RCC1-dependent activation of Ran accelerates cell cycle and DNA repair, inhibiting DNA damage–induced cell senescence

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ABSTRACT The coordination of cell cycle progression with the repair of DNA damage supports the genomic integrity of dividing cells. The function of many factors involved in DNA damage response (DDR) and the cell cycle depends on their Ran GTPase–regulated nuclear–cytoplasmic transport (NCT). The loading of Ran with GTP, which is mediated by RCC1, the guanine nucleotide exchange factor for Ran, is critical for NCT activity. However, the role of RCC1 or Ran GTP in promoting cell proliferation or DDR is not clear. We show that RCC1 overexpression in normal cells increased cellular Ran GTP levels and accelerated the cell cycle and DNA damage repair. As a result, normal cells overexpressing RCC1 evaded DNA damage–induced cell cycle arrest and senescence, mimicking colorectal carcinoma cells with high endogenous RCC1 levels. The RCC1-induced inhibition of senescence required Ran and exportin 1 and involved the activation of importin β–dependent nuclear import of 53BP1, a large NCT cargo. Our results indicate that changes in the activity of the Ran GTP–regulated NCT modulate the rate of the cell cycle and the efficiency of DNA repair. Through the essential role of RCC1 in regulation of cellular Ran GTP levels and NCT, RCC1 expression enables the proliferation of cells that sustain DNA damage.

INTRODUCTION

Regulation of the rate at which cells divide is critical to normal development and tissue homeostasis. Because many DNA-damaging events continuously challenge the genome integrity of dividing cells, conserved DNA damage response (DDR) signaling networks have evolved to coordinate DNA damage repair with the continuing cell cycle (Ciccia and Elledge, 2010; Smith et al., 2010). Central to DDR signaling is the activation of ATM (ataxia telangiectasia–mutated) and ATR (ataxia telangiectasia and Rad3–related) kinases. DNA damage checkpoints integrate the signaling of DNA damage, the cell cycle state, and other inputs to elicit a cell cycle pause, DNA repair or transcriptional changes, or if the damage is irreparable, programmed cell death or permanent exit from the cell cycle (Shaltiel et al., 2015).
Because separate signaling networks control the termination of DDR, the cell cycle reentry in the presence of DNA damage is also a possible outcome (Kleiblova et al., 2013; Shaltiel et al., 2015).

The activities of many factors involved in the control of cell cycle and DDR depend on their Ran GTPase–regulated nuclear–cytoplasmic transport (NCT; Pemberton and Paschal, 2005; Thompson, 2010; Blackinton and Keene, 2014), suggesting that NCT activation could promote cell cycle and DDR kinetics. The driving force of NCT is the concentration gradient of Ran-GTP across the nuclear envelope: nuclei contain high Ran-GTP concentrations, and the cytoplasmic Ran is mostly GDP bound. This striking Ran-GTP compartmentalization depends on the nuclear localization of RCC1, the guanine nucleotide exchange factor for Ran, and cytoplasmic localization of RanGAP1, the Ran GTPase–activating protein. Ran-GTP interacts with ~20 different nuclear transport receptors (NTRs), importins, and exportins (Pemberton and Paschal, 2005) and regulates either the loading or unloading of cargoes on NTRs. Cargoes are unloaded from importins that bind to Ran-GTP in the nucleus, while Ran-GTP is required for cargo loading onto exportins before their exit to the cytoplasm (Pemberton and Paschal, 2005). NTR cargoes include myriad proteins and several classes of nucleic acids, including the elf4e-dependent mRNAs for many cyclins (Culjovic et al., 2006; Muqbil et al., 2013). Following disassembly of the nuclear envelope in mitotic cells, mitotic chromosomes are surrounded by diffusional Ran-GTP gradients, which support the assembly and function of mitotic spindles through the continuing dynamics of Ran-GTP–NTR interactions (Kalab et al., 2002, 2006; Forbes et al., 2015). In addition to Ran-GTP-dependent NCT, Ran-GDP functions as the nuclear import receptor for proteins carrying the ankyrin repeat motif (Lu et al., 2014).

To date, the potential role of Ran activity in controlling the rate of the cell cycle or DNA repair is supported only by indirect or correlational evidence. Namely, since premature cell aging leads to decreased NCT activity (Snow et al., 2013), the reduced nuclear transport of various DDR factors could explain the decline in DDR function observed in aging cells (Klement and Goodarzi, 2014). The requirement of Ran for the continuation of the cell cycle is consistent with the evidence that Ran knockdown by RNA interference (RNAi) induces cell senescence (Nagai and Yoneda, 2012). Furthermore, we previously showed that, compared with slow-growing cells, rapidly proliferating tissue culture cells expressed approximately four times higher levels of RCC1 (Hasegawa et al., 2013), the guanine nucleotide exchange factor for Ran, suggesting the potential role of RCC1 and Ran-GTP in accelerating the cell cycle.

In this study, we set out to investigate the role of Ran activity in cell cycle and DDR regulation using RCC1 overexpression, RNAi, and chemical inhibitors. We manipulated the RCC1-dependent Ran-GTP levels in cells and the function of two major NTRs, importin β and exportin 1. Our results demonstrated that RCC1-dependent activation of Ran-regulated NCT accelerates cell cycle and DNA damage repair, leading to evasion of DNA damage–induced cell senescence.

