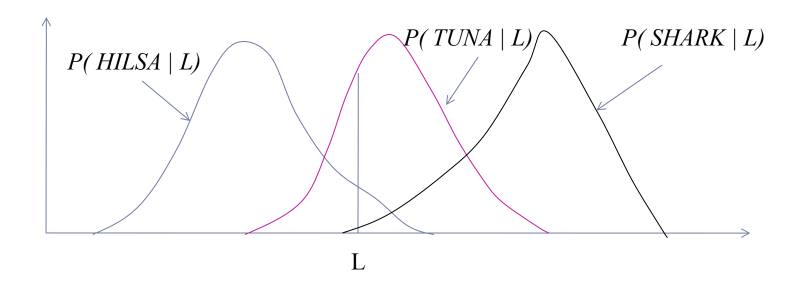




# K Nearest Neighbor Classification

## Bayes Classifier: Recap

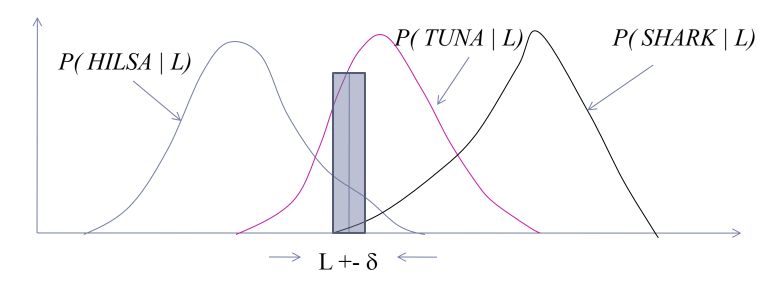


Maximum Aposteriori (MAP) Rule

Distributions assumed to be of particular family (e.g., Gaussian), and parameters estimated from training data.



### Bayes Classifier: Recap



Approximate Maximum Aposteriori (MAP) Rule

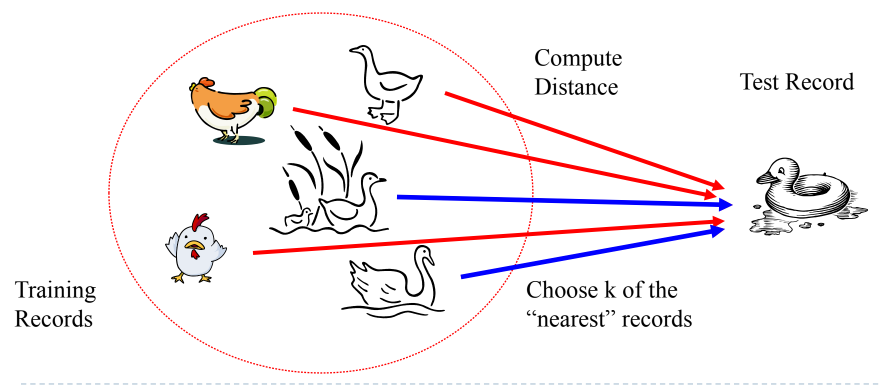
*Non-parametric (data driven) approach*: consider a small window around L, Find which class is most populous in that window.



