



# Data Mining

Practical Machine Learning Tools and Techniques

Slides for Chapter 7 of *Data Mining* by I. H. Witten and E. Frank

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## Engineering the input and output

- Attribute selection
  - Scheme-independent, scheme-specific
- Attribute discretization
  - Unsupervised, supervised, error- vs entropy-based, converse of discretization
- Data transformations
  - Principal component analysis, random projections, text, time series
- Dirty data
  - Data cleansing, robust regression, anomaly detection
- Meta-learning
  - Bagging (with costs), randomization, boosting, additive (logistic) regression, option trees, logistic model trees, stacking, ECOCs
- Using unlabeled data
  - Clustering for classification, co-training, EM and co-training

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## Just apply a learner? NO!

- Scheme/parameter selection  
*treat selection process as part of the learning process*
- Modifying the input:
  - Data engineering to make learning possible or easier
- Modifying the output
  - Combining models to improve performance

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## Attribute selection

- Adding a random (i.e. irrelevant) attribute can significantly degrade C4.5's performance
  - Problem: attribute selection based on smaller and smaller amounts of data
- IBL very susceptible to irrelevant attributes
  - Number of training instances required increases exponentially with number of irrelevant attributes
- Naïve Bayes doesn't have this problem
- *Relevant* attributes can also be harmful

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## Scheme-independent attribute selection

- *Filter* approach: assess based on general characteristics of the data
- One method: find smallest subset of attributes that separates data
- Another method: use different learning scheme
  - e.g. use attributes selected by C4.5 and 1R, or coefficients of linear model, possibly applied recursively (*recursive feature elimination*)
- IBL-based attribute weighting techniques:
  - can't find redundant attributes (but fix has been suggested)
- Correlation-based Feature Selection (CFS):
  - correlation between attributes measured by *symmetric uncertainty*:  

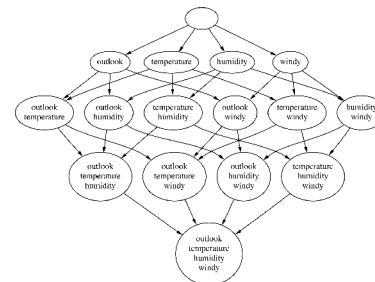
$$U(A, B) = 2 \frac{H(A) + H(B) - H(A, B)}{H(A) + H(B)} \in [0, 1]$$
  - goodness of subset of attributes measured by (breaking ties in favor of smaller subsets):  

$$\sum_j U(A_j, C) / \sqrt{(\sum_j \sum_j U(A_j, A_j))}$$

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## Attribute subsets for weather data



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## Searching attribute space

- Number of attribute subsets is exponential in number of attributes
- Common greedy approaches:
  - *forward selection*
  - *backward elimination*
- More sophisticated strategies:
  - *Bidirectional search*
  - *Best-first search*: can find optimum solution
  - *Beam search*: approximation to best-first search
  - *Genetic algorithms*

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## Scheme-specific selection

- *Wrapper* approach to attribute selection
- Implement “wrapper” around learning scheme
  - Evaluation criterion: cross-validation performance
- Time consuming
  - greedy approach,  $k$  attributes  $\Rightarrow k^2 \times \text{time}$
  - prior ranking of attributes  $\Rightarrow$  linear in  $k$
- Can use significance test to stop cross-validation for subset early if it is unlikely to “win” (*race search*)
  - can be used with forward, backward selection, prior ranking, or special-purpose *schemata search*
- Learning decision tables: scheme-specific attribute selection essential
- Efficient for decision tables and Naïve Bayes

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## Attribute discretization

- Avoids normality assumption in Naïve Bayes and clustering
- 1R: uses simple discretization scheme
- C4.5 performs *local* discretization
- *Global* discretization can be advantageous because it's based on more data
- Apply learner to
  - $k$ -valued discretized attribute *or* to
  - $k-1$  binary attributes that code the cut points

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## Discretization: unsupervised

- Determine intervals without knowing class labels
- When clustering, the only possible way!
- Two strategies:
  - *Equal-interval binning*
  - *Equal-frequency binning* (also called *histogram equalization*)
- Normally inferior to supervised schemes in classification tasks
- But equal-frequency binning works well with naïve Bayes if number of intervals is set to square root of size of dataset (*proportional k-interval discretization*)

