CDK control pathways integrate cell size and ploidy information to control cell division

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Abstract

Maintenance of cell size homeostasis is a property that is conserved throughout eukaryotes. Cell size homeostasis is brought about by the co-ordination of cell division with cell growth, and requires restriction of smaller cells from undergoing mitosis and cell division, whilst allowing larger cells to do so. Cyclin-CDK is the fundamental driver of mitosis and therefore ultimately ensures size homeostasis. Here we dissect determinants of CDK activity in vivo to investigate how cell size information is processed by the cell cycle network in fission yeast. We develop a high-throughput single-cell assay system of CDK activity in vivo and show that inhibitory tyrosine phosphorylation of CDK encodes cell size information, with the phosphatase PP2A aiding to set a size threshold for division. CDK inhibitory phosphorylation works synergistically with PP2A to prevent mitosis in smaller cells. Finally, we find that diploid cells of equivalent size to haploid cells exhibit lower CDK activity in response to equal cyclin-CDK enzyme concentrations, suggesting that CDK activity is reduced by increased DNA levels. Therefore, scaling of cyclin-CDK levels with cell size, CDK inhibitory phosphorylation, PP2A, and DNA-dependent inhibition of CDK activity, all inform the cell cycle network of cell size, thus contributing to cell-size homeostasis.
Introduction

Cells display homeostatic behavior in maintaining population cell size by controlling cell size at mitosis\(^1\)-\(^4\). This homeostasis is driven by larger cells being more likely to divide than smaller cells, resulting in the correction at cell division of cell size deviations\(^1\),\(^5\),\(^6\). Cyclin dependent kinase (CDK\(^{\text{Cdc2}}\)) is the master regulator of mitosis and cell division, and therefore the propensity for smaller cells not to divide must ultimately feed into the regulation of CDK activity\(^7\).

CDK activity is subject to several mechanisms of control: cyclin synthesis, and subsequent binding of cyclin to CDK which drives CDK into a catalytically competent form\(^8\); Wee1 kinase and Cdc25 phosphatase act to inhibit or activate CDK respectively through regulatory tyrosine phosphorylation\(^9\)-\(^11\); and PP2A phosphatase works to remove phosphates deposited by CDK reducing its net activity\(^12\)-\(^17\), and also controls the phosphorylation state of Wee1 and Cdc25 to regulate the level of CDK tyrosine phosphorylation\(^18\)-\(^21\). Finally, the CDK control network also co-ordinates cell division with cell growth through an unknown mechanism that responds to cell ploidy, with cell size generally doubling as ploidy doubles\(^3\).

It is likely that potential size control pathways will be integrated at the level of CDK activity control because CDK activity is the driver of mitosis. For example, in the fission yeast Schizosaccharomyces pombe, it has been proposed that size control was mediated by the DYRK kinase Pom1, which ultimately inhibits mitotic onset by causing the inhibitory tyrosine phosphorylation of CDK by signaling through the Wee1 kinase\(^22\),\(^23\). However, in both the absence of Pom1 itself or the absence of inhibitory tyrosine phosphorylation, cells are able to maintain cell size homeostasis\(^7\),\(^24\). Thus there must exist alternative mechanisms by which fission yeast cells integrate cell size information into the CDK control network.

Much of our understanding of Cyclin-CDK network regulation has been derived from in vitro studies, but these have limitations when considering cellular parameters such as cell size\(^17\),\(^25\)-\(^27\). Here, therefore, we have studied in vivo regulation of Cyclin-CDK activation at mitosis in the fission yeast. Using a novel CDK activity sensor, we have monitored cell size, CDK activity, and cyclin-CDK complex level simultaneously, whilst genetically varying
regulators of the Cyclin-CDK control system. We propose that CDK activity regulation through inhibitory tyrosine phosphorylation and PP2A work synergistically to communicate information about cell size to the CDK control network. Further, we show that cyclin-CDK complex level scales with cell size, and this aids in the prevention of division in small cells. Finally we show that in cells lacking PP2A and inhibitory tyrosine phosphorylation, haploid and diploid cells of equivalent size and similar Cyclin-CDK concentration have differing Cyclin-CDK activities, with diploid cells exhibiting a lower activity. This suggests that Cyclin-CDK activity is increased in cells of lower ploidy. These experiments inform our understanding of the regulation of Cyclin-CDK, and illuminate how cell size is integrated into this regulatory network.

Results

Given the complexity of the CDK regulatory network, we have used fission yeast cells containing a reduced CDK control system, with the cell cycle driven by a monomeric cyclin-CDK fusion-protein (C-CDK). This simplifies the CDK control network by eliminating cyclin binding to CDK as a regulatory component, and allows co-expression of both cyclin and CDK from a single promoter. This C-CDK fusion protein is expressed under the Cyclin B<sup>Cdc13</sup> promoter, and therefore C-CDK expression mimics endogenous Cyclin B expression. Using this system, inhibitory Wee1-dependent phosphoregulation can also be removed using a non-phosphorylatable C-CDK<sup>AF</sup> mutant. These C-CDK<sup>AF</sup> strains are healthy and viable, but have markedly distinct cell cycle profiles from C-CDK<sup>WT</sup> expressing strains, as they spend a significantly longer period in G1 than C-CDK<sup>WT</sup> cells. Nevertheless, C-CDK<sup>AF</sup> cells co-ordinate cell division with cell growth, and maintain cell-size homeostasis (Figure 1a).

To examine the relationship between cell size, C-CDK concentration, and mitosis, we performed quantitative fluorescence time-lapse microscopy on strains expressing C-CDK<sup>WT</sup> and C-CDK<sup>AF</sup> fluorescently tagged with YFP (Figure 1a-e) (Figure 1-figure supplement 1, figure supplement 2a). This analysis showed clear oscillations of C-CDK<sup>WT</sup> and C-CDK<sup>AF</sup>, with degradation of C-CDK occurring just before cell division (Figure 1b). C-CDK<sup>AF</sup> oscillations were more variable, and 5% of the C-CDK<sup>AF</sup> cells trigger C-CDK degradation in the absence of division (Figure 1-figure supplement 2), similar to what has been observed in CDK1<sup>AF</sup>.
expressing human cells. In both backgrounds, C-CDK concentration scaled with cell size, with C-CDK\textsuperscript{WT} exhibiting a higher amount of C-CDK at mitotic entry compared to C-CDK\textsuperscript{AF} (Figure 1c). On investigating the links between the probability of a given cell to divide, cell size, and C-CDK level, we found that for C-CDK\textsuperscript{WT} both cell size and C-CDK level reach sharp thresholds at which cell division rates increase (Figure 1d,e). In the absence of tyrosine phosphorylation a sharp threshold for C-CDK\textsuperscript{AF} levels still present (Figure 1e), but occurs at a lower level than C-CDK\textsuperscript{WT}. C-CDK\textsuperscript{AF} cells fail to generate a sharp threshold for cell size, but even without a clear size threshold, C-CDK\textsuperscript{AF} cells still restrict smaller cells from division (Figure 1d).

