

## 05-Molecular Modelling and Design for Proteins and Drugs

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After scanning the sequence against the library of 3D profile fragments, each residue in the given sequence normally has numerous matches with residues from different profile fragments with different scores. The score indicates the preference of this residue adopting the conformation of the matched residue in the profile fragment. We can make a scatter plot of the main chain conformation ( $\phi$  and  $\psi$  angles) of all the matched residues from the profile fragments in the library. The number of residues which fall into each of the three regions ( $\alpha$ -helix,  $\beta$ -strand or coil) are calculated, weighted by the scores matched by that fragment. The distribution of the three states for all the structures in the profile fragment library are also calculated. The preference of each residue in one of the three conformation states is calculated using a statistical inference algorithm. The secondary structure state of a residue is assigned to be the one with the highest preference. Tests using 40 structurally non-homologous structures indicate that the method has an overall prediction accuracy of 64%.

**MS-05.01.04 DESIGN STRATEGY FOR PROTEIN STABILITY: STABLE LOCAL CONFORMATION IN CONSISTENCY WITH THE GLOBAL CONFORMATION.** By K. Ishikawa, S. Kimura, K. Morikawa, S. Kanaya and H. Nakamura, Protein Engineering Research Institute, 6-2-3 Furuedai, Suita, Osaka 565, Japan.

Various mechanisms of the protein stability have so far been studied, and several strategies to enhance the stability are now proposed. We have recently made lots of mutant proteins of ribonuclease HI from *Escherichia coli* (*E. coli* RNase HI, 17.6kDa), and studied the conformational stability and their crystal structures. As a result, several of the mutant proteins obtained the remarkable thermal stability, due to only very local amino acid replacements; For example, Lys95 $\rightarrow$ Gly is considered to stabilize the local left-handed  $\alpha$ -helical conformation by a Gly residue (S. Kimura *et al.*, *J. Biol. Chem.*, 1992, **267**, 22014-22017). His62 $\rightarrow$ Pro may stabilize the short turn structure (K. Ishikawa *et al.*, *Protein Eng.*, 1993, **6**, 85-91). Val74 $\rightarrow$ Leu, fills the cavity in the hydrophobic core (K. Ishikawa *et al.*, *Biochemistry*, 1993, in press). All three mechanisms are localized, and the characteristic features of individual amino acids contribute the increase of the thermal stability. Analyses of the crystal structures of the wild-type and mutant proteins of *E. coli* RNase HI less than 1.8Å resolution reveal that the global conformations of all those mutant proteins deviate very little from that of the wild-type protein. It means that those local structural changes can be permitted and even suitable for the original global conformation. The additivity of the mutations was confirmed (S. Kimura *et al.*, *J. Biol. Chem.*, 1992, **267**, 21535-21542), and the structural analysis of the associated protein from *Thermus thermophilus* show that these local mechanisms are used in the thermophilic protein (K. Ishikawa *et al.*, *J. Mol. Biol.*, 1993, **230**, 529-542).

**MS-05.01.05 RULE-BASED APPROACHES TO COMPARATIVE MODELLING** by Z.Y. Zhu, M. S. Johnson, H. Wako, R. Sowdhamini, N. Srinivasan, K. Guruprasad, Z. Sun, B. Reddy, S. Rufino, Y. Edwards, T. Blundell\*. Imperial Cancer Research Fund Unit of Structural Molecular Biology, Department of Crystallography, Birkbeck College, University of London, Malet Street, London WC1E 7HX, UK

Comparative modelling can be envisaged as two steps. The first is to solve the inverse folding problem: to define all those sequences that can adopt a particular fold. Operationally this is more usefully posed as defining whether a new sequence belongs to any of the known folds. It involves projecting restraints from a three-dimensional structure on to a one dimensional sequence. For this step we have calculated amino acid substitution tables in terms of local structural environmental parameters, which can be used to generate sequence templates for secondary structures, structural motifs and tertiary folds. The second step is to use the sequence, together with the knowledge that the protein belongs to a family of known fold, to construct a model. This form of protein modelling and prediction involves placing constraints from a known fold on a related protein sequence. The two steps require similar knowledge of the structures of protein families, and this knowledge can be expressed as rules that relate both local and global three-dimensional structure to patterns in the sequence of amino acids in a polypeptide chain. The method is comparative but exploits a broader knowledge-base of non-homologous protein structures.

**PS-05.01.06 COMPARATIVE STRUCTURAL AND STEREOELECTRONIC STUDY OF PINACIDIL, DIAZOXIDE AND CROMAKALIM, POTASSIUM CHANNEL OPENERS BELONGING TO THREE DIFFERENT CHEMICAL CLASSES.** By L. Dupont<sup>a</sup>, B. Pirotte, P. de Tullio, B. Masereel, M. Schynts and J. Delarge<sup>b</sup>.

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Over the past few years the number of chemical agents with K<sup>+</sup> channel opening properties has greatly expanded. They are separated into distinct chemical classes typically exemplified by cromakalim (a benzopyran), pinacidil (a pyridylalkyl-cyanoguanidine), diazoxide (a benzothiadiazine), nicorandil (a pyridinic nitro compound), minoxidil sulfate (a pyrimidine derivative) and RP 49356 (a pyridinic thioformamide). For the three best studied K<sup>+</sup> channel openers, the rank order of potency for vascular smooth muscle relaxation was found to be cromakalim > pinacidil > diazoxide whereas for their activity on insulin secreting cells, the order was diazoxide > pinacidil > cromakalim (Newgreen, D.F., Bray, K.M., Mellarg, A.D., Weston, A.H., Duty, S., Brown, B.S., Kay, P.B., Edwards, G., Longmore, J. and Southerton, J.S., *Br. J. Pharmacol.*, 1990, **100**, 605-613; Lebrun, P., Antoine, M.H., Devreux, V., Hermann, M., Herschuelly, A., *J. Pharmacol. Exp. Theor.*, 1990, **255**, 948-954). Of particular interest is the intermediate position of pinacidil between diazoxide and cromakalim. Moreover, pinacidil could be regarded as a fairly good structural analog of diazoxide.

The present work tries to evaluate the level of structural analogy between these three classes of compounds by using crystallographic and infographic data. A systematic search was performed with SYBYL (Tripos Associates Inc., St Louis, Missouri, USA) starting from the X-ray conformation of pinacidil optimized by the Tripos force field maximin2 energy minimizer. The analysis of the search process exhibits four interesting low energy conformations. The four selected geometries have been optimized using the semiempirical method AM1 (MOPAC 5.0) and have been compared in terms of total energy calculation: the lowest energy conformation is actually the one found in crystal. A comparative study was undertaken on conformations and stereoelectronic properties of pinacidil, diazoxide and cromakalim to highlight the similarities which could be related to