2009

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Recommended Citation
Helmink, Beth A.; Bredemeyer, Andrea L.; Lee, Baeck-Seung; Huang, Ching-Yu; Sharma, Girdhar G.; Walker, Laura M.; Bednarski, Jeffrey J.; Lee, Wan-Ling; Pandita, Tej K.; Bassing, Craig H.; and Sleckman, Barry P., "MRN complex function in the repair of chromosomal Rag-mediated DNA double-strand breaks." Journal of Experimental Medicine. 206, 3. 669 - 679. (2009).
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MRN complex function in the repair of chromosomal Rag-mediated DNA double-strand breaks

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The Mre11–Rad50–Nbs1 (MRN) complex functions in the repair of DNA double-strand breaks (DSBs) by homologous recombination (HR) at postreplicative stages of the cell cycle. During HR, the MRN complex functions directly in the repair of DNA DSBs and in the initiation of DSB responses through activation of the ataxia telangiectasia-mutated (ATM) serine-threonine kinase. Whether MRN functions in DNA damage responses before DNA replication in G0/G1 phase cells has been less clear. In developing G1-phase lymphocytes, DNA DSBs are generated by the Rag endonuclease and repaired during the assembly of antigen receptor genes by the process of V(D)J recombination. Mice and humans deficient in MRN function exhibit lymphoid phenotypes that are suggestive of defects in V(D)J recombination. We show that during V(D)J recombination, MRN deficiency leads to the aberrant joining of Rag DSBs and to the accumulation of unrepaired coding ends, thus establishing a functional role for MRN in the repair of Rag-mediated DNA DSBs. Moreover, these defects in V(D)J recombination are remarkably similar to those observed in ATM-deficient lymphocytes, suggesting that ATM and MRN function in the same DNA DSB response pathways during lymphocyte antigen receptor gene assembly.

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activated in an MRN-independent fashion in response to some DSBs like those generated at stalled replication forks, suggesting that the requirement for MRN in initiating DSB responses may be context dependent (9). MRN also has several DNA damage response functions that are downstream of ATM activation. In this regard, analyses of Nbs1 mutants have implicated MRN in the regulation of cell cycle checkpoints and the activation of apoptotic pathways in response to DNA DSBs (13, 14). Mre11 has endonuclease and exonuclease activities that are important for DNA end processing, and Mre11 dimers may align and bridge two DNA ends during HR-mediated DSB repair (5, 6, 15). Rad50 also has DNA binding activities that may be involved in tethering sister chromatids during HR (16–20). In response to DSBs, ATM phosphorylates Mre11, Rad50, and Nbs1, which could potentially modulate their functions in DSB responses (21–24). Importantly, although Mre11, Nbs1, and Rad50 have distinct functions, these individual components are thought to function only in the context of the MRN holocomplex.

Whether the MRN complex functions in the response to and NHEJ-mediated repair of DSBs in G0/G1-phase cells has been much less clear. DNA end joining by NHEJ components purified from human cellular extracts was augmented by MRN in vitro; however, the repair activity of purified Xenopus laevis NHEJ components was not affected by the addition of MRN (25, 26). MRX, the yeast orthologue of MRN, functions during NHEJ in the budding yeast Saccharomyces cerevisiae but not in the fission yeast Schizosaccharomyces pombe (27–29). In mammalian cells, MRN is not recruited to the site of DSBs generated by the I-sceI endonuclease in G1-phase cells, and MRN does not appear to be required for the NHEJ-mediated repair of these DSBs (30, 31).

DSBs are generated in all developing lymphocytes during the assembly of the second exon of antigen receptor genes from component V, J, and, in some cases, D gene segments (32). This occurs through the process of V(D)J recombination, which is initiated by the Rag-1 and Rag-2 proteins, which together form an endonuclease, hereafter referred to as Rag (33, 34). Rag introduces DSBs at the borders of two recombining gene segments and their associated Rag recognition sequences, which are termed recombination signals (RSs). The generation of these DSBs is restricted to cells at the G1 phase of the cell cycle as a result of the rapid degradation of Rag-2 upon entry into S phase (35). DNA cleavage by Rag leads to the formation of two hairpin-sealed coding ends and two blunt phosphorylated signal ends. These DNA ends are processed and joined by the NHEJ pathway of DNA DSB repair into a coding joint and signal joint, respectively (36, 37). The critical dependence of V(D)J recombination on NHEJ is indicated by the severe joining defects in NHEJ-deficient cell lines and the profound immunodeficiency observed in mice deficient for NHEJ factors required for the repair of Rag-mediated DNA DSBs (36).

