

The Fragment Transformation Method to Detect the Protein Structural Motifs

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ABSTRACT To identify functional structural motifs from protein structures of unknown function becomes increasingly important in recent years due to the progress of the structural genomics initiatives. Although certain structural patterns such as the Asp-His-Ser catalytic triad are easy to detect because of their conserved residues and stringently constrained geometry, it is usually more challenging to detect a general structural motifs like, for example, the $\beta\beta\alpha$ -metal binding motif, which has a much more variable conformation and sequence. At present, the identification of these motifs usually relies on manual procedures based on different structure and sequence analysis tools. In this study, we develop a structural alignment algorithm combining both structural and sequence information to identify the local structure motifs. We applied our method to the following examples: the $\beta\beta\alpha$ -metal binding motif and the treble clef motif. The $\beta\beta\alpha$ -metal binding motif plays an important role in nonspecific DNA interactions and cleavage in host defense and apoptosis. The treble clef motif is a zinc-binding motif adaptable to diverse functions such as the binding of nucleic acid and hydrolysis of phosphodiester bonds. Our results are encouraging, indicating that we can effectively identify these structural motifs in an automatic fashion. Our method may provide a useful means for automatic functional annotation through detecting structural motifs associated with particular functions. *Proteins* 2006;63:636–643. © 2006 Wiley-Liss, Inc.

Key words: local structure alignment; DNA-binding proteins; structural motif; structural genomics

INTRODUCTION

Due to the progress of structural genomics research,¹ a significant number of protein structures of unknown function are solved in recent years—more than 700 structures annotated as “Structural Genomics Unknown Protein” are currently deposited in Protein Data Bank (PDB).² Hence, the need of function prediction tools from protein structures becomes increasingly important in the present structural genomic era. Sequence alignment and global fold comparison are valuable tools in inferring protein function through comparing sequence or fold similarity; however, these methods may not be applicable in certain situations:

because nonhomologous proteins with different folds may evolve convergently similar spatial dispositions of functional residues for similar functions, sequence or fold alignment may fail to identify the particular functional sites. It may also happen that nonhomologous proteins evolve similar overall conformations as a scaffold for different functions. The apparent fold similarity may lead to erroneous prediction of function. Hence, the ability to identify local structural motifs is important in predicting protein function and in providing a complementary tool to the fold alignment method.

The structural motif may be roughly divided into two types. The first type is characterized by a conserved spatial arrangement of a number of conserved residues that are functionally important. The distances among these residues are usually quite conserved, showing little variations. Hence, the sequence and distance constraints provide stringent criteria to identify this type of motif. A typical example is given by the catalytic triad (Asp-His-Ser) of serine protease. Study³ showed that the majority of the catalytic triads are within a root-mean-square (RMS) distance of 2.0 Å from the consensus template. Using this distance, that is, 2.0 Å, as a cutoff can effectively separate the catalytic triad from other noncatalytic Ser-His-Asp associations. Another example is given by the active site of enolase superfamily, which can be accurately characterized by the spatial arrangement of five residues.⁴ A number of methods^{5–16} were developed to identify this type of structural motif, taking advantage of the distance constraints of the conserved residues.

The second type of structural motifs is composed of a number of secondary structure elements (SSEs) with each SSE arranged in a particular orientation. These SSEs are usually separated by variable lengths of intervening sequences in different protein families. The motif sequences are usually much less conserved than those of the first type, and the RMS distances of the motif residues from the

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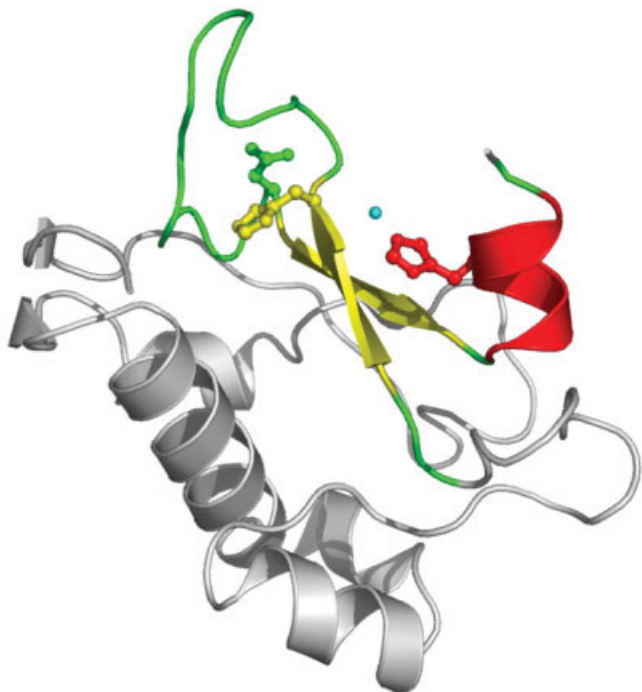


