DIFFERENTIATE EVERYTHING WITH A REVERSIBLE EMBEDED DOMAIN-SPECIFIC LANGUAGE

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ABSTRACT

Reverse-mode automatic differentiation (AD) suffers from the issue of having too much space overhead to trace back intermediate computational states for The traditional method to trace back states is called back-propagation. checkpointing that stores intermediate states into a global stack and restore state through either stack pop or re-computing. The overhead of stack manipulations and re-computing makes the general purposed (not tensor-based) AD engines unable to meet many industrial needs. Instead of checkpointing, we propose to use reverse computing to trace back states by designing and implementing a reversible programming eDSL, where a program can be executed bi-directionally without implicit stack operations. The absence of implicit stack operations makes the program compatible with existing compiler features, including utilizing existing optimization passes and compiling the code as GPU kernels. We implement AD for sparse matrix operations and some machine learning applications to show that the performance of our framework has state-of-the-art performance.

1 Introduction

Most of the popular automatic differentiation (AD) tools in the market, such as TensorFlow Abadi et al. (2015), Pytorch Paszke et al. (2017), and Flux Innes et al. (2018) implements reverse mode AD at the tensor level to meet the need in machine learning. Later, People in the scientific computing domain also realized the power of these AD tools, they use these tools to solve scientific problems such as seismic inversion Zhu et al. (2020), variational quantum circuits simulation Bergholm et al. (2018); Luo et al. (2019) and variational tensor network simulation Liao et al. (2019); Roberts et al. (2019). To meet the diverse need in these applications, one sometimes has to define backward rules manually, for example

- 1. To differentiate sparse matrix operations used in Hamiltonian engineering Hao Xie & Wang, people defined backward rules for sparse matrix multiplication and dominant eigensolvers Golub & Van Loan (2012),
- 2. In tensor network algorithms to study the phase transition problem Liao et al. (2019); Seeger et al. (2017); Wan & Zhang (2019); Hubig (2019), people defined backward rules for singular value decomposition (SVD) function and QR decomposition Golub & Van Loan (2012).

To avoid defining backward rules manually, one can also use a general purposed AD (GP-AD) software like Tapenade Hascoet & Pascual (2013), OpenAD Utke et al. (2008) and Zygote Innes (2018); Innes et al. (2019) to differentiate a general program. These tools have been used in non-tensor based applications such as bundle adjustment Shen & Dai (2018) and earth system simulation Forget et al. (2015). However, these tools have their limitations too. In many practical applications, differentiating a program might do billions of computations. Frequent caching of data slows down the program significantly, while the memory usage will become a bottleneck as well. Although most of them are source-code-transformation based, they can not be used to differentiate GPU kernel functions because implicit stack operations are not compatible with kernel functions.

We need a new GP-AD framework that does not cache for users automatically. Hence we propose to implement the reverse mode AD on a reversible (domain-specific) programming

language Perumalla (2013); Frank (2017), where intermediate states can be traced backward without accessing an implicit stack. Reversible programming provides flexible time-space trade-off. It also allows people to utilize the reversibility to reverse a program without any overhead. In machine learning, reversibility is proven to significantly decrease the memory usage in unitary recurrent neural networks MacKay et al. (2018), normalizing flow Dinh et al. (2014), hyper-parameter learning Maclaurin et al. (2015) and residual neural networks Gomez et al. (2017); Behrmann et al. (2018). Reversible programming will make these happen naturally.

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 to study reversible programming is to support reversible computing devices Frank & Knight Jr (1999) such as adiabatic complementary metal-oxide-semiconductor (CMOS) Koller & Athas (1992), molecular mechanical computing system Merkle et al. (2018) and superconducting system Likharev (1977); Semenov et al. (2003); Takeuchi et al. (2014; 2017), and these reversible computing devices are orders more energy-efficient. Landauer Landauer (1961) proves that only when a device does not erase information (i.e. reversible), its energy efficiency can go beyond the thermal dynamic limit. However, these reversible programming languages can not be used in real scientific computing, since most of them do not have basic elements like floating point numbers, arrays, and complex numbers. This motivates us to build a new embedded domain-specific language (eDSL) in Julia Bezanson et al. (2012; 2017) as a new playground of GP-AD.

In this paper, we first introduce the language design of NiLang in Sec. 2. In Sec. 3, we explain the implementation of automatic differentiation in NiLang. In Sec. 4, we benchmark the performance of NiLang's AD with other AD software and explain why it is fast.

