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Dephosphorylation of Iqg1 by Cdc14 regulates cytokinesis in budding yeast

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ABSTRACT

Cytokinesis separates cells by contraction of a ring composed of filamentous actin (F-actin) and type II myosin. Iqg1, an IQGAP family member, is an essential protein in Saccharomyces cerevisiae required for assembly and contraction of the actomyosin ring. Localization of F-actin to the ring occurs only after anaphase and is mediated by the calponin homology domain (CHD) of Iqg1, but the regulatory mechanisms that temporally restrict actin ring assembly are not well defined. We tested the hypothesis that dephosphorylation of four perfect cyclin-dependent kinase (Cdk) sites flanking the CHD promotes actin ring formation, using site-specific alanine mutants. Cells expressing the nonphosphorylatable iqg1-4A allele formed actin rings before anaphase and exhibited defects in myosin contraction and cytokinesis. The Cdc14 phosphatase is required for normal cytokinesis and acts on specific Cdk phosphorylation sites. Overexpression of Cdc14 resulted in premature actin ring assembly, whereas inhibition of Cdc14 function prevented actin ring formation. Cdc14 associated with Iqg1, dependent on several CHD-flanking Cdk sites, and efficiently dephosphorylated these sites in vitro. Of importance, the iqg1-4A mutant rescued the inability of cdc14-1 cells to form actin rings. Our data support a model in which dephosphorylation of Cdk sites around the Iqg1 CHD by Cdc14 is both necessary and sufficient to promote actin ring formation. Temporal control of actin ring assembly by Cdk and Cdc14 may help to ensure that cytokinesis onset occurs after nuclear division is complete.

INTRODUCTION

Cytokinesis, the final step in cell division, divides the cytoplasm between two daughter cells. Precise temporal control is necessary to coordinate cytokinesis and mitosis so that proper chromosome segregation can be completed. Cytokinetic failure results in tetraploid cells, and there is evidence that tetraploidy is an intermediate state leading to chromosomal instability, aneuploidy, and tumorigenesis (Ganem et al., 2007; Storchova and Kuffer, 2008). In animal and fungal cells, cytokinesis is achieved by a ring composed of filamentous actin (F-actin) and nonmuscle type II myosin (Satterwhite and Pollard, 1992; Fishkind and Wang, 1995). The actomyosin ring is a transient structure that is precisely positioned to bisect the elongating anaphase spindle, ensuring proper chromosome segregation (Pollard, 2010). In yeast, actomyosin ring contraction must be coordinated with septation, the process of adding new cell wall material between the dividing cells (Bi, 2001).

In budding yeast, both actomyosin ring assembly and contraction are cell cycle regulated. The type II myosin heavy chain, Myo1, forms a ring at the bud neck in G1 (Bi et al., 1998; Lippincott and Li, 1998b). Localization of F-actin and contraction of the ring occur after anaphase and require the essential protein Iqg1/Cyk1 (Epp and Chant, 1997; Lippincott and Li, 1998b; Lippincott et al., 2001). Iqg1 is a 173-kDa scaffolding protein homologous to mammalian IQGAPs. IQGAP family members are essential for actin-based processes such as phagocytosis, cell adhesion, migration, and cytokinesis (Shannon, 2012; White et al., 2012). Iqg1 shares with other IQGAPs multiple functional domains: an N-terminal calponin homology domain (CHD), IQ repeats, a GTPase-related domain (GRD), and a Ras-GAP C-terminus (RGCt; Epp and Chant, 1997; Lippincott and Li, 1998b; Lippincott et al., 2001).
the myosin regulatory light chain, Mlc1, via interactions with the IQ domains (Boyme et al., 2000; Shannon and Li, 2000). Iqg1 recruitment to the ring is restricted to mitosis even though Mlc1 localizes to the bud neck earlier in the cell cycle (Boyme et al., 2000; Shannon and Li, 2000). Iqg1 binds F-actin via the CHD and recruits F-actin to the bud neck during anaphase/telophase of mitosis, which completes actomyosin ring assembly (Shannon and Li, 1999). The C-terminus of Iqg1, containing both the GRD and RGCl domains, interacts with the GTPase Tem1 and is required for contraction of the actomyosin ring (Shannon and Li, 1999). Tem1 is part of the mitotic exit network (MEN), and temperature-sensitive alleles of Tem1 and other MEN proteins cause mitotic arrest in late anaphase before cytokinesis (Jaspersen et al., 1998; McComb and Gould, 2001).

The MEN, a signaling cascade that leads to the full release and activity of the phosphatase Cdc14, regulates mitotic exit, cytokinesis, and septation in budding yeast (Bardin and Amon, 2001; Meitinger et al., 2012). The MEN brings about exit from mitosis by inactivating the mitotic cyclin-dependent kinase 1 (Cdk1). Cdk1, or Cdc28-Clb2 in budding yeast, is inactivated durung anaphase via two partially redundant methods: inhibition of its kinase activity by Sic1 and degradation of the mitotic cyclin Clb2 by the APC^cdcl complex (Jaspersen et al., 1998; Visintin et al., 1998). Both Sic1 and Cdk1 become active once dephosphorylated by the MEN component Cdc14, leading to inhibition of Cdk1 activity (Visintin et al., 1998). When Cdk1 activity drops, MEN proteins Cdc15, Dbf2-Mob1, Dbf20-Mob1, and Cdc14 all accumulate to the bud neck by an unknown mechanism, where they are positioned to regulate cytokinesis and septation (Bembenek et al., 2005; Meitinger et al., 2012). Although Cdk1 must be inactivated in order for actomyosin ring assembly and contraction to occur, the MEN has a role in promoting cytokinesis in addition to Cdk1 inactivation (Meitinger et al., 2012; Sanchez-Diaz et al., 2012). Overexpression of Sic1 bypasses mitotic arrest in most MEN mutants, but cytokinesis defects persist. In cells expressing temperature-sensitive alleles of the MEN gene MOB1 and overexpressing Sic1, the actomyosin ring could assemble but not contract (Luca et al., 2001). Similarly, inducing Cdc14 release after Tem1 depletion led to formation of actin rings that did not contract (Lippincott et al., 2001), suggesting that the MEN regulates the timing of cytokinesis onset. Cdc14 can dephosphorylate the actin filament-nucleating formins Bni1 and Bnr1, which may affect the actin filament–nucleating formins Bni1 and Bnr1, which may affect the 

RESULTS

Effects of Iqg1 phosphorylation mutations on cytokinesis

To test the hypothesis that the phosphorylation state of Iqg1 during the cell cycle is important for regulation of cytokinesis, we constructed two plasmids containing mutations of the four perfect consensus Cdk sites (S/TPxR/K, where x is any amino acid; Li et al., 2008) that flank the CHD (Figure 1A). Using site-directed PCR mutagenesis, we mutated each serine or threonine to either alanine or glutamic acid. Two alleles were generated—iqg1-4A, with all four perfect Cdk sites mutated to alanine to prevent phosphorylation, and iqg1-4E, with all four amino acids mutated to glutamic acid to mimic phosphorylation. Both alleles were expressed using the endogenous IQG1 promoter and tagged at the 3′ end with 13 copies of the myc epitope. Because it was uncertain whether cells expressing only the mutant alleles would be viable, each plasmid was introduced into a yeast strain that contains the wild-type copy of IQG1 under the inducible GAL1 promoter. This allowed the cells to be grown while expressing the wild-type copy of IQG1 and for the wild-type copy to be repressed in order to see the effects of the mutations expressed using IQG1 native promoter. As we have previously shown, the GAL1-IQG1 is repressed after growth in yeast extract/peptone/dextrose (YPD) and phenocopies the null IQG1 allele (Figure 1C, lane 3; Lippincott and Li, 1998b; Shannon and Li, 1999). Both mutant proteins were expressed at levels comparable to a similarly tagged wild-type Iqg1 protein (Figure 1C).

