PRDM9 binding organizes hotspot nucleosomes and limits Holliday junction migration

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In mammals, genetic recombination during meiosis is limited to a set of 1- to 2-kb regions termed hotspots. Their locations are predominantly determined by the zinc finger protein PRDM9, which binds to DNA in hotspots and subsequently uses its SET domain to locally trimethylate histone H3 at lysine 4 (H3K4me3). This sets the stage for double-strand break (DSB) formation and reciprocal exchange of DNA between chromatids, forming Holliday junctions. Here we report genome-wide analyses of PRDM9-dependent histone modifications using two inbred mouse strains differing only in their PRDM9 zinc finger domain. We show that PRDM9 binding actively reorganizes nucleosomes into a symmetrical pattern, creating an extended nucleosome-depleted region. These regions are centered by a consensus PRDM9 binding motif, whose location and identity was confirmed in vitro. We also show that DSBs are centered over the PRDM9 binding motif within the nucleosome-depleted region. Combining these results with data from genetic crosses, we find that crossing-over is restricted to the region marked by H3K4me3. We suggest that PRDM9-modified nucleosomes create a permissible environment that first directs the location of DSBs and then defines the boundaries of Holliday junction branch migration.

[Supplemental material is available for this article.]

A hallmark of sexual reproduction is the exchange of genetic material between chromatids through the process of meiotic recombination. This exchange is essential for proper chromatid segregation during meiosis, and improper segregation can lead to germ cell death or the production of abnormal gametes that result in spontaneous abortion and developmental disorders (Paigen and Petkov 2010). Rather than randomly located along chromosomes, mammalian recombination sites are restricted to 1- to 2-kb regions, termed hotspots, whose locations are determined by the meiosis-specific protein PRDM9 (Baudat et al. 2010; Berg et al. 2010; Myers et al. 2010; Parvanov et al. 2010; Berg et al. 2011). PRDM9 is a DNA-binding, zinc finger protein that also trimethylates histone H3K4 (Hayashi et al. 2005), a chromatin modification enriched at mammalian hotspots (Buard et al. 2009; Grey et al. 2011; Billings et al. 2013). An irreversible commitment to meiotic recombination occurs when DNA double-strand breaks (DSBs) are introduced at hotspots by the conserved enzyme SPO11 (Keeney 2008; Cole et al. 2010a). These DSBs are preferentially repaired using a homologous chromatin as the template and either involve the reciprocal exchange of DNA, leading to crossing-over, or to repair without exchange, resulting in gene conversion (Kohl and Sekelsky 2013). After introduction of DSBs, the DNA is processed to produce single-stranded ends that are bound by the meiotic recombinase DMC1 to facilitate strand invasion of the homologous chromatin. Repair of DSBs involving reciprocal exchange of DNA proceeds by formation of double Holliday junctions which undergo enzymatically assisted branch migration, allowing the exact positions of crossing-over to be moved from the original site of the DSB (Neale and Keeney 2006; Handel and Schimenti 2010). Physical confirmation that sites of PRDM9 binding determine where recombination can occur came with the first genome-wide maps of DSBs, by mapping DMC1-associated DNA, showing that nearly all DSBs overlap with sites of PRDM9-dependent H3K4me3 (Smagulova et al. 2011; Brick et al. 2012). In the absence of Prdm9, DSBs are redirected to other H3K4me3-modified loci (Brick et al. 2012). These ectopic DSBs are inefficiently repaired, leading to meiotic arrest (Hayashi et al. 2005), thus highlighting another important role of PRDM9. Prdm9 is highly polymorphic in many mammalian species including primate and mouse populations (Thomas et al. 2009; Berg et al. 2010; Buard et al. 2014; Kono et al. 2014). These polymorphisms are almost entirely constrained to variation in the number of C-terminal zinc finger repeats and, within each zinc finger, nonsynonymous substitutions found at the three amino acids that contact DNA. Sequencing large populations of both wild-caught and laboratory strains of Mus, Kono et al. (2014) found 57 different alleles in a sample of 116 mice, while Buard et al. (2014) found 113 alleles in their sample of 257 mice. These allelespecific combinations of zinc fingers dictate the nucleotide sequences at which PRDM9 can bind DNA and locally trimethylate hotspot nucleosomes, ultimately marking the position of meiotic recombination (Grey et al. 2011; Billings et al. 2013).

