## Neural Networks

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## Logistics

- Need your presentation preference by the end of this class. Again, please give me three package names with order of preference. The final decision will be computed by minimizing the following cost function :)
- $\sum_{\text {student }}$ student cost $+\sum_{\text {package }}$ package cost
- student cost $= \begin{cases}0, & \text { first priority } \\ 2.5, & \text { second priority } \\ 5, & \text { third priority }\end{cases}$
- package cost $=\alpha \cdot 2^{(\text {num }}$ presentations covered $)$


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- Most popular package (in terms of first priority pick) will be presented first. If there is a tie, I will break it with popularity based all choices regardless of priority. If there is a tie, I will break it by random


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- Most popular package (in terms of first priority pick) will be presented first. If there is a tie, I will break it with popularity based all choices regardless of priority. If there is a tie, I will break it by random
- Students presenting the same packages will be ordered randomly


## Logistics

- HW1 due this Thursday
- Package choice due this Thursday


## Review

In the last couple classes, we discussed

- Basic concepts of regression and classification
- Examples of regularization such as ridge $\left(l_{2}\right)$ regression and lasso $\left(l_{1}\right)$
- Linear classifiers including logistic regression and softmax classifier


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- We described stochastic gradient descent and momentum trick for classification
- We also briefly went through SVM and hinge loss


## SVM



- Denote $\hat{\mathbf{w}}=\frac{\mathbf{w}}{\|\mathbf{w}\|}, \hat{\mathbf{w}} \cdot \mathbf{x}_{1}$ ( $\hat{\mathbf{w}} \cdot \mathbf{x}_{-1}$ ) is the distance of the boundary line of $\mathbf{x}_{1}\left(\mathbf{x}_{-1}\right)$ from the origin


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- Thus, the distance between the two boundary lines is $\hat{\mathbf{w}} \cdot\left(\mathbf{x}_{1}-\mathbf{x}_{-1}\right)=\frac{2}{\|\mathbf{w}\|}$


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- SVM: for all $\mathbf{x}_{i}$
$\max \frac{2}{\|\mathbf{w}\|} \quad$ s.t. $\quad y_{i}\left(\mathbf{w} \cdot \mathbf{x}_{i}-b\right) \geq 1$


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$\max \frac{2}{\|\mathbf{w}\|} \quad$ s.t. $\quad y_{i}\left(\mathbf{w} \cdot \mathbf{x}_{i}-b\right) \geq 1$
Equivalently,

$$
\min \|\mathbf{w}\| \quad \text { s.t. } \quad y_{i}\left(\mathbf{w} \cdot \mathbf{x}_{i}-b\right) \geq 1
$$

## Soft-margin SVM and hinge loss

- Hard-margin SVM

$$
\min \|\mathbf{w}\| \quad \text { s.t. } \quad y_{i}\left(\mathbf{w} \cdot \mathbf{x}_{i}-b\right)-1 \geq 0
$$



- Soft-margin SVM (allow constrain to be violate)
- Define "hinge" loss function

$$
h(z)=\max (0, z)
$$

- Want to minimize hinge loss

$$
\sum_{i} h\left(1-y_{i}\left(\mathbf{w} \cdot \mathbf{x}_{i}-b\right)\right)
$$

- Soft-margin SVM

$$
\min \lambda\|\mathbf{w}\|^{2}+\sum_{i} h\left(1-y_{i}\left(\mathbf{w} \cdot \mathbf{x}_{i}-b\right)\right)
$$

## Multi-class SVM

- We can easily extend soft-margin SVM to multi-class case. Let $s_{l}(\mathbf{x})=\mathbf{w}_{\mathbf{l}}^{T}\left[\begin{array}{l}1 \\ \mathbf{x}\end{array}\right]$ be the score for class $l$.


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\sum_{l \neq j} h\left(s_{l}(\mathbf{x})-s_{j}(\mathbf{x})+\Delta\right)=\sum_{l \neq j} \max \left(0, s_{l}(\mathbf{x})-s_{j}(\mathbf{x})+\Delta\right),
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where $j$ is the true label of $\mathbf{x}$ and $\Delta$ contributes a margin ensuring that the true label score has to be at least $\Delta$ more than the rest to be penalty free

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- Multi-class SVM:

$$
\min \lambda\|\mathbf{w}\|^{2}+\sum_{i} \sum_{l \neq j\left(\mathbf{x}_{i}\right)} h\left(s_{l}\left(\mathbf{x}_{i}\right)-s_{j\left(\mathbf{x}_{i}\right)}\left(\mathbf{x}_{i}\right)+\Delta\right)
$$

## Perceptron

- Perceptron is an artificial neuron with step function as activation function



## Perceptron



- Perceptron is an artificial neuron with step function as activation function
- It is impossible to extend perceptron to multilayer. Multilayer perceptron (MLP) is a misnomer. Step activation function is never used multilayer neural networks (not trainable)


## Perceptron



- Perceptron is an artificial neuron with step function as activation function
- It is impossible to extend perceptron to multilayer. Multilayer perceptron (MLP) is a misnomer. Step activation function is never used multilayer neural networks (not trainable)
- According to Hinton, perceptrons are still used in systems with large number (millions) of features. Other than that, it has relatively limited use since most problems are not linearly separable


## Perceptron

- In most cases, perceptron would be useful if only one manages to handcode inputs into separable features



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- That was the main area of research in many machine learning applications-finding efficient ways to generate good features


## Perceptron



- In most cases, perceptron would be useful if only one manages to handcode inputs into separable features
- That was the main area of research in many machine learning applications-finding efficient ways to generate good features
- One attractive characteristic of deep learning (neural networks) is that we not only can train the classifier but also can learn the appropriate features automatically


## Nomenclature of basic network architectures

## Neural Networks: Architectures



hidden layer 1 hidden layer 2
"3-layer Neural Net", or "2-hidden-layer Neural Net"
"Fully-connected" layers

Fei-Fei Li \& Andrej Karpathy \& Justin Johnson Lecture 4-77 13 Jan 2016

## Caveat: don't go too far for the brain analogy

Biological neurons:

- Many different types
- Dendrite can perform complex non-linear operations
- Synapses are not a single weight but a complex non-linear dynamical system
- Rate code may not be adequate

Also see London 2005 (Slide credit: CS231n)

## Back-propagation and computational graph

- As described in last lecture, training in supervised learning system often boils down to minimizing of loss function w.r.t. some parameters


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- Back-propagation (BP) is an efficient way to find such derivation. Actually it is in fact just another way of spelling out the chain rule $\frac{\partial L}{\partial x}=\frac{\partial L}{\partial y} \frac{\partial y}{\partial x}$ in calculus


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- It is often easier to explain BP in terms of computational graph
- Computational graph can be interpreted as generalization of a neural networks
- Neuron no longer will be restricted to summation and activation function but can be any computation as well (e.g., max)


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- Computational graph can be interpreted as generalization of a neural networks
- Neuron no longer will be restricted to summation and activation function but can be any computation as well (e.g., max)
- Let me try to explain through an example


## Multi-class SVM

## Computational graphs



Fei-Fei Li \& Justin Johnson \& Serena Yeung Lecture 4-8
April 13, 2017

## More complex example

## Convolutional network (AlexNet)



## A simple BP example

$$
\begin{aligned}
& f(x, y, z)=(x+y) z \\
& \text { e.g. } x=-2, y=5, z=-4
\end{aligned}
$$



## A simple BP example

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q=x+y \quad \frac{\partial q}{\partial x}=1, \frac{\partial q}{\partial y}=1
$$



$$
f=q z \quad \frac{\partial f}{\partial q}=z, \frac{\partial f}{\partial z}=q
$$

Want: $\frac{\partial f}{\partial x}, \frac{\partial f}{\partial y}, \frac{\partial f}{\partial z}$

Fei-Fei Li \& Andrej Karpathy \& Justin Johnson Lecture 4-11 13 Jan 2016

## A simple BP example

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\begin{aligned}
& f(x, y, z)=(x+y) z \\
& \text { e.g. } \mathrm{x}=-2, \mathrm{y}=5, \mathrm{z}=-4 \\
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\end{aligned}
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Fei-Fei Li \& Andrej Karpathy \& Justin Johnson Lecture 4-12 13 Jan 2016

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Fei-Fei Li \& Andrej Karpathy \& Justin Johnson Lecture 4-13 13 Jan 2016

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Fei-Fei Li \& Andrej Karpathy \& Justin Johnson Lecture 4-14 13 Jan 2016

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Fei-Fei Li \& Andrej Karpathy \& Justin Johnson Lecture 4-15 13 Jan 2016

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## A simple BP example

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## A simple BP example

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$$

$$
f=q z \quad \frac{\partial f}{\partial q}=z, \frac{\partial f}{\partial z}=q
$$

Chain rule:

$$
\frac{\partial f}{\partial y}=\frac{\partial f}{\partial q} \frac{\partial q}{\partial y}
$$

Want: $\frac{\partial f}{\partial x}, \frac{\partial f}{\partial y}, \frac{\partial f}{\partial z}$

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## A simple BP example

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## BP at one node



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## BP at one node



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## BP at one node



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## BP at one node



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## BP at one node



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## BP at one node



## Yet another BP example

Another example: $\quad f(w, x)=\frac{1}{1+e^{-\left(u_{0} x_{0}+w_{1}, x_{1}+w_{2}\right)}}$


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## Yet another BP example

Another example: $\quad f(w, x)=\frac{1}{1+e^{-\left(w_{0} x_{0}+w_{1} x_{1}+w_{2}\right)}}$


$$
\begin{array}{lll|lll}
f(x)=e^{x} & \rightarrow & \frac{d f}{d x}=e^{x} & f(x)=\frac{1}{x} & \rightarrow & \frac{d f}{d x}=-1 / x^{2} \\
f_{a}(x)=a x & \rightarrow & \frac{d f}{d x}=a & f_{c}(x)=c+x & \rightarrow & \frac{d f}{d x}=1
\end{array}
$$

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## Yet another BP example

Another example: $\quad f(w, x)=\frac{1}{1+e^{-\left(w_{0} x_{0}+w_{1} x_{1}+w_{2}\right)}}$


| $f(x)=e^{x}$ | $\rightarrow$ | $\frac{d f}{d x}=e^{x}$ | $f(x)=\frac{1}{x}$ | $\rightarrow$ |
| :--- | :--- | :--- | :--- | :--- |
| $f_{a}(x)=a x$ | $\rightarrow$ | $\frac{d f}{d x}=a$ | $f_{c}(x)=c+x$ | $\frac{d f}{d x}=-1 / x^{2}$ |

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13 Jan 2016

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Another example: $\quad f(w, x)=\frac{1}{1+e^{-\left(w_{0} x_{0}+w_{1} x_{1}+w_{2}\right)}}$


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\hline
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$$

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| :--- | :--- | :--- | :---: | :---: | :---: |
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Lecture 4-33
13 Jan 2016

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| $f(x)=e^{x}$ | $\rightarrow$ | $\frac{d f}{d x}=e^{x}$ | $f(x)=\frac{1}{x}$ | $\rightarrow$ |  |
| :--- | :--- | :--- | :--- | :--- | :--- |
| $f_{a}(x)=a x$ | $\rightarrow$ | $\frac{d f}{d x}=a$ |  | $\frac{d f}{d x}=-1 / x^{2}$ |  |
| $f_{c}(x)=c+x$ |  | $\rightarrow$ | $\frac{d f}{d x}=1$ |  |  |

[^3]
## Yet another BP example

Another example: $\quad f(w, x)=\frac{1}{1+e^{-\left(u_{0} x_{0}+w_{1}, x_{1}+w_{2}\right)}}$


$$
\begin{array}{lll|ll}
f(x)=e^{x} & \rightarrow & \frac{d f}{d x}=e^{x} & f(x)=\frac{1}{x} & \rightarrow
\end{array} \quad \begin{array}{ll}
\frac{d f}{d x}=-1 / x^{2} \\
f_{a}(x)=a x & \rightarrow
\end{array}
$$

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## Yet another BP example

Another example: $\quad f(w, x)=\frac{1}{1+e^{-\left(w_{0} x_{0}+w_{1} x_{1}+w_{2}\right)}}$


$$
\begin{array}{lll|ll}
f(x)=e^{x} & \rightarrow & \frac{d f}{d x}=e^{x} & f(x)=\frac{1}{x} & \rightarrow \\
f_{a}(x)=a x & \rightarrow & \frac{d f}{d x}=a & f_{c}(x)=c+x & \rightarrow
\end{array} \begin{aligned}
& \frac{d f}{d x}=-1 / x^{2} \\
&
\end{aligned}
$$

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## Yet another BP example

Another example: $\quad f(w, x)=\frac{1}{1+e^{-\left(w_{0} x_{0}+w_{1} x_{1}+w_{2}\right)}}$


$$
\begin{array}{lll|lll}
f(x)=e^{x} & \rightarrow & \frac{d f}{d x}=e^{x} & f(x)=\frac{1}{x} & \rightarrow & \frac{d f}{d x}=-1 / x^{2} \\
f_{a}(x)=a x & \rightarrow & \frac{d f}{d x}=a & f_{c}(x)=c+x & \rightarrow & \frac{d f}{d x}=1
\end{array}
$$

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## Yet another BP example

Another example: $\quad f(w, x)=\frac{1}{1+e^{-\left(w_{0} x_{0}+w_{1} x_{1}+w_{2}\right)}}$


| $f(x)=e^{x}$ | $\rightarrow$ | $\frac{d f}{d x}=e^{x}$ | $f(x)=\frac{1}{x}$ | $\rightarrow$ | $\frac{d f}{d x}=-1 / x^{2}$ |
| :--- | :--- | :--- | :--- | :--- | :--- |
| $f_{a}(x)=a x$ | $\rightarrow$ | $\frac{d f}{d x}=a$ | $f_{c}(x)=c+x$ | $\rightarrow$ | $\frac{d f}{d x}=1$ |

[^4]
## Breaking down at different granularities

$$
\begin{aligned}
& f(w, x)=\frac{1}{1+e^{-\left(w_{0} x_{0}+w_{1} x_{1}+w_{2}\right)}} \quad \sigma(x)=\frac{1}{1+e^{-x}} \quad \text { sigmoid function } \\
& \frac{d \sigma(x)}{d x}=\frac{e^{-x}}{\left(1+e^{-x}\right)^{2}}=\left(\frac{1+e^{-x}-1}{1+e^{-x}}\right)\left(\frac{1}{1+e^{-x}}\right)=(1-\sigma(x)) \sigma(x)
\end{aligned}
$$

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## Breaking down at different granularities

$$
\begin{aligned}
& f(w, x)=\frac{1}{1+e^{-\left(w_{0} x_{0}+w_{1} x_{1}+w_{2}\right)}} \quad \sigma(x)=\frac{1}{1+e^{-x}} \quad \text { sigmoid function } \\
& \frac{d \sigma(x)}{d x}=\frac{e^{-x}}{\left(1+e^{-x}\right)^{2}}=\left(\frac{1+e^{-x}-1}{1+e^{-x}}\right)\left(\frac{1}{1+e^{-x}}\right)=(1-\sigma(x)) \sigma(x)
\end{aligned}
$$

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## Think, pair, share

## Patterns in backward flow

add gate: gradient distributor


## Think, pair, share

## Patterns in backward flow

add gate: gradient distributor
Q: What is a max gate?


