Inhibition of cell membrane ingression at the division site by cell wall in

fission yeast

Ting Gang Chew¹²*, Tzer Chyn Lim¹*, Yumi Osaki³, Junqi Huang¹, Anton Kamnev¹, Tomoyuki
Hatano¹, Masako Osumi³⁴, Mohan K. Balasubramanian¹

Affiliations

¹Division of Biomedical Sciences, Warwick Medical School, University of Warwick, Gibbet Hill Rd,
Coventry, West Midlands, CV4 7AL, United Kingdom.
²ZJU-UoE Institute, Zhejiang University School of Medicine, International Campus, Zhejiang
University, 718 East Haizhou Road, Haining, Zhejiang 314400, P.R. China.
³Integrated Imaging Research Support, 1-7-5-103, Hirakawa-cho, Chiyoda-ku, Tokyo 102-0093,
Japan.
⁴Laboratory of Electron Microscopy/Bio-imaging Center, Japan Women's University, 2-8-1
Mejirodai, Bunkyo-ku, Tokyo 112-8681, Japan.

*Co-first authors

#Correspondence to: Mohan K. Balasubramanian (m.k.balasubramanian@warwick.ac.uk)

Total Characters: 19999
Abstract

Eukaryotic cells assemble an actomyosin ring during cytokinesis to function as a force-generating machine to drive membrane invagination, and to counteract the intracellular pressure and the cell surface tension. How the extracellular matrix affects actomyosin ring contraction has not been fully explored. While studying the *S. pombe* 1,3-β-glucan-synthase mutant *cps1*-191, which is defective in division septum synthesis and arrests with a stable actomyosin ring and, we found that weakening of the extracellular glycan matrix caused the generated spheroplasts to divide at the non-permissive condition. This non-medial slow division was dependent on a functional actomyosin ring and vesicular trafficking, but independent of normal septum synthesis. Interestingly, the high intracellular turgor pressure appears to play minimal roles in inhibiting ring contraction in the absence of cell wall remodeling in *cps1*-191 mutants as decreasing the turgor pressure alone did not enable spheroplast division. We propose that during cytokinesis, the extracellular glycan matrix restricts actomyosin ring contraction and membrane ingression, and remodeling of the extracellular components through division septum synthesis relieves the inhibition and facilitates actomyosin ring contraction.

Introduction

Animal cells and fungal cells require assembly and contraction of an actomyosin ring during cytokinesis. In fission yeast, the actomyosin ring contracts to drive membrane ingression, and coordinates with the septum assembly machinery to deposit cell wall materials at the division site (Ramos *et al.*, 2019). The fungal cell wall has been suggested as a functional equivalent of the extracellular matrix (ECM) in animal cells (Munoz *et al.*, 2013). The division septum is a special wall structure composed of primary and secondary septa. The primary septum is a structure that must be degraded to permit cell separation and the secondary septum is a structure that forms the cell wall once both cells are separated. The septum assembly machinery consists of α-glucan synthase Ags1 and β-glucan synthase Bgs1 and Bgs4. The β-glucan synthase Cps1/Bgs1 is essential for primary septum formation. Cps1 synthesizes specifically the linear β-glucan matrix of the primary septum at the division site and couples the extracellular glycan matrix to the
actomyosin ring via intermediate protein complexes (Munoz et al., 2013; Cortes et al., 2015; Davidson et al., 2016; Sethi et al., 2016). The β-glucan synthases Bgs4 and the α-glucan synthase Ags1 are primarily involved in the secondary septum formation and participate in the synthesis of primary septum (Garcia Cortes et al., 2016). The deposition of extracellular glycan matrix coordinates with actomyosin ring contraction and stabilizes the contracting actomyosin ring at the division site (Munoz et al., 2013; Arasada and Pollard, 2014).

How the extracellular glycan matrix influences actomyosin ring contraction (apart from its roles in ring stability during cytokinesis) has not been examined closely (Mishra et al., 2012; Munoz et al., 2013). In this study, we used the thermosensitive allele of β-1,3-glucan synthase, cps1-191 to address this question. The cps1-191 mutant is defective in β−1,3-glucan and septum synthesis and arrests with a non-contracting actomyosin ring at the non-permissive temperature (Liu et al., 2000). Interestingly, we found that weakening of the extracellular glycan matrix in cps1-191 mutant at the non-permissive temperature has enabled actomyosin ring contraction and membrane ingression.

