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The initiation of meiotic sex chromosome inactivation sequesters DNA damage signaling from autosomes in mouse spermatogenesis

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Summary

Meiotic sex chromosome inactivation (MSCI) is an essential event in the mammalian male germline. MSCI is directed by a DNA damage response (DDR) pathway centered on the phosphorylation of histone variant H2AX at serine 139 (termed γ H2AX). The failure to initiate MSCI is linked to complete meiotic arrest and elimination of germ cells; however, the mechanisms underlying this arrest and elimination remain unknown. To address this question, we established a new separation-of-function mouse model for *H2ax* that shows specific and complete defects in MSCI. The genetic change is a point mutation in which another H2AX amino acid residue important in the DDR, tyrosine 142 (Y142), is converted to alanine (*H2ax-Y142A*). In *H2ax-Y142A* meiosis, the establishment of DDR signals on the chromosome-wide domain of the sex chromosomes is impaired. The initiation of MSCI is required for stage progression, which enables crossover formation, suggesting that the establishment of MSCI permits the timely progression of male meiosis. Our results further suggest that normal meiotic progression requires the removal of

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Author Contributions

H.A. and S.H.N. designed the research. Y.C.H. and R.S.H. designed the *H2ax-Y142A* mice, and Y.C.H. supervised the generation of the *H2ax-Y142A* mice. H.A. performed the experiments. H.A., K.G.A., Y.C.H., Q.P., P.R.A., R.S.H., and S.H.N. analyzed the data. H.A., K.G.A., R.P.A., and S.H.N. wrote the manuscript. S.H.N. supervised the investigation. All authors reviewed the manuscript.

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ATR-mediated DDR signaling from autosomes. We propose a novel biological function for MSCI: The initiation of MSCI sequesters DDR factors from autosomes to the sex chromosomes at the onset of the pachytene stage, and the subsequent formation of an isolated XY nuclear compartment—the XY body—sequesters DDR factors to permit meiotic progression from the mid pachytene stage onward.

Graphical Abstract



eTOC Blurb

Abe *et al.* demonstrate that tyrosine 142 of histone variant H2AX is required for the initiation of meiotic sex chromosome inactivation (MSCI). Based on new genetic evidence, the study proposes a novel biological function for MSCI: MSCI sequesters DNA damage signaling from autosomes to permit timely progression of male meiosis.

Keywords

Meiosis; DNA damage signaling; checkpoint; germline; spermatogenesis

Introduction

Meiosis is a hallmark event in germline development, when paternal and maternal chromosomes undergo synapsis and a reshuffling of the genome prior to producing haploid gametes. During meiosis, the fidelity of meiotic recombination and chromosome synapsis is strictly monitored by checkpoint mechanisms. In coordinating these and other critical events in meiosis, checkpoint mechanisms facilitate timely progression of germ cells through meiosis. Importantly, evolutionarily conserved proteins in DNA damage response (DDR) pathways are implicated in meiotic checkpoint mechanisms of a variety of organisms, from yeast to worms to mammals [1]. Yet despite our understanding of meiotic checkpoints in yeast and worms, their molecular operation remains largely unknown in mammals [2].

In mammalian male meiosis, the X and Y chromosomes are subjected to meiotic sex chromosome inactivation (MSCI) [3, 4]. MSCI is an essential process in the male germline, as failure to initiate MSCI is linked to the complete arrest and timely elimination of male germ cells in the mid pachytene stage of meiotic prophase I [5, 6]. MSCI is a sex chromosome-specific manifestation of a general mechanism for transcriptional silencing in meiosis termed meiotic silencing of unsynapsed chromatin, which operates as a surveillance mechanism for chromosome asynapsis [7–10]. Mechanistically, the initiation of meiotic silencing is directed by a DDR pathway centered on the kinase Ataxia telangiectasia and Rad3 related (ATR) and its phosphorylation of histone H2AX at serine 139 (termed γ H2AX) [3, 5, 6, 11–15]. In response to meiotic chromosome asynapsis, a large γ H2AX "domain" forms through signal amplification of ATR-mediated γ H2AX—first from the unsynapsed axes, then to their protruding loops of chromatin; the signal amplification is directed by Mediator of DNA damage checkpoint protein 1 (MDCI), a γ H2AX-binding protein [3, 6]. A major question remains as to how the failure to initiate MSCI is related to complete meiotic arrest and the timely elimination of spermatocytes.

To understand the molecular events that occur in response to MSCI defects, we generated and studied a new separation-of-function mouse model for *H2ax*. This model contains a point mutation in which another H2AX amino acid residue important in the DDR, tyrosine 142 (Y142) [16, 17], is converted to alanine (*H2ax-Y142A*). In somatic cells, Y142 is constitutively phosphorylated under physiological conditions; Y142 becomes dephosphorylated upon DNA damage, enabling interactions between γ H2AX and other DDR factors, including MDC1, which promote DNA repair [17]. In meiosis, the dephosphorylation of Y142 takes place on the sex chromosomes at the onset of MSCI [18]. Here, we find that *H2ax-Y142A* mice show specific and complete defects in MSCI, supporting the notion that a common DDR pathway operates in both the somatic DDR and MSCI [3].

In this study, we sought to define the common molecular events that occur in response to defective MSCI. We analyzed the *H2ax-Y142A* mouse model and an *Mdc1* knockout (*Mdc1*KO) mouse model, both of which exhibit complete impairment of MSCI [6]. Our results suggest that the initiation of MSCI sequesters DDR signaling from autosomes to the sex chromosomes and that the establishment of MSCI permits the timely progression of male germ cells through meiotic prophase I. We propose a novel biological function for MSCI: The initiation of MSCI sequesters DDR factors from autosomes at the onset of the pachytene stage, and the subsequent formation of a distinctly compartmentalized XY body sequesters DDR factors to enable meiotic progression from the mid pachytene stage onward. This model postulates a new mechanism for the meiotic checkpoint: An ATR-mediated DDR pathway that operates in the checkpoint mechanism of the somatic cell cycle has direct regulatory roles in the MSCI checkpoint, which operates to induce cell death if DDR signals remain on autosomes.

Results

An H2AX Y142A point mutation disrupts spermatogenesis

We used CRISPR/Cas9 genome-editing technology to generate a knock-in mouse line harboring a point mutation that changes tyrosine 142 of H2AX into an alanine residue (*H2ax-Y142A*: Figure 1A). Homozygous males for *H2ax-Y142A* (hereafter referred to as "*H2ax-Y142A*") were infertile, had reduced body weight (Figure 1B), and had significantly smaller testes in comparison to wild-type littermate controls (Figure 1C). We confirmed the abrogation of phosphorylated Y142 (pY142) of H2AX in *H2ax-Y142A* testes through two methods: western blotting using wild-type and *H2ax-Y142A* testis lysates (Figure 1D), and immunofluorescence of wild-type pachytene spermatocytes, pY142 signals were detected throughout nuclei except at XY bodies [18]; pY142 signals were not detected in *H2ax-Y142A* mouse line.

Next, we examined the spermatogenic stage progression of H2ax-Y142A mice using two specific markers: H1T and γ H2AX. H1T is a testis-specific histone variant that accumulates in nuclei from the mid-pachytene stage onward [19]. γ H2AX is a prominent marker of the DDR at DNA double-strand breaks (DSBs), stalled replication forks, and MSCI [20, 21]. We observed complete spermatogenetic arrest at the mid pachytene stage (H1T-positive) in H2ax-Y142A testes (Figure 1F). Accordingly, the epididymides of H2ax-Y142A mice displayed a complete absence of sperm (Figure S1A). Apoptotic cell death is linked to germ cell loss in H2ax-Y142A testes (Figure S1B), while the point mutation did not noticeably change H2AX protein abundance (Figure S1C). Thus, the Y142A mutation reflects the biological importance of an H2AX residue rather than an effect mediated by destabilization of the H2AX protein. Together, these results suggest that Y142 of H2AX is an amino acid residue essential for the progression of spermatogenesis. In contrast to the severe H2ax-Y142A phenotype observed in males, H2ax-Y142A female mice were fertile (6 independent females were tested; mean litter size: 6.33 ± 0.48 , mean \pm s.e.m.) similar to H2ax null females [5, 22].

