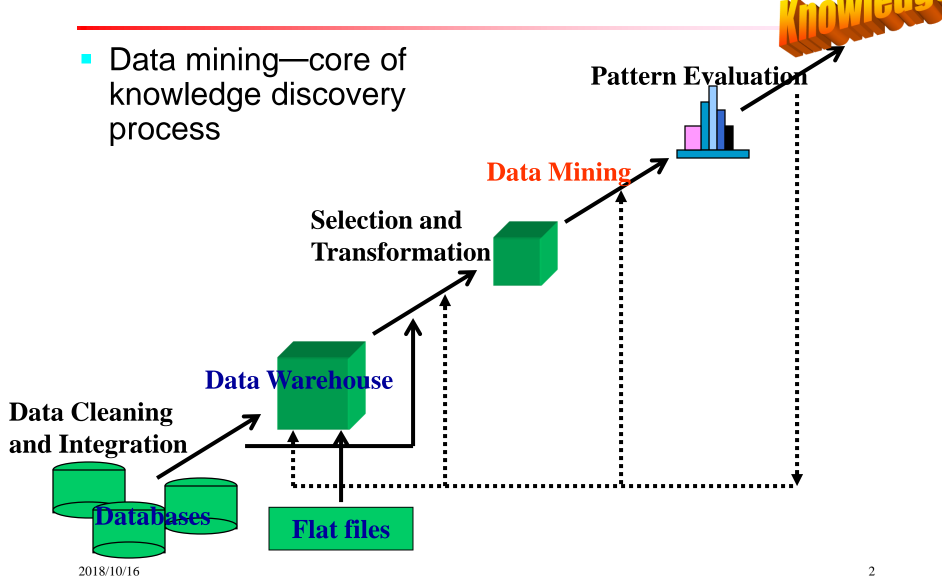
## **Data Mining**

Ying Liu, Prof., Ph.D

University of Chinese Academy of Sciences

#### **Review**



#### Classification and Prediction

- What is classification?
  What is prediction?
- Issues regarding classification and prediction
- Classification by decision tree induction
- Bayesian classification
- Classification by back propagation

- Other classification methods
- Prediction
- Accuracy and error measures
- Summary

#### Classification vs. Prediction

#### Classification

- Predict categorical class labels (discrete or nominal)
- Classify records (constructs a model) based on the training set and the class labels in a classifying attribute and then use the rules to classify new records

#### Prediction

 Model continuous-valued functions, i.e., predict unknown or missing values

#### Typical applications

- Credit approval
- Target marketing
- Medical diagnosis
- Fraud detection
- Intrusion detection

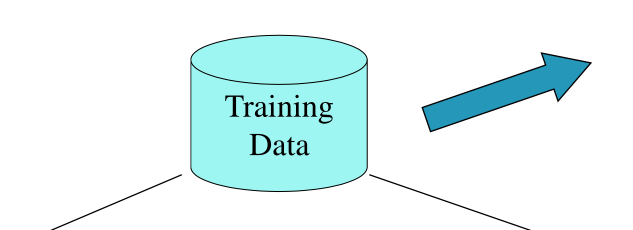
## Classification—A Two-Step Process

- Model construction: describing a set of predetermined classes
  - Each tuple/sample is assumed to belong to a predefined class, as determined by the class label attribute
  - The set of tuples used for model construction is training set
  - The model is represented as classification rules, decision trees, or mathematical formulae

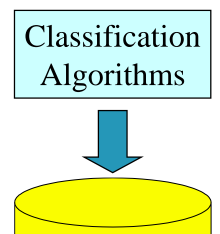
## Classification—A Two-Step Process

- Model usage: for classifying future or unknown objects
  - Estimate accuracy of the model
    - The known label of test sample is compared with the classified result from the model
    - Accuracy rate is the percentage of test set samples that are correctly classified by the model
    - Test set is independent of training set, otherwise over-fitting will occur
  - If the accuracy is acceptable, use the model to classify data tuples whose class labels are not known

### **Process (1): Model Construction**



NAME	RANK	YEARS	TENURED
Mike	Assistant Prof	3	no
Mary	Assistant Prof	7	yes
Bill	Professor	2	yes
Jim	Associate Prof	7	yes
Dave	Assistant Prof	6	no
Anne	Associate Prof	3	no

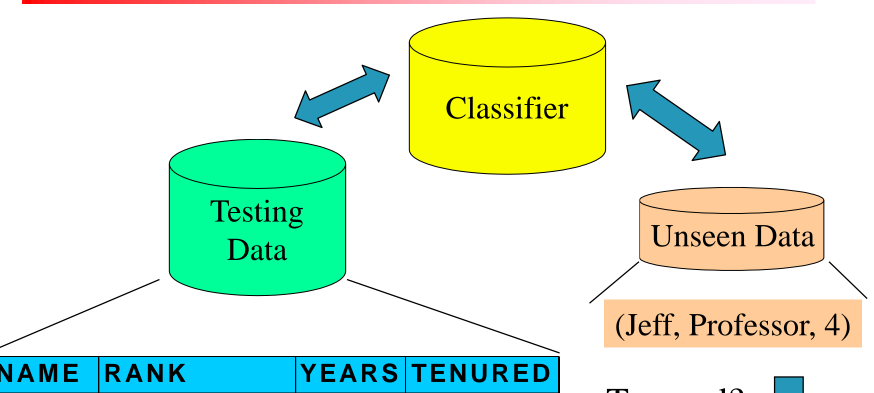


Classifier

(Model)

IF rank = 'professor' OR years > 6 THEN tenured = 'yes'

#### **Process (2): Using the Model in Classification**



NAME	RANK	YEARS	TENURED
Tom	Assistant Prof	2	no
Merlisa	Associate Prof	7	no
George	Professor	5	yes
Joseph	Assistant Prof	7	yes

Tenured?





