

The formins Cdc12 and For3 cooperate during contractile ring assembly in cytokinesis

Valerie C. Coffman,¹ Jennifer A. Sees,³ David R. Kovar,^{3,4} and Jian-Qiu Wu^{1,2}

¹Department of Molecular Genetics and ²Department of Molecular and Cellular Biochemistry, The Ohio State University, Columbus, OH 43210

³Department of Molecular Genetics and Cell Biology and ⁴Department of Biochemistry and Molecular Biology, The University of Chicago, Chicago, IL 60637

Both de novo-assembled actin filaments at the division site and existing filaments recruited by directional cortical transport contribute to contractile ring formation during cytokinesis. However, it is unknown which source is more important. Here, we show that fission yeast formin For3 is responsible for node condensation into clumps in the absence of formin Cdc12. For3 localization at the division site depended on the F-BAR protein Cdc15, and *for3* deletion was synthetic lethal with mutations that cause defects in contractile ring formation. For3 became essential in cells expressing N-terminal truncations

of Cdc12, which were more active in actin assembly but depended on actin filaments for localization to the division site. In tetrad fluorescence microscopy, double mutants of *for3* deletion and *cdc12* truncations were severely defective in contractile ring assembly and constriction, although cortical transport of actin filaments was normal. Together, these data indicate that different formins cooperate in cytokinesis and that de novo actin assembly at the division site is predominant for contractile ring formation.

Introduction

Cytokinesis is the final step of the cell division cycle that partitions cellular components into two daughter cells. The contractile ring containing actin filaments, myosin II, and many other proteins is required for cytokinesis in fungi and animal cells (Balasubramanian et al., 2004; Barr and Gruneberg, 2007; Pollard and Wu, 2010). Both de novo actin assembly at the division site and cortical transport/flow contribute actin filaments to the contractile ring (White and Borisy, 1983; Bray and White, 1988; Cao and Wang, 1990; Lee et al., 1998; Pelham and Chang, 2002; Chen et al., 2008; Zhou and Wang, 2008; Alsop et al., 2009; Huang et al., 2012; Subramanian et al., 2013). However, the relative importance of these sources of actin filaments for the contractile ring is unknown in any cell type. Mechanisms of actin accumulation at the division site hold the key to understanding different models for cleavage site selection and contractile ring assembly. However, determining the contributions from de novo assembly and cortical transport has been difficult because of the overlap between the two mechanisms (Zhou and Wang, 2008; Huang et al., 2012).

The fission yeast *Schizosaccharomyces pombe* is an excellent model organism for investigating molecular mechanisms

of cytokinesis (Roberts-Galbraith and Gould, 2008; Laporte et al., 2010). The anillin-like protein Mid1 is essential for division site specification (Chang et al., 1996; Sohrmann et al., 1996; Bähler et al., 1998a; Paoletti and Chang, 2000; Celton-Morizur et al., 2004; Almonacid et al., 2009, 2011). Mid1 acts as a scaffold and positional cue to assemble IQGAP Rng2, myosin-II Myo2 and its light chains Cdc4 and Rlc1, F-BAR protein Cdc15, and the formin Cdc12 into cytokinesis nodes and then the contractile ring (Coffman et al., 2009; Almonacid et al., 2011; Laporte et al., 2011; Padmanabhan et al., 2011; Lee and Wu, 2012). Search, capture, pull, and release (SCPR) is a stochastic ring assembly model whereby actin filaments nucleated in random directions by Cdc12 are captured by myosin-II motors in neighboring nodes to pull them together into the contractile ring (Vavylonis et al., 2008; Lee et al., 2012). The SCPR model describes ring assembly by de novo actin nucleation without considering cortical transport of actin filaments. The ring remains at a constant diameter as it matures by addition of more proteins during anaphase B (Wu et al., 2003). After anaphase, the contractile ring begins to disassemble as it constricts.

Correspondence to Jian-Qiu Wu: wu.620@osu.edu

Abbreviations used in this paper: FH, formin homology; fps, frames per second; Lat-A, latrunculin A; MT, microtubule; SCPR, search, capture, pull, and release; SPB, spindle pole body; ts, temperature-sensitive; wt, wild type.

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Formins are a family of conserved proteins that nucleate and elongate linear actin filaments (Castrillon and Wasserman, 1994; Evangelista et al., 2002; Pruyne et al., 2002; Sagot et al., 2002a,b; Kovar et al., 2003; Pring et al., 2003). All formins contain a highly conserved formin homology (FH) 2 domain that forms a stable homodimer with an actin binding surface for nucleation and processive barbed end association (Moseley et al., 2004; Xu et al., 2004) and an FH1 domain with proline-rich tracts to bind and rapidly elongate profilin-actin (Wasserman, 1998; Kovar et al., 2003; Li and Higgs, 2003; Romero et al., 2004; Kovar, 2006; Vavylonis et al., 2006; Neidt et al., 2009; Courtemanche and Pollard, 2012). GTPase binding domains and FH3 domains existing in some formins are involved in localization or activation (Petersen et al., 1998; Carnahan and Gould, 2003; Gorelik et al., 2011; Liu et al., 2012). Some formins remain bound to barbed ends of growing actin filaments for >1,000 s in vitro (Kovar and Pollard, 2004; Kovar, 2006), sufficient to make a >30- μ m filament, whereas actin filaments average only \sim 1 μ m in vivo (Karpova et al., 1998; Kamasaki et al., 2005; Coffman et al., 2009), suggesting that formin activity must be tightly regulated. Many formins are regulated by autoinhibition when the two ends of the protein interact to prevent actin nucleation. Inhibition is relieved by Rho GTPase binding, phosphorylation, or localization to cortical nucleation sites (Takeya et al., 2008; Wang et al., 2009; Ramalingam et al., 2010; Buttery et al., 2012; H. Chen et al., 2012; Maiti et al., 2012). In addition, inhibitors regulate nucleation activity or processivity of some formins by displacing them from barbed ends of actin filaments (Eisenmann et al., 2007; Quinlan et al., 2007; Chesarone et al., 2009).

S. pombe uses three formins for distinct actin structures: Fus1 assembles actin filaments for the mating projection (Petersen et al., 1998); For3 assembles actin filaments within long actin cables for cellular transport and polarization (Feierbach and Chang, 2001b); and Cdc12 is essential to assemble the contractile ring for cytokinesis (Chang et al., 1997). For3 has established roles in cell polarity (Feierbach and Chang, 2001b; Martin and Chang, 2006; Martin et al., 2007; Wang and Vavylonis, 2008; Scott et al., 2011), but it has not been studied in the contractile ring. For3 localizes to the division site but is not essential for cytokinesis (Pelham and Chang, 2002; Martin and Chang, 2006). The prevailing assumption is that For3 localization to the division site is to mark the new cell end (Feierbach and Chang, 2001a; Kamasaki et al., 2005). Cdc12 assembles actin filaments both from cytokinesis nodes at the division site (Coffman et al., 2009) and from nonmedial locations (Huang et al., 2012). Most knowledge of Cdc12 targeting and regulation stems from structure-function analyses of Cdc12 and F-BAR protein Cdc15. An N-terminal fragment of the first \sim 500 aa localizes normally to the contractile ring in the presence of native full-length Cdc12 and binds to Cdc15 (Carnahan and Gould, 2003; Yonetani et al., 2008). The regulation of Cdc12 activity remains a mystery because mutational analyses provided no obvious evidence of autoinhibition (Yonetani et al., 2008). However, the formation of interphase contractile rings when the C terminus is deleted indicates an activated allele, suggesting autoinhibition may occur by an unknown mechanism (Laporte et al., 2010; Yonetani and Chang, 2010).

Here we investigate the relative importance of different sources of actin filaments for the contractile ring in fission yeast cytokinesis by studying the formin For3 and N-terminal truncations of Cdc12. The Cdc12 truncations depend on actin and For3 for localization to the division site. Despite the truncations nucleating more actin filaments throughout the cell, without proper Cdc12 localization for de novo actin assembly at the division site, contractile ring assembly is inefficient and unreliable. Collectively, these results reveal a novel role for For3 at the division site and the cooperation between two formins for successful cytokinesis.

Results

Condensation of cytokinesis nodes without the essential formin Cdc12

Actin filaments are required for condensation of cytokinesis nodes (defined as 20–65 punctate structures at the cell equator containing formin Cdc12, myosin-II heavy chain Myo2, regulatory light chain Rlc1, and other proteins) into the contractile ring during early mitosis (Wu et al., 2006; Huang et al., 2008; Saha and Pollard, 2012). The formin Cdc12, a nucleator of linear actin filaments, is essential for cytokinesis (Chang et al., 1996, 1997; Kovar et al., 2003). Instead of remaining dispersed, Myo2 nodes condensed into clumps (hereafter we refer to partially condensed nodes as clumps) but not a contractile ring in temperature-sensitive (ts) *cdc12-112* cells at the restrictive temperature of 36°C (Fig. 1 A), similar to Rlc1 (Hachet and Simanis, 2008). It was suggested that clump formation might arise from the ts mutants being less efficient at actin filament nucleation or elongation at the restrictive temperature, resulting in fewer or shorter actin filaments at the division site (Ojkic and Vavylonis, 2010). To investigate the mechanism of node movement in the mutants, we first determined if the ts formins are defective in cortical localization. After 2 h at 36°C, Cdc12-112 and Cdc12-299 concentrated to the nodes or clumps, whereas wild-type (wt) Cdc12 localized to nodes or rings (Fig. 1 B). Thus, improper Cdc12 localization is not the main contributor to defective node condensation.