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April 13, 2017

## Think, pair, share

## Patterns in backward flow

add gate: gradient distributor max gate: gradient router


## Think, pair, share

## Patterns in backward flow

add gate: gradient distributor max gate: gradient router
Q: What is a mul gate?


## Think, pair, share

## Patterns in backward flow

add gate: gradient distributor max gate: gradient router
mul gate: gradient switcher


## Merging gradients

## Gradients add at branches



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## Handing vector variables

Gradients for vectorized code


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April 13, 2017

## Handing vector variables

## Vectorized operations



## Handing vector variables

Vectorized operations

4096-d
input vector
Q: what is the size of the Jacobian matrix?


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## Handing vector variables

Vectorized operations

4096-d
input vector
Q: what is the size of the Jacobian matrix? [4096 x 4096!]

$$
\frac{\partial L}{\partial x}=\frac{\partial f}{\partial x} \frac{\partial L}{\partial f}
$$

Jacobian matrix

4096-d
output vector

## Handing vector variables

## Vectorized operations



## Handing vector variables

## Vectorized operations



## Handing vector variables

A vectorized example: $f(x, W)=\|W \cdot x\|^{2}=\sum_{i=1}^{n}(W \cdot x)_{i}^{2}$

## Handing vector variables

A vectorized example: $f(x, W)=\|W \cdot x\|^{2}=\sum_{i=1}^{n}(W \cdot x)_{i}^{2}$

$$
\in \mathbb{R}^{n} \in \mathbb{R}^{n \times n}
$$

## Handing vector variables

A vectorized example: $f(x, W)=\|W \cdot x\|^{2}=\sum_{i=1}^{n}(W \cdot x)_{i}^{2}$


## Handing vector variables

A vectorized example: $f(x, W)=\|W \cdot x\|^{2}=\sum_{i=1}^{n}(W \cdot x)_{i}^{2}$
$\left[\begin{array}{cc}0.1 & 0.5 \\ -0.3 & 0.8\end{array}\right] \mathbf{W}$

$q=W \cdot x=\left(\begin{array}{c}W_{1,1} x_{1}+\cdots+W_{1, n} x_{n} \\ \vdots \\ W_{n, 1} x_{1}+\cdots+W_{n, n} x_{n}\end{array}\right)$
$f(q)=\|q\|^{2}=q_{1}^{2}+\cdots+q_{n}^{2}$
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## Handing vector variables

A vectorized example: $f(x, W)=\|W \cdot x\|^{2}=\sum_{i=1}^{n}(W \cdot x)_{i}^{2}$
$\left[\begin{array}{cc}0.1 & 0.5 \\ -0.3 & 0.8\end{array}\right] \mathbf{W}$

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## Handing vector variables

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$\left[\begin{array}{cc}0.1 & 0.5 \\ -0.3 & 0.8\end{array}\right] \mathbf{W}$

$q=W \cdot x=\left(\begin{array}{c}W_{1,1} x_{1}+\cdots+W_{1, n} x_{n} \\ \vdots \\ W_{n, 1} x_{1}+\cdots+W_{n, n} x_{n}\end{array}\right)$
$f(q)=\|q\|^{2}=q_{1}^{2}+\cdots+q_{n}^{2}$
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## Handing vector variables

A vectorized example: $f(x, W)=\|W \cdot x\|^{2}=\sum_{i=1}^{n}(W \cdot x)_{i}^{2}$
$\left.\begin{array}{cc}0.1 & 0.5 \\ -0.3 & 0.8\end{array}\right] \mathbf{W}$

$q=W \cdot x=\left(\begin{array}{c}W_{1,1} x_{1}+\cdots+W_{1, n} x_{n} \\ \vdots \\ W_{n, 1} x_{1}+\cdots+W_{n, n} x_{n}\end{array}\right) \quad \frac{\partial f}{\partial q_{i}}=2 q_{i}$
$f(q)=\|q\|^{2}=q_{1}^{2}+\cdots+q_{n}^{2}$
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## Handing vector variables

A vectorized example: $f(x, W)=\|W \cdot x\|^{2}=\sum_{i=1}^{n}(W \cdot x)_{i}^{2}$
$\left[\begin{array}{cc}0.1 & 0.5 \\ -0.3 & 0.8\end{array}\right] \mathbf{W}$

$q=W \cdot x=\left(\begin{array}{c}W_{1,1} x_{1}+\cdots+W_{1, n} x_{n} \\ \vdots \\ W_{n, 1} x_{1}+\cdots+W_{n, n} x_{n}\end{array}\right) \quad \frac{\partial f}{\partial q_{i}}=2 q_{i}$
$f(q)=\|q\|^{2}=q_{1}^{2}+\cdots+q_{n}^{2}$
Fei-Fei Li \& Justin Johnson \& Serena Yeung Lecture 4-65

## Handing vector variables

A vectorized example: $f(x, W)=\|W \cdot x\|^{2}=\sum_{i=1}^{n}(W \cdot x)_{i}^{2}$
$\left.\begin{array}{cc}0.1 & 0.5 \\ -0.3 & 0.8\end{array}\right] \mathbf{W}$
$\left[\begin{array}{l}0.2 \\ 0.4\end{array}\right] \xrightarrow{\left(2 q_{i, j}\right.}$

$$
\begin{aligned}
& q=W \cdot x=\left(\begin{array}{c}
W_{1,1} x_{1}+\cdots+W_{1, n} x_{n} \\
\vdots \\
W_{n, 1} x_{1}+\cdots+W_{n, n} x_{n}
\end{array}\right) \\
& f(q)=\|q\|^{2}=q_{1}^{2}+\cdots+q_{n}^{2}
\end{aligned}
$$

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## Handing vector variables

A vectorized example: $f(x, W)=\|W \cdot x\|^{2}=\sum_{i=1}^{n}(W \cdot x)_{i}^{2}$
$\left[\begin{array}{cc}0.1 & 0.5 \\ -0.3 & 0.8\end{array}\right] \mathbf{W}$


$$
\begin{aligned}
q=W \cdot x=\left(\begin{array}{c}
W_{1,1} x_{1}+\cdots+W_{1, n} x_{n} \\
\vdots \\
W_{n, 1} x_{1}+\cdots+W_{n, n} x_{n}
\end{array}\right) & \left.\begin{array}{rl}
\frac{\partial f}{\partial W_{i, j}} & =\sum_{k} \frac{\partial f}{\partial q_{k}} \frac{\partial q_{k}}{\partial W_{i, j}} \\
& =\sum_{k}\left(2 q_{k}\right)\left(\mathbf{1}_{k=i} x_{j}\right) \\
f(q)=\|q\|^{2}=q_{1}^{2}+\cdots+q_{n}^{2} &
\end{array}\right)=2 q_{i} x_{j}
\end{aligned}
$$

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## Handing vector variables

A vectorized example: $f(x, W)=\|W \cdot x\|^{2}=\sum_{i=1}^{n}(W \cdot x)_{i}^{2}$

$$
\left.\begin{array}{l}
{\left[\begin{array}{cc}
0.1 & 0.5 \\
-0.3 & 0.8
\end{array}\right] \mathrm{W}} \\
{\left[\begin{array}{cc}
0.088 & 0.176 \\
0.104 & 0.208
\end{array}\right]} \\
0.4
\end{array}\right] \text { ( }
$$

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## Handing vector variables

A vectorized example: $f(x, W)=\|W \cdot x\|^{2}=\sum_{i=1}^{n}(W \cdot x)_{i}^{2}$

$$
\begin{aligned}
& \begin{array}{cc}
{\left[\begin{array}{cc}
0.1 & 0.5 \\
-0.3 & 0.8
\end{array}\right] \mathrm{W}} & \nabla_{W} f=2 q \cdot x^{T}
\end{array} \\
& 0.104 \quad 0.208 \\
& {\left[\begin{array}{l}
0.2 \\
0.4
\end{array}\right] \mathrm{x} \ll\left[\begin{array}{l}
0.26
\end{array}\right]} \\
& q=W \cdot x=\binom{W_{1,1} x_{1}+\cdots+W_{1, n} x_{n}}{:} \quad \frac{\partial f}{\partial W_{i, j}}=\sum_{k} \frac{\partial f}{\partial q_{k}} \frac{\partial q_{k}}{\partial W_{i, j}} \\
& =\sum_{k}\left(2 q_{k}\right)\left(\mathbf{1}_{k=i} x_{j}\right) \\
& f(q)=\|q\|^{2}=q_{1}^{2}+\cdots+q_{n}^{2} \\
& =2_{q_{i}}^{k} x_{j}
\end{aligned}
$$

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## Handing vector variables

A vectorized example: $f(x, W)=\|W \cdot x\|^{2}=\sum_{i=1}^{n}(W \cdot x)_{i}^{2}$

$$
\begin{aligned}
& {\left[\begin{array}{cc}
0.1 & 0.5 \\
-0.3 & 0.8
\end{array}\right]} \\
& {\left[\begin{array}{cc}
0.088 & 0.176
\end{array}\right] \mathrm{W}} \\
& 0.104 \quad 0.208 \\
& \nabla_{W} f=2 q \cdot x^{T} \\
& \xrightarrow\left[(\text { L2 }]{\frac{\partial q_{k}}{\partial W_{i, j}}=\stackrel{1.00}{0.116}} \stackrel{\mathbf{1}_{k=i} x_{j}}{1}\right. \\
& \text { Always check: The } \\
& \text { gradient with } \\
& \text { respect to a variable } \\
& \text { should have the } \\
& \text { same shape as the } \\
& \text { variable } \\
& q=W \cdot x=\left(\begin{array}{c}
W_{1,1} x_{1}+\cdots+W_{1, n} x_{n} \\
\vdots \\
W_{n, 1} x_{1}+\cdots+W_{n, n} x_{n}
\end{array}\right) \\
& f(q)=\|q\|^{2}=q_{1}^{2}+\cdots+q_{n}^{2} \\
& \frac{\partial f}{\partial W_{i, j}}=\sum_{k} \frac{\partial f}{\partial q_{k}} \frac{\partial q_{k}}{\partial W_{i, j}} \\
& =\sum\left(2 q_{k}\right)\left(\mathbf{1}_{k=i} x_{j}\right) \\
& =2_{q_{i}}^{k} x_{j}
\end{aligned}
$$

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## Handing vector variables

A vectorized example: $f(x, W)=\|W \cdot x\|^{2}=\sum_{i=1}^{n}(W \cdot x)_{i}^{2}$


$$
\frac{\partial q_{k}}{\partial x_{i}}=W_{k, i}
$$

$q=W \cdot x=\left(\begin{array}{c}W_{1,1} x_{1}+\cdots+W_{1, n} x_{n} \\ \vdots \\ W_{n, 1} x_{1}+\cdots+W_{n, n} x_{n}\end{array}\right)$
$f(q)=\|q\|^{2}=q_{1}^{2}+\cdots+q_{n}^{2}$
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## Handing vector variables

A vectorized example: $f(x, W)=\|W \cdot x\|^{2}=\sum_{i=1}^{n}(W \cdot x)_{i}^{2}$


$$
\begin{aligned}
\frac{\partial q_{k}}{\partial x_{i}} & =W_{k, i} \\
\frac{\partial f}{\partial x_{i}} & =\sum_{k} \frac{\partial f}{\partial q_{k}} \frac{\partial q_{k}}{\partial x_{i}} \\
& =\sum_{k} 2 q_{k} W_{k, i}
\end{aligned}
$$

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## Handing vector variables

A vectorized example: $f(x, W)=\|W \cdot x\|^{2}=\sum_{i=1}^{n}(W \cdot x)_{i}^{2}$

$$
\begin{aligned}
& \frac{\partial q_{k}}{\partial x_{i}}=W_{k, i} \\
& \frac{\partial f}{\partial x_{i}}=\sum_{k} \frac{\partial f}{\partial q_{k}} \frac{\partial q_{k}}{\partial x_{i}} \\
& =\sum_{k} 2 q_{k} W_{k, i}
\end{aligned}
$$

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## Handing vector variables

A vectorized example: $f(x, W)=\|W \cdot x\|^{2}=\sum_{i=1}^{n}(W \cdot x)_{i}^{2}$

$$
\begin{aligned}
& \frac{\partial q_{k}}{\partial x_{i}}=W_{k, i} \\
& \frac{\partial f}{\partial x_{i}}=\sum_{k} \frac{\partial f}{\partial q_{k}} \frac{\partial q_{k}}{\partial x_{i}} \\
& =\sum_{k} 2 q_{k} W_{k, i}
\end{aligned}
$$

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## Implementation

## Modularized implementation: forward / backward API

## Graph (or Net) object (rough psuedo code)



```
class ComputationalGraph(object):
    #...
    def forward(inputs):
        # 1. [pass inputs to input gates...]
        # 2. forward the computational graph:
        for gate in self.graph.nodes_topologically_sorted():
        gate.forward()
        return loss # the final gate in the graph outputs the loss
    def backward():
        for gate in reversed(self.graph.nodes_topologically_sorted()):
        gate.backward() # little piece of backprop (chain rule applied)
    return inputs_gradients
```


## Implementation

## Modularized implementation: forward / backward API



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## Implementation

## Modularized implementation: forward / backward API

```
class MultiplyGate(object):
```

class MultiplyGate(object):
def forward(x,y):
def forward(x,y):
z = x*y
z = x*y
self.x = x \# must keep these around!
self.x = x \# must keep these around!
self.y = y
self.y = y
return z
return z
def backward(dz):
def backward(dz):
dx = self.y * dz \# [dz/dx * dL/dz]
dx = self.y * dz \# [dz/dx * dL/dz]
dy = self.x * dz \# [dz/dy * dL/dz]
dy = self.x * dz \# [dz/dy * dL/dz]
return [dx, dy]

```
    return [dx, dy]
```


## Remark of BP

- During the forward pass, each computing unit will evaluate the output and also the corresponding local derivatives of the output w.r.t. the inputs


## Remark of BP

- During the forward pass, each computing unit will evaluate the output and also the corresponding local derivatives of the output w.r.t. the inputs
- During the backward pass, the local derivatives and the evaluated outputs will be "consumed" to compute the overall derivatives


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- For a large network, there can be a large spike of memory consumption during the forward pass


## Remark of BP

- During the forward pass, each computing unit will evaluate the output and also the corresponding local derivatives of the output w.r.t. the inputs
- During the backward pass, the local derivatives and the evaluated outputs will be "consumed" to compute the overall derivatives
- For a large network, there can be a large spike of memory consumption during the forward pass
- Note that BP only computes the gradients. It does not perform the optimization. Sometimes you may hear people said that they trained their networks with BP. What they said was not literally right. We will discuss more on optimizer later today


## Activation functions

## Activation Functions



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## Activation functions

## Activation Functions

## Leaky ReLU $\max (0.1 x, x)$

Sigmoid
$\sigma(x)=1 /\left(1+e^{-x}\right)$

$\boldsymbol{\operatorname { t a n h }} \boldsymbol{\operatorname { t a n h }}(\mathrm{x})$

ReLU $\max (0, x)$


Maxout $\max \left(w_{1}^{T} x+b_{1}, w_{2}^{T} x+b_{2}\right)$

ELU

$$
f(x)= \begin{cases}x & \text { if } x>0 \\ \alpha(\exp (x)-1) & \text { if } x \leq 0\end{cases}
$$



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## Activation functions

## Activation Functions



## Sigmoid

$$
\sigma(x)=1 /\left(1+e^{-x}\right)
$$

- Squashes numbers to range $[0,1]$
- Historically popular since they have nice interpretation as a saturating "firing rate" of a neuron


## Activation functions

## Activation Functions



Sigmoid

$$
\sigma(x)=1 /\left(1+e^{-x}\right)
$$

- Squashes numbers to range $[0,1]$
- Historically popular since they have nice interpretation as a saturating "firing rate" of a neuron

3 problems:

1. Saturated neurons "kill" the gradients

## Activation functions



What happens when $x=-10$ ?
What happens when $x=0$ ?
What happens when $x=10$ ?

