Diverse protein kinase interactions identified by protein microarrays reveal novel connections between cellular processes

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Protein kinases are key regulators of cellular processes. In spite of considerable effort, a full understanding of the pathways they participate in remains elusive. We globally investigated the proteins that interact with the majority of yeast protein kinases using protein microarrays. Eighty-five kinases were purified and used to probe yeast proteome microarrays. One-thousand-twenty-three interactions were identified, and the vast majority were novel. Coimmunoprecipitation experiments indicate that many of these interactions occurred in vivo. Many novel links of kinases to previously distinct cellular pathways were discovered. For example, the well-studied Kss1 filamentous pathway was found to bind components of diverse cellular pathways, such as those of the stress response pathway and the Ccr4–Not transcriptional/translational regulatory complex; genetic tests revealed that these different components operate in the filamentation pathway in vivo. Overall, our results indicate that kinases operate in a highly interconnected network that coordinates many activities of the proteome. Our results further demonstrate that protein microarrays uncover a diverse set of interactions not observed previously.

[Keywords: Kss1 MAPK; interaction network; protein kinase; protein microarrays; proteomics; yeast]

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In our study, we investigated the proteins that interact with yeast protein kinases using protein microarrays. Protein microarrays offer the advantage that the arrays contain nearly all yeast proteins, and they can be assayed simultaneously for biochemical interactions in vitro [Zhu et al. 2001]. In contrast, many other methods, such as AC/MS, primarily detect interactions between the most abundant proteins, and may also be biased toward identification of those components in stable complexes. Therefore, it is likely that new interactions could be discovered using protein microarrays. However, thus far, only limited studies to globally analyze protein–protein interactions have been performed using this approach [Zhu et al. 2001; Popescu et al. 2007].

In this study, we employed a large-scale approach using protein microarrays to identify yeast kinase–protein interactions. We discovered and validated many novel interactions, revealing an entirely new set of functional links between kinases and cellular pathways. Such information was shown to provide new insights into even well-studied biological processes such as the filamentation signaling pathway.

Results

In vitro protein kinase-binding profiles

To identify proteins that interact with protein kinases using proteome microarrays, 109 of the ~125 yeast protein kinases [Zhu et al. 2001; Breitkreutz et al. 2010] were tagged with a C-terminal V5 epitope. Eighty-five of these were successfully purified in sufficient quantities and purity to probe a yeast proteome microarray containing ~4200 unique yeast proteins [Fig. 1; Table 1; see the Materials and Methods]. Each kinase was assayed using two protein microarrays, along with a negative control (see the Materials and Methods), and the overlapping interactions for each duplicate experiment were analyzed using modified ProCAT software [Zhu et al. 2006]. Using stringent criteria (described in the Materials and Methods), we identified 1023 interactions involving 425 different proteins [Supplemental Table S1]. The number of interacting proteins for each kinase ranged from one to 47 (median = 9). The majority of targets bound a single kinase, although 198 proteins bound more than one kinase (Supplemental Fig. S1).

Coimmunoprecipitation (co-IP) validation of targets in vivo

To evaluate whether the proteins bound on the array also interact with the protein kinases in vivo, we tested interactions for nine kinases from distinct subfamilies using co-IP experiments. Forty-five strains were constructed that contain a protein kinase, tagged at its C terminus with 13XMyC, and the interacting protein identified from the protein array, tagged at its C terminus with a TAP tag [Rigaut et al. 1999]. These included 20 proteins that interacted with 10 or more kinases (two of these were tested for interactions with multiple kinases). The 45 strains contained proteins whose kinase interactions were detected at a wide range of intensities on the protein microarrays. A strain with tagged Crz1, which was known previously to interact with Hrr25 [Kafadar et al. 2003], was also included as a positive control. In each case, the proteins were expressed from native loci to assess the binding interactions at their endogenous levels. Protein complexes were isolated from crude yeast cell extracts using IgG-coupled magnetic beads to capture the TAP-tagged target protein, and the immunoprecipitates were examined for the presence of 13XMyC protein kinases by immunoblot analysis. Negative controls included the untagged kinase and untagged interacting protein strains. A total of 32 novel protein–protein interactions identified in the protein microarray screen were confirmed by co-IP [Fig. 2], as was the positive control. The negative control strains did not coimmunoprecipitate. Of the 13 that failed [Supplemental Table S2], 10 involved the proteins that interacted with >10 kinases and are likely to be nonspecific interactions on the original array; the remaining three proteins interacted with <10 kinases. For the three proteins that appear to be specific (i.e., interacted with <10 proteins on the array) but did not validate using co-IP, their lack of interaction cannot be readily interpreted; they might interact under other growth/environmental conditions. In summary, 20
partner interactions in 887 distinct kinase-interacting data set of Breitkreutz et al. (2010) had 1844 partner–Supplemental Fig. S2A). In contrast, the affinity capture GRID (an example of the most connected one is shown in other interacting proteins of the same kinase from Bio-