The *cdc12-112* mutant (Chang et al., 1997) has a point mutation resulting in a substitution of phenylalanine to serine at residue 1355 (Fig. 1 C) in the last α helix of the FH2 domain (Xu et al., 2004). The *cdc12-299* mutant (Chang et al., 1996) has a similar mutation, F1361S (Fig. 1 C), as well as a deletion of three proline residues (948–950) in the FH1 domain. Both point mutations in the FH2 domain of Cdc12 alter residues that are highly conserved in evolutionarily diverse formins (Xu et al., 2004). To determine if the mutations affect actin nucleation/elongation activities, we purified wt and mutant Cdc12 FH1FH2 from *Escherichia coli* and tested their actin assembly properties in vitro. Unlike wt FH1FH2, the mutants were not capable of stimulating the assembly of actin or profilin-actin (Fig. 1, D–G). Although these assays were performed at a permissive temperature of 25°C, the purified proteins' inability to function is probably reflective of their state at the restrictive temperature because cells are normally viable at room temperature. To rule out the possibility that residual activity of the mutant Cdc12 at

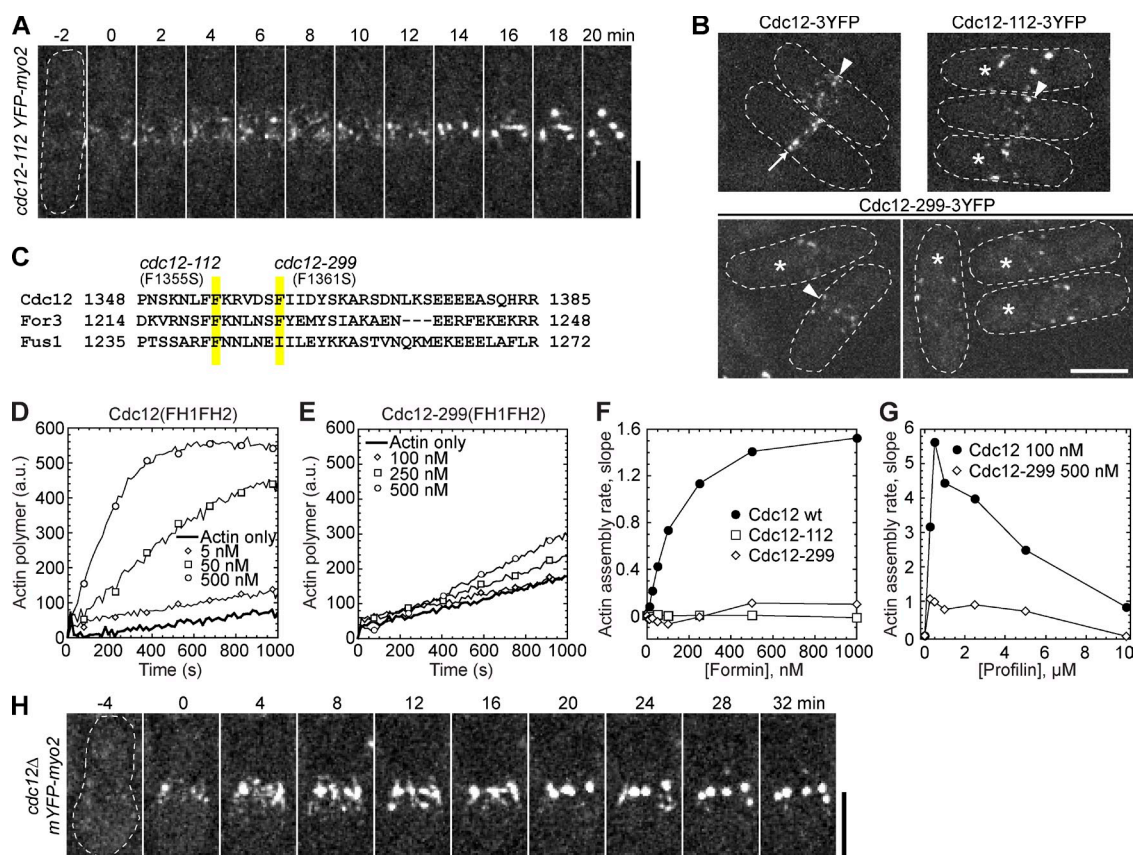


Figure 1. Condensation of cytokinesis nodes without functional formin Cdc12. (A and B) Cells were grown at 36°C for 2 h before imaging at 36°C. (A) Time course of Myo2 nodes condensing into clumps in *cdc12-112* mutant. Time 0 is when nodes appear. (B) Localization of Cdc12, Cdc12-112, and Cdc12-299. Nodes (arrowheads), contractile rings (arrows), and cells with scattered clumps (asterisks) are marked. (C) Alignment of the last α -helix of the FH2 domains (α T; Xu et al., 2004) of *S. pombe* formins. Positions of point mutations in *cdc12-112* and *cdc12-299* are highlighted. (D–G) In vitro spontaneous assembly of 2.5 μ M Mg-ATP actin monomers (10% pyrene labeled) using purified FH1-FH2 domains from wt and mutant Cdc12, in the absence (D–F) and presence (G) of profilin. The data are single representative experiments out of at least two repeats. (D and E) Time course of actin polymerization with the increasing concentrations of wt (D) or mutant (E) Cdc12. (F) Dependence of the actin assembly rate on formin concentration. (G) Dependence of Cdc12-mediated actin assembly rate on profilin concentration. (H) Time course of condensation of Myo2 nodes in *cdc12Δ* cells. Spores from *cdc12Δ/cdc12+* diploid strain were germinated and grown at 25°C for 12 h before imaging at 25°C. Time 0 is when nodes appear. Bars, 5 μ m.

36°C is responsible to condense the nodes into clumps, we deleted one copy of *cdc12* in a diploid strain (Laporte et al., 2011). Surprisingly, we found that germinated *cdc12Δ* spores also condensed nodes into clumps instead of a contractile ring (Fig. 1 H). These data from *cdc12* mutants reveal that other actin nucleators also function at the division site during contractile ring assembly.

Involvement of the formin For3 in cytokinesis

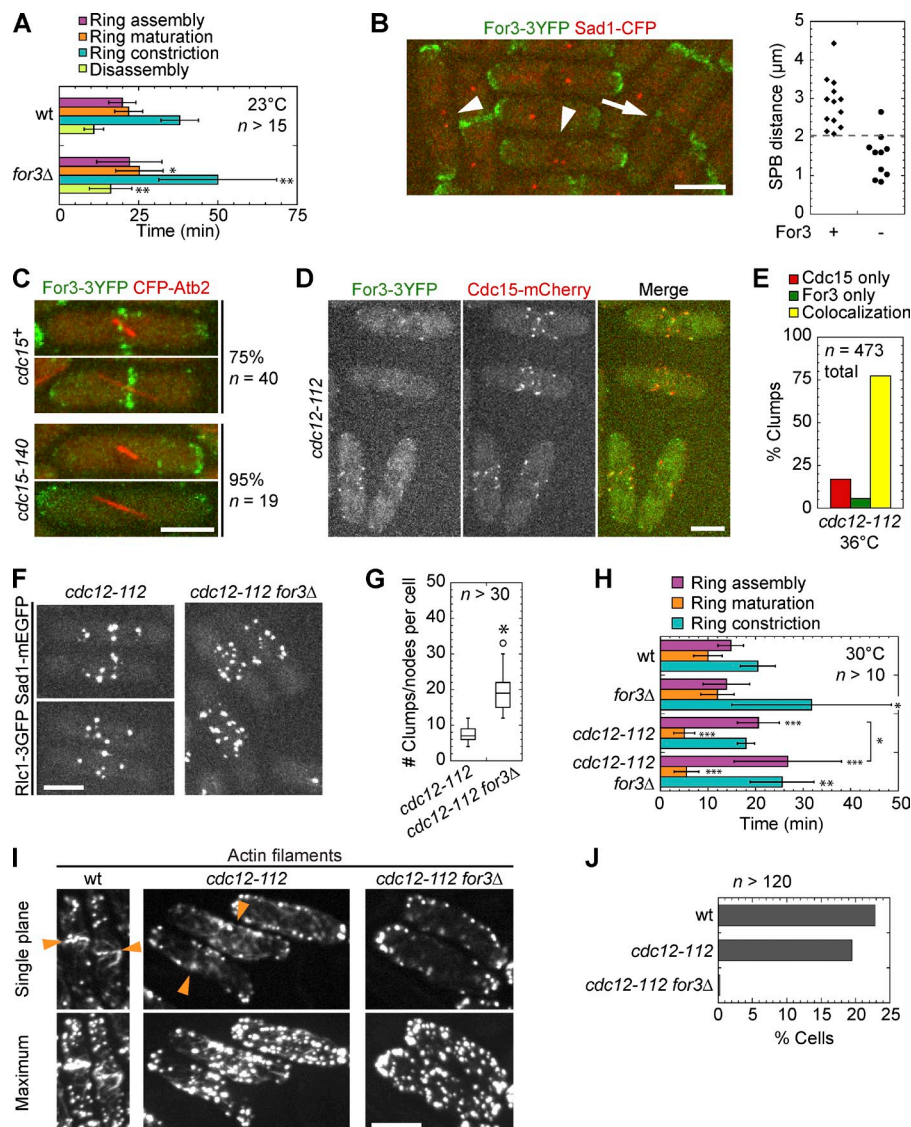
The formin For3 is the nucleator of linear actin filaments in actin cables (Feierbach and Chang, 2001b). It has been reported that the Arp2/3 complex, the nucleator of branched actin filaments, participates in the formation of actin rings at the division site and that the actin ring is not detected in fixed *arp3-c1 for3Δ* double mutant cells (Pelham and Chang, 2002). However, YFP-actin, which is used by the Arp2/3 complex but not by formins in fission yeast, is not visible in the contractile ring (Wu et al., 2006; Q. Chen et al., 2012). We examined live cells expressing Rlc1-3GFP and spindle pole body (SPB) protein Sad1-mEGFP in *for3Δ*, *arp3-c1*, or the *for3Δ arp3-c1* double mutant. After shifting cells to 19°C for 6–16 h, the restrictive temperature for *arp3-c1*,

both single and double mutant cells formed rings normally, despite polarity defects in the mutants (Fig. S1, A and B). These data suggest that neither the Arp2/3 complex nor For3 is required for contractile ring assembly in a wt *cdc12* background.