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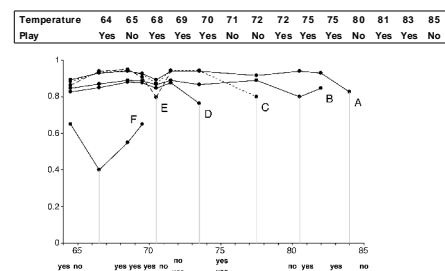
## Discretization: supervised

- *Entropy-based* method
- Build a decision tree with pre-pruning on the attribute being discretized
  - Use entropy as splitting criterion
  - Use minimum description length principle as stopping criterion
- Works well: the state of the art
- To apply min description length principle:
  - The “theory” is
    - the splitting point ( $\log_2[N-1]$  bits)
    - plus class distribution in each subset
  - Compare description lengths before/after adding splitting point

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## Example: temperature attribute



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## Formula for MDLP

- $N$  instances
  - Original set:  $k$  classes, entropy  $E$
  - First subset:  $k_1$  classes, entropy  $E_1$
  - Second subset:  $k_2$  classes, entropy  $E_2$

$$gain > \frac{\log_2(N-1)}{N} + \frac{\log_2(3^L-2) - kE + k_1E_1 + k_2E_2}{N}$$

- Results in *no* discretization intervals for temperature attribute

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## Supervised discretization: other methods

- Can replace top-down procedure by bottom-up method
- Can replace MDLP by chi-squared test
- Can use dynamic programming to find optimum  $k$ -way split for given additive criterion
  - Requires time quadratic in the number of instances
  - But can be done in linear time if error rate is used instead of entropy

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## Error-based vs. entropy-based

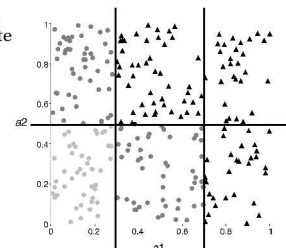
- Question: could the best discretization ever have two adjacent intervals with the same class?
- Wrong answer: No. For if so,
  - Collapse the two
  - Free up an interval
  - Use it somewhere else
  - (*This is what error-based discretization will do*)
- Right answer: Surprisingly, yes.
  - (*and entropy-based discretization can do it*)

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## Error-based vs. entropy-based

A 2-class, 2-attribute problem



Entropy-based discretization can detect change of class distribution

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## The converse of discretization

- Make nominal values into “numeric” ones
- 1. Indicator attributes (used by IB1)
  - Makes no use of potential ordering information
- 2. Code an ordered nominal attribute into binary ones (used by M5')
  - Can be used for any ordered attribute
  - Better than coding ordering into an integer (which implies a metric)
- In general: code subset of attributes as binary

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## Data transformations

- Simple transformations can often make a large difference in performance
- Example transformations (not necessarily for performance improvement):
  - Difference of two date attributes
  - Ratio of two numeric (ratio-scale) attributes
  - Concatenating the values of nominal attributes
  - Encoding cluster membership
  - Adding noise to data
  - Removing data randomly or selectively
  - Obfuscating the data

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## Principal component analysis

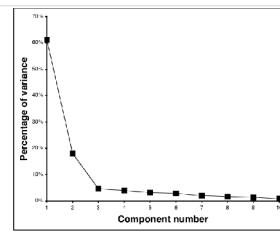
- Method for identifying the important “directions” in the data
- Can rotate data into (reduced) coordinate system that is given by those directions
- Algorithm:
  1. Find direction (axis) of greatest variance
  2. Find direction of greatest variance that is perpendicular to previous direction and repeat
- Implementation: find eigenvectors of covariance matrix by diagonalization
  - Eigenvectors (sorted by eigenvalues) are the directions

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## Example: 10-dimensional data

Axis	Variance	Cumulative
1	61.2%	61.2%
2	18.0%	79.2%
3	4.7%	83.9%
4	4.0%	87.9%
5	3.2%	91.1%
6	2.9%	94.0%
7	2.0%	96.0%
8	1.7%	97.7%
9	1.4%	99.1%
10	0.9%	100.0%