C-CDK level is not a direct measure of C-CDK activity because of the multiple regulatory networks affecting CDK\textsuperscript{25}. To investigate CDK activity, cell size, and C-CDK level at the same time we developed an in vivo single-cell assay of CDK activity. We used Cut3, the Smc4 homolog, as a CDK activity biosensor, because it translocates from the cytoplasm into the nucleus upon CDK-dependent phosphorylation of a single site in its N-terminus (Figure 1f).\textsuperscript{29} Thus, the Cut3 nuclear/cytoplasmic (N/C) ratio can be used to assess CDK activity, a method that has been applied to other protein kinases\textsuperscript{30,31}. As a test of this assay, we blocked cells expressing fluorescently tagged Cut3 in the background of a bulky ATP-analogue sensitive C-CDK\textsuperscript{32} using 1NM-PP1, and tracked single cells following their release from G2 arrest into a range of 1NM-PP1 doses (Figure 1g) (Figure 1-figure supplement 3). The response of the maximum nuclear Cut3 concentration to 1NM-PP1 was similar to the one measured in our previous phosphoproteomics study\textsuperscript{33}, confirming that the sensor reflects in vivo CDK activity (Figure 1h). Subsequently, we examined CDK activity in unperturbed cells measured by the Cut3 N/C ratio, and showed that it both rises to a higher level in C-CDK\textsuperscript{WT} cells in comparison to C-CDK\textsuperscript{AF} cells, and also that progress through mitosis in C-CDK\textsuperscript{AF} cells is slower and more variable (Figure 1i) (Figure 1-figure supplement 4).

We next investigated the links between C-CDK protein levels, CDK activity, and cell size in C-CDK\textsuperscript{WT} and C-CDK\textsuperscript{AF} cells, which have been enlarged beyond their physiological cell size. During a G2/M block (Figure 1g), cell size and C-CDK enzyme concentration (as measured by C-CDK-YFP fluorescence intensity) scaled with each other in both backgrounds (Figure 1j,k). After the release from CDK inhibition, C-CDK\textsuperscript{WT} activity correlated well with both cell size
and C-CDK protein level (Figure 1l,n). However, peak C-CDK<sup>AF</sup> activity correlated better with protein level than with cell size (Figure 1m,o). The link between cell size and CDK activity was much clearer for C-CDK<sup>WT</sup> than for C-CDK<sup>AF</sup> in these low throughput time-lapse assays (Figure 1m). Therefore we repeated this experiment using a high throughput assay based on imaging flow cytometry (Figure 1-figure supplement 5, 6) and observed that peak CDK activity in both C-CDK<sup>AF</sup> and C-CDK<sup>WT</sup> was clearly size dependent (Figure 1-figure supplement 5e). Thus, CDK tyrosine phosphorylation helps to inform the cell division machinery of cell size (Figure 1d,l). However, in the absence of tyrosine phosphorylation, C-CDK<sup>AF</sup> cells are still able to generate a threshold C-CDK level for division and prevent small cells from division (Figure 1e,o) (Figure 1-figure supplement 5e).

A complication of the above assay is that cell size scales with C-CDK level<sup>6,7,34</sup> (Figure 1c, j, k). To uncouple cell size from C-CDK level, and study if small cells are prevented from entering mitosis due to low C-CDK level or for some other reason, we developed a more flexible single cell CDK assay system. This assay was also based on Cut3 translocation into the nucleus (Figure 2a) but used a brighter synthetic C-CDK activity sensor, synCut3-mCherry to allow its co-detection with C-CDK in a high-throughput assay (Figure 2-figure supplement 1). This sensor was expressed in a strain where the endogenous CDK network can be switched off using a temperature sensitive CDK1 allele, cdc2<sup>TS</sup>. A tetracycline-inducible C-CDK tagged with Superfolder GFP (sfGFP) was constructed and made non-degradable<sup>35</sup> as well as sensitive to inhibition by 1NM-PP1. Induction of C-CDK at the cdc2<sup>TS</sup> restrictive temperature allows the study of the activity of the inducible C-CDK without either wild-type CDK activity or C-CDK-sfGFP proteolysis during mitosis. Using this assay, we acquired hundreds of thousands of images of single cells, which allowed us to study the in vivo biochemistry of CDK activity in response to a wide range of C-CDK concentrations in physiologically-sized cells. C-CDK level was uncoupled from cell size as induction of C-CDK was not dependent on cell size (Figure 2b,c). Results from this assay demonstrated that in vivo CDK activity was dependent on C-CDK level, and was reduced when CDK activity was inhibited using 1NM-PP1 (Figure 2d) (Figure 2-figure supplement 2).

Combining this system with genetic backgrounds in which major C-CDK regulation was altered, we analysed how mechanisms of CDK regulation affected C-CDK activity in relation
to cell size. We performed the assay in backgrounds lacking the major PP2A catalytic subunit (PP2A<sup>ppa2Δ</sup>)<sup>12,13</sup>, inhibitory CDK tyrosine phosphorylation, or both (Figure 2e). Following endogenous CDK1 inactivation after temperature shift, both PP2A<sup>+</sup> and PP2AΔ cells arrest in an almost uniform G2 state, ensuring that downstream analysis is not confounded by cells arresting in different phases of the cell cycle (Figure 2f). C-CDK levels increased similarly upon induction in all mutant backgrounds (Figure 2g). Population mean C-CDK activity was comparable between all conditions (Figure 2h), however C-CDK activity displayed differences at the single-cell level when activity was measured in cells of different sizes. In all genetic backgrounds at the same level of C-CDK enzyme, maximum C-CDK activity increases with cell size (Figure 2i). This is particularly noticeable when directly comparing the maximum C-CDK activity of cells with a C-CDK level of ~750 AU in the 8 μm bin to the 14 μm bin in all backgrounds (Figure 2i, dashed lines). The single cell dose-response of CDK activity on C-CDK<sup>WT</sup> concentration background is clearly bistable, with cells existing in either an ‘on’ or an ‘off’ state. The mean CDK activity is relevant directly for strains expressing C-CDK<sup>AF</sup>, as these cells exhibit little bistability in CDK activation. In the C-CDK<sup>WT</sup> expressing cells, there are two population distributions demonstrating bistability. We averaged the two population means as the gradient of this line shows the degree of bifurcation between the lower and the upper CDK activity populations. The C-CDK concentration required to switch cells ‘on’ decreases with increasing cell size, and the sharpness of the transition increases with size (Figure 2i,k). This bistable behavior is heavily dependent on CDK tyrosine phosphorylation (Figure 2i,k,l). Removal of PP2A allows the attainment of the ‘on’ state at lower cell sizes (Figure 2i), effectively shifting the C-CDK dose response curve towards lower sizes without altering the shape of the response (Figure 2k). In addition, PP2A also adds switch like behavior to the C-CDK activity dose-response, as bistable behavior with C-CDK<sup>AF</sup> is not present with C-CDK<sup>AF</sup> PP2AΔ (Figure 2i dashed box, inset and 2l).

When looking across all size bins, maximum C-CDK activity increases with cell size in all genetic backgrounds, but plateaus at about 12-13 μm in the absence of tyrosine phosphorylation (Figure 2j). However, it is clear that cell size is able to regulate C-CDK activity even in the absence of both tyrosine phosphorylation and PP2A (Figure 2i,j). These results are consistent with our previous observations (Figure 1), that although tyrosine...
phosphorylation has a role in informing the cell cycle machinery of cell size, small cells are still restricted from mitosis when tyrosine phosphorylation is absent.

