Protein homology modelling and its use in South Africa

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HOMOLOGY MODELLING IS AN IMPORTANT computational technique, within structural biology, to determine the 3D structure of proteins. It uses available high-resolution protein structures to produce a model of a protein of similar, but unknown, structure. We describe the essential steps in the process, and discuss the circumstances in which homology modelling is likely to lead to a useful result. Homology modelling plays a valuable role in drug design, and we illustrate this by one example, anti-SARS inhibitors. In South Africa, homology modelling has been applied to proteins that may be relevant for drug design in connection with diseases as well as in other potential industrial applications. The use to date has been limited, however, so this article aims to introduce this useful and cost-effective technique to a wider community.

Introduction

It is almost 50 years since the first protein crystal structure, of myoglobin, was solved.1,2 With advances in experimental structural biology techniques as well as in whole-genome sequencing in the late 1990s, the excitement of solving a single-crystal structure has been replaced by determining protein structures on a large scale. This was the beginning of a new field, structural genomics, with active centres around the world.3–4 The contribution of structural genomics to the Protein Data Bank (PDB, an electronic repository of 3D structures of proteins and nucleic acids) is considerable, and nowadays these initiatives contribute approximately half of the new structurally characterized families of proteins.5

There are currently (January 2008) 44 820 protein structures in the PDB (Table 1). Although this number is increasing rapidly, there remains a huge gap between the number of available gene sequences and experimentally solved protein structures. X-ray crystallography, NMR and electron microscopy proceed much more slowly than genome sequencing; and since many more genome sequences are on the way, this gap must surely grow. So, what could be the solution? Structural genomics projects aim to determine experimentally at least one representative 3D structure for every protein family, and then to exploit the fact that proteins from the same family are evolutionarily related and share similar sequences and structures.6 This means that instead of trying to characterize the structure of every protein experimentally, we may start from known representative structures and use computational methods to predict the structure of related proteins.

This method has become known as homology modelling. According to the New York Structural Genomics Research Consortium, for each new structure, on average about 100 protein sequences without any prior structural characterization could be modelled at least at the fold level.7 Why is there this great effort to solve protein structures? Because they carry large amounts of information, and influence drug discovery, as one example of an application, at every stage in the design process.8,9 HIV/AIDS drugs such as Agenerase and Viracept were developed using the crystal structure of HIV protease.10,11 Thus, no matter how obtained (experimentally, computationally or using both approaches), 3D protein structure is of undeniable importance.

Structural biology in South Africa is a relatively new field that cannot yet be regarded as fully established. In this review, we discuss one of the techniques, homology modelling, and its applications. Our aim is to familiarize more research groups in biology with homology modelling, as it has great potential and can be learnt relatively easily and quickly.

Steps in homology modelling

Homology modelling seeks to predict the 3D structure of a protein based on its sequence similarity to one or more proteins of known structure. The method relies on the observation that the structural conformation of a protein is more highly conserved than its amino acid sequence. Homology modelling can be divided into four steps: template identification, alignment, model building and refinement, and validation (Fig. 1), with various computational tools (Table 2) available for each step. More detailed information is available in recently published reviews.10–16

Template identification

Template identification is the critical first step. It lays the foundation by identifying appropriate homologue(s) of known protein structure, called template(s), which are sufficiently similar to the target sequence to be modelled. A simple search submits the target sequence to programs such as BLAST17 or FASTA.18 However, these programs work well only for alignment of sequences with high similarities. Methods such as PSI-BLAST19 and ScanPS20 have recently increased the possibility of detecting distant homologues.

These methods often suggest several candidate templates. The ideal is to identify the template(s) which has the highest percentage identity to the target, has the highest resolution, and has structures with (or without) appropriate ligands and/or cofactors. It may be that there is no candidate template that is best according to all criteria, in which case the choice is a matter of judgment and perhaps of trying different templates.

Alignment

The next step involves creating an alignment of the target sequence with the template structure(s). This is a vital step and there are various ways to ensure high accuracy. The target and template sequence can be aligned with a protein domain family alignment retrieved from Pfam,21 or a custom alignment can be generated from all relevant sequences retrieved via BLAST. Programs such as Clustal,22

Table 1. Number of protein and protein/nucleic acid complex structures obtained by various experimental methods, available in the PDB as of 29 January 2008 (modified from www.rcsb.org/pdb/statistics/holdings.do).

