Revealing static and dynamic modular architecture of the eukaryotic protein interaction network

Kakajan Komurov* and Michael White

Department of Cell Biology, University of Texas Southwestern Medical Center, Dallas, TX, USA
* Corresponding author. Department of Cell Biology, University of Texas Southwestern Medical Center, 5323 Harry Hines Blvd, Dallas, TX 75390-9039, USA.
Tel.: +1 214 648 3015; Fax: +1 214 648 8694; E-mail: kakajan.komurov@utsouthwestern.edu

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In an effort to understand the dynamic organization of the protein interaction network and its role in the regulation of cell behavior, positioning of proteins into specific network localities was studied with respect to their expression dynamics. First, we find that constitutively expressed and dynamically co-regulated proteins cluster in distinct functionally specialized network neighborhoods to form static and dynamic functional modules, respectively. Then, we show that whereas dynamic modules are mainly responsible for condition-dependent regulation of cell behavior, static modules provide robustness to the cell against genetic perturbations or protein expression noise, and therefore may act as buffers of evolutionary as well as population variations in cell behavior. Observations in this study refine the previously proposed model of dynamic modularity in the protein interaction network, and propose a link between the evolution of gene expression regulation and biological robustness.

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Introduction

Revealing complex patterns and organizational principles of biological systems is an important goal of systems biology. Recent studies have considerably expanded our understanding of organizational principles underlying biological networks. Much of the past effort in this field has focused on the topological properties of protein interaction and gene regulation networks and many of their design principles have been uncovered, such as the scale-free topology (Albert et al., 2000; Jeong et al., 2000), modularity (Ihmels et al., 2002; Ravasz et al., 2002; Spirin and Mirny, 2003; Han et al., 2004; Gavin et al., 2006), disassortativeness (Maslov and Sneppen, 2002) and enrichment for certain network motifs (Milo et al., 2002; Harbison et al., 2004; Luscombe et al., 2004). At the level of cell behavior, these properties are thought to promote robustness (Albert et al., 2000; Jeong et al., 2001; Maslov and Sneppen, 2002) and reliability of information processing (Klemm and Bornholdt, 2005). The studies described above mainly analyzed static protein interaction networks without accounting for dynamic properties that arise as a result of gene expression programs that modulate the expression of proteins in the network. Integration of protein interaction data with the gene expression data in recent years, however, has given some important insights into the dynamic organization of the eukaryotic protein interaction network (Ge et al., 2001; Han et al., 2004; Ihmels et al., 2004; Kharchenko et al., 2005; de Lichtenberg et al., 2005). For example, it has been found that co-regulated proteins frequently interact with each other (Ge et al., 2001), and metabolic enzymes topologically close to each other in the metabolic network (Kharchenko et al., 2005) or those that participate in the same metabolic pathway (Ihmels et al., 2004) are also frequently co-regulated. Another study has reported differential positioning of proteins in the protein interaction network based on their coexpression properties with their interacting neighbors in the network (Han et al., 2004). These studies have suggested that a strong correlation exists between topological positioning of proteins in the network and their expression properties.

Recent large-scale experimental and computational studies have delineated the action of gene expression programs under various conditions (Gasch et al., 2000; Segal et al., 2003a; Harbison et al., 2004; Luscombe et al., 2004). Importantly, these studies defined both condition-dependent and condition-independent (constitutive) expression patterns (Luscombe et al., 2004; de Lichtenberg et al., 2005). Given that proteins are subject to variable modes of regulation, we considered the topological positioning of proteins with different levels of transcriptional regulation in protein interaction networks. Previous studies have examined the correlation of co-regulation of proteins with their topological positioning in the network. Instead, we examined how the protein products of regulated versus nonregulated genes (dynamic and static proteins, respectively) are positioned in the protein interaction network relative to each other. We identified organizational principles in the protein interaction network that appear to dictate the specific relative topological positioning of dynamic and static proteins. This information expands our under-

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standing of protein network dynamics, gives systems-level insights into how gene expression programs may modulate the protein network architecture and cell behavior, and suggests a link between the evolution of cellular robustness and the evolution of gene expression regulation.

Results

Expression variance of genes across multiple conditions

To identify genes with condition-dependent or constitutive expression, we leveraged legacy genomic expression profiles derived from cells exposed to multiple conditions. A compendium of 272 microarray experiments from six different data sets from the *Saccharomyces* Genome Database (SGD, ftp://ftp.yeastgenome.org/yeast/) was compiled and used to calculate the statistical variance of expression profiles for each gene across these 272 experiments. The statistical variance of the expression profile of a gene was assumed to reflect the frequency and magnitude of modulation of its gene product under diverse conditions, such that a low variance would indicate that the protein is relatively static and a high variance would indicate that the protein is relatively dynamic.

For each gene in the genome, an expression variance (EV) was assigned, as defined by the quantile value of the variance of its expression profile in the genomic distribution of variances, such that the EV closest to 0 indicates the gene with the lowest variance in the genome (least dynamic), and the EV value of 1 indicates the gene with the most dynamic expression pattern in the genome. The expression dynamics of the bottom 10 and top 10 genes in the genome-wide EV distribution are shown in Supplementary Figure 1.