Results and discussion

Under the restrictive temperature, the β-glucan synthase mutant cps1-191 assembles actomyosin rings that do not contract (Liu et al., 2000). It has been suggested that β-glucan synthesis at the division site is required to overcome the high intracellular turgor pressure during cytokinesis, and that the actomyosin ring may not be able to overcome the high turgor (Proctor et al., 2012). To test if the turgor pressure inhibited ring contraction in cps1-191 mutants, we cultured cps1-191 cells in EMMA medium containing 0.8 M sorbitol to decrease the turgor pressure to that inside the cells at the restrictive temperature, and we added 2-deoxyglucose (2-DG) to this culture to prevent further glucan synthesis at the division site and elsewhere in the cell (Megnet, 1965; Svoboda and Smith, 1972; Osumi et al., 1998). A recent study showed that rings in cps1-191 mutant cells constricted slowly after shifting to the restrictive temperature for ~2 hours prior to microscopy at the restrictive temperature (Dundon and Pollard, 2020). To ensure a highly penetrant phenotype for cps1-191, we shifted the cps1-191 cells to the restrictive temperature for ~6 hours prior to microscopy, which
was performed at the restrictive temperature. As previously reported, actomyosin rings of \textit{cps1}-191 cells maintained in normal turgor pressure did not undergo contraction (Figure 1A). We occasionally observed that parts of the \textit{cps1}-191 cells swelled into a bump and the cells lysed eventually with a collapsing ring (Figure 1B). Culturing \textit{cps1}-191 cells in EMMA medium containing 0.8 M sorbitol, did not increase actomyosin ring contraction events and phenotypically these cells resembled \textit{cps1}-191 grown under normal growth conditions in EMMA medium, in which a high intracellular turgor pressure is maintained (Figure 1C and 1D). Thus, our results showed that a decreased turgor pressure does not allow ring contraction in \textit{cps1}-191 mutant cells.

Next, we considered the possibility that the extracellular glycan matrix inhibited ring contraction and membrane ingression in \textit{cps1}-191 mutants in the absence of cell wall remodeling. The Cps1 is a transmembrane protein that (along with other integral membrane proteins, such as Ags1 and Bgs4) links actomyosin rings underneath the cell membrane to the extracellular glycan matrix (Cortes et al., 2005; Cortes et al., 2012; Munoz et al., 2013; Arasada and Pollard, 2015; Davidson et al., 2016; Sethi et al., 2016; Martin-Garcia et al., 2018). It was possible that in the absence of division septum synthesis (and thereby cell wall remodeling), the actomyosin rings are stably fixed to the inactive \textit{cps1}-191 gene-product or other integral membrane proteins (such as mok1, sbg1, and bgs4) that link the cell wall to the actomyosin ring. To test if this was the case, we weakened the extracellular glycan matrix by treating the \textit{cps1}-191 cells with cell wall lytic enzymes and further blocking new cell wall and septum synthesis by supplementing the culture with 2-DG. Interestingly, upon weakening of the cell wall, myosin rings in \textit{cps1}-191 mutant expressing the regulatory light chain of myosin tagged with the fluorescent protein tdTomato (Rlc1-tdTomato) underwent contraction coupled with membrane ingression at the restrictive temperature of 36 °C (Figure 2A; GFP-tagged Syntaxin-like protein Psy1 was used as a cell membrane marker; n = 19/29 spheroplasts). Consistently, contracting actin rings labeled with the Lifeact-mCherry were also detected in the \textit{cps1}-191 mutant upon weakening of cell wall, suggesting that the actomyosin rings were driving the contraction and membrane ingression (Figure 2B; n = 5/41 spheroplasts). These mutant spheroplasts with weakened cell wall often divided non-medially into two, and the rings contracted at much reduced rate ($0.061 \pm 0.021$ μm/min, $n_{\text{spheroplast}} = 8$) compared to wild-type cells.
(0.299 ± 0.059 µm/min, n_cell = 14) (Figure 2C). The slow rate of ring contraction is comparable to that of in the wild-type spheroplasts in which the rings slide along the cell membrane during ring contraction (0.046 ± 0.031 µm/min, n_wild-type spheroplast = 40). We frequently observed that the rings contracted till mid-phase of division and disassembled before completion of cytokinesis. The spheroplasts however went on to divide into two entities (Figure 2A and 2B). The segregation of daughter nuclei in the cps1-191 spheroplasts was often not coordinated with the cytokinesis, with some spheroplasts having two nuclei in one of the daughter entities or have cleaved nuclei, presumably due to the non-medial division (Supplementary figure 1). The mis-coordination of cytofission and nuclear division spatially could arise from the variable dumb-bell shaped morphology of cps1-191 spheroplasts (Mishra et al., 2012). The functions of Mto1 and Mto2, which are involved in the assembly of post-anaphase microtubule arrays may also contribute to this mis-coordination defects (Dundon and Pollard, 2020). Since the cps1-191 mutant spheroplast division was morphologically different from normal fission yeast cell division and was mimicking the morphological changes of some animal cells during division, we have called this type of division as cytofission.

