A comprehensive model for the proliferation-quiescence decision in response to endogenous DNA damage in human cells

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Abstract

Human cells that suffer mild DNA damage can enter a reversible state of growth arrest known as quiescence. This decision to temporarily exit the cell cycle is essential to prevent the propagation of mutations and most cancer cells harbour defects in the underlying control system. Here we present a mechanistic mathematical model to study the proliferation-quiescence decision in non-transformed human cells. We show that two bistable switches, the restriction point (RP) and the G1/S transition, mediate this decision by integrating DNA damage and mitogen signals. In particular, our data suggests that the cyclin-dependent kinase inhibitor p21 (Cip1/Waf1), which is expressed in response to DNA damage, promotes quiescence by blocking positive feedback loops that facilitate G1 progression downstream of serum stimulation. Intriguingly, cells exploit bistability in the RP to convert graded p21 and mitogen signals into an all-or-nothing cell cycle response. The same mechanism creates a window of opportunity where G1 cells that have passed the RP can revert to quiescence if exposed to DNA damage. We present experimental evidence that cells gradually lose this ability to revert to quiescence as they progress through G1 and that the onset of rapid p21 degradation at the G1/S transition prevents this response altogether, insulating S-phase from mild, endogenous DNA damage. Thus, two bistable switches conspire in the early cell cycle to provide both sensitivity and robustness to external stimuli.

Significance

Controlled transitions of human cells between proliferating and non-proliferating states are essential for normal development and tissue homeostasis. To understand how the decision to proliferate is made in response to positive input from growth factors and negative input from the DNA damage response, we have built a mathematical model of the underlying molecular network, based on data from live-cell imaging experiments. Our model suggests that two major cell cycle transitions are crucial for decision making: the restriction point, which integrates pro- and anti-proliferative signals, and the G1/S transition, which temporarily insulates cells from some aspects of the DNA damage response. Together, our model gives mechanistic insight into how cells maintain both sensitivity and robustness to external signals.

Keywords:
Cell cycle; DNA damage; proliferation-quiescence decision; mathematical modelling; live-cell imaging
Introduction

Living systems strive to proliferate and to protect their genomic information from harm. On the cellular level, these two aims have to be balanced to avoid the accumulation of deleterious mutations that can lead to cell death or cancer. Therefore, cells have evolved mechanisms to postpone proliferation in response to DNA damage. In particular, they can enter a reversible state of growth arrest after mitosis known as quiescence (1), which provides time for DNA repair and prevents the propagation of damage to future generations. Yet, how and when the decision between proliferation and quiescence is made, and how decision-making is implemented on a molecular level, remains elusive.

Cell proliferation is regulated by a series of steps known as the cell cycle, which is driven by cyclin-dependent kinases (Cdks) and their regulatory counterparts, the cyclins (2, 3). The activation of Cdk2 by either cyclin E (CycE) or cyclin A (CycA) controls commitment to proliferation and initiation of DNA synthesis (4). However, Cdk2 activity can be opposed by p21 (Cip1/Waf1), a stoichiometric inhibitor that is under the control of the DNA damage-induced transcription factor p53 (5, 6). Thus, p21 acts as a damage-dependent brake on cell cycle progression. p21 can also bind to proliferating cell nuclear antigen (PCNA), a vital component of the DNA replication machinery (7), where it competes with DNA processivity factors for PCNA binding and thus blocks DNA synthesis directly (8–10). These mechanisms enable p21 to maintain genome stability downstream of DNA damage by preventing the propagation of mutations to future cellular generations, which could otherwise lead to decreased cell fitness or tumorigenesis. This is consistent with observations that p21–/– mouse models show enhanced tumorigenesis (6, 11) and that p21 is required for the maintenance of haematopoietic and forebrain neural stem cells (12, 13).

Using live single-cell imaging, Spencer and colleagues have shown that, following mitosis, proliferating cells bifurcate into one population that rapidly accumulates Cdk2 activity and re-enters the cell cycle and another, quiescent population where Cdk2 activity is downregulated (14). This bifurcation is controlled by both mitogen stimulation and p21 expression (14, 15). Recently, we and others have observed that p21 protein levels show high cell-to-cell variability in genetically-identical, non-transformed cells due to intrinsic DNA damage suffered during unperturbed growth (16, 17). Frequently, this damage occurs during DNA replication causing p21 accumulation in the G2- and M-phases of a mother cell’s cycle. p21 is then inherited by daughter cells at cell division and determines their decision to proliferate. These results suggest that p21 is part of a control system that integrates information across generations to determine cell fate. In addition, the proliferation-quiescence decision appears to be reversible since DNA damage or extrinsic stress applied in G1 can revert cells back to a quiescent state (18). This window of reversibility is thought to be closed by the inactivation of the anaphase-promoting complex / cyclosome (APC/C\(^{Cdh1}\)), a ubiquitin ligase that targets many cell cycle regulators for degradation during G1, including CycA.

Motivated by these experiments, we aimed to gain a systems-level understanding of the proliferation-quiescence decision and its response to endogenous, naturally occurring DNA damage. In a previous study, we developed and experimentally validated a mathematical model for p21 regulation, revealing a bistable switch at the G1/S transition that controls p21 degradation (16). Here, we augment this model with the restriction point (RP), a second major cell cycle transition that gates cell cycle entry. We show that, by promoting p21 synthesis, DNA damage can prevent passage through the RP and that, due to its bistable nature, the RP converts a graded p21 signal into an all-or-nothing cell cycle decision. Our model also suggests that DNA damage in post-RP G1 can reverse the decision to proliferate and reset cells to a pre-RP state. This window of reversibility ends at the G1/S transition when rapid p21 degradation prevents a response to DNA damage, which is vital to avoid premature S-phase exit. Finally, we show that G1 cells progressively lose the ability to revert
to quiescence as they approach the G1/S transition and that the loss of G1 regulators compromises the control system leading to different forms of dysregulated growth. Hence, our study demonstrates how cellular decision-making integrates information over time and how bistable switches can make cells both sensitive and tolerant to external stimuli.

Results

Developing a model for early cell cycle regulation

To understand the control of cell cycle entry and subsequent progression, we built a mechanistic mathematical model of the underlying regulatory network (Fig. 1A). Our model links the two main transitions of early cell cycle regulation: the RP, where cells commit to proliferation, and the G1/S transition marking the onset of DNA replication (see SI Appendix for details). The G1/S module of the model is based on our previous work on p21 dynamics (16) extended here by explicitly distinguishing between CycE and CycA. G1 progression was modelled as follows: Passage through the RP requires growth factor-dependent hyper-phosphorylation of the retinoblastoma protein (Rb), which alleviates inhibition of E2F transcription factors (19). Once active, E2Fs drive their own expression as well as the synthesis of CycE and CycA, both of which promote Cdk2 activity, further reinforcing Rb inhibition (20, 21). E2F-dependent transcription also causes expression of the early mitotic inhibitor (Emi1), an inhibitor of APC/C^Cdh1 (Cdh1), which is responsible for degradation of CycA during G1. Increasing levels of CycE:Cdk2 and CycA:Cdk2 then activate replication complexes (aRCs), initiating S-phase. DNA damage can halt this process by upregulating p53, which causes p21 expression, inhibiting both Cdk2 activity and DNA replication (5, 6). This is counteracted by two p21 degradation pathways depending on either Cdk2 or PCNA in aRCs (22).

Deterministic simulations of this model show a distinct peak in CycE:Cdk2 levels, as well as an abrupt increase in CycA:Cdk2, at the G1/S transition (Fig. 1B), in good agreement with quantitative live-cell imaging data (23). Here, the accumulation of CycA is facilitated by the rapid inactivation of Cdh1 at S-phase entry, which is caused by both Emi1 expression and increased Cdk2 activity. We also observe the presence of p21 both before and after S-phase, while the inhibitor is absent during DNA replication, as has been shown experimentally (24). To capture the large cell-to-cell heterogeneity in p21 protein levels that has been reported, we performed stochastic simulations (Fig. 1C, top). We assumed that DNA damage occurs randomly throughout the cell cycle but is more likely to occur during S-phase, when chromosome replication takes place (25). The resulting simulations recapitulate both the spread and general pattern of experimental p21 levels (Fig. 1C and D) as well as the length of G1- and S-phase (Fig. 1E), suggesting that we can use the model to further analyse p21 dynamics in the early cell cycle.

