



Enzyme: High-Performance, Cross-Language, and Parallel Automatic Differentiation



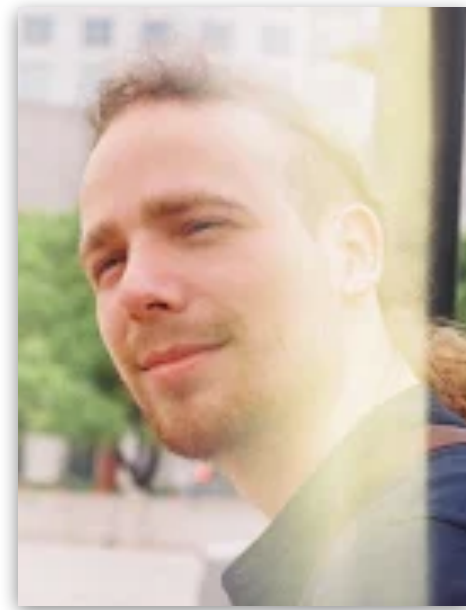
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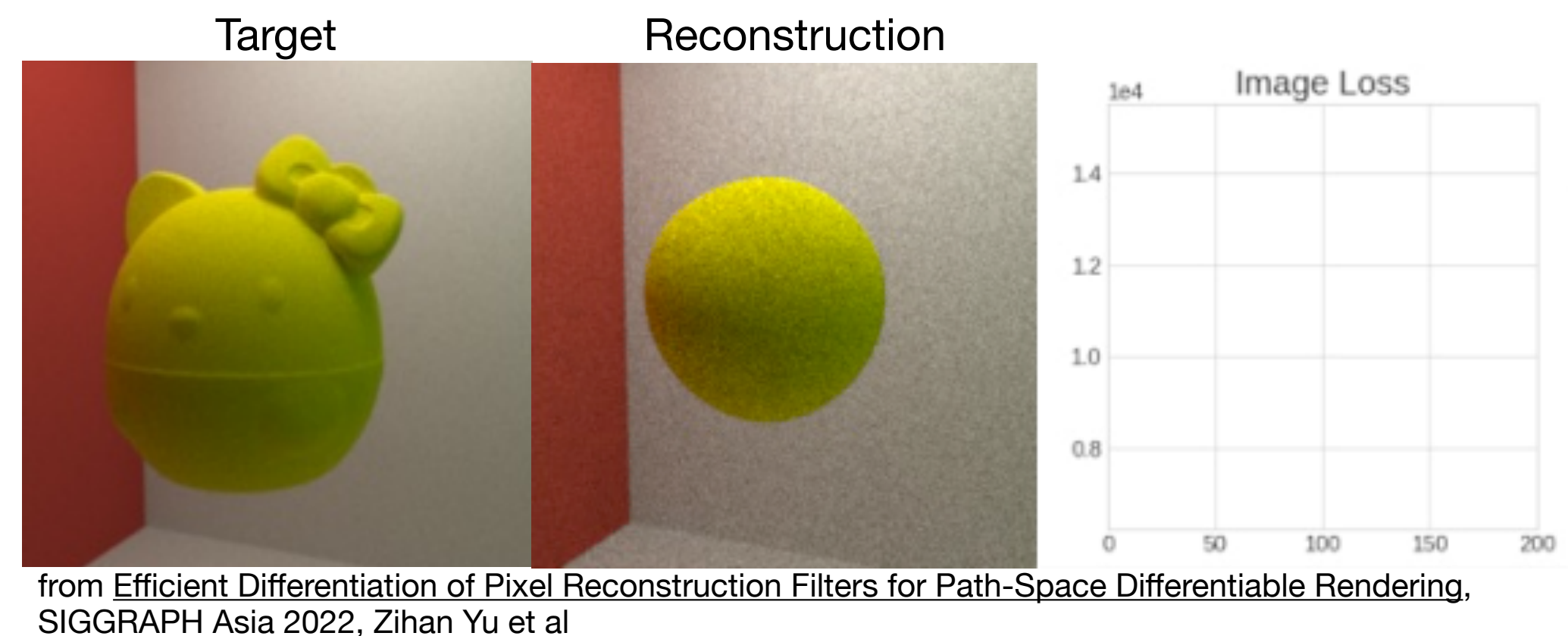
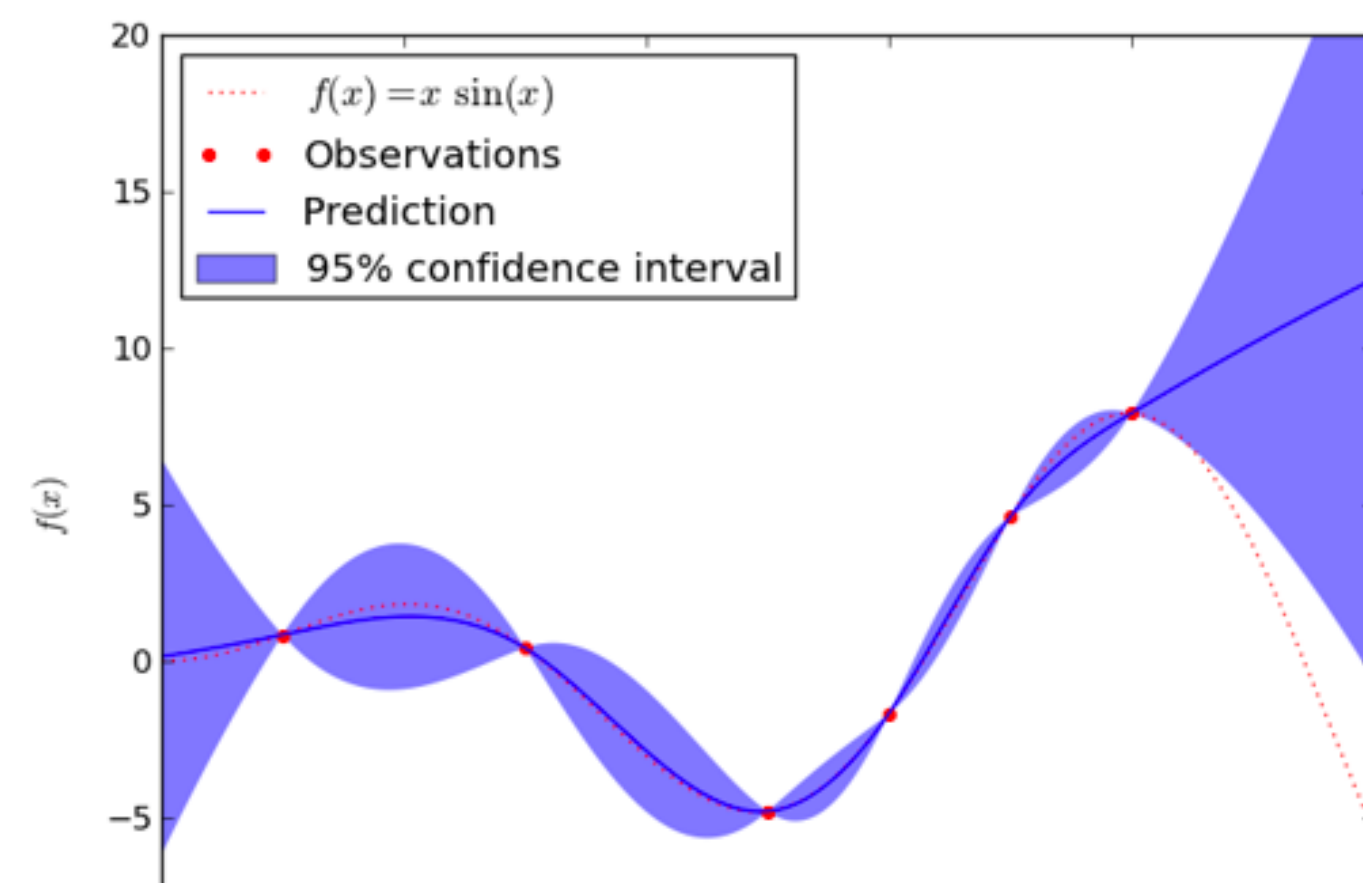
Manuel Drehwald

AP Calculus: Revisited

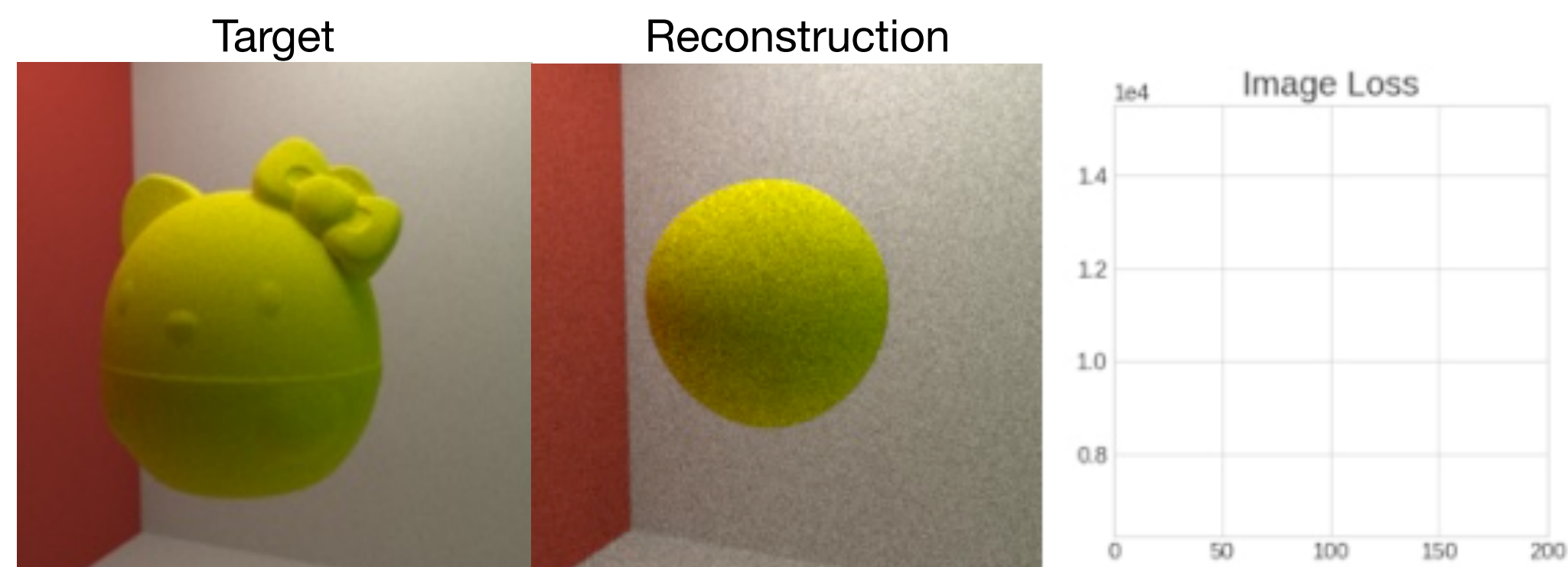
- Derivatives compute the rate of change of a function's output with respect to input(s)

$$f'(x) = \lim_{h \rightarrow 0} \frac{f(a+h) - f(a)}{h}$$

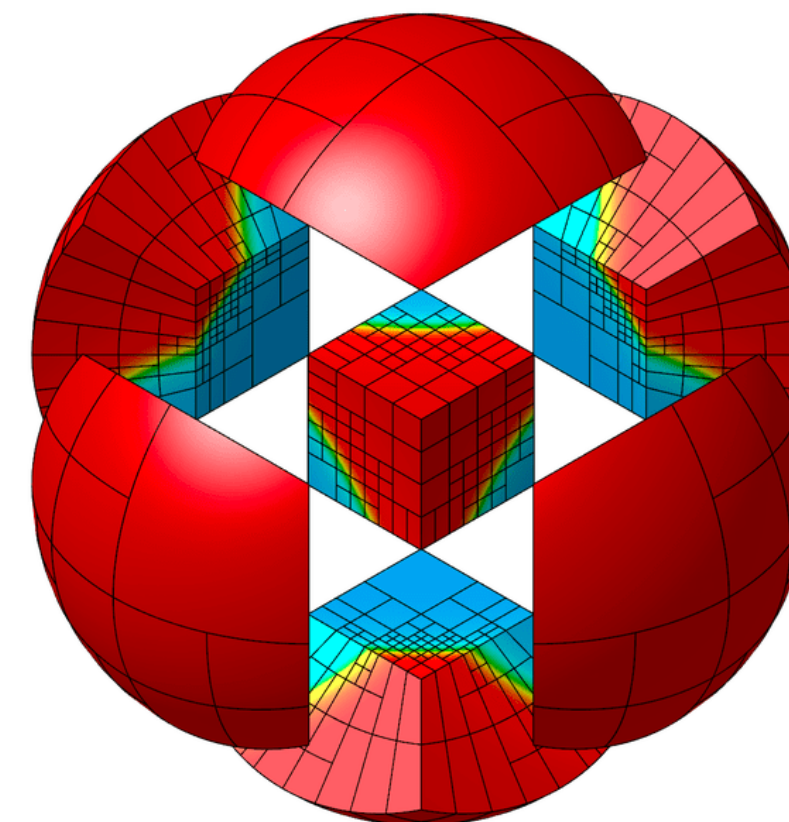
- Derivatives are used widely across science
 - Machine learning (back-propagation, Bayesian inference)
 - Scientific computing (modeling, simulation, uncertainty quantification)



AD-Powered Applications



from [Efficient Differentiation of Pixel Reconstruction Filters for Path-Space Differentiable Rendering](#), SIGGRAPH Asia 2022, Zihan Yu et al



from [MFEM Team at LLNL](#)



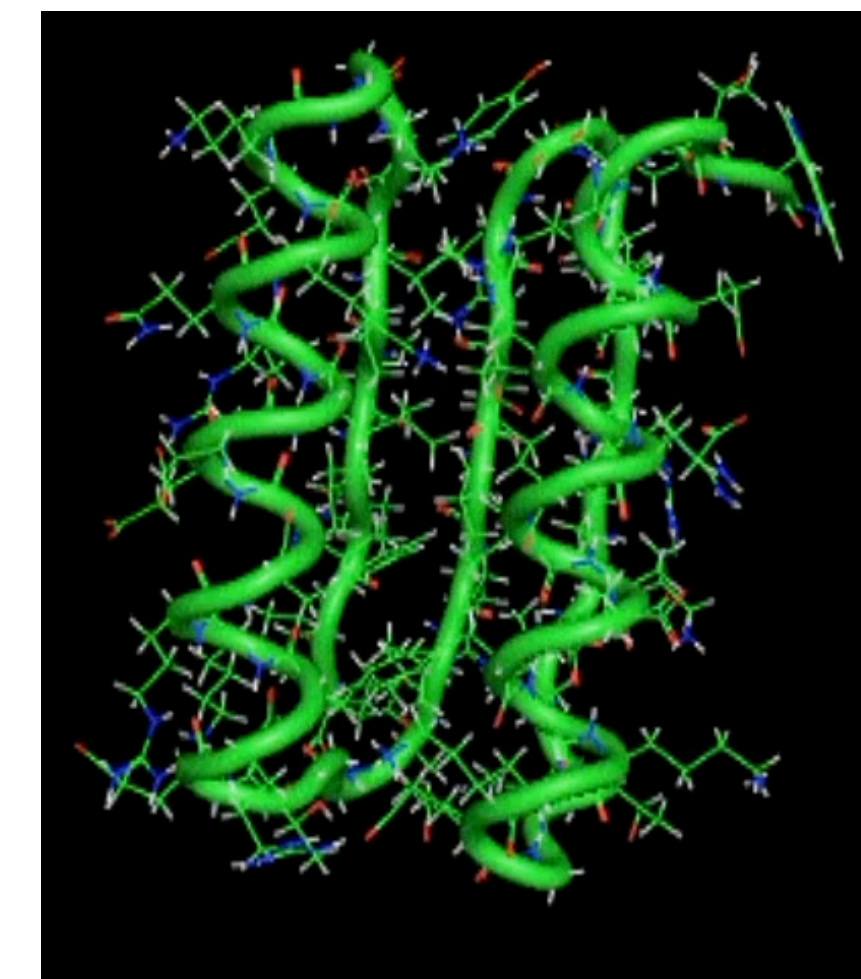
from [Comrade: High Performance Black-Hole Imaging](#) JuliaCon 2022, Paul Tiede (Harvard)



from [CLIMA & NSF CSSI: Differentiable programming in Julia for Earth system modeling \(DJ4Earth\)](#)



from [Center for the Exascale Simulation of Materials in Extreme Environments](#)

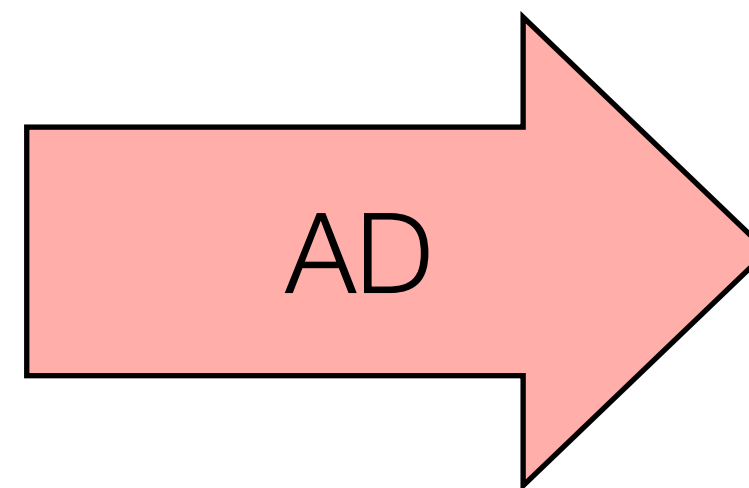


from [Differential Molecular Simulation with Molly.jl](#), EnzymeCon 2023, Joe Greener (Cambridge)

Automatic Derivative Generation

- Derivatives can be generated automatically from definitions within programs

```
double relu3(double x) {  
    if (x > 0)  
        return pow(x,3)  
    else  
        return 0;  
}
```



```
double grad_relu3(double x) {  
    if (x > 0)  
        return 3 * pow(x,2)  
    else  
        return 0;  
}
```

- Unlike numerical approaches, automatic differentiation (AD) can compute the derivative of ALL inputs (or outputs) at once, without approximation error!

```
// Numeric differentiation  
// f'(x) approx [f(x+epsilon) - f(x)] / epsilon  
double grad_input[100];  
  
for (int i=0; i<100; i++) {  
    double input2[100] = input;  
    input2[i] += 0.01;  
    grad_input[i] = (f(input2) - f(input))/0.001;  
}
```

```
// Automatic differentiation  
double grad_input[100];  
  
grad_f(input, grad_input)
```

Existing AD Approaches (1/3)

- Differentiable DSL (TensorFlow, PyTorch, DiffTaichi)
 - Provide a new language designed to be differentiated
 - Requires rewriting everything in the DSL and the DSL must support all operations in original code
 - Fast if DSL matches original code well

```
double relu3(double val) {  
    if (x > 0)  
        return pow(x,3)  
    else  
        return 0;  
}
```

Manually
Rewrite

```
import tensorflow as tf  
  
x = tf.Variable(3.14)  
  
with tf.GradientTape() as tape:  
    out = tf.cond(x > 0,  
                  lambda: tf.math.pow(x,3),  
                  lambda: 0  
    )  
print(tape.gradient(out, x).numpy())
```

Existing AD Approaches (2/3)

- Operator overloading (Adept, JAX)
 - Differentiable versions of existing language constructs (double => adouble, np.sum => jax.sum)
 - May require writing to use non-standard utilities
 - Often dynamic: storing instructions/values to later be interpreted

```
// Rewrite to accept either
//      double or adouble
template<typename T>
T relu3(T val) {
    if (x > 0)
        return pow(x,3)
    else
        return 0;
}
```

```
adept::Stack stack;
adept::adouble inp = 3.14;

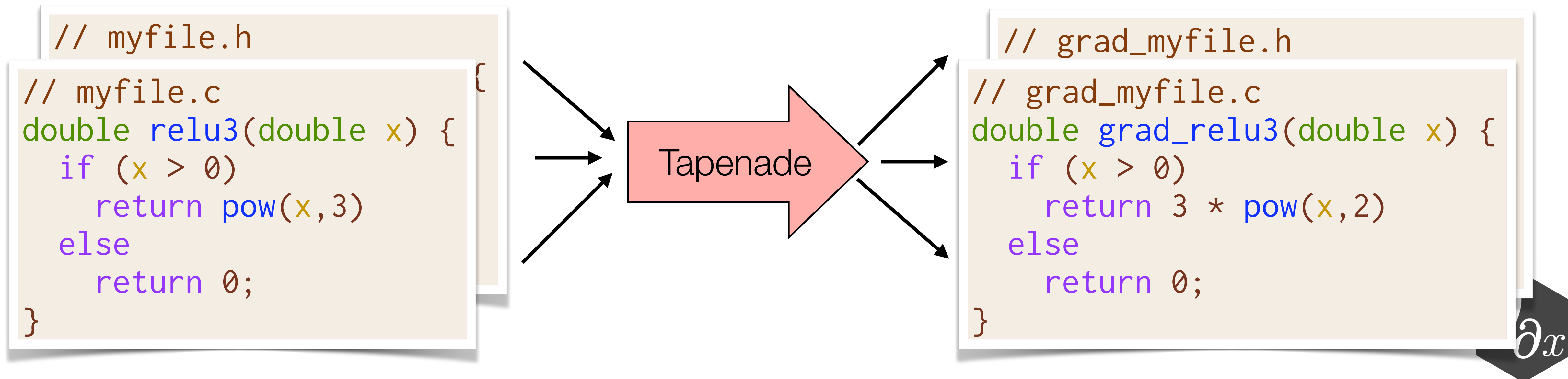
// Store all instructions into stack
adept::adouble out(relu3(inp));
out.set_gradient(1.00);

// Interpret all stack instructions
double res = inp.get_gradient(3.14);
```



Existing AD Approaches (3/3)

- Source rewriting
 - Statically analyze program to produce a new gradient function in the source language
 - Re-implement parsing and semantics of given language
 - Requires all code to be available ahead of time => hard to use with external libraries

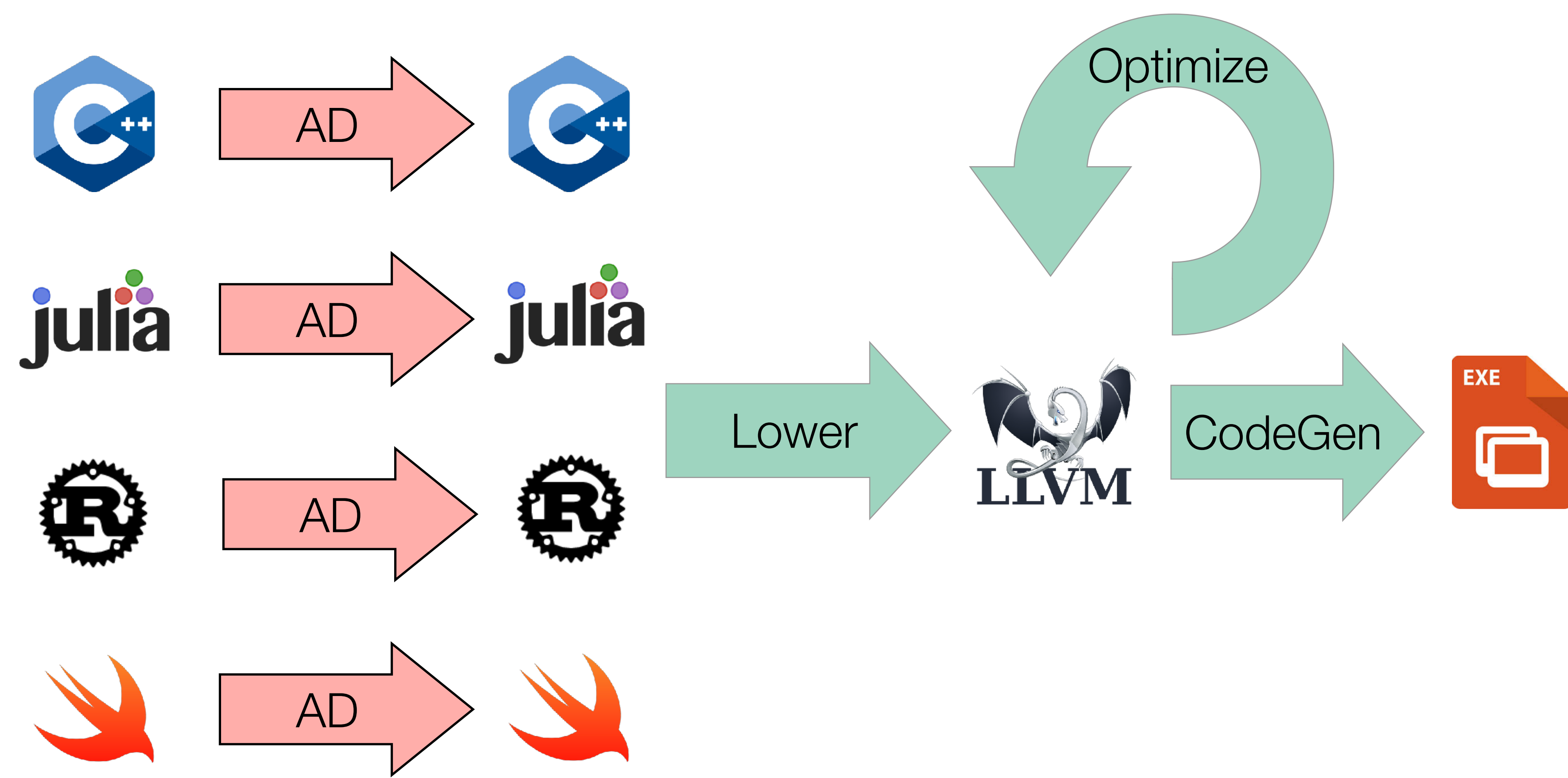


Existing AD Approaches (3/3)

