Topics:

- Jacobians
- Optimization

CS 4803-DL / 7643-A ZSOLT KIRA

• Assignment Due Feb 5th

- Resources:
 - These lectures
 - Matrix calculus for deep learning
 - <u>Gradients notes</u> and <u>MLP/ReLU Jacobian notes</u>.
 - <u>Assignment</u> (@41) and <u>matrix calculus</u> (@46)
- **Project:** Teaming thread on piazza

 Schedule 	9:
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Graded Deliverabl 💌	Release Date / 💌	Due Date ES 💌
Start of Term	10-Jan	
A1	14-Jan	5-Feb
A2	30-Jan	20-Feb
A3	20-Feb	6-Mar
Project Proposal	5-Feb	13-Mar
A4	9-Mar	3-Apr
Project Report	5-Apr	1-May

To develop a general algorithm for this, we will view the function as a **computation graph**

Graph can be any **directed acyclic** graph (DAG)

 Modules must be differentiable to support gradient computations for gradient descent

A **training algorithm** will then process this graph, **one module at a time**



Adapted from figure by Marc'Aurelio Ranzato, Yann LeCun



A General Framework













Note that we must store the intermediate outputs of all layers!

This is because we will need them to compute the gradients (the gradient equations will have terms with the output values in them)





Step 1: Compute Loss on Mini-Batch: Forward PassStep 2: Compute Gradients wrt parameters: Backward Pass







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Step 2: Compute Gradients wrt parameters: Backward Pass

Step 3: Use gradient to update all parameters at the end



$$w_i = w_i - \alpha \frac{\partial L}{\partial w_i}$$

Backpropagation is the application of gradient descent to a computation graph via the chain rule!





• We want to to compute:
$$\left\{\frac{\partial L}{\partial h^{\ell-1}}, \frac{\partial L}{\partial W}\right\}$$



• We will use the *chain rule* to do this:

Chain Rule: $\frac{\partial z}{\partial x} = \frac{\partial z}{\partial y} \cdot \frac{\partial y}{\partial x}$

Computing the Gradients of Loss





Adapted from figure by Marc'Aurelio Ranzato, Yann LeCun

Computing the Gradients of Loss



Conventions:

• Size of derivatives for scalars, vectors, and matrices: Assume we have scalar $s \in \mathbb{R}^1$, vector $v \in \mathbb{R}^m$, i.e. $v = [v_1, v_2, ..., v_m]^T$ and matrix $M \in \mathbb{R}^{k \times \ell}$



Dimensionality of Derivatives



 $\begin{array}{c} X \in \mathbb{R}' \xrightarrow{g_1(f)} 2 \in \mathbb{R}' \xrightarrow{g_2(f)} Y \in \mathbb{R}' \\ \xrightarrow{g_2(f)} Y = g_2(g_1(x)) \end{array}$











Vector Case



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Jacobian View of Chain Rule





Graphical View of Chain Rule

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- Binary label: $y \in \{-1, +1\}$
- Parameters: $w \in \mathbb{R}^D$

• Output prediction:
$$p(y = 1|x) = \frac{1}{1 + e^{-w^T x}}$$



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We have discussed **computation** graphs for generic functions

Machine Learning functions (input -> model -> loss function) is also a computation graph

We can use the **computed gradients from backprop/automatic differentiation** to update the weights!



Neural Network Computation Graph





Automatic differentiation:

- Carries out this procedure for us on arbitrary graphs
- Knows derivatives of primitive functions
- As a result, we just define these (forward) functions and don't even need to specify the gradient (backward) functions!

$$\bar{L} = 1$$

$$\bar{p} = \frac{\partial L}{\partial p} = -\frac{1}{p}$$

where $p = \sigma(w^T x)$ and $\sigma(x) = \frac{1}{1+e^{-x}}$

$$\bar{u} = \frac{\partial L}{\partial u} = \frac{\partial L}{\partial p} \frac{\partial p}{\partial u} = \bar{p} \sigma(1-\sigma)$$

$$\bar{w} = \frac{\partial L}{\partial w} = \frac{\partial L}{\partial u} \frac{\partial u}{\partial w} = \bar{u}x^T$$

We can do this in a combined way to see all terms together:

$$\overline{w} = \frac{\partial L}{\partial p} \frac{\partial p}{\partial u} \frac{\partial u}{\partial w} = -\frac{1}{\sigma(w^T x)} \sigma(w^T x) (1 - \sigma(w^T x)) x^T$$
$$= -(1 - \sigma(w^T x)) x^T$$

This effectively shows gradient flow along path from L to w





The chain rule can be computed as a **series of scalar, vector, and matrix linear algebra operations**



Extremely efficient in graphics processing units (GPUs)



