#### CIS 519/419 Applied Machine Learning www.seas.upenn.edu/~cis519

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Slides were created by Dan Roth (for CIS519/419 at Penn or CS446 at UIUC), Eric Eaton for CIS519/419 at Penn, or from other authors who have made their ML slides available. CIS419/519 Spring '18

#### Administration

Registration <sup>(C)</sup>

#### Questions

- Hw1 is due next week
  - You should have started working on it already...
- Hw2 will be out next week
- No lecture on Tuesday next Week (2/6)!!

#### Projects

- CIS 519 students need to do a team project
  - Teams will be of size 2-3
- Projects proposals are due on Friday 3/2/18
  - Details will be available on the website
  - We will give comments and/or requests to modify / augment/ do a different project.
  - There may also be a mechanism for peer comments.
- Please start thinking and working on the project now.
  - Your proposal is limited to 1-2 pages, but needs to include references and, ideally, some preliminary results/ideas.
- Any project with a significant Machine Learning component is good.
  - Experimental work, theoretical work, a combination of both or a critical survey of results in some specialized topic.
  - The work has to include some reading of the literature .
  - Originality is not mandatory but is encouraged.
- Try to make it interesting!

#### **Project Examples**

- KDD Cup 2013:
  - "Author-Paper Identification": given an author and a small set of papers, we are asked to identify which papers are really written by the author.
    - https://www.kaggle.com/c/kdd-cup-2013-author-paper-identification-challenge
  - "Author Profiling": given a set of document, profile the author: identification, gender, native language, ....
- Caption Control: Is it gibberish? Spam? High quality text?
  - Adapt an NLP program to a new domain
- Work on making learned hypothesis more comprehensible
  - Explain the prediction
- Develop a (multi-modal) People Identifier
- Identify contradictions in news stories
- Large scale clustering of documents + name the cluster
  - E.g., cluster news documents and give a title to the document
- Deep Neural Networks: convert a state of the art NLP program to a NN

#### This Lecture

- Decision trees for (binary) classification
  - Non-linear classifiers
- Learning decision trees (ID3 algorithm)
  - Greedy heuristic (based on information gain)
     Originally developed for discrete features
  - Some extensions to the basic algorithm
- Overfitting
  - Some experimental issues

# A Guide

- Learning Algorithms
  - (Stochastic) Gradient Descent (with LMS)
  - Decision Trees
  - Importance of hypothesis space (representation)
- How are we doing?
  - Quantification in terms of cumulative # of mistakes
  - Our algorithms were driven by a different metric than the one we care about.
- Today: Versions of Perceptron
  - How to deal better with large features spaces & sparsity?
  - Variations of Perceptron
    - Dealing with overfitting
  - Closing the loop: Back to Gradient Descent
  - Dual Representations & Kernels
- Multilayer Perceptron
- Beyond Binary Classification?
  - Multi-class classification and Structured Prediction
- More general way to quantify learning performance (PAC)
  - New Algorithms (SVM, Boosting)

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#### Today:

Take a more general perspective and think more about learning, learning protocols, quantifying performance, etc.

This will motivate some of the ideas we will see next.

## **Quantifying Performance**

- We want to be able to say something rigorous about the performance of our learning algorithm.
- We will concentrate on discussing the number of examples one needs to see before we can say that our learned hypothesis is good.

#### Learning Conjunctions

 There is a hidden (monotone) conjunction the learner (you) is to learn

 $f(x_1, x_2, ..., x_{100}) = x_2 \land x_3 \land x_4 \land x_5 \land x_{100}$ 

- How many examples are needed to learn it ? How ?
  - Protocol I: The learner proposes instances as queries to the teacher
  - Protocol II: The teacher (who knows f) provides training examples
  - Protocol III: Some random source (e.g., Nature) provides training examples; the Teacher (Nature) provides the labels (f(x))

- Protocol I: The learner proposes instances as queries to the teacher
- Since we know we are after a monotone conjunction:
- Is x<sub>100</sub> in? <(1,1,1...,1,0), ?> f(x)=0 (conclusion: Yes)
- Is x<sub>99</sub> in? <(1,1,...1,0,1), ?> f(x)=1 (conclusion: No)
- Is x<sub>1</sub> in ? <(0,1,...1,1,1), ?> f(x)=1 (conclusion: No)
- A straight forward algorithm requires n=100 queries, and will produce as a result the hidden conjunction (exactly).

• 
$$h(x_1, x_2, ..., x_{100}) = x_2 \wedge x_3 \wedge x_4 \wedge x_5 \wedge x_{100}$$

What happens here if the conjunction is not known to be monotone? If we know of a positive example, the same algorithm works.

Protocol II: The teacher (who knows f) provides training examples

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- To show you that all these variables are required...

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- To show you that all these variables are required...
  - <(0,0,1,1,1,0,...,0,1), 0> need x<sub>2</sub>
  - <(0,1,0,1,1,0,...,0,1), 0> need x<sub>3</sub>

Modeling Teaching Is tricky

- .....
- <(0,1,1,1,1,0,...,0,0), 0> need x<sub>100</sub>
- A straight forward algorithm requires k = 6 examples to produce the hidden conjunction (exactly).

$$h(x_1, x_2, ..., x_{100}) = x_2 \wedge x_3 \wedge x_4 \wedge x_5 \wedge x_{100}$$

 $f = x_2 \land x_3 \land x_4 \land x_5 \land x_{100}$ 

- Protocol III: Some random source (e.g., Nature) provides training examples
- Teacher (Nature) provides the labels (f(x))
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  - (1,1,1,0,0,0,...,0,0), 0>
  - (1,1,1,1,1,0,...0,1,1), 1>
  - (1,0,1,1,1,0,...0,1,1), 0>
  - (1,1,1,1,1,0,...0,0,1), 1>
  - (1,0,1,0,0,0,...0,1,1), 0>
  - (1,1,1,1,1,1,...,0,1), 1>
  - (0,1,0,1,0,0,...0,1,1), 0>
- How should we learn?
- Skip

- Protocol III: Some random source (e.g., Nature) provides training examples
  - Teacher (Nature) provides the labels (f(x))
- Algorithm: Elimination
  - Start with the set of all literals as candidates
  - Eliminate a literal that is not active (0) in a positive example

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  - <(1,1,1,0,0,0,...,0,0), 0>  $\leftarrow$  learned nothing: h= x<sub>1</sub> ^ x<sub>2</sub> ,..., ^ x<sub>100</sub>
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  - <(1,1,1,1,1,0,...0,1,1), 1>  $\leftarrow$  h= x<sub>1</sub>  $\wedge$  x<sub>2</sub>  $\wedge$  x<sub>3</sub>  $\wedge$  x<sub>4</sub>  $\wedge$  x<sub>5</sub>  $\wedge$  x<sub>99</sub>  $\wedge$  x<sub>100</sub>

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  - (1,0,1,0,0,0,...0,1,1), 0>
  - (1,1,1,1,1,1,...,0,1), 1>
- <(0,1,0,1,0,0,...0,1,1), 0>CIS419/519 Spring '18

 $f = x_2 \wedge x_3 \wedge x_4 \wedge x_5 \wedge x_{100}$ 

- Protocol III: Some random source (e.g., Nature) provides training examples
  - Teacher (Nature) provides the labels (f(x))
- Algorithm: Elimination