- Source rewriting (Zygote.jl -ish, Tapenade)
 - Statically analyze program to produce a new gradient function in the source language
 - Re-implement parsing and semantics of given language
 - Requires all code to be available ahead of time => hard to use with external libraries



Existing Automatic Differentiation Pipelines



Case Study: Vector Normalization

```
//Compute magnitude in O(n)
double mag(double[] x);

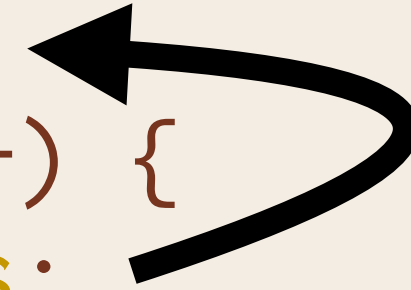
//Compute norm in O(n^2)
void norm(double[] out, double[] in) {

    for (int i=0; i<n; i++) {
        out[i] = in[i] / mag(in);
    }
}
```

Case Study: Vector Normalization

```
//Compute magnitude in O(n)
double mag(double[] x);

//Compute norm in O(n)
void norm(double[] out, double[] in) {
    double res = mag(in);
    for (int i=0; i<n; i++) {
        out[i] = in[i] / res;
    }
}
```



Optimization & Automatic Differentiation

$O(n^2)$

```
for i=0..n {  
  out[i] /= mag(in)  
}
```

Optimize

$O(n)$

```
res = mag(in)  
for i=0..n {  
  out[i] /= res  
}
```

AD

$O(n)$

```
d_res = 0.0  
for i=n..0 {  
  d_res += d_out[i]...  
}  
vmag(d_in, d_res)
```

Optimization & Automatic Differentiation

$O(n^2)$

```
for i=0..n {  
  out[i] /= mag(in)  
}
```

Optimize

$O(n)$

```
res = mag(in)  
for i=0..n {  
  out[i] /= res  
}
```

AD

$O(n)$

```
d_res = 0.0  
for i=n..0 {  
  d_res += d_out[i]...  
}  
∇mag(d_in, d_res)
```

$O(n^2)$

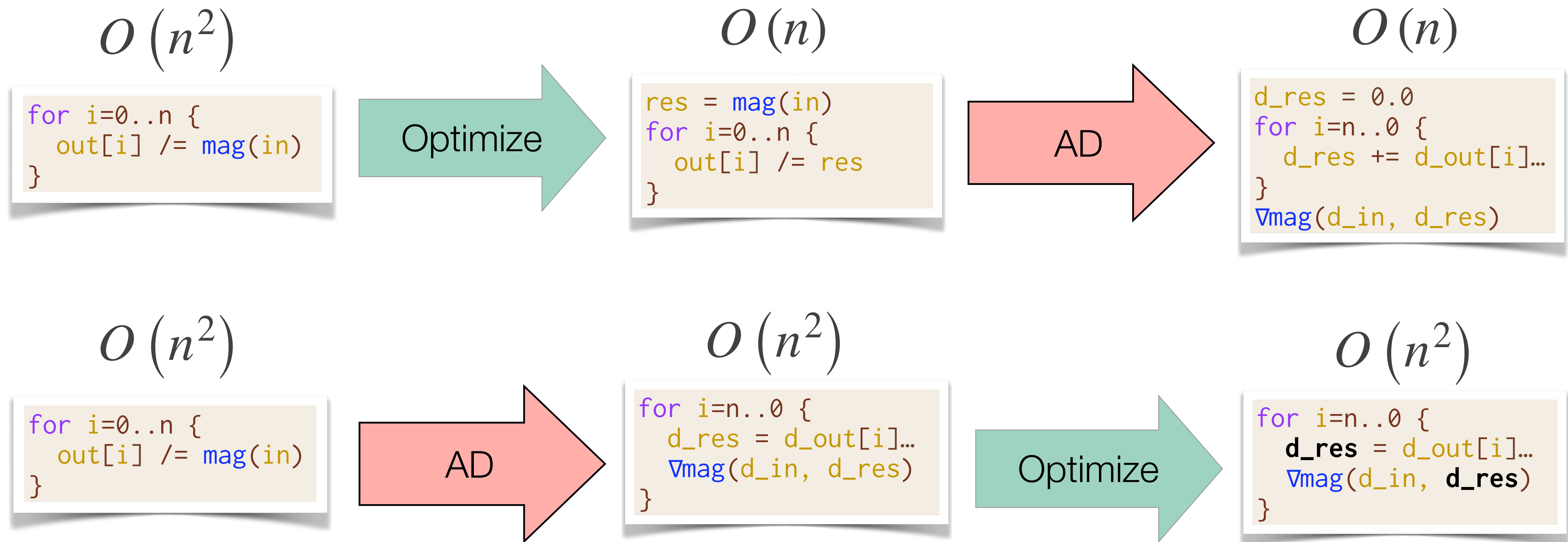
```
for i=0..n {  
  out[i] /= mag(in)  
}
```

AD

$O(n^2)$

```
for i=n..0 {  
  d_res = d_out[i]...  
  ∇mag(d_in, d_res)  
}
```

Optimization & Automatic Differentiation



Optimization & Automatic Differentiation

Differentiating after optimization can create *asymptotically faster* gradients!

$O(n^2)$

```
for i=0..n {  
  out[i] /= mag(in)  
}
```

Optimize

$O(n)$

```
res = mag(in)  
for i=0..n {  
  out[i] /= res  
}
```

AD

$O(n)$

```
d_res = 0.0  
for i=n..0 {  
  d_res += d_out[i]...  
}  
∇mag(d_in, d_res)
```

$O(n^2)$

```
for i=0..n {  
  out[i] /= mag(in)  
}
```

AD

$O(n^2)$

```
for i=n..0 {  
  d_res = d_out[i]...  
  ∇mag(d_in, d_res)  
}
```

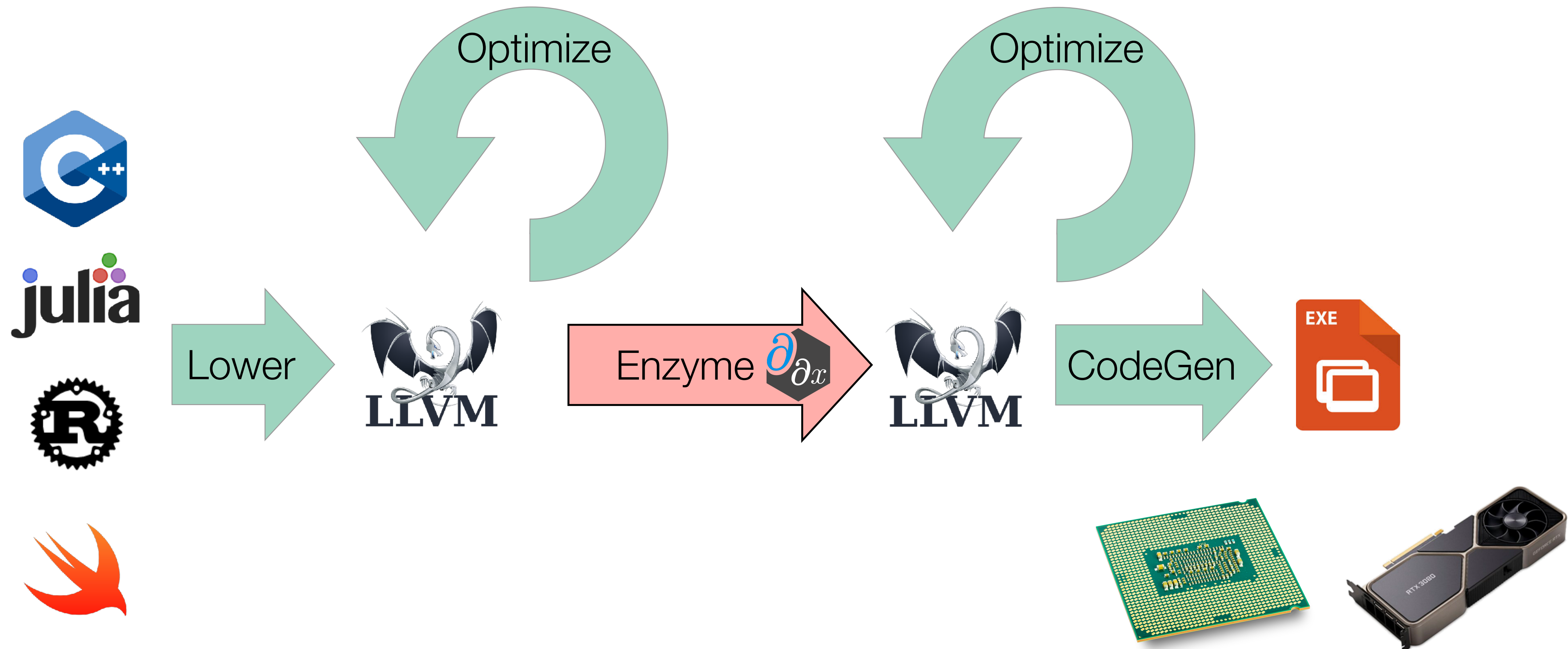
Optimize

$O(n^2)$

```
for i=n..0 {  
  d_res = d_out[i]...  
  ∇mag(d_in, d_res)  
}
```

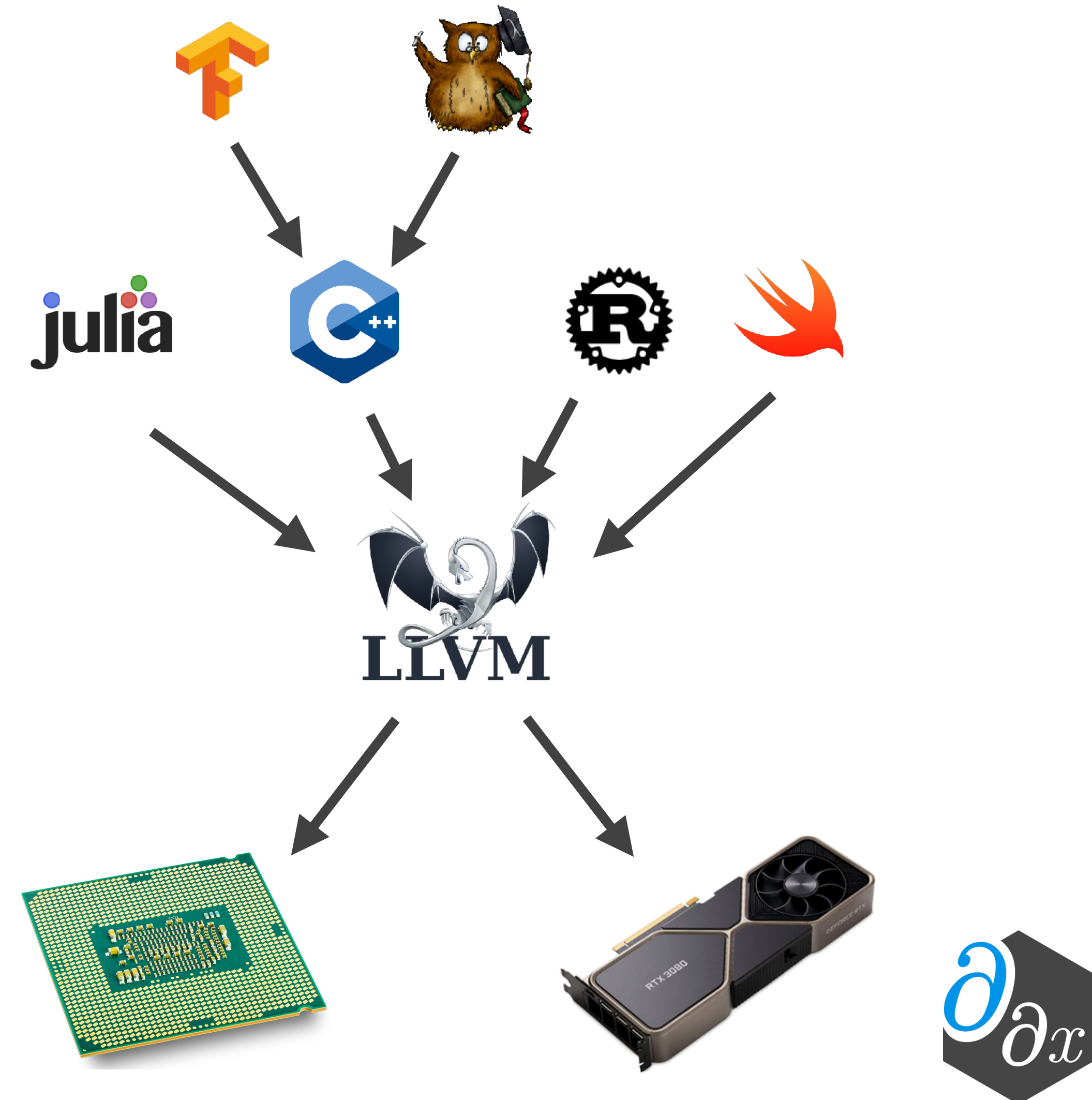
Enzyme Approach

Performing AD at low-level lets us work on ***optimized*** code!



Why Does Enzyme Use LLVM?

- Generic low-level compiler infrastructure with many frontends
 - “Cross platform assembly”
 - Many backends (CPU, CUDA, AMDGPU, etc)
- Well-defined semantics
- Large collection of optimizations and analyses



Case Study: ReLU3

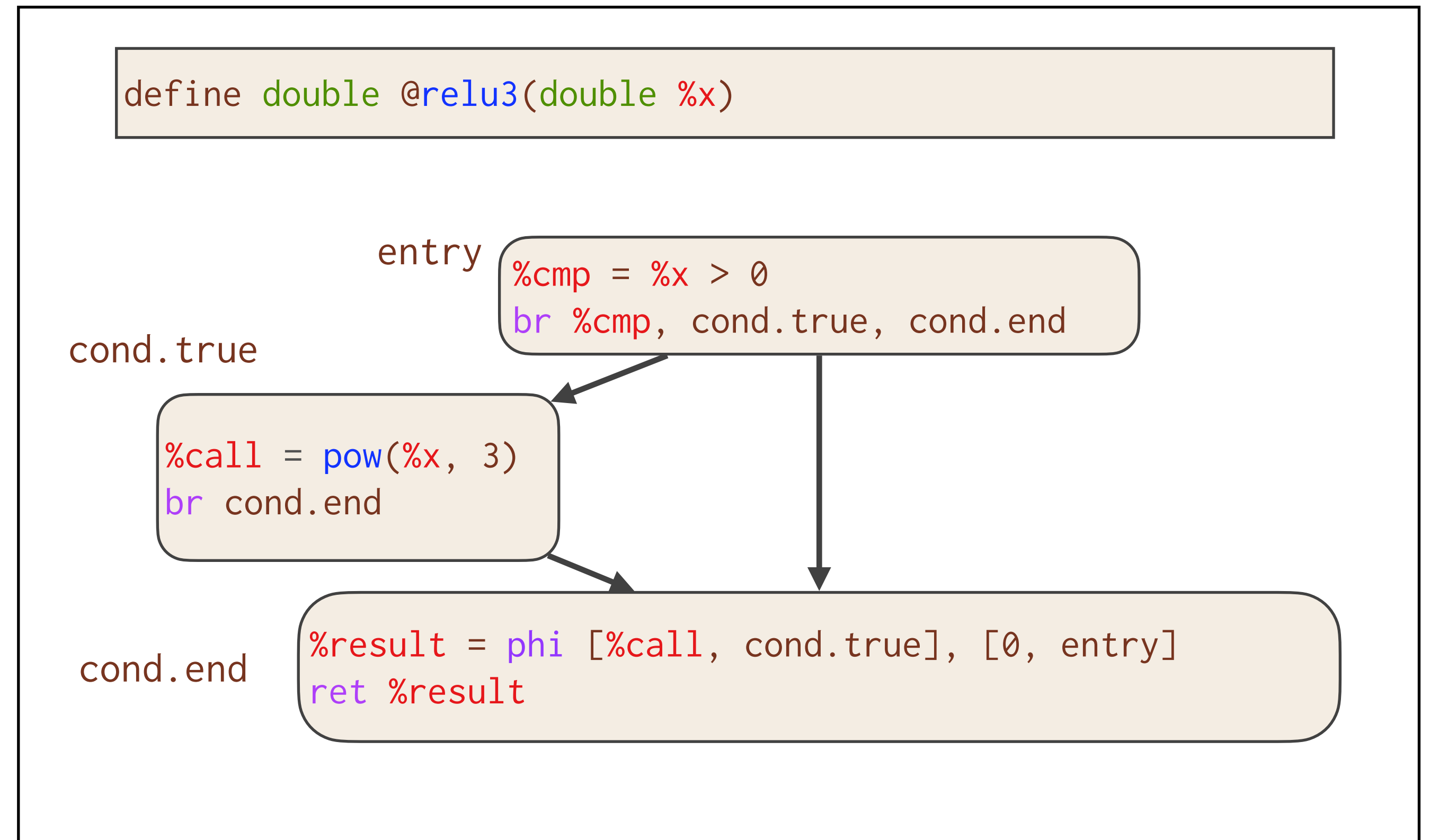
C Source

```
double relu3(double x) {  
    double result;  
    if (x > 0)  
        result = pow(x, 3);  
    else  
        result = 0;  
    return result;  
}
```

Enzyme Usage

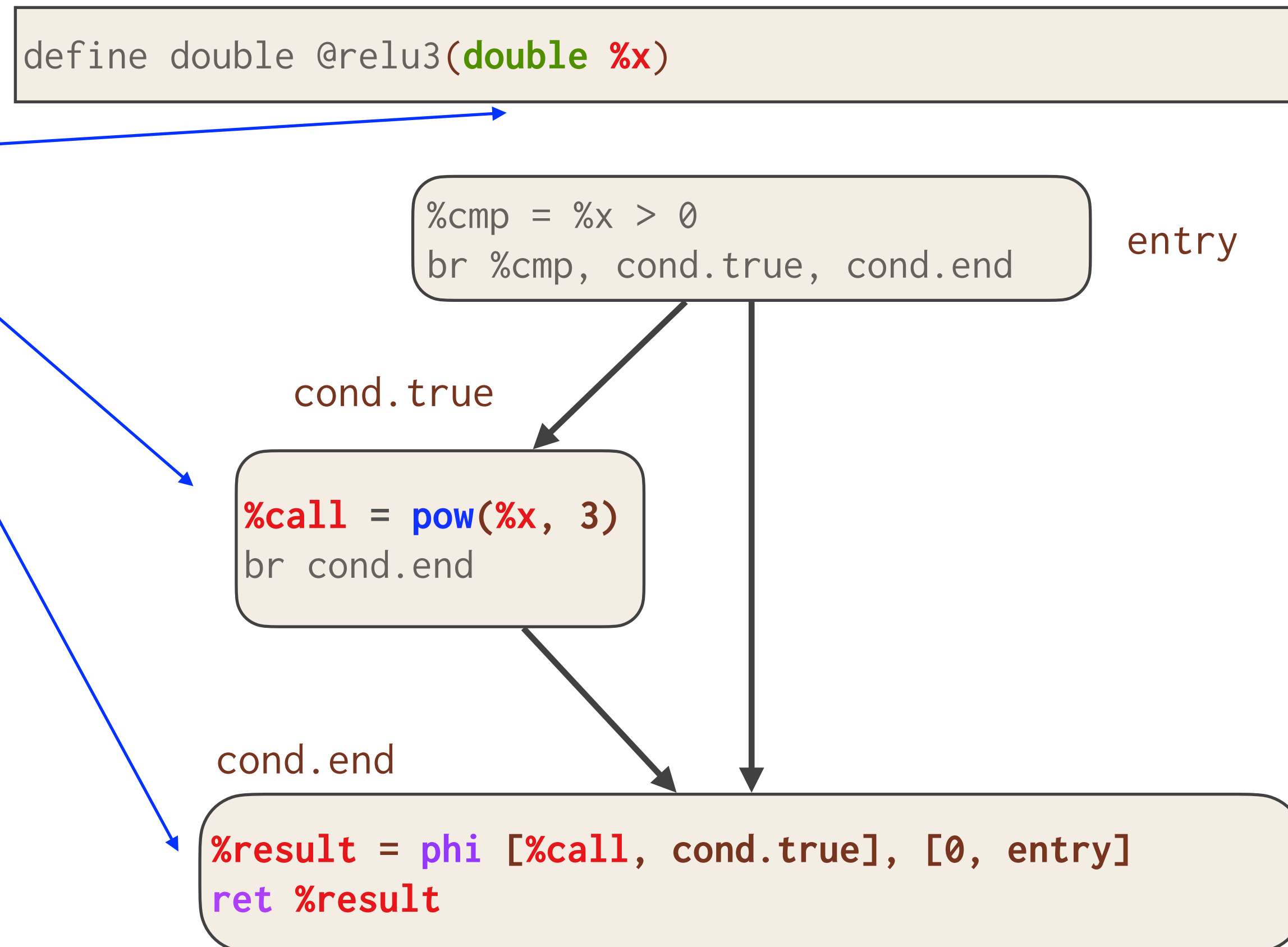
```
double diffe_relu3(double x) {  
    return __enzyme_autodiff(relu3, x);  
}
```

