

The ssDNA-binding protein MEIOB acts as a dosage-sensitive regulator of meiotic recombination

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ABSTRACT

Meiotic recombination enables reciprocal exchange of genetic information between parental chromosomes and is essential for fertility. MEIOB, a meiosis-specific ssDNA-binding protein, regulates early meiotic recombination. Here we report that the human infertility-associated missense mutation (N64I) in MEIOB causes protein degradation and reduced crossover formation in mouse testes. Although the MEIOB N64I substitution is associated with human infertility, the point mutant mice are fertile despite meiotic defects. *Meiob* mutagenesis identifies serine 67 as a critical residue for MEIOB. Biochemically, these two mutations (N64I and S67 deletion) cause self-aggregation of MEIOB and sharply reduced protein half-life. Molecular genetic analyses of both point mutants reveal an important role for MEIOB in crossover formation in late meiotic recombination. Furthermore, we find that the MEIOB protein levels directly correlate with the severity of meiotic defects. Our results demonstrate that MEIOB regulates meiotic recombination in a dosage-dependent manner.

INTRODUCTION

Meiotic recombination, a hallmark of meiosis, is essential for faithful segregation of homologous chromosomes during the first meiotic cell division and enhances genetic diversity in gametes (1,2). This process begins with the formation of a large number of programmed DNA double strand breaks (DSBs) in germ cells (~300 per cell in mouse), which are generated by SPO11 (3,4). DSBs are resected to produce 3' single-stranded DNA (ssDNA) overhangs (2). One DSB ssDNA end invades the homologous chromosomes to produce the displacement loop (D-loop). The capture of the second DSB end results in the formation of double Holliday junctions (dHJs). Resolution of dHJs produces crossovers

(~22 crossovers per cell in mouse). Failures in meiotic recombination entail blockade in meiosis and thus sterility.

Meiotic recombination is temporally executed by a large number of proteins including ssDNA-binding proteins (1,2). These proteins typically localize to DSBs as foci on meiotic chromosomes. RPA, consisting of RPA1, RPA2 and RPA3, is a ubiquitously expressed ssDNA-binding complex (5,6). RPA coats ssDNA to prevent formation of secondary structures and protect ssDNA from degradation. Loss of RPA1 causes a failure in meiotic recombination (7). RAD51 and DMC1, ssDNA-binding recombinases, bind to DSB ssDNA to drive strand invasion into homologues, resulting in the formation of D-loops (8–12). Hop2-Mnd1 proteins stimulate the RAD51/DMC1 recombinase activity (13). HSF2BP is a meiosis-specific localizer of BRCA2, which in turn recruits RAD51/DMC1 to DSBs (14,15). RPA interacts with RAD51 and DMC1, suggesting that RPA may recruit RAD51/DMC1 to meiotic DSB ssDNA (16). This notion is supported by the loss of RAD51/DMC1 foci in RPA1-deficient spermatocytes (7). MEIOB, an ssDNA-binding protein, is a meiosis-specific paralogue of RPA1 (17,18). MEIOB forms a heterodimer with SPATA22 and the MEIOB/SPATA22 heterodimer interacts with the RPA heterotrimer (17,19). The protein stability of MEIOB and SPATA22 depends on each other in both testis and transfected cells (17,19). When MEIOB is absent or reduced, SPATA22 is absent or reduced as well and vice versa. Although the MEIOB/SPATA22 complex and RPA colocalize as foci on meiotic chromosomes, their focal localizations are independent of each other (7,17). Unlike RPA, MEIOB/SPATA22 are not required for the recruitment of RAD51/DMC1 to DSB sites (17,18,20–23). Instead, MEIOB and SPATA22 appear to function in the second-end capture to stabilize recombination intermediates after strand invasion. Several proteins such as MSH4, MSH5 and TEX11 are involved in processing of recombination intermediates (24–26). In particular, the MSH4/MSH5 complex recognizes dHJ structures (24). The MLH1/MLH3 complex exhibits endonuclease activity and is required for for-

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mation of the majority of crossovers (27–30). Mus81 generates the remaining small subset of crossovers (31,32). Other proteins such as RNF212, HEI10 and CNTD1 also regulate crossover formation (33–36). Genetic studies have shown that lack of crossovers results in formation of univalent chromosomes, followed by chromosome mis-segregation and subsequent apoptosis of meiotic germ cells. In addition, HEI10, RNF212 and TEX11 function in crossover formation in a dosage-sensitive manner (33,37,38). This dosage sensitivity may have implications in modulation of genome-wide recombination rates.

Meiosis-specific proteins have been studied genetically in model organisms from yeast to mouse. However, our understanding of their role in human meiosis is limited (39). Mutations in several meiosis-specific genes have been found in humans with primary ovarian insufficiency or non-obstructive azoospermia (39). For example, a homozygous splice site mutation in human *MSH4* causes primary ovarian insufficiency (40). Mutations in the X-linked *TEX11* gene cause azoospermia in men (38,41). Several human studies reported homozygous mutations in the human *MEIOB* gene. A frame-shift mutation in *MEIOB* causes primary ovarian insufficiency (42). A different frame-shift *MEIOB* mutation is a recurrent cause of azoospermia in men (43). Interestingly, a homozygous missense mutation (N64I) in *MEIOB* was found in azoospermic men (44). In this study, we generated a mouse mutant to model the human *MEIOB* N64I mutation. Our biochemical studies reveal that the N64I mutation causes *MEIOB* self-dimerization and degradation. Unexpectedly, cell biological studies uncover a distinct role for *MEIOB* in crossover formation. Importantly, genetic studies of various combinations of *Meiob* mutant alleles demonstrate that *MEIOB* functions in meiotic recombination in a dosage-dependent manner.

MATERIALS AND METHODS

Generation of *Meiob*^{N64I} Mice

The *Meiob* N64I point mutation was generated by the CRISPR/Cas9-mediated genome editing (Supplementary Figure S1B). Guide RNA (sgRNA) was cloned into the px330 vector, in vitro transcribed and purified as described in the protocol (45). The sgRNA flanks codon 64 (underlined): ACTTTGTGAATGTATCCTCC. The single-stranded oligodeoxynucleotide (ssODN) template contains the N64I mutation (AAT to ATC): 5'-GGAATATAAAAGCGTTGGTTTTTCCTCAGATATTGGATCAGAAAGATACACTTTTCAGTTTCACCATTCGTACTCACCATGTCACCTTTGTGATCGTATCCTCCTGGGGCAGTGAGGACTACATCCGCTCGCTGTCCGAGAGCTTCAGGGTTGCTGAGTGTGGTAAGTAGGCAGGATTCTTTTGTGTTTGTG-3'. A mixture of Cas9 mRNA (Trilink, 100 ng/μl), sgRNA (50 ng/μl) and ssODN DNA (100 ng/μl) was used for microinjection of 50 zygotes on the DBA x C57BL/6 hybrid genetic background. Out of 20 founders, the following genotypes were obtained: 8 *Meiob*^{N64I/FS} (2 males, 6 females) (FS, frame-shift mutation), 2 *Meiob*^{N64I/N64I} (1 male, 1 female), 2 *Meiob*^{N64I/ΔS67} (1 male, 1 female), 2 *Meiob*^{ΔS67/ΔS67} (2 males) and 2 *Meiob*^{ΔS67/FS} (1 male,

1 female). Founder *Meiob*^{N64I/FS} females were mated to wild type C57BL/6J males to obtain heterozygous *Meiob*^{N64I/+} mice. Germline transmission of the point mutation was validated by PCR and sanger sequencing (PCR primers 5'-GCATGCAGAACCAAAGCTAAATAA-3' and 5'-GAGATAGGGTTTTACTATATAACA-3'; PCR product, 451 bp). The *Meiob*^{N64I} allele was genotyped by PCR (216 bp) with primers 5'-GACTCACCATGTCACTTTGTGATC-3' and 5'-GTGGCTGGATTGAACTGGTTATG-3'. The wild type *Meiob* allele was genotyped by PCR (216 bp) with primers 5'-GACTCACCATGTCACTTTGTGAAT-3' and 5'-GTGGCTGGATTGAACTGGTTATG-3'.

Generation of *Meiob*^{ΔS67} mice

The targeting vector was constructed as illustrated (Supplementary Figure S2C). The codon 67 deletion was introduced by overlapping PCR (46). The homologous arms were amplified from a *Meiob*-containing BAC clone (RP23–397O23) by PCR with high-fidelity DNA polymerase. V6.5 ES cells on the 129 × C57BL/6 hybrid genetic background were electroporated with the linearized targeting vector and cultured in the presence of 350 μg/ml G418 (postive selection) and 2 μM ganciclovir (negative selection). Homologously targeted ES clones were screened by long-distance PCR. Two targeted ES cell clones (1A12 and 1B11) were injected into B6C3F1 (Taconic) blastocysts. Male chimeras from both clones were mated with ubiquitous *Actb*-Cre females (FVB genetic background) to remove the floxed PGK-neo selection cassette and germline transmission were obtained from both clones (47). The codon 67 deletion was confirmed by sequencing. The wild type (397 bp) and *Meiob*^{ΔS67} (495 bp) alleles were genotyped by PCR with the following primers: 5'-CTTTCAGTTTCCATTTCGTGACT-3' and 5'-AAGGAAGTTACCTAGCAAGTCTCA-3'.

Histological, surface nuclear spread, and TUNEL analyses

For histological analysis, testes and ovaries were fixed in Bouin's solution overnight, embedded with paraffin, and sectioned. Sections were stained with hematoxylin and eosin. For TUNEL analysis, testes were fixed in 4% paraformaldehyde at 4°C overnight, dehydrated in sucrose, embedded in OCT and sectioned. Frozen sections were analyzed with the TUNEL Enzyme and Label Kit (Roche Boehringer Mannheim). Nuclear spread analysis of spermatocytes was performed as previously described (48). The primary antibodies used for immunofluorescence were listed in Supplementary Table S1.

