Finding Patterns in Three-Dimensional Graphs: Algorithms and Applications to Scientific Data Mining

Xiong Wang, *Member*, *IEEE*, Jason T.L. Wang, *Member*, *IEEE*, Dennis Shasha, Bruce A. Shapiro, Isidore Rigoutsos, *Member*, *IEEE*, and Kaizhong Zhang

Abstract—This paper presents a method for finding patterns in 3D graphs. Each node in a graph is an undecomposable or atomic unit and has a label. Edges are links between the atomic units. Patterns are rigid substructures that may occur in a graph after allowing for an arbitrary number of whole-structure rotations and translations as well as a small number (specified by the user) of edit operations in the patterns or in the graph. (When a pattern appears in a graph only after the graph has been modified, we call that appearance "approximate occurrence.") The edit operations include relabeling a node, deleting a node and inserting a node. The proposed method is based on the geometric hashing technique, which hashes node-triplets of the graphs into a 3D table and compresses the label-triplets in the table. To demonstrate the utility of our algorithms, we discuss two applications of them in scientific data mining. First, we apply the method to locating frequently occurring motifs in two families of proteins pertaining to RNA-directed DNA Polymerase and Thymidylate Synthase and use the motifs to classify the proteins. Then, we apply the method to clustering chemical compounds pertaining to aromatic, bicyclicalkanes, and photosynthesis. Experimental results indicate the good performance of our algorithms and high recall and precision rates for both classification and clustering.

Index Terms—KDD, classification and clustering, data mining, geometric hashing, structural pattern discovery, biochemistry, medicine.

1 Introduction

STRUCTURAL pattern discovery finds many applications in natural sciences, computer-aided design, and image processing [8], [33]. For instance, detecting repeatedly occurring structures in molecules can help biologists to understand functions of the molecules. In these domains, molecules are often represented by 3D graphs. The tertiary structures of proteins, for example, are 3D graphs [5], [9], [18]. As another example, chemical compounds are also 3D graphs [21].

In this paper, we study a pattern discovery problem for graph data. Specifically, we propose a geometric hashing technique to find frequently occurring substructures in a set of 3D graphs. Our study is motivated by recent advances in

- X. Wang is with the Department of Computer Science, California State University, Fullerton, CA 92834. E-mail: wang@ecs.fullerton.edu.
- J.T.L. Wang is with the Department of Computer and Information Science, New Jersey Institute of Technology, University Heights, Newark, NJ 07102. E-mail: jason@cis.njit.edu.
- D. Shasha is with the Courant Institute of Mathematical Sciences, New York University, 251 Mercer St., New York, NY 10012.
 E-mail: shasha@cs.nyu.edu.
- B.A. Shapiro is with the Laboratory of Experimental and Computational Biology, Division of Basic Sciences, National Cancer Institutes, Frederick, MD 21702. E-mail: bshapiro@ncifcrf.gov.
- I. Rigoutsos is with the IBM T.J. Watson Research Center, Yorktown Heights, NY 10598. E-mail: rigoutso@us.ibm.com.
- K. Zhang is with the Department of Computer Science, The University of Western Ontario, London, Ontario, Canada N6A 5B7.
 E-mail: kzhang@csd.uwo.ca.

Manuscript received 19 May 1999; revised 18 Oct. 2000; accepted 18 Jan. 2001; posted to Digital Library 7 Sept. 2001.

For information on obtaining reprints of this article, please send e-mail to: tkde@computer.org, and reference IEEECS Log Number 109849.

the data mining field, where automated discovery of patterns, classification and clustering rules is one of the main tasks. We establish a framework for structural pattern discovery in the graphs and apply our approach to classifying proteins and clustering compounds. While the domains chosen here focus on biochemistry, our approach can be generalized to other applications where graph data occur commonly.

1.1 3D Graphs

Each node of the graphs we are concerned with is an undecomposable or atomic unit and has a 3D coordinate. Each node has a label, which is not necessarily unique in a graph. Node labels are chosen from a domain-dependent alphabet Σ . In chemical compounds, for example, the alphabet includes the names of all atoms. A node can be identified by a unique, user-assigned number in the graph. Edges in the graph are links between the atomic units. In the paper, we will consider 3D graphs that are connected. For disconnected graphs, we consider their connected components [16].

A graph can be divided into one or more *rigid* substructures. A rigid substructure is a subgraph in which the relative positions of the nodes in the substructure are fixed, under some set of conditions of interest. Note that the rigid substructure as a whole can be rotated (we refer to this as a "whole-structure" rotation or simply a rotation when the context is clear). Thus, the relative position of a node in

 $1.\,\mathrm{More}$ precisely, the 3D coordinate indicates the location of the center of the atomic unit.

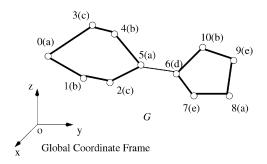


Fig. 1. A graph G.

the substructure and a node outside the substructure can be changed under the rotation. The precise definition of a "substructure" is application dependent. For example, in chemical compounds, a ring is often a rigid substructure.

Example 1. To illustrate rigid substructures in a graph, consider the graph G in Fig. 1. Each node is associated with a unique number, with its label being enclosed in parentheses. Table 1 shows the 3D coordinates of the nodes in the graph with respect to the Global Coordinate Frame. We divide the graph into two rigid substructures: Str_0 and Str_1 . Str_0 consists of nodes numbered 0, 1, 2, 3, 4, and 5 as, well as, edges connecting the nodes (Fig. 2a). Str_1 consists of nodes numbered 6, 7, 8, 9, and 10 as, well as, edges connecting them (Fig. 2b). Edges in the rigid substructures are represented by boldface links. The edge {5,6} connecting the two rigid substructures is represented by a lightface link, meaning that the two substructures are rotatable with respect to each other around the edge.² Note that a rigid substructure is not necessarily complete. For example, in Fig. 2a, there is no edge connecting the node numbered 1 and the node numbered 3.

We attach a local coordinate frame SF_0 (SF_1 , respectively) to substructure Str_0 (Str_1 , respectively). For instance, let us focus on the substructure Str_0 in Fig. 2a. We attach a local coordinate frame to Str_0 whose origin is the node numbered 0. This local coordinate frame is represented by three basis points P_{b_1} , P_{b_2} , and P_{b_3} , with coordinates $P_{b_1}(x_0, y_0, z_0)$, $P_{b_2}(x_0 + 1, y_0, z_0)$, and $P_{b_3}(x_0, y_0 + 1, z_0)$, respectively. The origin is P_{b_1} and the three basis vectors are \vec{V}_{b_1,b_2} , \vec{V}_{b_1,b_3} , and $\vec{V}_{b_1,b_2} \times \vec{V}_{b_1,b_3}$. Here, \vec{V}_{b_1,b_2} represents the vector starting at point P_{b_1} and ending at point P_{b_2} . $\vec{V}_{b_1,b_2} \times$ V_{b_1,b_3} stands for the cross product of the two corresponding vectors. We refer to this coordinate frame as Substructure Frame 0, or SF_0 . Note that the basis vectors of SF_0 are orthonormal. That is, the length of each vector is 1 and the angle between any two basis vectors has 90 degrees. Also note that, for any node numbered i in the substructure Str_0 with global coordinate $P_i(x_i, y_i, z_i)$, we can find a local coordinate of the node i with respect to SF_0 , denoted P'_i , where

$$P_i' = \vec{V}_{b_1,i} = (x_i - x_0, y_i - y_0, z_i - z_0).$$
 (1)

2. Such an edge is referred to as a rotatable edge. If two rigid substructures cannot be rotated with respect to each other around the edge connecting them, that edge is a nonrotatable edge.

TABLE 1
Identifiers, Labels and Global Coordinates of the Nodes of the Graph in Fig. 1

Node identifier	Node label	Node coordinates
0	a	(1.0178, 1.0048, 2.5101)
1	b	(1.2021, 2.0410, 2.0020)
2	c	(1.3960, 2.9864, 2.0006)
3	c	(0.7126, 2.0490, 3.1921)
4	b	(0.7610, 2.7125, 3.0124)
5	a	(1.0097, 3.6478, 2.2660)
6	d	(1.1329, 4.5002, 2.2024)
7	e	(1.5309, 5.2026, 1.7191)
8	a	(1.4529, 6.1015, 1.5712)
9	е	(1.0356, 6.0030, 2.2820)
10	b	(0.7359, 5.0571, 2.6857)

1.2 Patterns in 3D Graphs

We consider a pattern to be a rigid substructure that may occur in a graph after allowing for an arbitrary number of rotations and translations as well as a small number (specified by the user) of edit operations in the pattern or in the graph. We allow three types of edit operations: relabeling a node, deleting a node and inserting a node. Relabeling a node v means to change the label of v to any valid label that differs from its original label. Deleting a node v from a graph means to remove the corresponding atomic unit from the 3D Euclidean space and make the edges touching v connect with one of its neighbors v. Inserting a node v into a graph means to add the corresponding atomic unit to the 3D Euclidean space and make a node v and a subset of its neighbors become the neighbors of v.

We say graph G matches graph G' with n mutations if by applying an arbitrary number of rotations and translations as well as n node insert, delete or relabeling operations, 1) G and G' have the same size, 4 2) the nodes in G geometrically match those in G', i.e., they have coinciding 3D coordinates, and 3) for each pair of geometrically matching nodes, they have the same label. A substructure P approximately occurs in a graph G (or G approximately contains P) within n mutations if P matches some subgraph of G with G mutations or fewer where G is chosen by the user.

Example 2. To illustrate patterns in graphs, consider the set S of three graphs in Fig. 3a. Suppose only exactly coinciding substructures (without mutations) occurring

3. Here, exactly which subset of the neighbors is chosen is unimportant, as the proposed geometric hashing technique hashes nodes only, ignoring edges among the nodes. Notice that when a node v is inserted or deleted, the nodes surrounding v do not move, i.e., their coordinates remain the same. Note also that we do not allow multiple edit operations to be applied to the same node. Thus, for example, inserting a node with label m followed by relabeling it to n is considered as inserting a node with label n. The three edit operations are extensions of the edit operations on sequences; they arise naturally in graph editing [10] and molecule evolution [25]. As shown in Section 4, based on these edit operations, our algorithm finds useful patterns that can be used to classify and cluster 3D molecules effectively.

4. The size of a graph is defined to be the number of nodes in the graph, since our geometric hashing technique considers nodes only. Furthermore, in our target applications, e.g., chemistry, nodes are atomic units and determine the size of a compound. Edges are links between the nodes and have a different meaning from the atomic units; as a consequence, we exclude the edges from the size definition.

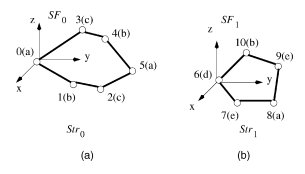


Fig. 2. The rigid substructures of the graph in Fig. 1.

in at least two graphs and having size greater than three are considered as "patterns." Then, \mathcal{S} contains one pattern shown in Fig. 3b. If substructures having size greater than four and approximately occurring in all the three graphs within one mutation (i.e., one node delete, insert, or relabeling is allowed in matching a substructure with a graph) are considered as "patterns," then \mathcal{S} contains one pattern shown in Fig. 3c.

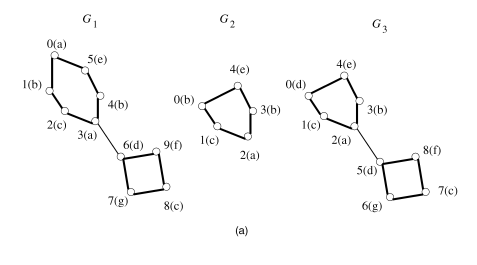
Our strategy to find the patterns in a set of 3D graphs is to decompose the graphs into rigid substructures and, then, use geometric hashing [14] to organize the substructures and, then, to find the frequently occurring ones. In [35], we applied the approach to the discovery of patterns in chemical compounds under a restricted set of edit operations including node insert and node delete, and tested the

quality of the patterns by using them to classify the compounds. Here, we extend the work in [35] by

- considering more general edit operations including node insert, delete, and relabeling,
- presenting the theoretical foundation and evaluating the performance and efficiency of our patternfinding algorithm,
- applying the discovered patterns to classifying 3D proteins, which are much larger and more complicated in topology than chemical compounds, and
- 4. presenting a technique to cluster 3D graphs based on the patterns occurring in them.

