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Constraints imposed by non-functional protein–protein interactions on gene expression and proteome size

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Crowded intracellular environments present a challenge for proteins to form functional specific complexes while reducing non-functional interactions with promiscuous non-functional partners. Here we show how the need to minimize the waste of resources to non-functional interactions limits the proteome diversity and the average concentration of co-expressed and co-localized proteins. Using the results of high-throughput Yeast 2-Hybrid experiments, we estimate the characteristic strength of non-functional protein–protein interactions. By combining these data with the strengths of specific interactions, we assess the fraction of time proteins spend tied up in non-functional interactions as a function of their overall concentration. This allows us to sketch the phase diagram for baker's yeast cells using the experimentally measured concentrations and subcellular localization of their proteins. The positions of yeast compartments on the phase diagram are consistent with our hypothesis that the yeast proteome has evolved to operate closely to the upper limit of its size, whereas keeping individual protein concentrations sufficiently low to reduce non-functional interactions. These findings have implication for conceptual understanding of intracellular compartmentalization, multicellularity and differentiation.

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Introduction

The properties of individual proteins ranging from their structure and folding to function and evolution have been extensively studied. Recently, protein-protein interactions (PPIs) became popular subjects of both experimental (Uetz et al, 2000; Ito et al, 2001; Giot et al, 2003; Li et al, 2004; Stelzl et al, 2005; Rual et al, 2005; Gavin et al, 2006) and theoretical scrutiny (Bogan and Thorn, 1998; Lo Conte et al, 1999; Kortemme and Baker, 2002; Aloy et al, 2004; Aloy and Russell, 2006; Kim et al, 2006). In addition to the traditional studies, which mainly focus on the detailed understanding of protein interactions within small sets of functionally related proteins, the new systems biology perspectives have emerged (Sear, 2004; Deeds et al, 2007; Maslov and Ispolatov, 2007), which address the behaviour of large sets of proteins of diverse types operating together inside living cells. However, our understanding of the coexistence of different proteins in crowded cellular environments remains quite limited.

The key issue we discuss here is whether the possibility to form numerous non-functional interactions (Box 1) in such environments impedes proteins' ability to engage in highly specific biologically functional interactions. Here, the non-functional interaction refers to a transient complex between two functionally unrelated proteins. Such complexes are formed following random encounters between proteins as they diffuse in the cell searching for their functional (specific) partners. Apparently, one way for nature to reduce non-functional interactions is to keep the concentrations of individual proteins in a cell to a minimum. However, there is a limit to how low this concentration can go, as it must allow for the efficient biological functioning of the cell as well as formation of the specific complexes (both transient and permanent). Obviously, under these conditions, the only remaining way to reduce the overall concentration of proteins and hence the formation of non-functional complexes is to limit the diversity of the proteome, that is, the number of different protein types that are co-expressed in a given subcellular

Box 1 A non-functional interaction refers to formation of a transient complex between two functionally unrelated proteins. Such complexes are formed following random encounters between proteins as they diffuse in the cell searching for their functional (specific) partners. If there are many types of coexisting proteins in a cell compartment, the chance to encounter an arbitrary partner in such random walk is much greater than to encounter a protein's specific partner. For functional interactions to dominate, this entropic factor has to be compensated by making functional interactions, on average, much stronger than random non-functional ones. This is achieved by evolutionary selection of interacting surfaces for functionally interacting proteins. Proteins expressed at lower concentrations are subject to stronger evolutionary pressure, as they need a smaller dissociation constant to form specific complexes. To enhance their strength and specificity, functional interactions may involve the formation of additional hydrogen bonds and, in some cases, salt bridges. On the other hand, non-functional interactions, which have not been evolutionarily optimized, result from random encounters between surfaces of two proteins. The physical nature of these interactions is hydrophobic: it is mainly due to burial of hydrophobic groups on the transient interface from water.

compartment. If such limit on the proteome diversity does indeed exist, how far below it are the environments inside real cells? For instance, among the ~4500 yeast protein types simultaneously expressed under the normal conditions (Ghaemmaghami *et al*, 2003; Huh *et al*, 2003), about 1800 types are known to be colocalized in the cytoplasm (Huh *et al*, 2003). How far is this number from the theoretical upper bound?

A related question is to what extent non-functional interactions affect the biochemical efficiency of formation of functional protein complexes? That is to say, by how much proteins' search for their specific partners is slowed down by transient non-functional complexes? The answer depends upon both the proteome diversity and individual protein concentrations. Were individual protein concentrations evolutionarily tuned to assure the sufficient efficiency to form specific protein complexes?

We hypothesize that in general the biological evolution pushes an organism towards higher protein diversity. The diversity increases until it reaches the limit beyond which nonfunctional interactions start to significantly interfere with proteins' functions. At the same time, the average concentration of individual proteins is pushed down in evolution to reduce the impairment of non-functional interactions to biochemical efficiencies, until it reaches the lower limit defined by stability of specific complexes as well as the overall functioning of the cell.

In this study, we quantitatively address these questions and check this hypothesis by comparing the equilibrium concentrations of specific and non-functional complexes and that of unbound proteins in the monomeric form. The typical binding energy of specific PPIs is obtained from a database (Kumar and Gromiha, 2006) generated by manual curation of the literature. As the information about non-functional PPIs is not usually reported in the literature, we infer the distribution of their binding energies from the data generated in high-throughput Yeast 2-Hybrid (Y2H) experiments (Uetz et al, 2000; Ito et al, 2001; Giot et al, 2003; Li et al, 2004; Rual et al, 2005; Stelzl et al, 2005). As presented in Box 2, the PPIs detected in these experiments are known to include many false-positives (Ito et al, 2001; Aloy and Russell, 2002; Deane et al, 2002; Vidalain et al, 2004; Huang et al, 2007), which are likely caused by nonfunctional interactions (Deeds et al, 2006). Therefore, the

Box 2 Interactions between many proteins are systematically tested in large-scale Y2H screens. In Y2H approach, proteins are overexpressed to high concentration of 1 µM or higher (Estojak et al, 1995). To this end, the Y2H approach detects PPI with binding affinities stronger than $K_{d}\!\approx\!1\,\mu\text{M}.$ Some reported interactions (up to 19%; Nakayama et al, 2002) may not be real, but owing to experimental artefacts such as bait self-activation (Vidalain et al, 2004). However, the remaining of the reported interactions, although being real in Y2H assay, may not all be observed in living cells. Many proteins are expressed in vivo at lower concentrations than baits and preys in Y2H experiments, and therefore they will not form complexes in cells, although in Y2H assay they can form complexes due to overexpression. Pairs of proteins with binding affinities close to K_d^* form transient complexes in Y2H, which may be detected in some experiments and missed in others, hence lack of reproducibility for weaker complexes. Although these limitations may affect the ability of Y2H to reliably detect functional PPIs, the Y2H data is very useful to estimate strength of non-functional PPI because it massively reports on relatively weak interactions between proteins. Here we use the Y2H data for that purpose, assuming that fraction α of PPIs are real yet non-functional, which are detected under high concentration conditions of the Y2H experiment. The uncertainty in the estimate of parameter α gives rise to the uncertainty in the estimate of m_{dead} —the maximal number of protein types that can be coexpressed in a cell. In the future, to obtain the complete set of non-functional PPIs within a given interactome, it will be necessary to systematically disrupt each interaction and test the impact of such a disruption on biological functioning of the cell. Eventually, when large-scale experimental information about the binding affinities of individual PPIs will become available, one would be able to exactly calculate the effect of nonfunctional interactions on biochemical efficiency of individual proteins.

results of proteome-wide Y2H assays provide a unique opportunity to estimate both the average strength and the distribution of non-functional interactions.

