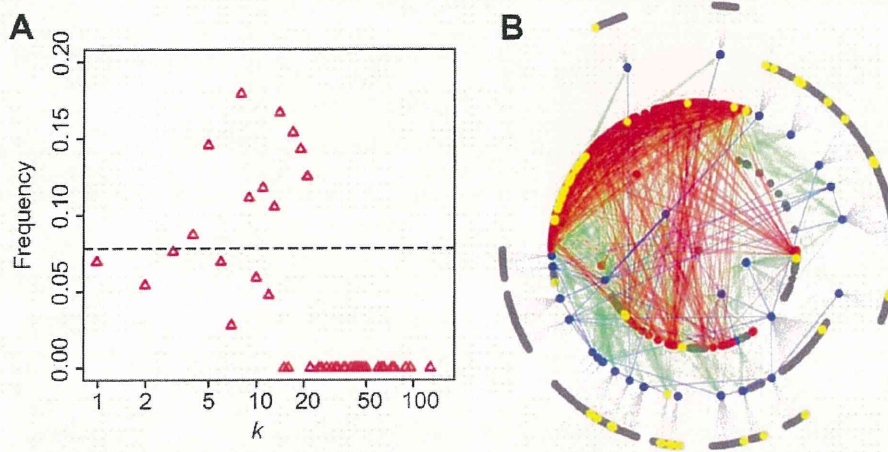


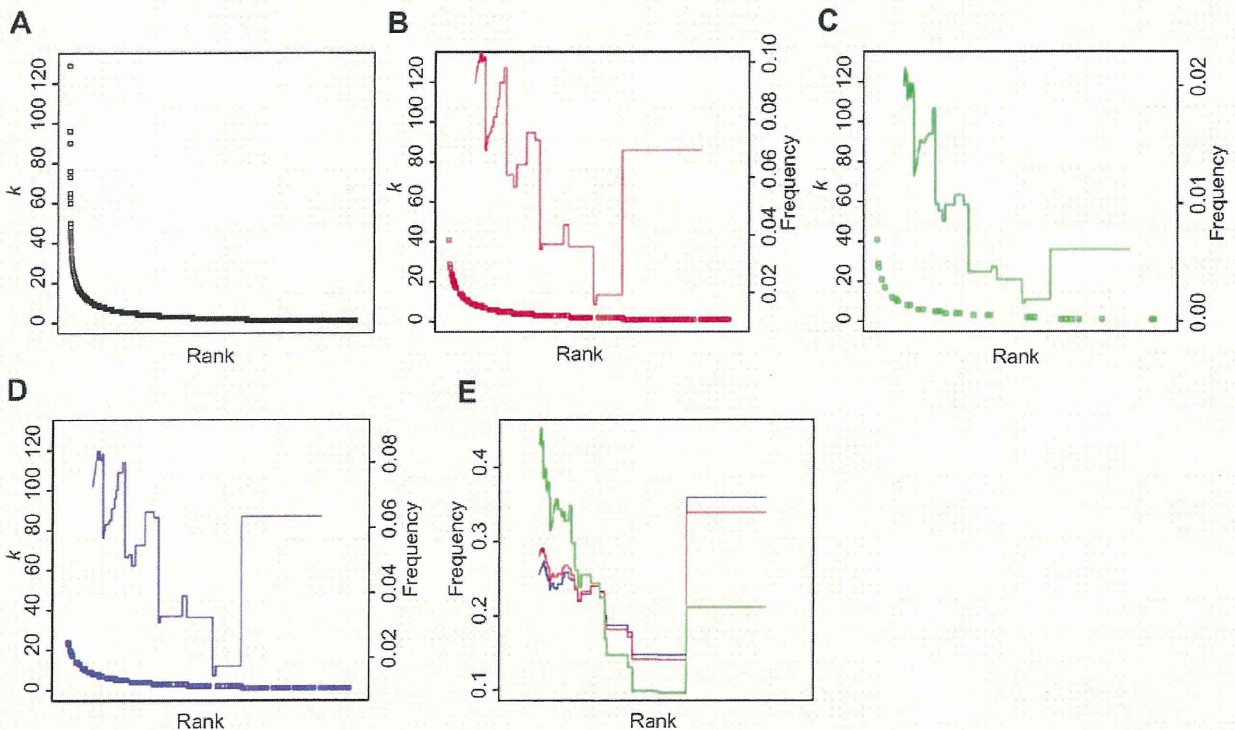
**Figure 4. Scale-richness in yeast PIN.** Each diagram shows cumulative degree distributions of proteins in each functional group. The name above each diagram denotes the name of the functional category with which the cumulative degree distribution was examined.  
doi:10.1371/journal.pcbi.1000550.g004



**Figure 5. Distribution of drug targets.** (A) Degree distribution. Red triangles represent fraction of drug-target proteins to all proteins with degree  $k$ . The dashed line in black gives the probability that a randomly selected protein is a drug target. (B) Distribution on network topology. Drugs targets (yellow circles) are mapped on human PIN network topology shown in Figure 2D. doi:10.1371/journal.pcbi.1000550.g005

robustness against node failure and optimized performance [21]. It has been reported that analysis using several possible router topologies found that a HOTnet configuration was most efficient,

providing more maximum overall bandwidth to users than that with other network-configuration approaches such as random and preferential attachment [21]. The implication is that biological



**Figure 6. The long tails in degree distribution of drug targets, targets for cancer, and those for non-cancerous diseases.** Proteins were ranked in decreasing order of their degree  $k$ . (A) Rank of a protein with degree  $k$ . (B) Rank of a drug target with degree  $k$ . (C) Rank of a target for cancer diseases with degree  $k$ . (D) Rank of a target for non-cancerous diseases with degree  $k$ . Red, green, and blue lines represent fraction of drug targets, targets for cancer diseases, and those for non-cancerous diseases to all proteins with rank from rank $-0.1N$  to rank $+0.1N$  ( $N$  represents number of proteins in the human PIN). (E) Red, green, and blue lines represent fraction of drug targets, targets for cancer disease, and those for non-cancerous disease with rank from rank $-0.1N$  to rank $+0.1N$  to all drug targets, all targets for cancer diseases, and those for non-cancerous diseases, respectively. doi:10.1371/journal.pcbi.1000550.g006

PINs have evolved to become efficient and error tolerant. The series of analyses presented in this report indicate that there are changes whereby we can rationally design drugs by taking into account network properties, and additional insights from engineering and physics may further extend our opportunities for exploring network-based biology.

## Materials and Methods

**PINs, GO data, and essential genes.** Yeast PIN data were obtained from the Munich Information Center for Protein Sequences (MIPS) database (<http://mips.gsf.de>) [22] and human PIN data were obtained from Rual et al. [23]. The yeast (or human) PIN contained 4,153 (or 3,023) proteins and 7,417 (or 6,149) non-redundant interactions. The GO slim dataset for the yeast PIN was from the ftp site of the Saccharomyces Genome Database (SGD) ([ftp://genome-ftp.stanford.edu/pub/yeast/literature\\_curation/](ftp://genome-ftp.stanford.edu/pub/yeast/literature_curation/)) [22] and that for the human PIN was from the European Bioinformatics Institute (EBI) (<ftp://ftp.ebi.ac.uk/pub/databases/GO/goa/HUMAN/>). The list of essential genes from SGD [22] contained 889 essential genes that were mapped to the yeast PIN.

**Synthetic lethal and synthetic sick proteins.** We obtained a list of synthetic lethal and sick interactions from Tong et al. [24]. There were 735 proteins having at least one synthetic lethal interaction and we defined these proteins as synthetic lethal proteins. However, there were 816 proteins having at least one synthetic sick interaction, of which 310 proteins had no synthetic lethal interactions. We defined these 310 proteins as synthetic sick proteins. 538 synthetic lethal proteins and 209 synthetic sick proteins were mapped to the yeast PIN.

**Drug-target proteins.** To analyze the statistical features of drug-target genes, we obtained a list of proteins that were targets of FDA-approved and experimental drugs from Yildirim et al. [3]. This list contained 1,013 drug-target proteins, of which 236 were mapped to the human PIN. To generate a list of drug-disease associations, we mapped drugs to diseases by investigating information on drugs obtained from the DrugBank database [25] (information on drugs is contained in the “indications” field in the DrugBank database). Then, by using the list of drug-disease associations, we divided drug-target proteins into two groups: target proteins for cancer drugs and those for non-cancerous diseases. The human PIN contained 33 target proteins for cancer and 203 for non-cancerous diseases.

**Random network.** We generated a random network by using the method proposed by Maslov and Sneppen [13], where the following procedures were performed. First, two links in a network were chosen randomly. Assume that one link connects nodes A and B, and the other connects nodes C and D. Second, these links were rewired by exchanging their connecting partners. That is, nodes A and D were connected, and nodes B and C were connected. We repeated these two procedures  $1,000E$  times ( $E$  is the number of interactions in the original network) to generate a random network. Note that the method did not alter the degree distribution.

**Cluster coefficient.** The cluster coefficient of node  $i$  is defined as  $C_i = 2e_i/k_i(k_i - 1)$ , where  $k_i$  is the degree of node  $i$  and  $e_i$  is the number of links connecting  $k_i$  neighbors of node  $i$  to one another [26]. When  $k_i$  is zero or one,  $C_i$  is defined as zero.  $C_i$  is equal to one when all neighbors of node  $i$  are fully connected to one another, while  $C_i$  is zero when none of the neighbors are connected to one another.

**Moving stratification by degree.** A method of analysis termed moving stratification by degree (MSD) was developed and

used to compare three networks, the budding yeast PIN, the human PIN, and a randomly generated network with exactly the same degree distribution as the PINs. Hubs were defined as nodes with degrees of more than six [10,11]. MSD was used to extract sub-networks consisting of hubs with degrees from  $k_c - \mu$  to  $k_c + \mu$ . In this study, we used  $\mu = 1, 3, 5, \text{ and } 7$ . Since a hub is defined as a node with degrees of more than 6, we used initial values of  $k_c = 7, 9, 11, \text{ and } 13$ . Then,  $k_c$  was scanned up to 300 with step size 1. For each initial value ( $k_c = 7, 9, 11, \text{ and } 13$ ), MSD extracted 293, 291, 289, and 287 sub-networks, respectively. For these sub-networks, only hub nodes were included. In the following analysis, data from  $\mu = 5$  were used because changing  $\mu$  did not significantly alter the results.