## Nearest Neighbor Classifiers

#### ► Basic idea:

If it walks like a duck, quacks like a duck, then it's probably a duck



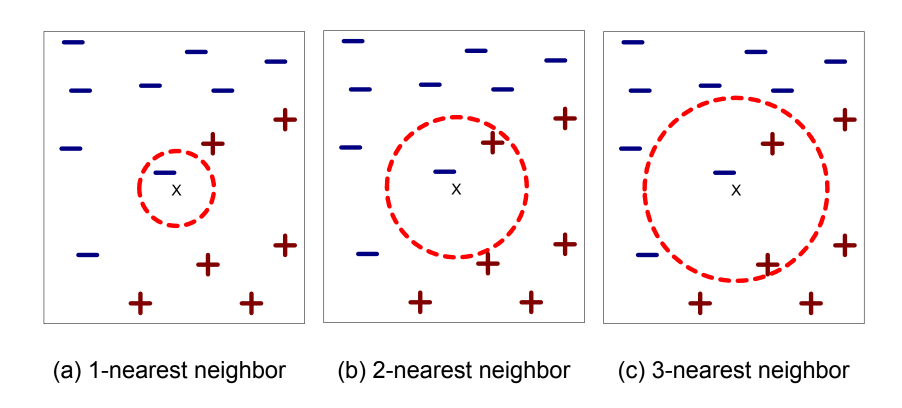


#### Basic Idea

- ▶ *k*-NN classification rule is to assign to a test sample the majority category label of its *k* nearest training samples
- In practice, k is usually chosen to be odd, so as to avoid ties
- The k = 1 rule is generally called the nearest-neighbor classification rule



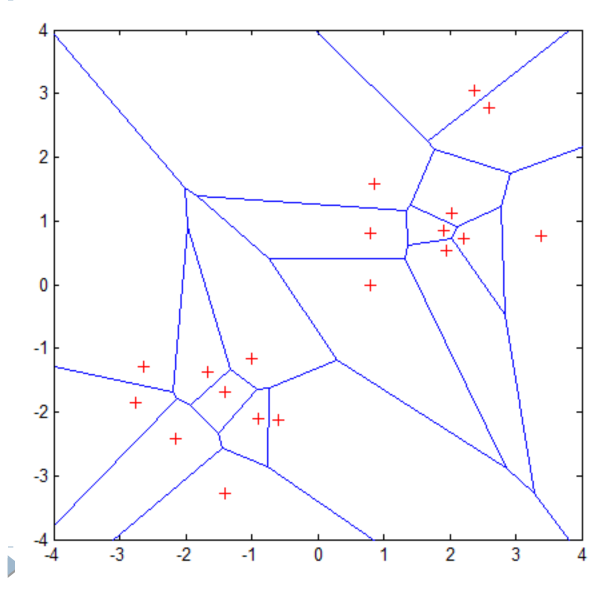
### Definition of Nearest Neighbor



K-nearest neighbors of a record x are data points that have the k smallest distance to x



## Voronoi Diagram



#### Properties:

- All possible points within a sample's Voronoi cell are the nearest neighboring points for that sample
- 2) For any sample, the nearest sample is determined by the closest Voronoi cell edge

## Distance-weighted k-NN

$$\hat{f}(q) = \underset{v \mid V}{\operatorname{argmax}} \stackrel{\hat{a}}{\underset{i=1}{\overset{k}{o}}} \frac{1}{d(x_i, x_q)^2} d(v, f(x_i))$$

General Kernel functions like Parzen Windows may be considered Instead of inverse distance.



### Predicting Continuous Values

$$\hat{f}(q) = rg \max_{v \in V} \mathop{\mathop{\rm all}}_{v} w_i d(v, f(x_i))$$

$$\hat{f}(q) = \mathop{\rm arg max}_{v \in V} \mathop{\mathop{\rm all}}_{v} w_i d(v, f(x_i))$$

$$\hat{f}(q) = \frac{k}{w_i f(x_i)}$$

$$\hat{f}(q) = \frac{i = 1}{k}$$

$$\mathop{\mathop{\rm all}}_{v \in V} w_i$$



### Nearest-Neighbor Classifiers: Issues

- The value of k, the number of nearest neighbors to retrieve
- Choice of Distance Metric to compute distance between records
- Computational complexity
  - Size of training set
  - Dimension of data



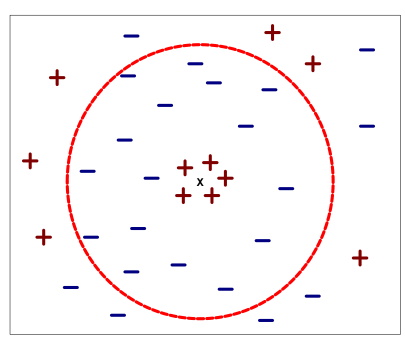
#### Value of K

- Choosing the value of k:
  - If k is too small, sensitive to noise points
  - If k is too large, neighborhood may include points from other classes

Rule of thumb:

K = sqrt(N)