Using these estimates, we find that the size of the yeast proteome is close to the theoretical upper limit imposed by competition between specific and non-functional interactions. Namely, the number of protein types co-expressed and colocalized in the cytoplasm, which is the largest compartment of the yeast cell, is close to its theoretical upper limit. It could be further increased only by the virtue of intracellular compartmentalization or (in multicellular organisms) by celltype differentiation. At the same time, we find that the average concentration of individual proteins expressed in the yeast cytoplasm is close to its theoretical lower limit at which functional interactions with a typical specific binding energy are possible. On average, the time proteins waste in various non-functional complexes is typically comparable with the time they spend in the monomeric (unbound) form. However, for some of the stickiest proteins, this lost time ratio can be significantly higher. We conjecture that the impact of nonfunctional PPIs on biochemical efficiencies of specific protein complexes is close to the tolerable limit. Other relatively large compartments of the yeast cell, the nucleus and mitochondria, have fewer protein types than the cytoplasm and higher average individual protein concentrations, although these differences do not exceed an order of magnitude.

Results

The upper limit on protein diversity: a simple estimate

Non-functional interactions are ubiquitous in crowded cellular environments. As a large variety of different proteins are forced to coexist in the same intracellular compartment, a protein encounters non-functional interaction partners much more frequently than the few protein types with which it has evolved to specifically interact. As was pointed out by Janin (1996) and Deeds et al. (2007), the specific interactions must be significantly stronger than non-functional ones to guarantee that specific complexes are more common than random, non-functional complexes. We begin the discussion of the role of non-functional interactions from a simplified conceptual example where we assume that: (1) all binding energies E_s in a small set (one interaction partner per protein) of specific PPIs are the same; (2) the binding energies E_n of non-functional interactions between all possible pairs of proteins are also identical to each other and (3) the concentrations C_i of every protein in the cell are the same. Furthermore, we consider the case in which the biologically functional state of each protein is when it is bound with its specific interaction partner. Although, as will be shown below, these assumptions are certainly an oversimplification, they represent a conceptually useful first step towards addressing the problem of the competition between specific and non-functional complexes.

Consider a protein of a certain type *i* that interacts with other proteins of a unique type *i'* through specific binding, and with all other proteins *R* through non-functional binding. We assume that proteins participating in pairwise specific and non-functional complexes do not bind to other proteins. Then the total concentration $C_i = \tilde{C}$ of the protein *i* is formed by

$$\tilde{C} = [i] + [ii'] + [iR]$$
 (1)

where [i], [ii'] and [iR] are the concentrations of the three states of the protein *i*: the monomeric (unbound), bound in a specific complex and bound in any of the promiscuous non-functional complexes. These three concentrations are related to each other by the law of mass action

$$[iR] = \frac{[i][R]}{C_0 \exp(E_n/kT)}$$
(2a)

$$[ii'] = \frac{[i][i']}{C_0 \exp(E_s/kT)}$$
(2b)

Here the normalization concentration $C_0=1M$ is the convention and $[R] = \sum_{j=1}^{m} [j]$ sums over nearly all of the *m* protein types co-localized in the same compartment. The binding energies are related to the dissociation constants by $K=C_0 \exp(E/kT)$ and are usually negative. Among the three terms in Equation (1), the concentration [ii'] should be the dominant one to ensure the abundance of specific complexes needed for proper functioning of the cell. We refer to a situation in which [ii'] is not the largest of the three terms as a biologically prohibitive 'dead zone' shown in red colour in Figure 1. One way, the system can fall into the dead zone if [ii'] < [i]. For a simple estimate of the boundary of this zone, let us ignore for a moment the contribution of non-functional interactions. In this case, precisely at the border of the dead zone, one has $[ii'] = [i] = [i'] = \tilde{C}/2$. Thus, according to Equation (2b), the system is in the dead zone if the individual protein concentration \tilde{C} is below the critical value

$$\tilde{C} < C_{\text{crit}} = 2K_s = 2C_0 \exp(E_s/kT) \tag{3}$$

We emphasize that the choice of [ii']=[i] to define the boundary of the dead zone is purely on the basis of



Figure 1 The conceptual illustration of possible states of a protein in a compartment of a living cell. The green, yellow and red regions represent the 'safe zone', the 'dangerous zone' and the 'dead zone', respectively. The boundaries between them are given by Equations (3–5).

convenience. In reality, the ratio [ii']/[i] changes continuously with \tilde{C} . Therefore, the dead zone is separated from the other zones by a smooth crossover rather than a sharp phase transition. The dead zone also includes the situation when [ii'] < [iR]. By comparing Equations (2a) and (2b), we find that this happens when the number m of co-expressed and co-localized protein types exceeds the critical value

$$m_{\text{dead}} = [R]/[i] = \exp[(E_n - E_s)/kT]$$
(4)

Similar to Equation (3), m_{dead} is a characteristic value of the crossover. The inequality $m_{\text{dead}} \ge 1$ follows from the observation (Janin, 1996; Deeds *et al*, 2007) that specific, functional, interactions are significantly stronger, statistically, than nonfunctional ones. We further divide the biologically allowed region into two parts: the 'safe zone' with [iR] < [i] < [ii'] (shown in green in Figure 1), and the 'dangerous zone' with [i] < [iR] < [ii'] (shown in yellow in Figure 1). The boundary between them in this simplified example is determined by Equation (2a) to be:

$$m_{\text{warning}} \approx K_n / (\tilde{C}/2) = (2C_0/\tilde{C}) \exp(E_n/kT)$$
 (5)

where K_n is the dissociation constant for non-functional interactions. If one fixes the concentration \tilde{C} of each individual protein type and increases the number of protein types m, starting from the safe zone, one first enters the dangerous zone at $m=m_{\text{warning}}$, and finally reaches the dead zone at $m=m_{\text{dead}}$. As \tilde{C} decreases, m_{warning} increases, and the separation between m_{warning} and m_{dead} shrinks. Eventually, when \tilde{C} reaches the lowest value C_{crit} given by Equation (3), the separation between m_{warning} and m_{dead} disappears altogether. In other words, the three lines corresponding to Equations (3)–(5) intersect approximately at the same 'triple point' (see Figure 1), where all three terms in Equation (1) are equal to each other.

A protein in the dangerous zone has disadvantages compared with those in the safe zone. First, to ensure the dominance of the specific complex, [ii'] > [iR], its specific interaction must be strong enough to resist the interference from non-functional interactions (see Supporting Information for details). Although this could be achieved in evolution, it is an extra requirement the cell needs to satisfy. Moreover, even if the specific interaction is strong enough, a protein in the

dangerous zone would waste a considerable amount of time on non-functional interactions with non-functional partners. This increases the time it takes for it to find its specific partner by the factor 1 + [iR]/[i] > 2 and thus affects the temporal efficiency of all biochemical processes involving this protein. Therefore, it indeed represents a 'dangerous', or suboptimal, situation and imposes a soft upper limit on the protein diversity. Outside of the dead zone, compartments of living cells need to minimize the average losses of biochemical efficiency experienced by proteins inside the dangerous zone.

