

Training Neural Networks: Optimization

Intro to Deep Learning, Spring 2019

Quick Recap

• Gradient descent, Backprop

Quick Recap: Training a network



- Define a total "loss" over all training instances
 - Quantifies the difference between desired output and the actual output, as a function of weights
- Find the weights that minimize the loss

Quick Recap: Training networks by gradient descent

$$L(W) = \frac{1}{N_X} \sum_X div(f(X; W), D(X))$$
$$\nabla_W L(W) = \frac{1}{N_X} \sum_X \nabla_W div(f(X; W), D(X))$$

Solved through gradient descent as $\widehat{W} = \arg\min_{W} L(W)$ \longrightarrow $W_k = W_{k-1} - \eta \nabla_W L(W)^T$

- The gradient of the total loss is the average of the gradients of the loss for the individual instances
- The total gradient can be plugged into gradient descent update to learn the network

Quick Recap: Training networks by gradient descent $L(W) = \frac{1}{N_X} \sum_{X} \begin{cases} \text{Computed using} \\ \text{backpropagation} \end{cases}$ $\nabla_W L(W) = \frac{1}{N_X} \sum_{W} \nabla_W div(f(X; W), D(X))$ Solved through gradient descen<u>t as</u>

- The gradient of the total loss is the average of the gradients of the loss for the individual instances
- The gradient can be plugged into gradient descent update to learn the network parameters

- x
- •
- •



• Forward pass: Compute output and all intermediate variables in the network, for the input *X*



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• Forward pass: Compute output and all intermediate variables in the network, for the input X11



• Forward pass: Compute output and all intermediate variables in the network, for the input *X*

The Forward Pass

- Set $y_0 = x$
- For layer k = 1 to N:
 - Recursion:

$$\mathbf{z}_{k} = \mathbf{W}_{k}\mathbf{y}_{k-1} + \mathbf{b}_{k}$$
$$\mathbf{y}_{k} = \mathbf{f}_{k}(\mathbf{z}_{k})$$

• Output:

$$\mathbf{Y}=\mathbf{y}_N$$

Quick Recap: Backprop. Forward pass



- Forward pass: Compute output and all intermediate variables in the network, for the input *X*
- Compute the divergence w.r.t. *desired* output

Quick Recap: Backpropagation



• Now work your way *backward* through the net to compute the derivative w.r.t each intermediate variable and each weight/bias

Backprop



First compute the gradient of the divergence w.r.t. Y. The actual gradient depends on the divergence function.

Backprop



 $\nabla_{\mathbf{z}_N} Div = \nabla_{\mathbf{Y}} Div J_{\mathbf{Y}}(\mathbf{z}_N)$

Chain rule (vector format; note order of multiplication)



 $\mathbf{z}_{N} = \mathbf{W}_{N}\mathbf{y}_{N-1} + \mathbf{b}_{N}$ $\mathbf{\nabla}_{\mathbf{W}_{N}}Div = \mathbf{y}_{N-1}\nabla_{\mathbf{z}_{N}}Div$ $\nabla_{\mathbf{b}_{N}}Div = \nabla_{\mathbf{z}_{N}}Div$

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Backprop $\mathbf{W}_{N-1}, \mathbf{b}_{N-1}$ **W**₁, **b**₁ d \mathbf{W}_N , \mathbf{b}_N Div(Y, d)<u>Softmax</u> Y Div \mathbf{z}_N Х \mathbf{Z}_1 $\mathbf{y_1}$ \mathbf{y}_{N-2} \mathbf{z}_{N-1} \mathbf{Z}_{N-2} $\nabla_{\mathbf{y}_{N-1}}Div$ $\nabla_{\mathbf{y}_{N-1}} Div = \nabla_{\mathbf{z}_N} Div \mathbf{W}_N$

Chain rule (vector format; note order of multiplication)



Chain rule (vector format; note order of multiplication)





$$\nabla_{\mathbf{y}_{N-2}} Div = \nabla_{\mathbf{z}_{N-1}} Div \mathbf{W}_{N-1}$$



 $\nabla_{\mathbf{z}_{N-2}} Div = \nabla_{\mathbf{y}_{N-2}} Div J_{\mathbf{y}_{N-2}}(\mathbf{z}_{N-2})$



 $\nabla_{\mathbf{z}_1} Div = \nabla_{\mathbf{y}_1} Div J_{\mathbf{y}_1}(\mathbf{z}_1)$



 $\nabla_{\mathbf{W}_{1}}Div = \mathbf{x}\nabla_{\mathbf{z}_{1}}Div$ $\nabla_{\mathbf{b}_{1}}Div = \nabla_{\mathbf{z}_{1}}Div$

In some problems we will also want to compute the derivative w.r.t. the input

The Backward Pass

- Set $\mathbf{y}_N = Y$, $\mathbf{y}_0 = \mathbf{x}$
- Initialize: Compute $\nabla_{\mathbf{y}_N} Div = \nabla_Y Div$
- For layer k = N downto 1:
 - Recursion:

$$\nabla_{\mathbf{z}_{k}} Div = \nabla_{\mathbf{y}_{k}} Div J_{\mathbf{y}_{k}}(\mathbf{z}_{k})$$
$$\nabla_{\mathbf{y}_{k-1}} Div = \nabla_{\mathbf{z}_{k}} Div \mathbf{W}_{k}$$

