Security and Fairness of Deep Learning

# Deep Feedforward Networks

Anupam Datta CMU

Spring 2019

#### **Announcements**

- Please Create an Xsede Account now (pinned Piazza post)
  - Will be used for HW2
- HW2 will be released shortly

### Story so far

- Image classification problem
- Linear models
  - Score function
  - Loss function
  - Learning
- Learning as optimization
  - Gradient descent (batch, mini-batch, stochastic)
  - Second-order methods (Newton's method)
  - Backpropagation

## Today

- From linear score functions to neural networks
  - Practical design choices
  - (Some) justification of design choices

#### Recall: Linear score function

$$f(x_i, W) = Wx_i$$

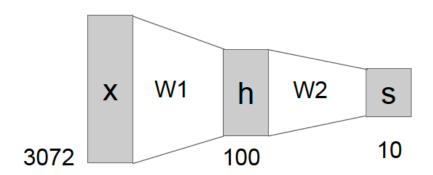
						Ι.	
	0.2	-0.5	0.1	2.0	1.1		56
	1.5	1.3	2.1	0.0	3.2		231
	0	0.25	0.2	-0.3	-1.2		24
W $b$							2
							1
							$x_i$

For CIFAR:

W: 10 x 3072 x: 3072 x 1 10 class scores

#### 2-Layer neural network

$$s = W_2 \max(0, W_1 x)$$



For CIFAR:

W1: 100 x 3072 W2: 10 x 100 x: 3072 x 1 10 class scores

- Iterated construction: linear function followed by non-linear function
- Training network: learn W1, W2 using stochastic gradient descent; use backpropagation to compute gradients

## Topic outline

- Setting up the architecture
- Setting up the data and the loss
- Learning and evaluation

Neural network architecture

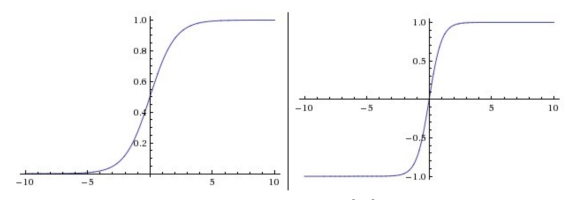
#### Outline

- Modeling one neuron
  - Single neuron as a linear classifier
  - Commonly used activation functions
- Neural Network architectures
  - Layer-wise organization
  - Example feed-forward computation
  - Representational power
  - Setting number of layers and their sizes

### Single neuron as a linear classifier

- Binary softmax classifier
  - Interpret  $\,\sigma(\sum_i w_i x_i + b)\,$  to be the probability of one of the classes  $\,P(y_i=1\mid x_i;w)\,$
  - Set threshold at 0.5
- Binary SVM classifier
  - Attach a max-margin hinge loss to the output of the neuron

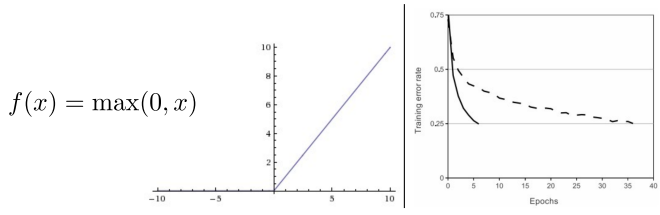
### Commonly used activation functions



Left: Sigmoid non-linearity squashes real numbers to range between [0,1] Right: The tanh non-linearity squashes real numbers to range between [-1,1].

- Sigmoid weaknesses:
  - saturate and kill gradients
  - outputs not zero-centered
- Tanh outputs are zero-centered

#### Commonly used activation functions



Left: Rectified Linear Unit (ReLU) activation function, which is zero when x < 0 and then linear with slope 1 when x > 0. Right: A plot from Krizhevsky et al. (pdf) paper indicating the 6x improvement in convergence with the ReLU unit compared to the tanh unit.