- Can transform data into space given by components
- Data is normally standardized for PCA
- Could also apply this recursively in tree learner

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## Random projections

- PCA is nice but expensive: cubic in number of attributes
- Alternative: use random directions (projections) instead of principle components
- Surprising: random projections preserve distance relationships quite well (on average)
  - Can use them to apply  $k$ D-trees to high-dimensional data
  - Can improve stability by using ensemble of models based on different projections

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## Text to attribute vectors

- Many data mining applications involve textual data (eg. string attributes in ARFF)
- Standard transformation: convert string into bag of words by *tokenization*
  - Attribute values are binary, word frequencies ( $f_{ij}$ ),  $\log(1+f_{ij})$ , or TF  $\times$  IDF:  $f_{ij} \log \frac{\# \text{ documents}}{\# \text{ documents that include word } i}$
- Only retain alphabetic sequences?
- What should be used as delimiters?
- Should words be converted to lowercase?
- Should *stopwords* be ignored?
- Should *hapax legomena* be included? Or even just the  $k$  most frequent words?

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## Time series

- In time series data, each instance represents a different time step
- Some simple transformations:
  - Shift values from the past/future
  - Compute difference (*delta*) between instances (ie. “derivative”)
- In some datasets, samples are not regular but time is given by *timestamp* attribute
  - Need to normalize by step size when transforming
- Transformations need to be adapted if attributes represent different time steps

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## Automatic data cleansing

- To improve a decision tree:
  - Remove misclassified instances, then re-learn!
- Better (of course!):
  - Human expert checks misclassified instances
- Attribute noise vs class noise
  - Attribute noise should be left in training set (*don't train on clean set and test on dirty one*)
  - Systematic class noise (e.g. one class substituted for another): leave in training set
  - Unsystematic class noise: eliminate from training set, if possible

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## Robust regression

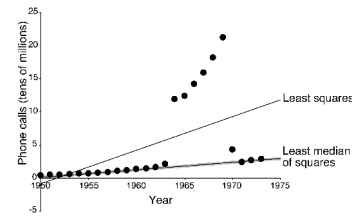
- “Robust” statistical method  $\Rightarrow$  one that addresses problem of *outliers*
- To make regression more robust:
  - Minimize absolute error, not squared error
  - Remove outliers (e.g. 10% of points farthest from the regression plane)
  - Minimize *median* instead of *mean* of squares (copes with outliers in  $x$  and  $y$  direction)
  - Finds narrowest strip covering half the observations

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## Example: least median of squares

Number of international phone calls from Belgium, 1950–1973



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## Detecting anomalies

- Visualization can help to detect anomalies
- Automatic approach: committee of different learning schemes
  - E.g.
    - decision tree
    - nearest-neighbor learner
    - linear discriminant function
  - Conservative approach: delete instances incorrectly classified by them all
  - Problem: might sacrifice instances of small classes

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## Combining multiple models

- Basic idea: build different “experts”, let them vote
- Advantage:
  - often improves predictive performance
- Disadvantage:
  - usually produces output that is very hard to analyze
  - but: there are approaches that aim to produce a single comprehensible structure

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## Bagging

- Combining predictions by voting/averaging
  - Simplest way
  - Each model receives equal weight
- “Idealized” version:
  - Sample several training sets of size  $n$  (instead of just having one training set of size  $n$ )
  - Build a classifier for each training set
  - Combine the classifiers’ predictions
- Learning scheme is *unstable*  $\Rightarrow$  almost always improves performance
  - Small change in training data can make big change in model (e.g. decision trees)

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## Bias-variance decomposition

- Used to analyze how much selection of any *specific* training set affects performance
- Assume infinitely many classifiers, built from different training sets of size  $n$
- For any learning scheme,
  - *Bias* = expected error of the combined classifier on new data
  - *Variance* = expected error due to the particular training set used
- Total expected error  $\approx$  bias + variance

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## More on bagging

- Bagging works because it reduces *variance* by voting/averaging
  - Note: in some pathological hypothetical situations the overall error might increase
  - Usually, the more classifiers the better
- Problem: we only have one dataset!
- Solution: generate new ones of size  $n$  by sampling from it *with replacement*
- Can help a lot if data is noisy
- Can also be applied to numeric prediction
  - Aside: bias-variance decomposition originally only known for numeric prediction

