

# **Machine Learning – Lecture 12**

### **Tricks of the Trade**

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# **Course Outline**

- Fundamentals
  - Bayes Decision Theory
  - Probability Density Estimation
- Classification Approaches
  - Linear Discriminants
  - Support Vector Machines
  - Ensemble Methods & Boosting
  - Random Forests
- Deep Learning
  - Foundations
  - Convolutional Neural Networks
  - Recurrent Neural Networks





# **Topics of This Lecture**

- Recap: Optimization
  - Effect of optimizers
- Tricks of the Trade
  - Shuffling
  - Data Augmentation
  - Normalization
- Nonlinearities
  - Initialization
  - Advanced techniques
    - Batch Normalization
    - Dropout



### **Recap: Computational Graphs**



- Forward differentiation needs one pass per node. Reverse-mode differentiation can compute all derivatives in one single pass.
- $\Rightarrow$  Speed-up in  $\mathcal{O}(\#$ inputs) compared to forward differentiation!

Slide inspired by Christopher Olah

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# **Recap: Automatic Differentiation**

Approach for obtaining the gradients



- > Convert the network into a computational graph.
- Each new layer/module just needs to specify how it affects the forward and backward passes.
- Apply reverse-mode differentiation.
- $\Rightarrow$  Very general algorithm, used in today's Deep Learning packages

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# Recap: Choosing the Right Learning Rate

- Convergence of Gradient Descent
  - Simple 1D example

$$W^{(\tau-1)} = W^{(\tau)} - \eta \frac{\mathrm{d}E(W)}{\mathrm{d}W}$$

- » What is the optimal learning rate  $\eta_{
  m opt}$ ?
- If E is quadratic, the optimal learning rate is given by the inverse of the Hessian

$$\eta_{\rm opt} = \left(\frac{\mathrm{d}^2 E(W^{(\tau)})}{\mathrm{d}W^2}\right)^{-1}$$

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- Advanced optimization techniques try to approximate the Hessian by a simplified form.
- If we exceed the optimal learning rate, bad things happen!





Don't go beyond

this point!



### Separate, Adaptive Learning Rates

- Problem
  - In multilayer nets, the appropriate learning rates can vary widely between weights.
  - The magnitudes of the gradients are often very different for the different layers, especially if the initial weights are small.
    - $\Rightarrow$  Gradients can get very small in the early layers of deep nets.
  - The fan-in of a unit determines the size of the "overshoot" effect when changing multiple weights simultaneously to correct the same error.
    - The fan-in often varies widely between layers
  - Solution
    - Use a global learning rate, multiplied by a local gain per weight (determined empirically)

Slide adapted from Geoff Hinton





### Better Adaptation: RMSProp

- Motivation
  - The magnitude of the gradient can be very different for different weights and can change during learning.
  - This makes it hard to choose a single global learning rate.
  - For batch learning, we can deal with this by only using the sign of the gradient, but we need to generalize this for minibatches.
- Idea of RMSProp
  - Divide the gradient by a running average of its recent magnitude

$$MeanSq(w_{ij}, t) = 0.9MeanSq(w_{ij}, t-1) + 0.1\left(\frac{\partial E}{\partial w_{ij}}(t)\right)^{2}$$

> Divide the gradient by  $sqrt(MeanSq(w_{ij},t))$ .

# **Other Optimizers**

AdaGrad

AdaDelta

[Zeiler '12]

[Duchi '10]

Adam

[Ba & Kingma '14]

- Notes
  - All of those methods have the goal to make the optimization less sensitive to parameter settings.
  - Adam is currently becoming the quasi-standard

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### Example: Behavior in a Long Valley



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# Example: Behavior around a Saddle Point



# Visualization of Convergence Behavior



Machine Learning Winter '18

B. Leibe Image source: Aelc Radford, http://imgur.com/SmDARzn



### **Trick: Patience**

Saddle points dominate in high-dimensional spaces!



 $\Rightarrow$  Learning often doesn't get stuck, you just may have to wait...

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# Reducing the Learning Rate

- Final improvement step after convergence is reached
  - Reduce learning rate by a factor of 10.
  - Continue training for a few epochs.
  - Do this 1-3 times, then stop training.