Analysis of the extracellular glycan matrix using calcofluor staining (a division septum-specific fluorochrome) (JC et al., 2018) in cells undergoing cytofission in EMMA containing sorbitol and 2-DG medium revealed that the division site of cps1-191 spheroplasts undergoing cytofission with 2-DG medium contained significantly reduced β-glucan materials (Figure 2D). Further study with the high-resolution scanning electron microscopy showed that the glucan fibrils regenerated in cps1-191 spheroplasts without 2-DG (Figure 2E; bottom panel) while the fibrils were not noticeable in cps1-191 spheroplasts with 2-DG (Figure 2E; top panel). The glucan fibrils commonly present at the division site of fission yeast was largely absent in cps1-191 spheroplasts undergoing cytofission (Supplementary figure 2). Taken together, we showed that weakening of cell wall in cps1-191 cells at non-permissive temperature and ensuing further inhibition of new cell wall and septum synthesis with 2-DG facilitates a novel cytofission event that leads to division of one spheroplast into two in the absence of detectable division-septum growth. Our results also suggested that the extracellular glycan matrix anchored to the actomyosin rings negatively
regulates the ring contraction and membrane ingression. This is consistent with a previous finding that the absence of the Bgs4-synthesized β-glucan in the septum promoted a faster ring contraction and membrane ingression than that of normal septa, and at the same time, the synthesis and ingression of septum wall progressed slower than that of a normal septum (Munoz et al., 2013).

A reduction of β-glucan may result in an increased amount of α-glucan in the cell wall of fission yeast. To test if the cytofission of cps1-191 spheroplasts were due to the synthesis of α-glucan at the division site, we prepared the cps1-191 mok1-664 double mutant spheroplasts containing the thermosensitive alleles of both α- and β- glucan synthases, and imaged the myosin rings and cell membrane in this double mutant spheroplasts at the non-permissive temperature. Similar to the cps1-191 spheroplast, the cps1-191 mok1-664 double mutant spheroplasts underwent cytofission (Figure 3A, n = 11/26), suggesting that α-glucan and β-glucan synthesis did not contribute significantly to the cytofission events.

Normal fission yeast cells that just complete ring contraction and membrane ingression are not entirely separated until the primary septum digestion of the division septum connecting the two newly-divided cells (Sipiczki, 2007). This process is achieved in fission yeast through the action of endoglucanases (Martin-Cuadrado et al., 2003; Dekker et al., 2004; Garcia et al., 2005). We tested if proteins involved in the separation of fission yeast cells were also involved in the cytofission, which would be expected if trace amounts of division septum had been deposited during ring contraction. To this end, we constructed double mutant spheroplasts of cps1-191 lacking the endoglucanases eng1 (β-glucanase) and agn1 (α-glucanase), respectively. Similar to the single mutant cps1-191, the double mutants lacking either of the two endoglucanases underwent cytofission upon weakening of the cell wall (Figure 3B, n = 8/23; Figure 3C, n = 11/38). The results indicated that the cytofission events of cps1-191 mutants does not require the break-down of cell wall materials by endoglucanases, even though cytofission leads to the complete separation of spheroplasts.
In ~53% of the cps1-191 spheroplasts (53 out of 99 spheroplasts) that underwent cytofission, the rings contracted till mid-phase of division and disassembled before division into two entities. We tested if the ESCRT abscission complex was involved in the cytofission by removing two of the ESCRT proteins Vps4 and Vps20 in the cps1-191 spheroplasts. The cps1-191 vps4Δ and cps1-191 vps20Δ double mutant spheroplasts underwent cytofission like in the single cps1-191 mutant spheroplast (Supplementary figure 3A, n = 14/14; Supplementary figure 3B, n = 8/8). It is possible that the completion of cytofission without the actomyosin rings was achieved via an unknown cell abscission mechanism.

Previous studies suggested under certain circumstances, some eukaryotic cells are able to divide without an actomyosin ring (Proctor et al., 2012; Choudhary et al., 2013; Flor-Parra et al., 2014; Dix et al., 2018; Ramos et al., 2019). To see if the cytofission was driven by contraction of the actomyosin ring, we first perturbed the functions of rings using Latrunculin-A (LatA) to inhibit actin polymerization (Morton et al., 2000; Fujiwara et al., 2018). cps1-191 spheroplasts treated with DMSO underwent ring contraction, membrane ingression, and completed cytofission (Figure 4A; n = 11/16 spheroplasts). By contrast, cps1-191 spheroplasts treated with LatA underwent ring disassembly and failed to divide into two entities or ingressed very slowly resembling dividing cells after long time incubation with LatA (Ramos et al., 2019). Interestingly, the smaller entity retracted into the bigger entity, probably due to the imbalance of intracellular pressures (Figure 4B; n = 27/33 spheroplasts).