To facilitate genome-wide identification of trimethylated hotspots, we used co-isogenic mouse strains carrying two different Prdm9 alleles, both in the C57BL/6J (B6) background (Fig. 1A). One strain was B6 itself, containing Prdm9<sup>Dom2</sup>. The other strain was constructed by replacing exon 12 of Prdm9<sup>Dom2</sup>, which encodes the entire zinc finger domain, with the corresponding exon from the Prdm9<sup>Dom</sup> allele found in the CAST/EiJ (CAST) strain. The only
which originated in erosion (Boulton et al. 1997). Insufficient time for its hotspot sequences to undergo evolutionary PRDM9Dom2-dependent ones, and these PRDM9Cst sites, on average, appreciably more PRDM9 Cst-dependent H3K4me3 sites than putative allele-specific sites of modification (Fig. 1B,C). We found zinc finger array of PRDM9, unique H3K4me3 positions represent strains, representing common features such as active promoters (Fig. 1B). Most H3K4me3-modified loci are shared between the two co-isogenic strains allowed by high-throughput DNA sequencing (ChIP-seq), to generate genome-wide H3K4me3 maps from the two co-isogenic strains (Fig. 1B). Most H3K4me3-modified loci are shared between the two strains of mice (in all figures, the Prdm9Cst allele originates from the CAST/EiJ strain and is represented in red, while Prdm9Dom2, the endogenous allele for C57BL/6J, is represented in blue). (B) H3K4me3 coverage profile from a representative 100-kb window of chromosome 1. H3K4me3 peaks can be unique or shared between the two strains of mice (in all figures, the Prdm9Cst allele originates from the CAST/EiJ strain and is represented in red, while Prdm9Dom2, the endogenous allele for C57BL/6J, is represented in blue). (C) Venn diagram showing quantity and identity of H3K4me3 peaks. Prdm9Dom2 results in more unique H3K4me3-marked hotspots. (D) H3K4me3 signal for strain-specific peaks. Prdm9Cst hotspots have higher methylation levels than Prdm9Dom2 in the B6 background (red: Prdm9Dom2; blue: Prdm9Cst).

Figure 1. H3K4me3 ChIP-seq allows detection of PRDM9-dependent sites using co-isogenic strains of mice. (A) PRDM9Dom2 and PRDM9Cst differ both in the number of zinc finger domains and sequence-specific identity of individual domains. The known domain structure of PRDM9 includes an N-terminal KRAB domain (Krüppel-associated box), an SSXRD (SSX repression domain), a PR/SET domain (methyltransferase activity), an invariant zinc finger domain, and a C-terminal zinc finger array. The three most polymorphic positions in the individual zinc finger domains are indicated by their amino acid identity. (B) H3K4me3 coverage profile from a representative 100-kb window of chromosome 1. H3K4me3 peaks can be unique or shared between the two strains of mice (in all figures, the Prdm9Cst allele originates from the CAST/EiJ strain and is represented in red, while Prdm9Dom2, the endogenous allele for C57BL/6J, is represented in blue). (C) Venn diagram showing quantity and identity of H3K4me3 peaks. Prdm9Dom2 results in more unique H3K4me3-marked hotspots. (D) H3K4me3 signal for strain-specific peaks. Prdm9Cst hotspots have higher methylation levels than Prdm9Dom2 in the B6 background (red: Prdm9Dom2; blue: Prdm9Cst).

Results

PRDM9-dependent chromatin modification is extensive in meiosis

We used immunoprecipitation of micrococcal nuclease (MNase)-digested chromatin from 12-d post-partum male germ cells, followed by high-throughput DNA sequencing (ChIP-seq), to generate genome-wide H3K4me3 maps from the two co-isogenic strains (Fig. 1B). Most H3K4me3-modified loci are shared between the strains, representing common features such as active promoters and enhancers. Because the only difference between strains is the zinc finger array of PRDM9, unique H3K4me3 positions represent putative allele-specific sites of modification (Fig. 1B,C). We found appreciably more PRDM9Dom2-dependent H3K4me3 sites than PRDM9Dom2-dependent ones, and these PRDM9Cst sites, on average, have a greater level of trimethylation (Fig. 1C,D). One contributing factor to this difference may be that the Prdm9Cst knock-in allele, which originated in Mus musculus castaneus, has been in contact with the M. m. domesticus genome of B6 for only a few generations, insufficient time for its hotspot sequences to undergo evolutionary erosion (Boulton et al. 1997).

We estimated the number of PRDM9Dom2-activated sites in an average meiosis. Using quantitative PCR of our H3K4me3 positions that varied in activity, we established the proportionality between ChIP results and the fraction of total “input” DNA from spermatocytes that was H3K4me3-modified in the same samples (Supplemental Fig. S1). This enabled conversion of the number of ChIP-seq reads at a hotspot to the fraction of chromatids with trimethylated nucleosomes. Summing across all hotspots, after correcting for the presence of four chromatids in a meiotic cell and for the estimated representation of early meiotic prophase cells (see Methods), we estimate that there are ~4700 ± 400 PRDM9-modified sites in the B6 background in an average meiosis. This conservative estimate of PRDM9-activated positions is ~15 times higher than the current estimated number of DSBs per meiotic cell based on immunohistochemistry (Cole et al. 2012), raising the question of what additional factors determine the likelihood of progression from PRDM9-activation to induction of a DSB.