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## Activation functions

## Activation Functions



Sigmoid

$$
\sigma(x)=1 /\left(1+e^{-x}\right)
$$

- Squashes numbers to range $[0,1]$
- Historically popular since they have nice interpretation as a saturating "firing rate" of a neuron

3 problems:

1. Saturated neurons "kill" the gradients
2. Sigmoid outputs are not zerocentered

## Activation functions

Consider what happens when the input to a neuron (x) is always positive:


What can we say about the gradients on w?

## Activation functions

Consider what happens when the input to a neuron is always positive...

$$
f\left(\sum_{i} w_{i} x_{i}+b\right)
$$

allowed
gradient
update
directions

> allowed
> gradient
> update
> directions
hypothetical optimal w vector

What can we say about the gradients on $\mathbf{w}$ ?
Always all positive or all negative :(
(this is also why you want zero-mean data!)
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## Activation functions

## Activation Functions



Sigmoid

$$
\sigma(x)=1 /\left(1+e^{-x}\right)
$$

- Squashes numbers to range $[0,1]$
- Historically popular since they have nice interpretation as a saturating "firing rate" of a neuron

3 problems:

1. Saturated neurons "kill" the gradients
2. Sigmoid outputs are not zerocentered
3. $\exp ()$ is a bit compute expensive

## Activation functions

## Activation Functions



- Squashes numbers to range [-1,1]
- zero centered (nice)
- still kills gradients when saturated :(


## $\tanh (x)$

[LeCun et al., 1991]
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## Activation functions

## Activation Functions



ReLU
(Rectified Linear Unit)
[Krizhevsky et al., 2012]

## Activation functions

## Activation Functions



ReLU
(Rectified Linear Unit)

- Computes $\mathbf{f}(\mathbf{x})=\max (\mathbf{0 , x})$
- Does not saturate (in +region)
- Very computationally efficient
- Converges much faster than sigmoid/tanh in practice (e.g. 6x)
- Not zero-centered output
- An annoyance:
hint: what is the gradient when $x<0$ ?

[^5]
## Activation functions




What happens when $x=-10$ ?
What happens when $x=0$ ?
What happens when $x=10 ?$

## Activation functions



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## Activation functions



## Activation functions

## Activation Functions

[Mass et al., 2013] [He et al., 2015]

- Does not saturate
- Computationally efficient
- Converges much faster than sigmoid/tanh in practice! (e.g. 6x)
- will not "die".

Leaky ReLU
$f(x)=\max (0.01 x, x)$

## Activation functions

## Activation Functions

[Mass et al., 2013] [He et al., 2015]

- Does not saturate
- Computationally efficient
- Converges much faster than sigmoid/tanh in practice! (e.g. 6x)
- will not "die".


## Parametric Rectifier (PReLU)

$$
f(x)=\max (\alpha x, x)
$$

backprop into \alpha (parameter)

Leaky ReLU
$f(x)=\max (0.01 x, x)$

Lecture 5-44
20 Jan 2016

## Activation functions

## Activation Functions

## Exponential Linear Units (ELU)



- All benefits of ReLU
- Does not die
- Closer to zero mean outputs
- Computation requires $\exp ()$

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## Activation functions

Maxout "Neurons" [Goodfellow et al., 2013]

- Try to generalize ReLU and leaky ReLU

$$
\max \left(\mathbf{w}_{1}^{T} \mathbf{x}+b_{1}, \mathbf{w}_{2}^{T} \mathbf{x}+b_{2}\right)
$$

## Activation functions

Maxout "Neurons" [Goodfellow et al., 2013]

- Try to generalize ReLU and leaky ReLU

$$
\max \left(\mathbf{w}_{1}^{T} \mathbf{x}+b_{1}, \mathbf{w}_{2}^{T} \mathbf{x}+b_{2}\right)
$$

Pros

- Linear regime
- Does not saturate
- Does not die


## Activation functions

Maxout "Neurons" [Goodfellow et al., 2013]

- Try to generalize ReLU and leaky ReLU

$$
\max \left(\mathbf{w}_{1}^{T} \mathbf{x}+b_{1}, \mathbf{w}_{2}^{T} \mathbf{x}+b_{2}\right)
$$

Pros

- Linear regime
- Does not saturate
- Does not die

Cons

- Double amount of parameters


## Activation functions

## TLDR: In practice:

- Use ReLU. Be careful with your learning rates
- Try out Leaky ReLU / Maxout / ELU
- Try out tanh but don't expect much
- Don't use sigmoid


## Input preprocessing

## Step 1: Preprocess the data


(Assume $\mathrm{X}[\mathrm{NxD}]$ is data matrix, each example in a row)

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## Input preprocessing

## Step 1: Preprocess the data

In practice, you may also see PCA and Whitening of the data


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## Input preprocessing

## TLDR: In practice for Images: center only

e.g. consider CIFAR-10 example with [32,32,3] images

- Subtract the mean image (e.g. AlexNet) (mean image $=[32,32,3]$ array)
- Subtract per-channel mean (e.g. VGGNet) (mean along each channel $=3$ numbers)

Not common to normalize variance, to do PCA or whitening

$$
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$$

## Weight initialization

- $Q$ : what happens when $W=0$ init is used?


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## Weight initialization

- First idea: Small random numbers
(gaussian with zero mean and 1e-2 standard deviation)

$$
W=0.01^{*} \text { np. random. } \operatorname{randn}(\mathrm{D}, \mathrm{H})
$$

## Weight initialization

- First idea: Small random numbers
(gaussian with zero mean and 1e-2 standard deviation)

$$
W=0.01^{*} \mathrm{np} . \text { random. } \operatorname{randn}(\mathrm{D}, \mathrm{H})
$$

Works ~okay for small networks, but can lead to non-homogeneous distributions of activations across the layers of a network.

## Weight initialization

## Lets look at some activation statistics

E.g. 10-layer net with 500 neurons on each layer, using tanh nonlinearities, and initializing as described in last slide.

```
# assume some unit gaussian 10-D input data
D = np.random. randn(1000, 500)
hidden_layer_sizes = [500]*10
nonlinearities = ['tanh']*len(hidden_layer_sizes)
act = {'relu':lambda x:np.maximum( }0,x\mathrm{ ), 'tanh':lambda x:np.tanh (x)}
Hs}={
for i in xrange(len(hidden layer_sizes)):
    x=D if i== else Hs[i-1] # Input at this layer
    fan in = X.shape[1]
    fan_out = hidden layer_sizes[i]
    W = np.random.randn(fan_in, fan_out) * 0.01 # layer initialization
    H=np.dot(X, W) # matrix multiply
    H}=\operatorname{act[nonlinearities[i]](H) # nonlinearity
    Hs[i] = H# cache result on this layer
```

\# look at distributions at each layer
print 'input layer had mean \%f and std \%f' \% (np.mean(D), np.std(D))
layer means $=[n p$. mean $(\mathrm{H})$ for $\mathrm{i}, \mathrm{H}$ in Hs.iteritems()]
layer_stds $=$ [np.std $(H)$ for $i, H$ in Hs.iteritems()]
layer stds $=$ [np.std $(\mathrm{H})$ fo
for i , H in Hs . iteritems():
print 'hidden layer \%d had mean sf and std \%f' \% ( $i+1$, layer_means[i], layer_stds[i])
\# plot the means and standard deviations
plt.figure()
plt.subplot(121)
plt.plot(Hs.keys(), layer_means, ob-1)
plt.title('layer mean')
plt.subplot(122)
plt.plot(Hs.keys(), layer stds, 'or-')
plt.title('layer std')
\# plot the raw distributions
plt.figure()
for $\mathrm{i}, \mathrm{H}$ in $\mathrm{H} s$.iteritems():
plt.subplot(1,len(Hs),1+1)
plt.hist(H.ravel(), 30 , range $=(-1,1))$

## Weight initialization

input layer had mean 0.000927 and std 0.998388
hidden layer 1 had mean -0.000117 and std 0.213081 hidden Layer 1 had mean -0.000117 and std 0.213081 hidden layer 2 had mean -0.000001 and std 0.047551 hidden layer 3 had mean $-\theta .000002$ and std $\theta .010636$
hidden layer 4 had mean 0.000001 and std 0.002378 hidden layer 4 had mean 0.000001 and std 0.002378 hidden layer 5 had mean 0.000002 and std 0.000532 hidden layer 6 had mean -0.000000 and std 0.000119 hidden layer 7 had mean 0.000000 and std 0.000026 hidden layer 8 had mean -0.000000 and std 0.000006 hidden layer 9 had mean 0.000000 and std 0.000001 hidden layer 10 had mean -0.000000 and std 0.000000


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## Weight initialization

input layer had mean 0.000927 and std 0.998388
hidden layer 1 had mean -0.000117 and std 0.213081 hidden layer 2 had mean -0.000001 and std 0.047551 hidden layer 3 had mean $-\theta .000002$ and std $\theta .010631$ hidden layer 4 had mean 0.000001 and std 0.002378 hidden layer 5 had mean 0.000002 and std 0.000532 hidden layer 6 had mean -0.000000 and std 0.000119 hidden layer 7 had mean 0.000000 and std 0.000026 hidden layer 8 had mean -0.000000 and std 0.000006 hidden layer 9 had mean 0.000000 and std 0.000001 hidden layer 10 had mean -0.000000 and std 0.000000


## All activations become zero!

Hint: think about backward pass for a $W^{*} X$ gate.

## Weight initialization


input layer had mean 0.001800 and std 1.001311 hidden layer 1 had mean -0.000430 and std 0.981879 hidden layer 2 had mean -0.000849 and std 0.981649 hidden layer 3 had mean $0.0 日 0565$ and std $\theta .981601$ hidden layer 4 had mean 0.000483 and std 0.981755 hidden layer 5 had mean -0.000682 and std 0.981614 hidden layer 6 had mean -0.000401 and std 0.981560 hidden layer 7 had mean -0.000237 and std 0.981520 hidden layer 8 had mean -0.000448 and std 0.981913 hidden layer 9 had mean -0.000899 and std 0.981728 hidden layer 10 had mean 0.000584 and std 0.981736




## Almost all neurons completely saturated, either -1 and 1. Gradients will be all zero.

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## Variance calibration for linear layer

Assume linear activation and zero-mean weights and inputs. And number of inputs is $n$. Then,

$$
\operatorname{Var}(y)=\operatorname{Var}\left(\sum_{i}^{n} w_{i} x_{i}\right)=\sum_{i}^{n} \operatorname{Var}\left(w_{i} x_{i}\right)
$$

## Variance calibration for linear layer

Assume linear activation and zero-mean weights and inputs. And number of inputs is $n$. Then,

$$
\begin{aligned}
\operatorname{Var}(y) & =\operatorname{Var}\left(\sum_{i}^{n} w_{i} x_{i}\right)=\sum_{i}^{n} \operatorname{Var}\left(w_{i} x_{i}\right) \\
& =\sum_{i}^{n}\left[E\left(w_{i}\right)\right]^{2} \operatorname{Var}\left(x_{i}\right)+E\left[\left(x_{i}\right)\right]^{2} \operatorname{Var}\left(w_{i}\right)+\operatorname{Var}\left(x_{i}\right) \operatorname{Var}\left(w_{i}\right)
\end{aligned}
$$

# $\operatorname{Var}(X Y)=$ $E[X]^{2} \operatorname{Var}(X)+E[Y]^{2} \operatorname{Var}(Y)+\operatorname{Var}(X) \operatorname{Var}(Y)$ 

$$
\operatorname{Var}(X Y)=E\left[(X Y)^{2}\right]-E[X Y]^{2}
$$

# $\operatorname{Var}(X Y)=$ $E[X]^{2} \operatorname{Var}(X)+E[Y]^{2} \operatorname{Var}(Y)+\operatorname{Var}(X) \operatorname{Var}(Y)$ 

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\begin{aligned}
\operatorname{Var}(X Y) & =E\left[(X Y)^{2}\right]-E[X Y]^{2} \\
& =E\left[X^{2}\right] E\left[Y^{2}\right]-E[X]^{2} E[Y]^{2}
\end{aligned}
$$

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& =E\left[X^{2}\right] E\left[Y^{2}\right]-E[X]^{2} E[Y]^{2}
\end{aligned}
$$

$$
\begin{aligned}
& \operatorname{Var}(X) \operatorname{Var}(Y) \\
= & \left(E\left[X^{2}\right]-E[X]^{2}\right)\left(E\left[Y^{2}\right]-E[Y]^{2}\right)
\end{aligned}
$$

## $\operatorname{Var}(X Y)=$ $E[X]^{2} \operatorname{Var}(X)+E[Y]^{2} \operatorname{Var}(Y)+\operatorname{Var}(X) \operatorname{Var}(Y)$

$$
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\end{aligned}
$$

$$
\begin{aligned}
& \operatorname{Var}(X) \operatorname{Var}(Y) \\
= & \left(E\left[X^{2}\right]-E[X]^{2}\right)\left(E\left[Y^{2}\right]-E[Y]^{2}\right) \\
= & E\left[X^{2}\right] E\left[Y^{2}\right]-E[X]^{2} E\left[Y^{2}\right]-E\left[X^{2}\right] E[Y]^{2}+E[X]^{2} E[Y]^{2}
\end{aligned}
$$

## $\operatorname{Var}(X Y)=$ $E[X]^{2} \operatorname{Var}(X)+E[Y]^{2} \operatorname{Var}(Y)+\operatorname{Var}(X) \operatorname{Var}(Y)$

$$
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& =E\left[X^{2}\right] E\left[Y^{2}\right]-E[X]^{2} E[Y]^{2}
\end{aligned}
$$

$$
\begin{aligned}
& \operatorname{Var}(X) \operatorname{Var}(Y) \\
= & \left(E\left[X^{2}\right]-E[X]^{2}\right)\left(E\left[Y^{2}\right]-E[Y]^{2}\right) \\
= & E\left[X^{2}\right] E\left[Y^{2}\right]-E[X]^{2} E\left[Y^{2}\right]-E\left[X^{2}\right] E[Y]^{2}+E[X]^{2} E[Y]^{2} \\
= & E\left[X^{2}\right] E\left[Y^{2}\right]-E[X]^{2}\left(E\left[Y^{2}\right]-E[Y]^{2}\right) \\
& E[Y]^{2}\left(E\left[X^{2}\right]-E[X]^{2}\right)-E[X]^{2} E[Y]^{2}
\end{aligned}
$$

