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What do we learn from high-throughput protein interaction data?

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The biological significance of protein interactions, their method of generation and reliability is briefly reviewed. Protein interaction networks adopt a scale-free topology that explains their error tolerance or vulnerability, depending on whether hubs or peripheral proteins are attacked. Networks also allow the prediction of protein function from their interaction partners and therefore, the formulation of analytical hypotheses. Comparative network analysis predicts interactions for distantly related species based on conserved interactions, even if sequences are only weakly conserved. Finally, the medical relevance of protein interaction analysis is discussed and the necessity for data integration is emphasized.

Currently, more than 150 bacterial and approximately 15 eukaryotic genomes have been completely sequenced [101]. These sequencing projects provide us with a wealth of information about these organisms. Theoretically, most gene products of these genomes can be predicted from their sequence. Nevertheless, the biochemical activities and biological roles of many gene products remain unclear. Surprisingly, even in new genome sequences approximately one third of the genes cannot be annotated functionally, either because there is no unambiguous homology or homologous genes lack sufficient annotation.

High-throughput functional analysis appears to be the perfect tool to turn the significant number of uncharacterized open reading frames (ORFs) into biological knowledge. Although high-throughput screening (HTS) usually fails to yield a detailed understanding of a protein's function, it often provides the first evidence for function and therefore, an inroute to further characterization.

Currently established HTS methodology includes expression profiling using DNA microarray technology, systematic knockout studies, high-throughput localization studies and protein–protein interaction mapping approaches [1].

This review focuses on protein–protein interaction mapping (interactomics), mainly by two-hybrid approaches. Three questions will be addressed:

- What can we learn from the interaction data generated for several organisms?
- What other information is needed to derive biological conclusions from these data?
- How can such additional data improve our conclusions?

Biological significance of protein–protein interactions

Protein–protein interactions greatly expand the flexibility of proteins beyond their individual activities. For example, the dimeric transcription factors Myc and Max must associate in order to recognize their DNA-binding motif. The Myc/Max dimer allows regulation by altering the concentration of each protein but also by the expression of competitive inhibitors, such as Mad, which binds to and blocks Max. Such combinatorial regulation also expands evolutionary flexibility since each gene's encoded binding partner can duplicate. These additional proteins can adopt different specificities and eventually biological roles. For an extensive discussion of

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protein interactions and their biological significance the reader is referred to standard textbooks of molecular biology [2].

Generation of protein–protein interaction data

Although a number of methods are available for high-throughput analysis of protein–protein interactions, the most commonly used are the yeast two-hybrid (Y2H) system and a combination of protein-complex purification with subsequent analysis by mass spectrometry (MS) [3–5].

The first genome-wide two-hybrid screen was performed by Bartel and coworkers for the study of protein interactions in bacteriophage T7 [6]. The first genome-wide protein–protein interaction studies of a free-living organism have been published by Uetz and coworkers [7] and Ito and collaborators [8] using the yeast *Saccharomyces cerevisiae*. These and other systematic screens have been reviewed by Uetz and Hughes [9].

Soon after these two-hybrid screens, Ho and coworkers [10] and Gavin and colleagues [11] used a large-scale strategy to purify protein complexes from yeast and identified them using MS. Some key differences of the resulting two-hybrid and MS data sets are illustrated in FIGURE 1.

These experimental approaches for high-throughput interaction analyses have already taught us one important lesson: Y2H and MS data sets are strikingly different but are also highly complementary. Interestingly, transient interactions are more often found by Y2H analysis, whereas stable interactions (such

as those in protein complexes) are more reliably identified by *in vivo* pull-down techniques [12]. This finding is not surprising, given the highly cooperative forces that stabilize a protein complex. Many interactions in a complex will not be detected by Y2H analysis, given that the pairs of proteins being tested are not stabilized by the other subunits of a complex.

Recently, the first comprehensive protein–protein interaction maps (PIMs) of flies and worms have been published by Giot and colleagues [13] and Li and coworkers [14]. These studies also used the Y2H system and obtained high confidence maps of approximately 5000 and 2200 unique interactions, respectively.

Protein complex purifications from these organisms have not been carried out successfully on a larger scale, although this may be possible with improved protocols and MS sensitivity.

No matter how they are generated, interaction data have been used by both experimentalists and theorists for further analysis. A breakdown of such uses is shown in FIGURE 2 and discussed below in more detail.

Reliability of high-throughput data

Before conclusions from high-throughput interaction data can be drawn, it is necessary to briefly discuss the quality of available data sets.

No method is able to identify all protein–protein interactions. That is, each experimental strategy generates a significant number of false negatives. The sources of this systematic error are poorly understood. Two-hybrid false negatives may be caused by sterical effects due to the use of two fusion proteins (two-hybrid) or it may involve weak interactions within complexes that require cooperative effects to be stabilized and therefore to generate a two-hybrid signal [12]. Conversely, a major bottleneck for MS analysis are low abundance proteins and proteins that are only weakly associated with protein complexes and hence tend to get lost during purification. False positives are usually a more serious problem since they result in erroneous data and thus misleading conclusions. In Y2H studies, some bait constructs activate the reporter gene without interacting with a prey and so may generate large numbers of technical false positives. On the other hand, biological false positives represent true interactions that take place in the Y2H system but have no biological relevance [15]. A case in point are interacting proteins that are usually expressed in different cell types.

Several approaches were used to minimize the number of false positives in high-throughput studies. Uetz and coworkers [7] discarded Y2H interactions that could not be reproduced, while Ito and collaborators [8] defined interacting protein pairs found three or more times as the (supposedly reliable) core data set.