Hotspots contain symmetrically organized nucleosomes around a central nucleosome-depleted region

In vivo, DNA is packaged by wrapping 147–150 bp of DNA around a histone octamer to form a nucleosome, with 38–75 bp of inter-nucleosome linker DNA (Kornberg 1977; Valouev et al. 2011; Yamada and Ohta 2013). Earlier studies are contradictory regarding the question of how nucleosomes are organized within hotspots. Getun et al. (2010) reported that hotspots could contain regions >1 kb nearly devoid of nucleosomes in an arrangement that is similar for meiotic and somatic cells; however, this is difficult to accept given the much repeated observation that hotspots sequences are enriched for H3K4me3 nucleosomes. Shenkar et al. (1991) reported one hotspot that contained a DNase I hypersensitivity site overlapping the position of recombination, indicating the absence of a nucleosome there; whereas in contrast, Smagulova et al. (2011) reported that sites of DSBs typically contain a well-placed nucleosome at their center. In our data, H3K4me3-modified nucleosomes at individual hotspots show a distinct, symmetrical oscillatory pattern, organized around a central nucleosome-depleted region (NDR) (Fig. 2A,B). Predicted PRDM9 DNA binding motifs for each allele, derived from in vivo binding in this study closely matches the region identified by in vitro studies as most sensitive to base-pair substitution (Billings et al. 2013). Together, these data suggest that PRDM9 binds hotspots in the central NDR.

To calculate hotspot NDR lengths, we located nucleosomes, at individual hotspots, in the –1 and +1 positions relative to our
PRDM9 binds predicted motifs in the NDRs

In order to confirm that the motifs found in the NDRs are the targets of PRDM9 binding, we chose four hotspots as models, two for each allele, and tested their binding specificity using an in vitro mobility shift assay. We found that only the allele that activates a hotspot in vivo binds DNA from that NDR in vitro, and the minimum DNA fragment required for binding always contains a close match to the consensus motif (Fig. 4; Supplemental Figs. S4, S5). These binding experiments also revealed several properties of PRDM9-DNA interactions. First, competition for PRDM9 binding in vitro is allele-specific. DNAs from different hotspots activated by PRDM9-DNA interactions. First, competition for PRDM9 binding match to the consensus motif (Fig. 4; Supplemental Figs. S4, S5).

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To characterize average hotspot nucleosome positioning for both Prdm9 alleles, this analysis was extended to the whole genome by sequencing all nucleosome-associated DNA in spermatocytes (MNase-seq). These data show that in C57BL/6J spermatocytes, nucleosomes are organized in a symmetrical pattern only within hotspots containing the DNA binding motif recognized by PRDM9Dom2 (Fig. 5C, blue line) and not at PRDM9Cst-activated hotspots (Fig. 5C, red line). The reverse is true in the B6-Prdm9Cst-KI strain, where hotspots containing the PRDM9Cst motif only show nucleosome organization (Fig. 5D, red line). Because the knock-in strain was created on the B6 background, hotspot sequences are identical between the two strains and would not contribute to a difference in nucleosome positioning (cis-control); the only difference is the identity of the Prdm9 allele (trans-control). In contrast, nucleosome organization at CTCF binding sites is similar in these two co-isogenic strains, highlighting the allele-specific nature of PRDM9 sites (Supplemental Fig. S3C,D). Collectively, these data show that PRDM9 binding actively reorganizes nucleosomes at hotspots.

**Figure 5.** Nucleosome positions at hotspots are variable prior to PRDM9 binding. (A) Schematic of an idealized PRDM9-modified hotspot showing signal intensity representing nucleosome positions. Primers for qPCR were designed to amplify DNA from either the −1 or +1 nucleosome flanking the NDR (peak) or from within the NDR (valley). (B) Ratio of nucleosome peak vs. nucleosome-depleted valley signal in two genetic backgrounds. qPCR was performed on MNase-digested spermatocyte chromatin DNA from both strains (blue: DNA from Prdm9Dom2; red: Prdm9Cst). Error bars represent standard deviation of triplicate measurements from three biological replicates. (**) P < 0.01, (****) P < 0.001; one-sided t-test. Hotspots Pbx1 and chr8:88,114 Mb are PRDM9Dom2-activated; hotspot Hlx1 is PRDM9Cst-activated. (C) Aggregation plot showing nucleosome positions at hotspots inferred from MNase-seq of C57BL/6J spermatocytes. Only hotspots containing a motif that is predicted to be recognized by the endogenous Prdm9 allele show organized nucleosomes (blue: aggregation of Prdm9Dom2 sites; red: Prdm9Cst sites). (D) Aggregation plot of Prdm9Dom2 and Prdm9Cst hotspots from sequencing of mono-nucleosome DNA prepared from B6-Prdm9Cst-KI spermatocytes.
chose two hotspots (Fig. 6A, C) containing SNPs distributed across the H3K4me3 interval and expanded the mapping of all crossovers at these hotspots using the repertoire of available SNPs. At both hotspots, the NDR and PRDM9 binding sites reside at the midpoint of all recombination events, and crossovers are limited to the region of H3K4me3 modification (Fig. 6B, D). These observations were confirmed for previously mapped recombinants at the HJv1 hotspot activated by PRDM9<sup>Cst</sup> (Parvanov et al. 2009) and the A3 hotspot activated by PRDM9<sup>Cdom2</sup> (Supplemental Fig. S6; Kelmenson et al. 2005; Cole et al. 2010b).

