# Ensemble methods

COMP5318 Machine Learning and Data Mining

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Reference: Tan ch.4.10, Witten ch.12, Géron ch.7







- Motivation for creating ensembles
- Ensemble methods
  - Bagging
  - Boosting AdaBoost
  - Random Forest



### What is an ensemble method?

- An ensemble combines the predictions of multiple classifiers
- The classifiers that are combined are called base classifiers, they are created using the training data
- There are various ways to make a prediction for new examples, e.g. by taking the majority vote of the base classifiers, the weighed vote, etc.
- Ensembles tends to work better than the base classifiers they combine
- Example of an ensemble:
  - 3 base classifiers are trained on the training data, e.g. k nearest neighbor, logistic regression and decision tree
  - To classify a new example, the individual predictions are combined by taking the majority vote



### Motivation for ensembles

- When do ensemble methods work?
- Let's consider an example which illustrates how ensembles can improve the performance of a single classifier:
- An ensemble of 25 binary classifiers. Each base classifier has an error rate ε =0.35 on the test set (i.e. accuracy=0.65). To predict the class of a new example, the predictions of the base classifiers are combined by majority vote.
- Case 1: The base classifiers are identical, i.e. make the same mistakes.
   What will be the error rate of the ensemble on the test set?
  - 0.35



### Motivation for ensembles (2)

- Case 2: The base classifiers are independent, i.e. their errors are not correlated. What will be the error rate of the ensemble on the test set?
- When will a new example be misclassified? Only if more than half of the base classifiers predict incorrectly.
- It can be shown that the error rate of the ensemble will be:

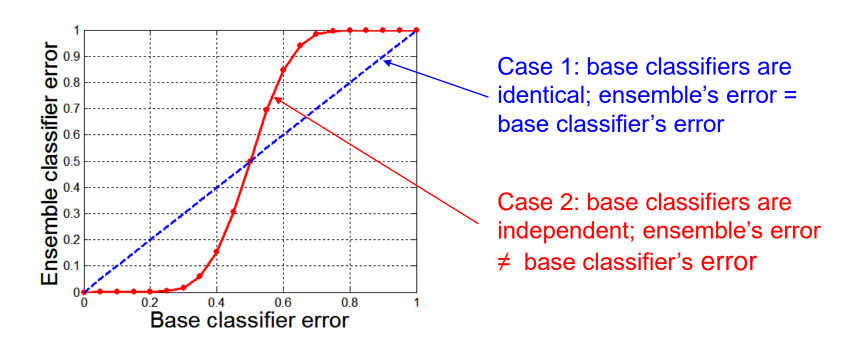
$$e_{ensemble} = \sum_{i=13}^{25} {25 \choose i} \varepsilon^{i} (1-\varepsilon)^{25-i} = 0.06$$

 0.06<0.35, i.e. the error rate of the ensemble is much lower than the error rate of the base classifiers



### Error rate graph – ensemble vs base classifier

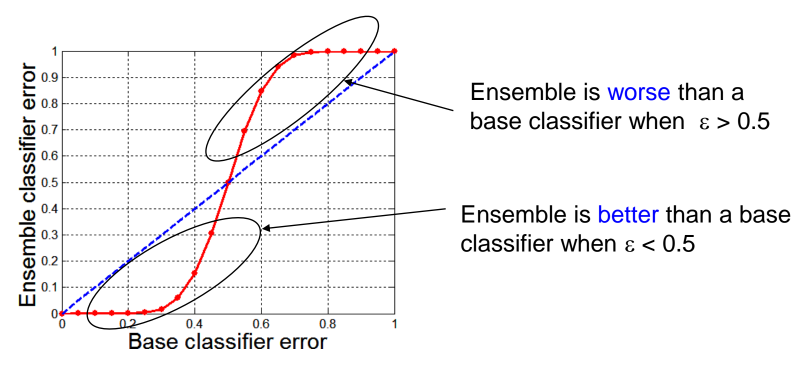
For our example of 25 binary classifiers:





### Error rate graph – ensemble vs base classifier (2)

When is the ensemble better and worse than the base classifier?



