Histone methyltransferase PRDM9 is not essential for meiosis in male mice

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A hallmark of meiosis is the rearrangement of parental alleles to ensure genetic diversity in the gametes. These chromosome rearrangements are mediated by the repair of programmed DNA double-strand breaks (DSBs) as genetic crossovers between parental homologs. In mice, humans, and many other mammals, meiotic DSBs occur primarily at hotspots, determined by sequence-specific binding of the PRDM9 protein. Without PRDM9, meiotic DSBs occur near gene promoters and other functional sites. Studies in a limited number of mouse strains showed that functional PRDM9 is required to complete meiosis, but despite its apparent importance, Prdm9 has been repeatedly lost across many animal lineages. Both the reason for mouse sterility in the absence of PRDM9 and the mechanism by which Prdm9 can be lost remain unclear. Here, we explore whether mice can tolerate the loss of Prdm9. By generating Prdm9 functional knockouts in an array of genetic backgrounds, we observe a wide range of fertility phenotypes and ultimately demonstrate that PRDM9 is not required for completion of male meiosis. Although DSBs still form at a common subset of functional sites in all mice lacking PRDM9, meiotic outcomes differ substantially. We speculate that DSBs at functional sites are difficult to repair as a crossover and that by increasing the efficiency of crossover formation at these sites, genetic modifiers of recombination rates can allow for meiotic progression. This model implies that species with a sufficiently high recombination rate may lose Prdm9 yet remain fertile.

[Supplemental material is available for this article.]

Meiotic recombination is essential for the production of euploid gametes. Recombination requires the repair of programmed double-stranded DNA breaks (DSBs), and this, in turn, results in an interchange of genetic information between homologous chromosomes (Bolcun-Filas and Schimenti 2012). In mice, humans, and several other mammals, the positions of recombination-initiating DSBs are primarily determined by DNA sequence-specific binding of the meiosis-specific histone methyltransferase PRDM9 (Baudat et al. 2010; Myers et al. 2010; Parvanov et al. 2010; Grey et al. 2018). All evidence to date has suggested that PRDM9 is essential for meiotic progression in mice because knocking out or disrupting the Prdm9 gene results in extensive asynapsis of homologous chromosomes and in complete meiotic arrest (Hayashi et al. 2005; Fairfield et al. 2011; Flachs et al. 2012; Weiss et al. 2012; Sun et al. 2015; Imai et al. 2017; Diagouraga et al. 2018; Thibault-Sennett et al. 2018). Nonetheless, dogs, swordtail fish, and even a human female (Muñoz-Fuentes et al. 2011; Narasimhan et al. 2016; Baker et al. 2017) have lost functionally essential functional sites. Studies in a limited number of mouse strains showed that functional PRDM9 is required to complete meiosis, but despite its apparent importance, Prdm9 has been repeatedly lost across many animal lineages. Both the reason for mouse sterility in the absence of PRDM9 and the mechanism by which Prdm9 can be lost remain unclear. Here, we explore whether mice can tolerate the loss of Prdm9. By generating Prdm9 functional knockouts in an array of genetic backgrounds, we observe a wide range of fertility phenotypes and ultimately demonstrate that PRDM9 is not required for completion of male meiosis. Although DSBs still form at a common subset of functional sites in all mice lacking PRDM9, meiotic outcomes differ substantially. We speculate that DSBs at functional sites are difficult to repair as a crossover and that by increasing the efficiency of crossover formation at these sites, genetic modifiers of recombination rates can allow for meiotic progression. This model implies that species with a sufficiently high recombination rate may lose Prdm9 yet remain fertile.