Inhibitory tyrosine phosphorylation directly results in a reduction of intrinsic CDK activity\textsuperscript{36}, whilst PP2A has a dual mechanism of CDK activity modulation: PP2A is able to regulate inhibitory tyrosine phosphorylation by controlling the phosphorylation state of Wee1 and Cdc25\textsuperscript{18}, and in addition can directly oppose the phosphorylation of CDK substrates\textsuperscript{37}. We therefore sought to calculate the contributions of tyrosine phosphorylation and PP2A in restricting CDK activity, both in contexts with and without tyrosine phosphorylation. This was carried out to examine if their combined contribution was greater than the sum of their parts. To calculate the individual contributions of tyrosine phosphorylation and PP2A in restricting C-CDK activity, first we measured the threshold C-CDK level required for 50% of cells to reach a C-CDK activity determined as being $>$5 in arbitrary units (See Figure 3 legend) in different strain backgrounds within different size bins (Figure 3a). This value was chosen as an approximate value of the C-CDK concentration required \textit{in vivo} to trigger mitotic entry in wild-type cells (Figure 1i). When this C-CDK threshold level was plotted across all size bins (Figure 3b) the threshold was seen to be size dependent in all strain backgrounds, with wild-type cells exhibiting the strongest capacity to raise the C-CDK level threshold for mitosis in smaller cells. By subtracting the curves of cell length vs. mitotic C-CDK level (Figure 3c) for various backgrounds we were able to estimate the contributions of tyrosine phosphorylation and PP2A in a given background. For example, C-CDK\textsuperscript{WT} PP2A\Delta - C-CDK\textsuperscript{AF} PP2A\Delta, estimates the ability of tyrosine phosphorylation alone to restrict mitotic entry in a background lacking PP2A. Inhibitory tyrosine phosphorylation alone is able to restrict cells with 600 units of C-CDK from entering mitosis at 8 $\mu$m cell length, but only 200 units of C-CDK at 10 $\mu$m (Figure 3c, yellow). If the different components of the CDK control network act separately, adding individual threshold contributions together would generate a threshold curve similar to the wild-type curve. However, when the contributions of tyrosine phosphorylation and PP2A were added to the C-CDK\textsuperscript{AF} PP2A\Delta curve, they did not recapitulate the wild-type curve (Figure 3d). Thus, this analysis suggests that there is synergy between the tyrosine phosphorylation network and PP2A activity, and that this synergy is important for establishing the C-CDK level threshold for division.
We have shown that small cells are normally prevented from division by their low C-CDK protein level (Figure 1) along with PP2A and tyrosine phosphorylation working synergistically to increase the level of C-CDK needed to trigger division in smaller cells (Figure 3). Strikingly however, in the absence of these major regulators, small cells are still able to restrict division by lowering CDK activity as a result of some other factor related to cell size (Figure 2h,i,j). This unknown factor is able to lower CDK activity in small cells despite high C-CDK levels, thus restricting them from division (Figure 2i).

Given the positive relationship between maximum C-CDK activity and increasing cell size in the C-CDK\textsuperscript{AF} PP2A\textDelta mutant (Figure 2i), we hypothesized that cells dilute a CDK inhibitor as they grow\textsuperscript{1}, perhaps through a titration based mechanism. Cell size is linked to ploidy through an unknown mechanism, and so we tested whether DNA concentration could influence CDK activity, and therefore be a candidate for the unknown factor able to lower C-CDK activity in small cells. We induced C-CDK\textsuperscript{AF} in haploid and diploid variants of the C-CDK\textsuperscript{AF} PP2A\textDelta strain, thereby eliminating all major CDK regulation at mitosis (Figure 4a). Both haploid and diplod cells were present almost uniformly in G2 after endogenous CDK1 inactivation, as cells with 2C and 4C DNA content respectively (Figure 4b), and expressed C-CDK\textsuperscript{AF}-sfGFP in a largely size independent manner (Figure 4c). Strikingly, diploid cells exhibited lower C-CDK activity in response to the same C-CDK enzyme concentration as haploids (Figure 4d). The EC50 of the diploid dose response curve was almost double that of the haploid (Figure 4e). Looking at single-cell, volume-resolved data, the inhibition of C-CDK activity is most marked in smaller diploid cells, with larger diploid cells having almost indistinguishable dose-response curves from their haploid equivalents (Figure 4f). The effect of cell size on CDK activation is much less marked in larger than normal haploids (Figure 4g). The diploids, which feature cells of physiological diploid size, still experience DNA concentration dependent inhibition of their CDK activity. The effect of equal C-CDK levels resulting in lower C-CDK activity in small diploids when compared to equivalent sized haploids is readily seen from the raw images (Figure 4h). Therefore, cells of different ploidies, but otherwise equivalent volume, experience variable CDK activity in response to equal C-CDK level. This suggests that even in the absence of all major CDK regulation, DNA concentration is able to lower CDK activity and prevents division in small cells. At higher
volumes this inhibition of CDK activity disappears, and so the regulation may operate through titrating out an inhibitor.

Discussion

The Cyclin-CDK complex, and its role in controlling mitotic onset, has been studied in many model eukaryotes from yeast\textsuperscript{38}, through marine invertebrates\textsuperscript{39}, to mammalian cells\textsuperscript{40}. A number of regulatory components have been shown to be conserved across these model systems, including the CDK activating Cyclin B, inhibitory CDK tyrosine phosphorylation, and the CDK-counteracting PP2A phosphatase which both opposes CDK substrate phosphorylation and regulates CDK inhibitory phosphorylation though the Wee1/Cdc25 control loop. Despite extensive study, these studies have yet to reveal a fully satisfactory mechanism for cell size homeostasis at the onset of mitosis. To improve our understanding of this control system, we have focused on how CDK activity itself is directly regulated in the context of cell size. Our approach has demonstrated that three mechanisms inform the cell cycle control network of cell size through CDK activity control: C-CDK enzyme concentration scaling with cell size, synergistic PP2A and tyrosine-phosphorylation dependent C-CDK threshold scaling, and DNA concentration dependent inhibition of CDK activity. Our results demonstrate that C-CDK activity vs. C-CDK level dose-response curves previously demonstrated in vitro operate in vivo, but in addition we show they are dependent on cell size in vivo\textsuperscript{26}. We also demonstrate a link between ploidy and CDK activity, with higher ploidy causing a reduction in CDK activity. We propose that CDK activity can be inhibited by a DNA-related mechanism in keeping with early work showing that increasing DNA content delays mitosis, with removal of DNA by irradiation causing acceleration of the following mitosis\textsuperscript{41-43}. Our experiments show that DNA inhibits CDK activity more in smaller cells, potentially reducing CDK activity by a titration mechanism. This may be related to the mechanism by which cell size is linked to ploidy across cell types\textsuperscript{43-46}. Finally, we show that tyrosine phosphorylation, PP2A activity, and DNA dependent inhibition of CDK activity act together to restrict small cells from division, forming a mechanism to generate the robust cell size threshold behavior observed in normal cells.
Our observations suggest that cell size control over the onset of mitosis involves several molecular mechanisms. If it is assumed that the accumulation of the C-CDK cyclin chimera driven by the cyclin promoter mimics the accumulation of cyclin, then one mechanism is the accumulation of cyclin through the cell cycle which scales with the increase in cell size. A second is a synergistic interaction between the inhibitory CDK tyrosine phosphorylation pathway and the PP2A phosphatase, which acts on both the tyrosine phosphorylation pathway and dephosphorylation of CDK substrates. The third is a DNA-concentration dependent inhibition of CDK activity. Given the conservation of all these molecular regulators, these mechanisms can be expected to have direct relevance in other eukaryotic cells.