N: number of training points





#### Distance Metrics

Minkowsky:

Manhattan / city-block:

$$D(x,y) = \left(\sum_{i=1}^{m} |x_i - y_i|^r\right)^{\frac{1}{r}} \qquad D(x,y) = \sqrt{\sum_{i=1}^{m} (x_i - y_i)^2} \qquad D(x,y) = \sum_{i=1}^{m} |x_i - y_i|$$

definite  $m \times m$  weight matrix

$$D(x,y) = \sqrt{\sum_{i=1}^{m} (x_i - y_i)}$$

$$D(x,y) = \sum_{i=1}^{m} |x_i - y_i|$$

Camberra:

$$D(x,y) = \sum_{i=1}^{m} \frac{|x_i - y_i|}{|x_i + y_i|}$$

$$D(x,y) = \sum_{i=1}^{m} \frac{|x_i - y_i|}{|x_i + y_i|}$$
 Chebychev: 
$$D(x,y) = \max_{i=1}^{m} |x_i - y_i|$$

Quadratic:

adratic: 
$$D(x,y) = (x - y)^T Q(x - y) = \sum_{j=1}^m \left(\sum_{i=1}^m (x_i - y_i)q_{ji}\right)(x_j - y_j)$$
  
Q is a problem-specific positive

**Mahalanobis:** 

$$D(x, y) = [\det V]^{1/m} (x - y)^{\mathrm{T}} V^{-1} (x - y)$$

V is the covariance matrix of  $A_1..A_m$ , and  $A_i$  is the vector of values for attribute *j* occuring in the training set instances 1..n.

Direlation: 
$$D(x,y) = \frac{\sum_{i=1}^{m} (x_i - \overline{x_i})(y_i - \overline{y_i})}{\sqrt{\sum_{i=1}^{m} (x_i - \overline{x_i})^2 \sum_{i=1}^{m} (y_i - \overline{y_i})^2}}$$

 $\overline{x}_i = \overline{y}_i$  and is the average value for attribute i occuring in the training set.

Chi-square:  $D(x,y) = \sum_{i=1}^{m} \frac{1}{sum_i} \left( \frac{x_i}{size_{ii}} - \frac{y_i}{size_{ii}} \right)^2$ 

sum; is the sum of all values for attribute i occurring in the training set, and  $size_x$  is the sum of all values in the vector x.

sign(x)=-1, 0 or 1 if x < 0,

$$sign(x)=-1, 0 \text{ or } 1 \text{ if } x < 0,$$
  
  $x = 0, \text{ or } x > 0, \text{ respectively.}$ 

Kendall's Rank Correlation:  $D(x,y) = 1 - \frac{2}{n(n-1)} \sum_{i=1}^{m} \sum_{j=1}^{i-1} \operatorname{sign}(x_i - x_j) \operatorname{sign}(y_i - y_j)$ 

#### Distance Measure: Scale Effects

- Different features may have different measurement scales
  - E.g., patient weight in kg (range [50,200]) vs. blood protein values in ng/dL (range [-3,3])
- Consequences
  - Patient weight will have a much greater influence on the distance between samples
  - May bias the performance of the classifier



#### Standardization

Transform raw feature values into z-scores

$$z_{ij} = \frac{x_{ij} - \underline{M}_{j}}{\underline{S}_{j}}$$

- $X_{i}$  is the value for the  $i^{th}$  sample and  $j^{th}$  feature
- $M_j$  is the average of all  $X_{ij}$  for feature j
- $^{S}$  j is the standard deviation of all  $^{X}$ gover all input samples
- Range and scale of z-scores should be similar (providing distributions of raw feature values are alike)

## Nearest Neighbor: Dimensionality

VS

- Problem with Euclidean measure:
  - High dimensional data
    - curse of dimensionality
  - Can produce counter-intuitive results
  - Shrinking density sparsification effect

011111111111

d = 1.4142

1000000000000

 $0\ 0\ 0\ 0\ 0\ 0\ 0\ 0\ 0\ 1$ 

d = 1.4142



#### Distance for Nominal Attributes

### Value Difference Metric (VDM)

[Stanfill & Waltz, 1986]

Providing appropriate distance measurements for nominal attributes.

$$vdm_{a}(x,y) = \sum_{c=1}^{C} \left( \frac{N_{a,x,c}}{N_{a,x}} - \frac{N_{a,y,c}}{N_{a,y}} \right)^{2}$$

 $N_{ax}$  = # times attribute a had value x

 $Na_{,x,c}$  = # times attribute a had value x and class was c C = # output classes

Two values are considered closer if they have more similar classifications, i.e., if they have more similar correlations with the output classes.