Results

PRDM9 is dispensable for sperm production in PWD males

A loss-of-function allele of Prdm9, Prdm9tm1Ymat (henceforth, Prdm9tm1), causes a complete arrest of spermatogenesis in C57BL/6 (B6) mice (B6tm1) (Table 1; Sun et al. 2015) and in mice with a mixed B6 and 129P2 genetic background (Hayashi et al. 2005). We crossed this allele into distantly related PWd/Ph (PWD) and...
found that male PWD mice homozygous for this loss-of-function Prdm9 allele (PWD*tm/tm) produced sperm (Table 1, column SC). In contrast, no sperm were found in B6*tm/tm males, indicating a partial rescue of meiotic progression in the PWD genetic background. No PRDM9 protein was detected in PWD*tm/tm males (Supplemental Fig. S1); therefore, the phenotypic differences between B6*tm/tm and PWD*tm/tm were not the result of Prdm9 trans-activation or duplication of the Prdm9 locus in PWD. We validated the partial rescue with a second Prdm9-loss-of-function allele (Prdm9em1Fore) (see B6*tm/tm, B6*em/tm, and (B6*tm/tm × PWD) × PWD*N3F1*tm/tm in Table 1). Testicular tubule sections confirmed the presence of tubules with sperm and spermatids in PWD*tm/tm but not in B6*tm/tm males (Fig. 1). Nonetheless, many tubules were devoid of spermatids in PWD*tm/tm and the sperm count in PWD*tm/tm was far lower than that in wild-type mice. None of the four tested PWD*tm/tm males produced offspring.

PRDM9 is important for meiotic synthesis in PWD males and females

Meiosis arrests at pachytene in B6*tm/tm male mice. This arrest manifests as a failure to synapse all pairs of homologous autosomes and a failure to form the specialized chromatin compartment for the sex chromosomes (sex body) (Hayashi et al. 2005; Flachs et al. 2012). Both full synopsis and sex-body formation are required for normal meiotic progression in spermatocytes (for review, see Burgoyne et al. 2009). Although just 2% of pachytene nuclei appeared to be normal (fully synapsed autosomes and observable sex body) in B6*tm/tm males (Table 1, column nPS; Supplemental Fig. S2), 41% were normal in PWD*tm/tm. Homologous chromosome synopsis in pachynema depends on the repair of programmed DSBs (Mahadevaiah et al. 2001). PWT*tm/tm males displayed 13% more DSB repair foci than equivalent nuclei from B6*tm/tm (B6*tm/tm = 193 ± 24 foci; PWD*tm/tm = 219 ± 35 foci; P = 0.003, Wilcoxon test). A similar increase in DSBs was seen in PWD*tm/tm compared to B6*tm/tm (B6*tm/tm = 201 ± 30 foci; PWT*tm/tm = 225 ± 34 foci; P = 0.001, Wilcoxon test) (also see Balcova et al. 2016), implying that the loss of Prdm9 function was not responsible for the increase (Fig. 2A). One speculative hypothesis is that more DSBs would allow more chromosomes to synapse; for example, if two extra chromosomes could synapse in B6*tm/tm pachytene cells (from the 13% increase in DSBs), close to 9% of pachytene nuclei would be fully synapsed. Finally, we found that the crossover rate (number of MLH1 foci) (see Methods) in normal PWT*tm/tm pachytene spermatocytes was similar to that in PWD*wt/wt males (Table 1; Fig. 2B). In addition, the density of MLH1 foci on synapsed chromosomes was lower in abnormal PWD*tm/tm pachytene spermatocytes (1.33 foci/autosome, n = 245 autosomes), than in normal PWD*wt/wt (1.56 foci/autosome, n = 1330) or PWD*tm/tm pachynemas (1.54 foci/autosome, n = 1311) (Supplemental Table S1), suggesting that cells with a higher recombination rate may better tolerate Prdm9 loss. MLH1 foci were rarely found (2/30 cells) on the synapsed chromosomes of abnormal B6*tm/tm pachynemas. In conclusion, PWD males lacking functional PRDM9 exhibit a partial meiotic arrest. This contrasts mice with the same PRDM9 defect in a B6 background, where the meiotic arrest is close to absolute.

We next examined meiotic progression in B6*tm/tm and PWD*tm/tm females. At birth, the number of oocytes in both B6*tm/tm and PWD*tm/tm was comparable to wild type (Fig. 3D; Supplemental Fig. S3). Two days after birth, there was a notable reduction in both backgrounds (Fig. 3C; Supplemental Fig. S3), and all oocytes were completely eliminated in Prdm9-deficient adults (2–5 mo), unlike in wild-type controls (Fig. 3B; Hayashi et al. 2005). The post-birth oocyte loss is likely mediated by mechanisms that detect oocytes in which DSB repair and/or homolog synopsis were aberrant (Di Giacomo et al. 2005; Kogo et al. 2012; Wojtasz et al. 2012; Bolis et al. 2014; Cloutier et al. 2015; Rinaldi et al. 2017; Qiao et al. 2018). Both B6*tm/tm (Hayashi et al. 2005; Diagouraga et al. 2018) and PWD*tm/tm adult females had small ovaries (Fig. 3A, B), and none of four mated PWD*tm/tm females produced offspring.