• The ensemble is better if the error of the base classifier  $\epsilon$  < 0.5, i.e. the predictions of the base classifier (which is binary) are better than random guess



### When ensembles work well

- Conditions for an ensemble to perform better than a single classifier:
  - The base classifiers should be good enough, i.e. better than a random guessing (ε <0.5 for binary classifiers)</li>
  - The base classifiers are independent of each other
- Independence in practice:
  - It is not possible to ensure total independence among the base classifiers
  - Good results have been achieved in ensemble methods when the base classifiers are slightly correlated



### Methods for constructing ensembles

- Effective ensemble consists of base classifiers that are reasonably correct and also diverse (i.e. independent)
- Methods for creating ensembles focus on generating disagreement among the base classifiers by:
  - Manipulating the training data creating multiple training sets by resampling the original data according to some sampling distribution and constructing a classifier for each training set (e.g. Bagging and Boosting)
  - Manipulating the attributes using a subset of input features (e.g. Random Forest and Random Subspace)
  - Manipulating the class labels will not be covered (e.g. errorcorrecting output coding)
  - Manipulating the learning algorithm e.g. building a set of classifiers with different parameters



# Manipulating training data - examples

- Creating multiple training sets from the original training set by sampling
- Examples: Bagging and Boosting



# **Bagging**



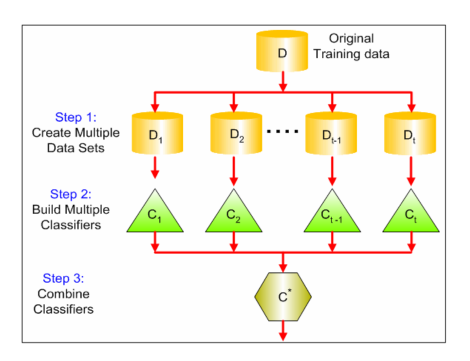
- Bagging is also called bootstrap aggregation
- A bootstrap sample definition:
  - Given: a dataset D with n example (the original dataset)
  - Bootstrap sample D' from D: contains also n examples, randomly chosen from D with replacement (i.e. some examples from D will appear more than once in D', some will not appear at all)
- For n=10, on average 65% of the examples in D will also appear in D' as
  it can be shown that the probability to choose an example is 1- (1-1/n)<sup>n</sup>

| Dataset with | 10 | examp | les: |
|--------------|----|-------|------|
|--------------|----|-------|------|

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



- Create M bootstrap samples
- Use each sample to build a classifier
- To classify a new example: get the predictions of each classifier and combine them with a majority vote
  - i.e. the individual classifiers receive equal weights





### Bagging - pseudocode

### **Model generation**

Let *n* be the number of examples in the training data For each of *M* iterations:

Sample *n* examples with replacement from training data Apply the learning algorithm to the sample Store the resulting model

### Classification

For each of the *M* models:

Predict class of testing example using model

Return class that has been predicted most often (majority vote)





- Typically performs significantly better than the single classifier and is never substantially worse
- Especially effective for unstable classifiers
- Unstable classifiers: small changes in the training set result in large changes in predictions, e.g. decision trees, neural networks are considered unstable classifiers
- Applying Bagging to regression tasks the individual predictions are averaged



# **Boosting**



# Boosting – main idea

- The most widely used ensemble method
- Idea: Make the classifiers complement each other
- How: The next classifier should be created using examples that were difficult for the previous classifiers

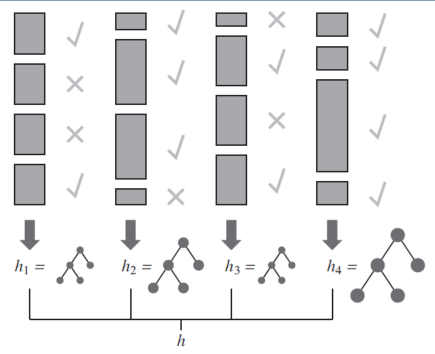




- Uses a weighed training set
- Each training example has an associated weight (≥0)
- The higher the weight, the more difficult the example was to classify by the previous classifiers
- Examples with higher weight will have a higher chance to be selected in the training set for the next classifier
- AdaBoost was proposed by Freund and Schapire in 1996







Combining decision trees

- 1 rectangle = 1 training example
- height of the rectangle corresponds to the weight of the example
- √ and X how the example was classified by the current classifier (correctly or incorrectly)
- size of the tree corresponds to the weight of its prediction in the ensemble



### AdaBoost algorithm

- Given: N samples  $\{x_n, y_n\}$ , where  $y_n \in \{+1, -1\}$ , and some way of constructing weak (or base) classifiers
- Initialize weights  $w_1(n) = \frac{1}{N}$  for every training sample
- 1. Train a weak classifier  $h_t(x)$  using current weights  $w_t(n)$ , by minimizing

$$\epsilon_t = \sum_n w_t(n) \mathbb{I}[y_n \neq h_t(x_n)]$$
 (the weighted classification error)

