### **Nonparametric Methods**

Introduction à l'apprentissage automatique – GIF-4101 / GIF-7005

Professor: Christian Gagné

Week 4



# 4.1 Histogram estimation

### Nonparametric methods

- Parametric methods (including mixture models)
  - ullet Probability densities  $(p(\mathbf{x}))$  selected in advance (typically,  $\mathbf{x} \sim \mathcal{N}_D(oldsymbol{\mu}, oldsymbol{\Sigma}))$
  - Search for the parameterization of these densities
- Nonparametric methods
  - Estimate the probability density directly from the data
  - No hypothesis a priori on the distribution of data
- Main approaches
  - Histogram estimation
  - Kernel density estimation
  - *k*-nearest neighbours (*k*-NN)

### Nonparametric density estimation

- ullet Probability that value x is less than or equal to a
  - $P(x \le a) = \int_{x=-\infty}^{a} p(x) dx$
  - Estimation by sampling  $\{x^t\}_{t=1}^N$ :  $\hat{P}(x \le a) = \frac{\#\{x^t \le a\}}{N}$
- Estimated value x in the range [a, a + h]

$$\hat{P}(a \le x \le (a+h)) = \frac{\#\{x^t \le (a+h)\} - \#\{x^t \le a\}}{N}$$

• Approximation of density p(x) in [a,a+h] by constant value  $\hat{p}(x|x \in [a,(a+h)]) \approx \hat{p}(a)$ 

$$\hat{P}(a \le x \le (a+h)) = \int_{x=a}^{a+h} \hat{p}(x) \ dx \approx \hat{p}(a)(a+h-a) = h\hat{p}(a)$$

$$\hat{p}(x|x \in [a,(a+h)]) \approx \frac{1}{h} \left[ \frac{\#\{x^t \le (a+h)\} - \#\{x^t \le a\}}{N} \right]$$

### Histogram estimation

- Histogram estimation (1D)
  - Divide the input space into compartments of equal size (bins)
  - Each bin is h wide and positioned with respect to an origin  $x_0$

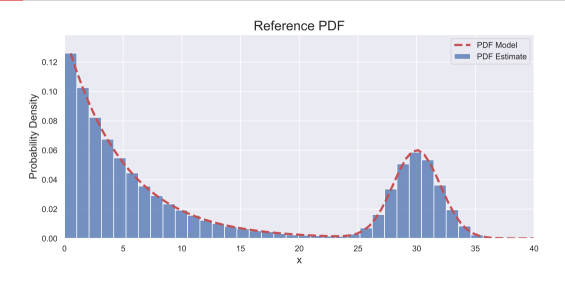
$$]x_0 + mh, x_0 + (m+1)h]$$
, with  $m$  n natural number

• Estimation in 1D, from a set  $\{x^t\}_{t=1}^N$ 

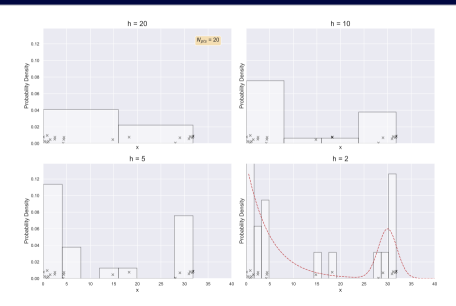
$$\hat{p}(x) = \frac{\#\{x^t \text{ in the same bin than } x\}}{Nh}$$

- Choice of origin  $x_0$  may slightly affect the estimator (boundary discontinuities)
- Choice of width h significantly affects the estimator
  - If the value of h is low, many peaks in the estimate
  - If the value of h is high, softer (less accurate) estimate

# Histogram density estimation



# Histogram density estimation



## **Estimation in many dimensions**

- Histogram estimation in many dimensions
  - Bins corresponding to equal hypervolume hypercubes
  - Highly impacted by the curse of dimensionality
- General conditions for estimates to converge to the true probability density,

$$\hat{p}(\mathbf{x}) 
ightarrow p(\mathbf{x})$$

• Volume  $V_n$  of each bin reduced

$$\lim_{n\to\infty}V_n=0$$

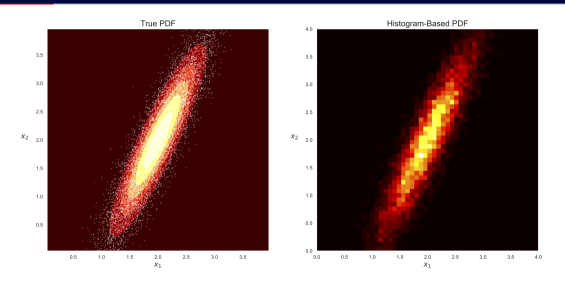
• Number of observations  $k_n$  per bin is very high