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\begin{aligned}
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& =E\left[X^{2}\right] E\left[Y^{2}\right]-E[X]^{2} E[Y]^{2}
\end{aligned}
$$

$$
\begin{aligned}
& \operatorname{Var}(X) \operatorname{Var}(Y) \\
= & \left(E\left[X^{2}\right]-E[X]^{2}\right)\left(E\left[Y^{2}\right]-E[Y]^{2}\right) \\
= & E\left[X^{2}\right] E\left[Y^{2}\right]-E[X]^{2} E\left[Y^{2}\right]-E\left[X^{2}\right] E[Y]^{2}+E[X]^{2} E[Y]^{2} \\
= & E\left[X^{2}\right] E\left[Y^{2}\right]-E[X]^{2}\left(E\left[Y^{2}\right]-E[Y]^{2}\right) \\
& E[Y]^{2}\left(E\left[X^{2}\right]-E[X]^{2}\right)-E[X]^{2} E[Y]^{2} \\
= & \operatorname{Var}(X Y)-E[X]^{2} \operatorname{Var}(Y)-E[Y]^{2} \operatorname{Var}(X)
\end{aligned}
$$

## Variance calibration for linear layer

Assume linear activation and zero-mean weights and inputs. And number of inputs is $n$. Then,

$$
\begin{aligned}
\operatorname{Var}(y) & =\operatorname{Var}\left(\sum_{i}^{n} w_{i} x_{i}\right)=\sum_{i}^{n} \operatorname{Var}\left(w_{i} x_{i}\right) \\
& =\sum_{i}^{n}\left[E\left(w_{i}\right)\right]^{2} \operatorname{Var}\left(x_{i}\right)+E\left[\left(x_{i}\right)\right]^{2} \operatorname{Var}\left(w_{i}\right)+\operatorname{Var}\left(x_{i}\right) \operatorname{Var}\left(w_{i}\right)
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& =\sum_{i}^{n} \operatorname{Var}\left(x_{i}\right) \operatorname{Var}\left(w_{i}\right) \\
& =(n \operatorname{Var}(w)) \operatorname{Var}(x)
\end{aligned}
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& =\sum_{i}^{n} \operatorname{Var}\left(x_{i}\right) \operatorname{Var}\left(w_{i}\right) \\
& =(n \operatorname{Var}(w)) \operatorname{Var}(x)
\end{aligned}
$$

Thus, output will have same variance as input if $n \operatorname{Var}(w)=1$

## Weight initialization

input layer had mean 0.001800 and std 1.001311 hidden layer 1 had mean 0.001198 and std 0.627953 hidden layer 2 had mean -0.000175 and std 0.486051 hidden layer 3 had mean 0.000055 and 5 td 0.407723 hidden layer 4 had mean - 0.000306 and std 0.357108 hidden layer 5 had mean 0.000142 and std 0.320917 hidden layer 6 had mean -0.000389 and std 0.292116 hidden layer 6 had mean -0.000389 and std 0.292116 hidden layer 7 had mean -0.000228 and std 0.273387 hidden layer 9 had mean 0.000361 and std 0.239266 hidden layer 9 had mean 0.000361 and std 0.239266
hidden layer 10 had mean 0.000139 and std 0.228008

W = np. random. randn(fan in, fan out) / np.sqrt(fan in) \# layer initialization
"Xavier initialization"
[Glorot et al., 2010]


Reasonable initialization. (Mathematical derivation assumes linear activations)


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## Weight initialization

input layer had mean $0.0005 \theta 1$ and std 0.999444 hidden layer 1 had mean 0.398623 and std 0.582273 hidden layer 2 had mean 0.272352 and std 0.403795 hidden layer 3 had mean 0.186076 and std 0.276912 hidden layer 4 had mean 0.136442 and std 0.198685 hidden layer 5 had mean 0.099568 and std 0.140299 hidden layer 6 had mean 0.072234 and std 0.103280 hidden layer 7 had mean 0.049775 and std 0.072748 hidden layer 8 had mean 0.035138 and std 0.051572 hidden layer 9 had mean 0.025404 and std 0.038583 hidden layer 10 had mean 0.018408 and std 0.026076

W = np. random. randn(fan in, fan out) / np.sqrt(fan in) \# layer initialization

## but when using the ReLU nonlinearity it breaks.

## Variance calibration for ReLU

Note that it doesn't work when the activation layer is ReLU. But... ${ }^{1}$

$$
\begin{aligned}
\operatorname{Var}\left(y^{(l)}\right) & =\operatorname{Var}\left(\sum_{i}^{n} w_{i}^{(l)} x_{i}^{(l)}\right)=\sum_{i}^{n} \operatorname{Var}\left(w_{i}^{(l)} x_{i}^{(l)}\right)=n \operatorname{Var}\left(w^{(l)} x^{(l)}\right) \\
& =n E\left(w^{(l)}\right)^{2} \operatorname{Var}\left(x^{(l)}\right)+n E\left(x^{(l)}\right)^{2} \operatorname{Var}\left(w^{(l)}\right)+n \operatorname{Var}\left(x^{(l)}\right) \operatorname{Var}\left(w^{(l)}\right)
\end{aligned}
$$

${ }^{1}$ Note that $y^{(l)}$ now denotes the sum of input before going through the activation function.

## Variance calibration for ReLU

Note that it doesn't work when the activation layer is ReLU. But... ${ }^{1}$

$$
\begin{aligned}
\operatorname{Var}\left(y^{(l)}\right) & =\operatorname{Var}\left(\sum_{i}^{n} w_{i}^{(l)} x_{i}^{(l)}\right)=\sum_{i}^{n} \operatorname{Var}\left(w_{i}^{(l)} x_{i}^{(l)}\right)=n \operatorname{Var}\left(w^{(l)} x^{(l)}\right) \\
& =n E\left(w^{(l)}\right)^{2} \operatorname{Var}\left(x^{(l)}\right)+n E\left(x^{(l)}\right)^{2} \operatorname{Var}\left(w^{(l)}\right)+n \operatorname{Var}\left(x^{(l)}\right) \operatorname{Var}\left(w^{(l)}\right) \\
& =n E\left(x^{(l)}\right)^{2} \operatorname{Var}\left(w^{(l)}\right)+n \operatorname{Var}\left(x^{(l)}\right) \operatorname{Var}\left(w^{(l)}\right)
\end{aligned}
$$

[^6]
## Variance calibration for ReLU

Note that it doesn't work when the activation layer is ReLU. But... ${ }^{1}$

$$
\begin{aligned}
\operatorname{Var}\left(y^{(l)}\right) & =\operatorname{Var}\left(\sum_{i}^{n} w_{i}^{(l)} x_{i}^{(l)}\right)=\sum_{i}^{n} \operatorname{Var}\left(w_{i}^{(l)} x_{i}^{(l)}\right)=n \operatorname{Var}\left(w^{(l)} x^{(l)}\right) \\
& =n E\left(w^{(l)}\right)^{2} \operatorname{Var}\left(x^{(l)}\right)+n E\left(x^{(l)}\right)^{2} \operatorname{Var}\left(w^{(l)}\right)+n \operatorname{Var}\left(x^{(l)}\right) \operatorname{Var}\left(w^{(l)}\right) \\
& =n E\left(x^{(l)}\right)^{2} \operatorname{Var}\left(w^{(l)}\right)+n \operatorname{Var}\left(x^{(l)}\right) \operatorname{Var}\left(w^{(l)}\right) \\
& =n E\left(\left(x^{(l)}\right)^{2}\right) \operatorname{Var}\left(w^{(l)}\right)
\end{aligned}
$$

[^7]
## Variance calibration for ReLU

Note that it doesn't work when the activation layer is ReLU. But... ${ }^{1}$

$$
\begin{aligned}
\operatorname{Var}\left(y^{(l)}\right) & =\operatorname{Var}\left(\sum_{i}^{n} w_{i}^{(l)} x_{i}^{(l)}\right)=\sum_{i}^{n} \operatorname{Var}\left(w_{i}^{(l)} x_{i}^{(l)}\right)=n \operatorname{Var}\left(w^{(l)} x^{(l)}\right) \\
& =n E\left(w^{(l)}\right)^{2} \operatorname{Var}\left(x^{(l)}\right)+n E\left(x^{(l)}\right)^{2} \operatorname{Var}\left(w^{(l)}\right)+n \operatorname{Var}\left(x^{(l)}\right) \operatorname{Var}\left(w^{(l)}\right) \\
& =n E\left(x^{(l)}\right)^{2} \operatorname{Var}\left(w^{(l)}\right)+n \operatorname{Var}\left(x^{(l)}\right) \operatorname{Var}\left(w^{(l)}\right) \\
& =n E\left(\left(x^{(l)}\right)^{2}\right) \operatorname{Var}\left(w^{(l)}\right) \\
& =n\left[\operatorname{Var}\left(y^{(l-1)}\right) / 2\right] \operatorname{Var}\left(w^{(l)}\right)=\left[\frac{n}{2} \operatorname{Var}\left(w^{(l)}\right)\right] \operatorname{Var}\left(y^{(l-1)}\right)
\end{aligned}
$$

Variance of $y$ conserved across a layer if $\frac{n}{2} \operatorname{Var}(w)=1$

[^8]
## Weight initialization

nput layer had mean 0.000501 and std 0.999444 hidden layer 1 had mean 0.562488 and std 0.82523 hidden layer 2 had mean 0.553614 and std 0.827835 hidden layer 3 had mean 0.545867 and std 0.813855 hidden layer 4 had mean 0.565396 and std 0.826902 hidden layer 5 had mean 0.547678 and std 0.834092 hidden layer 6 had mean 0.587103 and std 0.860035 hidden layer 7 had mean 0.596867 and std 0.870610 hidden layer 8 had mean 0.623214 and std 0.889348 hidden layer 9 had mean 0.567498 and std 0.845357 hidden layer 10 had mean 0.552531 and std 0.844523

```
W = np.random.randn(fan in, fan out) / np.sqrt(fan in/2) # layer initialization
```

He et al., 2015 (note additional /2)


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## Weight initialization

nput layer had mean 0.000501 and std 0.999444 hidden layer 1 had mean 0.562488 and std 0.82523 hidden layer 2 had mean 0.553614 and std 0.827835 hidden layer 3 had mean 0.545867 and std 0.813855 hidden layer 4 had mean 0.565396 and std 0.826902 hidden layer 5 had mean 0.547678 and std 0.834092 hidden layer 6 had mean 0.587103 and std 0.860035 hidden layer 7 had mean 0.596867 and std 0.870610 hidden layer 8 had mean 0.623214 and std 0.889348 hidden layer 9 had mean 0.567498 and std 0.845357 hidden layer 10 had mean 0.552531 and std 0.844523



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## Weight initialization

## Proper initialization is an active area of research...

Understanding the difficulty of training deep feedforward neural networks by Glorot and Bengio, 2010

Exact solutions to the nonlinear dynamics of learning in deep linear neural networks by Saxe et al, 2013

Random walk initialization for training very deep feedforward networks by Sussillo and Abbott, 2014

Delving deep into rectifiers: Surpassing human-level performance on ImageNet classification by He et al., 2015

Data-dependent Initializations of Convolutional Neural Networks by Krähenbühl et al., 2015
All you need is a good init, Mishkin and Matas, 2015

$$
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$$

## Batch normalization

## Batch Normalization

"you want unit gaussian activations? just make them so."
consider a batch of activations at some layer. To make each dimension unit gaussian, apply:

$$
\widehat{x}^{(k)}=\frac{x^{(k)}-\mathrm{E}\left[x^{(k)}\right]}{\sqrt{\operatorname{Var}\left[x^{(k)}\right]}}
$$

this is a vanilla differentiable function...

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## Batch normalization

## Batch Normalization

[loffe and Szegedy, 2015]
"you want unit gaussian activations?
just make them so."


1. compute the empirical mean and variance independently for each dimension.
[^9]
## Batch normalization

## Batch Normalization

[loffe and Szegedy, 2015]


Usually inserted after Fully
Connected / (or Convolutional, as we'll see soon) layers, and before nonlinearity.
 tanh layer?

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## Batch normalization

## Batch Normalization

[loffe and Szegedy, 2015]

Normalize:
$\widehat{x}^{(k)}=\frac{x^{(k)}-\mathrm{E}\left[x^{(k)}\right]}{\sqrt{\operatorname{Var}\left[x^{(k)}\right]}}$
And then allow the network to squash the range if it wants to:

$$
y^{(k)}=\gamma^{(k)} \widehat{x}^{(k)}+\beta^{(k)}
$$

Note, the network can learn:
$\gamma^{(k)}=\sqrt{\operatorname{Var}\left[x^{(k)}\right]}$
$\beta^{(k)}=\mathrm{E}\left[x^{(k)}\right]$
to recover the identity mapping.

## Batch normalization

## Batch Normalization

Input: Values of $x$ over a mini-batch: $\mathcal{B}=\left\{x_{1 \ldots m}\right\}$; Parameters to be learned: $\gamma, \beta$
Output: $\left\{y_{i}=\mathrm{BN}_{\gamma, \beta}\left(x_{i}\right)\right\}$

$$
\begin{array}{rlr}
\mu_{\mathcal{B}} & \leftarrow \frac{1}{m} \sum_{i=1}^{m} x_{i} & \text { // mini-batch mean } \\
\sigma_{\mathcal{B}}^{2} & \leftarrow \frac{1}{m} \sum_{i=1}^{m}\left(x_{i}-\mu_{\mathcal{B}}\right)^{2} & \text { // mini-batch variance } \\
\widehat{x}_{i} & \leftarrow \frac{x_{i}-\mu_{\mathcal{B}}}{\sqrt{\sigma_{\mathcal{B}}^{2}+\epsilon}} & \text { // normalize } \\
y_{i} & \leftarrow \gamma \widehat{x}_{i}+\beta \equiv \operatorname{BN}_{\gamma, \beta}\left(x_{i}\right) & \text { // scale and shift }
\end{array}
$$

- Improves gradient flow through the network
- Allows higher learning rates
- Reduces the strong dependence on initialization
- Acts as a form of regularization in a funny way, and slightly reduces the need for dropout, maybe


## Batch normalization

## Batch Normalization

Input: Values of $x$ over a mini-batch: $\mathcal{B}=\left\{x_{1 \ldots m}\right\}$; Parameters to be learned: $\gamma, \beta$
Output: $\left\{y_{i}=\mathrm{BN}_{\gamma, \beta}\left(x_{i}\right)\right\}$

$$
\begin{array}{rlr}
\mu_{\mathcal{B}} & \leftarrow \frac{1}{m} \sum_{i=1}^{m} x_{i} & \text { // mini-batch mean } \\
\sigma_{\mathcal{B}}^{2} & \leftarrow \frac{1}{m} \sum_{i=1}^{m}\left(x_{i}-\mu_{\mathcal{B}}\right)^{2} & \text { // mini-batch variance } \\
\widehat{x}_{i} & \leftarrow \frac{x_{i}-\mu_{\mathcal{B}}}{\sqrt{\sigma_{\mathcal{B}}^{2}+\epsilon}} & \text { // normalize } \\
y_{i} & \leftarrow \gamma \widehat{x}_{i}+\beta \equiv \operatorname{BN}_{\gamma, \beta}\left(x_{i}\right) & \text { // scale and shift }
\end{array}
$$

[loffe and Szegedy, 2015]

## Note: at test time BatchNorm layer functions differently:

The mean/std are not computed based on the batch. Instead, a single fixed empirical mean of activations during training is used.
(e.g. can be estimated during training with running averages)

## Reducing testing error

## How to improve single-model performance?




## Ensemble trick

## 1. Train multiple independent models <br> 2. At test time average their results

Enjoy 2\% extra performance

## Ensemble trick

## Fun Tips/Tricks:

- can also get a small boost from averaging multiple model checkpoints of a single model.


