

# Efficient Dual-Numbers Reverse AD via Well-Known Program Transformations

TOM J. SMEDING, Utrecht University, The Netherlands

MATTHIJS I. L. VÁKÁR, Utrecht University, The Netherlands

Where dual-numbers forward-mode automatic differentiation (AD) pairs each scalar value with its tangent value, dual-numbers *reverse-mode* AD attempts to achieve reverse AD using a similarly simple idea: by pairing each scalar value with a backpropagator function. Its correctness and efficiency on higher-order input languages have been analysed by Brunel, Mazza and Pagani, but this analysis used a custom operational semantics for which it is unclear whether it can be implemented efficiently. We take inspiration from their use of *linear factoring* to optimise dual-numbers reverse-mode AD to an algorithm that has the correct complexity and enjoys an efficient implementation in a standard functional language with support for mutable arrays, such as Haskell. Aside from the linear factoring ingredient, our optimisation steps consist of well-known ideas from the functional programming community. We demonstrate the practical use of our technique by providing a performant implementation that differentiates most of Haskell98.

Additional Key Words and Phrases: automatic differentiation, source transformation, functional programming

## 1 INTRODUCTION

An increasing number of applications requires computing derivatives of functions specified by a computer program. The derivative of a function gives more qualitative information of its behaviour around a point (i.e. the local shape of the function's graph) than just the function value at that point. This qualitative information is useful, for example, for optimising parameters along the function (because the derivative tells you how the function changes) or inferring statistics about the function (e.g. an approximation of its integral). These uses appear, respectively, in parameter optimisation in machine learning or numerical equation solving, and in Bayesian inference of probabilistic programs. Both application areas are highly relevant today.

Automatic differentiation (AD) is the most effective technique for efficient computation of derivatives of programs, and comes in two main flavours: forward AD and reverse AD. In practice, by far the most common case is that functions have many input parameters and few, or even only one, output parameters; in this situation, forward AD is inefficient while reverse AD yields the desired computational complexity. Indeed, reverse AD promises to compute the gradient of a function implemented as a program in time at most a constant factor more than runtime of the original program. However, reverse AD is also significantly more difficult to implement flexibly, correctly and efficiently than forward AD.

Many approaches exist for doing reverse AD on a higher-order language: using taping/tracing in an imperative language (e.g. [Paszke et al. 2017]) and in a functional language [Kmett and contributors 2021], using linearisation and transposition code transformations [Paszke et al. 2021], or sometimes specialised by taking advantage of common usage patterns in domain-specific languages [Schenck et al. 2022]. In the theory community, various algorithms have been described that apply to a wide variety of source languages, including approaches based on symbolic execution and tracing [Abadi and Plotkin 2020; Brunel et al. 2020] and on category theory [Vákár and Smeding

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Authors' addresses: Tom J. Smeding, Department of Information and Computing Sciences, Utrecht University, Utrecht, The Netherlands, t.j.smeding@uu.nl; Matthijs I. L. Vákár, Department of Information and Computing Sciences, Utrecht University, Utrecht, The Netherlands, m.i.l.vakar@uu.nl.

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$\lambda(x : \mathbb{R}, y : \mathbb{R}).$	$\lambda((x : \mathbb{R}, dx : \mathbb{R}), (y : \mathbb{R}, dy : \mathbb{R})).$	$\lambda((x : \mathbb{R}, dx : \mathbb{R} \multimap (\mathbb{R}, \mathbb{R})), (y : \mathbb{R}, dy : \mathbb{R} \multimap (\mathbb{R}, \mathbb{R}))).$
<b>let</b> $z = x + y$	<b>let</b> $(z, dz) = (x + y, dx + dy)$	<b>let</b> $(z, dz) = (x + y, \underline{\lambda}(d : \mathbb{R}). dx \cdot d + dy \cdot d)$
<b>in</b> $x \cdot z$	<b>in</b> $(x \cdot z, x \cdot dz + z \cdot dx)$	<b>in</b> $(x \cdot z, \underline{\lambda}(d : \mathbb{R}). dz \cdot (x \cdot d) + dx \cdot (z \cdot d))$
(a) The original program	(b) Dual-numbers forward AD	(c) Dual-numbers reverse AD

Fig. 1. An example program together with its derivative, both using dual-numbers forward AD and using dual-numbers reverse AD. The original program is of type  $(\mathbb{R}, \mathbb{R}) \rightarrow \mathbb{R}$ .

2022], as well as formalisations of existing implementations [Krawiec et al. 2022]. Despite the fact that all these source languages could, theoretically, be translated to a single generic higher-order functional language, each reverse AD algorithm takes a different approach to solve the same problem. It is unclear how exactly these algorithms relate to each other, meaning that correctness proofs (if any) need to be rewritten for each individual algorithm.

This paper aims to improve on the situation by providing a link from the elegant dual-numbers reverse AD algorithm analysed in [Brunel et al. 2020] to a functional taping approach as used in [Kmett and contributors 2021] and analysed in [Krawiec et al. 2022]. The key point made by Brunel, Mazza and Pagani [Brunel et al. 2020] is that one can attain the right computational complexity by starting from the very elegant dual-numbers reverse AD code transformation (Sections 2 and 3), and adding a *linear factoring* rule to the operational semantics of the output language of the code transformation. This linear factoring reduction rule states that for linear functions  $f$ , the expression  $f \ x + f \ y$  should be reduced to  $f \ (x + y)$ .

Our main contributions are the following:

- We show how the theoretical analysis based on the linear factoring rule can be used as a basis for an algorithm that assumes normal, call-by-value semantics. We do this by *staging calls to backpropagators* in Section 4.
- We show how this algorithm can be made efficient by using the standard functional programming techniques of Cayley transformation (Section 5) and (e.g. linearly typed or monadic) functional in-place updates (Section 7).
- We explain how our algorithm relates to classical taping-based approaches (Section 8).
- We give an implementation of the final algorithm of Section 8.2 that can differentiate most of Haskell98 (but using call-by-value semantics), and that exhibits the correct complexity in practice, as well as reasonable constant-factor performance (Section 10).
- We explain in detail how our technique relates to the functional taping AD of [Kmett and contributors 2021] and [Krawiec et al. 2022] as well as [Shaikhha et al. 2019]’s approach of trying to optimise forward AD to reverse AD at compile time (Section 11). We also briefly describe the broader relationship with related work.

## 2 KEY IDEAS

*Naive dual-numbers reverse AD.* In traditional dual-numbers *forward* AD, one pairs up the real scalars in the input of a program with their *tangent* (these tangents together form the *directional derivative* of the input), runs the program with overloaded arithmetic operators to propagate forward these tangents, and finally reads the tangent of the output from the tangents paired up with the output scalars of the program. For example, transforming the program in Fig. 1a using dual-numbers forward AD yields Fig. 1b.

For reverse AD, such an elegant formulation is also possible, but we have to somehow encode the “reversal” in the tangent scalars that we called  $dx$  and  $dy$  in Fig. 1b. A solution is to replace those tangent scalars with *linear functions* that take the *cotangent* (or *reverse derivative*, or *adjoint*) of the scalar it is paired with, and return the cotangent of the full input of the program. Transforming the

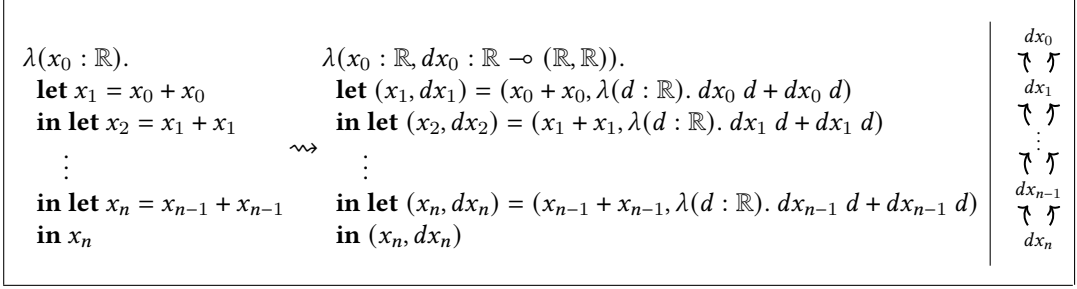


Fig. 2. Left: an example showing how naive dual-numbers reverse AD can result in exponential blow-up when applied to a program with sharing. Right: the dependency graph of the backpropagators  $dx_i$ .

same example program Fig. 1a using this style of reverse AD yields Fig. 1c. The linearity indicated by the  $\multimap$ -arrow here is that of a monoid homomorphism (a function preserving 0 and (+)); however, operationally, linear functions behave just like regular functions.

This naive dual-numbers reverse AD transformation, which we list in Fig. 6, is simple and it is easy to see that it is correct via a logical relations argument [Nunes and Vákár 2022]. The idea of this argument is to prove via induction that a backpropagator  $x' : \mathbb{R} \multimap c$  that is paired with an intermediate value  $x : \mathbb{R}$  in the program, holds the gradient of the computation that calculates  $x : \mathbb{R}$  from the global input of type  $c$ .

Dual-numbers forward AD has the very useful property that it generalises over many types (e.g. products, coproducts, recursive types) and program constructs (e.g. recursion, higher-order functions), thereby being applicable to e.g. all of Haskell98; the same property is inherited by the style of dual-numbers reverse AD exemplified here. However, unlike dual-numbers forward AD (which can propagate tangents through a program with only a constant-factor overhead over the original runtime), dual-numbers reverse AD is wildly inefficient: calling  $dx_n$  returned by the differentiated program in Fig. 2 takes time *exponential* in  $n$ . Such overhead would make reverse AD completely useless in practice—particularly because other (less flexible) reverse AD algorithms exist that indeed do a lot better. (See e.g. [Baydin et al. 2017].)

However, it turns out that this naive form of dual-numbers reverse AD can be *optimised* to be as efficient (in terms of time complexity) as these other algorithms—and most of these optimisations are just applications of standard functional programming techniques. This paper presents a sequence of changes to the code transformation (see the overview in Fig. 3) that fix all the complexity issues and, in the end, produce an algorithm with which the differentiated program has only a constant-factor overhead in runtime over the original program. This complexity is as desired from a reverse AD algorithm, and is best possible, while nevertheless being applicable to a wide range of programming language features. We explain how the result is essentially equivalent to classical taping techniques.

*Optimisation steps.* We present, in Fig. 3, an overview of the optimisations that we apply to the dual-numbers reverse AD algorithm to fix its complexity problems. We discuss these one by one.

We first apply *linear factoring*: for a linear function  $f$ , such as a backpropagator, we have that  $f \ x + f \ y = f \ (x + y)$ . Observing the form of the backpropagators in Fig. 6, we see that in the end all we produce is a giant sum of applications of backpropagators to scalars; hence, in this giant sum, we should be able to contract applications to the same backpropagator using this linear factoring rule.

We achieve this linear factoring by not returning a plain  $c$  (presumably the type of the program input) from our backpropagators, but instead a  $c$  wrapped in an object that can delay calls to linear functions producing a  $c$ . This object we call *Staged*; aside from changing the monoid that we are mapping into from  $(c, 0, (+))$  to  $(\text{Staged } c, 0_{\text{Staged}}, (+_{\text{Staged}}))$ , the only material change is that the

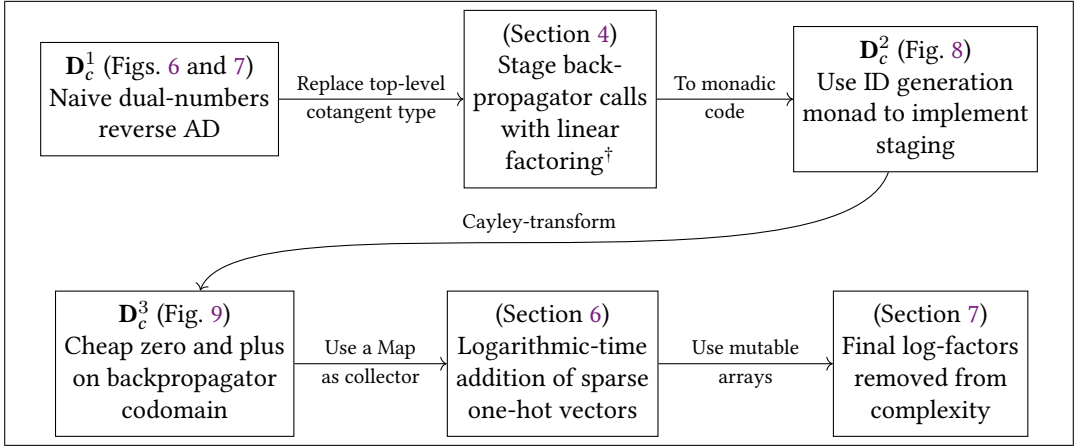


Fig. 3. Overview of the optimisations to dual-numbers reverse AD as a code transformation that are described in this paper. (<sup>†</sup> = inspired by [Brunel et al. 2020])

calls to  $d_i$  in  $\mathbf{D}_c^1[op]$  are now wrapped using a new function `SCall`, which delays the calls to  $d_i$  by storing the relevant metadata in the returned `Staged` object.

However, it is not obvious how to implement this `Staged` type: we need a linear order on (linear) function values if they are to serve as keys in a tree map, which the `Staged` interface currently prescribes. Furthermore, even if we can delay calls to backpropagators, we still need to call them at some point, and it is unclear in what order we should do so (and this order turns out to be very important). We solve these problems by generating, at runtime, a unique identifier (ID) for each backpropagator that we create, which we do by letting the differentiated program run in an ID generation monad (a special case of a state monad). The result is shown in Fig. 8, which is very similar to the previous version in Fig. 6 apart from threading through the next-ID-to-generate. (The code looks very different, but this is only due to monadic bookkeeping.)

At this point, the code transformation reaches a significant milestone: by staging (delaying) calls to backpropagators as long as possible, we can ensure that *every backpropagator is called at most once*. This milestone is achieved using the following observation: if we assign incrementing numeric IDs at runtime to lambda functions in a pure functional program, then the runtime closure of a lambda function can only refer to other functions with *smaller* IDs. Now, since our backpropagators only call other functions contained in their closure (and not functions contained in their input argument), we can use the observation to conclude that all backpropagator calls are to other backpropagators with smaller IDs! This allows us to resolve all backpropagator calls in the `Staged` object produced by the differentiated program by simply calling them from highest ID to lowest ID, and collecting and combining (using linear factoring) the calls made along the way. (See Section 4.)

But we are not done yet. The code transformation at this point ( $\mathbf{D}_c^2$  in Fig. 8) still has a glaring problem: orthogonal to the issue that backpropagators were called too many times (which we fixed), we are still creating one-hot input cotangents and adding those together. This problem is somewhat more subtle, because it is not actually apparent in the program transformation itself; indeed, looking back at Fig. 1c, there are no one-hot values to be found. However, the only way to *use* the program in Fig. 1c to do something useful, namely to compute the cotangent (gradient) of the input, is to pass  $(\lambda z. (z, 0))$  to  $dx$  and  $(\lambda z. (0, z))$  to  $dy$ ; it is easy to see that generalising this to larger input structures results in input values like  $(0, \dots, 0, z, 0, \dots, 0)$  that get added together. Adding many zeros together can hardly be the most efficient way to go about things, and indeed this is a complexity issue in the algorithm.

