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1	Ensuring meiotic DNA break formation in the mouse pseudoautosomal region			
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20				
21	Sex chromosomes in males of most eutherian species share only a diminutive homologous			
22	segment, the pseudoautosomal region (PAR), wherein double-strand break (DSB)			
23	formation, pairing, and crossing over must occur for correct meiotic segregation <sup>1,2</sup> . How			
24	cells ensure PAR recombination is unknown. Here we delineate an unexpected dynamic			
25	ultrastructure of the PAR and identify controlling cis- and trans-acting factors that make			
26	this the hottest area of DSB formation in the male mouse genome. Before break formation,			
27	multiple DSB-promoting factors hyper-accumulate in the PAR, its chromosome axes			
28	elongate, and the sister chromatids separate. These phenomena are linked to			
29	heterochromatic mo-2 minisatellite arrays and require ME14 and ANKRD31 proteins but			
30	not axis components REC8 or HORMADI. We propose that the repetitive PAR sequence			
31	confers unique chromatin and higher order structures crucial for recombination.			
32	Chromosome synapsis triggers collapse of the elongated PAR structure and, remarkably,			
33 24	deleving on proventing synapsis. Thus, sevuelly dimembia behavior of the DAD rests in			
34 25	nert on kinetic differences between the seves for a rese between meturation of DAD			
25 26	part on Kneue unterences between the sexes for a race between maturation of PAK structure DSR formation and completion of pairing and synapsis. Our findings establish a			
30 27	machanistic paradiam of say chromosome recombination during majosis			
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39	During meiotic recombination, DSBs must occur within the tiny ( $\sim$ 700 kb <sup>3,4</sup> ) mouse			

39	During metotic recombination, DSBs must occur within the tiny (~700 kb <sup>-+</sup> ) mouse
40	PAR <sup>2-6</sup> . Since on average one DSB forms per ten megabases, the PAR would risk frequent
41	recombination failure if it behaved like a typical autosomal segment <sup>2</sup> . Consequently, the PAR

has disproportionately frequent DSBs and recombination<sup>2,6-8</sup> (Supplementary Discussion).
 Mechanisms promoting such frequent DSBs are unknown in any species.

DSBs arise concomitantly with linear axial structures that anchor chromatin loops wherein DSBs occur<sup>9,10</sup>. Axes begin to form during replication and become assembly sites for proteins that promote SPO11 DSBs<sup>11-13</sup>. PAR chromatin in spermatocytes forms relatively short loops on a long axis<sup>2</sup>. However, only a low-resolution view of PAR structure was available and the controlling cis- and trans-acting factors were unknown. Moreover, it was unclear how spermatocytes but not oocytes make the PAR so hyperrecombinogenic.

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# A distinctive PAR ultrastructure

X and Y usually pair late, with PARs paired in less than 20% of spermatocytes at late
 zygonema when most autosomes are paired<sup>2,14</sup>. At this stage, unsynapsed PAR axes (SYCP2/3)
 appeared thickened relative to other unsynapsed axes and had bright HORMAD1/2 staining (Fig.
 **1a and Extended Data Fig. 1a,b**)<sup>15</sup>. Moreover, the PAR was highly enriched for REC114,
 MEI4, MEI1, and IHO1—essential for genome-wide DSB formation<sup>16-19</sup>—plus ANKRD31, a
 REC114 partner essential for PAR DSBs<sup>20,21</sup>.

All five proteins (RMMAI) colocalized in several bright "blobs" for most of prophase I (Fig. 1a and Extended Data Fig. 1c). Two blobs were on X and Y PARs and others highlighted specific autosome ends (Fig. 1a, Extended Data Fig. 1d), revisited below. Similar blobs in published micrographs were uncharacterized<sup>16,17,19,22</sup>. The proteins also colocalized in smaller foci along unsynapsed axes<sup>16,17,19-22</sup> (Extended Data Fig. 1c). Enrichment on the PAR was already detectable in pre-leptonema (Extended Data Fig. 1e)<sup>17,22</sup> but not in spermatogonia (Extended Data Fig. 1f). Mass spectrometry of testis immunoprecipitates identified ZMYM3 and PTIP as new ANKRD31 interactors also enriched on the PAR (Extended Data Fig. 1g-i).

Structured illumination microscopy (SIM) resolved the thickened PAR as two axial cores 66 (Fig. 1b and Extended Data Fig. 2a,b) decorated with RMMAI (Fig. 1c). PAR axes were 67 extended and separated in late zygonema before X and Y synapsis, then collapsed during X-Y 68 synapsis in early pachynema (Fig. 1b). Each axial core is a sister chromatid, with a "bubble" 69 from near the PAR boundary almost to the telomere (Extended Data Fig. 2c-h). This PAR 70 structure is distinct from what is seen at chromosome ends later in prophase I (Supplementary 71 Discussion). Axis splitting and REC114 enrichment occurred independently of DSB formation 72 (Extended Data Fig. 2i). 73

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# Dynamic remodeling of PAR structure

We investigated temporal patterns of axis differentiation, RMMAI composition, and 76 77 chromatin loop configuration on the PAR using SIM or conventional microscopy (Fig. 1d and Extended Data Fig. 3a,b). The SYCP3-defined axis was already long as soon as it was 78 79 detectable in leptonema, and the PARb FISH signal was compact and remained so while the axis lengthened further through late zygonema, when the sister axes separated. Throughout, abundant 80 ANKRD31 and REC114 signals stretched along the PAR axes, decorating the compact 81 chromatin (Extended Data Fig. 3a chromosomes a-h, and Extended Data Fig. 3b i-ii). After 82 83 synapsis, the axes shortened and chromatin loops decompacted, with concomitant RMMAI dissociation. A focus of the meiotic cohesin subunit REC8 was juxtaposed to ANKRD31 blobs 84

at pre-leptonema; REC8 was mostly restricted to the borders of the PAR as its axes elongated 85 and split, and remained highly enriched on the short axis after RMMAI proteins disappeared 86 (Extended Data Fig. 3a chromosomes i-o, and Extended Data Fig. 3b iii-iv). Collapse of the 87 88 loop-axis structure and REC114 dissociation also occurred when the PAR underwent nonhomologous synapsis in a  $Spo11^{-/-}$  mutant (Extended Data Fig. 3c), so synapsis without 89 recombination is sufficient for PAR reconfiguration. DSB formation without synapsis may also 90 be sufficient (Supplementary Discussion). These findings delineate large-scale reconfiguration of 91 loop-axis structure and establish spatial and temporal correlations between RMMAI proteins and 92 association of a long axis with compact PAR chromatin. 93

94

### 95 Heterochromatic mo-2 minisatellites

We deduced that specific DNA sequences might recruit RMMAI proteins because 96 autosomal blobs also hybridized to the PARb probe (Extended Data Fig. 1d). This repetitive 97 probe includes a ~20-kb tandem array of a minisatellite called mo-2, with a 31-bp repeat<sup>23,24</sup> 98 (Fig. 2a). Clusters of mo-2 are also present at the non-centromeric ends of chr4, chr9, and chr13 99 (Fig. 2a,b and Extended Data Fig. 4a,b)<sup>23,24</sup>. FISH with an mo-2 oligonucleotide probe showed 100 that RMMAI blobs colocalize completely with mo-2 arrays (Fig. 2b and Extended Data Fig. 101 **4c.d**). Mo-2 arrays become enriched at the onset of meiosis for heterochromatic histone 102 modifications (H3K9me3, H4K20me3) and proteins (HP1 $\beta$ , HP1 $\gamma$ , and others), independent of 103 DSB formation (Extended Data Fig. 5). 104

#### To test if mo-2 arrays are cis-acting determinants of RMMAI recruitment, we exploited the fact that the *Mus musculus molossinus* subspecies has substantially lower mo-2 copy number<sup>24</sup>. The MSM/MsJ strain (MSM) showed less hybridization signal than B6 with the mo-2 FISH probe and had lower REC114 intensity in blobs (**Extended Data Fig. 4e**).

To avoid confounding strain effects, we examined spermatocytes of F1 hybrids (Fig. 2c 109 and Extended Data Fig. 4f.g). Less ANKRD31 accumulated on MSM PARs: the Y<sup>MSM</sup> PAR 110 had 8-fold less ANKRD31 than the  $X^{B6}$  PAR in offspring from B6 mothers and MSM fathers 111 (Fig. 2c and Extended Data Fig. 4g), and the  $X^{MSM}$  PAR had 6.5-fold less than the  $Y^{B6}$  PAR in 112 the reciprocal cross (Extended Data Fig. 4f.g). Relative ANKRD31 levels matched mo-2 FISH. 113 Nevertheless, MSM PARs support sex chromosome pairing efficiency and timing similar to B6 114 (Extended Data Fig. 4h), not surprisingly since MSM is fertile. Interestingly, the ssDNA 115 binding protein RPA2 was present at lower intensity on MSM PARs (Fig. 2c and Extended 116 Data Fig. 4f), revisited below. 117

118

# 119 **Trans-acting determinants**

To identify factors important for PAR behavior, we eliminated RMMAI or axis 120 proteins<sup>16,20,25,26</sup>. Requirements for RMMAI blobs overlap with but are distinct from those for 121 smaller RMMAI foci, for which Hormad1 is important and Mei4 even more so, but Ankrd31 122 contributes only partially<sup>17,20,22</sup> (Fig. 3a). HORMAD1 and REC8 were dispensable for RMMAI 123 assembly on mo-2 regions, PAR axis elongation, splitting of sister axes, and formation of short 124 loops (i.e., compact mo-2 and REC114 signals) (Fig. 3a,b,c and Extended Data Fig. 6a,b). 125 Distal PAR axes were separated in  $Rec8^{-1}$  (Fig. 3c and Extended Data Fig. 6c), so REC8 is 126 essential for cohesion at the PAR end. 127

The smaller MEI4 and REC114 foci still formed in *Ankrd31<sup>-/-</sup>*, but fewer and weaker 128 (Fig. 3a and Extended Data Fig. 6a,d,e)<sup>20</sup>. On mo-2 in contrast, RMMAI proteins did not 129 accumulate detectably in *Mei4<sup>-/-</sup>* and *Ankrd31<sup>-/-</sup>* (Fig. 3a and Extended Data Fig. 6a,b). 130 ANKRD31 was dispensable for enrichment of heterochromatin factors (Extended Data Fig. 6f). 131 REC114, although not IHO1, is similarly essential for RMMAI blobs<sup>21</sup>. Normal PAR 132 ultrastructure was also absent in Mei4<sup>-/-</sup> and Ankrd31<sup>-/-</sup>: axes were short with no sign of splitting 133 and mo-2 was decompacted (Fig. 3b,c and Extended Data Fig. 6b). We conclude that PAR 134 RMMAI blobs share genetic requirements with autosomal mo-2 blobs, and presence of blobs 135 correlates with normal PAR structural differentiation. 136

137

#### 138 PAR(-like) axis remodeling and mo-2

If mo-2 arrays are cis-acting determinants of high-level RMMAI recruitment that in turn 139 governs PAR structural dynamics, then autosomal mo-2 should also form PAR-like structures. 140 Indeed, the distal end of chr9 underwent splitting in spermatocytes where this region was late to 141 synapse (Fig. 4a) and showed a PAR-like pattern of extended axes and compact chromatin 142 dependent on Ankrd31 (Extended Data Fig. 7a). Thus, mo-2 (and/or linked elements) may be 143 sufficient for both RMMAI recruitment and axis remodeling. Less axis remodeling for MSM 144 PARs (Extended Data Fig. 7b) reinforced the correlation between mo-2 copy number, RMMAI 145 levels, and PAR ultrastructure. 146

147

#### 148 **DSB formation in spermatocytes**

We hypothesized that RMMAI recruitment and axis remodeling create an environment conducive to high-level DSB formation. This idea predicts that mutations should affect all of these processes coordinately and that autosomal mo-2 regions should experience PAR-like DSB formation. We counted axial RPA2 foci as a proxy for global DSB numbers and assessed mo-2 overlap with RPA2 (**Fig. 4b and Extended Data Fig. 7c-f**).

