Re-engineering the zinc fingers of PRDM9 reverses hybrid sterility in mice

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The DNA-binding protein PRDM9 directs positioning of the double-strand breaks (DSBs) that initiate meiotic recombination in mice and humans. Prdm9 is the only mammalian speciation gene yet identified and is responsible for sterility phenotypes in male hybrids of certain mouse subspecies. To investigate PRDM9 binding and its role in fertility and meiotic recombination, we humanized the DNA-binding domain of PRDM9 in C57BL/6 mice. This change repositions DSB hotspots and completely restores fertility in male hybrids. Here we show that alteration of one Prdm9 allele impacts the behaviour of DSBs controlled by the other allele at chromosome-wide scales. These effects correlate strongly with the degree to which each PRDM9 variant binds both homologues at the DSB sites it controls. Furthermore, higher genome-wide levels of such ‘symmetric’ PRDM9 binding associate with increased fertility measures, and comparisons of individual hotspots suggest binding symmetry plays a downstream role in the recombination process. These findings reveal that subspecies-specific degradation of PRDM9 binding sites by meiotic drive, which steadily increases asymmetric PRDM9 binding, has impacts beyond simply changing hotspot positions, and strongly support a direct involvement in hybrid infertility. Because such meiotic drive occurs across mammals, PRDM9 may play a wider, yet transient, role in the early stages of speciation.

In spite of its central role in evolution, the molecular mechanisms underlying speciation are not well understood. Only a small number of genes involved in speciation have been documented1, with only one such gene, Prdm9, known in mammals2,3. Prdm9 contributes to hybrid sterility in male (PWD × B6)F1 mice from crosses between male Mus musculus domesticus C57BL/6 (hereafter B6) and female Mus musculus musculus PWD/Ph (hereafter PWD)4. Although its genetic basis is only partially understood5,6, this hybrid sterility is characterized by failure of pairing (synapsis) of homologous chromosomes and an arrested meiotic prophase owing to lack of repair of recombination intermediates2. Homologous recombination, and synapsis, are interdependent, essential meiotic processes2, and evidence suggests synapsis often nucleates at recombination sites8. Aside from the PWD × B6 cross, Prdm9 allele and dosage have been associated with variation in measures of fertility and successful meiosis in many additional mouse crosses8.

PRDM9 has several functional domains, including a DNA-binding zinc-finger array, and a PR/SET domain responsible for histone H3 lysine 4 trimethylation (H3K4me3)10 (Fig. 1a). By binding to specific DNA sequence targets, PRDM9 directs the positions of the DSB events that initiate meiotic recombination11. This results in these DSBs and downstream recombination events clustering into small discrete regions called hotspots12,13. The PRDM9 zinc-finger array, encoded by a minisatellite repeat, is highly polymorphic within and across mammalian species3,14–18 and is among the fastest evolving regions in the genome, with strong evidence of natural selection influencing this evolution3. It is unknown whether PRDM9 zinc-finger array polymorphism has additional impacts, aside from direct alterations of DSB hotspot positions.

Humanizing Prdm9 restores hybrid fertility

To explore the DNA-binding characteristics of PRDM9, we generated a line of humanized B6 mice, by replacing the portion of mouse Prdm9 exon 10 encoding the zinc-finger array with the orthologous sequence from the human reference PRDM9 allele (the ‘B’ allele) (Fig. 1a). A feature of PRDM9 (explored further below) is the co-evolution of its zinc-finger array with the genomic background in which it sits13,17. Minisatellite mutational processes at PRDM9 can produce new alleles with duplications, deletions or rearrangements within the zinc-finger array, yielding an almost complete change in PRDM9 binding sites, and thus hotspot locations14. Because the human PRDM9 zinc-finger array evolved on a lineage separated from mice for ~150 million years, our experimental approach allows assessment of functional properties of a PRDM9 zinc-finger array unaffected by changes it has induced in the background genome, similar to new alleles randomly arising in the population.

Humanization of the Prdm9 zinc-finger array in B6 inbred mice had no effect on fertility (Extended Data Fig. 2) and cytogenetic comparisons revealed no significant impact on zygote DSB counts (DMC1 immunoreactivity, Extended Data Fig. 2b), crossover counts (MLH1 foci, Extended Data Fig. 2c), normal sex body formation (H2AX immunostaining, Fig. 1b) or quantitative measures of fertility and successful synapsis (see later). The full fertility of humanized mice implies there are unlikely to be any specific essential PRDM9 binding sites. One mechanism underlying speciation in many settings involves Dobzhansky–Muller incompatibilities: hybrid dysfunction arising from incorrect epistatic interactions1. Based on the above, it seems likely that if such interactions involving PRDM9 occur, they do not reflect constrained co-evolution of Prdm9 with specific genes.

To explore the role of PRDM9 in fertility directly, we crossed PWD females with B6B6/H males. As expected18, male (PWD × B6)F1 PWD/B6 hybrids (we use superscripts to indicate Prdm9 genotypes and write the female strain first in crosses) exhibited hybrid sterility as evidenced by failures in siring pups (Extended Data Fig. 2e), sex body formation...
Humanizing the recombination landscape

To characterize the consequences of re-engineering the zinc-finger domain on recombination, we generated high-resolution DSB maps for mice with different Prdm9 alleles and genomic backgrounds, using ChIP-seq single-stranded DNA sequencing\(^{19}\) on adult testes. This approach identifies single-stranded 3′ sequence ends decorated with DMC1, which arise as intermediates following creation of DSBs with DMC1, which arise as intermediates following creation of DSBs

Humanization restores symmetric binding

Next we examined DSB hotspot maps for hybrid males: infertile (PWD × B6)F1PWD/B6, reciprocal semi-fertile (B6 × PWD)F1B6/PWD, humanized rescue (PWD × B6)F1PWD/H, and reciprocal humanized rescue (B6 × PWD)F1H/PWD, with wild-type PWD for comparison. Sequence differences between the PWD and B6 genomes allowed us to determine whether individual hotspots in these hybrids were ‘symmetric’, with DSBS occurring equally on both chromosomes, or ‘asymmetric’, with a preference towards either the PWD or B6 chromosome (Supplementary Information Section 5).

We found that most DMC1 signal (71.8%) in (PWD × B6)F1PWD/B6 or (B6 × PWD)F1B6/PWD hybrids occurs within asymmetric DSB hotspots (Fig. 2a, Extended Data Fig. 6). Further, DSBS associated with the PWD allele occur largely on the B6 chromosome and those associated with the B6 allele occur largely on the PWD chromosome. We also measured asymmetry of the H3K4me3 mark at each hotspot and found the same pattern, confirming that DSB asymmetry largely reflects underlying differences in Prdm9 binding and methylation between the two homologues. This H3K4me3 asymmetry resembles that previously described for (B6 × CAST)F1B6/C\(^{-}\) hybrids\(^{17}\), but is considerably more extreme. Sequence differences directly disrupting Prdm9 binding motifs explain almost all cases of binding asymmetry (83.4% of PWD hotspots; 91.3% of B6 hotspots), and result from rapid mutational accumulation along the separate lineages from the common ancestor of B6 and PWD (Extended Data Fig. 6).

