# Lecture 7 **Rules and ensembles**

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# Lecture plan

- Logical rules
- Comprehension
- Rules induction
- Decision trees (reminder)
- Composition of algorithms
- Boosting
- AdaBoost and its theoretical properties
- Random Algorithm synthesis
- Random Forest
- Stacking
- The presentation is prepared with materials of the K.V. Vorontsov's course "Machine Leaning".

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#### Concepts and rules

**Concept** is a predicate on an object set *X*:

$$\varphi: X \to \{0,1\}.$$

A concept **covers** an object x, if  $\varphi(x) = 1$ .

**Rule** is a logical predicate which covers many objects from one class and few objects from other classes, and which can be simply interpreted.

**Example** (from the Russian language): *If a word is an adverb* and it ends with a hissing sound ("ж", "ч" or "ш"), you should end it with "ь".

Examples: "вскачь", "настежь".

Exceptions: "уж", "замуж", "невтерпеж".

#### Interpretable concepts

Origins in knowledge discovery in databases.

A concept φ can be interpreted if

- 1) it is formulated in natural language;
- 2) depends on a small number of feature (1-7).

#### Informative concepts

A concept  $\varphi$  is **informative** for a class c, if  $p(\varphi) = |\{x_i | \varphi(x_i) = 1, y_i = c\}| \rightarrow \max;$ 

 $n(\varphi) = |\{x_i | \varphi(x_i) = 1, y_i \neq c\}| \rightarrow \min$ . Can be reformulated in a probabilistic sense.

- $p(\varphi)$  is True Positive;
- $n(\varphi)$  is False Positive;
- $p(\varphi) + n(\varphi)$  is coverage.

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### Convolution choice problem

It is not obvious, how to convolute these two parameters:

• Precision:

$$\frac{p}{p+n} \to \max;$$

Accuracy

$$p-n \rightarrow \max$$
;

Linear cost accuracy:

$$p - Cn \rightarrow \max$$
;

• Relative accuracy:

$$\frac{p}{P} - \frac{n}{N} \to \text{max.}$$

## Convolution choice comparison

Compare with P = N = 100:

p	n	p-n	p-5n	$\frac{p}{P} - \frac{n}{N}$	$\frac{p}{n+1}$	$\frac{p}{n+p}$	$I_c$	$\mathrm{IGain}_c$	$\sqrt{p} - \sqrt{n}$
50	0	50	50	0.25	50	1	22.65	23.70	7.07
100	50	50	-150	0	1.96	0.67	2.33	1.98	2.93
50	9	41	5	0.16	5	0.85	7.87	7.94	4.07
5	0	5	5	0.03	<b>5</b>	1	2.04	3.04	2.24
100	0	100	100	0.5	100	1	52.18	53.32	10.0
140	20	120	40	0.5	6.67	0.88	37.09	37.03	7.36

#### ε, δ-rule

 $E_c(\varphi, T^{\ell}) = \frac{n_c(\varphi)}{p_c(\varphi) + n_c(\varphi)}$  is a share of falsely covered objects.

 $D_c(\varphi, T^{\ell}) = \frac{p_c(\varphi)}{\ell}$  is a share of positive objects among the covered objects.

 $\varphi(x)$  is  $\varepsilon$ ,  $\delta$ -rule (for class c), if  $E_c(\varphi, T^{\ell}) \le \varepsilon$  and  $D_c(\varphi, T^{\ell}) \ge \delta$ .

If  $n_c(\varphi) = 0$ , then the rule is **exact**.

#### Statistical rule

**Assumption**: the sample is simple.

Probability of pair (p, n) is described with hyper-geometric distribution:

$$\mathcal{H}_{P,N}(p,n) = \frac{C_P^p C_N^n}{C_{P+N}^{p+n}}.$$

**Comprehension** of predicate  $\varphi$  with sample  $T^{\ell}$ :

$$I_c(\varphi, T^{\ell}) = -\ln \mathcal{H}_{P_c, N_c}(p_c(\varphi), n_c(\varphi)).$$

 $\varphi(x)$  is **statistical rule** (for class c), if

$$I_c(\varphi, T^{\ell}) \geq \alpha$$

with  $\alpha$  being high enough (Fisher exact test).

