c-Myc Accelerates S-Phase and Requires WRN to Avoid Replication Stress

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Abstract

c-Myc interacts with components of the pre-replication complex and directly regulates DNA replication [1]. However, the consequences of this novel c-Myc function on cell cycle dynamics and replication-associated damage are unknown. Here, we show that c-Myc overexpression in primary human fibroblasts markedly accelerates S-phase while c-Myc deficient fibroblasts exhibit a prolonged S-phase. We also show that the Werner DNA helicase protein (WRN) plays a critical role in supporting c-Myc-driven S-phase, as depletion of WRN in c-Myc overexpressing cells increases DNA damage specifically at sites of DNA synthesis. This excess DNA damage activates a “replication stress” pathway involving ATR, CHK1, CHK2, and p53, leading to rapid senescence of WRN deficient c-Myc overexpressing cells. Indeed, depletion of p53 rescues this senescence response. We propose that WRN functions to repair abnormal replication structures caused by the acceleration of DNA replication by c-Myc. This work provides an additional mechanistic explanation for c-Myc-induced DNA damage and senescence, and reveals a vulnerability of c-Myc overexpressing cells that could potentially be exploited in cancer therapy.

Introduction

The profound effects of c-Myc on cell growth, proliferation, apoptosis, and tumorigenesis have been mainly attributed to its ability to coordinate gene transcription. c-Myc is a basic helix-loop helix transcription factor that directly modulates transcription of a large number of genes controlled by all three RNA polymerases [2,3,4]. c-Myc can also indirectly regulate transcription through up-regulation of histone acetylases, with the potential to globally influence chromatin structure [5]. In addition, previous studies have demonstrated that c-Myc overexpression triggers the generation of double strand breaks and thus promotes chromosomal instability [6,7]. Several mechanisms have been proposed to explain this phenomenon, including overriding checkpoints [8]. c-Myc interference with DNA repair mechanisms [9] and increased production of reactive oxygen species, (ROS) [10]. However, a recent study has identified a novel function for c-Myc in the direct stimulation of DNA replication, independent of its transcriptional targets [1], thus providing yet another potential mechanism for c-Myc induced genomic instability. This new c-Myc function was demonstrated by its ability to bind to components of the pre-replication complex, both in mammalian as well as in Xenopus cells, and to influence the rate of DNA replication in Xenopus cell-free extracts [1,11]. In addition, c-Myc binds to known mammalian replication origins [1]. We hypothesized that overexpression of c-Myc might accelerate S-phase, leading to increased sensitivity to replication stress. As c-Myc-expressing cells show impaired proliferation and rapid senescence in the absence of the Werner DNA helicase, we further hypothesize that WRN is required to minimize replication stress during c-Myc driven S-phase.

The WRN gene encodes for a RecQ-DNA helicase, which has been shown to resolve DNA structures that form during S-phase, such as those generated at stalled of replication forks [12,13]. Loss of function mutations in WRN cause progeroid features in humans, a condition known as Werner Syndrome (WS) [14]. Fibroblasts isolated from WS patients exhibit genomic instability, increased sensitivity to specific DNA damaging agents, slow proliferation, lengthened S-phase, and accelerated replicative senescence [15,16]. These phenotypes have been in part recapitulated by acute knock-down of WRN protein by RNAi [17,12]. hTERT overexpression can immortalize WS fibroblasts, although it does not rescue their slow growth, or the S-phase defects ([18] and CG unpublished observations). These findings are consistent with a role for the WRN helicase in maintaining genome integrity during DNA replication, in addition to its described role in telomere maintenance and stability [19,20].

Here we monitored the effect of c-Myc expression on the rate of S-phase in human foreskin fibroblasts (HFF) and in c-myc deficient rat cells. Our experiments demonstrate that c-Myc overexpression markedly accelerates S-phase and conversely, c-myc deficient cells exhibit a greatly prolonged S-phase. Furthermore, WRN plays a critical role in this phenotype, as depleting WRN in c-Myc overexpressing cells leads to an increase in DNA damage, and cellular senescence. These results provide a mechanism for c-Myc-induced senescence and identify the DNA replication-repair machinery as a potential therapeutic target for c-Myc overexpressing tumor cells.

