A Step-by-step Introduction to the Implementation of Automatic Differentiation

Yu-Hsueh Fang*¹, He-Zhe Lin*¹, Jie-Jyun Liu¹, and Chih-Jen Lin^{1,2}

¹National Taiwan University
{d11725001, r11922027, d11922012}@ntu.edu.tw
cjlin@csie.ntu.edu.tw

²Mohamed bin Zayed University of Artificial Intelligence
chihjen.lin@mbzuai.ac.ae

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Abstract

Automatic differentiation is a key component in deep learning. This topic is well studied and excellent surveys such as Baydin et al. (2018) have been available to clearly describe the basic concepts. Further, sophisticated implementations of automatic differentiation are now an important part of popular deep learning frameworks. However, it is difficult, if not impossible, to directly teach students the implementation of existing systems due to the complexity. On the other hand, if the teaching stops at the basic concept, students fail to sense the realization of an implementation. For example, we often mention the computational graph in teaching automatic differentiation, but students wonder how to implement and use it. In this document, we partially fill the gap by giving a step by step introduction of implementating a simple automatic differentiation system. We streamline the mathematical concepts and the implementation. Further, we give the motivation behind each implementation detail, so the whole setting becomes very natural.

1 Introduction

In modern machine learning, derivatives are the cornerstone of numerous applications and studies. The calculation often relies on automatic differentiation, a classic method for efficiently and accurately calculating derivatives of numeric functions. For example, deep learning cannot succeed without automatic differentiation. Therefore, teaching students how automatic differentiation works is highly essential.

Automatic differentiation is a well-developed area with rich literature. Excellent surveys including Chinchalkar (1994), Bartholomew-Biggs et al. (2000), Baydin et al. (2018) and Margossian

^{*}These authors contributed equally to this work

(2019) review the algorithms for automatic differentiation and its wide applications. In particular, Baydin et al. (2018) is a comprehensive work focusing the automatic differentiation in machine learning. Therefore, there is no lack of materials introducing the concept of automatic differentiation.

On the other hand, as deep learning systems now solve large-scale problems, it is inevitable that the implementation of automatic differentiation becomes highly sophisticated. For example, in popular deep learning systems such as PyTorch (Paszke et al., 2017) and Tensorflow (Abadi et al., 2016), at least thousands of lines of code are needed. Because of this, many places of teaching automatic differentiation for deep learning stops at the basic concepts. Then students fail to sense the realization of an implementation. For example, we often mention the computational graph in teaching automatic differentiation, but students wonder how to implement and use it. In this document, we aim to partially fill the gap by giving a tutorial on the basic implementation.

In recent years, many works^{1,2,3,4,5} have attempted to discuss the basic implementation of automatic differentiation. However, they still leave room for improvement. For example, some are not self-contained – they quickly talk about implementations without connecting to basic concepts. Ours, which is very suitable for the beginners, has the following features:

- We streamline the mathematical concepts and the implementation. Further, we give the motivation behind each implementation detail, so the whole setting becomes very natural.
- We use the example from Baydin et al. (2018) for the consistency with past works. Ours is thus an extension of Baydin et al. (2018) into the implementation details.
- We build a complete tutorial including this document, slides and the source code at https://www.csie.ntu.edu.tw/~cjlin/papers/autodiff/.

2 Automatic Differentiation

There are two major modes of automatic differentiation. In this section, we introduce the basic concepts of both modes. Most materials in this section are from Baydin et al. (2018). We consider the same example function

$$y = f(x_1, x_2) = \log x_1 + x_1 x_2 - \sin x_2.$$

 $[\]overline{}^1 {
m https://towardsdatascience.com/build-your-own-automatic-differentiation-program-6ecd585eec2a}$

²https://sidsite.com/posts/autodiff

https://mdrk.io/introduction-to-automatic-differentiation-part2/

⁴https://github.com/dlsyscourse/lecture5/blob/main/5_automatic_differentiation_

implementation.ipynb

⁵https://github.com/karpathy/micrograd

2.1 Forward Mode

First, we discuss the forward mode. Before calculating the derivative, let us check how to calculate the function value. Assume that we want to calculate the function value at $(x_1, x_2) = (2, 5)$. Then, in the following table, we have a forward procedure.

