Comparative interactomics analysis of protein family interaction networks using PSIMAP (protein structural interactome map)

Daeui Park¹,†, Semin Lee²,†, Dan Bolser³, Michael Schroeder⁴, Michael Lappe⁵, Donghoon Oh¹, and Jong Bhak²,6*

¹Object Interaction Technologies Inc., CMS Building 705, 373-1, Guseong-dong, Yuseong-gu, Daejeon 305-701, KOREA, ²Department of BioSystems, KAIST, 373-1, Yuseong-gu, Daejeon 305-701, KOREA, ³Medical Research Council, Dunn Human Nutrition Unit, Cambridge, CB2 2QH, UK, ⁴Biotec and Department of Computing, TU Dresden, 108 01307 Dresden, GERMANY, ⁵Max-Planck-Institute for Molecular Genetics, Ihnestrasse 63-73, 14195 Berlin, GERMANY, ⁶NGIC, KRIBB, Daejeon 305-333, Korea.

ABSTRACT

Motivation: Many genomes have been completely sequenced. However, detecting and analyzing their protein-protein interactions by experimental methods such as co-immunoprecipitation, tandem affinity purification, and Y2H is not as fast as genome sequencing. Therefore, a computational prediction method based on the known protein structural interactions will be useful to analyze large-scale protein-protein interaction rules within and among complete genomes.

Results: We confirmed that all the predicted protein family interactomes (the full set of protein family interactions within a proteome) of 146 species are scale-free networks, and they share a small core network comprising 36 protein families related to indispensable cellular functions. We found two fundamental differences among prokaryotic and eukaryotic interactomes: 1) eukarya had significantly more hub families than archaea and bacteria, and 2) certain special hub families determined the topology of the eukaryotic interactomes. Our comparative analysis suggests that a very small number of expansive protein families led the evolution of interactomes and seemed to have played a key role in species diversification.

Contact: jong@kribb.re.kr

Supplementary Information: Supplementary data: Figure 1, Table 4

INTRODUCTION

Since 1995, over 250 genomes have been completely sequenced (Bernal, et al., 2001; Shendure, et al., 2004). The availability of such genomic sequence data allows us to conduct a comparative genomics study, yielding important information on developmental processes and disease defense mechanisms (Eichler and Sankoff, 2003; O’Brien, et al., 1999; Rubin, et al., 2000). Protein comparison using proteomes alone is, however, not sufficient to fully understand how the cellular machinery evolved over a long period of time. The fundamental differences among organisms cannot be explained fully by simply looking at the genes and proteins. A step forward would be to look at all the interactions among them (Ng, et al., 2003; von Mering, et al., 2002).

The full range of functional complexity and diversity in biological systems is probably the result of interactions among biological entities. The architectures and organization of such interactions are best represented as networks, for example, networks of interacting proteins that reflect biochemical pathways and genetic regulations. It has been reported that, due to functional constraints, biological interaction networks are tightly conserved (Bolser and Park, 2003).

The question is what is the basic similarity and difference that underlies the networks of interacting proteins in the completely sequenced genomes. To answer this question, the examination of many complete proteomes and their interactomes would be necessary. However, present experimental technology is not fast enough to map the molecular interactions of proteins for all the completely sequenced genomes. Consequently, computational methods for assigning and predicting protein interactions have been developed using genomic sequence data (Dandekar, et al., 1998; Enright, et al., 1999; Huynen and Bork, 1998; Marcotte, et al., 1999; Overbeek, et al., 1999; Pellegrini, et al., 1999; Tan, et al., 2004). The aim of these methods, however, has been more focused on discovering functional interactions rather than physical interactions.

Therefore, we introduced a structure-oriented protein interaction protocol: PSIMAP (Protein Structural Interactome Map) (Gong, et al., 2005; Park, et al., 2001). The interactions among structural protein families are
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fundamental to the workings of cells: in multi-domain polypeptide chains, in multi-subunit proteins, and in transient complexes among proteins that also exist independently. One critical aspect of PSIMAP is that it allows us to view interactions among protein domains in terms of their structural families to analyze the large-scale patterns and evolution of interactomes among species. (Fig. 1). PSIMAP extracts the exact molecular interaction information of proteins from the Protein Data Bank (PDB) (Berman, et al., 2000) and their domains from Structural Classification of Proteins (SCOP) (Murzin, et al., 1995). It has a predictive capacity that can be extended to a genomic scale with the assistance of bioinformatics.

