Topics:

• Optimization

# CS 4644-DL / 7643-A ZSOLT KIRA

- Assignment 1 Due Friday!!!
  - DO NOT SEARCH FOR CODE!!!!
- Note: Syllabus will shift!!!
- Assignment 2
  - Implement convolutional neural networks
- **Piazza:** Start with public posts so that others can benefit!
  - Doesn't mean don't post!
- Meta OH: Data wrangling Friday 02/2 3pm ET
  - Full schedule and discussions on <a href="https://ai-learning.org/">https://ai-learning.org/</a>
  - See dropbox link on piazza @109 for first office hours and link to their lessons





So Far



- Gradient Descent
- Compute gradients via chain rule
  - Backpropagation
  - Computation
    - Graph +Automatic

Differentiation



↓

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



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

#### Modularized implementation: forward / backward API



ass M	<pre>ultiplyGate(object):</pre>
def	forward(x,y):
	z = x*y
	<pre>self.x = x # must keep these around!</pre>
	self.y = y
	return z
def	backward(dz):
	dx = self.y * dz # [dz/dx * dL/dz]
	dy = self.x * dz # [dz/dy * dL/dz]
	return [dx, dy]





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



**Example** 



#### 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())
next_h = i2h + h2h
```



(Note above)









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 L to w

#### Computation Graph / Global View of Chain Rule





#### **Computational / Tensor View**



#### **Graph View**



Backpropagation View (Recursive Algorithm)

**Different Views of Equivalent Ideas** 



There are still many design decisions that must be made:

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



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
- Combination of only linear layers has same representational power as one linear layer
- Non-linear layers are crucial
  - Composition of non-linear layers enables complex transformations of the data

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







**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







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



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





Georg



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



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





- Activation functions is still area of research!
  - Though many don't catch on

 In Transformer architectures, other activations such as GeLU is common



From "Gaussian Error Linear Units (GELUs)", Hendrycks & Gimpel





### **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



# Demo

# <u>http://playground.tensorflow.org</u>





# Initialization



### **Initializing the Parameters**

# The parameters of our model must be initialized to something

- Initialization is extremely important!
  - Determines how statistics of outputs (given inputs) behave
  - Determines how well gradients flow in the beginning of training (important)
  - Could limit use of full capacity of the model if done improperly
- Initialization that is close to a good (local) minima will converge faster and to a better solution





Initializing values to a constant value leads to a degenerate solution!

- What happens to the weight updates?
- Each node has the same input from previous layers so gradients will be the same
- As a results, all weights will be updated to the same exact values

input layer 1 hidden layer 2

 $w_i = c \quad \forall i$ 





#### Common approach is small normally distributed random numbers

- E.g.  $N(\mu, \sigma)$  where  $\mu = 0, \sigma = 0.01$
- Small weights are preferred since no feature/input has prior importance
- Keeps the model within the linear region of most activation functions











# Deeper networks (with many layers) are more sensitive to initialization

- With a deep network, activations (outputs of nodes) get smaller
  - Standard deviation reduces significantly
- Leads to small updates smaller values multiplied by upstream gradients



Distribution of activation values of a network with tanh nonlinearities, for increasingly deep layers

