# Differentiate Everything with a Reversible Domain-Specific Language

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#### **Abstract**

This paper considers implementing automatic differentiation (AD) in a reversible embedded domain-specific language. We start by reviewing the limitations of traditional AD frameworks. To solve the issues in these frameworks, we developed an open source reversible eDSL NiLang in Julia that can differentiate a general reversible program while being compatible with Julia's ecosystem. It empowers users the flexibility to tradeoff time, space, and energy rather than caching data into a global tape. With examples, we show one can use reversible programming AD to obtain gradients and Hessians for a wide class of functions in scientific programming and machine learning, including elementary mathematical functions, sparse matrix operations and linear algebras. Manageable memory allocation makes it a good tool to differentiate GPU kernel functions. By benchmarking its performance in differentiating Bessel function, graph embedding, Gaussian mixture model and bundle adjustment, we demonstrate that the AD implemented in a reversible programming language can achieve state-of-the-art performance in both time and space. Finally, we will discuss the challenges that we face towards rigorous reversible programming, mainly from the instruction and hardware perspective.

#### 1 Introduction

Computing the gradients of a numeric model  $f: \mathbb{R}^m \to \mathbb{R}^n$  plays a crucial role in scientific computing. Consider a computing process

$$\mathbf{x}^{1} = f_{1}(\mathbf{x}^{0})$$

$$\mathbf{x}^{2} = f_{2}(\mathbf{x}^{1})$$

$$\dots$$

$$\mathbf{x}^{L} = f_{L}(\mathbf{x}^{L-1})$$

where  $x^0 \in R^m$ ,  $x^L \in R^n$ , L is the depth of computing. The Jacobian of this program is a  $n \times m$  matrix  $J_{ij} \equiv \frac{\partial x_i^L}{\partial x_j^0}$ , where  $x_j^0$  and  $x_i^L$  are single elements of inputs and outputs. Computing the Jacobian or part of the Jacobian automatically is what we called automatic differentiation (AD). It can be classified into three classes, the forward mode AD, the backward mode AD and the mixed mode AD. Hascoet and Pascual (2013) The forward mode AD computes the Jacobian matrix elements related to a single input using the chain rule  $\frac{\partial \mathbf{x}^k}{\partial x_j^0} = \frac{\partial \mathbf{x}^k}{\partial \mathbf{x}^{k-1}} \frac{\partial \mathbf{x}^{k-1}}{\partial x_j^0}$  with j the column index,

while a backward mode AD computes Jacobian matrix elements related to a single output using the chain rule in the reverse direction  $\frac{\partial \mathbf{x}_i^L}{\partial \mathbf{x}^{k-1}} = \frac{\partial \mathbf{x}_i^L}{\partial \mathbf{x}^k} \frac{\partial \mathbf{x}^k}{\partial \mathbf{x}^{k-1}}$  with i the row index. In variational applications where the loss function always outputs a scalar, the backward mode AD is preferred. However, implementing backward mode AD is harder than implementing its forward mode counterpart, because it requires propagating the gradients in the inverse direction of computing the loss. The backpropagation of gradients requires intermediate information of a program that includes

- 1. a representation of the computational process,
- 2. and variables required for computing gradients.

The computational process is often represented as a computational graph, which is a directed acyclic graph (DAG) representing the relationship between data and functions. Comparing with a general computer program, a computational graph does not mutate variables and does not have control flows. These limitations makes it a suitable representation to describe the data caching in the forward pass, at the cost of poor support to inplace operations. Most popular AD implementations in the market implements the computational graph AD by providing a finite set of primitive functions with predefined backward rules. In Pytorch Paszke *et al.* (2017) and Flux Innes *et al.* (2018), every variable has a tracker field. When applying a predefined primitive function on a variable, the variable's tracker field keeps track of this function as well as data needed in backpropagation. TensorFlow Abadi *et al.* (2015) also implements the computational graph, but it builds a static computational graph as a description of the program before actual computation happens. These frameworks sometimes fail to meet the diverse needs in research, for example, in physics research,

- 1. People need to differentiate over sparse matrix operations that are important for Hamiltonian engineering Hao Xie and Wang, like solving dominant eigenvalues and eigenvectors Golub and Van Loan (2012),
- 2. People need to backpropagate singular value decomposition (SVD) function and QR decomposition in tensor network algorithms to study the phase transition problem Golub and Van Loan (2012); Liao *et al.* (2019),
- 3. People need to differentiate over a quantum simulation where each quantum gate is an inplace function that changes the quantum register directly Luo *et al.* (2019).

Scientists put lots of effort to derive backward rules for these needs. In the backpropagation of dominant eigensolver Hao Xie and Wang, people managed to circumvent the sparse matrix issue by allowing users to provide the backward function for sparse matrix constructors. The backward rules for SVD and eigensolvers have been formulated in recent years Seeger *et al.* (2017); Wan and Zhang (2019); Hubig (2019). One can obtain the gradients correctly for both real numbers and complex numbers Wan and Zhang (2019) in most cases now. In variational quantum simulator Yao Luo *et al.* (2019), authors implemented a builtin cache free AD engine by utilizing the reversible nature of quantum computing. They derive and implement the backward rule for each type of quantum gate.

Some other tools like Tapenade Hascoet and Pascual (2013), ReverseDiff and Zygote Innes (2018); Innes *et al.* (2019) can provide program level reverse mode AD, that is differentiate a general program written by a user. However, these approaches have their own limitations too. In many practical applications, an elementary level differentiable program that might do billions of computations will cache intermediate results to a global storage. Frequent caching of data slows down the program significantly, and the memory will become a bottleneck as well. With these AD tools, it is still nearly impossible to automatically generate the backward rule for BLAS functions and sparse matrix operations with a performance comparable to the state-of-the-art.

We propose to differentiate a program by writting it reversiblly. So that the program can differentiate over inplace operations while can be traced backward with no extra effort. The overhead of reverse mode AD becomes the overhead of reversing a program, where the later has the advantage of efficient and controllable memory management. Reversibility has been used in reducing the memory allocations in machine learning models such as recurrent neural networks MacKay et al. (2018), Hyperparameter learning Maclaurin et al. (2015) and residual neural networks Behrmann et al. (2018). Our approach shares the same spirit, but from the programming level. There have been many prototypes of reversible languages like Janus Lutz (1986), R (not the popular one) Frank (1997), Erlang Lanese et al. (2018) and object-oriented

ROOPL Haulund (2017). In the past, the primary motivation of studying reversible programming is to support reversible devices Frank and Knight Jr (1999) like adiabatic complementary metal-oxide-semiconductor (CMOS) Koller and Athas (1992), molecular mechanical computing system Merkle et al. (2018) and superconducting system Likharev (1977); Semenov et al. (2003). Reversible computing are more energy-efficient from the perspective of information and entropy, or by the Landauer's principle Landauer (1961). After decades of efforts, reversible computing devices are very close to providing productivity now. As an example, adiabatic CMOS can be a better choice in a spacecraft Hänninen et al. (2014); DeBenedictis et al. (2017), where energy is more valuable than device itself. Reversible programming is interesting to software engineers too, because it is a powerful tool to schedule asynchronious events Jefferson (1985) and debug a program bidirectionally Boothe (2000). However, the field of reversible computing faces the difficulty of having not enough funding in recent decade Frank (2017). As a result, not many people studying AD know the marvelous designs in reversible computing. People have not connected it with automatic differentiation seriously, even though they have many similarities. We aim to break the information barrier between the machine learning community and the reversible programming community in our work and provide yet another strong motivation to develop reversible programming.

In this paper, we first introduce the language design of a reversible programming language and introduce our reversible eDSL NiLang in Sec. 2. In Sec. 3, we explain the implementation of automatic differentiation in this eDSL. In Sec. 4, we show several examples, and introduce the concept of memory oriented computational graph. In Sec. 5, we benchmark the performance of NiLang with other AD packages. In Sec. 6, we discuss several important issues, the time-space tradeoff, reversible instructions and hardware, and finally, an outlook to some open problems to be solved. In the appendix, we show the grammar of NiLang and other technical details.