In wild-type zygotene spermatocytes, RPA2 foci overlapped on average 35% of each 154 cell's mo-2 regions, increasing to 70% at pachynema (Extended Data Fig. 7e). Similar to the 155 PAR<sup>2</sup>, autosomal mo-2 often acquired DSBs late (Extended Data Fig. 7g). In contrast, Ankrd31<sup>-</sup> 156 <sup>-</sup> mutants had starkly reduced overlap of RPA2 foci with mo-2, so X and Y paired in only 6% of 157 mid-pachytene spermatocytes (Fig. 4b and Extended Data Fig. 7e,h). This is distinct from 158 autosomes: global RPA2 foci were only modestly reduced (Extended Data Fig. 7d) and most 159 Ankrd31<sup>-/-</sup> cells pair and synapse all autosomes<sup>20,21</sup>. (Ankrd31<sup>-/-</sup> mutants form fewer RPA2 foci 160 at leptonema and early zygonema, but normal numbers thereafter<sup>20,21</sup>.) 161

*Rec8* deficiency did not reduce RPA2 focus formation on mo-2 or more globally relative 162 to a synapsis-deficient control ( $Syce1^{-/-}$ ) (Extended Data Fig. 7c-e). However, X–Y pairing was 163 reduced (Extended Data Fig. 7h), presumably because REC8 promotes interhomolog 164 recombination<sup>27</sup>. *Hormad1*<sup>-/-</sup> spermatocytes had comparable or higher frequencies of mo-2-165 overlapping RPA2 foci and X–Y pairing as the  $Syce 1^{-/-}$  control (Extended Data Fig. 7e,h). The 166 high frequency of mo-2 RPA2 foci was striking given the global reduction in RPA2 foci 167 (Extended Data Fig. 7d.f) and DSBs<sup>28</sup>, but consistent with HORMAD1 dispensability both for 168 RMMAI recruitment to mo-2 and for PAR ultrastructure (Fig. 3a-c). 169

These findings establish a tight correlation of RMMAI recruitment and axis remodeling with high-frequency DSB formation. Further strengthening this correlation, we noted above that MSM PARs display lower RPA2 intensity (**Fig. 2c**), perhaps reflecting a lesser tendency to make multiple DSBs. Indeed, multiple PAR RPA2 foci were resolved by SIM more frequently in B6 than MSM (**Extended Data Fig. 7i,j**).

We used maps of ssDNA bound by the strand-exchange protein DMC1 (ssDNA 175 sequencing, or SSDS)<sup>7,29,30</sup> to test more directly whether autosomal mo-2 regions experience 176 PAR-like DSB formation, i.e., dependent on ANKRD31 but largely independent of the histone 177 methyltransferase PRDM9 (Fig. 4c and Extended Data Fig. 8a)<sup>7,20,21</sup>. Indeed, the region 178 encompassing the chr9 mo-2 cluster displayed accumulation of SSDS reads that was 179 substantially reduced in Ankrd31<sup>-/-</sup> but not in Prdm9<sup>-/-</sup>. A modest ANKRD31-dependent, 180 PRDM9-independent peak was also observed near the mo-2 cluster on chr13 (Extended Data 181 Fig. 8a). Thus, autosomal mo-2 regions not only accumulate PAR-like levels of RMMAI 182 proteins and undergo PAR-like axis remodeling in spermatocytes, they frequently form DSBs in 183 a PAR-like manner. 184

185

### 186 **Mo-2 regions in oocytes**

In females, recombination between the two X chromosomes is not restricted to the PAR, 187 so oocytes do not require PAR DSBs like spermatocytes<sup>31</sup>. We therefore asked whether the PAR 188 undergoes spermatocyte-like structural changes in oocytes. RMMAI proteins robustly 189 accumulated on PAR and autosomal mo-2 regions from leptonema to pachynema (Extended 190 Data Fig. 9a), consistent with studies of MEI4 and ANKRD31<sup>16,21</sup>. Oocytes also displayed an 191 extended PAR axis and compact PARb FISH signal from leptonema to zygonema and 192 transitioned to a shorter axis and more extended PARb signal in pachynema, with loss of 193 REC114 signal upon synapsis (Extended Data Fig. 9b). Heterochromatin factors were also 194 enriched (Extended Data Fig. 9c). However, we did not detect spermatocyte-like thickening or 195 196 splitting of the PAR axis or REC8 accumulation (Extended Data Fig. 9d), even in the absence of synapsis in  $Syce1^{-/-}$  mutants (Extended Data Fig. 9e). Moreover, similar to the PAR<sup>31</sup>, 197 autosomal mo-2 regions showed little enrichment for SSDS signal in wild-type ovaries 198 199 (Extended Data Fig. 8b,c).

Low SSDS signal despite RMMAI enrichment and long axes could indicate that oocytes lack a critical factor(s) that promotes PAR DSBs in spermatocytes. Alternatively, oocyte PARs may not realize their full DSB potential because of negative feedback tied to homolog engagement<sup>32,33</sup>: perhaps synapsis that initiated elsewhere on X often spreads into the PAR and disrupts the PAR ultrastructure before DSBs can form. To test this idea, we tested effects of delaying or blocking PAR synapsis using sex-reversed XY females<sup>34</sup> and *Syce1<sup>-/-</sup>* mutants.

206 XY oocytes pair and synapse their PARs relatively late: only 28% of late zygotene cells 207 had X and Y paired and/or synapsed (25 of 90 cells from two mice), increasing to 66% at 208 pachynema (115 of 174 cells). This late pairing and synapsis is reminiscent of spermatocytes, but 209 appears less efficient. Most pachytene XY oocytes that synapsed their PARs had a PAR-210 associated RPA2 focus, at twice the frequency and with higher immunofluorescence intensity 211 than in XX oocytes (**Fig. 4d, Extended Data Fig. 9f**). RPA2 foci were also seen on most PARs 212 that failed to synapse (**Extended Data Fig. 9g**). In contrast, chr9 and chr13 had lower RPA2 frequency and intensity that was comparable to XX PARs and that did not differ between XY
and XX (Extended Data Fig. 9f).

These findings suggest that delayed PAR synapsis allows oocytes to more efficiently form DSBs. Supporting this conclusion, absence of synapsis in  $Syce I^{-/-}$  oocytes was accompanied by an increase in both the frequency and intensity of RPA2 on PARs and autosomal mo-2 regions alike (**Extended Data Fig. 9h**). Our results do not exclude the possibility of spermatocyte-oocyte differences in trans-acting factors, but we infer that the ability to manifest high-level DSB formation depends substantially on the result of a race between DSB formation and completion of synapsis (Supplementary Discussion).

#### 223 Discussion

222

224 We demonstrate that the PAR in male mice undergoes a striking rearrangement of loop-225 axis structure prior to DSB formation involving recruitment of RMMAI proteins, dynamic axis elongation, and splitting of sister chromatid axes (Extended Data Fig. 10). Most of these 226 behaviors also occur in oocytes and can support high-level DSB formation if synapsis is delayed. 227 228 The mo-2 array may be a key cis-acting determinant and RMMAI proteins are crucial transacting determinants. Although the function of sister axis splitting is unclear (Supplementary 229 Discussion), the full suite of PAR behaviors appears essential for pairing, recombination, and 230 segregation of heteromorphic sex chromosomes. 231

Budding yeast also uses robust recruitment of Rec114 and Mer2 (the IHO1 ortholog) to ensure that its smallest chromosomes incur DSBs<sup>35</sup>. Thus, such preferential recruitment is an evolutionarily recurrent strategy for mitigating risk of recombination failure when the length of chromosomal homology is limited.

RMMAI hyper-accumulation may reflect binding of one or more of these proteins to an
 mo-2-associated chromatin structure and/or direct binding to mo-2 repeats or another tightly
 linked DNA element. We note that the repetitive mo-2 array imposes risks of unequal
 exchange<sup>23,36</sup>. Thus, paradoxically, the PAR DNA structure stabilizes the genome by supporting
 sex chromosome segregation but also promotes the rapid evolution of mammalian PARs<sup>4</sup>.

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#### 242 **References:**

- 2431Raudsepp, T. & Chowdhary, B. P. The eutherian pseudoautosomal region. Cytogenet244Genome Res 147, 81-94 (2015).
- 245 2 Kauppi, L. *et al.* Distinct properties of the XY pseudoautosomal region crucial for male
   246 meiosis. *Science* 331, 916-920 (2011).
- 247 3 Perry, J., Palmer, S., Gabriel, A. & Ashworth, A. A short pseudoautosomal region in
  248 laboratory mice. *Genome Res* 11, 1826-1832 (2001).
- 2494Raudsepp, T., Das, P. J., Avila, F. & Chowdhary, B. P. The pseudoautosomal region and250sex chromosome aneuploidies in domestic species. Sex Dev 6, 72-83 (2012).
- 2515Palmer, S., Perry, J., Kipling, D. & Ashworth, A. A gene spans the pseudoautosomal252boundary in mice. *Proc Natl Acad Sci U S A* 94, 12030-12035 (1997).
- 2536Soriano, P. et al. High rate of recombination and double crossovers in the mouse254pseudoautosomal region during male meiosis. Proc Natl Acad Sci U S A 84, 7218-7220255(1987).
- Prick, K., Smagulova, F., Khil, P., Camerini-Otero, R. D. & Petukhova, G. V. Genetic
   recombination is directed away from functional genomic elements in mice. *Nature* 485, 642-645 (2012).
- Lange, J. *et al*. The landscape of mouse meiotic double-strand break formation, processing, and repair. *Cell* **167**, 695-708 e616 (2016).
- 2619Kleckner, N. Chiasma formation: chromatin/axis interplay and the role(s) of the262synaptonemal complex. *Chromosoma* **115**, 175-194 (2006).
- 263 10 Zickler, D. & Kleckner, N. Meiotic chromosomes: integrating structure and function.
   264 Annu Rev Genet 33, 603-754 (1999).
- 26511Panizza, S. *et al.* Spo11-accessory proteins link double-strand break sites to the<br/>chromosome axis in early meiotic recombination. *Cell* **146**, 372-383 (2011).
- Lam, I. & Keeney, S. Mechanism and regulation of meiotic recombination initiation.
   *Cold Spring Harb Perspect Biol* 7, a016634 (2014).
- 26913de Massy, B. Initiation of meiotic recombination: how and where? Conservation and270specificities among eukaryotes. Annu Rev Genet 47, 563-599 (2013).
- 27114Kauppi, L., Jasin, M. & Keeney, S. The tricky path to recombining X and Y272chromosomes in meiosis. Ann N Y Acad Sci 1267, 18-23 (2012).
- 27315Page, J. *et al.* Inactivation or non-reactivation: what accounts better for the silence of sex274chromosomes during mammalian male meiosis? *Chromosoma* **121**, 307-326 (2012).
- Kumar, R., Bourbon, H. M. & de Massy, B. Functional conservation of Mei4 for meiotic
  DNA double-strand break formation from yeasts to mice. *Genes Dev* 24, 1266-1280
  (2010).
- 27817Stanzione, M. *et al.* Meiotic DNA break formation requires the unsynapsed chromosome279axis-binding protein IHO1 (CCDC36) in mice. Nat Cell Biol 18, 1208-1220 (2016).
- 18 Reinholdt, L. G. & Schimenti, J. C. *Meil* is epistatic to *Dmc1* during mouse meiosis. *Chromosoma* 114, 127-134 (2005).
- 28219Kumar, R. *et al.* Mouse REC114 is essential for meiotic DNA double-strand break283formation and forms a complex with MEI4. *Life Sci Alliance* 1, e201800259 (2018).
- 284 20 Boekhout, M. *et al.* REC114 partner ANKRD31 controls number, timing, and location of 285 meiotic DNA breaks. *Mol Cell* **74**, 1053-1068 e1058 (2019).

286 287	21	Papanikos, F. <i>et al.</i> Mouse ANKRD31 regulates spatiotemporal patterning of meiotic recombination initiation and ensures recombination between X and Y sex chromosomes
288		<i>Mol Cell</i> <b>74</b> , 1069-1085 e1011 (2019).
289	22	Kumar, R. <i>et al.</i> MEI4 - a central player in the regulation of meiotic DNA double-strand
290		break formation in the mouse. J Cell Sci <b>128</b> , 1800-1811 (2015).
291	23	Harbers, K., Francke, U., Soriano, P., Jaenisch, R. & Muller, U. Structure and
292		chromosomal mapping of a highly polymorphic repetitive DNA sequence from the
293		pseudoautosomal region of the mouse sex chromosomes. Cytogenet Cell Genet 53, 129-
294		133 (1990).
295	24	Takahashi, Y. et al. Methylation imprinting was observed of mouse mo-2 macrosatellite
296		on the pseudoautosomal region but not on chromosome 9. Chromosoma 103, 450-458
297		(1994).
298	25	Shin, Y. H. et al. Hormad1 mutation disrupts synaptonemal complex formation,
299		recombination, and chromosome segregation in mammalian meiosis. PLoS Genet 6,
300		e1001190 (2010).
301	26	Bannister, L. A., Reinholdt, L. G., Munroe, R. J. & Schimenti, J. C. Positional cloning
302		and characterization of mouse <i>mei8</i> , a disrupted allelle of the meiotic cohesin Rec8.
303		Genesis 40, 184-194 (2004).
304	27	Kim, K. P. et al. Sister cohesion and structural axis components mediate homolog bias of
305		meiotic recombination. Cell 143, 924-937 (2010).
306	28	Daniel, K. et al. Meiotic homologue alignment and its quality surveillance are controlled
307		by mouse HORMAD1. Nat Cell Biol 13, 599-610 (2011).
308	29	Brick, K., Pratto, F., Sun, C. Y., Camerini-Otero, R. D. & Petukhova, G. Analysis of
309		meiotic double-strand break initiation in mammals. Methods Enzymol 601, 391-418
310		(2018).
311	30	Khil, P. P., Smagulova, F., Brick, K. M., Camerini-Otero, R. D. & Petukhova, G. V.
312		Sensitive mapping of recombination hotspots using sequencing-based detection of
313		ssDNA. Genome Res (2012).
314	31	Brick, K. et al. Extensive sex differences at the initiation of genetic recombination.
315		<i>Nature</i> <b>561</b> , 338-342 (2018).
316	32	Thacker, D., Mohibullah, N., Zhu, X. & Keeney, S. Homologue engagement controls
317		meiotic DNA break number and distribution. <i>Nature</i> <b>510</b> , 241-246 (2014).
318	33	Kauppi, L. et al. Numerical constraints and feedback control of double-strand breaks in
319		mouse meiosis. <i>Genes Dev</i> 27, 873-886 (2013).
320	34	Arnold, A. P. Mouse models for evaluating sex chromosome effects that cause sex
321		differences in non-gonadal tissues. J Neuroendocrinol 21, 377-386 (2009).
322	35	Murakamı, H., Lam, I., Song, J., van Overbeek, M. & Keeney, S. Multilayered
323		mechanisms ensure that short chromosomes recombine in meiosis. <i>bioRxiv</i> doi:
324	•	<u>https://doi.org/10.1101/406892</u> (2018).
325	36	Harbers, K., Soriano, P., Muller, U. & Jaenisch, R. High frequency of unequal
326		recombination in pseudoautosomal region shown by proviral insertion in transgenic
327		mouse. <i>Nature</i> <b>324</b> , 682-685 (1986).
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Fig. 1: Ultrastructure of the PAR during male meiosis. (a) Axis thickening (SYCP2 and 330 SYCP3) and ANKRD31 accumulation on X and Y PARs (arrowheads) in late zygonema. The 331 asterisk shows an autosomal ANKRD31 blob. Scale bar: 2 µm. (b) Ultrastructure of the PAR 332 333 before and after synapsis (montage of representative SIM images). Dashed lines indicate where chromosomes are cropped. SIM: Structured Illumination Microscopy. Scale bar: 1 µm. (c) 334 RMMAI enrichment along split PAR axes in late zygonema. Scale bar: 1 µm. (d) Schematic 335 showing the dynamic remodeling of the PAR loop-axis ensemble during prophase I. See 336 measurements in Extended Data Fig. 3b and Data File S1. Scale bar: 1 µm. 337

