Mitotic kinase oscillation governs the latching of cell cycle switches

Graphical abstract

Highlights
- Mitotic kinases enforce once-per-cycle occurrence of cell cycle events
- Many cell cycle events can occur periodically in the absence of mitosis
- Are they autonomous oscillators or accidental consequences of faulty regulation?
- Regulatory mutations allow negative feedback loops to sustain oscillations

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In brief
Periodic cell cycle events could be coordinated by an entrainment or a suppression mechanism. Using a mathematical model of the yeast cell cycle, Novak and Tyson show that autonomous oscillations of S-phase kinase and mitotic exit-phosphatase activities are suppressed by a “latching-gate” mechanism imposed by a universal mitotic Cdk oscillator.
Mitotic kinase oscillation governs the latching of cell cycle switches

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SUMMARY

In 1996, Kim Nasmyth1 proposed that the eukaryotic cell cycle is an alternating sequence of transitions from G1 to S-G2-M and back again. These two phases correlate to high activity of cyclin-dependent kinases (CDKs) that trigger S-G2-M events and CDK antagonists that stabilize G1 phase. We associated these “alternative phases” with the coexistence of two stable steady states of the biochemical reactions among CDKs and their antagonists.2,3 Transitions between these steady states (G1-to-S and M-to-G1) are driven by “helper” proteins. The fact that the transitions are irreversible is guaranteed by a “latching” property of the molecular switches, as we have argued in previous publications.4,5 Here, we show that if the latch is broken, then the biochemical reactions can swing back-and-forth across the transitions; either G1-S-G1-S … (periodic DNA replication without mitosis or cell division) or M-G1-M-G1 … (periodic Cdc14 release, without fully exiting mitosis). Using mathematical modeling of the molecular control circuit in budding yeast, we provide a fresh account of aberrant cell cycles in mutant strains: endoreplication in the clb1-5Δ strain6 and periodic release and resequestration of Cdc14 (an “exit” phosphatase) in the CLB2kdΔ strain.7,8 In our opinion, these “endocycles” are not autonomous oscillatory modules that must be entrained by the CDK oscillator5,7 but rather inadvertent and deleterious oscillations that are normally suppressed by the CDK latching-gate mechanism.8

RESULTS AND DISCUSSION

As we have argued elsewhere,2,4,5 cell proliferation (growth, DNA replication, mitosis, and cell division) is an alternating sequence of “one-way” transitions from G1 (unreplicated chromosomes) into S-G2-M (replication and partitioning of chromosomes) and back to G1, which we picture dynamically (Figure 1A) as a series of flips of a bistable switch between two provisionally stable steady states, G1 and S-G2-M. Like a hinged gate with a latch in the “middle” position, the gate must be pushed sufficiently in one direction (by transcription factor, abbreviated as TF) to enter S-G2-M, after which the gate swings back into the latched position. To return to G1, the gate must be pushed sufficiently in the opposite direction (by exit protein, abbreviated as EP), after which it latches again. This latch has the unusual property that the gate must alternate between letting cells enter S-G2-M and letting them leave. The latch catches because of the antagonistic interactions between cyclin-dependent kinases (CDKs) and their antagonists (the ubiquitin ligases and stoichiometric inhibitors that oppose S phase-promoting and M phase-promoting CDKs; Figure 1B). For the gate to latch properly, the mitotic cyclins must oscillate between low activity (G1) and high activity (S-G2-M), and the inducers (TF and EP) must be under negative-feedback control (i.e., TF must be inactivated after the cell enters S-G2-M, and EP must be inactivated after the cell returns to G1).

