Efficient Mining of Graph-Based Data

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Abstract
With the increasing amount of structural data being collected, there arises a need to efficiently mine information from this type of data. The goal of this research is to provide a system that performs data mining on structural data represented as a labeled graph. We demonstrate how the graph-based discovery system Subdue can be used to perform structural pattern discovery and structural hierarchical clustering on graph data.

Introduction
The amount of data being generated and collected today is quickly overwhelming researchers’ abilities to interpret the data. In response to this problem, a number of researchers have developed techniques for discovering concepts in databases. Although much of the collected data has an explicit or implicit structural component (e.g., spatial or temporal), few discovery systems are designed to handle this type of data.

In this paper we demonstrate how a structural discovery system, Subdue, can be used to perform efficient mining on structural data. The discovery algorithm implemented in this system is used to perform two types of mining on graph-based data. First, we describe a method of discovering patterns from examples embedded in graph data. Second, we introduce a method of performing hierarchical clustering on structural data using this graph-based discovery system.

Subdue
Subdue (Cook & Holder 2000) is a tool that discovers patterns embedded in labeled graphs. Subdue searches through the space of possible substructure (subgraph) patterns, evaluating each possible substructure pattern based on how well it compresses the input graph in terms of description length.

Once a substructure is discovered, the substructure is used to simplify the data by replacing instances of the substructure with a pointer to the newly discovered substructure. The discovered substructures allow abstraction over detail in the original data. Iteration of the substructure discovery and replacement process constructs a hierarchical description of the structural data in terms of the discovered substructures.

Subdue represents structural data as a labeled graph. Objects in the data map to vertices or small subgraphs in the graph, and relationships between objects map to directed or undirected edges in the graph. This graphical representation serves as input to the substructure discovery system. Subdue has been used to discover patterns in domains such as CAD circuit analysis, chemical compound analysis, and scene analysis. User-defined limits on the number of reported patterns and on the graph match constrain the algorithm to be polynomial in the size of the input graph (Cook, Holder, & Djoko 1996).

Subdue uses a variant of beam search for its main search algorithm (see Figure 1). The goal of the search is to find the subgraph that compresses the input graph the best. A subgraph in Subdue consists of a substructure definition and all its occurrences throughout the graph. The initial state of the search is the set of subgraphs consisting of all uniquely labeled vertices, that is, each subgraph represents one uniquely labeled vertex. The only operator of the search is the Extend Subgraph operator. As its name suggests, it extends a subgraph in all possible ways by a single edge and a vertex, or by a single edge only if both vertices are already in the subgraph. Substructures are evaluated based on their ability to compress the description length of the input graph, following the Minimum Description Length (MDL) principle.

The search progresses by applying the Extend Subgraph operator to each subgraph in the current state. The resulting state, however, does not contain all the subgraphs generated by the Extend Subgraph operator. The subgraphs are kept on a queue and are ordered based on their ability to compress the graph. The length of the queue is partially user specified. The user chooses how many subgraphs of different value, in terms of compression, are to be kept on the queue. Several subgraphs, however, might have the same ability to compress the graph, therefore the actual queue length can vary.
Subdue (graph G, int Beam, int Limit)

queue Q = {v | v has a unique label in G}
bestSub = first substructure in Q

repeat
    newQ = {}
    for each S in Q
        newSubs = S extended by an adjacent edge from G in all possible ways
        newQ = newQ ∪ newSubs
        Limit = Limit - 1
    evaluate substructures in newQ by compression of G
    Q = substructures in newQ with top Beam compression scores
    if best substructure in Q better than bestSub
        then bestSub = best substructure in Q
    until Q is empty or Limit ≠ 0

return bestSub

Figure 1: Subdue’s discovery algorithm.

The search terminates upon reaching a user specified limit on the number of substructures extended, or upon exhaustion of the search space. Once the search terminates and returns the list of best subgraphs, the graph can be actually compressed using the best subgraph. The compression procedure replaces all instances of the subgraph in the input graph by a single vertex, which represents the subgraph. Incoming and outgoing edges to and from the replaced subgraph will point to, or originate in the new vertex that represents the subgraph. The Subdue algorithm can be called again on this compressed graph. This procedure can be repeated a user-specified number of times, and is referred to as an iteration. The maximum number of iterations that can be performed on a graph cannot be predetermined; however, a graph that has been compressed into a single vertex cannot be compressed further.