2 Language design

NiLang is an embedded domain-specific language (eDSL) NiLang on top of the host language Julia Bezanson et al. (2012; 2017). Julia is a popular language for scientific programming and machine learning. We choose Julia mainly for speed. Julia is a language with high abstraction, however, its clever design of type inference and just in time compiling make it has a C like speed. Meanwhile, it has rich features for meta-programming. Its package for pattern matching MLStyle allows us to define an eDSL in less than 2000 lines. Comparing with a regular reversible programming language, NiLang features array operations, rich number systems including floating-point numbers, complex numbers, fixed-point numbers, and logarithmic numbers. It also implements the compute-copy-uncompute Bennett (1973) macro to increase code reusability. Besides the above "nice" features, it also has some "bad" features to meet the practical needs. For example, it views the floating-point + and - operations as reversible. It also allows users to extend instruction sets and sometimes inserting external statements. These features are not compatible with future reversible hardware. NiLang's source code is available https://github.com/GiggleLiu/NiLang.jl, https://github.com/GiggleLiu/NiLangCore.jl. By the time of writing, the version of NiLang is v0.7.2.

2.1 REVERSIBLE FUNCTIONS AND INSTRUCTIONS

Mathematically, any irreversilbe mapping y = f(args...) can be trivially transformed to its reversible form y += f(args...) or y = f(args...) (y = f(args...)) is the bit-wise XOR), where y = f(args...) is a pre-emptied variable. But in numeric computing with finite precision, this is not always true. The reversibility of arithmetic instruction is closely related to the number system. For integer and fixed point number system, y += f(args...) and y -= f(args...) are rigorously reversible. For logarithmic number system and tropical number system Speyer & Sturmfels (2009), y = f(args...) and y -= f(args...) as reversible (not introducing the zero element). While for floating point numbers, none of the above operations are regorously reversible. However, for convenience, we ignore the rounding errors in floating point y = f(args...) and treat them on equal footing with fixed point numbers in the following discussion. Other reversible operations includes SWAP, ROT, NEG et. al., and this instruction set is extensible.

The following code defines a reversible multiplier.

Listing 1: A reversible multiplier

Macro @i generates two functions that are reversible to each other, multiplier and \sim multiplier, each defines a mapping $\mathbb{R}^3 \to \mathbb{R}^3$. The ! after a symbol is a part of the name, as a conversion to indicate the mutated variables.

2.2 REVERSIBLE MEMORY MANAGEMENT

A distinct feature of reversible memory management is that the content of a variable must be known when it is deallocated. We denote the allocation of a pre-emptied memory as $x \leftarrow 0$, and its inverse, deallocating a **zero emptied** variable, as $x \rightarrow 0$. An unknown variable can be pushed to a stack and used in the uncomputing stage with a pop statement. If a variable is allocated and deallocated in the local scope, we call it an ancilla.

Listing 2: Reversible complex valued log function $y += \log(|x|) + i\operatorname{Arg}(x)$.

Listing 3: Compute-copy-uncompute version of Listing. 2

```
@i @inline function (:+=)(log)(y!::Complex{T
     }, x::Complex{T}) where T
     @routine begin
     n ← zero(T)
     n += abs(x)
     end
     y!.re += log(n)
     y!.im += angle(x)
     ~@routine
end
```

Listing. 2 defines the complex valued accumulative log function. The macro @inline tells the compiler that this function can be inlined. One can input "\(-\)" by typing "\leftarrow[TAB KEY]" and "\rightarrow[TAB KEY]" respectively in a Julia editor or REPL. NiLang does not have immutable structs, so that the real part y!.re and imaginary y!.im of a complex number can be changed directly. It is easy to verify that the bottom two lines in the function body are the inverse of the top two lines. i.e., the bottom two lines uncomputes the top two lines. The motivation of uncomputing is to zero clear the contents in ancilla n so that it can be deallocated correctly. Compute-copy-uncompute is a useful design pattern in reversible programming so that we created a pair of macros @routine and ~@routine for it. One can rewrite the above function as in Listing. 3.

