Telomere shortening alters the kinetics of the DNA damage response after ionizing radiation in human cells

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Abstract

Studies of telomerase-deficient mice and human cell lines have demonstrated that telomere shortening enhances sensitivity to ionizing radiation (IR). The molecular basis for this observation remains unclear. To better understand the connection between telomere shortening and radiation sensitivity, we evaluated components of the DNA damage response pathway in normal human fibroblasts with short and long telomeres. Late-passage cells with short telomeres showed enhanced sensitivity to IR compared to early-passage cells with longer telomeres. Compared to early-passage cells, late-passage cells had a higher baseline level of phosphorylated H2AX protein ($\gamma$H2AX) before IR, but diminished peak levels of H2AX phosphorylation after IR. Both the appearance and disappearance of $\gamma$H2AX foci were delayed in late-passage cells, indicative of delayed DNA repair. In contrast to the situation with H2AX, ATM and p53 phosphorylation kinetics were similar in early and late-passage cells, but phosphorylation of the chromatin-bound ATM targets SMC1 and NBS1 was delayed in late-passage cells. Because impaired phosphorylation associated with short telomeres was restricted to chromatin-bound ATM targets, chromatin structure was assessed. DNA from cells with short telomeres was more resistant to digestion with micrococcal nuclease, indicative of compacted chromatin. Moreover, cells with short telomeres showed histone acetylation and methylation profiles consistent with heterochromatin. Together our data suggest a model in which short telomeres induce chromatin structure changes that limit access of activated ATM to its downstream targets on the chromatin, thereby providing a potential explanation for the increased radiation sensitivity seen with telomere shortening.
Introduction

There has been growing recognition of a close relationship between telomeres and the DNA damage response (1). Proteins involved in the DNA damage response localize to telomeres and are required for normal telomere maintenance, yet are also involved in the cellular response to telomere shortening and dysfunction (2). Cells with short or dysfunctional telomeres activate a damage response similar to the response observed with DNA double-strand breaks, including activation of ATM, phosphorylation of ATM targets, and formation of nuclear foci containing protein complexes involved in DNA repair (3-5). The telomere-induced damage response arises, in part, from direct uncapping of telomeres and is not solely attributed to end-to-end fusions of dysfunctional telomeres leading to chromosome breakage during anaphase.

The link between telomeres and the DNA damage response has been furthered by several studies, both at cellular and organismal levels, demonstrating that telomere shortening and dysfunction are determinants of radiation sensitivity (6-11). However, the molecular basis underlying this sensitivity remains unclear. Understanding how telomere integrity influences sensitivity to DNA damaging agents is especially relevant given the emergence of telomerase inhibition as a therapeutic modality for human cancers. In the present study, we compared the cellular response to ionizing radiation in isogenic human cells with short and long telomeres. Our data show that the kinetics of the DNA damage response is altered in cells with short telomeres and suggest that telomere shortening is associated with chromatin structure changes that limit access of activated ATM to its chromatin-bound downstream targets.
Materials and Methods:

Cell Culture

The ATCC CRL-2091 normal primary human foreskin fibroblast strain (HFF) and the HeLa human cervical carcinoma cell line were obtained from the American Type Culture Collection (Manassas, VA). 293T cells were a gift from Dr. David Baltimore; no authentication of 293T cells was done by the authors. Cells were propagated in DMEM (Biowhittaker) supplemented with 10% fetal bovine serum (HyClone), L-Glutamine, and Penicillin/Streptomycin (Gibco, Carlsbad, CA) at 37 °C and 5% CO₂ in a humidified incubator. Fibroblasts were trypsinized and counted every 7 days and reseeded at a density of 5 x 10⁵ cells per 75 cm² flask. HeLa cells were counted every 3-4 days and replated at a density of 0.5-1 X 10⁶ cells per 100 mm dish. The histone deacetylase (HDAC) inhibitor vorinostat (LC laboratories, Woburn, MA) was dissolved in DMSO, and applied to cells at a concentration of 2 µM. Cells were treated with vorinostat for 24 hours prior to irradiation.