## Supervised vs. Unsupervised Learning

- Supervised learning (classification)
  - Supervision: The training data (observations, measurements, etc.) are accompanied by labels indicating the class of the observations
  - New data is classified based on the training set
- Unsupervised learning (clustering)
  - The class labels of training data is unknown
  - Given a set of measurements, establish classes or clusters in the data

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## **Issues: Data Preparation**

- Data cleaning
  - Preprocess data in order to reduce noise and handle missing values
- Relevance analysis (feature selection)
  - Remove the irrelevant or redundant attributes
- Data transformation
  - Generalize and/or normalize data

#### **Issues: Evaluating Classification Methods**

#### Accuracy

- classifier accuracy: predicting class label
- predictor accuracy: guessing value of predicted attributes

#### Speed

- time to construct the model (training time)
- time to use the model (classification/prediction time)
- Robustness: handling noise and missing values
- Scalability: efficiency in disk-resident databases 可扩展性
- Interpretability 可解释性
  - understanding and insight provided by the model
- Other measures, e.g., goodness of rules, such as decision tree size or compactness of classification rules

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## Classification by Decision Tree Induction

- Decision tree
  - A flow-chart-like tree structure
  - Internal node denotes a splitting test on an attribute
  - Branch represents an outcome of the test
  - Leaf node represents class distribution
- Decision tree generation -- two phases
  - Tree construction
    - At start, all the training examples are at the root
    - Partition examples recursively based on selected attributes
  - Tree pruning
    - Identify and remove branches that reflect noise or outliers
- Use of decision tree: Classifying an unknown sample

## Classification by Decision Tree Induction

**Generate\_decision\_tree** (*D, attribute\_list*)

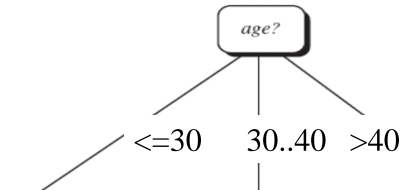
- (1) create a node N;
- (2) **if** tuples in *D* are all of the same class, *C* **then**
- (3) return N as a leaf node labeled with the class C;
- (4) **if** attribute\_list is empty **then**
- (5) return N as a leaf node labeled with the majority class in D; // majority voting
- (6) apply Attribute\_selection\_method(*D*, attribute\_list) to find the highest information gain;
- (7) label node *N* with *test-attribute*;
- (8) **for each** value  $a_i$  of *test-attribute* // partition the tuples and grow subtrees for each partition
- (9) Grow a branch from node N for test-attribute =  $a_i$ ; // a partition
- (10) Let  $s_i$  be the set of samples in D for which test-attribute =  $a_i$ ;
- (11) **if**  $s_i$  is empty **then**
- (12) attach a leaf labeled with the majority class in *D* to node *N*;
- (13) else attach the node returned by **Generate\_decision\_tree**(s<sub>i</sub>, attribute\_list) to node N;

<sup>2</sup>(計2) end for

## **Decision Tree Induction: Training Dataset**

age	income	student	credit_rating	buys_computer
<=30	high	no	fair	no
<=30	high	no	excellent	no
3140	high	no	fair	yes
>40	medium	no	fair	yes
>40	low	yes	fair	yes
>40	low	yes	excellent	no
3140	low	yes	excellent	yes
<=30	medium	no	fair	no
<=30	low	yes	fair	yes
>40	medium	yes	fair	yes
<=30	medium	yes	excellent	yes
3140	medium	no	excellent	yes
3140	high	yes	fair	yes
>40	medium	no	excellent	no

## **Decision Tree**

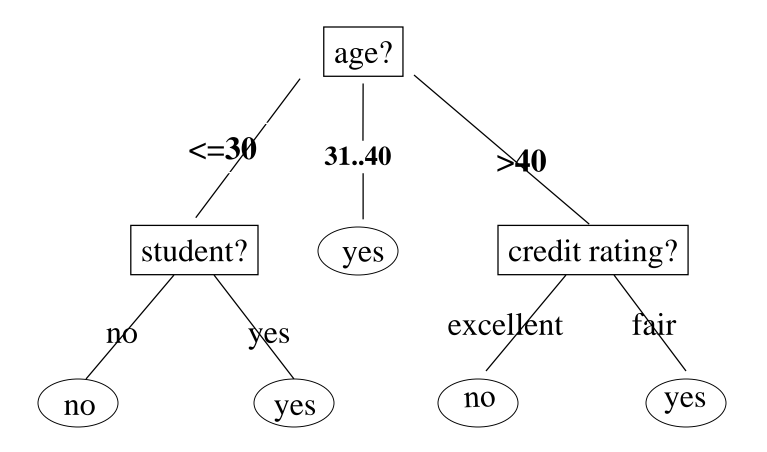


income	student	credit_rating	class
high	no	fair	no
high	no	excellent	no
medium	no	fair	no
low	yes	fair	yes
medium	yes	excellent	yes

income	student	credit_rating	class	
medium low low medium medium	no yes yes yes no	fair fair excellent fair excellent	yes yes no yes no	

income	student	credit_rating	class
high	no	fair	yes
low	yes	excellent	yes
medium	no	excellent	yes
high	yes	fair	yes

#### Output: A Decision Tree for "buys\_computer"



## Algorithm for Decision Tree Induction

- Basic algorithm (a greedy algorithm)
  - Tree is constructed in a top-down recursive divide-and-conquer manner
  - At start, all the training examples are at the root
  - Attributes are categorical (if continuous-valued, they are discretized in advance)
  - Examples are partitioned recursively based on selected attributes
  - Test attributes are selected on the basis of a heuristic or statistical measure (e.g., information gain, Gini index)
- Conditions for stopping partitioning
  - All samples for a given node belong to the same class
  - There are no remaining attributes for further partitioning majority voting is employed for classifying the leaf

#### **Information Gain (ID3/C4.5)**

- Select the attribute with the highest information gain
- Assume there are two classes, P and N
  - Let the set of examples S contain p elements of class P and n elements of class N
  - The amount of information, needed to classify sample

$$I(p,n) = -\frac{p}{p+n}\log_2\frac{p}{p+n} - \frac{n}{p+n}\log_2\frac{n}{p+n}$$

