

### **Machine Learning – Lecture 11**

### **Neural Networks II**

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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**

#### • Learning Multi-layer Networks

- Backpropagation
- Computational graphs
- Automatic differentiation
- Practical issues
- Gradient Descent
  - Stochastic Gradient Descent & Minibatches
  - Choosing Learning Rates
  - Momentum
  - > RMS Prop
  - > Other Optimizers
- Tricks of the Trade
  - Shuffling

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- Data Augmentation
- Normalization



### **Recap: Learning with Hidden Units**

- How can we train multi-layer networks efficiently?
  - > Need an efficient way of adapting all weights, not just the last layer.

- Idea: Gradient Descent
  - Set up an error function

$$E(\mathbf{W}) = \sum_{n} L(t_n, y(\mathbf{x}_n; \mathbf{W})) + \lambda \Omega(\mathbf{W})$$

with a loss  $L(\cdot)$  and a regularizer  $\Omega(\cdot)$ .

$$\text{E.g., } L(t, y(\mathbf{x}; \mathbf{W})) = \sum_{n} (y(\mathbf{x}_{n}; \mathbf{W}) - t_{n})^{2} \qquad \text{L}_{2} \text{ loss}$$

$$\Omega(\mathbf{W}) = ||\mathbf{W}||_{F}^{2} \qquad \text{L}_{2} \text{ regularizer}$$
("weight decay")
$$\partial E(\mathbf{W})$$

 $\Rightarrow$  Update each weight  $W_{ij}^{(k)}$  in the direction of the gradient  $\frac{\partial E(k)}{\partial W_{ij}}$ 



### **Gradient Descent**

- Two main steps
  - 1. Computing the gradients for each weight
  - 2. Adjusting the weights in the direction of the gradient



### **Obtaining the Gradients**

Approach 1: Naive Analytical Differentiation



$\frac{\partial E(\mathbf{W})}{\partial W_{10}^{(2)}}$	•••	$\frac{\partial E(\mathbf{W})}{\partial W_{kh}^{(2)}}$
$\frac{\partial E(\mathbf{W})}{\partial W_{10}^{(1)}}$	•••	$\frac{\partial E(\mathbf{W})}{\partial W_{hd}^{(1)}}$

- > Compute the gradients for each variable analytically.
- What is the problem when doing this?

#### **RWTHAACHEN** UNIVERSITY Excursion: Chain Rule of Differentiation

One-dimensional case: Scalar functions



$$\Delta z = \frac{\mathrm{d}z}{\mathrm{d}y} \Delta y$$
$$\Delta y = \frac{\mathrm{d}y}{\mathrm{d}x} \Delta x$$
$$\Delta z = \frac{\mathrm{d}z}{\mathrm{d}y} \frac{\mathrm{d}y}{\mathrm{d}x} \Delta x$$
$$\frac{\mathrm{d}z}{\mathrm{d}x} = \frac{\mathrm{d}z}{\mathrm{d}y} \frac{\mathrm{d}y}{\mathrm{d}x}$$

# Excursion: Chain Rule of Differentiation

• Multi-dimensional case: Total derivative



$$\frac{\partial z}{\partial x} = \frac{\partial z}{\partial y_1} \frac{\partial y_1}{\partial x} + \frac{\partial z}{\partial y_2} \frac{\partial y_2}{\partial x} + \dots$$
$$= \sum_{i=1}^k \frac{\partial z}{\partial y_i} \frac{\partial y_i}{\partial x}$$

 $\Rightarrow$  Need to sum over all paths that lead to the target variable x.



### **Obtaining the Gradients**

Approach 1: Naive Analytical Differentiation



- Compute the gradients for each variable analytically.
  - > What is the problem when doing this?
  - $\Rightarrow$  With increasing depth, there will be exponentially many paths!
- $\Rightarrow$  Infeasible to compute this way.



### **Obtaining the Gradients**

Approach 2: Numerical Differentiation



- > Given the current state  $\mathbf{W}^{(\tau)}$ , we can evaluate  $E(\mathbf{W}^{(\tau)})$ .
- > Idea: Make small changes to  $\mathbf{W}^{(\tau)}$  and accept those that improve  $E(\mathbf{W}^{(\tau)})$ .
- $\Rightarrow$  Horribly inefficient! Need several forward passes for each weight. Each forward pass is one run over the entire dataset!