The way we solve this problem of one-hots is less AD-specific: the most important optimisations that we perform are Cayley-transformation (Section 5) and using a better sparse vector representation (Map  $\text{Int } \mathbb{R}$  instead of a plain  $c$  value; Section 6). Cayley-transformation (also known as *difference lists* [Hughes 1986] in the Haskell community) is a classic technique in functional programming that represents an element  $m$  of a monoid  $M$  (in this paper, written additively) by the function  $m + - : M \rightarrow M$  it induces through addition. Cayley-transformation helps us because the monoid  $M \rightarrow M$  has very cheap zero and plus operations:  $\text{id}$  and  $(\circ)$ . Afterwards, using a better (sparse) representation for the value in which we collect the final gradient, we can ensure that adding a one-hot value to this gradient collector can be done in logarithmic time.

By now, the differentiated program can compute the gradient with a *logarithmic* overhead over the original program. If a logarithmic overhead is not acceptable, the log-factor in the complexity can be removed by using functional mutable arrays (Section 7).

And then we are done, because we have now obtained a code transformation with the right complexity: the differentiated program computes the gradient of the source program at some input with runtime proportional to the runtime of the source program.

*Comparison to other algorithms.* We can relate our technique to that of [Krawiec et al. 2022] by noting that we can replace  $\mathbf{D}_c^1[\mathbb{R}] = (\mathbb{R}, \mathbb{R} \multimap c)$  with the isomorphic definition  $\mathbf{D}_c^1[\mathbb{R}] = (\mathbb{R}, c)$ . This turns the linear factoring rule into a *distributive law*  $v \cdot x + v \cdot y \rightsquigarrow v \cdot (x + y)$  that is effectively applied at runtime by using an intensional representation of the cotangent expressions of type  $c$ . While their development is presented very differently and the equivalence is not at all clear at first sight, we explain the correspondence in Section 8.

This perspective also makes clear the relationship between our technique and that of [Shaikhha et al. 2019]. Where they try to optimise vectorised forward AD to reverse AD at *compile-time* by using a distributive law (which sometimes succeeds for sufficiently simple programs), our technique proposes a clever way of efficiently applying the distributive law in the required places at *run-time*, giving us the power to always achieve the desired reverse AD behaviour.

Finally, we are now in the position to note the similarity to taping-based AD as in [Kmett and contributors 2021; Krawiec et al. 2022]: the incrementing IDs that we attached to backpropagators earlier give a mapping from  $\{0, \dots, n\}$  to our backpropagators. Furthermore, each backpropagator corresponds to either a primitive arithmetic operation performed in the source program, or to an input value; this already means that we have a tape, in a sense, of all performed primitive operations, albeit in the form of a chain of closures. The optimisation using mutable arrays then eliminates also this last difference, because there we actually reify this tape in a large array.

### 3 NAIVE, UNOPTIMISED DUAL-NUMBERS REVERSE AD

We first describe the naive implementation of dual-numbers reverse AD: this algorithm is easy to define and prove correct compositionally, but it is wildly inefficient in terms of complexity. Indeed, it tends to blow up to exponential overhead over the original function, whereas the desired complexity is to have only a constant factor overhead over the original function. Later, we will apply a number of optimisations to this algorithm (in Section 4 and onwards) that fix the complexity issues, to derive an algorithm that does have the desired complexity.

#### 3.1 Source and target languages

The reverse AD methods in this paper are code transformations, and hence have a source language (in which input programs may be written) and a target language (in which gradient programs are expressed). While the source language will be identical for all versions of the transformation that we discuss, the target language will expand to support the optimisations that we perform.

<b>Types:</b>	$\sigma, \tau ::= \mathbb{R} \mid () \mid (\sigma, \tau) \mid \sigma \rightarrow \tau \mid \text{Int}$
<b>Terms:</b>	$s, t ::= x$ (variable references, where $x$ is an identifier) $  ()$ (unit constructor) $  (s, t)$ (pair constructor) $  \text{fst}(t) \mid \text{snd}(t)$ (pair projections) $  s t$ (function application) $  \lambda(x : \tau). t$ (lambda abstraction) $  \text{let } x : \tau = s \text{ in } t$ (let binding) $  r$ (literal $\mathbb{R}$ values) $  \text{op}(t_1, \dots, t_n)$ ( $\text{op} \in \text{Op}_n$ , primitive operation application ( $\mathbb{R}^n \rightarrow \mathbb{R}$ ))

Fig. 4. The source language of all variants of this paper's reverse AD transformation.

<b>Types:</b>	$\sigma, \tau ::= \mathbb{R} \mid () \mid (\sigma, \tau) \mid \sigma \rightarrow \tau \mid \text{Int}$ $  \sigma \multimap \tau$ (linear functions)
<b>Terms:</b>	$s, t ::= x \mid () \mid (s, t) \mid \text{fst}(t) \mid \text{snd}(t) \mid s t$ $  \lambda(x : \tau). t \mid \text{let } x : \tau = s \text{ in } t$ $  r \mid \text{op}(t_1, \dots, t_n)$ $  \underline{\lambda}(z : \tau). b$ (linear lambda abstraction ( $\tau$ plain-data type))
<b>Linear function bodies:</b>	$b ::= () \mid (b, b') \mid \text{fst}(b) \mid \text{snd}(b)$ (tupling) $  z$ (reference to $\underline{\lambda}$ -bound variable) $  x b$ (linear function application; $x : \sigma \multimap \tau$ is an identifier) $  \partial_i \text{op}(x_1, \dots, x_n)(b)$ ( $\text{op} \in \text{Op}_n$ , $i$ 'th partial derivative of $\text{op}$ ( $\mathbb{R}^n \rightarrow \mathbb{R}$ )) $  b + b'$ (elementwise addition of results) $  \underline{0}$ (zero of result type)

Fig. 5. The target language of the unoptimised variant of the reverse AD transformation. Components that are also in the source language (Fig. 4) are set in grey.

The source language is defined in Fig. 4; the initial target language is given in Fig. 5. The typing of the source language is completely standard, so we omit typing rules here. We assume call-by-value evaluation. The only part that warrants explanation is the treatment of primitive operations: for all  $n \in \mathbb{Z}_{>0}$  we presume the presence of a set  $\text{Op}_n$  containing primitive operations of type  $\mathbb{R}^n \rightarrow \mathbb{R}$  in the source language. The program transformation does not care what the contents of  $\text{Op}_n$  are, as long as their derivatives are available in the target language after differentiation.

In the target language in Fig. 5, we add linear functions with the type  $\sigma \multimap \tau$ : these functions are linear in the sense of being monoid homomorphisms, meaning that if  $f : \sigma \multimap \tau$ , then  $\llbracket f \rrbracket(0) = 0$  and  $\llbracket f \rrbracket(x + y) = \llbracket f \rrbracket(x) + \llbracket f \rrbracket(y)$ . Because it is not well-defined what the derivative of a function value (in the input or output of a program) should be, we disallow function types on either side of the  $\multimap$ -arrow.<sup>1</sup> (Note that higher-order functions *within* the program are fine; the full program should just have first-order input and output types.) Operationally, however, linear functions are just regular functions: the operational meaning of all code in this paper remains identical if all  $\multimap$ -arrows are replaced with  $\rightarrow$  (and partial derivative operations are allowed in regular terms).

<sup>1</sup>In Section 5 we will, actually, put endomorphisms ( $a \rightarrow a$ ) on both sides of a  $\multimap$ -arrow; for justification, see there.





Variable references, tuples, projections, function application, lambda abstraction and let-binding are mapped homomorphically, i.e., the code transformation simply recurses over the subterms of the current term. However, note that for variable references, lambda abstraction and let-binding, the types of the variables do change.

Scalar constants are transformed to a pair of that scalar constant and a backpropagator that produces the derivative of the input given the derivative of this subterm. This input derivative is zero, since the current subterm (a scalar constant) does not depend on the input.

Finally, primitive scalar operations are the most important place where this code transformation does something non-trivial. First, we compute the values and backpropagators of the (scalar) arguments to the operation, after which we can compute the original (scalar) result by applying the original operation to those argument values. Second, we return the backpropagator of the result of the operation, which applies all partial derivatives of the primitive operation to the incoming cotangent derivative, passes the results on to the corresponding backpropagators of the arguments, and finally adds the (top-level input derivative) results of type  $c$  together.

### 3.3 Wrapper of the AD transformation: exposing a usable API

*The ideal API of reverse AD.* Given a program  $(\lambda(x : \sigma). t) : \sigma \rightarrow \tau$  that computes a differentiable function  $\llbracket \lambda(x : \sigma). t \rrbracket : \llbracket \sigma \rrbracket \rightarrow \llbracket \tau \rrbracket$ , where  $\sigma, \tau$  do not contain function types, we expect the following type for its reverse derivative:

$$\text{Wrap}^1[\lambda(x : \sigma). t] : \sigma \rightarrow (\tau, \underline{\tau} \multimap \underline{\sigma})$$

Here, we write  $\underline{\tau}$  and  $\underline{\sigma}$  for the types of cotangent vectors to  $\tau$  and  $\sigma$ ; for example,  $\underline{\mathbb{R}} = \mathbb{R}$  and  $(\underline{\sigma}, \underline{\tau}) = (\underline{\sigma}, \underline{\tau})$ . The idea is that  $\text{Wrap}^1[\lambda(x : \sigma). t]$  computes the function  $x \mapsto (\llbracket \lambda(x : \sigma). t \rrbracket(x), v \mapsto D_x \llbracket \lambda(x : \sigma). t \rrbracket^t(v))$ , where we write  $D_x f^t$  for the transposed derivative of  $f$  at the point  $x$ .

*A step towards the right type.* This type seems quite different from that of our AD transformation of Fig. 6. Suppose that our AD transformation instead had the following type:

$$\tilde{\mathbf{D}}_c^1[\lambda(x : \sigma). t] : \forall c. (\sigma, \underline{\sigma} \multimap c) \rightarrow (\tau, \underline{\tau} \multimap c),$$

Then by the Yoneda lemma and choosing  $c = \sigma$ , we can build our desired wrapper with the definition  $\text{Wrap}^1[\lambda(x : \sigma). t] = \lambda(y : \sigma). \tilde{\mathbf{D}}_c^1[\lambda(x : \sigma). t](y, \text{id})$ .

*(De)interleaving backpropagators.* The typing of  $\tilde{\mathbf{D}}_c^1[\lambda(x : \sigma). t]$  feels similar to the typing of the transform in Fig. 6:

$$\mathbf{D}_c^1[\lambda(x : \sigma). t] : \mathbf{D}_c^1[\sigma] \rightarrow \mathbf{D}_c^1[\tau]$$

Indeed, the missing part is interleaving and deinterleaving of the backpropagators, implementing the isomorphism of linear function types  $(\underline{\sigma}, \underline{\tau}) \multimap c \cong (\underline{\sigma} \multimap c, \underline{\tau} \multimap c)$ . By doing such interleaving, we can define:

$$\tilde{\mathbf{D}}_c^1[\lambda(x : \sigma). t] = \lambda(y : (\sigma, \underline{\sigma} \multimap c)). \text{Deinterleave}_c^1(\mathbf{D}_c^1[\lambda(x : \sigma). t](\text{Interleave}_c^1 y))$$

Suitable definitions of  $\text{Interleave}^1$  and  $\text{Deinterleave}^1$  are shown in Fig. 7, including the resulting definition of  $\text{Wrap}^1$  (where we already inlined the transformation for the top-level lambda).

### 3.4 Complexity of the naive transformation

Reverse AD transformations like the one described in this section are well-known to be correct (e.g. [Brunel et al. 2020; Huot et al. 2020; Mazza and Pagani 2021; Nunes and Vákár 2022]). However, as given here, it does not at all have the right time complexity.

The forward pass is fine: computing the value of  $\text{Wrap}^1[\lambda(x : \sigma). t : \tau] : \sigma \rightarrow (\tau, \tau \multimap \sigma)$  takes time proportional to the original program  $t$ . However, the problem arises when we call the



$\text{Interleave}_{\tau}^1 : \forall c. (\tau, \tau \multimap c) \rightarrow \mathbf{D}_c^1[\tau]$ $\text{Interleave}_{\mathbb{R}}^1 = \lambda(x, d). (x, d)$ $\text{Interleave}_{\mathbb{0}}^1 = \lambda(). ((), d). ()$ $\text{Interleave}_{(\sigma, \tau)}^1 = \lambda((x, y), d). (\text{Interleave}_{\sigma}^1(x, \underline{\lambda}(z : \sigma). d(z, \mathbb{0})), \text{Interleave}_{\tau}^1(y, \underline{\lambda}(z : \tau). d(\mathbb{0}, z)))$ $\text{Interleave}_{\text{Int}}^1 = \lambda(n, d). n$ $\text{Interleave}_{\sigma \rightarrow \tau}^1 = \text{not defined!}$ $\text{Deinterleave}_{\tau}^1 : \forall c. \mathbf{D}_c^1[\tau] \rightarrow (\tau, \tau \multimap c)$ $\text{Deinterleave}_{\mathbb{R}}^1 = \lambda(x, d). (x, d)$ $\text{Deinterleave}_{\mathbb{0}}^1 = \lambda(). ((), \underline{\lambda}(z : ()). \mathbb{0})$ $\text{Deinterleave}_{(\sigma, \tau)}^1 = \lambda(x, y). \mathbf{let} (x_1, x_2) = \text{Deinterleave}_{\sigma}^1 x$ $\qquad \qquad \qquad \mathbf{in} \mathbf{let} (y_1, y_2) = \text{Deinterleave}_{\tau}^1 y$ $\qquad \qquad \qquad \mathbf{in} ((x_1, y_1), \underline{\lambda}(z : (\sigma, \tau)). x_2(\text{fst}(z)) + y_2(\text{snd}(z)))$ $\text{Deinterleave}_{\text{Int}}^1 = \lambda n. (n, \underline{\lambda}(z : \text{Int}). \mathbb{0})$ $\text{Deinterleave}_{\sigma \rightarrow \tau}^1 = \text{not defined!}$ $\text{Wrap}^1 : (\sigma \rightarrow \tau) \rightsquigarrow (\sigma \rightarrow (\tau, \tau \multimap \sigma))$ $\text{Wrap}^1[\lambda(x : \sigma). t] = \lambda(x : \sigma). \mathbf{let} x : \mathbf{D}_{\sigma}^1[\sigma] = \text{Interleave}_{\sigma}^1(x, \text{id}) \mathbf{in} \text{Deinterleave}_{\tau}^1(\mathbf{D}_{\sigma}^1[t])$
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Fig. 7. Wrapper around  $\mathbf{D}_c^1$  of Fig. 6.

top-level backpropagator returned by the wrapper. When we do so, we start a tree of calls to the linear backpropagators of all scalars in the program, where the backpropagator corresponding to a particular scalar value will be invoked once for each usage of that scalar as an argument to a primitive operation. This means that any sharing of scalars in the original program results in multiple calls to the same backpropagator in the derivative program. Fig. 2 displays an example program  $t$  with its naive derivative  $\mathbf{D}_c^1[t]$ , in which sharing of scalars results in exponential time complexity.

