Centrosomal MPF triggers the mitotic and morphogenetic switches of fission yeast

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Activation of mitosis-promoting factor (MPF) drives mitotic commitment¹. In human cells active MPF appears first on centrosomes². We show that local activation of MPF on the equivalent organelle of fission yeast, the spindle pole body (SPB), promotes Polo kinase activity at the SPBs long before global MPF activation drives mitotic commitment. Artificially promoting MPF or Polo activity at various locations revealed that this local control of Plo1 activity on G2 phase SPBs dictates the timing of mitotic commitment. Cytokinesis of the rod-shaped fission yeast cell generates a naive, new, cell end. Growth is restricted to the experienced old end until a point in G2 phase called new end take off (NETO) when bipolar growth is triggered³. NETO coincided with MPF activation of Plo1 on G2 phase SPBs (ref. 4). Both MPF and Polo activities were required for NETO and both induced NETO when ectopically activated at interphase SPBs. NETO promotion by MPF required polo. Thus, local MPF activation on G2 SPBs directs polo kinase to control at least two distinct and temporally separated, cell-cycle transitions at remote locations.

Mutations in the SPB component Cut12 exhibit a reciprocal relationship with mutations in the MPF-activating phosphatase Cdc25. The loss-of-function mutation cdc25.22 is suppressed by gain-of-function mutations in cut12, such as cut12.s11 (refs 5,6). Conversely, increased Cdc25 levels suppress the loss-of-function mutant cut12.1 (ref. 7). The polo kinase Plo1 associates with the SPB towards the end of G2 phase. The timing of this recruitment is advanced in cut12.s11 cells⁴. The antibody MPM2 recognizes SPBs in a Plo1-dependent manner⁴ and so offers an ideal tool with which to assess the local activity of Plo1 on the SPB. Unlike bulk in vitro kinase assays, MPM2 assessment of Plo1 activity on SPBs is independent of any changes in Plo1 activities at any other locations. MPM2 SPB staining indicates that the Plo1 that associates with SPBs during G2 phase of both wild-type and cut12.s11 cells is active⁵. Prompted by the fact that the enhancement of Plo1 activity that accompanies mitotic commitment is driven by MPF activation⁶, we now address the function of this SPB-associated Plo1 in mitotic control.

MPM2 or Plo1 immunofluorescence staining reveals single (late G2 phase) or paired foci (mitotic) of SPB staining. The ratio of 1:2 foci (G2/Mitotic) in cell populations gives an indication of the point in G2 at which Plo1 is recruited to (Plo1 staining), or activated at (MPM2), SPBs. Staining patterns in an established plo1.GFP strain⁷ revealed that the fusion of GFP sequences to the carboxy terminus of Plo1 did not alter the timing or activation of Plo1 at SPBs (Fig. 1a). Time-lapse imaging revealed that Plo1 was initially recruited around 30 min before mitosis (Fig. 1a–c). This initial recruitment was advanced by a further 30 min in cut12.s11 cells (Fig. 1a–c and Supplementary Video S1). Fluorescence recovery after photobleaching (FRAP) revealed a rapid turnover of Plo1.GFP (half-life 22–23 s) at the SPBs of both cut12.s11 and cut12.s11 cells (Fig. 1d and Supplementary Video S2). To assess the impact of MPF activity on Plo1 SPB recruitment we exploited the cdc2.2 mutation in the catalytic subunit of MPF that renders it sensitive to inhibition by ATP analogues⁸. plo1.GFP cdc2⁺ cells were labelled by transient immersion in fluorescent lectin⁹ and mixed with plo1.GFP cdc2.2 cells for live-cell imaging. Addition of the ATP analogue 1NA-PP1 prompted the immediate loss of Plo1.GFP from G2 SPBs of neighbouring cdc2⁺ cells (Fig. 1e,f and Supplementary Videos S3 and S4). Restoration of MPF activity by removal of the analogue promoted a rapid return (Fig. 1f and Supplementary Video S4), indicating that MPF activity is continuously required for Plo1 association with the late G2 SPB.

We extended the analogue-sensitive approach to assess the contribution of Plo1 activity to mitotic control in unperturbed
Table: Cell density (cell ml⁻¹) and Live cell Plo1 Recruitment duration (min) for different cell lines.

| Cell line         | Cell density (cell ml⁻¹) | Live cell Plo1 Recruitment duration (min) |
|-------------------|--------------------------|------------------------------------------|
| Wild type         | 1 × 10⁶                  | 1                                        |
| cut12.s11         | 2 × 10⁶                  | 2.5                                      |
| plo1.GFP          | 2 × 10⁶                  | 1.3                                      |
| Wild type         | 4 × 10⁶                  | 1.8                                      |
| plo1.GFP          | 4 × 10⁶                  | 1.8                                      |