- ReLU is often used in modern deep networks
  - Linear, non-saturating form speeds up convergence of stochastic gradient descent; efficient to compute (threshold operation)
  - If learning rate is high, then ReLU units can die i.e., never activate during subsequent training

### Commonly used activation functions

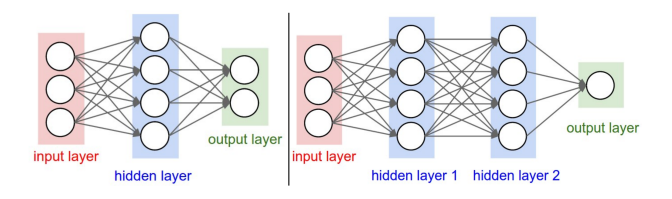
- Leaky ReLU
  - Function has small negative slope when x < 0 to avoid dying

$$f(x) = \mathcal{I}(x < 0)(\alpha x) + \mathcal{I}(x >= 0)(x)$$

- Maxout
  - Generalizes ReLU and Leaky ReLU; advantages of both but more parameters

$$\max(w_1^T x + b_1, w_2^T x + b_2)$$

#### Neural network architectures



- Neural network as a directed acyclic graph
- Examples above: 2-layer NN and 3-layer NN
- Fully connected layer

### Example feedforward computation

```
# forward-pass of a 3-layer neural network:
f = lambda x: 1.0/(1.0 + np.exp(-x)) # activation function (use sigmoid)
x = np.random.randn(3, 1) # random input vector of three numbers (3x1)
h1 = f(np.dot(W1, x) + b1) # calculate first hidden layer activations (4x1)
h2 = f(np.dot(W2, h1) + b2) # calculate second hidden layer activations (4x1)
out = np.dot(W3, h2) + b3 # output neuron (1x1)
```

- Repeated matrix multiplications interwoven with activation function
- x could hold a batch of training data evaluated in parallel
- Output layer neurons do not go through non-linear activation function

#### Representational power

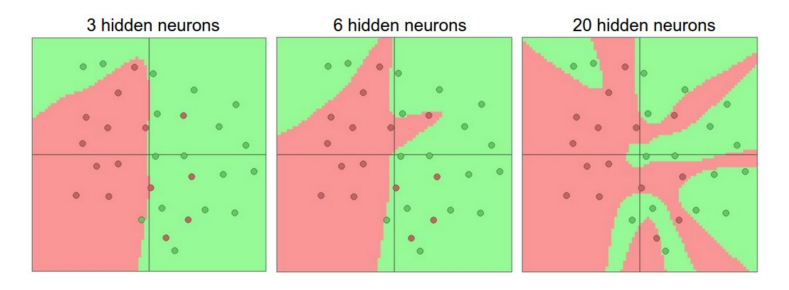
Neural Networks with at least one hidden layer are *universal* approximators:

Given any continuous function f(x) and some  $\epsilon>0$ , there exists a Neural Network g(x) with one hidden layer (with a reasonable choice of non-linearity, e.g. sigmoid) such that  $\forall x, |f(x)-g(x)| < \epsilon \forall x$ 

#### Representational power

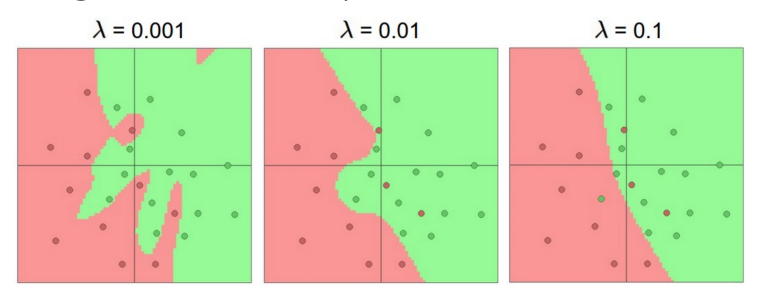
- Neural Networks work well in practice because they compactly express nice, smooth functions that fit well with the statistical properties of data we encounter in practice, and are also easy to learn using our optimization algorithms (e.g. gradient descent).
- The fact that deeper networks (with multiple hidden layers) can work better than a single-hidden-layer networks is an empirical observation, despite the fact that their representational power is equal.