H2AX-Y142 is critical for the establishment of MSCI

In the leptotene stage of meiotic prophase I, the topoisomerase II-like enzyme SPO11 induces programmed DNA double-strand breaks (DSBs) and, in the subsequent zygotene stage, the synapsis of homologs takes places [23–25]. In response to SPO11-dependent DSBs, an initial wave of γ H2AX signaling takes place throughout nuclei in the leptotene stage. In chromosome spreads of *H2ax-Y142A* leptotene spermatocytes, as in wild-type controls, γ H2AX displayed apparently normal nuclear accumulation patterns (Figure S2A). Upon homolog synapsis in normal meiosis, γ H2AX signals diminish from autosomes, and a chromosome-wide γ H2AX domain is established on the sex chromosomes—a hallmark of MSCI (Figure 2A, top panel). In *H2ax-Y142A* spermatocytes, we detected no sex chromosome-wide accumulation of γ H2AX; instead, we observed an intense, linear accumulation of γ H2AX along the XY axes (Fig. 2A, bottom panel). This conclusion was confirmed through immunostaining and imaging of testis sections (Fig. 1F, arrows in the

upper right panel). Importantly, the linear accumulation of γ H2AX along the XY axes comprised a completely penetrant phenotype: We observed no H2ax-Y142A spermatocytes with a chromosome-wide γ H2AX domain (>500 nuclei observed from >10 independent H2ax-Y142A mice). Additionally, in staging H2ax-Y142A spermatocytes, we did not observe spermatocytes beyond the mid pachytene stage, neither in our analyses of testis sections nor in our analyses of chromosome spreads of spermatocyte nuclei. In normal male meiosis, a feedforward mechanism mediates the formation of a γ H2AX domain on the sex chromosomes; this mechanism sees the accumulation of γ H2AX along unsynapsed axes spread to chromatin loops to form a chromosome-wide domain [3, 6] (Figure 2B). The mechanism is centered on MDC1, a yH2AX-binding protein that interacts with TOPBP1 [26]; the binding of MDC1 to γ H2AX enables the progressive activation of ATR kinase through its activator protein TOPBP1 [12, 27, 28], which then mediates adjacent yH2AX formation on chromatin loops. MDC1 binds γ H2AX again and the mechanism continues onward. In our previous study of the Mdc1KO mouse model, we observed a strikingly similar accumulation of γ H2AX signals along sex chromosome axes but not through chromatin loops [6].

Despite a completely penetrant sex chromosome phenotype, autosome synapsis was unimpaired in *H2ax-Y142A* spermatocytes as mutant chromosome spreads immunostained against SYCP1 (a marker of synapsed chromosome axes) revealed no apparent abnormalities (Figure 2C). We corroborated this observation by immunostaining against another marker of chromosome synapsis, SIX6OS1 [29] (Figure S2B). Together, these data confirm the unimpaired formation of the synaptonemal complex in *H2ax-Y142A* spermatocytes. Furthermore, the partial synapsis of the X and Y chromosomes at an area of homology known as the pseudoautosomal region was present in 87.86% of observed *H2ax-Y142A* spermatocytes, a proportion that is slightly lower in comparison to control spermatocytes (Figure 2C). A recent study revealed that autosome synapsis is partially defective in *H2ax* null mutants, suggesting that H2AX has additional roles in the regulation of autosome synapsis [30]. However, the role of H2AX-Y142 is restricted to the regulation of the sex chromosomes. Thus, we conclude that *H2ax-Y142A* is a separation-of-function mutation for the *H2ax* gene.

To determine the function of H2AX-Y142 in the regulation of DDR signals, we examined the localization of ATR and TOPBP1. In *H2ax-Y142A* spermatocytes, ATR and TOPBP1 are restricted to the axes of sex chromosomes with no apparent spreading to chromatin loops, unlike wild-type spermatocytes (Figures 2D and 2E). These data confirm a function for H2AX-Y142 in the establishment of a chromosome-wide domain. In contrast to these proteins, in *H2ax-Y142A* spermatocytes, the localization of BRCA1 and HORMAD2 (factors that localize to unsynapsed axes to regulate ATR signaling [13, 15, 31]) was unchanged (Figures 2F and 2G). Therefore, the function of H2AX-Y142 is specific to the XY chromatin domain. Given these findings, we examined the localization of MDC1 on the XY chromatin of *H2ax-Y142A* spermatocytes. Interestingly, we observed punctate MDC1 foci on *H2ax-Y142A* sex chromosome axes, unlike the sex chromosome-wide distribution of MDC1 in wild-type spermatocytes (Figure 2H). These data suggest that, while MDC1 was recruited to the axes of *H2ax-Y142A* sex chromosomes, MDC1 signals were not amplified throughout the XY chromatin. Taken together, we conclude that Y142 of H2AX is required

for the MDC1-dependent amplification of DDR signals throughout XY chromatin (Figure 2B).

To further determine the necessity for H2AX-Y142 in MSCI, we examined the status of active transcription on the sex chromosomes. Using slides that preserve the 3D chromatin organization of spermatocytes (3D slides) [32, 33], we judged the localization of RNA polymerase II (RNAPII) signals with respect to the bulbous chromo-nuclear compartment that encompasses XY chromatin in mid pachytene (H1T-positive) spermatocytes. In wildtype mid pachytene spermatocytes, RNAPII signals were largely excluded from the XY body (Figure 3A), consistent with previous reports [6, 32]. However, in stage-matched H2ax-Y142A spermatocytes, RNAPII signals were overtly present throughout XY chromatin, while DAPI-discernible XY bodies were absent (Figure 3A). Likewise, the compaction of the XY axes, another telltale signature of MSCI initiation and XY body formation, was not observed in H2ax-Y142A mid pachytene spermatocytes, neither in our analyses of testis sections (Figure 3B) nor in our analyses of 3D slides (Figure 3C). Interestingly, these observations are essentially the same as those made in studies of other mouse models with complete defects in the initiation of MSCI, including loss-of-function models for H2AX, MDC1, ATR, and TOPBP1 [5, 6, 14, 28]. Therefore, we conclude that H2AX-Y142 is required both for the establishment of MSCI and for formation of the XY body.

The initial steps of autosome DSB repair are normal in MSCI-defective mutants

The results presented here demonstrate the *H2ax-Y142A* model has a completely penetrant defect in MSCI initiation, a finding that corroborates the essentiality of MSCI in spermatogenesis [5, 6, 13–15, 28, 34]. In turn, this raises the possibility that failure of MSCI is a direct cause of meiotic arrest and germ cell demise. However, if DSB repair in meiotic recombination were defective, that would suggest a recombination-dependent checkpoint operates to trigger germ cell demise in meiotic prophase I [1, 35, 36]. To test the hypothesis that MSCI failure is a direct cause of meiotic arrest in *H2ax-Y142A* spermatocytes, we evaluated the status of meiotic recombination on *H2ax-Y142A* autosomes.

To assess the status of DSB repair in H2ax-Y142A spermatocytes, we evaluated RAD51 foci in spermatocyte nuclei. An established surrogate for DSBs, RAD51 foci appear in response to SPO11-induced DSBs in leptotene spermatocytes [37]. As leptotene spermatocytes advance into the zygotene and pachytene stages, there is a progressive decrease in the numbers of RAD51 foci concomitant with the repair of DSBs, which is completed in the pachytene stage [38]. In our analyses of H2ax-Y142A spermatocytes, we observed a normal accumulation of RAD51 foci in the leptotene and zygotene stages (Figure S3A); as well, we observed comparable numbers of RAD51 foci between wild-type and H2ax-Y142A mid pachytene spermatocytes (per nucleus, we observed <10 autosomal foci: Figure 4A). These data indicate that the initial steps of DSB repair are apparently normal on the autosomes of H2ax-Y142A spermatocytes. Interestingly, we observed a slight increase in the numbers of RAD51 foci on the sex chromosome axes of H2ax-Y142A mid pachytene spermatocytes (wild-type mid pachytene spermatocytes: 17.69 \pm 0.67, mean \pm s.e.m.; H2ax-Y142A mid pachytene spermatocytes: 22.01 \pm 0.76, mean \pm s.e.m.). This finding further confirms a sex

chromosome-specific function for H2AX-Y142—a function apparently extraneous to the formation and resolution of autosomal DSBs.

While we judged DSB resolution to be grossly normal, a small-but-vitally important proportion of DSBs resolve via a specialized crossover recombination repair pathway that facilitates the reshuffling of genetic information between maternal and paternal chromatids [23–25]. Thus, we sought to evaluate crossover recombination repair in *H2ax-Y142A* and control spermatocytes. In *H2ax-Y142A* spermatocytes, we noted a distinct absence of MLH1 foci (Figure S3B), which mark sites of crossover recombination [39]. We infer this is due to the inability of *H2ax-Y142A* spermatocytes to advance past the mid pachytene stage into the late pachytene stage, when the vast majority of MLH1 focal accumulation takes place (Figure S3B). In a previous report of *Mdc1*KO spermatocytes: The absence of MLH1 foci was concomitant with completely defective MSCI [6]. This commonality suggests that the initiation of MSCI is necessary for meiotic progression to the stage of MLH1 focal accumulation.

