The Clp1/Cdc14 phosphatase contributes to the robustness of cytokinesis by association with anillin-related Mid1

Dawn M. Clifford,1 Benjamin A. Wolfe,1 Rachel H. Roberts-Galbraith,1 W. Hayes McDonald,2 John R. Yates III,2 and Kathleen L. Gould1

1Howard Hughes Medical Institute and Department of Cell and Developmental Biology, Vanderbilt University School of Medicine, Nashville, TN 37232
2Department of Cell Biology, Scripps Research Institute, San Diego, CA 93037

Cd14 phosphatases antagonize cyclin-dependent kinase–directed phosphorylation events and are involved in several facets of cell cycle control. We investigate the role of the fission yeast Cdc14 homologue Clp1/Flp1 in cytokinesis. We find that Clp1/Flp1 is tethered at the contractile ring (CR) through its association with anillin-related Mid1. Fluorescent recovery after photobleaching analyses indicate that Mid1, unlike other tested CR components, is anchored at the cell midzone, and this physical property is likely to account for its scaffolding role. By generating a mutation in mid1 that selectively disrupts Clp1/Flp1 tethering, we reveal the specific functional consequences of Clp1/Flp1 activity at the CR, including dephosphorylation of the essential CR component Cdc15, reductions in CR protein mobility, and CR resistance to mild perturbation. Our evidence indicates that Clp1/Flp1 must interact with the Mid1 scaffold to ensure the fidelity of Schizosaccharomyces pombe cytokinesis.

Introduction

In eukaryotes, progression through the cell cycle is directed by the activity of CdkS. Reversal of Cdk-dependent phosphorylation events involves the highly conserved Cdc14 family of phosphatases. First characterized for their essential role in facilitating mitotic exit in budding yeast, Cdc14 family members are now known to play numerous cell cycle roles, ranging from regulating centrosome duplication to controlling the terminal events of cytokinesis (Stegmeier and Amon, 2004). For example, cells lacking the fission yeast Cdc14 homologue Clp1/Flp1 (hereafter referred to as Clp1) exhibit defects in chromosome segregation (Trautmann et al., 2004), cytokinesis (Cueille et al., 2001; Trautmann et al., 2001), and mitotic exit (Esteban et al., 2004; Wolfe and Gould, 2004). Although the roles of Cdc14 phosphatases in cell division have been best characterized in yeast, cytokinesis defects are also observed in human cells after depletion of hCdc14A (Kaiser et al., 2002; Mailand et al., 2002). Thus, a regulatory role for Cdc14 phosphatases in mitotic exit and cytokinesis appears conserved in multiple organisms.

Cdc14 phosphatases localize dynamically to many subcellular compartments to carry out their functions (Stegmeier and Amon, 2004). For example, Clp1 is primarily nucleolar and at the spindle pole body during interphase (Cueille et al., 2001; Trautmann et al., 2001). During mitosis, Clp1 disperses throughout the nucleus and cytoplasm, concentrating at kinetochores (Trautmann et al., 2004), the mitotic spindle, and the contractile ring (CR), a highly dynamic structure that controls ingression of the division plane. Clp1 remains at the CR during its constriction and then reconcentrates in the nucleolus.

Consistent with its localization at the CR, several findings have established a regulatory role for Clp1 in cytokinesis. Cells lacking Clp1 display strong negative genetic interactions with many cytokinesis mutants, including those which disrupt CR assembly, septum formation, and cell wall synthesis (Trautmann et al., 2001; Mishra et al., 2004). In addition, genetic studies indicate that Clp1 stabilizes CRs when actin polymerization is inhibited by latrunculin A (lat A) treatment (Mishra et al., 2004). However, it is not known how Clp1 affects the process of cytokinesis nor whether Clp1’s interaction with the CR is important for this function.

Proper assembly and localization of the CR for cell division in fission yeast requires Mid1, an anillin-related protein (Chang et al., 1996; Sohrmann et al., 1996). In early mitosis, Mid1 exits...
the nucleus and directly interacts with the cell cortex, anchoring the CR in the cell center (Celton-Morizur et al., 2004). In the absence of \textit{mid1}, CR structures are frequently off centered and/or tilted leading to unequal nuclear and cellular division (Chang et al., 1996; Sohrmann et al., 1996). Once the division site has been established, several other proteins are recruited to this region to assemble the CR. One essential ring component that Mid1 recruits to the cortical region is myosin heavy chain Myo2, and mitotic dephosphorylation of the Myo2 C terminus has been implicated in regulating this interaction (Motegei et al., 2004).