$$\lim_{n\to\infty}k_n=\infty$$

• Ratio of the number of observations per bin to total number of observations is high

$$\lim_{n\to\infty}\frac{k_n}{n}=0$$

# 2D density estimations



### Naive histogram density estimation

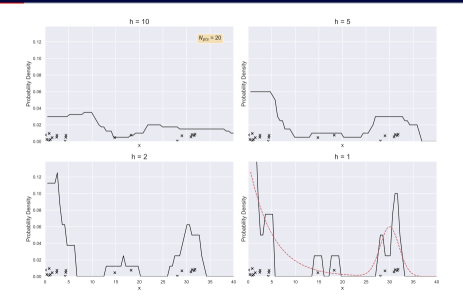
- Naive histogram estimator (also known as a Parzen window)
  - Estimate the density around x in a hypercube of width 2h
  - Formulation in 1D

$$\hat{p}(x) = \frac{\#\{(x-h) < x^t \le (x+h)\}}{2Nh}$$

$$= \frac{1}{2Nh} \sum_{t=1}^{N} w\left(\frac{x-x^t}{h}\right)$$
where  $w(u) = \begin{cases} 1 & \text{if } |u| < 1\\ 0 & \text{otherwise} \end{cases}$ 

- Removes the origin  $x_0$
- The estimation is not continuous and has steps at  $x^t \pm h$

# Naive histogram density estimation



# 4.2 Kernel density estimation

### Kernel density estimation

- Kernel density estimation: softer estimation than the naive histogram estimator
  - Use a softening kernel, typically a Gaussian kernel

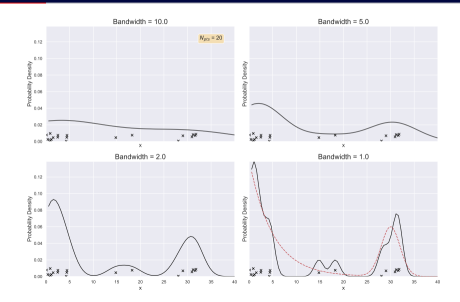
$$K(u) = \frac{1}{\sqrt{2\pi}} \exp\left[-\frac{u^2}{2}\right]$$

• Convolution of the softening kernel with data  $\{x^t\}_{t=1}^N$ 

$$\hat{\rho}(x) = \frac{1}{Nh} \sum_{t=1}^{N} K\left(\frac{x - x^{t}}{h}\right)$$

- Kernel  $K(\cdot)$  determines the shape of influence of the data
- Window width h determines the width of the data influence
- Generalizes the naive estimation, which uses a rectangular box as a kernel

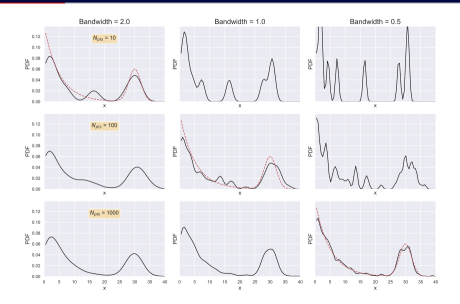
## Kernel density estimation



### Quality of the kernel density estimate

- Window width greatly influences the estimate
  - h small: each data has an important local effect
  - h large: smoother estimation, with overlapping between kernels
- Estimation  $\hat{p}(x) \to p(x)$  when  $N \to \infty$ 
  - ullet h has to o 0, but slower than N (i.e.  $Nh o \infty$ )
  - ullet Typically, we set  $h_N=h_1/\sqrt{N}$ , using a window of  $h_N$  for a dataset of size N

# Varying the number of observations

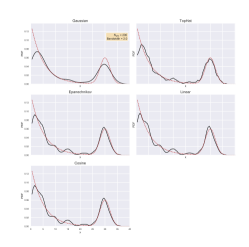


### Properties of softening kernels

- Desirable properties of a softening kernel
  - 1. Positive (or zero) values:  $K(x) \ge 0$ ,  $\forall x$
  - 2. Area under the curve is equal to 1:  $\int_{-\infty}^{\infty} K(x) dx = 1$
  - 3. Centred on the origin:  $\int_{-\infty}^{\infty} x K(x) dx = 0$
- If properties 1 and 2 are respected, K(u) corresponds to a valid density function and therefore  $\hat{p}(x)$  is also valid
- Moreover, if K(u) is continuous and differentiable,  $\hat{p}(x)$  also is
- Support: spreading of u values for which K(u) is non-zero