Imaging

Color histological images were captured on a Leica DM5500B microscope with a DFC450 digital color camera (Leica Microsystems). Immunolabeled chromosome spreads and testis TUNEL assay images were captured with an ORCA Flash4.0 digital monochrome camera (Hamamatsu Photonics) on a Leica DM5500B microscope (Leica Microsystems) and processed using the Photoshop software (Adobe).

Sperm count

Cauda epididymis was dissected and placed in a small tissue dish with 2 ml of 1× PBS buffer without calcium/magnesium. We minced epididymis with forceps and squeezed out sperm as much as possible. Samples were incubated at room temperature for 10 min to allow sperm to disperse and transferred to a 10-ml falcon tube. 2 ml PBS was used to wash the dish and then added to the same tube. 2 ml 4% paraformaldehyde (in PBS) was added to the tube to fix at room temperature for 10 min. 10 μ l was loaded to the Hausser Bright-Line hematocytometer (Fisher Scientific). Sperm in the 25 small squares (a total volume of 0.1 μ l) were counted and the number was multiplied by 6×10^4 to give rise to the total sperm count per male.

Expression constructs

The *Meiob* and *Spata22* expression constructs were previously described (19). Mouse and human *Meiob* cDNAs were subcloned into the pcDNA6 vector (Invitrogen) harboring an N-terminal Flag tag and a C-terminal Myc tag. Mouse and human *Spata22* cDNAs were subcloned into the pcDNA6 vector harboring an N-terminal Flag tag. All *Meiob* mutations were generated by PCR and subcloned into the pcDNA6 vector. Green fluorescent protein (GFP)-*Meiob*, N64I, Δ S67, *Mlh1*, *Msh4* and *Rnf212* constructs were generated by subcloning into the pEGFP-C1 vector (Clontech). *Tex11-V5* construct was generated by subcloning into the pcDNATM3.1/V5-His vector (Invitrogen). All the constructs were verified by Sanger sequencing on an ABI 3730 DNA analyzer.

Cell culture and transfections

HEK293T cells were maintained in DMEM/high glucose (Mediatech) supplemented with 10% FBS (Sigma) and penicillin/streptomycin (Invitrogen). Plasmid DNA transfections in HEK293T cells were carried out using a standard calcium phosphate method. Briefly, transfections were performed when HEK293T cells are 50–90% confluent. To make 100 μ l plasmid mixture, add 2 μ g plasmid and 12.5 μ l 2 M CaCl₂. Mix the HEPES buffered saline solution (2× HEPES, pH 7.15, 100 μ l) by vortexing at 300 RPM, add dropwise to the plasmid solution, and incubate at room temperature for 30 min. The entire 200 μ l mixture was added directly into one well of a six-well plate in a dropwise manner. Cells were collected after 24–48 h.

Cycloheximide chase assays

HEK293T cells in one well of a six-well plate were transfected with 1 μ g of flag-mMEIOB-myc, flag-mMEIOB^{N64I}-myc, flag-mMEIOB ^{Δ S67}-myc, flag-hMEIOB-myc, flag-hMEIOB^{N64I}-myc or flag-mMEIOB^{IAA}-myc, incubated at 37°C for 24 h, and treated with cycloheximide (20 μ g/ml) for 0–24 h to inhibit *de novo* protein synthesis. Cells were collected in 2× SDS PAGE buffer for immunoblotting analysis with anti-flag antibody (Figure 7).

For protein degradation assays, HEK293T cells in one well of a six-well plate were transfected with 1 μ g of flag-mMEIOB-myc, flag-mMEIOB^{N64I}-myc or flag-mMEIOB ^{Δ S67}-myc, incubated at 37°C for 24 h, and treated

with cycloheximide (20 μ g/ml) in the presence of 20 μ M MG132 (M8699, Sigma) or 200 nM bafilomycin A1 (B1793, Sigma) for 0–24 h (Supplementary Figure S10).

Immunoprecipitation and immunoblotting assays

A total of 1×10^7 cells were harvested at 24–36 h post transfection and lysed in 1 ml RIPA buffer (140 mM NaCl, 1 mM EDTA, 0.5 mM EGTA, 10 mM Tris-HCl, pH 8.0, and 1% Triton X-100, 0.1% sodium deoxycholate, 0.1% SDS) supplemented with 1× protease inhibitor cocktail (Sigma). For immunoprecipitation experiments, the lysates were incubated on a rocking platform for 1 h at 4°C, followed by centrifugation at 16 100 g for 15 min at 4°C. The supernatant was transferred to a new tube, incubated with primary antibodies overnight, and then incubated with protein A or G dynabeads (Thermo Fisher Scientific) for 2–3 h. The immunoprecipitated complexes were washed with the RIPA buffer four times, boiled in 50 μ l 2× SDS-PAGE loading buffer for 12 min, and subjected to PAGE electrophoresis and western blotting. For Western blot analysis, testes or cells were homogenized in 500 μ l protein extraction buffer (62.5 mM Tris-HCl (pH 6.8), 3% SDS, 10% glycerol, 5% 2-mercaptoethanol). About 30 μ g of protein extracts were resolved by SDS-PAGE, transferred onto nitrocellulose membranes using iBlot (Invitrogen), and immunoblotted with indicated antibodies (Supplementary Table S1).

Quantitative RT-PCR assays

Total RNA was extracted from postnatal day 20 testes with Trizol reagent (Thermo Fisher Scientific). 1 μ g of total RNA was treated with DNaseI (Amp grade 1 U/ μ l, Invitrogen RNase free) and was reverse transcribed with PrimeScriptTM RT Master Mix (RR036A, TaKaRa, Japan). For quantitative RT-PCR, diluted cDNA was used for real-time PCR reactions containing SYBR Green Premix Ex Taq II (RR820A, TaKaRa). Gene expression was normalized to *Actb* (β -actin). The *Meiob* mRNA was assayed using primers 5'-CTGCATGGCAGCAACTGTAATCTC-3' and 5'-GCATAAAGGATGCCATAGAAAGGA-3' (exon 9–11). The *Spata22* mRNA was assayed using primers 5'-GCCTGTGCCATTGTTCAATCAGAA-3' and 5'-CGCCCAATCTGTTGAAAAGAAGG-3' (exon 3–4).

Statistics

All data were analyzed using GraphPad Prism (GraphPad Software Inc). All data are reported as mean \pm SD. Statistical analysis was performed with Student's *t*-test.

RESULTS

The human *Meiob*^{N64I} mutation destabilizes the MEIOB protein in testis

The human *MEIOB*^{N64I} missense mutation was found in azoospermic men with meiotic arrest (44). The N64 residue is conserved in MEIOB orthologues across species including *Drosophila*, *Xenopus*, and mouse (Supplementary Figure S1A). To study the requirement of this residue, we generated *Meiob*^{N64I} mutation in mice by CRISPR-mediated genome editing (Supplementary Figure S1B). In

contrast with the meiotic arrest observed in infertile men, the *Meiob*^{N64I/N64I} males had normal testis size, weight, and sperm count (Figure 1A–C), highlighting the species-specific requirement of this residue. We reduced the gene dosage by crossing this point mutant allele with the *Meiob* null allele, which was previously generated (17). Interestingly, the *Meiob*^{N64I/-} males exhibited significant reduction in both testis weight and sperm count (Figure 1B and C), suggesting that MEIOB is a dosage-sensitive regulator of spermatogenesis. The abundance of MEIOB protein was reduced by >70% in the *Meiob*^{N64I/N64I} testes and by 90% in *Meiob*^{N64I/-} testes (Figure 1D). Accordingly, SPATA22, the obligate MEIOB-binding partner, was also dramatically reduced in abundance, whereas the abundance of RPA2, which interacts with MEIOB, remained relatively constant (Figure 1D) (17,19). These data suggest that the N64I substitution causes destabilization of the MEIOB protein and subsequent destabilization of SPATA22 *in vivo*.

Meiob^{N64I} mutation causes apoptosis in metaphase I spermatocytes in mice

Meiotic cell divisions occur at the stage XII of seminiferous tubules, which contain metaphase spermatocytes. Histological analysis revealed a sharply increased number of apparently apoptotic spermatocytes only in stage XII tubules in the *Meiob*^{N64I/N64I} testes at postnatal day 20 (P20; juvenile) and P60 (adult), in contrast with the absence or rare appearance of apoptotic cells in the *Meiob*^{N64I/+} testes (Figure 1E). TUNEL analysis and quantification confirmed this observation (Figure 1F). In addition, the number of apoptotic spermatocytes was higher in *Meiob*^{N64I/-} testes than in *Meiob*^{N64I/N64I} testes (Figure 1E–G), showing that the level of apoptosis increases with reduced MEIOB protein abundance.

MEIOB regulates the number of crossovers

As previously reported, inactivation of *Meiob* leads to an early failure in meiotic recombination, prior to the formation of crossovers (17,18). Crossovers are formed in mid to late pachytene spermatocytes. Whether MEIOB plays a second role in meiotic recombination such as crossover formation cannot be studied, because the *Meiob*^{-/-} mice lack mid to late pachytene spermatocytes, due to meiotic arrest (17,18). The increased apoptosis of metaphase I spermatocytes in *Meiob*^{N64I/N64I} and *Meiob*^{N64I/-} testes indicates a chromosome segregation defect. HEI10, an ubiquitin E3 ligase, is required for crossover formation and localizes as foci on meiotic chromosomes in pachytene spermatocytes (Figure 2A) (34,37). We found that the number of HEI10 foci was reduced in mid to late pachytene spermatocytes from *Meiob*^{N64I/N64I} testes in comparison to *Meiob*^{N64I/+} testes (Figure 2A). Each chromosome from *Meiob*^{N64I/+} spermatocytes had at least one HEI10 focus, in contrast, some chromosomes from *Meiob*^{N64I/N64I} testes lacked HEI10 foci (Figure 2A). Likewise, the number of MLH1 foci was reduced in *Meiob*^{N64I/N64I} spermatocytes than *Meiob*^{N64I/+} spermatocytes (Figure 2B). In addition, *Meiob*^{N64I/-} spermatocytes exhibited a further reduction in the number of HEI10 and MLH1 foci than *Meiob*^{N64I/N64I} spermatocytes (Figure 2A and B).