The G1-to-S transition (activation of TF) and the M-to-G1 transition (activation of EP) are controlled by “checkpoints” called Start (in yeast) and the spindle assembly checkpoint (SAC), respectively. To pass Start, a yeast cell must grow to a sufficiently large size, and to pass SAC, it must properly align its replicated chromosomes on the mitotic spindle. The mechanisms of these checkpoints have been described elsewhere.9

Our concept of the cell cycle as a latching gate has been verified by elegant experiments with budding yeast.10,11 In this paper, we explore how perturbing the latch mechanism by mutation can convert a latching gate (Figure 2A) into a swinging gate (Figure 2B). In many cell types, deletion of mitotic cyclins may induce repeated rounds of DNA replication without mitosis (“endoreplication”12). On the other hand, if a nondegradable mitotic cyclin is expressed in budding yeast (so that a cell cannot properly exit mitosis), then cells exhibit sustained oscillations of a mitotic-EP (Cdc14 phosphatase; hence, “Cdc14 endocycles”17,6). How do these perturbations generate endocycles?

Mathematical model

To answer this question, we investigate a simple model of the budding yeast cell cycle (Figure 1C). At the heart of the network are double-negative interactions between ClbS and ClbM and their inhibitory substrates, Cdh1 and Sic1.13 ClbS is regulated by two negative feedback loops involving its TF, MBF; namely MBF → Nrm1 → MBF and MBF → Cbs → I Cdh1 → I Nrm1 → MBF. ClbM is regulated by the negative feedback loop ClbM → Polo → Cdc14 → Cdh1 → CblM, and Cdc14 participates in its own negative feedback loop Cdc14 → Cdh1 → I Polo → Cdc14.
To simplify the mathematical model, we neglect the ubiquitin ligase activity of APC/C:Cdc20 (a mitotic-exit helper acting in conjunction with Cdc14). Indeed, in budding yeast, Cdc20 is dispensable, as demonstrated by viability of cdc20Δ pds1Δ clb5Δ triple-mutant cells.14 Evidently, the indispensable function of Cdc20 is to degrade Securin (Pds1) and Clb5 (an S-phase cyclin); in the triple mutant, the degradation of mitotic cyclin (ClbM) is dependent solely on APC/C:Cdh1, which is activated by Cdc14. Hence, for simplicity, we consider that Cdc14 is our generic EP.

In the STAR Methods section, we provide a set of ordinary differential equations (ODEs) that describe this simplified model of the budding yeast cell cycle. Numerical integration of the ODEs (Figure 3A) presents time courses that agree reasonably well with the observed fluctuations of cell cycle regulators in wild-type budding yeast cells. Note that we do not include the size control mechanism operating at Start in daughter cells; therefore, these oscillations represent cell division cycles of mother cells. Furthermore, the SAC is not limiting in wild-type yeast cells (Figure 3B). Hence, the latch catches the gate at the G1 steady state.

Having established that the mathematical model gives a reasonable description of the yeast cell cycle, we proceed to draw bifurcation diagrams for the model (Figure 3B), using Cdh1 activity as the dynamical variable and MBF (TF) and Cdc14 (EP) as the bifurcation parameters. The red Z-shaped curve in Figure 3B is the locus of steady-state activity of Cdh1 as a function of MBF (for Cdc14 = 0) to the right and as a function of Cdc14 (for MBF = 0) to the left. The Z-shaped curve here, computed from the mathematical model, confirms the hypothetical Z-shaped curve in Figure 1A.

It is important to recognize that each half of the Z-shaped curve is itself Z-shaped in the following sense. For Cdh1 as a function of MBF (to the right in Figure 3B), the middle branch of the red curve connects to the lower branch at a negative value of MBF, as illustrated in Figure S1A. Because this connection point is at negative activity of MBF, Cdh1 cannot be reactivated as MBF activity falls to zero. This effect is crucial to the “latching” behavior of the control system. As the “gate swings open” and the cell transits from G1 to S-G2-M, the “spring” (the negative feedback on MBF) pulls the gate closed again and the “latch” (the stability of the metaphase steady state) holds the cell in mitosis. To leave mitosis and return to G1, the cell must activate Cdc14 and exactly the same latching mechanism holds for this transition. As Cdc14 activity falls, Cdh1 does not inactivate because the threshold for Cdh1 inactivation occurs at a negative value of Cdc14 activity (Figure S1B). Hence, the latch catches the gate at the G1 steady state.