Thus such asymmetry can arise through meiotic drive to favor mutations disrupting Prdm9 binding motifs, within populations where these motifs are active. Any new mutation disrupting Prdm9 binding at a hotspot is preferentially transmitted to offspring: in individuals heterozygous for the mutation, DSBs occur preferentially on the non-mutant chromosome and are then repaired by copying from the mutant chromosome\(^{25}\). This phenomenon has been observed at Prdm9 binding motifs in human\(^{18}\) and mouse\(^{17}\) and causes a rapid accumulation of mutations disrupting Prdm9 binding. B6 and PWD Prdm9 alleles are largely subspecies-specific\(^{15}\), so only the B6 lineage has experienced strong erosion of the B6 binding motif, and only the PWD lineage has experienced strong erosion of the PWD binding motif. This asymmetric erosion explains the highly asymmetric Prdm9 binding sites in F1 hybrids.

Because the human allele has not been present in mice, its binding sites have not experienced erosion in the mouse genome. As a consequence, DSBS at hotspots attributable to the human allele occur...
mostly (57%) in symmetric hotspots, with the remaining, asymmetric hotspots mainly (84.2%) explained by the presence of mutations that coincidentally fall within the human PRDM9 binding motif (Fig. 2b). Conversely, only 30% of DSBs at hotspots attributable to the B6 allele occur in symmetric hotspots. An identical pattern is seen in the reciprocal crosses. Thus, a genome-wide effect of humanizing the mouse is to reprogram hotspot positions with the consequence that hotspot asymmetry is reduced in the hybrids.

Meiotic drive might also explain dominance, as seen for the human Prdm9 allele over the B6 allele in the B6B6/H mouse, because B6 motifs get across mice, without any additional information regarding other features such as SNP diversity, consistent with the sole difference between the infertile and rescue mice being the zinc-finger array of Prdm9. Moreover, only symmetric hotspots (in the infertile mouse, a minority) provide predictive power.

Next, we sought to understand the drivers of these chromosome-specific differences in DMC1 heat by testing various potential predictors of these differences between the infertile and humanized rescue mice (Supplementary Information). After an exhaustive search over possible models, given the predictors considered, the best-fitting model was highly predictive ($r^2 = 0.84$; Fig. 2f) and included only symmetric hotspot measures—the total H3K4me3 signal from PRDM9 binding on both homologues (that is, symmetrically) at the same hotspots, summed over the entire chromosome—for each of the three Prdm9 alleles ($P < 0.01$ in each case). The trans effect is thus explained by knowledge of only the direct differences in PRDM9 binding targets across mice, without any additional information regarding other features such as SNP diversity.
This excludes sequence differences at or near hotspots, or asynapsis itself, as a cause, and suggests that the total amount of symmetric binding on each chromosome, as opposed to a simple lack of asymmetric binding, plays an important role in predicting DMC1 heat. The direction of causality is reasonably clear (binding predates DSB formation, and the H3K4me3 mark lacks similar chromosome effects), while confounding influences should always be shared between the mice being compared and thus cannot alone explain the observed inter-chromosomal differences. It therefore appears that differences in the level of overall symmetric binding by PRDM9 drive downstream trans effects at chromosomal scales, with lower symmetric binding somehow increasing the number, or repair time, of DSBs even at distant hotspots.

### PRDM9 binding symmetry and synapsis

Sterile (PWD × B6)F1PWD/B6 hybrids show very high rates of asynapsis, particularly at specific chromosomes, and failure to form the sex body during early meiosis. By contrast, these phenotypes are completely rescued in (PWD × B6)F1PWD/H hybrids harbouring the humanized Prdm9 allele (Fig. 1c, d). Having seen a relationship between PRDM9 binding symmetry and the recombination process, we examined binding symmetry in relation to fertility. For different male mice, we measured three quantitative fertility phenotypes (Fig. 3a), and calculated several genome-wide measures of hotspot symmetry (Extended Data Fig. 8; Supplementary Information). We observed a significant correlation (P = 0.0083; rank correlation permutation test) between the DMC1 symmetry measures and the rate of proper synapsis among all nine mice studied. In humanized hybrid mice, the observed increase in symmetry was accompanied by improved fertility. Notably, this improvement effect is stronger than the Hstx2 modifier, responsible for the improvement effect is stronger than the Hstx2 modifier, responsible for the difference in asynapsis and fertility observed between the sterile and reciprocal hybrids (Fig. 3a). An additional mouse hybrid, (B6 × CAST)F1B6/CAST, showed intermediate PRDM9 binding symmetry and also an intermediate asynapsis level. Symmetry measures in homozygous mice (PWD, B6/B6, B6/H, B6/H) are, as expected, much higher than hybrids, and these mice show the highest symmetry rates and fertility measures.

Previous work showed that in the sterile (PWD × B6)F1 mouse, synapsis failure occurs at different rates among five chromosomes tested. We compared the reported asynapsis rates for these five chromosomes with the chromosome-specific DMC1 heat effects described above and found an identical ranking (P = 0.017 by rank correlation permutation test; Fig. 3b). Because these DMC1 heat effects are strongly predicted by symmetric H3K4me3 levels in the infertile mouse, this result implies that chromosomes with lower symmetric PRDM9 binding experience higher asynapsis rates. This may explain why lower symmetric PRDM9 binding genome-wide accompanies higher overall asynapsis rates among different mice.

Having found elevated DMC1 heat on chromosomes influenced by asynapsis (where homologous pairing fails), we examined DMC1 and H3K4me3 heats in two additional settings, where no homologue exists at all and thus homologous chromosome pairing cannot occur: the X chromosome in male mice, and separately in humanized hybrid mice at autosomal hotspots where the human PRDM9 binding motif lies within a region deleted in the PWD genome. In both these settings, we observed an elevation of DMC1 heat relative to autosomal hotspots bound symmetrically by PRDM9 (Extended Data Fig. 9). Elevation of DMC1 heat might, therefore, be a consistent signature of non-pairing of homologous chromosomes during meiosis. DMC1 elevation might be explained by an increased probability of a DSB occurring at that site, or by the DMC1 coating at breaks persisting for longer (delayed repair). However, the total number of RAD51-marked DSBs initiated per cell is tightly regulated, remaining unchanged even in Prdm9 knockouts, while in both knockouts and infertile hybrids, DSB marks indeed persist late into pachytene, suggesting a failure of repair. Therefore, the elevated DMC1 signals we observe may be explained by persistence of DMC1 where homologous repair is compromised or delayed.