<table>
<thead>
<tr>
<th>Experimental method</th>
<th>Protein type</th>
<th>Molecule type</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Proteins</td>
<td>Protein/nucleic acid complexes</td>
<td></td>
</tr>
<tr>
<td>X-ray diffraction</td>
<td>38 541</td>
<td>1770</td>
<td>40 311</td>
</tr>
<tr>
<td>NMR</td>
<td>6 080</td>
<td>137</td>
<td>6 217</td>
</tr>
<tr>
<td>Electron microscopy</td>
<td>112</td>
<td>41</td>
<td>153</td>
</tr>
<tr>
<td>Other methods</td>
<td>87</td>
<td>4</td>
<td>91</td>
</tr>
<tr>
<td>Total</td>
<td>44 820</td>
<td>1952</td>
<td>46 772</td>
</tr>
</tbody>
</table>
Muscle\(^{23}\) and TCoffee\(^{24}\) can be used to construct the alignment. Sometimes structural alignments are preferred, especially for distantly related sequences, because structure is more conserved than sequence. 3DCoffee\(^{25}\), FUGUE\(^{26}\) and mGenThreader\(^{27}\) are well-known structural alignment programs. MEME\(^{28}\) provides information about conserved motifs found in aligned sequences, and can be used to guide the alignment.

The alignment can and should be optimized manually. By including biological information such as the solvation environment of an amino acid, better-informed changes to the alignment can be made by the user. This type of information is not often available to the alignment program.

**Model building and refinement**

Although the theory behind building a protein homology model is complicated, using available programs is relatively easy. Several modelling programs are available, using different methods to construct the 3D structures. In segment matching methods, the target is divided into short segments, and alignment is done over segments rather than over the entire protein.\(^{29}\) Satisfying spatial restraints is the most common method. It uses either distances or optimization techniques to satisfy the spatial restraints. The method is implemented using the popular program, Modeller,\(^{30}\) which includes the CHARMM\(^{31}\) energy terms that ensure valid stereochemistry is combined with spatial restraints. There are several stand-alone modelling programs available such as WHAT IF\(^{32}\). Web servers such as SwissModel and the Rosetta server make it even easier to generate a model.

A problem regularly encountered in homology modelling is loops. A general guideline is that any insertion/loop longer than about five residues should be omitted. There are programs which try to model loops such as Modeller and its more specialized loop modelling server, Modloop.\(^{33}\)

The initial model may have suboptimal bond angles and lengths. Such deficiencies can be adjusted by an energy minimization procedure, but a difficulty is that it moves atoms towards a local minimum, which may not be a global minimum over all possible conformations. Alternatively, molecular dynamics can be used, in which the motion of the whole protein is modelled.\(^{34,35}\)

**Validation**

After being built, the model needs to be validated. One of the most thorough structure checking programs is WHATCHECK.\(^{36}\) Other programs such as Procheck,\(^{37}\) and ANOLEA\(^{38}\) at SwissModel evaluate fewer parameters. The best validation combines common sense, biological knowledge and results from analytical tools. Some models will need further refinement. There is a cycle between building–validating–refining. Most refinement involves adjusting the alignment.

**Advantages and limitations of homology modelling**

Homology modelling is a relatively easy technique. It takes much less time to learn, to do the calculations and obtain a result, than an experiment. Nor does it require expensive experimental facilities, just a standard desktop computer. In the absence of high-resolution experimental structures, therefore, homology modelling can be of much value.

However, the quality and accuracy of the homology model depend on several factors. The technique requires a high-resolution experimental protein structure as a template, the accuracy of which directly affects the quality of the model. Even more importantly, the quality of the model depends on the degree of sequence identity between the template and protein to be modelled.\(^{39-41}\) Alignment errors increase rapidly when the sequence identity is less than 30%. Medium accuracy homology models have between about 30% and 50% sequence identity to the template. They can facilitate structure-based prediction of target for ‘druggability’, the design of mutagenesis experiments and the construction of in vitro test assays. Higher accuracy models are typi-
homology modelling. Severe acute respiratory syndrome (SARS) was identified in China in 2002 and quickly spread to other countries. The cause was a new coronavirus (CoV). Soon afterwards, whole genomes of different SARS-CoV strains were solved. Main protease (Mpro), which has an important role in virus replication, became an immediate drug target. CoV-Mpro has 40% and 46% sequence identity to transmissible gastroenteritis coronavirus (TGEV) Mpro, and human coronavirus 229E, respectively, and X-ray structures were already available. Several groups released the homology model of the protease in May 2003. After a comparison of the inhibitor complexed with TGEV-Mpro with available inhibitor complexes in PDB gave a similar inhibitor-binding mode in the complex of human rhino-virus type 2 (HRV2) 3C proteinase with AG7088. At the time, AG7088 was in clinical trials for the treatment of the human rhino-virus that causes the common cold. AG7088 was docked into the substrate-binding site of the SARS-CoV-Mpro model, indicating that it would be a good starting point for the design of anti-SARS drugs. Shortly thereafter, it was shown that AG7088 does indeed have anti-SARS activity in vitro.