**RESULTS**

**RCC1 overexpression accelerates the cell cycle**

The cellular concentration of free Ran-GTP available for binding to NTRs is critical for most known Ran-regulated functions, including NCT. To determine changes in Ran regulation that could lead to increased Ran-GTP levels, we applied computational modeling of the minimal set of proteins controlling the GTP/GDP cycle on Ran. Within these models, we either increased or decreased the concentrations of their individual components and followed the changes of average cellular Ran-GTP concentration (Gorlich et al., 2003; Caudron et al., 2005; Kalab et al., 2006). The NCT rates for all Ran system components are not known. Therefore we used the previously published single-compartment models of mitotic HeLa cells (Caudron et al., 2005; Kalab et al., 2006) to simulate the Ran system in rapidly dividing mitotic cells with high Ran-GTP levels (Hasegawa et al., 2013; Figure 1B). From the relative expression in HeLa and human fibroblasts (HFF-1; Supplemental Figure S1), we deduced the composition of the Ran system in slowly dividing cells (Hasegawa et al., 2013; Figure 1A, Supplemental Figure S1, Supplemental Text S1, and Supplemental Tables S1 and S2). In both models, Ran-GTP levels were most responsive to simultaneous changes in Ran and RCC1 concentrations and reduced RanGAP1 expression (Figure 1, A and B). When overexpression of individual factors was considered, Ran-GTP was most sensitive to RCC1 concentration in the fibroblast model (Figure 1A) and to changes in Ran in the HeLa model (Figure 1B). Therefore, to examine how increased Ran-GTP levels affect the cell cycle in relatively slowly dividing cells, we tested RCC1 overexpression.

We chose the telomerase-immortalized normal epithelial RPE1 cells (hTERT-RPE1WT) as a model, because these cells display intermediate mitotic Ran-GTP gradients and Ran-GTP levels (Hasegawa et al., 2013). Because the models predicted that the effect of RCC1 on Ran-GTP would saturate at low micromolar RCC1 concentrations (Supplemental Figure S1), we selected human phosphoglycerate kinase (PGK) promoter to drive moderate overexpression of RCC1 with a C-terminal V5 tag in a stable hTERT-RPE1RCC1-V5 cell line. From immunobLOTS (Figure 1C), we estimated that the total RCC1 concentration in the hTERT-RPE1RCC1-V5 cells was four to six times higher than in the hTERT-RPE1WT cells. Such an increase in RCC1 levels was similar to the difference between HFF-1 and HeLa cells (Figure 1, A and B, and Supplemental Figure S1), suggesting that the RCC1 overexpression mimicked the physiologically relevant range. We then verified the effect of RCC1 overexpression using RBP-4, the Förster resonance energy transfer (FRET) biosensor for Ran-GTP in mitotic cells (Hasegawa et al., 2013). RBP-4 consists of a Ran-GTP-binding domain flanked by the cyan mTFP-1 donor and a nonfluorescent dsREACCh acceptor (Figure 1D; Hasegawa et al., 2013). RBP-4 binding to Ran-GTP extends the donor from the acceptor, leading to decreased FRET efficiency (E[290]FRET) and increased donor fluorescence lifetime (τdonor). Because the RBP-4 FRET activity decreases with Ran-GTP binding, the average (E[290]FRET)−1 value could be used to compare the Ran-GTP concentration between different mitotic cells quantitatively (Hasegawa et al., 2013). We applied fluorescence lifetime imaging microscopy (FLIM) to detect the changes of E[290]FRET in live mitotic cells transiently expressing the RBP-4 sensor. The mitotic hTERT-RPE1RCC1-V5 cells displayed larger differences in E[290]FRET between the chromatin and cytoplasmic areas (Figure 1F) and higher (E[290]FRET)−1 values (Figure 1G), confirming the expected increased mitotic Ran-GTP gradients and elevated Ran-GTP levels. The compartmentalization of Ran and its regulators RCC1 and RanGAP1 between the nucleus and cytoplasm precludes the direct measurements of Ran-GTP with RBP-4 FRET in interphase cells. However, previous studies showed that the expression of Ran, RCC1, and RanGAP1 remained stable during the exit from mitosis (Ciciarello et al., 2010), indicating that the mitotic Ran-GTP levels correspond to the Ran-GTP-generating activity in the interphase cells as well. Fluorescence-activated cell sorting (FACS) analysis showed that the fraction of cells in different cell cycle phases was not appreciably affected by RCC1 overexpression (Supplemental Figure S1). However, consistent with our initial hypothesis, the hTERT-RPE1RCC1-V5 cells proliferated significantly faster than the WT cells (Figure 1H; population doubling time [PDT] = 2.9 vs. 4.5 d).
Ran-GTP levels decline in nondividing cells

If the RCC1-dependent rise of Ran-GTP activates cell cycle progression, stimuli inducing cell cycle arrest should oppose RCC1 by lowering its activity or expression. To test this scenario, we induced cell cycle arrest in HFF-1 fibroblasts by a long-term in vitro culture or by inducing DNA damage through treatment with doxorubicin (Chang et al., 2002; Sliwinska et al., 2009). Both replicative exhaustion and doxorubicin treatment induced the onset of cell senescence, as indicated by the appearance of the senescence-associated β-galactosidase (SABG) staining (Debeaq-Chainiaux et al., 2009; Figure 2C) and led to a strong decline in RCC1 expression (Figure 2B). We also observed lower Ran and RanBP1 levels in senescent HFF-1 cells and a decrease in RCC1 levels in doxorubicin-treated Wi-38 and CRL-1474 primary human fibroblasts (Supplemental Figure S2). Quantification of immunoblots showed that, when normalized to the total cell protein concentration, the senescent HFF-1 cells contained ~25% RCC1 compared with early-passage cells (Figure 2B). However, because of their nearly three-times larger volume (1.27 ± 0.07 pl, passage 8 vs. 3.57 ± 0.30 pl, passage 30 HFF-1 fibroblasts), the late-passage cells contained <10% RCC1 concentration compared with the early-passage cells, confirming the strong reduction in the RCC1-dependent Ran-GTP generation in nondividing cells.

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**FIGURE 1:** RCC1-dependent increase in Ran GTP accelerates the cell cycle. (A and B) Changes in cellular Ran-GTP concentration analyzed by computational models of the minimal Ran system in a fibroblast-like cell (A) and a HeLa-like cell (B). (C) Immunoblots of RCC1 in the hTERT-RPE1WT and hTERT-RPE1RCC1-V5 cell lysates, using FLIM with RBP-4. The top row shows the donor intensity images and the bottom row shows the pseudocolor FLIM images. The range of the displayed values (corresponding to Tdonor values) is indicated beneath the FLIM panels. Scale bar: 10 µm. (F) Scatter plot of the mitotic Ran-GTP gradients quantified as the difference between the cytoplasmic and chromatin E in each cell (ΔE; single-cell data; means ± SD; t test). (G) Scatter plot of the inverse of the average cellular RBP-4 E, which is proportional to Ran-GTP concentration (E−1; single-cell data; means ± SD; t test). (H) Cell number in cultures of hTERT-RPE1WT and hTERT-RPE1RCC1-V5 cells grown in parallel. Means ± SD from two experiments performed in triplicate were fitted with exponential growth equations after 2 d from the start of culture (dashed lines) to calculate the PDT. The null hypothesis was tested: one curve for both data sets.

