullet]$ is not in WHNF (this sort of unevaluated closure is called a *thunk* [17, 25]), then if it is entered in either the evaluation of t or t_1 , the resulting value will overwrite the closure at p . The result of the computation is then shared with all other instances of x in t and t_1 . In the case that terms have a large number of shared variables, environment duplication can be expensive. Compile-time transformation [25] (tupling arguments) helps, but we show that the machine can avoid duplication completely.

Depending on g , either or both of the closures created for its arguments may not be evaluated. Therefore, it is possible that the work of creating the environment for that thunk will be wasted. This waste is well known, and existing approaches address it by avoiding thunks as much as possible [24, 18]. Unfortunately, in cases like the above example, thunks are necessary. Indeed, even if we tuple the closures for x , y , and z , that tuple could be wasted if neither t nor t' is used. Thus, we would like to minimize the cost of creating such thunks.

Thunks are special in another way. Recall that one advantage of the flat environments is quick variable lookups. In a lazy language, this advantage is reduced because *a thunk can only be entered once*. After it is entered, it is overwritten with a value, so the next time that heap location is entered it is entered with a value and a different environment. Thus, the work to ensure that the variable lookup is fast is used only once.

This is in contrast to a call by value language, in which every closure is constructed for a value, and can be entered an arbitrary number of times.

A more subtle drawback of the flat environment representation is that environments can vary in size, and thus a value in WHNF can be too large to fit in the space allocated for the thunk it is replacing. This problem is discussed in [24], where the proposed solution is to put the value closure in a fresh location in the heap where there is sufficient room. The original thunk location is then replaced with an indirection to the value at the freshly allocated location. These indirections are removed during garbage collection, but do impose some cost, both in runtime efficiency and implementation complexity [24].

We have thus far ignored a number of details with regard to current implementations. For example, the STG machine can split the flat environment, so that part is allocated on the stack and part on the heap. The TIM allocates its flat environments separately from its closures so that each closure is a code pointer, environment pointer pair [13] while the STG machine keeps environment and code co-located [24]. Still, the basic design principle holds: a flat environment for each closure allows quick variable indexing, but with an initial overhead.

To summarize, the flat environment representation in a call by need language is whenever a term might be needed, the necessary environment is constructed from the current environment. This operation can be expensive, and it is wasted if the variable is never entered. In this work, we aim to minimize this potentially unnecessary overhead.

Figure 1 depicts the design space relevant to this paper. There are existing call by value machines with both flat and shared environments, and call by need machines with flat environments. As far as we are aware, we are the first to use a shared environment to implement lazy evaluation.

	Flat Environment	Shared Environment
Call by need	STG [24], TIM [13], GRIN [7]	$\mathcal{C}\mathcal{E}$ Machine (this paper)
Call by value	ZAM [21], SML/NJ [3]	ZAM, SECD [20], SML/NJ

Fig. 1. Evaluation strategy and environment structure design space. Each acronym refers to an existing implementation. Some implementations use multiple environment representations.

4 Cactus Environment Calculus

This section shows how the shared environment approach can be applied to call-by-need evaluation. We start with a calculus that abstracts away environment representation, Curien’s calculus of closures, and we show how it can be modified to force sharing. See Curien’s call-by-name calculus of closures in Figure 2.⁵

⁵ Curien calls it a “lazy” evaluator, and there is some ambiguity with the term lazy, but we use the term only to mean call-by-need. We also remove the condition checking that $i < m$ because we are only concerned with evaluation of closed terms.

The LEval rule pushes a closure onto the environment, and the LVar rule indexes into the environment, entering the corresponding closure. We show in this section that by removing ambiguity about how the environments are represented, and forcing them to be represented in a *cactus stack* [30], we can define our novel call-by-need calculus.

