# Introduction to optimization and machine learning Lesson 3: basis of machine learning

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# General outline

- Introduction to optimization problems
- Introduction to machine learning
- Fundamentals of optimization methods
- Fundamentals of machine learning methods
- Practice of some algorithms using python

# Outline of the day

- Linear regression methods (supervised learning)
- Classification : decision tree
- Non-supervised classification
- Ensemble methods : random forest, gradient boosting

Introduction

# Bibliography



Linear regressions





- Data Science : fondamentaux et Ã(C)tudes de cas, Machine Learning avec Python et R, Eric Biernat, Michel Lutz, Eyrolles, 2015.
- Artificial Intelligence : A Modern Approach, Fourth edition, 2020. Stuart Russell and Peter Norvig.
- Vincent Barra, Antoine CornuÃ(C)jols, Laurent Miclet, "Apprentissage Artificiel. Concepts et algorithmes. De Bayes et Hume au Deep learning" Eyrolles. Mars 2021. 990 pages.

# Machine Learning

Introduction

E : set of all possible tasks.

Linear regressions

S: a system (a machine with parametrized algo)

# A more formal definition [T.M. Mitchell, 1997]

 $T \subset E$ : set of tasks called *training set* 

 $P: \mathcal{S} \times E \to \mathbb{R}$ : performance metric of a system on tasks.

A system S learns from an experience Exp if the performance of S on tasks T, measured by P, is improving.  $P(S_{\text{before Fxp}}, T) \leq P(S_{\text{after Fxp}}, T)$ 

## Example

Task T: Classifier of emails during one day Performance P : rejection rate of spams by S

Experience Exp: 1 week of emails from users

# Learning from L. Valliant, 1984 [Turing award, 2010]

# PAC ("Probably Approximately Correct")

In model of PAC Learning under the uniform distribution on X, a learning problem is defined with a concept class C, which is just a collection of functions  $f: X \to \mathbb{R}$ ; "We learn a class C of functions".

A learning algorithm A for C is a randomized algorithm which has limited access to an unknown target function  $f \in \mathcal{C}$  .

The two access models are:

Linear regressions

- random : A can draw pairs (x, f(x)) where  $x \in X$  is uniformly random
- queries : A can request the value f(x) for any  $x \in X$  of its choice.

A is given as input an accuracy parameter  $\epsilon \in [0, 1/2]$ .

Output of A: a hypothesis function  $h: X \to \mathbb{R}$ .

### PAC learning

A learns  $\mathcal{C}$  with error  $\epsilon$  if for any  $f \in \mathcal{C}$ , with high probability, A outputs an h which is  $\epsilon$ -close to f: dist $(f, h) < \epsilon$ .

Introduction

- Quantitative data
  - mesurable quantity, answer to "how much?" allow computation (mean, etc.),
  - comparaisons (equality, difference, inferior/superior)
  - Numerical : ∈ IR.
  - Discrete: number of values are limited
- Qualitative data
  - quality or features
  - answer to the "category"
  - Nominale (categorial), ex : eyes color comparison (equality / difference)
  - Ordinal
    - Order between elements (degree to test, etc.) comparison : superior / inferior
- Structured data
  - relations, etc.
  - Tree, graph, complex data, etc.

Supervised learning :

Learn from a set of examples:

$$\{(x_i,y_i)\mid i\in 1..n\}$$

Non-supervised learning :

Learn from a set of example without labels (cf. clustering)

$$\{x_i \mid i \in 1..n\}$$

Semi-supervised learning :

Learn from a set of examples with, and without labels

Reinforcement learning :

Learn when the actions on environment are rewarded by a score

Introduction

# Typology according to data

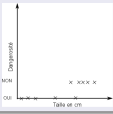
Introduction

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ullet Regression :  $(x_i,y_i)$  with  $y_i\in {\rm I\!R}$ 



• Classification :  $(x_i, y_i)$  with  $y_i$  discrete



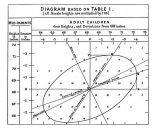
# Definition of the model [F. Galton, 1886]

model h

input value x output value y

With univariate linear regression:

$$h_{\theta}(X) = \theta_0 + \theta_1 X$$
 with  $\theta \in \mathbb{R}^2$ 



Define: Variance, co-variance, correlation.

