Fluctuations in p53 Signaling Allow Escape from Cell-Cycle Arrest

Graphical Abstract

Highlights

- Live single-cell profiling post-irradiation reveals heterogeneity in arrest maintenance
- Heterogeneity in arrest maintenance results from noisy p53 pulse amplitude
- The interplay between p21 and CDK2 translates p53 fluctuations into escape from arrest
- High p21 levels are sufficient to establish arrest but insufficient to maintain it

Authors

José Reyes, Jia-Yun Chen, Jacob Stewart-Ornstein, Kyle W. Karhohs, Caroline S. Mock, Galit Lahav

Correspondence

galit@hms.harvard.edu

In Brief

Reyes et al. show that individual human cells vary in their ability to maintain cell-cycle arrest in the course of 1 week after DNA damage. They show that fluctuations in the oscillatory dynamics of the tumor suppressor p53 can trigger a switch from an arrested to a proliferative state.
Fluctuations in p53 Signaling Allow Escape from Cell-Cycle Arrest

José Reyes,1,2 Jia-Yun Chen,1,3 Jacob Stewart-Ornstein,1 Kyle W. Karhohs,1,2,4 Caroline S. Mock,1 and Galit Lahav1,3,5,*

1Department of Systems Biology, Harvard Medical School, Boston, MA 02115, USA
2Systems Biology PhD Program, Harvard Medical School, Boston, MA 02115, USA
3Laboratory of Systems Pharmacology, Harvard Medical School, Boston, MA 02115, USA
4Present address: Broad Institute of MIT and Harvard, Cambridge, MA 02142, USA
5Lead Contact
*Correspondence: galit@hms.harvard.edu
https://doi.org/10.1016/j.molcel.2018.06.031

SUMMARY

Biological signals need to be robust and filter small fluctuations yet maintain sensitivity to signals across a wide range of magnitudes. Here, we studied how fluctuations in DNA damage signaling relate to maintenance of long-term cell-cycle arrest. Using live-cell imaging, we quantified division profiles of individual human cells in the course of 1 week after irradiation. We found a subset of cells that initially establish cell-cycle arrest and then sporadically escape and divide. Using fluorescent reporters and mathematical modeling, we determined that fluctuations in the oscillatory pattern of the tumor suppressor p53 trigger a sharp switch between p21 and CDK2, leading to escape from arrest. Transient perturbation of p53 stability mimicked the noise in individual cells and was sufficient to trigger escape from arrest. Our results show that the self-reinforcing circuitry that mediates cell-cycle transitions can translate small fluctuations in p53 signaling into large phenotypic changes.

INTRODUCTION

In response to DNA damage, proliferating cells can either repair the damage and resume growth or activate anti-proliferative programs such as cell death (apoptosis) or senescence, a state characterized by the long-term enforcement of cell-cycle arrest and the loss of recovery potential (Figure 1A). While pro-apoptosis therapy has been used for several decades as a tool for destroying the growth of cancerous cells, recent studies also highlighted the therapeutic potential of pro-senescence cancer therapy (Collado and Serrano, 2010; Nardella et al., 2011; Xue et al., 2007). However, as opposed to apoptosis, which is a terminal cell fate, senescing cells require continuous activation of the pathways responsible for maintaining the arrested state (Beausejour et al., 2003; Dirac and Bernards, 2003) (Figure 1A). It is unclear how senescing cells respond to fluctuations in these pathways over prolonged times.

The tumor-suppressor protein p53 is a master transcriptional regulator of the response of human cells to DNA damage (Lakin and Jackson, 1999). Upon cellular exposure to ionizing radiation, p53 stabilization leads to the transcriptional induction of hundreds of genes involved in DNA repair, cell-cycle arrest, apoptosis, and cellular senescence (Riley et al., 2008). In addition, p53 regulates the expression of proteins involved in controlling its levels. In particular, the direct p53 transcriptional target mouse double minute 2 (MDM2) E3 ubiquitin ligase tags p53 for proteosomal-dependent degradation (Haupt et al., 1997), forming a negative feedback loop. Dynamically, the interaction of p53 and MDM2 generates oscillatory dynamics of p53 activation characterized by a stereotyped frequency and noisy amplitude (Lahav et al., 2004). While pulsatile p53 dynamics have been quantified in multiple cell lines over 24 hr after DNA damage (Gev-a-Zatorsky et al., 2006; Stewart-Ornstein and Lahav, 2017), the long-term evolution of such dynamics has not been explored. In addition, it was shown that activation of p53 during G2 is sufficient to trigger entry into senescence (Krenning et al., 2014). However, it is not known the extent to which heterogeneity in p53 signaling over time affect the long-term maintenance of the senescence state in individual cells.

Here, we studied the way fluctuations in DNA damage signaling relate to cell fate heterogeneity in the long-term response of human cells to ionizing radiation. Using live-cell imaging, we identified a subpopulation of cells that initially established cell-cycle arrest but escaped such state in the presence of damage through sporadic cell-cycle reentry events spanning ~1 week after irradiation. Using fluorescent reporters for p53 and its downstream target, the CDK inhibitor p21, we showed that cell-to-cell variation in the level of these proteins contributes to heterogeneity in the ability of individual cells to maintain the arrested state over long timescales. We further showed that escape from cell-cycle arrest is characterized by a sharp switch between p21 levels and CDK2 activity. By analyzing natural variability in p53 signaling in the vicinity of this transition, we showed that transient fluctuations in p53 pulse amplitude are amplified into escape from arrest as a consequence of the double-negative feedback interaction between p21 and CDK2. Taken together, we identified two levels at which heterogeneity in DNA damage signaling shapes the long-term ability of human cells to maintain cell-cycle arrest in the presence of persistent DNA damage: (1) low historical average p53 amplitude defines...
Cycling cells and developed a semiautomated tracking method to quantify the number and timing of division events that an isogenic population of cells experience in the course of 1 week, during which they undergo senescence (Krenning et al., 2014) (Figures 1B and 1C; Video S1). Division profiles revealed large heterogeneity between single cells, ranging from cells that did not divide at all to cells that divided only once (single dividers, Figure 1D, red squares) to cells that showed multiple divisions (Figure 1D), the proportion of which changed with irradiation dose (Figure 1D). Single dividers exhibited bimodality in their mitosis timing; the first phase of divisions occurred within the first day after DNA damage, and the second phase started ~2 days post-irradiation and was broadly distributed, spanning an entire week (Figure 1E). Note that single dividers were present in every irradiation and was broadly distributed, spanning an entire week (Figure 1E). Note that single dividers were present in every irradiation and was broadly distributed, spanning an entire week (Figure 1E).

RESULTS

Long-Term Live-Cell Imaging Reveals Heterogeneity in Cell-Cycle Arrest Maintenance after DNA Damage

We used fluorescent live-cell imaging to study DNA-damage-induced senescence in individual human cells. We irradiated cells and developed a semiautomated tracking method to quantify the number and timing of division events that an isogenic population of cells experience in the course of 1 week, during which they undergo senescence (Krenning et al., 2014) (Figures 1B and 1C; Video S1). Division profiles revealed large heterogeneity between single cells, ranging from cells that did not divide at all to cells that divided only once (single dividers, Figure 1D, red squares) to cells that showed multiple divisions (Figure 1D), the proportion of which changed with irradiation dose (Figure 1D). Single dividers exhibited bimodality in their mitosis timing; the first phase of divisions occurred within the first day after DNA damage, and the second phase started ~2 days post-irradiation and was broadly distributed, spanning an entire week (Figure 1E). Note that single dividers were present in every dose of DNA damage, ranging from nonphysiological levels that lead to loss of recovery potential to those that are used in radiation therapy (Lomax et al., 2013). We identified single dividers even in nonirradiated controls, suggesting that heterogeneity in

Figure 1. DNA Damage Leads to Heterogeneous Division Profiles over Long Timescales

(A) DNA damage can lead to different cellular outcomes, including terminal cell fates. Cellular senescence requires active maintenance.

