Data Mining Techniques: Classification and Prediction

Mirek Riedewald

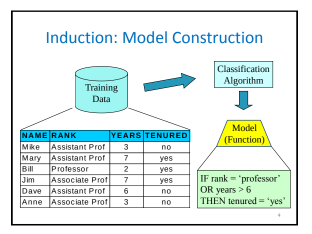
Some slides based on presentations by Han/Kamber/Pei, Tan/Steinbach/Kumar, and Andrew Moore

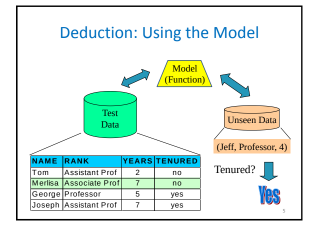
Classification and Prediction Overview

- Introduction
- Decision Trees
- Statistical Decision Theory
- Nearest Neighbor
- Bayesian Classification
- Artificial Neural Networks
- Support Vector Machines (SVMs)
- Prediction
- Accuracy and Error MeasuresEnsemble Methods

Classification vs. Prediction

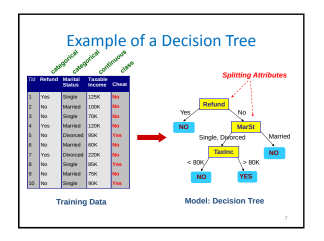
- Assumption: after data preparation, we have a data set where each record has attributes $X_1, ..., X_n$, and Y.
- Goal: learn a function $f:(X_1,...,X_n) \rightarrow Y$, then use this function to predict y for a given input record $(x_1,...,x_n)$.
 - Classification: Y is a discrete attribute, called the class label
 Usually a categorical attribute with small domain
 Prediction: Y is a continuous attribute
- Called supervised learning, because true labels (Yvalues) are known for the initially provided data
- Typical applications: credit approval, target marketing, medical diagnosis, fraud detection

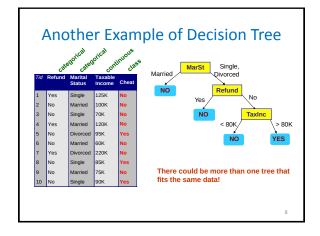


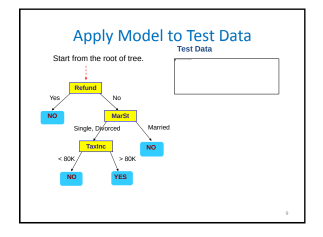


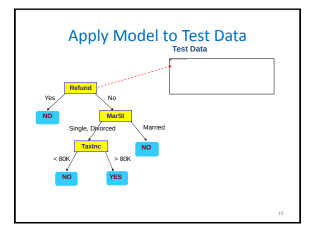
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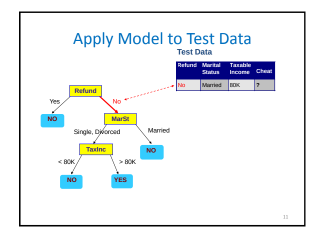
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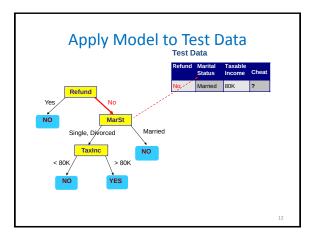


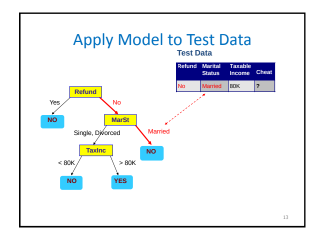


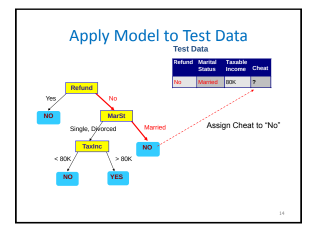


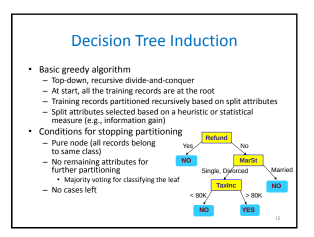


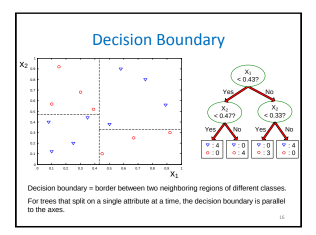


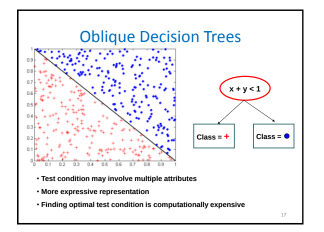


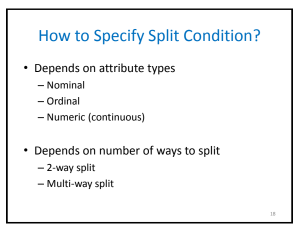


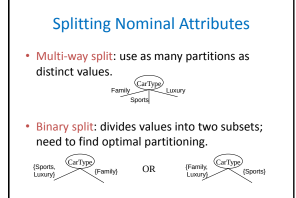


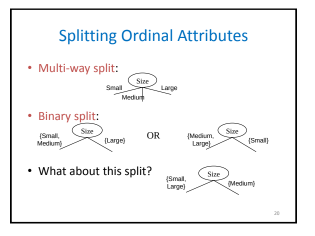








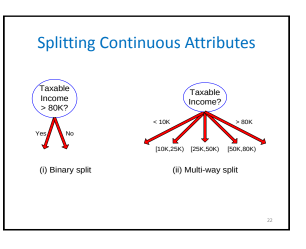


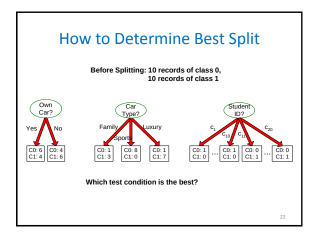


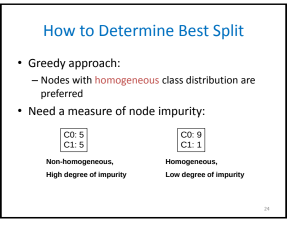
Splitting Continuous Attributes

• Different options

- Discretization to form an ordinal categorical attribute
 - Static discretize once at the beginning
 - Dynamic ranges found by equal interval bucketing, equal frequency bucketing (percentiles), or clustering.
- Binary Decision: (A < v) or $(A \ge v)$
 - Consider all possible splits, choose best one







Attribute Selection Measure: Information Gain

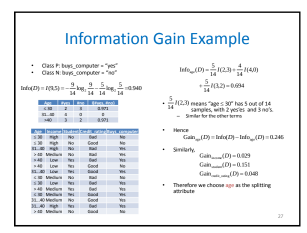
- · Select attribute with highest information gain
- \boldsymbol{p}_i = probability that an arbitrary record in D belongs to class C_{μ} i=1,...,m
- Expected information (entropy) needed to classify a record in D: $Info(D) = -\sum_{i=1}^{m} p_i \log_2(p_i)$
- Information needed after using attribute A to split D into v partitions $D_1, ..., D_v$:

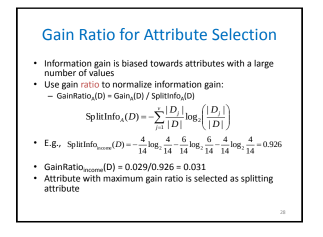
$$\operatorname{Info}_{A}(D) = \sum_{j=1}^{j} \frac{|D_{j}|}{|D|} \operatorname{Info}(D_{j})$$

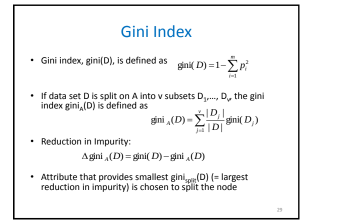
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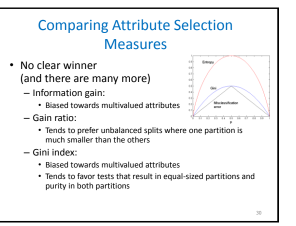
- Information gained by splitting on attribute A: ${\rm Gain}_{\scriptscriptstyle A}(D) {=} \, {\rm Info}(D) {-} \, {\rm Info}_{\scriptscriptstyle A}(D)$

Example								
Predict if somebo	ody v	vill bu	лу а с	compute	r			
 Given data set: 	Age	Income	Student	Credit_rating	Buys_computer			
• Given data set.	≤ 30	High	No	Bad	No			
	≤ 30	High	No	Good	No			
	3140	High	No	Bad	Yes			
	> 40	Medium	No	Bad	Yes			
	> 40	Low	Yes	Bad	Yes			
	> 40	Low	Yes	Good	No			
	3140	Low	Yes	Good	Yes			
	≤ 30	Medium	No	Bad	No			
	≤ 30	Low	Yes	Bad	Yes			
	> 40	Medium	Yes	Bad	Yes			
	≤ 30	Medium	Yes	Good	Yes			
	3140	Medium	No	Good	Yes			
	3140	High	Yes	Bad	Yes			
	> 40	Medium	No	Good	No			
					26			







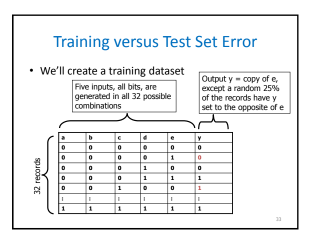


Practical Issues of Classification

- Underfitting and overfitting
- Missing values
- Computational cost
- Expressiveness

How Good is the Model?

- Training set error: compare prediction of training record with true value
 - Not a good measure for the error on unseen data. (Discussed soon.)
- Test set error: for records that were **not** used for training, compare model prediction and true value
 - Use holdout data from available data set

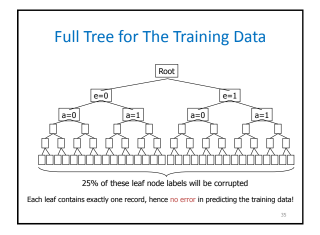


Test Data

 Generate test data using the same method: copy of e, 25% inverted; done independently from previous noise process

- Some y's that were corrupted in the training set will be uncorrupted in the testing set.
- Some y's that were uncorrupted in the training set will be corrupted in the test set.