Cytokinesis defects in budding yeast cause a distinct phenotype in which cells continue to divide and rebud despite failing to separate, producing chains of cells (Figure 1D). To determine whether mutation of the Cdk phosphorylation sites affected cytokinesis, we examined the morphology of cells expressing the iqg1-4A and
and either wild-type expressed under the or phosphorylation mutant IQG1 GAL1-IQG1, IQG1 GAL1-IQG1, three strains—wild-type copy of under the inducible promoter. The _iqg1-4E_ alleles. For comparison, the wild-type copy of IQG1 under the _GAL1_ promoter was also introduced into the strain with the _IQG1_ copy of IQG1 under the _GAL1_ promoter and cells expressing both wild-type _IQG1_ and mutant alleles affecting IQG1 expression from the _GAL1_ promoter did not differ significantly from control cells in YPGR, indicating that _iqg1-4A_ does not have a dominant-negative effect (Figure 1E). The cells expressing _iqg1-4A_ exhibited altered morphology when wild-type _GAL1-IQG1_ was repressed in YPD, forming chains in 44% of cells, significantly different both from controls expressing wild-type _IQG1_ from the endogenous promoter and cells expressing both _iqg1-4A_ and wild-type _IQG1_ from the _GAL1_ promoter (p = 0.0002 and 0.018; Figure 1E). Because yeast cells have a cell wall, it is possible that this chain phenotype is caused by defects in either septation or cytokinesis. To distinguish between the two possibilities, we used Zymolyase to remove the cell wall (Lippincott and Li, 1998b). After Zymolyase treatment, cells expressing only _iqg1-4A_ still exhibited chains with significantly higher frequency than controls (32% compared with 9.5% in _IQG1_ alone, p = 0.002, and 14% in _iqg1-4A_ _GAL1-IQG1_—expressing cells, p = 0.02; Figure 1E).

In the strain expressing _iqg1-4E_, defects in cytokinesis were high both in the presence and absence of wild-type IQG1 expressed from the _GAL1_ promoter and with and without Zymolyase (Figure 1, D and E). In all cases, the percentage of chains in _iqg1-4E_ cells was significantly higher than in control _IQG1_ cells under the same conditions (p ≤ 0.01). In addition to the high number of chains, the cell bodies in the chains were often elongated and appeared to be hyperpolarized (Figure 1D, right). _iqg1-4E_, but not _iqg1-4A_, caused slower growth and appeared hyperpolarized (Figure 1D, right). _iqg1-4E_, but not _iqg1-4A_, caused slower growth and appeared hyperpolarized (Figure 1D, right). _iqg1-4E_, but not _iqg1-4A_, caused slower growth and appeared hyperpolarized (Figure 1D, right). The reason for this is unknown, but it is possible that _iqg1-4E_ is acting as a dominant negative, and therefore we did not further investigate the _iqg1-4E_ allele.

Our results show that the _iqg1-4A_ and _iqg1-4E_ alleles affecting phosphorylation of Iqg1 cause a chain phenotype significantly above that of _IQG1_ control cells. Because the _IQG1_ mutant alleles...
were expressed from a plasmid rather than the endogenous locus, we integrated a copy of iqg1-4A into the chromosome as described in Materials and Methods. The cells with integrated iqg1-4A had a significantly higher percentage of chains than the parental strain (11% in control cells compared with 24% in iqg1-4A; \( p = 0.01 \); Supplemental Figure S1). Our results are similar to those found in Candida albicans, demonstrating that the regulation of cytokinesis by phosphorylation of IQG1 is conserved between these yeast species. These data demonstrate that mutation of the four perfect Cdk sites in lqg1 compromises the cell's ability to complete cytokinesis.

**Mutations that prevent phosphorylation of the iqg1 Cdk sites cause actin ring formation before anaphase**

In the yeast *C. albicans*, mutation of 15 minimal Cdk1 consensus sites in lqg1 to alanine resulted in premature assembly of the actomyosin ring and cytokinesis defects (Li et al., 2008). We also recently reported that mutation of 14 Cdk sites in budding yeast iqg1 caused actin ring formation before anaphase (Naylor and Morgan, 2014). We tested specifically whether the four perfect Cdk sites in lqg1 flanking the actin-binding CHD domain were important for controlling the timing of actin recruitment to the ring. To do this, we examined the timing of actin ring formation in synchronous cell cultures. Assembly of the actomyosin ring is regulated during the cell cycle, with lqg1 and F-actin localizing to the bud neck during anaphase (Lippincott and Li, 1998; Lippincott et al., 2001). Cells were arrested in G1 using mating factor. After release from the G1 arrest, a time course was performed with a sample of cells fixed every 20 min up to 100 min. The cells were stained using antibodies, phalloidin, and 4',6-diamidino-2-phenylindole (DAPI) to visualize lqg1, F-actin, and DNA, respectively. Images of >100 cells were analyzed at each time point under each condition in triplicate for the presence of actin and lqg1 rings (Figure 2). Experimental cultures were cultured in YPD to represses wild-type GAL1-IQG1, leaving iqg1-4A as the sole source of lqg1 (Shannon and Li, 1999). Cells expressing only iqg1-4A were compared with cells expressing iqg1-4A and GAL1-IQG1 (the same strain grown in YPGR), as well as to control cells with IQG1 under the endogenous promoter in YPD.