Syntax

$$\begin{array}{ll}
 t ::= i \mid \lambda t \mid t t & \text{(Term)} \\
 i \in \mathbb{N} & \text{(Variable)} \\
 c ::= t[\rho] & \text{(Closure)} \\
 \rho ::= \bullet \mid c \cdot \rho & \text{(Environment)}
 \end{array}$$

Semantics

$$\frac{t_1[\rho] \xrightarrow{*}_L \lambda t_2[\rho']}{t_1 t_3[\rho] \rightarrow_L t_2[t_3[\rho] \cdot \rho']} \quad \text{(LEval)}$$

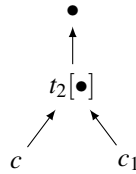
$$i[c_0 \cdot c_1 \cdot \dots \cdot c_i \cdot \rho] \rightarrow_L c_i \quad \text{(LVar)}$$

Fig. 2. Curien's call-by-name calculus of closures [8]

To start, consider again the example from Section 3, this time with deBruijn indices: $(\lambda(\lambda t) (\lambda t_1))t_2$. The terms t and t_1 , when evaluated in the closure calculus, would have the following environments, respectively:

$$\begin{array}{l}
 c \cdot t_2[\bullet] \cdot \bullet \\
 c_1 \cdot t_2[\bullet] \cdot \bullet
 \end{array}$$

Again, the second closure is identical in each environment. And again, we can represent these environments with a shared environment, this time keeping call-by-need evaluation in mind:



This inverted tree structure seen earlier with the leaves pointing toward the root is called a *cactus stack* (sometimes called a spaghetti stack or saguaro stack) [15, 16]. In this particular cactus stack, every node defines an environment as the sequence of closures in the path to the root. If $t_2[\bullet]$ is a thunk, and is updated in place with the value after its

first reference, then both environments would contain the resulting value. This is exactly the kind of sharing that is required by call-by-need, and thus we can use this structure to build a call-by-need evaluator. This is the essence of the cactus environment calculus and the cactus environment (\mathcal{CE}) machine.

Curien’s calculus of closures does not differentiate between flat and shared environment representations, and indeed, no calculus that we are aware of has had the need to. Therefore, we must derive a calculus of closures, forcing the environment to be shared. Because we can hold the closure directly in the environment, we choose to replace the standard heap of closures with a *heap of environments*. To enforce sharing, we extend Curien’s calculus of closures to include the heap of environments, which we refer to as a *cactus environment*.

See Figure 3 for the syntax and semantics of the cactus calculus. Recall that we are only concerned with evaluation of closed terms. The initial closed term t is placed in a $(t[0], \varepsilon[0 = \bullet])$ tuple, and evaluation terminates on a value. We define two different semantics, one for call-by-name and one for call-by-need, which makes the connection to Curien’s call-by-name calculus more straightforward. The rule for application (MEval and NEval) is identical for both semantics: each evaluates the left hand side to a function, then binds the variable in the cactus environment, extending the current environment. The only difference between this semantics and Curien’s is that if we need to extend an environment multiple times, the semantics *requires* sharing it among the extensions. This makes no real difference for call-by-name, but it is needed for the sharing of results in the NVar rule. The explicit environment sharing ensures that the closure that is overwritten with a value is shared correctly.

4.1 Correctness

Ariola et al. define the standard call-by-need semantics in [4]. To show correctness, we show that there is a strong bisimulation between \rightarrow_N and their operational semantics, \Downarrow (Figure 4).

Theorem 1. (Strong Bisimulation)

$$\rightarrow_N \sim \Downarrow$$

We start with a *flattening* relation between a configuration for \Downarrow and a configuration for \rightarrow_N . The deBruijn indexed terms and the standard terms are both converted to terms that use deBruijn indices for local variables and direct heap locations for free variables, locations for global variables. The flattening relation holds only when both terms are closed under their corresponding heaps. It holds trivially for the special case of initializing each configuration with a standard term and its corresponding deBruijn-indexed term, respectively. The proof is finished by induction on the step relation for each direction of the bisimulation.

4.2 Recursion

Correct sharing for recursive data values, as pointed out in [4], is not possible without adding explicit recursion. We return to this issue in Section 8, but note here that adding explicit recursion to this scheme is left for future work.

