Multiple autophosphorylation sites are dispensable for murine ATM activation in vivo

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Cellular responses to both physiological and pathological DNA double-strand breaks are initiated through activation of the evolutionarily conserved ataxia telangiectasia mutated (ATM) kinase. Upon DNA damage, an activation mechanism involving autophosphorylation has been reported to allow ATM to phosphorylate downstream targets important for cell cycle checkpoints and DNA repair. In humans, serine residues 367, 1893, and 1981 have been shown to be autophosphorylation sites that are individually required for ATM activation. To test the physiological importance of these sites, we generated a transgenic mouse model in which all three conserved ATM serine autophosphorylation sites (S367/1899/1987) have been replaced with alanine. In this study, we show that ATM-dependent responses at both cellular and organismal levels are functional in mice that express a triple serine mutant form of ATM as their sole ATM species. These results lend further support to the notion that ATM autophosphorylation correlates with the DNA damage-induced activation of the kinase but is not required for ATM function in vivo.

Introduction

Double-strand breaks (DSBs) in DNA can occur from chemical damage or γ irradiation as well as during normal physiological processes, including antigen receptor gene rearrangement and meiosis. Unrepaired DSBs are a source of genomic instability that can lead to uncontrolled cell proliferation and cancer (McKinnon and Caldecott, 2007). After sensor proteins recognize damaged chromatin, the ataxia telangiectasia mutated (ATM) kinase is recruited to the break site and initiates a cellular response through phosphorylation of consensus sites on numerous substrates that signal for repair of the DNA and/or prevent cell cycle progression (Shiloh, 2003). In addition to amplifying the DNA damage signal, ATM may also participate more directly in DNA repair by stabilizing repair complexes on damaged chromatin (Bredemeyer et al., 2006). The critical function of ATM in maintaining genomic stability is underscored by its high conservation in eukaryotes and its deficiency in the cancer-prone ataxia telangiectasia disease (Shiloh, 2003).

ATM kinase activity is directly stimulated by damaged DNA and the MRE11–RAD50–NBS1 (MRN) sensor complex, which also stabilizes ATM to the break site (Girard et al., 2002; Carson et al., 2003; Uziel et al., 2003; Lee and Paull, 2004, 2005; Difilippantonio et al., 2005; Ceresaletti et al., 2006; Dupre et al., 2006; Berkovich et al., 2007; You et al., 2007). The DNA damage–induced activation of ATM results in the dissociation of inactive ATM dimers and/or multimers into enzymatically active monomers (Bakkenist and Kastan, 2003; Lee and Paull, 2004; Dupre et al., 2006). An autophosphorylation mechanism has been proposed to initiate ATM activation upon DNA damage by promoting monomerization and subsequent localization of ATM to break sites (Bakkenist and Kastan, 2003; Berkovich et al., 2007). In human cells, serine residues 367, 1893, and 1981 (S367, S1893, and S1981) have been shown to be autophosphorylation sites that are individually required for ATM activation and function (Bakkenist and Kastan, 2003; Kozlov et al., 2006). In a reconstituted in vitro system, however, evidence suggests that ATM monomerization by MRN does not
require ATM S1981 autophosphorylation (Lee and Paull, 2005). Moreover, using *Xenopus laevis* egg extracts, high concentrations of damaged DNA can promote ATM monomerization without concomitant autophosphorylation (Dupre et al., 2006). Also, in a *Xenopus* system, there is evidence that ATM recruitment to DSBs can precede autophosphorylation (You et al., 2005), further questioning its importance in ATM activation. Because of the lack of physiologically relevant models, the significance of these autophosphorylation sites still remains poorly understood.

Using an ATM transgenic mouse model, we recently found that the most prominent S1981 autophosphorylation site in humans (S1987 in mice) is dispensable for murine ATM function in vivo (Pellegrini et al., 2006). To test whether other putative autophosphorylation sites compensate for the loss of murine S1987, we used bacterial artificial chromosome (BAC) recombineering to generate a transgenic mouse model in which all three conserved ATM serine autophosphorylation sites (S367/1899/1987) have been replaced with alanine. In this study, we show that ATM-dependent responses remain functionally intact in mice that express a double or triple serine mutant form of ATM as their sole ATM species. Collectively, our data strongly support the notion that DNA damage–induced activation of the ATM kinase does not depend on autophosphorylation.