On the bifurcation diagram (Figure 3B), we also plot the influence of Cdh1 activity on the steady-state levels of MBF and Cdc14. MBF becomes more active as Cdh1 activity increases (the blue curve) because Cdh1 degrades the MBF inhibitor, Nrm1.15 In contrast, Cdc14 activity decreases with increasing Cdh1 (the orange curve) because Cdh1 degrades the Cdc14 activator, Polo.16 We may think of these curves as “pseudo-nullclines”: on the red curves, Cdh1 is at a pseudo-steady state, $d\{\text{Cdh1}\}/dt = 0$; on the blue curve, $d\{\text{MBF}\}/dt = 0$; and on the orange curve, $d\{\text{Cdc14}\}/dt = 0$. We say “0” because the actual rates of change depend on what the other variables of the dynamical system are doing at any particular time in a simulation. Wherever two pseudo-nullclines intersect is a potential steady state of the full set of ODEs; the black dot at the top, at Cdh1 = 1, is a steady state if MBF = 0 and the black dot at the bottom, at Cdh1 = 0.0005, is a steady state if Cdc14 = 0. Taking the simulation in Figure 3A and projecting onto Figure 3B (dashed curves), we see that the pseudo-nullclines act as if they were “guiding” the temporal evolution of the dynamical system. In this case, the simulated mother-cell trajectory does not stop at the checkpoints (the black dots) because, as we have explained, the checkpoints (Start and SAC) are not operational in mother cells.
Cdc14 endocycles

To see how a proper latching gate (Figure 2A) in wild-type cells becomes an aberrant swinging gate (Figure 2B) in mutant cells, we first consider the case of Cdc14 endocycles (Figure 2C) in the mutant strain CLB2kdJ (“KEN and D-box deleted”) encoding a functional, nondegradable Clb2 (ndClbM). Degradation of mitotic cyclins is a necessary requirement for dephosphorylation of mitotic substrates by counter-acting phosphatases. In budding yeast, the bulk of mitotic dephosphorylation is catalyzed by Cdc14, which is transiently released from the nucleolus during mitotic exit.17 In the presence of sustained activity of nondegradable Clb2, exit from mitosis is incomplete and Cdc14 release becomes periodic.7,9

If we express a constant low level of nondegradable ClbM in our model (e.g., ndClbM = 0.1), cell cycle progression is hardly perturbed and mitotic exit is not compromised (Figure S2A). However, at higher levels of ndClbM, the cell is unable to leave mitosis and start a new cell division cycle. For example, at ndClbM = 0.4 (Figure 4A), total ClbM > 0.4 and Sic1 < 0.02; hence, there is sufficient ClbM activity at all times to block relicensing of replication origins. Therefore, the genome cannot be replicated, although ClbS is oscillating. Nonetheless, Cdc14 is periodically inactivated and activated (sequestered in and released from nucleoli) by the negative feedback loop Polo → ndClbM = 0.15. At this value of ndClbM (refer to Figure S3A), the stable steady state (●) collides with the unstable saddle point (○) underneath, and they both disappear. As a consequence, Cdh1 inactivation is no longer dependent on entering a new cell cycle and accumulating Cin and ClbS kinases; rather, the remnant ndClbM-kinase can inactivate Cdh1. In this case, there is only a single, unstable steady state of the control system, surrounded by a limit cycle oscillation (the dashed curve in Figure 4B). The limit cycle, which corresponds to the Cdc14 endocycles in Figure 4A, is independent of Cin and ClbS activities (Figure S2B). The oscillation is a consequence of the fact that as Cdc14 phosphatase activity drops, Cdh1 is inactivated by the remnant ClbM kinase activity, as evidenced by the Cdh1 inactivation-threshold at a positive value of Cdc14. It is an example of an “inhibitor-amplified negative feedback loop oscillator”.18

Cdc14 endocycles persist over a limited range of expression of CLB2kdJ, as observed in the study conducted by Lu and Cross,17 although our range (0.15 < ndClbM < 0.66) is somewhat different from theirs. Because oscillations start where a saddle (o) and node (●) coalesce, the period of oscillation is infinitely long at first and drops significantly as ndClbM increases and the amplitude changes little (Figure S4), as observed.