The average cluster coefficient  $\langle C(k_c) \rangle$ , average shortest path length  $\langle L(k_c) \rangle$ , betweenness centrality  $B_i(k_c)$ , and node ratio included in largest components  $G_c(k_c)$  in each sub-network from the PINs were compared with each value from random networks. The sub-networks were tightly connected when the average cluster coefficient was high. While there were no significant differences in the average cluster coefficient between the PINs and random networks for high  $k_c$  ( $k_c > 38$  for the yeast PIN and  $k_c > 30$  for the human PIN) (Figures S1A and S2A), the average cluster coefficient for PINs was significantly higher than that for the random networks. There were no significant differences in  $\langle L(k_c) \rangle$  and  $G_c(k_c)$  between the PINs and random networks (Figures S1B, S1C, S2B, and S2C). It is interesting to note that there were no significant differences in global properties (i.e., betweenness centrality  $B_i(k_c)$ ) between PINs and random networks (see Figures S1D and S2D), although difference in local properties (i.e., average cluster coefficient  $\langle C(k_c) \rangle$ ) were significant between PINs and random networks (see Figures S1A and S2A).

The fraction of essential proteins to all proteins in each sub-network ( $F_{LC}(k_c)$ ) was investigated for the budding yeast PIN (Figure S1E). The fraction of drug targets to all proteins in each sub-network ( $F_{DT}(k_c)$ ) was investigated for the human PIN (Figure S2E).

There were no known drug-target proteins when  $k_c$  was over 50 ( $F_{DT}(k_c) = 0$  for  $k_c > 50$ ). See Table S4). This means that high-degree proteins were unlikely to be drug targets. However,  $F_{DT}(k_c)$  is significantly higher than random expectation when  $k_c$  is between 11 and 32. Thus, middle-degree proteins are biologically important and can be drug-target proteins. Table S7 lists middle-degree proteins and their functions categorized by GO annotation. We can expect novel drug targets to be included in the list.

Further analyses were carried out by partitioning a network into three sub-networks, a sub-network consisting of low-degree nodes (degrees from 1 to 5), that consisting of middle-level degree nodes (degrees from 6 to 38 for the yeast PIN and from 6 to 30 for the human PIN), and that consisting of high-degree nodes (degrees more than 39 for the yeast PIN and more than 31 for the human PIN). Middle-level nodes formed a tightly coupled stratus structure whereas high-degree nodes formed a modularized altocumulus structure.

Tables S1 and S2 show that middle-degree nodes formed a high-density tightly coupled structure and a middle-degree sub-network had higher average cluster coefficients than other sub-networks. The average cluster coefficient of PINs without nodes in the middle-degree sub-network was substantially lower than that of the original PIN. In addition, the average shortest distance in the middle-degree sub-network was almost equal to that of the entire PINs. Most nodes in the entire PINs or middle-degree sub-network (over 95% of nodes) were included in the largest component. However, this is not a case for low-degree or high-degree sub-

networks. Thus, the characteristics of middle-degree sub-networks strongly influence the statistical characteristics of the whole PIN. The whole network architecture seems to have tightly connected middle-degree nodes that are connected to high-degree nodes, and a large number of low-degree nodes are mostly connected to high-degree nodes (see Figure 2). Moreover, we used more stringent thresholds for middle- and high-degree nodes and found that changing the thresholds did not essentially affect the results (i.e., the average cluster coefficient, average shortest path length, or  $G_C$ ) (see Tables S3 and S4).

The series of analyses thus far indicates that the functional role for proteins included in low-degree, middle-degree, and high-degree sub-networks are totally different. This means that the yeast and human PINs are not scale-free in terms of the composition of the functional role of proteins. Proteins with each functional group have a characteristic degree distribution. To investigate the degree distribution of proteins in each functional category, we annotated proteins in the yeast and human PINs by using the GO slim biological process ontology. As shown in Figures 4 and S5, there are different degree-distribution patterns for proteins from different functional categories. This suggests that a scale-free distribution emerges from the composition of different functional protein groups each of which has scale-dependent degree distributions. Thus, from the functional distribution, the yeast and human PINs are scale-rich.

**$S(g)$  value.** Before giving a definition for the  $S(g)$  value, let us first define some notations. Let  $n$  be the number of nodes in a network and  $k_i$  be the degree of node  $i$ .  $D = \{k_1, k_2, \dots, k_n\}$  represents a given degree distribution and  $G(D)$  denotes the set of all connected networks having the same degree distribution,  $D$ . For a network,  $g$ , having degree distribution  $D$ , graph-theoretic quantity  $s(g)$  is defined as  $s(g) = \sum_{(i,j) \in E(g)} k_i k_j$ , where  $E(g)$  is the set of links in the network.  $s_{\max}$  is defined as  $s_{\max} = \max\{s(g): g \in G(D)\}$  and we calculated the value of  $s_{\max}$  by using the algorithm devised by Alderson et al. [7].  $S(g)$ , the value normalized against  $s_{\max}$ , is defined as  $S(g) = s(g)/s_{\max}$  [8]. In this paper, we calculated the value of  $S(g)$  in the yeast and human PINs.

## Supporting Information

**Figure S1** Statistics of sub-networks generated by MSD (yeast PIN). Red triangles and black squares show the values for the yeast PIN and random network, respectively. The results for random network were obtained by taking the average among 100 random networks. (A) Distribution of  $\langle C(kC) \rangle$ . (B) Distribution of  $\langle L(kC) \rangle$ . (C) Distribution of  $G_C(kC)$ . (D) Distribution of  $PLC(kC)$ . The dashed line represents the probability that a randomly selected protein is a lethal protein.

Found at: doi:10.1371/journal.pcbi.1000550.s001 (1.37 MB TIF)

**Figure S2** Statistics of sub-networks generated by MSD (human PIN). Red triangles and black squares show the values for the human PIN and random network, respectively. The results for random network were obtained by taking the average among 100 random networks. (A) Distribution of  $\langle C(kC) \rangle$ . (B) Distribution of  $\langle L(kC) \rangle$ . (C) Distribution of  $G_C(kC)$ . (D) Distribution of  $PDT(kC)$ . The dashed line in black represents the probability that a randomly selected protein is a drug target.

Found at: doi:10.1371/journal.pcbi.1000550.s002 (1.35 MB TIF)

**Figure S3** Degree Dependent Connectivity Chart with stringent thresholds.  $P_n(k)$  gives the probability that a link of a  $k$ -degree node is a link to a node in each sub-network of the yeast (left) and human (right) PINs. The value of  $P_n(k)$  is calculated for a sub-network consisting of high-degree nodes, that consisting of middle-

degree nodes, and that consisting of low-degree nodes. (A) Distribution of  $P_n(k)$  for the high-degree sub-network. (B) Distribution of  $P_n(k)$  for the middle-degree sub-network. (C) Distribution of  $P_n(k)$  for the low-degree sub-network.

Found at: doi:10.1371/journal.pcbi.1000550.s003 (0.26 MB TIF)

**Figure S4** Cloud topologies in yeast and human PINs with stringent thresholds. Grey, red, and blue nodes correspond to low-, middle-, and high-degree nodes. Grey, red, green, and blue links correspond to links between low- and high-degree nodes, those between middle-degree nodes, those between middle- and high-degree nodes, and those between high-degree nodes. For clarity, low- and middle-degree nodes that have no links to high-degree nodes have been omitted. (A) Altocumulus and stratus structures in the yeast PIN. (B) Stratus structure in the yeast PIN. (C) Altocumulus structure in the yeast PIN. (D) Altocumulus and stratus structure in the human PIN. (E) Stratus structure in the human PIN. (F) Altocumulus structure in the human PIN.

Found at: doi:10.1371/journal.pcbi.1000550.s004 (2.83 MB TIF)

**Figure S5** Scale-richness in human PIN. Each diagram shows cumulative degree distributions of proteins in each functional group. The name above each diagram denotes the name of the functional category with which the cumulative degree distribution was examined.

Found at: doi:10.1371/journal.pcbi.1000550.s005 (0.45 MB TIF)

**Table S1** Statistics of sub-networks in the yeast PIN. a. number of nodes b. average shortest path length c. fraction of nodes contained in a largest component to all nodes contained in a sub-network d. average cluster coefficient e. betweenness centrality f. fraction of essential nodes to all nodes contained in a sub-network g. a sub-network consist of low-degree nodes h. a sub-network consist of middle-degree nodes i. a sub-network consist of high-degree nodes j. a sub-network consist of low- and middle-degree nodes k. a sub-network consist of low- and high-degree nodes.

Found at: doi:10.1371/journal.pcbi.1000550.s006 (0.04 MB DOC)

**Table S2** Statistics of sub-networks in the human PIN. a. See Table S1. b. fraction of drug-target nodes contained in a sub-network to all nodes contained in the sub-network.

Found at: doi:10.1371/journal.pcbi.1000550.s007 (0.04 MB DOC)

**Table S3** Statistics of sub-networks in yeast PIN with stringent thresholds for middle- and high-degree nodes. a. See Table S1.

Found at: doi:10.1371/journal.pcbi.1000550.s008 (0.04 MB DOC)

**Table S4** Statistics of sub-networks in human PIN with stringent thresholds for middle- and high-degree nodes. a. See Table S2.

Found at: doi:10.1371/journal.pcbi.1000550.s009 (0.04 MB DOC)

**Table S5** Degrees of the genes in yeast PIN belonging to each functional category. a. Mean degree among the proteins contained in each functional category. b. Number of proteins in each functional category. c. \*\*\*, \*\*, and \* represents that a given value is significantly higher (or lower) than average degree among proteins belonging other functional categories with  $P < 0.001$ ,  $P < 0.01$ , and  $P < 0.05$ , respectively, by the Wilcoxon rank-sum two-sample test with the Bonferroni correction.