Vectorized Computations





$$L = 1$$
$$\overline{p} = \frac{\partial L}{\partial p} = -\frac{1}{p}$$

where $p = \sigma(w^T x)$ and $\sigma(x) = \frac{1}{1+e^{-x}}$

 $\overline{u} = \frac{\partial L}{\partial u} = \frac{\partial L}{\partial p} \quad \frac{\partial p}{\partial u} = \overline{p} \; \sigma(1 - \sigma)$ $\overline{w} = \frac{\partial L}{\partial w} = \frac{\partial L}{\partial u} \quad \frac{\partial u}{\partial w} = \overline{u} x^{T}$

We can do this in a combined way to see all terms together:

$$\begin{split} \overline{w} &= \frac{\partial L}{\partial p} \frac{\partial p}{\partial u} \frac{\partial u}{\partial w} = -\frac{1}{\sigma(w^T x)} \sigma(w^T x) (1 - \sigma(w^T x)) x^T \\ &= -\left(1 - \sigma(w^T x)\right) x^T \end{split}$$

This effectively shows gradient flow along path from $\mathit{L}\, \mathrm{to}\, \mathit{w}$

Computation Graph / Global View of Chain Rule



Different Views of Equivalent Ideas

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Fully Connected (FC) Layer



Full Jacobian of ReLU layer is **large** (output dim x input dim)

- But again it is sparse
- Only diagonal values non-zero because it is element-wise
- An output value affected only by corresponding input value

Max function **funnels gradients through selected max**

Gradient will be zero if input
 <= 0







Backpropagation and Automatic Differentiation



Backpropagation does not really spell out how to **efficiently** carry out the necessary computations

But the idea can be applied to **any directed acyclic graph** (DAG)

 Graph represents an ordering constraining which paths must be calculated first

Given an ordering, we can then iterate from the last module backwards, **applying the chain rule**

- We will store, for each node, its gradient outputs for efficient computation
- We will do this automatically by computing backwards function for primitives and as you write code, express the function with them

This is called reverse-mode automatic differentiation







Computation = Graph

- Input = Data + Parameters
- Output = Loss
- Scheduling = Topological ordering

Auto-Diff

 A family of algorithms for implementing chain-rule on computation graphs

Deep Learning = Differentiable Programming





We want to find the **partial derivative of output f** (output) with respect to **all intermediate variables**

Assign intermediate variables

Simplify notation: Denote bar as: $\overline{a_3} = \frac{\partial f}{\partial a_3}$

Start at end and move backward



Example





$$\overline{a_1} = \frac{\partial f}{\partial a_1} = \frac{\partial f}{\partial a_3} \quad \frac{\partial a_3}{\partial a_1} = \frac{\partial f}{\partial a_3} \quad \frac{\partial (a_1 + a_2)}{\partial a_1} = \frac{\partial f}{\partial a_3} \quad \mathbf{1} = \overline{a_3}$$
$$\overline{a_2} = \frac{\partial f}{\partial a_2} = \frac{\partial f}{\partial a_3} \quad \frac{\partial a_3}{\partial a_2} = \overline{a_3}$$

Addition operation distributes gradients along all paths!

Patterns of Gradient Flow: Addition





Multiplication operation is a gradient switcher (multiplies it by the values of the other term)

$$\overline{x_2} = \frac{\partial f}{\partial a_2} \ \frac{\partial a_2}{\partial x_2} = \frac{\partial f}{\partial a_2} \ \frac{\partial (x_1 x_2)}{\partial x_2} = \overline{a_2} x_1$$

$$\overline{x_1} = \frac{\partial f}{\partial a_2} \ \frac{\partial a_2}{\partial x_1} = \overline{a_2} x_2$$

Patterns of Gradient Flow: Multiplication



Several other patterns as well, e.g.:

Max operation **selects** which path to push the gradients through

- Gradient flows along the path that was "selected" to be max
- This information must be recorded in the forward pass



The flow of gradients is one of the most important aspects in deep neural networks

• If gradients **do not flow backwards properly**, learning slows or stops!

Patterns of Gradient Flow: Other



- Key idea is to explicitly store computation graph in memory and corresponding gradient functions
- Nodes broken down to basic primitive computations (addition, multiplication, log, etc.) for which corresponding derivative is known







Note that we can also do **forward mode** automatic differentiation

Start from **inputs** and propagate gradients forward

Complexity is proportional to input size

- Memory savings (all forward pass, no need to store activations)
- However, in most cases our inputs (images) are large and outputs (loss) are small



Automatic Differentiation





See https://www.cc.gatech.edu/classes/AY2020/cs7643 spring/slides/autodiff forward reverse.pdf

Forward Mode Autodifferentiation



A graph is created on the fly from torch.autograd import Variable x = Variable(torch.randn(1, 20)) prev_h = Variable(torch.randn(1, 20)) W_h = Variable(torch.randn(20, 20)) W_x = Variable(torch.randn(20, 20)) i2h = torch.mm(W_x, x.t()) h2h = torch.mm(W_h, prev_h.t())



(Note above)

next h = i2h + h2h

Computation Graphs in PyTorch



Back-propagation uses the dynamically built graph

from torch.autograd import Variable

x = Variable(torch.randn(1, 20))
prev_h = Variable(torch.randn(1, 20))
W_h = Variable(torch.randn(20, 20))
W_x = Variable(torch.randn(20, 20))

```
i2h = torch.mm(W_x, x.t())
h2h = torch.mm(W_h, prev_h.t())
next_h = i2h + h2h
next_h = next_h.tanh()
```

next_h.backward(torch.ones(1, 20))



From pytorch.org









Slide Credit: Fei-Fei Li, Justin Johnson, Serena Yeung, CS 231n

Neural Turing Machine



Figure reproduced with permission from a Twitter post by Andrej Karpathy.