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**FIGURE 2:** RCC1 expression declines during cell cycle arrest. (A) Immunoblotting of total cell lysates of HFF-1 cells harvested from early (p 8) or late (p 30) passages of in vitro culture and of early-passage HFF-1 cells recovering from the treatment with doxorubicin. (B) Relative tubulin-normalized RCC1 protein expression in HFF-1 cells treated as in A (N = 3; means ± SD). (C) Micrographs of HFF-1 cells treated as in A and B and stained for the SABG. Scale bar: 100 µm.
RCC1 overexpression inhibits cell senescence

The cell cycle–promoting activity of RCC1 (Figure 1) indicated that increased RCC1 expression could attenuate DNA damage–induced cell cycle arrest. To test this idea, we compared the responses of hTERT-RPE1WT and hTERT-RPE1RCC1-V5 cells to doxorubicin treatment. Within the first 2–4 d after doxorubicin washout, most cells stopped dividing in both cultures, as indicated by the disappearance of interphase and mitotic markers (MCM2, Rad51, p-histone H3 [psH3]). At the same time, the increase in cyclin D1 levels remained stable, and SABG positivity increased over time in the hTERT-RPE1WT cells (Figure 3B). In contrast, SABG-negative and proliferating cells gradually prevailed in the hTERT-RPE1RCC1-V5 cultures (Figure 3B, arrows), concomitant with increased interphase and mitotic markers and the decline in cyclin D1 expression (Figure 3A). The quantitative capillary immunoblotting (Simple Western) analysis confirmed that, 8 d after doxorubicin treatment, hTERT-RPE1WT cells accumulated cyclin D1, while hTERT-RPE1RCC1-V5 cells resumed expression of cyclin B1 (Supplemental Figure S3). As in the senescing fibroblasts (Supplemental Figure S2), the expression of Ran decreased to ~65% in doxorubicin-treated hTERT-RPE1WT cells (Figure 3A). In contrast, Ran levels slightly increased in the hTERT-RPE1RCC1-V5 cells exposed to doxorubicin, indicating that RCC1 expression supported Ran stability in cells exposed to DNA damage (Figure 3A). Two months after the doxorubicin treatment, the hTERT-RPE1RCC1-V5 cells regained normal proliferation, while virtually no dividing cells were detectable in the hTERT-RPE1WT cells (Figure 3B). To monitor the progress of DNA damage repair, we used immunofluorescence (IF) to quantify the 53BP1 nuclear foci at 8 d of recovery from the doxorubicin. Individual cell data; means ± SD; t test representative of two experiments. (E) Column graph showing fractions of hTERT-RPE1WT and hTERT-RPE1RCC1-V5 cells that contained the indicated numbers of 53BP1 foci per nucleus during the recovery from doxorubicin treatment. Means ± SD from two independent experiments, adjusted p values from two-way analysis of variance (ANOVA) with Sidak’s multiple comparison tests.

FIGURE 3: RCC1 overexpression inhibits DNA damage–induced cell senescence in normal epithelial cells. (A) Immunoblotting of total lysates from the hTERT-RPE1WT and hTERT-RPE1RCC1-V5 cells recovering from doxorubicin treatment, showing the resumed expression of the cell cycle markers in the hTERT-RPE1RCC1-V5 cells. (B) Micrographs of cells treated as in A and stained for SABG. Scale bar: 100 μm. Arrows indicate dividing cells. (C) IF micrographs of hTERT-RPE1WT cells that contained the indicated numbers of 53BP1 nuclear foci after 8 d of recovery, and mitotic cells were already detectable (Figure 3C). In contrast, the nuclear 53BP1 foci persisted until at least 5 d after recovery, and mitotic cells were already detectable (Figure 3C). (D) Scatter plot of the cytoplasmic/nuclear ratios of the 53BP1 IF signal at 8 d of recovery from doxorubicin. Individual cell data; means ± SD; t test representative of two experiments. (E) Column graph showing fractions of hTERT-RPE1WT and hTERT-RPE1RCC1-V5 cells that contained the indicated numbers of 53BP1 foci per nucleus during the recovery from doxorubicin treatment. Means ± SD from two independent experiments, adjusted p values from two-way analysis of variance (ANOVA) with Sidak’s multiple comparison tests.
in doxorubicin-treated cells did not depend on accelerated drug efflux.

RCC1 promotes doxorubicin resistance in colorectal carcinoma cells

Because the overexpression of RCC1 prevented the onset of DNA damage–induced cell senescence in normal cells (Figure 3 and Supplemental Figure S4), RCC1 could play a role in cancer cell resistance to DNA damage. Consistent with this idea, the expression of RCC1 was found to be activated by a superenhancer element in colorectal carcinoma HCT116 cells (Hnisz et al., 2013), which is a well-studied model of resistance to DNA damage–inducing chemotherapy (Chang et al., 2002; Sliwinska et al., 2009). We used several approaches to test the role of RCC1 in HCT116 cells.