We have built a high-throughput, homology-based interaction prediction method utilizing PSIMAP. The key advantages of the PDB-derived predictive method are that 1) it covers a many times larger data set for probable protein interactions, 2) it can reveal the history of interaction in genomes with the limited amount of experimental interaction information available at present, and 3) it provides explicit models of structural interactions that can be used in the validation of the predicted interactions. A disadvantage is that the individual protein interaction information cannot be readily verified. However, we found that 340 out of 591 (57.5%) human protein interactions in the Database of Interacting Proteins could be explained by structural interactions (Moon, et al., 2004).

To investigate the broad evolutionary trend in protein interaction networks, using conserved protein family interactions is more appropriate than individual protein interactions. For this reason, in our comparative analysis using PSIMAP, the precise verification of individual

Fig. 1. Global view of protein family interaction networks. (a) An overview of PSIMAP for all the protein family interactions (1895 protein family nodes, 2655 protein family interaction pairs). Each family is color-coded by its connectivity. More than one-fifth of families are part of the large connected cluster centering c.37.1.8 (G proteins). Other highly connected nodes are b.47.1.2 (eukaryotic proteases) and d.58.1.5 (ferredoxin domain). Most of the protein families have only 1–3 interaction partners. (b) Spherical layout of protein family interactomes for three species from three different superkingdoms: Protein family interaction networks for Thermoplasma acidophilum (archaea, 432 protein families, 552 protein family interaction pairs), Escherichia coli (bacteria, 856 protein families, 1100 protein family interaction pairs), and Homo sapiens (eukarya, 1126 protein families, 1624 protein family interaction pairs). The protein families were taken from SCOP. The protein family interactions were assigned by PSIMAP. Depending on the number of protein families and their interactions, the density of the network varies, representing the complexity of the interactomes.
578,625 protein sequences were used (Pruess et al., 2003). The European Bioinformatics Institute (EBI) and their protein sequences. 146 completely sequenced species from structural families (on which PSIMAP is based) to the particular proteome, we first assigned the known 3D protein structure assignment.

**METHODS**

**Protein structure assignment**

To construct the protein family interaction network in a particular proteome, we first assigned the known 3D structural families (on which PSIMAP is based) to the protein sequences. 146 completely sequenced species from the European Bioinformatics Institute (EBI) and their 578,625 protein sequences were used (Pruess et al., 2003). The 146 genomes represented species of archaea (15), bacteria (122), and eukarya (9). Among the 578,625 proteins, 296,630 (52%) had at least one SCOP family assignment. The SCOP database (Version: 1.65) is a manually-curated protein structural domain classification system. The actual structure assignment of proteins was conducted by the PSI-BLAST algorithm (Altschul et al., 1997), using the default BLOSUM62 amino acid substitution matrix as a threshold with 0.01 E-value (expectation value scoring). Several thresholds ranging from 0.0001 to 0.01 E-values were tested, and 0.01 was chosen after a manual inspection of results, as there was no explosion of erroneous matches in the iterative searching process of PSI-BLAST. Our experiments showed that 0.01 E-value provided coverage 2-3% higher than that of 0.0001 without sacrificing the assignment quality.

Our homology assignments were made at the SCOP family level of well classified and clearly distinguishable SCOP classes: a (all alpha proteins), b (all beta proteins), c (alpha and beta proteins (a / b)), d (alpha and beta proteins (a + b)), e (multi-domain proteins), f (membrane and cell surface proteins and peptides), and g (small proteins). 2091 (90%) out of 2327 SCOP families were assigned to at least one protein sequence. Then, 371 SCOP families having no interaction information in PSIMAP were filtered out.

Multi-domain proteins contain more than one SCOP domain. A problem in structural assignment is that erroneous alignment overlap can occur between two domain sequences on the same region of a multi-domain protein. To overcome this, we regarded domain sequences with an overlap of 15 or less amino acid residues in the alignments as two separate domains in the protein sequence.

