Loss of pRB causes centromere dysfunction and chromosomal instability

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Chromosome instability (CIN) is a common feature of tumor cells. By monitoring chromosome segregation, we show that depletion of the retinoblastoma protein (pRB) causes rates of missegregation comparable with those seen in CIN tumor cells. The retinoblastoma tumor suppressor is frequently inactivated in human cancers and is best known for its regulation of the G1/S-phase transition. Recent studies have shown that pRB inactivation also slows mitotic progression and promotes aneuploidy, but reasons for these phenotypes are not well understood. Here we describe the underlying mitotic defects of pRB-deficient cells that cause chromosome missegregation. Analysis of mitotic cells reveals that pRB depletion compromises centromeric localization of CAP-D3/condensin II and chromosome cohesion, leading to an increase in intercentromeric distance and deformation of centromeric structure. These defects promote merotelic attachment, resulting in failure of chromosome congression and an increased propensity for lagging chromosomes following mitotic delay. While complete loss of centromere function or chromosome cohesion would have catastrophic consequences, these more moderate defects allow pRB-deficient cells to proliferate but undermine the fidelity of mitosis, leading to whole-chromosome gains and losses. These observations explain an important consequence of RB1 inactivation, and suggest that subtle defects in centromere function are a frequent source of merotely and CIN in cancer.

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High-throughput genomic profiling studies illustrate the fact that most human tumors are aneuploid and display abnormalities in the number of whole chromosomes or chromosome arms [Hanahan and Weinberg 2000; Albertson et al. 2003; Beroukhim et al. 2010]. Furthermore, many tumors have been shown to be chromosomally unstable [Lengauer et al. 1997]. Chromosome instability (CIN), defined as an elevated rate of gains and losses of whole chromosomes (10–100× more often than stable diploid cells) [Lengauer et al. 1997], has been proposed to promote the evolution of tumor cells. Such genomic changes potentially promote metastasis and chemotherapeutic resistance, and correlate with poorer patient prognosis [Nowell 1976; Kuukasjarvi et al. 1997; Rajagopalan and Lengauer 2004; Gao et al. 2007]. Recent studies show that aneuploidy and CIN can have a causal role in tumorigenesis and relapse [Rasnick and Duesberg 1999; Weaver et al. 2007; Baker et al. 2009, Sotillo et al. 2010].

CIN likely results from persistent defects in mitotic fidelity, and several mechanisms have been described that cause cells to missegregate whole chromosomes, including defects in bipolar spindle formation, chromosome–spindle association, chromosome cohesion, and the spindle assembly checkpoint (Cahill et al. 1998; Nigg 2002; Sotillo et al. 2007; Thompson and Compton 2008; Baker et al. 2009, Bakhoun et al. 2009; Ganem et al. 2009; Silkworth et al. 2009).

The retinoblastoma tumor susceptibility gene (RB1) is a key regulator of cell proliferation. RB1 was one of the first tumor suppressor genes to be identified, and its product (pRB) is functionally inactivated in most forms of cancer [Bookstein and Lee 1991; Marshall 1991; Weinberg 1995]. Although mutation of RB1 is a key, rate-limiting event in the development of most retinoblastoma, recent studies suggest that homozygous mutation of RB1 causes the appearance of benign retinoma, and these subsequently progress to retinoblastoma [Dimaras et al. 2008]. This malignant progression correlates with greatly increased levels of aneuploidy and genomic instability. The idea that mutation of RB1 may cause genomic instability is consistent with studies carried out using cultured cells. Populations of Rab−/− cells, or cells in which pRB is functionally inactivated, have elevated numbers of aneuploid cells, and are prone to increases in ploidy [Hernando et al. 2004; Iovino et al. 2006; Isaac et al. 2006; Mayhew et al. 2007; Srinivasan et al. 2007; Amato et al. 2009]. Together, these studies have lead to the hypothesis that much of
the aneuploidy seen in tumor cells is a by-product of the inactivation of the pRB pathway. Support for this idea comes from a completely independent line of investigation: Expression profiling has led to the identification of expression signatures that correlate with CIN. Remarkably, 10 of the 20 genes that show the highest correlation with CIN are genes that are known E2F targets, and are surrogate markers of pRB inactivation [Markey et al. 2002; Black et al. 2005; Carter et al. 2006].

Whether loss of pRB function leads to aneuploidy directly (through mechanisms that disrupt normal chromosome segregation) or indirectly (by allowing for the proliferation of spontaneously arising aneuploid cells) remains unclear. Evidence suggesting pRB loss may promote chromosome missegregation directly comes from a number of different observations; these include alterations in centrosome number [Iovino et al. 2006], defects in the spindle assembly checkpoint [Hernando et al. 2004], and formation of micronuclei [Amato et al. 2009], all of which are correlated with abnormal chromosome segregation. In addition, loss of pRB function alters the transcription of several genes with roles during mitosis [such as Aurora A, astrin, Cdc20, Mad2, Nek2, and Ndc80] [Iovino et al. 2006; Chakraborty et al. 2007], and several of these transcriptional changes likely have functional consequences. One potentially important target of pRB is Mad2 [Hernando et al. 2004]. Overexpression of Mad2 can cause aneuploidy, and can compromise the regulation of the spindle assembly checkpoint [Sotillo et al. 2007]. In contrast, it has also been reported Mad2 levels decrease, rather than increase, in pRB-depleted cells [Amato et al. 2009]. Currently, the link between pRB and aneuploidy is poorly understood because there is little data on how the mitotic machinery, which controls chromosome segregation, is altered when pRB is lost.