Fig. 1. The ribbon representation of colicin ColE7 (PDB code: 7CEI). The $\beta\alpha$ -metal motif is rendered in colors according to the secondary structural elements (helices in red, β -strands in yellow, and loops in green), while other parts of the protein in gray. The Zn metal ion is represented by a cyan sphere. The $\beta\alpha$ -metal motif of the colicin family (i. e., ColE2, ColE7, ColE8, and ColE9) contains the so-called conserved H-N-H motif, which is shown in a ball-and-stick model. However, only the first His residue of the sequence motif (i. e., on the first β -strand) is strictly conserved in the general $\beta\alpha$ -metal motif of other proteins. All molecular images in this work are created by PyMOL (<http://www.pymol.org>).

consensus template are more widely distributed. Examples are the $\beta\alpha$ -metal binding motif,^{17–23} the treble clef finger motif,²⁴ and the helix-turn-helix motif.²⁵ Figure 1 shows a typical $\beta\alpha$ -metal binding motif. The motif sequences are highly variable—there is only one strictly conserved His residue, which serves as a general base to activate a water nucleophile.²³ There do exist a special case of the $\beta\alpha$ -metal motif that can be identified by its particular sequence motif, the so-called the H-N-H motif,^{26,27} characterized by the consensus sequence of EXHHX_{1,4}NX₈HX₃H;²⁸ however, this instance does not reflect the general cases. At present, the identification of the $\beta\alpha$ -metal motifs mainly relies on manual inspection assisted with a number of structure and sequence analysis tools.²² Here, we present a structural alignment algorithm combining both structural and sequence information to identify the local structure motifs. As a practical application, we applied our method to detecting the $\beta\alpha$ -metal motif and the treble clef finger motif.

METHODS

In this section, we will first define some basic notations of our method. The symbol S denotes the query structural motif of length n , which may comprise several discontinuous peptide fragments. The C_α coordinates of S are given

by $(\mathbf{x}_1, \mathbf{x}_2, \dots, \mathbf{x}_n)$. The symbol T is used to denote the target protein of length m with the C_α coordinates given by $(\mathbf{y}_1, \mathbf{y}_2, \dots, \mathbf{y}_m)$. The basic unit of the structural alignment is a triplet composed of three consecutive C_α atoms. The structures S and T can be expressed in terms of these triplets, that is, $S = \{\sigma_1, \sigma_2, \dots, \sigma_{n-2}\}$ and $T = \{\tau_1, \tau_2, \dots, \tau_{m-2}\}$, where

$$\sigma_i = (\mathbf{x}_i, \mathbf{x}_{i+1}, \mathbf{x}_{i+2}) \text{ and } \tau_i = (\mathbf{y}_i, \mathbf{y}_{i+1}, \mathbf{y}_{i+2}).$$

We can construct an $(m-2) \times (n-2)$ matrix between the σ and τ triplets,

$$\mathbf{M} = \begin{bmatrix} M_{11} & M_{12} & \dots & M_{1,n-2} \\ M_{21} & M_{22} & \dots & M_{2,n-2} \\ \dots & \dots & \dots & \dots \\ M_{m-2,1} & M_{m-2,2} & \dots & M_{m-2,n-2} \end{bmatrix}, \quad (1)$$

where the element $M_{i,j}$ is a rigid-body transformation matrix from σ_i to τ_j , that is, $M_{i,j}\sigma_i = \tau_j$ (see Fig. 2 for schematic illustration).

Triplet Clustering

We use the symbol D_{kl}^{ij} to denote the Cartesian distance between the transformed triplet $M_{i,j}\sigma_k$ and the target triplet τ_l . The distance D_{kl}^{ij} is used as a measure of orientation similarity between the triplet pairs (σ_i, τ_j) and (σ_k, τ_l) . We cluster the triplet fragments using the single-linkage algorithm.²⁹ The procedures go as follows: for two triplets (σ_i, τ_j) and (σ_k, τ_l) under the criteria: $D_{kl}^{ij} < D_0$, and $i \neq k$ and $j \neq l$, the triplets will be clustered in the same group. Let G_1 and G_2 be two cluster groups, the former containing (σ_i, τ_j) and (σ_k, τ_l) , the latter containing $(\sigma_{i'}, \tau_{j'})$. If $D_{k'l'}^{ij} < D_0$, then G_1 and G_2 will be merged to form a new cluster $G_3 = G_1 \cup G_2$. The procedures will be carried out iteratively until no more new cluster groups are formed. For the final cluster G_μ , we can get the aligned substructure pair:

$$S_\mu = \bigcup_{\sigma_k \in G_\mu} \sigma_k$$

and

$$T_\mu = \bigcup_{\tau_k \in G_\mu} \tau_k.$$

Note that the substructure is not necessarily continuous in space. We define for each cluster G_μ the alignment ratio ϵ_μ as f/N , where f is the number of triplets of S_μ , that is, the aligned atoms, and N is the number of the triplets of the query structure.

