Timeless Maintains Genomic Stability and Suppresses Sister Chromatid Exchange during Unperturbed DNA Replication

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Genome integrity is maintained during DNA replication by coordination of various replisome-regulated processes. Although it is known that Timeless (Tim) is a replisome component that participates in replication checkpoint responses to genotoxic stress, its importance for genome maintenance during normal DNA synthesis has not been reported. Here we demonstrate that Tim reduction leads to genomic instability during unperturbed DNA replication, culminating in increased chromatid breaks and translocations (triradials, quadradrials, and fusions). Tim deficiency led to increased H2AX phosphorylation and Rad51 and Rad52 foci formation selectively during DNA synthesis and caused a 3–4-fold increase in sister chromatid exchange. The sister chromatid exchange events stimulated DNA synthesis and caused a 3–4-fold increase in sister chromatid exchange. These results indicate that Tim deficiency leads to genomic instability during unperturbed DNA replication, culminating in increased chromatin breaks and translocations (triradials, quadradrials, and fusions). Tim deficiency led to increased H2AX phosphorylation and Rad51 and Rad52 foci formation selectively during DNA synthesis and caused a 3–4-fold increase in sister chromatid exchange. The sister chromatid exchange events stimulated DNA synthesis and caused a 3–4-fold increase in sister chromatid exchange. Therefore, Tim deficiency leads to an increased reliance on homologous recombination for proper continuation of DNA synthesis. Together, these results indicate a pivotal role for Tim in maintaining genome stability throughout normal DNA replication.

DNA synthesis utilizes complex sets of genes and functionalities to prevent genome maintenance failures. These potential failures include the misincorporation of nucleotides, the collapse of replication forks, and the formation of secondary structures that lead to chromosome deletions, duplications, and other mutagenic events (1–4).

In addition to polymerases and their cofactors, several higher order protein complexes act in concert during DNA replication to promote chromatin decondensation, DNA duplex unwinding, and protection of the resulting single-stranded DNA (ssDNA). Failure to efficiently coordinate these processes can lead to stalling and collapse of the replication fork into double strand breaks (DSBs). For example, in budding yeast, deficiencies in DNA priming caused by low levels of pol α lead to a 22-fold increase in mitotic recombination (5). Similarly, in vertebrate cells, treatment with DNA polymerase inhibitors (e.g. aphidicolin) causes the generation of short segments of ssDNA, via polymerase-helicase uncoupling (6–11), and increases chromatid breaks at common fragile sites (12). The importance of polymerase processivity in genome stabilization raises the question to what degree the other components of the replication apparatus participate in genome maintenance.

Tim and its putative orthologs in yeast (Swi1 in Schizosaccharomyces pombe and Tof1 in Saccharomyces cerevisiae) facilitate DNA replication. Although less is known about Tim function in mammals, several functions of Swi1 and Tof1 have recently been described. Swi1 and Tof1 associate with chromatin and travel with the replication fork during S phase and are required for normal pausing at replication fork barriers (13–20). In addition, Tof1 associates with Cdc45 and MCM6, and deletion of Tof1 can lead to slowed replication fork progression (13, 17, 21).

Importantly, recent studies indicate that Swi1 and Tof1 coordinate DNA synthesis with DNA unwinding when replication is stalled (13). Using ChIP analysis, Tof1 deletion has been shown to cause the spatial uncoupling of replisome components from newly synthesized DNA when replication forks are stalled by hydroxyurea (HU) treatment. In addition, extended stretches of ssDNA have been observed in swi1 mutants, consistent with uncoupling of DNA unwinding from synthesis (22). Finally, loss of Swi1 increases inter-sister recombination during S phase, as determined by the accumulation of X-shaped DNA structures in swi1mus81 double mutants (15). These data, taken together, point to a role for Swi1 and Tof1 in both replisome maintenance and replication fork stability.

Several functions of Swi1 and Tof1 have been shown to be conserved either in Tim or in its vertebrate binding partner, Tipin. Tim and Tipin associate with chromatin as an interdependent complex (23) during S phase, putatively though the RPA34 binding domain of Tipin (24, 25). In addition, the Tim-Tipin complex associates with other components of the DNA replication apparatus, including Claspin, MCM subunits, pol δ, and pol ε (24, 26, 27). Furthermore, reduced expression of Tim modified Eagle’s medium; PBS, phosphate-buffered saline; PCNA, proliferating cell nuclear antigen; BrdUrd, bromodeoxyuridine; 7-AAD, 7-aminoactinomycin D.
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slows DNA replication rates, as measured by cell cycle profile changes and DNA fiber labeling (24, 25, 27, 28). Consistent with a role for the Tim-Tipin complex in suppressing the accumulation of ssDNA at replication forks, pharmacological inhibition of DNA replication in Tipin-depleted Xenopus extracts leads to a 2-fold increase in chromatin-associated RPA (26).