**Mapping the protein family interaction**

To construct species-specific protein family interactomes, we used PSIMAP (see http://psimap.org and http://psibase.kaist.ac.kr). PSIMAP is a global interaction map that describes domain-domain and protein-protein interaction information for known PDB structures. It considers every possible pair of structural domains within a protein or complex to see if there are at least five residue contacts within a five Angstrom distance (Bolser et al., 2003). Although the number of PDB structures is relatively small in comparison to the sequence data, PSIMAP can cover the majority of known protein structural information (Fig. 1a) (Aloy and Russell, 2002).

All the predicted protein family interactomes are species specific, and our aim was to find any evolutionary trend among all the interactomes. Hence, the interactomes are based at the protein family interaction level, instead of the individual protein level. Figure 1b shows how interaction intensity is unevenly distributed in a spherical interaction network layout of three model species interactomes. The unique topologies of protein family interaction networks enabled us to compare and analyze them, in order to hypothesize on how the interactomes have expanded.

In terms of the coverage of the interactomes, we supposed the present PDB (the source of interaction information for our analysis) represented the majority of protein folds in nature. This is because the number of PDB entries is growing exponentially while the number of new folds is increasing very slowly. It has been reported that there may be fewer than 2000 distinct protein architectures in nature (Alexandrov and Go, 1994; Chothia, 1992; Orengo et al., 1994; Wang, 1996; Zhang, 1997). Therefore, the present structural interactome data represents a relatively complete set of distinct protein families, although this does not imply that all the possible family interactions have been observed in PDB. Also, as a significant portion of the unassigned genes represents trans-membrane proteins yet to be determined experimentally, we suggest that PDB and PSIMAP cover the majority of the existing soluble families in nature. Table 1 shows the subdivision of the 1720 molecular interactions is not necessary, and the reliability of the family interaction predictions is mostly dependent on accurate family detection in the genomes. Although particular organisms may inevitably have false or missing predictions, in general this should not greatly affect the overall comparative family level analysis.

Based on the comparative analysis results of protein family interaction networks reconstructed by PSIMAP (see Supplementary Table 1), we were able to get insight into how species diversified in terms of computationally predicted protein family interactomes and their common features. First, we found that all 146 species’ protein family interactomes share a small core network comprised of 47 protein family interaction pairs (including self- or homo-interacting pairs). Although small and probably incomplete in number, we can infer that the core is well-conserved in diverse life forms. A notable outcome was that a very small number of expansive protein families played a key role in the interactome growth and determined the characteristics of the interactomes of prokarya (archaea and bacteria) and eukarya. Eukaryotic interactomes have a distinctly different topology from prokaryotic interactomes regardless of proteome size.
observed protein families and the 2404 observed protein family interactions.

Table 1. Summary of structure assignment and comparative analysis of protein family interaction networks at the superkingdom level.

<table>
<thead>
<tr>
<th>Description</th>
<th>Archaea</th>
<th>Bacteria</th>
<th>Eukarya</th>
<th>All</th>
</tr>
</thead>
<tbody>
<tr>
<td>No. of species</td>
<td>15</td>
<td>122</td>
<td>9</td>
<td>146</td>
</tr>
<tr>
<td>No. of proteins</td>
<td>35,197</td>
<td>362,484</td>
<td>180,944</td>
<td>578,625</td>
</tr>
<tr>
<td>No. of structure assigned proteins</td>
<td>16,085</td>
<td>186,539</td>
<td>94,006</td>
<td>296,630</td>
</tr>
<tr>
<td>No. of families assigned to proteins</td>
<td>986</td>
<td>1526</td>
<td>1616</td>
<td>2091</td>
</tr>
<tr>
<td>No. of families assigned to proteins that can be covered by PSIMAP</td>
<td>1281</td>
<td>1340</td>
<td>1720</td>
<td></td>
</tr>
<tr>
<td>No. of protein family interaction pairs</td>
<td>1086</td>
<td>1691</td>
<td>1916</td>
<td>2404</td>
</tr>
</tbody>
</table>

*non-redundant count.