Next, we perturbed the myosin component of actomyosin rings by deleting rlc1, the regulatory light chain of myosin II (Le Goff et al., 2000; Naqvi et al., 2000; Pollard et al., 2017), in cps1-191 mutants. It has been shown that the cells lacking rlc1 (rlc1Δ) is cold sensitive and fail to undergo cytokinesis at low temperature, but at high temperatures, the rlc1Δ cells assemble an intact actomyosin ring that contracts normally (Naqvi et al., 2000) (Supplementary figure 4; n = 31/31 cells). We used this differential temperature requirement to test the essentiality of actomyosin ring functions in cytofission. If the actomyosin ring was essential in driving cytofission, the absence of
**rlc1** might render the cells with weakened cell wall unable to undergo cytofission at the high temperature, which is normally permissive for cell division in *rlc1Δ* cells alone (Naqvi *et al.*, 2000). Consistent with the LatA experimental findings, the double mutant *cps1-191 rlc1Δ* with weakened cell wall did not undergo ring contraction at the high temperature (Figure 4C; n = 23/24 spheroplasts), whereas the single mutant of *cps1-191* could undergo cytofission.

Targeted membrane deposition is required in the cytokinesis of fission yeast (Wang *et al.*, 2016; Onwubiko *et al.*, 2019). Next, we tested if targeted membrane deposition at the division site facilitates actomyosin ring contraction in cytofission. When the vesicular trafficking across the endomembrane system was inhibited using brefeldin A in the *cps1-191* spheroplasts, the myosin rings were not able to contract to drive cytofission events (Figure 4D, n = 27/27). This result suggested that addition of cell membrane via targeted membrane trafficking at the division site is required to enable cytofission.

Our study reveals that the extracellular glycan matrix inhibits actomyosin ring contraction in the absence of cell wall remodeling and division septum synthesis. When the inhibition is relieved by experimental treatments like ones reported in this study, or by septum synthesis, the actomyosin ring contracts to drive the membrane ingression. A previous study by Proctor *et al.* analyzed *cps1-191* mutants and explained that the failure of membrane ingression in the mutant was due to a defect in division-septum assembly. The authors also proposed that the high intracellular turgor pressure prevents actomyosin ring contraction in fission yeast (Proctor *et al.*, 2012). We tested this model by lowering the turgor pressure in *cps1-191* mutant cells and found that it was not sufficient to enable membrane ingression in the absence of cell wall remodeling in the *cps1-191* cells. However, the ability of *cps1-191* mutant cells to divide upon weakening of cell wall indicates that the actomyosin ring in *cps1-191* mutant cells is capable of driving membrane ingression even when the division septum assembly is defective. When cell wall remodeling is normal, like in wild-type cells, ring contraction and membrane ingression coordinate with cell wall and septum growth.

The lowering of turgor pressure by sorbitol addition in wild type cells with normal cell wall remodeling may facilitate ring contraction, explaining the findings of Proctor et al (Proctor *et al.*, 2012).
The fact that ring contraction is slower during cytofission however, agrees better with the work of O’Shaughnessy and colleagues, who have proposed that the rate of septum synthesis sets the rate of cytokinesis (Stachowiak et al., 2014). It is possible that our work reveals the highest rate of actomyosin ring contraction when confronted with membrane drag and viscous drag of the cytosol. The slow ring contraction rate could be as a result of a reduced amount of Cps1-191 or cytokinetic proteins at the division site (Cortes et al., 2015). In fission yeasts, the actomyosin ring is probably required at the early phase of cytofission to drive spheroplasts into a dumbbell shape with high curvature. Although we cannot exclude the possibility that residual and undetectable actomyosin structures may facilitate division after seeming actomyosin ring disassembly, recent work suggests other potential mechanisms not involving actomyosin rings or ESCRT in division of dumbbell-shaped vesicles. It has been proposed that the spontaneous curvature in dumbbell-shape lipid vesicles generates constriction forces to induce membrane fission. This leads to the division of a dumbbell-shaped lipid vesicle into two with an increased curvature (Steinkuhler et al., 2020).

The yeast cell wall consists of mainly glycan matrix and glycosylated proteins and has been suggested as a functional equivalent of the extracellular matrix (ECM) in animal cells (Munoz et al., 2013). The mechanical interaction between the cytokinetic actomyosin ring and the ECM is not well understood. A recent study of zebrafish epicardial cells in the heart explants shows the cell-ECM adhesions at the division site. The cell-ECM adhesions lead to the traction forces at the cytokinetic ring that inhibit cytokinesis (Uroz et al., 2019). An early biophysical study also detected a large traction force at the cleavage furrow of the fibroblast cells cultured on an elastic substrate, suggesting an interaction of cytokinetic machinery and ECM (Burton and Taylor, 1997). When the cell-ECM adhesion is enhanced during mitosis, the cleavage furrow ingression is inhibited in the epithelial cells (Taneja et al., 2016). Consistently, our study shows that the anchoring of actomyosin rings to the extracellular glycan matrix that do not undergo remodeling (due to a defective Bgs1) prevents the actomyosin ring contraction and cell membrane ingestion. Weakening of the extracellular glycan matrix, presumably mimicking a decreased cell-ECM adhesion, has enabled cytofission events.
Materials and methods

Yeast strains, medium, and culture conditions

Table S1 lists the *S. pombe* strains used in our study. Standard fission yeast genetic techniques were used to prepare the strains. The rich medium YEA (5 g/l yeast extract, 30 g/l glucose, 225 mg/l adenine) was used to culture cells until mid-log phase at 24°C before the temperature shift. Latrunculin-A (latA) (Enzo Life Sciences; BML-T119) was used at the final concentration of 150 µM to perturb the actin dynamics in spheroplasts. Brefeldin A (Fisher Scientific; 15526276) was used at the final concentration of 75 µM to slow down plasma membrane invagination. Calcofluor White Stain for cell wall staining was purchased from Sigma.