In this study, we show that the depletion of pRB from nontransformed, primary epithelial cells increases rates of chromosome missegregation to levels comparable with CIN tumor cells. This change occurs despite an intact spindle checkpoint and normal centrosome numbers. We show that the loss of pRB causes defects in centromeric condensation and sister chromatid cohesion, leading to centromere dysfunction during mitosis. The resulting aneuploid cells remain viable and capable of further cell division. Taken together, these observations explain why pRB mutant cells are prone to CIN.

Results

Rb depletion induces CIN

To determine the influence of pRB on chromosome segregation, we used RPE-1 cells, a well-characterized line of h-TERT immortalized human retinal pigment epithelial cells that has a stable, diploid karyotype and is frequently used for studies of mitosis. RPE-1 cells were infected with empty vector, or vector encoding shRNA specific for GFP or pRB [Morris et al. 2008]. An efficient and stable knock down of pRB was evident in cells treated with pRB-specific shRNA (sh-Rb) 6 d post-infection [Supplemental Fig. S1A,B]. To determine if cells depleted of pRB become aneuploid, we used fluorescence in situ hybridization (FISH) to monitor chromosome numbers. Centromere-specific probes for chromosomes 6 and 8 revealed a high degree of chromosome missegregation to levels comparable with pRB-depleted RPE-1 cells that persisted following several weeks of chronic depletion. Both control and sh-Rb populations exhibited a modal copy number of two for both chromosomes 6 and 8. (C) Analysis of individual segregation events revealed high rates of chromosome segregation errors in pRB-depleted cells (0.87% and 0.64% segregation error/division for chromosomes 6 and 8, respectively). (D) Following chronic depletion of pRB, aneuploid cells continued to proliferate (prometaphase cell) (top panel) and to missegregate chromosomes (late anaphase/telophase cell) (bottom panel). Chromosome 6 is shown in green, and chromosome 8 is shown in red throughout. Bar, 5 μm.

A

B

C

D

Figure 1. Loss of pRB induces CIN. (A,B) FISH analysis with probes for chromosomes 6 and 8 revealed a high degree of aneuploidy in populations of pRB-depleted RPE-1 (sh-Rb) cells that persisted following several weeks of chronic depletion. Both control and sh-Rb populations exhibited a modal copy number of two for both chromosomes 6 and 8. (C) Analysis of individual segregation events revealed high rates of chromosome segregation errors in pRB-depleted cells (0.87% and 0.64% segregation error/division for chromosomes 6 and 8, respectively). (D) Following chronic depletion of pRB, aneuploid cells continued to proliferate (prometaphase cell) (top panel) and to missegregate chromosomes (late anaphase/telophase cell) (bottom panel). Chromosome 6 is shown in green, and chromosome 8 is shown in red throughout. Bar, 5 μm.
The overduplication of centrosomes has been observed in cells in which the pRB pathway is targeted (Duensing et al. 2001; Lovino et al. 2006), and extra centrosomes have been shown to cause chromosome missegregation by promoting passage through a multipolar spindle intermediate (Lovino et al. 2006; Ganem et al. 2009; Silkworth et al. 2009). To determine whether this phenomenon might underlie chromosome missegregation in pRB-depleted RPE-1 cells, we scored the incidence of multipolar spindles as well as the number of centrioles in these cells. To avoid differences that result indirectly from changes in the mitotic index, we scored nondividing cells and mitotic cells separately. In nondividing cells, the distribution of centriole number was unchanged by pRB depletion [Supplemental Fig. S3A,B]. In addition, no increase in multipolar spindle formation and no mitotic cells containing more than two centrioles per spindle pole were observed in pRB-depleted cells [Supplemental Fig. S3C]. Nontransformed cells do not tolerate supernumerary centrosomes; consequently, extra centrioles are quickly lost from such cells [Ganem et al. 2009]. It is possible that pRB depletion might transiently induce the formation of supernumerary centrosomes. However, at the time during which chromosome segregation events are scored, extra centrosomes are not present and therefore are not the cause of the observed segregation errors.

Centromeric dysfunction in cells lacking pRB

Having excluded defects in the spindle assembly checkpoint and centrosome overduplication as possible sources of the chromosome segregation defects, we noted three clear phenotypes in cells depleted of pRB. Similar changes were seen in cells depleted of pRB by treatment with lentiviral shRNA constructs or with siRNA constructs that target different sequences [see the Materials and Methods].