Results and Discussion

To examine the effect of c-Myc upon cell cycle dynamics we utilized primary human foreskin fibroblasts (HFF), as these cells...
tolerate c-Myc overexpression using a retroviral vector (pB-puro), and exhibit c-Myc-specific phenotypes, but do not develop the senescence response that is seen in other cell types [21]. HFF-Myc cells exhibit faster growth and increased size and prominent nucleoli, increased RNA synthesis, and a global gene expression signature typical of c-Myc overexpressing cells [3] and CG unpublished. To examine the duration of S-phase, HFF-pB and HFF-Myc cells were synchronized at the G1/S boundary by double thymidine block. Upon release from G1/S, cells were labeled for one hour with BrdU and then chased for the indicated times and subjected to FACS analysis (Fig. 1). The progression of BrdU/propidium iodide-stained cells through S-phase into G2 as shown by the accumulation of cells with a G2 DNA content, indicated that HFF-pB cells completed S-phase within five to six hours, as expected for the duration of S-phase of mammalian cells (Fig. 1). In contrast, HFF-Myc cells showed a greatly reduced duration of S-phase, completing S-phase within three to four hours (Fig. 1). This indicates that c-Myc overexpression accelerates the duration of S-phase, by about 40% in normal human fibroblasts. To determine if the loss of c-Myc slowed the duration of S-phase, we examined the c-myc−/− TGR rat cell line in comparison with the c-myc wild type TGR parental cell line [22]. After release from the G1/S block, the TGR parental line reached G2/M after eleven hours, at which time c-myc−/− cells were still in the middle of S-phase (Fig. 1b). An additional time course showed that the c-myc−/− cells completed S-phase at approximately fourteen hours from the time of release (data not shown), thus extending S-phase by three hours. In sharp contrast, the c-myc-reconstituted knockout TGR cells, (c-myc−/−;+pB-Myc) cells had an accelerated S-phase completed at six hours (Fig. 1b). Thus, loss of c-Myc extended S-phase, and this delay was not just rescued by c-Myc overexpression, but in fact was accelerated nearly two fold relative to the wild type cells.

These data are consistent with the results obtained in HFFs, although they differ in two ways. First, a substantial fraction of c-myc reconstituted TGR line did not arrest at G1/S after thymidine block (see Fig. 1b, time 0 hour). Second, the parental TGR cells exhibited a prolonged arrest after thymidine release, leading to an apparently longer S-phase then the HFFs (eleven hours vs. six hours, Fig. 1a and b). Nevertheless, this experiment indicates that the levels of c-Myc can dramatically influence not only entry into S-phase as previously described [22], but also the duration of S-phase. These results, together with the previous observation that c-Myc overexpressing cells show an increased number of replication foci [1], suggest that c-Myc may increase the number of active replication origins, thus reducing replication timing.

As WRN participates in the repair of abnormal DNA replication structures we next asked if WRN function precipitates DNA damage during S-phase. We thus compared γ-H2AX foci; a marker of DNA damage, in WRN deficient cells that have been immortalized by h-tERT overexpression, h-tERT-WRN+/− versus normal h-tERT immortalized human fibroblasts, h-tERT-WRN+/+ [24]. γ-H2AX foci coinciding with sites of newly replicated DNA were quantified by labeling cells with a short pulse of BrdU prior to fixation. The co-localization of BrdU and γ-H2AX was also compared in h-tERT-WRN+/− fibroblasts versus h-tERT-WRN−/− fibroblasts after c-Myc overexpression with the c-Myc retroviral vector, LXSN-Myc. c-Myc expression in h-tERT-WRN−/− cells led to a 3.5-fold increase in γ-H2AX foci co-localizing with BrdU, relative to h-tERT-WRN−/− cells with empty vector (Fig. 3a and b), indicating c-Myc induced DNA damage occurs at sites of DNA replication. In contrast, only a 1.5-fold increase in BrdU/γ-H2AX foci was caused by LXSN-Myc in h-tERT-WRN+/− control fibroblasts (Fig. 3a and b), indicating that WRN loss allowed c-Myc-induced DNA damage to accumulate, presumably due to lack of repair. The number of γ-H2AX/BrdU foci is likely an underestimate of S-phase DNA damage, as cells were labeled with BrdU for only 30 minutes to capture DNA-damage associated with newly replicated DNA. Thus, BrdU negative γ-H2AX foci could also represent damage incurred during S-phase. BrdU labeling was only detectable in cells undergoing S-phase and not in thymidine arrested or quiescent cells, thus excluding the possibility of labeling replication-associated repair (see Fig. 2a, 30 min after thymidine release and data not shown). These results indicated that c-Myc-induced DNA damage occurred during S-phase and was aggravated in WRN deficient cells. To determine if WRN protein in normal HFFs also caused replication-associated damage, two WRN specific siRNAs were expressed from retroviral vectors (siWRN I and II; Fig. 3c). WRN knockdown synergized with c-Myc overexpression to aggravate DNA-replication associated damage as shown by increased γ-H2AX/ BrdU foci (Fig. 3c and quantitated in Fig. 3d). These results confirm that both acute and chronic WRN deficiency exacerbates the accumulation of DNA damage at replication sites under conditions of c-Myc overexpression.