(B) Representative images of cells assayed for senescence associated β-galactosidase (SA-β-gal) activity 6 days post-irradiation.

(C) Frequency of SA-β-gal-positive cells 6 days post-irradiation as a function of damage dose.

(D) Division profiles obtained after tracking individual telomerase-immortalized primary cells and annotating mitoses in the course of 1 week after DNA damage. Panels aggregate single cells exposed to a particular irradiation dose. Each row represents the division profile of an individual cell over time. Colors change upon mitosis. Cells are grouped by their total number of mitoses and ordered by the timing of their first mitosis. Red boxes highlight the single divider populations.

(E) Distribution of mitosis timing in single dividers.

(F) Single cell quantification of mVenus-hGeminin(1–110) reporter for a multiple divider (top) and a late divider (bottom).

(G and H) Distributions of G1 (G) and S/G2 (H) duration in unirradiated cycling cells or irradiated late dividers (n = 77 cells per condition).
timing and number of divisions is present in cells that engage in a long-term arrested program due to DNA damage generated during regular cycling, even without an external source of stress. Using the mVenus-hGemini(1–110) fluorescent reporter for cell-cycle progression (Sakaue-Sawano et al., 2008) (Figure 1F), we found that heterogeneity in the second phase is mainly attributed to time spent in a G0/G1 state (Figures 1F–1H), suggesting that late-dividing cells initially arrested in G1 and then escaped from the arrest. Taken together, these results revealed that arrested cells hold the potential for reversion of cell-cycle arrest for prolonged timescales (~1 week).

**Escaper Cells Reenter the Cell Cycle in the Presence of DNA Damage**

We next sought to understand the molecular features that underlie heterogeneity in the long-term maintenance of cell-cycle arrest in response to ionizing radiation. Over long timescales, unrepaired sites of damage become large chromatin domains, known as persistent DNA damage foci (Fumagalli et al., 2014). Given the active role that residual DNA damage plays in the maintenance of cell-cycle arrest and senescence (Fumagalli et al., 2014), we asked whether complete repair of DNA damage was a necessary condition for cell-cycle reentry after a prolonged period of cell-cycle arrest. We used a previously established mCherry-53bp1 fluorescent fusion protein to quantify the amount of DNA damage in live individual cells (Dimitrova et al., 2008; Karanam et al., 2012). Upon irradiation, mCherry-53bp1 accumulated at sites of double-stranded DNA breaks, forming discrete foci (Figure 2A). We showed that the mCherry-53bp1 reporter co-localizes with the canonical double-stranded DNA breaks marker γ-H2AX after irradiation, making it a suitable proxy to quantify the persistence of DNA damage in our experimental live-cell system (Figure 2B).

To test whether late-dividing cells undergo G1-S transitions in the presence of DNA damage, we treated cells with a high dose of γ-irradiation and imaged them 4 days after DNA damage, for 48 hr. We selected a dose of 20 Gy, after which cells either do not divide or divide only once within the timescale of our observations (Figure 1D). This allowed us to avoid the confounding effect of a small subpopulation of cells that recover from damage and divide multiple times after lower doses of DNA damage. We used the mVenus-hGemini(1–100) reporter (Sakaue-Sawano et al., 2008) to identify entry into S phase and the mCherry-53bp1 to measure DNA damage in each cell. We found that cells that underwent G1-S transitions did so with DNA damage. Specifically, they exhibited a significantly higher number of 53bp1 foci than nonirradiated cells (Figures 2C–2E), implying that complete repair of DNA damage is not necessary to trigger cell-cycle reentry in long-term-arrested cells. Notably, within the irradiated population, there was no significant difference between the number of 53bp1 foci in cells that escaped the arrest and those that remained arrested (Figure 2E). These results suggest that late divisions do not reflect heterogeneity in the presence or amount of DNA damage; rather, they result from variability in the way individual cells process information about the presence of DNA damage through downstream signaling. We termed these late dividing cells “escapers”.

**Cell-to-Cell Variation in p53 and p21 Contributes to Heterogeneity in Arrest Maintenance**

We next sought to determine whether escape events can be explained by levels of the molecular players that control cell-cycle arrest. The p53 signaling pathway links DNA damage sensing with cell-cycle arrest (Lakin and Jackson, 1999) (Figure 3A). Upon irradiation, the tumor-suppressor protein p53 transcriptionally activates hundreds of genes involved in DNA repair, cell-cycle arrest, and apoptosis (Riley et al., 2008). One of such target genes, the cyclin-dependent kinase (CDK) inhibitor p21 (el-Deiry et al., 1993), directly enforces G1 arrest through sequestration of cyclin-CDK complexes (Brugarolas et al., 1995; Harper et al., 1993; Xiong et al., 1993). Both p53 and p21 are necessary to establish cell-cycle arrest after damage in our experimental system (Figure 3B). Note that knockdown of p53 either 2 days or 7 days post-irradiation triggered cell-cycle reentry in arrested cells (Figure 3C), demonstrating the important role that p53 signaling plays for both the establishment and maintenance of the arrested state.

To quantitatively measure the relationship between p53, p21, and escape from arrest in individual cells we established a cell line harboring a p53 fluorescence fusion protein (Lahav et al., 2004) and applied a recently developed CRISPR-Cas9-based method (Stewart-Ornstein and Lahav, 2016) to endogenously tag p21 with the fluorescent protein mKate2 (Figures 3D and S1). After irradiation, we observed that oscillatory p53 dynamics, which were previously described during the first 24 hr post-irradiation (Geva-Zatorsky et al., 2006), persisted for the entire experimental time period (5 days) (Figure 3E; Video S2). In agreement with studies on single-cell p53 dynamics in other cell lines (Loewer et al., 2013; Stewart-Ornstein and Lahav, 2017), p53 oscillations had a stereotyped period of ~5.5 hr (Figure 3F) and noisy amplitude (Figure 3E), which was previously attributed to arise from intrinsic fluctuations in the feedback control circuit that generates p53 pulses (Geva-Zatorsky et al., 2006). Following p53 activation, p21 levels increased (Figure 3E; Video S2). The derivative of p21 trajectories revealed a pulsatile rate of protein accumulation that matched upstream p53 dynamics with a time delay of ~1.5 hr (Figure S2). Beyond this tight relationship at the level of dynamics, we observed a correlation between the amplitude of individual p53 pulses and subsequent p21 protein production pulses (Spearman’s r = 0.41), suggesting that fluctuations in p53 pulse amplitude contribute to temporal variability in p21 protein levels (Figure S2).

We next asked whether general features of p53 and p21 trajectories can explain the difference between escaper and nondivider cells. While the period of p53 oscillations was similar between nondividing and escaper cells (Figure 3F), escapers fluctuated around a lower average amplitude of p53 compared to nondividers (Figure 3G). Similar behavior was observed for the average levels of p21 (Figure 3H), which correlated with p53 average amplitude at the single-cell level (Figure 3I). Consistently, we were able to increase the percentage of cell-cycle reentry of arrested cells by partial small interfering RNA (siRNA)-based knockdown of p53 (Figures 3J and 3K). Taken together, our results show that cell-to-cell variability in the levels of p53 and p21 contribute to heterogeneity in the long-term maintenance of cell-cycle arrest after irradiation, with escapers...
having overall lower average levels of p53 and p21 than nondividing cells.