2.3 Reversible control flows

One can define reversible if, for and while statements in a reversible program. Fig. 1 (a) shows the flow chart of executing the reversible if statement. There are two condition expressions in this chart, a precondition and a postcondition. The precondition decides which branch to enter in the forward execution. After executing the specific branch, the program checks the consistency between precondition and postcondition to make sure they are consistent. To reverse this statement, one can exchange the precondition and postcondition, and reverse the expressions in both branches. Fig. 1 (b) shows the flow chart of the reversible while statement. There are also two conditions expressions. Before executing the condition expressions, the program presumes the postcondition is false. After

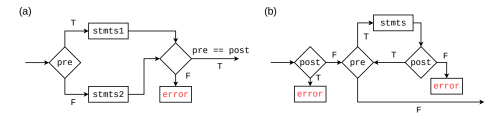


Figure 1: The flow chart for reversible (a) if statement and (b) while statement. "pre" and "post" represents precondition and postcondition respectively.

each iteration, the program asserts the postcondition to be true. To reverse this statement, one can exchange the precondition and postcondition, and reverse the body statements. The reversible for statement is similar to the irreversible one except that after execution, the program will assert the iterator to be unchanged. To reverse this statement, one can exchange start and stop and inverse the sign of step.

The following code computes the Fibonacci number recursively and reversibly.

Listing 4: Computing Fibonacci number recursively and reversibly.

Here, out! is an integer initialized to 0 for storing outputs. The precondition and postcondition are wrapped into a tuple. In the if statement, the postcondition is the same as the precondition, hence we omit the postcondition by inserting a "~" in the second field as a placeholder. In the while statement, the postcondition is true only for the initial loop. Once code is proven correct, one can turn off the reversibility check by adding @invcheckoff before a statement. This will remove the reversibility check and make the code faster and compatible with GPU kernels (kernel functions can not handle exceptions).

2.4 Reverse computing is not checkpointing

There are two approaches to trace back intermediate states of a computational process, one is checkpointing Griewank & Walther (2008); Chen et al. (2016) and another is reverse computing. Reverse computing and checkpointing share many similarities. When there is no time overhead, both have a space overhead $\sim O(T)$ (i.e. linear to time). They also have many differences. When a polynomial overhead in time is allowed, reversible computing has a minimum space overhead of $O(S \log(T))$ Bennett (1989); Levine & Sherman (1990); Perumalla (2013). For checkpointing, there can be no space overhead, since one can just recompute from beginning to obtain any intermediate state with time complexity $O(T^2)$. In practical using cases, reverse computing has the following advantages.

Reverse computing shows the advantage of handling effective codes with mutable structures and arrays. For example, the affine transformation can be implemented without any overhead.

Listing 5: Inplace affine transformation.

Here, the expression following the @safe macro is an external irreversible statement.

Reverse computing can utilize reversibility to trace back states without extra memory cost. For example, we can define the unitary matrix multiplication that can be used in a type of memory-efficient recurrent neural network Jing et al. (2016).

Listing 6: Two level decomposition of a unitary matrix.

Last but not least, reversible programming encourages users to code in a memory friendly style. Since allocations in reversible programming are explicit, programmers have the flexibility to control how to allocate memory and which number system to use. For example, to compute the power of a positive fixed-point number and an integer, one can easily write irreversible code as in Listing. 7

Listing 7: A regular power function.

```
function mypower(x::T, n::Int) where T
    y = one(T)
    for i=1:n
        y *= x
    end
    return y
end
```

Listing 8: A reversible power function.

Since the fixed-point number is not reversible under *=, naive checkpointing would require stack operations inside a loop. With reversible thinking, we can convert the fixed-point number to logarithmic numbers to utilize the reversibility of *= as shown in Listing. 8. Here, the algorithm to convert a regular fixed-point number to a logarithmic number can be efficient Turner (2010).

3 REVERSIBLE AUTOMATIC DIFFERENTIATION

3.1 Backpropagation

To backpropagate the program, we first reverse the code through source code transformation and then insert the gradient code through operator overloading. If we inline all the instructions in Listing. 3, the program would be like Listing. 9. The automatically generated inverse program (i.e. $(y, x) \rightarrow (y - \log(x), x)$) is like Listing. 10.

Listing 9: The function body of Listing. 3.

```
@routine begin
    nsq ← zero(T)
    n ← zero(T)
    nsq += x[i].re ^ 2
    nsq += x[i].im ^ 2
    n += sqrt(nsq)
end
y![i].re += log(n)
y![i].im += atan(x[i].im, x[i].re)
~@routine
```

Listing 10: The inverse of Listing. 9.

```
@routine begin
    nsq ← zero(T)
    n ← zero(T)
    nsq += x[i].re ^ 2
    nsq += x[i].im ^ 2
    n += sqrt(nsq)
end
y![i].re -= log(n)
y![i].im -= atan(x[i].im, x[i].re)
~@routine
```

To compute the adjoint of the computational process in Listing. 9, one simply insert the gradient code into its inverse in Listing. 10. The resulting inlined code is show in Listing. 11.