### Setting number of layers and their sizes



With more neurons, we have greater representation power but possibly more overfitting

## Setting number of layers and their sizes



Train large network; control overfitting with regularization

The Loss Surfaces of Multilayer Networks

Setting up the data and the model

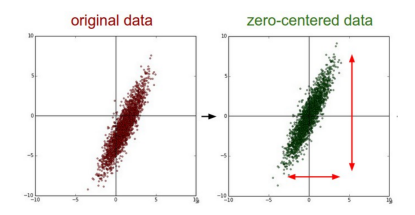
#### Outline

- Setting up the data and the model
  - Data Preprocessing
  - Weight Initialization
  - Regularization

• Loss functions

#### Data preprocessing

- Mean subtraction
  - Subtract the mean across every individual feature in the data

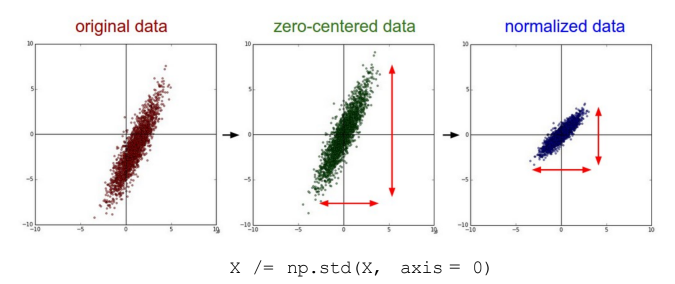


 $X \rightarrow np.mean(X, axis = 0)$ 

Data matrix X, where we will assume that X is of size  $[N \times D]$  (N is the number of data, D is their dimensionality)

#### Data preprocessing

- Normalization
  - Divide each zero-centered feature by its standard deviation
  - Bringing data dimensions to same scale helps SGD converge



#### Outline

- Setting up the data and the model
  - Data Preprocessing
  - Weight Initialization
  - Regularization

• Loss functions

## Weight initialization

- First attempt
  - Initialize all weights to 0
  - Not a good idea
    - Every neuron computes the same output => every neuron computes the same gradients and undergoes the same parameter updates

### Weight initialization

- Important to introduce asymmetry
  - Idea: Initialize weights to independent small random numbers

```
W = 0.01* np.random.randn(D, H)
```

where randn samples from a zero mean, unit standard deviation gaussian.

• Issue: Distribution of the outputs from a randomly initialized neuron has a variance that grows with the number of inputs

### Weight initialization

 Recommended practice for initializing weights of neurons in NNs with ReLU units

```
w = np.random.randn(n) * sqrt(2.0/n)
where n is the number of its inputs
```

• Every neuron's weight vector is sampled from a multi-dimensional gaussian normalized by its variance

<u>Delving Deep into Rectifiers: Surpassing Human-Level Performance on</u> ImageNet Classification

#### Weight initialization (under simplifying assumptions)

Initialize weights of NN as follows

```
w = np.random.randn(n) * sqrt(1.0/n)
where n is the number of its inputs
```

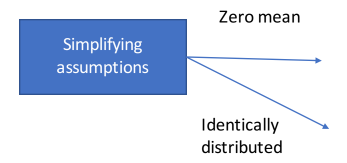
• Every neuron's weight vector is sampled from a multi-dimensional gaussian normalized by its variance

Based on: Understanding the difficulty of training deep feedforward neural networks

#### Weight initialization (under simplifying assumptions)

$$s = \sum_{i}^{n} w_i x_i$$
  $\operatorname{Var}(s) = \operatorname{Var}(\sum_{i}^{n} w_i x_i)$ 