In PWD*wt/wt embryonic oocytes (~18 d post coitum), pachytene synopsis was often incomplete. On average, 14/20

Table 1. The effect of Prdm9 removal on male mouse fertility

| Background (FF) and genotype | n  | TW/BW (mg/g) | SC (million) | nPS (%) | COR | Fertility |
|------------------------------|----|--------------|--------------|---------|-----|-----------|
| PWD*wt/wt                    | 12 | 6.4 ± 0.4    | 25 ± 7       | 94 ± 10 | 29.3* | F(9/9)    |
| PWD*tm/tm                    | 30 | 2.3 ± 0.3    | 0.4 ± 0.4    | 41 ± 24 | 29.6 | S(0/4)    |
| B6*tm/tm                     | 5  | 1.7 ± 0.1    | 0.00 ± 0.0   | 2 ± 2   | 24.4*b | S(0/1)    |
| B6*em/tm                     | 4  | 1.9 ± 0.1    | 0.00 ± 0.0   | 1 ± 1   | 24.4*b | S         |
| B6*em/tm                     | 4  | 2.0 ± 0.1    | 0.00 ± 0.0   | 2 ± 2   | 24.4*b | S         |
| ((B6*tm/tm × PWD)N3F1 em/em) | 7  | 2.3 ± 0.6    | 1.5 ± 4      | ND      | ND   | (3/6)     |
| ((B6K × B6*tm/tm) × PWD)N3F1 im/im | 8 | 4.0 ± 1.0    | 5.3 ± 6      | ND      | ND   | (2/6)     |
| C3H*tm/tm                    | 6  | 1.6 ± 0.1    | 0.00 ± 0.0   | 9 ± 3   | 22.7  | S         |
| (B6 × PWD)F1 em/em           | 7  | 2.3 ± 0.3    | 0.12 ± 0.2   | 35.6    | ND   |                |
| (B6 × PWD)F1 tm/tm           | 10 | 2.4 ± 0.3    | 0.19 ± 0.3   | 37 ± 17 | 28.9  | S(0/4)    |
| (PWW × P6)F1 tm/tm           | 4  | 1.9 ± 0.5    | 0.00 ± 0.0   | 2 ± 2   | 24.1*c | S         |
| (B6.PWD-ChrX.1 × PWD)F1 tm/tm | 9 | 3.1 ± 0.4    | 0.6 ± 1      | ND      | 28.9  | ND      |
| (B6.PWD-ChrX.1 × PWD)F1 tm/tm | 4 | 2.4 ± 0.2    | 0.00 ± 0.0   | ND      | ND   | S         |
| (B6 × PWD)F1 tm/tm            | 8  | 3.2 ± 0.8    | 7.3 ± 4      | 46 ± 10 | 26.8  | F(3/6)    |
| (C3H × PWD)F1 tm/tm           | 6  | 3.0 ± 0.7    | 7.6 ± 6      | 49 ± 7  | 29.5  | F(6/6)    |
| (PWW × C3H)F1 tm/tm           | 6  | 2.0 ± 0.1    | 0.00 ± 0.0   | 3 ± 1   | 24.7*b | S         |

FF, female parent shown first; tmm, the Prdm9tm1Fore null allele; emm, the Prdm9em1Fore null allele; n, number of males; TW, weight of paired testicles; TW/BW (mean ± SD), relative testis weight in mg of TW per gram of body weight (BW); SC, sperm count (millions) in the entire epididymis; nPS, percentage of normal (full synopsis) pachytene spermatocytes (of all pachynema); COR, mean crossover rate (autosomal MLH1 foci/cell; for details, see Supplemental Table S1); F, fertile (number of offspring-producing males per total mated males); S, sterile (no pups and/or SC = 0); ND, not determined. For males with other genotypes and for statistics, see Supplemental Table S4.

*Balcova et al. 2016.

bValues for the genetically matched wild-type males, because these Prdm9-deficient males lack pachynema.