In a previous report analyzing chromosome spreads of spermatocyte nuclei [40], it was demonstrated that future sites of crossover exhibit focal accumulation of another crossover marker, MLH3, prior to the accumulation of MLH1 foci. To evaluate the initial step of crossover recombination, prior to the arrest of mid pachytene spermatocytes, we scored the numbers of MLH3 foci in control and H2ax-Y142A spermatocytes. In early pachytene (H1T-negative) spermatocytes from both control and H2ax-Y142A mice, MLH3 foci were detected at similarly low levels (Figure 4B). However, in mid pachytene (H1T-positive) spermatocytes, the numbers of MLH3 foci differed between wild-type and H2ax-Y142A: In wild-type samples, MLH3 focal accumulation showed a marked increase (8.72 \pm 0.61, mean \pm s.e.m.; Figure 4B); in *H2ax-Y142A* samples, we noted reduced numbers of MLH3 foci $(2.01 \pm 0.23, \text{mean} \pm \text{s.e.m.};$ Figure 4B). In wild-type late pachytene spermatocytes, the numbers of MLH3 foci reached their zenith (>20, with at least one per chromosome; Figure S3C), similar to the focal accumulation of MLH1; this suggests MLH3 focal accumulation grows in and through the mid pachytene stage. Meiotic arrest likely takes place early in the mid pachytene stage, when MLH3 focal accumulation is in its initial growth phase (Figure 4C). Importantly, this timing coincides with the formation of the XY body, a distinct, DAPIdiscernible chromo-nuclear compartment at the periphery of the mid pachytene nucleus (Figure 4D) [32]. In the early pachytene stage, the diffuse, indistinct chromosome-wide accumulation of γ H2AX on XY chromatin represents the initiation of MSCI, and it is not until the mid pachytene stage that a bulbous, bounded, and distinctly compartmentalized XY body appears at the nuclear periphery [15, 41, 42]. We infer that the complete death of H2ax-Y142A spermatocytes is linked to the failure to form an XY body.

To further define the molecular events that occur in response to defective MSCI, we analyzed meiotic DSB repair in *Mdc1*KO mice, a separate, independent genetic model exhibiting complete failure of MSCI [6]. We did not examine other mutants for direct regulators of MSCI, such as H2AX, ATR, and TOPBP1, because they are directly involved in meiotic recombination [5, 14, 28, 30]. Similar to our analyses of *H2ax-Y142A* and control mice, the numbers of RAD51 foci were comparable on the mid pachytene autosomes

of Mdc1KO and control mice (Figure 4E). Interestingly, RAD51 foci on Mdc1KO sex chromosomes were comparable with wild-type sex chromosomes (Figure 4E); this is in contrast to the slight increase in RAD51 focal accumulation observed on H2ax-Y142A sex chromosomes (Figure 4A), suggesting that RAD51 foci on the sex chromosomes may be regulated by H2AX-Y142 but not MDC1. Taken together our findings indicate that the initial steps of meiotic DSB repair are grossly normal in *Mdc1*KO spermatocytes, thereby confirming that MSCI deficiency does not disturb the initial steps of meiotic recombination. Furthermore, reduced numbers of MLH3 foci were observed in mid pachytene spermatocytes of Mdc1KO mice (Figure 4F). Thus, based on findings from two independent mouse models, we conclude that meiotic arrest takes place in the midst of the mid pachytene stage prior to the full accumulation of MLH3 foci (Figure 4C). Importantly, we have excluded the possibility that the meiotic arrest observed in H2ax-Y142A mice is caused by the reduction of MLH3 foci because MLH3-deficient spermatocytes are capable of reaching the metaphase stage of meiosis I [43]. Also, our results further suggest the successful initiation of MSCI is required for stage progression, which enables crossover formation in male meiosis.

Finally, we evaluated meiosis in female *H2ax-Y142A* mice and confirmed normal meiotic recombination (Figure S4). Consistent with the normal fertility of *H2ax-Y142A* females, MLH1 foci were observed in *H2ax-Y142A* pachytene oocytes (Figure S4A). Therefore, Y142 of H2AX is dispensable for the formation of MLH1 foci in female meiosis. This result underscores specific functions for H2AX-Y142 in male-specific MSCI. In support of this notion, regulators of MSCI—such as MDC1, H2AX, BRCA1, and HORMAD2—are not required for female meiosis, and female models for the loss-of-function of these factors are fertile [6, 22, 31, 44, 45].

ATR-associated DDR signals are sequestered from autosomes to the sex chromosomes at the onset of MSCI

Our results suggest the establishment of MSCI permits the timely progression of male germ cells through meiotic prophase. By definition, checkpoints monitor the completion of essential cellular events, enabling cells to progress to their next stages. In meiosis, it is generally thought meiotic checkpoint mechanisms coordinate genetically distinct and/or independent meiotic processes [1]. Since genetic evidence, including the current study, shows that meiotic arrest and subsequent cellular demise ensue when MSCI is abrogated [5, 6, 13–15, 31, 45, 46], we reasoned that the initiation of MSCI functions as a checkpoint to coordinate and pace precise meiotic stage progression during normal meiosis.

A previous study proposed that, in response to defective MSCI, germ cell death results from the ectopic expression of sex chromosome-linked genes [34]. However, DDR signaling and particularly ATR signaling—is directly involved in the meiotic checkpoints of various organisms, including yeast, worms, and flies [1]. These observations, in combination with our new results, compelled us to hypothesize that DDR signaling itself functions as a checkpoint in MSCI. This is a novel model for the MSCI checkpoint (see Discussion). To test this hypothesis, we sought to identify DDR signaling phenotypes common to both the *H2ax-Y142A* and *Mdc1*KO models, which exhibit specific MSCI defects.

In our previous evaluation of mid pachytene *Mdc1*KO spermatocytes, we observed persistent yH2AX and TOPBP1 foci on autosomes axes when MSCI is abrogated [6]. In H2ax-Y142A spermatocytes in the mid pachytene stage, we observed that γ H2AX and TOPBP1 foci were present largely at autosome axes (Figures 2A and 5A). On the other hand, in the transition from the early-to-mid pachytene stages in wild-type spermatocytes, these DDR signals progressively faded from autosomes. This retention of TOPBP1 foci on autosome axes was comparable between H2ax-Y142A and Mdc1KO spermatocytes (Figures 5A–5C: $65.95 \pm$ 4.26, mean \pm s.e.m. for *H2ax-Y142A*; 76.09 \pm 2.76, mean \pm s.e.m. for *Mdc1*KO). Importantly, the retention of YH2AX and TOPBP1 foci in H2ax-Y142A and MdcIKO mice was largely independent of persistent DSBs: The numbers of autosomal yH2AX and TOPBP1 foci were much higher than those of RAD51 foci, and we did not observe differences in the numbers of RAD51 foci between controls and mutants (Figures 4A and 4E). Since TOPBP1 is an ATR activator [47], we also examined the retention of ATR foci. In normal meiosis, γ H2AX and ATR foci are confined to the XY body in the mid pachytene stage. In contrast, we confirmed the retention of γ H2AX and ATR foci on autosome axes in H2ax-Y142A and Mdc1KO pachytene spermatocytes (Figures 5D and 5E). Furthermore, we found that ATR interacting protein (ATRIP), a binding partner of ATR in checkpoint signaling [48] and a protein that binds unsynapsed meiotic chromosome axes [49], was retained on autosome axes in H2ax-Y142A pachytene spermatocytes (Figure S5). These results raise a compelling possibility: In the early pachytene stage, MDC1-dependent amplification of DDR signals on the XY chromatin sequesters the DDR signaling to the sex chromosomes and away from autosomes (Figure 6). Although another interpretation is possible: MDC1 and H2AX-Y142 are required for the release of DDR signals from autosomes axes. We believe this possibility is unlikely because the release of DDR signals from chromosome axes takes place normally in H2ax-Y142A pachytene oocytes (Figure S4B and S4C).

Discussion

In this study, we show that *H2ax-Y142A* is an *H2ax* separation-of-function mutation that exhibits specific defects in MSCI. We capitalized on this model to dissect the molecular events that occur in response to defective MSCI. In interpreting our results in the context of the literature, we propose a new model for the MSCI checkpoint wherein MSCI sequesters ATR-mediated DDR signaling from autosomes to permit meiotic progression (Figure 6).

The Y142 residue of H2AX is essential for MSCI

We show that H2AX-Y142 is important for the function of MDC1 in MSCI. In *H2ax-Y142A* mutants, the localization of MDC1 to the XY axes was significantly reduced (Fig. 2H); this raises the possibility that the binding of MDC1 to H2AX with an A142 point mutation is less than the binding of MDC1 to unphosphorylated H2AX-Y142. This interpretation is in line with results reported for somatic cells [50, 51]: In response to irradiation-induced DNA damage, an H2AX-Y142A point mutation abolishes the binding of MDC1 to γ H2AX (i.e., H2AX phosphorylated at S139). However, the PI3-kinases that act in response to irradiation-induced DNA damage in somatic cells differ from those that act in MSCI. At sites of irradiation-induced DSBs, the kinase ATM is thought to amplify γ H2AX

signals in conjunction with MDC1 [52]; in contrast, ATR-mediated γ H2AX is amplified in conjunction with MDC1 in MSCI. Since ATR-mediated γ H2AX is linked to stalled replication forks in somatic cells [20], the current study supports a model in which the ATR-mediated DDR pathway that induces MSCI is an adapted version of the ATR-mediated DDR pathway that recognizes replication stress in S phase of somatic cells [3, 6].