The finding of DNA replication associated damage in c-Myc expressing WRN deficient cells suggested that intra-S checkpoints
A first sensor of replication-associated damage might be triggered. A first sensor of replication-associated damage is the single stranded DNA binding protein complex, RPA [27]. The RPA complex, which consists of three subunits, RPA70, 32 and 14, coats excess single strand DNA generated by aberrant or collapsed replication forks. Indeed, transduction of WRN$^{-/-}$ cells with LXSN-Myc caused a striking increase in RPA nuclear speckles compared with c-Myc expressing WRN$^{+/+}$ cells or relative to WRN$^{-/-}$ cells transduced with an empty retroviral

Figure 1. c-Myc levels determine the rate of S-phase. a) c-Myc accelerates S-phase in human foreskin fibroblasts (HFFs). BrdU pulse-chase experiment in HFFs transduced with pB-Myc retrovirus or empty vector, pBabe. Cells were synchronized by thymidine block and released in BrdU for one hour, then chased in BrdU-free media for the indicated time. Progression of early labeled BrdU cells through S and G2 was monitored by FACS analysis. The BrdU positive HFF-Myc cells accumulated in G2 between three to four hours after release, while the control cells HFF+M strain accumulated in G2 beginning at five hours. Propidium iodide staining at the time of release, prior to BrdU addition to verify the G1/S arrest of HFF-Myc cells is shown in supplementary Figure S1. b) Knockout of c-myc lengthens S-phase. Propidium iodide profiles of the rat cell line TGR, its derivatives c-myc$^{-/-}$ [22] and the reconstituted c-myc$^{-/-}$;+pB-Myc upon release from thymidine block is shown. While TGR cells completed S-phase between eight and eleven hours, c-myc$^{-/-}$ cells were still in S-phase at eleven hours. Reconstituted c-myc$^{-/-}$;+pB-Myc cells showed a dramatic acceleration of S-phase (six hrs). c) Expression of c-Myc was analyzed by western blot. Actin is shown as loading control.

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Figure 2. Increased presence of WRN at replication sites in c-Myc overexpressing HFFs. HFF-pB (a) and HFF-Myc (b) were synchronized by thymidine block, labeled at the indicated times for 30 minutes with BrdU, and fixed for staining. Cells were stained for WRN (green) and BrdU (red), and DNA was stained with Hoechst (blue). Images were collected with a 100X oil-immersion objective. An increased co-localization of WRN with BrdU foci (yellow) is detectable at the 1.5 and 2.5 hr time point in HFF-Myc cells. This indicates recruitment of WRN at sites of newly replicated DNA caused by c-Myc overexpression. A 3-D movie reconstructed from this experiment can be seen as supplemental data (Supplementary Movies S1 and S2). c) Higher magnification of HFF-pB (top) and HFF-Myc (bottom) cells at 1–1.5 hr time point to better visualize WRN and BrdU co-localized foci. d), e) Percentage of cells in early, intermediate, and late S-phase determined by the BrdU patterns. HFF-Myc (e) progressed faster than HFF-pB (d) and only “late” patterns were present at 3.5 hrs post-release, similar to the result obtained by the pulse-chase experiment (Fig. 1).
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vector (Fig. 4a). RPA staining was increased in virtually every cell undergoing the first cellular division and thus likely during the first or second S-phase after c-Myc overexpression. However, RPA staining was not induced above background by LXSN-Myc transduction in fibroblasts expressing a functional WRN protein (Fig. 4a).

RPA is thought to signal activation of ATR and thereby trigger an intra-S checkpoint [27]. This leads to the tight association of ATR with chromatin surrounding the sites of damage [28]. Here we employed anti-phospho-ATR (Ser 428) specific antibody to stain the Triton-resistant nuclear fraction of cells, as an alternative to biochemical cell-fractionation techniques. Indeed this method