*Hattacharyya et al. 2013.
chromosomes were fully synapsed and just 14% of oocytes (9/64) exhibited a full complement of synapsed chromosomes. Like in males, the loss of functional Prdm9 exacerbated asynapsis at pachytene (Fig. 3E,F); PWD\textsuperscript{tm/tm} oocytes had, on average, 3/20 synapsed chromosomes, and none of the 149 PWD\textsuperscript{tm/tm} oocytes studied had more than 14 synapsed chromosomes (Fig. 3). In males, we only considered nuclei with more than nine synapsed chromosomes as “pachytene-like.” Only 10/149 PWD\textsuperscript{tm/tm} oocytes met this strict criterion because of the severity of the defect in females. To compare with B6 mice, we examined ovaries from newborn mice. Both PWD\textsuperscript{wt/wt} and B6\textsuperscript{wt/wt} mice had fewer fully synapsed pachytene nuclei (PWD\textsuperscript{wt/wt} = 0.28; B6\textsuperscript{wt/wt} = 2/85 oocytes) than their wild-type littersmates (PWD\textsuperscript{wt/wt} = 6/30; B6\textsuperscript{wt/wt} = 21/32 oocytes) (Fig. 3G,H; Supplemental Fig. S4). Thus, the lack of functional Prdm9 appears to result in synapsis defects and sterility both in B6 and PWD female mice.

**Functional Prdm9 is not required for fertility in male mice**

Having discovered that mice can make few sperm without functional Prdm9, we next hypothesized that hybrid vigor may help to restore full fertility to mice lacking Prdm9. By backcrossing (N) and intercrossing (F) B6 and PWD mice heterozygous for functional Prdm9 (for the breeding scheme, see Supplemental Fig. S5), we indeed obtained fertile Prdm9-deficient mice ((B6\textsuperscript{em/wt} × PWD) × PWD) N3F1\textsuperscript{em/em} (Table 1). In an effort to increase fertility of these mice, we bred (B6\textsuperscript{em/wt} × PWD) × PWD) N3F1\textsuperscript{em/em} males with their heterozygous sisters for several generations (for the breeding scheme, see Supplemental Fig. S6). These mice were designated (B6 × PWD) F\textsuperscript{em/em} (Table 1) and indeed had a higher sperm count (P = 0.014, Wilcoxon test) than (B6\textsuperscript{em/wt} × PWD) × PWD) N3F1\textsuperscript{em/em}. Three of six (B6 × PWD) F\textsuperscript{em/em} males sired pups. To test the generality of this phenomenon, we performed a similar crossing strategy with two other pairs of mouse strains: B6 crossed with PWK/Ph (PWK), and C3H crossed with PWD. Each strategy yielded fertile mice that lacked functional Prdm9; 2/6 (PWK × B6) N3F1\textsuperscript{em/em} males and 6/6 (C3H × PWD) F\textsuperscript{em/em} sired pups. In conclusion, mice can fully tolerate the loss of Prdm9 and remain fertile.

**DSB hotspot distribution is similar in fertile and sterile Prdm9-deficient mice**

The only known function of the PRDM9 protein is to designate the genomic locations at which meiotic DSBs occur. In the absence of functional PRDM9, meiotic DSBs occur at functional sites in the genome (Brick et al. 2012). It has been proposed that DSBs at functional sites may be difficult to repair, and that this may cause the sterility of B6\textsuperscript{Prdm9}\textsuperscript{-/-} males ((B6\textsuperscript{em/wt} × PWD) × PWD) N3F1\textsuperscript{em/em} (Table 1). As expected, these hotspots mostly coincided with non-PRDM9-mediated H3K4me3-marked histones, often at transcription start sites (Fig. 4A,C). Thus, we conclude that meiotic DSBs at functional sites are not a complete impediment to meiotic progression.
A locus on Chromosome X affects fertility of Prdm9-deficient male mice

Prdm9 governs hybrid sterility in male F1 hybrids of B6 and PWD mice (Mihola et al. 2009; Flachs et al. 2012). Genetic mapping identified variants of a locus on Chromosome X (Hstx2) that modulates fertility in these mice (Bhattacharyya et al. 2014). We therefore investigated whether this locus may also modulate fertility in mice lacking Prdm9.