$$x_{1} = 2$$

$$x_{2} = 5$$

$$v_{1} = \log x_{1} = \log 2$$

$$v_{2} = x_{1} \times x_{2} = 2 \times 5$$

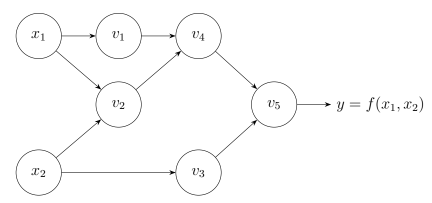
$$v_{3} = \sin x_{2} = \sin 5$$

$$v_{4} = v_{1} + v_{2} = 0.693 + 10$$

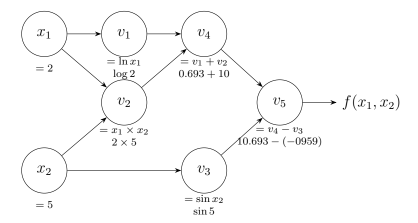
$$v_{5} = v_{4} - v_{3} = 10.693 + 0.959$$

$$y = v_{5} = 11.652$$

We use variables v_i to record the intermediate outcomes. First, we know log function is applied to x_1 . Therefore, we have $\log(x_1)$ as a new variable called v_1 . Similarly, there is a variable v_2 , which is $x_1 \times x_2$. Each v_i is related to a simple operation. The initial value of this member is zero when a node is created. In the end, our function value at (2,5) is $y=v_5$. As shown in the table, the function evaluation is decomposed into a sequence of simple operations. We have a corresponding computational graph as follows:



Because calculating both v_1 and v_2 needs x_1 , x_1 has two links to them in the graph. The following graph shows all the intermediate results in the computation.



The computational graph tells us the dependencies of variables. Thus, from the inputs x_1 and x_2 we can go through all nodes for getting the function value $y = v_5$ in the end.

Now, we have learned about the function evaluation. But remember, we would like to calculate the derivative. Assume that we target at the partial derivative $\partial y/\partial x_1$. Here, we denote

$$\dot{v} = \frac{\partial v}{\partial x_1}$$

as the derivative of the variable v with respect to x_1 . The idea is that by using the chain rule, we can obtain the following forward derivative calculation to eventually get $\dot{v}_5 = \partial f/\partial x_1$.

The table starts from \dot{x}_1 and \dot{x}_2 , which are $\partial x_1/\partial x_1 = 1$ and $\partial x_2/\partial x_1 = 0$. Based on \dot{x}_1 and \dot{x}_2 , we can calculate other values. For example, let us check the partial derivative $\partial v_1/\partial x_1$. From

$$v_1 = \log x_1$$

by the chain rule,

$$\dot{v}_1 = \frac{\partial v_1}{\partial x_1} \times \frac{\partial x_1}{\partial x_1} = \frac{1}{x_1} \times \frac{\partial x_1}{\partial x_1} = \frac{\dot{x}_1}{x_1}.$$

Therefore, we need \dot{x}_1 and x_1 for calculating \dot{v}_1 on the left-hand side. We already have the value of \dot{x}_1 from the previous step $(\partial x_1/\partial x_1 = 1)$. Also, the function evaluation gives the value of x_1 . Then, we can calculate $\dot{x}_1/x_1 = 1/2$. Clearly, the chain rule plays an important role here. The calculation of other \dot{v}_i is similar.

2.2 Reverse Mode

Next, we discuss the reverse mode. We denote

$$\bar{v} = \frac{\partial y}{\partial v}$$

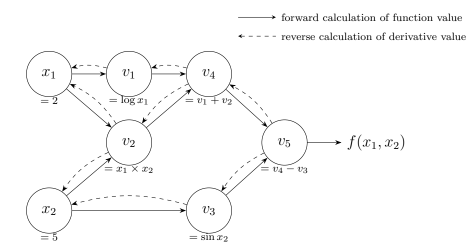
as the derivative of the function y with respect to the variable v. Note that earlier, in the forward mode, we considered

$$\dot{v} = \frac{\partial v}{\partial x_1},$$

so the focus is on the derivatives of all variables with respect to one input variable. In contrast, the reverse mode focuses on $\bar{v} = \partial y/\partial v$ for all v, the partial derivatives of one output with respect to all variables. Therefore, for our example, we can use \bar{v}_i 's and \bar{x}_i 's to get both $\partial y/\partial x_1$ and $\partial y/\partial x_2$ at once. Now we illustrate the calculation of

$$\frac{\partial y}{\partial x_2}$$
.