Preparation of *S. pombe* spheroplasts for live-cell imaging (spheroplasting)

The *cps1*-191 cells used in this study were first cultured in YEA medium at 24°C to mid-log phase (OD$_{595}$ = 0.2-0.5), and then were shifted to 36°C for 6 hours 15 minutes (non-permissive conditions). Twenty milliliters of culture were spun down at 3,000 r.p.m. for 1 minute and washed once with equal volume of E-buffer (50 mM sodium citrate, 100 mM sodium phosphate, [pH 6.0]). After spinning down the cells and resuspending cells in 5 ml of E-buffer containing 1.2 M sorbitol, the cell suspension was incubated with 30 mg of lysing enzyme Enzymes from Trichoderma harzianum or Glucanex (Sigma, L1412; an enzymatic mixture of at least glucanases, cellulase, protease, and chitinase activities) at 36°C with shaking at 80 r.p.m. for 90 minutes. This was followed by continuous incubation with 40 µl of LongLife Zymolyase (G-Biosciences, 1.5 U/µl; an enzymatic mixture with at least β-glucanase, protease and mannanase activities) at 36°C with shaking at 80 r.p.m. for 60 minutes. After enzymatic digestion, the cell suspensions were spun down at 450 xg for 2 minutes and washed once with 5 ml of E-buffer containing 0.6 M sorbitol. After spinning at 450 xg for 2 minutes, the spheroplasts were recovered in 10 ml EMMA medium (Edinburgh minimal medium with all amino acids and nucleotides supplements) containing 0.8 M sorbitol and 0.5% (v/v) of 1 M 2-deoxyglucose (Sigma, D6134) for 30 minutes at 36°C prior to microscopy imaging.
Sample preparation for light microscopy

One to two milliliters of spheroplast suspensions in EMMA medium containing 0.8 M sorbitol and 0.5% (v/v) of 1 M 2-deoxyglucose (Sigma, D6134) were concentrated to 20-100 µl by centrifugation at 450 xg for 2 minutes. About 10 µl of concentrated spheroplasts were loaded onto an Ibidi µ-Slide 8-Well glass bottom dish (Cat. No. 80827), and covered with mineral oil (Sigma, M5310) to prevent evaporation during imaging process.

To image cells in Figure 1 and supplementary Figure 4, the cps1-191 cells and rlc1Δ cells after shifting to non-permissive conditions were treated with buffers used to prepare spheroplasts but with the lysing and lytic enzymes omitted to preserve the cell wall integrity. After the buffer washing, the cps1-191 cells in the Figure 1 were recovered in EMMA medium with full supplements containing 0.8 M sorbitol and 0.5% 2-DG. For the rlc1Δ cells in the supplementary Figure 4, after the buffer washing, the cells were recovered in EMMA medium with full supplements containing 0.8 M sorbitol but not 2-DG to allow septation.

Sample preparation for electron microscopy

Two hundred and fifty milliliters of cells with OD_{595} 0.2 were collected for spheroplasting. Spheroplasts were prepared with the spheroplasting method described above. Spheroplasts were spun down from EMMA with 0.8 M sorbitol and resuspended in phosphate-buffered saline (PBS) with 2.5% glutaraldehyde and 1.2 M sorbitol. Fixation solution was prepared by adding 2% glutaraldehyde and dissolving 1.2 M sorbitol in PBS. After 2 hours incubation at room temperature, spheroplasts were spun down in round bottom tubes. The following procedures were done at 4°C and gently (vortex mixer was avoided). Spheroplasts were resuspended in fixation solution and stood on ice for 2 hours. The spheroplasts were separated into 2 tubes: washed and unwashed samples. Unwashed samples were stored at 4°C. The washed samples were washed with 1 mL PBS containing 1.2 M sorbitol for three times. Lastly the spheroplasts were resuspended in 1 mL PBS containing 1.2 M sorbitol and stored at 4°C before electron microscopy.
For electron microscopy, glutaraldehyde-fixed cells were placed on a slide glass whose surface was pre-treated with 0.1% poly-L-lysine. They were washed with 0.1 M phosphate buffer (pH 7.2), post-fixed with 1% osmium tetroxide at 4°C for 1 hour, dehydrated with graded series of ethanol, and critical point dried with a Leica EM CPD030 apparatus (Leica Microsystems, Vienna). The specimens were coated with osmium tetroxide by osmium coater (Vacuum Device.inc, Japan) and observed with S-3400N and SU8020 scanning electron microscope (Hitachi High Technologies, Tokyo) at 10.0 kV and 1.0 kV respectively (Namiki et al., 2011).