**Escape from Cell-Cycle Arrest Is Characterized by a Sharp Switch in p21 Levels and CDK2 Activity**

While the distributions of p53 and p21 differed significantly between non-dividers and escapers, there was an extensive overlap between them (Figures 3G and 3H), suggesting that properties of p53 and p21 trajectories beyond average levels might be linked to the escape from arrest. To further understand the series of molecular events leading to arrest escape, we followed p21 dynamics in individual cells several hours before cell-cycle reentry. While nondividers maintained sustained p21 levels through the entire duration of the experiment (Figure 4A), p21 was sharply degraded upon escape from G1 arrest and S phase entry (Figures 4B and S3). Note that this sharp degradation of p21 in escaper cells was preceded by significantly higher levels of p21 compared to regularly dividing cells that enter S phase (Figure S4), arguing that elevated levels of p21 are insufficient for preventing escape.
Figure 3. Cell-to-Cell Variation in DNA Damage Signaling Contributes to Heterogeneity in Arrest Maintenance

(A) In response to DNA damage, p53 transcriptionally activates p21, which arrests the cell cycle. (B) Fraction of cells that divide after siRNA-mediated depletion of p53 or p21 2 days post-DNA damage (20 Gy). n > 150 cells per condition. ***p < 0.001, two-tailed binomial proportion test. (C) siRNA-mediated depletion of p53 at 2 days or 7 days post-DNA damage leads to cell-cycle reentry. n > 90 cells per condition. **p < 0.01, two-tailed binomial proportion test. (D) A fluorescent fusion protein and endogenous tagging allow quantification of p53 and p21 dynamics in individual cells. (E) Representative p53 and p21 trajectories of a single cell exposed to 10 Gy γ-irradiation. (F) Autocorrelation functions estimated from p53 trajectories of nondividers (n = 1105 cells) or escapers (n = 503 cells). Bold lines and shaded areas correspond to median and interquartile range, respectively. (G and H) Distributions of mean p53 pulse amplitude (G) and mean p21-mKate2 intensity (H) after γ-irradiation in escapers (n = 503 cells) and non-dividers (n = 1105 cells). (I) Correlation of mean p53 pulse amplitude and mean p21-mKate2 intensity. Each dot represents a single cell. (J) Distributions of p53-mNeonGreen intensity in p53 (4 nM and 32 pM) or control siRNA-treated cells 3 days post-irradiation (20 Gy). (K) Fraction of cells that divide within 2 days after siRNA knockdown (I). n > 100 cells per condition. ***p < 0.001, two-tailed binomial proportion test.

p21 halts cell-cycle progression through direct inhibition of cyclin-CDK complexes. The cyclin E-CDK2 complex in turn drives transition into S phase and leads to p21 degradation, forming a double-negative feedback loop (Bornstein et al., 2003; Coleman et al., 2015) (Figure 4C). Using a recently established translocation-based fluorescent reporter of CDK2 activity (Spencer et al., 2013) (Figures 4D and 4E), we found that the onset of sharp p21 degradation was tightly correlated with an increase in CDK2 activity, leading to a switch in the abundance and activity of these two molecular players (Figure 4F; Video S3). This transition occurred within a time window of ~5 hr (Figures S5A and S5B).

To understand the relationship between p21 behavior and the probability of escape, we pooled single-cell trajectories from escapers and non-dividers and computed the fraction of cells that would reenter the cell cycle within a 5-hr escape window as a function of p21 levels or fold increase/ decrease over time. The probability of escape increased sharply with decreasing p21 levels and increasing fold degradation (STAR Methods; Figures S5D and S5E). Instantaneous levels and dynamics of p21 protein were most informative of escape in the vicinity of G1/S transitions (~20 hr). High p21 levels at any given time point increased the probability but did not guarantee the long-term maintenance of the arrested state (Figures S5F and S5G). Thus, while elevated levels of p21 in escapers are sufficient to establish and maintain the arrested state for several days, transient excursions into low levels can trigger the switch between p21 and CDK2. We sought to determine the molecular mechanism that triggers such an abrupt switch.

Escape from Arrest Is Preceded by a Transient Decrease in p53 Pulse Amplitude

The interplay between p21 and CDK2 can function as a bistable toggle switch in a deterministic dynamics framework (Tyson and Novák, 2010). To understand whether and how this core circuit might be affected by temporal fluctuations stemming from p53, we adapted a previously developed computational model that...
accounts for deterministic dynamics of p21 and CDK2 interactions (Overton et al., 2014) and introduced pulsatile p53 dynamics as the only source of variation in p21 production (Figure 5A; Supplemental Information). We used single-cell time series of paired p53/p21 and p21/CDK2 reporters to calculate the dynamics as the only source of variation in p21 production (Figure 5B), which serves as a quantitative proxy for CDK2 activity in live cells.

![Figure 5A](image1.png)

Figure 5A. Representative single-cell trajectories of p21-mKate2 levels in irradiated non-dividers (A) and escapers (B). Vertical lines denote mitosis events.

To test whether local fluctuations in p53 pulse amplitude indeed contribute to escape from cell-cycle arrest, we quantified p53 dynamics in >1,000 individual cells that either maintained or escaped the arrested state in the course of 5 days post-irradiation (Figure 5D). We used p21 degradation events to in silico align individual cells at the time of cell-cycle reentry, allowing a retrospective view at p53 fluctuations in the vicinity of G1-S transitions. We found that a transient decrease in p53 pulse amplitude preceded the onset of p21 degradation events, a pattern that was absent in trajectories from non-dividers randomly sampled to match the timing of unsynchronized escape events (Figure 5E).

**Transient Artificial Decrease of p53 Is Sufficient to Trigger Cell-Cycle Reentry in Arrested Cells**

We next sought to directly manipulate p53 levels in a way that mimics the natural noise and test the effect on cell-cycle reentry in long-term-arrested cells. To this end, we established a system that allows quantitative and reversible control of the p53 protein levels in single cells. Such system consists of two inducible components: (1) a doxycycline (dox)-inducible small hairpin RNA (shRNA) targeting the endogenous p53 mRNA (Figure 6A) and (2) a shRNA-resistant exogenous p53 (Gaglia et al., 2013) fused to the YFP fluorescent protein and to a DHFR destabilizing domain (Iwamoto et al., 2010) (Figure 6B), which render the protein stable only in the presence of trimethoprim (TMP). Addition of TMP in the absence of damage resulted in similar p53 levels to those of untreated cells, implying that the DHFR(DD)-YFP domains do not disrupt homeostatic control of p53 under basal conditions (Figure 6C). Addition of dox alone prevented the induction of p53 protein in response to ionizing radiation, which was rescued by TMP-mediated stabilization of p53-DHFR(DD)-YFP (Figure 6C). Treatment with increasing concentrations of TMP led to a gradual increase in the average p53 levels (Figure 6D), which translated into a gradual increase in p21 protein levels (Figure 6E) and a corresponding decrease in the fraction of cycling cells in the population, as quantified by pRb staining (Figure 6F). Moreover, we showed that p53 pulse amplitude could be reversibly modulated by changing the concentration of TMP over time (Figure 6G). Taken together, our inducible system allows the functional replacement of the endogenous p53 for an exogenous version of p53 that can be simultaneously quantified and modulated.