Generation of TERT constructs and retroviruses

Plasmids containing human TERT cDNA (pCI-neo-hEST2) were a gift from Dr. Robert Weinberg (Whitehead Institute for Biomedical Research, Cambridge, MA). Catalytically inactive TERT-DN (TERT-dominant negative) was generated by substituting the Aspartic acid residues with Valine residues at positions 868 and 869 by site-directed mutagenesis (Quick Change Site-directed Mutagenesis kit, Stratagene, La Jolla, CA). The TERT and TERT-DN mutants were sequenced completely and subcloned into the retroviral expression vector MSCV-IGFP, which expresses the green
fluorescent protein (GFP) reporter from an internal ribosomal entry site (IRES). Vesicular stomatitis virus-G (VSV-G) pseudotyped retroviruses were packaged in 293T cells transfected with the plasmids pEQ-PAM3-E (containing the helper proteins), VSV-G (containing the envelope), and MSCV-IGFP, MSCV-IGFP-TERT or MSCV-IGFP-TERT-DN. Virus-containing medium was applied to the cells of interest in three separate aliquots with polybrene, as previously described (12). Forty-eight hours after transduction, GFP-containing cells were isolated by fluorescence activated cell sorting (FACS). The experiments described were conducted on HFFs taken in the early passages after transduction with the hTERT retrovirus when a sufficient number of cells were available after FACS sorting (estimated to be several population doublings). HeLa cells transduced with the mutant TERT-DN constructs were cloned by the limiting dilution method. The process of generating visible HeLa cell colonies required 20 population doublings.

**Telomerase Activity and Telomere Length Assays**

Cell extracts were prepared according to protocols provided by the manufacturers. Telomerase activity was measured using the TRAPeze telomerase detection kit (Intergen, Purchase, NY). For telomere length analysis, 5 µg genomic DNA was digested with *Hinf* I and *Rsa* I and analyzed by southern blotting using 32P labeled (TTAGGG)6 probe. Signals were visualized using a PhosphorImager (Storm 860, Molecular Dynamics).

**Clonogenic survival assays**

Cells were seeded onto six-well plates in concentrations predicted to yield 100-200 colonies per well. Increasing cell concentrations were used for increasing doses of
ionizing radiation (IR). For the IR experiments, cells were subjected to the indicated doses of IR 20 hrs after seeding. Two weeks after irradiation, the cells were fixed and stained with 0.01% formaldehyde and 0.1% crystal violet. Colonies containing at least 50 cells were counted.

**Immunoprecipitation, Western Blotting and Immunofluorescence**

All procedures are described in Bakkenist, Drissi *et al.*(3) and in Kitagawa *et al.*(13). Briefly, cells extracts were prepared in RIPA buffer (200 mM NaCl, 10 mM Tris at pH 7.5, 0.1% SDS, 1% NP-40, 0.5% Deoxycholate, 1X protease inhibitor cocktail tablet from Roche), anti-SMC1 (BL308, Bethyl Laboratories, INC), anti-SMC1S957-P, anti-p53 (Ab-6, Oncogene Research Products), anti-p53S15-P (Cell Signaling Technology), anti-beta actin (Novus Biologicals), anti-acetyl-Histone H3K9 (Upstate), anti-dimethyl-Histone H3K9 (Upstate), anti-H3, CT, pan, clone A3S (Upstate), anti γH2AX (Millipore), anti-NBS1 (Novus Biologicals), anti-NBS1S343-P (Signalway Antibody).

**Histones Extraction**

Extracts containing soluble histone proteins were performed as described previously (13). In brief, cell pellets were resuspended in hypotonic buffer (10 mM HEPES at pH 7.9, 1.5 mM MgCl₂ and 10 mM KCl). Cell were then lysed in a hypertonic lysis buffer (10 mM HEPES at pH 7.9, 1.5 mM MgCl₂, 250 mM NaCl, 10 mM KCl, 0.2 M HCl, 0.5 mM DTT, 1.5 mM PMSF and 1X protease inhibitor cocktail tablet from Roche). Cell extracts were dialyzed against 0.1 M acetic acid twice for 1 h and three times against H₂O for 1h, 3 h and overnight respectively.
Micrococcal Nuclease Assay

Cells were permeabilized for 90 s with 0.01% egg-Lysolecithin (Sigma) diluted in 150 mM sucrose, 80 mM KCl, 35 mM HEPES, pH 7.4, 5 mM K2HPO4, 5 mM MgCl2 and 0.5 mM CaCl2. Cells were then digested with 2 U/ml micrococcal nuclease (Sigma) in 20 mM sucrose, 50 mM Tris-HCl, pH 7.5, 50 mM NaCl and 2 mM CaCl2 at room temperature for various time points. DNA was isolated and subjected to 6% polyacrylamide gel electrophoresis to analyze the digestion products.

