Neural Networks and Deep Learning

www.cs.wisc.edu/~dpage/cs760/
Goals for the lecture

you should understand the following concepts

• perceptrons
• the perceptron training rule
• linear separability
• hidden units
• multilayer neural networks
• gradient descent
• stochastic (online) gradient descent
• sigmoid function
• gradient descent with a linear output unit
• gradient descent with a sigmoid output unit
• backpropagation
Goals for the lecture

you should understand the following concepts

• weight initialization
• early stopping
• the role of hidden units
• input encodings for neural networks
• output encodings
• recurrent neural networks
• autoencoders
• stacked autoencoders
Neural networks

• a.k.a. *artificial neural networks, connectionist models*
• inspired by interconnected neurons in biological systems
  • simple processing units
  • each unit receives a number of real-valued inputs
  • each unit produces a single real-valued output
Perceptrons

[McCulloch & Pitts, 1943; Rosenblatt, 1959; Widrow & Hoff, 1960]

\[
o = \begin{cases} 
1 & \text{if } w_0 + \sum_{i=1}^{n} w_i x_i > 0 \\
0 & \text{otherwise}
\end{cases}
\]

input units: represent given \( x \)  
output unit: represents binary classification
Learning a perceptron: the perceptron training rule

1. randomly initialize weights

2. iterate through training instances until convergence

2a. calculate the output for the given instance

\[ o = \begin{cases} 
  1 & \text{if } w_0 + \sum_{i=1}^{n} w_i x_i > 0 \\
  0 & \text{otherwise}
\end{cases} \]

2b. update each weight

\[ \Delta w_i = \eta (y - o)x_i \]

\[ w_i \leftarrow w_i + \Delta w_i \]

\( \eta \) is learning rate; set to value \(< < 1 \)
Representational power of perceptrons

Perceptrons can represent only \textit{linearly separable} concepts.

$$o = \begin{cases} 
1 & \text{if } w_0 + \sum_{i=1}^{n} w_i x_i > 0 \\
0 & \text{otherwise}
\end{cases}$$

Decision boundary given by:

$$1 \text{ if } w_0 + w_1 x_1 + w_2 x_2 > 0$$

Also write as: \( \mathbf{w} \mathbf{x} > 0 \)

$$w_1 x_1 + w_2 x_2 = -w_0$$

$$x_2 = -\frac{w_1}{w_2} x_1 - \frac{w_0}{w_2}$$
Representational power of perceptrons

• in previous example, feature space was 2D so decision boundary was a line
• in higher dimensions, decision boundary is a hyperplane
Some linearly separable functions

**AND**

<table>
<thead>
<tr>
<th>$x_1$</th>
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**OR**

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XOR is not linearly separable

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A multilayer perceptron can represent XOR

Assume $w_0 = 0$ for all nodes.
Example multilayer neural network

input: two features from spectral analysis of a spoken sound

output: vowel sound occurring in the context “h__d”

figure from Huang & Lippmann, NIPS 1988
Decision regions of a multilayer neural network

input: two features from spectral analysis of a spoken sound

output: vowel sound occurring in the context “h__d”
Learning in multilayer networks

• work on neural nets fizzled in the 1960’s
  • single layer networks had representational limitations (linear separability)
  • no effective methods for training multilayer networks

• revived again with the invention of backpropagation method [Rumelhart & McClelland, 1986; also Werbos, 1975]
  • key insight: require neural network to be differentiable; use gradient descent

how to determine error signal for hidden units?
Gradient descent in weight space

Given a training set \( D = \{(x^{(1)}, y^{(1)})... (x^{(m)}, y^{(m)})\} \) we can specify an error measure that is a function of our weight vector \( w \)

\[
E(w) = \frac{1}{2} \sum_{d \in D} \left( y^{(d)} - o^{(d)} \right)^2
\]

This error measure defines a surface over the hypothesis (i.e. weight) space.
Gradient descent in weight space

gradient descent is an iterative process aimed at finding a minimum in the error surface

on each iteration
- current weights define a point in this space
- find direction in which error surface descends most steeply
- take a step (i.e. update weights) in that direction
**Gradient descent in weight space**

Calculate the gradient of $E$: \[
\nabla E(w) = \left[ \frac{\partial E}{\partial w_0}, \frac{\partial E}{\partial w_1}, \ldots, \frac{\partial E}{\partial w_n} \right]
\]