а	b	c	d	e	y (training data)	y (test data)
0	0	0	0	0	0	0
0	0	0	0	1	0	1
0	0	0	1	0	0	1
0	0	0	1	1	1	1
0	0	1	0	0	1	1
:	:	:	:	:	:	:
1	1	1	1	1	1	1



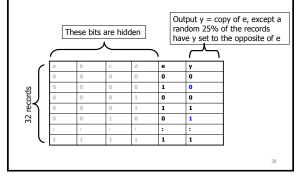
Testing The Tree with The Test Set

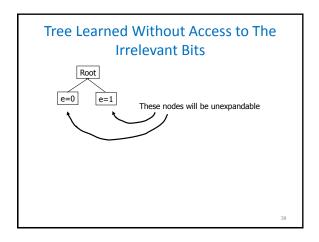
	1/4 of the tree nodes are corrupted	3/4 are fine
1/4 of the test set records are corrupted	1/16 of the test set will be correctly predicted for the wrong reasons	3/16 of the test set will be wrongly predicted because the test record is corrupted
3/4 are fine	3/16 of the test predictions will be wrong because the tree node is corrupted	9/16 of the test predictions will be fine

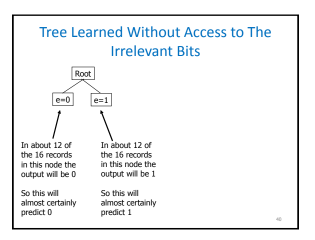
What's This Example Shown Us?

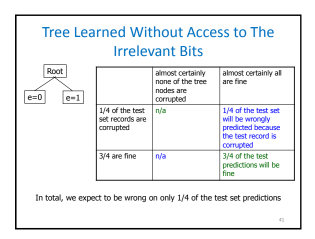
- Discrepancy between training and test set error
- But more importantly
 - ...it indicates that there is something we should do about it if we want to predict well on future data.

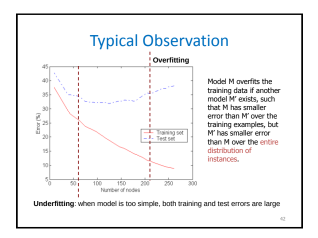
Suppose We Had Less Data











Reasons for Overfitting

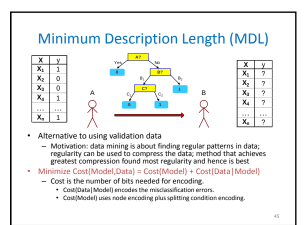
- Noise
 - Too closely fitting the training data means the model's predictions reflect the noise as well
- Insufficient training data
 - Not enough data to enable the model to generalize beyond idiosyncrasies of the training records
- Data fragmentation (special problem for trees)

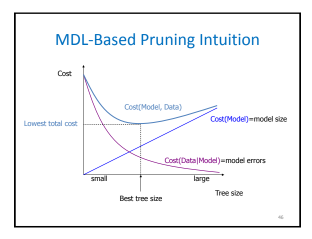
 Number of instances gets smaller as you traverse down the tree
 - Number of instances at a leaf node could be too small to make any confident decision about class

Avoiding Overfitting

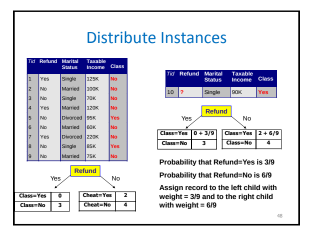
- General idea: make the tree smaller – Addresses all three reasons for overfitting
- Prepruning: Halt tree construction early

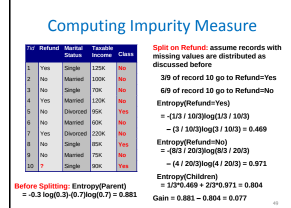
 Do not split a node if this would result in the goodness measure falling below a threshold
 - Difficult to choose an appropriate threshold, e.g., tree for XOR
- Postpruning: Remove branches from a "fully grown" tree
 Use a set of data different from the training data to decide when to stop pruning
 - Validation data: train tree on training data, prune on validation data, then test on test data

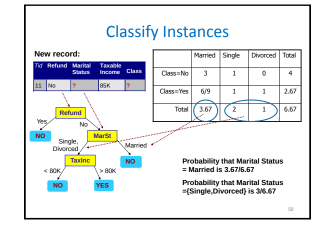




Handling Missing Attribute Values Missing values affect decision tree construction in three different ways: How impurity measures are computed How to distribute instance with missing value to child nodes How a test instance with missing value is classified







Tree Cost Analysis

- Finding an optimal decision tree is NP-complete Optimization goal: minimize expected number of binary tests to
 - uniquely identify any record from a given finite set
- Greedy algorithm
 - O(#attributes * #training_instances * log(#training_instances)) At each tree depth, all instances considered
 - Assume tree depth is logarithmic (fairly balanced splits)
 - Need to test each attribute at each node
 - What about binary splits?

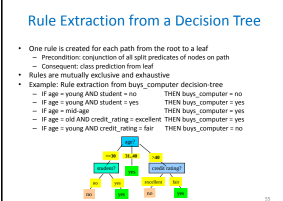
 - Sort data once on each attribute, use to avoid re-sorting subsets Incrementally maintain counts for class distribution as different split points are explored
- In practice, trees are considered to be fast both for training (when using the greedy algorithm) and making predictions



· Can represent any finite discrete-valued function

- But it might not do it very efficiently
 - Example: parity function Class = 1 if there is an even number of Boolean attributes with truth value = True
 - Class = 0 if there is an odd number of Boolean attributes with truth value = True
 - For accurate modeling, must have a complete tree
- · Not expressive enough for modeling continuous attributes

- But we can still use a tree for them in practice; it just cannot accurately represent the true function





- Scalability: Classify data sets with millions of examples and hundreds of attributes with reasonable speed
- Why use decision trees for data mining?
 - Relatively fast learning speed
 - Can handle all attribute types
 - Convertible to intelligible classification rules
 - Good classification accuracy, but not as good as newer methods (but tree ensembles are top!)

Scalable Tree Induction

- High cost when the training data at a node does not fit in memory
- Solution 1: special I/O-aware algorithm
 - Keep only class list in memory, access attribute values on disk
 - Maintain separate list for each attribute
- Use count matrix for each attribute Solution 2: Sampling

 - Common solution: train tree on a sample that fits in memory
 - More sophisticated versions of this idea exist, e.g., Rainforest Build tree on sample, but do this for many bootstrap samples
 - Combine all into a single new tree that is guaranteed to be almost identical to the one trained from entire data set
 - · Can be computed with two data scans

Tree Conclusions

- Very popular data mining tool
 - Easy to understand
 - Easy to implement
 - Easy to use: little tuning, handles all attribute types and missing values
 - Computationally relatively cheap
- Overfitting problem
- Focused on classification, but easy to extend to prediction (future lecture)

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- Statistical Decision Theory
- Nearest Neighbor
- **Bayesian Classification**
- Artificial Neural Networks
- Support Vector Machines (SVMs)
- Prediction
- Accuracy and Error Measures
- · Ensemble Methods

Theoretical Results

- Trees make sense intuitively, but can we get some hard evidence and deeper understanding about their properties?
- Statistical decision theory can give some answers
- Need some probability concepts first

Random Variables

- Intuitive version of the definition:
 - Can take on one of possibly many values, each with a certain probability
 - These probabilities define the probability distribution of the random variable
 - E.g., let X be the outcome of a coin toss, then Pr(X='heads')=0.5 and Pr(X='tails')=0.5; distribution is uniform
- Consider a discrete random variable X with numeric values x₁,...,x_k
 - Expectation: $E[X] = \sum x_i^* Pr(X=x_i)$
 - Variance: $Var(X) = E[(X E[X])^2] = E[X^2] (E[X])^2$

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Working with Random Variables • E[X + Y] = E[X] + E[Y] Var(X + Y) = Var(X) + Var(Y) + 2 Cov(X,Y) • For constants a, b - E[aX + b] = a E[X] + b- Var(aX + b) = Var(aX) = a² Var(X) Iterated expectation: - $E[X] = E_X[E_Y[Y | X]]$, where $E_Y[Y | X] = \sum y_i^* Pr(Y=y_i | X=x)$ is the expectation of Y for a given value x of X, i.e., is a function of X In general for any function f(X,Y): $E_{X,Y}[f(X,Y)] = E_X[E_Y[f(X,Y)|X]]$ 63

What is the Optimal Model f(X)?

Let X denote a real - valued random input variable and Y a real - valued random output variable

The squared error of trained model f(X) is $E_{X,Y}[(Y - f(X))^2]$

Which function f(X) will minimize the squared error?

Consider the error for a specific value of X and let $\overline{Y} = E_{y}[Y | X]$: $\mathbf{E}_{Y}\left[\left(Y-f(X)\right)^{2} \mid X\right] = \mathbf{E}_{Y}\left[\left(Y-\overline{Y}+\overline{Y}-f(X)\right)^{2} \mid X\right]$ $= \mathbf{E}_{Y} \left[(Y - \overline{Y})^{2} \mid X \right] + \mathbf{E}_{Y} \left[(\overline{Y} - f(X))^{2} \mid X \right] + 2 \mathbf{E}_{Y} \left[(Y - \overline{Y})(\overline{Y} - f(X)) \mid X \right]$ $= \mathbf{E}_{Y} \left[(Y - \overline{Y})^{2} \mid X \right] + \left(\overline{Y} - f(X) \right)^{2} + 2(\overline{Y} - f(X)) \mathbf{E}_{Y} \left[(Y - \overline{Y}) \mid X \right]$ $= \mathbf{E}_{Y} \left[(Y - \overline{Y})^{2} \mid X \right] + \left(\overline{Y} - f(X) \right)^{2}$

(Notice: $\mathbf{E}_{Y}[(Y - \overline{Y}) | X] = \mathbf{E}_{Y}[Y | X] - \mathbf{E}_{Y}[\overline{Y} | X] = \overline{Y} - \overline{Y} = 0$)

Optimal Model f(X) (cont.)

The choice of f(X) does not affect $E_{Y}\left[(Y-\overline{Y})^{2} \mid X\right]$ but $(\overline{Y} - f(X))^{2}$ is minimized for $f(X) = \overline{Y} = \mathbb{E}_{Y}[Y \mid X].$

Note that $\mathbb{E}_{XY}\left[(Y - f(X))^2\right] = \mathbb{E}_{X}\left[\mathbb{E}_{Y}\left[(Y - f(X))^2 \mid X\right]\right]$ Hence

 $E_{XY}[(Y - f(X))^2] = E_{Y}[E_{Y}[(Y - \overline{Y})^2 | X] + (\overline{Y} - f(X))^2]$

Hence the squared error is minimzed by choosing $f(X) = E_{y}[Y | X]$ for every X.