We first examined meiotic progression in Prdm9-deficient male F1 mice from reciprocal crosses of Prdm9wt/wt and Prdm9tm/tm. In F1 mice with a PWD mother (and therefore, a PWD Chromosome X and Hstx2PVD), just 2% of pachytene spermatocytes exhibited full synapsis and no sperm were found ((PWD × B6)F1tm/tm). In contrast, no sperm was found in mice carrying Hstx2B6 ((B6 × PWD)F1tm/tm). This mirrors a role of Hstx2B6 in facilitating fertility in otherwise sterile (PWD × B6)F1 hybrids (Bhattacharyya et al. 2014).

Discussion

In this work, we identified male mice with multiple genetic backgrounds that are fertile despite lacking a functional PRDM9 protein. This unambiguously demonstrates that PRDM9 is not required for completion of meiosis in males, and that its function is to target meiotic DSBs, and in its absence, meiotic DSBs occur at functional sites in the genome. We
Figure 3. Similar meiotic arrest in B6 and PWD females lacking functional Prdm9. (A) Reduced adult ovary weight in both B6<sup>tm/tm</sup> and PWD<sup>tm/tm</sup> females indicative of arrested meiosis. There is no significant difference between B6<sup>tm/tm</sup> and PWD<sup>tm/tm</sup>. Black dots with bars symbolize mean ± SD; color dots depict individuals. Tukey’s box-and-whisker plots show median values and quartiles. H/E-stained sections of paraffin-embedded ovaries of adult (B) and 2 d after birth (dpp) (C) mice. Arrows point to oocytes present in the controls but mostly absent in both PWD<sup>tm/tm</sup> and B6<sup>tm/tm</sup>. (D) Oocytes from newborn ovaries immunostained with anti-MSY2 (YBX2) antibody Abcam (ab33164) and DAPI. The number of oocytes was comparable in all tested females at birth (D), but decreased in both types of Prdm9-deficient mice compared to littermate controls 2 d after birth (C), suggesting pachytene arrest attributable to oocyte attrition. (E) Reduced homologous chromosome synapsis in PWD<sup>tm/tm</sup> oocytes (~18 d post coitum). Colors distinguish oocytes derived from individual embryos (2 PWD<sup>wt/wt</sup>, 4 PWD<sup>wt/tm</sup>, and 6 PWD<sup>tm/tm</sup>). Black dots represent mean values, and bars represent SDs. (F) Representative spread nuclei from the PWD<sup>tm/tm</sup> and PWD<sup>wt/tm</sup> oocytes in E. Nuclei were immunostained for the synaptonemal complex lateral element (SYCP3), central element (SYCP1; yellow color indicates synapsis), and gamma H2AFX (a chromatin marker of DNA damage). (G) Decreased chromosomal synapsis in both PWD and B6 newborn oocytes lacking functional PRDM9. (H) Representative nuclei from the PWD<sup>tm/tm</sup> and PWD<sup>wt/tm</sup> oocytes in G immunostained for the synaptonemal complex central element (SYCP1; green indicates synapsis), recombination nodules (MLH1), and centromeres (CENT). See Supplemental Fig. S4 for nuclei from B6<sup>tm/tm</sup> and B6<sup>wt/wt</sup>.
Figure 4. Hotspot locations are consistent in all mice lacking functional PRDM9. (A) DSB hotspots in wild-type mice (B6 × PWD)F7 and PWDwt/wt are absent in mice lacking functional PRDM9 (B6tm/tm, PWDtm/mtm, B6 × PWD)F7/mtm). Raw coverage is shown in 150-bp windows. Each panel is scaled to the maximum value. DSB hotspots are the peaks in DMC1-SSDS coverage. Red arrows represent Ensembl gene models. (B) Hotspot locations were conserved in all mice lacking functional PRDM9. The maximum reciprocal hotspot overlap (±500 bp) between pairs of samples is shown. (C) Hotspots in mice lacking functional PRDM9 occur at non-PRDM9-mediated H3K4me3 (non-hotspot) marks and transcription start sites (TSSs). All H3K4me3 peaks (Methods) that did not coincide with a wild-type DSB hotspot were considered non-hotspot H3K4me3. For TSSs, the overlap with Ensembl transcript’s 5′ ends ± 500 bp is shown. These features rarely coincide with DSB hotspots in wild-type mice. Note that B6vtm/vtm H3K4me3 ChIP-seq data are from (Baker et al. 2014). B6vtm/vtm and PWDvtm/vtm DMC1-SSDS data are from Smagulova et al. (2016).