From "Understanding the difficulty of training deep feedforward neural networks." AISTATS, 2010.



**Limitation of Small Weights** 



# Ideally, we'd like to maintain the variance at the output to be similar to that of input!

This condition leads to a simple initialization rule, sampling from uniform distribution:

Uniform 
$$\left(-\frac{\sqrt{6}}{n_j+n_{j+1}},+\frac{\sqrt{6}}{n_j+n_{j+1}}\right)$$

Where n<sub>j</sub> is fan-in

 (number of input nodes)
 and n<sub>j+1</sub> is fan-out
 (number of output nodes)



Distribution of activation values of a network with tanh nonlinearities, for increasingly deep layers

From "Understanding the difficulty of training deep feedforward neural networks." AISTATS, **2010.** 



## **Xavier Initialization**

In practice, simpler versions perform empirically well:

$$N(0,1) * \sqrt{\frac{1}{n_j}}$$

- This analysis holds for tanh or similar activations.
- Similar analysis for ReLU activations leads to:

$$N(0,1) * \sqrt{\frac{1}{n_j/2}}$$

"Delving Deep into Rectifiers: Surpassing Human-Level Performance on ImageNet Classification", ICCV, 2015.





### Summary

Key takeaway: Initialization matters!

- Determines the activation (output) statistics, and therefore gradient statistics
- If gradients are small, no learning will occur and no improvement is possible!
- Important to reason about output/gradient statistics and analyze them for new layers and architectures





Normalization, Preprocessing, and Augmentation



#### **Importance of Data**

In deep learning, **data drives learning** of features and classifier

- Its characteristics are therefore extremely important
- Always understand your data!
- Relationship between output statistics, layers such as nonlinearities, and gradients is important





Just like initialization, normalization can improve gradient flow and learning

Typically **normalization methods** apply:

- Subtract mean, divide by standard deviation (most common)
  - This can be done per dimension
- Whitening, e.g. through Principle Component Analysis (PCA) (not common)





Figure from slides by Fei-Fei Li, Justin Johnson, Serena Yeung, CS 231n





- We can try to come up with a *layer* that can normalize the data across the neural network
- Given: A mini-batch of data  $[B \times D]$  where B is batch size
- Compute mean and variance for each dimension d

**Input:** Values of x over a mini-batch:  $\mathcal{B} = \{x_{1...m}\}$ ; Parameters to be learned:  $\gamma, \beta$  **Output:**  $\{y_i = BN_{\gamma,\beta}(x_i)\}$   $\mu_{\mathcal{B}} \leftarrow \frac{1}{m} \sum_{i=1}^m x_i$  // mini-batch mean  $\sigma_{\mathcal{B}}^2 \leftarrow \frac{1}{m} \sum_{i=1}^m (x_i - \mu_{\mathcal{B}})^2$  // mini-batch variance

From: Batch Normalization: Accelerating Deep Network Training by Reducing Internal Covariate Shift, Sergey Ioffe, Christian Szegedy

#### Making Normalization a Layer

#### Normalize data

$$\widehat{x}_i = \frac{x_i - \mu_B}{\sqrt{\sigma_B^2 + \epsilon}}$$

**Note:** This part does not involve new parameters

**Input:** Values of x over a mini-batch:  $\mathcal{B} = \{x_{1...m}\};$ Parameters to be learned:  $\gamma$ ,  $\beta$ **Output:**  $\{y_i = BN_{\gamma,\beta}(x_i)\}$  $\mu_{\mathcal{B}} \leftarrow \frac{1}{m} \sum_{i=1}^{m} x_i$ // mini-batch mean  $\sigma_{\mathcal{B}}^2 \leftarrow \frac{1}{m} \sum_{i=1}^m (x_i - \mu_{\mathcal{B}})^2$ // mini-batch variance  $\widehat{x}_i \leftarrow \frac{x_i - \mu_{\mathcal{B}}}{\sqrt{\sigma_r^2 + \epsilon}}$ // normalize

From: Batch Normalization: Accelerating Deep Network Training by Reducing Internal Covariate Shift, Sergey Ioffe, Christian Szegedy





- We can give the model flexibility through
   learnable parameters γ (scale) and β (shift)
- Network can learn to not normalize if necessary!
- This layer is called a
   Batch Normalization
   (BN) layer