Fig. 2: Arrays of the mo-2 minisatellite are sites of RMMAI protein enrichment in the PAR 338 and on autosomes. (a) Left panel: Self alignment of the PARb FISH probe. The circled block is 339 a 20-kb mo-2 cluster. Right panel: Schematic showing the non-centromeric chromosome ends 340 identified by BLAST search using the mo-2 consensus sequence. (b) Colocalization of REC114 341 blobs with mo-2 oligonucleotide FISH signal (zygotene spermatocyte). Scale bar: 2 µm. (c) PAR 342 enrichment for ANKRD31 and RPA2 correlates with mo-2 copy number. Top panels: late 343 zygotene spermatocyte from F1 hybrid from crosses of B6  $\times$  MSM. Scale bars: 1  $\mu$ m. Bottom 344 panels: PAR-associated signals (A.U., arbitrary units) on B6-derived (X<sup>B</sup>) and MSM-derived 345 chromosomes  $(Y^M)$  from the indicated number of spermatocytes (N). Red lines: means  $\pm$  SD. 346 Differences between X and Y PAR intensities are significant for both proteins and for mo-2 347 FISH ( $p < 10^{-6}$ , paired t-test; exact two-sided p values are in **Data File S2**). 348

Fig. 3: Requirements for RMMAI recruitment and PAR axis remodeling. (a) Quantification 349 350 of REC114, ANKRD31, MEI4, and IHO1 foci along unsynapsed axes in leptotene/early zygotene spermatocytes. Error bars: means  $\pm$  SD. Comparisons to wild type are indicated (two-351 sided Student's t test): \* = p < 0.02, \*\* =  $p \le 10^{-7}$ , ns = not significant (p > 0.05); exact p values are 352 in Data File S3. Representative micrographs of REC114 staining are shown; other proteins are in 353 Extended Data Fig. 6a. Presence of mo-2 associated blobs (arrowheads) is indicated in the 354 bottom panel. Scale bars:  $2 \mu m$ . (b) Genetic requirements for PAR loop-axis organization 355 (length of REC114 and mo-2 FISH signals along the PAR axis and axis-orthogonal extension of 356 mo-2). Error bars: means  $\pm$  SD. (c) Representative SIM images of Y-PAR loop-axis structure in 357 each mutant at late zygonema. Scale bar: 1 µm. 358

Fig. 4: PAR-like structural reorganization and DSB formation on autosomal mo-2 arrays. 359 (a) The mo-2 region of chr9 undergoes axis elongation and splitting similar to PARs (SIM image 360 of a wild-type zygotene spermatocyte). Scale bar: 1  $\mu$ m. (b) ANKRD31 is required for high-level 361 DSB formation in mo-2 regions and XY pairing. Immuno-FISH for RPA2 and mo-2 was used to 362 detect DSBs. Illustration from Extended Data Fig. 7c. (c) PAR-like DSB formation near 363 autosomal mo-2 regions. Excerpt from Extended Data Fig. 8a. SSDS coverage<sup>6,19</sup> is shown for 364 the Y PAR (left) and the mo-2-adjacent region of chr9 (right). Positions of mo-2 repeats are 365 shown below. (d) Early pachytene XY oocyte showing bright RPA2 focus in the PAR. Scale bar: 366 367 2 µm.

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### 369 **METHODS**

#### 370 **Mice**

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Mice were maintained and sacrificed under U.S.A. regulatory standards and experiments 371 were approved by the Memorial Sloan Kettering Cancer Center (MSKCC) Institutional Animal 372 Care and Use Committee (IACUC, protocol number 01-03-007). Animals were fed regular 373 rodent chow with ad libitum access to food and water. The Ankrd31 knockout allele 374  $(Ankrd31^{em1Sky})$  is a single base insertion mutation (+A) in exon 3; its generation and phenotypic 375 characterization are described elsewhere<sup>20</sup>. Mice with the *Mei4* knockout allele<sup>16</sup> were kindly 376 provided by B. de Massy (IGH, Montpellier, France). All other mouse strains were purchased 377 from the Jackson Laboratory: C57BL/6J (stock #00664), MSM/MsJ (stock #003719), B6N(Cg)-378 *Syce1*<sup>tm1b(KOMP)Wtsi</sup>/2J (stock #026719), B6;129S7-*Hormad1*<sup>tm1Rajk</sup>/Mmjax (stock #41469-JAX), B6;129S4-*Rec8*<sup>mei8</sup>/JcsMmjax (stock #34762-JAX), B6.Cg-Tg(Sry)2Ei Sry<sup>d11Rlb</sup>/ArnoJ (stock 379 380 #010905). Mice were genotyped using Direct Tail lysis buffer (Viagen) following the 381 manufacturer's instructions. 382

B6.Cg-Tg(Sry)2Ei *Sry<sup>dl1Rlb</sup>*/ArnoJ males have a Y chromosome with a deletion of the sexdetermining *Sry* gene and also have an *Sry* transgene integrated on an autosome. When these males are crossed with C57BL/6J females, those XY and XX animals that do not inherit the *Sry* transgene develop as females.

#### 388 Generation of REC8 and REC114 antibodies

To produce antibodies against REC8, a fragment of the mouse Rec8 gene encoding amino 389 acids 36 to 253 (NCBI Reference Sequence: NP 001347318.1) was cloned into pGEX-4T-2 390 vector. The resulting fusion of the REC8 fragment fused to glutathione S transferase (GST) was 391 expressed in E. coli, affinity purified on glutathione Sepharose 4B, and cleaved with Precision 392 393 protease. Antibodies were raised in rabbits by Covance Inc. (Princeton NJ) against the purified recombinant REC8 fragment, and antibodies were affinity purified using GST-REC836-253 that 394 had been immobilized on glutathione sepharose by crosslinking with dimethyl pimelimidate; 395 bound antibodies were eluted with 0.1 M glycine, pH 2.5. Purified antibodies were tested in 396 western blots of testis extracts and specificity was validated by immunostaining of spread 397 meiotic chromosomes from wild type and  $Rec8^{-/-}$  mice. 398

399 To produce antibodies against REC114, a fragment of the mouse *Rec114* gene encoding a truncated polypeptide lacking the N-terminal 110 amino acids (NCBI Reference Sequence: 400 NP 082874.1) was cloned into pET-19b expression vector. The resulting hexahistidine-tagged 401 REC114<sub>111-259</sub> fragment was insoluble when expressed in *E. coli*, so the recombinant protein was 402 solubilized and affinity purified on Ni-NTA resin in the presence of 8 M urea. Eluted protein was 403 dialyzed against 100 mM NaH<sub>2</sub>PO<sub>4</sub>, 10 mM Tris-HCl, 6 M urea, pH 7.3 and used to immunize 404 rabbits (Covance Inc.). Antibodies were affinity purified against purified recombinant His<sub>6</sub>-405 REC114<sub>111-259</sub> protein immobilized on cyanogen bromide-activated sepharose and eluted in 0.2 406 M glycine pH 2.5. The affinity purified antibodies were previously used by Stanzione et al.<sup>17</sup> 407 who reported detection of a band of appropriate molecular weight in western blots of testis 408 extracts. However, subsequent analysis showed that this band is also present in extracts of 409 *Rec114<sup>-/-</sup>* testes, and thus is non-specific (C. Brun and B. de Massy, personal communication). 410 Importantly, however, Stanzione et al. also reported detection of immunostaining foci on spread 411 meiotic chromosomes similar to findings reported here and by Boekhout et al.<sup>20</sup>. This 412 immunostaining signal is absent from chromosome spreads prepared from Rec114<sup>-/-</sup> mutant mice 413

414 (C. Brun and B. de Massy, personal communication). Moreover, this immunostaining signal is 415 indistinguishable from that reported using independently generated and validated anti-REC114 416 antibodies<sup>19</sup>. We conclude that our anti-REC114 antibodies are highly specific for the cognate 417 antigen when used for immunostaining of meiotic chromosome spreads.

# 419 Chromosome spreads

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Testes were dissected and deposited after removal of the tunica albuginea in  $1 \times PBS pH$ 420 7.4. Seminiferous tubules were minced using forceps to form a cell suspension. The cell 421 suspension was filtered through a 70-um cell strainer into a 15 ml Falcon tube pre-coated with 422 3% (w/v) BSA, and was centrifuged at 1000 rpm for 5 min. The cell pellet was resuspended in 423 12 ml of 1× PBS for an additional centrifugation step at 1000 rpm for 5 min and the pellet was 424 resuspended in 1 ml of hypotonic buffer containing 17 mM sodium citrate, 50 mM sucrose, 30 425 mM Tris-HCl pH 8, 5 mM EDTA pH 8, 0.5 mM dithiothreitol (DTT), 10 µl of 100× Halt 426 protease inhibitor cocktail (Thermo Scientific), and incubated for 8 min. Next, 9 ml of 1× PBS 427 was added and the cell suspension was centrifuged at 1000 rpm for 5 min. The cell pellet was 428 429 resuspended in 100 mM sucrose pH 8 to obtain a slightly turbid cell suspension, and incubated for 10 min. Superfrost glass slides were divided into two squares using an ImmEdge 430 hydrophobic pen (Vector Labs), then 110 µl of 1% paraformaldehyde (PFA) (freshly dissolved 431 in presence of NaOH at 65°C, 0.15% Triton, pH 9.3, cleared through 0.22 um filter) and 30 ul of 432 cell suspension was added per square, swirled three times for homogenization, and the slides 433 434 were placed horizontally in a closed humid chamber for 2 h. The humid chamber was opened for 1 h to allow almost complete drying of the cell suspension. Slides were washed in a Coplin jar 2 435  $\times$  5 min in 1 $\times$  PBS on a shaker, and 2 min with 0.4% Photo-Flo 200 solution (Kodak), air dried 436 and stored in aluminum foil at  $-80^{\circ}$ C. 437

438 Ovaries were extracted from 14.5–18.5 d post-coitum mice, and collected in 1× PBS pH 439 7.4. After 15 min incubation in hypotonic buffer, the ovaries were placed on a slide containing 440 30  $\mu$ l of 100 mM sucrose pH 8, and dissected with forceps to form a cell suspension. The 441 remaining tissues were removed, 110  $\mu$ l of 1% paraformaldehyde-0.15% Triton was added, and 442 the slides were gently swirled for homogenization, before incubation in a humid chamber as 443 described above for spermatocyte chromosome spreads.

