Calcineurin regulates the yeast synaptojanin Inp53/Sjl3 during membrane stress

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Abstract

During hyperosmotic shock, Saccharomyces cerevisiae adjusts to physiological challenges, including large plasma membrane invaginations generated by rapid cell shrinkage. Calcineurin, the Ca²⁺/calmodulin-dependent phosphatase, is normally cytosolic but concentrates in puncta and at sites of polarized growth during intense osmotic stress; inhibition of calcineurin-activated gene expression suggests that restricting its access to substrates tunes calcineurin signaling specificity. Hyperosmotic shock promotes calcineurin binding to and dephosphorylation of the PI(4,5)P₂ phosphatase synaptojanin/Inp53/Sjl3 and causes dramatic calcineurin-dependent reorganization of PI(4,5)P₂-enriched membrane domains. Inp53 normally promotes sorting at the trans-Golgi network but localizes to cortical actin patches in osmotically stressed cells. By activating Inp53, calcineurin repolarizes the actin cytoskeleton and maintains normal plasma membrane morphology in synaptojanin-limited cells. In response to hyperosmotic shock and calcineurin-dependent regulation, Inp53 shifts from associating predominantly with clathrin to interacting with endocytic proteins Sla1, Bzz1, and Bsp1, suggesting that Inp53 mediates stress-specific endocytic events. This response has physiological and molecular similarities to calcineurin-regulated activity-dependent bulk endocytosis in neurons, which retrieves a bolus of plasma membrane deposited by synaptic vesicle fusion. We propose that activation of Ca²⁺/calcineurin and PI(4,5)P₂ signaling to regulate endocytosis is a fundamental and conserved response to excess membrane in eukaryotic cells.

Introduction

Cells dynamically maintain their surface by adding membrane through exocytosis and retrieving it through endocytosis. In growing cells, coordination of these processes ensures that surface area increases to keep pace with cellular volume. In contrast, a rapid decrease in cell size generates excess plasma membrane. During hypertonic shock, for example, cells of the budding yeast Saccharomyces cerevisiae lose up to 50% of their volume, which causes the plasma membrane to buckle and form large, sheet-like invaginations (Kopecka et al., 1973; Morris et al., 1986; Slaninova et al., 2000; Dupont et al., 2010). In response to these extreme conditions, the cell surface is rapidly remodeled, normal morphology is restored, and growth resumes. Much of this regulation must target the actin cytoskeleton, which is essential for both exocytosis and endocytosis in S. cerevisiae. However, how these processes are regulated in response to stress is not well understood.

During normal growth, the yeast actin cytoskeleton is polarized; actin cables, which deliver secretory vesicles, and mobile cortical actin patches, which are sites of clathrin-mediated endocytosis (CME) and cell wall synthesis, both concentrate in the growing bud (Mulholland et al., 1994; Utsugi et al., 2002; Pruyne et al., 2004; Kaksonen et al., 2005). Polarized growth is established and maintained through the activity of small G proteins, Cdc42 and Rhô1, which are activated by TOR/target of rapamycin and directly regulate exocytosis, cell wall synthesis, and actin polymerization through their targets Sec3, Fks1, protein kinase C, and the formins Bni1 and Bnr1 (Levin, 2011; Loewith and Hall, 2011; Bi and Park, 2012). In response to many types of environmental stress, including heat stress and hypertonic shock, the actin cytoskeleton rapidly reorganizes: within 30 min, actin cables disassemble, halting the delivery of...
secretory vesicles, and actin patches spread throughout the surface of both mother and daughter cells (Chowdhury et al., 1992; Palmer et al., 1992). These depolarized actin patches are presumed to function as sites of cell wall synthesis and endocytosis that repair and restore the cell surface. Indeed, after several hours, the cytoskeleton repolarizes, indicating that the cell has adapted to its new environment and is ready to resume growth (Chowdhury et al., 1992). The composition and function of actin patches in stressed cells are incompletely understood; however, in growing cells, the mechanisms that underlie CME have been well elucidated.

CME occurs through a coordinated series of protein–protein interactions, which allows incorporation of cargo into a membrane deformation that matures into an endocytic invagination, using force provided by actin polymerization, and is released into the cytosol as an endocytic vesicle after membrane scission. A well-defined set of proteins act in temporally discrete modules to carry out these events (Weinberg and Drubin, 2012). Coat proteins, including clathrin, are among the earliest to arrive at endocytic sites (Kaksonen et al., 2005). The coat matures with the arrival of cargo and adaptor proteins such as Sla1 (Mahadev et al., 2007; Di Pietro et al., 2010; Feliciano and Di Pietro, 2012). As the endocytic site matures, actin polymerization begins with recruitment of proteins in the WASP/MYO module (Sun et al., 2006). Actin polymerization drives the actin network and associated actin module proteins (such as Bsp1) into the cytosol (Tonikian et al., 2009); the force supplied by actin polymerization promotes formation of a membrane invagination. Scission module proteins are the last to arrive at the mature endocytic site. Actin polymerization, phosphatidylinositol-4,5-bisphosphate (PI(4,5)P2) hydrolysis (carried out by the yeast synaptojanin Inp52/Sjl2), and curvature induced by BAR-domain proteins (Bz1 and Rvs161/Rvs167) together promote membrane scission and release of an endocytic vesicle (Kishimoto et al., 2011). Regulation of PI(4,5)P2 plays an important role throughout endocytosis, as PI(4,5)P2 recruits proteins to the membrane, increases membrane fluidity, and facilitates curvature (Sun et al., 2007; Liu et al., 2009). In a growing cell, these events occur constitutively and lead to continual formation of endocytic vesicles.

Mammalian cells use many of these same principles to carry out endocytosis. The reliance on a network of SH3-dependent protein–protein interactions is highly conserved, as is the role of PI(4,5)P2 and BAR-domain proteins in membrane invagination (Taylor et al., 2011). However, whereas some core endocytic proteins and regulators are conserved, others are organism specific, and the SH3 network in particular evolved rapidly to conserve overall function rather than preserve the identity of participating proteins (Xin et al., 2013). Other differences between endocytosis in mammalian and yeast cells include the role of actin polymerization, which is apparently not essential for all endocytic pathways in animal cells but is universally required in yeast cells which are under turgor pressure (Aghamohammadzadeh and Asycough, 2009; Weinberg and Drubin, 2012). In addition, membrane scission depends on the dynamin GTPase in mammals, whereas the role of Vps1, the yeast dynamin relative, in endocytosis is unclear (Shupliakov et al., 1997; Smaczynska-de et al., 2010, 2012; Weinberg and Drubin, 2012).

How does this complex network of endocytic events respond to the challenges presented by hypertonic shock? When exposed to even mild hypertonic conditions, yeast cells immediately lose water, causing a decrease in turgor pressure, which activates the high-osmolarity glycerol (HOG) mitogen-activated protein kinase signaling pathway. HOG signaling increases intracellular glycerol and promotes long-term osmoadaptation (Schaber et al., 2010; Saito and Posas, 2012), but during extreme hypertonic shock, cells lose volume and decrease in size despite maximal activation of the HOG pathway. Although it is not yet clear how the massive plasma membrane invaginations that result from this cell shrinkage are resolved (Kopecka et al., 1973; Morris et al., 1986; Slaninova et al., 2000; Dupont et al., 2010), the endocytic machinery is likely involved. Many endocytic proteins promote the recovery of actin polarization after hyperosmotic shock, and depolarized actin patches may carry out endocytic events that restore the cell surface.

Several observations indicate that the composition of actin patches changes during osmotic stress. Yeast encodes three synaptojanin homologues—Inp51/Sjl1, Inp52/Sjl2, and Inp53/Sjl3 (Singer-Kruger et al., 1998; Stolz et al., 1998). As noted, Inp52 plays a vital role in CME that occurs in the bud during polarized growth (Stefan et al., 2005). In contrast, Inp53, which is closely related to Inp52, localizes to the trans-Golgi network (TGN), where it binds to clathrin and mediates endosomal protein sorting, a process that is phosphoinositide dependent (Ha et al., 2001, 2003; Daboussi et al., 2012). However, in response to osmotic stress, Inp53 localizes to actin patches, and overexpression of either Inp53 or its homologue Inp52 hastens repolarization of the actin cytoskeleton under these conditions (Ooms et al., 2000). Rvs167/ampiphysin and Bzz1/syndapin are also required to restore the polarized actin cytoskeleton after hyperosmotic shock (Bauer et al., 1993; Soulard et al., 2002), and several mutants lacking endocytic proteins are defective for growth in hypertonic conditions (including inp52A, inp53A, rvs167A, sla1Δ, and act1-1; Chowdhury et al., 1992; Srinivasan et al., 1997; Dudley et al., 2005; Yoshikawa et al., 2009). Thus hyperosmotic stress modulates actin patches, and these structures may resolve the membrane invaginations caused by volume loss through stress-specific endocytic events.

