# CLUSTERING

# What is Clustering

- Cluster: a collection of data objects
  - Similar to one another within the same cluster
  - Dissimilar to the objects in other clusters
- Cluster analysis
  - Grouping a set of data objects into clusters
- Clustering is unsupervised classification: no predefined classes
- Typical applications
  - As a stand-alone tool to get insight into data distribution
  - As a preprocessing step for other algorithms

# **Examples of Clustering Applications**

- <u>Marketing</u>: Help marketers discover distinct groups in their customer bases, and then use this knowledge to develop targeted marketing programs
- Land use: Identification of areas of similar land use in an earth observation database
- <u>Insurance</u>: Identifying groups of motor insurance policy holders with a high average claim cost
- <u>Urban planning</u>: Identifying groups of houses according to their house type, value, and geographical location
- <u>Seismology</u>: Observed earth quake epicenters should be clustered along continent faults

# What Is a Good Clustering?

- A good clustering method will produce clusters with
  - High intra-class similarity
  - Low inter-class similarity
- Precise definition of clustering quality is difficult
  - Application-dependent
  - Ultimately subjective

### **Similarity and Dissimilarity Between Objects**

• Euclidean distance:

$$d(i,j) = \sqrt{(|x_{i_1} - x_{j_1}|^2 + |x_{i_2} - x_{j_2}|^2 + \dots + |x_{i_p} - x_{j_p}|^2)}$$

- Properties of a metric *d*(*i*,*j*):
  - $d(i,j) \ge 0$
  - d(i,i) = 0
  - d(i,j) = d(j,i)
  - $d(i,j) \le d(i,k) + d(k,j)$

# **Major Clustering Approaches**

- <u>**Partitioning</u>**: Construct various partitions and then evaluate them by some criterion</u>
- <u>Hierarchical</u>: Create a hierarchical decomposition of the set of objects using some criterion
- <u>Model-based</u>: Hypothesize a model for each cluster and find best fit of models to data
- **Density-based**: Guided by connectivity and density functions

# **Partitioning Algorithms**

- **<u>Partitioning method</u>**: Construct a partition of a database *D* of *n* objects into a set of *k* clusters
- Given a *k*, find a partition of *k clusters* that optimizes the chosen partitioning criterion
  - Global optimal: exhaustively enumerate all partitions
  - Heuristic methods: *k-means* and *k-medoids* algorithms
  - <u>k-means</u> (MacQueen, 1967): Each cluster is represented by the center of the cluster
  - <u>*k-medoids*</u> or PAM (Partition around medoids) (Kaufman & Rousseeuw, 1987): Each cluster is represented by one of the objects in the cluster

# **K-Means Clustering**

- Given *k*, the *k*-means algorithm consists of four steps:
  - Select initial centroids at random.
  - Assign each object to the cluster with the nearest centroid.
  - Compute each centroid as the mean of the objects assigned to it.
  - Repeat previous 2 steps until no change.

# **Algorithm Definition**

- The K-Means algorithm is an method to cluster objects based on their attributes into k partitions.
- It assumes that the *k* clusters exhibit Gaussian distributions.
- It assumes that the object attributes form a vector space.
- The objective it tries to achieve is to minimize total intra-cluster variance.

### **Algorithm Fitness Function**

- The K-Means algorithm attempts to minimize the squared error for all elements in all clusters.
- The error equation is:

$$E = \sum_{i=1}^{k} \sum_{p \in C_i} \left| p - m_i \right|^2$$

Where E is the sum of the square error for all elements in the data set;
p is a given element; and m<sub>i</sub> is the mean of cluster C<sub>i</sub>

# **K-Means Algorithm**

#### • Input

- k: the number of clusters
- D: a dataset containing *n* elements
- **Output:** a set of *k* clusters

#### • Method

- (1) arbitrarily choose k elements from D as the initial cluster mean values
- (2) **repeat**
- (3) assign each element to the cluster whose mean the element is *closest* to
- (4) once all of the elements are assigned to clusters, calculate the *actual* cluster means
- (5) **until** there is no change between the new and old cluster means

#### **K-Means Clustering (Example)**



#### **Comments on the K-Means Method**

#### • <u>Strengths</u>

- *Relatively efficient*: O(tkn), where n is # objects, k is # clusters, and t is # iterations. Normally, k, t << n.
- Often terminates at a *local optimum*. The *global optimum* may be found using techniques such as *simulated annealing* and *genetic algorithms*

#### Weaknesses

- Applicable only when *mean* is defined (what about categorical data?)
- Need to specify *k*, the *number* of clusters, in advance
- Trouble with noisy data and *outliers*
- Not suitable to discover clusters with *non-convex shapes*