(Notice that for minimizing absolute error $E_{yy} \left[|Y - f(X)| \right]$, one can show that the best model is $f(X) = median(X \mid Y).)$

Interpreting the Result

- To minimize mean squared error, the best prediction for input X=x is the mean of the Y-values of all training records $\chi(i), \chi(i)$ with $\chi(i)=x$ = E.g. assume there are training records (5.2.1) (5.24), (5.26), (5.28). The optimal prediction for input X=5 would be estimated as (22+24+26+28)/4 = 25.
- Problem: to reliably estimate the mean of Y for a given X=x, we need sufficiently many training records with X=x. In practice, often there is only one or no training record at all for an X=x of interest.
- If there were many such records with X=x, we would not need a model and could just return the average Y for that X=x.
- The benefit of a good data mining technique is its ability to interpolate and extrapolate from known training records to make good predictions even for X-values that do not occur in the training data at all. Classification for two classes: encode as 0 and 1, use squared error as before
- Then f(X) = E[Y | X=x] = 1*Pr(Y=1 | X=x) + 0*Pr(Y=0 | X=x) = Pr(Y=1 | X=x)Classification for k classes: can show that for 0-1 loss (error = 0 if correct class, error = 1 if wrong class predicted) the optimal choice is to return the majority class for a given input X=x

This is called the Bayes classifier

Implications for Trees

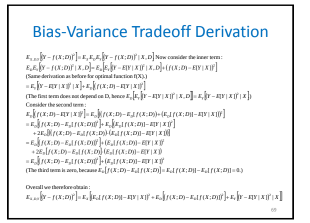
- Since there are not enough, or none at all, training records with X=x, the output for input X=x has to be based on records "in the neighborhood"
 - A tree leaf corresponds to a multi-dimensional range in the data space
- Records in the same leaf are neighbors of each other Solution: estimate mean Y for input X=x from the training
- records in the same leaf node that contains input X=x - Classification: leaf returns majority class or class probabilities
- (estimated from fraction of training records in the leaf) - Prediction: leaf returns average of Y-values or fits a local model
- Make sure there are enough training records in the leaf to obtain reliable estimates

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Bias-Variance Tradeoff

- Let's take this one step further and see if we can understand overfitting through statistical decision theory
- As before, consider two random variables X and Y
- · From a training set D with n records, we want to construct a function f(X) that returns good approximations of Y for future inputs X Make dependence of f on D explicit by writing f(X; D)
 - Goal: minimize mean squared error over all X, Y,
- and D, i.e., $E_{X,D,Y}[(Y f(X; D))^2]$

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Bias-Variance Tradeoff and Overfitting

 $(E_D[f(X;D)] - E[Y | X])^2$: bias $E_D[(f(X;D) - E_D[f(X;D)])^2]$: variance $E_D[(Y - E[Y + Y])^2 + X]$, investigating a surgery

 $E_{Y}\left[(Y - E[Y | X])^{2} | X\right]$: irreducible error (does not depend on f and is simply the variance of Y given X.)

- Option 1: f(X;D) = E[Y| X,D] - Bias: since E_D[E[Y| X,D]] = E[Y| X], bias is zero
- Bias: since E_D[E[Y] X,D]] = E[Y] X], bias is zero
 Variance: (E[Y] X,D]-E_D[E[Y] X,D]]² = (E[Y] X,D]-E[Y] X])² can be very large since E[Y] X,D] depends heavily on D
 Might overfit!
- Option 2: f(X;D)=X (or other function independent of D)
- Variance: (X-E_D[X])²=(X-X)²=0
- Bias: (E₀[X]-E[Y|X])²[X-E[Y|X])² can be large, because E[Y|X] might be completely different from X
 Might underfit!
- Find best compromise between fitting training data too closely (option 1) and completely ignoring it (option 2)

Implications for Trees

- Bias decreases as tree becomes larger
 Larger tree can fit training data better
- Variance increases as tree becomes larger

 Sample variance affects predictions of larger tree more
- Find right tradeoff as discussed earlier
 - Validation data to find best pruned tree
 - MDL principle

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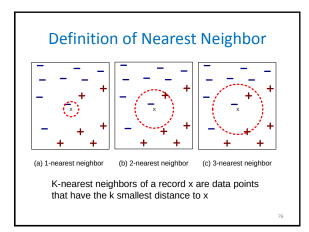
Lazy vs. Eager Learning

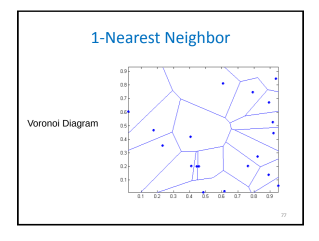
- Lazy learning: Simply stores training data (or only minor processing) and waits until it is given a test record
- Eager learning: Given a training set, constructs a classification model before receiving new (test) data to classify
- General trend: Lazy = faster training, slower predictions
- Accuracy: not clear which one is better!
 - Lazy method: typically driven by local decisions
 - Eager method: driven by global and local decisions

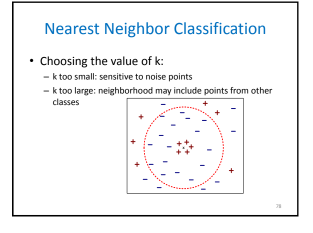
Nearest-Neighbor Recall our statistical decision theory analysis: Best prediction for input X=x is the mean of the Y-values of all records (x(i),y(i)) with x(i)=x (majority class for classification) Problem was to estimate E[Y| X=x] or majority class for X=x from the training data

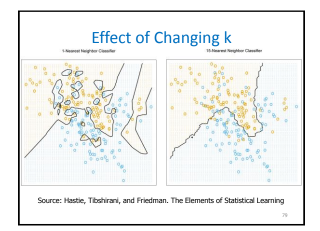
- Solution was to approximate it
 - Use Y-values from training records in neighborhood around X=x

Nearest-Neighbor Classifiers Unknown tuple Requires: . Set of stored records Distance metric for pairs of records on choice: Euclidean • Comr $d(\mathbf{p},\mathbf{q}) = \sqrt{\sum_{i} (p_i - q_i)^2}$ Parameter k Number of nearest neighbors to retrieve To classify a record: - Find its k nearest neighbors + - Determine output based on (distance-weighted) average of neighbors' output









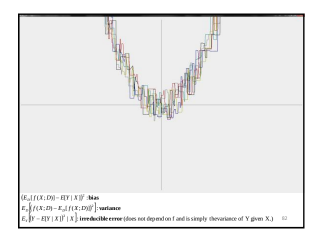
Explaining the Effect of k

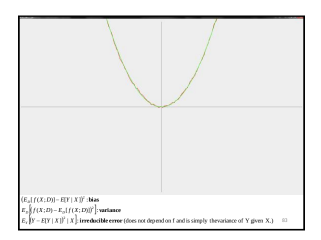
- · Recall the bias-variance tradeoff
- Small k, i.e., predictions based on few neighbors
 - High variance, low bias
- Large k, e.g., average over entire data set
 Low variance, but high bias
- Need to find k that achieves best tradeoff
- · Can do that using validation data

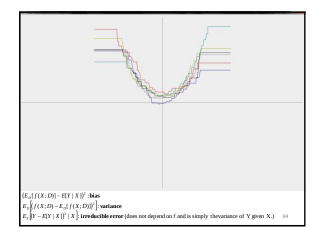
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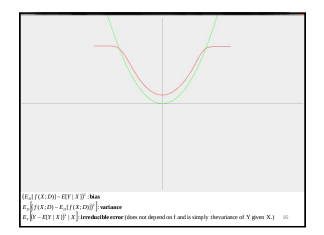
Experiment

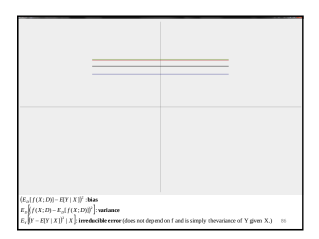
- 50 training points (x, y)
 - $-2 \le x \le 2$, selected uniformly at random
 - $-y = x^2 + \varepsilon$, where ε is selected uniformly at random from range [-0.5, 0.5]
- Test data sets: 500 points from same distribution as training data, but $\varepsilon=0$
- Plot 1: all (x, NN1(x)) for 5 test sets
- Plot 2: all (x, AVG(NN1(x))), averaged over 200 test data set
 - Same for NN20 and NN50

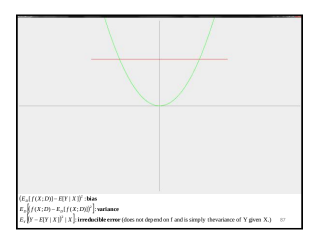












Scaling Issues

- Attributes may have to be scaled to prevent distance measures from being dominated by one of the attributes
- Example:
 - Height of a person may vary from 1.5m to 1.8m
 - Weight of a person may vary from 90lb to 300lb
 - Income of a person may vary from \$10K to \$1M
 - Income difference would dominate record distance

Other Problems

- Problem with Euclidean measure:
 - High dimensional data: curse of dimensionality - Can produce counter-intuitive results

111111111110	VS	1000000000000				
011111111111		0000000000000				
d = 1.4142		d = 1.4142				
 Solution: Normalize the vectors to unit length 						
Irrolovant attributo	c mia	at dominato distanco				

Irrelevant attributes might dominate distance Solution: eliminate them

Computational Cost

- Brute force: O(#trainingRecords)
 - For each training record, compute distance to test record, keep if among top-k
- Pre-compute Voronoi diagram (expensive), then search spatial index of Voronoi cells: if lucky O(log(#trainingRecords))
- Store training records in multi-dimensional search tree, e.g., R-tree: if lucky O(log(#trainingRecords))
- Bulk-compute predictions for many test records using spatial join between training and test set
 - Same worst-case cost as one-by-one predictions, but usually much faster in practice

Classification and Prediction Overview

- Introduction
- **Decision Trees**
- Statistical Decision Theory
- Nearest Neighbor
- **Bayesian Classification**
- Artificial Neural Networks
- Support Vector Machines (SVMs)
- Prediction
- Accuracy and Error Measures
- **Ensemble Methods**

Bayesian Classification

- Performs probabilistic prediction, i.e., predicts class membership probabilities
- Based on Bayes' Theorem
- Incremental training
 - Update probabilities as new training records arrive
- Can combine prior knowledge with observed data • Even when Bayesian methods are computationally intractable, they can provide a standard of optimal decision making against which other methods can be measured

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Bayesian Theorem: Basics

- X = random variable for data records ("evidence")
- H = hypothesis that specific record X=x belongs to class C
- Goal: determine P(H | X=x)
- Probability that hypothesis holds given a record x
- P(H) = prior probability- The initial probability of the hypothesis
- E.g., person x will buy computer, regardless of age, income etc. P(X=x) = probability that data record x is observed
- $P(\mathbf{X}=\mathbf{x} | \mathbf{H}) = \text{probability of observing record } \mathbf{x}$, given that the
- hypothesis holds
 - E.g., given that **x** will buy a computer, what is the probability that **x** is in age group 31...40, has medium income, etc.?

