

1 Article

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3 **CTCF-mediated 3D chromatin predetermines the gene expression**  
4 **program in the male germline**

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21 **Abstract**

22

23 **Spermatogenesis is a unidirectional differentiation process that generates haploid sperm, but how**  
24 **the gene expression program that directs this process is established is largely unknown. Here we**  
25 **determine the high-resolution 3D chromatin architecture of male germ cells during**  
26 **spermatogenesis and show that CTCF-mediated 3D chromatin predetermines the gene expression**  
27 **program required for spermatogenesis. In undifferentiated spermatogonia, CTCF-mediated**  
28 **chromatin contacts on autosomes pre-establish meiosis-specific super-enhancers (SE). These**  
29 **meiotic SE recruit the master transcription factor A-MYB in meiotic spermatocytes, which**  
30 **strengthens their 3D contacts and instructs a burst of meiotic gene expression. We also find that at**  
31 **the mitosis-to-meiosis transition, the germline-specific Polycomb protein SCML2 resolves**  
32 **chromatin loops that are specific to mitotic spermatogonia. Moreover, SCML2 and A-MYB**  
33 **establish the unique 3D chromatin organization of sex chromosomes during meiotic sex**  
34 **chromosome inactivation. We propose that CTCF-mediated 3D chromatin organization enforces**  
35 **epigenetic priming that directs unidirectional differentiation, thereby determining the cellular**  
36 **identity of the male germline.**

## 37 Introduction

38 Eukaryotic genomes are folded into a dynamic three-dimensional (3D) architecture within the  
39 nucleus that influences gene expression<sup>1-4</sup>. The development of genome-wide chromosome conformation  
40 capture methods, especially Hi-C<sup>5</sup>, has accelerated our understanding of the 3D genome and the interplay  
41 between 3D genome organization and cell-fate decisions<sup>6</sup>. For example, disruption of 3D chromatin  
42 architecture has been shown to lead to disturbed gene expression, incomplete cell differentiation, and  
43 conversion to other cell types, at least in cell culture systems<sup>7,8</sup>. Still, there is a major gap in our  
44 understanding of how the 3D genome defines gene expression programs during development and  
45 differentiation in vivo.

46  
47 The mammalian male germline provides an ideal model to decipher the relationship between the  
48 3D genome and gene expression programs. In spermatogenesis, after sex determination, male germ cells  
49 undergo a unidirectional differentiation process that comprises the maintenance of spermatogonia stem  
50 cells, commitment to meiosis, and production of haploid sperm<sup>9</sup>. Male germ cell differentiation is defined  
51 by chromatin-based mechanisms that instruct stage-specific gene expression both on autosomes and on  
52 sex chromosomes<sup>10-12</sup>. In spermatogonia, histone modifications are preset to regulate later gene expression  
53 programs<sup>13-16</sup>. Specifically, on autosomes, dimethylation of histone H3 at lysine 4 (H3K4me2) is pre-  
54 established on meiotic super-enhancer (SE) loci that drive a genome-wide burst of transcription after the  
55 mitosis-to-meiosis transition<sup>17</sup>. The sex chromosomes, on the other hand, undergo meiotic sex  
56 chromosome inactivation (MSCI), an essential event in the male germline<sup>11,18</sup>. They form a distinct  
57 nuclear compartment called the XY body (sex body) that is physically segregated from the autosomes<sup>19</sup>.

58  
59 In this study, we elucidated how the 3D genome architecture of male germ cells is regulated to  
60 define the gene expression programs that drive spermatogenesis. Drawing on the recent analysis of basic  
61 3D chromatin features in spermatogenesis<sup>20-24</sup>, we performed high-resolution Hi-C analysis using cell  
62 types representative of major stages of spermatogenesis to decipher the detailed pictures of the 3D  
63 genome in the male germline at unprecedented resolution. To determine how the 3D genome regulates the  
64 dynamic transcriptional transition from the mitotic to meiotic stages, we also performed Hi-C analyses  
65 using mouse mutants lacking key transcriptional regulators of spermatogenesis. One of these factors is  
66 SCML2, a germ cell-specific component of Polycomb Repressive Complex 1 (PRC1) that is critical for  
67 the suppression of the mitotic program in late spermatogenesis<sup>2</sup>. The other is A-MYB (MYBL1), a master  
68 transcription factor that regulates the burst of meiotic gene expression at the pachytene stage<sup>25</sup>. These  
69 functional analyses reveal that SCML2 resolves the mitotic 3D chromatin organization, whereas A-MYB  
70 drives the establishment of meiotic 3D chromatin. Importantly, we show that the unidirectional  
71 differentiation program during spermatogenesis is predetermined by CTCF-mediated 3D chromatin

72 contacts. These results provide a molecular basis for the cellular identity of male germ cells defined by  
73 the 3D genome.