Mice with homozygous-null mutations in the Mre11, Nbs1, or Rad50 genes exhibit early embryonic lethality; however, mice and humans with hypomorphic Mre11 or Nbs1 mutations are viable and exhibit mild immunodeficiency, suggesting that the MRN complex could function in the response to or repair of Rag-mediated DSBs generated during V(D)J recombination (9, 38–45). In developing lymphocytes, Rag DSBs activate ATM, which initiates a broad genetic program and functions in the repair of these breaks (46–53). In this regard, MRN could function in the activation of ATM in response to Rag-mediated DSBs and also downstream of ATM in the repair of these DSBs. Consistent with this notion, Nbs1 associates with Rag-mediated DSBs generated at T cell receptor loci in thymocytes and MRX function is required for the joining of signal ends generated by Rag cleavage in yeast (54, 55). However, analyses of V(D)J recombination of extrachromosomal substrates in mammalian nonlymphoid cells deficient in MRN have failed to reveal any significant defects in coding or signal joint formation (56–58). In this paper, we show that MRN-deficient lymphocytes exhibit defects in V(D)J recombination at endogenous antigen receptor loci and chromosomally introduced recombination substrates. These findings establish a function for MRN in the response to Rag-mediated DSBs generated in G1-phase lymphocytes.

**RESULTS**

**Generation of Nbs1<sup>m/m</sup> and Mre11<sup>ATLD1/ATLD1</sup> abl pre–B cells**

We have developed an approach for analyzing the response to and repair of chromosomal Rag-mediated DSBs in v-abl-transformed pre–B cell lines, which are hereafter referred to as abl pre–B cells (48–50). Treatment of these cells with the v-abl kinase inhibitor STI571 leads to G1 cell cycle arrest, induction of Rag expression, and the generation of Rag DSBs at the endogenous IgL-κ locus and at chromosomally introduced retroviral recombination substrates (48–50, 59). In WT G1-phase abl pre–B cells these DSBs are rapidly repaired, whereas in cells deficient in specific DNA repair proteins these DSBs can persist unrepaired for substantial periods of time (48–50).

Mice with homozygous-null mutations at the Mre11, Nbs1, or Rad50 loci exhibit early embryonic lethality, preventing the generation of abl pre–B cells that are completely deficient of any of these proteins (9, 43–45). However, mice that express hypomorphic Nbs1 or Mre11 alleles are viable (9, 38–40). We generated several abl pre–B cell lines from mice homozygous for hypomorphic Mre11 (Mre11<sup>ATLD1</sup>) and Nbs1 (Nbs1<sup>+</sup>) alleles (38, 40). The Mre11<sup>ATLD1</sup> allele has a gene-targeted point mutation that generates a premature stop codon, resulting in a C-terminal truncation of the Mre11 protein, which mimics a mutation that causes ataxia-telangiectasia–like disease (40). Exons 2 and 3 of the Nbs1 allele were replaced with a neomycin resistance gene generating the Nbs1<sup>+</sup> allele, which encodes a truncated Nbs1 protein (38).

Independently derived WT, Atm<sup>−/−</sup>, Nbs1<sup>m/m</sup>, and Mre11<sup>ATLD1/ATLD1</sup> (referred to as Mre11<sup>A/A</sup> in figures) abl pre–B cell lines were transduced with the pMX-INV retroviral recombination substrate (Fig. 1a). The RSs in pMX-INV are oriented such that rearrangement occurs by inversion, with the coding and signal joints remaining in the chromosome...
(Fig. 1 a). Bulk populations of abl pre–B cells with pMX-INV integrants were purified by flow cytometric cell sorting for the expression of human CD4, which is encoded by the retrovirus (50). Abl pre–B cells with single pMX-INV integrants were isolated by limiting dilution. We analyze pMX-INV rearrangement both in clones with single pMX-INV integrants (Fig. 1, b and c; and Fig. S1, available at http://www.jem.org/cgi/content/full/jem.20081326/DC1) and in original bulk populations (Fig. 1, d and e) of abl pre–B cells that have pMX-INV integrants at broadly heterogeneous genomic locations (50).