We next examined metaphase I spermatocytes. While normal metaphase I spermatocytes with all 20 bivalents were observed in the mutant testes, the number of univalent chromosomes was significantly higher in *Meiob*^{N64I/N64I} testes than *Meiob*^{N64I/+} testes (Figure 2C). Univalent chromosomes were even more prevalent in spermatocytes from *Meiob*^{N64I/-} testes (Figure 2C). Both autosomes and sex chromosomes were affected by univalence. The presence of univalent chromosomes in *Meiob*^{N64I/N64I} and *Meiob*^{N64I/-} testes was consistent with the reduced number of MLH1 foci (the sites of crossovers) in the mutant testes. Taken together, the presence of univalent chromosomes leads to chromosome mis-segregation, activates the spindle assembly checkpoint, and thus causes increased apoptosis of metaphase I spermatocytes in *Meiob*^{N64I/N64I} and *Meiob*^{N64I/-} testes (Figure 1E–G). These results demonstrate that MEIOB plays a second and late role in meiotic recombination - crossover formation.

Serine 67 is critical for MEIOB function in meiosis

In the process of generating the N64I allele using the CRISPR/Cas9 approach, the serine 67 deletion allele (*Meiob*^{ΔS67}) was also produced. From the CRISPR injection, we obtained *Meiob*^{N64I/ΔS67} and *Meiob*^{ΔS67/ΔS67} founders, both of which were sterile (Supplementary Figure S2). The adult *Meiob*^{ΔS67/ΔS67} testes showed complete meiotic arrest at the pachytene-like stage (Supplementary Figure S2A) and the adult *Meiob*^{ΔS67/ΔS67} ovaries lacked oocytes (Supplementary Figure S2B). The adult *Meiob*^{N64I/ΔS67} testes exhibited metaphase I arrest (Supplementary Figure S2A), a phenotype different from the *Meiob* null or *Meiob*^{ΔS67/ΔS67} phenotypes. Therefore, the serine 67 residue may provide important insights into the function of MEIOB. However, the *Meiob*^{ΔS67} allele could not be transmitted through the germline from the founders due to sterility and *Meiob*^{ΔS67/+} founder mice (heterozygous) were never produced from the CRISPR injections. To validate the finding on the critical role of serine 67, we generated *Meiob*^{ΔS67} allele through homologous recombination in embryonic stem (ES) cells (Supplementary Figure S2C and S2D).

While *Meiob*^{ΔS67/+} males and females were fertile, both *Meiob*^{ΔS67/ΔS67} males and females were sterile (Figure 3). Histological analysis showed meiotic arrest at the pachytene-like stage in the testis and lack of spermatozoa in the epididymis from *Meiob*^{ΔS67/ΔS67} males (Figure 3A). The adult *Meiob*^{ΔS67/ΔS67} ovary exhibited a complete loss of oocytes (Figure 3B). We further analyzed the spermatocytes from the adult mutant testes by assessing chromosomal synapsis with immunostaining of synaptonemal complex (SC) proteins SYCP1 and SYCP3 (Figure 3C–H). The percentage of zygotene-like spermatocytes was significantly higher in the *Meiob*^{ΔS67/ΔS67} testes than the *Meiob*^{ΔS67/+} testes. Pachytene spermatocytes with normal synapsis were abundant in *Meiob*^{ΔS67/+} testes (Figure 3G and H) but absent in *Meiob*^{ΔS67/ΔS67} testes. Instead, pachytene-like spermatocytes with well-formed SC lateral elements but lack of synapsis were observed in *Meiob*^{ΔS67/ΔS67} testes (Figure 3F and H). γ H2AX-positive XY body was present in wild type pachytene spermatocytes but absent in *Meiob*^{ΔS67/ΔS67}

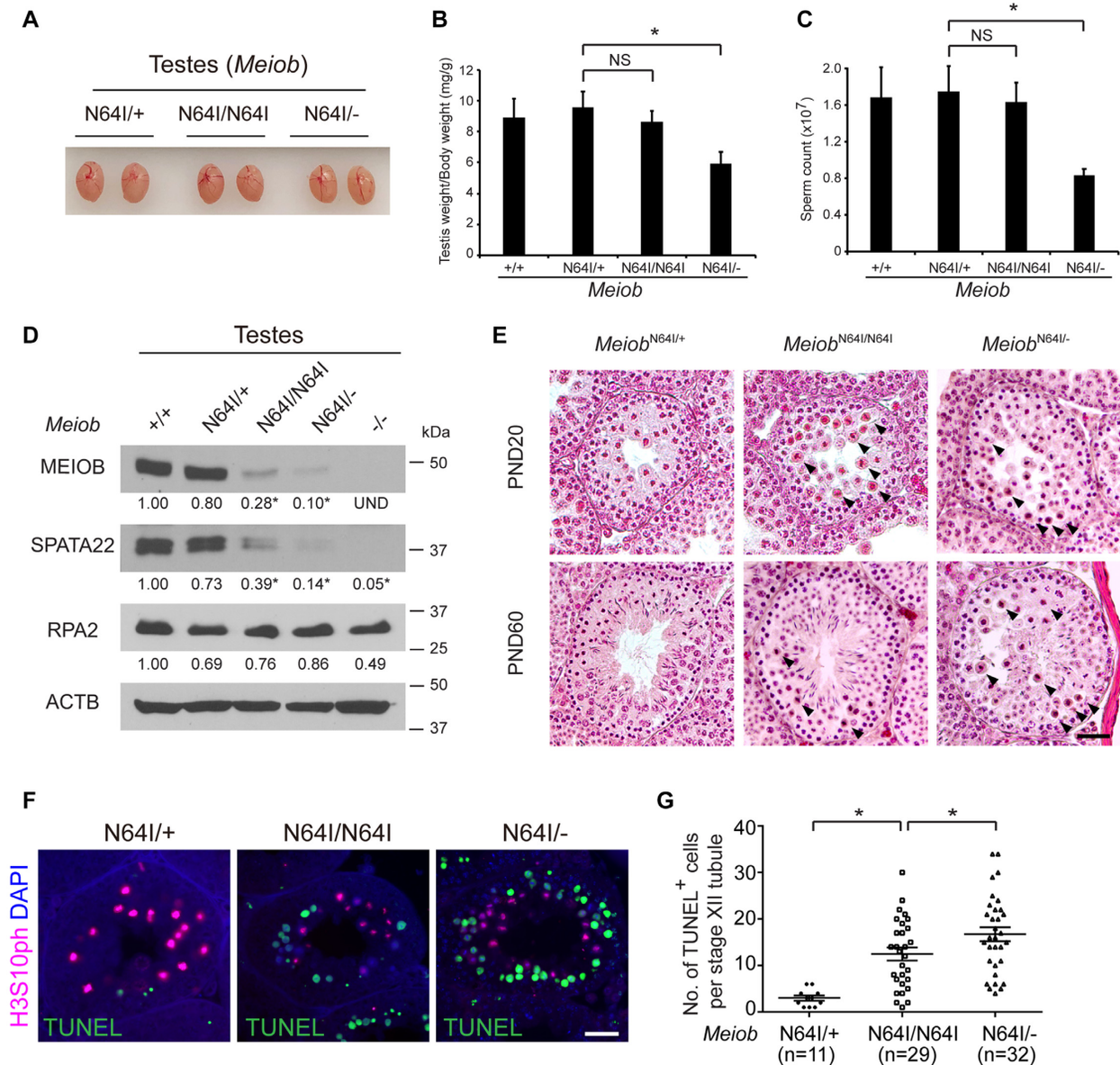


Figure 1. The *MeioB* N64I mutation causes stage XII-specific meiotic defects in males. (A) Testis size of 8-week-old *MeioB*^{N64I/+}, *MeioB*^{N64I/N64I} and *MeioB*^{N64I/-} mice. (B, C) Reduced testis weight (B) and sperm count (C) in *MeioB*^{N64I/-} males. $n = 3$, 8-week-old males. Data are presented as mean \pm S.D., * $P < 0.05$ by Student's *t*-test. NS, non-significant. (D) The N64I substitution destabilizes the MEIOB protein in testes (postnatal day 20, P20). Band quantification was normalized to β -actin (ACTB). Western blotting was performed in triplicates. * $P < 0.05$. (E) Histological analysis of testes at P20 and P60. Stage XII tubules are shown. Apoptotic metaphase spermatocytes are marked by black arrowheads. Scale bar, 20 μ m. (F) TUNEL analysis of testicular sections at P20. H3S10ph (Phosphorylated H3 at serine 10) is a metaphase marker. Scale bar, 20 μ m. (G) Quantification of TUNEL-positive spermatocytes at P20. n , the number of TUNEL-positive stage XII tubules (TUNEL-negative stage XII tubules were excluded). Three mice per genotype were examined. * $P < 0.05$ by Student's *t*-test.

pachytene-like spermatocytes (Figure 3I) (49). These results demonstrate a failure in chromosome synapsis in the *MeioB* ^{Δ S67/ Δ S67} mice.

