Histone H2AX stabilizes broken DNA strands to suppress chromosome breaks and translocations during V(D)J recombination

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Histone H2AX stabilizes broken DNA strands to suppress chromosome breaks and translocations during V(D)J recombination

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The H2AX core histone variant is phosphorylated in chromatin around DNA double strand breaks (DSBs) and functions through unknown mechanisms to suppress antigen receptor locus translocations during V(D)J recombination. Formation of chromosomal coding joins and suppression of translocations involves the ataxia telangiectasia mutated and DNA-dependent protein kinase catalytic subunit serine/threonine kinases, each of which phosphorylates H2AX along cleaved antigen receptor loci. Using Abelson transformed pre-B cell lines, we find that H2AX is not required for coding join formation within chromosomal V(D)J recombination substrates. Yet we show that H2AX is phosphorylated along cleaved Igκ DNA strands and prevents their separation in G1 phase cells and their progression into chromosome breaks and translocations after cellular proliferation. We also show that H2AX prevents chromosome breaks emanating from unrepaird RAG endonuclease-generated TCR-α/δ locus coding ends in primary thymocytes. Our data indicate that histone H2AX suppresses translocations during V(D)J recombination by creating chromatin modifications that stabilize disrupted antigen receptor locus DNA strands to prevent their irreversible dissociation. We propose that such H2AX-dependent mechanisms could function at additional chromosomal locations to facilitate the joining of DNA ends generated by other types of DSBs.

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et al., 2002a; Celeste et al., 2002), H2ax−/− cells are defective in the G2/M checkpoint after induction of only a few DSBs (Fernandez-Capetillo et al., 2002). The phenotypes of H2ax−/− cells suggest that the ability of γ-H2AX to retain repair and checkpoint proteins around DSBs may promote accessibility of DNA ends, stabilize disrupted DNA strands, and/or amplify checkpoint signals (Bassing and Alt, 2004; Stucki and Jackson, 2006; Bonner et al., 2008; Kinner et al., 2008).

The health and survival of humans and mice depends on the ability of their adaptive immune systems to generate lymphocytes with receptors capable of recognizing and eliminating large varieties of pathogens. In developing lymphocytes, Ig and TCR variable region exons are assembled from germ-line V (variable), D (diversity), and J (joining) gene segments by the lymphoid-specific RAG1/RAG2 (RAG) endonuclease and the ubiquitously expressed nonhomologous end-joining (NHEJ) DSB repair factors (Bassing et al., 2002b). The RAG proteins catalyze the coupled cleavage of DNA strands between a pair of gene segments and their flanking recombination signal sequences to generate covalently sealed coding ends (CEs) and blunt signal ends (SEs; Fugmann et al., 2000). RAG-mediated cleavage occurs only in G1 phase because of cell cycle phase-restricted expression of RAG2 (Lee and Desiderio, 1999). The DNA-PKcs/Artemis endonuclease opens CEs (Ma et al., 2002), which are then processed by nucleases and polymerases (McEllhinney and Ramsden, 2004). Core NHEJ factors join together CEs and SEs to form coding joints (CJs) and signal joints (SJs), respectively (Bassing et al., 2002b). RAG1/RAG2 can hold CEs and SEs within stable synaptic complexes (Agrawal and Schatz, 1997; Lee et al., 2004); however, ATM and, likely, the Mre11–Rad50–Nbs1 (MRN) complex maintain chromosomal CEs in proximity to facilitate end-joining in G1 phase cells (Bredemeyer et al., 2006; Deriano et al., 2009; Helmark et al., 2009). The large combination of V(D)J joining events and the imprecision in CJ formation cooperate to generate a diverse repertoire of antigen receptor specificities.

Despite its benefits, V(D)J recombination poses substantial threats to the viability and genomic integrity of lymphocytes and lymphoma predisposition in host organisms. For example, DNA-PKcs−/− mice lack mature lymphocytes as a result of inability to repair RAG-generated DSBs, but they only occasionally develop lymphoma (Bosma et al., 1993; Gao et al., 1998; Taccioli et al., 1998). However, DNA-PKcs−/−mice rapidly succumb to pro-B lymphomas with RAG-dependent IgH/c-myc translocations (Barlow et al., 1996, 2002; Bredemeyer et al., 2008), which is consistent with the notion that H2AX does function within chromosomal locations in G1-arrested cells to test our hypothesis of H2AX function in chromosomal end joining during V(D)J recombination. Consequently, we have used a cell line–based system that enables the controlled induction of RAG DSBs at single defined chromosomal locations in G1-arrested cells to test our hypothesized functions of H2AX during V(D)J recombination.

RESULTS
H2AX-deficient cells exhibit normal coding join formation within chromosomal substrates