Our model attributes the heterogeneity in p21 levels primarily to the stochasticity of DNA damage, i.e., the spontaneous occurrence and subsequent repair of DNA lesions. However, we also observe a dependence of p21 expression on the cell cycle stage (Fig. 1F and G). In particular, DNA damage in G1-phase causes an immediate increase in p21 (Fig. 1F), while S-phase cells do not show such an increase and delay p21 accumulation until after the completion of DNA replication (Fig. 1G). As previously described, this is due to the presence of a bistable switch at the G1/S transition ((16) and SI Appendix, Fig. S1A, B). This switch promotes rapid p21 degradation during DNA replication (SI Appendix, Fig. S1C), while it allows p21 levels to vary in G1 and G2-phase. Thus, intrinsic DNA damage that occurs during S-phase results in elevated p21 protein levels in G2 and these elevated levels may persist for some time, even after the damage has been repaired (Fig. 1F).
Fig. 1 | A model of early cell cycle progression. (A) Influence diagram of the cell cycle model. Components associated with the RP and G1/S transition are shown separately for visual clarity, but are connected in the model. GF, growth factors; aRC, active replication complexes. (B) Deterministic simulation of the model in A. (C) p21 levels from stochastic simulations (top; n = 30) and experiments in individual hTert-RPE1 cells (bottom; n = 29). (D) Maximum p21 level in G1- and G2-phase in stochastic simulations (Sim; n = 200) and experiments (Exp; n = 186). Values were normalised to the median in G1-phase for each case. A two-sided Wilcoxon rank sum test shows no significant difference between experiment and simulation (G1: p = 0.65; G2: p = 0.33). (E) Comparison of G1- and S-phase lengths in stochastic simulations (Sim; n = 200) and experiments (Exp; n = 186). A two-sided Wilcoxon rank sum test yields p = 0.78 (G1) and p = 0.64 (S). (F, G) Stochastic simulations of two cells that suffer DNA damage in different cell cycle phases.

Inherited p21 promotes quiescence by blocking passage through the RP

Having observed that DNA damage generated during DNA synthesis can lead to increased p21 expression only after S-phase exit, we wanted to understand how this affects cell cycle progression. Single-cell imaging data during unperturbed growth suggests that a moderate increase in p21 levels in G2- or M-phase does not impair progression through these stages (Fig. 2A). Instead, p21 protein is transferred from mother to daughter cells, where it can promote quiescence (16). Consistent with this, quiescent cells have been shown to
contain high amounts of p21, as well as increased levels of hypo-phosphorylated Rb (14). The Rb protein is an essential component of a positive feedback system that creates the RP (Fig. 2B). An inherent property of positive feedback is its ability to create discontinuous switches (26) and a bistable Rb-E2F switch has been reported to underlie the RP (27). Our model supports this notion showing bistability in E2F activity in the absence of p21, with low concentrations of growth factors attracting cells to the quiescent state (Fig. 2C). Once a certain threshold of mitogens is reached, Rb is hyper-phosphorylated and E2F rapidly activated, facilitating progression through the RP. Note that this transition is irreversible in the absence of DNA damage such that cells become independent of growth factor stimulation once in the proliferating state, a hallmark of the RP (28).

Extending on the previously known bistability of the RP, our model suggests that p21 can modulate the Rb-E2F switch such that cells can enter quiescence even under high growth factor stimulation if they contain increased p21 levels (Fig. 2D). This mechanism primarily depends on p21’s ability to inhibit Cdk2 activity and thereby prevent activation of the positive
feedback loops that are required to pass the RP (SI Appendix, Fig. S2A). Due to bistability, cells on either side of the p21 threshold exhibit very different fates. Whilst cells with sub-threshold p21 levels continue to cycle, their supra-threshold counterparts enter quiescence (Fig. 2E). This dividing behaviour is even more apparent when considering Cdk2 activity (Fig. 2F). Both populations show an initial decrease in Cdk2 activity at the end of mitosis as CycA is degraded by the APC/C^{Cdc20}. However, Cdk2 activity rapidly recovers in cells which contain low p21 levels, while high p21 levels prevent this recovery, in excellent agreement with live-cell imaging (14, 16). The stark contrast between both populations, and the fact the intermediate states are rarely observed, provide further evidence that bistability is indeed at the core of the proliferation-quiescence decision.

In summary, our model suggests that p21 interferes with a bistable switch that controls RP passage to convert a graded p21 signal, which can be inherited from the mother cell (SI Appendix, Fig. S2B, C), into an all-or-nothing cell fate decision. It also shows that mitogen stimulation is necessary, but not sufficient, for proliferation.

**p21 degradation controls whether the proliferation decision can be reversed**

In addition to mitogen stimulation and past DNA damage that control whether a cell enters the cell cycle in the first place, it has been shown that the decision to proliferate can be reversed by exposure to stress and DNA damage in G1-phase (18). One mechanism that may mediate this response is an increased expression of p21 which inhibits cyclin:Cdk2 activity and DNA synthesis (Fig. 3A). Indeed, our simulations show that DNA damage incurred during G1-phase can reverse Cdk2 activation, pushing cells back into a pre-RP state with low Cdk2 activity and unphosphorylated Rb (SI Appendix, Fig. S3A). However, whether the decision to proliferate is reversible depends on both the strength of DNA damage and its timing (Fig. 3B). Specifically, our model predicts that cells in early G1-phase are particularly susceptible to arrest, with low levels of DNA damage being sufficient to reset the cell cycle. As cells move towards S-phase they progressively lose this sensitivity such that more, or more sustained, DNA damage is required to revert to quiescence. Intriguingly, our simulations suggest that S-phase cells completely lose the ability to enter p21-induced quiescence in response to the low levels of intrinsic DNA damage we consider here.

To test these model predictions, we introduced DNA damage into asynchronously cycling hTert-RPE1 cells with endogenously labelled mRuby-PCNA and p21-GFP, and followed their cell cycle fates and p21-GFP levels by time-lapse imaging. We imaged cells for 5 h to determine when they exited mitosis, before treating them with the DNA damage-inducing drug Camptothecin (CPT). We then tracked cells for a further 20 h to see whether they would enter S-phase or arrest. Control cells that were treated with DMSO almost exclusively entered S-phase, showing no bias for the length of time spent in G1 before treatment (Fig. 3C and SI Appendix, Fig. S3B). In sharp contrast, CPT-treated cells became increasingly less likely to enter quiescence the more time they had spent in G1. This is consistent with our model prediction that cells become less sensitive to DNA damage as they progress towards S-phase. Note that while we see an all-or-none response on the single-cell level, i.e., a cell either enters quiescence or fully commits to the cell cycle, experimental results compiled from multiple cells show a fractional response due to cell-to-cell variability.

Our model specifically predicts that cells lose the ability to enter quiescence because of an increasing rate of p21 degradation. Two ubiquitin ligases can mediate p21 degradation: SCF^{Skp2} and CRL4^{Cdt2} (Fig. 3A, (22)). We recently showed that it is Skp2 that controls the turnover of p21 in G1 (16). Simulating our model in the absence of Skp2 suggests that this perturbation would cause cells to remain highly sensitive to DNA damage until late G1 (Fig. 3D). To test this prediction, we depleted Skp2 using siRNA and repeated the above
Fig. 3 | p21 degradation controls the ability to enter DNA damage-induced quiescence.
(A) Subnetwork of DNA damage-induced control of early cell cycle progression including Skp2- and Cdt2-dependent p21 degradation pathways. (B) Effect of DNA damage of variable strength and timing on the decision to proliferate (bottom) in deterministic simulations of the full model in Fig. 1A. Damage was present from the indicated time until the end of the simulation. Time course of an undamaged cell is shown for comparison (top). (C, E, G) Experimental data showing the percentage of control-siRNA (C; n = 55, 77), Skp2-depleted (E; n = 71, 107) and Cdt2-depleted (G; n = 32, 52) cells that enter S-phase or arrest after treatment with DMSO or CPT at the indicated time post-mitosis. (D, F) Same as in the lower panel of B for Skp2-depleted (D) and Cdt2-depleted (F) cells.