In cells expressing only iqg1-4A (iqg1-4A GAL1-IQG1 cells grown in YPD, henceforth referred to as iqg1-4A cells), formation of the actin ring occurred 20 min earlier in the time course than in both sets of control cells (Figure 2A). Approximately 13% of iqg1-4A cells had both an lqg1 ring and an actin ring 40 min after release from arrest, whereas the same cells in YPGR had <1% at this time point, and cells expressing only IQG1 had no rings at 40 min (\( p = 9.2 \times 10^{-6} \) and 4.5 \( \times 10^{-4} \); Figure 2, A and B). The iqg1-4A cells that formed lqg1 and actin rings at 40 min had not yet entered anaphase, indicated by the single DNA mass (Figure 2B). It is noteworthy that, contrary to a previous report, overexpression of wild-type GAL1-IQG1 in YPGR control cells did not induce premature ring formation (Epp and Chant, 1997). At the 60-min time point, iqg1-4A and control cells contained actin rings. Control cells with both an lqg1 ring and an actin ring either contained two DNA masses (Figure 2C) or an elongated mass of DNA spanning the bud neck, indicating that the cell is in anaphase. In iqg1-4A cells that had formed lqg1 and actin rings at 60 min, some cells had yet to go through anaphase, since they had only a single nucleus (Figure 2, C and D). There was a significant increase in rings observed in iqg1-4A cells at the 60-min time point compared with controls, likely due to the fact that rings in iqg1-4A cells could be formed before anaphase, whereas in control cells, only those that had initiated anaphase contained rings (\( p = 0.01 \) compared with YPGR control and \( p = 9.5 \times 10^{-6} \) compared with YPD control; Figure 2A). At the 20-min time point, no lqg1 or actin rings were seen in control or experimental cells. At the 80- and 100-min time points, there was no statistically significant difference in the number of cells with lqg1 and actin rings between the iqg1-4A cells and YPGR controls, but these cells did have a greater amount of rings than YPD controls (Figure 2A). These data show that expression of iqg1-4A at endogenous levels increased ring formation similar to overexpression of IQG1 using the GAL1 promoter, showing that iqg1-4A is constitutively active for actin ring formation. Overexpression of wild-type IQG1 increases the number of rings relative to YPD controls but does not change the timing of ring assembly.

To analyze whether the lqg1/actin rings formed before anaphase onset, we scored the DNA content of >100 iqg1-4A cells and YPDR and YPGR control cells with rings at the 40-, 60-, and 80-min time points to determine whether there was one nucleus, two nuclei, or a nucleus in the process of division. Cells were judged to be in the process of division if the DAPI signal was elongated across the bud neck. Of iqg1 and actin rings formed in control cells, 92–98% had a nucleus in the process of division or completely separated, indicating that actomyosin ring formation occurred after anaphase onset (Figure 2D). In contrast, 55% of iqg1-4A cells with rings had only a single nucleus, indicating that preventing phosphorylation of lqg1 accelerated actin ring formation (Figure 2D). These results are consistent with the finding that wild-type cells form actin rings only when there is an anaphase spindle, whereas lqg1 phosphomutants formed actin rings in cells with both preanaphase and anaphase spindles (Naylor and Morgan, 2014). Our data support the hypothesis that phosphorylation of the four perfect Cdk sites flanking the CHD of lqg1 negatively regulates actin ring formation.

**Changing Cdc14 levels affect actomyosin ring formation**

Cdc14 is the final protein in the MEN signaling cascade and has been shown to dephosphorylate Cdk1 targets (Chin et al., 2012; Visintin et al., 1998). To test the hypothesis that Cdc14 regulates the timing of actin ring formation, we overexpressed CDC14 in metaphase cells and examined formation of the actin ring. For these experiments, we obtained the GAL1-CDC14 construct used by Sanchez-Diaz et al. (2012). Cells were arrested using the microtubule disruptornocodazole, which activates the spindle assembly checkpoint and arrests cells before anaphase. Cells that had been cultured in YPD overnight were resuspended into YPD (control, GAL1-CDC14 repressed) or YPGR (experimental, GAL1-CDC14 expressed) simultaneously with nocodazole (5 \( \mu \)g/ml) and incubated for 2.5 h. Cells were analyzed for actin rings using A568 phalloidin. Cells overexpressing CDC14 formed twice as many actin rings as control cells (\( p = 0.008 \); Figure 3, A and B). The increase in actin rings in nocodazole-arrested cells overexpressing CDC14 indicates that Cdc14 activity can promote premature actin ring formation. Control cells exhibited a higher percentage of actin rings than expected, since wild-type cells do not form actin rings in nocodazole (Naylor and Morgan, 2014), but this may be due to incomplete inhibition of the GAL1 promoter in YPD and elevated Cdc14 protein levels. It was reported that overexpression of Cdc14 improves the efficiency of cytokinesis when Cdk is inactivated before anaphase (Sanchez-Diaz et al., 2012).

If Cdc14 regulates the timing of actin ring formation by reversing inhibitory phosphorylation by Cdk1, then cdc14-1 mutants will be defective for actin ring formation. Although actin ring formation has been seen in many MEN mutants after bypassing mitotic arrest, a careful analysis of actin ring formation in CDC14 mutant cells has not been reported. To determine whether actin ring formation requires Cdc14 function, we used a temperature-sensitive mutant
allele of CDC14, cdc14-1, and bypassed mitotic arrest using a 2μ plasmid containing SIC1. Cells were cultured overnight and synchronized in mitosis using nocodazole for 90 min at room temperature. Next they were incubated at room temperature (control) or 37°C (experimental) to inhibit cdc14-1 activity, for an additional 90 min. Samples were taken during nocodazole arrest, and then the nocodazole was removed, and two time points were taken after release from mitotic arrest. Cells were fixed, stained for actin, and examined by epifluorescence microscopy.

In nocodazole, cdc14-1 cells almost completely failed to form actin rings at the nonpermissive temperature (2% of cells; \( p = 0.001 \); Figure 3C). In the room temperature controls, ~18% of cells formed rings due to the overexpression of SIC1, as previously reported (Sanchez-Diaz et al., 2012). After release from mitotic arrest, actin ring formation in cdc14-1 cells remained significantly higher in room temperature controls than in the nonpermissive experimental conditions at both 10- and 20-min time points (Figure 3C). These data show that Cdc14 activity is important for actin ring formation in addition to its role in inactivating Cdk1.

**FIGURE 2:** Mutations preventing phosphorylation of Iqg1 lead to formation of the actin ring before anaphase. (A) Timing of actin ring formation. Cells were synchronized in G1 using α factor, and time points were taken at 20-min intervals after release from arrest. During α factor arrest, IQG1 GAL1-IQG1 or iqg1-4A GAL1-IQG1 cells were cultured in medium containing galactose (YPGR) to allow expression of wild-type GAL1-IQG1 or glucose (YPD) to repress expression of GAL1-IQG1. Labels in the key show which genes were expressed. Average of three experiments with 100 cells analyzed at each time point. Error bars are SDs, and \( p \) values were calculated using the Student’s \( t \) test in Excel. *\( p \leq 0.01 \)

(B) Examples of cells at 40 min. Actin, Iqg1, and DNA in iqg1-4 GAL1-IQG1 cells grown in YPGR as a control (top) and iqg1-4A GAL1-IQG1 experimental cells grown in YPD (bottom) at the 40-min time point.