More elaborate statistical scores were proposed by Rain and coworkers [16] for the *Helicobacter* interaction map and by Bader and colleagues [17] for yeast and other data sets.

Rain and coworkers screened bait proteins against a genomic fragment prey library and considered overlapping prey fragments as the most reliable. This approach combines reproducibility and identifies the interacting domain at the same time.

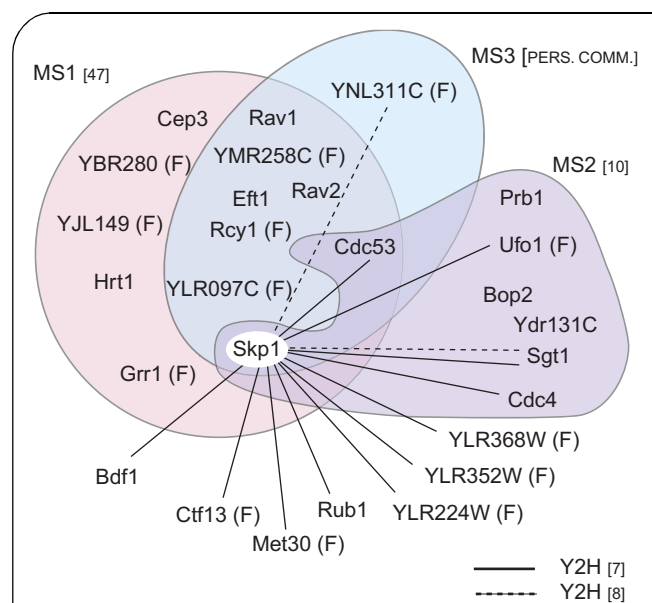


Figure 1. Interaction data gained by Y2H and MS are complementary.

Skp1 is a protein involved in ubiquitin-mediated protein degradation and has been epitope tagged for both Y2H screens and MS analysis. The purified complexes of Skp1 from three independent MS studies and the binary interactions from two Y2H studies are compared. Despite the differences in the data sets, most of the discovered interactions seem to be plausible: most proteins are known to be involved in protein degradation. Skp1 is directed to its target proteins via so-called F-box proteins, which contain a short peptide motif, the F-box (F) [8–10,47, MS3: ELLEDGE S & AEBERSOLD R, PERS. COMM.]. MS: Mass spectrometry; Y2H: Yeast two-hybrid.

The critical point of any attempt to estimate the number of true and false positives in an HTS interaction study is the choice of the true positive data set against which the new interactions are evaluated. Bader and coworkers used the data set of known protein complexes to derive other parameters that allow the scoring of Y2H data [17]. A similar statistical model was applied to the whole *Drosophila* data set resulting in a high confidence protein interaction network, which the authors estimated to retain 40% interactions of biological significance [13].

Edwards and coworkers selected known interactions from 3D structures (RNA polymerase II, proteasome and the Arp2/3 complex) and additional complexes from the literature [18]. The crystal structures of complexes approximate the absolute truth regarding stable protein interactions since they reveal all interactions in atomic detail, at least for the proteins that have been cocrystallized. Based on crystal structures, Edwards and coworkers found a false-negative rate of 51–96% for Y2H and 15–50% for *in vivo* pull-down experiments, respectively. In this context it is remarkable that conventional low-throughput methods also produce a large fraction of false positives, for example, 61% in a pull-down study of RNA polymerase II [18].

Several studies showed that interacting proteins tend to be coexpressed at the messenger RNA (mRNA) level under various experimental conditions [19,20]. However, while coexpression of the two partners increases the confidence in a protein–protein interaction, it is only an indirect measure of its reliability. While proteins in a complex must be expressed at similar levels in order to maintain their stoichiometric ratios, this is not necessarily true for transient interactions that are often found in Y2H screens.

Topology of protein interaction networks

Protein interactions identified on a genome-wide scale are commonly visualized as protein interaction networks. Such networks are graphs with proteins as nodes and interactions as edges (FIGURE 3). Although this representation does not reflect the true nature of protein interactions (which is rather composed of dynamically forming complexes), it serves as a useful mental map and allows for the analysis of certain network properties.

Many biological networks, including protein interaction networks and metabolic networks, have a so-called scale-free topology [21]. Scale-free networks are characterized by a few highly connected nodes (hubs) and many less-well connected peripheral nodes. The distribution of the node degree k follows a power law ($P[k] \sim k^{-\gamma}$) (FIGURE 3) [22,23].

The scale-free nature explains several properties of protein interaction networks. For example, highly connected hubs often appear to have central roles in a network, which would make them vulnerable to attack by mutation or drugs. Jeong and coworkers have shown that the lack of homogeneity of a network results in tolerance to errors [24]. Random mutations in the yeast genome do not appear to affect the overall topology of the network. By contrast, when the most connected proteins are computationally eliminated, the network diameter increases rapidly (i.e., the minimum number of nodes between

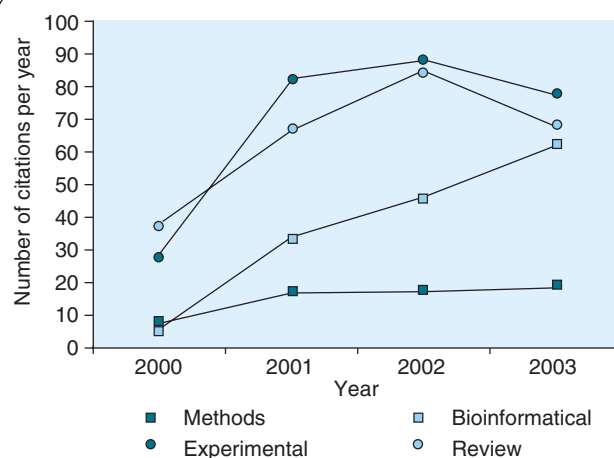


Figure 2. The use of large-scale protein interaction data sets as shown by the number of citations that received during the past 4 years (grouped into four categories) [7]. The high level of citation by experimental (small scale, in depth) studies indicate the usefulness of high-throughput interaction data for more focused analyses. The increasing citation rate by bioinformatics studies, which mainly focus on the high level organization of protein interaction networks, however, illustrates that both a bottom-up, as well as a top-down, view of biological systems are encouraged by these high-throughput studies.