Physical model of non-functional interactions

What happens inside intracellular compartments of real cells is more complicated than the simplified example presented above. Most importantly, the binding energies of both specific and non-functional interactions as well as the concentrations of individual proteins vary in a broad range. Therefore, different proteins in a cell are characterized by different 'danger levels' [*iR*]/[*i*]. In spite of these complications, can we still quantitatively analyse the general situation inside compartments of real cells, for example, in the cytoplasm of yeast cells under normal conditions? How many protein types fall inside the dangerous zone? What is the largest value of [*iR*]/[*i*] among all protein types present in the yeast cytoplasm? What is the phase diagram in terms of the proteome diversity m and the distribution of individual protein concentrations? To address these questions, we start with a basic physical model of non-functional interactions.

To enhance their strength, functional interactions may involve the formation of additional hydrogen bonds and, in some cases, salt bridges. On the other hand, non-functional interactions, which have not been evolutionarily selected, are formed primarily due to the hydrophobic effect. Indeed, nonfunctional interactions result from random encounters between surfaces of two proteins, and the free energy gain associated with such an encounter was shown to be proportional to the total amount of buried hydrophobic residues on these two surfaces (Janin, 1995; Bahadur et al, 2004). It has become standard to relate the binding (free) energy to surface hydrophobicity (Eisenberg and McLachlan, 1986; Noskov and Lim, 2001; Bahadur et al, 2004; Deeds et al, 2006). Here, we adopt a simple model from Deeds et al (2006) to describe the distribution of non-functional interactions. The major premise of the model is that the free energy of a non-functional PPI is determined by the total hydrophobic surface that is screened from the water upon the formation of a complex. This assumption is strongly supported by the structural and energetic analysis of many functional and non-functional protein complexes (Bahadur et al, 2004). The interaction energy between any two proteins *i* and *j* is therefore additive:

$$E_{ij} = E_i + E_j + \Delta G_{(0)} \tag{6}$$

Here, E_i and E_j describe the 'stickiness' of proteins *i* and *j*, indicating how hydrophobic their surfaces are. In support of this model, Figure 3 of Janin (1995) showed that experimental result of binding energies can be described by the sum of stickiness terms and a constant term. The constant term $\Delta G_{(0)}$, determined as ~6 kcal/mol ≈ 10kT from experiments (Janin, 1995; Tamura and Privalov, 1997), comes from the transla-



Figure 2 Gaussian distributions: (**A**) $f_1(E_i)$ of 'stickiness' parameters E_i of individual proteins (**B**) $f_2(E_{ij})$ of binding energies $E_{ij}=E_i+E_j+\Delta G_{(0)}$ of pairwise non-functional interactions. The dashed line in the panel (A) shows the fraction of proteins in the monomer defined by Equation (10) with the chemical potential μ . The shadowed area is the product of $f_1(E_i)$ and Equation (10), and represents the distribution of E_i in protein monomers. The shadowed region in panel (B) corresponds to non-functional PPIs that are stronger than the detection threshold E^* and thus appear in high-throughput Y2H screens. The fraction of such interactions among $\sim N^2/2$ pairs is given by Equation (16).

tional and rotational entropy the two proteins lose while binding to each other. In other words, the effective 'interaction volume' is much smaller than the convention of $1/C_0=1M^{-1}$ in Equation (2). The fraction *P* of hydrophobic residues on proteins' surfaces has an approximately Gaussian distribution, with the average $\vec{P} \cong 0.22$ and the standard deviation $\sigma_P \cong 0.058$ (Deeds *et al*, 2006). As E_i is proportional to the surface hydrophobicity, $E_i/E_j=P_i/P_j$, the distribution of E_i is also expected to be Gaussian (Figure 2a)

$$f_1(E_i) = \frac{1}{\sqrt{2\pi\sigma^2}} \exp\left[-\frac{(E_i - \bar{E})^2}{2\sigma^2}\right]$$
 (7)

where

$$\bar{E}/\sigma = -\bar{P}/\sigma_P = -0.22/0.058 = -3.8$$
 (8)

(Deeds *et al*, 2006). The distribution of binding energy $E_{ij}=E_i+E_j+\Delta G_{(0)}$ of two-body non-functional interactions is then given by

$$f_2(E_{ij}) = \frac{1}{\sqrt{4\pi\sigma^2}} \exp\left[-\frac{(E_{ij} - 2\bar{E} - \Delta G_{(0)})^2}{4\sigma^2}\right]$$
(9)

with the average value $2\bar{E} + \Delta G_{(0)}$ and the standard deviation $\sqrt{2}\sigma$ (Figure 2b). This model has been used to successfully explain most global topological properties of PPI network (Deeds *et al*, 2006).

In fact, the details of the (hydrophobic) physical nature of non-functional interactions are not crucial for our model. What is indeed important is the assumption that the free energy of such interactions is proportional to the total hydrophobic surface covered upon formation of a nonfunctional complex involving these two proteins. There exists a significant evidence (Janin, 1995; Tamura and Privalov, 1997) that this may indeed be the case. The key aspect underlying this assumption is that non-functional interactions, having not been evolutionarily selected, represent random encounters between protein surfaces and as such adequately reflect the average surface composition of proteins. We use the Y2H data (see Materials and methods) to evaluate the significant parameters of this free energy model. In contrast, protein surfaces participating in functional PPIs have been presumably selected by evolution for stability and specificity, and various types of interactions were employed to achieve that (Lukatsky *et al*, 2007). As a result, these evolutionarily selected interactions are not adequately described by the sum of hydrophobicities of covered surfaces.

Fraction of proteins wasted in non-functional complexes: the phase diagram

If a protein with the surface energy E_i is not bound in a specific complex, it can either exist in a monomeric form or be bound to any of the non-functional partners. The ratio $[iR]/[i] = e^{-(E_i - \mu)/kT}$ between the last two concentrations depends on the chemical potential μ of the pool of all non-functional interaction partners (see Supporting Information for details). Then the probability for a protein to be in a monomeric form obeys the Fermi–Dirac distribution

$$[i]/([iR] + [i]) = \frac{1}{1 + e^{-(E_i - \mu)/kT}}$$
(10)

shown by the dashed line in Figure 2a. The value of μ can be found by solving

$$e^{-(E_{i}-\mu)/kT} = \frac{[iR]}{[i]} = \sum_{j} \frac{[j]}{C_{0}} e^{-E_{ij}/kT}$$
$$\cong \frac{\gamma \cdot m\bar{C}}{C_{0}} e^{-E_{i}/kT} \int e^{-(E_{j}+\Delta G_{(0)})/kT} \frac{f_{1}(E_{j})}{1 + e^{-(E_{j}-\mu)/kT}} dE_{j}$$
(11)

self-consistently. Here, we implicitly assume that the possible correlation between individual protein concentrations C_i and their stickiness parameters E_i can be neglected (see Supporting Information for the justification). The parameter γ is defined as the effective fraction of the total protein concentration $m\bar{C}$ that is not tied up inside the specific complexes and thus could participate in non-functional interactions. Although we assumed above that specific complexes do not participate in non-functional PPIs, in fact some of them do (however, with a reduced affinity). Indeed, their constituent proteins typically have more than one 'surface side' (Kim et al, 2006) and could use their other (usually less hydrophobic) surface sides for non-functional interactions. This would push the effective fraction γ closer to 1. For simplicity, throughout the manuscript we use $\gamma = 1$. This is justified, as the value of μ determined by the solution of Equation (11) only depends weakly on the value of γ within its plausible range (see Supporting Information for details).