### K-medoids Clustering

- *K-means* is appropriate when we can work with Euclidean distances
- Thus, *K*-means can work only with numerical, quantitative variable types
- Euclidean distances do not work well in at least two situations
  - Some variables are categorical
  - Outliers can be potential threats
- A general version of **K-means** algorithm called *K-medoids* can work with any distance measure
- *K-medoids* clustering is computationally more intensive

#### K-medoids Algorithm

• Step 1: For a given cluster assignment *C*, find the observation in the cluster minimizing the total distance to other points in that cluster:

$$i_k^* = \underset{\{i:C(i)=k\}}{\operatorname{arg\,min}} \sum_{C(j)=k} d(x_i, x_j).$$

• **Step 2:** Assign 
$$m_k = x_{i_k^*}, k = 1, 2, ..., K$$

• Step 3: Given a set of cluster centers  $\{m_1, ..., m_K\}$ , minimize the total error by assigning each observation to the closest (current) cluster center:  $C(i) = \arg \min d(x, m), i = 1, N$ 

$$L(i) = \arg \min_{1 \le k \le K} d(x_i, m_k), i = 1, ..., N$$

• Iterate steps 1 to 3

### K-medoids Summary

- Generalized *K*-means
- Computationally much costlier that *K*-means
- Apply when dealing with categorical data
- Apply when data points are not available, but only pair-wise distances are available
- Converges to local minimum

# **Hierarchical Clustering**

- Two types: (1) agglomerative (bottom up), (2) divisive (top down)
- **Agglomerative**: two groups are merged if distance between them is less than a threshold
- **Divisive**: one group is split into two if intergroup distance more than a threshold
- Can be expressed by an excellent graphical representation called "dendogram", when the process is monotonic: dissimilarity between merged clusters is increasing. Agglomerative clustering possesses this property. Not all divisive methods possess this monotonicity.
- Heights of nodes in a **dendogram** are proportional to the threshold value that produced them.

#### **An Example Hierarchical Clustering**



# **Hierarchical Clustering**

- Use distance matrix as clustering criteria.
- This method does not require the number of clusters *k* as an input, but needs a termination condition



# **Agglomerative Nesting (Bottom Up)**

- Produces tree of clusters (nodes)
- Initially: each object is a cluster (leaf)
- Recursively merges nodes that have the least dissimilarity
- Criteria: min distance, max distance, avg distance, center distance
- Eventually all nodes belong to the same cluster (root)



# A *Dendrogram* Shows How the Clusters are Merged Hierarchically

- Decompose data objects into several levels of nested partitioning (tree of clusters), called a *dendrogram*.
- A clustering of the data objects is obtained by cutting the *dendrogram* at the desired level. Then each connected component forms a cluster.



# **Divisive Analysis (Top Down)**

- Inverse order of Agglomerative
- Start with root cluster containing all objects
- Recursively divide into subclusters
- Eventually each cluster contains a single object



# **Linkage Functions**

- We know how to measure the distance between two objects, but defining the distance between an object and a cluster, or defining the distance between two clusters is non obvious.
  - Single linkage (nearest neighbor): In this method the distance between two clusters is determined by the distance of the two closest objects (nearest neighbors) in the different clusters.  $d_{SL}(G, H) = \min_{\substack{i \in G \\ j \in H}} d_{ij}$
  - **Complete linkage (furthest neighbor):** In this method, the distances between clusters are determined by the greatest distance between any two objects in the different clusters (i.e., by the "furthest neighbors").  $d_{CL}(G, H) = \max_{\substack{i \in G \\ i \in H}} d_{ij}$
  - Group average linkage: In this method, the distance between two clusters is calculated as the average distance between all pairs of objects in the two different clusters.  $\frac{1}{1}$   $\sum \sum d$

$$d_{GA}(G,H) = \frac{1}{N_G N_H} \sum_{i \in G} \sum_{j \in H} d_{ij}$$

# **Linkage Functions**

- SL considers only a single pair of data points; if this pair is close enough then action is taken. So, SL can form a "chain" by combining relatively far apart data points.
- SL often violates the compactness property of a cluster. SL can produce clusters with large diameters  $(D_G)$ .

$$D_G = \max_{i \in G, \, j \in G} d_{ij}$$

- CL is just the opposite of SL; it produces many clusters with small diameters.
- CL can violate "closeness" property- two close data points may be assigned to different clusters.
- GA is a compromise between SL and CL

# **Linkage Functions**



# **Density-Based Clustering Methods**

- Clustering based on density (local cluster criterion), such as densityconnected points
- Major features:
  - Discover clusters of arbitrary shape
  - Handle noise
  - One scan
  - Need density parameters as termination condition

# Model based clustering

- Assume data generated from K probability distributions
- Typically Gaussian distribution Soft or probabilistic version of K-means clustering
- Need to find distribution parameters.
- EM Algorithm