MEIOB and SPATA22 interact with each other and form foci on SC in meiotic germ cells (Figure 3J) (17). In addition, the stability of these two proteins depends on each other in both cultured cells and testis (19,50). Interestingly, MEIOB foci were not detected in *MeioB* ^{Δ S67/ Δ S67} spermatocytes (Figure 3J). As expected, the SPATA22 foci were also

absent in *MeioB* ^{Δ S67/ Δ S67} spermatocytes (Figure 3J). These *MeioB* ^{Δ S67/ Δ S67} phenotypes were similar to the *MeioB*^{-/-} phenotypes (17), showing that serine 67 is required for the MEIOB function.

Impaired spermatogenesis in *MeioB*^{N64I/ Δ S67} males

To test the dosage effect of these two *MeioB* mutant alleles, we generated *MeioB*^{N64I/ Δ S67} males through breeding of

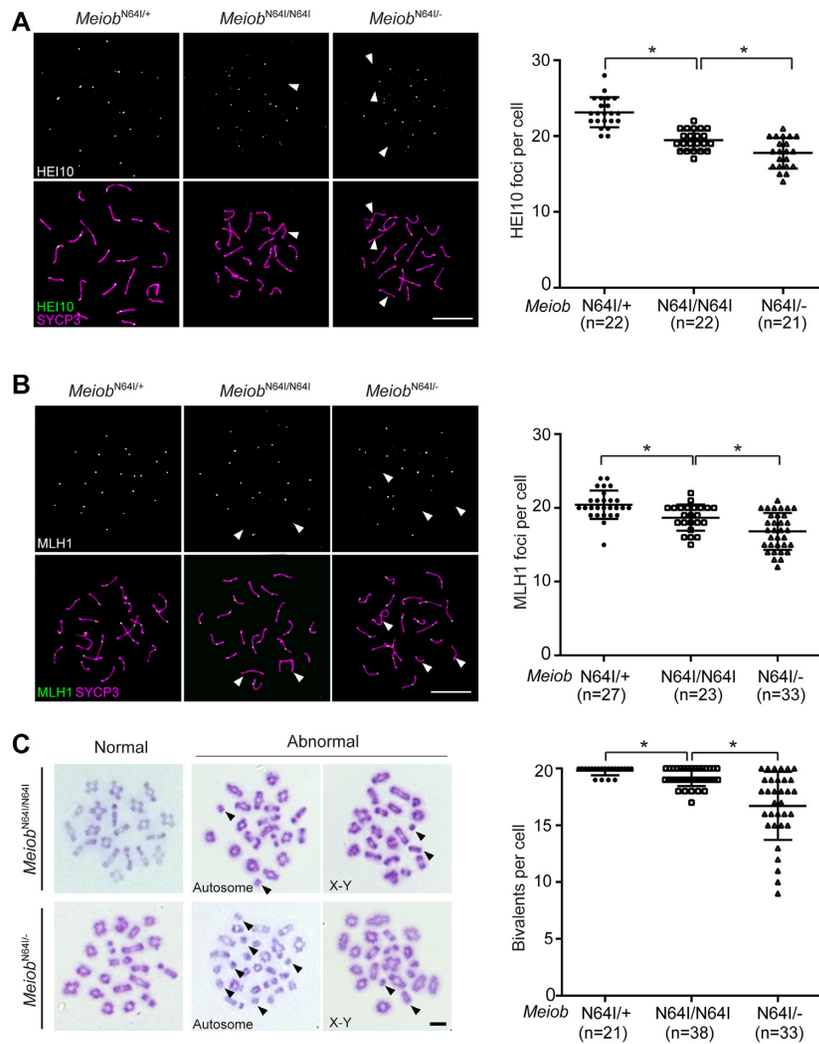


Figure 2. The *MeioB* N64I mutation reduces crossover formation. (A, B) Analysis of HEI10 foci (A), and MLH1 foci (B) in pachytene spermatocytes from wild type, *MeioB*^{N64I/N64I}, and *MeioB*^{N64I/-} mice at P20. *n*, number of mid to late pachytene spermatocytes counted. (C) Analysis of chiasmata in metaphase I spermatocytes from *MeioB*^{N64I/N64I} and *MeioB*^{N64I/-} testes at P20. Univalents are indicated by arrowheads. *n*, number of metaphase I spermatocytes counted. The data are from three to five mice per genotype. **P* < 0.05 by Student's *t*-test. Scale bars, 10 μ m.

MeioB ^{Δ S67/+} and *MeioB*^{N64I/N64I} mice in anticipation that the MEIOB protein levels in *MeioB*^{N64I/ Δ S67} testis may lie between those in *MeioB*^{N64I/N64I} and *MeioB* ^{Δ S67/ Δ S67} testes. The *MeioB*^{N64I/ Δ S67} testis was smaller than *MeioB*^{N64I/+} testis but larger than *MeioB* ^{Δ S67/ Δ S67} testis (Figure 4A). Consistently, the testis weight of adult *MeioB*^{N64I/ Δ S67} males was significantly lower than that of *MeioB*^{N64I/+} males but higher than that of *MeioB* ^{Δ S67/ Δ S67} males (Figure 4B). The sperm count in *MeioB*^{N64I/ Δ S67} males was much lower than *MeioB*^{N64I/+} males, whereas *MeioB* ^{Δ S67/ Δ S67} males lacked sperm (Figure 4C). We performed mating tests for both males and females (Supplementary Table S2). While all four *MeioB*^{N64I/+} males were fertile, two of five *MeioB*^{N64I/ Δ S67} males did not sire any offspring but the remaining three *MeioB*^{N64I/ Δ S67} males produced fewer litters but with normal litter size (Supplementary Table S2A). The *MeioB*^{N64I/ Δ S67} females produced normal litter size but

MeioB ^{Δ S67/ Δ S67} females were sterile (Supplementary Table S2B).

In spermatogenesis, the pachytene checkpoint functions at stage IV tubules and the spindle checkpoint operates at stage XII tubules. Histological analysis revealed increased apoptosis of spermatocytes in stage XII but not stage IV seminiferous tubules from *MeioB*^{N64I/ Δ S67} males at P24, 28 and 35 (Figure 4D). The increased apoptosis at stage XII was confirmed by TUNEL (Figure 4E and F). However, *MeioB*^{N64I/ Δ S67} males (generated through ES cells) still produced sperm and thus their meiotic defects were similar to but less severe than those observed in *MeioB*^{N64I/ Δ S67} founder males (generated through CRISPR injection), which showed complete metaphase I arrest in stage XII (Supplementary Figure S2A). It was possible that CRISPR founders harbored unknown off-target mutations or had a different (DBA \times C57BL/6) hybrid genetic back-

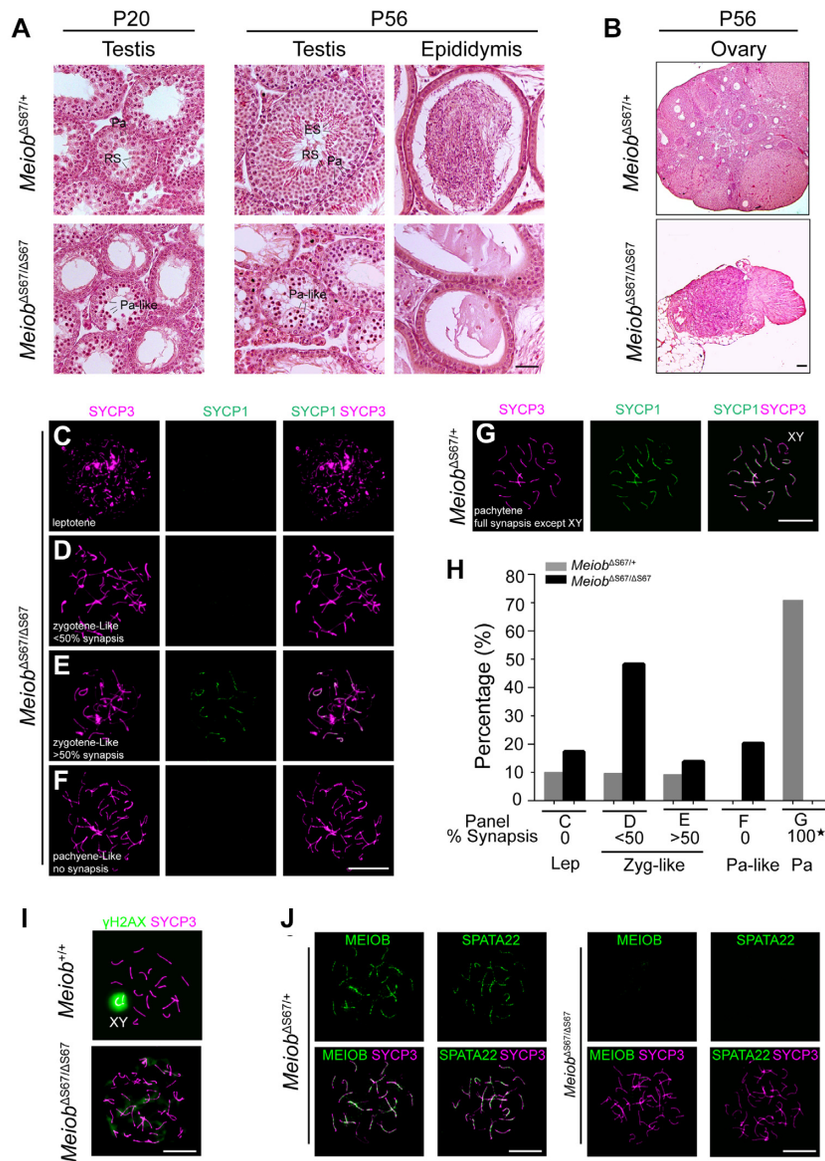


Figure 3. The S67 residue of MEIOB is essential for meiosis and fertility in mice of both sexes. (A) Histological analysis of testis and epididymis from P20 and 8-week-old mice. Zyg-like, zygotene-like spermatocytes; Pa, pachytene spermatocytes; RS, round spermatids; ES, elongated spermatids. Scale bar, 50 μ m. (B) Histological analysis of ovaries from 8-week-old mice. Scale bar, 100 μ m. (C–H) Spread nuclei of spermatocytes from 8-week-old males were immunostained with anti-SYCP1 and anti-SYCP3 antibodies. 247 *MeioB^{ΔS67/+}* and 201 *MeioB^{ΔS67/ΔS67}* spermatocytes were counted and categorized into different groups (three mice per genotype): leptotene (C), zygotene-like with <50% synapsis (D), zygotene-like with >50% synapsis (E), pachytene-like with no synapsis (F) and *MeioB^{ΔS67/+}* pachytene spermatocytes (G). (H) Percentage of spermatocytes at each stage of meiotic prophase I with representative images shown in panels C–G. The 100% of synapsis (asterisk) in *MeioB^{ΔS67/+}* pachytene spermatocytes refer to the full synapsis of all autosomes excluding XY chromosomes. (I) γ H2AX localization in spermatocytes from 8-week-old males. Note the γ H2AX-positive XY body in the wild type pachytene spermatocyte and cloud-like γ H2AX signals in pachytene-like *MeioB^{ΔS67/ΔS67}* spermatocyte. (J) MEIOB and SPATA22 foci are present in *MeioB^{ΔS67/+}* late zygotene spermatocytes but absent in *MeioB^{ΔS67/ΔS67}* pachytene-like spermatocytes (I). Scale bars (C–J), 10 μ m.