Importantly, the initial steps of DSB repair and ATM-mediated γ H2AX signaling take place normally in both *H2ax-Y142A* and *Mdc1*KO meiosis [6]. Therefore, we conclude H2AX-Y142 and MDC1 are not required for ATM-dependent processes in meiosis, such as the control of DSB numbers [53]. This is in contrast to a recent study postulating H2AX and MDC1 are involved in ATM-dependent processes in meiosis [30]. However, the normal DSB repair found in the current study suggests the ATM-related recombination-dependent checkpoint—triggered by persistent DSBs in early pachytene spermatocytes [35, 36, 54, 55] —may not be activated in *H2ax-Y142A* and *Mdc1*KO spermatocytes. Interestingly, *H2ax null* mice exhibited additional defects beyond those of *H2ax-Y142A* mice, including chromosome synapsis abnormalities [30]. Such defects may represent H2AX functions genetically distinct from H2AX-Y142.

The numbers of RAD51 foci on the male sex chromosomes in *H2ax-Y142A* spermatocytes were slightly increased in comparison to wild-type sex chromosomes. In somatic cells, H2AX controls recombination both of homologous chromosomes and sister chromatids [56]. Since sister chromatids comprise the only templates for DSB repair of the male sex chromosomes, we postulate that H2AX-Y142 has a critical role in the control of H2AX-mediated sister chromatid recombination. Alternatively, H2AX may be a key factor in a proposed role for MSCI in the suppression of illegitimate recombination between unsynapsed regions of the X and Y chromosomes [57]; as well, H2AX may be critical for proposed roles of meiotic silencing in suppressing nonhomologous recombination on unsynapsed axes [4]. However, in the above scenarios, the function of H2AX is likely independent of MDC1, since we did not observe increased numbers of RAD51 foci on sex chromosomes in *Mdc1*KO mice.

A novel model for the MSCI checkpoint

The biological function of MSCI remains a major unsolved question. We propose a novel function for MSCI: Initiation of MSCI sequesters DDR factors from autosomes to the sex chromosomes. According to this model, when MSCI is defective, the retention of ATR-mediated DDR signaling on autosomes directly induces cell arrest and demise (Figure 6). We postulate that this ATR-mediated surveillance mechanism directly functions in the MSCI checkpoint, which, in establishing MSCI, results in an XY body that sequesters ATR-mediated DDR signaling to permit the timely progression of male meiosis (Figure 6). In our reasoning, MSCI can be interpreted as a checkpoint since classical checkpoints function to coordinate essential cell cycle-related events [58, 59]. In support of this, ATR-mediated DDR signaling—including its regulatory mechanisms (e.g., those enabled by BRCA1 and HORMAD1/2)—is tightly linked to the control of meiotic progression, as has been demonstrated in many previous studies [11, 13, 15, 31, 45, 46, 60]. In concert with a possible role for ATR in the MSCI checkpoint, ATR functions to regulate checkpoints in

other contexts too, including the intra-S replication checkpoint in somatic cells as well as the G2 DNA damage checkpoint in somatic cells [61, 62]. Importantly, CHEK2, a downstream effector of ATR, was shown to be critical in an oogenic checkpoint [63], further indicating the direct function of DDR signaling in certain checkpoints.

Interestingly, there may exist a specific threshold for the amount of ATR-mediated DDR signaling that triggers mid pachytene cell death. The retention of small amounts of ATRmediated DDR signaling on autosomes is compatible with meiotic progression through the mid pachytene stage as in, for example, small genomic regions of asynapsis; yet apparently, larger amounts lead to arrest and apoptosis [64, 65]. Furthermore, MSCI is attenuated but not defective in response to a hypomorphic mutation of the Atr gene, and persistent DDR signaling on autosomes is associated with cell death beyond the mid pachytene stage [66]. The current work reveals an intriguing finding: The retention of ATR-mediated DDR signaling on autosomes is largely independent of DSB repair. Therefore, the MSCI checkpoint is functionally separable from the recombination-dependent checkpoint resulting from persistent DSBs [35, 36, 54, 55]. In relation to the MSCI checkpoint model presented here, the depletion of CHEK1, another downstream effector of ATR, results in the accumulation of DSB-independent DDR signaling on autosomes as well as abnormal meiotic progression [41]. A previous study demonstrated that the amounts of DDR factors present in pachytene spermatocytes are fixed [46]; thus, extensive autosome asynapsis may attenuate MSCI by overdrawing from a limited bank of DDR proteins. Thus, our new model provides a logical explanation for why an ATR-mediated DDR pathway recognizes the asynapsis of autosomes and then eliminates germ cells with chromosome abnormalities.

Although the unique characteristics of the MSCI checkpoint do not invalidate other kinds of checkpoints, we need to carefully distinguish between our new model of the MSCI checkpoint and previously described meiotic checkpoint mechanisms. This is because no protein has been identified that, when absent, bypasses the arrest linked to MSCI defects. ATR itself has functions in meiotic recombination [66, 67], and ATR loss-of-function is associated with cell death in the mid pachytene stage [14]. According to our MSCI checkpoint model, sequestration of DDR signaling from autosomes to the XY body, a chromo-nuclear compartment isolated from the rest of the nucleus, permits meiotic progression at the mid pachytene stage. It can be reasonably assumed that, in spermatocytes, a variety of proteins must be sequestered to the XY body at certain thresholds to ensure continued stage progression and gamete development. Notably, in such a scenario, then, the depletion or mutation of a given DDR protein cannot bypass the meiotic checkpoint to permit meiotic progression.

MSCI defects are compatible with cell survival in the early pachytene stage but result in meiotic arrest and death with impressive exactitude in the mid pachytene stage. Indeed, XY bodies are invariably absent in mutants with defective MSCI and/or chromosome synapsis abnormalities [5, 6, 15, 46], indicating a tight, underlying link between the formation of an isolated XY body and meiotic progression. We infer the biological function of the XY body is to sequester specific nuclear components, including DDR signals/checkpoint machinery, from the remainder of the nucleus. The phenotypes of *Spo11*–/– spermatocytes, which display extensive chromosome asynapsis abnormalities and undergo cell death, could arise

from an inability to isolate such nuclear components. While SPO11-deficient spermatocytes manifest one or more diffuse sites of ectopic meiotic silencing—termed "pseudo-XY bodies"—they are not compatible with meiotic progression since pseudo-XY bodies do not constitute distinct, bounded, physically isolated chromo-nuclear compartments like true XY bodies [11, 15]. On the other hand, mutants deficient for XY-body markers that function downstream of the ATR-mediated DDR pathway that initiates MSCI—including, but not limited to, AGO4, RNF8, SCML2, FANCB—are not associated with mid pachytene arrest [68–71]. These XY-body markers may function subsequent to progression through the MSCI checkpoint.

Interestingly, our recent study suggests the 3D chromatin organization of the sex chromosomes during MSCI arises from phase separation [72], a physical process in which membraneless organelles form and behave as liquid droplets [73, 74]. Following on this, it is intriguing to speculate that MDC1-dependent amplification of γ H2AX gives rise to nuclear environments in which it is energetically favorable for DDR signals to coalesce, a liquid-liquid condensation that sequesters DDR signals/checkpoint machinery from their initial sites of accumulation on autosome axes and chromatin. In the absence of such a mechanism, DDR signaling is retained on autosomes.

In contrast to our new model, a previous study suggested another potential cause of meiotic cell death in mutants with defective MSCI: Derepression of toxic genes from sex chromosomes such as the Y-linked genes Zfy1 and Zfy2 [34]. Given our growing knowledge of spermatogenesis, and given that both ZFY1 and ZFY2 function in normal spermatogenesis [75–77], derepression of sex-linked genes is unlikely to be the sole mechanism inducing cell death. As well, the extended timespans, noise, and cell-to-cell variability of transcription- and translation-dependent biological processes suggest toxic gene derepression is likely not compatible with the complete, clockwork nature of mid pachytene cell death. In our previous study of Scml2KO mice, the abnormal expression of sex-linked genes in the pachytene stage was found to be compatible with meiotic progression [70], and Zfy1 and Zfy2 were derepressed in Scml2KO pachytene spermatocytes (Figure S6). Furthermore, SCML2 and FANCB are examples of X-linked gene products that function on the XY body [70, 71], indicating that sex-linked gene products can exert physiological functions even after transcriptional silencing via MSCI. Since gene dosage of the sex chromosomes is a critical determinant of fertility [78], it is possible this gene dosage is monitored by quality control mechanisms apart from the MSCI checkpoint. The monitoring of gene dosages and checkpoints are unlikely to be mutually exclusive. Thus, continued investigation will be important to more precisely identify the molecular mechanisms underlying the MSCI checkpoint.