Found at: doi:10.1371/journal.pcbi.1000550.s010 (0.06 MB DOC)

**Table S6** Degrees of the genes in human PIN belonging to each functional category. a. See Table S5.

Found at: doi:10.1371/journal.pcbi.1000550.s011 (0.04 MB DOC)

**Table S7** Middle degree proteins in human PIN and their functions.

Found at: doi:10.1371/journal.pcbi.1000550.s012 (2.79 MB DOC)

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## Author Contributions

Conceived and designed the experiments: TH HT YS SN HK. Performed the experiments: TH YS SN. Analyzed the data: TH HT YS SN. Wrote the paper: TH HK.

## The Systems Biology Graphical Notation

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**Circuit diagrams and Unified Modeling Language diagrams are just two examples of standard visual languages that help accelerate work by promoting regularity, removing ambiguity and enabling software tool support for communication of complex information. Ironically, despite having one of the highest ratios of graphical to textual information, biology still lacks standard graphical notations. The recent deluge of biological knowledge makes addressing this deficit a pressing concern. Toward this goal, we present the Systems Biology Graphical Notation (SBGN), a visual language developed by a community of biochemists, modelers and computer scientists. SBGN consists of three complementary languages: process diagram, entity relationship diagram and activity flow diagram. Together they enable scientists to represent networks of biochemical interactions in a standard, unambiguous way. We believe that SBGN will foster efficient and accurate representation, visualization, storage, exchange and reuse of information on all kinds of biological knowledge, from gene regulation, to metabolism, to cellular signaling.**

*“Un bon croquis vaut mieux qu’un long discours”* (“A good sketch is better than a long speech”), said Napoleon Bonaparte. This claim is nowhere as true as for technical illustrations. Diagrams naturally engage innate cognitive faculties<sup>1</sup> that humans have possessed since before the time of our cave-drawing ancestors. Little wonder that we find ourselves turning to them in every field of endeavor. Just as with written human languages, communication involving diagrams requires that authors and readers agree on symbols, the rules for arranging them and the interpretation of the results. The establishment and widespread use of standard notations have permitted many fields to thrive. One can

hardly imagine today’s electronics industry, with its powerful, visually oriented design and automation tools, without having first established standard notations for circuit diagrams. Such was not the case in biology<sup>2</sup>. Despite the visual nature of much of the information exchange, the field was permeated with *ad hoc* graphical notations having little in common between different researchers, publications, textbooks and software tools. No standard visual language existed for describing biochemical interaction networks, inter- and intracellular signaling gene regulation—concepts at the core of much of today’s research in molecular, systems and synthetic biology. The closest to a standard is the notation long used in many metabolic and signaling pathway maps, but in reality, even that lacks uniformity between sources and suffers from undesirable ambiguities (Fig. 1). Moreover, the existing tentative representations, however well crafted, were ambiguous, and only suitable for specific needs, such as representing metabolic networks or signaling pathways or gene regulation.

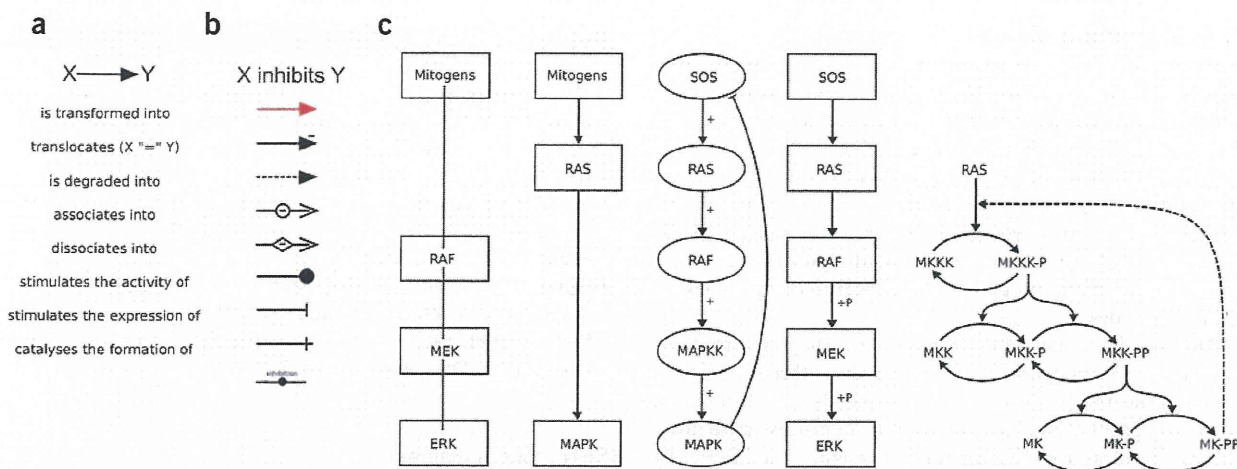
The molecular biology era, and more recently the rise of genomics and other high-throughput technologies, have brought a staggering increase in data to be interpreted. It also favored the routine use of software to help formulate hypotheses, design experiments and interpret results. As a group of biochemists, modelers and computer scientists working in systems biology, we believe establishing standard graphical notations is an important step toward more efficient and accurate transmission of biological knowledge among our different communities. Toward this goal, we initiated the SBGN project in 2005, with the aim of developing and standardizing a systematic and unambiguous graphical notation for applications in molecular and systems biology.

### Historical antecedents

Graphical representation of biochemical and cellular processes has been used in biochemical textbooks as far back as sixty years ago<sup>3</sup>, reaching an apex in the wall charts hand drawn by Nicholson<sup>4</sup> and Michal<sup>5</sup>. Those graphs describe the processes that transform a set of inputs into a set of outputs, in effect being process, or state transition, diagrams. This style was emulated in the first database systems that depicted metabolic networks, including EMP<sup>6</sup>, EcoCyc<sup>7</sup> and KEGG<sup>8</sup>. More notations have been ‘defined’ by virtue of their implementation in specialized software tools such as pathway and network designers (e.g., NetBuilder<sup>9</sup>, Patika<sup>10</sup>, JDesigner<sup>11</sup>, CellDesigner<sup>12</sup>). Those

A list of affiliations appears at the end of the paper.

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**Figure 1** Inconsistency and ambiguity of current nonstandardized notations. (a) Eight different meanings associated with the same symbol in a chart describing the role of cyclin in cell regulations ([http://www.abcam.com/ps/pdf/nuclearsignal/cell\\_cycle.pdf](http://www.abcam.com/ps/pdf/nuclearsignal/cell_cycle.pdf)). (b) Nine different symbols found in the literature to represent the same meaning. (c) Five different representations of the MAP kinase cascade found in the scientific literature, depicting progressive levels of biological and biochemical knowledge. From left to right: relations<sup>30</sup>, directionality of influence<sup>31</sup>, directionality of effect<sup>32</sup>, biochemical effect<sup>33</sup>, chemical reactions<sup>34</sup>. In the last diagram, different instances of an identical arrowhead style represent catalysis, production and inhibition.

graphical notations were not standardized, and their understanding relied mainly on relating examples with one's preexisting knowledge of biochemical processes. Although the classical graphs adequately conveyed information about biochemistry, other types of diagrams were needed to represent signaling pathways, and incomplete or indirect information, as coming from molecular biology or genomics. Those conventions effectively mimicked the empirical notations used by biologists, describing either the relationships between elements<sup>13,14</sup> or the flow of activity or influence<sup>15–17</sup>. Lists of standard glyphs (Box 1) to represent identified concepts were then provided. The efforts to create rigidly defined schema were pioneered by Kurt Kohn with his Molecular Interaction Maps (MIM), which defined not only a set of symbols but also a syntax to describe interactions and relationships of molecules<sup>18,19</sup>. The MIM notation influenced other proposals<sup>14</sup>. Several proposals followed to describe process diagrams, not only with standard symbols but also defined grammars<sup>20–23</sup>.

**The SBGN project**

Despite the popularity of some of the efforts mentioned above, none of the notations has acquired the status of a community standard. This can be attributed partly to the fact that the efforts only went as far as to propose notations, or implement them in software. Several of us have been involved in the development of the Systems Biology Markup Language (SBML)<sup>24</sup>, from which we learned that establishing a standard is extremely difficult without an explicit, concerted, effort to engage a community and build a consensus among participants. We organized the SBGN project with this lesson in mind.

For SBGN to be successful, it must satisfy a majority of technical and practical needs and be embraced by a diverse community of biologists, biochemists, bioinformaticians, geneticists, theoreticians and software engineers. Early in the project's history, we established the following overarching principles to help steer SBGN toward those aims, ranked by rough hierarchical order of precedence.

The notation should

- be free of intellectual property restrictions to allow free use by the community;
- be syntactically and semantically consistent and unambiguous;

- support representation of diverse common biological objects, their properties and their interactions;
- keep the number of symbols and syntax to a minimum to help comprehension and learning by humans;
- be visually consistent and concise, using discriminable symbols;
- support modularity to help cope with diagram size and complexity;
- support the automated generation of diagrams by software starting from mathematical models.

Many of the design principles above resonate with research on visual languages<sup>25,26</sup> and studies aimed at understanding end-user needs in pathway visualization<sup>27</sup>, although we derived them from our collective hands-on experiences with developing notations and software. In addition to these principles, we also sought to avoid many problems (Table 1) that affect some existing notations.

SBGN aims to specify the connectivity of the graphs and the types of the nodes and edges, but not the precise layout of the graphs. The semantics of an SBGN diagram does not depend on the relative position of the symbols. Furthermore, it does not depend on colors, patterns, shades, shapes and thickness of edges (Fig. 2). Similarly, the labels of symbols are not regulated and are only required to be unique within a map.

