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PAIRWISE PROTEIN STRUCTURE ALIGNMENT BASED ON AN ORIENTATION-INDEPENDENT BACKBONE REPRESENTATION

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Determining structural similarities between proteins is an important problem since it can help identify functional and evolutionary relationships. In this paper, an algorithm is proposed to align two protein structures. Given the protein backbones, the algorithm finds a rigid motion of one backbone onto the other such that large substructures are matched. The algorithm uses a representation of the backbones that is independent of their relative orientations in space and applies dynamic programming to this representation to compute an initial alignment, which is then refined iteratively. Experiments indicate that the algorithm is competitive with two well-known algorithms, namely DALI and LOCK.

Keywords: Protein structure alignment; backbone representation; dynamic programming; optimization.

1. Introduction

Three-dimensional (3D) structure plays a central role in research directed towards understanding evolutionary and functional relationships among proteins. It is wellknown that structural information is better conserved than sequence information in the evolution of proteins,¹⁴ hence it can be used in the construction of phylogenetic trees.¹⁶ Protein-protein interactions are governed in large part by the shape, location, and composition of the so-called active sites.² The assignment of proteins to fold families is accomplished via structural analysis.^{15,18} The need for effective structural analysis techniques has increased with the rapid growth in the number of 3D structures available now in the Protein Data Bank (PDB).¹

A key problem in protein structure analysis is pairwise protein structure alignment: Given the C_{α} backbones of two proteins, the goal is to find a rigid motion of one backbone onto the other such that large, contiguous regions of the backbones are matched. (A formal definition is given later in Sec. 2.) Besides the applications mentioned above, pairwise structural alignment is also a key component of algorithms that seek to align multiple protein structures in order to find a core structure that captures essential structural information for the whole set.⁴

Pairwise alignment of protein structures has been a subject of much research. Holm and Sander^{13,14} propose an algorithm, called DALI, which works with the distance matrices obtained from the interatomic $(C_{\alpha}-C_{\alpha})$ distances on each backbone. The algorithm decomposes each matrix into submatrices that represent so-called elementary contact patterns, aligns a pair of patterns from the two matrices, and iteratively builds a connected chain of such pairwise aligned patterns using a Monte Carlo method to optimize the similarity score. Singh and Brutlag²⁰ give an algorithm, called LOCK, which represents the secondary structure elements $(\alpha$ -helices and β -strands) as vectors and computes a local alignment of these via dynamic programming. A suite of seven different scoring functions is used to score the alignment. This yields an initial superposition, which is then improved iteratively by operating at the atomic level, on the 3D coordinates of the C_{α} atoms, until the root-mean squared deviation (RMSD)⁶ of the aligned atoms converges. Chew et $al.^3$ propose an approach which represents the backbone by a chain of unit-vectors. These vectors are then translated to the origin, yielding a representation of the protein as a set of points on the unit-sphere. To compute a structural alignment, they first compute a small set of shifts in sequence space that are likely to bring 3D structures into correspondence. For each such shift, they compute substructures that are contiguous on the backbone and for which there is a 3D alignment, and then combine such substructures into large (possibly non-contiguous) substructures called domains. They use a variant of the RMSD measure, called unit-vector RMSD, which is robust to outliers.

A non-exhaustive list of other related work includes the combinatorial extension (CE) method of Shindyalov and Bourne,¹⁹ a method based on geometric hashing by Fischer, Nussinov, and Wolfson,⁹ the method of Falicov and Cohen⁷ which attempts

to minimize the so-called soap-bubble surface area between the backbones, and the double-dynamic programming technique of Orengo and Taylor.^{21,22}

In this paper, we propose a new approach to the pairwise structural alignment problem. Our algorithm computes a representation of each backbone in terms of certain angles defined by consecutive C_{α} - C_{α} bonds. The resulting backbone representation, which consists of a sequence of triples of angles, is independent of the relative orientation of the two proteins in space. We apply dynamic programming on this representation (not on the protein sequence) and compute an initial alignment of the two proteins. We then refine this alignment iteratively. Our algorithm takes time that is quadratic in the sum of the lengths of the two proteins. We have implemented this alignment and tested it against LOCK and DALI, on various protein data sets. We have found that our algorithm is quite competitive with these algorithms, as discussed in detail in Sec. 4.

We note that our angle-based representation is related to the pseudodihedral angle used by Dewitte and Shakhnovich.⁵ The pseudodihedral angle is the torsion angle between planes defined by four consecutive C_{α} atoms. Indeed, as stated by Dewitte and Shakhnovich,⁵ the pseudodihedral angle provides a simplified backbone representation that manifests information about secondary structure elements. Moreover, the distribution of pseudodihedral angles is highly correlated to the identity of the central pair of amino acids. These observations provide empirical support for our use of a local angle-based representation of the backbone to compute pairwise alignments. The efficacy of this representation is also borne out by our experimental results.

The rest of the paper is organized as follows. Section 2 gives an overview of our method. Section 3 describes in detail the five steps that comprise our algorithm. We report on experimental results in Sec. 4 and conclude in Sec. 5.

2. Overview of the Approach

Let A and B be the two proteins under consideration, each represented by a chain of C_{α} atoms (the backbone) in \mathbb{R}^3 . (As is customary,^{13,20} we consider only the backbone, not the amino acid residues themselves.) Assuming B is fixed in space, we would like to find a rigid motion (translation and rotation) of A that aligns large substructures of A and B. Specifically, we would like to find substructures A' of A and B' of B, a bijection between atoms of A' and B', which preserves their order on the backbones, and a rigid motion of A onto B such that |A'| = |B'| is as large as possible and the Euclidean distance between atom pairs in the bijection is at most a user-specified threshold ε . We present a heuristic for this problem which is very competitive with other known methods.