- 2. Compute contribution for this classifier:  $\beta_t = \frac{1}{2} \log \frac{1-\epsilon_t}{\epsilon_t}$ 3. Update weights on training points

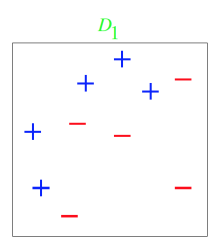
$$w_{t+1}(n) \propto w_t(n) e^{-\beta_t y_n h_t(\mathbf{x}_n)}$$

and normalize them such that  $\sum_n w_{t+1}(n) = 1$ Output the final classifier

$$h[\mathbf{x}] = \operatorname{sign}\left[\sum_{t=1}^{T} \beta_t h_t(\mathbf{x})\right]$$



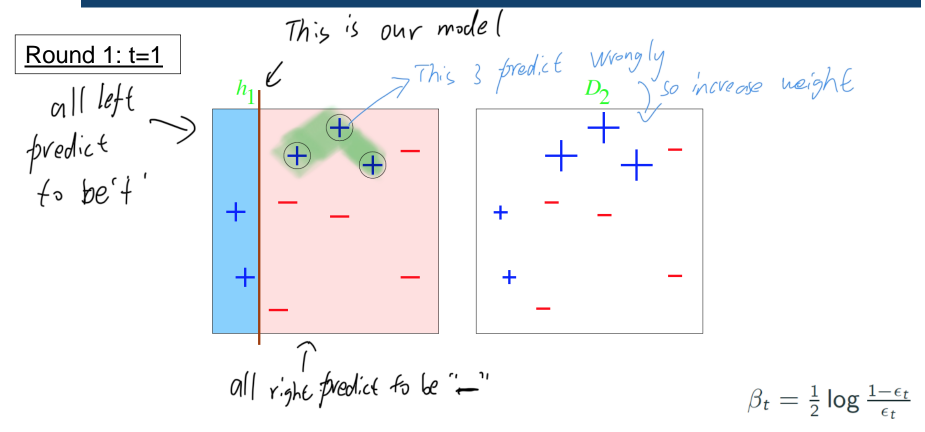




- The data points are clearly not linearly separable
- In the beginning, all data points have equal weights (the size of the data markers "+" or "-")
- Base classifier  $h(\cdot)$ : horizontal or vertical lines ('decision stumps')
  - Depth-1 decision trees, i.e., classify data based on a single attribute.



### AdaBoost: Example

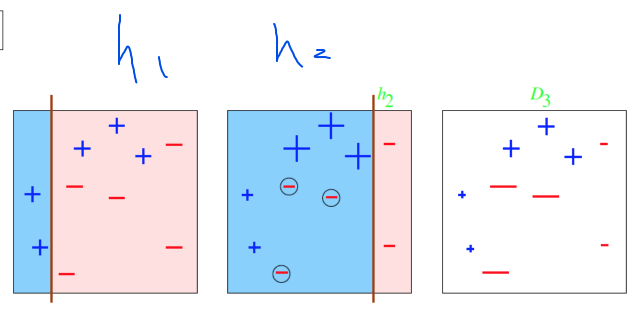


- 3 misclassified (with circles):  $\epsilon_1 = 0.3 \rightarrow \beta_1 = 0.42$ .
- Recompute the weights; the 3 misclassified data points receive larger weights



# AdaBoost: Example





- 3 misclassified (with circles):  $\epsilon_2 = 0.21 \rightarrow \beta_2 = 0.65$ .  $\epsilon_2 < 0.3$  as those 3 misclassified data points have weights < 0.1
- 3 newly misclassified data points get larger weights
- Data points classified correctly in both rounds have small weights

$$\epsilon_t = \sum_n w_t(n) \mathbb{I}[y_n \neq h_t(x_n)]$$

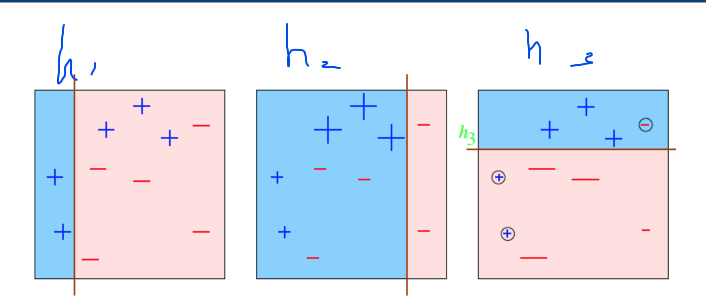
$$\beta_t = \frac{1}{2} \log \frac{1 - \epsilon_t}{\epsilon_t}$$