We used TMP-mediated control of p53 levels to transiently disrupt p53 function in long-term-arrested cells and to test the effect of such perturbation on the maintenance of the arrested state. We exposed cells to a high dose of ionizing radiation and treated them with TMP and dox 4 days after...
damage. Five days post-irradiation, we washed the media and either fully removed TMP or reduced its concentration (0.5 μM) for 10 hr, after which TMP was replenished to its original concentration (Figure 6H). We found that transient reduction or depletion of p53 levels was sufficient to disrupt cell-cycle arrest, as measured by an increase in the fraction of cells incorporating the thymidine analog 5-Iodo-2'-deoxyuridine (IdU) (Figure 6I). Moreover, lowering the basal level of p53 before porating the thymidine analog 5-Iodo-2'-deoxyuridine (IdU) rest, as measured by an increase in the fraction of cells incorporating IdU, was sufficient to disrupt cell-cycle arrest. Critical time windows but amounted to an appreciable frequency as long-term observations allowed us to identify cell-cycle arrest differed in their ability to maintain cell-cycle arrest. Critically, long-term observations allowed us to identify cell-cycle arrest escape events in real time. These events were rare within short time windows but amounted to an appreciable frequency as the timescale of escape events. Median trajectories ± interquartile range are compared in the vicinity of G1-S transitions.

The variability in cellular outcomes after DNA damage is often conceptualized in the form of discrete cellular states (i.e., arrested and cycling) that are quantifiable at fixed time points or short-term observations. By studying the response of single cells to ionizing radiation, we discovered that arrested cells differed in their ability to maintain cell-cycle arrest. Critically, long-term observations allowed us to identify cell-cycle arrest escape events in real time. These events were rare within short time windows but amounted to an appreciable frequency as the timescale of escape events. Median trajectories ± interquartile range, respectively. n = 503 cells per group. ***p < 0.001, two tailed t test. Escapers show overall lower levels of p53 than nondividers and a dip in p53 pulse amplitude prior to the p21/Cdk2 switch that leads to the escape from arrest.

DISCUSSION

The variability in cellular outcomes after DNA damage is often conceptualized in the form of discrete cellular states (i.e., arrested and cycling) that are quantifiable at fixed time points or short-term observations. By studying the response of single cells to ionizing radiation, we discovered that arrested cells differed in their ability to maintain cell-cycle arrest. Critically, long-term observations allowed us to identify cell-cycle arrest escape events in real time. These events were rare within short time windows but amounted to an appreciable frequency as the timescale of escape events. Median trajectories ± interquartile range, respectively. n = 503 cells per group. ***p < 0.001, two tailed t test. Escapers show overall lower levels of p53 than nondividers and a dip in p53 pulse amplitude prior to the p21/Cdk2 switch that leads to the escape from arrest.
Stable p53

40

70

30

0.5

2.5

80

10

60

0

1

20

50

603.0x783.0

Molecular Cell

588

active proliferation, as quantified by colony formation in vitro which sporadic escape events lead to the reestablishment of Moreover, it will be important to determine the conditions under from studying the arrested state at these two different levels. Moreover, it will be important to determine the conditions under which sporadic escape events lead to the reestablishment of active proliferation, as quantified by colony formation in vitro and tumor growth in vivo.

Early efforts to determine the molecular players required for the maintenance of the senescent phenotype identified a critical role for p53 signaling (Beauséjour et al., 2003; Dirac and Bernard, 2003; Gire and Wynford-Thomas, 1998). While it has been shown that other signaling pathways can compensate for p53 loss after the establishment of senescence (Jacobs and de Lange, 2004; Mallette et al., 2004; Sugrue et al., 1997), it has also been shown that loss of p53 could lead to the establishment of senescence identified a critical role for p53 signaling (Beauséjour et al., 2003; Dirac and Bernard, 2003; Gire and Wynford-Thomas, 1998). While it has been shown that other signaling pathways can compensate for p53 loss after the establishment of senescence (Jacobs and de Lange, 2004; Mallette et al., 2004; Sugrue et al., 1997), it was also shown that the resumption of active proliferation (Beauséjour et al., 2003). In our system, active p53 signaling was necessary for the establishment and maintenance of cell-cycle arrest after

from studying the arrested state at these two different levels. Moreover, it will be important to determine the conditions under which sporadic escape events lead to the reestablishment of active proliferation, as quantified by colony formation in vitro and tumor growth in vivo.

Early efforts to determine the molecular players required for the maintenance of the senescent phenotype identified a critical role for p53 signaling (Beauséjour et al., 2003; Dirac and Bernard, 2003; Gire and Wynford-Thomas, 1998). While it has been shown that other signaling pathways can compensate for p53 loss after the establishment of senescence (Jacobs and de Lange, 2004; Mallette et al., 2004; Sugrue et al., 1997), it was also shown that the resumption of active proliferation (Beauséjour et al., 2003). In our system, active p53 signaling was necessary for the establishment and maintenance of cell-cycle arrest after

...from studying the arrested state at these two different levels. Moreover, it will be important to determine the conditions under which sporadic escape events lead to the reestablishment of active proliferation, as quantified by colony formation in vitro and tumor growth in vivo.

Early efforts to determine the molecular players required for the maintenance of the senescent phenotype identified a critical role for p53 signaling (Beauséjour et al., 2003; Dirac and Bernard, 2003; Gire and Wynford-Thomas, 1998). While it has been shown that other signaling pathways can compensate for p53 loss after the establishment of senescence (Jacobs and de Lange, 2004; Mallette et al., 2004; Sugrue et al., 1997), it was also shown that the resumption of active proliferation (Beauséjour et al., 2003). In our system, active p53 signaling was necessary for the establishment and maintenance of cell-cycle arrest after

...from studying the arrested state at these two different levels. Moreover, it will be important to determine the conditions under which sporadic escape events lead to the reestablishment of active proliferation, as quantified by colony formation in vitro and tumor growth in vivo.

Early efforts to determine the molecular players required for the maintenance of the senescent phenotype identified a critical role for p53 signaling (Beauséjour et al., 2003; Dirac and Bernard, 2003; Gire and Wynford-Thomas, 1998). While it has been shown that other signaling pathways can compensate for p53 loss after the establishment of senescence (Jacobs and de Lange, 2004; Mallette et al., 2004; Sugrue et al., 1997), it was also shown that the resumption of active proliferation (Beauséjour et al., 2003). In our system, active p53 signaling was necessary for the establishment and maintenance of cell-cycle arrest after

...from studying the arrested state at these two different levels. Moreover, it will be important to determine the conditions under which sporadic escape events lead to the reestablishment of active proliferation, as quantified by colony formation in vitro and tumor growth in vivo.

Early efforts to determine the molecular players required for the maintenance of the senescent phenotype identified a critical role for p53 signaling (Beauséjour et al., 2003; Dirac and Bernard, 2003; Gire and Wynford-Thomas, 1998). While it has been shown that other signaling pathways can compensate for p53 loss after the establishment of senescence (Jacobs and de Lange, 2004; Mallette et al., 2004; Sugrue et al., 1997), it was also shown that the resumption of active proliferation (Beauséjour et al., 2003). In our system, active p53 signaling was necessary for the establishment and maintenance of cell-cycle arrest after

...from studying the arrested state at these two different levels. Moreover, it will be important to determine the conditions under which sporadic escape events lead to the reestablishment of active proliferation, as quantified by colony formation in vitro and tumor growth in vivo.