**Figure 3. DNA damage accumulates at replication sites in WRN deficient cells in response to c-Myc overexpression.** a) hTERT-immortalized WRN−/− fibroblasts and normal hTERT-control fibroblasts were transduced with LXSN-Myc expressing retrovirus and labeled with BrdU for 30 minutes during S-phase. Cells were stained with anti-BrdU (red) and anti-γ-H2AX (green). Images were collected with a 100X oil-immersion objective. Note the increased appearance of γ-H2AX and BrdU spots (yellow) in WRN−/− cells expressing c-Myc versus vector, LXSN, transduced WRN−/− cells or WRN+/+ cells also expressing c-Myc. b) BrdU and γ -H2AX co-localized foci from (a) were counted in 10 cells/condition (Imaris software). c) HFF-Myc and HFF-pB cells were transduced with retroviruses encoding two different short hairpins to WRN, siWRN I and II, or scrambled control and labeled with BrdU at the first passage after infection. Knockdown using either siWRN virus caused increased co-localization of γ -H2AX and BrdU of experiment shown in (c). d) Quantitation of γ -H2AX and BrdU of experiment shown in (c). e) Western analysis of a parallel transduction of HFFs indicating the WRN knockdown achieved with siWRN I and II. Actin is shown as loading control.
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Figure 4. Replication stress induced in WRN deficient cells by c-Myc overexpression. a) \textit{hTERT-WRN}^{+/--} cells or control \textit{hTERT}-immortalized fibroblasts were transduced with LXSN-Myc and stained with anti-RPA (subunit 32) antibodies. Images were collected with a 40X oil-immersion objective. WRN^{+/--} cells expressing c-Myc showed prominent RPA nuclear staining. b) Cells as in (a) were subjected to a Triton extraction prior to fixation and stained for p-ATR (Ser428) to reveal the p-ATR associated with the chromatin resistant fraction. Images were collected with a 100X oil-immersion objective. Specific staining occurred in WRN^{+/--} cells upon c-Myc overexpression and was not detectable in the control \textit{hTERT}-immortalized normal fibroblasts. c) Increased p-ATR (Ser428) and RPA (subunit 70) in WRN^{+/--} cells upon c-Myc overexpression was confirmed by Western analysis. In addition increased levels of p-CHK1, p-CHK2, and p-p53 by c-Myc were detected in WRN^{+/--} cells. HFFs exposed to UV are shown as positive control of DNA damage response.

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has been employed to selectively detect relocation of the Mre11 complex upon irradiation of cells [29]. WRN−/− cells overexpressing c-Myc showed prominent phospho-ATR nuclear staining that was resistant to Triton extraction (Fig. 4b), confirming the activation of checkpoints by replication-associated damage. The increased level of p-ATR and RPA in c-Myc expressing WRN−/− cells was confirmed by Western blot analysis (Fig. 4c).

DNA damage checkpoint signaling is mediated through the CHK1 and CHK2 kinases that, through phosphorylation of targets such as p53, CDG25A, and CDG25C, lead to cell cycle arrest [27]. Thus, we examined the levels of phospho-CHK1, CHK2, and p53 by Western blot analysis. The results shown in Fig. 4c indicate that indeed c-Myc overexpression led to increased phosphorylation of these downstream mediators selectively in WRN deficient cells, (Fig. 4c). These results, together with the increased accumulation of γ-H2AX foci at sites of newly replicated DNA, indicate the presence of replication stress. Thus, WRN is required to minimize replication-associated damage in response to c-Myc driven S-phase.

p53 is required for the intra-S checkpoint activated by low doses of radiation [30]. This is consistent with our findings of increased phosphorylation of p53. To test if the activation of p53 consequent to replication-associated DNA damage in c-Myc expressing WRN−/− cells was responsible for the senescence response seen in these cells [24], we utilized stable RNA interference to reduce p53 (sh-p53). Simultaneous transduction of WRN−/− cells with c-Myc and sh-p53 expressing retroviruses led to their continuous proliferation and bypassing of senescence, as shown by the lack of β-galactosidase staining (SA β-Gal), when compared to WRN−/− + c-Myc cells with intact p53 (Fig. 5b and 5c). Silencing of the major transcriptional target of p53, the Cdk inhibitor p21, also allowed proliferation and rescued senescence of the WRN−/− cells expressing c-Myc (data not shown). These experiments indicate that DNA damage caused by c-Myc in WRN deficient cells ultimately leads to activation of p53, cell-cycle arrest and senescence. A role for p53 in mediating cell-cycle arrest in response to c-Myc activation has been shown previously, albeit independently of abrogation of DNA repair mechanisms [31]. This is contrast with our results, obtained in both primary HFFs and hTERT-immortalized fibroblasts, where cell-cycle arrest and senescence are only observed if WRN is depleted (this work and [24]). Possible explanations to this discrepancy are discussed below.

In summary, using primary human fibroblasts we have shown that c-Myc directly regulates S-phase duration. This was shown by both overexpression as well as loss of function experiments. The acceleration of S-phase caused by c-Myc leads to increased replication-associated damage that is exacerbated by loss of WRN function, consistent with the role for WRN in recovery of damaged DNA [32,7], multiple underlying mechanisms have been proposed, one of which is increased generation of ROS [3,9,33] and [10]. However, it was recently shown that DNA damage in c-Myc overexpressing cells could accumulate independently of ROS [34]. Here, we propose that the dramatic acceleration of S-phase in c-Myc overexpressing cells leads to an accumulation of abnormal replication structures that, if left unrepaired because of WRN depletion, become sites of double stranded breaks. It is important to notice that in normal early passage HFFs, such as those used here, c-Myc constitutive overexpression leads to low level of DNA damage unless WRN is depleted. This is true both in early passage cells and upon c-Myc mediated immortalization, where no chromosomal translocations are generated [Benanti et al, 2007 and Fig. 4b]. These observations differ from the rapid occurrence of chromatin breaks and chromosomal translocations detected by acute activation of Myc-ER [7]. We attribute this difference to both the cell type as well as to the constitutive versus conditional expression of c-Myc. Consistently, HFFs do not display cell-cycle arrest and senescence in response to either Ras or c-Myc expression [35,21]. Finally, in these early passage HFFs, c-Myc induced DNA damage and senescence only occurs if there is interference with a DNA-repair pathway, indicating that it is likely that different tissue types and perhaps the age of the organism, may determine the outcome of oncogenic stimulation and to what extent senescence might be an effective barrier to cancer development in vivo.

