Ste11p MEKK signals through HOG, mating, calcineurin and PKC pathways to regulate the FKS2 gene

Wang et al.
Ste11p MEKK signals through HOG, mating, calcineurin and PKC pathways to regulate the FKS2 gene

Xiaoyan Wang1,2, Mark A Sheff1, David M Simpson1,3 and Elaine A Elion1*

Abstract

Background: The S. cerevisiae MAPKKK Ste11p, a homologue of mammalian MEKK1, regulates three MAPK cascades for mating, invasive growth and osmotic stress and provides functions that are additive with the cell wall integrity pathway. Cell wall integrity requires the FKS2 gene that encodes a stress-induced alternative subunit of beta-1,3 glucan synthase that is the target of echinocandin 1,3- beta glucan synthase inhibitors. The major signal transduction pathways that activate transcription of the FKS2 gene include the cell wall integrity and calcineurin pathways, and the Ste11p pathway.

Results: Here it is shown that catalytically active Ste11p regulates FKS2-lacZ reporter genes through Ste12, calcineurin/Crz1p- and PKC pathways and the high osmolarity pathway. Ste11p stimulated the cell wall integrity MAPK Mpk1p (Erk5 homologue) and FKS2 independently of the mating pathway. Ste11p regulated FKS2 through all known and putative substrates: Pbs2p MAPKK, Ste7 MAPKK, Cmk2p calmodulin dependent kinase and Ptk2p kinase. Ste11p increased the expression level of Cmk2p through transcription-dependent and -independent mechanisms.

Conclusions: The data suggest Ste11p regulates the FKS2 gene through all its known and putative downstream kinase substrates (Pbs2p, Ste7p, Cmk2p, and Ptk2p) and separately through Mpk1p MAPK. The patterns of control by Ste11p targets revealed novel functional linkages, cross-regulation, redundancy and compensation.

Background

Ste11p encodes a mitogen activated protein kinase kinase (MAPKKK) that activates conserved MAPK pathways controlling mating, high osmolarity glycerol (HOG), invasive growth and the FKS2 gene in the cell wall integrity pathway [1-3]. The catalytic domain of Ste11p is most homologous to mammalian MEKK1 [4]. During mating, high osmolarity growth and invasive growth, Ste11p is phosphorylated and activated by Ste20p, a p21 activated kinase that binds to Cdc42p. Once activated, Ste11p has the potential to phosphorylate and activate two MAPKKs, either Ste7p for mating and invasive growth pathways, or Pbs2p for the high osmolarity sensing pathway [5]. The activation of Ste7p leads to activation of Fus3p and Kss1p MAPKs that activate shared and unique transcription factors among other substrates [6,7]. Pbs2p activates Hog1p MAPK, which also activates transcription factors and other substrates [8,9].

Prior work suggests that Ste11p MAPKKK signals through MAPK Kss1p to positively regulate cell wall integrity [2]. This analysis revealed that Ste11p has functional redundancy with the Bck1p MAPKKK in the PKC pathway and can activate the expression of the FKS2 gene [2]. FKS2 encodes a stress induced beta-1,3 glucan synthase subunit similar to constitutively expressed FKS1 that is important for cell wall integrity under conditions of cell wall stress [1]. The Fks1p and Fks2p subunits of glucan synthase are major therapeutic targets of anti-fungal inhibitors and acquire resistance mutations during fungal infections in people [10,11]. FKS2 is expressed at low levels and is upregulated under conditions of stress, higher temperature, when FKS1 is mutated, reduced carbon source [1], and when glycosylation is disrupted [2].
The calcineurin and protein kinase C and mating pathways are the major signaling pathways that regulate FKS2 expression and cell wall integrity. Calcineurin (Cna1p catalytic subunit and a regulatory subunit Cnb1) is activated by increases in intracellular calcium by influx of extracellular calcium through a Mid1/Cch1-Ca\(^{2+}\) channel [1,12,13]. Many of its responses are mediated through the transcription factor Crz1p. Calcineurin dephosphorylates Crz1p/Tcn1p, leading to nuclear localization that activates many genes [14]. During conditions of high extracellular calcium or pheromone, the calcineurin pathway signals Crz1p/Tcn1p to bind the FKS2 promoter at a calcineurin-dependent response element (CDRE) site within residues -928 to -706 [1,14]. During polarized growth, cell wall damage or temperature stress, the cell wall integrity pathway plasma membrane sensors signal through Rhop1 to Pkc1p, which activates Bck1p MAPKKK to activate Mkk1/2p MAPKK, which signals Slt2p/Mpk1p MAPK and Mlp1p pseudokinase to activate several transcription factors. Slt2p/Mpk1p, its human homolog Erk5p, and Mlp1p activate FKS2 through the cell cycle transcription factor SBF (Swi4p/Swi6p) at a SCB consensus site at -385 to -391 [15]. There are three potential Ste12 TGAAACA binding sites starting at -894 to -899, but Ste12 has not been found to bind the FKS2 promoter in two independent CHIP studies. Mating pheromone induces FKS2 at late times [16], supporting the possibility that it is due to secondary events including activation of the calcineurin and PKC pathways from calcium influx and polarized growth.

One gene that is induced by Crz1p is CMK2, which encodes one of two redundant calmodulin dependent kinases, possibly an orthologue of human CaM kinase II [17,18]. Cmk2p prevents death of calcineurin-deficient cells under conditions of low calcium [19]. A genomic in vitro screen for kinase substrates suggests that Cmk2p could be a direct substrate of Ste11p [20], Ptk2p was the only other kinase substrate of Ste11p identified [20]. Ptk2p is a putative serine/threonine protein kinase involved in regulation of plasma membrane ATPase, spermine and spermidine transport [21-23] and stimulates the PKC pathway MAPK (Pbs2p), Cmk2p, and Ptk2p in addition to Ste7p MAPKK and stimulates the FKS2 gene independently of Ste12p. These findings reveal new functional links between Ste11p downstream targets to calcineurin and PKC pathways and illustrate a high level of signaling flexibility.

**Results**

**Ste11p is required for basal Slt2p/Mpk1p activation**

Prior work indicated that Ste11p is functionally redundant with the Bck1p MAPKKK for cell integrity during vegetative growth and could basally activate the cell wall integrity target FKS2 gene using a well established FKS2 (-928 to +6)-lacZ reporter gene [2]. To test the possibility that Ste11p regulates cell integrity through activation of the PKC pathway during vegetative growth, we assessed the basal level of active Mpk1p in three yeast strains backgrounds, S288c, W303a and Σ1278b by probing whole cell extracts with a mammalian phospho-p42p44 antibody that cross-reacts with S. cerevisiae Slt2p/Mpk1p [25]. We compared the level of active Slt2p/Mpk1p in exponentially dividing cells (Figure 1A, a longer exposure that also shows active Kss1p is in Additional file 2, Figure S2). The level and gel mobility of basal active Slt2p/Mpk1p varied tremendously. W303a and S288c had an equivalently sized short form of Slt2p/Mpk1p, whereas Σ1278b had a longer form. W303a had the highest basal level of active Slt2p/Mpk1p, whereas S288c had the lowest basal level, indicating variability at the level of pathway flux. The basal level of active Slt2p/Mpk1p was clearly reduced in a ste11Δ, mutant (Figure 1B, S288c background shown). Less obvious partial reductions in the level of active Slt2p/Mpk1p were detected in ste4Δ, ste5Δ, ste50Δ, fus3Δ, kss1Δ, and ste5Δ fus3Δ strains (Figure 1A-1B, and data not shown). These observations suggested that Ste11p had a more critical role in regulating the PKC pathway than other mating pathway components.