### **Obtaining the Gradients**

Approach 3: Incremental Analytical Differentiation



- > Idea: Compute the gradients layer by layer.
- > Each layer below builds upon the results of the layer above.
- $\Rightarrow$  The gradient is propagated backwards through the layers.
- $\Rightarrow$  Backpropagation algorithm



- Core steps
  - Convert the discrepancy between each output and its target value into an error derivate.
  - 2. Compute error derivatives in each hidden layer from error derivatives in the layer above.
  - 3. Use error derivatives *w.r.t.* activities to get error derivatives *w.r.t.* the incoming weights



### **Backpropagation Algorithm**



Input of layer k

 $\frac{\partial E}{\partial z_i^{(k)}} = \frac{\partial y_j^{(k)}}{\partial z_i^{(k)}} \frac{\partial E}{\partial y_i^{(k)}} \qquad = \frac{\partial g\left(z_j^{(k)}\right)}{\partial z_i^{(k)}} \frac{\partial E}{\partial y_i^{(k)}}$ 



Slide adapted from Geoff Hinton

 $\succ z_i^{(k)}$ 

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Connections:  $z_{j}^{(k)} = \sum_{i} w_{ji}^{(k-1)} y_{i}^{(k-1)}$  $y_{j}^{(k)} = g(z_{j}^{(k)})$ 16







- Efficient propagation scheme
  - >  $y_i^{(k-1)}$  is already known from forward pass! (Dynamic Programming)
- $\Rightarrow$  Propagate back the gradient from layer k and multiply with  $y_i^{(k-1)}$ .



### Summary: MLP Backpropagation

• Forward Pass

$$\mathbf{y}^{(0)} = \mathbf{x}$$
  
for  $k = 1, ..., l$  do  
$$\mathbf{z}^{(k)} = \mathbf{W}^{(k)} \mathbf{y}^{(k-1)}$$
  
$$\mathbf{y}^{(k)} = g_k(\mathbf{z}^{(k)})$$

endfor

$$\mathbf{y} = \mathbf{y}^{(l)}$$

$$E = L(\mathbf{t}, \mathbf{y}) + \lambda \Omega(\mathbf{W})$$

#### Backward Pass

 $\mathbf{h} \leftarrow \frac{\partial E}{\partial \mathbf{y}} = \frac{\partial}{\partial \mathbf{y}} L(\mathbf{t}, \mathbf{y}) + \lambda \frac{\partial}{\partial \mathbf{y}} \Omega$ for k = l, l-1, ..., 1 do  $\mathbf{h} \leftarrow \frac{\partial E}{\partial \mathbf{z}^{(k)}} = \mathbf{h} \odot g'(\mathbf{y}^{(k)})$  $\frac{\partial E}{\partial \mathbf{W}^{(k)}} = \mathbf{h} \mathbf{y}^{(k-1)\top} + \lambda \frac{\partial \Omega}{\partial \mathbf{W}^{(k)}}$  $\mathbf{h} \leftarrow \frac{\partial E}{\partial \mathbf{y}^{(k-1)}} = \mathbf{W}^{(k)\top} \mathbf{h}$ endfor

- $\,\,$  For efficiency, an entire batch of data  ${\bf X}$  is processed at once.
- ➤ ⊙ denotes the element-wise product



### Analysis: Backpropagation

- Backpropagation is the key to make deep NNs tractable
   However...
- The Backprop algorithm given here is specific to MLPs
  - It does not work with more complex architectures, e.g. skip connections or recurrent networks!
  - Whenever a new connection function induces a different functional form of the chain rule, you have to derive a new Backprop algorithm for it.



- $\Rightarrow$  Tedious...
- Let's analyze Backprop in more detail
  - This will lead us to a more flexible algorithm formulation



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  - Stochastic Gradient Descent & Minibatches
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  - Momentum
  - > RMS Prop
  - > Other Optimizers
- Tricks of the Trade
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  - Data Augmentation
  - Normalization

### **Computational Graphs**

- We can think of mathematical expressions as graphs
  - E.g., consider the expression

$$e = (a+b)*(b+1)$$

 We can decompose this into the operations



e = c \* d



and visualize this as a computational graph.

- Evaluating partial derivatives  $\frac{\partial Y}{\partial X}$  in such a graph
  - General rule: sum over all possible paths from Y to X and multiply the derivatives on each edge of the path together.