- Gradient computation:

$$\nabla_{\mathbf{W}_{k}} Div = \mathbf{y}_{k-1} \nabla_{\mathbf{z}_{k}} Div$$
$$\nabla_{\mathbf{b}_{k}} Div = \nabla_{\mathbf{z}_{k}} Div$$

Neural network training algorithm

- Initialize all weights and biases $(\mathbf{W}_1, \mathbf{b}_1, \mathbf{W}_2, \mathbf{b}_2, \dots, \mathbf{W}_N, \mathbf{b}_N)$
- Do:
 - Err = 0
 - For all k, initialize $\nabla_{\mathbf{W}_k} Err = 0$, $\nabla_{\mathbf{b}_k} Err = 0$
 - For all t = 1:T
 - Forward pass : Compute
 - Output $Y(X_t)$
 - Divergence $Div(Y_t, d_t)$
 - $Err += Div(Y_t, d_t)$
 - Backward pass: For all k compute:
 - $\nabla_{\mathbf{W}_k} Div(Y_t, d_t); \nabla_{\mathbf{b}_k} Div(Y_t, d_t)$
 - $\nabla_{\mathbf{W}_{k}} Err += \nabla_{\mathbf{W}_{k}} \mathbf{D}i\boldsymbol{\nu}(\boldsymbol{Y}_{t}, \boldsymbol{d}_{t}); \quad \nabla_{\mathbf{b}_{k}} Err += \nabla_{\mathbf{b}_{k}} \mathbf{D}i\boldsymbol{\nu}(\boldsymbol{Y}_{t}, \boldsymbol{d}_{t})$
 - For all *k*, update:

$$\mathbf{W}_{k} = \mathbf{W}_{k} - \frac{\eta}{T} \left(\nabla_{\mathbf{W}_{k}} Err \right)^{T}; \qquad \mathbf{b}_{k} = \mathbf{b}_{k} - \frac{\eta}{T} \left(\nabla_{\mathbf{W}_{k}} Err \right)^{T}$$

• Until *Err* has converged

Quick Recap

- Gradient descent, Backprop
- The issues with backprop and gradient descent
 - 1. Minimizes a *loss* which *relates* to classification accuracy, but is not actually classification accuracy
 - The divergence is a continuous valued proxy to classification error
 - Minimizing the loss is *expected* to, but not *guaranteed* to minimize classification error
 - 2. Simply minimizing the loss is hard enough...

Quick recap: Problem with gradient descent



 $\frac{W_k = W_{k-1} - \eta \nabla_w L(W)^T}{W_k = W_{k-1} - \eta \nabla_w L(W)^T}$ A step size that assures fast convergence for a given eccentricity can result in

divergence at a higher eccentricity

• .. Or result in extremely slow convergence at lower eccentricity

Quick recap: Problem with gradient descent



- The loss is a function of many weights (and biases)
 - Has different eccentricities w.r.t different weights
- A fixed step size for all weights in the network can result in the convergence of one weight, while causing a divergence of another

L(W)

Solutions for problem with gradient descent

- Try to normalize curvature in all directions
 - Second order methods, e.g. Newton's method
 - Too expensive: require inversion of a giant Hessian
- Treat each dimension independently:
 - Rprop, quickprop
 - Works, but ignores dependence between dimensions
 - Can result in unexpected behavior
 - Can still be too slow

Quick Recap

- Gradient descent, Backprop
- The issues with backprop and gradient descent
- Momentum methods..

A closer look at the convergence problem



• With dimension-independent learning rates, the solution will converge smoothly in some directions, but oscillate or diverge in others

A closer look at the convergence problem



- With dimension-independent learning rates, the solution will converge smoothly in some directions, but oscillate or diverge in others
- Proposal:
 - Keep track of oscillations
 - Emphasize steps in directions that converge smoothly
 - Shrink steps in directions that bounce around..

The momentum methods

- Maintain a running average of all past steps
 - In directions in which the convergence is smooth, the average will have a large value
 - In directions in which the estimate swings, the positive and negative swings will cancel out in the average
- Update with the running average, rather than the current gradient



Momentum Update



• The momentum method maintains a running average of all gradients until the *current* step

$$\Delta W^{(k)} = \beta \Delta W^{(k-1)} - \eta \nabla_W Err(W^{(k-1)})$$
$$W^{(k)} = W^{(k-1)} + \Delta W^{(k)}$$

- Typical β value is 0.9
- The running average steps
 - Get longer in directions where gradient stays in the same sign
 - Become shorter in directions where the sign keeps flipping
Training by gradient descent