STAR METHODS

CONTACT FOR REAGENT AND RESOURCE SHARING

Further information and requests for reagents should be directed to the Lead Contact, Satoshi H. Namekawa (satoshi.namekawa@cchmc.org). *H2ax-Y142A* mice are available for sharing.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Mouse lines—H2AX tyrosine 142-to-alanine 142 point mutation mice (H2ax-Y142A mice or H2ax-Y142A) used in this study were generated via CRISPR-Cas9 gene editing technology in the Transgenic Animal and Genome Editing Core Facility of Cincinnati Children's Hospital Medical Center. The methods for the design of sgRNAs, the design of donor oligos, and the production of animals have been described [79]. The sgRNA target sequences (5' GCCTCTCAGGAGTACTGAGG 3') were cloned through established methods [80] into a modified pX458 vector (Addgene, 48138); the pX458 vector contains an optimized sgRNA scaffold and a high-fidelity Cas9 [81, 82]. Editing activity was validated via T7 Endonuclease 1 mismatch assays in mouse mK4 cells [83] in side-by-side comparisons with Tet2 sgRNA, which is known to work efficiently in mouse embryos [84]. Validated sgRNA was transcribed in vitro using the MEGAshortscript T7 Transcription Kit (ThermoFisher Scientific, AM1354), purified by the MEGAclear Transcription Clean-UP Kit (ThermoFisher Scientific, AM1908), and then stored at -80°C. To prepare the injection mix, we incubated sgRNA and Cas9 protein (ThermoFisher Scientific, A36498) at 37°C for 10 minutes (min) to form ribonucleoproteins; then, we added the single-strand DNA donor oligo (IDT) to the mixture. The final concentrations were as follows: 50 ng/µl sgRNA, 100 ng/µl Cas9 protein, and 100 ng/µl DNA donor oligo. Using a piezoelectric microinjection technique [85], mutant mice were generated via microinjections of the mix into the cytoplasm of fertilized eggs on a C57BL/6 background. On the same day as the microinjections, the eggs were transferred into the oviduct ampulla of pseudopregnant CD-1 female mice.

Pups were born and genotyped by PCR with AfIII (NEB, R0520S) enzyme digestion. Animals were housed in a controlled environment with a 12-hour (h) light/12-h dark cycle and with free access to water and a standard chow diet. For genotyping, the wild-type allele and point mutation allele —which codes alanine instead of tyrosine at H2AX residue 142 were distinguished via the following forward primers: wild-type allele, 5' CGC AGG CCT CTC AGG AGT AC 3'; mutant allele, 5' CGC AGG CCT CTC AGG AGG CT 3'. The reverse primer was 5' CTG CGG AGG GAC TAA CCT TC 3'. Heterozygous males and females were bred to produce homozygous mutant males (i.e., H2ax-Y142A mice). The *Mdc1* knockout (*Mdc1*KO) mice used in this study are described in our previous report [6]. For analyses of *H2ax-Y142A* and *Mdc1*KO mice, homozygous mice bearing wild-type alleles were used as littermate controls unless otherwise described in the figures and/or figure legends. All mouse tissue samples used in this study were harvested at 5-12 weeks of age unless otherwise noted in the figures and/or figure legends. For analysis of female meiocytes, fetal ovaries were harvested at day 17.5 of pregnancy (where observation of vaginal plug is counted as day 0.5). All subsequent experimental work was performed under protocol no. IACUC2018-0040 approved by the Institutional Animal Care and Use Committee of Cincinnati Children's Hospital and Medical Center.

METHOD DETAILS

Western blotting—The following western blot experiments were replicated twice; for each experiment, all samples were run on the same gel. To confirm the absence of phosphorylated Y142 of H2AX via western blots, detunicated testis pieces were

homogenized in RIPA buffer (50 mM Tris-HCl pH 7.5, 150 mM NaCl, 0.1% SDS, 1% Triton X-100, 1% sodium deoxycholate) containing cOmplete Protease Inhibitor Cocktail (Roche, 11697498001) and Phosphatase Inhibitor Cocktail 2 (Sigma, P5726); then, the homogenate was incubated on ice for 30 min. After DNA fragmentation by sonication and subsequent centrifugation at $10,000 \times g$ at 4°C, the supernatant was transferred to a new tube before total protein concentration was quantified via Bradford assays. Volumes of lysates containing 20 µg proteins were separated by electrophoresis on 10% SDS-PAGE gels. Then, the proteins were transferred onto a PVDF membrane (EMD Millipore, IPVH00010). The membranes were blocked with StartingBlockTM T20 (TBS) Blocking Buffer (ThermoFisher Scientific, 37543) at room temperature (RT) for 30 min before incubation with anti-H2AXpY142 antibody (Millipore, 07–1590), diluted 1/1000 in Tris-buffered saline containing 0.1% Tween 20 detergent (TBST), at 4°C overnight. On the next day, after washing three times in TBST, 5 min per wash, the blot was incubated with VeriBlot for IP Detection Reagent (HRP) (Abcam, ab131366), diluted 1/5000 in TBST at RT, for 1 h. The blot was washed three times in TBST, 5 min per wash, before incubation in Immobilon Western Chemiluminescent HRP Substrate (EMD Millipore, WBKLS0500) at RT for 1 min; then, the blot was imaged using Super RX-N x-ray film (Fujifilm) and a FluorChemQ MultiImage III instrument (Alpha Innotech). To blot loading controls, the initial blot was stripped with Restore Western Blot Stripping Buffer (ThermoFisher Scientific, 21059) at RT for 10 min; then, the stripped blot was washed two times in TBST, 5 min per wash, prior to incubation with anti-Lamin B1 antibody (Abcam, ab16048), diluted 1/2000 in TBST, at RT for 1 h. After washing the blot three times in TBST, 5 min per wash, the blot was incubated with VeriBlot for IP Detection Reagent (HRP), diluted 1/5000 in TBST, at RT for 1 h, and bands were visualized through the procedures described above.

To detect H2AX protein expression in wild-type and H2ax-Y142A mouse heart, liver, and testis tissues, whole tissue lysates were isolated and prepared from mice at two weeks of age. We selected this timepoint because spermatogenesis is in its semi-synchronized "first wave;" thus, the vast majority of testis cells are early pachytene spermatocytes. To prepare tissue lysates, ~ 20 mg of heart tissue, ~ 20 mg of liver tissue, and two whole testes were processed as described above. Volumes of lysates containing 20 µg proteins were separated by electrophoresis on a 15% SDS-PAGE gel. After transferal to a PVDF membrane and blocking, the membrane was cut into two pieces and trimmed. One piece was used for the detection of H2AX, and the other piece was used for the detection of a-tubulin, a loading control. Thus, one portion of the split membrane was incubated with anti-H2AX antibody (Cell signaling technology, 2595), diluted 1/2000 in TBST; and the other portion was incubated with anti-a-tubulin antibody (Abcam, ab7291), diluted 1/5000 dilution in TBST. The membranes were incubated at 4°C overnight. The next day, after washing (described above), the membranes were incubated with VeriBlot for IP Detection Reagent (HRP; described above). After additional washing (described above), the membranes were incubated with Immobilon Western Chemiluminescent HRP Substrate (described above); then, the blots were imaged using an Amersham Imager 680 (GE Healthcare).

Preparation and immunofluorescence of meiotic chromosome spreads-

Meiotic chromosome spreads from testes were prepared essentially as described [42, 86];

here, we briefly describe our protocol, including a small number of deviations from the cited literature. Testes were excised, detunicated, and placed in $1 \times$ phosphate-buffered saline (PBS). Seminiferous tubules were dissociated from whole testes in the following amounts: from wild-type or heterozygous models, approximately one-quarter of an adult testis; from an experimental model—i.e., H2ax-Y142A or Mdc1KO—a whole adult testis. Seminiferous tubules were transferred to a four-well dish (Nunc 4-Well Dishes: ThermoFisher Scientific, 144444) kept on ice. Three of the four wells contained 1 mL PBS, while the fourth well contained 1 mL hypotonic extraction buffer [HEB: 30 mM Tris base, 17 mM trisodium citrate, 5 mM ethylenediaminetetraacetic acid (EDTA), 50 mM sucrose, 5 mM dithiothreitol (DTT), 1× cOmplete Protease Inhibitor Cocktail (Sigma, 11836145001), 1× phosphatase inhibitor cocktail 2 (Sigma, P5726–5ML), pH. 8.2]. In the first well containing 1 mL PBS, seminiferous tubules were gently unraveled into small clumps with fine-point tweezers; care was taken not to tear or mince the tubules. The clumps of seminiferous tubules were transferred to the second and third wells of 1 mL PBS for additional unraveling before transfer to the fourth well containing 1 mL HEB. Once there, fine-point tweezers were used to carefully expose the surface area of tubules to HEB. The seminiferous tubules were incubated in HEB on ice for approximately 2 h with gentle stirring every 30-45 min. After incubation, a small clump of seminiferous tubules-approximately four-to-six seminiferous tubules—was gently pulled and mashed between tweezer tips in 30 µL of sucrose (100 mM) on a plain, uncharged microscope slide (Gold Seal: ThermoFisher Scientific, 3010-002). After approximately 15–25 mashes, a semi-translucent cell suspension was formed. An additional 30 μ L of sucrose (100 mM) was added to the suspension, and the suspension was mixed via gentle pipetting up and down several times. The diluted cell suspension was applied to positively charged slides (Probe On Plus: ThermoFisher Scientific, 22-230-900) in $30 \,\mu\text{L}$ volumes; before application of the suspension, the slides had been incubating in chilled fixation solution (2% paraformaldehyde, 0.1% Triton X-100, 0.02% sodium monododecyl sulfate, adjusted to pH 9.2 with sodium borate buffer) for a minimum of 2 min. After applying the cell suspension/sucrose mixture, the slide was slowly, gently tilted up and down at slight angles (<10°) to mix the cell suspension/sucrose mixture with remaining fixation solution. The slides were placed in "humid chambers" (closed pipet tip boxes filled to approximately two-thirds volume with water) at RT for a minimum of 1 h (maximum overnight). Then, the slides were washed in a low-concentration surfactant, 0.4% Photo-Flo 200 (Kodak, 146-4510), at RT two times, 2 min per wash. Slides were dried completely at RT (~30 min) before staining or storage in slide boxes at -80°C.

