Data Mining, Lecture 13 Mining Graph Data

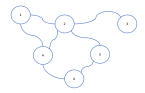
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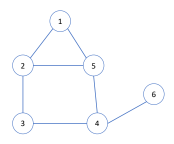
Introduction

- The structure may be more important compared to content.
- Applications: physics, biology, social studies.



Non oriented graph representation

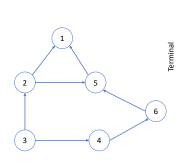
Described by the list or by adjacency matrix



		1	2	3	4	5	6
Ī	1	0	1	0	0	1	0
Ī	2	1	0	1	0	1	0
Ī	3	0	1	0	1	0	0
Ī	4	0	1	1	0	1	1
Ī	5	1	1	0	1	0	0
	6	0	0	0	1	0	0

Oriented graph description

Adjacent matrix is non symmetric.

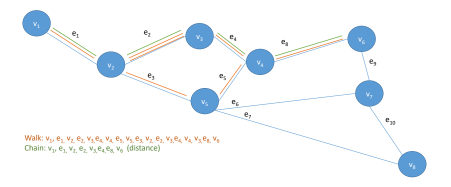


Origin (who)										
	1	2	3	4	5	6				
1	0	1	0	0	1	0				
2	0	0	1	0	0	0				
3	0	0	0	0	0	0				
4	0	0	1	0	0	0				
5	0	1	0	0	0	1				
6	0	0	0	1	0	0				

Path & Walk (chain)

- Walk in the graph G is the sequence $v_0, e_1, v_1, \dots e_l, v_l$, where v_i are nodes (vertexes) and e_i are the ages between the vertexes.
- Vertex v_0 is referred as initial vertex and v_l terminal vertex.
- Path is the walk with no repetitions.
- Vertex v_i is reachable from the vertex v_j if thehere is a walk from v_i to v_j .
- The distance between v_i and v_j is defined as the shortest path between them.

Path & Walk (chain)



Graph database

Definition

- ullet Graph data base ${\mathcal D}$ is defined as the collection of different undirected graphs $G_1 = (N_1, A_1), \dots, G_n = (N_n, A_n).$
 - The set of nodes in ith graph is denoted by N_i and the set of edges by A_i .
 - Each node $p \in N_i$ is associated with the label l(p).

Matching and distance computation

- The term matching is used in two distinct contexts for graph mining.
- Pairing up nodes in a single graph with the use of edges is also referred to as matching.
- Within the frameworks of the present lecture the term matching is used with conjunction to graph matching, the problem is also referred as graph isomorphism.

Matching and distance computation

Definition

Two graphs $G_1=(N_1,A_1)$ and $G_2=(N_2,A_2)$ are said to be isomorphic if there exists a bijection f between the sets of nodes N_1 and N_2 , such that following two conditions are satisfied.

- For each pair of corresponding nodes their labels are the same.
- 2 The edge between the nodes $p_{i,1}$ and $p_{j,1}$ exists in G_1 if and only if the edge exists between the nodes $f(p_{i,2})$ and $f(p_{i,2})$ in G_2 .

Definition

A node induced subgraph of graph G=(N,A) is a graph $G_s=(N_s,A_s)$ satisfying two properties:

- $\mathbf{0}$ $N_s \subseteq N$.
- $A_s = A \cap (N_s \times N_s).$

Matching and distance computation

Definition

A query graph $G_q = (N_q, A_q)$, is said to be a subgraph isomorphism of the data graph G = (N, A) if two following conditions are satisfied:

- For each node $p_i \in N_q$ there is exist a node $p_j \in N$ such that $l(p_i) = l(p_j)$.
- 2 The edge a_{i_1,j_1} , between the nodes $p_{i,1}$ and $p_{j,1}$, exists in G_q if and only if corresponding edge exists in G.

Definition

A Maximal Common Subgraph between a pair of subgraphs $G_1=(N_1,A_1)$ and $G_2=(N_2,A_2)$ is a graph $G_0=(N_0,A_0)$ such that it is a subgraph isomorphism for the both G_1 and G_2 , whereas the power of N_0 is the maximal (of all possible).

Ullmans algorithm may be used to determine all possible subgraph isomorphisms between a query graph and a data graph.

MCG-based distances

NB! Not all of the MCG-based distances satisfy condition to be a metric.