consistent with our model, only those Skp2-depleted cells that were most advanced in G1 before CPT addition avoided quiescence (Fig. 3E). Furthermore, we find that while increasingly higher CPT doses are required to trigger quiescence in control cells as they advance in G1 (SI Appendix, Fig. S3C), Skp2 depletion leads to an abrupt, less drug dose-dependent transition (SI Appendix, Fig. S3D), as suggested by our model.
A further model prediction is that cells become insensitive to p21-induced quiescence during S-phase because of strong Cdt2-mediated p21 degradation. More precisely, our simulations suggest that Cdt2 depletion would weaken the G1/S bistable switch (SI Appendix, Fig. S1D). This would allow p21 to re-accumulate during S-phase (SI Appendix, Fig. S3E), such that now even S-phase cells become susceptible to DNA damage-induced quiescence (Fig. 3F). Yet, susceptibility of Cdt2-depleted cells to quiescence in early and mid G1 would be unaffected. To test this, we depleted Cdt2 in hTert-RPE1 cells. Indeed, we observed that when these cells were treated with CPT in early and mid G1 they were still able to enter S-phase (Fig. 3G). However, these cells only spend a short amount of time in S-phase and, more importantly, they re-accumulate p21 during DNA replication (SI Appendix, Fig. S3F, G), indicative of a reversion to quiescence.

Taken together, these data confirm our model predictions that Skp2-dependent p21 degradation leads to a progressive loss of sensitivity to DNA damage-induce quiescence in G1, which culminates in the Cdt2-mediated tolerance of DNA damage during S-phase.

**Loss of p21 and Rb causes uncontrolled proliferation**

The results above indicate that the Rb-p21 control network integrates signals from mitogen stimulation and DNA damage to mediate the proliferation-quiescence decision. Considering that many cancer cells have lost the ability to properly respond to these stimuli (29, 30), we used our model to explore which changes in the molecular network would compromise decision-making. With intact control, mitogen stimulation is necessary for proliferation such that cells below a certain threshold remain quiescent (Fig. 4A). Once this threshold is exceeded, the ratio between growth factor signalling and DNA damage, i.e., between the stimulation of Cdk2 activity by E2F-mediated cyclin synthesis and its inhibition by p21, controls proliferation. The molecular network integrates both stimuli and high amounts of damage or reduced mitogen levels will promote quiescence. Due to the presence of a bistable switch these continuous inputs are converted into a discontinuous, all-or-nothing cell fate decision, resulting in a sharp boundary between fates.

**Fig. 4** | **Protein loss leads to deregulated proliferation.** (A-D) Simulated effect of the presence of growth factors and DNA damage on the proliferation (blue)-quiescence (red) decision in unperturbed cells (A), and cells depleted for Rb (B), p21 (C) and Cdh1 (D). The indicated amount of damage and growth factors was present from the start of the simulation.
According to our model, a missing or non-functional Rb protein removes the necessity for mitogen stimulation with cells proliferating in the absence of growth factors (Fig. 4B), as has been found experimentally (31, 32). Yet, the system retains its ability to respond to DNA damage, albeit at much reduced sensitivity, i.e., the transition to a quiescent regime occurs at higher DNA damage levels. The latter is due to uninhibited E2F promoting substantial cyclin synthesis and Cdk2 activity, which requires a higher level of p21 induction to overcome. Loss of p21 protein has the opposite effect, abolishing DNA damage-induced quiescence, with the control by mitogens remaining in place (Fig. 4C). Finally, removal of Cdh1, another negative regulator of Cdk2 activity, makes cells less prone to enter quiescence in general. This is due to the increased accumulation of CycA:Cdk2 in G1-phase that provides another source of kinase activity that can promote RP passage (Fig. 4D). Yet, the Cdh1-depleted system still responds to both mitogen stimulation and DNA damage.

We decided to test the predictive value of our model by measuring the response of p21-proficient versus p21-knockout (p21KO, (16)) cells to a range of concentrations of growth factors and DNA damage. Consistent with our model prediction, we find that p21KO cells proliferate much more readily (SI Appendix, Fig. S4). Quiescence is only triggered at very high drug doses, most likely by DNA damage-sensing signalling pathways that can cause cell cycle arrest under high levels of exogenous DNA damage, which we did not consider in the model presented here of the cellular response to relatively lower levels of endogenous DNA damage.

In summary, our model suggests that the molecular network that controls the proliferation-quiescence decision integrates growth factor signalling and DNA damage into an all-or-nothing cell fate decision. Loss of Rb and p21 compromises the ability to enter quiescence in response to loss of mitogens and DNA damage, respectively, while Cdh1-depleted cells are more likely to proliferate in general.

**Discussion**

In this study, we developed a unifying model for the proliferation-quiescence decision in mammalian cells based on live single-cell imaging experiments. We show that two bistable switches underpin this decision: the RP, which mediates quiescence in response to serum starvation and DNA damage, and the G1/S transition allowing cells to tolerate intrinsic DNA damage during S-phase. The interplay between these switches establishes a window of reversibility, where cells otherwise committed to proliferation can revert to quiescence by expressing the Cdk inhibitor protein p21.

The RP has long been recognised as a prototype example of a bistable commitment point in cell cycle progression, converting a graded mitogen signal into an all-or-none decision to proliferate (27). Our model suggests that p21 expression interferes with this switch by interrupting a positive Cdk2-Rb-E2F feedback loop. This establishes a p21 threshold for RP passage. Above this threshold, p21 prevents E2F activation by blocking Rb hyper-phosphorylation, while at sub-threshold levels, Cdk2 can overcome p21 inhibition and phosphorylate Rb. Indeed, cells that exit mitosis with low p21 levels have been shown to maintain residual Cdk2 activity and Rb hyper-phosphorylation, which facilitates their re-entry into the cell cycle (14, 16). By contrast, intrinsic DNA damage incurred during S- or G2-phases can increase p21 to supra-threshold levels, promoting quiescence in the subsequent cell cycle. We thus suggest that p21 connects the DNA damage response to the RP switch, where a continuous p21 signal is converted into an all-or-none cell cycle response (Fig. 5A). Spontaneous DNA damage during unperturbed growth can thus lead to a quiescent subpopulation of cells characterised by low Cdk2 activity and high p21 levels. However, this
The RP establishes a bistable switch with two distinct cell fates: quiescence and proliferation. (A) p21 carried over from the previous cycle can drive cells into quiescence by tipping the balance toward low Cdk2 activity, counteracting growth factor-mediated passage through the RP. (B) If initial p21 levels are low, growth factors can induce the re-accumulation of cyclin:Cdk2. Yet, DNA damage incurred during G1-phase may reverse the decision to proliferate by promoting p21 accumulation and Cdk2 inhibition. (C) In S-phase, cells are insensitive to p21-induced quiescence as the Cdk inhibitor can no longer accumulate due to strong degradation. quiescent sub-population is absent in cell lines that have lost Rb or p53 function (23), as is the case in many cancers (29, 30).

When first proposed, the RP was described as a transition that renders cell cycle progression independent of growth-factor stimulation (33, 34). In our model, the RP is indeed an irreversible switch that can maintain the proliferative state in the absence of mitogens. Yet, for the transition to occur in the first place a certain mitogen threshold has to be exceeded such that Rb can become hyper-phosphorylated and E2F activated. We show that this threshold is controlled by p21 levels. However, growth factors remain a necessary condition for cell cycle entry even in the complete absence of p21. Consistent with this, p21-knockout cells enter quiescence under serum starvation conditions (15, 16). Our model also suggests that once a critical serum level is reached, the decision to proliferate depends on the ratio of mitogen stimulation and DNA damage, in agreement with experimental results showing that competition between p21 and mitogen-regulated cyclin D1:Cdk4 controls cell cycle entry (35). The RP, hence, serves as a hub that integrates growth signals, which promote cyclin:Cdk2 activity, and DNA damage, which counteracts Cdk2 through p21 expression. In support of this, increasing serum concentrations drive more cells into the proliferating state, while DNA damage and high p21 levels cause quiescence (14–16).