**Materials and Methods**

**S. pombe genetics and cell culture**

S. pombe media and standard methods are as previously described. After nitrogen and glucose addition, EMM was filter sterilised. This process allows for the generation of clear un-caramelised media. Nutritional supplements for auxotrophic yeast strains were added at a concentration of 0.15 mg/ml. Temperature-sensitive mutant strains were grown at temperatures as specified in the text. The temperature-sensitive allele of Cdc2 (CDK) used was Cdc2-M26. To modulate inducible promoters, anhydrotetracycline hydrochloride (Sigma) in DMSO at specified concentrations was added to 0.03125 μg/ml final concentration unless otherwise specified. To alter Cdc2(as) activity, 1NM-PP1 diluted in DMSO was used at concentrations specified in the text. To stain for septa, calcofluor (Fluorescent Brightener 28 (Sigma Aldrich)) was made up in water at 1 g/L and stored as 500x stock. SynCut3 was constructed by Gibson assembly of a codon optimised fragment consisting of the first 528 amino acids of Cut3, a linker region, and a fluorescent protein (mCherry or mNeongreen). YFP was tagged onto C-CDK at the C-terminus of the protein. Where the sfGFP labelled C-CDK was used, the sfGFP was present internally within the Cdc13 component. Cut3-mCherry was generated by C-terminal tagging and Cut3-GFP was developed previously. Details of the TetR promoter and linearised variants can be found in a previous publication.
Microscopic imaging

All imaging was performed using a Deltavision Elite (Applied Precision) microscope – an Olympus IX71 wide-field inverted fluorescence microscope with a PLAN APO 60x oil, 1.42 NA objective and a Photometrics CoolSNAP HQ2 camera. To maintain specified temperatures during imaging, an IMSOL incubator Environment control system and an objective heater was used. SoftWoRx was used to set up experiments. 5 z-stacks were acquired, with 1 μm spacing. Image analysis was performed using custom Matlab scripts that executed the steps outlined in Figure 1, figure supplement 1.

The ONIX Microfluidics platform allows for long-term time-lapse imaging of live cells. Plate details can be found at http://www.cellasic.com/ONIX_yeast.html. 50 μl of cell culture at density 1.26x10^6/ml was loaded into the plate, and imaged in the 3.5 μm chamber. Cells were loaded at 8 psi for 5 seconds. Media was perfused at a flow rate of 3 psi. The imaging chamber was washed with media for 1 minute at 5 psi before cells were loaded.

Mattek glass bottom dishes were used for some time-lapse imaging applications with drugs that were incompatible with Cellasics plates, primarily for the purpose of release from a 1NM-PP1/Cdc2(as) cell cycle block. Dishes were pre-treated with soybean lectin to permit cell adherence (Sigma Aldrich). Before addition of cells Mattek dishes were pre-warmed on a heatblock at appropriate temperature. Cells were grown and blocked in liquid culture before 2 ml were pelleted (5000 rpm/30 seconds). Cell pellets were then pooled and resuspended in 1 ml of release media (at which time a stop watch was started) in a new microcentrifuge tube before pelleting (5000 rpm/30 seconds) and resuspended in 5 μl of media. This concentrated cell suspension was then applied to the centre of the Mattek dish, and allowed to settle for ~5 seconds. The dish was then washed with 1 ml of release media 3x. The dish was then filled with 3 ml of release media before rapid imaging. In general the wash process required 1.5 minutes, and imaging setup requires 5 minutes for ~8 fields of view. The levels of Cut3 and C-CDK referred to in the paper reference the concentration of both these proteins. This was measured by finding the mean intensity of the brightest group of 9 pixels within the cell, to give a measurement of the concentration.

Imaging flow cytometry
Imaging flow cytometry was performed with an Imagestream Mark X two-camera system (Amnis), using the 60x objective. Cells were concentrated by centrifugation (5000 rpm/30 seconds) and resuspended in ~25 μl of media before sonication in a sonicating water bath. In focus single-cells were then gated by applying the following gating methods in sequence:

1. Gradient RMS>65 (a measure of cell focus).
2. Area/Aspect ratios consistent with single cells.

To avoid any autofocus based drift within an experiment, cell were imaged at fixed, empirically determined focal points, designed to maximise the number of cells with gradient RMS>65. Data was analysed using custom Matlab scripts. The steps these scripts executed were similar to the image processing pipeline for widefield imaging (Figure 1, figure supplement 1) albeit slightly simplified given the presence of only a single cell per image.

The Imagestream acquires two brightfield images of a cell, allowing definition of a cell region using the standard deviation of pixels between the two. This is analogous to the approach used for timelapse cell region segmentation (Figure 1, figure supplement 1).

A line was then drawn through the middle of the cell, by finding the middle pixel on the horizontal axis of the cell. The line was widened by one pixel either side, and the mean fluorescence intensity within that line extracted. The standard peak finding algorithm within Matlab was then used to identify nuclear Cut3 fluorescence (either a dip in the case of a low CDK activity cell or a peak in the case of a high CDK activity cell). Background or “cytoplasmic” fluorescence was defined by the intensity of the flat regions of the curve either side of the peak. The levels of Cut3 and C-CDK referred to in the paper reference the concentration of both these proteins. This was measured by finding the mean intensity of the brightest group of 9 pixels within the cell, to give a robust measurement of the concentration.

To perform time-lapse imaging flow cytometry (IMS), water baths at specified temperatures for the experiment were set up with cultures next to the IMS. Time was measured from the point of drug addition to liquid culture or as described during a wash protocol for drug release. Samples were collected as above from the waterbath, and sample time-points defined as the time at which acquisition on the IMS began (as opposed to time when sample
was collected – although this was consistently ~3 minutes apart). Samples were imaged for ~1 minute unless otherwise stated.

**Data analysis and plotting**

**Boxplots**

The top of box is the 25th percentile of the data, the bottom is the 75th percentile. The line in the middle of the box is the median. Whisker lengths are either the distance to the furthest point outside of the box, or 1.5x the interquartile range, whichever is lower. If data exists that is greater than 1.5x the interquartile range from the top or bottom of the box, this is shown as a red “+”.

**Statistical testing**

Statistical testing was performed where appropriate using a two tailed two sample t-test. P values below 0.05 were considered significant. Replicates are shown where appropriate by N numbers.

**Cell size measurement**

Cell size was measured by three different metrics. In timelapse microscopy assays, cell size was determined as the area of the 2D surface segmented by our segmentation algorithm. In the high-throughput imagestream assays, cell size was measured as length of the cell. The difference in metric choice between these two systems was due to improved ability of measuring cell length in the high-throughput assay, where it was less affected by focal dependent changes in cell volume. In the haploid vs. diploid experiments, a measure of cell volume was used, where cells were assumed to behave as cylinders, and volume was calculated from the measured radius and length. This was done as diploids are wider than haploids and thus a simple length metric cannot be employed for size binning.

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410 ^Cdc2(F84G)

412 §Cdc2(F84G, K79E)

413 *TetR1 – CMVP:TetOx1:TetR-tup11Δ70 (Described originally by Patterson et al.6)

414

Data availability statement

Analysed data and custom scripts have been made available on Figshare, and can be accessed with the handle: 10779/crick.14633037

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References


Wilson, E. B. The cell in development and heredity. 3d edn, (The Macmillan company, 1925).