Figure 1 Interaction pattern of proteins according to their EV. (A) Boxplot of proteins in each of 50 bins with the given EV versus their neighborhood EV. (B) Interaction preference matrix of the yeast network. Each square represents the number of interactions between corresponding bins. Left panel: Interaction preference matrix of the actual network, right panel: Interaction preference matrix of randomized network achieved by randomly shuffling the positions of proteins in the network (right panel). (C) Interaction preference matrix of proteins with different node degrees corresponding to the four quartiles of the node degree distribution. Proteins were binned according to their EVs and node degrees. Each square represents the normalized number of interactions between proteins with given node degree (k) and EV. Normalization of a square (i, j) in the matrix was carried out by calculating the number of interactions between proteins in the bins i and j, and dividing that number with the total number of interactions that proteins in bins i and j have. Color key shows the normalized number of interactions between bins.
We considered that this metric may capture random noise in the expression of genes with low mRNA levels as high EV, and/or may capture the low overall variations in the levels of highly abundant mRNAs as low EV. However, we found that there is a significantly high positive correlation between the calculated EV values of genes and their mRNA abundance values in the cell (Spearman’s \( r = 0.21, P < 1 \times 10^{-16} \)), indicating that low-EV genes are expressed at lower levels than the high-EV genes. This positive correlation indicates that the high-EV gene set is not enriched for low-abundance genes and reflects the extent of gene regulation rather than measuring artifacts owing to low mRNA abundance.

A high-confidence yeast protein interaction network of 2315 proteins connected with 5356 interactions was derived using the confidence scores assigned to each interaction by the study of Bader et al (2004). Importantly, all findings presented below were reproducible using a scoring scheme different from the same study to obtain a slightly different network, or by using an independent high-quality network from a recent large-scale study (Krogan et al., 2006) (data not shown).

**Interaction preferences of static and dynamic proteins**

In order to understand how proteins with different expression dynamics assemble within the network, proteins were analyzed in the context of their network neighborhoods; here, defined as the set of binding partners of a protein. A comparison of EVs of proteins with the EVs of their neighbors was used to examine the relative distribution of static and dynamic proteins within the network. For this purpose, we defined neighborhood EV of a protein to be the average of EV values of its interacting partners (neighbors) in the network. There is a strikingly high positive correlation between EVs of proteins and their neighborhood EVs (Spearman’s \( r = 0.32, n = 2245; P = 1 \times 10^{-54} \)) (Figure 1A), which suggests that proteins have similar expression dynamics as their immediate neighbors in the protein interaction network. To examine this correlation at a higher resolution, proteins were grouped into 50 bins according to similarity of EV scores and a 50 \( \times \) 50 interaction preference matrix was constructed. The density of interactions between every bin pair was displayed as the total number of ‘interbin’ protein interactions. Consistent with the statistical correlation above (Figure 1A), there is a high density of interactions among the low-EV bins or among the high-EV bins (Figure 1B). Moreover, the interaction densities between low-EV and high-EV bins appear to be extremely low. This interaction pattern was not observed in a randomized network (Figure 1B). These results suggest that the protein interaction network is enriched for sub-networks that are primarily composed of either static proteins or dynamic proteins, but not both.

The interaction preference matrix shows the number of interactions between bins, which may bias the matrix for the interaction profiles of highly connected proteins. In order to check this, we tested the interaction profiles of proteins with different numbers of interactions (node degrees) and different EVs. We found that the strong correlation in Figure 1B is most apparent for proteins with high node degrees (hubs) (Figure 1C), although the statistical correlation between EV and neighborhood EV in less-connected proteins is also significantly high (data not shown). These observations suggest that proteins having more interaction partners in the network are segregated into distinct network neighborhoods that are characterized by low and high EVs, respectively.

**Functional specialization of static and dynamic neighborhoods**

In order to understand what these distinct protein neighborhoods represent, the sub-network formed by hubs, defined as proteins with greater than six interactions, was visualized. A plot of interactions between hubs reveals that the static and dynamic neighborhoods represented distinct densely connected large clusters of static and dynamic proteins (Figure 2A). A densely connected cluster of proteins in the network is likely to represent a functional module (Pereira-Leal et al., 2004), that is, a set of interacting proteins dedicated to a specific cellular process such as the mRNA splicing machinery or the proteasome. Dynamically expressed proteins within a functional module are expected to be coexpressed with each other (Ge et al., 2001; Segal et al., 2003b). In order to check this, we defined neighborhood Pearson correlation coefficient (PCC) as a measure of how well proteins in a neighborhood are coexpressed with each other (see Materials and methods). In agreement with previous studies (Ge et al., 2001; Segal et al., 2003b), we found that neighborhoods of dynamic proteins in clusters are highly coexpressed with each other (Figure 2B). Consistent with their static nonvariant expression pattern, static proteins within clusters had lower neighborhood PCC (nPCC) (Figure 2B). These observations suggest the existence of two distinct types of large modules in the cell: those composed of static proteins (static modules), which are presumably always present in the cell as the expression of their members does not seem to be regulated, and those composed of co-regulated dynamic proteins (dynamic modules), which are expressed in a condition-dependent manner.