Endoreplication cycles

In fission yeast cells, shutting off the synthesis of mitotic cyclin leads to periodic rounds of DNA replication in the absence of mitosis.19 These endoreplication cycles are driven by periodic expression of fission yeast’s S-phase cyclin. In budding yeast, deletion of the four mitotic cyclins (Clb1–4) causes a G2 block because of persisting activity of the S-phase cyclin ClbS.20 However, deletion of the CLBS gene supports periodic endoreplication...
cycles in the clb1-5Δ strain,⁶ which must be driven by an oscillation of the other S-phase cyclin, Clb5, in budding yeast (Figure 2D). Interestingly, Clb6 is the only B-type cyclin that is targeted for degradation by an SCF-dependent mechanism,²¹ which evidently allows the relicensing of replication origins in the absence of Clbs 1–5.

We simulate the clb1-5Δ strain by reducing the synthesis of mitotic cyclins (ClbM) to zero. In this case, the cell cycle control network exhibits limit-cycle oscillations of ClbS activity driven by periodic synthesis and degradation (Figure 4C). Cln-kinase activity is maintained at a constantly high level because SBF is constitutively active in the absence of inhibition by mitotic cyclins (Clb1–4).²²

In the absence of mitotic cyclins, the irreversible (latching) nature of the G1-S transition is lost (Figure 4D) because the lower threshold of the Z-shaped Cdh1 balance curve, as a function of MBF, moves to a positive value of MBF activity (Figure S1A shows that the lower threshold moves into positive territory when the synthesis rate of ClbM falls below 20% of its wild-type value). Consequently, the saddle and node points coalesce (Figure S3B), the remaining steady state (at the intersection of the two balance curves in Figure 4D) becomes unstable, and the control system executes limit cycle oscillations (the dashed curve).

**Other periodic events in the absence of mitotic cycles**

Periodic cell cycle events uncoupled from the CDK “master” control system are observed in other circumstances. In the absence of B-type cyclin Cdk1 activity (clb1-6Δ) or in mutants (cdc4 etc.) defective in degrading Sic1 (a stoichiometric inhibitor of Cdk1), cells arrest in G1 with high Cln2-kinase activity and periodic bud formation.²³ Because constitutive Cln2-expression does not suppress this phenotype,²³ it is doubtful that periodic budding is driven by an autonomous oscillator (e.g., the small amplitude fluctuations of Cln2-kinase activity). It is more likely that the high level of Cln2-kinase activity drives continuous accumulation of bud precursors, and buds appear at regular intervals of time as bud primordia are assembled, as in the “structural model” of Fantes et al.²⁴ or the “dripping faucet” model of Tyson et al.²⁵

Duplication of the spindle pole body (SPB) happens concomitantly with bud formation as yeast cells pass through Start. clb1-6Δ cells, although they bud periodically, duplicate their SPB only once in the presence of high Cln-kinase activity.²⁶ In contrast, endoreplicating clb1-5Δ cells, with oscillating activity of Clb6-kinase, overduplicate their SPBs, and overduplication becomes more pronounced in the presence of Cib5 cyclin in clb1-4Δ mutant cells.²⁶ DNA endoreplication is blocked by persistent Cib5 activity in clb1-4Δ cells, indicating that SPB overduplication is independent of periodic DNA replication and oscillating G1- and S-phase Cdk1 activities. This conclusion is consistent with the observation that cells constitutively overexpressing Cln2 or Clb5 reduplicate their SPBs.²⁶ It is unclear whether SPB duplication in budding yeast is driven by (1) an autonomous endocycle similar to Plk4 oscillations driving centriole duplication in Drosophila embryos²⁷ or (2) a “dripping faucet” mechanism, as suggested for centrosome reduplication in S phase-arrested frog egg extracts in the presence of high cyclin E kinase activity.²⁸