First, we compared the effects of doxorubicin on HCT116 WT and an HCT116 RCC1-V5 cell line that stably expressed V5-tagged RCC1 (Figure 4). As previously reported (Chang et al., 2002; Sliwinska et al., 2009), the doxorubicin-treated HCT116 WT cells initially stopped dividing, increased in size, and became SABG-positive (Figure 4A). Small, rapidly dividing, and SABG-negative cells prevailed in the cultures 5–7 d later (Figure 4A, arrows). A transient peak in cyclin D1 expression and a decline in S and G2/M markers (MCM2, pS10H3; Figure 4B, left) indicated a temporary cell cycle arrest. The phenotypic changes induced by doxorubicin in the HCT116 RCC1-V5 cells were similar to those seen in the HCT116 WT cells. However, the shorter duration of cyclin D1 expression and earlier recovery of the S and G2/M markers indicated that the overexpressed RCC1 slightly accelerated cell cycle reentry after DNA damage in the HCT116 RCC1-V5 cell line (Figure 4B).

We used clonogenic assays to validate the role of RCC1 expression in DNA damage recovery. After doxorubicin treatment, the fraction of colony-forming cells was nearly twice as high in HCT116 RCC1-V5 compared with HCT116 WT cells (Figure 4C), demonstrating that RCC1 overexpression strongly increased cell survival following DNA damage. By analyzing cultures derived from single doxorubicin-treated HCT116 WT cells, we found that rapidly proliferating clones contained RCC1 concentrations similar to those seen in the HCT116 WT cells, in contrast to a slow-proliferating clone (Supplemental Figure S5), indicating that the overexpressed RCC1 slightly accelerated cell cycle reentry after DNA damage in the HCT116 RCC1-V5 cell line (Figure 4B).

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Ran regulates NCT to promote the cellular response to DNA damage. Immunoblotting of total cell lysates of hTERT-RPE1 WT cells treated with control, RCC1-, or Ran-directed RNAi oligos and harvested 1 h after increasing doses of γ-irradiation. Notice the decreased intensity of the p-KAP1 and γH2AX signals in the RCC1 or Ran RNAi-treated cells. (B) IF micrographs of the γ-irradiated (5 Gy) hTERT-RPE1 WT cells showing the reduced 53BP1 recruitment to the chromatin (arrows) upon Ran or RCC1-directed RNAi. (C) Scatter plot of the 53BP1/γH2AX IF signal ratios in cells treated as in B. Individual cell data; means ± SD; ANOVA with Tukey’s test. (D) IF micrographs of hTERT-RPE1 WT, RCC1-V5, and Ran overexpression was sufficient to accelerate cell cycle and DNA damage repair (Figures 1 and 3), relatively small increases in tumor RCC1 could correspond to the activation of Ran pathways restricted to the proliferative fraction of tumor cells.

Ran promotes DNA damage signaling and repair

Apart from accelerating the cell cycle, RCC1 and Ran-GTP could support the survival from DNA damage by activating DNA repair. Consistent with the requirement for Ran activity in DNA damage signaling, immunoblotting showed that the knockdown of Ran or RCC1 by RNAi reduced the ATM-dependent phosphorylation of histone H2AX (γH2AX, Ser-139) and KAP1 (Iyengar and Farnham, 2011; p-KAP1, Ser-824) in γ-irradiated hTERT-RPE1 WT cells (Figure 5A). We applied IF in cells that were fixed with paraformaldehyde and co-permeabilized with Triton X-100 to detect changes in the recruitment of 53BP1 to the chromatin, following DNA damage (Jackson and Bartek, 2009). Although the 53BP1 expression was not affected (Figure 5A), both RCC1 and Ran knockdowns significantly reduced the ratio of nuclear 53BP1 and γH2AX signals in the irradiated cells (Figure 5C), indicating defects in 53BP1 association with the DNA damage sites marked by γH2AX signal. The knockdown of Ran had a stronger effect (Figure 5C) and caused the virtual disappearance of 53BP1 foci in a fraction of cells (Figure 5B, arrows). These results indicate that Ran activity is required for the recruitment of 53BP1 to DNA double-strand breaks, which is an essential step for their subsequent repair (Schultz et al., 2000; Callen et al., 2013).

Next we used inhibitors of importin β and exportin 1 to test the role of two major NTRs in DNA damage signaling. To that end, we

carboplatin-resistant cervical tumors (Peters et al., 2005), compared with normal tissues (Supplemental Figure S6). Because moderate RCC1 overexpression was sufficient to accelerate cell cycle and DNA damage repair (Figures 1 and 3), relatively small increases in tumor RCC1 could correspond to the activation of Ran pathways restricted to the proliferative fraction of tumor cells.

FIGURE 5: Ran-regulated NCT promotes the cellular response to DNA damage. (A) Immunoblotting of total cell lysates of hTERT-RPE1 WT cells treated with control, RCC1-, or Ran-directed RNAi oligos and harvested 1 h after increasing doses of γ-irradiation. Notice the decreased intensity of the p-KAP1 and γH2AX signals in the RCC1 or Ran RNAi-treated cells. (B) IF micrographs of the γ-irradiated (5 Gy) hTERT-RPE1 WT cells showing the reduced 53BP1 recruitment to the chromatin (arrows) upon Ran or RCC1-directed RNAi. (C) Scatter plot of the 53BP1/γH2AX IF signal ratios in cells treated as in B. Individual cell data; means ± SD; ANOVA with Tukey’s test. (D) IF micrographs of hTERT-RPE1 WT, RCC1-V5, and Ran overexpression was sufficient to accelerate cell cycle and DNA damage repair (Figures 1 and 3), relatively small increases in tumor RCC1 could correspond to the activation of Ran pathways restricted to the proliferative fraction of tumor cells.

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pretreated cells for 2 h with the exportin 1 inhibitor KPT-330 (Muqbil et al., 2013) or the importin β inhibitor importazole (IPZ; Soderholm et al., 2011) and collected samples for IF and immunoblots 1 h after γ-irradiation. IF indicated an increased number of p-KAP1 nuclear foci associated with the heterochromatin in IPZ-treated cells, while KPT-330 treatment had an opposite effect (Figure 5D). Although this trend was reproducible, the frequency of the IPZ-induced p-KAP1 foci varied between experiments. However, consistent with the IF data, immunoblotting showed that treatment with IPZ induced increased p-KAP1 signal upon irradiation, particularly in hTERT-RPE1 ΔCCT/V5 cells (Figure 5, E and F). Also in agreement with the IF data, KPT-330 caused a strong reduction in p-KAP1 signal and a more moderate decrease in γH2AX signal on immunoblots (Figure 5, E and F). Reciprocal effects of the inhibitors indicate that the interplay between exportin 1– and importin β–regulated NCT modulates the ATM-dependent phosphorylation of KAP1, which has a role in the heterochromatin DNA repair (White et al., 2012).