The key to our approach is a geometric representation of the backbone structure that is independent of the relative orientations of the proteins A and B. Let the C_{α} atoms of A and B be numbered 1 through n and 1 through m, respectively, in order along the backbone. In effect, the geometric representation of A (respectively B) may be viewed as a collection $\{A_i\}_{i=2}^{n-2}$ (respectively $\{B_i\}_{i=2}^{m-2}$) of points in \mathbb{R}^3 , where each point A_i (respectively B_i) captures the local geometry of the (virtual) bond joining the *i*th and (i + 1)th C_{α} atoms of A (respectively B) in relation to the preceding and succeeding bond on the backbone. A major advantage of this representation is that it is independent of the relative positions and orientations of the backbones. Indeed, as we will see in Sec. 3, the question of whether there exists a rigid motion of A that aligns it to B can be framed as the question of determining the similarity of the (static) point-sets representing A and B. In particular, A can be aligned exactly with B if and only if the sets $\{A_i\}_{i=2}^{n-2}$ and $\{B_i\}_{i=2}^{m-2}$ are identical.

In practice, of course, an exact alignment is unlikely to exist; instead the goal is to align approximately one or more substructures of A to substructures of B. Towards this end, we use dynamic programming to compute an optimal global alignment of the sequences A_2, \ldots, A_{n-2} and B_2, \ldots, B_{m-2} , using the Euclidean distance between the A_i 's and B_j 's as the scoring function. The resulting alignment consists of runs of matches between elements of the two sequences, with gaps in between. (See Fig. 1.)

However, it may not be possible to simultaneously align, in 3D, all of the substructures corresponding to the different runs of matches found in the sequence alignment. This is because the transformation matrix that aligns the structures corresponding to one run may not be consistent with the matrix for another run. For instance, in Fig. 1, the matrix that aligns atoms 1-5 in A with atoms 1-5 in Bmay be different from the one that aligns atoms 9-13 in A with atoms 5-9 in B.

To overcome this problem, we compute a subset of the runs such that the transformation matrices for any two runs in the subset are similar (as measured by the



Fig. 1. A local alignment of the sequences A_2, \ldots, A_{19} and B_2, \ldots, B_{13} and the corresponding matched substructures of proteins A and B, shown in solid lines of varying thickness. Dashed lines denote unmatched substructures.

Frobenius norm¹¹) and the total number of atoms in the runs is large. We show how this can be reduced to the problem of finding a large weighted clique in an undirected graph and solve this using a greedy heuristic. For the subset of consistent runs thus found, we compute a transformation matrix that aligns the corresponding substructures. This alignment is further improved via dynamic programming on the coordinates of the C_{α} atoms, during which pairs that are not within the threshold of ε are discarded. This yields an initial alignment of the proteins. The closeness of the overall alignment is measured by the RMSD.⁶

The algorithm then enters an iterative refinement phase. From the matches in the initial alignment, a new transformation matrix is derived. This is used to realign the proteins via dynamic programming, as above. The process is repeated until the difference in the RMSD values of two successive alignments drops below a preset threshold.

3. Details of the Algorithm

3.1. Step 1: Computing a local geometric representation

Recall that protein A consists of $n C_{\alpha}$ atoms, numbered $1, \ldots, n$ along the backbone. We define a sequence of vectors \vec{a}_i , $1 \leq i \leq n-1$ on the backbone, where \vec{a}_i is the vector from the *i*th C_{α} atom to the (i + 1)th C_{α} atom. Each \vec{a}_i has the same length as the corresponding (virtual) bond; this is about 3.8 Å. We represent the geometry of the backbone in the vicinity of $\vec{a}_i, 2 \leq i \leq n-2$ by a triple of angles $A_i = (\alpha_i^A, \beta_i^A, \gamma_i^A)$. Here $\alpha_i^A \in [0, \pi]$ is the angle between $-\vec{a}_{i-1}$ and \vec{a}_i and $\beta_i^A \in [0, \pi]$ is the angle between $-\vec{a}_i$ and \vec{a}_{i+1} . Both of these angles can be computed via dot products of the corresponding vectors. We define γ_i^A as follows: Consider the vectors $\vec{n}_i = -\vec{a}_{i-1} \times \vec{a}_i$ and $\vec{n}_{i+1} = -\vec{a}_i \times \vec{a}_{i+1}$. Note that \vec{a}_i is perpendicular to both \vec{n}_i and \vec{n}_{i+1} . Let $\theta \in [0, 2\pi]$ be the angle between \vec{n}_i and \vec{n}_{i+1} . Then $\gamma_i^A = \theta$ if $\vec{n}_i \times \vec{n}_{i+1}$ has the same direction as \vec{a}_i ; otherwise $\gamma_i^A = 2\pi - \theta$. In other words, γ_i^A is the dihedral angle between the plane containing $-\vec{a}_{i-1}$ and \vec{a}_i and the plane containing $-\vec{a}_i$ and \vec{a}_{i+1} (Fig. 2 illustrates these angles). Thus, $A_i = (\alpha_i^A, \beta_i^A, \gamma_i^A)$ captures the orientation of \vec{a}_i relative to its predecessor \vec{a}_{i-1} and its successor \vec{a}_{i+1} . The set $\{A_i\}_{i=2}^{n-2}$ is the local geometric representation of the backbone A. We can define similarly a sequence of vectors \vec{b}_i , $2 \leq i \leq m-2$, for the backbone of protein B, and a triple $B_i = (\alpha_i^B, \beta_i^B, \gamma_i^B)$ for \vec{b}_i . The local geometric representation of B is then the set $\{B_i\}_{i=2}^{m-2}$.

The following lemma establishes the orientation independence of this representation.