### PRDM9–dependent homologue interactions

Given our chromosome-scale observations, we next asked whether symmetric binding at individual hotspots might also influence DMC1 heat. At each human-controlled hotspot in the humanized rescue, we measured the component of total DMC1 heat contributed by the B6 chromosome only, and compared this to the DMC1 heat for the same hotspot in B6/H. The comparison revealed (Fig. 4a) a remarkably strong, and clear, elevation in DMC1 heat in the hybrid mouse for the asymmetric hotspots (>90% asymmetry, towards binding of only the B6 chromosome), relative to the symmetric hotspots (those within 10% of complete symmetry). However, similar to the chromosomal analysis, H3K4me3 enrichment showed no difference whatsoever between symmetric and asymmetric sites in these mice (Fig. 4b). Indeed a comparison of H3K4me3 and DMC1 heat revealed a far higher ratio of average DMC1 heat to H3K4me3 enrichment for asymmetric relative to symmetric hotspots, across all hybrid mice, backgrounds, and Prdm9 alleles tested (Extended Data Fig. 9d). This effect reflects a consistent elevation of DMC1 heat at DSB sites on individual chromosomes when the homologue is not bound strongly (Extended Data Fig. 9e, f). This phenomenon cannot easily be explained by factors including local heterozygosity within or outside the PRDM9 motif, the type of mutation(s) disrupting PRDM9 binding, or outlier effects (Extended Data Figs 9, 10; Supplementary Information Section 13). Thus, elevation of DMC1 heat on the bound chromosome appears to be a universal feature of hotspots where PRDM9 binds asymmetrically, relative to symmetrically bound hotspots. By contrast, the results for H3K4me3 suggest the mark is deposited in an independent manner on each homologue (Supplementary Information Section 12.2). This implies the DMC1 heat elevation depends on a process involving symmetric PRDM9 binding, downstream of H3K4me3 deposition, involving both homologues. While we cannot exclude the possibility

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**Figure 3** | Humanizing PRDM9 restores proper synapsis and rescues fertility in hybrids. a. Fertility metrics in hybrid mice. Error bars represent bootstrap 95% confidence intervals (symmetry metric), or ± 1 s.e. (other metrics). b. Chromosome effects in DMC1 signals (as Fig. 2d) versus previously reported asynapsis rates for five chromosomes in infertile (PWD × B6)F1PWD/B6. Error bars, ± 1 s.e.

### Table 1

| Chromosome | Symmetry metric | Normal sex body | Proper synapsis | Testes weight |
|------------|----------------|----------------|----------------|--------------|
| Chr 1      |                |                |                |              |
| Chr 2      |                |                |                |              |
| Chr 3      |                |                |                |              |
| Chr 4      |                |                |                |              |
| Chr 5      |                |                |                |              |

| Chromosome | Observed DMC1 ratio (infertile/rescue) | Synapsis rate in infertile (%) |
|------------|---------------------------------------|-------------------------------|
| Chr 1      |                                       |                               |
| Chr 2      |                                       |                               |
| Chr 3      |                                       |                               |
| Chr 4      |                                       |                               |
| Chr 5      |                                       |                               |
that somehow more DSBs occur at asymmetric hotspots, this would require early, precise pairing of homologues, at least at hotspots, before DSB formation, to determine which hotspots are symmetrically bound. Although there is some evidence of pre-meiotic homologue association, current data do not suggest the existence of precise pairing before DSB formation. The alternative and more plausible explanation is that sites where PRDM9 binds asymmetrically simply experience a delay in DSB processing, delaying DMC1 removal compared to symmetric DSB hotspots. While our data represent the collective behaviour of populations of cells, this model suggests a mechanism of PRDM9-dependent interaction between homologues influencing downstream DSB processing operating within individual cells, which we discuss below (also Supplementary Information Section 14).

Discussion

Only one mammalian speciation gene, Prdm9, has so far been identified. Humanizing the zinc-finger array of Prdm9 redirects binding, thereby entirely reprogramming recombination hotspots, and in doing so reverses the hybrid infertility between musculus and domesticus subspecies. This modification mimics the consequences of a newly arising mutation in either or both subspecies would also restore hybrid fertility.

Multiple lines of evidence in our data, at chromosomal, whole-organism, and individual hotspot scales, strongly support novel roles for PRDM9 in the formation or processing of DSBs downstream of H3K4me3 deposition, dependent upon symmetric binding. Several aspects of our, and published, data (comparison between B6/B6 and B6/H mice, see also Supplementary Information) also mean that our results cannot be fully explained simply by sequence differences within or around hotspots, which do not specifically impact binding symmetry.

Pervasive asynapsis is proposed to be the underlying cause of infertility in hybrid mice. We observed a positive relationship between symmetric PRDM9 binding and correct synopsis of homologous chromosomes later in meiosis. Replacing the B6 allele with the humanized allele in hybrids greatly increases symmetric binding, restoring proper synopsis and fertility. Many apparently complex relationships have previously been reported between naturally occurring mouse Prdm9 alleles, allelic dosage, and quantitative fertility measures in hybrids. Each of ten manipulations shown or predicted to increase PRDM9 binding symmetry also increases meiotic success and fertility (Supplementary Information), supporting the idea that the link between binding symmetry and fertility might be very general, and causal.

The erosion of PRDM9 binding sites through meiotic drive also occurs at human hotspots, and probably across many mammals. In two populations separated for sufficient time, differential PRDM9 binding site erosion will decrease symmetry in hybrids, which is likely to decrease fertility levels (though not necessarily to the extreme of sterility). Therefore, PRDM9 may affect hybrid fertility levels across many mammalian species and so might repeatedly act in driving early speciation steps, although the rapid evolution of the zinc-finger array of PRDM9 implies an unexpected transience of this direct role. However, even subtle or transient PRDM9-driven reductions in fertility might still provide a selective advantage to additional mutations contributing towards speciation. This mechanism is different from the previously characterized causes of intrinsic hybrid incompatibilities, such as differences in ploidy, chromosomal rearrangements, or incompatibilities between genes. The extent to which it has been responsible for speciation in the natural world appears an interesting question for further research.

One plausible mechanism for the impacts of (a)symmetry involves a role for PRDM9 binding in aiding homology search—a process thought to involve invasion of the homologous chromosome to probe for homology by single-stranded DNA formed around DSBs. It has been suggested that synaptonemal complex proteins are loaded at some DSB sites and synopsis begins to spread. Extending this model, to incorporate the property that asymmetrically bound sites are less favourable for homology search, would parsimoniously predict each symmetry-related phenomenon we observed: DSBs at asymmetric hotspots would repair more slowly, elevating their DMC1 signal, and chromosomes with fewer symmetric hotspots overall would show delayed DSB repair and higher asynapsis rates, ultimately causing subfertility or sterility in animals with low symmetric binding. It is not known how homology search occurs efficiently in the nuclear environment, given the enormous potential search space of the genome, or why hotspots exist at all. Both phenomena could be explained by the above model in which homology search is focused at least partly on hotspot positions. Indeed hotspots might massively increase search efficiency by directing homology search to PRDM9 binding sites.