Examining the response to hyperosmotic stress may therefore reveal signaling mechanisms that regulate endocytosis. We noted that high osmolarity triggers a rise in cytosolic Ca2+ in yeast, mediated in part by release of vacuolar Ca2+ through the Yec1 channel (Denis and Cyert, 2002). To investigate possible targets of this Ca2+ signal, we examined whether calcineurin (CN), the conserved Ca2+/calmodulin–activated protein phosphatase and target of the immunosuppressant drugs FK506 and cyclosporine A (Roy and Cyert, 2013), was active during hyperosmotic shock. The CN heterodimer is made up of a catalytic subunit (Cna1 or Cna2) and a required Ca2+-binding regulatory subunit (Cnb1). CN is not active under standard growth conditions but promotes survival during environmental stress conditions, such as high concentrations of Na+ and Li+ and high pH (Cyert and Philpott, 2013). In response to a Ca2+ signal, Ca2+-bound calmodulin interacts with and activates CN, which must physically associate with its substrates via conserved docking motifs during phosphorylation (Roy and Cyert, 2009; Grigoriu et al., 2013). CN regulates gene expression by promoting accumulation of the Cza1 transcription factor in the nucleus (Stathopoulos and Cyert, 1997; Stathopoulos-Gerontides et al., 1999) and stimulates specific endocytic events mediated by the α-arrestins Aly1 and Rod1 (O’Donnell et al., 2013; Alvaro et al., 2014). Furthermore, recent systematic identification of CN substrates in yeast revealed several proteins involved in polarized growth and actin regulation, including the CN-interacting protein synaptojanin Inp52 (Goldman et al., 2014).

In this work, we demonstrate that CN accumulates at sites of polarized growth during hyperosmotic shock and dephosphorylates synaptojanin/Inp52 to promote actin polarization and maintain membrane morphology. Localized activation of synaptojanin is also suggested by a dramatic redistribution of PI(4,5)P2 in the membrane observed during osmotic shock that is calcineurin regulated.
Osmotic stress induces Inp53 to associate with multiple endocytic proteins, and some of these interactions are promoted by CN. Synaptotagmin is one of only two proteins identified as CN substrates in both yeast and mammals. In neurons, CN dephosphorylates synaptotagmin and other proteins, termed dephosphins, to stimulate the activity-dependent bulk endocytosis (ADBE) that follows multiple rounds of synaptic vesicle fusion events (Clayton and Cousin, 2009). We suggest that in both neurons and osmotically stressed yeast cells, a sudden excess of plasma membrane triggers a conserved, CN-regulated response that restores the cell surface.

RESULTS
Calcineurin localizes to sites of polarized growth during hyperosmotic stress
Mammalian cells encode scaffold proteins that interact with CN and substrates to target the phosphatase to specific subcellular compartments. For example, in neurons, AKAP79 localizes CN to L-type Ca^{2+} channels at the plasma membrane (Oliveria et al., 2007). In contrast, under standard growth conditions, yeast CN localized diffusely throughout the cell, as visualized in cells expressing the functional Cna1-3x green fluorescent protein (Cna1-3GFP) fusion protein from its endogenous locus (Figure 1A). Because CN is active during environmental stress, we also examined its distribution under a range of stress conditions. These analyses revealed that Cna1-3GFP distribution changed dramatically 10 min after addition of 1.25 M KCl to the growth medium, accumulating at foci throughout the cell body, the bud tip, and the bud neck. Cna2-3GFP, a functionally redundant catalytic subunit isoform, displayed similar localization changes under these conditions (Supplemental Figure S1A). This redistribution of CN was triggered by hypertonic shock, as NaCl (1.25 M), sorbitol (2 M), and KCl (1.25 M) all provoked the same response (Figure 1A), which occurred only during intense hyperosmotic stress (KCl > 0.75 M) and became more pervasive as osmotic strength increased (Supplemental Figure S1B). However, CN remained cytosolic in cells exposed either to 200 mM LiCl, a condition under which CN is required for survival that does not induce hyperosmotic shock (Figure 1A and Supplemental Figure S1C), or 50 mM CaCl_{2}, which activates CN-dependent transcription without inducing osmotic stress (Figure 1A). Furthermore, hypertonic shock caused redistribution of Cna1-3GFP in cells treated with the CN inhibitor FK506 (Figure 1A) and in a CN-deficient mutant that lacks the regulatory subunit (cnb1; Supplemental Figure S1D). Therefore, the cellular localization of CN changes specifically in response to intense hyperosmotic shock, and this redistribution in independent of CN activity. Surprisingly, the observed changes in CN localization did not depend on the HOG pathway and occurred robustly in hog1, pbs2, and ssh2 mutants exposed to hyperosmotic challenge (Supplemental Figure S1E). The requirement for intense osmotic shock to redistribute CN also distinguishes this response from HOG pathway activation, which occurs at much lower osmolarities (Schaber et al., 2010).

After hyperosmotic shock, CN localization to cytoplasmic foci, the bud neck, and the bud tip followed different kinetics (p < 0.0001; Figure 1C). Ten minutes after hyperosmotic shock, CN localized to foci in 100% of cells, but this pattern was transient and disappeared completely by 4 h. Some foci showed colocalization with clathrin (Chc1–red fluorescent protein [RFP]; Supplemental Figure S1G). CN-containing foci also showed partial colocalization with the actin-binding protein Abp1-RFP, which serves as a marker for endocytic patches (Supplemental Figure S2A). In contrast, CN localization to the bud tip and bud neck peaked later (60% of budded cells at 30 min) and was more prolonged than the observed foci; a significant fraction of cells retained Cna1-3GFP at the bud neck after 4 h of hyperosmotic stress (Figure 1, B and C). Thus cellular CN foci may reflect a response to hyperosmotic stress that is functionally distinct from its localization to the bud neck and bud tip. Overall, changes in CN distribution suggest that its interactions substantially change during hyperosmotic shock and in particular include colocalization with the actin cytoskeleton.

In *S. cerevisiae*, the bud tip and bud neck are sites of polarized growth, which stimulate polymerization and organization of the polarized actin cables that deliver the bulk of cellular secretory vesicles to the bud (Pruyne et al., 2004). To determine whether the actin cytoskeleton is required for polarized localization of CN, we exposed a uniform population of large-budded cells, obtained after release from mating pheromone arrest, to osmotic stress. Under these conditions, CN localization to the bud neck was extremely consistent (Figure 1D). Fluorescence intensity of Cna1-3GFP in a line along the mother/bud axis was averaged, and after a 10-min hyperosmotic shock, two clear Cna1-3GFP peaks on either side of the bud neck were visible (Figure 1E; N = 104 cells). However, depolymerization of the actin cytoskeleton with latrunculin A abolished localization of Cna1-3GFP to the bud neck (Figure 1E and Supplemental Figure S1F). Thus localization of CN to sites of polarized growth requires an intact actin cytoskeleton.

Hypertonic shock induces distinct calcineurin signaling events
A role for CN in the response to hyperosmotic stress had not been previously appreciated. Although CN is required for survival during both NaCl and LiCl stress, in part due to CN-dependent expression of the ENA1-encoded plasma membrane Na^+/Li^+-ATPase (Mendoza et al., 1994), loss of CN has no effect on long-term growth under nontoxic hyperosmotic stresses such as KCl (Supplemental Figure S1C). Despite this, the cytoplasmic Ca^{2+} signal (Denis and Cyert, 2002) and dramatic localization changes induced by hyperosmotic stress suggested that CN might function during the early response to these conditions.

Under many environmental conditions, CN ensures cell survival by activating the Crz1 transcription factor, promoting its translocation from the cytosol to the nucleus (Stathopoulos and Cyert, 1997; Stathopoulos-Gerontides et al., 1999). Therefore we measured CN-Crz1–dependent expression of 4x-CDRE-LacZ, a reporter gene controlled by the Crz1- binding site, during hyperosmotic stress. As expected, addition of CaCl_{2} caused a 40-fold induction of CDRE-LacZ expression (Figure 2). In contrast, no activation of CN-Crz1 signaling was observed in response to hyperosmotic stress despite the increase in intracellular Ca^{2+} that occurs under these conditions (Denis and Cyert, 2002). Instead, hyperosmotic stress blocked Ca^{2+}-dependent activation of Crz1 (Figure 2). This surprising effect suggested that hypertonic conditions inhibit CN/Crz1 signaling, perhaps in part by relocating CN and preventing its access to Crz1.