Lemma 1. Let A' and B' be substructures of A and B, respectively, where A' contains the pth through $(p + \ell)$ th C_{α} atoms of A and B' contains the qth through $(q + \ell)$ th C_{α} atoms of B, $\ell \geq 3$, $1 \leq p \leq n - \ell$, and $1 \leq q \leq m - \ell$. Then there is a rigid motion (translation and rotation) of A to B which aligns exactly the (p + j)th C_{α} atom of A with the (q + j)th C_{α} atom of B if and only if $||A_{p+i} - B_{q+i}||_2 = 0$,



Fig. 2. Vector representation of backbone of protein A and associated angles. The triple $A_i = (\alpha_i^A, \beta_i^A, \gamma_i^A)$ is associated with the vector $\vec{a}_i, 2 \le i \le n-2$.

where $0 \leq j \leq \ell$ and $1 \leq i \leq \ell - 2$. (Here $||A_{p+i} - B_{q+i}||_2$ is the Euclidean distance between A_{p+i} and B_{q+i} when they are viewed as points in \mathbb{R}^3 .)

Proof. (\Rightarrow) Obvious.

(\Leftarrow) Consider first the case $\ell = 3$. W.l.o.g., we may assume that the qth, (q+1)th, and (q+2)th C_{α} atoms of B lie in the xy-plane, with the (q+1)th atom at the origin, and the qth atom on the positive x-axis. Clearly, we can transform A so that the pth, (p+1)th, and (p+2)th C_{α} atoms of A also lie on the xy-plane, with the (p+1)th atom at the origin and the pth atom on the positive x-axis. Since \vec{a}_p and \vec{b}_q have the same length, \vec{a}_{p+1} and \vec{b}_{q+1} have the same length, and $\alpha_{p+1}^A = \alpha_{q+1}^B$, it follows that the pth and qth C_{α} atoms coincide and (p+2)th and (q+2)th C_{α} atoms coincide.

Now, $\gamma_{p+1}^A = \gamma_{q+1}^B$ implies that the (p+3)th and (q+3)th C_{α} atoms of A and B, respectively, lie in a common half-plane. (This half-plane makes an angle of γ_{p+1}^A with the xy-plane and its intersection with the xy-plane is the line containing the (coincident) vectors \vec{a}_{p+1} and \vec{b}_{q+1} .) Then since $\beta_{p+1}^A = \beta_{q+1}^B$ and \vec{a}_{p+2} and \vec{b}_{q+2} have the same length, it follows that the (p+3)th and (q+3)th C_{α} atoms coincide. This completes the proof for the case $\ell = 3$.

Next, let $\ell > 3$. The argument above shows that once the first three pairs of atoms of A' and B' coincide, then the fourth pair also coincides. But now since the second through fourth pairs of atoms of A' and B' coincide, by a similar argument, the fifth pair must also coincide. This argument can be repeated until the $(p + \ell)$ th and $(q + \ell)$ th atoms are shown to coincide.

3.2. Step 2: Aligning the local representation

As mentioned in Sec. 2, a perfect alignment between A and B (Lemma 1) is unlikely to exist. Instead we try to align substructures of A and B. Towards this end, we treat the sets $\{A_i\}_{i=2}^{n-2}$ and $\{B_i\}_{i=2}^{m-2}$ as sequences A_2, \ldots, A_{n-2} and B_2, \ldots, B_{m-2} and compute an optimal global alignment for them using dynamic programming.¹² This alignment is produced using a global alignment method with zero end-gap penalties. Internal gaps are penalized using an affine gap penalty of the form: h(k) = a + bk, where k is the length of the gap, and a and b are gap opening and extension penalties, respectively. We score the alignment using a scoring function S_0 , defined as:

$$S_0(A_i, B_j) = K - d(A_i, B_j).$$
(1)

Here $d(A_i, B_j)$ is the Euclidean distance between the points $A_i = (\alpha_i^A, \beta_i^A, \gamma_i^A)$ and $B_j = (\alpha_j^B, \beta_j^B, \gamma_j^B)$. (A similar function is used in DALI.¹³) Our goal is to compute a global alignment whose total score is maximum. It is tempting to write

$$d(A_i, B_j) = \left((\alpha_i^A - \alpha_j^B)^2 + (\beta_i^A - \beta_j^B)^2 + (\gamma_i^A - \gamma_j^B)^2 \right)^{1/2}.$$
 (2)

However, if one of the angles γ_i^A and γ_j^B is close to zero and the other one is close to 2π , then A_i and B_j are actually close to each other in the γ -dimension, even though $|\gamma_i^A - \gamma_j^B|$ is large (i.e. close to 2π). Thus we modify $d(A_i, B_j)$ as follows:

$$d(A_i, B_j) = \left((\alpha_i^A - \alpha_j^B)^2 + (\beta_i^A - \beta_j^B)^2 + g(|\gamma_i^A - \gamma_j^B|)^2 \right)^{1/2}$$
(3)

where $g(x) = \min(2\pi - x, x)$.

What about the term K in the definition of S_0 and the gap opening and extension penalties a and b? We have done several experiments and found that K = 1.4, a = 0.2, and b = 0.2 work very well, and this is what we use throughout.

3.3. Step 3: Computing consistent runs of alignments

The alignment computed in step 2 yields runs of matched elements of A_2, \ldots, A_{n-2} and B_2, \ldots, B_{m-2} , interspersed with gaps (see Fig. 1). Let R_1, \ldots, R_k be the runs and let N_1, \ldots, N_k be their lengths. For each R_i , we can compute a transformation matrix $T_i = (T_i^t, T_i^r)$ that aligns the substructures of A and B corresponding to R_i such that the RMSD is minimized. Here T_i^t is the translation matrix and T_i^r is the rotation matrix. T_i can be computed using the Singular Value Decomposition (SVD).^{11,17} However, the matrices T_1, T_2, \ldots, T_k will not necessarily be mutually consistent, in the sense that the transformation specified by one matrix T_i may "interfere" with that specified by another matrix T_j , so that it may not be possible to align simultaneously the structures corresponding to all the runs R_1, \ldots, R_k .