Subdue’s search is guided by the Minimum Description Length (MDL) principle, originally developed by Rissanen (Rissanen 1989). According to this evaluation heuristic, the best substructure is the one that minimizes the description length of the graph when compressed by the substructure. This compression is calculated as follows:

\[ \text{Compression} = \frac{DL(S) + DL(G \mid S)}{DL(G)} \]

where \( DL(G) \) is the description length of the input graph, \( DL(S) \) is the description length of the subgraph, and \( DL(G \mid S) \) is the description length of the input graph compressed by the subgraph. The search algorithm is looking to maximize the Value of the subgraph which is simply the inverse of the Compression.

Because variations in the data exist, Subdue employs an inexact graph match when finding the instances of a substructure definition. If the cost of transforming the instance to become isomorphic to the substructure definition is less than a user-defined threshold (in the range \([0...1]\)) multiplied by the size of the larger graph, the subgraph is considered as an instance of the substructure.

Because the goal of this research is to apply the substructure discovery algorithm to large, complex databases, the algorithm is constrained by the values of Beam and Limit to run in polynomial time. Parallel and distributed versions of the system have also been developed to provide further scalability (Cook et al. 2000). The distributed algorithm partitions the graph into as many subgraphs as available processors. Each processor searches for subgraphs that compress the local graph well. After broadcasting the best local substructures to all processors and collecting values based on compressing the description length of all graph partitions, the best global substructures are determined. Processors locally expand substructures that are selected to be among the best globally.

Structural Discovery in Earthquake Data

In this study, we demonstrate Subdue’s ability to discovery structural patterns in earthquake activity databases. The earthquake data we analyze contains information from several catalogs (www.neic.cr.usgs.gov), provided by sources like the National Geophysical Data Center of the National Oceanic and Atmospheric Administration. The database has records of earthquakes from 2000 B.C. to present data. An earthquake record consists of 35 fields: source catalog, date, time, latitude, longitude, magnitude, intensity and seismic related information such as cultural effects, isoseismal map, geographic region and stations used for the computations. In this experiment we analyze data covering one year.

In the graph representation of the data, we used two types of edges to connect the events (earthquakes). The first type of edge is the “near_in_distance” edge, which is defined between two events whose distance is at most 75 kilometers. The second type of edge is the “near_in_time” edge that is defined between two events occurring within 36 hours of each other. These parameters were suggested by a professor in the UTA Geology department, Dr. Burke Burkart. An earthquake event in graph form is shown in Figure 2.

Earthquake Data Discovery Results

Our first sampling of data consisted of 10,135 events, comprising a graph with 136,077 vertices and 883,358 edges. The first several discovered substructures linked events with near_in_time and near_in_distance edges. The fourth discovered substructure is more complex and is shown in Figure 3. This substructure is interesting because several earthquakes happened in a short period of time and could be related to a fault placement.
From the discovered results, it was also possible to conclude from the data that most earthquakes occurred in the months of May and June and that a frequent depth for the related earthquakes was 33km and 10km. These depths are validated as frequent depths for earthquake activity. In next experiment, we use Subdue to determine the earthquake activity of a specific area of Mexico. Dr. Burke Burkart, a Geologist at UTA, has extensively studied the Orizaba Fault in Mexico (Burkart 1994; Burkart & Self 1985), whose location is shown in Figure 4.

In processing this data, Subdue discovered not only a subarea with a high concentration of earthquakes, but also some of the area’s characteristics. The first few discovered substructures indicate that region 59, located in the state of Guerrero, has the greatest concentration of earthquakes and that spatial relationships do exist between earthquakes in this region. Dr. Burkart identified this area as very active.

Figure 5 shows substructures discovered using two iterations of Subdue. The second substructure, which is defined in terms of the first substructure, represents a pattern of some of the events at a depth of 33 Km. This is a very interesting pattern, because it might give us information about the cause of those earthquakes. If the earthquake is not caused by subduction (caused by the Pacific plate, which effects depth based on the closeness to the Pacific Ocean), then there is more possibility that it is related to the fault. However, we first have to evaluate and determine the depth of earthquakes caused by subduction in that zone.