74

## 75 **Results**

### 76 **High-resolution Hi-C data sets reveal 3D chromatin reprogramming during spermatogenesis**

77 To determine high-resolution 3D chromatin structures of germ cells during the course of  
78 spermatogenesis, we performed in-depth Hi-C analysis of spermatogenic cells isolated at four  
79 representative developmental stages (Fig. 1a). Specifically, we isolated THY1<sup>+</sup> undifferentiated  
80 spermatogonia and KIT<sup>+</sup> differentiating spermatogonia from the testes of 7-day-old male C57BL/6 mice  
81 using magnetic-activated cell sorting (MACS)<sup>26</sup>. In addition, we isolated pachytene spermatocytes (PS),  
82 which are in the meiotic prophase, and round spermatids (RS), which are in the postmeiotic stage, from  
83 the testes of adult male C57BL/6 mice using BSA gradient sedimentation<sup>20</sup>. We confirmed high purity for  
84 all cell types (Extended Data Fig. 1a, b) and performed Hi-C experiments on two biological replicates for  
85 each cell type (Supplementary Table 1). These replicates showed a high correlation (Extended Data Fig.  
86 1c) and we merged them for downstream analysis, yielding ~430-670 million Hi-C contact reads for each  
87 stage and exceeding the total read depths of previous Hi-C analyses in spermatogenesis<sup>20-24</sup>. Comparing  
88 the four developmental stages, we detected a relatively high correlation between mitotic spermatogonia  
89 (THY1<sup>+</sup> and KIT<sup>+</sup>) compared to PS and RS (Extended Data Fig. 1d). This reflects the biological similarity  
90 between THY1<sup>+</sup> and KIT<sup>+</sup> spermatogonia and the presence of a dynamic transition from the mitotic  
91 stages to the meiotic and postmeiotic stages.

92

93 Next, we examined the Hi-C maps of an entire representative chromosome (chromosome 7) at  
94 each stage, including a zoom-in to a specific chromosomal region (Fig. 1b). Previous high-resolution Hi-  
95 C studies in other cellular systems revealed the presence of point interactions that represent stable  
96 chromatin-loops (dots) in addition to topologically associating domains (TADs), which manifest in  
97 triangular patterns<sup>27-29</sup>. We also detected these stable chromatin loops ("chromatin loops" hereafter) (Fig.  
98 1b, bottom, marked by circles), confirming the high resolution of our new data sets. The Hi-C interaction  
99 contact matrices show that the genomes of THY1<sup>+</sup> and KIT<sup>+</sup> cells are enriched in distal interactions (>10  
100 Mb), a typical feature of interphase nuclei. These interactions are abolished in PS, where proximal  
101 interactions (a range between 1-10 Mb) dominate (Fig. 1b). This tendency was confirmed by a contact  
102 probability  $P(s)$  analysis, which is indicative of the general polymer state of chromatin<sup>5,30</sup> (Fig. 1c). These  
103 results corroborate previous Hi-C studies<sup>21,22</sup> and confirm that the typical interphase pattern of high-order  
104 chromatin present in spermatogonia is reprogrammed when cells enter meiosis.

105

### 106 **Formation of inter-TAD chromatin loops for meiotic gene regulation**

107 Chromatin is spatially organized into TADs, which restrict interactions of cis-regulatory  
108 sequences and thereby contribute to gene regulation<sup>31-33</sup>. Accordingly, chromatin loops, which are critical  
109 for gene expression regulation<sup>1,34,35</sup>, are typically observed within TADs. Indeed, in THY1<sup>+</sup>  
110 spermatogonia, chromatin loops were largely detected within TADs (intra-TADs: Fig. 1d). By contrast, in  
111 PS, many chromatin loops were detected beyond TAD borders (inter-TADs, Fig. 1d). From mitotic  
112 spermatogonia to meiotic PS, the total number of chromatin loops decreased (Fig. 1e). Of note, the  
113 chromatin loops present in THY1<sup>+</sup> spermatogonia and PS are mostly unique, and the same holds true for  
114 the subsequent PS to RS transition (Fig. 1f). This feature presumably reflects formation and resolution of  
115 meiotic chromosome structure, organized into chromatin loop arrays along chromosome axes. Moreover,  
116 the length of chromatin loops increased from mitotic spermatogonia to meiotic PS (Fig. 1g), while  
117 average contact strengths decreased based on a pile-up analysis<sup>36</sup> (Fig. 1h). These results demonstrate that  
118 the structural changes that occur at the mitosis to meiosis transition are based on the resolution of intra-  
119 TAD chromatin loops and the de novo establishment of inter-TAD chromatin loops.