**STE11-4 activates the FKS2 promoter through -928 to -706 and -706 to -540**

The FKS2 promoter is complex with multiple promoter elements that include the calcineurin-dependent response element (CDRE), Swi4/Swi6 SBF element, glucose repression elements and potential pheromone response elements [1,14]. We examined how Ste11p regulates the FKS2 gene by comparing the ability of hyperactive **STE11-4** to stimulate established **FKS2-lacZ** promoter truncations of the promoter: FKS2(-928+6)-lacZ (pDM5; [26]), FKS2(-928+1)CYC1-lacZ [14], FKS2(-706+1)CYC1-lacZ [14] and FKS2(-540-375) CYC1-lacZ (p2052; [15]). The FKS2 -928 to +1 promoter has an SBF site at -385 CACGAAA-391 that binds a
Swi4/Swi6/Mpk1 complex (assayed in YPD or supplemented SD [15]), the CDRE (calcineurin-dependent response element) within -762 to -705 (that includes a Crz1p -740 CAGTCGGTGGCTGTGCGCTTG-760 element that supports LacZ expression when present in 2 copies and assayed in synthetic medium containing 200 mM CaCl2 and ammonium chloride substituted for ammonium sulfate; [27]), a glucose repression element overlapping two putative Mig1p repressor consensus sites at -847 and -785 [15], and one of three Ste12p consensus binding sites, TGAAACA (-899-894). Consistent with prior work [2], STE11-4 activated both FKS2 (-968 to +6)-CYC1-lacZ and (-928 to +1)-CYC1-lacZ in 1X synthetic medium containing 2% dextrose and lacking uracil ([(Figure 2A, B; negative control ("neg cn") is pLGΔ-178 pCYC1-lacZ)]. STE11-4 also activated the FKS2(-706 to +1)CYC1-lacZ promoter that contains the SBF recognition site that is positively regulated by Mpk1p/Slt2p and Mlp1 (Figure 2A). STE11-4 activation was three-fold greater on the FKS2(-928 to +1) promoter than on -706 to +1, consistent with the presence of a putative Ste12p binding site and the CDRE element. STE11-4 failed to stimulate the FKS2(-540 to -375)-lacZ reporter that contains the SBF consensus site (-385 CACGAAA -391) that is regulated by Swi4p/Swi6p/Mpk1p/Mlp1p complexes and stimulated by...
temperature stress and congo red [15]. The absence of stimulation of the SBF site is consistent with STE11-4 and mating pheromone-induced inhibition of Cdc28 cyclin dependent kinase [28]. Thus, under our growth conditions, Ste11-4p positively regulated FKS2 reporter genes through at least two promoter elements, from -928 to -706 containing a Crz1p binding site, two potential Mig1 binding sites and 1 potential Ste12p binding site, and another element between -706 and -540, but did not stimulate transcription through the Mpk1p/ Mpl1p responsive SBF site within -540 to -375.

**STE11-4, α factor, and Ste12 differentially regulate FKS2-lacZ**

Mating pheromone stimulates FKS2 gene expression at late times, with optimal expression occurring 90 minutes after α factor addition in YEPD medium [16] and less obvious induction in synthetic medium [27]. α factor
induction of genomic *FKS2* mRNA is blocked by *FKS2Δ* [16] and a *cze1Δ/ten1Δ* mutation [16,27]. We compared the ability of α factor to stimulate *FKS2Δ* (-928 to +1)-lacZ and *FKS2* (-706 to +1)-lacZ and two Ste12-dependent reporter genes *FUS1-lacZ* and *PRM5-lacZ*. In a wild type *STE11* background, α factor addition caused only a modest 2-fold increase in expression of the *FKS2* (-928 to +1)-lacZ reporter and no effect on the *FKS2* (-706 to +1)-lacZ reporter. Much larger increases occurred for *PRM5-lacZ* (20-fold) and *FUS1-lacZ* (>100-fold) reporter genes that harbor multiple Ste12p consensus binding sites (Figure 2C). In wild type, *far1Δ*, and the *STE11*-*4* *far1Δ* background (which has reduced levels of active Cin2p/Cdc28p compared to wild type [28]), a factor caused little increase in *FKS2* (-928 to +1)-lacZ levels (i.e. 0.2-<2-fold). The *far1Δ* strain (which has increased basal levels of G1 cyclin dependent kinases compared to wild type [28]), had lowest basal levels of *FKS2*- (-928 to +1)-lacZ and blocked α factor (Figure 2C,E). This contrasted up to >100 fold increases in *FUS1-lacZ* and *PRM5-lacZ* (Figure 2C-E).

By comparison, α factor stimulated the pDM5 *FKS2* (-968 to +6)-lacZ approximately 4-fold in a different wild type background [26]. Thus, regulation of reporter genes with -928 and -706 cut offs was relatively independent of α factor, consistent with only one Ste12p TGAAACA site very close to the 5’end of the -928 *FKS2-lacZ* constructs near the CDRE.

The absence of a strong α factor effect suggested that Ste12p might not be required for *STE11-4* activation of *FKS2*. To determine whether the Ste12p transcription factor was required for Ste11p activation of *FKS2*, we tested the effect of a ste12Δ mutation in *STE11-4 far1Δ* strains treated or not with α factor compared with that in no ste12Δ mutation strains (Figure 2E, F). The ste12Δ null mutation blocked the ability of *STE11-4* to activate both the *FKS2* (-928 to +1)-lacZ and *FKS2* (-706 to +1)-lacZ reporter genes similar to the blocks that occurred for *PRM5-lacZ* and *FUS1-lacZ* reporter genes. The effect was similar to that seen for α factor. Thus, Ste12p is critically important for *STE11-4* to activate the *FKS2-lacZ* reporter genes, but its function is likely indirect.

**Ste12 stimulates the expression of *FKS2* (-928 to +1)-lacZ but not *FKS2* (-706 to +1)-lacZ**

A ste12Δ mutation blocks the expression of pheromone response pathway genes that lead to activation of Ste11p, Ste7p and Fus3p and Kss1p MAPKs. Over expression of Ste12p can bypass positive and negative control by Fus3p and Kss1p MAPKs. We examined whether a GAL1prom-STE12 gene would encode the expression of the *FKS2* reporter genes in the absence of calcineurin and mating pathway components. Cells were shifted from dextrose medium to raffinose medium and finally to galactose medium to induce expression of GAL1prom-STE12 (Materials and Methods). *FKS2* gene expression is derepressed in poor carbon sources including raffinose and galactose, most likely through Snf1p kinase inhibition of Mig1p, which may bind consensus sites at -847 and -785 [14].

Ste12p stimulated the expression of *FKS2* (-928 to +1)-lacZ ≈5-fold in wild type, ste11Δ, fus3Δ kss1Δ and *far1Δ* strains, but not if the strain lacked the Cnb1 regulatory subunit of calcineurin (compare wild type and *cnb1Δ*, Figure 3A, C). In sharp contrast, GAL1p-STE12 did not activate the expression of *FKS2* (-706 to +1)-lacZ to a great extent (Figure 3B, C), although *STE11-4* was able to stimulate (Figure 2). Thus, Ste12p activated the *FKS2-lacZ* promoter by a mechanism involving the calcineurin pathway, suggesting it may mimic the late onset of activation by mating pheromone that leads to an increase in calcium influx that activates the calcineurin pathway CDRE element. In contrast, Ste11-4p activated the *FKS2* promoter through the CDRE element as well as a second element within -706 to +1.

**Ste11ΔNp activates the *FKS2* promoter from -928 to +1 through Pbs2p and Ste7p**

It was unlikely that the block in expression of the -928 and -706 *FKS2-lacZ* constructs in the *STE11-4* ste12Δ *far1Δ* strain was the result of a direct effect of Ste12p on *FKS2* expression. However, the activity of Ste11-4p requires proteins that are regulated by Ste12p, such as Ste5p; for example, a dominant active STE5hyp1 gene also increased the level of active Mpk1 in anti-active MAPK westerns (data not shown). To bypass dependence on upstream regulators of Ste11p, we used a *STE11ΔN* allele (pGAL1-STE11ΔN) that lacks the N-terminal regulatory domain that represses the catalytic kinase domain and also binds Ste5p scaffold, Ste50p and Sho1p [29-31]. The *STE11ΔN* mutation largely, but not completely, bypasses a requirement for Ste20p phosphorylation to be active. Ste11ΔNp activates Kss1p and more weakly activates Fus3p due to the absence of Ste5p binding to link Ste7p to Fus3p [32], but can activate Pbs2p MAPKK down to Hog1p efficiently [33].

We tested dependence on Pbs2p and Ste7p MAPKKs that are activated by Ste11p. Strikingly, in a pbs2Δ strain, basal *FKS2* (-928 to +1)-lacZ expression decreased and remained greatly decreased in the presence of *STE11ΔN* compared to the wild type strain. In contrast, a ste7Δ mutation had little effect on pGAL1-STE11ΔN induced expression of *FKS2* (-928 to +1)-lacZ. However, the level of *FKS2* (-928 to +1)-lacZ was lower in a ste7Δ pbs2Δ double mutant than in a pbs2Δ single mutant (compare Figure 3D, G), revealing contribution from Ste7p. Thus, Ste11ΔNp regulates the *FKS2* promoter gene through Pbs2p and Ste7p, but Pbs2p is more crucial.
Figure 3 STE12 and STE11ΔN activation of FKS2 promoter in mating and HOG pathway mutants. GAL1prom-STE12 over expressed in presence of FKS2 (-928 to +1)-lacZ (A), FKS2 (-706 to +1)-lacZ (B) and Δ-178 pCYC1-lacZ (C). GAL1prom-STE11ΔN over expressed in presence of FKS2 (-928 to +1)-lacZ (D), FKS2 (-706 to +1)-lacZ (E), PRM5-lacZ (F) and Δ-178 pCYC1-lacZ (G) and FKS2(-928 to -706)-lacZ (H). Strains were pregrown in raffinose- dextrose medium and then in galactose medium for 6 hours at 30°C. Data were expressed as mean ± SE of at least three independent experiments. Statistical significance was computed by the unpaired Student's t test. *p < 0.05, **p < 0.01.
Fus3p activated FKS2 (-928 to +1)-lacZ expression whereas Kss1p inhibited