Take a step in the opposite direction:

\[
\Delta w = -\eta \ \nabla E(w)
\]

\[
\Delta w_i = -\eta \ \frac{\partial E}{\partial w_i}
\]
The sigmoid function

- to be able to differentiate $E$ with respect to $w_i$, our network must represent a continuous function
- to do this, we use sigmoid functions instead of threshold functions in our hidden and output units

$$f(x) = \frac{1}{1 + e^{-x}}$$
The sigmoid function

for the case of a single-layer network

\[ f(x) = \frac{1}{1 + e^{-\left( w_0 + \sum_{i=1}^{n} w_i x_i \right)}} \]
**Batch neural network training**

**given:** network structure and a training set \( D = \{(x^{(1)}, y^{(1)}) \ldots (x^{(m)}, y^{(m)})\} \)

initialize all weights in \( w \) to small random numbers

until stopping criteria met do

initialize the error \( E(w) = 0 \)

for each \((x^{(d)}, y^{(d)})\) in the training set

input \( x^{(d)} \) to the network and compute output \( o^{(d)} \)

increment the error \( E(w) = E(w) + \frac{1}{2}(y^{(d)} - o^{(d)})^2 \)

calculate the gradient

\[
\nabla E(w) = \left[ \frac{\partial E}{\partial w_0}, \frac{\partial E}{\partial w_1}, \ldots, \frac{\partial E}{\partial w_n} \right]
\]

update the weights

\[
\Delta w = -\eta \nabla E(w)
\]
Online vs. batch training

• Standard gradient descent (batch training): calculates error gradient for the entire training set, before taking a step in weight space

• *Stochastic gradient descent* (online training): calculates error gradient for a single instance, then takes a step in weight space
  – much faster convergence
  – less susceptible to local minima
Online neural network training
(stochastic gradient descent)

given: network structure and a training set  \( D = \{(x^{(1)}, y^{(1)}) \ldots (x^{(m)}, y^{(m)})\} \)

initialize all weights in \( w \) to small random numbers

until stopping criteria met do

for each \((x^{(d)}, y^{(d)})\) in the training set

input \( x^{(d)} \) to the network and compute output \( o^{(d)} \)

calculate the error  \( E(w) = \frac{1}{2} (y^{(d)} - o^{(d)})^2 \)

calculate the gradient

\[
\nabla E(w) = \left[ \frac{\partial E}{\partial w_0}, \frac{\partial E}{\partial w_1}, \ldots, \frac{\partial E}{\partial w_n} \right]
\]

update the weights

\[
\Delta w = -\eta \nabla E(w)
\]
Convergence of gradient descent

- gradient descent will converge to a minimum in the error function

- for a multi-layer network, this may be a local minimum (i.e. there may be a “better” solution elsewhere in weight space)

- for a single-layer network, this will be a global minimum (i.e. gradient descent will find the “best” solution)
Taking derivatives in neural nets

recall the chain rule from calculus

\[ y = f(u) \]
\[ u = g(x) \]

\[ \frac{\partial y}{\partial x} = \frac{\partial y}{\partial u} \frac{\partial u}{\partial x} \]

we’ll make use of this as follows

\[ \frac{\partial E}{\partial w_i} = \frac{\partial E}{\partial o} \frac{\partial o}{\partial net} \frac{\partial net}{\partial w_i} \]

\[ net = w_0 + \sum_{i=1}^{n} w_i x_i \]
Gradient descent: simple case

Consider a simple case of a network with one linear output unit and no hidden units:

\[ o^{(d)} = w_0 + \sum_{i=1}^{n} w_i x_i^{(d)} \]

let’s learn \( w_i \)'s that minimize squared error

\[ E(w) = \frac{1}{2} \sum_{d \in D} \left( y^{(d)} - o^{(d)} \right)^2 \]

**batch case**

\[ \frac{\partial E}{\partial w_i} = \frac{\partial}{\partial w_i} \frac{1}{2} \sum_{d \in D} \left( y^{(d)} - o^{(d)} \right)^2 \]