## 2 Language design

#### 2.1 Introductions to reversible language design

In a modern programming language, functions are pushed to a global stack for scheduling. The memory layout of a function consists of input arguments, a function frame with information like the return address and saved memory segments, local variables, and working stack. After the call, the function clears run-time information, only stores the return value. In reversible programming, this kind of design is no longer the best practice. One can not discard input variables and local variables easily after a function call, since discarding information may ruin reversibility. For this reason, reversible functions are very different from irreversible ones from multiple perspectives.

## 2.1.1 Memory management

A distinct feature of reversible memory management is, the content of a variable must be known when it is deallocated. We denote the allocation of a zero emptied memory as  $x \leftarrow 0$ , and the corresponding deallocation as  $x \rightarrow 0$ . A variable x can be allocated and deallocated in a local scope, which is called an ancilla. It can also be pushed to a stack and used later with a pop statement. This stack is similar to a traditional stack, except it zero-clears the variable after pushing and presupposes that the variable being zero-cleared before popping.

Knowing the contents in the memory when deallocating is not easy. Hence Charles H. Bennett introduced the famous compute-copy-uncompute paradigm Bennett (1973) for reversible programming. To explain how it works, we introduce the memory oriented computational graph, as shown in Fig. 1. Notations are highly inspired by quantum circuit representations. A vertical line is a variable, and it can be used by multiple operations. Hence it is a hypergraph rather than a simple graph (i.e. the DAG). When a variable is used by a function, depending on whether its value is changed or not, we put a box or a dot at the line cross. In panel (a). The subprogram in dashed box X is executed on space  $x_{1:3}$  represents the computing stage. In the copying stage, the content in  $x_3$  is read out to a pre-emptied memory  $x_4$  through inplace add +=. Since this copy operation does not change contents of  $x_{1:3}$ , we can use the uncomputing operation  $\sim$ X to undo all the changes to these registers. Now we computing the result  $x_4$  without modifying the contents in  $x_{1:3}$ . If any of them is in a known state, it can be deallocated immediately. In panel (b), we can use the subprogram defined in (a) maked as Y to generate  $x_{5:n}$  without modifying the contents of variables  $x_{1:4}$ . It is

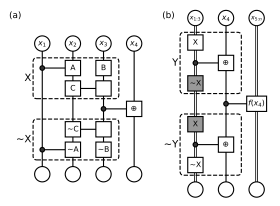


Figure 1: Two computational processes represented in memory oriented computational graph, where (a) is a subprogram in (b). In these graphs, a vertical single line represents one variable, a vertical double line represents multiple variables, and a parallel line represents a function. A dot at the cross represents a control parameter of a function and a box at the cross represents a mutable parameter of a function.

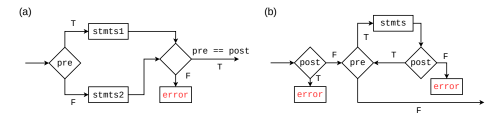


Figure 2: The flow chart for reversible (a) if statement and (b) while statement. "pre" and "post" represents precondition and postconditions respectively.

easy to see that although this uncompute-copy-uncompute design pattern can restore memories to known state, it has computational overhead. Both X and  $\sim X$  are executed twice in the program (b), which is not necessary. We can cancel a pair of X and  $\sim X$  (the gray boxes). By doing this, we are not allowed to deallocate the memory  $x_{1:3}$  during computing  $f(x_{5:n})$ . This is the famous time-space tradeoff that playing the central role in reversible programming. The trade off strategy will be discussed in detail in Sec. 6.1.

#### 2.1.2 Control flows

One can define reversible if, for and while statements in a slightly different way comparing with its irreversible counterpart. The reversible if statement is shown in Fig. 2 (a). Its condition statement contains two parts, a precondition and a postcondition. The precondition decides which branch to enter in the forward computing, while the postcondition decides which branch to enter in the backward computing. After executing the specific branch, the program checks the consistency between precondition and postcondition to make sure they are consistent. The reversible while statement in Fig. 2 (b) also has two condition fields. Before executing the condition expressions, the program preassumes the postcondition is false. After each iteration, the program asserts the postcondition to be true. In the reverse pass, we exchange the precondition and postcondition. The reversible for statement is similar to irreversible ones except that after executing the loop, the program checks the values of these variables to make sure they are not changed. In the reverse pass, we exchange start and stop and inverse the sign of step.

#### 2.1.3 Arithmetic instructions

Every arithmetic instruction has a unique inverse that can undo the changes.

- For logical operations,  $y \subseteq f(args...)$  is self reversible.
- For integer and floating point arithmetic operations, we treat y += f(args...) and y -= f(args...) as reversible to each other. Here f can be an arbitrary pure function such as identity, \*, / and ^. Here, we ignore the effect of floating point rounding errors for the moment and discuss in detail in Sec. 6.2.
- For logartihmic number and Tropical number algebra Speyer and Sturmfels (2009), y \*= f(args...) and y /= f(args...) as reversible to each other. The irreversibility of the zero element (-∞) in the Tropical number multiplication is ignored here.

Besides the above two types of operations, SWAP operation that exchanges the contents in two memory spaces is also widely used in reversible computing systems.

#### 2.2 Differetiable Reversible eDSL: NiLang

We develop an embedded domain-specific language (eDSL) NiLang in Julia language Bezanson et al. (2012, 2017) that implements reversible programming. Perumalla (2013); Frank (2017). One can write reversible control flows, instructions, and memory managements inside this macro. We choose Julia as the host language for multiple purposes. Julia is a popular language for scientific programming. Its meta-programming and its package for pattern matching MLStyle MLS allow us to define an eDSL conveniently. Its type inference and just in time compiling can remove most overheads introduced in our eDSL, providing the state-of-the-art performance. Most importantly, its multiple-dispatch provides the polymorphism that will be used in our AD engine. The main feature of NiLang is contained in a single macro @i that compiles a reversible function. The allowed statements in this eDSL are shown in Appendix A. We can use macroexpand to show the compiling a reversible function to the native Julia function.

```
julia> using NiLangCore, MacroTools
julia> MacroTools.prettify(@macroexpand @i function f(x, y)
            SWAP(x, y)
       end)
quote
    $(Expr(:meta, :doc))
    function $(Expr(:where, :(f(x, y))))
        dove = wrap_tuple(SWAP(x, y))
        x = dove[1]
        y = dove[2]
         (x, y)
    if NiLangCore._typeof(f) != _typeof(~f)
  function $(Expr(:where, :((hummingbird::_typeof(~f))(x, y))))
             toad = wrap_tuple((~SWAP)(x, y))
             x = toad[1]
             y = toad[2]
             (x. v)
        end
    end
end
```

Here, the version of NiLang is v0.4.0. Macro @i generates two functions that reversible to each other f and  $\sim f$ .  $\sim f$  is an callable of type Inv{typeof(f)}, where the type parameter typeof(f) stands for the type of the function f. In the body of f, NiLangCore.wrap\_tuple is used to unify output data types to tuples. The outputs of SWAP are assigned back to its input variables. At the end of this function, this macro attaches a return statement that returns all input variables.

The compilation of a reversible function to native Julia functions is consisted of three stages: *preprocessing*, *reversing* and *translation*. Fig. 3 shows the compilation of the complex valued log function body, which is originally defined as follows.

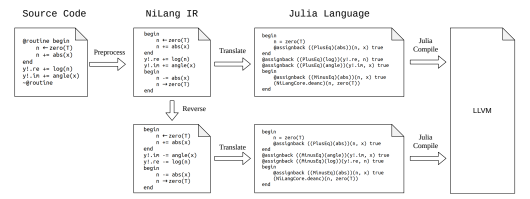


Figure 3: Compiling the body of the complex valued log function defined in Listing. 1.

Listing 1: Reversible implementation of the complex valued log function.

In the *preprocessing* stage, the compiler pre-processes human inputs to reversible NiLang IR. The preprocessor removes redundant grammars and expands shortcuts. In the left most code box in Fig. 3, one uses @routine <stmt> statement to record a statement, and ~@routine to insert the corresponding inverse statement for uncomputing. The computing-uncomputing macros @routine and ~@routine is expanded in this stage. Here, one can input " $\leftarrow$ " and " $\rightarrow$ " by typing "\leftarrow[TAB KEY]" and "\rightarrow[TAB KEY]" respectively in a Julia editor or REPL.