## Results and discussion

### Triple S1987/367/1899A mutant ATM displays normal kinase activity in response to DNA DSBs

All three human ATM autophosphorylation sites are conserved in the mouse (Fig. 1A). Using BAC recombineering, we generated double S1987/367A mutant (2SA) ATM mice by creating a second S367A mutation in the S1987A murine ATM BAC described previously (Fig. 1B; Pellegrini et al., 2006). Subsequently, a third mutation was introduced into the 2SA mutant BAC to change S1899 to alanine and was used to generate triple S1987/367/1899A mutant (3SA) ATM mice (Fig. 1B). All serine mutations were confirmed in vivo by sequencing 3SA mutant founder Q7 transgenic mouse tail DNA (unpublished data). The A7 and Q7 founder lines were analyzed in detail.
To investigate the role of these autophosphorylation sites in the activation of the ATM kinase, ATM-dependent phosphorylation events were assessed in lysates from cultured splenic B cells irradiated with 10 Gy. As expected, an antibody directed against ATM S1987 autophosphorylation did not recognize ATM in  

\[ \text{Atm}^{TeSSA} \text{Atm}^{-/-} \] or  

\[ \text{Atm}^{TeSSA} \text{Atm}^{-/-} \] mice, reconfirming the original S1987A mutation in vivo (Fig. 1 C). Splenic B cells from  

\[ \text{Atm}^{TeSSA} \text{Atm}^{-/-} \] and  

\[ \text{Atm}^{TeSSA} \text{Atm}^{-/-} \] mice responded to γ irradiation similarly to  

\[ \text{Atm}^{TeSSA} \text{Atm}^{-/-} \] and  

\[ \text{Atm}^{TeSSA} \text{Atm}^{-/-} \] by displaying irradiation-induced phosphorylation of SMC1, KAP1, p53, and CHK2 (Fig. 1 C), which are events largely dependent on the ATM kinase (Banin et al., 1998; Canman et al., 1998; Matsuoka et al., 1998; Chaturvedi et al., 1999; Kim et al., 2002; Yazdi et al., 2006). Another 3SA mutant founder (line Q2), which overexpressed ATM, also displayed similar irradiation-induced substrate phosphorylation (Fig. 1 C). Similar results were found in thymocytes (unpublished data). Moreover, no differences were found in  

\[ \text{Atm}^{TeSSA} \text{Atm}^{-/-} \] transgenic mutants with one wild-type (WT) allele of  

\[ \text{Atm} \] (Fig. 1 C and not depicted), arguing against a predicted dominant interfering activity (Bakkenist and Kastan, 2003; Pellegrini et al., 2006). Together, these results demonstrate that disruption of all three autophosphorylation sites does not affect DNA damage–induced ATM kinase activity in vivo.

To complement our findings in primary mouse cells, we also tested whether all three autophosphorylation sites are required for ATM kinase activity in a completely reconstituted system. WT and 3SA mutant dimeric human ATM complexes were purified by sequential anti-Flag and anti-HA immunoprecipitation, and kinase assays were performed using p53 as a substrate (Lee and Paull, 2005). Consistent with previous findings that mutation of S1981 does not affect monomerization of ATM dimers or kinase activity (Lee and Paull, 2005), 3SA mutant ATM dimers also displayed kinase activity for p53 S15 in an MRN- and DNA-dependent manner similar to WT ATM dimers (Fig. 1 D). Collectively, our data suggest that autophosphorylation events at the conserved S1987, 367, and 1899 sites are dispensable for the kinase activity of ATM in response to DNA breaks.

**Triple S1987/367/1899A mutant Atm mice display irradiation-induced chromatin retention, cell cycle checkpoint activation, and genomic stability**

Evidence for the requirement of S1981 autophosphorylation in the accumulation of ATM at DSB sites is controversial (You et al., 2005; Pellegrini et al., 2006; Berkovich et al., 2007). In this study, using a biochemical method to detect chromatin retention of ATM protein after DNA damage (Andegeko et al., 2001), we found that mutant ATM from  

\[ \text{Atm}^{TeSSA} \text{Atm}^{-/-} \] B cells accumulates onto chromatin after 10-Gy irradiation similarly to that from  

\[ \text{Atm}^{TeSSA} \text{Atm}^{-/-} \] (Fig. 2 A). These results demonstrate that autophosphorylation at these three sites is dispensable for ATM retention on damaged chromatin.