ground. Interestingly, increased apoptosis of spermatocytes was also observed in stage IV tubules from *MeioB^{N641/ΔS67}* males at a younger age (P20), which did not have stage XII tubules possibly due to a delay in spermatogenesis (Figure 4D). In contrast, spermatogenesis progressed to stage XII and there was increased apoptosis in stage XII but not stage IV tubules in *MeioB^{N641/-}* testes at P20 (Figure 1E), suggesting that the *MeioB^{N641/ΔS67}* testes exhibited slightly more severe meiotic defects than *MeioB^{N641/-}* testes.

To investigate the cause of increased apoptosis in stage IV tubules in P20 *MeioB^{N641/ΔS67}* testis, we analyzed chro-

mosome synapsis. P20 *MeioB^{N641/+}* testis contained spermatocytes from leptotene to metaphase I, however, P20 *MeioB^{N641/ΔS67}* testis lacked diplotene and metaphase I spermatocytes, suggesting a delay in spermatogenesis (Supplementary Figure S3A). In P20 *MeioB^{N641/ΔS67}* testis, two types of abnormal spermatocytes were observed: zygotene-like (no synapsis) and pachytene-like (condensed SC with <50% synapsis) (Supplementary Figure S3A). The pachytene-like spermatocytes were γ H2AX-positive throughout the chromatin and lacked the typical XY body (Supplementary Figure S3B). Consistent with the histolog-

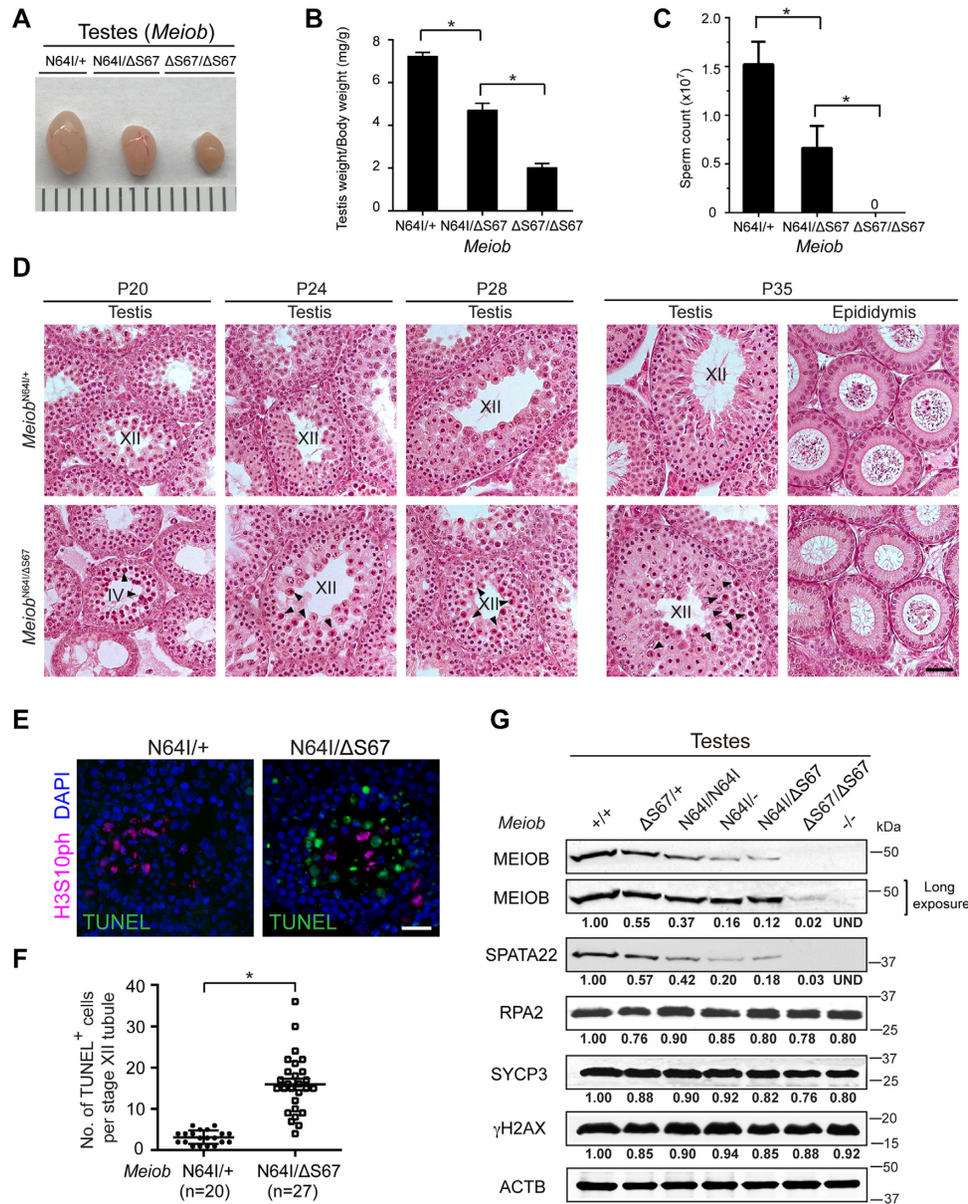


Figure 4. *MeioB*^{N641/ Δ S67} males exhibit age-dependent meiotic defects in juvenile spermatogenesis. (A) Testis size of adult (8-week-old) males. (B, C) Testis weight (B) and sperm count (C) of adult (8-week-old) males. n, number of males. Data are shown as mean \pm S.D., **P* < 0.05 by Student's *t*-test. (D) Histological analysis of testis and epididymis at P20, P24, P28 and P35. Stage IV and stage XII tubules are labeled. Apoptotic spermatocytes are marked by arrowheads. Scale bar, 50 μ m. (E, F) TUNEL analysis of *MeioB*^{N641/+} and *MeioB*^{N641/ Δ S67} testes at P24 (E) and quantification of TUNEL-positive spermatocytes (F). n, the number of TUNEL-positive stage XII tubules (TUNEL-negative stage XII tubules were excluded). Three mice per genotype were examined. **P* < 0.05 by Student's *t*-test. Scale bar, 20 μ m. (G) Western blot analysis of MEIOB, SPATA22, and RPA2 in P20 testes. The levels of SYCP3 and γ H2AX are constant across samples. Relative band intensity (normalized to ACTB) is shown.

ical result (Figure 4D), massive apoptosis was observed in stage IV tubules from P20 *MeioB*^{N641/ Δ S67} testis, presumably to eliminate the pachytene-like spermatocytes (Supplementary Figure S3C). However, zygotene-like and pachytene-like spermatocytes were absent in P24 *MeioB*^{N641/ Δ S67} testis (Supplementary Figure S3D). One possible explanation is that meiosis in the first wave of spermatogenesis at P20 is more sensitive to perturbation than the subsequent waves (P24 and beyond).

To probe the nature of gene dosage effect, Western blot analysis was performed on testes of various combinations

of *MeioB* alleles (Figure 4G). The MEIOB protein was nearly absent but detectable in *MeioB* ^{Δ S67/ Δ S67} testes, providing an explanation for the complete meiotic arrest phenotype. As expected, the MEIOB protein abundance was reduced by nearly 50% in *MeioB* ^{Δ S67/+} testes compared to wild type. *MeioB*^{N641/ Δ S67} testes exhibited significantly lower MEIOB protein abundance than *MeioB*^{N641/N641} testes and slightly lower than *MeioB*^{N641/-} testes. The SPATA22 abundance correlated with the MEIOB abundance (Figure 4G). Consistent with their low protein abundance, MEIOB and SPATA22 foci on meiotic chromosomes were much weaker

in *Meiob*^{N64I/ΔS67} spermatocytes (Supplementary Figure S4A). In contrast, RPA2, another MEIOB-interacting protein, did not change in protein abundance in these *Meiob* mutant testes (Figure 4G). The *Meiob* and *Spata22* transcript levels were measured by real-time RT-PCR (Supplementary Figure S4B). The slight decrease in the transcript abundance could not account for the near absence of MEIOB and SPATA22 proteins in *Meiob*^{ΔS67/ΔS67} testes. Therefore, these results demonstrate that the S67 deletion severely affects the MEIOB protein stability *in vivo*.