Specifically, we conducted two experiments. In the first experiment, we applied the proposed method to locating frequently occurring motifs (substructures) in two families of proteins pertaining to RNA-directed DNA Polymerase and Thymidylate Synthase, and used the motifs to classify the proteins. Experimental results showed that our method achieved a 96.4 percent precision rate. In the second experiment, we applied our pattern-finding algorithm to discovering frequently occurring patterns in chemical compounds chosen from the Merck Index pertaining to aromatic, bicyclicalkanes and photosynthesis. We then used the patterns to cluster the compounds. Experimental results showed that our method achieved 99 percent recall and precision rates.

The rest of the paper is organized as follows: Section 2 presents the theoretical framework of our approach and



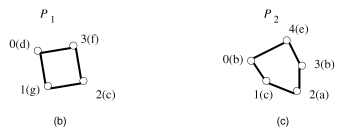


Fig. 3. (a) The set S of three graphs, (b) the pattern exactly occurring in two graphs in S, and (c) the pattern approximately occurring, within one mutation, in all the three graphs.

describes the pattern-finding algorithm in detail. Section 3 evaluates the performance and efficiency of the pattern-finding algorithm. Section 4 describes the applications of our approach to classifying proteins and clustering compounds. Section 5 discusses related work. Section 6 concludes the paper.

2 PATTERN-FINDING ALGORITHM

2.1 Terminology

Let \mathcal{S} be a set of 3D graphs. The occurrence number of a pattern P is the number of graphs in \mathcal{S} that approximately contain P within the allowed number of mutations. Formally, the occurrence number of a pattern P with respect to mutation d and set \mathcal{S} , denoted $occur_no_S^d(P)$, is k if there are k graphs in \mathcal{S} that contain P within d mutations. For example, consider Fig. 3 again. Let \mathcal{S} contain the three graphs in Fig. 3a. Then, $occur_no_S^0(P_1) = 2$; $occur_no_S^1(P_2) = 3$.

Given a set $\mathcal S$ of 3D graphs, our algorithm finds all the patterns P where P approximately occurs in at least Occur graphs in $\mathcal S$ within the allowed number of mutations Mut and $|P| \geq Size$, where |P| represents the size, i.e., the number of nodes, of the pattern P. (Mut, Occur, and Size are user-specified parameters.) One can use the patterns in several ways. For example, biologists or chemists may evaluate whether the patterns are significant; computer scientists may use the patterns to classify or cluster molecules as demonstrated in Section 4.

Our algorithm proceeds in two phases to search for the patterns: 1) find candidate patterns from the graphs in S; and 2) calculate the occurrence numbers of the candidate patterns to determine which of them satisfy the user-specified requirements. We describe each phase in turn below.

2.2 Phase 1 of the Algorithm

In phase 1 of the algorithm, we decompose the graphs into rigid substructures. Dividing a graph into substructures is necessary for two reasons. First, in dealing with some molecules such as chemical compounds in which there may exist two substructures that are rotatable with respect to each other, any graph containing the two substructures is not rigid. As a result, we decompose the graph into substructures having no rotatable components and consider the substructures separately. Second, our algorithm hashes node-triplets within rigid substructures into a 3D table. When a graph as a whole is too large, as in the case of proteins, considering all combinations of three nodes in the graph may become prohibitive. Consequently, decomposing the graph into substructures and hashing node-triplets of the substructures can increase efficiency. For example, consider a graph of 20 nodes. There are

$$\binom{20}{3} = 1140$$
 node-triplets.

On the other hand, if we decompose the graph into five substructures, each having four nodes, then there are only

$$5 \times {4 \choose 3} = 20$$
 node-triplets.

There are several alternative ways to decompose 3D graphs into rigid substructures, depending on the application at hand and the nature of the graphs. For the purposes of exposition, we describe our pattern-finding algorithm based on a partitioning strategy. Our approach assumes a notion of atomic unit which is the lowest level of description in the case of interest. Intuitively, atomic units are fundamental building elements, e.g., atoms in a molecule. Edges arise as bonds between atomic units. We break a graph into maximal size rigid substructures (recall that a rigid substructure is a subgraph in which the relative positions of nodes in the substructure are fixed). To accomplish this, we use an approach similar to [16] that employs a depth-first search algorithm, referred to as DFB, to find blocks in a graph.⁵ The DFB works by traversing the graph in a depth-first order and collecting nodes belonging to a block during the traversal, as illustrated in the example below. Each block is a rigid substructure. We merge two rigid substructures B_1 and B_2 if they are not rotatable with respect to each other, that is, the relative position of a node $n_1 \in B_1$ and a node $n_2 \in B_2$ is fixed. The algorithm maintains a stack, denoted STK, which keeps the rigid substructures being merged. Fig. 4 shows the algorithm, which outputs a set of rigid substructures of a graph G. We then throw away the substructures P where |P| < Size. The remaining substructures constitute the candidate patterns generated from G. This pattern-generation algorithm runs in time linearly proportional to the number of edges in G.

Example 3. We use the graph in Fig. 5 to illustrate how the Find_Rigid_Substructures algorithm in Fig. 4 works. Rotatable edges in the graph are represented by lightface links; nonrotatable edges are represented by boldface links. Initially, the stack STK is empty. We invoke DFB to locate the first block (Step 4). DFB begins by visiting the node numbered 0. Following the depth-first search, DFB then visits the nodes numbered 1, 2, and 5. Next, DFB may visit the node numbered 6 or 4. Without loss of generality, assume DFB visits the node numbered 6 and, then, the nodes numbered 10, 11, 13, 14, 15, 16, 18, and 17, in that order. Then, DFB visits the node numbered 14, and realizes that this node has been visited before. Thus, DFB goes back to the node numbered 17, 18, etc., until it returns to the node numbered 14. At this point, DFB identifies the first block, B_1 , which includes nodes numbered 14, 15, 16, 17, and 18. Since the stack STK is empty now, we push B_1 into STK (Step 7).

In iteration 2, we call DFB again to find the next block (Step 4). DFB returns the nonrotatable edge $\{13,14\}$ as a block, denoted B_2 .⁶ The block is pushed into STK (Step 7). In iteration 3, DFB locates the block B_3 , which includes nodes numbered 10, 11, 12, and 13 (Step 4). Since B_3 and the top entry of the stack, B_2 are not rotatable with respect to each other, we push B_3 into

^{5.} A block is a maximal subgraph that has no cut-vertices. A cut-vertex of a graph is one whose removal results in dividing the graph into multiple, disjointed subgraphs [16].

^{6.} In practice, in graph representations for molecules, one uses different notation to distinguish nonrotatable edges from rotatable edges. For example, in chemical compounds, a double bond is nonrotatable. The different notation helps the algorithm to determine the types of edges.

```
{\bf Procedure}\ {\sf Find\_Rigid\_Substructures}
Input: Graph G.
Output: A set of maximal size rigid substructures generated from G.
1. STK := \emptyset:
   while G is not empty do
3.
        begin
            locate the next block B_1 in G using the DFB algorithm;
5.
            delete B_1 from G;
            /* let the top entry of STK be B_2 */
            {f if} (STK is empty) or (B_1 and B_2 are not rotatable with respect to each other) {f then}
6.
7.
                push the nodes of B_1 into STK;
8.
            else begin
Q
                pop out all nodes in STK, merge them and output the resulting substructure;
10.
                push the nodes of B_1 into STK;
11.
            end:
        end;
12.
13. pop out all nodes in STK, merge them and output the resulting substructure;
```

Fig. 4. Algorithm for finding rigid substructures in a graph.

STK (Step 7). In iteration 4, we continue to call DFB and get the single edge $\{6,10\}$ as the next block, B_4 (Step 4). Since $\{6,10\}$ is rotatable and the stack STK is nonempty, we pop out all nodes in STK, merge them and output the rigid substructure containing nodes numbered 10, 11, 12, 13, 14, 15, 16, 17, 18 (Step 9). We then push B_4 into STK (Step 10).

In iteration 5, DFB visits the node numbered 7 and, then, 8 from which DFB goes back to the node numbered 7. It returns the single edge $\{7,8\}$ as the next block, B_5 (Step 4). Since $\{7, 8\}$ is connected to the current top entry of STK, $\{6, 10\}$, via a rotatable edge $\{6, 7\}$, we pop out $\{6, 10\}$, which itself becomes a rigid substructure (Step 9). We then push $\{7,8\}$ into STK (Step 10). In iteration 6, DFB returns the single edge $\{7,9\}$ as the next block, B_6 (Step 4). Since B_6 and the current top entry of STK, B_5 = {7,8}, are not rotatable with respect to each other, we push B_6 into STK (Step 7). In iteration 7, DFB goes back from the node numbered 7 to the node numbered 6 and returns the single edge $\{6,7\}$ as the next block, B_7 (Step 4). Since $\{6,7\}$ is rotatable, we pop out all nodes in STK, merge them and output the resulting rigid substructure containing nodes numbered 7, 8, and 9 (Step 9). We then push $B_7 = \{6, 7\}$ into STK (Step 10).

In iteration 8, DFB returns the block $B_8 = \{5,6\}$ (Step 4). Since $\{5,6\}$ and $\{6,7\}$ are both rotatable, we pop out $\{6,7\}$ to form a rigid substructure (Step 9). We then push B_8 into STK (Step 10). In iteration 9, DFB returns the block B_9 containing nodes numbered 0, 1, 2, 3, 4, and 5 (Step 4). Since the current top entry of STK, $B_8 = \{5,6\}$,

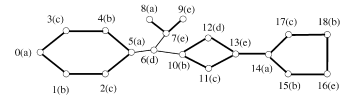


Fig. 5. The graph used for illustrating how the Find_Rigid_Substructures algorithm works.

is rotatable with respect to B_9 , we pop out $\{5,6\}$ to form a rigid substructure (Step 9) and push B_9 into STK (Step 10). Finally, since there is no block left in the graph, we pop out all nodes in B_9 to form a rigid substructure and terminate (Step 13).

2.3 Phase 2 of the Algorithm

Phase 2 of our pattern-finding algorithm consists of two subphases. In subphase A of phase 2, we hash and store the candidate patterns generated from the graphs in phase 1 in a 3D table \mathcal{H} . In subphase B, we rehash each candidate pattern into \mathcal{H} and calculate its occurrence number. Notice that in the subphase B, one does not need to store the candidate patterns in \mathcal{H} again.

In processing a rigid substructure (pattern) of a 3D graph, we choose all three-node combinations, referred to as nodetriplets, in the substructure and hash the node-triplets. We hash three-node combinations, because to fix a rigid substructure in the 3D Euclidean space one needs at least three nodes from the substructure and three nodes are sufficient provided they are not collinear. Notice that the proper order of choosing the nodes i, j, k, in a triplet is significant and has an impact on the accuracy of our approach, as we will show later in the paper. We determine the order of the three nodes by considering the triangle formed by them. The first node chosen always opposes the longest edge of the triangle and the third node chosen opposes the shortest edge. For example, in the triangle in Fig. 6, we choose i, j, k, in that order. Thus, the order is unique if the triangle is not isosceles or equilateral, which usually holds when the coordinates are floating point

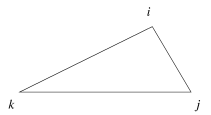


Fig. 6. The triangle formed by the three nodes i, j, k.

TABLE 2 The Node Labels of the Graph in Fig. 1 and Their Indices in the Array $\mathcal A$

index	0	1	2	3	4
label	a	b	С	d	е

numbers. On the other hand, when the triangle is isosceles, we hash and store the two node-triplets [i,j,k] and [i,k,j], assuming that node i opposes the longest edge and edges $\{i,j\}$, $\{i,k\}$ have the same length. When the triangle is equilateral, we hash and store the six node-triplets [i,j,k], [i,k,j], [j,i,k], [j,k,i], [k,i,j], and [k,j,i].

The labels of the nodes in a triplet form a label-triplet, which is encoded as follows: Suppose the three nodes chosen are v_1 , v_2 , v_3 , in that order. We maintain all node labels in the alphabet Σ in an array A. The code for the labels is an unsigned long integer, defined as

$$((L_1 \times Prime + L_2) \times Prime) + L_3,$$

where $Prime > |\Sigma|$ is a prime number, L_1 , L_2 , and L_3 are the indices for the node labels of v_1 , v_2 , and v_3 , respectively, in the array A. Thus, the code of a label-triplet is unique. This simple encoding scheme reduces three label comparisons into one integer comparison.

Example 4. To illustrate how we encode label-triplets, consider again the graph G in Fig. 1. Suppose the node labels are stored in the array \mathcal{A} , as shown in Table 2. Suppose Prime is 1,009. Then, for example, for the three nodes numbered 2, 0, and 1 in Fig. 1, the code for the corresponding label-triplet is $((2 \times 1,009 + 0) \times 1,009) + 1 = 2,036,163$.