Therefore, in this step, we compute a subset of the runs such that the total number of atoms in the runs is as large as possible and the transformation matrices for all the runs are "similar". Formally, we wish to compute a subset I of $\{1, \ldots, k\}$ such that $\sum_{i \in I} |R_i|$ is as large as possible and for all $i, j \in I$, $||T_i^t - T_j^t||_F < \tau^t$ and $||T_i^r - T_j^r||_F < \tau^r$. Here τ^t and τ^r are similarity thresholds and $||C||_F = (\sum_{i=1}^3 \sum_{j=1}^3 c_{ij}^2)^{1/2}$ is the Frobenius norm of matrix $C = (c_{ij})$.¹¹

To compute *I*, we first build an undirected graph G = (V, E), where $V = \{1, 2, ..., k\}$ and $E = \{(i, j) \in V \times V : ||T_i^t - T_j^t||_F < \tau^t$ and $||T_i^r - T_j^r||_F < \tau^r\}$. With each vertex *i* of *G*, we associate a weight $w_i = |R_i|$. Clearly, our problem now is equivalent to finding a clique (complete subgraph) of *G* whose total vertex weight is maximum. This problem is NP-hard,¹⁰ so we solve it approximately via a greedy heuristic, as follows. Among all the vertices of *G*, we find a vertex, *v*, such that the total weight of *v* and all its neighbors in *G* is maximum. We add *v* to an initially empty list *L*, which accumulates the growing clique. We then repeat the above step on the subgraph, *G'*, of *G* induced by the neighbors of *v*. That is, among the vertices of *G'*, we find a vertex, *v'*, such that the total weight of *v'* and its neighbors in *G'* is maximum, and add *v'* to *L*. And so on, until at some point the current induced subgraph becomes empty. At this point, the vertices in *L* form a clique of *G* of large (but not necessarily maximum) total weight.

What should be the values of the similarity thresholds τ^t and τ^r ? If τ^t and τ^r are very small, the matrix T_i and T_j are required to be very similar; this yields a subset of runs of small total size. If τ^t and τ^r are large then the matrices can be quite different, so the quality of the structural alignment is poor. We found via experiments that choosing τ^t between 10 and 40, and τ^r between 1 and 1.5 works very well. Our experiments also show that τ^r is much more sensitive than τ^t . The degree of similarity of two rotation matrices is determined by the threshold τ^r . As shown in Lemma 2 below, the expected value of the Frobenius norm of the difference of two rotation matrices whose elements are chosen randomly is about 2.4. Thus our choice of $\tau^r \in [1, 1.5]$ ensures that the associated matrices are all quite similar and not random.

Lemma 2. The expected value of the Frobenius norm of the matrix that is the difference of two rotation matrices whose elements are chosen randomly is about 2.4.

Proof. Let C and D be two rotation matrices. Assume $C = M_z M_y M_x$, where M_x rotates by an angle c_x about the x-axis, M_y rotates by an angle c_y about the y-axis, and M_z rotates by an angle c_z about the z-axis. Similarly, let $D = N_z N_y N_x$, where N_x , N_y , and N_z rotate by angles d_x , d_y , and d_z about the x-, y-, and z-axes, respectively.

It follows that

$$C - D$$

$$= \begin{pmatrix} \cos(c_z) & -\sin(c_z) & 0\\ \sin(c_z) & \cos(c_z) & 0\\ 0 & 0 & 1 \end{pmatrix} \begin{pmatrix} \cos(c_y) & 0 & \sin(c_y)\\ 0 & 1 & 0\\ -\sin(c_y) & 0 & \cos(c_y) \end{pmatrix} \begin{pmatrix} 1 & 0 & 0\\ 0 & \cos(c_x) & -\sin(c_x)\\ 0 & \sin(c_x) & \cos(c_x) \end{pmatrix} \\ - \begin{pmatrix} \cos(d_z) & -\sin(d_z) & 0\\ \sin(d_z) & \cos(d_z) & 0\\ 0 & 0 & 1 \end{pmatrix} \begin{pmatrix} \cos(d_y) & 0 & \sin(d_y)\\ 0 & 1 & 0\\ -\sin(d_y) & 0 & \cos(d_y) \end{pmatrix} \begin{pmatrix} 1 & 0 & 0\\ 0 & \cos(d_x) & -\sin(d_x)\\ 0 & \sin(d_x) & \cos(d_x) \end{pmatrix}.$$

Since each of the six rotation angles is chosen at random from $[0, 2\pi]$, the expected value of $||C - D||_F$ then is

$$\iiint \iiint \int_{0 \le c_x, c_y, c_z, d_x, d_y, d_z \le 2\pi} \|C - D\|_F \frac{1}{(2\pi)^6} d(c_x) d(c_y) d(c_z) d(d_x) d(d_y) d(d_z).$$

Performing this calculation, via numerical integration in MATLAB, yields an expected value of about 2.3958.

We also computed the expected value experimentally using MATLAB, by picking 40,000 pairs of rotation matrices whose elements are chosen at random and computing the Frobenius norm of the difference of each pair. This yielded an expected value of 2.3982.

3.4. Step 4: Computing an initial structural alignment

The previous step yields a subset of runs with pairwise similar transformation matrices. This means that there is now a single transformation matrix T, which can simultaneously align well the substructures corresponding to all runs in the subset. We compute T by SVD and use this to transform protein A and obtain an initial alignment with protein B.