In this study we demonstrate that Subdue can be used to discover patterns in structural data containing spatio-temporal relations. In this next section we will describe a different use of the Subdue system, to perform structural hierarchical clustering.

Hierarchical Clustering Using Subdue

Clustering techniques provide a useful means of gaining better understanding of the data, in many cases through revealing hierarchical topologies. Clustering has been applied in diverse fields such as analytical chemistry, geology, biology, zoology and archeology, and is a key component in model fitting, hypothesis generation and testing, data exploration and data reduction. A simple example of hierarchical clustering is the classification of vehicles into groups such as cars, trucks, motorcycles, tricycles, and so on, which are then further subdivided into smaller and smaller groups based on some other traits.

Current clustering techniques have some intrinsic disadvantages. Statistical and syntactic approaches have trouble expressing structural information, and neural approaches are greatly limited in representing semantic information (Schalkoff 1992). Despite these limitations, a number of clustering algorithms have demonstrated success including Cobweb (Fisher 1987), Labyrinth (Thompson & Langley 1991), and AutoClass (Cheeseman & Stutz 1996). Hierarchical approaches to cluster-
ing including agglomerative methods that merge clusters until an optimal separation of clusters is achieved, and divisive approaches that split existing clusters until an optimal clustering is found. These approaches usually have the disadvantage of being applicable only to metric data, which excludes discrete-valued and structured databases (Guha, Rastogi, & Shim 1998; Krypis, Han, & Kumar 1999).

Our Graph-Based Hierarchical Conceptual Clustering algorithm uses Subdue to construct the hierarchical lattice of clusters. The discovered substructure after a single iteration comprises a cluster. The identified cluster is inserted into the classification lattice and used to compress the input graph. The compressed graph is passed again to Subdue to find another substructure. This iteration allows Subdue to find new substructures defined in terms of previously discovered substructures.

Previous work suggested the use of classification trees; however, in structured domains a strict tree is inadequate. In these domains a lattice-like structure may emerge instead of a tree. When substructures are added to the lattice, their parents may include other, non-root nodes in the lattice. If a substructure is composed of two of the same previously-discovered substructures, then there will be two links from the parent to the child in the lattice.

Subdue searches the hypothesis space of all classification lattices. During each iteration of the search process, numerous local minima are encountered, where the global minimum tends to be one of the first few minima. For clustering purposes the first local minimum is used as the best partial hypothesis. The reason for this is easy to see. Subdue starts with all the single-vertex instances of all unique substructures, and iteratively expands the best ones by a single vertex. The local minimum encountered first is therefore caused by a smaller substructure with more instances than the next local minimum, which must be larger, and have fewer instances. A smaller substructure is more general than a larger one, and should represent a parent node in the classification lattice for any more specific clusters. Even though it is entirely possible to use the global minimum as the best substructure, we found that if the global minimum is not the first local minimum it may produce overlapping clusters.

To provide an example of the explanation above, the generation of the hierarchical conceptual clustering for the artificial domain (shown in Figure 6) is demonstrated here. The resulting classification lattice is shown in Figure 7.

Subdue in the first iteration discovers the substructure that describes the pentagon pattern in the input graph. This comprises the first cluster, $C_p$. This cluster is inserted as a child for the root node. In iterations 2 and 3, the square shape (cluster $C_s$) and the triangle shape (cluster $C_t$) are discovered, respectively. These are inserted at the root as well, since $C_s$ does not contain $C_p$ in its definition, $C_t$ does not contain either $C_p$ or $C_s$.

All of the basic shapes (pentagon, square and triangle) appear four times in the input graph. So why is it then that they are discovered in the order described above? Since all of them have the same number of instances in the graph, the size of the subgraph will decide how much they are capable of compressing the input graph. The subgraph describing the pentagon has five vertices and five edges, that of the square has four vertices and four edges, and that of the triangle has three vertices and three edges. Given the same number of instances, the bigger subgraph will compress the input graph better.