The current notion of functional modules predicts that a set of interacting proteins that are highly coexpressed is likely to be specialized to a specific process (Ihmels et al., 2002; Segal et al., 2003b; Kharchenko et al., 2005), and some studies suggest that the protein interaction network is enriched for interactions between co-regulated proteins (Ge et al., 2001; Ihmels et al., 2004). We show that sets of interacting static proteins, which are supposedly constitutively present in the cell but do not have high statistical correlation in their expressions, also may represent specialized functional modules, and that they are at least as abundant in the cell as the sets of interacting dynamic proteins that are highly coexpressed. In order to test this hypothesis, a simple function that compares a protein’s Gene Ontology (GO) (Ashburner et al., 2000) annotations with that of its neighbors was derived to quantitate the functional specialization of a protein’s neighborhood. This function, ‘neighborhood function homology’ (see Materials and methods), generates values in the range from 0 (no shared GO terms between a protein and its neighbors) to 1 (all of GO terms assigned to a protein are shared with its neighbors).
Neighborhood function homology of static proteins (EV<0.25, i.e. lower quartile of genomic distribution) in the network negatively correlates with their neighborhood EV with a high significance (Spearman’s $\rho = -0.41$, $P = 1.8 \times 10^{-18}$), suggesting that static proteins interacting with other static proteins are found in functionally specialized neighborhoods. On the other hand, neighborhood function homology of dynamic proteins (EV>0.75, higher quartile of genomic distribution) in the network negatively correlates with their neighborhood EV with a high significance (Spearman’s $\rho = -0.41$, $P = 1.8 \times 10^{-18}$), suggesting that dynamic proteins interacting with other dynamic proteins are found in functionally specialized neighborhoods. Figure 2 Static and dynamic neighborhoods resemble functional modules. (A) Plot of the sub-network formed by hubs. Proteins are colored according to their EV. (B) Plot of hubs’ nPCC values against their neighborhood density. Neighborhood density (see Materials and methods) is a measure of how densely the neighbors of a protein are connected to each other, and ranges from 0, for the least dense neighborhood, to 1, for a maximally densely connected neighborhood. Proteins within dense clusters are expected to have high neighborhood densities. Dots (hubs) are colored according to the hub EVs (left panel) and neighborhood EVs (right panel).
network (see Materials and methods). The static network measure functional specialization of the interactions in a coherence in a network generated by random draws of and to see if it is possible to achieve a similar level of functional significance of modular composition of the static network is enriched for functional modules. In order to test the apparent functional coherence suggests that the static network cell-cycle regulation (see Supplementary Table 1). This ing mRNA transcription and splicing, vesicle transport and functionally coherent, representing various functions includ- ing proteins (Supplementary Table 1). The functional annotations disconnected sub-networks ranging in size from 2 to 86 interactions). The static network consists of 82 distinct network (static network, 491 proteins connected by 897 compiling all the interactions between static proteins in the neighborhoods also seem to be functionally coherent, it should be possible to assign proteins to specific modules by the virtue of their associations with static neighborhoods. To this end, all static neighborhoods in our network were identified by compiling all the interactions between static proteins in the network (static network, 491 proteins connected by 897 interactions). The static network consists of 82 distinct disconnected sub-networks ranging in size from 2 to 86 proteins (Supplementary Table 1). The functional annotations associated with these static sub-networks appear to be functionally coherent, representing various functions including mRNA transcription and splicing, vesicle transport and cell-cycle regulation (see Supplementary Table 1). This apparent functional coherence suggests that the static network is enriched for functional modules. In order to test the significance of modular composition of the static network and to see if it is possible to achieve a similar level of functional coherence in a network generated by random draws of interactions, a network modularity metric was defined to measure functional specialization of the interactions in a network (see Materials and methods). The static network shows significantly higher network modularity than what would be expected by random draws of interactions from the large network (Figure 3), suggesting that the association of functionally coherent sets of proteins with each other within static neighborhoods reflects a biological phenomenon. We compared the static network modularity with that of the network formed by highly coexpressed proteins, which is expected to be enriched for functional modules, in agreement with the previous studies showing modularity of coexpressed proteins (Segal et al., 2003b; Han et al., 2004). We identified the dynamic network by taking all interacting pairs of proteins that also have pairwise PCCs of at least 0.65 (383 proteins connected by 777 interactions). This dynamic network, therefore, contains interactions between those proteins that are also highly transcriptionally co-regulated. The dynamic network consists of 77 sub-networks mainly composed of dynamic proteins (data not shown) and, as expected from previous publications, the dynamic sub-networks are highly functionally coherent (see Supplementary Table 2). The dynamic network also shows a significantly high network modularity that is comparable to that of the static network (Figure 3). This indicates that both networks are enriched for functional modules. The fact that only 15 proteins and six interactions are common to both networks indicates that the modules in the two networks are distinct, and that the high modularity of the static network is not a consequence of a significant overlap with the dynamic network. These observations argue that the static protein neighborhoods represent functional modules, and it should be possible to assign proteins to functional modules by virtue of their association with static proteins.

In order to see if either network is specifically enriched for certain cellular functions, we performed enrichment analyses of the two networks for overrepresentations of MIPS functional categories (Mewes et al., 2004). Interestingly, the most significant relative enrichment is seen in the functional categories related to mRNA transcription and processing (static network) and rRNA transcription and processing as well as translation (dynamic network) (see Supplementary Table 3). The static network is enriched for general mRNA transcription (RNA polymerase II holoenzyme complexes), splicing (the pre-mRNA splicing complex) and processing (CPF and CCR4-NOT) as well as co-regulator complexes like the chromatin remodeling complexes (SWI/SNF and INO80),

Identification of static and dynamic modules and their functions
Past studies have measured statistical correlation of gene expression in order to assign proteins to specific modules and also to assign new functions to previously uncharacterized proteins (Ihmels et al., 2002; Segal et al., 2003b, c). As static neighborhoods also seem to be functionally coherent, it should be possible to assign proteins to specific modules by the virtue of their associations with static neighborhoods. To this end, all static neighborhoods in our network were identified by compiling all the interactions between static proteins in the network (static network, 491 proteins connected by 897 interactions). The static network consists of 82 distinct disconnected sub-networks ranging in size from 2 to 86 proteins (Supplementary Table 1). The functional annotations associated with these static sub-networks appear to be functionally coherent, representing various functions including mRNA transcription and splicing, vesicle transport and cell-cycle regulation (see Supplementary Table 1). This apparent functional coherence suggests that the static network is enriched for functional modules. In order to test the significance of modular composition of the static network and to see if it is possible to achieve a similar level of functional coherence in a network generated by random draws of interactions, a network modularity metric was defined to measure functional specialization of the interactions in a network (see Materials and methods). The static network shows significantly higher network modularity than what would be expected by random draws of interactions from the large network (Figure 3), suggesting that the association of functionally coherent sets of proteins with each other within static neighborhoods reflects a biological phenomenon. We compared the static network modularity with that of the network formed by highly coexpressed proteins, which is expected to be enriched for functional modules, in agreement with the previous studies showing modularity of coexpressed proteins (Segal et al., 2003b; Han et al., 2004). We identified the dynamic network by taking all interacting pairs of proteins that also have pairwise PCCs of at least 0.65 (383 proteins connected by 777 interactions). This dynamic network, therefore, contains interactions between those proteins that are also highly transcriptionally co-regulated. The dynamic network consists of 77 sub-networks mainly composed of dynamic proteins (data not shown) and, as expected from previous publications, the dynamic sub-networks are highly functionally coherent (see Supplementary Table 2). The dynamic network also shows a significantly high network modularity that is comparable to that of the static network (Figure 3). This indicates that both networks are enriched for functional modules. The fact that only 15 proteins and six interactions are common to both networks indicates that the modules in the two networks are distinct, and that the high modularity of the static network is not a consequence of a significant overlap with the dynamic network. These observations argue that the static protein neighborhoods represent functional modules, and it should be possible to assign proteins to functional modules by virtue of their association with static proteins.