Thus several unique features of CN signaling, including its localization to sites of polarized growth and failure to activate Crz1-dependent gene expression, suggest that hypertonic shock activates distinct calcineurin-regulated events. We next sought to identify these CN-regulated processes and substrates.

Calcineurin regulates actin rearrangements during hyperosmotic shock
The depolarization and subsequent repolarization of the actin cytoskeleton is a prominent and acute cellular response to hyperosmotic shock (Chowdhury et al., 1992). CN localization to sites of polarized growth and partial colocalization with Abp1 suggested that the phosphatase might regulate the actin cytoskeleton during hyperosmotic shock. To test this idea, cells were exposed to
FIGURE 1: Calcineurin localizes to sites of polarized growth during hyperosmotic stress. (A) Micrographs of wild-type (BY4741) cells with integrated Cna1-3GFP. Cells were treated as indicated for 10 min before visualization; where indicated, cells were pretreated with 1 μg/ml FK506 for 30 min; scale bars, 5 μm. (B) Localization of Cna1-3GFP during prolonged 1.25 M KCl stress; arrows: bud neck, arrowheads: bud tip, scale bars, 5 μm. (C) Quantification of percentage of cells with foci or bud neck/bud tip CN during 1.25 M KCl stress; N > 100 at each time point. Loss of foci and neck/tip localization was evaluated by fitting one-phase exponential decay, yielding $t_{1/2}$ foci = 26.8 ± 0.6 min, $t_{1/2}$ neck/tip = 36 ± 3.4 min;
Calcineurin-activated gene expression is inhibited during hyperosmotic shock. Ca^{2+}/CN-dependent Crz1 transcription was measured with a CDRE-LacZ reporter. β-Galactosidase activity normalized to protein concentration is reported. Cells were treated for 90 min with 200 mM CaCl_{2} and/or 1.25 M KCl as indicated. Error bars are SD; **p < 0.001.

1.25 M KCl, fixed at 30-min intervals, stained for actin with rhodamine–phalloidin, and polarization quantified using image-analysis software (see Materials and Methods). Control cells were maximally depolarized (50%) 2 h after exposure to 1.25 M KCl, and polarity was largely restored by 4 h (Figure 3, A and B). Inactivation of CN with FK506 exacerbated depolarization at 120 and 150 min; however, as in control cells, polarity was reestablished by 4 h (Figure 3, A and B). Analyzing the distributions of cellular fluorescence ratios at 120 and 150 min after hyperosmotic shock showed that FK506 significantly decreased polarization compared with the control (Figure 3C; p < 10^{-6}). Thus CN promotes repolarization of the actin cytoskeleton after hyperosmotic challenge.

Inp53 is a calcineurin substrate

Because the altered subcellular distribution of CN likely reflected a change in protein–protein interactions, we sought to identify proteins whose association with CN increased during hyperosmotic stress. Mass spectrometry was used to identify proteins that copurified with endogenously expressed, epitope-tagged CN (Cna2-TEV-ZZ) isolated from either unstressed cells or cells exposed to 1.25 M KCl, as compared with purifications from untagged cells, which controlled for nonspecific background. These analyses suggested a possible interaction between CN and Inp53/Sjl3, one of three yeast synaptojanins (Figure 4A; Srinivasan et al., 1997; Stolz et al., 1998), which was enhanced during hyperosmotic stress. In unstressed cells, Inp53 contributes to protein sorting at the TGN but localizes to actin patches and stimulates repolarization of the actin cytoskeleton during osmotic stress (Ooms et al., 2000). Thus we hypothesized that CN might promote actin polarization during hyperosmotic stress by regulating Inp53.

To examine CN-Inp53 interaction, we isolated glutathione S-transferase (GST)–Inp53 from yeast extracts and confirmed that it copurified with Cna2 and that more Cna2 associated with GST-Inp53 purified from cells exposed to hyperosmotic stress (Figure 4B). Inp53 contains two distinct phosphoinositide phosphatase domains (SAC and IP5Pase; Guo et al., 1999; Figure 4A), as well as a C-terminal proline-rich tail domain (PRD) that mediates protein–protein interactions (Ha et al., 2003). Expressing each domain of Inp53 separately and assessing its copurification with CN established that the CN-Inp53 interaction was mediated by the SAC domain (unpublished data). CN interacts with many of its substrates and regulators via a conserved PxixIT docking motif (Roy and Cyert, 2009). A match to the PxixIT consensus (Goldman et al., 2014; PRVQIIT_{300-305}, occurs in a surface-exposed loop of the Inp53 SAC domain (Manford et al., 2010), as well as in this region of Inp52 and synaptojanin homologues from a broad range of fungal species (Figure 4A and Supplemental Figure S3C; Wapinski et al., 2007). Mutation of this sequence to ARAQAA substantially reduced, but did not completely eliminate, copurification of CN with Inp53 (Figure 4B). This mutated allele is hereafter referred to as Inp53\_ARAQAA. Despite the physical interaction between Inp53 and CN, substantial colocalization between Cna1-3GFP and Inp53-3RFP was not observed during osmotic shock (Supplemental Figure S2, B and D), nor did disruption of CN binding with Inp53\_ARAQAA block Cna1-3GFP localization to sites of polarized growth (Supplemental Figure S2, C and D).

Purification of GST-Inp53 from cell extracts revealed that its phosphorylation increased in cells exposed to hyperosmotic stress (Figure 4C). Examination of each Inp53 domain showed that only phosphorylation of the PRD increased under these conditions (Supplemental Figure S3A). Furthermore, enhanced PRD phosphorylation was observed in extracts of FK506-treated cells, demonstrating that CN regulates PRD phosphorylation both in the presence and absence of hyperosmotic stress. Surprisingly, this phosphorylation was not due to Hog1, as osmotic stress-induced phosphorylation of PRD appeared to be equivalent in wild-type (WT) and hog1 yeast (Supplemental Figure S3B). No effect of FK506 on the phosphorylation of full-length GST-Inp53 was detected, possibly due to its large size and high level of overall phosphorylation. Thus, during hyperosmotic stress, the Inp53 PRD becomes heavily phosphorylated, and CN, also active under these conditions, promotes its dephosphorylation.

Taken together, the phosphorylation and interaction data strongly suggest that CN directly regulates Inp53 in vivo and that their association is specifically enhanced during hyperosmotic stress. Furthermore, because the CN-Inp53 interaction is disrupted in Inp53\_ARAQAA, this allele should display impaired regulation by CN when expressed in vivo.

Calcineurin promotes actin polarization via regulation of Inp53

To examine regulation of Inp53 by CN in the absence of partially redundant Inp51 and Inp52, we used an inp51 inp52 triple mutant; the viability of this strain (hereafter inp53\_ΔΔΔ) was maintained by a galactose-regulated copy of INP53 that was stably integrated into the genome. As expected, inp53\_ΔΔΔ cells expressing a control vector were inviable when grown in dextrose-containing medium (Figure 5A). INP53 or INP53\_ARAQAA was introduced into inp3\_ΔΔΔ on either a low- or high-copy plasmid and provided the sole source of synaptojanin during growth on dextrose. In the
absence of hyperosmotic stress, inpΔΔΔ cells expressing high-copy INP53 or INP53ARAQAA grew at a rate similar to wild-type cells and, like wild-type cells, showed no change in growth rate in the presence of FK506. In contrast, inpΔΔΔ cells expressing low-copy Inp53 were viable but grew slowly, indicating that synaptojanin activity was limiting. Expression of INP53ARAQAA at low-copy resulted in even slower growth, suggesting that interaction with CN was required for full activity of Inp53. FK506 further inhibited growth of cells expressing either INP53 or INP53ARAQAA, and neither strain grew appreciably in the presence of 1.25 M KCl (unpublished data). These data suggest that under synaptojanin-limiting conditions, CN is required for growth and that one of its essential functions is to positively regulate Inp53.

We then examined these cells to determine whether CN promotes actin polarization by regulating Inp53. Polarization of the actin cytoskeleton was compared in inpΔΔΔ cells expressing either INP53 or INP53ARAQAA on a low-copy plasmid (Figure 5, B and C). Under normal growth conditions, Inp53 was sufficient to polarize actin in these cells. However, inhibiting CN with FK506 resulted in actin depolarization. Furthermore, the actin cytoskeleton was depolarized in the majority of inpΔΔΔ cells expressing low-copy INP53ARAQAA, with little further depolarization when CN was inhibited with FK506 (p < 0.0001; Figure 5C, inset). Inp53 and Inp53ARAQAA were expressed equivalently (Supplemental Figure S4). These results show that when synaptojanin activity is limiting, activation of Inp53 by CN is required to maintain a normally polarized actin cytoskeleton.