The Scoring Function

To assess the alignment quality of a given cluster G_μ , we define the following scoring function

$$C_\mu = w_1 \text{RANK}_a(C_\mu^R) + w_2 \text{RANK}_d(C_\mu^B) + w_3 \text{RANK}_d(C_\mu^S) \quad (2)$$

where $\text{RANK}_a(x)$ is a ranking function of x in the ascending order, $\text{RANK}_d(x)$ is a ranking function in the descending order, and w_i is the weight of the associated rank score. The weights w_i of Equation (2) are positive and satisfies

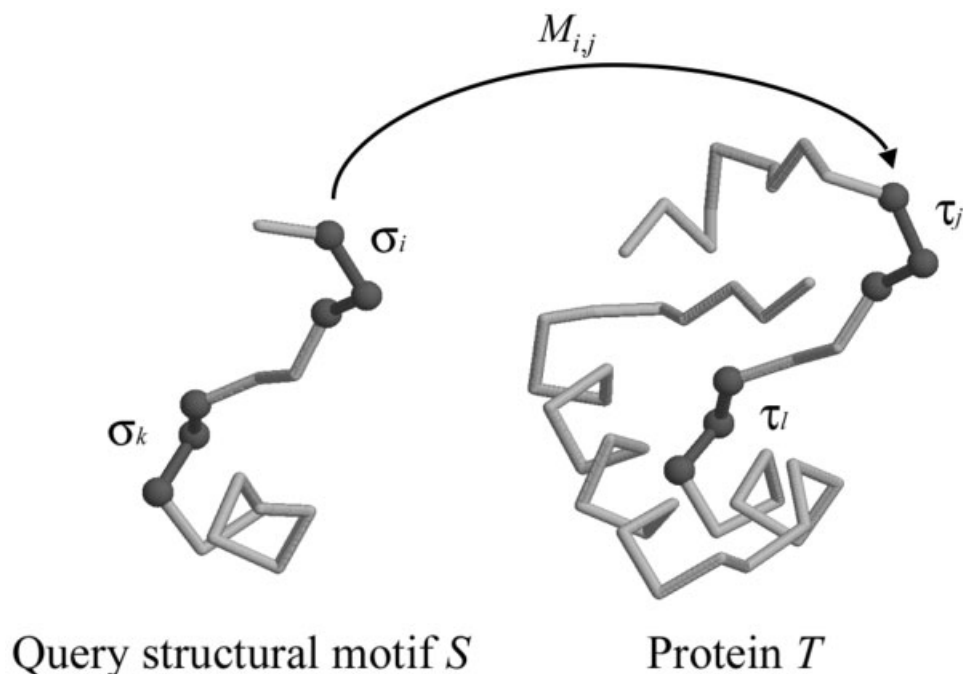


Fig. 2. σ_i and σ_k are two arbitrary triplet units of the query structural motif *S*, and τ_i and τ_j are two arbitrary triplet units of the target protein *T*. The triplet σ_i is transformed to τ_j through the transformation matrix $M_{i,j}$.

TABLE I. Substitution Matrix of Secondary Structural Elements

	H	G	I	E	B	T	S	U
H	5	3	3	-5	-5	0	0	0
G	3	5	3	-5	-5	0	0	0
I	3	3	5	-5	-5	0	0	0
E	-5	-5	-5	5	3	0	0	0
B	-5	-5	-5	3	5	0	0	0
T	0	0	0	0	0	3	0	0
S	0	0	0	0	0	0	3	0
U	0	0	0	0	0	0	0	1

the condition that $\sum_i^3 w_i = 1$. The $\beta\beta\alpha$ -metal motifs identified by Li et al.²² are used as the positive examples for the manual adjustment of the parameters: the weight w_i , and the clustering parameters D_0 and ϵ_μ . The final values of these parameters are: $(w_1, w_2, w_3) = (0.4, 0.2, 0.4)$, $D_0 = 3 \text{ \AA}$ and $\epsilon_\mu > 0.75$. The same parameters are used throughout all calculations in this work. The alignment scores C_μ^R , C_μ^B , and C_μ^S are defined as follows,

$$C_\mu^R = \frac{1}{\epsilon_\mu} \text{RMSD}(S, T_\mu) \quad (3)$$

where $\text{RMSD}(S, T_\mu)$ is the root-mean-square deviation of C_α atoms between *S* and T_μ ;

$$C_\mu^B = \frac{1}{\epsilon_\mu} (\text{BLOSUM}(S, T_\mu) + d_0^B) \quad (4)$$

where $\text{BLOSUM}(S, T_\mu)$ is the sequence alignment score based on BLOSUM62³⁰ between *S* and T_μ and d_0^B is a constant to make C_μ^B negative;