First, pRB-depleted cells have a striking increase (>20%) in intercentromeric distance (1.24 ± 0.09 vs. 1.51 ± 0.07, P < 0.0001) [Fig. 2A,B]. This increase is somewhat dependent on the presence of microtubules, since treatments that depolymerize microtubules [hypotonic treatment or addition of nocodazole] partially alleviate this change [Fig. 2Bj data not shown]. Such treatments failed to restore the intercentromeric distance of pRB-depleted cells to that of control cells, suggesting that the loss of pRB causes a fundamental centromeric defect that is exaggerated by the forces associated with functional kinetochore–microtubule attachments.

Second, pRB-depleted cells exhibit a defect in chromosome congression, and chromosomes have difficulty aligning in a tight metaphase plate. Quantitation of this defect showed that the width of the metaphase plate increased 40% in pRB-depleted cells compared with that seen in control cells (7.06 ± 0.78 vs. 9.86 ± 2.31, P < 0.03) [Figs. 2C; Supplemental Fig. S4]. Live-cell imaging of RPE-1 cells expressing H2B-GFP [to allow visualization of chromosomes] or RFP-CENPA [to allow visualization of centromeres] supported fixed cell analysis and confirmed that chromosomes in cells depleted of pRB congress to
a disorganized metaphase plate prior to anaphase onset (data not shown). Lack of kinetochore microtubule attachment and chromosome biorientation could prevent proper chromosome congression. However, pRB-depleted cells form calcium-stable microtubules similarly to control cells, suggesting they are proficient to form stable kinetochore–microtubule interactions, and chromosomes are capable of biorientation [Supplemental Fig. S4B]. Recently, kinetochore congression and alignment within the metaphase plate have been shown to be influenced by centromere stiffness [Jaqaman et al. 2010], suggesting the idea that centromere structure may be compromised in pRB-depleted cells.

Third, a close analysis of sister kinetochore pairs revealed that, while properly bioriented chromosomes align with sister kinetochores oriented along the pole-to-pole axis, bioriented sister kinetochores in cells lacking pRB often deviate from this axis [Fig. 2D]. This phenotype has been reported for merotelically attached chromosomes where one or both sister kinetochores are associated with microtubules emanating from both spindle poles [Draviam et al. 2006]. Merotelic attachments are a common feature of CIN cells [Thompson and Compton 2008], and are not sensed by the spindle assembly checkpoint. If left uncorrected, these attachment errors give rise to lagging chromosomes during anaphase and missegregation events [Cimini et al. 2001, 2003]. Staining of sister kinetochore pairs with the anticientromeric antigen (ACA) labels sister kinetochores and the adjoining centromeric region in a barbell pattern. In pRB-depleted cells, the normal barbell pattern is dramatically bent at many kinetochore pairs [Fig. 2D, inset]. This type of centromeric distortion is reminiscent of the distortion seen with merotelic kinetochore attachments [Cimini et al. 2003; Draviam et al. 2006].

Taken together, these observations show that there is an underlying defect in centromeric structure in pRB-depleted cells in which decreased centromeric rigidity allows for improper kinetochore–microtubule attachments and defects in congression.

**Figure 2.** pRB loss causes centromeric dysfunction. (A) RPE-1 cells lacking pRB exhibited >20% increase in intercentromeric distance. This increase was not restored by inhibition of PLK1, but was partially corrected by microtubule depolymerization. (C) Chromosome alignment was also impaired, and the width of the metaphase plate was increased ~40% in cells lacking pRB. Error bars represent standard error (SEM) of three independent experiments. (D) Centromeres were distorted when pRB was depleted, such that the barbell configuration [staining the kinetochores and associated centromeric region] seen with ACA staining in control cells [insets in top panels] was often bent [insets in bottom panels] in cells lacking pRB. Kinetochore staining was shown in red, microtubules are shown in green, and centrioles are shown in yellow [A] or blue [D]. Similar results were obtained with siRNAs or shRNAs, the data shown were obtained from shRNA-treated cells. Bars, 10 μm. Insets are 4× enlargements.

**pRB loss influences sister chromatid cohesion**

The depletion of cohesin complex components has been shown to cause defects in centromeric structure and function that are similar to the properties of pRB-depleted cells [Kenney and Heald 2006; Jaqaman et al. 2010]. To determine whether the chromosomal abnormalities seen in pRB-depleted cells might also lead to defects in chromosome cohesion, we examined metaphase chromosome spreads. Cells treated with shRNA specific for pRB were found to have an increased incidence of premature loss of sister chromatid cohesion [Fig. 3A,D]. Furthermore, arms of sister chromatids that maintained cohesion in pRB-depleted cells were found to be further apart than in control cells [Fig. 3B]. Strikingly, both of these defects were exacerbated by prolonged mitotic arrest [Fig. 3A [bottom panel], C]. These results suggest that chromosome cohesion is impaired in pRB-depleted cells, and that sister chromatids are prone to separate in the absence of pRB, particularly when cells are delayed in mitosis.
To test the idea that the centromeric defects of pRB-deficient cells can compromise mitotic fidelity, we examined progression through mitosis following release from a mitotic arrest. Completion of mitosis following release from nocodazole-induced microtubule depolymerization promotes transit through a multipolar spindle intermediate, which results in formation of syntelic and merotelic kinetochore attachments (Cimini et al. 2001). Because merotelic attachments are not sensed by the spindle assembly checkpoint, cells may enter anaphase without correcting such erroneous attachments, as evidenced by lagging chromosomes (Cimini et al. 2001). In this sensitized system, cells lacking pRB exhibit a delay in the formation of a bipolar spindle, a delay in anaphase progression, an increased number of anaphase cells exhibiting lagging chromosomes, and an increased number of laggards per cell [Fig. 4].