RESULTS AND DISCUSSION

The core protein family network of life

We found 36 commonly present protein families in the 146 species. They produced 47 protein family interaction pairs (1.3 links per family) that are predicted to be conserved across all species (Fig. 2c), while a third of protein family interaction pairs (31%, 734 out of 2404 total pairs) were counted in over 80 species (Fig. 2a, Supplementary Table 2). The statistical likelihood of forming the core network (36 families) in the 1720 protein families is $5e^{-9}$ under a Poisson distribution (i.e., not random).

A notable aspect of the core network is that there are only 17 hetero-interaction pairs which reflected that many protein families are self- or homo-interacting (e.g., homo-dimer proteins). There were 1358 homo- and 1251 hetero-interaction pairs in PDB. 60% of archaeal interactomes had homo-interaction pairs. Bacteria had 59%. Eukarya had the lowest rate at 53%. Eukaryotic homo-interaction is statistically significantly lower (Kruskal-Wallis test) than the rest. This indicates that eukaryotic interactomes may have expanded their interaction partner repertoires more diversely than archaea and bacteria.

16 out of the 36 core protein families (44%) were related to protein translation. Notably, c.37.1.8, the most highly interactive protein family (Bolser, et al., 2003; Bolser and Park, 2003), contained domain variations that were directly related to protein translation such as elongation factors Tu/1-alpha/2 and initiation factors IF2/eIF2/eIF5b. Seven protein families (19%) were related to DNA binding proteins. The last five protein families (14%) were related to ATP metabolism (see Supplementary Table 3). Our results corroborate previous studies on well-conserved and minimal gene sets. The functions of protein families constituting a core network are mostly related to protein translation, ribosomal structure, and biogenesis (Aravind, et al., 2000; Koonin, 2000; Mushegian, 1999; Tatusov, et al., 1997).

At the other end of the scale, many protein family interactions appear species or lineage specific (Fig. 2a right-hand side, Fig. 2b left-hand side). This U-shape trend is, however, not common throughout the superkingdoms. Archaea and bacteria show more unique protein family interactions, while eukarya have fewer unique family interactions. Eukarya have a higher ratio of common protein families without many unique interaction pairs (Fig. 2b). To check if this trend is found in the occurrences of protein families, we also plotted protein family numbers without considering their interactions (data not shown). We found the same U-shape distribution for prokarya. This suggests faster evolution rates of prokarya (Nancy, et al., 1995) with more room for new interaction links. This would result in a high ratio of species-specific protein families and their interactions. Also, the eukaryotic era is shorter than that of prokarya, and it is bound to have fewer unique protein family interactions. We suspect that some portion of the unique protein families and their interactions is due to an artifact from incomplete protein structure assignment. When the protein structural assignment ratio for proteomes increases, their portion will decrease gradually to give an accurate number of highly species-specific protein interactions.

Functional coverage of structure assigned proteins

We carried out a test to quantify the coverage of biological functions with the structurally assigned proteins. Using EBI’s GOA-slim (Camon, et al., 2004), a selected set of Gene Ontology (GO) (Harris, et al., 2004), we found that most GO annotations (97% on average) for a complete proteome can be covered by structurally assigned proteins. Although the coverage of our structural assignment is around 50% of whole proteomes, its functional coverage was high enough to deal with overall biological functions. The functional assignment comparison is found in Supplementary Figure 1.

We also carried out a functional analysis of the core protein families. Out of 296,630 proteins, 45,164 belonged to the core protein families, and they covered 83% of GO annotations at the level of EBI’s GOA-slim. In a GO ‘biological process’ mapping, over 91% of the core protein family assigned proteins were associated with central biochemical processes categories such as physiological process (GO:0007582), cellular process (GO:0009987), cellular physiological process (GO:0005875), metabolism (GO:0008152), nucleobase, nucleoside, nucleotide and nucleic acid metabolism (GO:0006139), biosynthesis (GO:0009058), macromolecule metabolism (GO:0043170) and transport (GO:0006810). In the ‘molecular function’ category of GO, over 90% of the proteins were associated with binding (GO:0005488), catalytic activity (GO:0008224), hydrolase activity (GO:0016787), nucleic acid binding (GO:0003676), ligase activity (GO:0016874), transporter activity (GO:0005215), transferase activity (GO:0003676).
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Given the functional coverage, it seems likely that the core protein family network spanning all types of life forms was formed in the very early stage of evolution, occupying the core biochemical processes for life. After the initial formation, a gradual attachment of the interactome seems to have occurred for a long period of time as peripheral functions, such as cell motility (GO:0006928), membrane fusion (GO:0006944), extracellular structure organization and biogenesis (GO:0043062), and pathogenesis (GO:0009405), were needed.