Figure 1: Cell size and C-CDK concentration dictate probability of division and CDK activity in C-CDK\textsuperscript{WT} and C-CDK\textsuperscript{AF} cells

a Schematic of major components influencing C-CDK activity at mitosis, and in red the pathways that do not influence C-CDK\textsuperscript{AF}. The negative relationship between C-CDK activity and cell growth refers to the block of cell length extension in mitosis. PP2A opposes CDK activity by dephosphorylating CDK substrates, and also by opposing the activation of CDK at mitosis by opposing the phosphorylation of Wee1 and Cdc25. Reciprocally, CDK causes the downregulation of PP2A activity in mitosis.

b Example cell lineage traces from timelapse microscopy. Cell size in pixels\textsuperscript{2} is given in orange, and C-CDK-YFP fluorescence intensity is given in purple. Steep decreases in cell size traces correspond to cell division.

c Scatter plot of mean C-CDK level vs. cell size from timelapse microscopy data. C-CDK level is a measure of C-CDK-YFP fluorescence intensity. Colours indicate density of data. Inset boxplot is mean nuclear C-CDK concentration immediately prior to degradation at anaphase. Boxes represent IQR, with whiskers delimiting 5\textsuperscript{th} to 95\textsuperscript{th} percentiles. C-CDK\textsuperscript{WT} n=28, C-CDK\textsuperscript{AF} n=44 full cycles.

d Plot of the probability of division at the next timepoint (P(Div)) vs cell length for CDK\textsuperscript{WT} and CDK\textsuperscript{AF}. Cells were followed through timelapse microscopy with measurements taken each frame. P(Div) defined as the proportion of cells that undergo C-CDK degradation at anaphase by the next timepoint, given as rate per minute. Points represent cells binned by size, with points plotted at bin centre. C-CDK\textsuperscript{WT} n=685, C-CDK\textsuperscript{AF} n=961 timepoints.

e Plot of P(Div) function vs C-CDK level for CDK\textsuperscript{WT} and CDK\textsuperscript{AF}. C-CDK\textsuperscript{WT} n=685, C-CDK\textsuperscript{AF} n=961 timepoints. C-CDK-YFP intensity measurements taken every frame from timelapse microscopy, and binned by C-CDK level.

f Schematic of Cut3 as a CDK activity reporter. Mitotic CDK dependent phosphorylation of Cut3 on T19 results in nuclear translocation of the protein.

g Experimental outline of block and release timelapse experiment for panels (h),(j)-(o). Asynchronous cells possessing an analogue sensitive (as) CDK were blocked in G2 using 1 \textmu M 1NM-PP1 for 5 hours, and then released into a range of 1NM-PP1 concentrations. Cells were then followed and monitored for their Cut3-tdTomato nuclear/cytoplasmic (N/C) ratio (C-CDK activity) and C-CDK-YFP level using fluorescence timelapse microscopy (see methods). Data for panels (l)-(o) were acquired 15 minutes following release from 1NM-PP1.

h Maximum CDK activity (normalized against maximum level, obtained by release into DMSO) against 1NM-PP1 concentration. Red points are the median of the data sets for each drug concentration (N=324), green point is median in DMSO. Black line is the Hill equation fit to the median data by a nonlinear fitting algorithm (IC50=115.4, Hill coefficient=-1.71).
Purple dashed line is Hill curve derived from Swaffer et al. (2016) dose response data (IC50=133.4, Hill coefficient=-1.47).

i Timelapse quantification of CDK activity in asynchronous cells. Traces are aligned so that 0 minutes corresponds to peak Cut3-tdTomato N/C ratio. Curve smoothing could move Cut3 peak earlier/later than exactly 0 min. Trace colour indicates cell size. Red X indicates automatically defined mitotic entry point. C-CDK<sup>WT</sup> n=23 and C-CDK<sup>AF</sup> n=14.

j Scatter plot of C-CDK-YFP levels against cell size. Experiment described in (g), with measurements taken before release from 1NM-PP1 block. Black points indicate binned data, bin window size 500 pixels<sup>2</sup>. n=324. Pearson correlation coefficient: 0.55.

k As in (j), but with C-CDK<sup>AF</sup>, n=312. Pearson correlation coefficient: 0.62.

l Scatter plot of peak Cut3-tdTomato level vs cell size. Experiment described in (g), with measurements taken 20 minutes after release from 1NM-PP1 block into DMSO. Black points indicate binned data, bin window size 500 pixels<sup>2</sup>. Points are coloured by YFP C-CDK levels at release. n=83. R<sup>2</sup> = 0.5040. Pearson correlation coefficient: 0.50

m As in (l), but with C-CDK<sup>AF</sup>, n=81. R<sup>2</sup> = 0.2150. Pearson correlation coefficient: 0.22.

n Scatter plot of peak Cut3-tdTomato level vs. C-CDK-YFP intensity level 20 minutes after release from 1NM-PP1 block into DMSO. Black points indicate binned data, bin window size 15 AU. Points are coloured by cell size at release. n=83. R<sup>2</sup> = 0.3668. Pearson correlation coefficient: 0.60

o As in (n), but with C-CDK<sup>AF</sup>, n=81. R<sup>2</sup> = 0.5501. Pearson correlation coefficient: 0.74.
Figure 2: Cell size is able to modulate CDK activity independently of canonical CDK regulation

a Experimental outline for figure for panels (b)-(d). Cells were held at 36°C for 1 hour to ablate the function of the temperature sensitive (TS) cdc2 allele. C-CDK-sfGFP expression was induced by addition of tetracycline, and ectopic C-CDK concentration and CDK activity were measured by sequential sampling during induction. Induced C-CDK-sfGFP lacks its degron box sequence, and therefore is not degraded at anaphase. Sequential sampling during C-CDK-sfGFP induction begins at the point of tetracycline addition, with roughly one sample taken every 3 minutes after the start of C-CDK production. Sampling is conducted using an imaging flow cytometer (IMS).

b Expression of C-CDKWT from point of tetracycline addition. Different coloured lines represent different size bins. Black dots represent mean C-CDK-sfGFP level over all size bins for given timepoint. After lag period of ~1000 seconds after tetracycline addition, samples are taken roughly every 3 minutes. n=759633.

c Scatter plot of cell length vs. C-CDK-sfGFP levels. Coloured by density of data points. Data collected throughout induction. n=759633.

d Mean CDK activity dose response against C-CDK-sfGFP in the presence of annotated levels 1NM-PP1. Circles represent average CDK activities across all cells from a single sample taken after induction. 0 nM n=166081, 125 nM n=60759, 250 nM n=165128, 500 nM n=135670 and 1000 nM n=231995.

e Experimental outline for panels (f)-(k). Cells were held at 36°C for 1 hour to ablate cdc2TS function. After 1 hour, C-CDKWT or C-CDKAF fused to sfGFP was induced with tetracycline in cells with either the major PP2A catalytic subunit (encoded by the ppa2 gene) deleted or present. Induced C-CDK-sfGFP lacks its degron box sequence, and therefore is not degraded at anaphase. Sequential sampling during C-CDK-sfGFP induction begins at the point of tetracycline addition, with timepoints taken roughly every 3 minutes after 1000 second lag period in C-CDK-sfGFP induction.

f Flow cytometric DNA content analysis for wild-type cells, cdc2-M26 cells and cdc2-M26 PP2AΔ cells. The major PP2A, ppa2, was deleted in PP2AΔ cells. Cells were fixed for sampling after the block lengths specified in (e), before the addition of tetracycline.

g Induction of C-CDK after tetracycline addition. Points represent mean concentration of C-CDK-sfGFP across all size bins at indicated time points. CDKWT n=166081. C-CDKWT PP2AΔ n=175247. C-CDKAF n=177292. C-CDKAF PP2AΔ n=174847.