Results

Cells with short telomeres are more sensitive to ionizing radiation (IR)

To evaluate the sensitivity of cells with short telomeres to IR, we used clonogenic survival assays at different population doublings of primary human foreskin fibroblasts (HFF). We previously had shown that telomeres shorten in HFF at each population doubling (3). Late-passage fibroblasts with short telomeres had significantly enhanced sensitivity to IR compared to early-passage fibroblasts (Fig. 1). To confirm that IR sensitivity is associated with short telomeres, we also introduced a catalytically inactive dominant-negative hTERT construct (hTERT-DN) to inhibit telomerase activity in HeLa cells. Transduced cells were cloned by the limiting dilution method. IR sensitivity was assessed at different population doublings using clonogenic survival assays. The introduction of hTERT-DN into HeLa cells resulted in complete inhibition of detectable telomerase activity (Fig. S1A) and telomere shortening with successive population
doublings (PD) (Fig. S1B). Cells with short telomeres were markedly more sensitive to IR than their counterparts with longer telomeres (Fig. S1C).

**Kinetics of γH2AX nuclear focus formation and resolution are delayed in cells with short telomeres**

Cellular survival following DNA damage depends on the integrity of the signaling pathways to repair DNA double-strand breaks (DSBs) and restore genomic integrity. H2AX phosphorylation is an early step in the response to DNA damage. Upon IR, H2AX is rapidly phosphorylated by activated ATM and DNA-PK on serine 139 to generate γH2AX, a robust marker of DSBs (15-16) that can be used to monitor DSBs repair (17). Both activated ATM and γH2AX form nuclear foci upon DSBs. The extent of DNA damage post-IR and the kinetics of γH2AX foci formation and resolution were evaluated in late and early-passage HFF by immunofluorescence using γH2AX-S139 phospho-specific antibodies. Late-passage cells were pre-senescent, as evidenced by continued growth and lack of senescence-associated β-galactosidase staining (supplemental Figure S2). Although the cells were pre-senescent, they were growing at a slower rate than early passage cells. To control for differences in cell growth, most of the experiments outlined below were conducted on contact-inhibited cells that were 3 weeks past the point of confluence.

As expected, at baseline before subjecting cells to IR, late-passage (PD63) HFF cultures had a higher percentage of cells with γH2AX foci compared to early-passage cultures (PD11). However, upon IR, a higher peak percentage of early-passage HFF manifested
γH2AX foci compared late-passage HFF (Fig. 2A). Moreover, the kinetics of γH2AX focus appearance and disappearance was faster in early-passage compared to late-passage HFF (Fig. 2A, 2B). Similar results were obtained using western blot analysis to assess γH2AX protein levels (Fig. 2C). 30 min after 1 Gy of IR, early-passage HFF showed a robust induction of H2AX phosphorylation, while late passage HFF did not increase γH2AX protein levels until 60 minutes after IR (Fig 2C, lane 3 and 4). At 5 hours post-IR, while γH2AX protein levels were still elevated in late-passage HFF, γH2AX protein levels returned to near baseline levels in early-passage HFF (Fig 2C, lane 9 and 10). γH2AX foci disappeared at later time point in late-passage cells, 16 hours post-IR, (Fig 2C, lane 15 and 16).