To investigate the role of Clp1 at the division site, we used a proteomics approach to identify Mid1 as a Clp1-interacting protein and defined Mid1 association as the mechanism of Clp1 recruitment to the CR. By selectively disrupting Clp1–Mid1 interaction, we have been able to determine the functional consequence of Clp1 recruitment to the CR. We find that the phosphorylation status and dynamic properties of key CR components are altered in the absence of Clp1 activity and that otherwise silent mutations of CR components lead to cytokinetic failures in the absence of Clp1 CR localization. Our data offer a mechanistic explanation for Clp1’s requirement in the fidelity of \textit{Schizosaccharomyces pombe} cytokinesis.

\section*{Results}

\subsection*{Clp1 and Mid1 interact in vivo}
To identify proteins that might link Clp1 to the CR, we isolated tandem affinity purification (TAP) complexes from \textit{nda3} (β-tubulin)-\textit{KM311}–arrested cells producing C-terminally TAP-tagged Clp1. These cells arrest in a premetaphase-like state with a CR but lack a mitotic spindle (Hiraoka et al., 1984). The protein composition of the complexes was analyzed by 2D liquid chromatography tandem mass spectrometry. In addition to Clp1 at nearly 100\% sequence coverage, this strategy identified Mid1 with 39.9\% sequence coverage. Confirming the TAP analysis, reciprocal coimmunoprecipitation experiments demonstrated that Clp1 and Mid1 associate (Fig. S1 A, available at http://www.jcb.org/cgi/content/full/jcb.200709060/DC1). Yeast two-hybrid analysis and in vitro binding assays showed that an internal region of Mid1 (aa 331–534) interacted directly with the N-terminal catalytic domain of Clp1 (Fig. S1, B–E). Other proteins were identified in Clp1–TAP complexes and the validity of these hits is being investigated. No other CR components were recovered with such high coverage (unpublished data).

\subsection*{Clp1 requires \textit{mid1} for localization to the CR}
During early mitosis, both Clp1 and Mid1 localize to the CR (Fig. 1 A; Sohrmann et al., 1996; Cueille et al., 2001; Trautmann et al., 2001). Therefore, we examined whether the CR localization of either protein is dependent on the other. Live-cell imaging of Mid1-GFP cells demonstrated that Clp1 does not affect Mid1 ring localization (Fig. 1 A). In \textit{nda3-KM311}–arrested cells, Clp1-GFP was observed at kinetochores and in rings (Fig. 1 A). In \textit{nda3-KM311 mid1Δ} cells, however, Clp1-GFP rings were not detected, although Clp1-GFP was still observed at kinetochores (Fig. 1 A). Lack of Clp1-GFP rings in \textit{mid1Δ} cells was not because of a change in Clp1 levels (Fig. 2 E and Fig. S1 F). To verify that CRs could be detected in \textit{mid1Δ} cells, we examined the localization of Clp1-YFP together with Cdc15-CFP, which localizes to the CR independently of Mid1 (Sohrmann et al., 1996). In 97\% (193/198) of \textit{nda3-KM311}–arrested cells examined, Clp1-YFP and Cdc15-CFP colocalized in rings (Fig. 1 B). Although Cdc15-CFP formed rings in \textit{mid1Δ} cells, Clp1-YFP could not be detected in these rings (0/102 cells). Time-lapse videomicroscopy of cells progressing through mitosis confirmed that Clp1-GFP localized to kinetochores and the mitotic spindle in \textit{mid1Δ} cells with similar timing and intensity as in \textit{mid1Δ} cells but failed to form a ring (Fig. 1 C; Videos 1 and 2, available at http://www.jcb.org/cgi/content/full/jcb.200709060/DC1). These data indicate that Mid1 is necessary to recruit Clp1 to the CR.

The \textit{S. pombe} CR is a highly dynamic structure (Pelham and Chang, 2002; Wong et al., 2002). However, if Mid1 serves as a scaffold to link the cortex with CR components, such as Clp1, we reasoned that Mid1 might remain stably bound to the cortex. This was tested by FRAP analysis of Mid1-GFP. After spindle pole body separation, which was detected with Sid4-GFP (Chang and Gould, 2000), Mid1-GFP rings were bleached and recovery was monitored for the duration of Mid1 ring localization in a nonbleached cell (Fig. 1 D–E; and Video 3, available at http://www.jcb.org/cgi/content/full/jcb.200709060/DC1). Bleached Mid1-GFP rings showed little signal recovery after 15 min, indicating that Mid1 is very immobile while at the CR.