Persistence oscillations in energy metabolism can be detected in both budding²⁹ and fission yeast³⁰ after a block of the cell division cycle. These metabolic cycles might be driven by an autonomous oscillator in nondividing cells, but they are clearly coupled with Cdk oscillations in dividing cells, presumably by Cdk1 phosphorylation of metabolic enzymes.³¹,³²

As described here, periodic DNA replication and Cdc14 endocycles in budding yeast are “autonomous oscillators” only when the latching mechanism of cell cycle control fails. During normal cycles, helper proteins push the gate open, then negative feedback pulls the gate closed and the latchcatches the gate at a stable steady state (either G1 or M). In our view, endocycles are inadvertent consequences of this “logic” of cell cycle control, when the latch fails to catch.

Our view is consistent with the proposal of Manzoni et al.⁸ that “events capable of repeating themselves multiple times [like Cdc14 endocycles or endoreplication cycles] are restrained to occur once per cycle by their coupling to the cyclin-Cdk engine.”
It differs from the paradigm suggested by Lu and Cross that "Cdc14 release, and likely other cell cycle processes, is controlled by intrinsically oscillatory modules that are entrained to a single occurrence at appropriate cell cycle positions by cyclin-Cdk cycles through a 'phase-locking' mechanism." We do not view ClbS activation and Cdc14 release as intrinsically oscillatory modules that happen to be entrained by a complementary ClbM oscillator in a phase-locked relationship but rather as inadvertent and deleterious oscillations that are normally suppressed by the CDK/Cdh1 latching-gate mechanism.

STAR METHODS

Detailed methods are provided in the online version of this paper and include the following:

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SUPPLEMENTAL INFORMATION

Supplemental information can be found online at https://doi.org/10.1016/j.cub.2022.04.016.

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AUTHOR CONTRIBUTIONS

Conceptualization, B.N. and J.J.T.; software: B.N.; formal analysis, B.N. and J.J.T.; writing – original draft, J.J.T.; writing – reviewing & editing, B.N. and J.J.T.; visualization, B.N.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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STAR METHODS

KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| Software and algorithms | XPPAUT v8.0 | Ermentrout[^33] | http://www.math.pitt.edu/~bard/xpp/xpp.html |

RESOURCE AVAILABILITY

Lead contact
Further information about modelling should be directed to the lead contact, Bela Novak (bela.novak@bioch.ox.ac.uk).

Materials availability
This study did not generate any new reagents.

Data and code availability
The computer code is provided in Methods S1 and any additional information required to reanalyze the model reported in this paper is available from B.N. upon request.

METHOD DETAILS

A simple model of the budding yeast cell cycle
Our mathematical model provides a description of the cdc20Δ pds1Δ clb5Δ triple-mutant, in which Securin (Pds1), one of the S-phase cyclins (Clb5) and an APC/C activator (Cdc20) are deleted. We assume that the level of Cdc28 (Cdk1) is not limiting and the rate of Cdk1:Cyclin dimer accumulation is driven by the synthesis of different cyclins. The various Cdk1:Cyclin complexes are labelled by their cyclin component. The stoichiometric inhibitor, Sic1, binds to B-type cyclin dimers (Cdk1:Clb) to form inactive trimers. To distinguish these forms, we introduce the notation: ClbM = [Cdk1:ClbM] = ‘free ClbM’, ClbMt = [Cdk1:ClbM] + [Cdk1:ClbM:Sic1] = ‘free ClbM’ + ‘complex with Sic1’, and similarly for ClbS and ClbSt. Furthermore, Clb = ‘free ClbS’ + ‘free ClbM’, and Clb = ClbS + ClbM + ndClbM, where Clb refers to the total Clb concentration, including non-degradable ClbM (ndClbM) in some simulations. To keep the model simple, we assume that Sic1 binds reversibly to both S-phase and M-phase Cdk1 complexes with the same equilibrium dissociation constant (K_{diss}). In this case, the level of inactive Sic1:Cbind complexes is given by:

\[
\text{Sic1Clb} = \frac{2 \cdot \text{Sic1} \cdot \text{Clb}}{B + \sqrt{B^2 - 4 \cdot \text{Sic1} \cdot \text{Clb}}} \]

where \(B = \text{Sic1} + \text{Clb} + K_{diss}\). The overall Clb activity (ClbS + ClbM) is calculated by subtracting the inactive complexes with Sic1 (Sic1Clb) from total Clb (Clb):

\[
\text{Clb} = \text{Clb} - \text{Sic1Clb}.
\]