**Effect of ectopic hTERT expression on γH2AX focus formation and resolution**

To confirm that the altered kinetics of DNA repair observed in late-passage HFF was telomere dependent, we transduced HFF at late passage (PD 64) with a retroviral vector that expresses hTERT. Transduced cells exhibited telomerase activity and elongated telomeres in early passages after the cells were harvested after the transduction (Figure S3). Ectopic telomerase expression in late-passage cells completely restored the kinetics of γH2AX focus formation and resolution to the levels observed with early-passage cells (Fig. 3A), demonstrating that telomere dysfunction, rather than other effects of prolonged cell culture such as oxidative damage, was responsible for the altered H2AX phosphorylation Similar results were obtained using western blot analysis to assess γH2AX protein levels (Fig. 3B).
ATM, p53, SMC1 and NBS1 phosphorylation in early and late-passage HFF

To further characterize the delayed DNA repair kinetics observed in late-passage HFF, we assessed other members of the DNA damage response pathway. ATM is rapidly activated in response to DSBs and phosphorylates key proteins leading to cell cycle arrest, senescence and apoptosis (18). ATM substrates include p53, NBS1, H2AX, 53BP1, BRCA1, SMC1 and Chk2 (19-20). Because late-passage cells showed delayed H2AX phosphorylation and γH2AX focus formation, we surmised that ATM activation may also be delayed. To test this hypothesis, the kinetics of ATM activation (measured by ATM-S1981P) after IR was evaluated in early and late-passage HFF that were contact inhibited for several weeks. Similar to the observation with H2AX, late-passage HFF had a higher baseline level of ATM activation compared to early-passage cells. However, in contrast to the situation with H2AX phosphorylation, there was no striking difference in the level of ATM activation between early and late-passage HFF in response to IR (Fig. 4A). Moreover, the kinetics of ATM-S1981P nuclear focus formation was similar in early and late-passage cells (Fig. 4B). We also evaluated the phosphorylation of p53, SMC1 and NBS1 in early and late-passage HFF. The accumulation of p53 protein and its phosphorylation on S15 occurred with the same kinetics after IR in early and late-passage cells (Fig. 4C). However, the phosphorylation of SMC1 on S957 and NBS1 on S343 was delayed in late-passage cells (Fig. 4C and 4D). Together these results suggest that late-passage cells exhibit a delay in phosphorylation of ATM target proteins bound to chromatin, like H2AX, SMC1 and NBS1. Proteins that are not chromatin bound, such as p53 and ATM itself, have similar levels of activation in response to IR in late and early-passage cells.
Changes in Chromatin structure is associated with short telomeres

The delay in phosphorylation of H2AX, SMC1 and NBS1 might be induced by chromatin structure change, triggered by short telomeres, resulting in limited access of activated ATM to its chromatin-bound downstream targets. To test this hypothesis, we used the micrococcal nuclease assay to analyze chromatin structure in late and early-passage cells. Consistent with the resistance of compacted chromatin to limited nuclease digestion (21), at an early time point (1 min), chromatin from late-passage pre-senescent cells was more resistant to micrococcal nuclease digestion compared to early-passage cells, suggesting that late-passage chromatin is compacted (Fig 5A and 5B). The resistance to micrococcal nuclease was completely abolished in late-passage cells expressing hTERT, linking this resistance to telomere dysfunction.

Chromatin from late-passage cells has features of heterochromatin

To determine whether late-passage compacted chromatin is related to heterochromatin, we examined H3K9 acetylation and methylation levels. Consistent with the features seen with compact chromatin (22), our results indicated reduced levels of H3 acetyl K9 and increased levels of H3 dimethyl K9 in late-passage cells. (Fig 6, lane 1 and 3). Ectopic expression of hTERT in late-passage cells reversed H3K9 acetylation and methylation profiles, suggesting that these histone modifications are caused by short telomeres (Fig. 5, lane 4). Moreover, this reversal required an active hTERT on telomeres, as the ectopic expression of hTERT-DN or hTERT-HA (a form of hTERT that is catalytically active
but unable to elongate telomeres and prevent senescence) (23) did not reverse these profiles (Fig. 6, lane 5 and 6).

To further assess whether chromatin state was responsible for the diminished DNA damage response in late-passage cells, we treated late-passage HFF with the histone deacetylase (HDAC) inhibitor vorinostat, which promotes histone acetylation and an open chromatin state. Treatment with the HDAC inhibitor restored the intensity and rapidity of H2AX phosphorylation in late-passage cells (Figure 6B). Together, these results suggest a model in which short telomeres induce chromatin structure change consistent with heterochromatin, causing chromatin compaction and limited access of ATM kinase to its chromatin targets, thereby potentially contributing to enhanced radiosensitivity in late-passage cells.