Atypical joining of Rag-mediated DSBs in MRN-deficient abl pre–B cells

Induction of Rag expression leads to robust pMX-INV rearrangement in Nbs1+/−:INV and Mre11ΔTLD1/ΔTLD1:INV abl pre–B cell clones (Fig. 1, b and c; and Fig. S1). Strikingly, pMX-INV hybrid joints form at high levels in both Nbs1+/−:INV and Mre11ΔTLD1/ΔTLD1:INV abl pre–B cell clones (Fig. 1, b and c; and Fig. S1). Hybrid joints are atypical nonproductive rearrangements formed by the ligation of a signal end generated at one Rag DSB to the coding end generated at the other (Fig. 1 a) (60, 61). pMX-INV hybrid joint formation in Nbs1+/−:INV and Mre11ΔTLD1/ΔTLD1:INV abl pre–B cell clones occurs at levels approximating those observed in Atm−/−:INV clones (Fig. 1, b and c; and Fig. S1). In contrast, Southern blot analyses did not reveal pMX-INV hybrid joint formation in WT abl pre–B cell clones (WT:INV; Fig. 1, b and c; and Fig. S1). Similar results were observed when analyzing rearrangement in bulk populations of abl pre–B cells with pMX-INV integrants at numerous heterogeneous chromosomal locations (Fig. 1, d–f; and Fig. S2). As was observed in Atm−/− abl pre–B cells, the increase in hybrid joint formation in Nbs1+/− and Mre11ΔTLD1/ΔTLD1 abl pre–B cells occurs during rearrangements by inversion but not during rearrangements by deletion (unpublished data) (50).

Coding joint formation in MRN-deficient abl pre–B cells

To assess the impact of MRN during signal joint formation, we introduced the pMX-DEL3 retroviral recombination substrate into Nbs1+/−:INV and Mre11ΔTLD1/ΔTLD1 (Mre11ΔTLD1/ΔTLD1:DEL3) abl pre–B cells (Fig. 2 a). pMX-DEL3 is a derivative of pMX-INV in which one RS has been inverted so that rearrangement occurs by deletion with the signal joint remaining in the chromosome (Fig. 2 a) (50). As was previously observed in Atm−/−:DEL3 abl pre–B cells, induction of Rag expression in these cells leads to robust pMX-DEL3 signal joint formation with no detectable unrepaired signal ends (Fig. 2 b) (50). In contrast, unrepaired signal ends were readily detected after Rag induction in Ku70-deficient abl pre–B cells (Ku70−/−:DEL3), as would be expected given the requirement for Ku70 in signal joint formation (Fig. 2 b). The majority of pMX-DEL3 signal joint PCR products amplified from Nbs1+/−:DEL3, Mre11ΔTLD1/ΔTLD1:DEL3, WT:DEL3, and Atm−/−:DEL3 abl pre–B cells can be digested with ApaLI, which is indicative of precise formation (Fig. 2 c). Indeed, sequence analyses confirmed that most of the signal joints were generated by the precise joining of two signal ends; however, a small fraction of imprecise joints were present in all cells (Fig. 2 c and Fig. S4, available at http://www.jem.org/cgi/content/full/jem.20081326/DC1). Together, these findings demonstrate that the compromised MRN function in the Nbs1+/− and Mre11ΔTLD1/ΔTLD1 abl pre–B cells does not lead to substantial quantitative or qualitative defects in signal joint formation.