# 445 Immunostaining

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446 Slides of meiotic chromosome spreads were blocked for 30 min at room temperature horizontally in a humid chamber with an excess of blocking buffer containing  $1 \times PBS$ , pH 7.4 with 0.05% 447 Tween-20, 7.5% (v/v) donkey serum, 0.5 mM EDTA, pH 8.0, and 0.05% (w/v) sodium azide, 448 and cleared by centrifugation at 13,000 rpm for 15 min. Slides were incubated with primary 449 antibody overnight in a humid chamber at 4°C, or for at least 3 hours at room temperature. Slides 450 were washed  $3 \times 5$  min in  $1 \times PBS$ , 0.05% Tween-20, then blocked for 10 min, and incubated 451 with secondary antibody for 1–2 hours at 37°C in a humid chamber. Slides were washed  $3 \times 5$ 452 min in the dark on a shaker with 1× PBS, 0.05% Tween-20, rinsed in H<sub>2</sub>O, and mounted before 453 air drying with Vectashield (Vector Labs). Antibody dilutions were centrifuged at 13,000 rpm 454 for at least 5 min before use. Primary antibodies used were rabbit and guinea pig anti-455 ANKRD31<sup>20</sup> (1:200 dilution), rabbit anti-HORMAD2 (Santa Cruz, sc-82192, 1:50), guinea pig 456 anti-HORMAD2 (1:200) and guinea pig anti-IHO1 (1:200) (gifts from A. Toth (Technical 457

University of Dresden)), goat anti-MEI1 (Santa Cruz, sc-86732, 1:50), rabbit anti-MEI4 (gift 458 from B. de Massy, 1:200), rabbit anti-REC8 (this study, 1:100), rabbit anti-REC114 (this study, 459 1:200), rabbit anti-RPA2 (Santa Cruz, sc-28709, 1:50), goat anti-SYCP1 (Santa Cruz, sc-20837, 460 1:50), rabbit anti-SYCP2 (Atlas Antibodies, HPA062401, 1:100), mouse anti-SYCP3 (Santa 461 Cruz, sc-74569, 1:100), goat anti-SYCP3 (Santa Cruz, sc-20845, 1:50), rabbit anti-TRF1 (Alpha 462 Diagnostic, TRF12-S, 1:100), rabbit anti-H4K20me3 (Abcam, ab9053, 1:200), rabbit anti-463 H3K9me3 (Abcam, ab8898, 1:200), mouse anti-macroH2A1.2 (Active motif, 61428, 1:100), 464 mouse anti-HP-1 gamma (Millipore, MAB3450, 1:100), mouse anti-HP1-beta (Millipore, 465 MAB3448, 1:100), rabbit anti-HP1-beta (Genetex, GTX106418, 1:100), rabbit anti-Mi2 466 (recognizes CHD3 and CHD4; Santa Cruz, sc-11378, 1:50), rabbit anti-ATRX (Santa Cruz, sc-467 15408, 1:50), mouse anti-DMRT1 (Santa Cruz, sc-377167, 1:50), rabbit anti-ZMYM3 (Abcam, 468 ab19165, 1:300), rabbit anti-PAXIP1 (EMD Millipore, ABE1877, 1:300). Secondary antibodies 469 used were CF405S anti-guinea pig (Biotium, 20356), CF405S anti-rabbit (Biotium, 20420), 470 CF405S anti-mouse (Biotium, 20080), Alexa Fluor488 donkey anti-mouse (Life technologies, 471 A21202), Alexa Fluor488 donkey anti-rabbit (Life technologies, A21206), Alexa Fluor488 472 donkey anti-goat (Life technologies, A11055), Alexa Fluor488 donkey anti-guinea pig (Life 473 technologies, A11073), Alexa Fluor568 donkey anti-mouse (Life technologies, A10037), Alexa 474 Fluor568 donkey anti-rabbit (Life technologies, A10042), Alexa Fluor568 goat anti-guinea pig 475 (Life technologies, A11075), Alexa Fluor594 donkey anti-mouse (Life technologies, A21203), 476 Alexa Fluor594 donkey anti-rabbit (Life technologies, A21207), Alexa Fluor594 donkey anti-477 goat (Life technologies, A11058), Alexa Fluor647 donkey anti-rabbit (Abcam, ab150067), Alexa 478 Fluor647 donkey anti-goat (Abcam, ab150131), all at 1:250 dilution. 479

#### 480 481

#### ImmunoFISH and DNA probe preparation

All steps were performed in the dark to prevent loss of fluorescence from prior 482 immunostaining. After the last washing step in the immunostaining protocol, slides were placed 483 horizontally in a humid chamber and the chromosome spreads were re-fixed with an excess of 484 2% (w/v) paraformaldehyde in 1× PBS (pH 9.3) for 10 min at room temperature. Slides were 485 rinsed once in H<sub>2</sub>O, washed for 4 min in  $1 \times PBS$ , sequentially dehydrated with 70% (v/v) ethanol 486 for 4 min, 90% ethanol for 4 min, 100% ethanol for 5 min, and air dried vertically for 5-10 min. 487 Next, 15 µl of hybridization mix was applied containing the DNA probe(s) in 70% (v/v) 488 deionized formamide (Amresco), 10% (w/v) dextran sulfate, 2× SSC buffer (saline sodium 489 citrate), 1× Denhardt's buffer, 10 mM EDTA pH 8 and 10 mM Tris-HCl pH 7.4. Cover glasses 490 (22 x 22 mm) were applied and sealed with rubber cement (Weldwood contact cement), then the 491 slides were denatured on a heat block for 7 min at 80°C, followed by overnight incubation (>14 492 h) at 37°C. Cover glasses were carefully removed using a razor blade, slides were rinsed in  $0.1 \times$ 493 SSC buffer, washed in 0.4× SSC, 0.3% NP-40 for 5 min, washed in PBS-0.05% Tween-20 for 3 494 min, rinsed in H<sub>2</sub>O, and mounted with Vectashield before air drying. 495

To generate FISH probes, we used the nick translation kit from Abbott Molecular following
the manufacturer's instructions and using CF dye-conjugated dUTP (Biotium), on BAC DNA
from the clones RP24-500I4 (maps to the region of the PAR boundary, PARb probe) CH25592M6 (maps to the distal PAR, PARd probe), RP23-139J18, RP24-136G21, and CH36-200G6
(centromere-distal ends of chr4, chr9, and chr13, respectively). BAC clones were obtained from
the BACPAC Resource Center (CHORI). Labeled DNA (500 ng) was precipitated during 30 min
incubation at -20°C after adding 5 µl of mouse Cot-1 DNA (Invitrogen), 0.5 volume of 7.5 M

ammonium acetate and 2.5 volumes of cold 100% ethanol. After washing with 70% ethanol and air drying in the dark, the pellet was dissolved in 15  $\mu$ l of hybridization buffer.

<sup>505</sup> Mo-2 oligonucleotide probes were synthesized by Integrated DNA Technologies, with 6-<sup>506</sup> FAM or TYE<sup>TM</sup> 665 fluorophores added to both 5' and 3' ends of the oligonucleotide. The DNA <sup>507</sup> sequence was designed based on the previously defined consensus sequence<sup>24</sup>, and the probe was <sup>508</sup> used at a final concentration of 10 pmol/ $\mu$ l in hybridization buffer without Cot-1 DNA. The Y-<sup>509</sup> chromosome paint probe was purchased from IDLabs and used at 1:30 dilution in hybridization <sup>510</sup> buffer without Cot-1 DNA.

### 512 EdU incorporation

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Seminiferous tubules were incubated in DMEM with 10% FCS and 10 μM EdU at 37°C for
1 h for *in vitro* labeling. EdU incorporation was detected using the Click-iT EdU Alexa Fluor
647 imaging kit (Invitrogen) according to the manufacturer's instructions.

### 517 Image acquisition

518 Images of spread spermatocytes were acquired on a Zeiss Axio Observer Z1 Marianas 519 Workstation, equipped with an ORCA-Flash 4.0 camera and DAPI, CFP, FITC, TEXAS red and 520 Cy5 filter sets, illuminated by an X-Cite 120 PC-Q light source, with either 63×/1.4 NA oil 521 immersion objective or 100×/1.4 NA oil immersion objective. Marianas Slidebook 5.0 522 (Intelligent Imaging Innovations) software was used for acquisition.

Structured illumination microscopy (3D-SIM) was performed at the Bio-Imaging Resource 523 Center in Rockefeller University using an OMX Blaze 3D-SIM super-resolution microscope 524 (Applied Precision), equipped with 405 nm, 488nm and 568 nm lasers, and 100×/1.40 NA 525 UPLSAPO oil objective (Olympus). Image stacks of several µm thickness were taken with 526 0.125 µm z-steps, and were reconstructed in Deltavision softWoRx 6.1.1 software with a 527 Wiener filter of 0.002 using wavelength specific experimentally determined OTF functions. 528 Slides were prepared and stained as described above, except that chromosomes were spread only 529 on the central portion of the slides, and the slides mounted using  $18 \times 18$  mm coverslips (Zeiss). 530

# 532 Image analysis

3D-SIM images are shown either as a z-stack using the sum slices function in Fiji/ImageJ,
or as a unique slice. The X and/or Y chromosomes were cropped, rotated and further cropped for
best display. For montage display, the X and Y chromosome images were positioned on a black
background using Adobe Illustrator 2020 (version 24.1). In the instances where the axes of the X
and Y chromosomes were cropped, the area of cropping was labeled with a light gray dotted line.
Loop/axis measurements, foci counts, and fluorescence intensity quantification were only
performed on images from conventional microscopy using the original, unmodified data.

To measure the colocalization between RMMAI proteins, we costained for SYCP3 and
 ANKRD31 along with either MEI4, REC114, or IHO1, and manually counted the number of
 ANKRD31 foci overlapping with SYCP3 and colocalizing or not with MEI4, REC114 or IHO1.
 These counts were performed in 16 spermatocytes from leptonema to early/mid zygonema.

544 To quantify the total number of RPA2, MEI4, REC114, ANKRD31, and IHO1 foci, single 545 cells were manually cropped and analyzed with semi-automated scripts in Fiji<sup>37</sup> (version 2.0.0-546 rc-69/1.52p) as described in detail elsewhere<sup>20</sup>. Briefly, images were auto-thresholded on SYCP3 staining, which was used as a mask to use 'Find Maxima' to determine the number of foci.
Images were manually inspected to determine that there were no obvious defects in determining
SYCP3 axes, that no axes from neighboring cells were counted, that no artifacts were present,
and that no foci were missed by the script.

To test for colocalization between RPA2 and mo-2 FISH signals, we manually scored the 551 percentage of mo-2 FISH signals colocalizing at least partly with RPA2. Depending on the 552 progression of synapsis during prophase I, between eight and four discrete mo-2 FISH signals 553 could be detected, corresponding to (with increasing signal intensity) the chr4, chr13, chr9, and 554 the PAR (two signals for each when unpaired, or a single signal for each after homologous 555 pairing/synapsis). Notably, the RPA2 focus was most often found in a slightly more centromere-556 proximal position compared to the bulk of mo-2 FISH signals, and therefore colocalized partly 557 with mo-2 FISH signals. In the case of the PAR, this position corresponds closely to the region 558 of the PAR boundary (PARb probe). A similar trend was observed on autosomal mo-2 clusters. 559

560 For estimates of chromatin extension, we measured the maximal axis-orthogonal distance 561 between the FISH signal and the center of the PAR axis, or the centromere-distal axis for chr9 562 stained by SYCP3. In mutant mice defective for RMMAI protein recruitment in the mo-2 563 regions, the PAR axis was defined as the nearest SYCP3 segment adjacent to the telomeric 564 SYCP3 signal.

For quantification of RPA2, ANKRD31, REC8, and mo-2 signal intensity in  $B6 \times MSM$ and  $MSM \times B6$  F1 hybrids, late zygotene spermatocytes with at least one RPA2 focus on X or Y PAR were analyzed. We used the elliptic selection tool in Fiji to define a region of interest around the largest signal in the PAR, and the same selection tool was then positioned on the other PAR axis for comparison. The fluorescence intensity was measured as the integrated density with background substraction.

#### 572 Prophase I sub-staging and identification of the PAR

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Nuclei were staged according to the dynamic behavior of the autosome and sex 573 chromosome axes during prophase I, using SYCP3 staining. Leptonema was defined as having 574 short stretches of SYCP3 but no evidence of synapsis, early/mid-zygonema as having longer 575 stretches of SYCP3 staining and some synapsis, and late zygonema as having fully assembled 576 chromosome axes and substantial (>70%) synapsis. The X and Y chromosomes generally can be 577 identified at this stage, and the PAR axis is distinguishable because it appears thicker than the 578 centromeric end, particularly near the end of zygonema when autosomes are almost fully 579 synapsed. Early pachynema was defined as complete autosomal synapsis, whereas the X and Y 580 chromosomes could display various configuration: i) unsynapsed, with thickened PAR axes, ii) 581 engaged in PAR synapsis, iii) synapsed in the PAR and non-homologously synapsed along the 582 full (or nearly full) Y chromosome axis. Mid pachynema was defined as showing bright signal 583 from autosome axes, desynapsing X and Y axes remaining synapsed only in the PAR, with short 584 PAR axis. During this stage, the autosomes and the non-PAR X and Y axes are initially short and 585 thick, and progressively become longer and thinner. Late pachynema was defined as brighter 586 autosome axes with a characteristic thickening of all autosome ends. The X and Y non-PAR axes 587 are then long and thin and show excrescence of axial elements. Diplonema was defined as 588 brighter axes and desynapsing autosome, associated with prominent thickening of the autosome 589 ends, particularly the centromeric ends. In early diplonema, the non-PAR axes of X and Y 590 chromosomes are still long and thin and progressively condense to form bright axes, associated 591 with bulges. Most experiments were conducted using SYCP3 in combination with a RMMAI 592

protein, which allows easier distinction between synapsing and desynapsing X and Ychromosomes.