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histone acetyl-transferase complexes (SAGA and NuA4), histone methylase (COMPASS) as well as mRNA nuclear export (TREX) (see Supplementary Table 1). The dynamic network, in addition to RNA polymerase I and III components, contains modules like the SSU processome, involved in rRNA processing, and translation initiation factor complexes (see Supplementary Table 2), which is consistent with studies reporting extensive regulation of these modules under various stress conditions (Warner, 1999; Gasch et al, 2000). In addition, the dynamic network contains most of the proteosomal proteins, whereas the static network also contains many of the mitochondrial ribosomal proteins. There are many modules in the two networks that also seem to perform similar functions. For example, components of the mitotic cohesin complex, which holds sister chromatids together, and the septin ring complex, which is required for cytokinesis, are in the dynamic network (sub-networks 63 and 55; Supplementary Table 2), whereas the DASH complex, which plays a role in chromosome segregation, and the COMA complex, which is involved in the kinetochore assembly, are in the static network (sub-networks 51 and 58; Supplementary Table 1). Components of the anaphase-promoting complex (APC) are also static (sub-network 4), as reported previously (de Lichtenberg et al, 2005). These complexes are all involved in the final stages of cell division, yet their regulation is markedly different. Another potentially interesting correlation relates to vesicle trafficking, where proteins associated with clathrin-coated vesicles (AP-1 and AP-3 complex proteins) seem to be static (sub-networks 9 and 69), whereas those associated with coatomer protein-coated vesicles that are involved in vesicle transport between Golgi and ER (COPI and COPII complex proteins) are dynamic (sub-networks 10, 45 and 76). The dynamic expression pattern of the latter may stem from the involvement of the early secretory pathway in various stress responses like unfolded protein response or osmotic stress (Lee and Linstedt, 1999; Higashio and Kohno, 2002; Sato et al, 2002), whereas clathrin-coated vesicles may play role in constitutive transport. These examples suggest that although some functions in the cell can be classified as static or dynamic (like mRNA and rRNA synthesis, respectively), many others are carried out through dynamic interplay between distinct static and dynamic modules. A closer observation of expression dynamics of functional modules under various conditions may provide an in-depth insight into the regulation of cellular behavior by transcriptional programs.

Expression properties of centrally positioned proteins

A recent study classified hub proteins into two based on their coexpression with their neighbors. They reported that hubs that do not statistically correlate with their neighbors in expression are positioned centrally in the network, meaning that they function between modules as organizers of cellular processes rather than having a specialized function inside a module (Han et al, 2004). However, as shown above, hubs that are found within static neighborhoods also do not statistically correlate with their neighbors in expression even though they are within modules, and therefore are not central. The *bona fide* central hubs, therefore, could be proteins that have low avPCC (i.e. those that do not belong to dynamic modules) and relatively high neighborhood EV (i.e. those that do not belong to static modules). In order to test this hypothesis, we tested the ‘betweenness’ centralities of hub proteins with different avPCC as well as neighborhood EV values. Betweenness centrality is a graph theoretic measure of network centrality that measures how frequently a node is found ‘on the path’ between other nodes in the network, and therefore scores how ‘important’ a node is for communication between other nodes in the network (Wasserman and Faust, 1994) (see Materials and methods). As expected, proteins with low avPCC and relatively high neighborhood EV have high betweenness centralities and low neighborhood densities, strongly suggesting that these hubs are positioned centrally in the network (Figure 4A). Their low neighborhood function homologies, in turn, suggest that they are involved in multifunctional interactions, indicating that they may have roles as integrators of multiple processes in the cell (Figure 4A). Unlike hubs in modules, central hubs and their neighbors have a broad distribution of EV values (Figure 4B), suggesting that true central hubs interact with proteins of diverse expression patterns.

In their study, Han et al (2004) defined hubs that are highly coexpressed with their neighbors as ‘party’ hubs, which are modular, and those that are not coexpressed with their neighbors as ‘date’ hubs, which they reported as central. However, the set of date hubs also contains hubs that are found within static modules (where there is also no statistical correlation of expression among neighbors). Therefore, based on our findings, we propose that static hubs interacting with static proteins within static modules be excluded from date hubs and, in analogy to the party-date hub terminology, be named ‘family’ hubs, as they are always present in the network and interact with their neighbors constitutively. Therefore, family and party hubs form static and dynamic modules, respectively, whereas date hubs (family hubs excluded) organize the network. In concordance with their central functions, our date hubs are enriched for signal transducing and signal regulating proteins like protein kinases, phosphatases, small G-proteins and molecular chaperones (Table 1). Date hubs contain 20 out of 22 hub protein kinases (the two hub kinases excluded from date hubs being YAK1, a kinase in the glucose sensing pathway, and SSN3, a C-terminal kinase in the RNA polymerase holoenzyme complex), five out of six hub phosphatases and all of hub small GTPases in the network, indicating that these proteins constitute the true central coordinators of the cellular network.