Finally, it was clear at the outset that it would be impossible to design a perfect and complete language from the beginning. Apart from the prescience this would require, it also would likely require a vast language that most newcomers would shun as being too complex. Thus, the SBGN community decided to stratify language development into levels. A level in SBGN represents a usable set of functionalities that the user community agrees is sufficient for a reasonable set of tasks and goals. Capabilities and features that cannot be agreed upon and are judged insufficiently critical to require inclusion in a given level are postponed to a higher level. In this way, SBGN development is envisioned to proceed in stages, with each higher SBGN level adding richness compared to the levels below it, while maintaining compatibility whenever possible. Furthermore only the actual usage of SBGN languages will tell us how well they work for the diverse communities involved, and this experience will certainly shape the evolution of the notation.

### The three languages of SBGN

Molecular entities possess many properties that affect their interactions with other entities. Attempting to represent all the possible reactions and interactions in the same diagram is often futile, usually resulting in an incomprehensible jumble. The different styles of notations described above were attempts to control this complexity by presenting only what was needed in a specific context, or what was available through specific views of the system<sup>14</sup>. Each view focuses on only a portion of the semantics of the overall system, trading off diagram comprehensibility against completeness of biological knowledge.

SBGN follows this strategy and defines three orthogonal and complementary types of diagrams that can be seen as three alternative projections of the underlying more complex biological information. The process diagram draws its inspiration from process-style notations, borrowing ideas from the work of CellDesigner<sup>28</sup> and EPE<sup>22</sup>. By contrast, the entity relationship diagram is based to a large extent on Kohn's MIM notation<sup>18,19</sup>. The SBGN activity flow diagram depicts only the cascade of activity, thus making the notation similar to the reduced representations often used in the current literature to describe signaling pathways and gene regulatory networks. In Figure 2, we illustrate the three views applied to a very simple example. The characteristics of the SBGN languages are summarized in Table 2.

The idea of having three diagram types naturally begs the question of whether they could be merged into one, at least in paper form. The answer is no, for at least two reasons. First, a single diagram type would bring us back to the problem of dealing with unreasonable numbers of interactions as described above. Second, each SBGN language reflects fundamental differences in the underlying formal description of the phenomena. The meanings are so different that merging diagram types would compromise their representational robustness.

Having multiple visual languages is not uncommon in engineering (consider, for example, block diagrams and circuit diagrams in electronics, UML class, state sequence and deployment diagrams in software engineering), and this supports the idea that having three sublanguages in SBGN will be manageable in practice. In SBGN, the sharing of symbols representing identical concepts further reduces the differences between the three languages to differences in syntax and semantics. We believe that this, combined with careful design, will mitigate some of the difficulties of learning SBGN. However, it is to be noted that the clean orthogonality of the languages makes their overlap very limited, mostly to modulatory arcs, and node decorations.

### Box 1 Glossary

SBGN diagrams are a specific set of graphs and thus make use of concepts from graph theory. The following list defines the terms used most often. We are aware of the unavoidable circularity of such definitions.

- **Arc.** A directed edge, that is, an edge that is not symmetrical in shape.
- **Edge.** A line joining two nodes.
- **Glyph.** A symbol that conveys information nonverbally.
- **Graph.** A set of nodes connected with edges.
- **Node.** A point that terminates a line or curve or comprises the intersection of two or more lines or curves.

### SBGN process diagram

A process diagram represents all the molecular processes and interactions taking place between biochemical entities, and their results. This type of diagram depicts how entities transition from one form to another as a result of different influences; thus, it portrays the temporal qualities of molecular events occurring in biochemical reactions. In this way, the approach underlying process diagrams is the same as in the familiar textbook drawings of metabolic pathways. The main drawback of process diagrams is that a given entity must appear multiple times in the same diagram if it exists under several states; therefore, the notation is sensitive to the combinatorial explosion of possible entities and reactions, as is often the case in signaling pathways.

The SBGN process diagram level 1 specification defines six major classes of glyphs: entity pool nodes, process nodes, container nodes, reference nodes, connecting arcs and logical operators (Supplementary Note 1). In Figure 3a, we show a complete example of an SBGN process diagram. The number of symbols in level 1 of the SBGN process diagram notation has been purposefully limited so that they could be easily memorized. The notation may be enriched (perhaps using subclasses of symbols) in higher levels of SBGN.

Table 3 lists software projects that are already developing support for SBGN process diagram level 1 (see also Supplementary Note 2). Some of these rely on manual design of the pathways, whereas others, such as Arcadia, automatically generate SBGN PD from SBML models that have been annotated with terms from the Systems Biology Ontology<sup>29</sup>. The encoding of SBGN diagrams using computer-readable formats, a crucial step toward exchange and reuse of SBGN

**Table 1 Features of *ad hoc* graphical notations, and the problems they create**

Feature	Problem(s)
Different line thicknesses distinguish different types of processes or elements Dotted or dashed line styles distinguish different types of processes or elements	1. Rescaling a diagram can make line thicknesses and styles impossible to discern 2. Photocopying or faxing a diagram can cause differences in line thicknesses and styles to disappear 3. Differences in line thickness and style are difficult to make consistent in diagrams drawn by hand
Different colors distinguish different types of processes or elements	1. Photocopying or faxing a diagram will cause color differences to be indistinguishable 2. Color characteristics are difficult to achieve and keep consistent when drawing diagrams by hand
Identical line terminators (e.g., a single arrow) indicate different effects or processes depending on context	1. Greater ambiguity is introduced into a diagram 2. Interpreting a diagram requires more thought on the part of the reader 3. Automated verification of diagrams is more difficult due to lack of distinction between different processes or elements
<i>Ad hoc</i> symbols introduced at will by author	Interpreting a diagram requires the reader to search for additional information explaining the meaning of the symbols



diagrams, is currently supported in different formats such as SBML, GML and GraphML by different tools, and a general XML-based exchange format for SBGN is currently under discussion.

**SBGN entity relationship diagram**

The SBGN notation for entity relationships puts the emphasis on the influences that entities have upon each other's transformations rather than the transformations themselves. One can imagine that each of the relationships represents a specific conclusion of a scientific experiment or article. Their addition on a map represents the knowledge we have of the effects the entities have upon each other. Contrary to the process diagrams, where the different processes affect each other, the relationships are independent, and this independence is the key to avoiding the combinatorial explosion inherent to process diagrams. Unlike in process diagrams, a given entity may appear only once. Readers can better grasp at first sight all the possible influences and interactions affecting an entity, without having to explore the whole diagram to discover the different states an entity may be in, or to trace all the edges to find the relevant process nodes.

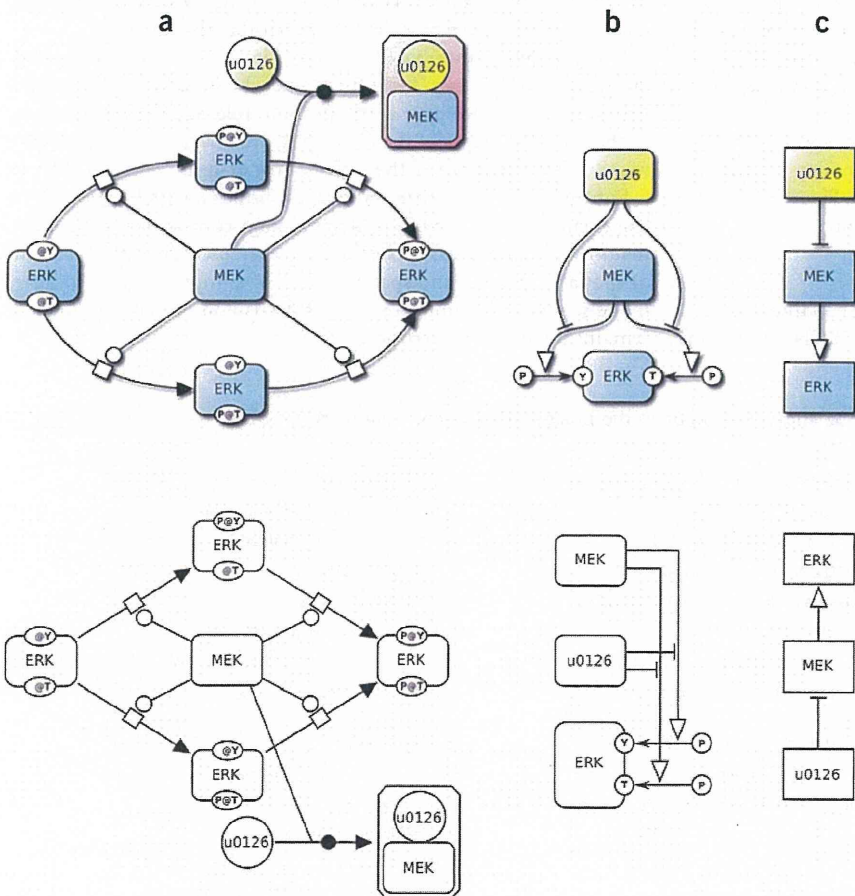
The relationship symbols in entity relationship diagrams support the representation of interactions and state variable assignments, thus allowing the notation to describe certain processes that cannot be expressed in process diagrams, such as allosteric modulation. In process diagrams, one can represent the formation of a ligand-receptor complex, but it is not possible to state that the complex is more active than the receptor alone without additional markup; entity relationships

support this by allowing the interaction with the ligand to modulate the assignment of the variable representing the activity. The trade-off is that the temporal course is difficult to follow in entity relationships, because the sequence of events is not explicitly described (Fig. 2a,b).

The specification of SBGN entity relationship diagram level 1 defines three major classes of glyphs: entity nodes, statements and influences (logical operators are entity nodes). We summarize the symbols and the rules for their assembly (Supplementary Note 3). In Fig. 3b, we show a complete example of an SBGN entity relationship diagram.