By checking the variable x_2 in the computational graph, we see that variable x_2 affects y by affecting v_2 and v_3 .



This dependency, together with the fact that x_1 is fixed, means if we would like to calculate $\partial y/\partial x_2$, then it is equal to calculate

$$\frac{\partial y}{\partial x_2} = \frac{\partial y}{\partial v_2} \frac{\partial v_2}{\partial x_2} + \frac{\partial y}{\partial v_3} \frac{\partial v_3}{\partial x_2}.$$
 (1)

We can rewrite Equation 1 as follows with our notataion.

$$\bar{x}_2 = \bar{v}_2 \frac{\partial v_2}{\partial x_2} + \bar{v}_3 \frac{\partial v_3}{\partial x_2}.$$
 (2)

If \bar{v}_2 and \bar{v}_3 are available beforehand, all we need is to calculate $\partial v_2/\partial x_2$ and $\partial v_3/\partial x_2$. From the operation between x_2 and v_3 , we know that $\partial v_3/\partial x_2 = \cos(x_2)$. Similarly, we have $\partial v_2/\partial x_2 = x_1$. Then, the evaluation of \bar{x}_2 is done in two steps:

$$\bar{x}_2 \leftarrow \bar{v}_3 \frac{\partial v_3}{\partial x_2}$$
$$\bar{x}_2 \leftarrow \bar{x}_2 + \bar{v}_2 \frac{\partial v_2}{\partial x_2}.$$

These steps are part of the sequence of a reverse traversal, shown in the following table.

To get the desired \bar{x}_1 and \bar{x}_2 (i.e., $\partial y/\partial x_1$ and $\partial y/\partial x_2$), we begin with

$$\bar{v}_5 = \frac{\partial y}{\partial v_5} = \frac{\partial y}{\partial y} = 1.$$

From the computational graph, we then get \bar{v}_4 and \bar{v}_3 . Because v_4 affects y only through v_5 , we have

$$\bar{v}_4 = \frac{\partial y}{\partial v_4} = \frac{\partial y}{\partial v_5} \frac{\partial v_5}{\partial v_4} = \bar{v}_5 \frac{\partial v_5}{\partial v_4} = \bar{v}_5 \times 1$$

The above equation is based on that we already know $\partial y/\partial v_5 = \bar{v}_5$. Also, the operation from v_4 to v_5 is an addition, so $\partial v_5/\partial v_4$ is a constant 1. By such a sequence, in the end, we obtain

$$\frac{\partial y}{\partial x_1} = \bar{x}_1 \text{ and } \frac{\partial y}{\partial x_2} = \bar{x}_2$$

at the same time.

3 Implementation of Function Evaluation and the Computational Graph

With the basic concepts ready in Section 2, we move to the implementation of the automatic differentiation. For simplicity, we consider the forward mode. The reverse mode can be designed in a similar way. Consider a function $f: \mathbb{R}^n \to \mathbb{R}$ with

$$y = f(\boldsymbol{x}) = f(x_1, x_2, \dots, x_n).$$

For any given \boldsymbol{x} , we show the computation of

$$\frac{\partial y}{\partial x_1}$$

as an example.

3.1 The Need to Calculate Function Values

We are calculating the derivative, so at the first glance, function values are not needed. However, we show that function evaluation is necessary due to the function structure and the use of the chain rule. To explain this, we begin with knowing that the function of a neural network is usually a nested composite function

$$f(\mathbf{x}) = h_k(h_{k-1}(\dots h_1(\mathbf{x})))$$

due to the layered structure. For easy discussion, let us assume that f(x) is the following general composite function

$$f(\boldsymbol{x}) = g(h_1(\boldsymbol{x}), h_2(\boldsymbol{x}), \dots, h_k(\boldsymbol{x})).$$