Previous studies have argued in favour of a singular decision point or window for cell cycle entry either located at the end of mitosis, in mid G1 or at the G1/S transition (14, 18, 33). Our simulations suggest that the interaction between the RP switch and the G1/S transition creates a decision window that extends at least throughout the entirety of G1 phase, where it integrates signals from mitogen-related pathways and DNA damage. During this window, cells that are committed to proliferation based on growth factor stimulation may revert to quiescence through DNA damage-induced p21 expression (Fig. 5B). Once cells do enter S-phase, strong p21 degradation by Cdt2 prevents p21 re-expression, insulating S-phase from mild, intrinsic DNA damage (Fig. 5C). This provides a mechanistic explanation for recent data showing that, in contrast to G1 and G2, S-phase is comparatively insensitive to DNA damage (36). We speculate that this insensitivity is important to prevent cells from entering growth arrest with partly replicated DNA, from which they could later re-emerge starting a second round of replication.

A recent study proposed that APC/C\textsuperscript{Cdh1} inactivation constitutes a point of no return for cell cycle entry, preventing cells from entering quiescence in response to exogenous stress (18). However, Cdh1 inactivation temporally overlaps with the G1/S transition and thus with the onset of Cdt2-mediated p21 degradation. Moreover, by facilitating CycA:Cdk2 accumulation,
the inactivation of Cdh1 not only promotes the G1/S transition but also strengthens p21 degradation via the Skp2-dependent pathway. Hence, Cdh1 regulation and p21 control are intimately linked. Nevertheless, our data suggest that it is p21 degradation that ultimately mediates the tolerance of intrinsic DNA damage. Two observations support this notion. First, compromising p21 degradation by Cdt2 depletion allows p21 re-accumulation during S-phase and premature S-phase exit despite Cdh1 control remaining in place. Second, Emi1-depleted cells that are treated with a Cdk inhibitor around the time of Cdh1 inactivation, i.e., at S-phase entry, show an immediate re-activation of Cdh1 (18). Yet, when the same experiment is conducted in the absence of Cdk inhibition but presence of a DNA-damaging drug, Cdh1 re-activation occurs only after a delay of several hours (18). We propose that, in this experiment, Cdh1 re-activation is caused by DNA damage-induced p21 inhibiting Cdk activity, and that the observed delay results from S-phase preventing p21 accumulation (SI Appendix, Fig. S3H, I). Thus, p21 degradation protects S-phase progression even when Cdh1 control is compromised.

Taken together, we show that bistable transitions serve two different purposes during cell cycle entry. The RP switch integrates growth and damage signals, translating them into an all-or-none decision to proliferate, while the G1/S transition temporarily insulates cell cycle progression from the p21-mediated DNA damage response during S-phase. Hence, these ubiquitous control motifs can provide both sensitivity and robustness to external stimuli.
Methods

Live cell imaging

Both alleles of the endogenous CDKN1A locus were labelled at the C-terminus with GFP in hTert-RPE1 cells, generating a cell line where all p21 protein was labelled with GFP (16). In the same cells, a single allele of PCNA was labelled at the N-terminus with mRuby, in order to determine the timing of the G1/S transition, S-phase exit and mitosis (37). To follow p21-GFP dynamics across the cell cycle, asynchronously cycling cultures growing in phenol-red free DMEM plus 10% FBS, were imaged every 10 mins on the Opera HCS microscope, using a 20x objective (NA 0.45) for up to 72 hrs. Automated image analysis was used to extract p21-GFP levels over time, relative to cell cycle transitions (38).

In order to test the sensitivity of G1 cells to DNA damage, cells were reverse-transfected with either control, Skp2- or Cdt2-targeting siRNAs (Dharmacon ONTargetPlus siRNA pools) 24 hrs before time-lapse imaging, using Lipofectamine RNAiMax (Invitrogen), according to the manufacturer’s instructions. Efficient depletion of Skp2 and Cdt2 protein levels by these siRNAs has been previously evaluated in this cell line (see (16)). Cells were filmed for 5 hours prior to the addition of Camptothecin (CPT) or DMSO and filming was continued for 20 hrs post-drug addition. mRuby-PCNA expression in these cells was used to determine the time of mitotic exit and the time of S-phase entry. A cell was classified as quiescent if it spend more than 600 min in G1, which we previously showed is consistent with a lack of both Rb hyper-phosphorylation and Cdk2 activity (16).

Analysis of proliferation-quiescence decisions in response to p21 knockout

hTert-RPE1 mRuby-PCNA p21-wild-type or hTert-RPE1 mRuby-PCNA p21KO cells (16) were plated in a 384-well CellCarrier plate at a density of 2000 cells/well in DMEM with 10% FBS. Once cells had attached, cells were washed three times in PBS and replaced with DMEM containing no FBS. Cells were serum-starved in this way for 72 hrs. We then added a range of concentrations of FBS and CPT in DMEM to cells and incubated for a further 24 hrs before fixing in 4% paraformaldehyde. Cells were permeabilised in PBS/0.2% TritonX-100, blocked in 2% BSA/PBS and immunostained with phospho-807/811 Rb antibody (CST 9308), in order to differentiate between quiescent and proliferating cells. Phospho-Rb antibody was detected by goat anti-rabbit Alexa 647 secondary antibody (ThermoFisher) and Hoescht was used to label nuclei. Plates were imaged on an Opera HCS (PerkinElmer) using a 20x objective, NA 0.45. Automated image analysis was performed using Columbus to determine the percentage of quiescent cells per condition. Briefly, nuclei were segmented using the Hoescht stain and mitotic, dead and poorly segmented cells were excluded, using intensity and morphology thresholds. The phospho-Rb intensity was quantified for each nucleus, and by calculating a threshold for proliferating cells, we could extract the percentage of quiescent cells per condition.

Mathematical modelling

Our model of cell cycle progression (see Fig. 1A), comprises a set of ordinary differential equations (see SI Appendix for details), which provide the basis for deterministic and stochastic simulations shown in the main text. A deterministic version of the model was prepared using the Systems Biology Toolbox 2 (39) for MatLab (version 9.1.0 R2016b) and simulated with the CVODE routine (40). Bifurcation diagrams were calculated using the freely available software XPP-Auto (41). We simulated a stochastic version of the model using custom-made MatLab code of the stochastic simulation algorithm, also known as Gillespie’s
algorithm (reviewed in (42)), according to the sorting direct method (43). The model is provided as Dataset S1 and different versions of it are available at www.cellcycle.org.uk/publication. The model has also been deposited in BioModels (44) and assigned the identifier MODEL1703030000.

Author contributions


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**A** Subnetwork of p21 regulation comprising three mutual inhibition motifs that involve Skp2- and Cdt2-dependent degradation pathways. **B** Stable (solid) and unstable (dashed) steady states of p21 during S-phase entry (left) and exit (right), calculated from the model using the subnetwork in A. Each line represents a different level of constant DNA damage. The G1/S transition is indicated for intermediate damage levels. Note the difference in y-axis scales. **C** Dynamics of cell cycle regulators (top), Skp2-dependent p21 degradation (middle) and Cdt2-dependent p21 degradation (bottom) in deterministic simulations. p21 degradation via Skp2 is linked to CycE- and CycA-associated Cdk2 activity, while Cdt2-dependent degradation depends on active replication complexes (aRC). Note the differences in y-axis scales between middle and bottom panels. **D** G1/S transition for low levels of DNA damage in a control simulation (unperturbed) and in the absence of Skp2 (-Skp2) and Cdt2 (-Cdt2). Note that the -Cdt2 curve overlaps with the control simulation for low cyclin:Cdk2 levels.