h C-CDK activity against C-CDK-sfGFP level in given genetic backgrounds defined in (g). Points represent mean C-CDK activity of all cells. Data is pooled from experiment in (e), from all time points following tetracycline induction. Key is the same as (g).
i Violin plots of single cell C-CDK-sfGFP level against CDK activity in annotated size bins and strain backgrounds. Solid line through violin plot indicates the mean CDK activity within the C-CDK level bin.

j Maximum mean CDK activity vs. cell length in annotated strain backgrounds. Max mean CDK activity is the maximum mean CDK activity within a C-CDK fluorescence level bin for a given cell size. The mean CDK activity level across all fluorescence bins is shown by the solid line in the violin plots in panel (i).

k Maximum gradient of the mean lines in panel (i) plotted against cell length. Maximum gradient of change is derived from a spline fit to the mean CDK activity vs. C-CDK-sfGFP level trace.

l Linear regression lines were fit to data in (k), and residuals were plotted (actual value – predicted value). Non-linear residuals indicate bistability in CDK activation.
Figure 3: CDK Tyrosine phosphorylation and PP2A act synergistically to restrict division in small cells

a Scatter plots of C-CDK level against CDK activity. Either C-CDK$^{WT}$ or C-CDK$^{AF}$ fused to sfGFP was induced in backgrounds with PP2A either lacking or present. PP2AΔ refers to a deletion of the $ppa2$ gene. Red line indicates the C-CDK-sfGFP level at which 50% of cells have a CDK activity greater than 5. Black dashed line marks CDK activity of 5. Data taken from Figure 2i.

b C-CDK-sfGFP level at which 50% of cells have C-CDK activity > 5. Data is taken from (a) across all size bins. Y-axis represents the C-CDK-sfGFP threshold at which 50% of cells will have a C-CDK activity of 5. Dashed lines indicate values where this C-CDK-sfGFP threshold level is undefined due to the threshold being unattainable in experimental conditions.

c Piecewise dissection of the amount of C-CDK-sfGFP a particular component of the cell cycle network is able to prevent from switching to an ‘on’ state (C-CDK activity level of 5) in different size bins. Bar chart shown is of subtractions of curves described in key (from inset). For example, C-CDK$^{WT}$ - C-CDK$^{AF}$ gives the C-CDK threshold tyrosine phosphorylation alone (in a background with PP2A present) is able to generate to restrict C-CDK activation. Values that are undefined due to undefined original threshold values from (a) are taken to be 1000 units, and are marked above the axis (pink).

d Cell length against C-CDK level threshold of annotated curves. Here, a synthetic threshold curve is built (pink), by adding the individual component regulatory contributions of CDK tyrosine phosphorylation (panel (c), yellow) and PP2A (panel (c), orange) to the base curve of C-CDK$^{AF}$ PP2AΔ (green) to try and re-capitulate the WT behaviour (blue). Dashed line indicates undefined threshold values.
Figure 4: Cellular DNA content inhibits CDK activity independently of tyrosine phosphorylation or PP2A activity

a Experimental outline for panels (b)-(h). PP2A Δ/Δ diploids and PP2AΔ haploids were arrested using cdc2TS. PP2AΔ refers to a deletion of the ppa2 gene. Diploids were held at 36°C for 1 hour, whilst haploids were held for 3 hours to generate blocked cell populations with similar cell volumes despite ploidy differences. C-CDKAF expression was induced by addition of tetracycline, and C-CDKAF-sfGFP concentration and CDK activity were measured by sequential sampling from time of induction in an imaging flow cytometer.

b Flow cytometric DNA content analysis for wild-type cells, haploid cdc2-M26 PP2AΔ cells and diploid cdc2-M26/cdc2-M26 PP2AΔ/PP2AΔ cells. PP2AΔ refers to a deletion of the ppa2 gene. Cells were fixed for sampling after the block lengths specified in (a), before the addition of tetracycline.

c Expression of C-CDKAF fused to sfGFP from point of tetracycline addition in haploid and diploid strains. Different coloured lines represent different size bins. Haploid n=125021, Diploid n=139557.

d Mean CDK activity against C-CDKAF-sfGFP level in haploids and diploids. Solid line is a sigmoid fit to data.

e EC50 from sigmoid curves in (d). Haploid EC50: 372 AU. Diploid EC50: 663 AU. Haploid EC50 is 56% of diploid EC50.

f Violin plots of single cell C-CDKAF-sfGFP level against CDK activity in annotated volume bins and ploidy status. Solid line through violin plot indicates the mean CDK activity within the C-CDK-sfGFP level bin. Volume bins span a physiological range of diploid cell sizes. Volume bin 17 corresponds to a haploid cell length of 12.1 μm and a diploid cell length of 9.53 μm. Volume bin 36 corresponds to a haploid length of 18.7 μm and a diploid length of 14.4 μm.

g Mean intra volume-bin dose response of C-CDK-sfGFP level vs. CDK activity in annotated ploidy level. Lines are sigmoid curves fit to raw data. Cell volume bin indicated by line colour.

h Example raw images from experiment. Brightfield (BF) channel displaying cell morphology, C-CDK-sfGFP channel and synCut3-mCherry CDK activity indicator are shown. C-CDK level is the same across all images. Scale bars = 3 μm.
**Supplementary Figure Legends**

**Figure 1 - Figure Supplement 1: Automated image analysis pipeline for widefield imaging.**

a Definition of the initial gross cell region and removal of excess background. A threshold was automatically computed using the Otsu algorithm from the standard deviation projection image. The blue region shows the binary mask of all pixels above this threshold.

b The top z-stack image is both normalised and filtered to remove excess background and intensity variations using a Gaussian filter that removes most of the cell character, but retains information on gross brightfield intensity differences. The original image is then divided by this filter, homogenising the signal from the image. Once the brightfield image is processed, the previously generated cell region mask is applied. To generate an initial mask of single cells, a threshold is computed on the pixels within the initial cell region mask.

c. True cell-cell contacts exhibit concavities at both sides of any false contact, and the presence of the concavities can be used to license potential “cut” lines. To generate potential cut lines a watershed algorithm is applied to the distance image of the initial single cell segmentation mask.

d To remove any spuriously segmented background regions the standard deviation of the difference between the top and bottom z-stack images are used. A difference image generates heterogeneous pixel values in cell regions but has almost no effect on background regions. An example of an image with this standard deviation metric quantified in each mask can be seen is given, where non-cell masks feature a low value.

e An example of the final segmented image after steps outlined in panels (a)-(d).
Figure 1 - Figure Supplement 2: Fluorescence time-lapse quantification of C-CDK dynamics in unperturbed cell cycles

a Schematics of C-CDK<sup>WT</sup> and C-CDK<sup>AF</sup> regulation by Wee1 kinase and Cdc25 phosphatase. C-CDK<sup>AF</sup> has T14 mutated to A and Y15 mutated to F to mimic constitutive dephosphorylation of both residues. Example images of a FOV from time-lapse movie is shown. Cells were grown in a Cellasics microfluidics plate following 2 days of culture in YE4S at 32 °C. C-CDK-YFP is seen in purple. Scale bar=10 μm.

b Purple lines indicate C-CDK levels (mean nuclear concentration) and yellow indicates cell size (measured by cell mask area in pixels<sup>2</sup>). Cell mask and lineage tracing generated by Pomseg and Pomtrack (see methods). DD=Double dip cell, hDD=half double dip cell. DD cells undergo complete cyclin degradation without cell division. hDD cells undergo incomplete cyclin degradation without division. Trace marked (a) represents an aberrant cycle in a C-CDK<sup>WT</sup> expressing cell.