Most studies to date have focused on the functions of Tim-Tipin under genotoxic stress (24–29). Although it is known that Tim can associate with the replisome components during S phase, and that its depletion leads to decreased rates of DNA synthesis, the role of Tim in genome stability during normal DNA replication is not well characterized. Here we demonstrate that Tim is required to maintain genome stability even in the absence of exogenous DNA-damaging agents. We show that Tim reduction leads to increased chromatic breaks, translocations, and inter-sister recombination events, as revealed by Rad52 and Rad51 focal accumulation and increased rates of sister chromatid exchange (SCE). Increased SCEs in Tim-deficient cells are at least partially dependent on Brca2 and Rad51, indicating that Tim dysfunction leads to an increased reliance on homologous recombination for continuation of DNA synthesis. These data demonstrate that Tim, an important component of the DNA replication apparatus, is required for maintaining genome integrity during unperturbed DNA replication.

EXPERIMENTAL PROCEDURES

Cell Culture—Blm−/− and Rad52−/− embryos were generated by intercrossing mice heterozygous for the Blm+/− gene and Rad52 null alleles, respectively (30, 31). Blm−/− and Rad52−/− murine embryonic fibroblasts (MEFs) were generated from E12.0 and E14.5 embryos, respectively, and immortalized by stable expression of SV40 large T antigen (32). MEFs and NIH3T3 cells were grown in Dulbecco’s modified Eagle’s medium (DMEM) containing 10% fetal bovine serum (FBS) in a 5% CO2 atmosphere. NIH3T3 cells were infected in 0.1% FBS/DMEM. Twenty eight hours after infection, MEFs were stimulated with 10% FBS to re-enter the cell cycle.

BrdUrd Flow Cytometric Analysis—The percentage of cells in S phase (Fig. 1B) was quantified by pulsing similarly cultured NIH3T3 cells with 10 μM BrdUrd (Sigma) for 30 min. Cells were then fixed in 70% ethanol, and DNA was denatured using 3 N HCl, 0.5% Tween 20. Cells were stained with anti-BrdUrd antibodies (1:200, Pharmingen) for 20 min, followed by detection with fluorescein isothiocyanate-conjugated goat anti-mouse secondary antibodies (1:200, Pharmingen) and propidium iodide (2.5 μg/ml) for DNA content. BrdUrd-positive cells were quantified by flow cytometry using a FACSCalibur (BD Biosciences).

Western Blots and Quantitative Real Time PCR—Whole cell extracts were prepared by washing suspended cells with ice-cold PBS, followed by direct lysis in Laemmli sample buffer, and boiling for 5 min to denature proteins and DNA. Proteins were separated by SDS-PAGE and transferred to polyvinylidene difluoride (Millipore). Primary antibodies for protein detection include the following: anti-actin (1:2000, Santa Cruz Biotechnology, Inc.), anti-glyceraldehyde-3-phosphate dehydrogenase (1:2000, USBiological), anti-phospho-H2AX clone JBW301 (1:2000, Upstate), anti-Rad51 (1:500, Santa Cruz Biotechnology, Inc.), and anti-Stat3 (1:1000, Cell Signaling). The Timless antibody (36) was generously provided by P. Minoo (University of Southern California, Los Angeles). Secondary horseradish peroxidase-conjugated species-specific antibodies were diluted 1:2000 (Santa Cruz Biotechnology, Inc.). For real time PCR quantification, RNA from Rad52−/+ and Rad52−/− immortalized MEFs was isolated by TRIzol (Invitrogen) and chloroform extractions, and purified using an RNeasy kit (Qiagen). cDNA was prepared using a high capacity cDNA reverse transcription kit (Applied Biosystems). Primer and probes sets for β-actin (Mm 00607939 s1), Brca2 (Mm 00464783 m1), and Tim (Mm 00495610 m1) along with Taqman universal master mix were purchased from Applied Biosystems. Samples were run on an Applied Biosystems 7900HT Fast Real-Time PCR System, and data were analyzed using Applied Biosystems software.