This overhead is unacceptable: we can do much better. For first-order programs, we understand well how to write a code transformation such that the output program computes the gradient in only a constant factor overhead over the original program [Griewank and Walther 2008]. This is less immediately clear for higher-order programs, as we consider here, but it is nevertheless possible.

In [Brunel et al. 2020], this problem of exponential complexity is addressed by observing that calling a linear backpropagator multiple times is actually a waste of work: indeed, linearity of a backpropagator  $f$  means that  $f x + f y = f(x + y)$ . Hopefully, applying this *linear factoring rule* from left to right (thereby taking together two calls into one) allows us to ensure that every backpropagator is executed at most once.

And indeed, should we achieve this, the complexity issue described above (the exponential blowup) is fixed: every created backpropagator corresponds to some computation in the original program (either a primitive operation, a scalar constant or an input value), so with maximal application of linear factoring, the number of backpropagator executions would become proportional to the runtime of the original program. If we can further make the body of a single backpropagator (not counting its callees) constant-time,<sup>2</sup> the differentiated program will compute the gradient with only a constant-factor overhead over the original program—as it should be for reverse AD.

<sup>2</sup>Obstacles to this are e.g.  $\mathbb{0}$  and  $(+)$  on the type  $c$ ; we will fix this in Sections 5 to 7.

However, this argument crucially depends on us being able to ensure that every backpropagator gets invoked at most once. The solution of [Brunel et al. 2020] is to symbolically evaluate the output program of the transformation to a straight-line program with the input backpropagators still as symbolic variables, and afterwards symbolically reduce the obtained straight-line program in a very specific way, making use of the linear factoring rule ( $f x + f y = f (x + y)$ ) in judicious places.

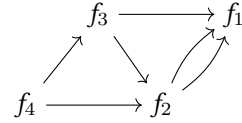
In this paper, we present an alternative way to use linear factoring to make standard, call-by-value evaluation of the target language have the correct computational complexity without any need for symbolic execution. We achieve this by changing the type  $c$  that the input backpropagators map to, to a more intelligent type than the space of cotangents of the input that we have considered so far.

#### 4 LINEAR FACTORING BY STAGING FUNCTION CALLS

As observed above in Section 3.4, the most important complexity problem of the reverse AD algorithm is solved if we ensure that all backpropagators are invoked at most once, and for that we must use that every linear function  $f$  satisfies  $f x + f y = f (x + y)$ . This means that we must find a way to “merge” all invocations of a single backpropagator using this linear factoring rule so that in the end only one invocation remains (or zero if it was not invoked at all in the first place).

*Evaluation order.* Ensuring this complete merging of linear function calls is really a question of choosing an order of evaluation for the tree of function calls created by the backpropagators. Consider for example the (typical) situation where a program generates the following backpropagators:

$$\begin{aligned} f_1 &= \lambda(z : \mathbb{R}). (0, (z, 0)) \\ f_2 &= \lambda(z : \mathbb{R}). f_1 (2 \cdot z) + f_1 (3 \cdot z) \\ f_3 &= \lambda(z : \mathbb{R}). f_2 (4 \cdot z) + f_1 (5 \cdot z) \\ f_4 &= \lambda(z : \mathbb{R}). f_2 z + f_3 (2 \cdot z) \end{aligned}$$



and where  $f_4$  is the (only) backpropagator contained in the result. Normal call-by-value evaluation of  $f_4$  would yield two invocations of  $f_2$  and five invocations of  $f_1$ , following the displayed call graph.

However, taking inspiration from symbolic evaluation and moving away from standard call-by-value for a moment, we could also first invoke  $f_3$  to expand the body of  $f_4$  to  $f_2 z + f_2 (4 \cdot (2 \cdot z)) + f_1 (5 \cdot (2 \cdot z))$ . Now we can take the two invocations of  $f_2$  together using linear factoring to produce  $f_2 (z + 4 \cdot (2 \cdot z)) + f_1 (5 \cdot (2 \cdot z))$ ; then invoking  $f_2$  first, producing two more calls to  $f_1$ , we are left with three calls to  $f_1$  which we can take together to a single call using linear factoring, which we can then evaluate. With this alternate evaluation order, we have indeed ensured that every linear function is invoked at most (in this case, exactly) once.

If we want to obtain something like this evaluation order, the first thing that we must achieve is to *postpone* invocation of linear functions until we conclude that we have merged all calls to that function and that its time for evaluation has arrived. To achieve this goal, we would like to change the representation of  $c$  to a dictionary mapping linear functions to the argument at which we intend to later call them.<sup>3</sup> Note that this uniform representation in a dictionary works because all backpropagators have the same codomain. The idea is that we replace what are now applications of linear functions with creation of a dictionary containing one key-value (function-argument) pair, and to replace addition of values in  $c$  with taking the union of dictionaries, where arguments for common keys are added together.

<sup>3</sup>This is the intuition; it will not go through precisely as planned, but something similar will.

*Initial Staged object.* More concretely, we want to replace the  $c$  in  $\mathbf{D}_c^1[\mathbb{R}] = (\mathbb{R}, \mathbb{R} \multimap c)$  with ‘Staged  $c$ ’ (our “dictionary”), which we define as follows:

$$\text{Staged } c = (c, \text{Map } (\mathbb{R} \multimap \text{Staged } c) \mathbb{R})$$

This type can represent both literal  $c$  values (necessary for the one-hot vectors returned by the input backpropagators created in `Interleave1`) and staged (delayed) calls to linear functions. We use `Map` to denote a standard (persistent) tree-map as found in every functional language. The intuitive semantics of a value  $(x, \{f_1 \mapsto a_1, f_2 \mapsto a_2\}) : \text{Staged } c$  is its *resolution*  $x + f_1 a_1 + f_2 a_2 : c$ .

To be able to replace  $c$  with `Staged  $c$`  in  $\mathbf{D}_c^1$ , we must support all operations that we perform on  $c$  also on `Staged  $c$` . We implement them as follows:

- $\underline{0} : c$  becomes simply  $0_{\text{Staged}} := (\underline{0}, \{\}) : \text{Staged } c$ .
- $(+) : c \rightarrow c \rightarrow c$  becomes  $(+_{\text{Staged}})$ , adding  $c$  values using  $(+)$  and taking the union of the two `Maps`. **Here we apply linear factoring:** if the two `Maps` both have a value for the same key (i.e. we have two staged invocations to the same linear function  $f$ ), the resulting map will have *one* value for that same key  $f$ : the sum of the arguments stored in the two separate `Maps`. For example:

$$(c_1, \{f_1 \mapsto a_1, f_2 \mapsto a_2\}) +_{\text{Staged}} (c_2, \{f_2 \mapsto a_3\}) = (c_1 + c_2, \{f_1 \mapsto a_1, f_2 \mapsto a_2 + a_3\})$$

- The one-hot  $c$  values created in the backpropagators from `Interleave1` are stored in the  $c$  component of `Staged  $c$` .
- An application  $f x$  of a backpropagator  $f : \mathbb{R} \multimap c$  to an argument  $x : \mathbb{R}$  now gets replaced with `SCall  $f x := (\underline{0}, \{f \mapsto x\}) : \text{Staged } c$` . This occurs in  $\mathbf{D}_c^1[\text{op}(\dots)]$  and in `Deinterleave1`.

What is missing from this list is how to “resolve” the final `Staged  $c$`  value produced by the derivative computation down to a plain  $c$  value—we need this at the end of the wrapper. This resolve algorithm will need to call functions in the `Staged  $c$`  object in the correct order, ensuring that we only invoke a backpropagator when we are sure that we have collected all calls to it in the `Map`. For example, in the example at the beginning of this section,  `$f_4 1$`  returns  $(\underline{0}, \{f_2 \mapsto 1, f_3 \mapsto 2\})$ . At this point, “resolving  $f_3$ ” means calling  $f_3$  at 2, observing the return value  $(\underline{0}, \{f_2 \mapsto 8, f_1 \mapsto 10\})$ , and adding it to the remainder (i.e. without the  $f_3$  entry) of the previous `Staged  $c$`  object to get  $(\underline{0}, \{f_2 \mapsto 9, f_1 \mapsto 10\})$ .

But as we observed above, the choice of which function to invoke first is vital to the complexity of the reverse AD algorithm: if we chose  $f_2$  first instead of  $f_3$ , the later call to  $f_3$  would produce another call to  $f_2$ , forcing us to evaluate  $f_2$  twice—something that we must avoid. There is currently no information in a `Staged  $c$`  object from which we can deduce the correct order of invocation, so we need something extra.

There is another problem with the current definition of `Staged  $c$` : it contains a `Map` keyed by functions, meaning that we need equality—actually, even an ordering—on functions! This is nonsense in general. Fortunately, both problems can be tackled with the same solution.

*Resolve order.* The backpropagators that occur in the derivative program (as produced by  $\mathbf{D}_c^1$  from Fig. 6) are not just arbitrary functions. Indeed, taking the target type  $c$  of the input backpropagators to be equal to the input type  $\sigma$  of the original program (of type  $\sigma \rightarrow \tau$ ), as we do in `Wrap1` in Fig. 7, all backpropagators in the derivative program have one of the following three forms:

- (1)  $(\underline{\lambda}(z : \mathbb{R}). t)$  where  $t$  is a tuple (of type  $\sigma$ ) filled with zero scalars except for one position, where it places  $z$ ; we call such tuples *one-hot tuples*. These backpropagators result from `Interleave $\sigma$ 1` (Fig. 7) after trivial beta-reduction of the intermediate linear functions.
- (2)  $(\underline{\lambda}(z : \mathbb{R}). \underline{0})$  occurs as the backpropagator of a scalar constant  $r$ . Note that since this  $\underline{0}$  is of type  $\sigma$ , operationally it is equivalent to a tuple filled completely with zero scalars.

- (3)  $(\underline{\lambda}(z : \mathbb{R}). d_1 (\partial_1 op(x_1, \dots, x_n)(z)) + \dots + d_n (\partial_n op(x_1, \dots, x_n)(z)))$  for an  $op \in \text{Op}_n$  where  $d_1, \dots, d_n$  are other linear backpropagators: these occur as the backpropagators generated for primitive operations.

**Insight:** Hence, we observe that all other backpropagators that a backpropagator  $f$  will ever call are contained in its closure, and were hence created (at runtime of the derivative program) before  $f$  itself was created. Therefore, if we give all backpropagators at runtime sequentially incrementing integer IDs, where a  $\underline{\lambda}$  allocated later in time gets a higher ID, we obtain that a called backpropagator always has a lower ID than the backpropagator it was called from.

This provides an answer to the question of in what order to resolve backpropagators: if they have sequentially increasing IDs, just start with the one with the largest ID! After all, any calls to other backpropagators that it produces in the returned Staged  $c$  value will have lower IDs, and so cannot be functions that we have already resolved (i.e. called) before. And as promised, giving backpropagators IDs also solves the issue of using functions as keys in a Map: we can just use the (integer) ID as the Map key, which is perfectly valid and efficient.

With this knowledge, we rewrite Staged  $c$  and SCall to the following:

$$\begin{aligned} \text{Staged } c &= (c, \text{Map Int } (\mathbb{R} \multimap \text{Staged } c, \mathbb{R})) \\ \text{SCall} &: (\text{Int}, \mathbb{R} \multimap \text{Staged } c) \rightarrow \mathbb{R} \multimap \text{Staged } c \\ \text{SCall } (i, f) \ x &= (\underline{0}, \{i \mapsto (f, x)\}) \end{aligned}$$

We call the second component of a Staged  $c$  value, which has type  $\text{Map Int } (\mathbb{R} \multimap \text{Staged } c, \mathbb{R})$ , the *staging map*, after its function to stage (linear) function calls.

The only thing that remains is to actually generate the IDs for the backpropagators at runtime; this we do using an ID generation monad (a state monad with a state of type Int). The resulting new program transformation, modified from Figs. 6 and 7, is shown in Fig. 8.

*New program transformation.* In Fig. 8, the term transformation now produces a term in the ID generation monad ( $\text{Int} \rightarrow (-, \text{Int})$ ); therefore, all functions in the original program will also need to run in the same monad. This gives the second change in the type transformation (aside from  $\mathbf{D}_c^2[\mathbb{R}]$ , which now tags backpropagators with an ID):  $\mathbf{D}_c^2[\sigma \rightarrow \tau]$  now produces a monadic function type instead of a plain function type.

On the term level, notice that the backpropagator for primitive operations (in  $\mathbf{D}_c^2[op(\dots)]$ ) now no longer calls  $d_1, \dots, d_n$  (the backpropagators of the arguments to the operation) directly, but instead registers the calls as pairs of function and argument in the Staged  $c$  returned by the backpropagator. The  $\cup$  in the definition of  $(+_{\text{Staged}})$  refers to map union including linear factoring; for example:

$$\{i_1 \mapsto (f_1, a_1), i_2 \mapsto (f_2, a_2)\} \cup \{i_2 \mapsto (f_2, a_3)\} = \{i_1 \mapsto (f_1, a_1), i_2 \mapsto (f_2, a_2 + a_3)\}$$

Note that the transformation assigns consistent IDs to backpropagators: it will never occur that two staging maps have an entry with the same key (ID) but with a different function in the value.

In the wrapper,  $\text{Interleave}^2$  is lifted into the monad and generates IDs for scalar backpropagators;  $\text{Deinterleave}^2$  is essentially unchanged. The initial backpropagator provided to  $\text{Interleave}^2$  in  $\text{Wrap}^2$ , which was  $\text{id} : \sigma \multimap \sigma$  in Fig. 7, has now become  $\text{SCotan} : \sigma \multimap \text{Staged } \sigma$ , which injects a cotangent into a Staged  $c$  object.  $\text{Interleave}^2$  will “split” this function up into individual  $\mathbb{R} \multimap \text{Staged } \sigma$  backpropagators for each of the individual scalars in  $\sigma$ .

