IRView: a database and viewer for protein interacting regions

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ABSTRACT

Summary: Protein–protein interactions (PPIs) are mediated through specific regions on proteins. Some proteins have two or more protein interacting regions (IRs) and some IRs are competitively used for interactions with different proteins. IRView currently contains data for 3417 IRs in human and mouse proteins. The data were obtained from different sources and combined with annotated region data from InterPro. Information on non-synonymous single nucleotide polymorphism sites and variable regions owing to alternative mRNA splicing is also included. IRView web interface displays all IR data, including user-uploaded data, on reference sequences so that the positional relationship between IRs can be easily understood. IRView should be useful for analyzing underlying relationships between the proteins behind the PPI networks.

Availability: IRView is publicly available on the web at http://ir.hgc.jp/.

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1 INTRODUCTION

Protein–protein interactions (PPIs) and their networks play central roles in governing cellular processes. Recently, much effort has been put into collecting binary interaction data (e.g. Rual et al., 2005) to dissect PPI networks. These interaction data have been compiled in public biomolecular interaction databases. For example, BioGRID (Breitkreutz et al., 2008) and IntAct (Aranda et al., 2009) are major molecular interaction databases of PPIs. These databases primarily contain PPI data at the protein level, namely pairs of protein names. Two recent articles, one of them from our group, have pointed out that there are major molecular interaction databases of PPIs which focus on the regions required for PPIs. IRView contains, as a primary data source, IRs that were determined using the in vitro virus (IVV) method (human data: Miyamoto-Sato et al., 2010; mouse data: Horisawa et al., 2004 and Miyamoto-Sato et al., 2005) and the yeast two-hybrid (Y2H) method. DOMINO (Ceol et al., 2007) is a database of domain–domain interactions that is similar in scope to IRView. DOMINO stores data on IRs described in the scientific literature and applies the existing domain/motif names from the InterPro (Hunter et al., 2000) to dissect PPI networks. IRView also supports a viewer that allows users to compare the positional relationships of IRs in protein reference sequences and in 3D structures (when available). IRView should be useful for investigating the hidden relationships between the proteins behind protein interaction networks.

2 CONTENTS AND FEATURES

2.1 The IR and other functional region data

To comprehend the complicated relations underlying PPIs in more detail, further refined interaction data are required. In this article, we describe IRView, a database and viewer for IRs of proteins, which focuses on the regions required for PPIs. IRView contains, as a primary data source, IRs that were determined using the in vitro virus (IVV) method (human data: Miyamoto-Sato et al., 2010; mouse data: Horisawa et al., 2004 and Miyamoto-Sato et al., 2005) and Y2H (Sugaya et al., 2007) methods. Over half the IR data (2629 IRs) were derived from the results obtained using the IVV method (Miyamoto-Sato et al., 2010). Although the current data and sources for IRView are limited, we plan to add our original, experimental data and data extracted from the literature. The IR data can be downloaded in the PSI-MITAB format (Kerrien et al., 2007) file.

The conserved domains and motifs data that were used to visualize the positional relationships of the IRs were retrieved from the InterPro database (Hunter et al., 2009). Data on the nsSNPs that can lead to amino acid changes and potentially affect protein interactions (Mendelsohn, 2004; Schuster-Bockler and Bateman, 2008) were

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obtained from the dbSNP (Smigielski et al., 2000). Variable regions derived from alternative mRNA splicing that may potentially affect protein interactions (Resch et al., 2004) were defined based on the results of pair-wise alignments between the various isoforms. The 3D structure data were downloaded from the Protein Data Bank. When 3D models of complexes were available, information about the interacting amino acid residues on different peptide chains (defined as amino acids that were within a distance <4.0 Å of each other) were also added to the IR data.

2.2 Reference sequence-centered map

One of the main features of IRView is that all positional data are standardized to positions in reference sequences. IRView uses the NCBI RefSeq sequences as the reference protein sequences. When different isoforms of a protein are recorded in the RefSeq database, the longest sequence was selected as the representative sequence (RS) and the others were treated as related sequences. Standardization of the position data was achieved by pair-wise alignments between the RS and the other related sequences using ClustalW 2.0 (Larkin et al., 2007). As a result of this standardization, users can easily capture the positional relationships between independently annotated regions from different sources. Using the standardized position data, IRView can provide information about the positional relationships between the IRs in one protein sequence that interact with different proteins. In addition, IRView provides information on the positional relationships between IRs and other annotated regions (e.g. InterPro regions). Whether or not an IR overlaps with any other annotated region is indicated by special icons that accompany each IR entry. Of the 3417 IRs in the current version of IRView, 1492 IRs overlap with known domain/motif regions, 521 IRs overlap with nsSNPs, 207 IRs overlap with structured regions, 102 IRs overlap with variable regions derived from alternative mRNA splicing and 691 IRs overlap with other IRs.

3 DESCRIPTION OF THE IRVIEW INTERFACE

IRView supports searches by protein name, gene symbol, NCBI Entrez GeneID, RefSeq ID, species name and free keywords. IRView also supports the use of field specifiers to limit the scope of searches. Query results are returned as a list of RSs which correspond to a ‘Region information’ page consisting of a number of sections: ‘Representative sequence’, ‘Related sequences’, ‘Structured regions’, ‘Domain/Motif regions’, ‘Variable regions’, ‘Non-synonymous SNPs’, ‘Interacting regions’ and ‘Contacting amino acids’. To make it easier to compare two or more annotated regions (e.g. comparing variant regions with IRs to infer the impact of alternative splicing), unnecessary subsections can be collapsed. Details of these subsections are described in the Help page of IRView.

IRView also possesses a system for mapping specific regions to 3D structure(s) when the corresponding structure data are available. Users can map regions of interest to the 3D structure individually or simultaneously via the in-lined Jmol applet (http://www.jmol.org/) on any Java-enabled web browser.

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