Table 1. Kinases tested in this study sorted by major kinase groups

<table>
<thead>
<tr>
<th>Group</th>
<th>Kinases tested</th>
</tr>
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<tbody>
<tr>
<td>AGC</td>
<td>Chk1, Dbf2, Dbf20, Kin82, Phk2, Phk3, Sch9, Tpk1, Tpk2, Tpk3, Yhr028c, Ypk1</td>
</tr>
<tr>
<td>Atypical PK</td>
<td>Rpr1</td>
</tr>
<tr>
<td>CAMK</td>
<td>Cmk1, Cmk2, Elm1, Frk1, Gin4, Hsl1, Kcc4, Kin1, Kin2, Mek1, Rck1, Rck2, Snf1, Ymr291w, Tos3, Ypl150w</td>
</tr>
<tr>
<td>CAMK/EMK</td>
<td>Prr1</td>
</tr>
<tr>
<td>CK1</td>
<td>Hr25, Yck2, Yck3</td>
</tr>
<tr>
<td>CMGC</td>
<td>Cak1, Cdc28, Ctk1, Ctk2*, Fus3, Kin28, Kss1, Mrk1, Mrk1, Rim11, Sky1, Slr2, Smk1, Sn3, Yak1, Ygk3</td>
</tr>
<tr>
<td>Other</td>
<td>Ak1, Ark1, Cdc7, Chk1, Gcn2, Hal5, Ip11, Kkg8, Ksp1, Mps1, Npr1, Prr2, Psk2, Ptk2, Rtk1, Sat4, Scy1, Sks1, Sxe1, Vhs1</td>
</tr>
<tr>
<td>STE</td>
<td>Bck1, Cdc15, Cia4, Kin3, Mkk1, Pbs2, Skm1, Sps1, Ssk2, Ste7, Ste11, Ste20</td>
</tr>
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Eighty-five kinases were used to screen for protein–protein interactions on the protein arrays. The kinases were sorted according to their taxonomical grouping: AGC (cyclic nucleotide-dependent family [PKA and PKG] and the protein kinase C), Atypical PK, CAMK (calcium/calmodulin), CAMK/EMK (CAMK/ELKL motif kinase), CK1 [casein kinases], CDKs [cyclin-dependent kinases], MAP kinases [mitogen-activated protein kinases], CMGC (glycogen synthase kinases [GSK] and CDK-like kinases), other [kinases that do not have sequence similarity with any other kinase], STE [sterile homologs], and Yeast PK (kinases unique to yeast). Ctk2* is the β subunit of C-terminal domain kinase 1 (CTDK1) kinase.

out of 23 (83%) specific interaction sets [those that bound <10 kinases] were validated, whereas only half [10 out of 20] of the less specific interaction sets [those that bound >10 kinases] were validated. Given the success rate for validation of targets bound more specifically to kinases tested in our screen, the data were filtered to include only those interactions occurring between <10 kinases on the protein microarray, our data suggest that the majority of these interactions occur in vivo.

The interactions identified by protein microarrays are distinct from those found by other methods

Our study is the largest interaction screen performed to date using protein microarrays. Several other global studies to identify proteins that interact with protein kinases have been performed, notably AC/MS studies and two-hybrid studies. Surprisingly, in spite of the extensive in vivo validation we performed, only a small fraction of the interacting proteins found using the protein microarrays overlap with those of the other studies. We found that only two of 1023 interactions overlapped with earlier genome-wide co-IP studies, and nine overlapped with a very recent and thorough kinase AC/MS study [Fig. 3A; Breitkreutz et al. 2010]. Similarly, only three of 1023 of our interactions overlap with a recent two-hybrid study [Yu et al. 2008]. These results indicate that the interacting partners identified using protein microarrays are distinct from those found using other methods.