### **Examples of softening kernels**

- Gaussian
  - Differentiable, but support is not bounded
- Boxcar / TopHat: Naive histogram estimation
  - Bounded support, non-differentiable function
- Epanechnikov:  $K(u) = (3/4)(1 u^2)$  for  $u \in [-1,1]$ 
  - Bounded support, non-derivable function
- Linear / triangle: K(u) = 1 |u| for  $u \in [-1,1]$ 
  - Bounded support, non-derivable function
- Cosinus:  $K(u) = \cos(u \pi/2)$  for  $u \in [-1,1]$ 
  - Bounded support, non-derivable function



#### Kernel estimation, multidimensional case

General equation of the kernel estimation in D dimensions

$$\hat{p}(\mathbf{x}) = \frac{1}{Nh^D} \sum_{t=1}^{N} K\left(\frac{\mathbf{x} - \mathbf{x}^t}{h}\right)$$

- Kernel constraint:  $\int_{\mathbb{R}^D} K(\mathbf{x}) \ d\mathbf{x} = 1$
- Multivariate Gaussian kernel

$$K(\mathbf{u}) = \left(\frac{1}{\sqrt{2\pi}}\right)^D \exp\left[-\frac{\|\mathbf{u}\|^2}{2}\right]$$

- Sensitive to dimensionality and normalization of values in different dimensions
- ullet Kernel including a normalization based on the covariance estimation  $\Sigma$

$$K(\mathbf{u}) = \frac{1}{(2\pi)^{0.5D} |\mathbf{\Sigma}|^{0.5}} \exp\left[-0.5\mathbf{u}^{\top}\mathbf{\Sigma}^{-1}\mathbf{u}\right]$$

#### Kernel estimation for classification

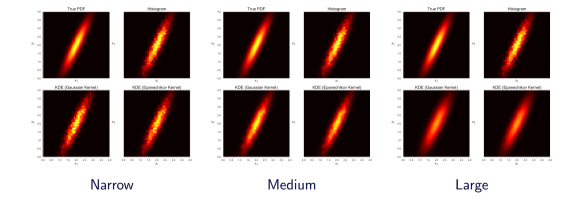
• Kernel estimation of  $\hat{p}(\mathbf{x}|C_i)$ 

$$\hat{p}(\mathbf{x}|C_i) = \frac{1}{N_i h^D} \sum_{t=1}^{N} K\left(\frac{\mathbf{x} - \mathbf{x}^t}{h}\right) r_i^t$$

• Corresponding discriminant function

$$\hat{P}(C_i) = \frac{N_i}{N} 
h_i(\mathbf{x}) = \hat{p}(\mathbf{x}|C_i)\hat{P}(C_i) 
= \frac{1}{Nh^D} \sum_{t=1}^{N} K\left(\frac{\mathbf{x} - \mathbf{x}^t}{h}\right) r_i^t$$

### Kernel width: impact the classification



# 4.3 *k*-nearest neighbours

### *k*-NN density estimation

- *k*-nearest neighbours (*k*-NN)
  - Reference dataset  $\mathcal{X} = \{x^t\}_{t=1}^N$
  - Adapt the window width according to the local data density (k closest data)

$$\hat{p}(x) = \frac{k}{2Nd_k(x,\mathcal{X})}$$

- $h = d_k(x, \mathcal{X})$ : distance from the k-th neighbour to the x data in  $\mathcal{X}$
- Non-continuous estimator, similar to the naive histogram estimator, with adaptive h
  width

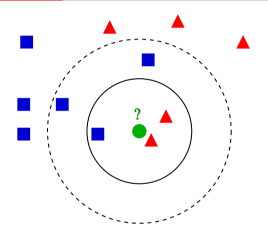
### *k*-NN parameters

- k-NN is defined by three main parameters
  - Number of neighbours *k* 
    - k low: narrow space division based on the reference dataset
    - k high: smoother, larger divisions, average depending on the neighbourhood
  - Distance measurement  $D(\mathbf{x},\mathbf{y})$ 
    - Defines the neighbourhood relationship between the data
  - ullet Reference dataset  ${\mathcal X}$ 
    - Dataset size
    - Density of distribution in the data space
    - Data representativeness (filtering)