We tested whether Ste11p required Fus3p and Kss1p MAPK targets of Ste7p, to stimulate FKS2 (Figure 3D, G). Fus3p and Kss1p have many targets including Ste12p and its repressors Rst1p/Dig1p and Rst2p/Dig2p. In a fus3A strain, the basal level of the FKS2 (-928 to +1)-lacZ reporter gene decreased to the same degree as in a ste12A strain. Fus3p was required for STE11ΔN to activate FKS2 (-928 to +1)-lacZ expression; the block in the fus3A strain was similar to that of the ste12A mutant. In contrast, the kss1Δ mutation caused a modest decrease in expression on its own and, conversely, restored expression in the fus3A mutant background (Figure 3D, G compare fus3 kss1 with fus3). Therefore, Fus3p and Ste12p activated FKS2 (-928 to +1)-lacZ whereas Kss1p had more inhibitory effect than positive. Ste11ΔNp activated the FKS2 gene via Pbs2p in the absence of Kss1p and Fus3p or Ste7p. A pbs2Δ mutation reduced the level of STE11ΔN activation of the FKS2 (-928 to +1)-lacZ reporter gene in a kss1Δ fus3Δ strain similar to in a ste7Δ (Figure 3D, G). Therefore, Ste11ΔNp regulates the FKS2-lacZ gene through both Ste7p MAPKK and Pbs2p MAPKK, with Pbs2p signaling being more significant and Ste7p positive signaling being dependent on Fus3p, Kss1p and Ste12p.

Ste11ΔNp activates FKS2 (-706 to +1)-lacZ through Pbs2p and Fus3p and Kss1p, whereas Ste7p blocks signaling by Pbs2p

STE11ΔN was strictly dependent on Pbs2p and Fus3p/ Kss1p to stimulate the FKS2 gene in the absence of the CDRE element. All three single deletions, pbs2Δ, fus3Δ and kss1Δ, decreased FKS2(-706 to +1)-lacZ expression with nearly complete blocks occurring with pbs2Δ and fus3Δ (Figure 3E, G). In sharp contrast, Ste7p had primarily negative function in regulating the FKS2 (-706 to +1)-lacZ reporter gene, with 3-fold increase in lacZ level occurring in the ste7Δ strain compared to wild type (Figure 3E). The increased signaling in the absence of Ste7p was primarily dependent on Pbs2p, and on Fus3p/Kss1p (Figure 3E, i.e. the level of FKS2(-706 to +1)-lacZ for pbs2Δ ste7Δ is equivalent to ste12Δ and wild type strains, whereas the pbs2Δ fus3Δ kss1Δ strain is even lower). Thus, we can define two separable branches of control that are inhibited by Ste7p, one through Pbs2p and another through Fus3p and Kss1p. Loss of Ste7p may increase the ability of Ste11ΔNp to signal through Pbs2p and may also confer catalytically inactive functions to Fus3p and Kss1p.

The relative contributions of Ste7p to Ste11ΔNp stimulation of the FKS2(-928 to +1)-lacZ and FKS2(-706 to +1)-lacZ reporters suggested Ste11p/Ste7p’s major positive regulatory function was within the -928 to -706 region that responds to pheromone and its major negative regulatory function was between -706 to +1. The analysis with STE11-4 (Figure 2A) suggested a target region between -706 to -540 rather than the -385 CAC-GAAA-391 site that binds the SBF-Mpk1 complex [15].

Ste11ΔNp exhibits little dependence on Bni1p and Ste20p

Bni1p and Ste20p are required for mating morphogenesis and pheromone activation of the PKC pathway and filamentation induced by Kss1p [1,33,34]. In both bni1Δ and ste20Δ strains, the basal level of the FKS2 (-928 to +1) reporter gene was the same as wild type strain and the level induced by Ste11ΔNp was reduced by ~25% compared to the wild type control (Figure 4A, D). Bni1p and Ste20p were required for full activation of FKS2 by Ste11ΔNp, but their contribution was much less than that of Ste7p, Fus3p, Kss1p and Pbs2p. There was no difference in expression of FKS2 (-706 to +1) reporter gene for wild type, ste20Δ and bni1Δ strains with STE11ΔN (Figure 4B, D). Thus, the STE11ΔN mutation largely bypasses a dependence on morphogenesis and filamentation to induce the FKS2 reporter genes.

Ste11ΔNp activated the FKS2 promoter from -928 to +1 through cell wall integrity and calcineurin pathways

We determined whether Ste11ΔNp required the calcineurin and cell wall integrity pathways to stimulate FKS2(-928 to +1)-lacZ and FKS2(-706 to +1)-lacZ genes. Prior work demonstrated functional redundancy between Bck1p and Ste11p (based on synthetic sickness of bck1Δ ste11Δ double mutants [2]). In the course of this analysis, we found that a bck1Δ strain transformed with the STE11ΔN plasmid grew much better than bck1Δ cells without STE11ΔN, based on faster and greater density of growth in streak outs on SC selective plates (data not shown). This result supports functional redundancy between Bck1p and Ste11p [2] being mediated by targets of Ste11ΔNp.

The calcineurin pathway was required for Ste11ΔNp to induce expression of the FKS2 (-928 to +1)-lacZ reporter gene, as lacZ induction was blocked by mutations in calcineurin cnb1Δ and the target transcription factor crz1Δ (Figure 4E, H). Bck1p was also required for Ste11ΔNp to activate the FKS2 (-928 to +1) reporter gene, as was Mpk1p (Figure 4A, D). The mpk1Δ mutation reduced expression somewhat less than the bck1Δ mutation, presumably due to MLP1 functional redundancy [15]. The addition of an osmotic support, KCl, had no effect on the ability of Ste11ΔNp to support better growth or induce expression of the FKS2 (-928 to +1)-lacZ reporter gene, ruling out a trivial explanation of cell lysis for reduced expression in the bck1Δ strain or a block in signaling through the HOG pathway.
Figure 4 Effect of mutations in PKC and calcineurin pathways and in CMK2 and PTK2 on Ste11ΔNp stimulation of reporter genes

STE11ΔN activation of FKS2 (-928 to +1)-lacZ (A), FKS2 (-706 to +1)-lacZ (B), PRM5-lacZ (C) and Δ-178 pCYC1-lacZ (D) in cell wall integrity and filamentous pathway mutants. STE11ΔN activation of FKS2 (-928 to +1)-lacZ (E), FKS2 (-706 to +1)-lacZ (F), PRM5-lacZ (G) and Δ-178 pCYC1-lacZ (H) in calcineurin pathway mutants. Data were expressed as mean ± SE of at least three independent experiments. Statistical significance was computed by the unpaired Student’s t test. *p < 0.05, **p < 0.01.
Ste11ΔNp-induced expression of FKS2 (-706 to +1)-lacZ and PRM5-lacZ was also dependent upon both Cnb1p/Crz1p and Bck1p/Slt2p/Mpk1p pathways (Figure 4F, G, H), even though the FKS2(-706 to +1)-lacZ reporter lacks the calcineurin-dependent response element. The dependence on Bck1p and Slt2p/Mpk1p was as expected for both FKS2(-706 to +1)-lacZ and PRM5-lacZ [1] whereas dependence on Cnb1p/Crz1p was unanticipated and revealed additional cross-regulation.

Ste11ΔNp activated FKS2 (-928 to +1) and FKS2(-706 to +1) through Cmk2p and Ptk2p

We tested whether the two potential Ste11p substrates Cmk2p and Ptk2p [20] were required for STE11ΔNp to activate FKS2-lacZ reporter genes. Loss of Cmk2p blocked the ability of STE11ΔNp to activate the FKS2 (-928 to +1) reporter gene to a greater extent than crz1Δ or cnb1Δ mutations (Figure 4E, H). A cnb1Δ cmk2Δ double mutant was as equivalently blocked as a cmk2Δ single mutant. Thus, Cmk2p is important for Ste11ΔNp activation of FKS2 (-928 to +1)-lacZ.

Similar results were found for the FKS2 (-706 to +1)-lacZ reporter gene, except in this instance the inhibitory effects of the cmk2Δ, cnb1Δ and crz1Δ single mutations were more similar to each other (Figure 4F, H). Cmk2p and Cnb1p provided additive functions based on a greater block in cmk2Δ cnb1Δ double mutant than either single mutant. Moreover, dependence on Cmk2p and Ptk2p was also apparent using the PRM5-lacZ reporter gene that senses the PKC pathway (Figure 4G, H). Cmk2p was more critical than Cnb1p and clearly greater in cells that express Ste11ΔNp activation was blocked Ste11ΔNp induction of the FKS2 (-706 to +1)-lacZ reporter that lacks the calcineurin CDRE (Figure 4F). Therefore, Cmk2p and Ptk2p are both required for Ste11ΔNp to signal to calcineurin and PKC pathway reporter genes, but Cmk2p plays a greater role than Ptk2p.