TABLE II. The Top Ranking Hits of the $\beta\beta\alpha$ -Metal Motif

Proteins	PDB ID	Structural aligned sequences ^b
Colicin E7 ^a	7CEI:B	SFELHHE . (13) .NISVV-TPKR <u>H</u> IDI
Colicin E9	1EMV:B	VYELHHD . (13) .NISVV-TPKR <u>H</u> IDI
Vnn	1OUO:A	RLEWEHV . (36) .NLTPA-IGE <u>V</u> NGDR
I-HmuI	1U3E:M	GLVVDHK . (10) .NLRWV-TQK <u>I</u> NVEN
I-PpoI	1EVW:A	TCTASHL . (10) .HLCWE-SADD <u>N</u> KGR
Serratia	1Q10:A	KVDRGHQ . (9) .NITPQ-KSD <u>L</u> NGAW
EndoVII	1E71:A	ANHLDDH . (10) .VRGLL-CNLC <u>D</u> AAE
CAD/DFP40	1V0D:A	TWNLDHI . (34) .NLKLV-HIACH <u>H</u> KKT
HIV integrase	1K6Y:A	QLDCTHL . (2) .KVILV*IESM <u>N</u> KEL

Only the representative proteins of each protein family are listed.

^aThe $\beta\beta\alpha$ -metal motif of this protein is used as the structural template.

^bIn the alignment, The symbol “.(n).” indicates a sequence of length, *n*, “-” a gap and “*” a sequence of unspecified length. The letter in bold face indicates the strictly conserved residue the His residue, while the letter underlined the less conserved His, Asn or Glu residue.

$$C_\mu^S = \frac{1}{\epsilon_\mu} (SS(S, T_\mu) + d_0^S) \quad (5)$$

where $SS(S, T_\mu)$ is the secondary structure alignment score between *S* and T_μ and d_0^S is a constant to make C_μ^S negative. The secondary structure assignment is based on the DSSP method,³¹ which assigns secondary structures according to their hydrogen bonding patterns. The DSSP method defines eight secondary structures: α -helix (H), 3_{10} -helix (G), π -helix (I), extended β -strand (E), isolated β -strand (B), turn (T), bend (S), and coil (U). The substitution matrix is given in Table I.