**DSBs are located at the center of the NDR**

To determine the locations of DSBs in relation to the PRDM9-dependent NDRs, we combined our present data with the DSB data generated from DMC1 ChIP-seq in testis from B6 mice (Brick et al. 2012). Aggregating the genome-wide DSB signals in relation to the center of the NDR, we find that the peak of the mean DMC1 tag distribution signal is precisely aligned with the center of the PRDM9 NDR (Fig. 6E). The shape of the DSB distribution curve is expected if DSBs are located at the center of the NDR, given the 100- to 200-bp size of the DNA fragments from which sequences were obtained (Brick et al. 2012) and the ~1-kb length of DMC1 coated filaments (Neale and Keeney 2006). Previous estimates found that, for a particular hotspot, the median of the DMC1 tag distribution could predict the location of meiotic DSBs with a precision of ~180 nt (Smagulova et al. 2011); we now show that the peak of the average signal aligns with the center of PRDM9-dependent NDRs. These data support the idea that meiotic DSBs occur within NDRs and fit well with the above observation that crossing-over occurs symmetrically around the PRDM9 binding site.

**Discussion**

We have characterized the genome-wide positions of PRDM9-dependent chromatin modifications by comparing two inbred mouse strains that only differ in the zinc finger array of PRDM9. Because PRDM9 is an H3K4 methyltransferase, our fine-scale mapping of H3K4me3-modified nucleosomes reflects nucleosome positions after PRDM9 binding. These post-PRDM9 binding snapshots of nucleosome positions reveal several previously unknown features of hotspot chromatin architecture and provide mechanistic insight into how PRDM9 activates hotspots (Fig. 6E).

Hotspots contain a central nucleosome-depleted region that, on average, covers ~120 bp (Fig. 3). PRDM9 can trimethylate 1–2 nucleosomes beyond the ~1 and +1 nucleosomes flanking the binding sites. The ability of the catalytic domain of PRDM9 to reach this far may depend on supercoiling of chromatin to bring nucleosomes into closer proximity to PRDM9 in three-dimensional space. Previously, Valouev et al. (2011) reported a 130-bp NDR for the zinc finger protein CTCF and a 93-bp NDR for the REST (also known as NRSE) repressor in human cell lines. We have found a 134-bp NDR for mouse CTCF in spermatocytes (Supplemental Fig. S3), indicating that our MNase-seq methodology is robust.

Nucleosomes surrounding the PRDM9 NDR are organized in an oscillatory pattern, with increasing variability as a function of distance from the binding site, similar to CTCF. This symmetrical distribution of nucleosomes around an NDR fits with the previously established barrier model for nucleosome organization first proposed in yeast (Mavrich et al. 2008) and subsequently confirmed in mammalian cells (Valouev et al. 2011), which proposes that nucleosomes are packed against a well-positioned barrier whose location depends on either primary DNA sequence or the presence of a sequence-dependent DNA-binding factor. NDRs, such as the one reported for PRDM9 here, are likely general features of sequence-specific DNA-binding factors, with the length of the depleted region being dependent on the protein complex examined.

We also find that the PRDM9-dependent H3K4me3 level is positively correlated with NDR length. If H3K4me3 level is a measure of the frequency with which a hotspot binds PRDM9 and becomes trimethylated, which in turn depends on the affinity of PRDM9 for its binding site in that hotspot, it appears that the extent to which PRDM9 reorganizes nucleosomes at a hotspot depends on its affinity for the PRDM9 binding site in that hotspot. That is, the stronger the binding, the greater the effect on nucleosome positioning.