**STE11ΔN increased PRM5 mRNA and FKS2 mRNA**

We tested whether Ste11ΔNp increased FKS2 and PRM5 mRNAs in wild type and cnb1Δ strains. The mRNA levels for FKS2, PRM5, PMC1 and ACT1 were assessed by cDNA synthesis and real-time PCR using ACT1 as a normalization control. FKS2 mRNA increased in the presence of STE11ΔN, but a much larger increase was detected for PRM5, and no increase was detected for PMC1 mRNA (Figure 5A). The effect on FKS2 mRNA was less than what we detected with the FKS2-lacZ reporter and may reflect posttranscriptional effects or the fact that the promoter is complex with multiple signaling elements less easily distinguished than by using the reporter genes. A cnb1Δ mutation decreased the level of FKS2 and PRM5 mRNAs as expected but had no effect on basal mRNA of PMC1. Ste11ΔNp still increased FKS2 and PRM5 mRNAs in the cnb1Δ mutant, consistent with a calcineurin-independent pathway that could involve the PKC pathway.

**Ste11ΔNp increased the abundance of Cmk2p**

The observation that Ste11ΔNp activation was blocked in cmk2Δ strains (Figure 4E, H) supported the possibility that Cmk2p might be an in vivo substrate of Ste11p [20]. To look for a posttranslational effect of Ste11p on Cmk2p, we determined whether Ste11ΔNp influenced the mobility of Cmk2p in a native polyacrylamide gel mobility experiment. Triple and double GFP tags were inserted at the carboxyl-terminal end of CMK2 which were confirmed by PCR (data not shown) and western blotting (Figure 5B). Immunoblot analysis of whole cell extracts revealed that the abundance of Cmk2p was greater in cells that express Ste11ΔNp (Figure 5C). This finding was consistent with an effect of Ste11p on the CMK2 promoter that is controlled by Cmk2p and a post-transcriptional effect on the level of Cmk2p that could...
arise from feedback stimulation of its own expression [13]. A comparison of the mobility of CMK2-3XGFP and CMK2-2XGFP in wild type and Ste11ΔNp cells also suggested a faster mobility in the presence of Ste11ΔNp, which would be consistent with a posttranslational modification (Figure 5C). These data were consistent with the possibility that Ste11p regulated Cmk2p directly.

Ste11ΔNp activated CMK2-lacZ through Cmk2p

The calcium signal is transmitted to both Cnb1p and Cmk2p, but only Cnb1p appears to activate Crz1p [19]. Calcium activation of a CMK2-lacZ reporter is blocked by a crz1Δ cmk2Δ double mutation but not by a cmk2Δ single mutation, indicating that Crz1p activates the CMK2-lacZ gene independently of Cmk2p [19]. To test for an effect of Ste11ΔNp on the CMK2 promoter that...
might be mediated through its effects on Cmk2p protein, we measured CMK2-lacZ expression in wild type and cmk2Δ strains. The CMK2-lacZ reporter is stimulated by 200 mM CaCl2 in the medium in a Crz1- and FK506-dependent manner [19]. Basal and Ste11ΔNp induction of CMK2-lacZ were blocked by cnb1Δ and crz1Δ single mutations (Figure 5D). Strikingly, Cmk2p was also required for Ste11ΔNp to activate the CMK2-lacZ reporter gene, suggesting feedback control. The cmk2Δ mutation caused a large block in Ste11ΔNp induction of CMK2-lacZ expression compared with the wild type CMK2 strain. Ste7p and Ste12p were also required for Ste11ΔNp to efficiently activate the CMK2 promoter as was Ptk2p (Figure 5D). Therefore, Ste11ΔNp uses multi-pathway crosstalk to regulate the CMK2 promoter. The strict dependence on Cmk2p suggests Ste11p activates the level or activity of Cmk2p protein, which then activates the CMK2 promoter through positive feedback.

Ste11ΔNp activated morphogenesis in cmk2Δ strain
Ste11p and its known targets Ste7p and Pbs2p positively regulate morphogenesis. To determine whether Cmk2p had a role in morphogenesis, we counted the number of polarized and nonpolarized cells that were enlarged in wild type and cmk2Δ cells with and without Ste11ΔNp. Interestingly, in the cmk2Δ strain lacking Ste11ΔNp, only 2.1% cells displayed an enlarged morphology compared to 9.9% of wild type cells grown in YEPD medium (Table 1). These findings supported a role for Cmk2p in the control of morphogenesis. As expected, wild type cells expressing Ste11ΔNp grew bigger than control. In the cmk2Δ strain, fewer cells enlarged with Ste11ΔNp. However, Ste11ΔNp was still able to induce cell enlargement in the cmk2Δ strain. Collectively, these findings revealed that Cmk2p was important for morphogenesis and that Ste11ΔNp retained a strong ability to activate morphogenesis independent of Cmk2p, presumably through activation of one or more downstream targets (e.g. Pbs2p, Ste7p, Ptk2p, and Ste12p indirectly).

Ste11ΔNp also required Pbs2p, Ste7p and Cmk2p to activate a PRM5 reporter gene
To confirm the validity of our findings with FKS2, we examined the effects of mutations on PRM5, a second gene that senses both PKC pathway and mating pathway inputs (Figure 4C, D). Ste11ΔNp induction of expression of the PRM5-lacZ reporter gene was blocked by fus3Δ kss1Δ and ste12Δ mutations, but not by fus3Δ and kss1Δ single mutations (Figure 3F, G) as expected. In contrast, ste7Δ and pbs2Δ single mutations had no effect but Ste11ΔNp was blocked by the ste7Δpbs2Δ double mutation (Figure 3F, G). Thus, Pbs2p and Ste7p are both equivalently required for Ste11ΔNp to signal to PRM5-lacZ, a second PKC pathway reporter gene.

Among the various calcineurin pathway mutant strains examined, the cmk2Δ mutation caused the biggest block in Ste11ΔNp activation of PRM5-lacZ. A strong block occurred with a crz1Δ mutation, but only a partial decreased occurred for cnb1Δ and ptk2Δ mutations. The overall pattern of effects was similar to the FKS2 (-706 to +1) reporter gene, with Ptk2p being required for Cmk2p to regulate the PRM5-lacZ reporter (Figure 4G, H). The pattern of effects of mutations in CMK2, PTK2, CNB1 and CRZ1 resembled that found for FKS2 (-706 to +1)-lacZ reporter, which primarily senses the PKC pathway. Collectively, these findings supported Ste11p crosstalk of the PKC pathway as observed with the FKS2-lacZ gene and dependence on Pbs2p, Ste7p, Cmk2p and Ptk2p.

Slt2p/Mpk1p blocked Ste11ΔNp activation of the PMC1-lacZ reporter gene
Since Ste11p crosstalked with downstream targets of the calcineurin pathway, we also tested PMC1-lacZ, another reporter gene of this pathway whose expression is stimulated by Ca2+ [26]. When mutant strains were treated with Ca2+, PMC1-lacZ expression increased as reported (Figure 4G). However, Ste11ΔNp did not significantly increase PMC1-lacZ expression (Figure 5F). Instead, we found unexpectedly that Ste11ΔNp increased the level of PMC1-lacZ expression 5-fold in a strain lacking Mpk1p MAPK (Figure 5F; MPK1 and mpk1Δ strains grown in YEPD medium). This interesting result revealed that Sltp/Mpk1p normally suppresses Ste11ΔNp activation of PMC1-lacZ expression. We examined the level of PMC1 mRNA by RT-qPCR and found that expression of Ste11ΔNp decreased PMC1 mRNA levels in both wild type and cnb1Δ strains (Figure 5A), which would be consistent with the ability of Ste11ΔNp to increase the level of active Slt2p/Mpk1p. Thus, Slt2p/Mpk1p may prevent Ste11p cross-regulation of the calcineurin pathway and feedback inhibit at some step upon activation by Ste11p.

Discussion
The activation of the calcineurin, cell wall integrity and high osmolarity pathways in S. cerevisiae and other fungi is important for cell survival and response to cell
wall damage and other stimuli including calcium influx, polarized growth, hypo-osmolarity, pheromone, heat shock and perturbation of the actin cytoskeleton [1,35]. FKS2 is important for survival under stress conditions linked to the calcinerin and PKC pathways [1,14,15]. Prior analysis had suggested a potential link between mating pathway activation and the calcineurin and cell wall integrity pathways, but the means by which this signaling occurs was not fully understood [1,36]. We discovered that Ste11ΔNp crosstalks with cell wall integrity and calcineurin pathways through Pbs2p MAPKK, and two putative kinase targets Cmk2p and Ptk2p, with separable contribution from Ste7p MAPKK, Fus3p and Kss1p MAPKs. Support for the relevance of our findings is the logical pattern of effects of mutations in the genes encoding these protein kinases, and in calcineurin and cell wall integrity components (Additional File 1, Figure S1; Figure 6). Moreover, similar patterns of control were found for other physiologically relevant promoters, including PRM5 and CMK2. Additional support comes from the observation that the activation of FKS2 reporter genes by Ste11p did not require upstream activators or Ste20p or Bni1p-mediated morphogenesis, and the finding that activation of Mpk1p kinase (Figure 1) was

Figure 6 Summary of Ste11p crosstalk through kinase targets to FKS2. (A) Ste11p crosstalks with mating, calcineurin, HOG, and cell wall integrity pathways. (B) Slt2p/Mpk1p may inhibit Ste11p activation of PMC1.
more dependent on Ste11p than it was on mating pathway components such as Ste5p and Ste2p.

