UNIT-IV

What is Cluster Analysis?

- 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

General Applications of Clustering

- Pattern Recognition
- Spatial Data Analysis
 - create thematic maps in GIS by clustering feature spaces
 - detect spatial clusters and explain them in spatial data mining
- Image Processing
- Economic Science (especially market research)
- WWW
 - Document classification
 - Cluster Weblog data to discover groups of similar access patterns

Examples of Clustering Applications

- Marketing: Help marketers discover distinct groups in their customer bases, and then use this knowledge to develop targeted marketing programs
- <u>Land use</u>: Identification of areas of similar land use in an earth observation database

- Insurance: Identifying groups of motor insurance policy holders with a high average claim cost
- City-planning: Identifying groups of houses according to their house type, value, and geographical location
- Earth-quake studies: Observed earth quake epicenters should be clustered along continent faults

Types of Data in Cluster Analysis

- Data Matrix: This represents n objects, such as persons, with p variables (also called measurements or attributes), such as age, height, weight and so on.
 - The structure is in the form of a relational table, or n-by-p matrix (n objects x p variables). $\begin{bmatrix} x \\ y \\ z \end{bmatrix}$

| ^x 11 | ^x lf | ••• | x_{lp} |
|-----------------------|---------------------|-----|---|
| | | | |
| <i>x</i> <i>i1</i> | ^x if | | x_{ip} |
| | | | |
| x_{n1} | ^x nf | | $\begin{bmatrix} x \\ np \end{bmatrix}$ |

Dissimilarity Matrix: This stores a collection of proximities that are available for all pairs of n objects.

Measure the Quality of Clustering

- Dissimilarity/Similarity metric: Similarity is expressed in terms of a distance function, which is typically metric: d(i, j)
- There is a separate "quality" function that measures the "goodness" of a cluster.
- The definitions of distance functions are usually very different for interval-scaled, boolean, categorical, ordinal and ratio variables.
- Weights should be associated with different variables based on applications and data semantics.

- It is hard to define "similar enough" or "good enough"
 - the answer is typically highly subjective.

Type of data in clustering analysis

- Interval-scaled variables:
- Binary variables:
- Nominal, ordinal, and ratio variables
- Variables of mixed types

Interval-valued variables

- Standardize data $S_f = \frac{1}{n}(|x_{1f} m_f| + |x_{2f} m_f| + ... + |x_{nf} m_f|)$
 - Calculate the mean absolute deviation:

where

$$m_f = \frac{1}{n} (x_{1f} + x_{2f} + \dots + x_{nf})$$

■ Calculate the standardized measurement (*z*-score)

$$z_{if} = \frac{x_{if} - m_f}{s_f}$$

■ Using mean absolute deviation is more robust than using standard deviation

Similarity and Dissimilarity Between Objects

- Distances are normally used to measure the <u>similarity</u> or <u>dissimilarity</u> between two data objects
- Some popular ones include: *Minkowski distance*:

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

where $i = (x_{i1}, x_{i2}, ..., x_{ip})$ and $j = (x_{j1}, x_{j2}, ..., x_{jp})$ are two *p*-dimensional data objects, and *q* is a positive integer

• If q = 1, d is Manhattan distance

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

If $q = 2$, d is Euclidean distance:

$$d(i, j) = \sqrt{(|x_{i1} - x_{j1}|^2 + |x_{i2} - x_{j2}|^2 + ... + |x_{ip} - x_{jp}|^2)}$$

• Properties
• $d(i,j) \ge 0$
• $d(i,i) = 0$
• $d(i,j) = d(j,i)$
• $d(i,j) \le d(i,k) + d(k,j)$

Also one can use weighted distance, parametric Pearson product moment correlation, or other disimilarity measures.