PRDM9 binding alters hotspot nucleosome organization. Nucleosomes are only symmetrically phased around PRDM9 binding sites when the appropriate Prdm9 allele is present (Fig. 5). This process probably requires the activity of the SWI/SNF family of ATP-dependent remodeling enzymes (Narlikar et al. 2013),
which are often recruited to reposition or remove existing nucleosomes. Currently, it is unclear whether accessibility of PRDM9 to its binding site prior to nucleosome remodeling ultimately influences the recombination rate. Two hotspots assayed here, \textit{Pbx1} and \textit{Hlx1}, are among the most highly active for recombination in male mice on chromosome 1 (2.6 and 3.0 CM, respectively) (Parvanov et al. 2009; Billings et al. 2013). However, in the absence of PRDM9 binding, the two hotspots do differ in chromatin configuration; the \textit{Pbx1} binding site is more susceptible to MNase digestion than the \textit{Hlx1} binding site, suggesting differing chromatin states and therefore accessibility to trans-acting factors. To fully answer this question, nucleosome positions need to be resolved in the absence of PRDM9 binding at the individual hotspot level and genome-wide hotspot recombination rates measured to correlate the two.

The position of crossing-over is restricted to the area modified by PRDM9 (Fig. 6). This suggests that not only does PRDM9 direct the DSB machinery to its genomic binding locations (Brick et al. 2012), but that it is also necessary for creating an environment permissive to branch migration. This limiting of branch migration to the area of trimethylated nucleosomes was seen when the span of H3K4me3 nucleosomes among hotspots varied from 1000 to 2500 bp (Fig. 6A,C). Since DSBs are introduced in close proximity to PRDM9 binding sites (Fig. 6E), we conclude that Holliday junctions can migrate equally well in either direction but only within the space defined by trimethylated nucleosomes. Whether other factors help limit the locations of Holliday junctions is not known, but it is clear that branch migration does not extend beyond the chromatin environment created by PRDM9-dependent modification.

While our data show a clear relationship between H3K4me3 track length and crossover track length, most meiotic DSBs are repaired as noncrossover (NCO) gene conversions. Previous work has shown that gene conversion tracks are generally short (10 to 120 bp), found within hotspots, and are limited to a narrower window than crossovers (Ng et al. 2008; Paigen et al. 2008; Cole et al. 2010b). It has been proposed that the position of noncrossovers may correlate with the position of DSBs (Cole et al. 2010b). If this is true, they should cluster within the NDR at hotspots. Indeed, for the hotspots \textit{Hlx1} and \textit{A3}, where noncrossovers have been extensively mapped, the majority of noncrossovers are concentrated at the center of hotspots near the NDR, although they can still be found outside of the NDR (Ng et al. 2008; Cole et al. 2010b). It is possible that some DSBs occur outside the NDR, given the DMC1 distribution (Fig. 6E); alternatively, NCOs outside the NDR could result from extended single-strand resection, followed by limited repair synthesis. Co-conversion events, when multiple SNPs within a hotspot are converted together in the same meiotic cell, are rare. Co-conversions at the \textit{Hlx1} hotspot were only found within the PRDM9*^{Dom2} binding site, and 92% of co-conversions found at the \textit{A3} hotspot are in a 100-bp window within the \textit{A3} NDR, which includes the predicted PRDM9*^{Dom2} binding site (Supplemental Fig. 56B; Cole et al. 2010b).

Our findings relate to the question of what determines the likelihood that a methylated hotspot will transition to DSB formation in mammals. In yeast, a species that lacks \textit{Prdm9}, Pan et al. (2011) found that Spo11 preferentially cuts DNA within the NDRs found near promoters. In addition, two recent studies demonstrated the molecular link between H3K4me3-dependent binding of the yeast protein Spp1 in proximity to promoter NDRs and the meiotic chromosome axis containing DSB machinery, including Spo11 (Acquaviva et al. 2013; Sommermeyer et al. 2013). Similar to yeast, the DSBs found in mutant mice lacking \textit{Prdm9} are frequently located at H3K4me3 sites near promoters (Brick et al. 2012). Here we show that PRDM9 binding actively creates an NDR that overlaps with the position of DSBs in mice, thereby potentially providing Spo11 access to DNA.

Taken together, the data presented here demonstrate that PRDM9 has a large influence over the state of hotspot chromatin. This influence is most apparent at the initiation of recombination, when it reorganizes hotspot nucleosomes, and later in its ability to constrain the migration of Holliday junctions and the location of genetic crossing-over.

**Methods**

**Mouse strains and genotyping**

C57BL/6j (stock number 000664) and CAST/Eij (stock number 009928) mice were obtained from The Jackson Laboratory. All animal experiments were approved by the Animal Care and Use Committee of The Jackson Laboratory (Summary #040008).

