Supervised Learning: Classification

Sept. 24, 2018

Classification: Basic concepts

- Classification: Basic Concepts
- Decision Tree Induction
- Bayes Classification Methods
- Model Evaluation and Selection
- Techniques to Improve Classification
 Accuracy: Ensemble Methods
- Summary

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, observations, etc. with the aim of establishing the existence of classes or clusters in the data

Prediction Problems: Classification vs. Numeric Prediction

- Classification
 - predicts categorical class labels (discrete or nominal)
 - classifies data (constructs a model) based on the training set and the values (class labels) in a classifying attribute and uses it in classifying new data
- Numeric Prediction
 - models continuous-valued functions, i.e., predicts unknown or missing values
- Typical applications
 - Credit/loan approval:
 - Medical diagnosis: if a tumor is cancerous or benign
 - Fraud detection: if a transaction is fraudulent
 - Web page categorization: which category it is

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
- 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)
 - If the accuracy is acceptable, use the model to classify new data
- Note: If *the test set* is used to select models, it is called validation (test) set

Process (1): Model Construction



Process (2): Using the Model in Prediction



Chapter 8. Classification: Basic Concepts

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Decision Tree Induction: An Example

- Training data set: Buys_computer •
- The data set follows an example • of Quinlan's ID3 (Playing Tennis)
- **Resulting tree:**



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

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)
- 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
 - There are no samples left

Brief Review of Entropy

- Entropy (Information Theory)
 - A measure of uncertainty associated with a random variable
 - Calculation: For a discrete random variable Y taking m distinct values {y₁, ..., y_m},
 - $H(Y) = -\sum_{i=1}^{m} p_i \log(p_i)$, where $p_i = P(Y = y_i)$
 - Interpretation:
 - Higher entropy => higher uncertainty
 - Lower entropy => lower uncertainty
- Conditional Entropy

• $H(Y|X) = \sum_{x} p(x)H(Y|X = x)$



Attribute Selection Measure: Information Gain (ID3/C4.5)

- Select the attribute with the highest information gain
- Let p_i be the probability that an arbitrary tuple in D belongs to class C_i , estimated by $|C_{i, D}|/|D|$
- Expected information (entropy) needed to classify a tuple in D: $Info(D) = -\sum_{i=1}^{m} p_i \log_2(p_i)$
- Information needed (after using A to split D into v partitions) to classify D:

$$Info_A(D) = \sum_{j=1}^{\nu} \frac{|D_j|}{|D|} \times Info(D_j)$$

• Information gained by branching on attribute A

 $Gain(A) = Info(D) - Info_A(D)$

Attribute Selection: Information Gain

- Class P: buys_computer = "yes"
- Class N: buys_computer = "no"

$$Info(D) = I(9,5) = -\frac{9}{14}\log_2(\frac{9}{14}) - \frac{5}{14}\log_2(\frac{5}{14}) = 0.940$$

age	p _i	n _i	l(p _i , n _i)
<=30	2	3	0.971
3140	4	0	0
>40	3	2	0.971

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

 $\frac{5}{14}I(2,3)$ means "age <=30" has 5 out of 14 samples, with 2 yes' es and 3 no's. Hence,

 $Gain(age) = Info(D) - Info_{age}(D) = 0.246$

Similarly,

Gain(income) = 0.029Gain(student) = 0.151 $Gain(credit_rating) = 0.048$

Computing Information-Gain for Continuous-Valued Attributes

- Let attribute A be a continuous-valued attribute
- Must determine the *best split point* for A
 - Sort the value A in increasing order
 - Typically, the midpoint between each pair of adjacent values is considered as a possible *split point*
 - $(a_i+a_{i+1})/2$ is the midpoint between the values of a_i and a_{i+1}
 - The point with the *minimum expected information* requirement for A is selected as the split-point for A
- Split:
 - D1 is the set of tuples in D satisfying A ≤ split-point, and D2 is the set of tuples in D satisfying A > split-point

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)

$$SplitInfo_{A}(D) = -\sum_{j=1}^{\nu} \left(\frac{|D_{j}|}{|D|} \right) \times \log_{2} \left(\frac{|D_{j}|}{|D|} \right)$$

– GainRatio(A) = Gain(A)/SplitInfo(A)

• Ex.