**online case**

\[ \frac{\partial E^{(d)}}{\partial w_i} = \frac{\partial}{\partial w_i} \frac{1}{2} \left( y^{(d)} - o^{(d)} \right)^2 \]
Stochastic gradient descent: simple case

let’s focus on the online case (stochastic gradient descent):

\[
\frac{\partial E^{(d)}}{\partial w_i} = \frac{\partial}{\partial w_i} \frac{1}{2} \left( y^{(d)} - o^{(d)} \right)^2 \\
= \left( y^{(d)} - o^{(d)} \right) \frac{\partial}{\partial w_i} \left( y^{(d)} - o^{(d)} \right) \\
= \left( y^{(d)} - o^{(d)} \right) \left( - \frac{\partial o^{(d)}}{\partial w_i} \right) \\
= - \left( y^{(d)} - o^{(d)} \right) \frac{\partial o^{(d)}}{\partial net^{(d)}} \frac{\partial net^{(d)}}{\partial w_i} \\
= - \left( y^{(d)} - o^{(d)} \right) \left( x^{(d)}_i \right)
\]
Gradient descent with a sigmoid

Now let’s consider the case in which we have a sigmoid output unit and no hidden units:

\[ \text{net}^{(d)} = w_0 + \sum_{i=1}^{n} w_i x_i^{(d)} \]

\[ o^{(d)} = \frac{1}{1 + e^{-\text{net}^{(d)}}} \]

useful property:

\[ \frac{\partial o^{(d)}}{\partial \text{net}^{(d)}} = o^{(d)} (1 - o^{(d)}) \]
Stochastic GD with sigmoid output unit

\[ \frac{\partial E^{(d)}}{\partial w_i} = \frac{\partial}{\partial w_i} \frac{1}{2} \left( y^{(d)} - o^{(d)} \right)^2 \]

\[ = \left( y^{(d)} - o^{(d)} \right) \frac{\partial}{\partial w_i} \left( y^{(d)} - o^{(d)} \right) \]

\[ = \left( y^{(d)} - o^{(d)} \right) \left( - \frac{\partial o^{(d)}}{\partial w_i} \right) \]

\[ = -\left( y^{(d)} - o^{(d)} \right) \frac{\partial o^{(d)}}{\partial net^{(d)}} \frac{\partial net^{(d)}}{\partial w_i} \]

\[ = -\left( y^{(d)} - o^{(d)} \right) o^{(d)} (1 - o^{(d)}) \frac{\partial net^{(d)}}{\partial w_i} \]

\[ = -\left( y^{(d)} - o^{(d)} \right) o^{(d)} (1 - o^{(d)}) x_i^{(d)} \]
now we’ve covered how to do gradient descent for single-layer networks with
  • linear output units
  • sigmoid output units

how can we calculate \( \frac{\partial E}{\partial w_i} \) for every weight in a multilayer network?

\( \Rightarrow \) backpropagate errors from the output units to the hidden units
Backpropagation notation

Let’s consider the online case, but drop the \(^{(d)}\) superscripts for simplicity.

We’ll use:

- Subscripts on \(y\), \(o\), \(net\) to indicate which unit they refer to.
- Subscripts to indicate the unit a weight emanates from and goes to.
Backpropagation

each weight is changed by

$$\Delta w_{ji} = -\eta \frac{\partial E}{\partial w_{ji}}$$

$$= -\eta \frac{\partial E}{\partial net_j} \frac{\partial net_j}{\partial w_{ji}}$$

$$= \eta \delta_j o_i$$

where

$$\delta_j = -\frac{\partial E}{\partial net_j}$$

$x_i$ if $i$ is an input unit
Backpropagation

Each weight is changed by

$$\Delta w_{ji} = \eta \ \delta_j \ o_i$$

where

$$\delta_j = -\frac{\partial E}{\partial net_j}$$

$$\delta_j = o_j (1 - o_j)(y_j - o_j) \quad \text{if } j \text{ is an output unit}$$

$$\delta_j = o_j (1 - o_j) \sum_k \delta_k w_{kj} \quad \text{if } j \text{ is a hidden unit}$$

Same as single-layer net with sigmoid output

Sum of backpropagated contributions to error
1. calculate error of output units
\[ \delta_j = o_j (1 - o_j) (y_j - o_j) \]

2. determine updates for weights going to output units
\[ \Delta w_{ji} = \eta \delta_j o_i \]
3. calculate error for hidden units
\[ \delta_j = o_j(1 - o_j) \sum_k \delta_k w_{kj} \]

4. determine updates for weights to hidden units using hidden-unit errors
\[ \Delta w_{ji} = \eta \delta_j o_i \]
Neural network jargon

- **activation**: the output value of a hidden or output unit

- **epoch**: one pass through the training instances during gradient descent

- **transfer function**: the function used to compute the output of a hidden/output unit from the net input
Initializing weights