**Protein homology modelling in South Africa**

Homology modelling in South Africa has involved projects on disease-related proteins as well as on proteins with industrial applications. We highlight some of these below.

Sleeping sickness is a severe disease in Africa, caused by trypanosomes. There are problems with current drugs such as their high toxicity and ineffectiveness due to drug resistance. The enzyme, glycerol-3-phosphate dehydrogenase (GPDH), from the glycolytic metabolic pathway of *Leishmania mexicana*, has been elucidated as a possible new drug target.49 At Rhodes University, this information, together with the availability of a homologue crystal structure of *Leishmania mexicana*, led to the construction of a homology model of NAD-dependent glycerol-3-phosphate dehydrogenase of *Trypanosoma brucei rhodesiensese*.50 The model was further used in molecular dynamic simulations to show that the protein can be used for drug design.

Ticks transmit pathogens and toxic compounds, severely affecting the health of the host, even leading to death. The development of tick control methods requires an understanding of tick–host interactions. A group at the University of Pretoria used homology modelling to study the structure of a thrombin inhibitor, savignin, from the tick, *Ornithodoros savignyi*.51 The aim was to understand the anti-clotting function of savignin, with the long-term goal of the design of anti-haemostatic pharmaceuticals.

Malaria affects 300–500 million people annually and kills approximately 2 million.52 *Plasmodium falciparum* is the most lethal malaria parasite infecting humans, and the emergence of drug-resistant strains demands a search for new targets. This depends on structural information. However, there are difficulties with malaria proteins. Crystalization trials are limited due to the difficulty of expressing malaria proteins in high yield and pure state, as well as by the presence of long and unordered inserts. These *Plasmodium*-specific inserts...
also make it difficult to construct homology models. A group at the University of Pretoria has computed homology models of malaria proteins. In the polyamine pathway, models of ornithine decarboxylase (ODC)\(^{39}\) and S-adenosylmethionine decarboxylase\(^{41}\) were prepared. The comparison of these models with the human enzyme showed differences in the active site, which might allow identification of parasite-specific inhibitors. \(P.\) falciparum spermidine synthase (PfSpdSyn) is another enzyme in this pathway. Molecular dynamics of the homology model demonstrated the mechanism of the amino-propyltransferase action of PfSpdSyn.\(^{36}\) The folate biosynthetic pathway is also important for drug targeting. Homology models of the bifunctional enzymes hydroxymethylpterin pyrophosphokinase-dihydropterate synthase (PPPK-DHPS) were constructed. The aim was to investigate possible causes of the development of resistance in mutations.\(^{36}\) In the vitamin B\(_6\) system, an homology model of pyridoxal kinase (PdxK) showed the development of resistance in mutations. In over two decades, mostly due to its biocatalytic activity (in acylamide and nicotinamide production). Its importance extends also to its use for environmental remediation by removing nitriles from waste streams. The group at UWC generated the homology model of the enzyme from thermophile \(Bacillus pallidus\) RAPc8 as well as a \(122\)-amino-acid accessory protein involved in thermostable NHase expression (P14K), a homologue of the \(2\text{Fe-2S}\) class of ferredoxins.\(^{36}\) Modelling of the P14K protein structure suggested that the protein functions as a subunit-specific chaperone, and helps with the folding of the NHase and the formation of NHase heterotetramer.

Lipases are a widely used group of biocatalysts,\(^{37}\) and can be produced in large quantities from fungi and bacteria. They do not require cofactors nor do they catalyse side reactions, and are thus very attractive for industrial applications. They hydrolyze triacylglycerols to fatty acids and acylglycerols. Lipases are used for the synthesis of biopolymers, biofuel, and the production of enantiopure pharmaceuticals, agrochemicals, cosmetics and flavour compounds. A group from the University of the Free State investigated an extracellular lipase from \(Pseudomonas luteola.\)\(^{41}\) Homology modelling of this lipase enabled them to compare it with the other available lipase structures, concluding that this enzyme could also have biocatalytic applications.

**Conclusion**

We have given an overview of what homology modelling is all about: procedure, applications, advantages and limitations, as well as its current use in South Africa. Homology modelling is entirely a computational process and much easier to implement than the experimental path to structural information about a protein, although it relies on suitable experimental structures being already known. Applications of homology modelling can range from design of the next experiment in an ongoing biochemical investigation, to the discovery of drugs with important disease control properties. On the other hand, in circumstances where the homology model may be of only limited accuracy, the results may require experimental verification.

Because homology modelling has been used in South Africa in only a limited way, we hope that this article will introduce a broader community to this relatively simple and cost-effective technique, and so lead to practical applications, especially in tackling our own problems—of HIV/AIDS, tuberculosis and malaria, for example—rather than waiting for solutions from elsewhere.

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