two arbitrary proteins). Although proteins with five or fewer links constitute approximately 93% of the total number of proteins in the data set of Jeong and coworkers, they found that only approximately 21% of them are essential. By contrast, only some 0.7% of the yeast proteins with known phenotypic profiles had more than 15 links, but a deletion of 62% of these proves lethal.

Experimentally derived interaction networks, such as that shown in FIGURE 3B, can be extremely complex and biological meaning is not immediately obvious in them. However, biological systems are hierarchically organized into functional modules and submodules [25]. For example, cells produce ATP via a set of modules, such as the glycolytic pathway, the Krebs cycle and the protein complexes involved in oxidative phosphorylation. Even if their annotation cannot be used for clustering as shown in FIGURE 3C, several groups have developed algorithms to identify functional clusters (cliques) in protein interaction networks. For example, Spirin and Mirny developed an algorithm that was able to recover many previously known protein complexes (e.g., the anaphase-promoting complex) and functional modules (e.g., the yeast pheromone response pathway) [26]. In addition, new complexes (e.g., a complex of six proteins including a YIP1 Golgi membrane protein) and new members of complexes (e.g., two 40S small ribosomal subunits in the Lsm splicing complex) were identified and thus these methods can provide information about single proteins and their biological context.

The interconnections between different modules can be derived from individual protein interactions and their functional annotation (FIGURE 3D). When all proteins of a certain functional class (or module) are collapsed into one node each, the protein interactions can be used to visualize their relationships. For