Is it good

- Performance ?
- # of examples ?
- Start with the set of all literals as candidates
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- <(1,1,1,1,1,1,...,1,1), 1>
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<(0,1,0,1,0,0,...0,1,1), 0> CIS419/519 Spring '18

- Final hypothesis:

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#### Learning Conjunctions (III)

- Protocol III: Some random source (e.g., Nature) provides training examples
  - Teacher (Nature) provides the labels (f(x))
- Algorithm: .....
  - (1,1,1,1,1,1,...,1,1), 1>
  - <(1,1,1,0,0,0,...,0,0), 0>
  - (1,1,1,1,1,0,...0,1,1), 1>
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  - <(1,0,1,0,0,0,...0,1,1), 0> Final hypothesis:
  - <(1,1,1,1,1,1,1,...,0,1), 1> h= x<sub>1</sub>  $\land$  x<sub>2</sub>  $\land$  x<sub>3</sub>  $\land$  x<sub>4</sub>  $\land$  x<sub>5</sub>  $\land$  x<sub>100</sub>
  - <(0,1,0,1,0,0,...0,1,1), 0>
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#### Performance ?

# of examples ?

 $f = x_2 \wedge x_3 \wedge x_4 \wedge x_5 \wedge x_{100}$ 

- With the given data, we only learned an "approximation" to the true concept
- We don't know how many examples we need to see to learn **exactly**. (do we care?)
- But we know that we can make a limited # of mistakes.



#### **Two Directions**

- Can continue to analyze the probabilistic intuition:
  - Never saw x<sub>1</sub>=0 in positive examples, maybe we'll never see it?
  - And if we will, it will be with small probability, so the concepts we learn may be pretty good
  - Good: in terms of performance on future data
  - PAC framework
- Mistake Driven Learning algorithms/On line algorithms
  - Now, we can only reason about #(mistakes), not #(examples)
    - any relations?
  - Update your hypothesis only when you make mistakes
    - Not all on-line algorithms are mistake driven, so performance measure could be different.

#### **On-Line Learning**

- New learning algorithms

   (all learn a linear function over the feature space)
  - Perceptron (+ many variations)
  - General Gradient Descent view
- Issues:
  - Importance of Representation
  - Complexity of Learning
  - Idea of Kernel Based Methods
  - More about features

#### Generic Mistake Bound Algorithms

- Is it clear that we can bound the number of mistakes ?
- Let C be a finite concept class. Learn f 2 C
- CON:
  - In the ith stage of the algorithm:
  - C<sub>i</sub> all concepts in C consistent with all i-1 previously seen examples
  - Choose randomly f 2 C<sub>i</sub> and use to predict the next example
  - Clearly, C<sub>i+1</sub> μ C<sub>i</sub> and, if a mistake is made on the ith example, then |C<sub>i+1</sub>| < |C<sub>i</sub>| so progress is made.
- The CON algorithm makes at most |C|-1 mistakes
- Can we do better ?

# The Halving Algorithm

- Let C be a concept class. Learn f 2 C
- Algorithm:
- In the ith stage of the algorithm:
  - C<sub>i</sub> all concepts in C consistent with all i-1 previously seen examples
- Given an example e<sub>t</sub> consider the value f<sub>j</sub> (e<sub>t</sub>) for all f<sub>j</sub> 2 C<sub>i</sub> and predict by majority.
- Predict 1 iff

 $|\{f_j \in C_i; f_j (e_i) = 0\}| < |\{f_j \in C_i; f_j (e_i) = 1\}|$ 

- Clearly  $C_{i+1} \subseteq C_i$  and if a mistake is made in the ith example, then  $|C_{i+1}| < 1/2 |C_i|$
- The Halving algorithm makes at most log(|C|) mistakes
  - Of course, this is a theoretical algorithm; can this ne achieved with an efficient algorithm?

#### Administration

#### **Questions?**

- Hw1 is done ③
  - Recall that this is an Applied Machine Learning class.
  - We are not asking you to simply give us back what you've seen in class.
  - The HW will try to simulate challenges you might face when you want to apply ML.
  - Allow you to experience various ML scenarios and make observations that are best experienced when you play with it yourself.

#### Hw2 will be out tomorrow

- Please start to work on it early.
- This way, you will have a chance to ask questions in time.
- Come to the recitations and to office hours.
- Be organized you will run a lot of experiments, but a good script can do a lot of the work.
- Recitations

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# Learning Conjunctions

There is a hidden conjunctions the learner is to learn

 $f(x_1, x_2, ..., x_{100}) = x_2 \wedge x_3 \wedge x_4 \wedge x_5 \wedge x_{100}$ 

The number of (all; not monotone) conjunctio

Can mistakes be bounded in the **nonfinite case?** 

Can this bound be

achieved?

Iog(|C|) = n

Last time:

- Talked about various learning protocols & on algorithms for conjunctions.
- Discussed the performance of the algorithms in terms of bounding the number of mistakes that algorithm makes.
- Gave a "theoretical" algorithm with log |C| mistakes.
  - The number of k-conjunctions:  $\binom{n}{k} 2^k$ 
    - log(|C|) = klog n
    - Can we learn efficiently with this number of mistakes ?

#### Representation

- Assume that you want to learn conjunctions. Should your hypothesis space be the class of conjunctions?
  - **Theorem:** Given a sample on n attributes that is consistent with a conjunctive concept, it is NP-hard to find a pure conjunctive hypothesis that is both consistent with the sample and has the minimum number of attributes.
  - [David Haussler, AIJ'88: "Quantifying Inductive Bias: AI Learning Algorithms and Valiant's Learning Framework"]
- Same holds for Disjunctions.
- Intuition: Reduction to minimum set cover problem.
  - Given a collection of sets that cover X, define a set of examples so that learning the best (dis/conj)junction implies a minimal cover.
- Consequently, we cannot learn the concept efficiently as a (dis/con)junction.
- But, we will see that we can do that, if we are willing to learn the concept as a Linear Threshold function.
- In a more expressive class, the search for a good hypothesis sometimes becomes combinatorially easier.
   So, there

So, there is a tradeoff! (recall your DT results)

#### Linear Threshold Functions

$$f(\mathbf{x}) = \operatorname{sgn} \{ w^T \cdot x - \theta \} = \operatorname{sgn} \{ \sum_{i=1}^n w_i x_i - \theta \}$$

- Many functions are Linear
  - Conjunctions:
    - $y = x_1 \wedge x_3 \wedge x_5$
    - $y = sgn\{1 \cdot x_1 + 1 \cdot x_3 + 1 \cdot x_5 3\};$   $w = (1, 0, 1, 0, 1) \theta = 3$

Probabilistic Classifiers as well

- At least m of n:
  - y = at least 2 of {x<sub>1</sub>, x<sub>3</sub>, x<sub>5</sub>}
  - $y = sgn\{1 \cdot x_1 + 1 \cdot x_3 + 1 \cdot x_5 2\}\};$   $w = (1, 0, 1, 0, 1) \theta = 2$
- Many functions are not
  - Xor:  $y = (x_1 \land x_2) \lor (\neg x_1 \land \neg x_2)$
  - Non trivial DNF:  $y = (x_1 \land x_2) \lor (x_3 \land x_4)$
- But can be made linear
- Note: all the variables above are Boolean variables



