Incorporating high-throughput proteomics experiments into structural biology pipelines

Recent years have seen the emergence of many large-scale proteomics initiatives that have identified thousands of new protein interactions and macromolecular assemblies. However, unfortunately, only a few of the discovered complexes meet the high-quality standards required to be used promptly in structural studies. Consequently, there is an increasing gap between the number of known protein interactions and complexes and those for which a high-resolution three-dimensional structure is available. We have developed and validated a computational strategy to distinguish complexes found in high-throughput affinity purification experiments that will stand the best chances to be successfully expressed, purified and crystallized with little further intervention. Our study suggests that there are some 50 complexes recently discovered in yeast that could readily enter the structural biology pipelines. Indeed, we have used our target selection strategy to draw up a list of 20 complex candidates whose structural determinations will be attempted by groups within 3D Repertoire, a large European Integrated Project that seeks to solve the structures of all amenable protein complexes in yeast at the best possible resolution. The web version of the system is publicly available at http://targetselection.pcb.ub.es.

Contextual specificity in peptide-mediated protein interactions

Protein interactions are central to virtually every major cellular function. While large protein-protein interfaces are typical in tightly associated macromolecular complexes, in most signalling events there is a globular domain in one protein that recognises a linear peptide from another, thereby creating a relatively small interface. These interactions are predominantly found in regulatory networks and, due to their transient nature, are much more difficult to handle biochemically. Recently, large-scale experiments for the determination of peptide recognition profiles of interaction domains, and derivation of the corresponding patterns, have been developed, although transient peptide-mediated interactions are still underrepresented in high-throughput experiments. Although binding is mediated by a small number of contacts formed by the residues in linear motifs, this type of interaction is extremely specific in vivo. For instance, it has been shown that the Pbs2 peptide is recognised only by the SH3 domain of Sho1 (its biological partner) and does not cross-react with any of the other 26 SH3 domains in yeast, although interactions with SH3 domains from other species are biophysically possible. More recently, another study has also shown that the binding specificity of PDZ domains is optimised across the 157 domains contained in the mouse proteome. However,
bonds created between residues in linear motifs and globular domains, while sufficient to ensure binding, are too few to explain the high degree of specificity observed \textit{in vivo}. It is thus, as occurs in phosphorylation events, the biological context that will ultimately determine the interaction specificity. This context has several aspects; certain subcellular localisation or expression patterns will determine whether proteins that are potential competitors for an interaction \textit{in vitro} actually meet \textit{in vivo} and thus evolve into niches of molecular recognition that allow them to bind only the desired target domain. Nevertheless, even within a cellular compartment several interaction domains and their complementary ligands are regularly expressed simultaneously, so yet more contextual information is required to achieve the observed specificity. This information is, to a great extent, contained in the residues surrounding the motif.

In the lab, we have systematically identified all instances of peptide-mediated protein interactions of known 3D structure and used them to study the individual contribution of motif and context to the global binding energy. We found that, on average, the context is responsible for roughly 20\% of the binding and plays a crucial role in determining interaction specificity, by either improving the affinity with the native partner or impeding non-native interactions. We also have examined and quantified the topological and energetic variability of interaction interfaces, and have found a much higher heterogeneity in the context residues than in the consensus binding motifs (Figure 1). Our analysis partially revealed the molecular mechanisms responsible for the dynamic nature of peptide-mediated interactions, and suggested a global evolutionary mechanism to maximize the binding specificity. Finally, we have investigated the viability of non-native interactions and highlight cases of potential cross-reaction that might compensate for individual protein failures and establish backup circuits to increase the robustness of cell networks.

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\textbf{Figure 1.} Peptide exchange results for human SH3 domains. Upper heat maps show the topological distortion (left) and the energy variations (right) of all artificial (ie, non-native) interaction pairs constructed between human SH3 domains and their ligand proteins, with respect to the native topologies and the average native binding energy. SH3-binding peptides 1-9 correspond to class I and 10-16 to Class II. The lower figure shows the energy variation of motif and context, compared to the native binding energies, for each individual residue in the native (n) and artificial (a) interactions.
Exploiting gene deletion fitness effects to understand the modular architecture of protein complexes under different growth conditions

An understanding of how individual genes contribute towards the fitness of an organism is a fundamental issue in biology. Although recent genome-wide screens have generated abundant data on quantitative fitness for single gene knock-outs, very few studies have systematically integrated other types of biological information to study how and why deletion of specific genes gives rise to a particular fitness effect. In a recent study, we combined quantitative fitness data for single gene knock-outs in yeast with large-scale interaction discovery experiments to examine the effect of gene deletion on the modular architecture of protein complexes, under a range of growth conditions. Our analysis revealed that genes in complexes show more severe fitness effects upon deletion than other genes (Figure 2). However, in contrast to what has been observed in binary protein-protein interaction networks, we found that this was not related to the number of complexes in which they are present. We also observed that, in general, the modular components of protein complexes (ie, core and attachment proteins) are equally relevant for the complex machinery to function. However, when quantifying the importance of core and attachments in single complex variations, or isoforms, we observed that this global trend originates from a combination of apparently unrelated factors, thereby indicating the presence of distinct fitness patterns in a single complex across growth conditions. Finally, our study also highlighted several interesting cases of potential functional compensation between protein paralogs and, perhaps, a new piece to fit in the histone-code puzzle.