#### k-NN classification

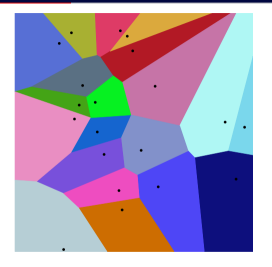
- k-nearest neighbours classification
  - Reference (training) dataset  $\mathcal{X} = \{\mathbf{x}^t, r^t\}_{t=1}^N$
  - To classify an unknown data x, compute the k-closest neighbours in  $\mathcal{X}$  using a distance measure (e.g., Euclidean distance)
  - Assign to x the most frequent label among those of the k-nearest neighbours
- Very simple and direct method for classification
- With k = 1, divide the input space according to a Voronoi diagram based on  $\mathcal{X}$ .

### *k*-NN classification



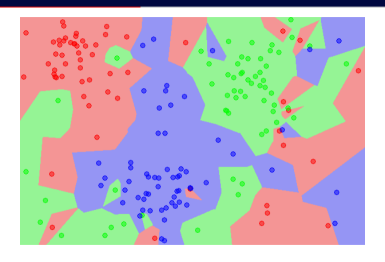
 ${\tt By\ Antti\ Ajanki,\ CC-BY-SA\ 3.0,\ https://en.wikipedia.org/wiki/File:KnnClassification.svg} \\$ 

# Voronoï diagram (1-NN)



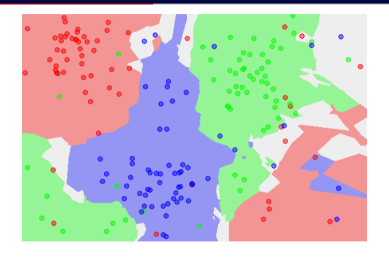
 $By \ Balu.ertl, \ CC-BY-SA\ 4.0, \ https://commons.wikimedia.org/wiki/File: Euclidean\_Voronoi\_diagram.svg$ 

# Regions and borders for 1-NN



By Agor153, CC-BY-SA 3.0, https://en.wikipedia.org/wiki/File:Map1NN.png

# Regions and borders for 5-NN

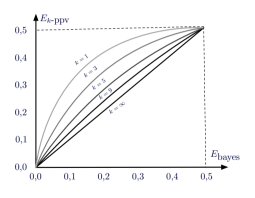


By Agor153, CC-BY-SA 3.0, https://en.wikipedia.org/wiki/File:Map5NN.png

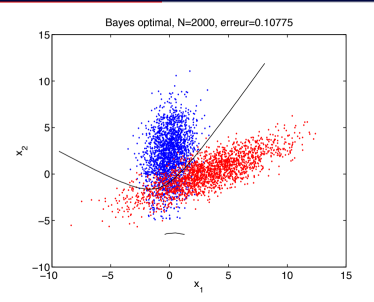
# 4.4 Notions about k-NN

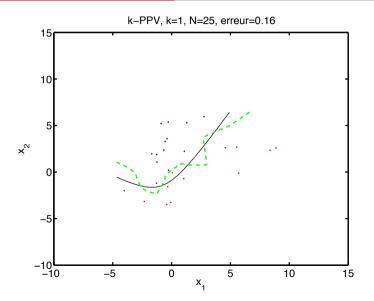
#### Bounds of the *k*-NN classifier

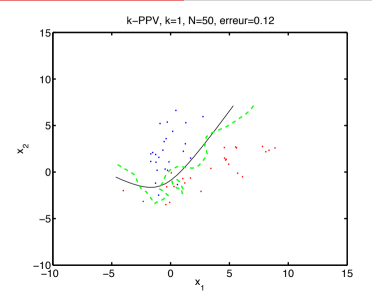
- Optimal Bayesian error rate (E<sub>bayes</sub>)
  - Error rate when true class probability densities are known
  - Optimal, impossible to do better in generalization
- Two bounds on the k-NN error rate
  - With k=1 and  $N \to \infty$  then  $E_{1\text{-ppv}} \le 2E_{\text{baves}}$
  - With  $k \to \infty$  and  $N \to \infty$  then  $E_{k\text{-ppv}} \to E_{\text{baves}}$