Protein-expression noise and evolutionary rate in the static and dynamic networks

Based on the classification of hubs by Han et al (2004), it was suggested that centrally positioned hubs in the network evolve faster than hubs in modules, and that modularity imposes a constraint on the evolvability of proteins, suggesting an evolutionary scenario where protein networks evolve mainly by modifying their central coordinators (Fraser, 2005). We examined this hypothesis in the context of our modified hub
classification, and also found that party hubs evolve at a significantly slower rate than other hubs (Figure 5A). However, surprisingly, family hubs do not evolve slower than date hubs (Figure 5A). By extrapolation, this suggests that hubs present in dynamic modules are evolutionarily constrained, whereas those present in static modules are not. Accordingly, proteins in the dynamic network have significantly lower evolutionary rates than proteins in the static network ($P<1 \times 10^{-16}$, Wilcoxon’s test), and there is a significant negative correlation between EV values of proteins and their rates of evolution (Spearman’s $\rho=-0.21$, $P=4.5 \times 10^{-15}$). These results suggest that proteins in static modules have more freedom of variation than proteins within dynamic modules.

![Image of Figure 4](https://example.com/figure4.png)

**Figure 4** Distinct neighborhood EV and avPCC characteristics of central and modular hubs. (A) Plots of hubs' avPCC versus neighborhood EV; each dot (hub) is colored according to the value of the corresponding measure. (B) Hubs are colored according to the neighborhood EV variation, which is the standard deviation of the neighbors' EV values of a protein and shows how diverse the EVs of proteins in the neighborhood of a protein are.

| MIPS protein classes                        | Number in date hubs | Total number in hubs | P-value |
|----------------------------------------------|---------------------|----------------------|---------|
| Small GTP-binding proteins (RAS superfamily)| 4                   | 4                    | 0       |
| Molecular chaperones                         | 3                   | 4                    | 0.02    |
| Protein kinases                              | 20                  | 22                   | <1.41E-07 |
| CaMK group                                   | 2                   | 2                    | 0       |
| CMGC group                                   | 7                   | 9                    | 0.003   |
| OPG group                                    | 7                   | 8                    | 0.0005  |
| Unique Saccharomyces cerevisiae kinases      | 3                   | 3                    | 0       |
| Protein phosphatases                         | 5                   | 6                    | 0.003   |
| Ubiquitin system proteins                    | 3                   | 3                    | 0       |

Date hubs were defined as hubs that have neighborhood EVs greater than 0.3 and avPCC less than 0.45. For protein classes, we used the MIPS (Mewes et al., 2004) protein classification catalogue. Only the classes with most significant P-values are shown. P-values of enrichment were calculated using the cumulative hypergeometric distribution function.
robustness in cell regulatory networks that allows for the eukaryotic protein interaction network are a source of evolution. These observations argue that static components of expression patterns are most variable between cells and during surprising to find that proteins with least variable mRNA variations within a population. Accordingly, family hubs have levels of static proteins are the ones that show most cell-to-cell and date hubs are those with neighborhood EVs of $<0.3$. party hubs are hubs with avPCC $>0.45$. (B) Fractions of proteins in the static and dynamic networks whose gene deletion is lethal to yeast. (C) Boxplot of protein expression noise in the different hub classes.

Figure 5  Evolutionary rate and expression noise of the static and dynamic modules. Evolutionary rates of yeast proteins derived by Hirsh et al (2005) were used. (A) Boxplot of evolutionary rates of family, party and date hubs. Family hubs are static hubs with neighborhood EVs of $<0.3$, party hubs are hubs with avPCC $>0.45$ and date hubs are those with neighborhood EV $>0.3$ and avPCC $<0.45$. (B) Fractions of proteins in the static and dynamic modules. Using the coefficients of variation of protein expression when compared to proteins in static modules (Figure 5B), which is also true for party hubs when compared to family hubs (data not shown), indicating that the cell is highly tolerant of the loss of proteins in static modules, a property that may allow them to evolve at a faster rate than proteins in dynamic modules.

The significant correlation of protein EVs with their evolutionary rates suggests that the static network may be a buffer of evolutionary variations in the protein interaction network, granting static proteins a role as evolutionary modifiers of cell behavior. We reasoned that if the cell is more tolerant of genetic variations in the components of static modules, then the cell may also be more tolerant of variations in the expression of these proteins within a cell population. Expression variation of proteins between cells within a population, or protein expression noise, is a major factor contributing to the variations of cell behavior among cells within a cell population (Blake et al, 2003; Raser and O’Shea, 2005). Therefore, we compared the expression noise of proteins in static modules with that of proteins in dynamic modules. Using the coefficients of variation of protein expression levels (CV values) within a clonal cell population derived by a recent study (Newman et al, 2006), we found that proteins in dynamic modules are significantly less noisy in their expression when compared to proteins in static modules ($P=3 \times 10^{-15}$, Wilcoxon’s test), indicating that the expression levels of static proteins are the ones that show most cell-to-cell variations within a population. Accordingly, family hubs have significantly higher CV values than other hubs (Figure 5C). It is surprising to find that proteins with least variable mRNA expression patterns are most variable between cells and during evolution. These observations argue that static components of the eukaryotic protein interaction network are a source of robustness in cell regulatory networks that allows for evolutionary as well as populational variations in cell behavior (see Discussion).