**Regulation of Inp53 by calcineurin affects membrane morphology**

Inp53 exerts its effects on actin polarity by dephosphorylating PI(4,5)P2. This phospholipid is a specific component of the plasma membrane (Ooms et al., 2000; Stefan et al., 2002), and mutants lacking multiple synaptojanins display aberrant plasma membrane invaginations due to increased levels of PI(4,5)P2 and a defect in synaptojanin-mediated membrane scission (Singer-Kruger et al., 1998; Sun et al., 2007). To determine whether regulation of Inp53 by CN affects plasma membrane morphology, we visualized the plasma membrane in inpΔΔΔ cells using the PI(4,5)P2-binding biosensor GFP-2xPH(PLCδ) (Stefan et al., 2002). In contrast to the normal, smooth plasma membranes observed in inpΔΔΔ cells expressing high-copy INP53, inpΔΔΔ cells expressing a vector, and thus depleted of INP53, formed massive, PI(4,5)P2-containing membrane deformations (Figure 6A). To quantify these structures, we computationally defined the membrane region of each cell using GFP-2xPH(PLCδ) intensity and counted regions that extended >10 pixels (~0.5 μm) into the cytoplasm (Figure 6, A and B). Synaptojanin-limited cells expressing low-copy INP53 had slightly more membrane invaginations per cell than the high-copy control, and inhibiting CN with FK506 exacerbated this defect. inpΔΔΔ cells expressing low copy INP53ARAQAA, which disrupts CN binding, displayed even more severe membrane deformation, which was further exacerbated by incubation with FK506. Thus analyses of both polarized actin distribution and plasma membrane morphology in cells expressing Inp53 as the sole synaptojanin revealed that CN must bind to and dephosphorylate Inp53 to achieve full synaptojanin function.
Calcineurin regulates PI(4,5)P<sub>2</sub> distribution in the membrane during hyperosmotic shock

In wild-type cells, CN interaction with and dephosphorylation of Inp53 is specifically stimulated during hyperosmotic stress, suggesting that CN regulates PI(4,5)P<sub>2</sub> pools under these conditions. Total PI(4,5)P<sub>2</sub> levels do not change significantly during osmotic stress (Bonangelino et al., 2002), suggesting that CN-dependent activation of Inp53 may instead result in altered subcellular distribution of PI(4,5)P<sub>2</sub>. We used the GFP-2xPH(PLCδ) biosensor to investigate PI(4,5)P<sub>2</sub> distribution during osmotic shock.

In unstressed WT cells, GFP-2xPH(PLCδ) was smoothly distributed throughout the plasma membrane and uniformly brighter than the adjacent cytoplasm (Figure 7A). In contrast, after a 10-min hyperosmotic shock, distribution of GFP-2xPH(PLCδ) at the membrane dramatically changed; fluorescence was confined to a few bright patches in each cell, which were separated by large regions depleted of GFP-2xPH(PLCδ), where fluorescence was not significantly higher than in the adjacent cytoplasm (Figure 7A). This change in GFP-2xPH(PLCδ) distribution was quantified by determining the percentage of total membrane area in each cell that displayed significant enrichment of GFP-2xPH(PLCδ) fluorescence relative to the adjacent cytoplasm (Figure 7B; see Materials and Methods). This analysis captured the uniform, bright GFP-2xPH(PLCδ) localization in unstressed cells: on average, >70% of the membrane showed enriched GFP-2xPH(PLCδ) fluorescence. In contrast, after hyperosmotic shock, only 17% of the membrane, on average, showed enriched GFP-2xPH(PLCδ) (Figure 7; p < 0.0001). These results document a major redistribution of PI(4,5)P<sub>2</sub> into enriched and depleted membrane domains during osmotic shock, which may also underlie changes in membrane morphology previously described by electron microscopy (Dupont et al., 2010).

The extent of GFP-2xPH(PLCδ) depletion during osmotic shock also depended on CN: when CN was inhibited, 24% of the membrane retained enriched GFP-2xPH(PLCδ), a significantly greater area than in control cells (Figure 7, A–C; p < 0.0001). This result is consistent with CN-dependent activation of Inp53 during osmotic shock enhancing the depletion of PI(4,5)P<sub>2</sub> from large regions of the membrane and likely represents an important mechanism by which the cells adjust to the membrane stress induced by rapid volume loss.
Molecular mechanisms of Inp53 regulation by calcineurin

The phenotypic effects of the CN-binding deficient Inp53<sup>ARAGAA</sup> mutant and of CN inhibition suggest that in vivo, CN acts to activate Inp53. We next investigated whether CN activates Inp53 by altering its intrinsic catalytic activity or by regulating its interactions with other proteins.

To examine whether hyperphosphorylation induced by hyperosmotic stress or dephosphorylation by CN directly altered Inp53 catalytic activity, we purified Inp53 from yeast under conditions that maintained its phosphorylation state and SAC and IP5Pase domain activities (Figure 8A; Guo et al., 1999) and analyzed the kinetics of PI(4,5)P<sub>2</sub> dephosphorylation by Inp53 in vitro. GST-Inp53 and GST were purified from cells treated either with vehicle or FK506 and grown in the presence or absence of hypertonic shock. GST-Inp53, but not GST, exhibited PI(4,5)P<sub>2</sub> phosphatase activity in vitro (Figure 8B and Supplemental Figure S5) which was statistically equivalent in all preparations tested (Figure 8B). Thus we find no evidence that the intrinsic catalytic activity of Inp53 is altered either by the massive hyperphosphorylation that occurs after osmotic shock or by subsequent dephosphorylation via CN. However, changes in PI(4,5)P<sub>2</sub> distribution in the membrane suggest that sites of Inp53 action are reorganized.

Therefore we tested whether any of the established or predicted protein–protein interactions for Inp53 were altered by hyperosmotic shock or its regulation by CN. In unstressed cells, Inp53 functions primarily to regulate phosphoinositide-dependent sorting of proteins at the TGN and interacts directly with clathrin via an LLDID motif in the Inp53-PRD (Ha et al., 2001, 2003; Daboussi et al., 2012). Equivalent amounts of clathrin heavy chain (Chc1) copurified with Inp53 and Inp53<sup>ARAGAA</sup>, but copurification of Chc1 with both proteins decreased after hyperosmotic shock (Figure 9A). Hyperphosphorylation of the Inp53-PRD may be responsible for reduced clathrin binding; we identified numerous phosphorylation sites in the Inp53-PRD that increased during osmotic stress, including S<sub>914</sub>, which immediately precedes LLDID<sub>915-919</sub> (unpublished data). Loss of interaction with clathrin may indicate a repurposing of Inp53 from its normal role at the TGN to a function at actin patches and is consistent with reports of Inp53 translocation to actin patches during hyperosmotic shock (Ooms et al., 2000).

We next investigated whether Inp53 interacts with actin patch/endocytic proteins. Like synaptojanins Inp52 and Inp53, Rvs167, a yeast homologue of amphiphysin/endophilin, promotes repolarization of the actin cytoskeleton during hyperosmotic stress, and rvs167 mutants are sensitive to hyperosmotic stress (Bauer et al., 1993). Inp53 and Rvs167 show genetic interactions (Aguilar et al., 2010), but no physical interaction between either Inp52 or Inp53 and Rvs167 has been reported. We found that Rvs167-GFP copurified equivalently with both GST-Inp53 and GST-Inp53<sup>ARAGAA</sup> (Figure 9B). There was no significant change in Rvs167-Inp53 association under hyperosmotic conditions, but Rvs167 showed slower electrophoretic mobility during hyperosmotic stress, suggesting that it may be phosphorylated or otherwise modified under these conditions.