At the end of the wrapper, we apply the insight that we had earlier: by resolving (calling and eliminating) the backpropagators in the final Staged  $c$  returned by the differentiated program in order from the highest ID to the lowest ID, we ensure that every backpropagator is called at most

**On types:**

$$\mathbf{D}_c^2[\mathbb{R}] = (\mathbb{R}, (\text{Int}, \mathbb{R} \multimap \text{Staged } c)) \quad \mathbf{D}_c^2[()] = () \quad \mathbf{D}_c^2[(\sigma, \tau)] = (\mathbf{D}_c^2[\sigma], \mathbf{D}_c^2[\tau])$$

$$\mathbf{D}_c^2[\sigma \rightarrow \tau] = \mathbf{D}_c^2[\sigma] \rightarrow \text{Int} \rightarrow (\mathbf{D}_c^2[\tau], \text{Int}) \quad \mathbf{D}_c^2[\text{Int}] = \text{Int}$$

**On terms:**

If  $\Gamma \vdash t : \tau$  then  $\mathbf{D}_c^2[\Gamma] \vdash \mathbf{D}_c^2[t] : \text{Int} \rightarrow (\mathbf{D}_c^2[\tau], \text{Int})$

$\mathbf{D}_c^2[x : \tau] = \lambda i. (x : \mathbf{D}_c^2[\tau], i)$

$\mathbf{D}_c^2[(s, t)] = \lambda i. \text{let } (x, i') = \mathbf{D}_c^2[s] \text{ } i \text{ in let } (y, i'') = \mathbf{D}_c^2[t] \text{ } i' \text{ in } ((x, y), i'')$

$\mathbf{D}_c^2[\text{let } x : \tau = s \text{ in } t] = \lambda i. \text{let } (x : \mathbf{D}_c^2[\tau], i') = \mathbf{D}_c^2[s] \text{ } i \text{ in } \mathbf{D}_c^2[t] \text{ } i'$

*etc.*

$\mathbf{D}_c^2[r] = \lambda i. ((r, (i, \underline{\lambda}(z : \mathbb{R}). 0_{\text{Staged}})), i + 1)$

$\mathbf{D}_c^2[\text{op}(t_1, \dots, t_n)] =$

$\lambda i. \text{let } ((x_1, d_1), i_1) = \mathbf{D}_c^2[t_1] \text{ } i \text{ in } \dots \text{ in let } ((x_n, d_n), i_n) = \mathbf{D}_c^2[t_n] \text{ } i_{n-1}$   
 $\text{in } ((\text{op}(x_1, \dots, x_n), (i_n, \underline{\lambda}(z : \mathbb{R}). \text{SCall } d_1 (\partial_1 \text{op}(x_1, \dots, x_n)(z)) +_{\text{Staged}} \dots +_{\text{Staged}}$   
 $\text{SCall } d_n (\partial_n \text{op}(x_1, \dots, x_n)(z))))$   
 $, i_n + 1)$

**Changed wrapper:**

$\text{Wrap}^2 : (\sigma \rightarrow \tau) \rightsquigarrow (\sigma \rightarrow (\tau, \tau \multimap \sigma))$

$\text{Wrap}^2[\lambda(x : \sigma). t] = \lambda(x : \sigma). \text{let } (x : \mathbf{D}_\sigma^2[\sigma], i) = \text{Interleave}_\sigma^2(x, \text{SCotan } 0)$   
 $\text{in let } (y, d) = \text{Deinterleave}_\tau^2(\text{fst } (\mathbf{D}_\sigma^2[t] \text{ } i))$   
 $\text{in } (y, \underline{\lambda}(z : \tau). \text{SResolve } (d \ z)) \quad \text{— see main text for SResolve}$

$\text{Interleave}_{\tau}^2 : \forall c. (\tau, \tau \multimap \text{Staged } c) \rightarrow \text{Int} \rightarrow (\mathbf{D}_c^2[\tau], \text{Int})$

$\text{Interleave}_{\mathbb{R}}^2 = \lambda(x, d). \lambda i. ((x, (i, d)), i + 1)$

$\text{Interleave}_{()}^2 = \lambda((), d). \lambda i. ((), i)$

$\text{Interleave}_{(\sigma, \tau)}^2 = \lambda((x, y), d). \lambda i. \text{let } (x', i') = \text{Interleave}_\sigma^2(x, \underline{\lambda}(z : \sigma). d \ (z, \underline{0})) \text{ } i$   
 $\text{in let } (y', i'') = \text{Interleave}_\tau^2(y, \underline{\lambda}(z : \tau). d \ (\underline{0}, z)) \text{ } i'$   
 $\text{in } ((x', y'), i'')$

$\text{Interleave}_{\text{Int}}^2 = \lambda(n, d). \lambda i. (n, i)$

$\text{Deinterleave}_\tau^2$  gets type  $\forall c. \mathbf{D}_c^2[\tau] \rightarrow (\tau, \tau \multimap \text{Staged } c)$  and ignores the new  $\text{Int}$  in  $\mathbf{D}_c^2[\mathbb{R}]$ .  $\underline{0}$  changes to  $0_{\text{Staged}}$  and  $(+)$  changes to  $(+_{\text{Staged}})$ .

**Staged interface:**

$\text{Staged } c = (c, \text{Map Int } (\mathbb{R} \multimap \text{Staged } c, \mathbb{R}))$

$0_{\text{Staged}} : \text{Staged } c \quad (+_{\text{Staged}}) : \text{Staged } c \rightarrow \text{Staged } c \rightarrow \text{Staged } c$

$0_{\text{Staged}} = (\underline{0}, \{\}) \quad (c, m) +_{\text{Staged}} (c', m') = (c + c', m \cup m') \quad \text{— with linear factoring}$

$\text{SCotan} : c \multimap \text{Staged } c \quad \text{SCall} : (\text{Int}, \mathbb{R} \multimap \text{Staged } c) \rightarrow \mathbb{R} \multimap \text{Staged } c$

$\text{SCotan } c = (c, \{\}) \quad \text{SCall } (i, f) \ x = (\underline{0}, \{i \mapsto (f, x)\})$

Fig. 8. The monadically transformed code transformation (from Fig. 4 to Fig. 5 plus Staged operations), based on Fig. 6. Parts of  $\mathbf{D}_c^2$  and  $\text{Interleave}^2$  that were simply lifted to monadic code are set in grey.

once. This is done in the following function:

```
SResolve (c : σ, m : Map Int (ℝ → Staged σ, ℝ)) :=
  if m is empty then c
  else let i = highest key in m
        in let (f, a) = lookup i in m
        in let m' = delete i from m
        in SResolve (f a +Staged (c, m'))
```

The three operations on  $m$  are standard logarithmic-complexity tree-map operations.

#### 4.1 Remaining complexity challenges

We have gained a lot with the function call staging so far: where the naive algorithm from Section 3 easily ran into exponential blowup of computation time if the results of primitive operations were used in multiple places, the updated algorithm from Fig. 8 completely solves this issue. For example, the program of Fig. 2 now results in the call graph displayed on the right: each backpropagator is called exactly once. However, some other complexity problems still remain.<sup>4</sup>



In general, for a reverse AD algorithm to have the right complexity, we want the produced derivative program  $P'$  to compute the gradient of the original program  $P$  at a given input  $x$  with runtime only a constant factor times the runtime of  $P$  itself on  $x$ —and this constant factor should work for all programs  $P$ . However, as-is, this requirement is untenable:  $P = \text{id} : \tau \rightarrow \tau$  always takes constant time whereas its gradient program must at the very least construct the value of  $P$ 's full (non-zero) gradient, which might be large (indeed, its size is  $\text{size}(x)$ ). Hence, we require that:

$$\exists c > 0. \forall P \in \text{Programs}(\sigma \rightarrow \tau). \forall x : \sigma, dy : \tau. \\ \text{cost}(\text{snd}(\text{Wrap}[P] x) dy) \leq c \cdot (\text{cost}(P x) + \text{size}(x))$$

where  $\text{cost}(E)$  is the time taken to evaluate  $E$  to normal form, and  $\text{size}(x)$  is the time taken to read all of  $x$  sequentially.

So, what is  $\text{cost}(\text{snd}(\text{Wrap}[P] x) dy)$ ? First, the primal pass  $(\text{Wrap}[P] x)$  consists of interleaving, running the differentiated program, and deinterleaving.

- Interleave<sup>2</sup> itself runs in  $O(\text{size}(x))$ . (The backpropagators it creates are more expensive, but those are not called just yet.)
- For the differentiated program,  $\mathbf{D}_\sigma^2[P]$ , we can see that in all cases of the transformation  $\mathbf{D}_c^2$ , the right-hand side does the work that  $P$  would have done, plus threading of the next ID to generate, as well as creation of backpropagators. Since this additional work is a constant amount per program construct,  $\mathbf{D}_\sigma^2[P]$  runs in  $O(\text{cost}(P x))$ .
- Deinterleave<sup>2</sup> runs in  $O(\text{size}(P x))$ , i.e. the size of the program output; this is certainly in  $O(\text{cost}(P x) + \text{size}(x))$  but likely much less.

Summarising, the primal pass as a whole runs in  $O(\text{cost}(P x) + \text{size}(x))$ , which is as required.

Then, the dual pass  $(f dy)$ , where  $f$  is the linear function returned by  $\text{Wrap}^2$ ) first calls the backpropagator returned by  $\text{Deinterleave}^2$  on the output cotangent, and then passes the result through  $\text{SResolve}$  to produce the final gradient. Let  $t$  be the function body of  $P$  (i.e.  $P = \lambda(x : \sigma). t$ ).

<sup>4</sup>This section does not provide a proof that  $\text{Wrap}^2$  does *not* have the correct complexity; rather, it argues that the expected complexity analysis does not go through. The same complexity analysis *will* go through for  $\text{Wrap}^3$  after the improvements of Sections 6 and 7.



- Because the number of scalars in the output is potentially as large as  $O(\text{cost}(P x) + \text{size}(x))$ , the backpropagator returned by  $\text{Deinterleave}^2$  is only allowed to perform a constant-time operation for each scalar. However, looking back at Fig. 7, we see that this function calls all scalar backpropagators contained in the result of  $\mathbf{D}_\sigma^2[t]$  once, and adds the results using  $(+\text{Staged})$ . Assuming that the scalar backpropagators run in constant time (we will cover this point later), we are left with the many uses of  $(+\text{Staged})$ ; if these are constant-time, we are still within our complexity budget. However:

**Problem:**  $(+\text{Staged})$  (see Fig. 8) is not constant-time: it adds values of type  $c$  and takes the union of staging maps, both of which may be large.

- Afterwards, we use  $\text{SResolve}$  on the resulting Staged  $\sigma$  to call every scalar backpropagator in the program (created in  $\mathbf{D}_\sigma^2[r]$ ,  $\mathbf{D}_\sigma^2[op(\dots)]$  and  $\text{Interleave}^2$ ) at most once; this is accomplished using three  $\text{Map}$  operations and one call to  $(+\text{Staged})$  per backpropagator. However, each of the scalar backpropagators corresponds to either a constant-time operation<sup>5</sup> in the original program  $P$  or to a scalar in the input  $x$ ; therefore, in order to stay within the time budget of  $O(\text{cost}(P x) + \text{size}(x))$ , we are only allowed a constant-time overhead per backpropagator here. Since  $(+\text{Staged})$  was covered already, we are left with:

**Problem:** the  $\text{Map}$  operations in  $\text{SResolve}$  are not constant-time.

- While we have arranged to invoke each scalar backpropagator at most once, we still need those backpropagators to individually run in constant-time too: our time budget is  $O(\text{cost}(P x) + \text{size}(x))$ , but there could be  $O(\text{cost}(P x) + \text{size}(x))$  distinct backpropagators. Recall from earlier that we have three kinds of scalar backpropagators:

- (1)  $(\lambda(z : \mathbb{R}). \text{SCotan}(0, \dots, 0, z, 0, \dots, 0))$  created in  $\text{Interleave}^2$  (with  $\text{SCotan}$  from  $\text{Wrap}^2$ ).

**Problem:** The interleave backpropagators take time  $O(\text{size}(x))$ , not  $O(1)$ .

- (2)  $(\lambda(z : \mathbb{R}). 0_{\text{Staged}})$  created in  $\mathbf{D}_\sigma^2[r]$ .

**Problem:**  $0_{\text{Staged}}$  takes time  $O(\text{size}(x))$ , not  $O(1)$ .

- (3)  $(\lambda(z : \mathbb{R}). \text{SCall } d_1 (\partial_1 op(\dots)(z)) +_{\text{Staged}} \dots +_{\text{Staged}} \text{SCall } d_n (\partial_n op(\dots)(z)))$  created in  $\mathbf{D}_\sigma^2[op(\dots)]$ . Assuming that primitive operation arity is bounded, we are allowed a constant-time operation for each operation argument.

**Problem:**  $\text{SCall}$  creates a  $\underline{0} : c$  and therefore runs in  $O(\text{size}(x))$ , not  $O(1)$ . (The problem with  $(+\text{Staged})$  was already covered above.)

Summarising again, we see that we have three categories of complexity problems to solve:

- (A) We are not allowed to perform monoid operations on  $c$  as often as we do. (This affects  $0_{\text{Staged}}$ ,  $(+\text{Staged})$  and  $\text{SCall}$ ). Our fix for this (in Section 5) will be to Cayley-transform the Staged  $c$  object, including the contained  $c$  value, turning zero into  $\text{id}$  and plus into  $(\circ)$  on the type  $\text{Staged } c \rightarrow \text{Staged } c$ .
- (B) The  $\text{Interleave}$  backpropagators that create a one-hot  $c$  value should avoid touching parts of  $c$  that they are zero on. After Cayley-transforming Staged  $c$  in Section 5, this problem becomes less pronounced: the backpropagators now *update* a Staged  $c$  value, where they can keep untouched subtrees of  $c$  fully as-is. However, the one-hot backpropagators will still do work proportional to the *depth* of the program input type  $c$ . We will turn this issue into a simple log-factor in the complexity in Section 6 by replacing the  $c$  in Staged  $c$  with a more efficient structure (namely,  $\text{Map } \text{Int } \mathbb{R}$ ). This log-factor can optionally be further eliminated using mutable arrays as described in Section 7.
- (C) The  $\text{Map}$  operations in  $\text{SResolve}$  are logarithmic in the size of the staging map. Like in the previous point, mutable arrays (Section 7) can eliminate this final log-factor in the complexity.

<sup>5</sup> Assuming primitive operations all have bounded arity and are constant-time. A more precise analysis, omitted here, lifts these restrictions—as long as the gradient of a primitive operation can be computed in the same time as the original.

From the analysis above, we can conclude that after we have solved each of these issues, the algorithm attains the correct complexity for reverse AD.

## 5 CAYLEY-TRANSFORMING THE COTANGENT COLLECTOR

The classical “difference list” trick in functional programming, originally designed to improve the performance of repeated application of the list-append operation [Hughes 1986], is an instance of a more general theorem: any monoid  $(M, 0, +)$  is isomorphic to the submonoid of  $(M \rightarrow M, \text{id}, \circ)$  containing only the functions  $\lambda m'. m + m'$  for all  $m \in M$ . In other words, the map that sends  $m \in M$  to  $(\lambda m'. m + m') \in M \rightarrow M$  is a monoid homomorphism, and it has a left-inverse:  $\lambda f. f 0$ .

In the original application on lists, the intent of moving from  $[\tau]$  to  $[\tau] \rightarrow [\tau]$  (an action that we call *Cayley-transforming* the  $[\tau]$  type) was to ensure that the list-append operations are consistently associated to the right. In our case, however, the primary remaining complexity issues are not due to operator associativity, but instead because our monoid has very expensive  $0$  and  $+$  operations (namely,  $0_{\text{Staged}}$  and  $(+_{\text{Staged}})$ ). If we Cayley-transform Staged  $c$ , i.e. if we replace Staged  $c$  with  $\text{Staged } c \rightarrow \text{Staged } c$ , all occurrences of  $0_{\text{Staged}}$  in the code transformation turn into  $\text{id}$  and all occurrences of  $(+_{\text{Staged}})$  turn into  $(\circ)$ . Since  $\text{id}$  is a constant and the composition of two functions can be constructed in constant time, this makes the monoid operations on the codomain of backpropagators (which now becomes  $\text{Staged } c \rightarrow \text{Staged } c$ ) constant-time.