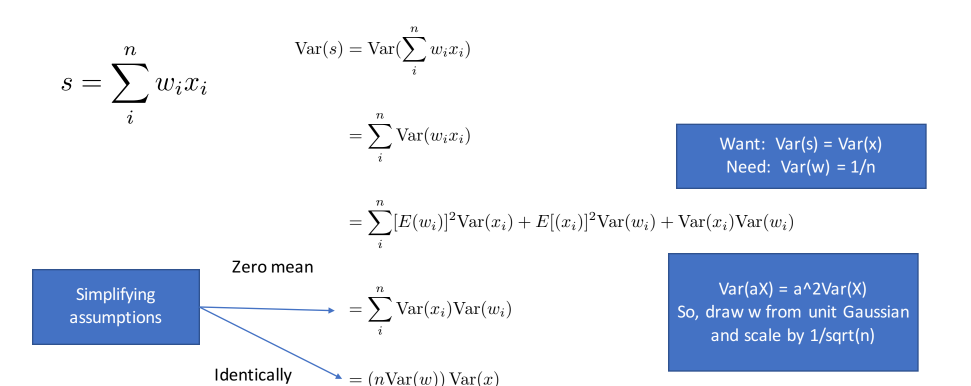
Want: Var(s) = Var(x) Need: Var(w) = 1/n



Var(aX) = a^2Var(X)
So, draw w from unit Gaussian
and scale by 1/sqrt(n)

#### Weight initialization (under simplifying assumptions)

distributed



#### Bias initialization

• Initialize biases to 0

#### Outline

- Setting up the data and the model
  - Data Preprocessing
  - Weight Initialization
  - Regularization

• Loss functions

#### Recall: loss function

$$L = \frac{1}{N} \sum_{i} L_{i} + \underbrace{\lambda R(W)}_{\text{regularization loss}}$$

### Regularization

• L2 regularization

For every weight w in the network, we add the term  $\frac{1}{2}\lambda w^2$  to the objective, where  $\lambda$  is the regularization strength

- Encourages the network to use all of its inputs a little rather than some of its inputs a lot
- During gradient descent parameter update, every weight is decayed linearly toward zero

#### Regularization

• L1 regularization

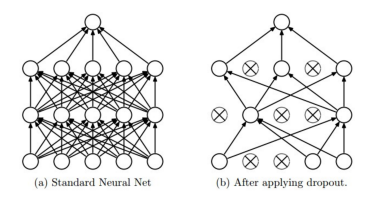
For every weight w in the network, we add the term  $\lambda |w|$  to the objective, where  $\lambda$  is the regularization strength

- Encourages the network to use some of its inputs a lot (i.e. sparse weight vectors)
- If explicit feature selection is not a goal, L2 regularization usually performs better than L1 regularization

#### Regularization

#### • Dropout

- Sample a neural network within the full network and only update its parameters
- Typically hidden units retained with p = 0.5, input units with p close to 1



Dropout: A Simple Way to Prevent Neural Networks from Overfitting

## Regularization: Dropout

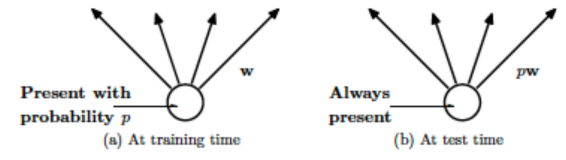


Figure 2: Left: A unit at training time that is present with probability p and is connected to units in the next layer with weights w. Right: At test time, the unit is always present and the weights are multiplied by p. The output at test time is same as the expected output at training time.