#### **Canonical Representation**

$$f(\mathbf{x}) = \operatorname{sgn} \{ w^T \cdot x \cdot \theta \} = \operatorname{sgn} \{ \sum_{i=1}^n w_i x_i - \theta \}$$

- Note: sgn { $w^T \cdot x \theta$ } = sgn { $w'^T \cdot x'$ }
- Where:
  - x' = (x, -1) and w' = (w, θ)
- Moved from an n dimensional representation to an (n+1) dimensional representation, but now can look for hyperplanes that go through the origin.
- Basically, that means that we learn both w and  $\theta$


# Perceptron learning rule

- On-line, mistake driven algorithm.
- Rosenblatt (1959) suggested that when a target output value is provided for a single neuron with fixed input, it can incrementally change weights and learn to produce the output using the <u>Perceptron learning rule</u>
- Perceptron == Linear Threshold Unit)



# Perceptron learning rule

- We learn f:X $\rightarrow$ {-1,+1} represented as f =sgn{w<sup>T</sup>•x}
- Where X=  $\{0,1\}^n$  or X=  $\mathbb{R}^n$  and  $w \in \mathbb{R}^n$
- Given Labeled examples:  $\{(x_1, y_1), (x_2, y_2), ..., (x_m, y_m)\}$ 
  - 1. Initialize w=0 $\in \mathbb{R}^n$
  - 2. Cycle through all examples [multiple times]
    - a. Predict the label of instance x to be  $y' = sgn\{w^T \bullet x\}$
    - b. If  $y' \neq y$ , update the weight vector:

**w** = **w** + **r y x** (r - a constant, learning rate)

Otherwise, if y'=y, leave weights unchanged.

## Perceptron in action



PositiveNegative

(Figures from Bishop 2006)



(Figures from Bishop 2006)

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Positive Negative

# Perceptron learning rule

- If x is Boolean, only weights of active features are updated
- Why is this important?



•  $w^T x > 0$  is equivalent to:

$$\frac{1}{1+e^{-w^T x}} > \frac{1}{2}$$

# **Perceptron Learnability**

- Obviously can't learn what it can't represent (???)
  - Only linearly separable functions
- Minsky and Papert (1969) wrote an influential book demonstrating Perceptron's representational limitations
  - Parity functions can't be learned (XOR)
  - In vision, if patterns are represented with local features, can't represent symmetry, connectivity
- Research on Neural Networks stopped for years
- Perceptron
  - Rosenblatt himself (1959) asked,
    - *"What pattern recognition problems can be transformed so as to become linearly separable?"*



## Perceptron Convergence

- Perceptron Convergence Theorem:
- If there exist a set of weights that are consistent with the data (i.e., the data is linearly separable), the perceptron learning algorithm will converge
  - How long would it take to converge ?
- Perceptron Cycling Theorem:
- If the training data is not linearly separable the perceptron learning algorithm will eventually repeat the same set of weights and therefore enter an infinite loop.
  - How to provide robustness, more expressivity ?

## Perceptron

Input set of examples and their labels  $\mathcal{Z} = ((x_1, y_1), \dots, (x_m, y_m)) \in (\mathbb{R}^n \times \{-1, 1\})^m$ ,  $\eta$ ,  $\theta_{Init}$ 

- Initialize  $\mathbf{w} \leftarrow \mathbf{0}$  and  $\mathbf{ heta} \leftarrow \mathbf{ heta}_{Init}$
- for every training epoch:
- for every  $x_j \in \mathcal{X}$ :

$$- \hat{y} \leftarrow sign(\langle \mathbf{w}, x_j \rangle - \theta)$$
  
- if  $(\hat{y} \neq y_j)$   
\*  $\mathbf{w} \leftarrow \mathbf{w} + \eta y_j x_j$   
\*  $\theta \leftarrow \theta + \eta y_j \theta_{Init}$ 

Just to make sure we understand that we learn both w and  $\mu$ 

### Perceptron: Mistake Bound Theorem

- Maintains a weight vector  $w \in \mathbb{R}^N$ ,  $w_0 = (0, ..., 0)$ .
- Upon receiving an example  $X \in \mathbb{R}^{N}$
- Predicts according to the linear threshold function  $w^T \bullet x \ge 0$ .
- Theorem [Novikoff,1963] Let  $(x_1; y_1), ..., : (x_t; y_t)$ , be a sequence of labeled examples with  $x_i \in <^N$ ,  $||x_i|| \le R$  and  $y_i \in \{-1,1\}$  for all i. Let  $u \in <^N$ ,  $\gamma > 0$  be such that, ||u|| = 1 and

 $y_i u^T \bullet x_i \geq \gamma$  for all i.

**Complexity Parameter** 

Then Perceptron makes at most  $R^2 / \gamma^2$  mistakes on this example sequence.

(see additional notes)

# Perceptron-Mistake Bound

**Proof:** Let  $v_k$  be the hypothesis before the *k*-th mistake. Assume that the *k*-th mistake occurs on the input example  $(x_{i'}, y_i)$ .

$$\therefore y_i(\vec{v_k} \cdot \vec{x_i}) \le 0$$

Assumptions  $v_1 = \mathbf{0}$ 

 $y_i u^T \bullet \mathbf{x}_i \geq \gamma$ 

*||u||* = 1

- $\begin{aligned} v_{\vec{k}+1} &= \vec{v_k} + y_i \vec{x_i} \\ v_{\vec{k}+1} \cdot \vec{u} &= \vec{v_k} \cdot \vec{u} + y_i (\vec{u} \cdot \vec{x_i}) \\ &\geq \vec{v_k} \cdot \vec{u} + \gamma \\ \therefore \quad v_{\vec{k}+1} \cdot \vec{u} &\geq k\gamma \end{aligned}$
- 1. Note that the bound does not depend on the dimensionality nor on the number of examples.
- 2. Note that we place weight vectors and examples in the same space.
- 3. Interpretation of the theorem

$$\begin{aligned} \|v_{\vec{k}+1}\|^2 &= \|\vec{v}_{\vec{k}}\|^2 + 2y_i(\vec{v}_{\vec{k}} \cdot \vec{x}_i) + \|\vec{x}_i\|^2 \\ &\leq \|\vec{v}_{\vec{k}}\|^2 + R^2 \\ \cdot \|\vec{v}_{\vec{k}+1}\|^2 &\leq kR^2 \end{aligned}$$

Therefore,

The second inequality follows because  $\|\vec{u}\| \le 1$ . CIS419/519 Spring '18

# **Robustness to Noise**

 In the case of non-separable data , the extent to which a data point fails to have margin Υ via the hyperplane w can be quantified by a slack variable

$$\xi_i = \max(0, \Upsilon - y_i w^T x_i).$$

- Observe that when ξ<sub>i</sub> = 0, the example x<sub>i</sub> has margin at least Υ.
   Otherwise, it grows linearly with y<sub>i</sub> w<sup>T</sup> x<sub>i</sub>
- Denote:  $D_2 = [\sum {\xi_i^2}]^{1/2}$
- Theorem: The perceptron is guaranteed to make no more than ((R+D<sub>2</sub>)/Y)<sup>2</sup> mistakes on any sequence of examples satisfying ||x<sub>i</sub>||<sup>2</sup><R</li>
- Perceptron is expected to have some robustness to noise.

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itness to noise.