ATM activation in response to Rag DSBs in MRN-deficient abl pre–B cells

To determine whether ATM is activated in response to Rag DSBs in MRN-deficient abl pre–B cells, we generated several independently derived Mre11ΔTLD1/ΔTLD1 and Nbs1+/− abl pre–B cells that are also deficient in Artemis (Mre11ΔTLD1/ΔTLD1:Artemis−/− and Nbs1+/−:Artemis−/− abl pre–B cells, respectively). As we have previously demonstrated, induction of Rag in Artemis−/− abl pre–B cells leads to the accumulation of unrepaired coding ends at the endogenous IgL-κ locus as a result of the function of Artemis in opening hairpin-sealed DNA ends (Fig. 3 a) (48, 49, 62). These unrepaired coding
Figure 1. Rearrangement of pMX-INV in Nbs1<sup>m/m</sup> and Mre11<sup>ATLD1/ATLD1</sup> abl pre-B cells. (a) Schematic of the pMX-INV retroviral recombination substrate, rearrangement intermediates, and products. The retroviral packaging signal (φ), GFP complementary DNA, and IRES-human CD4 (I-hCD4)
ends activate ATM, as indicated by the phosphorylation of
KAP-1 and H2AX and the nuclear translocation of NF-kB in
response to these DSBs in Artemis−/−, but not Artemis−/−:Artemis−/−, abl pre-B cells (Fig. 3 and Fig. 4 c) (48, 63).

The induction of Rag DSBs in Mre11ATLD1/ATLD1:Artemis−/− and Nbs1 m/m:Artemis−/− abl pre-B cells led to robust but variable levels of KAP-1 phosphorylation (Fig. 4 a and Fig. S5, available at http://www.jem.org/cgi/content/full/jem.20081326/DC1). The variability in KAP-1 phosphorylation in these cells does not appear to be a result of differences in the levels of KAP-1, Nbs1, or Mre11 protein but may, in part, reflect differences in the levels of unrepaired coding ends generated in these cells after Rag induction (Fig. 4 a and Figs. S5–S7). Importantly, KAP-1 phosphorylation in Artemis−/−, Mre11ATLD1/ATLD1:Artemis−/−, and Nbs1 m/m:Artemis−/− abl pre-B cells is nearly completely abrogated by the Atm inhibitor KU-55933, demonstrating that ATM is activated by Rag DSBs in all of these cells (Fig. 4 a and Fig. S5). Moreover, Rag DSBs promote an Atm-dependent phosphorylation of H2AX and nuclear translocation of NF-kB in Mre11ATLD1/ATLD1:Artemis−/− and Nbs1 m/m:Artemis−/− abl pre-B cells (Fig. 4 b and c; Fig. S8; and not depicted). Finally, treatment of Mre11ATLD1/ATLD1:INV and Nbs1 m/m:INV abl pre-B cells with KU-55933 results in a significant increase in the accumulation of unrepaired pMX-INV coding ends (Fig. 5). Collectively, our findings demonstrate that ATM is activated by Rag DSBs in Mre11ATLD1/ATLD1 and Nbs1 m/m abl pre-B cells.

Defects in V(DJ) recombination in developing MRN-deficient lymphocytes

We have demonstrated that deficiencies in MRN function lead to defects in V(DJ) recombination in Mre11ATLD1/ATLD1 and Nbs1 m/m abl pre-B cells. To determine whether similar defects occur in developing MRN-deficient lymphocytes, we first analyzed thymocyte development in Nbs1 m/m mice. Nbs1 m/m mice have a mild reduction in total thymocyte number but a significant reduction in CD4+ and CD8+ (single positive) thymocytes (Fig. 6 a) (38). In addition, they have diminished numbers of preselection βα−TcR−expressing (TcR-βα [TcR-β intermediate]) CD4+/CD8+ (double positive [DP]) thymocytes (Fig. 6 b). Defects in thymocytes that are similar in nature but more severe are observed in Atm−/− mice (Fig. 6) (9, 46, 51, 52, 64). The reduction in TcR-βα DP thymocytes in Nbs1 m/m mice could be a result of defects in the survival of these cells or in the developmental transition of thymocytes through this compartment. However, the reduction in TcR-βα DP thymocytes in Nbs1 m/m mice could also be explained by defects in TcR-α chain gene assembly as have been observed previously in Atm−/− thymocytes (46, 51, 52).

To directly determine whether MRN deficiency leads to defects in the repair of Rag-mediated DSBs in developing lymphocytes in vivo, we generated Mre11ATLD1/ATLD1 and Nbs1 m/m mice homozygous for a modified TcR-α allele, TcR-αβ (Mre11ATLD1/ATLD1:TcR-αβ and Nbs1 m/m:TcR-αβ). The TcR-αβ allele is identical to the TcR-α allele, except that the 61 Jα gene segments have been replaced by two closely linked Jα gene segments (Jα61 and Jα56) (65). As all Vα rearrangements must use one of these two Jα gene segments, accumulation of unrepaired Rag-mediated DSBs generated at these Jα gene segments can be readily detected by Southern blotting of genomic DNA from developing TcR-αβ thymocytes (46).