The data hint at the existence of a distinct promoter element within the FKS2 (-706 to +1) promoter region that is regulated by Ste11p, but not by Ste12p. STE11-4 did not activate a FKS2(-540 to -375)-lacZ reporter that strictly overlaps the Mpk1p regulated SBF element (Figure 2A), but could stimulate FKS2(-706 to +1)-lacZ and Mpk1p kinase activity. The Ste11p-inducible promoter element operates independently of Ste12p, because Ste12p could not activate FKS2(-706 to +1)-lacZ although it could activate FKS2(-928 to +1)-lacZ with the CDRE. Collectively, these observations suggest the Ste11p-inducible element overlaps the -706 to -540 region.

Multiple lines of evidence support a functional link between Ste11p and Cmk2p and the calcineurin pathway (Figure 6). While support for direct physical interaction between Cmk2p and Ste11p comes from the prior in vitro analysis [17], there has been no functional connection between these protein kinases. Four lines of evidence support Cmk2p being positively regulated by Ste11p. First, the effect of Ste11ΔNp on FKS2 expression was almost blocked without Cmk2p; Ste11ΔNp was functionally dependent on Cmk2p. Second, the cmk2Δ mutation caused a larger effect block of Ste11ΔNp phosphorylation. Third, there is a hint of a shift in the mobility of Cmk2-GFP when Ste11p is expressed (Figure 5C), supporting the possibility of posttranslational modification by phosphorylation. Fourth, Ste11ΔNp clearly increased the abundance of Cmk2-GFP (Figure 5C) and increased the expression of CMK2-lacZ in a manner dependent on Cmk2p (Figure 5D), consistent with the possibility of greater Cmk2p activity. These observations support direct positive control of Cmk2p by Ste11p phosphorylation, although more work is needed.

In vitro analysis suggests Cmk2p may be a target of phosphorylation by a number of kinases including Ste11p, Fus3p and Pbs2p and a central node in cellular control [20]. Cmk2p was the substrate of Ste11p, Pbs2p, Fus3p, Cka1p, Kns1p, Pho85p, Pho85-Pcll9, Slt2p/Mpk1p, Pkh2p, Tpk2p, Tpk3p, Ygl059w and Yol128c. Thus, the affects of fus3Δ and pbs2Δ mutations on expression of the various reporter genes in our study may be partly due to loss of regulation of Cmk2p by Fus3p and Pbs2p. The expression levels of FKS2 (-928 to +1)-lacZ and FKS2 (-706 to +1)-lacZ reporter genes were equivalently reduced to the same degree by the pbs2Δ or fus3Δ mutations. It is curious that a ste7Δ mutation had little effect since Ste7p activates Fus3p. However, Ste7p contribution could be detected in the ste7Δ pbs2Δ double mutant strain with Ste11ΔNp; FKS2 expression was 3-fold lower than in pbs2Δ or fus3Δ single mutant strains. Perhaps without its regulatory domain, Ste11p preferentially phosphorylates Pbs2p over Ste7p; this could provide an explanation for the lesser effect in a ste7Δ strain. An alternative explanation is that loss of Ste7p leads to an increase in Ste11ΔNp pathway flux through Pbs2p. In fission yeast, Cmk2p is phosphorylated in response to oxidative stress [37], supporting the possibility that Ste11p regulates Cmk2p in part through Pbs2p. Further work is needed to understand how Cmk2p and Ptk2p regulate the A FKS2 gene. Ptacek et al., 2005 identified 9 in vitro substrates for Cmk2p and 194 for Ptk2p [20]. Remarkably, four of these substrates overlap, supporting functional redundancy between Cmk2p and Ptk2p.

Our analysis reveals a possible feedback loop whereby Cmk2p activates its own expression through calcineurin (Cnb1). CMK2-lacZ expression was also partially dependent on Ste7p and Ste12p. Because Ste12p is a transcription factor, there are multiple genes through which it could regulate CMK2-lacZ. However, one gene could be FUS3, since Fus3p is both activated by Ste7p and may directly regulate Cmk2p [20]. The multi-pathway crosstalk through the CMK2 promoter that requires Cmk2p could reflect a cooperative activity of one or more transcription factors. Potential shared physical linkages between Cmk2p, calmodulin and calcineurin (Cmk2-Cmd1, [38-40]; Cmd1-Cna1p and Cmd1-Cnb1p, [38,39,41]), raises the possibility of a direct link between Cmk2p and calcineurin signaling to Crz1p.

An unexpected interesting observation was that Slt2p/Mpk1p inhibits the ability of Ste11p to regulate a calcineurin pathway gene, PMC1. In the W303a background, PMC1 expression could be activated in Ca2+ medium (Figure 5E), but there was only modest activation by Fus3p, since Fus3p is both activated by Ste7p and may directly regulate Cmk2p [20]. The multi-pathway crosstalk through the CMK2 promoter that requires Cmk2p could reflect a cooperative activity of one or more transcription factors. Potential shared physical linkages between Cmk2p, calmodulin and calcineurin (Cmk2-Cmd1, [38-40]; Cmd1-Cna1p and Cmd1-Cnb1p, [38,39,41]), raises the possibility of a direct link between Cmk2p and calcineurin signaling to Crz1p.

In vitro analysis suggests Cmk2p may be a target of phosphorylation by a number of kinases including Ste11p, Fus3p and Pbs2p and a central node in cellular control [20]. Cmk2p was the substrate of Ste11p, Pbs2p, Fus3p, Cka1p, Kns1p, Pho85p, Pho85-Pcll9, Slt2p/Mpk1p, Pkh2p, Tpk2p, Tpk3p, Ygl059w and Yol128c. Thus, the affects of fus3Δ and pbs2Δ mutations on expression of the various reporter genes in our study may be partly due to loss of regulation of Cmk2p by Fus3p and Pbs2p. The expression levels of FKS2 (-928 to +1)-lacZ and FKS2 (-706 to +1)-lacZ reporter genes were equivalently reduced to the same degree by the pbs2Δ or fus3Δ mutations. It is curious that a ste7Δ mutation had little effect since Ste7p activates Fus3p. However, Ste7p contribution could be detected in the ste7Δ pbs2Δ double mutant strain with Ste11ΔNp; FKS2 expression was 3-fold lower than in pbs2Δ or fus3Δ single mutant strains. Perhaps without its regulatory domain, Ste11p preferentially phosphorylates Pbs2p over Ste7p; this could provide an explanation for the lesser effect in a ste7Δ strain. An alternative explanation is that loss of Ste7p leads to an increase in Ste11ΔNp pathway flux through Pbs2p. In fission yeast, Cmk2p is phosphorylated in response to oxidative stress [37], supporting the possibility that Ste11p regulates Cmk2p in part through Pbs2p. Further work is needed to understand how Cmk2p and Ptk2p regulate the A FKS2 gene. Ptacek et al., 2005 identified 9 in vitro substrates for Cmk2p and 194 for Ptk2p [20]. Remarkably, four of these substrates overlap, supporting functional redundancy between Cmk2p and Ptk2p.

Our analysis reveals a possible feedback loop whereby Cmk2p activates its own expression through calcineurin (Cnb1). CMK2-lacZ expression was also partially dependent on Ste7p and Ste12p. Because Ste12p is a transcription factor, there are multiple genes through which it could regulate CMK2-lacZ. However, one gene could be FUS3, since Fus3p is both activated by Ste7p and may directly regulate Cmk2p [20]. The multi-pathway crosstalk through the CMK2 promoter that requires Cmk2p could reflect a cooperative activity of one or more transcription factors. Potential shared physical linkages between Cmk2p, calmodulin and calcineurin (Cmk2-Cmd1, [38-40]; Cmd1-Cna1p and Cmd1-Cnb1p, [38,39,41]), raises the possibility of a direct link between Cmk2p and calcineurin signaling to Crz1p.

An unexpected interesting observation was that Slt2p/Mpk1p inhibits the ability of Ste11p to regulate a calcineurin pathway gene, PMC1. In the W303a background, PMC1 expression could be activated in Ca2+ medium (Figure 5E), but there was only modest activation by Fus3p, since Fus3p is both activated by Ste7p and may directly regulate Cmk2p [20]. The multi-pathway crosstalk through the CMK2 promoter that requires Cmk2p could reflect a cooperative activity of one or more transcription factors. Potential shared physical linkages between Cmk2p, calmodulin and calcineurin (Cmk2-Cmd1, [38-40]; Cmd1-Cna1p and Cmd1-Cnb1p, [38,39,41]), raises the possibility of a direct link between Cmk2p and calcineurin signaling to Crz1p.