## Perceptron for Boolean Functions

- How many mistakes will the Perceptron algorithms make when learning a k-disjunction?
- Try to figure out the bound
- Find a sequence of examples that will cause Perceptron to make O(n) mistakes on k-disjunction on n attributes.
- Where is n coming from?)
- Recall that halving suggested the possibility of a better bound – klog(n).
- This can be achieved by Winnow
  - A multiplicative update algorithm [Littlestone'88]
  - See HW2

# **Practical Issues and Extensions**

- There are many extensions that can be made to these basic algorithms.
- Some are necessary for them to perform well
  - Regularization (next; will be motivated in the next section, COLT)
- Some are for ease of use and tuning
  - Converting the output of a Perceptron/Winnow to a conditional probability

$$P(y = +1|x) = \frac{1}{1 + e^{-Aw^T x}}$$

- The parameter A can be tuned on a development set
- Multiclass classification (later)
- Key efficiency issue: Infinite attribute domain
  - Sparse representation on the input

### **Regularization Via Averaged Perceptron**

- An Averaged Perceptron Algorithm is motivated by the following considerations:
  - In real life, we want more guarantees from our learning algorithm
  - In the mistake bound model:
    - We don't know when we will make the mistakes.
  - Every Mistake-Bound Algorithm can be converted efficiently to a PAC algorithm to yield global guarantees on performance.
  - In the PAC model:
    - Dependence is on number of examples seen and not number of mistakes.
    - Being consistent with more examples is better
    - Which hypothesis will you choose...??
- To convert a given Mistake Bound algorithm (into a global guarantee algorithm):
  - Wait for a long stretch w/o mistakes (there must be one)
  - Use the hypothesis at the end of this stretch.
  - Its PAC behavior is relative to the length of the stretch.
- Averaged Perceptron returns a weighted average of a number of earlier hypotheses; the weights are a function of the length of no-mistakes stretch.

### **Regularization Via Averaged Perceptron**

**Training:** 

[m: #(examples); k: #(mistakes) = #(hypotheses);  $c_i$ : consistency count for  $v_i$ ]

- Input: a labeled training set {(x<sub>1</sub>, y<sub>1</sub>),...(x<sub>m</sub>, y<sub>m</sub>)}
- Number of epochs T
- Output: a list of weighted perceptrons {(v<sub>1</sub>, c<sub>1</sub>),...,(v<sub>k</sub>, c<sub>k</sub>)}
- Initialize: k=0;  $v_1 = 0$ ,  $c_1 = 0$
- Repeat T times:
  - For i =1,...m:
  - Compute prediction y' = sgn( $v_k^T \mathbf{x}_i$ )
  - If y' = y, then  $c_k = c_k + 1$

else:  $v_{k+1} = v_k + y_i x$ ;  $c_{k+1} = 1$ ; k = k+1

**Prediction:** 

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Given: a list of weighted perceptrons  $\{(v_1, c_1), \dots, (v_k, c_k)\}$ ; a new example x Predict the label(x) as follows:

 $y(x) = sgn \left[ \sum_{i,k} c_i (v_i^T x) \right]$ 

Averaged version of Perceptron /Winnow is as good as any other linear learning algorithm, if not better.

- This can be done on top of any online mistake driven algorithm.
- three different algorithms.

• In HW two you will run it over

# Perceptron with Margin

- Thick Separator (aka as Perceptron with Margin) (Applies both for Perceptron and Winnow)
- Promote if:
  - $\mathbf{w}^{\mathsf{T}} \mathbf{x} \mathbf{\theta} < \gamma$
  - Demote if:
    - $\mathbf{w}^{\mathsf{T}} \mathbf{x} \mathbf{\theta} > \gamma$



Note:  $\gamma$  is a functional margin. Its effect could disappear as w grows. Nevertheless, this has been shown to be a very effective algorithmic addition. (Grove & Roth 98,01; Karov et. al 97)

# **Other Extensions**

- Assume you made a mistake on example x.
- You then see example x again; will you make a mistake on it?
- Threshold relative updating (Aggressive Perceptron)
- $w \leftarrow w + rx$  $r = \frac{\theta - w^T x}{||x||^2}$ Equivalent to updating
- on the same example
  - multiple times
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# LBJava

- Several of these extensions (and a couple more) are implemented in the LBJava learning architecture that supports several linear update rules (Winnow, Perceptron, naïve Bayes)
- Supports
  - Regularization(averaged Winnow/Perceptron; Thick Separator)
  - Conversion to probabilities
  - Automatic parameter tuning
  - True multi-class classification
  - Feature Extraction and Pruning
  - Variable size examples
  - Good support for large scale domains in terms of number of examples and number of features.
  - Very efficient
  - Many other options
- [Download from: http://cogcomp.org/page/software/]

### General Stochastic Gradient Algorithms

The loss Q: a function

of x, w and y

Given examples {z=(x,y)}<sub>1, m</sub> from a distribution over X×Y, we are trying to learn a linear function, parameterized by a weight vector w, so that we minimize the expected risk function

$$J(w) = E_z Q(z,w) \sim = \sim 1/m \sum_{i,m} Q(z_i, w_i)$$

In Stochastic Gradient Descent Algorithms we approximate this minimization by incrementally updating the weight vector w as follows:

$$w_{t+1} = w_t - r_t g_w Q(z_t, w_t) = w_t - r_t g_t$$

- Where  $g_t = g_w Q(z_t, w_t)$  is the gradient with respect to w at time t.
- The difference between algorithms now amounts to choosing a different loss function Q(z, w)



$$w_{t+1} = w_t - r_t g_w Q(x_t, y_t, w_t) = w_t - r_t g_t$$

- LMS:  $Q((x, y), w) = 1/2 (y w^T x)^2$
- leads to the update rule (Also called Widrow's Adaline):

$$w_{t+1} = w_t + r (y_t - w_t^T x_t) x_t$$

- Here, even though we make binary predictions based onsgn (w<sup>T</sup> x) we do not take the sign of the dot-product into account in the loss.
- Another common loss function is:
- Hinge loss:

 $Q((x, y), w) = max(0, 1 - y w^T x)$ 

This leads to the perceptron update rule:



If 
$$y_i w_i^T \cdot x_i > 1$$
 (No mistake, by a margin): No update  
Otherwise (Mistake, relative to margin):  $w_{t+1} = w_t + r y_t x_t$   
Here  $g = -yx$   
Good to think about the  
case of Boolean examples 58

### New Stochastic Gradient Algorithms

 $w_{t+1} = w_t - r_t g_w Q(z_t, w_t) = w_t - r_t g_t$ 

(notice that this is a vector, each coordinate (feature) has its own  $w_{t,i}$  and  $g_{t,i}$ )

- So far, we used fixed learning ratesr = r<sub>t</sub>, but this can change.
- AdaGrad alters the update to adapt based on historical information
  - Frequently occurring features in the gradients get small learning rates and infrequent features get higher ones.
  - The idea is to "learn slowly" from frequent features but "pay attention" to rare but informative features.
- Define a "per feature" learning rate for the feature j, as:

$$r_{t,j} = r/(G_{t,j})^{1/2}$$

- where  $G_{t,j} = \sum_{k=1, t} g_{k,j}^2$  the sum of squares of gradients at feature j until time t.
- Overall, the update rule for Adagrad is:

$$w_{t+1,j} = w_{t,j} - g_{t,j} r/(G_{t,j})^{1/2}$$

Easy to think about the case of Perceptron, and on Boolean examples.