**Fig. S1.** A bistable switch controls p21 levels (related to Fig. 1). (A) Subnetwork of p21 regulation comprising three mutual inhibition motifs that involve Skp2- and Cdt2-dependent degradation pathways. (B) Stable (solid) and unstable (dashed) steady states of p21 during S-phase entry (left) and exit (right), calculated from the model using the subnetwork in A. Each line represents a different level of constant DNA damage. The G1/S transition is indicated for intermediate damage levels. Note the difference in y-axis scales. (C) Dynamics of cell cycle regulators (top), Skp2-dependent p21 degradation (middle) and Cdt2-dependent p21 degradation (bottom) in deterministic simulations. p21 degradation via Skp2 is linked to CycE- and CycA-associated Cdk2 activity, while Cdt2-dependent degradation depends on active replication complexes (aRC). Note the differences in y-axis scales between middle and bottom panels. (D) G1/S transition for low levels of DNA damage in a control simulation (unperturbed) and in the absence of Skp2 (-Skp2) and Cdt2 (-Cdt2). Note that the -Cdt2 curve overlaps with the control simulation for low cyclin:Cdk2 levels.
Fig. S2. p21 impairs RP passage (related to Fig. 2). (A) Same as in Fig. 2C except that total p21 was set to the indicated level. (B) Deterministic simulations of cells suffering increasing amounts of DNA damage (bottom to top) during S-phase. (C) Final p21 level of cells shown in (B) over the strength of DNA damage. Dashed line indicates the p21 level that would cause quiescence in daughter cells assuming that p21 protein is inherited but the damage repaired.
Fig. S3. p21 degradation controls reversibility of proliferation decision (related to Fig. 3). (A) Deterministic simulation of a cell that suffers DNA damage during G1-phase and enters quiescence (solid) using the full model in Fig. 1A. An undamaged cell is shown for comparison (dashed). (B) Percentage of unperturbed cells that enter S-phase or arrest after treatment with DMSO or CPT at the indicated time post-mitosis (n = 45, 59). Note that cells in this experiment were only imaged for 4 h prior to drug addition. (C, D) Percentage of control siRNA-treated (B; n = 292) and Skp2-depleted (C; n = 279) cells that enter S-phase (red to blue) in experiments after treatment with the indicated amount of CPT at the indicated time post-mitosis. Note that Fig. 3 and S3B show a different set of experiments, which were conducted at a single concentration of 1.25 μM CPT. (E) p21 levels from stochastic simulations of Cdt2-depleted cells in the absence of extrinsic DNA damage (n = 30). (F, G) Experimental p21 levels in control-siRNA (F; n = 10) and Cdt2-depleted (G; n = 10) hTert-RPE1 cells that entered S-phase after being treated with CPT at the times indicated by black dots. (H, I) Deterministic simulations of an unperturbed (H) and an Emi1-depleted (I) cell that suffer damage after S-phase entry. Note the re-activation of Cdh1 in G2-phase in response to DNA damage in the absence of Emi1.
Fig. S4. Loss of p21 leads to deregulated proliferation (related to Fig. 4). (A, B) Percentage of wild-type (A; n = 41224) and p21-knockout (B; n = 133004) cells that enter S-phase (red to blue) in experiments after treatment with the indicated amount of CPT and growth factors (FBS) that were present from the beginning until the end of the experiment. Cells were serum-starved for 72 h before the experiment.
Mathematical modelling

Restriction point

Entry into the cell cycle is control by the restriction point. In order to pass through it, cells hyper-phosphorylate the Rb protein \( R_b \), an inhibitor of the transcription factor E2F (19).

\[
\frac{d(Rb_t)}{dt} = -r_{Rb}^{Ph} \cdot Rb_t + k_{Rb}^{Dp} \cdot Rb_p,
\]

with \( r_{Rb}^{Ph} = k_{Rb,Cd}^{Ph} \cdot Cd + k_{Rb,Ce}^{Ph} \cdot Ce + k_{Rb,Ca}^{Ph} \cdot Ca, \)

and \( Rb_t = Rb_u + Rb_p = Rb + RbE2F + Rb_p, \)

where \( Rb_t, Rb_p, \) and \( Rb_u \) denote total, hyper- and hypo-phosphorylated Rb, respectively. The latter comprises both free Rb \( (Rb) \) and Rb:E2F complexes \( (RbE2F) \). In our model, phosphorylation occurs with rate \( r_{Rb}^{Ph} \) and is mediated by three cyclin:Cdk complexes: CycD:Cdk4/6 \( (Cd) \), CycE:Cdk2 \( (Ce) \) and CycA:Cdk2 \( (Ca) \) with their specific phosphorylation rates \( k_{Rb,Cd}^{Ph}, k_{Rb,Ce}^{Ph} \) and \( k_{Rb,Ca}^{Ph} \), respectively (4). The dephosphorylation rate is \( k_{Rb}^{Dp} \). In its hypo-phosphorylated state Rb binds free E2F \( (E2F) \) impairing its ability to promote transcription (19). Hence, we assumed that Rb (hyper-)phosphorylation prevents E2F binding and leads to the dissociation of Rb:E2F complexes.

\[
\frac{d(E2F_t)}{dt} = r_{E2F}^{Sy} - k_{E2F}^{De} \cdot E2F_t,
\]

\[
\frac{d(E2F)}{dt} = r_{E2F}^{Sy} - k_{E2F}^{De} \cdot E2F - k_{RbE2F}^{As} \cdot Rb \cdot E2F + \left( k_{RbE2F}^{Ds} + r_{Rb}^{Ph} \right) \cdot RbE2F,
\]

with \( r_{E2F}^{Sy} = k_{E2F}^{Sy} + k_{E2F,E2F}^{Sy} \frac{E2F}{j_{E2F} + E2F} \)

and \( E2F_t = E2F + RbE2F. \)

Here, \( E2F_t \) and \( E2F \) are total and free E2F, respectively, and \( r_{E2F}^{Sy} \) and \( k_{E2F}^{De} \) denote the E2F synthesis and degradation rate, respectively. Note that E2F promotes its own transcription (20). Hence, we assumed that synthesis occurs with a constitutive rate \( (k_{E2F}^{Sy}) \) and an E2F-dependent rate \( (k_{E2F,E2F}^{Sy}, j_{E2F}) \), where half-maximal synthesis is reach at \( j_{E2F}. \)

Association and dissociation of Rb and E2F are described by the rates \( k_{RbE2F}^{As} \) and \( k_{RbE2F}^{Ds} \), respectively.

Synthesis and degradation of cyclins and Emi1

Activation of E2F drives the synthesis of a battery of genes that are essential for early cell cycle progression (4, 20). Among these are Emi1 \( (E1) \), CycE \( (Ce) \) and CycA \( (Ca) \).

\[
\frac{d(E1_t)}{dt} = k_{E1}^{Sy} \cdot E2F - k_{E1,C1}^{De} \cdot E1C1 - k_{E1}^{De} \cdot E1,
\]

\[
\frac{d(Ce_t)}{dt} = k_{Ce}^{Sy} \cdot E2F - r_{Ce}^{De} \cdot Ce_t,
\]

\[
\frac{d(Ca_t)}{dt} = k_{Ca}^{Sy} \cdot E2F - r_{Ca}^{De} \cdot Ca_t,
\]

with \( r_{Ce}^{De} = k_{Ce}^{De} + k_{Ce,Ca}^{De} \cdot Ca, \)

\( r_{Ca}^{De} = k_{Ca}^{De} + k_{Ca,C1}^{De} \cdot C1, \)

and \( E1_t = E1 + E1C1, \)

where the index \( t \) denotes total protein levels including all complexes that contain the protein. \( C1 \) and \( E1C1 \) correspond to APC/C\(^Cdh1\) and its complex with Emi1, respectively (see
Inactivation of APC/C\textsuperscript{Cdh1}

APC/C\textsuperscript{Cdh1} is an ubiquitin E3 ligase that is active from late mitosis to late G1-phase and targets key cell cycle regulators for degradation (45, 49). Its inhibition by Emi1 and by cyclin:Cdk2-mediated phosphorylation is crucial for G1/S progression (23, 45). Therefore, we assumed that APC/C\textsuperscript{Cdh1} can exists in a free, active form (C1) as well as two inactive forms: as part of an Emi1:APC/C\textsuperscript{Cdh1} complex (E1C1) and as a free but Cdk2-phosphorylated form (C1\textsubscript{p}).

\[
\frac{d(C1)}{dt} = -r_{C1}^{Ph} \cdot C1 + k_{C1}^{Dp} \cdot C1\textsubscript{p} - k_{E1C1}^{As} \cdot E1 \cdot C1 + (k_{E1C1}^{De} + k_{E1C1}^{De}) \cdot E1C1, \\
\frac{d(E1C1)}{dt} = -r_{C1}^{Ph} \cdot E1C1 + k_{E1C1}^{As} \cdot E1 \cdot C1 - (k_{E1C1}^{De} + k_{E1C1}^{De}) \cdot E1C1, \\
\text{with } r_{C1}^{Ph} = k_{C1}^{Ph} + k_{C1,Ce}^{Ph} \cdot Ce + k_{C1,Ca}^{Ph} \cdot Ca,
\]

and C1\textsubscript{t} = C1 + C1\textsubscript{p} + E1C1.