## **Information Gain in Decision Tree Induction**

- Assume that attribute A have v distinct values, {a<sub>1</sub>, a<sub>2</sub>, ..., a<sub>v</sub>}
- Training set S will be partitioned into sets  $\{S_1, S_2, ..., S_{\nu}\}$ 
  - If S<sub>i</sub> contains p<sub>i</sub> examples of P and n<sub>i</sub> examples of N, the entropy, or the expected information based on the partitioning into subsets by attribute A is

$$E(A) = \sum_{i=1}^{\nu} \frac{p_i + n_i}{p + n} I(p_i, n_i)$$

Information gain of A

$$Gain(A) = I(p,n) - E(A)$$

## **Attribute Selection by Information Gain Computation**

- Class P: buys\_computer = "yes"
- Class N: buys\_computer = "no"
- I(p, n) = I(9, 5) = 0.940
- Compute the entropy for age:

age	p <sub>i</sub>	n <sub>i</sub>	I(p <sub>i</sub> , n <sub>i</sub> )
<=30	2	3	0.971
3040	4	0	0
>40	3	2	0.971

$$E(age) = \frac{5}{14}I(2,3) + \frac{4}{14}I(4,0) + \frac{5}{14}I(3,2) = 0.694$$

Hence

$$Gain(age) = I(p,n) - E(age)$$
$$= 0.246$$

#### **Exercise**

- Please calculate the information gain of income, student, and credit\_rating, respectively.
- Gain(income) = 0.029
- Gain(Student) = 0.151
- Gain(credit\_rating) = 0.048

#### **Gain Ratio for Attribute Selection (C4.5)**

- Information gain measure is biased towards attributes with a large number of values
- C4.5 (a successor of ID3) uses gain ratio to overcome the problem (normalization to information gain)  $SplitInfo_A(D) = -\sum_{j=1}^{\nu} \frac{|D_j|}{|D|} \times \log_2(\frac{|D_j|}{|D|})$

$$SplitInfo_{A}(D) = -\frac{2}{|D|} \times \log_{2}(\frac{1}{|D|})$$

$$SplitInfo_{A}(D) = -\frac{4}{14} \times \log_{2}(\frac{4}{14}) - \frac{6}{14} \times \log_{2}(\frac{6}{14}) - \frac{4}{14} \times \log_{2}(\frac{4}{14}) = 0.926$$

- GainRatio(A) = Gain(A)/SplitInfo(A)
- Ex.
  - gain\_ratio(income) = 0.029/0.926 = 0.031
- The attribute with the maximum gain ratio is selected as the splitting attribute

#### **Gini Index (CART, IBM Intelligent Miner)**

- If a data set T contains examples from n classes, gini index, gini(T) is defined as  $gini(T) = 1 \sum_{j=1}^{n} p_j^2$ 
  - where  $p_i$  is the relative frequency of class j in T.
- If a data set T is split into two subsets T<sub>1</sub> and T<sub>2</sub> with sizes N<sub>1</sub> and N<sub>2</sub> respectively, the gini index of the split data contains examples from n classes, the gini index of the split is defined as

$$gini_{split}(T) = \frac{N_1}{N}gini(T_1) + \frac{N_2}{N}gini(T_2)$$

■ The attribute provides the smallest gini<sub>split</sub>(T) is chosen to split the node (need to enumerate all possible splitting points for each attribute).

#### Gini index (CART, IBM IntelligentMiner)

- The lowest is the best
- All attributes are assumed continuous-valued
- Can be modified for categorical attributes
- Ex. D has 9 tuples in buys\_computer = "yes" and 5 in "no"

$$gini(D) = 1 - \left(\frac{9}{14}\right)^2 - \left(\frac{5}{14}\right)^2 = 0.459$$

Suppose the attribute income partitions D into 10 in D<sub>1</sub>: {medium, high} and 4 in D<sub>2</sub>

$$\begin{aligned} &gini_{income \in \{medium, \ high\}} \ (D) = & \left(\frac{10}{14}\right) Gini(D_1) + \left(\frac{4}{14}\right) Gini \ (D_2) \\ &= \frac{10}{14} (1 - (\frac{6}{10})^2 - (\frac{4}{10})^2) + \frac{4}{14} (1 - (\frac{1}{4})^2 - (\frac{3}{4})^2) \\ &= 0.450 \end{aligned}$$

#### **Extracting Classification Rules from Trees**

- Represent the knowledge in the form of IF-THEN rules
- One rule is created for each path from the root to a leaf
- Each attribute-value pair along a path forms a conjunction
- The leaf node holds the class prediction
- Rules are easier for humans to understand
- Example

```
IF age = "<=30" AND student = "no" THEN buys_computer = "no"

IF age = "<=30" AND student = "yes" THEN buys_computer = "yes"

IF age = "31...40" THEN buys_computer = "yes"

IF age = ">40" AND credit_rating = "excellent" THEN buys_computer = "no"

IF age = "<=30" AND credit_rating = "fair" THEN buys_computer = "yes"
```

## **Overfitting and Tree Pruning**

- Overfitting: An induced tree may overfit the training data
  - Too many branches, some may reflect anomalies due to noise or outliers
  - Poor accuracy for unseen samples
- Two approaches to avoid 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
  - Postpruning: Remove branches from a "fully grown" tree—get a sequence of progressively pruned trees
    - Use a set of data different from the training data to decide which is the "best pruned tree"

2018/10/16 28

## **Evaluating Classifier Accuracy**

- Holdout 保留法
  - Train on 2/3
  - Test on 1/3
- Cross validation: k-fold cross validation
  - Partition data set into k parts
  - Train on random (k-1) parts, test on 1 part
  - Repeat k times
  - Average accuracy

#### **Comments on Decision Tree Induction**

- Relatively faster learning speed (than other classification methods)
- Convertible to simple and easy to understand classification rules
- Comparable classification accuracy with other methods
- Comparably scalable to large database