Mitotic Spreads and SCE—For quantification of chromosome breaks, mitotic spreads were prepared as described previously (37) and stained with Giemsa. To visualize SCE, synchronized cells stimulated with 10% FBS were incubated with 10 μM BrdUrd (Sigma) for two consecutive S phases and treated with 0.5 μM nocodazole 3–4 h before collection. For aphidicolin treatment experiments, 0.2 μM aphidicolin (Calbiochem) was added prior to the second S phase in BrdUrd-containing media (15–16 h before nocodazole addition); 5 μM aphidicolin was added for 1 h, at a time point corresponding to the second S phase in BrdUrd-containing media, and was subsequently removed to allow cells to reinitiate cell cycle progression for 3 h before addition of nocodazole. For UV treatment experiments, cells in the second S phase of BrdUrd incorporation were treated with 5 or 10 J/m2 UV light (λ = 365 nm) 4 h before the addition of nocodazole. Nocodazole-treated cells were resuspended by trypsinization, washed with PBS, incubated in 75 mM KCl for 20 min, fixed in methanol/acetic acid (3:1), and dropped onto prewarmed slides at 37 °C. Slides were incubated in 10 μg/ml Hoechst 33258 (Invitrogen) in H2O for 20 min and then
washed in MacIlvaine solution (164 mM Na₂HPO₄, 16 mM citric acid, pH 7.0). Slides were then covered with glass coverslips and exposed to UV light (λ = 365 nm) for 30 min followed by a 1-h incubation in 1× SSC (20× = 3 mM NaCl, 0.3 mM Na₃citrate*2H₂O, pH 7.0) at 55 °C. Metaphase spreads were stained with Giemsa and visualized using an Olympus BX41 microscope (model U-DO3) with a 100× oil objective. SCEs and chromatid breaks were quantified from photographs taken with a Spot 3-shot insight QE camera (model 4.3) using Spot advanced software.

**Cell Proliferation and Apoptosis Assays**—Cells were infected in 10% FBS/DMEM. Twenty four hours later, cells were plated at low density in 6-well plates in triplicate for cell proliferation or on 6-cm dishes for apoptosis. Forty eight hours after plating, cell number was quantified using a Coulter Z2 counter or, for apoptosis, staining with annexin-V (Pharmingen) and 7-AAD (Pharmingen) in 1× annexin-V binding buffer (10× = 0.1 M Hepes-NaOH, pH 7.4, 1.4 mM NaCl, 25 mM CaCl₂) for 30 min. Apoptosis rates were determined by flow cytometry on a BD Biosciences FACSCalibur and gating for annexin-V positive/7-AAD negative events.

**Immunofluorescence Staining**—Cells were fixed at peak S phase, in 3% paraformaldehyde, 2% sucrose buffered with PBS or, for PCNA detection, in ice-cold methanol for 5 min, followed by 1 min in acetone. Cells were permeabilized in 0.5% Triton X-100/PBS for 20–60 min and immunostained with antibodies detecting PCNA (1:500, Santa Cruz Biotechnology, Inc.), γ-H2AX (1:1000, Upstate), and/or Rad51 (1:100, Santa Cruz Biotechnology, Inc.). Fluorophore-conjugated, species-specific secondary antibodies include the following: α-mouse Rhodamine Red and α-rabbit fluorescein isothiocyanate (Jackson ImmunoResearch) and α-rabbit Alexa Fluor 594 (Invitrogen), each diluted 1:1000. Foci were visualized by using a Nikon Eclipse E-800 microscope with a 100× oil objective. Rad51 and Rad52 foci, identified by significant distinction from nuclear background fluorescence, were quantified from images acquired using Metamorph software in Adobe Photoshop. Double blind methods were used for all quantifications.

**Statistical Tests and Foci Quantification**—For all figures, error bars represent standard errors, and p values were calculated by Student’s t test. At least three independent replicates were performed for each experiment.