Since SBF is inhibited only by the mitotic Clb, ClbM needs to be calculated as a fraction of Clb’s not complexed by Sic1:

\[
\text{ClbM} = \frac{\text{ndClbM} + \text{ClbM}}{\text{Clb}} \cdot (\text{Clb} - \text{Sic1Clb})
\]

The synthesis of Cln’s (Cln1 & Cln2) is dependent on the SBF transcription factor, which is regulated by Cln activity (see below):

\[
\frac{d\text{Cln}}{dt} = k_{\text{cln}} \cdot SBF - k_{\text{cln}} \cdot \text{Cln}
\]

The synthesis of S-phase cyclin ClbS (Clb6) is driven by the MBF transcription factor, which is activated by Cln-phosphorylation and inhibited by Nrm1, a transcriptional repressor synthesized by MBF itself. The degradation of both ClbS and Nrm1 is controlled by APC/C in complex with Cdh1:

\[
\frac{d\text{ClbS}_i}{dt} = k_{\text{scibs}} \cdot \text{MBF}_a - (k_{\text{debs}} + k_{\text{debs} \cdot \text{Cdh1}}) \cdot \text{ClbS}_i
\]
\[
\frac{d\text{MBF}}{dt} = k_{\text{MBF}} \cdot (\text{MBF}_{\text{tot}} - \text{MBF}) - k_{\text{MBF}} \cdot \text{MBF} \cdot (Nrm_1 - (\text{MBF}_{\text{tot}} - \text{MBF}))
\]

\[
\frac{dNrm_1}{dt} = k_{\text{MBF}} \cdot \text{MBF} - k_{\text{MBF}} \cdot \text{Cdh1} \cdot \text{Nrm1}.
\]

In these equations, \(Nrm_1 = Nrm + Nrm1MBF\) is ‘free Nrm1’ and ‘complex with MBF’, \(MBF_{\text{tot}} = MBF + Nrm1MBF\), and \(MBF_a = (MBF - \text{Cln} + J_{MBF} + \text{Cln})\) is ‘active MBF’, i.e., the fraction of free MBF that is phosphorylated by Cln (assuming rapid phosphorylation and dephosphorylation of MBF).

The synthesis of M-phase cyclins (Clb1-4) is autacatalytic and their degradation is APC/C:Cdh1-dependent:

\[
\frac{d\text{ClbM}}{dt} = k_{\text{ClbM}} + k_{\text{ClbM}} \cdot \text{ClbM} - (k_{\text{ClbM}} + k_{\text{ClbM}} \cdot \text{Cdh1}) \cdot \text{ClbM}.
\]

The synthesis of Polo-kinase is dependent on the activity of mitotic Cdk1 complexes, and it is degraded in a Cdh1-dependent fashion:

\[
\frac{d\text{Polo}}{dt} = k_{\text{Polo}} \cdot \text{ClbM} - (k_{\text{Polo}} + k_{\text{Polo}} \cdot \text{Cdh1}) \cdot \text{Polo}.
\]

We assume that Sic1 is synthesized at a constant rate and that its degradation is controlled by dual phosphorylation. The first phosphorylation step, which is assumed to be in equilibrium with dephosphorylation, is catalyzed by both Cln- and Clb-kinases, while the second step is Clb-dependent:

\[
\frac{d\text{Sic1}}{dt} = k_{\text{Sic1}} - \left(k_{\text{Sic1}} + k_{\text{Sic1}} \cdot \text{Clb} \cdot \frac{\text{Clb} + \text{ClbM}}{\text{Clb} + \text{ClbM}}\right) \cdot \text{Sic1}.
\]