Recall that we also require that each aligned pair of atoms from A and B to be within distance ε of each other. One possibility is to take the alignment provided by T and simply discard any pairs that do not meet the ε threshold. However, a large number of pairs that are close to the threshold could get discarded. A better strategy is to first re-align A and B by using dynamic programming on the coordinates just computed by the application of T. In the process, we also enforce the ε threshold automatically by choosing the gap penalty suitably.

Specifically, we compute a minimum-score global alignment of A and B.¹² We score the alignment of atom $i \in A$ and atom $j \in B$ using the Euclidean distance d(i, j) between them. The score for matching a gap with an atom (i.e., the gap penalty) is a constant equal to $\varepsilon/2$. (In the experiments reported in Sec. 4, we used $\varepsilon = 8$ Å.) At the end of this step, we get an initial structural alignment, \mathcal{I} , of A and B such that its score $s(\mathcal{I})$ is minimum. We then compute its RMSD.

We argue now that \mathcal{I} satisfies the ε threshold. Suppose, for a contradiction, that, in \mathcal{I} , atoms $i \in A$ and $j \in B$ are aligned but that $d(i, j) > \varepsilon$. We modify \mathcal{I} locally by inserting two gaps as shown in Fig. 3 to get a new alignment \mathcal{I}' . Since



Fig. 3. Local insertion of gaps in \mathcal{I} .

the gap penalty is $\varepsilon/2$, $s(\mathcal{I}') = s(\mathcal{I}) - d(i, j) + \varepsilon/2 + \varepsilon/2 < s(\mathcal{I})$, which contradicts the optimality of \mathcal{I} .

3.5. Step 5: Refining the alignment iteratively

Our experiments have shown that the size of the alignment (i.e. the number of matched pairs) can be increased as follows. For the initial alignment computed in step 4, we recompute a new transformation matrix T' corresponding to all of the matched pairs in the initial alignment, again using SVD. We then use dynamic programming as in step 4 to obtain a new alignment, with a new RMSD value.

We iterate on the above transform-and-realign process until one of the following two conditions is met: Either the number of iterations exceeds a specified limit or the absolute value of the difference in RMSD values of two successive alignments drops below a preset threshold η . (Recall that the RMSD of an alignment \mathcal{I} containing matched pairs (i, j) of atoms, where $i \in A$ and $j \in B$ have coordinates (x_i, y_i, z_i) and (x_j, y_j, z_j) , respectively, is $(\frac{1}{|\mathcal{I}|} \sum_{(i,j) \in \mathcal{I}} ((x_i - x_j)^2 + (y_i - y_j)^2 + (z_i - z_j)^2))^{1/2}$.) In our experiments, we limited the number of iterations to ten and chose $\eta = 0.1$ Å.

Complexity analysis: Step 1 clearly takes O(m + n) time. Step 2 takes O(mn) time, and produces an alignment of length at most m + n. In step 3, the computation of T_i , using the SVD, takes time $O(N_i)$, so the total time for all the T_i 's is $O(\sum_{i=1}^k N_i) = O(m + n)$. Computing the clique takes the time proportional to the size of the graph G, which is $O((m + n)^2)$, since G has at most m + n vertices. In step 4, the computation of the transformation matrix T, via the SVD, takes O(m + n) time, and the dynamic programming takes an additional $O((m + n)^2)$ time. In step 5, each iteration takes $O((m + n)^2)$ time, similar to step 4. Since the number of iterations is bounded by a constant (ten in our experiments), the total time for step 5 is $O((m + n)^2)$. Thus, the overall running time of the algorithm is $O((m + n)^2)$.

4. Experimental Results

We implemented our algorithm (using MATLAB) and tested it against two wellknown structural alignment algorithms, namely LOCK²⁰ and DALI¹³. The code for our algorithm, all the test datasets, and color versions of all the figures in this paper may be accessed at http://www.cs.umn.edu/~jieping/Research.html.

Our first experiment was similar to that done for LOCK,²⁰ in that we aligned a query protein with each member of a set of proteins identified from the FSSP¹⁴ and SCOP¹⁸ databases as structural neighbors of the query.

The first query protein was the protein *Sperm whale myoglobin* (PDB ID: 1mbc), from the Globin family. The results of the alignment (i.e., the number of matched atoms and the RMSD value) are shown in Table 1. Note that the upper half of