Hence, all non-trivial work with Staged  $c$  objects that we still perform is limited to: 1. the single  $0_{\text{Staged}}$  value that the full composition is in the end applied to (to undo the Cayley-transform), and 2. the implementation of the non-monoid operations on Staged  $c$ : SCall, SCotan and SResolve. We do not have to worry about one single zero of type  $c$ , hence we focus only on SCall, SCotan and SResolve, which get the following updated types after the Cayley-transform:<sup>6</sup> (the changed parts are shown in red)

$$\begin{aligned} \text{SCall} & : (\text{Int}, \mathbb{R} \multimap (\text{Staged } c \rightarrow \text{Staged } c)) \rightarrow \mathbb{R} \multimap (\text{Staged } c \rightarrow \text{Staged } c) \\ \text{SCotan} & : c \multimap (\text{Staged } c \rightarrow \text{Staged } c) \\ \text{SResolve} & : (\text{Staged } c \rightarrow \text{Staged } c) \multimap c \end{aligned}$$

The definition of Staged  $c$  itself also gets changed accordingly:

$$\text{Staged } c = (c, \text{Map Int } (\mathbb{R} \multimap (\text{Staged } c \rightarrow \text{Staged } c), \mathbb{R}))$$

The new definition of SCall arises from simplifying the composition of the old SCall with  $(+_{\text{Staged}})$ :

$$\begin{aligned} \text{SCall } (i, f) \ x \ (c, m) & = (c, \text{if } i \notin m \text{ then insert } i \mapsto (f, x) \text{ into } m \\ & \quad \text{else update } m \text{ at } i \text{ with } (\lambda \_, x'). (f, x + x')) \end{aligned}$$

Note that  $(+_{\text{Staged}})$  has been eliminated, and we do not use  $(+)$  on  $c$  here anymore. For SCotan we have to modify the type further (Cayley-transforming its  $c$  argument as well) to lose all  $(+)$  operations on  $c$ :

$$\begin{aligned} \text{SCotan} & : (c \rightarrow c) \multimap (\text{Staged } c \rightarrow \text{Staged } c) \\ \text{SCotan } f \ (c, m) & = (f \ c, m) \end{aligned}$$

SResolve simply applies its argument to  $0_{\text{Staged}}$  (undoing the Cayley transform—this is now the only remaining  $0_{\text{Staged}}$ ) and runs the old code from Section 4, only changing  $f \ a \ +_{\text{Staged}} \ (c, m')$  to  $f \ a \ (c, m')$  on the last line:  $f$  from the Map now has type  $\mathbb{R} \multimap (\text{Staged } c \rightarrow \text{Staged } c)$ .

<sup>6</sup>Despite the fact that we forbade it in Section 3.1, we are putting function types on both sides of a  $\multimap$ -arrow here. The monoid structure here is the one from the Cayley transform (i.e. with  $\text{id}$  and  $(\circ)$ ). Notice that this monoid structure is indeed the one we want in this context: the “sum” (composition) of two values of type  $(\text{Staged } c \rightarrow \text{Staged } c)$  corresponds with the sum (with  $(+_{\text{Staged}})$ ) of the Staged  $c$  values that they represent. (This is the Cayley isomorphism described above.)

**On types:**

$$\mathbf{D}_c^3[\mathbb{R}] = (\mathbb{R}, (\text{Int}, \mathbb{R} \multimap (\text{Staged } c \rightarrow \text{Staged } c))) \quad \mathbf{D}_c^3[()] = () \quad \mathbf{D}_c^3[(\sigma, \tau)] = (\mathbf{D}_c^3[\sigma], \mathbf{D}_c^3[\tau])$$

$$\mathbf{D}_c^3[\sigma \rightarrow \tau] = \mathbf{D}_c^3[\sigma] \rightarrow \text{Int} \rightarrow (\mathbf{D}_c^3[\tau], \text{Int}) \quad \mathbf{D}_c^3[\text{Int}] = \text{Int}$$

**On terms:**

$$\text{If } \Gamma \vdash t : \tau \text{ then } \mathbf{D}_c^3[\Gamma] \vdash \mathbf{D}_c^3[t] : \text{Int} \rightarrow (\mathbf{D}_c^3[\tau], \text{Int})$$

Same as  $\mathbf{D}_c^3$ , except with ‘id’ in place of  $0_{\text{Staged}}$  and ‘o’ in place of  $(+_{\text{Staged}})$ .

**Changed wrapper:**

$$\text{Wrap}^3 : (\sigma \rightarrow \tau) \rightsquigarrow (\sigma \rightarrow (\tau, \tau \multimap \sigma))$$

$$\text{Wrap}^3[\lambda(x : \sigma). t] = \lambda(x : \sigma). \text{let } (x : \mathbf{D}_\sigma^3[\sigma], i) = \text{Interleave}_\sigma^3(x, \text{SCotan } 0) \\ \text{in let } (y, d) = \text{Deinterleave}_\tau^3(\text{fst } (\mathbf{D}_\sigma^3[t] i)) \\ \text{in } (y, \underline{\lambda}(z : \tau). \text{SResolve } (d z))$$

$$\text{Interleave}_{\tau}^3 : \forall c. (\tau, (\tau \rightarrow \tau) \multimap (\text{Staged } c \rightarrow \text{Staged } c)) \rightarrow \text{Int} \rightarrow (\mathbf{D}_c^3[\tau], \text{Int})$$

$$\text{Interleave}_{\mathbb{R}}^3 = \lambda(x, d). \lambda i. ((x, (i, \underline{\lambda}(z : \mathbb{R}). d (\lambda(a : \mathbb{R}). z + a))), \\ , i + 1)$$

$$\text{Interleave}_{()}^3 = \lambda((), d). \lambda i. ((), i)$$

$$\text{Interleave}_{(\sigma, \tau)}^3 = \lambda((x, y), d). \lambda i.$$

$$\text{let } (x', i') = \text{Interleave}_\sigma^3(x, \lambda(f : \sigma \rightarrow \sigma). d (\lambda((v, w) : (\sigma, \tau)). (f v, w))) i \\ \text{in let } (y', i'') = \text{Interleave}_\tau^3(y, \lambda(f : \tau \rightarrow \tau). d (\lambda((v, w) : (\sigma, \tau)). (v, f w))) i' \\ \text{in } ((x', y'), i'')$$

$$\text{Interleave}_{\text{Int}}^3 = \lambda(n, d). \lambda i. (n, i)$$

$$\text{Deinterleave}_\tau^3 : \forall c. \mathbf{D}_c^3[\tau] \rightarrow (\tau, \tau \multimap (\text{Staged } c \rightarrow \text{Staged } c))$$

(Same as  $\text{Deinterleave}^2$  in Fig. 8, except with id and  $(\circ)$  in place of  $0_{\text{Staged}}$  and  $(+_{\text{Staged}})$ )

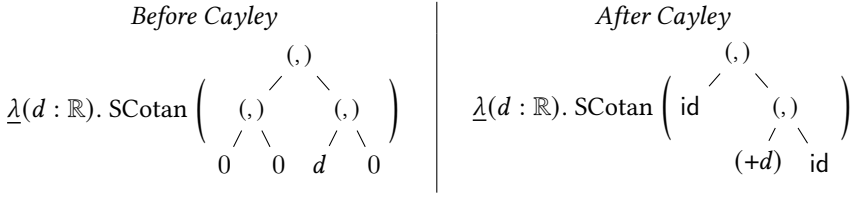
Fig. 9. The Cayley-transformed code transformation, based on Fig. 8. Grey parts are unchanged.

## 5.1 Code transformation

The new code transformation is shown in Fig. 9. Aside from the changes to types and to the target monoid of the backpropagators, the only additional change is in  $\text{Interleave}^3$ , which is adapted to accommodate the additional Cayley-transform to the  $c$  argument of  $\text{SCotan}$ . Note that the backpropagators in  $\text{Interleave}^3$  do not create any  $\underline{0}$  values for untouched parts of the collected cotangent of type  $c$ , as promised, and that the new type of  $\text{SCotan}$  has indeed eliminated all uses of  $(+)$  on  $c$ , not just moved them around.

## 5.2 Remaining complexity challenges

In Section 4.1, we pinpointed the three remaining complexity issues with the reverse AD algorithm after function call staging: costly monoid operations on  $c$ , costly one-hot backpropagators from  $\text{Interleave}$ , and logarithmic  $\text{Map}$  operations in  $\text{SResolve}$ . The first issue has been solved by Cayley-transforming  $\text{Staged } c$ : only  $\underline{0} : c$  is still used, and that only once (in the new  $\text{SResolve}$ ). For the second issue, although performance of the one-hot backpropagators has improved in most cases, it is still unsatisfactory; for example, given the input type  $\sigma = ((\mathbb{R}, \text{Int}), (\mathbb{R}, \mathbb{R}))$ , the backpropagator for the second scalar looks as follows before and after the Cayley transform:



This yields complexity logarithmic in the size of the input if the input is balanced, but can degrade to linear in the size of the input in the worst case—which is no better than the previous version. We will make these backpropagators properly logarithmic in the size of the input in Section 6, after which one can remove the final log-factors from the algorithm’s complexity by introducing mutable arrays as in Section 7.

## 6 KEEPING JUST THE SCALARS: EFFICIENT GRADIENT UPDATES

The codomain of the backpropagators is currently  $\text{Staged } c \rightarrow \text{Staged } c$ , with  $\text{Staged } c$  defined as:

$$\text{Staged } c = (c, \text{Map Int } (\mathbb{R} \multimap (\text{Staged } c \rightarrow \text{Staged } c), \mathbb{R}))$$

The final cotangent of the input to the program is collected in the first component of the pair, of type  $c$ . This collector is initialised with  $\underline{0} : c$  in  $0_{\text{Staged}}$ , and added to by the one-hot input backpropagators from `Interleave`, called in `SResolve`. The task of these input backpropagators is to add the cotangent (of type  $\mathbb{R}$ ) that they receive in their argument, to a particular scalar in the collector.

Hence, all we need of  $c$  in  $\text{Staged } c$  is really the collection of its scalars: the rest simply came from  $\underline{0} : c$  and is never changed.<sup>7</sup> Furthermore, the reason why the one-hot input backpropagators currently do not finish in logarithmic time is that  $c$  may not be a balanced tree of its scalars. But if we are interested only in the scalars anyway, we can *make* the collector balanced—by replacing it with  $\text{Map Int } \mathbb{R}$ :

$$\text{Staged } c = (\text{Map Int } \mathbb{R}, \text{Map Int } (\mathbb{R} \multimap (\text{Staged } c \rightarrow \text{Staged } c), \mathbb{R}))$$

`Interleave` changes to number all the scalars in the input with distinct IDs (for example with the same IDs as their corresponding input backpropagators, but this is not required); the cotangent of the input scalar with ID  $i$  is stored in the `Map` at key  $i$ . The input backpropagators can then modify the correct scalar in the collector (now of type  $\text{Map Int } \mathbb{R}$ ) in time logarithmic in the size of the input. To be able to construct the final gradient from this collection of just its scalars, `Interleaveτ` can additionally build a *reconstruction* function of type  $(\text{Int} \rightarrow \mathbb{R}) \rightarrow \tau$ , which we pass a function that looks up the ID in the final collector `Map` to compute the actual gradient value.<sup>8</sup>

*Complexity.* Now that we have fixed (in Section 5) the first complexity problem identified in Section 4.1 (expensive monoid operations) and reduced the second (expensive input backpropagators) to a logarithmic overhead over the original program, we have reached the point where we satisfy the complexity requirement stated in Section 4.1 apart from log-factors. More precisely, `Wrap[P]` computes the gradient of  $P$  at  $x$  not in time  $O(\text{cost}(P\ x) + \text{size}(x))$  but in time  $O((\text{cost}(P\ x) + \text{size}(x)) \log(\text{cost}(P\ x) + \text{size}(x)))$ . If we are okay with logarithmic overhead, we can stop here: the algorithm is already very close to efficient. However, if we wish to strictly conform to the required complexity, we need to make the input backpropagators and `Map` operations in `SResolve` constant-time; we do this using mutable arrays in Section 7.

<sup>7</sup>If  $c$  contains coproducts (sum types), this  $\underline{0} : c$  becomes dependent on the actual input to the program, copying the structure from there.

<sup>8</sup>For an example implementation of this idea, see the `IArray`  $\mathbb{R} \rightarrow \tau$  in the return type of `Interleave4` in Fig. 12 in Appendix A.

## 7 USING MUTABLE ARRAYS TO SHAVE OFF LOG FACTORS

The analysis in Section 4.1 showed that after Cayley-transform in Section 5, the strict complexity requirements are met if we make the input backpropagators constant-time and make SResolve not do non-constant extra work over the actual backpropagators that it must call. Luckily, in both cases the only component that is not constant-time is the interaction with one of the Maps in Staged  $c$ :

$$\text{Staged } c = (\text{Map Int } \mathbb{R}, \text{Map Int } (\mathbb{R} \multimap (\text{Staged } c \rightarrow \text{Staged } c), \mathbb{R}))$$

The input backpropagators perform (logarithmic-time) updates to the first Map (the cotangent collector), and SResolve reads, deletes and updates entries in the second Map (the staging map for recording delayed backpropagator calls). Both of these Maps are keyed by increasing, consecutive integers starting from 0, and are thus ideal candidates to be replaced by an array:

$$\text{Staged } c = (\text{Array } \mathbb{R}, \text{Array } (\mathbb{R} \multimap (\text{Staged } c \xrightarrow{R} \text{Staged } c), \mathbb{R}))$$

To allocate an array, one must know how large it should be. Fortunately, at the time when we allocate the initial Staged  $c$  value using  $0_{\text{Staged}}$  in SResolve, the primal pass has already been executed and we know (from the output ID of Interleave) how many input scalars there are, and (from the output ID of the transformed program) how many backpropagators there are. Hence, the size of these arrays is indeed known when they are allocated.

To get any complexity improvements from replacing a Map with an Array (indeed, to not pessimise the algorithm completely!), the write operations to the arrays need to be done *mutably*. These write operations occur in two places: in the updater functions produced by backpropagators ( $\text{Staged } c \rightarrow \text{Staged } c$ ) and in SResolve. Hence, in these two places we need an effectful function type; options include a resource-linear function type, written  $\xrightarrow{R}$  above (as e.g. available in Rust<sup>9</sup> and Haskell [Bernardy et al. 2018]) and a monad for local side-effects such as the ST monad in Haskell [Launchbury and Jones 1994] (where one would get  $\text{Staged } c \rightarrow \text{ST } s ()$  instead). Our implementation (Section 10) uses resource-linear types; this approach is described in more detail in Appendix A.