**SBGN activity flow diagram**

A strategy often used for coping with biochemical network complexity or with incomplete or indirect knowledge is to selectively ignore the biochemical details of processes, instead representing the influences between entities directly. SBGN's activity flow diagrams permit modulatory arcs to directly link different activities, rather than entities and processes or relationships as described previously. Instead of displaying the details of biochemical reactions with process nodes and connecting arcs, the activity flow diagrams show only influences such as 'stimulation' and 'inhibition' between the activities displayed by the molecular entities (Fig. 2c). For example, a signal 'stimulates' the activity of a receptor, and this activity in turn 'stimulates' the activity of an intracellular transducing protein (note that activity flow retains the sequential chains of influences). Because most signaling pathway diagrams in the current literature are essentially activity flow diagrams, we expect many biologists will find this type of diagram familiar.



**Figure 2** Simple example of protein phosphorylation catalyzed by an enzyme and modulated by an inhibitor. The semantics of an SBGN diagram does not depend on the relative position of the symbols, or on colors, patterns, shades, shapes and thickness of edges. Therefore, the upper and lower diagrams are identical as far as SBGN is concerned, and have to be interpreted exactly the same way. (a) Process diagrams, explicitly displaying the four forms of ERK, phosphorylated and nonphosphorylated on the tyrosine and the threonine, as well as the processes of phosphorylation by MEK and the inhibition of MEK by complexation with u0126. Note that the inhibition in this diagram emerges from the sequestration of MEK and is not explicitly represented. The phosphorylation sites are represented by variables, which in this example are labeled simply as 'Y' and 'T' (but in general could be anything desired by the diagram author), shown adorning the main symbols for ERK. (b) Entity relationship diagrams, showing ERK and the assignment of its phosphorylations (at the tyrosine and threonine residues), as well as the relationships between those and MEK and u0126. Note that ERK appears only once in this diagram; the different possible states are not explicitly depicted. (c) Activity flow diagrams depicting the activation of ERK by MEK and the inhibition of MEK by u0126. In this notation, only the relevant activities of u0126, MEK and ERK are represented, as well as abstract representations of the influences of activities upon each other, whereas the biochemical details are omitted.

**Table 2 Comparison between the three languages of SBGN**

	Process diagram	Entity relationship diagram	Activity flow diagram
<b>Purpose</b>	Represent processes that convert physical entities into other entities, change their states or change their location	Represent the interactions between entities and the rules that control them	Represent the influence of biological activities on each other
<b>Building block</b>	Different states of physical entities are represented separately	Physical entities are represented only once	Different activities of physical entities are represented separately
<b>Ambiguity</b>	Unambiguous transcription into biochemical events	Unambiguous transcription into biochemical events	Ambiguous interpretation in biochemical terms
<b>Level of description</b>	Mechanistic descriptions of processes	Mechanistic description of relationships	Conceptual description of influences
<b>Temporality</b>	Representation of sequential events	Absence of sequentiality between events	Representation of sequential influences
<b>Pitfalls</b>	Sensitive to combinatorial explosion of states and processes	Creation, destruction and translocation are not easily represented	Not suitable to represent association, dissociation, multistate entities
<b>Advantages</b>	The best for representing temporal/mechanistic aspects of processes such as metabolism	The best for representing signaling involving multistate entities	The best for functional genomics and signaling with simple activities

By ignoring processes and entity states, the number of nodes in an activity flow diagram is greatly reduced compared to an equivalent process diagram (Fig. 2a,c). Activity flow diagrams are also especially convenient for representing the effects of perturbations, whether genetic or environmental, because the complete mechanisms of the perturbations may not be known, or are irrelevant to the goals of a given study. The drawback is that activity flow diagrams may contain a high level of ambiguity. For instance, the biochemical basis of a positive or negative influence in a given system is left undefined. For this reason, this type of SBGN diagram should not exist alone; it should be associated, when possible, with detailed entity relationship and process diagrams, and used only for viewing purposes. We expect it will often be possible to generate activity flow diagrams mechanically from process diagrams and entity relationships, and have already performed preliminary work in that direction.

The SBGN activity flow diagram level 1 specification defines four major classes of glyphs: activity nodes, container nodes, modulating arcs and logical operators (Supplementary Note 4). Figure 3c shows a complete example of an SBGN activity flow diagram.

### Participation and future prospects

The SBGN website (<http://sbgn.org/>) is a portal for all things related to SBGN. Interested persons can get involved in SBGN discussions

by joining the SBGN discussion list ([sbgn-discuss@sbgn.org](mailto:sbgn-discuss@sbgn.org)). Face-to-face meetings of the SBGN community, generally held as satellite workshops of larger conferences, are announced on the website as well as the mailing list.

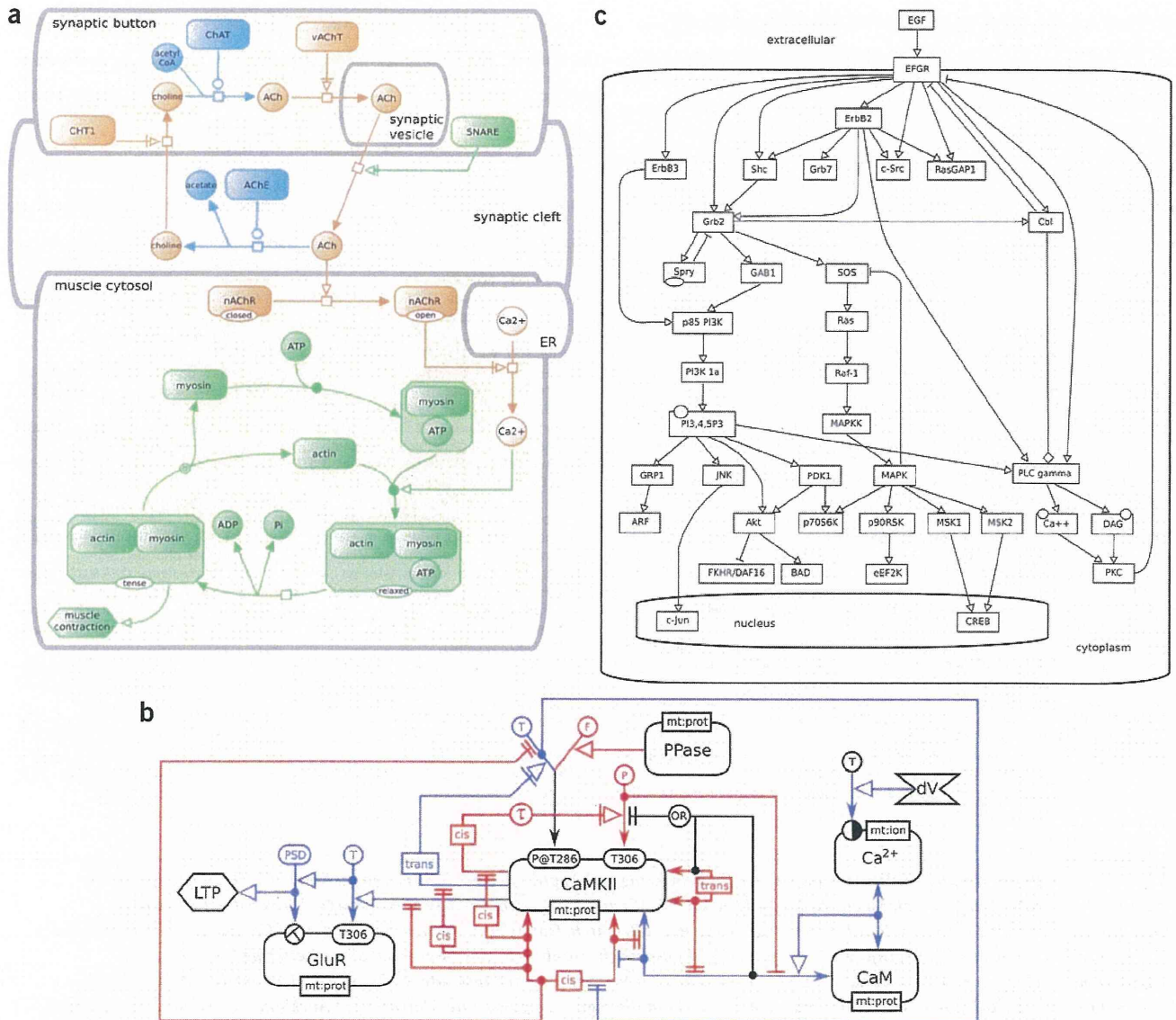
Standardizing a notation for depicting networks of biochemical interactions has so far remained an elusive goal, despite numerous but isolated efforts in that direction. Only with such a standardized notation will biologists, modelers and computer scientists be able to exchange accurate descriptions of complex systems—a task that continues to grow more demanding as our collective knowledge expands. SBGN blends many influences from past efforts, and also introduces many new ideas designed to overcome limitations of other notations.

Using a community-based approach involving many interested groups and individuals (including some who have been involved in previous efforts), we have developed and released the first version of the three languages of the SBGN, the process diagram, the entity relationship diagram and the activity flow diagram.

Future levels of the three languages should address major challenges currently faced by the systems biology community, as the field matures and diversifies. To cite but a few examples, the representation of spatial structures and spatial events, of composed and modular models, and of dynamic creation or destruction of compartments remains uncharted territory.