**Figure 3.** pRB loss disrupts sister chromatid cohesion. (A) Cells depleted of pRB (shRb) have an increased incidence of premature loss of sister chromatid cohesion (examples of separated sister are indicated by asterisks [*]). (B) Line scans through arms of sisters that maintain cohesion revealed an increase in interchromosomal distance in response to pRB loss. Each colored line represents a line scan through the arms of paired sister chromatids, with each chart depicting five line scans within a single representative cell. Prolonged mitotic arrest increased the interchromosomal distance (C) (error bars represent standard error [SEM]; P < 0.05 between control and sh-Rb cells at both 3 and 20 h) and promoted the complete loss of cohesion (D), as measured by the number of cells with one to three or four or more unpaired chromosomes.

**Centromeric defects promote chromosome malattachment and segregation errors**

As described above, the depletion of pRB promotes premature loss of sister chromatid cohesion, and this defect is enhanced by prolonged mitotic delay [Fig. 3]. We therefore expected that individual unpaired kinetochores should be apparent by immunofluorescence following release from nocodazole arrest. Indeed, single kinetochores located near spindle poles [Supplemental Fig. S5A] were evident in pRB-deficient cells but not control cells following nocodazole washout. Under these experimental conditions, spindle formation occurs following drug washout, after chromatid cohesion has been lost between some sister pairs, and lone chromatids are unable to efficiently biorient and align at the spindle equator. However, such lone kinetochores were not routinely observed near spindle poles in control or pRB-depleted cells that had not been subjected to prolonged mitotic arrest. We infer that the reduced level of sister chromatid cohesion seen in pRB-depleted cells is sufficient for chromosome bi-orientation. Although sister chromatids are more likely to separate when pRB-depleted cells are delayed in mitosis,
weakened sister chromatid cohesion also promotes merotelic attachment, and this may allow single chromosomes to maintain alignment that is achieved prior to loss of cohesion (Supplemental Fig. S5B,C). The delays in mitotic progression and the increase in lagging chromosomes following release from a nocodazole arrest indicate that pRB-depleted cells are especially sensitive to conditions that promote merotelic attachments.

pRB loss disrupts centromeric cohesin and condensin II localization

In vertebrate cells, components of both the cohesin and condensin complexes are important for normal centromeric structure and function (Ono et al. 2004; Kenney and Heald 2006; Samoshkin et al. 2009). Cohesin and condensin complexes first associate with DNA during telophase (for review, see Peters et al. 2008). Cohesin is the primary determinant of sister chromatid cohesion, and cohesion is established during S phase concurrent with DNA replication. As cells progress through mitosis, cohesin complexes are removed in two steps. During the prophase-to-metaphase transition, cohesin is first removed from chromosome arms by phosphorylation of cohesin subunits. Subsequently, during the metaphase-to-anaphase transition, cohesin is removed from the centromeric region by cleavage of the Rad21/Scc1 subunit (Peters et al. 2008).

The overall levels of the cohesin proteins Rad21 and SMC3 were unchanged in pRB-depleted cells, and the levels of chromatin-associated Rad21 and SMC3 were only slightly reduced when pRB was depleted (Fig. 5A; Supplemental Fig. S6A). This is consistent with the observation that the degree of chromatid cohesion in these cells is sufficient to maintain cohesion upon mitotic entry. However, centromeric-localized cohesin represents only ~10% of all cohesin complexes (Peters et al. 2008), and centromere-specific changes in cohesin localization

Figure 4. pRB loss compromises the fidelity of mitotic progression. Formation of a bipolar spindle (A,D), and progression into anaphase (B) following release from nocodazole-induced mitotic arrest is delayed when pRB is depleted (shRb). (A) The state of spindle organization is characterized as disorganized microtubule arrays (dark blue), multipolar spindles (light blue), and bipolar spindles (yellow). Numbers represent percentage of mitotic cells exhibiting specified spindle structure. P < 0.01 for differences in spindle structure and anaphase progression at time points 40 and 60 min between control and shRb samples. (C) pRB-depleted cells that enter anaphase exhibit an increase in the incidence of lagging chromosomes over control cells. (D) Examples of representative spindle structure at specified time points following nocodazole washout. Microtubules are shown in green, and DNA is shown in blue. Bar, 10 μm.
may cause only a minor change in the overall pool of chromatin-associated protein. Using immunofluorescence, we observed that the punctate pattern of Rad21 seen in cells that have undergone nuclear envelope breakdown was specifically reduced in pRB-depleted cells (Fig. 5B). Polo-like kinase 1 (PLK1) promotes the removal of cohesin during the transition to metaphase (Lenart et al. 2007). Since PLK1 is an E2F1 target gene and its expression has been shown to increase in the absence of pRB (Amato et al. 2009), we tested whether inhibition of PLK1 could rescue the effects of pRB depletion. Addition of the PLK1 inhibitor BI 2536 (Chemie Tek) induced a strong mitotic arrest and prevented loss of sister chromatid cohesion in both control and pRB-depleted cells. However, PLK1 inhibition did not suppress the increased intercentromeric distance seen in pRB-depleted cells (Fig. 2C), suggesting that this defect is not due to an inappropriate PLK1-dependent removal of cohesin. Because centromeric cohesin is removed by Separase-dependent cleavage of the Rad21 subunit, we compared the levels of cleaved Rad21 in control and pRB-depleted RPE-1 cells. No increase in Rad21 cleavage was observed in pRB-depleted cells (Fig. 5C), suggesting that the reduction in centromeric cohesin is more likely due to a defect in the loading or maintenance of cohesin at the centromere rather than to excessive Rad21 cleavage.