Protein		NEW			OCK	DALI	
PDB ID	PDB Header (partial)	#	RMSD	#	RMSD	#	RMSD
5mbn	MYOGLOBIN (DEOXY)	153	0.48	152	0.30	153	0.50
1mbn	MYOGLOBIN (FERRIC IRON)	153	0.48	152	0.47	151	0.50
1myh-A	MYOGLOBIN (AQUOMET, PH 7.1)	153	0.56	153	0.56	139	0.50
1hds-B	HEMOGLOBIN (SICKLE CELL)	145	1.62	130	1.26	145	1.70
2dhb-A	HEMOGLOBIN (HORSE, DEOXY)	141	1.53	133	1.34	140	1.60
1mba	MYOGLOBIN (MET)	143	1.86	124	1.49	142	1.90
1dm1	MYOGLOBIN	143	1.90	130	1.54	143	2.00
1hlm	HEMOGLOBIN (CYANO-MET)	144	2.05	124	1.50	144	2.20
2lhb	HEMOGLOBIN V (CYANO, MET)	137	1.49	129	1.10	135	1.40
2fal	MYOGLOBIN (FERRIC)	143	1.85	122	1.43	143	2.00
1hbg	HEMOGLOBIN (CARBON MONOXY)	139	1.83	125	1.41	140	2.10
1ith-A	HEMOGLOBIN (CYANOMET)	139	1.66	126	1.30	139	1.70
1flp	HEMOGLOBIN I (MONOMERIC)	138	1.63	129	1.35	137	1.70
1eca	HEMOGLOBIN (ERYTHROCRUORIN)	136	1.62	131	1.50	136	1.70
2hbg	HEMOGLOBIN (DEOXY)	139	1.84	125	1.42	138	2.00
1ash	HEMOGLOBIN (DOMAIN ONE)	138	1.86	122	1.43	138	1.90
1hbi-B	HEMOGLOBIN I (OXYGENATED)	135	1.77	122	1.46	141	2.20
1gdi	LEGHEMOGLOBIN	144	2.14	115	1.53	144	2.60
1hlb	HEMOGLOBIN (SEA CUCUMBER)	142	2.18	121	1.48	145	2.50
1lh2	LEGHEMOGLOBIN (AQUO,MET)	143	2.23	120	1.49	144	2.40
1h97-A	HEMOGLOBIN	141	2.18	110	1.48	141	2.20
1dly-A	HEMOGLOBIN	111	2.68	86	1.69	110	2.70
1idr-A	HEMOGLOBIN HBN	107	2.63	84	1.74	106	2.70
1dlw-A	HEMOGLOBIN	107	2.90	80	1.72	106	2.70
1all-A	ALLOPHYCOCYANIN	117	2.95	79	1.69	121	3.50
1phn-A	PHYCOCYANIN	117	3.01	82	1.78	119	3.30
1cpc-A	C-PHYCOCYANIN	113	3.19	80	1.79	117	3.20
1lia-A	R-PHYCOERYTHRIN	117	3.10	73	1.73	120	3.30
1cpc-B	C-PHYCOCYANIN	118	3.29	73	1.70	120	3.80
1qgw-C	CRYPTOPHYTAN PHYCOERYTHRIN	110	3.59	66	1.95	125	4.00
1lia-B	R-PHYCOERYTHRIN	113	3.54	59	1.71	121	3.90
1col-A	COLICIN *A (C-TERMINAL)	88	3.69	61	1.82	113	3.20
2cp4	CYTOCHROME P450CAM	62	4.31	50	1.95	83	4.40
1eum-A	FERRITIN 1	75	4.13	32	1.76	83	7.00
1fpo-A	CHAPERONE PROTEIN HSCB	71	3.74	18	1.72	80	4.20
1oxa	CYTOCHROME P450 ERYF	70	4.01	57	1.85	79	4.60
1le2	APOLIPOPROTEIN-*E2	75	4.13	46	1.71	71	4.90
2fha	FERRITIN	70	3.48	22	1.36	84	6.80
1nfn	APOLIPOPROTEIN E3 FRAG	60	2.88	48	1.59	75	7.90
1grj	GREA TRANSCRIPT CLEAVAGE	55	4.37	27	0.92	62	6.20

Table 1. Query protein: 1mbc (Sperm whale myoglobin).

Table 1 contains proteins that are in the same family as the query, hence closelyrelated, whereas the lower half contains proteins that are in different families, but still related. (Nearly the same set was used for LOCK²⁰ also.) The results of Table 1 are shown graphically in Fig. 4. Note that here the proteins are listed on the x-axis in non-increasing order of the matches found by our method. The alignment of 1mbc with each of the proteins 5mbn, 2fal, and 1lia-A is shown in Fig. 6 (left half).



Fig. 4. Comparison between the proposed method (NEW), LOCK, and DALI using query protein 1mbc. (Data used is from Table 1.) The proteins on the x-axis are ordered by non-increasing number of matches in the NEW method.

The second query protein that we used was *Thioredoxin-Reduced Form* (PDB ID: 3trx), from the Thioltransferase family. The results for this are shown in Table 2 and Fig. 5, and sample alignments are illustrated in Fig. 6 (right half).

These results show that when the proteins in question are closely related, then all three methods (ours, LOCK, and DALI) are able to detect large structural matches.

Protein		Ν	IEW	LOCK		D	ALI
PDB ID	PDB Header (partial)	#	RMSD	#	RMSD	#	RMSD
4trx	THIOREDOXIN (REDUCED FORM)	105	0.40	105	0.41	105	0.40
1mdi-A	THIOREDOXIN MUTANT	105	0.73	105	0.73	105	1.70
1aiu	HUMAN THIOREDOXIN (MUTANT)	104	1.11	102	0.96	105	1.30
1erv	THIOREDOXIN	104	1.11	103	0.97	105	1.30
1f9m-B	THIOREDOXIN F	103	1.36	101	1.28	104	1.50
1f9m-A	THIOREDOXIN F	103	1.38	100	1.25	104	1.50
1gh2-A	THIOREDOXIN-LIKE PROTEIN	104	1.27	101	1.14	105	1.50
1ep7-A	THIOREDOXIN CH1, H-TYPE	104	1.35	101	1.66	105	1.60
1ep7-B	THIOREDOXIN CH1, H-TYPE	104	1.39	98	1.15	105	1.60
1faa	THIOREDOXIN F	103	1.43	100	1.28	104	1.60
1tof	THIOREDOXIN H	105	1.71	96	1.37	104	1.70
2tir	THIOREDOXIN MUTANT	101	1.44	96	1.26	103	1.80
1thx	THIOREDOXIN-2	101	1.56	95	1.30	103	1.80
1fb6-B	THIOREDOXIN M	100	1.37	97	1.20	102	1.70
1quw	THIOREDOXIN	101	1.94	89	1.47	102	2.10
1kte	THIOLTRANSFERASE	86	3.54	47	1.94	87	3.20
1jhb	GLUTAREDOXIN	80	3.67	40	2.01	86	3.40
3grx	GLUTAREDOXIN 3	74	2.21	50	1.73	73	2.20

Table 2. Query protein: 3trx (Thioredoxin-Reduced Form).