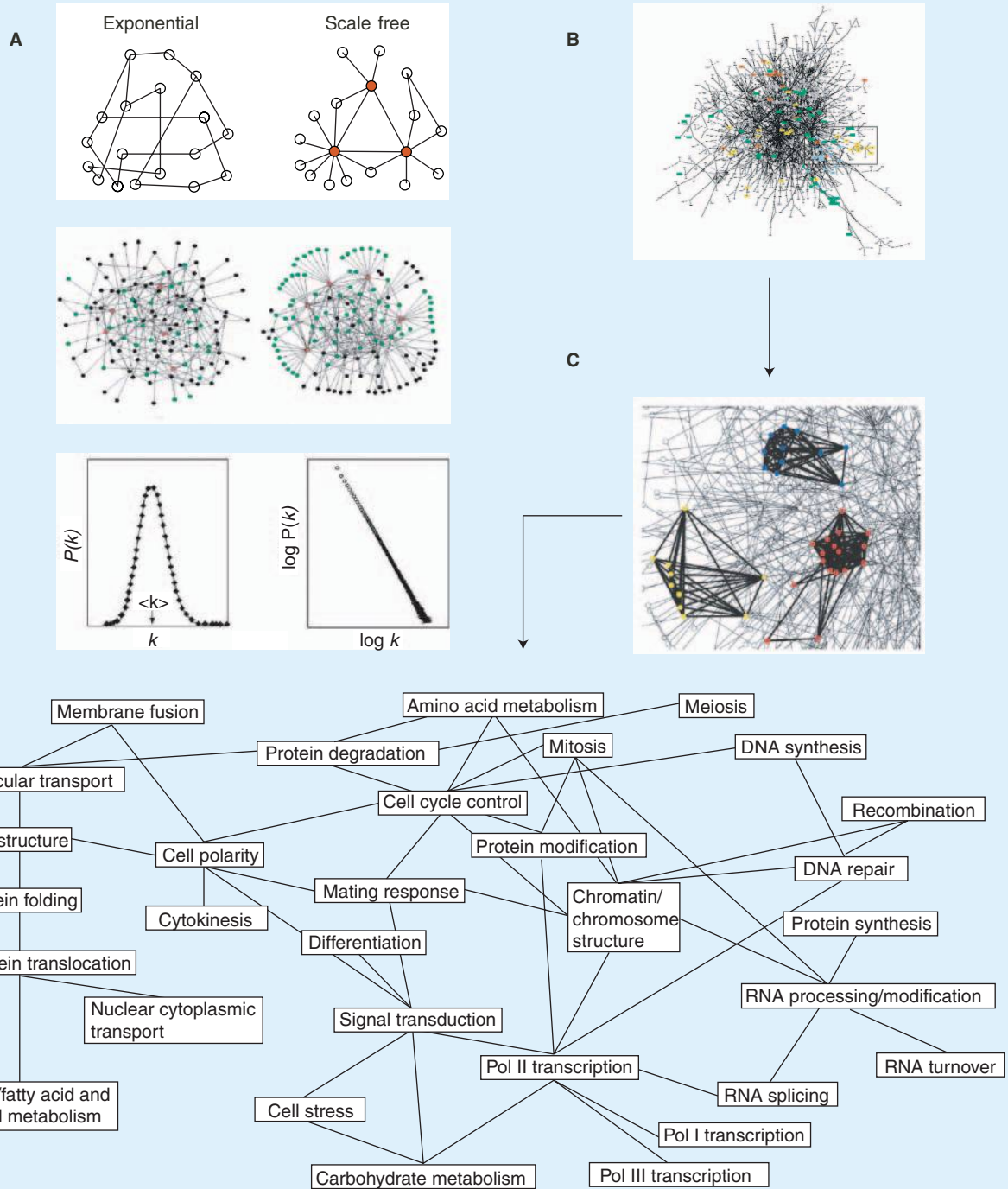


Figure 3. Network classification and analysis. (A) Protein interaction networks are scale-free networks. In contrast to exponential random networks, in which all proteins (nodes) are regarded as equal, scale-free networks have highly connected proteins which are more likely to interact with new proteins added to the network. Exponential networks are therefore statistically homogenous, whereas scale-free networks have a few highly connected proteins (hubs) and many proteins with few interactions. The signature of scale-free networks is the power law distribution of the node degree (k ; number of interacting partners of a protein), $P(k) \sim k^{-\gamma}$, whereas the node degree follows a Poisson distribution in the exponential network model. Reprinted with permission from [23] and [48]. (B) The protein interaction network of yeast reveals different levels of organization. (C) Computer algorithms can deduce molecular modules (protein complexes and pathways) directly from the topology of protein interaction networks [26,49]. (D) Complex protein interaction networks can be collapsed into a meta-network showing the interactions between functional categories. (B) and (D) reprinted with permission from [32].

example, in FIGURE 3D (top middle) the 68 proteins involved in amino acid metabolism are connected by 23 protein interactions. More importantly, this class of proteins also interacts with proteins involved in protein degradation (arguably to generate amino acids), the cell cycle (which controls almost everything and therefore is highly connected by definition) and, surprisingly, chromatin structure. Unexpected interactions such as the one between amino acid metabolism and chromatin structure point to hitherto unnoticed crosstalk between biological pathways and functions which in this case may be regulatory in nature. The fact that some groups (such as the cell cycle proteins) are highly connected indicates their central regulatory role for most other processes in a cell.

Another method for the detection of complexes in protein interaction networks based on k-cores was used to detect a novel nucleolar network in yeast [27,28]. A k-core is a subnetwork of the protein interaction network in which each protein is connected to at least k proteins of this subnetwork. Therefore, this set of proteins forms a highly connected complex in the protein interaction network. The identified nucleolar protein interaction network showed a structure corresponding to the known electron microscopic substructure of the nucleolus (fibrillar component, dense fibrillar component and granular component) [28]. This illustrates that the close examination of protein interaction networks can reveal molecular structures, without *a priori* knowledge of protein functions.

Lessons from single interactions

The ultimate goal of molecular biology is the mechanistic explanation of specific biological phenotypes. For such explanations a detailed understanding of single proteins is necessary. Protein interaction data often provide critical information on the molecular behavior of a protein and almost always allow the formulation of some biological hypothesis. The chromosome cohesin proteins illustrate this point (FIGURE 4): a few interactions of the Smc and Scc proteins in yeast and their predicted coiled-coil structure suggested a model that explained their ability to hold chromatids together.

Obviously, the lower reliability of high-throughput interaction data has to be taken into account and hypothesis building should start with the most plausible interaction and then proceed to less likely ones. However, the power of interaction mapping is also based on the fact that it is not dependent on previous knowledge of a certain protein. Therefore, completely unexpected interactions may lead to spectacular new discoveries. For example, interactions between membrane proteins and transcription factors have usually been considered as false positives. However, during the past couple of years it has been shown in a number of cases that such interactions represent novel methods of regulating transcription directly by membrane receptors. Well-studied examples include the sterol regulatory element-binding proteins [29], Alzheimer protein amyloid precursor protein and the signaling protein Notch [30]. In this manner protein interactions can uncover new connections between previously unlinked processes or pathways. Striking

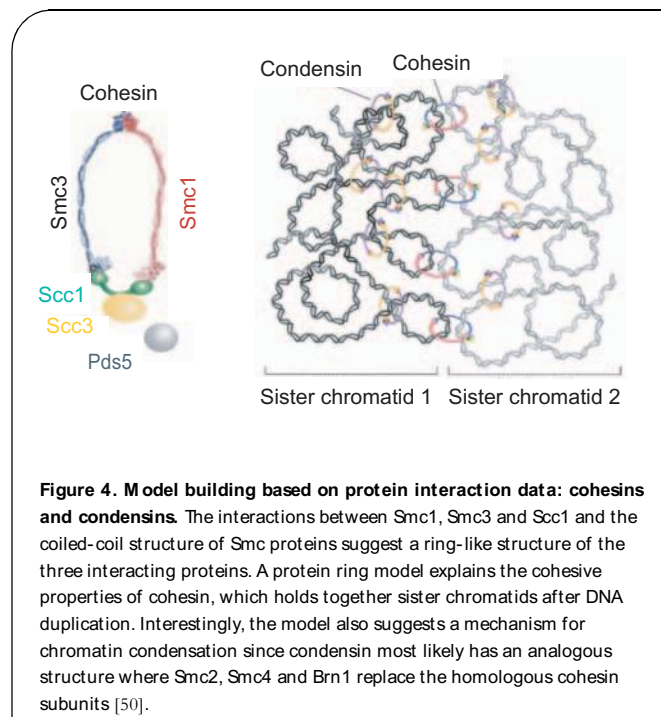


Figure 4. Model building based on protein interaction data: cohesins and condensins. The interactions between Smc1, Smc3 and Scc1 and the coiled-coil structure of Smc proteins suggest a ring-like structure of the three interacting proteins. A protein ring model explains the cohesive properties of cohesin, which holds together sister chromatids after DNA duplication. Interestingly, the model also suggests a mechanism for chromatin condensation since condensin most likely has an analogous structure where Smc2, Smc4 and Brn1 replace the homologous cohesin subunits [50].

examples are moonlighting proteins. These proteins possess multiple functions that are not due to gene fusions, splice variants or multiple proteolytic fragments. Clf1p, for example, is a protein involved in pre-mRNA splicing in yeast. In addition to its interaction with the U5 and U6 subunits of the spliceosome, an interaction with the replication initiation protein Orc2p was shown in a two-hybrid assay. This interaction, together with a DNA replication phenotype, makes Clf1p a protein involved in splicing and in DNA replication initiation and thus represents a link between these putative unrelated processes. More generally, many proteins appear to have several functions. New interactions may suggest such additional functions [31].