Reduced crossover formation in *Meiob*^{N64I/ΔS67} testes

We examined meiotic recombination in P24 *Meiob*^{N64I/ΔS67} spermatocytes. The number of RPA2 foci was comparable between *Meiob*^{N64I/ΔS67} and *Meiob*^{N64I/+} spermatocytes (Supplementary Figure S5A). The number of γH2AX foci was also comparable between *Meiob*^{N64I/ΔS67} and *Meiob*^{N64I/+} spermatocytes (Supplementary Figure S5B). The increased apoptosis of metaphase spermatocytes in stage XII tubules from *Meiob*^{N64I/ΔS67} testes suggested a defect in chromosome segregation. Therefore, we assessed crossover formation in spermatocytes from *Meiob*^{N64I/ΔS67} testes. TEX11, a component of intermediate meiotic recombination nodules, forms foci on meiotic chromosomes and regulates crossover formation (26). The number of TEX11 foci was reduced in both early- and mid-pachytene spermatocytes from *Meiob*^{N64I/ΔS67} testes (Figure 5A). The number of HEI10 and MLH1 foci was also significantly reduced in mid-late pachytene spermatocytes from *Meiob*^{N64I/ΔS67} testes (Figure 5B and C). Metaphase spread analysis revealed a significant increase in the number of univalent chromosomes in *Meiob*^{N64I/ΔS67} testes and univalence affected both autosomes and sex chromosomes (Figure 5D).

At the diplotene stage, homologous chromosomes are connected by chiasmata (the sites of crossover) and remain as bivalents (Supplementary Figure S6). However, univalent chromosomes were present in diplotene spermatocytes from *Meiob*^{N64I/N64I}, *Meiob*^{N64I/-}, and *Meiob*^{N64I/ΔS67} testes at an increasing frequency (Supplementary Figure S6). These data demonstrate that both *Meiob*^{N64I} and *Meiob*^{ΔS67} mutations reduce the formation of crossovers in spermatocytes.

Females are usually more tolerant of meiotic errors than males (51). We examined meiosis in oocytes from *Meiob*^{N64I/+} and *Meiob*^{N64I/ΔS67} ovaries at E18.5, when pachytene oocytes are most abundant (Supplementary Figure S7). Meiotic progression was slightly delayed in *Meiob*^{N64I/ΔS67} ovaries, indicated by the lower percentage of pachynema and the concurrently increased percentage of leptoneuma and zygonema (Supplementary Figure S7A). However, the numbers of RPA2 foci and MLH1 foci, and γH2AX disappearance were comparable between *Meiob*^{N64I/+} and *Meiob*^{N64I/ΔS67} ovaries at E18.5, showing normal crossover formation in females (Supplementary Figure S7B–D).

Reduced crossover formation in spermatocytes was also reported in other meiotic mutants such as *Rpa1*, *Tex11* and *Rnf212* (7,26,33). MEIOB colocalizes and interacts with RPA1 (17,19). MEIOB colocalizes with MSH4 in a subset of foci (17). Immunofluorescent analysis showed that the majority of MEIOB foci (83%) overlapped with TEX11 foci

in pachytene spermatocytes (Supplementary Figure S8A). We performed co-transfection/co-immunoprecipitation experiments in HEK293T cells. MEIOB was not associated with any of the proteins tested (TEX11, MSH4, RNF212 and MLH1) (Supplementary Figure S8). Although it is possible that there are unknown protein interactions involving MEIOB, the MEIOB-RPA1 interaction might be the main contributor to crossover regulation.

N64I and ΔS67 mutations destabilize the MEIOB protein

The MEIOB protein abundance was dramatically reduced in *Meiob*^{N64I/N64I} testes and further reduced in *Meiob*^{N64I/ΔS67} testes (Figure 4G), suggesting that the mutant MEIOB proteins are unstable. To further test the effect of these mutations on protein stability, we constructed expression plasmids with various mutations for expression in HEK293T cells (Figure 6A). In transfected cells, the MEIOB^{N64I} protein was less abundant than the wild type, and the MEIOB^{ΔS67} protein abundance was even lower (Figure 6B). Therefore, these two mutations reduced the MEIOB protein stability both in testes and in transfected somatic cells. It is puzzling that the N64I mutation causes meiotic arrest in the infertile man but not in mice (44). The difference in mouse and human phenotypes could be attributed to the context of the mutation, because the mouse and human MEIOB proteins exhibit only 85% identity and noticeably have two different residues adjacent to N64 (Supplementary Figure S1A). To mimic the human protein sequence, we made the partially humanized IAA mouse mutant (Figure 6A). Indeed, the mouse MEIOB^{IAA} protein was much less stable than the MEIOB^{N64I} protein, and was even less stable than the MEIOB^{ΔS67} protein (Figure 6B). In the presence of SPATA22, the abundance of these MEIOB mutant proteins was still significantly reduced (Figure 6B). Human MEIOB increased human SPATA22 abundance when co-expressed (Figure 6C). The human MEIOB^{N64I} protein was less stable than wild type in the absence or presence of hSPATA22 (Figure 6D). In conclusion, the severity of these mutations on MEIOB protein stability provides an explanation for the meiotic arrest phenotypes in *Meiob*^{ΔS67/ΔS67} male mice and *MEIOB*^{N64I/N64I} man.

N64I and ΔS67 mutations cause MEIOB self-association

To investigate the destabilizing effects of N64I and ΔS67 mutations, we tested protein aggregation. The wild type MEIOB protein did not dimerize in the absence of SPATA22 (Figure 6E) but were associated with each other in the presence of SPATA22 (Figure 6F). However, in the absence of SPATA22, MEIOB^{N64I} protein alone formed a complex (Figure 6G). MEIOB^{ΔS67} alone also formed a complex (Figure 6H). In addition, MEIOB^{N64I} and MEIOB^{ΔS67} interacted with each other (Figure 6I) and each mutant protein formed a complex with the wild type MEIOB (Figure 6J and K). We performed reciprocal immunoprecipitations using GFP antibody and obtained the same results (Supplementary Figure S9).

We measured the half-life of wild type mouse MEIOB, mouse MEIOB^{N64I}, and mouse MEIOB^{ΔS67} proteins in transfected HEK293T cells (Figure 7A). The wild type

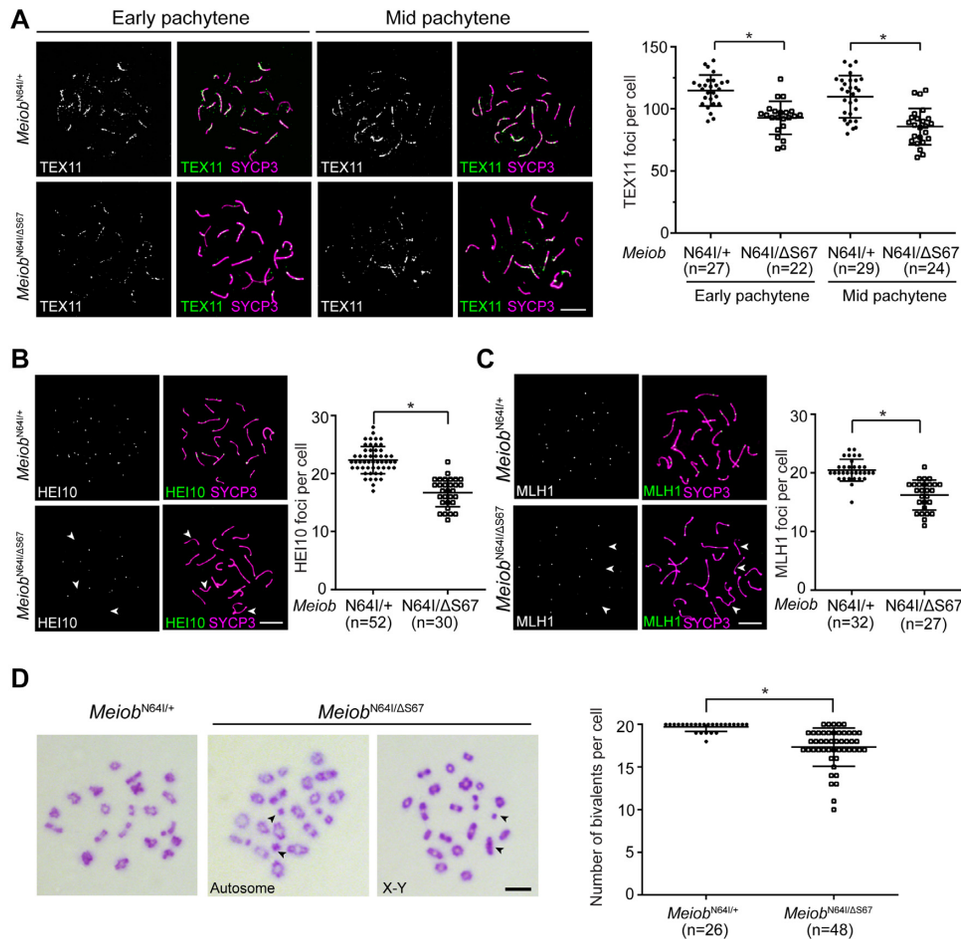


Figure 5. MEIOB dosage-dependent regulation of crossover formation. (A–C) Analysis of TEX11, HEI10, and MLH1 foci in pachytene spermatocytes at P24. n, number of pachytene spermatocytes counted. HEI10 and MLH1 foci were counted in mid-late pachytene spermatocytes. (D) Spread analysis of metaphase I spermatocytes at P24. Univalents are indicated by arrowheads. n, number of metaphase I spermatocytes counted. Data are from three to five mice per genotype. * $P < 0.05$ by Student's *t*-test. Scale bars, 10 μ m.