In the fourth iteration, Subdue deems the subgraph the best that describes two pentagon shapes connected by a single edge. There are two of these formations in the graph, not four, as one might think, since no overlapping of instances are permitted. This cluster is inserted into the classification lattice as the child of the cluster describing the pentagon, since that cluster appears in its definition. There are two links connecting this new cluster to its parent. This is because the parent cluster definition appears twice.

During iteration five, a subgraph is discovered that contains a pair of squares connected by an edge, a pair of triangles connected by an edge, and these two pair are connected by a single edge. This subgraph has two instances in the input graph. This cluster is inserted as a child for two clusters in the first level of the lattice,
which appear in the definition of this new cluster. The new lattice is depicted in Figure 7. Since both parent cluster definitions appear twice in the new cluster, there are two links from each of those parents to the new node.

**Evaluation Metrics**

Conventional (non-hierarchical) clusterings have been evaluated by the observation which says that instances within a cluster should be similar to each other, and instances in adjacent clusters should be significantly different. The measure suggested by this observation is

\[
ClusteringQuality = \frac{InterClusterDistance}{IntraClusterDistance},
\]

where \( InterClusterDistance \) is the average dissimilarity between members of different clusters, and \( IntraClusterDistance \) is the average dissimilarity between members of the same cluster. The larger the clustering quality, the more distinctly defined the clusters, hence the better the clustering.

In hierarchical clusterings the previously mentioned metric cannot be applied. The main reason for this is that clusters are organized into a hierarchy, and are not completely disjoint. Therefore it does not make sense to compare all of the clusters to all other clusters to compute the average inter-cluster distance. Instead, only clusters that have a common parent may be meaningfully compared.

To develop a good metric for hierarchical conceptual clusterings, first we need to define what characteristics such clusterings should have. One of the things we would like to have is the best coverage by the smallest number of clusters possible. This would imply that clusters are general enough to describe the data while still defining concepts well. In other words, larger clusters provide better generality.

Another desirable property is big cluster description. The more features a cluster has the more its inferential power (Lebowitz 1987). The tradeoff to this is, of course, less coverage. A third thing we would like a clustering to have is minimal or no overlap between its clusters. No overlap indicates disjoint concepts. Clearly defined concepts are of primary importance in conceptual clustering. The above mentioned points can be applied recursively to all clusters of a classification hierarchy. Therefore quality of the root cluster of the hierarchy equals to the quality of the entire hierarchy.

According to our new metric, evaluation of the quality of the clustering \( C (CQ_c) \) is computed by the equation shown in Figure 8. In this equation, \( c \) represents the number of child clusters of \( C \), \( H_i \) represents the \( i \)th child of cluster \( C \), \( H_{i,k} \) represents the \( k \)th instance of the \( i \)th child of cluster \( C \), \( | H_i | \) represents the number of instances of the child cluster \( H_i \), and \( || H_i || \) represents the size of the child cluster definition (number of edges plus the number of vertices). The \( \text{distance} (H_{i,k}, H_{j,l}) \) operation in the numerator is the difference between the two instances of the two child clusters as measured by the number of transformations required to transform the smaller cluster instance into the larger one.

The computation of the quality of a hierarchical clustering is recursive, as expressed by the formula. Because of this recursive nature of the calculation, the quality of the root node of the classification lattice is the quality of the entire clustering.

To compute the quality of a single cluster, all of its child clusters are pairwise compared and normalized. A pairwise comparison between child clusters is performed by comparing each instance of the larger cluster to each instance of the smaller cluster using the inexact graph matching algorithm. The value returned by the inexact graph matching is an integer signifying the number of transformations required to transform one graph into the other. This value is normalized between 0 and 1 by dividing it by the size of the larger graph. The dissimilarity between any two graphs is never greater than the size of the larger graph. In addition, each cluster inherits the quality of its children which are simply added to the total quality.

As suggested by the pairwise comparison of child clusters, this metric measures the dissimilarity of child clusters of a cluster. A larger number signifies a better quality.

This evaluation heuristic rewards the desirable properties of hierarchical clusters. Big clusters are rewarded, since two big disjoint clusters need more transformations to transform one cluster into the other. This is in spite of the normalization, since two 5-vertex and 5-edge clusters having 1 vertex in common are 90% different, while two 2-vertex and 1-edge clusters having one vertex in common are only 66% different.