Although the timing of ring assembly was not significantly different in *for3Δ* and wt cells, later stages of ring constriction and disassembly were slowed in *for3Δ* (Fig. 2 A), consistent with its relocalization to the septation site during ring constriction (Fig. S1 C). However, we found that *for3Δ* was synthetic lethal with *rng2* and *cdc4* mutants, two genes essential for contractile ring assembly (Table 1), suggesting that For3 might play an overlapping role in contractile ring assembly with other proteins. Consistently, For3 arrived at the division site shortly after SPB separation during ring assembly and usually before the spindle began to elongate during anaphase B (Fig. 2 B). This localization did not depend on actin (Fig. S2 A) because For3 still localized to the division site during ring assembly after Latrunculin A (Lat-A) treatment or in the actin-binding mutant *for3-1930A* that is defective for actin assembly (Martin and Chang, 2006; Scott et al., 2011). Of the cytokinesis mutants tested, only *cdc15-140* affected For3 localization to the division site (Fig. 2 C

Figure 2. The formin For3 localization at the division site and its involvement in cytokinesis.

(A) Timing (mean \pm SD) of cytokinesis in *for3Δ* and wt at 23°C. *, $P < 0.05$; **, $P < 0.005$ compared with wt. This and subsequent p-values were determined using a two-tailed *t* test. Here and in later figures, ring assembly is from Rlc1 node appearance until a compact ring forms; ring maturation is from a compact ring to the start of ring constriction; ring constriction is from the start of constriction until the ring becomes a dot; and disassembly is from a dot until all the signal disappears from the division site. (B) For3 (green) localization to the division site. Arrowheads indicate cells with two SPBs (red) but no For3, whereas the arrow indicates a cell with two SPBs and For3. Graph (right) depicts 3D SPB distance in cells with short (<5 μ m) spindles with (+) and without (–) For3 at the division site. (C–G) Cells were grown at 36°C for 2 h before imaging at 36°C. (C) For3 (green) localization in wt and *cdc15-140* cells expressing CFP-Atb2 (red). Cells with similar spindle lengths are shown. Percentage of cells (with 2–7- μ m spindles) with (wt) or without (*cdc15-140*) For3 at the division site is given. (D) For3 and Cdc15 localization in clumps in *cdc12-112* mutants. (E) Quantification of clumps/nodes in (D) with Cdc15 only, For3 only, or colocalization of For3 and Cdc15. (F) Node condensation is impaired in *cdc12-112 for3Δ* double mutants compared with *cdc12-112*. (G) Quantification of clumps/nodes per cell in images as in F, subtracting two SPBs from the total. *, $P < 0.0001$ compared with *cdc12-112*. Box plots include 50% of the data inside the box, the line is the median, the whiskers indicate upper and lower bounds, and the circles outside the whiskers are outliers. (H) Timing of contractile ring stages (mean \pm SD) in indicated Rlc1-3GFP labeled strains grown at 30°C for 2 h before imaging at 30°C. *, $P < 0.05$; **, $P < 0.005$; ***, $P < 0.0005$ in *t* tests compared with wt (except where the bracket indicates). (I) Staining of actin filaments in wt, *cdc12-112*, and *cdc12-112 for3Δ* cells. Cells were fixed at 36°C after growing at 36°C for 2 h. Single planes near the top of the cells and maximum intensity projections are shown. The meshwork/ring of filaments at the division site is marked (arrowheads). (J) Quantification of cells as in I. Percent cells with actin meshwork and rings (wt only) at the division site. Bars, 5 μ m.



and Fig. S2, B and C). Proteins that affect For3 localization to cell tips or interact with For3 at tips were not required for its localization to the division site (Fig. S2 D). For3 localized to the division site later than F-BAR protein Cdc15 in wt cells (Wu et al., 2003), but it colocalized with Cdc15 in clumps in *cdc12-112* cells at 36°C, although at the elevated temperature 3YFP is more difficult to detect (Fig. 2, D and E). Double mutant *cdc12-112 for3Δ* cells had significantly more but dimmer node-like structures at the division site in cells with two nuclei than in single mutant *cdc12-112* cells (Fig. 2, F and G), suggesting that actin filaments nucleated by For3 might be responsible for node movement and node condensation into clumps. Indeed, at a semipermissive temperature of 30°C, ring assembly was significantly delayed in *cdc12-112 for3Δ* compared with *cdc12-112* (Fig. 2 H). Consistently, meshwork-like actin filaments existed at the cell

equator in *cdc12-112* cells, similar to a previous study (Arai and Mabuchi, 2002), but not in *cdc12-112 for3Δ* double mutants (Fig. 2, I and J). Together, these data suggest that For3 and Cdc12 may work together to nucleate actin filaments for the contractile ring during cytokinesis.

N-terminal truncations of Cdc12 affect actin cytoskeleton organization

To further explore the relationship between the formins Cdc12 and For3 during cytokinesis, we tested whether altering Cdc12 activity or localization by truncating Cdc12 would make For3 essential. We truncated the first 503 aa of Cdc12 (Δ 503-Cdc12; Fig. 3 A) to eliminate the binding site for Cdc15 (Carnahan and Gould, 2003) and another localization domain (Yonetani et al., 2008). We also made a truncation that only contained FH1, FH2,

Table 1. Genetic interactions of *for3Δ* with cytokinesis mutants

Mutant ^a	Viable double mutants at 25°C	Viable single mutants at 25°C ^b	Total number of tetrads
	%	%	
<i>cdc12-299</i>	100	100	11
<i>rng2-D5</i>	0	100	10
<i>cdc4-8</i>	0	100	9
<i>rlc1Δ</i>	36	100	8
<i>cdc15-140</i>	100	100	11
<i>mid1-366</i>	100	100	11

^aParent strains are given in Table S1.^bSingle mutants of *for3Δ* were 100% viable on tetrad plates in this group of crosses. Some ts mutants had been backcrossed before these crosses.

and the C terminus ($\Delta 841$ -Cdc12; Fig. 3 A). The $\Delta 841$ -Cdc12 fragment can rescue *cdc12-112* or *cdc12Δ* when overexpressed from a high-copy plasmid (Yonetani et al., 2008). Both N-terminal truncations were expressed from the *cdc12* promoter at its native locus replacing the wt Cdc12. Cells with the truncations resembled wt cells. The truncations localized to node-like structures and contractile rings at the division site, although fewer nodes were detectable than in wt (Fig. 3 B and Video 1). We used fluorescence intensity to estimate the expression of the truncations (Wu and Pollard, 2005; Coffman et al., 2011; Laporte et al., 2011).

$\Delta 503$ -Cdc12 levels in the whole cell and the contractile ring were not significantly different from those in wt cells; however, the levels of $\Delta 841$ -Cdc12 were reduced to $\sim 60\%$ in the whole cell and $\sim 15\%$ at the contractile ring (Fig. S3 A). The contractile ring assembled slightly faster in both truncations, whereas ring constriction was delayed (Fig. 3 C and Video 2).

Next we asked if their actin assembly activities are altered. Actin staining revealed a significant increase in total actin filaments (Fig. 3, D and E), whereas actin patch (Arp2/3 complex-nucleated, branched actin filaments) intensities were unchanged (mean actin patch intensities were $\sim 97\%$ of wt in both truncations; $n \geq 50$ patches each). In addition, actin cables per cross section increased from 3.3 ± 1.3 in wt to 5.0 ± 1.1 and 4.9 ± 1.2 in $\Delta 503$ -*cdc12* and $\Delta 841$ -*cdc12* cells (mean \pm SD; Fig. 3 E). Thus, $\Delta 841$ -Cdc12 even at the reduced level is sufficient to form excess actin cables to the same extent as $\Delta 503$ -Cdc12. Together, these data show that the Cdc12 truncations are activated alleles for actin assembly and are sufficient for contractile ring formation.

For3 is essential for contractile ring assembly and constriction in Cdc12 truncations

Surprisingly, unlike full-length Cdc12 (Wu et al., 2003, 2006), localization of both truncations to the division site