\subsection*{Clp1 regulates Cdc15 dephosphorylation at the CR}
A plausible role for the Mid1–Clp1 interaction is to position Clp1 at the CR so that it can dephosphorylate CR components and thereby modify CR properties. Although several CR components are phosphoproteins (Wolfe and Gould, 2005) and might be Clp1 substrates, one essential CR component whose phosphorylation is known to vary during the cell cycle is Cdc15 (Fankhauser et al., 1995). Cdc15 is essential for cytokinesis (Fankhauser et al., 1995) and is responsible for recruiting F-actin nucleators to the division site (Fankhauser et al., 1995; Carnahan and Gould, 2003). To investigate whether Cdc15 dephosphorylation depends on Clp1, Cdc15 phosphorylation status was monitored in the presence and absence of \textit{clp1}. As observed previously (Fankhauser et al., 1995), Cdc15 became progressively dephosphorylated as cells progressed through mitosis. Corresponding with the time of CR formation, Cdc15 was partially dephosphorylated as cells progressed through mitosis. Corresponding with the time of CR formation, Cdc15 was partially dephosphorylated as cells progressed through mitosis.
Clp1 localization to the CR does not occur (Fig. 1). Indeed, Cdc15 in nda3-KM311–arrested mid1/H9004 cells was in the hyperphosphorylated form, comigrating with Cdc15 from nda3-KM311–arrested clp1/H9004 cells (Fig. 2C). Furthermore, Clp1-dependent Cdc15 dephosphorylation was not detected in synchronized mid1/H9004 cells progressing through mitosis (Fig. 2D; and Fig. S2B). Clp1 phosphatase activity assays verified that the difference in Cdc15 phosphorylation was not a result of reduced Clp1 phosphatase activity in the absence of mid1 (Fig. 2E). Collectively, our results indicate that Mid1 is required to properly localize mitosis (Trautmann et al., 2004). Further supporting the idea that Cdc15 is a Clp1 target, recombinant Clp1 dephosphorylated the partially phosphorylated Cdc15 in immunocomplexes purified from nda3-KM311–arrested cells (Fig. 2F) and a small amount of Clp1 specifically associated with Cdc15 immune complexes isolated from nda3-KM311–arrested cells (Fig. S2C). These results indicate that Clp1 phosphatase activity is required for maximal Cdc15 dephosphorylation during mitosis.

We then reasoned that Clp1-dependent Cdc15 dephosphorylation should also be compromised in mid1Δ cells, in which Clp1 localization to the CR does not occur (Fig. 1). Indeed, Cdc15 in nda3-KM311–arrested mid1Δ cells was in the hyperphosphorylated form, comigrating with Cdc15 from nda3-KM311–arrested clp1Δ cells (Fig. 2C). Furthermore, Clp1-dependent Cdc15 dephosphorylation was not detected in synchronized mid1Δ cells progressing through mitosis (Fig. 2D; and Fig. S2B). Clp1 phosphatase activity assays verified that the difference in Cdc15 phosphorylation was not a result of reduced Clp1 phosphatase activity in the absence of mid1 (Fig. 2E). Collectively, our results indicate that Mid1 is required to properly localize mitosis (Trautmann et al., 2004). Further supporting the idea that Cdc15 is a Clp1 target, recombinant Clp1 dephosphorylated the partially phosphorylated Cdc15 in immunocomplexes purified from nda3-KM311–arrested cells (Fig. 2F) and a small amount of Clp1 specifically associated with Cdc15 immune complexes isolated from nda3-KM311–arrested cells (Fig. S2C). These results indicate that Clp1 phosphatase activity is required for maximal Cdc15 dephosphorylation during mitosis.

Figure 1. Clp1 depends on Mid1 for localization to the CR. (A) Live-cell images of nda3-KM311 mid1-GFP, nda3-KM311 mid1-GFP clp1Δ, nda3-KM311 clp1-GFP, and nda3-KM311 clp1-GFP mid1Δ cells after incubation at 18°C for 7 h. (B) Live-cell images of nda3-KM311 clp1-YFP cdc15-CFP and nda3-KM311 clp1-YFP cdc15-CFP mid1Δ cells after arrest by incubation at 18°C for 7 h. Over 100 cells were examined for each strain. (C) Time-lapse confocal microscopy of live cdc25-22 clp1-GFP and cdc25-22 clp1-GFP mid1Δ cells during mitosis. Exponentially growing cells were incubated at the restrictive temperature (36°C) for 3.5 h, and then released to the permissive temperature (25°C) for 15 min. Indicated time points are from time of release. Arrows indicate Clp1 ring. Images are from Videos 1 and 2 (available at http://www.jcb.org/cgi/content/full/jcb.200709060/DC1). (D and E) Representative images (D) and fluorescence recovery curves (E) for Mid1-GFP FRAP in mid1-GFP sid4-GFP cells. Sid4-GFP signal determined mitotic stage and vertical lines indicate the window when Mid1 begins to dissociate from the CR. B, bleach region; NB, nonbleached region (used to correct for overall bleaching); BKGD, background region (used to correct for overall bleaching). Images are from Video 3 (available at http://www.jcb.org/cgi/content/full/jcb.200709060/DC1). Bars, 5 μm.
Clp1 to the CR where Clp1 substrates, such as Cdc15, undergo Clp1-dependent dephosphorylation.