2.3.1 Subphase A of Phase 2

In this subphase, we hash the candidate patterns generated in phase 1 of the pattern-finding algorithm into a 3D table. For the purposes of exposition, consider the example substructure Str_0 in Fig. 2a, which is assumed to be a candidate pattern. We choose any three nodes in Str_0 and calculate their 3D hash function values as follows: Suppose the chosen nodes are numbered i, j, k, and have global coordinates $P_i(x_i, y_i, z_i)$, $P_j(x_j, y_j, z_j)$, and $P_k(x_k, y_k, z_k)$, respectively. Let l_1 , l_2 , l_3 be three integers where

$$l_{1} = ((x_{i} - x_{j})^{2} + (y_{i} - y_{j})^{2} + (z_{i} - z_{j})^{2}) \times Scale$$

$$l_{2} = ((x_{i} - x_{k})^{2} + (y_{i} - y_{k})^{2} + (z_{i} - z_{k})^{2}) \times Scale .$$

$$l_{3} = ((x_{k} - x_{j})^{2} + (y_{k} - y_{j})^{2} + (z_{k} - z_{j})^{2}) \times Scale$$
(2)

Here, $Scale = 10^p$ is a multiplier. Intuitively, we round to the nearest pth position following the decimal point (here p is the last accurate position) and, then, multiply the numbers by 10^p . The reason for using the multiplier is that we want some digits following the decimal point to contribute to the distribution of the hash function values. We ignore the digits after the position p because they are inaccurate. (The appropriate value for the multiplier can be calculated once data are given, as we will show in Section 3.)

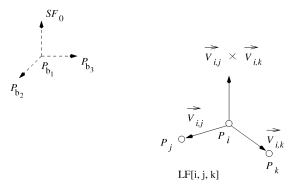


Fig. 7. Calculation of the coordinates of the basis points P_{b_1} , P_{b_2} , P_{b_3} of Substructure Frame 0 (SF_0) with respect to the local coordinate frame LF[i,j,k].

Let

$$d_1 = (l_1 + l_2) \mod Prime_1 \mod Nrow$$

 $d_2 = (l_2 + l_3) \mod Prime_2 \mod Nrow$.
 $d_3 = (l_3 + l_1) \mod Prime_3 \mod Nrow$

 $Prime_1$, $Prime_2$, and $Prime_3$ are three prime numbers and Nrow is the cardinality of the hash table in each dimension. We use three different prime numbers in the hope that the distribution of the hash function values is not skewed even if pairs of l_1 , l_2 , l_3 are correlated. The node-triplet [i,j,k] is hashed to the 3D bin with the address $h[d_1][d_2][d_3]$. Intuitively, we use the squares of the lengths of the three edges connecting the three chosen nodes to determine the hash bin address. Stored in that bin are the graph identification number, the substructure identification number, and the label-triplet code. In addition, we store the coordinates of the basis points P_{b_1} , P_{b_2} , P_{b_3} of Substructure Frame 0 (SF_0) with respect to the three chosen nodes.

Specifically, suppose the chosen nodes i, j, k are not collinear. We can construct another local coordinate frame, denoted LF[i,j,k], using $\vec{V}_{i,j}$, $\vec{V}_{i,k}$ and $\vec{V}_{i,j} \times \vec{V}_{i,k}$ as basis vectors. The coordinates of P_{b_1} , P_{b_2} , P_{b_3} with respect to the local coordinate frame LF[i,j,k], denoted $SF_0[i,j,k]$, form a 3×3 matrix, which is calculated as follows (see Fig. 7):

$$SF_0[i,j,k] = \begin{pmatrix} \vec{V}_{i,b_1} \\ \vec{V}_{i,b_2} \\ \vec{V}_{i,b_3} \end{pmatrix} \times A^{-1}, \tag{3}$$

where

$$A = \begin{pmatrix} \vec{V}_{i,j} \\ \vec{V}_{i,k} \\ \vec{V}_{i,j} \times \vec{V}_{i,k} \end{pmatrix}. \tag{4}$$

Thus, suppose the graph in Fig. 1 has identification number 12. The hash bin entry for the three chosen nodes i, j, k is $(12, 0, Lcode, SF_0[i, j, k])$, where Lcode is the label-triplet code. Since there are 6 nodes in the substructure Str_0 , we have

$$\binom{6}{3} = 20$$
 node-triplets

generated from the substructure and, therefore, 20 entries in the hash table for the substructure.⁷

Example 5. To illustrate how we hash and store patterns in the hash table, let us consider Table 1 again. The basis points of SF_0 of Fig. 2a have the following global coordinates:

$$P_{b_1}(1.0178, 1.0048, 2.5101),$$

 $P_{b_2}(2.0178, 1.0048, 2.5101),$
 $P_{b_2}(1.0178, 2.0048, 2.5101).$

The local coordinates, with respect to SF_0 , of the nodes numbered 0, 1, 2, 3, and 4 in substructure Str_0 of Fig. 2a are as follows:

$$\begin{split} &P_0'(0.0000,0.0000,0.0000),\\ &P_1'(0.1843,1.0362,-0.5081),\\ &P_2'(0.3782,1.9816,-0.5095),\\ &P_3'(-0.3052,1.0442,0.6820),\\ &P_4'(-0.2568,1.7077,0.5023). \end{split}$$

Now, suppose Scale, $Prime_1$, $Prime_2$, $Prime_3$ are 10, 1,009, 1,033, and 1,051, respectively, and Nrow is 31. Thus, for example, for the nodes numbered 1, 2, and 3, the hash bin address is h[25][12][21] and,

$$SF_0[1,2,3] = \begin{pmatrix} -1.056731 & 0.357816 & 0.173981 \\ -0.875868 & 0.071949 & 0.907227 \\ -0.035973 & 0.417517 & 0.024041 \end{pmatrix}.$$
 (5)

As another example, for the nodes numbered 1, 4, and 2, the hash bin address is h[24][0][9] and

$$SF_0[1,4,2] = \begin{pmatrix} 0.369457 & -1.308266 & -0.242990 \\ -0.043584 & -0.857105 & -1.006793 \\ 0.455285 & -0.343703 & -0.086982 \end{pmatrix}. (6)$$

Similarly, for the substructure Str_1 , we attach a local coordinate frame SF_1 to the node numbered 6, as shown in Fig. 2b. There are 10 hash table entries for Str_1 , each having the form $(12, 1, Lcode, SF_1[l, m, n])$ where l, m, n are any three nodes in Str_1 .

7. One interesting question here is regarding the size limitation of patterns (rigid substructures) found by our algorithm. Suppose the graph identification number and the substructure identification number are both short integers of 2 bytes. Lcode is a long integer of 4 bytes. $SF_0[i,j,k]$ is a 3×3 matrix of floating point numbers, each requiring 4 bytes. Thus, each node-triplet requires 44 bytes. For a set of M patterns (substructures), each having T nodes on average, the hash table requires $M\times \binom{T}{3}\times 44$ bytes of disk space. Suppose there are 10,000 patterns, i.e., M=10,000. If T=20, the hash table requires about 502 Megabytes. If T=100, the hash table requires over 71 Gigabytes. This gives an estimate of how large patterns can be in practice. In our case, the pattern sizes are between 5 and 13, so the size demands are modest. Note that the time and memory space needed also increase dramatically as patterns become large.

Recall that we choose the three nodes i,j,k based on the triangle formed by them—the first node chosen always opposes the longest edge of the triangle and the third node chosen opposes the shortest edge. Without loss of generality, let us assume that the nodes i,j,k are chosen in that order. Thus, $\vec{V}_{i,j}$ has the shortest length, $\vec{V}_{i,k}$ is the second shortest and $\vec{V}_{j,k}$ is the longest. We use node i as the origin, $\vec{V}_{i,j}$ as the X-axis and $\vec{V}_{i,k}$ as the Y-axis. Then, construct the local coordinate frame LF[i,j,k] using $\vec{V}_{i,j},\vec{V}_{i,k}$, and $\vec{V}_{i,j} \times \vec{V}_{i,k}$ as basis vectors. Thus, we exclude the longest vector $\vec{V}_{j,k}$ when constructing LF[i,j,k]. Here is why.

The coordinates (x,y,z) of each node in a 3D graph have an error due to rounding. Thus, the real coordinates for the node should be $(\overline{x},\overline{y},\overline{z})$, where $\overline{x}=x+\epsilon_1$, $\overline{y}=y+\epsilon_2$, $\overline{z}=z+\epsilon_3$ for three small decimal fractions $\epsilon_1,\epsilon_2,\epsilon_3$. After constructing LF[i,j,k] and when calculating $SF_0[i,j,k]$, one may add or multiply the coordinates of the 3D vectors. We define the *accumulative error* induced by a calculation C, denoted $\Delta(C)$, as

$$\Delta(C) = |\overline{f} - f|,$$

where \bar{f} is the result obtained from C with the real coordinates and f is the result obtained from C with rounding errors.

Recall that in calculating $SF_0[i,j,k]$, the three basis vectors of LF[i,j,k] all appear in the matrix A defined in (4). Let

$$\overline{|\vec{V}_{i,j}|} = |\vec{V}_{i,j}| + \delta_1, \overline{|\vec{V}_{i,k}|} = |\vec{V}_{i,k}| + \delta_2,$$
 and
$$\overline{|\vec{V}_{j,k}|} = |\vec{V}_{j,k}| + \delta_3. \text{ Let } \delta = \max\{|\delta_1|, |\delta_2|, |\delta_3|\}. \text{ Notice}$$

$$|\vec{V}_{i,j} \times \vec{V}_{i,k}| = |\vec{V}_{i,j}| |\vec{V}_{i,k}| \sin \theta,$$

where $|\vec{V}_{i,j}|$ is the length of $\vec{V}_{i,j}$, and θ is the angle between $\vec{V}_{i,j}$ and $\vec{V}_{i,k}$. Thus,

$$\begin{split} &\Delta(|\vec{V}_{i,j} \times \vec{V}_{i,k}|) \\ &= ||\vec{V}_{i,j}||\vec{V}_{i,k}|\sin\theta - |\vec{V}_{i,j}||\vec{V}_{i,k}|\sin\theta| \\ &= |(|\vec{V}_{i,j}| + \delta_1)(|\vec{V}_{i,k}| + \delta_2)\sin\theta - |\vec{V}_{i,j}||\vec{V}_{i,k}|\sin\theta| \\ &= |(|\vec{V}_{i,j}|\delta_2 + |\vec{V}_{i,k}|\delta_1 + \delta_1\delta_2)||\sin\theta| \\ &\leq (|\vec{V}_{i,j}||\delta_2| + |\vec{V}_{i,k}||\delta_1| + |\delta_1||\delta_2|)|\sin\theta| \\ &\leq (|\vec{V}_{i,j}| + |\vec{V}_{i,k}| + \delta)\delta \\ &= U_1 \end{split}$$

Likewise,

$$\Delta(|\vec{V}_{i,j} \times \vec{V}_{i,k}|) \le (|\vec{V}_{i,j}| + |\vec{V}_{i,k}| + \delta)\delta = U_2$$

and

$$\Delta(|\vec{V}_{i,k} \times \vec{V}_{j,k}|) \le (|\vec{V}_{i,k}| + |\vec{V}_{j,k}| + \delta)\delta = U_3.$$

Among the three upperbounds U_1 , U_2 , U_3 , U_1 is the smallest. It's likely that the accumulative error induced by calculating the length of the cross product of the two corresponding vectors is also the smallest. Therefore, we choose $\vec{V}_{i,j}$, $\vec{V}_{i,k}$, and exclude the longest vector $\vec{V}_{j,k}$ in constructing the local coordinate frame LF[i,j,k], so as to minimize the accumulative error induced by calculating $SF_0[i,j,k]$.

2.3.2 Subphase B of Phase 2

Let \mathcal{H} be the resulting hash table obtained in subphase A of phase 2 of the pattern-finding algorithm. In subphase B, we calculate the occurrence number of each candidate pattern P by rehashing the node-triplets of P into \mathcal{H} . This way, we are able to match a node-triplet tri of P with a node-triplet tri' of another substructure (candidate pattern) P' stored in subphase A where tri and tri' have the same hash bin address. By counting the node-triplet matches, one can infer whether P matches P' and, therefore, whether P occurs in the graph from which P' is generated.