Bayes' Theorem

 Given data record x, the posterior probability of a hypothesis H, P(H | X=x), follows from Bayes theorem:

$$P(H | \mathbf{X} = \mathbf{x}) = \frac{P(\mathbf{X} = \mathbf{x} | H)P(H)}{P(\mathbf{X} = \mathbf{x})}$$

- Informally: posterior = likelihood * prior / evidence
 Among all candidate hypotheses H find the maxima
- Among all candidate hypotheses H, find the maximally probably one, called maximum a posteriori (MAP) hypothesis
- Note: P(X=x) is the same for all hypotheses
- If all hypotheses are equally probable a priori, we only need to compare P(X=x| H)
 Winning hypothesis is called the maximum likelihood (ML) hypothesis
- Winning hypothesis is called the maximum likelihood (ML) hypothesis
 Practical difficulties: requires initial knowledge of many probabilities and has high computational cost

Towards Naïve Bayes Classifier

- Suppose there are m classes C₁, C₂,..., C_m
- Classification goal: for record **x**, find class C_i that has the maximum posterior probability $P(C_i | X=x)$
- Bayes' theorem:

$$P(C_i | \mathbf{X} = \mathbf{x}) = \frac{P(\mathbf{X} = x | C_i) P(C_i)}{P(\mathbf{X} = \mathbf{x})}$$

 Since P(X=x) is the same for all classes, only need to find maximum of P(X=x|C_i)P(C_i)

Computing $P(\mathbf{X}=\mathbf{x} | C_i)$ and $P(C_i)$

- Estimate $P(C_i)$ by counting the frequency of class C_i in the training data
- Can we do the same for P(X=x|C_i)?
 - Need very large set of training data - Have $|X_1|^*|X_2|^*...^*|X_d|^*m$ different combinations of
 - possible values for X and C_i
 Need to see every instance x many times to obtain reliable estimates
- Solution: decompose into lower-dimensional problems

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Example: Computing $P(\mathbf{X}=\mathbf{x} | C_i)$ and $P(C_i)$

- P(buys_computer = yes) = 9/14
- P(buys_computer = no) = 5/14
- P(age>40, income=low, student=no, credit_rating=bad| buys_computer=yes) = 0 ?

Age	Income	Student	Credit_rating	Buys_computer
≤ 30	High	No	Bad	No
≤ 30	High	No	Good	No
3140	High	No	Bad	Yes
> 40	Medium	No	Bad	Yes
> 40	Low	Yes	Bad	Yes
> 40	Low	Yes	Good	No
3140	Low	Yes	Good	Yes
≤ 30	Medium	No	Bad	No
≤ 30	Low	Yes	Bad	Yes
> 40	Medium	Yes	Bad	Yes
≤ 30	Medium	Yes	Good	Yes
3140	Medium	No	Good	Yes
3140	High	Yes	Bad	Yes
> 40	Medium	No	Good	No

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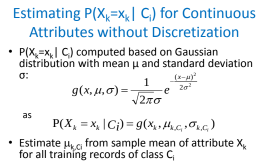
X, Y, Z random variables X is conditionally independent of Y, given Z, if P(X| Y,Z) = P(X| Z) Equivalent to: P(X,Y| Z) = P(X| Z) * P(Y| Z) Example: people with longer arms read better Confounding factor: age Young child has shorter arms and lacks reading skills of adult

 If age is fixed, observed relationship between arm length and reading skills disappears

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Derivation of Naïve Bayes Classifier • Simplifying assumption: all input attributes conditionally independent, given class $P(\mathbf{X} = (x_1, ..., x_d) | C_i) = \prod_{k=1}^{d} P(X_k = x_k | C_i) = P(X_1 = x_1 | C_i) \cdot P(X_2 = x_2 | C_i) \cdots P(X_d = x_d | C_i)$ • Each P($\mathbf{X}_k = \mathbf{x}_k | C_i$) can be estimated robustly – If \mathbf{X}_k is categorical attribute

- P(X_k = x_k | C_i) = #records in C_i that have value x_k for X_k, divided by #records of class C_i in training data set
- If X_k is continuous, we could discretize it
 - Problem: interval selection
 - Too many intervals: too few training cases per interval
 - Too few intervals: limited choices for decision boundary



- Estimate $\sigma_{\text{k,Ci}}$ similarly from sample

Naïve Bayes Example

Classes:					
– C ₁ :buys_computer = yes	5				
– C ₂ :buys computer = no	Age	Income	Student	Credit rating	Buys compute
2 7 = 1	≤ 3 0	High	No	Bad	No
	≤ 3 0	High	No	Good	No
	3140	High	No	Bad	Yes
	> 40	Medium	No	Bad	Yes
 Data sample x 	> 40	Low	Yes	Bad	Yes
Duta sumple x	> 40	Low	Yes	Good	No
$-$ age \leq 30,	3140		Yes	Good	Yes
0 ,		Medium		Bad	No
 income = medium, 	≤ 3 0	Low	Yes	Bad	Yes
المست مستنا فستاب فم		Medium		Bad	Yes
— student = yes, and		Medium		Good	Yes
– credit rating = bad		Medium		Good	Yes
- credit_lating = bad	3140		Yes	Bad	Yes
	> 40	Medium	No	Good	No
					117

Naïve Bayesian Computation

- Compute P(C_i) for each class: P(buys_computer = "yes") = 9/14 = 0.643 P(buys_computer = "no") = 5/14= 0.357

- Therefore we predict buys_computer = "yes" for input x = (age = "≤30", income = "medium", student = "yes", credit_rating = "bad")

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Zero-Probability Problem

Naïve Bayesian prediction requires each conditional probability to be non-zero (why?)

 $P(\mathbf{X} = (x_1, \dots, x_d) \mid C_i) = \prod_{k=1}^{n} P(X_k = x_k \mid C_i) = P(X_1 = x_1 \mid C_i) \cdot P(X_2 = x_2 \mid C_i) \cdots P(X_d = x_d \mid C_i)$

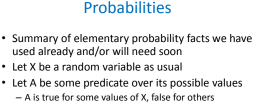
- Example: 1000 records for buys_computer=yes with income=low (0), income= medium (990), and income = high (10) For input with income=low, conditional probability is zero
- Use Laplacian correction (or Laplace estimator) by adding 1 dummy record to each income level
 - Prob(income = low) = 1/1003
 - Prob(income = medium) = 991/1003

 - Prob(income = high) = 11/1003
 "Corrected" probability estimates close to their "uncorrected" counterparts, but none is zero

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Naïve Bayesian Classifier: Comments

- · Easy to implement
- · Good results obtained in many cases
 - Robust to isolated noise points
 - Handles missing values by ignoring the instance during probability estimate calculations
 - Robust to irrelevant attributes
- Disadvantages
 - Assumption: class conditional independence, therefore loss of accuracy
 - Practically, dependencies exist among variables
- How to deal with these dependencies?



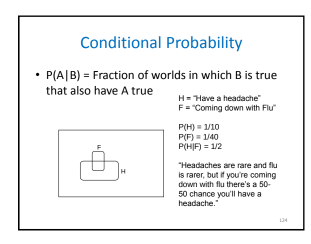
- E.g., X is outcome of throw of a die, A could be "value is greater than 4"
- P(A) is the fraction of possible worlds in which A is true
 - P(die value is greater than 4) = 2 / 6 = 1/3

Axioms

- $0 \le P(A) \le 1$
- P(True) = 1
- P(False) = 0
- $P(A \lor B) = P(A) + P(B) P(A \land B)$

Theorems from the Axioms

- $0 \le P(A) \le 1$, P(True) = 1, P(False) = 0
- $P(A \lor B) = P(A) + P(B) P(A \land B)$
- From these we can prove:
 P(not A) = P(~A) = 1 P(A)
 P(A) = P(A \land B) + P(A \land ~B)



Definition of Conditional Probability

 $P(A|B) = \frac{P(A \land B)}{P(B)}$

Corollary: the Chain Rule

$$P(A \land B) = P(A|B) P(B)$$

Multivalued Random Variables

- Suppose X can take on more than 2 values
- X is a random variable with arity k if it can take on exactly one value out of {v₁, v₂,..., v_k}
- Thus

$$P(X = v_i \land X = v_i) = 0 \text{ if } i \neq j$$

$$P(X = v_1 \lor X = v_2 \lor ... \lor X = v_k) = 1$$

Easy Fact about Multivalued Random Variables

• Using the axioms of probability $- 0 \le P(A) \le 1$, P(True) = 1, P(False) = 0 $- P(A \lor B) = P(A) + P(B) - P(A \land B)$

• And assuming that X obeys

$$P(X = v_i \land X = v_j) = 0 \text{ if } i \neq j$$

$$P(X = v_1 \lor X = v_2 \lor ... \lor X = v_k) = 1$$
• We can prove that

$$P(X = v_1 \lor X = v_2 \lor ... \lor X = v_i) = \sum_{j=1}^{i} P(X = v_j)$$

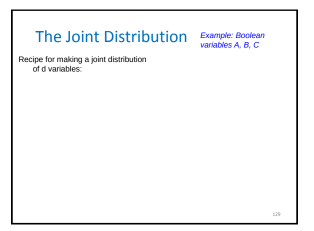
And therefore:
$$\sum_{j=1}^{N} P(X = v_j) = 1$$

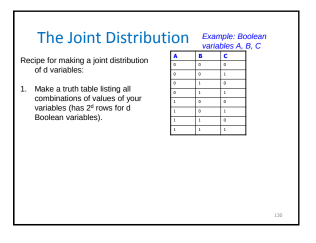
Useful Easy-to-Prove Facts

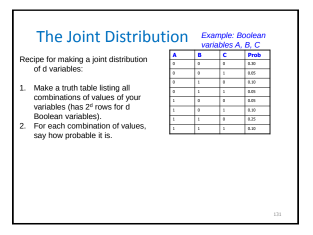
$$P(A | B)+P(\sim A | B) = 1$$

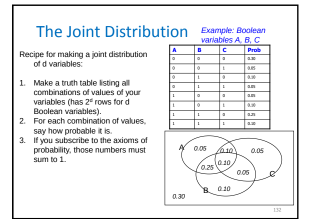
$$\sum_{j=1}^{k} P(X = v_j | B) = 1$$

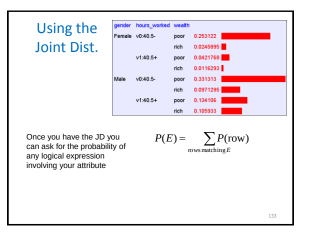
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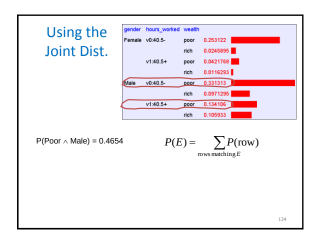