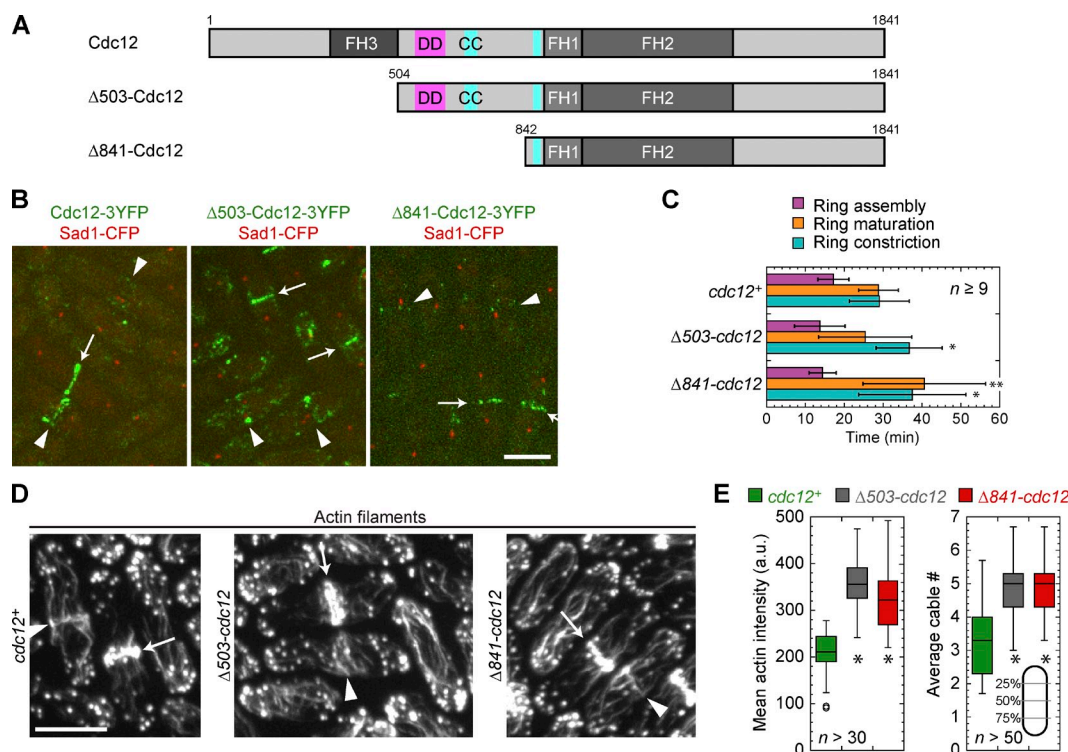
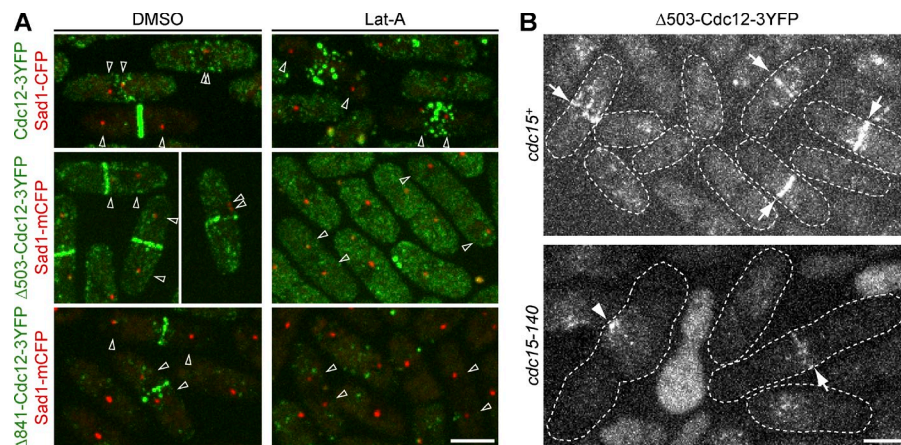


Figure 3. N-terminal truncations of Cdc12 affect actin cytoskeleton organization. (A) Domain architecture of full-length Cdc12 and two truncations, approximately to scale. $\Delta 503$ -Cdc12 and $\Delta 841$ -Cdc12 delete the first 503 aa and the first 841 aa, respectively. FH domains 1, 2, and 3 are in darker shades of gray. Putative dimerization domain (DD, pink) and coiled-coil domains (CC, teal) are shown (Yonetani et al., 2008). (B) Localization of N-terminal truncations of Cdc12 (green) and SPB protein Sad1 (red) as a cell cycle marker. Localizations to ring precursors (arrowheads) and rings (arrows) are shown. (C) Rlc1-3GFP ring assembly, maturation, and constriction (mean \pm SD) in Cdc12 N-terminal truncations and wt. *, $P < 0.05$; **, $P < 0.005$. (D and E) Staining of actin filaments in wt, $\Delta 503$ -*cdc12*, and $\Delta 841$ -*cdc12* cells to reveal an actin meshwork at the division site (arrowheads), actin rings (arrows), and more and brighter actin cables in the mutants. (E, left) Box plots of cell size-corrected integrated actin intensity from the sum intensity projection of 0.4- μ m-spaced sections. (right) Box plots of actin cable numbers per cross section in interphase cells. Cables were counted in three line scans of the maximum intensity projection at the positions indicated (gray lines) and the mean number from those three positions for each cell is plotted. *, $P < 0.0001$ compared with wt. Bars, 5 μ m.

Figure 4. N-terminal truncations of Cdc12 depend on actin and Cdc15 for localization. (A) Cdc12 N-terminal truncations or wt were treated with DMSO or Lat-A for 30 min. Cells with similar SPB distances are marked by open arrowheads for comparison. (B) Localization of $\Delta 503$ -Cdc12 to the contractile ring/forming ring (arrows) or clump (arrowhead) in wt or *cdc15-140* after growth at 36°C for 4 h and then imaged at 36°C. Bars, 5 μ m.



was abolished after Lat-A treatment (Fig. 4 A). We hypothesized that For3-nucleated actin filaments are crucial for the localization of the Cdc12 truncations. To test this hypothesis, we abolished For3 localization to the division site using *cdc15-140* (Fig. 2 C). The *cdc15-140* $\Delta 503$ -*cdc12* cells were viable at 25°C but synthetic lethal from 30 to 36°C (Table 2). After 4 h at 36°C, $\Delta 503$ -Cdc12-3YFP localized weakly to a contractile ring in 1% of living cells in *cdc15-140*, compared with 18% in wt (Fig. 4 B; $n > 100$ cells each). Some cells also had large $\Delta 503$ -Cdc12 clumps at the division site. The perturbed localization is not likely to be a direct effect of the *cdc15* mutation because the Cdc12 truncation lacks the Cdc15 binding site.

Crossing the Cdc12 truncations to *for3* Δ yielded no visible double mutant colonies despite 80–100% single mutant and wt spores forming colonies (Fig. 5 A and Table 2; $n > 40$ tetrads for each cross), indicating that For3 is essential in these Cdc12 truncations. Because neither $\Delta 503$ -*cdc12* *for3* Δ nor $\Delta 841$ -*cdc12* *for3* Δ double mutants could be recovered from the

crosses, we observed tetrads 2–4 d after dissection to determine the lethal defects (Fig. 5, B–H), using a confocal microscope with an automated stage (tetrad fluorescence microscopy; see Materials and methods). Many double mutants died with only one or two elongated cells ($\sim 30\%$; Fig. 5 B), suggesting a cytokinesis defect. Although we found colonies of up to 32 cells, most had 3–8 cells ($\sim 52\%$; Fig. 5 C). We mainly imaged the double mutants with 3–8 cells because the long cells had likely failed cytokinesis. Because *for3* Δ cells have polarity and spindle orientation defects (Feierbach and Chang, 2001b), we used CRIB-3GFP and mRFP-Atb2 as markers for polarized growth and the mitotic spindle, respectively, to observe tetrad colonies. The CRIB domain from *Saccharomyces cerevisiae* Cdc42 effector Gic2 is a marker for active Cdc42 (Tatebe et al., 2008; H. Chen et al., 2012). Although cell polarization was affected in the double mutants based on abnormal cell shapes, the majority of interphase cells had one or two cell tips with CRIB localization (Fig. 5 D and Video 3). Mitosis occurred in $\sim 40\%$ of double

Table 2. Genetic interactions of $\Delta 503$ -*cdc12* with cytokinesis mutants

Mutant ^a	Viable double mutants at 25°C	Viable single mutants at 25°C ^b	Total number of tetrads	Lowest synthetic lethal temperature ^c
	%	%		°C
<i>for3</i> Δ	0	90	42	N/A
<i>for3-1930A</i>	0	42	43	N/A
<i>mid1</i> Δ	0	64	18	N/A
<i>rng2-D5</i>	7	89	11	30
<i>rng2-346</i>	0	88	11	N/A
<i>cdc4-8</i>	0	64	10	N/A
<i>myo2-E1</i>	5 ^d	82	19	N/A
<i>rlc1</i> Δ	0	93	11	N/A
<i>rng3-65</i>	100	100	10	30
<i>cdc15-140</i>	60	80	10	30
<i>mid1-6</i>	50	86	14	ND
<i>mid1-366</i>	67	100	9	ND
<i>blt1</i> Δ	93	100	10	ND

^aParent strains are given in Table S1.

^bSingle mutants of $\Delta 503$ -*cdc12* were 93% viable on tetrad plates in this group of crosses. Ts single mutants were more viable after backcrossing; see Table 1.

^cTested by streaking wt, single mutants, and double mutant on YE5S plates and incubating at temperatures from 25 to 36°C. The $\Delta 503$ -*cdc12* single mutants grew normally from 25 to 36°C.

^dThe only viable double mutant isolated likely had a suppressor based on cell morphology at 36°C.