However, autophosphorylation could still play a role in downstream functions, including checkpoint activation and maintaining genomic stability. To test whether multiple autophosphorylation sites are important for checkpoint activation, we irradiated B cells with 0, 1, 2.5, or 5 Gy and assessed the DNA damage–induced G2/M checkpoint by quantifying histone H3 S10 phosphorylation (Fig. 2 B). For all three autophosphorylation-positive cells containing diploid DNA content. Compared with  

\[ \text{Atm}^{-/-} \], which displays a defect in the G2/M checkpoint after treatment with ionizing radiation, B cells from

Figure 2. **Triple S1987/367/1899A mutant Atm mice display normal irradiation-induced chromatin retention, cell cycle checkpoint activation, and genomic stability.** (A) B cells stimulated with LPS for 48 h were harvested 30 min after no treatment (0 Gy) or after γ irradiation (10 Gy) and processed for detergent fractionation and Western blot analysis. The numbers to the right of the blot represent the measurements in kilodaltons. (B) B cells stimulated with LPS for 48 h were harvested 1 h after no treatment (unirradiated controls) or after the indicated doses of γ irradiation (grays), and mitotic cells positive for histone H3 S10 phosphorylation were quantified by flow cytometry. Data are normalized with respect to the number of mitotic cells in unirradiated samples from the same genotype. (C) LN T cells stimulated with anti-CD28 + anti-TCR-β for 48 h were treated with colcemid for 1 h and harvested for metaphase chromosome preparation. FISH was performed on slides with probes for TCR-α, chromosome 14, and telomeres and counterstained with DAPI. The percentages of metaphases (n = 80 for each genotype) with abnormalities specifically associated with the TCR locus on chromosome 14 [TCR-associated] or with all other chromosomes (general) are shown.
Triple S1987/367/1899A mutant Atm mice display normal physiological responses during recombination and exposure to ionizing radiation

In the developing thymus, ATM functions in TCR-α gene rearrangement by facilitating the resolution of recombination activation gene–induced DSBs to promote TCR-α/H9251 surface expression for subsequent CD4 and CD8 single-positive (SP) thymocyte selection (Matei et al., 2006; Huang et al., 2007; Vacchio et al., 2007). To begin examining the physiological role of multiple ATM autophosphorylation sites, we performed flow cytometry on freshly isolated thymocytes from Atm+/+, Atm+/−, Atm Tg2SA Atm−/−, Atm Tg3SA Atm−/−, and Atm−/− mice to assess thymocyte development. Percentages of CD4 and CD8 SP thymocytes in Atm Tg2SA Atm−/− and Atm Tg3SA Atm−/− mice were comparable with those found in Atm+/+ mice, whereas thymocytes from Atm−/− mice showed characteristic reductions in both SP compartments (Fig. 3 A). Moreover, thymocytes from both Atm Tg2SA Atm−/− and Atm Tg3SA Atm−/− mice displayed normal distributions of TCR-β surface expression, whereas Atm−/− thymocytes showed only low levels (Fig. 3 B), indicating that mice mutant for ATM autophosphorylation sites S1987, S367, and S1899 maintain competency for TCR-α rearrangement and thymocyte development.
A separate antigen receptor gene rearrangement process called class switch recombination (CSR) confers different effector functions for antibodies during an immune response (Stavnezer et al., 2008). In mature B lymphocytes undergoing class switching, ATM functions in recombination of switch regions that lie upstream of the immunoglobulin heavy chain constant region genes (Lumsden et al., 2004; Reina-San-Martin et al., 2004). To further understand the physiological role of multiple autophosphorylation sites for ATM function, we stimulated B cells to undergo CSR at the γ-1 constant region locus. Although Atm−/− mice displayed a reduction in IgG1-positive cells (Lumsden et al., 2004; Reina-San-Martin et al., 2004). B cells from both AtmTg2SAAtm−/− and AtmTg3SAAtm−/− mice exhibited levels of IgG1 switching similar to Atm+− (Fig. 3 C). These results suggest that ATM activation and function during CSR is intact in the absence of S1987, S367, and S1899 autophosphorylation.

The infertility of both male and female Atm−/− mice results from a lack of mature gametes caused by chromosomal fragmentation and arrest during meiosis (Barlow et al., 1996; Elson et al., 1996; Xu et al., 1996). AtmTg2SAAtm−/− mice had normal testes size and numbers of spermatids within the seminiferous tubules compared with the degeneration and complete absence of spermatids in Atm−/− testes (Fig. 3 D). Similar results were obtained from ovaries of AtmTg3SAAtm−/− mice (unpublished data). Thus, spermatogenesis and oogenesis disrupted in Atm−/− mice appear to be normal in AtmTg3SAAtm−/− mice. Collectively, our results demonstrate that physiological DNA rearrangements that require ATM function to resolve DSBs during V(D)J, CSR, and meiotic recombination are not dependent on ATM autophosphorylation at S1987, S367, and S1899.