**RESULTS**

**Tim Deficiency Increases Genomic Instability**—Although Tim and Tipin have been shown to play partial roles in checkpoint responses to UV and HU treatment (24, 25, 27–29), they are also required for efficient DNA replication in the absence of such genotoxic stresses (24, 25, 27, 28). To investigate the role of Tim in maintaining genome stability during normal cellular proliferation, two independent shRNAs were used to reduce Tim expression to near undetectable levels in synchronized NIH3T3 cells (Fig. 1A). Genome stability was then assessed by two methods: H2AX phosphorylation and chromosome spreads. H2AX phosphorylation, a marker of DSB formation (38), was quantified in G₁-enriched cells (Fig. 1B, 1st and 2nd lanes) and in cells at peak S phase following growth factor stimulation (Fig. 1B, 3rd and 4th lanes). As expected, H2AX phosphorylation increased upon passage from G₁ into S phase in control cells (39); however, this increase was substantially enhanced by Tim reduction (Fig. 1B), consistent with the reported effect of Tipin knockdown (27). The increase in H2AX phosphorylation following Tim reduction was slightly greater than that observed following 1 gray ionizing radiation of asynchronous cells (data not shown), indicating that the level of DSB formation upon entry of Tim-deficient cells into S phase was considerable.

To determine whether the instability caused by Tim reduction persisted into mitosis, chromosome spreads were prepared. Reduction of Tim expression utilizing two independent shRNAs resulted in a significant increase in chromatid breaks in comparison with controls (Fig. 1C, D). In addition, whereas control cells exhibited few abnormal chromosome structures (triradials, quadiradials, and fusions), Tim reduction caused a 5–7-fold increase in these types of chromosome rearrangements (Fig. 1D). Taken together, these data indicate that Tim reduction is sufficient to cause genomic instability in the absence of exogenous genotoxic stress.
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**FIGURE 2.** Rad51 and Rad52 foci accumulate following Tim reduction. A, analysis of Rad52 and γH2AX foci accumulation in NIH3T3 cells stably expressing GFP-Rad52. Serum-starved cells were infected with control or Tim #5 shRNA-expressing lentivirus as described in Fig. 1 and collected during peak S phase (16–17 h after growth factor stimulation). The number of Rad52 foci per cell was detected and quantified by GFP fluorescence; γH2AX was immunodetected using anti-phospho-H2AX antibodies. B, quantification of the average number of Rad51 and Rad52 foci from A. C, quantification of S phase and non-S phase cells exhibiting Rad52 foci. D, quantification of the percentage of PCNA-positive cells exhibiting Rad52 foci. E, quantification of the percentage of PCNA-positive cells exhibiting Rad52 foci.

**Tim Reduction Increases Rad51 and Rad52 Foci Formation during S Phase**—Previous studies have shown that abnormalities during DNA replication can lead to DSB formation through replication fork collapse, causing a reliance on homologous recombination (HR)-mediated restart mechanisms (40). In addition, Rad52-dependent HR has also been proposed to counter the untoward effects of ssDNA accumulation upon helicase-polymerase uncoupling (15).

To assess whether Tim reduction leads to the accumulation of factors that participate in HR, Rad51 and Rad52 foci formation were quantified (41–43) in synchronized NIH3T3 cells. As shown in Fig. 2A, Tim reduction led to a significant increase in cells exhibiting high numbers of Rad52 foci, as determined by visualization of stably expressed GFP-Rad52. In contrast to controls, which generally exhibited few foci (<5 per cell), the vast majority of Tim knockdown cells (89.1%) had greater than 5 Rad52 foci, with 41.0% exhibiting more than 20 foci per cell. Rad52 foci predominantly overlapped with γH2AX (86.8% in control cells and 92.4% in Tim knockdown cells), indicating that Rad52 foci formed at sites of DNA damage. On average, 4.3 and 21.5 Rad52 foci were observed in control and Tim-deficient cells, respectively, a 5.7-fold increase (Fig. 2C).

Rad51 foci also overwhelmingly overlapped with phospho-H2AX (>92%) and were increased in Tim knockdown cells, although more modestly in comparison with Rad52 foci (Fig. 2B). In comparison with controls, a significant increase (p < 0.05) in Rad51 foci was observed only in Tim knockdown cells with >20 Rad51 foci per cell. On average, 13.3 Rad51 foci were observed in control cells compared with 19.0 foci in Tim knockdown cells (Fig. 2C), a 1.5-fold increase. Thus, a lesser absolute number of Rad51 foci was stimulated by Tim reduction in comparison with Rad52 foci (5.7 Rad51 versus 17.2 Rad52). This result combined with the frequency of Rad51 and Rad52 overlap (controls, 58.7% ± 4.8, and Tim-deficient, 53.2% ± 3.2; supplemental Fig. 1) suggest that only approximately half the Rad52 foci that formed as a consequence of Tim reduction were accompanied by Rad51.