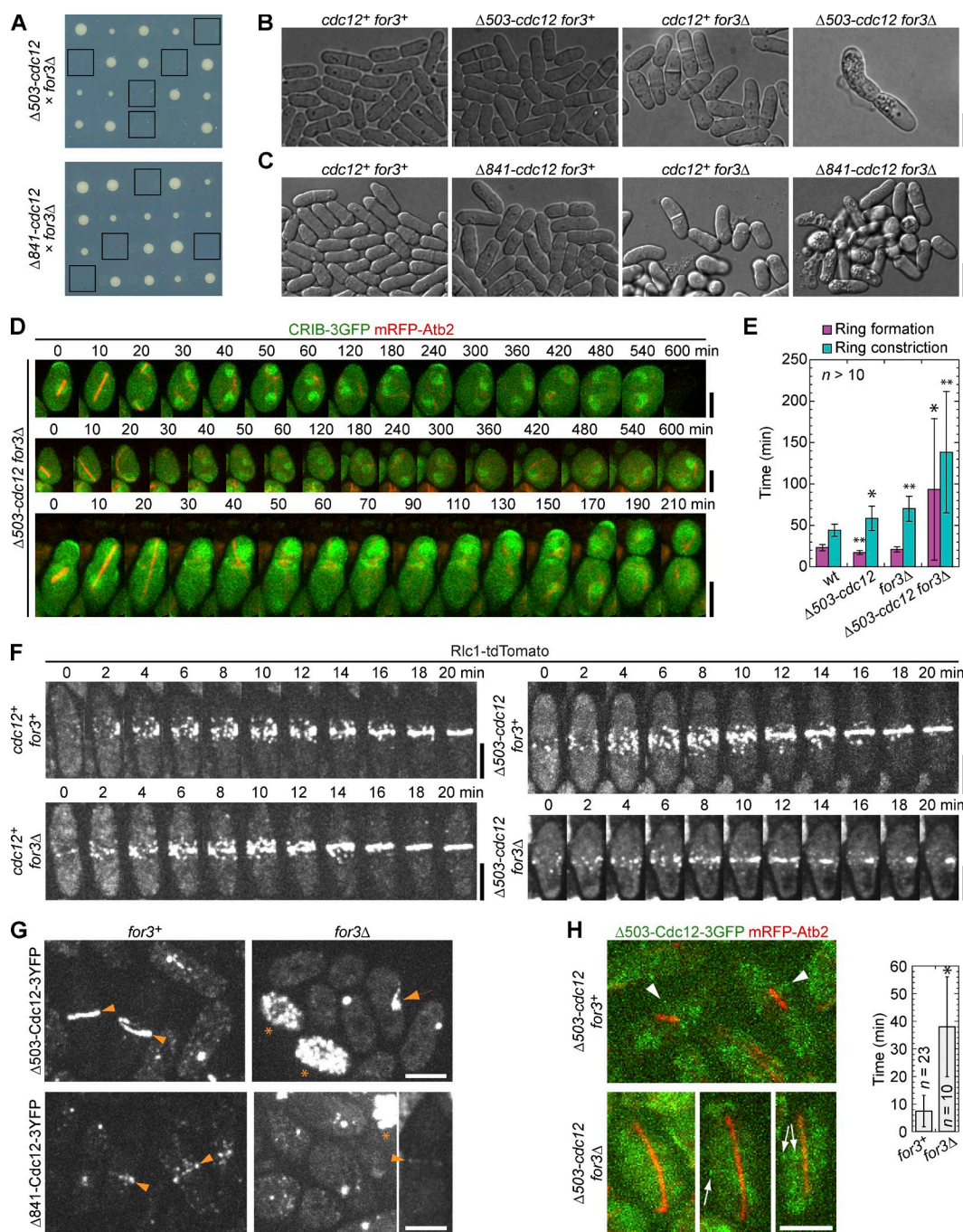


Figure 5. Tetrad fluorescence microscopy reveals that *for3Δ cdc12* truncation double mutants are defective in contractile ring assembly and constriction. (A) Sections of tetrad plates from crosses between *for3Δ* and *cdc12* truncations grown at 25°C. Double mutants from each tetrad (vertical) are boxed. (B) Differential interference contrast images of a tetratype tetrad from the *for3Δ* × *Δ503-cdc12* cross (JW1665 × JW5124) 3 d after tetrad dissection. The entire colony of the double mutant is shown, while the wt and single mutants are focused at the edge of the colonies where the cells are in a single layer. (C) Same as B for *for3Δ* × *Δ841-cdc12* cross (JW5281 × JW5283). (D) Time courses showing three *Δ503-cdc12 for3Δ* CRIB-3GFP mRFP-*atb2* cells that undergo mitosis during the movies. Time 0 is the presence of a short spindle. Tetrads were dissected from a cross JW5281 × JW5282-2 and imaged on YE5S agar (see Materials and methods) at 23°C. CRIB-3GFP is a marker for cell polarization and mRFP-*Atb2* is a marker for MTs and the spindle. (E and F) *for3Δ Δ503-cdc12* cells are defective in contractile ring assembly and constriction. Tetrad fluorescence microscopy of a cross between *for3Δ rlc1-tdTomato* and *Δ503-cdc12 rlc1-tdTomato*. (E) Timing (mean ± SD) of ring assembly and constriction as shown in F from tetrad colonies. Ring constriction here is from the appearance of a compact ring to ring constriction to a dot (including the ring maturation time). *, *P* < 0.05; **, *P* < 0.005 compared with wt. (F) Time courses of four representative cells from the same tetratype tetrad. (G) For3 is crucial for the localization of Cdc12 N-terminal truncations. Tetrad fluorescence microscopy of crosses between *Δ503-cdc12-3YFP* or *Δ841-cdc12-3YFP* with *for3Δ*. The localizations to rings (arrowhead) and clumps (arrow) are marked. Dead cells with high autofluorescence in *for3Δ* are marked with asterisks. (H) Micrographs (left) of mRFP-*Atb2* spindles and *Δ503-Cdc12-3GFP* localization to faint rings (arrowhead) or clumps (arrow). Graph (right) shows timing (mean ± SD) of *Δ503-cdc12-3GFP* localization to the medial region of the cell after the appearance of a spindle using a 5–10-min delay in tetrad fluorescence microscopy. *, *P* < 0.001. Bars: (D and F–H) 5 μm; (B and C) 10 μm.

mutants during the movies, but less than half of those succeeded in cytokinesis. The low rate of mitosis in 12 h is likely caused by the polarity and growth defects. However, these defects alone are not sufficient to explain the cytokinesis defect, because *arp3-c1 for3Δ* cells were not well polarized but assembled contractile rings normally (Fig. S1 A).

To observe contractile ring assembly directly, we crossed *for3Δ rlc1-tdTomato* to *Δ503-cdc12 rlc1-tdTomato* cells and observed them 2–3 d after tetrad dissection (Fig. 5, E and F). Of the cells that attempted contractile ring formation during imaging, 39% of *Δ503-cdc12 for3Δ* cells successfully assembled a compact ring (Fig. 5 F), but it took approximately three times as long to assemble and constrict the ring compared with wt cells (Fig. 5 E). The single mutants were only slightly delayed in ring constriction, similar to our previous results from liquid cultures (Figs. 2 A and 3 C). In the double mutants, 51% of cytokinesis attempts ($n = 37$) resulted in formation of large Rlc1 clumps at the division site instead of a compact ring (Fig. 5 F and Video 4). The large variation in times to condense the nodes into a compact ring and to constrict the ring in the double mutants suggests stochastic and unreliable cytokinesis pathways without normal formin function or localization. Because Rlc1 localizes to nodes and other upstream node proteins are independent of Cdc12 for localization (Laporte et al., 2011; Padmanabhan et al., 2011), we expect that the nodes only lack Cdc12.

To test whether ring assembly might fail in the double mutants because of defective localization of the Cdc12 truncations to the division site, we crossed tagged truncations to *for3Δ* and observed their localization in the double mutants. Both truncations localized to the division site mainly at a large spot or clump in *for3Δ* cells in contrast to a ring in *for3⁺* cells (Fig. 5 G). The time from spindle appearance to spindle breakdown was similar in wt (38.0 ± 5.6 min, $n = 15$ cells) and *for3Δ Δ503-cdc12* cells (37.9 ± 10.5 min, $n = 14$), indicating that the duration of mitosis is normal. However, in cells that formed spindles, 83% cells formed a *Δ503-Cdc12* spot/clumps at the division site 38.0 ± 18.1 min after spindle appearance in *for3Δ* (no signal in other cells) compared with 100% cells forming a *Δ503-Cdc12* ring after 7.5 ± 5.7 min in *for3⁺* (Fig. 5 H). In two cells that managed to form formin rings out of >40 cells (Fig. 5 G), the ring assembly took >2.5 h. The actin-binding mutant *for3-I930A* was also synthetic lethal with *cdc12* truncations, indicating that actin filament assembly by For3 is essential for the localization of the truncations. The expression level and localization of For3 in *cdc12* truncation strains were similar to wt (Fig. S3, B–D). Thus, the dependence on For3 was not an artifact of abnormal For3 behavior. Together, these data indicate that Cdc12 lacking its known N-terminal localization domains (Carnahan and Gould, 2003; Yonetani et al., 2008) can be recruited to the division site by actin filaments nucleated by For3.

Genetic interactions further support For3's roles in cytokinesis. Both *Δ503-cdc12* and *for3Δ* exhibited synthetic lethal interactions with several cytokinesis mutants involved in contractile ring assembly and constriction including *rng2*, *cdc4*, *myo2*, *rlc1Δ*, and *mid1Δ* (Tables 1 and 2). Thus, For3 has a direct role in contractile ring assembly in cooperation with Cdc12.

Contributions of actin filaments assembled at or transported to the division site for contractile ring assembly

Cdc12 concentrates to cytokinesis nodes from scattered cytoplasmic speckles (Coffman et al., 2009) and For3 relocates from cell tips to the division site (Fig. 2 A) during contractile ring assembly in wt cells. Cdc12 and For3 assemble actin filaments at the division site and at other places in the cell (Fig. 3 D; Martin and Chang, 2006; Coffman et al., 2009; Huang et al., 2012). One open question is the relative contribution of de novo actin filament assembly at the division site and cortical transport/flow of actin filaments assembled elsewhere to the contractile ring (Huang et al., 2012). The actin binding peptide, Lifeact (Riedl et al., 2008), is reported to label actin structures with the least impact on actin dynamics and cell growth (Huang et al., 2012). We compared the timing of ring assembly using Rlc1 as a marker in wt and strains expressing tagged Lifeact or GFP alone from the strong actin promoter. The Lifeact tagged with a GFP that has a tendency to dimerize at high concentration (Yang et al., 1996) doubled the times needed for ring assembly, unlike the monomeric GFP (mGFP L221K)-tagged Lifeact (Zacharias et al., 2002; Huang et al., 2012) or overexpressed GFP alone (Fig. S4 A). We also compared the intensity of the actin meshwork at the cell equator to the regions just outside the broad band (to avoid actin patches concentrated at cell tips) in fixed wt cells stained for actin filaments and in live cells expressing Lifeact-GFP or Lifeact-mGFP during early contractile ring assembly (Fig. S4 B). All three strains were similar with approximately three times more actin filaments at the cell equator than at the adjacent regions. In later stages, cells expressing Lifeact-GFP usually had thick, wavy bundles outside of the compact ring; whereas in wt cells (Fig. S4 C) or Lifeact-mGFP cells (Fig. S4 D and Video 5), we rarely detected prominent nonmedial cables after a compact actin ring had formed. Together these data suggest that the excess cables in Lifeact-GFP-expressing cells are not crucial for contractile ring assembly or maturation (see Discussion).