# Regularization: Dropout

```
""" Vanilla Dropout: Not recommended implementation (see notes below) """
p = 0.5 # probability of keeping a unit active. higher = less dropout
def train step(X):
  """ X contains the data """
 # forward pass for example 3-layer neural network
 H1 = np.maximum(0, np.dot(W1, X) + b1)
 U1 = np.random.rand(*H1.shape) 
 H1 *= U1 # drop!
 H2 = np.maximum(0, np.dot(W2, H1) + b2)
 U2 = np.random.rand(*H2.shape) 
 H2 *= U2 # drop!
 out = np.dot(W3, H2) + b3
 # backward pass: compute gradients... (not shown)
 # perform parameter update... (not shown)
def predict(X):
 # ensembled forward pass
 H1 = np.maximum(0, np.dot(W1, X) + b1) * p # NOTE: scale the activations
 H2 = np.maximum(0, np.dot(W2, H1) + b2) * p # NOTE: scale the activations
 out = np.dot(W3, H2) + b3
```

#### Regularization: Inverted Dropout

```
Inverted Dropout: Recommended implementation example.
We drop and scale at train time and don't do anything at test time.
p = 0.5 # probability of keeping a unit active. higher = less dropout
def train step(X):
 # forward pass for example 3-layer neural network
 H1 = np.maximum(0, np.dot(W1, X) + b1)
 U1 = (np.random.rand(*H1.shape) < p) / p # first dropout mask. Notice /p!</pre>
 H1 *= U1 # drop!
 H2 = np.maximum(0, np.dot(W2, H1) + b2)
 U2 = (np.random.rand(*H2.shape) < p) / p # second dropout mask. Notice /p!
  H2 *= U2 # drop!
  out = np.dot(W3, H2) + b3
  # backward pass: compute gradients... (not shown)
  # perform parameter update... (not shown)
def predict(X):
 # ensembled forward pass
 H1 = np.maximum(0, np.dot(W1, X) + b1) # no scaling necessary
 H2 = np.maximum(0, np.dot(W2, H1) + b2)
  out = np.dot(W3, H2) + b3
```

#### Regularization

#### In practice

It is most common to use a single, global L2 regularization strength that is cross-validated. It is also common to combine this with dropout applied after all layers. The value of p=0.5 is a reasonable default, but this can be tuned on validation data.

#### Outline

- Setting up the data and the model
  - Data Preprocessing
  - Weight Initialization
  - Regularization

• Loss functions

#### Loss functions for classification

**Data loss** 

$$L = \frac{1}{N} \sum_{i} L_{i}$$

SVM loss

$$L_i = \sum_{j \neq y_i} \max(0, f_j - f_{y_i} + 1)$$

Cross-entropy loss

$$L_i = -\log\left(\frac{e^{f_{y_i}}}{\sum_j e^{f_j}}\right)$$

Learning and evaluation

#### Outline

- Gradient checks
- Monitoring the learning process
- Parameter updates
- Hyperparameter Optimization
- Evaluation
  - Model Ensembles

#### Gradient checks

- SGD uses gradients that we computed analytically using calculus
- Issue: How do we check that we did not make errors?
- Compare analytic gradient implementation to the numerical gradient

$$\frac{df(x)}{dx} = \lim_{h \to 0} \frac{f(x+h) - f(x)}{h}$$

$$\frac{f(x+h)-f(x)}{h}$$
 where  $h \approx 10^{-5}$ 

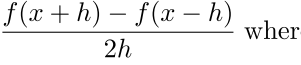
## Gradient checks: tip

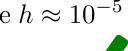
Use centered formula

$$\frac{df(x)}{dx} = \lim_{h \to 0} \frac{f(x+h) - f(x)}{h} \qquad \frac{df(x)}{dx} = \lim_{h \to 0} \frac{f(x+h) - f(x-h)}{2h}$$

$$\frac{df(x)}{dx} = \lim_{h \to 0} \frac{f(x+h) - f(x-h)}{2h}$$

$$\frac{f(x+h) - f(x)}{h} \text{ where } h \approx 10^{-5} \qquad \frac{f(x+h) - f(x-h)}{2h} \text{ where } h \approx 10^{-5}$$