# Univariate linear regression

# Definition of the model [F. Galton, 1886]

model h

input value  $x \longrightarrow \text{output value } y$ 

With univariate linear regression :

$$h_{ heta}(X) = heta_0 + heta_1 X$$
 with  $heta \in \mathbb{R}$ 

Find parameter  $\beta$  such that  $h_{\beta}(X)$  is the closest to Y

# Mean square error (MSE) function

$$J_{x,y}(\theta) = \frac{1}{2n} \sum_{i=1}^{n} (h_{\theta}(x_i) - y_i)^2$$

Gradient descent :  $\theta_j := \theta_j - \alpha \frac{\partial}{\partial \theta_i} J(\theta)$ 

# Definition of the model

# model h

input value  $x \longrightarrow \text{output value } y$ 

With multivariate linear regression:

$$h_{\theta}(X) = \theta_0 + \theta_1 X_1 + \theta_2 X_2 + \ldots + \theta_n X_n$$

# Mean square error function

$$J(\theta) = \frac{1}{2m} \sum_{i=1}^{m} (h_{\theta}(x_i) - y_i)^2$$

Gradient descent :  $\theta_j := \theta_j - \alpha \frac{\partial}{\partial \theta_i} J(\theta)$ 

Normalization (scaling) of the predictors : reduced centered variable ( $\mu = 0$ ,  $\sigma^2 = 1$ )

Scikit-learn is a library in python with MLmodels, and related tools. Open source, BSD license, https://scikit-learn.org/

```
From example Ordinary Least Square
from sklearn import linear_model
reg = linear_model.LinearRegression()
reg.fit ([[0, 0], [1, 1], [2, 2]], [0, 1, 2])
print(reg.coef_)
```

```
see code : linear_reg.ipynb and url :
https://scikit-learn.org/stable/auto_examples/linear_
model/plot_ols.html#
sphx-glr-download-auto-examples-linear-model-plot-ols-py
```

# Ordinary Least Square (OLS)

After algebraic computation,

$$\beta = (X^{\mathsf{T}}X)^{-1}X^{\mathsf{T}}y$$

where X is the matrix of predictor values with n lines of ppredictors, y the vector of predicted values

## Complexity

 $\mathcal{O}(p^2n)$  for multiplication of  $X^{T}X$ 

 $\mathcal{O}(pn)$  for multiplication of  $X^{\mathsf{T}}y$ 

 $\mathcal{O}(p^3)$  to compute the LU factorization of  $X^{\mathsf{T}}X$  and compute the product  $(X^{T}X)^{-1}X^{T}y$ 

if  $n \ge p$  then complexity is  $\mathcal{O}(p^2 n)$ , if n < p then complexity is  $\mathcal{O}(p^3)$ .

## Iterative method: gradient descent

Minimize  $J_{x,y}(\beta) = \frac{1}{2n} \sum_{i=1}^{n} (h_{\beta}(x_i) - y_i)^2$  with gradient descent Gradient of  $J_{x,y}(\beta)$  is a close formula (time complexity  $\mathcal{O}(np)$ ) Gradient step for each variable j:

$$\beta_j := \beta_j - \alpha \frac{\partial}{\partial \beta_j} J(\beta)$$

The time complexity for each "classical" gradient step  $\mathcal{O}(pn)$ , and we can expect around p steps...

Any gradient variant can be used : stochastic gradient, nesterov, adam, nadam, etc. Conjugate gradient  $(\mathcal{O}(p^2n))$ 

# Linear models are simple to interpret:

- Effect of a single predictor : when the predictor  $x_i$  increases by a factor 2, then the response is increased by the term  $2\beta_i$
- Importance of predictors can be compared : when the model is  $y = 0.001x_1 + 10x_2 + \epsilon$ then variable  $x_2$  is more important than  $x_1$ (if the scale of predictors are similar!!!)

# Sometime, it is more useful:

to understand the relation between variables, and response than having an accurate model of prediction of response

⇒ Tradeoff between Explanation vs. Prediction, toward XAI...