We suspect that the minimal overlap of our results with those identified by other methods is likely due to the fact that our method identifies binary interactions, whereas AC/MS identifies entire complexes. To explore this possibility further, we analyzed the frequency with which kinase-interacting proteins had interactions among themselves, as reported in BioGRID [Stark et al. 2006]. We found that, of the 425 interacting proteins, 12 interacted with other interacting proteins of the same kinase from Bio-

proteins [an example is shown in Supplemental Fig. S2A]. Thus, these results suggest that the AC/MS experiments identify interacting sets of proteins that are part of the same complex, whereas the protein array identifies binary interacting partners.

We analyzed our interaction sets to see whether they might be targets of phosphorylation of the interacting kinase using two methods. First, we compared the interaction data with the set of kinase substrates identified using similar protein microarrays [Ptacek et al. 2005]. We found only 34 protein targets in common between the interaction and phosphorylation data sets, indicating that these are mostly distinct sets of proteins; in addition, the phosphorylation data set generated from protein microarrays is likely incomplete. To further explore whether the kinase partners are potentially phosphorylated by their interacting kinase, we computationally examined each interacting partner for the presence of high-scoring phosphorylation motifs using the Mok et al. [2010] data set [see the Materials and Methods]. We found that 25% of the interacting proteins had high-confidence phosphorylation motifs for their partner kinase; this figure is significantly enriched over random [P-value < 10^-20], but is lower than that observed by Breitkreutz et al. [2010] [50%]. It nonetheless indicates that many interacting partners are likely to be kinase substrates. We do not expect 100% overlap, since interacting proteins identified by protein microarrays may contain divergent phosphorylation sites or contain domains that bind the kinase and are not phosphorylated. Markov clustering [van Dongen 2000] was performed on the BioGRID protein–protein interaction set [Stark et al. 2006] to determine cluster membership for kinase interactions with high-confidence phosphorylation motifs in both the Breitkreutz et al. [2010] and the protein microarray data sets [Supplemental Table S3]. In both cases, a majority of the kinases’ interacting partners belong to the same functional cluster, indicating that kinase interactions with high-confidence phosphorylation motifs are highly likely to function in the same cellular process.
Distinct and overlapping sets of interactions revealed by different members of kinase families

Yeast has a number of protein kinases that are closely related by sequence and/or function to one another. An example is the Nim1-like kinases Gin4, Hsl1, and Kcc4, which are functionally redundant in the cell cycle checkpoint that responds to defects in the peripheral cytoskeleton (Barral et al. 1999). We found that the Nim1-like kinases shared some targets, but the majority of targets are kinase-specific (Fig. 4A). One interesting example is Ygr017w, which was bound by Hsl1 and Gin4. Swe1, which is in the same pathway as Hsl1 and Gin4, phosphorylates Ygr017w (Ptacek et al. 2005). Nonetheless, the vast majority of Gin4, Hsl1, and Kcc4 have distinct targets, which is consistent with the unique phenotypes of the different kinases in pseudohyphal growth and cell cycle control (La Valle and Wittenberg 2001).

Analyses of the targets from very different kinase families revealed an unexpected result: Different types of kinases can often bind the same targets. One example is Akl1, in which 10 of 24 targets overlap with the 29 targets of Snf1 ($P < 1.15E-16$) [Fig. 4B]. Akl1 is involved in cytoskeletal function and Snf1 is involved in carbon metabolism; these two processes are themselves linked (see Costigan and Snyder 1994). Another example is Ipl1, for which nine targets bind one or more members of the Nim1 kinase family (Hsl1, Gin4, and Kcc4; $P < 1.20E-12$; two of these nine were validated to interact with Ipl1) [Fig. 4A]. Thus, distinct types of kinases often share common targets, suggesting that they may coordinate the same types of processes, either cooperatively or antagonistically. To our knowledge, this is the first report of different families of kinases sharing significant overlap in their binding of proteins involved in multiple cellular processes, which are expected to be coordinated.

Novel interactions with Kss1 suggest the filamentation pathway is associated with a diverse set of biological processes

Many interesting interactions were observed, as shown in Figures 5 and Supplemental Figure S3. One example is the Rio1 protein kinase involved in rRNA processing (Vanrobays et al. 2001). We found that Rio1 interacts with proteins involved in rRNA processing and translation; several of these interactions were validated [Supplemental Fig. S3]. Examples include Tif11 (a translation initiation factor), Rkm4 (a ribosomal lysine methyltransferase), Nop53 (a protein involved with the biogenesis of the 60S subunit of the ribosome), and Bfr1 (a component of mRNP complexes). Thus, Rio1 is likely to mediate its
functions, at least in part, through these components, and may help coordinate rRNA processing and translation.