Figure 1: MPF activity controls Plo1.GFP recruitment to the SPB in late G2 phase. (a) Timing of Plo1.GFP recruitment before SPB separation (living cells) and the ratios of 1 dot (G2, early mitotic SPB) to two dots (mitotic SPB) for MPM2 staining. (b) Plots of the intensity of the Plo1.GFP signals averaged for three cells. The single plot for each strain splits into two different plots after SPB separation to indicate the signals arising from two individual SPBs that emerge from this single focus of paired SPB signals in G2. intensity of the two (see also Supplementary Video S1). (c) Time-lapse images of plo1.GFP and cut12.s11 plo1.GFP cells. (d) FRAP of the signal from a late G2 Plo1.GFP SPB in wild-type cells (left). The FRAP profile of a cut12.s11 plo1.GFP strain was indistinguishable from this wild-type plot (right; see also Supplementary Video S2). (e,f) Wild-type (TRITC–lectin pre-treated) and analogue-sensitive (cdc2.as) cells (no coating) recorded side by side in the same field of view at 25°C. The timing of the addition of 25 μM 1NA-PP1 is indicated in the panel. Control cdc2 cells advanced through mitosis irrespective of the presence of analogue with Plo1.GFP on their SPBs whereas Plo1.GFP signal left the SPBs of neighbouring cdc2.as mutant cells following analogue addition. White arrows indicate Plo1.GFP signals on SPBs. See also Supplementary Videos S3 and S4. (g,h) Size-selected cultures of either plo1.as8 (g) or plo1 (h) cells were split into three following completion of the first synchronous division after size selection and treated as indicated in the legend to the graphs before processing for immunofluorescence microscopy with phospho-histone H3 Ser 10 antibodies. The analogue 3-BrB-PP1 was added to a final concentration of 20 μM. The numbers in time-lapse images indicate minutes.
cell cycles. In strains harbouring the plo1.as1, plo1.as2, plo1.as3 or plo1.as4 mutations that are predicted to confer sensitivity\(^1\), addition of analogue had little impact on the most sensitive readout of Plo1 function, spindle formation\(^2\). However, seption was compromised in the first division after the addition of 40 \(\mu\)M 3MB-PP1 to plo1.as3 cells (data not shown), indicating a modest degree of sensitization\(^2\).

We therefore studied the structure of AMP-PNP-bound human Plk1 (Supplementary Fig. S1c) to seek notable differences between the structure of this mammalian kinase that can be sensitized to ATP analogue inhibition and its refractory fission yeast counterpart, Plk1. Plk1 harbours a phenylalanine at position 170 at the bottom of the ATP-binding pocket that is replaced by methionine in Plo1. Mutation of this methionine to phenylalanine in a plo1.as3 backbone created the plo1.as8 allele that responded to 3MB-PP1 or 3-BrB-PP1 addition by arresting mitotic progression with the monopolar spindle phenotype that is a characteristic of severely compromised Plo1 function\(^4\) (Supplementary Fig. S1a,b). Addition of 20 \(\mu\)M 3-BrB-PP1 to G2 plo1.as8 cells delayed the phosphorylation of histone H3 Ser 10 that accompanies mitotic commitment by 40 min (Fig. 1g,h).

Plo1 comprises a catalytic domain and two polo boxes that direct the kinase to target molecules that are either direct substrates or platforms from which neighbouring substrates can be phosphorylated\(^6\) (Supplementary Fig. S1c). We reasoned that Plo1 could be directed to a site of choice by replacement of the polo boxes with a surrogate targeting sequence. We replaced the polo box domain of plo1.as8 with sequences encoding the GFP-binding protein (GBP), in a transgene that was expressed from an ectopic location. We incorporated mutations conferring constitutive activation\(^8\) into the catalytic domain of this hybrid molecule to override the controls that normally coordinate Plo1 activity with cell-cycle progression (Supplementary Fig. S1c). Ectopic expression of this Plo1.RL chimaera in the presence of inhibitory ATP analogue targets a restrained kinase to any location that hosts GFP (Supplementary Fig. S2a). Subsequent analogue removal releases sustained Plo1 kinase activity at the target site. This approach enabled us to uncouple kinase activation from its normal cell-cycle context and to examine which events are triggered by forced Plo1 activation at a particular location.

Plo1.RL activation at SPBs through recruitment to Cut12.NEGFP promoted mitotic commitment (Fig. 2a). A lower level of mitotic induction arose from recruitment to the SPB components Pcp1.GFP and Sid4.GFP whereas none followed targeting to any other location (Fig. 2a and Supplementary Fig. S2b). Bleaching the fluorescence signal derived from either the host molecule Cut12.NEGFP or the associated Plo1.RL.Tom chimaeric kinase revealed no turnover of the Cut12.NEGFP/Plo1.RL.Tom complex at SPBs (Fig. 2b). We therefore investigated whether the anchored Plo1.RL chimaeric kinase relied on the endogenous, mobile, Plo1 to promote mitotic commitment. plo1-ts41 cells (Supplementary Fig. S1e,f) harbouring Cut12.NEGFP-anchored Plo1.RL were shifted from 25 \(^\circ\)C to 36 \(^\circ\)C to inhibit Plo1 function as the inhibitory analogue was removed. Different induction times produced different levels of Plo1.RL at this point of release. Levels of Plo1.RL (18 or 20 h induction) that were competent to induce mitosis when the endogenous Plo1 kinase was a functional wild-type molecule were unable to do so when it was the incapacitated Plo1.ts41 (Fig. 2c).