Syntax

$t ::= i \mid \lambda t \mid t t$	(Term)
$i \in \mathbb{N}$	(Variable)
$c ::= t[l]$	(Closure)
$v ::= \lambda t[l]$	(Value)
$\mu ::= \varepsilon \mid \mu[l \mapsto \rho]$	(Heap)
$\rho ::= \bullet \mid c \cdot l$	(Environment)
$l, f \in \mathbb{N}$	(Location)
$s ::= (c, \mu)$	(State)

Call-by-Name Semantics

$\frac{(t[l], \mu) \rightarrow_M (\lambda t_2[l'], \mu') \quad f \notin \text{dom}(\mu')}{(t t_3[l], \mu) \rightarrow_M (t_2[f], \mu'[f \mapsto t_3[l] \cdot l'])}$	(MEval)
$\frac{\mu(l, i) = l' \mapsto c \cdot l''}{(i[l], \mu) \rightarrow_M (c, \mu)}$	(MVar)

Call-by-Need Semantics

$\frac{(t[l], \mu) \xrightarrow{*}_N (\lambda t_2[l'], \mu') \quad f \notin \text{dom}(\mu')}{(t t_3[l], \mu) \rightarrow_N (t_2[f], \mu'[f \mapsto t_3[l] \cdot l'])}$	(NEval)
$\frac{\mu(l, i) = l' \mapsto c \cdot l'' \quad (c, \mu) \xrightarrow{*}_N (v, \mu')}{(i[l], \mu) \rightarrow_N (v, \mu'[l' \mapsto v \cdot l''])}$	(NVar)

Fig. 3. Cactus calculus syntax and semantics.

$\frac{\langle \Phi \rangle t \Downarrow \langle \Psi \rangle \lambda x. t'}{\langle \Phi, x \mapsto t, \Upsilon \rangle x \Downarrow \langle \Psi, x \mapsto \lambda x. t', \Upsilon \rangle \lambda x. t'}$	(Id)
$\frac{}{\langle \Phi \rangle \lambda x. t \Downarrow \langle \Phi \rangle \lambda x. t}$	(Abs)
$\frac{\langle \Phi \rangle t_l \Downarrow \langle \Psi \rangle \lambda x. t_n \quad \langle \Psi, x' \mapsto t_m \rangle [x'/x] t_n \Downarrow \langle \Upsilon \rangle \lambda y. t'}{\langle \Phi \rangle t_l t_m \Downarrow \langle \Upsilon \rangle \lambda y. t'}$	(App)

Fig. 4. Ariola et. al's Operational Semantics

5 \mathcal{CE} Machine

Using the calculus of cactus environments defined in the previous section, we derive an abstract machine: the \mathcal{CE} machine. The syntax and semantics are defined in Figure 5.

Syntax

$s ::= \langle c, \sigma, \mu \rangle$	(State)
$t ::= i \mid \lambda t \mid t t$	(Term)
$i \in \mathbb{N}$	(Variable)
$c ::= t[l]$	(Closure)
$v ::= \lambda t[l]$	(Value)
$\mu ::= \varepsilon \mid \mu[l \mapsto \rho]$	(Heap)
$\rho ::= \bullet \mid c \cdot l$	(Environment)
$\sigma ::= \square \mid \sigma c \mid u := \sigma$	(Context)
$l, u, f \in \mathbb{N}$	(Location)

Semantics

$\langle v, u := \sigma, \mu[u \mapsto c \cdot l] \rangle \rightarrow_{\mathcal{CE}} \langle v, \sigma, \mu[u \mapsto v \cdot l] \rangle$	(Upd)
$\langle \lambda t[l], \sigma c, \mu \rangle \rightarrow_{\mathcal{CE}} \langle t[f], \sigma, \mu[f \mapsto c \cdot l] \rangle, f \notin \text{dom}(\mu)$	(Lam)
$\langle t t'[l], \sigma, \mu \rangle \rightarrow_{\mathcal{CE}} \langle t[l], \sigma t'[l], \mu \rangle$	(App)
$\langle 0[l], \sigma, \mu \rangle \rightarrow_{\mathcal{CE}} \langle c, l := \sigma, \mu \rangle$ where $c \cdot l' = \mu(l)$	(Var1)
$\langle i[l], \sigma, \mu \rangle \rightarrow_{\mathcal{CE}} \langle (i-1)[l'], \sigma, \mu \rangle$ where $c \cdot l' = \mu(l)$	(Var2)

Fig. 5. Syntax and semantics of the \mathcal{CE} machine.