Here, phosphorylation occurs with rate \(r_{C1}^{Ph}\) comprising the constitutive rate \(k_{C1}^{Ph}\) as well as the cyclin:Cdk2-dependent rates \(k_{C1,Ce}^{Ph}\) and \(k_{C1,Ca}^{Ph}\). We assumed that Emi1:APC/C\textsuperscript{Cdh1} complexes dissociate upon phosphorylation. Dephosphorylation is described by \(k_{C1}^{Dp}\). The rates of association and dissociation of APC/C\textsuperscript{Cdh1} and Emi1 are \(k_{E1C1}^{As}\) and \(k_{E1C1}^{De}\), respectively. Note that Emi1 is assumed to act as a pseudo-substrate for the APC/C (50). Hence, we chose \(k_{E1C1}^{De} > k_{E1}^{De}\) (see Eq. (4)).

\textbf{p21 dynamics and Cdk2 inhibition}

Activation of Cdk2 drives G1/S progression but its kinase activity is opposed by the stoichiometric inhibitor p21 (51). Hence, our model accounts for free p21 (P21) and its ability to bind active CyclE:Cdk2 (Ce) and CyclA:Cdk2 (Ca) complexes. The resulting complexes of CyclE:Cdk2:p21 (CeP21) and CyclA:Cdk2:p21 (CaP21) are considered inactive.

\[
\frac{d(P21)}{dt} = k_{P21}^{Sy} k_{P21,P53}^{Sy} \cdot P53 - r_{P21}^{De} \cdot P21, \\
\frac{d(CeP21)}{dt} = k_{CyP21}^{As} \cdot P21 \cdot Ce - (k_{CyP21}^{Ds} + r_{Cy}^{De} + r_{Cy}^{De}) \cdot CeP21, \\
\frac{d(CaP21)}{dt} = k_{CyP21}^{As} \cdot P21 \cdot Ca - (k_{CyP21}^{Ds} + r_{Cy}^{De} + r_{Cy}^{De}) \cdot CaP21,
\]

with \(r_{P21}^{De} = k_{P21}^{De} + k_{P21,Cy}^{De} \cdot Skp2 \cdot (Ce + Ca) + k_{P21,RCa}^{De} \cdot Cdt2 \cdot R_c\)

\[
P21_t = P21 + CeP21 + CaP21 + Pca1 + R_c, \\
Ce_t = Ce + CeP21, \\
Ca_t = Ca + CaP21.
\]
Here, synthesis of p21 occurs with the constitutive rate $k_{P_{21}}^{Sy}$ and the p53-dependent rate $k_{P_{21},P53}^{Sy}$, where P53 denotes the transcription factor p53 (5). The degradation rate of p21 is denoted $r_{P_{21}}^{De}$ and comprises a constitutive rate ($k_{P_{21}}^{De}$) as well as a cyclin:Cdk2-dependent rate ($k_{P_{21},Cy}^{De}$) that is linked to the ubiquitin E3 ligase SCF$^{Skp2}$ (Skp2, (22, 52)). In addition, CRL4$^{Cdt2}$ (Cdt2), which is recruited to chromatin-bound PCNA within active replication complexes ($R_{Ca}$, see below), can mediate p21 degradation with rate $k_{P_{21},R_{Ca}}^{De}$ (22, 53). In our model, free p21 can bind to cyclin:Cdk2 complexes with rate $k_{CyP_{21}}^{As}$ and dissociate from these complexes with rate $k_{CyP_{21}}^{Ds}$. Furthermore, the total amount of p21 ($P_{21}$) comprises free p21 ($P_{21}$) as well as p21 in complexes with cyclin:Cdk2 ($CeP_{21}$ and $CaP_{21}$), PCNA ($Pca_{i}$), and replication complexes ($Rc_{i}$, see below).

**Activation of DNA replication**

During S-phase, DNA is synthesised by specialised replication complexes that assemble on licensed replication origins once there is sufficient Cdk2 activity (54). Therefore, we assumed that Cdk2 activates pre-replication complexes ($Rc$) such that they are primed ($Rc_{p}$) for the loading of DNA processivity factors.

$$\frac{d(Rc)}{dt} = -r_{Rc}^{Ph} \cdot Rc + k_{Rc}^{dp} \cdot Rc_{p} - r_{Rc}^{Ds} \cdot Rc,$$

with $r_{Rc}^{Ph} = k_{Rc}^{Ph} \cdot \frac{(Ce+Ca)^n}{(J_{Cy})^n + (Ce+Ca)^n}$

where priming occurs with rate $k_{Rc}^{Ph}$ in an ultrasensitive fashion with hill coefficient $n$ at a cyclin:Cdk2 threshold of $J_{Cy}$.

Primed replication complexes can revert back to an unprimed state with a small rate $k_{Rc}^{dp}$ and replication complexes are disassembled upon completion of S-phase with rate $r_{Rc}^{Ds}$ (see below). Once primed, replication complexes can bind PCNA, which forms a trimeric complex around the DNA that serves as a sliding platform for other processivity factors (7). We assumed that either free PCNA ($Pca_{a}$) or PCNA:p21 complexes ($Pca_{i}$) are bound leading to the formation of active ($Rc_{a}$) or inactive ($Rc_{i}$) replication complexes, respectively.

$$\frac{d(Rc_{p})}{dt} = r_{Rc}^{Ph} \cdot Rc - k_{Rc}^{dp} \cdot Rc_{p} - k_{Rcpc}^{As} \cdot (Pca_{a} + Pca_{i}) \cdot Rc_{p} + k_{Rcpc}^{Ds} \cdot (Rc_{a} + Rc_{i})$$

$$\frac{d(Rc_{a})}{dt} = -k_{PcP_{21}}^{As} \cdot P21 \cdot Rc_{a} + (k_{PcP_{21}}^{Ds} + r_{P21}^{As}) \cdot Rc_{i} + k_{Rcpc}^{As} \cdot Pca_{a} \cdot Rc_{p} - k_{Rcpc}^{Ds} \cdot Rc_{a}$$

$$\frac{d(Rc_{i})}{dt} = k_{PcP_{21}}^{As} \cdot P21 \cdot Rc_{a} - (k_{PcP_{21}}^{Ds} + r_{P21}^{As}) \cdot Rc_{i} + k_{Rcpc}^{As} \cdot Pca_{i} \cdot Rc_{p} - k_{Rcpc}^{Ds} \cdot Rc_{i}$$

The rates of PCNA loading and unloading are $k_{Rcpc}^{As}$ and $k_{Rcpc}^{Ds}$, respectively. Also note that p21 is the strongest known binding partner of PCNA (9) and it was proposed to compete with other PCNA-binding proteins (7, 8). Thus, we assumed that binding of p21 to active replication complexes with rate $k_{PcP_{21}}^{As}$ (or binding of PCNA:p21 to primed replication complexes) creates inactive replication complexes (10, 55). The dissociation of p21 from PCNA occurs with rate $k_{PcP_{21}}^{Ds}$. In our model, active replication complexes synthesise DNA ($DNA$) and once DNA replication is finished, replication complexes disassemble with rate $r_{Rc}^{Ds}$.

$$\frac{d(DNA)}{dt} = k_{DNA}^{Sy} \cdot Rc_{a},$$

and $r_{Rc}^{Ds} = H(DNA - 1)$.
where $k_{\text{DNA}}^{\text{SY}}$ is the DNA synthesis rate and $H$ the Heaviside function.