(C) Examples of cells with actin and Iqg1 rings at 60 min. Control iqg1-4 GAL1-IQG1 cells in YPGR (top) and iqg1-4A GAL1-IQG1 experimental cells grown in YPD (bottom) at the 60-min time point. For B and C, labels on the left show which genes were expressed. Images are deconvolved single-plane projections of a Z-series. Scale bar, 5 μm.

(D) IQG1 GAL1-IQG1 or iqg1-4 GAL1-IQG1 cells containing actin and Iqg1 rings were examined using DAPI staining to determine whether they contained one nucleus, two nuclei, or a nucleus in the process of dividing. The x-axis is labeled to reflect the genes expressed: IQG1 is IQG1 GAL1-IQG1 cells grown in YPD, iqg1-4A is iqg1-4 GAL1-IQG1 cells grown in YPD, and iqg1-4A GAL1-IQG1 cells grown in YPGR.
The failure of CDC14 mutants to form actin rings can be rescued by expression of iqg1-4A

Because iqg1-4A and overexpression of CDC14 affect the timing of actin ring formation, and cdc14-1 cells overexpressing SIC1 fail to form actin rings at the nonpermissive temperature, Cdc14 might regulate actin ring formation through dephosphorylation of Iqg1 at Cdk sites. To test this hypothesis, we used control cells that contained cdc14-1 and GAL-SIC1ΔNT-myc to bypass mitotic arrest (Chin et al., 2012). Experimental cells combined the cdc14-1 and GAL-SIC1ΔNT-myc alleles with iqg1-4A and a chromosomal deletion of wild-type IQG1, leaving iqg1-4A as the sole source of Iqg1.

Cells were synchronized in mitosis using nocodazole for 90 min at room temperature and then resuspended in YPGR (to overexpress SIC1) with nocodazole and incubated at 37°C to inhibit

**FIGURE 3:** Cdc14 regulates actin ring formation. (A) Overexpression of Cdc14 causes formation of actin rings during mitotic arrest. Cells containing CDC14 under the GAL1 promoter were used to selectively overexpress Cdc14. Cells grown in glucose were pelleted and resuspended in medium with nocodazole containing either glucose (control) or galactose (experimental) for 2.5 h at 30°C and then fixed for imaging. Actin rings were quantified in three replicates of 100 cells/treatment type. Data are the averages with SDs. A p value of 0.008 was obtained using the Student’s t test. (B) Actin and DNA staining during nocodazole arrest. Top, lack of actin ring in control cell (YPD); bottom, actin ring formation in GAL1-CDC14 cells. Arrow indicates actin ring. Scale bar, 5 μm. (C) Effect of inhibiting Cdc14 activity on actin ring formation. Cells with a temperature-sensitive cdc14-1 allele and SIC1 expressed from a 2μ plasmid were diluted into YPD with addition of nocodazole for 90 min to arrest cells in mitosis. Cells were then placed at room temperature or 37°C for 90 min and fixed during and after release from nocodazole arrest. Noc, cells in nocodazole; 10 and 20 min are minutes after release from nocodazole arrest. Actin rings were quantified in three replicates of 100 cells/treatment type. Data are the averages with SDs. *p ≤ 0.05.
contained actin rings (p = 0.0003 and 0.001; Figure 4A). The cdc14-1 GAL-SIC1ΔNT-myc iqg1-4A cells 10 min after release from metaphase arrest typically had a single DNA mass, indicating preanaphase cells (Figure 4B), whereas 20 min after release, some cells had segregated DNA (Figure 4C). The iqg1-4A mutant rescues the cdc14-1 mutant allele’s inability to assemble an actin ring. These results suggest that lqg1 is the primary Cdc14 target involved in temporal regulation of actin ring formation.

Iqg1 and Cdc14 interact in vivo, and lqg1 is a Cdc14 substrate in vitro

Because our experiments suggested that lqg1 is the target of Cdc14 that regulates actin ring formation, we looked for an interaction between lqg1 and a substrate trap Cdc14 mutant in yeast extracts by coimmunoprecipitation (coIP). Wild-type lqg1 copurified with the Cdc14 substrate-trap protein (Figure 5A). However, an lqg1 mutant (lqg1-3T) in which the Cdk phosphorylation sites at positions 7, 354, and 404 were changed from serines to threonines did not copurify with the Cdc14 substrate trap. Cdc14 is highly selective for phosphoserines within consensus Cdk sequences (Bremmer et al., 2012). This result suggests that Cdc14 recognizes lqg1 as a substrate and that recognition is dependent on just three N-terminal Cdk sites that flank the CHD and contain the optimal Cdc14 recognition motif.

We also directly tested whether the N-terminal Cdk sites flanking the CHD could be efficiently dephosphorylated by Cdc14 in vitro, using phosphopeptide substrates. Cdc14 specificity can be effectively evaluated using phosphopeptides, and catalytic efficiency of Cdc14 toward phosphopeptides containing different Cdk phosphorylation-site sequences can vary by several orders of magnitude (Bremmer et al., 2012).

As expected, phosphorylated Ser-354 and Ser-404 peptides were very efficient substrates, comparable to a previously characterized optimal site in the Acm1 protein (Figure 5B). The peptide containing phosphorylated Thr-299 exhibited essentially no activity, consistent with Cdc14’s selectivity for serine Cdk sites. Two additional Ser Cdk sites that lack the optimal Lys/Arg at position +3, pS365 and pS1347, were used as controls and were dephosphorylated at lower rates, as expected. Surprisingly, phosphorylated Ser-7 was not dephosphorylated as efficiently as Ser-354 and Ser-404, even though this serine is followed by a lysine at the +3 position (Figure 4B). Nonetheless, these data confirm that at least two of the four perfect Cdk sites near the lqg1 N-terminal CHD behave as optimal Cdc14 substrate sites in vitro.
Actomyosin ring contraction defects in iqg1-4A cells

Because cells expressing iqg1-4A formed actin rings early and had cytokinesis defects, we used live-cell imaging to examine actomyosin ring contraction. We considered that actomyosin ring contraction might be occurring early, since the ring is assembled earlier, or that the rate or symmetry of contraction might be affected. For these experiments, we used cells expressing iqg1-4A, MYO1-green fluorescent protein (GFP), and TUB1-mCherry (Shannon and Li, 1999; Khmelinskii et al., 2007). Contraction of the actomyosin ring did not occur earlier in the cell cycle in iqg1-4A cells, as Tub1-mCherry showed that the spindle was completely elongated before contraction began (Figure 6A). In control cells, Myo1-GFP exhibited symmetric contraction to a single dot over an average of 8.1 ± 1.0 min, consistent with previous results (N = 6; Figure 6B; see Supplemental Figure 6B.mov for another example; Lippincott and Li, 1998b; Bi, 2001; Stockstill et al., 2013). However, Myo1-GFP failed to contract normally in iqg1-4A cells. Cells expressing iqg1-4A exhibited what appeared to be disassembly of Myo1-GFP, with the GFP signal becoming dimmer without the diameter of the ring decreasing (N = 6; Figure 6C; see Supplemental Figure 6C.mov for another example). This loss of Myo1-GFP signal occurred over the same length of time as contraction in control cells, 8.3 ± 2.2 min. To look at disassembly of Myo1-GFP, we generated a fluorescence intensity plot. In contrast to control cells, the Myo1-GFP signal rapidly decreased in intensity, without the peaks moving closer together as during contraction in control cells (Figure 6C). It was shown previously that preventing nuclear export of Cdc14 to the cytoplasm caused cytokinesis defects and a failure of Myo1-GFP to contract (Bembenek et al., 2005). Therefore timely dephosphorylation of Cdc14 targets at the bud neck may be essential for normal myosin contraction. A defect in Myo1-GFP contraction was recently reported in cells in which the Cdc14 target Inn1 was forced to undergo constitutive Cdk phosphorylation (Kuilman et al., 2015). Therefore there are at least two Cdc14 targets—iqg1 and Inn1—required for proper contraction of myosin during cytokinesis.