The machine operates on a similar syntax as the calculus, extended only with a context to implement the updates from the NVar subderivation ($u := \sigma$) and the operands from the NEval subderivation (σc). We also add f explicitly, which is the fresh heap location. Thus, we now have a 4-tuple, with initial state $\langle t[0], \square, \varepsilon[0 = \bullet], 1 \rangle$. The four parts of the tuple are the current closure, the context, the heap, and the fresh heap location. On the update rule, the current closure is a value, and there is an update marker as the outermost context. This implies that a variable was entered and that the current closure represents the corresponding value for that variable. Thus, we update the location u that the variable entered, replacing whatever term was entered with the current closure. The Lam rule takes an argument off the context and binds it to a variable, allocating a fresh heap location for the bound variable. This ensures that every instance of the variable will point to this location, and thus the bound term will be evaluated at most once. The App rule simply pushes an argument term in the current environment. The Var1 rule enters the closure pointed to by the environment location, while the Var2 rule traverses the cactus environment to locate the correct closure.

To get some intuition for the \mathcal{CE} machine and how it works, consider the depiction in Figure 6 of its state during evaluation of the term: $(\lambda a. (\lambda b. b a) (\lambda c. c a)) ((\lambda i. i) (\lambda j. j))$. We leave the variable names for readability. For thoroughness, the occurrences of variable a may be replaced with index 1, and the rest can be replaced with index 0. At this point in the computation, the closure for a is shared, and has been updated with its value. The closure in the context (the a from the application $c a$) now points to the updated value.

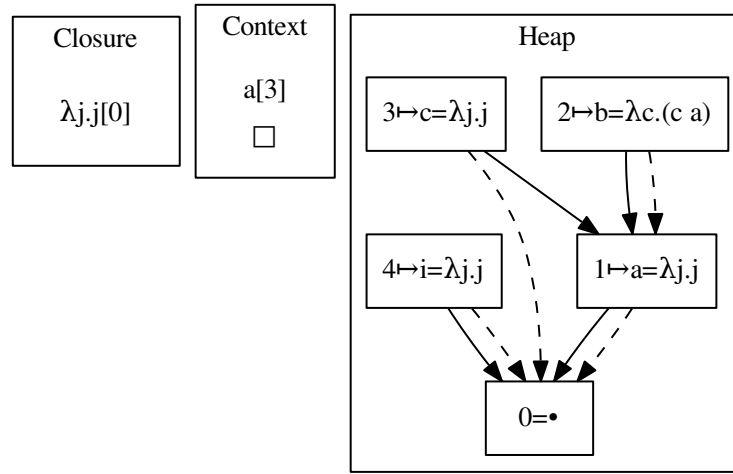


Fig. 6. \mathcal{CE} machine state example. The dotted lines represent the environment for the bound closure, and the solid lines represent the next environment cell: $t[\text{dotted}] \cdot \text{solid}$. Variables are left in for readability, but can be trivially replaced with deBruijn indices.

5.1 Correctness

We prove that the reflexive transitive closure of the \mathcal{CE} machine step relation evaluates to a value and heap and empty context iff \rightarrow_N evaluates to the same value and heap.

Theorem 2. (Equivalence)

$$(c, \mu) \rightarrow_N (v, \mu') \leftrightarrow \langle c, \square, \mu \rangle \xrightarrow{*}_{\mathcal{CE}} \langle v, \square, \mu' \rangle$$

By induction on the \rightarrow_N step relation and the reflexive transitive closure of the $\rightarrow_{\mathcal{CE}}$ step relation.