In vitro analysis suggests Cmk2p may be a target of phosphorylation by a number of kinases including Ste11p, Fus3p and Pbs2p and a central node in cellular control [20]. Cmk2p was the substrate of Ste11p, Pbs2p, Fus3p, Cka1p, Kns1p, Pho85p, Pho85-Pcll9, Slt2p/Mpk1p, Pkh2p, Tpk2p, Tpk3p, Ygl059w and Yol128c. Thus, the affects of fus3Δ and pbs2Δ mutations on expression of the various reporter genes in our study may be partly due to loss of regulation of Cmk2p by Fus3p and Pbs2p. The expression levels of FKS2 (-928 to +1)-lacZ and FKS2 (-706 to +1)-lacZ reporter genes were equivalently reduced to the same degree by the pbs2Δ or fus3Δ mutations. It is curious that a ste7Δ mutation had little effect since Ste7p activates Fus3p. However, Ste7p contribution could be detected in the ste7Δ pbs2Δ double mutant strain with Ste11ΔNp; FKS2 expression was 3-fold lower than in pbs2Δ or fus3Δ single mutant strains. Perhaps without its regulatory domain, Ste11p preferentially phosphorylates Pbs2p over Ste7p; this could provide an explanation for the lesser effect in a ste7Δ strain. An alternative explanation is that loss of Ste7p leads to an increase in Ste11ΔNp pathway flux through Pbs2p. In fission yeast, Cmk2p is phosphorylated in response to oxidative stress [37], supporting the possibility that Ste11p regulates Cmk2p in part through Pbs2p. Further work is needed to understand how Cmk2p and Ptk2p regulate the A FKS2 gene. Ptacek et al., 2005 identified 9 in vitro substrates for Cmk2p and 194 for Ptk2p [20]. Remarkably, four of these substrates overlap, supporting functional redundancy between Cmk2p and Ptk2p.

Our analysis reveals a possible feedback loop whereby Cmk2p activates its own expression through calcineurin (Cnb1). CMK2-lacZ expression was also partially dependent on Ste7p and Ste12p. Because Ste12p is a transcriptio...
Ste11ΔNp on activating various reporter genes. These findings suggest significant compensation and redundancy at the level of control and support functional relationships among the many targets of Ste11p.

The study also reveals that a transcription factor (i.e. Ste12p) can strongly regulate the expression of a gene through indirect means. Prior CHIP studies failed to find evidence of Ste12p binding to FKS2 and α factor induction of FKS2 mRNA is completely blocked by FK506 and erg11Δ [16,27]. It has been suggested that mating pheromone stimulates the FKS2 gene indirectly through effects on calcium influx [14,27]. However, multiple pathways are activated by α factor mating pheromone and Ste12p has the capacity to bind to multiple transcription factors to regulate distinct pathways. Ste12p and Ste11ΔNp stimulated FKS2(-928 to +1)-lacZ reporter harboring the CRE more than one of three Ste12p consensus TGGAAACA sites (-1374GTGAAACA-1367, -928CTGAAACA-919, -901TTGAAACA-894), but stimulation was largely blocked by a calcineurin regulatory subunit cnb1Δ mutation. Thus, Ste12p is critically important for activation of FKS2, but its function is indirect within the promoter region examined in this study.

Conclusions

Our results lead us to several conclusions: (1) Ste11p crosstalks with calcineurin and cell wall integrity pathways independently of the mating pathway. (2) Ste11p crosstalks through all known or potential downstream kinase targets, Ste7p, Pbs2p, Cmk2p and Ptk2p, to regulate calcineurin and PKC pathway reporter genes, FKS2, PRM5, and CMK2. (3) Cmk2p is likely to be a direct substrate of Ste11p in the calcineurin pathway in vivo based on functional dependence, (4) Ste11p increases the abundance of Cmk2p in part through a positive effect on the CMK2 gene promoter which is dependent on Cmk2p. An additional unexpected observation is that Slt2p/Mpk1p may prevent Ste11p from stimulating PMCI expression, which could arise through Ste11p activation of Slt2p/Mpk1p. A summary of these observations is shown in Figure 6. Ste12p is important but its role is indirect.

Methods

Media, Plasmids, and Strains

Yeast extract/peptone/dextrose (YPD) and synthetic complete (SC) media with dextrose, raffinose or galactose were made according to standard laboratory methods. Yeast strains and plasmids used in this study are described in Table 2. Yeast strains are isogenic to W303a or Sigma or S288c strains. All yeast deletion strains derivatives were made either with published deletion plasmids or through PCR oligonucleotide cassette disruption with pFA6a-3xGFP-KAN as described [44]. EY957 (W303a MATa bar1Δ background) was engineered to harbor C-terminally tagged CMK2 using pFA6a-3xGFP-KAN following the experimental approach in [44]. Two CMK2-GFP-KAN integrants were recovered, one with 3xGFP-KAN and the other with 2xGFP-KAN.

Detection of active MAPK in whole cell extracts

Active Slt2p/Mpk1p, Fus3p and Kss1p were detected with anti-phospho-p42p44 antibodies using 200 μg whole cell extract exactly as described [25].

Galactose induction

Cells were grown at 30°C (unless noted otherwise) to logarithmic phase in medium containing 2% dextrose. Cells were then pelleted and resuspended in medium containing 2% raffinose and 0.1% dextrose and grown to an OD600 ~ 0.8, then collected and resuspended in medium containing 2% galactose at an OD600 ~ 0.2, and then grown for another 6 hours.

β-galactosidase assay

Cells were harvested at 4°C and frozen at -80°C. Pellets were thawed on ice in 0.25 ml breaking buffer (0.1 M Tris-HCl pH 8.0, 20% Glycerol v/v, 1 mM DTT) containing 12.5 μl of 40 mM PMSF (phenyl methyl sulphonyl fluoride in 95% ethanol). Glass beads were added and samples were vortexed three times at 4°C, added 0.25 ml breaking buffer into tubes and vortexed again. The supernatant was transferred to a new tube, centrifuged at 12,000 g for 15 min at 4°C and clarified supernatant was transferred to a new tube; protein concentration was determined by BIORAD™ assay. A linear range of protein concentration and incubation time was established. Standardized amounts of extracts were mixed with Z buffer (0.06 M Na2HPO4 · 7H2O, 0.04 M NaH2PO4 · H2O, 0.01 M KCl, 0.001 M MgSO4 · 7H2O) and incubated for 15 min at 28°C. Reactions were initiated by addition of 0.2 ml ONPG (4 mg/ml) and stopped by addition of 0.5 ml Na2CO3 (1 M) after the color had changed into light yellow. Read samples at OD420, then calculated as nanomoles of ONPG cleaved per min per milligram protein as described [45]. These experiments were performed at least in triplicate from independent yeast transformants.

RNA preparation for yeast

Cells were grown in the appropriate medium, then 10 ml (OD600 ~ 0.6-0.8) of cells were collected, washed once with sterile deionized water, and then resuspended in 600 μl TES buffer (10 mM Tris pH 7.5, 10 mM EDTA, 0.5% SDS). Samples were mixed with 500 μl acid washed phenol and incubated 45 min to 1 hour at 65°C.
### Table 2 Yeast strains and plasmids used in this study.