Binary Variables

■ A contingency table for binary data

| | | Object <i>j</i> | | | |
|----------|-----|-----------------|---------------------|-----|--|
| | | 1 | 0 | sum | |
| | 1 | a | b | a+b | |
| Object i | 0 | c | d | c+d | |
| | sum | a+c | <i>b</i> + <i>d</i> | p | |

■ Simple matching coefficient (invariant, if the binary variable is *symmetric*):

$$d(i,j) = \frac{b+c}{a+b+c+d}$$

■ Jaccard coefficient (noninvariant if the binary variable is *asymmetric*):

$$d(i,j) = \frac{b+c}{a+b+c}$$

Dissimilarity between Binary Variables

Example

| Name | Gender | Fever | Cough | Test-1 | Test-2 | Test-3 | Test-4 |
|------|--------|-------|-------|--------|--------|--------|--------|
| Jack | М | Y | Ν | Р | Ν | Ν | Ν |
| Mary | F | Y | Ν | Р | Ν | Р | Ν |
| Jim | Μ | Y | Р | Ν | Ν | Ν | Ν |

- gender is a symmetric attribute
- the remaining attributes are asymmetric binary
- let the values Y and P be set to 1, and the value N be set to 0

$$d(jack, mary) = \frac{0+1}{2+0+1} = 0.33$$
$$d(jack, jim) = \frac{1+1}{1+1+1} = 0.67$$
$$d(jim, mary) = \frac{1+2}{1+1+2} = 0.75$$

Nominal Variables

- A generalization of the binary variable in that it can take more than 2 states, e.g., red, yellow, blue, green
- Method 1: Simple matching
 - m: # of matches, p: total # of variables

$$d(i,j) = \frac{p-m}{p}$$

- Method 2: use a large number of binary variables
 - creating a new binary variable for each of the *M* nominal states

Ordinal Variables

- An ordinal variable can be discrete or continuous
- order is important, e.g., rank

$$r_{if} \in \{1, \dots, M_f\}$$

- Can be treated like interval-scaled
 - replacing x_{if} by their rank
 - map the range of each variable onto [0, 1] by replacing *i*-th object in the *f*-th variable by

$$z_{if} = \frac{r_{if} - 1}{M_f - 1}$$

• compute the dissimilarity using methods for interval-scaled variables

Ratio-Scaled Variables

■ <u>Ratio-scaled variable</u>: a positive measurement on a nonlinear scale, approximately at exponential scale, such as Ae^{Bt} or Ae^{-Bt}

■ Methods:

- treat them like interval-scaled variables not a good choice! (why?)
- apply logarithmic transformation

 $y_{if} = log(x_{if})$

■ treat them as continuous ordinal data treat their rank as interval-scaled.

Variables of Mixed Types

- A database may contain all the six types of variables
 - symmetric binary, asymmetric binary, nominal, ordinal, interval and ratio.
- One may use a weighted formula to combine their effects.

$$d(i,j) = \frac{\sum_{f=1}^{p} \delta_{ij}^{(f)} d_{ij}^{(f)}}{\sum_{f=1}^{p} \delta_{ij}^{(f)}}$$

• f is binary or nominal:

 $d_{ij}{}^{(f)} = 0 \ \ if \ x_{if} = x_{jf} \ , \ or \ d_{ij}{}^{(f)} = 1 \ o.w.$

- f is interval-based: use the normalized distance
- f is ordinal or ratio-scaled

$$Z_{if} = \frac{r_{if} - 1}{M_f - 1}$$

• and treat z_{if} as interval-scaled

A Categorization of Major Clustering Methods

- Partitioning methods: Construct various partitions and then evaluate them by some criterion
- Hierarchy methods: Create a hierarchical decomposition of the set of data (or objects) using some criterion
- <u>Density-based</u>: based on connectivity and density functions
- <u>Grid-based</u>: based on a multiple-level granularity structure
- Model-based: A model is hypothesized for each of the clusters and the idea is to find the best fit of that model to each other

Partitioning Methods

- Partitioning method: 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'67): Each cluster is represented by the center of the cluster
 - <u>k-medoids</u> or PAM (Partition around medoids) (Kaufman & Rousseeuw'87): Each cluster is represented by one of the objects in the cluster

The K-Means Clustering Method

- Given *k*, the *k*-means algorithm is implemented in 4 steps:
 - Partition objects into *k* nonempty subsets
 - Compute seed points as the centroids of the clusters of the current partition. The centroid is the center (mean point) of the cluster.
 - Assign each object to the cluster with the nearest seed point.
 - Go back to Step 2, stop when no more new assignment.