The switch-like activation and inactivation of three regulatory proteins (SBF, Cdh1 and Cdc14) is described by ‘Goldbeter-Koshland kinetics’. Cln-kinases phosphorylate and inactivate the inhibitor (Whi5) of the SBF transcription factor, i.e., SBF is activated as the variable and \(\text{J}_{\text{MBF}}\) as the bifurcation parameter, with \(\text{J}_{\text{MBF}} = 0\) and calculated the one-parameter bifurcation diagram for \(\text{Cdh1}\) and reactivated by Cdc14 phosphatase:

\[
\frac{d\text{Cdh1}}{dt} = \left(k_{\text{cach1}} + k_{\text{cach1}} \cdot \text{Cdc14}\right) \cdot \frac{(1 - \text{Cdh1})}{\text{J}_{\text{cach1}} + 1 - \text{Cdh1}} - \left(k_{\text{cach1}} + k_{\text{cach1}} \cdot \text{Cdc14}\right) \cdot \frac{(1 - \text{Cdh1})}{\text{J}_{\text{cach1}} + 1 - \text{Cdh1}}.
\]

The nucleolar release of Cdc14 is induced by Polo-kinase:

\[
\frac{d\text{Cdc14}}{dt} = k_{\text{cach1}} \cdot \text{Polo} \cdot \frac{(1 - \text{Cdc14})}{\text{J}_{\text{cach1}} + 1 - \text{Cdc14}} - k_{\text{cach1}} \cdot \frac{\text{Cdc14}}{\text{J}_{\text{cach1}} + \text{Cdc14}}.
\]

Computation

This set of ordinary differential equations was implemented in freely available software XPPAUT in the form of an ‘ode’ file (see Methods S1). The dynamics of the wild-type budding yeast cell cycle were computed for the kinetic parameter values given in Table S1. Time-course simulations were performed by numerical integration of the model using the ‘stiff’ integrator of XPPAUT. The simulation of the endoreplication cycle was performed by setting the rate constants of CibM synthesis \((k_{\text{cibm}})\), and \((k_{\text{cibm}})\) to zero. The calculation of the Cdc14-endocycle was done by setting ndCibM = 0.4.

Calculation of bifurcation diagrams and pseudo-nullclines

The pseudo-nullclines were calculated using the capacity of XPPAUT to plot one-parameter bifurcation diagrams for a reduced set of ODEs. To this end, we first removed MBF and Cdc14 from the system of ODEs and treated them both as parameters. To plot the Cdh1 pseudo-nullcline for the G1/S module, we set Cdc14 = 0 and calculated the one-parameter bifurcation diagram for Cdh1ss as a function of MBF as the bifurcation parameter. To plot the Cdh1 pseudo-nullcline for the M/G1 module, we set MBF = 0 and calculated the one-parameter bifurcation diagram for Cdh1ss as a function of Cdc14. For the MBF pseudo-nullcline of the G1/S module, we removed the Cdh1 and Cdc14 ODEs, treating them as parameters, and calculated the one-parameter bifurcation diagram for the remaining ODEs, with MBF as the variable and Cdh1 as the bifurcation parameter, with Cdc14 = 0. The Cdc14 pseudo-nullcline of the M/G1 module was calculated in a similar fashion.
Supplemental Information