Recent literature indicates that a hyper-replication phenotype leading to a DNA damage response and senescence is a common consequence to expression of activated ras and mos [36,37]. However, the effect of these oncoproteins upon the rate of S-phase and the co-localization of damage with replication sites was not reported. Thus, we add a new facet to this work. c-Myc shortens the duration of S-phase, which is different from the previous notion of c-Myc causing accelerated entry into S, through simultaneous activation of GI cyclins [38]. There are very few examples in the literature of genetic regulation of S-phase duration. Loss of function of Acfl, a chromatin assembly factor that influences nucleosome periodicity in Drosophila, leads to a fast S-phase transit [39]. Similarly, depletion of the histone linker H1 in slime mold also leads to a dramatic acceleration of S-phase, consistent with the observed inhibitory activity of chromatin packaging upon DNA replication [40]. These findings suggest that higher order chromatin structure influences replication dynamics and origin usage [reviewed by [41]]. The possibility that c-Myc may exert its effect upon DNA replication though alteration of chromatin structure is intriguing in light of the role of c-Myc in recruiting histone acetylase complexes [42] and its direct binding to known replication origins [1]. For example, studies in Drosophila have shown that tethering histone acetylases to chromatin increases origin activity [43].

Finally, the finding that c-Myc sensitizes cells to replication-stress indicates that replication-coupled DNA repair might represent an Achilles’ heel of c-Myc-driven tumors to which new therapeutic approaches might be developed.

Methods

Cell Culture and viral transductions

For cell cycle analysis shown in Fig. 1, 2 and 3, primary human foreskin fibroblasts (HFF) were generated from fresh neonatal human foreskins and infected with retroviruses within the first ten population doublings as described [21]. The TGR c-myc−/− rat fibroblast line and TGR parental cells, shown in Fig. 1b, were obtained from J. Sedivy’s Laboratory [22]. The TGR c-myc−/− cells were reconstituted to express c-Myc by transduction with the retroviral vector pBabe-puro expressing human c-Myc. All cells were cultured in Dulbecco’s Modified Eagles Medium (DMEM) supplemented with 10% FBS, penicillin and streptomycin. WRN deficient fibroblasts, shown in Fig. 4 and 5, immortalized with hTert as well as normal hTert-immortalized fibroblasts were previously described [24]. Overexpression of c-Myc in these cells was obtained using the retroviral vector pLXSN expressing human c-Myc. For UV irradiation, HFFs were exposed to 20 mJoules/cm² for two minutes and then collected six hrs post treatment.
Stable RNA interference and SA-β-galactosidase staining

Retroviral vectors pB-siWRN#1 and pB-sip53 were previously described [24]. pB-siWRN#2 contained a hairpin corresponding to the human WRN cDNA bps 2524–42 (Genbank #L76937): 5′-ggtccaacaatcatctact-3′. For the experiment shown in Fig. 5, cells were exposed to both LXSN-Myc and pB-si-p53 viruses and selected with G418 to allow selection of c-Myc overexpressing cells. At passage two and three after selection cells were stained for SA-β-Gal as described [44].

FACS analysis and BrdU labeling

For the pulse chase experiments shown in Fig. 1 cells were synchronized by thymidine block (2.5 mM) for sixteen hours, released and labeled with 200 uM BrdU for 30 min (HFFs) or...
50 uM for one hour (TGR cells), and either fixed immediately in 70% ethanol or chased in BrdU-free media and collected at the indicated times. Nuclei were isolated from fixed cells, stained with fluorescein isothiocyanate (FITC) conjugated anti-BrdU antibody (Becton Dickinson), and resuspended in 50 ug propidium iodide per ml. Nuclei were then analyzed on a FACs Scan instrument (Becton Dickinson), and cell cycle fractions were quantified with CellQuest software (Becton Dickinson).