Although H2AX is not required for coding join formation in extrachromosomal substrates (Bassing et al., 2002a), the phosphorylation of H2AX could function downstream of ATM to facilitate chromosomal coding join formation. To investigate potential H2AX function in chromosomal end joining during V(D)J recombination, we generated multiple independently derived immortalized abl pre−B cell lines from two different H2axf/f mice containing floxed H2ax loci on both alleles.
H2axF/F cells express normal amounts of H2AX and exhibit phenotypes indistinguishable from those of wild-type cells (Bassing et al., 2002a). Treatment of abl pre–B cells with STI571, an inhibitor of the abl kinase, causes G1 arrest, induction of RAG expression, and robust rearrangement of endogenous Igk loci and V(D)J recombination substrates (Muljo and Schlissel, 2003; Bredemeyer et al., 2006). V(D)J recombination of the chromosomally integrated pMX-DEL3 retroviral substrate results in formation of a coding join within the chromosone and generation of a signal join on an extrachromosomal circle (Fig. 1a). We transduced H2axF/F abl pre–B cell lines with the pMX-DEL3 retroviral recombination substrate and used limiting dilution to isolate independent H2axF/F clones with single pMX-DEL3 substrates integrated into their genome (H2axF/F:DEL3 cells). We next incubated individual H2axF/F:DEL3 lines with TAT-Cre protein to delete the floxed H2ax alleles and again used limiting dilution to recover H2axF/F:DEL3 and H2axΔ/Δ:DEL3 clones with pMX-DEL3 substrates integrated at identical genomic locations. The genotypes of these clones were confirmed by both PCR and Southern blot analyses that distinguish between the H2axF/F and H2axΔ/Δ alleles (unpublished data).

To evaluate potential H2AX function in chromosomal end-joining during V(D)J recombination, we sought to monitor the repair of RAG-generated CEs within identical pMX-DEL3 integrants in two different H2axF/F:DEL3 and H2axΔ/Δ:DEL3 clones of the same passage. The induction and repair of RAG-generated DSBs within integrated pMX-DEL3 substrates can be monitored by Southern blot analysis that distinguishes between uncleaved (GL) substrates, cleaved but not repaired (CE) substrates, and cleaved and repaired (CJ) substrates (Fig. 1a). Thus, we conducted Southern blotting of identical pMX-DEL3 integrants in two H2axF/F:DEL3 and H2axΔ/Δ:DEL3 clones treated with STI571 for increasing amounts of time. As a control for the accumulation of unrepaired chromosomal CEs, we also conducted Southern blot analysis of Artemis−/−:DEL3 clones. After 48 h of STI571 treatment, we observed loss of the pMX-DEL3 GL fragment and appearance of the pMX-DEL3 CJ fragment to similar extents in the two H2axF/F:DEL3 and H2axΔ/Δ:DEL3 clones assayed (Fig. 1b). In contrast, we detected loss of the pMX-DEL3 GL fragment and appearance of the pMX-DEL3 CE fragment at 24 h of STI571 treatment in Artemis−/−:DEL3 cells (Fig. 1b). At each time point assayed between 48 and 96 h of STI571 treatment, we observed similar increasing loss of the pMX-DEL3 GL fragment and appearance of the pMX-DEL3 CJ fragment in H2axF/F:DEL3 and H2axΔ/Δ:DEL3 cells (Fig. 1b) but increasing loss of the pMX-DEL3 GL fragment and appearance of the pMX-DEL3 CE fragment in Artemis−/−:DEL3 cells (Fig. 1b). Notably, despite conducting Southern analysis on twice as much genomic DNA for H2axF/F:DEL3 and H2axΔ/Δ:DEL3 cells than for Artemis−/−:DEL3 cells, we were unable to detect the pMX-DEL3 CE fragment in H2axΔ/Δ:DEL3 cells. These data demonstrate that, unlike deficiency in ATM or DNA-PKcs (Bredemeyer et al., 2006), H2AX deficiency does not lead to an observable accumulation of unrepaired coding ends during V(D)J recombination of chromosomal substrates within G1-phase cells.

Because lymphomas with RAG-dependent antigen receptor locus translocations arise at a higher frequency in H2ax−/− p53−/− mice than in H2ax−/− mice (Bassing et al., 2003; Celeste et al., 2003), we considered the possibility that p53 deficiency might reveal a role of H2AX in formation of chromosomal coding joins in G1-phase cells. To investigate this issue, we generated two independently derived abl pre–B cell lines from different H2axF/F mice also containing floxed p53 exons on both alleles. We used TAT-Cre to generate H2axF/F/p53F/F: DEL3 and H2axΔ/Δ/p53Δ/Δ:DEL3 clones of the same passage with pMX-DEL3 substrates integrated at identical genomic locations. Southern blot analysis of identical pMX-DEL3 integrants in two different H2axF/F/p53F/F:DEL3 and H2axΔ/Δ/p53Δ/Δ:DEL3 clones treated with STI571 for increasing amounts of time revealed similar increasing loss of the pMX-DEL3 GL fragment and appearance of the pMX-DEL3 CJ fragment in the two H2axF/F/p53F/F:DEL3 and H2axΔ/Δ/p53Δ/Δ:DEL3 cells (Fig. 1c). Again, despite conducting Southern analysis on twice as much genomic DNA for H2axF/F/p53F/F:DEL3 and H2axΔ/Δ/p53Δ/Δ:DEL3 as for Artemis−/−:DEL3 cells, we did not observe detectable levels of the pMX-DEL3 CE fragment in H2axΔ/Δ/p53Δ/Δ:DEL3 cells (Fig. 1c). These data further support the notion that H2AX is not required for end joining of chromosomal coding ends in G1-phase lymphocytes.