This algorithm is supposed to update weights faster than Perceptron or LMS when needed.

# Regularization

 The more general formalism adds a regularization term to the risk function, and minimize:

 $J(w) = \sum_{i, m} Q(z_i, w_i) + \lambda R_i(w_i)$ 



- Where R is used to enforce "simplicity" of the learned functions.
- LMS case: Q((x, y), w) =(y w<sup>T</sup> x)<sup>2</sup>
  - $R(w) = ||w||_2^2$  gives the optimization problem called **Ridge Regression**.
  - $R(w) = ||w||_1$  gives a problem called the LASSO problem
- Hinge Loss case:  $Q((x, y), w) = max(0, 1 y w^T x)$ 
  - $R(w) = ||w||_{2}^{2}$  gives the problem called **Support Vector Machines**
- Logistics Loss case: Q((x,y),w) = log (1+exp{-y w<sup>T</sup> x})
  - $R(w) = ||w||_2^2$  gives the problem called **Logistics Regression**
- These are convex optimization problems and, in principle, the same gradient descent mechanism can be used in all cases.
- We will see later why it makes sense to use the "size" of w as a way to control "simplicity".

# **Algorithmic Approaches**

- Focus: Two families of algorithms (one of the on-line representative)
  - Additive update algorithms: Perceptron
    - SVM is a close relative of Perceptron
  - Multiplicative update algorithms: Winnow
    - Close relatives: Boosting, Max entropy/Logistic Regression

### Which algorithm is better? How to Compare?

### Generalization

- Since we deal with linear learning algorithms, we know (???) that they will all converge eventually to a perfect representation.
  - All can represent the data
- So, how do we compare:
  - 1. How many examples are needed to get to a given level of accuracy?
  - 2. Efficiency: How long does it take to learn a hypothesis and evaluate it (per-example)?
  - 3. Robustness (to noise);
  - 4. Adaptation to a new domain, ....
- With (1) being the most fundamental question:
  - Compare as a function of what?
    - One key issue is the characteristics of the data

## Sentence Representation

S= I don't know whether to laugh or cry

- Define a set of features:
  - features are relations that hold in the sentence
- Map a sentence to its feature-based representation
  - The feature-based representation will give some of the information in the sentence
- Use this feature-based representation as an example to your algorithm

## Sentence Representation

S= I don't know whether to laugh or cry

- Define a set of features:
  - features are properties that hold in the sentence
- Conceptually, there are two steps in coming up with a feature-based representation
  - What are the information sources available?
    - Sensors: words, order of words, properties (?) of words
  - What features to construct based on these?

Why is this distinction needed?



# **Domain Characteristics**

- The number of potential features is very large
- The instance space is sparse
- Decisions depend on a small set of features: the function space is sparse
- Want to learn from a number of examples that is small relative to the dimensionality

# Generalization

- Dominated by the sparseness of the function space
  - Most features are irrelevant
- # of examples required by multiplicative algorithms depends mostly on # of relevant features
  - (Generalization bounds depend on the target ||u||)
- # of examples required by additive algorithms depends heavily on sparseness of features space:
  - Advantage to additive. Generalization depend on input ||x||
    - (Kivinen/Warmuth 95).
- Nevertheless, today most people use additive algorithms.

# Which Algorithm to Choose?

#### Generalization

The  $I_1$  norm:  $||x||_1 = \sum_i |x_i|$ The  $I_2$  norm:  $||x||_2 = (\sum_1^n |x_i|^2)^{1/2}$ The  $I_p$  norm:  $||x||_p = (\sum_1^n |x_i|^P)^{1/p}$ The  $I_1$  norm:  $||x||_1 = \max_i |x_i|$ 

- Multiplicative algorithms:
  - Bounds depend on ||u||, the separating hyperplane; i: example #)
  - $M_w = 2\ln n ||u||_1^2 \max_i ||x^{(i)}||_1^2 / \min_i (u |x^{(i)}|)^2$
  - Do not care much about data; advantage with sparse target u
- Additive algorithms:
  - Bounds depend on ||x|| (Kivinen / Warmuth, '95)
  - $M_p = ||u||_2^2 \max_i ||x^{(i)}||_2^2 / \min_i (u x^{(i)})^2$
  - Advantage with few active features per example

### Examples

$$\begin{split} \mathsf{M}_{w} =& 2\ln n \| \|u\|_{1}^{2} \max_{i} \| x^{(i)} \|_{1}^{2} / \min_{i} (u \ x^{(i)})^{2} \\ \mathsf{M}_{p} =& \| \|u\|_{2}^{2} \max_{i} \| \|x^{(i)} \|_{2}^{2} / \min_{i} (u \ x^{(i)})^{2} \end{split}$$

- Extreme Scenario 1: Assume the u has exactly k active features, and the other n-k are 0. That is, only k input features are relevant to the prediction. Then:
  - $||\mathbf{u}||_2, = k^{1/2} ; ||\mathbf{u}||_1, = k ; \max ||\mathbf{x}||_2, = n^{1/2} ; \max ||\mathbf{x}||_1, = 1$
  - We get that:  $M_p = kn;$   $M_w = 2k^2 \ln n$
  - Therefore, if k<<n, Winnow behaves much better.
- Extreme Scenario 2: Now assume that u=(1, 1,....1) and the instances are very sparse, the rows of an n×n unit matrix. Then:
  - $||\mathbf{u}||_2, = \mathsf{n}^{1/2} ; ||\mathbf{u}||_1, = \mathsf{n} ; \max ||\mathbf{x}||_2, = \mathsf{1} ; \max ||\mathbf{x}||_1 , = \mathsf{1}$
  - We get that:  $M_p = n$ ;  $M_w = 2n^2 \ln n$
  - Therefore, Perceptron has a better bound.



# Summary



- Introduced multiple versions of on-line algorithms
- All turned out to be Stochastic Gradient Algorithms
  - For different loss functions
- Some turned out to be mistake driven

A term that minimizes error on the training data

- We suggested generic improvements via:
  - Regularization via adding a term that forces a "simple hypothesis"

 $J(w) = \sum_{i, m} Q(z_i, w_i) + \lambda R_i(w_i)$ 

- Regularization via the Averaged Trick
  - "Stability" of a hypothesis is related to its ability to generalize
- An improved, adaptive, learning rate (Adagrad)
- Dependence on function space and the instance space properties.
- Today:
  - A way to deal with non-linear target functions (Kernels)
  - Beginning of Learning Theory.

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A term that forces simple hypothesis

# Efficiency

- Dominated by the size of the feature space
- Most features are functions (e.g. conjunctions) of raw attributes

 $X(x_1, x_2, x_3, \dots, x_k) \to X(\chi_1(\mathbf{x}), \chi_2(\mathbf{x}), \chi_3(\mathbf{x}) \dots \chi_n(\mathbf{x})) \qquad \mathbf{n} >> \mathbf{k}$ 

- Additive algorithms allow the use of Kernels
  - No need to explicitly generate complex features

$$f(x) = \sum_{i} c_i K(x, x_i)$$

 Could be more efficient since work is done in the original feature space, but expressivity is a function of the kernel expressivity.