6 Implementation

This section describes how the \mathcal{CE} machine can be mapped directly to x64 instructions. Specifically, we re-define the three instructions given by the TIM [13]: TAKE, ENTER, and PUSH, and implement them with x64 assembly. We also describe several design decisions, as well as some optimizations. All implementation and benchmark code is available at <http://cs.unm.edu/~stelleg/cem-tfp2016.tar.gz>.

Each closure is represented as a code pointer, environment pointer tuple. The Context is implemented as a stack, with updates represented as a null pointer, environment pointer tuple to differentiate them from closure arguments. The Heap, or cactus environment, is implemented as a heap of environments.

6.1 Compilation

The three instructions are given below, with descriptions of their behavior.

- TAKE: Pops a context item off the stack. If the item is an update $u :=$, the instruction updates the location u with the current closure. If it is an argument c , the instruction binds the closure c to the fresh location in the cactus environment.
- ENTER i : Enters the closure defined by variable index i , the current environment pointer, and the current cactus environment.
- PUSH m : Pushes the code location m along with the current environment pointer.

These operations correspond directly to lambda terms, and to the transition functions of the \mathcal{CE} machine. The mapping from lambda terms can be seen in Figure 7, which defines the compiler. Unlike the TIM, our version of TAKE doesn't have an arity; we compile a sequence of lambdas as a sequence of TAKE instructions. While we have not compared performance of the two approaches directly, we suspect that a n inlined TAKE instructions should be roughly as fast as a TAKE n instruction. Similarly, the ENTER i instruction can be implemented either as a loop or unrolled, depending on i , and more performance comparisons are needed to determine the trade off between code size and speed.

$$\begin{aligned} C[t \ t'] &= \text{PUSH } \text{label}_{C[t']} : C[t] ++ C[t'] \\ C[\lambda .t] &= \text{TAKE} : C[t] \\ C[i] &= \text{ENTER } i \end{aligned}$$

Fig. 7. \mathcal{CE} machine compilation scheme. C compiles a sequence of instructions from a term. The *label* represents a code label: each instruction is given a unique label. The $:$ operator denotes prepending an item to a sequence and $++$ denotes concatenating two sequences.

We compile to x64 assembly. Each of the three instructions is mapped onto x64 instructions with a macro. The PUSH instruction is particularly simple, consisting of only

two x64 instructions (two pushes, one for the code pointer and one for the environment pointer). This is actually an important point: *thunk creation is only two hardware instructions, regardless of environment size.*

6.2 Machine Literals and Primitive Operations

Following Sestoft [28], we extend the $\mathcal{C}\mathcal{E}$ machine to include machine literals and primitive operations. Figure 8 shows the parts of syntax and semantics that are new or modified.

Syntax

$t ::= i \mid \lambda t \mid t t \mid m$	(Term)
$m ::= n \mid op$	(Machine)
$n \in \mathbb{I}$	(Integer)
$op ::= + \mid - \mid * \mid / \mid = \mid > \mid <$	(PrimOp)
$v ::= \lambda t[l] \mid m[l]$	(Value)

Integer and Primop Semantics

$\langle n[l], \sigma c, \mu, k \rangle \rightarrow_{\mathcal{C}\mathcal{E}} \langle c, \sigma n[l], \mu, k \rangle$	(Int)
$\langle op[l], \sigma n' n, \mu, k \rangle \rightarrow_{\mathcal{C}\mathcal{E}} \langle op(n', n)[l], \sigma, \mu, k \rangle$	(Op)

Fig. 8. Extensions to the syntax and semantics of the $\mathcal{C}\mathcal{E}$ machine.

6.3 Omitted Extensions

Our implementation omits a few other standard extensions. Here we address some of these omissions.

Data types are a common extension that we omit [24, 7]. We take the approach of Sestoft [28] that these can be efficiently implemented with pure lambda terms. For example, consider a list data type (in Haskell syntax): `data List a = Cons a (List a) | Nil`. This can be represented in pure lambda terms with $Cons = \lambda h. \lambda t. \lambda c. \lambda n. c h t$ and $Nil = \lambda c. \lambda n. n$.