Expression levels of static and dynamic modules

EV of genes positively correlates with their mRNA abundance (Spearman’s $\rho=0.21$, $P<1 \times 10^{-16}$), and accordingly, proteins in static modules are expressed at a significantly lower level than those in dynamic modules (Wilcoxon’s test, $P<1 \times 10^{-19}$). This may suggest that the correlation of EV with the organizational layout in the protein interaction network may be a reflection of the effect of expression levels of proteins rather than their EV. The expression levels of proteins does seem to contribute to the protein network layout, as there is a high positive correlation between mRNA abundance values of hub proteins and that of their neighbors in the network (i.e. average neighborhood mRNA abundance, Spearman’s $\rho=0.43$), although the correlation is significantly less than that between EV and neighborhood EV (Spearman’s $\rho=0.61$). This correlation is not surprising given that the expression levels of proteins participating in the same protein complex are generally similar (Papp et al, 2003). The relatively low correlation between EV and mRNA abundance and the fact that the correlation of mRNA abundance between neighboring proteins is less than that of EV suggests that our observations with EV above are not an artifact of the underlying mRNA abundance values. In order to rule out the possibility that our observations with EV values of proteins presented above are an artifact of their expression levels, we performed partial correlation analyses (see Materials and methods). Between EV, neighborhood EV, neighborhood function homology and mRNA abundance values of proteins. Partial correlation between EV and neighborhood EV while controlling for mRNA abundance is almost as high ($r_{EV \sim \text{neigh.EV}, \text{mRNA}}=0.66$) as their normal correlation ($r_{EV \sim \text{neigh.EV}}=0.67$). Similarly, partial correlation between neighborhood function homologies of static proteins with their neighborhood EV while controlling for mRNA abundance or average neighborhood mRNA abundance is almost as high as their normal correlations (data not shown). These observations argue that the observed
static and dynamic modules in the protein network
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Discussion
A proper stoichiometry in the expression levels of components of a module is essential as an imbalance in the levels of the module constituents can be deleterious (balance hypothesis) (Papp et al., 2003). A priori, there are two simple ways to control stoichiometry of module components at the level of transcription: by maintaining constant expression and by co-regulated expression of all components. Both mechanisms are apparently employed for the design of cellular modules, leading to an organizational model of the network resembling a circuit board with integrated ‘built-in’ as well as removable ‘plug-and-play’ components. For example, the functionally ubiquitous process of mRNA synthesis and splicing is carried out by proteins organized in modules with apparent invariant expression. The highly dynamic nature of ribosome biogenesis modules, on the other hand, has been suggested to be a mechanism of energy preservation for the cell under stress, as transcription of ribosomal genes accounts for around 80% of all RNA synthesis in the cell (Warner, 1999).

Although the expression variations of modular proteins are constrained by those of their neighbors, central proteins, which are versatile in their functions, are also more versatile in their expression patterns. The existence of both static and dynamic central hubs, which are presumably the coordinators of cellular processes, suggests that some connections between processes in the cell are ‘hard-wired’, whereas some are adjustable depending on the cellular requirements. For example, the sub-network 2 in our static network (Supplementary Table 1) indicates that the TFIID/SAGA complex is hard wired to the nuclear proteasomal complex (Supplementary Figure 2), suggesting an integral function of the proteasome in sequence-specific transcription, consistent with previous reports (Lee et al., 2005; Auld et al., 2006). This sub-network also indicates an integral connection of vesicle trafficking with general mRNA synthesis, a relationship that to our knowledge has not yet been explored. Therefore, in addition to revealing some novel architectural characteristics of the protein network, the analysis employed in this study also helps reveal how local dynamics of the network architecture may shape cell behavior.

The faster evolutionary rate and higher expression noise in static modules suggests that robustness to variations in these modules may largely enhance cell-to-cell variations within a population and consequently increase robustness of the population to environmental fluctuations. Similarly, genetic variations in static modules during evolution may result in the phenotypic enhancement of other mutations in the cell, which may facilitate adaptation. As mRNA abundance is a major factor contributing to protein expression noise (Newman et al., 2006) and evolutionary rate (Pal et al., 2001), it is conceivable that relatively lower expression levels of static modules is an evolutionarily selected trait to maximize variations in these modules.

A future comparison of the expression dynamics of the protein network of yeast with that of higher eukaryotes should give an insight into the evolution of expression dynamics in concordance with the evolution of protein network connectivity and robustness.

Materials and methods
Microarray data sets
The microarray gene expression data sets from various conditions (cell cycle, sporulation, stress response, unfolded protein response and diauxic shift) were obtained from the Saccharomyces Genome Database (ftp://ftp.yeastgenome.org/yeast/). In order for the data to account for true fold differences in the expression of genes relative to the control (i.e. 0’ time point), 0’ time points were removed from the data sets, and the corresponding later time points were zero-transformed by subtracting the expression values at these time points from those at the 0 time point.

Protein interaction network
The protein interaction network was obtained using the confidence scores assigned to potential interactions in the study of Bader et al. (2004). Following the original study (Bader et al., 2004), a high cutoff of 0.65 was used to obtain a high confidence network. The giant connected component of the network was used in the analyses.

Neighborhood PCC
nPCC of a protein is defined as the average of all pairwise PCCs between all the proteins in its neighborhood including itself;

\[
nPCC_a = \frac{\sum_{i=1}^{n} PCC_{ij}}{n^2 - n}
\]

where nPCC_a is the nPCC of a protein a, n is the node degree of protein a plus 1 (for itself) and PCC_{ij} is the PCC between proteins i and j.

Neighborhood function homology
Let G_i be the set of GO terms assigned to protein i that has a node degree of k. Neighborhood function homology F_i of the protein i is defined as

\[
F_i = \frac{\sum_{j=1}^{k} |G_i \cap G_j|}{|G_i|}
\]

where G_i is the set of GO terms assigned to the jth neighbor of protein i. F_i ranges from 0, where there are no shared GO terms between protein i and its neighbors, to 1, where all GO terms assigned to the protein i are also present in all of its neighbors.

Average interactor PCC
Following Han et al. (2004), we defined avPCC as the average of pairwise PCC values between a protein and its neighbors. Differently
from npCC, which is a measure of how the proteins in the neighborhood are coexpressed, avpCC measures how a protein is coexpressed with its neighbors.

**Neighborhood density**

Neighborhood density of a protein is its clustering coefficient. Clustering coefficient \( C_C \) of a protein \( i \) is defined as

\[
C_C = \frac{2N}{k(k+1)}
\]

where \( N \) is the number of interactions between the neighbors of protein \( i \), and \( k \) is its node degree.