LLVM

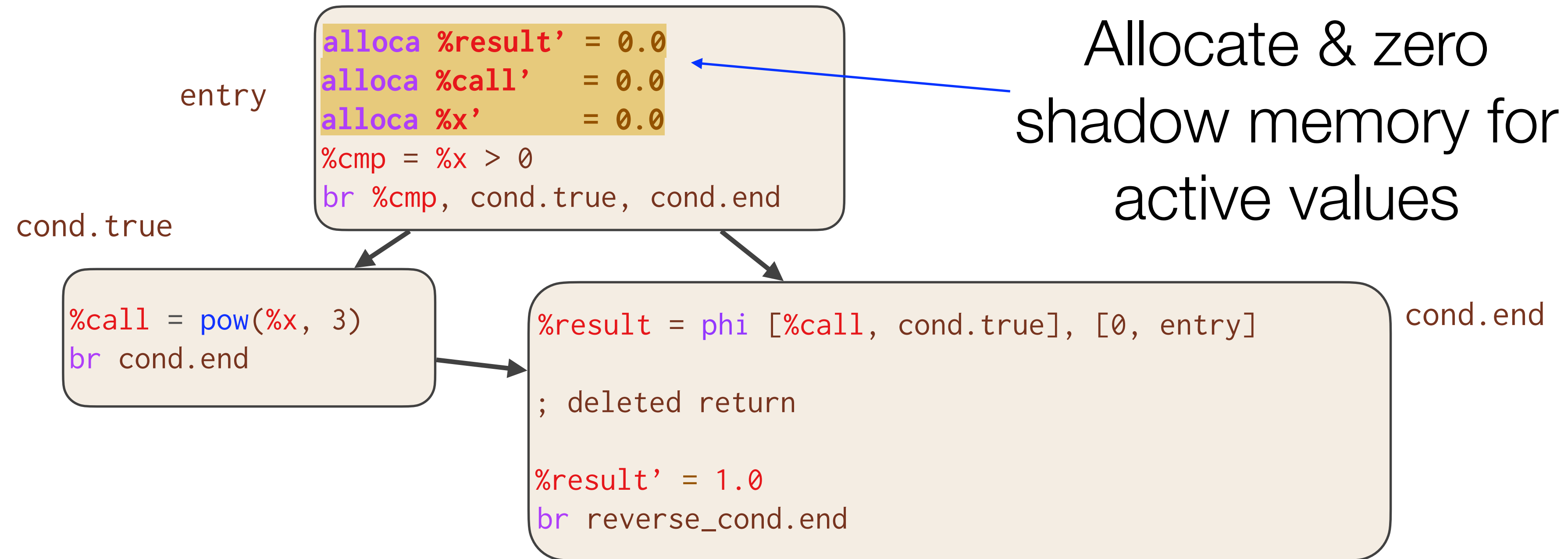


Case Study: ReLU3

Active Instructions

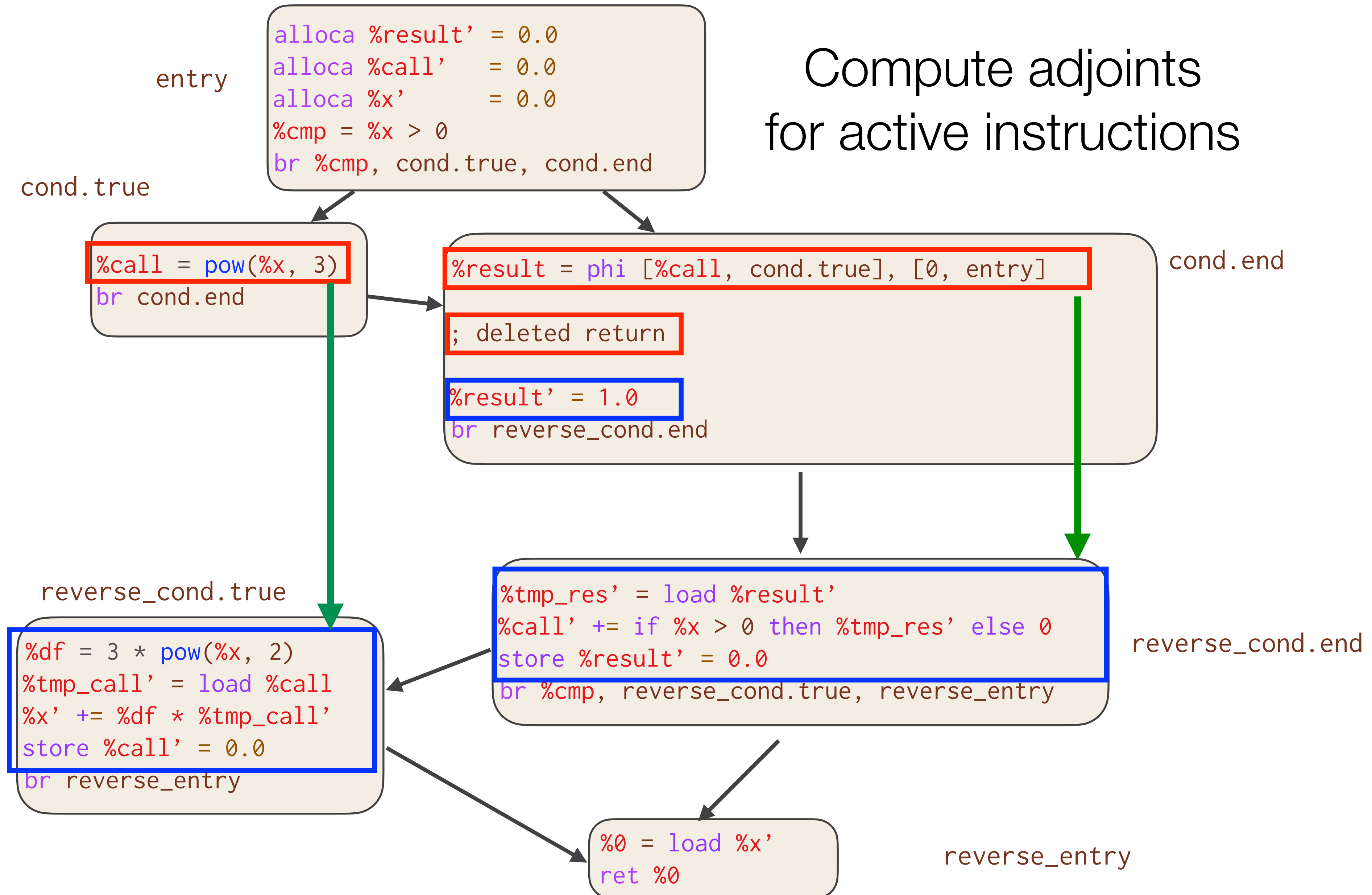


```
define double @diffe_relu3(double %x, double %differet)
```



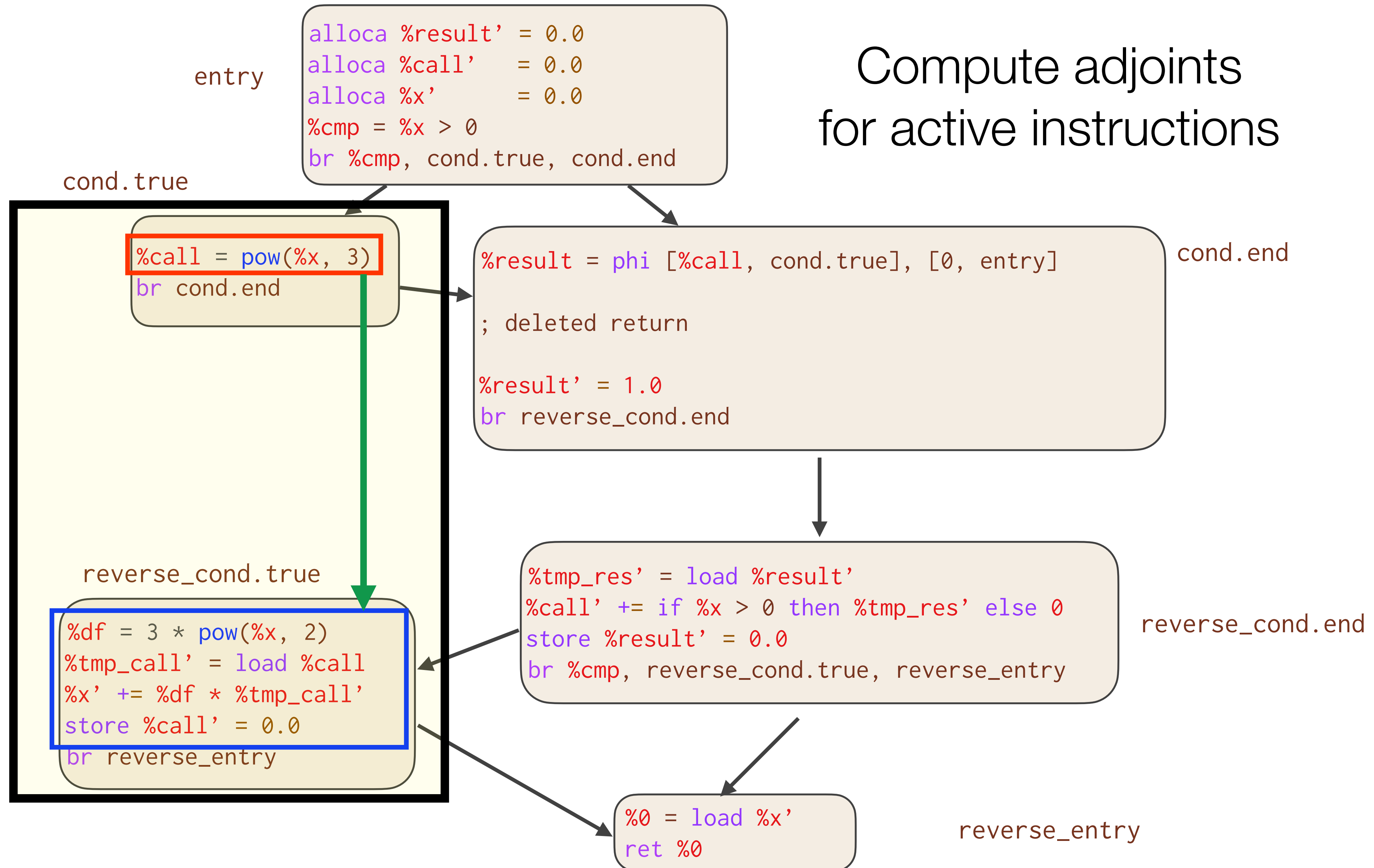
```
define double @diffe_relu3(double %x, double %differet)
```

Compute adjoints
for active instructions



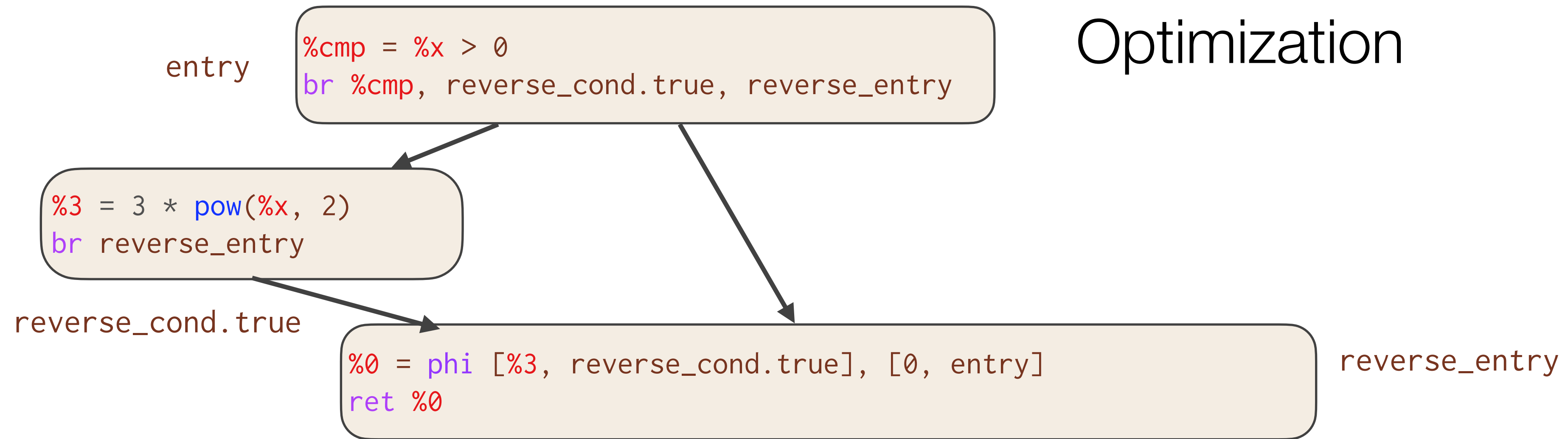
```
define double @diffe_relu3(double %x, double %differet)
```

Compute adjoints
for active instructions



```
define double @diffe_relu3(double %x)
```

Post Optimization

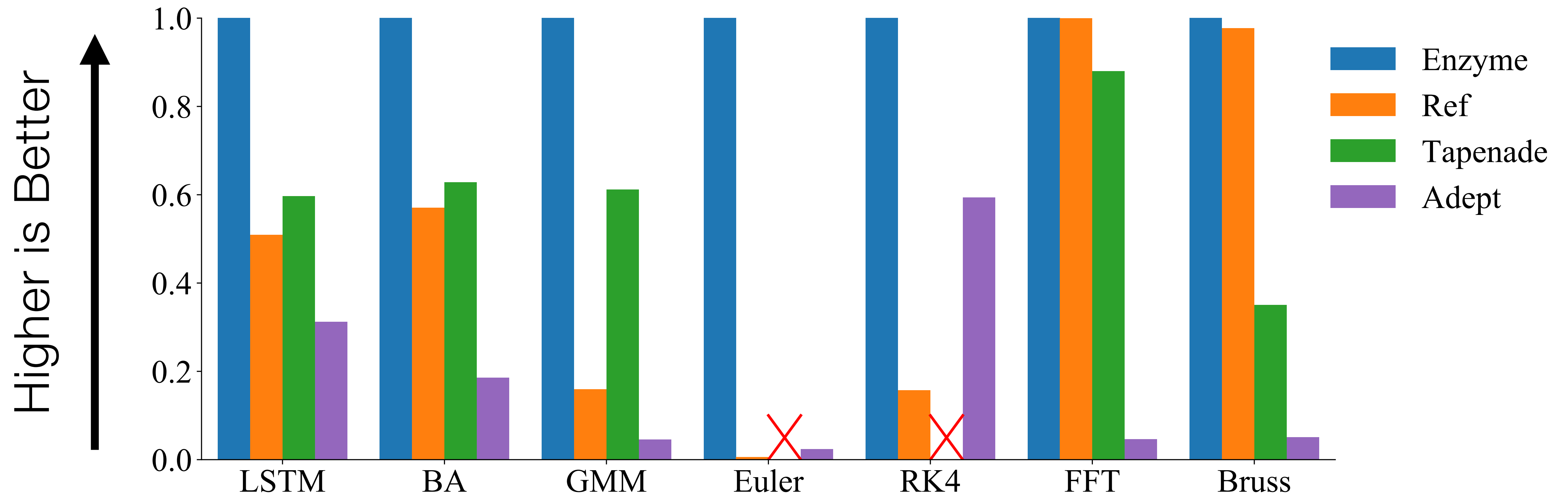


Essentially the optimal hand-written gradient!

```
double diffe_relu3(double x) {  
    double result;  
    if (x > 0)  
        result = 3 * pow(x, 2);  
    else  
        result = 0;  
    return result;  
}
```



Speedup of Enzyme [MC @ NeurIPS 2020]



Enzyme is **4.2x faster** than Reference!

Automatic Differentiation & GPUs [MCPHNSD @ SC'21]

- Prior work has not explored reverse mode AD of existing GPU kernels
 1. Reversing parallel control flow can lead to incorrect results
 2. Complex performance characteristics make it difficult to synthesize efficient code
 3. Resource limitations can prevent kernels from running at all



Challenges of Parallel AD

- The adjoint of an instruction increments the derivative of its input
- Benign read race in forward pass => Write race in reverse pass (undefined behavior)

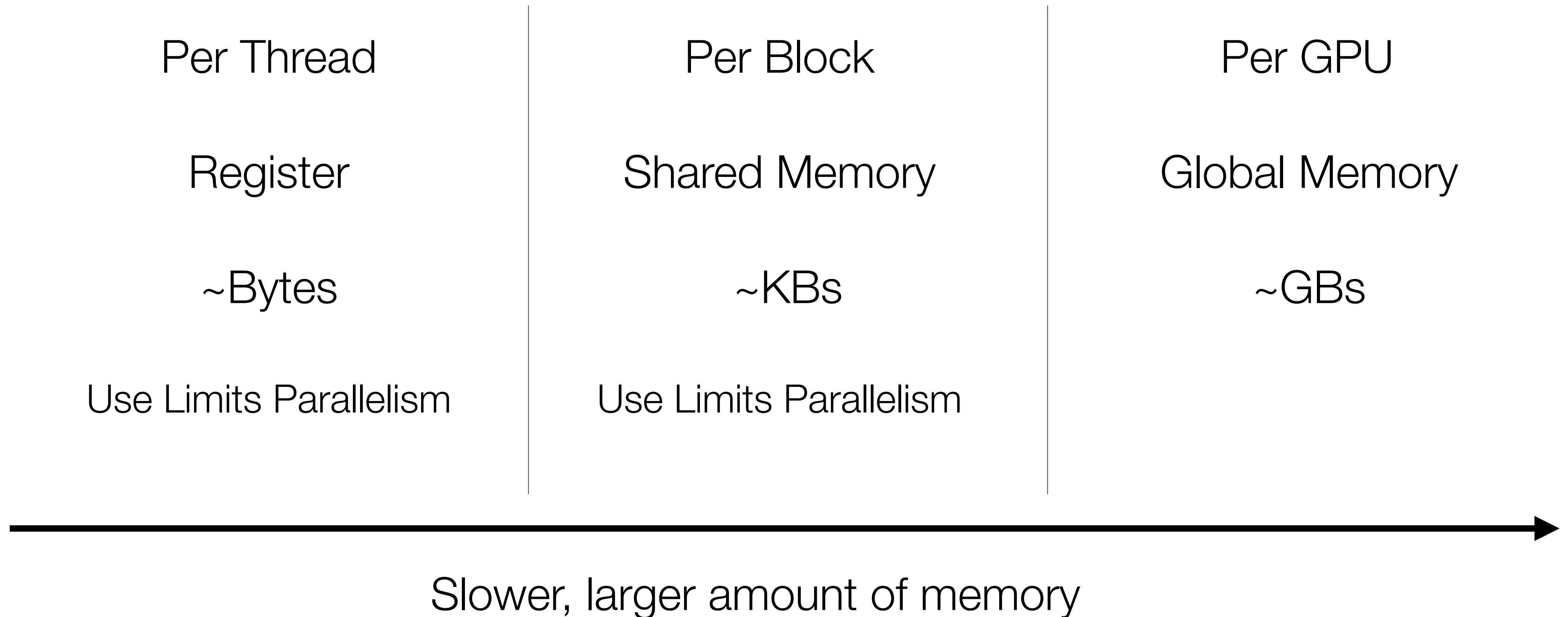
```
void set(double* ar, double val) {  
    parallel_for(int i=0; i<10; i++)  
        ar[i] = val;  
}
```

Read Race

```
double gradient_set(double* ar, double* d_ar,  
                   double val) {  
    double d_val = 0.0;  
    parallel_for(int i=0; i<10; i++)  
        ar[i] = val;  
    parallel_for(int i=0; i<10; i++) {  
        d_val += d_ar[i];  
        d_ar[i] = 0.0;  
    }  
    return d_val;  
}
```

Write Race

GPU Memory Hierarchy



Correct and Efficient Derivative Accumulation

Thread-local memory

- Non-atomic load/store

```
__device__  
void f(...) {  
  
    // Thread-local var  
    double y;  
  
    ...  
  
    d_y += val;  
}
```

Same memory location across
all threads (some shared mem)

- Parallel Reduction

```
// Same var for all threads  
double y;  
  
__device__  
void f(...) {  
  
    ...  
  
    reduce_add(&d_y, val);  
}
```

Others [always legal fallback]

- Atomic increment

```
__device__  
// Unknown thread-aliasing  
void f(double* y) {  
  
    ...  
  
    atomic { d_y += val; }  
}
```

Slower



Synchronization Primitives

- Synchronization (`sync_threads`) ensures all threads finish executing `codeA` before executing `codeB`
- Sync is only necessary if A and B may access to the same memory
- Assuming the original program is race-free, performing a sync at the corresponding location in the reverse ensures correctness
- Prove correctness of algorithm by cases

```
codeA();  
sync_threads;  
codeB();
```

Case 1: Store, Sync, Load

```
codeA(); // store %ptr
sync_threads;

codeB(); // load %ptr
...

diffe_codeB(); // atomicAdd %d_ptr
sync_threads;

diffe_codeA(); // load %d_ptr
                // store %d_ptr = 0
```



Correct

- Load of d_ptr must happen after all atomicAdds have completed

CUDA Example

```
__device__
void inner(float* a, float* x, float* y) {
    y[threadIdx.x] = a[0] * x[threadIdx.x];
}

__device__
void __enzyme_autodiff(void*, ...);

__global__
void daxpy(float* a, float* da,
           float* x, float* dx,
           float* y, float* dy) {
    __enzyme_autodiff((void*)inner,
                      a, da, x, dx, y, dy);
}
```

```
__device__
void diffe_inner(float* a, float* da,
                 float* x, float* dx,
                 float* y, float* dy) {
    // Forward Pass

    y[threadIdx.x] = a[0] * x[threadIdx.x];

    // Reverse Pass

    float dy = dy[threadIdx.x];
    dy[threadIdx.x] = 0.0f;

    float dx_tmp = a[0] * dy;
    atomic { dx[threadIdx.x] += dx_tmp; }

    float da_tmp = x[threadIdx.x] * dy;
    atomic { da[0] += da_tmp; }
}
```



CUDA Example

```
__device__
void inner(float* a, float* x, float* y) {
    y[threadIdx.x] = a[0] * x[threadIdx.x];
}

__device__
void __enzyme_autodiff(void*, ...);

__global__
void daxpy(float* a, float* da,
           float* x, float* dx,
           float* y, float* dy) {
    __enzyme_autodiff((void*)inner,
                      a, da, x, dx, y, dy);
}
```

```
__device__
void diffe_inner(float* a, float* da,
                 float* x, float* dx,
                 float* y, float* dy) {
    // Forward Pass

    y[threadIdx.x] = a[0] * x[threadIdx.x];

    // Reverse Pass

    float dy = dy[threadIdx.x];
    dy[threadIdx.x] = 0.0f;

    float dx_tmp = a[0] * dy;
    dx[threadIdx.x] += dx_tmp;

    float da_tmp = x[threadIdx.x] * dy;
    reduce_accumulate(&da[0], da_tmp);
}
```

Efficient GPU Code

- For correctness, Enzyme may need to cache values in order to compute the gradient
 - The complexity of GPU memory means large caches slow down the program by several orders of magnitude, if it even fits at all
- Like the CPU, existing optimizations reduce the overhead
- Unlike the CPU, existing optimizations aren't sufficient
- Novel GPU and AD-specific optimizations can speedup by several orders of magnitude

```
// Forward Pass
out[i] = x[i] * x[i];
x[i] = 0.0f;

// Reverse (gradient) Pass
...
grad_x[i] += 2 * x[i] * grad_out[i];
...
```

Efficient Correct GPU Code

- For correctness, Enzyme may need to cache values in order to compute the gradient
 - The complexity of GPU memory means large caches slow down the program by several orders of magnitude, if it even fits at all
- Like the CPU, existing optimizations reduce the overhead
- Unlike the CPU, existing optimizations aren't sufficient
- Novel GPU and AD-specific optimizations can speedup by several orders of magnitude

```
double* x_cache = new double[...];

// Forward Pass

out[i] = x[i] * x[i];
x_cache[i] = x[i];

x[i] = 0.0f;

// Reverse (gradient) Pass

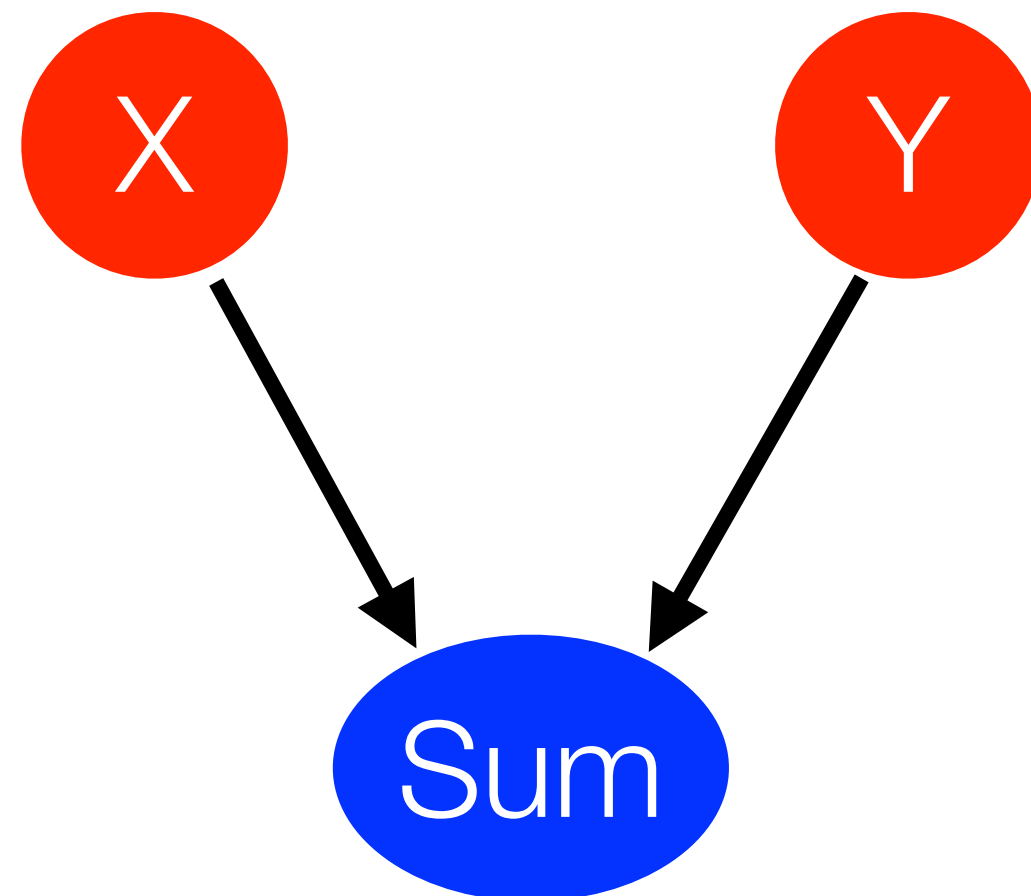
...
grad_x[i] += 2 * x_cache[i]
             * grad_out[i];
...

delete[] x_cache;
```

Cache Reduction Example

- By considering the dataflow graph we can perform a min-cut to approximate smaller cache sizes.