Because RAG-dependent formation of Ig-γH2AX occurs at Jκ segments and over adjacent sequences extending away from the Igk locus (Savic et al., 2009), we also considered that H2AX might be required for resolution of Jκ CEs. The mouse Igk locus resides on chromosome 6 and is composed of 140 Vκs spanning 2 Mb and four functional Jκs spanning 1.8 kb and residing 60 kb from the Vκs. Igk locus V(D)J recombination occurs through the coupled cleavage and subsequent joining of a Vκ and Jκ segment. Because of the small size of the Jκ cluster, Southern blot analysis with a 3′ Jκ probe can be used to monitor and quantify the induction of RAG-generated Igk locus DSBs (Fig. 2a). Thus, we next conducted Southern blotting of H2ax−/− pre–B cells either untreated or treated with STI571 for 72 h. As a control for the accumulation of unrepaired Jκ CEs, we also conducted Southern blot analysis of previously described Artemis−/− cells (Helming et al., 2009). We observed decreased intensity of the Jκ GL band in both H2ax−/− and Artemis−/− cells treated with STI571, and appearance of bands corresponding to Jκ CEs in STI571-treated Artemis−/− cells but not in STI571-treated H2ax−/− cells (Fig. 2a). These data indicate that H2AX deficiency does not result in the detectable accumulation of unrepaired Igk coding ends in G1-phase cells. Consequently, we conclude that H2AX function is not required for chromosomal end joining during V(D)J recombination in G1-phase lymphocytes.

H2AX-dependent chromatin changes prevent separation of RAG-cleaved Igk DNA strands in G1-phase cells

We first sought to define the entire γ-H2AX chromatin domain formed along Igk loci during V(D)J recombination in G1-phase cells. For this purpose, we conducted chromatin
Figure 1. H2AX-deficient cells exhibit normal coding join formation within chromosomal substrates. (a) Shown are schematic diagrams of the pMX-DEL substrate in the uncleaved (GL), cleaved but not repaired (CE), and cleaved and repaired (CJ) configurations. The recombination signal sequences are represented by triangles. Arrows represent the LTR sequences. Indicated are the relative positions of the EcoRV sites (E) and C4 probe used for Southern blot analysis and the sizes of the C4-hybridizing EcoRV fragments in pMX-DEL substrates of the GL, CE, and CJ configurations. (b and c) Southern blot analysis of recombination products generated in cells of two different H2ax F/F :DEL and H2ax F/F :DEL abl pre–B cell lines (b) or H2ax F/F p53 F/F :DEL and H2ax F/F p53 / p53 / :DEL abl pre–B cell lines (c) treated with STI571 for the indicated times. EcoRV-digested genomic DNA was hybridized with the C4 probe. The bands corresponding to pMX-DEL substrates of the GL, CE, and CJ configurations are indicated. Blots were stripped and then probed with a TCR-β probe as a control for DNA content. Artemis F/F :DEL abl pre–B cell lines were used as a positive control for detection of pMX-DEL CEs, with half as much genomic DNA loaded to increase the sensitivity of detection for pMX-DEL CEs in experimental cells. These data are representative of experiments performed more than three independent times.
immunoprecipitation (ChIP) to measure γ-H2AX densities in nucleosomes on DNA strands located within and adjacent to Igκ loci in Artemis<sup>−/−</sup> pre-B cells, either untreated or treated with STI571 for 96 h to ensure complete RAG-mediated cleavage. We used Artemis<sup>−/−</sup> cells rather than wild-type cells because the accumulation of unrepaired coding ends enhances ability to detect γ-H2AX along RAG-cleaved antigen receptor loci (Savic et al., 2009). Using ChIP, we detected significant increases in γ-H2AX densities within Igκ and for ~500 kb on both sides of Igκ in STI571-treated Artemis<sup>−/−</sup> cells.

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**Figure 2.** No accumulation of Jκ coding ends in the absence of H2AX phosphorylation along RAG-cleaved Igκ DNA strands. (a) Shown are schematic diagrams of the Jκ cluster of the Igκ locus in the uncleaved (GL) and cleaved but not repaired (CE) configurations. Open boxes represent the Jκ segments and triangles their recombination signal sequences. Indicated are the relative positions of the EcoRI and SacI sites and 3′ Jκ probe used for Southern blot analysis. (b) Southern blot analysis of recombination products generated in H2ax<sup>−/−</sup> and Artemis<sup>−/−</sup> abl pre-B cell lines, either untreated or treated with STI571 for 72 h. EcoRI–SacI-digested genomic DNA was hybridized with the 3′ Jκ probe. The bands corresponding to Jκ loci of the GL and CE configurations are indicated. The STI571-treated H2ax<sup>−/−</sup> cells harbor a band that likely represents a predominant VκJκ rearrangement. Blots were stripped and then probed with a TCR-β probe as a control for DNA content. These data are representative of experiments performed >10 independent times. (c) Schematic diagram of the mouse Igκ locus and graphical representation of γ-H2AX densities as determined by ChIP at locations along DNA strands within and adjacent to Igκ in Artemis<sup>−/−</sup> abl pre-B cells treated with STI571 for 96 h. The 0-kb value of the x-axis corresponds to the 3′ end of the Jκ5 coding segment. The negative and positive values represent the distances centromeric and telomeric, respectively, from the 3′ end of Jκ5. Red and green bars indicate the approximate genomic locations to which the 5′ Vκ (RP24-243E11) and 3′ Cκ (RP23-341DS) BACs hybridize. The lengths of these bars are not drawn to scale. These data are representative of experiments performed >20 independent times. Error bars indicate standard deviation of three independent experiments.
Figure 3. H2AX suppresses separation of RAG-cleaved Igκ locus DNA strands. (a) Shown are representative fluorescent light microscopy images of 2C-FISH analysis conducted on G1-phase nuclei of Rag2−/−, Artemis−/−, and Artemis+/− H2ax−/− abl pre-B cells treated with STI571 for 96 h. Nuclei were hybridized with the 5′ Vκ (red) and 3′ Cκ (green) BACs and stained with DAPI to visualize DNA. The representative Rag2−/− image shows a nucleus with coincident probe hybridization signals on both Igκ alleles. The top nucleus contains paired Igκ alleles and the bottom unpaired Igκ alleles. The representative Artemis−/− and Artemis+/− H2ax−/− images each shows nuclei with coincident probe signals on both Igκ alleles and paired Igκ alleles (top) or noncoincident probe signals on one Igκ allele, overlapping probe hybridization on the other Igκ allele, and unpaired Igκ alleles (bottom). Bars, ∼3 μm. (b) Shown are representative scatter plots depicting the distances between red and green signals on allele 1 (shorter distance) and allele 2 (longer distance).
pre–B cells (Fig. 2 c). The γ-H2AX densities formed telomeric of the Jk segments were greater than those formed over the Vk cluster (Fig. 2 c). The numbers of Vk and Jk DSBs within our experimental population of cells must be equal because V(D)J recombination proceeds through coupled cleavage. Because the Igk locus contains a cluster of 140 Vk segments spanning 2 Mb and a cluster of four functional Jk segments spanning 1.8 kb, the chromosomal density of Vk cluster DSBs should be less than the chromosomal density of Jk cluster DSBs within our experimental population of cells. Thus, the simplest explanation for the observed γ-H2AX pattern across the Igk locus is that it reflects the density of DSBs induced at the assayed genonic locations. However, these observations are also consistent with the notion that different features of local chromatin environment among the Vk segments and downstream of the Jk segments may influence γ-H2AX dynamics. Regardless, our data demonstrate that a γ-H2AX chromatin domain forms along chromosomal DNA strands containing RAG-cleaved Igk loci for distances extending at least 500 kb from Igk coding ends.