To test whether the link between pRB and sister chromatid cohesion might be conserved during evolution, we compared neuroblasts of wild-type Drosophila larvae with larvae that are strongly hypomorphic for the pRB homolog RBF1. As in pRB-depleted human cells, rbf1120 mutant animals exhibit defects in sister chromatid cohesion and an increase in aneuploidy (Fig. 6A–C). Consistent with the idea that pRB family members promote loading of the cohesin complex onto DNA, immunostaining experiments show reduced dRad21 banding on the polytene chromosomes of rbf1120/D14 mutant larvae (Fig. 6D). Interestingly, defects in sister chromatid cohesion have also been observed in mouse embryonic fibroblasts lacking the three Rb family proteins (TKO MEFs) (van Harn et al. 2010).

Our previous studies using Drosophila showed that RBF1 interacts physically with the condensin II subunit dCAP-D3 and promotes its association with chromatin (Longworth et al. 2008). This physical interaction is
conserved in mammalian cells, and requires the LXCXE-binding cleft of pRB; however, the significance of this interaction in mammalian cells was unclear. Consistent with the published data, fractionation experiments showed that the level of chromatin-associated CAP-D3 (condensin II) was reduced in pRB-depleted cells (Figs. 5A, 7A; Supplemental Fig. S6C). In comparison, the level of chromatin-associated CAP-D2 (condensin I) was unaffected by pRB depletion [Fig. 5A]. Intriguingly, condensin II complexes have been shown to be enriched in the centromeric region of mitotic chromosomes [Ono et al. 2004]. We therefore tested whether pRB might be required specifically for normal centromeric recruitment of CAP-D3, and whether the loss of CAP-D3 could cause defects in centromere function similar to pRB-depleted cells.

Immunofluorescence experiments confirmed that CAP-D3 is strongly enriched at the centromere of control RPE-1 cells. This localization was strikingly reduced in cells treated with pRB-specific siRNA, but not cells depleted of Rad21 (Fig. 7B,C). To assess the functional significance of reduced CAP-D3, RPE-1 cells were treated with siRNAs specific for pRB, CAP-D2, or CAP-D3, and examined for centromeric defects. pRB- and CAP-D3-depleted cells showed a similar increase in intercentromeric distance that was not evident in control-treated cells, nor in cells specifically depleted of CAP-D2 [Fig. 7C,D]. Previous studies have shown that the depletion of CAP-D3 enhances the loss of sister chromatid cohesion in a similar manner to that reported here in pRB-depleted cells [Hirota et al. 2004]. Taken together, our results show that pRB-depleted cells have a defect in chromosome cohesion, that this defect is most evident at the centromere, and that the defects in centromere function appear to be caused by a reduction in the binding and/or regulation of condensin II and cohesin complexes.

Discussion

The loss of pRB causes a consistent, high rate of chromosome segregation errors, leading to the generation of aneuploid, chromosomally unstable cells. The results described here reveal why this occurs: The loss of pRB compromises centromere function and cohesion, and these changes undermine the fidelity of mitosis.

*pRB, CIN, and tumorigenesis*

Some cancer cells have a near-tetraploid karyotype, suggesting that missegregation of single chromosomes occurs in conjunction with, or is facilitated by, doubling of the genome (through cell fusion, endoreduplication, or failure of cytokinesis, etc.). However, most human tumor cells are near-diploid [Weaver and Cleveland 2006], and seem unlikely to have progressed through a tetraploid intermediate. We show that loss of pRB results in near-diploid aneuploid cells, and that the rate of chromosome missegregation resulting from the inactivation of pRB is remarkably similar to that measured in human tumor cells. Given that the pRB pathway is disrupted in most tumor cells, the idea that pRB inactivation causes CIN provides a simple explanation for the prevalence of chromosome missegregation in human cancer. pRB inactivation may be especially relevant during tumorigenesis because its loss both increases segregation errors and
compromises cell cycle arrest. Previous studies have focused on the role of pRB in G1/S-phase progression, but understanding the basis for chromosome segregation defects in pRB-deficient cells is equally important: This information gives insight into the processes and pathways that are misregulated in human tumors. Moreover, the characterization of these defects may reveal ways to restore chromosome stability, thereby reducing metastatic potential. Conversely, treatments that can further exaggerate these changes may cause mitotic catastrophe and eventual death in cells that lack pRB function.