Genomic DNA isolated from WT:TcR-αβ, Atm−/−:TcR-αβ, Nbs1 m/m:TcR-αβ, and Mre11ATLD1/ATLD1:TcR-αβ thymocytes was digested with StuI and hybridized to the Cox1 probe, which is directed to the TcR-α constant region gene immediately downstream of the Jα56 gene segment on the TcR-αβ allele (Fig. 7, a and b). These analyses revealed many nongermline-sized hybridizing fragments as a result of diverse Vα to Jα rearrangements in thymocytes from all of these mice (Fig. 7 b). In addition, the Atm−/−:TcR-αβ, Nbs1 m/m:TcR-αβ, and Mre11ATLD1/ATLD1:TcR-αβ thymocytes each have a distinct 5.9-kb band, which was not present in WT:TcR-αβ thymocytes, that we have previously demonstrated is generated by unrepaired Jα56 coding ends (Fig. 7 b) (46). The presence of unrepaired Jα56 coding ends in Nbs1 m/m:TcR-αβ and Mre11ATLD1/ATLD1:TcR-αβ thymocytes was confirmed by ligation-mediated PCR (unpublished data). Notably, as was observed for pMX-INV coding ends in the abl pre-B cells, the level of unrepaired Jα56 coding ends was generally lower in the Nbs1 m/m:TcR-αβ and Mre11ATLD1/ATLD1:TcR-αβ thymocytes, as compared with the Atm−/−:TcR-αβ thymocytes (Fig. 7 b). Unrepaired Jα56 signal ends do not accumulate at higher levels in Atm−/−:TcR-αβ, Nbs1 m/m:TcR-αβ, and
Like MRN-deficient abl pre–B cells, \( \text{Mre11 ATLD1/ATLD1} \) and \( \text{Nbs1}^{m/m} \) thymocytes also exhibited an increase in hybrid joint formation during rearrangements that occur by inversion as compared with WT:TCR-\( \alpha^{d/d} \) thymocytes (unpublished data) (46).

In this regard, we find that hybrid joints involving the VB14 gene segment (TCR-\( \beta \) locus) and the V\( \delta5 \) gene segment (TCR-\( \delta \) locus), which both rearrange by inversion, are increased in \( \text{Mre11 ATLD1/ATLD1} \) and \( \text{Nbs1}^{m/m} \) thymocytes as compared with WT thymocytes (Fig. 7 c). Furthermore, the level of VB14 and V\( \delta5 \) hybrid joints is similar to that observed in \( \text{Atm}^{-/-} \) thymocytes (Fig. 7 c). The increase in hybrid joint formation during rearrangements that occur by inversion was not restricted to MRN-deficient T cells as it is also observed during IgL-\( \kappa \) locus rearrangements in \( \text{Mre11 ATLD1/ATLD1} \) and \( \text{Nbs1}^{m/m} \) B cells (Fig. 7 d). Together, these findings demonstrate that developing MRN-deficient lymphocytes exhibit defects in V(D)J recombination similar to those observed in Atm-deficient lymphocytes, which are characterized by an accumulation of unrepaired coding ends.

Figure 2. Signal joint formation in MRN-deficient abl pre–B cells. (a) Schematic of the pMX-DEL\(^{SJ} \) retroviral recombination substrate. All components of the retrovirus are as indicated in the Fig. 1 legend. The signal end (SE) and signal joint (SJ) are indicated. (b) Southern blot analysis of EcoRV-digested genomic DNA from \( \text{Nbs1}^{m/m} \) (line 675.3), \( \text{Mre11 ATLD1/ATLD1} \) (line 48.1), and \( \text{Ku70}^{-/-} \) (line 0.2) abl pre–B cells containing the pMX-DEL\(^{SJ} \) (DEL\(^{SJ} \)) retroviral recombination substrate that had been treated with STI571 for the indicated number of hours. (c) PCR analysis of pMX-DEL\(^{SJ} \) signal joints generated in WT:DEL\(^{SJ} \), \( \text{Atm}^{-/-}:\text{DEL}^{SJ} \), \( \text{Mre11 ATLD1/ATLD1}:\text{DEL}^{SJ} \), and \( \text{Nbs1}^{m/m}:\text{DEL}^{SJ} \) abl pre–B cells treated with STI571 for 72 h. PCR was performed using primers \( \text{pB} \) and \( \text{pC} \) (a), and PCR products were either not digested (–) or digested with ApaLI (+). Products were visualized by ethidium bromide staining, and undigested (SJ) and digested (CP) signal joint products are indicated. The data presented are representative of at least two experiments.