In the *reversing* stage, based on this symmetric and reversible IR, the compiler generates reversed statements according to table Table 1.

In the *translation* stage, the compiler translates this reversible IR as well as its inverse to native Julia code. It adds @assignback before each function call, inserts codes for reversibility check, and handle control flows. We can expand the @assignback macro to see the compiled expression.

```
julia> macroexpand(Main, :(@assignback PlusEq(log)(y!.re, n)))
quote
    var"##277" = (PlusEq(log))(y!.re, n)
begin
        y! = chfield(y!, Val{:re}(), ((NiLangCore.wrap_tuple)(var"##277"))[1])
        n = ((NiLangCore.wrap_tuple)(var"##277"))[2]
end
end
```

Here, the function chfield returns a complex number with an updated re field. This updated value is then assigned back to y!. In other words, this macro simulates "inplace" operations on immutable types. Except immutable field, one can also simulate mutability on function call and indexing. We call an expression that directly modifiable in NiLang a *dataview*, it can be a variable itself, a field or an element of a dataview, or a bijective mapping of a dataview.

The following are some examples of dataviews

statement	inverse
<f>(<args>)</args></f>	(~ <f>)(<args>)</args></f>
<y> += <f>(<args>)</args></f></y>	<y> -= <f>(<args>)</args></f></y>
<y> .+= <f>.(<args>)</args></f></y>	<y>= <f>.(<args>)</args></f></y>
<y> ⊻= <f>(<args>)</args></f></y>	<y> ⊻= <f>(<args>)</args></f></y>
<y> . ⊻= <f>. (<args>)</args></f></y>	<y> .⊻= <f>.(<args>)</args></f></y>
<a> ← <expr></expr></a>	<a> → <expr></expr></a>
<pre>begin      <stmts> end</stmts></pre>	begin $\sim$ ( <stmts>) end</stmts>
<pre>if (<pre></pre></pre>	<pre>if (<post>, <pre>)</pre></post></pre>
while ( <pre>, <post>)   <stmts> end</stmts></post></pre>	while ( <post>, <pre>)      ~(<stmts>) end</stmts></pre></post>
<pre>for <i>=<m>:<s>:<n></n></s></m></i></pre>	<pre>for <i>=<m>:-<s>:<n></n></s></m></i></pre>

Table 1: The statements in NiLang IR, where statements in the left column and those in the right column are reversible to each other. "~" is the symbol for reversing a statement or a function. "." is the symbol for the broadcasting magic in Julia, pre> stands for precondition</pr>, and post> stands for postcondition "begin <stmts> end" is the code block statement in Julia. It can be inverted by reversing the order as well as each element in it.

o real(x)	o real.(x)
o x.re	$\circ (x, y, z)$
o x[3]	o tget(x, 2) # tuple get index
∘ x'	• -x[2].re'

One can skip the assign back operation by inserting @skip! macro before a variable like f(y, @skip! x) to avoid automatic assigning back.

As a final step, the compiler attaches a return statement that returns all updated input arguments at the end of a function definition. Now, the function is ready to execute on the host language. One can also define a reversible constructor and destructor, we put this part in Appendix C.

## 3 Reversible automatic differentiation

## 3.1 First order gradient

The computation of gradients in NiLang contains two parts, computing and uncomputing. In the computing stage, the program marches forward and computes outputs. In the uncomputing stage, we attach each scalar with an extra gradient field and obtain a composite data type called GVar. We binding the adjoint program of a forward function to its inverse as shown in Fig. 4. When a instruction is uncalled, we first uncompute the value field of GVars to  $x_1$  and  $y_1$ , using the input information, we then update the gradient fields according to the formula in the right panel. The binding utilizes the multiple dispatch in Julia, where a function can be dynamically dispatched based on the run time type of more than one of its arguments. Here, we dispatch a inverse instruction with input type GVar to a function that corresponds to the above uncall procedure. Similar approach has been used in the forward mode AD package ForwardDiff Revels *et al.* (2016). For this example,

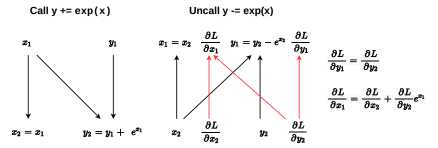


Figure 4: Binding the adjoint rule of  $y=\exp(x)$  to its uncomputing program.

to bind the backward rules for instructions (:+=)(exp) (or PlusEq(exp)) and (:-=)(exp) (or MinusEq(exp)). One can overload either of them as follows.

```
@i @inline function (:-=)(exp)(out!::GVar, x::GVar{T}) where T
    @routine @invcheckoff begin
        anc1 ← zero(value(x))
        anc1 += exp(value(x))
    end
    value(out!) -= identity(anc1)
    grad(x) += grad(out!) * anc1
    ~@routine
end
```

Here, the first four lines (the @routine statement) in the function body computes  $e^{x_2}$  and store it in an ancilla. The 5th line update the value dataview of out!. The 6th line updates the gradient fields of x and y, where update rule corresponds to the adjoint rule of (:+=)(exp). Finally, @routine uncomputes anc1 so that it can be returned to the "memory pool". The update rule defined on (:+=)(exp) is automatically generated by macro @i, which reflects the fact that taking inverse and computing gradients commute McInerney (2015).

#### 3.2 Second-order gradient

Combining the uncomputing program in NiLang with dual-numbers is a simple yet efficient way to obtain Hessians. By wrapping the elementary type with Dual defined in package ForwardDiff Revels *et al.* (2016) and throwing it into the gradient program defined in NiLang, one obtains one row/column of the Hessian matrix straightforward. We will show this point with an example of optimizing the graph embedding loss with Newton's trust region optimization in Sec. 4.3.

## 3.3 Differentiating complex numbers

To differentiate complex numbers, we re-implemented complex instructions reversiblly. For example, with the definition of complex valued log function in Listing. 1, the complex valued log can be differentiated with no extra effort.

## 3.4 Differentiating CUDA kernels

CUDA programming is playing a significant role in high-performance computing. In Julia, one can write kernel functions in native Julia language with CUDAnative Besard *et al.* (2017). NiLang is compatible with KernelAbstractions. With KernelAbstractions, one can insert a predefined routine in a kernel provided it does not have allocations inside. Since NiLang does not push variables into stack automatically for users, so one can write differentiable GPU kernels safely. We will show this feature in the benchmark of Bundle adjustment (BA) in Sec. 5.4. Here, one should notice that the shared read in forward will become shared write in the backward pass, which may result in an incorrect gradient. We will review this issue in Sec. 6.5.

## 4 Examples

In this section, we introduce several examples.

- o sparse matrix dot product,
- o first kind bessel function and memory oriented computational graph,
- o solving the graph embedding problem.

All codes used in this section as well as the next benchmark section are available in NiLang's paper repository.

#### 4.1 Sparse Matrices

Differentiating sparse matrices is useful in many applications, however, it can not benefit directly from generic backward rules for dense matrix because general rules do not keep the sparse structure. In the following, we will show how to convert a normal Frobenius dot product between two sparse matrices to a reversible one and differentiate it. Here, the Frobenius dot product is defined as trace(A'B). In SparseArrays code base, it is implemented as follows.

```
ra = rowvals(A)[ia]
function dot(A::AbstractSparseMatrixCSC{T1,S1},B
                                                                  elseif ra > rb
     ::AbstractSparseMatrixCSC{T2,S2}) where {T1
                                                                   ib += oneunit(S2)
     ,T2,S1,S2}
                                                                    ib < ib nxt || break
 m, n = size(A)
                                                                    rb = rowvals(B)[ib]
 size(B) == (m,n) || throw(DimensionMismatch("
                                                                  else # ra == rb
     matrices must have the same dimensions"))
                                                                    r += dot(nonzeros(A)[ia],
  r = dot(zero(T1), zero(T2))
                                                                          nonzeros(B)[ib])
 @inbounds for j = 1:n
                                                                    ia += oneunit(S1)
   ia = getcolptr(A)[j]
                                                                    ib += oneunit(S2)
   ia_nxt = getcolptr(A)[j+1]
                                                                    ia < ia_nxt && ib < ib_nxt || break</pre>
   ib = getcolptr(B)[j]
                                                                    ra = rowvals(A)[ia]
   ib\_nxt = getcolptr(B)[j+1]
                                                                    rb = rowvals(B)[ib]
   if ia < ia_nxt && ib < ib_nxt</pre>
                                                                  end
     ra = rowvals(A)[ia]
                                                                end
      rb = rowvals(B)[ib]
                                                              end
     while true
                                                            end
        if ra < rb</pre>
                                                           return r
          ia += oneunit(S1)
                                                          end
          ia < ia_nxt || break</pre>
```