Rb loss: a subtle defect that undermines the fidelity of mitosis

Here we show that pRB loss causes defects in centromere function and reduces sister chromatid cohesion. These changes promote merotelic attachment and increase the incidence of unpaired sister chromatids. While the complete loss of cohesion would have catastrophic consequences, moderate defects in centromeric condensation and cohesion have been shown to cause chromosome segregation errors in tissue culture and various model systems (for review, see Nasmyth and Haering 2009). Hence, the functional inactivation of pRB is a subtle enemy during tumorigenesis because it reduces the fidelity of mitosis without causing more dramatic changes that would compromise cell proliferation.

pRB is necessary for the proper action of condensin II and cohesin components at the centromere. Condensins and cohesins are both required for normal centromere structure and function. Failure to recruit these components leads to a decrease in rigidity between sister centromeres and promotes merotelic kinetochore attachment—defects that are evident in pRB-depleted cells in the increased intercentromeric distance, deformation of centromeric structure, failure of chromosome congression, and increased propensity for lagging chromosomes following mitotic arrest. This combination of centromeric cohesion defects and merotelic kinetochore orientation is highly consistent with studies showing the importance of pericentromeric cohesion in promoting kinetochore biorientation (Ng et al. 2009). Many studies have shown that merotelic attachments are a dominant cause of chromosome missegregation in CIN tumor cell lines (Cimini et al. 2001; Thompson and Compton 2008; Ganem et al. 2009). Given the frequency of lesions in the pRB pathway, and the evidence that pRB loss promotes merotelic attachments, it is likely that much of the CIN in tumors is initiated by functional inactivation of pRB. This underscores an important caveat to the widespread use of human tumor cells in the study of normal mitotic progression and chromosome segregation.

The link between pRB and chromosome segregation

We suggest that the mitotic phenotype of pRB-deficient cells may have multiple components. First, as illustrated here, are the changes resulting from the physical connection between pRB and condensin II. pRB is required for the concentration of CAP-D3 at the centromere, and the depletion of either pRB or CAP-D3 gives similar centromeric defects. These observations are consistent with evidence that condensin II proteins are necessary for centromere function (Hirota et al. 2004; Samoshkin et al. 2009), that pRB associates physically with CAP-D3, and that pRB can promote the association of CAP-D3 with chromatin as cells exit mitosis and enter G1 (Longworth et al. 2008). Taken together, these results suggest that the interaction between pRB and CAP-D3 is important for normal centromere function.

Second, the centromeric defects of pRB-depleted cells are associated with reduced cohesion. Since condensation and cohesion are interrelated processes, it is possible that the cohesion defects are secondary to reduced condensin II activity. Indeed, the depletion of CAP-D3 has been shown to enhance the loss of sister chromatid cohesion in a manner similar to that reported here (Hirota et al. 2004). Additionally, we note that the tight association of cohesin with DNA and the establishment of sister chromatid cohesion are intimately linked to the initiation of DNA replication (Terret et al. 2009), for review, see Nasmyth and Haering 2009). The loss of pRB not only alters the expression of replication proteins, but also changes the spatial organization of DNA synthesis within the nucleus (Barbie et al. 2004), a change that may also impact the loading of cohesin and the maintenance of condensin II complexes. The changes described here add to the emerging view that defects in chromosome cohesion are important in human disease. Mutations in cohesin components and mutations in regulators of cohesin association with DNA have been identified in various human disorders, including Cornelia de Lange syndrome (CdLS) and Roberts syndrome (RBS) (Mannini et al. 2009), as well as colorectal cancer (Barber et al. 2008). Interestingly, many of the mutations identified in human disease do not alter cohesin protein levels, but instead affect the loading of the cohesin complex or change its affinity for DNA (Mannini et al. 2009; Revenkova et al. 2009).

Third, it is important to remember that the centromere defects resulting from the loss of pRB do not occur in a normal cell, but in a cell in which the deregulation of E2F has caused numerous mitotic proteins to be misexpressed. Unlike the change seen when pRB or CAP-D3 are depleted, the overexpression of E2F1 did not increase the intercentromeric distance, even though it increased the expression of several E2F-regulated genes (Supplemental Fig. S6F–I). This suggests that changes seen in centromere structure and function are not directly caused by elevated E2F activity. However, we note that the deregulation of E2F may enhance the importance of these structural changes. Several groups have shown that the length of mitosis increases in pRB-deficient cells (Hernando et al. 2004; Amato et al. 2009). This mitotic delay has been suggested to be due to an E2F-dependent up-regulation of Mad2 and hyperactivity of the spindle assembly checkpoint (Hernando et al. 2004). Although the centromeric defects described are not apparently caused by the deregulation of E2F, and therefore are unlikely to be closely connected to the up-regulation of Mad2, the finding that pRB-deficient cells have defects in cohesion that are...
exaggerated by prolonged mitosis may help to explain why such a delay can have important consequences. One could argue that the inactivation of pRB has two syner-
gistic effects: a condensation defect that weakens sister
chromatid cohesion and promotes merotelic attachment,
and a mitotic delay that enhances these defects.