# **Enhancements to Basic Decision Tree Induction**

- Allow for continuous-valued attributes
  - Dynamically define new discrete-valued attributes that partition the continuous attribute value into a discrete set of intervals
- Handle missing attribute values
  - Assign the most common value of the attribute
- Attribute construction
  - Create new attributes based on existing ones

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## **Bayesian Classification**

#### A statistical classifier

Perform *probabilistic prediction, i.e.,* predict class membership probabilities

#### Foundation

Based on Bayes' Theorem

#### Assumption

The effect of an attribute on a given class is independent of other attributes

#### Performance

A simple Bayesian classifier, *naïve Bayesian classifier*, has comparable performance with decision tree and selected neural network classifiers

## **Bayesian Theorem: Basics**

- Let X be a data sample, class label is unknown
- Let H be a hypothesis, e.g. X belongs to class C
- Classification is to determine P(H|X), the probability that the hypothesis holds given the observed data sample X
- $\blacksquare$  P(*H*), the initial probability
  - E.g., X will buy computer, regardless of age, income, ...
- $\blacksquare$  P(X): probability that sample data is observed
- Arr P(X|H), the probability of observing the sample X, given that the hypothesis holds
  - E.g., Given that X will buy computer, what is the prob. that X is 31..40?

## **Bayesian Theorem**

Given training data X, probability of a hypothesis H,
 P(H|X), follows the Bayes Theorem

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

- Predict X belongs to  $C_i$  iff the probability  $P(C_i|X)$  is the highest among all the  $P(C_k|X)$  for all the k classes
- Practical difficulty: require initial knowledge of many probabilities, significant computational cost

## Na we Bayesian Classifier

- Let D be a training set of tuples and their associated class labels, and each tuple is represented by an n-D attribute vector  $X = (x_1, x_2, ..., x_n)$
- Suppose there are m classes C<sub>1</sub>, C<sub>2</sub>, ..., C<sub>m</sub>
- Classification is to derive the maximum posteriori, i.e., the maximal P(C<sub>i</sub>|X)
- This can be derived from Bayes Theorem

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

Since P(X) is constant for all classes, only

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

needs to be maximized

P(C<sub>i</sub>) can be obtained from training data set s<sub>i</sub>/s

36

## Derivation of Na we Bayes Classifier

Assumption: attributes are conditionally independent (i.e., no dependence relation between attributes):

$$P(\mathbf{X} \mid C_i) = \prod_{k=1}^{n} P(x_k \mid C_i) = P(x_1 \mid C_i) \times P(x_2 \mid C_i) \times ... \times P(x_n \mid C_i)$$

- This greatly reduces the computation cost: Only counts the class distribution
- If  $A_k$  is categorical,  $P(x_k|C_i) = s_{ik}/s_i$ , count the distribution
- If A<sub>k</sub> is continuous-valued, P(x<sub>k</sub>|C<sub>i</sub>) is usually computed based on Gaussian distribution with a mean μ and standard deviation σ

$$g(x,\mu,\sigma) = \frac{1}{\sqrt{2\pi}\sigma} e^{-\frac{(x-\mu)^2}{2\sigma^2}}$$

and  $P(x_k|C_i)$  is

$$P(X_k|C_i) = g(x_k, \mu_{C_i}, \sigma_{C_i})$$

### **Exercise**

Predict what class does the data sample

X = (age <=30,
Income = medium,
Student = yes
Credit\_rating = Fair) belong

### Class:

to?

C1:buys\_computer = 'yes' C2:buys\_computer = 'no'

age	income	student	credit_rating	_com <sub> </sub>
<=30	high	no	fair	no
<=30	high	no	excellent	no
3140	high	no	fair	yes
>40	medium	no	fair	yes
>40	low	yes	fair	yes
>40	low	yes	excellent	no
3140	low	yes	excellent	yes
<=30	medium	no	fair	no
<=30	low	yes	fair	yes
>40	medium	yes	fair	yes
<=30	medium	yes	excellent	yes
3140	medium	no	excellent	yes
3140	high	yes	fair	yes
>40	medium	no	excellent	no

### **Solution**

- P( $C_i$ ): P(buys\_computer = "yes") = 9/14 = 0.643 P(buys\_computer = "no") = 5/14= 0.357
- Compute P(X|C<sub>i</sub>) for each class

```
P(age = "<=30" | buys_computer = "yes") = 2/9 = 0.222
P(age = "<= 30" | buys_computer = "no") = 3/5 = 0.6
P(income = "medium" | buys_computer = "yes") = 4/9 = 0.444
P(income = "medium" | buys_computer = "no") = 2/5 = 0.4
P(student = "yes" | buys_computer = "yes) = 6/9 = 0.667
P(student = "yes" | buys_computer = "no") = 1/5 = 0.2
P(credit_rating = "fair" | buys_computer = "yes") = 6/9 = 0.667
P(credit_rating = "fair" | buys_computer = "no") = 2/5 = 0.4
```

X = (age <= 30, income = medium, student = yes, credit\_rating = fair)</p>

```
P(X|C_i): P(X|buys\_computer = "yes") = 0.222 x 0.444 x 0.667 x 0.667 = 0.044 
 <math>P(X|buys\_computer = "no") = 0.6 x 0.4 x 0.2 x 0.4 = 0.019

P(X|C_i)*P(C_i): P(X|buys\_computer = "yes") * <math>P(buys\_computer = "yes") = 0.028

P(X|buys\_computer = "no") * <math>P(buys\_computer = "no") = 0.007

Therefore, X belongs to class ("buys computer = yes")
```

2018/10/16

39

## Na we Bayesian Classifier: Comments

### Advantages

- Easy to implement
- Good results obtained in most of the cases

### Disadvantages

- Assumption: class conditional independence, therefore loss of accuracy
- Practically, dependencies do exist among variables
  - E.g., hospitals: patients: Profile: age, family history, etc. Symptoms: fever, cough etc., Disease: lung cancer, diabetes, etc.
  - Dependencies among these cannot be modeled by Naïve Bayesian Classifier
- How to deal with these dependencies?
  - Bayesian Belief Networks