Because the activation of Ran promotes cell cycle progression (Figure 1), the RNAi knockdowns or inhibitor treatments described above could have induced cell cycle changes that contribute to the observed effects on DNA damage signaling and repair processes. However, the above results demonstrate that, whether directly or also through modulating the cell cycle, the activity of Ran-GTP–regulated NCT mechanisms strongly affects essential steps of DDR in cell populations.

RCC1-induced activation of Ran-regulated NCT terminates DDR

The perturbations of DNA damage signaling induced by Ran or RCC1 knockdowns (Figure 5) indicate that the mechanism of RCC1-induced recovery from DNA damage could require high Ran activity during the initial response to the damage. We examined this possibility by manipulating Ran function in cells that already sustained γ-irradiation–induced (10 Gy) DNA damage. First, we followed the irradiation of the hTERT-RPE1 WT cells by transduction with control or RCC1-V5–expressing lentiviruses (Figure 6A). As shown by the strongly reduced expression of S-phase and G2/M markers (MCM2, pS10H3), the majority of irradiated control cells stopped proliferating, as expected. The rise of transiently expressed RCC1-V5 was paralleled by a drop in cyclin D1 levels, and the renewed expression of interphase and mitotic markers (MCM2, pS10H3; Figure 6A), indicating cell cycle reentry. This result showed that the activation of Ran in cells recovering from DNA damage is sufficient to terminate the DDR and enable the continuation of cell proliferation (Figure 5).

To determine whether RCC1-induced DDR termination requires the function of exportin 1, we treated γ-irradiated hTERT-RPE1ΔCCT/V5 cells with KPT-330. The untreated hTERT-RPE1ΔCCT/V5 cells returned to cell cycle after irradiation, as shown by the expression of MCM2, pS10H3, and cyclin B1 (Figure 6B). KPT-330 treatment robustly reversed these trends and induced the accumulation of cyclin D1, indicating that DDR termination requires exportin 1–dependent nuclear export.

To verify that Ran mediates RCC1-induced DDR termination, we treated irradiated hTERT-RPE1ΔCCT/V5 cells with control or Ran-directed RNAi. As expected, RNAi controls resumed the cell cycle, as indicated by increased pS10H3 signal and reexpression of the Chk1 kinase, whose activity is essential for S-phase progression (Toledo et al., 2013; Figure 6C). The treatment with Ran RNAi in hTERT-RPE1ΔCCT/V5 cells reduced Ran levels by ~50% (Figure 6C). However, even the partial reduction in Ran expression induced severe delays in DNA repair, as shown by the persistence of phosphorylated ATM on immunoblots (Figure 6C) and 53BP1 nuclear foci in cells stained for IF (Figure 6D). Cell cycle arrest was also confirmed by continuously low levels of Chk1 and pS10H3 (Figure 6C). Similar to hTERT-RPE1 WT cells treated with doxorubicin (Figure 3), the irradiated and Ran RNAi–treated hTERT-RPE1ΔCCT/V5 cells accumulated 53BP1 in the cytoplasm (Figure 6, D and E). Therefore the reduced nuclear import of large molecular cargoes, such as 53BP1, appears to be a hallmark of cells failing to complete DNA repair as a consequence of insufficient Ran activity. In summary, these results demonstrate that RCC1-dependent activation of Ran-regulated nuclear import and export pathways facilitated the termination of DDR and cell cycle reentry.

DISCUSSION

The key finding of our study is that increased RCC1 expression accelerates cell cycle progression and DNA damage repair in a mechanism that involves raised cellular Ran-GTP levels and the function of importin β and exportin 1. Our results indicate that the overall activity of Ran-GTP–regulated nuclear export and import pathways modulates the cell cycle pace and cellular responses to DNA damages. Many of the underlying cellular and molecular mechanisms have yet to be clarified. However, our results indicate that, through the role of Ran-GTP in NCT of the cell cycle and DDR regulators, the interphase cell cycle checkpoints are sensing the RCC1-dependent Ran-GTP levels. If the cellular Ran-GTP concentration reaches sufficient threshold, Ran-GTP–regulated NCT reactions approach rates that support normal DDR function and cell cycle transitions. Consequently, DNA damage in such cells is successfully repaired, followed by termination of DDR and reentry into the cell cycle. If Ran-GTP levels fall below the required threshold, or a required NTR pathway is defective, the mechanisms of DNA repair or cell cycle reentry are delayed, triggering cell cycle arrest, as we documented, or possibly cell death.

The identity of the cell cycle transition(s) that was enabled by the RCC1 overexpression is an outstanding question raised by our results. Upon the induction of DNA damage, cell cycle reentry was induced by RCC1 in cells that exited mitosis (Figures 3 and 4), indicating that, in the majority of cells, RCC1 induced DNA damage repair and cell cycle reentry through its interphase role in NCT. The transient accumulation of cyclin D1 and absence of mitotic markers indicated that RCC1 overexpression enabled the exit from G1/S arrest. However, cells can enter senescence also from G2/M cell cycle arrest associated with high cyclin D1 expression (Gire and Dunic, 2015). Moreover, single-cell imaging studies showed that the timing of the individual cell’s commitment to S-phase reentry from the G0/G1 boundary depends on the mitosis-promoting signals in the preceding cell cycle (Spencer et al., 2013). Therefore it remains possible that, in RCC1-overexpressing cells exposed to DNA damage, activated mitotic function of Ran (Kalab and Heald, 2008; Forbes et al., 2015) enables G2/M transition and accelerates the S-phase commitment in the next cell cycle. The involvement of Ran in facilitating NCT of DNA replication factors (Liku et al., 2005; Ghosh et al., 2011) suggests that the intra S-phase cell cycle checkpoint also could be sensitive to the disruptions or activation of NCT as well. Because the WT and RCC1-overexpressing hTERT-RPE1 cells contain similar fractions of cells in G1, S, and G2/M stages (Supplemental Figure S1), it is possible that RCC1 accelerates several cell cycle phases. Future research involving synchronized cells in single-cell studies with cell cycle markers is needed to identify the specific cell cycle transitions targeted by Ran activation.