Clp1 phosphatase activity regulates Cdc15 and myosin ring stability

Given the previously established role for Clp1 in stabilizing CRs in response to mild perturbation (Mishra et al., 2004), we reasoned that even in the absence of CR disruption there might be an intrinsic difference in the dynamics of CR components in the absence of Clp1 function. To address this possibility, we examined the dynamics of four CR components involved in CR assembly and stability: Cdc15, the myosin II components, Myo2 (heavy chain) and Rlc1 (regulatory light chain), and the actin binding IQGAP, Rng2. Myosin II was of particular interest, as its recruitment to the CR is partially dependent on Mid1, and Myo2 dephosphorylation has been implicated in mediating a Mid1–Myo2 interaction (Motegi et al., 2004). In particular, a regulatory dephosphorylation event has been shown to occur in a C-terminal Myo2 fragment (Motegi et al., 2004) and Clp1 can dephosphorylate this fragment (Fig. S2 D), suggesting Myo2 as a C-terminal Myo2 fragment (Motegi et al., 2004) and Clp1 can dephosphorylate this fragment (Motegi et al., 2004). In particular, a regulatory dephosphorylation event has been shown to occur in a C-terminal Myo2 fragment (Motegi et al., 2004) and Clp1 can dephosphorylate this fragment (Motegi et al., 2004).

As in clp1Δ cells, the half-times of Cdc15-GFP, GFP-Myo2, and Rlc1-GFP were not significantly altered in clp1Δ-C286S cells relative to wild type (Fig. 3 B, and Fig. 4, A and B) but the mean mobile fraction was higher (Fig. 3, B and C; and Fig. 4, A and B). Interestingly, there was no difference in the half-time or mean mobile fraction of YFP-Rng2 in the absence of Clp1 activity (Fig. 4 C). GFP-Myo2, Rlc1-GFP, and YFP-Rng2 levels were similar in wild-type and clp1 mutant cells (Fig. S2 E). In addition, a significant increase in Cdc15 levels was not observed in clp1-C286S cells (Fig. S2 A), indicating that the observed differences in Cdc15 dynamics in clp1Δ cells are unlikely to result from the modest increase in overall Cdc15 levels in this strain. These results not only indicate that Clp1 influences CR component dynamics selectively to provide stability to the CR but provide the first evidence that the dynamics of individual CR components are different.

Clp1 at the CR regulates cytokinesis

To determine if physical association of Clp1 with the CR is required for CR stability, we developed a strain expressing a Mid1 mutant that disrupts its interaction with Clp1. Yeast two-hybrid analysis was used to narrow the region within Mid1 necessary for Clp1 binding. Mid1 deleted of aa 431–481 failed to interact with Clp1 (unpublished data). When integrated at the endogenous mid1 locus and C-terminally tagged with GFP, mid1Δ431–481-GFP cells formed medial septa and Mid1Δ431–481-GFP localized at the CR (Fig. 5 A), which was centrally located and perpendicular to the cell length. In addition, protein levels of Mid1Δ431–481-GFP were similar to those of wild-type Mid1-GFP (Fig. 5 B). Myo2 and the polo-like kinase Plo1 both depend on Mid1 for proper CR localization and tight ring formation (Bahrle et al., 1998a; Motegi et al., 2004). GFP-Myo2, Rlc1-GFP, and Plo1-GFP CR localization were...
Genetic interactions with \( `{clp1}\) (Mishra et al., 2004). At the semipermissive temperature (32 °C), \( `{cdc15-140}\) (Fankhauser et al., 1995), formin \( `{cdc12-112}\) (Chang et al., 1996), profilin \( `{cdc3-124}\) (Balasubramanian et al., 1994), and 1,3-\( \beta\)-glucan synthase \( `{cps1-191}\) (Liu et al., 1999) cells are viable and the majority of the population contains one nucleus, as is the case for wild-type cells (Fig. 6, A–C). However, under the same conditions, double mutants with \( `{mid1}\) did not form colonies (Fig. 6 A) and the majority of cells accumulated two or more nuclei (Fig. 6, B and C), which is indicative of cytokinetic failure. These results indicate that \( `{clp1}\) activity is required at the CR for successful cytokinesis when specific CR or septum assembly components are compromised.