Comments on the K-Means Method

- <u>Strength</u>
 - Relatively efficient: O(tkn), where n is # objects, k is # clusters, and t is # iterations. Normally, $k, t \ll n$.
 - Often terminates at a *local optimum*. The *global optimum* may be found using techniques such as: *deterministic annealing* and *genetic algorithms*
- Weakness
 - Applicable only when *mean* is defined, then what about categorical data?
 - Need to specify *k*, the *number* of clusters, in advance
 - Unable to handle noisy data and *outliers*
 - Not suitable to discover clusters with *non-convex shapes*

Variations of the *K-Means* Method

- A few variants of the *k*-means which differ in
 - Selection of the initial k means
 - Dissimilarity calculations
 - Strategies to calculate cluster means
- Handling categorical data: *k-modes* (Huang'98)
 - Replacing means of clusters with <u>modes</u>
 - Using new dissimilarity measures to deal with categorical objects
 - Using a <u>frequency</u>-based method to update modes of clusters
 - A mixture of categorical and numerical data: *k-prototype* method

The K-Medoids Clustering Method

- Find *representative* objects, called <u>medoids</u>, in clusters
- *PAM* (Partitioning Around Medoids, 1987)

- starts from an initial set of medoids and iteratively replaces one of the medoids by one of the non-medoids if it improves the total distance of the resulting clustering
- PAM works effectively for small data sets, but does not scale well for large data sets
- CLARA (Kaufmann & Rousseeuw, 1990)
- CLARANS (Ng & Han, 1994): Randomized sampling
- Focusing + spatial data structure (Ester et al., 1995)

PAM (Partitioning Around Medoids) (1987)

- PAM (Kaufman and Rousseeuw, 1987), built in Splus
- Use real object to represent the cluster
 - Select k representative objects arbitrarily
 - For each pair of non-selected object h and selected object i, calculate the total swapping cost TC_{ih}
 - For each pair of i and h,
 - If $TC_{ih} < 0$, *i* is replaced by *h*
 - Then assign each non-selected object to the most similar representative object
 - repeat steps 2-3 until there is no change

CLARA (Clustering Large Applications) (1990)

- *CLARA* (Kaufmann and Rousseeuw in 1990)
 - Built in statistical analysis packages, such as S+
- It draws *multiple samples* of the data set, applies *PAM* on each sample, and gives the best clustering as the output
- <u>Strength</u>: deals with larger data sets than *PAM*
- Weakness:
 - Efficiency depends on the sample size

A good clustering based on samples will not necessarily represent a good clustering of the whole data set if the sample is biased

CLARANS ("Randomized" CLARA)

- *CLARANS* (A Clustering Algorithm based on Randomized Search) (Ng and Han'94)
- CLARANS draws sample of neighbors dynamically
- The clustering process can be presented as searching a graph where every node is a potential solution, that is, a set of *k* medoids
- If the local optimum is found, *CLARANS* starts with new randomly selected node in search for a new local optimum
- It is more efficient and scalable than both *PAM* and *CLARA*
- Focusing techniques and spatial access structures may further improve its performance (Ester et al.'95)

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



AGNES (Agglomerative Nesting)

- Introduced in Kaufmann and Rousseeuw (1990)
- Implemented in statistical analysis packages, e.g., Splus

- Use the Single-Link method and the dissimilarity matrix.
- Merge nodes that have the least dissimilarity
- Go on in a non-descending fashion
- Eventually all nodes belong to the same cluster

DIANA (Divisive Analysis)

- Introduced in Kaufmann and Rousseeuw (1990)
- Implemented in statistical analysis packages, e.g., Splus
- Inverse order of AGNES
- Eventually each node forms a cluster on its own

More on Hierarchical Clustering Methods

- Major weakness of agglomerative clustering methods
 - <u>do not scale</u> well: time complexity of at least $O(n^2)$, where *n* is the number of total objects
 - can never undo what was done previously
- Integration of hierarchical with distance-based clustering
 - BIRCH (1996): uses CF-tree and incrementally adjusts the quality of sub-clusters
 - <u>CURE (1998</u>): selects well-scattered points from the cluster and then shrinks them towards the center of the cluster by a specified fraction
 - <u>CHAMELEON (1999)</u>: hierarchical clustering using dynamic modeling