Because Ran-GTP regulates ~20 different NTRs, multiple NTRs acting in parallel could contribute to the RCC1-induced acceleration
of the cell cycle. However, it remains possible that the rate of cell cycle transitions could be particularly sensitive to NCT of only a subset of essential cargoes. The identity of such cell cycle rate-limiting NTR cargoes is not known, although published literature offers a list of many candidates. During the G1/S transition, such cargoes could include the mRNAs for many cyclins, including cyclin D1, whose elf4e adaptor-dependent nuclear export requires exportin 1 (Culjovic et al., 2006; Osborne and Borden, 2015). The S-phase entry and transition also involve Ran in the nuclear–cytoplasmic shuttling of proteins of the cyclin D, A, and E families (Malumbres and Barbacid, 2001; Jackman et al., 2002). Exportin 1 is required for the nuclear–cytoplasmic shuttling of cyclin B1 (Toyoshima et al., 1998; Gavet and Pines, 2010) and the Gwl/Mastl kinase that supports the function of the CDK1/cyclin B1 kinase (Alvarez-Fernandez et al., 2013; Wang et al., 2013). Potentially, Ran activity could modulate the timing of the activation of the cyclin B/CDK1 and Gwl/Mastl kinases during entry into mitosis. The loss of RCC1 function in temperature-sensitive tsBN2 cells induced premature entry to mitosis via the nuclear accumulation of the CDC25B phosphatase (Ohtsubo et al., 1987; Nishijima et al., 1997; Karlsson et al., 1999). Therefore the CDC25 phosphatase family could have a role in the Ran-regulated cell cycle rate. Because the phosphatases Wip1 and PP4 promote the termination of DDR signaling (Kleiblova et al., 2013; Shaltiel et al., 2014), the potential involvement of NCT in their activation is also of interest.

As indicated by the studies of Ran function in aging cells (Snow et al., 2013), large size could be one of the common features of cargoes whose NCT is particularly sensitive to Ran activity. Consistent with this idea, we observed that the knockdown of Ran induced cytoplasmic accumulation of 53BP1 (240–250 kDa; Figures 3, 5, and...
6). In addition to 53BP1, many other prominent DNA repair regulators are large, multidomain proteins whose sufficient Ran-GTP levels. However, the NCT of only a few of these regulators was studied in detail (53BP1; Moudry et al., 2012), TopBP1 (179 kDa; Bai et al., 2014), and BRCA1 (220 kDa; Thompson, 2010).

In addition to its role in NCT, the activation of Ran could promote changes in cell cycle or DDR indirectly, through gene expression changes induced by the NCT of transcriptional and epigenetic factors (Nemerug et al., 2001; Turner et al., 2012; Mcqibli et al., 2014). Such transcriptionally mediated effects of Ran possibly contribute to the activation of the cell cycle and DDR in RCC1-overexpressing cell lines. However, our results indicate that, even within the potentially diverse cellular context of cells with different RCC1 expression (Figures 5 and 6), the activity of the Ran-regulated NCT pathways was still required for the cellular response to DNA damage.

The crucial role of cellular RCC1 concentrations is a salient feature of the Ran system, as shown by computational modeling (Figure 1 and Supplemental Figure S1) and our results. Because RCC1-mediated guanine nucleotide exchange reaction on Ran is the slowest step in the GTP/GDP cycle on Ran (Supplemental Figure S1), excessive RCC1 concentration sequesters Ran, explaining the saturable effect of RCC1 expression on Ran-GTP (Supplemental Figure S1). Interestingly, RCC1 overexpression in normal cells (Figure 3) or high endogenous RCC1 expression in carcinoma cells (Figure 4) stabilized endogenous Ran levels in cells recovering from DNA damage. The underlying mechanism remains to be determined. However, the computational models of the Ran system predict that changes in RCC1 expression are amplified through such coregulation, expanding the amplitude of cellular Ran-GTP levels and their cellular effects. Consequently, DNA damage–induced decline of RCC1 in normal cells could lead to a severe impairment of NCT function as documented by the stalled nuclear import of 53BP1 (Figures 3 and 6). Because cell cycle reentry requires Ran-GTP-dependent NCT activity, the depletion of Ran-GTP levels could function as a universal mechanism that enforces stable cell cycle arrest in senescent cells.

MATERIALS AND METHODS
Cell culture
Human foreskin fibroblasts (HFF-1) from the American Type Culture Collection (ATCC, Manassas, VA), hTERT-immortalized retinal pigment epithelial cells (hTERT-RPE1; ATCC), and human embryonic kidney cells, 293FT (HEK-293FT; Invitrogen) were cultured as previously described (Kalab et al., 2006; Hasegawa et al., 2013). Human colorectal carcinoma HCT116 cells were cultured in DMEM with 10% fetal bovine serum (FBS). Normal human fibroblasts CRL1474 (ATCC) were cultured in MEM with 15% FBS and normal lung fibroblasts WI38 (ATCC) in MEM with 10% FBS. Media of cells expressing RCC1-V5 was supplemented with 10 μg/ml blasticidin (Invitrogen). All cells were grown in a 37°C humidified incubator with 5% CO₂ and in media supplemented with 100 U/ml penicillin and 10 μg/ml streptomycin. Cells were treated with Importazole (IPZ; Sigma-Aldrich, St. Louis, MO; final concentration 40 μM) and KPT-330 (Karyopharm, Newton, MA; final concentration 500 nM) by supplementing their complete cell culture with aliquots of 10 mM solutions of the drugs in dimethyl sulfoxide.