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### Factoring Paths

- Problem: Combinatorial explosion
  - > Example:



- > There are 3 paths from X to Y and 3 more from Y to Z.
- If we want to compute  $\frac{\partial Z}{\partial X}$ , we need to sum over  $3 \times 3$  paths:  $\frac{\partial Z}{\partial X} = \alpha \delta + \alpha \epsilon + \alpha \zeta + \beta \delta + \beta \epsilon + \beta \zeta + \gamma \delta + \gamma \epsilon + \gamma \zeta$
- > Instead of naively summing over paths, it's better to factor them

$$\frac{\partial Z}{\partial X} = (\alpha + \beta + \gamma) * (\delta + \epsilon + \zeta)$$

Slide inspired by Christopher Olah

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### **Efficient Factored Algorithms**



- Efficient algorithms for computing the sum
  - Instead of summing over all of the paths explicitly, compute the sum more efficiently by merging paths back together at every node.

Slide inspired by Christopher Olah



### Why Do We Care?

- Let's consider the example again
  - Using forward-mode differentiation from b up...
  - > Runtime:  $\mathcal{O}(\text{#edges})$
  - Result: derivative of every node with respect to b.



 $\partial a$ 



### Why Do We Care?

- Let's consider the example again
  - Using reverse-mode differentiation from e down...
  - > Runtime:  $\mathcal{O}(\text{#edges})$
  - Result: derivative of *e* with respect to every node.



- $\Rightarrow$  This is what we want to compute in Backpropagation!
- Forward differentiation needs one pass per node. With backward differentiation we can compute all derivatives in one single pass.
- $\Rightarrow$  Speed-up in  $\mathcal{O}($ #inputs) compared to forward differentiation!

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### **Obtaining the Gradients**

Approach 4: Automatic Differentiation



- > 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



### **Modular Implementation**

- Solution in many current Deep Learning libraries
  - Provide a limited form of automatic differentiation
  - Restricted to "programs" composed of "modules" with a predefined set of operations.
- Each module is defined by two main functions
  - 1. Computing the outputs  ${\bf y}$  of the module given its inputs  ${\bf x}$

 $\mathbf{y} = \text{module.fprop}(\mathbf{x})$ 

where  $\mathbf{x}, \mathbf{y}$ , and intermediate results are stored in the module.

2. Computing the gradient  $\partial E/\partial \mathbf{x}$  of a scalar cost w.r.t. the inputs  $\mathbf{x}$  given the gradient  $\partial E/\partial \mathbf{y}$  w.r.t. the outputs  $\mathbf{y}$ 

$$\frac{\partial E}{\partial \mathbf{x}} = \text{module}.\mathbf{bprop}(\frac{\partial E}{\partial \mathbf{y}})$$

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### Implementing Softmax Correctly

- Softmax output
  - De-facto standard for multi-class outputs

$$E(\mathbf{w}) = -\sum_{n=1}^{N} \sum_{k=1}^{K} \left\{ \mathbb{I}\left(t_n = k\right) \ln \frac{\exp(\mathbf{w}_k^{\top} \mathbf{x})}{\sum_{j=1}^{K} \exp(\mathbf{w}_j^{\top} \mathbf{x})} \right\}$$

#### Practical issue

- Exponentials get very big and can have vastly different magnitudes.
- Trick 1: Do not compute first softmax, then log, but instead directly evaluate log-exp in the nominator and log-sum-exp in the denominator.
- > Trick 2: Softmax has the property that for a fixed vector  ${\bf b}$   ${\rm softmax}({\bf a}+{\bf b})={\rm softmax}({\bf a})$
- $\Rightarrow$  Subtract the largest weight vector  $\mathbf{w}_j$  from the others.

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### Gradient Descent

- Two main steps
  - 1. Computing the gradients for each weight
  - 2. Adjusting the weights in the direction of the gradient
- Recall: Basic update equation

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

- Main questions
  - On what data do we want to apply this?
  - > How should we choose the step size  $\eta$  (the learning rate)?
  - In which direction should we update the weights?



### Stochastic vs. Batch Learning

- Batch learning
  - Process the full dataset at once to compute the gradient.

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

- Stochastic learning
  - Choose a single example from the training set.
  - Compute the gradient only based on this example
  - This estimate will generally be noisy, which has some advantages.