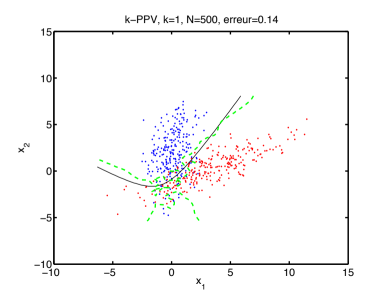


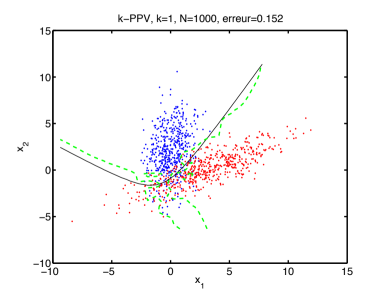
# Optimal Bayesian classification (N = 2000)



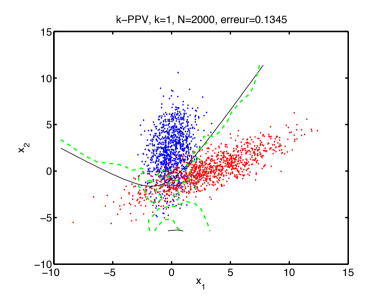




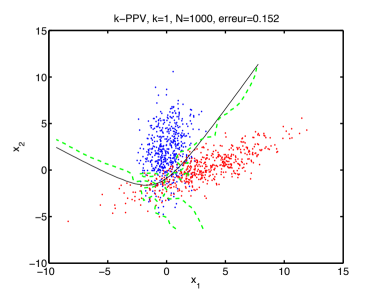




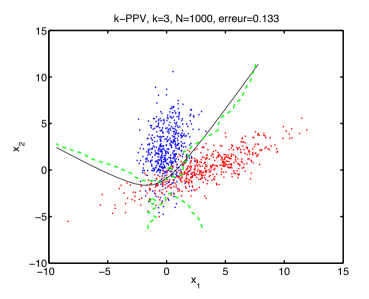
# Varying the number of observations, k = 1, N = 2000



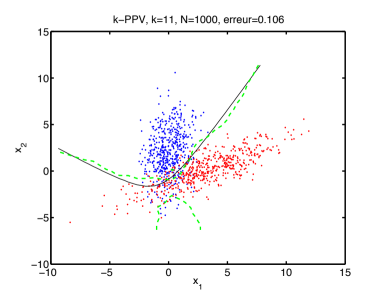
# Varying the number of neighbours, k = 1, N = 1000



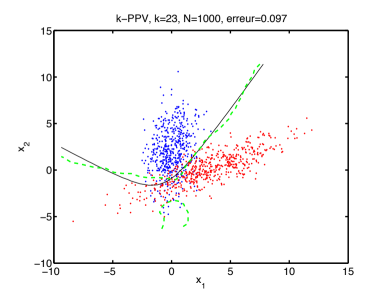
# Varying the number of neighbours, k = 3, N = 1000



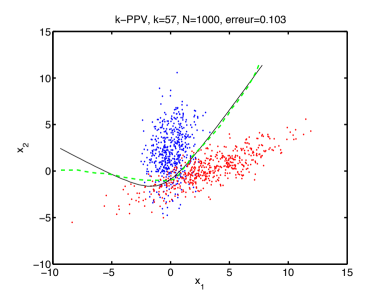
# Varying the number of neighbours, k = 11, N = 1000



# Varying the number of neighbours, k = 23, N = 1000



# Varying the number of neighbours, k = 57, N = 1000



#### **Distances**

- The distance measurement gives the neighbourhood relationship between the data
- Mathematical definition of a metric  $D: X \times X \mapsto \mathbb{R}$ 
  - Non-negativity:  $D(\mathbf{x},\mathbf{y}) \geq 0$
  - Reflexivity:  $D(\mathbf{x},\mathbf{y}) = 0$  iff  $\mathbf{x} = \mathbf{y}$
  - Symmetry:  $D(\mathbf{x},\mathbf{y}) = D(\mathbf{y},\mathbf{x})$
  - Inequality of the triangle:  $D(\mathbf{x},\mathbf{z}) \leq D(\mathbf{x},\mathbf{y}) + D(\mathbf{y},\mathbf{z})$
- Different distance measurements are possible
  - Euclidean distance
  - Minkowsky distance
  - Tanimoto distance (distance between sets)
  - Tangent distance