120  
121 The weakening of TADs in PS<sup>20-24</sup> indicates that this feature might drive meiotic gene regulation  
122 mediated by inter-TAD chromatin loops. To test this possibility, we investigated the relationship between  
123 TAD strength and chromatin loop formation during spermatogenesis. First, we detected TADs in each  
124 stage of spermatogenesis. The number of TADs decreased during the transition from spermatogonia to  
125 meiotic PS and recovered in postmeiotic RS (Fig. 2a). This is consistent with previous reports showing  
126 the attenuation of TADs in meiotic prophase<sup>21,22</sup>. TAD boundaries were largely shared between THY1<sup>+</sup>  
127 and KIT<sup>+</sup> spermatogonia, with 3,550 common TAD boundaries (85% of 4,192 THY1<sup>+</sup> TAD boundaries),  
128 and they progressively changed from KIT<sup>+</sup> to PS and from PS to RS (Fig. 2b). Among 4,039 TAD  
129 boundaries in KIT<sup>+</sup> spermatogonia, only 1,447 (36%) were maintained in PSs. In contrast, from PSs to  
130 RSs, 1,935 (80%) out of 2,420 PS TAD boundaries were maintained, and 1,659 TAD boundaries were  
131 newly generated in RS (Fig. 2b). To define stage-specific features of TADs, we next examined the  
132 average contact strength of TADs using a pile-up analysis of the Hi-C matrices, which visualizes average  
133 insulation strengths of the regions around TADs and their boundaries<sup>36</sup>. We found that while insulation at  
134 TAD boundaries was weakened during the transition from KIT<sup>+</sup> spermatogonia to PS (Fig. 2c), the  
135 interaction strengths between adjacent TADs increased (Fig. 2d). Thus, during the transition from mitotic  
136 spermatogonia to meiotic PS, TADs and TAD borders are reprogrammed. Interactions beyond TAD  
137 boundaries increase in meiosis, not only through the de novo formation of inter-TAD chromatin loops but  
138 also through TAD-TAD interactions.

139  
140 There are various types of chromatin loops, including CTCF-CTCF loops, enhancer-promoter  
141 loops, and Polycomb-dependent loops. Since these different types contribute to gene activation or

142 silencing<sup>28,37</sup>, it is possible to infer their functions. Therefore, we examined the modifications on anchor  
143 sites of chromatin loops and distinguished the three loop classes based on the presence of CTCF,  
144 H3K4me3 (a promoter mark)/H3K27ac (an active enhancer mark), or H3K27me3 (the Polycomb  
145 repressive complex2 (PRC2)-mediated mark). First, we performed CTCF ChIP-seq in THY1<sup>+</sup>  
146 spermatogonia and PS (Extended Data Fig. 2a). CTCF binding sites overlapped with 35.6% of anchor  
147 sites of THY1<sup>+</sup> chromatin loops, while this overlap decreased to 17.3 % in PS, suggesting the resolution  
148 of CTCF-pair loops during meiosis (Fig. 2e). Indeed, at some of the sites of CTCF-pair loops in THY1<sup>+</sup>  
149 spermatogonia, CTCF enrichment was reduced or lost in PS (Fig. 2f). This suggests that the loss of CTCF  
150 might underly the resolution of some CTCF-pair loops in meiosis. Second, we used our previous  
151 H3K4me3, H3K27ac, and H3K27me3 ChIP-seq datasets in THY1<sup>+</sup> and PSs<sup>17,38,39</sup> and defined  
152 enhancer/promoter-pair loops and Polycomb-dependent loops. The proportion of enhancer/promoter-pair  
153 loops was in the range of ~20-30 %, and the proportion of Polycomb-dependent loops was less than 10 %  
154 of total chromatin loops in THY1<sup>+</sup> and PS (Fig. 2e). Since chromatin loops in PS are mostly PS-specific  
155 (Fig. 1f), all three types of chromatin loops are largely de novo generated during meiosis. Contact  
156 strengths were comparable within each class of loops in THY1<sup>+</sup> and PS (Extended Data Fig. 2b),  
157 indicating that contact strength is not class- but stage-dependent.