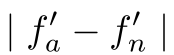


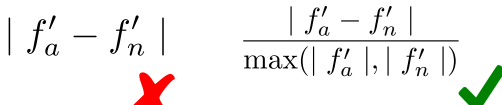
Use Taylor expansion of f(x+h) and f(x-h): first formula has an error on order of O(h), while the second formula only has error terms on order of O(h^2)

# Gradient checks:tip

#### Use relative error

- Analytical gradient:  $f_a^\prime$
- Numerical gradient:  $f'_n$





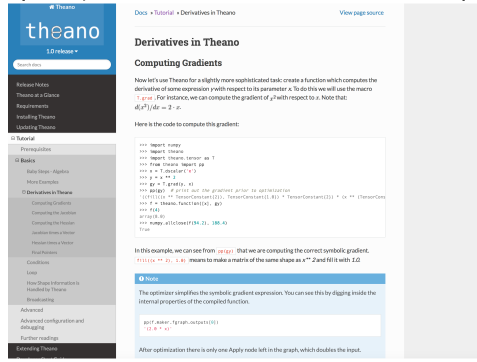


#### • In practice:

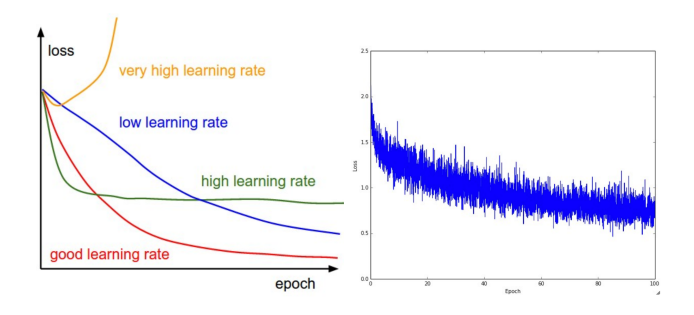
- relative error > 1e-2 usually means the gradient is probably wrong
- 1e-2 > relative error > 1e-4 should make you feel uncomfortable
- 1e-4 > relative error is usually okay for objectives with kinks. But if there are no kinks (e.g. use of tanh nonlinearities and softmax), then 1e-4 is too high.
- 1e-7 and less you should be happy.

## Gradient computation

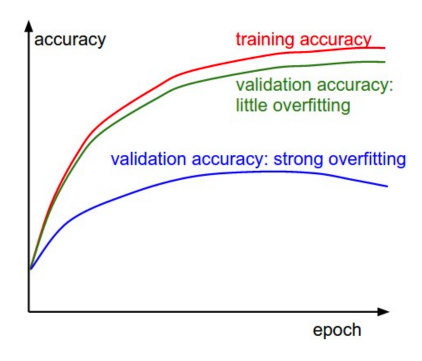
• Symbolic (analytical) differentiation available in deep learning libraries



#### • Loss function



Train/Val accuracy



Ratio of updates:weights

A rough heuristic is that this ratio should be somewhere around 1e-3.

If it is lower than this then the learning rate might be too low. If it is higher then the learning rate is likely too high.

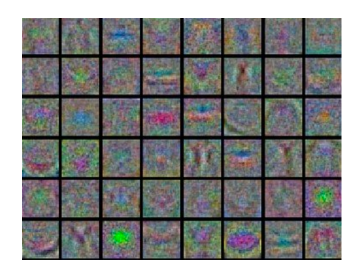
- Activation / Gradient distributions per layer
  - plot activation/gradient histograms for all layers of the network
  - not a good sign to see any strange distributions
  - e.g. with tanh neurons we would like to see a distribution of neuron activations between the full range of [-1,1], instead of seeing all neurons outputting zero, or all neurons being completely saturated at either -1 or 1.