Chromosome spreads of oocytes at embryonic day 17.5 were prepared as described [42]. Briefly, fetal ovaries at embryonic day 17.5, which contain oocytes in the midst of meiotic prophase, were harvested. Ovaries were in incubated in HEB on ice for 45–60 min with gentle stirring every 15 min. Then, a suspension of oocytes was generated by pulling and mashing an ovary between tweezer tips in 30 μ L of sucrose (100 mM) on an uncharged microscope slide. 30 μ L of the suspension was applied to positively charged slides and all subsequent steps are the same as those described in the previous paragraph.

For immunostaining experiments, testis and ovary chromosome spreads were incubated in PBS containing 0.1% Tween 20 (PBST) for 5–30 min before blocking in antibody dilution buffer (PBST containing 0.15% BSA) for an additional 30–60 min. Primary and secondary

antibodies (described below) were diluted in antibody dilution buffer. Then, chromosome spreads were coated with 100 µL of the antibody/antibody dilution buffer solution, gently covered with Parafilm (Parafilm M All-Purpose Laboratory Film, Bemis Company, Inc.), and stored in "humid chambers" (described in [87]) at RT or 4°C for a minimum timespan of 6 h to a maximum timespan of overnight (~15 h). This study made use of the following primary antibodies at the following dilutions [format: host anti-protein (source or company with product/catalog number if applicable), dilution]: rabbit anti-H2AX-pY142 (Millipore, 07-1590), 1/200; rabbit anti-SYCP3 (Novus, NB300-232), 1/500; mouse anti-H2AX-pS139 (yH2AX: Millipore, 05–636), 1/5000; rabbit anti-SYCP1 (Abcam, ab15090), 1/200; mouse anti-SYCP3 (Abcam, ab97672), 1/5000; rabbit anti-TOPBP1 (gift from Dr. Junjie Chen [88]), 1/2000; rabbit anti-BRCA1 (generated in the Namekawa Lab [6]), 1/500; sheep anti-MDC1 (Bio-Rad, AHP799), 1/500; rabbit anti-RAD51 (Millipore, PC130), 1/50; guinea pig anti-H1T (gift from Dr. Mary Ann Handel [19]), 1/2000; rabbit anti-ATR (Millipore, PC538), 1/2000; rabbit anti-SIX6OS1 (gift from Dr. Alberto M. Pendás [89]), 1/4000; rabbit anti-HORMAD2 (gift from Dr. Attila Tóth [90]), 1/800; rabbit anti-MLH3 (gift from Dr. Paula E. Cohen [43]), 1/1000; rabbit anti-MLH1 (Santa Cruz, sc-11442), 1/500; mouse anti-SYCP3 conjugated with Alexa 488 fluorophore (Abcam, ab205846), 1/500 in Figure 5D, 5E; mouse anti-H2AX-pS139 (yH2AX) conjugated to Alexa 647 fluorophore (Millipore, 05–636-AF647), 1/500 in Figure 5D, 5E; rabbit anti-ATRIP (gift from Dr. Stephen J. Elledge [48]), 1/100. After incubation of the primary antibodies, slides were washed three times in PBST, 5 min per wash. Then, the slides were incubated with appropriate secondary antibodies conjugated to Alexa 488, 555, and/or 647 fluorophores (ThermoFisher Scientific). All secondary antibodies were diluted 1/500 in antibody dilution buffer. Slides were coated with 100 μ L of the antibody/antibody dilution buffer solution; then, they were gently covered with Parafilm for 1-h incubation at RT in humid chambers in darkness. After slides were washed three times in PBST in darkness, 5 min per wash, they were counterstained with the DNA-binding chemical 4',6-diamidino-2-phenylindole (DAPI; Sigma, D9542–5MG) diluted to 1 µg/mL concentration in PBS. Finally, slides were mounted using 20 µL undiluted ProLong Gold Antifade Mountant (ThermoFisher Scientific, P36930). Slides were either imaged immediately or stored at 4°C in darkness. For long-term storage, stained slides were kept at 4°C in darkness.

Images were obtained with an ECLIPSE Ti-E microscope (Nikon) equipped with a Zyla 5.5 sCMOS camera (Andor Technology) and an 60× CFI Apochromat TIRF oil immersion objective NA 1.4 (Nikon), and were processed with NIS-Elements Basic Research (Nikon), Photoshop (Adobe), and Illustrator (Adobe).

3D slide preparation and immunostaining—To conserve the gross conformation of meiotic chromatin, specialized slides that preserve the relative three-dimensional nuclear organization of testis germ cells (3D slides) were prepared as described [33]. Briefly, seminiferous tubules underwent permeabilization in cytoskeleton buffer [CSK buffer: 100 mM sodium chloride, 300 mM sucrose, 10 mM 1,4-Piperazinediethanesulfonic acid (PIPES), 3 mM magnesium chloride, 0.5% Triton X-100], fixation by 4% paraformaldehyde in PBS, and then mechanical dissociation with fine-point tweezers before being cytospun via specialized centrifuge (Shandon, Cytospin 2) onto positively charged slides (Superfrost Plus:

ThermoFisher Scientific, 12-550-15). Immunostaining was performed following procedures described above. Briefly, 3D slides were blocked with antibody dilution buffer at RT for 30 min, then the following primary antibodies were diluted in antibody dilution buffer and applied to the slides: rabbit anti-BRCA1 antibody (generated in the Namekawa Lab), diluted 1/2000; mouse anti-RNA Polymerase II antibody (Millipore, 05–952), diluted 1/200. 3D slides were incubated at 4°C overnight in a humid chambers. On the following day, 3D slides were washed three times in PBST, 5 min per wash, before incubation with secondary antibodies conjugated to Alexa 488 or Alexa 555 fluorophores (ThermoFisher Scientific); the secondary antibodies were diluted 1/500 in antibody dilution buffer, then incubated at RT for 1 h in darkness. After washing the 3D slides three times in PBST, 5 min per wash, they were counterstained with DAPI as described above and then mounted using 20 μ L undiluted ProLong Gold Antifade Mountant (ThermoFisher Scientific, P36930). Slides were imaged immediately thereafter.

Images were obtained with an ECLIPSE Ti-E microscope (Nikon) equipped with a Zyla 5.5 sCMOS camera (Andor Technology) and an 100× CFI Apochromat TIRF oil immersion objective NA 1.4 (Nikon), and were processed with NIS-Elements Basic Research (Nikon), Photoshop (Adobe), and Illustrator (Adobe).

Immunohistochemistry and TUNEL assay—For the preparation of testis paraffin blocks, excised testes in tunicae albugineae were fixed with 4% paraformaldehyde at 4°C overnight. Testes were dehydrated and embedded in paraffin. For histological analyses, 6 um-thick paraffin sections were deparaffinized. Standard periodic acid-Schiff staining was performed; for immunostaining, sections were autoclaved in Target Retrieval Solution, Citrate pH 6.1 (DAKO, S-1700), 121°C, 100 kPa (15 psi) for 10 min. The sections were blocked with Blocking One Histo (Nacalai USA, 06349-64) at RT for 10 min; then, the sections were incubated with primary antibodies diluted in PBS at 4°C overnight. The following antibodies were used in the following dilutions [format: host anti-protein (source or company with product/catalog number), dilution]: mouse anti- γ H2AX (Millipore, 05– 636), 1/2500; guinea pig anti-H1T (gift from Dr. Mary Ann Handel), 1/2500; rabbit anti-BRCA1 (generated in the Namekawa Lab), 1/2000. The resulting signals were detected with appropriate secondary antibodies conjugated to Alexa 488 and 555 (ThermoFisher Scientific), diluted 1/1000 in PBS and incubated at RT for 1 h. TUNEL assays were performed using an In Situ Cell Death Detection Kit (Roche, 11684795910) per the instructions in the manual. Sections were counterstained with DAPI as described above. Images were obtained with an A1RSi Inverted Confocal Microscope (Nikon) and processed with NIS-Elements Basic Research (Nikon), Photoshop (Adobe), and Illustrator (Adobe).