Overwritten:

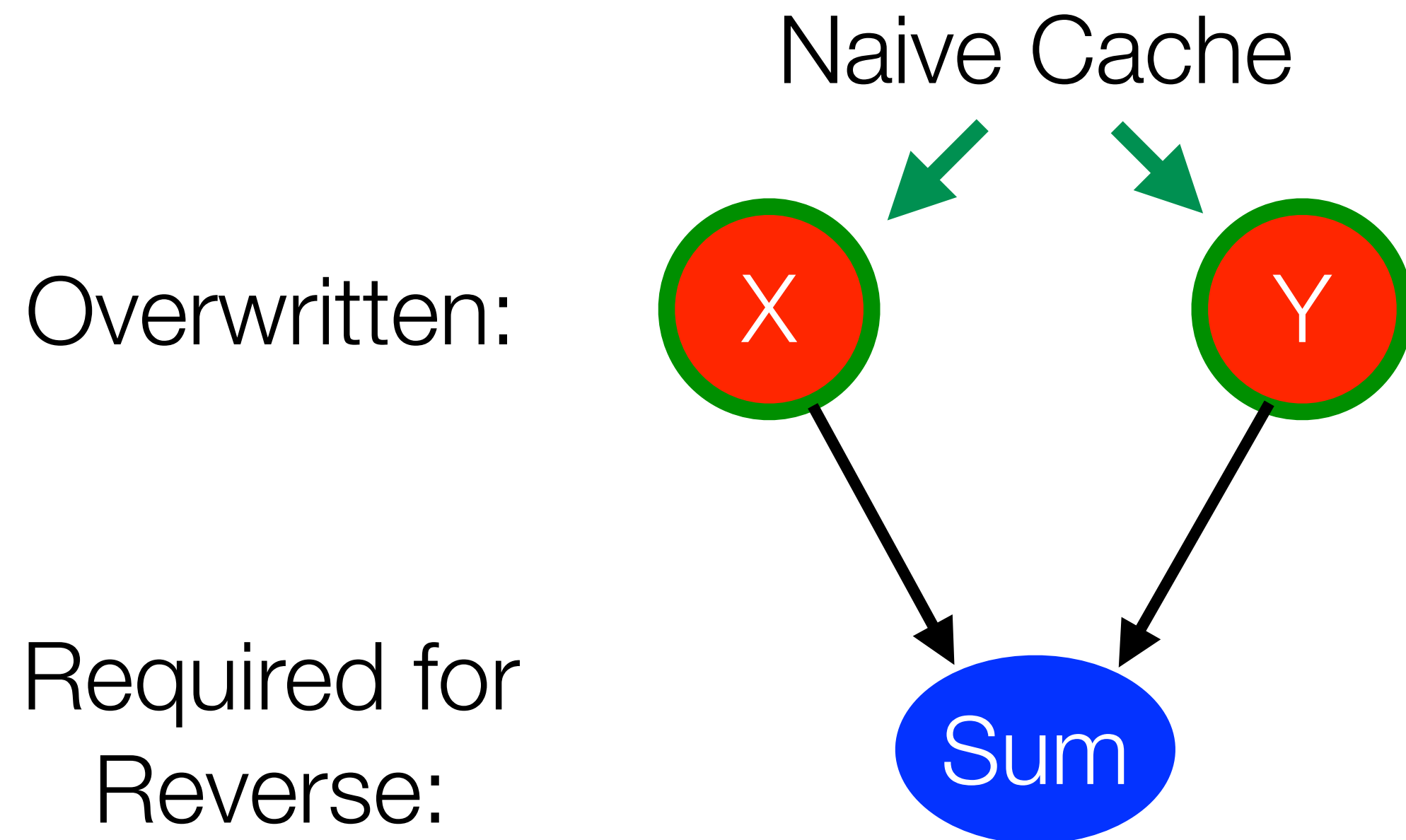


Required for
Reverse:

```
for(int i=0; i<10; i++) {  
    double sum = x[i] + y[i];  
  
    use(sum);  
}  
  
overwrite(x, y);  
grad_overwrite(x, y);  
  
for(int i=9; i>=0; i--) {  
    ...  
    grad_use(sum);  
}
```

Cache Reduction Example

- By considering the dataflow graph we can perform a min-cut to approximate smaller cache sizes.



```
double* x_cache = new double[10];
double* y_cache = new double[10];

for(int i=0; i<10; i++) {
    double sum = x[i] + y[i];
    x_cache[i] = x[i];
    y_cache[i] = y[i];
    use(sum);
}

overwrite(x, y);
grad_overwrite(x, y);

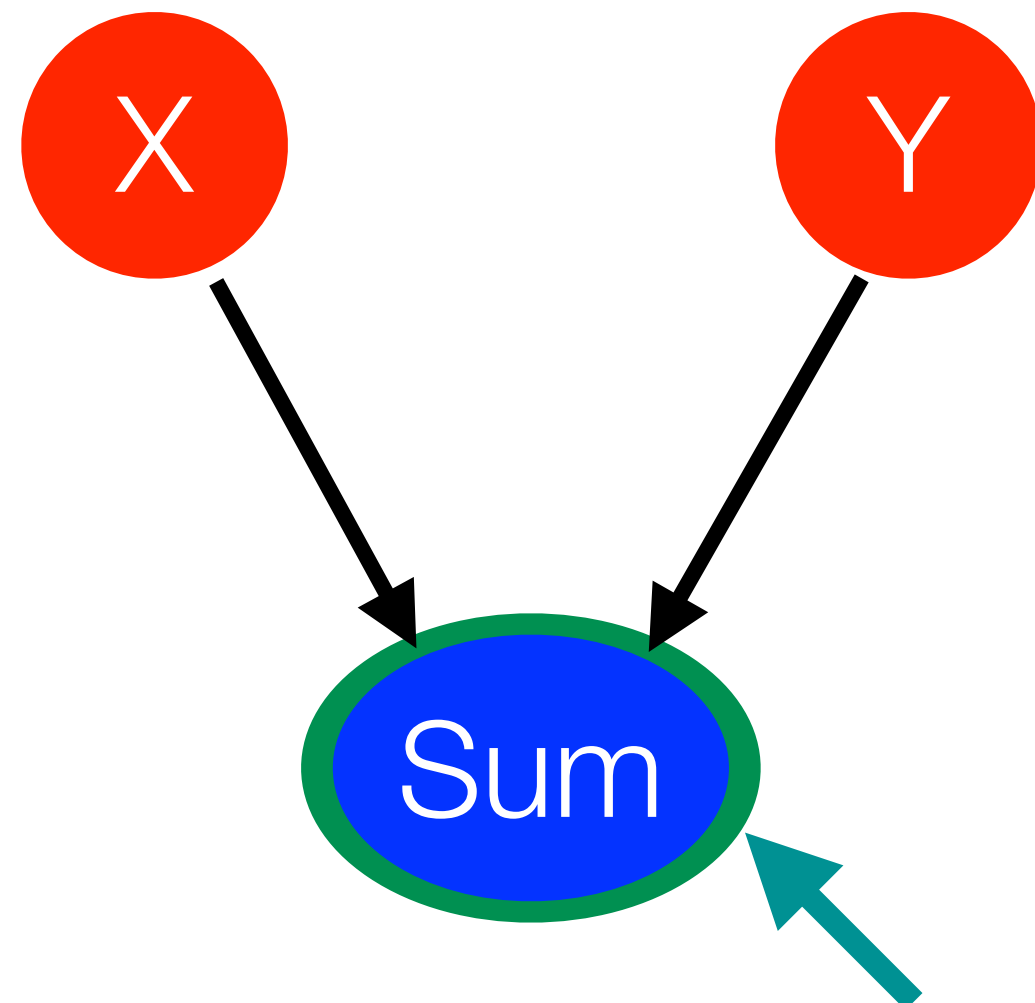
for(int i=9; i>=0; i--) {
    double sum = x_cache[i] + y_cache[i];
    grad_use(sum);
}
```

Cache Reduction Example

- By considering the dataflow graph we can perform a min-cut to approximate smaller cache sizes.

Overwritten:

Required for
Reverse:



Smallest Cache

```
double* sum_cache = new double[10];

for(int i=0; i<10; i++) {
    double sum = x[i] + y[i];
    sum_cache[i] = sum;

    use(sum);
}

overwrite(x, y);
grad_overwrite(x, y);

for(int i=9; i>=0; i--) {
    grad_use(sum_cache[i]);
}
```

Allocation Merging

- Allocations (and any calls) on the GPU are expensive
- Given two allocations in the same scope, replace uses with a single allocation
- Beneficial for not just AD, but any GPU programs!

```
double* var1 = new double[N];  
double* var2 = new double[M];  
  
use(var1, var2);  
  
delete[] var1;  
delete[] var2;
```

```
double* var1 = new double[N + M];  
double* var2 = var1 + N;  
  
use(var1, var2);  
  
delete[] var1;
```

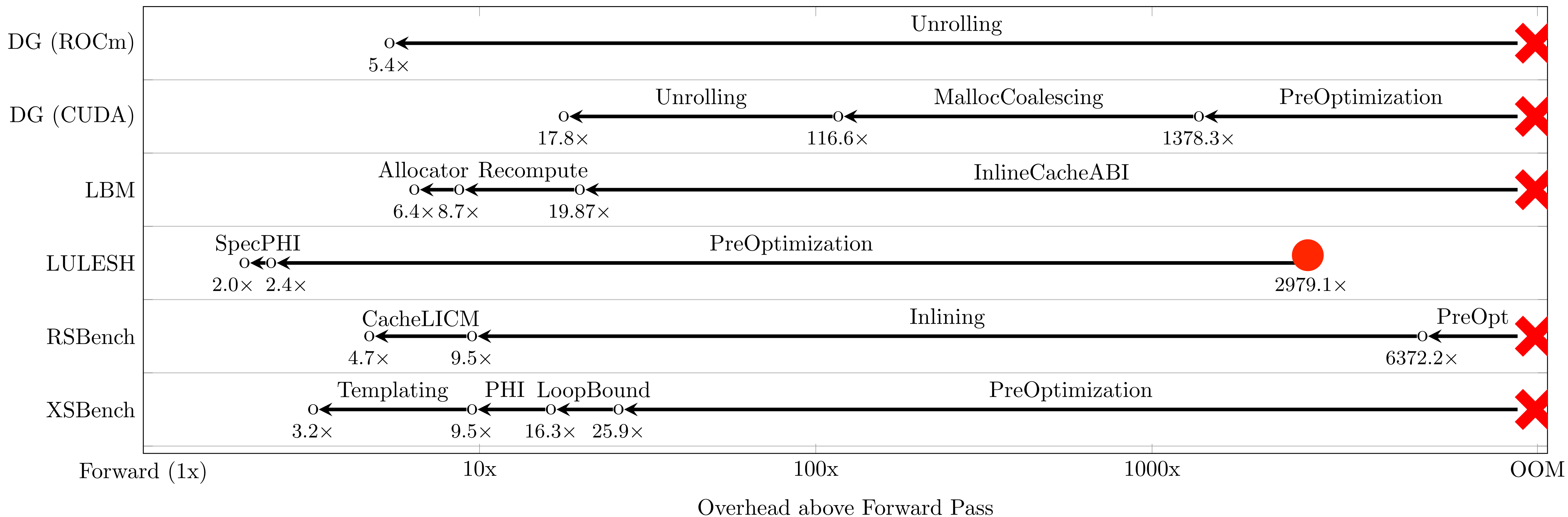
Novel AD + GPU Optimizations

- See our SC'21 paper for more (<https://c.wsmoses.com/papers/EnzymeGPU.pdf>)
Reverse-Mode Automatic Differentiation and Optimization of GPU Kernels via Enzyme. SC, 2021
- [AD] Cache LICM/CSE
- [AD] Min-Cut Cache Reduction
- [AD] Cache Forwarding
- [GPU] Merge Allocations
- [GPU] Heap-to-stack (and register)
- [GPU] Alias Analysis Properties of SyncThreads

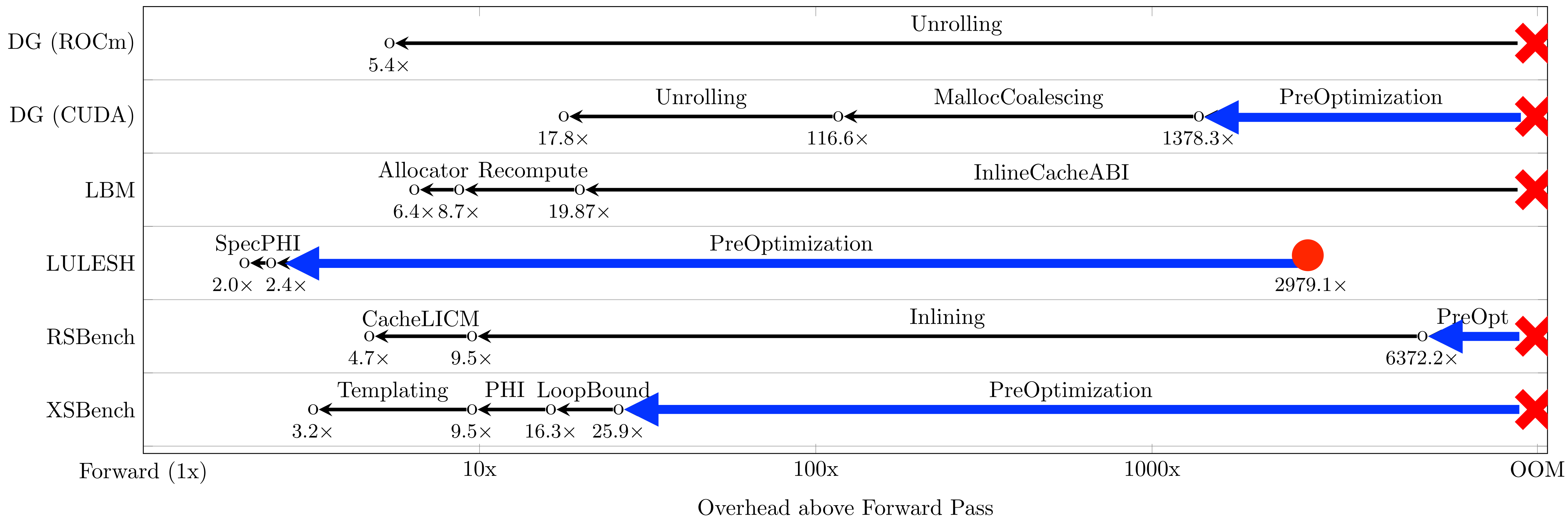
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...

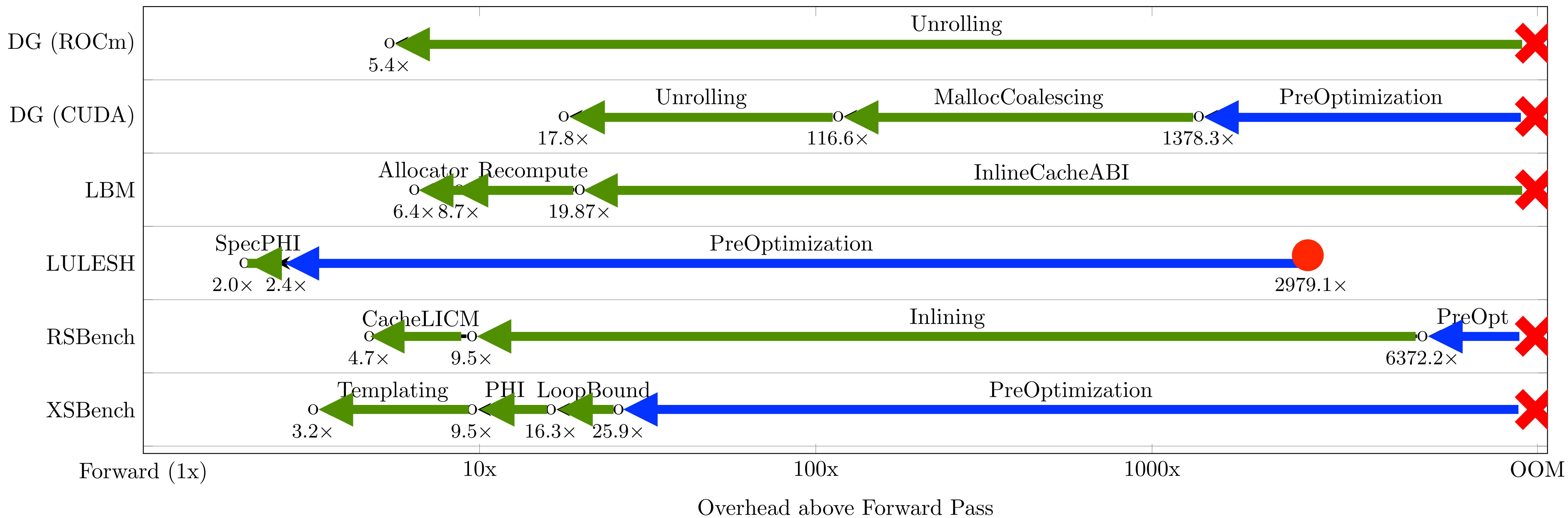
Ablation Analysis of Optimizations



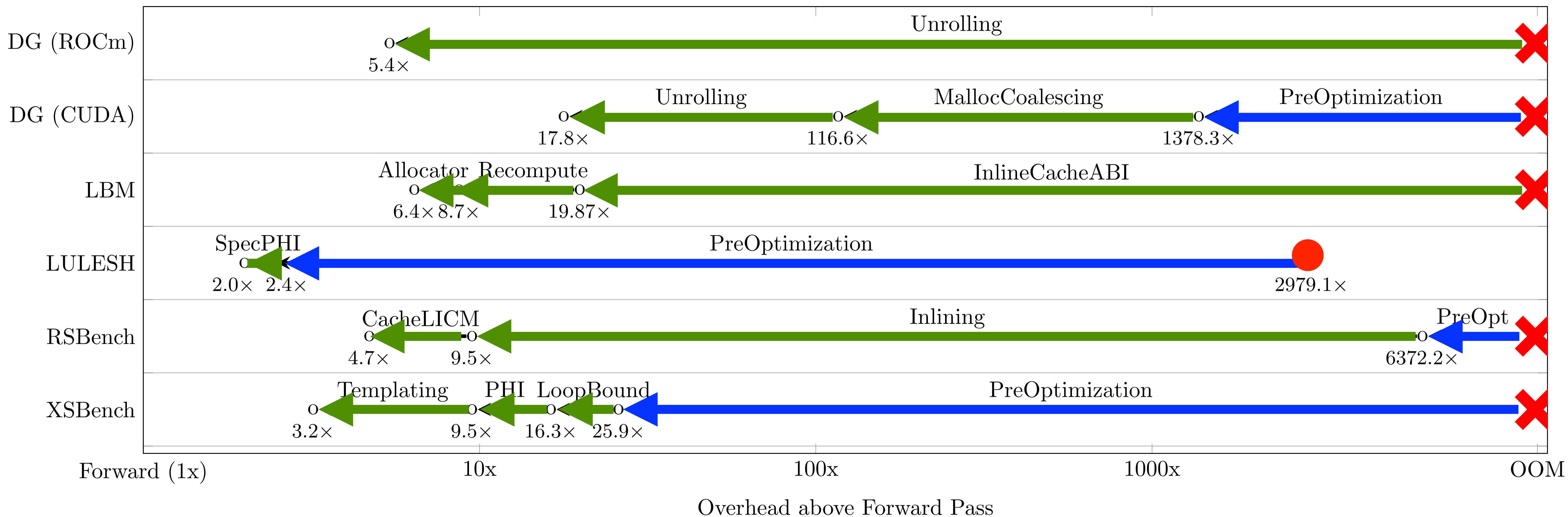
Ablation Analysis of Optimizations



Ablation Analysis of Optimizations



Ablation Analysis of Optimizations



GPU AD is Intractable Without Optimization!

CUDA.jl / AMDGPU.jl Example

```
function compute!(inp, out)
    s_D = @cuStaticSharedMem eltype(inp) (10, 10)
    ...
end

function grad_compute!(inp, out)
    Enzyme.autodiff_deferred(compute!, inp, out)
    return nothing
end

@cuda grad_compute!(Duplicated(inp, d_inp),
                    Duplicated(out, d_out))
```

```
function compute!(inp, out)
    s_D = AMDGPU.alloc_special(...)
    ...
end

function grad_compute!(inp, out)
    Enzyme.autodiff_deferred(compute!, inp, out)
    return nothing
end

@rocm grad_compute!(Duplicated(inp, d_inp),
                    Duplicated(out, d_out))
```

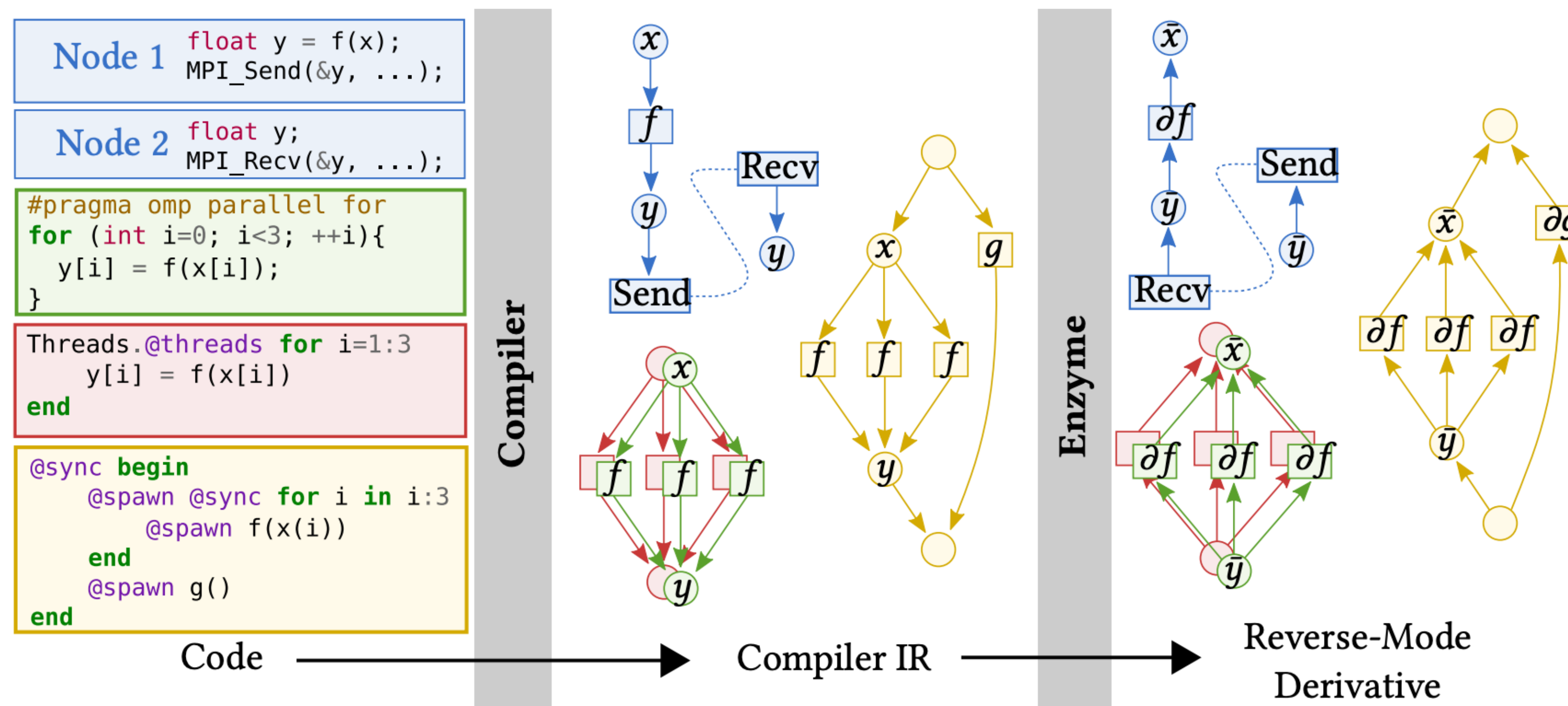
See Below For Full Code Examples

<https://github.com/wsmoses/Enzyme-GPU-Tests/blob/main/DG/>

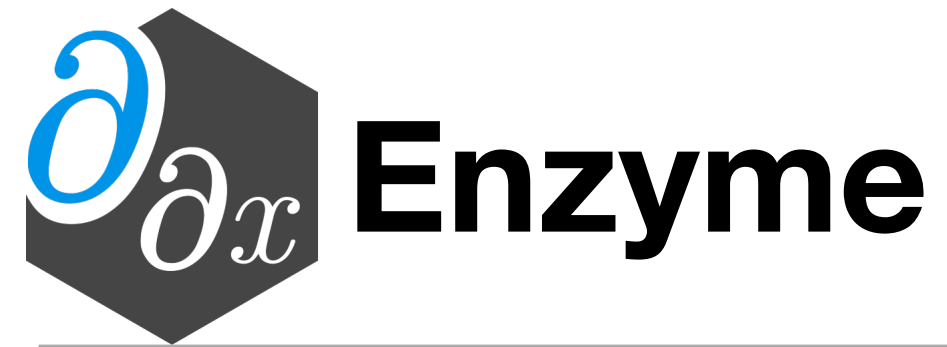


Common Framework for Parallel AD [SC'22]

- Common infrastructure for supporting parallel AD (caching, race-resolution, gradient accumulation) enables parallel differentiation independent of framework or language.