# Scaling the data

# Don't forget to scale your data :

- Between minimum and maximum :
  - $z_i = rac{x_i \min[x]}{\max[x] \min[x]}$ , then  $z_i \in [0, 1]$
- Using mean, and standard deviation :

 $z_i=rac{x_i-ar{x}}{\sigma}$  where  $ar{x}$ , and  $\sigma$  are mean, and std dev. of  $x_i$  for all i, then, E[z]=0, and  ${
m Var}[z]=1$ 

Robust scaling :

```
z_i = \frac{x_i - \text{med}[x]}{q_3[x] - q_1[x]} where \text{med}[x] is the median, and q_1[x], q_3[x] first, and third quartile of x_i for all i, then, \text{med}[z] = 0
```

See StandardScaler, and Pipeline in scikit learn See also the exo01.py

### Definition of the model

With polynomial linear regression:

$$h_{\theta}(X) = \theta_0 + \theta_1 X_1 + \theta_2 X_2 + \theta_3 X_1^2 + \theta_4 X_2^2 + \theta_5 X_1 X_2 + \dots$$

# Mean square error function

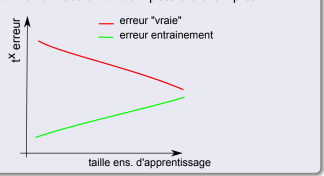
$$J(\theta) = \frac{1}{2m} \sum_{i=1}^{m} (h_{\theta}(x_i) - y_i)^2$$

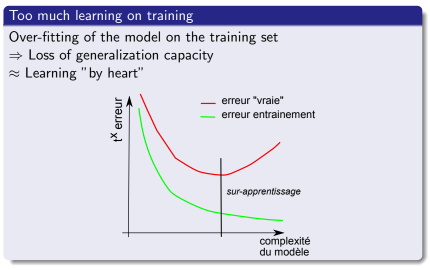
Linear regression is simple, and fun, but...

# Errors of models

### Relation between errors

- Learning error : error rate on the training set
- "True" error : error rate on the all possible examples





Complexity metrics such as the polynomial degree

Suppose a function f to learn such that  $y_i = f(x_i) + \epsilon_i$ , where  $\epsilon_i$  is the noise a null mean, and variance  $\sigma^2$ .  $\{(x_i, y_i)\}$  a training set, h a learnt model, and  $\bar{h}$  the "average" model learn on all possible sets.

The variance of a model can be decomposed (see proof) by :

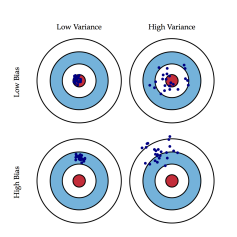
$$E[(h(x_i) - y_i)^2] = E[(h(x_i) - f(x_i))^2] + E[(f(x_i) - y_i)^2]$$

# Relation biais/variance

$$E[(h(x_i)-y_i)^2]=$$

$$E[(\bar{h}(x_i) - f(x_i))^2] + E[(h(x_i) - \bar{h}(x_i))^2] + E[(f(x_i) - y_i)^2]$$

Variance of model =  $Bias^2 + Variance + Variance$  of Noice



- Error from the bias: difference between predictions and true values
- Erreur from the variance: variability of the prediction for one given data x

Source Scott Fortmann-Roe :

http://scott.fortmann-roe.com/docs/BiasVariance.html

### Definition

$$R^2 = 1 - \frac{\text{Variance of the residus}}{\text{Variance of the data}}$$

with residus :  $r_i = h_\beta(x_i) - y_i$ 

The  $R^2$  is the part of variance of f explained by the model h

- $R^2 < 1$ , but  $R^2$  can be negative
- $R^2 = 0$  when the model h is equal to mean value, i.e. when  $h(x) = E[y_i]$ .
- $R^2 < 0$  when the model h is worst than  $h(x) = E[y_i]$
- $R^2 = 1$  when the model h is perfect, h(x) = f(x) on the set to estimate  $R^2$
- In general, but depending on the context, a relevant  $R^2$  is above 0.8.

# Goodness of fit: Coefficient of determination

Compute the  $R^2$  for the linear regression example.

# Evaluation of a model quality

### Technique

Partitionning the set into:

- Training set ( $\approx 70\%$ )
- **Test** set, independent one ( $\approx 30\%$ )

The error rate can be estimated without bias on the test set.

### Drawback

- An initial large set is requiered
- Dilemma :
  - The larger the test set, the better the estimation is
  - The larger the training set, the better the model is

# Resampling method

Allow to estimate the generalization error

### K-folds cross-validation

Partition randomly the set into K blocks

For each blok k.