We can see that the example function considered earlier

$$f(x_1, x_2) = \log x_1 + x_1 x_2 - \sin x_2 \tag{3}$$

can be written in the following composite function

$$g(h_1(x_1, x_2), h_2(x_1, x_2))$$

with

$$g(h_1, h_2) = h_1 - h_2,$$

$$h_1(x_1, x_2) = \log x_1 + x_1 x_2,$$

$$h_2(x_1, x_2) = \sin(x_2).$$

To calculate the derivative at $x = x_0$ using the chain rule, we have

$$\frac{\partial f}{\partial x_1}\Big|_{\boldsymbol{x}=\boldsymbol{x}_0} = \sum_{i=1}^k \left(\frac{\partial g}{\partial h_i} \Big|_{\boldsymbol{h}=\boldsymbol{h}(\boldsymbol{x}_0)} \times \frac{\partial h_i}{\partial x_1} \Big|_{\boldsymbol{x}=\boldsymbol{x}_0} \right),$$

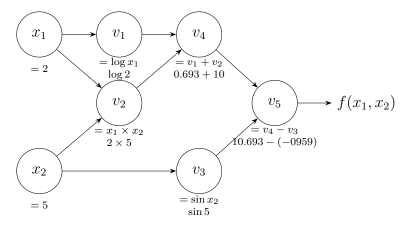
where the notation

$$\left. \frac{\partial g}{\partial h_i} \right|_{\boldsymbol{h} = \boldsymbol{h}(\boldsymbol{x}_0)}$$

means the derivative of g with respect to h_i evaluated at $\mathbf{h}(\mathbf{x}_0) = \begin{bmatrix} h_1(\mathbf{x}_0) & \cdots & h_k(\mathbf{x}_0) \end{bmatrix}^T$. Clearly, we must calculate the inner function values $h_1(\mathbf{x}_0), \dots, h_k(\mathbf{x}_0)$ first. The process of computing all $h_i(\mathbf{x}_0)$ is part of (or almost the same as) the process of computing $f(\mathbf{x}_0)$. This explanation tells why for calculating the partial derivatives, we need the function value first.

Next, we discuss the implementation of getting the function value. For the function (3), recall we have a table recording the order to get $f(x_1, x_2)$:

Also, we have a computational graph to generate the computing order



Therefore, we must check how to build the graph.

3.2 Creating the Computational Graph

A graph consists of nodes and edges. We must discuss what a node/edge is and how to store information. From the graph shown above, we see that each node represents an intermediate expression:

$$v_1 = \log x_1,$$

 $v_2 = x_1 \times x_2,$
 $v_3 = \sin x_2,$
 $v_4 = v_1 \times v_2,$
 $v_5 = v_4 - v_3.$

The expression in each node is from an operation to expressions in other nodes. Therefore, it is natural to construct an edge

$$u \to v$$
,

if the expression of a node v is based on the expression of another node u. We say node u is a parent node (of v) and node v is a child node (of u). To do the forward calculation, at node v we should

store v's parents. Additionally, we need to record the operator applied on the node's parents and the resulting value. For example, the construction of the node

$$v_2 = x_1 \times x_2,$$

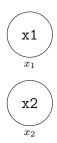
requires to store v_2 's parent nodes $\{x_1, x_2\}$, the corresponding operator " \times " and the resulting value. Up to now, we can implement each node as a class Node with the following members.

member	data type	example for Node v_2
numerical value	float	10
parent nodes	List[Node]	$[x_1, x_2]$
child nodes	List[Node]	$[v_4]$
operator	string	"mul" $(for \times)$

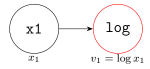
At this moment, it is unclear why we should store child nodes in our Node class. Later we will explain why such information is needed. Once the Node class is ready, starting from initial nodes (which represent x_i 's), we use nested function calls to build the whole graph. In our case, the graph for $y = f(x_1, x_2)$ can be constructed via

$$y = sub(add(log(x1), mul(x1, x2)), sin(x2)).$$

let us see this process step by step and check what each function must do. First, our starting point is the root nodes created by the Node class constructor.



These root Nodes have empty members "parent nodes," "child nodes," and "operator" with only "numerical value" respectively set to x_1 and x_2 . Then, we apply our implemented log(node) to the node x_1 .