Let bindings are another term commonly included in functional language compilers, even in the internal representation [7, 24]. Non-recursive let is syntactic sugar for a lambda binding and application, and we treat it as such. This approach helps ensure that we can compile arbitrary lambda terms, while some approaches require pre-processing [28, 13].

Recursive let bindings are a third omission. Here we follow Rozas [26]: If it can be represented in pure lambda terms, it should be. Thus, we implement recursion using the standard Y combinator. In the case of mutual recursion, we use the Y combinator

in conjunction with a tuple of the mutually recursive functions. Without the appropriate optimizations [26], this approach has high overhead, as we discuss in Section 7.1.

Our philosophy regarding extensions is: when an extension can be represented with pure lambda calculus, then optimized to achieve equal performance to implementations that represent them explicitly, then do so. This approach should allow more general optimizations.

6.4 Optimizations

The \mathcal{CE} implementation described in the previous section is completely unoptimized. For example, no effort is expended to discover global functions to avoid costly jumps to pointers in the heap [24]. Indeed, every variable reference will look up the code pointer in the shared environment and jump to it. There is also no implementation of control flow analysis as used by Rozas to optimize away the Y combinator. Thus, every recursive call exhibits the large overhead involved in re-calculating the fixed point of the function.

We do, however, implement two basic optimizations, primarily to reduce the load on the heap:

- POP: A TAKE instruction can be converted to a POP instruction that throws away the operand on the top of the stack if there are no variables bound to the λ term in question. For example, the function $\lambda x.\lambda y.x$ can be implemented with TAKE, POP, ENTER 0.
- ENTERVAL: An ENTER instruction, when entering a closure that is already a value, should not push an update marker onto the stack. This shortcut prevents unnecessary writes to the stack and heap [24, 14, 28].

6.5 Garbage Collection

We have implemented a simple mark and sweep garbage collector with the property that it does not require two spaces, due to constant sized closures in the heap, allowing a linked list representation for the free cells. Indeed, while the abstract machine from Section 5 increments a free heap counter, the actual implementation uses the next free cell in the linked list.

Because the focus of this paper is not on the performance of garbage collection, we ensure the benchmarks in Section 7 are not dominated by GC time.

7 Performance Evaluation

This section describes experiments that assess the strengths and weaknesses of the \mathcal{CE} machine. We evaluate using benchmarks from the `nofib` benchmark suite. Because we have implemented only machine integers, and must translate the examples by hand, we use a subset of the `nofib` suite that and exclude floating point values or arrays. A list of the benchmarks used and a brief description is given in Figure 9.

We compare the \mathcal{CE} machine to two existing implementations:

- **exp3**: A Peano arithmetic benchmark. It computes 3^8 in Peano arithmetic, and prints the result.
- **queens**: Computes the number of solutions to the nqueens problem for an n by n board.
- **primes**: A simple primes sieve that computes the nth prime.
- **digits-of-e1**: A calculation of the digits of e using continued fractions. Computes the first n digits of e .
- **digits-of-e2**: Another calculation of the digits of e using an infinite series. Computes the first n digits of e .
- **fib**: Naively computes the nth Fibonacci number.
- **fannkuch**: A counting benchmark counting the number of reverses of a subset of a list.
- **tak**: A synthetic benchmark involving basic recursion.

Fig. 9. Description of Benchmarks

- **GHC**: The Glasgow Haskell compiler. A high performance, optimizing compiler based on the STG machine [24].
- **UHC**: The Utrecht Haskell compiler. An optimizing compiler based on the GRIN machine [7, 12].

We use GHC version 7.10.3 and UHC version 1.1.9.3. We compile with -O0 and -O3, and show the results for both. Where possible, we pre-allocate a heap of 1GB to avoid measuring the performance of GC implementations. The tests were run on an Intel(R) Xeon(R) CPU E5-4650L @ 2.60GHz, running Linux version 3.16.

7.1 Results

Figure 10 gives the benchmark results. In general, we are outperformed by GHC, sometimes significantly, and we outperform UHC. We spend the remainder of the section analyzing these performance differences.