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demonstrated that this redirection of DSBs away from functional sites is not sufficient for the successful completion of meiosis, because DSB hotspots occurred at broadly similar sites in all mice lacking functional PRDM9. Indeed, this is consistent with the fertility of species that lack PRDM9 such as dogs, swordtail fish, and birds (Muñoz-Fuentes et al. 2011; Axelsson et al. 2012; Singhal et al. 2015; Baker et al. 2017), where recombination is favored and more pachytene nuclei successfully synapse all pairs of homologs (Table 1; Supplemental Fig. S8). These data imply that the loss of functional Prdm9 reduces the efficiency at which DSBs can be repaired from the homolog in such a way as to allow synopsis (likely as a crossover). If this efficiency dips below a critical threshold, one or more chromosomes fail to synapse and pachynema cannot be completed. In our model, the shortest chromosomes would be least likely to receive a repair-competent DSB, because the length of chromosomes is proportionate to the number of DSBs per meiosis (Supplemental Fig. S9). Thus, short chromosomes would be less likely to synapse with their homolog during pachytene. In mice with reduced DSB formation (Spo11-hypomorph) (Kauppi et al. 2013), short chromosomes asynapsis occurs disproportionately frequently. This is consistent with our model.

The extent of the pachytene defect resulting from Prdm9 deficiency does not linearly correlate with the observed sperm count and/or fertility of the animals; for example, 40% of pachytene spermatocytes synapse fully in PWDvtm/mtm mice, but the sperm count is just 1.2% of that in wild-type mice (sperm count SC = 0.4 million). In contrast, sperm production is far more robust in mice with just slightly more normal pachytene cells ((C3H × PWD)F1/wt/mice, nPS = 49%, sperm count = 7.6 million; and (B6 × PWD)F7/mtm/wt, nPS = 46%, sperm count = 7.3 million). Previous studies have demonstrated that the failure to complete pachytene in a critical percentage of spermatocytes can effectively shut down spermatogenesis. This is thought to result from apoptotic signaling from the spindle assembly checkpoint (SAC; senses misaligned chromosomes), and this signal may affect "normal cells" because of communication among spermatocytes in the germinal syncytium (Faisal and Kauppi 2016). Indeed, Faisal and Kauppi (2016) found that the critical percentage of normal pachytene cells required for sperm production is somewhere between 30% (infertile) and 50% (fertile). Thus, we conclude that the removal of functional Prdm9 makes it more difficult for cells to complete full chromosome synopsis in pachytene. How this is translated into fertility is likely dependent on other factors, such as SAC potency, and can result in a broad range of fertility phenotypes ranging from sterility to fertility.
Figure 5. Sperm-producing Prdm9-deficient mice have a relatively high crossover rate. Black dots depict means, and error bars represent SDs. Each all mice carrying epididymal sperm (sperm count above the detection limit represents a single mouse genetic background (data from Table 1). Hstx2 is not the only locus affecting crossover rate in these mice, as judged from the distribution of its alleles in the groups. The mean crossover rate in animals that “Produce Sperm” was significantly higher than that in animals that did not (P, probability, Wilcoxon test).