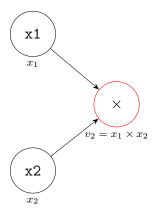
The implementation of our log function should create a Node instance to store $log(x_1)$. Therefore, what we have is a wrapping function that does more than the log operation; see details in Section 3.3. The created node is the v_1 node in our computational graph. Next, we discuss details of the node creation. From the current log function and the input node x_1 , we know contents of the following members:

parent nodes: [x₁]operator: "log"

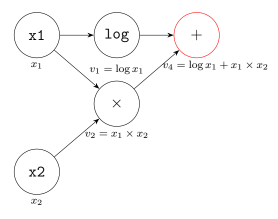
• numerical value: log 2

However, we have no information about children of this node. The reason is obvious because we have not had a graph including its child nodes yet. Instead, we leave this member "child nodes" empty and let child nodes to write back the information. By this idea, our \log function should add v_1 to the "child nodes" of x_1 . See more details later in Section 3.3.

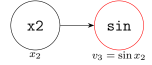
We move on to apply mul(node1, node2) on nodes x1 and x2.



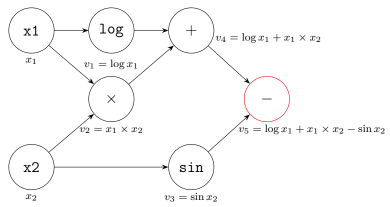
Similarly, the mul function generates a Node instance. However, different from log(x1), the node created here stores two parents (instead of one). Then we apply the function call add(log(x1), mul(x1, x2)).



Next, we apply sin(node) to x2.



Last, applying sub(node1,node2) to the output nodes of add(log(x1), mul(x1, x2)) and sin(x1) leads to



We can conclude that each function generates exactly one Node instance; however, the generated nodes differ in the operator, the number of parents, etc.

3.3 Wrapping Functions

We mentioned that a function like "mul" does more than calculating the product of two numbers. Here we show more details. These customized functions "add", "mul" and "log" in the previous pages are wrapping functions, which "wrap" numerical operations with additional codes. An important task is to maintain the relation between the constructed node and its parents/children. This way, the information of graph can be preserved.

For example, we may implement the following "mul" function.

Listing 1: The wrapping function "mul"

```
def mul(node1, node2):
    value = node1.value * node2.value
    parent_nodes = [node1, node2]
    newNode = Node(value, parent_nodes, "mul")
    node1.child_nodes.append(newNode)
    node2.child_nodes.append(newNode)
    return newNode
```

In this code, we add the created node to the "child nodes" lists of the two input nodes: node1 and node2. As we mentioned earlier, when node1 and node2 were created, their lists of child nodes were unknown and left empty. Thus, in creating each node, we append the node to the list of its parent(s).

The output of the function should be the created node. This setting enables the nested function call. That is, calling

```
y = sub(add(log(x1), mul(x1, x2)), sin(x2))
```

finishes the function evaluation. At the same time, we build the computational graph.

4 Topological Order and Partial Derivatives

Once the computational graph is built, we want to use the information in the graph to compute

$$\frac{\partial y}{\partial x_1} = \frac{\partial v_5}{\partial x_1}.$$

4.1 Finding the Topological Order

Recall that $\partial v/\partial x_1$ is denoted by \dot{v} . From the chain rule,

$$\dot{v}_5 = \frac{\partial v_5}{\partial v_4} \dot{v}_4 + \frac{\partial v_5}{\partial v_3} \dot{v}_3. \tag{4}$$

We are able to calculate

$$\frac{\partial v_5}{\partial v_4}$$
 and $\frac{\partial v_5}{\partial v_3}$, (5)

because the node v_5 stores the needed information related to its parents v_4 and v_3 . We defer the details on calculating (5), so the focus is now on calculating \dot{v}_4 and \dot{v}_3 . For \dot{v}_4 , we further have

$$\dot{v}_4 = \frac{\partial v_4}{\partial v_1} \dot{v}_1 + \frac{\partial v_4}{\partial v_2} \dot{v}_2,\tag{6}$$

which, by the same reason, indicates the need of \dot{v}_1 and \dot{v}_2 . On the other hand, we have $\dot{v}_3 = 0$ since v_3 (i.e., $\sin(x_2)$) is not a function of x_1 . The discussion on calculating \dot{v}_4 and \dot{v}_3 leads us to find that

$$v$$
 is not reachable from x_1 in the graph $\Rightarrow \dot{v} = 0$. (7)

We say a node v is reachable from a node u if there exists a path from u to v in the graph. From (7), now we only care about nodes reachable from x_1 . Further, we must properly order nodes reachable from x_1 so that, for example, in (4.2), \dot{v}_4 and \dot{v}_3 are ready before calculating \dot{v}_5 . Similarly, \dot{v}_1 and \dot{v}_2 should be available when calculating \dot{v}_4 .