Design a model using the k-1 other training blocks

Compute the test error  $e_k$  on the block k

Compute the mean of errors  $e_k$ 

# Other techniques:

- Leave-one-out (K = n)
- Bootstrap, bagging, etc.

See example in scikit-learn "Underfitting vs. Overfitting" :

```
Polynomial regression of f(x) = \cos(\frac{3\pi}{2}x) with polynomial of
degree 1, 4, and 10
from sklearn.pipeline import Pipeline
from sklearn.preprocessing import PolynomialFeatures
from sklearn.linear_model import LinearRegression
polynomial_features = PolynomialFeatures(degree=degrees[i],
                  include bias=False)
linear_regression = LinearRegression()
pipeline = Pipeline([
   ("polynomial_features", polynomial_features),
   ("linear_regression", linear_regression) ])
pipeline.fit(X[:, np.newaxis], y)
```

# Sparse approach: regularization methods

# Objective function (nota: penalty approach)

$$J(\theta) = L(\theta) + \Omega(\theta)$$

- $L(\theta)$ : training error, distance between data and prediction of the model
- $\Omega(\theta)$ : regularization, cost of the model complexity

### Goal of sparse model

Reduce the complexity of the model in order to reduce the variance of prediction, and to reduce the generalization error.

# Regularization: model complexity metrics

- Norme  $L_2: \Omega(\theta) = \lambda \|\theta\|^2 = \lambda \sum_{i=1}^n \theta_i^2$
- Norme  $L_1: \Omega(\theta) = \lambda \|\theta\|_1 = \lambda \sum_{i=1}^n |\theta_i|$

# Ridge:

$$J(\theta) = L(\theta) + \lambda \sum_{i=1}^{\infty} \theta_j^2$$

Lasso:

$$J(\theta) = L(\theta) + \lambda \sum_{i=1}^{\infty} |\theta_i|$$

ElasticNet:

$$J(\theta) = L(\theta) + \lambda \sum_{i=1} ((1 - \alpha)\theta_j^2 + \alpha |\theta_j|)$$

Warnings: parameters tuning

(use cross-validation to select relevant parameter values)

See also: forward/backward selection, LARS, dropout, etc. Indeed, the selection of a model is also an optimization problem that combined combinatorial, and numerical optimization!...

### Goal

Find a function :  $f(x) \in \{0, 1\}$ 

# Approach

Transform the model h into a binary response :

$$r(x) = \begin{cases} 0 & \text{si } h(x) < 0.5\\ 1 & \text{si } h(x) \geqslant 0.5 \end{cases}$$

The model h is interpreted as probability function :

$$h(x) \in [0,1]$$
, and  $h(x) = Pr(y = 1|x)$ .

# Sigmoid fonctions

Transform a real number from  ${\rm I\!R}$  into [0,1] using;

• hyperbolic tangent, inverse of normal density, logistic function

# Logistic regression

# Model: linear model composed with logistic function

$$h_{\theta}(X) = g(\theta_0 + \theta_1 X_1 + \theta_2 X_2 \dots + \theta_n X_n)$$
 with g logistic function



# Loss function: cross-entropy

$$j(h, y) = -\Pr(y = 1) \log \Pr(h = 1) - \Pr(y = 0) \log \Pr(h = 0)$$
 which gives :

$$i(h(x), y) = -y \log(h(x)) - (1 - y) \log(1 - h(x))$$

$$J(\theta) = \frac{1}{m} \sum_{i=1}^{m} j(h_{\theta}(x_i), y_i)$$

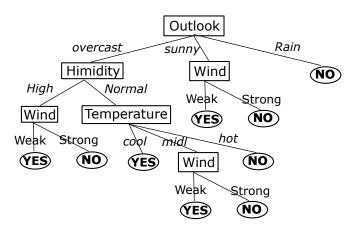
### Goal

Find a function :  $f(x) \in \{0, 1, ..., k - 1\}$ 

- For each class c, we build a binary classifier  $h^{(c)}$  which measures the probability  $y \in c$  (and  $y \notin c$ )
- The predicted class is the class with the highest probability :

$$y^{\mathsf{pred}} = \mathsf{argmax}_{c \in \mathcal{C}} \ h^{(c)}(x)$$

# Decision tree classifier One of the most explainable ML model



if Outlook = "overcast" and Humidity = ... then playball = Yes Compact way to represent a set of inference relations

# Learning with decision tree

# Design a tree :

- Nodes : **select** one attribute (decision variable  $X_j$ ) as label, the edges are labelled by the value of the attributes
- Leaves : cut the tree with target attribute y

# Targeted quality of the tree

- Low error rate
- Low generalization error
- Small tree in order to be explainable
- etc.

