## Learning II

## Linda Shapiro

ECE/CSE 576

## More on Learning

- Neural Nets
- Support Vectors Machines
- Unsupervised Learning (Clustering)
- K-Means
- Expectation-Maximization


## Neural Net Learning

- Motivated by studies of the brain.
- A network of "artificial neurons" that learns a function.
- Doesn't have clear decision rules like decision trees, but highly successful in many different applications. (e.g. face detection)
- We use them frequently in our research.
- I'll be using algorithms from http://www.cs.mtu.edu/~nilufer/classes/cs4811/2016-spring/lecture-slides/cs4811-neural-net-algorithms.pdf
$10^{11}$ neurons of $>20$ types, $10^{14}$ synapses, $1 \mathrm{~ms}-10 \mathrm{~ms}$ cycle time Signals are noisy "spike trains" of electrical potential


Output is a "squashed" linear function of the inputs:

$$
a_{i} \leftarrow g\left(i n_{i}\right)=g\left(\sum_{j} W_{j, i} a_{j}\right)
$$



A gross oversimplification of real neurons, but its purpose is to develop understanding of what networks of simple units can do

## Activation functions


(a)

(b)
(a) is a step function or threshold function
(b) is a sigmoid function $1 /\left(1+e^{-x}\right)$

Changing the bias weight $W_{0, i}$ moves the threshold location

## Simple Feed-Forward Perceptrons


in $=\left(\sum W_{j} x_{j}\right)+\theta$
out $=g[$ in $]$
$g$ is the activation function
It can be a step function:
$g(x)=1$ if $x>=0$ and $0($ or -1$)$ else.
It can be a sigmoid function:
$g(x)=1 /(1+\exp (-x))$.

The sigmoid function is differentiable and can be used in a gradient descent algorithm to update the weights.


## Gradient Descent

takes steps proportional to the negative of the gradient of a function to find its local minimum

- Let $\mathbf{X}$ be the inputs, y the class, $\mathbf{W}$ the weights
- in $=\sum W_{j} x_{j}$
- $E r r=y-g(i n)$
- $\mathrm{E}=1 / 2 \mathrm{Err}{ }^{2}$ is the squared error to minimize
- $\partial \mathrm{E} / \partial \mathrm{W}_{\mathrm{j}}=\mathrm{Err} * \partial \mathrm{Err} / \partial \mathrm{W}_{\mathrm{j}}=\mathrm{Err} * \partial / \partial \mathrm{W}_{\mathrm{j}}(\mathrm{g}(\mathrm{in}))(-1)$
- $=-E r r{ }^{*} \mathrm{~g}^{\prime}(\mathrm{in})^{*} \mathrm{x}_{\mathrm{j}}$
- The update is $W_{j}<-W_{j}+\alpha^{*} E r r$ * $g^{\prime}(i n)^{*} x_{j}$
- $\alpha$ is called the learning rate.


## Simple Feed-Forward Perceptrons



```
repeat
    for each e in examples do
        in=(\sum W ( }\mp@subsup{\textrm{x}}{\textrm{j}}{})+
    Err = y[e]-g[in]
    W}\mp@subsup{\textrm{W}}{\textrm{j}}{=}\mp@subsup{\textrm{W}}{\textrm{j}}{}+\alpha\operatorname{Errg}\mp@subsup{g}{}{\prime}(\textrm{in})\mp@subsup{\textrm{x}}{\textrm{j}}{[e]
    until done
```

Examples: $\mathrm{A}=[(.5,1.5),+1], \mathrm{B}=[(-.5, .5),-1], \mathrm{C}=[(.5, .5),+1]$
Initialization: $W_{1}=1, W_{2}=2, \theta=-2$

Note1: when g is a step function, the $\mathrm{g}^{\prime}(\mathrm{in})$ is removed. Note2: later in back propagation, Err ${ }^{*} \mathrm{~g}^{\prime}(\mathrm{in})$ will be called $\Delta$ We'll let $\mathrm{g}(\mathrm{x})=1$ if $\mathrm{x}>=0$ else -1

## Graphically

Examples: $\mathrm{A}=[(.5,1.5),+1], \mathrm{B}=[(-.5, .5),-1], \mathrm{C}=[(.5, .5),+1]$ Initialization: $W_{1}=1, W_{2}=2, \theta=-2$


Examples:

$$
\begin{aligned}
& A=[(.5,1.5),+1], \\
& B=[(-.5, .5),-1], \\
& C=[(.5, .5),+1]
\end{aligned}
$$