595 By using only SYCP3 staining, the PARs can only be identified unambiguously from the 596 late zygonema-to-early pachynema transition through to diplonema. From pre-leptonema to 597 mid/late-zygonema, the PARs were identified as the two brightest RMMAI signals, the two 598 brightest mo-2 FISH signals, the two brightest PARb FISH signals, or the two FISH signals from 599 the PARd probe. The Y PAR could be distinguished from the X PAR using the PARb probe, as 500 this probe also weakly stains the chromatin of the non-PAR portion of the Y chromosome.

601 PAR loop/axis measurements in oocytes were performed on two 14.5–15.5 dpc (days post-602 coitum) (enriched for leptotene and zygotene oocytes) and two 18.5 dpc female fetuses (enriched 603 for pachytene oocytes).

We found significant variability in the X or Y PAR axis length between different animals in 604 our mouse colony maintained in a C57BL/6J congenic background, and even between different 605 C57BL/6J males obtained directly from the Jackson Laboratory. This is in agreement with 606 previous reports about the hypervariable nature of the mo-2 minisatellite and its involvement in 607 unequal crossing over in the mouse<sup>6,24,36,38,39</sup> (mo-2 was also named DXYmov15 or Mov15 608 flanking sequences). However, the RMMAI signal intensity/elongation and the PAR axis length 609 were always correlated with mo-2 FISH signal intensity. Importantly, despite this variability, 610 mo-2 and RMMAI proteins were enriched in the PAR and autosome ends of all mice analyzed. 611

613 Analysis of SSDS data

SSDS sequencing data were from previously described studies<sup>7,20,31</sup> and are all available at 614 the Gene Expression Omnibus (GEO) repository under accession numbers GSE35498, 615 GSE99921, GSE118913. To define enrichment values presented in Extended Data Fig. 8b, the 616 SSDS coverage was summed across the indicated coordinates adjacent to the mo-2 repeats. A 617 chromosomal mean and standard deviation for chr9 was estimated by dividing the chromosome 618 into 4-kb bins, summing the SSDS coverage in each bin, and calculating the mean and standard 619 620 deviation after excluding those bins that overlapped a DSB hotspot. The enrichment score was then defined as the difference between the coverage in the mo-2-adjacent region and the chr9 621 mean coverage, divided by the chr9 standard deviation. 622

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# 624 Immunoprecipitation/mass spectrometry

Immunoprecipitations were carried out on samples from wild type and  $Ankrd31^{-/-}$  animals 625 using two separate polyclonal anti-ANKRD31 antibodies raised in rabbit and guinea pig<sup>20</sup> (four 626 samples total). Two additional immunoprecipitations were performed using an anti-Cyclin B3 627 antibody on either wild-type or *Ccnb3* knockout testes<sup>40,41</sup>; these samples serve as additional 628 negative controls for the ANKRD31 interaction screen. For each sample, protein extracts were 629 prepared from testes of three 12-dpp mice in 1 ml of RIPA buffer (50 mM Tris-HCl, 150 mM 630 NaCl, 0.1% SDS, 0.5% Na-deoxycholate, 1% NP-40, 10 mM MgCl<sub>2</sub>, 100 units of Benzonase for 631 1h at 4°C. After centrifugation at 13,000rpm for 20 min at 4°C, the lysate was pre-cleared using 632 30µl of a slurry of protein A/G Dynabeads for 1h at 4°C. Next, 50µl of protein A/G beads 633 coupled for 30 min with 10µg of anti-ANKRD31 or anti-Cyclin B3 antibody (monoclonal 634 antibody #5 from ref. 43) were added and the solution incubated overnight at 4°C on a rotating 635 rack. Beads were washed 3 times in 1 ml of RIPA buffer and once with 1 ml of 50 mM 636 ammonium bicarbonate. Samples were then digested overnight with 2µg trypsin in 80 µl of 50 637

mM ammonium bicarbonate at 37°C on a thermo mixer (850 rpm). Peptides were desalted using 638 C18 zip tips, and then dried by vacuum centrifugation. Each sample was reconstituted in 10 µl 639 0.1% (vol/vol) formic acid and 4 µl was analyzed by microcapillary liquid chromatography with 640 tandem mass spectrometry using the NanoAcquity (Waters) with an ACQUITY UPLC BEH 641 C18 Column (Waters) configured with an ACQUITY UPLC M-Class Symmetry C18 trap 642 column (Waters) coupled to a QExactive Plus mass spectrometer (Thermo Fisher Scientific). 643 Peptides were eluted with a linear gradient of 0-35% acetonitrile (0.1% formic acid) in water 644 (0.1% formic acid) over 150 min with a flow rate of 300 nl/min. The QE Plus was operated in 645 automatic, data dependent MS/MS acquisition mode with one MS full scan (380-1800 m/z) at 646 70,000 mass resolution and up to ten concurrent MS/MS scans for the ten most intense peaks 647 selected from each survey scan. Survey scans were acquired in profile mode and MS/MS scans 648 were acquired in centroid mode at 17,500 resolution and isolation window of 1.5 amu and 649 normalized collision energy of 27. AGC was set to  $1 \times 10$  for MS1 and  $5 \times 10$  and 100 ms IT for 650 MS2. Charge exclusion of unassigned and greater than 6 enabled with dynamic exclusion of 15 651 652 s. All MS/MS samples were analyzed using MaxQuant (Max Planck Institute of Biochemistry, Martinsried, Germany; version 1.5.3.3) at default settings with a few modifications. 653

# 654655 Yeast two-hybrid assay

Mouse testis cDNAs for Ptip, Zmym3, and Ankrd31 were amplified and cloned in vectors to 656 generate fusion proteins with the Gal4 DNA-binding domain (Gal4BD) or activation domain 657 (Gal4AD). Assays were conducted according to manufacturer's instructions (Clontech). Briefly, 658 Y2HGold and Y187 (Clontech) yeast haploid strains were transformed with constructs encoding 659 Gal4BD and Gal4AD fusion proteins. After mating on YPD plates, diploid cells expressing 660 Gal4BD and Gal4AD fusion proteins were selected on double dropout medium (DDO) lacking 661 leucine and tryptophan. Protein interactions were assayed by spotting diploid cell suspensions on 662 selective medium lacking leucine, tryptophan, histidine, and adenine (quadruple dropout, QDO), 663 and QDO containing X-α-gal (5-bromo-4-chloro-3-indolyl α-D-galactopyranoside) and 664 aureobasidin A and growing for 3 days at 30°C. 665

### 667 Statistical analysis

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668 All statistical tests were performed in R (version 3.4.4)<sup>42</sup> and RStudio (Version 1.1.442). 669 Negative binomial regression was calculated using the glm.nb function from the MASS package 670 (version 7.3-49)<sup>43</sup>.

### 672 Statistics and reproducibility

The pictures shown in this article are representative images that aim to illustrate the findings in the clearest manner. Any conclusion or statement regarding the results that is not associated with explicit quantification is based on the imaging and analysis of at least 20 cells, sometimes hundreds, usually from multiple mice. Details for main figures are as follows.

Fig. 1a: The thickening of the PAR axis (using SYCP3 staining) and the elongation of the
RMMAI signal along the PAR axis have been observed in more than three different mice in
hundreds of late zygotene spermatocytes using mostly our homemade antibodies against
REC114 and ANKRD31. Other antibodies such as anti-SYCP2 and anti-HORMAD2 were used
to confirm the PAR axis thickening, and anti-MEI1, anti-MEI4 and anti-IHO1 were used to

- confirm the elongation of the REC114/ANKRD31 signal along the PAR axis, in more than 20
   spermatocytes for each antibody.
- Fig. 1b: The PAR axis splitting, the extension of the RMMAI signal and the collapse of the
   PAR structure during X-Y synapsis have been observed by SIM in more than 60 spermatocytes
   in more than 3 different mice.
- Fig. 2b: The colocalization between REC114 blobs (or RMMAI blobs in general) and mo-2
   FISH signals has been observed in all spermatocytes analyzed (N>200), from leptotene to early
   pachytene in more than three different mice.
- Fig. 3c: Axis splitting on the Y PAR has been observed by SIM in more than 100 late 690 zygotene spermatocytes and in more than 20 zygotene-like spermatocytes from Hormad1<sup>-/-</sup> 691 mice. The fork-shaped PAR structure in  $Rec8^{-7-}$  mice has been observed in more than 20 692 spermatocytes. The absence of PAR differentiation and decompaction of mo-2-containing 693 chromatin was observed in more than 30 Ankrd31<sup>-/-</sup> spermatocytes and 20 Mei4<sup>-/-</sup> 694 spermatocytes. This specific pattern has been confirmed in at least three different mice of each 695 genotype using conventional microscopy. The differentiation of the PAR axis becomes hardly 696 detectable in *Hormad1<sup>-/-</sup>* at later stage in some pachytene-like spermatocytes as cells enter 697 apoptosis, similar to Spo11<sup>-/-</sup>. 698
- Fig. 4a: The differentiation of the non-centromeric end of the chr9 was observed in 6
   spermatocytes by SIM and was observed in more than 20 late zygotene spermatocytes by
   conventional microscopy in three different mice.
- 703 Data and code availability

Image analysis scripts are available on Github: https://github.com/Boekhout/ImageJScripts\_
 SSDS data are publicly available at GEO under the accession numbers indicated above. The
 mass spectrometry proteomics data have been deposited to the ProteomeXchange Consortium
 via the PRIDE partner repository<sup>44</sup> with the dataset identifier PXD017191.

- 70937Schindelin, J. et al. Fiji: an open-source platform for biological-image analysis. Nat710Methods 9, 676-682 (2012).
- 71138Kipling, D. *et al.* Structural variation of the pseudoautosomal region between and within712inbred mouse strains. *Proc Natl Acad Sci U S A* **93**, 171-175 (1996).
- Kipling, D., Salido, E. C., Shapiro, L. J. & Cooke, H. J. High frequency de novo
  alterations in the long-range genomic structure of the mouse pseudoautosomal region. *Nat Genet* 13, 78-80 (1996).
- Karasu, M. E. & Keeney, S. Cyclin B3 is dispensable for mouse spermatogenesis.
   *Chromosoma* 128, 473-487 (2019).
- Karasu, M. E., Bouftas, N., Keeney, S. & Wassmann, K. Cyclin B3 promotes anaphase I onset in oocyte meiosis. *J Cell Biol* 218, 1265-1281 (2019).
- 72042R Core Team. R: A language and environment for statistical computing., (R Foundation721for Statistical Computing, 2018).
- Venables, W. N. & Ripley, B. D. *Modern Applied Statistics with S.* 4 edn, (Springer, 2002).
- 72444Perez-Riverol, Y. *et al.* The PRIDE database and related tools and resources in 2019:725improving support for quantification data. Nucleic Acids Res 47, D442-D450 (2019).
- 726 727

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#### 728 Main Text Statements

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Author contributions: LA designed and conducted all of the cytogenetic experiments presented 742 and analyzed the data. MEK generated Ankrd31 mutant mice and anti-ANKRD31 antibodies. 743 MB and MEK provided *Ankrd31* mutant mice and unpublished data. MEK performed the IP/MS 744 experiment and TL validated the ANKRD31 interacting proteins. KB and FP performed SSDS 745 and analyzed the data under the supervision of RDC with input from LA and SK. MvO generated 746 REC8 and REC114 antibodies. LK performed initial characterization and provided unpublished 747 data on PAR ultrastructure and cohesin enrichment. MJ and SK designed and supervised the 748 research, analyzed data, and secured funding. LA and SK wrote the manuscript with input from 749 MJ. All authors edited the manuscript. 750

- 751 **Author Information:** Reprints and permissions information is available at
- www.nature.com/reprints. Authors declare no competing interests. Correspondence and requests
   for materials should be addressed to s-keeney@ski.mskcc.org.
- 754 **Data and code availability:** Image analysis scripts are available on Github:

https://github.com/Boekhout/ImageJScripts\_SSDS data are publicly available at GEO under the
 accession numbers indicated in Methods. Underlying data for all graphs in figures are provided
 in Data Files S1, S2, S3, and S4. The mass spectrometry proteomics data have been deposited to
 the ProteomeXchange Consortium via the PRIDE partner repository<sup>44</sup> with the dataset identifier
 PXD017191. Processed mass spectrometry data are provided in Data File S5.

- 760 Supplementary Materials:
- 761 Extended Data Figures 1–10
- Data File S1: Excel file containing underlying data for Fig. 1c,d and Extended Data Fig. 1c,d,
   3b,c
- 764 Data File S2: Excel file containing underlying data for Fig. 2c and Extended Data Fig. 4f,g.
- 765 Data File S3: Excel file containing underlying data for Fig. 3a,b and Extended Data Fig. 6b,d.
- Data File S4: Excel file containing underlying data for Fig. 4 and Extended Data Fig. 7a,d,e and
- 767 9b,f,h.

- 768 Data File S5: Excel file containing results of anti-ANKRD31 immunoprecipitation/mass
- 769 spectrometry.
- Supplemental Information: PDF file containing Supplementary Discussion and Supplementary
- 771 References.