B6-Prdm9*^{Dom2}/Kpgn was generated by targeted replacement of exon 12 of B6 \textit{Prdm9*}^{Dom2}. The targeting construct was assembled by recombineering technology (Liu et al. 2003) such that exon 12 coding sequencing of the \textit{Prdm9*}^{Dom2} from the C57BL/6j strain (1,388 bp), along with the 5’ flanking sequence from intron 11 (55 bp) and the 3’ flanking sequence from the 3’ UTR sequence of exon 12 (106 bp), were replaced by the corresponding \textit{Prdm9*}^{Dom2} sequence from CAST/Eij. In addition, a neomycin-resistance cassette flanked by the FRT sites was inserted 621 bp downstream from the end of exon 12. The targeting construct was electroporated into an embryonic stem cell line derived from B6(Cg)-Tyr^a2/J (albino B6, JAX stock number 000058), and clones surviving G418 and ganciclovir were screened for the successfully recombined allele that had retained \textit{Prdm9*}^{Dom2} exon 12. Targeted clones with normal chromosome count were injected into C57BL/6j, and male chimeras were bred with B6(Cg)-Tyr^a2/J female mice to transmit the targeted allele. Subsequent mice were backcrossed with B6.Cg-Tg(ActFlpe)9205Dym/j (JAX stock number 005703) to excise the neomycin-resistance cassette. Mice were bred to homozygosity for the knock-in allele for this study.

For fine-mapping hotspots genetically, we used a library of DNA samples stored at –80°C taken from progeny of F1 B6 × CAST backcrossed to B6 (Paigen et al. 2008; Billings et al. 2010). SNPs between B6 and CAST were selected from publicly available databases for further genotyping crossovers. Recombination was detected as a transition from homozygous to heterozygous genotype using PCR-based KASP genotyping reagents (LGc Genomics) to narrow the interval to hotspot resolution. The final position of crossovers was confirmed by DNA sequencing of the entire hotspot.

**Spermatocyte isolation and crosslinking**

Crude isolation of germ cells was conducted from male mice at 12 d post-partum (dpp). Testes were isolated from six mice and the tunica albuginea removed. Tubules were placed in 20 mL DMEM (Life Technologies) with 0.5 mg of Liberase TM added (Roche) and incubated with shaking at 32°C for 15 min. Tubules were washed with 20 mL DMEM and resuspended in the same volume with 0.5 mg Liberase and 100 units (U) DNase I (Affymetrix). Tubules were disrupted by pipetting several times and incubated another 15 min at 32°C. After final incubation, tubules were disrupted by gentle pipetting and passed through an 80-μm Nitex nylon mesh filter (Sefar America). The resulting cell solution was pelleted by centrifugation at 1500 rpm. The cell pellet was washed with 10 mL DMEM supplemented with 25 U DNase I by gentle pipetting and
spun again. The cell pellet was resuspended in 5 mL DMEM and passed through the nylon filter a second time, and the number of cells was counted by hemocytometer. Typical yield of crude spermatocytes after isolation was 1 million cells per mouse.

For crosslinking, cells were pelleted and resuspended in PBS at 1 million cells per mL. Formaldehyde (Sigma-Aldrich) was added to a final concentration of 1%, and the cells were incubated with rotation at room temperature for 10 min. Crosslinking was stopped by drop-wise addition of glycine to 125 mM, followed by incubation for 5 min at room temperature with rotational mixing. The cells were pelleted, washed once with PBS, and snap-frozen in liquid nitrogen for storage at −80°C until use.

**Chromatin immunoprecipitation**

ChiP was performed using antibody against H3K4me3 (Millipore, #07-473) as previously described (Billings et al. 2013). To obtain sufficient material for high-throughput sequencing, four ChiP reactions from a single preparation of spermatoocyte chromatin were pooled after final DNA elution. ChiP DNA was concentrated using Agencourt AMPure XP beads (Beckman Coulter) following the manufacturer’s protocol. Quantitation of ChiP and input DNA was measured using the Qubit dsDNA HS assay (Life Technologies). Typically, 7–10 ng of ChiP DNA was obtained after concentration and used for the sequencing library preparation. For all samples, an equal amount of MNase-treated input DNA was sequenced as a control. Biological replicates for each genotype were prepared independently from different litters of mice. Libraries were prepared for sequencing using Bioo Scientific’s NEXTflex ChiP-Seq Kit (protocol version V11.11) without size selection. Amplification of the libraries was done with 20 μl ligation product, 16 μl water, 12 μL NEXTflex ChiP PCR master mix, 2 μL NEXTflex ChiP primer mix, and 14–18 cycles of PCR.