#### Entropy-based rule

Entropy of two outcomes:

$$H(q_0, q_1) = -q_0 \log_2 q_0 - q_1 \log_2 q_1.$$

Entropy of the sample:

$$\widehat{H}(P,N) = H\left(\frac{P}{P+N}, \frac{N}{P+N}\right).$$

$$\begin{split} \widehat{H}_{\varphi}(P,N,p,n) &= \\ &= \frac{p+n}{P+N} \widehat{H}(p,n) + \frac{P+N-p-n}{P+N} \widehat{H}(P-p,N-n). \\ &\operatorname{IGain}_{c} \left( \varphi, T^{\ell} \right) = \widehat{H}(P,N) - \widehat{H}_{\varphi}(P,N,p,n). \end{split}$$

 $\varphi(x)$  is **entropy-based rule** (for class c), if  $IGain_c(\varphi, T^{\ell}) \ge G_0$  with a certain  $G_0$  being high enough.

#### Good criterions (convolutions)

•  $IGain_c$ ,  $I_c$ 

**Theorem**: 
$$\lim_{\ell \to \infty} IGain_c(\varphi, T^{\ell}) = \frac{1}{\ell \ln 2} I_c(\varphi, T^{\ell})$$
.

• Boosting criterion:

$$\sqrt{p} - \sqrt{n} \rightarrow \max$$
.

Normalized boosting criterion:

$$\sqrt{\frac{p}{P}} - \sqrt{\frac{n}{N}} \to \max.$$

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## Rule definition and examples (1/2)

**Rule** is an interpretable highly informative singleclass classifiers with refusals.

#### Examples

1. Conjunction of boundaries (terms):

$$R(x) = \bigwedge_{j \in J} [a_j \le f(x_j) \le b_j].$$

## Rule definition and examples (2/2)

2. **Syndrome** is at least *d* terms of *J* are true:

$$R(x) = \left[\sum_{j \in J} \left[ a_j \le f(x_j) \le b_j \right] \ge d \right],$$

(when d = |J|, it is conjunction, when d = 1, it is disjunction).

3. Half-plane:

$$R(x) = \left[\sum_{j \in J} w_j f_j(x) \ge w_0\right].$$

4. **Ball** is threshold similarity function:

$$R(x) = [r(x, x_0) \le r_0].$$

# Where to get concepts and how to choose rules

#### Concepts can be:

- created by yourself;
- learnt;
- given from experts.

#### Rules can be learnt with

- optimization problem solvers;
- heuristic methods;
- special machine learning algorithms.

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#### **Decision trees**

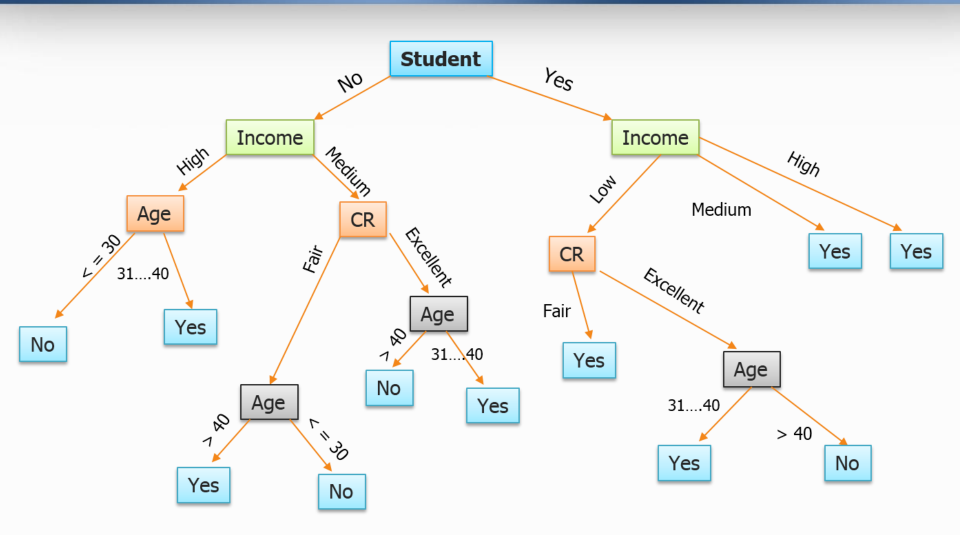
**Decision tree** is a classifier and regression algorithm.