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#### 773 Extended Data Fig. 1: PAR axis thickening and accumulation of RMMAI proteins.

(a) Axis thickening (SYCP3 and HORMAD2 staining) on the PAR (arrowhead) in a late 774 775 zygotene spermatocyte. Scale bar: 2 µm. HORMAD2 staining in the PAR at late zygonema mimics SYCP3 staining in all late zygonema spermatocytes analyzed (N>20) in three mice. (b) 776 Image adapted under Creative Commons CC-BY license from ref.<sup>45</sup> showing enrichment of 777 HORMAD1 on the thick PAR axis of the Y chromosome. (c) Colocalization of ANKRD31 and 778 MEI4, REC114, IHO1, and MEI1. Representative zygotene spermatocytes are shown. 779 Arrowheads indicate densely staining blobs. Areas indicated by dashed boxes are shown at 780 higher magnification. The graphs show the total number of foci colocalized in 781 782 leptotene/zygotene spermatocytes (error bars are mean  $\pm$  SD). N.D., not determined: The low immunofluorescence signal for MEI1 did not allow us to quantify the colocalization with 783 ANKRD31, although MEI1 showed clear colocalization with ANKRD31 in the blobs and at least 784 some autosomal foci (insets). Scale bars: 2 µm. Underlying data for all graphs are in Data Files 785 S1-4. Further evidence for extensive colocalization with ANKRD31 is documented in separate 786 studies<sup>20,21</sup>. (d) PARb FISH probe colocalizes with REC114 blobs. Two blobs are on PAR, as 787 judged by chromosome morphology and bright fluorescence in situ hybridization (FISH) with a 788 PAR boundary probe (PARb) and others highlight specific autosome ends. Scale bar: 2 µm. The 789 colocalization between REC114 blobs and PARb FISH signals has been observed in all 790 791 spermatocytes analyzed (N>60), from pre-leptonema to early pachynema, in more than three mice. (e) ANKRD31, REC114, and MEI1 immunostaining starts to appear in pre-leptonema. 792 Seminiferous tubules were cultured with 5-ethynyl-3'-deoxyuridine (EdU) to label replicating 793 cells, then chromosome spreads were stained for SYCP3 and either MEI1 plus REC114 or 794 ANKRD31 plus PARb FISH. Colocalized foci appear in pre-leptonema (EdU-positive cells that 795 are weakly SYCP3-positive), as previously shown for MEI4 and IHO1<sup>17,22</sup>. Because we can 796 already detect ANKRD31 accumulation at sites of PARb-hybridization, we infer that the 797 798 stronger sites of accumulation of MEI1 and REC114 also include PARs. Scale bars: 2 µm. PARb colocalized with ANKRD31 blobs (top panel) and MEI1 with REC114 (bottom panel) in all pre-799 leptotene spermatocytes analyzed (N>20) in one mouse. (f) REC114 is not detected in the mo-2 800 regions in spermatogonia. Seminiferous tubules were cultured with EdU, and chromosome 801 spreads were stained for DMRT1 (a marker of spermatogonia<sup>46</sup>) and REC114 plus mo-2 FISH. 802 REC114 blobs colocalized with mo-2 FISH signals in the preleptotene spermatocyte (bottom) 803 804 but were not apparent in the DMRT1-positive spermatogonium (top). Both cells shown were captured in a single microscopic field. Scale bar: 2 µm. Mo-2 FISH signals do not colocalize 805 with REC114 signal in all the spermatogonia analyzed (N>20) in one mouse. (g) Candidate 806 ANKRD31 interacting proteins. To identify other PAR-associated proteins, ANKRD31 was 807 808 immunoprecipitated from extracts made from whole testes of 12-dpp-old mice using two different polyclonal antibodies. This table shows a subset of proteins that were identified by 809 mass spectrometry in immunoprecipitates from wild-type testes but not from  $Ankrd31^{-/-}$  animals, 810 and not from immunoprecipitates using an irrelevant antibody (anti-Cyclin B3). Full results are 811 in Data File S5. LFQ, label-free quantification. REC114, MEI4, and MEI1 were recovered, 812 confirming specificity. REC114 is known to interact directly with ANKRD31<sup>20</sup> and MEI4 is a 813 direct partner of REC114<sup>16,47</sup>. MEI1 colocalizes with ANKRD31 on chromatin (panel c). We 814 also identified ZMYM3 and PTIP. ZMYM3 (zinc finger, myeloproliferative, and mental 815 retardation-type 3) is a component of LSD1-containing transcription repressor complexes<sup>48</sup> and 816 has incompletely understood functions in DNA repair in somatic cells<sup>49</sup>. Mutation of Zmym3 817 results in adult male infertility from unknown causes<sup>50</sup>. However, the spermatocyte metaphase I 818

arrest in this mutant<sup>50</sup> may be consistent with presence of achiasmate chromosomes, possibly 819 including X and Y. PTIP (Pax transactivation domain interacting protein; also known as 820 PAXIP1) contains multiple BRCT (BRCA1 C-terminal) domains and regulates gene 821 transcription, class switch recombination, and DNA damage responses in somatic cells<sup>51-53</sup>. 822 Conditional knockout of *Ptip* causes spermatogenic arrest, but the function of PTIP during 823 meiosis remains unclear<sup>54</sup>. Neither ZMYM3 nor PTIP was implicated previously in sex 824 chromosome recombination. (h) Enrichment of ZMYM3 (top) and PTIP (bottom) on the PAR. 825 Sex chromosomes of representative early pachytene spermatocytes are shown. Scale bars: 1 µm. 826 ZMYM3 and PTIP were enriched in the PAR in all spermatocytes analyzed (N>20) in three 827 mice. (i) Yeast two-hybrid assays testing interaction of full-length ANKRD31 fused to the Gal4 828 activating domain (AD) with either full-length PTIP or the C-terminal 191 amino acids of 829 ZMYM3 fused to the Gal4 DNA binding domain (BD). (Full-length ZMYM3 autoactivates in 830 this assay.) DDO (double dropout) medium selects for presence of both the AD and BD vectors 831 (positive control for growth); QDO (quadruple dropout) and QXA (QDO plus X-α-gal and 832 aureobasidin A) media select for a productive two-hybrid interaction at lower and higher 833 stringency, respectively. Image is representative of two experiments using the same yeast strains. 834

### 835 Extended Data Fig. 2: PAR ultrastructure.

(a) Comparison of conventional microscopy and SIM, showing that the thickened PAR axis in 836 conventional microscopy is resolved as separated axial cores (arrowheads). Scale bars: 2 µm. 837 The thickening of the PAR axis in conventional microscopy and the splitting of the PAR axis in 838 SIM was observed in more than 60 spermatocytes at late zygonema in at least three mice. (b) 839 Ultrastructure of axis proteins SYCP2, SYCP3, and HORMAD2 in the PAR. Scale bars: 1 µm. 840 SYCP2 (left) and HORMAD2 (right) staining mimic SYCP3 staining in late zygonema by 841 conventional microscopy in all cells analyzed (N>30) in at least three mice, and by SIM (N=5, 842 one mouse) (except that HORMAD2 appears rather depleted at the telomeres compared to 843 SYCP3 and SYCP2). (c-d) Ruling out a crozier configuration. In principle, sister chromatid axes 844 could be split apart or the PAR could adopt a crozier configuration in which a single conjoined 845 axis for both sister chromatids is folded back on itself. A crozier (cartooned in c) was ruled out 846 because the telomere binding protein TRF1<sup>55</sup> decorates the tip of the PAR bubble (d) and FISH 847 signal for the PARb probe is arrayed relatively symmetrically on both axial cores (e), consistent 848 with separated sister chromatid axes (a bubble configuration). Scale bars: 1 µm. We conclude 849 that each axis is a sister chromatid, with a "bubble" from near the PAR boundary almost to the 850 telomere. The presence of TRF1 at the distal tip of the PAR was observed in all spermatocytes 851 analyzed, in one mouse (by conventional microscopy, N>20; by SIM, N=3). PARb FISH signals 852 were relatively symmetrically arranged along the split PAR axes (by conventional microscopy, 853 N>100 in at least three mice, or by SIM, N=9 in three mice). (f) Schematic of PAR ultrastructure 854 and distribution of axis and RMMAI proteins at late zygonema. (g, h) Paired PARs with 855 elongated and split axes occur in late zygonema to early pachynema. Shown are electron 856 micrographs adapted with permission from ref.<sup>56</sup> in comparison with SIM immunofluorescence 857 images of spermatocytes at early pachynema (panel g) or late zygonema (panel h; cyan 858 arrowheads indicate examples of incomplete autosomal synapsis). The spermatocytes in the 859 electron micrographs were originally considered to be in mid-to-late pachynema<sup>56</sup>. However, in 860 our SIM experiments, we can only detect this structure (paired X and Y with elongated and split 861 axes, resembling a crocodile's jaws) around the zygotene-to-pachytene transition, when RMMAI 862 proteins are still highly abundant on the PAR axes, and when most or all autosomes are 863

completely synapsed. Moreover, other published electron micrographs from mid-to-late 864 pachytene spermatocytes show diagnostic ultrastructural features that are not present in the 865 electron micrographs reproduced here, including a short PAR axis length, multi-stranded 866 stretches of axis on non-PAR portions of the X and Y chromosomes with excrescence of axial 867 elements, and a clear thickening of autosomal telomeres<sup>15,57</sup>. These observations allow us to 868 conclude definitively that the elongation and splitting of PAR axes are a hallmark of cells from 869 late zygonema into early pachynema. Scale bars in SIM images: 1 µm in panel g, 2 µm in panel 870 h. Extended and split PAR axes were observed by SIM (N>30 spermatocytes) around the 871 zygonema-pachynema transition in more than three mice. (i) REC114 enrichment and axis 872 splitting occurs in the absence of SPO11, thus neither is provoked by DSB formation. Scale bar: 873 1  $\mu$ m. PAR axis splitting and extension of the RMMAI signal were observed by SIM in Spo11<sup>-/-</sup> 874 mice in more than 20 late zygotene-like spermatocytes in more than three mice. The 875 differentiation of the PAR axis became hardly detectable at later stages in some pachytene-like 876 spermatocytes as cells entered apoptosis. 877

# Extended Data Fig. 3: Time course of the spatial organization of the PAR loop-axis ensemble.

- 880 (a) Time course of REC8 and ANKRD31 immunostaining along the PAR axis from preleptonema (preL, left) to mid pachynema (right). A montage of representative SIM images is 881 shown. Chromosomes a-e are presumptive X or Y, but could be the distal end of chr9. 882 883 Chromosomes at later stages were unambiguously identified by morphology. Chromosomes i-k show examples where the initial pairing (probably synaptic) contact between X and Y is (i) 884 centromere-proximal (that is, closer to the PAR boundary), (k) distal (closer to the telomere), or 885 (i) interstitial. Scale bar: 1 um. The preferential enrichment of REC8 at the border of the PAR 886 split axes was observed in more than 30 zygotene spermatocytes by SIM in more than three 887 mice. (b) We collected three measurements of conventional immuno-FISH images from 888 889 leptonema through mid-pachynema: length of the REC114 signal along the PAR axis; maximal distance from the PARb FISH signal to the distal end of the SYCP3-defined axis; and axis-890 orthogonal extension of FISH signal for the PARb probe (a proxy for loop sizes). Data were 891 collected on three males. Insets show examples of each type of measurement at each stage. 892 893 Horizontal black lines indicate means. Means of each measurement for each mouse at each stage are given below, along with the means across all three mice. Means are rounded to two 894 895 significant figures; the grand means were calculated using unrounded values from individual mice. The number of cells of each stage from each mouse is given (N). Modest variability in the 896 897 apparent dimensions of the Y chromosome PAR between different mice may be attributable to variation in copy number of mo-2 and other repeats because of unequal exchange during meiosis. 898 Nonetheless, highly similar changes in spatial organization over time in prophase were observed 899 in all mice examined, namely progressive elongation then shortening of axes and concomitant 900 lengthening of loops. Scale bar: 1 µm. 901
- Briefly, panels a and b show the following. At pre-leptonema, ANKRD31 blobs had a closely juxtaposed focus of the meiotic cohesin subunit REC8 (chromosome a). In leptonema and early zygonema, ANKRD31 and REC114 signals stretched along the presumptive PAR axes, with REC8 restricted to the borders (panel a, chromosomes b–e). The SYCP3-defined axis was already long as soon as it was detectable (0.73  $\mu$ m) and the PARb FISH signal was compact (0.52  $\mu$ m) (panel bi). At late zygonema, the PAR axis had lengthened still further (1.0  $\mu$ m), while the PARb signal remained compact (panel bi). The PAR split into separate axes during

- 909 this stage, each with abundant RMMAI (panel a, chromosomes f–h). The split was a REC8-poor
- 2010 zone bounded by REC8 foci (panel a, chromosomes f–h and Extended Data Fig. 2f). After
- 911 synapsis, axes shortened and chromatin loops decompacted, with concomitant RMMAI
- dissociation. As cells transitioned into early pachynema and the X and Y PARs synapsed (panel
- a, chromosomes i–m), the PAR axes began to shorten slightly (0.85  $\mu$ m) while the PARb signal expanded (0.85  $\mu$ m) (panel biii). Meanwhile, the elongated ANKRD31 signals progressively
- 914 expanded (0.85 µm) (panel bin). Meanwine, the elongated ANKKD51 signals progressively 915 decreased in intensity, collapsed along with the shortening axes, and separated from the axis
- while remaining nearby (panel a, chromosomes l-m). By mid-pachynema, PAR axes collapsed
- still further, to about half their zygotene length ( $0.50 \,\mu$ m) and the PARb chromatin expanded to
- more than twice the zygotene measurement  $(1.3 \ \mu\text{m})$ . ANKRD31 and REC114 enrichment largely disappeared, leaving behind a bright bolus of REC8 on the short remaining axis (panel a,
- 920 chromosomes n–o and panel biv).
- 921 (c) Non-homologous synapsis appears sufficient to trigger collapse of the PAR loop-axis
- structure. We measured REC114 signal length along the PAR axis and extension of mo-2
- 923 chromatin orthogonal to the axis in  $Spo11^{-/-}$  spermatocytes in which the X PAR had non-
- homologously synapsed with an autosome while the Y PAR remained unsynapsed. Within any
- given cell, the unsynapsed Y PAR maintained the characteristic late zygotene configuration
   (long axis, short loops) whereas the synapsed X PAR adopted the configuration characteristic of
- mid-pachynema (short axis, long loops). Error bars are mean  $\pm$  SD. Scale bar: 2µm. We do not exclude that DSB formation without synapsis may also be sufficient (Supplementary
- 928 exclude that DSB form929 Discussion).