Disjoint clusters are also rewarded. The less overlap two clusters have, the more distant they are according to the fuzzy matching algorithm, therefore contributing more into the sum of the normalized distances. A small number of clusters is rewarded by taking the average of the comparisons of all the instances, this way offsetting the summing effect, which would normally reward a large number of clusters. As we can see, this evaluation heuristic achieves all of our goals set forth for a hierarchical clustering metric.

**Experimental Results**

In our first experiment, we compare the lattice generated by Subdue with the classification tree produced by Cobweb on an animal database (Fisher 1987). The data used for the experiment is given in Table. The animal data is represented in Subdue as a graph, where attribute names (like \textit{Name} and \textit{BodyCover}) are mapped to labeled edges, and attribute values (like \textit{mammal} and \textit{hair}) are mapped to labeled vertices. The hierarchical clustering generated by Cobweb is shown in Figure 9.

Using the unstructured data, both systems generated similar hierarchies. The lattice generated by Subdue is shown in Figure 10. Using the evaluation measure introduced in the previous section, the clustering generated by Subdue has a quality of 2.6. The classification
\[ CQ_C = \frac{\sum_{i=1}^{c-1} \sum_{j=i+1}^{c} \sum_{k=1}^{c} |H_i| \sum_{j=1}^{|H_j|} \frac{distance(H_{ij}, H_{jkl})}{\max_{jk}(|H_{ij}, H_{jkl}|)}}}{\sum_{i=1}^{c-1} \sum_{j=i+1}^{c} |H_i| + \sum_{i=1}^{c} CQ_{H_i}} \]

Figure 8: New metric for evaluating hierarchical clusters.

Figure 9: Hierarchical clustering over animal descriptions by Cobweb.

Figure 10: Hierarchical clustering over animal descriptions by Subdue.
Table 1: Animal descriptions.

<table>
<thead>
<tr>
<th>Name</th>
<th>Body Cover</th>
<th>Heart Chamber</th>
<th>Body Temp</th>
<th>Fertilization</th>
</tr>
</thead>
<tbody>
<tr>
<td>mammel</td>
<td>hair</td>
<td>four</td>
<td>regulated</td>
<td>internal</td>
</tr>
<tr>
<td>bird</td>
<td>feathers</td>
<td>four</td>
<td>regulated</td>
<td>internal</td>
</tr>
<tr>
<td>reptile</td>
<td>cornified-skin</td>
<td>imperfect-four</td>
<td>unregulated</td>
<td>internal</td>
</tr>
<tr>
<td>amphibian</td>
<td>moist-skin</td>
<td>three</td>
<td>unregulated</td>
<td>external</td>
</tr>
<tr>
<td>fish</td>
<td>scales</td>
<td>two</td>
<td>unregulated</td>
<td>external</td>
</tr>
</tbody>
</table>

Figure 11: Portion of a DNA molecule.

To illustrate the clusters that Subdue generates for structured data, we apply the algorithm to a portion of a DNA data shown in Figure 11. In the input graph, vertices represent atoms and small molecules, and edges represent bonds. A portion of the classification lattice generated by Subdue is shown in Figure 12. As the figure shows, the first level of the lattice represents small commonly-occurring substructures (covering 61% of the data). Subsequently identified clusters are based on these smaller clusters that are combined with each other or with other atoms or molecules to form a new cluster.

Conclusions

The increasing structural component of current databases requires efficient methods for mining structural data. In this paper, we described a graph-based discovery system called Subdue that is capable of mining structural data stored as a labeled graph.

The experiments presented in this paper demonstrate that Subdue can discover interesting and repetitive structural patterns in data. These results were described based on spatio-temporal earthquake data, and ongoing efforts are focusing on other structural databases such as molecular biology data and chemical compound data. In addition, we detail methods of using Subdue to generate structural hierarchical clusters. This algorithm has been demonstrated on animal description data and DNA molecule data. By using the graph-based discovery and mining techniques found in Subdue, structural information can be learned and utilized for a variety of domains and discovery tasks.

References