The bud necks of the iqg1-4A cells were significantly larger than control cells, 1.9 compared with 1.3 μm (p = 0.0002). Effects on the size of the bud neck have been seen after perturbation of many other cytokinesis and septation genes, including Hof1, Cyk3, Inn1, and Chs2 (Lippincott and Li, 1998a; Korinek et al., 2000; Stockstill et al., 2013; Kuilman et al., 2015). Mechanistically, the reason for the altered bud neck size is unknown, but it is likely due to the coupling between cytokinesis, septation, and selection of the future bud site in budding yeast. Our results suggest that timely phosphorylation and dephosphorylation of iqg1 is needed to ensure normal contraction of the ring during cytokinesis and maintenance of normal bud neck morphology.

**DISCUSSION**

Dephosphorylation of iqg1 by Cdc14 regulates actin ring assembly

Actin ring assembly in budding yeast is restricted to anaphase, but the mechanism by which this is accomplished has been unclear. Expression of iqg1 is cell cycle regulated, and Epp and Chant (1997) reported that overexpression of iqg1 was able to drive premature actin ring assembly. However, in our strain background, overexpression of IQG1 using the GAL1 promoter did not affect the timing of actin ring assembly. This can be seen from the data in Figure 2, where control cells cultured in YPGR that overexpress IQG1 do not form actin rings before anaphase.

In C. albicans, phosphorylation of iqg1 by Cdk1 at 15 perfect and minimal Cdk consensus sites was shown to affect the timing of actin ring formation (Li et al., 2008). During the course of our study, it was shown that inhibition of Cdk1 activity in metaphase cells and iqg1 mutations that prevent phosphorylation at N-terminal Cdk consensus sites total. This suggested that iqg1 function in directing final actomyosin ring maturation is controlled by multiple Cdk site clusters cause premature actin ring assembly in Saccharomyces cerevisiae (Naylor and Morgan, 2014). In that study, mutation of two distinct Cdk site clusters chosen based on proximity within the primary sequence resulted in partial actin ring assembly phenotypes, whereas a more severe effect on actin ring formation was observed with simultaneous mutation of both clusters, containing 14 perfect and imperfect Cdk sites total. This suggested that iqg1 function in directing final actomyosin ring maturation is controlled by multiple Cdk phosphorylation events spread over a relatively large primary sequence space. A role for Cdc14, or another phosphatase, in the dephosphorylation and regulation of iqg1 was not addressed in either of these studies. Here, we showed that 1) preventing phosphorylation of just the four perfect Cdk sites in the N-terminus of...
Iqg1 (which come from both of the clusters analyzed by Naylor and Morgan, 2014) is sufficient to cause premature actin ring formation and 2) Iqg1 is a novel substrate of Cdc14 whose dephosphorylation at a few perfect Cdk sites helps to regulate the timing of actin ring assembly and the fidelity of actomyosin ring function.

Our results are largely consistent with the conclusions of Naylor and Morgan (2014). Both studies found that Cdk phosphorylation of sites in the Iqg1 N-terminus restrain actin ring formation until late anaphase after chromosome segregation is completed. Both studies likewise observed early recruitment of Iqg1 to the division site when Cdk phosphosites were mutated. Also consistent with their work, we did not observe premature contraction of Myo1 in cells expressing the iqg1-4A mutant, confirming their conclusion that actomyosin ring assembly and constriction are regulated independently. Our results differ from theirs in that they did not report cytokinesis phenotypes or Myo1 contraction defects in their Iqg1 phosphomutant cells, and the reason for this difference is unclear, since both studies were done in the W303 background and involved integrated IQG1 alleles expressed under the endogenous promoter (Naylor and Morgan, 2014). However, our results are consistent with analysis of Iqg1 in C. albicans, where premature actin ring assembly correlated with cytokinesis defect and aberrant Myo1-GFP contraction (Li et al., 2008).

Of the four amino acids mutated in our study, serines at 354 and 404 of Iqg1 have been shown to be phosphorylated by Cdk1/Cdc28 in a cell cycle–dependent manner in vivo (Holt et al., 2009). We demonstrated that these two serines are Cdc14 substrates in vitro. Although phosphorylation of serine has not been observed at S7, this peptide may be poorly detected by MS. Serine 7 is predicted to be phosphorylated by the NetPhos2.0 neural network predictor of eukaryotic phosphorylation sites (Blom, 1999). This serine was not an effective Cdc14 substrate in vitro, so whether the phosphorylation and dephosphorylation of this residue occurs in vivo is unclear. Phosphorylation of the threonine at position 299 has also not been detected in vivo but is predicted to occur, although this threonine is not a Cdc14 substrate (Blom, 1999). Further studies are needed to determine whether the threonine at position 299 remains phosphorylated after the serines are dephosphorylated by Cdc14 and to define its functional importance.

Regulation of cytokinesis may be a conserved function of Cdc14 phosphatases. In addition to its requirement for cytokinesis in budding yeast, Cdc14 homologues have been reported to affect cytokinesis also in fission yeast, Caenorhabditis elegans, Xenopus laevis, and humans, although few molecular targets affecting cytokinesis have been identified in these systems (Mocciaro and Schiebel, 2010). In fission yeast, the Cdc14 homologue Clp1 localizes to the division site through binding to the anillin orthologue Mid1, where it stabilizes the contractile ring (Clifford et al., 2008). One key substrate of Clp1 at the contractile ring is Cdc15, a Hof1 homologue and an important regulator of actomyosin ring assembly, function, and septation (Clifford et al., 2008). The Iqg1 orthologue Rng2 may be another contractile ring substrate of Clp1, because it was identified as a Clp1 binding partner in a recent proteomics analysis, although the functional importance of the Clp1 interaction with Rng2 has not been demonstrated (Chen et al., 2013). Overall, few targets of Cdc14 orthologues directly involved in execution of cytokinesis...
are known. Our data reveal that Cdc14 functions in the final stage of actomyosin ring assembly in budding yeast and suggest that Iqg1 is the sole Cdc14 target that regulates the timing of actin ring assembly.