Inp53 was previously reported to interact with the actin patch components Bsp1, Bzz1, and Slal in yeast two-hybrid analyses (Wicky et al., 2003; Tonikian et al., 2009). Bzz1 and Slal both contain SH3 domains that are predicted to interact with the Inp53-PRD (Tonikian et al., 2009). Bsp1 is an endocytic protein of unclear function that binds the SAC domain of Inp52 and Inp53 and arrives late during internalization at endocytic sites along with other components of the actin module (Wicky et al., 2003; Tonikian et al., 2009). Bzz1/syndapin stimulates Las17/WASP-mediated actin polymerization and recognizes shallow membrane curvature through its F-BAR domain. We were unable to detect either Bsp1 or Bzz1 copurification with Inp53 in unstressed cells but saw substantial copurification of both proteins with Inp53 during hyperosmotic...
shock (Figure 9, C and D); neither the Inp53-Bsp1 nor the Inp53-Bzz1 interaction was substantially altered by disruption of the CN binding site in Inp53\(^{\text{ARAQAA}}\) (Figure 9, C and D). Bzz1 levels did not change during hyperosmotic shock, suggesting that its ability to interact with Inp53 was enhanced during stress. By contrast, the amount of Bsp1 in the soluble fraction of our extracts changed dramatically. Under all conditions, the majority of Bsp1 was present in the insoluble pellet fraction. However, the soluble pool of Bsp1 increased significantly in extracts of cells exposed to hyperosmotic stress, which may explain its copurification with Inp53 under these conditions (Figure 9C). Thus interaction of Inp53 with both Bsp1 and Bzz1 occurs specifically during osmotic shock.

Finally, we found that Inp53 interacts with the endocytic coat protein Sla1/intersectin. Sla1 has diverse functions during endocytosis, including roles in cargo recognition, where it binds the NPFXD motif, as a regulator of the actin nucleation-promoting factor Las17/WASP, and as a clathrin-binding adaptor (Mahadev et al., 2007; Piao et al., 2007; Feliciano and Di Pietro, 2012). Inp53 copurified with Sla1/intersectin in unstressed cells, and this interaction was substantially enhanced during hyperosmotic stress. By contrast, the CN-binding–deficient Inp53\(^{\text{ARAQAA}}\) copurified with the same low amount of Sla1 when purified from either stressed or unstressed cells (Figure 9E). Thus the Inp53-Sla1 interaction is enhanced during hyperosmotic stress, and this effect is dependent on CN. Despite the prediction that SH3 domains in both Bzz1 and Sla1 should bind to proline-rich motifs in the Inp53-PRD, the Inp53-PRD alone was insufficient to copurify either Bzz1 or Sla1 (Supplemental Figure S6). Instead, interactions between Inp53, Sla1, and CN likely occur in part of a larger complex. Consistent with this idea, Sla1 contains numerous phosphorylation sites that are regulated by CN in vivo but lacks a predicted CN docking motif (Goldman et al., 2014). Thus, after hypertonic shock, Inp53 may form a complex with both Sla1 and CN that facilitates Sla1 and Inp53 dephosphorylation.

**DISCUSSION**

Membrane structure must be restored in response to hyperosmotic shock

Hyperosmotic shock causes dramatic changes throughout the cell and requires a broad response coordinated by multiple signaling pathways. At the center of this response, the HOG pathway acts to restore turgor pressure and control gene expression (Saito and Posas, 2012). Independently, the cell wall sensor Wsc1 promotes depolarization of the actin cytoskeleton (Balguerie et al., 2002) and induces a widespread secretory arrest (Nanduri and Tartakoff, 2001a,b); depolymerization of actin cables liberates monomeric
For each condition, bars represent SEM. ***membrane in each cell that was enriched for GFP-2xPH(PLC)
cytoplasm are marked in green; see Materials and Methods. (C) The percentage of total membrane in
each cell that was enriched for GFP-2xPH(PLC) was averaged and plotted. Error bars represent SEM. ***p < 0.0001; one-way ANOVA and Tukey's multiple comparison posttest.
For each condition, N > 700 cells, from four independent experiments.

actin, which stimulates Ssk2 to activate the polarisome (Yuzuyk, 2002, 2003). We show here that the Ca²⁺/calmodulin–dependent phosphatase CN also functions during osmotic stress, which was not previously appreciated.

Hypertonic shock induces distinct physiological responses, depending on its severity. As osmotic challenge increases to ~500 mM KCI, loss of cell volume is mild because turgor pressure is only partially depleted. Over this range, the HOG pathway becomes increasingly active to restore and maintain turgor pressure (Schaber et al., 2010; Saito and Posas, 2012; Lee et al., 2013). More intense osmotic stress leads to complete loss of turgor, and for increasing osmotic challenge beyond this point, volume loss is extreme. Dramatic cell shrinkage confines the plasma membrane to a much-reduced surface area, causing deep invaginations to form (Dupont et al., 2010) and presenting an acute challenge that must be resolved rapidly and with precise spatial control.

Ca²⁺ signaling is well suited to regulate such responses. Hyperosmotic shock triggers a rapid and transient Ca²⁺ signal (Denis and Cyert, 2002) and promotes an immediate redistribution of CN, mediating its interaction with specific targets, such as Inp53. Similarly, the uniform distribution of PI(4,5)P₂ in the membrane quickly changes into a mosaic of depleted and PI(4,5)P₂–enriched domains, which CN regulates, potentially through Inp53. These PI(4,5)P₂

Juvvadi et al., 2011; Kozubowski et al., 2011). The mechanism(s) responsible for the CN localization changes caused by hyperosmotic stress are unknown, as significant colocalization of CN and Inp53 was not observed (Supplemental Figure S2, B and C). However, similar approaches to those described here will likely identify additional scaffold candidates. We also suggest that interactions between Inp53 and CN during hyperosmotic stress bring CN into proximity with substrates like Bsp1 and Sla1 that lack canonical docking motifs but show CN-dependent dephosphorylation (Goldman et al., 2014). Thus hyperosmotic shock illustrates how a specific environmental condition can tune CN signaling by modifying localization and substrate access.

Regulation of endocytosis during hyperosmotic shock
This study provides some initial insights into molecular events that occur at depolarized actin patches in cells responding to intense osmotic shock. Endocytosis is regulated by a densely interconnected network of proteins (Johnson and Hummer, 2013). During osmotic stress, we show that components of this network are dephosphorylated by CN and are also antagonistically hyperphosphorylated. Key interactions between Inp53 and other proteins involved in scission and actin force generation (Sla1 and Bzz1) are altered, and the interaction of Inp53 with clathrin is reduced, presumably

FIGURE 7: Calcineurin regulates PI(4,5)P₂ distribution during osmotic shock. (A) Representative images of wild-type (BY4741) cells expressing GFP-2xPH(PLC). Cells were treated with 1 μg/ml FK506 or vehicle for 30 min; osmotic shock was induced with 1.25 M KCl for 10 min where indicated. (B) Membrane regions with >10% enriched GFP-2xPH(PLC) compared with adjacent cytoplasm are marked in green; see Materials and Methods. (C) The percentage of total membrane in each cell that was enriched for GFP-2xPH(PLC) was averaged and plotted. Error bars represent SEM. ***p < 0.0001; one-way ANOVA and Tukey’s multiple comparison posttest.

patches are well suited, by binding and activating a host of known effector proteins, to pattern the localized events required to restore the integrity of the cell surface.

New insights into CN signaling specificity in S. cerevisiae
A diverse array of >40 proteins involved in polarized growth, cell cycle regulation, mating, autophagy, and many other cellular processes have been identified as CN targets (Goldman et al., 2014). Where and when these proteins are regulated by CN have yet to be elucidated. We show here that CN undergoes major localization changes during hyperosmotic shock, which may restrict the substrates it dephosphorylates. In particular, we find that CN colocalizes with a subset of actin patches, perhaps indicating transient localization to sites of endocytosis, and binds preferentially to Inp53 during hyperosmotic shock. In contrast, activation of the CN-dependent Crz1 transcription factor is blocked (Figure 10A). Intense osmotic shock delays multiple signaling events, including Crz1 nuclear translocation, possibly because of impaired diffusion caused by water loss (Miermont et al., 2013). However, changing CN’s binding partners also affects downstream signaling; artificially strengthening the CN–Crz1 interaction, for example, causes growth defects by blocking the interaction of CN with other substrates (Roy et al., 2007). Therefore CN signaling outcomes during hyperosmotic stress are likely specified by its subcellular distribution and protein–protein interactions, in agreement with important roles for CN localization in other fungi and in mammals (Oliveria et al., 2007;

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directing it away from the TGN (Figure 10B). This stress-specific remodeling may adjust the endocytic machinery to the unique physiological circumstances that accompany intense hypertonic stress. For example, loss of turgor reduces the requirement for actin-generated force during endocytosis (Aghanomhamedazadeh and Ayscough, 2009), and the deep membrane invaginations observed after intense osmotic stress suggest that a large amount of membrane may need to be internalized. Although the mechanisms by which osmotically stressed yeast resolve these invaginations have not been elucidated, they may be similar to the process of ADBE, which has been well studied in neurons.