Figure 3. Atm activation by Rag DSBs. (a) Southern blot analysis showing Jk1 and Jk2 coding ends (CE) generated in Rag\(^{2-/-} \), Artemis\(^{-/-} \), and Artemis\(^{-/-}:\text{Atm}^{-/-} \) abl pre–B cells treated with STI571 for the indicated times (hours). Sacl- and EcoRI-digested genomic DNA was hybridized to the JkII probe. The bands corresponding to the IgL-\( \kappa \) locus in the germline configuration (GL) and Jk1 and Jk2 CEs are indicated. (b) Western blot analysis of phospho-KAP-1 (\( \alpha\)-p-KAP-1) and KAP-1 (\( \alpha\)-KAP-1) from STI571-treated cells shown in a. (c) Quantification of \( \gamma\)-H2AX nuclear foci in the cells from a after 24 h of STI571 treatment. Shown is the percentage of cells containing one, two, or three to five \( \gamma\)-H2AX foci. The total number of cells analyzed for each genotype was 500 and p-values were calculated using a two-tailed Fisher’s exact test. Note that \( \gamma\)-H2AX foci were not detected in any of the STI571-treated Rag\(^{2-/-} \) abl pre–B cells. The data presented are representative of at least two experiments.
and an increase in hybrid joint formation during rearrangements by inversion.

DISCUSSION

In this paper, we have demonstrated that the MRN complex functions in the repair of chromosomal Rag-mediated DNA DSBs generated during V(D)J recombination in cultured pre–B cell lines and in developing lymphocytes. This is indicated by the accumulation of unrepaired coding ends and the increase in aberrant hybrid joint formation during rearrangements by inversion.

The V(D)J recombination defects observed in MRN-deficient lymphocytes are remarkably similar to those observed in ATM-deficient lymphocytes, suggesting that MRN and ATM function in the same pathway during the repair of Rag-mediated DSBs (50). Although the defects in V(D)J recombination and an increase in hybrid joint formation during rearrangements by inversion.

Figure 4. Rag-DSB-mediated Atm activation in MRN-deficient cells. (a) Western blot analysis for phospho–KAP-1 (α–p-KAP-1) and KAP-1 (α–KAP-1) in Artemis−/− (line 5.1), Nbs1+/−Artemis−/− (line 37.1), and Mre11ATLD1/ATLD1:Artemis−/− (line 64.1) abl pre–B cell lines treated with STI571 for 96 h and either DMSO (−) or the Atm inhibitor (iAtm) KU-55933 (+). (b) Percentage of Artemis−/−, Nbs1+/−, and Mre11ATLD1/ATLD1:Artemis−/− abl pre–B cells with one, two, or three nuclear γ-H2AX foci after treatment with STI571 for 24 h and either DMSO (−) or the Atm inhibitor (iAtm) KU-55933 (+). The total number of cells analyzed for each genotype was 500 and p-values were calculated using a two-tailed Fisher’s Exact test. (c) NF-κB EMSA of nuclear lysates from Artemis−/−, Mre11ATLD1/ATLD1:Artemis−/−, and two independent Nbs1+/−Artemis−/− and Mre11ATLD1/ATLD1:Artemis−/− abl pre–B cell lines treated with STI571 for 36 h. NF-κB EMSA is shown as a control. The data presented are representative of at least two experiments.

Figure 5. Atm inhibition in MRN-deficient cells leads to increased coding end accumulation. Southern blot analysis of EcoRV-digested genomic DNA from WT, Atm−/−, and two Mre11ATLD1/ATLD1 and Nbs1+/− abl pre–B clones containing pMX-INV. Cells were treated with STI571 for the indicated time (hours) and either DMSO (−) or the Atm inhibitor (iAtm) KU-55933 (+). Expected sizes for bands generated by nonrearranged substrate (NR), coding joints (CJ), hybrid joints (HJ), and coding ends (CE) are indicated. The data presented are representative of at least two experiments.