- Enables differentiation of a combination of GPU (e.g. CUDA, ROCm), CPU (OpenMP, Julia Tasks, RAJA), Distributed (MPI, MPI.jl), and more



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Case 1: Store, Sync, Load

```
codeA(); // store %ptr
sync_threads;

codeB(); // load %ptr
...

diffe_codeB(); // atomicAdd %d_ptr
sync_threads;

diffe_codeA(); // load %d_ptr
                // store %d_ptr = 0
```



Correct

- Load of d_ptr must happen after all atomicAdds have completed

CUDA Example

```
__device__
void inner(float* a, float* x, float* y) {
    y[threadIdx.x] = a[0] * x[threadIdx.x];
}

__device__
void __enzyme_autodiff(void*, ...);

__global__
void daxpy(float* a, float* da,
           float* x, float* dx,
           float* y, float* dy) {
    __enzyme_autodiff((void*)inner,
                      a, da, x, dx, y, dy);
}
```

```
__device__
void diffe_inner(float* a, float* da,
                 float* x, float* dx,
                 float* y, float* dy) {
    // Forward Pass

    y[threadIdx.x] = a[0] * x[threadIdx.x];

    // Reverse Pass

    float dy = dy[threadIdx.x];
    dy[threadIdx.x] = 0.0f;

    float dx_tmp = a[0] * dy;
    atomic { dx[threadIdx.x] += dx_tmp; }

    float da_tmp = x[threadIdx.x] * dy;
    atomic { da[0] += da_tmp; }
}
```



CUDA Example

```
__device__
void inner(float* a, float* x, float* y) {
    y[threadIdx.x] = a[0] * x[threadIdx.x];
}

__device__
void __enzyme_autodiff(void*, ...);

__global__
void daxpy(float* a, float* da,
           float* x, float* dx,
           float* y, float* dy) {
    __enzyme_autodiff((void*)inner,
                      a, da, x, dx, y, dy);
}
```

```
__device__
void diffe_inner(float* a, float* da,
                 float* x, float* dx,
                 float* y, float* dy) {
    // Forward Pass

    y[threadIdx.x] = a[0] * x[threadIdx.x];

    // Reverse Pass

    float dy = dy[threadIdx.x];
    dy[threadIdx.x] = 0.0f;

    float dx_tmp = a[0] * dy;
    dx[threadIdx.x] += dx_tmp;

    float da_tmp = x[threadIdx.x] * dy;
    reduce_accumulate(&da[0], da_tmp);
}
```



Efficient GPU Code

- For correctness, Enzyme may need to cache values in order to compute the gradient
 - The complexity of GPU memory means large caches slow down the program by several orders of magnitude, if it even fits at all
- Like the CPU, existing optimizations reduce the overhead
- Unlike the CPU, existing optimizations aren't sufficient
- Novel GPU and AD-specific optimizations can speedup by several orders of magnitude

```
// Forward Pass
out[i] = x[i] * x[i];
x[i] = 0.0f;

// Reverse (gradient) Pass
...
grad_x[i] += 2 * x[i] * grad_out[i];
...
```



Efficient Correct GPU Code

- For correctness, Enzyme may need to cache values in order to compute the gradient
 - The complexity of GPU memory means large caches slow down the program by several orders of magnitude, if it even fits at all
- Like the CPU, existing optimizations reduce the overhead
- Unlike the CPU, existing optimizations aren't sufficient
- Novel GPU and AD-specific optimizations can speedup by several orders of magnitude

```
double* x_cache = new double[...];

// Forward Pass

out[i] = x[i] * x[i];
x_cache[i] = x[i];

x[i] = 0.0f;

// Reverse (gradient) Pass

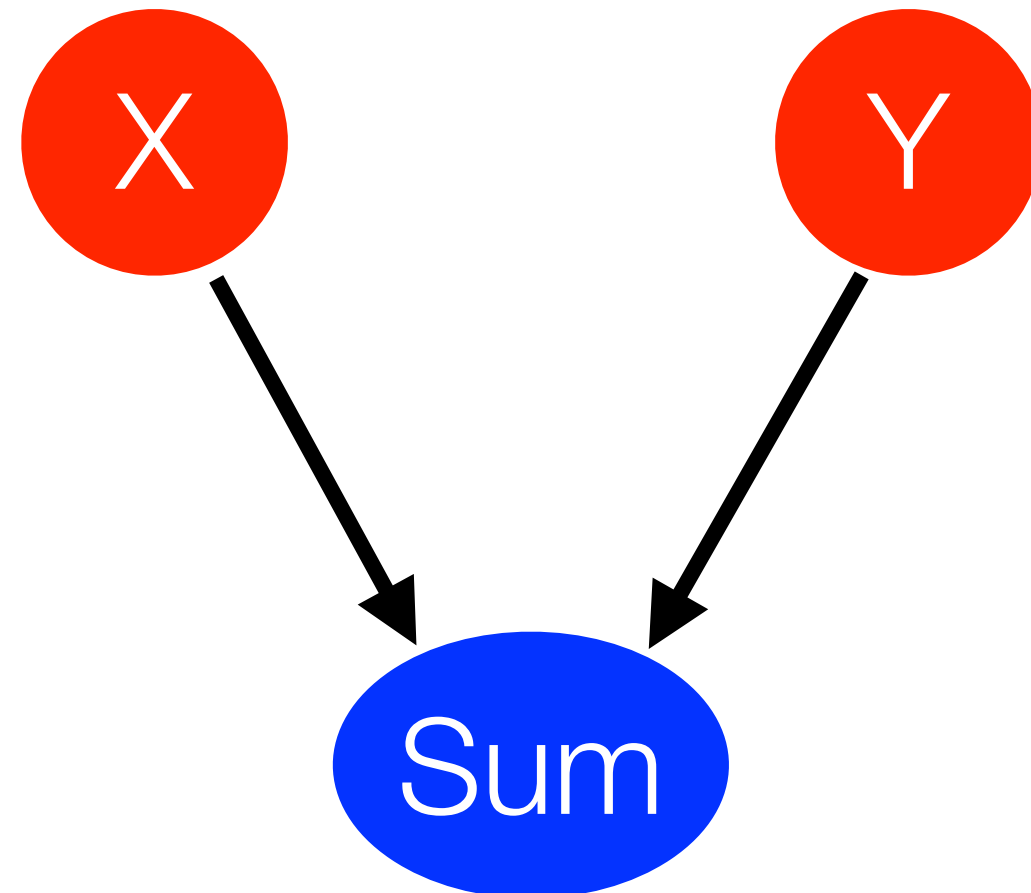
...
grad_x[i] += 2 * x_cache[i]
             * grad_out[i];
...

delete[] x_cache;
```

Cache Reduction Example

- By considering the dataflow graph we can perform a min-cut to approximate smaller cache sizes.

Overwritten:

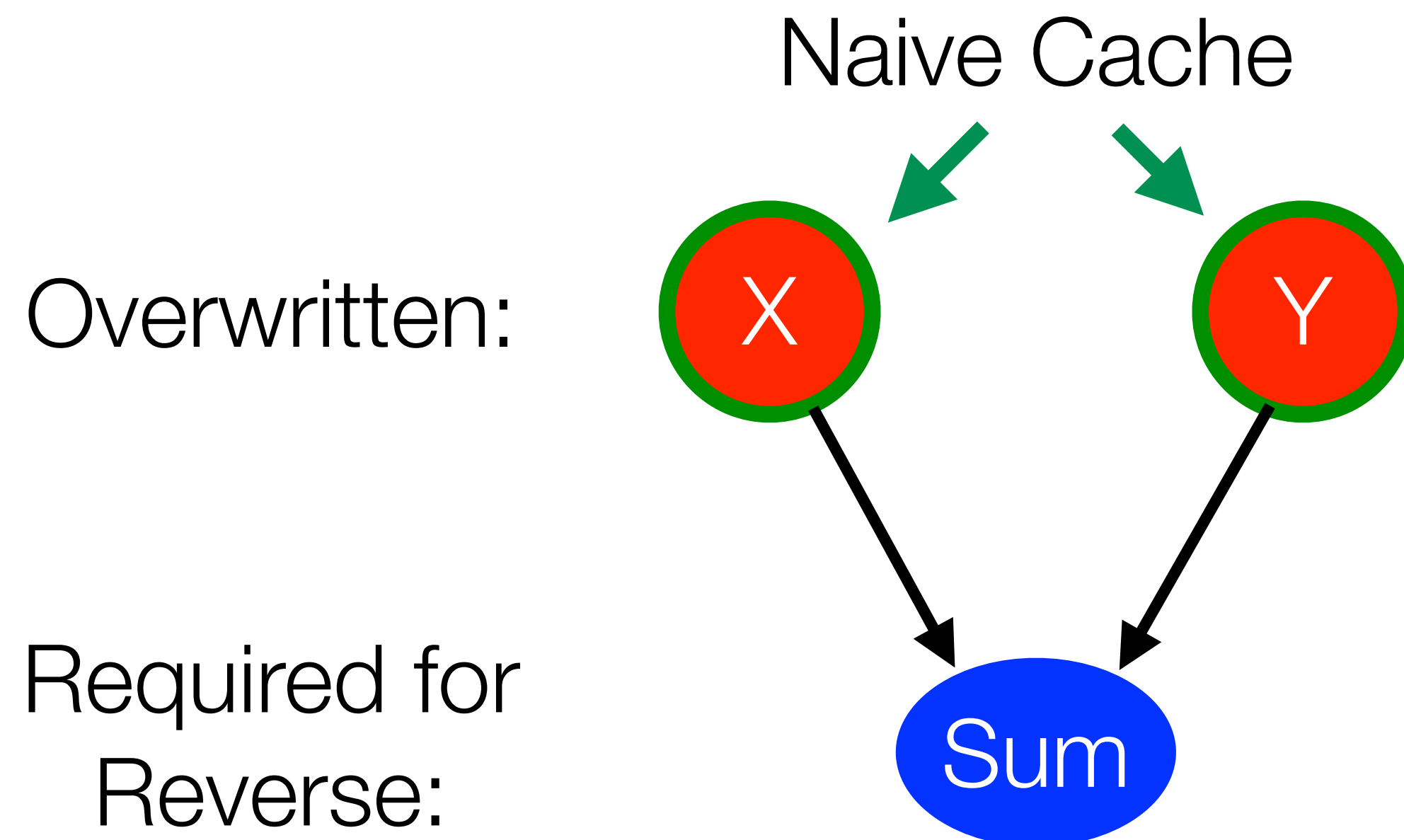


Required for
Reverse:

```
for(int i=0; i<10; i++) {  
    double sum = x[i] + y[i];  
  
    use(sum);  
}  
  
overwrite(x, y);  
grad_overwrite(x, y);  
  
for(int i=9; i>=0; i--) {  
    ...  
    grad_use(sum);  
}
```

Cache Reduction Example

- By considering the dataflow graph we can perform a min-cut to approximate smaller cache sizes.



```
double* x_cache = new double[10];
double* y_cache = new double[10];

for(int i=0; i<10; i++) {
    double sum = x[i] + y[i];
    x_cache[i] = x[i];
    y_cache[i] = y[i];
    use(sum);
}

overwrite(x, y);
grad_overwrite(x, y);

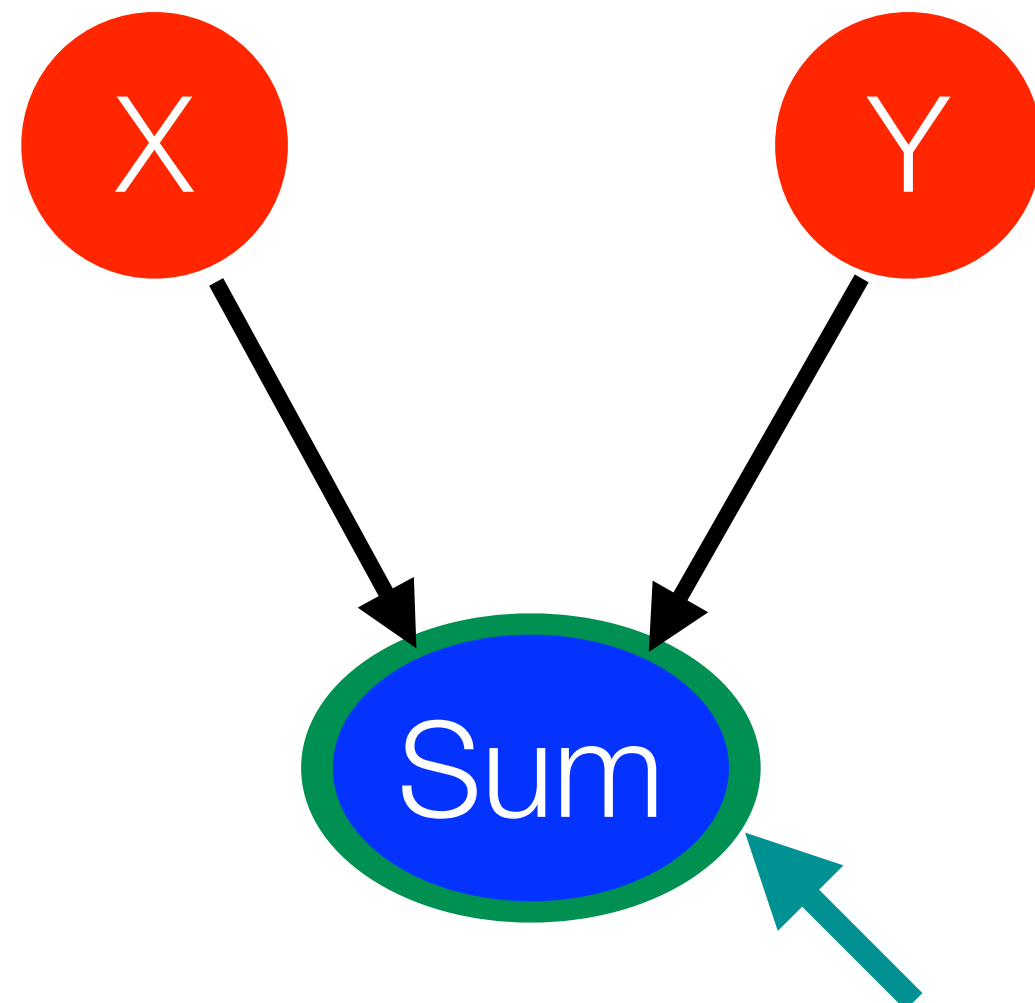
for(int i=9; i>=0; i--) {
    double sum = x_cache[i] + y_cache[i];
    grad_use(sum);
}
```

Cache Reduction Example

- By considering the dataflow graph we can perform a min-cut to approximate smaller cache sizes.

Overwritten:

Required for
Reverse:



Smallest Cache

```
double* sum_cache = new double[10];

for(int i=0; i<10; i++) {
    double sum = x[i] + y[i];
    sum_cache[i] = sum;

    use(sum);
}

overwrite(x, y);
grad_overwrite(x, y);

for(int i=9; i>=0; i--) {
    grad_use(sum_cache[i]);
}
```

Allocation Merging

- Allocations (and any calls) on the GPU are expensive
- Given two allocations in the same scope, replace uses with a single allocation
- Beneficial for not just AD, but any GPU programs!

```
double* var1 = new double[N];  
double* var2 = new double[M];  
  
use(var1, var2);  
  
delete[] var1;  
delete[] var2;
```

```
double* var1 = new double[N + M];  
double* var2 = var1 + N;  
  
use(var1, var2);  
  
delete[] var1;
```

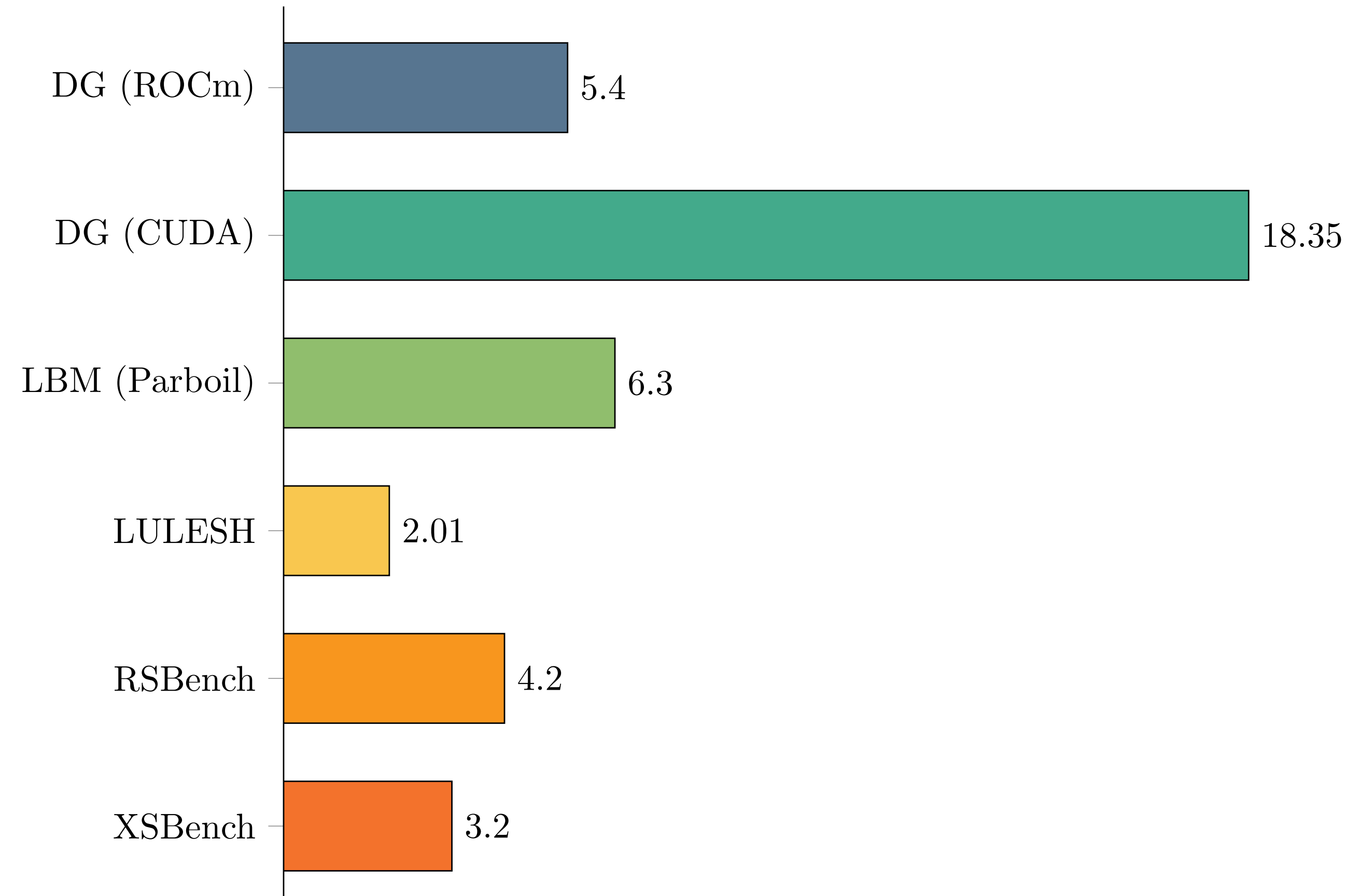
Novel AD + GPU Optimizations

- See our SC paper (Nov 17) for more (<https://c.wsmoses.com/papers/EnzymeGPU.pdf>)
Reverse-Mode Automatic Differentiation and Optimization of GPU Kernels via Enzyme. SC, 2021
- [AD] Cache LICM/CSE
- [AD] Min-Cut Cache Reduction
- [AD] Cache Forwarding
- [GPU] Merge Allocations
- [GPU] Heap-to-stack (and register)
- [GPU] Alias Analysis Properties of SyncThreads
- ...



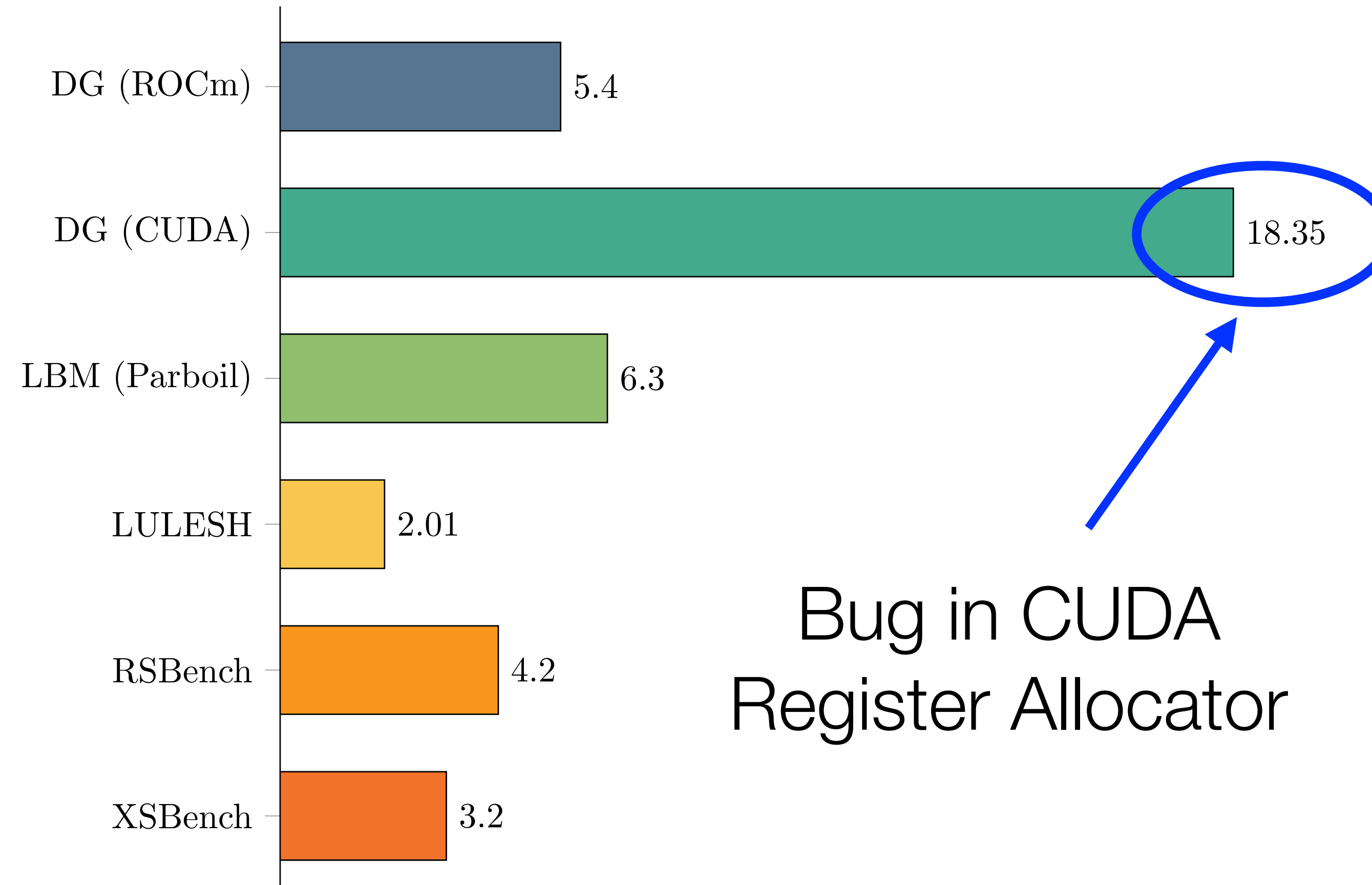
GPU Gradient Overhead [MCPHNMJ'21]

- Evaluation of both original code and gradient
 - DG: Discontinuous-Galerkin integral (Julia)
 - LBM: particle-based fluid dynamics simulation
 - LULESH: unstructured explicit shock hydrodynamics solver
 - XSBench & RSBench: Monte Carlo simulations of particle transport algorithms (memory & compute bound, respectively)