Western blotting

Whole-cell lysates for Western blotting were prepared by trypsinization of cells, followed by PBS wash, and then lysing cell pellets directly in SDS-sample loading buffer. Samples were normalized by using equivalent cell numbers. SDS-polyacrylamide gels were transferred to Immobilon-P membranes (Millipore). Blots were probed with antibodies to WRN (611169, BD PharMingen); p33 (Ab-6 OP43, Oncogene Research Products); c-Myc (Mab C-33, Santa Cruz); actin (I-19, Santa Cruz). All other antibodies were from Cell Signaling: anti-p-ATR (S-428 #3853S); anti-p-CHK1 (Ser 296, #2549); anti-p-CHK1 (Thr 68, #2661); anti-p-H2AX (Ser 139, #2577); anti-p-p53 (Ser 15, #9286); anti-RPA 70 (#2267).

Immunofluorescence microscopy

Immunofluorescence was performed on cells grown on coverslips (#1 1/2 x 18 mm). Cells were synchronized in S-phase by either growing to confluence and then fixing fourteen -sixteen hrs after re-plating (Fig. 3 and 4) or by Thymidine block (Fig. 2), release and labeling with BrdU (200 uM) for 30 min at the indicated times prior to fixation. Cells were fixed with 4% paraformaldehyde for fifteen minutes at room temperature, washed with PBS and permeabilized with ice cold MeOH/ Acetone (1:1) for 30 sec. After blocking with PBS containing 0.005% TWEEN-20 and 5% goat serum for 30 min at 37°C, cover-slips were incubated for one-two hrs at 37°C with the following antibodies: mouse anti-BrdU (SIGMA B-23531), rabbit anti-WRN rabbit (gift from Ashwin Kamath-Loeb), mouse anti-RPA-32 (#NA18, Calbiochem), rabbit anti-c-Myc (N-262, Santa Cruz), rabbit anti-p-ATR (S-428 #3853S, Cell Signaling), anti-γH2AX (#07164, Upstate). The anti-p-ATR was designed to recognize p-ser428, a phospho-site predicted by Scansite algorithm (Scansite,.mit.edu)

Verifed as in vivo phospho-site upon UV irradiation of cultured cells (Cell Signaling). Rhodamine or FITC-conjugated goat anti-rabbit IgG or goat anti-mouse IgG (Jackson Laboratories) were used as secondary antibodies at 1:1000 in blocking buffer. For nuclear staining, cover-slips were incubated with Hoechst for five min, washed and mounted on glass slides with DAPCO (1,4-Diazobicyclo-(2,2,2) octane (Sigma), spectrophotometric grade glycerol (Kodak). 625 mg DAPCO, 22.5 ml spectrosglycerol, 2.5 ml 1XPBS pH 8.6). Staining of Triton resistant nuclear fractions with anti-p-ATR (Fig. 5) was accomplished by washing cells twice with ice-cold PBS, extraction in Cytoskeleton Buffer (10 mM PIPES pH 6.8, 100 mM NaCl, 300 mM sucrose, 3 mM MgCl2, 1 mM EGTA, 0.5% Triton X-100) for five minutes on ice, followed by two cold rinses in PBS prior to fixing with 4% paraformaldehyde fifteen minutes. Staining was then carried on as above. Confocal images (stacks) were acquired at 0.2 micron spacing with an Olympus 100 x and a 40 x Oil immersion objective as specified in the figure legends with an Applied Precision DeltaVision RT microscope system (Applied Precision, Issaquah, WA). The exposure times were kept constant for each fluorescence channel within each experiment and antibody used. Stacks were deconvolved using a constrained iterative algorithm with DeltaVision SoftWorx software.

Supporting Information

Movie S1 Co-localization of WRN and BrdU in HFF-Myc. HFF-Myc synchronized by double thymidine block were released in S phase and labeled after 1 hr with BrdU for 30’ prior to fixation. WRN and BrdU were detected with specific antibodies followed by a FITC or Rhodamine conjugate secondary, respectively. Yellow spots present throughout the nucleus indicate overlap of WRN protein with newly replicated DNA. Images were collected using an Olympus 100x immersion objective with an Applied Precision DeltaVision RT microscope system (Applied Precision, Issaquah, WA). 3D reconstruction was obtained using Imaris software (Bitplane).

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Movie S2 Predominant nuclear localization of WRN in HFF-pB. HFF-pB were synchronized and processed as for Movie 1. In these cells WRN staining (green) is shown mostly in nucleoli and only a few extra-nuclear sites coincide with BrdU.

Found at: doi:10.1371/journal.pone.0005951.s002 (10.18 MB MPG)

Figure S1 Cell-cycle profile of HFF-pB and HFF-Myc after Thymidine block. HFFs, synchronized by double thymidine block, were stained with Propidium iodide and analyzed by FACS. Both cell populations HFF-pB and HFF-Myc show an arrest at the G1/ S boundary. HFF’s exponentially growing cell cycle profiles are shown as reference.