One of the most interesting sets of interactions involved the Kss1 kinase, which controls filamentous growth and the pheromone response and has been highly studied (Cook et al. 1997; Erdman and Snyder 2001). Kss1 interacted with a diverse set of proteins on the proteome array, many of which were validated by co-IP/immunoblot analysis. These Kss1 interactions include two phosphatases [Ptc1 which is involved in the Hog1-osmosensing pathway] and Ptp1 [a phosphotyrosine-specific phosphatase with broad substrate specificity], a protein component of the CCR4–NOT transcriptional regulatory complex [Caf16], a component of the MIND kinetochore complex [Nsl1, which has been implicated in cell cycle progression], and two proteins with unknown functions [Ycl047c and Nba1]. As demonstrated below, Ycl047c promotes filamentation, and we renamed it Pof1 [for promoter of filamentation]. The complexes identified in our screen between Kss1 and this group of targets had not been identified previously using AC/MS or two-hybrid approaches, nor have they been directly implicated in filamentation signaling. However, these proteins are highly connected among themselves, and they also interact with other proteins implicated in filamentation, such as the phosphatase Glc7 (Fig. 5). Our results indicate that Kss1 operates with a diverse suite of components, such as those involved in transcriptional regulation, the Hog1 stress response, and the kinetochore/cell cycle progression.

We did not identify many known binding partners of Kss1 because they either were not present on the array [Ste7, Ste11, Ste12, and Tec1] or were present at very low abundance [Dig1 and Dig2, which were ~175-fold and ~11-fold less, respectively, than other binding proteins on the array]. Nonetheless, our ability to detect and validate many new Kss1-interacting partners indicates a much broader involvement in other pathways (see the Discussion).

**Kss1-interacting proteins operate in the filamentation pathway**

The results presented above suggest a highly interactive network between Kss1 and a variety of different cellular components. We examined whether any of the newly identified interacting partners operate in the same pathways as KSS1 using genetic tests. We specifically tested the validated targets from the Kss1 data sets for their ability to operate in the haploid invasive growth pathway, which requires Kss1 (Cook et al. 1997; Erdman and Snyder 2001). When haploid yeast cells are deprived of a fermentable carbon source, the cells adopt an elongated cellular morphology and invade agar (Cook et al. 1996; Lorenz et al. 2000). We tested whether the protein factors that interact with Kss1 in our screen affected filamentation and/or suppressed the filamentation defects of a kss1 deletion strain. *PTC1, CAF16, POF1, NSL1, and PTP1* [along with vector, *SNU71, CTA1*, and *RIM9*, which proved to be negative controls] were expressed from the *GAL1* promoter on a high-copy plasmid and transformed into wild-type (S1278b) and kss1Δ haploid yeast. Expression was induced by growth on galactose, and the ability of the cells to invade agar was examined (Fig. 6A). Wild-type cells invaded agar, whereas kss1Δ cells did not. Overexpression of *PTC1, CAF16, POF1*, and *NSL1* each partially suppressed the phenotype of kss1Δ cells, suggesting that they function independently of or downstream from Kss1. Overexpression of *SNU71, CTA1*, and *RIM9* in the kss1Δ background did not facilitate invasive growth [Fig. 6, data not shown]. Furthermore, microscopic examination of kss1Δ yeast overexpressing CAF16, NBA1, NSL1, and PTC1 revealed the elongated pseudohyphal cellular morphology, consistent with haploid invasive growth [Fig. 6B], negative control strains did not. The fact that *PTC1*, a negative regulator of the HOG1-osmosensing pathway, also operates downstream from *KSS1* indicates...
a further genetic link between these pathways [see the Discussion].

Interestingly, overexpression of PTP1 in wild-type cells suppressed haploid invasion of wild-type cells, suggesting that it is a negative regulator of filamentation [Fig. 6A]. PTP1 overexpression also suppressed pseudohyphae in the wild-type background [Fig. 6B]. Overexpression of genes in wild-type cells for the other target proteins (PTC1, CAF16, YCL047c, NSL1, SNU71, CTA1, NBA1, and RIM9) had minimal to no effect on invasion or morphology. Overall, our results suggest that Kss1 operates directly or indirectly through a diverse array of cellular activities, which include elements of the kinetochore and Ccr4–Not-regulated transcriptional/translational control. We also identified interactions with several important protein phosphatases that have a broad array of substrates in vivo [Fig. 6].