MEIOB protein was very stable with a half-life of 19.9 h (Figure 7A). However, the MEIOB^{N64I} protein had a significantly shortened half-life (9.4 h). The S67 deletion further reduced the MEIOB half-life to 7.5 h, showing that S67 deletion is more deleterious to the MEIOB protein than the N64I substitution. We also measured the half-life of wild type human MEIOB, human MEIOB^{N64I}, and mouse MEIOB^{IAA} proteins in transfected HEK293T cells (Figure 7B). The half-life of human wild-type MEIOB (19.1 h) was comparable to that of mouse MEIOB. The human MEIOB^{N64I} had a shortened half-life (9.1 h), which was comparable to the half-life of mouse MEIOB^{N64I}. In addition, the mouse MEIOB^{IAA} protein had the shortest half-life (3.7 h). Taken together, these data demonstrate that N64I and Δ S67 mutations cause abnormal self-aggregation of MEIOB, which could be targeted for degradation.

We sought to investigate the mechanism of accelerated degradation of mutant MEIOB proteins in HEK293T cells by inhibiting the proteasome or the autophagy protein degradation pathways. Treatment with MG132, a proteasome inhibitor, caused the accumulation of c-Jun, an endogenous proteasome target (52), but did not extend the

half-life of wild type MEIOB, MEIOB^{N64I} and MEIOB ^{Δ S67} proteins (Supplementary Figure S10A). Likewise, treatment with bafilomycin A1, an autophagy inhibitor, increased abundance of LC3A/B-II, an autophagosome marker (53), but lacked any effect on the turnover of wild type MEIOB, MEIOB^{N64I}, and MEIOB ^{Δ S67} proteins (Supplementary Figure S10B). These results indicate that instability of MEIOB^{N64I} and MEIOB ^{Δ S67} proteins is dependent on proteases, which are likely not associated with proteasome or lysosome activities. While many examples of such proteases (including calpain and caspases) are known, the exact nature of enzymes facilitating rapid degradation of MEIOB mutants is yet to be determined.

DISCUSSION

Our results demonstrate that MEIOB is a dosage-sensitive regulator of meiotic recombination (Figure 8). As described in this study and previous studies (17,18), *MeioB*^{-/-} and *MeioB* ^{Δ S67/ Δ S67} spermatocytes lack or nearly completely lose the MEIOB protein, show a complete failure in DSB repair in early meiotic recombination, and as a result,

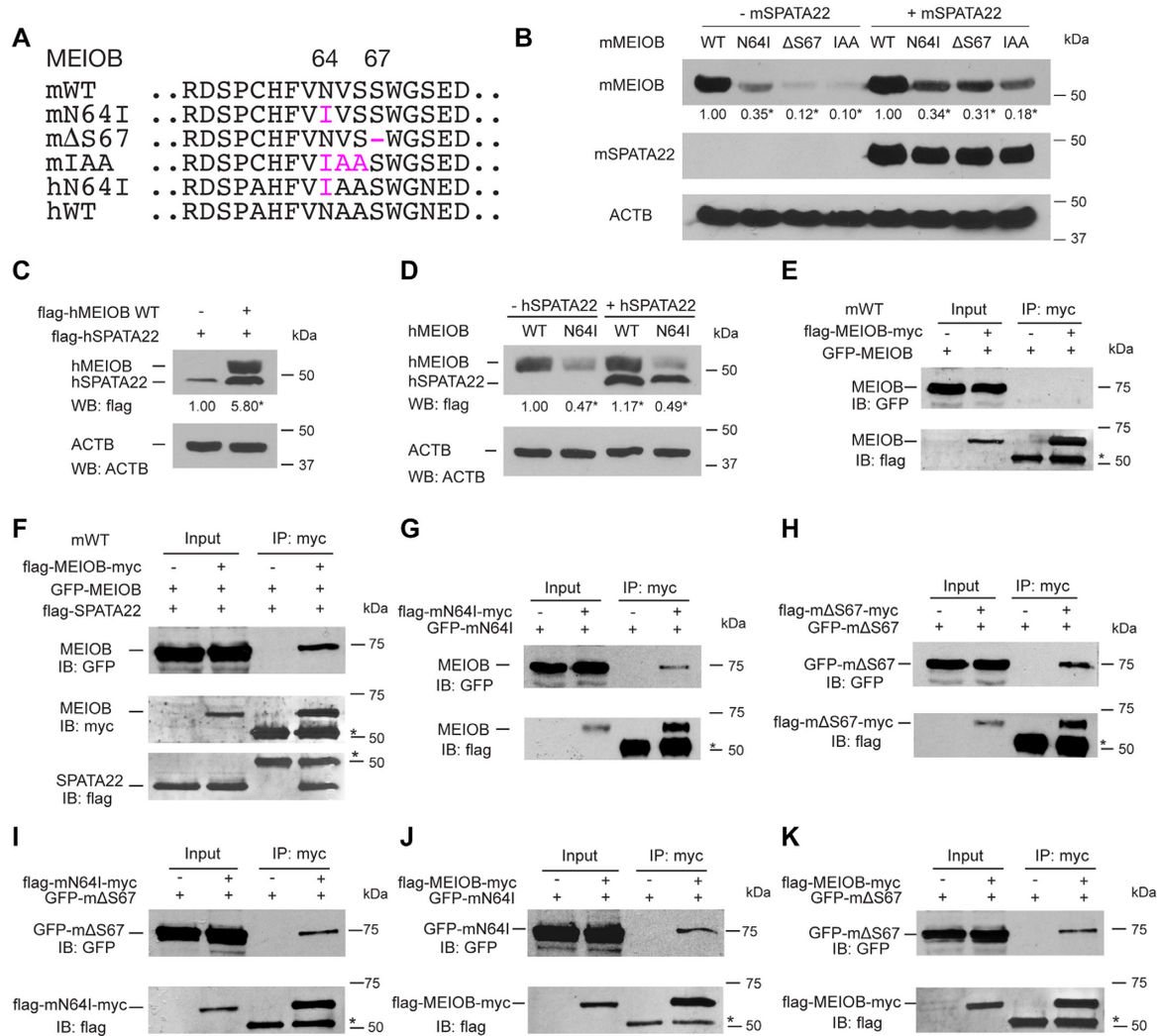


Figure 6. *MeioB* mutations destabilize the MEIOB protein and cause self-association. Full-length mouse and human MEIOB proteins were expressed in HEK293T cells in all transfection experiments. The number below each band is the fold change of protein abundance compared to the wild type. (A) Wild type and mutant MEIOB proteins. m, mouse; h, human. (B) Decreased stability of mutant mouse MEIOB proteins in the presence and absence of mSPATA22. The experiments were performed three times. **P* < 0.05 by Student's *t*-test. (C) Human MEIOB stabilizes hSPATA22. (D) The N64I substitution destabilizes hMEIOB. hMEIOB bands were quantified. The experiments were performed three times. **P* < 0.05 by Student's *t*-test. (E, F) Wild type MEIOB does not self-associate (E) but associate in the presence of SPATA22 (F). (G–K) MEIOB^{N64I} interacts with itself (G), MEIOB^{ΔS67} interacts with itself (H), MEIOB^{N64I} interacts with MEIOB^{ΔS67} (I), MEIOB^{N64I} interacts with wild type MEIOB (J), MEIOB^{ΔS67} interacts with wild type MEIOB (K).

undergo apoptosis at stage IV due to the activation of the pachytene checkpoint (Figure 8A). In contrast, the *MeioB*^{N64I/N64I} spermatocytes retaining 30% of the protein progress through the pachytene stage but exhibit reduced crossovers and thus undergo apoptosis at stage XII due to activation of the spindle assembly checkpoint. The reduction in crossovers is more pronounced in *MeioB*^{N64I/-} and *MeioB*^{N64I/ΔS67} spermatocytes, which have ~10% of the protein. It is striking that even 10% of the MEIOB protein is sufficient to complete early meiotic recombination and most crossovers in late meiotic recombination. We postulate that, at a low MEIOB/SPATA22 abundance, second ends are captured but a subset of captured second ends are not stable enough to form double Holliday junctions, resulting in a lower number of crossovers (Figure 8B). Notably,

meiosis-specific proteins RNF212, HEI10 and TEX11 are also dosage-sensitive regulators of crossovers (33,37,38). Therefore, crossover formation appears to be particularly sensitive to protein abundance.

The *MEIOB*^{N64I} mutation caused male infertility in humans but the biochemical basis of the deleterious effect of this missense mutation was unknown (44). Our previous study identifies D383 in MEIOB as a critical residue for interaction with SPATA22. The D383A mutation abolishes the MEIOB-SPATA22 interaction and causes sterility in mice (19,50). Asparagine 64 and serine 67 reside in one of the three OBDs (oligonucleotide-binding domain) but outside of the SPATA22-binding domain (19,44). We find that the N64I missense mutation and deletion of S67 result in self-aggregation of MEIOB and sharply re-