In the above implementation, many statements are not reversible. However, it is easy to adapt it to a reversible style with NiLang without sacrificing much performance.

```
@i function dot(r::T, A::SparseMatrixCSC{T}, B::
      SparseMatrixCSC{T}) where {T}
    m \leftarrow size(A, 1)
    n \leftarrow size(A, 2)
    @invcheckoff branch_keeper ← zeros(Bool, 2*m)
    @safe size(B) == (m,n) || throw(
      DimensionMismatch("matrices must have the
      same dimensions"))
    @invcheckoff @inbounds for j = 1:n
         ia1 ← A.colptr[i]
         ib1 ← B.colptr[j]
         ia2 ← A.colptr[j+1]
         \texttt{ib2} \leftarrow \texttt{B.colptr[j+1]}
         ia ← ia1
         ib \leftarrow ib1
         @inbounds for i=1:ia2-ia1+ib2-ib1-1
              ra ← A.rowval[ia]
rb ← B.rowval[ib]
              if (ra == rb, ~)
                  r += A.nzval[ia]'*B.nzval[ib]
                                                                        end
              # b move -> true, a move -> false branch_keeper[i] \underline{V}= ia==ia2-1 ||
```

```
ra > rb
        ra → A.rowval[ia]
        rb \rightarrow B.rowval[ib]
        if (branch_keeper[i], ~)
             ib += identity(1)
             ia += identity(1)
         end
    ~@inbounds for i=1:ia2-ia1+ib2-ib1-1
         # b move -> true, a move -> false
        branch_keeper[i] ⊻= ia==ia2-1 ||
A.rowval[ia] > B.rowval[ib]
        if (branch_keeper[i], ~)
             ib += identity(1)
         else
             ia += identity(1)
        end
    end
@invcheckoff branch_keeper → zeros(Bool, 2*m
```

Here, all = are replaced with  $\leftarrow$  to indicate that these variables must be returned at the end of this function scope. We put a " $\sim$ " symbol in the postcondition field of if statements to indicate this postcondition is a dummy one that takes the same value as the precondition, i.e. the condition is not changed inside the loop body. If the precondition is changed by the loop body, one can

use a branch\_keeper vector to cache branch decisions. The memory of branch\_keeper can be returned through uncomputing (the "" statement above) so that it does not have to appear in the function argument list. Finally, after checking the correctness of the program, one can off the reversibility checks by using the macro @invcheckoff macro for better performance. We provide the benchmark of this function in Sec. 5.1, where the reversible sparse matrix matrix multiplication is also benchmarked.

#### 4.2 The first kind Bessel function

A Bessel function of the first kind of order  $\nu$  can be computed using Taylor expansion

$$J_{\nu}(z) = \sum_{n=0}^{\infty} \frac{(z/2)^{\nu}}{\Gamma(k+1)\Gamma(k+\nu+1)} (-z^2/4)^n$$
 (1)

where  $\Gamma(n) = (n-1)!$  is the Gamma function. One can compute the accumulated item iteratively as  $s_n = -\frac{z^2}{4}s_{n-1}$ . The irreversible implementation is

```
function besselj(v, z; atol=1e-8)
    k = 0
    s = (z/2)^v / factorial(v)
    out = s
    while abs(s) > atol
        k += 1
        s *= (-1) / k / (k+v) * (z/2)^2
        out += s
    end
    out
end
```

This computational process could be diagrammatically represented as a DAG as shown in Fig. 5 (a). In this diagram, the data is represented as an edge. It connects at most two nodes. One generates this data, and one consumes it. A computational graph is more likely a mathematical expression, and it can not describe inplace functions or control flows conveniently because it does not have the notation for memory and loops.

In the following, we introduce the reversible implementation and the memory oriented computational graph. The above Bessel function contains a loop with irreversible "\*=" operation inside. Intuitively, consequtive multiplication requires an increasing size of tape to cache the intermediate state  $s_n$ , since one can not release state  $s_{n-1}$  directly after computing  $s_n$  Perumalla (2013). To reduce the memory allocation without increasing the time complexity of the program, we introduce the following reversible approximate multiplier.

```
1 @i @inline function imul(out!, x, anc!)
2     anc! += out! * x
3     out! -= anc! / x
5WAP(out!, anc!)
end
```

Here, instruction SWAP exchanges values of the two variables, and anc!  $\approx 0$  is a *dirty ancilla*. Line 2 computes the result and accumulates it to the dirty ancilla, and we get an approximately correct output in anc!. Line 3 uncomputes out! approximately by using the information stored in anc!, leaving a dirty zero state in register out!. Line 4 swaps the contents in out! and anc!. Finally, we have an approximately correct output and a dirtier ancilla. The "approximate uncomputing" trick can be extensively used in practice. It mitigates the artificial irreversibility brought by the number system that we have adopted at the cost of output precision. The reason why this trick works here lies in the fact that from the mathematics perspective the state in nth step  $\{s_n, z\}$  contains the same amount of information as its previous state  $\{s_{n-1}, z\}$  except for some particular points, and it is highly possible to find an equation to uncompute the previous state from the current state. With this approximate multiplier, we implement  $J_{\gamma}$  as follows.

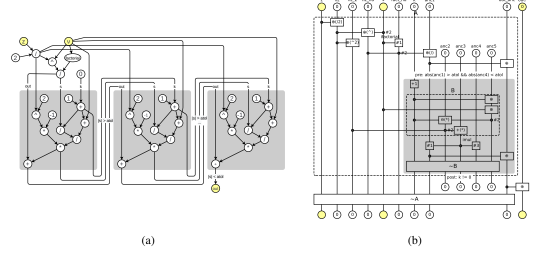


Figure 5: (a) The traditional computational graph for the irreversible implementation of the first kind Bessel function. A vertex (circle) is an operation, and a directed edge is a variable. The gray regions are the body of the unrolled while loop. (b) The memory oriented computational graph for the reversible implementation of the first kind Bessel function. Notations are explained in Fig. 1. The gray region is the body of a while loop. Its precondition and postcondition are positioned on the top and bottom, respectively.

```
anc5 += identity(k)
using NiLang, NiLang.AD
                                                                                 anc5 += identity(\nu)
                                                                                 anc2 -= k * anc5
@i function ibesselj(out!, v, z; atol=1e-8)
                                                                                 anc3 += halfz_power_2 / anc2
    k \leftarrow 0
    fact_nu \leftarrow zero(v)
                                                                            imul(anc1, anc3, anc4)
    halfz \leftarrow zero(z)
                                                                            out_anc += identity(anc1)
    halfz_power_nu \leftarrow zero(z)
                                                                             ~@routine
    halfz power 2 \leftarrow zero(z)
                                                                        end
    out\_anc \leftarrow zero(z)
                                                                   end
    anc1 \leftarrow zero(z)
                                                                   out! += identity(out_anc)
    anc2 \leftarrow zero(z)
                                                                   ~@routine
    anc3 \leftarrow zero(z)
                                                               end
    anc4 \leftarrow zero(z)
    anc5 \leftarrow zero(z)
                                                               @i function ifactorial(out!, n)
                                                                   anc \leftarrow zero(n)
    @routine begin
                                                                   out! += identity(1)
        halfz += z / 2
                                                                   for i=1:n
         halfz_power_nu += halfz ^ v
                                                                       imul(out!, i, anc)
        halfz power 2 += halfz ^ 2
                                                                   end
        ifactorial(fact_nu, v)
                                                               end
         anc1 += halfz_power_nu/fact_nu
                                                               @i @inline function imul(out!::T, x::T, anc!::T)
         out_anc += identity(anc1)
                                                                      where T<:Integer
         while (abs(unwrap(anc1))>atol && abs(
                                                                   anc! += out! * x
      unwrap(anc4)) < atol, k!=0)
                                                                   out! -= anc! ÷ x
             k += identity(1)
                                                                   SWAP(out!, anc!)
             @routine begin
```

The above algorithm uses a constant number of ancilla, while the time overhead factor is also a constant. Ancilla anc4 plays the role of *dirty ancilla* in multiplication, and it is uncomputed rigorously in the uncomputing stage marked by ~@routine.