We detected more actin filaments at the cell equator than at the adjacent regions during contractile ring assembly (Fig. S4 B), but it is challenging to tell whether these filaments are assembled de novo at the division site or transported from elsewhere by cortical flow. We predicted that double mutants of *for3Δ* and *cdc12* truncations are ideal for testing the importance of de novo actin assembly at the division site because no formins could localize to the cell equator efficiently at early mitosis (Fig. 5, G and H). We crossed *for3Δ* and *Δ503-cdc12* cells both expressing Lifeact-mGFP for tetrad fluorescence microscopy. Consistent with previous experiments (Fig. 5), 100% of wt and single mutant cells assembled and constricted the contractile ring with normal timing (Fig. 6 A and Video 6). However, only ~25% of double mutant cells assembled actin rings ($n = 28$ cells), and the timing of assembly and constriction were severely delayed, despite these cells having an excess of actin filaments throughout the cells (Fig. 6 A and Video 6). Because myosin-II marked with Rlc1 still localized to the division site (Fig. 5 F), cortical transport might be normal. Indeed, in tetrad fluorescence microscopy, the timing of actin accumulation at the division site was not significantly different in *Δ503-cdc12* and *for3Δ*

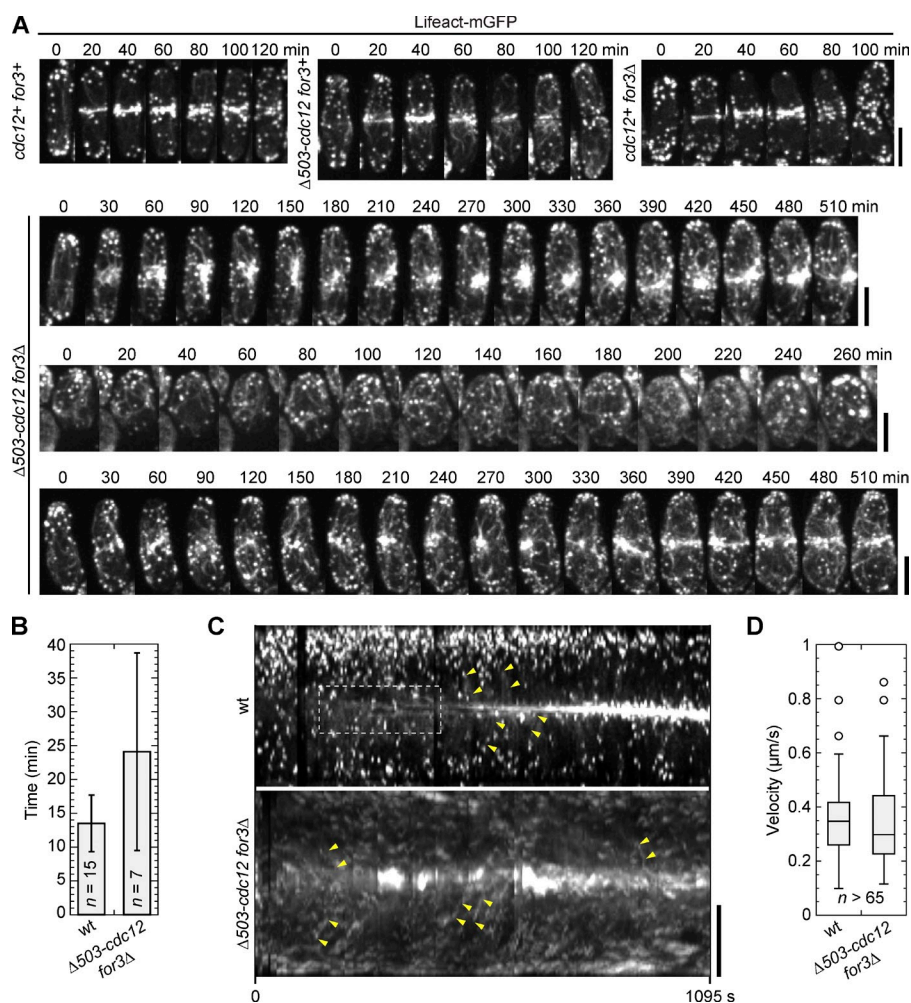


Figure 6. Medial actin filaments are predominant for contractile ring assembly. (A) Time courses of contractile ring assembly and constriction by tetrad fluorescence microscopy in cells expressing Lifeact-mGFP from the cross JW5463 \times JW5481. The interval of the time-lapse movie is 10 min and time 0 is the time point just before actin filaments concentrate near the division site (except for the double mutant with no accumulation at the division site). Three classes of double mutant cells are shown: large clump at the division site (top); no specific structures (middle); and severely delayed ring assembly (bottom). (B) Timing of actin accumulation in a ring/meshwork (wt) or aster ($\Delta 503\text{-cdc12 for3}\Delta$) after Rlc1-tdTomato node appearance in tetrad fluorescence microscopy. Many double mutants could not be included in this quantification because they did not attempt division during the imaging. Error bars show SDs. (C and D) Actin filament movements toward the division site. (C) Kymographs of wt and double mutant ($\Delta 503\text{-cdc12 for3}\Delta$) cells expressing Lifeact-mGFP. De novo actin filament nucleation (dashed box, wt only) and cortical flow (examples are marked by pairs of yellow arrowheads) are evident in tetrad fluorescence microscopy from the same cross as B. (D) Box plots of velocity of actin movements toward the division site as shown in C. Bars, 5 μm .

$\Delta 503\text{-cdc12}$ cells ($P = 0.10$; Fig. 6 B). Of those double mutant cells that formed an actin aster, 43% eventually formed rings. Because $\Delta 503\text{-Cdc12}$ accumulation was delayed (Fig. 5 H), this result suggests that actin accumulates at the division site by cortical transport before $\Delta 503\text{-Cdc12}$ does. The velocity of actin filaments moving toward the division site in wt and $for3\Delta$ $\Delta 503\text{-cdc12}$ was also similar (Fig. 6, C and D; and Video 7). Thus, cortical transport is normal, whereas de novo actin filament assembly at the division site could not occur until after $\Delta 503\text{-Cdc12}$ localized to the clump in $for3\Delta$ $\Delta 503\text{-cdc12}$. These results suggest that although the Cdc12 truncation assembles actin filaments efficiently in $for3\Delta$ cells, cortical transport alone is not effective and reliable for contractile ring formation and constriction. Together, these data indicate that de novo assembly of actin at the medial cortex is crucial for contractile ring assembly and constriction in fission yeast cytokinesis.

Discussion

The formin For3 participates in assembly of actin filaments for the contractile ring and becomes essential in the absence of other pathways to efficiently localize the formin Cdc12 to the division site. In addition, cells that cannot anchor the activated Cdc12 to the division site assemble contractile rings

inefficiently, indicating that de novo nucleated actin filaments at the cleavage site are crucial for contractile ring assembly and constriction.

Cooperation between the formins Cdc12 and For3 in fission yeast cytokinesis

Our data support a previously uncharacterized role of For3 at the division site that is redundant with Cdc12. In the presence of normal Cdc12 function, For3 has little effect on the fidelity of contractile ring assembly and only affects ring constriction. However, when the function of Cdc12 is affected by point mutations or N-terminal truncations, actin assembly by For3 is needed for node movement and cell survival. The lower efficiency of actin nucleation by For3 (Scott et al., 2011) might explain node clumping in $cdc12$ mutants, based on the predictions of the SCPR model (Vavylonis et al., 2008; Ojic and Vavylonis, 2010).

The localization of Cdc12 N-terminal truncations depends on actin and For3, but the mechanism is unclear. There are at least four possibilities for the recruitment of Cdc12 truncations to the division site: (1) Cdc12 displaces For3 from the barbed end of actin filaments at the division site. This is likely because the barbed end affinity of Cdc12 is $\sim 10\times$ higher than that of For3 (Scott et al., 2011) and capping protein localizes to the division site after a compact ring is formed (Wu et al., 2003).

(2) Cdc12 travels on actin filaments. Both For3 and the formin Bni1 are released from actin nucleation sites and incorporate into actin cables by remaining bound to the barbed ends of short bundled filaments (Kamasaki et al., 2005; Martin and Chang, 2006; Buttery et al., 2007). Thus, Cdc12 might travel to the division site by displacing For3 within actin cables. (3) The actin filaments nucleated by Cdc12 truncations could be bundled with those assembled by For3 at the division site, which recruits Cdc12 to the division site. (4) Actin nucleated by the Cdc12 truncations might be captured and pulled into the division site by Myo2 or Rng2 in the nodes, which accumulate there independently of actin filaments and Cdc12 (Wu et al., 2006; Laporte et al., 2011; Padmanabhan et al., 2011). This mechanism could also explain the limited localization of Cdc12 truncations in *for3Δ* to the contractile ring or large clump (Fig. 5, G and H) and might be similar to myosin-II-dependent retrograde flow of actin filaments in budding yeast (Huckaba et al., 2006; Gao and Bretscher, 2009). These four possibilities are not mutually exclusive, but further experiments are needed to distinguish them. We cannot exclude the possibility that For3 has additional roles at the division site beyond localizing Cdc12.

The increase of actin filaments in the Cdc12 N-terminal truncations is consistent with these being activated Cdc12 alleles. A Cdc12 fragment missing the C terminus (downstream of the FH2 domain) is not sufficient for viability, but induces the formation of contractile rings during interphase when overexpressed in the presence of native protein (Yonetani and Chang, 2010). These data suggest that the N and C termini of Cdc12 might be involved in an autoinhibitory interaction, but the regulation of such an interaction remains a mystery.