**Input:** Values of x over a mini-batch:  $\mathcal{B} = \{x_{1...m}\}$ ; Parameters to be learned:  $\gamma$ ,  $\beta$ **Output:**  $\{y_i = BN_{\gamma,\beta}(x_i)\}$  $\mu_{\mathcal{B}} \leftarrow \frac{1}{m} \sum_{i=1}^{m} x_i$ // mini-batch mean  $\sigma_{\mathcal{B}}^2 \leftarrow \frac{1}{m} \sum_{i=1}^m (x_i - \mu_{\mathcal{B}})^2$ // mini-batch variance  $\widehat{x}_i \leftarrow \frac{x_i - \mu_{\mathcal{B}}}{\sqrt{\sigma_{\mathcal{B}}^2 + \epsilon}}$ // normalize

 $y_i \leftarrow \gamma \widehat{x}_i + \beta \equiv \mathrm{BN}_{\gamma,\beta}(x_i)$  // scale and shift

From: Batch Normalization: Accelerating Deep Network Training by Reducing Internal Covariate Shift, Sergey Ioffe, Christian Szegedy

### Learnable Scaling and Offset



### **Some Complexities of BN**

**During inference**, stored mean/variances calculated on training set are used

**Sufficient batch sizes** must be used to get stable per-batch estimates during training

- This is especially an issue when using multi-GPU or multi-machine training
- Use torch.nn.SyncBatchNorm to estimate batch statistics in these settings





# Normalization especially important before **non-linearities**!

 Very low/high values (unnormalized/imbalanced data) cause saturation









#### Batch normalization unstable for small batch sizes



From: Group Normalization, Wu et al.





### **Generalization of BN**

# There are many variations of batch normalization

 See Convolutional Neural Network lectures for an example

#### **Resource:**

ML Explained - Normalization



# **Optimizers**



# Deep learning involves complex, compositional, non-linear functions

The **loss landscape** is extremely **nonconvex** as a result

There is **little direct theory** and a **lot of intuition/rules of thumbs** instead

 Some insight can be gained via theory for simpler cases (e.g. convex settings)







It used to be thought that existence of local minima is the main issue in optimization

# There are other **more impactful issues**:

- Noisy gradient estimates
- Saddle points
- Ill-conditioned loss surface



From: Identifying and attacking the saddle point problem in highdimensional non-convex optimization, Dauphi et al., 2014.





We use a subset of the data at each iteration to calculate the loss (& gradients)

- This is an unbiased estimator but can have high variance
- This results in noisy steps in gradient descent

 $L = \frac{1}{M} \sum_{i} L(f(x_i, W), y_i)$ 





Several **loss surface geometries** are difficult for optimization

**Several types of minima:** Local minima, plateaus, saddle points

**Saddle points** are those where the gradient of orthogonal directions are zero

 But they disagree (it's min for one, max for another)









- Gradient descent takes a step in the steepest direction (negative gradient)
- Intuitive idea: Imagine a ball rolling down loss surface, and use momentum to pass flat surfaces

 $v_i = \beta v_{i-1} + \frac{\partial L}{\partial w_{i-1}}$  Update Velocity (starts as 0,  $\beta = 0.99$ )

 $w_i = w_{i-1} - \alpha v_i$  Update Weights

• Generalizes SGD ( $\beta = 0$ )

$$w_i = w_{i-1} - \alpha \frac{\partial L}{\partial w_i}$$



Adding Momentum



Velocity term is an exponential moving average of the gradient

$$v_i = \beta v_{i-1} + \frac{\partial L}{\partial w_{i-1}}$$

$$v_{i} = \beta (\beta v_{i-2} + \frac{\partial L}{\partial w_{i-2}}) + \frac{\partial L}{\partial w_{i-1}}$$
$$= \beta^{2} v_{i-2} + \beta \frac{\partial L}{\partial w_{i-2}} + \frac{\partial L}{\partial w_{i-1}}$$

There is a general class of accelerated gradient methods, with some theoretical analysis (under assumptions)

**Accelerated Descent Methods** 



## **Equivalent formulation:**

$$egin{aligned} &v_i = eta v_{i-1} - lpha rac{\partial L}{\partial w_{i-1}} & ext{Update Velocity} \ & ext{(starts as 0)} \end{aligned}$$







**Key idea:** Rather than combining velocity with current gradient, go along velocity **first** and then calculate gradient at new point

 We know velocity is probably a reasonable direction

$$\widehat{w}_{i-1} = w_{i-1} + \beta v_{i-1}$$

$$v_i = \beta v_{i-1} + \frac{\partial L}{\partial \widehat{w}_{i-1}}$$

$$w_i = w_{i-1} - \alpha v_i$$



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