This reversible program can be diagrammatically represented as a memory oriented computational graph as shown in Fig. 5 (b). In this graph, a variable is a vertical line, while a function is a parallel line. The critical difference comparing with the traditional computational graph is that it adopts a variable oriented view. A variable can be accessed by multiple functions. Hence it represents a hypergraph rather than a simple graph. If a function uses a variable but does not change the contents in it, we call this variable a control parameter of this function and put a dot at the cross. Otherwise, if the content is changed, we put a square. This diagram can be used to analyse uncomputable variables. In this example routine "B" uses  $hz_2$ ,  $\nu$  and k as control parameters, and changes

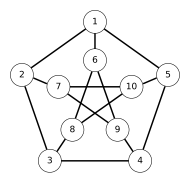


Figure 6: The Petersen graph has 10 vertices and 15 edges. We want to find a minimum embedding dimension for it.

the contents in anc2, anc3 and anc5. while the following operation imul does not change these variables. Hence we can apply the inverse routine ~B to safely restore contents in anc2, anc3 and anc5, and this is what people called compute-copy-uncompute paradigm.

One can obtain gradients of this function by calling Grad(ibesselj).

```
julia> out!, x = 0.0, 1.0
(0.0, 1.0)
julia> Grad(ibesselj)(Val(1), out!, 2, x)
(Val{1}(), GVar(0.0, 1.0), 2, GVar(1.0, 0.2102436))
```

Here, Grad(ibesselj) returns a callable instance of type Grad{typeof(ibesselj)}. The first parameters Val(1) specifies the position of loss in argument list. The Hessian can be obtained by feeding dual-numbers into this gradient function.

Here, the gradient field of hxx is defined as  $\frac{\partial \text{out!}}{\partial x}$ , which is a Dual number. It has a field partials that store the derivative for x. It corresponds to the Hessian  $\frac{\partial \text{out!}^2}{\partial x^2}$  that we need.

## 4.3 Solving a graph embedding problem

Graph embedding can be used to find representation for an order parameter Takahashi and Sandvik (2020) in condensed matter physics. Ref. Takahashi and Sandvik (2020) considers a problem of finding the minimum Euclidean space dimension k that a Petersen graph can fit into, with extra requirements that the distance between a pair of connected vertices has the same value  $l_1$ , and the distance between a pair of disconnected vertices has the same value  $l_2$  and  $l_2 > l_1$ . The Petersen graph is ten vertices graph, as shown in Fig. 6. Let us denote the set of connected and disconnected vertex pairs as  $L_1$  and  $L_2$ , respectively. This problem can be variationally solved by differential

programming by designing the subsequent loss.

$$\mathcal{L} = \text{Var}(\text{dist}(L_1)) + \text{Var}(\text{dist}(L_2)) + \exp(\text{relu}(\overline{\text{dist}(L_1)} - \overline{\text{dist}(L_2)} + 0.1))) - 1$$
 (2)

The first line is a summation of distance variances in two sets of vertex pairs, where VarX means taking the variance of samples in X. The second line is used to guarantee  $l_2 > l_1$ , where  $\overline{X}$  means taking the average of samples in X. Its reversible implementation could be found in our benchmark repository.

We repeat the training for each dimension k from 1 to 10 and search for possible solutions by variationally optimizing the positions of vertices. In each training, we fix two of the vertices and train the rest. Otherwise, the program will find the trivial solution with overlapped vertices. For k = 5, we can get a loss close to machine precision with high probability, while for k < 5, the loss is always much higher than 0. From the solution, it is easy to see  $l_2/l_1 = \sqrt{2}$  is the solution. For k = 5, an Adam optimizer with a learning rate 0.01 Kingma and Ba requires  $\sim 2000$  steps training. The trust region Newton's method converges much faster, which requires  $\sim 20$  computations of Hessians to reach convergence. Although training time is comparable, the converged precision of the later is much better.

#### 5 Benchmarks

In the following benchmarks the CPU device is Intel(R) Xeon(R) Gold 6230 CPU @ 2.10GHz, and the GPU device is Nvidia Titan V. For NiLang benchmarks, we have turned off the reversibility check off to achieve better performance.

## 5.1 Sparse matrices

We benchmarked the call, uncall and backward propagation time used for sparse matrix dot product and matrix multiplication. Here, we estimate the time for back propagating gradients rather than including both forward and backward, since mul! does not output a scalar as loss.

	dot	mul! (complex valued)
Julia-O	3.493e-04	8.005e-05
NiLang-O	4.675e-04	9.332e-05
NiLang-B	5.821e-04	2.214e-04

Table 2: Absolute runtimes in seconds for computing the objectives (O) and the backward pass (B) of sparse matrix operations. The matrix size is  $1000 \times 1000$ , and the element density is 0.05. The total time used in computing gradient can be estimated as a sum of "O" and "B".

The time used for computing backward pass is approximately 1.5-3 times the Julia's native forward pass. This is because the instruction length of differentiating basic arithmetic instructions is longer than pure computing by a factor of 2 or more.

#### 5.2 Bessel Function

We differentiate the first type Bessel function in Sec. 4.2 and show the benchmarks in Table 3. In the table, Julia is the CPU time used for running the irreversible forward program. It is the baseline for benchmarking. NiLang (call/uncall) is the time of reversible call or uncall. Both of them are ~ 3 times slower than its irreversible counterpart. Since Bessel function has only one input argument, forward mode AD tools are faster than reverse mode AD, both source-to-source framework ForwardDiff and operator overloading framework Tapenade have the a comparable computing time with the pure function call.

NiLang.AD is the reverse mode AD submodule in NiLang, and it takes 13.6 times the native Julia program, and is also 2 times slower than Tapenade. However, the key point is, there is no extra

	$T_{\min}/ns$	Space/KB
Julia-O	17	0
NiLang-O	53	0
Tapenade-O	32	0
ForwardDiff-G	39	0
NiLang-G	231	0
NiLang-G (CUDA)	1.4	0
ReverseDiff-G	7198	7.3
Zygote-G	22561	13.47
Tapenade-G (Forward)	30	0
Tapenade-G (Backward)	111	> 0

Table 3: Time and space used for computing objective (O) and gradient (G) of the first kind Bessel function  $J_2(1.0)$ . The CUDA time is averaged over a batch size of 4000, which is not a fair comparison but shows how much performance can we get from GPU in the parallel computing context.

memory allocation like stack operations in the whole computation. The controllable memory allocation of NiLang makes it compatible with CUDA program. In other backward mode AD like Zygote, ReverseDiff and Tapenade, the memory allocation in heap is nonzero due to the checkpointing and possible failure of type inference.

## 5.3 Graph embedding problem

Since one can combine ForwardDiff and NiLang to obtain Hessians, it is interesting to see how much performance we can get in differentiating the graph embedding program in Sec. 4.3.

k	2	4	6	8	10
Julia-O	4.477e-06	4.729e-06	4.959e-06	5.196e-06	5.567e-06
NiLang-O	7.173e-06	7.783e-06	8.558e-06	9.212e-06	1.002e-05
NiLang-U	7.453e-06	7.839e-06	8.464e-06	9.298e-06	1.054e-05
NiLang-G	1.509e-05	1.690e-05	1.872e-05	2.076e-05	2.266e-05
ForwardDiff-G	1.518e-05	4.053e-05	6.732e-05	1.184e-04	1.701e-04
ReverseDiff-G	1.384e-04	1.928e-04	2.392e-04	2.893e-04	3.556e-04
Zygote-G	5.315e-04	5.570e-04	5.811e-04	6.096e-04	6.396e-04
(NiLang+F)-H	4.528e-04	1.025e-03	1.740e-03	2.577e-03	3.558e-03
ForwardDiff-H	2.378e-04	2.380e-03	6.903e-03	1.967e-02	3.978e-02
(ReverseDiff+F)-H	1.966e-03	6.058e-03	1.225e-02	2.035e-02	3.140e-02

Table 4: Absolute runtimes in seconds for computing the objectives (O), uncall objective (U), gradients (G) and Hessians (H) of the graph embedding program. k is the embedding dimension, the number of parameters is 10k.