Protein	otein		IEW	LOCK		DALI	
PDB ID	PDB Header (partial)	#	RMSD	#	RMSD	#	RMSD
1h75-A	GLUTAREDOXIN-LIKE PROTEIN	72	2.63	48	1.80	70	2.50
1ego	GLUTAREDOXIN (OXIDIZED)	73	2.66	42	1.31	72	2.80
1ilo	HYPOTHETICAL PROTEIN MTH895	71	3.04	43	1.51	71	3.10
1aba	GLUTAREDOXIN MUTANT	72	2.65	51	1.80	67	2.90
1fo5-A	THIOREDOXIN	77	3.01	53	1.56	74	3.40
1mek	PROTEIN DISULFIDE ISOMERASE	102	2.48	82	1.44	101	2.50
1a8y	CALSEQUESTRIN	100	2.96	80	1.58	95	2.20
1e2y-A	TRYPAREDOXIN PEROXIDASE	96	2.10	82	1.53	96	2.30
1e2y-C	TRYPAREDOXIN PEROXIDASE	96	2.10	82	1.54	96	2.30
1qmv-A	HUMAN THIOREDOXIN PEROXID.	93	2.78	79	1.60	97	2.40
1bjx	PROTEIN DISULFIDE ISOMERASE	95	2.19	80	1.53	93	2.20
1gp1-B	GLUTATHIONE PEROXIDASE	92	2.82	81	1.57	95	2.40
1qq2-A	THIOREDOXIN PEROXIDASE 2	93	2.86	73	1.59	94	2.60
1ezk-A	TRYPAREDOXIN I	96	2.77	69	1.60	90	2.50
1ewx-A	TRYPAREDOXIN I	95	2.73	70	1.59	87	2.70
1qk8-A	TRYPAREDOXIN-I	95	2.82	70	1.58	89	2.80
1fg4-A	TRYPAREDOXIN II	96	2.89	67	1.55	87	2.50
1a8l	PROTEIN DISULFIDE OXIDORED.	91	2.11	72	1.36	92	2.30
1fg4-B	TRYPAREDOXIN II	91	2.60	67	1.51	87	2.60
1fvk-A	DISULFIDE BOND FORMATION	80	2.79	58	1.54	78	2.90
1f37A	FERREDOXIN [2FE-2S]	73	2.56	46	1.59	74	3.00
1f37B	FERREDOXIN [2FE-2S]	73	2.60	50	1.57	73	3.10
1ghh-A	DNA-DAMAGE-INDUCIBLE PROTEIN	45	3.38	30	1.44	50	4.00

Table 2. (Continued)



Fig. 5. Comparison between the proposed method (NEW), LOCK, and DALI using query protein 3trx (Data used is from Table 2.) The proteins on the *x*-axis are ordered by non-increasing number of matches in the NEW method.



Fig. 6. Left side: Query protein 1mbc aligned with 5mbn, 2fal, and 1lia-A. Right side: Query protein 3trx aligned with 1erv, 1quw, and 1a8l.

In most cases, the number of matches found by our method is close to that found by DALI and fairly larger than that found by LOCK. Correspondingly, the RMSD value of our method is quite a bit smaller than that of DALI and larger than that of LOCK. When the proteins in question are distantly-related or unrelated, then the number of matches is much smaller (as is to be expected) and none of these methods perform consistently better than the others. Overall, our method appears to be competitive with LOCK and DALI.

	Prote	ein Pair		Ν	EW	L	OCK	D	ALI
PDB ID	Length	PDB ID	Length	#	RMSD	#	RMSD	#	RMSD
1mdc	133	1ifc	131	128	1.84	33	1.91		
1npx	447	3grs	461	384	2.65	32	1.62	395	3.50
1onc	104	7rsa	124	98	1.90	88	1.22	97	1.90
1osa	148	4cpv	108	68	1.69	11	2.01	67	1.40
1pfc	111	3hla-B	99	90	2.61	49	1.35	79	2.90
2cmd	312	6ldh	329	287	2.47	46	2.30	286	2.50
2pna	104	1sha-A	103	91	2.35	52	1.61	91	2.50
1bbh-A	131	2ccy-A	127	124	1.88	101	1.44	125	2.00
1c2r-A	116	1ycc	108	98	1.77	80	1.24	96	1.60
1chr-A	370	2mnr	357	347	1.83	302	1.48	347	1.90
1dxt-B	147	1hbg	147	136	1.70	119	1.36	135	2.00
2fbj-L	213	8fab-B	214	194	2.10	152	1.43	194	2.30
1gky	186	3adk	194	151	2.63	85	1.74	154	3.00
1hip	85	2hip-A	71	69	2.05	15	1.72	67	1.80
2sas	185	2scp-A	174	160	3.02	95	1.45	167	3.50
1fc1-A	206	2fb4-H	229	128	2.72	95	1.30	127	3.00
2hpd-A	457	2 cpp	405	361	2.88	217	1.75	367	3.30
1aba	87	1ego	85	73	2.08	64	1.52	72	2.20
1eaf	243	4cla	213	177	2.63	132	1.55	176	2.90
2sga	181	$5 \mathrm{ptp}$	222	149	2.44	92	1.32	145	2.70
2hhm-A	278	1fbp-A	316	225	2.56	161	1.73	219	2.90
1aaj	105	1paz	120	83	2.04	72	1.19	80	1.70
5fd1	106	1iqz	81	44	2.87	46	1.48	57	2.60
1isu-A	62	2hip-A	71	58	2.37	_	_	58	2.30
1gal	581	3cox	500	400	2.65	260	1.53	402	3.10
1cau-B	184	1cau-A	181	162	2.01	120	1.35	161	2.20
1hom	68	11 fb	77	56	1.80	48	1.11	56	1.90
1tlk	103	2rhe	114	91	1.89	60	1.17	89	2.00
20mf	340	2 por	301	264	2.54	207	1.60	261	2.80
1lga-A	343	2 cyp	293	259	2.20	210	1.54	262	2.50
1 mio-C	525	2min-B	522	396	2.86	260	1.85	412	3.60
4sbv-A	199	2tbv-A	287	163	1.98	49	2.10	162	2.10
8i1b	146	4fgf	124	115	2.42	88	1.26	118	2.50
1hrh-A	125	$1 \mathrm{rnh}$	148	114	1.92	99	1.23	110	1.90
1mup	157	1 rbp	174	141	2.69	88	1.55	140	2.90
1cpc-L	172	1col-A	197	118	3.35	75	1.65	116	3.40
2ak3-A	226	1gky	186	150	2.71	10	1.51	150	3.10
1atn-A	373	$1 \mathrm{atr}$	383	289	2.61	193	1.58	291	3.00
1arb	263	$5 \mathrm{ptp}$	222	191	2.68	128	1.43	187	2.80
2pia	321	$1 {\rm fnb}$	296	215	2.31	173	1.52	216	2.50
3rub-L	441	6xia	387	159	3.68	81	1.82	206	4.10
2sar-A	96	$9\mathrm{rnt}$	104	75	3.01	41	1.66	72	3.20
3cd4	178	2rhe	114	95	2.00	65	1.07	94	2.60
1aep	153	256b-A	106	55	2.28	49	1.74	74	1.80
2mnr	357	4enl	436	285	3.13	189	1.96	284	3.40
1lts-D	103	1bov-A	69	69	2.16	51	1.36	68	2.10
2gbp	309	2liv	344	216	3.46	88	1.91	260	6.70
1bbt	186	2 plv	288	170	2.58	96	1.56	168	2.60