To determine whether foci formation occurred during DNA synthesis, GFP-Rad52-expressing cells were stained for PCNA. Indeed, nearly all cells, either control or Tim knockdown, that exhibited Rad52 foci also had profuse PCNA staining, a hallmark of S phase (89.1% for control and 92.2% with Tim reduction; Fig. 2D). As expected, the percentage of S phase cells with Rad52 foci increased significantly in Tim-reduced cells compared with controls (Fig. 2E). Taken together, these results indicate that Tim reduction increases Rad51 and Rad52 foci, and that Rad52 foci form during S phase.

**Sister Chromatid Exchange Increases with Tim Reduction**—The increase in Rad52 and Rad51 foci in Tim-reduced cells (Fig. 2, A and B) raised the question whether the chromosomal abnormalities observed (Fig. 1, C and D) under-represent the frequency of genomic instability resulting from Tim deficiency and that HR effectively counters such instability in S phase. SCE detects recombinatorial repair events between sister chromatids during the S and G2 phases of the cell cycle and are generated by the resolution of HR intermediates (Holliday junctions) in a manner that causes crossovers (2). Recombination structures that lead to SCEs can arise during replication fork restart following collapse (2) and, purportedly, as a means to compensate for extended lagging strand gaps (15).
SCE frequency was determined in synchronized NIH3T3 control and Tim-reduced cells as a means to assess HR-mediated repair. As shown in Fig. 3A, the number of SCEs per mitotic spread increased more than 3-fold upon Tim reduction. Tim #1 and Tim #5 shRNAs increased the number of SCEs per chromosome from 0.33 in wild-type controls to 0.92 and 1.09 in Tim knockdown cells, respectively. Thus, Tim is required to prevent recombination events that lead to SCEs.

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Tim deficiency has previously been shown to slow DNA replication (24, 25, 27, 28). We next sought to compare the rate of SCE caused by Tim deficiency with that caused by other treatments known to inhibit replication rates, specifically aphidicolin and UV light exposure (Fig. 3B). Consistent with previous findings, low dose aphidicolin and UV treatment caused marginal, but statistically significant, increases in SCE rates in wild-type cells ($p = 0.0007$ and $0.015$, respectively). However, these increased rates of SCE were far less than that caused by Tim reduction alone (Fig. 3B). Moreover, treatment of Tim-deficient cells with low dose aphidicolin (0.2 mm), pulsed high-dose aphidicolin, or UV light exposure did not appreciably increase SCEs, indicating that the inherent role of Tim in preventing recombination events in S phase is not associated with excessive DNA damage or polymerase stalling. Together, these studies strongly argue that Tim has an important function in preventing recombination events during unperturbed DNA replication.

**Tim and Blm Are Non-epistatic**—Bloom syndrome, caused by a mutation in the *BLM* gene, is characterized by increased frequencies of chromosomal aberrations (chromatid breaks, triradials, and quadriradials) and, in particular, SCEs (44, 45). As a member of the RecQ helicase family, Blm aids in Holliday branch migration and a resolution pathway that does not lead to SCE (46–48). The cytogenetic similarities between Blm mutants and Tim-reduced cells, in conjunction with recent studies implying regulation of Blm through the ATR-Chk1 pathway (49, 50), led us to investigate whether increased SCEs in Tim-deficient cells was partly attributable to Tim and Blm functioning in the same pathway.

To examine Tim and Blm epistasis, Tim protein levels were reduced by shRNA (Fig. 4A) in transformed Blm wild-type and null MEFs. Changes in proliferation rates in Blm$^{+/+}$ and Blm$^{-/-}$ MEFs 72 h (~3 doublings) after lentiviral infection were determined. Although Tim reduction in Blm$^{+/+}$ MEFs led to a 2.1-fold reduction in proliferation in comparison with control shRNA-expressing Blm$^{+/+}$ and Blm$^{-/-}$ MEFs, Tim sup-
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FIGURE 4. Tim and Blm perform distinct functions. A, Western blot quantification of Tim knockdown efficiency in transformed Blm+/+ and Blm−/− MEFs. Protein samples were collected 48 h after infection. Immunodetection was performed using anti-Tim and anti-Stat3 antibodies. B, quantification of cell proliferation. Cells were infected with control or Tim #5 shRNA-expressing lentiviruses. Twenty four hours after infection, cells were replated at low density and counted 48 h later. A statistically significant decrease in proliferation was found between Blm−/− and Blm+/+ MEFs infected with Tim #5 shRNA-expressing lentiviruses (p = 0.03). C, analysis of apoptosis rates. Cells were infected with control or Tim #5 shRNA-expressing lentiviruses and co-stained with allophycocyanin-annexin-V and 7-AAD 72 h after infection; annexin-V-positive and 7-AAD-negative cells were scored as apoptotic. A statistically significant increase in apoptosis was found comparing Blm−/− and Blm+/+ MEFs infected with Tim #5 shRNA-expressing lentivirus (p = 0.02). D, example of SCE rates in Blm−/− MEFs expressing Tim #5 shRNA. The enlarged inset shows a chromosome with five SCEs, a level of SCE that was frequently observed in Tim-reduced Blm−/− MEFs. E, quantification of SCEs per chromosome in transformed Blm+/+ and Blm−/− MEFs expressing control or Tim #5 shRNAs. A statistically significant increase in SCEs per chromosome for Blm−/− cells expressing Tim #5 shRNA was observed compared with similar cells expressing control shRNA (p = 0.003).