## Ensemble trick

## Model Ensembles: Tips and Tricks

Instead of training independent models, use multiple snapshots of a single model during training!


## Ensemble trick

## Model Ensembles: Tips and Tricks

Instead of training independent models, use multiple snapshots of a single model during training!


Loshchilov and Hutter, "SGDR: Stochastic gradient descent with restarts", arXiv 2016
Huang et al, "Snapshot ensembles: train 1, get M for free", ICLR 2017
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Cyclic learning rate schedules can make this work even better!

## Ensemble trick

## Model Ensembles: Tips and Tricks

Instead of using actual parameter vector, keep a moving average of the parameter vector and use that at test time (Polyak averaging)

```
while True:
    data_batch = dataset.sample_data_batch()
    loss = network.forward(data_batch)
    dx = network.backward()
    x += - learning_rate * dx
    x_test = 0.995*x_test + 0.005*x # use for test set
```


## Dropout

## Regularization: Dropout

"randomly set some neurons to zero in the forward pass"

(a) Standard Neural Net

(b) After applying dropout.

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## Dropout

$\mathrm{p}=0.5$ \# 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) $<\mathrm{p}$ \# first dropout mask
H1 *= U1 \# drop!
H2 = np.maximum(0, np. dot(W2, H1) + b2)
$\mathrm{U} 2=\mathrm{np}$. random. rand $(* \mathrm{H} 2$. shape $)<\mathrm{p} \#$ second dropout mask
H2 *= U2 \# drop!
out $=$ np.dot (W3, H2) + b3
\# backward pass: compute gradients... (not shown)
\# perform parameter update... (not shown)

Example forward pass with a 3layer network using dropout


## Dropout

## Regularization: Dropout

How can this possibly be a good idea?


Forces the network to have a redundant representation;
Prevents co-adaptation of features


## Dropout

## Regularization: Dropout

How can this possibly be a good idea?


Another interpretation:
Dropout is training a large ensemble of models (that share parameters).

Each binary mask is one model
An FC layer with 4096 units has $2^{4096} \sim 10^{1233}$ possible masks!
Only $\sim 10^{82}$ atoms in the universe...

[^10]
## Dropout

## Dropout: Test time

Dropout makes our output random!

| Output (label) | $\begin{gathered} \text { Input } \\ \text { (image) } \end{gathered}$ |
| :---: | :---: |
| $y=$ | (x, 2 |

Want to "average out" the randomness at test-time

$$
y=f(x)=E_{z}[f(x, z)]=\int p(z) f(x, z) d z
$$

But this integral seems hard ...

[^11]
## Dropout

## Dropout: Test time

Want to approximate the integral

$$
y=f(x)=E_{z}[f(x, z)]=\int p(z) f(x, z) d z
$$

Consider a single neuron.


## Dropout

## Dropout: Test time

Want to approximate the integral

$$
y=f(x)=E_{z}[f(x, z)]=\int p(z) f(x, z) d z
$$

Consider a single neuron.


At test time we have: $E[a]=w_{1} x+w_{2} y$

## Dropout

## Dropout: Test time

Want to approximate the integral

$$
y=f(x)=E_{z}[f(x, z)]=\int p(z) f(x, z) d z
$$

Consider a single neuron.


At test time we have: $E[a]=w_{1} x+w_{2} y$ During training we have: $E[a]=\frac{1}{4}\left(w_{1} x+w_{2} y\right)+\frac{1}{4}\left(w_{1} x+0 y\right)$

$$
\begin{aligned}
& +\frac{1}{4}(0 x+0 y)+\frac{1}{4}\left(0 x+w_{2} y\right) \\
= & \frac{1}{2}\left(w_{1} x+w_{2} y\right)
\end{aligned}
$$

## Dropout

## Dropout: Test time

Want to approximate the integral

$$
y=f(x)=E_{z}[f(x, z)]=\int p(z) f(x, z) d z
$$

Consider a single neuron.


At test time we have: $E[a]=w_{1} x+w_{2} y$
During training we have: $\quad E[a]=\frac{1}{4}\left(w_{1} x+w_{2} y\right)+\frac{1}{4}\left(w_{1} x+0 y\right)$

At test time, multiply
by dropout probability

$$
\begin{aligned}
& +\frac{1}{4}(0 x+0 y)+\frac{1}{4}\left(0 x+w_{2} y\right) \\
= & \frac{1}{2}\left(w_{1} x+w_{2} y\right)
\end{aligned}
$$

## Dropout

## Dropout: Test time

```
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
```

At test time all neurons are active always
=> We must scale the activations so that for each neuron: output at test time $=$ expected output at training time

[^12]
## Dropout

```
""" Vanilla Dropout: Not recommended implementation (see notes below) """
p = 0.5 # probability of keeping a unit active. higher = less dropout
Dropout Summary
def train_step(X):
    """ X contains the data """
    # forward pass for example 3-layer neural network
    H1 = np.maximum(0, np.dot (W1, X) + bl)
    U1 = np.random. rand(*H1.shape) < p # first dropout mask
    H1 *= U1 # drop!
H2 = np.max1mum(0, np.dot(W2, H1) + b2)
    U2 = np, random, rand(*H2.shape) < p # second dropout mask
    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
    scale at test time
    out = np.dot(W3, H2) + b3
```


## Dropout

## More common: "Inverted dropout"

```
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) + bl)
    U1 = (np.random. rand(*H1.shape) < p) / p # first dropout mask. Notice /p!
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)
                                    test time is unchanged!
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
```


## Data augmentation

## Regularization: Data Augmentation



[^13]
## Data augmentation

## Regularization: Data Augmentation



Transform image
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Data augmentation

## Data Augmentation Horizontal Flips



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## Data augmentation

## Data Augmentation <br> Random crops and scales

Training: sample random crops / scales
ResNet:

1. Pick random $L$ in range $[256,480]$
2. Resize training image, short side $=\mathrm{L}$
3. Sample random $224 \times 224$ patch

[^14]
## Data augmentation

## Data Augmentation <br> Random crops and scales

Training: sample random crops / scales
ResNet:

1. Pick random $L$ in range $[256,480]$
2. Resize training image, short side $=\mathrm{L}$
3. Sample random $224 \times 224$ patch

Testing: average a fixed set of crops
ResNet:

1. Resize image at 5 scales: $\{224,256,384,480,640\}$
2. For each size, use $10224 \times 224$ crops: 4 corners + center, + flips
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## Data augmentation

## Data Augmentation <br> Color Jitter

Simple: Randomize contrast and brightness


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## Data augmentation

## Data Augmentation Color Jitter

Simple: Randomize contrast and brightness


## More Complex:

1. Apply PCA to all $[R, G, B]$ pixels in training set
2. Sample a "color offset" along principal component directions
3. Add offset to all pixels of a training image
(As seen in [Krizhevsky et al. 2012], ResNet, etc)
[^15]
## Data augmentation

## Data Augmentation <br> Get creative for your problem!

Random mix/combinations of :

- translation
- rotation
- stretching
- shearing,
- lens distortions, ... (go crazy)

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## Other regularization techniques

## Regularization: A common pattern

Training: Add some kind of randomness

$$
y=f_{W}(x, z)
$$

Testing: Average out randomness (sometimes approximate)
$y=f(x)=E_{z}[f(x, z)]=\int p(z) f(x, z) d z$

## Other regularization techniques

## Regularization: A common pattern

Training: Add random noise
Testing: Marginalize over the noise

Examples:

Dropout
Batch Normalization
Data Augmentation

## Other regularization techniques

## Regularization: A common pattern

Training: Add random noise
Testing: Marginalize over the noise

Examples:<br>Dropout<br>Batch Normalization Data Augmentation DropConnect



Wan et al, "Regularization of Neural Networks using DropConnect", ICML 2013
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## Other regularization techniques

## Regularization: A common pattern

Training: Add random noise
Testing: Marginalize over the noise

Examples:<br>Dropout<br>Batch Normalization<br>Data Augmentation DropConnect<br>Fractional Max Pooling



Graham, "Fractional Max Pooling", arXiv 2014

$$
\text { Fei-Fei Li \& Justin Johnson \& Serena Yeung Lecture 7-84 April 25, } 2017
$$

## Other regularization techniques

## Regularization: A common pattern

Training: Add random noise
Testing: Marginalize over the noise

Examples:

Dropout
Batch Normalization
Data Augmentation
DropConnect
Fractional Max Pooling Stochastic Depth


[^16]
## Optimizers

## Optimization

```
# Vanilla Gradient Descent
while True:
    weights_grad = evaluate_gradient(loss_fun, data, weights)
    weights += - step_size * weights_grad # perform parameter update
```



## Optimizers

## Optimization: Problems with SGD

What if loss changes quickly in one direction and slowly in another?
What does gradient descent do?


Loss function has high condition number: ratio of largest to smallest singular value of the Hessian matrix is large

[^17]
## Optimizers

## Optimization: Problems with SGD

What if loss changes quickly in one direction and slowly in another?
What does gradient descent do?
Very slow progress along shallow dimension, jitter along steep direction


Loss function has high condition number: ratio of largest to smallest singular value of the Hessian matrix is large

[^18]
## Optimizers

## Optimization: Problems with SGD

What if the loss<br>function has a<br>local minima or saddle point?

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## Optimizers

## Optimization: Problems with SGD

What if the loss<br>function has a<br>local minima or saddle point?

Zero gradient, gradient descent gets stuck

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## Optimizers

## Optimization: Problems with SGD

What if the loss<br>function has a<br>local minima or saddle point?

Saddle points much more common in high dimension

## Optimizers

## Optimization: Problems with SGD

Our gradients come from
minibatches so they can be noisy!

$$
\begin{aligned}
L(W) & =\frac{1}{N} \sum_{i=1}^{N} L_{i}\left(x_{i}, y_{i}, W\right) \\
\nabla_{W} L(W) & =\frac{1}{N} \sum_{i=1}^{N} \nabla_{W} L_{i}\left(x_{i}, y_{i}, W\right)
\end{aligned}
$$



## Exponential moving average

- $S_{t}= \begin{cases}Y_{1}, & t=1 \\ \alpha \cdot Y_{t}+(1-\alpha) \cdot S_{t-1}, & t>1\end{cases}$


## Exponential moving average

- $S_{t}= \begin{cases}Y_{1}, & t=1 \\ \alpha \cdot Y_{t}+(1-\alpha) \cdot S_{t-1}, & t>1\end{cases}$
- $S_{t}=\alpha\left[Y_{t-1}+(1-\alpha) Y_{t-2}+(1-\alpha)^{2} Y_{t-3}+\cdots\right]$


## Exponential moving average

- $S_{t}= \begin{cases}Y_{1}, & t=1 \\ \alpha \cdot Y_{t}+(1-\alpha) \cdot S_{t-1}, & t>1\end{cases}$
- $S_{t}=\alpha\left[Y_{t-1}+(1-\alpha) Y_{t-2}+(1-\alpha)^{2} Y_{t-3}+\cdots\right]$
$=\frac{Y_{t-1}+(1-\alpha) Y_{t-2}+(1-\alpha)^{2} Y_{t-3}+\cdots}{1+(1-\alpha)+(1-\alpha)^{2}+\cdots}$



## Optimizers

## Momentum update

```
# Gradient descent update
x += - learning_rate * dx
```



- Physical interpretation as ball rolling down the loss function + friction (mu coefficient).
- mu $=$ usually $\sim 0.5,0.9$, or 0.99 (Sometimes annealed over time, e.g. from $0.5->0.99$ )

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## Optimizers

## Momentum update



- Allows a velocity to "build up" along shallow directions
- Velocity becomes damped in steep direction due to quickly changing sign

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## Optimizers



## Optimizers

## Nesterov Momentum update

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

Ordinary momentum update:

gradient
step

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## Optimizers

## Nesterov Momentum update



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## Optimizers

## Nesterov Momentum update



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## Optimizers

## Nesterov Momentum update

$$
\begin{gathered}
v_{t}=\mu v_{t-1}-\epsilon \nabla f\left(\theta_{t-1}+\mu v_{t-1}\right) \\
\theta_{t}=\theta_{t-1}+v_{t}
\end{gathered}
$$

Slightly inconvenient... usually we have :

$$
\theta_{t-1}, \nabla f\left(\theta_{t-1}\right)
$$

## Optimizers

## Nesterov Momentum

$$
\begin{aligned}
v_{t+1} & =\rho v_{t}-\alpha \nabla f\left(x_{t}+\rho v_{t}\right) \\
x_{t+1} & =x_{t}+v_{t+1}
\end{aligned}
$$

Annoying, usually we want update in terms of $x_{t}, \nabla f\left(x_{t}\right)$

Change of variables $\tilde{x}_{t}=x_{t}+\rho v_{t}$ and rearrange:

$$
\begin{aligned}
v_{t+1} & =\rho v_{t}-\alpha \nabla f\left(\tilde{x}_{t}\right) \\
\tilde{x}_{t+1} & =\tilde{x}_{t}-\rho v_{t}+(1+\rho) v_{t+1} \\
& =\tilde{x}_{t}+v_{t+1}+\rho\left(v_{t+1}-v_{t}\right)
\end{aligned}
$$

[^19]
## Optimizers



## AdaGrad update

[Duchi et al., 2011]

```
# Adagrad update
cache += dx**2
x += - learning_rate * dx / (np.sqrt(cache) + le-7)
```

Added element-wise scaling of the gradient based on the historical sum of squares in each dimension

## Optimizers

## AdaGrad update

```
# Adagrad update
cache += dx**2
x += - learning_rate * dx / (np.sqrt(cache) + le-7)
```



Q: What happens with AdaGrad?
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## Optimizers

## AdaGrad update

```
# Adagrad update
cache += dx**2
x += - learning_rate * dx / (np.sqrt(cache) + le-7)
```



Q2: What happens to the step size over long time?
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## Optimizers

number of steps $=10$


## Optimizers

## RMSProp update

[Tieleman and Hinton, 2012]
\# Adagrad update

| cache $+=\mathrm{dx} * * 2$ |  |
| :--- | :--- |
| $\mathrm{x}+=-$ learning_rate $* \mathrm{dx} /(\mathrm{np} . \operatorname{sqrt}($ cache $)+1 \mathrm{e}-7)$ |  |



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25 Jan 2016

## Optimizers

## rmsprop: A mini-batch version of rprop

- rprop is equivalent to using the gradient but also dividing by the size of the gradient.
- The problem with mini-batch rprop is that we divide by a different number for each mini-batch. So why not force the number we divide by to be very similar for adjacent mini-batches?
- rmsprop: Keep a moving average of the squared gradient for each weight $\operatorname{MeanSquare}(w, t)=0.9 \operatorname{MeanSquare}(w, t-1)+0.1(\partial E / \partial w(t))^{2}$
- Dividing the gradient by $\sqrt{\operatorname{MeanSquare}(w, t)}$ makes the learning work much better (Tijmen Tieleman, unpublished).

