Application of Sequence Alignment Methods to Multiple Structural Alignment and Superposition¹

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Abstract. With the goal of developing efficient multiple structural alignment methods, we have asked which of the pairwise structure alignment methods lends itself most readily to generalization to multiple structure alignment. A simple linear encoding of the sequence and associated residue conformation can be treated by standard multiple *sequence* alignment methods.

Key words: Protein structure, multiple alignment

1 Introduction

One often wishes to analyse proteins that have similar folding patterns but too little sequence similarity to permit the alignment of their residues by sequence-based methods. Such proteins may be very distant relatives, or independently-evolved examples of the same folding pattern. For only two structures, it is possible to perform a structural alignment; that is, to identify residues that occupy similar spatial positions within the structure [GL98]. However, just as multiple sequence alignments are far more informative than pairs of aligned sequences, so the analysis of protein structures requires alignment of more than two sequences.

Most previous approaches to multiple structure alignment have been based on pairwise structural alignments. The simplest approach is to choose a master structure and align all the others to it. This has the obvious limitations of dependence on the choice of the master structure, and failure to make use of relationships between pairs of nonmaster sequences. Lesk & Fordham [LF96], in a study of the chymotrypsin-like serine proteases, did structural alignments of all pairs of structures, and collated the results into a common alignment table. However, the experience with those calculations suggests that it would be useful to ask whether any of the known pairwise structural

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superposition methods lends itself to generalisation to a true multiple superposition approach. The problem is only to determine the residue-residue correspondences, that is, the alignment. Once the alignment is known methods are available for the multiple superposition of the molecules [SBPL92],[D92].

There have been numerous approaches to the problem of structural superposition (for a review see [GL98]). Some operate in three-dimensional space, and are based on detection of small well-fitting pieces and combining them [VS91],[ATG92]; others are based on similarity of contact matrices [HS93],[NRTZ95],[L95].

However, the methods that would seem to be most directly generalizable to multiple alignment are those that reduce the three-dimensional structural superposition problem to a one-dimensional problem. There are several ways to achieve this. One is to characterise each residue by its pattern of neighbours [LVW85],[TO89]. Another is to characterize each residue by its mainchain conformation [LSW84],[KdHN89]. (It is clear that these approaches depend on the linear nature of the polypeptide chain.) Still another is to classify each position in a polypeptide chain by its environment; this also has application to structure prediction by asking whether a particular sequence is compatible with a succession of encoded environments [BLE91].

In this report we pursue the idea that after encoding a protein by a one-dimension characterization of the successive residues, together with limited amino acid sequence information, multiple sequence alignment methods can be applied to produce a multiple structure alignment. We use a set of distantly-related globins as an example and test of feasibility of the method.

Other approaches to multiple structure alignment have been published by Russell & Barton [RB92], Taylor, Flores & Orengo [TFO94], and May & Johnson [MJ95]. Our approach is similar to that of Šali & Blundell [ŠB90].

2 Co-ordinates and Calculations

All co-ordinates are taken from the Protein Data Bank [B77]. For multiple sequence alignment we used the program map, by Huang [H94].

We assign to each residue a symbol that combines information from the amino acid sequence and from the residue conformation.

2.1 Encoding the sequence: reduced amino acid alphabet

We encode the amino acid sequence according to a reduced alphabet corresponding to physico-chemical classes of amino acids:

Table 1. Reduced alphabet based on classifying amino acids into types of similar physicochemical properties

GAST	small nonpolar
CVILP	small/medium hydrophobic
FYMW	large hydrophobic
NQH	polar
DE	charged, negative
KR	charged, positive

2.2 Encoding the conformation

We make use of Efimov's dissection of the Sasisekhan-Ramachandran diagram [E93], with modifications: The conformation of the mainchain of a protein is specified by conformational angles ϕ , ψ and ω . Values of ω are limited to narrow ranges around +180° and -180°. Allowed ranges for ψ and ϕ are limited by steric constraints to discrete regions which can be charted in the Sasisekharan-Ramachandran plot. We use the nomenclature of Efimov [E93] but extend his regions to assign to each residue a symbol for the region to which it is closest. (Efimov's definitions cover only a subset of the possible values of ϕ and ψ .) In this way we encode the structure of a protein as a sequence of conformation states of the individual residues:

Table 2. Classification of mainchain conformations based on that of A.V. Efimov [E93]

- A α_r right-handed α -helix
- B β extended strand
- D throat between α and β regions
- L α_l left-handed α -helix
- E bottom of +/- region (in which $\phi > 0, \psi < 0$)
- C cis-peptide
- X other

From the previous two tables we have assigned to each residue one of six symbols based on its amino acid identity, and one of 7 symbols based on its conformation. By assigning a unique symbol to each possible combination of these we represent each residue by a single character in a 42-character alphabet. Each element of the substitution matrix associated with this alphabet is the sum of a contribution from change in amino acid class (see Table 1) and a contribution from change in conformation class (see Table 2) according to the following rules:

Contribution from amino acid classification:

Same class uncharged \leftrightarrow uncharged uncharged \leftrightarrow charged (including polar) 10 5 0

Contribution from conformational classification:

Same class different class 0 -10

The initiate-gap penalty was 20 and the extend-gap penalty 5.