**Table 3 List of software systems known to provide support, or to be in the process of developing support, for SBGN**

Name	Organization	Link
Arcadia	Manchester Centre for Integrative Systems Biology, Manchester, UK	<a href="http://arcadiapathways.sourceforge.net/">http://arcadiapathways.sourceforge.net/</a>
Athena	University of Washington, Seattle, WA, USA	<a href="http://www.codeplex.com/athena/">http://www.codeplex.com/athena/</a>
BioModels Database	European Bioinformatics Institute, Cambridge, UK	<a href="http://www.ebi.ac.uk/biomodels/">http://www.ebi.ac.uk/biomodels/</a>
BioUML	Institute of Systems Biology, Novosibirsk, Russia	<a href="http://www.biouml.org/">http://www.biouml.org/</a>
ByoDyn	Institut Municipal d'Investigació Mèdica, Barcelona, Spain	<a href="http://byodyn.imim.es/">http://byodyn.imim.es/</a>
CellDesigner	The Systems Biology Institute, Tokyo	<a href="http://www.celldesigner.org/">http://www.celldesigner.org/</a>
Dunnart	Monash University, Melbourne, Australia	<a href="http://www.csse.monash.edu.au/~mwybrow/dunnart/">http://www.csse.monash.edu.au/~mwybrow/dunnart/</a>
Edinburgh Pathway Editor	Edinburgh Centre for Bioinformatics, Edinburgh, UK	<a href="http://www.pathwayeditor.org/">http://www.pathwayeditor.org/</a>
JWS Online	Stellenbosch University, Stellenbosch, South Africa	<a href="http://jwj.biochem.sun.ac.za/">http://jwj.biochem.sun.ac.za/</a>
NetBuilder	STRI, University of Hertfordshire, Hatfield, UK	<a href="http://strc.herts.ac.uk/bio/maria/Apostrophe/">http://strc.herts.ac.uk/bio/maria/Apostrophe/</a>
PANTHER	Artificial Intelligence Center, SRI international, Menlo Park, CA, USA	<a href="http://www.pantherdb.org/pathway/">http://www.pantherdb.org/pathway/</a>
Reactome	European Bioinformatics Institute, Cambridge, UK	<a href="http://www.reactome.org/">http://www.reactome.org/</a>
Vanted	IPK Gatersleben, Gatersleben, Germany	<a href="http://vanted.ipk-gatersleben.de/">http://vanted.ipk-gatersleben.de/</a>
VISIOweb	Bilkent University, Ankara, Turkey	<a href="http://www.bilkent.edu.tr/~bcbi/pvs.html">http://www.bilkent.edu.tr/~bcbi/pvs.html</a>



**Figure 3** Example of complete SBGN diagrams. (a) Process diagram representing the synthesis of the neurotransmitter acetylcholine in the synaptic bouton of a nerve terminal, its release in the synaptic cleft, degradation in the synaptic cleft, the post-synaptic stimulation of its receptors and the subsequent effect on muscle contraction. Colors are used to enhance the biological semantics, blue representing catalytic reactions, orange for transport between compartments (including unrepresented ions, through channels) and green for the function of contractile proteins. However, it is important to note that those colors are not part of SBGN process diagram notation, and must not change the interpretation of the graph. (b) SBGN entity relationship diagram representing the transduction, by calcium/calmodulin kinase II, of the effect of voltage-induced increase of intracellular calcium onto the long-term potentiation (LTP) of the neuronal synapses, triggered by a translocation of glutamate receptors. The diagram describes the various relationships between the phosphorylations of the kinase monomers and their conformation. Colors highlight the direction of the relationships relative to the phenotype; blue relationships enhance LTP whereas red ones preclude this enhancement. (c) SBGN activity flow diagram representing the cascade of signals triggered by the epidermal growth factor, and going from the plasma membrane to the nucleus. The diagram is derived from reference 30.

Note: Supplementary information is available on the Nature Biotechnology website.

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## REPORT

# Robustness and fragility in the yeast high osmolarity glycerol (HOG) signal-transduction pathway

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Cellular signalling networks integrate environmental stimuli with the information on cellular status. These networks must be robust against stochastic fluctuations in stimuli as well as in the amounts of signalling components. Here, we challenge the yeast HOG signal-transduction pathway with systematic perturbations in components' expression levels under various external conditions in search for nodes of fragility. We observe a substantially higher frequency of fragile nodes in this signal-transduction pathway than that has been observed for other cellular processes. These fragilities disperse without any clear pattern over biochemical functions or location in pathway topology and they are largely independent of pathway activation by external stimuli. However, the strongest toxicities are caused by pathway hyperactivation. *In silico* analysis highlights the impact of model structure on *in silico* robustness, and suggests complex formation and scaffolding as important contributors to the observed fragility patterns. Thus, *in vivo* robustness data can be used to discriminate and improve mathematical models.

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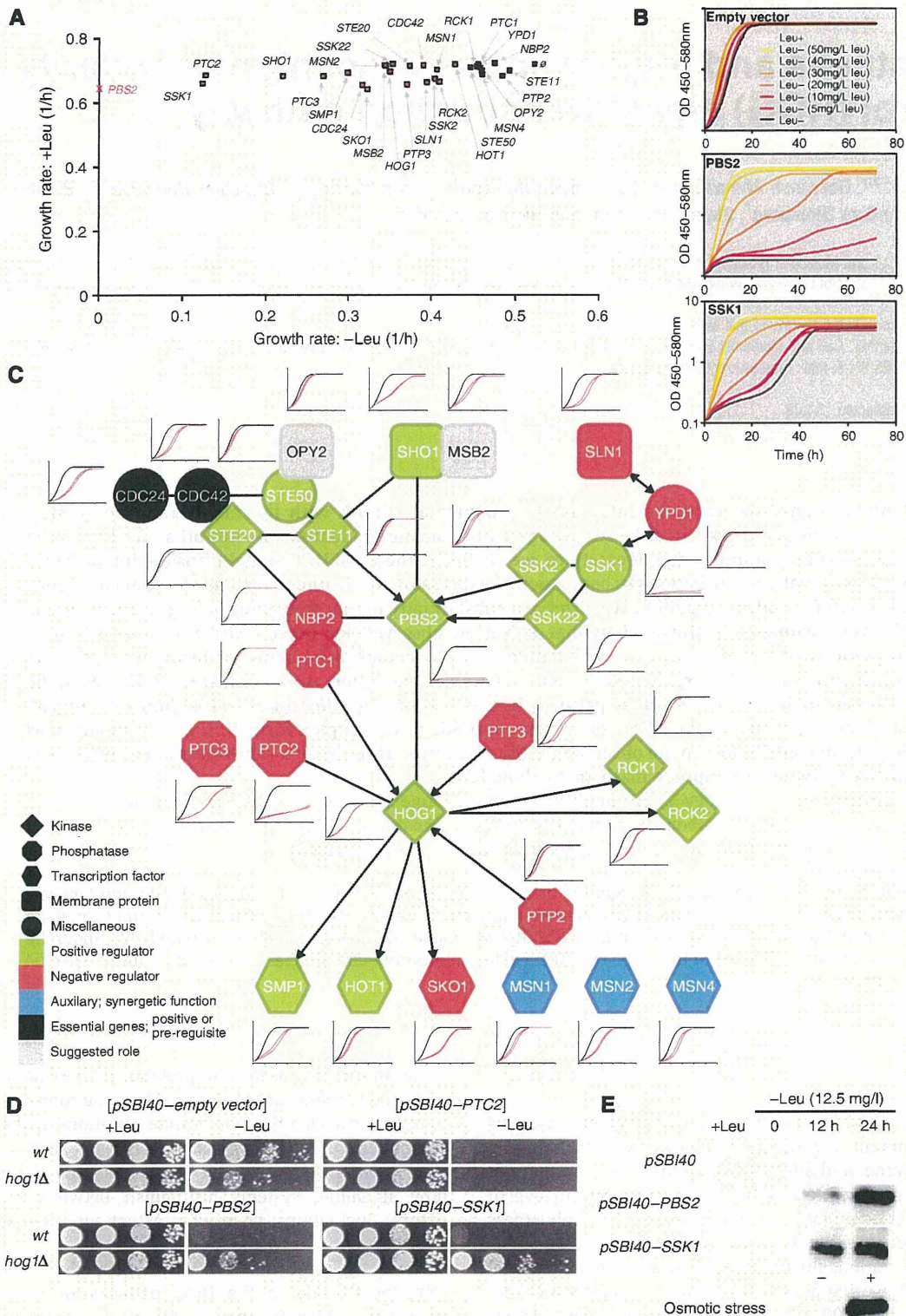
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## Introduction

Robustness is an intrinsic feature of life as all cellular systems have to maintain functionality in the face of naturally occurring external and internal fluctuations. The resilience of cellular genetic networks lets the cell tolerate a certain level of environmental or mutational perturbations. This robustness can be achieved either by maintaining the cellular status stable against various fluctuations, or by adapting to external changes by triggering a series of cellular responses (Kitano, 2004; Stelling *et al.*, 2004). The decision to respond and adapt is relayed via signal transduction systems, which, upon activation by specific stimuli, produce distinct regulatory signals in the form of changes in levels of activated signal-pathway components. A critical aspect of such processes is distinguishing a genuine signal from stochastic fluctuations in protein

levels and activity, as misinterpretation of these has potentially disastrous fitness consequences. Thus, the robustness of cells to maintain such a function despite variations in dosage of the components is of primary importance for survival. Despite their importance for viability and fitness, little is known about how signalling systems distinguish between signals and natural fluctuations, or to what extent such fluctuations are tolerated.

Here, we approach this issue through a system-wide robustness study of the HOG pathway of *Saccharomyces cerevisiae*, which is one of the most extensively studied eukaryotic signal-transduction cascades. It is activated by high osmolarity and is essential under this condition. The signalling pathway, which is depicted in Figure 1, consists of a MAP kinase (MAPK) core module, upon which two independent upstream branches converge. The first of these consists of



**Figure 1** (A) gToW growth phenotypes occur in the absence of leucine, and red squares indicate a significant growth-rate defect as compared with the empty plasmid control ( $\emptyset$ ). *PBS2*'s growth rate could only be determined in the presence of leucine. (B) The severity of the growth defect increases with the level of leucine starvation and (C) they spread over different pathway functions. Graphs in (C) indicate growth with (black) or without (red) leucine. (D) Phenotypes caused by *PBS2* and *SSK1*, unlike *PTC2*, are partially suppressed by the deletion of *HOG1*. (E) Overexpression of *Pbs2p* and *Ssk1p* causes dual phosphorylation of *Hog1p*, which after leucine limitation (12.5 mg/l), reaches levels comparable to those caused by osmotic stress (+) within 24 h. The empty plasmid control (*pSBI40*) remains similar to unstressed cells (-).

a phosphotransfer module, including the histidine kinase and presumed osmosensor Sln1p, the phosphotransfer protein Ypd1p and the response regulator Ssk1p. When active, this module keeps Ssk1p phosphorylated and inactive. When the module is inactive, dephosphorylated Ssk1p binds to and activates the MAPK kinase kinases Ssk2p and Ssk22p (Saito and Tatebayashi, 2004). Two mucin-like proteins Msb2p and Hkr1p were recently suggested as putative osmosensors of the second input branch (Tatebayashi *et al*, 2007). On activation, the transmembrane protein Sho1p is believed to receive signals from these sensors and convey these signals to the interior of the cell. Sho1p also assembles the MAPK kinase Pbs2p and the MAPK kinase kinase Ste11p through its cytoplasmic domain. The Sho branch also requires Cdc42p and Ste20p for the transmission of the signal. Once active, Ste11p, Ssk2p and Ssk22p are each able to phosphorylate Pbs2p, which also acts as a scaffold for the MAPK module. Pbs2p in turn phosphorylates and activates the MAPK Hog1p, which has numerous targets, including the cytoplasmic kinase Rck2p and several transcription factors such as Hot1p, Sko1p and Smp1p. Active Hog1p accumulates in the nucleus and partakes in transcription (Hohmann, 2002).