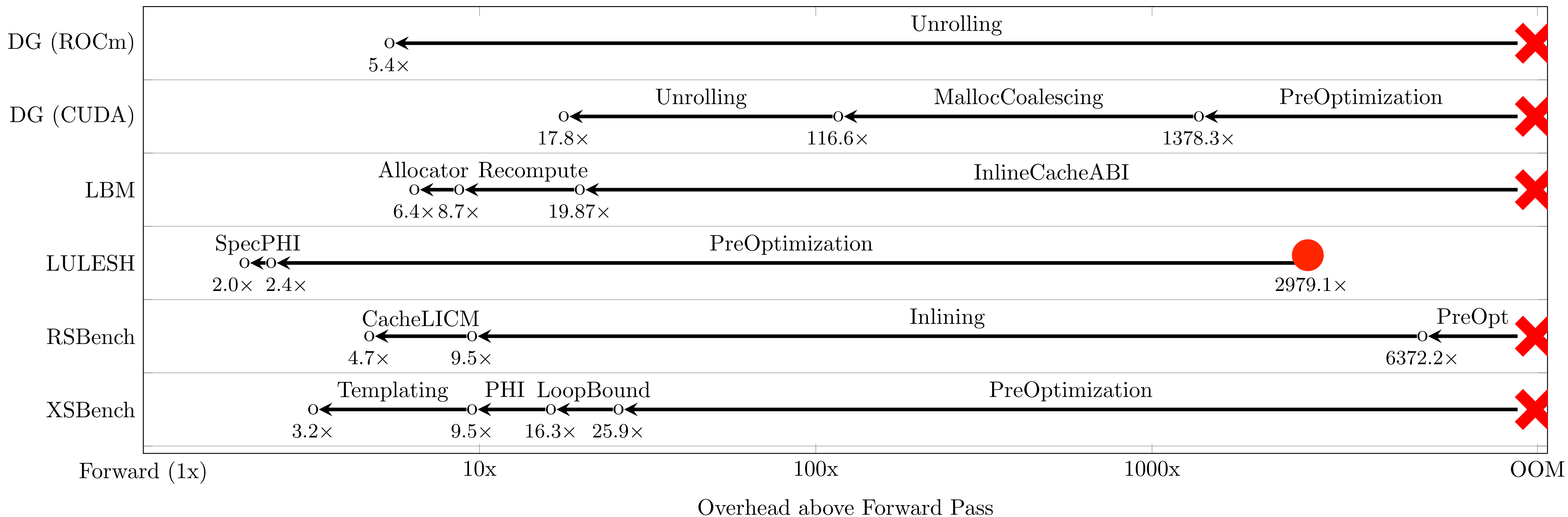


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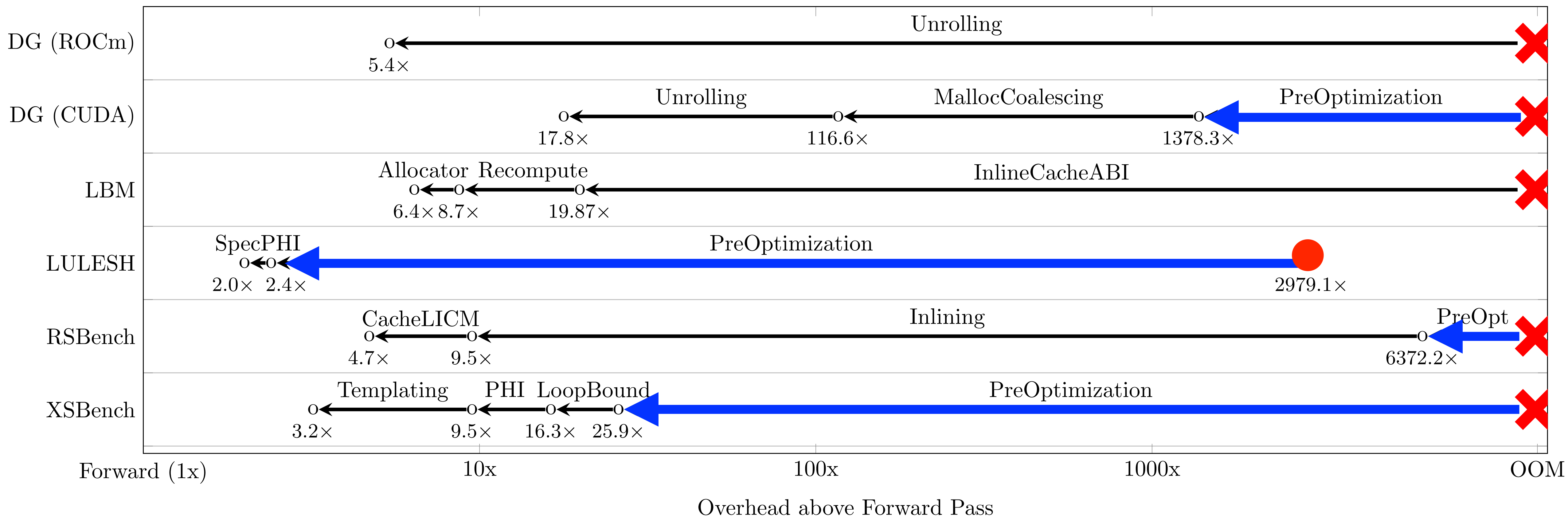
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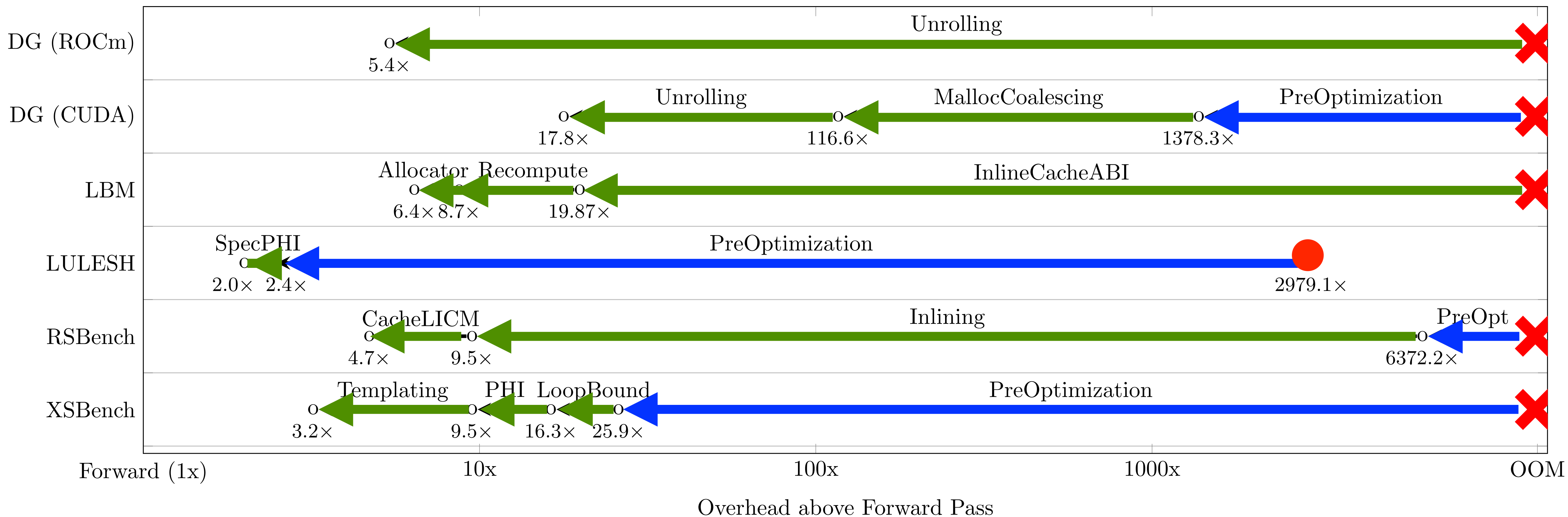
Ablation Analysis of Optimizations [MCPHNMJ'21]



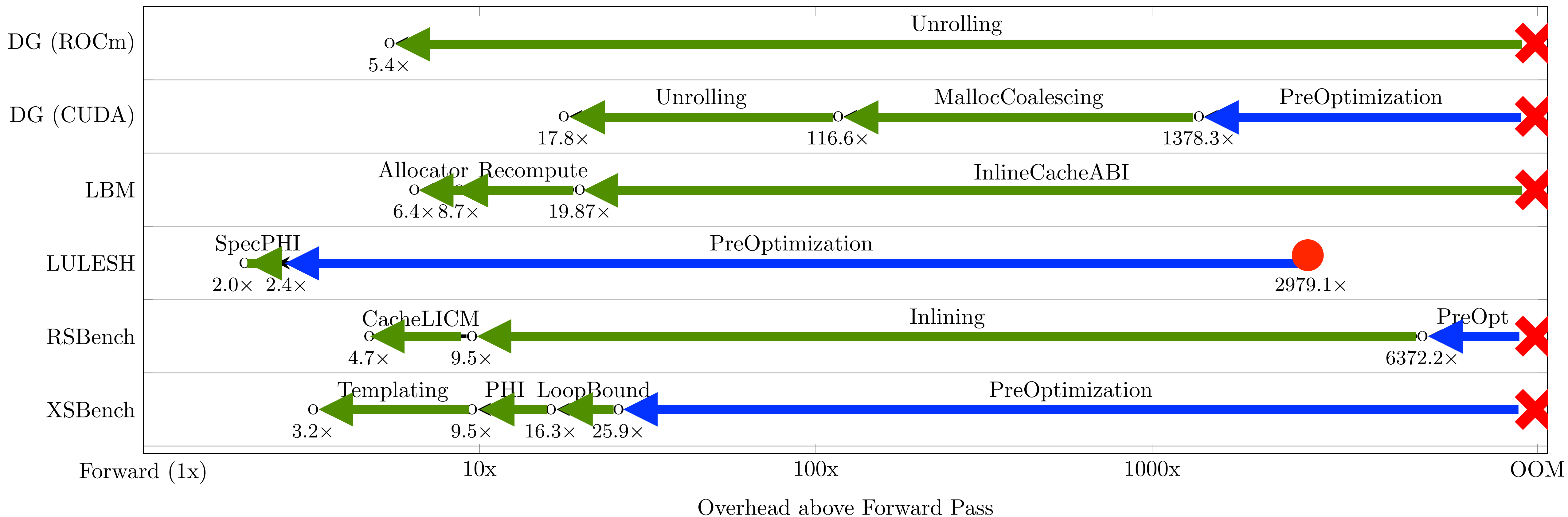
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GPU AD is Intractable Without Optimization!



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PyTorch-Enzyme & TensorFlow-Enzyme

```
import torch
from torch_enzyme import enzyme

# Create some initial tensor
inp = ...

# Apply foreign function to tensor
out = enzyme("test.c", "f").apply(inp)

# Derive gradient
out.backward()
print(inp.grad)
```

```
import tensorflow as tf
from tf_enzyme import enzyme

# Create some initial tensor
inp = tf.Variable(...)

# Use external C code as a regular TF op
out = enzyme(inp, filename="test.c",
              function="f")

# Results is a TF tensor
out = tf.sigmoid(out)
```

```
// Input tensor + size, and output tensor
void f(float* inp, size_t n, float* out);

// diffe_dupnneed specifies not recomputing the output
void diffef(float* inp, float* d_inp, size_t n, float* d_out) {
    __enzyme_autodiff(f, diffe_dup, inp, d_inp, n, diffe_dupnneed, (float*)0, d_out);
}
```



Cache

- Adjoint instructions may require values from the forward pass
 - e.g. $\nabla(x * y) \Rightarrow x \, dy + y \, dx$
- For all values needed in the reverse, allocate memory in the forward pass to store the value
- Values computed inside loops are stored in an array indexed by the loop induction variable
 - Array allocated statically if possible; otherwise dynamically realloc'd



When LLVM Doesn't Cut It

- Enzyme relies on optimizations such as LICM and CSE to eliminate redundant loads, and thus redundant caches.
- Since we instead need to preserve values for the reverse pass, these optimizations may not apply

```
for(int i=0; i<N; i++) {  
    for(int j=0; j<M; j++) {  
        use(array[j]);  
    }  
}  
overwrite(array);
```

When LLVM Doesn't Cut It

- Enzyme relies on optimizations such as LICM and CSE to eliminate redundant loads, and thus redundant caches.
- Since we instead need to preserve values for the reverse pass, these optimizations may not apply
- This requires far more caching than necessary

```
double* cache = new double[N*M];

for(int i=0; i<N; i++) {
    for(int j=0; j<M; j++) {
        cache[i*M+j] = array[j];
        use(array[j]);
    }
}

overwrite(array);
grad_overwrite(array);

for(int i=0; i<N; i++) {
    for(int j=M-1; j<M; j++) {
        grad_use(cache[i*M+j], d_array[j]);
    }
}
```

When LLVM Doesn't Cut It

- Enzyme relies on optimizations such as LICM and CSE to eliminate redundant loads, and thus redundant caches.
- Since we instead need to preserve values for the reverse pass, these optimizations may not apply
- This requires far more caching than necessary
- By analyzing the read/write structure, we can hoist the cache.

```
double* cache = new double[M];
memcpy(cache, array, sizeof(double)*M);
for(int i=0; i<N; i++) {
    for(int j=0; j<M; j++) {

        use(array[j]);
    }
}

overwrite(array);
grad_overwrite(array);

for(int i=0; i<N; i++) {
    for(int j=M-1; i<M; i++) {
        grad_use(cache[j], d_array[j]);
    }
}
```

Cache

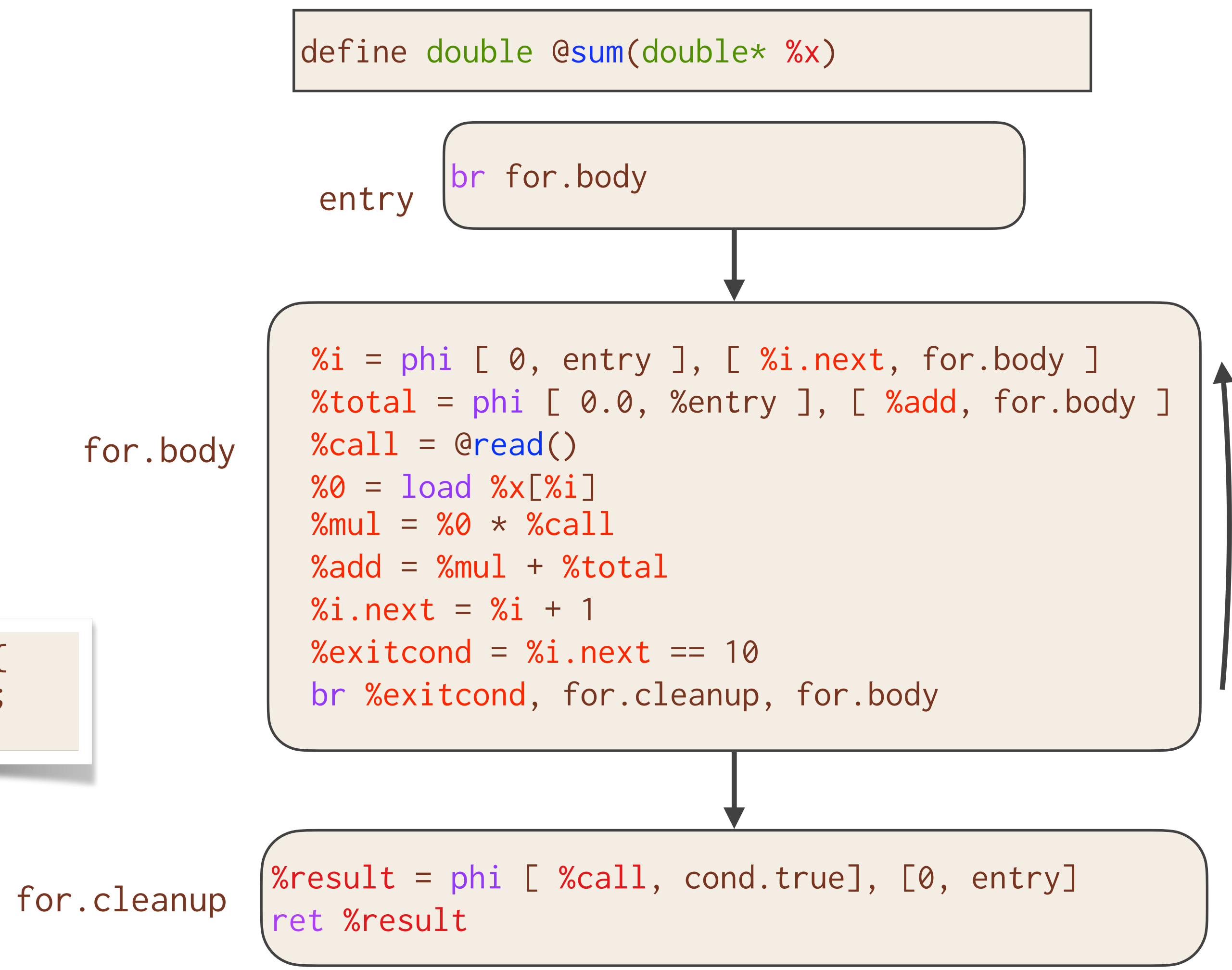
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Case Study: Read Sum

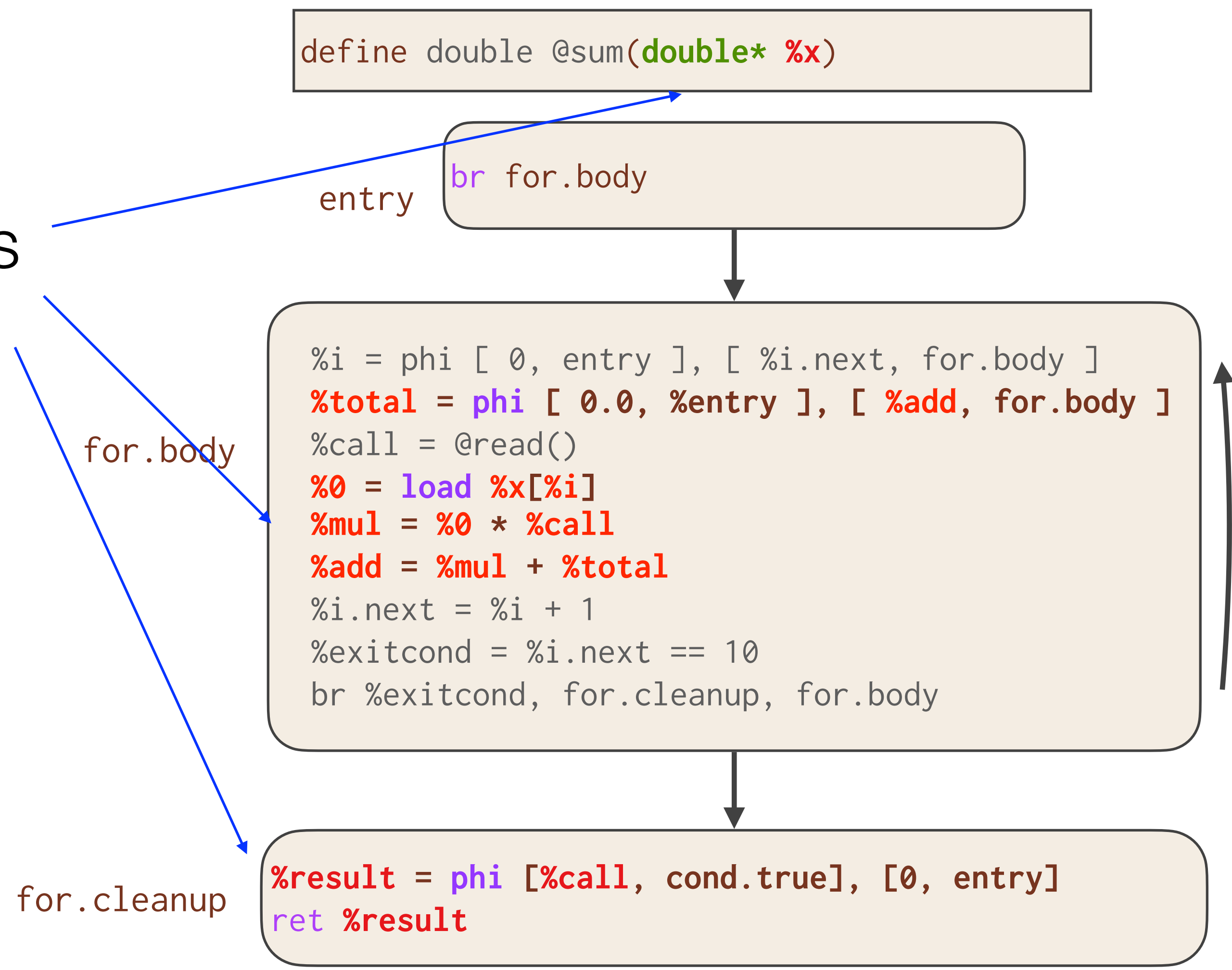
```
double sum(double* x) {  
    double total = 0;  
  
    for(int i=0; i<10; i++)  
        total += read() * x[i];  
  
    return total;  
}
```

```
void diffe_sum(double* x, double* xp) {  
    return __enzyme_autodiff(sum, x, xp);  
}
```



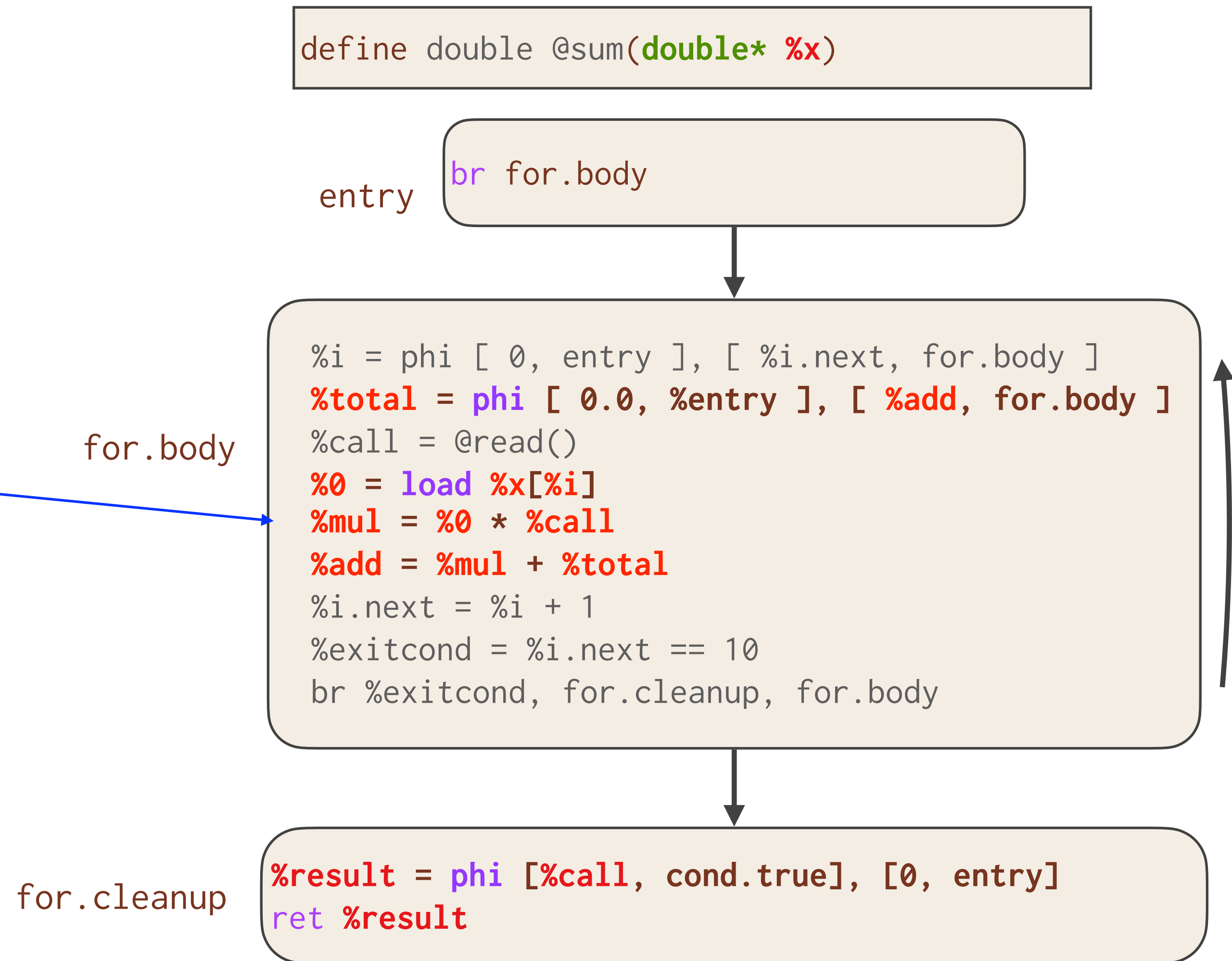
Case Study: Read Sum

Active Variables



Case Study: Read Sum

Each register in the
for loop represents a
distinct active variable
every iteration



Allocate & zero
shadow memory
per active value

```
define double @diffe_sum(double* %x, double* %xp)
```

entry

```
alloca %x'      = 0.0  
alloca %total'   = 0.0  
alloca %0'       = 0.0  
alloca %mul'     = 0.0  
alloca %add'     = 0.0  
alloca %result'  = 0.0  
br for.body
```

for.body

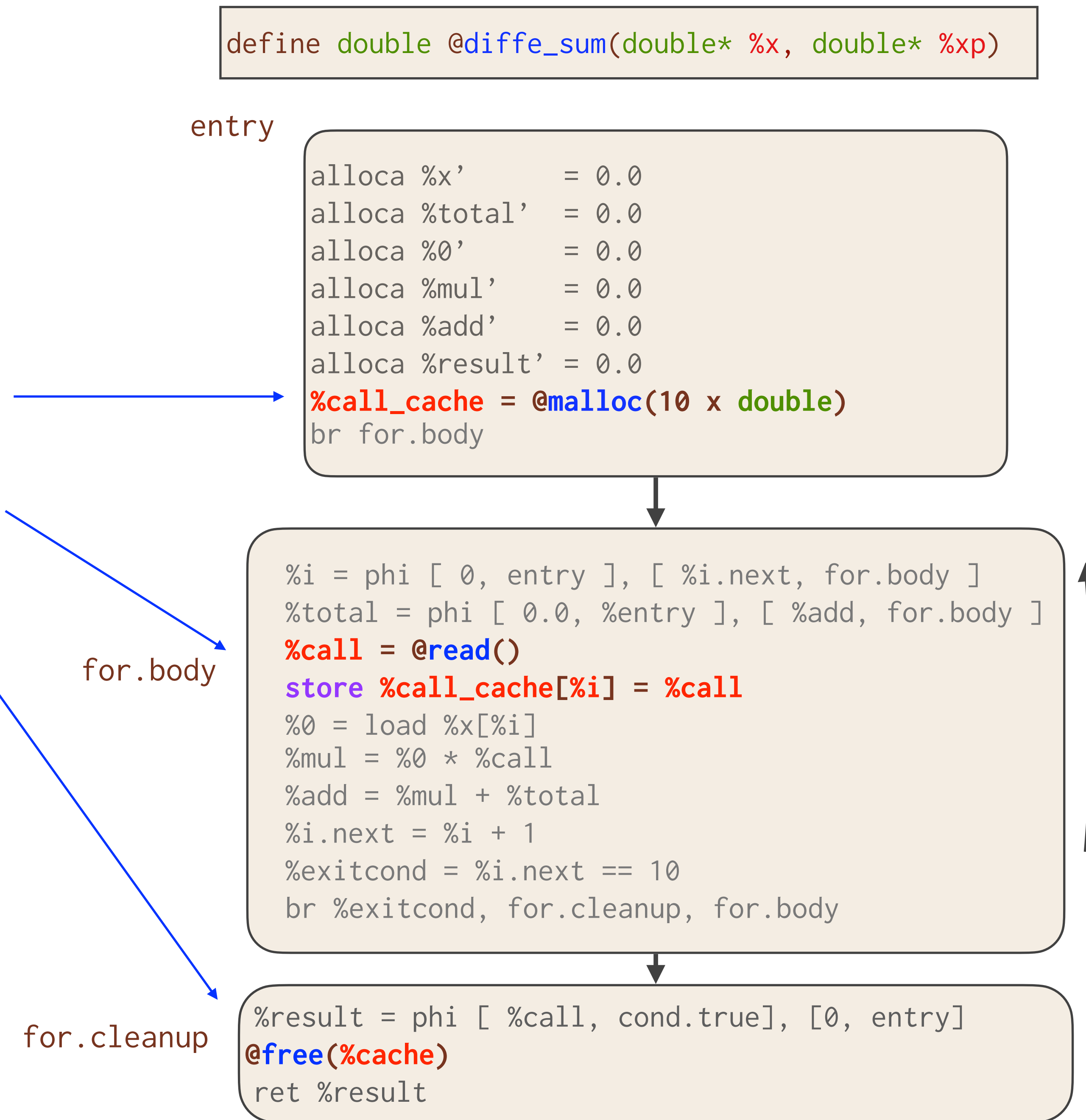
```
%i = phi [ 0, entry ], [ %i.next, for.body ]  
%total = phi [ 0.0, %entry ], [ %add, for.body ]  
%call = @read()  
%0 = load %x[%i]  
%mul = %0 * %call  
%add = %mul + %total  
%i.next = %i + 1  
%exitcond = %i.next == 10  
br %exitcond, for.cleanup, for.body
```

for.cleanup

```
%result = phi [ %call, cond.true ], [ 0, entry ]  
ret %result
```