**Global interaction network and network modules**

The interactions identified in our screen were used to generate a first-generation protein microarray kinase physical interaction network [Fig. 7]. Raw kinase interactions were processed through a two-step learning algorithm to produce a final high-quality kinase interaction set [see the Materials and Methods, data analysis]. As noted above, almost half of the targets [198 out of 425] interact with multiple kinases, and further analysis of kinase–kinase interactions and those with common interacting proteins revealed that, overall, the kinases are highly connected with one another. In fact, there are more interactions with multiple kinases than those that interacted uniquely with a single kinase, consistent with the recent study by Breitkreutz et al. [2010]. However, our data are mostly unique relative to those of Breitkreutz et al. [2010], suggesting that the density of interactions is much higher than reported previously.

We subsequently determined that the protein microarray kinase interaction data set was enriched for network modules [Supplemental Table S4]. First, we used the Mok et al. [2010] data set to identify interactions due to kinase phosphorylation [kphos] or binary protein–protein interactions [kbin]. Second, we integrated our kinase interaction data [Fig. 7A] with transcription factor binding and protein interactions between kinase-interacting partners [Fig. 7B] to generate a “metanetwork” of 1374 interactions [Fig. 7C; Lee et al. 2002; Stark et al. 2006]. Searching the metanetwork resulted in the identification of six modules that were significantly enriched over random, shown in Figure 7D [see the Materials and Methods]. Modules such as two kinases binding a common protein (module 2) are suggestive of either scaffold protein or coordinate regulation. Others, such as modules 5 and 6, indicate that proteins bound by the same kinase are often bound by the same transcription factor, suggesting they are subjected to common regulatory mechanisms. A number of kinase–protein interactions from these modules were validated, as summarized in the legend for Figure 7.

**Gene Ontology (GO) analysis of protein–kinase interactions**

We also searched for enrichment of GO terms in the sets of all kinase targets [see Supplemental Table S5]; the cytoplasm was the only significant GO term in the cellular component GO category. We further analyzed the targets that interacted with the taxonomically related kinases to search for enrichment across biological processes, functions, and cellular components. The CAMK
study to date using proteome microarrays and identified we performed the largest protein–protein interaction novel protein–protein interactions. In the present study, vide a useful, rapid, and unbiased approach to identifying comprehensive network models. Protein microarrays pro-
egenerated accelerated the rate at which genetic and biochemical wide experiments with 

The information derived from genome- and proteome-

Identification of new interacting partners using protein microarrays

The interaction between Kss1 and Nsl1, a kinetochore component, was unexpected. When yeast undergoes haploid-invasive growth, there is an accompanying G2/M cell cycle delay [Kron et al. 1994], similar to ns1-16 and ns11-42 mutants [Scharfenberger et al. 2003]. Perhaps Kss1 functions with kinetochore components to help mediate this delay/checkpoint. These results suggest
links with many cellular processes, as might be expected for a key regulatory kinase that controls filamentation. Filamentation is activated upon nutrient stress responses, which would be expected to exert a global effect on cellular activities. The broad interactions of Kss1 with components of different cellular pathways are consistent with this global regulatory response.

Interestingly, our results also suggest cross-regulation of the Kss1 and Hog1 MAPK cascades through the activities of Ptc1. A requirement of Hog1 pathway activation is that the Kss1 pathway is rendered inactive and vice versa: Activation of the Kss1 pathway inactivates Hog1 (Shock et al. 2009). Ptc1 inactivates the osmosensing MAPK cascade by dephosphorylating Hog1 (Krantz, et al. 2009). Overexpression of PTC1 rescues invasive growth in kss1Δ cells, indicating that it also functions either downstream from or in parallel to the Kss1 pathway.

Conclusions

The myriad of interactions contributing to the complexities of eukaryotic cell signaling will require a wide variety of approaches to better understand these networks. Our study to identify a vast repertoire of protein–protein interactions that have not been found previously reveals many connections between cellular pathways. Our results are consistent with a highly connected web of interactions that coordinates cellular activities with intercellular and extracellular responses. Finally, since signaling components are conserved throughout eukaryotes, the results from our study are expected to be applicable to the understanding of signaling networks in other eukaryotes. MAPKs have been proposed to have an evolutionarily conserved role in the cross-regulation of cellular signaling pathways to effect global control in response to external stimuli (Weston