In Table 4, we show the the performance of different implementations by varying the dimension k. As the baseline, (a) shows the time for computing the function call. We have reversible and irreversible implementations, where the reversible program is slower than the irreversible native Julia program by a factor of  $\sim 2$  due to the uncomputing overhead. The reversible program shows the advantage of obtaining gradients when the dimension  $k \geq 3$ . The larger the number of inputs, the more advantage it shows due to the overhead proportional to input size in forward mode AD. The same reason applies to computing Hessians. The combining NiLang and ForwardDiff gives better performance when  $k \geq 3$  comparing with other AD frameworks.

#### 5.4 Gaussian mixture model and bundle adjustment

We reproduced the benchmarks for Gaussian mixture model (GMM) and bundle adjustment (BA) in Ref. Srajer *et al.* (2018) by re-writing the programs in a reversible style. We show the results in Table 5 and Table 6. In our new benchmarks, we also rewrite the ForwardDiff program for a fair benchmark, this explains the difference between our results and the original benchmark. The Tapenade data is obtained by executing the docker file provided by the original benchmark, which is the baseline of our benchmark and the original benchmark.

# parameters	3.00e+1	3.30e+2	1.20e+3	3.30e+3	1.07e+4	2.15e+4	5.36e+4	4.29e+5
Julia-O	9.844e-03	1.166e-02	2.797e-01	9.745e-02	3.903e-02	7.476e-02	2.284e-01	3.593e+00
NiLang-O	3.655e-03	1.425e-02	1.040e-01	1.389e-01	7.388e-02	1.491e-01	4.176e-01	5.462e+00
Tapende-O	1.484e-03	3.747e-03	4.836e-02	3.578e-02	5.314e-02	1.069e-01	2.583e-01	2.200e+00
ForwardDiff-G	3.551e-02	1.673e+00	4.811e+01	1.599e+02	-	-	-	-
NiLang-G	9.102e-03	3.709e-02	2.830e-01	3.556e-01	6.652e-01	1.449e+00	3.590e+00	3.342e+01
Tapenade-G	5.484e-03	1.434e-02	2.205e-01	1.497e-01	4.396e-01	9.588e-01	2.586e+00	2.442e+01

Table 5: Absolute runtimes in seconds for computing the objective (O) and gradients (G) of GMM with 10k data points.

In the GMM benchmark, NiLang's objective function has more overhead than irreversible programs in most cases. Except the uncomputing overhead, it is also because the bottleneck of this program is the matrix-vector multiplication, where our naive reversible implementation is much slower than the highly optimized BLAS function. The forward mode AD suffers from too large input dimension in this case, hence some data are missing. Although ForwardDiff is able to batch the gradient fields, the overhead proportional to input size still dominates. The source to source AD framework Tapenade is faster than NiLang in all scale, but the ratio between computing gradients and objective function are close. The memory consumption of our reversible implementation is quite low. The peak memory is only slightly more than twice of the original program, where the factor 2 comes from wrapping each variable with an extra gradient field.

# measurements	3.18e+4	2.04e+5	2.87e+5	5.64e+5	1.09e+6	4.75e+6	9.13e+6
Julia-O	2.020e-03	1.292e-02	1.812e-02	3.563e-02	6.904e-02	3.447e-01	6.671e-01
NiLang-O	2.708e-03	1.757e-02	2.438e-02	4.877e-02	9.536e-02	4.170e-01	8.020e-01
Tapenade-O	1.632e-03	1.056e-02	1.540e-02	2.927e-02	5.687e-02	2.481e-01	4.780e-01
ForwardDiff-J	6.579e-02	5.342e-01	7.369e-01	1.469e+00	2.878e+00	1.294e+01	2.648e+01
NiLang-J	1.651e-02	1.182e-01	1.668e-01	3.273e-01	6.375e-01	2.785e+00	5.535e+00
NiLang-J (GPU)	1.354e-04	4.329e-04	5.997e-04	1.735e-03	2.861e-03	1.021e-02	2.179e-02
Tapenade-J	1.940e-02	1.255e-01	1.769e-01	3.489e-01	6.720e-01	2.935e+00	6.027e+00

Table 6: Absolute runtimes in seconds for computing the objective (O) and Jacobians (J) in bundle adjustment.

In the BA benchmark, reverse mode AD shows slight advantage over ForwardDiff. The bottleneck of computing this large sparse Jacobian is computing the Jacobian of a elementary function with 15 input arguments and 2 output arguments, where input space is larger than the output space. In this instance, our reversible implementation is even faster than the source code transformation based AD framework Tapenade. With KernelAbstractions, we run the reversible program on GPU, which provides more than 200x speed up. Running codes on GPU (in Julia) requires not introducing any memory allocation at the kernel level. NiLang provides user the flexibility to manage the memory allocation instead of using a global stack.

## 6 Discussion and outlooks

In this paper, we show how to realize a reversible programming eDSL and implement an instruction level backward mode AD on top of it. It gives the user more flexibility to tradeoff memory and computing time comparing with traditional checkpointing. The Julia implementation NiLang gives the state-of-the-art performance and memory efficiency in obtaining first and second-order gradients

in applications, including first type Bessel function, sparse matrix manipulations, solving graph embedding problem, Gaussian mixture model and bundle adjustment.

In the following, we discuss some practical issues about reversible programming, and several future directions to go.

#### 6.1 Time Space Tradeoff

In history, there have been many discussions about time-space tradeoff on a reversible Turing machine (RTM). In the most straightforward g-segment tradeoff scheme Bennett (1989); Levine and Sherman (1990), an RTM model has either a space overhead that is proportional to computing time T or a computational overhead that sometimes can be exponential to the program size comparing with an irreversible counterpart. This result stops many people from taking reversible computing seriously as a high-performance computing scheme. In the following, we try to explain why the overhead of reversible computing is not as terrible as people thought.

First of all, the overhead of reversing a program is upper bounded by the checkpointing Chen *et al.* (2016) strategy used in many traditional machine learning package that memorizes inputs of primitives because similar strategy can also be used in reversible programming. Perumalla (2013) Reversible programming provides some alternatives to reduce the overhead. For example, accumulation is reversible, and it does not require checkpointing. Meanwhile, the memory allocation in some iterative algorithms can often be reduced with the "arithmetic uncomputing" trick without sacrificing reversibility, as shown in the ibesselj example in Sec. 4.2. Clever compiling based on memory oriented computational graphs (Fig. 1 and Fig. 5 (b)) can also be used to help user tradeoff between time and space. The overhead of a reversible program mainly comes from the uncomputing of ancillas. It is possible to automatically uncompute ancillas by analyzing variable dependency instead of asking users to write @routine and ~@routine pairs. In a hierarchical design, uncomputing can appear in every memory deallocation (or symbol table reduction). To quantify the overhead of uncomputing, we introduce the term uncomputing level as bellow.

**Definition 1** (uncomputing level). The log-ratio between the number of instructions of a reversible program and its irreversible counterpart.

From the lowest instruction level, whenever we reduce the symbol table in the tree of computation (or the abstract syntax tree), the computational cost doubles. In sparse matrix multiplication and dot product, we don't introduce uncomputing in the most time consuming part, so it is  $\sim 0$ . in Gaussian mixture model, the most time consuming matrix-vector multiplication is doubled, so it is  $\sim 1$ . In the first kind Bessel function, the most time consuming part is (nested) uncomputed twice, hence its uncomputing level is  $\sim 2$ . In the bundle adjustment program, its uncomputing level is also  $\sim 2$ . The computational overhead grows exponentially as the uncomputating level increases, which can be seen from some of the benchmarks in the main text.