| Strain | Relevant genotype | Source or reference |
|--------|-------------------|---------------------|
| EYL582 | MATα mpk1Δ::TRP1 ura3-1 trp1-1 leu2-3,112 ade2-1 ade3-1 his3 can1-100 lys2 (Gal+) | Elion lab collection |
| EYL780 | MATα bar1Δ far1Δ ste11-4 ste12Δ::URA3 trp1-1 his3Δ200 leu2-3,112 ade2-1 ura3-1 can1-100 (Gal+) FUS1p::HIS3::lys2 | Elion lab collection |
| EY957  | MATα bar1Δ trp1-1 his3-11,15 leu2-3,112 ade2-1 ura3-1 can1-100 (Gal+) | Elion lab collection |
| EY1298 | MATα bar1Δ far1Δ ste11-4 trp1-1 his3Δ200 leu2-3,112 ade2-1 ura3-1 can1-100 (Gal+) FUS1p::HIS3::lys2 | Elion lab collection |
| EYL4661| MATα bar1Δ trp1-1 his3-11,15 leu2-3,112 ade2-1 ura3-1 can1-100 (Gal+) ste11Δ::Hygro | Elion lab collection |
| EYL328 | MATα bar1Δ trp1-1 his3-11,15 leu2-3,112 ade2-1 ura3-1 can1-100 (Gal+) pbs2Δ::Hygro | This study |
| EYL529 | MATα bar1Δ trp1-1 his3-11,15 leu2-3,112 ade2-1 ura3-1 can1-100 (Gal+) kss1Δ::His3 fus3Δ::Hygro pbs2Δ::kan | This study |
| EYL530 | MATα bar1Δ trp1-1 his3-11,15 leu2-3,112 ade2-1 ura3-1 can1-100 (Gal+) ste12Δ::Hygro | This study |
| EYL531 | MATα bar1Δ trp1-1 his3-11,15 leu2-3,112 ade2-1 ura3-1 can1-100 (Gal+) bni1Δ::Hygro | This study |
| EYL532 | MATα bar1Δ trp1-1 his3-11,15 leu2-3,112 ade2-1 ura3-1 can1-100 (Gal+) ste7Δ::Hygro pbs2Δ::kan | This study |
| EYL533 | MATα bar1Δ trp1-1 his3-11,15 leu2-3,112 ade2-1 ura3-1 can1-100 (Gal+) cmk2Δ::Hygro | This study |
| EYL534 | MATα bar1Δ trp1-1 his3-11,15 leu2-3,112 ade2-1 ura3-1 can1-100 (Gal+) ptk2Δ::Hygro | This study |
| EYL535 | MATα bar1Δ trp1-1 his3-11,15 leu2-3,112 ade2-1 ura3-1 can1-100 (Gal+) crb1Δ::Hygro | This study |
| EYL536 | MATα bar1Δ far1Δ trp1-1 his3Δ200 leu2-3,112 ade2-1 ura3-1 can1-100 (Gal+) FUS1p::HIS3::lys2 pbs2Δ::Hygro | This study |
| EYL537 | MATα bar1Δ far1Δ trp1-1 his3Δ200 leu2-3,112 ade2-1 ura3-1 can1-100 (Gal+) FUS1p::HIS3::lys2 cmk1Δ::Hygro | This study |
| EYL538 | MATα bar1Δ trp1-1 his3-11,15 leu2-3,112 ade2-1 ura3-1 can1-100 (Gal+) kss1Δ::His3 fus3Δ::Hygro | This study |
| EYL539 | MATα bar1Δ trp1-1 his3-11,15 leu2-3,112 ade2-1 ura3-1 can1-100 (Gal+) ptk2Δ::Hygro cmk2Δ::kan | This study |
| EYL530 | MATα bar1Δ trp1-1 his3-11,15 leu2-3,112 ade2-1 ura3-1 can1-100 (Gal+) cmk2::tag3::GFP::kan | This study |
| EYL532 | MATα bar1Δ trp1-1 his3-11,15 leu2-3,112 ade2-1 ura3-1 can1-100 (Gal+) crz1Δ::kan | This study |

**S288c deletion strains in Figure 1 are isogenic to S288c BY4741.**

BY4741 | MATα his3Δ ura3Δ leu2Δ met15-0 | [46]
--- | --- | ---
RG2468 | ste4Δ::KAN leu2 ura3 his3 met15 | Research Genetics
--- | --- | ---
RG4038 | ste5Δ::KAN leu2 ura3 his3 met15 | Research Genetics
--- | --- | ---
RG3439 | ste50Δ::KAN leu2 ura3 his3 met15 | Research Genetics
--- | --- | ---
RG5271 | ste11Δ::KAN leu2 ura3 his3 met15 | Research Genetics
--- | --- | ---
RG6981 | kss1Δ::KAN leu2 ura3 his3 met15 | Research Genetics
--- | --- | ---
JAY408 | fus3Δ::KAN ste52Δ::KAN | [25]

**Isogenic to S1278b**

L5528 | ura3-52, his3::hisG | [48]
--- | --- | ---
L5554 | ste5::LEU2, ura3-52, leu2::hisG | [48]

**Plasmids**

pYGU-11A | pGAL1::STE11::LEU2 CEN | [49]
--- | --- | ---
GAL1 vector | pGAL1, LEU2 CEN | Elion lab collection
--- | --- | ---
pYEE133 | pGAL1::STE12 TRP1 CEN | Elion lab collection
--- | --- | ---
GAL1 vector | pGAL1, TRP1 CEN | Elion lab collection
--- | --- | ---
pO01 | pCMK2-loaZ, URA3, 2 μ | [19]
--- | --- | ---
EBL05 | pDL1460 pFUS1::ubi1-loaZ, URA3, 2 μ | Elion lab collection
--- | --- | ---
EBL183 | PMCl-loaZ, URA3, 2 μ | [50]
--- | --- | ---
EBL185 | pDM5, FS2 (-968-+6)-CYC1-loaZ, URA3, 2 μ | [26]
--- | --- | ---
EBL201 | FS2 (-928-+1)-CYC1-loaZ, URA3, 2 μ | [14]
--- | --- | ---
EBL200 | FS2 (-706-+6)-CYC1-loaZ, URA3, 2 μ | [14]
--- | --- | ---
EBL1258 | FS2 (-540-375)-CYC1-loaZ, URA3, 2 μ | [15]
--- | --- | ---
EBL513 | YKL161c (MLP1)-loaZ, URA3, 2 μ | Elion lab collection
--- | --- | ---
EBL514 | YIL117c (PRM5)-loaZ, URA3, 2 μ | [51]
--- | --- | ---
EBL517 | pLGB-178 pCYC1-loaZ, URA3, 2 μ | [52]
with vortexing for 30 sec every 10 min. Samples were then cooled on ice and centrifuged 5 min. and then 0.5 ml aqueous phase was transferred into new tubes and re-extracted with 400 μl acid phenol. These steps were repeated once again. Then 400 μl aqueous phase was transferred into a new tube and mixed with 40 μl sodium acetate (3M, pH 5.3) and 800 μl ethanol and incubated for 1 hour at -20°C. RNA and other precipitated material was washed and the RNA was purified with a Qiagen kit following manufacturer’s specifications. 1-10 μg of RNA was used for cDNA synthesis according to protocol (SuperScript III, Invitrogen) and the cDNA was used for real-time quantitative PCR (RT-qPCR).

The following forward and reverse primers were used, respectively: ACT1, 5’-TGATCTCGGTAGTTGTTT-3’ and 5’-AAATGGCGTGAAGAGAAAC-3’; FKS2, 5’- GACATCTGTAAAGATAGAG-3’ and 5’- CATGACAAAACCATACAG-3’; PMC1, 5’- ACTGTGGTGATAGTGTC-3’ and 5’- TCGAGTCCAAATACG-3’; PRMS, 5’- TGGGTGTCTACATCTCTTC-3’ and 5’- TGGTTACGATTACGCTAC-3’.

**Immunoblot analysis**

Cmk2p was monitored by western blot analysis of 3xGFP-tagged and 2xGFP-tagged Cmk2p. Cells were lysed by vortexing with glass beads in lysis buffer (50 mM Tris-HCl, pH 7.5, 100 mM NaCl, 0.5% triton-X 100, 10 mM NaF, 5 μg/ml aprotinin, 5 μg/ml leupeptin, 0.2 mM Na3VO4). Cell extracts containing 100 μg of total protein were run on 6% native polyacrylamide gels in 307 mM Tris base, 76 mM Bis, pH 8.8 at 100 V constant voltage. After blotting onto PVDF membranes, anti-Cmk2p antibody was used for real-time quantitative PCR (RT-qPCR).

**Statistics**

Data were expressed as mean ± SE of at least three independent experiments. Statistical significance was computed by the unpaired Student’s t test. A p < 0.05 was considered statistically significant.

**Additional material**

Additional file 1: Figure S1. Ste11 in vitro phosphorylation links to Cmk2p and Ptk2p overlap other MAPK pathway kinases. From Ptacek J, Devgan G, Michaud G, Zhu H, Zhu X, Fasolo J, Guo H, Jona G, Breitkreutz A, Sopko R, et al. Global analysis of protein phosphorylation in yeast. Nature 2005, 438:679-684.

Additional file 2: Figure S2. Level of active Mpk1p and Kss1p in different yeast backgrounds. This figure shows short and long exposures of immunoblot in Figure 1A to better visualize the relative amount of phosphorylated Mpk1p in W303a compared to S288c. The data in Figure 1A are from a re-probing of the same normalized immunoblot shown in Supplemental Figure Two in Anderson et al., 2004 [25].

**Acknowledgements**

We thank Steve Buratowski (Harvard Medical School) and Zhijie Chang (Tsinghua University) for comments on the manuscript, and Kyle Cunningham (Johns Hopkins University) and David Levin (Boston University) for plasmids. This research was supported by a fellowship to X.Y.W. from the China Scholarship Council Postgraduate Scholarship Program (2008-2009) and prior funding to E.A.E. from the N.I.H. (R01 GM46962).

**Author details**

1. Department of Biological Chemistry and Molecular Pharmacology, Harvard Medical School, Boston, MA 02115, USA.
2. Protein Science Key Laboratory of the Ministry of Education, School of Medicine, Tsinghua University, Beijing 100084, PR China.
3. Martek Biosciences, Nautilus Court North, Boulder, CO 80301, USA.