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Author Contributions

Conceived and designed the experiments: CG. Performed the experiments: KR NA CG. Analyzed the data: KR NA CG. Contributed reagents/ materials/analysis tools: DG CG. Wrote the paper: CG.

References

1. Dominguez-Sola D, Ying CY, Grandori C, Ruggerio L, Chen B, et al. (2007) Non-transcriptional control of DNA replication by c-Myc. Nature 448: 445–451.
2. Gomez-Roman N, Grandori C, Eisenman RN, White RJ (2003) Direct activation of RNA polymerase III transcription by c-Myc. Nature 421: 290–294.
3. Grandori C, Gomez-Roman N, Felton-Edkins ZA, Ngouenet C, Galloway DA, et al. (2003) c-Myc binds to human ribosomal DNA and stimulates transcription of rRNA genes by RNA polymerase I. Nat Cell Biol 7: 311–318.
4. Dang CV, O’Donnell KA, Zeller KI, Nguyen T, Othuser RC, et al. (2006) The c-Myc target gene network. Semin Cancer Biol 16: 253–264.
5. Knoepfler PS, Zhang XY, Cheng PF, Galen PK, McMahon SR, et al. (2006) Myc influences global chromatin structure. Embo J 25: 2723–2734.

6. Mai S, Hanley-Hyde J, Fluri M (1999) c-Myc overexpression associated DHFR gene amplification inhamster, rat, mouse and human cells. Oncogene 12: 277–283.

7. Felscher DW, Bishop JM (1999) Transient excess of MYC activity can elicit genomic instability and tumorigenesis. Proc Natl Acad Sci U S A 96: 3940–3944.

8. Li Q, Dang CV (1999) c-Myc overexpression uncouples DNA replication from mitosis. Mol Cell Biol 19: 5339–5351.

9. Karlsson A, Deh-Basu D, Cherry A, Turner S, Ford J, et al. (2003) Defective double-strand DNA break repair and chromosomal translocations by MYC overexpression. Proc Natl Acad Sci U S A 100: 9974–9979.

10. Vafa O, Wade M, Kern S, Beeche M, Paulita TK, et al. (2002) c-Myc Can Induce DNA Damage, Increase Reactive Oxygen Species, and Mitigate p53 Function. A Mechanism for Oncogene-Induced Genetic Instability. Mol Cell 9: 1031–1044.

11. Koch HB, Zhang R, Verdoold B, Bailey A, Zhang CD, et al. (2007) Large-scale identification of c-MYC-associated proteins using a combined TAP/MudPIT approach. Cell Cycle 6: 205–217.

12. Sidorova JM, Li N, Folch A, Momrat RJ Jr (2008) The RecQ helicase WRN is required for normal replication fork progression after DNA damage or replication fork arrest. Cell Cycle 7.

13. Brosh RM Jr, Bohr VA (2007) Human premature aging, DNA repair and RecQ helicases. Nucleic Acids Res 35: 7527–7544.

14. Martin GM, Oshima J (2000) Lessons from human progeroid syndromes. Nature 408: 263–266.

15. Poot M, Hoehn H, Runger TM, Martin GM (1992) Impaired S-phase transit of Werner syndrome cells expressed in lymphoblastoid cell lines. Exp Cell Res 202: 267–273.

16. Salk D, Bryant E, Hoehn H, Johnston P, Martin GM (1985) Growth Differ 8: 1039–1048.

17. Dhillon KK, Sidorova J, Saintigny Y, Poot M, Gollahon K, et al. (2007) Functional role of the Werner syndrome RecQ helicase in human fibroblasts. Aging Cell 6: 53–61.

18. Choi D, Whittier PS, Oshima J, Funk WD (2001) Telomerase expression prevents replicative senescence but does not fully reset mRNA expression patterns in Werner syndrome cell strains. FEBS J 15: 1014–1020.

19. Chang S, Midtiani AS, Cabrera NG, Naylor ML, Laid P, et al. (2004) Essential role of limiting telomeres in the pathogenesis of Werner syndrome. Nat Genet 36: 877–882.

20. Opresko PL, Oterlei M, Graakjaer J, Bruheim P, Dawut L, et al. (2004) The Werner syndrome helicase and exusuducase cooperate to resolve telomeric D loops in a manner regulated by TRF1 and TRF2. Mol Cell 14: 763–774.

21. Benanti JA, Wang ML, Myers HE, Robinson KL, Grandori C, et al. (2007) Epigenetic down-regulation of ARF expression is a selection step in immortalization of human fibroblasts by c-Myc. Mol Cancer Res 5: 1181–1189.

22. Mateyak MK, Obaya AJ, Adachi S, Sedivy JM (1997) Phenotypes of c-Myc-deficient rat fibroblasts isolated by targeted homologous recombination. Cell Growth Differ 8: 1039–1048.