During periods of intense stimulation, massive release of synaptic vesicles deposits a large amount of membrane at the cell surface (Meunier et al., 2012; Kittelmann et al., 2013), which cannot be efficiently retrieved by CME. Instead, ADBE is induced to pinch off large membrane invaginations, forming endosome-like compartments (Cousin, 2009; Saheki and De Camilli, 2012). ADBE is activated by Ca\(^{2+}\)/CN (Evans and Cousin, 2007) and may be physiologically similar to the CN-dependent processes that are required to maintain yeast viability after intense hypertonic shock.

To carry out ADBE, the cell repurposes many of the same components that participate in CME, that is, synaptojanin, endophilin, syndapin, dynamin, and intersectin (Mani et al., 2007; Clayton et al., 2009; Winther et al., 2013). CN initiates ADBE by dephosphorylating synaptojanin, dynamin, and other proteins collectively termed the dephosphins. Dephosphorylation promotes synaptojanin–endophilin and dynamin–syndapin binding (Mani et al., 2007; Clayton et al., 2009). Both of these interactions occur and are specifically required only during ADBE (Figure 9C). In particular, syndapin, which contains an F-BAR domain, stabilizes shallow membrane curvature and may facilitate formation and/or resolution of the large membrane invaginations unique to ADBE. In addition, Cdk5 kinase and CN phosphatase activity are simultaneously required during ADBE (Evans and Cousin, 2007). Similarly, we observe both hyperphosphorylation and CN-mediated dephosphorylation of the Inp53-PRD during hypertonic shock, suggesting that coordination between CN and antagonistic kinases may be required to temporally or spatially regulate the activity of Inp53. Thus the parallel recruitment/interaction with the homologous yeast proteins Inp53 (synaptojanin), Rvs167 (amphiphysin/endophilin), Bzz1 (syndapin), Sia1 (intersectin) and CN during hypertonic stress suggest that this yeast response is strikingly similar to ADBE at the molecular level.

Although CN is well conserved between yeast and humans, with 44% sequence identity (Cyert et al., 1991), its functions and substrates have significantly diverged. Among >70 CN targets, synaptojanin/Inp53 is one of only two proteins identified as CN substrates in both yeast and mammals (Goldman et al., 2014). Thus regulation of synaptojanin may be one of CN’s primordial functions, and we hypothesize that the activation of Ca\(^{2+}\)/CN and PI(4,5)P\(_2\) signaling to regulate endocytosis is a fundamental and conserved response to excess membrane in eukaryotic cells.

**MATERIALS AND METHODS**

**Growth media and general methods**

Yeast media and culture conditions were essentially as previously described, except that twice the levels of amino acids and nucleotides were used in synthetic complete medium (Sherman, 1991). Yeast transformations were carried out by the lithium acetate method (Ausubel et al., 1991). For 4x-CDRE-lacZ reporter assays, cells were grown to mid log phase, and CaCl\(_2\) or KCI was added for 2 h as noted. β-Galactosidase activity was measured as previously described (Bultynck et al., 2006). Values represent an average of three independent extracts, each measured in triplicate. Error bars indicate the SD. Yeast strains are listed in Supplemental Table S1a; strains from the yeast deletion collection were purchased from Open Biosystems (Huntsville, AL). Plasmids used in this study are listed in Supplemental Table S1b. FK506 was used from a 10 mg/ml stock dissolved in 9:1 EtOH/Tween-20 and used at 1 μg/ml (LC Laboratories, Woburn MA).

**Yeast strain construction**

YEg1 was created from YMJ38 (provided by Maria Molina, Universidad Complutense, Madrid, Spain) by transformation with the pCNA1::URA3hisG deletion cassette and counterselected on 5-fluoroorotic acid to restore URA-. YEG2 was created from YEG1 by transformation with the 5-TEV-ZZ-URA3 cassette from pKW953, PCR amplified with Vent DNA Polymerase (New England BioLabs, Ipswich, MA) and primers 5′-GGCAGAAAGATCATGAAACTATGCAAGAAATTGATAGCAAGAGCGGTATCGACG GATCCCCGGG-3′ and 5′-CTTACTTATTGAGATGTACAGTGGAAATAGGAGCTTTCTGAATTC-3′ and served both hyperphosphorylation and CN-mediated dephosphorylation of the Inp53-PRD during hypertonic shock, suggesting that coordination between CN and antagonistic kinases may be required to temporally or spatially regulate the activity of Inp53. Thus the parallel recruitment/interaction with the homologous yeast proteins Inp53 (synaptojanin), Rvs167 (amphiphysin/endophilin), Bzz1 (syndapin), Sia1 (intersectin) and CN during hypertonic stress suggest that this yeast response is strikingly similar to ADBE at the molecular level.

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FIGURE 9: Molecular mechanism of Inp53 regulation by calcineurin. Wild-type cells (A) or cells expressing the indicated GFP fusion from endogenous loci (B–E) and expressing GST-Inp53, GST-Inp53 AraQAA, or GST were treated with or without 1.25 M KCl for 10 min, and GST-tagged proteins were purified with glutathione-Sepharose. (C) Soluble (Input) and insoluble fractions (20,000 × g pellet). Copurification of the GFP fusion proteins with indicated GST-fusions was assessed by Western blotting; representative blots from more than three independent experiments are shown.
FIGURE 10: Model. (A) CN is inactive in unstressed cells, and in response to many stress conditions, remains cytoplasmic and is activated by Ca\(^{2+}\)/calmodulin, promoting dephosphorylation of Crz1 and resulting in gene expression. In response to hyperosmotic shock, CN relocation promotes activation of Inp53 but blocks Crz1 dependent transcription. (B) In unstressed cells, constitutive endocytosis is polarized and occurs with the involvement of clathrin, Slal, Inp53, Rvs167, Bz1, Bsp1, and many other proteins not shown; Inp53 associates with clathrin and Rvs167 and functions primarily at the TGN. During hyperosmotic shock, the composition of actin patches changes, with Inp53 losing interactions with clathrin and gaining interactions with Slal, Bz1, Bsp1, and CN; Inp53 also becomes hyperphosphorylated, and CN dephosphorylates it and Slal. (C) Many endocytic proteins and their interactions are conserved in mammals, where CN promotes association between synaptopgin, dynamin, amphiphysin, syndapin, and intersectin during synaptic vesicle endocytosis.

Plasmid construction

A C-terminal fragment of CNA1 was amplified with primers 5′-CCCGCTCGAGCAGTGCGCCAAACTACCTGGAC-3′ and 5′-CGCGGATCCTCCAGTTGTGGCTTTTTCTCCGCTCCC-3′ and inserted as a Xhol/BamHI fragment into pB1963 to create pCNA1-3GFP. A C-terminal fragment of CNA2 was amplified with primers 5′-CCCGCTCGAGGGGAGATTCCCTCTCATGGCCTG-3′ and 5′-CGCGGATCCTCTTTGCTATCATTCTTTGCATCATGTT-3′ and inserted as a Xhol/BamHI fragment into pB1963 to create pCNA2-3GFP.

Inp53 was amplified with primers 5′-GCTCTAGACATGATTATCTTTGTTTCAGAAGAACCTGAAAG-3′ and 5′-ACGCGTCGACTCATTTTGGGGTCAATGGCTGCC-3′ from genomic DNA and ligated into frame into pEG(KT) as an XbaI/SalI fragment to create pEG6(GST-Inp53). Inp53 SAC domain was amplified with primers 5′-GCTCTAGACATGATTATCTTTGTTTCAGAAGAACCTGAAAG-3′ and 5′-ACGCGTCGACTCATTTTGGGGTCAATGGCTGCC-3′ and ligated in-frame into pEG(KT) as an XbaI/SalI fragment to create pEG7. The Inp53 IP5P domain was amplified with primers 5′-GCTCTAGACAGGTTACCGTATCAGAAAGCAG-3′ and 5′-ACGCGTCGACTCATTCTTCGGGATGCTCTTGTTTGTATTC-3′ and ligated in-frame into pEG(KT) as an XbaI/SalI fragment to create pEG8. The Inp53 PRD domain was amplified with primers 5′-GCTCTAGACCCGGGTTCTCTGATATCGGATTCTTC-3′ and 5′-ACGCGTCGACTCATTTTGGGGTCAATGGCTGCC-3′ and ligated in-frame into pEG(KT) as an XbaI/SalI fragment to create pEG9.

pEG12 was created using PCR mutagenesis from pGST-Inp53 with internal primers 5′-GGGAGCAAGGTACCTCTTTAATCAATGCTAGAGCACAAGCAGCAGATCATTTGAAGCCACCCAACCG-3′ and 5′-CGGTTGGGTGGCTTCAAATGATCTGGCTGCTTGTGCTCTAGCATTTTGGGGTCAATGGCTGCC-3′ and inserted into pEG(KT) as an XbaI/SalI fragment to create pEG7. The Inp53 IP5P domain was amplified with primers 5′-GCTCTAGACAGGTTACCGTATCAGAAAGCAG-3′ and 5′-ACGCGTCGACTCATTCTTCGGGATGCTCTTGTTTGTATTC-3′ and ligated in-frame into pEG(KT) as an XbaI/SalI fragment to create pEG7.