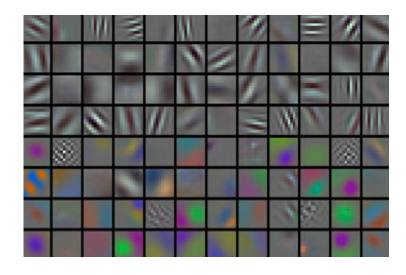
## Annealing the learning rate

#### Step decay

- Reduce the learning rate by some factor every few epochs.
- Typical values might be reducing the learning rate by a half every 5 epochs, or by 0.1 every 20 epochs. These numbers depend heavily on the type of problem and the model.
- One heuristic you may see in practice is to watch the validation error while training with a fixed learning rate, and reduce the learning rate by a constant (e.g. 0.5) whenever the validation error stops improving.

• First-layer Visualizations





# Parameter updates

• Vanilla SGD

```
# Vanilla update
x += - learning_rate * dx
```

where learning\_rate is a hyperparameter - a fixed constant.

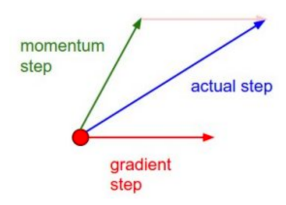
#### Parameter updates

#### • Momentum update

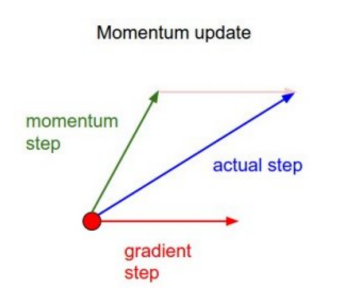
```
# Momentum update
v = mu * v - learning_rate * dx # integrate velocity
x += v # integrate position
```

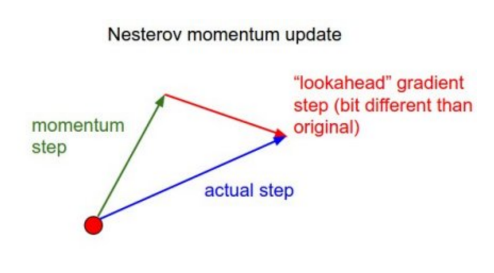
A typical setting is to start with momentum of about 0.5 and anneal it to 0.99 or so over multiple epochs (cf. learning rate is decreased over time)

#### Momentum update



#### Nesterov momentum





# Per-parameter adaptive learning rate methods

- Adaptively tune learning rate per parameter (instead of single global learning rate)
  - Adagrad
  - RMSprop

#### Adagrad

```
# Assume the gradient dx and parameter vector x
cache += dx**2
x += - learning_rate * dx / (np.sqrt(cache) + eps)
```

- cache has size equal to the size of the gradient, and keeps track of perparameter sum of squared gradients
- Reduce effective learning rate more for weights with high gradients
- eps (usually set somewhere in range from 1e-4 to 1e-8) avoids division by zero
- Monotonically decreasing learning rate may stop learning too early

#### Example

- <w1, w2>
- w1: 1, 1, 1
- w2: 0.2, 0.2, 0.2
- cache =  $\sqrt{3}$ ,  $\sqrt{.12}$
- Effective gradient for w1 =  $1/\sqrt{3}$  = 0.76
- Effective gradient for w2 = 0.2/  $\sqrt{.12}$  = 0.35

#### **RMSprop**

```
cache = decay_rate * cache + (1 - decay_rate) * dx**2
x += - learning_rate * dx / (np.sqrt(cache) + eps)
```

- Adjusts Adagrad to reduce aggressive, monotonically decreasing learning rate
- Uses a moving average of squared gradients
- decay\_rate is a hyperparameter and typical values are [0.9, 0.99, 0.999]
- Unlike Adagrad the updates do not get monotonically smaller

#### Adam

```
m = beta1*m + (1-beta1)*dx
v = beta2*v + (1-beta2)*(dx**2)
x += - learning_rate * m / (np.sqrt(v) + eps)
```

- RMSProp with momentum
- smooth version of the gradient m used instead of gradient dx
- Recommended values: eps = 1e-8, beta1 = 0.9, beta2 = 0.999