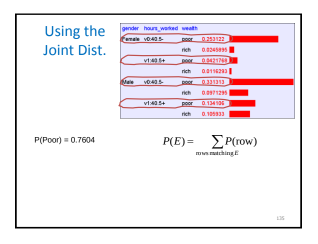


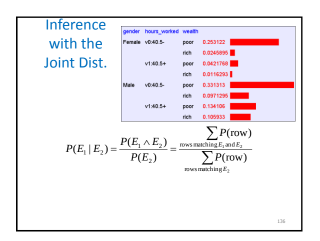


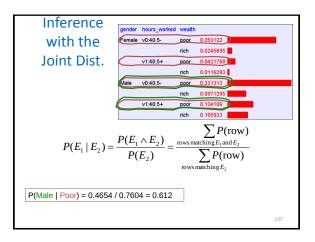


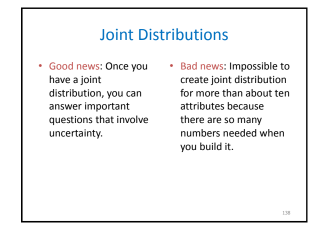








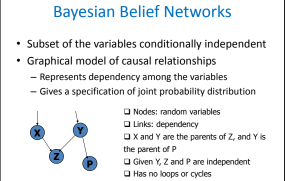




What Would Help?

- Full independence
 - P(gender=g ^ hours_worked=h ^ wealth=w) = P(gender=g) * P(hours_worked=h) * P(wealth=w)
 - Can reconstruct full joint distribution from a few marginals
- Full conditional independence given class value

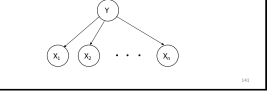
 Naïve Bayes
- What about something between Naïve Bayes and general joint distribution?

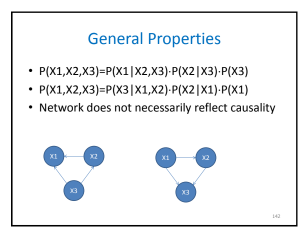


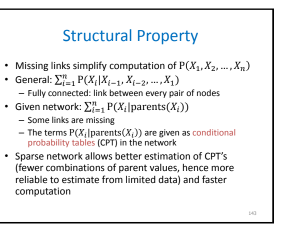
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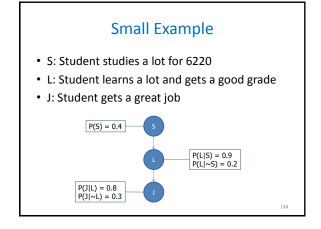
Bayesian Network Properties

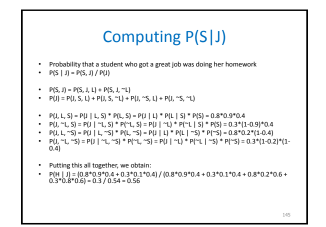
- Each variable is conditionally independent of its non-descendents in the graph, given its parents
- Naïve Bayes as a Bayesian network:

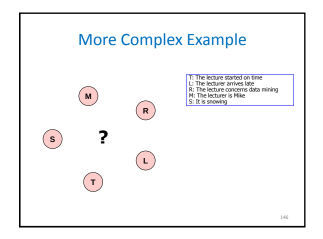


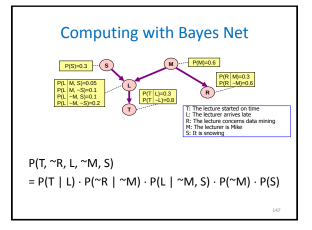


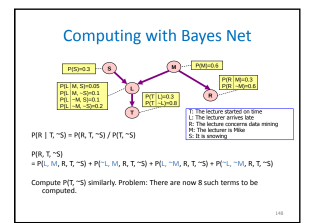


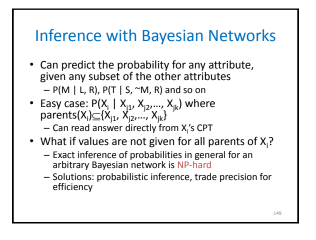












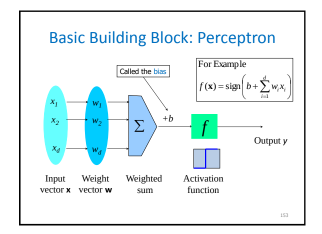
Training Bayesian Networks

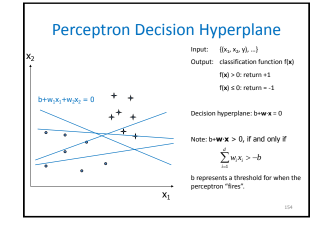
- Several scenarios:
 - Network structure known, all variables observable: learn only the CPTs
 - Network structure known, some hidden variables: gradient descent (greedy hill-climbing) method, analogous to neural network learning
 - Network structure unknown, all variables observable: search through the model space to reconstruct network topology
 - Unknown structure, all hidden variables: No good algorithms known for this purpose
- Ref.: D. Heckerman: Bayesian networks for data mining

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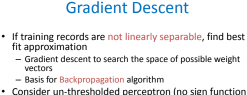
Representing Boolean Functions

- AND with two-input perceptron - b=-0.8, w₁=w₂=0.5
- OR with two-input perceptron - b=-0.3, w₁=w₂=0.5
- m-of-n function: true if at least m out of n inputs are true
 - All input weights 0.5, threshold weight b is set according to m, n
- Can also represent NAND, NOR
- What about XOR?

Perceptron Training Rule

- Goal: correct +1/-1 output for each training record
- Start with random weights, constant $\boldsymbol{\eta}$ (learning rate)
- While some training records are still incorrectly classified do
 - For each training record (x, y)
 - Let $f_{old}(\pmb{x})$ be the output of the current perceptron for \pmb{x}
 - Set b:= b + Δb , where Δb = η (y f_{old}(x))
 - + For all i, set w_i := w_i + Δw_i , where Δw_i = $\eta($ y $f_{old}(\textbf{x}))x_i$
- Converges to correct decision boundary, if the classes are linearly separable and a small enough η is used

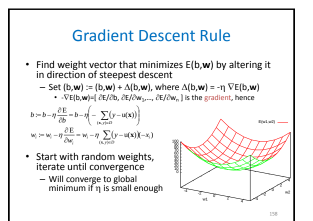
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- Consider <u>un-thresholded</u> perceptron (no sign function applied), i.e., $u(\mathbf{x}) = b + \mathbf{w} \cdot \mathbf{x}$
- Measure training error by squared error $E(b, \mathbf{w}) = \frac{1}{2} \sum_{(\mathbf{x}, y) \in D} (y \mathbf{u}(\mathbf{x}))^2$

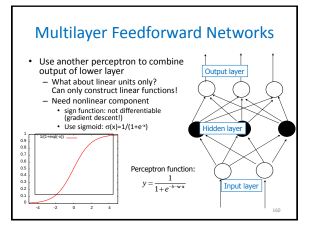
- D = training data

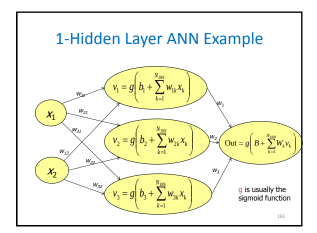
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Gradient Descent Summary

- Epoch updating (batch mode)
 - Compute gradient over entire training set
 - Changes model once per scan of entire training set
- Case updating (incremental mode, stochastic gradient descent)
 - Compute gradient for a single training record
 - Changes model after every single training record immediately
- Case updating can approximate epoch updating arbitrarily close if $\boldsymbol{\eta}$ is small enough
- What is the difference between perceptron training rule and case updating for gradient descent?
 - Error computation on thresholded vs. unthresholded function





Making Predictions

- Input record fed simultaneously into the units of the input layer
- Then weighted and fed simultaneously to a hidden layer
- Weighted outputs of the last hidden layer are the input to the units in the output layer, which emits the network's prediction
- The network is feed-forward
 None of the weights cycles back to an input unit or to an output unit of a previous layer
- Statistical point of view: neural networks perform nonlinear regression

Backpropagation Algorithm

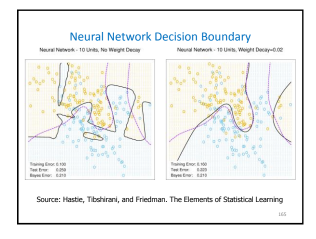
- Earlier discussion: gradient descent for a *single* perceptron using a simple un-thresholded function
- If sigmoid (or other differentiable) function is applied to weighted sum, use complete function for gradient descent
- Multiple perceptrons: optimize over all weights of all perceptrons
 - Problems: huge search space, local minima
- Backpropagation
- Initialize all weights with small random values
- Iterate many times
 - Compute gradient, starting at output and working back
 Error of hidden unit h: how do we get the true output value? Use weighted
 sum of errors of each unit influenced by h
 - Update all weights in the network

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Overfitting

- When do we stop updating the weights?
- · Overfitting tends to happen in later iterations
 - Weights initially small random values
 - Weights all similar => smooth decision surface
 - Surface complexity increases as weights diverge
- Preventing overfitting
 - Weight decay: decrease each weight by small factor during each iteration, or
 - Use validation data to decide when to stop iterating



Backpropagation Remarks

- · Computational cost
 - Each iteration costs O(|D|*|w|), with |D| training records and |w| weights
 - Number of iterations can be exponential in n, the number of inputs (in practice often tens of thousands)
- Local minima can trap the gradient descent algorithm: convergence guaranteed to *local* minimum, not *global*
- Backpropagation highly effective in practice

 Many variants to deal with local minima issue, use of case updating

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Defining a Network

- 1. Decide network topology
 - #input units, #hidden layers, #units per hidden layer, #output units (one output unit per class for problems with >2 classes)
- Normalize input values for each attribute to [0.0, 1.0]
 Nominal/ordinal attributes: one input unit per domain value
 For attribute grade with values A. B. C. have 3 input that are set to be a set of the s
 - For attribute *grade* with values A, B, C, have 3 inputs that are set to 1,0,0 for grade A, to 0,1,0 for grade B, and 0,0,1 for C
 - Why not map it to a single input with domain [0.0, 1.0]?
- 3. Choose learning rate $\eta, \, e.g., \, 1 \, / \, (\text{\#training iterations}) \\ \, \text{Too small: takes too long to converge}$
 - Too large: might never converge (oversteps minimum)
- 4. Bad results on test data? Change network topology, initial weights, or learning rate; try again.