**Discussion**

In the present study we evaluated the mechanism of enhanced radiosensitivity in cells with short telomeres. As expected, primary and transformed cells with short telomeres had markedly enhanced sensitivity to IR compared to cells with longer telomeres. These results mirror the results observed in *Terc* knock-out mice, which showed that progressive telomere shortening was associated with enhanced cellular and organismal sensitivity to ionizing radiation (7, 11). Similar results have been observed in human cells (24-27). Although some studies have suggested that telomerase expression *per se* influences radiosensitivity (8, 28), our results and the results of other groups suggest that telomere length is the primary determinant of radiosensitivity (7, 11, 14, 25, 29-30).
these studies, ectopic telomerase expression or telomerase inhibition had no effect on sensitivity to IR until a measurable change in telomere length was observed.

The biological mechanism underlying the relationship between telomere integrity and radiation sensitivity remains to be determined. A simple model to explain this correlation is that cells with short telomeres possess a baseline level of DNA damage from uncapped telomeres even before they are subjected to IR. In this model, the pre-existing damage would be additive with the extra damage induced by IR, thereby enhancing radiation sensitivity. Arguing against this model is our observation that the baseline number of $\gamma$H2AX foci in late-passage fibroblasts is equivalent to an ionizing radiation dose of only 0.1 Gy (data not shown). Adding 0.1 Gy to the levels of ionizing radiation that were used in the clonogenic survival assays would be insufficient to explain the enhanced sensitivity to IR that we observed, though the protracted 0.1 Gy-equivalent exposure in late-passage cells may be biologically different from the short-lived exposure associated with a one-time dose of IR. Although the baseline number of $\gamma$H2AX foci in late-passage cells was elevated compared to early-passage HFF, the late-passage cells had diminished H2AX phosphorylation in response to IR at early time points (Figure 2). If the enhanced radiation sensitivity in late-passage cells were merely the result of an additive effect of radiation damage to telomere damage, we would have expected the peak levels of H2AX phosphorylation to be higher in the late-passage cells than the early passage cells. A second model that has been proposed is that telomeres serve as repositories of proteins that participate in the DNA damage response, which can readily translocate to DNA breaks (11). According to this premise, once telomeres become short, repair proteins can
no longer bind telomeres and are no longer immediately available to sites of DNA damage. However, the DSB repair proteins that associate with telomeres, such as the MRE11/RAD50/NBS1 (MRN) complex, are uniformly distributed in the nuclei of unirradiated cells (31). Because the telomere-bound component comprises just a small fraction of the abundant pool of repair proteins, it seems unlikely that a decrease in telomere length would have a major effect on the availability of repair proteins to be recruited to DNA breaks.

The present data support a third model for increased radiation sensitivity in cells with short telomeres. One of the earliest events in the DNA damage response is ATM phosphorylation/activation and swift phosphorylation of H2AX (γH2AX) on Ser139, primarily by activated ATM and DNA-PK (16, 19). Upon DSBs, phosphorylated H2AX (γH2AX) rapidly forms DNA repair foci that include the MRN complex, 53BP1 and BRCA1 (32). Our results showed that late-passage cells had attenuated peak levels of H2AX phosphorylation and that the kinetics of H2AX phosphorylation and dephosphorylation were delayed. Because γH2AX plays a crucial role in the recruitment of proteins to the repair focus and in its stabilization to recruit late factors like cohesins to tether sister chromatids for homologous recombination repair pathway (33-36), the altered H2AX phosphorylation kinetics may account for the enhanced radiosensitivity in late-passage cells. Recently, it was shown that telomerase-null and telomerase-inhibited mouse embryo fibroblasts (MEFs) formed fewer γH2AX foci and had delayed appearance and disappearance of such foci following heat shock and IR exposure compared to telomerase positive MEFs (6). Although these results focused on telomerase
expression and not telomere length, they are concordant with our results. Furthermore, delayed kinetics of DNA DSB processing, including the rates of recruitment of DSB repair proteins to γH2AX foci, has been observed in normal and pathological aging (9).