An important goal of proteomics is a functional assignment for proteins which cannot be annotated by homology alone. Several approaches for automated functional assignment from protein interaction networks have been developed. The majority rule assignment is based on the observation that 70–80% of the interacting proteins share at least one function, therefore an unclassified protein is assigned the most common function in the set of characterized interacting proteins [32,33]. A disadvantage of this simple method is that interactions between two uncharacterized proteins are not taken into account.

Such predictions have also been experimentally tested. Kemmeren and coworkers verified the predicted function of five proteins that had interactions with known proteins that were also coexpressed [34]. For example, a deletion strain of an uncharacterized ORF (YLR270W) shown to interact with a protein required for thermotolerance (NTH1, neutral trehalase gene) showed sensitivity to heat shock.

Ideally, high-throughput interaction data are used by more traditional cell biological studies (FIGURE 2). For example, Tesse and coworkers examined the role of Ski8p in *Soradia* meiosis [35].

A role of Ski8p in meiotic DNA recombination was suggested by the mutational phenotype. However, due to its known role in cytoplasmic RNA degradation (nonpoly[A] and double-stranded RNA), an indirect role of Ski8p was assumed. However, a direct interaction between Ski8p and a protein involved in meiotic recombination, Spo11, in a comprehensive Y2H study led the authors to examine a direct effect of Ski8p on meiotic recombination which was subsequently proven [7].

Evolution of protein interaction networks

It has been suggested that proteins involved in interactions are more conserved than proteins that participate with a smaller number of interaction partners [36]. However, Jordan and coworkers demonstrated that only proteins with the largest number of interactions (the hubs of the protein interaction network) show a slower evolution rate [37]. Thus, the correlation found by Fraser and colleagues may be an artefact caused by a small subset of proteins rather than a general phenomenon [36].

Comparative interactomics: predicting homologous interactions

Proteins evolve and so do their interactions. If interacting proteins have a weak homology to another pair of interacting proteins, the interaction will support both their functional and evolutionary homology (FIGURE 5A). In order to detect such homologous interactions and pathways, Kelley and coworkers [38] developed the program PathBlast, which aligns two protein–protein interaction networks combining interaction topology and sequence similarity [102]. Using this approach, it was possible to show that the protein–protein interaction networks of yeast and *Helicobacter pylori* harbor a significant number of evolutionarily conserved pathways. A spectacular example among the conserved subnetworks is a group of proteins involved in bacterial membrane transport and nuclear–cytoplasmic transport in yeast. This finding indicates that nuclear–cytoplasmic transport may have originated from a homologous system in bacterial plasma membranes.

Pathway comparison cannot only uncover conserved pathways but can also identify additional components that have been found in one organism but not in another. For example, the homologous proteins shown in FIGURE 5A have different interaction partners. This information can be exploited to predict unknown interaction partners based on homologs in another model. Such predictions are particularly supported by protein complexes that tend to be well conserved, especially as they usually require several conserved subunits for stability.

Integrating protein interaction data with other HTS data

Obviously, high-throughput data are not sufficient to explain complex biological processes. However, it has been demonstrated that the combination of several data sets can contribute significantly to the understanding of certain processes [39]. In addition, high-throughput approaches can also be used to improve data quality and therefore, their predictive power. For

example, it has been shown that the intersection of high-throughput interaction data sets contains more interactions from the same MIPS complex than single data sets [18].

A major drawback of this method is that all high-throughput data sets are far from being comprehensive, which results in a very small intersection between different data sources (e.g., 133 common interactions between Uetz's and Ito's core data sets) [28]. Therefore, a very limited number of interactions are marked as reliable using this method.

A more elaborate approach is the use of a Bayesian network, which allows for the probabilistic combination of multiple data sets. It has been shown that the fraction of false positives and false negatives can be reduced using this method [18]. This approach has also been used in a comprehensive study by Jansen and coworkers [40], in which the high-throughput interaction data sets for the yeast proteome (Y2H and *in vivo* pull-down) were combined with genomic features only weakly associated with an interaction (e.g., coexpression of two proteins) to generate a more reliable interaction data set.

Can a combination of high-throughput data replace traditional experiments?