- Initialize all weights $W_1, W_2, ..., W_K$
- Do:
 - For all *i*, *j*, *k*, initialize $\nabla_{W_k} Err = 0$
 - For all t = 1:T
 - For every layer k:
 - Compute $\nabla_{W_k} Div(Y_t, d_t)$
 - Compute $\nabla_{W_k} Err + = \frac{1}{T} \nabla_{W_k} Div(Y_t, d_t)$

- For every layer k:

 $W_k = W_k - \eta \nabla_{W_k} Err$

• Until *Err* has converged

Training with momentum

- Initialize all weights $W_1, W_2, ..., W_K$
- Do:
 - For all layers k, initialize $\nabla_{W_k} Err = 0$, $\Delta W_k = 0$
 - For all t = 1:T
 - For every layer k:
 - Compute gradient $\nabla_{W_k} Div(Y_t, d_t)$

 $-\nabla_{W_k} Err += \frac{1}{T} \nabla_{W_k} \mathbf{Div}(Y_t, d_t)$

- For every layer k

$$\Delta W_{k} = \beta \Delta W_{k} - \eta \nabla_{W_{k}} Err$$
$$W_{k} = W_{k} + \Delta W_{k}$$

Until *Err* has converged



- The momentum method $\Delta W^{(k)} = \beta \Delta W^{(k-1)} - \eta \nabla_W Err(W^{(k-1)})$
- At any iteration, to compute the current step:



The momentum method

$$\Delta W^{(k)} = \beta \Delta W^{(k-1)} - \eta \nabla_W Err(W^{(k-1)})$$

• At any iteration, to compute the current step:

- First computes the gradient step at the current location



• The momentum method

$$\Delta W^{(k)} = \beta \Delta W^{(k-1)} - \eta \nabla_W Err(W^{(k-1)})$$

- At any iteration, to compute the current step:
 - First computes the gradient step at the current location
 - Then adds in the scaled *previous* step
 - Which is actually a running average



• The momentum method

$$\Delta W^{(k)} = \beta \Delta W^{(k-1)} - \eta \nabla_W Err(W^{(k-1)})$$

- At any iteration, to compute the current step:
 - First computes the gradient step at the current location
 - Then adds in the scaled *previous* step
 - Which is actually a running average
 - To get the final step



- Takes a step along the past running average *after* walking along the gradient
- The procedure can be made more optimal by reversing the order of operations..



- Change the order of operations
- At any iteration, to compute the current step:



- Change the order of operations
- At any iteration, to compute the current step:

First extend the previous step



- Change the order of operations
- At any iteration, to compute the current step:
 - First extend the previous step
 - Then compute the gradient step at the resultant position



- Change the order of operations
- At any iteration, to compute the current step:
 - First extend the previous step
 - Then compute the gradient step at the resultant position
 - Add the two to obtain the final step



• Nestorov's method

$$\Delta W^{(k)} = \beta \Delta W^{(k-1)} - \eta \nabla_W Err \left(W^{(k-1)} + \beta \Delta W^{(k-1)} \right)$$
$$W^{(k)} = W^{(k-1)} + \Delta W^{(k)}$$



- Comparison with momentum (example from Hinton)
 - Blue: Momentum
 - Dotted line is the final update at each iteration
 - Brown/ochre/green: Nestorov
 - Brown is (scaled) previous update, ochre is gradient, green is the final update
- Converges much faster

Training with Nestorov

- Initialize all weights $W_1, W_2, ..., W_K$
- Do:
 - For all layers k, initialize $\nabla_{W_k} Err = 0$, $\Delta W_k = 0$
 - For every layer k

 $W_k = W_k + \beta \Delta W_k$

- For all t = 1:T
 - For every layer k:
 - Compute gradient $\nabla_{W_k} Div(Y_t, d_t)$

$$- \nabla_{W_k} Err += \frac{1}{T} \nabla_{W_k} \mathbf{Div}(Y_t, d_t)$$

– For every layer k

$$W_{k} = W_{k} - \eta \nabla_{W_{k}} Err$$
$$\Delta W_{k} = \beta \Delta W_{k} - \eta \nabla_{W_{k}} Err$$

• Until *Err* has converged

Momentum and trend-based methods..

• We will return to this topic again, very soon..

Story so far : Convergence

- Gradient descent can miss obvious answers
 - And this may be a good thing
- Vanilla gradient descent may be too slow or unstable due to the differences between the dimensions
- Second order methods can normalize the variation across dimensions, but are complex
- Adaptive or decaying learning rates can improve convergence
- Methods that decouple the dimensions can improve convergence
- Momentum methods which emphasize directions of steady improvement are demonstrably superior to other methods

Momentum methods: principle



- Ideally: Have component-specific step size
 - Too many independent parameters (maintain a step size for every weight/bias)
- Adaptive solution: Start with a common step size
 - Shrink step size in directions where the weight oscillates
 - Expand step size in directions where the weight moves consistently in one direction

Quick recap: Momentum methods

Momentum

Nestorov



- Momentum: Retain gradient value, but smooth out gradients by maintaining a running average
 - Cancels out steps in directions where the weight value oscillates
 - Adaptively increases step size in directions of consistent change