pression in Blm−/− MEFs reduced proliferation 19.4-fold (Fig. 4B). To determine whether increased cell death contributed to the proliferative failure of Tim-deficient Blm−/− MEFs, apoptotic cells were quantified 72 h after infection (≈3 cell doublings). Although apoptosis was slightly elevated in cells deficient for Tim or Blm alone in comparison with wild-type controls (Fig. 4C), the increase in apoptotic cells increased 8-fold in Tim-reduced Blm−/− MEFs over wild-type controls. These results indicate that combined deficiency in Tim and Blm causes synthetic lethality.

To investigate whether Tim and Blm suppress inter-sister recombination as part of the same or different pathways, SCE analysis was performed in Tim-deficient Blm wild-type and null MEFs. To avoid any potential confounding effects of proliferative failure and apoptosis, SCEs were analyzed at an earlier point after Tim reduction (≈2 cell doublings). Indeed, reduction of Tim in Blm−/− cells increased SCE rates to levels greater than that observed under any other condition (Fig. 4, D and E). Tim reduction in Blm−/− MEFs led to a rate of SCE (2.26 SCEs per chromosome) that was significantly greater than that observed in either Tim-reduced Blm+/+ cells or control shRNA-infected Blm−/− cells (Fig. 4E), which exhibited SCE levels similar to previous determinations (Fig. 3) (51, 52). Of note, it is likely that the SCE rate in Tim-deficient Blm−/− MEFs is an underestimate of the actual SCE level, as it was near the upper limit of quantification (Fig. 4D, inset) and may have been influenced by inhibition of metaphase formation in the most severely affected cells. These results indicate that Tim and Blm perform distinct functions in suppressing SCE rates. Thus, the function of Tim in preventing recombination in S phase likely lies in avoiding the formation of recombination intermediates and not in regulating Blm-mediated resolution.

Sister Chromatid Exchange in Tim-deficient Cells Is Dependent on Brca2 and Rad51—Previous studies in S. pombe suggest that recombination events generated in swi1 mutants require Rad22 (mammalian Rad52 homolog), purportedly to anneal parent strands within extended lagging strand gaps in a Rad51-independent manner (15). Our results suggest that recombination events generated by Tim deficiency could be through a similar mechanism as in yeast because both Rad52 foci and SCE increased significantly when Tim was reduced.

To assess if inter-sister recombination in Tim-reduced cells requires Rad52, we determined SCE rates in transformed Rad52+/+ and Rad52−/− MEFs. Once again, Tim reduction led to a 3–4-fold increase in SCE in Rad52+/+ MEFs in comparison with control shRNA-infected Rad52+/+ and Rad52−/− MEFs (Fig. 5A). However, the absence or presence of Rad52 led to no significant differences in either SCE rates (Fig. 5A) or the number of chromatid breaks generated upon Tim reduction (Fig. 5B). These results demonstrate that recombination events generated by loss of Tim are not dependent on Rad52, and suggest that Tim deficiency in mammals leads to recombination via mechanisms distinct from those operating in swi1 mutants.