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## Bagging classifiers

### Model generation

```
Let  $n$  be the number of instances in the training data
For each of  $t$  iterations:
  Sample  $n$  instances from training set
  (with replacement)
  Apply learning algorithm to the sample
  Store resulting model
```

### Classification

```
For each of the  $t$  models:
  Predict class of instance using model
Return class that is predicted most often
```

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## Bagging with costs

- Bagging unpruned decision trees known to produce good probability estimates
  - Where, instead of voting, the individual classifiers' probability estimates are averaged
  - Note: this can also improve the success rate
- Can use this with minimum-expected cost approach for learning problems with costs
- Problem: not interpretable
  - *MetaCost* re-labels training data using bagging with costs and then builds single tree

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## Randomization

- Can randomize learning algorithm instead of input
- Some algorithms already have a random component: eg. initial weights in neural net
- Most algorithms can be randomized, eg. greedy algorithms:
  - Pick from the  $N$  best options at random instead of always picking the best options
  - Eg.: attribute selection in decision trees
- More generally applicable than bagging: e.g. random subsets in nearest-neighbor scheme
- Can be combined with bagging

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## Boosting

- Also uses voting/averaging
- Weights models according to performance
- Iterative: new models are influenced by performance of previously built ones
  - Encourage new model to become an "expert" for instances misclassified by earlier models
  - Intuitive justification: models should be experts that complement each other
- Several variants

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## AdaBoost.M1

### Model generation

```
Assign equal weight to each training instance
For  $t$  iterations:
  Apply learning algorithm to weighted dataset,
  store resulting model
  Compute model's error  $e$  on weighted dataset
  If  $e = 0$  or  $e \geq 0.5$ :
    Terminate model generation
  For each instance in dataset:
    If classified correctly by model:
      Multiply instance's weight by  $e/(1-e)$ 
  Normalize weight of all instances
```

### Classification

```
Assign weight = 0 to all classes
For each of the  $t$  models (or fewer):
  For the class this model predicts
    add  $-\log e/(1-e)$  to this class's weight
Return class with highest weight
```

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## More on boosting I

- Boosting needs weights ... but
- Can adapt learning algorithm ... or
- Can apply boosting *without* weights
  - resample with probability determined by weights
  - disadvantage: not all instances are used
  - advantage: if error > 0.5, can resample again
- Stems from *computational learning theory*
- Theoretical result:
  - training error decreases exponentially
- Also:
  - works if base classifiers are not too complex, and
  - their error doesn't become too large too quickly

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## More on boosting II

- Continue boosting after training error = 0?
- Puzzling fact: generalization error continues to decrease!
  - Seems to contradict Occam's Razor
- Explanation: consider *margin* (confidence), not error
  - Difference between estimated probability for true class and nearest other class (between -1 and 1)
- Boosting works with *weak* learners only condition: error doesn't exceed 0.5
- In practice, boosting sometimes overfits (in contrast to bagging)

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## Additive regression I

- Turns out that boosting is a greedy algorithm for fitting additive models
- More specifically, implements *forward stagewise additive modeling*
- Same kind of algorithm for numeric prediction:
  1. Build standard regression model (eg. tree)
  2. Gather residuals, learn model predicting residuals (eg. tree), and repeat
- To predict, simply sum up individual predictions from all models

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## Additive regression II

- Minimizes squared error of ensemble if base learner minimizes squared error
- Doesn't make sense to use it with (multiple) linear regression, why?
- Can use it with *simple* linear regression to build multiple linear regression model
- Use cross-validation to decide when to stop
- Another trick: shrink predictions of the base models by multiplying with pos. constant < 1
  - Caveat: need to start with model 0 that predicts the mean

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## Additive logistic regression

- Can use the logit transformation to get algorithm for classification
  - More precisely, class probability estimation
  - Probability estimation problem is transformed into regression problem
  - Regression scheme is used as base learner (eg. regression tree learner)
- Can use forward stagewise algorithm: at each stage, add model that maximizes probability of data
- If  $f_j$  is the  $j$ th regression model, the ensemble predicts probability  $p(1 | \vec{a}) = \frac{1}{1 + \exp(-\sum f_j(\vec{a}))}$  for the first class