We associate each substructure with several counters, which are created and updated as illustrated by the following example. Suppose the two substructures (patterns) of graph G with identification number 12 in Fig. 1 have already been stored in the hash table \mathcal{H} in subphase A. Suppose i, j, k, are three nodes in the substructure Str_0 of G. Thus, for this node-triplet, its entry in the hash table is $(12, 0, Lcode, SF_0[i, j, k])$. Now, in subphase B, consider another pattern P; we hash the node-triplets of P using the same hash function. Let u, v, w, be three nodes in P that have the same hash bin address as i, j, k; that is, the nodetriplet [u, v, w] "matches" the node-triplet [i, j, k]. If the nodes u, v, w, geometrically match the nodes i, j, k respectively, i.e., they have coinciding 3D coordinates after rotations and translations, we call the node-triplet match a true match; otherwise it is a false match. For a true match, let

$$SF_P = SF_0[i, j, k] \times \begin{pmatrix} \vec{V}_{u,v} \\ \vec{V}_{u,w} \\ \vec{V}_{u,v} \times \vec{V}_{u,w} \end{pmatrix} + \begin{pmatrix} P_u \\ P_u \\ P_u \end{pmatrix}. \tag{7}$$

This SF_P contains the coordinates of the three basis points of the Substructure Frame 0 (SF_0) with respect to the global coordinate frame in which the pattern P is given. We compare the SF_P with those already associated with the substructure Str_0 (initially none is associated with Str_0). If the SF_P differs from the existing ones, a new counter is created, whose value is initialized to 1, and the new counter is assigned to the SF_P . If the SF_P is the "same" as an existing one with counter value Cnt_1^8 and the code of the label-triplet of nodes i, j, k, equals the code of the label-triplet of nodes u, v, w, then Cnt is incremented by one. In general, a substructure may be associated with several different SF_P s, each having a counter.

We now present the theory supporting this algorithm. Below, Theorem 1 establishes a criterion based on which one can detect and eliminate a false match. Below, Theorem 2 justifies the procedure of incrementing the counter values.

Theorem 1. Let P_{c_1} , P_{c_2} , and P_{c_3} be the three basis points forming the SF_P defined in (7), where P_{c_1} is the origin. \vec{V}_{c_1,c_2} , \vec{V}_{c_1,c_3} , and $\vec{V}_{c_1,c_2} \times \vec{V}_{c_1,c_3}$ are orthonormal vectors if and only if the

8. By saying SF_P is the same as an existing SF_P' , we mean that for each entry $e_{i,j}, 1 \leq i, j \leq 3$, at the ith row and the jth column in SF_P and its corresponding entry $e'_{i,j}$ in SF_P' , $|e_{i,j}-e'_{i,j}| \leq \epsilon$, where ϵ is an adjustable parameter depending on the data. In the examples presented in the paper, $\epsilon=0.01$.

nodes u, v, and w geometrically match the nodes i, j, and k, respectively.

Proof. (If) Let A be as defined in (4) and let

$$B = \begin{pmatrix} \vec{V}_{u,v} \\ \vec{V}_{u,w} \\ \vec{V}_{u,v} \times \vec{V}_{u,w} \end{pmatrix}. \tag{8}$$

Note that, if u, v, and w geometrically match i, j, and k, respectively, then |B| = |A|, where |B| (|A|, respectively) is the determinant of the matrix B (matrix A, respectively). That is to say, $|A^{-1}||B| = 1$.

From (3) and by the definition of the SF_P in (7), we have

$$SF_P = \begin{pmatrix} \vec{V}_{i,b_1} \\ \vec{V}_{i,b_2} \\ \vec{V}_{i,b_3} \end{pmatrix} \times A^{-1} \times B + \begin{pmatrix} P_u \\ P_u \\ P_u \end{pmatrix}. \tag{9}$$

Thus, the SF_P basically transforms P_{b_1} , P_{b_2} , and P_{b_3} via two translations and one rotation, where P_{b_1} , P_{b_2} , and P_{b_3} are the basis points of the Substructure Frame 0 (SF_0). Since \vec{V}_{b_1,b_2} , \vec{V}_{b_1,b_3} , and $\vec{V}_{b_1,b_2} \times \vec{V}_{b_1,b_3}$ are orthonormal vectors, and translations and rotations do not change this property [27], we know that \vec{V}_{c_1,c_2} , \vec{V}_{c_1,c_3} , and $\vec{V}_{c_1,c_2} \times \vec{V}_{c_1,c_3}$ are orthonormal vectors.

(Only if) If u, v, and w do not match i, j and k geometrically while having the same hash bin address, then there would be distortion in the aforementioned transformation. Consequently, \vec{V}_{c_1,c_2} , \vec{V}_{c_1,c_3} , and $\vec{V}_{c_1,c_2} \times \vec{V}_{c_1,c_3}$ would no longer be orthonormal vectors. \square

Theorem 2. If two true node-triplet matches yield the same SF_P and the codes of the corresponding label-triplets are the same, then the two node-triplet matches are augmentable, i.e., they can be combined to form a larger substructure match between P and Str_0 .

Proof. Since three nodes are enough to set the SF_P at a fixed position and direction, all the other nodes in P will have definite coordinates under this SF_P . When another nodetriplet match yielding the same SF_P occurs, it means that geometrically there is at least one more node match between Str_0 and P. If the codes of the corresponding label-triplets are the same, it means that the labels of the corresponding nodes are the same. Therefore, the two node-triplet matches are augmentable.

Fig. 8 illustrates how two node-triplet matches are augmented. Suppose the node-triplet [3,4,2] yields the SF_P shown in the figure. Further, suppose that the node-triplet [1,2,3] yields the same SF_P , as shown in Fig. 8. Since the labels of the corresponding nodes numbered 3 and 2 are the same, we can augment the two node-triplets to form a larger match containing nodes numbered 1, 2, 3, 4.

Thus, by incrementing the counter associated with the SF_P , we record how many true node-triplet matches are augmentable under this SF_P . Notice that in cases where two node-triplet matches occur due to reflections, the directions

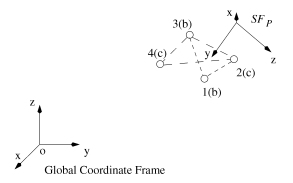


Fig. 8. Illustration of two augmentable node-triplet matches.

of the corresponding local coordinate systems are different, so are the SF_P s. As a result, these node-triplet matches are not augmentable.

Example 6. To illustrate how we update counter values for augmentable node-triplet matches, let us consider the pattern P in Fig. 9. In P, the nodes numbered 0, 1, 2, 3, 4 match, after rotation, the nodes numbered 5, 4, 3, 1, 2 in the substructure Str_0 in Fig. 2a. The node numbered 0 in Str_0 does not appear in P (i.e., it is to be deleted). The labels of the corresponding nodes are identical. Thus, P matches Str_0 with 1 mutation, i.e., one node is deleted.

Now, suppose in *P*, the global coordinates of the nodes numbered 1, 2, 3, and 4 are

$$P_1(-0.269000, 4.153153, 2.911494),$$

 $P_2(-0.317400, 4.749386, 3.253592),$
 $P_3(0.172100, 3.913515, 4.100777),$
 $P_4(0.366000, 3.244026, 3.433268).$

Refer to Example 5. For the nodes numbered 3, 4, and 2 of P, the hash bin address is h[25][12][21], which is the same as that of nodes numbered 1, 2, 3 of Str_0 , and

$$SF_P = \begin{pmatrix} -0.012200 & 5.005500 & 4.474200 \\ 0.987800 & 5.005500 & 4.474200 \\ -0.012200 & 4.298393 & 3.767093 \end{pmatrix}. \tag{10}$$

The three basis vectors forming this SF_P are

$$\vec{V}_{c_1,c_2} = (1.000000, 0.000000, 0.000000),$$

$$\vec{V}_{c_1,c_3} = (0.000000, -0.707107, -0.707107),$$

$$\vec{V}_{c_1,c_3} \times \vec{V}_{c_1,c_3} = (0.000000, 0.707107, -0.707107),$$

which are orthonormal.

For the nodes numbered 3, 1, and 4 of P, the hash bin address is h[24][0][9], which is the same as that of nodes numbered 1, 4, 2 of Str_0 , and

$$SF_P = \begin{pmatrix} -0.012200 & 5.005500 & 4.474200 \\ 0.987800 & 5.005500 & 4.474200 \\ -0.012200 & 4.298393 & 3.767093 \end{pmatrix}. \tag{11}$$

These two true node-triplet matches have the same SF_P and, therefore, the corresponding counter associated with the substructure Str_0 of the graph 12 in Fig. 1 is updated to 2. After hashing all node-triplets of P, the counter value will be

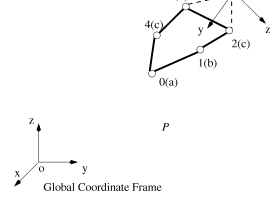


Fig. 9. A substructure (pattern) P.

$$\binom{5}{3} = 10,$$

since all matching node-triplets have the same SF_P as in (10) and the labels of the corresponding nodes are the same.

Now, consider again the SF_P defined in (7) and the three basis points P_{c_1} , P_{c_2} , P_{c_3} forming the SF_P , where P_{c_1} is the origin. We note that for any node i in the pattern P with global coordinate $P_i(x_i, y_i, z_i)$, it has a local coordinate with respect to the SF_P , denoted P'_i , where

$$P_i' = \vec{V}_{c_i,i} \times E^{-1}.$$
 (12)

Here, E is the base matrix for the SF_P , defined as

$$E = \begin{pmatrix} \vec{V}_{c_1, c_2} \\ \vec{V}_{c_1, c_3} \\ \vec{V}_{c_1, c_2} \times \vec{V}_{c_1, c_3} \end{pmatrix}$$
 (13)

and $\vec{V}_{c_1,i}$ is the vector starting at P_{c_1} and ending at P_i .

Remark. If \vec{V}_{c_1,c_2} , \vec{V}_{c_1,c_3} , and $\vec{V}_{c_1,c_2} \times \vec{V}_{c_1,c_3}$ are orthonormal vectors, then |E|=1. Thus, a practically useful criterion for detecting false matches is to check whether or not |E|=1. If $|E|\neq 1$, then \vec{V}_{c_1,c_2} , \vec{V}_{c_1,c_3} , and $\vec{V}_{c_1,c_2} \times \vec{V}_{c_1,c_3}$ are not orthonormal vectors and, therefore, the nodes u,v, and w do not match the nodes i,j and k geometrically (see Theorem 1).

Intuitively, our scheme is to hash node-triplets and match the triplets. Only if one triplet tri matches another tri' do we see how the substructure containing tri matches the pattern containing tri'. Using Theorem 1, we detect and eliminate false node-triplet matches. Using Theorem 2, we record in a counter the number of augmentable true node-triplet matches. The following theorem says that the counter value needs to be large (i.e., there are a sufficient number of augmentable true node-triplet matches) in order to infer that there is a match between the corresponding pattern and substructure. The larger the Mut, the fewer node-triplet matches are needed.

Theorem 3. Let Str be a substructure in the hash table $\mathcal H$ and let G be the graph from which Str is generated. Let P be a pattern where $|P| \geq Mut + 3$. After rehashing the node-triplets of P,

suppose there is an SF_P associated with Str whose counter value $Cnt > \Theta_P$, where

$$\Theta_P = \binom{N-1}{3} = \frac{(N-1) \times (N-2) \times (N-3)}{6}$$
 (14)

and N = |P| - Mut. Then, P matches Str within Mut mutations (i.e., P approximately occurs in G, or G approximately contains P, within Mut mutations).

Proof. By Theorem 2, we increase the counter value only when there are true node-triplet matches that are augmentable under the SF_P . Thus, the counter value shows currently how many node-triplets from the pattern P are found to match with the node-triplets from the substructure Str in the hash table. Note that Θ_P represents the number of distinct node-triplets that can be generated from a substructure of size N-1. If there are N-1 node matches between P and Str, then $Cnt \leq \Theta_P$. Therefore, when $Cnt > \Theta_P$, there are at least N node matches between Str and P. This completes the proof.

Notice that our algorithm cannot handle patterns with size smaller than 3, because the basic processing unit here is a node-triplet of size 3. This is reflected in the condition $|P| \geq Mut + 3$ stated in Theorem 3. Since Mut can only be zero or greater than zero, this condition implies that the size of a pattern must be greater than or equal to 3.

Example 7. To illustrate how Theorem 3 works, let us refer to Example 6. Suppose the user-specified mutation number Mut is 1. The candidate pattern P in Fig. 9 has size |P|=5. After rehashing the node-triplets of P, there is only one counter associated with the substructure Str_0 in Fig. 2a; this counter corresponds to the SF_P in (10) and the value of the counter, Cnt, is 10. Thus, Cnt is greater than $\Theta_P=(5-2)(5-3)(5-4)/6=1$. By Theorem 3, P should match the substructure Str_0 within 1 mutation. This means that there are at least four node matches between P and Str_0 .