# Minkowsky distance

Minkowsky distance

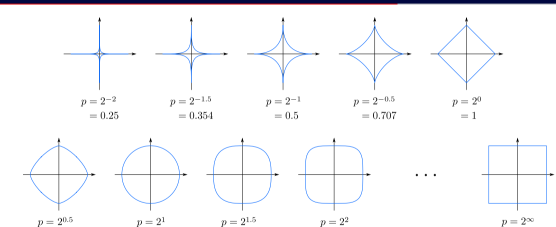
$$\mathrm{D}_p(\mathbf{x},\mathbf{y}) = \left(\sum_{i=1}^D |x_i - y_i|^p\right)^{1/p}$$

- Parameter p controls the emphasis on the dimensions where the magnitude is greatest
- Manhattan distance (p = 1), equal weight for all the dimensions:  $D_1(\mathbf{x}, \mathbf{y}) = \sum_{i=1}^{D} |x_i y_i|$
- Distance  $D_{\infty}$ , using only the dimension where the difference is of maximum magnitude:  $D_{\infty}(\mathbf{x},\mathbf{y}) = \max_{i=1}^{D} |x_i y_i|$
- Euclidean distance (p=2), trade-off between these extremes:  $D_2(\mathbf{x},\mathbf{y}) = \sqrt{\sum_{i=1}^{D} (x_i y_i)^2}$

# Illustration of the Minkowsky distance

=2

= 1.414



By Waldir, CC-BY-SA 3.0, https://commons.wikimedia.org/wiki/File:2D\_unit\_balls.svg

= 4

= 2.828

 $=\infty$ 

#### Data normalization

- Distance measurement sensitive to data scale of all the dimensions
  - Values in a dimension where the scale is large relative to the other dimensions will absorb the value of the other dimensions

$$|x_j - y_j| \gg |x_i - y_i|, \ \forall i \neq j \quad \Rightarrow \quad \mathrm{D}(\mathbf{x}, \mathbf{y}) \approx |x_j - y_j|$$

- Standardization of the data is necessary if the scales are different according to the dimensions
  - Standardization according to the meaning of the data (physical units)
  - Standardization according to max-min value of each dimension
  - Whitening transformation

#### Performance evaluation leave-one-out

- No training required with k-NN
  - ullet Training simply consists in storing the dataset  ${\mathcal X}$
- Leave-one-out performance evaluation
  - Takes advantage of zero cost training
  - Corresponds to K-folds cross validation, with K = N
- 1. For each data  $\mathbf{x}^t \in \mathcal{X}$ :
  - 1.1 Calculate the k-NN of  $\mathbf{x}^t$  among the  $\mathcal{X}\setminus\{\mathbf{x}^t\}$  set
  - 1.2 Determine the most common label of these k closest neighbours as a classification label of  $\mathbf{x}^t$
- 2. For computing the error rate, return the ratio between the number of misclassified data in  $\mathcal X$  and the size of  $\mathcal X$

# 4.5 Computational efficiency of

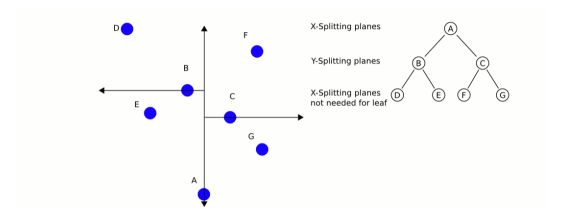
k-NN

# Algorithmic complexity of *k*-NN

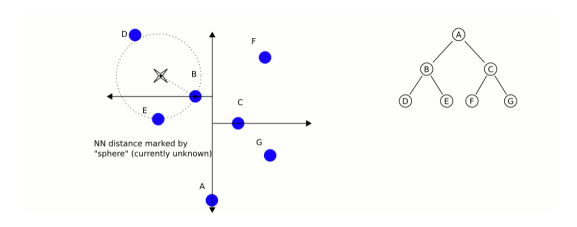
- Training: data storage in memory
  - Complexity in time and memory: O(N)
- Data processing (test/validation): get the *k* neighbours
  - Get the k closest neighbours of  $\mathbf{x}$  in  $\mathcal{X}$ : complexity in time  $O(N \log N)$  (naive algorithm)
  - Classifying M data: complexity in time  $O(MN \log N)$
- It is possible to optimize the calculations by using more sophisticated methods