At last, making reversible programming an eDSL rather than an independent language allows flexible choices between reversibility and computational overhead. For example, to deallocate the memory that stores gradients in a reversible language, one has to uncompute the whole process of obtaining them. As an eDSL, one has an alternative to deallocate the memory irreversibly outside the scope of a reversible program, i.e., trade energy with time.

## 6.2 Differentiability as a Hardware Feature

So far, our eDSL is compiled to Julia language. It relies on Julia's multiple dispatch to differentiate a program, which requires users to write generic programs. A more liable AD should be a hardware or micro instruction level feature. In the future, we can expect NiLang being compiled to reversible instructions Vieri (1999) and executed on a reversible device. A reversible devices can play a role of differentiation engine as shown in the hetero-structural design in Fig. 7. It defines a reversible instruction set and has a switch that controls whether the instruction calls a normal instruction or an instruction that also updates gradients. When a program calls a reversible differentiable subroutine, the reversible co-processor first marches forward, compute the loss and copy the result to the main memory. Then the co-processor execute the program backward and uncall instructions, initialize and updating gradient fields at the same time. After reaching the starting point of the program, the gradients are transferred to the global memory. Running AD program on a reversible device can

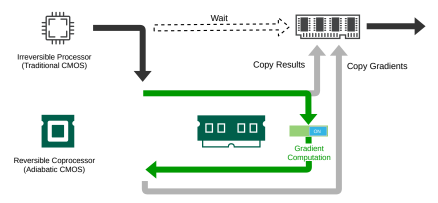


Figure 7: Energy efficient AI co-processor. Green arrows represents energy efficient operations on reversible devices.

save energy. Theoretically, the reversible routines do not necessarily cost energy, the only energy bottleneck is copying gradient and outputs to the main memory.

#### 6.3 The connection to Quantum programming

A Quantum device Nielsen and Chuang (2002) is a special reversible hardware that features quantum entanglement. The instruction set of classical reversible programming is a subset of quantum instruction set. However, building a universal quantum computer is difficult. Unlike a classical state, a quantum state can not be cloned. Meanwhile, it loses information by interacting with the environment. Classical reversible computing does not enjoy the quantum advantage, nor the quantum disadvantages of non-cloning and decoherence, but it is a model that we can try directly with our classical computer. It is technically smooth to have a reversible computing device to bridge the gap between classical devices and universal quantum computing devices. introducing entanglement little by little, we can accelerate some elementary components in reversible computing. For example, quantum Fourier transformation provides an alternative to the reversible adders and multipliers by introducing the Hadamard and CPHASE quantum gates Ruiz-Perez and Garcia-Escartin (2017). From the programming languages's perspective, most quantum programming language preassumes the existence of a classical coprocessor to control quantum devices Syore et al. (2018). It is also interesting to know what is a native quantum control flow like, and does quantum entanglement provide speed up to automatic differentiation? We believe the reversible compiling technologies will open a door to study quantum compiling.

#### 6.4 Gradient on ancilla problem

In this subsection, we introduce an easily overlooked problem in our reversible AD framework. An ancilla can sometimes carry a nonzero gradient when it is deallocated. As a result, the gradient program can be irreversible in the local scope. In NiLang, we drop the gradient field of ancillas instead of raising an error. In the following, we justify our decision by proving the following theorem.

**Theorem 1.** Deallocating an ancilla with constant value field and nonzero gradient field does not introduce incorrect gradients.

*Proof.* Consider a reversible function  $\mathbf{x}^i, b = f_i(\mathbf{x}^{i-1}, a)$ , where a and b are the input and output values of an ancilla. Since both a, b are constants that are independent of input  $\mathbf{x}^{i-1}$ , we have

$$\frac{\partial b}{\partial \mathbf{x}^{i-1}} = \mathbf{0}.\tag{3}$$

Discarding gradients should not have any effect on the value fields of outputs. The key is to show  $grad(b) \equiv \frac{\partial x^L}{\partial b}$  does appear in the grad fields of the output. It can be seen from the back-propagation rule

$$\frac{\partial \mathbf{x}^{L}}{\partial \mathbf{x}^{i-1}} = \frac{\partial \mathbf{x}^{L}}{\partial \mathbf{x}^{i}} \frac{\partial \mathbf{x}^{i}}{\partial \mathbf{x}^{i-1}} + \frac{\partial \mathbf{x}^{L}}{\partial b} \frac{\partial b}{\partial \mathbf{x}^{i-1}},\tag{4}$$

where the second term with  $\frac{\partial \mathbf{x}^L}{\partial b}$  vanishes naturally. We emphasis here, the value part of discarded ancilla must be a constant.

## 6.5 Shared read and write problem

One should be careful about shared read in reversible programming AD, because the shared read can introduce shared write in the adjoint program. Let's begin with the following expression.

```
y += x * y
```

Most people will agree that this statement is not reversible and should not be allowed because it changes input variables. We call it the *simultaneous read-and-write* issue. However, the following expression with two same inputs is a bit subtle.

```
y += x * x
```

It is reversible, but should not be allowed in an AD program because of the *shared write* issue. It can be seen directly from the expanded expression.

```
julia> macroexpand(Main, :(@instr y += x * x))
quote
    var"##253" = ((PlusEq)(*))(y, x, x)
begin
    y = ((NiLangCore.wrap_tuple)(var"##253"))[1]
    x = ((NiLangCore.wrap_tuple)(var"##253"))[2]
    x = ((NiLangCore.wrap_tuple)(var"##253"))[3]
end
end
```

In an AD program, the gradient field of x will be updated. The later assignment to x will overwrite the former one and introduce an incorrect gradient. One can get free of this issue by avoiding using same variable in a single instruction

```
anc ← zero(x)
anc += identity(x)
y += x * anc
anc -= identity(x)
```

or equivalently,

```
y += x ^ 2
```

Share variables in an instuction can be easily identified and avoided. However, it becomes tricky when one runs the program in a parallel way. For example, in CUDA programming, every thread may write to the same gradient field of a shared scalar. How to solve the shared write in CUDA programming is still an open problem, which limits the power of reversible programming AD on GPU.

#### 6.6 Outlook

We can use NiLang to solve some existing issues related to AD. We can use it to generate AD rules for existing machine learning packages like ReverseDiff, Zygote Innes *et al.* (2019), Knet Yuret (2016), and Flux Innes *et al.* (2018). Many backward rules for sparse arrays and linear algebra operations have not been defined yet in these packages. We can also use the flexible time-space

tradeoff in reversible programming to overcome the memory wall problem in some applications. A successful, related example is the memory-efficient domain-specific AD engine in quantum simulator Yao Luo *et al.* (2019). This domain-specific AD engine is written in a reversible style and solved the memory bottleneck in variational quantum simulations. It also gives so far the best performance in differentiating quantum circuit parameters. Similarly, we can write memory-efficient normalizing flow Kobyzev *et al.* (2019) in a reversible style. Normalizing flow is a successful class of generative models in both computer vision Kingma and Dhariwal (2018) and quantum physics Dinh *et al.* (2016); Li and Wang (2018), where its building block bijector is reversible. We can use a similar idea to differentiate reversible integrators Hut *et al.* (1995); Laikov (2018). With reversible integrators, it should be possible to rewrite the control system in robotics Giftthaler *et al.* (2017) in a reversible style, where scalar is a first-class citizen rather than tensor. Writing a reversible control program should boost training performance. Reversibility is also a valuable resource for training.

To solve the above problems better, reversible programming should be improved from multiple perspectives. First, we need a better compiler for compiling reversible programs. To be specific, a compiler that admits mutability of data, and handle shared read and write better. Then, we need a reversible number system and instruction set to avoid rounding errors and support reversible control flows better. There are proposals of reversible floating point adders and multipliers, however these designs require allocating garbage bits in each operation Nachtigal et al. (2010, 2011); Nguyen and Meter (2013); Häner et al. (2018). In NiLang, one can simulate rigorous reversible arithmetics with the fixed-point number package FixedPointNumbers. However, a more efficient reversible design requires instruction-level support. Some other numbers systems are reversible under \*= and /= rather than += and -=, including LogarithmicNumbers Taylor et al. (1988) and TropicalNumbers. They are powerful tools to solve domain specific problems, for example, we have an upcoming work about differentiating over tropical numbers to solve the ground state configurations of a spinglass system efficiently. We also need come from like instruction as a partner of goto to specify the postconditions in our instruction set. Finally, although we introduced that the adiabatic CMOS as a better choice as the computing device in a spacecraft DeBenedictis et al. (2017). There are some challenges in the hardware side too, one can find a proper summary of these challenges in Ref. Frank (2005).