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## LogitBoost

### Model generation

```

For j = 1 to t iterations:
  For each instance a[i]:
    Set the target value for the regression to
    z[i] = (y[i] - p(1|a[i])) / [p(1|a[i]) * (1-p(1|a[i]))]
    Set the weight of instance a[i] to p(1|a[i]) * (1-p(1|a[i]))
  Fit a regression model f[j] to the data with class
  values z[i] and weights w[i]
  
```

### Classification

Predict 1<sup>st</sup> class if  $p(1 | \vec{a}) > 0.5$ , otherwise predict 2<sup>nd</sup> class

- Maximizes probability if base learner minimizes squared error
- Difference to AdaBoost: optimizes probability/likelihood instead of exponential loss
- Can be adapted to multi-class problems
- Shrinking and cross-validation-based selection apply

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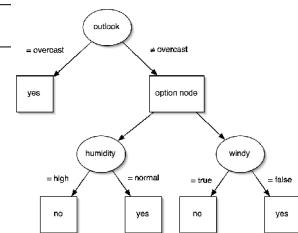
## Option trees

- Ensembles are not interpretable
- Can we generate a single model?
  - One possibility: “cloning” the ensemble by using lots of artificial data that is labeled by ensemble
  - Another possibility: generating a single structure that represents ensemble in compact fashion
- *Option tree*: decision tree with option nodes
  - Idea: follow all possible branches at option node
  - Predictions from different branches are merged using voting or by averaging probability estimates

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## Example



- Can be learned by modifying tree learner:
  - Create option node if there are several equally promising splits (within user-specified interval)
  - When pruning, error at option node is average error of options

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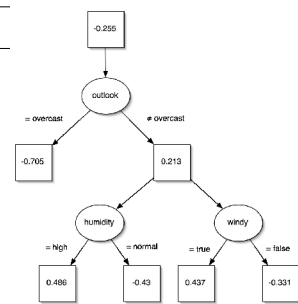
## Alternating decision trees

- Can also grow option tree by incrementally adding nodes to it
- Structure called *alternating decision tree*, with *splitter nodes* and *prediction nodes*
  - Prediction nodes are leaves if no splitter nodes have been added to them yet
  - Standard alternating tree applies to 2-class problems
  - To obtain prediction, filter instance down all applicable branches and sum predictions
    - Predict one class or the other depending on whether the sum is positive or negative

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## Example



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## Growing alternating trees

- Tree is grown using a boosting algorithm
  - Eg. LogitBoost described earlier
  - Assume that base learner produces single conjunctive rule in each boosting iteration (note: rule for regression)
  - Each rule could simply be added into the tree, including the numeric prediction obtained from the rule
  - Problem: tree would grow very large very quickly
  - Solution: base learner should only consider candidate rules that extend existing branches
    - Extension adds splitter node and two prediction nodes (assuming binary splits)
  - Standard algorithm chooses best extension among all possible extensions applicable to tree
  - More efficient heuristics can be employed instead

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## Logistic model trees

- Option trees may still be difficult to interpret
- Can also use boosting to build decision trees with linear models at the leaves (ie. trees without options)
- Algorithm for building logistic model trees:
  - Run LogitBoost with simple linear regression as base learner (choosing the best attribute in each iteration)
  - Interrupt boosting when cross-validated performance of additive model no longer increases
  - Split data (eg. as in C4.5) and resume boosting in subsets of data
  - Prune tree using cross-validation-based pruning strategy (from CART tree learner)

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## Stacking

- To combine predictions of base learners, don't vote, use *meta learner*
  - Base learners: *level-0 models*
  - Meta learner: *level-1 model*
  - Predictions of base learners are input to meta learner
- Base learners are usually different schemes
- Can't use predictions on training data to generate data for level-1 model!
  - Instead use cross-validation-like scheme
- Hard to analyze theoretically: "black magic"