#### Effect

- Turning down the learning rate will reduce the random fluctuations in the error due to different gradients on different minibatches.
- Be careful: Do not turn down the learning rate too soon!
  - Further progress will be much slower/impossible after that.



### Summary

- Deep multi-layer networks are very powerful.
- But training them is hard!
  - Complex, non-convex learning problem
  - Local optimization with stochastic gradient descent
- Main issue: getting good gradient updates for the lower layers of the network
  - Many seemingly small details matter!
  - Weight initialization, normalization, data augmentation, choice of nonlinearities, choice of learning rate, choice of optimizer,...
  - > In the following, we will take a look at the most important factors

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#### • Tricks of the Trade

- Shuffling
- Data Augmentation
- Normalization

#### Nonlinearities

- Initialization
- Advanced techniques
  - Batch Normalization
  - > Dropout

# Shuffling the Examples

Ideas

- Networks learn fastest from the most unexpected sample.
- $\Rightarrow$  It is advisable to choose a sample at each iteration that is most unfamiliar to the system.
  - E.g. a sample from a *different class* than the previous one.
  - This means, do not present all samples of class A, then all of class B.
- A large relative error indicates that an input has not been learned by the network yet, so it contains a lot of information.
- $\Rightarrow$  It can make sense to present such inputs more frequently.
  - But: be careful, this can be disastrous when the data are outliers.

#### Practical advice

When working with stochastic gradient descent or minibatches, make use of shuffling.

### **Data Augmentation**

- Idea
  - Augment original data with synthetic variations to reduce overfitting
- Example augmentations for images
  - Cropping
  - Zooming
  - Flipping
  - Color PCA













# **Data Augmentation**

- Effect
  - Much larger training set
  - Robustness against expected variations
- During testing
  - When cropping was used during training, need to again apply crops to get same image size.
  - Beneficial to also apply flipping during test.
  - Applying several ColorPCA variations can bring another ~1% improvement, but at a significantly increased runtime.



Augmented training data (from one original image)



### **Practical Advice**





### Normalization

- Motivation
  - Consider the Gradient Descent update steps

$$w_{kj}^{(\tau+1)} = w_{kj}^{(\tau)} - \eta \left. \frac{\partial E(\mathbf{w})}{\partial w_{kj}} \right|_{\mathbf{w}^{(\tau)}}$$

From backpropagation, we know that

$$\frac{\partial E}{\partial w_{ij}} = \frac{\partial z_j}{\partial w_{ij}} \frac{\partial E}{\partial z_j} = \mathbf{y}_i \frac{\partial E}{\partial z_j} \qquad \qquad \mathbf{O} \qquad \mathbf{O}$$

- > When all of the components of the input vector y<sub>i</sub> are positive, all of the updates of weights that feed into a node will be of the same sign.
   ⇒ Weights can only all increase or decrease together.
- $\Rightarrow$  Slow convergence



# Normalizing the Inputs

- Convergence is fastest if
  - The mean of each input variable over the training set is zero.
  - The inputs are scaled such that all have the same covariance.
  - Input variables are uncorrelated if possible.



- Advisable normalization steps (for MLPs only, not for CNNs)
  - Normalize all inputs that an input unit sees to zero-mean, unit covariance.
  - If possible, try to decorrelate them using PCA (also known as Karhunen-Loeve expansion).

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- Normalization is also important for intermediate layers
  - Symmetric sigmoids, such as tanh, often converge faster than the standard logistic sigmoid.
  - Recommended sigmoid:

$$f(x) = 1.7159 \tanh\left(\frac{2}{3}x\right)$$

 $\Rightarrow$  When used with transformed inputs, the variance of the outputs will be close to 1.



### Usage

#### Output nodes

- > Typically, a sigmoid or tanh function is used here.
  - Sigmoid for nice probabilistic interpretation (range [0,1]).
  - tanh for regression tasks
- Internal nodes
  - Historically, tanh was most often used.
  - > tanh is better than sigmoid for internal nodes, since it is already centered.
  - Internally, tanh is often implemented as piecewise linear function (similar to hard tanh and maxout).
  - > More recently: ReLU often used for classification tasks.