• Unnormalized non-matching measure:

$$U(G_1, G_2) = |G_1| + |G_2| - 2 \cdot |MCS(G_1, G_2)|.$$

• Union-normalized distance:

$$U_n = (G_1, G_2) = 1 - \frac{|MCS(G_1, G_2)|}{|G_1| + |G_2| - MCS(G_1, G_2)}.$$

Max-normalized distance:

$$U_n^{max} = 1 - \frac{|MCS(G_1, G_2)|}{\max\{|G_1|, |G_2|\}}.$$

Edit based distances

Definition

The graph edit distance $E(G_1, G_2)$ it the minimum cost of the edit operations to be applied to G_1 in order to transform it to G_2 .

item Not necessarily symmetric.

Topological descriptors

Topological descriptors convert structural graphs to multidimensional data by using quantitative measures of important structural characteristics as dimensions.

- Morgan index: equal to the number of nodes reachable from the node within a distance of k.
- Wiener index:equal to the sum of the pairwise shortest path distances between all pairs of nodes.

$$W(G) = \sum_{i,j \in G} d(i,j).$$

- Hosoya index: is equal to the number of valid pairwise node-node matchings in the graph.
- Circuit rank: is equal to the minimum number of edges that need to be removed from a graph in order to remove all cycles.

Frequent Substructure Mining in Graphs

The idea of frequent subgraph is identical to the case of association pattern mining, except that a subgraph relationship is used to count the support rather than a subset relationship.

- ullet Let ${\cal G}$ Graph Database, minsup minimum support.
- begin
- $F_1 = \{ \text{ All Frequent singleton graphs } \};$
- k = 1:
- while F_k is not empty do begin
- Generate C_{k+1} by joining pairs of graphs in F_k that share a subgraph of size (k 1) in common;
- Prune subgraphs from C_{k+1} that violate downward closure;
- Determine F_{k+1} by support counting on (C_{k+1}, \mathcal{G}) and retaining subgraphs from C_{k+1} with support at least minsup;
- k = k + 1;
- end:
- return $(\bigcup_{i=1}^k F_i)$;
- end

Graph clustering

- The graph clustering problem partitions a database of n graphs into groups.
- Distance-based methods.
 - k-medoids
 - "community detection" (will be discussed during the next lecture)
- Frequent substructure-based methods.
 - Generic Transformational Approach
 - ▶ XProj: Direct Clustering with Frequent Subgraph Discovery

Graph Classification

- Distance-based methods.
- Frequent substructure-based methods.
 - Generic Transformational Approach
 - XRules: A Rule-Based Approach

Ullmans algorithm

- Let G_q query graph, G data graph, ${\mathcal M}$ currently partially matched node pairs.
- begin
- ullet if $|\mathcal{M}|=|N_q|$ then return successful match \mathcal{M}
- else
- ullet $\mathcal{C}=\mathsf{Set}$ of all label matching node pairs from (G_q,G) not in \mathcal{M}
- (Optional efficiency optimization)
- ullet for each pair $(p_{i_q},p_i)\in\mathcal{C}$ do
- if $\mathcal{M} \cup \{(p_{i_q}, p_i)\}$ is valid partial matching
- then subgraph match $(G_q, G, \mathcal{M} \cup \{(p_{i_q}, p_i)\});$
- end for
- end

Maximum common subgraph algorithm

- Let G_1 and G_2 graphs, \mathcal{M} currently partially matched node pairs, \mathcal{M}_b currently best match .
- begin
- ullet $\mathcal{C}=$ Set of all label matching node pairs from (G_1,G_2) not in \mathcal{M}
- (Optional efficiency optimization)
- for each pair $(p_{i,1}, p_{i,2}) \in \mathcal{C}$ do
- if $\mathcal{M} \cup \{(p_{i,1}, p_{j,2})\}$ is valid matching
- then $\mathcal{M}_b = \text{MCG } (G_1, G_2, \mathcal{M} \cup \{(p_{i,1}, p_{j,2})\});$
- end for
- ullet if $(|\mathcal{M}|>|\mathcal{M}_b|)$ then return \mathcal{M} else return \mathcal{M}_b
- end

Graph matching methods and distance computations

- Pairs of graphs that share large subgraphs in common are likely to be more similar.
- Edit distance.
- Transformation based distance computation.