c Boxplot of C-CDK oscillation period. Period was calculated by measuring the peak to peak (P2P) distance on the autocorrelation function of each C-CDK level lineage trace. C-CDK<sup>WT</sup>, N=32; C-CDK<sup>AF</sup>, N=57. Box represents median value delimited by 25<sup>th</sup> and 75<sup>th</sup> percentiles. See methods for outlier points.

d Boxplot of intra-lineage standard deviation of period length. C-CDK<sup>WT</sup>, N=32; C-CDK<sup>AF</sup>, N=57. Box represents median value delimited by 25<sup>th</sup> and 75<sup>th</sup> percentiles. See methods for outlier points.
Figure 1 - Figure Supplement 3: A time-lapse block and release assay to measure the effect of CDK inhibition on CDK activity in single cells

a Experimental outline for panels B-G. 1NM-PP1 sensitive C-CDK\textsuperscript{WT} and C-CDK\textsuperscript{AF} cells are blocked by addition of 1NM-PP1. C-CDK\textsuperscript{AF} cells were block for longer (7 hours against 5 hours) to allow cells to reach a similar size distribution as C-CDK\textsuperscript{WT} cells. Cells were then released into a range of 1NM-PP1 concentrations. After release, images were acquired every minute. Time between washing and image acquisition is ~5 minutes. Cells were grown in EMM at 32°C.

b Left: Schematic demonstrating that as cells are blocked at G2/M, they continue to grow and accumulate C-CDK but do not translocate Cut3 into the nucleus or alter their levels of Cut3. Right: Density plot demonstrates the overlap population cell lengths of C-CDK\textsuperscript{WT} and C-CDK\textsuperscript{WT} cells after variable block times.

c Black traces indicate raw data. Red traces indicate exponential curve fit to data. Photobleaching curves were derived from the 1000 nM release using C-CDK\textsuperscript{WT}-YFP and Cut3-tdTomato. All subsequent measurements were corrected for photobleaching from derived curves.

d Images of Cut3-GFP channel from representative FoV ~25 minutes after release from a 1 μM block into indicated drug concentrations.

e Plots of nuclear Cut3-GFP levels against time after release over a range of 1NM-PP1 concentrations. Lines are coloured by cell size at T=0 of the release.

f Single cell C-CDK-YFP traces in DMSO and 20 nM of release. Red x indicates end of anaphase. Traces are coloured by cell size at Time=0. Only traces which undergo anaphase are shown. End of anaphase defined as first time-point at which C-CDK-YFP trace is equal to post anaphase YFP plateau level +10 AU.

g Boxplot of anaphase time in WT and AF strains. Anaphase time is calculated as end of anaphase time – peak Cut3 time. Difference is non-significant. C-CDK\textsuperscript{WT}, N=69 and C-CDK\textsuperscript{AF}, N=47. Lower panel, scatter plot of anaphase time vs cell size, with strain indicated by colour. Box represents median value delimited by 25\textsuperscript{th} and 75\textsuperscript{th} percentiles. See methods for outlier points.
Figure 1 - Figure Supplement 4: Cut3-GFP as a marker of CDK activity in WT and AF cell strains

a Still images of Cut3-GFP tagged in strains expressing C-CDK\textsuperscript{WT} and C-CDK\textsuperscript{AF}. Cells were grown in a Cellasics microfluidics device in YE4S at 32°C. Scale bar=10 μm.

b Example cell length and Cut3-GFP single cell lineages. Quantification is performed by Pomseg and Pomtrack (see methods). Cut3-GFP nuclear/cytoplasmic (N/C) ratio is calculated by dividing mean cytoplasmic Cut3 intensity by mean nuclear Cut3 intensity after background subtraction. Orange lines= cell size, green lines= CDK activity (measured by Cut3 N/C ratio).

c Montage of tagged C-CDK\textsuperscript{WT} and C-CDK\textsuperscript{AF} strains from time-lapse. Colour outline indicates strain and is derived from Pomseg based segmentation of the brightfield image. Scale bar=5 μm.

d Boxplot of mitotic times in C-CDK\textsuperscript{WT} and C-CDK\textsuperscript{AF} strains. Mitotic time is calculated as peak time – mitotic entry time. Difference is significant by two sample t-test (p=0.006). Box represents median value delimited by 25\textsuperscript{th} and 75\textsuperscript{th} percentiles. See methods for outlier points.

e Boxplot of cell size at mitotic entry (cell size sampled at red x position in Figure 1i). Note high variability in the C-CDK\textsuperscript{AF} population (CoV=0.18 vs 0.08 in WT). Box represents median value delimited by 25\textsuperscript{th} and 75\textsuperscript{th} percentiles. See methods for outlier points.
Figure 1 - Figure Supplement 5: An imaging flow cytometry assay reveals that size, C-CDK level and tyrosine phosphorylation dictate the rate and timing of CDK activation at mitosis

a Schematic of the high-throughput imaging flow cytometry block and release assay. Cells are arrested in G2 using 1NM-PP1 for various lengths of time, before being washed of 1NM-PP1 and sampled on an imaging flow cytometer.

b Representative images of single cells with computed cell masks overlaid on fluorescent Cut3 images in red. Top row of images is from the brightfield channel of the top row of fluorescent images. Representative images taken from Cut3-GFP cells in EMM at 32°C. Scale bar = 10 μm.

c Experimental outline for panels (D-G). C-CDKWT/AF cells sensitive to the CDK inhibitor 1NM-PP1 are blocked for variable amounts of time. Cells are then washed of 1NM-PP1 and released into mitosis. After release, cells are monitored via sequential sampling using imaging flow cytometry. Block performed using 1 μM 1NM-PP1. Cells were grown in EMM at 32°C.

d Quantification of C-CDK-YFP levels after indicated block time. Colours indicate density of data; yellow represents high density. Red data points indicate mean of binned data, bin widths 0.33 μm.

e Plots of mean CDK activity (as measured by Cut3 N/C ratio) within size bins indicated by line colours. Red dots indicate points of maximum Cut3 N/C ratio change, as derived from the first derivative of a smoothing spline fit to raw data (raw data is shown). Each point on line has >50 cells. N=3000-12000 per time point, with ~400,000 single cell images analysed in total. Background subtraction for N/C ratio performed using wild-type cells lacking Cut3-GFP after indicated block time.

f Maximum Cut3 N/C ratio change against cell size or C-CDK level. C-CDK level is predicted from data in d. Data is taken from 2,3 and 4 hour releases. Black line represents linear regression line.

g Time of maximum Cut3 N/C ratio change against cell size or C-CDK level. C-CDK level is predicted from data in d. Data is taken from 2,3 and 4 hour releases. Black line is the linear regression line. Colours represent the same as panel (f).
Figure 1 - Figure Supplement 6: Size dependent grading of mitotic entry rates and timing are dose responsively dependent on CDK inhibition

a Experimental outline for panels B-D. 1NM-PP1 sensitive C-CDK\(^{WT}\) and C-CDK\(^{AF}\) cells are blocked by addition of 1NM-PP1. C-CDK\(^{AF}\) cells were blocked for longer (7 hours against 5 hours) to allow cells to reach a similar size distribution to C-CDK\(^{WT}\) cells. Cells were then released into a range of 1NM-PP1 concentrations. After release, images were acquired every minute. Time between washing and image acquisition is ~5 minutes. Cells were grown in EMM at 32°C. Cells are sampled during the region marked time-lapse.