**PCNA dynamics**

We assumed that free PCNA in the nucleus ($PCn_a$) is replenished from a cytoplasmic pool and that p21 can bind to PCNA creating PCNA:p21 complexes ($PCn_ap21$, (9)).

$$\frac{d(PCn_{ap21})}{dt} = k_{PCn}^{\text{p21}} \cdot P21 \cdot PCn_a + (k_{PCn}^{\text{Ds}} + r_{PCn}^{\text{p21}}) \cdot PCn_{ap21}$$

$$- (k_{PCn}^{\text{Ex}} + k_{RCPC}^{\text{p21}} \cdot Rc_p) \cdot PCn_{ap21} + (k_{RCPC}^{\text{p21}} + r_{RC}^{\text{p21}}) \cdot Rc_a.$$  

$$\frac{d(PCn_a)}{dt} = k_{PCn}^{\text{Ap21}} \cdot P21 \cdot PCn_a - (k_{PCn}^{\text{Ds}} + r_{PCn}^{\text{p21}}) \cdot PCn_{ap21}$$

$$- (k_{PCn}^{\text{Ex}} + k_{RCPC}^{\text{p21}} \cdot Rc_p) \cdot PCn_a + (k_{RCPC}^{\text{p21}} + r_{RC}^{\text{p21}}) \cdot Rc_i.$$  

Here, $k_{PCn}^{\text{p21}}$ and $k_{PCn}^{\text{Ex}}$ denote the nuclear import and export rate PCNA, respectively. The association and dissociation of p21 and PCNA occur with rate $k_{PCn}^{\text{Ap21}}$ and $k_{RCPC}^{\text{p21}}$, respectively. The binding and unbinding rates of PCNA and replication complexes are $k_{RCPC}^{\text{p21}}$ and $k_{RCPC}^{\text{p21}}$, respectively.

**DNA damage and repair**

In our model, DNA damage ($Dam$) can either occur independent of the cell cycle stage or specifically during DNA replication. It induces the expression of p53 ($P53$), which promotes damage repair processes (56).

$$\frac{d(Dam)}{dt} = k_{\text{Dam}}^{\text{Ge}} + k_{\text{Dam,RCa}}^{\text{Ge}} \cdot Rc_a - r_{\text{Dam}}^{\text{Re}} \cdot Dam.$$  

$$\frac{d(P53)}{dt} = k_{P53}^{\text{Sy}} - r_{P53}^{\text{De}} \cdot P53,$$

with $r_{\text{Dam}}^{\text{Re}} = k_{\text{Dam}}^{\text{Re}} + k_{\text{Dam,P53}}^{\text{Re}} \cdot \frac{p53}{j_{\text{Dam}} + p53}$

and $r_{P53}^{\text{De}} = \frac{k_{P53}^{\text{De}}}{j_{P53} + Dam}$.

where $k_{\text{Dam}}^{\text{Ge}}$ and $k_{\text{Dam,RCa}}^{\text{Ge}}$ correspond to the constitutive and DNA replication-dependent rates of damage induction, respectively. Damage is repaired with a constitutive rate ($k_{\text{Dam}}^{\text{Re}}$) and a p53-dependent rate ($k_{\text{Dam,P53}}^{\text{Re}}$) assuming that p53 expression triggers repair processes. The latter reaches its half-maximal rate at a damage level of $j_{\text{Dam}}$. Since DNA damage has been shown to stabilize p53 (57), we assumed that p53 synthesis occurs with a constitutive rate ($k_{P53}^{\text{Sy}}$), while degradation occurs with rate $k_{P53}^{\text{De}}$, which is inversely proportional to the sum of DNA damage and an inhibition constant $j_{P53}$.

**Cdh1 activity reporter**

In order to compare our simulation results with measurements of Cdh1 activity obtained by following the degradation of a model APC/C<sup>Cdh1</sup> substrate (18), we simulated such an activity probe ($Pr$).

$$\frac{d(Pr)}{dt} = k_{Pr}^{\text{Sy}} - (k_{Pr}^{\text{De}} + k_{\text{Ca,C1}}^{\text{De}} \cdot C1) \cdot Pr,$$

where the constitutive synthesis and degradation rates of the probe are $k_{Pr}^{\text{Sy}}$ and $k_{Pr}^{\text{De}}$, respectively, and, without loss of generality, the Cdh1-dependent degradation rate $k_{\text{Ca,C1}}^{\text{De}}$ is the same than for CycA. Based on the analysis of experimental data (18), we calculated Cdh1 activity ($C1_{\text{act}}$) using the following equation.
\[ C_{1\text{act}} = \frac{k_{Pr}^{Sy} \Delta (Pr + Bg)}{\Delta t (Pr + Bg)}. \] (22)

Here, we added a small background signal \((Bg)\) to reflect that while small changes in probe concentration can be accurately calculated from numerical data they might be below the signal-to-noise ratio in experiments.

**Computation**

A deterministic version of the model was prepared using the Systems Biology Toolbox 2 (39) for MatLab (version 9.1.0 R2016b) and simulated with the CVODE routine (40). Bifurcation diagrams were calculated using the freely available software XPP-Aut (41). The model is provided as Dataset S1 and different versions of it are available at www.cellcycle.org.uk/publication. The model was also deposited in BioModels (44) and assigned the identifier MODEL1703030000.

We simulated a stochastic version of the model using custom-made MatLab code of the stochastic simulation algorithm, also known as Gillespie’s algorithm (reviewed in (42)), according to the sorting direct method (43). To this end, the rate expressions of the deterministic model were converted into propensity functions, which requires the transformation of the relative levels of cell cycle regulators into numbers of molecules. Here, we followed the system in a control volume containing on average 1000 molecules/AU of protein and 1 event/AU of DNA damage. Hence, most of the simulated cell-to-cell variability originates in the stochastic infliction of DNA damage, while transcriptional and translational noise play only a minor role. To account for the extrinsic noise observed in our experiments, i.e., the variability in p21 levels after cell division, we chose the initial p21 level for each cell (simulation run) from a log-normal distribution with \(\mu = -0.5\) and \(\sigma = 0.3\). This distribution is based on the distribution of initial p21 levels estimated from experiments \((\mu = 2.4, \sigma = 0.3)\), which was rescaled to account for the difference in p21 level between experiments and simulations. For the simulation of depletion experiments \(Skp2\) and/or \(Cdt2\) were reduced to 1%.

To classify cells as either proliferating or quiescent we used the fact that these two fates represent attractors in state space in our model (see Fig. 2C-F). In particular, quiescent cells are characterised by high p21 levels as well as low Cdk2 activity, Rb hypo-phosphorylation and low E2F activity, which prevents entry into S-phase. Hence, simulated cells that maintain these features were classified as quiescent, while cells that did enter S-phase were deemed proliferating.

**Model parameters**

Parameter values and non-zero initial conditions of the model are listed in Tables S1 and S2, respectively. A large part of the parameter set is based on our previously published study of p21 dynamics (16), where the model was parameterised to capture p21 dynamics in individual, unperturbed hTert-RPE1 cells as well as in perturbation experiments such as upon depletion of Skp2 and Cdt2. This includes parameters for p21 synthesis and degradation, the import, export, phosphorylation and binding of PCNA, and the dynamics of p53 and DNA damage. Parameter values for the newly added restriction point module are based on experimental observations and were adapted to reproduce cell cycle progression in RPE1 cells (see Fig. 1C-E). In particular, we assumed that in its hypo-phosphorylated state Rb binds tightly to E2F as it was shown to act as a stoichiometric inhibitor (19). Synthesis and degradation of E2F were chosen to yield a bistable response, which was found experimentally (27). Inhibition of \(APC/C^{Ch1}\) by Emi1 was modelled via a pseudo-substrate...
inhibitor mechanism (50), implying tight binding of both and a slow turnover of Emi1 by Cdh1. Synthesis and degradation of Emi1 and phosphorylation of APC/C<sup>Cdh1</sup> were adjusted to yield a rapid, switch-like inactivation of APC/C<sup>Cdh1</sup> at the G1/S transition based on measurements of APC/C activity (18). Parameters for CycE and CycA synthesis and degradation were chosen to qualitatively match the measurements of both proteins in live, single cells (23) and of overall Cdk2 activity (14, 16), while accounting for the length of cell cycle phases in RPE1 cells (see Fig. 1E). Local parameter sensitivities (see Table S1) were calculated according to

\[
S_\theta = \frac{\Delta M_\theta}{M \Delta \theta}
\]

with

\[
\frac{\Delta M}{M} = \frac{1}{N_V N_t} \sum_{V,t} \left( \frac{y_V(\theta, t) - y_V(\theta^*, t)}{\sigma_V} \right)^2
\]

where \( S_\theta \) is the local sensitivity of parameter \( \theta \), comprising the relative change in model output \( M \) with respect to the relative change in parameter \( \theta \). The relative change in model output was defined as the squared change in state variable \( V \) at time \( t \) given the original parameter set \( y_V(\theta, t) \) and the same output in response to a parameter change \( y_V(\theta^*, t) \), normalised by the maximum value of variable \( V \) \( (\sigma_V) \), which was summed over all time points and state variables and normalised to the number of simulated time points \( (N_t) \) and state variables \( (N_V) \), following (58).
# Supporting tables