BIRCH:-

- Birch: Balanced Iterative Reducing and Clustering using Hierarchies, by Zhang, Ramakrishnan, Livny (SIGMOD'96)
- Incrementally construct a CF (Clustering Feature) tree, a hierarchical data structure for multiphase clustering
 - Phase 1: scan DB to build an initial in-memory CF tree (a multi-level compression of the data that tries to preserve the inherent clustering structure of the data)

- Phase 2: use an arbitrary clustering algorithm to cluster the leaf nodes of the CFtree
- *Scales linearly*: finds a good clustering with a single scan and improves the quality with a few additional scans

Weakness: handles only numeric data, and sensitive to the order of the data record

Clustering Feature Vector

Clustering Feature: CF = (N, LS, SS)

N: Number of data points

 $LS: \sum_{i=1}^{N} = \overline{X_i}$

SS: $\sum_{i=1}^{N} = \overline{X_i^2}$





Clustering Categorical Data: ROCK

ROCK: Robust Clustering using linKs,

by S. Guha, R. Rastogi, K. Shim (ICDE'99).

Use links to measure similarity/proximity

• Computational complexity: $O(n^2 + nm_m m_a + n^2 \log n)$

■ Basic ideas:

Similarity function and neighbors: $Sim(T_1, T_2) = \frac{|T_1 \cap T_2|}{|T_1 \cup T_2|}$

Let $T_1 = \{1, 2, 3\}, T_2 = \{3, 4, 5\}$

$$Sim(T1, T2) = \frac{|\{3\}|}{|\{1, 2, 3, 4, 5\}|} = \frac{1}{5} = 0.2$$

CHAMELEON

- CHAMELEON: hierarchical clustering using dynamic modeling, by G. Karypis, E.H. Han and V. Kumar'99
- Measures the similarity based on a dynamic model
 - Two clusters are merged only if the *interconnectivity* and *closeness (proximity)* between two clusters are high *relative to* the internal interconnectivity of the clusters and closeness of items within the clusters
- A two phase algorithm
 - I. Use a graph partitioning algorithm: cluster objects into a large number of relatively small sub-clusters
 - 2. Use an agglomerative hierarchical clustering algorithm: find the genuine clusters by repeatedly combining these sub-clusters

Overall Framework of CHAMELEON



Density-Based Clustering Methods

- Clustering based on density (local cluster criterion), such as density-connected points
- Major features:
 - Discover clusters of arbitrary shape
 - Handle noise
 - One scan
 - Need density parameters as termination condition
- Several interesting studies:
 - <u>DBSCAN:</u> Ester, et al. (KDD'96)
 - <u>OPTICS</u>: Ankerst, et al (SIGMOD'99).
 - <u>DENCLUE</u>: Hinneburg & D. Keim (KDD'98)
 - <u>CLIQUE</u>: Agrawal, et al. (SIGMOD'98)
- Two parameters:
 - *Eps*: Maximum radius of the neighbourhood
 - *MinPts*: Minimum number of points in an Eps-neighbourhood of that point
- $N_{Eps}(p)$: {q belongs to $D \mid dist(p,q) \leq Eps$ }
- Directly density-reachable: A point p is directly density-reachable from a point q wrt. Eps, MinPts if
 - 1) p belongs to $N_{Eps}(q)$
 - 2) core point condition:

 $|N_{Eps}(q)| \ge MinPts$



Density-reachable:

A point *p* is density-reachable from a point *q* wrt. *Eps*, *MinPts* if there is a chain of points $p_1, ..., p_n, p_1 = q, p_n = p$ such that p_{i+1} is directly density-reachable from p_i



Density-connected

■ A point *p* is density-connected to a point *q* wrt. *Eps*, *MinPts* if there is a point *o* such that both, *p* and *q* are density-reachable from *o* wrt. *Eps* and *MinPts*.