Nodes contain splitting rules (questions).

Each edge is a possible answer to its node question.

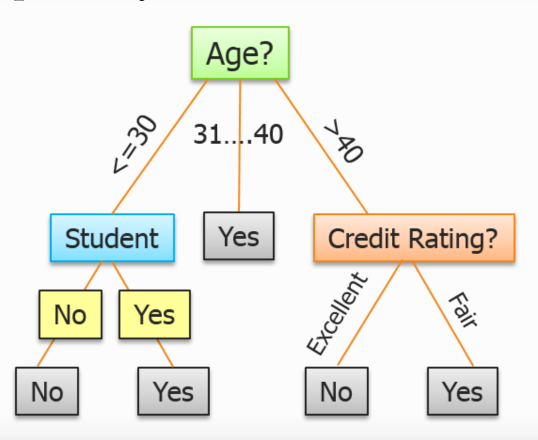
Leafs contain decisions (a class for classification problem and a number for regression problem).

### Decision tree example (1/2)



### Decision tree example (2/2)

The same classification can be achieved in a much more simpler way



#### General scheme

With splitting rules space  $\mathcal{B}$  and split quality functional  $\Phi$ .

- 1. Sent *S* to the root.
- 2. On each step process sample *S*.
- 2.1. If *S* contains objects from a single class *c*, create a leaf of the class *c* and stop.
- 2.2. Else choose a splitting rule  $\mathscr{E} \in \mathcal{B}$ , the most informative with respect to  $\Phi$ , and split the sample to  $S_1, \dots, S_k$ .
- 2.3. If stop-criterion is true, then return the most popular class in S, otherwise create k children with samples  $S_i$ .
- 3. Prune the resulting tree.

#### Three main questions

How to choose splitting rules? How to choose a stopping criterion? How to prune the tree?

## Selecting splitting rules family

Can be any family of classifiers.

• In most of the cases, it is single-feature-based rules like:

$$f_i(x) > m_i$$
;  
 $f_i(x) = d_i$ .

• Sometimes, it can be a combination.

#### Selection of feature-based rules

There are  $\ell - 1$  options to split the sample.

- Check each and pick the most informative.
- Join diapasons of values.
- Skip small diapasons.

This is how you can synthesize a rule for each feature.

## Sample splitting

- If a sample is split each time into 2 (k = 2), then  $\mathcal{B}$  is a family of binary rules, tree is binary.
- If a feature is categorical, several edges can be added.
- If a feature is real, discretization / binarization is applied.

On each step, the number of edges can differ, but usually k is fixed.

## Selecting split quality criterion

Split quality  $\Phi$  can be sometimes represented as:

$$\Phi(S) = \Phi(S) - \sum_{i=1}^k \frac{|S_i|}{|S|} \Phi(S_i).$$

The most popular:

- $\phi_h(S)$  is entropy,  $\Phi_h(S)$  is IGain;
- $\phi_g(S) = 1 \sum_{i=1}^m p_i^2$ , where  $p_i$  is probability (frequency) of ith class in sample S is **Gini index**.  $\Phi_h(S)$  is GiniGain.

Many other are used

$$GainRatio = IGain(S) / Enthropy(S)$$
.

Split quality criterion usually does not matter.

#### Stop-criteria

The most popular stop-criteria:

- one of classes is empty after splitting
- $\Phi(S)$  is lower than a certain threshold
- |*S*| is lower than a certain threshold
- tree height is higher than a certain threshold

## Pruning

**Premises:** just first node impact on performance; decision trees tend to be overfitted.

Main idea: to cut lower branches.

**Pruning** is processing of created trees, when branches are delated consequently with a certain criterion (reduction number of errors, for example).

#### Pruning algorithm scheme

Split sample to train and control in proportion 2:1. For each tree node apply operation, which is the best in terms of number of errors:

- 1) Don not change anything;
- 2) Replace node with its child (for each child);
- 3) Replace node with a leaf (for each class).

#### Examples

ID3 (Quinlan, 1986):

IGain; only  $\Phi(S) < 0$ ; no pruning.

C4.5 (Quinlan, 1993):

GainRatio; pruning.