A hallmark of ataxia telangiectasia is extreme sensitivity to the effects of ionizing radiation (Taylor and Byrd, 2005). Upon radiation exposure, the death of Atm−/− mice results from acute radiation toxicity to the gastrointestinal tract (Barlow et al., 1996). To further investigate whether multiple ATM autophosphorylation sites are important for ATM function in vivo, mice were subjected to 8 Gy of whole-body irradiation. Intestinal tissues were histologically examined 4 d after irradiation. Recovery from irradiation in the small intestines was similar in Atm+−, AtmTg3SAAtm−/−, and AtmTg3SAAtm−/− mice, whereas Atm−/− mice displayed characteristic toxicity indicated by severe epithelial crypt degeneration and loss of villi (Fig. 4). Examination of the large intestines yielded similar results (unpublished data). These data indicate that ATM autophosphorylation at S1987, S367, and S1899 are not essential for ATM function in vivo to promote tissue recovery after exposure to ionizing radiation.

The differences among results of ATM autophosphorylation mutants, mostly from human cell culture and reconstituted in vitro systems, underscore the necessity for appropriate animal models to ascertain the physiological relevance of ATM post-translational modifications. Although the possibility of different requirements in humans and mice exists, high conservation of the ATM protein sequence between the two organisms as well as the conservation of other phosphatidyl-inositol-3 kinase–related kinase family members strongly suggests mechanistic similarity in their response to DNA damage (Shiloh, 2003).
affect the irradiation-induced increase in ATM kinase activity (Pellegrini et al., 2006), providing additional evidence against an essential role for autophosphorylation in ATM activation.

Collectively with our in vivo evidence, we propose that ATM autophosphorylation occurs simultaneously or as a consequence of ATM activation and may be the combined result of ATM localization to break sites along with promiscuous phosphorylation of consensus sites exposed on the surface of ATM. By proposing that autophosphorylation is not the mechanism of ATM activation, we stimulate speculation on the relevant way in which damaged chromatin activates the ATM kinase. The MRN complex clearly plays a critical role in ATM activation by retaining ATM to DNA breaks, stimulating substrate binding, and directly stimulating kinase activity through activities on DNA by the RAD50 ATPase (Lee and Paull, 2007). Growing evidence suggests that ATM activation at DNA damage sites is mediated through direct recruitment by MRN, and to what extent other posttranslational modifications of ATM, such as acetylation at lysine 3016 (Sun et al., 2007), may function remains to be determined.

Materials and methods

Generation of mice

The S1987A mutant murine Atm BAC (Pellegrini et al., 2006) was recombined as described previously (Yang and Sharan, 2003) to contain an EcoRI site between exons 35 and 36 for a PCR-based method to distinguish between Atm<sup>S1987A</sup> and Atm<sup>Δ</sup> genotypes. The S367A and S1899A mutations were sequentially targeted into the S1987A EcoRI site-containing BAC as described previously (Yang and Sharan, 2003), and both double serine and triple serine mutant BACs were used to generate transgenic mice. The presence of the Tg was determined by PCR as previously described (Pellegrini et al., 2006). Transgenic founders were crossed to Atm<sup>−/−</sup> mice. All experiments were performed with 25A A7 and 35A QA founder lines unless otherwise noted. All experiments were performed in compliance with the National Institutes of Health Intramural Animal Care and Use program.

Lymphocyte cultures

B cells were isolated from spleens of 6–12-wk-old mice by immunomagnetic depletion with anti-CD43 beads (Miltenyi Biotech) and stimulated with either 25 μg/ml lipopolysaccharide (LPS) alone (Sigma-Aldrich) or in combination with 5 ng/ml interleukin 4 (IL4; Sigma-Aldrich) for 2–4 d as described previously (Lee and Paull, 2005). Transgenic lymphocytes were co-stained with a 10x NA 0.45 objective lens (Plan-Apochromat; Nikon) and a color charge-coupled device camera (Axioskop MRC5; Carl Zeiss, Inc.) using AxiosVision software (version 4.6.3.0; Carl Zeiss, Inc.).

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