SCEs can also result from the HR-mediated reinitiation of replication forks that have collapsed into DSBs. These repair events are highly dependent on Rad51 (2). In addition, Brca2, a putative mammalian counterpart to yeast Rad52, facilitates recruitment of Rad51 to resected DSBs (53) and has also been shown to play a key role in replication fork recovery (54).
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To determine whether Rad51 and Brca2 mediate SCEs stimulated by Tim deficiency, Rad51 and Brca2 levels were suppressed by shRNA expression in combination with Tim reduction (supplemental Fig. 2, A and B). Strikingly, the majority of the SCEs resulting from Tim deficiency were suppressed by reduction of Rad51 or Brca2 (Fig. 5A). Similar effects were observed utilizing independent shRNAs targeting different regions of the Rad51 and Brca2 transcripts (supplemental Fig. 2C). Complete suppression of SCE in Tim-deficient cells was not achieved by Rad51 or Brca2 reduction (Fig. 5A), even in the absence of Rad52 (Rad52-/-, Fig. 5A). These results argue against Rad52 playing a minor compensatory function in SCE under conditions of Rad51 or Brca2 suppression. The inability of shRNA-mediated suppression of Rad51 and Brca2 to prevent all SCEs could be due to incomplete protein reduction (supplemental Fig. 2, A and B) or reflect the existence of an unknown alternative pathway that is required for this minor fraction of recombination events following Tim reduction.

Failure to undergo HR-mediated repair in Rad51-deficient cells, as indicated by reduced SCEs, would be predicted to lead to persistence of DSBs. Consistent with this expectation, dual suppression of Tim and Rad51 significantly increased chromatic breaks (Fig. 5B). Similar trends were observed upon Brca2 suppression (Fig. 5B). Together, these data strongly argue that Tim deficiency leads to a reliance on Rad51- and Brca2-mediated HR repair to maintain genome stability in S phase.

**DISCUSSION**

Prevention and repair of DNA damage during DNA replication is crucial for maintaining genome integrity and cell viability. Replisome deregulation can lead to genome instability, both through replication fork collapse and the generation of single strand gaps created by the uncoupling of polymerases from helicases (1, 6, 55). In this study, we find that Tim, a regulator of DNA replication efficiency (24, 25, 27, 28), suppresses DSBs and chromosome abnormalities during unperturbed DNA synthesis. Furthermore, we demonstrate that Tim deficiency results in an increased reliance on the HR repair factors, Rad51 and Brca2, to mediate SCE and maintain genome stability. Because the stimulatory effect of Tim reduction on SCE far exceeds that caused by exogenous DNA damage or replication inhibitors, the function of Tim in preventing recombination events during DNA synthesis is considerable, even during normal replication.

Previously, the Tim-Tipin complex has been shown to interact with proteins in the replisome and influence both Chk1 phosphorylation and the intra-S phase checkpoint in response to genotoxic stress (23–29). Because deficiencies in the ATR-Chk1 pathway lead to increased DSB formation (3), the facilitative role of Tim in Chk1 regulation under genotoxic conditions raises the possibility that the checkpoint function of Tim is the sole mechanism by which Tim/Tipin suppresses recombination during normal S phase. However, several lines of evidence argue against this possibility. First, although a partial decrease in Chk1 phosphorylation (>50%) has been reported in Tim- and/or Tipin-deficient cells under conditions of genotoxic stress, the basal levels of Chk1 phosphorylation in unchallenged cells either has been shown to increase or not change appreciably (25, 26, 28, 29). Increased Chk1 phosphorylation in Tim-deficient cells is in fact expected, given evidence both in yeast and *Xenopus* extracts that Tim/Tipin deficiency leads to the accumulation of ssDNA during DNA replication (15, 22, 26), which stimulates the ATR-Chk1 pathway (3). Therefore, the function of Tim in preventing DSBs during unperturbed S phase does not appear to be solely linked to assisting Chk1 activation under genotoxic stress.

Evidence both in yeast and vertebrates suggests that ssDNA accumulation resulting from defects in replisome efficiency may be the root cause of genome instability in Tim-reduced cells. For example, using ChIP analysis, it has been shown that
Tof1 is required to maintain coupling between helicase complexes (Cdc45 and MCM7) and newly synthesized daughter strands following HU treatment of S. cerevisiae (13). Deficiencies in coupling were correlated with an expansion of RPA-coated DNA, a hallmark of ssDNA generation. These effects were not observed in mec1tel1sm1l mutants, arguing against checkpoint deficiency being the primary cause of uncoupling (13). Generation of ssDNA has also been observed in swi1 and swi3 mutants, S. pombe orthologs of Tim and Tipin (22), and in Tipin-depleted Xenopus extracts (26). Together these data argue that primary genome stabilizing function of Tim in S phase is to ensure that daughter strand synthesis remains coupled to helicase progression. Loss of this capability alone, or in combination with partial checkpoint reduction, likely contributes to DSB formation in Tim-depleted cells.