#### **Momentum**

Note there are **several equivalent formulations** across deep learning frameworks!

#### **Resource:**

https://medium.com/the-artificialimpostor/sgd-implementation-inpytorch-4115bcb9f02c





- Various mathematical ways to characterize the loss landscape
- If you liked **Jacobians**... meet:



 Gives us information about the curvature of the loss surface

Hessian and Loss Curvature



**Condition number** is the ratio of the largest and smallest eigenvalue

 Tells us how different the curvature is along different dimensions

If this is high, SGD will make **big** steps in some dimensions and **small** steps in other dimension

Second-order optimization methods divide steps by curvature, but expensive to compute







### **Per-Parameter Learning Rate**

**Idea:** Have a dynamic learning rate for each weight

# Several flavors of **optimization algorithms**:

- RMSProp
- Adagrad
- Adam

SGD can achieve similar results in many cases but with much more tuning



Idea: Use gradient statistics to reduce learning rate across iterations

**Denominator:** Sum up gradients over iterations

Directions with **high curvature will have higher gradients**, and learning rate will reduce  $G_{i} = G_{i-1} + \left(\frac{\partial L}{\partial w_{i-1}}\right)^{2}$  $w_{i} = w_{i-1} - \frac{\alpha}{\sqrt{G_{i} + \epsilon}} \frac{\partial L}{\partial w_{i-1}}$ 

As gradients are accumulated learning rate will go to zero

Duchi, et al., "Adaptive Subgradient Methods for Online Learning and Stochastic Optimization"





**Solution:** Keep a moving average of squared gradients!

Does not saturate the learning rate

$$G_{i} = \beta G_{i-1} + (1 - \beta) \left(\frac{\partial L}{\partial w_{i-1}}\right)^{2}$$

$$\alpha \qquad \partial L$$

$$w_i = w_{i-1} - \frac{\partial}{\sqrt{G_i + \epsilon}} \frac{\partial u}{\partial w_{i-1}}$$





**Combines ideas** from above algorithms

Maintains both first and second moment statistics for gradients

$$v_i = \beta_1 v_{i-1} + (1 - \beta_1) \left( \frac{\partial L}{\partial w_{i-1}} \right)$$

$$G_{i} = \beta_{2} G_{i-1} + (1 - \beta_{2}) \left(\frac{\partial L}{\partial w_{i-1}}\right)^{2}$$

$$w_i = w_{i-1} - \frac{\alpha v_i}{\sqrt{G_i + \epsilon}}$$

But unstable in the beginning (one or both of moments will be tiny values)

Kingma and Ba, "Adam: A method for stochastic optimization", ICLR 2015





**Solution:** Time-varying bias correction

Typically  $\beta_1 = 0.9$ ,  $\beta_2 = 0.999$ 

So  $\hat{v}_i$  will be small number divided by (1-0.9=0.1) resulting in more reasonable values (and  $\hat{G}_i$  larger)

$$v_{i} = \beta_{1} v_{i-1} + (1 - \beta_{1}) \left(\frac{\partial L}{\partial w_{i-1}}\right)$$
$$G_{i} = \beta_{2} G_{i-1} + (1 - \beta_{2}) \left(\frac{\partial L}{\partial w_{i-1}}\right)^{2}$$

$$\widehat{v}_{i} = \frac{v_{i}}{1 - \beta_{1}^{t}} \quad \widehat{G}_{i} = \frac{G_{i}}{1 - \beta_{2}^{t}}$$
$$w_{i} = w_{i-1} - \frac{\alpha \,\widehat{v}_{i}}{\sqrt{\widehat{G}_{i} + \epsilon}}$$





Optimizers behave differently depending on landscape

Different behaviors such as **overshooting**, **stagnating**, **etc**.

Plain SGD+Momentum can generalize better than adaptive methods, but requires more tuning

See: Luo et al., Adaptive Gradient Methods with Dynamic Bound of Learning Rate, ICLR 2019



From: https://mlfromscratch.com/optimizers-explained/#/





First order optimization methods have learning rates

Theoretical results rely on **annealed** learning rate

Several schedules that are typical:

- Graduate student!
- Step scheduler
- Exponential scheduler
- Cosine scheduler



From: Leslie Smith, "Cyclical Learning Rates for Training Neural Networks"

## Learning Rate Schedules



First order optimization methods have learning rates

Theoretical results rely on **annealed** learning rate

Several schedules that are typical:

- Graduate student!
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From: Leslie Smith, "Cyclical Learning Rates for Training Neural Networks"

## Learning Rate Schedules