Early efforts to determine the molecular players required for the maintenance of the senescent phenotype identified a critical role for p53 signaling (Beauséjour et al., 2003; Dirac and Bernard, 2003; Gire and Wynford-Thomas, 1998). While it has been shown that other signaling pathways can compensate for p53 loss after the establishment of senescence (Jacobs and de Lange, 2004; Mallette et al., 2004; Sugrue et al., 1997), it was also shown that the resumption of active proliferation (Beauséjour et al., 2003). In our system, active p53 signaling was necessary for the establishment and maintenance of cell-cycle arrest after

...from studying the arrested state at these two different levels. Moreover, it will be important to determine the conditions under which sporadic escape events lead to the reestablishment of active proliferation, as quantified by colony formation in vitro and tumor growth in vivo.

Early efforts to determine the molecular players required for the maintenance of the senescent phenotype identified a critical role for p53 signaling (Beauséjour et al., 2003; Dirac and Bernard, 2003; Gire and Wynford-Thomas, 1998). While it has been shown that other signaling pathways can compensate for p53 loss after the establishment of senescence (Jacobs and de Lange, 2004; Mallette et al., 2004; Sugrue et al., 1997), it was also shown that the resumption of active proliferation (Beauséjour et al., 2003). In our system, active p53 signaling was necessary for the establishment and maintenance of cell-cycle arrest after...
DNA damage (Figure 3C); however, even in the presence of functional p53, fluctuations in its levels led to heterogeneity in arrest maintenance. Future work will address the extent to which combinatorial fluctuations in distinct tumor-suppressor pathways collectively shape the propensity for cells to maintain or escape cell-cycle arrest.

Whether cellular noise is beneficial for cells or an obstacle they need to buffer and cope with is an unsolved question. On one hand, fluctuations in signaling have been functionally linked to the generation of phenotypic diversity that shapes multicellular development (Kalmar et al., 2009) and allows cells to adapt to changes in their environment (Balaban et al., 2004; Spencer et al., 2009). On the other hand, noise was shown to impose limits to cellular control circuits (Lestas et al., 2010), and cells have evolved mechanisms to distinguish bona fide signals from noise (Sokolik et al., 2015). In the context of DNA damage signaling, it is surprising that cells reenter the cell cycle even in the presence of high DNA damage (Figure 2). One potential explanation is that p53 pulses emerged as a trade-off between the need to respond sensitively to DNA damage while maintaining the potential to recover. Indeed, previous work showed that perturbation of p53 from pulsatile to sustained dynamics led to loss of proliferation potential in cells that would otherwise recover from damage (Purvis et al., 2012). While periodic p53 activation allows recovery, it also creates the risk of escape from arrest as a byproduct of its fluctuations.

Self-reinforcing feedback loops confer stability to cellular states but can also be engaged in triggering switches from noisy input signals. The double-negative feedback between p21 and CDK2 has been implicated in the establishment of quiescence after mitosis by reinforcing initial accumulation of p21 during pre-CDK2 has been implicated in the establishment of quiescence states but can also be engaged in triggering switches from noisy signaling, it is surprising that cells reenter the cell cycle even in the presence of high DNA damage (Figure 2). One potential explanation is that p53 pulses emerged as a trade-off between the need to respond sensitively to DNA damage while maintaining the potential to recover. Indeed, previous work showed that perturbation of p53 from pulsatile to sustained dynamics led to loss of proliferation potential in cells that would otherwise recover from damage (Purvis et al., 2012). While periodic p53 activation allows recovery, it also creates the risk of escape from arrest as a byproduct of its fluctuations.

Self-reinforcing feedback loops confer stability to cellular states but can also be engaged in triggering switches from noisy input signals. The double-negative feedback between p21 and CDK2 has been implicated in the establishment of quiescence after mitosis by reinforcing initial accumulation of p21 during previous S and G2 phases (Barr et al., 2017; Spencer et al., 2013). Moreover, it was shown that this circuit translates heterogeneity in p21 levels into fractional quiescence entry during unperturbed proliferation (Overton et al., 2014). Here, we found that the same circuit architecture is engaged during escape from arrest. Critically, levels of p21 that were sufficient to trigger entry into the arrested state, were not necessarily sufficient to maintain such state in the presence of fluctuations from upstream signaling. Specifically, the double-negative interaction between p21 and CDK2 amplified transient decreases in p53 pulse amplitude, ultimately leading to the disruption of cell-cycle arrest homeostasis. A similar noise-driven switch has been shown in Bacillus subtilis, in which an excitable system of negative and positive feedback loops led to transient differentiation (Süel et al., 2006). Thus, the interaction between noisy signaling and self-reinforcing circuits may be a common mechanism that translates fluctuations that occur over short timescales in phenotypic changes that occur over long timescales. Careful quantitative measurements in single cells will be required to determine whether other systems show similar responses to fluctuations and specifically whether cells can more efficiently filter fluctuations in oscillatory systems than non-oscillatory systems, giving some hints to the puzzling function of oscillations in transcriptional regulation.

The long-term maintenance of cell-cycle arrest has emerged as a critical factor for the success of cytotoxic chemotherapy (Collado and Serrano, 2010; Nardella et al., 2011). However, it has been recognized that specific genetic alterations, including p53 loss, can lead to the reversion of the senescent phenotype and the resumption proliferation (Beauséjour et al., 2003; Dirac and Benard, 2003). This raises the possibility that therapy-induced senescent cells may constitute a latent cellular reservoir of damaged cells that could contribute to tumor repopulation and the emergence of resistance to DNA damaging therapies (Kahlem et al., 2004; Schmitt et al., 2002). Little is known about the dynamics by which cells could transition from a senescent state into active proliferation. It will be important to understand the extent to which noise-driven cell-cycle reentry can facilitate early stages of senescence escape and the emergence of resistance to cytotoxic chemotherapy.
Children Memorial Fund for Medical Research. J.S.-O. received support from the NIH (grant CA207727). Materials and data are available upon request.

AUTHOR CONTRIBUTIONS


DECLARATION OF INTERESTS

The authors declare no competing interests.

Received: November 13, 2017
Revised: April 10, 2018
Accepted: June 20, 2018
Published: July 26, 2018