# Functions Can be Made Linear

- Data are not linearly separable in one dimension
- Not separable if you insist on using a specific class of functions


### **Blown Up Feature Space**

Data are separable in <x, x<sup>2</sup>> space



### Making data linearly separable

Original feature space



## Making data linearly separable



\_ \_ \_

f(x') = 1 iff  $x'_1 + x'_2 \le 1$ 

### **Dual Representation**

Examples  $x \in \{0,1\}^N$ ; Learned hypothesis  $w \in \mathbb{R}^N$  $f(x) = \text{sgn} \{w^T \cdot x\} = \text{sgn}\{\sum_{i=1}^n w_i x_i\}$ 

Perceptron Update:

#### If y'≠y, update: **w** = **w** + **ry x**

- Let w be an initial weight vector for perceptron. Let (x<sup>1</sup>,+), (x<sup>2</sup>,+), (x<sup>3</sup>,-), (x<sup>4</sup>,-) be examples and assume mistakes are made on x<sup>1</sup>, x<sup>2</sup> and x<sup>4</sup>.
- What is the resulting weight vector?

$$w = w + x^1 + x^2 - x^4$$

In general, the weight vector w can be written as a linear combination of examples:

w = 
$$\sum_{i,m} r \alpha_i y_i x^i$$

• Where  $\alpha_i$  is the number of mistakes made on  $x^i$ .

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Note: We care about the dot product:  $f(x) = w^T x =$ =  $(\sum_{1,m} r \alpha_i y_i x^i)^T x$ =  $\sum_{1,m} r \alpha_i y_i (x^{iT} x)$ 

### Kernel Based Methods

 $f(\mathbf{x}) = \operatorname{sgn} \{ w^T \cdot x \} = \operatorname{sgn} \{ \sum_{i=1}^n w_i x_i \}$ 

- A method to run Perceptron on a very large feature set, without incurring the cost of keeping a very large weight vector.
- Computing the dot product can be done in the original feature space.
- Notice: this pertains only to efficiency: The classifier is identical to the one you get by blowing up the feature space.
- Generalization is still relative to the real dimensionality (or, related properties).
- Kernels were popularized by SVMs, but many other algorithms can make use of them (== run in the dual).
  - Linear Kernels: no kernels; stay in the original space. A lot of applications actually use linear kernels.

### Kernel Base Methods

Examples  $x \in \{0,1\}^N$  ; Learned hypothesis  $w \in \, {\mathcal R}^{\! N}$ 

$$f(\mathbf{x}) = \operatorname{sgn} \{ w^T \cdot x \} = \operatorname{sgn} \{ \sum_{i=1}^n w_i x_i (\mathbf{x}) \}$$

- Let I be the set t<sub>1</sub>,t<sub>2</sub>,t<sub>3</sub> ... of monomials (conjunctions) over the feature space x<sub>1</sub>, x<sub>2</sub>... x<sub>n</sub>.
- Then we can write a linear function over this new feature space.

$$f(\mathbf{x}) = \operatorname{sgn} \{ w^T \cdot x \} = \operatorname{sgn} \{ \sum_I w_i t_i(\mathbf{x}) \}$$

Example:  $x_1x_2x_4(11010) = 1$   $x_3x_4(11010) = 0$   $x_1x_2x_4(11011) = 1$ 

### Kernel Based Methods

Examples  $x \in \{0,1\}^N$ ; Learned hypothesis  $w \in \mathbb{R}^N$  $f(x) = sgn \{w^T \cdot x\} = sgn\{\sum_I w_i t_i(x)\}$ 

Perceptron Update:

If y'≠y, update: **w** = **w** + **ry x** 

- Great Increase in expressivity
- Can run Perceptron (and Winnow) but the convergence bound may suffer exponential growth.
- Exponential number of monomials are true in each example.
- Also, will have to keep many weights.

### Embedding



### The Kernel Trick(1)

Examples  $x \in \{0,1\}^N$ ; Learned hypothesis  $w \in \mathbb{R}^N$  $f(x) = sgn \{w^T \cdot x\} = sgn\{\sum_I w_i t_i(x)\}$ 

Perceptron Update:

If y'≠y, update: **w** = **w** + **ry x** 

- Consider the value of w used in the prediction.
- Each previous mistake, on example z, makes an additive contribution of +/-1 to some of the coordinates of w.
  - Note: examples are Boolean, so only coordinates of w that correspond to ON terms in the example z (t<sub>i</sub>(z) = 1) are being updated.
- The value of w is determined by the number and type of mistakes.

## The Kernel Trick(2)

Examples  $x \in \{0,1\}^N$ ; Learned hypothesis  $w \in \mathbb{R}^N$  $f(x) = \text{sgn} \{w^T \cdot x\} = \text{sgn}\{\sum_I w_i t_i(x)\}$ 

Perceptron Update:

If y'≠y, update: **w** = **w** + **ry x** 

- P set of examples on which we Promoted
- D set of examples on which we Demoted
  M = P [ D

 $f(x) = \operatorname{sgn}_{\Sigma_{I}} w_{i} t_{i}(x) = \sum_{I} \left[ \sum_{z \in P, t_{i}(z)=1} 1 - \sum_{z \in D, t_{i}(z)=1} 1 \right] t_{i}(x) =$ 

$$= \sum_{I} \left[ \sum_{z \in M} S(z) t_i(z) t_i(x) \right]$$

### The Kernel Trick(3)

$$f(\mathbf{x}) = \operatorname{sgn} \{ w^T \cdot x \} = \operatorname{sgn} \{ \sum_I w_i t_i(\mathbf{x}) \}$$

- P set of examples on which we Promoted
- D set of examples on which we Demoted
- M = P [ D

$$f(\mathbf{x}) = \operatorname{sgn}\sum_{I} w_{i}t_{i}(\mathbf{x}) = \sum_{I} \left[\sum_{z \in P, t_{i}(z)=1} 1 - \sum_{z \in D, t_{i}(z)=1} 1\right]t_{i}(\mathbf{x})$$
$$= \operatorname{sgn}\left\{\sum_{I} \left[\sum_{z \in M} S(z)t_{i}(z)t_{i}(x)\right]\right\}$$

- Where S(z)=1 if  $z \in P$  and S(z) = -1 if  $z \in D$ .
- Reordering:

$$f(\mathbf{x}) = \operatorname{sgn}\{\sum_{z \in M} S(z) \sum_{I} S(z) t_i(z) t_i(x)\}$$

## The Kernel Trick(4) $f(x) = Th_{\theta}(\sum_{i \in I} w_i t_i(x))$

- $S(y)=1 \text{ if } y \in P \text{ and } S(y) = -1 \text{ if } y \in D.$  $f(x) = Th_{\theta} \left( \sum_{z \in M} S(z) \sum_{i \in I} t_i(z) t_i(x) \right)$
- A mistake on z contributes the value +/-1 to all monomials satisfied by z. The total contribution of z to the sum is equal to the number of monomials that satisfy both x and z.
- Define a dot product in the t-space:

$$\textbf{K(x,z)} = \sum_{i \in I} \textbf{t}_i(\textbf{z})\textbf{t}_i(\textbf{x})$$

• We get the standard notation:

$$f(x) = \mathsf{Th}_{\theta}(\sum_{z \in M} \mathsf{S}(z)\mathsf{K}(x,z))$$

$$\begin{aligned} \textbf{Kernel Based Methods} \\ \textbf{f}(\textbf{x}) = \textbf{Th}_{\theta}(\sum_{\textbf{z} \in M} \textbf{S}(\textbf{z})\textbf{K}(\textbf{x},\textbf{z})) \end{aligned}$$