To test our hypothesis that H2AX-mediated chromatin changes function to hold together broken DNA strands, we sought to monitor the positional stability of RAG-cleaved Igk loci in G1-phase cells. We have previously used two-dimensional two-color DNA fluorescent in situ hybridization (FISH [2D–2C–FISH]) with a 5’ Vk BAC (RP24-243E11) and a 3’ Ck BAC (RP23-341D5) probe to monitor the positional stability of RAG-cleaved Igk locus DNA strands in STI571-treated Artemis−/−p53−/− and Artemis−/−p53−/−/H11002 abl pre–B cells (Hewitt et al., 2009). These 5’ Vk and 3’ Ck probes hybridize to genomic sequences located near the ends of γ-H2AX domain that forms along RAG-cleaved Igk loci within STI571 treated pre–B cells (Fig. 2 c). 2D-FISH cannot distinguish between signals that appear on top of each other that are really separate and, therefore, likely underestimates the numbers of cells with unstabilized Igk locus DNA ends. However, because this should be equal across cells of different genotypes, conclusions about different genotypes relative to each other are still possible with 2D–2C–FISH.

We first conducted 2D–2C–FISH using these 5’ Vk and 3’ Ck probes on G1 interphase nuclei prepared from three independent Rag2−/− abl pre–B cells treated with STI571 for 96 h (Fig. 3 a). We measured the distances between 5’ Vk (red) and 3’ Ck (green) signals on both alleles in ~200 nuclei of each cell line assayed, designated the shorter distance from allele 1 and the longer distance from allele 2, and plotted these values onto scatter plots. We observed overlapping or coincident probe hybridization signals (<1 μm apart) on both alleles in >95% of nuclei and noncoincident signals on one allele in <4% of nuclei (Fig. 3 b). Using three independent Artemis−/− cell lines, we observed overlapping or coincident probe signals on both alleles in ~80% of nuclei and noncoincident probe signals on a single allele in ~20% of nuclei (Fig. 3 b). With three independent Artemis−/−H2ax−/− abl cells, we observed overlapping or coincident probe signals on both alleles in ~60% of nuclei and noncoincident probe signals on a single allele in ~40% of nuclei (Fig. 3 b). Although similar levels of unrepaired Igk locus CEs accumulated in all Artemis−/− and Artemis−/−H2ax−/− cells assayed (not depicted), we also normalized the percentage of nuclei with noncoincident probe hybridization signals to the extent of Igk locus cleavage (Fig. 3 c). These data show that RAG-cleaved Igk locus DNA strands physically separate in a significantly higher percentage of Artemis−/−H2ax−/− cells than in Artemis−/− cells. Similar results were obtained using a larger distance (>1.5 μm) to score noncoincident hybridization (Fig. 3 c). Consequently, we conclude that γ-H2AX–mediated chromatin changes suppress physical separation of RAG-cleaved antigen receptor loci in G1-phase cells to prevent their irreversible dissociation or aberrant joining.