Figure 6. Phenotypic analysis of Kss1 target overexpression in the S1278b background reveals new signaling components contributing to haploid filamentous growth. (A) Yeast cells were transformed with 2μ pYES-DEST52 plasmids containing ORFs for the proteins that were validated (at their endogenous levels) by co-IP with Kss1. The effects of overexpressing target proteins on haploid-invasive growth were tested in kss1Δ and wild-type strains (see the Materials and Methods). The yeast strains labeled vector (empty vector) and RIM9 are negative controls (as were CTA1 and SNU71) [data not shown], and KSS1 overexpression serves as a positive control for filamentous growth. CAF16, NSL1, PTC1, and POF1 were sufficient to induce the filamentous pathway in the kss1Δ cells. The invasive growth phenotype was not observed for overexpression of PTP1 in wild-type cells compared with the negative controls. PTP1 overexpression inhibited the invasive growth phenotype in wild-type cells. (B) DIC images of yeast cells pre-induction and post-induction with 3XYEP/2% galactose in liquid culture exhibited the haploid pseudohyphal filamentous phenotype, in agreement with proteins that induce invasive growth. The kss1Δ yeast cells transformed with pYES-DEST52 high-copy CAF16, KSS1, NSL1, PTC1, and POF1 constructs all formed pseudohyphae compared with the vector control. The cells transformed with similar constructs for PTP1 and RIM9 did not form pseudohyphae. Cells transformed with pYES-DEST52 PTP1 suppressed pseudohyphal growth in the S1278b wild-type background. SNU71 and CFA1 overexpression had no effect on the filamentous/haploid-invasive growth phenotype in either the knockout or wild-type strains [data not shown].
et al. 2002; Breitkreutz et al. 2010), and our studies with Kss1 strongly support this concept.

Materials and methods

Kinase-V5 probe preparation

Full-length kinase ORFs were cloned into the pYES-DEST52 gateway vector [Invitrogen] from constructs either obtained from the MORF collection or cloned de novo as described previously by Gelperin et al. (2005). The pYES-DEST52 plasmid contains an LR cloning site flanked by a V5-His6X C-terminal tag under control of a GAL1 promoter. The resulting kinase-V5-His6X fusion protein has a His6X tag for protein purification and a 14-amino-acid V5 epitope tag (GKPIPNPLLGLDST) for antibody detection. Sequence-verified plasmids were transformed into yeast (Y258: MATa, pep4-3, his4-580, ura3-52, leu2-3, 112) using a standard lithium acetate transformation protocol as described previously (Kumar et al. 2000), and the resulting strains were used to purify the V5-kinase fusion proteins (see Supplemental Table S7 for strains used in this study). Yeast cultures were grown overnight at 30°C in 6 mL of Sc-Ura/2% dextrose and diluted to OD600 = 0.1 the next day in 400 mL of Sc-Ura/2% raffinose. Cultures were grown at 30°C to OD600 = 0.6 and induced with 3XYEP [3% yeast extract, 6% peptone, 6% galactose] for 6 h, followed by centrifugation, washing with ice-cold water, and freezing overnight at −80°C. Crude lysates were prepared by lysing cells twice with 350 mL of lysis buffer [1× PBS at pH 7.4, 150 mM NaCl, 2 mM MgCl2, 0.5 mM DTT, 0.1% Triton X-100, 0.3% sarkosyl, 0.5% deoxycholate, 1 mM PMSF, 1× complete EDTA-free protease inhibitors [Roche], 1:100 dilution phosphatase inhibitor cocktail 1 [Sigma], 100 μM sodium orthovanadate] by shaking for 6 min in a paint shaker [SG-HD, Harbell] at 4°C with 250 μL of 0.5-mm glass beads [BioSpec Products], followed by centrifugation at 2500g for 5 min. Between 2 g and 10 g of cell pellets was used to normalize kinase-specific protein yield [microgram per milliliter] from extraction. The lysate was then centrifuged at 150,000g for 1 h at 4°C and passed through a nickel (Ni2+) sepharose column (His Spintrap, GE Life Sciences) to bind the kinase-V5-His6X fusion protein. The column was washed three times with wash buffer [1× PBS at pH 7.4, 500 mM NaCl, 2 mM MgCl2, 0.1 mM Trition X-100, 0.3% sarkosyl, 0.5% deoxycholate, 1 mM PMSF, 500 mM imidazole, 20% glycerol] and the protein was eluted with elution buffer [1× PBS at pH 7.4, 500 mM NaCl, 2 mM MgCl2, 0.1% Trition X-100, 0.5 mM DTT, 500 mM imidazole, 20% glycerol].