What does this representation give us?

$$\mathbf{K}(\mathbf{x},\mathbf{z}) = \sum_{i \in I} \mathbf{t}_i(\mathbf{z}) \mathbf{t}_i(\mathbf{x})$$

- We can view this Kernel as the distance between x,z in the t-space.
- But, K(x,z) can be measured in the original space, without explicitly writing the t-representation of x, z

 $x_{1}x_{3}(001) = x_{1}x_{3}(011) = 1$   $x_{1}(001) = x_{1}(011) = 1; \quad x_{3}(001) = x_{3}(011) = 1$  $f(x) = n_{\theta} \sum_{z \in M} S(z)K(x,z)) \quad K(x,z) = \sum_{i=1}^{n} t_{i}(z)t_{i}(x)$ 

sider the space of all 3<sup>n</sup> monomials (allowing both tive and negative literals). Then,

$$\mathbf{K}(\mathbf{x},\mathbf{z}) = \sum_{i=1}^{n} t_i(z) t_i(x) = \mathbf{2}^{\mathsf{same}(\mathbf{x},\mathbf{z})}$$

- N n same(x,z) is the n<sup>i</sup><sup>[f]</sup>mber of features that have the same value for both x and z.
- We get:

$$f(x) = \mathbf{Th}_{\theta} \left( \sum_{z \in M} \mathbf{S}(z) (2^{\mathsf{same}(x, z)}) \right)$$

Example: Take n=3; x=(001), z=(011), monomials of size 0,1,2,3

Proof: let k=same(x,z); construct a "surviving" monomials by: (1) choosing to include one of these k literals with the right polarity in the monomial, or (2) choosing to not include it at all. Monomials with literals outside this set disappear.

## Example

 $f(x) = Th_{\theta}\left(\sum_{z \in M} S(z)K(x,z)\right) K(x,z) = \sum_{i \in I} t_i(z)t_i(x)$ 

- Take  $X = \{x_1, x_2, x_3, x_4\}$
- I = The space of all 3<sup>n</sup> monomials; | I | = 81
- Consider x=(1100), z=(1101)
- Write down I(x), I(z), the representation of x, z in the I space.
- Compute I(x) ·I(z).
- Show that
- $K(x,z) = I(x) \cdot I(z) = \sum_{I} t_{i}(z) t_{i}(x) = 2^{same(x,z)} = 8$
- Try to develop another kernel, e.g., where I is the space of all conjunctions of size 3 (exactly).

### **Implementation:** Dual Perceptron

$$f(x) = Th_{\theta} \left( \sum_{z \in M} S(z)K(x,z) \right)$$
$$K(x,z) = \sum_{i \in I} t_i(z)t_i(x)$$

- Simply run Perceptron in an on-line mode, but keep track of the set M.
- Keeping the set M allows us to keep track of S(z).
- Rather than remembering the weight vector w, remember the set M (P and D) – all those examples on which we made mistakes.
- Dual Representation

### Example: Polynomial Kernel

- Prediction with respect to a separating hyper planes (produced by Perceptron, SVM) can be computed as a function of dot products of feature based representation of examples.
- We want to define a dot product in a high dimensional sp<sup>2</sup>.e.
- Given two examples x = (x<sub>1</sub>, x<sub>2</sub>, ...x<sub>n</sub>) and y = (y<sub>1</sub>, y<sub>2</sub>, ...y<sub>n</sub>) we want to map them to a high dimensional space [example- quadratic]:
  - $\Phi(\mathbf{x}_1, \mathbf{x}_2, ..., \mathbf{x}_n) = (\mathbf{1}, \mathbf{x}_1, ..., \mathbf{x}_n, \mathbf{x}_1^2, ..., \mathbf{x}_n^2, \mathbf{x}_1 \mathbf{x}_2, ..., \mathbf{x}_{n-1} \mathbf{x}_n)$
  - $\Phi(y_1, y_2, ..., y_n) = (1, y_1, ..., y_n, y_1^2, ..., y_n^2, y_1y_2, ..., y_{n-1}y_n)$

and compute the dot product  $A = \Phi(x)^T \Phi(y)$  [takes time]

Instead, in the original space, compute

• 
$$B = k(x, y) = [1 + (x_1, x_2, ..., x_n)^T (y_1, y_2, ..., y_n)]^2$$

Theorem: A = B (Coefficients do not really matter)

# Kernels – General Conditions

- Kernel Trick: You want to work with degree 2 polynomial features, Φ(x). Then, your dot product will be in a space of dimensionality n(n+1)/2. The kernel trick allows you to save and compute dot products in an n dimensional space.
- Can we use any K(.,.)?

$$f(\mathbf{x}) = \mathbf{Th}_{\theta} \left( \sum_{\mathbf{z} \in \mathbf{M}} \mathbf{S}(\mathbf{z}) \mathbf{K}(\mathbf{x}, \mathbf{z}) \right)$$

We proved that K is a valid kernel by explicitly

- A function K(x,z) is a valid kernel if it corresponds to an inner product in some (perhaps infinite dimensional) feature space.
   K(x,z) = ∑it<sub>i</sub>(z)t<sub>i</sub>(x)
- Take the quadratic kernel:  $k(x,z) = (x^Tz)^2$
- Example: Direct construction (2 dimensional, for simplicity):
- $K(x,z) = (x_1 z_1 + x_2 z_2)^2 = x_1^2 z_1^2 + 2x_1 z_1 x_2 z_2 + x_2^2 z_2^2$
- $= (x_1^2, sqrt\{2\} x_1 x_2, x_2^2) (z_1^2, sqrt\{2\} z_1 z_2, z_2^2)^{\top}$
- =  $\Phi(x)^T \Phi(z) \rightarrow A$  dot product in an expanded space.
- It is not necessary to explicitly show the feature function Φ.
- General condition: construct the kernel matrix {k(x<sub>i</sub>, z<sub>j</sub>)}; check that it's positive semi definite.

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1.5

0.5

0.5

1.5 -2

-2 -1 5 -1

-0.5

## Polynomial kernels

- Linear kernel: k(x, z) = xz
- Polynomial kernel of degree d: k(x, z) = (xz)<sup>d</sup> (only dth-order interactions)

 Polynomial kernel up to degree d: k(x, z) = (xz + c)<sup>d</sup> (c>0) (all interactions of order d or lower)

### **Constructing New Kernels**

- You can construct new kernels k'(x, x') from existing ones:
  - Multiplying k(x, x') by a constant c:
     k'(x, x') = ck(x, x')
  - Multiplying k(x, x') by a function f applied to x and x': k'(x, x') = f(x)k(x, x')f(x')
  - Applying a polynomial (with non-negative coefficients) to k(x, x'):
     k'(x, x') = P(k(x, x')) with P(z) = ∑<sub>i</sub> a<sub>i</sub>z<sup>i</sup> and a<sub>i</sub>≥0
  - Exponentiating k(x, x'):
     k'(x, x') = exp(k(x, x'))

### Constructing New Kernels (2)

- You can construct  $k'(\mathbf{x}, \mathbf{x'})$  from  $k_1(\mathbf{x}, \mathbf{x'}), k_2(\mathbf{x}, \mathbf{x'})$  by:
  - Adding  $k_1(\mathbf{x}, \mathbf{x'})$  and  $k_2(\mathbf{x}, \mathbf{x'})$ :  $k'(\mathbf{x}, \mathbf{x'}) = k_1(\mathbf{x}, \mathbf{x'}) + k_2(\mathbf{x}, \mathbf{x'})$
  - Multiplying  $k_1(\mathbf{x}, \mathbf{x'})$  and  $k_2(\mathbf{x}, \mathbf{x'})$ :  $k'(\mathbf{x}, \mathbf{x'}) = k_1(\mathbf{x}, \mathbf{x'})k_2(\mathbf{x}, \mathbf{x'})$
- Also:
  - If φ(x) ∈ R<sup>m</sup> and k<sub>m</sub>(z, z') a valid kernel in R<sup>m</sup>, k(x, x') = k<sub>m</sub>(φ(x), φ(x')) is also a valid kernel
  - If A is a symmetric positive semi-definite matrix, k(x, x') = xAx' is also a valid kernel
- In all cases, it is easy to prove these directly by construction.