H2AX prevents transition of RAG-cleaved DNA strands into chromosome breaks and translocations during cellular proliferation

Molecular characterization of IgH/c-myc translocations in H2ax−/−p53−/− pro–B lymphomas revealed that these lesions occurred after the replication of chromosomes containing unrepaired RAG-initiated IgH locus DNA breaks (Bassing et al., 2003; Celeste et al., 2003). Thus, we hypothesized that γ-H2AX formation along RAG-cleaved antigen receptor loci promotes chromatin changes that hold together broken DNA strands to prevent unrepaired coding ends from transitioning into chromosome breaks and translocations during S phase (Bassing and Alt, 2004). To test our hypothesis, we sought to quantify the frequency of Igk chromosome breaks and translocations in H2ax−/−, Artemis−/−, and Artemis−/−H2ax−/− abl pre–B cell lines treated with STI571 and then released back into cell cycle by STI571 removal. For this purpose, we hybridized chromosome 6-specific paints (red) and conducted FISH with the 5’ Vk and 3’ Ck BACs (green) on 100 or more metaphase spreads prepared from untreated cells or cells that had been treated with STI571 and then released back into cell cycle (Fig. 4 a). We found chromosome 6 chromosome breaks...
or translocations in <1% of metaphases prepared from untreated H2ax−/−, Artemis−/−, or Artemis−/−H2ax−/− cells of three different cell lines for each genotype (Fig. 4, a and b). These data indicate that structural abnormalities of chromosome 6 are not frequent occurrences in abl pre-B cells with deficiencies in H2AX and/or Artemis. We also detected chromosome 6 abnormalities in <1% of metaphases prepared from H2ax−/− or Artemis−/− cells treated and then released from STI571 (Fig. 4, a and b). However, we observed chromosome 6 breaks or translocations involving Igκ in 4–5% of metaphases prepared from Artemis−/−H2ax−/− cells treated and then released from STI571 (Fig. 4, a and b). Southern blot analysis of STI571-treated cells before removal of the STI571 revealed similar levels of unrepaired Igκ locus DSBs in all cells (unpublished data). These data suggest that unrepaired Igκ locus coding ends progress into chromosome breaks and translocations at a significantly higher frequency in Artemis−/−H2ax−/− cells than in Artemis−/− or H2ax−/− cells after STI571 treatment and release. We also found chromosome 6 breaks and translocations involving Igκ in a significantly higher percentage of STI571-treated and released H2ax−/−Artemis−/−/p53−/− abl pre-B cells, as compared with H2ax−/−/p53−/− and Artemis−/−/p53−/− abl pre-B cells (unpublished data). Although these data are consistent with a role of H2AX in promoting chromatin changes that hold together unrepaired coding ends persisting into S phase, potential effects of mutations in cell cycle checkpoints that arise during v-abl-mediated immortalization more frequently or profoundly in H2ax−/− cells than in H2ax+/− cells could contribute to this observation.

H2AX prevents chromosome breaks emanating from unrepaired TCR-α/δ locus coding ends

We next sought to test our hypothesis that H2AX prevents unrepaired coding ends from transitioning into chromosome breaks and translocations during continued cell cycle progression using primary lymphocytes. This approach also enables assessment of this potential H2AX function during a physiological cell cycle rather than during recovery from the prolonged G1 arrest associated with STI571 treatment and release. Although a small fraction of unrepaired coding ends persists into S phase in normal primary thymocytes (Pedraza-Alva et al., 2006), we decided to dramatically increase the percentage of cells with unrepaired coding ends for analyses by using Artemis−/− thymocytes. Because Artemis−/− cells with unrepaired coding ends are eliminated by p53-mediated apoptosis (Rooney et al., 2004), we also needed to use p53 deficiency to allow thymocytes with unrepaired coding ends to survive and proliferate. We first generated germline H2ax−/−/p53−/−, Artemis−/−/p53−/−, and H2ax−/−Artemis−/−/p53−/− mice to test our hypothesis. Unfortunately, H2ax−/−Artemis−/−/p53−/− mice rapidly succumbed to multiple malignancies, preventing analysis of unrepaired coding ends in nonmalignant thymocytes. We reasoned that the specific deletion of H2ax and/or p53 in Artemis−/− thymocytes should prevent early onset of lymphomas and provide the requisite in vivo experimental system. Transgenic mice expressing Cre under control of the proximal Lek promoter (Lek-Cre mice) direct excision of target genes such as floxed p53 (p53fl ) in thymocytes before the initiation of V(D)J recombination (Orban et al., 1992; Lee et al., 2001). Thus, we generated Lek-CreArtemis−/−/p53fl/fl (LAP), Lek-CreH2ax−/−/p53fl/fl (LHP), and Lek-CreArtemis−/−H2ax−/−/p53fl/fl (LAHP) mice. All mice were created heterozygous for Lek-Cre to avoid potential complications associated with transgene integration site. The low cellularity of LAP and LAHP thymocytes posed an insurmountable obstacle for quantitative cytogenetic analyses. Thus, we cultured bone marrow from LHP, LAP, and LAHP mice on OP9-DL1 stromal cells and prepared metaphase spreads from skit+CD25−CD4− CD8− thymocytes in which normal TCR rearrangements occur (Schmitt and Zúñiga-Pflücker, 2002).