As MPF associates with G2 SPBs (refs 22,23) and controls SPB recruitment of Plo1 (Fig. 1e,f), we investigated whether it was also responsible for generating the pro-mitotic Plo1 signal at SPBs. We examined this possibility with a Cdc2–GBP fusion protein, Cdc2.RL (cdc2.F84GY15F, Supplementary Fig. S1c) that is insensitive to inhibition by either Mtk1 or Wee1 (ref. 24) and sensitive to ATP analogues\(^11\). Activation of Cdc2.RL by analogue removal from strains in which the chimaeric kinase had been recruited to a variety of locations (Supplementary Fig. S3a) mirrored the Plo1.RL data: SPB recruitment through Cut12.NEGFP gave a strong induction of mitosis; weaker levels followed SPB recruitment with either Pcp1.GFP or Sid4.GFP and no induction was seen following recruitment to any other location (Fig. 2d and Supplementary Fig. S3b). Bleaching Cut12.NEGFP-anchored Cdc2.RL.Tom revealed a static association with the SPBs (Supplementary Fig. S3c). MPM2 reactivity of SPBs showed that the wave of mitosis that arose from activation of Cut12.NEGFP-anchored Cdc2.RL was preceded by Plo1 activation (Supplementary Fig. S3d). Consistently, this ability of Cut12.NEGFP-anchored Cdc2.RL to induce mitosis relied on Plo1 activity (Fig. 2e).

Three further Cut12.NEGFP/Cdc2.RL experiments were informative. When the cdc2 encoding sequence retained Tyr 15 (Cdc2.RLY15, Supplementary Fig. S1c), mitosis was not induced (Fig. 2f), indicating that the balance of Wee1 and Cdc25 activities normally determines the timing at which SPB-bound MPF activity is enhanced to trigger mitosis. Mitotic induction by SPB-tethered Cdc2-RL was blocked in a cdc2.33 background in which the bulk population of MPF within the nucleus\(^22,23\) was inactivated (Fig. 2g,h), even though Plo1 activity was promoted on the SPBs of these interphase cells (Fig. 2i). Thus, division is not driven from the SPB, rather MPF activation at the SPB serves as a trigger event that promotes conversion of the bulk population MPF throughout the cell\(^22,23\) to the active state. The second wave of mitosis after analogue removal was abolished by the inclusion of hydroxyurea in the replacement, analogue-free, medium (Fig. 2f), indicating that the checkpoint remains intact.

The point in G2 phase at which Plo1 was initially recruited to G2 SPBs is strongly influenced by cell density (Fig. 1a). Consistently, nutrient provision and TOR signalling is closely tied to SPB recruitment of Plo1 in mitotic control\(^25\)-\(^28\). This context dependency is highly reminiscent of the way by which the timing of the G2 event NETO changes in response to changes in nutrient supply\(^7\). We therefore examined the relationship between Plo1 recruitment to the G2 SPB and NETO. Strikingly, the two events were coincident: in counts of 100 cells, every post-NETO cell had none (Fig. 3a). The correlation between NETO and Plo1 recruitment was even maintained when Plo1 SPB recruitment was advanced by 30 min by cut12.s11 (Fig. 3b).

To determine whether there was a direct causal relationship between Plo1 activity and NETO, NETO was monitored when small G2 plo1.ts41 cells (Supplementary Fig. S1e,f) were shifted from 25 \(^\circ\)C to 36 \(^\circ\)C. NETO execution was severely compromised by Plo1 inactivation (Fig. 3c and Supplementary Fig. S1g). To determine whether it was the SPB-associated pool of Plo1 that triggered NETO, we targeted Plo1.RL to various locations in cdc10.v50 cells. At 36 \(^\circ\)C, cdc10.v50 arrests cell-cycle progression in G1 phase before NETO (ref. 3). NETO was triggered when Plo1.RL was activated at the SPBs of cdc10.v50 cells but nowhere else, including recruitment to the cell end itself through either Tea1.v5GFP or For3.GFP fusion proteins (Fig. 3d and Supplementary Fig. S2a).
The observation that G2-arrested cdc2.33 cells undergo NETO (ref. 3), yet NETO can be triggered by Plo1 activation on SPBs (an MPF-dependent event; Fig. 1e–f), prompted us to re-examine the relationship between NETO and Cdc2 with the cdc2.33 allele. Addition of 1 μM BrB-PP1 to G2 cells mimicked temperature-mediated inactivation of cdc2.33 in having no impact on NETO yet arresting cell-cycle progression in G2. However, a stronger level of inhibition (20 μM BrB-PP1) did block NETO (Fig. 4a and Supplementary Fig. S4), indicating that the threshold of MPF activity required for NETO was lower than that for mitotic commitment.