The chemical potential μ goes up when the total protein concentration $m\bar{C}$ in the cytoplasm increases. As shown in Figure 2a, proteins with too sticky surfaces $E_i < \mu$ actively



Figure 3 The phase diagram for baker's yeast. The area in shade is the 'dead zone'. Its boundaries, defined by Equations (13) and (15), are blurred to indicate they are crossover rather than sharp transitions. The colours indicate the average fraction of proteins tied up inside non-functional complexes in the yeast cytoplasm as defined in Equation (12). The white circle shows the \bar{C} and m for a group of proteins coexpressed and colocalized in the yeast cytoplasm, whereas the white star and cross represents these parameters for the cell nucleus and mitochondria.

participate in non-functional bindings, and the interference of non-functional bindings for such proteins is strong: [iR] > [i]. On the other hand, less sticky proteins with $E_i > \mu$ have [iR] < [i] and thus belong to the safe zone in which the effects of non-functional interactions are weak. As will be described in the Materials and methods section, the high-throughput Y2H experiments allow us to estimate the parameters $E\bar{\cong} -7.0kT$ and $\sigma \cong 1.8kT$ of the distribution $f_1(E_i)$. For any given number of protein types m and average concentration \bar{C} , one could use Equation (11) to calculate the chemical potential μ and the individual danger levels [iR]/[i] for proteins with any energy E_i . The total fraction of proteins wasted in the non-functional complexes

$$\frac{\sum [iR]}{\sum ([i] + [iR])} \cong \int f_1(E_i) \frac{e^{-(E_i - \mu)/kT}}{1 + e^{-(E_i - \mu)/kT}} dE_i$$
(12)

is shown in colour in Figure 3. For the cytoplasm of a yeast cell with $m \cong 1800$ and $\bar{C} \cong 1 \,\mu\text{M}$ (Ghaemmaghami *et al*, 2003), Equation (11) gives $\mu \cong -8.9kT$. Thus, according to Equation (12), at any time point about 22% of proteins that are not in specific complexes are bound in non-functional complexes. Figure 3 also shows the nucleus and the mitochondria of a yeast cell by the white star and the white cross.

The next question is whether these compartments, especially the cytoplasm, fall in the dead zone. The range of the dead zone depends on specific and non-functional binding energies as well as concentrations, and these quantities vary greatly for proteins within a compartment. Therefore, Equations (3) and (4) can only find the boundaries of the dead zone for the simplified example system. However, for a realistic compartment such as the cytoplasm, we can still find the representative boundaries of the dead zone using the median values. Namely, we focus on a protein with median values of concentration, specific interaction $E_s \approx -16.5 kT$ and stickiness $E_i = E = -7.0 kT$, which correspond to the median non-functional interaction $E_n = 2E + \Delta G_{(0)} = -4.0kT$ (see the next section and Materials and methods for these values). Owing to the variability of stickiness and concentrations of proteins, Equations (3) and (4) should be modified. First, as the distribution of individual protein concentrations is nearly

lognormal, the median concentration of individual proteins is higher than the average value. We find $C_{\text{med}} \cong \overline{C}/6$ from data of the yeast cytoplasm. So Equation (3) becomes

$$C_{\text{med}} \cong \bar{C}/6 < C_{\text{crit}} = 2K_s = 2C_0 \exp(E_s/kT)$$
(13)

This gives $\bar{C}=0.82 \ \mu M$ as the boundary of the dead zone for the protein with median-specific interactions. Second, variation of surface 'stickiness' changes Equation (2a) to

$$[iR] = \frac{[i][R]}{C_0} \int \frac{f_1(E_j) dE_j}{\exp(E_{ij}/kT)} = \frac{[i][R]}{C_0} \exp\left[-\left(2\bar{E} + \Delta G_{(0)} - \frac{\sigma^2}{2}\right)/kT\right]$$
(14)

Comparing Equations (2b) and (14), we obtain a modification of Equation (4)

$$m_{\text{dead}} = \frac{mC}{6C_{\text{med}}} \approx \frac{[R]}{6[i']}$$
$$= \frac{1}{6} \exp\left[\left(2\bar{E} + \Delta G_{(0)} - \frac{\sigma^2}{2} - E_s\right)/kT\right] \qquad (15)$$

This results in $m_{dead} \cong 8200$ for the cytoplasm. This value is not necessarily very precise, as shown in Table I. m_{dead} is smaller if we use other Y2H data sets, or we obtain $m_{dead} \cong 4900$ if the full Ito data set is used. Figure 3 shows the dead zone in shade. The boundaries of the dead zone, characterized by Equations (13) and (15), are blurred to indicate that they are crossover rather than sharp transitions. We can see from Figure 3 that the cytoplasm of a yeast cell avoids the dead zone but locates near the corner corresponding to the upper limit of m and the lower limit of C. The nucleus and the mitochondria are also away but not far from the dead zone.

The phase diagram shown in Figure 3 represents all proteins in a given subcellular compartment by a single point. This is just an approximation, as different proteins would have different distances from the boundaries of the dead zone due to the variation in their energies of specific and non-functional interactions as well as in their concentrations. For instance, proteins with relatively strong specific binding could afford to have smaller concentrations C_i or, conversely, proteins with low concentrations have more difficulty maintaining the stability of specific complexes and thus need to have a stronger

Table I The summary of data from high-throughput Y2H experiments in different organisms and calculation results using the databases

Species	v	Ν		Fraction	$\frac{E{*}{-}2\bar{E}{-}\Delta G_{(0)}}{\sqrt{2}\sigma}$	$m_{\rm dead}$	$\frac{\sum [iR]}{\sum ([i]+[iR])}$
		$N_{\rm bait}$	$N_{\rm prey}$				
Yeast	1600	6000		6.2 e-5	3.8	8200	22%
Worm	4000	1900	1000	1.5 e-4	3.6	5300	26%
Fly	2000	1100	1000	2.4 e-4	3.5	4100	28%
Human1	3200	4500	5600	1.8 e−4	3.6	4900	26%
Human2	2800	7200		7.4 e−5	3.8	7500	23%

The data set of yeast merges interaction data in Uetz *et al* (2000) and the core data of Ito *et al* (2001), which contain interactions identified by at least three interaction sequence tags. The source of other data sets are as follows: worm (Li *et al*, 2004), fly (Giot *et al*, 2003), human1 (Stelzl *et al*, 2005) and human2 (Rual *et al*, 2005). The fraction is calculated as $\alpha v/(N^2/2)$ if $N_{\text{bait}} \approx N_{\text{prey}}$ and $\alpha v/(N_{\text{bait}} N_{\text{prey}})$ otherwise. The parameter α =0.7 is used in calculations.

binding energy of specific interactions. Also, sticky proteins have stronger than average non-functional interactions and thus tend to be more sequestered in non-functional complexes than the average fraction colour-coded in Figure 3. The complete description of the system should include the ratio [i]:[ii']:[iR] for every protein. However, the presently available experimental data (most notably the scarcity of data on binding energies of individual-specific interactions) does not allow for such a complete description.