Recap

- Neural networks are universal approximators
- We must *train* them to approximate any function
- Networks are trained to minimize total "error" on a training set
 - We do so through empirical risk minimization
- We use variants of gradient descent to do so
 - Gradients are computed through backpropagation

Recap

- Vanilla gradient descent may be too slow or unstable
- Better convergence can be obtained through
 - Second order methods that normalize the variation across dimensions
 - Adaptive or decaying learning rates that can improve convergence
 - Methods like Rprop that decouple the dimensions can improve convergence
 - Momentum methods which emphasize directions of steady improvement and deemphasize unstable directions

Moving on: Topics for the day

- Incremental updates
- Revisiting "trend" algorithms
- Generalization
- Tricks of the trade
 - Divergences..
 - Activations
 - Normalizations

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- Incremental updates
- Revisiting "trend" algorithms
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 - Divergences..
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 - Normalizations



• Given input output pairs at a number of locations, estimate the entire function



• Start with an initial function



- Start with an initial function
- Adjust its value at *all* points to make the outputs closer to the required value
 - Gradient descent adjusts parameters to adjust the function value at *all* points
 - Repeat this iteratively until we get arbitrarily close to the target function at the training points



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Effect of number of samples



- Problem with conventional gradient descent: we try to simultaneously adjust the function at *all* training points
 - We must process *all* training points before making a single adjustment
 - "Batch" update



- Alternative: adjust the function at one training point at a time
 - Keep adjustments small



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- Alternative: adjust the function at one training point at a time
 - Keep adjustments small
 - Eventually, when we have processed all the training points, we will have adjusted the entire function
 - With *greater* overall adjustment than we would if we made a single "Batch" update

Incremental Update: Stochastic Gradient Descent

- Given $(X_1, d_1), (X_2, d_2), ..., (X_T, d_T)$
- Initialize all weights W_1, W_2, \ldots, W_K
- Do:
 - For all t = 1:T
 - For every layer k:
 - Compute $\nabla_{W_k} Div(Y_t, d_t)$
 - Update

 $W_k = W_k - \eta \nabla_{W_k} \mathbf{Div}(\mathbf{Y}_t, \mathbf{d}_t)$

Until *Err* has converged










- If we loop through the samples in the same order, we may get cyclic behavior
- We must go through them *randomly* to get more convergent behavior



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Story so far

- In any gradient descent optimization problem, presenting training instances incrementally can be more effective than presenting them all at once
 - Provided training instances are provided in random order
 - "Stochastic Gradient Descent"
- This also holds for training neural networks

Explanations and restrictions

- So why does this process of incremental updates work?
- Under what conditions?
- For "why": first consider a simplistic explanation that's often given
 - Look at an extreme example

The expected behavior of the gradient



- The individual training instances contribute different directions to the overall gradient
 - The final gradient points is the average of individual gradients
 - It points towards the *net* direction



• Extreme instance of data clotting: all the training instances are exactly the same

The expected behavior of the gradient



- The individual training instance contribute identical directions to the overall gradient
 - The final gradient points is simply the gradient for an individual instance



- Batch gradient descent operates over T training instances to get a *single* update
- SGD gets T updates for the same computation



• Also holds if all the data are not identical, but are tightly clumped together

Clumpy data..



• As data get increasingly diverse, the benefits of incremental updates decrease, but do not entirely vanish

When does it work

- What are the considerations?
- And how well does it work?

Incremental Update: Stochastic Gradient Descent

- Given $(X_1, d_1), (X_2, d_2), ..., (X_T, d_T)$
- Initialize all weights $W_1, W_2, \dots, W_K; j = 0$
- Do:
 - Randomly permute $(X_1, d_1), (X_2, d_2), ..., (X_T, d_T)$
 - For all t = 1:T
 - j = j + 1
 - For every layer k:
 - Compute $\nabla_{W_k} Div(Y_t, d_t)$
 - Update

 $W_k = W_k - \eta_j \nabla_{W_k} \mathbf{Div}(\mathbf{Y_t}, \mathbf{d_t})$

• Until *Err* has converged



- Except in the case of a perfect fit, even an optimal overall fit will look incorrect to *individual* instances
 - Correcting the function for individual instances will lead to never-ending, non-convergent updates
 - We must *shrink* the learning rate with iterations to prevent this
 - Correction for individual instances with the eventual miniscule learning rates will not modify the function

Incremental Update: Stochastic Gradient Descent

- Given $(X_1, d_1), (X_2, d_2), ..., (X_T, d_T)$
- Initialize all weights $W_1, W_2, \dots, W_K; j = 0$
- Do:



• Until *Err* has converged

Stochastic Gradient Descent

- The iterations can make multiple passes over the data
- A single pass through the entire training data is called an "epoch"
 - An epoch over a training set with T samples results in T updates of parameters