• Weights should be initialized to
  • **small values** so that the sigmoid activations are in the range where the derivative is large (learning will be quicker)
  • **random values** to ensure symmetry breaking (i.e. if all weights are the same, the hidden units will all represent the same thing)
  • **typical initial weight range** [-0.01, 0.01]
Setting the learning rate

convergence depends on having an appropriate learning rate

\[ - \frac{\partial E}{\partial w_{ij}} \]

\( \eta \) too small (error goes down a little)

\( \eta \) too large (error goes up)
Stopping criteria

- conventional gradient descent: train until local minimum reached

- empirically better approach: *early stopping*
  - use a validation set to monitor accuracy during training iterations
  - return the weights that result in minimum validation-set error
Input (feature) encoding for neural networks

nominal features are usually represented using a *1-of-k* encoding

\[
A = \begin{bmatrix} 1 \\ 0 \\ 0 \\ 0 \end{bmatrix} \quad C = \begin{bmatrix} 0 \\ 1 \\ 0 \\ 0 \end{bmatrix} \quad G = \begin{bmatrix} 0 \\ 0 \\ 1 \\ 0 \end{bmatrix} \quad T = \begin{bmatrix} 0 \\ 0 \\ 0 \\ 1 \end{bmatrix}
\]

ordinal features can be represented using a *thermometer* encoding

small = \[
\begin{bmatrix} 1 \\ 0 \\ 0 \end{bmatrix}
\]
medium = \[
\begin{bmatrix} 1 \\ 1 \\ 0 \end{bmatrix}
\]
large = \[
\begin{bmatrix} 1 \\ 1 \\ 1 \end{bmatrix}
\]

real-valued features can be represented using individual input units (we may want to scale/normalize them first though)

precipitation = \[
\begin{bmatrix} 0.68 \end{bmatrix}
\]
Output encoding for neural networks

regression tasks usually use output units with linear transfer functions

binary classification tasks usually use one sigmoid output unit

$k$-ary classification tasks usually use $k$ sigmoid or softmax output units

$$O_i = \frac{e^{net_i}}{\sum_{j \in \text{outputs}} e^{net_j}}$$
Recurrent neural networks are sometimes used for tasks that involve making sequences of predictions

- Elman networks: recurrent connections go from hidden units to inputs
- Jordan networks: recurrent connections go from output units to inputs
Alternative approach to training deep networks

• use unsupervised learning to find useful hidden unit representations
Learning representations

- The feature representation provided is often the most significant factor in how well a learning system works.

- An appealing aspect of multilayer neural networks is that they are able to change the feature representation.

- Can think of the nodes in the hidden layer as new features constructed from the original features in the input layer.

- Consider having more levels of constructed features, e.g., pixels -> edges -> shapes -> faces or other objects.
Competing intuitions

• Only need a 2-layer network (input, hidden layer, output)
  – Representation Theorem (1989): Using sigmoid activation functions (more recently generalized to others as well), can represent any continuous function with a single hidden layer
  – Empirically, adding more hidden layers does not improve accuracy, and it often degrades accuracy, when training by standard backpropagation

• Deeper networks are better
  – More efficient representationally, e.g., can represent $n$-variable parity function with polynomially many (in $n$) nodes using multiple hidden layers, but need exponentially many (in $n$) nodes when limited to a single hidden layer
  – More structure, should be able to construct more interesting derived features
The role of hidden units

- Hidden units transform the input space into a new space where perceptrons suffice
- They numerically represent “constructed” features
- Consider learning the target function using the network structure below:

<table>
<thead>
<tr>
<th>Input</th>
<th>Output</th>
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<tbody>
<tr>
<td>100000000</td>
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<tr>
<td>00000001</td>
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</table>
The role of hidden units

- In this task, hidden units learn a compressed numerical coding of the inputs/outputs

<table>
<thead>
<tr>
<th>Input</th>
<th>Hidden Values</th>
<th>Output</th>
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<tbody>
<tr>
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<td>→ 0.89 0.04 0.08 → 100000000</td>
<td></td>
</tr>
<tr>
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<td>001000000</td>
<td>→ 0.01 0.97 0.27 → 001000000</td>
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<td>000100000</td>
<td>→ 0.99 0.97 0.71 → 000100000</td>
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<tr>
<td>000010000</td>
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<tr>
<td>000000001</td>
<td>→ 0.60 0.94 0.01 → 000000001</td>
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How many hidden units should be used?