Thus, after rehashing the node-triplets of each candidate pattern P into the 3D table \mathcal{H} , we check the values of the counters associated with the substructures in \mathcal{H} . By Theorem 3, P approximately occurs in a graph G within Mut mutations if G contains a substructure Str and there is at least one counter associated with Str whose value $Cnt > \Theta_P$. If there are less than Occur graphs in which P approximately occurs within Mut mutations, then we discard P. The remaining candidates are qualified patterns. Notice that Theorem 3 provides only the "sufficient" (but not the "necessary") condition for finding the qualified patterns. Due to the accumulative errors arising in the calculations, some node-triplets may be hashed to a wrong bin. As a result, the pattern-finding algorithm may miss some node-triplet matches and, therefore, miss some

qualified patterns. In Section 3, we will show experimentally that the missed patterns are few compared with those found by exhaustive search.

Theorem 4. Let the set S contain K graphs, each having at most N nodes. The time complexity of the proposed pattern-finding algorithm is $O(KN^3)$.

Proof. For each graph in \mathcal{S} , phase 1 of the algorithm requires $\mathcal{O}(N^2)$ time to decompose the graph into substructures. Thus, the time needed for phase 1 is $\mathcal{O}(KN^2)$. In subphase A of phase 2, we hash each candidate pattern P by considering the combinations of any three nodes in P, which requires time

$$\binom{|P|}{3} = \mathcal{O}(|P|^3).$$

Thus, the time needed to hash all candidate patterns is $\mathcal{O}(KN^3)$. In subphase B of phase 2, we rehash each candidate pattern, which requires the same time $\mathcal{O}(KN^3)$.

3 Performance Evaluation

We carried out a series of experiments to evaluate the performance and the speed of our approach. The programs were written in the C programming language and run on a SunSPARC 20 workstation under the Solaris operating system version 2.4. Parameters used in the experiments can be classified into two categories: those related to data and those related to the pattern-finding algorithm. In the first category, we considered the size (in number of nodes) of a graph and the total number of graphs in a data set. In the second category, we considered all the parameters described in Section 2, which are summarized in Table 3 together with the base values used in the experiments.

Two files were maintained: One recording the hash bin addresses and the other containing the entries stored in the hash bins. To evaluate the performance of the pattern-finding algorithm, we applied the algorithm to two sets of data: 1,000 synthetic graphs and 226 chemical compounds obtained from a drug database maintained in the National Cancer Institute. When generating the artificial graphs, we randomly generated the 3D coordinates for each node. The node labels were drawn randomly from the range A to E. The size of the rigid substructures in an artificial graph ranged from 4 to 10 and the size of the graphs ranged from 10 to 50. The size of the compounds ranged from 5 to 51.

In this section, we present experimental results to answer questions concerning the performance of the pattern-finding algorithm. For example, are all approximate patterns found, i.e., is the recall high? Are any uninteresting patterns found, i.e., is the precision high? In Section 4, we study the applications of the algorithm and intend to answer questions such as whether graphs having some common phenomenological activity (e.g., they are proteins with the same function) share structural patterns in common and whether these patterns can characterize the graphs as a whole.

^{9.} Notice that we cannot use $Cnt = \Theta_P$ here. The reason is that we augment node-triplet matches only when they yield the same SF_P . In practice, it's likely that two node-triplet matches could be augmented, but yield different SF_P s due to the errors accumulated during the calculation of the SF_P s. As a consequence, our algorithm fails to detect those node-triplet matches and update the counter values appropriately.

T .	77.1	
Parameter	Value	Description
Mut	1	Allowed mutation between a pattern and a graph
Occur	3	Minimum occurrence number of an interesting pattern
Size	6	Minimum size of an interesting pattern
ϵ	0.01	Allowed error in comparing the entries of two coordinate matrices
Scale	10	the multiplier used in calculating the hash bin address
$Prime_1$	1,009	the 1st prime number used in calculating the hash bin address
$Prime_2$	1,033	the 2nd prime number used in calculating the hash bin address
$Prime_3$	1,051	the 3rd prime number used in calculating the hash bin address
Nrow	101	the cardinality of the hash table along each dimension

TABLE 3
Parameters in the Pattern-Finding Algorithm and Their Base Values Used in the Experiments

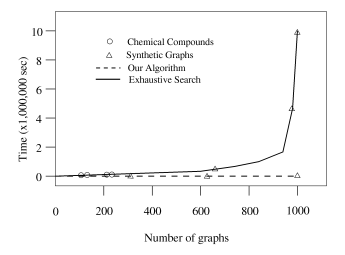
3.1 Effect of Data-Related Parameters

To evaluate the performance of the proposed pattern-finding algorithm, we compared it with exhaustive search. The exhaustive search procedure works by generating all candidate patterns as in phase 1 of the pattern-finding algorithm. Then, the procedure examines if a pattern P approximately matches a substructure Str in a graph by permuting the node labels of P and checking if they match the node labels of Str. If so, the procedure performs translation and rotation on P and checks if P can geometrically match Str.

The speed of the algorithms was measured by the running time. The performance was evaluated using three measures: recall (RE), precision (PR), and the number of false matches, N_{fm} , arising during the hashing process. (Recall that a false match arises, if a node-triplet [u,v,w] from a pattern P has the same hash bin address as a node-triplet [i,j,k] from a substructure of graph G, though the nodes u,v,w do not match the nodes i,j,k geometrically, see Section 2.3.2.) Recall is defined as

$$RE = \frac{UPFound}{TotalP} \times 100\%.$$

Precision is defined as



$$PR = \frac{UPFound}{PFound} \times 100\%,$$

where PFound is the number of patterns found by the proposed algorithm, UPFound is the number of patterns found that satisfy the user-specified parameter values, and TotalP is the number of qualified patterns found by exhaustive search. One would like both RE and PR to be as high as possible. Fig. 10 shows the running times of the algorithms as a function of the number of graphs and Fig. 11 shows the recall. The parameters used in the proposed pattern-finding algorithm had the values shown in Table 3. As can be seen from the figures, the proposed algorithm is 10,000 times faster than the exhaustive search method when the data set has more than 600 graphs while achieving a very high (> 97%) recall. Due to the accumulative errors arising in the calculations, some node-triplets may be hashed to a wrong bin. As a result, the proposed algorithm may miss some node-triplet matches in subphase B of phase 2 and, therefore, cannot achieve a 100 percent recall. In these experiments, precision was 100 percent.

Fig. 12 shows the number of false matches introduced by the proposed algorithm as a function of the number of graphs. For the chemical compounds, N_{fm} is small. For the synthetic graphs, N_{fm} increases as the number of graphs

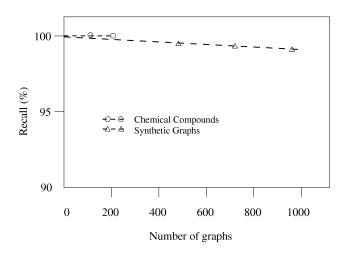
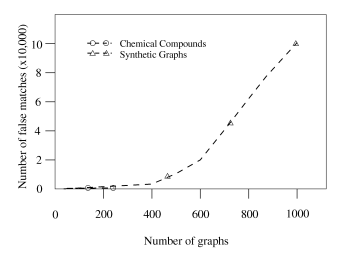
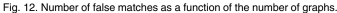


Fig. 11. Recall as a function of the number of graphs.





becomes large. Similar results were obtained when testing the size of graphs for both types of data.

3.2 Effect of Algorithm-Related Parameters

The purpose of this section is to analyze the effect of varying the algorithm-related parameter values on the performance of the proposed pattern-finding algorithm. To avoid the mutual influence of parameters, the analysis was carried out by fixing the parameter values related to data graphs—the 1,000 synthetic graphs and 226 compounds described above were used, respectively. In each experiment, only one algorithm-related parameter value was varied; the other parameters had the values shown in Table 3.

We first examined the effect of varying the parameter values on generating false matches. Since few false matches were found for chemical compounds, the experiments focused on synthetic graphs. It was observed that only Nrow and Scale affected the number of false matches. Fig. 13 shows N_{fm} as a function of Scale for Nrow = 101, 131, 167, 199, respectively. The larger the Nrow, the fewer entries in a hash bin, and consequently

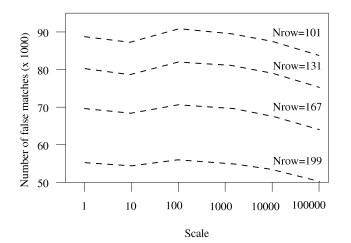


Fig. 13. Number of false matches as a function of Scale.

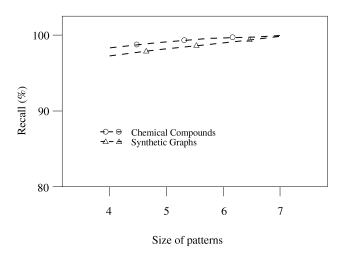


Fig. 14. Effect of Size.

the fewer false matches. On the other hand, when *Nrow* is too large, the running times increase substantially, since one needs to spend a lot of time in reading the 3D table containing the hash bin addresses.

Examining Fig. 13, we see how Scale affects N_{fm} . Taking an extreme case, when Scale = 1, a node-triplet with the squares of the lengths of the three edges connecting the nodes being 12.4567 is hashed to the same bin as a nodetriplet with those values being 12.0000, although the two node-triplets do not match geometrically, see Section 2.3.1. On the other hand, when Scale is large (e.g., Scale = 10,000), the distribution of hash function values is less skewed. which reduces the number of false matches. It was observed that N_{fm} was largest when Scale was 100. This happens because with this Scale value, inaccuracy was being introduced in calculating hash bin addresses. A node-triplet being hashed to a bin might generate k false matches where k is the total number of node-triplets already stored in that bin—k would be large if the distribution of hash function values is skewed.

Figs. 14, 15, and 16 show the recall as a function of *Size*, *Mut*, and *Scale*, respectively. In all three figures, precision

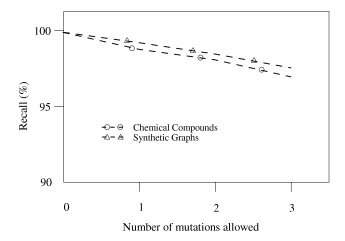


Fig. 15. Effect of Mut.

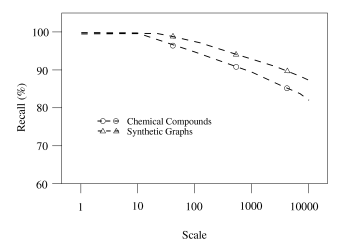


Fig. 16. Impact of Scale.

is 100 percent. From Fig. 14 and Fig. 15, we see that Size and Mut affect recall slightly. It was also observed that the number of interesting patterns drops (increases, respectively) significantly as Size (Mut, respectively) becomes large. Fig. 16 shows that the pattern-finding algorithm yields a poor performance when Scale is large. With the data we tested, we found that setting Scale to 10 is the best overall for both recall and precision.