Connectivity of protein family interaction networks

A simple method of characterizing an interactome is to calculate the degree (the total number of connections) of an interacting protein family. In the protein family interactomes, degree \(k\) is the total number of connecting partners at the protein family level (in physics, this quantity is often called "connectivity" and has a different meaning in graph theory) (Dorogovtsev and Mendes, 2002; Fraser, et al., 2002). The degree is the actual number of nearest neighbors of a node (protein family). The degree distribution, the total distribution of degrees of a network, \(P(k)\), has been reported to be of a power-law form: \(P(k) \sim k^{-\gamma}\) in most biological networks (Fraser, et al., 2002; Wagner, 2001).

We found that the \(\gamma\) is 2.57 ± 0.11 for the 15 archaeal species and 2.55 ± 0.11 for the 122 bacterial species, on average. We found the average \(\gamma\) of nine eukaryotic species had a much lower value of 2.08 ± 0.09. This means protein family interactomes are scale-free (Jeong, et al., 2001; Steffen, et al., 2002) in all superkingdoms with a distinction between eukarya and the rest (see Supplementary Table 1). Eukarya have a higher number of hub families (Barabasi and Oltvai, 2004) than those of archaea and bacteria. Also, in eukarya, the hub families have higher numbers of interaction partners, although the criterion for what constitutes a hub family is arguable. The highest degree

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ranges from 24 ~ 38 for eukarya, 11 ~ 17 for bacteria, and 9 ~ 11 for archaea.

Eukarya have more multi-domain proteins than prokarya in general (Apic, et al., 2001). We can explain the eukaryotic protein family interaction networks with many factors including being multi-domained. However, being multi-domained alone can not fully account for this. For example, Pseudomonas syringae (bacteria) and Saccharomyces cerevisiae (eukarya) have similar assignable proteome sizes of 2812 (52% of total) and 2784 (45% of total) and have multi-domain proteins of 752 and 720, respectively; however, they have different \( \gamma \) of 2.67 and 2.03. That is, S. cerevisiae has more hub protein families than P. syringae. Although being multi-domained influences the difference, it does not account for the total architectural difference between the two types.

This high number of interaction partners is not a simple function of proteome size. In the case of Encephalitozoon cuniculi (a eukaryotic parasite protozoan, proteome size: 839 proteins, genome size: 2.9 Mb), the largest hub family has 24 interaction partners, while 80% of other protein families have one or two interaction partners. Although its proteome is smaller than that of E. coli (2338 proteins, 5.2 Mb), the connectivity of the main hub family is much larger than that of E. coli (15 interaction partners). This implies that there is a fundamental difference between prokaryotic and eukaryotic interactomes.

**Network topology and interactome complexity of prokarya and eukarya**

The function of a protein is often affected by its interacting partners. The topology of interaction networks among protein families is determined by 1) the number of protein families, 2) the number of their interactions, and 3) the topology of the interaction link patterns.

Having calculated the number of all the interaction pairs for the 146 species, we found that the number of interaction pairs increases linearly along with the number of protein families without any particular deviation from species to species (Fig. 3a). This indicates that the size of a protein interaction network itself does not determine the topology of the evolving network. Rather, it is closely related to the presence and number of extreme hub families that are capable of continuous growth.