$$w_{t+1}(n) \propto w_t(n)e^{-\beta_t y_n h_t(\mathbf{x}_n)}$$



# AdaBoost: Example



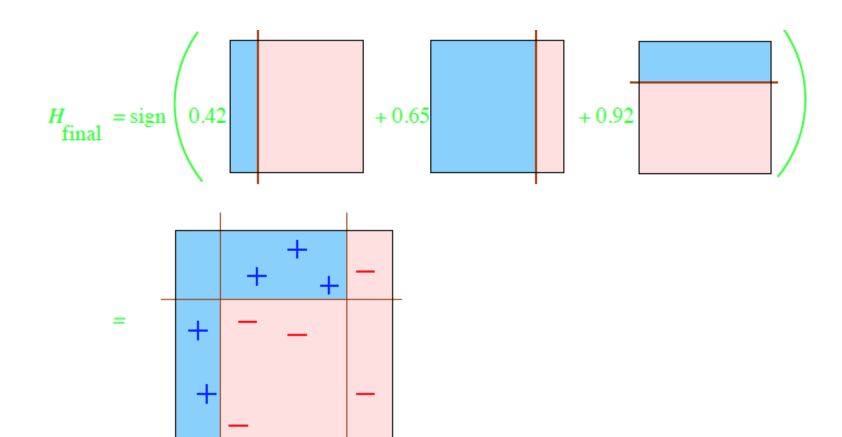


- 3 misclassified (with circles):  $\epsilon_3 = 0.14 \rightarrow \beta_3 = 0.92$ .
- Previously correctly classified data points are now misclassified, hence our error is low. Why?
  - Since they have been consistently classified correctly, this round's mistake will hopefully not have a huge impact on the overall prediction





### Combine 3 Classifier





### Why does AdaBoost work?

It minimizes a loss function related to classification error.

#### Classification loss

Suppose we want to have a classifier

$$h(\mathbf{x}) = \operatorname{sign}[f(\mathbf{x})] = \begin{cases} 1 & \text{if } f(\mathbf{x}) > 0 \\ -1 & \text{if } f(\mathbf{x}) < 0 \end{cases}$$

One seemingly natural loss function is 0-1 loss:

$$\ell(h(\mathbf{x}), y) = \begin{cases} 0 & \text{if } yf(\mathbf{x}) > 0 \\ 1 & \text{if } yf(\mathbf{x}) < 0 \end{cases}$$

Namely, the function f(x) and the target label y should have the same sign to avoid a loss of 1.



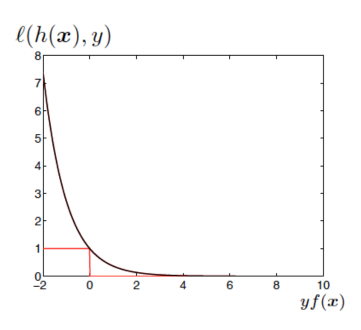
# Surrogate function of 0-1 loss?

0-1 loss function  $\ell(h(x),y)$  is non-convex and difficult to optimize.

We can instead use a surrogate loss – what are examples?

#### **Exponential Loss**

$$\ell^{\text{EXP}}(h(\mathbf{x}), y) = e^{-yf(\mathbf{x})}$$





### How to choose the *t*-classifier?

Suppose a classifier  $f_{t-1}(x)$ , and want to add a weak learner  $h_t(x)$ 

$$f(\mathbf{x}) = f_{t-1}(\mathbf{x}) + \beta_t h_t(\mathbf{x})$$

Note:  $h_t(\cdot)$  outputs -1 or 1, as does sign  $[f_{t-1}(\cdot)]$ 

How can we 'optimally' choose  $h_t(\mathbf{x})$  and combination coefficient  $\beta_t$ ?