Learning algos: ID3, C4.5, CART, CHAID, evolutionary algo., etc.

# One classical learning approach

```
Top-down approach (binary tree)
function buildTree(X)
if all examples of X are in the same class, or stop(X) then
  Create a leaf with the label of examples
else
  Select one attribute to create a node
  Positive, and negative values in X create two sets X_0 et X_1
  buildTree(X_0)
  buildTree(X_1)
end if
```

# Greedy top-down approach

For each node.

Select the best attribute according to a metric on X

### Recursive call until:

- All examples are in the same class
- A new partitioning does not improve the prediction quality
- Top-down : from root node
- Greedy approach: best "local" choice without backtrack Beware of local optima...

#### Pros

- Easy to understand : explainable IA
- White-box model (neural net. is black-box)
- Data preparation is minimal: no scaling, etc.
- Numerical, and categorial data are possible
- Robust to outliers

#### Cons

- Learning a decision tree is NP-complet
- Greedy approach is sub-optimal
- Complex tree, overfitting
- Bias toward small tree, attribute with more values, etc.
- Interaction between is difficult to detect
- Problems are difficult to learn with tree (xor, parity, multiplexer)

Ross Quinlan, University of New South Wales, Australia, 1986

Top-down greedy algorithm based on information gain

### Principle

- Compute entropy of all attributes  $X_j$  using training set X
- Select the attribute with maximum information gain
- Partitionning the set X
- Recursive call on each subset of attribute values  $X_0, X_1, \ldots$

## Entropy H

Quantity of information (disorder) of a set

$$H(X) = -\sum_{s \in D_X} p(s) \log_2 p(s)$$

- X : data set
- $D_X$ : set of classes y of X
- p(s): proportion of class  $s \in D_X$  in X

When H(X) = 0, X is perfectly classified.

### Mutual Information (cross-entropy)

Measure the joint information of 2 random variables (information one variable from the other one)

$$I(S,T) = -\sum_{s \in D_S, t \in D_T} p(s,t) \log_2 \frac{p(s,t)}{p(s)p(t)}$$

### Information gain ( - mutual information)

Measure the entropy difference before, and after the partitionning

$$IG(S, T) = -I(S, T) = H(S) - \sum_{t} p(S_{t})H(S_{t})$$

- $T = \{S_1, \ldots\}$  partitionning of  $S = \bigcup_t S_t$
- $p(S_t) = \sharp S_t/\sharp S$
- H(S),  $H(S_t)$ : entropies of S, and  $S_t$

```
ID3(examples, target, attributes):

if attributes is empty then

return a leaf the most frequent label

elseif all examples are positive (resp. negatives) then

return a leaf positive label (resp. negative)

else

A \leftarrow attribute the highest information gain

create a node with label A

for each values v_i of A
```

**add** a branch v; to the node

ID3(examples( $A = v_i$ ), target, attributs -A)

### C4.5 algorithm Ross Quinlan, 1993

### Improvement of ID3

- Use the information gain ratio instead of IG:
  - IG(S, T) is bias toward attributes vers a larger number of values ratioIG(S,T) = IG(S,T)/H(T)
- "null" values are possible :

Example are ignored when compute the node

- Can handle attribute with real number :
  - Discretization with  $P(A < a_i)$

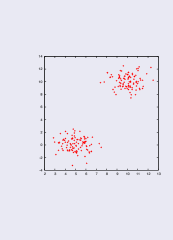
for all possible values of A, IG computation

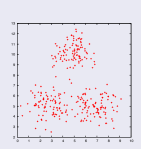
- Pruning to reduce the tree size :
  - Bottom-up Technique: final branches are pruned when the error rate is larger than with a simple leaf



# Clustering: graphical examples

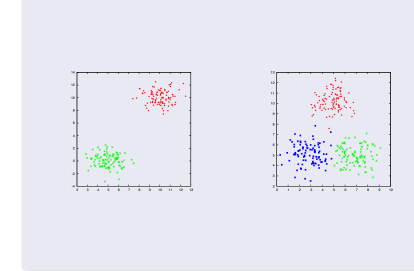
In decision / variable / attribute space :