REFERENCES


590 Molecular Cell 71, 581–591, August 16, 2018


### KEY RESOURCES TABLE

<table>
<thead>
<tr>
<th>REAGENT or RESOURCE</th>
<th>SOURCE</th>
<th>IDENTIFIER</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Antibodies</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mouse monoclonal anti-p21WAF1</td>
<td>EMD Millipore</td>
<td>Ab-1; cat#OP64; RRID: AB_213423</td>
</tr>
<tr>
<td>Mouse monoclonal anti-H2A.X-P(Ser139)</td>
<td>Millipore</td>
<td>JBW301; RRID: AB_310795</td>
</tr>
<tr>
<td>Mouse monoclonal anti-p53</td>
<td>Santa Cruz Biotechnology</td>
<td>sc-126 (DO1); RRID: AB_628082</td>
</tr>
<tr>
<td>Rabbit polyclonal anti-RFP</td>
<td>MBL international</td>
<td>PM005; RRID: AB_591279</td>
</tr>
<tr>
<td>Mouse monoclonal anti-β-Actin</td>
<td>Sigma</td>
<td>A5316; RRID: AB_476743</td>
</tr>
<tr>
<td><strong>Bacterial and Virus Strains</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>pRRL-UbCp-p53-mNeonGreen (lentiviral)</td>
<td>This work</td>
<td>N/A</td>
</tr>
<tr>
<td>CSII-EF-H2B-mTurquoise (lentiviral)</td>
<td>Spencer et al., 2013</td>
<td>N/A</td>
</tr>
<tr>
<td>CSII-EF-mVenues-hGeminin(1–110) (lentiviral)</td>
<td>Sakaue-Sawano et al., 2008</td>
<td>N/A</td>
</tr>
<tr>
<td>pRRL-EF1α-mCerulean3--NLS (lentiviral)</td>
<td>This work</td>
<td>N/A</td>
</tr>
<tr>
<td>CSII-EF-DHB-mVenus (lentiviral)</td>
<td>Spencer et al., 2013</td>
<td>N/A</td>
</tr>
<tr>
<td>Tet-on-pLKO -p53siRNA (lentiviral)</td>
<td>This work</td>
<td>N/A</td>
</tr>
<tr>
<td>pRRL-UbCp-p53-DHFR(DD)-YFP (lentiviral)</td>
<td>This work</td>
<td>N/A</td>
</tr>
<tr>
<td><strong>Chemicals, Peptides, and Recombinant Proteins</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>DAPI</td>
<td>Sigma</td>
<td>D9542</td>
</tr>
<tr>
<td>Trimethoprim (TMP)</td>
<td>Sigma</td>
<td>T7883-5G</td>
</tr>
<tr>
<td>(+)-5-Iodo-2′-deoxyuridine (IdU)</td>
<td>VWR</td>
<td>AAA11542-03</td>
</tr>
<tr>
<td>SMARTpool: siGENOME CDKN1A (p21) siRNA</td>
<td>Dharmacon</td>
<td>M-003471-00-0005</td>
</tr>
<tr>
<td>SMARTpool: siGENOME TP53 (p53) siRNA</td>
<td>Dharmacon</td>
<td>M-003329-03-0005</td>
</tr>
<tr>
<td><strong>Critical Commercial Assays</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>RNAi MAX transfection reagent</td>
<td>Thermo Fisher Scientific</td>
<td>cat#13778075</td>
</tr>
<tr>
<td>Senescence Associated β-Galactosidase staining kit</td>
<td>Cell Signaling Technology</td>
<td>cat#9860</td>
</tr>
<tr>
<td><strong>Experimental Models: Cell Lines</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Human: RPE hTERT</td>
<td>S.J. Elledge Lab (Harvard Medical School)</td>
<td>N/A</td>
</tr>
<tr>
<td>Human: RPE + mVenues-hGeminin(1–110) + H2B-mTurquoise</td>
<td>This work</td>
<td>N/A</td>
</tr>
<tr>
<td>Human: RPE p21-mKate2 + UbCp-p53-mNeonGreen</td>
<td>This work</td>
<td>N/A</td>
</tr>
<tr>
<td>Human: RPE p21-CFP + mVenues-hGeminin(1–110) + EF1α-mCherry-53bp1</td>
<td>This work</td>
<td>N/A</td>
</tr>
<tr>
<td>Human: RPE p21-mKate2 + DHB-mVenus</td>
<td>This work</td>
<td>N/A</td>
</tr>
<tr>
<td>Human: RPE + tel-p53 shRNA + p53(shRNA-resistant)-DHFR(DD)-YFP</td>
<td>This work</td>
<td>N/A</td>
</tr>
<tr>
<td><strong>Oligonucleotides</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Homology donor p21 REV: CCACTCCAAACGCCGGCTGA</td>
<td>Stewart-Ornstein and Lahav, 2016</td>
<td>N/A</td>
</tr>
<tr>
<td>TCTTCTACAGAGGGGAGCCC GGAGCCTGTGACGCG</td>
<td>Stewart-Ornstein and Lahav, 2016</td>
<td>N/A</td>
</tr>
<tr>
<td>Homology donor p21 REV: GGCCTCCTCTCCTCGAGAC GAGGTGAGGGGCGGAG GGGGAGCCC GGAGCCTGTGACGCG</td>
<td>Stewart-Ornstein and Lahav, 2016</td>
<td>N/A</td>
</tr>
<tr>
<td><strong>Recombinant DNA</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>pEF1α-mCherry-53BP1</td>
<td>Karanam et al., 2012</td>
<td>N/A</td>
</tr>
<tr>
<td>pCas9-T2A-p21 gRNA; GGCTTCTGTGGGGATTA</td>
<td>Stewart-Ornstein and Lahav, 2016</td>
<td>N/A</td>
</tr>
<tr>
<td><strong>Software and Algorithms</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Custom MATLAB script – Model</td>
<td>This work</td>
<td>Methods S1</td>
</tr>
</tbody>
</table>
CONTACT FOR REAGENT AND RESOURCE SHARING

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Galit Lahav (galit@hms.harvard.edu).

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Cell Lines
RPE cells immortalized with telomerase overexpression (RPE-hTERT, a kind gift from S.J. Elledge, Harvard Medical School) were grown in DMEM/F12 supplemented with 10% fetal bovine serum (FBS), 100U/mL penicillin, 100mg/mL streptomycin and 250ng/mL fungizone (Gemini Bio-Products). For microscopy, DMEM/F12 + 5% FBS lacking phenol red and riboflavin, was used. When necessary, media was supplemented with selective antibiotics (400 μg/mL G418, 2 μg/mL puromycin). Cells were grown in an incubator with controlled temperature (37°C, CO2 (5%) and relative humidity (80%). RPE cells were authenticated by Short-Tandem-Repeats at the Dana Farber Molecular Diagnostics Laboratory.

Cell Line Construction
To endogenously tag p21 with the fluorescent protein mKate2, we used the eFlut toolset as previously described (Stewart-Ornstein and Lahav, 2016). The resulting tagged coding sequence includes a P2A cleavable Neomycin resistance protein, allowing selection for cells harboring the tag without disrupting the endogenous p21 3’UTR. To establish the p21-mKate2/p53-mNeonGreen dual reporter cell line, p21-mKate2 cells were infected with a p53-mNeonGreen lentiviral vector. Single cell clones were expanded through limited dilution and subsequently screened for marker expression. To establish the p21-mKate2/DHB-mVenus reporter line, clonal p21-mKate2 tagged cells were infected with the DHB-mVenus lentiviral vector (a kind gift from S. Spencer, University of Colorado Boulder; Spencer et al., 2013). Cells were further infected with mCerulean3-NLS lentiviral vector, a constitutive nuclear marker.

To establish the RPE p53-DHFR(DD)-YFP cell line, RPE cells were first infected with Tet-on-pLKO-shp53 which allows doxycyclin inducible expression of an shRNA targeting endogenous p53 (Brummelkamp et al., 2002; Wiederschain et al., 2009). Next, we infected cells with a construct harboring the p53 coding sequence fused to the DHFR destabilizing domain (Iwamoto et al., 2010) and YFP. The exogenous p53 has 7 synonymous point mutations that allow it to escape from the dox inducible shRNA silencing. This modified p53 was functionally validated in a previous study (Gaglia et al., 2013). Single cell clones were expanded and the effects of doxycyclin and TMP on p53 function were tested as described in Figures 6C–6F.

METHOD DETAILS

Irradiation
To acutely induce DNA damage, cells were exposed to time-controlled doses of γ-irradiation in a 60Co irradiator or X-ray in a RS-2000 irradiator (Radsource).

