# Fiche Machine Learning

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### General Stuff

- Supervised learning, learning from humain supervision
- Unsupervised Learning, learn without humain supervision
- Step for Classification:
  - Trainig phase: to give the concept of classes to a machine using labeled data
  - Testing phase: to determine the calss of new unseen (unlabeled) data

## Nearest Neighbour

- Binary classification
- Find the nearest neighbour, and classify x to the same class
- k-nearest neighbour:
  - Algo:
    - \* Compute distances to all samples from new data x
    - \* Pick k-neighbours that are nearest to x
    - \* Majoruty vote to classify x
  - Other
    - \* The boundary becomes smoother as k increases
    - \* Lower computational cost for lower k

- \* k-NN better generalizes given may samples
- Pros
  - \* Simple, only with a single parameter k
  - \* Applicable to multi-class problems
  - \* Good performance, effective in low dimension data
- Cons
  - \* Costly to compute distances to search for the nearest
  - \* Memory requirement: must store all the training set

### **Decision Trees**

- Test the attributes (features) sequentially
- Each leaf node bears a category label, and the test pattern is assigned the category of the leaf node reached.
- Tree Construction:
  - Choose the best question (according to the information gain), and split the input data into subsets
  - Terminate: Call branches with a unique class labels leaves
  - Grow: Recursively extedn other branches
- Entropy measure of uncertainty number of bit of information

$$ENTROPY = \sum_{i} -p_i \log_2 p_i$$

• Information gain:

As about attribute A for a data set S that has Entropy ENT(S) and get subsets  $S_v$  according to the value of A

$$GAIN = ENT(S) - \sum_{v \in Valuers(A)} \frac{|S_v|}{|S|} ENT(S_v)$$

• Gini impurity: Another definition of predicatabiliby (impurity)

$$\sum_{i} p_i (1 - p_i) = 1 - \sum_{i} p_i^2$$

- Overfitting, when the learned models are overly specialized for the training smaples (Good result on training data, but generalizes poorly)
- The simplest explanation compatible with data tends to be the right one
- To avoid overfitting we can use validation set and pruning. Pruning means simplifying/compressing and
  optimizing a decision tree by removing sections of the tree that are uncritical and redundant to calssify
  instances.

## Challenge in machine learning

• The missclasification of a model f rate on a training data D:

$$err(f, D) = \frac{1}{N} \sum_{i=1}^{N} Ind(f(x_i) \neq y_i)$$

- Overfitting occurs due to:
  - Non-representative sample
  - Noisy examples
- K-fold cross validation (training set and validation set)
- Inuitions in low-dimensions do not apply in hight-dimensions. Techniques for dimensionality reduction / feature selection exist.
- Error due to **Bias**: the difference between the average (expected) prediction of our model and the correct value
  - It is the discrepancy between its averaged estimated and true funcion
  - Low model complexity ⇒ High-bias
  - High model complexity  $\implies$  Low-bias
- Error due to **Variance**: The variability of a model prediction for a given data point between different realizations of the model
  - It is the expected divergence of the estimated predicton from its average value.
  - Low model complexity  $\implies$  Low-variance
  - High model complexity ⇒ High-variance
- $MSE = Variance + Bias^2$

## Regression

- Regression = Real valued output
- Lienar Regression tries to estimate the function f(x) and predict the output by

$$\hat{f}(x) = \sum_{i=0}^{d} w_i x_i = w^T x$$

- To measure the error for N samples we use the mean square error

$$E_{in}(\hat{f}) = \frac{1}{N} \sum_{n=1}^{N} (\hat{f}(x_n) - y_n)^2$$

The wieght vector that sets the gradient to zero minimizes the errors. RSS is the sum of squared errors

$$w = (X^T X)^{-1} X^T Y$$

- RANSAC: RANdom SAmpling Consensus
  - Randomly select a (minimum number of) sample of s data points from S and instantiate the model from this subset
  - Determine the set of data points  $S_i$  which are within a distance threshold t of the model. The set  $S_i$  is the consensus set of samples and defines the inliers of S
  - If the subset is  $S_i$  is greter than some threshold T, re-estimate the model using all the points  $S_i$  and terminate
  - If the size of  $S_i$  is less than T select a new subset and repeat the above
  - After N trials the largest consensus set  $S_i$  is selected and the model is re-estimated using all the points in the subset  $S_i$