Table 3. Comparison of our algorithm with LOCK and DALI on the Fischer benchmark. A "—" indicates that the method (LOCK or DALI) did not return an answer.

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Protein pair			1	NEW		OCK	DALI		
PDB ID	Length	PDB ID	Length	#	RMSD	#	RMSD	#	RMSD
2mta-C	147	1ycc	108	83	2.11	53	1.30	81	2.20
1tah-A	318	1tca	317	191	2.37	119	1.57	187	2.40
1rcb	129	2gmf-A	121	98	3.52	63	1.44	82	3.30
1sac-A	204	2ayh	214	97	4.48	94	1.86	134	3.30
1dsb-A	188	2trx-A	109	85	2.56	19	2.14	83	2.90
1stf-I	98	1mol-A	94	85	1.90	65	1.26	85	1.90
2afn-A	331	1aoz-A	552	128	2.41	161	1.77	248	2.60
1fxi-A	96	1ubq	76	63	2.62	40	1.48	61	2.70
1bge-B	159	2gmf-A	121	101	3.09	59	1.90	94	3.30
3hla-B	99	2rhe	114	82	3.13	32	1.64	71	3.30
3chy	128	2fox	138	107	2.90	57	1.96	103	3.00
2aza-A	129	1paz	120	83	2.36	63	1.65	81	2.50
1cew	108	1mol-A	94	81	2.12	46	1.72	81	2.30
1cid	177	2rhe	114	98	2.61	35	1.79	97	3.20
1crl	534	1ede	310	208	2.99	133	1.87	211	3.50
2 sim	381	1nsb-A	390	292	2.97	154	1.78	291	3.30
1ten	89	3hhr-B	195	87	1.82	29	1.87	86	2.00
1tie	166	4fgf	124	90	3.85	86	1.51	114	3.10
$2 \mathrm{snv}$	151	5 ptp	222	132	2.87	82	1.79	130	3.10
1gp1-A	183	2trx-A	109	97	2.40	25	2.08	97	2.60

Table 3. (Continued)



Fig. 7. Comparison between the proposed method (NEW), LOCK, and DALI using the Fischer benchmark. (Data used is from Table 3.) The proteins on the *x*-axis are ordered by non-increasing number of matches in the NEW method.

Our second experiment was to compare our method with LOCK and DALI using the well-known Fischer benchmark,⁸ which contains 68 pairs of proteins. The proteins in each pair are known to be structurally similar, but have low sequence identity, ranging from 8% to 31% with an average of 18.6% and a standard deviation



Fig. 8. Left side: Protein 1onc aligned with 7rsa. Right side: Protein 2fbj-L with 8fab-B.

of 4.4. The results of the alignment (i.e. the number of matched atoms and the RMSD value) are summarized in Table 3 and shown graphically in Fig. 7, where the protein pairs are listed on the *x*-axis in non-increasing order of the matches found by our method. The alignment of protein 1 onc with 7 rsa is shown in Fig. 8 (left side), and the alignment of protein 2 fbj-L with 8 fab-B is shown in Fig. 8 (right side). We observe the same trend as before with this dataset also, i.e., the number of matches found by our method is close to that found by DALI and much larger than that found by LOCK, while the RMSD value of our method is quite a bit smaller than that of DALI and larger than that of LOCK.

5. Conclusions

We have presented an iterative refinement algorithm for pairwise protein structure alignment. The algorithm uses an angle-based representation of the protein backbones, which is independent of the relative orientation of the proteins in space. An initial alignment is found using dynamic programming (on the backbone representation) and a graph-based method and then refined iteratively, such that the number of matched C_{α} atoms is large and the distance between matched atoms is within a prescribed threshold. The heuristic has been implemented and found to be competitive with two other algorithms (LOCK and DALI), especially for closely-related proteins.

The algorithm proposed in this paper uses protein backbone geometry alone for comparison. However, other information, such as chemical properties of residues and secondary structure information could be used to improve the quality of the alignment further, by using a suitable scoring function in Eq. (1). We plan to study the effect of including this additional information in our algorithm.

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A shorter version of this paper, which omits some experimental results, the proof of Lemma 2, and some figures, appears in the Proceedings of the 15th IEEE International Conference on Tools with AI, Sacramento, CA, Nov. 2003.

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