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# Effect of Sigmoid Nonlinearities

- Effects of sigmoid/tanh function
  - Linear behavior around 0
  - Saturation for large inputs



- If all parameters are too small
  - Variance of activations will drop in each layer
  - Sigmoids are approximately linear close to 0
  - Good for passing gradients through, but...
  - Gradual loss of the nonlinearity
  - $\Rightarrow$  No benefit of having multiple layers
- If activations become larger and larger
  - > They will saturate and gradient will become zero



- Squared error on sigmoid/tanh output function
  - Avoids penalizing "too correct" data points.
  - > But: zero gradient for confidently incorrect classifications!
  - $\Rightarrow$  Do not use L<sub>2</sub> loss with sigmoid outputs (instead: cross-entropy)!



# Extension: ReLU

- Another improvement for learning deep models
  - > Use Rectified Linear Units (ReLU)

$$g(a) = \max\left\{0, a\right\}$$

 Effect: gradient is propagated with a constant factor

$$\frac{\partial g(a)}{\partial a} = \begin{cases} 1, & a > 0\\ 0, & \text{else} \end{cases}$$



#### Advantages

- Much easier to propagate gradients through deep networks.
- We do not need to store the ReLU output separately
  - Reduction of the required memory by half compared to tanh!

#### $\Rightarrow$ ReLU has become the de-facto standard for deep networks.



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- Disadvantages / Limitations
  - > A certain fraction of units will remain "stuck at zero".
    - If the initial weights are chosen such that the ReLU output is 0 for the entire training set, the unit will never pass through a gradient to change those weights.
  - > ReLU has an offset bias, since its outputs will always be positive





### Further Extensions

Rectified linear unit (ReLU)
 g(a) = max{0, a}

- Leaky ReLU  $g(a) = \max{\beta a, a}$ 
  - > Avoids stuck-at-zero units
  - Weaker offset bias
  - ELU  $g(a) = \begin{cases} a, & x < 0\\ e^a - 1, & x \ge 0 \end{cases}$ 
    - No offset bias anymore
    - > BUT: need to store activations



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#### Initialization

- Advanced techniques
  - Batch Normalization
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# Initializing the Weights

- Motivation
  - The starting values of the weights can have a significant effect on the training process.
  - Weights should be chosen randomly, but in a way that the sigmoid is primarily activated in its linear region.
- Guideline (from [LeCun et al., 1998] book chapter)
  - Assuming that
    - The training set has been normalized

– The recommended sigmoid  $f(x) = 1.7159 \tanh\left(\frac{2}{3}x\right)$  is used the initial weights should be randomly drawn from a distribution (e.g., uniform or Normal) with mean zero and variance

$$\sigma_w^2 = \frac{1}{n_{in}}$$

where  $n_{in}$  is the fan-in (#connections into the node).



### **Historical Sidenote**

- Apparently, this guideline was either little known or misunderstood for a long time
  - A popular heuristic (also the standard in Torch) was to use

$$W \sim U\left[-\frac{1}{\sqrt{n_{in}}}, \frac{1}{\sqrt{n_{in}}}\right]$$

- This looks almost like LeCun's rule. However...
- When sampling weights from a uniform distribution [a,b]
  - Keep in mind that the standard deviation is computed as

$$\sigma^2 = \frac{1}{12}(b-a)^2$$

If we do that for the above formula, we obtain

$$\sigma^{2} = \frac{1}{12} \left( \frac{2}{\sqrt{n_{in}}} \right)^{2} = \frac{1}{3} \frac{1}{n_{in}}$$

 $\Rightarrow$  Activations & gradients will be attenuated with each layer! (bad)



### **Glorot Initialization**

- Breakthrough results
  - In 2010, Xavier Glorot published an analysis of what went wrong in the initialization and derived a more general method for automatic initialization.
  - This new initialization massively improved results and made direct learning of deep networks possible overnight.
  - Let's look at his analysis in more detail...

X. Glorot, Y. Bengio, <u>Understanding the Difficulty of Training Deep</u> <u>Feedforward Neural Networks</u>, AISTATS 2010.