**Discussion**

In this paper, we investigate the function of \( `{clp1}\) at the division site and pinpoint a phosphatase-dependent regulatory mechanism that influences CR dynamics and stability. We identify anillin-related \( `{mid1}\) as the key targeting molecule recruiting \( `{clp1}\) to the CR but not affecting \( `{clp1}\)’s other localizations, making \( `{mid1}\) the first reported cytoplasmic Cdc14 phosphatase tether.
bilayers (Itoh et al., 2005), and Cdc15 also uses this domain to bind the formin Cdc12 (Carnahan and Gould, 2003). In addition to Cdc15, other *pombe cdc15* homology family members are phosphoproteins (Chitu and Stanley, 2007), and phosphoregulation may be a conserved mechanism to control their functions. Analysis of Cdc15 phosphosite mutants in vivo will be necessary to test the precise molecular mechanism of its regulation, and an investigation of other potential Clp1 CR targets will likely contribute to a more complete picture of the complexity of CR regulation.

Actin and myosin are initially recruited to the division site independently (Motegi et al., 2000). Several lines of evidence indicate that later mitotic events, including their turnover at the division site, require their cooperation. In mammalian cells, myosin II light chain influences the turnover of actin at the CR via a phosphorylation-dependent mechanism (Murthy and Wadsworth, 2005). In rat cells, inhibition of myosin II motor activity causes a twofold increase in the recovery time of actin, as assessed by FRAP, which leads to an accumulation of actin at the division site and eventual cytokinesis failure (Guha et al., 2005). We observed an approximate twofold increase in Cdc15 and myosin II mobility when Clp1 phosphatase activity was defective. In addition, we observed an increase in cytokinesis failure in cells lacking Clp1 activity at the CR when CR formation was disturbed. Given that Clp1 influences two major components that provide structural integrity to the CR and are among the most abundant proteins concentrated at the division site (Wu and Pollard, 2005),

Once localized at the CR through Mid1 binding, Clp1 regulates the kinetics with which Cdc15 and myosin II are maintained at the CR. CR-associated Clp1 becomes essential for successful cytokinesis when specific components of the cytokinetic machinery are challenged, adding necessary robustness to the process of cell division.

Although we have a good understanding of the temporal assembly of the *S. pombe* CR (Wu et al., 2003), the mechanisms that regulate this ordered process are still unclear. Our FRAP results strongly support a scaffolding role for Mid1. Probably its stable association with the cortex, in marked contrast to the dynamic nature of other tested CR components (Pelham and Chang, 2002; Wong et al., 2002; this study), allows Mid1 to effectively recruit and organize other CR proteins into a focused ring structure and to secure the CR in place. Recruiting Clp1 is apparently one function of Mid1 in the CR assembly process.

The involvement of Clp1 phosphatase activity in regulating *S. pombe* CR dynamics is indicative of a phosphorylation/dephosphorylation-driven mechanism, and we have investigated Cdc15 as one of probably several Clp1 targets at the CR. Cdc15 phosphorylation is altered in *clp1Δ, clp1-C286S, mid1Δ*, and *mid1Δ* cells, and an alteration in Cdc15 phosphorylation status in *clp1Δ* cells was also reported by Wachtler et al. (2006). What is the function of Clp1-mediated Cdc15 dephosphorylation? Clp1 might regulate interactions between Cdc15 and other CR proteins or plasma membrane lipids. The F-BAR/EFC domain that is present in Cdc15 generally binds phospholipid
any alteration to their regulation is likely to disrupt the fidelity of cytokinesis. FRAP analysis indicated that Rng2 dynamics were unaffected by Clp1 and that the recovery of Cdc15 after photobleaching was measurably faster than myosin II or Rng2. These data are the first to indicate variable CR component dynamics.