$$SplitInfo_{income}(D) = -\frac{4}{14} \times \log_2\left(\frac{4}{14}\right) - \frac{6}{14} \times \log_2\left(\frac{6}{14}\right) - \frac{4}{14} \times \log_2\left(\frac{4}{14}\right) = 1.557$$

- gain_ratio(income) = 0.029/1.557 = 0.019

• The attribute with the maximum gain ratio is selected as the splitting attribute

Gini Index (CART, IBM IntelligentMiner)

 If a data set D contains examples from n classes, gini index, gini(D) is defined as

$$gini(D) = 1 - \sum_{j=1}^{n} p_j^2$$

where p_i is the relative frequency of class *j* in *D*

If a data set D is split on A into two subsets D₁ and D₂, the gini index gini(D) is defined as

$$gini_{A}(D) = \frac{|D_{1}|}{|D|}gini(D_{1}) + \frac{|D_{2}|}{|D|}gini(D_{2})$$

• Reduction in Impurity:

$$\Delta gini(A) = gini(D) - gini_A(D)$$

The attribute provides the smallest gini_{split}(D) (or the largest reduction in impurity) is chosen to split the node (need to enumerate all the possible splitting points for each attribute)

Computation of Gini Index

Ex. D has 9 tuples in buys_computer = "yes" and 5 in "no"

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

 $=\frac{10}{14}\left(1-\left(\frac{7}{10}\right)^2-\left(\frac{3}{10}\right)^2\right)+\frac{4}{14}\left(1-\left(\frac{2}{4}\right)^2-\left(\frac{2}{4}\right)^2\right)$

 $= Gini_{income \in \{high\}}(D).$

• Suppose the attribute income partitions D into 10 in D₁: {low, medium} and 4 in D₂ $gini_{income \in \{low, medium\}}(D) = \left(\frac{10}{14}\right)Gini(D_1) + \left(\frac{4}{14}\right)Gini(D_2)$

Gini_{low,high} is 0.458; Gini_{medium,high} is 0.450. Thus, split on the {low,medium} (and {high}) since it has the lowest Gini index

- All attributes are assumed continuous-valued
- May need other tools, e.g., clustering, to get the possible split values
- Can be modified for categorical attributes

Comparing Attribute Selection Measures

- The three measures, in general, return good results but
 - Information gain:
 - biased towards multivalued attributes
 - Gain ratio:
 - tends to prefer unbalanced splits in which one partition is much smaller than the others
 - Gini index:
 - biased to multivalued attributes
 - has difficulty when # of classes is large
 - tends to favor tests that result in equal-sized partitions and purity in both partitions

Other Attribute Selection Measures

- <u>CHAID</u>: a popular decision tree algorithm, measure based on χ^2 test for independence
- <u>C-SEP</u>: performs better than info. gain and gini index in certain cases
- <u>G-statistic</u>: has a close approximation to χ^2 distribution
- <u>MDL (Minimal Description Length) principle</u> (i.e., the simplest solution is preferred):
 - The best tree as the one that requires the fewest # of bits to both (1) encode the tree, and (2) encode the exceptions to the tree
- Multivariate splits (partition based on multiple variable combinations)
 - <u>CART</u>: finds multivariate splits based on a linear comb. of attrs.
- Which attribute selection measure is the best?
 - Most give good results, none is significantly superior than others

Overfitting and Tree Pruning

- <u>Overfitting</u>: 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
 - <u>Prepruning</u>: *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
 - <u>Postpruning</u>: *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"

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
 - Assign probability to each of the possible values
- Attribute construction
 - Create new attributes based on existing ones that are sparsely represented
 - This reduces fragmentation, repetition, and replication

Classification in Large Databases

- Classification—a classical problem extensively studied by statisticians and machine learning researchers
- Scalability: Classifying data sets with millions of examples and hundreds of attributes with reasonable speed
- Why is decision tree induction popular?
 - relatively faster learning speed (than other classification methods)
 - convertible to simple and easy to understand classification rules
 - can use SQL queries for accessing databases
 - comparable classification accuracy with other methods

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Bayesian Classification: Why?