In our previous studies, we found that all three pRB
family members are able to interact with CAP-D3
[Longworth et al. 2008]. The up-regulation of p107 and
p130 has been shown to allow them to compensate for the
absence of pRB, although these effects are often tissue-
or context-specific [Pacal and Bremner 2006; Burkhart
and Sage 2008]. We know that the depletion of pRB causes
centromeric defects and aneuploidy in nontransformed
cells as diverse as RPE1 epithelial cells and IMR90 fibro-
basts [data not shown], however, it remains to be deter-
dined whether p107 and/or p130 also influence cen-
tromere function and the fidelity of chromosome segregation,
and whether there are specific tissues or contexts in which
these family members impact the degree of chromosome
missegregation resulting from the inactivation of pRB.

Loss of pRB promotes genomic instability
Genomic instability has been implicated recently in the
development of retinoblastoma [Dimaras et al. 2008], and
the results reported here outline one way in which the
inactivation of pRB promotes genetic change. Consistent
with these observations, Coschi et al. [2010] found that
mutation of the LXXCE cleft, which reduces the ability of
pRB to interact with CAP-D3, enhances tumorigenesis and
genomic instability in mouse tumor models. It is tem-
pering to speculate that the elevated levels of DNA damage
observed in pRB-deficient cells [Pickering and Kowalik
2006] and in pRB, p107, p130 TKO cells [van Harn et al.
2010] may further increase the likelihood that cells will
termitosis with damaged DNA and activate mitotic
checkpoints [Mikhailov et al. 2002]. While further studies
are clearly needed to characterize the full extent of the
changes resulting from reduced condensin II and cohesin
function in cells lacking pRB, work presented here de-
fines an important and clinically relevant mechanistic
role for pRB in the maintenance of genome stability.

Materials and methods
Cell culture and RNAi
hTERT-RPE-1 cells were grown in Dulbecco’s Modified Essential
Medium (DMEM) supplemented with 10% fetal bovine serum
(FBS) and 1% penicillin/streptomycin. Cells were infected with
LNP empty vector, LNP-shGFP, or LNP-shRb[GD] (targeting se-
quence GGUGAAAAUGGCAAUACGG) as described previ-
ously [Morris et al. 2008], placed under puromycin selection for
4–5 d, and analyzed at day 6 or later as specified. For all ex-
periments involving the shRNA depletion of pRB (as indicated
as shRb), cells infected with empty vector and cells infected with
a shGFP-containing vector were both used as controls.

For experiments in which the short-term depletion of targets
was achieved using siRNAs, cells were transfected with RNAi
MAX [Invitrogen] according to the manufacturer’s directions,
with 50 nM pool of four pRB-specific siRNA constructs [Dhar-
macon Smartpool; target sequence 1, CGAAAUCCAGUCCA
UAAG; target sequence 2, GAGUGUACCUGAUGAGAU; tar-
get sequence 3, AAACUACCUGUAGUAUG; target sequence 4,
GAAUCGUCCUGCUCUA], a CAPD2-specific siRNA construct
(CCAUACUGUGCUAGUACCA) [Hirota et al. 2004],
a CAPD3-specific siRNA construct (CAUGGACGUAGGAGA
GU) [Hirota et al. 2004], or a Rad21-specific siRNA construct
(GGUGAAUAAGCUAUAGCG) [Watrin et al. 2006]. mRNA
samples were collected and cells were fixed and stained for
analysis as described below, 36–40 h after siRNA transfection.

For consistency, in the experiments where cohesin and conden-
sin components were depleted by siRNA, we also used siRNAs
to deplete pRB (indicated as siRb in the text). As described for the
shRNA treatments, the treatment with siRb strongly decreased
the level of pRB mRNA and the level of pRB protein detected
by immunofluorescence. As expected, the depletion of pRB
increased the levels of pRB/E2F-regulated mRNAs p107 and
SMC2 [Supplemental Fig. S4A]. To test the effects of E2F1
overexpression, RPE-1 cells were transfected with HA-E2F1
and examined 36 h later. Real-time quantitative PCR was performed
to determine relative mRNA levels as published previously
[Morris et al. 2008].

Drosophila stocks
All fly stocks and crosses were maintained at 25°C. Female
progeny of the Rbf1[120a];CA14° and male progeny of the Rbf1[120a]
stocks were analyzed for mitotic defects, chromosome structure,
and aneuploidy.