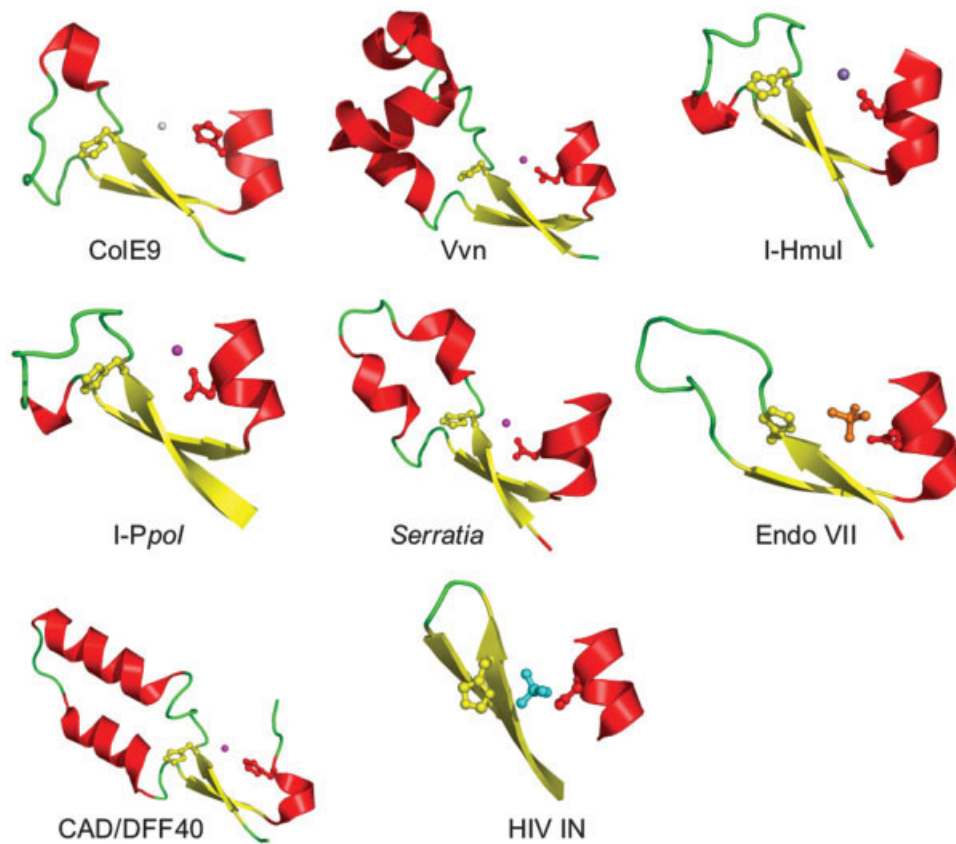


Fig. 3. The identified $\beta\alpha$ -metal motifs (using ColE9 (1EMV:B) as the reference) in Vvn (1OOU:A), I-Hmul (1U3E:M), I-Ppol (1EVW), *Serratia* nuclease (1QL0), the T4 Endo VII (1E7L:A), CAD/DFF40 (1V0D), and HIV IN (1EX4:A). The conserved His residue, the ligand and the binding residue (His, Asn or Asp), which is the underlined residue in Table II, are shown in the ball-and-stick model.

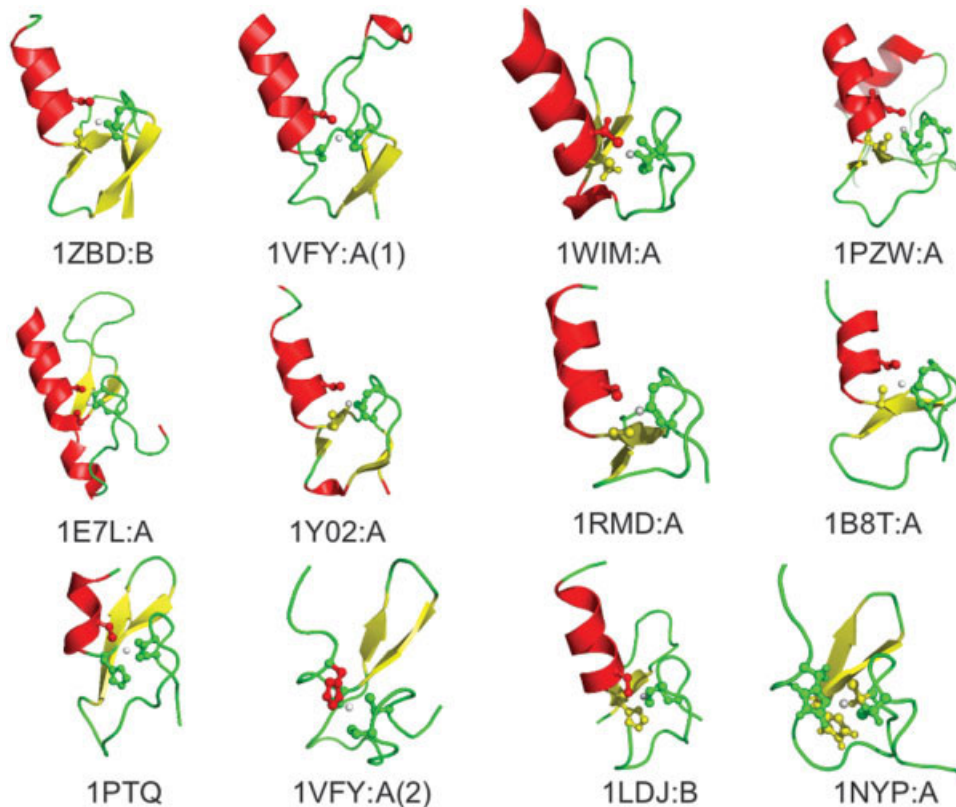


Fig. 4. The identified treble clef finger motifs [using G protein Rab3A (1ZBD:B) as the reference] in Vps27p (1Vfy:A(1)), UbcM4-interacting protein 4 (1WIM:A), transcription factor Grauzone (1PZW:B), the T4 Endo VII (1E7L:A), CARP2 (1Y02:A), Hrs (1DVP:A), RAG1 dimerization domain (1RMD), CRP1 (1B8T:A), Protein kinase C (1PTQ:A), Vps27p (1Vfy:A(2)), ring-box protein 1 (1LDJ:B), and LIM4 (1NYP:A).