SGD convergence

- SGD converges "almost surely" to a global or local minimum for most functions
 - Sufficient condition: step sizes follow the following conditions

$$\sum_k \eta_k = \infty$$

• Eventually the entire parameter space can be searched

$$\sum_k \eta_k^2 < \infty$$

- The steps shrink
- The fastest converging series that satisfies both above requirements is

$$\eta_k \propto \frac{1}{k}$$

- This is the optimal rate of shrinking the step size for strongly convex functions
- More generally, the learning rates are heuristically determined
- If the loss is convex, SGD converges to the optimal solution
- For non-convex losses SGD converges to a local minimum

SGD convergence

- We will define convergence in terms of the number of iterations taken to get within ϵ of the optimal solution
 - $\left| f \left(W^{(k)} \right) f (W^*) \right| < \epsilon$
 - Note: f(W) here is the error on the *entire* training data, although SGD itself updates after every training instance
- Using the optimal learning rate 1/k, for strongly convex functions,

$$|W^{(k)} - W^*| < \frac{1}{k} |W^{(0)} - W^*|$$

- Strongly convex \rightarrow Can be placed inside a quadratic bowl, touching at any point
- Giving us the iterations to ϵ convergence as $O\left(\frac{1}{\epsilon}\right)$
- For generically convex (but not strongly convex) function, various proofs report an ϵ convergence of $\frac{1}{\sqrt{k}}$ using a learning rate of $\frac{1}{\sqrt{k}}$.

Batch gradient convergence

In contrast, using the batch update method, for strongly convex functions,

$$|W^{(k)} - W^*| < c^k |W^{(0)} - W^*|$$

– Giving us the iterations to ϵ convergence as $O\left(log\left(\frac{1}{\epsilon}\right)\right)$

- For generic convex functions, iterations to ϵ convergence is $O\left(\frac{1}{\epsilon}\right)$
- Batch gradients converge "faster"
 - But SGD performs T updates for every batch update

SGD Convergence: Loss value

If:

- f is λ -strongly convex, and
- at step t we have a noisy estimate of the subgradient \hat{g}_t with $\mathbb{E}[\|\hat{g}_t\|^2] \leq G^2$ for all t,
- and we use step size $\eta_t = 1/\lambda_t$

Then for any T > 1:

$$\mathbb{E}[f(w_T) - f(w^*)] \le \frac{17G^2(1 + \log(T))}{\lambda T}$$

SGD Convergence

- We can bound the expected difference between the loss over our data using the optimal weights w^* and the weights w_T at any single iteration to $\mathcal{O}\left(\frac{\log(T)}{T}\right)$ for strongly convex loss or $\mathcal{O}\left(\frac{\log(T)}{\sqrt{T}}\right)$ for convex loss
- Averaging schemes can improve the bound to $\mathcal{O}\left(\frac{1}{T}\right)$ and $\mathcal{O}\left(\frac{1}{\sqrt{T}}\right)$
- Smoothness of the loss is not required

SGD Convergence and weight averaging

Polynomial Decay Averaging:

$$\overline{w}_t^{\gamma} = \left(1 - \frac{\gamma + 1}{t + \gamma}\right) \overline{w}_{t-1}^{\gamma} + \frac{\gamma + 1}{t + \gamma} w_t$$

With γ some small positive constant, e.g. $\gamma = 3$ Achieves $\mathcal{O}\left(\frac{1}{T}\right)$ (strongly convex) and $\mathcal{O}\left(\frac{1}{\sqrt{T}}\right)$ (convex) convergence

SGD example



- A simpler problem: K-means
- Note: SGD converges slower
- Also note the rather large variation between runs
 - Lets try to understand these results..

Recall: Modelling a function





To learn a network f(X; W) to model a function g(X) we minimize the *expected divergence*

$$\widehat{\boldsymbol{W}} = \underset{W}{\operatorname{argmin}} \int_{X} div(f(X; W), g(X))P(X)dX$$
$$= \underset{W}{\operatorname{argmin}} E\left[div(f(X; W), g(X))\right]$$

Recall: The *Empirical* **risk**





• In practice, we minimize the *empirical error*

$$Err(f(X;W),g(X)) = \frac{1}{N} \sum_{i=1}^{N} div(f(X_i;W),d_i)$$
$$\widehat{W} = \underset{W}{\operatorname{argmin}} Err(f(X;W),g(X))$$

• The expected value of the empirical error is actually the expected divergence $E\left[Err(f(X;W),g(X))\right] = E\left[div(f(X;W),g(X))\right]$

Recap: The Empirical risk





• In practice, we minimize the *empirical error*

$$Err(f(X;W),g(X)) = \frac{1}{N} \sum_{i=1}^{N} div(f(X_i;W),d_i)$$

The empirical error is an unbiased estimate of the expected error Though there is no guarantee that minimizing it will minimize the expected error

value of the empirical error is actually the expected erro

 $E\left[Err(f(X;W),g(X))\right] = E\left[div(f(X;W),g(X))\right]$



$$Err(f(X;W),g(X)) = \frac{1}{N}\sum_{i=1}^{N}div(f(X_i;W),d_i)$$

The empirical error is an unbiased estimate of the expected error Though there is no guarantee that minimizing it will minimize the expected error

E[Err(f(X;W),g(X))] = E[div(f(X;W),g(X))]