All this imposes additional evolutionary constraints on the system. As was proposed by one of us (Maslov and Ispolatov, 2007), the result of such evolutionary pressure is that interactions between proteins with lower C_i would be stronger, that is, have lower (more negative) binding energy E_s translating into smaller dissociation constant K_s . However, from the evolutionary standpoint, the strength of these interactions must not be much stronger than the minimum required to ensure a sufficient concentration of a specific complex [ii'] (Maslov and Ispolatov, 2007).

Variability of individual protein concentrations and specific binding energies

Let us compare the histogram of specific binding energies with that of individual protein concentrations (Figure 4). The concentrations of individual proteins in yeast cells vary greatly (Ghaemmaghami *et al*, 2003; Belle *et al*, 2006) (open bars in Figure 4), with the median (typical) value $C_{\text{med}} \cong 0.2 \,\mu\text{M}$ or $\ln(C_{\text{med}}/2C_0) \approx -16.1$.

The dissociation constants (or corresponding binding energies) of specific interactions between yeast proteins have not been measured systematically. The binding energy data available from the PINT (Protein–protein Interaction Thermodynamic) database (Kumar and Gromiha, 2006) is quite limited. To gather sufficient statistics, we had to combine the binding energy data from different species including yeast, rat, human and so on (see Table S1, in Supporting Information). The resulting histogram is shown with filled bars in Figure 4.



Figure 4 The histograms of individual protein concentrations $\ln (C/2C_0)$ (white bars) and binding energies E_s of specific interactions (filled bars) (Kumar and Gromiha, 2006). Note that the peaks of these two distributions almost precisely agree with each other, whereas the distribution of binding energies is broader, especially on the negative side.

The data we choose are interactions between two types of wild-type proteins from the same species. Although most data are not from yeast cells, it could still provide some helpful information, because protein systems in different species have certain similarities.

Comparing the two histograms shown in Figure 4, we find that the median of $\ln (C_i/2C_0) \sim -16.1$ is just slightly above that of $E_s/kT \sim -16.5$. Therefore, most individual protein concentrations C_i are only slightly larger than the typical limiting value given by Equation (3). This would be enough for proteins within the safe zone where $[iR] \ll [i]$. On the other hand, the distribution of E_s/kT below the median is noticeably broader than that of $\ln (C_i/2C_0)$. This corresponds to the contribution of proteins that are inside the dangerous zone ([iR] > [i]) where specific interactions are required to be even stronger than required as per Equation (3).

This validates our observation that concentrations of most proteins are kept at or near the lower limit to ensure the formation of specific complexes. Maintaining low individual protein concentrations allows the cell to reduce the interference of non-functional interactions and, therefore, enhance the evolutionarily important protein diversity (the number of protein types that are simultaneously expressed and colocalized in the same compartment).

Discussion

In this work, we compared the concentrations of the three states of protein inside compartments of living cells: bound in specific complexes [*ii'*], bound in non-functional complexes [*iR*] and free (monomeric) [*i*]. As shown in Figure 1, the lower limit of the individual protein concentration is controlled by the requirement that [ii'] > [i], and the upper limit on the number of types of proteins that are co-expressed in the same intracellular compartment is set by the requirement that [ii'] > [iR]. The average concentration of a single protein and the protein diversity inside the yeast cytoplasm are in fact rather close to these two limits (see Figure 3). The size of the proteome is likely to be pushed up to the upper limit by evolution, because higher protein diversity allows for more complex biological functions. The average concentration of individual proteins is then pushed down towards the lower limit to reduce the waste of proteins in non-functional complexes and to improve the spatial and temporal efficiency of biochemical processes in a cell. Therefore, we suggest that the systems of coexisting proteins inside compartments of living cells, especially the cytoplasm, are located near the corner of the allowed region in the phase diagram. This requires the value of $m_{\rm cytoplasm}/m_{\rm dead}$ to fall in the interval 0.1–0.8 and $\sum [iR] / \sum ([i] + [iR])$ in the interval 0.1–0.35, and for Ito core data this further requires α to be in the interval 0.05–15.0. Our estimated ranges $\alpha \sim 0.3-0.9$, $m_{\rm cytoplasm}/$ $m_{\rm dead} \sim 0.11$ -0.4 and $\sum [iR] / \sum ([i] + [iR]) \sim 0.18 - 0.29$ are consistent with the hypothesis. Another argument in favour of keeping the specifically bound concentration [ii'] comparable with the free concentration [i] relies on the observation that along this horizontal line of the phase diagram in Figure 3, the functionally important concentration [ii'] is most sensitive to changes in total concentration C_i . This can be used for regulatory purposes allowing the cell to effectively turn on and off this particular biological function by changing the expression level of the protein *i*.

The phase diagram presented in Figure 3 highlights the major trade-off faced by compartments of living cells. To ensure the robust function, concentrations of individual proteins should be high enough to provide for sufficient formation of specific complexes. On the other hand, higher concentrations lead to an increase and possible dominance of non-functional interactions sequestering proteins and wasting precious resources. To overcome this limitation, the evolution could either strengthen the specific interactions, or weaken the non-functional ones, or (better yet) do both things at once. Nature has certainly exercised the first possibility by evolving stronger specific PPIs between proteins in functional complexes (e.g., by the virtue of cooperative binding). However, this route has its limitations, as designing strong specific interactions requires a prolonged evolutionary search in sequence space, which might be quite challenging to achieve. An alternative possibility is to evolve proteins with less hydrophobic surfaces. An interesting manifestation of this possibility could be seen in comparison of mesophilic with (hyper-)thermophilic proteins. It is clear from Equation (3) that the gap between specific and non-functional interaction energies should increase for organisms living at higher temperatures to keep the same proteome size. In this regard, it is important to note that surfaces of proteins from hyperthermophilic organisms are enriched with charged residues (Glyakina et al, 2007) and that majority of those residues do not form salt bridges (Goldstein, 2007). It is therefore tempting to speculate that the main reason for this enrichment with charged residues is to reduce non-functional PPIs in hyperthermophiles to keep their proteome size at a physiologically acceptable level.

Multicellular organisms, however, can go around the upper limit of protein diversity. By separating different kinds of proteins in different cell types, they can avoid coexistence of too many protein types and reduce the amount of nonfunctional interactions. Therefore, they are allowed to develop more complicated functions through cellular differentiation. This could be one of the crucial reasons for multicellular organisms to emerge in the course of the evolution. It will be also interesting to study whether there is a systematic difference in stickiness of proteins from prokaryotes with less compartmentalized cells and higher organisms whose cells are highly compartmentalized. It could be an interesting undertaking to systematically compare surface hydrophobicities with specific interaction energies between groups of proteins from simplest-not compartmentalized-cells to single-cell eukaryotes to multicellular organisms.

The specific estimates of tolerable concentrations and proteome sizes made in this study are on the basis of assessment of the specific and non-functional interactions, which are not necessarily very accurate. An important source of uncertainty lies in the estimate of the parameter α , the fraction of non-functional interactions among all interactions detected in Y2H data sets. To see the impact of α on the results, if α is in the interval 0.4–0.9, the value of m_{dead} is in the interval 7300–10500 and $\sum [iR]/\sum ([i] + [iR])$ is in the interval 21–23%. We also face a dilemma in choosing either the full

or the core data from Ito et al (2001). As the filtered Ito core data requires three interaction sequence tags, many nonfunctional interactions that are only slightly stronger than the threshold $K_d^* \approx 1 \,\mu\text{M}$ (Estojak *et al*, 1995) may not be included in the data set. The number of non-functional PPIs is better reflected by the full Ito data set, which contains ~ 4500 interactions and results in m_{dead} =4900 and average fraction [iR]/([iR] + [i]) = 26% for yeast cytoplasm. However, the full Ito data set might contain many false interactions due to experimental artefacts such as bait self-activation. To be cautious, we choose the united data of Uetz et al (2000) and Ito core rather than the full Ito data set for yeast PPIs to avoid experimental artefacts, and keep in mind that the result m_{dead} =8200 might be overestimated and the average fraction [iR]/([iR] + [i]) = 22% might be slightly underestimated. In consistence with this expectation, the result of m_{dead} is also smaller if we try the calculation with Y2H data sets of other organisms (see Table I).