Light microscopy

The Andor Revolution XD spinning disk confocal microscope was used to image the spheroplasts and cells at 36°C. The microscope was equipped with a Nikon ECLIPSE Ti inverted microscope, Nikon Plan Apo Lambda 100×/1.45N.A. oil immersion objective lens, a spinning-disk system (CSU-X1; Yokogawa), and the Andor iXon Ultra EMCCD camera 897 or 888. The Andor IQ3 software was used to acquire images at the pixel size of 80 nm/pixel or 69 nm/pixel, depending on the camera models. Laser lines at wavelengths of 405 nm, 488 nm or 561 nm were used for the excitation of fluorophores. Most images were acquired with Z-step sizes of 0.5 µm as listed here: Figure 2A (12 µm / 25 Z-sections); Figure 2B (10 µm / 21 Z-sections); Figure 3A (10 µm / 21 Z-sections); Figure 3B (15 µm / 31 Z-sections); Figure 3C (15 µm / 31 Z-sections); Figure 4A (15 µm / 31 Z-sections); Figure 4B (12 µm / 25 Z-sections); Figure 4C (13 µm / 27 Z-sections); Figure 4D (10 µm / 21 Z-sections); Supplementary figure 1 (10 µm / 21 Z-sections); Supplementary figure 3A (15 µm / 31 Z-sections); Supplementary figure 3B (13 µm / 27 Z-sections); Supplementary figure 4 (13 µm / 27 Z-sections).

Image analysis

Images were processed using Fiji. The time-lapse montages are maximum intensity projections of Z-stack of specified time points. All images analyzed were prepared in this study, except images for quantification of the rate of ring sliding in wild-type spheroplasts in which the data was based on the time-lapse images acquired in a previous study (Lim et al., 2018).
Acknowledgements

This work was supported by research grants from Wellcome Trust (WT101885MA) and ERC (GA 671083 - ACTOMYOSIN RING) to MKB. Part of the work was supported by the Fundamental Research Funds for the Central Universities (K20200099) to TGC.

Figure Legends

Figure 1. Lowering down turgor pressure does not allow cell membrane ingression in cps1 mutant cells.
(A) cps1-191 GFP-psy1 rlc1-tdTomato cells were cultured in YEA at the restrictive temperature of 36°C for 6.5 hours and were processed similarly using the spheroplasting protocol but omitting lysing enzymes and Zymolyase. Cells in the EMMA medium with 2-DG were imaged at 36°C. Green: GFP-psy1. Red: rlc1-tdTomato.

(B) Some cps1-191 GFP-psy1 rlc1-tdTomato cells lysed after more than 6.5 hours of incubation at the restrictive temperature. Treatment of cells was same as in Figure 1(A). Green: rlc1-tdTomato. Red: GFP-psy1.

(C) cps1-191 GFP-psy1 rlc1-tdTomato cells were cultured in YEA at the restrictive temperature for 6.5 hours and were processed similarly using the spheroplasting protocol but omitting lysing enzymes and Zymolyase. Cells were imaged at 36°C in the EMMA medium containing 2-DG and 0.8 M sorbitol to lower down the turgor pressure. Green: GFP-psy1. Red: rlc1-tdTomato.

(D) cps1-191 GFP-psy1 rlc1-tdTomato cells treated as in Figure 1(C) were stained with calcofluor dye to reveal the cell wall.

Scale bar: 5 µm

Figure 2. Weakening of cell wall allows ring contraction and cell membrane ingression.
(A) Two examples of cps1-191 spheroplasts underwent cytofission at 36°C. The cps1-191 GFP-psy1 rlc1-tdTomato cells were cultured at 36°C for 6.5 hours, processed into spheroplasts, and recovered for 1 hour at 36°C prior to imaging.
(B) Two examples of *cps1*-191 spheroplasts expressing Lifeact-mCherry underwent cytofission at 36°C. The *cps1*-191 GFP-psy1 lifeact-mCherry cells were cultured at 36°C for 6.5 hours, processed into spheroplasts, and recovered for 1 hour at 36°C prior to imaging.

(C) Quantification of the rate of ring contraction in wild type cells and *cps1*-191 spheroplasts undergoing cytofission.

(D) Wild type cells and *cps1*-191 GFP-psy1 *rlc1*-tdTomato spheroplasts were stained with the calcofluor dye. The image was pseudo-colored in green to represent calcofluor staining.

(E) Electron micrographs of *cps1*-191 GFP-psy1 *rlc1*-tdTomato spheroplasts regenerated in medium with or without 2-DG.

Scale bar: 5 µm except Figure 2E, which is 1 µm; Error bars: standard deviation

**Figure 3.** The *cps1*-191 mutant spheroplasts undergo cytofission independent of the α-glucan synthase and endoglucanases.

(A) Cytofission in *cps1*-191 *mok1*-664 GFP-psi1 *rlc1*-tdTomato.