- <u>A statistical classifier</u>: performs *probabilistic prediction, i.e.,* predicts class membership probabilities
- <u>Foundation</u>: Based on Bayes' Theorem.
- <u>Performance</u>: A simple Bayesian classifier, *naïve Bayesian classifier*, has comparable performance with decision tree and selected neural network classifiers
- <u>Incremental</u>: Each training example can incrementally increase/ decrease the probability that a hypothesis is correct — prior knowledge can be combined with observed data
- <u>Standard</u>: Even when Bayesian methods are computationally intractable, they can provide a standard of optimal decision making against which other methods can be measured

Bayes' Theorem: Basics

- Total probability Theorem: $P(B) = \sum_{i=1}^{M} P(B|A_i)P(A_i)$
- Bayes' Theorem: $P(H|\mathbf{X}) = \frac{P(\mathbf{X}|H)P(H)}{P(\mathbf{X})} = P(\mathbf{X}|H) \times P(H)/P(\mathbf{X})$
 - Let **X** be a data sample ("evidence"): class label is unknown
 - Let H be a hypothesis that X belongs to class C
 - Classification is to determine P(H|X), (i.e., *posteriori probability):* the probability that the hypothesis holds given the observed data sample X
 - P(H) (*prior probability*): the initial probability
 - E.g., X will buy computer, regardless of age, income, ...
 - P(X): probability that sample data is observed
 - P(X|H) (likelihood): the probability of observing the sample X, given that the hypothesis holds
 - E.g., Given that **X** will buy computer, the prob. that X is 31..40, medium income

Prediction Based on Bayes' Theorem

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

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

Informally, this can be viewed as

posteriori = likelihood x prior/evidence

- Predicts **X** belongs to C_i iff the probability $P(C_i | \mathbf{X})$ is the highest among all the $P(C_k | \mathbf{X})$ for all the *k* classes
- Practical difficulty: It requires initial knowledge of many probabilities, involving significant computational cost

Classification Is to Derive the Maximum Posteriori

- 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 = (x₁, x₂, ..., x_n)
- Suppose there are *m* classes C₁, C₂, ..., C_m.
- Classification is to derive the maximum posteriori, i.e., the maximal P(C_i | 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

nee

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

ds to be maximized

Naïve Bayes Classifier

• A simplified assumption: attributes are conditionally independent (i.e., no dependence relation between attributes):

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

- This greatly reduces the computation cost: Only counts the class distribution
- If A_k is categorical, P(x_k|C_i) is the # of tuples in C_i having value x_k for A_k divided by |C_{i, D}| (# of tuples of C_i in D)
- If A_k is continous-valued, $P(x_k | C_i)$ 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 $P(\mathbf{x}_k | \mathbf{C}_i)$ is $P(\mathbf{X} | C_i) = g(x_k, \mu_{C_i}, \sigma_{C_i})$

Naïve Bayes Classifier: Training Dataset

Class:

C1:buys_computer = 'yes' C2:buys_computer = 'no'

Data to be classified: X = (age <=30, Income = medium, Student = yes Credit_rating = Fair)

age	income	student	credit_rating	com
<=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
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<=30	medium	yes	excellent	yes
3140	medium	no	excellent	yes
3140	high	yes	fair	yes
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Naïve Bayes Classifier: An Example

- P(C_i): P(buys_computer = "yes") = 9/14 = 0.643
 P(buys_computer = "no") = 5/14 = 0.357
- Compute P(X|C_i) 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 = 0.444P(income = "medium" | buys_computer = "no") = 2/5 = 0.4P(student = "yes" | buys_computer = "yes) = 6/9 = 0.667P(student = "yes" | buys_computer = "no") = 1/5 = 0.2P(credit_rating = "fair" | buys_computer = "yes") = 6/9 = 0.667P(credit_rating = "fair" | buys_computer = "no") = 2/5 = 0.4

X = (age <= 30, income = medium, student = yes, credit_rating = fair)
 P(X|C_i): P(X|buys_computer = "yes") = 0.222 x 0.444 x 0.667 x 0.667 = 0.044

 $P(X|buys_computer = "no") = 0.6 \times 0.4 \times 0.2 \times 0.4 = 0.019$

P(X|C_i)*P(C_i): P(X|buys_computer = "yes") * P(buys_computer = "yes") = 0.028 P(X|buys_computer = "no") * P(buys_computer = "no") = 0.007 Therefore, X belongs to class ("buys_computer = yes")

age	income	student	credit_rating	_com
<=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
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Avoiding the Zero-Probability Problem