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Representational Power

• Boolean functions

- Each can be represented by a 2-layer network
- Number of hidden units can grow exponentially with number of inputs
 - Create hidden unit for each input record
 - Set its weights to activate only for that input
 - Implement output unit as OR gate that only activates for desired output patterns
- · Continuous functions
 - Every bounded continuous function can be approximated arbitrarily close by a 2-layer network
- Any function can be approximated arbitrarily close by a 3-layer network

Neural Network as a Classifier

- Weaknesses
 - Long training time
 - Many non-trivial parameters, e.g., network topology
 - Poor interpretability: What is the meaning behind learned weights and hidden units?
 Nets bidden units are alternative representation of input value
- Note: hidden units are alternative representation of input values, capturing their relevant features
 Strengths
- High tolerance to noisy data
- Well-suited for continuous-valued inputs and outputs
- Successful on a wide array of real-world data
- Techniques exist for extraction of rules from neural networks

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SVM—Support Vector Machines

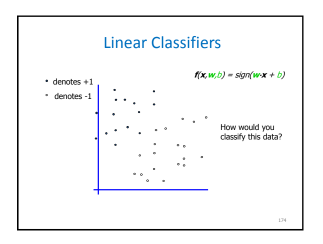
- Newer and very popular classification method
- Uses a nonlinear mapping to transform the original training data into a higher dimension
- Searches for the optimal separating hyperplane (i.e., "decision boundary") in the new dimension
- SVM finds this hyperplane using support vectors ("essential" training records) and margins (defined by the support vectors)

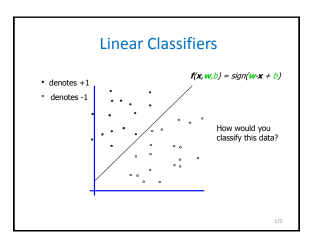
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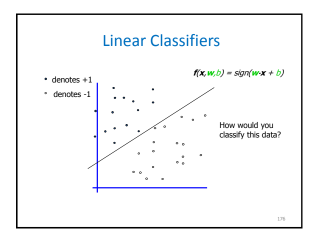
SVM—History and Applications

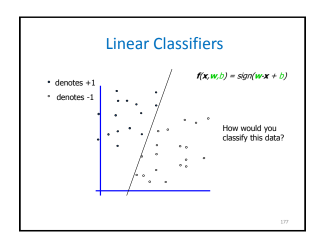
- Vapnik and colleagues (1992)
 - Groundwork from Vapnik & Chervonenkis' statistical learning theory in 1960s
- Training can be slow but accuracy is high

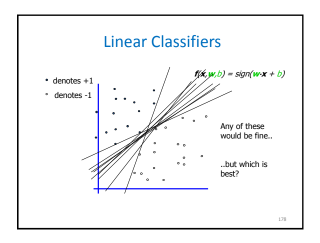
 Ability to model complex nonlinear decision boundaries (margin maximization)
- Used both for classification and prediction
- Applications: handwritten digit recognition, object recognition, speaker identification, benchmarking time-series prediction tests

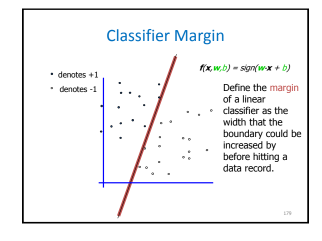


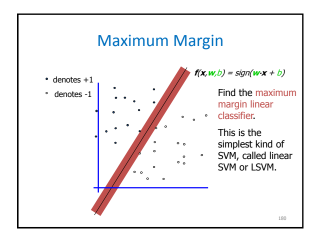


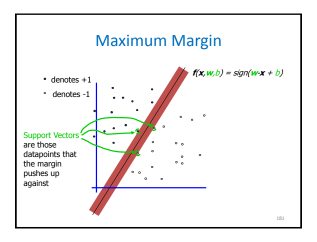


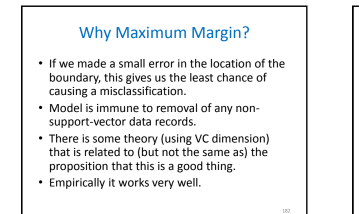


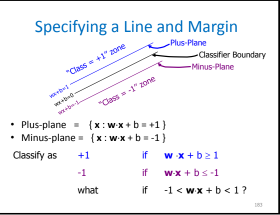


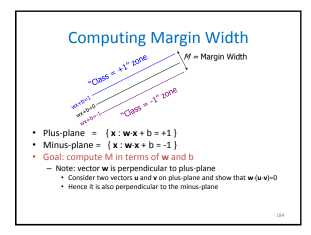


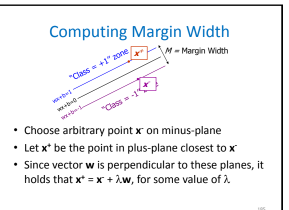




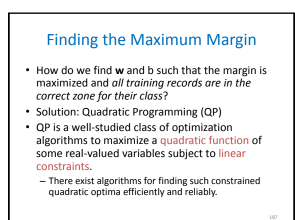


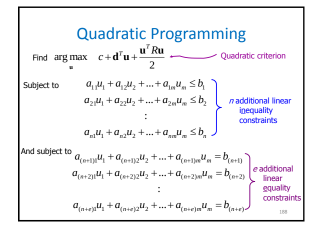


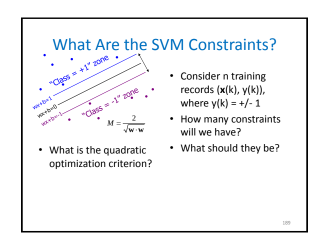


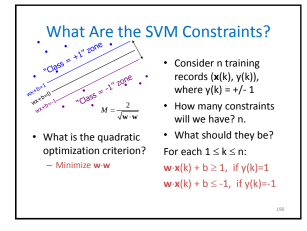


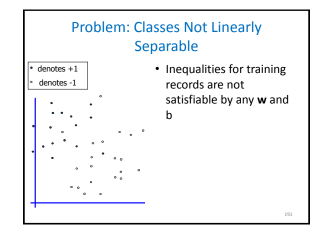
Putting It All Together • We have so far: $- w \cdot x^* + b = +1$ and $w \cdot x^* + b = +1$ $- x^* = x^* + \lambda w$ $- |x^* - x^*| = M$ • **Derivation**: $- w \cdot (x^* + \lambda w) + b = +1$, hence $w \cdot x^* + b + w \lambda w = 1$ $- 1 his implies <math>\lambda w \cdot w = 2$, i.e., $\lambda = 2 / w \cdot w$ $- 1 his implies <math>\lambda w \cdot w = 2$, i.e., $\lambda = 2 / w \cdot w$ $- 2 his m plies (w \cdot w)^{0.5} / (w \cdot w) = \lambda ((w \cdot w))^{0.5}$

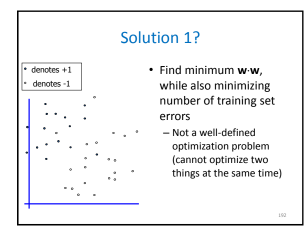


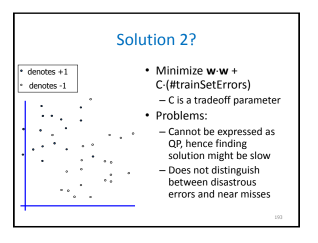


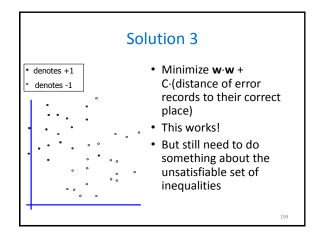


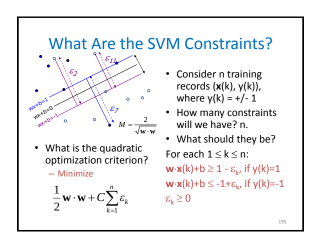






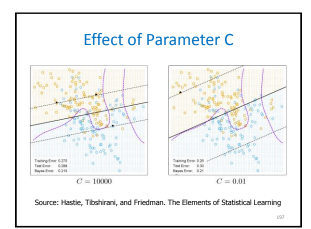


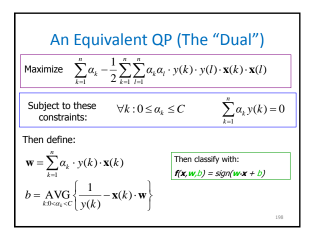


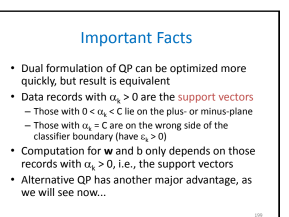


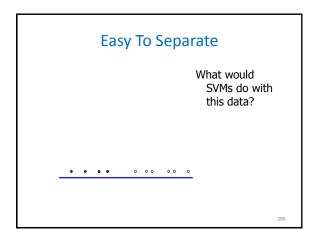
Facts About the New Problem Formulation

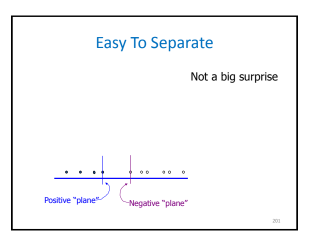
- Original QP formulation had d+1 variables
 - w_1 , w_2 ,..., w_d and b
- New QP formulation has d+1+n variables
 - $-w_1, w_2, ..., w_d$ and b
 - **-** ε₁, ε₂,..., ε_n
- C is a new parameter that needs to be set for the SVM
 - Controls tradeoff between paying attention to margin size versus misclassifications