Based on our results, Clp1 at the CR regulates the dynamic properties of specific CR components during an unperturbed mitosis, and this mechanism is likely to account in part for the observed cytokinesis failure in 3–5% of a clp1 Δ population (Cueille et al., 2001; Trautmann et al., 2001). When the natural assembly of the CR or septum is challenged, Clp1 activity is specifically required at the CR to ensure CR integrity and completion of cytokinesis. In budding yeast, Cdc14 export from the nucleus to the cytoplasm is required for CR constriction (Bembenek et al., 2005), raising the possibility that Cdc14, like Clp1, needs to anchor with the CR for proper ring function. However, our data also indicate that Clp1 must be able to influence CR stability from other locations. Cell cycle progression in the presence of low doses of lat A requires septation initiation network (SIN) signaling and Clp1 activity (Mishra et al., 2004). Although Clp1 functions to maintain SIN activity, the SIN is required for nuclear exclusion of Clp1 until cytokinesis is complete (Trautmann et al., 2001; Mishra et al., 2004). We found that Clp1 activity at the CR is dispensable for Clp1-SIN signaling in response to low-dose lat A treatment (Fig. S3). This is consistent with the finding that Clp1 is dispersed throughout the cytoplasm rather than concentrated at the CR in the presence of low-dose lat A (Mishra et al., 2004). Our analyses of the mid1Δ431-481 mutant demonstrate that Clp1 utilizes at least two mechanisms to influence cytokinesis,
| Strain     | Genotype                                                                 | Reference                  |
|-----------|---------------------------------------------------------------------------|----------------------------|
| KGY246    | h-- ade6-M210 leu1-32 ura4-D18                                           | Laboratory stock           |
| KGY648    | h+ cdc3-124 clp1::ura4 ade6-M21X leu1-32 ura4-D18                         | Laboratory stock           |
| KGY1278   | h+ rcl1-GFP:ura4 ade6-M210 leu1-32 ura4-D18                               | Laboratory stock           |
| KGY1985   | h+ cdc25-22 cdc15-HA:kan5 clp1::ura4 ade6-M210 leu1-32 ura4-D18           | This study                 |
| KGY2417   | h- mid1-GFP:kan5 ade6-M210 leu1-32 ura4-D18                               | Laboratory stock           |
| KGY2882   | h- clp1-MYC:kan5 ade6-M218 leu1-32 ura4-D18                               | Trautmann et al. (2001)    |
| KGY3019   | h- cdc15-GFP:kan5 ade6-M210 leu1-32 ura4-D18                               | Carnahan and Gould (2003)   |
| KGY3078   | h- cdc15-140 mid1-GFP:kan5 ade6-M21X leu1-32 ura4-D18                      | Laboratory stock           |
| KGY3079   | h- mid1-GFP:kan5 mid1-GFP:kan5 ade6-M21X leu1-32 ura4-D18                 | Laboratory stock           |
| KGY3155   | h- cdc15-140 mid1-GFP:kan5 ade6-M21X leu1-32 ura4-D18                      | Laboratory stock           |
| KGY3173   | h+ nda3-KM311 mid1-GFP:kan5 ade6-M210 leu1-32                              | This study                 |
| KGY3350   | h- cdc25-22 cdc15-HA:kan5 ade6-M21X leu1-32 ura4-D18                      | This study                 |
| KGY3381   | h- clp1::ura4 ade6-M216 leu1-32 ura4-D18                                  | This study                 |
| KGY3382   | h- cdc25-22 cdc15-HA:kan5 ade6-M21X leu1-32 ura4-D18                      | This study                 |
| KGY3388   | h+ nda3-KM311 clp1-MYC:kan5 leu1-32                                       | This study                 |
| KGY3612   | h- nda3-KM311 ura4-D18 leu1-32                                            | Laboratory stock           |
| KGY3783   | h+ nda3-KM311 clp1::ura4 ade6-M21X leu1-32 ura4-D18                        | Wölfe and Gould (2004)      |
| KGY4410   | h- nda3-KM311 cdc15-HA:kan5 ade6-M21X leu1-32                              | This study                 |
| KGY4630   | h+ nda3-KM311 cdc15-HA:kan5 ade6-M21X leu1-32                              | This study                 |
| KGY4791   | h- cdc25-22 cdc15-HA:kan5 ade6-M21X leu1-32                               | This study                 |
| KGY4815   | h+ cdc25-22 cdc15-HA:kan5 ade6-M21X leu1-32                               | This study                 |
| KGY4833   | h+ cdc15-140 clp1::ura4 ade6-M21X leu1-32 ura4-D18                        | Laboratory stock           |
| KGY4881   | h+ clp1-MYC:kan5 mid1::ura4 ade6-M216 leu1-32 ura4-D18                   | This study                 |
| KGY5298   | h+ kan5::GFP-myo2 ade6-M210 leu1-32 ura4-D18                              | Wu et al. (2003)           |