Introduced in a slide in Geoff Hinton's Coursera class, lecture 6

## Optimizers

## rmsprop: A mini-batch version of rprop

- rprop is equivalent to using the gradient but also dividing by the size of the gradient.
- The problem with mini-batch rprop is that we divide by a different number for each mini-batch. So why not force the number we divide by to be very similar for adjacent mini-batches?
- rmsprop: Keep a moving average of the squared gradient for each weight $\operatorname{MeanSquare}(w, t)=0.9 \operatorname{MeanSquare}(w, t-1)+0.1(\partial E / \partial w(t))^{2}$
- Dividing the gradient by $\sqrt{\operatorname{MeanSquare}(w, t)}$ makes the learning work much better (Tijmen Tieleman, unpublished).

Introduced in a slide in Geoff Hinton's Coursera class, lecture 6

Cited by several papers as:
[52] T. Tieleman and G. E. Hinton. Lecture 6.5-rmsprop: Divide the gradient by a running average of its recent magnitude., 2012.

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## Optimizers

number of steps=10


## Optimizers

## Adam update

[Kingma and Ba , 2014]
(incomplete, but close)

```
# Adam
m = betal*m + (1-betal)*dx # update first moment
v = beta2*v + (1-beta2)*(dx**2) # update second moment
x += - learning_rate * m / (np.sqrt(v) + le-7)
```


## Optimizers

## Adam update

[Kingma and Ba, 2014]
(incomplete, but close)

| \#\# $\mathrm{m}=$ betal*m + (1-betal)*dx \# update first mome |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| $\mathrm{v}=\mathrm{beta}^{*} \mathrm{v}+(1-\mathrm{beta2}) *(\mathrm{dx} * * 2)$ \# update second moment |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| x += - learning_rate * m / (np.sqrt(v) + le-7) |  |  |  |  |  |  |  |  |  |  |  |  |  |  |

Looks a bit like RMSProp with momentum

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## Optimizers

## Adam update

[Kingma and Ba, 2014]
(incomplete, but close)

| \# Adam |
| :--- |
| $\mathrm{m}=$ beta ${ }^{*} \mathrm{~m}+(1$-betal $) * \mathrm{dx} \#$ update first moment <br> $\mathrm{V}=$ beta $2 * \mathrm{v}+(1-$ beta 2$) *(\mathrm{dx*2}) ~$ update second moment <br> $\mathrm{x}+=-$ learning_rate $* \mathrm{~m} /(\mathrm{np} . \operatorname{sqrt}(\mathrm{v})+1 \mathrm{e}-7)$ |

momentum
$\mathrm{V}=\mathrm{beta}^{*} \mathrm{~V}+(1$-beta2 $) *(\mathrm{dX} * * 2)$ \# update second moment
x += - learning_rate * m / (np.sqrt(v) + le-7)
RMSProp-like

## Looks a bit like RMSProp with momentum

```
# RMSProp
cache = decay_rate * cache + (1 - decay_rate) * dx**2
x += - learning_rate * dx / (np.sqrt(cache) + le-7)
```

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## Optimizers

## Adam (full form)

```
    first_moment = 0
    second_moment = 0
    for t in range(1, num_iterations):
    dx = compute gradient (x)
    first_moment = beta1 * first_moment + (1 - beta1) * dx
    second_moment = beta2 * second_moment + (1 - beta2) * dx * dx
    first_unbias = first_moment / (1 - beta1 ** t)
    second_unbias = second_moment / (1 - beta2 ** t)
```


## Momentum

## Bias correction

AdaGrad / RMSProp

Bias correction for the fact that first and second moment estimates start at zero

Adam with beta1 $=0.9$, beta $2=0.999$, and learning_rate $=1 e-3$ or $5 e-4$ is a great starting point for many models!

## Optimizers

number of steps $=10$


## Optimizers

SGD, SGD+Momentum, Adagrad, RMSProp, Adam all have learning rate as a hyperparameter.


## Q: Which one of these learning rates is best to use?

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## Optimizers

## SGD, SGD+Momentum, Adagrad, RMSProp, Adam all have learning rate as a hyperparameter.


=> Learning rate decay over time!
step decay:
e.g. decay learning rate by half every few epochs.
exponential decay:

$$
\alpha=\alpha_{0} e^{-k t}
$$

1/t decay:

$$
\alpha=\alpha_{0} /(1+k t)
$$

## Optimizers

## SGD, SGD+Momentum, Adagrad, RMSProp, Adam all have learning rate as a hyperparameter.




[^20]
## Optimizers

## Second order optimization methods

second-order Taylor expansion:

$$
J(\boldsymbol{\theta}) \approx J\left(\boldsymbol{\theta}_{0}\right)+\left(\boldsymbol{\theta}-\boldsymbol{\theta}_{0}\right)^{\top} \nabla_{\boldsymbol{\theta}} J\left(\boldsymbol{\theta}_{0}\right)+\frac{1}{2}\left(\boldsymbol{\theta}-\boldsymbol{\theta}_{0}\right)^{\top} \boldsymbol{H}\left(\boldsymbol{\theta}-\boldsymbol{\theta}_{0}\right)
$$

Solving for the critical point we obtain the Newton parameter update:

$$
\boldsymbol{\theta}^{*}=\boldsymbol{\theta}_{0}-\boldsymbol{H}^{-1} \nabla_{\boldsymbol{\theta}} J\left(\boldsymbol{\theta}_{0}\right)
$$

Q: what is nice about this update?

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## Optimizers

## Second order optimization methods

$$
\boldsymbol{\theta}^{*}=\boldsymbol{\theta}_{0}-\boldsymbol{H}^{-1} \nabla_{\boldsymbol{\theta}} J\left(\boldsymbol{\theta}_{0}\right)
$$

- Quasi-Newton methods (BGFS most popular): instead of inverting the Hessian ( $O\left(n^{\wedge} 3\right)$ ), approximate inverse Hessian with rank 1 updates over time (O( $n^{\wedge} 2$ ) each).
- L-BFGS (Limited memory BFGS): Does not form/store the full inverse Hessian.


## Quasi-Newton methods (watch this)

- Ref:
(1) https://www.youtube.com/watch?v=uo2z0AT_83k
(2) Nocedal \& Wright - Numerical Optimization $(B \leftrightarrow H)$
(3) http://users.ece.utexas.edu/ cmcaram/EE381V_2012F/Lecture_10_Scribe_Notes.final.pdf
- The inverse of Hessian $H$ is expensive to compute. Want to approximate it iteratively instead


## Quasi-Newton methods (watch this)

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- The inverse of Hessian $H$ is expensive to compute. Want to approximate it iteratively instead
- Quasi-Newton methods:
(1) Approximate Newton direction

$$
d_{k}=-B_{k} g_{k},
$$

where $B_{k} \approx H^{-1}$ and $g_{k}=\nabla J\left(\theta_{k}\right)$

## Quasi-Newton methods (watch this)

- Ref:
(1) https://www.youtube.com/watch?v=uo2z0AT_83k
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caram/EE381V_2012F/Lecture_10_Scribe_Notes.final.pdf
- The inverse of Hessian $H$ is expensive to compute. Want to approximate it iteratively instead
- Quasi-Newton methods:
(1) Approximate Newton direction

$$
d_{k}=-B_{k} g_{k},
$$

where $B_{k} \approx H^{-1}$ and $g_{k}=\nabla J\left(\theta_{k}\right)$
(2) Line search: $\theta_{k+1}=\theta_{k}+\alpha_{k} d_{k}$

## Quasi-Newton methods (watch this)

- Ref:
(1) https://www.youtube.com/watch?v=uo2z0AT_83k
(2) Nocedal \& Wright - Numerical Optimization $(B \leftrightarrow H)$
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caram/EE381V_2012F/Lecture_10_Scribe_Notes.final.pdf
- The inverse of Hessian $H$ is expensive to compute. Want to approximate it iteratively instead
- Quasi-Newton methods:
(1) Approximate Newton direction

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where $B_{k} \approx H^{-1}$ and $g_{k}=\nabla J\left(\theta_{k}\right)$
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## Quasi-Newton methods (watch this)

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(9) Approximate inverse Hessian

$$
B_{k+1}=\text { update_formula }\left(B_{k}, \theta_{k+1}-\theta_{k}, g_{k+1}-g_{k}\right)
$$

## Approximation with rank-1 update

- As Hessian is essentially the "derivative" of $\nabla J$, we have $\nabla J\left(\theta_{k+1}\right) \approx \nabla J\left(\theta_{k}\right)+H\left(\theta_{k+1}-\theta_{k}\right)$


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## Inverse Hessian update for BFGS

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- $\left(D-\frac{H p p^{T} H}{p^{T} H^{T} p}\right)^{-1}=D^{-1}-\frac{D^{-1} H p p^{T} H D^{-1}}{p^{T} q q^{T} p\left(q^{T} p-q^{T} B q\right)} \cdots$


## Inverse Hessian update for BFGS

- Recall $H_{k+1}=H_{k}+\frac{q_{k} q_{k}^{T}}{q_{k}^{T} p_{k}}-\frac{H_{k} p_{k} p_{k}^{T} H_{k}}{p_{k}^{T} H_{k}^{T} p_{k}}$ and

$$
\left(A+u v^{T}\right)^{-1}=A^{-1}+\frac{A^{-1} u v^{T} A^{-1}}{1-v^{T} A^{-1} u}
$$

- $D^{-1}=\left(H+\frac{q q^{T}}{q^{T} p}\right)^{-1}=H^{-1}+\frac{H^{-1} q q^{T} H^{-1}}{\left(q^{T} p\right)\left(1-q^{T} H^{-1} q /\left(q^{T} p\right)\right)}=B+\frac{B q q^{T} B}{q^{T} p-q^{T} B q}$
- $\left(D-\frac{H p p^{T} H}{p^{T} H^{T} p}\right)^{-1}=D^{-1}-\frac{D^{-1} H p p^{T} H D^{-1}}{p^{T} H^{T} p\left(1-p^{T} H D^{-1} H p /\left(p^{T} H^{T} p\right)\right)}$
$=D^{-1}-\frac{D^{-1} H p p^{T} H D^{-1}}{p^{T} H p-p^{T} H D^{-1} H p}$
- $D^{-1} H p=\left(B H p+\frac{B q q^{T} B H p}{q^{T} p-q^{T} B q}\right)=\left(p+\frac{B q q^{T} p}{q^{T} p-q^{T} B q}\right)$
- $\left(D-\frac{H p p^{T} H}{p^{T} H^{T} p}\right)^{-1}=D^{-1}-\frac{D^{-1} H p p^{T} H D^{-1}}{p^{T} q q^{T} p\left(q^{T} p-q^{T} B q\right)} \cdots$
- $\left(D-\frac{H p p^{T} H}{p^{T} H^{T} p}\right)^{-1}=\left(I-\frac{p q^{T}}{q^{T} p}\right) B\left(I-\frac{q p^{T}}{q^{T} p}\right)+\frac{p p^{T}}{q^{T} p}$
$\Rightarrow B_{k+1}=\left(I-\frac{p_{k} q_{k}^{T}}{q_{k}^{T} p_{k}}\right) B_{k}\left(I-\frac{q_{k} p_{k}^{T}}{q_{k}^{T} p_{k}}\right)+\frac{p_{k} p_{k}^{T}}{q_{k}^{T} p_{k}}$


## Inverse Hessian update for BFGS

- Recall $H_{k+1}=H_{k}+\frac{q_{k} q_{k}^{T}}{q_{k}^{T} p_{k}}-\frac{H_{k} p_{k} p_{k}^{T} H_{k}}{p_{k}^{T} H_{k}^{T} p_{k}}$ and

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\left(A+u v^{T}\right)^{-1}=A^{-1}+\frac{A^{-1} u v^{T} A^{-1}}{1-v^{T} A^{-1} u}
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- $D^{-1}=\left(H+\frac{q q^{T}}{q^{T} p}\right)^{-1}=H^{-1}+\frac{H^{-1} q q^{T} H^{-1}}{\left(q^{T} p\right)\left(1-q^{T} H^{-1} q /\left(q^{T} p\right)\right)}=B+\frac{B q q^{T} B}{q^{T} p-q^{T} B q}$
- $\left(D-\frac{H p p^{T} H}{p^{T} H^{T} p}\right)^{-1}=D^{-1}-\frac{D^{-1} H p p^{T} H D^{-1}}{p^{T} H^{T} p\left(1-p^{T} H D^{-1} H p /\left(p^{T} H^{T} p\right)\right)}$
$=D^{-1}-\frac{D^{-1} H p p^{T} H D^{-1}}{p^{T} H p-p^{T} H D^{-1} H p}$
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- $\left(D-\frac{H p p^{T} H}{p^{T} H^{T} p}\right)^{-1}=D^{-1}-\frac{D^{-1} H p p^{T} H D^{-1}}{p^{T} q q^{T} p\left(q^{T} p-q^{T} B q\right)} \cdots$
- $\left(D-\frac{H p p^{T} H}{p^{T} H^{T} p}\right)^{-1}=\left(I-\frac{p q^{T}}{q^{T} p}\right) B\left(I-\frac{q p^{T}}{q^{T} p}\right)+\frac{p p^{T}}{q^{T} p}$
$\Rightarrow B_{k+1}=\left(I-\frac{p_{k} q_{k}^{T}}{q_{k}^{T} p_{k}}\right) B_{k}\left(I-\frac{q_{k} p_{k}^{T}}{q_{k}^{T} p_{k}}\right)+\frac{p_{k} p_{k}^{T}}{q_{k}^{T} p_{k}}$
- Bounty: $3 \%$ bonus to complete the algebra


## Summary of BFGS

Initialize Initialize inverse Hessian approximation $B \leftarrow B_{0}$. Can set $B \leftarrow I$ if no initial estimate; $k \leftarrow 0$; Pick a random starting point $\theta_{0}$
Loop (1) Get search direction $d_{k}=-B_{k} \nabla J\left(\theta_{k}\right)$
(2) Conduct line search to find optimum

$$
\theta_{k+1}=\theta_{k}+\alpha_{k} d_{k}
$$

(3) $p_{k} \leftarrow \theta_{k+1}-\theta_{k} ; q_{k} \leftarrow \nabla J\left(\theta_{k+1}\right)-\nabla J\left(\theta_{k}\right)$; $B_{k+1}=\left(I-\frac{p_{k} q_{k}^{T}}{q_{k}^{T} p_{k}}\right) B_{k}\left(I-\frac{q_{k} p_{k}^{T}}{q_{k}^{T} p_{k}}\right)+\frac{p_{k} p_{k}^{T}}{q_{k}^{T} p_{k}}$
(1) $k \leftarrow k+1$; Exit if $\left\|\nabla J\left(\theta_{k}\right)\right\|>\epsilon$

## Inverse Hessian update for BFGS

- Like rank-1 update, we can also rearrange the variables to obtain an update rule for $B=H^{-1}$
- Instead of $H_{k+1} p_{k}=q_{k}$, we want $B_{k+1} q_{k}=p_{k}$.