Adaboost greedily minimizes the exponential loss function!

$$(h_t^*(\mathbf{x}), \beta_t^*) = \operatorname{argmin}_{(h_t(\mathbf{x}), \beta_t)} \sum_n e^{-y_n f(\mathbf{x}_n)}$$

$$= \operatorname{argmin}_{(h_t(\mathbf{x}), \beta_t)} \sum_n e^{-y_n [f_{t-1}(\mathbf{x}_n) + \beta_t h_t(\mathbf{x}_n)]}$$

$$= \operatorname{argmin}_{(h_t(\mathbf{x}), \beta_t)} \sum_n w_t(n) e^{-y_n \beta_t h_t(\mathbf{x}_n)}$$

where we have used  $w_t(n)$  as a shorthand for  $e^{-y_n f_{t-1}(x_n)}$ 



### The New Classifier

We can decompose the *weighted* loss function into two parts

$$\begin{split} & \sum_{n} w_{t}(n) e^{-y_{n}\beta_{t}h_{t}(\mathbf{x}_{n})} \\ & = \sum_{n} w_{t}(n) e^{\beta_{t}} \mathbb{I}[y_{n} \neq h_{t}(\mathbf{x}_{n})] + \sum_{n} w_{t}(n) e^{-\beta_{t}} \mathbb{I}[y_{n} = h_{t}(\mathbf{x}_{n})] \\ & = \sum_{n} w_{t}(n) e^{\beta_{t}} \mathbb{I}[y_{n} \neq h_{t}(\mathbf{x}_{n})] + \sum_{n} w_{t}(n) e^{-\beta_{t}} (1 - \mathbb{I}[y_{n} \neq h_{t}(\mathbf{x}_{n})]) \\ & = (e^{\beta_{t}} - e^{-\beta_{t}}) \sum_{n} w_{t}(n) \mathbb{I}[y_{n} \neq h_{t}(\mathbf{x}_{n})] + e^{-\beta_{t}} \sum_{n} w_{t}(n) \end{split}$$

We have used the following properties to derive the above

- $y_n h_t(\mathbf{x}_n)$  is either 1 or -1 as  $h_t(\mathbf{x}_n)$  is the output of a binary classifier
- The indicator function  $\mathbb{I}[y_n = h_t(\mathbf{x}_n)]$  is either 0 or 1, so it equals  $1 \mathbb{I}[y_n \neq h_t(\mathbf{x}_n)]$

### Finding the Optimal Weak Learner

#### **Summary**

$$(h_t^*(\mathbf{x}), \beta_t^*) = \operatorname{argmin}_{(h_t(\mathbf{x}), \beta_t)} \sum_n w_t(n) e^{-y_n \beta_t h_t(\mathbf{x}_n)}$$

$$= \operatorname{argmin}_{(h_t(\mathbf{x}), \beta_t)} (e^{\beta_t} - e^{-\beta_t}) \sum_n w_t(n) \mathbb{I}[y_n \neq h_t(\mathbf{x}_n)]$$

$$+ e^{-\beta_t} \sum_n w_t(n)$$

What term(s) must we optimize to choose  $h_t(x_n)$ ?

$$h_t^*(\mathbf{x}) = \operatorname{argmin}_{h_t(\mathbf{x})} \epsilon_t = \sum_n w_t(n) \mathbb{I}[y_n \neq h_t(\mathbf{x}_n)]$$

Minimize weighted classification error as noted in step 1 of Adaboost!



### Finding the Optimal Weak Learner

#### Summary

$$(h_t^*(\boldsymbol{x}), \beta_t^*) = \operatorname{argmin}_{(h_t(\boldsymbol{x}), \beta_t)} \sum_n w_t(n) e^{-y_n \beta_t h_t(\boldsymbol{x}_n)}$$

$$= \operatorname{argmin}_{(h_t(\boldsymbol{x}), \beta_t)} (e^{\beta_t} - e^{-\beta_t}) \sum_n w_t(n) \mathbb{I}[y_n \neq h_t(\boldsymbol{x}_n)]$$

$$+ e^{-\beta_t} \sum_n w_t(n)$$

#### What term(s) must we optimize?

We need to minimize the entire objective function with respect to  $\beta_t$ !