Role of the MEN in actomyosin ring assembly
In light of our results that Cdc14 function is needed for normal actin ring formation before cytokinesis, it is somewhat surprising that previous studies with conditional MEN mutants upstream of Cdc14 did not observe a similar defect (Vallen et al., 2000; Lippincott et al., 2001; Luca et al., 2001). A critical function of the MEN in promoting cytokinesis is to release Cdc14 from nucleolar sequestration and target it to the cytoplasm, where it can dephosphorylate its cytokinetic targets (Bembenek et al., 2005; Mohl et al., 2009; Sanchez-Diaz et al., 2012; Kuliman et al., 2015). In the previous studies of the MEN proteins Tem1 and Mob1, mitotic exit was induced by using either net1-1 temperature-sensitive allele or SIC1 overexpression, and it is likely that active Cdc14 was released from the nucleolus as a result (Luca and Winey, 1998; Lippincott et al., 2001). In the case in which dbf2-2 was examined at the semipermissive temperature or a CDC15 deletion mutant was used and actin ring formation was normal, there may not be a complete absence of Cdc14 activity (Menssen et al., 2001; Hwa Lim et al., 2003). One prior study did report that actin rings form normally in cdc14 cells (Vallen et al., 2000). However, the results were based on single time points after a long incubation at restrictive temperature and therefore would not reveal the changes in the kinetics of ring formation that we report here.

It will be important to determine in future studies the activity and localization of Cdc14 relative to actomyosin ring formation and contraction. We are evaluating actin ring formation in FEAR network mutants to test whether FEAR-released Cdc14 can support actin ring formation.

Mechanism by which phosphorylation regulates Iqg1 function
Another question raised by our study is how the phosphorylation of Iqg1 by Cdk1 prevents actin ring assembly before anaphase. One attractive hypothesis is that phosphorylation of the Cdk sites alters the ability of Iqg1 to bind to F-actin, since the Cdk sites flank the CHD. Alternatively, rather than affect the binding to F-actin directly, the phosphorylation mutants could affect Iqg1's interactions with formin proteins Bni1 and Bnr1, as has been shown in C. albicans (Li et al., 2008). Iqg1, Bni1, and Bnr1 are all required for actin ring assembly in budding yeast, but how they might cooperate in generating the actin ring is not known (Shannon and Li, 1999; Tolliday et al., 2002). Phosphorylation of Iqg1 could also prevent interaction with other proteins, such as those that help recruit Iqg1 to the contractile ring. This would be consistent with the observation that the Iqg1-4A mutant protein appears earlier than the wild-type protein at the bud neck. The IQ domains of Iqg1 interact with Mic1, and this interaction is important for recruiting Iqg1 to the contractile ring (Shannon and Li, 2000). The IQ domains are also situated in the Iqg1 N-terminal region just downstream of several of the optimal Cdc14 target sites (Figure 1A). However, the recent study by Naylor and Morgan (2014) did not find any evidence that Cdk site mutations affected the interaction between Iqg1 and Mic1, and the interaction between Iqg1-4A and Iqg1-4E mutants and GST-Mic1 in a GST pull-down assay was indistinguishable from wild-type Iqg1 (our unpublished data).

How mutation of the phosphorylation sites in Iqg1 causes cytokinesis defects is unclear. The interaction between Iqg1 and Tem1 is required for myosin ring contraction, but the GRD of Iqg1 that mediates the interaction with Tem1 is not near the Cdk sites (Shannon and Li, 1999). The mechanism by which the Iqg1-4A mutant affects contraction of the actomyosin ring could be due to loss of coordination between cytokinesis and septation (Bi, 2001). Iqg1 is assumed to have a function in septation, since it is an essential gene, and cytokinesis is not an essential process in budding yeast, but exactly how Iqg1 might function to link septation and cytokinesis is unclear.

One possibility is through an interaction with the Hof1 protein, which promotes primary septum deposition by Chs2 and is required for normal myosin contraction and bud neck size (Lippincott and Li, 1998a; Nishihama et al., 2009). Iqg1 interacts with Hof1 directly, and Hof1 appeared at the bud neck prematurely in cells expressing Cdk1 phosphorylation-deficient Iqg1 (Naylor and Morgan, 2014; Tian et al., 2014). Thus it is conceivable that the misregulated assembly of proteins in the contractile ring in the absence of normal Iqg1 phosphorylation results in a structurally and functionally compromised ring.

There are other possible mechanisms by which the Iqg1-4A mutant protein could perturb ring contraction. A quantitative model for generation of force in the actomyosin ring suggested that Iqg1 could augment cofilin function in the ring by acting as a cross-linker during actin filament depolymerization (Mendes Pinto et al., 2012). If mutation of the phosphorylation sites disrupts the ability of Iqg1 to act as an actin cross-linker during actomyosin ring contraction, it could result in lack of contraction seen. To test this hypothesis, the effect of mutations on Iqg1's cross-linking function and modeling studies will be needed.

Regardless of the mechanisms by which phosphorylation acts, a general conclusion of our work is that Cdc14 regulation of Iqg1 function contributes to the proper temporal coordination of cytokinesis after nuclear division is complete.

**MATERIALS AND METHODS**

**Strains and media**

All *S. cerevisiae* strains were derived from the W303 background and are listed in Table 1. Cultures were grown at 30°C in YEP + 2% D-glucose (YPD) or YEP + 2% galactose and raffinose (YPGR), except where noted.

**Plasmid construction and mutagenesis**

Plasmids are listed in Table 2. To make the IQG1 phosphomutant plasmids, the QuikChange II XL Multi Site-Directed Mutagenesis Kit (Agilent Technologies, Santa Clara, CA) was used to mutate pTL12, which contains full-length IQG1 under its endogenous promoter with a 13-myc tag on the C-terminus (Lippincott and Li, 1998b). The kit was used per manufacturer’s instruction with a few modifications. The PCR primer amount was sensitive and needed to be 200 ng with template DNA concentration at 30 ng. The *Pfu* was added 5 min after starting the following program: 95°C for 6 min, 1 cycle, 98°C for 1 min, 55°C for 1 min, 65°C for 20 min, 18 cycles, 72°C for 7 min, 1 cycle.