FISH, chromosome spreads, drug treatment, and cell
cycle arrest
Cells were prepared and fixed, and α-satellite-specific probes
for chromosomes 6 and 8 (Cytocell) were hybridized according
to protocols in Thompson and Compton [2008]. Chromosome
spreads were prepared as described in Ganem et al. [2009]. Cells
were treated with 100 ng/mL nocodazole or 100 ng/mL colcemid
for 5 h for karyotype analysis, or for indicated times for analysis
of sister chromatid cohesion. To measure interchromosomal
distances, line scans to measure pixel intensity were performed
on five chromosomes in each of 10 cells per condition. Centriole
numbers were counted in nonproliferating cells that were grown
to confluency. The lack of proliferation was confirmed by the
appearance of confluency and the lack of mitotic cells, as deter-
mined by phase-contrast imaging prior to fixation, and by mi-
crotubule and chromatin structure following fixation. For noco-
dazole washout experiments, cells were treated with 100 ng/mL
nocodazole for 16 h, then washed three times in drug-free me-
dium and fixed as described below at the indicated time points.
Mitotic cells were staged and spindle structure was analyzed
according to DAPI and α-tubulin staining. More than 90 mitotic
cells were scored for spindle structure and mitotic progression
for each condition and at each time point in three independent
experiments. Dissected third instar larval Drosophila neu-
roblasts were treated in 1 mM colchicine for 2 h prior to fixation
in 4% paraformaldehyde for 30 min. Brains were then incubated
sequentially in ice-cold methanol for 2 min and 45% acetic acid
for 3 min, squashed between slide and coverslip in 60% acetic
acid, and frozen in liquid nitrogen. Chromosome numbers were
scored for at least 100 mitotic cells for each of three neuroblasts
of w1118 and rbf1 mutant animals.

Immunofluorescence microscopy
Cells were extracted in either microtubule-stabilizing buffer
[4 M glycerol, 100 mM PIPES at pH 6.9, 1 mM EGTA, 5 mM
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Dharmacon Smartpool; target sequence 1, CGAAAUCCAGUCCA
UAAG; target sequence 2, GAGUGUACCUGAUGAGAU; target
sequence 3, AAACUACCUGUAGUAUG; target sequence 4,
GAAUCGUCCUGCUCUA], a CAPD2-specific siRNA construct
(CCAUACUGUGCUAGUACCA) [Hirota et al. 2004],
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GU) [Hirota et al. 2004], or a Rad21-specific siRNA construct
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and examined 36 h later. Real-time quantitative PCR was performed
to determine relative mRNA levels as published previously
[Morris et al. 2008].
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MgCl₂, 0.5% Triton X-100) or calcium-containing buffer (100 mM PIPES at pH 6.8, 1 mM MgCl₂, 0.1% Triton X-100, 1 mM CaCl₂), followed by fixation in 1% glutaraldehyde [microtubule staining: dim.1a [Sigma] or cold methanol [ACA Antibodies, Inc.], BubR1 [Abcam], CAPD2 [Bethyl Laboratories], CAPD3 [Bethyl Laboratories], CENP-A [Cell Signaling], Centrin-2 [Santa Cruz Bio-technology], Rad21 [Abcam], and pRB [Cell Signaling and BD Pharmingen]]. Alternatively, cells were fixed in cold methanol without additional pre-extraction. Subsequent antibody incubations and washes were done in TBS-BSA [10 mM Tris at pH 7.5, 150 mM NaCl, 1% bovine serum albumin]. DNA was detected with 0.2 μg/mL DAPI (Sigma-Aldrich). Coverslips were mounted with ProLong Antifade mounting medium [Molecular Probes]. Fluorescent images of fixed and live cells were captured with a Hamamatsu Orca AG cooled CCD camera mounted on a Nikon Ti/Yokagawa CSU-10 spinning-disk confocal microscope with a 100×, 1.4 NA objective or a Hamamatsu EM CCD camera mounted on an Olympus IX81 microscope with a 100×, 1.4 NA objective. A series of 0.25-μm optical sections were collected in the Z-axis for each channel [DAPI, fluorescein, Cy5, and/or Texas red]. Iterative restoration was performed on images using NIS elements for images acquired on the Nikon. Selected planes from the Z-series were then overlaid to generate the final image. Mitotic index and mitotic progression were quantified by immunofluorescence using chromatin compaction (DAPI stain) and spindle structure (α-tubulin) as indicators of mitotic state. For calculation of the mitotic index, n > 5000 cells per condition were used. For evaluation of mitotic progression, n > 300 mitotic cells were examined per condition. Measurements of intercentromere distances and metaphase plate width were made with Slidebook analysis software; line scans measuring interchromome distances and metaphase plate width were made with NIS elements for images acquired on the Nikon. Selected planes from the Z-series were then overlaid to generate the final image. Mitotic index and mitotic progression were quantified by immunofluorescence using chromatin compaction (DAPI stain) and spindle structure (α-tubulin) as indicators of mitotic state. For calculation of the mitotic index, n > 5000 cells per condition were used. For evaluation of mitotic progression, n > 300 mitotic cells were examined per condition. Measurements of intercentromere distances and metaphase plate width were made with Slidebook analysis software; line scans measuring interchromosomal distances were acquired with NIS elements software. Measurements of chromatin-bound or centromere-localized cohesin and condensin components by immunofluorescence were made with Slidebook software by selecting nuclei based on DAPI staining or kinetochores/centromeres based on ACA staining, measuring pixel intensities in >50 selected areas for each condition, and normalizing all intensities to average intensities in control cells. All measurements were performed for three independent experiments. All error bars represent standard errors (SEM). The Student’s t-test was used to calculate the significance of differences between samples.

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Loss of pRB causes centromere dysfunction and chromosomal instability

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