b Plots of mean CDK activity (as measured by Cut3-GFP N/C ratio) against time from release in indicated size bins at annotated 1NM-PP1 levels. N=1000-4000 cells per time-point, >10 cells averaged within each bin.

c Plots of maximum Cut3 nuclear translocation rates against cell size in C-CDK\(^{WT}\) and C-CDK\(^{AF}\) cells. Maximum rates were taken from the first derivative of a smoothing spline fit to data in b. Line colours indicate 1NM-PP1 concentration. Key given on the right hand side.

d Plots of time of maximum Cut3 translocation rate timing vs cell size in WT and AF cells. Maximum rates were taken from the first derivative of a smoothing spline fit to data in b. Line colours indicate 1NM-PP1 concentration.
**Figure 2 - Figure Supplement 1: A new synthetic CDK sensor for S. pombe**

**a** Design of the synthetic Cut3 (synCut3) sensor. The design includes the first 528 amino acids of Cut3 (and has previously been shown to translocate into the nucleus at mitosis\(^1\)).

**b** Example images of synCut3-mNeonGreen expressed from the eno101 promoter, in the presence or absence of 1NM-PP1 (for 1 hour) or a mutated T19 residue. The T19V mutation does not allow CDK phosphorylation, therefore preventing nuclear translocation. Scale bar = 20 μm.

**c** Examples images of exogenous synCut3-mCherry and endogenous Cut3-GFP expressing cells. Scale bar = 20 μm.

**d** Detailed view of two mitotic cells expressing both synCut3-mCherry and Cut3-GFP.

**e** Quantification of exogenous synCut3 signal vs endogenous Cut3 nuclear levels. Data points coloured to indicate cell size. Note endogenous Cut3 signal is smoothed to remove foci containing condensed chromatin regions.
**Figure 2 - Figure Supplement 2:** A single cell *in vivo* biochemistry approach permits decoupling of cell size from C-CDK concentration

**a** Experimental outline for panels B-D. Cells were held at 36°C for 1 hour to ablate *cdc2*-M26 function. After 1 hour, C-CDK\textsuperscript{WT} or C-CDK\textsuperscript{AF} was induced with tetracycline. Induced C-CDK lacks its degron box sequence, and therefore is not degraded at anaphase. Sequential sampling during C-CDK induction begins at the point of tetracycline addition. Concurrent with tetracycline addition, 1NM-PP1 was added to the specified concentration to inhibit the induced C-CDK.

**b** Mean CDK activity against C-CDK level, within specified size bins. Colours within subplot indicate cell size bin (see colour bar). Different subplots represent cells released into different 1NM-PP1 concentrations.

**c** Violin plots of single cell C-CDK level against CDK activity data. Individual subplots are the single cell data from a given size bin and 1NM-PP1 level. Rows correspond to the same size bin, columns to the same 1NM-PP1 level. Although bistable behaviour is observed, lines through data represent the population mean C-CDK activity level within a given C-CDK level bin.

**d** Heatmap of annotated features, extracted from the single cell dose response data. Max mean CDK activity is the maximum mean CDK activity within a C-CDK fluorescence level bin. C-CDK slope breadth is the change in C-CDK between the C-CDK bin at which CDK activity is greater than 1.1x of minimum, and less than 0.8x of maximum. C-CDK level when \( P(\text{CDK}>5)>0.1 \) indicates the C-CDK level required to increase CDK activity in 10% of cells to a level greater than 5.

**e** Experimental outline for panels F and G. Cells were held at 36°C for 1 hour to ablate *cdc2*-M26 function. After 1 hour, C-CDK\textsuperscript{WT} or C-CDK\textsuperscript{AF} was induced with tetracycline to different levels by adding variable amounts of tetracycline. C-CDK was induced in the presence of 10 \( \mu \text{M} \) 1NM-PP1 to inhibit the induced C-CDK. After 60 minutes, 1NM-PP1 was washed from cells and cells were sequentially sampled using imaging flow cytometry (IMS). All time measurements are given as time from washing 1NM-PP1.

**f** Scatter plot of C-CDK levels against cell size after C-CDK induction. Data represent pooled data from all cells encompassing all 1NM-PP1 release concentrations. Colours indicate local data point density. \( N>10000 \).

**g** synCut3 N/C ratio (representing CDK activity) against time in the presence of induced C-CDK\textsuperscript{WT} or C-CDK\textsuperscript{AF}. Line colours indicate size bins. \( N>50 \) cells per data point.
Figure 1

a) Proteomics

b) Cell size (pixels²)

c) P(Div) vs. C-CDK

d) P(Div) vs. length

f) Active C-CDK

j) C-CDK activity

k) C-CDK activity

m) C-CDK activity

n) C-CDK activity

h) Normalised Max Cut3 level vs. [1NM-PP1] (nM)

i) C-CDKWT vs. C-CDKAF

l) C-CDKWT vs. C-CDKAF

o) C-CDKWT vs. C-CDKAF

Figure 1
**Figure 2**

**Panel a**

- TetP::C-CDK^WT/AF-sfGFP
- synCut3-mCherry
- cdc2^ts
- 25°C
- + Tetracycline
- 1 hour
- 36°C
- DNA Content

**Panel b**

- Time (seconds)
- C-CDK level (AU)

**Panel c**

- Cell length (μm)
- C-CDK level (AU)

**Panel d**

- C-CDK activity
- [1NM-PP1]

**Panel e**

- TetP::C-CDK^WT/AF-sfGFP
- synCut3-mCherry
- cdc2^ts
- 25°C
- + Tetracycline
- 1 hour
- 36°C
- DNA Content

**Panel f**

- Wild Type
- DNA Content

**Panel g**

- Time (seconds)
- C-CDK level (AU)

**Panel h**

- C-CDK activity

**Panel i**

- Cell length bin
- 8 μm
- 10 μm
- 12 μm
- 14 μm

**Panel j**

- Max mean CDK activity
- Cell length (μm)

**Panel k**

- Max CDK activity gradient
- Cell length (μm)

**Panel l**

- Residuals
- Cell length (μm)
A Image acquisition and initial cell region definition

Brightfield Z-Stack acquisition
Standard deviation projection
Cell region defined by threshold of std. proj.

B Brightfield image processing and first pass thresholding

Unaltered top Z-slice
Gaussian filter of top Z-slice used for intensity normalisation
Intensity normalised, cellular region defined. Image to be used for segmentation.
Otsu thresholding and mask hole filling allow first guess at cell masks

C Concavity licensed watershed cell segmentation algorithm

Example mask requiring segmentation
Masks generated during the "concavity licensed watershed algorithm"
Overlay of proposed watershed lines and concavities
Licensed watershed lines are used to split cells

D Non-cell mask removal

Top Z-slice
Bottom Z-slice
Top/bottom Z-slice background normalised
Standard deviation of difference image intensities within the segmented region allows for true cell identification

E Final masked image

Figure 1, Figure Supplement 1
Figure 1, Figure Supplement 3
Figure 1, Figure Supplement 4
Figure 1, Figure Supplement 6
Endogenous Cut3

T19V

Max Endo-Cut3-GFP (AU)

Max synCut3-mCherry (AU)

Cell size (pixels²)

DMSO 1000 nM DMSO 1000 nM

Fluorescent protein

synCut3-
mNeon
greenCalcofluor

Figure 2, Figure Supplement 1