CART (Breinman, 1984):

binary; GiniGain; pruning. Can solve regression (values in leafs).

#### Trees discussion

#### **Advantages:**

- easily understandable and interpretable;
- learning is quick;
- can work with different data type.

#### Disadvantage:

- sensitive to noise;
- easily get overfitted.

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## Weak and strong learnability

Weak learnability means that one can find an algorithm in polynomial time, performance of which would be more than 0.5.

**Strong learnability** means that one can find an algorithm in polynomial time, performance of which would be any high.

What is true: weak or strong learnability?

## Weak and strong learnability

**Weak learnability** means that one can find an algorithm in polynomial time, performance of which would be more than 0.5.

**Strong learnability** means that one can find an algorithm in polynomial time, performance of which would be any high.

#### Theorem (Schapire, 1990)

Strong learnability is equivalent to weak learnability, because any model can be strengthen with algorithm composition.

### Simple example

We have n algorithms with probabilities of correct classification answer  $p_1, p_2, ..., p_n \approx p$ . These probabilities are independent.

New algorithm will choose a class label with respect to the most preferable class within these algorithms.

Then probability of the correct classification answer is:

$$P_{vote} = p^{n} + np^{n-1}(1-p) + \frac{n(n-1)}{2}p^{n-2}(1-p)^{2} + \dots + C_{n}^{n/2}p^{n/2}(1-p)^{n/2}.$$

#### Problem formulation

Object set *X*, answer set *Y*.

Training sample  $X^{\ell} = \{x_i\}_{i=1}^{\ell}$  with known labels  $\{y_i\}_{i=1}^{\ell}$ .

Family of basic algorithms

$$H = \{h(x, a): X \to R | a \in A\},\$$

a is a parameter vector, which describes an algorithm, R is codomain (usually  $\mathbb{R}$  or  $\mathbb{R}^{M}$ ).

**Problem**: find (synthesize) a algorithm which is the most precise in forecasting label of object of *X*.

### Composition of algorithms

**Composition** of *N* basic algorithms  $h_1, ..., h_N: X \to R$  is

$$H_T(x) = C(F(h_1(x), ..., h_T(x))),$$

where  $C: R \to Y$  is a **decision rule**,  $F: R^T \to R$  is an **adjusting function**.

*R* is usually wider than *Y*.

#### Decision rule

#### **Decision rule:** $C(H(x)) \rightarrow Y$ :

- for regression,  $Y = \mathbb{R}$ C(H(x)) = H(x), or with a transformation.
- for classification on k classes,  $Y = \{1, ..., k\}$

$$C(F(h_1(x), ..., h_k(x))) = \operatorname{argmax}_{y \in Y} h_y(x)$$

• for binary classification,  $Y = \{-1, +1\}$ 

$$C(H(x)) = sign(H(x))$$

Usually this function is used:

$$L(H(x), y) = L(H(x)y)$$

### Voting

The simplest example of the adjusting function is **voting**.

Two types of voting:

- majority voting (count votes)
- soft voting (count probabilities)

We can add weights for voters (better with soft voting).

### Implementation (in Python)

### Python: VotingClassifier

#### previously

```
clf1 = LogisticRegression(random_state=1)
clf2 = RandomForestClassifier(random_state=1)
clf3 = GaussianNB()
```

### Implementation (in WEKA)

#### WEKA: Vote

```
Vote voter = new Vote();
voter.setClassifiers(classifiers)
previously
Classifier[] classifiers = {
    new J48(),
    new NaiveBayes(),
    new RandomForest()
};
```

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### Boosting problem formulation

Let's synthesize an algorithm described as

$$H_T(x) = \sum_{t=1}^T b_t h(x, a_t),$$

where  $b_t \in \mathbb{R}$  are the coefficients minimizing empirical risk

$$Q = \sum_{i}^{\ell} L(H_T(x_i), y_i) \to \min$$

for a loss function  $L(H_T(x_i), y_i)$ .

#### Gradient descent

It is hard to find an exact solution  $\{(a_t, b_t)\}_{t=1}^T$ .

We will develop function step by step

$$H_t(x) = H_{t-1}(x) + b_t h(x, a_t)$$

To do that, we estimate gradient of error function  $Q^{(t)} = \sum_{i=1}^{\ell} L(H_t(x_i), y_i)$  incremently.