| KGY5318   | h+ kan5::YFp02 ade6-M210 leu1-32 ura4-D18                                 | Wu et al. (2003)           |
| KGY5482   | h+ cdc25-22 cdc15-HA:kan5 ade6-M21X leu1-32 ura4-D18                      | This study                 |
| KGY5594   | h- nda3-KM311 cdc15-HA:kan5 clp1::ura4 leu1-32                            | This study                 |
| KGY5598   | h+ nda3-KM311 cdc15-HA:kan5 mid1::ura4 ade6-M21X leu1-32 ura4-D18         | This study                 |
| KGY5615   | h+ cdc25-22 cdc15-HA:kan5 mid1::ura4 ade6-M21X leu1-32 ura4-D18           | This study                 |
| KGY5616   | h+ cdc25-22 cdc15-HA:kan5 mid1::ura4 ade6-M21X leu1-32 ura4-D18           | This study                 |
| KGY5617   | h+ cdc25-22 cdc15-HA:kan5 mid1::ura4 ade6-M21X leu1-32 ura4-D18           | This study                 |
| KGY5626   | h+ cdc25-22 cdc15-HA:kan5 mid1::ura4 ade6-M21X leu1-32 ura4-D18           | This study                 |
| KGY5627   | h- cdc15-GFP:kan5 clp1::ura4 ade6-M21X leu1-32 ura4-D18                   | This study                 |
| KGY5630   | h+ cdc25-22 cdc15-GFP:kan5 clp1::ura4 ade6-M21X leu1-32 ura4-D18          | This study                 |
| KGY5631   | h+ cdc25-22 cdc15-GFP:kan5 clp1::ura4 ade6-M21X leu1-32 ura4-D18          | This study                 |
| KGY5667   | h+ cdc25-22 cdc15-GFP:kan5 clp1::ura4 ade6-M21X leu1-32 ura4-D18          | This study                 |
| KGY5812   | h+ cdc25-22 cdc15-GFP:kan5 mid1::ura4 ade6-M21X leu1-32 ura4-D18          | This study                 |
| KGY5907   | h+ cdc15-GFP:kan5 C2865-MYC:kan5 ade6-M21X leu1-32 ura4-D18               | This study                 |
| KGY5924   | h- rcl1-GFP::ura4 clp1::ura4 ade6-M21X leu1-32 ura4-D18                   | This study                 |
| KGY5925   | h- rcl1-GFP::ura4 C2865-MYC:kan5 ade6-M21X leu1-32 ura4-D18               | This study                 |
| KGY5944   | h- kan5::YFP-myo2 ade6-M21X leu1-32 ura4-D18                              | This study                 |
| KGY5946   | h- kan5::YFP-myo2 clp1::ura4 ade6-M21X leu1-32 ura4-D18                   | This study                 |
| KGY5956   | h+ kan5::GFP-myo2 clp1::ura4 ade6-M21X leu1-32 ura4-D18                   | This study                 |
| KGY5958   | h+ kan5::GFP-myo2 ade6-M21X leu1-32 ura4-D18                              | This study                 |
| KGY6039   | h+ cdc25-22 cdc15-HA:kan5 ade6-M21X leu1-32 ura4-D18                      | This study                 |
| KGY6040   | h- cdc25-22 cdc15-HA:kan5 clp1::ura4 ade6-M21X leu1-32 ura4-D18           | This study                 |
| KGY6462   | h- mid1::ura4 ade6-M210 leu1-32 ura4-D18                                  | This study                 |
| KGY6485   | h+ mid1::ura4 ade6-M210 leu1-32 ura4-D18                                  | This study                 |
| KGY6486   | h+ mid1::ura4 ade6-M210 leu1-32 ura4-D18                                  | This study                 |
| KGY6487   | h+ mid1::ura4 ade6-M210 leu1-32 ura4-D18                                  | This study                 |
| KGY6549   | h+ cdc15-140 mid1::ura4 ade6-M21X leu1-32 ura4-D18                        | This study                 |
| KGY6597   | h+ cdc15-140 mid1::ura4 ade6-M21X leu1-32 ura4-D18                        | This study                 |
| KGY6599   | h+ cdc15-GFP::kan5 mid1::ura4 ade6-M21X leu1-32 ura4-D18                  | This study                 |
| KGY6605   | h+ cdc25-22 cdc15-HA:kan5 ade6-M21X leu1-32 ura4-D18                      | This study                 |
| KGY6612   | h+ cdc15-140 mid1::ura4 ade6-M21X leu1-32 ura4-D18                        | This study                 |
| KGY6624   | h+ cdc15-140 mid1::ura4 ade6-M21X leu1-32 ura4-D18                        | This study                 |
| KGY6626   | h+ cdc15-140 mid1::ura4 ade6-M21X leu1-32 ura4-D18                        | This study                 |
one of which requires direct association with the CR and the other of which involves signaling to the cytokinetic machinery. The dual activities of Clp1 are further indications of the robustness of cytokinesis, a process protected by several redundant mechanisms. Cdk activity inhibits cytokinesis in both human and yeast cells (Murray, 2004; Wolfe and Gould, 2005) but the mechanism is unknown. Given that Clp1 dephosphorylates sites of Cdk phosphorylation, these studies reveal at least one likely mechanism by which Cdk activity antagonizes cytokinesis and suggest that through similar mechanisms other Cdc14 family members will be found to regulate CR dynamics.