- Some hyperparameters
  - the initial learning rate
  - learning rate decay schedule (such as the decay constant)
  - regularization strength (L2 penalty, dropout strength)

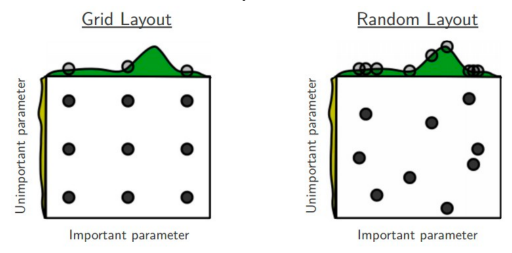
- Hyperparameter ranges
  - Search for multiplicative hyperparameters (e.g., learning rate, regularization strength) on a log scale

```
learning_rate = 10 ** uniform(-6, 1)
```

• Dropout parameter search on original scale

```
dropout = uniform(0,1)
```

 Random search better than grid search on high dimensional spaces with low effective dimensionality



Random Search for Hyper-Parameter Optimization

- Search from coarse to fine ranges
  - First search in coarse ranges (e.g. 10 \*\* [-6, 1]), and then depending on where the best results are turning up, narrow the range.

#### Model ensembles

- Approach
  - Train multiple independent models, and at test time average their predictions
- Training"independent" models
  - Same model, different initializations
  - Top models discovered during cross-validation
  - Different checkpoints of a single model
  - Running average of parameters during training

Hinton et al., Distilling the Knowledge in a Neural Network

# Many other tools to improve training

- Second-order methods (Newton, L-BFGS)
- Momentum
- Batch normalization
- Adam
- •

# Acknowledgment

Based in part on material from Stanford CS231n <a href="http://cs231n.github.io/">http://cs231n.github.io/</a>

#### **Batch Normalization**

• Batch Normalization is a technique that alleviates problems with proper initialization of neural networks

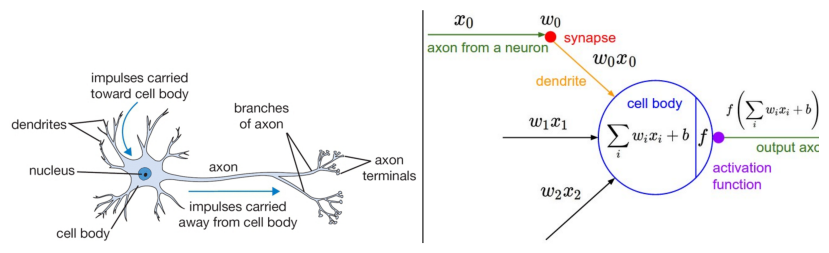
• We will discuss it in a later lecture

## Numerical gradient

```
def eval_numerical_gradient(f, x):
 a naive implementation of numerical gradient of f at x
 - f should be a function that takes a single argument
 - x is the point (numpy array) to evaluate the gradient at
 fx = f(x) # evaluate function value at original point
 grad = np.zeros(x.shape)
 h = 0.00001
 # iterate over all indexes in x
 it = np.nditer(x, flags=['multi index'], op flags=['readwrite'])
 while not it.finished:
   # evaluate function at x+h
   ix = it.multi_index
   old value = x[ix]
   x[ix] = old value + h # increment by h
   fxh = f(x) # evalute f(x + h)
   x[ix] = old value # restore to previous value (very important!)
   # compute the partial derivative
   grad[ix] = (fxh - fx) / h # the slope
   it.iternext() # step to next dimension
 return grad
```

A problem of efficiency

## Biological motivation



A cartoon drawing of a biological neuron (left) and its mathematical model (right).

Imperfect analogy!

#### Second-order methods

- Newton's method does not scale
  - Computing inverse Hessian explicitly is too expensive
- Quasi-newton method L-BFGS works quite well
  - Iteratively build up limited memory approximation of Hessian

Dean et al. Large Scale Distributed Deep Networks