Live-Cell Microscopy
Cells were grown in poly-D-lysine-coated glass bottom plates (MatTek Corporation) and imaged using a Nikon Eclipse TE-2000 inverted microscope with a 10X Plan Apo objective and a Hammamatsu Orca ER camera, equipped with environmental chamber controlling temperature, atmosphere (5% CO2) and humidity. For long-term live cell imaging experiments (> 5 days), media was replaced every day to maintain growth factor stimulation throughout the entire duration of imaging. Images were acquired every 30 min using the MetaMorph Software.

siRNA Knockdown
Small interfering RNA (siRNA) targeting p53, p21 or control siRNA were delivered to irradiated cells using RNAiMAX reagent (Thermo Fisher Scientific), following manufacturer protocol. Media was replaced 5h post-delivery, and cells were imaged for >2 days to quantify cell cycle re-entry. Live reporters allowed confirmation of target knockdown.

Senescence Associated β-Galactosidase Assay
Senescence Associated β-Galactosidase activity was assayed using the corresponding staining kit purchased from Cell Signaling (cat#9860), following manufacturer protocol. Cells received γ-irradiation, were replated 3 days post-irradiation adjusting for differences in cell density associated with different damage doses, and stained for β-Galactosidase activity. Cells were imaged using a Nikon Eclipse TE-2000 inverted microscope equipped with an IDEA digital color camera and a 10X Plan Apo objective.

Immunofluorescence
Cells were plated in poly-D-lysine coated glass bottom dishes (MatTek Corporation) and fixed with 4% paraformaldehyde for 10min at room temperature at the appropriate time after treatment. Cells were permeabilized with 0.1% Triton-X and blocked with 2% BSA in 1X PBS. Cells were incubated overnight with primary antibody (1:100 dilution), washed, and stained with secondary fluorescent
antibodies and DAPI (1:10^4 dilution) for 1h, followed by a final wash. Cells were imaged using a Nikon Eclipse TE-2000 inverted microscope with a 20X Plan Apo objective and a Hamamatsu Orca ER camera.

**Western Blot**

Cells were harvested with RIPA buffer (Cold Spring Harbor Protocols) containing protease and phosphatase inhibitors. Protein samples were separated by electrophoresis on 4%–12% Bis-Tris gradient gels (Invitrogen) and transferred to Pure Nitrocellulose Blotting Membrane (Life Sciences). Membranes were blocked with 4% BSA, incubated with primary antibody overnight (1:1000-1:2000 dilution), washed, and incubated with secondary fluorescent conjugate antibodies, followed by a last wash. Membranes were scanned using Li-COR Odyssey CLX.

**Transient Perturbation of p53**

To transiently perturb p53 levels, we used trimethoprim (TMP)-mediated stabilization of an exogenous p53-DHFR(DD)-YFP construct, coupled with doxycyclin (dox)-inducible shRNA depletion of endogenous p53 mRNA. Cells received 8Gy X-ray and media was changed 4 days post-irradiation into doxycyclin (250ng/mL) and trimethoprim (basal concentration of 10uM or 2.5uM) supplemented media. Depending on treatment schedule, TMP was added to a final concentration of 10uM, 2.5uM or 0.5uM immediately after wash, and replenished to its base concentration 10h after wash. Cells were fixed 1 day after this transient perturbation. To probe for IdU incorporation, cells were treated with 2N HCl for 20min, followed by a standard immunofluorescence protocol (described above) with primary anti-BrdU staining. Cells were imaged using the Cytell Cell Imaging System (GE Healthcare Life Sciences) with 10X magnification.

**Mathematical Modeling**

The aim of our computational model was to develop an intuition regarding the way fluctuations in p21 levels stemming from variable p53 oscillations could give place to stochastic switching from a p21 high/CDK2low (arrest) to a p21 low/CDK2high (escape) state. We consider a deterministic model with four molecular species: p53 protein, p21 mRNA, p21 protein and CDK2 activity, which depends on p21 levels. This model consists of a simplified architecture of the core double negative feedback loop between p21 and CDK2, and is adapted from a previously develop computational model (Overton et al., 2014). The dynamics of this system are described by the following coupled ordinary differential equations:

\[
\frac{dp21_{mRNA}}{dt} = f(p53) - \beta_1 p21_{mRNA}
\]

\[
\frac{dp21_{prot}}{dt} = -\alpha_2 p21_{mRNA} - \beta_2 p21_{prot} - h(\text{CDK2}_{activity}) p21_{prot}
\]

\[h(\text{CDK2}_{activity}) = \lambda_3 \left( \frac{CDK2^{ag}_{activity}}{CDK2^{ag}_{activity} + K_2^{ag}} \right)\]

We assume that CDK2 activity depends on the instantaneous levels of p21:

\[\text{CDK2}_{activity} = \alpha_3 \left( \frac{K_1^{ag}}{\log(p21_{prot}) + K_1^{ag}} \right)\]

The model receives p53 time trajectories with fixed period and variable amplitude as an input. To generate trains of p53 pulses with random amplitudes, we used a custom MATLAB script that sequentially generates individual cosine peaks, sets the base at 0 and scales the amplitude to a randomly sampled value from a log normal distribution: \(\mu = \frac{m}{\sqrt{(m-CV)^2 + m^2}}\) and \(\sigma = \sqrt{\log(CV^2 + 1)}\), with m and CV specified in Table S1 and Figure S6B. Each pulse is sampled independently. Computational analysis of the effect of fluctuations in p53 pulse amplitude on the core p21-CDK2 double negative feedback loop was performed using ode45 solver in MATLAB (Mathworks). Statistics on features of p53 trajectories in the vicinity of transitions were estimated from 5,000 simulated cells (Figure 5).

**Determination of Model Parameters**

p21 mRNA half-life was previously estimated to be between 1.2 h and 3.26 h (Hafner et al., 2017; Kim et al., 2012; Yang et al., 2017). We set p21 mRNA half-life to 1.75h in our simulations.

To estimate the half-life of p21 protein in damaged cells, we quantified the decay of p21-mKate2 5h after knockdown of p53 (Figure S6C). We excluded the time points corresponding to the sharp degradation of p21 during the G1-S transition and trajectories that had residual p53 activity, as quantified by the p53-mNeonGreen reporter. We estimated a median protein half-life of ~5h (Figures S6D and S6E). This estimate is in disagreement with previous reports in the literature, which resulted in much shorter half-lives.
(30min – 1h) using population level assays (Chang and Eastman, 2012; Yang et al., 2017). This disagreement could potentially arise as a result of averaging of cells in distinct stages of the cell cycle (p21 degradation rate is markedly increased during S-phase) and thus, we think that our estimate better represents p21 half-life in our experimental conditions. However, we note that our measurements could overestimate p21 protein half-life due to the effect of residual p21 mRNA after p53 depletion, and thus represent an upper bound of the actual half-life.