158  
159 We further examined how the resolution of TADs at the mitosis-to-meiosis transition is regulated.  
160 As opposed to the resolution of CTCF-pair loops (Fig. 2e, f), there was no change in the proportion of  
161 CTCF-associated TAD boundaries among all TAD boundaries (Extended Data Fig. 2c). This suggests  
162 that distinct mechanisms operate between the resolution of TAD boundaries and chromatin loops in  
163 meiosis. Taken together, these results demonstrate that attenuation of TADs and resolution of CTCF-pair  
164 loops take place at the mitosis-to-meiosis transition to establish long inter-TAD loops during meiosis  
165 (Fig. 2g).

### 166 167 **SCML2 is required for the resolution of spermatogonia-type 3D chromatin and gene repression**

168 To further understand the mechanisms that underlie the resolution of TAD boundaries and  
169 chromatin loops at the transition from mitotic spermatogonia to meiotic spermatocytes, we focused on the  
170 germline-specific Polycomb protein SCML2. SCML2 is responsible for the suppression of genes that are  
171 highly expressed in mitotic spermatogonia after the mitosis-to-meiosis transition<sup>14</sup>. It is expressed in  
172 undifferentiated spermatogonia and forms part of PRC1, which deposits H2AK119ub<sup>14</sup> and facilitates  
173 PRC2-mediated H3K27me3 during meiosis<sup>39</sup>. H3K27me3 counteracts the active enhancer mark  
174 H3K27ac, thereby resolving spermatogonia-type enhancers<sup>17</sup>. We therefore hypothesized that SCML2 is  
175 involved in the resolution of spermatogonia-type 3D chromatin. To test this hypothesis, we performed Hi-  
176 C analysis using *Scml2* knockout (*Scml2*-KO) PSs and RSs (Extended Data Fig. 1c). *Scml2*-KO PSs and

177 RSs showed increased distal interactions compared to wild-type cells, and this pattern resembles Hi-C  
178 maps in THY1<sup>+</sup> and KIT<sup>+</sup> cells (Fig. 3a). Pearson correlation analysis also showed that *Scml2*-KO PSs  
179 and RSs are more similar to wild-type THY1<sup>+</sup> and KIT<sup>+</sup> than wild-type PSs and RSs (Extended Data Fig.  
180 1d). This suggests that the mitotic 3D chromatin organization of spermatogonia is retained in *Scml2*-KO  
181 cells during meiosis.

182  
183 To determine whether SCML2 mediates the resolution of spermatogonia-type TAD boundaries,  
184 we detected TADs in the Hi-C dataset of *Scml2*-KO PS and RS. The number of TADs increased in *Scml2*-  
185 KO PS compared to wild-type PS (Fig. 3b). Of note, TAD boundaries in the *Scml2*-KO PSs overlapped  
186 with the TAD boundaries in wild-type KIT<sup>+</sup> spermatogonia at 2,523 loci (Fig. 3c). This overlap is more  
187 abundant than that between KIT<sup>+</sup> and wild-type PS TAD boundaries (1,447 loci; Fig. 3c), suggesting that  
188 spermatogonia-type TAD boundaries are retained in *Scml2*-KO PS. Indeed, among 2,613 KIT<sup>+</sup>-specific  
189 TAD boundaries (which do not overlap with wild-type PS TAD boundaries), 1,191 loci remain in *Scml2*-  
190 KO PS TAD boundaries. Pile-up analysis further confirmed that KIT<sup>+</sup>-specific TAD boundaries remain in  
191 *Scml2*-KO PS (Fig. 3d). These results indicate that SCML2 is required for the resolution of KIT<sup>+</sup>-specific  
192 TADs in PS.