**High-throughput sequencing and data processing**

High-throughput sequencing was performed using the Illumina HiSeq 2000 platform. Raw sequences were aligned to the mouse genome NCBI Build 37 (mm9) using BWA (v. 1.2.3) (Li and Durbin 2009) with default settings. The alignments were subsequently filtered to retain only uniquely mapped reads. Peak calling was performed using MACS (v.1.4.2) (Zhang et al. 2008a) with input DNA for control and ChiP samples as treatment, setting the P-value to 0.01, and nodup = “all.” Because nucleosomes are protected from micrococcal nuclease digestion, and nucleosomes appear to be sharply positioned at hotspots, we chose to keep all mapped reads with the same 5’ starting position in the final coverage profiles and when quantifying ChiP-seq signal. All coverage profiles presented in figures are from MACS bedgraph output after tag shifting.

The only difference between the co-isogenic strains created here was the identity of the PRDM9 zinc finger array. Therefore, we were able to identify PRDM9-dependent H3K4me3 positions by categorizing peaks as unique to one strain. Biological replicates for both B6 and B6-Prdm9Cst mouse samples over Prdm9Dom2 samples were highly reproducible (Pearson’s r = 0.956 and 0.946, respectively), so data sets were pooled prior to final peak-calling. In our samples, MACS tended to identify more H3K4me3 peaks with low activity in the Prdm9Dom2 samples over Prdm9Dom2 samples. By inspecting the bedgraph coverage profiles manually, it was clear that many of the corresponding PRDM9-dependent regions contained visually clear uncalled peaks, thus creating many false positive unique PRDM9Dom2 positions. To overcome this, we implemented a conservative bioinformatics strategy to maximize identification of common peaks between strains by performing two rounds of peak calling. The first round of peak calling was performed using MACS with standard treatment and control samples with a P-value 0.01. After identifying common peaks between the two genotypes, all reads falling within those intervals were subtracted from both treatment and control alignment files and MACS run a second time to identify peaks with the same P-value. Common peaks between Prdm9Dom2 and Prdm9Dom2 from the second run were identified and pooled with common peaks from the first run. We then used this pooled common set of peaks to filter the original MACS results, removing common peaks found in both rounds, leaving peaks unique to the Prdm9 genotype.

This had the effect of enriching the sample for potential PRDM9 signals by eliminating all of the common H3K4me3 sites, even if they were low activity. While this strategy may miss potential quantitative differences in H3K4me3 strength due to PRDM9, such as sites that are coincident with other H3K4me3 sites, this strategy did satisfy our primary concern of identifying where in the genome PRDM9 modifies chromatin with the fewest false positive positions.

**Electrophoretic mobility shift assay**

EMSA was performed as previously described (Billings et al. 2013). DNA was 5’-labeled for EMSA using biotinylated primers (Eurofins MWG Operon) for PCR of genomic DNA. For the competition assay, synthetic DNA oligos from both strands were annealed to create unlabeled dsDNA and used at 20× concentration of labeled DNA. All oligo sequences are listed in Supplemental Table S1.

**Quantitative PCR**

Qualitative PCR was carried out using the Eppendorf Realplex cycler and the QuantiFast SYBR Green PCR kit (Qiagen) following the manufacturer’s protocols, except that primers were used at a final concentration of 0.5 μM to help eliminate primer dimers in later cycles. qPCR primers were designed using OligoPerfect (Life Technologies) with 40%–60%GC and 80- to 120-bp product size (primer sequences are listed in Supplemental Table S1). PCR was carried out for 40 cycles followed by melting curve analysis, and all samples were run in triplicate. Cycle number was determined using the automated threshold analysis.

For estimating the total number of PRDM9-dependent hotspots per meiosis, an equal volume of MNase-digested input DNA was analyzed alongside the ChiP DNA. Cycle threshold (Ct) values were calculated by averaging the technical replicates. Percent input was calculated by using the formula: % Input = 2^(-ΔCt Input − Ct ChiP)/100.

To determine peak-to-valley ratios, the peak and valley primer pairs were assayed using five 10-fold dilutions of genomic DNA to compare amplification efficiencies. For each hotspot, after qPCR the Ct values for peaks and valleys were subtracted at each dilution and plotted against the template amount. Only peak and valley primer sets that had a delta-Ct slope ~ 0.1 or less were used for subsequent analysis of the peak-to-valley ratio.

**Estimating the number of PRDM9-dependent H3K4me3 sites per meiosis**

qPCR of ChiP DNA allows us to estimate the average number of modified chromatids per meiosis. Using the hotspot Pbx1 as an example, we measured that 25% of the total input DNA had H3K4me3 at this hotspot (Supplemental Fig. S1). Given that there are four chromatids per meiotic cell, we would estimate that one chromatid in four is bound by PRDM9 and modified. However, the actual fraction of early prophase I meiotic cells in 12-dpp mouse testis is ~46% of the total cell population (Bellve et al. 1977). Therefore, we adjust the estimated number of H3K4me3 sites per meiosis by the fraction of cells in leptotene/zygotene. From this we
conclude that ~2.2 chromatids are modified at Pbx1 per average meiosis. To estimate the genome-wide number of sites, we extended the principle of this analysis to all hotspots.