### 7.1 Complexity

We now satisfy all the requirements of the analysis in Section 4.1, and hence have the correct complexity for reverse AD. In particular, let  $I$  denote the size of the input and  $T$  the runtime of the original program. We can observe the following:

- The number of operations performed by  $\mathbf{D}_c^3[t]$  (with the improvements from Sections 6 and 7) is only a constant factor times the number of operations performed by  $t$ , and hence in  $O(T)$ . This was already observed for  $\mathbf{D}_c^2[t]$  in Section 4.1, and still holds.
- The number of backpropagators created while executing  $\mathbf{D}_c^3[t]$  is clearly also in  $O(T)$ .
- The number of operations performed in any one backpropagator is constant. This is new, and only true because  $\text{id}$  (replacing  $0_{\text{Staged}}$ ),  $(\circ)$  (replacing  $(+_{\text{Staged}})$ ), SCotan (with a constant-time mutable array updater as argument) and SCall are now all constant-time.
- Hence, because every backpropagator is invoked at most once, and because the overhead of SResolve is constant per invoked backpropagator, the amount of work performed by calling the top-level input backpropagator is again in  $O(T)$ .
- Finally, the (non-constant-time) extra work performed in Wrap<sup>3</sup> is interleaving ( $O(I)$ ), deinterleaving ( $O(\text{size of output})$  and hence  $O(T + I)$ ), resolving ( $O(T)$ ) and reconstructing the gradient from the scalars in the Array  $\mathbb{R}$  in Staged  $c$  ( $O(I)$ ); all this work is in  $O(T + I)$ .

<sup>9</sup><https://www.rust-lang.org>

Hence, calling  $\text{Wrap}^3[t]$  with an argument and calling its returned top-level derivative once takes time  $O(T + I)$ , i.e. at most proportional to the runtime of calling  $t$  with the same argument, plus the size of the argument itself. This is indeed the correct time complexity for an efficient reverse AD algorithm, as discussed in Section 4.1.

## 8 WAS IT TAPING ALL ALONG?

In this section we first apply one more optimisation to our algorithm to improve its complexity by a constant factor (Section 8.1). Next, we show that defunctionalising the backpropagators (Section 8.2) essentially reduces the technique to classical taping approaches (Section 8.3).

### 8.1 Dropping the cotangent collection array

Recall that the final transformation of Section 7 used two mutable arrays threaded through the backpropagators in the Staged  $c$  pair: a cotangent collection array of type  $\text{Array } \mathbb{R}$  and a backpropagator call staging array of type  $\text{Array } (\mathbb{R} \multimap (\text{Staged } c \multimap \text{Staged } c), \mathbb{R})$ . The first array is modified by  $\text{Interleave}_{\mathbb{R}}$ , and the second by  $\text{SCall}$ . No other functions modify these arrays.

Looking at the function of  $\text{Interleave}_{\mathbb{R}}$  in the algorithm, all it does is produce input backpropagators with some ID  $i$ , which act by adding their argument to index  $i$  in the cotangent collection array. This means that if  $(c, m)$  is the input to  $\text{SResolve}$  for which the recursion terminates, we have  $c[i] = \text{snd}(m[i])$  for all  $i$  for which  $c[i]$  is defined. Therefore, the cotangent collection array is actually unnecessary: its information is directly readable from the backpropagator staging array.

With this knowledge, we can instead use  $\text{Staged } c = \text{Array } (\mathbb{R} \multimap (\text{Staged } c \multimap \text{Staged } c), \mathbb{R})$  as our definition. The reconstruction functions of Section 6 simply take the second projection of the corresponding array element.

### 8.2 Defunctionalisation of backpropagators

In the core code transformation ( $\mathbf{D}_c$ , excluding the wrapper), all backpropagators are (now) of type  $\mathbb{R} \multimap (\text{Staged } c \multimap \text{Staged } c)$ , and, as observed earlier in Section 4, these backpropagators come in only a limited number of forms:

- (1) the input backpropagators, as created in  $\text{Interleave}_{\mathbb{R}}$ , reduced to  $(\underline{\lambda}(z : \mathbb{R}). \text{id})$  in Section 8.1;
- (2)  $(\underline{\lambda}(z : \mathbb{R}). \text{id})$ , as created in  $\mathbf{D}_c[r]$  for scalar constants  $r$ ;
- (3)  $(\underline{\lambda}(z : \mathbb{R}). \text{SCall } d_1 (\partial_1 \text{op}(x_1, \dots, x_n)(z)) \circ \dots \circ \text{SCall } d_n (\partial_n \text{op}(x_1, \dots, x_n)(z)))$ , as created in  $\mathbf{D}_c[\text{op}(x_1, \dots, x_n)]$  for primitive operations  $\text{op}$ .

Furthermore, the information contained in an operator backpropagator of form (3) can actually be described without reference to the value of its argument  $z$ : because our operators return a single scalar (as opposed to e.g. a vector), we have  $\frac{\partial f(\text{op}(x_1, \dots, x_n))}{\partial x_i} = \frac{\partial f(u)}{\partial u} \cdot \frac{\partial \text{op}(x_1, \dots, x_n)}{\partial x_i}$ , which can also be written as  $\partial_i \text{op}(x_1, \dots, x_n)(z) = z \cdot \partial_i \text{op}(x_1, \dots, x_n)(1)$ .

Hence, we can defunctionalise [Reynolds 1998] and change all occurrences of the type  $\mathbb{R} \multimap (\text{Staged } c \multimap \text{Staged } c)$  to  $\text{Contrib}$ , where  $\text{Contrib} = [(\mathbb{R}, (\text{Int}, \text{Contrib}))]$ : a list of triples of a scalar, an integer ID, and a recursive  $\text{Contrib}$  structure. (The recursive  $\text{Contrib}$  structures are interpreted as having sharing as encoded by their IDs, similarly to how the references to existing backpropagators in the closures of operator backpropagators (3) already had sharing.) The meaning of  $[(a_1, (i_1, cb_1)), \dots, (a_n, (i_n, cb_n))]$  of type  $\text{Contrib}$  would then be:

$$\underline{\lambda}(z : \mathbb{R}). \text{SCall } (i_1, cb_1) (z \cdot a_1) \circ \dots \circ \text{SCall } (i_n, cb_n) (z \cdot a_n)$$

For example, suppose we differentiate the following simple program:

$$\lambda(x, y). \text{let } z = x + y \text{ in } x \cdot z$$



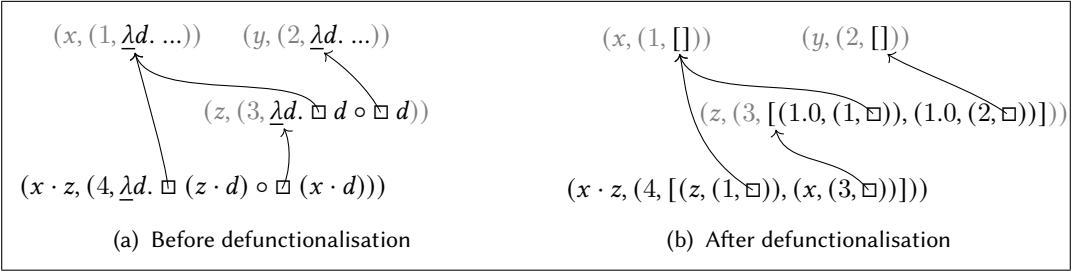


Fig. 10. The sharing structure before and after defunctionalisation. SCall is elided here; in Fig. 10a, the backpropagator calls are depicted as if they are still normal calls. Boxes (□) are the same in-memory value as the value their arrow points to; two boxes pointing to the same value indicates that this value is *shared*: referenced in two places.

using the final algorithm of Section 7. The return value from the  $\mathbf{D}_{\mathbb{R}}$ -transformed code (when applied to the output from  $\text{Interleave}_{\mathbb{R}}$ ) has the sharing structure shown in Fig. 10a. This shows how the backpropagators refer to each other in their closures.

If we perform the type replacement and the defunctionalisation,  $\text{Interleave}$  simplifies and SCall disappears, backpropagators of forms (1) and (2) become [] (the empty list) and those of form (3) become:

$$[(\text{fst } d_1, (\text{snd } d_1, \partial_1 \text{op}(x_1, \dots, x_n)(1))), \dots, (\text{fst } d_n, (\text{snd } d_n, \partial_n \text{op}(x_1, \dots, x_n)(1)))]$$

SResolve then interprets a list of such  $(i, (cb, a))$  by iterating over the list and for each such triple, replacing  $(cb', a')$  at index  $i$  in the staging array with  $(cb, a' + a)$ .

### 8.3 Was it taping all along?

After the improvements from Sections 8.1 and 8.2, what previously was a tree of (staged) calls to backpropagator functions is now a tree of Contrib values with attached IDs<sup>10</sup> that are interpreted by SResolve. This interpretation (eventually) writes the Contrib value with ID  $i$  to index  $i$  in the staging array (possibly multiple times), and furthermore accumulates argument cotangents in the second component of the pairs in the staging array. While the argument cotangents must be accumulated in reverse order of program execution (indeed, that is the whole point of *reverse* AD), the mapping from ID to Contrib value can be fully known in the forward pass: the partial derivatives of operators,  $\partial_i \text{op}(x_1, \dots, x_n)(1)$ , can be computed in the forward pass already.

This means that if we change the ID generation monad that the differentiated code already lives in (which is a state monad with a single Int as state) to additionally carry the staging array, and furthermore change the monad to thread its state through resource-linearly,<sup>11</sup> we can already compute the Contrib lists and write them to the array in the forward pass. All that SResolve then has to do is loop over the array in reverse order (as it already does) and add cotangent contributions to the correct positions in the array according to the Contrib lists that it finds there.

At this point, there is no meaningful difference any more between this algorithm and what is classically known as taping: we have a tape (the staging array) that we write the performed operations to in the forward pass (automatically growing the array as necessary)—although the tape entries are the already-differentiated operations in this case, and not the original ones. In this way, we have related the naive version of dual-numbers reverse AD, which admits neat correctness proofs, to the classical, very imperative approach to reverse AD based on taping, which is used in industry-standard implementations of reverse AD (e.g. PyTorch [Paszke et al. 2017]).

<sup>10</sup>Note that we now have  $\mathbf{D}[\mathbb{R}] = (\mathbb{R}, (\text{Int}, \text{Contrib}))$ , the integer being the ID of the Contrib value.

<sup>11</sup>This is possible and results in a linear variant of standard Haskell monads, as described in [Bernardy et al. 2018].

## 9 EXTENDING THE SOURCE LANGUAGE

The source language (Fig. 4) that the algorithm discussed so far works on, is a higher-order functional language including product types and primitive operations on scalars. However, dual-numbers reverse AD generalises to much richer languages in a very natural way, because most of the interesting work happens in the scalar primitive operations. The efficiency of the algorithm is independent of the language constructs in the source language. Indeed, in the forward pass, the code transformation is fully structure-preserving outside of the scalar constant and primitive operation cases; and in the reverse pass (in SResolve), all program structure is forgotten anyway, because the computation is flattened to a linear sequence of primitive operations on scalars.

*(Mutual) recursion.* For example, we can allow recursive functions in our source language by adding the syntax **letrec**  $f : \sigma \rightarrow \tau = \lambda(x : \sigma). s$  **in**  $t$ . The code transformation  $\mathbf{D}^i$  for all  $i$  then treats **letrec** exactly the same as **let**—note that the only syntactic difference between **letrec** and **let** is the scoping of  $f$ —and the algorithm remains both correct and efficient.

*Coproducts.* To support dynamic control flow (necessary to make recursion useful), we can easily add coproducts to the source language. First add coproducts to the syntax for types ( $\sigma, \tau ::= \dots \mid \sigma + \tau$ ) both in the source language and in the target language, and add constructors and eliminators to all term languages (both linear and non-linear):

$$s, t ::= \dots \mid \text{inl}(t) \mid \text{inr}(t) \mid \mathbf{case\ s\ of\ \{inl}(x) \rightarrow t_1; \text{inr}(y) \rightarrow t_2\}}$$

where  $x$  and  $y$  are in scope in  $t_1$  and  $t_2$ . Then the type and code transformations extend in the unique structure-preserving manner:

$$\mathbf{D}_c^1[\sigma + \tau] = \mathbf{D}_c^1[\sigma] + \mathbf{D}_c^1[\tau]$$

$$\mathbf{D}_c^1[\text{inl}(t)] = \text{inl}(\mathbf{D}_c^1[t]) \quad \mathbf{D}_c^1[\text{inr}(t)] = \text{inr}(\mathbf{D}_c^1[t])$$

$$\mathbf{D}_c^1[\mathbf{case\ s\ of\ \{inl}(x) \rightarrow t_1; \text{inr}(x) \rightarrow t_2\}}] = \mathbf{case\ D}_c^1[s] \mathbf{of\ \{inl}(x) \rightarrow \mathbf{D}_c^1[t_1]; \text{inr}(x) \rightarrow \mathbf{D}_c^1[t_2\}}$$

The type transformation stays unchanged when moving to  $\mathbf{D}_c^3$ , and the only change for the term definitions is to transition to monadic code in  $\mathbf{D}_c^2$ . Lifting a computation to monadic code is a well-understood process. The corresponding cases in Interleave and Deinterleave are the only reasonable definitions that type-check.

*Polymorphic and (mutually) recursive types.* In Haskell one can define (mutually) recursive data types e.g. as follows:

$$\mathbf{data\ } T_1 \alpha = C_1 \alpha (T_2 \alpha) \mid C_2 \mathbb{R}$$

$$\mathbf{data\ } T_2 \alpha = C_3 \text{Int} (T_1 \alpha) (T_2 \alpha).$$

If the user has defined some data types, then we can allow these data types in the code transformation. We add new *data type declarations* that simply apply  $\mathbf{D}_c^1[-]$  to all parameter types of all constructors:

$$\mathbf{data\ } DT_1 \alpha = DC_1 \alpha (DT_2 \alpha) \mid DC_2 (\mathbb{R}, \mathbb{R} \rightarrow c)$$

$$\mathbf{data\ } DT_2 \alpha = DC_3 \text{Int} (DT_1 \alpha) (DT_2 \alpha).$$

and we add one rule for each data type that simply maps:<sup>12</sup>

$$\mathbf{D}_c^1[T_1 \tau] = DT_1 \mathbf{D}_c^1[\tau] \quad \mathbf{D}_c^1[T_2 \tau] = DT_2 \mathbf{D}_c^1[\tau].$$

<sup>12</sup>As declaring new data types is inconvenient in Template Haskell, our current implementation only handles recursive data types that do not contain explicit scalar values. As we can pass all required scalar types by instantiating their type parameters with a type containing  $\mathbb{R}$ , this is not a real restriction.

	TH	ad	TH / ad
scalar mult.	0.146 $\mu$ s $\pm$ 0.000	0.536 $\mu$ s $\pm$ 0.002	0.27
dot product	2.21 $\mu$ s $\pm$ 0.10	2.07 $\mu$ s $\pm$ 0.06	$\approx$ 1.1
sum-mat-vec	2.05 $\mu$ s $\pm$ 0.14	1.32 $\mu$ s $\pm$ 0.05	$\approx$ 1.5
rotate_vec_by_quat	8.77 $\mu$ s $\pm$ 0.01	6.13 $\mu$ s $\pm$ 0.02	$\approx$ 1.43

Table 1. Benchmark results of Section 7+ Sections 8.1 and 8.2 versus ad-4.5. The ‘TH’ and ‘ad’ columns indicate runtimes on one machine for our implementation and the ad library, respectively. The last column shows the ratio between the previous two columns. We give the size of the largest side of `criterion`’s 95% confidence interval. Setup: GHC 9.2.2 on Linux, Intel i9-10900K CPU. Benchmarks are single-threaded.