- At each iteration, SGD focuses on the divergence of a single sample div(f(X_i; W), d_i)
- The expected value of the sample error is **still** the expected divergence E[div(f(X; W), g(X))] 106



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- The blue curve is the function being approximated
- The red curve is the approximation by the model at a given W
- The heights of the shaded regions represent the point-by-point error
 - The divergence is a function of the error
 - We want to find the W that minimizes the average divergence



 Sample estimate approximates the shaded area with the average length of the lines



- Sample estimate approximates the shaded area with the average length of the lines
- This average length will change with position of the samples



- Sample estimate approximates the shaded area with the average length of the lines
- This average length will change with position of the samples



- Having more samples makes the estimate more robust to changes in the position of samples
 - The variance of the estimate is smaller



- Having very few samples makes the estimate swing wildly with the sample position
 - Since our estimator learns the W to minimize this estimate, the learned W too can swing wildly



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SGD example



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SGD vs batch

- SGD uses the gradient from only one sample at a time, and is consequently high variance
- But also provides significantly quicker updates than batch
- Is there a good medium?

Alternative: Mini-batch update



- Alternative: adjust the function at a small, randomly chosen subset of points
 - Keep adjustments small
 - If the subsets cover the training set, we will have adjusted the entire function
- As before, vary the subsets randomly in different passes through the training data

Incremental Update: Mini-batch update

- Given $(X_1, d_1), (X_2, d_2), ..., (X_T, d_T)$
- Initialize all weights $W_1, W_2, \dots, W_K; j = 0$
- Do:
 - Randomly permute $(X_1, d_1), (X_2, d_2), ..., (X_T, d_T)$
 - For t = 1: b: T
 - j = j + 1
 - For every layer k:
 - $-\Delta W_k = 0$
 - For t' = t : t+b-1
 - For every layer k:
 - » Compute $\nabla_{W_k} Div(Y_t, d_t)$
 - » $\Delta W_k = \Delta W_k + \frac{1}{b} \nabla_{W_k} Div(Y_t, d_t)$
 - Update
 - For every layer k:

$$W_k = W_k - \eta_j \Delta W_k$$

• Until *Err* has converged

Incremental Update: Mini-batch update

- Given $(X_1, d_1), (X_2, d_2), ..., (X_T, d_T)$
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 - Update
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$$W_k = W_k - \eta_j \Delta W_k$$

• Until *Err* has converged

Mini-batch size

Shrinking step size

Mini Batches



• Mini-batch updates compute and minimize a *batch error*

$$BatchErr(f(X;W),g(X)) = \frac{1}{b}\sum_{i=1}^{b} div(f(X_i;W),d_i)$$

• The expected value of the batch error is also the expected divergence E[BatchErr(f(X; W), g(X))] = E[div(f(X; W), g(X))]



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• The expected value of the batch error is also the expected divergence $E\left[BatchErr(f(X; W), g(X))\right] = E\left[div(f(X; W), g(X))\right]$

Minibatch convergence

- For convex functions, convergence rate for SGD is $\mathcal{O}\left(\frac{1}{\sqrt{k}}\right)$.
- For *mini-batch* updates with batches of size *b*, the convergence rate is $O\left(\frac{1}{\sqrt{bk}} + \frac{1}{k}\right)$
 - Apparently an improvement of \sqrt{b} over SGD
 - But since the batch size is b, we perform b times as many computations per iteration as SGD
 - We actually get a *degradation* of \sqrt{b}
- However, in practice
 - The objectives are generally not convex; mini-batches are more effective with the right learning rates
 - We also get additional benefits of vector processing

SGD example



• Mini-batch performs comparably to batch training on this simple problem

But converges orders of magnitude faster

Measuring Error

 Convergence is generally defined in terms of the *overall training* error

Not sample or batch error



- Infeasible to actually measure the overall training error after each iteration
- More typically, we estimate is as
 - Divergence or classification error on a held-out set
 - Average sample/batch error over the past N samples/batches

Training and minibatches

- In practice, training is usually performed using minibatches
 - The mini-batch size is a hyper parameter to be optimized
- Convergence depends on learning rate
 - Simple technique: fix learning rate until the error plateaus, then reduce learning rate by a fixed factor (e.g. 10)
 - Advanced methods: Adaptive updates, where the learning rate is itself determined as part of the estimation



- SGD: Presenting training instances one-at-a-time can be more effective than full-batch training
 - Provided they are provided in random order
- For SGD to converge, the learning rate must shrink sufficiently rapidly with iterations
 - Otherwise the learning will continuously "chase" the latest sample
- SGD estimates have higher variance than batch estimates
- Minibatch updates operate on *batches* of instances at a time
 - Estimates have lower variance than SGD
 - Convergence rate is theoretically worse than SGD
 - But we compensate by being able to perform batch processing

Training and minibatches

- Convergence depends on learning rate
 - Simple technique: fix learning rate until the error plateaus, then reduce learning rate by a fixed factor (e.g. 10)
 - Advanced methods: Adaptive updates, where the learning rate is itself determined as part of the estimation