The proportionality established between qPCR measurement of the fraction of modified loci (Supplemental Fig. S1) and ChIP-seq reads was applied to all PRDM9-dependent H3K4me3 positions from B6 mice. These converted values were summed, multiplied by four to correct for the number of chromatids per cell, and divided by 0.46, the estimated fraction of cells in early prophase I. The standard error for the slope of the regression in Supplemental Figure S1 is 0.00039, or ~8%, equaling ~400 hotspots. We believe this is a conservative estimate. If our crude isolation of germ cells resulted in 100% efficiency for recovering only early meiotic prophase I cells, then, at most, our estimate here is twice the actual value, still nearly 10 times the number of SBs seen by immunofluorescence. However, if our estimate of early prophase I meiotic cells is too high, and our actual recovery of germ cells is much lower, a more likely scenario, then we would have underestimated the number of PRDM9-dependent modifications per meiosis.

**Motif identification and aggregation plot**

We used the MEME Suite (v4.9.0, accessed online) for motif discovery and sequence searching (Bailey et al. 2009). MEME was used to identify allele-specific motifs with default parameters. Reasoning that PRDM9 would have the most access to DNA in the NDR, we narrowed the search space by restricting the DNA sequence to 500 bp surrounding the local minimum of ChIP-seq tag density near the center of H3K4me3 tracks. Only a single significant motif was identified for each Prdm9 allele. Once the motifs were identified using this narrow search space, FIMO was subsequently used to scan the entire 2 kb surrounding each NDR to determine if the motifs (1) were found outside of the NDR, and (2) showed allele-specific enrichment. To calculate fold enrichment, the number of 5’ start positions for each motif was counted for a sliding 20-bp window with steps of 1 bp. The number of occurrences of the active motif was divided by the number of occurrences of the inactive motif at each bp position. MEME was also used to identify the de novo CTCF motif in Supplemental Figure S3. CTCF binding sites in adult mouse testis were obtained from the Mouse ENCODE Project and are publicly available at The University of California, Santa Cruz Genome Browser or at GEO GSE918711 (The Mouse ENCODE Project Consortium et al. 2012).

To determine the nucleosome profile at hotspots, the ChIP-seq tag densities were aggregated using the ag-gpy script from the Aggregation and Correlation Toolbox (Jee et al. 2011) with nbins = 200 and radius = 1000. The first aggregation plots were anchored at the local minimum of H3K4me3 signal found within the NDR. Once the position of the allele-specific motifs were located using FIMO, aggregation plots were anchored using the 5’ starting position and strand orientation of the motif to center each hotspot. These motif-center aggregation plots are shown in Figure 2. Aggregation of DMC1 ChIP-seq was performed using the PRDM9snip motif positions. Sequencing data for DMC1 ChIP-seq were previously published (Brick et al. 2012) and can be found at NCBI GEO GSE35498.

**NDR length calculation**

The Aggregation and Correlation Toolbox script generates the aggregate profile for the PRDM9-dependent H3K4me3 sites. This only allows a single calculation of the distance between nucleosome peaks and is biased toward higher active hotspots. To examine the distribution of NDRs at hotspots in relation to H3K4me3 signal strength, nucleosome positions at individual hotspots were identified using NPS (v. 1.3.2) with default settings (Zhang et al. 2008b). The centers of the resulting genomic intervals align with the subpeaks in the coverage profile within hotspots (Fig. 3A). Hotspots with nucleosomes surrounding the identified motifs further than 500 bp apart were discarded, because manual spot inspection showed that these cases were skewed, having only one nucleosome per interval. NDR lengths at each hotspot were calculated by subtracting 150 from the distance between the centers of the −1 and +1 nucleosomes flanking identified motifs. This analysis was extended to the next pair of nucleosomes to find the linker distances to the left and right of the NDR.

**Data access**

Raw sequencing data, bedgraph files, and unique PRDM9 intervals generated in this study have been submitted to the NCBI Gene Expression Omnibus (GEO; http://www.ncbi.nlm.nih.gov/geo/) under accession number GSE52628.

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**Author contributions:** C.L.B. and S.K. performed all experiments. C.L.B. and M.W. performed computational data analyses. C.L.B., P.M.P., and K.P. contributed to the experimental design. K.P. supervised the study. C.L.B. and K.P. wrote the manuscript. All authors discussed the results and interpretation, read the paper, and approved the final manuscript.

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