Furthermore, for plain type variables, we set  $\mathbf{D}_c^1[\alpha] = \alpha$  (we apply  $\mathbf{D}_c^1$  where  $\alpha$  is *instantiated*, not where it is *used*).

The code transformation on terms is completely analogous to a combination of coproducts (given above in this section, where we take care to match up constructors as one would expect:  $C_i$  gets sent to  $DC_i$ ) and products (given already in Fig. 6). The wrapper also changes analogously: Interleave and Deinterleave get clauses for Interleave<sub>( $T_i \tau$ )</sub> and Deinterleave<sub>( $T_i \tau$ )</sub>.

Finally, we note that with the mentioned additional rule that  $\mathbf{D}_c^1[\alpha] = \alpha$ , polymorphic functions can also be differentiated transparently, similarly to how the above handles polymorphic data types.

## 10 IMPLEMENTATION

To show the practicality of our method, we provide a prototype implementation<sup>13</sup> of the resulting algorithm of Section 7, together with the improvements from Sections 8.1 and 8.2, that differentiates a sizeable fragment of Haskell98 including recursive types (reinterpreted as a strict, call-by-value language) using Template Haskell. We realise the mutable arrays of Section 7 using resource-linearly typed arrays of Linear Haskell [Bernardy et al. 2018] (Appendix A), which are similar in intent, though not identical in design, to those of the Rust language. The implementation does not incorporate the changes given in Section 8.3 that transform the algorithm into classical taping, but it does include support for recursive functions and user-defined data types as described in Section 9.

Template Haskell [Sheard and Jones 2002] is a built-in metaprogramming facility in GHC Haskell that (roughly) allows the programmer to write a Haskell function that takes a block of user-written Haskell code, do whatever it wants with the AST of that code, and finally splice the result back into the user’s program. The resulting code is still type-checked as usual. The AST transformation that we implement is, of course, differentiation.

*Benchmarks.* To check that our implementation has reasonable performance in practice, we benchmark (in `bench/Main.hs`) against Kmett’s ad library [Kmett and contributors 2021] (version 4.5) on a few basic functions. These functions are:

- A single scalar multiplication of type `(Double, Double) -> Double`;
- Dot product of type `([Double], [Double]) -> Double`;
- Matrix–vector multiplication, then sum: of type `([[Double]], [Double]) -> Double`;
- The `rotate_vec_by_quat` example from [Krawiec et al. 2022] of type `(Vec3 Double, Quaternion Double) -> Vec3 Double`, with data `Vec3 s = Vec3 s s s` and data `Quaternion s = Quaternion s s s s`.

The last case has a non-trivial return type.

<sup>13</sup>The code is available at <https://github.com/tomsmeding/ad-dualrev-th>.

The benchmark results are shown in Table 1. The benchmarks are timed using the `criterion`<sup>14</sup> library. To get statistically significant results, we measure how the timings scale with increasing values of  $n$ :

- Scalar multiplication and `rotate_vec_by_quat` are simply differentiated  $n$  times;
- Dot product is performed on lists of length  $n$ ;
- Matrix multiplication is done for a matrix and vector of size  $\sqrt{n}$ , to get linear scaling in  $n$ .

By the results in Table 1, we see that for less trivial programs, our implementation is around 50% slower than the highly-optimised `ad` library. Because our code is simply a proof-of-concept, we conclude from this that the algorithm described in this paper indeed admits a work-efficient implementation.

## 11 ORIGINS OF DUAL-NUMBERS REVERSE AD, RELATIONSHIP WITH VECTORISED FORWARD AD AND OTHER RELATED WORK

The literature about automatic differentiation spans many decades and academic subcommunities (scientific computing, machine learning and—most recently—programming languages). Important early references are [Linnainmaa 1970; Speelpenning 1980; Wengert 1964]. Good surveys can be found in [Baydin et al. 2017; Margossian 2019]. In the rest of this section, we focus on the more recent literature that studies AD from a programming languages (PL) point of view, to extend the scope of our discussion of Section 8.

### 11.1 Theoretical foundations for our algorithm

The first mention that we know of the naive dual-numbers reverse mode AD algorithm that we analyse in this paper is [Pearlmutter and Siskind 2008, page 12], where it is quickly dismissed before a different technique is pursued. The algorithm is first thoroughly studied by [Brunel et al. 2020] using operational semantics and [Huot et al. 2020, Section 6] using denotational semantics. [Brunel et al. 2020] introduces the key idea that underlies the efficient implementation of our paper: the linear factoring rule, stating that a term  $f x + f y$ , with  $f$  a linear function, may be reduced to  $f (x + y)$ . We build on their use of this rule as a tool in a complexity proof to make it a suitable basis for a performant implementation. We achieve this by noting that it can be efficiently implemented using our Staged  $c$  datastructure combined with runtime numbering of backpropagators and next observing that we obtain a performant implementation if we apply a Cayley transformation (and use mutable arrays to shave off log-factors).

[Mazza and Pagani 2021] extends the work of [Brunel et al. 2020] to apply to a language with term recursion. Similarly, [Nunes and Vákár 2022] extends the work of [Huot et al. 2020] to apply to recursive types, thus giving a correctness proof for initial dual-numbers reverse AD transformation of Fig. 6 when applied to idealised Haskell98.

### 11.2 Vectorised forward AD

Furthermore, there are strong parallels with the derivation in [Krawiec et al. 2022]. Like the present paper, they give a sequence of steps that optimise a simple algorithm to an efficient implementation—but the starting algorithm is vectorised forward AD (VFAD) instead of backpropagator-based dual-numbers reverse AD (DNRAD). In our notation, their initial type transformation does not have  $\mathbf{D}_c^1[\mathbb{R}] = (\mathbb{R}, \mathbb{R} \multimap c)$ , but instead  $\mathbf{D}_c^1[\mathbb{R}] = (\mathbb{R}, c)$ . (As befits a dual-numbers algorithm, the rest of the type transformation is simply structurally recursive.)

Linear algebra tells us that the vector spaces  $\mathbb{R} \multimap c$  and  $c$  are isomorphic, and indeed inspection of the term transformations shows that both naive algorithms compute the same thing. Their

<sup>14</sup>By Bryan O’Sullivan: <https://hackage.haskell.org/package/criterion>

operational behaviour, on the other hand, is very different: the complexity problem with DNRAD is exponential blowup in the presence of sharing, whereas VFAD is “simply”  $n$  times too slow, where  $n$  is the number of scalars in the input.

But the first optimisation on VFAD, which defunctionalises the zero, one-hot, addition and scaling operations on the  $c$  tangent vector, introduces the same sharing-induced complexity problem as we have in naive DNRAD as payment for fixing the factor- $n$  overhead. The two algorithms are now on equal footing: we could defunctionalise the backpropagators in DNRAD just as easily.

Afterwards, VFAD is lifted to a combination (stack) of an ID-generation monad and a Writer monad. Each scalar result of a primitive operation gets an ID, and the Writer monad records for each ID (hence, scalar in the program) its defunctionalised tangent vector (i.e. an expression) in terms of other already-computed tangent vectors from the Writer record. These expressions correspond to our primitive operation backpropagators with calls replaced with SCall: where we replace calls with ID-tagged pairs of function and argument, VFAD replaces the usage of already-computed tangent vectors with scaled references to the IDs of those vectors. The choice in our SResolve of evaluation order from highest ID to lowest ID (Section 4) is encoded in VFAD’s definitions of *runDelta* and *eval*, which process the record back-to-front.

Finally, our Cayley-transform is encoded in the type of VFAD’s *eval* function, which interprets the defunctionalised operations on tangent vectors (including explicit sharing using the Writer log) into an actual tangent vector—the final gradient: its gradient return type is Cayley-transformed. Our final optimisation to mutable arrays to eliminate log-factors in the complexity is also mirrored in VFAD.

*Distributive law.* Under the isomorphism  $\mathbb{R} \multimap c \cong c$ , the type Staged  $c$  can be thought of as a type Expr  $c$  of ASTs of expressions (with sharing) of type  $c$ .<sup>15</sup> The linear factoring rule  $f\ x + f\ y \rightsquigarrow f\ (x + y)$  for a linear function  $f : \mathbb{R} \multimap c$  that rescales a vector  $v : c$  with a scalar then corresponds to the distributive law  $v \cdot x + v \cdot y \rightsquigarrow v \cdot (x + y)$ . This highlights the relationship between our work and that of [Shaikhha et al. 2019], which tries to (statically) optimise vectorised forward AD to reverse AD using precisely this distributive law. A key distinction is that we apply this law (in the form of the linear factoring rule) at runtime rather than compile time, allowing us to always achieve the complexity of reverse AD, rather than merely on some specific programs with straightforward control and data flow. The price we pay for this generality is a runtime overhead, similar to the usual overhead of taping.

### 11.3 Other PL literature about AD

*CHAD and category theory inspired AD.* Rather than interleaving backpropagators by pairing them with scalars in a type, we can also try to directly implement reverse AD as a structurally recursive code transformation that does not need a (de)interleaving wrapper. This is the approach taken by Elliott [Elliott 2018]. It pairs vectors (and values of other composite types) with a single composite backpropagator, rather than decomposing to the point where each scalar is paired with a mini-backpropagator like in our dual-numbers approach. The resulting algorithm is extended to source languages with function types in [Vákár 2021; Vákár and Smeding 2022; Vytiniotis et al. 2019] and to sum and (co)inductive types in [Nunes and Vákár 2021]. Like our dual-numbers reverse AD approach, the algorithm arises as a canonical structure-preserving functor on the syntax of a programming language. However, due to a different choice in target category (a Grothendieck construction of a linear  $\lambda$ -calculus for CHAD rather than the syntax of a plain  $\lambda$ -calculus for dual-numbers AD), the resulting algorithm looks very different.

<sup>15</sup>In [Krawiec et al. 2022], Expr  $c$  is called Delta.

*Approaches utilising non-local control flow.* Another category of approaches to AD recently taken by the PL community are those that rely on forms of non-local control flow such as delimited continuations [Wang and Rompf 2018] or effect handlers [de Vilhena and Pottier 2021; Sigal 2021]. These techniques are different in the sense that they generate code that is not purely functional. This use of non-local control flow makes it possible to achieve an efficient implementation of reverse AD that looks strikingly simple compared to alternative approaches. Where the CHAD approaches and our dual-numbers reverse AD approach both have to manually invert control flow at compile time by making use of continuations that get passed around, combined with smart staging of execution of those continuations in our case, this inversion of control can be deferred to run time by clever use of delimited control operators or effect handlers.



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## A MUTABLE ARRAYS USING RESOURCE-LINEAR TYPES

As announced in Section 7, a possible way of modelling efficient mutable arrays is using resource-linear types. This appendix illustrates how this looks when using the recent support for linear types in Haskell (Linear Haskell [Bernardy et al. 2018]), which is similar in intent, though not identical in design, to that of the Rust language.<sup>16</sup> For an introduction to Linear Haskell, we refer to the cited article.

By introducing resource-linearity into our type system, we now have three flavours of function arrows: regular functions ( $\rightarrow$ ), monoid-linear functions ( $\multimap$ ) and the new resource-linear functions, which we write as  $\multimap_R$ . Where regular functions used  $\lambda$  and monoid-linear functions used  $\underline{\lambda}$ , resource-linear functions use  $\underline{\lambda}_R$ . Both monoid-linearity and resource-linearity are essential for achieving the right time complexity in this paper, but in somewhat different ways: monoid-linearity shows that the optimisations that we performed up to this point preserve semantics, while resource-linearity shows that the transition to mutable arrays preserves referential transparency. In Haskell, resource-linearity must be explicitly annotated in the output program of the code transformation for GHC to accept the code, while monoid-linearity need not be reflected in the implementation. In a hypothetical implementation in OCaml, where mutable updates are allowed anywhere, neither form of linearity would need to be reflected.

For working with mutable arrays using resource-linearity, we use a standard interface taken from the `linear-base`<sup>17</sup> Haskell library. (For background information on implementation as well as usage idioms, we refer again to [Bernardy et al. 2018].) This library is written by the authors of Linear Haskell as a general standard library for usage of linear types in Haskell. The subset of the array interface that we use is listed and briefly explained in Fig. 11.<sup>18</sup> In the same figure, we also give a derived operation `modify`.

*Transformation using arrays.* As described in Section 7, we use arrays for two things: as replacement for the  $c$  value in a Staged  $c$  (in which the final cotangent is collected by invocation of the injector backpropagators, and which we had already replaced with `Map Int  $\mathbb{R}$`  in Section 6)—this solves problem (B) of Section 4.1—, and as a replacement for the staging `Map` in a Staged  $c$ —this solves problem (C). Hence, we change Staged  $c$  from:

$$(\text{Map Int } \mathbb{R}, \text{Map Int } (\mathbb{R} \multimap (\text{Staged } c \rightarrow \text{Staged } c), \mathbb{R}))$$

to:

$$(\text{Array } \mathbb{R}, \text{Array } (\mathbb{R} \multimap (\text{Staged } c \multimap_R \text{Staged } c), \mathbb{R})).$$

### A.1 Code transformation

Using this new type, we change the code transformation once more, this time from Fig. 9 to the version given in Fig. 12. While the transformation on types and terms simply gains resource-linearity on the arrows in scalar backpropagators, and thus does not materially change, some important changes occur in the Staged  $c$  interface (previously the Staged  $c$  interface) and the wrapper. Let us first look at the algorithm from the top, by starting with `Wrap`<sup>4</sup>; after understanding the high-level idea, we explain how the other components work.

In basis, `Wrap`<sup>4</sup> does the same as `Wrap`<sup>3</sup> from Fig. 9: interleave injector backpropagators with the input of type  $\sigma$ , execute the transformed function body using the interleaved input, and then

<sup>16</sup>In Rust, values may be implicitly dropped, making its version of linear types really *affine* instead of purely linear. Furthermore, Rust extends the system with a complex but convenient system of ownership and lifetime tracking.

<sup>17</sup><https://hackage.haskell.org/package/linear-base-0.2.0/docs/Data-Array-Mutable-Linear.html>

<sup>18</sup>In `linear-base`, the final  $\rightarrow \sigma$  in the type of `alloc` is instead  $\multimap !\sigma$ , but this is relevant only if the `Array  $\tau \multimap !\sigma$`  function closes over a linear value, which ours does not. Hence we choose the simpler presentation.

**Array interface:**

`alloc` :  $(length : \text{Int}) \rightarrow (initval : \tau) \rightarrow (\text{Array } \tau \xrightarrow{R} !\sigma) \rightarrow \sigma$

Allocate an array of the given *length* filled with *initval* in all positions, and pass it to the given function. Referential transparency is ensured because linearity guarantees that the  $\text{Array } \tau$  cannot be returned as part of the  $!\sigma$ .