# Clustering: graphical examples

In decision / variable / attribute space :



# Clustering: definition

## **Partitionning**

#### input:

Set of *n* points / examples / observations

$$E = \{e_1, e_2, \ldots, e_n\}$$

### output:

Partition of E

$$P = \{P_1, P_2, \dots, P_k\}$$

equivalent of coloring function  $c: E \to \{1, ..., k\}$ 

# Clustering: definition

# Partitionning

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Set of *n* points / examples / observations

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### output:

Partition of E

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equivalent of coloring function  $c: E \rightarrow \{1, ..., k\}$ 

How many possible partitions with k clusters?

# **Partitionning**

#### input:

Set of *n* points / examples / observations

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#### output:

Partition of E

$$P = \{P_1, P_2, \dots, P_k\}$$

equivalent of coloring function  $c: E \to \{1, ..., k\}$ 

How many possible partitions with k clusters?

$$k^n/k!$$

Huge even for n = 100, and k = 2, how to select one?...

# Associated optimization problem

### Optimisation problem

Criterium quality of the partition:

$$U:\mathcal{P}(E)\to\mathbb{R}$$

But without using labels (non-supervised learning)!

Find a good partition is equivalent to maximize a criterium quality:  $\operatorname{argmax}_{P \in \mathcal{P}(E)} U(P)$ 

Use an optimization method (local, greedy, etc.)

# Quality criterium

### Additive expression

Usually, the criterium is additive on clusters:

$$U(P) = \sum_{i=1}^{K} w(P_i)$$

where w is quality metric of one cluster

#### Examples

Sum of square distance between points of the cluster :

$$w(P_i) = \sum_{x_1 \in P_i} \sum_{x_2 \in P_i} d^2(x_1, x_2)$$

Probability of observation of points in the cluster:

$$w(P_i) = \prod_{x \in P_i} Pr(x|\theta_i)$$

# Clustering algorithms

# Differents approaches :

Hierarchical partitionning :

Agglomeration (or division) of clusters according to criterium (distance, etc.)

Dendrogramm

- Centroid partitionning:
   Use the center of cluster
   k-means (cf. after)
- Partitionning based on probability distribution
   A cluster is encoded by probability (parametric) distribution
   Algorithm E-M, Gaussian mixture models
- Partitionning based on density :
   According to the local density of points, cluster growth

# The famous k-means method: Pseudo-code

#### k-means

Select (randomly) k centers  $\mu_1^{(1)}, \dots, \mu_k^{(1)}$  repeat

Set the examples to each center according to distance :

$$P_i^{(t)} = \{e_j : d(e_j, \mu_i^{(t)}) \le d(e_j, \mu_a^{(t)}) \ \forall a = 1..k\}$$

Update the center of each cluster (mean of clusters) :

$$\mu_i^{(t+1)} = \frac{1}{\sharp P_i^{(t)}} \sum_{e_j \in P_i^{(t)}} e_j$$

until no more modification (convergence)

#### Comments:

The algorithm is based on parameter k, and a distance metric k-means is a local algorithm such that  $U(P^{(t+1)}) \leq U(P^{(t)})$ 

# Advantageous / drawbacks

#### Advantageous

- Easy to interpret
- Easy to code
- Polynomial complexity

#### Drawbacks

- Parameter k of the number of the clusters
- Shape of cluster are supposed to be "spherical"
- Clusters are supposed separable

# Ensemble methods

### **Principles**

Instead of one learner, use a team of learner!

Unity is strength

... if the team learns well together....

#### example

Majority voting:

A binary "stupid" predictor with error rate 0.4

What is the error rate of n independent "stupid" predictors with a majority voting mechanism?

- Efficient algorithm
- "easy" to train, parallelisation
- Less easy interpretable than decision tree, but better prediction performance
- Ensemble method

Breiman, Leo (2001). "Random Forests". Machine Learning 45 (1): 5-32.

- Ensemble method : use a team of "simple" predictors
- Simple predictor : decision tree
- Prediction :
  - Majority voting for clustering
  - Mean of predictions for regression
- Key of the learning mechanism : design a set of heterogeneous trees

# Principle

 ${\sf random\ forest} = {\sf tree\ bagging} + {\sf features\ sampling}$ 

Decision tree are too much dependent of the training set.