Initialization: $W_{1}=1, W_{2}=2, \theta=-2$

## Learning

repeat
for each e in examples do
in $=\left(\Sigma W_{j} \mathrm{x}_{\mathrm{j}}\right)+\theta$
Err $=y[e]-g[i n]$
$\mathrm{W}_{\mathrm{j}}=\mathrm{W}_{\mathrm{j}}+\alpha \operatorname{Errg}^{\prime}(\mathrm{in}) \mathrm{x}_{\mathrm{j}}[\mathrm{e}]$
until done

```
A=[(.5,1.5),+1]
in =.5(1)+(1.5)(2)-2 = 1.5
g(in) = 1; Err = 0; NO CHANGE
B=[(-.5,.5),-1]
In = (-.5)(1) + (.5)(2) -2 = -1.5
g(in) =-1; Err = 0; NO CHANGE
```

$$
\begin{aligned}
& \mathrm{C}=[(.5, .5),+1] \\
& \text { in }=(.5)(1)+(.5)(2)-2=-.5 \\
& \mathrm{~g}(\text { in })=-1 ; \text { Err }=1-(-1)=2
\end{aligned}
$$

## Graphically

Examples: $\mathrm{A}=[(.5,1.5),+1], \mathrm{B}=[(-.5, .5),-1], \mathrm{C}=[(.5, .5),+1]$ Initialization: $W_{1}=1, W_{2}=2, \theta=-2$


## Back Propagation

- Simple single layer networks with feed forward learning were not powerful enough.
- Could only produce simple linear classifiers.
- More powerful networks have multiple hidden layers.
- The learning algorithm is called back propagation, because it computes the error at the end and propagates it back through the weights of the network to the beginning.


## The backpropagation algorithm

The following is the backpropagation algorithm for learning in multilayer networks.
function BACK-PROP-LEARNING(examples, network)
returns a neural network

## inputs:

examples, a set of examples, each with input vector $\mathbf{x}$ and output vector $\mathbf{y}$. network, a multilayer network with $L$ layers, weights $W_{j, i}$, activation function $g$
local variables: $\Delta$, a vector of errors, indexed by network node
for each weight $w_{i, j}$ in network do
$w_{i, j} \leftarrow$ a small random number
repeat
for each example ( $\mathbf{x}, \mathbf{y}$ ) in examples do

## Let's break it into steps.

/* Propagate the inputs forward to compute the outputs. */
for each node $i$ in the input layer do // Simply copy the input values. $a_{i} \leftarrow x_{i}$
for $l=2$ to $L$ do $\quad / /$ Feed the values forward.
for each node $j$ in layer $l$ do

$$
\begin{aligned}
& i n_{j} \leftarrow \sum_{i} w_{i, j} a_{i} \\
& a_{j} \leftarrow g\left(i n_{j}\right)
\end{aligned}
$$

for each node $j$ in the output layer do // Compute the error at the output.

$$
\Delta[j] \leftarrow g^{\prime}\left(i n_{j}\right) \times\left(y_{j}-a_{j}\right)
$$

/* Propagate the deltas backward from output layer to input layer */
for $l=L-1$ to 1 do
for each node $i$ in layer $l$ do

$$
\Delta[i] \leftarrow g^{\prime}\left(i n_{i}\right) \sum_{j} w_{i, j} \Delta[j] \quad / / \text { "Blame" a node as much as its wei§ }
$$

/* Update every weight in network using deltas. */
for each weight $w_{i, j}$ in network do

$$
w_{i, j} \leftarrow w_{i, j}+\alpha \times a_{i} \times \Delta[j] \quad \text { // Adjust the weights. }
$$

until some stopping criterion is satisfied

```
return network
```


## The backpropagation algorithm

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for each weight $w_{i, j}$ in network do $w_{i, j} \leftarrow$ a small random number


## Forward Computation

## repeat

for each example ( $\mathbf{x}, \mathbf{y}$ ) in examples do
/* Propagate the inputs forward to compute the outputs. */
for each node $i$ in the input layer do Simply copy the input values.

$$
\begin{aligned}
& a_{i} \leftarrow x_{i} \\
& \text { for } l=2 \text { to } L \text { do } \\
& \text { for each node } j \text { in layer } l \text { do } \\
& \quad i n_{j} \leftarrow \sum_{i} w_{i, j} a_{i} \\
& \quad a_{j} \leftarrow g\left(i n_{j}\right)
\end{aligned}
$$



## Backward Propagation 1

for each node $j$ in the output layer do // Compute the error at the output.

$$
\Delta[j] \leftarrow g^{\prime}\left(i n_{j}\right) \times\left(y_{j}-a_{j}\right)
$$

- Node nf is the only node in our output layer.
- Compute the error at that node and multiply by the derivative of the weighted input sum to get the change delta.