In sum, the value of Scale has a significant impact on the performance of our algorithm. The choice of Scale is domain and data dependent. In practice, how would one select a value of Scale for a particular database? Recall that the coordinates (x,y,z) of each node in a 3D graph have an error due to rounding, see Section 2.3.1. The real coordinates for the node at (x_i,y_i,z_i) should be $(\overline{x_i},\overline{y_i},\overline{z_i})$, where $\overline{x_i}=x_i+\epsilon_{x_i},\ \overline{y_i}=y_i+\epsilon_{y_i},\ \overline{z_i}=z_i+\epsilon_{z_i}.$ Similarly, the real coordinates for the node at (x_j,y_j,z_j) should be $(\overline{x_j},\overline{y_j},\overline{z_j})$, where $\overline{x_j}=x_j+\epsilon_{x_j},\ \overline{y_j}=y_j+\epsilon_{y_j},\ \overline{z_j}=z_j+\epsilon_{z_j}.$ The real value of l_1 in (2) should be

$$\begin{split} \overline{l_1} &= ((\overline{x_i} - \overline{x_j})^2 + (\overline{y_i} - \overline{y_j})^2 + (\overline{z_i} - \overline{z_j})^2) \times Scale \\ &= ((x_i + \epsilon_{x_i} - x_j - \epsilon_{x_j})^2 + (y_i + \epsilon_{y_i} - y_j - \epsilon_{y_j})^2 \\ &+ (z_i + \epsilon_{z_i} - z_j - \epsilon_{z_j})^2) \times Scale \\ &= ((x_i - x_j)^2 + (y_i - y_j)^2 + (z_i - z_j)^2) \times Scale \\ &+ 2 \times ((x_i - x_j)(\epsilon_{x_i} - \epsilon_{x_j}) + (y_i - y_j)(\epsilon_{y_i} - \epsilon_{y_j}) \\ &+ (z_i - z_j)(\epsilon_{z_i} - \epsilon_{z_j})) \times Scale \\ &+ ((\epsilon_{x_i} - \epsilon_{x_j})^2 + (\epsilon_{y_i} - \epsilon_{y_j})^2 + (\epsilon_{z_i} - \epsilon_{z_j})^2) \times Scale \\ &= l_1 + \epsilon_{l_1}, \end{split}$$

where ϵ_{l_1} is an accumulative error,

$$\epsilon_{l_1} = 2 \times ((x_i - x_j)(\epsilon_{x_i} - \epsilon_{x_j}) + (y_i - y_j)(\epsilon_{y_i} - \epsilon_{y_j})$$

$$+ (z_i - z_j)(\epsilon_{z_i} - \epsilon_{z_j}) \times Scale$$

$$+ ((\epsilon_{x_i} - \epsilon_{x_j})^2 + (\epsilon_{y_i} - \epsilon_{y_j})^2 + (\epsilon_{z_i} - \epsilon_{z_j})^2) \times Scale$$

$$(15)$$

Since we used l_1 to calculate the hash bin addresses, it is critical that the accumulative error would not mislead us to a wrong hash bin. Assuming that the coordinates are accurate

to the nth digit after the decimal point, i.e., to 10^{-n} , the error caused by eliminating the digits after the (n+1)th position is no larger than 0.5×10^{-n} . Namely, for any coordinates (x,y,z), we have $\epsilon_x\leq 0.5\times 10^{-n}$, $\epsilon_y\leq 0.5\times 10^{-n}$, and $\epsilon_z\leq 0.5\times 10^{-n}$. Thus,

$$\begin{split} \epsilon_{l_1} & \leq 2 \times (|x_i - x_j|| \epsilon_{x_i} - \epsilon_{x_j}| + |y_i - y_j|| \epsilon_{y_i} - \epsilon_{y_j}| \\ & + |z_i - z_j|| \epsilon_{z_i} - \epsilon_{z_j}|) \times Scale \\ & + (|\epsilon_{x_i} - \epsilon_{x_j}|^2 + |\epsilon_{y_i} - \epsilon_{y_j}|^2 + |\epsilon_{z_i} - \epsilon_{z_j}|^2) \times Scale \\ & \leq 2 \times (|x_i - x_j|(|\epsilon_{x_i}| + |\epsilon_{x_j}|) + |y_i - y_j|(|\epsilon_{y_i}| \\ & + |\epsilon_{y_j}|) + |z_i - z_j|(|\epsilon_{z_i}| + |\epsilon_{z_j}|)) \times Scale \\ & + ((|\epsilon_{x_i}| + |\epsilon_{x_j}|)^2 + (|\epsilon_{y_i}| + |\epsilon_{y_j}|)^2 + (|\epsilon_{z_i}| + |\epsilon_{z_j}|)^2) \times Scale \\ & \leq 2 \times (|x_i - x_j| + |y_i - y_j| + |z_i - z_j|) \\ & \times 2 \times 0.5 \times 10^{-n} \times Scale \\ & + 3 \times (2 \times 0.5 \times 10^{-n})^2 \times Scale \\ & = (|x_i - x_j| + |y_i - y_j| + |z_i - z_j|) \\ & \times 2 \times Scale \times 10^{-n} + 3 \times Scale \times 10^{-2n}. \end{split}$$

Assume that the range of the coordinates is [-M, M]. We have $|x_i - x_j| \le 2M$, $|y_i - y_j| \le 2M$, and $|z_i - z_j| \le 2M$. Thus,

$$\epsilon_{l_1} \leq 12 \times M \times Scale \times 10^{-n} + 3 \times Scale \times 10^{-2n}$$
.

The second term is obviously negligible in comparison with the first term. In order to keep the calculation of hash bin addresses accurate, the first term should be smaller than 1. That is, *Scale* should be chosen to be the largest possible number such that

$$12 \times M \times Scale \times 10^{-n} < 1$$

or

$$Scale < \frac{1}{12 \times M} \times 10^n.$$

In our case, the coordinates of the chemical compounds used were accurate to the $4\mathrm{th}$ digit after the decimal point and the range of the coordinates was [-10, 10]. Consequently,

$$Scale < \frac{10^4}{12 \times 10} \times = \frac{500}{6}$$

Thus, choosing Scale=10 is good, as validated by our experimental results. Notice that Scale can be determined once the data are given. All we need to know is how accurate the data are, and what the ranges of their coordinates are, both of which are often clearly associated with the data.

Figs. 17 and 18 show the recall and precision as a function of ϵ . It can be seen that when ϵ is 0.01, precision is 100 percent and recall is greater than 97 percent. When ϵ becomes smaller (e.g., $\epsilon=0.0001$), precision remains the same while recall drops. When ϵ becomes larger (e.g., $\epsilon=10$), recall increases slightly while precision drops. This happens because some irrelevant node-triplet matches were included, rendering unqualified patterns returned as an answer. We also tested different values for Occur, Nrow, $Prime_1$, $Prime_2$, and $Prime_3$. It was found that varying

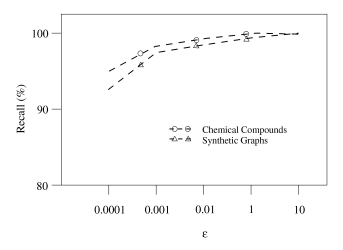


Fig. 17. Recall as a function of ϵ .

these parameter values had little impact on the performance of the proposed algorithm.

Finally, we examined the effect of sensor errors, which are caused by the measuring device (e.g., X-ray crystallography), and are proportional to the values being measured. Suppose that the sensor error is 10^{-n} of the real value for any coordinates (x,y,z), i.e., $\epsilon_x=x\times 10^{-n}$, $\epsilon_y=y\times 10^{-n}$, and $\epsilon_z=z\times 10^{-n}$. We have

$$\overline{x} = x \times (1 + 10^{-n}), \quad \overline{y} = y \times (1 + 10^{-n}),$$

and $\overline{z} = z \times (1+10^{-n})$. Let γ_{l_1} represent the sensor error induced in calculating l_1 in (2). In a way similar to the calculation of the accumulative error ϵ_{l_1} in (15), we obtain

$$\gamma_{l_1} = ((x_i - x_j)^2 + (y_i - y_j)^2 + (z_i - z_j)^2) \times 2 \times 10^{-n} \times Scale + ((x_i - x_j)^2 + (y_i - y_j)^2 + (z_i - z_j)^2) \times 10^{-2n} \times Scale.$$

Let us focus on the first term since it is more significant. Clearly, the proposed approach is not feasible in the environment with large sensor errors. Taking an example, suppose that the sensor error is 10^{-2} of the real value. Again, in order to keep the calculation of hash bin addresses accurate, the accumulative error should be smaller than 1. That is,

$$((x_i - x_j)^2 + (y_i - y_j)^2 + (z_i - z_j)^2) \times 2 \times 10^{-2} \times Scale < 1$$

$$((x_i - x_j)^2 + (y_i - y_j)^2 + (z_i - z_j)^2) \times Scale < 50.$$

Even when Scale = 1, the Euclidean distance between any two nodes must be smaller than 7, since the square of the distance, viz. $((x_i - x_j)^2 + (y_i - y_j)^2 + (z_i - z_j)^2)$, must be less than 50. Most data sets, including all the chemical compounds we used, do not satisfy this restriction.

To illustrate how this would affect recall, consider a substructure P already stored in the hash table. We add a 10^{-2} sensor error to the coordinates of a node v in P. Call the resulting substructure P'. Then, we use P' as a pattern to match the original substructure P in the hash table. Those node-triplets generated from the pattern P' that include the node v would not match the corresponding node-triplets in

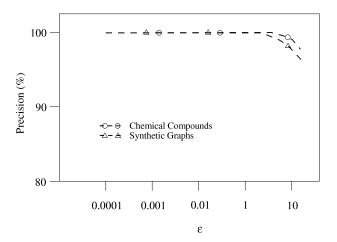


Fig. 18. Precision as a function of ϵ .

the original substructure P. Supposing the pattern P' has N nodes, the total number of node-triplets that include the node v is

$$\binom{N-1}{2}$$
.

Thus, we would miss the same number of node-triplet matches. However, if all other node-triplets that do not include the node v are matched successfully, we would still have

$$\binom{N}{3} - \binom{N-1}{2} = \binom{N-1}{3}$$

node-triplet matches. According to Theorem 3, there would be N-1 node matches between the pattern P' and the original substructure P (i.e., P' matches P with 1 mutation). This example reveals that, in general, if we allow mutations to occur in searching for patterns and the number of nodes with sensor errors equals the allowed number of mutations, the recall will be the same as the case in which there are no sensor errors and we search for exactly matched patterns without mutations. On the other hand, if no mutations are allowed in searching for patterns and sensor errors occur, the recall will be zero.

4 DATA MINING APPLICATIONS

One important application of pattern finding involves the ability to perform classification and clustering as well as other data mining tasks. In this section, we present two data mining applications of the proposed algorithm in scientific domains: classifying proteins and clustering compounds.¹⁰

4.1 Classifying Proteins

Proteins are large molecules, comprising hundreds of amino acids (residues) [18], [33]. In each residue the C_{α} , C_{β} , and N atoms form a backbone of the residue [17]. Following [28], we represent each residue by the three atoms. Thus, if we consider a protein as a 3D graph, each node of the graph is an atom. Each node has a label, which

10. Another application of the proposed algorithm to flexible chemical searching in 3D structural databases can be found in [34].

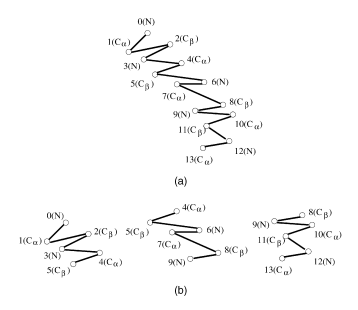


Fig. 19. (a) A 3D protein. (b) The three substructures of the protein in (a).

is the name of the atom and is not unique in the protein. We assign a unique number to identify a node in the protein, where the order of numbering is obtained from the Protein Data Bank (PDB) [1], [2], accessible at http://www.rcsb.org.

In the experiments, we examined two families of proteins chosen from PDB pertaining to RNA-directed DNA Polymerase and Thymidylate Synthase. Each family contains proteins having the same functionality in various organisms. We decompose each protein into consecutive substructures, each substructure containing six nodes. Two adjacent substructures overlap by sharing the two neighboring nodes on the boundary of the two substructures (see Fig. 19). Thus, each substructure is a portion of the polypeptide chain backbone of a protein where the polypeptide chain is made up of residues linked together by peptide bonds. The peptide bonds have strong covalent bonding forces that make the polypeptide chain rigid. As a consequence, the substructures used by our algorithm are rigid. Notice that in the proteins there are other atoms such as O and H (not shown in Fig. 19) lying between two residues. Since these atoms are not as important as C_{α} , C_{β} , and N in determining the structure of a protein, we do not consider them here. Table 4 summarizes the number of proteins in each family, their sizes and the frequently occurring patterns (or motifs) discovered from the proteins. The parameter values used were as shown in Table 3. In the experiments, 2,784 false matches were detected and eliminated during the process of finding the

motifs. That is, there were 2,784 times in which a node-triplet [u, v, w] from a pattern was found to have the same hash bin address as a node-triplet [i, j, k] from a substructure of a protein, though the nodes u, v, w did not match the nodes i, j, k geometrically, see Section 2.3.2.

To evaluate the quality of the discovered motifs, we applied them to classifying the proteins using the 10-way cross-validation scheme. That is, each family was divided into 10 groups of roughly equal size. Specifically, the RNA-directed DNA Polymerase family, referred to as family 1, contained five groups each having five proteins and five groups each having four proteins. The Thymidylate Synthase family, referred to as family 2, contained seven groups each having four proteins and three groups each having three proteins, and 10 tests were conducted. In each test, a group was taken from a family and used as test data; the other nine groups were used as training data for that family. We applied our pattern-finding algorithm to each training data set to find motifs (the parameter values used were as shown in Table 3). Each motif M found in family i was associated with a weight dwhere

$$d = r_i - r_j \qquad 1 \le i, j \le 2, \qquad i \ne j.$$

Here r_i is M's occurrence number in the training data set of family i. Intuitively, the more frequently a motif occurs in its own family and the less frequently it occurs in the other family, the higher its weight is. In each family we collected all the motifs having a weight greater than one and used them as the characteristic motifs of that family.

