Structural bioinformatics

PSIbase: a database of Protein Structural Interactome map (PSIMAP)

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Received on November 1, 2004; revised on January 27, 2005; accepted on February 28, 2005

Advance Access publication March 3, 2005

ABSTRACT

Summary: Protein Structural Interactome map (PSIMAP) is a global interaction map that describes domain–domain and protein–protein interaction information for known Protein Data Bank structures. It calculates the Euclidean distance to determine interactions between possible pairs of structural domains in proteins. PSIbase is a database and file server for protein structural interaction information calculated by the PSIMAP algorithm. PSIbase also provides an easy-to-use protein domain assignment module, interaction navigation and visual tools. Users can retrieve possible interaction partners of their proteins of interests if a significant homology assignment is made with their query sequences.


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Supplementary information: Supplementary material is available at http://psibase.kaist.ac.kr/Doc/supplementary_material.htm

INTRODUCTION

Most proteins function by interacting with other molecules. Therefore, it is important to investigate the interaction partners of proteins. Recently, high-throughput experiments, such as yeast (Uetz et al., 2000) and fly (Giot et al., 2003) proteomes, have enabled us to elucidate the interaction networks on a large scale. These large-scale experiment results are collected and well-curated into interaction databases such as the Database of Interacting Proteins (DIP) (Salwinski et al., 2000), Biomolecular Interaction Network Database (BIND) (Bader et al., 2003) and Molecular INTeraction database (MINT) (Zanzoni et al., 2002). There have also been computational approaches to map and predict the protein interactome in a genomic context using gene fusion and gene neighborhood methods (Huynen et al., 2000).

In parallel with the above methods, PSIMAP (Protein Structural Interactome map) has introduced a new mapping protocol in protein structural interactome study. An underlying concept of PSIMAP is homologous interaction: the interaction among protein structures is conserved as closely as the protein structures themselves (Park et al., 2001; Aloy and Russell, 2002; Aloy et al., 2003). With PSIMAP, we can view protein interactions in terms of family–family interactions, as well as individual protein–protein interactions. PSIMAP covers interaction information from both gene fusion style protein sequence level interaction and physical interaction within complexes or multi-domain proteins.

Here, we introduce PSIbase: the PSIMAP web server and database. It contains (1) domain–domain and protein–protein interaction information from proteins whose 3D-structures are identified, (2) a protein interaction map and its viewer at protein superfamily and family levels, (3) protein interaction interface viewers and (4) structural domain prediction tools for possible interactions by detecting homologous matches in the Protein Data Bank (PDB) from query sequences. Structural interaction data, in flat file format, can be downloaded from PSIbase (http://psibase.kaist.ac.kr/Download/download.shtml) for further analyses. It contains the smallest distance between two domains and the number of residue pairs that is within the threshold distance according to the PSIMAP algorithm. It not only provides raw data files, but it also serves biologists who need to look up the interaction partners of their proteins of interest. Simply putting a protein sequence is enough to search for possible interaction partners (interlogs). As the possible predicted domains of query sequence are based on a structural assignment protocol, users can see the interlogs’ 3D structures if they accept the prediction made by PSIbase. For structural domain assignment, we used two databases and two algorithms. They were the SCOP (http://scop.kaist.ac.kr/scop) and PSI-BLAST (http://www.ncbi.nlm.nih.gov/BLAST/) with a hidden Markov model package (HMMER, http://hmmer.wustl.edu/). We believe that PSIbase is useful for those in the fields of structural bioinformatics and molecular biology.

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**PSIMAP ALGORITHM**

The basic mechanism to check interactions between any two domains or proteins is the calculation of the Euclidean distance in order to see if they are within a certain distance threshold. PSIMAP checks every possible pair of structural domains in a protein to see if there are at least five residue contacts within a 5 Å distance (5–5 rule). The current PSIMAP protocol has three methods. They are the Full Atom Contact (FAC) PSIMAP, Sampled Atom Contact (SAC) PSIMAP and Bounding Box Contact (BBC) PSIMAP (Dafas et al., 2004). (The supplementary material provides in-depth information about the three different PSIMAP algorithms.)

The FAC calculates all the atomic contacts among two or more protein structural domains. FAC PSIMAP is the most accurate of the three, as we take into account all the atoms in domain pairs. The SAC and BBC algorithms are approximations of FAC. Their main purpose is to reduce the time taken in constructing PSIMAP. The BBC algorithm is a radically different approach, using a bounding box algorithm to dramatically reduce the time of computation. Dafas et al. introduced a bounding box and convex hull algorithm that can reduce the search space.

**DATABASE ACCESS**

The PSIbase server is available at http://psibase.kaist.ac.kr/. There are three different query interfaces to access the PSIbase. All queries are funneled into a web page that shows protein domain interactions with their partners.

First, PSIbase provides a simple search interface that looks up keywords or database accession IDs. Figure 1 shows the search result of ‘ligase’ as a query against 12 annotated DB resources (listed on the PSIbase webpage). Out of the 12, multiple matches for the query ‘ligase’ are listed up from the following databases: PDB, SCOP, TIGRFAMs, Swiss-Prot, ProDom, Pfam, Prosite and Interpro. There are three tools to view interaction interface structures: Chime (http://www.mdli.com), Imol (http://jmol.sourceforge.net) and Interfacer (http://www.interfacer.org). Interfacer is a slow but advanced protein interface viewer with surface representation capability.

The second PSIbase query interface is a protein structural domain assignment utility that accepts protein sequences from users. There are two domain assignment algorithms available in PSIbase. One is a homology-based sequence search by PSI-BLAST utilizing the ISL (see Introduction) and the other is the HMMER profile search algorithm. These two are complementary in terms of the coverage in the assignment.

The last PSIbase query interface accepts specific domain IDs at SCOP family or superfamily levels. There are several levels to determine interactions among query domains. For example, interacting partners of a specific query domain can be identified within a specified interaction depth (the maximum depth limit is 4). Interactions between two or more input query domains can also be identified. Additionally, PSIbase is equipped with a simple open-source Java applet program that shows the interaction network of each query.

**CONCLUDING REMARKS**

There are 1294 superfamilies and 2327 families in SCOP 1.65. On average, PSIbase covers 87% (1136/1294) of SCOP superfamily interactions, indicating that the majority of SCOP superfamilies have interacting partner information. In the supplementary material, Table 2 shows the 20 most interactive superfamilies in PSIbase. These can be regarded as the most central interaction components in interactomes, so we call them the ‘interactome core’. This core contains proteins with energy metabolism, RNA and DNA binding, and other key biological processes that have existed since the very early
days of interaction networks (Bolser et al., 2003). The interactions of non-protein molecules in cells are critical in biological functions. In the next version, PSIbase and PSIMAP will cover interactions between proteins and non-proteins such as nucleic acids and small molecules.

**ACKNOWLEDGEMENTS**

We thank Mr. Chung MoonSoul for donating $25 million to the Department of Biosystems at KAIST. This project was funded by IMT-2000 C3-4 grants from the Ministry of Information and Communication of Korea and a grant from KRRIB Research Initiative Program. J.B. is supported by Biogreen21. We thank Maryana Bhak for editing and commenting on this manuscript. We also send our loving gratitude to the anonymous reviewers for their precious comments.

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