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2018/10/16 41

# Classification by Backpropagation

- Backpropagation: A neural network learning algorithm
- Started by psychologists and neurobiologists to develop and test computational analogues of neurons
- A neural network: A set of connected input/output units where each connection has a weight associated with it
- During the learning phase, the network learns by adjusting the weights so as to be able to predict the correct class label of the input tuples

## A Multi-Layer Feed-Forward Neural Network

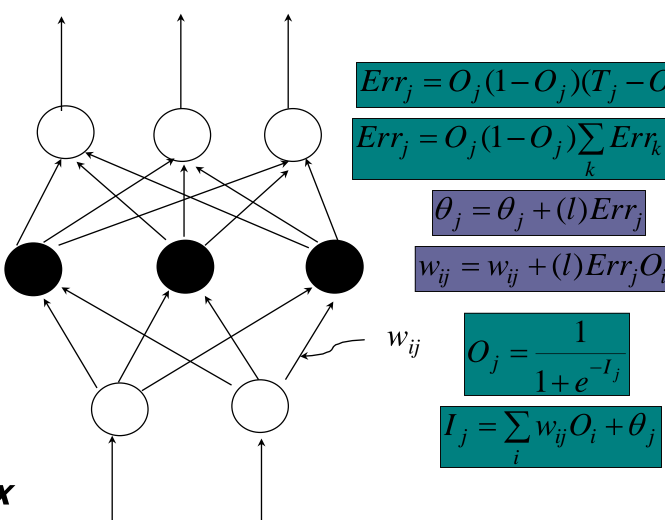


**Output layer** 

**Hidden layer** 

**Input layer** 

Input vector: X

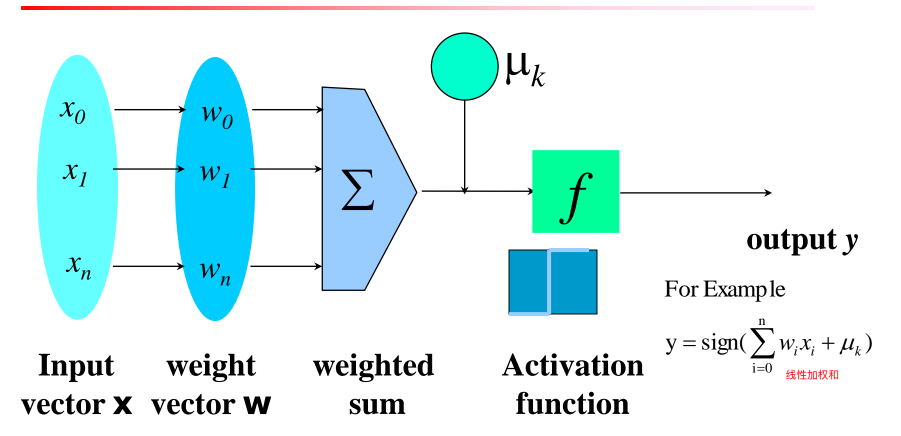


43

# **Defining a Network Topology**

- First decide the network topology: # of units in the input layer, # of hidden layers (if > 1), # of units in each hidden layer, and # of units in the output layer
- Normalizing the input values for each attribute measured in the training tuples to [0.0—1.0]
- For categorical attribute, one input unit per domain value, each initialized to 0
- Output, if for classification and more than two classes, one output unit per class is used
- Once a network has been trained and its accuracy is unacceptable, repeat the training process with a *different* network topology or a different set of initial weights

## A Neuron (= a perceptron)



45

The n-dimensional input vector x is mapped into variable y by means of the scalar product and a nonlinear function mapping

## **How A Multi-Layer Neural Network Works?**

- The **inputs** to the network correspond to the attributes measured for each training tuple
- Inputs are fed simultaneously into the units making up the input layer
- They are then weighted and fed simultaneously to a **hidden layer**
- The number of hidden layers is arbitrary, although usually only one
- The weighted outputs of the last hidden layer are input to units making up the **output layer**, which emits the network's prediction
- The network is **feed-forward** in that none of the weights cycles back to an input unit or to an output unit of a previous layer
- From a statistical point of view, networks perform **nonlinear** regression: Given enough hidden units and enough training samples, they can closely approximate any function

46

## **Backpropagation**

- Initialize weights and biases as random numbers
- Iteratively process a set of training tuples & compare the network's prediction with the actual known target value
- For each training tuple, the weights are modified to minimize the mean squared error between the network's prediction and the actual target value
- Modifications are made in the "backwards" direction: from the output layer, through each hidden layer down to the first hidden layer, hence "backpropagation"

2018/10/16 47

## **Backpropagation**

Algorithm: Backpropagation. Neural network learning for classification or prediction, using the backpropagation algorithm.

### Input:

- D, a data set consisting of the training tuples and their associated target values;
- l, the learning rate;
- · network, a multilayer feed-forward network.

Output: A trained neural network.