# Extended Data Fig. 4: RMMAI enrichment at mo-2 minisatellite arrays in the PAR and on specific autosomes.

932 (a) Top panel: Self alignment of the PARb FISH probe (reproduced from Fig. 2a). The circled block is a 20-kb mo-2 cluster. Bottom panel: Schematic depicting the last 1.4 Mb of the non-933 934 centromeric ends of the indicated chromosomes, showing the positions of mo-2 repeats (green) adjacent to assembly gaps (mm10); mo-2 repeats were identified by BLAST search using the 935 mo-2 consensus sequence. Mo-2 repeats also appear at the distal end of chr4 in the Celera 936 assembly (Mm Celera, 2009/03/04). PARb and PARd BAC clones are indicated. (b) 937 938 Confirmation that autosomal mo-2 FISH signals match the chromosomal locations indicated by mm10 or Celera genome assemblies. FISH was performed using an oligonucleotide probe 939 940 containing the mo-2 consensus sequence in combination with BAC probes for adjacent segments of chromosomes 13, 9 and 4, as indicated. Magenta arrows point to concordant FISH signals. 941 942 The chr9 BAC probe also hybridizes to the PAR. Scale bars: 2µm. The colocalization of mo-2 and the three autosomal FISH signals was observed in two mice (N>20 spermatocytes). (c) 943 Comparison of mo-2 FISH with REC114 localization relative to the PAR boundary (PARb FISH 944 probe) and the distal PAR (PARd probe). In mid zygonema, the mo-2 FISH signal colocalizes 945 well with REC114 staining in between the PARb and PARd FISH signals. In late zygonema, mo-946 2 and REC114 are similar to one another and are elongated along the thickened SYCP3 staining 947 948 of the PAR axis. From early to mid pachynema, REC114 progressively disappears, whereas the mo-2 FISH signal becomes largely extended away from the PAR axes. Note that the relative 949 positions of the PARb and PARd probes reinforce the conclusion that the PAR does not adopt a 950 crozier configuration. Scale bar: 1 µm. The different positioning of PARb and PARd FISH 951 signals compared to mo-2 or REC114 signals was observed in more than 30 spermatocytes in at 952 least three mice. (d) Illustration of the compact organization of the PAR chromatin (mo-2 FISH 953

954 signal) compared to a whole-Y-chromosome paint probe. Scale bar: 2 um. The costaining of mo-955 2 and full chrY probe was evaluated in one mouse (N>20 spermatocytes). (e) Lower mo-2 copy number in the *M. m. molossinus* subspecies correlates with lower REC114 staining in mo-2 956 957 regions. The left panels compare MSM and B6 mice for the colocalization between REC114 immunostaining and mo-2 FISH in leptotene spermatocytes. The REC114 and SYCP3 channels 958 are shown at equivalent exposure for the two strains, whereas a longer exposure is shown for the 959 mo-2 FISH signal in the MSM spermatocyte. Note that the mo-2-associated REC114 blobs are 960 much brighter relative to the smaller dispersed REC114 foci in the B6 spermatocyte than in 961 MSM. The right panel shows representative pachytene spermatocytes to confirm the locations of 962 mo-2 clusters at autosome ends and the PAR in the MSM background. Scale bars: 2 um. The 963 lower intensity of REC114 blobs in MSM compared to B6 was observed in N>30 spermatocytes 964 in three different pairs of mice. (f) PAR enrichment for ANKRD31 and RPA2 correlates with 965 mo-2 copy number. Top panel: late zygotene spermatocytes from MSM x B6 F1 hybrid. Scale 966 bar: 1 µm. Bottom panel: PAR-associated signals (A.U., arbitrary units) on B6-derived (Y<sup>B</sup>) and 967 MSM-derived chromosomes  $(X^M)$  from the indicated number of spermatocytes (N). Red lines: 968 means  $\pm$  SD. Differences between X and Y PAR intensities are significant for both proteins and 969 for mo-2 FISH in both F1 hybrids ( $p < 10^{-13}$ , paired t-test; exact two-sided p values are in **Data** 970 File S2). (g) Representative micrographs of late zygotene spermatocytes from reciprocal F1 971 hybrid males from crosses of B6 (high mo-2 copy number) and MSM (low mo-2 copy number) 972 parents. Scale bar: 1 µm. (h) Frequency of paired X and Y at late zygonema and mid pachynema 973 analyzed in three MSM and three B6 males. Differences between strains were not statistically 974 significant at either stage (p = 0.241 for late zygonema and p = 0.136 for mid pachynema; two-975 sided Student's t test). Note also that MSM X and Y are late-pairing chromosomes, as in the B6 976 background. The similar pairing kinetics indicates that the lower intensity of RMMAI staining on 977 the MSM PAR is not attributable to earlier PAR pairing and synapsis in this strain. The number 978 979 of spermatocytes analyzed is indicated (N).

#### 980 Extended Data Fig. 5: Mo-2 regions accumulate heterochromatin factors.

(a) Costaining of ANKRD31 or mo-2 with the indicated proteins and histone marks known to 981 localize at the pericentromeric heterochromatin (mouse major satellite), in zygotene 982 983 spermatocytes (left) and pre-leptotene spermatocytes (right). Each of the heterochromatin factors shows locally enriched signal coincident with mo-2 regions (arrowheads), in addition to broader 984 985 staining of other sub-nuclear regions. Scale bars: 2 µm. The CHD3/4 antibody recognizes both proteins<sup>58</sup>. The colocalization of ANKRD31 blobs with heterochromatin blobs was observed in 986 all zygotene spermatocytes analyzed (N>20) in at least three mice for each antibody (left panel) 987 and in one mouse for pre-leptotene spermatocytes (N>10) for each antibody (right panel). (b) 988 CHD3/4, ATRX, HP1B, H4K20me3, H3K9me3 and macroH2A1.2 are not detectably enriched at 989 mo-2 regions in spermatogonia (small, DMRT1-positive cells). These factors may be present at 990 mo-2 regions in these cells, but do not appear to accumulate to elevated levels. Scale bars: 2 µm. 991 992 The absence of colocalization between mo-2 FISH signals and heterochromatin factors was 993 noted in all spermatogonia analyzed (N>30) from one mouse. (c) Heterochromatin factors can be detected in the PAR up to late pachynema. Each of the assayed proteins and histone marks 994 showed staining on the autosomal and X-specific pericentromeric heterochromatin, the sex body, 995 and euchromatin, albeit with variations between sites in the timing and level of accumulation. 996 Importantly, however, they also showed enriched staining at all mo-2 regions up to early/mid-997 pachynema, as shown for H4K20me3 (top panel). By mid-to-late pachynema, as shown for 998

999 H3K9me3 here, the signal persisted in the PAR but was usually barely detectable on chr9 or 1000 chr13 mo-2 regions. This observation indicates that, at least for the PAR, the heterochromatin factors can continue to be enriched on mo-2 chromatin after RMMAI proteins have dissociated. 1001 1002 These results substantially extend previous observations about CHD3/4 colocalizing with PAR FISH signals; H4K20me3 being localized in the PAR and other chromosome ends; and 1003 H3K9me3, HP1β and macroH2A1.2 detection in the PAR in late pachynema<sup>58-61</sup>. Scale bars: 2 1004 µm. The colocalization between Maj sat and H4K20me3 and H3K9me3 was observed in all 1005 spermatocytes analyzed (N>20) in one mouse. The colocalization between H4K20me3 and mo-2 1006 FISH signals was observed in all spermatocytes analyzed (N>60), from preleptotene to mid 1007 pachytene in more than three mice. (d) Enrichment of the heterochromatin factors is independent 1008 of SPO11. Representative images of Y chromosomes from a Spo11<sup>-/-</sup> mouse are shown. Scale 1009 bar: 1 µm. The colocalization between PAR mo-2 FISH signals and heterochromatin factors was 1010 observed in all  $Spo11^{-/-}$  spermatocytes analyzed (N>30) in more than three mice for CHD3/4 and 1011 at least one mouse each for ATRX, HP1β, HP1γ, macroH2A1.2, H3K9me3, and H4K20me3. 1012

# Extended Data Fig. 6: Genetic requirements for RMMAI assembly on chromosomes and for PAR loop-axis organization.

(a) Representative micrographs of ANKRD31, MEI4, IHO1 and MEI1 staining in wild type and 1015 the indicated mutants (quantification is in Fig. 3a). Scale bars: 2 µm. (b) Measurements of PAR 1016 loop-axis organization, as in Fig. 3b, on two additional males. Data from mouse 1 are 1017 reproduced from Fig. 3b to facilitate comparison. Means of each measurement for each mouse at 1018 each stage are given below, along with the means across all three mice. Means are rounded to 1019 two significant figures; the grand means were calculated using unrounded values from individual 1020 mice. The number of cells of each stage from each mouse is given (N). (c) REC8 is dispensable 1021 for splitting apart of PAR sister chromatid axes, but is required to maintain the connection 1022 between sisters at the distal tip of the chromosome. A representative SIM image is shown of a Y 1023 chromosome from a late zygotene  $Rec8^{-/-}$  spermatocyte. The SYCP3-labeled axes adopt an 1024 open-fork configuration. Note that the distal FISH probe (PARd) shows that there are clearly 1025 disjoined sisters whereas the PAR boundary (PARb) shows only a single compact signal 1026 comparable to wild type. The disposition of the probes and SYCP3 further rules out the crozier 1027 configuration as an explanation for split PAR axes. Scale bar: 1 µm. The Y or X PAR structure 1028 was resolved by SIM as "fork-shaped" in all spermatocytes analyzed (N>20) from three mice. 1029 (d) Quantification of REC114 and MEI4 foci in two additional pairs of wild-type and Ankrd31<sup>-/-</sup> 1030 mice. Horizontal lines indicate means. Fewer foci were observed in the Ankrd31<sup>-/-</sup> mutant (two-1031 sided Student's t tests for each comparison of mutant to wild type:  $p = 5.6 \times 10^{-6} (2^{nd} \text{ set},$ 1032 REC114);  $p = 1.1 \times 10^{-5}$  (2<sup>nd</sup> set, MEI4);  $p = 2.1 \times 10^{-6}$  (3<sup>rd</sup> set, REC114); p = 0.017 (3<sup>rd</sup>, MEI4)). 1033 (e) Reduced REC114-staining intensity of axis-associated foci in  $Ankrd31^{-/-}$  mutants. To 1034 rigorously control for slide-to-slide and within-slide variation in immunostaining, we mixed 1035 together wild-type and Ankrd31<sup>-/-</sup> testis cell suspensions before preparing chromosome spreads. 1036 1037 A representative image is shown of a region from a single microscopic field containing two wildtype zygotene spermatocytes (left) and two  $Ankrd31^{-/-}$  spermatocytes of equivalent stage (right). 1038 Note the diminished intensity of REC114 foci in the Ankrd31<sup>-/-</sup> spermatocytes. Scale bar: 2  $\mu$ m. 1039 REC114 (non-blob) foci showed lower fluorescence intensity in Ankrd31<sup>-/-</sup> compared to wild 1040 type in all pairs of spermatocytes captured in the same imaging field (N=8 pairs), from one pair 1041 1042 of mice. (f) PAR enrichment of heterochromatin-associated factors is independent of ANKRD31. Representative images of the Y chromosome at late zygonema/early pachynema showing 1043

1044 colocalization between the decompacted mo-2 chromatin and the indicated proteins. Note that 1045 both the FISH and immunofluorescence signals are localized mostly off the axis. Compare with 1046 the same signals in absence of SPO11 (**Extended Data Fig. 5d**). Scale bar: 1  $\mu$ m. Mo-2 FISH 1047 signal colocalized off the axis with the heterochromatin factors in *Ankrd31<sup>-/-</sup>* mice in all 1048 spermatocytes analyzed (N>30) in more than three mice for CHD3/4 and at least one mouse for 1049 ATRX, HP1 $\beta$ , HP1 $\gamma$ , macroH2A1.2, H3K9me3, and H4K20me3.