Materials and methods

Strains, media, and molecular biology methods

The S. pombe strains used in this study (Table I) were grown in yeast extract (YE) as described in the figure legends. Induction of the nmt promoter (Maudelin, 1993) was achieved by growing cells in thiamine (repressing conditions) and then washing cells three times in medium (YE) as described in the figure legends. Induction of the nmt promoter was PCR amplified with oligos containing NdeI and BamHI sites on the endogenous GFP:kan R , CFP:kan R , or TAP:kan R cassettes as previously described (Tasto et al., 2003). Strains used in this study (Table I) were grown in yeast extract (YE) as described (Wolfe et al., 2006). In vitro phosphatase assays

Immunoprecipitated Cdc15-HA and HA-Myo2-Ct were incubated in the presence of recombinant MBP, MBP-Clp1, or phosphatase-dead MBP-Clp1 (MBP-Clp1Δ96) at 30°C for 30–45 min in phosphatase assay buffer (50 mM imidazole, pH 6.9, 1 mM EDTA, and 1 mM DTT). Reactions were terminated by the addition of SDS sample buffer and by boiling for 5 min. Proteins were resolved by SDS-PAGE and detected by Western blot with anti-HA. Clp1-MYC phosphatase activity from cells was determined by DIFMUP (Invitrogen) continuous assays as previously described (Wolfe et al., 2006).

Microscopy

GFP, YFP, or GFP-tagged proteins were visualized using a spinning disk confocal microscope (UltraView LCI; PerkinElmer) equipped with a 100×/1.40 Plan-Apochromat oil immersion objective. All images were acquired at 25°C. For time-lapse images, cells were placed on a hanging drop glass slide containing YE agar and covered with a coverslip. Images were captured with a charge-coupled device camera (Orcar E; Hamamatsu) using Metamorph software (1.7, MDS Analytical Technologies) or Ultraview LCI software (PerkinElmer) and then processed using Velocity software (version 3.5.1). For time-lapse and static images, z-series optical sections were taken at 0.5-μm spacing. Time-lapse images were obtained at an interval of 60 s.

FRAP experiments in Figs. 3 and 4 were performed on a confocal microscope (LSM 510; Carl Zeiss, Inc.) equipped with a 63×/1.4 Plan-Apochromat oil immersion objective. A 1.12-μm² circular region of the ring was bleached with a sequence of 10 high-intensity laser iterations and then pictures were taken every 1.5 (Cdc15-GFP and Ric1-GFP) or 3 s (GFP-Myo2 and YFP-Rng2). Fluorescence intensities were analyzed using S10 LSM software (Carl Zeiss, Inc.), and values were normalized to a region within the ring of a nonbleached cell and the background to correct for overall bleaching. Mid1-GFP FRAP was performed on a spinning disk confocal microscope (UltraView LCI) equipped with a 100×/1.40 Plan-Apochromat oil immersion objective and Micropoint laser ablation system (Photonic Instruments). After bleach of a 2.84-μm² circular region of the Mid1-GFP ring, z-series optical sections were captured with a charged-coupled device camera (Orcar E) at 0.5-μm spacing every 3 min. Normalized data were plotted using Prism 4.0c software (Graphpad Software, Inc.). Mobile fraction and half-time values were calculated from the best-fit curve equation as the difference in final and bleach intensity and the time to reach half of the final intensity after photobleaching, respectively. Student’s t test and standard error were calculated to determine significant differences. Differences reported as significant have p-values <0.05 and nonoverlapping standard error intervals.

Online supplemental material

Proteins were resolved by SDS-PAGE, followed by Coomassie blue staining or Western blot analysis with anti-GST or anti-MBP to visualize proteins.

In vitro binding assays

All recombinant bacterially produced proteins were purified on either glutathione beads (GST) or amylose beads (MBP) as previously described (Carnahan and Gould, 2003). Excess binding buffer consisted of 20 mM Tris-HCl, pH 7.4, 150 mM NaCl, 2 mM EDTA, and 0.1% Triton X-100.
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