#### Method:

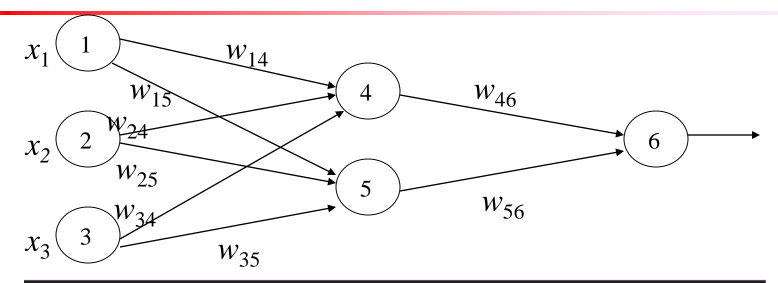
```
Initialize all weights and biases in network;
 (1)
        while terminating condition is not satisfied {
 (2)
 (3)
             for each training tuple X in D {
 (4)
                      // Propogate the inputs forward:
                      for each input layer unit j {
 (5)
                               O_i = I_i; // output of an input unit is its actual input value
 (6)
                      for each hidden or output layer unit j {
 (7)
                               I_i = \sum_i w_{ii}O_i + \theta_i; // compute the net input of unit j with respect to the previous layer, i
 (8)
                               O_j = \frac{1}{1 + e^{-I_j}}; } // compute the output of each unit j
 (9)
(10)
                      // Backpropogate the errors;
(11)
                      for each unit j in the output layer
                               Err_i = O_i(1 - O_i)(T_i - O_i); // compute the error
(12)
                      for each unit j in the hidden layers, from the last to the first hidden layer
(13)
                               Err_i = O_i(1 - O_i)\sum_k Err_k w_{ik}; // compute the error with respect to the next higher layer, k
(14)
                      for each weight wij in network {
(15)
                               \Delta w_{ij} = (l)Err_iO_i; // weight increment
(16)
                               w_{ij} = w_{ij} + \Delta w_{ij}; \} // weight update
(17)
                      for each bias \hat{\theta}_i in network {
(18)
                               \Delta\theta_{i} = (l)Err_{i}; // bias increment
(19)
                               \theta_i = \theta_i + \Delta \theta_i; } // bias update
(20)
                      } }
(21)
```

## **Backpropagation**

### Steps

- Initialize weights (to small random #s) and biases in the network
- Propagate the inputs forward (by applying activation function)
- Backpropagate the error (by updating weights and biases)
- Terminating condition (when error is very small, etc.)

## **Exercise**



$x_1$	$x_2$	$x_3$	w <sub>14</sub>	$w_{15}$	$w_{24}$	$w_{25}$	$w_{34}$	$w_{35}$	w <sub>46</sub>	$w_{56}$	$ heta_4^{^{ m Bias}}$	$\theta_5$	$\theta_6$
1	0	1	0.2	-0.3	0.4	0.1	-0.5	0.2	-0.3	-0.2	-0.4	0.2	0.1

Unit j	Net input, $I_j$	Output, $O_j$
4	0.2 + 0 - 0.5 - 0.4 = -0.7	$1/(1 + e^{0.7}) = 0.332$
5	-0.3 + 0 + 0.2 + 0.2 = 0.1	$1/(1 + e^{-0.1}) = 0.525$
6	(-0.3)(0.332) - (0.2)(0.525) + 0.1 = -0.105	$1/(1 + e^{0.105}) = 0.474$

# **Exercise**

Unit j	Err j
6	(0.474)(1 - 0.474)(1 - 0.474) = 0.1311
5	(0.474)(1 - 0.474)(1 - 0.474) = 0.1311 (0.525)(1 - 0.525)(0.1311)(-0.2) = -0.0065
4	(0.332)(1 - 0.332)(0.1311)(-0.3) = -0.0087

Weight or bias	New value
w <sub>46</sub>	-0.3 + (0.9)(0.1311)(0.332) = -0.261
w <sub>56</sub>	-0.2 + (0.9)(0.1311)(0.525) = -0.138
$w_{14}$	0.2 + (0.9)(-0.0087)(1) = 0.192
<i>w</i> <sub>15</sub>	-0.3 + (0.9)(-0.0065)(1) = -0.306
$w_{24}$	0.4 + (0.9)(-0.0087)(0) = 0.4
$w_{25}$	0.1 + (0.9)(-0.0065)(0) = 0.1
$w_{34}$	-0.5 + (0.9)(-0.0087)(1) = -0.508
$w_{35}$	0.2 + (0.9)(-0.0065)(1) = 0.194
$\theta_6$	0.1 + (0.9)(0.1311) = 0.218
$\theta_5$	0.2 + (0.9)(-0.0065) = 0.194
$\theta_4$	-0.4 + (0.9)(-0.0087) = -0.408

# **Backpropagation and Interpretability**

- Rule extraction from networks: network pruning
  - Simplify the network structure by removing weighted links that have the least effect on the trained network
  - The set of input and activation values are studied to derive rules describing the relationship between the input and hidden unit layers
- Sensitivity analysis: assess the impact that a given input variable has on a network output. The knowledge gained from this analysis can be represented in rules

### Neural Network as a Classifier

## Strength

- High tolerance to noisy data
- Well-suited for continuous-valued inputs and outputs
- Successful on a wide array of real-world data
- Techniques have recently been developed for the extraction of rules from trained neural networks

### Neural Network as a Classifier

### Weakness

- Long training time
- Require a number of parameters typically best determined empirically, e.g., the network topology or "structure"
- Poor interpretability: Difficult to interpret the symbolic meaning behind the learned weights and of "hidden units" in the network

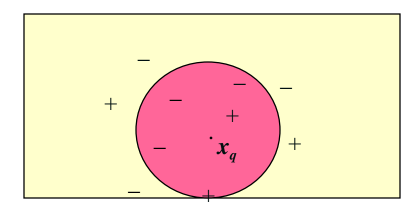
### **Classification and Prediction**

- What is classification?
  What is prediction?
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## The k-Nearest Neighbor Algorithm

- All instances correspond to points in the n-D space
- The nearest neighbor is defined in terms of Euclidean distance,  $dist(X_1, X_2)$
- Target function could be discrete- or real- valued
- For discrete-valued, k-NN returns the most common value among the k training examples nearest to  $X_q$



2018/10/16 56

### **Exercise**

1. Consider the one-dimensional data set. Please classify the data point x=5.0 according to its 1-, 3-, and 5-nearest neighbors (using majority vote).