3 Results

We have implemented the methods described and applied them to three distantlyrelated globin structures: sperm whale myoglobin, bloodworm globin and leghaemoglobin from yellow lupin. The results are as follows. (The symbols, which correspond to the assignment of a unique character to each ordered pair of reduced amino acid alphabet and residue conformation, should be considered arbitrary.)

Sperm whale	HBYAYMSGGGSGMA4GYAOGAASASYGGG4GM4AUGYAGY4NY4M4SH4BYAY
Bloodworm	HBAAS4SGGAAAM4YGAECOVDAAGA44GGG4MGAAUGSMAAGMDNAEACZGA
Yellow lupin	EJBYASAAGG4AAMYYMSAUGG4SAS4MMGGGGYGCGAA4YGNAMG5EBAZHHSUTGY
-	
Sperm whale	M4ABYYG44SAGAGGAAGAAGG447EUUYAYG4GGASASAA4S7HHG4MGYMGAYAGG
Bloodworm	GAAGAA4GGASGAGAGAUGAOYA4MGASM4AGAG4S4CNES5TH5ASMMYGGAAAGG
Yellow lupin	GSA-SAA4GM4GGMYAAGSGYGAEHHHBZAAG4SGAAGSGA6-DHBYASMGGG4YAGG
-	
Sperm whale	SGGSA4UGAYOBAYASAAMS4AGYGM44YGAA4M4YGDNV
Bloodworm	AAMYS4GEA4OTAAA4YAMAAAMAYGAAAGGAAGS
Yellow lupin	4AG4YGGEA6NBYYGSAAMAGAMYYGAGGG44YMYYA
Atnonalation	of these popults heads to the amine acid sequence follows:
A translation	f of these results back to the annuo acid sequence follows:
Sperm whale	VLSEGEWQLVLHVWAKVEADVAGHGQDILIRLFKSHPETLEKFDRFKHLKTEAE
Bloodworm	GLSAAQRQVIAATWKDIAGADNGAGVGKKCLIKFLSAHPQMAAVFGFSGASDPG
Yellow lupin	GALTESQAALVKSSWEEFNANIPKHTHRFFILVLEIAPAAKDLFSFLKGTSEVPQNNPE
Sperm whale	MKASEDLKKHGVTVLTALGAILKKKGHHEAELKPLAQSHATKHKIPIKYLEFISEAII
Bloodworm	VAALGAKVLAQIGVAVSHLGDEGKMVAQMKAVGVRHKGYGNKHIKAQYFEPLGASLL
Yellow lupin	LQA-HAGKVFKLVYEAAIQLEVTGVVVTDATLKNLGSVHVSK-GVADAHFPVVKEAIL
Sperm whale	HVLHSRHPGDFGADAQGAMNKALELFRKDIAAKYKELGYQG
Bloodworm	SAMEHRIGGKMNAAAKDAWAAAYADISGALISGLQS
Yellow lupin	KTIKEVVGAKWSEELNSAWTIAYDELAIVIKKEMDDAA
In contract t	he following regults are from applying the same multiple sequence alignment
n contrast, t	ne following results are nom apprying the same multiple sequence angliment
program to the s	equences alone.
Sperm whale	VLSEGEWQLVLHVWAKVE-ADV-AGHGQDILIRLFKSHPETLEKFDRFKHLKTEAEMKA
Bloodworm	GLSAAQRQVIAATWKDIAGADNGAGVGKKCLIKFLSAHPQMAAVFG-FSGA
Yellow lupin	GALTESQAALVKSSWEEFN-ANI-PKHTHRFFILVLEIAPAAKDLFS-FLKGTSEVPQ

	•	:	•	:	•	:		:	•	:	•	:
Sperm whale	SE-DLK	KHGVTV	'LTALG-	-AII	KKKGI	HEAE-	LKPL	AQSH-	AT	KHKIP	IKYLF	EFIS
Bloodworm	SDPGVA	ALGAKV	LAQIG	/AVSHI	GDEGI	KMVAQ-	-MKAV	GVRHM	(GYGN)	KH-IK.	AQYFI	EPLG
Yellow lupin	NNPELQ	AHAGKV	FKLVYE	EAAIQI	EVTGV	/VVTDA	TLKNL	GSVHV	/SKG-	VA	DAHFF	PVVK
	•	:	•	:	•	:	•	:	•	:	•	:
Sperm whale	EAIIHV	LHSRHF	GDFGAI	DAQGAM	INKALI	ELFRKE	JAAKY	KELGY	/QG			
Bloodworm	ASLLSA	MEHRIG	GKMNAA	AKDAV]		-AAAY	ADIS-	GAL	ISGLQ	3	

The results were checked against the published structural alignments [LC80],[BCL87], and it can be stated that the structure-based calculation performed somewhat better than the purely sequence-based one. However, extensive tests on a variety of systems are required to evaluate the effectiveness of the method properly. We suggest that the results presented here encourage further development of the approach.

Yellow lupin

EAILKTIKEVVGAKWSEELNSAW-----TIAYDEL----AIV--IKKEMDDAA

Conclusions

We have designed and implemented a simple method for multiple structural alignment, using a one-dimensional representation of the conformation of a polypeptide chain, combined with the sequence, and standard multiple *sequence* alignment methods to perform the alignment.

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