Cost:

- RANSAC can be vlnarable of the correct choice if the threshod
- k-NN Regression (non parametric)
  - Similar to the k-NN classifier
  - To regress Y for a given value of X, consider k closest point to X in training data and take the average of the responses

$$f(x) = \frac{1}{k} \sum_{x_i \in N_i} y_i$$

- Larger values of k provide a smoother and less variable fit (lower variance)
- Parametric vs non parametric
  - If the parametric forlm is close to the true form of f, the parametric approach will outperform the non parametric
  - As a general ruen parametric methods will tends of outperform nonparametric when there is a small number of observation per per predictor
  - Interpretability stand point: Linezar regression preferred to KNN if the test MSEs are similar or slightly lower
- Ridge regression
  - Similar la least squares but minimizes different quantity instead os sum of squared

$$RSS + \lambda \sum_{i=1}^{d} w_i^2$$

- The Lasso
  - Similar to ridge regression but with slightly different term

$$RSS + \lambda \sum_{i=1}^{d} |w_i|$$

## Learning as Inference

- Classification: Y is discrete
- Regression: Y is continuous
- $Pr(Y = y) \equiv Pr(y) \leftarrow Prior$
- $Pr(x \mid Y = y) \equiv Pr(x \mid y) \leftarrow \text{Likehood}$
- $Pr(X = x) \equiv Pr(x) \leftarrow \text{Evidence}$
- $Pr(y \mid X = x) = \frac{Pr(X|Y=y)Pr(Y=y)}{Pr(X=x)} \leftarrow Posterior$
- Parametric Inference:
  - Estimate  $\theta$  unsing D
  - Compute  $Pr(y \mid x, \hat{\theta})$  to make inference
  - Learning correspond to estimate  $\theta$

#### Approaches:

- Maximum Likehood (ML) Estimation

- Maximum A Posteriori (MAP) Extimation
- Non-parametric Inference
  - Estimate  $Pr(\theta \mid D)$
  - Compute  $Pr(y \mid x, D)$  from  $Pr(y \mid x, \theta, D)Pr(\theta \mid D)$  by marginalizing out  $\theta$

#### Approaches:

- Bayesian methods
- Assumption: Observation are independent and identically distributed
- Maximum Likehood Estimate

### Other

- Classification is discrete
- Regression is continuous
- Probabilistic learning involves estimating P(x,y) from a Dataset
- Probabilistic learning cannot be used to create generative model
- Naive Bayes classifier is a generative model (generative model learns joint probability distribution)
- Logistic regression is a discriminative model (discriminative model learns conditional probability distribution)
- For MLE we assume all observations are independently and identically ditributed given y
- An artificial neuron generates an output signal based on the integrated wieghted input
- Structural Risk Minimization is selecting a seperating hyperplane such that future data is most likely classified correctly
- Boosting is: Each training example has a weight which is re-weight through iterations
- Covariance matrix of given vectors is a useful tool to apply the maximum variance in PCA
- Expectation Maximisation : Algarithm to learn with latent variable
- Posterior probability: Conditional probability taking into account the evidence
- RANSAC : Robust method to fit a model to data with outliers
- Dropout : A method for preventing artificial neural networks from overfitting
- The Lasso: An approach to regression that results in feature selection
- Bagging: Booststrpa aggregating
- Error back propagation: Algorithm to train artificial neural networks
- k-fold cross validation : A technique for assessing a model while exploiting available data for training and testing
- In DEcision Forests, randomness is used in feature selection at each node and in generating bootstrap reoplicas
- In PCA and subspace method, x should belong to the class where the projection lengh to the corresponding subspaces is maximised
- Maximum likehood maximizes the probability of the observation conditioned on the class
- Naive Bayes assumes all D dimensions of an obervation are conditionally independent given Y

- perceptron learning stop modifying weights when all training data is correctly classified
- The kernel function correspond to the scalar product between two data points transformed into a higher dimensional space
- Adaboost algorithm : A weight is given to each training sample, and it is iteratively updated
- The main purpose of PCA is to reduce the effective number of variable