## Backward Propagation 2

/* Propagate the deltas backward from output layer to input layer */
for $l=L-1$ to 1 do
for each node $i$ in layer $l$ do

$$
\Delta[i] \leftarrow g^{\prime}\left(i n_{i}\right) \sum_{j} w_{i, j} \Delta[j] \quad / / \text { "Blame" a node as much as its wei } \S
$$

- At each of the other layers, the deltas use
- the derivative of its input sum
- the sum of its output weights
- the delta computed for the output error



## Backward Propagation 3

/* Update every weight in network using deltas. */
for each weight $w_{i, j}$ in network do

$$
w_{i, j} \leftarrow w_{i, j}+\alpha \times a_{i} \times \Delta[j] \quad \text { // Adjust the weights. }
$$

Now that all the deltas are defined, the weight updates just use them.


## Back Propagation Summary

- Compute delta values for the output units using observed errors.
- Starting at the output-1 layer
- repeat
- propagate delta values back to previous layer
- update weights between the two layers
- till done with all layers
- This is done for all examples and multiple epochs, till convergence or enough iterations.

Time taken to build model: 16.2 seconds

| Correctly Classified Instances | 307 | $80.3665 \%$ (did not boost) |
| :--- | :---: | :--- |
| Incorrectly Classified Instances | 75 | $19.6335 \%$ |
| Kappa statistic | 0.6056 |  |
| Mean absolute error | 0.1982 |  |
| Root mean squared error | 0.41 |  |
| Relative absolute error | $39.7113 \%$ |  |
| Root relative squared error | $81.9006 \%$ |  |
| Total Number of Instances | 382 |  |



## Handwritten digit recognition



3-nearest-neighbor $=2.4 \%$ error
400-300-10 unit MLP $=1.6 \%$ error
LeNet: 768-192-30-10 unit MLP $=0.9 \%$ error
Current best (kernel machines, vision algorithms) $\approx 0.6 \%$ error

## Kernel Machines

- A relatively new learning methodology (1992) derived from statistical learning theory.
- Became famous when it gave accuracy comparable to neural nets in a handwriting recognition class.
- Was introduced to computer vision researchers by Tomaso Poggio at MIT who started using it for face detection and got better results than neural nets.
- Has become very popular and widely used with packages available.


## Support Vector Machines (SVM)

- Support vector machines are learning algorithms that try to find a hyperplane that separates the different classes of data the most.
- They are a specific kind of kernel machines based on two key ideas:
- maximum margin hyperplanes
- a kernel 'trick'


## The SVM Equation

- $\mathrm{y}_{\mathrm{SVM}}\left(\mathrm{x}_{\mathrm{q}}\right)=\operatorname{argmax} \sum \mathrm{a}_{\mathrm{i}, \mathrm{C}} \mathrm{K}\left(\mathrm{x}_{\mathrm{i}}, \mathrm{x}_{\mathrm{q}}\right)$

$$
\text { c } \quad i=1, m
$$

- $x_{q}$ is a query or unknown object
- c indexes the classes
- there are m support vectors $x_{i}$ with weights $\alpha_{i, c}, i=1$ to $m$ for class $c$
- $K$ is the kernel function that compares $x_{i}$ to $X_{q}$
*** This is for multiple class SVMs with support vectors for every class; we'll see a simpler equation for 2 class.


## Maximal Margin (2 class problem)

| In 2D space, |
| :--- |
| a hyperplane is |
| a line. |
| In 3D space, |
| it is a plane. |



Find the hyperplane with maximal margin for all the points. This originates an optimization problem which has a unique solution.

## Support Vectors

- The weights $\alpha_{i}$ associated with data points are zero, except for those points closest to the separator.
- The points with nonzero weights are called the support vectors (because they hold up the separating plane).
- Because there are many fewer support vectors than total data points, the number of parameters defining the optimal separator is small.


## A Geometric Interpretation



## Kernels

- A kernel is just a similarity function. It takes 2 inputs and decides how similar they are.
- Kernels offer an alternative to standard feature vectors. Instead of using a bunch of features, you define a single kernel to decide the similarity between two objects.


## Kernels and SVMs

- Under some conditions, every kernel function can be expressed as a dot product in a (possibly infinite dimensional) feature space (Mercer's theorem)
- SVM machine learning can be expressed in terms of dot products.
- So SVM machines can use kernels instead of feature vectors.


## The Kernel Trick

The SVM algorithm implicitly maps the original data to a feature space of possibly infinite dimension in which data (which is not separable in the original space) becomes separable in the feature space.