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pEG12 was created using PCR mutagenesis from pGST-Inp53 with internal primers 5′-GGGAGCAAGGTACCTCTTTAATCAATGCTAGAGCACAAGCAGCAGATCATTTGAAGCCACCCAACCG-3′ and 5′-CGGTTGGGTGGCTTCAAATGATCTGGCTGCTTGTGCTCTAGCATTTTGGGGTCAATGGCTGCC-3′ and ligated in-frame into pEG(KT) as an XbaI/SalI fragment to create pEG9.

pEG12 was then cut with EagI/Sall, and EagI/BamHI and BamHI/Sall fragments of Inp53(IP5P) and Inp53(PR5D) amplified from pEG6 template with

BY4741 with Nde1-linearized pEG4 and with Xbal-linearized pEG38 or Bsu36I-linearized pEG27 or pEG28, respectively.

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primers 5′-GCAGATCTCTCGGCGGGTACCGTACGAAAGCA
GTGCAA-3′ and 5′-AAGGCTGATCCAGTGCCTGTTCGG
GGATGCTTCTTTTGTATTCG-3′, and primers 5′-GCAGATCTC
CAGGAACCGGATACATGGCTCCTGGACGCC-3′ and 5′-AGCCGTC
GACTCATTTTGGGTTAGTCTGGCC-3′, respectively, were in
serted to create pEG11 (GST-Inp53ΔC). A BsaBI/HindIII fragment
from pEG11 (GST-Inp53ΔC) was subcloned into pRS426-Inp53
and pRS316-Inp53 to create prS426-Inp53ΔC and pRS316-Inp53ΔC,
respectively. pRS426-GFP-2xPhI (PLC8) was converted from URA3
to ADE2 by gap repair, yielding pEG30. mKate was amplified (Shercher
et al., 2007) with primers 5′-GGTTGAACTTCACTGATCCATACATCG
GTTACTGATGTTAGG-3′, 5′-CGTAAAG
GCTTTCTTGCCAGACCTAGTAG-3′, 5′-CGTAAAG
GCTTTCTTGCCAGACCTAGTAG-3′, and 5′-GGTTGAACTTCACTGATCCATACATCG
GTTACTGATGTTAGG-3′ inserted into prS303 as a EcoRI-ApaI fragment. mKate was
amplified with 5′-CGTTTGAATTCGTCACTTTTGGGGTCAATGGCTG
GTTACTGATGTTAGG-3′, 5′-CGTAAAG
GCTTTCTTGCCAGACCTAGTAG-3′, and 5′-GGTTTGAATTC
GTTACTGATGTTAGG-3′ amplified with 5′-CGTTTGA
ATTCGTCACTTTTGGGGTCAATGGCTG
GTTACTGATGTTAGG-3′, 5′-CGTAAAG
GCTTTCTTGCCAGACCTAGTAG-3′, and 5′-GGTTTGAATTC
GTTACTGATGTTAGG-3′ and inserted as EcoRI-ApaI fragment, HindIII,
XhoI, and Xhol-Apal fragments into prS303. Inp53 and Inp53ARA
AAGA C-terminal fragments (amino acids 203–1107) were ampli
cified with 5′-GTCTAGctgtagATGTAAGGTGTTTGTG
GTTACTGATGTTAGG-3′ and 5′-GGCTGACATCGTGTTT
GTTACTGATGTTAGG-3′ and inserted as an XbaI-SphI fragment into prS303-3xmKate. Apb1 C-term was ampli
cified with 5′-GGTGTAGAGCTCAGGATTGGCCGCTTCAGAAAA
AGGAGG-3′ and 5′-CCACATAGTTCGTGTGCCCAAAACATACAT
TGCTGGG-3′ and inserted as a SacI-EcoRI fragment into prS303-
mKate. All plasmids were confirmed by sequence analysis.

Fluorescence microscopy and analysis
Cell imaging was performed at 25°C with a Zeiss Axio Imager M1
microscope (Carl Zeiss, Jena, Germany) with a mercury arc lamp and
a 100x/1.3 numerical aperture oil immersion objective, and images
were captured with an Orca-ER digital camera (Hamamatsu, Bridge
water, NJ) coupled to Openlab Software 5.0.1 (PerkinElmer-Cetus,
Waltham, MA). A 475/40 excitation and 530/50 emission filter set
was used for GFP and fluorescence isothiocyanate (FITC)–concanava
lin A (Sigma-Aldrich, St. Louis, MO) images, and a 560/40 excitation
and 630/75 emission filter set was used for RFP and rhodamine-phalloidin
(Invtrogen, Carlsbad, CA) images.

For calcineurin localization studies, Cna1-3GFP was grown to
mid log phase in synthetic complete dextrose (SCD), and hyperos
totic stress was initiated by adding NaCl, KCl, sorbitol, CaCl
2, or LiCl. Cells were concentrated by centrifugation and imaged. Cells
were identified from fluorescence images by Otsu’s threshold
method. Pearson’s r was determined for each GFP-RFP pair from all
analysis, one-phase decay was fitted to each time series with Prism
(GraphPad, La Jolla, CA) and the two fits were compared with the
extra sum-of-squares F test.

For the latrunculin A experiments, Cna1-3GFP cells were grown
to mid log phase in SCD, synchronized with 15 μg/ml α-factor for 2
h, and then washed and released into SCD for 40 min. Latrunculin A
(Sigma-Aldrich) stock solution was 20 mM in dimethyl sulfoxide
(DMSO); cells were treated with 200 μM latrunculin A or 1% DMSO
for 10 min, and then an equal volume of SCD with 2.5 M KCl and
200 μM latrunculin A or DMSO was added for an additional 10 min,
att which point cells were imaged or fixed with 3.7% formaldehyde
and stained with rhodamine–phalloidin (Invtrogen). Quantification of
Cna1-3GFP from unfixed cells, N > 100, was conducted using ImageJ (National Institutes of Health, Bethesda MD): pixel intensity
along a line parallel to the mother–bud axis and centered at the bud
neck was averaged.

Actin polarization and GFP-2xPhI (PLC8) were measured in
BY4741 and LSY312 (Inp53ΔA) cells. BY4741 cells were grown to mid
log phase, and LSY312 (Inp53ΔA) were grown to saturation in selec
tive media with 2% galactose and then washed and grown in 2%
dextrose for at least 10 doublings. Cells were pretreated for
between 30 min and 2 h with or without FK506, as indicated, and
hyperosmotic stress was initiated by adding KCl to 1.25 M.

For actin polarization measurements, cells were fixed with 3.7%
formaldehyde buffered with 100 mM KPI, pH 6.5, and stained with
FITC-concanavalin A and rhodamine–phalloidin. Images were ana
lyzed with CalMorph (http://scmd.gi.k.u-tokyo.ac.jp/datamine/
calmorph/; Ohtani et al., 2004), and actin polarity was calculated for small-
and medium-budded cells as the ratio of actin brightness in the
bud to actin brightness in the mother cell. Differences between
distribution of actin polarity were evaluated for significance with
the Kruskal–Wallis rank sum test for multiple comparisons using the
MATLAB statistics tool pack (MathWorks, Natick, MA).

GFP-2xPhI (PLC8) invaginations (Figure 6) were measured with a
custom GFP membrane identification algorithm implemented in MAT
LAB; briefly, we used a Laplacian of Gaussian kernel to identify con
tours defining the inner and outer borders of the cell membrane and
restricted the analysis to well-defined cells. Number of large invagi
nations was defined as the number of membrane regions that identi
fied extended >10 pixels (~0.5 μm) from the outer edge of the cell.

LSY312 (Inp53ΔA) cells with pRS426-Inp53, pRS316, pRS316-Inp53,
and pRS316-Inp53ARAAGA were treated with or without KCl for 10 min
before imaging; >200 cells from each condition from three indepen
dent experiments were analyzed.