X	0.5	3.0	4.5	4.6	4.9	5.2	5.3	5.5	7.0	9.5
У	-	-	+	+	+	-	-	+	-	-

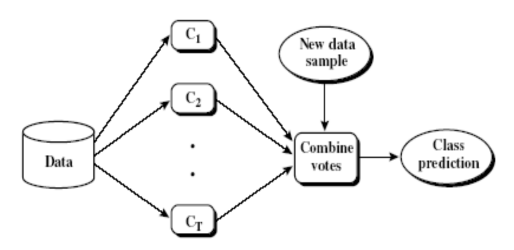
## Discussion on the k-NN Algorithm

- k-NN for real-valued prediction for a given unknown tuple
  - Returns the mean values of the k nearest neighbors
- Robust to noisy data by averaging k-nearest neighbors
- Distance between neighbors could be dominated by irrelevant attributes
  - To overcome it, eliminate irrelevant attributes
- Lazy-learner
  - Not build a classifier
  - Store all the training samples
  - High computational cost for each new tuple

2018/10/16 58

# **Ensemble Methods: Increasing the Accuracy**

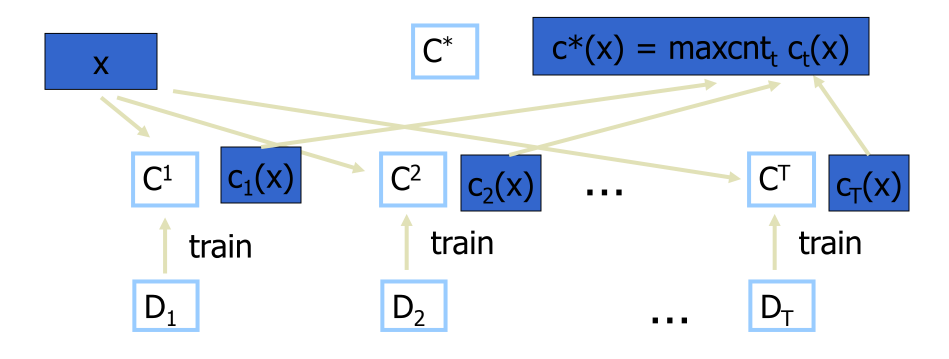
- Ensemble methods
  - Use a combination of models to increase accuracy
  - Combine a series of k learned models, M<sub>1</sub>, M<sub>2</sub>, ..., M<sub>k</sub>, with the aim of creating an improved model M\*
- Popular ensemble methods
  - Bagging
  - Boosting



## **Bagging: Boostrap Aggregation**

- Analogy: Diagnosis based on multiple doctors' majority vote
- Training
  - Given a set D of d tuples, at each iteration i, a training set D<sub>i</sub> is sampled with replacement from D
  - A classifier model M<sub>i</sub> is learned for each training set D<sub>i</sub>
- Classification: classify an unknown sample X
  - Each classifier M<sub>i</sub> returns its class prediction
  - The bagged classifier M\* counts the votes and assigns the class with the most votes to X

# **Bagging: Boostrap Aggregation**



## **Bagging: Boostrap Aggregation**

- Prediction: can be applied to the prediction of continuous values by taking the average value of each prediction for a given test tuple
- Accuracy
  - Often significant better than a single classifier derived from D
  - For noise data: not considerably worse, more robust
  - Proved improved accuracy in prediction

### **Exercise**

1. Following is a data set to construct a bagging classifier.

X	0.1	0.2	0.3	0.4	0.5	0.6	0.7	8.0	0.9	1
У	1	1	1	-1	-1	-1	-1	1	1	1

Examples chosen for training in each round are shown below:

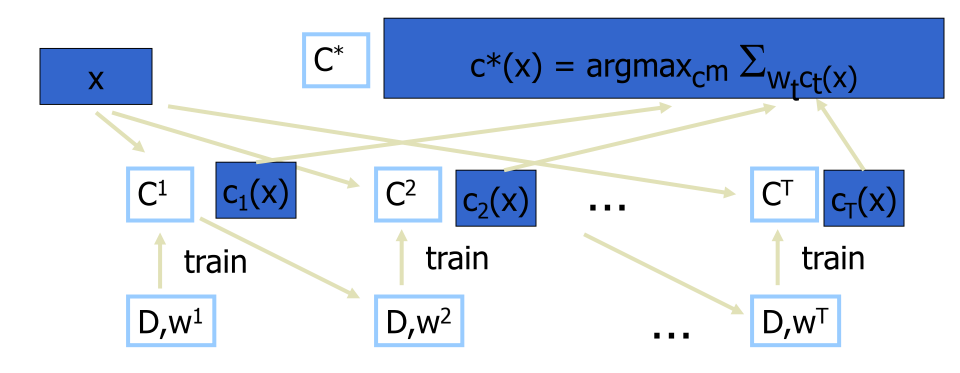
X	0.1	0.2	0.2	0.3	0.4	0.4	0.5	0.6	0.9	0.9	
У	1	1	1	1	-1	-1	-1	-1	1	1 0	$.35 < x \le 0.75 = y = -1$
Х	0.1	0.2	0.3	0.5	0.5	8.0	0.9	1	1	1	$0.4 < x \le 0.65 => y = -1$
У	1	1	1	-1	-1	1	1	1	1	1	,
X	0.1	0.2	0.3	0.4	0.4	0.5	0.7	0.7	8.0	0.90	$.35 < x \le 0.75 => y=-1$
У	1	1	1	-1	-1	-1	-1	-1	1	1	

Please predict the class label for the record x=0.38.