The contribution of SPB-associated MPF to the promotion of NETO cannot be assessed by the Cdc2.RL approach in wild-type cultures because the immediate induction of mitotic commitment (Fig. 2d) obscures any impact on NETO. We therefore blocked the propagation of the mitotic commitment signal with the cdc2.33 mutation (Fig. 2h) before assessing NETO induction. Hydroxyurea...
Figure 3 NETO is triggered by Plo1 activation/recruitment on the G2 SPB. (a) Fluorescence imaging of cell wall signals (calcofluor, top), Plo1.GFP and Sid4.Tom of living cells of the indicated strains. Arrows identify cells with Plo1.GFP signals on SPBs. (b) Size-selected small G2 cells were fixed and scored for NETO status and septation index as they transited the cell division cycle. (c) Cultures in which cell-cycle progression had been synchronized by the selection of small cells at 25°C before temperature shift to 36°C for 1 h to inactivate Plo1.RL (Supplementary Fig. S1g). (d) Mutants harbouring cdc10.v50 and the indicated GFP fusion genes were grown to early log phase at 25°C before the plo1.RL allele was induced by removal of thiamine in the presence of 40 μM 3MB-PP1. After dilution and a further 24 h at 25°C the early log phase cells were filtered into analogue- and thiamine-free medium and split into two, one half being kept at 25°C (upper), the other being shifted to 36°C (lower). Five hours later, the cells were fixed and the frequency of NETO was scored (left). The panels on the right show examples of the wild-type control (upper) and cut12.NEGFP (lower) cdc10.v50 strains after incubation at 36°C stained with calcofluor to highlight cell wall material or TRITC phalloidin to stain F-actin.

| Time (min) | Post-NETO cells (%) | Septating cells (%) |
|------------|---------------------|---------------------|
| 0          | 100                 | 0                   |
| 100        | 80                  | 20                  |
| 200        | 60                  | 40                  |
| 300        | 40                  | 60                  |
| 400        | 20                  | 80                  |
| 500        | 0                   | 100                 |

Post-NETO: no growth at new end
Pre-NETO: growth at new end

Netotic cell wall attracted by Cdc2.RL at the new end

Netotic F-actin stained by TRITC phalloidin

Netotic growth from new end

Netotic delay in mitosis

Netotic mitotic index

Netotic mitotic spindle index

Netotic Fig. 3 NETO is triggered by Plo1 activation/recruitment on the G2 SPB. (a) Fluorescence imaging of cell wall signals (calcofluor, top), Plo1.GFP and Sid4.Tom of living cells of the indicated strains. Arrows identify cells with Plo1.GFP signals on SPBs. (b) Size-selected small G2 cells were fixed and scored for NETO status and septation index as they transited the cell division cycle. (c) Cultures in which cell-cycle progression had been synchronized by the selection of small cells at 25°C before temperature shift to 36°C for 1 h to inactivate Plo1.RL (Supplementary Fig. S1g). (d) Mutants harbouring cdc10.v50 and the indicated GFP fusion genes were grown to early log phase at 25°C before the plo1.RL allele was induced by removal of thiamine in the presence of 40 μM 3MB-PP1. After dilution and a further 24 h at 25°C the early log phase cells were filtered into analogue- and thiamine-free medium and split into two, one half being kept at 25°C (upper), the other being shifted to 36°C (lower). Five hours later, the cells were fixed and the frequency of NETO was scored (left). The panels on the right show examples of the wild-type control (upper) and cut12.NEGFP (lower) cdc10.v50 strains after incubation at 36°C stained with calcofluor to highlight cell wall material or TRITC phalloidin to stain F-actin.
Figure 4 MPF activation on the SPB triggers NETO in a Plo1-dependent manner. (a) Cell-cycle progression was synchronized by the selection of small, G2, cdc2.33 cells at 25 °C. After transit of one cell cycle at 25 °C (Supplementary Fig. S4a) the culture was split into four equal aliquots that were all shifted to 32 °C. Either 1 or 20 μM 3-BrB-PP1, solvent alone or nothing was added before NETO was scored in fixed samples at the indicated intervals. Samples were also processed for FACS (fluorescence-activated cell sorting) analysis to monitor DNA content (right). For a repeat of this FACS analysis see Supplementary Fig. S4c. (b) The scheme used to assess whether analogue removal from strains harbouring cdc2.33 targeted to the SPB by Cut12.NEGFP targeted proteins was able to induce NETO when the bulk population of cdc2.33 cdc2.RL plo1.ts41 MPF in the cell was inactivated by incubation of cdc2.33 cells at 36 °C. HU, hydroxyurea. (c,d) Whereas NETO was induced when Cdc2.RL was targeted to the SPB by Cut12.NEGFP (c), no induction occurred when Plo1 kinase was inhibited (d). (e) Left, cut12.NEGFP cells containing the indicated additional mutations were treated as for Fig. 3d except that cells were maintained at 25 °C throughout. The time interval is in hours. Activation of Plo1 or Cdc2 on the SPB did not induce NETO in the polarity mutants bud6.Δ, tea1.Δ or sla2.Δ despite the fact that processing of the cut12.NEGFP mutant strains as described for Fig. 2a revealed that none of them compromised the ability of activation of either kinase on SPBs to induce mitosis (right).

ability of Cut12.NEGFP-anchored Cdc2.RL to induce NETO was reliant on Plo1 activity (Fig. 4d). It also required the polarity factors Tea1, Bud6 and Sla2 (Fig. 4e).