### Gaussian Kernel

(aka radial basis function kernel)

- $k(x, z) = \exp(-(x z)^{2/c})$ 
  - (x z)<sup>2</sup>: squared Euclidean distance between x and z
  - $c = \sigma^2$ : a free parameter
  - very small c: K ≈ identity matrix (every item is different)
  - very large c: K ≈ unit matrix (all items are the same)
  - k(x, z) ≈ 1 when x, z close
  - k(x, z) ≈ 0 when x, z dissimilar



### **Gaussian Kernel**

- $k(x, z) = \exp(-(x z)^{2/c})$
- Is this a kernel?
- $k(\mathbf{x}, \mathbf{z}) = \exp(-(\mathbf{x} \mathbf{z})^2/2\sigma^2)$   $= \exp(-(\mathbf{x}\mathbf{x} + \mathbf{z}\mathbf{z} - 2\mathbf{x}\mathbf{z})/2\sigma^2)$   $= \exp(-\mathbf{x}\mathbf{x}/2\sigma^2) \exp(\mathbf{x}\mathbf{z}/\sigma^2)\exp(-\mathbf{z}\mathbf{z}/2\sigma^2)$  $= f(\mathbf{x}) \exp(\mathbf{x}\mathbf{z}/\sigma^2) f(\mathbf{z})$
- $exp(xz/\sigma^2)$  is a valid kernel:
- **xz** is the linear kernel;
- we can multiply kernels by constants  $(1/\sigma^2)$
- we can exponentiate kernels

Unlike the discrete kernels discussed earlier, here you cannot easily explicitly blow up the feature space to get an identical representation.

# Summary – Kernel Based Methods $f(x) = Th_{\theta}(\sum_{z \in M} S(z)K(x,z))$

- A method to run Perceptron on a very large feature set, without incurring the cost of keeping a very large weight vector.
- Computing the weight vector can be done in the original feature space.
- Notice: this pertains only to efficiency: the classifier is identical to the one you get by blowing up the feature space.
- Generalization is still relative to the real dimensionality (or, related properties).
- Kernels were popularized by SVMs but apply to a range of models, Perceptron, Gaussian Models, PCAs, etc.

### Explicit & Implicit Kernels: Complexity

- Is it always worthwhile to define kernels and work in the dual space?
- Computationally:
  - Dual space  $-t_1 m^2 vs$ , Primal Space  $-t_2 m$
  - Where m is # of examples, t<sub>1</sub>, t<sub>2</sub> are the sizes of the (Dual, Primal) feature spaces, respectively.
  - Typically, t<sub>1</sub> << t<sub>2</sub>, so it boils down to the number of examples one needs to consider relative to the growth in dimensionality.
- Rule of thumb: a lot of examples  $\rightarrow$  use Primal space
- Most applications today: People use explicit kernels. That is, they blow up the feature space explicitly.

### Kernels: Generalization

- Do we want to use the most expressive kernels we can?
  - (e.g., when you want to add quadratic terms, do you really want to add all of them?)
- No; this is equivalent to working in a larger feature space, and will lead to overfitting.
- It's possible to give simple arguments that show that simply adding irrelevant features does not help.

### **Conclusion- Kernels**

- The use of Kernels to learn in the dual space is an important idea
  - Different kernels may expand/restrict the hypothesis space in useful ways.
  - Need to know the benefits and hazards
- To justify these methods we must embed in a space much larger than the training set size.
  - Can affect generalization
- Expressive structures in the input data could give rise to specific kernels, designed to exploit these structures.
  - E.g., people have developed <u>kernels over parse trees</u>: corresponds to features that are sub-trees.
  - It is always possible to trade these with explicitly generated features, but it might help one's thinking about appropriate features.

### Functions Can be Made Linear

- Data are not linearly separable in one dimension
- Not separable if you insist on using a specific class of functions



### **Blown Up Feature Space**

Data are separable in <x, x<sup>2</sup>> space



## Multi-Layer Neural Network

- Multi-layer network were designed to overcome the computational (expressivity) limitation of a single threshold element.
- The idea is to stack several layers of threshold elements, each layer using the output of the previous layer as input.



 Multi-layer networks can represent arbitrary functions, but building effective learning methods for such network was [thought to be] difficult.

## **Basic Units**

- Linear Unit: Multiple layers of linear functions
   o<sub>j</sub> = w ¢x produce linear functions. We want to represent nonlinear functions.
- Need to do it in a way that facilitates learning
- Threshold units: o<sub>j</sub> = sgn(w ¢x) are not differentiable, hence unsuitable for gradient descent.



The key idea was to notice that the discontinuity of the threshold element can be represents by a smooth non-linear approximation: o<sub>i</sub> = [1+ exp{-w ¢x}]<sup>-1</sup>

(Rumelhart, Hinton, William, 1986), (Linnainmaa, 1970), see: <u>http://people.idsia.ch/~juergen/who-invented-backpropagation.html</u>)

# Model Neuron (Logistic)

 Us a non-linear, differentiable output function such as the sigmoid or logistic function



- Net input to a unit is defined a  $\mathbf{pet}_{j} = \sum \mathbf{W}_{ij} \cdot \mathbf{X}_{i}$
- Output of a unit is defined as:

$$\mathbf{D}_{\mathbf{j}} = \frac{1}{1 + e^{-(\mathbf{net}_{\mathbf{j}} - \mathbf{T}_{\mathbf{j}})}}$$

### Learning with a Multi-Layer Perceptron

- It's easy to learn the top layer it's just a linear unit.
- Given feedback (truth) at the top layer, and the activation at the layer below it, you can use the Perceptron update rule (more generally, gradient descent) to updated these weights.
- The problem is what to do with the other set of weights – we do not get feedback in the intermediate layer(s).



### Learning with a Multi-Layer Perceptron

- The problem is what to do with the other set of weights – we do not get feedback in the intermediate layer(s).
- Solution: If all the activation functions are differentiable, then the output of the network is also



a differentiable function of the input and weights in the network.

- Define an error function (multiple options) that is a differentiable function of the output, that this error function is also a differentiable function of the weights.
- We can then evaluate the derivatives of the error with respect to the weights, and use these derivatives to find weight values that minimize this error function. This can be done, for example, using gradient descent.
- This results in an algorithm called back-propagation.