## Distance for Heterogeneous Data

In this section, we define a heterogeneous distance function HVDM that returns the distance between two input vectors x and y. It is defined as follows:

$$HVDM(x,y) = \sqrt{\sum_{a=1}^{m} d_a^2(x_a, y_a)}$$
 (11)

where m is the number of attributes. The function  $d_a(x,y)$  returns a distance between the two values x and y for attribute a and is defined as:

$$d_{a}(x,y) = \begin{cases} 1, & \text{if } x \text{ or } y \text{ is unknown; otherwise...} \\ normalized\_vdm_{a}(x,y), & \text{if } a \text{ is nominal} \\ normalized\_diff_{a}(x,y), & \text{if } a \text{ is linear} \end{cases}$$
(12)

Wilson, D. R. and Martinez, T. R., Improved Heterogeneous Distance Functions, Journal of Artificial Intelligence Research, vol. 6, no. 1, pp. 1-34, 1997



# Nearest Neighbour : Computational Complexity

- Expensive
  - To determine the nearest neighbour of a query point q, must compute the distance to all N training examples
    - + Pre-sort training examples into fast data structures (kd-trees)
    - + Compute only an approximate distance (LSH)
    - + Remove redundant data (condensing)
- Storage Requirements
  - Must store all training data P
    - + Remove redundant data (condensing)
    - Pre-sorting often increases the storage requirements
- High Dimensional Data
  - "Curse of Dimensionality"
    - Required amount of training data increases exponentially with dimension
    - Computational cost also increases dramatically
    - Partitioning techniques degrade to linear search in high dimension



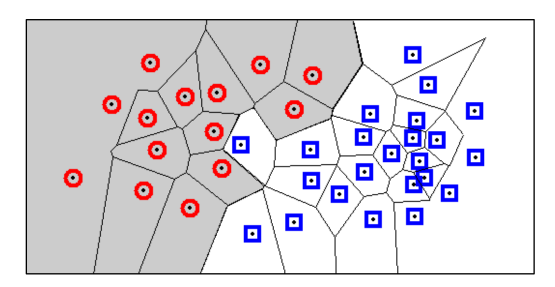
# Reduction in Computational Complexity

- Reduce size of training set
  - Condensation, editing

Use geometric data structure for high dimensional search



### Condensation: Decision Regions



Each cell contains one sample, and every location within the cell is closer to that sample than to any other sample.

A Voronoi diagram divides the space into such cells.

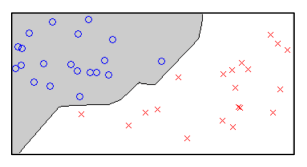
Every query point will be assigned the classification of the sample within that cell. The *decision boundary* separates the class regions based on the 1-NN decision rule.

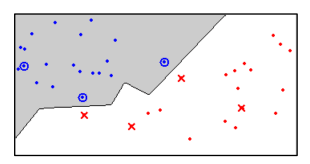
Knowledge of this boundary is sufficient to classify new points.

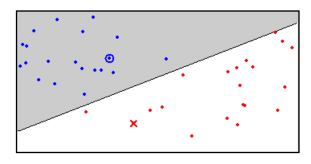
The boundary itself is rarely computed; many algorithms seek to retain only those points necessary to generate an identical boundary.