Online Content Methods, along with any additional Extended Data display items and Source Data, are available in the online version of the paper; references unique to these sections appear only in the online paper.

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Supplementary Information is available in the online version of the paper.

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Author Contributions S.R.M. and P.D. designed the study. B.D., N.A., E.B., D.B., R.D. and C.P. generated and bred the transgenic mice. F.P., G.Z. and R.D.C.O. performed and oversaw DMC1 ChIP-seq. N.A. performed H3K4me3 ChIP-seq. D.M. and C.G. performed cytological analysis. E.H., J.G.H., N.A., A.G.H., R.L. and K.B. performed and oversaw DMC1 ChIP-seq. N.A. performed H3K4me3 ChIP-seq. R.D. and C.P. generated and bred the transgenic mice. F.P., G.Z. and R.D.C.O. performed and oversaw DMC1 ChIP-seq. N.A. performed H3K4me3 ChIP-seq. D.M. and C.G. performed cytological analysis. E.H., J.G.H., N.A., A.G.H., R.L. and K.B. analysed the data. B.D., E.H., J.G.H., N.A., A.G.H., S.R.M. and P.D. wrote the paper, with input from all authors.

Additional Information The source data generated in this publication has been deposited in NCBI’s Gene Expression Omnibus under accession number GSE73833. Reprints and permissions information is available at www.nature.com/reprints. The authors declare no competing financial interests. Readers are welcome to comment on the online version of the paper. Correspondence and requests for materials should be addressed to S.R.M. (myers@stats.ox.ac.uk) or P.D. (donnelly@well.ox.ac.uk).
METHODS

Gene targeting in embryonic stem cells. A C57BL/6J (B6) mouse genomic BAC clone (RP23-159N6) encompassing the Prdm9 gene was used for subcloning of homology regions. A 7 kb Xmal/SpeI fragment upstream of exon 10 and a 2.5 kb BamHI/SpeI fragment downstream of exon 10 were used as S′ and 3′ homology regions, respectively. The intervening 4 kb SpeI/BamHI encoding exon 10 and flanking intronic regions were subcloned and an internal 1.4 kb BglII–HindIII fragment, containing the coding region of the zinc-finger array, was replaced with a synthesized fragment (Life Technologies) encoding the zinc-finger array from the human B allele. All coding sequence S′ of the first zinc finger and all 3′ untranslated regions (UTR) downstream of the stop codon were left as mouse. This humanized fragment was then assembled between the two homology arms, upstream of a neomycin selection cassette. Phix31 attP sites were incorporated immediately downstream of the S′ homology arm and between the PGK promoter and the neomycin phosphotransferase open reading frame to equip the locus with Phix31 integrase cassette exchange machinery for subsequent manipulations.

The completed targeting vector was linearized with Apal and electroporated into mycoplasma free C57BL/6J Jm8F6 embryonic stem cells (Extended Data Fig. 1a). Jm8F6 cells were a gift from B. Skarnes, Welcome Trust Sanger Institute. Following selection in 210 μg/ml G418, recombinant clones were screened by PCR to detect homologous recombination over the 3′ arm. A forward primer (5′-TACCGGTTGAGTGGAAATCTGTTGG-3′) binding within the PGK promoter was used together with a reverse primer (5′-TGACAGCAGAAAAACCCCTCCT-3′) binding downstream of the 3′ homology arm to amplify a 2.7 kb fragment from correctly recombined clones. Positive clones were examined for correct recombination at the S′ end by long range PCR using a forward primer (5′-CACAGGGACCTTATGTCGATG-3′) upstream of the 3′ homology arm and a reverse primer (5′-AGCAGAGGCTTTCACTCCTCACAA-3′) binding with the flanked cassette. Correctly targeted clones yielded a 10.4 kb ampiclon. Sanger sequencing analysis of the 10.4 kb ampiclon encompassing the 3′ homology arm with primer 5′-CCCTTCTCAATGATGATCCAAAT-3′ confirmed the correct integration of the S′ attP sequence, necessary for future manipulations of the locus. Southern blotting using a probe against neomycin was used to confirm that only a single integration event had occurred.

Mouse production and matings. Mice were housed in individually ventilated cages and received food and water ad libitum. All studies received local ethical review approval and were performed in accordance with the UK Home Office Animals (Scientific Procedures) Act 1986. Experimental groups were determined by genotype and were therefore not randomized, with no animals excluded from the analysis. Sample size for fertility studies and cytogenetics (see below) were selected on the basis of previously published studies.5,32 No statistical methods were used to predetermine sample size. All phenotypic characterization was performed blind to experimental group.

ES cells from correctly targeted clones were injected into albino C57BL/6J blastocysts and the resulting chimaeras were mated with albino C57BL/6J females. Successful germline transmission yielded black pups and F1 mice harbouring the humanized Prdm9 allele were identified using the above attP screening PCR. F1 heterozygous male mice were bred with C57BL/6J female mice. Paired testes weight was recorded and normalized against lean body weight, as assessed using EchoMRI-100 Small Animal Body Composition Analyzer. Immunohistochemistry analyses. Spermatoocytes from mice at approximately 9 weeks of age were prepared for immunohistochemistry by surface spreading33,34. In brief, the testis tunica was removed, the tubules cut with a razor blade and disassembled by pipetting, in PBS, containing protease inhibitors (Complete, Roche). Following centrifugation at 5,800 × g for 5 min, the cells were resuspended in 0.1 M sucrose, and spread onto the surface of slides in a drop of 1% paraformaldehyde

in PBS. The slides were left to dry for 3 h at room temperature, in a humidified box, then washed in 0.4% Photo-Flo 200 (Kodak), and then either immediately used, or stored at −80 °C. For immunohistochemistry the following antibodies were used: mouse anti-MLH1 (BD, 51-1327GR); mouse anti-phospho-H2AX (Millipore 05–636, clone JBW301); rabbit anti-SYCP1 (Novus Biological, NB300–229); rabbit anti-DMC1 (Santa Cruz Biotechnology sc-22768, H-100); mouse anti-SYCP3 (Santa Cruz Biotechnology sc-74569, D-1); rabbit anti-SYCP3 (Abcam ab15093). Non-specific binding sites were blocked by incubating the cells with 0.2% BSA in PBS for 40 min, followed by incubation in the primary antibodies overnight at 4 °C. Following washings in B/ABD buffer and detection with secondary antibodies, the slides were mounted in DAPI/Vectashield (Oncor) and analyzed with an Olympus BX60 microscope for epifluorescence, equipped with a Sensys CCD camera (Photometrics, USA), using Genus Cytovision software (Leica).