When classifying a test protein Q, we first decomposed Q into consecutive substructures as described above. The result was a set of substructures, say, Q^1,\ldots,Q^p . Let $n_i^k, 1\leq i\leq 2, 1\leq k\leq p$, denote the number of characteristic motifs in family i that matched Q^k within one mutation. Each family i obtained a score N_i where

$$N_i = \frac{\sum_{k=1}^p n_i^k}{m_i}$$

and m_i is the total number of characteristic motifs in family i. Intuitively, the score N_i here was determined by the number of the characteristic motifs in family i that occurred in Q, divided by the total number of characteristic motifs in family i. The protein Q was classified into the family i with maximum N_i . If the scores were 0 for both families (i.e., the test protein did not have any substructure that matched any characteristic motif), then the "no-opinion" verdict was given. This algorithm is similar to those used in [30], [35] to classify chemical compounds and sequences.

TABLE 4
Statistics Concerning the Proteins and Motifs Found in Them

Family	Number of proteins	Maximum protein size	Minimum protein size	Number of motifs	Motif size
RNA-directed DNA Polymerase	45	1,812	146	42	6
Thymidylate Synthase	37	2,128	1,000	33	6

As in Section 3, we use recall (RE_c) and precision (PR_c) to evaluate the effectiveness of our classification algorithm. Recall is defined as

$$RE_{c} = \frac{TotalNum - \sum_{i=1}^{2} NumLoss_{c}^{i}}{TotalNum} \times 100\%,$$

where TotalNum is the total number of test proteins and $NumLoss_c^i$ is the number of test proteins that belong to family i but are not assigned to family i by our algorithm (they are either assigned to family j, $j \neq i$, or they receive the "no-opinion" verdict). Precision is defined as

$$PR_{c} = \frac{TotalNum - \sum_{i=1}^{2} NumGain_{c}^{i}}{TotalNum} \times 100\%,$$

where $NumGain_c^i$ is the number of test proteins that do not belong to family i but are assigned by our algorithm to family i. With the 10-way cross validation scheme, the average RE_c over the 10 tests was 92.7 percent and the average PR_c was 96.4 percent. It was found that 3.7 percent test proteins on average received the "no-opinion" verdict during the classification. We repeated the same experiments using other parameter values and obtained similar results, except that larger Mut values (e.g., 3) generally yielded lower RE_c .

The binary classification problem studied here is concerned with assigning a protein to one of two families. This problem arises frequently in protein homology detection [31]. The performance of our technique degrades when applied to the n-ary classification problem, which is concerned with assigning a protein to one of many families [32]. For example, we applied the technique to classifying three families of proteins and the RE_c and PR_c dropped to 80 percent.

4.2 Clustering Compounds

In addition to classifying proteins, we have developed an algorithm for clustering 3D graphs based on the patterns occurring in the graphs and have applied the algorithm to grouping compounds. Given a collection $\mathcal S$ of 3D graphs, the algorithm first uses the procedure depicted in Section 2.2 to decompose the graphs into rigid substructures. Let $\{Str_p|p=0,1,...,N-1\}$ be the set of substructures found in the graphs in $\mathcal S$ where $|Str_p| \geq Size$. Using the proposed pattern-finding algorithm, we examine each graph G_q in $\mathcal S$ and determine whether each substructure Str_p approximately occurs in G_q within Mut mutations. Each graph G_q is represented as a bit string of length N, i.e., $G_q = (b_q^0, b_q^1, ..., b_q^{N-1})$, where

$$b_q^p = \begin{cases} 1 & \text{if } Str_p \text{ occurs in } G_q \text{ within } Mut \text{ mutations} \\ 0 & \text{otherwise.} \end{cases}$$

For example, consider the two patterns P_1 and P_2 in Fig. 3 and the three graphs in Fig. 3a. Suppose the allowed number of mutations is 0. Then, G_1 is represented as 10, G_2 as 01, and G_3 as 10. On the other hand, suppose the allowed number of mutations is 1. Then, G_1 and G_3 are represented as 11 and G_2 as 01.

The distance between two graphs G_x and G_y , denoted $d(G_x, G_y)$, is defined as the Hamming distance [12] between their bit strings. The algorithm then uses the well-known

average-group method [13] to cluster the graphs in \mathcal{S} , which works as follows.

Initially, every graph is a cluster. The algorithm merges two nearest clusters to form a new cluster, until there are only K clusters left where K is a user-specified parameter. The distance between two clusters C_1 and C_2 is given by

$$\frac{1}{|C_1||C_2|} \sum_{G_x \in C_1, G_y \in C_2} |d(G_x, G_y)|, \tag{16}$$

where $|C_i|$, i = 1, 2, is the size of cluster C_i . The algorithm requires $O(N^2)$ distance calculations where N is the total number of graphs in S.

We applied this algorithm to clustering chemical compounds. Ninety eight compounds were chosen from the Merck Index that belonged to three groups pertaining to aromatic, bicyclicalkanes and photosynthesis. The data was created by the CORINA program that converted 2D data (represented in SMILES string) to 3D data (represented in PDB format) [22]. Table 5 lists the number of compounds in each group, their sizes and the patterns discovered from them. The parameter values used were Size=5, Occur=1, Mut=2; the other parameters had the values shown in Table 3.

To evaluate the effectiveness of our clustering algorithm, we applied it to finding clusters in the compounds. The parameter value K was set to 3, as there were three groups. As in the previous sections, we use recall (RE_r) and precision (PR_r) to evaluate the effectiveness of the clustering algorithm. Recall is defined as

$$RE_r = \frac{TotalNum - \sum_{i=1}^{K} NumLoss_r^i}{TotalNum} \times 100\%,$$

where $NumLoss_r^i$ is the number of compounds that belong to group G_i , but are assigned by our algorithm to group G_j , $i \neq j$, and TotalNum is the total number of compounds tested. Precision is defined as

$$PR_r = \frac{TotalNum - \sum_{i=1}^{K} NumGain_r^i}{TotalNum} \times 100\%,$$

where $NumGain_r^i$ is the number of compounds that do not belong to group G_i , but are assigned by our algorithm to group G_i . Our experimental results indicated that $RE_r = PR_r = 99\%$. Out of the 98 compounds, only one compound in the photosynthesis group was assigned incorrectly to the bicyclicalkanes group. We experimented with other parameter values and obtained similar results.¹¹

5 RELATED WORK

There are several groups working on pattern finding (or knowledge discovery) in molecules and graphs. Conklin et al. [4], [5], [6], for example, represented a molecular structure as an image, which comprised a set of parts with their 3D coordinates and a set of relations that were preserved for the image. The authors used an incremental, divisive approach to discover the "knowledge" from a data set, that is, to build a

11. The *Occur* value was fixed at 1 in these experiments because of the fact that all the compounds were represented as binary bit strings.

Group	Number of compounds	Minimum compound size	Maximum compound size	Number of patterns	Minimum pattern size	maximum pattern size
aromatic	36	12	42	58	5	13
bicyclicalkanes	26	16	40	53	5	11
photosynthesis	36	31	44	114	5	11

TABLE 5
Statistics Concerning the Chemical Compounds and Patterns Found in Them

subsumption hierarchy that summarized and classified the data set. The algorithm relied on a measure of similarity among molecular images that was defined in terms of their largest common subimages. In [8], Djoko et al. developed a system, called SUBDUE, that utilized the minimum description length principle to find repeatedly occurring substructures in a graph. Once a substructure was found, the substructure was used to simplify the graph by replacing instances of the substructure with a pointer to the discovered substructure. In [7], Dehaspe et al. used DATALOG to represent compounds and applied data mining techniques to predicting chemical carcinogenicity. Their techniques were based on Mannila and Toivonen's algorithm [15] for finding interesting patterns from a class of sentences in a database.

In contrast to the above work, we use the geometric hashing technique to find approximately common patterns in a set of 3D graphs without prior knowledge of their structures, positions, or occurrence frequency. The geometric hashing technique used here originated from the work of Lamdan and Wolfson for model based recognition in computer vision [14]. Several researchers attempted to parallelize the technique based on various architectures, such as the Hypercube and the Connection Machine [3], [19], [20]. It was observed that the distribution of the hash table entries might be skewed. To balance the distribution of the hash function values, delicate rehash functions were designed [20]. There were also efforts exploring the uncertainty existing in the geometric hashing algorithms [11], [26].

In 1996, Rigoutsos et al. employed geometric hashing and magic vectors for substructure matching in a database of chemical compounds [21]. The magic vectors were bonds among atoms; the choice of them was domain dependent and was based on the type of each individual graph. We extend the work in [21] by providing a framework for discovering approximately common substructures in a set of 3D graphs, and applying our techniques to both compounds and proteins. Our approach differs from [21] in that instead of using the magic vectors, we store a coordinate system in a hash table entry. Furthermore, we establish a theory for detecting and eliminating false matches occurring in the hashing.

More recently, Wolfson and his colleagues also applied geometric hashing algorithms to protein docking and recognition [23], [24], [29]. They attached a reference frame to each substructure, where the reference frame is an orthonormal coordinate frame of arbitrary orientations, established at a "hinge" point. This hinge point can be any

point or a specific point chosen based on chemical considerations. The 3D substructure can be rotated around the hinge point. The authors then developed schemes to generate node-triplets and hash table indices. However, based on those schemes, false matches cannot be detected in the hash table and must be processed in a subsequent, additional verification phase. Thus, even if a pattern appears to match several substructures during the searching phase, one has to compare the pattern with those substructures one by one to make sure they are true matches during the verification phase. When mutations are allowed, which were not considered in [23], [24], [29], a brute-force verification test would be very time-consuming. For example, in comparing our approach with their approach in performing protein classification as described in Section 4, we found that both approaches yield the same recall and precision, though our approach is 10 times faster than their approach.

6 CONCLUSION

In this paper, we have presented an algorithm for finding patterns in 3D graphs. A pattern here is a rigid substructure that may occur in a graph after allowing for an arbitrary number of rotations and translations as well as a small number of edit operations in the pattern or in the graph. We used the algorithm to find approximately common patterns in a set of synthetic graphs, chemical compounds and proteins. This yields a kind of alphabet of patterns. Our experimental results demonstrated the good performance of the proposed algorithm and its usefulness for pattern discovery. We then developed classification and clustering algorithms using the patterns found in the graphs, and applied them to classifying proteins and clustering compounds. Empirical study showed high recall and precision rates for both classification and clustering, indicating the significance of the patterns.

We have implemented the techniques presented in the paper and are combining them with the algorithms for acyclic graph matching [36] into a toolkit. We use the toolkit to find patterns in various types of graphs arising in different domains and as part of our tree and graph search engine project. The toolkit can be obtained from the authors and is also accessible at http://www.cis.njit.edu/~jason/sigmod.html on the Web.

In the future, we plan to study topological graph matching problems. One weakness of the proposed geometric hashing approach is its sensitivity to errors as we discussed in Section 3.2. One has to tune the *Scale*

parameter when the accuracy of data changes. Another weakness is that the geometric hashing approach doesn't take edges into consideration. In some domains, edges are important despite the nodes matching. For example, in computer vision, edges may represent some kind of boundaries and have to be considered in object recognition. (Notice that when applying the proposed geometric hashing approach to this domain, the node label alphabet could be extended to contain semantic information, such as color, shape, etc.) We have done preliminary work in topological graph matching, in which our algorithm matches edge distances and the software is accessible at http://cs.nyu.edu/cs/faculty/shasha/papers/agm.html. We plan to combine geometric matching and topological matching approaches to finding patterns in more general graphs.

ACKNOWLEDGMENTS

The authors would like to thank the anonymous reviewers for their constructive suggestions that helped improve both the quality and the presentation of this paper. We also thank Dr. Carol Venanzi, Bo Chen, and Song Peng for useful discussions and for providing chemical compounds used in the paper. This work was supported in part by US National Science Foundation grants IRI-9531548, IRI-9531554, IIS-9988345, IIS-9988636, and by the Natural Sciences and Engineering Research Council of Canada under Grant No. OGP0046373. Part of this work was done while X. Wang was with the Department of Computer and Information Science, New Jersey Institute of Technology. Part of this work was done while J.T.L. Wang was visiting Courant Institute of Mathematical Sciences, New York University.