In mouse thymocytes, V(D)J recombination occurs at TCR-α/δ loci on chromosome 14, TCR-β loci on chromosome 6, TCR-γ loci on chromosome 13, and, to a limited extent, Igκ loci on chromosome 12. TCR-γ translocations are rarely observed in human and mouse αβ T cells and T lineage lymphomas. Thus, to assay for potential chromosome breaks and translocations initiating from unrepaired RAG-generated coding ends, we hybridized whole chromosome 14, 6, and 12 paints and conducted FISH with BAC probes that hybridize to genomic sequences 5′ or 3′ of TCR-α/δ, TCR-β, or Igκ loci on metaphase spreads prepared from LHP, LAP, or LAHP thymocytes (Fig. 5 a). For this purpose, we prepared metaphase spreads from independent thymocyte cultures from two different 3–5-wk-old mice of each genotype. We analyzed 200 or more metaphases for each chromosome paint and FISH probe set on each culture. Chromosome breaks or translocations involving antigen receptor loci were scored when the 5′ and 3′ FISH probe signals were noncoincident (Fig. 5 a). Through this approach, we did not find any chromosome 12 or Igκ loci breaks or translocations in metaphases prepared from LAHp, LHP, or LAP thymocytes (Fig. 5 a). However, we observed a modest increase in the frequency of chromosome 6 breaks and translocations involving TCR-β loci in metaphases prepared from LAHP thymocytes, as compared with metaphases prepared from LHP or LAP thymocytes (Fig. 5 a and b). All chromosome 14 abnormalities in LAHP cells were replicated TCR-α/δ chromosome breaks and, among the metaphases with these lesions, ~90% contained replicated breaks on a single copy of chromosome 14 (Fig. 5 c). Although we cannot exclude the possibility that these TCR-α/δ chromosome abnormalities represent rare events in a few DN thymocytes that preferentially expand, our data still support a role of H2AX in preventing accumulation of such cells. Based upon these data, we conclude that
H2AX-mediated chromatin changes along RAG-cleaved antigen receptor loci prevent unrepaired coding ends from transitioning into chromosome breaks during continued cell cycle progression.

**DISCUSSION**

The mechanisms by which H2AX suppresses translocations during V(D)J recombination have remained enigmatic since the findings that a significant percentage of $H2ax^{-/-}$ $\alpha\beta$

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**Figure 4.** H2AX prevents transition of RAG-cleaved Igκ DNA strands into chromosome breaks and translocations during cellular proliferation. (a) Shown are representative fluorescent light microscopy images of whole chromosome (Chr.) 6 paints (red) and FISH analysis using the 5¢' Vκ and 3¢' Cκ BACs (both green) and DAPI (blue) to visualize DNA on metaphases prepared from STI571-treated and released Artemis$^{-/-}$ H2ax$^{-/-}$ abl pre-B cells. Below each image is a schematic representation of the observed chromosome 6 configurations. Images show from left to right: two intact chromosome 6; an intact chromosome 6 and a chromosome 6 with an Igκ locus break; an intact chromosome 6 and a chromosome 6 broken into two fragments with one resolved as an Igκ locus translocation; and an intact chromosome 6 and a broken chromosome 6 that resolved as two distinct Igκ locus translocations. Bars, $\sim$1 μm. (b) Shown is a bar graph with quantification of RAG-initiated Igκ chromosome breaks or translocations in metaphases prepared from cells of three independent H2ax$^{-/-}$, Artemis$^{-/-}$, and Artemis$^{-/-}$ H2ax$^{-/-}$ pre-B cell lines released from STI571 treatment. Metaphases prepared from cells of the same lines without STI571 treatment were assayed for Igκ chromosome breaks or translocations to control for potential spontaneous rearrangement and genomic instability accumulated during cell culture. The data represents the percentage of total metaphases analyzed that contained Igκ chromosome abnormalities. The numbers of metaphases assayed to generate the representative data are indicated. 0, Igκ abnormalities were not observed; n.d., not determined. The $p$-values for comparison between cells of the indicated different genotypes are shown. These data were obtained from the same experiment performed three independent times.
Figure 5. H2AX stabilizes broken DNA strands | Yin et al.

(a) Shown are representative fluorescent light microscopy images of whole chromosome paints and FISH analysis of antigen receptor loci on metaphases prepared from OP9 primary thymocytes. The insets contain schematics depicting the chromosome abnormalities within each metaphase. Chromosome 14 paints (green) and FISH with 5’ Vα and 3’ Cα BACs (both red) and DAPI (blue) to visualize DNA are shown in the top. Chromosome 6 paint (red) and FISH with 5’ Vβ and 3’ Cβ BACs (both green) and DAPI (blue) to visualize DNA are shown in the bottom left. Chromosome 12 paint (red) and FISH with a 5’ Vβ BAC (green) and DAPI (blue) to visualize DNA are shown in the bottom right. Bars, ~2 μm. (b) Shown are bar graphs with quantification of chromosome abnormalities involving TCR-α/δ loci (left) or TCR-β loci (right) in metaphases prepared from primary thymocytes of two independent LHP, LAP, and LAHP