Spermatoocytes were staged based on SYCP3 staining. For MLH1 analysis, only pachytene with 19 or more foci, co-localizing with SYCP3, were considered, according to criteria defined by ref. 35. For DMC1 analysis, randomly selected cells, from any stage, were scored. The number of DMC1 foci per cell was counted using the PointPacker macro in ImageJ (64 bit). For SYCP1 analysis, only cells in pachytene were considered. Cells with 19 fully synapsed autosomes, with co-localizing SYCP1 and SYCP3 signals, and one XY body, were considered normal. For characterization of γH2AX, cells in pachytene or diplotene were scored, and we considered normal those where only a clearly identifiable XY body was covered by γH2AX signal.

Prdm9 expression via RT–PCR analysis. To verify the correct expression of the humanized Prdm9, we performed exon-spanning endpoint RT–PCR on whole testis cDNA prepared using Tetro reverse transcriptase (Bioline) using a forward primer binding to exon 9 (5′-CATTAGTGGGGGAACGAAAG-3′) and a reverse primer binding within the 3′ UTR, immediately downstream of the humanized zinc-finger domain encoded by exon 10 (5′-GGGATTTAATTCCCCCTCTAGTCA-3′) (Extended Data Fig. 1b). Q-expression analysis of Prdm9 transcripts was performed using two primer pairs (5′-GAATGAGAAGGCAAGACAGCA-3′ and 5′-GGACAACACACTGCTACAGA-3′; 5′-AGCCAGACGAAAACGCAACCA-3′ and 5′-GGGATTTAATTCCCCCTCTAGTCA-3′), amplifying regions within the 3′ UTR, normalizing against a housekeeping gene (Hprt; 5′-AGCTACTGTAATGATCAGTCAACG-3′ and 5′-CATTAAG) and using the Power SYBR Green PCR Master mix (Applied Biosystems) and a BioRad CFX96 cycler as per the manufacturer’s instructions. Relative expression was calculated using the Livak method. Expression of the humanized Prdm9 allele was unaffected by the genetic manipulation (Extended Data Fig. 1b, c).

Single-stranded DNA sequencing and double-strand break (DSB) detection. Testis cells from B6C3He/J, B6He3/6, wild-type PWD, the infertile (PWD × B6) F1PVD/B6, the reciprocal semi-fertile (B6 × PWD) F1PVD/B6, the humanized rescue (B6 × PWD) F1PVD/B6, (B6/CAST)F1B6/CAST, (B6/CAST)F2B6/H males were subjected to single-stranded DNA sequencing (SSD) as previously described.39

In addition, we used the sample C57BL/6J (sample 1) from ref. 21 aligned to mm9/NCBI38. This sample was also re-mapped to mm10/NCBI38 with a modified BWA mapper.39 Other samples from ref. 21, 9R (sample 2), 13R (samples 1 and 2) and Prdm9 knockout (B6E-/-) (sample 1)10, were also used in the comparative analysis of DNA sequences (Extended Data Fig. 3e). B6H101 and B6H51 libraries were prepared in D. Camerini-Otero’s lab (NIH) and sequenced on a HiSeq 2000 platform, using paired-end reads (read 1: 36 bp; read 2: 40 bp). These samples were aligned to the mouse mm9/NCBI38 reference genome. Wild-type PWD, the infertile (PWD × B6) F1PVD/B6, the reciprocal semi-fertile (B6 × PWD) F1PVD/B6, the reciprocal rescue (PWD × B6) F1PVD/B6, (B6/CAST)F1B6/CAST, (B6/CAST)F2B6/H males were subjected to single-stranded DNA sequencing (SSD) as previously described.39

In addition, we used the sample C57BL/6J (sample 1) from ref. 21 aligned to mm9/NCBI38. This sample was also re-mapped to mm10/NCBI38 with a modified BWA mapper.39 Variation in the number of sequenced fragments results from the difficulty to precisely assess the DNA concentration before sequencing. Only fragments with high mapping quality (at least 20) were retained for DSB hotspot calling, and only one copy of each duplicate fragment was conserved (here, a fragment is duplicated if there exists at least one other fragment mapping to the same genomic position). Supplementary Table 1 gives details about the samples considered in this study.

For ChIP-seq, ChIP-seq was performed as previously described39 with several modifications (noted here). In brief, the testis tunica was removed, the tubules dissociated with tweezers and fixed in 1% formaldehyde in PBS for 5 min followed by glycine quenching (125 mM final conc.) for 5 min at room temperature. Following washing steps, pellets were resuspended in 900 μl cold RIPA lysis

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buffer, dounced 20 times and sonicated in 300 µl aliquots in a Bioruptor Twin sonication bath at 4°C for three 10-min periods of 30 s on, 30 s off at high power, then cell debris was pelleted and removed and aliquots were pooled. For each sample, 50 µl of equilibrated magnetic beads were resuspended in 100 µl PBS/BSA and added to the chromatin samples for pre-clearing for 2 h at 4°C with rotation. Beads were removed, and 100 µl of pre-cleaned chromatin was set aside for the input control. 5 µl rabbit polyclonal anti-H3K4me3 antibody (Abcam ab8580) was added to the remaining pre-cleaned chromatin and incubated overnight at 4°C with rotation. 50 µl beads were washed and resuspended as before, then incubated with the chromatin samples for 2 h at 4°C with rotation. Beads were then washed and de-crosslinked at 65°C as described (26), and for input controls, 50 µl of pre-cleaned chromatin was used. After de-crosslinking, samples were further incubated with 80 µg RNase A at 37°C for 60 min and then with 80 µg Proteinase K at 55°C for 90 min. DNA was purified using a Qiagen MinElute cleanup kit.

ChIP and total chromatin DNA samples were sequenced in multiplexed paired-end Illumina libraries, yielding 51-bp reads. We prepared two biological replicates plus one genomic input control each for the ChIP (PWT + B6)F1 ChIP library, reciprocal (B6 × PWT)F1 ChIP library, and rescue (PWT × B6)F1 ChIP library. For the B6ChIP and B6ChIP mice, we prepared one biological replicate each (yielding 70–80 million usable read pairs per sample) and later split read pairs into pseudoreplicates. Sequencing reads were aligned to mm10 using BWA aln (v. 0.7.0) followed by Stampy (v. 1.0.23, option bmapkeepgoodreads), and reads not mapped in a proper pair with insert size smaller than 10 kb were removed. Read pairs representing likely PCR duplicates were also removed by samtools rmdup. Pairs for which neither read had a mapping quality score greater than 0 were removed. Fragment coverage was computed at each position in the genome and in 100-bp non-overlapping bins using in-house code and the sambamtools and bedtools packages.