REFERENCES

- E.E. Abola, F.C. Bernstein, S.H. Bryant, T.F. Koetzle, and J. Weng, "Protein Data Bank," Data Comm. Int'l Union of Crystallography, pp. 107-132, 1987.
- [2] F.C. Bernstein, T.F. Koetzle, G.J.B. Williams, E.F. Meyer, M.D. Brice, J.R. Rodgers, O. Kennard, T. Shimanouchi, and M. Tasumi, "The Protein Data Bank: A Computer-Based Archival File for Macromolecular Structures," J. Molecular Biology, vol. 112, 1977.
- [3] O. Bourdon and G. Medioni, "Object Recognition Using Geometric Hashing on the Connection Machine," Proc. 10th Int'l Conf. Pattern Recognition, pp. 596-600, 1990.
- [4] D. Conklin, "Knowledge Discovery in Molecular Structure Databases," doctoral dissertation, Dept. Computing and Information Science, Queen's Univ., Canada, 1995.
- [5] D. Conklin, "Machine Discovery of Protein Motifs," Machine Learning, vol. 21, pp. 125-150, 1995.
- [6] D. Conklin, S. Fortier, and J. Glasgow, "Knowledge Discovery in Molecular Databases," *IEEE Trans. Knowledge and Data Eng.*, vol. 5, no. 6, pp. 985-987, 1993.
- [7] L. Dehaspe, H. Toivonen, and R.D. King, "Finding Frequent Substructures in Chemical Compounds," Proc. Fourth Int'l Conf. Knowledge Discovery and Data Mining, pp. 30-36, 1998.
- [8] S. Djoko, D.J. Cook, and L.B. Holder, "An Empirical Study of Domain Knowledge and Its Benefits to Substructure Discovery," IEEE Trans. Knowledge and Data Eng., vol. 9, no. 4, pp. 575-586, 1997
- [9] D. Fischer, O. Bachar, R. Nussinov, and H.J. Wolfson, "An Efficient Automated Computer Vision Based Technique for Detection of Three Dimensional Structural Motifs in Proteins," J. Biomolecular and Structural Dynamics, vol. 9, no. 4, pp. 769-789, 1992.

- [10] H.N. Gabow, Z. Galil, and T.H. Spencer, "Efficient Implementation of Graph Algorithms Using Contraction," Proc. 25th Ann. IEEE Symp. Foundations of Computer Science, pp. 347-357, 1984.
- [11] W.E.L. Grimson, D.P. Huttenlocher, and D.W. Jacobs, "Affine Matching with Bounded Sensor Error: Study of Geometric Hashing and Alignment," Technical Memo AIM-1250, Artificial Intelligence Laboratory, Massachusetts Inst. of Technology, 1991.
- [12] R.W. Hamming "Error Detecting and Error Correcting Codes," *The Bell System Technical J.*, vol. 29, no. 2, pp. 147-160, 1950, Reprinted in E.E. Swartzlander, *Computer Arithmetic*. vol. 2, Los Alamitos, Calif.: IEEE Computer Soc. Press Tutorial, 1990.
- [13] L. Kaufman and P.J. Rousseeuw, Finding Groups in Data: An Introduction to Cluster Analysis. New York: John Wiley & Sons, 1990
- [14] Y. Lamdan and H. Wolfson, "Geometric Hashing: A General and Efficient Model-Based Recognition Scheme," Proc. Int'l Conf. Computer Vision, pp. 237-249, 1988.
- [15] H. Mannila and H. Toivonen, "Levelwise Search and Borders of Theories in Knowledge Discovery," Data Mining and Knowledge Discovery, vol. 1, no. 3, pp. 241-258, 1997.
- Discovery, vol. 1, no. 3, pp. 241-258, 1997.

 [16] J.A. McHugh, Algorithmic Graph Theory. Englewood Cliffs, N.J.: Prentice Hall, 1990.
- [17] X. Pennec and N. Ayache, "An $O(n^2)$ Algorithm for 3D Substructure Matching of Proteins," Proc. First Int'l Workshop Shape and Pattern Matching in Computational Biology, pp. 25-40, 1994.
- [18] C. Pu, K.P. Sheka, L. Chang, J. Ong, A. Chang, E. Alessio, I.N. Shindyalov, W. Chang, and P.E. Bourne, "PDBtool: A Prototype Object Oriented Toolkit for Protein Structure Verification," Technical Report CUCS-048-92, Dept. Computer Science, Columbia Univ., 1992.
- [19] I. Rigoutsos and R. Hummel, "Scalable Parallel Geometric Hashing for Hypercube (SIMD) Architectures," Technical Report TR-553, Dept. Computer Science, New York Univ., 1991.
- [20] I. Rigoutsos and R. Hummel, "On a Parallel Implementation of Geometric Hashing on the Connection Machine," Technical Report TR-554, Dept. Computer Science, New York Univ., 1991.
- [21] I. Rigoutsos, D. Platt, and A. Califano, "Flexible Substructure Matching in Very Large Databases of 3D-Molecular Information," research report, IBM T.J. Watson Research Center, Yorktown Heights, N.Y., 1996.
- [22] J. Sadowski and J. Gasteiger, "From Atoms and Bonds to Three-Dimensional Atomic Coordinates: Automatic Model Builders," Chemical Rev., pp. 2567-2581, vol. 93, 1993.
- [23] B. Sandak, R. Nussinov, and H.J. Wolfson, "A Method for Biomolecular Structural Recognition and Docking Allowing Conformational Flexibility," J. Computational Biology, vol. 5, no. 4, pp. 631-654, 1998.
- [24] B. Sandak, H.J. Wolfson, and R. Nussinov, "Flexible Docking Allowing Induced Fit in Proteins: Insights from an Open to Closed Conformational Isomers," *Proteins*, vol. 32, pp. 159-74, 1998.
- [25] Time Warps, String Edits, and Macromolecules: The Theory and Practice of Sequence Comparison. D. Sankoff and J.B. Kruskal, eds., Reading, Mass.: Addison-Wesley, 1983.
- [26] K.B. Sarachik, "Limitations of Geometric Hashing in the Presence of Gaussian Noise," Technical Memo AIM-1395, Artificial Intelligence Laboratory, Massachusetts Inst. of Technology, 1992.
- [27] R.R. Stoll, *Linear Algebra and Matrix Theory*, New York: McGraw-Hill, 1952.
- [28] I.I. Vaisman, A. Tropsha, and W. Zheng, "Compositional Preferences in Quadruplets of Nearest Neighbor Residues in Protein Structures: Statistical Geometry Analysis," Proc. IEEE Int'l Join Symp. Intelligence and Systems, pp. 163-168, 1998.
- [29] G. Verbitsky, R. Nussinov, and H.J. Wolfson, "Flexible Structural Comparison Allowing Hinge Bending and Swiveling Motions," Proteins, vol. 34, pp. 232-254, 1999.
- [30] J.T.L. Wang, G.-W. Chirn, T.G. Marr, B.A. Shapiro, D. Shasha, and K. Zhang, "Combinatorial Pattern Discovery for Scientific Data: Some Preliminary Results," *Proc. 1994 ACM SIGMOD Int'l Conf. Management of Data*, pp. 115-125, 1994.
- [31] J.T.L. Wang, Q. Ma, D. Shasha, and C.H. Wu, "Application of Neural Networks to Biological Data Mining: A Case Study in Protein Sequence Classification," Proc. Sixth ACM SIGKDD Int'l Conf. Knowledge Discovery and Data Mining, pp. 305–309, 2000.
- [32] J.T.L. Wang, T.G. Marr, D. Shasha, B. A. Shapiro, G.-W. Chirn, and T.Y. Lee, "Complementary Classification Approaches for Protein Sequences," *Protein Eng.*, vol. 9, no. 5, pp. 381-386, 1996.

- [33] J.T.L. Wang, B.A. Shapiro, and D. Shasha, eds., *Pattern Discovery in Biomolecular Data: Tools, Techniques and Applications*, New York: Oxford Univ. Press, 1999.
- [34] X. Wang and J.T.L. Wang, "Fast Similarity Search in Three-Dimensional Structure Databases," *J. Chemical Information and Computer Sciences*, vol. 40, no. 2, pp. 442-451, 2000.
- [35] X. Wang, J.T.L. Wang, D. Shasha, B.A. Shapiro, S. Dikshitulu, I. Rigoutsos, and K. Zhang, "Automated Discovery of Active Motifs in Three Dimensional Molecules," Proc. Third Int'l Conf. Knowledge Discovery and Data Mining, pp. 89-95, 1997.
- [36] K. Zhang, J.T.L. Wang, and D. Shasha, "On the Editing Distance Between Undirected Acyclic Graphs," Int'l J. Foundations of Computer Science, special issue on Computational Biology, vol. 7, no. 1, pp. 43-57, 1996.



Xiong Wang received the MS degree in computer science from Fudan University, China, and the PhD degree in computer and information science from the New Jersey Institute of Technology, in 2000 (thesis advisor: Jason T.L. Wang). He is currently an assistant professor of computer science in California State University, Fullerton. His research interests include databases, knowledge discovery and data mining, pattern matching, and bioinfor-

matics. He is a member of ACM, ACM SIGMOD, IEEE, and IEEE Computer Society.



Jason T.L. Wang received the BS degree in mathematics from National Taiwan University, and the PhD degree in computer science from the Courant Institute of Mathematical Sciences, New York University, in 1991 (thesis advisor: Dennis Shasha). He is a full professor in the Computer and Information Science Department at the New Jersey Institute of Technology and director of the University's Data and Knowledge Engineering Laboratory. Dr. Wang's research

interests include data mining and databases, pattern recognition, bioinformatics, and Web information retrieval. He has published more than 100 papers, including 30 journal articles, in these areas. He is an editor and author of the book *Pattern Discovery in Biomolecular Data* (Oxford University Press), and co-author of the book *Mining the World Wide Web* (Kluwer). In addition, he is program cochair of the 2001 Atlantic Symposium on Computational Biology, Genome Information Systems & Technology held at Duke University. He is a member of the IEEE



Dennis Shasha received the PhD degree from Harvard in 1984 (thesis advisor: Nat Goodman). He is a professor at Courant Institute, New York University, where he does research on biological pattern discovery, combinatorial pattern matching on trees and graphs, software support for the exploration of unfamiliar databases, database tuning, and database design for time series. After graduating from Yale in 1977, he worked for IBM designing circuits and microcode for the

3090. He has written a professional reference book *Database Tuning: A Principled Approach*, (1992, Prentice-Hall), two books about a mathematical detective named Dr. Ecco entitled *The Puzzling Adventures of Dr. Ecco*, (1988, Freeman, and republished in 1998 by Dover) and *Codes, Puzzles, and Conspiracy*, (1992, Freeman) and a book of biographies about great computer scientists called *Out of Their Minds: The Lives and Discoveries of 15 Great Computer Scientists*, (1995, Copernicus/Springer-Verlag). Finally, he is a co-author of *Pattern Discovery in Biomolecular Data: Tools, Techniques, and Applications*, published in 1999 by Oxford University Press. In addition, he has co-authored more than 30 journal papers and 40 conference papers and spends most of his time in building data mining and pattern recognition software these days. He writes a monthly puzzle column for *Scientific American*.



Bruce A. Shapiro received the BS degree from Brooklyn College studying mathematics and physics, the MS and PhD degrees in computer science from the University of Maryland, College Park. He is a principal investigator and computer specialist in the Laboratory of Experimental and Computational Biology in the National Cancer Institute, National Institutes of Health. Dr. Shapiro directs research on computational nucleic acid structure prediction and analysis

and has developed several algorithms and computer systems related to this area. His interests include massively parallel genetic algorithms, molecular dynamics, and data mining for molecular structure/function relationships. He is the author of numerous papers on nucleic acid morphology, parallel computing, image processing, and graphics. He is an editor and author of the book *Pattern Discovery in Biomolecular Data*, published by Oxford University Press in 1999. In addition, he is a lecturer in the Computer Science Department at the University of Maryland, College Park.

Isidore Rigoutsos received the BS degree in physics from the University of Athens, Greece, the MS and PhD degrees in computer science from the Courant Institute of Mathematical Sciences of New York University. He manages the Bioinformatics and Pattern Discovery group in the Computational Biology Center of the Thomas J. Watson Research Center. Currently, he is visiting lecturer in the Department of Chemical Engineering of the Massachusetts Institute of Technology. He has been the recipient of a Fulbright Foundation Fellowship and has held an adjunct professor appointment in the Computer Science Department of New York University. He is the author or co-author of eight patents and numerous technical papers. Dr. Rigoutsos is a member of the IEEE, the IEEE Computer Society, the International Society for Computational Biology, and the American Association for the Advancement of Science.

Kaizhong Zhang received the MS degree in mathematics from Peking University, Beijing, People's Republic of China, in 1981, and the MS and PhD degrees in computer science from the Courant Institute of Mathematical Sciences, New York University, New York, in 1986 and 1989, respectively. Currently, he is an associate professor in the Department of Computer Science, University of Western Ontario, London, Ontario, Canada. His research interests include pattern recognition, computational biology, image processing, sequential, and parallel algorithms.

⊳ For more information on this or any computing topic, please visit our Digital Library at http://computer.org/publications/dlib.