To consider nodes reachable from x_1 , from the whole computational graph $G = \langle V, E \rangle$, where V and E are respectively sets of nodes and edges, we define

$$V_R = \{ v \in V \mid v \text{ is reachable from } x_1 \}$$

and

$$E_R = \{(u, v) \in E \mid u \in V_R, \ v \in V_R\}.$$

Then,

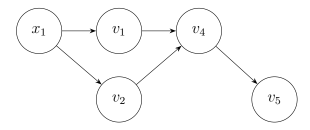
$$G_R \equiv \langle V_R, E_R \rangle$$

is a subgraph of G. For our example, G_R is the following subgraph with

$$V_R = \{x_1, v_1, v_2, v_4, v_5\}$$

and

$$E_R = \{(x_1, v_1), (x_2, v_2), (v_1, v_4), (v_2, v_4), (v_4, v_5)\}.$$



We aim to find a "suitable" ordering of V_R satisfying that each node $u \in V_R$ comes before all of its child nodes in the ordering. By doing so, \dot{u} can be used in the derivative calculation of its child nodes; see (6). For our example, a "suitable" ordering can be

$$x_1, v_1, v_2, v_4, v_5.$$

In graph theory, such an ordering is called a topological ordering of G_R . Since G_R is a directed acyclic graph (DAG), a topological ordering must exist.⁶ We may use depth first search (DFS) to traverse G_R to find the topological ordering. For the implementation, earlier we included a member "child nodes" in the Node class, but did not explain why. The reason is that to traverse G_R from x_1 , we must access children of each node.

Based on the above idea, we can have the following code to find a topological ordering.

Listing 2: Using depth first search to find a topological ordering

```
def topological_order(rootNode):
    def add_children(node):
        if node not in visited:
            visited.add(node)
            for child in node.child_nodes:
                add_children(child)
            ordering.append(node)
    ordering, visited = [], set()
    add_children(rootNode)
    return list(reversed(ordering))
```

The function add_children implements the depth-first-search of a DAG. From the input node, it sequentially calls itself by using each child as the input. This way explores all nodes reachable from the input node. After that, we append the input node to the end of the output list. Also, we must maintain a set of visited nodes to ensure that each node is included in the ordering exactly once. For our example, the input in calling the above function is x_1 , which is also the root node of G_R . The left-most path of the depth-first search is

$$x_1 \to v_1 \to v_4 \to v_5, \tag{8}$$

so v_5 is added first. In the end, we get the following list

$$[v_5, v_4, v_1, v_2, x_1].$$

⁶We do not get into details, but a proof can be found in Kleinberg and Tardos (2005).

Then, by reversing the list, a node always comes before its children. One may wonder whether we can add a node to the list before adding its child nodes. This way, we have a simpler implementation without needing to reverse the list in the end. However, this setting may fail to generate a topological ordering. We obtain the following list for our example:

$$[x_1, v_1, v_4, v_5, v_2].$$

A violation occurs because v_2 does not appear before its child v_4 . The key reason is that in a DFS path, a node may point to another node that was added earlier through a different path. Then this node becomes after one of its children. For our example,

$$x_1 \rightarrow v_2 \rightarrow v_4 \rightarrow v_5$$

is a path processed after the path in (8). Thus, v_2 is added after v_4 and a violation occurs. Reversing the list can resolve the problem. To this end, in the function add_children, we must append the input node in the end.

In automatic differentiation, methods based on the topological ordering are called *tape-based* methods. They are used in some real-world implementations such as Tensorflow. The ordering is regarded as a tape. We read the nodes one by one from the beginning of the sequence (tape) to calculate the derivative value.

Based on the obtained ordering, subsequently let us see how to compute each \dot{v} .