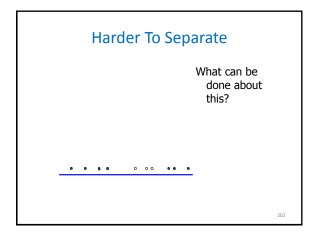


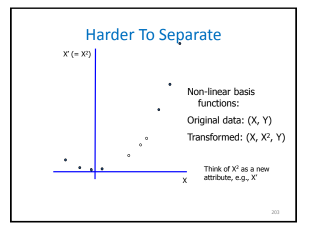


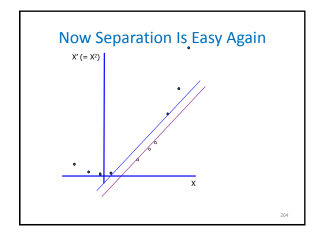


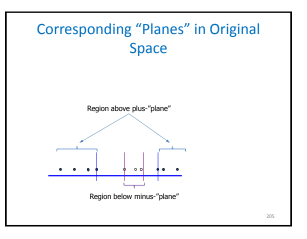


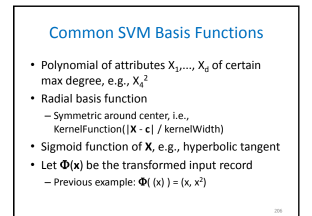


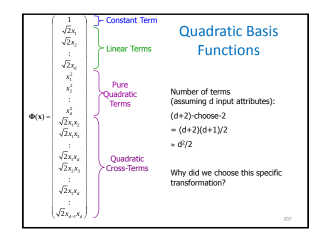


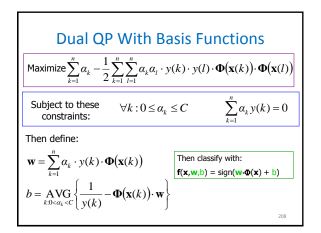




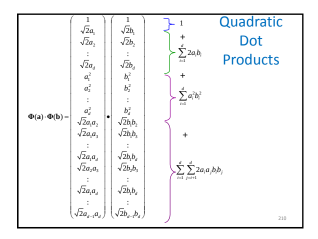


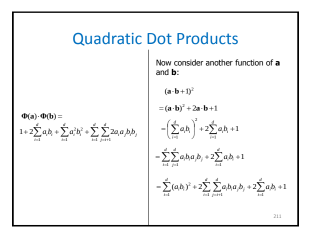






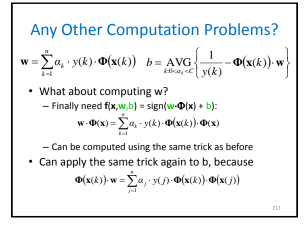
Computation Challenge Input vector x has d components (its d attribute values) The transformed input vector Φ(x) has d²/2 components Hence computing Φ(x(k))·Φ(x(l)) now costs order d²/2 instead of order d operations (additions, multiplications) ...or is there a better way to do this? Take advantage of properties of certain transformations

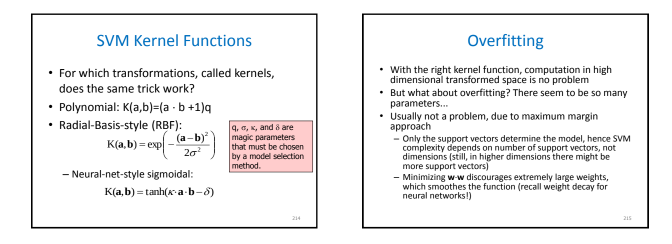


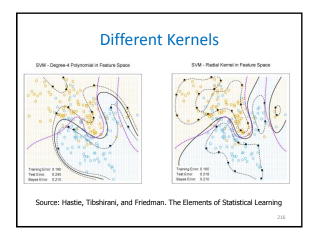


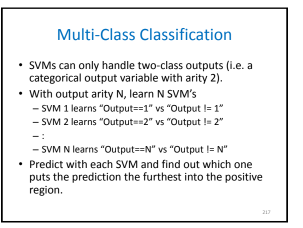
Quadratic Dot Products

- The results of $\Phi(\textbf{a}) \cdot \Phi(\textbf{b})$ and of $(\textbf{a} \cdot \textbf{b} + 1)^2$ are identical
- Computing Φ(a)·Φ(b) costs about d²/2, while computing (a·b+1)² costs only about d+2 operations
- This means that we can work in the high-dimensional space (d²/2 dimensions) where the training records are more easily separable, but pay about the same cost as working in the original space (d dimensions)
- Savings are even greater when dealing with higherdegree polynomials, i.e., degree q>2, that can be computed as (a·b+1)^q



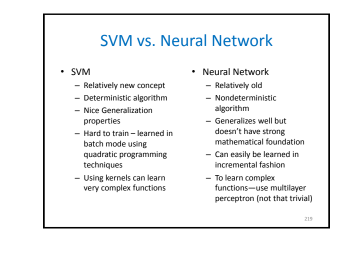






Why Is SVM Effective on High Dimensional Data?

- Complexity of trained classifier is characterized by the number of support vectors, not dimensionality of the data
- If all other training records are removed and training is repeated, the same separating hyperplane would be found
- The number of support vectors can be used to compute an upper bound on the expected error rate of the SVM, which is independent of data dimensionality
- Thus, an SVM with a small number of support vectors can have good generalization, even when the dimensionality of the data is high



Classification and Prediction Overview

- Introduction
- Decision Trees
- Statistical Decision Theory
- · Nearest Neighbor
- Bayesian Classification
- Artificial Neural Networks
- Support Vector Machines (SVMs)
- Prediction
- Accuracy and Error Measures
- Ensemble Methods

What Is Prediction?

- Essentially the same as classification, but output is continuous, not discrete
 - Construct a model, then use model to predict continuous output value for a given input
- Major method for prediction: regression

 Many variants of regression analysis in statistics literature; not covered in this class
- Neural network and k-NN can do regression "outof-the-box"
- · SVMs for regression exist
- What about trees?

Regression Trees and Model Trees

- Regression tree: proposed in CART system (Breiman et al. 1984)
 - CART: Classification And Regression Trees
 - Each leaf stores a continuous-valued prediction
 Average output value for the training records in the leaf
- Model tree: proposed by Quinlan (1992)

 Each leaf holds a regression model—a multivariate linear equation
- Training: like for classification trees, but uses variance instead of purity measure for selecting split predicates

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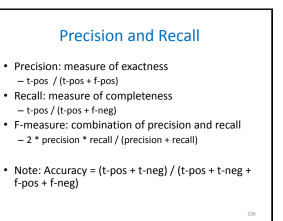
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Classification and Prediction Overview

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Classifier Accuracy Measures Predicted class total buy_computer = yes buy_computer = no 7000 True class | buy_computer = yes 6954 46 412 2588 3000 buy_computer = no total 7366 2634 10000 Accuracy of a classifier M, acc(M): percentage of test records that are correctly classified by M Error rate (misclassification rate) of M = 1 – acc(M) - Given m classes, CM[i,j], an entry in a confusion matrix, indicates # of records in class i that are labeled by the classifier as class j C_1 C₂ True positive False negative

C2 False positive True negative



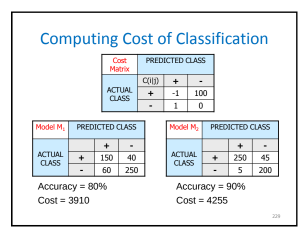
Limitation of Accuracy

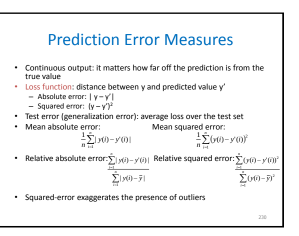
- · Consider a 2-class problem
 - Number of Class 0 examples = 9990
 - Number of Class 1 examples = 10
- If model predicts everything to be class 0, accuracy is 9990/10000 = 99.9 %
 - Accuracy is misleading because model does not detect any class 1 example
- Always predicting the majority class defines the baseline
 - A good classifier should do better than baseline

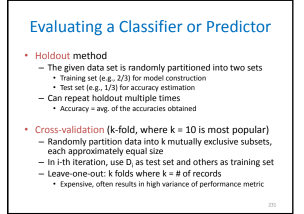
Cost-Sensitive Measures: Cost Matrix

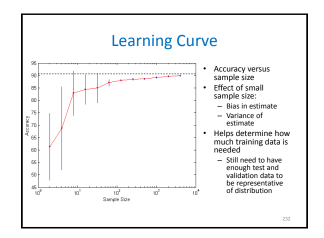
	PI	REDICTED	CLASS		
	C(i j)	Class=Yes	Class=No		
ACTUAL	Class=Yes	C(Yes Yes)	C(No Yes)		
CLASS	Class=No		C(No No)		

C(i| j): Cost of misclassifying class j example as class i



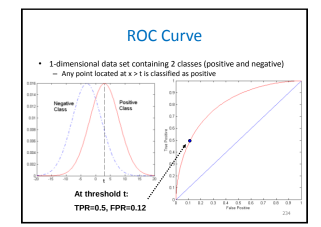


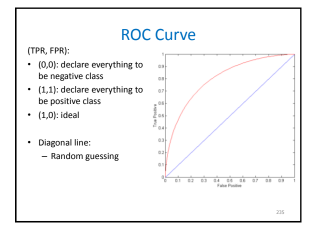




ROC (Receiver Operating Characteristic)

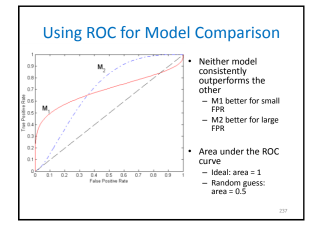
- Developed in 1950s for signal detection theory to analyze noisy signals
 - Characterizes trade-off between positive hits and false alarms
- ROC curve plots T-Pos rate (y-axis) against F-Pos rate (x-axis)
- Performance of each classifier is represented as a point on the ROC curve
 - Changing the threshold of the algorithm, sample distribution or cost matrix changes the location of the point

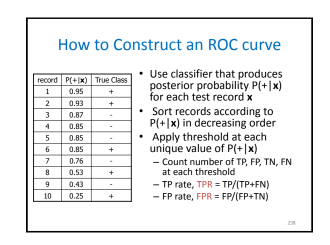


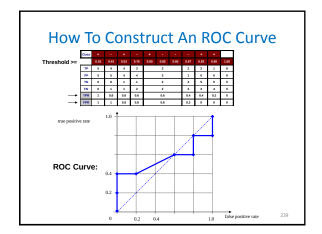


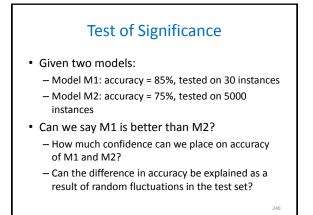
Diagonal Line for Random Guessing

- Classify a record as positive with fixed probability p, irrespective of attribute values
- Consider test set with a positive and b negative records
- True positives: p*a, hence true positive rate = (p*a)/a = p
- False positives: p*b, hence false positive rate = (p*b)/b = p
- For every value 0≤p≤1, we get point (p,p) on ROC curve









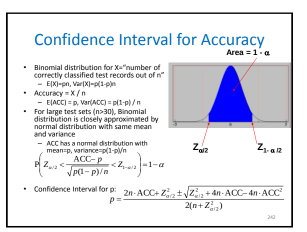
Confidence Interval for Accuracy

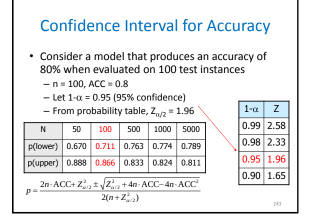
- - Collection of Bernoulli trials has a Binomial distribution
 - Probability of getting c correct predictions if model accuracy is p (=probability to get a single prediction right):

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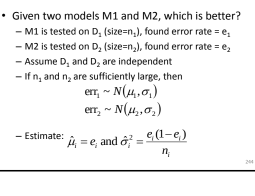
$$\binom{n}{c}p^{c}(1-p)$$

 Given c, or equivalently, ACC = c / n and n (#test records), can we predict p, the true accuracy of the model?