• Naïve Bayesian prediction requires each conditional prob. be **non-zero**. Otherwise, the predicted prob. will be zero

$$P(X \mid C_i) = \prod_{k=1}^{n} P(x_k \mid C_i)$$

- Ex. Suppose a dataset with 1000 tuples, income=low (0), income= medium (990), and income = high (10)
- Use Laplacian correction (or Laplacian estimator)
 - Adding 1 to each case

Prob(income = low) = 1/1003

Prob(income = medium) = 991/1003

Prob(income = high) = 11/1003

 The "corrected" prob. estimates are close to their "uncorrected" counterparts

Naïve Bayes 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 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 Bayes Classifier

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Model Evaluation and Selection

- Evaluation metrics: How can we measure accuracy? Other metrics to consider?
- Use validation test set of class-labeled tuples instead of training set when assessing accuracy
- Methods for estimating a classifier's accuracy:
 - Holdout method, random subsampling
 - Cross-validation
 - Bootstrap
- Comparing classifiers:
 - Confidence intervals
 - Cost-benefit analysis and ROC Curves

Classifier Evaluation Metrics: Confusion Matrix

Confusion Matrix:

Actual class\Predicted class	C ₁	¬ C ₁	
C ₁	True Positives (TP)	False Negatives (FN)	
¬ C ₁	False Positives (FP)	True Negatives (TN)	

Example of Confusion Matrix:

Actual class\Predicted	buy_computer	buy_computer	Total
Class	= yes	= 10	
buy_computer = yes	6954	46	7000
buy_computer = no	412	2588	3000
Total	7366	2634	10000

- Given *m* classes, an entry, *CM*_{i,j} in a confusion matrix indicates # of tuples in class *i* that were labeled by the classifier as class *j*
- May have extra rows/columns to provide totals

Classifier Evaluation Metrics: Accuracy, Error Rate, Sensitivity and Specificity

A\P	С	−C	
С	ТР	FN	Ρ
¬C	FP	ΤN	Ν
	Ρ'	N'	All

• Classifier Accuracy, or recognition rate: percentage of test set tuples that are correctly classified

Accuracy = (TP + TN)/All

Error rate: 1 – accuracy, or
 Error rate = (FP + FN)/All

- **Class Imbalance Problem**:
 - One class may be *rare*, e.g. fraud, or HIV-positive
 - Significant *majority of the negative class* and minority of the positive class
 - Sensitivity: True Positive recognition rate

Sensitivity = TP/P

- Specificity: True Negative recognition rate
 - Specificity = TN/N

Classifier Evaluation Metrics: Precision and Recall, and F-measures

- **Precision**: exactness what % of tuples that the classifier labeled as positive are actually positive $precision = \frac{TP}{TP + FP}$
- **Recall:** completeness what % of positive tuples did the classifier label as positive? $= \frac{TP}{TP + FN}$ recall
- Perfect score is 1.0
- Inverse relationship between precision & recall
- F measure (F₁ or F-score): harmonic mean of precision and recall,
- F_{β} : weighted measure of precision and recall <u>precision + recall</u> assigns ß times as much weight to recall as to precision

$$F_{\beta} = \frac{(1+\beta^2) \times precision \times recall}{\beta^2 \times precision + recall}$$

Classifier Evaluation Metrics: Example

Actual Class\Predicted class	cancer = yes	cancer = no	Total	Recognition(%)
cancer = yes	90	210	300	30.00 (sensitivity
cancer = no	140	9560	9700	98.56 (specificity)
Total	230	9770	10000	96.40 (<i>accuracy</i>)

– Precision = 90/230 = 39.13%

Recall = 90/300 = 30.00%

Evaluating Classifier Accuracy: Holdout & Cross-Validation Methods

Holdout method

- Given data is randomly partitioned into two independent sets
 - Training set (e.g., 2/3) for model construction
 - Test set (e.g., 1/3) for accuracy estimation
- <u>Random sampling</u>: a variation of holdout
 - Repeat holdout k times, accuracy = avg. of the accuracies obtained
- **Cross-validation** (*k*-fold, where k = 10 is most popular)
 - Randomly partition the data into k mutually exclusive subsets, each approximately equal size
 - At *i*-th iteration, use D_i as test set and others as training set
 - <u>Leave-one-out</u>: k folds where k = # of tuples, for small sized data
 - <u>*Stratified cross-validation</u>^{*}: folds are stratified so that class dist. in each fold is approx. the same as that in the initial data