23. Gray MD, Wang L, Youssoufiyan H, Martin GM, Oshima J (1998) Werner helicase is localized to transcriptionally active nuclei of cycling cells. Exp Cell Res 242: 487–494.

24. Grandori C, Wu KJ, Fernandez P, Nguyenet C, Grimal J, et al. (2003) Werner syndrome protein limits MYC-induced cellular senescence. Genes Dev 17: 1569–1574.

25. Sakamoto S, Nishikawa K, Hase SJ, Goto M, Furuchi Y, et al. (2003) Werner helicase relocates into nuclear foci in response to DNA damage and associates with RPA and Rad51. Genes Cells 6: 421–430.

26. Feijoo C, Hall-Jackson C, Wu R, Jenkins D, Leitch J, et al. (2001) Activation of mammalian Chk1 during DNA replication arrest: a role for Chk1 in the intra-S phase checkpoint monitoring replication origin firing. J Cell Biol 154: 913–923.

27. Li L, Zou L (2005) Sensing, signaling, and responding to DNA damage: organization of the checkpoint pathways in mammalian cells. J Cell Biochem 94: 290–306.

28. Cuadrado M, Martinez-Pastor B, Maroga M, Toledano ML, Gutierrez-Martinez P, et al. (2006) ATM regulates ATR chromatin loading in response to DNA double-strand breaks. J Exp Med 203: 297–303.

29. Mizzone OK, Perrini JH (2001) DNA damage-dependent nuclear dynamics of the Mvr11 complex. Mol Cell Biol 21: 281–288.

30. Shimura T, Toyoshima M, Adiga SK, Kuno T, Naga I, et al. (2006) Suppression of replication fork progression in low-dose-specific p53-dependent S-phase DNA damage checkpoint. Oncogene 25: 3921–3932.

31. Felscher DW, Zetterberg A, Zhou J, Tsy T, Bishop JM (2008) Overexpression of MYC causes p53-dependent G2 arrest of normal fibroblasts. Proc Natl Acad Sci U S A 105: 10544–10548.

32. Mai S, Fluri M, Staverski D, Huppi K (1996) Genomic instability in MycER-activated Rat1A-MycER cells. Chromosoma Res 4: 365–371.

33. Louis SF, Vermolen BJ, Grunin Y, Young IF, Guffee A, et al. (2005) c-Myc induces chromosomal rearrangements through telomere and chromosome remodeling in the interphase nucleus. Proc Natl Acad Sci U S A 102: 9613–9618.

34. Ray S, Askari KR, Deh-Basu D, Adler AS, Chang HY, et al. (2006) MYC can induce DNA breaks in vivo and in vitro independent of reactive oxygen species. Cancer Res 66: 6598–6605.

35. Benanti JA, Galloway DA (2004) Normal human fibroblasts are resistant to RA-induced senescence. Mol Cell Biol 24: 2812–2815.

36. Di Meoco R, Fumagalli M, Giralde A, Piccinin S, Gasparini P, et al. (2006) Oncogene-induced senescence is a DNA damage response triggered by DNA hyper-replication. Nature 444: 630–636.

37. Bartkova J, Rezaei N, Liontos M, Karakaidos P, Kletsas D, et al. (2006) Oncogene-induced senescence is part of the tumorigenesis barrier imposed by DNA damage checkpoints. Nature 444: 633–637.

38. Puch O, Bernaschek G, Eilers M, Hengstschlager M (1997) Activation of c-Myc uncouples DNA replication from activation of G1-cyclin-dependent kinases. Oncogene 15: 649–656.

39. Fyodorov DV, Blower MD, Karpen GH, Kadonaga JT (2004) Acf1 confers unique activities to ACF/CHRAC and promotes the formation rather than disruption of chromatin in vivo. Genes Dev 18: 170–183.

40. Thiriet C, Hayes JF (2000) Linker histone phosphorylation regulates global timing of replication origin firing. J Biol Chem.

41. Gerbi SA, Biehlmyer AK (2002) DNA replication and chromatin. Curr Opin Genet Dev 12: 243–248.

42. McMahon SB, Wood MA, Cole MD (2000) The essential cofactor TRRAP recruits the histone acetyltransferase HGCN5 to c-Myc. Molecular & Cellular Biology 20: 556–562.

43. Aggarwal BD, Calvi BR (2004) Chromatin regulates origin activity in Drosophila follicle cells. Nature 430: 572–576.

44. Dinur GP, Lee X, Basle G, Acosta M, Scott G, et al. (1995) A biomarker that identifies senescence human cells in culture and in aging skin in vivo. Proc Natl Acad Sci U S A 92: 9363–9367.