This error function  $Q^{(t)}$  is a vector with the length equal to the number of objects,  $\ell$ :

$$Q^{(t)} = \left(Q_1^{(t)}, \dots, Q_\ell^{(t)}\right).$$

#### Gradient

Gradient (for *i*th element of  $Q^{(t-1)}$ ):

$$\nabla Q_i^{(t-1)} = \frac{\delta Q_i^{(t-1)}}{\delta H_{t-1}(x_i)} = \frac{\delta \left(\sum_{i=1}^{\ell} L(H_{t-1}(x_i), y_i)\right)}{\delta H_{t-1}(x_i)} = \frac{\delta L(H_{t-1}(x_i), y_i)}{\delta H_{t-1}(x_i)}.$$

Thus, we will add

$$H_t(x) = H_{t-1}(x) - b_t \nabla Q^{(t-1)}.$$

#### Parameters selection

$$H_{t}(x) = H_{t-1}(x) - b_{t} \nabla Q^{(t-1)}$$

$$b_{t} = \operatorname{argmin}_{b} \sum_{i=1}^{\ell} L(H_{t-1}(x_{i}) - b \nabla Q^{(t-1)}, y_{i}).$$

Vector  $\nabla Q^{(t-1)}$  is not a basic algorithm, so

$$a_t = \operatorname{argmin}_{a \in A} \sum_{i=1}^{\ell} L(h(x_i, a), \nabla Q^{(t-1)}) \equiv$$

$$\equiv \text{LEARN}\Big(\{x_i\}_{i=1}^{\ell}, \Big\{\nabla Q_i^{(t-1)}\Big\}_{i=1}^{\ell}\Big).$$

We can find it by linear search

$$b_t = \operatorname{argmin}_b \sum_{i=1}^{\ell} L(H_{t-1}(x_i) - bh(x_i, a_t), y_i).$$

### Generalized algorithm

**Input:**  $T^{\ell}$ , N

$$H_0(x) = \text{LEARN}(\{x_i\}_{i=1}^{\ell}, \{y_i\}_{i=1}^{\ell})$$

1. **for** t = 1 **to** T **do** 

2. 
$$\nabla Q^{(t-1)} = \left[ \frac{\delta L(H_{t-1}, y_i)}{\delta H_{t-1}} (x_i) \right]_{i=1}^{\ell}$$

3. 
$$a_t = \text{LEARN}\left(\{x_i\}_{i=1}^{\ell}, \left\{\nabla Q_i^{(t)}\right\}_{i=1}^{\ell}\right)$$

4. 
$$b_t = \operatorname{argmin}_b \sum_{i=1}^{\ell} L(y_i, h_{t-1}(x_i) - bh(x_i, a_t))$$

5. 
$$H_t(x) = H_{t-1}(x) + b_t h(x, a_t)$$

6. return  $H_N$ 

### Smoothness of *Q*

Typical *Q* is piecewise linear:

$$Q = \sum_{i=1}^{\ell} M = \sum_{i=1}^{\ell} \left[ y_i \sum_{t=1}^{N} \alpha_t H_t(x_i) < 0 \right]$$

Smooth approximation of margian loss function  $[M \le 0]$ :

- $E(M) = \exp(-M)$  is exponential (in AdaBoost)
- $L(M) = \log_2(1 + e M)$  is logarithmic (in LogitBoost)
- $Q(M) = (1 M)^2$  is quadratic (in GentleBoost)
- $G(M) = \exp(-cM(M + s))$  is Gaussian (in BrownBoost)

### Well-known algorithms

- AdaBoost
- AnyBoost
- LogitBoost
- BrownBoost
- ComBoost
- Stochastic gradient boosting

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#### **AdaBoost Basis**

$$H_T(x) = \sum_{t=1}^T b_t h(x, a_t),$$

It is classification, therefore L(H(x), y) = L(H(x)y).

Loss function is  $E(M) = \exp(-M)$ 

Term "weights" appeared earlier than "gradient".