Towards a molecular characterisation of pathological pathways

In the last century, biomedical sciences were clearly immersed in a conceptual reductionism induced by the success of molecular biology. The development of methods to isolate and study individual cells and molecules has significantly increased our understanding of the nature of life and has led to considerable social advances, including the development of new medicines. Recent years have witnessed how the many genome sequencing projects have provided nearly complete lists of the macromolecules present in an organism, including humans. However, biological systems are often complex in nature, and the knowledge of the components reveals relatively little about their function and organisation. The scientific community is now aware of the difficulties of predicting the behaviour of an intact organism from the individual actions of its parts in isolation and is rapidly moving to systems approaches, where global properties are also considered. In fact, most follow-up initiatives to the sequencing projects have been directed towards solving the systems’ complexity and have focused on unveiling the millions of interrelationships between macromolecules in an organism or monitoring how they coordinately change in response to a particular stimulus (ie, disease). Indeed, functional genomics initiatives are already delivering the first drafts of whole organism interactomes, gene expression profiles for many tissues and conditions and the initial quantifications of metabolites in humans.

Figure 2. Comparison of the fitness of yeast strains upon deletion of genes in complexes and other genes. Distributions of strain fitness upon deletion of genes in complexes (red) and genes not part of complexes (blue) in two fermentable and three non-fermentable media. Genes with a fitness of zero are essential. The fitness values of individual genes are partitioned into four categories: ‘strong negative effect’ (−), ‘moderate negative effect’ (−), ‘weak or no effect’ (0) and ‘positive effect’ (+).
Pharmacological sciences have gone through a similar process, with traditional approaches being mostly reduced to the study, at the molecular level, of the target-compound duet. However, the truth is that phenotypic observations (i.e., disease symptoms) are often the result of an incredibly complex combination of molecular events. This is because virtually every major biological process is not performed by a single molecule but by large macromolecular assemblies and is often regulated through a complex network of transient interactions. Moreover, since most pathways are interconnected, slight changes in these transient regulatory networks can trigger one process or another, with completely different outcomes.

This reductionism has had striking consequences, for instance many promising drug candidates have failed the last, and most expensive, clinical phases because of the poorly understood action mechanisms of the pathways they target or an inappropriate choice of the animal models, which proved ineffective at predicting off-target effects. It is therefore clear that to increase the revenues of drug discovery, we need to improve our knowledge of the molecular mechanisms of disease by considering the full biological context of a drug target and moving beyond individual genes and proteins.

Probably, the main goal of our laboratory can be considered the global molecular characterisation of pathological pathways through a combination of computational biology and interaction discovery techniques, in a real dry-wet cycle, where we use computational modelling to design the experiments required to complement and complete the initial models (Figure 3). To this end, the recent creation of the Experimental Bioinformatics Lab, a joint initiative between IRB Barcelona and the Barcelona Supercomputing Center (BSC), has been crucial. The power of our approach is that we start from in silico modelling and therefore we are not restricted to the study of one or a few patho-physiological pathways. On the contrary, the first steps will involve a global modelling of all the human routes that might arise from known data, and this process will reveal novel and unexpected connections between them. We will then choose to further study those of most relevance from an academic or clinical perspective. We have already started to implement our approach to study the molecular bases of aging-related diseases and metabolic syndrome in collaboration with academic and industrial partners.

It is our belief that a deeper understanding of the molecular mechanisms underlying a disease phenotype will permit the discovery of new potential targets, lead to more effective combinations of already marketed products, and help to select the best model organisms to study a patho-physiological pathway or to identify specific biomarkers.
Publications

Research Networks and Grants
*Aproximación bioinformática al estudio de la especificidad contextual en redes de interacciones entre proteínas y sus posibles aplicaciones biomédicas y biotecnológicas*
Ministerio de Educación y Ciencia, BIO2007-62426: 2007-2010
Research Director: Patrick Aloy

*Identificación de dianas secundarias y diseño de fármacos para enfermedades relacionadas con el envejecimiento mediante el análisis estructural y funcional de sus rutas biológicas*
Research Director: Patrick Aloy

Collaborations
*Identification of potential phosphorylation targets for AURORA A kinase in human*
Isabelle Vernos, Centre for Genomic Regulation (Barcelona, Spain)

*Modular architecture of protein complexes and gene deletion fitness in yeast*
Madan Babu Mohan, MRC-LMB (Cambridge, UK)

*Molecular characterisation of biological pathways related to aging*
José Manuel Mas, InfoCiencia Clinical Research (Barcelona, Spain), Xavier Gomis-Rueth, IBMB-CSIC (Barcelona, Spain), Baldo Oliva, GRIB-UPF (Barcelona, Spain) and Xavier Daura, IBB-UAB (Barcelona, Spain)

*Molecular characterisation of biological pathways related to metabolic syndrome, type II diabetes and obesity*
José Manuel Mas, InfoCiencia Clinical Research (Barcelona, Spain), Andrés Fernández, Salvat Biotech (Barcelona, Spain), Albert Barberà and Ramón Gomis, IDIBAPS-Hospital Clinic de Barcelona (Barcelona, Spain)

*Structural characterisation of macromolecular machines in yeast*
Luis Serrano, Centre for Genomic Regulation (Barcelona, Spain) and Rob Russell (European Molecular Biology Laboratory, Heidelberg, Germany)