Figure 6. Thymocyte development in Nbs1+/− mice. (a) Number of thymocytes in Atm−/− (n = 16) and Nbs1+/− (n = 9) mice at the indicated stages of development, expressed as a percentage of the number of thymocytes found at each stage in WT Atm+/+ (n = 12) and Nbs1+/+ (n = 3) littermate controls. Calculations were done using mean numbers of thymocytes from mice of each genotype and propagated error. Error bars represent SEM. DN (double negative), CD4+CD8+; DP, CD4+CD8−; SP, single positive. (b) Representative flow cytometric analysis of TCR-β expression on CD4+CD8+ DP thymocytes from WT, Nbs1+/−, and Atm−/− mice. Numbers indicate percentage of DP cells that are TCR-β+.

The data presented are representative of at least three mice.
How does MRN function during V(D)J recombination? ATM phosphorylates a large number of proteins, including all of the components of the MRN complex, which participate in diverse and broadly functional DNA damage response pathways.
(2, 21–24). MRN functions to activate ATM, and thus the primary function of MRN in V(D)J recombination may be to activate ATM in response to Rag-mediated DSBs (10–12). However, we find that ATM is activated in response to Rag DSBs generated in Mre11A/−/− and Nbs1−/− pre-B cells, as indicated by the robust ATM-dependent phosphorylation of KAP-1 and H2AX and the nuclear translocation of NF-κB. Thus, the V(D)J recombination defects in MRN-deficient cells are not a result of a general inability of Rag DSBs to activate ATM in these cells. Importantly, although we find that ATM is activated by Rag-mediated DSBs in Mre11A/−/− and Nbs1−/− pre-B cells, these cells could have isolated defects in the phosphorylation of other specific targets required for the repair of Rag-mediated DSBs.

In addition to activating ATM, MRN could have downstream functions in the repair of Rag-mediated DSBs. In this regard, ATM promotes coding joint formation by stabilizing coding ends in postcleavage complexes, and although ATM could perform this function directly, it likely regulates the activity of downstream proteins, such as MRN, that could perform these functions (50). Rad50 has N- and C-terminal Walker A and B nucleotide binding motifs, respectively, that form a DNA binding domain upon their intramolecular association (5, 6). This association leads to the formation of a central Rad50 hook domain, which can facilitate the tethering of two distinct MRN-bound DNA molecules through an intermolecular interaction between two Rad50 hook domains (16–20). The interaction between hook domains of Rad50 proteins bound to two coding ends could stabilize these DNA ends until they are joined. Moreover, it has recently been shown that Mre11 dimers provide an interface that can bridge and juxtapose two broken DNA ends (15). Thus, it is possible that the MRN complex functions to stabilize coding ends in postcleavage complexes through the activities of both Rad50 and Mre11. This stabilizing activity could be modulated by the phosphorylation of Rad50, Mre11, and/or Nbs1 by ATM in response to Rag-mediated DSBs (21–24).

MATERIALS AND METHODS

Mice. Animals were housed in a specific pathogen-free animal facility at Washington University. Animal protocols were approved by the Washington University Institutional Animal Care and Use Committee. All mice were analyzed between 4 and 8 wk of age. The Nbs1−/−, Mre11A/−/−, Atm−/−, Artemis−/−, TCRβ−/−, and Kab70−− mice have been described previously (38, 40, 65, 69, 70). The Nbs1−/− mice were obtained from Y. Xu (University of California San Diego, La Jolla, CA). The Mre11A/−/− mice were obtained from J. Petri (Memorial Sloan Kettering Cancer Institute, New York, NY).