The Hstx2 locus, also known as meiotic recombination 1 (Meir1), is a 4.7-Mb region (containing multiple genes) on Chromosome X that regulates the crossover rate in mice (Balcova et al. 2016). Consistent with published data, we found that in male mice differing at the Hstx2 locus, those carrying the Hstx2<sup>Bo</sup> allele had a 30% higher crossover rate than those carrying the Hstx2<sup>PWD</sup> allele. We argue that this difference resulted in sperm production in the Prdm9-deficient Hstx2<sup>Bo</sup> carrying mice, but not in mice with the Hstx2<sup>PWD</sup> allele. The Hstx2<sup>Bo</sup> allele was previously shown to alleviate hybrid incompatibilities governed by Prdm9 genotype that result in the sterility of male F1 hybrid mice from a cross between PWD females and B6 males (Bhattacharyya et al. 2014). We propose that this alludes to a unifying principle of fertility rescue in these two systems in which sterility arises in a Prdm9-dependent manner. In both systems, there is a clear DSB repair defect; in Prdm9 knockout mice, this may result from difficulties in repairing DSBs at functional sites; in sterile hybrids, DSB repair as a crossover is compromised at loci where Prdm9 binding sites have been destroyed in one parental lineage (Davies et al. 2016; Smagulova et al. 2016). In both systems, some but not all homologous chromosomes synapse at pachytene, and in both systems, fertility can apparently be rescued by increasing the recombination rate. We propose that in both systems, increasing the crossover rate (or perhaps the number of DSBs) elevates the likelihood of a crossover on every pair of homologous chromosomes, and thus increases the number of viable pachytene spermatocytes. Sufficient rescue can overcome the critical threshold of “normal” pachytene cells to assure robust sperm production and fertility.

Females lacking functional PRDM9 were sterile, with a similar pachytene asynapsis defect as in males. However, the number of pachytene nuclei with asynapsed chromosomes was far higher in females than in males. In our working model, this defect could theoretically be rescued by increasing the DSB/crossover rate or by subverting the SAC. The alternate explanation for the observed sex dimorphism is that the requirement for Prdm9 differs between males and females.

One implication from our model is that DSB repair as a crossover is more efficient in the presence of functional PRDM9. Thus, one of the roles of PRDM9 may be to facilitate a lower recombination rate than would otherwise be possible. This would help to explain why Prdm9 is widely conserved but repeatedly lost during evolutionary history, because species with a sufficiently high recombination rate could lose functional PRDM9 with little or no reproductive consequences. It remains to be seen whether this is the case.

Methods

Ethics statement

The European Community Council Directive 86/609/EEC, Appendix A of the Council of Europe ConventionETS123, and the Czech Republic Act for Experimental Work with Animals (Decree No. 207/2004 Sb, and the Acts Nos. 246/92 Sb and 77/2004 Sb) were obeyed during the laboratory animal care and experiments. Permissions Nos. 61/2013 and 10/2016 were issued by the ethics committee for the work with animals of the Institute of Molecular Genetics in Prague.

Mice

The PWD/Ph and PWK/Ph mouse strains were described in Gregorová and Forejt (2000). The C57BL/6J strain was from The Jackson Laboratory (Stock no. 000664), C3H/N was from Velaz, Czech Republic. The knockout line Prdm9<sup>em1Fore</sup> (MGI allele No. 3623909; here Prdm9<sup>em</sup>) was generated in a 129P2/OlaHsd background by replacement of the first five coding exons with LacZ. This was maintained on the pure B6 background. The Prdm9<sup>em</sup> line was transferred by 10-times-repeated backcrossing to PWD; the N7 generation was devoid of 129- and B6-specific sequences except for Chr 17 (proximal 31.4 Mb harboring Prdm9), and this differential segment was reduced in the N12 generation to about 11.7 Mb (position 15.1–26.8 Mb). The fertility parameters of the Prdm9-deficient mice resulting from the strains carrying the differential segments of 31.4 and 11.7 Mb were similar (Supplemental Table S2). The Prdm9<sup>em1Fore</sup> allele (abbreviated Prdm9<sup>em</sup>) strain was prepared by injecting programmed artificial zinc-finger endonuclease (Sigma-Aldrich) into C57BL/6Ncdl zygotes, which resulted in a 4-bp deletion in the ninth exon of Prdm9 encoding the PR/SET domain (MGI allele No. 5501109). The (B6 × PWD)F7 line was prepared by repeated intercrossing of fertile Prdm9-deficient (B6<sup>em</sup> × PWD) × PWD)<sup>em</sup> males to heterozygous littersmates (for the breeding scheme, see Supplemental Fig. S6). The resulting (B6 × PWD)F7 mice carried the Hstx2<sup>Bo</sup> allele. C57BL/6J mice carrying different sections of the PWD Chromosome X were previously generated (Gregorová et al. 2008). We abbreviate these mice as B6.PWD-Chr# (where # indicates the introgressed chromosome fragment from PWD). F1 hybrids carrying regions of Chr X differing at Hstx2 and lacking PRDM9 were constructed by outcrossing the subconsonamic strains B6.PWD-ChrX.1 (carries Hstx2<sup>Bo</sup>) and B6.PWD-ChrX.1s (carries Hstx2<sup>PWD</sup>) to B6<sup>em</sup> animals, intercrossing the Prdm9<sup>em1</sup> female offspring to PWD<sup>em</sup> males, and then genotyping their F1 offspring for Prdm9<sup>em</sup> and Chr X (for the breeding scheme, see Supplemental Fig. S7).
Genotyping and phenotyping