There are other sources of uncertainties in the calculation besides the estimate of non-functional interactions in Y2H data sets. For binding energies of specific interactions, we used the PINT database (Kumar and Gromiha, 2006), which contains the experimentally measured strength of specific interactions between some of the proteins from various organisms, which we subsequently extrapolated to yeast. The data on such interactions are still scarce and its accuracy is not entirely clear. For non-functional interactions, our estimates are on the basis of a straightforward analysis of Y2H experiments that assumes a 'hard cutoff' E^* on binding energy E_{ii} of detected PPIs. A more sophisticated method that uses a 'soft' cutoff assumption (Shi et al, 2006) for the detection of PPI in Y2H experiments may give somewhat different estimates for strength and variation of non-functional interactions. For example, using the soft-cutoff model modifies our estimates for the non-functional interaction energy, but still predicts that around 15% of proteins are sequestered in non-functional complexes (Maslov, unpublished data). However, the fundamental finding of this work concerns the deep interrelation between specific and non-functional interactions and sizes of proteomes in living cells and is independent on the specific numeric values used in our estimates.

Above, we assumed that the biologically active state of all proteins is in the complex with their specific binding partner. Although this is certainly the case for a significant fraction of all proteins in the cell, there also exist proteins that are biologically active in their monomeric form. Our definition of death zone for such proteins should be adjusted to require [i] > [iR]. Instead of Equation (3), the lower bound on concentrations of monomeric proteins is imposed by purely physiological factors. For example, the abundance of metabolic enzymes is likely to be dictated by typical concentrations of their substrates. There are also cases of proteins with more than one specific partner. Under these circumstances, the condition [ii'] > [iR] should hold independently for each of the functionally important partners.

The interference of non-functional PPIs with biochemical efficiency of all intracellular functional processes is quantified by the ratio [iR]/[i], indicating whether proteins that do not participate in specific complexes are free in cytoplasm or non-functionalally bound to non-functional partners. Although the

average ratio [iR]/([iR] + [i])=22% does not seem particularly large, due to the variation in stickiness, certain proteins would have much greater ratio [iR]/[i] than average (see Materials and methods for details). Thus, the requirement of biochemical efficiency is pushing individual protein concentrations down in the cytoplasm. As a result, the cytoplasm of a cell can only stay at the corner near the triple point in the phase diagram. This conclusion might be generalized to apply to many other cells.

Materials and methods

Obtaining individual protein concentrations from experimental results

The expression level of proteins in yeast cell cytoplasm, nucleus and mitochondria are taken from experiments (Ghaemmaghami *et al*, 2003; Huh *et al*, 2003; Belle *et al*, 2006). The average volume of a haploid yeast cytoplasm is ~25 fL (Jorgensen *et al*, 2007), which is similar to the cell volume. The volume of nucleus (Jorgensen *et al*, 2007) and mitochondria (Visser *et al*, 1995) are both about 7% of cell volume, although early results of mitochondria (Grimes *et al*, 1974; Stevens, 1977) vary from 3 to 14%.

Most proteins within a given subcellular compartment diffuse over its whole volume. Although some proteins enhance their local concentrations through co-localization (Kuriyan and Eisenberg, 2007) near the membrane, such membrane-bound proteins are only a small fraction among all protein types, for example, no more than 15% for the cytoplasm (Kumar *et al*, 2002). Thus, this effect constitutes just a small correction to our estimates for individual subcellular compartments and will not be considered in this study.

These individual protein concentrations were obtained mainly for the cells in the G1 stage in the cell cycle. Are they really representative of what happens at other stages of the cell cycle? To address this concern, we point out that the median half-life of yeast proteins is ~40 min (Belle *et al.*, 2006) whereas the cell cycle takes ~90 min. Therefore, the concentrations of most yeast proteins do not change dramatically in the course of a cell cycle, and the above estimate using the average concentration in the G1 stage remains valid.

Our calculations are quite robust with respect to random errors in measurements of concentrations of individual proteins, as we use only the total concentration of all proteins and the number of protein types. Even large systematic errors in protein concentrations will not significantly impact our qualitative results. For example, a factor of 2 or 1/2 error in the total protein concentration will change the fraction $\sum [iR] / \sum ([i] + [iR])$ from 22% to only 28 or 17% correspondingly.

Estimating the strength of non-functional interactions from Y2H experiments

In this section, we estimate the mean and the standard deviation of the binding energy of all non-functional PPI from the Y2H experiments. This estimate was used in the previous section to derive the phase diagram (Figure 3) for the fraction of non-functional complexes in the cytoplasm of yeast cells.

The distribution of binding energy of non-functional interactions is determined by its average $2\vec{E} + \Delta G_{(0)}$ and variation $\sqrt{2\sigma}$ as defined by Equation (7). High-throughput Y2H experiments systematically inspect PPIs between nearly all pairs of proteins encoded in the genome of a given organism. Thus, they contain a useful information, allowing one to estimate the values of \vec{E} and σ .

A positive signal is detected in a Y2H experiment if a PPI is sufficiently strong, and the detection threshold for the dissociation constant is experimentally estimated as $K_d^* \approx 1 \,\mu\text{M}$ or $E^* \approx kT \ln(K_d^*/C_0) = -14kT$ (Estojak *et al*, 1995).

The interactions detected in Y2H experiments are not necessarily biologically functional. For one thing, the *in vivo* concentrations of individual proteins (Ghaemmaghami *et al*, 2003; Huh *et al*, 2003; Belle *et al*, 2006) are often considerably lower than those of bait-and-prey



Figure 5 The fraction of Y2H interactions from the full set of Ito *et al* (2001) confirmed in a curated set of biologically functional interactions that were independently reported in two or more publications not using the Y2H technique. The *x* and *y* axis are the *in vivo* protein concentrations (Ghaemmaghami *et al*, 2003) in yeast cytoplasm. One can see that Y2H interactions between proteins with relatively high *in vivo* concentrations ($>0.1 \mu$ M) have considerably higher chances of being biologically functional.

proteins used in Y2H experiments. As a result, there is no guarantee that even a reproducible interaction detected in a Y2H screen will occur with any significant probability for in vivo concentrations of interacting proteins. This view is supported by our observation (Figure 5) that the overlap between Y2H interactions from the full set of Ito et al (2001) and the curated set of biologically functional interactions is biased towards proteins with high in vivo concentrations. Below, we use the large-scale Y2H data sets to determine the parameters of non-functional interactions necessary for our model. As pointed out above, the Y2H technique detects interactions between baits and prevs at elevated concentrations, and therefore it samples a much broader set of PPIs than those that are occurring with high probability at in vivo concentration of proteins. Another advantage of large-scale Y2H screens is that they check for possible interactions among all possible pairs of proteins without any bias towards functionally related pairs. As a result, large-scale Y2H data sets contain ample information about the distribution of affinities for nonfunctional interactions, especially for proteins with low in vivo concentrations (see Figure 5).