Minirep plasmid isolation on the Site-Directed Mutagenesis transformation colonies was performed according to the Wizard Plus SV Miniprep DNA Purification System (Promega, Madison, WI) protocol, and resulting purified DNA was sequenced by Missouri S&T cDNA Resource Center for verification of mutations. The four consensus Cdk sites are located at amino acids S7, T299, S345, and S404. All four sites were changed either to alanine or to glutamic acid, creating two final plasmids, expressing *iqg1-4A* (*pDM9*) with all sites mutated to alanine or *iqg1-4E* (*pDM15*) with all sites mutated to glutamic acid (Table 2). To integrate *iqg1-4A* at the
Table 2:

| Strain      | Genotype                      | Source                      |
|-------------|-------------------------------|-----------------------------|
| KSY184      | MATa ade2-1 can1-100 his3-11,15 leu2-3112 trp1-1 ura3-1 IQG1-3HA::His3MX6 | Ko et al. (2007)             |
| KSY286      | MATa ura3-1 his3-11,15 leu2-3112 lys-2-801 trp1-1 Δbar1 ΔIQG1:LEU2 pGAL1-IQG1-4myc, ura3 | This work                    |
| KSY378      | MATa ura3-1 his3-11,15 leu2-3112 lys-2-801 trp1-1 Δbar1 ΔIQG1:LEU2 pGAL1-IQG1-4myc, URA3 | This work                    |
| KSY396      | MATa ura3-1 his3-11,15 leu2-3112 lys-2-801 trp1-1 Δbar1 ΔIQG1:LEU2 pGAL1-IQG1-4myc, URA3 | This work                    |
| KSY472      | MATa ura3-1 his3-11,15 leu2-3112 lys-2-801 trp1-1 Δbar1 ΔIQG1:LEU2 pGAL1-IQG1-4myc, ura3 (pRL170) | This work                    |
| KSY473      | MATa ura3-1 his3-11,15 leu2-3112 lys-2-801 trp1-1 Δbar1 ΔIQG1:LEU2 pGAL1-IQG1-4myc, ura3 (pRL170) | This work                    |
| KSY482      | MATa cdc14-1 ura3 leu2 SIC1 2μ:LEU2 | This work                    |
| KSY496      | MATa ura3-1 his3-11,15 leu2-3112 lys-2-801 trp1-1 Δbar1 ΔIQG1:LEU2 pGAL1-IQG1-4myc, URA3 | This work                    |
| KSY504      | MATa cdc14-1 ura3 leu2 GAL-SIC1ΔNT:URA (pFM160) | This work                    |
| KSY508      | MATa ura3-1 his3-11,15 leu2-3112 lys-2-801 trp1-1 Δbar1 ΔIQG1:LEU2 pGAL1-IQG1-4myc, URA3 | This work                    |
| KSY509      | MATa ura3-1 his3-11,15 leu2-3112 lys-2-801 trp1-1 Δbar1 ΔIQG1:LEU2 pGAL1-IQG1-4myc, URA3 (pKL1341) | This work                    |
| KSY510      | MATa cdc14-1 ura3 leu2 ΔIQG1:LEU2 (pRL170) Δmyc-4A, HIS3 (pDM9) GAL-SIC1ΔNT:URA (pFM160) | This work                    |
| KSY520      | MATa ura3-1 his3-11,15 leu2-3112 lys-2-801 trp1-1 Δbar1 ΔIQG1:LEU2 pGAL1-IQG1-4myc, ura3 (pRL170) | This work                    |
| KSY522      | MATa ade2-1 can1-100 his3-11,15 leu2-3112 trp1-1 ura3-1 ΔIQG1:HA::His3MX6 | This work                    |
| YKA830      | MATa leu2::GAL1-3HA-CDC14(C283S) ura3::IQG1-3FLAG:URA3 | This work                    |
| YKA832      | MATa leu2::GAL1-3HA-CDC14(C283S) ura3::IQG1-3FLAG:URA3 | This work                    |

The background for all was W303.

Table 1: Yeast strains used in this study.

| Name  | Genotype                                      | Source                      |
|-------|-----------------------------------------------|-----------------------------|
| pKT36 | Myo1-GFP:TRP1 integrated with Adel            | Shannon and Li (1999)       |
| pTL12 | IQG1-myc CEN plasmid HIS3                    | Lippincott and Li (1998b)   |
| pDM9  | iqtg1-4A-myc CEN plasmid HIS3                 | This work                   |
| pDM15 | iqtg1-4E-myc CEN plasmid HIS3                 | This work                   |
| pFM160| GAL1-SIC1∆NT:URA3 integrated with EcoRV       | Chin et al. (2012)          |
| pKL1341| GAL1-CDC14:HIS integrated with Nhel           | Sanchez-Diaz et al. (2012)  |
| pAK011| pRS306-mCherry-TUB1:URA3 integrated with Apal | Khmelinskii et al. (2007)   |
| YEp13-SIC1| SIC1::LEU2 2μ plasmid                         | Luca et al. (2001)          |
| pHIP082| iqtg1-3A-3xFLAG:URA3                          | This work                   |
| pMM1  | iqtg1-4A-3xFLAG:URA3 integrated with AffII    | This work                   |

Table 2: Plasmids used in this study.

chromosomal locus, plasmid pHIP082 was used as template for site-directed mutagenesis using the primers to introduce the T299A mutation, and the resulting plasmid, pMM1, was sequenced to verify the mutation. pMM1 was cut with AffII and transformed into KSY184, and URA+ transformants containing the integrated plasmid were then plated on 5-fluoroorotic acid media to select for recombination between the IQG1-hemagglutinin (HA) and iqtg1-4A-3xFLAG alleles (Scherer and Davis, 1979). Genomic DNA from URA- colonies was sequenced to determine that KSY522 contained the iqtg1-4A-HA allele.

Yeast transformation

Yeast transformations were performed by a modified lithium acetate method (Gietz et al., 1992). After 30 min incubation at 30°C, 50 μl of dimethyl sulfoxide was added and mixed. Cells were plated on appropriate media and incubated at 30 or 25°C for 2–3 d.

Analysis of cell morphology

Strains KSY378, KSY396, and KSY520 have wild-type IQG1 under the GAL1-inducible promoter, allowing wild-type IQG1 to be shut off in the presence of glucose, and express iqtg1-4A, iqtg1-4E, and IQG1 respectively, under the IQG1 promoter. Cells were cultured overnight in 5 ml YPGR at 30°C. The cells were then diluted (1 ml into 5 ml) into YPGR (control) or YPD (experimental) media and were allowed to continue growing at 30°C for 5–7 h. Cell morphology was then observed using an Olympus CH2 (Olympus Scientific Solutions Americas Corp., Waltham, MA) with objective EA40, numerical aperture (NA) 0.65. For each strain and treatment, 200 cells were counted and scored as chains if they contained three or more connected cell bodies. Experiments were repeated three times.
Zymolyase treatment was performed as described (Lippincott and Li, 1998b).

**Collection of time points**

To examine the timing of actin ring formation, KSY378 or KSY520 cells were grown overnight in 5 ml YPGR, diluted into 50 ml, and grown for 3 h at 30°C. The α mating factor was added to a final concentration of 100 μg/ml and cells incubated for additional 3 h to arrest cells in G1. The cells were then pelleted and washed three times with sterile water to remove α factor. They were then resuspended in 35 ml ofYPD or YPGR. Time points were taken by removing 5 ml at 20, 40, 60, 80, and 100 min after resuspension into YPD or YPGR.