#### **KD-Tree**

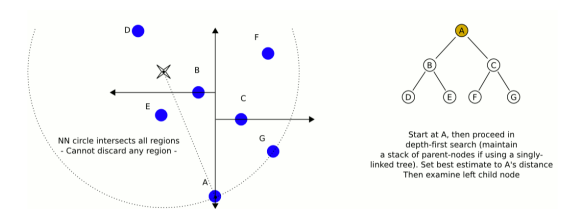
- Structure/topology of data in space can be exploited for the search of the k-NN
  - Avoid calculating the distance with some data, which are anyway too far from the data under test
- KD-tree: tree-like data structure capturing data topology in a Euclidean space
  - Construction of the KD-Tree for N data:  $O(N \log N)$ .
  - Required storage space of KD-Tree: O(N).
  - Querying the k-NN of a data in a KD-Tree
    - $O(N^{\frac{D-1}{D}} + k)$  in D dimensions
    - $O(\sqrt{N}+k)$  in 2D
    - $O(\log N)$  with k=1
  - Processing of M data:  $O(M(N^{\frac{D-1}{D}} + k))$
- Efficient implementations of KD-tree are available (e.g., CGAL in C++, scipy.spatial.KDTree in Python)



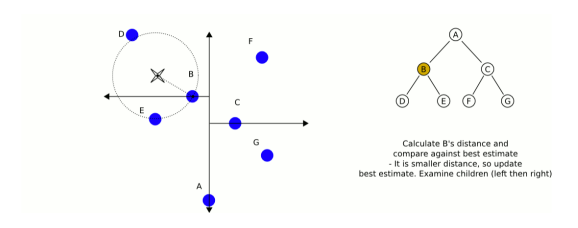
By User\_A1, CC-BY-SA 3.0, https://commons.wikimedia.org/wiki/File:KDTree-animation.gif



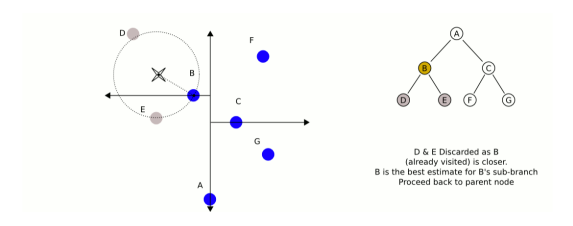
By User\_A1, CC-BY-SA 3.0, https://commons.wikimedia.org/wiki/File:KDTree-animation.gif



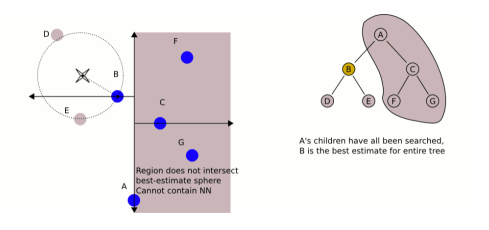
 $By\ User\_A1,\ CC-BY-SA\ 3.0,\ https://commons.wikimedia.org/wiki/File:KDTree-animation.gif$ 



 $By\ User\_A1,\ CC-BY-SA\ 3.0,\ https://commons.wikimedia.org/wiki/File:KDTree-animation.gif$ 



 $By\ User\_A1,\ CC-BY-SA\ 3.0,\ https://commons.wikimedia.org/wiki/File:KDTree-animation.gif$ 



 $By\ User\_A1,\ CC-BY-SA\ 3.0,\ https://commons.wikimedia.org/wiki/File:KDTree-animation.gif$ 

# 4.6 Prototype selection

# Size of training dataset

- Trade-off to make on the size of the training set
  - ullet With  $N o \infty$ , the algorithm tends toward optimal performance
  - ullet But with  $N o\infty$ , processing time and storage needs are huge
- Depending on the position, data density may vary
  - Far from decision boundaries, point density can be reduced
  - Outliers or noisy data in a different class region could be removed
- Approximation of decision boundaries by selecting a few representatives

#### Hart condensation

- Hart condensation
  - ullet Objective: select only  ${\mathcal X}$  data contributing to the classification
  - Heuristics making an incremental construction of the set of prototypes
- Approach
  - Start with an almost empty set of prototypes (a randomly chosen data)
  - Add data only if they are misclassified according to the NN
  - Repeat as long as there are misclassified unselected data

#### Hart condensation

- 1. Create a set of prototypes selected from an  $\mathbf{x}$  data randomly chosen in  $\mathcal{X}$ ,  $\mathcal{Z} = \{\mathbf{x}\}$
- 2. As long as the  $\mathcal{Z}$  set is modified relative to the previous iteration:
  - 2.1 For each data  $\mathbf{x}^t \in \mathcal{X}$ , processed in random order:
    - 2.1.1 Determine the closest neighbour of  $\mathbf{x}^t$  in  $\mathcal{Z}$

$$\mathbf{x}' = \operatorname*{argmin}_{\mathbf{x} \in \mathcal{Z}} \|\mathbf{x}^t - \mathbf{x}\|$$