We define α as the fraction of non-functional interactions among all interactions detected in large-scale Y2H screens. The list of all interactions detected in the largest Y2H screen of yeast proteins (Ito et al, 2001) contains many false-positives (Deane et al, 2002; Vidalain et al, 2004; Huang et al, 2007) due to reproducible yet non-functional interactions as well as methodological artefacts such as bait selfactivations (Nakayama et al, 2002). The Ito core data set, formed by the interactions repeatedly detected in the course of the experiment (three times or more) is more reliable and reproducible, but is still thought to contain over 40% false-positives (Deane et al, 2002). Most importantly, the internal reproducibility condition used to generate this data set likely eliminates most of the artefacts due to bait selfactivation. Indeed, the interaction partners of a self-activating bait are essentially randomly selected from a large pool of prey proteins. Hence, it is exceedingly improbable to repeatedly detect the same prey protein three or more times. We assume that the remaining falsepositives in the Ito core data set are real and reproducible, but not formed under the in vivo conditions inside a living cell and thus do not contribute to its biological functioning. The lower bound on the fraction of non-functional interactions in the Ito core data set is set by the following observation: among the 522 interacting pairs with known subcellular localizations of both proteins (Huh et al, 2003), 15% correspond to pairs of proteins that are never co-localized in the

same intracellular compartment, and thus in principle cannot bind each other *in vivo*. We expect the fraction of all non-functional interactions (including those caused by concentration effects of colocalized proteins) to be significantly higher than 15%.

On the other hand, many strong and functional interactions must be missed in large-scale Y2H screens as false-negatives, because another filtered Y2H data set (Uetz *et al*, 2000) of yeast proteins has few overlaps with the Ito core data set. Thus, we expect that many strong enough non-functional interactions can be missed in Y2H data sets. To take this effect into account, we merge the Ito core data set and the filtered Uetz data set to obtain a more complete data set of yeast proteins. Combining the estimate of false-positives in Y2H data sets reported by Deane *et al* (2002) with this false-negative effect, we propose α =0.7 as the ballpark fraction of non-functional PPIs among all interactions detected in Y2H screens. The impact of uncertainty in our estimation of this parameter α is assessed in Discussion.

For *N* types of proteins tested in a high-throughput experiment, there are $N(N+1)/2 \approx N^2/2$ possible interactions. If *v* distinct interacting pairs pass the dissociation constant cutoff and αv interactions are non-functional, their fraction among all pairs is given by

$$\frac{\alpha v}{N^2/2} = \int_{-\infty}^{E^*} f_2(E_{ij}) dE_{ij} = [Erf\left(\frac{E^* - 2\bar{E} - \Delta G_{(0)}}{\sqrt{2} \cdot \sqrt{2}\sigma}\right) + 1]/2$$
(16)

where *Erf* is the error function. This equation defines a relationship between the parameters E and σ of the distribution of binding energies of non-functional interactions and the parameters v and N of a high-throughput Y2H experiment. The genome-wide Y2H studies in baker's yeast *Saccharomyces cerevisiae* (Uetz *et al*, 2000; Ito *et al*, 2001) contain $v \approx 1600$ interacting pairs between about $N \cong 6000$ tested proteins. We use $\alpha = 0.7$ to find the number of non-functional interactions and solve Equation (16) to get

$$(E^* - 2\bar{E} - \Delta G_{(0)})/\sqrt{2}\sigma \approx 3.8$$
 (17)

We solve Equations (8) and (17) to find $\vec{E} \cong -7.0kT$ and $\sigma \cong 1.8kT$. Hence, the estimated median value of non-functional interactions is $2\vec{E} + \Delta G_{(0)} = -4.0kT$ or $K_n \sim 18$ mM. This value is in agreement with the suggestion (Kuriyan and Eisenberg, 2007) that K_n is a little larger than 10 mM for non-functional PPIs.

In Table I, we also show the value of $(E^* - 2\bar{E} - \Delta G_{(0)})/\sqrt{2}\sigma$ for several large-scale Y2H experiments, which are strikingly similar for different species. Although in this study we focus on the baker's yeast for which protein concentrations were systematically measured, our estimates of the parameters of non-functional interactions could be applicable to other organisms listed in Table I.

Protein-to-protein variability of non-functional sequestration ratio

As estimated above for yeast cytoplasm, the average fraction [iR]/([iR] + [i]) of proteins sequestered inside non-functional complexes is about 22%. As the chemical potential $\mu \cong -8.9kT$ is lower than the median energy $\vec{E} \cong -7.0kT$, the ratio $[iR]/[i] = \exp[-(E_i - \mu)/kT]$ is below one for most of the proteins. Around 20 of the most sticky types of proteins have [iR]/[i] > 10, and for the protein with highest hydrophobicity, the ratio reaches $[iR]/[i] \approx 60$. If we add the correction that proteins in specific complexes have reduced affinity for nonfunctional interactions, the highest ratio is reduced to $[iR]/[i] \approx 35$ (see Supporting Information).

Supplementary information

Supplementary information is available at the *Molecular Systems Biology* website (www.nature.com/msb).