PCR genotyping conditions have been published (Flachs et al. 2012), except for genotyping primers for the Prdm9em allele: 5′-ACCTTAGTGTGACTGTGC-3′ and 5′-CAAACTCTGTC TGAAACCC-3′; these primers amplify 111 bp and 107 bp from the wild type and mutant, respectively; annealing is carried out at 59°C, and products can be resolved in 5% agarose. Body weight (BW) and testis weight (TW) were taken for adult males (older than 11 wk). Sperm count (SC) was obtained from both epididymides. Embryos were dissected from mothers 18 d after the beginning of mating (mating lasted 1–3 d). Chromosome spreads (slides with surface-spread nuclei) were prepared from embryonic ovaries or whole adult testis using hypotonic treatment (Anderson et al. 1999). Spreads were stained with anti-DMC1 (Santa Cruz sc-22768), anti-RAD51 (Santa Cruz sc-8349), anti-SYCP3 (Abcam ab151500), and anti-centromere (see above) antibodies. The MLH1 protein marks the sites of all genetic crossovers (Anderson et al. 1999; Wang et al. 2017). MLH1 foci were analyzed by confocal microscopy (Leica TCS SP5 AOBS Tandem) to ensure that only MLH1 foci that colocalized with the chromosomal axis (SYCP1) were counted. In cells with fully synapsed autosomes, we omitted cells in which any chromosome lacked an MLH1 focus. This allows for counting of properly stained, closely matched, and staged cells. In abnormal pachytene cells (carrying 1–4 asynapsed autosomal pairs), MLH1 foci were counted per synapsed autosomal pair. In the studies of male pachytene synapsis, cells carrying more than 10 asynapsed chromosomal pairs were considered as zygotene. Immunocytochemistry and histology were performed as described previously (Baker et al. 2012, 2014; Bhattacharyya et al. 2013, 2014). The anti-MSY2 (YBX2) antibody was from Abcam (ab33164).

Chromatin immunoprecipitation (ChIP) and western blotting

DMC1 ChIP-SSDS was done using snap-frozen adult testes as described previously (Khil et al. 2012; Brick et al. 2018). H3K4me3 was assayed by ChIP-seq in 13- to 15-d-old testes according to Baker et al. (2015b). Western blotting was performed as described (Baker et al. 2015a).

Sequencing data analyses

DMC1-SSDS samples were aligned to the mm10 reference genome, peak calling was performed, and hotspot strength was calculated as described (Brick et al. 2018; for sample quality metrics and hotspot counts, see Supplemental Table S3). H3K4me3 ChIP-seq data were aligned to the reference mm10 genome using BWA 0.7.12 (Li 2013). Peaks for H3K4me3 (used in Fig. 4) were called as follows. H3K4me3 ChIP-seq data from 12 dpp B6 (SRR1035576) (Baker et al. 2014) and corresponding input DNA data (SRR1035578) (Baker et al. 2014) were aligned to the reference mm10 genome using BWA 0.7.12 (Li 2013). Aligned BAM files were converted to BED files with BEDTools (Quinlan and Hall 2010) and duplicate reads were discarded. NCIS (Liang and Koles 2012) was used to determine the background correction ratio \(R_{bg}\). MACS2 version 2.1.0.20150731 (Zhang et al. 2008) was used with the following arguments to call peaks: --ratio \(R_{bg}\) -g mm -bw 1000 --keep-dup all --local 5000 -t [SRR1035576 BED file] -c [SRR1035578 BED file].

Data access

ChIP data generated in this study have been submitted to the NCBI Gene Expression Omnibus (GEO; https://www.ncbi.nlm.nih.gov/geo/) under accession number GSE109874.

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