DSB hotspot detection and comparison. To analyse DMC1 data, we developed a novel ChIP-seq peak caller, specific to DSB hotspots, which takes advantage of the shift in the mapping of single stranded DNA (ssDNA) reads between the 5’ and the 3’ DNA strands to call hotspots. These ssDNA segments are a consequence of the resection of DNA ends that accompanies a DSB and are isolated by DMC1 ChIP (19). For each hotspot, the caller estimates in particular the centre of the hotspot, and its heat, loosely defined as the number of reads mapping to this hotspot, and its heat, loosely defined as the number of reads mapping to this hotspot, and its heat, loosely defined as the number of reads mapping to this hotspot. The caller handles sample replicates and is able to call hotspots using several samples jointly. Details are given in Supplementary Information. DSB hotspots from two different samples are considered to overlap if their centres are at most 600 bp apart. DMC1 hotspot heats have been normalized so that the sum of hotspot heats is identical in each sample (and equals the sum of hits on a given chromosome). DMC1 hotspot heats are identical in each sample (and equals the sum of hotspot heats in B6ChIP sample (1)).

H3K4me3 enrichments have been computed at DSB hotspots identified by DMC1 ChIP-seq, using our previously published method (26) (Supplementary Information, Subsection 7.1). H3K4me3 hotspots have also been called de novo, without using DSB hotspots, using the same approach (26). The de novo calls were used to generate a list of regions likely to be trimethylated independently of PRDM9, by intersecting calls in mice with different Prdm9 alleles. In comparisons involving both DMC1 and H3K4me3 data, we excluded DSB hotspots contained in any of the PRDM9-independent trimethylated regions, and we used H3K4me3 enrichments computed at DSB hotspots (Supplementary Information). We only used de novo calls for analysis in Extended Data Fig. 6d, e.

DNA binding motif analyses. We developed a new, Bayesian, approach to identify DNA motifs enriched at DSB hotspots (Supplementary Information). We used FIMO (MEME Suite version 4.9.1) to find the locations of those motifs genome-wide. Using Mus farnsida and Mus caroli as outgroups, we reconstructed an ancestral reference genome for B6 and PWD. We could therefore identify on which lineage (B6 or PWD) mutations between these two mouse strains occurred. See Supplementary Information for details.

DSB hotspot assignment in hybrids. Using SNPs between the B6 and PWD genomes, each read pair from a hybrid DSB library (DMC1 ChIP-seq) is assigned to one of the categories ‘B6’, ‘PWD’, ‘unclassified’ or ‘uninformative’ using criteria detailed in Supplementary Information. For each DSB hotspot, the ratio of informative reads from the B6 chromosome was then computed as the fraction of ‘B6’ reads mapped within 1 kb of the hotspot centre, over the sum of ‘B6’ and ‘PWD’ reads in that region. We followed a similar approach for H3K4me3 ChIP-seq, but we further corrected for background signal.

Chromosome effects. To test for statistically significant differential elevation of DMC1 (or H3K4me3) hotspots between chromosomes following Prdm9 humanization of the infertile (PWD × B6)(F1) F1(PWD/B6)ChIP library, we fitted a quasi-Poisson model to these heats, including predictors for each chromosome. Specifically, we estimated parameters α, β and γ by fitting the model \[ \log(\pi_\text{hotspots}) = \alpha + \gamma \log(\text{decrease}) + \sum_{i=1}^{14} \beta_i \chi_i, \] where \( \text{decrease} \) and \( \chi_i \) are the DMC1 heats of a particular hotspot which is shared between the infertile (PWD × B6)(F1) F1(PWD/B6)ChIP library and rescue (PWD × B6)(F1) F1(PWD/B6)ChIP library and c is a categorical variable which represents the chromosome on which the DSB hotspot occurs. Furthermore, for one of the hybrid mice we considered, for a given autosome, we defined the ‘total H3K4me3 signal from PRDM9 binding on both homologues (that is, symmetrically) at the same hotspots, summed over the entire chromosome’, also referred to as ‘the sum of symmetric heat’, as \( \sum_i 4r_i(1-r_i)h_i \), where \( r_i \) is the fraction of DMC1 reads coming from the B6 chromosome for hotspot \( i \), \( h_i \) is the H3K4me3 heat of that hotspot, and the sum is taken over all the hotspots on that chromosome which are under the control of a specific (PWD, B6, or humanized) PRDM9. (Our analyses always refer to this sum of symmetric heats for a specific allele.) When we considered the B6 mouse (which of course has two B6 chromosomes), we defined this sum of symmetric heats to be \( \sum_i h_i^2 \) (which is the special case of the formula above with \( r_i = \frac{1}{2} \), corresponding to all hotspots being fully symmetric). Under the assumptions we describe in the Supplementary Information, this can also be interpreted as being proportional to the expected number of hotspots with PRDM9 bound on both homologues. Details and motivations for defining this quantity are given in Supplementary Information Section 8, together with a slight adjustment we used in practice to provide robustness against outliers in the value of \( h_i^2 \).

We proceeded similarly in the B6ChIP × B6ChIP comparison. The observed effects reported in Fig. 2d–f and Extended Data Fig. 7 are normalized to the effect for Chromosome 1. Precise definitions for the model, and for the 14 chromosome effect predictors tested, are given in Supplementary Information.

Analysis code availability and source data. Analysis code used for analysis in this study is available at https://github.com/anjali-hinch/hybrid-rescue. The source data generated in this publication has been deposited in NCBI’s Gene Expression Omnibus (accession number GSE73833).

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Extended Data Figure 1 | Humanization of the zinc-finger domain of Prdm9. a, Top, the targeting vector used for the humanization of the zinc-finger array encoded by a portion of exon 10. Middle, wild-type Prdm9 allele. Bottom, the targeted humanized allele, following the action of Flp recombinase which removes the FRT-flanked neomycin selection cassette. The positions of primers used for the exon spanning RT–PCR are shown along with the sizes of the predicted amplification products from cDNA.

b, RT–PCR analysis using the exon spanning primers shown in a from testis cDNA prepared from wild-type (B6B6/B6) and heterozygous humanized (B6B6/H) mice. For gel source data, see Supplementary Fig. 1. c, Relative expression of the Prdm9 transcript from testis cDNA prepared from wild-type (B6B6/B6), heterozygous (B6B6/H) and humanized (B6H/H) testis cDNA, normalized to Hprt (n = 2 for each genotype). Error bars, ±1 s.e.
Extended Data Figure 2 | Effects of the humanization of the Prdm9 zinc-finger domain on fertility parameters. a, The average litter size is shown for all combinations of genotype matings. Error bars, ± 1 s.e.