## Table S1. Parameters of the mathematical model.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
<th>Value</th>
<th>Unit</th>
<th>Local Sensitivity (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$B_g$</td>
<td>background of Cdh1-activity probe</td>
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<td>–</td>
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<td>$C_{11}$</td>
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<td>$k_{E1C1}$</td>
<td>association of Emi1 and APC/C$^{Cdh1}$</td>
<td>10</td>
<td>1/(AU-min)</td>
<td>0.1</td>
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<tr>
<td>$k_{P_{21}}$</td>
<td>association of PCNA and p21</td>
<td>100</td>
<td>1/(AU-min)</td>
<td>0.001</td>
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<tr>
<td>$k_{R_b&amp;E2F}$</td>
<td>association of Rb and E2F</td>
<td>5</td>
<td>1/(AU-min)</td>
<td>0.34</td>
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<tr>
<td>$k_{Skp2}$</td>
<td>association of primed RCs and PCNA</td>
<td>0.01</td>
<td>1/(AU-min)</td>
<td>0.45</td>
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<tr>
<td>$k_{C_{y}}$</td>
<td>constitutive CycA degradation</td>
<td>0.01</td>
<td>1/min</td>
<td>0.15</td>
</tr>
<tr>
<td>$k_{C_{a}}$</td>
<td>constitutive CycE degradation</td>
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<td>1/min</td>
<td>0.89</td>
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<td>$k_{C_{a,C1}}$</td>
<td>APC/C$^{Cdh1}$-mediated CycA degradation</td>
<td>2</td>
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<tr>
<td>$k_{E_{1}}$</td>
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<td>1/min</td>
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<td>$k_{E_{1,C1}}$</td>
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<td>0.01</td>
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<td>1/min</td>
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<td>$k_{p21}$</td>
<td>constitutive p21 degradation</td>
<td>0.0025</td>
<td>1/min</td>
<td>0.05</td>
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<td>$k_{p21,Cy}$</td>
<td>cyclin:Cdk2-mediated p21 degradation</td>
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<td>$k_{p21,RCa}$</td>
<td>RC-mediated p21 degradation</td>
<td>1</td>
<td>1/(AU-min)</td>
<td>0.005</td>
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<td>$k_{P_{53}}$</td>
<td>DNA damage-dependent p53 degradation</td>
<td>0.05</td>
<td>AU/min</td>
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<td>$k_{p21}$</td>
<td>constitutive degradation of Cdh1-activity probe</td>
<td>0.0001</td>
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<td>1·10^-5</td>
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<td>$k_{C_{y}}$</td>
<td>dephosphorylation of APC/C$^{Cdh1}$</td>
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<td>0.05</td>
<td>1/min</td>
<td>0.001</td>
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<td>$k_{D_{a}}$</td>
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<td>1/min</td>
<td>1·10^-5</td>
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<td>$k_{P_{21}}$</td>
<td>dissociation of PCNA:p21 complexes</td>
<td>0.01</td>
<td>1/min</td>
<td>1·10^-4</td>
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<td>1/min</td>
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<td>$k_{R_b&amp;C}$</td>
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<td>0.001</td>
<td>1/min</td>
<td>0.01</td>
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<td>$k_{E}E_{1}$</td>
<td>PCNA export from the nucleus</td>
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<td>0.91</td>
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<td>$k_{D_{am}}$</td>
<td>replication-independent DNA damage</td>
<td>0.001</td>
<td>AU/min</td>
<td>0.01</td>
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<td>$k_{P_{c}}$</td>
<td>replication-dependent DNA damage</td>
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<td>1/min</td>
<td>0.01</td>
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<td>$k_{P_{c}}$</td>
<td>PCNA import into the nucleus</td>
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<td>$k_{C_{y}}$</td>
<td>constitutive APC/C$^{Cdh1}$ phosphorylation</td>
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<td>1/min</td>
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<td>Cyca:Cdk2-mediated APC/C$^{Cdh1}$ phosphorylation</td>
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<td>1/(AU-min)</td>
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<td>$k_{C_{y}}$</td>
<td>Cyca:Cdk2-mediated Rb phosphorylation</td>
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<td>0.13</td>
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<td>Variable</td>
<td>Description</td>
<td>Value</td>
<td>Unit</td>
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<td>( k_{RhCd} )</td>
<td>CycD:Cdk4/6-mediated Rb phosphorylation</td>
<td>0.2</td>
<td>1/(AU-min)</td>
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<tr>
<td>( k_{RbCe} )</td>
<td>CycE:Cdk2-mediated Rb phosphorylation</td>
<td>0.3</td>
<td>1/(AU-min)</td>
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<tr>
<td>( k_{Rc} )</td>
<td>Cyclin:Cdk2-mediated priming of RCs</td>
<td>0.1</td>
<td>1/min</td>
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<td>( k_{Re} )</td>
<td>p53-independent DNA damage repair</td>
<td>0.001</td>
<td>1/min</td>
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<tr>
<td>( k_{Re, Dam} )</td>
<td>p53-dependent DNA damage repair</td>
<td>0.005</td>
<td>1/min</td>
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<tr>
<td>( k_{Sy} )</td>
<td>constitutive CycA synthesis</td>
<td>0.02</td>
<td>1/min</td>
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<tr>
<td>( k_{Ca} )</td>
<td>constitutive CycE synthesis</td>
<td>0.01</td>
<td>1/min</td>
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<td>( k_{Sy} )</td>
<td>DNA synthesis by active RCs</td>
<td>0.0093</td>
<td>1/min</td>
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<tr>
<td>( k_{Sy} )</td>
<td>constitutive Emi1 synthesis</td>
<td>0.005</td>
<td>1/min</td>
<td></td>
</tr>
<tr>
<td>( k_{E2F} )</td>
<td>constitutive E2F synthesis</td>
<td>0.03</td>
<td>AU/min</td>
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<tr>
<td>( k_{E2F,E2F} )</td>
<td>E2F-dependent E2F synthesis</td>
<td>0.04</td>
<td>AU/min</td>
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</tr>
<tr>
<td>( k_{Sy} )</td>
<td>constitutive p21 synthesis</td>
<td>0.002</td>
<td>AU/min</td>
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<tr>
<td>( k_{P21} )</td>
<td>p53-dependent p21 synthesis</td>
<td>0.008</td>
<td>1/min</td>
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<tr>
<td>( k_{Ps} )</td>
<td>constitutive p53 synthesis</td>
<td>0.05</td>
<td>AU/min</td>
<td></td>
</tr>
<tr>
<td>( n )</td>
<td>hill coefficient for priming of RCs</td>
<td>6</td>
<td>-</td>
<td></td>
</tr>
</tbody>
</table>

*In simulations of depletion experiments*

**Table S2. Non-zero initial conditions of the mathematical model.**

<table>
<thead>
<tr>
<th>Variable</th>
<th>Description</th>
<th>Value</th>
<th>Unit</th>
</tr>
</thead>
<tbody>
<tr>
<td>( C_{at} )</td>
<td>CycA:Cdk2 level</td>
<td>1.2</td>
<td>AU</td>
</tr>
<tr>
<td>( C_{et} )</td>
<td>CycE:Cdk2 level</td>
<td>0.5</td>
<td>AU</td>
</tr>
<tr>
<td>( P21_{at} )</td>
<td>total p21 level</td>
<td>0.6</td>
<td>AU</td>
</tr>
<tr>
<td>( Pcn_{t} )</td>
<td>free PCNA in the nucleus</td>
<td>0.5</td>
<td>AU</td>
</tr>
<tr>
<td>( Rc )</td>
<td>pre-replication complexes</td>
<td>1</td>
<td>AU</td>
</tr>
</tbody>
</table>

*These values correspond to a newly-born cell.*