- 2.1.2 If the class label of  $\mathbf{x}'$  does not match the class label of  $\mathbf{x}^t$   $(r' \neq r^t)$ , then select the data as a prototype,  $\mathcal{Z} = \mathcal{Z} + \{\mathbf{x}^t\}$
- 3. Return the set  ${\mathcal Z}$  as the prototypes selected in  ${\mathcal X}$

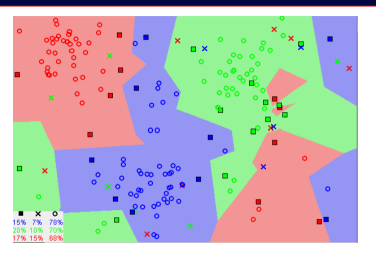
#### Wilson edition

- Wilson edition
  - ullet Heuristics to remove misclassified data from  ${\mathcal X}$  according to leave-one-out
  - Eliminates data that is thought to be aberrant or noisy
- 1. Create the set of prototypes  $\mathcal Z$  from all the data,  $\mathcal Z=\mathcal X$
- 2. For each data  $\mathbf{x}^t \in \mathcal{Z}$ , processed in random order:
  - 2.1 Determine V, which are the k-NN of  $\mathbf{x}^t$  in  $\mathcal{Z}\setminus\{\mathbf{x}^t\}$
  - 2.2 Determine the classification label  $r_{\mathcal{V}}^t$  of  $\mathbf{x}^t$  according to the most common label of the data in  $\mathcal{V}$
  - 2.3 If the  $r_{\mathcal{V}}^t$  label is different from the  $r^t$  label of  $\mathbf{x}^t$ , then remove the data from  $\mathcal{Z}$ ,  $\mathcal{Z} = \mathcal{Z} \setminus \{\mathbf{x}^t\}$
- 3. Return the set  ${\mathcal Z}$  as the prototypes selected in  ${\mathcal X}$

# Other approaches to generate prototypes

- Aggressive selection of prototypes: Wilson's edition followed by Hart's condensation
  - ullet Filter  ${\mathcal X}$  by first eliminating aberrant or noisy data (Wilson edition)
  - Select only the data contributing to the classification (Hart condensation)
- Prototype building
  - ullet Determine prototypes that are not data in  ${\mathcal X}$
  - ullet Possible approach (unsupervised): K-means of  ${\mathcal X}$  with high K value

# Wilson + Hart illustration



 ${\tt By\ Agor153,\ CC-BY-SA\ 3.0,\ https://en.wikipedia.org/wiki/File:Map1NNReducedDataSet.png} \\$ 

 $\times$ : data removed by Wilson (k=3)  $\square$ : prototypes selected by Hart O: data removed by Hart

4.7 Nonparametric methods in

scikit-learn

#### Scikit-learn: density estimation

- neighbors.KernelDensity: kernel density estimation
  - Parameters
    - bandwidth (float): kernel width
    - algorithm (string): neighbourhood algorithm to use, can be 'kd\_tree',
       'ball\_tree' or 'auto' (default: 'auto')
    - kernel (string): noyau utilisé, peut être 'gaussian', 'tophat', 'epanechnikov',
       'exponential', 'linear' ou 'cosine' (default: 'gaussian')
  - Methods
    - fit(X): learn density from data
    - sample(n\_samples=1): generates samples of the distribution (only for Gaussian and tophat kernels)
    - score(X): returns the log-likelihood of the data
    - score\_samples(X): returns the density of data

## Scikit-learn: *k*-nearest neighbours

- neighbors.KneighborsClassifier: classification with the *k*-nearest neighbours method
  - Parameters
    - n\_neighbors (int): number of neighbours used (default: 5)
    - algorithm (string): neighbourhood algorithm to use, can be 'kd\_tree',
       'ball\_tree', 'brute' or 'auto' (default: 'auto')
    - metric (string or object neighbors.DistanceMetric): distance metric used. By default 'minkowski' with p = 2, which returns to a Euclidean distance. For other metrics, see documentation of neighbors.DistanceMetric.
    - p (int): value of p for the Minkowski distance (default: 2)
  - Methods
    - fit(X,y): learn classification model from data
    - kneighbors(X, n\_neighbors): returns the k-nearest neighbours to the data
    - predict(X): does the data classification
- neighbors.KneighborsRegressor: regression by k-nearest neighbours