b, Numbers of DMC1 foci co-localizing with SYCP3 immunoreactivity per cell, grouped according to meiotic stage (wild-type (B6/B6), n = 5 mice; heterozygous (B6/B6/H), n = 7 mice; homozygous (B6/H/H), n = 5 mice; cell numbers counted: zygotene: 32, 38, 37; zygotene/pachytene: 55, 96, 90; pachytene: 188, 210, 176; signals on XY in pachytene: 188, 210, 175 for B6/B6/B6, B6/B6/H and B6/H/H, respectively). Mean values are shown.

c, Number of MLH1 foci per cell in pachytene stage meiotic spreads. (B6 B6/B6: n = 6 mice, 180 cells; B6 B6/H: n = 6 mice, 185 cells; B6/H/H: n = 6 mice, 183 cells). Mean values are shown.

d, Comparison of fertility metrics in four mice with homozygous genetic background (B6 or PWD). Across all four mice, there is no statistically significant evidence of differences in these fertility parameters (ANOVA, Bonferroni-corrected P values > 0.08). Error bars, ± 1 s.e.

e, Average litter sizes in F1 crosses. Error bars, ± 1 s.e.
Extended Data Figure 3 | Further features revealed by DMC1 signal analysis in mice with homozygous genetic background. a, Effect of humanization of the Prdm9 zinc-finger domain on DSB hotspots. A total of 16,225 and 17,517 DSB hotspots were localized in the homozygous humanized and wild-type mice, respectively. Only 2.6% of these hotspots overlap. b, Correlations between DSB hotspot maps at different scales. Autosomes are divided into bins of given length, and correlations between the sums of the heats of the hotspots falling into each bin are reported, for different bin sizes. The grey region indicates the empirical 95% confidence envelope for the correlation under the null hypothesis of no association between the B6H/H and B6H/H DSB maps. DSB maps for B6H/H, 13R, 9R and Prdm9 knockout (B6–/–) mice come from ref. 21. B6H/H and 9R have the same Prdm9 allele, but different genomic backgrounds. c, Breakdown of hotspot provenance (defined by overlap) in the heterozygous humanized mouse for all DSB hotspots (left panel) and for the hottest 20% of hotspots (right panel). d, Distributions of hotspot provenance in the heterozygous humanized mouse as a function of the estimated hotspot heats (blue, wild-type B6 mouse; red, humanized homozygous mouse; green, humanized heterozygous mouse; purple, undetermined). The human allele dominates over the mouse allele in terms of heat, as the proportion of DSB hotspots found in the heterozygous mouse that are shared with the homozygous humanized mouse increases with estimated heat. The relative heat/strength of a hotspot is the ratio of this hotspot’s estimated heat to the sum of all the estimated heats (on autosomes). e, Hotter hotspots present a PRDM9 binding motif more often than weaker hotspots in all samples (same colour legend).
Extended Data Figure 4 | Inferred PRDM9 binding motifs are enriched at DSB hotspot centres. a–d, Refined PRDM9 binding motifs detected in the wild-type B6 mouse (a), in the homozygous humanized mouse (b), in the heterozygous humanized mouse (c) and in wild-type PWD (d). Percentages above each motif indicate the fraction of DSB hotspots that are found to harbour this motif, with each DSB hotspot assigned at most to one motif. In logo plots, letter height in bits of information determines degree of base specificity. e–g, Enrichment of the most prevalent 15-bp wild-type (blue) and humanized homozygous (red) motifs within 100-bp bins across a 5-kb window centred on the DSB hotspot centres. Enrichments were computed for the wild-type (e), homozygous humanized (f) and heterozygous humanized (g) mice DSB hotspots.
Extended Data Figure 5 | Differential epigenetic mark distributions at PRDM9 binding motifs. a, Enrichment of H3K4me3 marks at mouse motifs that are either within a B6 (left) or human (right) PRDM9 allele controlled DSB hotspot, or outside such a hotspot. The enrichment is relative to a control genomic track. Given the spread of the distributions, the interaction range between the histones and the DSB hotspot seems to be ~1.5 kb on each side of the motif. b, As a, but for H3K36me3 marks. c, Mean coverage of H3K4me3 (left) or H3K36me3 (right) signal around the mouse motif nearest to each B6 DSB hotspot, split according to the strand on which the motif lies. d–h, As a, but for H3K9ac (d), H3K27ac (e), H3K27me3 (f), H3K4me1 (g) or H3K79me2 (h) marks. All ChIP-seq data for histone modifications used in this analysis were obtained from the Mouse ENCODE Project.
Extended Data Figure 6 | Further features of DSB hotspot asymmetry.

a–c, DSB hotspot asymmetry in (B6 × PWD)F1B6/PWD and in (B6 × PWD)F1HPWD. a, Distribution of the fraction of (DMC1) informative reads originating from the B6 chromosome in the reciprocal (B6 × PWD)F1B6/PWD mouse. PRDM9 control at each DSB is attributed either to the B6 allele (blue) or the PWD allele (pink) or is undeterminable (grey). b, c, As a, but showing fractions only for non-shared hotspots, unique to either the reciprocal (B6 × PWD)F1B6/PWD (b) or the reciprocal rescue (B6 × PWD)F1HPWD (c) mice. d, e, Comparison of the levels of asymmetric binding in the (PWD × B6)F1PWD/B6 and (B6 × CAST)F1B6/CAST mice, using H3K4me3 signal. d, Distributions of the fraction of H3K4me3 reads from the B6 chromosome in the two mice. We used raw data from ref. 17 for the (B6 × CAST)F1B6/CAST mouse, and processed both data sets in the same way. H3K4me3 heats were capped at the 95th percentile in each case, and only H3K4me3 binding peaks not inferred to be independent of PRDM9 binding (Supplementary Information Section 7), and overlapping with a DMC1 hotspot in the same mouse, were considered. e, Quantile–quantile plot for the distributions shown in d (blue). Dark grey: 95% confidence band. f, Density plot comparing, for each hotspot in the (PWD × B6)F1PWD/B6 mouse, its DMC1 and H3K4me3 asymmetries. The correlation between the two measures is 0.93. g, Mutations within 1-kb regions around B6 and PWD PRDM9 motifs, on the B6 and PWD genomes. Main plot, for each combination of motif and lineage (PWD or B6), we plot the fraction of 30-bp windows, along the 1-kb regions surrounding motif occurrences within DSB hotspots, where at least one SNP or indel mutation occurred along the respective lineage. Inset plot, distribution of motif score differences (derived-ancestral) for motif changes shown in the main plot. Motif score was defined as the logarithm of the probability that a motif was drawn from the motif’s position weight matrix, in the ancestral sequence and in the current-day mouse. A negative difference indicates the motif match worsened along the corresponding lineage. h, Mutations within 1-kb regions around B6 PRDM9 motifs, on the B6 and CAST genomes, as in g, using the (B6 × CAST)F1B6CAST DMC1 map. We see no evidence of erosion of B6 PRDM9 motifs on the CAST genome.
Extended Data Figure 7 | Chromosome effects following Prdm9 humanization in B6^{6+/6}. a, Individual chromosome effects (relative to chromosome 1) when comparing DMC1 signals in the B6^{6/6} mouse relative to the B6^{6+/6} mouse, for the DSB hotspots that are shared between these two mice. b, Comparison of the observed chromosome effects for DMC1 signals with the fitted chromosome effects, using the two-predictor model including the sum of symmetric H3K4me3 heats in B6^{6+/6} and in B6^{6+/H}. Bars conservatively show 3 standard errors in both plots.
Extended Data Figure 8 | Value by chromosome and sensitivity analysis for the symmetry metric. a, Symmetry metric, as defined in the main text, for each sample (ALL), and for each autosome amongst those samples. Error bars represent bootstrap 95% confidence intervals in all panels. b, Alternative symmetric metrics (to the ones reported in the main text), using only 10,000 hotspots per sample, or without weighting each chromosome specific metric, to compute the average metric genome-wide. Both metrics are computed using the DMC1 maps. c, Alternative symmetric metrics using H3K4me3 maps, similarly to b. The threshold of 12,540 hotspots per sample corresponds to the number of hotspots with ratio estimates in the (PWD × B6)F1 mouse, which was the lowest amongst the three samples shown here.
Extended Data Figure 9 | See next page for figure caption.
Asymmetric hotspots, hotspots on the X chromosome and hotspots opposite deletions show systematic increase of DMC1 heat, relative to symmetric hotspots. 