- Aim is to reduce the number of training samples
- Retain only the samples that are needed to define the decision boundary
- <u>Decision Boundary Consistent</u> a subset whose nearest neighbour decision boundary is identical to the boundary of the entire training set
- ► <u>Minimum Consistent Set</u> the smallest subset of the training data that correctly classifies all of the original training data







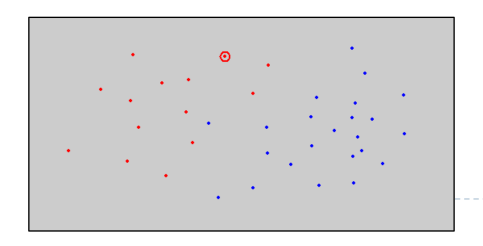
Original data

Condensed data

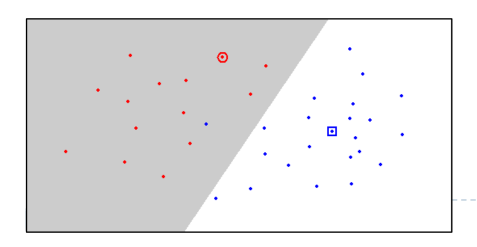
Minimum-Consistent Set

- Condensed Nearest Neighbour (CNN)
  - 1. Initialize subset with a single (or K) training example
  - 2. Classify all remaining samples using the subset, and transfer any incorrectly classified samples to the subset
  - 3. Return to 2 until no transfers occurred or the subset is full

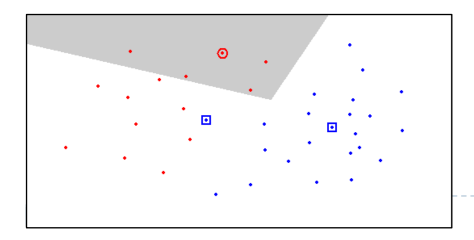
- Incremental
- Order dependent
- Neither minimal nor decision boundary consistent
- O(n³) for brute-force method



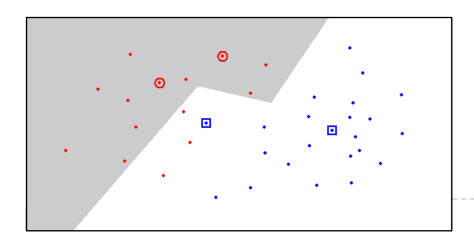
- Condensed Nearest Neighbour (CNN)
  - 1. Initialize subset with a single training example
  - 2. Classify all remaining samples using the subset, and transfer any incorrectly classified samples to the subset
  - 3. Return to 2 until no transfers occurred or the subset is full



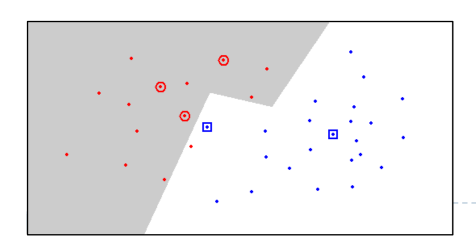
- Condensed Nearest Neighbour (CNN)
  - 1. Initialize subset with a single training example
  - 2. Classify all remaining samples using the subset, and transfer any incorrectly classified samples to the subset
  - 3. Return to 2 until no transfers occurred or the subset is full



- Condensed Nearest Neighbour (CNN)
  - 1. Initialize subset with a single training example
  - 2. Classify all remaining samples using the subset, and transfer any incorrectly classified samples to the subset
  - 3. Return to 2 until no transfers occurred or the subset is full

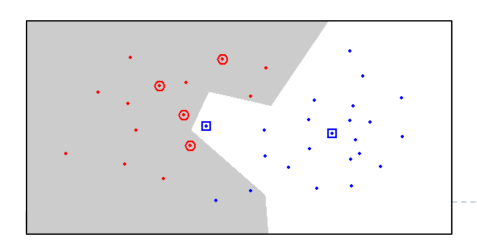


- Condensed Nearest Neighbour (CNN)
  - 1. Initialize subset with a single training example
  - 2. Classify all remaining samples using the subset, and transfer any incorrectly classified samples to the subset
  - 3. Return to 2 until no transfers occurred or the subset is full