Because we observed that the rate of SCE seen in Tim-reduced Blm−/− cells was greater than that observed in cells deficient for either protein alone, our results indicate that Tim deficiency does not lead to increased SCEs through a defect in Blm function. Furthermore, dual deficiency in Tim and Blm results in a decrease in cell proliferation and an increase in apoptosis, consistent with the growth defects observed upon combined mutation of yeast orthologs of Tim (Swi1 and Tof1) with Blm counterparts (Rqh1 and Sgs1) (14, 56). Blm, a RecQ helicase, aids in Holliday junction migration and resolution, which in turn suppress sister chromatid crossover events (46 – 48). Our studies are consistent with the hypothesis that Tim reduction leads to an increased frequency of Holliday junctions, which can undergo Blm-mediated migration and coalescence to suppress SCE (Fig. 6). Previous studies have indicated that the formation of recombination structures in S phase is associated with decreased rates of DNA synthesis (57). Thus, it is possible that increased recombination contributes to the slowed rate of DNA synthesis previously noted in Tim-deficient cells (24, 25, 27, 28).

An important study of Tim and Tipin orthologs in S. pombe has indicated that increased recombination in Swi1 and Swi3 mutant strains occurs independently of DSB generation and Rad51 (15). This study demonstrated that combined deletion of Swi1 and Mus81 (a protein required for Holliday junction resolution) caused the accumulation of unresolved recombination intermediates in a manner that was dependent on Rad22 (mammalian Rad52). Rad52 mutation, but not mutation of Rhp51 or Rhp54 (mammalian Rad51 and Rad54), suppressed swi1 mus81 synthetic lethality. From these data, the authors proposed that the absence of Swi1 during replication leads to replisome uncoupling from newly synthesized DNA, resulting in single strand gaps in the lagging strand that are predominantly repaired by exchange events between the intact parental strands (15). According to this model, fork collapse and DSB formation are averted by the single strand annealing function of Rad22, implying an important specific function in replication fork stability that does not involve Rad51 recruitment (Fig. 6).

Although Rad52 is required for HR in yeast, it is dispensable in mammals, and its loss has little effect on cell viability (31, 41, 42, 58). This difference is largely attributed to the existence of genes that function redundantly with Rad52 in recruiting Rad51 to breaks. These Rad52 analogs include Brca2 and Xrcc3 (41, 43, 59). However, such redundancy in Rad51 recruitment would not exclude a specific role for Rad52 in mediating exchange through single strand annealing, which appears to be conserved in vertebrate cells (60, 61). However, Tim knockdown in Rad52−/− cells had no significant effect on SCE rates or the appearance of chromatid breaks. Thus, unlike in yeast, Tim reduction does not induce a dependence on Rad52 for recombinatorial exchange.

Our results indicate that in mammals Rad52 is not required for SCE or genome maintenance in Tim-deficient cells. Instead, Rad51-mediated SCE is utilized as a means to avert the genomic instability resulting from Tim absence (Fig. 6). Interestingly, our data also indicate a prominent role for Brca2 in this process, consistent with the proposed role for Brca2 as a functional equivalent to yeast Rad52. A Rad51/Brca2-dependent process that averts fork collapse by mechanisms similar to S. pombe should not be excluded. However, given the increase in chromatin breaks and phosphorylation of H2AX in response to Tim deficiency (Figs. 1, 2, and 5), and the known functions of Brca2 and Rad51 in DNA repair, our data suggest that some fraction of SCEs observed in Tim-deficient cells are the product of replication fork collapse into DSBs (Fig. 6). Together, these results indicate that Tim deficiency leads to a reliance on Brca2- and Rad51-mediated repair for genome stability during normal DNA replication.
Although the mechanisms governing SCE are becoming increasingly understood, only a handful of genes that influence this process have been identified (2). One of these genes, Blm, leads to a >150-fold increase in cancer risk when mutated in humans. Importantly, the cytogenetic abnormalities (chromatid breaks, quadriradials, and triradials) and sister chromatid exchange rates observed in Blm null cells (30, 51, 52, 62) are only modestly greater than those observed in Tim-reduced cells (Fig. 3). Thus, it is conceivable that Tim, Tipin, and other yet to be described genes that contribute to resectase efficiency, may be important for suppressing cancer in humans. Furthermore, in light of our studies, it is conceivable that the penetrance of cancer-causing mutations in HR regulators, such as Blm and Brca2, may be strongly influenced by mutation of Tim or Tipin.

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