For weight vector  $U^{\ell}$ :

- $P(h, U^{\ell})$  is the number of correctly classified objects (TP+TN)
- $N(h, U^{\ell})$  is the number of incorrectly classified objects (FP+FN)

### Main boosting theorem

#### Theorem (Freund, Schapire, 1995)

For all normalized weights vector  $U^{\ell}$ , such algorithm H = h(x, a) exist that classifies sample better than randomly:

$$N = N(H, U^{\ell}) < 1/2.$$

Then the minimum of  $Q^{(t)}$  is reached with

$$H_t = \operatorname{argmin}_H N(H, U^{\ell}),$$

$$b_t = \frac{1}{2} \ln \frac{1 - N(H_t, U^{\ell})}{N(H_t, U^{\ell})}.$$

### Objects weights

For 
$$L(H(x), y) = L(H(x)y)$$
.  

$$\nabla Q_i^{(t)} = \frac{\delta L(H_{t-1}(x_i)y_i)}{\delta H_{t-1}(x_i)} = y_i \frac{\delta L(H_{t-1}(x_i)y_i)}{\delta (H_{t-1}(x_i)y_i)} = y_i w_i,$$

where  $w_i = \frac{\delta L(H_{t-1}(x_i)y_i)}{\delta(H_{t-1}(x_i)y_i)}$  is a **weight** of object  $x_i$ .

Then the forth algorithm step is  $a_t = \text{LEARN}\left(\{x_i\}_{i=1}^{\ell}, \left\{\nabla Q_i^{(t)}\right\}_{i=1}^{\ell}\right)$ :

$$h(x, a_t) = \operatorname{argmin}_{a \in A} \sum_{i=1}^{\ell} L(h(x_i, a_t), \nabla Q_i^{(t)}) =$$

$$= \operatorname{argmin}_{a \in A} \sum_{i=1}^{\ell} L(y_i w_i h(x_i, a_t)).$$

#### AdaBoost

#### Input: $T^{\ell}$ , T

1. for 
$$i = 1$$
 to  $\ell$  do

2. 
$$w_i = \frac{1}{\ell}$$

3. **for** 
$$t = 1$$
 **to**  $T$  **do**

4. 
$$a_t = \operatorname{argmin}_A N(h(x, a_t), U^{\ell})$$

5. 
$$N_t = \sum_{i=1}^{\ell} w_i [y_i h(x_i, a_t) < 0]$$

6. 
$$b_t = \frac{1}{2} \ln \frac{1 - N_t}{N_t}$$

7. for 
$$i = 1$$
 to  $\ell$  do

8. 
$$w_i = w_i \exp(-b_t y_t h(x_i, a_t))$$

9. NORMALIZE(
$$\{w_i\}_{i=1}^{\ell}$$
)

10. return 
$$H_N = \sum_{t=1}^{T} b_t h(x, a_t)$$

#### Classification refusals

Let  $P + N \neq \ell$ . The algorithm can **refuse** to classify.

#### Theorem (Freund, Schapire, 1996)

Let for every normalized weight vector  $U^{\ell}$  an algorithm H = h(x, a) exists such that it classifies a sample at least a bit better than randomly:

$$N(H, U^{\ell}) < P(H, U^{\ell}).$$

The minimum of  $Q^{(t)}$  is reached with

$$H_t = \operatorname{argmin}_H \sqrt{P(H, U^{\ell})} - \sqrt{N(H, U^{\ell})},$$
 
$$b_t = \frac{1}{2} \ln \frac{P(H_t, U^{\ell})}{N(H_t, U^{\ell})}.$$

### Convergence

#### Theorem (Freund, Schapire, 1996)

If on each step the family H and the learning method allow to synthesize such  $H_t$  that

$$\sqrt{P(H,U^{\ell})} - \sqrt{N(H,U^{\ell})} = \gamma_t > \gamma$$

with a certain  $\gamma > 0$ , then  $H_N$  is built in a fixed number of steps.

What is the number of steps? *N*, is such that

$$Q^{(1)}(1-\gamma)^N < 1.$$

### Boosting foundamentals

$$\nu_{\theta}(a, T^{\ell}) = \frac{1}{\ell} \sum_{i}^{\ell} [H(x_i) y_i \le \theta]$$

#### Theorem (Freund, Schapire, Barlett, 1998)

If  $|H| < \infty$ , then  $\forall \theta > 0$ ,  $\forall \eta \in (0,1)$  with probability  $1 - \eta$ 

$$\leq \nu_{\theta}(a, T^{\ell}) + C \sqrt{\frac{\ln|H| \ln \ell}{\ell \theta^2} + \frac{1}{\ell} \ln \frac{1}{\eta}}.$$

It does not depend on *T*.