Listing 11: Insert the gradient code into Listing. 10, the original computational processes are highlighted in yellow background.

```
@routine begin
  nsg \leftarrow zero(GVar\{T,T\})
  n \leftarrow zero(GVar\{T,T\})
   gsga ← zero(T)
   gsqa += x[i].re.x * 2
   x[i].re.g -= gsqa * nsq.g
   gsqa -= nsq.x * 2
   gsqa -= x[i].re.x * 2
   gsqa \rightarrow zero(T)
   nsq.x += x[i].re.x ^2
   qsqb \leftarrow zero(T)
   gsqb += x[i].im.x * 2
   x[i].im.g = gsqb * nsq.g
   gsqb = x[i].im.x * 2
   qsqb \rightarrow zero(T)
   nsq.x += x[i].im.x ^2
   @zeros T ra rb
   rta += sqrt(nsq.x)
   rb += 2 * ra
   nsq.g -= n.g / rb
  rb -= 2 * ra
```

```
ra -= sqrt(nsq.x)
  ~@zeros T ra rb
  n.x += sqrt(nsq.x)
end
y![i].re.x -= log(n.x)
n.g += y![i].re.g / n.x
y![i].im.x-=atan(x[i].im.x,x[i].re.x)
@zeros T xy2 jac_x jac_y
xy2 += abs2(x[i].re.x)
xy2 += abs2(x[i].im.x)
jac_y += x[i].re.x / xy2
jac_x += (-x[i].im.x) / xy2
x[i].im.g += y![i].im.g * jac_y
x[i].re.g += y![i].im.g * jac_x
jac_x = (-x[i].im.x) / xy2
jac_y = x[i].re.x / xy2
xy2 = abs2(x[i].im.x)
xy2 -= abs2(x[i].re.x)
~@zeros T xy2 jac_x jac_y
~@routine
```

Here, @zeros TYPE var1 var2... is the macro to allocate multiple variables of the same type. Its inverse operations starts with ~@zeros deallocates zero emptied variables. In practise, "inserting gradients" is not achieved by source code transformation, but by changing the element type to GVar, a composite type with two fields, value x and gradient g. With multiple dispatching primitive instructions on this new type, values and gradients can be updated simultaneously. Although the code looks much longer, the computing time (with reversibility check closed) is not.

Listing 12: Time and allocation to differentiate complex valued log.

The performance is unreasonably good because the generated Julia code is further compiled to LLVM so that it can enjoy existing optimization passes. For example, the optimization passes can find out that for an irreversible device, uncomputing local variables n and nsq does not affect return values, so that it will ignore the code for uncomputing automatically. Unlike checkpointing based approaches that focus a lot in the optimization of data caching on a global stack, NiLang does not have any optimization pass in itself. Instead, it throws itself to existing optimization passes in Julia. Without accessing the global stack, NiLang's code is quite friendly to optimization passes. In this case, we also see the boundary between source code transformation and operator overloading can be vague in a Julia, in that the generated code can be very different from how it looks.

One can define the adjoint of a primitive instruction as a reversible function on **either** the function itself or its inverse, because the adjoints of reversible functions are reversible to each other too.

$$f: (\vec{x}, \vec{g}_x) \to (\vec{y}, \frac{\partial \vec{y}}{\partial \vec{x}} \vec{g}_x)$$
 (1)

$$f^{-1}: (\vec{y}, \vec{g}_y) \to (\vec{x}, \frac{\partial \vec{x}}{\partial \vec{y}} \vec{g}_y)$$
 (2)

It can be easily verified by applying the above two mappings consecutively, which turns out to be an identity mapping considering $\frac{\partial \vec{y}}{\partial \vec{x}} \frac{\partial \vec{x}}{\partial \vec{y}} = 1$. As an example, the joint functions for primitive instructions (:+=)(sqrt) and (:-=)(sqrt) used above can be defined as in Listing. 13.

Listing 13: Adjoints for primitives (:+=)(sqrt) and (:-=)(sqrt).

```
@i @inline function (:-=)(sqrt)(out!::GVar, x::GVar{T}) where T
    @routine @invcheckoff begin
        @zeros T a b
        a += sqrt(x.x)
        b += 2 * a
    end
    out!.x -= a
    x.g += out!.g / b
    ~@routine
end
```

3.2 Hessians

Combining forward mode AD and reverse mode AD 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. We will use this approach to compute Hessians in the graph embedding benchmark in Sec. A.2.