To build each tree, use a fragmented view of the problem :

- Draw (with replacement) a sub-set of examples
  - $\Rightarrow$  bagging
  - Draw (with replacement) a sub-set of features/attributes
    - $\Rightarrow$  features sampling

# Reduce the variance using a forest

$$V_{forest} = \rho \sigma^2 + \frac{1 - \rho}{B} \sigma^2$$

where:

- $\sigma^2$  : variance of one decision tree
- B : number of trees
- ullet ho correlation between trees

The feature selection could reduce the coefficient  $\rho$  ( $\sqrt{n}$  features are randomly selected in feature selection step)

## Others versions

- Extremely randomized trees:
   For each selected variables, the split is also random (division of attribute values)
- rotation forest : Principle component analysis (PCA) before build the tree

Geurts, Pierre and Ernst, Damien and Wehenkel, Louis, Extremely randomized trees, Machine Learning, vol. 6, 1, pp. 3-42, 2006.

Rodriguez JJ1, Kuncheva LI, Alonso CJ, Rotation forest : A new classifier ensemble method, IEEE Trans Pattern Anal Mach Intell. 2006 Oct;28(10):1619-30.

### Bibliography

- adaBoost pour adaptive boosting [GÃPrdel price 2003] : Yoav Freund et Robert Schapire, A desicion-theoretic generalization of on-line learning and an application to boosting, Journal of Computer and System Sciences, vol. 55, Num. 1, 1997, p. 119-139
- Gradient boosting : J.H. Friedman, Greedy function approximation a gradient boosting machine, Ann. Statist., Vol., Num. 5 (2001), 1189-1232.

# Boosting principle

### Principle

- Iterative design of meta-algorithm based on trees (or another "weak" learner)
- The errors of the previous iteration are taken into account to build the next one :

The next weak learner learn the residus of the error of the previous ones

The weak leaners are "boosted"

- Random forest :
  - Majority voting

Weak learners are built in parallel

- Boosting :
  - Weighted sum of the weak learners Weak learners are built iteratively

$$H(x) = \operatorname{sign}(\sum_{i=1}^{B} \alpha_i h_i(x))$$

with  $h_i(x) \in \{-1, 1\}$  instead of  $\{0, 1\}$ 

Initialize the distribution of examples  $\forall i = 1, ..., m, D_1(i) = \frac{1}{m}$ 

# iboost algorithm

```
\begin{aligned} & \textbf{for } t = 1, \dots, T \textbf{ do} \\ & h_t = \underset{h \in \mathcal{H}}{\operatorname{argmin}} \ \epsilon_t \\ & \text{ with } \epsilon_t \text{ the weighted classification error (residues)} : \\ & \epsilon_t = \sum_{i=1}^m D_t(i).[y_i \neq h(x_i)] \end{aligned} Weights of classifier : \alpha_t = \frac{1}{2} \ln \frac{1-\epsilon_t}{\epsilon_t} Update weights : \forall i = 1, \dots, m, \ D_{t+1}(i) = \frac{D_t(i)e^{-\alpha_t y_i h_t(x_i)}}{Z_t}
```

end for

$$H(x) = \text{sign}(\sum_{t=1}^{T} \alpha_t h_t(x)) \text{ with } h(t) \in \{-1, 1\}, y_i \in \{-1, 1\}$$

with  $Z_t = 2\sqrt{\epsilon_t(1-\epsilon_t)}$ 

# Gradient boosting: Principle

 $\mathsf{gradient}\ \mathsf{boosting} = \mathsf{descente}\ \mathsf{de}\ \mathsf{gradient} + \mathsf{boosting}$ 

### Boosting:

sequential learning :  $H(x) = sign(\sum_{i=1}^{B} \alpha_i h_i(x))$ 

### Gradient descent :

Generalization the loss function : residues are replaced by the negative gradient of previous ones

```
See: http:
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//orbi.ulg.ac.be/bitstream/2268/163521/1/slides.pdf

There is many techniques, algorithms.

The principle is to use:

- a good model according to data, and available information
- an efficient optimization algorithm to find the parameters of the model according to data, and available information

Have fun with optimization, and machine learning!

To go futher: https://fidle.cnrs.fr/w3/