Protein microarray screen

Commercial yeast protein microarrays [Invitrogen] containing ~4200 full-length GST fusion proteins spotted in duplicate were
used in this screen (Zhu et al. 2001). Arrays were blocked for 1 h at 4°C with shaking in blocking buffer [1× PBS at pH 7.4, 1% BSA, 0.1% Tween-20]. Purified kinase-V5 probe was then diluted in probing buffer (1× PBS, 5 mM MgCl₂, 0.5 mM DTT, 0.05% Triton X-100, 1% glycerol, 1% BSA) augmented with 150 mM NaCl and 200 μM ATP. After blocking, 90 μL of the diluted kinase-V5 was loaded onto the microarray membrane at a final concentration of ∼10–50 μg/mL with a pipette tip, and the membrane was covered using a Lifterslip (Electron Microscopy Sciences) during probing. The arrays were then incubated in a humidified chamber for 1.5 h static (no shaking) at 4°C, followed by washing three times for 1 min (static) in 25 mL of probe buffer on ice. The arrays were then probed with 90 μL of anti-V5-AlexaFluor647 antibody [Invitrogen] at a concentration of 260 ng/mL diluted in probing buffer, covered with a Lifterslip as before, and placed back into the humidified chamber for 30 min static at 4°C. Following antibody detection, the arrays were washed in 25 mL of probe buffer as before and spun at 800g for 5 min in a tabletop centrifuge to dry. The protein-protein interactions were detected using a 4200AL microarray scanner (Axon Instruments). The image file was then processed using GenePix software [Axon Instruments] to determine signal intensity of the bound kinase probe detected by the anti-V5 AlexaFluor647-conjugated antibody.

Data analysis

An algorithm called ProCAT (Zhu et al. 2006) was used to identify proteins bound to the kinase probe. Briefly, the program normalizes the background by considering a local window to account for spatial and other artifacts. After a set of filtering steps, the signal of each spot is computed by accounting for the background and foreground signal of the neighboring spots in a three-by-three-feature window. To detect positive spots, the normalized signal is then compared with the neighboring spots in a nine-by-nine-feature window. The number of normalized standard deviations (SDs) above the median signal in the neighboring spots was used to set the threshold to detect positive signals above background. ProCAT has been shown to improve the signal-to-noise ratio of functional protein microarrays (Zhu et al. 2006). We then implemented a two-step learning algorithm to determine the optimal thresholds for the high-quality kinase interactions. For the training set, we used all known kinase interactions in the literature from the BioGRID database as gold-standard positives, and protein pairs with different subcellular localizations as the gold-standard negatives. We employed the naive Bayesian method as the learning kernel. At the first step, we determined the optimal normalized SD value of intervals for high-quality kinase interactions. Then, in the second step, we determined the specific interactions among the high-quality ones from the first step. The optimal conditions were determined to be a signal-to-noise interval that corresponded to targets validated by co-IP/immunoblot corresponding to ProCAT normalized SD values (5 ≤ SD ≤ 20)]. Interactions bound by this interval were filtered by the number of kinases that interact with the targets (≥10 kinases) and subcellular localization. The overlapping targets between kinases formed the hubs of interconnectivity, connecting the edges of each kinase to its corresponding targets.

Interaction validation by co-IP

Co-IP strains were generated by introducing a 13XMyt genomic tag onto the C terminus of each kinase in the BY4741 (MATα his3Δ1 leu2Δ0 ura3Δ0 met15Δ0) yeast strain containing a genomic TAP tag (Open Biosystems) for each of the interacting proteins. Additionally, negative control strains were generated for all kinases tested that contained the genomically tagged 13XMyt kinase without TAP-tagged proteins to control for cross-reactivity of the anti-Myc antibody. A genomic 13XMyt tag was generated using pFA6a-13Myc-KanMX6, and the reverse primer 5′- [gene-specific sequence] GAATTCGAGCTCCTTAAAC-3′ and forward primer 5′- [gene-specific sequence] CCGATCCCCGGTTAAT TAA-3′ (Longtine et al. 1998) were used generating genomic BY4741 gDNA as a template and Pfx platinum polymerase [Invitrogen].

The co-IP strains were used to inoculate 10-mL YPAD starter cultures overnight, followed by inoculating 500-mL YPAD cultures to OD600 = 0.1. Cultures were grown for 7 h at 30°C, harvested by centrifugation, washed with cold water, and frozen at −80°C. The next day, 2 g of cell pellet for both the control strain [TAP tag] and the experimental strain (co-IP strain) was parsed over four separate lysis tubes containing 0.5 g of pellet (2 g total), and each was used to prepare the lysates for co-IP.