We can take the derivative with respect to  $\beta_t$ , set it to zero, and solve for  $\beta_t$ . After some calculation and using  $\sum_n w_t(n) = 1...$ 

$$\beta_t^* = \frac{1}{2} \log \frac{1 - \epsilon_t}{\epsilon_t}$$

which is precisely step 2 of Adaboost! (Exercise – verify the solution)



# Updating the Weights

Once we find the optimal weak learner we can update our classifier:

$$f(\mathbf{x}) = f_{t-1}(\mathbf{x}) + \beta_t^* h_t^*(\mathbf{x})$$

We then need to compute the weights for the above classifier as:

$$\begin{aligned} w_{t+1}(n) &\propto e^{-y_n f(\mathbf{x}_n)} = e^{-y_n [f_{t-1}(\mathbf{x}) + \beta_t^* h_t^*(\mathbf{x}_n)]} \\ &= w_t(n) e^{-y_n \beta_t^* h_t^*(\mathbf{x}_n)} = \begin{cases} w_t(n) e^{\beta_t^*} & \text{if } y_n \neq h_t^*(\mathbf{x}_n) \\ w_t(n) e^{-\beta_t^*} & \text{if } y_n = h_t^*(\mathbf{x}_n) \end{cases} \end{aligned}$$

**Intuition** Misclassified data points will get their weights increased, while correctly classified data points will get their weight decreased



### Meta Algorithm

Note that the AdaBoost algorithm itself never specifies how we would get  $h_t^*(\mathbf{x})$  as long as it minimizes the weighted classification error

$$\epsilon_t = \sum_n w_t(n) \mathbb{I}[y_n \neq h_t^*(\boldsymbol{x}_n)]$$

In this aspect, the AdaBoost algorithm is a meta-algorithm and can be used with any type of classifier



# Other Boosting Algorithms

- AdaBoost is by far the most popular boosting algorithm.
- Gradient boosting generalizes AdaBoost by substituting another (smooth) loss function for exponential loss.
  - Squared loss, logistic loss, ...
  - Choose the candidate learner that greedily minimizes the error (mathematically, it should maximize the gradient of the residuals calculated with this loss function).
  - Reduce overfitting by constraining the candidate learner (e.g., computing the residuals only over a sample of points).
- LogitBoost minimizes the logistic loss instead of exponential loss.
- Gentle AdaBoost bounds the step size  $(\beta)$  of each learner update.



# Bagging and Boosting - comparison

#### Similarities

- Use voting (for classification) and averaging (for prediction) to combine the outputs of the individual learners
- Combine classifiers of the same type, typically trees e.g. decision stumps or decision trees

#### Differences

- Creating base classifiers:
  - Bagging separately
  - Boosting iteratively the new ones are encouraged to become experts for the misclassified examples by the previous base learners (complementary expertise)
- Combination method
  - Bagging equal weighs to all base learners
  - Boosting (AdaBoost) different weights based on the performance on training data



# **Random Forest**



# Creating ensembles by manipulating the attributes

- Each base classifier uses only a subset of the features
- E.g. training data with K features, create an ensemble of M classifiers each using a smaller number of features L (L<K)</li>
  - 1) Create feature subsets by random selection from the original feature set => creating multiple versions of the training data, each containing only the selected features
- 2) Build a classifier for each version of the training data
- 4) Combine predictions with majority vote
- Example: Random Forest
  - Combines decision trees
  - Uses 1) bagging + 2) subset of features (during decision tree building, when selecting the most important attribute)



# Random Forest algorithm

n - number of training examples, m - number of all features, k - number of features to be used by each ensemble member (k<m), M - number of ensemble members

#### Model generation:

For each of *M* iteration

- 1. Bagging generate a bootstrap sample Sample *n* instances with replacement from training data
- 2. Random feature selection for selecting the best attribute Grow decision tree without pruning. At each step select the best feature to split on by considering only *k* randomly selected features and calculating information gain

#### Classification:

Apply the new example to each of the *t* decision trees starting from the root. Assign it to the class corresponding to the leaf. Combine the decisions of the individual trees by majority voting.



### Comments on Random Forests

- Performance depends on
  - Accuracy of the individual trees (strength of the trees)
  - Correlation between the trees
- Ideally: accurate individual trees but less correlated
- Bagging and random feature selection are used to generate diversity and reduce the correlation between the trees
- As the number of features k increases, both the strength and correlation increase
- Random Forest typically outperforms a single decision tree
- Robust to overfitting
- Fast as only a subset of the features are considered



### Ensembles - summary

- Ensembles combine the predictions of several classifiers
- They work when the individual classifiers are accurate and diverse
- Diversity is generated by manipulating the
  - training data (Bagging, Boosting)
  - attributes (Random Forest = bagging + random selection of attributes)
  - learning algorithm
- Some ensembles combine classifiers of the same type, some not
- Most ensembles use a majority vote to make predictions on new data, others used a weighted vote
- Ensembles have shown excellent performance often the winning solution in ML competitions is an ensemble method