LAHP Mouse 1

4 (12.1%) 3 (8.3%

29 33

Break on Only One Allele of Chr. 14

Breaks on Both Alleles of Chr. 14

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T cells contain TCR-α/β locus translocations and H2ax−/− p53−/− mice develop pro-B lymphomas with RAG-initiated IgH/c-myc translocations (Bassing et al., 2003; Celeste et al., 2003). We have demonstrated in this paper that H2AX is not required for the joining of chromosomal coding ends during V(D)J recombination in G1-phase lymphocytes. In this context, we have shown that H2AX deficiency in abl pre-B cells does not lead to accumulation of unrepaird coding ends during the rearrangement of chromosomal substrates, as we previously observed in abl pre-B cells deficient for ATM or DNA-PKcs (Bredemeyer et al., 2006). These data are consistent with the observations that lymphocyte development in H2ax−/− mice is not blocked or impaired at stages in which antigen receptor variable region genes are assembled, as is the case in DNA-PKcs−/− and Atm−/− mice (Bosma et al., 1983; Barlow et al., 1996; Elson et al., 1996; Xu et al., 1996; Gao et al., 1998; Taccioli et al., 1998; Borghesani et al., 2000; Matei et al., 2007; Vacchio et al., 2007). Consequently, we conclude that γ-H2AX formation is not critical for ability of DNA-PKcs and ATM to process and join chromosomal coding ends during variable region gene assembly in G1-phase cells. Although generation of γ-H2AX has been proposed to facilitate DSB repair kinetics by promoting accessibility of broken DNA ends and recruiting repair factors (Rogakou et al., 1998), we did not observe a difference in the kinetics of chromosomal coding join formation between H2ax+/+ and H2ax−/−/abl pre-B cells. However, potential accessibility and repair functions of γ-H2AX might not be evident during V(D)J recombination because initiation of RAG-generated DSBs requires prior opening of chromosomal substrates and the RAG proteins may recruit end-joining factors (Agrawal and Schatz, 1997; Lee et al. 2004) and ATM maintains chromosomal CEs in proximity (Bredemeyer et al., 2006), our finding that Igκ locus strands with unrepaird CEs separated in Artemis−/− cells was unexpected. In this context, ATM alone is not sufficient to prevent disassociation of CEs that are held much less tightly than SEs in the RAG postcleavage complex. However, our observation is in accord with live cell imaging experiments demonstrating mobility of DNA ends at DSBs generated in Ku80-deficient cells (Soutoglou et al., 2007). Our data could reflect baseline mobility of unrepaird Igκ locus DSBs or indicate a role of Artemis in maintaining RAG-generated DNA ends within synaptic complexes, either directly or indirectly through Ku70/Ku80, DNA-PKcs, and/or other end-binding factors. Our finding that H2AX suppresses the physical separation of broken Igκ locus strands appears in contrast with previously published experiments demonstrating a role of Ku80, but not H2AX, in promoting immobility of DSBs (Soutoglou et al., 2007). Upon DSB induction, more unrepaird DNA ends should accumulate in Ku80−/− cells as compared with H2ax−/− cells as a result of the differential functions of these proteins in end joining. In our study, we used Artemis deficieny to compare the positional stability of disrupted chromosomal DNA strands between H2ax+/+ and H2ax−/− cells that have accumulated equivalent levels of unrepaird DSBs. Thus, we suggest that the disparte conclusions regarding the role of H2AX in maintaining broken chromosomal DNA ends in proximity is the result of a greater level of unrepaird DNA ends that can physically separate in cells deficient for Ku80 versus H2AX.
Although V(D)J recombination occurs in G1-phase cells, some fraction of developing lymphocytes with unrepaired CEs progress into S phase (Rooney et al., 2004). We have shown here that H2AX prevents accumulation of chromosome breaks from RAG-initiated antigen receptor locus DNA breaks that are not repaired before DNA replication. In this context, we have demonstrated that H2AX deficiency in Artemis/p53-deficient thymocytes leads to a substantial increase in the percentage of cells with replicated TCR-α/δ chromosome breaks. Replication through hairpin-sealed CEs should lead to either dicentric or ring chromosomes in metaphase cells. The absence of such chromosome abnormalities in metaphases prepared from LAP or LAHP thymocytes suggests that TCR-α/δ CEs persisting into S phase are opened before DNA replication. Detection of replicated TCR-α/δ chromosome breaks in LAHP, but not LAP, metaphases indicates that H2AX holds together DNA strands with unrepaired TCR-α/δ CEs to facilitate end-joining and/or activates the G2/M checkpoint to prevent mitosis. ATM similarly prevents the continued proliferation of lymphocytes with unrepaired RAG-generated coding ends (Callén et al., 2007). Thus, we conclude that ATM-mediated H2AX-dependent chromatin changes along RAG-cleaved antigen receptor loci prevent unrepaired coding ends from transitioning into chromosome breaks during continued cell cycle progression.

The data presented here further contribute to understanding the mechanisms by which chromosomal DSBs are repaired through NHEJ in G1-phase mammalian cells (Fig. 6). Our findings that H2AX, but not ATM or the MRN complex (Bredemeyer et al., 2006; Helmink et al., 2009), is dispensable for end joining of chromosomal CEs in G1 phase cells is consistent with the lower frequency of interlocus V(D)J recombination events and the milder lymphopenia of H2ax−/− mice, as compared with Atm−/−, Nbs1−/−, and Mre11 ΔΔLD/ΔΔLD mice (Kang et al., 2002; Theunissen et al., 2003). ATM (Bredemeyer et al., 2006), and likely MRN (Deriano et al., 2009; Helmink et al., 2009), facilitates end joining by maintaining chromosomal CEs within RAG postcleavage synaptic complexes, which engage the proximal ends of cleaved DNA strands. Our observations that γ-H2AX densities are low near Jk CEs, but elevated over sequences extending at least 500 kb on both sides of Igk CEs, indicates that γ-H2AX formation stabilizes broken DNA strands at locations distal to breakage sites (Bassing and Alt, 2004). During V(D)J recombination, this ATM-dependent formation of γ-H2AX along RAG-cleaved DNA strands (Savic et al., 2009) is dispensable for end joining of chromosomal CEs in G1 phase cells as a result of cooperation between the DNA end synaptic functions of RAG1/RAG2 and ATM/MRN. In contrast, the ATM-dependent generation of γ-H2AX along DNA strands would
be more critical for the NHEJ-mediated repair of DSBs generated by genotoxic insults, such as ionizing radiation, or nucleases lacking DNA end synaptic functions, such as those that initiate IgH locus CSR (Fig. 6).