Although formins often have distinct functions (Kovar et al., 2011; Scott et al., 2011), cooperation or redundancy between them is also common. Mouse formins mDia1 and mDia2 have both been implicated in cytokinesis (Tominaga et al., 2000; Watanabe et al., 2008). In budding yeast, both formins Bni1 and Bnr1 localize to the bud neck and are involved in cytokinesis (Vallen et al., 2000). In addition, cells can survive with only one formin (Pruyne et al., 2004; Gao and Bretscher, 2009), suggesting functional redundancy. Most remarkably, budding yeast can polarize and divide when only the FH1FH2 fragment of one formin is expressed in *bni1Δ bnrΔ* (Gao and Bretscher, 2009). We find some differences in fission yeast, where Cdc12 N-terminal truncations alone are not viable in *for3Δ* and Cdc12 is essential for cytokinesis, which might reflect the fact that the contractile ring is more important for cytokinesis in cells with a larger cleavage site diameter (Bi et al., 1998; Tolliday et al., 2003; Laporte et al., 2010).

Sources of actin filaments for contractile ring assembly during cytokinesis

One of the outstanding and challenging questions in fungal and animal cytokinesis is the relative importance of the sources of actin filaments for the contractile ring (Zhou and Wang, 2008; Huang et al., 2012). The defect in *for3Δ cdc12* truncation double mutants and our analysis of actin dynamics labeled by Lifeact can address this important question. Our data reveal for the first time the effect of mislocalizing Cdc12 but not upstream

proteins in node and contractile ring assembly pathways (Almonacid et al., 2011; Laporte et al., 2011; Padmanabhan et al., 2011). These activated alleles of Cdc12 are efficient at nucleating actin filaments throughout the cell (Figs. 3 D and 6 A), but when not anchored properly to the division site, severe cytokinesis defects emerge. These data indicate that actin filaments assembled at nonmedial sites are not efficient for contractile ring assembly even when the cortical transport is normal. It is important to note that the lethality of the *for3Δ cdc12* truncation double mutant might not arise solely from the contractile ring assembly defect, because polarity defects, slow growth, and cell lysis were also observed.

An interesting question is whether de novo actin assembly or cortical transport alone is sufficient for contractile ring formation. Cdc12 truncation cells have more actin cables (~3–7 per cross section) than wt cells (Fig. 3 E) but most *cdc12* truncation *for3Δ* mutants form large aggregates of actin and other ring components instead of a contractile ring (Fig. 5, F and G; and Fig. 6 A). The inefficiency of cortical transport for ring assembly might be a result of the asymmetric distribution of actin cables in contrast to symmetric distribution of cytokinesis nodes, the sites of de novo assembly at the division site. Collectively, eliminating de novo actin assembly in the presence of normal cortical transport during early stages of contractile ring formation results in failure of contractile ring assembly in most cells, suggesting that de novo actin assembly at the division site is more crucial than cortical transport in fission yeast cytokinesis. Indeed, after Cdc12 accumulates at the clump by cortical transport, de novo actin nucleation might also contribute to the successful ring assembly in some cells.

The actin promoter used for the Lifeact constructs is a strong constitutive promoter (Huang et al., 2012). High levels of Lifeact expression in plants altered actin filament dynamics (van der Honing et al., 2011). Although Huang et al. (2012) demonstrated that Lifeact-GFP colocalized with actin staining in the same cells, our data suggest that Lifeact-GFP might promote the formation of aberrant actin structures that inhibit contractile ring assembly (Fig. S4). Using the concentration of actin in fission yeast (63.2 μM; Wu and Pollard, 2005) and the dissociation constants measured for YFP and mYFP dimers (0.11 mM and 9.7 mM, respectively; Zacharias et al., 2002), we estimate that the concentration of Lifeact-GFP dimer may be as high as 12.8 μM, whereas Lifeact-mGFP dimer is only 0.4 μM in the cells with the respective Lifeact constructs. The range of dissociation constants given by different measurements for Lifeact and F-actin is 1.3 to 2.3 μM and for G-actin is 40 to 280 nM (Riedl et al., 2008). The combination of an actin-binding motif and a dimerization motif is a feature shared by actin filament nucleators such as formins and bundlers such as α-actinin. Thus, dimerization of GFP at high concentrations might transform the Lifeact-GFP peptide into an actin filament nucleator or bundler because of its high affinity with actin, which may result in thick actin cable formation throughout the cell.

In conclusion, our results revealed the contributions of different sources of actin to the contractile ring. Both formins Cdc12 and For3 nucleate actin filaments at the division site, with the essential Cdc12 playing the major role. Nonmedial

filaments nucleated by activated Cdc12 truncation alleles are unable to efficiently assemble a contractile ring when formin localization to the division site is impaired. Together, these data suggest that de novo actin nucleation at the division site by Cdc12 and For3 is the most important source of actin for contractile ring assembly in fission yeast. It will be interesting to know if de novo actin nucleation at the cleavage furrow is also the predominant source of actin filaments for the contractile ring in other cell types.

Materials and methods

Strains, growing conditions, and genetic, molecular, and cellular methods

Table S1 lists the *S. pombe* strains used in this study. All tagged genes are integrated at their native chromosomal loci under the control of endogenous promoters unless noted otherwise (Bähler et al., 1998b). Cells were grown in exponential phase for 36–48 h before microscopy as described previously (Wu et al., 2006). Spore germination for *cdc12* deletion was performed as described previously (Laporte et al., 2011). In brief, one copy of *cdc12* was deleted using the KS-ura4⁺ marker (Bähler et al., 1998b) in a diploid strain with two copies of the *mYFP-myo2*. The resulting strain was grown for 2 d on SPA5S to induce sporulation, and then tetrads were incubated 12 h in 2% glucuronase at 25°C to kill remaining diploids and release the spores. Spores were washed 5× in 1.7 g/liter of rinsing medium (Yeast Nitrogen Base; BD) and germinated in YE5S-ura liquid medium 12–24 h before imaging. Because fission yeast diploid cells are not stable for long-term storage, only the parental strains are listed in Table S1. Lat-A treatments were performed at a final concentration of 100 μM (Wu et al., 2001) and imaging was done on bare slides to maintain the drug concentration.

The N-terminal truncations of *cdc12* at the native locus were made by first cloning the *cdc12* promoter (−800 to +3 bp from the ATG) into the KS-ura4 vector at the multicloning sites upstream of the *ura4* promoter (Bähler et al., 1998b) to obtain plasmid JQW213. A *cdc12* truncation strain was constructed with an mYFP tag at the N terminus under control of the *81nmt1* promoter and selected by G418 resistance (Bähler et al., 1998b). Then the *ura4⁺-Pcdc12* region of plasmid JQW213 was amplified for targeting to the *cdc12* locus to replace the *kanMX6-P81nmt1-mYFP* portion, selected for growth on EMM-uracil, and screened for loss of YFP signal. The resulting *ura4⁺-Pcdc12* truncation strains were then tagged at the C terminus with 3YFP. In some truncation strains, the *ura4⁺* marker was looped out during crosses because it resided between 800 bp of identical *cdc12* promoter sequence. Control strains were matched to the same auxotrophic state of the truncations for quantifications. Sequencing of the mutants was done by cloning Cdc12 ORF into plasmids and sequencing the plasmids in both directions for duplicate coverage.

For actin staining, 300 U BODIPY-phalloidin 488 (Molecular Probes) was dissolved in methanol at a final concentration of 0.2 U/μl, and then dispensed in 25-μl aliquots and dried in a vacuum in the dark and stored at −20°C. 1 ml of log phase cells was fixed by adding 333 μl of 16% paraformaldehyde (Electron Microscopy Sciences) at the growing temperature for 5–15 min, washed in 1 ml PEM buffer (0.1 M Na Pipes, pH 6.8, 1 mM EGTA, and 1 mM MgCl₂) 3×, permeabilized with 1% Triton X-100 in PEM buffer for 2 min, and washed in 1 ml PEM buffer 3×. Each phalloidin aliquot was resuspended in 50 μl PEM buffer and 10 μl was combined with 10 μl of permeabilized cells. After ~30 min the stained cells were washed once in 1 ml PEM buffer and then centrifuged at 7,000 rpm for 30 s to collect cells. Most of the supernatant was removed and ~5 μl of cell suspension was added to a gelatin slide for imaging.

Plasmid constructs, protein purification, and actin assembly assays

wt and mutant FH1FH2 domain constructs (residues 882–1390) were cloned between BamHI and XhoI (wt and Cdc12-112) or BamHI and NotI (Cdc12-299) restriction sites into the bacterial expression vector pET21a, under control of the IPTG-inducible T7 promoter (EMD Millipore; Kovar and Pollard, 2004). Cdc12-112 (F1355S) was cloned by QuikChange site-directed mutagenesis (Agilent Technologies). Cdc12-299 (Δ948–950; F1361S) was cloned by amplification of genomic DNA and Infusion (Takara Bio Inc.). wt and mutant Cdc12(FH1FH2)-His proteins were purified as described previously (Kovar and Pollard, 2004; Scott et al., 2011). Constructs were expressed in *E. coli* strain BL21-Codon Plus (DE3)-RP (Agilent Technologies) with 0.5 mM IPTG for 16 h at 16°C, broken with an EmulsiFlex-C3 (Avestin) in extraction buffer (50 mM NaH₂PO₄, pH 8.0, 500 mM NaCl, 10% glycerol, 10 mM imidazole,

and 10 mM β-mercaptoethanol supplemented with protease inhibitors), clarified by spinning at 30,000 and 50,000 g for 15 and 30 min, incubated with Talon Metal Affinity Resin (Takara Bio Inc.) for 2 h at 4°C, and then loaded onto a disposable column. After a 50-ml wash with extraction buffer, formin was eluted with 250 mM imidazole in extraction buffer. Pure protein was dialyzed with formin buffer (20 mM Hepes, pH 7.4, 1 mM EDTA, 200 mM KCl, 0.01% Na₂S₂O₈, and 1 mM DTT), concentrated, flash frozen in liquid nitrogen, and stored at −80°C. Ca-ATP actin was purified from chicken skeletal muscle acetone powder by extraction with G Buffer (2 mM Tris, pH 8.0, 0.2 mM ATP, 0.1 mM CaCl₂, and 0.5 mM DTT) and a cycle of polymerization and depolymerization (Spudich and Watt, 1971), gel filtered on Sephacryl S-300 in G Buffer, and labeled on Cys-374 with pyrenyliodoacetamide (Life Technologies). *S. pombe* profilin Cdc3 was expressed and purified from bacteria by binding to poly-L-proline Sepharose, elution with 7.0 M urea, and dialysis in profilin buffer (20 mM Tris, 150 mM KCl, and 0.2 mM DTT, pH 7.5; Kovar et al., 2003). The spontaneous assembly of actin was measured from the fluorescence of 10% pyrene-actin with a Safire 2 fluorescence plate reader (Tecan) as described in detail previously (Scott et al., 2011). In brief, pyrene-labeled Mg-ATP-actin monomers were placed in an upper row of a 96-well black plate (Corning). Cdc12, profilin, and 10× KMEI (500 mM KCl, 10 mM MgCl₂, 10 mM EGTA, and 100 mM imidazole, pH 7.0) were placed in a lower row. Reactions were initiated by mixing contents of the lower wells with actin monomers in the upper wells.