Moving on: Topics for the day

- Incremental updates
- Revisiting "trend" algorithms
- Generalization
- Tricks of the trade
 - Divergences..
 - Activations
 - Normalizations

Recall: Momentum



• The momentum method

$$\Delta W^{(k)} = \beta \Delta W^{(k-1)} - \eta \nabla_W Err(W^{(k-1)})$$

• Updates using a running average of the gradient

Momentum and incremental updates



- The momentum method $\Delta W^{(k)} = \beta \Delta W^{(k-1)} - \eta \nabla_W Err(W^{(k-1)})$
- Incremental SGD and mini-batch gradients tend to have high variance
- Momentum smooths out the variations
 - Smoother and faster convergence

Incremental Update: Mini-batch update

- Given $(X_1, d_1), (X_2, d_2), ..., (X_T, d_T)$
- Initialize all weights W_1, W_2, \dots, W_K ; $j = 0, \Delta W_k = 0$
- Do:
 - Randomly permute $(X_1, d_1), (X_2, d_2), ..., (X_T, d_T)$
 - For t = 1:b:T
 - j = j + 1
 - For every layer k:

$$- \nabla_{W_k} Err = 0$$

- For t' = t : t+b-1
 - For every layer k:
 - » Compute $\nabla_{W_k} Div(Y_t, d_t)$

»
$$\nabla_{W_k} Err + = \frac{1}{b} \nabla_{W_k} Div(Y_t, d_t)$$

- Update
 - For every layer k:

$$\Delta W_k = \beta \Delta W_k - \eta_j \nabla_{W_k} Err$$
$$W_k = W_k + \Delta W_k$$

• Until *Err* has converged

Nestorov's Accelerated Gradient



- At any iteration, to compute the current step:
 - First extend the previous step
 - Then compute the gradient at the resultant position
 - Add the two to obtain the final step
- This also applies directly to incremental update methods
 - The accelerated gradient smooths out the variance in the gradients

Nestorov's Accelerated Gradient



• Nestorov's method $\Delta W^{(k)} = \beta \Delta W^{(k-1)} - \eta \nabla_W Err(W^{(k-1)} + \beta \Delta W^{(k-1)})$ $W^{(k)} = W^{(k-1)} + \Delta W^{(k)}$

Incremental Update: Mini-batch update

- Given $(X_1, d_1), (X_2, d_2), ..., (X_T, d_T)$
- Initialize all weights $W_1, W_2, ..., W_K; j = 0, \Delta W_k = 0$
- Do:
 - Randomly permute $(X_1, d_1), (X_2, d_2), ..., (X_T, d_T)$
 - For t = 1: b: T
 - j = j + 1
 - For every layer k:
 - $W_k = W_k + \beta \Delta W_k$
 - $\nabla_{W_k} Err = 0$
 - For t' = t : t+b-1
 - For every layer k:
 - » Compute $\nabla_{W_k} Div(Y_t, d_t)$
 - » $\nabla_{W_k} Err + = \frac{1}{h} \nabla_{W_k} \mathbf{Div}(Y_t, d_t)$
 - Update
 - For every layer k:

$$W_{k} = W_{k} - \eta_{j} \nabla_{W_{k}} Err$$
$$\Delta W_{k} = \beta \Delta W_{k} - \eta_{j} \nabla_{W_{k}} Err$$

• Until *Err* has converged

More recent methods

- Several newer methods have been proposed that follow the general pattern of enhancing longterm trends to smooth out the variations of the mini-batch gradient
 - RMS Prop
 - Adagrad
 - AdaDelta

- ...

- ADAM: very popular in practice
- All roughly equivalent in performance

Smoothing the trajectory

	Step	X component	Y component
	1	1	+2.5
	2	1	-3
$\left(\left(\frac{1}{\sqrt{3}} \right) \right)$	3	3	+2.5
	4	1	-2
	5	2	1.5

- Simple gradient and acceleration methods still demonstrate oscillatory behavior in some directions
- Observation: Steps in "oscillatory" directions show large total movement
 - In the example, total motion in the vertical direction is much greater than in the horizontal direction
- Improvement: Dampen step size in directions with high motion
 - Second order term

Variance-normalized step



- In recent past
 - Total movement in Y component of updates is high
 - Movement in X components is lower
- Current update, modify usual gradient-based update:
 - Scale *down* Y component
 - Scale *up* X component
 - According to their variation (and not just their average)
- A variety of algorithms have been proposed on this premise
 - We will see a popular example

RMS Prop

- Notation:
 - Updates are by parameter
 - Sum derivative of divergence w.r.t any individual parameter w is shown as $\partial_w D$
 - The **squared** derivative is $\partial_w^2 D = (\partial_w D)^2$
 - Short-hand notation represents the squared derivative, not the second derivative
 - The *mean squared* derivative is a running estimate of the average squared derivative. We will show this as $E[\partial_w^2 D]$
- Modified update rule: We want to
 - scale down updates with large mean squared derivatives
 - scale up updates with small mean squared derivatives