QUANTIFICATION AND STATISTICAL ANALYSIS

The details for statistical analyses performed in this study are described in relevant portions of the Results section, figures, figure legends, supplementary figures, and/or supplementary figure legends. Sample sizes used for analyses are described in relevant portions of the Results section, figures, figure legends, supplementary figures, and/or supplementary figure legends. In predetermining sample sizes, we sought to analyze a minimum of three independent control-mutant littermate pairs except for western blot experiments, in which

we analyzed two independent control-mutant littermate pairs; no statistical calculations were used to predetermine sample sizes. No data were excluded from analyses. The experiments were not randomized, and investigators were not blinded to allocation during experiments and outcome assessment. Measurements were recorded in Excel (Microsoft) and Prism 8.0 (GraphPad). Statistical tests were performed with Prism 8.0 (GraphPad).

Quantification of RAD51, TOPBP1, and MLH3 foci in the pachytene stage-To carefully detail the phenotypes of MSCI-deficient mice, we scored foci numbers for the proteins RAD51, TOPBP1, and MLH3 in wild-type and mutant spermatocytes in the initial portion of the mid pachytene stage. In H2ax-Y142A mice, spermatogenesis is arrested in epithelial stages IV, when tubules bear H1T-positive spermatocytes that have just entered the mid pachytene stage of meiotic prophase I. Thus, in our analyses of mutant meiotic chromosome spreads, we judged nuclei to be in the initial portion of the mid pachytene stage based on H1T staining-i.e., H1T-positive H2ax-Y142A nuclei were considered to be mid pachytene spermatocytes. However, we did not rely solely on H1T staining for identification of wild-type mid pachytene spermatocytes: Since all spermatocytes at and after the mid pachytene stage—as well as all spermiogenic cells—are H1T-positive, an H1T-positive status alone is insufficient for this task. To make direct comparisons with H1T-positive H2ax-Y142A spermatocytes, we precisely identified wild-type spermatocytes in the initial portion of the mid pachytene stage drawing on techniques described in our recent chapter in the Methods in Molecular Biology series (SpringerProtocols, Humana Press) [87]. Briefly, we focused on the following chromo-nuclear features to stage spermatocytes in the initial portion of the mid pachytene stage: (1) After immunostaining (described above), spermatocyte nuclei exhibited positive-but-weak H1T signal intensity relative to late pachytene and diplotene spermatocyte nuclei; (2) the pseudoautosomal region (PAR) of the sex chromosomes was clearly visible. After determining the substages of pachytene spermatocytes, we scored clearly visible RAD51, TOPBP1, and/or MLH3 foci overlapping SYCP3-labeled chromosomes axes.

Quantification of XY chromatin compaction—To evaluate chromatin compaction of the sex chromosomes, 3D slides stained with antibodies raised against BRCA1 and H1T, and counterstained with DAPI, were used as described [6]. Wild-type and *H2ax-Y142A* H1T-positive mid pachytene spermatocytes were evaluated. After obtaining 10 z-sections (0.2 µm interval) for each spermatocyte, we selected a z section containing in-focus XY centromeres for analysis. To quantify chromatin compaction of the sex chromosomes, a straight line was drawn to represent the widest spans of distance between the axial ends of the sex chromosomes as detected via anti-BRCA1 immunostaining (a) in H1T-positive nuclei; then, the widest spans were divided by the diameters of the nuclei (b). Distances were measured using NIS-Elements Basic Research (Nikon). The relative distances (i.e., the a/b distance ratios)in Figure 3 represent distances between XY centromeres with respect to nuclear diameters.

RNA-seq analysis—RNA-seq datasets were obtained from published work (GEO GSE55060) [70]. .fastq files were aligned to the *Mus musculus* mm10 reference genome via Spliced Transcripts Alignment to a Reference (STAR; version 2.4.2a)[91]; only unique

alignments were allowed with a maximum of two errors per alignment. Datasets were processed and visualized through the BioWardrobe Experiment Management Platform [92].

DATA AND CODE AVAILABILITY

The RNA-seq datasets analyzed in this study are available at NCBI Gene Expression Omnibus (GEO; https://www.ncbi.nlm.nih.gov/geo/) under accession number GSE55060. Software used in this study is described in context under portions of the METHOD DETAILS and/or QUANTIFICATION AND STATISTICAL ANALYSIS headings, and in the Key Resources Table. Software used for this study: NIS-Elements Basic Research (Nikon), Excel (Microsoft), Prism 8.0 (GraphPad), and the BioWardrobe Experiment Management Platform (biowardrobe.com, https://github.com/Barski-lab/biowardrobe) [92]. The manuscript was constructed using Word (Microsoft), Photoshop (Adobe), and Illustrator (Adobe).

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgements

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Highlights

The Y142 residue of H2AX is essential for MSCI and XY-body formation.

The initial steps of autosomal DSB repair are normal in MSCI-defective mutants.

ATR-associated DDR signals are sequestered from autosomes at the onset of MSCI.

Novel model for the MSCI checkpoint: MSCI permits meiotic progression.



$Figure \ 1. \ H2AX \text{-} Y142 \ is \ an \ essential \ amino \ residue \ for \ completing \ spermatogenesis.$

(A) Schematic: Induction of a point mutation and resulting sequence alteration. The introduction of an AfIII target site was used to screen mutant mice.

(**B**) Wild-type (WT) and *H2ax-Y142A* mice at 36 days post-partum (dpp). Body weights are shown as mean \pm s.e.m. from 4 independent pairs of WT and *H2ax-Y142A* mice. ** p < 0.01, unpaired t test.

(C) Testes from WT and *H2ax-Y142A* mice at 87 dpp. Scale bar: 1 cm. Ratios of testis weight (mg) to body weight (g) shown as mean \pm s.e.m. for 4 independent pairs of WT and *H2ax-H142A* mice. **** p < 0.0001, unpaired t test.

(**D**) Western blot of WT and *H2ax-Y142A* lysates from whole mouse testes with anti-H2AX-pY142 antibody. 20 µg protein samples were loaded in each lane. Two independent samples for WT and *H2ax-Y142A* are shown. Loading control: Lamin B1.

(E) Chromosome spreads of WT and *H2ax-Y142A* pachytene spermatocytes immunostained with antibodies raised against SYCP3 and H2AX-pY142. Dashed circles indicate the sex chromosomes. Scale bars: 10 µm.

(F) Testis sections from WT and *H2ax-Y142A* mice immunostained with antibodies raised against H1T and γ H2AX. Dashed squares are magnified in panels to the right. White arrowheads indicate γ H2AX signals on sex chromosomes in mid pachytene (H1T-positive) spermatocytes. Scale bars in larger panels: 100 µm; scale bars in smaller panels: 10 µm. See also Figure S1.



Figure 2. H2AX-Y142 is required for the initiation of MSCI

(A, C-H) Chromosome spreads of wild-type (WT) and *H2ax-Y142A* pachytene spermatocytes immunostained with antibodies raised against the following proteins: SYCP3
(A, C-H), γH2AX (A), SYCP1 (C), ATR (D), TOPBP1 (E), BRCA1 (F), HORMAD2 (G), and MDC1 (H). Dashed squares are magnified in the panels to the right.
(B) Model of the initiation of MSCI. ATR and its activator, TOPBP1, are recruited to unsynapsed axes in an MDC1- and Y142-H2AX-independent manner, resulting in phosphorylation of H2AX to generate γH2AX on axes (left). γH2AX then recruits MDC1,

which facilitates progressive recruitment of ATR and TOPBP1, resulting in γ H2AX and MDC1 spreading throughout loops (right).

(C) Percentages of mid pachytene chromosome synapsis for autosome and sex chromosomes, shown as mean \pm s.e.m.. Total numbers of analyzed nuclei obtained from 3

independent littermate pairs are indicated in the panel. n.s., not significant; * p < 0.05, unpaired t test. Scale bars: 10 μ m unless otherwise described in the panels. See also Figure S2.



Figure 3. Impaired XY body formation in H2ax-Y142A mouse.

(A) Wild-type (WT) littermate control and *H2ax-Y142A* pachytene spermatocytes on 3D slides (see STAR METHODS) immunostained with antibodies raised against RNAPII and BRCA1. Although not shown in the panel, the spermatocytes were also immunostained with an anti-H1T antibody to determine their stage; the spermatocytes shown are H1T-positive. Dashed circles indicate the sex chromosomes.

(**B**) WT littermate control and *H2ax-Y142A* testis sections immunostained with antibodies raised against BRCA1 and H1T. Dashed squares are magnified in the panels to the right. White arrowheads indicate the axes of sex chromosomes in H1T-positive pachytene spermatocytes. Nuclei were counterstained with DAPI. Scale bars: 100 μm and, in the panels to the right, 10 μm.