Microscopy and data analysis

Cells for microscopy were collected from liquid cultures as described previously (Wu et al., 2006; Coffman et al., 2009). In brief, they were centrifuged at 5,000 rpm, washed into EMM5S, placed on a thin layer of EMM5S liquid medium with 20% gelatin (Sigma-Aldrich) and 25 nM *n*-propyl-gallate, and observed at 23–24°C except where noted. Figs. 4 A and 5 D were taken from single stacks or time-lapse imaging at 23°C in a Delta TPG dish (Bioptechs) with YE5S agar on top of the cells to maintain a single cell layer. For imaging of ts mutants at 36°C, cells were collected and placed on EMM5S + 2% agar pads prewarmed for 10 min at 36°C. Precautions were taken to maintain cells at 36°C using a benchtop incubator (Boekel) and an Objective Heater (Bioptechs; Laporte et al., 2011). Imaging in Figs. 2 C and 4 B was done in a prewarmed Delta TPG dish from a small incubator with prewarmed YE5S agar placed on top of the cells. The dish was transferred to the preheated imaging chamber (Stage Top Incubator [INUB-PPZ12-F1] equipped with dish holder [UNIV2-D350; Tokai Hit]) on the microscope. Microscopy at 19°C was done by cooling the entire room to 19°C. Cells were grown at 19°C in a heated water bath in a 4°C cold room and centrifuged in the cold room before imaging.

We imaged cells using 100×/1.4 NA Plan-Apo objective lenses (Nikon) on two microscopes. For most imaging, we used a spinning disk confocal microscope (UltraVIEW ERS; PerkinElmer) with 440, 568-nm solid state lasers and 488, 514-nm argon ion lasers and a cooled charge-coupled device camera (ORCA-AG; Hamamatsu Photonics) on an Eclipse (TE2000-U; Nikon) with 2 × 2 binning. Actin staining was imaged with 1 × 1 binning. Some imaging (Fig. 2, C and D; Figs. 4–6; and Fig. S4, A, B, and D) was performed on another spinning disk confocal microscope (UltraVIEW Vox CSUX1 system; PerkinElmer) with 440-, 488-, 515-, and 561-nm solid state lasers and back-thinned EMCCD camera (Hamamatsu Photonics) on a Ti-E inverted microscope (Nikon) without binning.

Tetrad fluorescence microscopy was performed as follows on a Ti-E inverted microscope equipped with the UltraVIEW Vox spinning disk confocal microscope system. The cross was performed as normal on SPA5S for 2 d, followed by tetrad dissection on YE5S agar plate during which spores are placed 5 mm apart in both X and Y directions. Imaging was done after 2–4 d of colony growth at 25°C and genotypes were determined by cell morphology compared with previous crosses for synthetic interaction. Before imaging, the tetrad dissection needle was used to mark the agar around the double mutant colonies to enable us to find them using the 100× objective lens. For tetrad imaging before even the wt colonies were visible to the naked eye, the plate and the agar were marked at the first and last colony in each tetrad as a guide. A section of the agar with two to four tetrads was then cut out of the tetrad plate and inverted onto a large coverslip (24×60-1.5; Thermo Fisher Scientific) that was clamped into place on an automated stage on the microscope (Nikon). The marks in the agar were used to focus and to assist in finding the first colony. Subsequent colonies were found using the X-Y stage position based on the 5-mm distance of colony positions. Each imaging position was saved in the Volocity software so that they could be revisited for imaging at each time point. Tetrads were imaged with 2–10-min delays, and the autofocus drive in the software was used to ensure that the focus was maintained for ~12 h of imaging. For viable genotypes, cells at

the edges of the colonies were imaged, whereas the entire colony was imaged for the double mutants. Time-lapse images of fluorescence and differential interference contrast were acquired at each time point for each colony. The parent strains are listed in the strain table, because the double mutants could not be isolated and kept. The tetrad fluorescence microscopy we developed here is a promising new approach to determine the cause of cell death in synthetic lethal double mutants by following the progeny of a meiotic event. Using both the tetrad microscope and a sensitive fluorescence microscope, this imaging technique enabled several possible causes of cell death to be distinguished with much greater certainty and resolution than traditional random spore assays.

Maximum-intensity projections and other image analyses were performed using UltraVIEW, Velocity, and/or ImageJ. Images in figures are maximum-intensity projections of z-sections spaced at 0.2–0.8 μm except where noted. Two tagged proteins were considered in the same nodes if their centers were within one radius of the nodes. Images in Fig. 5 H are of 3GFP-tagged truncation instead of 3YFP, and thus the GFP signal is not as brightly visible above the higher autofluorescence at 488 nm. Rotated regions in Fig. S1 C were scaled to 400% in ImageJ using interpolation and resized in Adobe Illustrator to match the scale bar of the maximum intensity projection. In Fig. S4 B, cells with partially condensed actin meshworks at the division site were excluded from our analysis. Statistical analysis was done using Microsoft Excel, and *t* tests were performed using the “*t* test” function in Excel, with two tails and type 2 (homoscedastic) or 3 (heteroscedastic) as appropriate. Given numbers (*n*) in the figures are for each bar/box unless otherwise noted.

Analysis of actin filament movement was done using ImageJ as described previously (Huang et al., 2012) on ~20-min movies taken with 5-s delay, visualizing only a single slice through the cell. Up to half the cell width was used to make a kymograph along the long axis of the cell, and the angle and the distance traveled by actin filaments that moved toward the division site was measured. The velocity was calculated from these measurements based on the distance traveled and the time between frames. The kymographs in Fig. 6 C cover the entire cell width.

Supplemental videos were made with the QuickTime movie writer plug-in in ImageJ with Video Compression and Best Quality, with key frame every frame (Video 1) or every 5–10 frames. Videos 1 and 7 were scaled to 400% in ImageJ using interpolation. Side-by-side images were automatically adjusted for brightness and contrast the same way so the intensities are comparable for data collected on the same day (Videos 1–4, 6, and 7), whereas the left and right portions of Video 5 were taken on different days and might not be comparable.

Counting numbers of molecules in whole cells and at the division site

Cells were imaged at 0.4- μm z spacing, the full width at half maximum of the point spread function on our system (Coffman et al., 2011; Laporte et al., 2011; Coffman and Wu, 2012), and measurements were taken using the sum intensity of all sections with signal (Wu and Pollard, 2005; Wu et al., 2008). wt cells in the same field were measured the same way and intensity was subtracted to remove autofluorescence and background from the measurement. Global intensity of Cdc12 truncations tagged with 3YFP was compared with global intensity of Cdc12-3YFP, setting Cdc12 to 600 molecules per cell on average and then adjusting for cell size (Wu and Pollard, 2005). The division site intensity was measured using a rectangle region of interest over the ring area and a 2 \times larger rectangle along the cell's long axis to determine the background (Wu et al., 2008).

Online supplemental material

Table S1 lists the *S. pombe* strains used in this study. Fig. S1 shows contractile ring formation in *arp3-c1 for3 Δ* cells at 19°C. Fig. S2 shows For3 localization to the division site independent of the cytokinesis and polarity proteins. Fig. S3 shows Cdc12 truncations and For3 expression levels and For3 localization in *cdc12* truncation strains. Fig. S4 shows the characterization of Lifeact-mGFP and Lifeact-GFP. Video 1 shows contractile ring assembly from a band of Cdc12 nodes or node-like structures in wt and $\Delta 503\text{-cdc12}$. Video 2 shows contractile ring assembly from a broad band of Rlc1-3GFP nodes in wt and *cdc12* truncations. Video 3 shows tetrad fluorescence microscopy from a cross between *for3 Δ* and $\Delta 503\text{-cdc12}$ cells expressing CRIB-3GFP and mRFP-Atb2 as markers for active Cdc42 and microtubules (MTs). Video 4 shows tetrad fluorescence microscopy from a cross between *for3 Δ* and $\Delta 503\text{-cdc12}$ cells expressing Rlc1-tdTomato as a marker for cytokinesis nodes and the contractile ring. Video 5 shows the visualization of Lifeact-GFP and Lifeact-mGFP during contractile ring assembly. Video 6 shows Lifeact-mGFP in tetrad fluorescence microscopy from a cross between *for3 Δ* and $\Delta 503\text{-cdc12}$. Video 7 shows actin filaments moving toward the division site

in cells attempting cell division. Online supplemental material is available at <http://www.jcb.org/cgi/content/full/jcb.201305022/DC1>.

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