	\mathcal{L}	GHC -O0	UHC -O0	GHC -O3	UHC -O3
exp3	8	1.530	1.176	3.318	1.038
tak	16 8 0	0.366	0.146	1.510	0.006
primes	1500	0.256	0.272	1.518	0.230
queens	9	0.206	0.050	0.600	0.012
fib	35	2.234	0.872	10.000	0.110
digits-of-e1	1000	3.576	1.274	21.938	0.118
digits-of-e2	1000	0.404	0.792	3.430	0.372
fannkuch	8	0.560	0.084	2.184	0.048

Fig. 10. Machine Literals Benchmark Results. Measurement is wall clock time, units are seconds. Times averaged over 5 runs.

There are many optimizations built in to the abstract machine underlying GHC, but profiling indicates that three in particular lead to much of the performance disparity:

- **Register allocation:** The \mathcal{CE} machine has no register allocator. In contrast, by passing arguments to functions in registers, GHC avoids a great deal of heap thrashing.
- **Unpacked Literals:** Related to register allocation, this allows GHC to keep machine literals without tags in registers for tight loops. In contrast, the \mathcal{CE} machine operates entirely on the stack, and has a code pointer associated with every machine literal.
- **Y combinator:** Because recursion in the \mathcal{CE} machine is implemented with a Y combinator, it performs poorly. This could be alleviated with CFA-based techniques, similar to those used in [26].

Lack of register allocation is the primary current limitation of the \mathcal{CE} machine. The STG machine pulls the free variables into registers, allowing tight loops with everything kept in registers. However, it is less clear how to effectively allocate registers in a fully shared environment setting. That said, we believe being lazier about register allocation, e.g. not loading values into registers that may not be used, could have some performance benefits.

To isolate the effect of register allocation and unpacked machine literals, we replace the use of machine integers with Church numerals for a compatible subset of the evaluation programs. Figure 11 shows the performance results with this modification, which are much improved, with the \mathcal{CE} machine occasionally even outperforming optimized GHC.

	\mathcal{CE}	GHC -O0	UHC -O0	GHC -O3	UHC -O3
tak 14 7 0	1.610	2.428	7.936	1.016	7.782
primes 32	0.846	1.494	4.778	0.666	5.290
queens 8	0.242	0.374	1.510	0.154	1.508
fib 23	0.626	0.940	5.026	0.468	5.336
digits-of-e2 6	0.138	1.478	5.056	0.670	5.534
fannkuch 7	0.142	0.124	0.796	0.040	0.808

Fig. 11. Church Numeral Benchmark Results. Measurement is wall clock time, units are seconds. Times averaged over 5 runs.

Next, we consider the disparity due to the Y-combinator, by running a simple exponentiation example with church numerals, calculating $3^8 - 3^8 = 0$. We see that in this case, the \mathcal{CE} machine significantly outperform both GHC and UHC, as seen in Figure 12 .

	\mathcal{CE}	GHC -O0	UHC -O0	GHC -O3	UHC -O3
pow 3 8	0.564	1.994	4.912	0.906	4.932

Fig. 12. Church Numeral Exponentiation Benchmark Results. Measurement is wall clock time, units are seconds. Times averaged over 5 runs

These results give us confidence that by adding the optimizations mentioned above, among others, the $\mathcal{C}\mathcal{E}$ machine has the potential to be the basis of a real world compiler. We discuss how some of these optimizations can be applied to the $\mathcal{C}\mathcal{E}$ machine in Section 8.

7.2 The Cost of the Cactus

Recall that variable lookup is linear in the index of the variable, following pointers until the index is zero. As one might guess, the lookup cost is high. For example, for the queens benchmark without any optimizations, variable lookup took roughly 80 – 90% of the $\mathcal{C}\mathcal{E}$ machine runtime, as measured by profiling. Much of that cost was for lookups of known combinators, however, so for the benchmarks above we added the inlining mentioned in the previous section. Still, even with this simple optimization, variable lookup takes roughly 50% of the runtime. There is some variation across benchmarks, but this is a rough approximation for the average cost. We discuss how this cost could be addressed in future work in Section 8.