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## More on stacking

- If base learners can output probabilities, use those as input to meta learner instead
- Which algorithm to use for meta learner?
  - In principle, any learning scheme
  - Prefer "relatively global, smooth" model
    - Base learners do most of the work
    - Reduces risk of overfitting
- Stacking can be applied to numeric prediction too

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## Error-correcting output codes

- Multiclass problem  $\Rightarrow$  binary problems
- Simple scheme: One-per-class coding
- Idea: use *error-correcting codes* instead
  - base classifiers predict 1011111, true class = ??
- Use code words that have large *Hamming distance* between any pair
- Can correct up to  $(d-1)/2$  single-bit errors

class	class vector
a	1000
b	0100
c	0010
d	0001

class	class vector
a	1111111
b	0000111
c	0011001
d	0101010

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## More on ECOCs

- Two criteria :
  - *Row separation*: minimum distance between rows
  - *Column separation*: minimum distance between columns (and columns' complements)
  - Why? Because if columns are identical, base classifiers will likely make the same errors
  - Error-correction is weakened if errors are correlated
- 3 classes  $\Rightarrow$  only  $2^3$  possible columns
  - (and 4 out of the 8 are complements)
  - Cannot achieve row and column separation
- Only works for problems with  $> 3$  classes

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## Exhaustive ECOCs

- *Exhaustive* code for  $k$  classes:
  - Columns comprise every possible  $k$ -string ...
  - ... except for complements and all-zero/one strings
  - Each code word contains  $2^{k-1} - 1$  bits
- Class 1: code word is all ones
- Class 2:  $2^{k-2}$  zeroes followed by  $2^{k-2} - 1$  ones
- Class  $i$ : alternating runs of  $2^{k-i}$  0s and 1s
  - last run is one short

Exhaustive code,  $k = 4$

class	class vector
a	1111111
b	0000111
c	0011001
d	0101010

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## More on ECOCs

- More classes  $\Rightarrow$  exhaustive codes infeasible
  - Number of columns increases exponentially
- Random code words have good error-correcting properties on average!
- There are sophisticated methods for generating ECOCs with just a few columns
- ECOCs don't work with NN classifier
  - But: works if different attribute subsets are used to predict each output bit

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## Using unlabeled data

- *Semisupervised learning*: attempts to use unlabeled data as well as labeled data
  - The aim is to improve classification performance
- Why try to do this? Unlabeled data is often plentiful and labeling data can be expensive
  - Web mining: classifying web pages
  - Text mining: identifying names in text
  - Video mining: classifying people in the news
- Leveraging the large pool of unlabeled examples would be very attractive

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## Clustering for classification

- Idea: use naïve Bayes on labeled examples and then apply EM
  - First, build naïve Bayes model on labeled data
  - Second, label unlabeled data based on class probabilities ("expectation" step)
  - Third, train new naïve Bayes model based on all the data ("maximization" step)
  - Fourth, repeat 2<sup>nd</sup> and 3<sup>rd</sup> step until convergence
- Essentially the same as EM for clustering with fixed cluster membership probabilities for labeled data and #clusters = #classes

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## Comments

- Has been applied successfully to document classification
  - Certain phrases are indicative of classes
  - Some of these phrases occur only in the unlabeled data, some in both sets
  - EM can generalize the model by taking advantage of co-occurrence of these phrases
- Refinement 1: reduce weight of unlabeled data
- Refinement 2: allow multiple clusters per class

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## Co-training

- Method for learning from *multiple views* (multiple sets of attributes), eg:
  - First set of attributes describes content of web page
  - Second set of attributes describes links that link to the web page
- Step 1: build model from each view
- Step 2: use models to assign labels to unlabeled data
- Step 3: select those unlabeled examples that were most confidently predicted (ideally, preserving ratio of classes)
- Step 4: add those examples to the training set
- Step 5: go to Step 1 until data exhausted
- Assumption: views are independent

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## EM and co-training

- Like EM for semisupervised learning, but view is switched in each iteration of EM
  - Uses all the unlabeled data (probabilistically labeled) for training
- Has also been used successfully with support vector machines
  - Using logistic models fit to output of SVMs
- Co-training also seems to work when views are chosen randomly!
  - Why? Possibly because co-trained classifier is more robust

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