Our data suggest a mechanism for the altered kinetics of γH2AX focus formation and disappearance in late-passage cells. Late-passage chromatin was resistant to micrococcal nuclease digestion and had reduced levels of H3K9 acetylation and higher levels H3K9 dimethylation, indicative of heterochromatinization. These results are consistent with the observation that senescent cells accumulate foci of heterochromatin (37), though the late-passage fibroblasts employed in our experiments were pre-senescent (Figure S2). The compacted chromatin could explain why phosphorylation and activation of chromatin-bound DNA damage proteins (H2AX, SMC1, and NBS1) were delayed in late-passage cells, while activation of non-chromatin-bound proteins (p53 and ATM) was unchanged.

Moreover, we observed that dephosphorylation of γH2AX was delayed in late-passage cells. Dephosphorylation of γH2AX is achieved by protein phosphatase 2A (38), by the HTP-C phosphatase complex in budding-yeast (39) and by Wip1 phosphatase (40-42). Like delayed phosphorylation of chromatin-bound proteins, delayed dephosphorylation may be caused by reduced access of phosphatases to late-passage chromatin.

Furthermore, we and others have shown that H3K9Ac is a DNA-damage-responsive modification (Zhang and Drissi, unpublished data; (43), suggesting this histone modification is a mark of heterochromatin and DNA damage.

Telomerase protein has also been described to regulate chromatin state and the DNA damage response (28). Suppression of hTERT in human fibroblasts led to enhanced
radiosensitivity and delayed kinetics of $\gamma$H2AX focus formation, without measurable changes in telomere length. Abrogation of hTERT was associated with a pattern of euchromatin (enhanced sensitivity to micrococcal nuclease-induced DNA digestion, increased H3K9 acetylation, and decreased H3K9 dimethylation). It is interesting that although the end-result of telomerase suppression (enhanced radiosensitivity and delayed DNA repair kinetics) was similar to the effect we observed with telomere shortening, telomerase suppression increased euchromatin content whereas telomere shortening increased heterochromatin content. These results indicate that although telomerase repression and telomere shortening are closely intertwined, the two processes do not have identical consequences for the cell.

It remains to be determined how telomere length influences chromatin structure and, in turn, how chromatin structure influences the DNA repair process. It has recently been shown that ATM-mediated phosphorylation of heterochromatin-promoting proteins such as KAP-1 increases the repair of DNA breaks within heterochromatin (44). Our data showed that ATM activation kinetics in late-passage cells were not affected by short telomeres, but that phosphorylation of H2AX, SMC1, and NBS1 was delayed. It is possible that telomere shortening-induced heterochromatin introduces an impediment to ATM-dependent chromatin modifications that facilitate DNA double-strand break repair.

Finally, our results support several predictions regarding the use of telomerase inhibitors as therapeutic agents for human cancer. First, telomerase inhibition in itself can induce apoptosis and growth arrest in cancer cells, though this response may be delayed, as
observed in our HeLa cells. Second, the concurrent administration of telomerase inhibitors and ionizing radiation may have a synergistic effect once sufficient telomere shortening has occurred. The concurrent application of DNA-damaging agents and telomerase inhibitors may provide a therapeutic benefit.
References


Figure Legends

Fig. 1. Cells with short telomeres are more sensitive to IR. Early (PD 15) and late (PD 69) passage HFF were irradiated at the indicated IR doses. The surviving fraction of cells was determined using a clonogenic survival assay.

Fig. 2. Kinetics of $\gamma$H2AX focus formation and resolution are delayed in cells with short telomeres. A, Early (PD 11) and late-passage (PD 63) HFF were irradiated with 0.4 Gy and processed for immunofluorescence using anti-$\gamma$H2AX antibodies at the indicated time post-IR. A total of 100 cells were analyzed, cells with 3 or more foci were considered $\gamma$H2AX-positive. B, time course of $\gamma$H2AX resolution, the percentage of cells with unresolved $\gamma$H2AX IR-induced foci was assessed in early and late-passage at the indicated time following irradiation with 0.4 Gy. C, Early and late-passage HFF were irradiated with 1 Gy and harvested at the indicated time points post-IR for histone protein extraction. Phosphorylated H2AX protein levels were assessed using anti-$\gamma$H2AX antibodies.