- Computation graphs are not limited to mathematical functions!
- Can have control flows (if statements, loops) and backpropagate through algorithms!
- Can be done dynamically so that gradients are computed, then nodes are added, repeat
- Differentiable programming



Adapted from figure by Andrej Karpathy

Power of Automatic Differentiation



Optimization of Deep Neural Networks Overview



Backpropagation, and automatic differentiation, allows us to optimize **any** function composed of differentiable blocks

- No need to modify the learning algorithm!
- The complexity of the function is only limited by **computation and memory**



A network with two or more hidden layers is often considered a **deep** model

Depth is important:

- Structure the model to represent an inherently compositional world
- Theoretical evidence that it leads to parameter efficiency
- Gentle dimensionality reduction (if done right)





Importance of Depth

There are still many design decisions that must be made:

- Architecture
- Data Considerations
- Training and Optimization
- Machine Learning Considerations







We must design the **neural network** architecture:

- What modules (layers) should we use?
- How should they be connected together?
- Can we use our domain knowledge to add architectural biases?





















Example Architectures



As in traditional machine learning, **data** is key:

- Should we pre-process the data?
- Should we **normalize** it?
- Can we augment our data by adding noise or other perturbations?



Even given a good neural network architecture, we need a **good optimization algorithm to find good weights**

- What optimizer should we use?
 - Different optimizers make different weight updates depending on the gradients
- How should we initialize the weights?
- What regularizers should we use?
- What loss function is appropriate?

Optimization Considerations

Machine Learning Considerations

The practice of machine learning is complex: For your particular application you have to **trade off** all of the considerations together

- Trade-off between model capacity (e.g. measured by # of parameters) and amount of data
- Adding appropriate biases based on knowledge of the domain

Architectural Considerations 00

Determining what modules to use, and how to connect them is part of the **architectural design**

- Guided by the type of data used and its characteristics
 - Understanding your data is always the first step!
- Lots of data types (modalities) already have good architectures
 - Start with what others have discovered!
- The flow of gradients is one of the key principles to use when analyzing layers

Combination of linear and non-linear layers

$$w_1^T(w_2^T(w_3^Tx)) = w_4^Tx$$

-0.2

-4

-2

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Non-linear layers are crucial

 Composition of non-linear layers enables complex transformations of the data

Linear and Non-Linear Modules

Several aspects that we can **analyze**:

- Min/Max
- Correspondence between input & output statistics
- **Gradients**
 - At initialization (e.g. small values)
 - At extremes
- Computational complexity

Analysis of Non-Linear Function

- Min: 0, Max: 1
- Output always positive
- Saturates at both ends
- Gradients
 - Vanishes at both end
 - Always positive
- Computation: Exponential term

$$\sigma(x)=\frac{1}{1+e^{-x}}$$

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Sigmoid Function

- Min: -1, Max: 1
 - Centered
- Saturates at **both ends**
- Gradients
 - Vanishes at both end
 - Always positive
- Still somewhat computationally heavy

$$h^{\ell} = tanh(h^{\ell-1})$$

- Min: 0, Max: Infinity
- Output always positive
- **No saturation** on positive end!
- Gradients
 - 0 if $x \le 0$ (dead ReLU)
 - Constant otherwise (does not vanish)
- Cheap to compute (max)

$$h^{\ell} = max(0, h^{\ell-1})$$

Rectified Linear Unit

Min: -Infinity, Max: Infinity

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- Learnable parameter!
- No saturation
- Gradients
 - No dead neuron
- Still cheap to compute

$$h^{\ell} = max(\alpha h^{\ell-1}, h^{\ell-1})$$

Leaky ReLU

Selecting a Non-Linearity

Which **non-linearity** should you select?

- Unfortunately, no one activation
 function is best for all applications
- ReLU is most common starting point
 - Sometimes leaky ReLU can make a big difference
- Sigmoid is typically avoided unless clamping to values from [0,1] is needed

- **Backpropagation:** Recursive, modular algorithm for chain rule + gradient descent
- When we move to vectors and matrices:
 - Computation graph (composition of functions) => Multiplication of Jacobians along path
- Automatic differentiation:
 - Reduction of modules to simple operations we know (simple multiplication, etc.)
 - Automatically build computation graph in background as write code
 - Automatically compute gradients via backward pass
- We now have a generic algorithm! Considerations:
 - Architecture
 - Data Considerations
 - Training and Optimization
 - Machine Learning Considerations