GFP-2xPhI (PLC8) enrichment/depletion (Figure 7) in wild-type
(BY4741) cells, pretreated with FK506 or vehicle for 30 min with or
without osmotic shock, was analyzed using a custom MATLAB script;
we identified cell bodies and defined the outside five pixels as the
membrane region; GFP enrichment relative to the adjacent cyto
plasm was calculated in a sliding window along the membrane: a
location on the membrane was scored as GFP enriched if the aver
age fluorescence in the surrounding 10 × 5 pixel membrane region
was >10% more than the average fluorescence in a similarly sized
adjoining region of cytoplasm after image-wide background correc
tion. Details and code are available on request. Statistical testing
was with one-way analysis of variance (ANOVA) and Tukey’s multiple
comparison posttest (Prism), and >700 cells were analyzed from
each condition from four independent experiments.

Analysis of correlation between Cna1-3GFP, Apb1-RFP, Inp53-3
RF, and Inp53ARAAGA-3RF was conducted as follows. Cell bodies
were identified from fluorescence images by Otso’s threshold
method. Pearson’s r was determined for each GFP-RFP pair from all
pixels in cell bodies. To calculate statistical significance, correlation
scores from >10 images from at least two independent experiments
were analyzed by one-way ANOVA with Tukey’s posttest in Prism.

Mass spectrometry
Stable isotope labeling by amino acids in cell culture triple label
ing was conducted following Gruler and Kratchmarova (2008); cells
were grown in minimal media supplemented with 30 mg/l in 1.3
1-cultures. YEG1 was grown with Lys3-1-l-lysine, and YEG2 was
grown with Lys14 or Lys18 (4,4,5,5-D4-l-lysine or U-13C6, U-15N2
l-lysine; Cambridge Isotope Laboratories, Andover MA). Cells
were grown to OD of 0.8; 121 g of KCl was added to the Lys3/
YEG2 culture (1.25 M) for 10 min. Cells were washed, equalized
d by OD600, and pooled, then resuspended in lysis buffer (100 mM

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NaCl, 20 mM Tris, pH 7.6, 2 mM EDTA, 0.5% Triton, 1 mM dithiothreitol (DTT), and protease inhibitors (phenylmethylsulfonyl fluoride, aprotinin, leupeptin, pepstatin, benzamidine), and droplets were frozen directly in liquid N2. Frozen cell droplets were lysed with a Retsch MM301 Ball-Mill (5 × 2 min, 20 Hz, N2(I) cooled), and then extracts were clarified with 10,000, 50,000, and 100,000 × g sp. Cna2-S-TEV-2Z and associated proteins were purified with immunoglobulin G (IgG)–Sepharose and eluted by TEV cleavage at 4°C (reaction buffer: 50 mM Tris, pH 8, 0.5 mM EDTA, 1 mM DTT, 50 U AcTeV [Invitrogen]; elution buffer: 250 mM NaCl, 20 mM Tris, pH 7.6, 2 mM EDTA, 0.5% Triton, 1 mM DTT). Eluted proteins were trichloroacetic acid (TCA) precipitated (15% TCA, −20°C overnight; washed three times with acetone), digested with LysC (50 mM ammonium bicarbonate, 10% acetonitrile, 20 ng/μl LysC, 37°C overnight; stop with 10% trifluoroacetic acid), and cleaned with a C18 StageTip as described (Rappsilber et al., 2007; wash, 5% formic acid; elute, 50% acetonitrile, 5% formic acid). Samples were analyzed by liquid chromatography–tandem mass spectrometry (LTQ-Velos Orbitrap; Thermo Scientific, Waltham, MA). Peptides were identified with the SEQUEST algorithm (Yates et al., 1995), against the yeast proteome concatenated with reversed sequence and quantified with the VISTA algorithm (Bakalarski et al., 2008).

**Yeast protein purification and Western blotting**

Copurification and Western blot studies with GST-tagged Inp53, Inp5334ACAA, and Inp53-SAC and IP5P and PRD domains were conducted in JRY11, BY4741, and Rvs167-GFP, Bsp1-GFP, Bzz1-GFP, and Sla1-GFP cells transformed with the appropriate plasmids. Cells were grown in synthetic complete (SC)–Ura 2% raffinose medium to mid log phase and treated with 1 μM FK506 for 30 min where indicated and then induced with 2% galactose for 4 h. A 10-min hyperosmotic stress was induced by adding KCl to 1.25 M where indicated. Cells were harvested and washed in prechilled medium containing FK506 and 1.25 M KCl where appropriate and then pellet and flash frozen in liquid N2. Cells were resuspended in copurification buffer (150 mM NaCl, 20 mM Tris 7.6, 2 mM EDTA, 0.5% Triton X-100, 1 mM DTT, and protease inhibitors, with or without FK506 as appropriate) and were lysed with glass bead lysis, and extracts were clarified with 5000 and 20,000 × g sp.; extracts were incubated with glutathione-Sepharose beads (GE Healthcare, Little Chalfont, United Kingdom) at 4°C for 2 h with end-over-end rotation and washed three times for 5 min in copurification buffer. Samples were boiled in 2× SDS loading buffer for 10 min at 95°C, resolved with SDS–PAGE using standard methods, and transferred to nitrocellulose membranes (Bio-Rad, Hercules, CA) for Western blotting. Membranes were blocked with SuperBlock (Thermo Scientific, Rockford, IL) and probed with rabbit polyclonal anti-GFP IgG fraction (Invitrogen), anti-5′ (a gift from Ron Kopito, Stanford University), anti-Chc1 (SKL-1, a gift from Greg Payne, University of California, Los Angeles), anti-Inp53 (a gift from John York, Vanderbilt University), anti-phosphoTyr7/Y (Invitrogen, Beverly, MA), or mouse monoclonal anti-GST (Covance, Berkeley, CA) antibodies. Anti-rabbit or anti-mouse secondary antibodies conjugated to horseradish peroxidase (GE Healthcare) and SuperSignal West Dura Extended Duration Substrate (Thermo Scientific, Rockford, IL) or ECL Western Blotting Substrate (Pierce, Rockford, IL) were used to visualize proteins on Biomax XAR film (Eastman Kodak, Rochester, NY), except in Supplemental Figures S3 and S4, where anti-rabbit Alexa 780 and anti-mouse Alexa 680 secondary antibodies (Invitrogen) were visualized with an Odyssey scanner (LI-COR Biosciences, Lincoln, NE).

**In vitro phosphatidylinositol phosphatase assays**

For purification of Inp53 for in vitro assays, we transformed JRY11 with pEG6 (GST-Inp53wt) and grew cells to mid log phase in SC-Ura 2% raffinose medium and treated with or without 1 μg/ml FK506 for 30 min and then induced with 2% galactose for 4 h. A 10-min hyperosmotic stress was induced by adding KCl to 1.25 M. A 500-ml amount of cells was harvested and washed in prechilled medium containing FK506 and 1.25 M KCl where appropriate, then resuspended in lysis buffer (150 mM NaCl, 20 mM Tris, pH 7.6, 2 mM EDTA, 1 mM DTT, 5 mM Na pyrophosphate, protease inhibitors). To preserve SAC domain activity (with or without FK506), no detergent was used and droplets were frozen directly in liquid N2. Frozen cell droplets were lysed with a Retsch MM301 Ball-Mill (5 × 2 min, 20 Hz, N2(I) cooled), and then extracts were clarified at 5000 and 20,000 × g. GST-Inp53 was purified with glutathione-Sepharose beads (GE Healthcare) for 2 h at 4°C with end-over-end rotation and washed three times for 5 min, then eluted with 150 mM NaCl, 50 mM Tris, pH 7.6, 40 mM glutathione, and 1 mM DTT and dialyzed with 50 mM Tris, pH 7.6, 1 mM DTT, and frozen with 15% glycerol at −80°C. Proteins were analyzed by SDS–PAGE and Coomassie stained to assess purity or transferrable to polyvinylidenyl fluoride membrane and Western blotted with mouse monoclonal anti-GST primary (Covance, Emeryville, CA) and anti-mouse Alexa 680 secondary antibodies (Invitrogen); immunoblots were observed with an Odyssey scanner, and relative protein amounts were calculated from serial dilutions using ImageJ (unpublished data). Equal amounts of each enzyme were used (approximately 10 nM) for each reaction. Phosphatidylinositol phosphatase activity was assayed in 50 mM 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid, pH 7.4, 50 mM KCl, 3 mM ethylene glycol tetraacetic acid, and 2 mM MgCl2, with 0–250 μM di-C8-phosphatidylinositol(4,5)bis-phosphate (Echelon Biosciences, Salt Lake City UT); phosphate release was measured with Biomol Green (Enzo Life Sciences, Farmingdale, NY); rates were calculated from at least four linear time points and fitted with an allosteric-sigmoidal model and tested for significance with an extra sum-of-squares F test (p < 0.13, one curve adequately explained all data).

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