4.2 Computing the Partial Derivative

Earlier, by the chain rule, we have

$$\dot{v}_5 = \frac{\partial v_5}{\partial v_4} \dot{v}_4 + \frac{\partial v_5}{\partial v_3} \dot{v}_3.$$

In Section 4.1, we mentioned that

$$\dot{v}_4$$
 and \dot{v}_3

should be ready before calculating \dot{v}_5 . For

$$\frac{\partial v_5}{\partial v_4}$$
 and $\frac{\partial v_5}{\partial v_3}$,

we are able to calculate and store them when v_5 is created. The reason is that from

$$v_5(v_4, v_3) = v_4 - v_3,$$

we know

$$\frac{\partial v_5}{\partial v_4} = 1$$
 and $\frac{\partial v_5}{\partial v_3} = -1$.

A general form of our calculation is

$$\dot{v} = \sum_{u \in v' \text{s parents}} \frac{\partial v}{\partial u} \dot{u}. \tag{9}$$

The second term, $\dot{u} = \frac{\partial u}{\partial x_1}$, comes from past calculation due to the topological ordering. We can calculate the first term because u is one of v's parent(s) and we know the operation at v. For example, we have $v_4 = v_1 \times v_2$, so

$$\frac{\partial v_4}{\partial v_1} = v_2 \text{ and } \frac{\partial v_4}{\partial v_2} = v_1.$$

These values can be immediately computed and stored when we construct the computational graph. Therefore, we add one member "gradient w.r.t. parents" to our Node class. In addition, we need a member "partial derivative" to store the accumulated sum in the calculation of (9). In the end, this member stores the \dot{v}_i value. Details of the derivative evaluation are in Listing 4. The complete list of members of our node class is in the following table.

member	data type	example for Node v_2
numerical value	float	10
parent nodes	List[Node]	$[x_1, x_2]$
child nodes	List[Node]	$[v_4]$
operator	string	"mul"
gradient w.r.t parents	List[float]	[5,2]
partial derivative	float	5

We update the mul function accordingly.

Listing 3: The wrapping function "mul". The change from Listing 1 is in red color.

```
def mul(node1, node2):
    value = node1.value * node2.value
    parent_nodes = [node1, node2]
    newNode = Node(value, parent_nodes, "mul")

    newNode.grad_wrt_parents = [node2.value,node1.value]
    node1.child_nodes.append(newNode)
    node2.child_nodes.append(newNode)
    return newNode
```

As shown above, we must compute

$$\frac{\partial \text{ newNode}}{\partial \text{ parentNode}}$$

for each parent node in constructing a new child node. Here are some examples other than the mul function:

• For add(node1, node2), we have

$$\frac{\partial \ \text{newNode}}{\partial \ \text{Node1}} = \frac{\partial \ \text{newNode}}{\partial \ \text{Node2}} = 1,$$

so the red line is replaced by

newNode.grad_wrt_parents = [1., 1.].

• For log(node), we have

$$\frac{\partial \text{ newNode}}{\partial \text{ Node}} = \frac{1}{\text{Node.value}},$$

so the red line becomes

```
newNode.grad_wrt_parents = [1/node.value].
```

Now, we know how to get each term in (9), i.e., the chain rule for calculating \dot{v} . Therefore, if we follow the topological ordering, all \dot{v}_i (i.e., partial derivatives with respect to x_1) can be calculated. An implementation to compute the partial derivatives is as follows. Here we store the resulting value in the member partial_derivative of each node.

Listing 4: Evaluating derivatives

```
def forward(rootNode):
    rootNode.partial_derivative = 1
    ordering = topological_order(rootNode)
    for node in ordering[1:]:
        partial_derivative = 0
        for i in range(len(node.parent_nodes)):
            dnode_dparent = node.grad_wrt_parents[i]
            dparent_droot = node.parent_nodes[i].partial_derivative
            partial_derivative += dnode_dparent * dparent_droot
            node.partial_derivative = partial_derivative
```

4.3 Summary

The procedure for forward mode includes three steps:

- 1. Create the computational graph
- 2. Find a topological order of the graph associated with x_1
- 3. Compute the partial derivative with respect to x_1 along the topological order

We discuss not only how to run each step but also what information we should store. This is a minimal implementation to demonstrate the forward mode automatic differentiation.

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