**Network modularity**

First, a function similarity matrix was constructed by measuring all pairwise function similarities between proteins in the network. The pairwise function similarity between proteins \( i \) and \( j \) was defined as

\[
S_{ij} = \frac{|G_i \cap G_j|}{|G_i \cup G_j|}
\]

where \( G_i \) and \( G_j \) are the sets of GO terms assigned to proteins \( i \) and \( j \), respectively. For a network of \( n \) proteins, the function similarity matrix \( S \) would be a matrix of dimensions \( n \times n \). The network modularity \( M \) is calculated by summing the pairwise function similarities between every interacting pair of proteins in this network and dividing by the total number of interactions in the same network:

\[
M = \frac{\sum_{i \neq j} A_{ij} S_{ij}}{\sum_{i \neq j} A_{ij}}
\]

where \( A \) is the adjacency matrix of the network and has the same dimensions as \( S \). \( A \) is boolean, \( A_{ij} \) being 1 only if proteins \( i \) and \( j \) interact, and 0 otherwise.

**Betweenness centrality**

Betweenness centrality of a node \( i \) in the network, shown as \( C_B(i) \), is given by

\[
C_B(i) = \frac{1}{|G|} \sum_{j \neq k} \frac{g_{jk}(i)}{g_{jk}}
\]

where \( g_{jk}(i) \) is the number of shortest paths between nodes \( j \) and \( k \) that pass through node \( i \) and \( g_{jk} \) is the total number of shortest paths connecting nodes \( j \) and \( k \) (Wasserman and Faust, 1994).

**Protein-expression noise values**

For protein-expression noise values, we used the values derived by a large-scale single-cell proteomic analysis of Newman et al (2006). They defined protein-expression noise as CV (s.d. divided by mean expression) of protein expression between cells in a population.

**Partial correlation analysis**

Linear correlation between two variables \( a \) and \( b \) is given by

\[
r_{a-b} = \frac{\text{cov}(a,b)}{\sqrt{\text{var}(a) \text{var}(b)}}
\]

where \( \text{cov}(a,b) \) is covariance between \( a \) and \( b \), and \( \text{var}(a) \) is variance of \( a \). Partial correlation between \( a \) and \( b \) while controlling for a variable \( c \) is given by

\[
r_{a-b|c} = \frac{r_{a-b} - r_{a-c} r_{b-c}}{\sqrt{(1 - r_{a-c}^2)(1 - r_{b-c}^2)}}
\]

Equations were taken from (de la Fuente et al, 2004).

**Supplementary information**

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

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**References**

Albert R, Jeong H, Barabasi AL (2000) Error and attack tolerance of complex networks. Nature 406: 378–382

Ashburner M, Ball CA, Blake JA, Botstein D, Butler H, Cherry JM, Davis AP, Dolinski K, Dwight SS, Eppig JT, Harris MA, Hill DP, Issel-Tarver L, Kasarskis A, Lewis S, Matese JC, Richardson JE, Ringwald M, Rubin GM, Sherlock G (2000) Genome ontology: tool for the unification of biology. The Gene Ontology Consortium. Nat Genet 25: 25–29

Auld KL, Brown CR, Casolari JM, Komili S, Silver PA (2006) Genomic association of the proteasome demonstrates overlapping gene regulatory activity with transcription factor substrates. Mol Cell 21: 861–871

Bader JS, Chaudhuri A, Rothberg JM, Chant J (2004) Gaining confidence in high-throughput protein interaction networks. Nat Biotechnol 22: 78–85

Blake WJ, MKA, Cantor CR, Collins JJ (2003) Noise in eukaryotic gene expression. Nature 422: 633–637

de la Fuente A, Bing N, Hoeschele I, Mendes P (2004) Discovery of meaningful associations in genomic data using partial correlation coefficients. Bioinformatics 20: 3565–3574

de Lichtenberg U, Jensen LJ, Brunak S, Bork P (2005) Dynamic complex formation during the yeast cell cycle. Science 307: 724–727

Fraser HB (2005) Modularity and evolutionary constraint on proteins. Nat Genet 37: 351–352

Gasch AP, Spellman PT, Kao CM, Carmel-Harel O, Eisen MB, Storz G, Botstein D, Brown PO (2000) Genomic expression programs in the response of yeast cells to environmental changes. Mol Biol Cell 11: 4241–4257

Gavin AC, Aloy P, Grandi P, Krause R, Boesche M, Marzioch M, Rau C, Jensen LJ, Bastuck S, Dumplitz B, Edelmann A, Heurter MA, Hofmann V, Hoefert C, Klein K, Hudak M, Michon AM, Schelder M, Schirle M, Remor M, Rudi T, Hooper S, Bauer A, Bouwmeester T, Casari G, Drewes G, Neuberger G, Rick CM, Kuster B, Bork P, Russell RB, Superti-Furga G (2006) Proteome survey reveals modularity of the yeast cell machinery. Nature 440: 631–636

Ge H, Liu Z, Church GM, Vidal M (2001) Correlation between transcriptome and interactome mapping data from Saccharomyces cerevisiae. Nat Genet 29: 482–486

Han JD, Bertin N, Hao T, Goldberg DS, Berriz GF, Zhang LV, Dupuy D, Walhout AJ, Cusick ME, Roth FP, Vidal M (2004) Evidence for dynamically organized modularity in the yeast protein–protein interaction network. Nature 430: 88–93

HARBISON CT, Gordon DB, Lee TI, Rinaldi NJ, Macisaac KD, Danford TW, Hannett NM, Tagne JB, Reynolds DB, Yoo J, Jennings EG, Zeitlinger J, Pokholok DK, Kellis M, Rolfe PA, Takusagawa KT, Lander ES, Gifford DK, Fraenkel E, Young RA (2004) Transcriptional regulatory code of a eukaryotic genome. Nature 431: 99–104