Flow cytometric analyses and sorting. Flow cytometric analysis of thymocyte development was performed on a FACSCalibur (BD) using FITC-conjugated anti-CD8, PE-conjugated anti-CD4, and PE-conjugated anti-TCR-β chain (all obtained from BD). Cells containing pMX-INV or pMX-DEl9 were isolated by FACS (FACSVantage; BD) based on expression of the human sequences in the different Artemis−/−, Nbs1−/−, and Artemis−/−:Mre11A/−/− abl pre-B cells analyzed. Fig. S7 shows Southern blot analyses of thymocytes from mice expressing the TCR-α/β allele were performed using the Stul restriction enzyme and the Cod probe as reported previously (65). Southern blot analyses of Igκ-lac locus Jκ coding ends was performed on Saci- and EcoRi-digested genomic DNA using the JκIII probe as previously described (48). pMX-INV coding joints were amplified using the pκ and pβ oligonucleotides and pMX-DEl9 signal joints were amplified using primers pκ and pβ oligonucleotides as previously described (50). Vβ5, Vβ14, and Vw6-23 coding and hybrid joints and the IL-2 gene were amplified by PCR as previously described (50).

Immunofluorescent detection of γ-H2AX foci. Nuclear γ-H2AX foci were detected using standard protocols with minor modifications (54, 72, 73). In brief, cells were cytospun onto poly-L-lysine-coated slides (Sigma-Aldrich), fixed in 4% paraformaldehyde for 10 min, washed in PBS, permeabilized with 0.15% Triton X-100 in PBS, and blocked in PBS with 2% bovine serum albumin. Cells were incubated in anti-γ-H2AX antibody (Millipore) at a 1–2 μg/ml concentration for 3 h at 37°C in a moist chamber, washed with PBS, and further incubated with anti-mouse FITC conjugate (Vector Laboratories) for 45 min. After washing with PBS, slides were mounted in Vectashield mounting medium with DAPI (Vector Laboratories). Foci were observed and imaged as described previously on a microscope (Axioplan 2; Carl Zeiss, Inc.) using ISIS imaging software (MetaSystems) (72).

Western blot and EMSA analyses. Western blots were done on whole cell lysates using antibodies to mouse-1 (Genex, Inc.), phosphorylated Kap-1 (Bethyl Laboratories, Inc.), and Nbs1 (Cell Signaling Technology). The secondary reagents were horseradish peroxidase-conjugated goat anti-mouse IgG (Invitrogen) or donkey anti-rabbit IgG (GE Healthcare). NF-κB EMSA were run as described previously and analyzed with an Infrared Scanner (Odyssey; LI COR Biosciences) (48).

Online supplemental material. Fig. S1 shows Southern blot analyses of pMX-INV rearrangement on additional abl pre-B cell clones. Fig. S2 shows longer exposure of the WT:INV and Atm−/−:INV Southern blot analyses. Fig. S3 shows sequences of pMX-INV coding joints from WT:INV, Artemis−/−:INV, Mre11A/−/−:INV, and Nbs1−/−:INV abl pre-B cells. Fig. S4 shows sequences of pMX-DEl9 signal joints from WT:DEL9, Artemis−/−:DEL9, Mre11A/−/−:DEL9, and Nbs1−/−:DEL9 abl pre-B cells. Fig. S5 shows analysis of Kap-1 phosphorylation carried out as described in Fig. 4 a on additional Artemis−/−:Nbs1−/− and Artemis−/−:Mre11A/−/− abl pre-B cell lines. Fig. S6 shows analysis of Artemis−/−:Nbs1−/−, and Artemis−/−:Mre11A/−/− abl pre-B cell lines. Fig. S7 shows Southern blot analyses of Igκ-lac locus Jκ coding ends after Rag induction in the different Artemis−/−, Nbs1−/−, and Artemis−/−:Mre11A/−/− abl pre-B cell lines analyzed. Fig. S8 shows EMSA analyses of NF-κB translocation to the nucleus in the different Artemis−/−, Nbs1−/−, and Artemis−/−:Mre11A/−/− abl pre-B cells not analyzed in Fig. 4 c. Online supplemental material is available at http://www.jem.org/cgi/content/full/jem.20081326/DC1.

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We thank Dr. Yang Xu for providing the Nbs1−/− mice and Dr. John Petri for providing the Mre11ATLD1 mice.

This work is supported by the National Institute of Health grants A074953 and AI47829 (B.P. Sleckman) and CA123232 (T.K. Pandita). C.-Y. Huang and A.L. Bredemeyer were supported by a postdoctoral training grant from the National Institutes of Health. J.J. Bednarski is supported by a National Institutes of Health Research Grant 5T32 HD007499 from the National Institute of Child Health and Human Development. The authors have no conflicts of financial interests.

Submitted: 19 June 2008
Accepted: 26 January 2009

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