Cache forward pass
variables for use in
reverse



```
define void @diffe_sum(double* %x, double* %xp)
```

After lowering &
some optimizations

entry

```
%call_cache = @malloc(10 x double)
br for.body
```

for.body

```
%i = phi [ 0, entry ], [ %i.next, for.body ]
%total = phi [ 0.0, %entry ], [ %add, for.body ]
%call = @read()
store %call_cache[%i] = %call
%i.next = %i + 1
%exitcond = %i.next == 10
br %exitcond, reversefor.body, for.body
```

reversefor.body

```
%i' = phi [ 9, for.body ], [ %i'.next, reversefor.body ]
%i'.next = %i' - 1
%cached_read = load %call_cache[%i']
store %xp[%i'] = %cached_read + %xp[%i']
%exit2 = %i = 0
br %exitcond, %exit2, reversefor.body
```

exit

```
@free(%cache)
ret
```



Case Study: Read Sum

```
define void @diffe_sum(double* %x, double* %xp)
```

entry

```
%call0 = @read()
store %xp[0] = %call0
%call1 = @read()
store %xp[1] = %call1
%call2 = @read()
store %xp[2] = %call2
%call3 = @read()
store %xp[3] = %call3
%call4 = @read()
store %xp[4] = %call4
%call5 = @read()
store %xp[5] = %call5
%call6 = @read()
store %xp[6] = %call6
%call7 = @read()
store %xp[7] = %call7
%call8 = @read()
store %xp[8] = %call8
%call9 = @read()
store %xp[9] = %call9
ret
```

After more
optimizations

```
void diffe_sum(double* x, double* xp) {
    xp[0] = read();
    xp[1] = read();
    xp[2] = read();
    xp[3] = read();
    xp[4] = read();
    xp[5] = read();
    xp[6] = read();
    xp[7] = read();
    xp[8] = read();
    xp[9] = read();
}
```



Enzyme on the GPU

- Care must be taken to both ensure correctness and maintain parallelism.
- GPU programs have much lower memory limits. Performance is highly dependent on the number of memory transfers.
- Without first running optimizations reverse-mode AD of large kernels is intractable (OOM).
- Novel GPU and AD-specific optimizations can make a difference of several orders of magnitude when computing gradients.

Test	Overhead
Forward	1
AD, Optimized	4.4
AD, No CacheLICM	343.7
AD, Bad Recompute Heuristic	1275.6
AD, No Inlining	6372.2
AD, No PreOptimization	OOM



CUDA Automatic Differentiation

- Enzyme enables differentiation of CPU programs without rewriting them in a DSL.
- Similarly, GPU programs cannot currently be differentiated without being rewritten in a differentiable language (e.g. PyTorch).
- Enzyme enables reverse-mode AD of general existing GPU programs by:
 - Resolving potential data race issues
 - Differentiating parallel control (syncthreads)
 - Differentiating CUDA intrinsics (e.g. threadIdx.x /llvm.nvvm.read.ptx.sreg.tid.x)
 - Handling shared memory



CUDA Automatic Differentiation

- Most CUDA intrinsics [e.g. threadIdx.x] are inactive and recomputable and thus are incorporated into Enzyme without any special handling
- Derivative of syncthreads is a syncthreads at the corresponding place in reverse pass
- Shared memory is handled by making a second shared memory allocation to act as the shadow for any potentially active uses



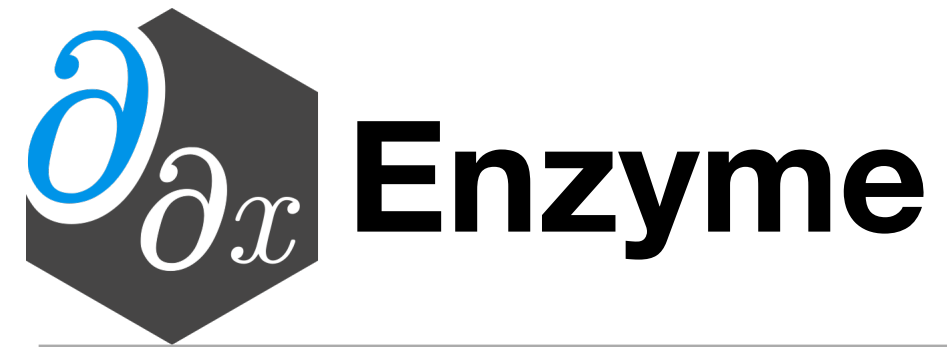


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- Differentiates code in a variety of languages (C, C++, Fortran, Julia, Rust, Swift, etc)
- 4.2x speedup over AD before optimization
- State-of-the art performance with existing tools
- Differentiate GPU kernels
- Open Source (enzyme.mit.edu / github.com/wsmoses/Enzyme)
- PyTorch-Enzyme & TensorFlow-Enzyme imports foreign code in ML workflow

GPU Automatic Differentiation

- Prior work has not explored reverse mode AD of GPU kernels
- Similarly, GPU programs cannot currently be differentiated without being rewritten in a differentiable language (e.g. PyTorch).
- Enzyme enables reverse-mode AD of general existing GPU programs by:
 - Resolving potential data race issues
 - Differentiating parallel control (syncthreads)
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- State-of-the art performance with existing tools
- Differentiate GPU kernels
- Open Source (enzyme.mit.edu / github.com/wsmoses/Enzyme)
- PyTorch-Enzyme & TensorFlow-Enzyme imports foreign code in ML workflow

Custom Derivatives & Multisource

- One can specify custom forward/reverse passes of functions by attaching metadata

```
__attribute__((enzyme("augment", augment_func)))  
__attribute__((enzyme("gradient", gradient_func)))  
double func(double n);
```

- Enzyme leverages LLVM's link-time optimization (LTO) & “fat libraries” to ensure that LLVM bitcode is available for all potential differentiated functions before AD



CUDA Performance Improvements

- Introduce optimizations to reduce the use of memory
 - Alias Analysis to determine legality of recomputing an instruction
 - More aggressive alias analysis properties of syncthreads
- Don't cache unnecessary values
 - Move cache outside of loops when possible
- Heap-to-stack [and to register]
- Don't cache memory itself acting as a cache [such as shared memory]



Enzyme Differentiation Algorithm

- Type Analysis
- Activity Analysis
- Synthesize derivatives
 - Forward pass that mirrors original code
 - Reverse pass inverts instructions in forward pass (adjoints) to compute derivatives
- Optimize



Activity Analysis

- Determines what instructions could impact derivative computation
- Avoids taking meaningless or unnecessary derivatives (e.g. d/dx cpuid)
- Instruction is active iff it can propagate a differential value to its return or memory
- Build off of alias analysis & type analysis
 - E.g. all read-only function that returns an integer are inactive since they cannot propagate adjoints through the return or to any memory location



Compiler Analyses Better Optimize AD

- Existing
- Alias analysis results that prove a function does not write to memory, we can prove that additional function calls do not need to be differentiated since they cannot impact the output
- Don't cache equivalent values
- Statically allocate caches when a loop's bounds can be determined in advance



Decomposing the “Tape”

- Performing AD on a function requires data structures to compute
 - All values necessary to compute adjoints are available [cache]
 - Place to store adjoints [shadow memory]
 - Record instructions [we are static]
- Creating these directly in LLVM allows us to explicitly specify their behavior for optimization, unlike approaches that call out to a library
- For more details look in paper



Conventional Wisdom: AD Only Feasible at High-Level

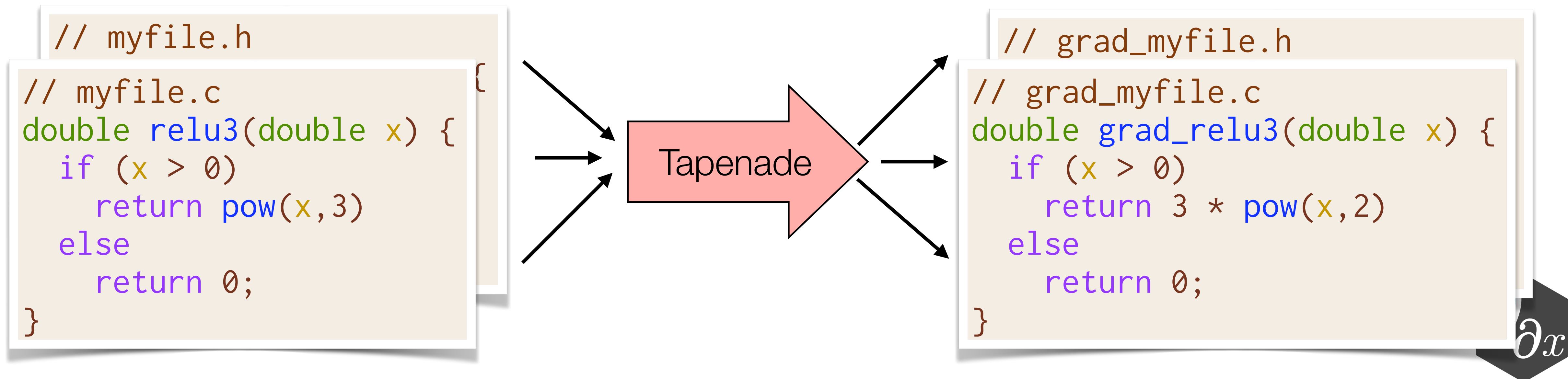
- Automatic Differentiation requires high level semantics to produce gradients
- Lack of high-level information can hinder performance of low-level AD
 - “AD is more effective in high-level compiled languages (e.g. Julia, Swift, Rust, Nim) than traditional ones such as C/C++, Fortran and LLVM IR [...]” -Innes^[1]

[1] Michael Innes. Don't Unroll Adjoint: Differentiating SSA-Form Programs. arXiv preprint arXiv:1810.07951, 2018



Existing AD Approaches (3/3)

- Source rewriting (Zygote.jl -ish, Tapenade)
 - Statically analyze program to produce a new gradient function in the source language
 - Re-implement parsing and semantics of given language
 - Requires all code to be available ahead of time => hard to use with external libraries



Differentiation Is Key To Machine Learning

```
// C++ nbody simulator

void step(std::array<Planet> bodies, double dt) {
    vec3 acc[bodies.size()];
    for (size_t i=0; i<bodies.size(); i++) {
        acc[i] = vec3(0, 0, 0);
        for (size_t j=0; j<bodies.size(); j++) {
            if (i == j) continue;
            acc[i] += force(bodies[i], bodies[j]) /
                       bodies[i].mass;
        }
    }
    for (size_t i=0; i<bodies.size(); i++) {
        bodies[i].vel += acc[i] * dt;
        bodies[i].pos += bodies[i].vel * dt;
    }
}
```

```
// PyTorch rewrite of nbody simulator
import torch

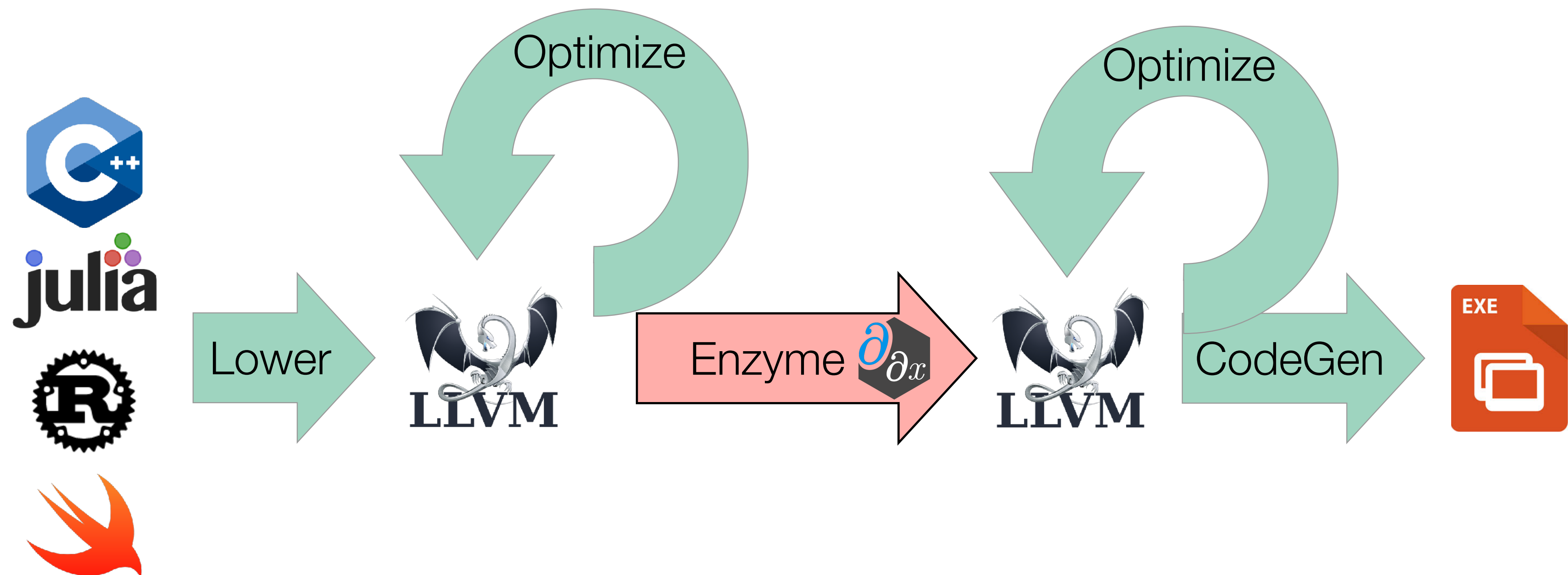
def step(bodies, dt):
    acc = []
    for i in range(len(bodies)):
        acc.push(torch.zeros([3]))
        for j in range(len(bodies)):
            if i == j: continue
            acc[i] += force(bodies[i], bodies[j]) /
                       bodies[i].mass

    for i, body in enumerate(bodies):
        body.vel += acc[i] * dt
        body.pos += body.vel * dt
```

- Hinders application of ML to new domains
- Synthesizing gradients aims to close this gap

Enzyme Overturns Conventional Wisdom

- As fast or faster than state-of-the-art tools
 - Running after optimization enables a **4.2x speedup**
- Necessary semantics for AD derived at low-level (with potential cooperation of frontend)



Parallel Memory Detection

- Thread-local memory
 - Non-atomic load/store
- Same memory location across all threads
 - Parallel Reduction
- Others [always legal fallback]
 - Atomic increment

```
%tmp = load %d_res  
store %d_res = 0  
atomic %d_ptr += %tmp
```

AD-Specific Cache

- Some optimizations require domain-specific knowledge
- Not all values are needed for the reverse pass. By considering the dataflow graph we can perform a min-cut to approximate smaller cache sizes.
- Not all (loop) sizes are known at compile-time, so this must be a heuristic

```
double xy_cache=x[0] + y[0];  
  
use(x[0] + y[0]);  
  
overwrite(x, y);  
grad_overwrite(x, y);  
  
grad_use(xy_cache);
```

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grad_overwrite(x, y);  
  
grad_use(xy_cache);
```

Differentiation Is Key To Machine Learning And Science

- Computing derivatives is key to many algorithms
 - Machine learning (back-propagation, Bayesian inference, uncertainty quantification)
 - Scientific computing (modeling, simulation)
- When working with large codebases or dynamically-generated programs, manually writing derivative functions becomes intractable
- Community has developed tools to create derivatives automatically



Existing AD Approaches

- Differentiable DSL (TensorFlow, PyTorch, DiffTaichi)
 - Provide a new language designed to be differentiated
 - Requires rewriting everything in the DSL and the DSL must support all operations in original code
 - Fast if DSL matches original code well
- Operator overloading (Adept, JAX)
 - Provide differentiable versions of existing language constructs (double => adouble, np.sum => jax.sum)
 - May require writing to use non-standard utilities
 - Often dynamic: storing instructions/values to later be interpreted



Existing AD Approaches

- Source rewriting
 - Statically analyze program to produce a new gradient function in the source language
 - Re-implement parsing and semantics of given language
 - Requires all code to be available ahead of time
 - Difficult to use with external libraries



Case Study: ReLU3

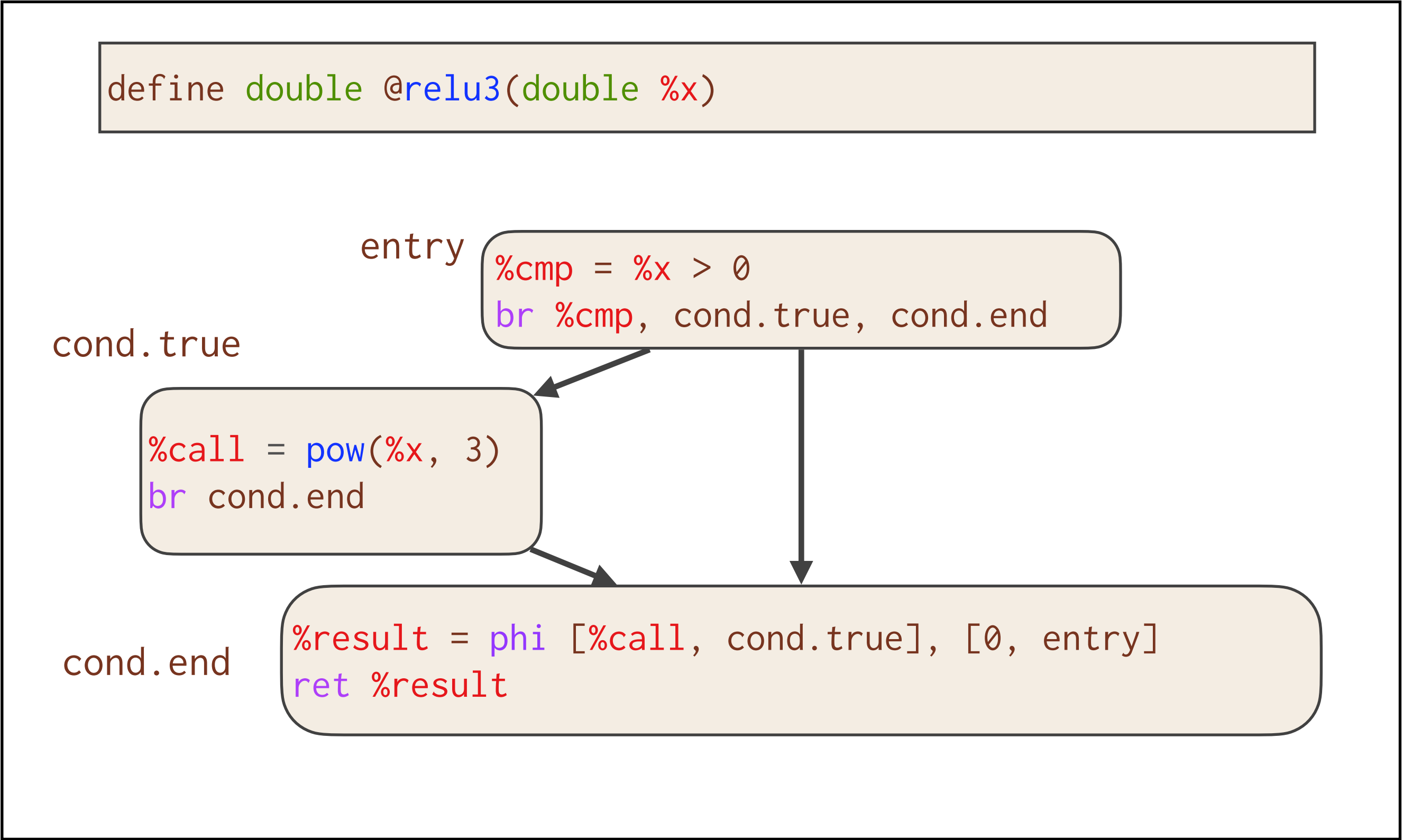
C Source