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References

- Aloy P, Bottcher B, Ceulemans H, Leutwein C, Mellwig C, Fischer S, Gavin AC, Bork P, Superti-Furga G, Serrano L, Russell RB (2004) Structure-based assembly of protein complexes in yeast. *Science* 303: 2026–2029
- Aloy P, Russell RB (2002) Potential artefacts in protein-interaction networks. *FEBS Lett* **530**: 253–254
- Aloy P, Russell RB (2006) Structural systems biology: modelling protein interactions. *Nat Rev Mol Cell Biol* **7:** 188–197
- Bahadur RP, Chakrabarti P, Rodier F, Janin J (2004) A dissection of specific and non-specific protein–protein interfaces. *J Mol Biol* **336**: 943–955
- Belle A, Tanay A, Bitincka L, Shamir R, O'Shea EK (2006) Quantification of protein half-lives in the budding yeast proteome. *Proc Natl Acad Sci USA* **103**: 13004–13009
- Bogan AA, Thorn KS (1998) Anatomy of hot spots in protein interfaces. *J Mol Biol* **280**: 1–9
- Deane CM, Salwinski L, Xenarios I, Eisenberg D (2002) Protein interactions: two methods for assessment of the reliability of high throughput observations. *Mol Cell Proteomics* 1: 349–356
- Deeds EJ, Ashenberg O, Gerardin J, Shakhnovich EI (2007) Robust protein protein interactions in crowded cellular environments. *Proc Natl Acad Sci USA* 104: 14952–14957
- Deeds EJ, Ashenberg O, Shakhnovich EI (2006) A simple physical model for scaling in protein–protein interaction networks. *Proc Natl Acad Sci USA* **103:** 311–316
- Eisenberg D, McLachlan AD (1986) Solvation energy in protein folding and binding. *Nature* **319**: 199–203
- Estojak J, Brent R, Golemis EA (1995) Correlation of two-hybrid affinity data with *in vitro* measurements. *Mol Cell Biol* **15**: 5820–5829
- Gavin AC, Aloy P, Grandi P, Krause R, Boesche M, Marzioch M, Rau C, Jensen LJ, Bastuck S, Dumpelfeld B, Edelmann A, Heurtier MA, Hoffman V, Hoefert C, Klein K, Hudak M, Michon AM, Schelder M, Schirle M, Remor M *et al* (2006) Proteome survey reveals modularity of the yeast cell machinery. *Nature* **440**: 631–636
- Ghaemmaghami S, Huh WK, Bower K, Howson RW, Belle A, Dephoure N, O'Shea EK, Weissman JS (2003) Global analysis of protein expression in yeast. *Nature* **425**: 737–741
- Giot L, Bader JS, Brouwer C, Chaudhuri A, Kuang B, Li Y, Hao YL, Ooi CE, Godwin B, Vitols E, Vijayadamodar G, Pochart P, Machineni H, Welsh M, Kong Y, Zerhusen B, Malcolm R, Varrone Z, Collis A, Minto M *et al* (2003) A protein interaction map of *Drosophila melanogaster. Science* **302**: 1727–1736
- Glyakina AV, Garbuzynskiy SO, Lobanov MY, Galzitskaya OV (2007) Different packing of external residues can explain differences in the thermostability of proteins from thermophilic and mesophilic organisms. *Bioinformatics* 23: 2231–2238
- Goldstein RA (2007) Amino-acid interactions in psychrophiles, mesophiles, thermophiles, and hyperthermophiles: insights from the quasi-chemical approximation. *Protein Sci* **16**: 1887–1895
- Grimes GW, Mahler HR, Perlman RS (1974) Nuclear gene dosage effects on mitochondrial mass and DNA. *J Cell Biol* **61**: 565–574
- Huang H, Jedynak BM, Bader JS (2007) Where have all the interactions gone? Estimating the coverage of two-hybrid protein interaction maps. *PLoS Comput Biol* 3: e214
- Huh WK, Falvo JV, Gerke LC, Carroll AS, Howson RW, Weissman JS, O'Shea EK (2003) Global analysis of protein localization in budding yeast. *Nature* **425**: 686–691

- Ito T, Chiba T, Ozawa R, Yoshida M, Hattori M, Sakaki Y (2001) A comprehensive two-hybrid analysis to explore the yeast protein interactome. *Proc Natl Acad Sci USA* 98: 4569–4574
- Janin J (1995) Protein-protein recognition. Prog Biophys Mol Biol 64: 145–166
- Janin J (1996) Quantifying biological specificity: the statistical mechanics of molecular recognition. *Proteins* **25**: 438–445
- Jorgensen P, Edgington NP, Schneider BL, Rupes I, Tyers M, Futcher B (2007) The size of the nucleus increases as yeast cells grow. *Mol Biol Cell* **18**: 3523–3532
- Kim PM, Lu LJ, Xia Y, Gerstein MB (2006) Relating three-dimensional structures to protein networks provides evolutionary insights. *Science* 314: 1938–1941
- Kortemme T, Baker D (2002) A simple physical model for binding energy hot spots in protein-protein complexes. *Proc Natl Acad Sci* USA 99: 14116–14121
- Kumar A, Agarwal S, Heyman JA, Matson S, Heidtman M, Piccirillo S, Umansky L, Drawid A, Jansen R, Liu Y, Cheung KH, Miller P, Gerstein M, Roeder GS, Snyder M (2002) Subcellular localization of the yeast proteome. *Genes Dev* 16: 707–719
- Kumar MD, Gromiha MM (2006) PINT: protein–protein interactions thermodynamic database. Nucleic Acids Res 34: D195–D198
- Kuriyan J, Eisenberg D (2007) The origin of protein interactions and allostery in colocalization. *Nature* **450**: 983–990
- Lo Contc L, Chothia C, Janin J (1999) The atomic structure of proteinprotein recognition sites. J Mol Biol **285:** 2177–2198
- Li S, Armstrong CM, Bertin N, Ge H, Milstein S, Boxem M, Vidalain PO, Han JD, Chesneau A, Hao T, Goldberg DS, Li N, Martinez M, Rual JF, Lamesch P, Xu L, Tewari M, Wong SL, Zhang LV, Berriz GF *et al* (2004) A map of the interactome network of the metazoan *C. elegans. Science* **303**: 540–543
- Lukatsky DB, Shakhnovich BE, Mintseris J, Shakhnovich EI (2007) Structural similarity enhances interaction propensity of proteins. *J Mol Biol* **365**: 1596–1606
- Maslov S, Ispolatov I (2007) Propagation of large concentration changes in reversible protein-binding networks. *Proc Natl Acad Sci USA* **104:** 13655–13660
- Nakayama M, Kikuno R, Ohara O (2002) Protein–protein interactions between large proteins: two-hybrid screening using a functionally classified library composed of long cDNAs. *Genome Res* **12**: 1773–1784
- Noskov SY, Lim C (2001) Free energy decomposition of protein-protein interactions. *Biophys J* 81: 737–750
- Rual JF, Venkatesan K, Hao T, Hirozane-Kishikawa T, Dricot A, Li N, Berriz GF, Gibbons FD, Dreze M, Ayivi-Guedehoussou N, Klitgord N, Simon C, Boxem M, Milstein S, Rosenberg J, Goldberg DS, Zhang LV, Wong SL, Franklin G, Li S *et al* (2005) Towards a proteome-scale map of the human protein–protein interaction network. *Nature* 437: 1173–1178
- Scar RP (2004) Specific protein-protein binding in many-component mixtures of proteins. *Phys Biol* 1: 53–60
- Shi YY, Miller GA, Qian H, Bomsztyk K (2006) Free-energy distribution of binary protein–protein binding suggests cross-species interactome differences. *Proc Natl Acad Sci USA* 103: 11527–11532
- Stelzl U, Worm U, Lalowski M, Haenig C, Brembeck FH, Goehler H, Stroedicke M, Zenkner M, Schoenherr A, Koeppen S, Timm J, Mintzlaff S, Abraham C, Bock N, Kietzmann S, Goedde A, Toksoz E, Droege A, Krobitsch S, Korn B *et al* (2005) A human protein–protein interaction network: a resource for annotating the proteome. *Cell* **122:** 957–968
- Stevens BJ (1977) Variation in number and volume of the mitochondria in yeast according to growth conditions. *Biologie Cellulaire* **28**: 37–56
- Tamura A, Privalov PL (1997) The entropy cost of protein association. *J Mol Biol* **273:** 1048–1060
- Uetz P, Giot L, Cagney G, Mansfield TA, Judson RS, Knight JR, Lockshon D, Narayan V, Srinivasan M, Pochart P, Qureshi-Emili A,

Li Y, Godwin B, Conover D, Kalbfleisch T, Vijayadamodar G, Yang M, Johnston M, Fields S, Rothberg JM (2000) A comprehensive analysis of protein-protein interactions in *Saccharomyces cerevisiae*. *Nature* **403**: 623–627

- Vidalain PO, Boxem M, Ge H, Li S, Vidal M (2004) Increasing specificity in high-throughput yeast two-hybrid experiments. *Methods* **32:** 363–370
- Visser W, van Spronsen EA, Nanninga N, Pronk JT, Gijs Kuenen J, van Dijken JP (1995) Effects of growth conditions on mitochon-

drial morphology in *Saccharomyces cerevisiae*. Antonie Van Leeuwenhoek **67**: 243–253

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