Mitotic kinase oscillation governs the lathing
of cell cycle switches

Bela Novak and John J. Tyson
Figure S1. Bifurcation diagram for the G1/S and M/G1 modules. Related to Figure 3. (A) The steady-state level of Cdh1 as a function of MBF (when Cdc14=0) is Z-shaped, but the lower kink of the ‘Z’ lives in negative territory of MBF, so the switch latches at Cdh1≈0.0005 as MBF falls to zero. As the synthesis rate of ClbM decreases from its wild-type value (100%), the lower kink moves to larger values of MBF and eventually enters positive territory. Now the switch can flip back to Cdh1≈1 as MBF falls. (B) Similarly, the steady-state level of Cdh1 as a function of Cdc14 (when MBF=0) is Z-shaped, with the upper kink at a negative value of Cdc14. As the level of ndClbM increases from zero, the upper kink moves to larger values of Cdc14, eventually entering positive territory, and the switch fails to latch.
Figure S2. Properties of Cdc14 endocycles. Related to Figure 4. (A) Endocycles require a sufficiently high level of non-degradable ClbM. For ndClbM=0.1, the large post-mitotic peak of Cln, whose synthesis by SBF is very sensitive to inhibition by ClbM, indicates that ClbM activity must be very low (close to 0) as these cells exit mitosis. Indeed, the non-degradable fraction of ClbM is inhibited by the peak of Sic1 in early G1. Low ClbM activity allows re-licensing of chromosome replication origins and the subsequent rise of ClbS activity is sufficient to initiate DNA replication and drive viable cell cycles. (B) Endocycles are independent of G1- and S-phase cyclins (Cln=0 and ClbS=0).
Figure S3. Endocycles arise by SNIPER (‘saddle-node infinite-period’) bifurcations. Related to Figure 4. Pseudo-phase plane for Cdc14 endocycles (A) and endoreplication cycles (B), respectively. The red curves plot the locus of Cdh1 steady states for parameter values just below (solid) and just above (dotted) the SNIPER bifurcation points. As the parameter passes through the bifurcation point, a stable node (●) coalesces with an unstable saddle point (○), giving way to a limit cycle oscillation of ‘infinite period’. For the solid and dotted red curves ndClbM=0.1 and 0.15 (A) and the rate of ClbM synthesis is reduced to 21% and 22.5% (B).
Figure S4. Bifurcation diagram for Cdc14 endocycles. Related to Figure 4. We plot the period and amplitude of the limit cycle oscillations as a function of the parameter ndClbM. Notice how the period diverges to infinity as ndClbM approaches the bifurcation point (ndClbM=0.135).
Table S1. Parameter values of the budding yeast cell cycle model. Related to Star Methods.

| Parameter values of the budding yeast cell cycle model |  |
|-------------------------------------------------------|---|
| **Cln synthesis/degradation & SBF regulation:**      |  |
| $k_{scln} = 0.2$, $k_{dcln} = 0.2$, $k'_{asbf} = 1$, $k_{asbf} = 10$, $k_{isbf} = 25$, $J_{sbf} = 1$ |  |
| **ClbS synthesis and degradation:**                   |  |
| $k_{scibs} = 0.15$, $k'_{dclbs} = 0.1$, $k_{dclbs} = 0.05$ |  |
| **MBF regulation by Nrm1:**                          |  |
| $k_{snrm1} = 0.05$, $k_{dnnrm1} = 0.1$, $MBF_{tot} = 0.5$, $k'_{ass} = 1$, $k'_{diss} = 0.001$, $J_{mbf} = 0.01$ |  |
| **ClbM synthesis and degradation:**                   |  |
| $k'_{scibm} = 0.01$, $k_{scibm} = 0.01$, $k'_{dclbm} = 0.01$, $k_{dclbm} = 1$, $J_{clbm} = 0.01$, $n = 2$ |  |
| **Polo synthesis/degradation & Cdc14 regulation:**    |  |
| $k_{spolo} = 0.01$, $k'_{dpolo} = 0.01$, $k_{dpolo} = 1$, $k_{acdc14} = 1$, $k_{icdc14} = 0.25$, $J_{cdcc14} = 0.01$ |  |
| **Sic1 synthesis/degradation & Clb binding:**         |  |
| $k_{ssic} = 0.02$, $k'_{dsic} = 0.01$, $k_{dsic} = 2$, $J_{sic1} = 0.01$, $K_{diss} = 0.05$ |  |
| **CdH1 activation & inactivation:**                   |  |
| $k'_{acdh1} = 1$, $k_{acdh1} = 10$, $k'_{icdh1} = 0.2$, $k_{icdh1} = 10$, $J_{cdh1} = 0.01$ |  |
| **Nondegradable ClbM:**                              |  |
| ndClbM=0 for mitotic and endoreplication cycle and nddbM >0 for Cdc14 endocycle. |  |