As has been seen, HTS data are often of lower quality than individually obtained data. On the other hand, HTS data are often better controlled internally since they have been collected under standard conditions. What if all kinds of data were collected under such standardized conditions and were subsequently combined? For example, why are intracellular transport processes not studied by:

- Localizing all proteins in organelles such as the Golgi
- Identifying all protein interactions and complexes
- Measuring their transcription, degradation and post-translational modifications under various conditions
- Their mutant phenotypes

Such data can easily be collected but will not explain any biological mechanism unless experiments that explicitly address defined causal relations are performed. Most importantly, cause and effect cannot be distinguished in advance. For example, deleting all genes in a genome is useful for investigating which proteins are essential, but if a protein is not essential under the tested conditions it will not tell us much. For instance, it is assumed that a protein of previously unknown function (e.g., YHR105W) is involved in vesicular transport since it interacts with other transport proteins in two-hybrid assays. However, one screen of a yeast mutant collection has not found YHR105W as being defective in transport [41]. For further clarification, other hypotheses are needed that reconcile the interaction data and the mutant phenotype. Such hypotheses are often not foreseeable by standardized HTS analysis: the interaction screen was most likely not comprehensive (i.e., there are probably false positives and false negatives) and the mutant screen has only looked at one transport phenotype, namely carboxypeptidase Y

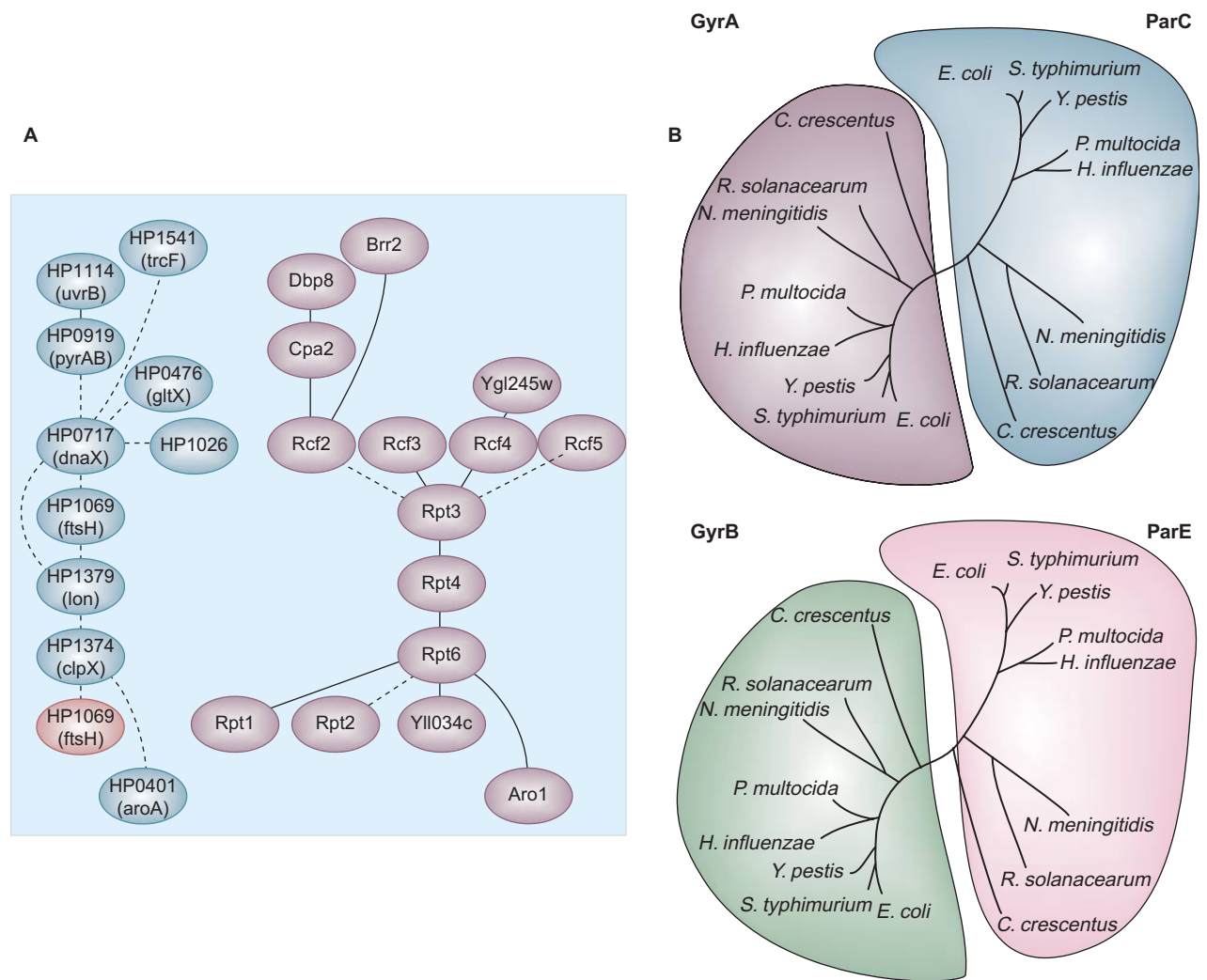


Figure 5. Comparison and evolution of protein interactions. (A) The comparison of protein interaction networks of different species reveals conserved pathways. PathBlast, an algorithm for the alignment of protein interaction networks, was used to identify conserved pathways between *Helicobacter pylori* and yeast [38]. As an example, a protein degradation/DNA replication pathway is shown. Proteins with a certain sequence similarity are placed in one row. Direct protein interactions appear as solid lines and gaps or mismatches are dotted. This pathway alignment demonstrates an association of two pathways which were not previously known to be linked. The network contains proteins associated with DNA polymerase (Rfc2, 3, 4, 6) and subunits of the 19S proteasome regulatory cap (Rpt1, 2, 3, 4, 6) and thereby provides evidence from both yeast and bacteria that the protein degradation and the DNA replication pathways associate *in vivo*. This method can be helpful for predicting protein functions and identifying functional orthologs from among multiple homologous sequences. Furthermore, the comparison of pathways and functional modules helps to understand and visualize protein network evolution [38]. **(B)** Interacting proteins show coevolution. The phylogenetic tree of the GyrA and ParC look strikingly similar to the trees of their interaction partners, GyrB and ParE (i.e., GyrA and GyrB form a complex as do ParC and ParE). Ramani and Marcotte used that similarity to predict interaction partners because the evolution of interacting proteins often shows a similar pattern [51].
C. crescentus: *Caulobacter crescentus*; *E. coli*: *Escherichia coli*; *H. influenzae*: *Haemophilus influenzae*; *N. meningitidis*: *Neisseria meningitidis*;
P. multocida: *Pasteurella multocida*; *R. solanacearum*: *Ralstonia solanacearum*; *S. typhimurium*: *Salmonella typhimurium*; *Y. pestis*: *Yersinia pestis*.