TABLE III. Top Ranking Hits of the Treble Clef Finger Motif

Proteins	PDB ID	Structural aligned sequences ^b
G protein Rab3A ^a	1ZBD:B	SVVCEDCCKNVC*GVE*WLCKICLEQREVWK
Vps27p	1VFX:A	KHHCRSCGGVFC*SSN*RVCDSCFEDYEFIV
UbcM4-interacting protein 4	1WIM:A	SSGCKLCLGEYP*MTT*IFCTLCLKQYVELL
Transcription factor Grauzone	1PZW:B	-ICRLCLRGVS*CLQ*VICNVQWTVQSEFH
Endo VII	1E7L:A	-GKCLICQRELN*NHL*LLCNLCDAEAGQMK
CARP2	1Y02:A	KQTCLDCKKNFC*SS*RLCLLQCRFRATFQ
Hrs	1DVP:A	KHHCRNCGQVFC*TAK*RVCDGCFALQR-
RAG1 dimerization domain	1RMD	-ISQICEHILA*PVE*LFRCICILRCLKVM
CRP1	1B8T:A	-FLCMVCKKNLD*VAV*TYCMVYCGKKG-P
Protein kinase C	1PTQ:A	PTFCDHCGSLQ*GLK*NVHHKCREKV-A-
Vps27p	1VFX:A	SDACMICSKKFS*KHH*VFCQEHSS-NS-
Ring-box protein 1	1LDJ:B	-NCAICRNM-D*VAW*AFHFHCISRWLKTR
LIM4 domain of PINCH protein	1NYP:A	VPICGACRRPIE*VVN*QWHVEHFCAKCE-

Only the representative proteins are listed. In addition, we list the proteins (the bottom 4), although not among the highest ranks, that contain motifs other than the C2C2 sequence motif of the query proteins, that is, C2HC, C2CH, and C2H2. The residue in bold indicates the conserved residues of the motif.

^aThe treble clef finger motif of this protein is used as the structural template.

^bIn the alignment, The symbol “.(n)” indicates a sequence of length *n*, “-” a gap and “*” a sequence of unspecified length.

RESULTS AND DISCUSSION

The $\beta\beta\alpha$ -Metal Motif

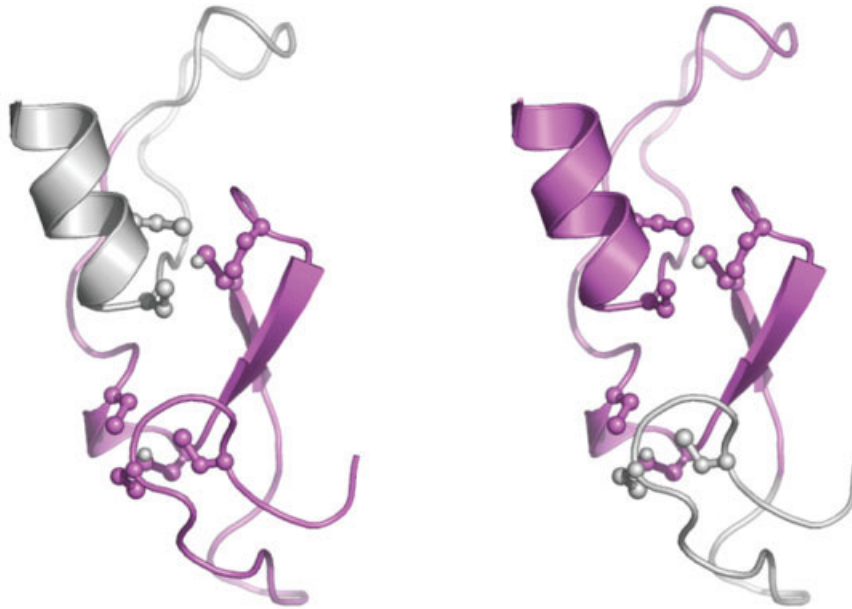
The $\beta\beta\alpha$ -metal motif has recently received much attention due to its important role in nonspecific DNA interactions and cleavage in host defense and apoptosis.^{17–22,27,32,23} The $\beta\beta\alpha$ -metal binding motif is characterized by one α -helix and an antiparallel β -sheet, which are separated from each other by a variable length of intervening sequences from 9 to 36. This motif usually binds to divalent metal ions like Zn or Mg. The lengths and sequences of the $\beta\beta\alpha$ -metal motif vary greatly in different proteins. For example, there is only one strictly conserved His residue located at the first β -strand in the motif sequence. We used the $\beta\beta\alpha$ -metal motif of the nuclease domain of colicin E7 (PDB ID: 7CEI)³³ as the query motif, which is composed of three fragments: E542–H545 (β -strand), I561–V564 (β -strand) and P566–D571 (α -helix). The query fragments are compared against the nonredundant PDB (nrpdb) chain set composed of nonidentical proteins from NCBI (<http://www.ncbi.nlm.nih.gov/Structure/VAST/nrpdb.html>).