Higashio H, Kohno K (2002) A genetic link between the unfolded protein response and vesicle formation from the endoplasmic reticulum. Biochem Biophys Res Commun 296: 568–574

Hirsh AE, Fraser HB, Wall DP (2005) Adjusting for selection on synonymous sites in estimates of evolutionary distance. Mol Biol Evol 22: 174–177

Ihmels J, Friedlander G, Bergmann S, Sarig O, Ziv Y, Barkai N (2002) Revealing modular organization in the yeast transcriptional network. Nat Genet 31: 370–377
Ihmels J, Levy R, Barkai N (2004) Principles of transcriptional control in the metabolic network of Saccharomyces cerevisiae. Nat Biotechnol 22: 86–92

Jeong H, Mason SP, Barabasi AL, Oltvai ZN (2001) Lethality and centrality in protein networks. Nature 411: 41–42

Jeong H, Tombor B, Albert R, Oltvai ZN, Barabasi AL (2000) The large-scale organization of metabolic networks. Nature 407: 651–654

Kærn M, Elston TC, Blake WJ, Collins JJ (2005) Stochasticity in gene expression: from theories to phenotypes. Nat Rev Genet 6: 451–464

Kharchenko P, Church GM, Vitiuk D (2005) Expression dynamics of a cellular metabolic network. Mol Syst Biol 1: 2005.0016

Klemm K, Bornholdt S (2005) Topology of biological networks and reliability of information processing. Proc Natl Acad Sci USA 102: 18414–18419

Krogan NJ, Cagney G, Yu H, Zhong G, Guo X, Ignatchenko A, Li J, Pu S, Datta N, Tikuisis AP, Punna T, Peregrin-Alvarez JM, Shales M, Zhang X, Davey M, Robinson MD, Paczianar A, Bray JE, Sheung A, Beattie B, Richards DP, Canadien V, Lalov A, Mena F, Wong P, Starostine A, Canete MM, Vlasblom J, Wu S, Orsi C, Collins SR, Chandran S, Haw R, Rilstone J, Gandi K, Thompson NJ, Musso G, St Onge P, Channy S, Lam MH, Butland G, Altaf-Ul AM, Kanaya S, Shilatifard A, O’Shea E, Weissman JS, Ingles CJ, Hughes TR, Parkinson J, Gerstein M, Wodak SJ, Emili A, Greenblatt JF (2006) Global landscape of protein complexes in the yeast Saccharomyces cerevisiae. Nature 440: 637–643

Lee D, Ezhkova E, Li B, Pattenden SG, Tansey WP, Workman JL (2005) The proteasome regulatory particle alters the SAGA coactivator to enhance its interactions with transcriptional activators. Cell 123: 423–436

Lee TH, Linstedt AD (1999) Osmotically induced cell volume changes alter anterograde and retrograde transport, Golgi structure, and COPI dissociation. Mol Biol Cell 10: 1445–1462

Lehner B, Crombie C, Tischler J, Fortunato A, Fraser AG (2006) Systematic mapping of genetic interactions in Caenorhabditis elegans identifies common modifiers of diverse signaling pathways. Nat Genet 38: 896–903

Luscombe NM, Babu MM, Yu H, Snyder M, Teichmann SA, Gerstein M (2004) Genomic analysis of regulatory network dynamics reveals large topological changes. Nature 431: 308–312

Maslov S, Sneppen K (2002) Specificity and stability in topology of protein networks. Science 296: 910–913

Mewes HW, Amid C, Arnold R, Frishman D, Guldener U, Mannhaupt G, Munsterkotter M, Pagel P, Strack N, Stumpflen V, Warfmann J, Ruepp A (2004) MIPS: analysis and annotation of proteins from whole genomes. Nucleic Acids Res 32: D41–D44

Milos R, Shen-Ors S, Itzkovitz S, Kashtan N, Chklovskii D, Alon U (2002) Network motifs: simple building blocks of complex networks. Science 298: 824–827

Newman JR, Ghaemmaghami S, Ihmels J, Breslow DK, Noble M, DeRisi JL, Weissman JS (2006) Single-cell proteomic analysis of S. cerevisiae reveals the architecture of biological noise. Nature 441: 840–846

Pal C, Papp B, Hurst LD (2001) Highly expressed genes in yeast evolve slowly. Genetics 158: 927–931

Papp B, Pal C, Hurst LD (2003) Dosage sensitivity and the evolution of gene families in yeast. Nature 424: 194–197

Pereira-Leal JB, Enright AJ, Ouzounis CA (2004) Detection of functional modules from protein interaction networks. Proteins 54: 49–57

Raser JM, O’Shea EK (2005) Noise in gene expression: origins, consequences, and control. Science 309: 2010–2013

Ravasz E, Somera AL, Mongru DA, Oltvai ZN, Barabasi AL (2002) Hierarchical organization of modularity in metabolic networks. Science 297: 1551–1555

Sato M, Sato K, Nakano A (2002) Evidence for the intimate relationship between vesicle budding from the ER and the unfolded protein response. Biochem Biophys Res Commun 296: 560–567

Segal E, Shapira M, Regev A, Pe’er D, Botstein D, Coller H, Friedman N (2003a) module networks: identifying regulatory modules and their condition-specific regulators from gene expression data. Nat Genet 34: 166–176

Segal E, Wang H, Koller D (2003b) Discovering molecular pathways from protein interaction and gene expression data. Bioinformatics 19 (Suppl 1): i264–i271

Segal E, Yelensky R, Koller D (2003c) Genome-wide discovery of transcriptional modules from DNA sequence and gene expression. Bioinformatics 19 (Suppl 1): i273–i282

Spirin V, Mirny LA (2003) Protein complexes and functional modules from protein interaction networks. Proteins 431: 437–440