#### 1050 Extended Data Fig. 7: PAR-associated RPA2 foci.

1051 (a) Loop-axis organization of the mo-2 region of chr9 in late zygonema. Compare with the PAR (Fig. 3b). Scale bars: 1  $\mu$ m. Error bars: means  $\pm$  SD. (b) Low mo-2 copy number correlates with 1052 less loop-axis reorganization (SIM images of late-zygotene F1-hybrid spermatocytes). Scale 1053 bars: 1 µm. The differentiation of the B6 PAR was observed in both hybrids B6 × MSM and 1054 MSM × B6 in 3 and 4 spermatocytes, respectively by SIM (1 mouse for each) and in more than 1055 1056 20 spermatocytes by conventional microscopy in two mice of each genotype. (c,d,e) Immuno-FISH for RPA2 and mo-2 was used to detect DSBs cytologically in wild type and the indicated 1057 mutants. To analyze Rec8 and Hormad1 mutations, we compared to mutants lacking SYCE1 (a 1058 svnaptonemal complex central element component<sup>62</sup>) because  $Syce1^{-/-}$  mutants show similar 1059 meiotic progression defects without defective RMMAI recruitment. Panel c shows representative 1060 images. Scale bars: 2 µm, inset 1 µm. Panel d shows the global counts of RPA2 foci for 1061 1062 zygotene (zyg) or zygotene-like cells and for pachytene (pach) or pachytene-like cells. Panel e shows, for each cell, the fraction of mo-2 regions that had a colocalized RPA2 focus. Red lines: 1063 means  $\pm$  SD. Statistical significance is indicated in panels c and d for comparisons (two-sided 1064 Student's t tests) of wild type to Ankrd31<sup>-/-</sup> or of Syce1<sup>-/-</sup> to either  $Rec8^{-/-}$  or Hormad1<sup>-/-</sup> for 1065 matched stages. Exact p values are in Data File S4. Note that the number of discretely scorable 1066 mo-2 regions in panel e varied from cell to cell depending on pairing status. (f) Frequent DSB 1067 1068 formation at mo-2 regions in the PARs and on autosomes does not require HORMAD1. Micrograph at left shows two adjacent spermatocytes (boundary indicated by dashed line). Scale 1069 bar: 2 µm. Insets at right show higher magnification views of the numbered mo-2 regions, all of 1070 which are associated with RPA2 immunostaining of varying intensity. This picture illustrates the 1071 preferential RPA2 focus formation in mo-2 regions in a Hormad1<sup>-/-</sup> mouse; quantification is in 1072 panel e. (g) Autosomal mo-2 regions often form DSBs late. Immuno-FISH for RPA2, mo-2, and 1073 1074 PARb was used to detect DSBs cytologically in wild type from leptonema to mid-pachynema, and to distinguish the X or Y PAR from chromosomes 9 and 13. Chr4 was not assaved because 1075 1076 the mo-2 FISH signal was often barely detectable. The top panel shows the global number of RPA2 foci per cell. Black lines are means  $\pm$  SD. The bottom panel shows the percentage of 1077 1078 spermatocytes with an RPA2 focus overlapping the PAR (X, Y, or both) or overlapping chr9 or chr13. A representative image of an early pachytene spermatocyte is shown. Note that, as 1079 previously shown for the PAR<sup>2</sup>, autosomal mo-2 regions continue to accumulate RPA2 foci 1080 beyond the time when global RPA2 foci have largely or completely ceased accumulating. Scale 1081 1082 bar: 2 µm. (h) X–Y pairing status, quantified by immuno-FISH for SYCP3 and the PARd probe. (i) Montage of SIM images from a B6 male showing that multiple, distinct RPA2 foci can be 1083 detected from late zygonema to mid pachynema, suggesting that multiple PAR DSBs can be 1084 formed during one meiosis (see also ref.<sup>2</sup> for further discussion). Scale bar: 1um. The presence 1085 of multiple RPA2 foci in the PAR was observed by SIM in more than 20 spermatocytes from late 1086 1087 zygonema to mid pachynema in one mouse. (j) Percentage of spermatocytes at the zygotenepachytene transition with no (0), 1, 2 or 3 distinguishable RPA2 foci on the unsynapsed Y 1088

1089 chromosome PAR of MSM and B6 mice. The difference between the strains is statistically 1090 significant (negative binomial regression,  $p = 7.2 \times 10^{-5}$ ). N indicates the number of 1091 spermatocytes analyzed. A representative picture is shown for each genotype, with one RPA2 1092 focus on the MSM PAR and two apparent sites of RPA2 accumulation on the B6 PAR. The 1093 detection of multiple foci is consistent with reported double crossovers<sup>6</sup>. Scale bar: 1 um.

### 1094 Extended Data Fig. 8: DSB maps on the PAR and autosomal mo-2 regions.

(a) SSDS sequence coverage (data from refs.<sup>7,20</sup>) is shown for the X PAR (shown previously in 1095 different form in ref.<sup>20</sup>), the Y PAR, and the mo-2-adjacent regions of chr9 and chr13. The 1096 dashed segments indicate gaps in the mm10 genome assembly. We did not assess chr4 because 1097 available assemblies are too incomplete. (b) Regions adjacent to the mo-2 region on chr9 show 1098 1099 SSDS signal that is reproducibly elevated relative to chr9 average in wild-type testis samples but not in maps from Ankrd31<sup>-/-</sup> testes or wild-type ovaries. Two of the SSDS browser tracks are 1100 reproduced from panel a. The bar graph shows enrichment values from individual SSDS maps 1101 1102 (T1–T9 are maps from wild-type testes; O1 and O2 are from wild-type ovaries<sup>31</sup>). Enrichment values are defined as coverage across the indicated coordinates relative to mean coverage for 1103 chr9 (see Methods for details). Note that ovary sample O1 and the Ankrd31<sup>-/-</sup> adult sample are 1104 known to have poorer signal:noise ratios than the other samples<sup>20,31</sup>. For all SSDS coverage 1105 tracks, reads mapping to multiple locations are included after random assignment to one of their 1106 mapped positions. However, the same conclusions are reached about ANKRD31-dependence 1107 1108 and PRDM9-independence of signal on chr9 and chr13 if only uniquely mapped reads are used. (c) Oocytes incur substantially less DSB formation than spermatocytes near the mo-2 region on 1109 chr9. SSDS signal is from ref.<sup>31</sup> (samples T1 and O2). The X-PAR is shown for comparison 1110 (previously shown to be essentially devoid of DSBs in ovary samples<sup>31</sup>). See panel **b** for 1111 quantification. 1112

# Extended Data Fig. 9: RMMAI accumulation and low-level DSB formation on mo-2 regions in oocytes.

(a) Examples of zygotene oocytes showing the colocalization between blobs of IHO1 and 1115 REC114, MEI4 and MEI1, or ANKRD31 and mo-2 FISH signal (arrowheads). Scale bars: 2µm. 1116 RMMAI blobs colocalized with mo-2 FISH signals in all zygotene oocytes analyzed (N>30) 1117 1118 from at least three mice. (b) PAR ultrastructure in oocytes, quantified as in Extended Data Fig. **3b**. Late zygotene cells with PAR synapsis are compiled separately from other zygotene cells. 1119 Error bars: means  $\pm$  SD. Scale bar: 1  $\mu$ m. (c) Examples of zygotene oocytes showing 1120 colocalization of ANKRD31 blobs with enrichment for heterochromatin factors. Scale bars: 2 1121 μm. ANKRD31 colocalized with heterochromatin factors blobs in all zygotene oocytes analyzed 1122 (N>20) from one mouse. (d) Representative SIM image of a wild-type late zygotene oocyte 1123 1124 showing neither detectable splitting of the PAR axis nor REC8 enrichment. Scale bar: 2 µm. The absence of spermatocyte-like differentiation of the PAR axis was observed (N>30 zygotene 1125 1126 oocytes) in more than three mice. A modest degree of differentiation was observed in a minority of oocytes (5/45) analyzed by SIM, but this did not resemble the typical PAR axis splitting found 1127 in spermatocytes. (e) Prolonged asynapsis does not allow axis splitting to occur in oocytes. 1128 Because synapsis appears sufficient to trigger collapse of PAR ultrastructure in spermatocytes 1129 1130 (Extended Data Fig. 3b), we asked if preventing synapsis (i.e., in a Syce 1<sup>-/-</sup> mutant) could reveal a cryptic tendency toward axis splitting in oocytes. However, whereas axis splitting was 1131 clearly observed by SIM in Syce  $l^{-l-}$  mutant spermatocytes, PAR axes were not detectably split in 1132

oocytes. Scale bars: 2 µm for main micrograph, 1 µm for insets. Axis splitting of chr9 was 1133 observed by SIM in multiple (N>20)  $Syce1^{-/-}$  spermatocytes from three different mice. The 1134 chr13 or chr4 centromere-distal axes were also occasionally seen to be split, but we did not 1135 1136 quantify this for these chromosomes. In males, the differentiation of the PAR or the chr9 axes becomes hardly detectable at later stages in some pachytene-like spermatocytes as cells enter 1137 apoptosis, similar to Spo11<sup>-/-</sup> or Hormad1<sup>-/-</sup> mice. However, in Syce1<sup>-/-</sup> oocytes, no significant 1138 axis differentiation or splitting was observed by conventional microscopy or by SIM in multiple 1139 1140 spermatocytes (N>30) from three different mice, similar to what we observed in wild-type oocytes. (f,h) Delaying synapsis promotes PAR DSB formation in oocytes. Top panels: 1141 representative micrographs of pachytene XY (f) and  $Syce 1^{-/-}$  XX oocytes (h). Middle panels: 1142 RPA2 fluorescence intensity at the border of mo-2 FISH signals from PAR, chr9, and chr13. 1143 Bottom panels: Percentage of oocytes with RPA2 focus colocalizing with mo-2 regions on PAR, 1144 chr9, and chr13. Graphs show data only for pachytene oocytes in which PARs are synapsed (two 1145 mice of each genotype). Error bars: means  $\pm$  SD. Scale bars: 2  $\mu$ m. (g) Percentage of pachytene 1146 oocvtes with one or more RPA2 foci colocalizing with mo-2 FISH signal from PAR, chr9 and 1147 1148 chr13 in XY pachytene oocytes that had unsynapsed X and Y chromosomes. Scale bar: 2 µm, 1149 inset: 1 µm.

# Extended Data Fig. 10: Summary of PAR ultrastructure and molecular determinants of axis remodeling and DSB formation.

- 1152 Schematic representation of the meiotic Y chromosome loop/axis structure before X-Y pairing/synapsis at the transition between zygonema and pachynema. The chromosome axis 1153 comprises the meiosis-specific axial proteins SYCP2, SYCP3, HORMAD1, and HORMAD2; 1154 1155 cohesin subunits (only REC8 is represented); and the RMMAI proteins (REC114, MEI4, MEI1, ANKRD31, and IHO1). On the non-PAR portion of the Y chromosome axis (left), RMMAI 1156 protein loading and DSB formation are partly dependent on HORMAD1 and ANKRD31, and 1157 strictly dependent on MEI4, REC114<sup>19</sup>, IHO1<sup>21</sup>, and presumably MEI1<sup>18</sup>. The DNA is organized 1158 into large loops, with a low number of axis-associated RMMAI foci. By contrast, in the PAR 1159 (right), the hyper-accumulation of RMMAI proteins at mo-2 minisatellites (possibly spreading 1160 into adjacent chromatin) promotes the elongation and subsequent splitting of the PAR sister 1161 1162 chromatid axes. Short mo-2-containing chromatin loops stretch along this extended PAR axis, increasing the physical distance between the PAR boundary and the distal PAR sequences, 1163 1164 including the telomere. The degree of RMMAI protein loading, PAR axis differentiation, and DSB formation are proportional to the mo-2 FISH signal (which we interpret as reflecting mo-2 1165 1166 copy number), and depend on MEI4, ANKRD31, and presumably REC114.
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#### 1168 Data S1.(Separate file)

- 1169 Excel file containing underlying data for Fig. 1c,d and Extended Data Fig. 1c,d, 3b,c
- 11701171 Data S2.(Separate file)
- 1172 Excel file containing underlying data for Fig. 2c and Extended Data Fig. 4f,g.

# 11731174 Data S3.(Separate file)

1175 Excel file containing underlying data for Fig. 3a,b and Extended Data Fig. 6b,d.

# 11761177 Data S4.(Separate file)

1178 Excel file containing underlying data for Fig. 4 and Extended Data Fig. 7a,d,e and 9b,f,h.

# 11791180 Data S5.(Separate file)

- 1181 Excel file containing results of anti-ANKRD31 immunoprecipitation/mass spectrometry
- analysis.
- 1183
- 1184