Feature space $\mathrm{R}^{\mathrm{n}}$


## Kernel Functions

- The kernel function is designed by the developer of the SVM.
- It is applied to pairs of input data to evaluate dot products in some corresponding feature space.
- Kernels can be all sorts of functions including polynomials and exponentials.
- Simplest is just the plain dot product: $\mathrm{xi} \bullet \times \mathrm{j}$
- The polynomial kernel $K(x i, x j)=(x i \bullet x j+1)^{p}$, where $p$ is a tunable parameter.


## Kernel Function used in our 3D Computer Vision Work

- $\mathrm{k}(\mathrm{A}, \mathrm{B})=\exp \left(-\theta_{\mathrm{AB}}^{2} / \sigma^{2}\right)$
- $A$ and $B$ are shape descriptors (big vectors).
- $\theta$ is the angle between these vectors.
- $\sigma^{2}$ is the "width" of the kernel.



## What does SVM learning solve?

- The SVM is looking for the best separating plane in its alternate space.
- It solves a quadratic programming optimization problem $\underset{\alpha}{\operatorname{argmax}} \sum_{j} \alpha_{j}-1 / 2 \sum_{j, k} \alpha_{j} \alpha_{k} y_{j} y_{k}\left(\mathbf{x}_{j} \bullet \mathbf{x}_{k}\right)$ subject to $\alpha_{j}>0$ and $\sum \alpha_{j} y_{j}=0$.
- The equation for the separator for these optimal $\alpha_{j}$ is

$$
h(x)=\operatorname{sign}\left(\sum_{j} \alpha_{j} y_{j}\left(x \cdot x_{j}\right)-b\right)
$$

## Simple Example of Classification

- $K(A, B)=A \bullet B$
- known positive class points $\{(3,1),(3,-1),(6,1),(6,-1)\}$
- known negative class points $\{(1,0),(0,1),(0,-1),(-1,0)\}$
- support vectors: $s=\{(1,0),(3,1), 3,-1)\}$ with weights $\alpha$
= 3.5, .75, . 75
- classifier equation: $f(x)=\operatorname{sign}\left(\Sigma_{i}\left[\alpha_{i}{ }^{*} K\left(s_{i}, x\right)\right]-b\right) \quad b=2$


$$
\begin{aligned}
& f(1,1)=\operatorname{sign}\left(\sum_{i} \alpha_{i} s_{i} \bullet(1,1) \quad-2\right) \\
&= \operatorname{sign}\left(.75^{*}(3,1) \bullet(1,1)+.75^{*}(3,-1) \bullet(1,1)+(-3.5)^{*}(1,0) \bullet(1,1)-2\right) \\
&= \operatorname{sign}(1-2)=\operatorname{sign}(-1)=- \text { negative class } \\
& \text { CORRECT }
\end{aligned}
$$

Time taken to build model: 0.15 seconds


## Unsupervised Learning

- Find patterns in the data.
- Group the data into clusters.
- Many clustering algorithms.
- K means clustering
- EM clustering
- Graph-Theoretic Clustering
- Clustering by Graph Cuts
- etc


## Clustering by K-means Algorithm

Form K-means clusters from a set of $n$-dimensional feature vectors

1. Set ic (iteration count) to 1
2. Choose randomly a set of $K$ means $m_{l}(1), \ldots, m_{K}(1)$.
3. For each vector $x_{i}$, compute $D\left(x_{i} m_{k}(i c)\right), k=1, \ldots K$ and assign $x_{i}$ to the cluster $C_{j}$ with nearest mean.
4. Increment $i c$ by 1 , update the means to get $m_{l}(i c), \ldots, m_{K}(i c)$.
5. Repeat steps 3 and 4 until $C_{k}(i c)=C_{k}(i c+1)$ for all $k$.


## K-Means Classifier (shown on RGB color data)


original data
one RGB per pixel

color clusters

## K-Means $\rightarrow$ EM

The clusters are usually Gaussian distributions.

- Boot Step:
- Initialize $K$ clusters: $\mathrm{C}_{1}, \ldots, C_{K}$

$$
\left(\mu_{j}, \Sigma_{j}\right) \text { and } P\left(C_{j}\right) \text { for each cluster } j .
$$



- Iteration Step:
- Estimate the cluster of each datum

$$
p\left(C_{j} \mid x_{i}\right)
$$

$\Rightarrow$ Expectation

- Re-estimate the cluster parameters

$$
\left(\mu_{j}, \Sigma_{j}\right), p\left(C_{j}\right) \quad \text { For each cluster } j
$$

$\Rightarrow$ Maximization
The resultant set of clusters is called a mixture model; if the distributions are Gaussian, it's a Gaussian mixture.