Table II lists the top ranking hits of proteins and the corresponding structure-aligned sequences. We list only the representative proteins of the protein families, which are colicin E9 (ColE9),³⁴ the periplasmic nuclease Vvn,²² the HNH homing endonuclease I-HmuI,³⁵ the His–Cys box endonuclease I-*PpoI*,¹⁸ the endonucleases such as *Serratia* nuclease,¹⁷ CAD/DFP40,³² the T4 Endo VII,²⁰ and HIV integrase (IN).³⁶ The lengths of the intervening sequence between the two β -strands vary from 2 to 39. The structures of the $\beta\beta\alpha$ -metal binding motifs of these proteins are shown in Figure 3. We have identified all proteins that have experimentally confirmed $\beta\beta\alpha$ -metal binding motifs. The only exception is HIV IN, which has not been reported to have a structural motif similar to the $\beta\beta\alpha$ -metal binding motif. The identified structural motif is located in the catalytic core domain of HIV IN.³⁶ Two of the three catalytic residues³⁶ of HIV IN, that is, D64 and E152, are

located in this motif, while the third one (D116) is in the vicinity of the motif. Although we did not find a divalent metal ion in the center of the motif in the crystal structures of HIV IN, we did find a phosphate group (PO_4^-) in 1K6Y, which is close to the supposed metal-binding site. However, conspicuous differences do exist: in HIV IN, the β -strands of the structural motif are separated by only two residues, while the usual intervening length is much longer in the range between 9–36 (Table II and Fig. 3); furthermore, the second β -strand does not directly connect to the α -helix like others do. Obviously, the question of whether HIV IN did contain an authentic $\beta\beta\alpha$ -metal motif needs further study.

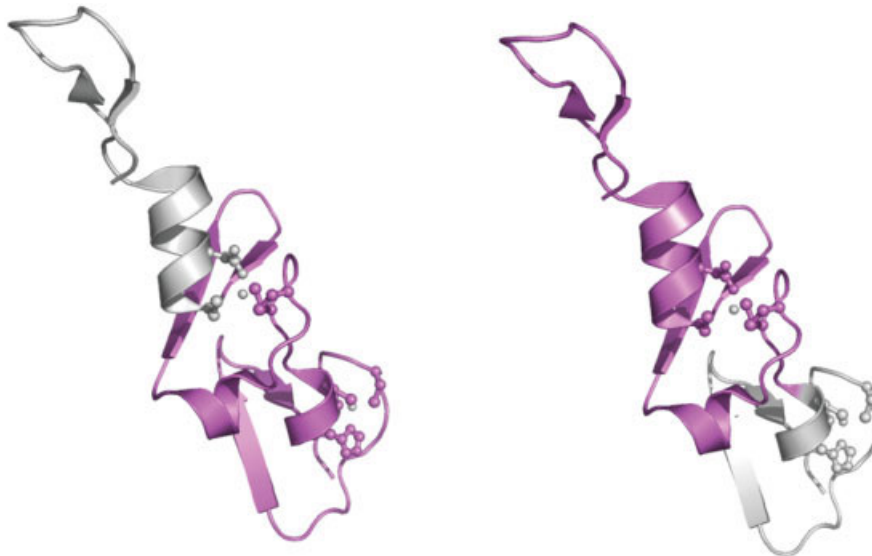
The Treble Clef Finger Motifs

The treble clef finger motif²⁴ is a zinc-binding structural motif composed of a zinc knuckle, a loop, a β -hairpin and an α -helix. It is so named because of its projection of the domain C_α trace resemble the treble clef sign.²⁴ The treble clef finger structures show highly variable sequences and appear in many folds. For example, in terms of the SCOP protein folds, the treble clef finger motif occurs in a number of folds such as the RING/U-box, glucocorticoid receptor-like folds, FYVE/PHD zinc fingers, His-Me finger endonucleases, C2H2 and C2HC zinc fingers, and cysteine-rich domains. It has been noted³⁷ that fold similarity search programs such as DALI,³⁸ and CE,³⁹ and VAST⁴⁰ are not consistent in detecting the treble clef finger motif due to its relatively smaller size. Previously, a combination of fold alignment methods, together with the PSI-BLAST and the manual procedures, was used to identify the treble clef finger motifs.²⁴ We used the zinc finger motif of the G protein Rab3A (PDB ID: 1ZDB:B)³³ as the query motif, which is composed of three fragments: S108–C119, G123–E125, and W135–K148. The top ranking hits are shown in Table III, and the corresponding structures of the zinc fingers are shown in Figure 4. Both sequence and struc-



161 196
GRVCHRCRVEFTFTNRKHHCRNCGQVFCGQCTAKQCPLPKYGIEKEVRVCDGCFAALQR
 176 219
 GRVCHRCRVEFTFTNRKHHCRNCGQVFCGQCTAKQCPLPKYGIEKEVRVCDGCFAALQR

Fig. 5. The ribbon drawing of the two overlapping zinc fingers in the FYVE domain of Hrs (1DVP:A). The finger motifs are colored in violet. Shown in the bottom are the sequences of the first zinc finger (G161–C196, which are underlined in the upper sequence) and second zinc finger (K176–R219, which are underlined in the lower sequence). The zinc ligands are shown in bold face.