For the PWD allele in the humanized rescue (PWD × B6)F1™ mouse, mean DMC1 signal is plotted in decile bins of H3K4me3 enrichment on the B6 chromosome (or the PWD X chromosome), with error bars showing 95% confidence intervals and lines of best fit (as in Fig. 4c). The slope of the line for asymmetric hotspots is 2.5-fold greater than that of the symmetric hotspots, and the slope for hotspots on the X chromosome is 5.2-fold greater, illustrating that the DMC1 signal at asymmetric sites is elevated in a similar fashion to hotspots on the X chromosome, which do not repair until late in meiosis. We found similar results in all cases tested.

Comparison of DMC1 heats on B6 chromosome for hotspots shared between the humanized B6/H and the humanized rescue (PWD × B6) F1™ mouse, under humanized PRDM9 control. We show symmetric hotspots (fraction of DMC1 informative reads between 0.4 and 0.6, green), and hotspots opposite a deletion on the PWD chromosome (deletion of at least 200 bp, encompassing a human PRDM9 binding motif, black). The black line represents the median DMC1 heat for symmetric hotspots. Hotspots opposite PWD deletion show a significant elevation in DMC1 heat relative to symmetric hotspots (14 of 16 hotspots above the symmetric median line, \( P = 0.004 \)).

Bar plot showing the genome-wide autosomal ratio of mean DMC1 heat to mean H3K4me3 enrichment for asymmetric hotspots relative to symmetric hotspots in 9 scenarios studied, each for a different combination of mouse, Prdm9 allele, and haplotype, with error bars representing 95% bootstrap confidence intervals for the ratio of means. In all cases, asymmetric hotspots show an elevation in DMC1 signal for a given H3K4me3 signal. 

Comparison of DMC1 heats on B6 chromosome relative to the PWD chromosome compared with the H3K4me3 signal on the PWD chromosome (log scale) in the infertile mouse. Asymmetric hotspots were defined as those with H3K4me3 fraction on the B6 chromosome > 0.9, and symmetric hotspots were those with the fraction between 0.1 and 0.9. Hotspots that we estimated to be completely asymmetric (H3K4me3 fraction = 0 on either chromosome) or those with H3K4me3 enrichment on either chromosome close to zero (enrichment < 0.05) were excluded to avoid singularities on either axis. Asymmetric hotspots were binned into four bins of equal size and symmetric hotspots were binned into ten bins of equal size. Different numbers of bins were used for asymmetric and symmetric hotspots to get approximately similar confidence intervals (error bars represent 95% confidence intervals) to enable comparison. We did not observe many weak symmetric hotspots as we have limited power to detect such hotspots, which is why there are no symmetric bins with very low H3K4me3 levels on the homologue (right). Right, as left, but with the ratio determined for the PWD chromosome relative to H3K4me3 on the B6 chromosome. Accordingly, asymmetric hotspots are defined as those with H3K4me3 fraction on the PWD chromosome > 0.9.
Extended Data Figure 10 | Elevation of DMC1 asymmetric heat is not explained by GC content, local heterozygosity, differences in binding-motif-disrupting mutations or by outliers. a–f, Comparison of DMC1 signals in the infertile (PWD × B6)F1PWD/B6 mouse, at symmetric and asymmetric hotspots respectively, binned by H3K4me3 enrichment, after matching symmetric and asymmetric hotspots on various features: a, DMC1 heat in B6B6/B6; b, local heterozygosity outside the PRDM9 binding motif, in a 500-bp window; c, as b, but for a 1-kb window; d, number of SNPs in binding motif; e, number of indels in binding motif; f, local GC content, computed in a 200-bp window around hotspot centre.

g, h, i, Distributions of the ratios of H3K4me3 heats on the B6 chromosome, in the rescue (PWD × B6)F1PWD/B6 versus humanized B6H/H mice, for the symmetric (fraction of informative DMC1 reads in the range 0.4–0.6, green) and asymmetric (fraction 0.9–1, orange) PRDM9 control shared between the two mice. The distributions are very close, suggesting similar trimethylation by PRDM9 on the B6 chromosome in both mice. h, As g, but for the DMC1 heats. Despite similar trimethylation marking by PRDM9 in both mice, we observed notable changes in the distribution of DMC1 ratios. This could be due to either more breaks occurring at the asymmetric sites, or a longer time taken to repair them. i, Quantile–quantile plots of DMC1 heats for hotspots under the control of the humanized allele on the B6 chromosome in the rescue (PWD × B6)F1PWD/B6 (y axis, left) versus the humanized B6H/H (x axis) mice, for symmetric (green) and asymmetric (orange) hotspots. Dotted line represents the ratios of asymmetric to symmetric quantiles (excluding distribution tails; y axis, right). Dashed line represents expected ratio if there were no differences between symmetric and asymmetric hotspots. The observed ratio of DMC1 quantiles is constant across DMC1 heats, emphasizing that the increase in DMC1 heat at asymmetric sites is very similar across the whole range of DMC1 heats, and does not simply result from a few outlying hotspots.