(B) Cytofission in *cps1*-191 *agn1*Δ GFP-psy1 *rlc1*-tdTomato.

(C) Cytofission in *cps1*-191 *eng1*Δ GFP-psy1 *rlc1*-tdTomato.

Scale bar: 5 µm

**Figure 4.** The function of actomyosin rings is required in the *cps1*-191 mutant spheroplasts to undergo cytofission.

(A) *cps1*-191 GFP-psy1 *rlc1*-tdTomato spheroplasts underwent cytofission in the presence of DMSO. Left panel shows the DIC images; right panel shows the fluorescence micrographs.

(B) *cps1*-191 GFP-psy1 *rlc1*-tdTomato spheroplasts were incubated with 150 µm Lat-A. Left panel shows the DIC images; right panel shows the fluorescence micrographs.

(C) *rlc1*Δ *cyk3*-GFP spheroplasts failed to undergo cytofission at 36°C. The *rlc1*Δ *cyk3*-GFP cells were cultured at 36°C for 6.5 hours, processed into spheroplasts, and then recovered in minimal medium containing sorbitol prior to imaging at 36°C. Top panel shows the DIC images; bottom panel shows the fluorescence micrographs.
(D) The *cps1-191 GFP-psy1 rlc1-tdTomato* spheroplasts failed to undergo cytofission in the presence of 75 µM brefeldin A.

Scale bar: 5 µm
References

Arasada, R., and Pollard, T.D. (2014). Contractile ring stability in S. pombe depends on F-BAR protein Cdc15p and Bgs1p transport from the Golgi complex. Cell Rep 8, 1533-1544.

Arasada, R., and Pollard, T.D. (2015). A role for F-BAR protein Rga7p during cytokinesis in S. pombe. J Cell Sci 128, 2259-2268.

Burton, K., and Taylor, D.L. (1997). Traction forces of cytokinesis measured with optically modified elastic substrata. Nature 385, 450-454.

Choudhary, A., Lera, R.F., Martowicz, M.L., Oxendine, K., Laffin, J.J., Weaver, B.A., and Burkard, M.E. (2013). Interphase cytofission maintains genomic integrity of human cells after failed cytokinesis. Proc Natl Acad Sci U S A 110, 13026-13031.

Cortes, J.C., Carnero, E., Ishiguro, J., Sanchez, Y., Duran, A., and Ribas, J.C. (2005). The novel fission yeast (1,3)beta-D-glucan synthase catalytic subunit Bgs4p is essential during both cytokinesis and polarized growth. J Cell Sci 118, 2259-2268.

Garcia Cortes, J.C., Ramos, M., Osumi, M., Perez, P., and Ribas, J.C. (2016). The Cell Biology of Fission Yeast Septation. Microbiol Mol Biol Rev 80, 779-791.

Garcia, I., Jimenez, D., Martin, V., Duran, A., and Sanchez, Y. (2005). The alpha-glucanase Agn1p is required for cell separation in Schizosaccharomyces pombe. Biol Cell 97, 569-576.

Lim, T.C., Hatano, T., Kamnev, A., Balasubramanian, M.K., and Chew, T.G. (2018). Equatorial Assembly of the Cell-Division Actomyosin Ring in the Absence of Cytokinetic Spatial Cues. Curr Biol 28, 955-962 e953.

Liu, J., Wang, H., and Balasubramanian, M.K. (2000). A checkpoint that monitors cytokinesis in Schizosaccharomyces pombe. J Cell Sci 113 (Pt 7), 2289-2299.
Ring Is Required for the Correct Progression of Cytokinesis in Fission Yeast. Cell Rep 25, 772-783.

Megnet, R. (1965). Effect of 2-deoxyglucose on Schizosaccharomyces pombe. J Bacteriol 90, 1032-1035.

Mishra, M., Huang, Y., Srivastava, P., Srinivasan, R., Sevugan, M., Shlomovitz, R., Gov, N., Rao, M., and Balasubramanian, M. (2012). Cylindrical cellular geometry ensures fidelity of division site placement in fission yeast. J Cell Sci 125, 3850-3857.

Morton, W.M., Ayscough, K.R., and McLaughlin, P.J. (2000). Latrunculin alters the actin-monomer subunit interface to prevent polymerization. Nat Cell Biol 2, 376-378.

Munoz, J., Cortes, J.C., Sipiczki, M., Ramos, M., Clemente-Ramos, J.A., Moreno, M.B., Martins, I.M., Perez, P., and Ribas, J.C. (2013). Extracellular cell wall beta(1,3)glucan is required to couple septation to actomyosin ring contraction. J Cell Biol 203, 265-282.

Naqvi, N.I., Wong, K.C., Tang, X., and Balasubramanian, M.K. (2000). Type II myosin regulatory light chain relieves auto-inhibition of myosin-heavy-chain function. Nat Cell Biol 2, 855-858.