RMS Prop

- This is a variant on the *basic* mini-batch SGD algorithm
- Procedure:
 - Maintain a running estimate of the mean squared value of derivatives for each parameter
 - Scale update of the parameter by the *inverse* of the *root mean* squared derivative

$$E[\partial_w^2 D]_k = \gamma E[\partial_w^2 D]_{k-1} + (1-\gamma)(\partial_w^2 D)_k$$
$$w_{k+1} = w_k - \frac{\eta}{\sqrt{E[\partial_w^2 D]_k + \epsilon}} \partial_w D$$

RMS Prop

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$$w_{k+1} = w_k - \frac{\eta}{\sqrt{E[\partial_w^2 D]_k + \epsilon}} \partial_w D$$

Note similarity to RPROP The magnitude of the derivative is being normalized out

RMS Prop (updates are for each weight of each layer)

- Do:
 - Randomly shuffle inputs to change their order
 - Initialize: k = 1; for all weights w in all layers, $E[\partial_w^2 D]_k = 0$
 - For all t = 1: B: T (incrementing in blocks of B inputs)
 - For all weights in all layers initialize $(\partial_w D)_k = 0$
 - For b = 0: B 1
 - Compute
 - » Output $Y(X_{t+b})$
 - » Compute gradient $\frac{dDiv(Y(X_{t+b}), d_{t+b})}{dw}$

» Compute
$$(\partial_w D)_k + = \frac{1}{B} \frac{dDiv(Y(X_{t+b}), d_{t+b})}{dw}$$

$$E[\partial_w^2 D]_k = \gamma E[\partial_w^2 D]_{k-1} + (1-\gamma)(\partial_w^2 D)$$
$$w_{k+1} = w_k - \frac{\eta}{\sqrt{E[\partial_w^2 D]_k + \epsilon}} \partial_w D$$

• k = k + 1

• Until $E(W^{(1)}, W^{(2)}, ..., W^{(K)})$ has converged
ADAM: RMSprop with momentum

- RMS prop only considers a second-moment normalized version of the current gradient
- ADAM utilizes a smoothed version of the *momentum-augmented* gradient
- Procedure:
 - Maintain a running estimate of the mean derivative for each parameter
 - Maintain a running estimate of the mean squared value of derivatives for each parameter
 - Scale update of the parameter by the *inverse* of the *root mean squared* derivative

$$m_{k} = \delta m_{k-1} + (1 - \delta)(\partial_{w}D)_{k}$$

$$v_{k} = \gamma v_{k-1} + (1 - \gamma)(\partial_{w}^{2}D)_{k}$$

$$\widehat{m}_{k} = \frac{m_{k}}{1 - \delta^{k}}, \qquad \widehat{v}_{k} = \frac{v_{k}}{1 - \gamma^{k}}$$

$$w_{k+1} = w_{k} - \frac{\eta}{\sqrt{\widehat{v}_{k} + \epsilon}} \widehat{m}_{k}$$

ADAM: RMSprop with momentum

- RMS prop only considers a second-moment normalized version of the current gradient
- ADAM utilizes a smoothed version of the *momentum-augmented* gradient
- Procedure:
 - Maintain a running estimate of the mean derivative for each parameter
 - Maintain a running estimate of the mean squared value parameter
 - Scale update of the parameter by the *inverse* of the derivative

$$m_k = \delta m_{k-1} + (1 - \delta) (\partial_w D)_k$$

Ensures that the δ and γ terms do not dominate in early iterations

$$v_k = \gamma v_{k-1} + (1 - \gamma) (\partial_w^2 D)_k$$
$$\widehat{m}_k = \frac{m_k}{1 - \delta^k}, \qquad \widehat{v}_k = \frac{v_k}{1 - \gamma^k}$$

 $w_{k+1} = w_k - \frac{\eta}{\sqrt{\hat{v}_k + \epsilon}} \hat{m}_k$

Other variants of the same theme

- Many:
 - Adagrad
 - AdaDelta
 - ADAM
 - AdaMax
 - ...
- Generally no explicit learning rate to optimize
 - But come with other hyper parameters to be optimized
 - Typical params:
 - RMSProp: $\eta = 0.001, \gamma = 0.9$
 - ADAM: $\eta = 0.001, \delta = 0.9, \gamma = 0.999$

Visualizing the optimizers: Beale's Function



• http://www.denizyuret.com/2015/03/alec-radfords-animations-for.html

Visualizing the optimizers: Long Valley



http://www.denizyuret.com/2015/03/alec-radfords-animations-for.html

Visualizing the optimizers: Saddle Point



http://www.denizyuret.com/2015/03/alec-radfords-animations-for.html

Story so far

- Gradient descent can be sped up by incremental updates
 - Convergence is guaranteed under most conditions
 - Learning rate must shrink with time for convergence
 - Stochastic gradient descent: update after each observation. Can be much faster than batch learning
 - Mini-batch updates: update after batches. Can be more efficient than SGD
- Convergence can be improved using smoothed updates
 - RMSprop and more advanced techniques

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