Issues Affecting Model Selection

- Accuracy
 - classifier accuracy: predicting class label
- 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
- Interpretability
 - 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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Ensemble Methods: Increasing the Accuracy



- Ensemble methods
 - Use a combination of models to increase accuracy
 - Combine a series of k learned models, M_1 , M_2 , ..., M_k , with the aim of creating an improved model M^*
- Popular ensemble methods
 - Bagging: averaging the prediction over a collection of classifiers
 - Boosting: weighted vote with a collection of classifiers
 - Ensemble: combining a set of heterogeneous classifiers

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_i of *d* tuples is sampled with replacement from D (i.e., bootstrap)
 - A classifier model M_i is learned for each training set D_i
- Classification: classify an unknown sample **X**
 - Each classifier M_i returns its class prediction
 - The bagged classifier M* counts the votes and assigns the class with the most votes to X
- 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 significantly better than a single classifier derived from D
 - For noise data: not considerably worse, more robust
 - Proved improved accuracy in prediction

Boosting

- Analogy: Consult several doctors, based on a combination of weighted diagnoses—weight assigned based on the previous diagnosis accuracy
- How boosting works?
 - Weights are assigned to each training tuple
 - A series of k classifiers is iteratively learned
 - After a classifier M_i is learned, the weights are updated to allow the subsequent classifier, M_{i+1}, to pay more attention to the training tuples that were misclassified by M_i
 - 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 algorithm can be extended for numeric prediction
- Comparing with bagging: Boosting tends to have greater accuracy, but it also risks overfitting the model to misclassified data

Adaboost (Freund and Schapire, 1997)

- Given a set of d class-labeled tuples, $(X_1, y_1), ..., (X_d, y_d)$
- Initially, all the weights of tuples are set the same (1/d)
- Generate k classifiers in k rounds. At round i,
 - Tuples from D are sampled (with replacement) to form a training set
 D_i of the same size
 - Each tuple's chance of being selected is based on its weight
 - A classification model M_i is derived from D_i
 - Its error rate is calculated using D_i as a test set
 - If a tuple is misclassified, its weight is increased, o.w. it is decreased

 $\log \frac{1 - error(M_i)}{error(M_i)}$

• Error rate: err(**X**_j) is the misclassification error of tuple **X**_j. Classifier M_i error rate is the sum of the weights of the misclassified tuples:

$$error(M_i) = \sum_{j}^{d} w_j \times err(\mathbf{X_j})$$

• The weight of classifier M_i's vote is

Random Forest (Breiman 2001)

- Random Forest:
 - Each classifier in the ensemble is a *decision tree* classifier and is generated using a random selection of attributes at each node to determine the split
 - During classification, each tree votes and the most popular class is returned
- Two Methods to construct Random Forest:
 - Forest-RI (*random input selection*): Randomly select, at each node, F attributes as candidates for the split at the node. The CART methodology is used to grow the trees to maximum size
 - Forest-RC (random linear combinations): Creates new attributes (or features) that are a linear combination of the existing attributes (reduces the correlation between individual classifiers)
- Comparable in accuracy to Adaboost, but more robust to errors and outliers
- Insensitive to the number of attributes selected for consideration at each split, and faster than bagging or boosting

Summary (I)

- Classification is a form of data analysis that extracts models describing important data classes.
- Effective and scalable methods have been developed for decision tree induction, Naive Bayesian classification, rule-based classification, and many other classification methods.
- Evaluation metrics include: accuracy, sensitivity, specificity, precision, recall, F measure, and F_β measure.
- Stratified k-fold cross-validation is recommended for accuracy estimation. Bagging and boosting can be used to increase overall accuracy by learning and combining a series of individual models.

Summary (II)

- There have been numerous comparisons of the different classification methods; the matter remains a research topic
- No single method has been found to be superior over all others for all data sets
- Issues such as accuracy, training time, robustness, scalability, and interpretability must be considered and can involve tradeoffs, further complicating the quest for an overall superior method
- References: http://hanj.cs.illinois.edu/