8 Discussion and Related Work

Some related work was discussed earlier to provide background and context. Here, we briefly and explicitly compare our approach to earlier work. We also discuss areas for future work.

8.1 Closure Representation

Appel and Shao [29] and Appel and Jim [2] both cover the design space for closure representation, and develop an approach called *safely linked closures*. Their approach uses flat closures when there is no duplication, and links in a way that preserves liveness, to prevent violation of the *safe for space complexity* (SSC) rule [1]. While we don't address SSC or garbage collection in general, understanding the relationship between SSC and shared environment call-by-need is an interesting area for future work.

8.2 Eval/Apply vs. Push/Enter

Marlow and Peyton Jones describe two approaches to the implementation of function application: *eval/apply*, where the function is evaluated and then passed the necessary arguments, and *push/enter*, where the arguments are pushed onto the stack and the function code is entered [22].

They conclude that despite *push/enter* being a standard approach to lazy machines, *eval/apply* performs better. While our current approach uses *push enter*, investigating whether or not *eval/apply* could be usefully implemented for the $\mathcal{C}\mathcal{E}$ machine is an interesting avenue for future work.

8.3 Collapsed Markers

Friedman et al. show how a machine can be designed to prevent multiple adjacent update markers being pushed onto the stack [14]. This property is desirable because multiple adjacent update markers are always updated with the same value. They give examples showing that in some cases, these redundant update markers can cause an otherwise constant-spaced stack to grow unbounded. They implement an optimization that collapses update markers by adding a layer of indirection between heap locations and closures. We propose a similar approach, but without the performance hit caused by an extra layer of indirection. Upon a variable dereference the \mathcal{CE} machine checks if the top of the stack is an update. If it is, instead of pushing a redundant update marker onto the stack, it replaces the closure in the heap at the desired location with an update marker. Then, the variable dereference rule checks for an update marker upon dereference, and will update accordingly. We have begun to implement this optimization, but leave the full implementation and description for future work.

8.4 Register Allocation

One advantage of flat environments is that register allocation is straightforward [1, 24, 31]. It is less obvious how to do register allocation with the \mathcal{CE} machine. We speculate that *only strict free variables should be loaded into registers*. That is to say, the environment variables may not be used, and only the ones we know will be used should be loaded into registers and the rest should be loaded on demand.

8.5 Verification

One attractive property of our implementation is its simplicity, which makes it an attractive target for a verified compiler. Because it avoids complexities required for flat environment implementations, e.g. black hole updates, basing a verified compiler on this machine is another exciting area for future work.

9 Conclusion

Lazy evaluation has long suffered from high overhead when delaying computations. While strictness analysis helps to alleviate this issue, there are and will always be cases it cannot catch. Existing implementations choose to pay an up-front cost of constructing a flat environment to ensure efficient variable lookup if a delayed computation is used. When a delayed computation is never used, this overhead is wasted. In this paper we have presented a novel approach that minimizes this overhead. We have achieved this overhead minimization by taking an old idea, shared environments, and using them in a novel way. By leveraging the structure inherent in a shared environment to share results of computation, we have avoided some of the overheads involved in delaying a computation.

We conclude by summarizing the key points of this paper. First, a shared environment, explicitly represented as a cactus stack, is a natural way to share the results of

computation as required by lazy evaluation. Second, this approach is in a sense *lazier* about lazy evaluation than existing implementations because it avoids some unnecessary packaging. Third, this approach can be formalized in both big-step and small-step semantics. Lastly, the abstract machine can be implemented as a compiler in a straightforward way, yielding performance comparable to existing implementations.

10 Acknowledgments

This material is based upon work supported by the National Science Foundation under grant CCF-1422840. Sandia National Laboratories is a multi-program laboratory managed and operated by Sandia Corporation, a wholly owned subsidiary of Lockheed Martin Corporation, for the U.S. Department of Energys National Nuclear Security Administration under contract DE-AC04-94AL85000.

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