The methodology used here links the expression of each target gene of interest to that of a defective allele of a metabolic gene. This allele, *leu2-d*, has a defective promoter and needs to be present in a high copy number to support high Leu2p levels and thus unperturbed growth in the absence of leucine (Schneider and Guarente, 1991). By placing a target gene of interest on the same episomal plasmid as *leu2-d*, the copy number and thus the expression level of this target gene can be controlled via the leucine concentration in the media. The gene in question is still under the control of its normal promoter, allowing expression that is regulated but increases proportionally with the increase in copy number (Moriya *et al*, 2006; Torres *et al*, 2007). If the increase in target gene product interferes with cellular function, a negative pressure on plasmid copy number will balance the positive pressure conferred by the metabolic gene, resulting in a genetic tug-of-war (gToW; Supplementary Figure S1). Such a compromise will result in a decrease in cellular fitness, which can be measured precisely using high-resolution microcultivation (Warringer *et al*, 2003).

## Results and discussion

### The HOG pathway shows a low level of robustness particularly during the adaptation phase

On applying the gToW methodology to investigate the HOG-pathway robustness, we found a high prevalence of negative impacts from gene dosage perturbations within the HOG pathway (Figure 1A). We used three physiological 'windows' to assess the robustness; growth rate, growth adaptation time and growth efficiency. Growth rate was considered the primary readout because of its strong correlation to plasmid copy number (Moriya *et al*, 2006). Adaptation defects (prolonged growth lag) turned out to be similar to growth-rate defects, although even more pronounced and frequent (Supplementary Figure S2a). There was almost no effect on growth efficiency (Supplementary Figure S2b). Altogether, overexpression of 22

out of 29 HOG-pathway components caused a significant defect in at least one of these three growth variables ( $P < 0.001$ , see Supplementary information). These phenotypes were strongly linked to leucine starvation and thus to selective pressure for high plasmid copy numbers (Figure 1B). They were also observed for all protein classes, and for both positive and negative regulators of the pathway (Figure 1C). The negative effects from increases in gene dosage were emphasized by a complete lack of positive fitness effects mediated by any of the gToW constructs. The highest frequency and strength of defects were observed during adaptation, which may reflect the delicate balance of signal transmission in initiating proliferation.

The high frequency (76%) of HOG-pathway gToW-imposed defects stands in stark contrast to the cell-cycle system, for which only 25–30% of the gToW constructs caused a clear growth retardation (Moriya *et al*, 2006), and to a global *GAL1* promoter-driven overexpression study using galactose induction in which a mere 15% of the targets conferred detectable growth defects (Sopko *et al*, 2006). Interestingly, little correlation in terms of cellular toxicity was found between the gToW- and galactose-driven overexpressions of the same, but GST tagged proteins, even on galactose (Supplementary Figure S3). The sole exception was Ssk1p that scored as highly toxic with both methods. The lack of correlation may be explained by the varying absolute levels of overexpressions (Supplementary Figure S3e) in the different screens and by the influence of the GST tag on protein function.

The high prevalence of nodes of fragility within the HOG pathway may be partly explained by the very nature of signal transduction, as overexpression defects are enriched among components that transduce adaptation signals, that is kinases, phosphatases and transcription factors (Sopko *et al*, 2006). However, the high fraction of such components in the HOG pathway cannot be the sole reason for the high frequency of fitness defects, as the system is also sensitive to overexpression of more than half of the components, which do not partake in (de)phosphorylation or transcription. In fact, each such component that does not impair normal growth when overexpressed is either known or presumed to be a targeting or activating partner of catalytic components within the pathway, that is Nbp2p for Ptc1p, Ste50p for Ste11p and Opy2p for Ste50p/Ste11p (Posas *et al*, 1998; Mapes and Ota, 2004; Wu *et al*, 2006). Overexpression of their catalytic partners, Ste11p and Ptc1p, also failed to cause significant growth defects ( $P > 0.001$ ; see Supplementary information). Overall, these paired components are more robust against overexpression ( $P = 0.022$ , Fisher's exact test).

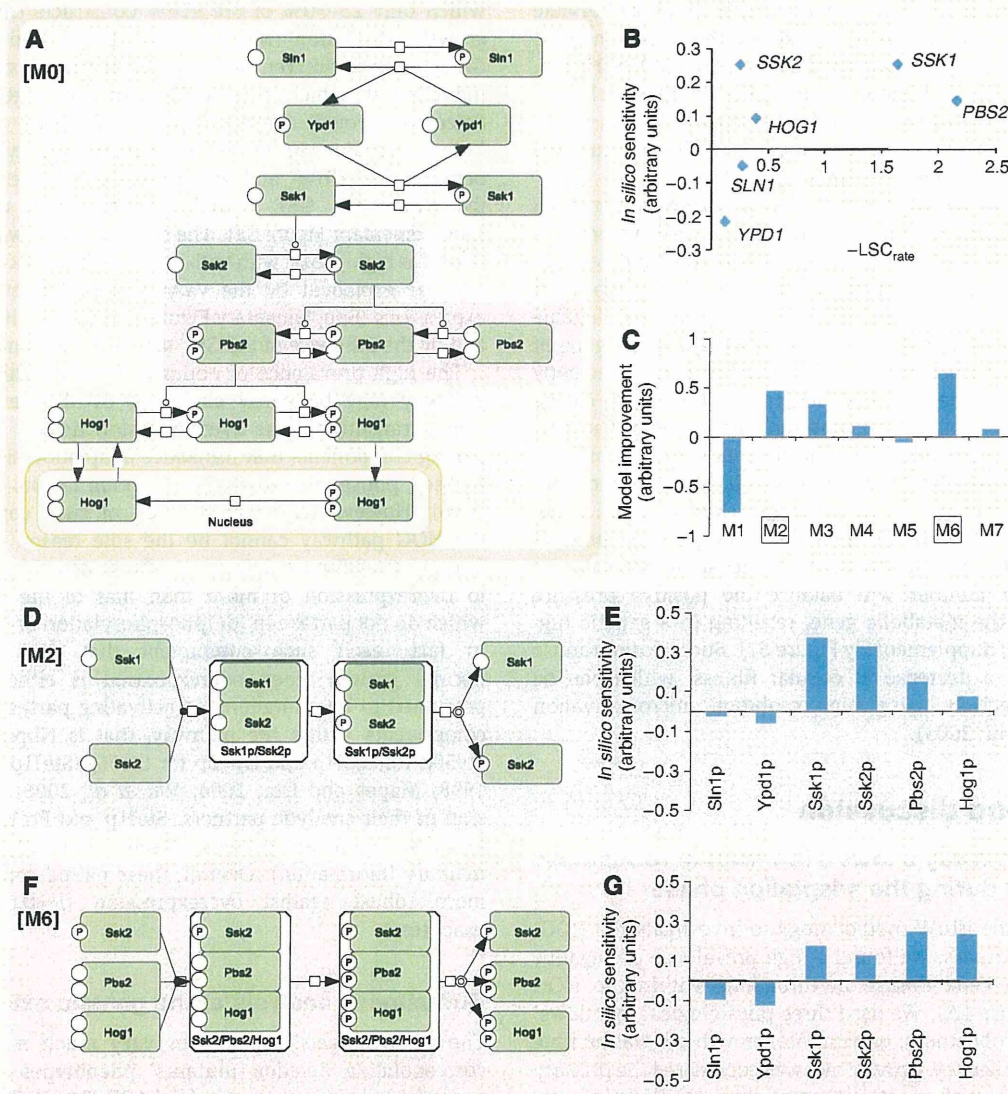
### Robustness analysis at the genetic extremes

The gToW targets' phenotypes vary much more than the corresponding deletion mutants' phenotypes. Four of the genes included here are essential (*CDC24*, *CDC42*, *SLN1* and *YPD1*), but the twenty-five viable deletion strains show at the most mild growth defects during normal conditions. Only in the presence of NaCl stress does deletion of those genes cause strong but viable phenotypes. However, these deletion phenotypes have little in common with those caused by

overexpression (Supplementary Figure S4). The strongest gToW phenotype is caused by Pbs2p, which is well known to be highly important for growth on salt as well as severely toxic when overexpressed. Hog1p is likewise important for osmotic tolerance but, unlike Pbs2p, not toxic when overexpressed. In addition, Ssk1p is severely toxic when overexpressed, but dispensable for osmotic tolerance. Although their roles in the osmotic stress response are well known, the toxicity mechanisms of Ssk1p and Pbs2p overexpression remain to be mechanistically resolved. However, both activate Hog1p constitutively and are suppressed by the deletion of *HOG1*, indicating that most, if not all, of the toxicity stems from the downstream pathway hyperactivation (Figure 1D and E).