We propose that Plo1 recruitment to SPBs in G2 phase both triggers the cell-cycle-dependent morphogenetic switch to bipolar growth and constitutes a priming event that licenses the cell to regulate the timing of mitotic commitment. The MPF dependency and activity of the Plo1 recruited to these G2 SPBs suggests a local activation of the feedback loops that will later be employed at remote locations to drive the global cell cycle during mitotic commitment (Fig. 5a). As NETO happens as soon as the feedback loop is running on the SPB the target/s for Plo1 in NETO control probably differ from those in G2/M commitment control. The fact that it is only constitutively active forms of MPF that can override the normal controls to induce mitosis and NETO
when targeted to the SPB (Fig. 2f and Supplementary Figs S2d,e and S3f) indicates that Wee1 activity blocks the propagation of the mitotic commitment signal from the SPB.

Once the restraining activity of Wee1 is removed/overcome, cells can execute mitosis. Given the relative location of the different signalling components, it would seem likely that the anti-mitotic activity of the equatorial belt of inhibitory nodes that contain Cdr1, Cdr2 and Wee1 and encircle the nucleus blocks the propagation of the signals emanating from the SPB (Fig. 5b)\(^29,30\).

The well-documented ability of TOR signalling to influence the timing of mitotic commitment in fission yeast relies on its ability to promote Plo1 recruitment to the SPB (refs 25–28), indicating that, in some situations, enhancing the level of Plo1 signalling from the SPB is sufficient to advance the timing of mitotic commitment. Perhaps mitotic commitment is also finally triggered when Plo1 activity exceeds a critical threshold in unperturbed cell cycles.

As NETO ensures that each daughter cell will receive an experienced, old, cell tip that will be optimal for growth in the subsequent cell cycle, Plo1 recruitment to G2 SPBs could be seen as the first phase of mitotic commitment. However, the question remains as to how a signal that emanates from the cell equator triggers an event at the cell extremity (NETO). A recent study may hold the clue\(^31\). The Cdc42 activation that drives tip growth is not a persistent signal at one end. Rather, it alternates between the two cell tips throughout interphase. Thus, Cdc42 is activated at new ends every 3 min throughout early G2 up to NETO. However, NETO is triggered only once global Cdc42 activity exceeds a critical threshold level of global Cdc42 activation. A general signal emanating from the SPB could set the level of this general threshold by targeting Cdc42 or its regulators globally.

Given the impact of TOR and stress pathways on SPB recruitment of Plo1 (refs 26,27,32) and the coordination of cytokinetic ring formation and function by a second cell-cycle regulatory network on the same SPBs (the septum initiation network\(^23\)), we propose that the SPB (and by implication the centrosome) acts as a platform at which signals from diverse signalling pathways are integrated to generate coordinated and coherent signals to control cell-cycle progression.

**METHODS**

Methods and any associated references are available in the online version of the paper.

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AUTHOR CONTRIBUTIONS
I.M.H. conceived the study. A.K. and V.S. generated cdc2. A.P. generated the conditional polo kinase alleles, including Plo1.RL. V.A.T. and K.Y.C. carried out the imaging work in Fig. 1. A.G. performed the remainder of the experiments with occasional assistance from A.P. and K.Y.C. for more complex experiments and cloning. I.M.H. wrote the manuscript with input and discussions from all authors.

COMPETING FINANCIAL INTERESTS
The authors declare no competing financial interests.

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METHODS

Cell culture and growth. Strains used in this study are listed in Supplementary Table S1. Cells were grown and maintained according to ref. 34. Appropriately supplemented EMM2 synthetic medium was used for all experiments. As cell density alters the nutritional environment of the cell, which, in turn, influences the timing of Plo1 recruitment to the SPB (Fig. 1a), all measurements of Plo1.GFP recruitment were conducted with prototrophs in filter-sterilized supplemented EMM2 at 25 °C. Centrifugal elutriation was used to isolate small G2 cells35. All such synchronized cultures were diluted to a concentration of 1 × 10^7 cells ml^-1 after elutriation. ATP analogues36 (Toronto Research Chemicals, Dalton Pharma Services) were dissolved in methanol to generate 50 mM stock solutions that were subsequently added to cultures. Removal of analogue from the cell environment was achieved by filtration of the cells from culture followed by re-suspension in an identical volume of pre-warmed analogue-free medium.