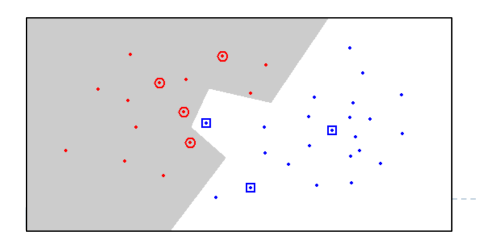


Condensed Nearest Neighbour (CNN)

- 1. Initialize subset with a single training example
- 2. Classify all remaining samples using the subset, and transfer any incorrectly classified samples to the subset
- 3. Return to 2 until no transfers occurred or the subset is full

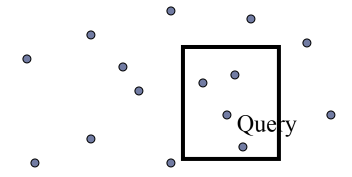


- Condensed Nearest Neighbour (CNN)
  - 1. Initialize subset with a single training example
  - 2. Classify all remaining samples using the subset, and transfer any incorrectly classified samples to the subset
  - 3. Return to 2 until no transfers occurred or the subset is full



## High dimensional search

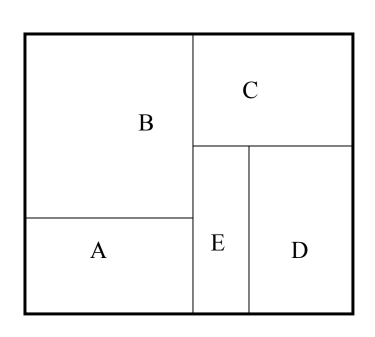
- Given a point set and a nearest neighbor query point
- Find the points enclosed in a rectangle (range) around the query
- Perform linear search for nearest neighbor only in the rectangle

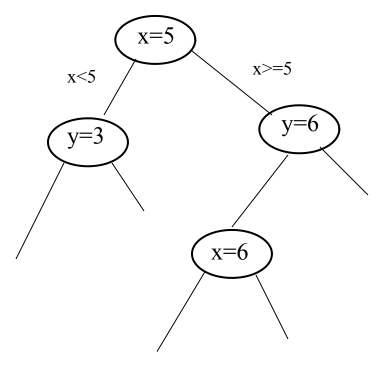




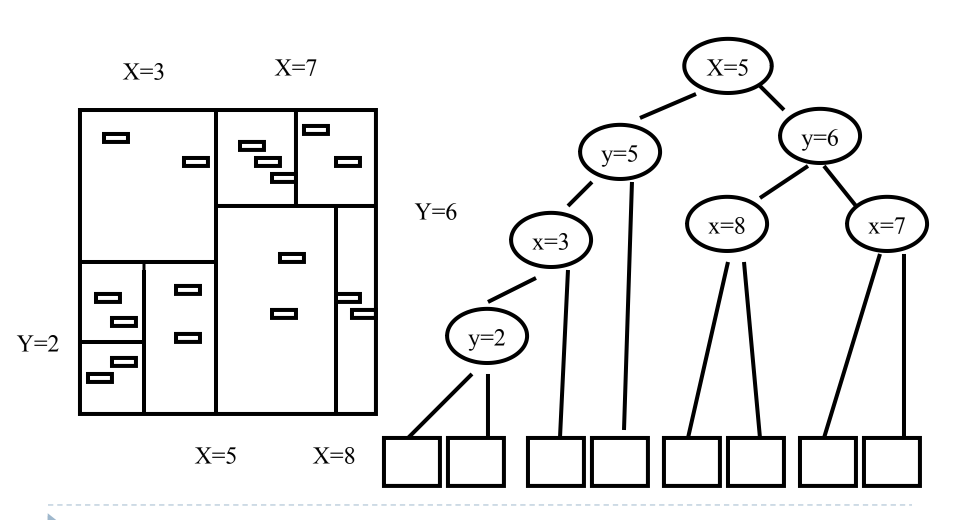
## kd-tree: data structure for range search

- Index data into a tree
- Search on the tree
- Tree construction: At each level we use a different dimension to split





# kd-tree example



## KNN: Alternate Terminologies

- Instance Based Learning
- Lazy Learning
- Case Based Reasoning
- Exemplar Based Learning