To examine the effect of Cdc14 overexpression on actin ring formation, strain KSY09 was grown overnight in 5 ml YPD at 30°C, and cells were pelleted and resuspended into YPD or YPGR with simultaneous addition of nocodazole at a final concentration of 5μg/ml and incubated for additional 2.5 h to arrest cells in mitosis.

To determine whether cdc14-1 cells could make actin rings at the nonpermissive temperature when mitotic arrest was bypassed with SIC1 expressed at high level using a 2μ plasmid, strain KSY482 was used (Tables 1 and 2). Cells were grown overnight in 5 ml YPD at room temperature (25°C). After resuspension into fresh YPD, nocodazole (5 μg/ml) was added and cells incubated for additional 2 h to arrest cells in mitosis. Cells were then placed at room temperature (control) and 37°C (experimental) for 90 min. A sample was collected before nocodazole was removed by washing cells twice with distilled H2O. After resuspension in fresh medium, samples were collected at 10- and 20-min time points were collected.

Strains KSY504 and KSY510, containing the temperature-sensitive cdc14-1 allele and integrated GAL1-SIC1ΔNT, were grown overnight in 5 ml of YPD at room temperature, resuspended into YPGR (to express SIC1 under the GAL1 promoter) with nocodazole (5 μg/ml), and incubated for 90 min to arrest cells in mitosis. Cells were then placed in YPGR with nocodazole at 37°C for an additional 90 min to allow for inactivation of cdc14-1. A sample was collected before nocodazole was removed by washing twice with distilled H2O. After resuspension in fresh medium, samples were collected at 10- and 20-min time points.

**Immunofluorescence**

Cells were fixed by addition of formaldehyde to 5% and rotation for 1 h at room temperature. Cells were then washed twice with sorbitol buffer (1 M sorbitol in 50 mM KPO₄, pH 7.5) to remove formaldehyde and stored at 4°C for up to 1 wk. Cells were permeabilized by Zymolyase treatment and then affixed to a 10-well microscope slide coated with polylysine. Cells were pipetted off, and the slide was allowed to air dry before being washed three times with 1 μg/ml bovine serum albumin in phosphate-buffered saline (Lippincott and Li, 1998b).

Cells were stained using primary antibody mouse anti-myc 9E10 (Covance, Princeton, NJ) and fluorescein isothiocyanate (FITC)–conjugated secondary antibody (Jackson Immunoresearch Laboratories, West Grove, PA) as described (Lippincott and Li, 1998b). Actin staining using A568 phalloidin (Invitrogen, Thermo Fisher Scientific, Waltham, MA) was also performed as described (Lippincott and Li, 1998b). Mounting solution containing 1 μg/ml DAPI was added before the coverslip was sealed to the slide.

Images were captured using an Olympus IX51 inverted microscope at 1000x total magnification using a UPLSAPO 100x/NA 1.4 objective. A BrightLine DA/Fl/TX-3x3M-a triple-band Sedat Filter set was used (Semrock, Rochester, NY). Images were captured with a Hamamatsu ORCA285 charge-coupled device (CCD) camera (Hamamatsu, Japan). A Prior motorized Z-drive was used to capture image stacks of 20 Z-planes in 0.2-μm steps (Prior Scientific, Rockland, MA). Shutter filters, and camera were controlled using Slide-Book software (Intelligent Imaging Innovations, Denver, CO), and this software was used to perform deconvolution of the image stacks and create projection images.

**Yeast protein extracts**

The protocol used is a modified version of Rigaut et al. (1999). KSY286, KSY472, KSY473, and KSY520 cells were grown overnight in YPGR and then diluted into YPD and arrested with α factor for 3 h. Cells were washed three times in sterile water and then released from α factor for 60 min before cells were pelleted and resuspended in 100 μl U of buffer (50 mM 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid, pH 7.5, 100 mM KCl, 3 mM MgCl₂, 1 mM ethylene glycol tetraacetic acid) plus protease inhibitors (0.5 μg/ml pepstatin, chymostatin, antipain, aprotinin, leupeptin) plus phenylmethylsulfonyl fluoride (1 mM). For cell lysis, 0.3 mg of acid-washed glass beads were added to cells and chilled on ice for 10 min. The cell suspension was vortexed five times (1 min on vortexer at maximum speed and 1 min on ice). The cell suspension was then centrifuged at 4°C at a relative centrifugal force of 16,000 × g for 5 min. The supernatant containing soluble protein was removed and the protein concentration determined using a NanoDrop 1000 (Thermo Scientific). An equal volume of 2× Laemmli sample buffer was added, and tubes were boiled for 5 min and then centrifuged for 5 min and frozen until use.

**Western blotting**

Protein samples were separated on 7.5 or 12.5% SDS–PAGE gels and then transferred to nitrocellulose and blocked in 5% milk in Tris-buffered saline/Tween before antibody staining. Mouse monoclonal anti-myc 9E10 (Covance) and mouse monoclonal anti-actin mAbGEa (Thermo Scientific Pierce) were used at 1:1000 dilution. Mouse anti-FLAG M2 (Sigma-Aldrich, St. Louis, MO) was used at 1:1000 dilution. EZview anti-HA-7 agarose resin (Sigma-Aldrich) was used for substrate trap coIP experiments. Donkey anti-mouse secondary antibody conjugated to horseradish peroxidase (Jackson Immunoresearch Laboratories) was used at 1:5000 concentration. The blot was then developed using an ECL kit (SuperSignal West Pico Chemiluminescent Substrate; Thermo Scientific) and imaged and analyzed on a Bio-Rad ChemiDoc MP system with Image Lab software (Bio-Rad Laboratories, Hercules, CA).

**Phosphatase assays and coimmunoprecipitation**

Recombinant Cdc14 was purified and assayed using synthetic phosphopeptide substrates (GenScript, Piscataway, NJ) as described (Bremmer et al., 2012). Peptide sequences were SG(pS)PSKPQGN (lqg1pS7), EY(pS)PIKNKL (lqg1pS354), HY(pS)MRERRM (lqg1pS404), YY(pS)PTISKYL (lqg1pS365), DF(pS) PVHKSFS (lqg1pS1347), and LIP(pT)PKNITD (lqg1pT299), where pS and pT are phosphoserine and phosphothreonine, respectively. Crude phosphopeptides were purified using Sep-Pak C18 cartridges (Waters Corporation, Milford, MA). Substrate trap coIP assays were performed as described (Eissler et al., 2014).

**Live-cell imaging**

KSY508 cells were grown overnight and placed on agarose pads made by melting 0.2 g of agarose in 1 ml TRP media (Waddle et al., 1996). The cells were then viewed using the Olympus Inverted Epifluorescent Microscope with a 100× Plan Apo/NA 1.4 DIC.
Objective. An FITC filter (EX 482/35 506DM EM 536/40) was used (Brightline). Images were captured with a Hamamatsu ORCA285 CCD camera. Shutters, filters, and camera were controlled using SlideBook software.

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