Cell count and cell volume measurement
For cell count measurement, aliquots of 50,000 cells/well were seeded in six-well plates. At different time points, cells were collected by trypsinization, washed in Dulbecco’s phosphate-buffered saline (DPBS), and fixed in Trypan Blue (15250; Invitrogen) before the cell count and average diameter were determined with Cellometer Auto T4 (Nexcelom, Lawrence, MA). The PDT was calculated from the growth curves using the equation PDT = t × log(2)/(log(C₁) – log(C₀)), where t is the time of culture, C₁ is the cell count at time t, and C₀ is the cell count at the initial t = 0. t = 0 was selected as the first time point after the initial decline in cell number after plating.

Recombinant constructs
The pLenti-PGK-RCC1-V5 construct (pK439) was prepared by the modification of the PGK-histone H2B-mCherry (21217; Addgene, Cambridge, MA; Kita-Matsuo et al., 2009). Using PCR cloning and restriction digestions, the histone H2B was replaced with an RCC1 open reading frame. The resulting pLenti-PGK-RCC1-mCherry was digested with Scal and SacI and ligated with the Scal and SacⅠ fragment from pLenti X2 Blast (17391; Addgene; Campeau et al., 2014), introducing blasticidin resistance. mCherry was excised with Age 1 and BsRG1 and replaced with a V5 sequence-coding oligo. The construct was validated by sequencing.

Lentiviral expression of RCC1-V5
Lentiviruses for the expression from pLenti-PGK-RCC1-V5 were produced as previously described (Kalab et al., 2006; Hasegawa et al., 2013). For transduction, normal cell media was replaced with media supplemented with lentiviral suspension (50–100 μl/ml) and 0.6/μl Polybrene/ml (Sigma-Aldrich). Following overnight treatment, cells were washed with warm DPBS (Life Technologies) and returned to normal growth medium. Stably expressing cells were selected by culture in media containing 10 μg/ml blasticidin. For transient expression, cells were transduced with pLenti-PGK-RCC1-V5 lentivirus 6 h after irradiation.

Induction of senescence by replicative exhaustion or DNA damage
For induction of senescence by replicative exhaustion, HFF-1 cells were kept in culture until passage 30. For induction of DNA damage by doxorubicin, normal cell media was supplemented with doxorubicin; this was followed by washes with warm DPBS and return to normal media. HFF-1, HCT116(B), HCT116(RCC1-V5), CRL1474, and WI38 cells were treated with 100 nM doxorubicin for 24 h. The hTERT-RPE1(B) and hTERT-RPE1(RCC1-V5) cells were treated for 16 h with 500 nM doxorubicin. For γ-irradiation–induced DNA damage, we used a J. L. Shepherd & Associates (San Fernando, CA) Mark-1-Model 68 Irradiator with a concealed 137 Cs source and calibrated lead shields to modulate the dose.

SABG staining
We used a previously described protocol (Debaq-Chaixnaia et al., 2009) to detect β-galactosidase activity. Bright-field micrographs were acquired using a Nikon Digital Sight color camera on a Nikon Eclipse TS100 microscope.

Electrophoresis and immunoblotting
For Supplemental Figures S1 and S2, cells were harvested and cell lysates were prepared for electrophoresis as described previously (Kalab et al., 2006; Hasegawa et al., 2013). For all other experiments, cells were trypsinized, washed in 37°C FBS-containing growth media, resuspended in 30–100 μl Clear Sample Buffer (CSB; 120 mM Tris, pH 6.8, 4% SDS, 20% glycerol), and immediately heated at 100°C for 5 min. Protein concentration was measured using a Micro BCA Protein Assay Kit (Fierce Thermo-Fisher, Carlsbad, CA) and adjusted to 4 mg/ml with CSB. Before electrophoresis, samples were mixed 1:1 with SDS-free CSB, 10% β-mercaptoethanol, and 0.5% bromophenol
blue and heated at 100°C for 2 min. For Supplemental Figures S1 and S2, proteins were separated by homemade SDS–PAGE gels (Hasegawa et al., 2013). In all other experiments, we used 4–20% Criterion gels (Bio-Rad, Hercules, CA). Gels were blotted to polyvinylidene difluoride transfer membrane (Immobilon-FL; Millipore, Bedford, MA) or Li-COR Odyssey (Li-COR Biotechnology, Lincoln, NE) nitrocellulose. Immunoblots were blocked and incubated with primary antibodies and IRDye 800– and/or IRDye 680–conjugated donkey secondary antibodies using Li-COR Odyssey buffers and protocols. The fluorescence emission was acquired using the Odyssey Imaging System (Li-COR). For each primary antibody, tubulin, actin, GAPDH, or Ponceau S–stained images of the blots were used as a loading control. The primary antibodies used for immunoblotting were from BD Biosciences (San Jose, CA; Ran, 610340; importin β, 610559; Bethyl (Montgomery, TX; Ran, A304-298A-M; p-KAP1(Ser-824), A300-767A; ATM, A300-136A), Genetex (Irvine, CA; RCC1, GTX63356; MCM2, GTX6238; cyclin B1, GTX6139; p-ATM (Ser-1981), GTX61739), Novus Biologicals (Littleton, CO; 53BP1, NB100-304), Santa Cruz Biotechnology (Dallas, TX; RanGAP1, sc-1862; cyclin D1, sc-753; GAPDH, sc-22233; Rad51, sc-8349; RanBP2, sc-74518; 53BP1, sc-22760; Chk1, sc-8408), Abcam (Cambridge, MA; RCC1, ab109379; TPX2 ab23785), Cell Signaling Technology (Danvers, MA; RanBP1, 8780; Chk2, 3440), Epitomics ( Burlingame, CA; pS101H, 1173-1), Developmental Studies Hybridoma Bank (Iowa City, IA; tubulin E7, E7), Millipore (γH2AX(Ser-193), 05-636), and GenScript (Piscataway, NJ; β-actin, A00702).