export, which mainly affects Golgi-to-vacuole transport. More subtle effects of YHR105W on protein transport must now be studied, as it is entirely possible that the interaction has a modulatory role in transport as opposed to being absolutely essential. One needs to remember that most mutations are not deleterious but rather show no, or only subtle, defects. This is due to the fact that gene functions can be substituted on the single gene level by duplicate or redundant genes. Such special circumstances usually cannot be identified by HTS and thus have

to be analyzed by a painstaking hypothesis-driven approach, where the hypothesis is refined by each additional experiment.

As an interesting new development, King and coworkers have devised algorithms to automate such hypothesis-driven research [42]. Computer algorithms can replace human reasoning to a certain extent and it may be possible to push HTS to a degree that its experimental conditions can be automatically refined based on previous experiments and therefore, do simulate hypothesis-driven experimentation.

Protein interaction networks for medical research

Most diseases are caused by malfunctioning proteins in one way or another. However, there are only a few known examples of disease-causing defects in protein interactions. The best-studied cases are probably receptors that bind (or do not bind) to peptide hormones or oncoproteins, such as Ras, which may cause cancer when their signaling interactions are defective.

When analyzing mutant proteins it is usually not easy to tell an impaired protein interaction apart from some unrelated effect, such as a folding problem. Hence it is difficult to say if a certain phenotype arises from a defective protein interaction or some indirect cause, such as an instability that prevents a protein from interacting.

For a detailed understanding of disease-causing mutations it would be desirable to have the crystal structures of proteins and their mutants. This would tell us if the structure is really unaffected by a mutant or if the mutant affects an exposed interaction surface.

Interestingly, Giot and colleagues present a human disease protein view in their *Drosophila* PIM, in which proteins with sequence similarity to human disease genes are highlighted (FIGURE 6A) [13]. 74% of human disease genes in the Online Mendelian Inheritance in Man (OMIM) database have strong matches (BLAST e-value $<10^{-10}$) to one or more sequences in the *Drosophila* database [43]. This clearly shows the utility of PIMs in model organisms for medical research.

Using protein interaction networks for drug discovery

The goal of drug discovery is to design or identify small molecular compounds which help to cure or at least ameliorate disease. Protein interaction mapping can be useful at several levels of the drug discovery process. The first step should be the drug target identification. PIMs can help to identify proteins of relevant molecular pathways or complexes which are involved in a specific disease. For example, a highly connected protein (hub) may be a suitable target for an antibiotic whereas a more peripheral protein with few interactions may be more appropriate for a highly specific drug that needs to avoid side effects.

Proteins and their protein–protein interaction surfaces are promising targets for specific drugs, although only a few