## Inverse Hessian update for BFGS

- Like rank-1 update, we can also rearrange the variables to obtain an update rule for $B=H^{-1}$
- Instead of $H_{k+1} p_{k}=q_{k}$, we want $B_{k+1} q_{k}=p_{k}$. Thus we have

$$
B_{k+1}=B_{k}+\frac{p_{k} p_{k}^{T}}{p_{k}^{T} q_{k}}-\frac{B_{k} q_{k} q_{k}^{T} B_{k}}{q_{k}^{T} B_{k}^{T} q_{k}}
$$

- Note that this update rule of $B$ is different from before. Actually this is the update rule of DFP. An older approach that is considered worse compared with BFGS


## Some theoretical notes

- A prettier but more technical explanation of BFGS/DFP involves weighted matrix norm


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(1) $B_{k+1} q_{k}=p_{k}$ (secant equation)
(2) $B_{k+1} \succ 0$ (symmetric and positive definite),
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where $\|A\|_{W}=\left\|W^{1 / 2} A W^{1 / 2}\right\|_{F}$ is the weighted Frobenius norm
$\bullet \Rightarrow \begin{cases}\text { BFGS } & W=H \\ \text { DFP } & W=H^{-1}\end{cases}$

## LBFGS

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## LBFGS

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- Instead of storing $B_{k}$, we can store the previous last several $p$ and $q$ to estimate $B_{k+1}$
- Let say we store the last $r$ pairs, we need to iterate $r$ times (instead of just once) and the estimate is less accurate
- Storage requirement decreases drastically


## Optimizers

## L-BFGS

- Usually works very well in full batch, deterministic mode i.e. if you have a single, deterministic $f(x)$ then L-BFGS will probably work very nicely
- Does not transfer very well to mini-batch setting. Gives bad results. Adapting L-BFGS to large-scale, stochastic setting is an active area of research.

$$
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$$

## Optimizers

## In practice:

- Adam is a good default choice in most cases
- If you can afford to do full batch updates then try out L-BFGS (and don't forget to disable all sources of noise)


## Babysitting learning process

## Step 1: Preprocess the data


(Assume $\mathrm{X}[\mathrm{NxD}]$ is data matrix, each example in a row)

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## Babysitting learning process

## Step 2: Choose the architecture: say we start with one hidden layer of 50 neurons:



## Babysitting learning process

## Double check that the loss is reasonable:

```
def init_two_layer_model(input_size, hidden_size, output_size):
    # initialize a model
    model = {}
    model['W1'] = 0.0001 * np.random.randn(input_size, hidden_size)
    model['b1'] = np.zeros(hidden_size)
    model['W2'] = 0.0001 * np.random.randn(hidden_size, output_size)
    model['b2'] = np.zeros(output_size)
    return model
```



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## Debugging optimizer

## Double check that the loss is reasonable:

```
def init_two_layer_model(input_size, hidden_size, output_size):
    # initialize a model
    model = {}
    model['W1'] = 0.0001 * np.random.randn(input_size, hidden_size)
    model['b1'] = np.zeros(hidden_size)
    model['W2'] = 0.0001 * np.random.randn(hidden_size, output_size)
    model['b2'] = np.zeros(output_size)
    return model
```

model = init_two_layer_model $(32 * 32 * 3,50,10)$ \# input size, hidden size, number of classes
loss, grad =-two_layer_net (X_train, model, y_train, 1e3) crank up regularization
loss, grad = two_layer_net (X_train, model, y_train, 1e3) crank up regularization
print loss
3.06859716482


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## Debugging optimizer

Lets try to train now...

Tip: Make sure that you can overfit very small portion of the training data

```
model = init two layer_model(32*32*3, 50, 10) # input size, hidden size, number of classes
```

trainer $=$ ClāssifierTräner()

X_tiny $=$ X_train $[: 20]$ \#take 20 examples
$y$ tiny $=y$ train [:20]
 mōdel, twō_layer_net,
num epochs $=200, \bar{r} \mathrm{eg}=0.0$,
updāte='sgd', learning rate decay=1,
sample batches $=$ False,
learning_rate $=1 e-3$, verbose=True)

The above code:

- take the first 20 examples from CIFAR-10
- turn off regularization (reg = 0.0)
- use simple vanilla 'sgd'


## Debugging optimizer

## Lets try to train now...

## Tip: Make sure that you can overfit very small portion of the training data

> Very small loss, train accuracy 1.00, nice!
model $=$ init two layer model $(32 * 32 * 3,50,10)$ input size, hidden size, number of classes trainer $=$ ClāssifierTräiner()
X_tiny $=$ X_train $[: 20]$ \#take 20 examples
$y \operatorname{tin} y=y \operatorname{train}[: 20]$
best_model, stats = trainer. train (X_tiny, y_tiny, X_tiny, y_tiny, mōdel, twō layer net,
num_epochs $=200, \bar{r} \mathrm{reg}=0.0$,
updāte='sgd', learning rate decay=1,
sample batches $=$ False,
learning rate=1e-3, verbose=True)
Finished epoch $1 / 200:$ cost 2.302603 , train: 0.400000 , val 0.400000 , lr $1.000000 \mathrm{e}-03$ Finished epoch $2 / 200:$ cost 2.302258, train: 0.450000 , val 0.450000 , lr $1.000000 \mathrm{e}-03$ Finished epoch $3 / 200$ : cost 2.301849, train: 0.600000 , val 0.600000 , lr $1.000000 \mathrm{e}-03$ Finished epoch $4 / 200$ : cost 2.301196 , train: 0.650000 , val 0.650000 , lr $1.000000 \mathrm{e}-03$ Finished epoch $5 / 200$ : cost 2.300044 , train: 0.650000 , val 0.650000 , lr $1.000000 \mathrm{e}-03$ Finished epoch $6 / 200:$ cost 2.297864 , train: 0.550000 , val 0.550600 , lr $1.000000 \mathrm{e}-03$ Finished epoch $7 / 200$ : cost 2.293595, train: 0.600000 , val 0.600000, lr 1.000000e-03 Finished epoch $8 / 200$; cost 2.285096 , train: 0.550000 , val 0.550000 , ir $1.000000 \mathrm{e}-03$ Finished epoch $9 / 200$ : cost 2.268094 , train: 0.550000 , val 0.550000 , ir $1.000000 e-03$ Finished epoch $10 / 200$ : cost 2.234787 , train: 0.500000 , val 0.500000 , Ir $1.000000 e-03$ Finished epoch $11 / 200:$ cost 2.173187, train: 0.500000 , val 0.500000 , lr $1.000000 e-03$ Finished epoch $12 / 200$ : cost 2.076862 , train: 0.500000 , val 0.500000 , lr $1.000000 \mathrm{e}-03$ Finished epoch $13 / 200$ : cost 1.974090 , train: 0.400000 , val 0.400000 , lr $1.000000 \mathrm{e}-03$ Finished epoch $14 / 200$ : cost 1.895885 , train: 0.400000 , val 0.400000 , ir $1.000000 \mathrm{e}-03$ Finished epoch $15 / 200:$ cost 1.820876 , train: 0.450000 , val 0.450000 , lr $1.000000 \mathrm{e}-03$ Finished epoch $16 / 200$ : cost 1.737430 , train: 0.450000 , val 0.450000 , lr $1.000000 \mathrm{e}-03$ Finished epoch $17 / 200$ : cost 1.642356 , train: 0.500000 , val 0.500000 , lr $1.000000 e-03$ Finished epoch $18 / 200$ : cost 1.535239 , train: 0.600000 , val 0.600000 , lr $1.000000 \mathrm{e}-03$ Finished epoch 19/, 200: cost 1.421527, train: 0.600000, val 0.600000, ir 1.000000e-03 Finished epoch 195 / 200 : cost 0.002694 , train: 1.000000, val 1.000000, lr $1.000000 \mathrm{e}-03$ Finished epoch $196 / 200$ : cost 0.002674 , train: 1.000000 , val 1.000000 , lr $1.000000 \mathrm{e}-03$ Finished epoch 197 / 200 : cost 0.002655 , train: 1.000000 , val 1.000000 , ir $1.000000 e-03$ Finished epoch $198 / 200$ : cost 0.002635 , train: 1.000000, val 1.000000, ir 1.000000e-03 Finished epoch 199 / 200 : cost 0.002617 , train: 1.000000, val 1.000000, Ir 1.000000e-03 Finished epoch 200 / 200 : cost 0.002597 , train: 1.000000, val 1.000000, ir $1.000000 e-03$ finished optimization. best validation accuracy: 1.000000

## Debugging optimizer

Lets try to train now...

I like to start with small regularization and find learning rate that makes the loss go down.

mōdel, two_layer_net̄, num epochs $\overline{=}=10$, $\overline{\mathrm{re}} \mathrm{g}=0.000001$,
update='sgd', learning_rate_decay=1, sample batches $=$ True,
learning_rate $=1 \mathrm{e}-6$, verbose=True)

## Debugging optimizer

Lets try to train now...

## I like to start with small regularization and find learning rate that makes the loss go down.

model $=$ init two layer_model $(32 * 32 * 3,50,10)$ \# input size, hidden size, number of classes trainer $=$ ClāssifierTräner()
best_model, stats = trainer.train(X_train, y_train, $x_{-}$val, y_val, mōdel, two_layer_nē, num epochs $=10$, reg=0.000001, update='sgd', learning_rate_decay=1, sampte batches - True,
learning_rate $=1 e-6$, verbose=True)
Finished epoch 1 / 10:
Finished epoch $2 / 10$ : Finished epoch 3 / 10 : Finished epoch 4 / 10: Finished epoch 5 / 10 : Finished epoch 6/10: Finished epoch $7 / 10$ : Finished epoch 8/10: Finished epoch 9/10: Finished epoch $10 / 10$ : cost 2.302420 train: 0.206000 , yal $0.192000, \operatorname{lr} 1.000000 \mathrm{e}-06$ finished optimization. best validation accuracy: 0.192000

Loss barely changing

## Debugging optimizer

Lets try to train now...

> I like to start with small regularization and find learning rate that makes the loss go down.
model $=$ init two layer_model $(32 * 32 * 3,50,10)$ \# input size, hidden size, number of classes trainer = ClassifierTrainer()
best_model, stats $=$ trainer. $\operatorname{train}\left(X_{\_}\right.$train, $y \_t r a i n, ~ X \_v a l, ~ y \_v a l, ~$ módel, two_layer_net, num_epochs=10, reg=0.000001, update='sgd', learning_rate_decay=1, sampte batehes - True,

Finished epoch $1 / 10$ :
Finished epoch 2 / 10: Finished epoch 3/10: Finished epoch 4 / 10 : Finished epoch 5 / 10 : Finished epoch 6/10: Finished epoch 7 / 10: Finished epoch 8/10: Finished epoch 9/10: finished optimization. best validation accuracy: 0.192000
learning_rate $=1 \mathrm{e}-6$, verbose $=$ True) Finished epoch $9 / 10:$ cost 2.302459 , train: 0.206000, y y 0.192000 , $\operatorname{lr} 1.000000 \mathrm{e}-06$
Finished epoch $10 / 10$, cost 2.302420 train: 0.190000 , val 0.192000 , $\operatorname{lr} 1.000000 \mathrm{e}-06$
trai 0.08000,

## cost 2.302576 , train: 0.080000, yal 0.103000, lr 1.000000e-06

 cost 2.302582, train: 0.121000, yal 0.124000, lr 1.000000e-06 cost 2.302558, trair: 0.119000, yal 0.138000, ir 1.000000e-06 cost 2.302519, train: 0.127000, yal 0.151000, lr 1.000000e-06 cost 2.302517, train: 0.158000, yal 0.171000, lr 1.000000e-06 cost 2.302518, trair: 0.179000, yal 0.172000, ir 1.000000e-06 cost 2.302466, train: 0.180000, yal 0.176000, lr 1.000000e-06 cost 2.302452, trair: 0.175000, yal 0.185000, lr 1.000000e-06 cost 2.302459, train: 0.206000, yal 0.192000, ir 1.000000e-06

Loss barely changing: Learning rate is probably too low

## loss not going down: learning rate too low

## Debugging optimizer

Lets try to train now...

> I like to start with small regularization and find learning rate that makes the loss go down.
model $=$ init two layer_model $(32 * 32 * 3,50,10)$ \# input size, hidden size, number of classes trainer = ClassifierTrainer()
best_model, stats $=$ trainer. $\operatorname{train}\left(X_{\_}\right.$train, $y \_t r a i n, ~ X \_v a l, ~ y \_v a l, ~$ model, two_layer_net, num epochs=10, reg=0.000001, update='sgd', learning_rate_decay=1, sample batches - True,
learning_rate $=1 e-6$, verbose $=$ True)
Finished epoch 1 / 10:
Finished epoch 2/10: Finished epoch 3/10: Finished epoch 4 / 10 : Finished epoch 5 / 10 : Finished epoch 6/10: Finished epoch $7 / 10$ : Finished epoch 8/10: Finished epoch 9/10: Finished epoch $10 / 10$; cost 2.302420 train: 0.190000 , val 0.192000 , $\operatorname{lr} 1.000000 \mathrm{e}-06$ finished optimization. best validation accuracy: 0.192000

Loss barely changing: Learning rate is

## loss not going down: learning rate too low

probably too low

Notice train/val accuracy goes to $20 \%$ though, what's up with that? (remember this is softmax)

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## Debugging optimizer

Lets try to train now...
I like to start with small regularization and find learning rate that makes the loss go down.
model $=$ init two layer model $(32 * 32 * 3,50,10) \#$ input size, hidden size, number of classes trainer = ClassifierTrainer()
best model, stats $=$ trainer, train ( $x$ train, $y$ train, $X$ val, y val,
model, two layer net,
num epochs $=10$, $\mathrm{reg}=0.000001$,
update='sgd', learning_rate_decay=1,
sample batches $=$ True,
learning_rate=le6, verbose=True)
$\uparrow$
Okay now lets try learning rate 1 e . What could possibly go wrong?

## loss not going down: learning rate too low

## Debugging optimizer

Lets try to train now...
I like to start with small regularization and find learning rate that makes the loss go down.
model $=$ init two layer model $(32 * 32 * 3,50,10) \#$ input size, hidden size, number of classes trainer $=$ clāssif̄ierTräiner()
best model, stats $=$ trainer. $\operatorname{train}(X$ train, $y$ train, $x$ val, $y$ val,
model, two_layer_net,
num epochs $=10$, reg=0.000001,
update='sgd', learning_rate_decay=1,
sample batches $=$ True,
learning_rate=le6, verbose=True)
/home/karpathy/cs 231 n /code/cs231n/classifiers/neural_net.py:50: RuntimeWarning: divide by zero en countered in $\log$
data loss $=-n p . \operatorname{sum}(n p . \log ($ probs $[$ range $(N), y])) / N$
/home/karpathy/cs231n/code/cs231n/classifiers/neural_net.py:48: RuntimeWarning: invalid value enc ountered in subtract
probs $=n p . \exp (s c o r e s ~-n p . \max (s c o r e s, ~ a x i s=1$, keepdims=True))
Finished epoch $1 / 10:$ cost nan, train: 0.091000, val 0.087000, $\operatorname{lr} 1.000000 \mathrm{e}+66$ Finished epoch $2 / 10:$ cost nan, train: 0.095000 , val 0.087000 , lr $1.000000 \mathrm{e}+06$ Finished epoch $3 / 10$ : cost nan, train: 0.100000 , val 0.087000 , $\operatorname{lr} 1.000000 \mathrm{e}+06$
loss not going down: learning rate too low loss exploding: learning rate too high

## cost: NaN almost always means high learning rate...

## Debugging optimizer

Lets try to train now...