3.3 CUDA KERNELS

CUDA programming is playing a significant role in high-performance computing. In Julia, one can write GPU compatible functions in native Julia language with KernelAbstractions Besard et al. (2017). Since NiLang does not push variables into stack automatically for users, it is safe to write differentiable GPU kernels with NiLang. We will differentiate CUDA kernels with no more than extra 10 lines in the bundle adjustment (BA) benchmark in Sec. 4.1.

4 Benchmarks

We benchmark our framework with the state-of-the-art GP-AD frameworks, including source code transformation based Tapenade and Zygote and operator overloading based ForwardDiff and ReverseDiff. Since most tensor based AD software like famous TensorFlow and PyTorch are not designed for the using cases used in our benchmarks, we do not include those package to avoid an unfair comparison. 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 the reversibility check off to achieve a better performance.

4.1 Gaussian mixture model and bundle adjustment

We reproduced the benchmarks for Gaussian mixture model (GMM) and bundle adjustment (BA) in Srajer et al. (2018) by re-writing the programs in a reversible style. We show the results in Table 1 and Table 2. The Tapenade data is obtained by executing the docker file provided by the original benchmark, which provides a baseline for comparison.

# 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 1: Absolute runtimes in seconds for computing the objective (O) and gradients (G) of GMM with 10k data points. "-" represents missing data due to not finishing the computing in limited time.

NiLang's objective function is $\sim 2\times$ slower than normal code due to the uncomputing overhead. In this case, NiLang does not show advantage to Tapenade in obtaining gradients, the ratio between computing the gradients and the objective function are close. This is because the bottleneck of this model is the matrix vector multiplication, traditional AD can already handle this function well. We emphasis the extra memory usage in NiLang is approximately 0.5% of the original program, this should be better than most existing AD frameworks.

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

NiLang performs the best on CPU, while having zero allocation during computing local Jacobians of size 15×2 . We also compiled our adjoint program to GPU with no more than 10 lines of code with KernelAbstractions, which provides another $\sim 200 \times$ speed up. Parallelizing the adjoint code

requires the forward code not reading the same variable simultaneously in different threads, and this requirement is satisfied here.

You can find more benchmarks in Appendix A, including differentiating sparse matrix dot product and obtaining Hessians in the graph embedding application.

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A More Benchmarks

A.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 3: Absolute runtimes in seconds for computing the objectives (O) and the backward pass (B) of sparse matrix operations. The matrix size is 1000×1000 , and the element density is 0.05. The total time used in computing gradient can be estimated by summing "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.

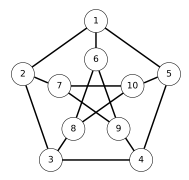


Figure 2: The Petersen graph has 10 vertices and 15 edges. We want to find a minimum embedding dimension for it.

A.2 Graph embedding problem

Graph embedding can be used to find a proper representation for an order parameter Takahashi & Sandvik (2020) in condensed matter physics. People want to find a minimum Euclidean space dimension k that a Petersen graph can embed into, that the distances between pairs of connected vertices are l_1 , and the distance between pairs of disconnected vertices are l_2 , where $l_2 > l_1$. The Petersen graph is shown in Fig. 2. Let us denote the set of connected and disconnected vertex pairs as L_1 and L_2 , respectively. This problem can be variationally solved with the following loss.

$$\mathcal{L} = \text{Var}(\text{dist}(L_1)) + \text{Var}(\text{dist}(L_2)) + \exp(\text{relu}(\overline{\text{dist}(L_1)} - \overline{\text{dist}(L_2)} + 0.1))) - 1$$
(3)

The first line is a summation of distance variances in two sets of vertex pairs, where Var(X) is 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 dimension k from 1 to 10. In each training, we fix two of the vertices and optimize the positions of the rest. Otherwise, the program will find the trivial solution with overlapped vertices. For k < 5, the loss is always much higher than 0, while for $k \ge 5$, we can get a loss close to machine precision with high probability. From the k = 5 solution, it is easy to see $l_2/l_1 = \sqrt{2}$. An Adam optimizer with a learning rate 0.01 Kingma & Ba requires ~ 2000 steps training. The trust region Newton's method converges much faster, which requires ~ 20 computations of Hessians to reach convergence. Although training time is comparable, the converged precision of the later is much better.

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 Table 4, we show the the performance of different implementations by varying the dimension k. The number of parameters is 10k. 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 ~ 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, where the combo of NiLang and ForwardDiff gives the best performance for $k \geq 3$.

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
ReverseDiff-G	2.823e-05	4.582e-05	6.045e-05	7.651e-05	9.666e-05
ForwardDiff-G	1.518e-05	4.053e-05	6.732e-05	1.184e-04	1.701e-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 times 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.