We next estimated the function describing p53-dependent p21 production rate. In non-divider cells, where CDK2 activity is low we assumed that

$$\beta_2 p_{21}_{prot} \gg h(CDK2_{activity}) p_{21}_{prot}$$

And therefore p21 protein dynamics are dominated by basal degradation, yielding a simplified form of Equation 2:

$$\frac{dp_{21}_{prot}}{dt} = \alpha_3 p_{21}_{mRNA} - \beta_2 p_{21}_{prot}$$

(5)

Solving for p21 mRNA from Equation 5, we obtained

$$p_{21}_{mRNA} = \frac{1}{\alpha_2} \left( \frac{dp_{21}_{prot}}{dt} + \beta_2 p_{21}_{prot} \right)$$

(6)

Finally, solving for f(p53) in Equation 1 and substituting p21 mRNA from Equation 6 we obtained

$$f(p53) = \frac{dp_{21}_{mRNA}}{dt} + \beta_1 p_{21}_{mRNA}$$

(7)

$$\alpha_2 f(p53) = \frac{dp_{21}_{prot}}{dt^2} + \beta_2 \frac{dp_{21}_{prot}}{dt} + \beta_1 \frac{dp_{21}_{prot}}{dt} + \beta_1 \beta_2 p_{21}_{prot}$$

(8)

Note that since we don’t directly measure p21 mRNA, we can only estimate the product of p21 transcription and translation rates as a function of p53 (referred here as p21 production rate). We computed the first and second derivatives from p21 protein trajectories and estimated the temporal evolution of p21 production accounting for the previously defined p21 mRNA and protein half-lives. For each pulse of p53 we identified the corresponding pulse of p21 production and used paired pulse amplitudes to infer the input-output function linking p53 levels with p21 production. Our analysis suggests that p53 dependent p21 production is well described by Michaelis-Menten kinetics (Figure S6F):

$$f(p53) = \frac{\alpha_1 p53}{K_1 + p53}$$

(9)

We computed the average CDK2 activity as a function of p21 levels from paired measurements of p21-mKate2 and DHB-mVenus in single cells (Figure 4F). This function was well described by a decaying hill function with a hill coefficient of ~5 (Figure S6G).

Lastly, we used paired p21-mKate2 and DHB-mVenus trajectories to estimate the CDK2-dependent contribution to p21 degradation. Solving for h(CDK2_{activity}) in Equation 2,

$$h(CDK2_{activity}) = -\frac{1}{p_{21}_{prot}} \left( \frac{dp_{21}_{prot}}{dt} + \alpha_2 p_{21}_{mRNA} \right) - \beta_2$$

(10)

Figure S6H shows the way the first term in this equation, which can be inferred using only p21-mKate2 trajectories, varies as a function of CDK2_{activity}. As expected, p21 degradation increases as a function of CDK2 activity. Moreover, we don’t find evidence for saturation of p21 degradation rate within the range of CDK2 activity that cells experience during G1-S transitions. Note that since we don’t have experimental data that simultaneously monitors p53, p21 and CDK2 activity in the same cell, our measurements are expected to underestimate the contribution of CDK2 activity to p21 degradation due to the continuous production of p21 protein. In our simulations, we introduced a scaling factor λ to find a regime in which we observed heterogeneity in long-term fate (arrest maintenance or escape). Our model recapitulated the sharp increase in the probability of escape with decreasing p21 levels (Figures S6l and S6J) and increasing relative p21 degradation rate (Figure S6K). The discrepancy between simulations and experimental data could be attributed to unaccounted sources of heterogeneity in p21 dynamics (for instance, cell-to-cell variation in p21 production and degradation rates) and in other molecular players known to play a role in orchestrating G1-S transitions (for instance CyclinD/CDK4 complexes). A summary of the parameters used in our simulations is presented in Table S1.

**QUANTIFICATION AND STATISTICAL ANALYSIS**

**Single-Cell Tracking and Quantification**

A semi-automated method was developed that allows tracking and cell fate annotation of individual, fast moving cells over long timescales. Our method relies on (i) automatic identification of single cell centroids using intensity and shape information of a constitutive
nuclear marker; (ii) centroid linkage and track propagation using nearest-neighbor criteria; and (iii) real-time user correction of tracking, and annotation of cell fate events. Tracking data were then used to quantify intensity of fluorescent reporters from background subtracted images by averaging 10 pixels within the cell nucleus. In order to quantify cytoplasmic-nuclear ratio of fluorescence intensity, cytoplasmic signal was estimated by averaging the intensity of a 3 pixels-wide ring surrounding segmented nuclei, which was then divided by the average nuclear intensity. Cells were tracked using only information about a constitutively expressed nuclear marker and were thus blind to the dynamics of molecular players of interest, such as p53 and p21. Only cells that remained within the field of view throughout the entire duration of the experiment were considered for downstream analyses.

Statistical Analysis of Single-Cell Trajectories
We operationally defined escapers as those cells that (i) arrested for at least 2 days after irradiation and (ii) divided only once after this prolonged phase of cell cycle arrest within long-term observations (> 5 days). These stand in contrast with non-divider cells, which remain arrested throughout the entire duration of the experiment. We compared the levels and dynamics of p53, p21 and CDK2 between these two subpopulations, which differed in fate but were otherwise exposed to the same environment. Since escape events were unsynchronized and broadly distributed throughout the experiment, we used in silico synchronization to project trajectories into comparable time axes in relation to escape events. In addition, for each escape event we randomly selected a non-divider cell trajectory and shifted it relatively to the timing of cell cycle re-entry. This allowed us to compare non-divider and escaper cells within the same data analysis framework.

Owing to the strong p21 degradation program active during this S-phase, the onset of p21 protein accumulation is delayed in cells damaged during S-phase, (Stewart-Ornstein and Lahav, 2016). To compute historical averages of p21 levels in single cells, we considered only time points after p21 induced above basal levels, effectively controlling for cell cycle related differences in induction timing.

Estimating Escape Probability from Single-Cell Trajectories
To estimate the probability that a cell would escape from cell cycle arrest within a given time window (τ), we pooled single cell measurements of p21-mKate2 from both non-dividers and escapers from the time of p21 induction up until the time of p21 degradation. For each instantaneous measurement, we recorded the time until escape. In the case of non-dividers, this timing was set to infinity. We then quantified the fraction of cells that would escape in τ hours or less given that we observed p21 in a specific intensity bin. For our initial analysis we chose τ = 5h, as this time-frame encompassed the duration from CDK2 activation to complete p21 degradation in most escaper cells (Figures S5A and S5B). We observed an ultrasensitive response curve, with a sharp increase in escape probability as cells approached low levels of p21 (Figure S5D). Interestingly, an instantaneous measurement of the fold change in p21 (Δp21/p21) yielded a sharper change in escape probability than absolute p21 levels over this 5h timescale (Figure S5E).

To understand how fast our ability to predict cell fate decayed over time, we repeated this analysis varying τ from 30min to 64h. Since the marginal probability of escape within τ hours (independent of any information of p21) increases with τ, we normalized the conditional escape probabilities (when we consider p21) by such marginal probability. This allowed us to quantify the extent to which knowing information about p21 increased our ability to predict whether a cell would escape or not within a particular time-frame. As shown in Figures S5F and S5G, both absolute and fold change values of p21 are most predictive when we measure them close to the transition. Such predictive power decays rapidly as we increase τ up to 10h~20h and stabilize at longer timescales (Figures S5H and S5I). Intuitively, the fact that instantaneous measurements of p21 protein levels are informative of cell fate over long timescales mirrors our observation that escaper cells induce p21 to lower average levels than non-dividers, and therefore operate closer to the transition (Figure 3). While variability in levels or fold change in p21 provide additional information about the timing of escape over short timescales, this additional information is quickly lost within a time-frame corresponding to 2-4 pulses of p53.

DATA AND SOFTWARE AVAILABILITY
Custom MATLAB scripts are available upon request. The p53Cinema software package for tracking and quantifying single cell data can be downloaded at https://github.com/balvahal/p53CinemaManual.