IF
Cells were grown on cleaned 12-mm round coverslips or on 32-mm ibidi chambers for RNAi-treated cells. For most experiments, cells were prepared for IF as previously described (Hasegawa et al., 2013), using antibodies to γH2AX (05-636; Millipore), 53BP1 (sc-22760; Santa Cruz Biotechnology), and p-KAP1 (A300-767A; Bethyl). For the detection of 53BP1 recruitment to γH2AX (Figure 5B), cells were fixed in 4% paraformaldehyde and 1% Triton X-100 in PBS for 10 min at room temperature.

53BP1 nuclear foci counting
The cells stained for 53BP1 and γH2AX were photographed on a wide-field Olympus IX81 microscope (Olympus USA, Waltham, MA) equipped with a UplanSApo 60×/1.35 oil-immersion lens and a Hamamatsu C4742 charge-coupled device (CCD) camera. For each sample, we acquired images of the DNA stain (Hoechst-33342), 53BP1 and γH2AX in at least 85 cells in randomly selected fields. Using the γH2AX foci as a control for a DNA damage–specific signal, the number of 53BP1 foci detected in each nucleus was manually counted and recorded in Excel spreadsheets by at least two independent observers.

IF quantification
MetaMorph 7.0.5 (Molecular Devices, Sunnyvale, CA) was used to quantify the IF signal intensity in images acquired using the UplanSApo 60×/1.35 oil-immersion lens and the Hamamatsu C4742 CCD camera. IF staining was performed as described above. Before quantification, the image-specific average out of cell fluorescence was subtracted from all images. For quantification of the cellular distribution of 53BP1, DNA images were thresholded for light objects to enable automatic creation of nuclear regions that were then transferred to 53BP1 images. After cytoplasmic 53BP1 regions for each cell were manually selected, region measurement data for each cell was logged into Excel, and the ratio of the average cytoplasmic and nuclear signal was calculated. Similarly, for Figure 5C, the DNA signal was used to create nuclear regions in the 53BP1 and γH2AX images, and Excel was used to calculate the ratios of average 53BP1 and γH2AX signal for each nucleus.

Immunoblotting quantification
To quantify protein expression on immunoblots, we used Odyssey, version 3 (Li-COR). The background-subtracted signal of individual protein bands was normalized to the GADPH or tubulin signal that was detected in the same lane.

Simple Western analysis
Simple Western (ProteinSimple, San Jose, CA) analyses were performed as previously described (Chen et al., 2013). Cell lysates (20 or 40 ng protein/capillary) were mixed with 1X SDS sample buffer, dithiothreitol, and fluorescent standards (ProteinSimple); heated at 70°C for 10 min before loading into the Peggy Sue instrument; separated by molecular weight; and immobilized using UV light. The capillaries were incubated with a blocking reagent (ProteinSimple), followed by primary and horseradish peroxidase–conjugated secondary antibodies (Jackson ImmunoResearch, West Grove, PA). A chemiluminescence signal was developed, captured by a CCD camera, and quantified as the peak area by the Compass software according to the ProteinSimple protocols. The primary antibodies used were Ran (610340; BD Biosciences), RCC1 (63356; GeneTex), cyclin D1 and cyclin B1 (sc-753 and sc-752; Santa Cruz Biotechnology), and vinculin (V9131; Sigma-Aldrich).

FACS analysis
Cells (500 μl of suspension in DPBS) were fixed with 4.5 ml ice-cold 70% ethanol, centrifuged (1200 rpm, 5 min, 4°C), washed with 5 ml ice-cold DPBS (1200 rpm, 5 min, 4°C), resuspended in 1 ml staining solution (20 μg/ml propidium iodide [Pi; Sigma-Aldrich], 200 μg/ml RNaseA [Sigma-Aldrich], 0.1% Triton X-100/ PBS), and incubated for 30 min at 37°C. After filtration with 70-μm nylon mesh, cells were sorted using FACS Calibur (Becton-Dickinson, Franklin Lakes, NJ). Cell debris and doublets were gated out, and G0/G1, S, and G2/M populations were quantified using the ModFit, version 3.0 (Becton-Dickinson). Images showing the total PI signal count versus count distribution were displayed using FlowJo-VX.

RCC1 and Ran RNAi
Control siRNA oligo (5′-UGGUUUAACUGUGACUA-3′; ONTARGET plus nontargeting siRNA #1), RCC1 siRNA SMARTpools (5′-GGAGAACCCUGUGGCUCUACCCG-AUGA-3′, 5′-GCACAGAACCUCGGCUUGGU-3′ and 5′-GGACUAACCGUGUGAUUU-3′; Chen et al., 2007), and Ran siRNA (5′-GAAUUUGGGACUGUGACAGA-3′ and 5′-CCACCAGAGAGCCUAAUA-3′) were obtained from Thermo-Fisher/Dharmacon. HC-T116 or hTERT-RPE1 cells were plated in six-well plates or 100-mm dishes in antibiotic-free medium and transfected with a 5 nM final siRNA concentration using Dharmafect (Thermo-Fisher).

FLIM imaging, analysis, and display
Spatially resolved, time-correlated single-photon counting (TCSPC) data sets were acquired as described previously (Kalab et al., 2006; Hasegawa et al., 2013), using a Zeiss LSM710 NLO (Zeiss USA, Thornwood, NY) microscope equipped with an SPC-830 (Becker & Hickl GmbH, Berlin, Germany) TCSPC controller. Imaging conditions were chosen to limit the emission to (0.2–1)×10· counts/s. Images of 128×128 pixels (1024 time bins/pixel) were acquired over 60–120 s and analyzed with SCI software (Becker & Hickl).
while assuming incomplete two-exponential decay and using trap-zoid integration and no prefixed parameters. We used SPCI to prepare the RGB pseudocolor images of the median \( t_{donor} \) values of RBP-4 detected in cells. The pseudocolor \( t_{donor} \) images were displayed within the range of 300 ps that was centered at cell-specific average \( t_{donor} \).

**Statistical analysis and image processing**

We analyzed data with Excel or GraphPad Prism, version 6.05 (GraphPad Software, La Jolla, CA). Images were prepared for publication with Adobe Photoshop and Illustrator CC (Adobe Systems, San Jose, CA).

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