Solutions to these issues requires the participation of people from multiple fields.

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## A NiLang Grammar

To define a reversible function one can use "@i" plus a standard function definition like bellow

definition of "<stmts>" are shown in the grammar page bellow. The following is a list of terminologies used in the definition of grammar

- ident, symbols
- num, numbers
- $\epsilon$ , empty statement
- JuliaExpr, native Julia expression
- [], zero or one repetitions.

Here, all *JuliaExpr* should be pure. Otherwise, the reversibility is not guaranteed. Dataview is a view of data. It can be a bijective mapping of an object, an item of an array, or a field of an object.

```
⟨Stmts⟩
                   ::=
                          \epsilon
                          | (Stmt)
                          |\langle Stmts \rangle \langle Stmt \rangle
         ⟨Stmt⟩
                         ⟨BlockStmt⟩
                          | (IfStmt)
                          | (WhileStmt)
                          | (ForStmt)
                          | (InstrStmt)
                          | (RevStmt)
                          | (AncillaStmt)
                          | \langle TypecastStmt \rangle
                          | (@routine) (Stmt)
                          | (@safe) JuliaExpr
                          | (CallStmt)
   ⟨BlockStmt⟩
                         begin (Stmts) end
    ⟨RevCond⟩
                          ( JuliaExpr , JuliaExpr )
       \langle IfStmt \rangle
                         if (RevCond) (Stmts) [else (Stmts)] end
  ⟨WhileStmt⟩
                         while (RevCond) (Stmts) end
       \langle Range \rangle
                   ::=
                          JuliaExpr : JuliaExpr [: JuliaExpr]
                         for\ ident = \langle Range \rangle \langle Stmts \rangle \ end
     ⟨ForStmt⟩
                   ::=
      ⟨KwArg⟩
                   ::=
                         ident = JuliaExpr
     \langle KwArgs \rangle
                         [\langle KwArgs \rangle,] \langle KwArg \rangle
                   ::=
     ⟨CallStmt⟩
                          JuliaExpr ( [(DataViews)] [; (KwArgs)] )
     ⟨Constant⟩
                         num \mid \pi \mid true \mid false
                   ::=
  ⟨InstrBinOp⟩
                   ::=
                          += | -= | ⊻=
  (InstrTrailer)
                         [.] ( [(DataViews)] )
    ⟨InstrStmt⟩
                          ⟨DataView⟩ ⟨InstrBinOp⟩ ident [⟨InstrTrailer⟩]
     \langle RevStmt \rangle
                         ~ (Stmt)
 ⟨AncillaStmt⟩
                         ident \leftarrow JuliaExpr
                   ::=
                          | ident \rightarrow JuliaExpr
(TypecastStmt)
                         ( JuliaExpr => JuliaExpr ) ( ident )
                   ::=
    ⟨@routine⟩
                          @routine ident (Stmt)
       ⟨@safe⟩
                   ::=
                          @safe JuliaExpr
  ⟨DataViews⟩
                          \epsilon
                          | (DataView)
                          | (DataViews), (DataView)
                          | (DataViews), (DataView) ...
   ⟨DataView⟩ ::=
                         ⟨DataView⟩ [ JuliaExpr ]
                          | (DataView) . ident
                          | JuliaExpr ( (DataView) )
                          | (DataView) '
                          | - (DataView)
                          | (Constant)
                          | ident
```

## B A table of instructions

A table instructions used in the main text

instruction	output
SWAP(a, b)	b, a
$ROT(a, b, \theta)$	$a\cos\theta - b\sin\theta, b\cos\theta + a\sin\theta, \theta$
$IROT(a, b, \theta)$	$a\cos\theta + b\sin\theta, b\cos\theta - a\sin\theta, \theta$
y += a * b	y + a * b, a, b
y += a/b	y + a/b, a, b
$y += a^{\wedge}b$	$y + a^b, a, b$
y += identity(x)	y + x, x
$y += \exp(x)$	$y + e^x, x$
$y += \log(x)$	$y + \log x, x$
$y += \sin(x)$	$y + \sin x, x$
$y += \cos(x)$	$y + \cos x, x$
y += abs(x)	y +  x , x
NEG(y)	-y
CONJ(y)	y'

Table 7: Predefined reversible instructions in NiLang.

## C Reversible Constructors

So far, the language design is not too different from a traditional reversible language. To port Julia's type system better, we introduce dataviews. The type used in the reversible context is just a standard Julia type with an additional requirement of having reversible constructors. The inverse of a constructor is called a "destructor", which unpacks data and deallocates derived fields. A reversible constructor is implemented by reinterpreting the new function in Julia. Let us consider the following statement.

```
x \leftarrow \text{new}\{TX, TG\}(x, g)
```

The above statement is similar to allocating an ancilla, except that it deallocates g directly at the same time. Doing this is proper because new is special that its output keeps all information of its arguments. All input variables that do not appear in the output can be discarded safely. Its inverse is

```
x \rightarrow \text{new}\{TX, TG\}(x, g)
```

It unpacks structure x and assigns fields to corresponding variables in the argument list. The following example shows a non-complete definition of the reversible type GVar.

```
end
julia> using NiLangCore
                                                                          @i function GVar(x::AbstractArray)
                                                                               GVar.(x)
iulia> @i struct GVar{T.GT} <: IWrapper{T}</pre>
                                                                           end
            x::T
                                                                      end
            a::GT
            function GVar{T,GT}(x::T, g::GT)
                                                              julia> GVar(0.5)
                          where {T,GT}
                                                              GVar{Float64,Float64}(0.5, 0.0)
                new{T,GT}(x, g)
            end
                                                              julia> (~GVar)(GVar(0.5))
            function GVar(x::T, g::GT)
                                                              0.5
                          where {T,GT}
                new{T,GT}(x, g)
                                                              iulia> (~GVar)(GVar([0.5, 0.6]))
            end
                                                              2-element Array{Float64,1}:
            @i function GVar(x::T) where T
                                                               0.5
                g \,\leftarrow\, \text{zero}(\textbf{x})
                                                               0.6
                x \leftarrow \text{new}\{T,T\}(x, g)
```

GVar has two fields that correspond to the value and gradient of a variable. Here, we put @i macro before both struct and function statements. The ones before functions generate forward and backward functions, while the one before struct moves ~GVar functions to the outside of the type definition. Otherwise, the inverse function will be ignored by Julia compiler.

Since an operation changes data inplace in NiLang, a field of an immutable instance should also be "modifiable". Let us first consider the following example.

```
julia> arr = [GVar(3.0), GVar(1.0)]
2-element Array{GVar{Float64,Float64},1}:
    GVar{Float64,Float64}(3.0, 0.0)
    GVar{Float64,Float64}(1.0, 0.0)

julia> x, y = 1.0, 2.0
    (1.0, 2.0)

julia> @instr -arr[2].g += x * y
2.0

julia> arr
2-element Array{GVar{Float64,Float64},1}:
    GVar{Float64,Float64}(3.0, 0.0)
    GVar{Float64,Float64}(1.0, -2.0)
```

In Julia language, the assign statement above will throw a syntax error because the function call "-" can not be assigned, and GVar is an immutable type. In NiLang, we use the macro @assignback to modify an immutable data directly. It translates the above statement to

The first line PlusEq(\*)(-arr[3].g, x, y) computes the output as a tuple of length 3. At lines 2-3, chfield(x, Val{:g}, val) modifies the g field of x and chfield(x, -, res[1]) returns -res[1]. Here, modifying a field requires the default constructor of a type not overwritten. The assignments in lines 4 and 5 are straightforward. We call a bijection of a field of an object a "dataview" of this object, and it is directly modifiable in NiLang. The definition of dataview can be found in Appendix A.