Lysates were obtained by lysing each of the 0.5-g cell pellets twice in 400 mL of RIPA buffer (0.5 M Tris-HCl at pH 7.4, 1.5 M NaCl, 2.5% deoxycholic acid, 10% NP-40, 10 mM EDTA, 1% PMSE, 1× complete protease inhibitors [Roche], 1:100 dilution phosphatase inhibitor cocktail 1 [Sigma], 100 μM sodium orthovanadate) by shaking for 1 min, followed by 2 min on ice in a Fastprep [MP Biomedicals] at 4°C with 250 μL of glass beads (0.5 mm), BioSpec Products) for a total of three intervals. The homogenate was centrifuged at 14,000 rpm using a microfuge, and the supernatant for each sample was collected. This process was repeated for a total of two extractions, and the lysates from 2 g of cell pellets were combined in a 15-mL conical tube for co-IP. The co-IPs were performed by incubating 25 μL of dynabeads [Invitrogen] conjugated with rabbit IgG (Sigma) with the lysates for 1.5 h on a nutator at 4°C. The dynabeads were then washed five times for 10 min with ice-cold wash buffer (50 mM Tris-HCl at pH 7.4, 0.1% Triton X-100, 150 mM NaCl), and beads were boiled at 95°C in 65 μL of 2× SDS sample buffer.

The samples were then run out on 10% NuPAGE gels [Invitrogen] and transferred to nitrocellulose membrane for immunoblotting. The membranes were incubated with 1:1000 dilution of clone 9E10 anti-Myc antibody [Millipore] in 3% milk/TBS overnight at 4°C. The next day, the membranes were washed, followed by incubation for 1 h at room temperature with 1:10,000 dilution of secondary anti-mouse antibodies conjugated to HRP [GE Biosciences] and exposed to Picowest [Pierce] for detection. Co-IP/immunoblot-validated kinase protein-protein interactions were scanned from multiple experiments to create the panels shown in Figure 2. Those interactions that did not validate by co-IP are shown in Supplemental Table S2.

Kinase phosphorylation and binary protein interactions

To determine which kinase-binding mechanism gave rise to an interaction found by the protein array, S. cerevisiae phosphosite data from the literature (Ficarro et al. 2002; Gruhler et al. 2005; Li et al. 2007; Albuquerque et al. 2008; Beltrao et al. 2009; Stark et al. 2010) were incorporated into the protein array-derived kinase network. Kinase position weight matrices [PWMs] from Mok et al. (2010) were used to score every phosphosite. For each kinase PWM, the best score for each phosphoprotein was used to establish a best score background distribution. We assumed the background distribution follows a Student’s t-distribution and applied a stringent threshold of P < 10−20 to annotate an interaction as a kinase-phosphorylating interaction. Interactions that failed the threshold were annotated as kinase-binding interactions.
Network modules analysis

The kinase network was augmented with transcription factor binding data [Lee et al. 2002]. BioGRID protein interactions version 3.1.71 was used to attain interactions between kinase and transcription factor protein targets [Stark et al. 2006]. The interaction network was mined for network modules by a custom in-house C++ program. The program uses a motif-mapping approach by starting from a set of directed nonisomorphic graphs provided by gools toolset version 2.4 [McKay 1981]. A depth-first approach was used to find each motif in the network. To establish network module significance, 500 random networks were generated. Each class of network interaction was randomized independently, while maintaining each protein’s in degree and out degree. Since the kinase network was comprised of two interaction types, a similar ratio of the two interaction types was maintained in the random networks. The one-sided Wilcoxon signed-rank test was used to determine network module significance.

Phenotypic analysis of Kss1 targets

The ORFs coding for the proteins confirmed to interact with Kss1 by co-IP/immunoblot were cloned into the pYES-DEST52 gateway plasmid [Invitrogen] as described previously (Gelperin et al. 2005) and transformed into the yeast strain Y825 [MATa ura3-52 leu2-0] derived from the filamentous Σ1278b background [Jin et al. 2008]. The pYES-DEST52 plasmid contained a GALL1-inducible promoter to drive expression of the cloned ORF as a V5-His6X fusion protein. The Y825 cells transformed with pYES-DEST52 plasmids containing the ORFs identified in the screen were grown overnight in Sc-Ura/2%/2% dextrose and were used to inoculate 5-mL Sc-Ura/2% raffinose/0.1% dextrose cultures grown in 5 mL of Sc-Ura/2%/0.6. The cultures were induced for an additional 7 h at 30°C, plated on Sc-Ura/2% galactose/0.2% raffinose agar plates, and incubated for 3 d at 30°C. The presence of the peripheral cytoskeleton in yeast.