Fig. 3. Delayed kinetics of $\gamma$H2AX foci formation and resolution is suppressed by ectopic hTERT expression. A, Early, late and late-passage HFF transduced with a hTERT-expressing retrovirus vector were irradiated with 0.4 Gy and analyzed by immunofluorescence with anti-$\gamma$H2AX at the indicated time points post-IR. Cells were considered positive if 3 or more foci were present. B, Kinetics of H2AX phosphorylation
after 1 Gy IR was assessed by immunoblot with anti-γH2AX in early, late, and late-passage cells transduced with the hTERT retrovirus.

Fig. 4. Delayed phosphorylation kinetics of chromatin bound proteins. A, ATM phosphorylation kinetics post-IR is similar in early and late-passage HFF. HFF at PD 20 and 65 were irradiated with 0.2 Gy. Cells were harvested at the indicated time points post-IR and whole-cell extracts were prepared. Cleared supernatants were immunoprecipitated with anti-ATM, and the samples were resolved by gel electrophoresis and immunoblotted with anti-total ATM and anti-ATM 1981S-P. B, ATM1981S-P foci formation kinetics are similar in early and late-passage HFF. HFF at PD 11 and 63 were irradiated with 0.5 Gy and the kinetics of ATMS1981-P foci formation were assessed at the indicated time points post-IR. A total of 100 cells per time point were analyzed, cells with 3 or more foci were considered ATMS1981-P positive. The values shown are the mean +/- S.D. of 5 fields analyzed of 100 cells each. C and D, HFF at PD 16 and 58 were irradiated with 10 Gy and harvested at the indicated time points for immunoblotting. Total and phosphorylated p53 SMC1 and NBS1 were assessed.

Fig. 5. Compaction of chromatin structure in late-passage cells is reversed by hTERT ectopic expression. A, Resistance of late-passage cells to micrococcal nuclease digestion. Micrococcal nuclease digestion of permeabilized HFF cells at PD 14, PD 60 and PD 60 transduced with retrovirus expressing hTERT. DNA was isolated from cells after micrococcal digestion for the indicated time in minutes and resolved by
polyacrylamide gel electrophoresis. B, represent quantification of the DNA amounts in A entering the gel relative to undigested DNA at time 0 minute. The values shown are the mean +/- S.D. of three independent experiments.

**Fig. 6. Reduced H3K9 acetylation and increased methylation levels in cells with short telomeres.** A, HFF at the indicated PD numbers were processed for acid cellular extraction of histone proteins (see materials and methods), lysates were analyzed by immunoblotting with anti-acetyl-Histone H3K9, anti-dimethyl-Histone H3K9 and anti-H3. HFF at PD 14 were transduced with retrovirus expressing hTERT HA-tagged at C-terminal and passaged to PD 41 after transduction. HA-hTERT was shown to have *in vitro* telomerase enzymatic activity but did not maintain telomeres (23). HFF at PD 14 were transduced with retrovirus expressing a dominant negative of hTERT (hTERT-DN) and passaged to PD 35. B, Cells were treated with 2 µM of the histone deacetylase inhibitor vorinostat prior to IR. After 1 Gy IR, H2AX protein was extracted and assessed for phosphorylation by immunoblotting with anti-γH2AX.
Early and late-passage primary fibroblasts

Drissi, Fig. 1

% Survival vs. Gray
Drissi, Fig. 3

A

![Bar graph showing cells with three or more γH2AX foci over time after IR.]

- PD 64 + hTERT
- PD 18
- PD 64

Time Point (Hours) After IR:
- 0
- 1
- 2
- 4
- 6
- 8

B

<table>
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<th>hr after 1 Gy</th>
<th>PD 66</th>
<th>PD 13</th>
<th>PD 66 + hTERT</th>
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<td>no-IR</td>
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<td>no-IR</td>
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Western Blot Analysis:
- γH2AX
- Tubulin
Figure 4

A. IP: Total ATM

Min. post IR
0.2 Gy

HFF PD 20
HFF PD 65

Anti-ATM-S1981P

Anti-ATM

WB

B. Percent of ATM-S1981-P foci in contact inhibited HFF

Hours after IR at 0.5 Gy

C. PD 16

PD 58

min. post IR
IR

- + +
- + +

Anti-SMC1S957-P

Anti-SMC1

Anti-p53S15-P

Anti-p53

Anti-β Actin

D. PD 16

PD 58

min. post IR
IR

30 60
30 60

Anti-NBS1S343P

Anti-NBS1

Anti-Actin