## I like to start with small regularization and find learning rate that makes the loss go down.

## loss not going down: learning rate too low loss exploding: learning rate too high

model $=$ init two layer model (32*32*3, 50, 10) \# input size, hidden size, number of classes trainer = ClassifierTräiner()
best model, stats $=$ trainer. train $(X$ train, $y$ train, $X$ val, $y$ val,
model, two_layer_net,
num_epochs=10, reg=0.000001,
update='sgd', learning_rate_decay=1,
sample_batches $=$ True,
learning_rate $=3 \mathrm{e}-3$, verbose $=$ True)

## Finished epoch $1 / 10:$ cost 2.186654, train: 0.308000, val 0.306000, lr 3.000000e-03

 Finished epoch 2 / 10: cost 2.176230, train: 0.330000, val 0.350000, lr 3.000000e-03 Finished epoch $3 / 10$ : cost 1.942257, train: 0.376000, val 0.352000, Ir 3.000000e-03 Finished epoch 4 / 10: cost 1.827868 , train: 0.329000, val 0.310000, lr 3.000000e-03 Finished epoch 5 / 10: cost inf, train: 0.128000, val 0.128000, lr 3.000000e-03 Finished epoch $6 / 10$ : cost inf, train: 0.144000 , val 0.147000 , lr $3.000000 \mathrm{e}-03$$3 \mathrm{e}-3$ is still too high. Cost explodes....
=> Rough range for learning rate we should be cross-validating is somewhere [1e-3 ... 1e-5]

```
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```


## Hyperparameter optimization

## Hyperparameter Optimization

## Hyperparameter optimization

## Cross-validation strategy

I like to do coarse -> fine cross-validation in stages
First stage: only a few epochs to get rough idea of what params work Second stage: longer running time, finer search
... (repeat as necessary)

Tip for detecting explosions in the solver:
If the cost is ever > 3 * original cost, break out early

$$
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$$

## Hyperparameter optimization

## For example: run coarse search for 5 epochs

$\max$ count $=100$
for count in xrange(max_count) :
reg $=10^{* *}$ uniform( $-5,5$ )
$\mathrm{lr}=10^{* *}$ uniform $(-3,-6)$
trainer = ClassifierTrainer()
model $=$ init two layer_model $(32 * 32 * 3,50,10)$ \# input size, hidden size, number of classes trainer = ClāssifierTrāiner()
best_model_local, stats = trainer.train(X_train, y_train, X_val, y_val, model, two_layer_net, num epochs=5, reg=reg, update $=$ 'momentum', learning_rate_decay $=0.9$, sample batches $=$ True, batch siz $\overline{\mathrm{e}}=100$, learning rate=lr, verbose=False)

| val_acc: | $0.412000, \operatorname{lr}: 1.405206 \mathrm{e}-04$, reg: $4.793564 \mathrm{e}-01,(1 / 100)$ |
| :--- | :--- | :--- | :--- | :--- | :--- |
| val_acc: $0.214000, \operatorname{lr}: 7.231888 \mathrm{e}-06$, reg: $2.321281 \mathrm{e}-04,(2 / 100)$ |  |
| val_acc: $0.208000, \operatorname{lr}: 2.119571 \mathrm{e}-06$, reg: $8.011857 \mathrm{e}+01,(3 / 100)$ |  |
| val_acc: $0.196000, \operatorname{lr}: 1.551131 \mathrm{e}-05$, reg: $4.374936 \mathrm{e}-05,(4 / 100)$ |  |
| val_acc: $0.079000, \operatorname{lr}: 1.753300 \mathrm{e}-05$, reg: $1.200424 \mathrm{e}+03,(5 / 100)$ |  |
| val_acc: $0.223000, \operatorname{lr}: 4.215128 \mathrm{e}-05$, reg: $4.196174 \mathrm{e}+01,(6 / 100)$ |  |
| val_acc: $0.441000, \operatorname{lr}: 1.750259 \mathrm{e}-04$, reg: $2.110807 \mathrm{e}-04,(7 / 100)$ |  |
| val acc: $0.241000, \operatorname{lr}: 6.749231 \mathrm{e}-05$, reg: $4.226413 \mathrm{e}+01,(8 / 100)$ |  |
| val_acc: $0.482000, \operatorname{lr}: 4.296863 \mathrm{e}-04$, reg: $6.642555 \mathrm{e}-01,(9 / 100)$ |  |
| val_acc: $0.079000, \operatorname{lr}: 5.401602 \mathrm{e}-06$, reg: $1.599828 \mathrm{e}+04,(10 / 100)$ |  |
| val_acc: $0.154000, \operatorname{lr}: 1.618508 \mathrm{e}-06$, reg: $4.925252 \mathrm{e}-01,(11 / 100)$ |  |

## Hyperparameter optimization

## Now run finer search...

```
max count = 100
for count in xrange(max_count):
    reg = 10**uniform( }-5,5
    lr = 10**uniform( -3, -6)
```

$\xrightarrow{\text { adjust range }}$

```
max count = 100
for count in xrange(max count):
    reg = 10**uniform( }-4,0
    lr = 10**uniform( -3, -4)
```

val acc: 0.527000, lr: $5.340517 \mathrm{e}-04$, reg: $4.097824 \mathrm{e}-01,(0 / 100)$
val acc: 0.492000 , lr: $2.279484 \mathrm{e}-04$, reg: $9.991345 \mathrm{e}-04,(1 / 100)$
val acc: 0.512000, lr: 8.680827e-04, reg: 1.349727e-02, (2 / 100)
val_acc: 0.461000, $\imath r: 1.028377 e-04$, reg: $1.220193 e-02$, ( $3 / 100$ )
val_acc: 0.460000 , lr: $1.113730 \mathrm{e}-04$, reg: $5.244309 \mathrm{e}-02,(4 / 100)$
val_acc: 0.498000, lr: 9.477776e-04, reg: 2.001293e-03, (5 / 100)
val acc: 0.469000 , lr: 1.484369e-04, reg: 4.328313e-01, ( $6 / 100$ )
val_acc: 0.522000, 1r: 5.586261e-04, reg: 2.312685e-04, (7/100)
val acc: 0.530000, $\quad \mathrm{r}: 5.808183 \mathrm{e}-04$, reg: $8.259964 \mathrm{e}-02$, $(8 / 100)$
val_acc: 0.489000, lr: 1.979168e-04, reg: 1.010889e-04, (9 / 100)
val-acc: 0.490000, lr: 2.036031e-04, reg: 2.406271e-03, (10 / 100)
val_acc: 0.475000, lr: 2.021162e-04, reg: 2.287807e-01, ( $11 / 100$ )
val_acc: 0.460000, $\operatorname{lr}: 1.135527 \mathrm{e}-04$, reg: $3.905040 \mathrm{e}-02$, ( $12 / 100$ )
val acc: $0.515000, \operatorname{lr}: 6.947668 \mathrm{e}-04, \mathrm{reg}: 1.562808 \mathrm{e}-02$, ( $13 / 100$ )
val acc: 0.531000, $\operatorname{lr}: 9.471549 \mathrm{e}-04$, reg: $1.433895 \mathrm{e}-03$, ( $14 / 100$ )
val_acc: 0.509000, lr: 3.140888e-04, reg: $2.857518 \mathrm{e}-01,(15 / 100)$
val acc: 0.514000, lr: 6.438349e-04, reg: 3.033781e-01, (16 / 100)
val acc: 0.502000, $\mathrm{lr}: 3.921784 \mathrm{e}-04$, reg: $2.707126 \mathrm{e}-04$, ( 17 / 100)
val_acc: 0.509000, lr: 9.752279e-04, reg: $2.850865 \mathrm{e}-03$, ( $18 / 100$ )
val_acc: 0.500000, $1 r: 2.412048 \mathrm{e}-04$, reg: $4.997821 \mathrm{e}-04$, ( $19 / 100$ )
val acc: $0.466000, \operatorname{lr}: 1.319314 \mathrm{e}-04, \mathrm{reg}: 1.189915 \mathrm{e}-02$, ( $20 / 100$ )
val_acc: 0.516000, $\operatorname{lr}: 8.039527 \mathrm{e}-04, \mathrm{reg}: 1.528291 \mathrm{e}-02,(21 / 100)$
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## Hyperparameter optimization

## Now run finer search...

$\max$ count $=100$
for count in xrange(max count): reg $=10 * *$ uniform $(-5,5)$ $1 r=10 * *$ uniform ( $-3,-6$ )

max count $=100$
for count in xrange(max count):
reg $=10 * *$ uniform $(-4,0)$
lr $=10 * *$ uniform $(-3,-4)$
val_acc: 0.527000, lr: 5.340517e-04, reg: 4.097824e-01, (0 / 100) val_acc: 0.492000 , lr: $2.279484 e-04$, reg: $9.991345 e-04,(1 / 100)$ val acc: 0.512000, lr: 8.680827e-04, reg: 1.349727e-02, (2 / 100) val_acc: 0.461000, lr: 1.028377e-04, reg: 1.220193e-02, (3/100) val_acc: 0.460000, $\mathrm{lr}: 1.113730 \mathrm{e}-04$, reg: $5.244309 \mathrm{e}-02,(4 / 100)$ val acc: 0.498000, lr: 9.477776e-04, reg: 2.001293e-03, (5 / 100) val acc: 0.469000, lr: 1.484369e-04, reg: 4.328313e-01, (6 / 100) val_acc: $0.522000, \operatorname{lr}: 5.586261 \mathrm{e}-04$, reg: $2.312685 \mathrm{e}-04,(7 / 100)$ val acc: 0.530000 , $\mathrm{lr}: 5.808183 \mathrm{e}-04$, reg: $8.259964 \mathrm{e}-02,(8 / 100)$ val_acc: 0.489000, lr: 1.979168e-04, reg: $1.010889 \mathrm{e}-04,(9 / 100)$ val-acc: 0.490000, lr: 2.036031e-04, reg: 2.406271e-03, ( $10 / 100$ ) val_acc: 0.475000, lr: 2.021162e-04, reg: 2.287807e-01, ( $11 / 100$ ) val_acc: 0.460000, $\mathrm{lr}: 1.135527 \mathrm{e}-04, \mathrm{reg}: 3.905040 \mathrm{e}-02,(12 / 100)$ val acc: $0.515000, \mathrm{lr}: 6.947668 \mathrm{e}-04$, reg: $1.562808 \mathrm{e}-02,(13 / 100)$ val acc: 0.531000, $\operatorname{lr}: 9.471549 \mathrm{e}-04$, reg: $1.433895 \mathrm{e}-03,(14 / 100)$ val_acc: 0.509000, $\operatorname{lr}: 3.140888 \mathrm{e}-04$, reg: $2.857518 \mathrm{e}-01,(15 / 100)$ val-acc: $0.514000, \operatorname{lr}: 6.438349 \mathrm{e}-04$, reg: 3.033781e-01, (16 / 100) val-acc: 0.502000, $\mathrm{lr}: 3.921784 \mathrm{e}-04$, reg: $2.707126 \mathrm{e}-04,(17 / 100)$ val_acc: 0.509000 , $\operatorname{lr}: 9.752279 \mathrm{e}-04$, reg: $2.850865 \mathrm{e}-03,(18 / 100)$ val_acc: 0.500000, $1 \mathrm{r}: 2.412048 \mathrm{e}-04$, reg: 4.997821e-04, ( $19 / 100$ ) val acc: $0.466000, \mathrm{lr}: 1.319314 \mathrm{e}-04, \mathrm{reg}: 1.189915 \mathrm{e}-02$, $(20 / 100)$ val_acc: $0.516000, \operatorname{lr}: 8.039527 e-04, r e g: 1.528291 e-02,(21 / 100)$

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## Hyperparameter optimization

## Random Search vs. Grid Search




Random Search for Hyper-Parameter Optimization Bergstra and Bengio, 2012

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Hyperparameter optimization

## Hyperparameters to play with: <br> - network architecture <br> - learning rate, its decay schedule, update type <br> - regularization (L2/Dropout strength)

neural networks practitioner music = loss function


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## Hyperparameter optimization

My cross-validation "command center"


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## Hyperparameter optimization

## Monitor and visualize the loss curve




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## Hyperparameter optimization



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## Hyperparameter optimization



## Hyperparameter optimization

## lossfunctions.tumblr.com Loss function specimen




Training Loss


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## Hyperparameter optimization

## lossfunctions.tumblr.com



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## Hyperparameter optimization

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## Hyperparameter optimization

## Monitor and visualize the accuracy:



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## Hyperparameter optimization

## Track the ratio of weight updates / weight magnitudes:

```
# assume parameter vector W and its gradient vector dW
param_scale = np.linalg.norm(W.ravel())
update = -learning_rate*dW # simple SGD update
update_scale = np.linalg.norm(update.ravel())
W += update # the actual update
print update_scale / param_scale # want ~1e-3
```

ratio between the values and updates: $\sim 0.0002 / 0.02=0.01$ (about okay)
want this to be somewhere around 0.001 or so

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## Conclusions (What we know in 2017)

- BP is just chain rule in calculus
- Use ReLU. Never use Sigmoid (use Tanh instead)
- Input preprocessing is no longer very important
- Do subtract mean
- Whitening and normalizing are not much needed
- Weight initialization on the other hand is extremely important for deep networks
- Use batch normalization if you can
- Use dropout
- Use Adam (or maybe RMSprop) for optimizer. If you don't have much data, can consider LBFGS
- Need to babysit your learning for real-world problems
- Never use grid search for tuning your hyperparameters


[^0]:    Fei-Fei Li \& Andrej Karpathy \& Justin Johnson Lecture 4-34 13 Jan 2016

[^1]:    Fei-Fei Li \& Andrej Karpathy \& Justin Johnson Lecture 4-35 13 Jan 2016

[^2]:    Fei-Fei Li \& Andrej Karpathy \& Justin Johnson Lecture 4-36 13 Jan 2016

[^3]:    Fei-Fei Li \& Andrej Karpathy \& Justin Johnson Lecture 4-37 13 Jan 2016

[^4]:    Fei-Fei Li \& Andrej Karpathy \& Justin Johnson Lecture 4-41 13 Jan 2016

[^5]:    Fei-Fei Li \& Andrej Karpathy \& Justin Johnson Lecture 5-39 20 Jan 2016

[^6]:    ${ }^{1}$ Note that $y^{(l)}$ now denotes the sum of input before going through the activation function.

[^7]:    ${ }^{1}$ Note that $y^{(l)}$ now denotes the sum of input before going through the activation function.

[^8]:    ${ }^{1}$ Note that $y^{(l)}$ now denotes the sum of input before going through the activation function.

[^9]:    Fei-Fei Li \& Andrej Karpathy \& Justin Johnson Lecture 5-66 20 Jan 2016

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