`allocBeside` :  $(length : \text{Int}) \rightarrow (initval : \tau) \rightarrow (witness : \text{Array } \sigma) \xrightarrow{R} (\text{Array } \sigma, \text{Array } \tau)$

Using a “linearity witness”, allocate another array. In particular: `allocBeside l x w` returns two arrays: *w* and a newly allocated array of length *l* filled with *xes*.

`get` :  $(index : \text{Int}) \rightarrow \text{Array } \tau \xrightarrow{R} (!\tau, \text{Array } \tau)$

Note: The returned value of type  $\tau$  does not need to be used linearly. While this means that this interface is unsuitable for arrays of (mutable) arrays, we will not need those here.

`set` :  $(index : \text{Int}) \rightarrow (value : \tau) \rightarrow \text{Array } \tau \xrightarrow{R} \text{Array } \tau$

`size` :  $\text{Array } \tau \xrightarrow{R} (!\text{Int}, \text{Array } \tau)$

`dealloc` :  $\text{Array } \tau \xrightarrow{R} \text{Array } \sigma \xrightarrow{R} \text{Array } \sigma$

`dealloc a b` deallocates *a* and returns *b*. (This is `lseq` from `linear-base` with a more specific type.)

`freeze` :  $\text{Array } \tau \xrightarrow{R} !(\text{IArray } \tau)$

Permanently convert a mutable array to an immutable array. `IArray` corresponds to `Vector` in the Haskell `linear-base` API.

`@` :  $\text{IArray } \tau \rightarrow \text{Int} \rightarrow \tau$

Index an immutable array—no linearity required. (`(Data.Vector.!)` in Haskell.)

**Sequencing:**

**let**  $(x_1, \dots, x_n) = t_1$  **in**  $t_2$

Compute  $t_1$ , bind (and, if  $n > 1$ , destructure) its result to  $x_1, \dots, x_n$ , and finally compute  $t_2$ .

$t_1$  is allowed to consume linearly-bound values, as long as they are unused in  $t_2$ . In Haskell using GHC 9.2, this is written as case  $t_1$  of  $(x_1, \dots, x_n) \rightarrow t_2$ .

**Derived array operations**

`modify` :  $(index : \text{Int}) \rightarrow (\tau \rightarrow \tau) \rightarrow \text{Array } \tau \xrightarrow{R} \text{Array } \tau$

`modify i f =  $\lambda_R(a : \text{Array } \tau)$ . let  $(!x, a')$  = get i a in set i (f x) a'`

Modify the array at the given index.

Fig. 11. Features from Linear Haskell that we use, including a subset of `Data.Array.Mutable.Linear` from the Haskell library `linear-base`, as well as a linear `let`-binding. We write the Haskell type `Ur` (“unrestricted” or “of course”) as  $!$ , in types as well as in pattern matches.

deinterleave the result. However, because we (since Section 6) represent the final cotangent not directly as a value of type  $\sigma$  in a `Staged`  $\sigma$  but instead as an array of only the embedded scalars ( $\text{Array } \mathbb{R}$ ), some more work needs to be done.

Firstly, `Interleave4` (monadically) produces, in addition to the interleaved input, also a *reconstruction* function<sup>19</sup> of type  $\text{IArray } \mathbb{R} \rightarrow \sigma$ . This rebuilder takes an array with precisely as many scalars as were in the input, and produces a value of type  $\sigma$  with the structure (and discrete-typed values) of the input, but the scalars from the array. The mapping between locations in  $\sigma$  and indices in the array is the same as the numbering performed by `Interleave4`.

<sup>19</sup>Implementing the  $(\text{Int} \rightarrow \mathbb{R})$  in  $(\text{Int} \rightarrow \mathbb{R}) \rightarrow \tau$  from Section 6 as `IArray`  $\mathbb{R}$ .

**On types:**

$$\mathbf{D}_c^4[\mathbb{R}] = (\mathbb{R}, (\text{Int}, \mathbb{R} \multimap (\text{Staged } c \xrightarrow{\mathbb{R}} \text{Staged } c))) \quad \mathbf{D}_c^4[()] = () \quad \mathbf{D}_c^4[(\sigma, \tau)] = (\mathbf{D}_c^4[\sigma], \mathbf{D}_c^4[\tau])$$

$$\mathbf{D}_c^4[\sigma \rightarrow \tau] = \mathbf{D}_c^4[\sigma] \rightarrow \text{Int} \rightarrow (\mathbf{D}_c^4[\tau], \text{Int}) \quad \mathbf{D}_c^4[\text{Int}] = \text{Int}$$

**On terms:**

If  $\Gamma \vdash t : \tau$  then  $\mathbf{D}_c^4[\Gamma] \vdash \mathbf{D}_c^4[t] : \text{Int} \rightarrow (\mathbf{D}_c^4[\tau], \text{Int})$

Same as  $\mathbf{D}_c^3$ . ('id' now has a resource-linear type,  $\text{Staged } c \xrightarrow{\mathbb{R}} \text{Staged } c$ . Note further that the usages of `SCall` remain type-correct.)

**New Staged interface:**

`Staged c` = (Array  $\mathbb{R}$ , Array ( $\mathbb{R} \multimap (\text{Staged } c \xrightarrow{\mathbb{R}} \text{Staged } c)$ ),  $\mathbb{R}$ )

`SAlloc` : Int  $\rightarrow$  Int  $\rightarrow$  (Staged  $c \xrightarrow{\mathbb{R}} !c$ )  $\rightarrow$   $c$

`SAlloc`  $i_{\text{inp}} i_{\text{out}} f = \text{alloc } i_{\text{inp}} 0 (f \circ \text{allocBeside } i_{\text{out}} (\underline{\lambda}(z : \mathbb{R}). \text{id}, 0))$

`SCall` : (Int,  $\mathbb{R} \multimap (\text{Staged } c \xrightarrow{\mathbb{R}} \text{Staged } c)$ )  $\rightarrow$   $\mathbb{R} \multimap (\text{Staged } c \xrightarrow{\mathbb{R}} \text{Staged } c)$

`SCall` ( $i, f$ )  $a = \underline{\lambda}_R(c, m). (c, \text{modify } i (\underline{\lambda}(\_, a'). (f, a + a')) m)$

`SOneHot` : Int  $\rightarrow$   $\mathbb{R} \multimap (\text{Staged } c \xrightarrow{\mathbb{R}} \text{Staged } c)$

`SOneHot`  $i a = \underline{\lambda}_R(c, m). (\text{modify } i (\underline{\lambda}(a' : \mathbb{R}). a + a') c, m)$

`SResolve` : Int  $\rightarrow$  Staged  $c \xrightarrow{\mathbb{R}}$  Array  $\mathbb{R}$

`SResolve`  $i_{\text{out}} = \underline{\lambda}_R(c, m). \mathbf{let} (c, m) = \text{loop } (i_{\text{out}} - 1) (c, m) \mathbf{in} \text{dealloc } m c$   
 where `loop` : Int  $\rightarrow$  Staged  $c \xrightarrow{\mathbb{R}}$  Staged  $c$   
`loop` 0 =  $\underline{\lambda}_R(s : \text{Staged } c). s$   
`loop`  $i = \underline{\lambda}_R(c, m). \mathbf{let} (!f, a), m = \text{get } i m \mathbf{in} \text{loop } (i - 1) (f a (c, m))$

**Wrapper:**

`Interleave` $_{\tau}^4$  :  $\tau \rightarrow \text{Int} \rightarrow ((\mathbf{D}_c^4[\tau], \text{IArray } \mathbb{R} \rightarrow \tau), \text{Int})$

`Interleave` $_{\mathbb{R}}^4 = \underline{\lambda}x. \underline{\lambda}i. (((x, (i, \text{SOneHot } i)), \underline{\lambda}a. a @ i), i + 1)$

`Interleave` $_{()}^4 = \underline{\lambda}(). \underline{\lambda}i. ((((), \underline{\lambda}a. ()), i)$

`Interleave` $_{(\sigma, \tau)}^4 = \underline{\lambda}(x, y). \underline{\lambda}i. \mathbf{let} ((x', f_1), i') = \text{Interleave}_{\sigma}^4 x i$   
 $\mathbf{in} \mathbf{let} ((y', f_2), i'') = \text{Interleave}_{\tau}^4 y i'$   
 $\mathbf{in} (((x', y'), \underline{\lambda}a. (f_1 a, f_2 a)), i'')$

`Interleave` $_{\text{Int}}^4 = \underline{\lambda}n. \underline{\lambda}i. ((n, \underline{\lambda}a. n), i)$

`Interleave` $_{\sigma \rightarrow \tau}^4 = \text{not defined!}$

`Deinterleave` $_{\tau}^4 : \mathbf{D}_c^4[\tau] \rightarrow (\tau, \tau \multimap (\text{Staged } c \xrightarrow{\mathbb{R}} \text{Staged } c))$

Same as `Deinterleave` $^3$  except for types.

`Wrap` $^4 : (\sigma \rightarrow \tau) \rightsquigarrow (\sigma \rightarrow (\tau, \tau \multimap \sigma))$

`Wrap` $^4[\underline{\lambda}(x : \sigma). t] = \underline{\lambda}(x : \sigma).$

$\mathbf{let} ((x : \mathbf{D}_c^4[\sigma], \text{rebuild} : \text{IArray } \mathbb{R} \rightarrow \sigma), i) = \text{Interleave}_{\sigma}^4 x 0$

$\mathbf{in} \mathbf{let} (y', i') = \mathbf{D}_c^4[t] i$

$\mathbf{in} \mathbf{let} (y, d : \tau \multimap (\text{Staged } c \xrightarrow{\mathbb{R}} \text{Staged } c)) = \text{Deinterleave}_{\tau}^4 y'$

$\mathbf{in} (y, \underline{\lambda}(z : \tau). \text{rebuild } (\text{SAlloc } i i' (\text{freeze} \circ \text{SResolve } i' \circ d z)))$

Fig. 12. Code transformation plus wrapper using mutable arrays, modified from Fig. 9. Grey parts are unchanged.

Having  $x$ , *rebuild* and  $i$  (the next available ID), we execute the transformed term  $t$  monadically (with  $x$  in scope), resulting in an output  $y' : \mathbf{D}^4[\tau]$ . This output we deinterleave to  $y : \tau$  and  $d : \tau \multimap (\text{Staged } c \xrightarrow{R} \text{Staged } c)$ .

The final result then consists of the regular function result ( $y$ ) as well as the top-level derivative function of type  $\tau \multimap \sigma$ . This  $\tau$  we can pass to  $d$  to get a Staged  $c$  updater that (because of how Deinterleave<sup>4</sup> works) stages calls to the top-level backpropagators contained in  $y'$  in a Staged  $c$ . Assuming that we can pass  $d z$  an empty Staged  $c$ , we then use the new SResolve to propagate the cotangent contributions backwards, by invoking each backpropagator in turn in descending order of IDs. Like before in the Cayley version, those backpropagators update the state (now mutably) to record their own contributions to (i.e. invocations of) other backpropagators. As listed in Fig. 12, SResolve does not return the entire state but only the cotangent collection array of type  $\text{Array } \mathbb{R}$  (corresponding to the  $c$  value in a Staged  $c$  for the Cayley version in Fig. 9); the other array is deallocated before returning. Passing this  $\text{Array } \mathbb{R}$  to *freeze* gives us an unrestricted  $\text{IArray } \mathbb{R}$  containing the cotangents of all scalars in the top-level input.

At this point we have built  $z : \tau \vdash (\text{freeze} \circ \text{SResolve} \circ d z) : \text{Staged } c \xrightarrow{R} !(\text{IArray } \mathbb{R})$ , which we need to pass an empty Staged  $c$ . This we do using SAlloc from Fig. 12, which uses an idiom in Linear Haskell (also used in *alloc*): to allow code to work with a mutable data structure, enforce that said code takes the mutable data structure resource-linearly to an *unrestricted* return value; with that typing, the mutable data structure cannot escape through the return value, meaning that the mutation is invisible from outside, as required. The additional  $i$  and  $i'$  arguments to SAlloc are the sizes of the arrays to allocate: the cotangent array of type  $\text{Array } \mathbb{R}$  will be indexed by the IDs of the scalars in the interleaved input (which are in  $\{0, \dots, i - 1\}$ ), and the backpropagator staging array (of type  $\text{Array } (\mathbb{R} \multimap (\text{Staged } c \xrightarrow{R} \text{Staged } c), \mathbb{R})$ ) will be indexed by the IDs of all backpropagators created in both Interleave<sup>4</sup> and  $\mathbf{D}^4[t]$  (which are in  $\{0, \dots, i' - 1\}$ ). Hence,  $i$  and  $i'$  are suitable array sizes.

Finally, the  $\text{IArray } \mathbb{R}$  returned by *freeze* through SAlloc is passed to *rebuild* from Interleave<sup>4</sup> to put all scalar cotangents in the correct locations in the input; the resulting final cotangent is returned.

*Implementation of the components.* Having discussed the high-level sequence of operations, let us briefly discuss the implementation of the Staged  $c$  interface and (de)interleaving. In Interleave<sup>4</sup>, instead of passing structure information down in the form of a setter ( $(\tau \rightarrow \tau) \multimap (\text{Staged } c \rightarrow \text{Staged } c)$ ) like we did in Interleave<sup>3</sup> in Fig. 9, we build structure information up in the form of a getter ( $\text{IArray } \mathbb{R} \rightarrow \tau$ ). This results in a somewhat more compact presentation, but in some sense the same information is still communicated.

The program text of Deinterleave<sup>4</sup> is again unchanged, because it is agnostic about the codomain of the backpropagators (apart from being a monoid, in this case over  $\text{id}$  and  $(\circ)$ ).

On the Staged interface the transition to mutable arrays had a significant effect. The  $0_{\text{Staged}}$  created by SResolve in Section 5 is now created in SAlloc, which uses *alloc* to allocate the cotangent collection array of size  $i_{\text{inp}}$  filled with zeros, and then *allocBeside* to allocate the backpropagator staging array of size  $i_{\text{out}}$  filled with zero-backpropagators with an accumulated argument of zero. (Recall Fig. 11 for the types of *alloc* and *allocBeside*.)

S $\text{Call}$  has essentially the same type, but its implementation differs: instead of performing a logarithmic-complexity immutable Map update, we perform a constant-time mutable update on the backpropagator staging array. Note that, unlike in Section 5, there is no special case if  $i$  is not yet in the array, because unused positions are already filled with zeros.

SOneHot takes the place of SCotan, except we have specialised SOneHot with the knowledge that all relevant  $c \rightarrow c$  functions add a particular scalar to a particular index in the input, and that



these functions can hence be defunctionalised to a pair  $(\text{Int}, \mathbb{R})$ . The monoid-linearity here is in the real scalar, as it was before, hence the placement of the  $\rightarrow$ -arrow.

Finally, `SResolve` takes an additional `Int` argument that should contain the output ID of  $\mathbf{D}^4[t]$ , i.e. one more than the largest ID generated. `loop` then does what the original `SResolve` did directly, iterating over all IDs in descending order and applying the state updaters in the backpropagator staging array one-by-one to the state. After the loop is complete, the backpropagator staging array is deallocated and the cotangent collection array is returned, to be passed to `freeze` in `Wrap`<sup>4</sup>.