- conventional wisdom in the early days of neural nets: prefer small networks because fewer parameters (i.e. weights & biases) will be less likely to overfit
- somewhat more recent wisdom: if early stopping is used, larger networks often behave as if they have fewer “effective” hidden units, and find better solutions

Figure from Weigend, *Proc. of the CMSS 1993*
Another way to avoid overfitting

• Allow many hidden units but force each hidden unit to output mostly zeroes: tend to meaningful concepts

• Gradient descent solves an optimization problem—add a “regularizing” term to the objective function

• Let $X$ be vector of random variables, one for each hidden unit, giving average output of unit over data set. Let target distribution $s$ have variables independent with low probability of outputting one (say 0.1), and let $\hat{s}$ be empirical distribution in the data set. Add to the backpropagation target function (that minimizes $\delta$’s) a penalty of $\text{KL}(s(X) || \hat{s}(X))$
Backpropagation with multiple hidden layers

- In principle, backpropagation can be used to train arbitrarily deep networks (i.e. with multiple hidden layers).

- In practice, this doesn’t usually work well.

  - There are likely to be lots of local minima.

  - Diffusion of gradients leads to slow training in lower layers.

    - Gradients are smaller, less pronounced at deeper levels.

    - Errors in credit assignment propagate as you go back.
Autoencoders

- one approach: use *autoencoders* to learn hidden-unit representations
- in an autoencoder, the network is trained to reconstruct the inputs
Autoencoder variants

• how to encourage the autoencoder to generalize

  • bottleneck: use fewer hidden units than inputs

  • sparsity: use a penalty function that encourages most hidden unit activations to be near 0 [Goodfellow et al. 2009]

  • denoising: train to predict true input from corrupted input [Vincent et al. 2008]

  • contractive: force encoder to have small derivatives (of hidden unit output as input varies) [Rifai et al. 2011]
Stacking Autoencoders

- can be stacked to form highly nonlinear representations [Bengio et al. NIPS 2006]

Train autoencoder to represent $x$

Discard output layer; train autoencoder to represent $h_1$

Repeat for $k$ layers

discard output layer; train weights on last layer for supervised task

Each $W_i$ here represents the matrix of weights between layers
Fine-Tuning

• After completion, run backpropagation on the entire network to fine-tune weights for the supervised task.

• Because this backpropagation starts with good structure and weights, its credit assignment is better and so its final results are better than if we just ran backpropagation initially.
Why does the unsupervised training step work well?

- *regularization hypothesis*: representations that are good for $P(x)$ are good for $P(y \mid x)$

- *optimization hypothesis*: unsupervised initializations start near better local minima of supervised training error
Deep learning not limited to neural networks

- First developed by Geoff Hinton and colleagues for *belief networks*, a kind of hybrid between neural nets and Bayes nets
- Hinton motivates the unsupervised deep learning training process by the credit assignment problem, which appears in belief nets, Bayes nets, neural nets, restricted Boltzmann machines, etc.
  - d-separation: the problem of evidence at a converging connection creating competing explanations
  - backpropagation: can’t choose which neighbors get the blame for an error at this node
Room for Debate

• many now arguing that unsupervised *pre-training* phase not really needed…

• backprop is sufficient if done better
  – wider diversity in initial weights, try with many initial settings until you get learning
  – don’t worry much about exact learning rate, but add *momentum*: if moving fast in a given direction, keep it up for awhile
  – *Need a lot of data for deep net backprop*
Dropout training

• On each training iteration, drop out (ignore) 90% of the units (or other %)

• Ignore for forward & backprop (all training)

Figures from Srivastava et al., *Journal of Machine Learning Research* 2014

(a) Standard Neural Net  
(b) After applying dropout.
At Test Time

- Final model uses all nodes
- Multiply each weight from a node by fraction of times node was used during training

Figures from Srivastava et al., *Journal of Machine Learning Research* 2014
Some Deep Learning Resources

• *Nature*, Jan 8, 2014: http://www.nature.com/news/computer-science-the-learning-machines-1.14481

• Ng Tutorial: http://deeplearning.stanford.edu/wiki/index.php/UFLDL_Tutorial

• Hinton Tutorial: http://videolectures.net/jul09_hinton_deeplearn/

Comments on neural networks

- stochastic gradient descent often works well for very large data sets

- backpropagation generalizes to
  - arbitrary numbers of output and hidden units
  - arbitrary layers of hidden units (in theory)
  - arbitrary connection patterns
  - other transfer (i.e. output) functions
  - other error measures

- backprop doesn’t usually work well for networks with multiple layers of hidden units; recent work in deep networks addresses this limitation