```
double relu3(double x) {  
    double result;  
    if (x > 0)  
        result = pow(x, 3);  
    else  
        result = 0;  
    return result;  
}
```

Enzyme Usage

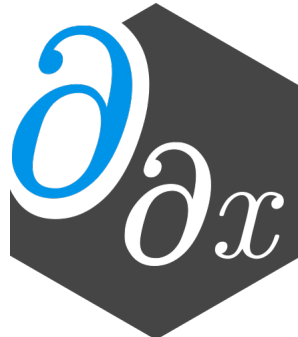
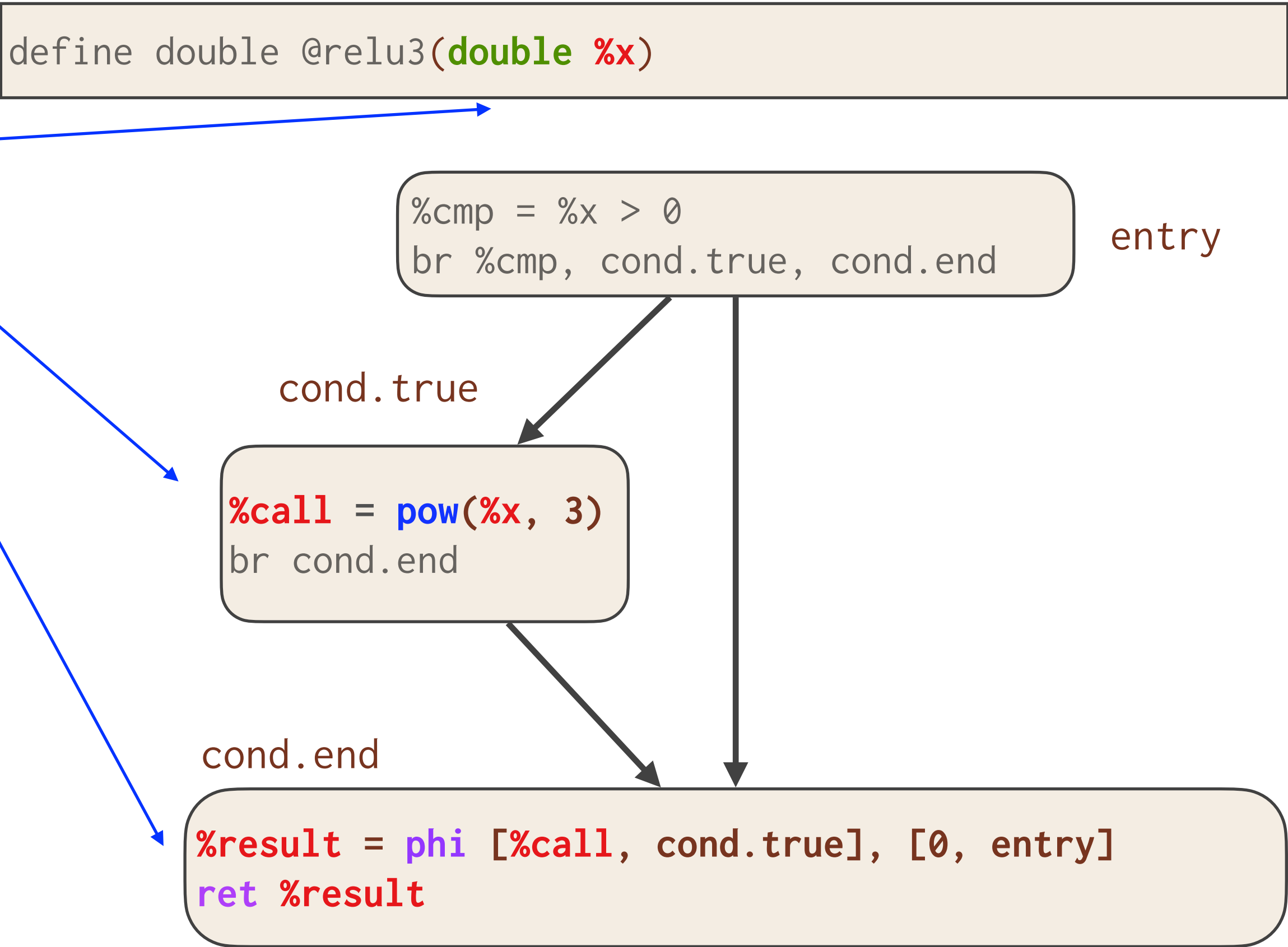
```
double diffe_relu3(double x) {  
    return __enzyme_autodiff(relu3, x);  
}
```

LLVM

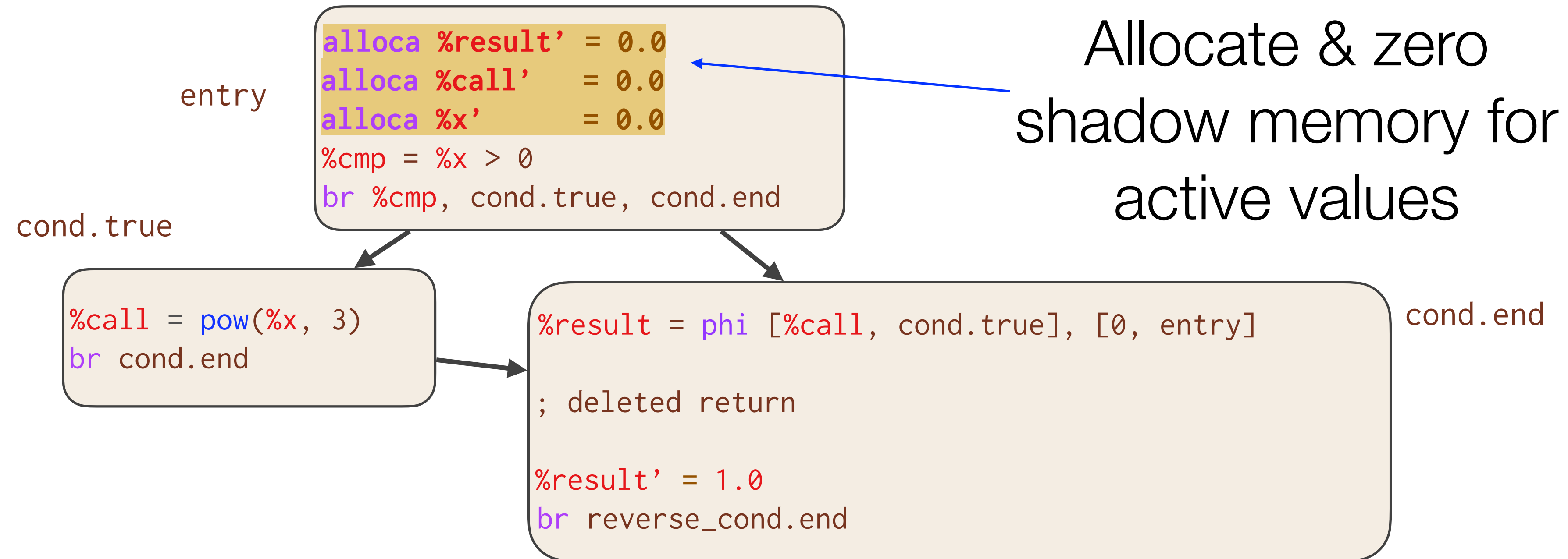


Case Study: ReLU3

Active Instructions

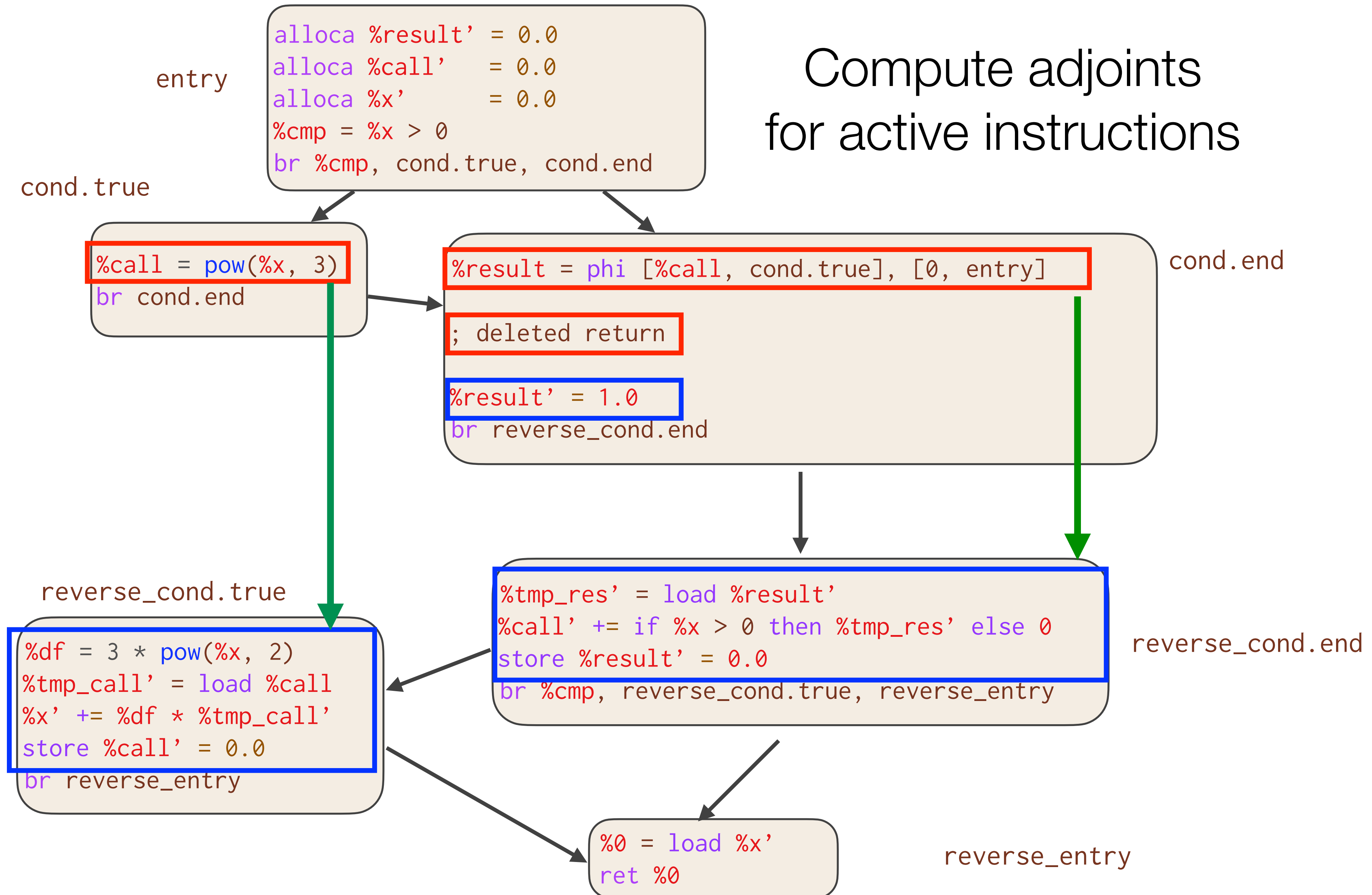


```
define double @diffe_relu3(double %x, double %differet)
```



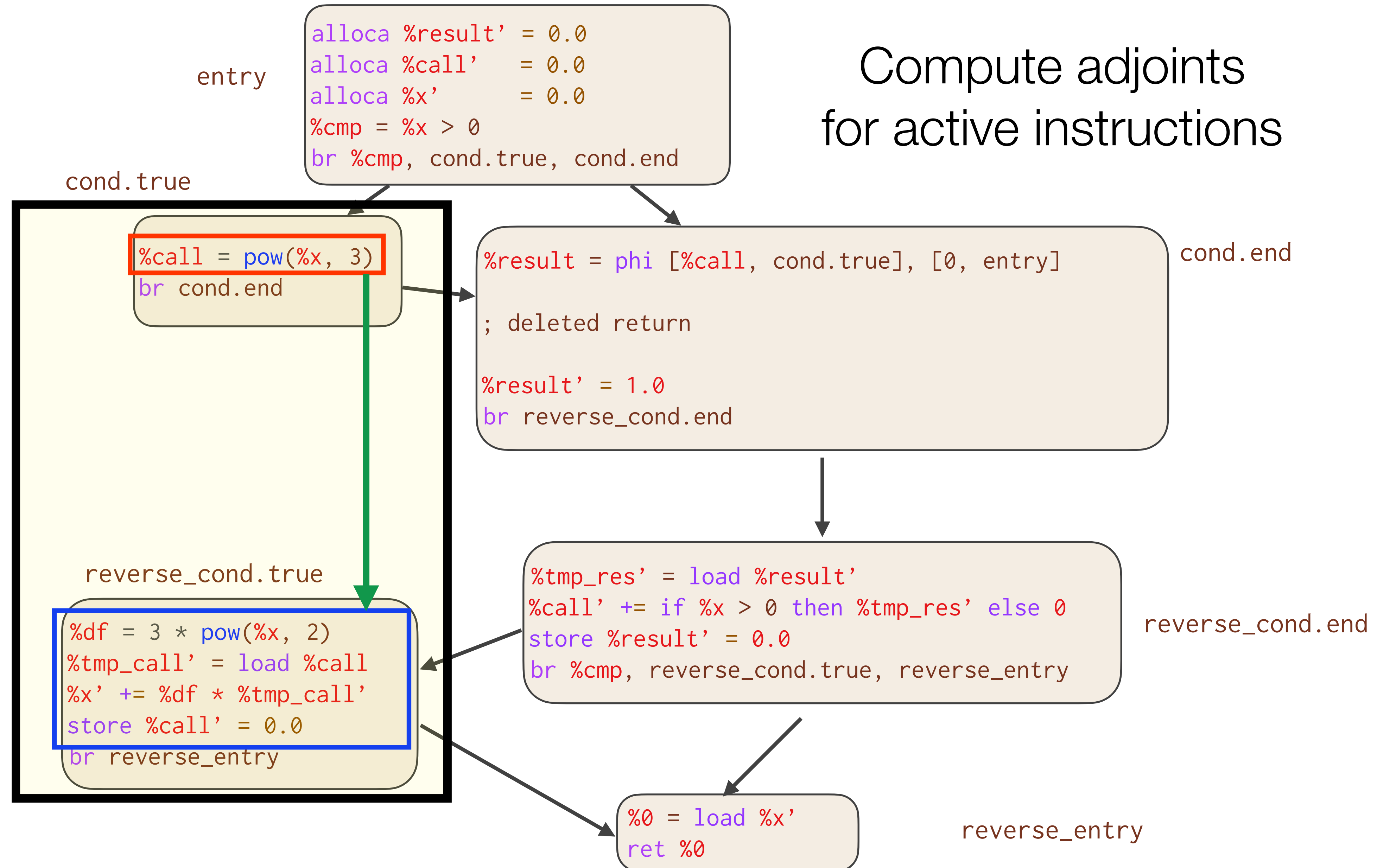
```
define double @diffe_relu3(double %x, double %differet)
```

Compute adjoints
for active instructions



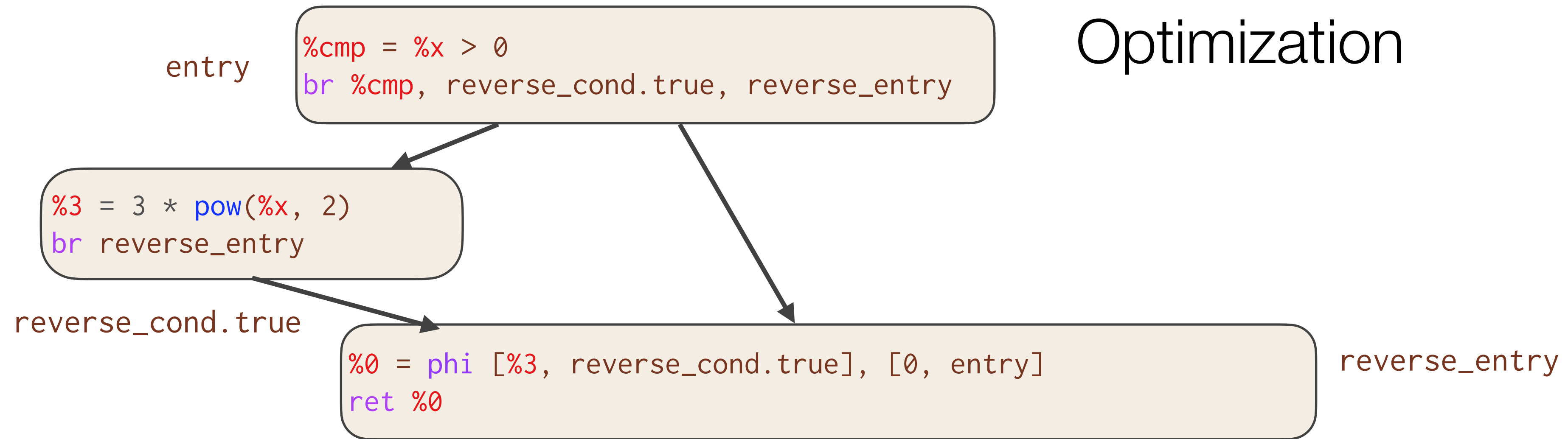
```
define double @diffe_relu3(double %x, double %differet)
```

Compute adjoints for active instructions



```
define double @diffe_relu3(double %x)
```

Post Optimization



Essentially the optimal hand-written gradient!

```
double diffe_relu3(double x) {  
    double result;  
    if (x > 0)  
        result = 3 * pow(x, 2);  
    else  
        result = 0;  
    return result;  
}
```



Challenges of Low-Level AD

- Low-level code lacks information necessary to compute adjoints

```
void f(void* dst, void* src) {  
    memcpy(dst, src, 8);  
}
```

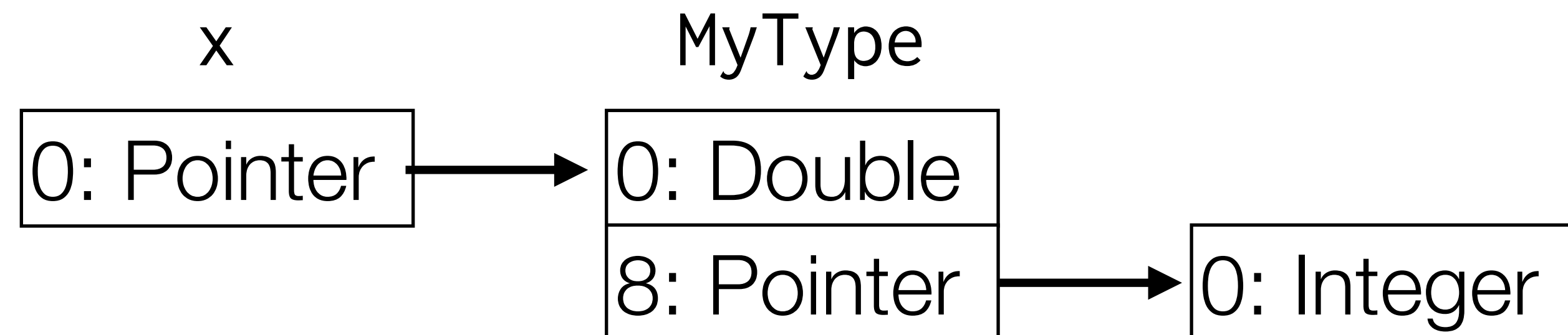
```
void grad_f(double* dst, double* dst',  
            double* src, double* src') {  
    // Forward Pass  
    memcpy(dst, src, 8);  
  
    // Reverse Pass  
    src'[0] += dst'[0];  
    dst'[0] = 0;  
}
```

```
void grad_f(float* dst, float* dst',  
            float* src, float* src') {  
    // Forward Pass  
    memcpy(dst, src, 8);  
  
    // Reverse Pass  
    src'[0] += dst'[0];  
    dst'[0] = 0;  
    src'[1] += dst'[1];  
    dst'[1] = 0;  
}
```

Type Analysis

- New interprocedural dataflow analysis that detects the underlying type of data
- Each value has a set of memory offsets : type
- Perform series of fixed-point updates through instructions

```
struct MyType {  
    double;  
    int*;  
}  
  
x = MyType*;
```



$\text{types}(x) = \{[0]:\text{Pointer}, [0,0]:\text{Double}, [0,8]:\text{Pointer}, [0,8,0]:\text{Integer}\}$

Case 3: Store, Sync, Store

```
codeA(); // store %ptr
sync_threads;

codeB(); // store %ptr
...

diffe_codeB(); // load %d_ptr
                // store %d_ptr = 0

sync_threads;

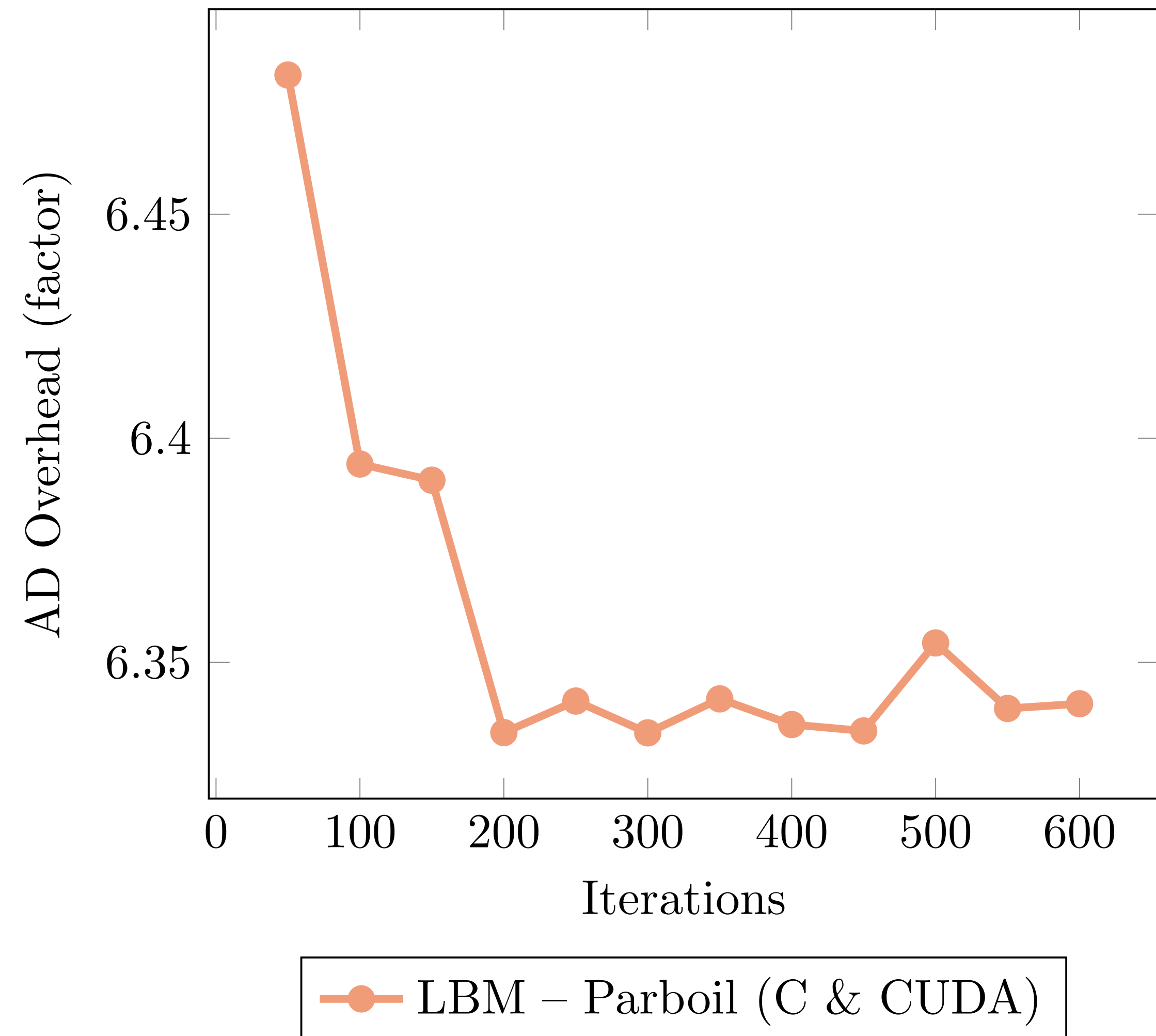
diffe_codeA(); // load %d_ptr
                // store %d_ptr = 0
```



Correct

- All stores to d_ptr in diffe_B will complete prior to diffe_A, ensuring only the clobbering store has its derivative incremented

Scalability Analysis (Fixed Thread Count)



CUDA Example

```
__device__ void inner(float* a, float* x, float* y) {
    y[threadIdx.x] = a[0] * x[threadIdx.x];
}
__device__ void __enzyme_autodiff(void*, ...);

__global__ void daxpy(float* a, float* da, float* x, float* dx, float* y, float* dy) {
    __enzyme_autodiff((void*)inner, a, da, x, dx, y, dy);
}
```

```
__device__ void diffe_inner(float* a, float* da, float* x, float* dx, float* y, float* dy) {
    y[threadIdx.x] = a[0] * x[threadIdx.x];

    float dy = dy[threadIdx.x];
    dy[threadIdx.x] = 0.0f;

    float dx_tmp = a[0] * dy;
    atomic { dx[threadIdx.x] += dx_tmp; }

    float da_tmp = x[threadIdx.x] * dy;
    atomic { da[0] += da_tmp; }
}
```



Existing AD Approaches (1/3)

- Differentiable DSL (TensorFlow, PyTorch, DiffTaichi)
 - Provide a new language designed to be differentiated
 - Requires rewriting everything in the DSL and the DSL must support all operations in original code
 - Fast if DSL matches original code well

```
double square(double val) {  
    return val * val;  
}
```

Manually
Rewrite

```
import tensorflow as tf  
  
x = tf.Variable(3.14)  
  
with tf.GradientTape() as tape:  
    out = tf.math.square(x)  
  
print(tape.gradient(out, x).numpy())
```

Existing AD Approaches (3/3)

- Source rewriting
 - Statically analyze program to produce a new gradient function in the source language
 - Re-implement parsing and semantics of given language
 - Requires all code to be available ahead of time => hard to use with external libraries

```
double square(double val) {  
    return val * val;  
}
```

Tool
Rewrite

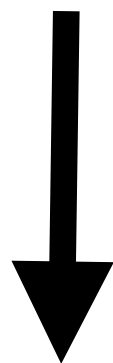
```
double grad_square(double val) {  
    return 2 * val;  
}
```

```
$ tapenade -b -o out.c -head "square(val)/(out)" square.c
```



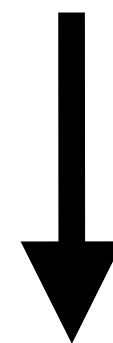
Parallel Automatic Differentiation in LLVM

```
%res = load %ptr
```



```
%tmp = load %d_res  
store %d_res = 0  
atomic %d_ptr += %tmp
```

```
store %ptr = %val
```



```
%tmp = load %d_ptr  
store %d_ptr = 0  
load/store %d_val += %tmp
```

- Shadow Registers `%d_res` and `%d_val` are **thread-local** as they shadow thread-local registers.
- No risk of races and no special handling required.
- Both `%ptr` and shadow `%d_ptr` might be raced upon and require analysis.

Case 2: Load, Sync, Store

```
codeA(); // load %ptr
sync_threads;
codeB(); // store %ptr
...
diffe_codeB(); // load %d_ptr
                // store %d_ptr = 0
sync_threads;
diffe_codeA(); // atomicAdd %d_ptr
```



Correct

- All of the stores of d_ptr will complete prior to any atomicAdds

No cross-thread race here since that's equivalent to a write race in B

Differentiation of SyncThreads

Case 3 [write sync write]

```
codeA(); // store %ptr
sync_threads;
codeB(); // store %ptr
...
diffe_codeB(); // load %d_ptr
                // store %d_ptr = 0
sync_threads;
diffe_codeA(); // load %d_ptr
                // store %d_ptr = 0
```

All uses of stores to d_ptr in diffe_B will correctly complete prior to diffe_A



Case 4 [read sync read]

```
codeA(); // load %ptr
sync_threads;
codeB(); // load %ptr
...
diffe_codeB(); // atomicAdd %d_ptr
sync_threads;
diffe_codeA(); // atomicAdd %d_ptr
```

Original and differential sync unnecessary and legal to include



Scalability Analysis (Fixed Work Per Thread)