Comparing Performance of Two Models

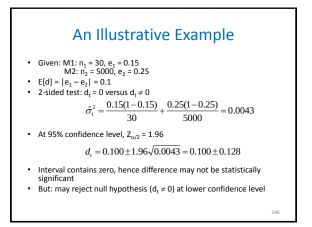


Testing Significance of Accuracy Difference

- Consider random variable d = err₁-err₂
 - Since err₁, err₂ are normally distributed, so is their difference
 - Hence d ~ N (d_t, σ_t) where d_t is the true difference
- Estimator for d_t:
 - $\mathsf{E}[\mathsf{d}] = \mathsf{E}[\mathsf{err}_1 \operatorname{-} \mathsf{err}_2] = \mathsf{E}[\mathsf{err}_1] \mathsf{E}[\mathsf{err}_2] \approx \mathsf{e}_1 \mathsf{e}_2$
 - Since D_1 and D_2 are independent, variance adds up:

$$\hat{\sigma}_{t}^{2} = \hat{\sigma}_{1}^{2} + \hat{\sigma}_{2}^{2} = \frac{e_{1}(1-e_{1})}{n_{1}} + \frac{e_{2}(1-e_{2})}{n_{2}}$$

– At (1- α) confidence level, $d_t = \mathbb{E}[d] \pm Z_{\alpha/2} \hat{\sigma}_t$



Significance Test for K-Fold Cross-Validation

- Each learning algorithm produces k models:
 - L1 produces M11 , M12, ..., M1k
 - L2 produces M21 , M22, ..., M2k
- Both models are tested on the same test sets $\mathsf{D}_1,$ $\mathsf{D}_2, ..., \mathsf{D}_k$
 - For each test set, compute $d_j = e_{1,j} e_{2,j}$
 - For large enough k, d_j is normally distributed with mean d_t and variance σ_t - Estimate: $\sum_{k=1}^{k} \sum_{j=1}^{k-1} \sum_{$

stimate:

$$\hat{\sigma}_t^2 = \frac{\sum_{j=1}^{j=1} (d_j - \overline{d})^2}{k(k-1)}$$

$$d_t = \overline{d} \pm t_{1-\alpha,k-1} \hat{\sigma}_t$$

t-distribution: get t coefficient $t_{1-\alpha,k-1}$ from table by looking up confidence level $(1-\alpha)$ and degrees of freedom (k-1)

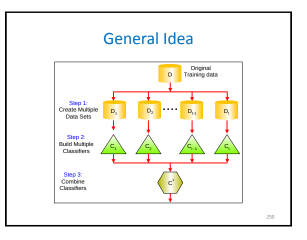
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Ensemble Methods

- Construct a set of classifiers from the training data
- Predict class label of previously unseen records by aggregating predictions made by multiple classifiers



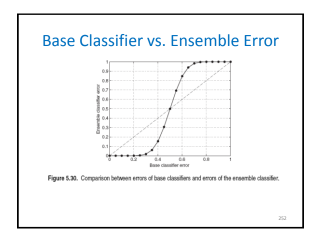
Why Does It Work?

- Consider 2-class problem
- Suppose there are 25 base classifiers

 Each classifier has error rate ε = 0.35
 - Assume the classifiers are independent
- Return majority vote of the 25 classifiers

 Probability that the ensemble classifier makes a wrong prediction: 25 (25)

 $\sum_{i=12}^{25} {\binom{25}{i}} \varepsilon^{i} (1-\varepsilon)^{25-i} = 0.06$



Model Averaging and Bias-Variance Tradeoff

- Single model: lowering bias will usually increase variance
 - "Smoother" model has lower variance but might not model function well enough
- · Ensembles can overcome this problem
 - 1. Let models overfit
 - Low bias, high variance
 - 2. Take care of the variance problem by averaging many of these models
- This is the basic idea behind bagging

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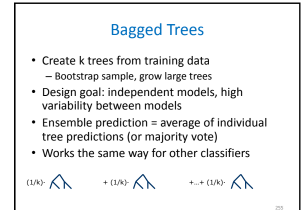
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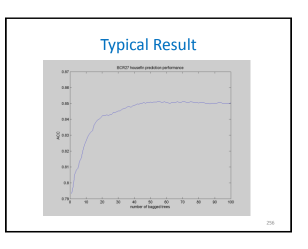
Bagging: Bootstrap Aggregation

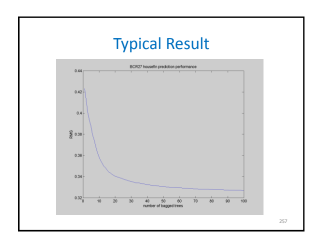
• Given training set with n records, sample n records randomly with replacement

Original Data	1	2	3	4	5	6	7	8	9	10
Bagging (Round 1)	7	8	10	8	2	5	10	10	5	9
Bagging (Round 2)	1	4	9	1	2	3	2	7	3	2
Bagging (Round 3)	1	8	5	10	5	5	9	6	3	7

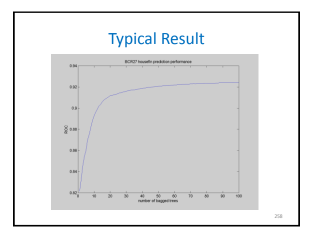
- Train classifier for each bootstrap sample
- Note: each training record has probability 1 – (1 – 1/n)ⁿ of being selected at least once in a sample of size n



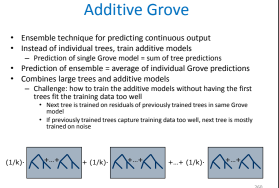


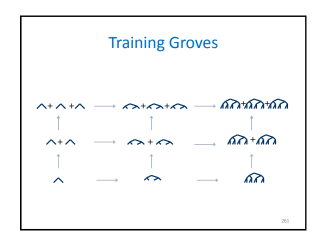


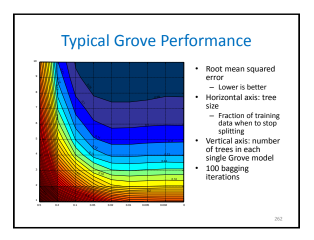
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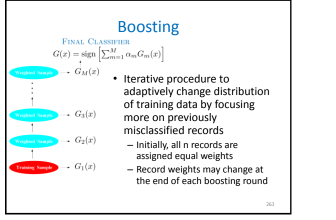


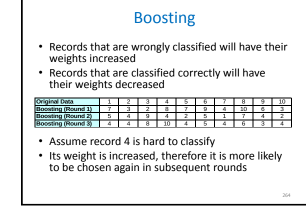
Bagging Challenges · Ideal case: all models independent of each other • Train on independent data samples - Problem: limited amount of training data Training set needs to be representative of data distribution Bootstrap sampling allows creation of many "almost" independent training sets · Diversify models, because similar sample might result in similar tree - Random Forest: limit choice of split attributes to small random subset of attributes (new selection of subset for each node) when training tree Use different model types in same ensemble: tree, ANN, SVM, regression models

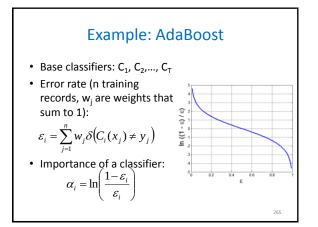


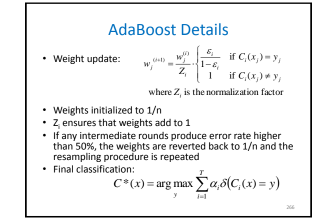


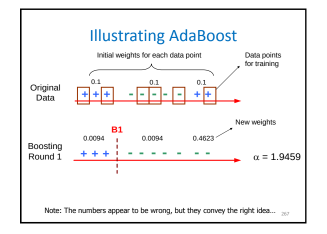


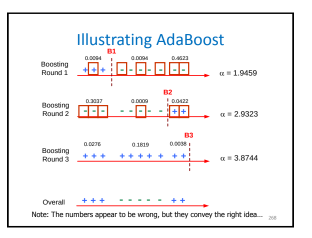












Bagging vs. Boosting

- Analogy
 - Bagging: diagnosis based on multiple doctors' majority vote - Boosting: weighted vote, based on doctors' previous diagnosis accuracy
 - Sampling procedure
 - Bagging: records have same weight; easy to train in parallel
 - Boosting: weights record higher if model predicts it wrong; inherently sequential process
- Overfitting

 - Bagging robust against overfitting
 - Boosting susceptible to overfitting: make sure individual models do not overfit
- · Accuracy usually significantly better than a single classifier Best boosted model often better than best bagged model
- Additive Grove
 - Combines strengths of bagging and boosting (additive models)
 - Shown empirically to make better predictions on many data sets
 Training more tricky, especially when data is very noisy

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Classification/Prediction Summary

- Forms of data analysis that can be used to train models • from data and then make predictions for new records
- Effective and scalable methods have been developed for decision tree induction, Naive Bayesian classification, Bayesian networks, rule-based classifiers, Backpropagation, Support Vector Machines (SVM), nearest neighbor classifiers, and many other classification methods
- Regression models are popular for prediction. Regression trees, model trees, and ANNs are also used for prediction.

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Classification/Prediction Summary

- . K-fold cross-validation is a popular method for accuracy estimation, but determining accuracy on large test set is equally accepted
 - If test sets are large enough, a significance test for finding the best model is not necessary Area under ROC curve and many other common performance
- measures exist
- Ensemble methods like bagging and boosting can be used to increase overall accuracy by learning and combining a series of individual models
- Often state-of-the-art in prediction quality, but expensive to train, store, use · No single method is superior over all others for all data sets
- Issues such as accuracy, training and prediction time, robustness, interpretability, and scalability must be considered and can involve trade-offs