### Analysis

- Variance of neuron activations
  - > Suppose we have an input X with n components and a linear neuron with random weights W that spits out a number Y.
  - > What is the variance of Y?

$$Y = W_1 X_1 + W_2 X_2 + \dots + W_n X_n$$

If inputs and outputs have both mean 0, the variance is  $Var(W_iX_i) = E[X_i]^2 Var(W_i) + E[W_i]^2 Var(X_i) + Var(W_i) Var(X_i)$ 

 $= Var(W_i)Var(X_i)$ 

- > If the  $X_i$  and  $W_i$  are all i.i.d, then  $Var(Y) = Var(W_1X_1 + W_2X_2 + \dots + W_nX_n) = nVar(W_i)Var(X_i)$
- $\Rightarrow$  The variance of the output is the variance of the input, but scaled by  $n \ {\rm Var}(W_i).$



# Analysis (cont'd)

- Variance of neuron activations
  - > if we want the variance of the input and output of a unit to be the same, then  $n \operatorname{Var}(W_i)$  should be 1. This means

$$\operatorname{Var}(W_i) = rac{1}{n} = rac{1}{n_{ ext{in}}}$$

If we do the same for the backpropagated gradient, we get

$$\operatorname{Var}(W_i) = rac{1}{n_{ ext{out}}}$$

> As a compromise, Glorot & Bengio proposed to use

$$\mathrm{Var}(W) = rac{2}{n_\mathrm{in} + n_\mathrm{out}}$$

 $\Rightarrow$  Randomly sample the weights with this variance. That's it.



### Sidenote

- When sampling weights from a uniform distribution [a,b]
  - Again keep in mind that the standard deviation is computed as

$$\sigma^2 = \frac{1}{12}(b-a)^2$$

Glorot initialization with uniform distribution

$$W \sim U\left[-\frac{\sqrt{6}}{\sqrt{n_{in}+n_{out}}}, \frac{\sqrt{6}}{\sqrt{n_{in}+n_{out}}}\right]$$

Or when only taking into account the fan-in

$$W \sim U\left[-\frac{\sqrt{3}}{\sqrt{n_{in}}}, \frac{\sqrt{3}}{\sqrt{n_{in}}}\right]$$

If this had been implemented correctly in Torch from the beginning, the Deep Learning revolution might have happened a few years earlier...



# Extension to ReLU

- Important for learning deep models
  - Rectified Linear Units (ReLU)

$$g(a) = \max\left\{0, a\right\}$$

 Effect: gradient is propagated with a constant factor

$$\frac{\partial g(a)}{\partial a} = \begin{cases} 1, & a > 0\\ 0, & \text{else} \end{cases}$$



- We can also improve them with proper initialization
  - However, the Glorot derivation was based on tanh units, linearity assumption around zero does not hold for ReLU.
  - > He et al. made the derivations, derived to use instead

$$\operatorname{Var}(W) = rac{2}{n_{\mathrm{in}}}$$

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#### UNIVERSIT Batch Normalization [loffe & Szegedy '14]

- Motivation
  - > Optimization works best if all inputs of a layer are normalized.
- Idea
  - Introduce intermediate layer that centers the activations of the previous layer per minibatch.
  - I.e., perform transformations on all activations and undo those transformations when backpropagating gradients
  - Complication: centering + normalization also needs to be done at test time, but minibatches are no longer available at that point.
    - Learn the normalization parameters to compensate for the expected bias of the previous layer (usually a simple moving average)
  - Effect
    - Much improved convergence (but parameter values are important!)
    - Widely used in practice

### Dropout

#### **RNTHAACHEN** UNIVERSITY [Srivastava, Hinton '12]



#### Idea

- Randomly switch off units during training (a form of regularization).
- Change network architecture for each minibatch, effectively training many different variants of the network.
- When applying the trained network, multiply activations with the probability that the unit was set to zero during training.
- $\Rightarrow$  Greatly improved performance



### **References and Further Reading**

 More information on many practical tricks can be found in Chapter 1 of the book

> G. Montavon, G. B. Orr, K-R Mueller (Eds.) Neural Networks: Tricks of the Trade Springer, 1998, 2012



Yann LeCun, Leon Bottou, Genevieve B. Orr, Klaus-Robert Mueller Efficient BackProp, Ch.1 of the above book., 1998.



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