**Authors’ contributions**

X.Y.W. helped design and carried out most of the experiments and wrote the manuscript. M.A.S. helped to do the experiments and edited the manuscript. D.M.S. performed experiments and edited the manuscript. E.A.E. conceived the project, supervised experiments and revised the manuscript. All authors read and approved relevant portions of the manuscript.

Received: 21 September 2011. Accepted: 24 November 2011. Published: 24 November 2011.

**References**

1. Levin DE. Cell wall integrity signaling in Saccharomyces cerevisiae. *Microbiol Mol Biol Rev* 2005, 69:262-291.
2. Lee BN, Elion EA. The MAPKKK Ste11 regulates vegetative growth through a kinase cascade of shared signaling components. *Proc Natl Acad Sci USA* 1999, 96:12679-12684.
3. Westfall PJ, Ballon DR, Thorer J. When the stress of your environment makes you go HOG wild. *Science* 2004, 306:1511-1512.
4. Lange-Carter CA, Pleiman CM, Gardner AM, Blumer KJ, Johnson GL: A divergence in the MAP kinase regulatory network defined by MEK kinase and Raf. *Science* 1993, 260:315-319.
5. Chen RE, Thorer J. Function and regulation in MAPK signaling pathways: lessons learned from the yeast Saccharomyces cerevisiae. *Biochim Biophys Acta* 2007, 1773:1311-1340.
6. Qi M, Elion EA: MAP kinase pathways. *J Cell Sci* 2005, 118:3569-3572.
7. Elion EA, Qi M, Chen W: Signal transduction. Signaling specificity in yeast. *Science* 2005, 307:687-688.
8. Rep M, Kranz M, Thevelein JM, Hofmann S: The transcriptional response of Saccharomyces cerevisiae to osmotic shock, Hot1p and Msn2p/Msn4p are required for the induction of subsets of high osmolarity glycerol pathway-dependent genes. *J Biol Chem* 2000, 275:8290-8300.
9. Possas F, Chambers JR, Heyman JA, Hoeffer JP, de Nadal E, Arino J: The transcriptional response of yeast to saline stress. *J Biol Chem* 2000, 275:17249-17255.
10. Sucher AJ, Chahine EB, Baker HE: Echinocandins: the newest class of antifungals. *Ann Pharmacother* 2009, 43:1647-1657.
11. Garcia-Effron G, Lee S, Park S, Cleary JD, Perlin DS: Effect of Candida glabrata FKS1 and FKS2 mutations on echinocandin sensitivity and kinetics of 1,3-beta-D-glucan synthase: implication for the existing susceptibility breakpoint. *Antimicrob Agents Chemother* 2009, 53:3690-3699.
12. Iida H, Nakamura H, Ono T, Okumura MS, Anraku Y: MID1, a novel Saccharomyces cerevisiae gene encoding a plasma membrane protein, is required for Ca2+ influx and mating. *Mol Cell Biol* 1994, 14:8259-8271.
13. Moser MJ, Geiser JR, Davis TN: Ca2+-calmodulin promotes survival of Ca2+-calmodulin-dependent protein kinase. *Mol Cell Biol* 1996, 16:4804-4813.
14. Zhao C, Jang US, Garrett-Engele P, Roe T, Cyert MS, Levin DE: Temperature-induced expression of yeast FKS2 is under the dual control of protein kinase C and calcineurin. *Mol Cell Biol* 1998, 18:1013-1022.
15. Kim KY, Truman AW, Levin DE: Yeast Mpk1 mitogen-activated protein kinase activates transcription through Swi4/Swi6 by a noncatalytic mechanism that requires upstream signal. *Mol Cell Biol* 2000, 20:2579-2589.
Deletions of SKY1 or PTK2 in the Saccharomyces cerevisiae wall integrity pathway in stress response and cross talk with other stress mitogen-activated protein kinase pathways in Saccharomyces cerevisiae. Concepcion ML, Rivas C, Arias P, Nombela C, Posas F, Arroyo J. Functional coscaffolds linking components in the yeast high osmolarity MAP kinase pathway and its regulation in Saccharomyces cerevisiae. Genes Dev 2001, 15:3445-3458.

Stathopoulos AM, Cyert MS. Calcium ions act through the CRZ1/TCN1-encoded transcription factor to regulate gene expression in yeast. Genes Dev 1997, 11:3445-3458.

Cherkasova V, Lyons DM, Elion EA. Fus3p and Kss1p control G1 arrest in Saccharomyces cerevisiae through a balance of distinct arrest and proliferative functions that operate in parallel with Far1p. Genetics 1999, 151:899-1004.

Slaughter BD, Huff JM, Wiegraeb L, Schwartz JW, Li R. SAM domain-based protein oligomerization observed by live-cell fluorescence fluctuation spectroscopy. PLoS One 2008, 3:e1931.

Bhunia A, Domadina PN, Mohamad H, Bhattacharya S. NMR structural studies of the Ste11 SAM domain in the dodecyl phosphocholine micelle. Proteins 2009, 74:328-343.

Zamparam A, Bhattacharya RP, Nittler MP, Lim WA. Sho1 and Pbs2 act as csc scaffolding links in the components of the yeast high osmolarity MAP kinase pathway. Mol Cell Biol 2004, 14:825-832.

Lefrancois P, Euskirchen GM, Auerbach RK, Rozowsky J, Gibson T, Breuer EM, Errede B. Coordination of the mating and cell integrity mitogen-activated protein kinase pathways in Saccharomyces cerevisiae. Mol Cell Biol 1997, 17:6571-6575.

Bermjejov C, Rodriguez E, Garcia R, Rodriguez-Pena JM, Rodriguez de la Concepcion ML, Rivas C, Arias P, Nomberla C, Posas F, Arroyo J. The sequential activation of the yeast HOG and SLT2 pathways is required for cell survival to cell wall stress. Mol Biol Cell 2008, 19:1113-1123.

Fuchs BB, Mylonakis E. Our paths might cross: the role of the fungal cell wall integrity pathway in stress response and cross talk with other stress response pathways. Eukaryot Cell 2009, 8:1616-1625.

Sanchez-Piris M, Posas F, Alemany V, Winge I, Hidalgo E, Bachs O, Aligre R. The serine/threonine kinase Cmk2 is required for oxidative stress response in fission yeast. J Biol Chem 2002, 277:17722-17727.

Ho T, Guthrie A, Hellobt A, Bader GD, Moore L, Adams SL, Millar A, Taylor P, Bennett K, Boutilier K, et al. Systematic identification of protein complexes in Saccharomyces cerevisiae by mass spectrometry. Nature 2002, 141:180-183.

Krogan NJ, Cagney G, Yu H, Zhong G, Guo X, Ingachowski A, Li J, Pu S, Datta N, Tikuisis AP, et al. Global landscape of protein complexes in the yeast Saccharomyces cerevisiae. Nature 2006, 440:637-643.

Zhu H, Brigin M, Bangham R, Hall D, Casamayor A, Bertone P, Lan N, Jansen R, Bidlingmaier S, Houfek T, et al. Global analysis of protein activities using proteome chips. Science 2001, 293:210-215.

Jiang B, Cyert MS. Identification of a novel region critical for calcineurin function in vivo and in vitro. J Biol Chem 1999, 274:18543-18551.

Zheng W, Zhao H, Mancena E, Sternmutter LM, Snyder M. Genetic analysis of variation in transcription factor binding in yeast. Nature 2010, 464:1187-1191.

Ren R, Robert F, Wyrick JJ, Aparicio O, Jennings EG, Simon I, Zeitlinger J, Schreiber J, Hannett N, Kanin E, et al. Genome-wide location and function of DNA binding proteins. Science 2000, 290:2306-2309.

Yu L, Qi M, Sheff MA, Elion EA. Counteractive control of polarized morphogenesis during mating by mitogen-activated protein kinase Fus3 and G1 cyclin-dependent kinase. Mol Biol Cell 2008, 19:1739-1752.

Farley FW, Satterberg B, Goldsmith J, Elion EA. Relative dependence of different outputs of the Saccharomyces cerevisiae pheromone response pathway on the MAP kinase Fus3p. Genetics 1999, 151:1425-1444.

Winzelek EA, Shomeroff DD, Astromoff A, Liang H, Anderson K, Andrie B, Bangham R, Benito R, Boeke JD, Bussey H, et al. Functional characterization of the S. cerevisiae genome by gene deletion and parallel analysis. Science 1999, 285:901-906.

Brachmann CB, Davies A, Cost GJ, Caputo E, Li J, Hieter P, Boeke JD. Designer deletion strains derived from Saccharomyces cerevisiae S288C: a useful set of strains and plasmids for PCR-mediated gene disruption and other applications. Yeast 1998, 14:115-132.

Roberts RL, Fink GR. Elements of a single MAP kinase cascade in Saccharomyces cerevisiae mediate two developmental programs in the same cell type: mating and invasive growth. Genes Dev 1998, 12:1187-1191.