Microscopy/FACS analysis. Tubulin, Sad1, MPM2 and anti-histone H3 phospho-Ser 10 immunofluorescence and calcifiuor or phalloidin staining were conducted using established procedures37,38,39. Affinity-purified polyclonal Sad1 antibodies37 were used at a concentration of 1 in 25 and the dilution of tissue culture supernatant containing the TAT1 monoclonal antibody40 (gift from K. Gull, University of Oxford) was 1 in 80. Antibodies that recognized histone H3 when phosphorylated on Ser 10 were generated and affinity purified by Eurogentec and used at a concentration of 1 in 100. MPM2 antibody (Upstate 05-368) that recognizes a phospho-epitope on the SPB that is dependent on Plo1 phosphorylation44 was used at a concentration of 1 in 100. For detection of the V5 epitope in Plo1.RL and Cdc2.RL molecules polyclonal antibodies against the epitope (Bethyl Laboratories A190-120A) were used at a concentration of 1 in 100 following fixation with 4% formaldehyde for either 10 (Cut12, Pcp1, Sid4, Cnp1) or 30 (Ctn1, Cut3, Nup85, Nup189) minutes. Thirty minutes in 4% formaldehyde was also used to detect Plo1.RL in Tea1.GFP cells in which the presence of the SV5 epitope in the fusion protein forced us to detect the hybrid molecule with Plo1 polyclonal sera at a concentration of 1 in 100 (ref. 4). DNA content analysis through FACS was conducted using Invitrogen green (Invitrogen) according to published procedures45,46.

Western blotting. Plo1 polyclonal antisera47 were used at a concentration of 1 in 500. PN24 anti-Cdc2 polyclonal antisera were used at a concentration of 1 in 500 to detect Cdc2. Polyclonal antibodies against the SV5 epitope (V5) were used at a concentration of 1 in 500.

Live-cell imaging. All quantitative microscopy was conducted on a DeltaVision system (Applied Precision) fitted with a Zeiss ×100, 1.45 NA objective in conjunction with Softworx (Applied Precision) and Imaris (Bitplane) software. Cells were mounted in a Bioptechs FC52 chamber. The images shown are maximal projections of 20 sections, with 0.3 μm between the slices; images were taken every 3 min. For the FRAP analysis images were taken every second. Qualitative assessment of analogue addition on Plo1.GFP recruitment in cdc2Δ cells was conducted on either DeltaVision or on an Olympus/Photometrics spinning-disc confocal microscope with Metamorph software. Cells were mounted in glass-based culture plates (Iwaki S826-024) and media exchanged by aspiration with a Pasteur pipette. A 532 nm 25 mW laser was used to bleach the red fluorescence signal of the Tomato constructs and a 488 nm 25 mW laser bleached the green fluorescence emanating from Plo1 and Cut12 GFP fusion proteins.

Genetic manipulation. plo1-tds1 was generated as described previously48 with the exception that screening for temperature-sensitive mutants was conducted at 32 °C rather than 36 °C. The plo1Δas8 allele incorporates the canonical as3 mutations49 alongside a mutation of methionine to phenylalanine at position 170. This last change was selected because the crystal structure of human Plk1 AMPPNP (ref. 15) identified this residue as being a major difference between the established Plk1 structure (Plk1.as3 can be strongly inhibited by ATP analogues) and the predicted structure of Plo1 (Plo1.as3 is not strongly inhibited by ATP analogues; data not shown). plo1Δas8 was integrated at the native locus using the marker switch approach49. To create Plo1.RL the S124D and T197D mutations that confer constitutive activation1 were introduced into plo1Δas8, and the polo box domain (position 502–608) was replaced with a GBP (refs 19–21)–SV5 epitope double tag, and integrated the resultant plo1.RL gene into the leu1 locus under the control of the nmt81 promoter. cdc2.RL and cdc2.FRAG15S were created by site-directed mutagenesis and tagged at their C termini with GBP–SV5 before each cassette was introduced into a non-transcribed region of chromosome 3 under the control of the nmt81 promoter. To create cdc2.RLTom and Plo1.RLTom, an NdeI fragment encoding a functional Tomato fluorescence unit (a dimer of two subunits) was inserted in the amino terminus of cdc2.RL and plo1.RL. All mutant alleles were generated by PCR amplification and QuikChange mutagenesis (Stratagene).