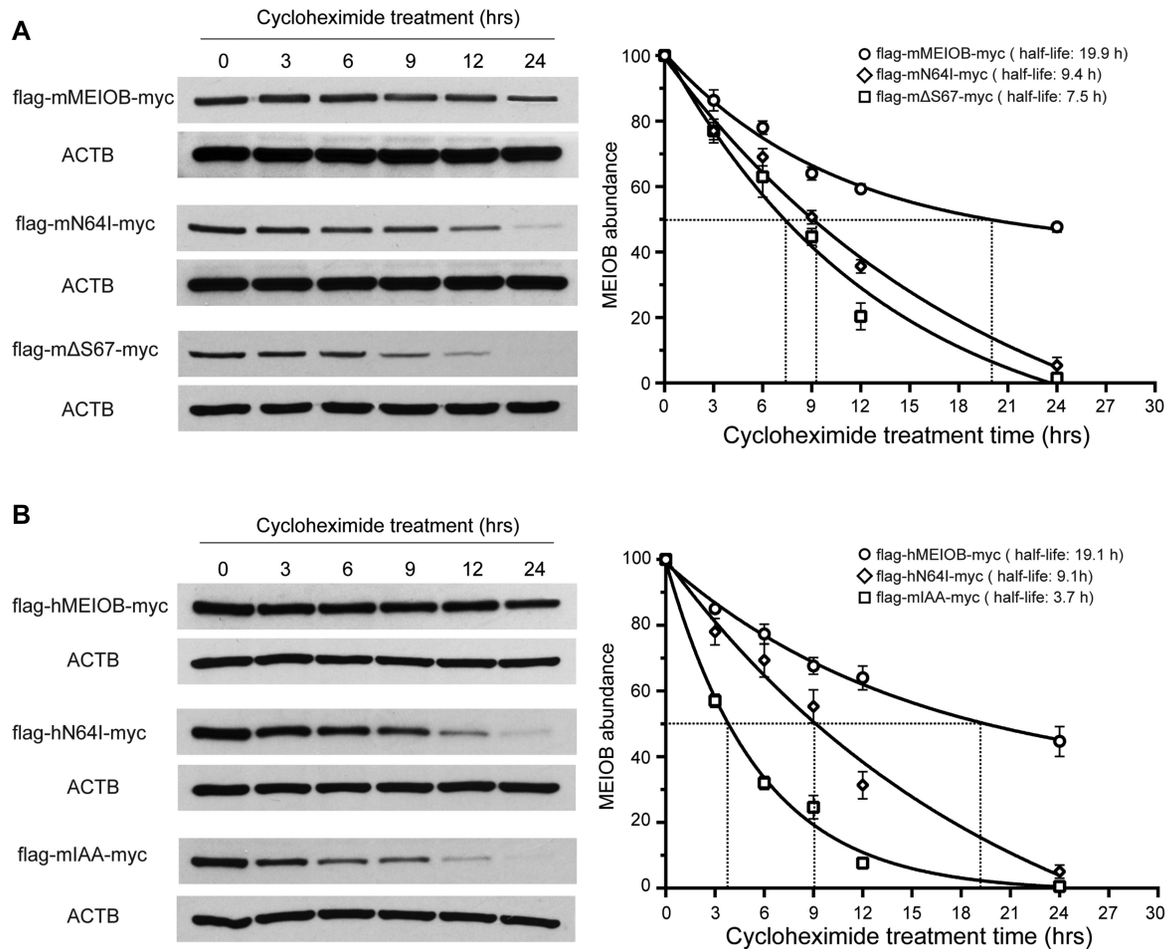


Figure 7. Reduction in the half-life of MEIOB mutant proteins. (A) Western blot analysis and half-life of mouse MEIOB (wt), MEIOB^{N64I} and MEIOB^{ΔS67} in HEK293T cells treated with cycloheximide. The plot on the right shows the half-life. (B) Western blot analysis of human MEIOB (wt), human MEIOB^{N64I} and mouse MEIOB^{ΔIAA} (as in Figure 6A) in HEK293T cells treated with cycloheximide. The plot on the right shows the half-life. ACTB serves as a loading control. hrs, hours after addition of cycloheximide. Experiments were done three times. Curve fitting was performed with one-phase exponential decay.

duced protein half-life. We also generated *Meiob*^{R235A} mice through CRISPR/Cas9-mediated genome editing. Arginine 235 (R235) in MEIOB resides in one of its OBDs and is critical for its ssDNA-binding activity *in vitro* (17). However, the *Meiob*^{R235A/R235A} mice had normal fertility, spermatogenesis, and sperm count (data not shown), showing that R235 is dispensable. Therefore, S67 deletion is not likely to affect meiosis through its effect on ssDNA-binding. Given the dramatic reduction in MEIOB^{ΔS67} protein abundance, the reduced protein stability is most likely to be the main cause of the meiotic arrest phenotype in *Meiob*^{ΔS67/ΔS67} testis. A large number of missense SNPs in humans have been identified from the thousand genome project and other genome sequencing projects. Each human genome contains 10 000–12 000 protein sequence-altering variants (54). However, the functional consequences of these variants are usually difficult to predict. A study of ~2000 missense human variants show that ~10% of them disrupt protein interactions (55). Our study of *Meiob* mutations suggest that some of human missense variants may affect protein stability.

The *Meiob* N64I substitution exerts different effects on fertility in human and mouse. Men with the homozygous MEIOB^{N64I} mutation exhibit meiotic arrest and infertility (44), whereas MEIOB^{N64I/N64I} mice display mild meiotic defects but are fertile. The human and mouse MEIOB proteins share 85% sequence identity. One possible explanation is that the N64I substitution may destabilize the human protein more than the mouse protein. However, this is not the case, because the human and mouse MEIOB^{N64I} proteins have comparable half-lives in cultured cells (Figure 7). It is possible that the MEIOB threshold required for fertility in humans is much higher than in mouse. Intriguingly, both a homozygous mutation in *PTX4* (400 kb from MEIOB) and MEIOB^{N64I/N64I} co-segregated with infertility in the family (44). Therefore, the meiotic arrest phenotype in azoospermic men could be due to the compound effect of the MEIOB^{N64I} mutation and the *PTX4* mutation (44).

Intriguingly, both MEIOB and SPATA22 protein levels were reduced, but their transcript levels were increased in *Meiob*^{N64I/N64I} testes (Figure 4G and Supplementary Figure S4B). Codon usage can affect transcript abundance through

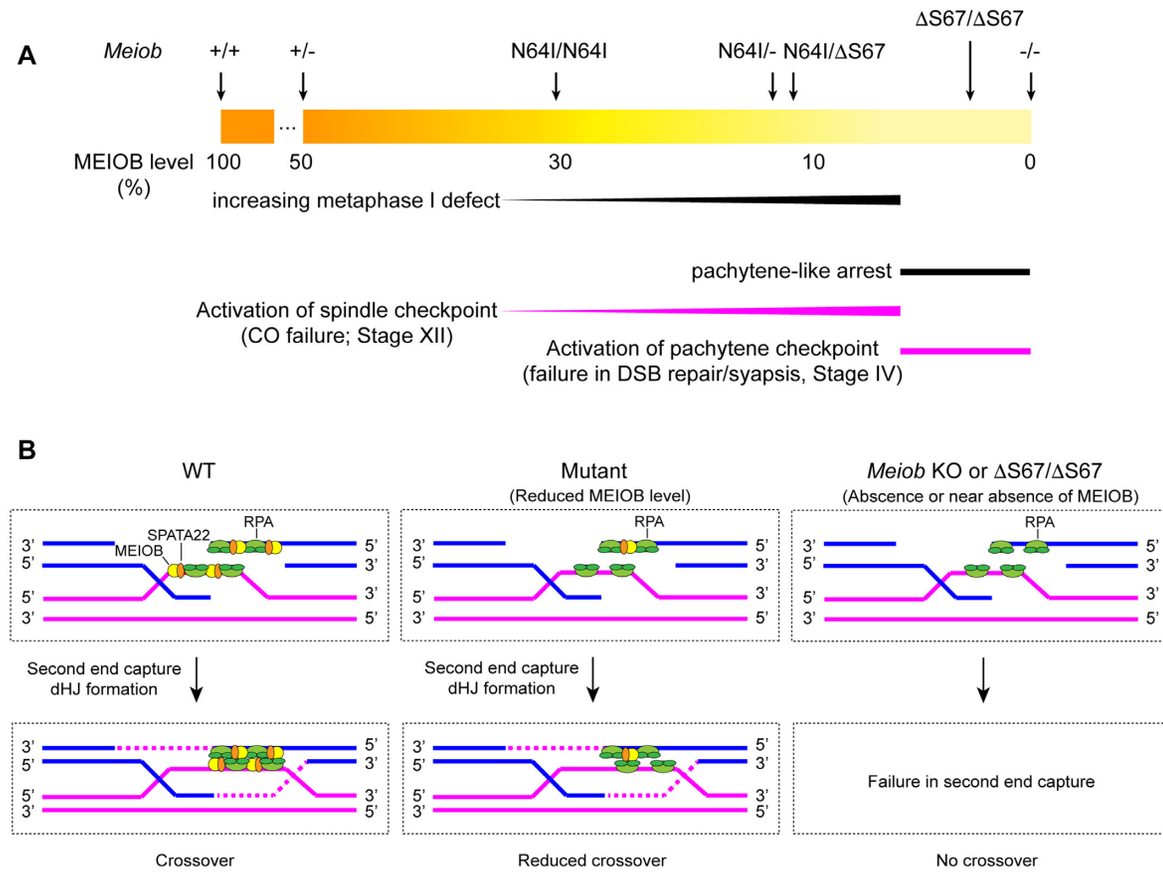


Figure 8. Dosage-dependent and stage-specific regulation of meiotic recombination by MEIOB. (A) The gradient of the MEIOB protein level in testis is depicted (WT, 100%; *Meiob*^{-/-}, 0%). Two stages of meiotic defects are diagrammed: pachytene defect at stage IV and metaphases I defect at stage XII. (B) Working model on the role of MEIOB/SPATA22 in crossover formation. MEIOB/SPATA22 (dimer) and RPA (trimer) interact with each other but bind to ssDNA independently. The interaction between these two complexes facilitates second end capture and enhances dHJ formation. Reduced MEIOB/SPATA22 abundance decreases the stability of captured second ends and thus fewer crossovers. Lack of MEIOB/SPATA22 blocks the formation of dHJs.

its effect on transcription (56). It is formally possible that the N64I codon change (AAT→ATC) causes increased transcription of *Meiob* through local chromatin modification. SPATA22 and MEIOB are binding partners with the same expression profiles. Functionally related mammalian genes could be co-regulated transcriptionally through topographically associated domains (TADs) even if located on different chromosomes (57). It is possible that the *Spata22* transcription might be increased in *Meiob*^{N64I/N64I} testes through high-order chromatin interactions. These possibilities warrant further investigation.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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