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

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### Stochastic vs. Batch Learning

- Batch learning advantages
  - Conditions of convergence are well understood.
  - Many acceleration techniques (e.g., conjugate gradients) only operate in batch learning.
  - Theoretical analysis of the weight dynamics and convergence rates are simpler.
- Stochastic learning advantages
  - Usually much faster than batch learning.
  - Often results in better solutions.
  - Can be used for tracking changes.
- Middle ground: Minibatches



### Minibatches

- Idea
  - Process only a small batch of training examples together
  - Start with a small batch size & increase it as training proceeds.

#### Advantages

- Gradients will be more stable than for stochastic gradient descent, but still faster to compute than with batch learning.
- > Take advantage of redundancies in the training set.
- > Matrix operations are more efficient than vector operations.
- Caveat
  - Error function should be normalized by the minibatch size, s.t. we can keep the same learning rate between minibatches

$$E(\mathbf{W}) = \frac{1}{N} \sum_{n} L(t_n, y(\mathbf{x}_n; \mathbf{W})) + \frac{\lambda}{N} \Omega(\mathbf{W})$$

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### Choosing the Right Learning Rate

- Analyzing the convergence of Gradient Descent
  - Consider a simple 1D example first

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

> What is the optimal learning rate  $\eta_{\text{opt}}$ ?



> If E is quadratic, the optimal learning rate is given by the inverse of the Hessian

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

What happens if we exceed this learning rate?



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### Choosing the Right Learning Rate

• Behavior for different learning rates



B. Leibe Image source: Yann LeCun et al., Efficient BackProp (1998)



### Learning Rate vs. Training Error



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### Batch vs. Stochastic Learning

- Batch Learning
  - Simplest case: steepest decent on the error surface.
  - ⇒ Updates perpendicular to contour lines





#### Stochastic Learning

- Simplest case: zig-zag around the direction of steepest descent.
- ⇒ Updates perpendicular to constraints from training examples.



43 Image source: Geoff Hinton

### Why Learning Can Be Slow

- If the inputs are correlated
  - > The ellipse will be very elongated.
  - The direction of steepest descent is almost perpendicular to the direction towards the minimum!



#### This is just the opposite of what we want!

Slide adapted from Geoff Hinton

### The Momentum Method

• Idea

- Instead of using the gradient to change the position of the weight "particle", use it to change the velocity.
- Intuition
  - Example: Ball rolling on the error surface
  - It starts off by following the error surface, but once it has accumulated momentum, it no longer does steepest decent.

### Effect

- Dampen oscillations in directions of high curvature by combining gradients with opposite signs.
- Build up speed in directions with a gentle but consistent gradient.

## The Momentum Method: Implementation

- Change in the update equations
  - > Effect of the gradient: increment the previous velocity, subject to a decay by  $\alpha < 1.$

$$\mathbf{v}(t) = \alpha \mathbf{v}(t-1) - \varepsilon \frac{\partial E}{\partial \mathbf{w}}(t)$$

Set the weight change to the current velocity

$$\Delta \mathbf{w} = \mathbf{v}(t)$$
  
=  $\alpha \mathbf{v}(t-1) - \varepsilon \frac{\partial E}{\partial \mathbf{w}}(t)$   
=  $\alpha \Delta \mathbf{w}(t-1) - \varepsilon \frac{\partial E}{\partial \mathbf{w}}(t)$ 



### The Momentum Method: Behavior

- Behavior
  - If the error surface is a tilted plane, the ball reaches a terminal velocity

$$\mathbf{v}(\infty) = \frac{1}{1-\alpha} \left( -\varepsilon \frac{\partial E}{\partial \mathbf{w}} \right)$$

- If the momentum  $\alpha$  is close to 1, this is much faster than simple gradient descent.
- > At the beginning of learning, there may be very large gradients.
  - Use a small momentum initially (e.g.,  $\alpha = 0.5$ ).
  - Once the large gradients have disappeared and the weights are stuck in a ravine, the momentum can be smoothly raised to its final value (e.g.,  $\alpha = 0.90$  or even  $\alpha = 0.99$ ).
- $\Rightarrow$  This allows us to learn at a rate that would cause divergent oscillations without the momentum.



### 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.





### 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





### **Behavior in a Long Valley**



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### **Behavior around a Saddle Point**



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## Visualization of Convergence Behavior



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.



#### Be careful: Do not turn down the learning rate too soon!

Further progress will be much slower/impossible after that.  $\geq$ 

Effect

 $\geq$ 



### 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 (to be continued in the next lecture...)

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### • Tricks of the Trade

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### 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

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### **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).



### **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.