As shown in Figure 3a, increasing one interaction node resulted in one additional interaction edge on average, regardless of the superkingdoms. Figure 3b shows that eukaryotic proteomes have distinctively higher \( \gamma \) values upon increasing the number of interaction pairs. This is due to the different complexity levels they have with a small number of large hub families. Figure 3c shows the degree of interaction for the most highly interacting protein family (G-proteins, c.37.1.8 SCOP family). The number of interacting partners of G-proteins is very high in eukarya. This is a major distinction between eukarya and prokarya, where certain families exploded in the number of interaction partners as the overall interactome size increased. In contrast to the G-proteins, 2Fe-2S ferredoxin domains (d.15.4.2 SCOP family) did not show many extra interaction partners in eukarya, although the interactome size increased (Fig 3d). This is because, while the G-proteins grew rapidly in eukarya, the 2Fe-2S ferredoxin domains remained conservative in incorporating new interacting partners. In other words, these two families could be important components distinguishing eukarya from other superkingdoms. From a functional view, it is plausible that G-proteins have evolved to transduce signals and mediated multicellularity, resulting in a homogeneous environment in eukarya, while 2Fe-2S ferredoxin domains have evolved under the pressure of necessity for diverse metabolism in prokarya.

Specifically, in the comparison of E. coli and S. cerevisiae, the number of interaction pairs is similar in size (1120 and 1097, respectively), even though they belong to different superkingdoms (see Supplementary Table 1). However, the value of \( \gamma \) representing connectivity distribution of the network was 2.58 in E. coli and only 2.03 in S. cerevisiae. This is because yeast has more hub families, genetic hub families are larger despite the similar number of nodes and edges. We suggest that the difference between their interaction network topologies is responsible for their organismal complexity. One mechanism for the higher number of interactions with the same number of interacting families could be compartmentalization within cells. The same kind of structures can have different interacting partners if they are located in different compartments without interfering with other similar molecular interactions. If we regard all the cellular functions as a part of information processing, this could be viewed as an optimization strategy of information processing in S. cerevisiae.

**Expansion of protein family networks**
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As shown above, interactomes can reveal the differences among species in terms of their network topologies. To analyze the evolution of protein family networks, we measured the correlation between the interaction degree of each protein family in each of the 146 predicted interaction networks. The correlation was calculated by Pearson’s correlation coefficient method which set the score 0.5 ~ 1 on positive correlations, -0.49 ~ 0.49 on non-correlations, and -1 ~ -0.5 on negative correlations (see Supplementary Table 4).

We found that only a limited number of protein families have positive correlations (199 out of 1720). Therefore, most of the protein families in a species do not have a positive correlation with the number of total interaction pairs in interactomes. An example for the positive correlations is c.37.1.8 (G-proteins) showed positive correlation especially in eukarya. d.15.4.2 (2Fe-2S ferredoxin domains) showed positive correlation in all three superkinds.

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CONCLUSION

We introduced an analysis protocol that was based on protein family interactions, PSIMAP. Using this protocol, we identified the core network of 47 protein family interaction pairs in all the 146 species. The functions of families constituting the core network are protein translation, ribosomal structure, DNA binding, and ATP metabolism. The results confirmed previous studies that all species share...
the same basic protein families and family interactions critical to cellular functions.

We noted topological characteristics in the interactomes across species: the protein family networks of eukarya had more hub families than archaea and bacteria. This implies an architectural difference between prokaryotic and eukaryotic interactomes.

The number of protein family interaction pairs increased linearly with the number of protein families, regardless of species. In this respect, we suggest that the increase of network size itself does not determine the characteristics of an evolving network. Only a small number of protein families have a very large number of protein family interaction partners, especially in eukarya. We suggest that big hub families continuously increase their number of interaction partners. Therefore, the addition of new protein families to the evolving network is driven not by a random process to all the protein families but by a selective process to special hub families. The recent empirical data (Eriksen and Hornquist, 2002; Jeong, et al., 2001; Newman, 2001) on the dynamics of the attachment of new edges in various growing networks support this mechanism. Interestingly, this indicates that a very small number of special protein families (G-proteins, c.37.1.8 SCOP family for example) play a key role in driving all species diversification, especially in higher organisms.

We suggest a unique approach to detecting an “interaction” core for many species without expensive experiments. As the structural assignment rate using PDB rises, we expect to expand and eventually complete the core. At this stage, our finding confirms the known small core network.

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