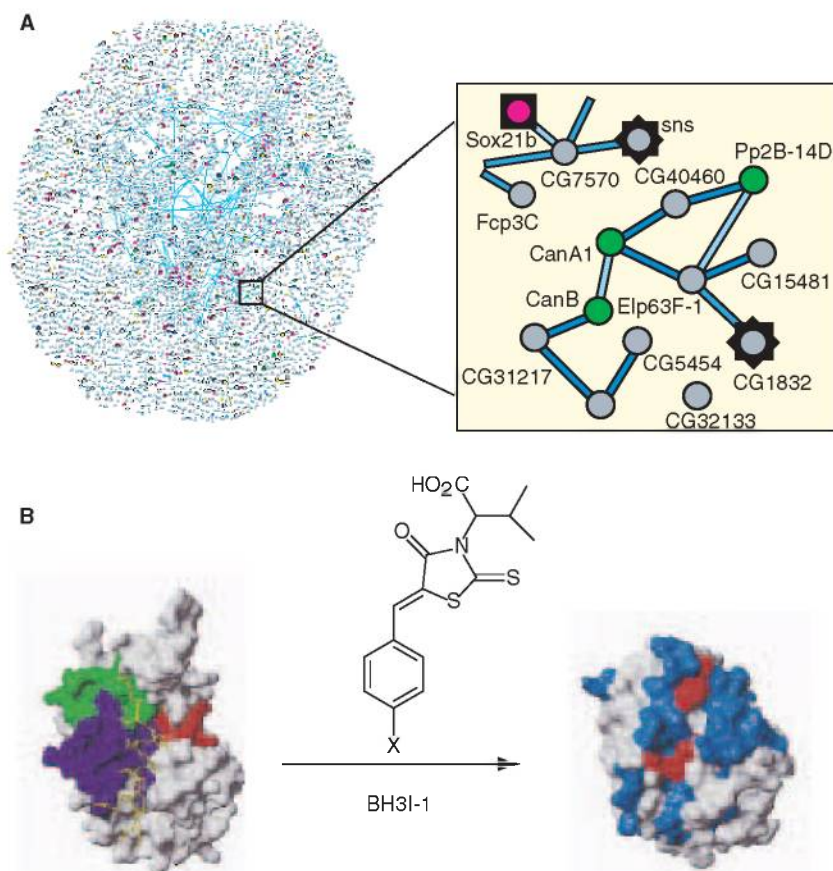


Figure 6. Protein interaction networks can be used for drug discovery. (A) *Drosophila* interaction map with proteins that have a human ortholog and which have been reported to be involved in human disease (shown as stars). Such homologous interactions can help to identify potential drug targets. The insert shows the association of CG1832 with two calcium-dependent phosphatases, CanA1 and Pp2B-14D, which is mediated by the calcium-binding protein Eip63F-1. The homolog of CG1832 (BCL6) is a transcription factor involved in the pathogenesis of human B-cell non-Hodgkin lymphoma. Its association with phosphatases might point to a regulation of this factor akin to nuclear factor of activated T cells, which translocates into the nucleus after dephosphorylation. These phosphatases are potential new targets for the treatment of B-cell lymphomas. Figure adapted with permission from [13]. **(B)** Protein interaction surfaces have been proven to be potential drug targets. Degterev and coworkers [44] designed a compound 5-Benzylidene- α -isopropyl-4-oxo-2-thioxo-3-thiazolidineacetic acid (BH3I-1) which inhibits the interaction of Bak to Bcl-xL (left: Structure of Bcl-xL in complex with the Bak BH3 peptide). It is hypothesized that BH3I-1 binds to residues of Bcl-xL (shaded on the Bcl-xL structure on the right-hand panel) and thereby blocks binding of Bak. Note that the structure of Bcl-xL differs depending on bound compounds (left with Bak peptide, right without bound interactor). Inhibition of Bcl-2 interactions leads to apoptosis. Inhibitors such as BH3I-1 can offer new options for antitumor therapy by sensitizing transformed cells to chemotherapy. Figure adapted and reprinted with permission from [44].

published examples of interaction inhibitors are available. One example are agents which inhibit the interaction between the BH3 domain and Bcl-xL (FIGURE 6B) [44].

Another recently published example is the hepatitis C virus protease, which cleaves the virus-encoded polyprotein. Lamarre and coworkers used interactions of the protease with a substrate to identify short peptides that are recognized by the protease [45]. Starting from a six amino acid peptide which acted as a weak enzyme inhibitor, three amino acid inhibitors were selected. These short peptides could then be used to design a

specific chemical inhibitor of similar structure. The inhibitor had to be designed to enter the cell and appears to have anti-viral activity in preliminary clinical trials. Theoretically this approach can also be applied to other interactions. The limiting problem is to find compounds that mimic peptides and are able to enter epithelia or cells [46].

The diversity of interactions of a targeted protein could also help to estimate or explain side effects of a drug. PIMs indicate immediately which other proteins or processes may be affected by inhibiting a certain interaction. Therefore PIMs can help to design selective agents which target specific interactions of a protein but do not affect others.

Conclusions

High-throughput protein–protein interaction data provide a starting point for the analysis of complexity, signaling and the structural and dynamic organization of cells. In addition, it illuminates an important aspect of the evolution of molecular systems.

If combined with results from other high-throughput methods, such as microarray analysis, a systematic, global view of the molecular functioning of organisms can be gained which for the first time gives us a glimpse of an organism as a whole. By contrast, conventional biological methods are hardly comprehensive, no matter how detailed they are since they always have to focus on certain selected aspects.

Knowledge about biological networks will help us to understand the complexity of biological systems not just as an intellectual achievement. Systems biology will eventually facilitate the simulation and even manipulation of living systems, for example to cure diseases or for the generation of safe and healthy food.

Expert opinion & five-year view

Today, only a limited number of protein interaction studies have been completed. Moreover, the available studies are far from being complete. The most comprehensive data set is

available for the yeast proteome, for which several Y2H and *in vivo* pull-down studies have been published. Protein interaction data sets for other organisms would not only provide insights into the biology of these organisms, but would also tell us about the evolution and general structure of protein interaction networks. In 5 years, many new PIMs for viruses and bacteria as well as other eukaryotes will become available. This will permit the assessment of the diversity of organisms from a systems perspective.

Of course, the protein–protein interaction map of the human proteome is an important goal since this knowledge would promote the understanding of human biology and the therapy of diseases. In 5 years, the human protein interaction map will be far from being complete but there will be several partial interaction maps which elucidate specific pathways and modules, such as those related to human diseases.

While a plethora of data are already available, today's protein interaction networks only give a static view of the molecular organization of the cell. In contrast, the dynamic regulation of protein interactions, for example, in signal transduction cascades, is central to the understanding of biological processes. Small-scale studies have succeeded in analyzing the dynamics of single protein–protein interactions, for example, of the bacterial chemotaxis system. In 2009, the investigation of the dynamics of several biological subsystems (e.g., specific signal transduction cascades) will provide a deeper insight into the complex temporal regulation of the interactome. In addition, new high-throughput techniques which capture these dynamic properties of protein–protein interactions will be available and thus make it possible to initiate projects to understand their dynamics on a proteome-wide scale.

Last but not least, improved databases and visualization tools are urgently needed to make available data more accessible, ideally even to nonspecialists.

Only when we have a clear idea of what is known can we imagine what we do not know.

Key issues

- Protein interaction maps (PIMs) are generated either by the yeast two-hybrid system or by mass spectrometric analysis of protein complexes.
- Both methods produce a certain number of false negatives and false positives. However, the reliability can be improved by combining several data sets.
- Visualization of protein interaction networks as a graph with nodes (proteins) and edges (interactions) reveals the scale-free topology of these networks.
- Biological, meaningful and functional modules can be identified in these networks and interconnections between these modules can be explored.
- A function can be assigned to an unknown protein by examining its binding partners (guilt-by-association approach).
- Protein interaction networks help to identify evolutionary conserved pathways.
- PIMs can be applied in drug discovery to identify target proteins and to minimize side effects.

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