### **Boosting discussion**

#### Advantages:

- hard to get overfitted
- can be applied for different loss functions

#### Disadvantages:

- no noise processing
- cannot be applied for powerful algorithm
- it is hard to explain result

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### **Empirical observations**

- 1. Algorithms weights are not very important for achieving equal margins.
- 2. Objects weights are not very important for achieving difference.

### Key idea

**Idea**: we can build diverse algorithms by learning one model in different conditions.

**Precise idea**: we can use different bites of dataset.

### Synthesis of random algorithms

- Subsampling: learn algorithm on subsample.
- **Bagging:** learn algorithm on subsamples of the same length with bootstrap (random choice with returns)
- Random subspace method: learn algorithms of subspaces of features
- Filtering (next slide)

## Filtering

Let we have a sample of infinite size.

Learn first algorithm on  $X_1$ , which are first  $m_1$  objects.

Then toss a coin  $m_2$  times:

- head: add in  $X_2$  first incorrectly classified object;
- tail: add in  $X_2$  first correctly classified object.

Learn second algorithm on  $X_2$ .

Add in  $X_3$  first  $m_3$  object, on which first two classifiers give different answers.

Learn third algorithm on  $X_3$ .

### Implementation

Python: BaggingClassifier

BaggingRegressor

WEKA: Bagging

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#### **Random Forest**

For sample  $T^{\ell} = \{x_i, y_i\}_{i=1}^{\ell}$  with n features.

- 1. Choose a subsample size of  $\ell$  with bootstrap.
- 2. Synthesize decision tree for that sample, for each vertex choose n' (usually  $n' = \sqrt{n}$ ) random features.
- 3. No pruning is applied.

This can be done 100, 1000, ... times.

### Why does it work?

- It is voting
- Trees are easily get overfitted to very different subsamples
- With the growth of the sample, its performance converges

#### More details

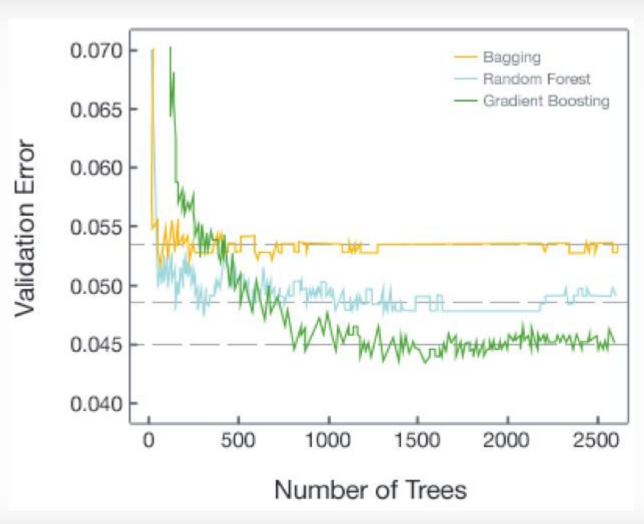
#### What to combine?

- Simple votes of trees
- Probabilities (frequency of class in the resulting leaf)

#### How to improve?

• Extremely randomized trees: use random values for splitting (it is faster).

### Ensemble method comparison



### Implementation

Python: RandomForestClassifier

RandomForestRegressor

**ExtraTreesClassifier** 

ExtraTreesRegressor

**WEKA: RandomForest** 

# Lecture plan

- Logical rules
- Comprehension
- Rules induction
- Decision trees (reminder)
- Composition of algorithms
- Boosting
- AdaBoost and its theoretical properties
- Random Algorithm synthesis
- Random Forest
- Stacking

### Stacking key idea

Instead of combining algorithms, use their predictions as new features and learn a model.

This idea can be generalized to using classification results as new features of objects.

### Blending key idea

Learn algorithms for stacking on a small (10%) hold-out data subset.

#### What also can be used?

- Algorithm mixture
- Ranking aggregation
- Model selection
- Combining several ensemble techniques