DBSCAN: Density Based Spatial Clustering of Applications with Noise

- Relies on a *density-based* notion of cluster: A *cluster* is defined as a maximal set of density-connected points
- Discovers clusters of arbitrary shape in spatial databases with noise



OPTICS: A Cluster-Ordering Method

- OPTICS: Ordering Points To Identify the Clustering Structure
 - Ankerst, Breunig, Kriegel, and Sander (SIGMOD'99)
 - Produces a special order of the database wrt its density-based clustering structure
 - This cluster-ordering contains info equiv to the density-based clusterings corresponding to a broad range of parameter settings
 - Good for both automatic and interactive cluster analysis, including finding intrinsic clustering structure
 - Can be represented graphically or using visualization techniques



Grid-Based Clustering Method

- Using multi-resolution grid data structure
- Several interesting methods
 - STING (a STatistical INformation Grid approach) by Wang, Yang and Muntz (1997)
 - WaveCluster by Sheikholeslami, Chatterjee, and Zhang (VLDB'98)
 - A multi-resolution clustering approach using wavelet method
 - CLIQUE: Agrawal, et al. (SIGMOD'98)

STING: A Statistical Information Grid Approach

- Wang, Yang and Muntz (VLDB'97)
- The spatial area area is divided into rectangular cells
- There are several levels of cells corresponding to different levels of resolution



- Each cell at a high level is partitioned into a number of smaller cells in the next lower level
- Statistical info of each cell is calculated and stored beforehand and is used to answer queries
- Parameters of higher level cells can be easily calculated from parameters of lower level cell
 - *count, mean, s, min, max*
 - type of distribution—normal, *uniform*, etc.
- Use a top-down approach to answer spatial data queries
- Start from a pre-selected layer—typically with a small number of cells
- For each cell in the current level compute the confidence interval
- Remove the irrelevant cells from further consideration
- When finish examining the current layer, proceed to the next lower level
- Repeat this process until the bottom layer is reached

- Advantages:
 - Query-independent, easy to parallelize, incremental update
 - O(K), where *K* is the number of grid cells at the lowest level
- Disadvantages:
 - All the cluster boundaries are either horizontal or vertical, and no diagonal boundary is detected

WaveCluster:

- Sheikholeslami, Chatterjee, and Zhang (VLDB'98)
- A multi-resolution clustering approach which applies wavelet transform to the feature space
 - A wavelet transform is a signal processing technique that decomposes a signal into different frequency sub-band.
- Both grid-based and density-based
- Input parameters:
 - # of grid cells for each dimension
 - the wavelet, and the # of applications of wavelet transform.



- How to apply wavelet transform to find clusters
 - Summaries the data by imposing a multidimensional grid structure onto data space
 - These multidimensional spatial data objects are represented in a n-dimensional feature space
 - Apply wavelet transform on feature space to find the dense regions in the feature space

Apply wavelet transform multiple times which result in clusters at different scales from fine to coarse



Quantization



Figure 1: A sample 2-dimensional feature space.

Transformation



CLIQUE (Clustering In QUEst)

- Agrawal, Gehrke, Gunopulos, Raghavan (SIGMOD'98).
- Automatically identifying subspaces of a high dimensional data space that allow better clustering than original space
- CLIQUE can be considered as both density-based and grid-based
 - It partitions each dimension into the same number of equal length interval
 - It partitions an m-dimensional data space into non-overlapping rectangular units

- A unit is dense if the fraction of total data points contained in the unit exceeds the input model parameter
- A cluster is a maximal set of connected dense units within a subspace

CLIQUE: The Major Steps

- Partition the data space and find the number of points that lie inside each cell of the partition.
- Identify the subspaces that contain clusters using the Apriori principle
- Identify clusters:
 - Determine dense units in all subspaces of interests
 - Determine connected dense units in all subspaces of interests.
- Generate minimal description for the clusters
 - Determine maximal regions that cover a cluster of connected dense units for each cluster
 - Determination of minimal cover for each cluster

Model-Based Clustering Methods

- Attempt to optimize the fit between the data and some mathematical model
- Statistical and AI approach
 - Conceptual clustering
 - A form of clustering in machine learning
 - Produces a classification scheme for a set of unlabeled objects
 - Finds characteristic description for each concept (class)
 - COBWEB (Fisher'87)
 - A popular a simple method of incremental conceptual learning
 - Creates a hierarchical clustering in the form of a classification tree
 - Each node refers to a concept and contains a probabilistic description of that concept