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Supplementary information

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Figure S1 pl01 mutants. a) Alignment of the sequences of human Plk1 and Plo1. Plk1 is rendered sensitive to ATP analogues by mutation of the gatekeeper and compensating residues predicted by Zhang et al. while the equivalent pl01.as3 mutation displays only moderate analogue sensitivity (data not shown). The candidates for changes that might enhance the association with AMPPNP in the crystal structure of Plk1 are highlighted. The three arginine residues in the hinge region after the back of the ATP binding pocket that contains the gatekeeper residue (blue) and the phenylalanine at the bottom of the pocket (red). To make the pocket even larger isoleucine 47 was replaced by a smaller glycine. Below the alignment is a summary of the four mutations based on these changes that were introduced into the native pl01 locus and assessed for impact on Plk1 function in response to analogue addition by monitoring the frequency of monopolar spindles in (b). c) The molecular structure of the pl01 and cdc2 alleles, their targeting specificities alongside the scheme used to induce the gene and subsequently activate the “caged” kinase by analogue removal. 3xV5 indicates three copies of the SV5 epitope. The level of Plk1.RL/Cdc2, RL attained in the 24 hour inductions used in all experiments were similar to those of the native kinases (with the exception of Figure 2c). (d) A cartoon of the approach used to generate a mutant library of temperature sensitive pl01 alleles. The significant deviation from the approach taken previously to generate such conditional alleles was setting the initial temperature for screening on plates to 32°C, rather than 36°C. Several highly penetrant mutants were identified following this simple modification. (e, f) Combined Sad1 and tubulin immunofluorescence was used to monitor the appearance of monopolar spindles following shift of an asynchronous culture to the restrictive temperature of 36°C to the restrictive temperature of 36°C at time 0 in the plot in d. f) Tubulin/Sad1/chromatin/DIC micrographs of cells at the 180 min in (d). g) Temperature mediated inactivation of Plk1 and NETO. Datasets from the experiments shown in Figure 3c. Small G2 mutants were isolated from pl01.ts41 (upper) or pl01+ (lower) cultures at time 0. After completion of the first division each culture was split and one half shifted to 36°C (Figure 3c) while the other was maintained at 25°C (right panels). Calcofluor staining was used to score NETO and septation status.
Figure S2  *plo1.RL* localization and impact upon mitotic control. a) Immunofluorescent localization of the *plo1.RL* fusion protein (left panel) in cells harbouring different GFP fusion proteins (central panel (the signal is the endogenous fluorescence of GFP)). The overlap between the two signals can be seen in the false coloured merged image in the right hand panels. In all cases, with the exception of Tea1, the immunolocalisation of *plo1.RL* in the left panel is via the detection of 3 SV5 epitopes that were inserted alongside the GBP sequences (Figure 2a). Polyclonal Plo1 antibodies were used to detect the Tea1.V5GFP fusion protein. 

Key to target locations: *sid4* encodes a fission yeast restricted SPB component. *cut3* encodes a condensin subunit. *cnp1.4* encodes the centromeric histone H3 variant cenpA. *tea1* encodes a kelch domain protein that is delivered to cell tips by microtubules where it regulates polarized cell growth. *for3* encodes one of the three fission yeast members of the formin family of actin nucleating proteins. *nup85* and *nup189* encode nuclear core components. *brr6* encodes a nuclear envelope protein that is recruited to the SPB during mitotic commitment to drive integration into the nuclear envelope at mitotic commitment and once more during mitotic exit to co-ordinate the expulsion from the envelope. b-e) Analogue (40µM 3MB-PP1) wash out experiments of the indicated strains as described for Figure 2a. (b) The moderate induction of mitosis following analogue washout in a *sid4.GFP* background was of a similar amplitude to that seen in *pcp1.GFP* background, while no induction was seen when *plo1.RL* was recruited to any other location. Importantly, no induction was seen upon recruitment to Br6 indicating that the timing of *plo1.RL* recruitment to the SPB is critical. The increase in mitotic index in the *cnp1.GFP* background arose from compromised chromosome segregation as lagging chromosomes and chromosomes falling off spindles were frequently observed following analogue removal (data not shown). Similarly, cytokinesis was severely disrupted by analogue removal when *plo1.RL* was recruited to the actin binding protein coronin (crn1.GFP (Figure 2a) data not shown). (c) MPM2 staining to monitor Plo1 activity at SPBs 20 minutes after analogue removal in the experiments shown in panel b and Figure 2a. Analogue removal clearly induced Plo1 activity on the G2/early mitotic SPBs (single dots). (d) Analogue wash out experiments of the indicated strains as described for Figure 2a with the exception that the *plo1.RLX* allele (Supplementary Figure 1c) that was used in these experiments lacked the S124D and T197D mutations that conferred constitutive activation (yet retained the analogue sensitizing mutations). No mitotic induction was observed, indicating that Plo1 activity at the SPB is normally subject to tight control to prevent premature mitotic commitment. (e) Calcofluor staining to monitor NETO state (left) and MPM2 staining to monitor Plo1 activity at SPBs (right) as described in Figure 3d. In stark contrast to the data with the constitutively active plo1.RL allele, analogue removal had no impact upon NETO or Plo1 activity on the G2/early mitotic SPBs (single dots) when the constitutive activation mutations were not present in the ectopically expressed chimera.
Figure S3  *cdc2.RL* localization and impact upon mitotic control.  
(\textit{a}) Immunofluorescent localization of the *Cdc2.RL* fusion protein (left panel) in cells harbouring different GFP fusion proteins (central panel (the signal is the endogenous fluorescence of GFP)). The overlap between the two signals can be seen in the false coloured merged image in the right hand panels. Immunolocalisation of *Cdc2.RL* is via the detection of the 3 V5 epitopes that had been fused to the end of the GBP sequences (Supplementary Figure 1c).  
(\textit{b}) Analogue wash out experiments of the indicated strains scoring the frequency of mitotic commitment as described for Figure 2b, e. 
(\textit{c}) Photobleaching of *Cdc2.RL*Tom anchored *Cut12.NEGFP* reveals stable associations with the SPB. 
(\textit{d}) MPM2 staining to monitor Plo1 activity at SPBs after analogue removal in the experiments shown in Figure 2b. 1 MPM2 dot represents Plo1 activity on the single SPB foci of late G2/early mitotic cells, while 2 MPM2 dots arise from staining of the separated SPBs in the period between SPB separation at the start of prophase until the metaphase anaphase transition. Analogue removal induced Plo1 activity on the G2/early mitotic SPBs (single dots) when the *Cdc2.RL* chimera had been recruited to the SPB via *Cut12.NEGFP*, but had no impact when no recruitment occurred. 
(\textit{e}) Analogue wash out experiments of the indicated strains scoring the frequency of NETO status as described for Figure 4 b,c. 
(\textit{f}) Analogue wash out experiments of the indicated strains as described for Figure 4b,c with the exception that the *cdc2.RLY15* allele that was used in these experiments, while retaining the F84G analogue sensitizing mutation, contained the wild type Y15 in place of the F15 mutation of *cdc2.RL* (Supplementary Figure 1c). This *cdc2.RLY15* allele therefore retained the ability to be subject to inhibitory phosphorylation by Wee1 kinase. Notably, there was no alteration of NETO status in the culture following the removal of analogue from cells in which this chimeric molecule had been recruited to SPBs via *Cut12.NEGFP*, establishing that MPF activity on the SPB is normally restrained by Wee1/Mik1 activity to prevent premature triggering of NETO. 40\textmu M 3MB-PP1 was used for the Analogue wash out experiments presented here.
**Figure S4** The MPF activity threshold for NETO induction lower than for mitotic induction. The control experiments for Figure 4a. (a) The same size selected culture as shown in Figure 4a. The 25°C panel shows the phenotypes from initial size selection until the splitting of the culture into four equal aliquots at 240 minutes, All cultures were shifted to 32°C to exploit the moderate temperature sensitivity of the cdc2.as mutation and the addition of the ATP analogue 3-BrB-PP1 was used to compromise Cdc2 activity, as indicated, in the right hand panels. (b) Wild type control treatment that mirrored the cdc2.as regime for panel revealed that the addition of the ATP analogue 3-BrB-PP1 had no impact upon the timing of NETO even though 20 µM does lead delay the timing of mitotic commitment to a small degree. (c) A repeat of the experiment presented in Figure 4a in which size selected early G2 cells were allowed to transit one cell cycle before the early G2 cells were split into three aliquots and treated with the ATP analogue 3-BrB-PP1, the solvent methanol, or subjected to no treatment as indicated. The FACS profiles of DNA content show that the two analogue treated cultures remain in a G2 arrested state for the duration of the experiment. This shows that the inhibition of NETO does not arise from a secondary consequence of premature commitment to DNA replication which then blocks NETO via a checkpoint mechanism.
Supplementary Movies:

Movie 1: Plo1.GFP recruitment in cut12<sup>+</sup> and cut12.s11
Time lapse sequences of Plo1.GFP signals in cut12<sup>+</sup> (left) and cut12.s11 (right) cells.

Movie 2: Photobleaching of Plo1.GFP signal at G2 SPBs; cut12<sup>+</sup> and cut12.s11
Time lapse sequences of Plo1.GFP signals on cut12<sup>+</sup> (left) and cut12.s11 (right) SPBs following laser mediated photobleaching.

Movie 3: Plo1.GFP leaves the SPB of cdc2.as cells in a mixed cdc2.as cdc2<sup>+</sup> culture
The panel on the left identifies the cdc2<sup>+</sup> (red lectin) cells visualised in the first frame of the movie of this mixed culture. “+ATP analogue” indicates the timing at which 25mM 1NA-PP1 was added. Analogue remained present for the rest of the sequence. Analogue addition promoted the departure of the Plo1.GFP signal from the SPBs of cdc2.as but not the neighbouring cdc2<sup>+</sup> cells.

Movie 4: Plo1.GFP leaves the SPB of cdc2.as cells in a mixed cdc2.as-cdc2<sup>+</sup> culture at 32°C
The panel on the left/right identifies the cdc2<sup>+</sup> (red lectin) cells visualised in the first frame of the movie of this mixed culture. “+ DRUG” indicates the timing of analogue addition (25mM 1NA-PP1). Analogue remained present for the rest of the sequence. Analogue addition promoted the departure of the Plo1.GFP signal from the SPBs of the longer cdc2.as but not the neighbouring cdc2<sup>+</sup> cells. The moderate temperature sensitivity of the cdc2.as cells delayed mitotic commitment at 32°C.

Movie 5: Photobleaching reveals a stable association of Cut12.NEGFP anchored Plo1.RLTom and Cdc2.RLTom
Two cells are shown for each strain. Each cell shows a single focus of SPB associated fluorescence. The SPB of one of the cells was bleached. Each sequence runs for 20 minutes. There was no recovery. A single focal plane was captured every 4 seconds for 20 minutes for each sequence. Bleaching employed a 532nm 25mW laser to bleach the red fluorescence signal of the “Tomato” constructs and 488nm 25mW to bleach the green fluorescence of the “GFP” Cut12.NEGFP signal.