MATERIALS AND METHODS

Mice. All mice were bred and maintained under specific pathogen-free conditions at the Children’s Hospital of Philadelphia and were handled according to Institutional Animal Care and Use Committee guidelines. The Institutional Animal Care and Use Committee of the Children’s Hospital of Philadelphia approved all of the animal experiments.

Generation and culture of abl pre-B cell lines. Bone marrow from 3–5-wk-old H2ax/G−/−, H2ax/F−/−, Artemis−/−/H2ax/F−/−, Artemis−/−, p53/F−/−, Lck-Cre transgene, and Eμ-Bcl-2 transgene mice have been previously described (Strasser et al., 1991; Jonkers et al., 2001; Lee et al., 2001; Bassing et al., 2002a; Bussing et al., 2003; Rooney et al., 2002).

Southern blot analysis. Southern blot analyses were performed on genomic DNA using the indicated restriction enzymes and the C4 probe for pMX-DEJ CJ Southern blot analysis.

STI571 and cultured for experiments, cells were collected, washed, and then plated into media without cell lines with single pMX-DEJ CJ integrants. Cells were treated with 3 μM dilution into 96-well plates and Southern blot analysis was used to isolate clonal recombination substrate by cocentrifugation at 1,800 rpm for 90 min. Limiting dilution into 96-well plates and Southern blot analysis was used to isolate clonal cell lines with single pMX-DEF3 integrants. Cells were treated with 3 μM STIS71 for the indicated times at a density of 106 cells/ml. For STIS71 release experiments, cells were collected, washed, and then plated into media without STIS71 and cultured for ~10-14 d before metaphase preparation.

Southern blot analysis. Southern blot analyses were performed on genomic DNA using the indicated restriction enzymes and the C4 probe for pMX-DEL3, the 3’ Jκ probe for Igκ loci, and a TCR-β VDJ probe for loading control (Bassing et al., 2000; Bredemeyer et al., 2006; Helminik et al., 2009). The extent of Jκ cleavage was calculated by the following formula:

\[
\text{Percentage of Jκ cleavage} = \left( 1 - \frac{\text{Igκ GL}^\text{STI} \text{ VDJ} \text{ GL}^\text{STI}}{\text{TCR-β GL}^\text{STI} \text{ VDJ} \text{ GL}^\text{STI}} \right) \times 100\% 
\]

where the Igκ germline hybridization intensity using the 3’ Jκ probe before or after STIS71 treatment [Igκ GL STI and Igκ GL STI, respectively] was normalized to the corresponding TCR-β germline hybridization intensity using the VDJ probe (TCR-β GL STI and TCR-β GL STI, respectively).

ChIP. All data points represent mean values with standard deviation error bars obtained from experiments conducted three times on cells of two independently derived Artemis−/− pre-B cell lines. ChIP assays were conducted and data analyzed exactly as described previously (Savic et al., 2009). The sequences of the primers used for quantitative PCR analysis of each genomic location relative to Jκ1 were either described previously (Savic et al., 2009) or are as follows: −250 kb, 5’-CCATCATCAGAAGCCCGAATTGATT-3’ and 5’-GCTCCAATAGCATACGTCTTCATT-3’; −660 kb, 5’-AATGCGCGCTTGGCTTGGTCTCTA-3’, 5’-ATTGGC- CATCTTACAGCTGAAGAG-3’, 5’-GGTGGTTGTCCTTTCAAGT-3’, and 5’-ACGCAAGGATTCCATGGAGCT-3’. Cyto genetic analyses. Kits for spectral karyotyping and whole chromosome painting were used for metaphase analysis according to the manufacturer’s instructions (Applied Spectral Imaging). BAC FISH probes were labeled with Biotin (Biotin-Nick Translation Mix; Roche), and were applied to G1-phase nuclei or metaphase spreads prepared as previously described (Hewitt et al., 2009). The 5′ Vv (RP24-243E11) and 3′ Cc (RP23-341D5) BACs used for 2C-FISH analysis were previously described (Hewitt et al., 2009). The TCR-β-164G11, TCR-Vβ17a-23N16, TCR-Co-232F19, TCR-Vβ6/66-46G9, and 5′ IgH BAC207 BACs used for FISH analysis of translocations in primary thymocytes have been previously described (Liyang et al., 2000; Franco et al., 2006). All images were captured and analyzed on a spectral karyotyping workstation using Case Data Manager Version 5.5, installed and configured by Applied Spectral Imaging. The 2C-FISH experiments to assay for DSB positional stability were conducted independently on three different lines of Rag2−/−, Artemis−/−, and Artemis−/−/H2ax−/− abl pre-B cells. To avoid observer bias, these data of these experiments were conducted and obtained blind of the genotypes, with the exception of one Artemis−/− line that was not scored blind.

OP9-DL1 co-culture. Bone marrow cells from 4–6-wk-old mice were harvested and red blood cell lysed. Progenitor cells were enriched using MACS purification columns (Miltenyi Biotec) and antibodies specific for mature hematopoietic lineages (CD24, B220, Mac-1, Gr-1, NK1.1, CD11c, and Thy1.2). Approximately 2 million cells were cultured per well using 24-well plates. Cultures were otherwise treated as previously described (Schnitt and Zülgica-Pflücker, 2002), except for supplementation with 5 ng/ml of mouse IL-7 and human Flt-3L (R&D Systems).

Statistical analysis. All statistical analysis was performed with two-tailed unpaired Student’s t tests assuming equal variation in Excel (Microsoft). P < 0.05 was considered to be statistically significant.

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