## EM A gorithn Sunnnary

- Boot Step:
- Initialize $K$ clusters: $\mathrm{C}_{1}, \ldots, C_{K}$

$$
\left(\mu_{j}, \Sigma_{j}\right) \text { and } p\left(C_{j}\right) \text { for each cluster } j \text {. }
$$

- Iteration Step:

$$
\begin{aligned}
& \text { Normal Probability Density Function } \\
& F(x)=\frac{1}{\sigma \sqrt{2 \pi}} e^{-(x-\mu)^{2} / 2 \sigma^{2}}
\end{aligned}
$$

$$
f(\mathbf{x})=\frac{1}{\sqrt{(2 \pi)^{k}|\boldsymbol{\Sigma}|}} \exp \left(-\frac{1}{2}(\mathbf{x}-\boldsymbol{\mu})^{\mathrm{T}} \boldsymbol{\Sigma}^{-1}(\mathbf{x}-\boldsymbol{\mu})\right)
$$

- Expectation Step

$$
\begin{aligned}
& p\left(C_{j} \mid x_{i}\right)=\frac{p\left(x_{i} \mid C_{j}\right) \cdot p\left(C_{j}\right)}{p\left(x_{i}\right)}=\frac{p\left(x_{i} \mid C_{j}\right) \cdot p\left(C_{j}\right)}{\sum_{j} p\left(x_{i} \mid C_{j}\right) \cdot p\left(C_{j}\right)} \\
& \text { Maximization Step }
\end{aligned}
$$

- Maximization Step

$$
\mu_{j}=\frac{\sum_{i} p\left(C_{j} \mid x_{i}\right) \cdot x_{i}}{\sum_{i} p\left(C_{j} \mid x_{i}\right)} \quad \Sigma_{j}=\frac{\sum_{i} p\left(C_{j} \mid x_{i}\right) \cdot\left(x_{i}-\mu_{j}\right) \cdot\left(x_{i}-\mu_{j}\right)^{T}}{\sum_{i} p\left(C_{j} \mid x_{i}\right)}
$$

$$
p\left(C_{j}\right)=\frac{\sum_{i} p\left(C_{j} \mid x_{i}\right)}{N}
$$

## EM Clustering using color and texture information at each pixel

(from Blobworld)


## EM for Classification of Images in Terms of their Color Regions

Initial Model for "trees"


Initial Model for "sky"


Final Model for "trees"


Final Model for "sky"


## Sample Results

## cheetah



## Sample Results (Cont.)

## grass



## Sample Results (Cont.)

## lion



# Haar Random Forest Features Combined with a Spatial Matching Kernel for Stonefly Species Identification 

Natalia Larios*<br>Bilge Soran*<br>Linda Shapiro*<br>Gonzalo Martinez-Munoz^<br>Jeffrey Lin+<br>Tom Dietterich+<br>*University of Washington<br>+Oregon State University<br>^Universidad Autónoma de Madrid

Goal: to identify the species of insect specimens rapidly and accurately


## Haar-like Features



> Value =
> $\Sigma$ (pixels in white area) -
> $\Sigma$ (pixels in black area)

There can be hundreds or even thousands of them.

So the classifier has to choose.

## Overview of our Classification Method



## RESULTS:

Stonefly Identification: Classification Error [\%]

| Task | SET | CIELAB <br> color | CIELAB+G |
| :---: | :---: | :---: | :---: |
| Cal vs Dor | 6.26 | 10.16 | $4.60 \quad 96.4 \%$ accuracy |
| Hes vs Iso | 3.74 | 9.05 | 3.55 |
| Pte vs Swe | 2.71 | 8.75 | 2.80 |
| Dor vs Hes | 2.25 | 8.09 | 2.20 |
| Mos vs Pte | 2.06 | 7.95 | 1.92 |
| Yor vs Zap | 1.52 | 6.89 | 1.60 |
| Zap vs Cal | 1.52 | 7.02 | 1.76 |
| Swe vs Yor | 1.44 | 6.85 | 1.50 |
| Iso vs Mos | 1.29 | 6.90 | 1.30 |
| Average | $\mathbf{2 . 5 3}$ | $\mathbf{7 . 9 6}$ | 2.25 |

## Finale

- We have looked at
- decision trees
- random decision forests
- boosting (and other metaclassifiers)
- neural nets
- SVMs
- unsupervised learning
- We will now go into object recognition of different types, ending with deep neural nets called Convolutional Neural Networks.