(C) WT and *H2ax-Y142A* pachytene spermatocytes on 3D slides immunostained with an antibody raised against BRCA1. Although not shown in the panel, the spermatocytes were also immunostained with an anti-H1T antibody to determine their stage; the spermatocytes shown are H1T-positive. The relative distances are shown in a box-and-whisker plot: The central line is the median, the bottom edge of the box is the first quartile, the top edge of the box is the third quartile, and the whiskers encompass, from top to bottom, the first to ninth decile. Total numbers of analyzed nuclei, obtained from 3 independent wild-type mice and 4 independent *H2ax-Y142A* mice, are indicated in the panel. **** p < 0.0001, Mann-Whitney U test. Nuclei were counterstained with DAPI. Scale bars: 10 µm.



Figure 4. Initial steps of DSB repair take place normally on autosomes of MSCI defective mutants.

(A, B, E, F) Chromosome spreads of wild-type (WT) littermate control and *H2ax-Y142A* (A, B) or *Mdc1*KO (E, F) mid pachytene spermatocytes immunostained with antibodies raised against SYCP3 (A, B, E, F), HIT (A, B, E, F), RAD51 (A, E), or MLH3 (B, F). Sex chromosomes in dashed squares are magnified in the panels to the right (A, E). Autosomes in dashed squares are magnified in the panels to the right, and the dashed circles indicate the sex chromosomes (B, F). Dot plots indicate the numbers of autosome RAD51 foci (A, E), sex chromosome RAD51 foci (A, E), or MLH3 foci (B, F) per mid pachytene (HIT-positive) spermatocyte, shown as mean \pm s.e.m. for 3 independent *H2ax-Y142A* littermate pairs (A, B) and 3 independent *Mdc1*KO littermate pairs (E, F). Total numbers of analyzed nuclei are indicated in the panels. n.s.: not significant; **** p < 0.0001, unpaired t tests. XY chr.: XY chromosomes. Scale bars: 10 µm.

(C) Model of the MSCI checkpoint and its relationship to meiotic recombination.

(**D**) DAPI counterstaining of a 3D slide (see STAR METHODS). The dashed circle indicates the XY body. Scale bar: 10 μ m. See also Figures S3 and S4.



Figure 5. DDR factors centered on ATR signaling are sequestered from autosomes to the sex chromosomes at the onset of MSCI.

(A, B, D, E) Chromosome spreads of wild-type (WT) littermate control and *H2ax-Y142A* (A, D) or *Mdc1*KO (B, E) mid pachytene spermatocytes immunostained with antibodies raised against SYCP3 (A, B, D, E), TOPBP1 (A, B), H1T (A, B), γ H2AX (D, E), and ATR (D, E). Yellow arrowheads indicate the sex chromosomes, and red arrowheads indicate ATR foci that persist on *H2ax-Y142A* autosomes (D, E). Scale bars: 10 µm.

(C) Numbers of TOPBP1 foci on autosomes in mid pachytene (H1T-positive)

spermatocytes, shown as mean \pm s.e.m. for 3 independent *H2ax-Y142A* littermate pairs (left) and 3 independent *Mdc1*KO littermate pairs (right). Total numbers of analyzed nuclei are indicated in the panels. **** p < 0.0001, Mann-Whitney U test. See also Figure S5.



Figure 6. A model of the MSCI checkpoint: The physical sequestration of DDR factors from autosomes to the XY body is a critical checkpoint in meiosis progression and gamete development.

At the onset of MSCI, DDR factors (shown as red spheres) are sequestered from autosomes to the sex chromosomes. The physical sequestration of DDR factors on/at a sex chromosome-specific chromo-nuclear compartment, the XY body, is a critical step in the MSCI checkpoint in the mid pachytene stage of meiotic prophase I. While the MSCI checkpoint ensures meiotic stage progression in normal meiosis, the abolishment of MSCI enables the ectopic retention of DDR signals on/at autosomes; these, in turn, trigger complete meiotic arrest and cell death in response to the checkpoint. See also Figure S6.

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies	•	•
Rabbit Anti-Histone H2A.X, phospho (Tyr142)	Millipore	Cat# 07-1590, RRID:AB_1977237
Rabbit Anti-Lamin B1	Abcam	Cat# ab16048, RRID:AB_443298
Mouse Anti-alpha Tubulin [DM1A]	Abcam	Cat# ab7291, RRID:AB_2241126
Rabbit Anti-Histone H2A.X	Cell Signaling Technology	Cat# 2595, RRID:AB_10694556
Mouse Anti-SYCP3	Abcam	Cat# ab97672, RRID:AB_10678841
Guinea pig Anti-H1t	[19]	N/A
Mouse Anti-Histone H2A.X, phospho (Ser139)	Millipore	Cat# 05-636, RRID:AB_309864
Rabbit Anti-SYCP1	Abcam	Cat# ab15090, RRID:AB_301636
Rabbit Anti-TOPBP1	[88]	N/A
Rabbit Anti-BRCA1	[6]	N/A
Sheep Anti-Human MDC1	Bio-Rad	Cat# AHP799, RRID:AB_323725
Mouse Anti-RNA Polymerase II, C-Terminus Domain	Millipore	Cat# 05–952, RRID:AB_492629
Rabbit Anti-Rad51	Millipore	Cat# PC130, RRID:AB_2238184
Rabbit Anti-ATR	Millipore	Cat# PC538, RRID:AB_2063178
Mouse Anti-SCP3 conjugated with Alexa Fluor 488	Abcam	Cat# ab205846, RRID: N/A
Mouse Anti-phospho Histone H2A.X (Ser139) conjugated with Alexa Fluor 647	Millipore	Cat# 05–636-AF647, RRID: N/A
Rabbit Anti-MLH3	[43]	N/A
Rabbit Anti-MLH1(H-300)	Santa Cruz Biotechnology	Cat# sc-11442, RRID:AB_2145332
Rabbit Anti-ATRIP	[48]	N/A
Rabbit Anti- SIX6OS1	[89]	N/A
Rabbit Anti-HORMAD2	[90]	N/A
VeriBlot for IP Detection Reagent (HRP)	Abcam	N/A
Donkey Anti-Sheep IgG (H+L) Alexa Fluor 647	Jackson ImmunoResearch Labs	Cat# 713-606-147, RRID:AB_2340752
Goat Anti-Guinea Pig IgG (H+L) Alexa Fluor 555	Thermo Fisher Scientific	Cat# A-21435, RRID:AB_2535856
Donkey Anti-Mouse IgG (H+L) Alexa Fluor 555	Thermo Fisher Scientific	Cat# A-31570, RRID:AB_2536180
Donkey Anti-Mouse IgG (H+L) Alexa Fluor 488	Thermo Fisher Scientific	Cat# A-21202, RRID:AB_141607
Donkey Anti-Rabbit IgG (H+L) Alexa Fluor 555	Thermo Fisher Scientific	Cat# A-31572, RRID:AB_162543
Donkey Anti-Rabbit IgG (H+L) Alexa Fluor 488	Thermo Fisher Scientific	Cat# A-21206, RRID:AB_2535792
Donkey Anti-Sheep IgG (H+L) DyLight 488	Jackson ImmunoResearch Labs	Cat# 713-486-147, RRID:AB_2340741
Goat Anti-Rabbit IgG (H+L) antibody F(ab')2 Fragment Cy3	Jackson ImmunoResearch Labs	Cat# 111-166-003, RRID:AB_2338007
Donkey Anti-Guinea Pig IgG (H+L) antibody F(ab')2 Fragment Alexa Fluor 647	Jackson ImmunoResearch Labs	Cat# 706-606-148, RRID:AB_2340477
Chemicals, Peptides, and Recombinant Proteins		
Phosphatase inhibitor cocktail 2	Sigma	Cat# P5726–5ML
cOmplete [™] Protease Inhibitor Cocktail	Sigma	Cat# 11836145001

REAGENT or RESOURCE	SOURCE	IDENTIFIER
StartingBlockTM T20 (TBS) Blocking Buffer	ThermoFisher Scientific	Cat# 37543
Restore TM Western Blot Stripping Buffer	ThermoFisher Scientific	Cat# 21059
Critical Commercial Assays		
In Situ Cell Death Detection Kit, Fluorescein	Sigma	Cat# 11684795910
Deposited Data		
RNA-seq data (Scml2KO)	[70]	GEO: GSE55060
Experimental Models: Organisms/Strains		
Mouse: H2ax-Y142A	This study	N/A
Mouse: Mdc1KO	[52]	N/A
Oligonucleotides		
Primer: H2ax-Y142A WT Forward: CGC AGG CCT CTC AGG AGT	This study	N/A
Primer: H2ax-Y142A KI Forward: CGC AGG CCT CTC AGG AGG CT	This study	N/A
Primer: H2ax-Y142A Common Reverse: CTG CGG AGG GAC TAA CCT TC	This study	N/A
sgRNA for generating H2ax-Y142A: GCCTCTCAGGAGTACTGAGG	This study	N/A
Software and Algorithms		-
BioWardrobe Experiment Management Platform	[92]	https://biowardrobe.cchmc.org/frontend/ Home
Prism 8	GraphPad software	https://www.graphpad.com/