Onwubiko, U.N., Mlynarczyk, P.J., Wei, B., Habiyaremye, J., Clack, A., Abel, S.M., and Das, M.E. (2019). A Cdc42 GEF, Gef1, through endocytosis organizes F-BAR Cdc15 along the actomyosin ring and promotes concentric furrowing. J Cell Sci 132.

Osumi, M., Sato, M., Ishijima, S.A., Konomi, M., Takagi, T., and Yaguchi, H. (1998). Dynamics of cell wall formation in fission yeast, Schizosaccharomyces pombe. Fungal Genet Biol 24, 178-206.

Pollard, T.D., Bookwalter, C.S., Tang, Q., Kremenskova, E.B., Trybus, K.M., and Lowey, S. (2017). Fission yeast myosin Myo2 is down-regulated in actin affinity by light chain phosphorylation. Proc Natl Acad Sci U S A 114, E7236-E7244.

Proctor, S.A., Minc, N., Boudaoud, A., and Chang, F. (2012). Contributions of turgor pressure, the contractile ring, and septum assembly to forces in cytokinesis in fission yeast. Curr Biol 22, 1601-1608.

Ramos, M., Cortes, J.C.G., Sato, M., Rincon, S.A., Moreno, M.B., Clemente-Ramos, J.A., Osumi, M., Perez, P., and Ribas, J.C. (2019). Two S. pombe septation phases differ in ingression rate, septum structure, and response to F-actin loss. J Cell Biol 218, 4171-4194.

Sethi, K., Palani, S., Cortes, J.C., Sato, M., Sevugan, M., Ramos, M., Vijaykumar, S., Osumi, M., Naqvi, N.I., Ribas, J.C., and Balasubramanian, M. (2016). A New Membrane Protein Sbg1 Links the Contractile Ring Apparatus and Septum Synthesis Machinery in Fission Yeast. PLoS Genet 12, e1006383.

Sipiczki, M. (2007). Splitting of the fission yeast septum. FEMS Yeast Res 7, 761-770.

Stachowiak, M.R., Laplante, C., Chin, H.F., Guirao, B., Karatekin, E., Pollard, T.D., and O'Shaughnessy, B. (2014). Mechanism of cytokinetic contractile ring constriction in fission yeast. Dev Cell 29, 547-561.

Steinkuhler, J., Knorr, R.L., Zhao, Z., Bhatia, T., Bartelt, S.M., Wegner, S., Dimova, R., and Lipowsky, R. (2020). Controlled division of cell-sized vesicles by low densities of membrane-bound proteins. Nat Commun 11, 905.

Svoboda, A., and Smith, D.G. (1972). Inhibitory effect of 2-deoxy-glucose on cell wall synthesis in cells and protoplasts of Schizosaccharomyces pombe. Z Allg Mikrobiol 12, 685-699.

Taneja, N., Fenix, A.M., Rathbun, L., Millis, B.A., Tyska, M.J., Hehnly, H., and Burnette, D.T. (2016). Focal adhesions control cleavage furrow shape and spindle tilt during mitosis. Sci Rep 6, 29846.

Uroz, M., Garcia-Puig, A., Tekeli, I., Elosequi-Artola, A., Abenza, J.F., Marin-Llaurado, A., Pujals, S., Conte, V., Albertazzi, L., Roca-Cusachs, P., Raya, A., and Trepot, X. (2019). Traction forces at the cytokinetic ring regulate cell division and polyplody in the migrating zebrafish epicardium. Nat Mater 18, 1015-1023.

Wang, N., Lee, I.J., Rask, G., and Wu, J.Q. (2016). Roles of the TRAPP-II Complex and the Exocyst in Membrane Deposition during Fission Yeast Cytokinesis. PLoS Biol 14, e1002437.
Figure 1

(A) cps1-191 cells in normal turgor

(B) cps1-191 cell lysis in normal turgor

(C) cps1-191 cells in low turgor

(D) cps1-191 cells in low turgor
Figure 2

(A) *cps1-191 GFP-psy1 rlc1-tdTomato*
Spheroplast 1

(B) *cps1-191 lifeact-mCherry*
Spheroplast 1

(C) Rate of ring contraction (µm/min)

(D) DIC

(E) In 2-DG medium

Wild-type cells + *cps1-191* spheroplasts after 2h 50min in 2-DG medium
Figure 3

(A) *cps1-191 mok1-664 GFP-psy1 rlc1-tdTomato*

(B) *cps1-191 eng1Δ GFP-psy1 rlc1-tdTomato*

(C) *cps1-191 agn1Δ GFP-psy1 rlc1-tdTomato*
Figure 4

(A) *cps1-191 GFP-psy1 rlc1-tdTomato + DMSO*

(B) *cps1-191 GFP-psy1 rlc1-tdTomato + Lat-A*

(C) *cps1-191 rlc1Δ cyk3-GFP spheroplast*

(D) *cps1-191 GFP-psy1 rlc1-tdTomato + Brefeldin A*