### Understanding toxicity mechanisms

In order to gain more insight into the mechanisms of toxicity of the two main nodes of fragility, Ssk1p and Pbs2p, we compared the observed *in vivo* sensitivity profiles with the *in silico* sensitivities with respect to nuclear, dually phosphorylated Hog1p predicted by the Hog model by Klipp *et al* (2005) (Figure 2A). As the *in vivo* differences in fragility between Ssk1p and Ssk2p and the fragility of the Pbs2p node cannot be captured using the original model, we studied seven variants with alternative motifs of regulation involving Ssk1p and Pbs2p (Supplementary Figure S5) and scored the relative improvements of each in the light of our data on overexpression (Figure 2C). We found that the *in silico* sensitivity



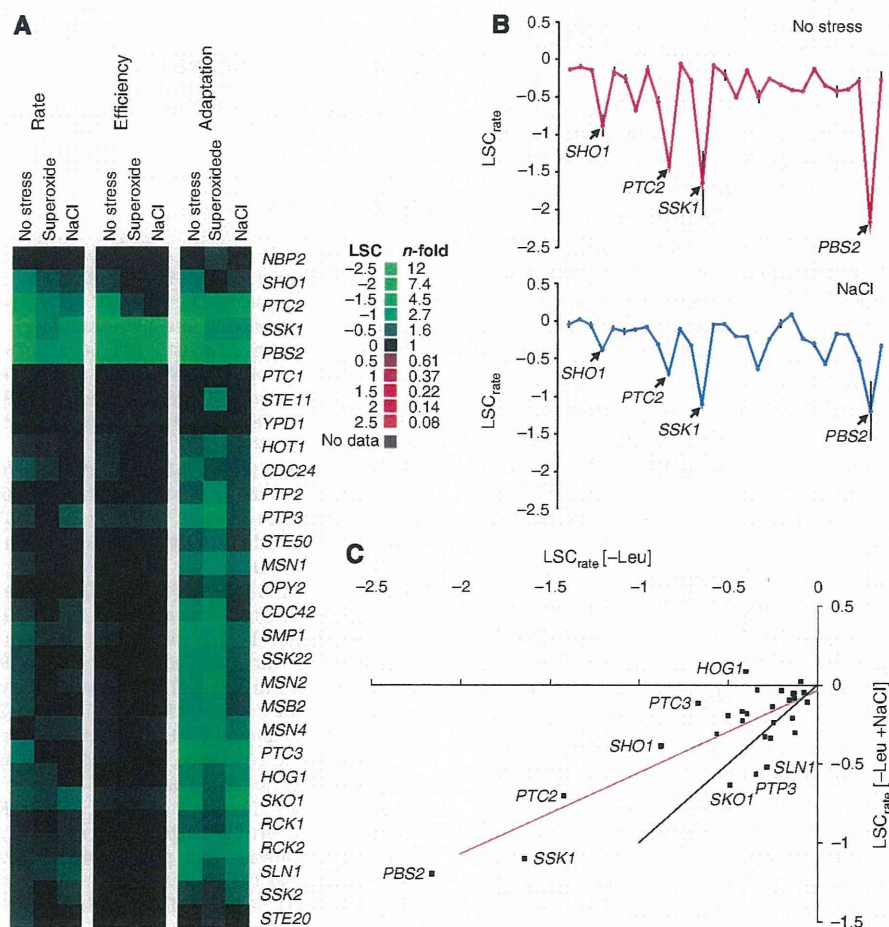
**Figure 2** (A) The Hog part of the mathematical osmoregulatory model by Klipp *et al* (2005) (B) *In vivo* growth-rate defects in leucine-free medium are compared with *in silico* increases in basal levels of nuclear, dually phosphorylated Hog1p as a result of gene overexpression. The model does not capture the fragility of the Pbs2p node or distinguish the sensitivities of SSK1 and SSK2. (C) The relative improvement in model performance by the inclusion of regulatory motifs around Ssk1p (M1–M3) and Pbs2p (M4–M7) (Supplementary Figure S5). (D) Requirement of dimerization before the phosphorylation of Ssk2p yielded the best improvements for Ssk1p (E; the corresponding sensitivity profile). (F) Explicit modelling of the scaffold function yields the best improvements for Pbs2p (G; the corresponding sensitivity profile). The LSC scores the growth difference compared with wild type, with a negative value, indicating a growth defect.



of Ssk1p is enhanced most when the dimerization of Ssk1p with Ssk2p is required for the phosphorylation and activation of Ssk2p (Figure 2D and E). In addition, explicit modelling of Pbs2p's function as a scaffold best improves its performance regarding the fragility of the Pbs2p node (Figure 2F and G). However, we observed no improvement of the *in silico* robustness through the implementation of the known dimerization of Ssk1p alone (Supplementary Figure S5; M1), suggesting that it is unlikely to contribute to the robustness pattern.

If *in vivo* toxicity stemmed from the indiscriminate interaction between protein pairs, we would expect the effect of overexpression to be roughly symmetrical for transient interactions or biased towards the component with lower expression levels in case of sustained interactions. Here, we see neither. Ssk1p has both a much stronger phenotype and higher basal expression level than Ssk2p (Supplementary Figure S6a). Ssk22p is even less abundant and the effect of Ssk1p overexpression is suppressed in *ssk2Δ* (Supplementary Figure S6d). As the phosphorylated, inactive state of Ssk1p has

been reported to be stabilized by Ypd1p (Janiak-Spens *et al*, 2000), and the gToW overexpression brings Ssk1p into parity with Ypd1p levels (Supplementary Figure S6b), it may be the depletion of the stabilizing Ypd1p that leads to an accumulation of dephosphorylated and active Ssk1p (Supplementary Figure S7). Consistently, deletion of either *SLN1* or *YPD1* is lethal owing to the resulting constitutive activity of Ssk1p/Ssk2p and the HOG pathway. Pbs2p likewise stands out as being much more sensitive than its neighbours. Although this toxicity may stem from a disrupted balance with negative regulators, such as the Nbp2p–Ptc1p phosphatase complex, the high basal abundance of Pbs2p argues against the depletion of Nbp2–Ptc1 as the sole source of toxicity (Supplementary Figure S6a). However, the toxicity stems from the amplification of an existing residual signal, as it can be suppressed by deletion of the upstream kinase Ssk2p (Supplementary Figure S6d). As for Ssk1p, it leads to hyperactivation of Hog1p (Figure 1E) and this is the source of its toxicity (Figure 1D). In contrast to Ssk1p and Pbs2p, the phenotype caused by the phosphatase Ptc2p is not mediated



**Figure 3** (A) Hierarchical clustering of the growth-phenotype profiles in the presence or absence of environmental perturbations. The phenotypic effect is indicated by colour. (B) Experimentally measured growth-rate-toxicity profiles in the presence or absence of an external pathway activator ( $LSC_{rate} \pm s.d.$ ,  $n=2$ ; gene order as in Supplementary Table S1). (C) There is a strong correlation between the phenotypic effect under the different conditions, although the relative gToW effect is milder under adverse growth conditions—NaCl ( $r^2=-0.57$ ,  $k=0.5$  (red line; the black line indicates 1:1 correlation)).

through the activation of the HOG pathway and cannot be suppressed by the deletion of *HOG1* (Figure 1D), as would be expected because its overexpression phenotype is stronger than the deletion phenotype of Hog1p (Supplementary Figure S4b). Instead, the mechanism of its toxicity should be found outside the context of the HOG pathway.

### The robustness of the HOG pathway is partly dependent on the environmental stress

To determine whether the HOG-pathway robustness is dependent on pathway activation status, we probed the HOG-pathway robustness during NaCl stress, which is known to activate the pathway, and superoxide stress (paraquat addition), which does not activate the pathway. Both stresses were applied in doses causing a similar (40–50%) reduction in reference strain growth rate. Interestingly, the relative gToW sensitivity patterns during these two stresses were very similar to that observed during unstressed conditions, indicating that the nodes of fragility remain regardless of pathway activation. The phenotypic correlations between these growth conditions ranged from 0.57 to 0.81 (Figure 3;  $r^2$  for LSC rate). The impact of the HOG gToW perturbations were significantly stronger under normal growth conditions than under either NaCl or paraquat stress ( $P=0.005$  and  $7.4 \times 10^{-5}$ , respectively, paired *t*-test of LSC rate). Apart from this general dampening effect, which is observed under different stress conditions, robustness is largely independent of pathway activation by environmental perturbations. However, we see an indication of interaction between the genetic and environmental perturbations. Although paraquat and NaCl stress give similar trends in the dampening of the phenotypes, the variance around this trend seems higher under NaCl (Supplementary Figure S8). An appealing interpretation would be conditional alleviation or aggravation, which would be expected if the effect of the genetic and environmental perturbation cancel out or act synergistically, respectively. We find it interesting that the targets furthest from the trend line under NaCl stress are *SLN1*, *SKO1* and *PTP3*, all known negative regulators of the osmotic stress response, on the negative side and *HOG1* and *PTC3* on the positive side. *PTC3* is equally and surprisingly alleviated by both paraquat and NaCl stress.

In summary, we used the gToW method to qualitatively capture nodes of fragility from overexpression within the HOG pathway. The quantitative correlation to the level of overexpression is more difficult to assess due to additional levels of gene, mRNA and protein regulation. However, previous results by Moriya *et al* show a correlation between growth phenotype, plasmid copy number and relative protein overexpression. Here, we report that expression changes have very strong impact on signalling. The system robustness against overexpression is heavily dependent on the target component and neighbouring nodes show very different fragility. In the HOG pathway, overexpression of Pbs2p and Ssk1p yield the strongest effects, whereas none of their neighbours; Ssk2p, Ste11p or Hog1p, are similarly sensitive. The *in silico* analysis of model variants clearly shows that model structure has a strong impact on the fragility of different nodes. Our results suggest that the stable formation of an Ssk1p–Ssk2p dimer and

Pbs2p's scaffold function contribute to the fragility of their respective nodes. Although robustness information alone cannot be used to reject model structures, it provides information complementary to dynamic data that can be used to discriminate models, and should prove a valuable tool in any modelling endeavour.

### Supplementary information

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

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### Conflict of interest

The authors declare that they have no conflict of interest.

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## Consistent design schematics for biological systems: standardization of representation in biological engineering

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- Supplementary data**      ["Data Supplement"](#)  
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