20 57
WKRCAGCGGKIADRFLLYAMDSYWHSRCLKCSSCQAQLGDIGTSSYTKSGMILCRNDYIRLFGNSGACSACGQSIPA
 48 96
 WKRCAGCGGKIADRFLLYAMDSYWHSRCLKCSSCQAQLGDIGTSSYTKSGMILCRNDYIRLFGNSGACSACGQSIPA

Fig. 6. The ribbon drawing of the consecutive zinc fingers in the tandem LIM domains of LMO4 (1RUT:X). The finger motifs are colored in violet. Shown in the bottom are the motif sequences of the first zinc finger (the sequence underlined in the upper row—W20–L57) and the second zinc finger (the sequence underlined in the upper row—L48–A96). The zinc ligands are shown in bold face. Note that LMO4 has an unusual CCCD Zn-binding module.

tures of the treble clef zinc fingers show significant variations; for example, the motif sequences show little similarity except the sequence motif like C2C2, C2HC, C2CH, C2H2, or C2CD (see next section) involved in zinc binding.

The Double Treble Clef Finger Motif

The treble clef finger motif in many cases occurs as an overlapping treble-clef structure,²⁴ where the two zinc binding fingers, each finger showing a typical treble clef motif, overlap each other over a significant fraction of the whole motif sequence. In the overlapping zinc binding module, the first and the third Cys pairs bind to one zinc ion, while the second and the fourth Cys pairs bind to the other zinc ion. The overlapping treble-clef structure is typical of the FYVE domains.⁴¹ Our method successfully identified this structural motif in the FYVE domain of the proteins such as endosomal autoantigen 1 (1JOC:B), the Vps27p protein (1Vfy:A), Hrs (1DVP:A), and the protein 19 from *Mus musculus* (1WFK). As an example, Figure 5 shows the detected overlapping treble-clef structure in the FYVE domain of Hrs.⁴² Note that, although the two zinc fingers overlap over more than two-thirds of the motif sequence, our method is still able to identify these two overlapping fingers.

The treble-clef fingers also occurs as the so-called double treble-clef structure²⁴ where two fingers are arranged in a consecutive, instead of overlapping, fashion. In contrast to the previous overlapping treble-clef structure, the first two pair of the ligands bind to one zinc ion, while the last two pairs bind to a second zinc ion. The double treble-clef structure is typical of LIM domains.⁴³ Our method detects an interesting case of such motif in the tandem LIM domains of the LMO4 protein shown in Figure 6. Note that LMO4 has an unusual CCCD Zn-binding sequence motif.

Comparison with Other Approaches

Currently, there are three popular methods for structure alignment available on the Internet: DALI, CE, and VAST. However, because DALI and CE are mainly designed for the structural alignment of global folds, they failed to return results for both the $\beta\beta\alpha$ -metal and the zinc finger motifs, which are composed of relatively short and discontinuous fragments. On the other hand, the VAST algorithm, using a hierarchical alignment based on the SSEs, is able to identify local structural motifs. In the case of $\beta\beta\alpha$ -metal motifs, the VAST tool, using the same query fragment defined in the previous section against the medium-redundancy subset of PDB from NCBI, correctly identifies 4 out of the top 10 ranks (40%) based on the VAST score, while our approach yields 7 of 10 (70%). In the case of the zinc finger motifs, VAST correctly identify 5 out of the top 10 ranks (50%), while our approach is able to yield 9 of 10 (90%). It should be noted that our program can be easily extended to identify the type 1 structural motifs such as the catalytic triads, which can be identified through stringent distance constraints. Using the data set comprising 88 serine hydrolase entries in enzyme class E.C. 3.4.21 from Chou and Cai,⁴⁴ our approach can achieve a overall success rate of 6187/6189 (99.92%) for the active

site identification, which is comparable to that of the covariant discriminant approach designed for the identification of the catalytic triad.

In summary, we have developed an approach to identify the local structural motifs, which may be useful in the automatic functional annotation for structural genomics research. The Linux binary codes of our method are available upon request.

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