# **Boosting**

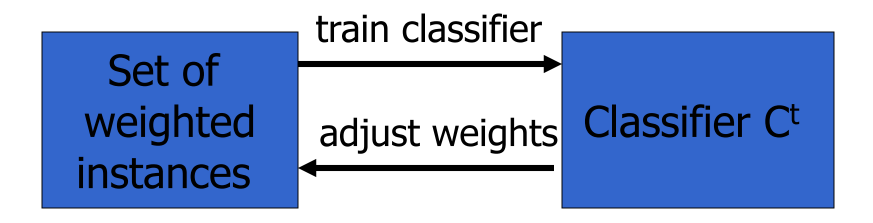
- Analogy: Consult several doctors, based on a combination of weighted diagnoses — weight assigned based on the previous diagnosis accuracy
- How boosting works?
  - After a classifier M<sub>i</sub> is learned, the weights are updated to allow the subsequent classifier, M<sub>i+1</sub>, pay more attention to the training tuples that were misclassified by M<sub>i</sub>
  - A series of k classifiers is iteratively learned
  - The final M\* combines the votes of each individual classifier, where the weight of each classifier's vote is a function of its accuracy

# **Boosting**



# **Boosting**

- The boosting algorithm can be extended for the prediction of continuous values
- Comparing with bagging: boosting tends to achieve greater accuracy, but it also risks overfitting the model to misclassified data



2018/10/16 66

# Bagging vs. Boosting

### Model training:

- Bagging: random sampling, independent classifiers
- Boosting: subsequent classifier, M<sub>i+1</sub>, pay more attention to the training tuples that were misclassified by M<sub>i</sub>

### Model usage:

- Bagging: equal weight
- Boosting: different weight assigned

## **Ensemble Methods**

- Text mining
- Video pattern recognition
- Audio pattern recognition

### Classification and Prediction

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### What Is Prediction?

- (Numerical) prediction is similar to classification
  - construct a model
  - use model to predict continuous or ordered value for a given input
- Prediction is different from classification
  - Classification refers to predict categorical class label
  - Prediction models continuous-valued functions
- Major method for prediction: regression
  - model the relationship between one or more independent or predictor variables and a dependent or response variable
- Regression analysis
  - Linear and multiple regression
  - Non-linear regression
- Other regression methods: generalized linear model, Poisson regression, log-linear models, regression trees, logistic regression

# **Linear Regression**

Linear regression: a response variable y and a single predictor variable x

$$y = w_0 + w_1 x$$

where w<sub>0</sub> (intercept) and w<sub>1</sub> (slope) are regression coefficient

Method of least squares: estimates the best-fitting straight line  $\sum_{i=1}^{|D|} (x_i - \bar{x})(y_i - \bar{y})$ 

 $w_{1} = \frac{\sum_{i=1}^{|D|} (x_{i} - \bar{x})(y_{i} - \bar{y})}{\sum_{i=1}^{|D|} (x_{i} - \bar{x})^{2}} \qquad w_{0} = \bar{y} - w_{1}\bar{x}$ 

- Multiple linear regression: more than one predictor variable
  - Training data is of the form (X<sub>1</sub>, y<sub>1</sub>), (X<sub>2</sub>, y<sub>2</sub>),..., (X<sub>|D|</sub>, y<sub>|D|</sub>)
  - Ex. For 2-D data, we may have:  $y = w_0 + w_1 x_1 + w_2 x_2$
  - Solvable by extension of least square method or using SAS, S-Plus, R, Matlab
- Many nonlinear functions can be transformed into the above 71

## **Nonlinear Regression**

 A polynomial regression model can be transformed into linear regression model. For example,

$$y = w_0 + w_1 x + w_2 x^2 + w_3 x^3$$
convertible to linear with new variables:  $x_2 = x^2$ ,  $x_3 = x^3$ 

$$y = w_0 + w_1 x + w_2 x_2 + w_3 x_3$$

- Some models are intractable nonlinear (e.g., sum of exponential terms)
  - possible to obtain least square estimates through extensive calculation on more complex formulae

## **Other Regression-Based Models**

### Generalized linear model:

- Foundation on which linear regression can be applied to modeling categorical response variables
- Logistic regression: models the prob. of some event occurring as a linear function of a set of predictor variables

$$Log(p/1-p) = W_0 + W_1 X_1 + W_2 X_2 + ... + W_n X_n$$
, p is probability Y=1

- Poisson regression: models the data that exhibit a Poisson distribution
- Log-linear models: (for categorical data)
  - Approximate discrete multidimensional prob. distributions
  - Also useful for data compression and smoothing

$$log(y) = W_0 + W_1 X + W_2 X_2 + ... + W_n X_n$$

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## Classifier Accuracy Measures

- Accuracy of a classifier M, acc(M): percentage of test set tuples that are correctly classified by the model M
  - Given m classes, CM<sub>i,j</sub>, an entry in a confusion matrix, indicates # of tuples in class i that are labeled by the classifier as class j
  - Accuracy = (t-pos + t-neg)/ (pos + neg)
  - Error rate (misclassification rate) of M = 1 acc(M)

### Predicted class

Actual class

	C <sub>1</sub>	$C_2$	Total
$C_1$	True positive	False negative	pos
$C_2$	False positive	True negative	neg
Total	t-pos+f-pos	t-neg+f-neg	pos+neg

## Classifier Accuracy Measures

Alternative accuracy measures

```
sensitivity = t-pos/pos /* true positive recognition rate */
specificity = t-neg/neg /* true negative recognition rate */
precision = t-pos/(t-pos + f-pos)
accuracy = sensitivity * pos/(pos + neg) + specificity * neg/(pos + neg)
= (t-pos + t-neg)/ (pos + neg)
```

### **Exercise**

1. Please compute the sensitivity, specificity, precision and accuracy of the classifier.

classes	buy_computer = yes	buy_computer = no	total	recognition(%)
buy_computer = yes	6954	46	7000	99.34
buy_computer = no	412	2588	3000	86.27
total	7366	2634	10000	95.42

### Classification and Prediction

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# Summary (I)

- Classification and prediction are two forms of data analysis that can be used to extract models describing important data classes or to predict future data trends.
- Effective and scalable methods have been developed for decision trees induction, Naive Bayesian classification, Backpropagation, knearest neighbor classifiers.

## Summary (II)

- Linear, nonlinear, and generalized linear models of regression can be used for prediction. Many nonlinear problems can be converted to linear problems by performing transformations on the predictor variables.
- k-fold cross-validation is a recommended method for accuracy estimation. ####
- Bagging and boosting can be used to increase overall accuracy by learning and combining a series of individual models.
- No single method has been found to be superior over all others for all data sets.
- Issues such as accuracy, training time, robustness, interpretability, and scalability must be considered.