193  
194 Next, we examined the role of SCML2 in the resolution of chromatin loops. The number of  
195 chromatin loops increased in *Scml2*-KO PSs and RSs compared to wild-type PS and RS (Fig. 3e). While  
196 wild-type KIT<sup>+</sup>-specific chromatin loops did not show high contact strengths in wild-type PS and RS,  
197 they remained in *Scml2*-KO PS and RS (Fig. 3f). On the other hand, chromatin loops detected in wild-  
198 type PS and RS did not show high contact strengths in *Scml2* KO PS and RS (Fig. 3f). Comparison of  
199 chromatin loops between wild-type PSs and *Scml2*-KO PSs revealed that there were 1,358 *Scml2*-KO-  
200 specific chromatin loops, 677 of which are shared with chromatin loops present in wild-type KIT<sup>+</sup> (Fig.  
201 3g). The persistence of KIT<sup>+</sup> chromatin loops in *Scml2* KO PS and RS was confirmed with Hi-C maps  
202 (Extended Data Fig. 3a). Therefore, SCML2 is also involved in the resolution of spermatogonia-type  
203 chromatin loops.

204  
205 To determine how SCML2 resolves spermatogonia-type 3D chromatin, we next examined  
206 whether SCML2 is required for the resolution of CTCF sites by performing CTCF ChIP-seq in *Scml2*-KO  
207 PS. The Pearson correlation between *Scml2*-KO PS and wild-type THY1<sup>+</sup> spermatogonia (0.79) is higher  
208 than the Pearson correlation between wild-type PS and wild-type THY1<sup>+</sup> (0.73; Extended Data Fig. 3c),  
209 suggesting that CTCF distribution in *Scml2*-KO PS is more similar to wild-type THY1<sup>+</sup> than that in wild-  
210 type PS. In wild-type PS, CTCF enrichment at the anchor sites of the CTCF pair loops detected in THY1<sup>+</sup>

211 was reduced (Fig. 2f), but CTCF enrichment at these loci remained high in *Scml2*-KO PS (Fig. 3h). This  
212 suggests that SCML2 is involved in the resolution of at least a fraction of CTCF sites.

213

214 Since chromatin conformation is implicated in the regulation of gene expression<sup>31,33</sup>, we  
215 examined the effect of the KIT<sup>+</sup>-specific loops that persist in *Scml2*-KO PS on gene expression.  
216 Therefore, we examined the expression profile of 1,243 genes present in the anchor sites of the *Scml2*-KO  
217 PS-specific loops defined in Fig. 3g. The overall expression level of these genes decreased from wild-type  
218 KIT<sup>+</sup> to wild-type PSs, but remained high in *Scml2*-KO PS compared to wild-type PSs (Fig. 3i). We thus  
219 conclude that SCML2 is required for the resolution of spermatogonia-type 3D chromatin, thereby  
220 suppressing spermatogonia-type gene expression in meiosis (Fig. 3j). Importantly, we did not observe a  
221 significant change in contact strengths of chromatin loops at SCML2-dependent bivalent promoters  
222 marked by both active (H3K4me2/3) and repressive (H3K27me3) histone modifications (Extended Data  
223 Fig. 3d). Therefore, the function of SCML2 in resolving spermatogonia-type 3D chromatin is independent  
224 of its regulation of bivalent promoters<sup>39</sup>.

225

### 226 **A-MYB is required for the formation of meiotic-type 3D chromatin and gene activation**

227 Because meiosis-specific chromatin loops are *de novo* generated after the resolution of  
228 spermatogonia-type chromatin loops (Fig. 2g), we next sought to determine the mechanism driving  
229 meiosis-specific chromatin loops. To this end, we focused on A-MYB, a transcription factor responsible  
230 for the activation of pachytene-specific genes<sup>25</sup>. A-MYB is required to establish H3K27ac on pachytene-  
231 specific enhancers, thereby activating these enhancers<sup>17</sup>. We suspected a role of A-MYB in the formation  
232 of meiosis-specific chromatin loops because of the establishment of specific enhancer/promoter-pair  
233 loops in PS. We therefore isolated PS from *A-myb* mutant (*Mybl1<sup>repro9</sup>*) mice and performed Hi-C  
234 analysis. We found that distal interactions were increased in the *A-myb* mutant PS as shown in a Hi-C  
235 heat map (Fig. 4a) and in a contact probability analysis (Extended Data Fig. 4a), suggesting that  
236 spermatogonia-type 3D chromatin is retained in the *A-myb* mutant PS. In accordance with this notion, the  
237 number of TADs also increased (Fig. 4b); in fact, more than 70% of the TAD boundaries in the *A-Myb*  
238 mutant PS were common to those detected in KIT<sup>+</sup> (Fig. 4c: left), and KIT<sup>+</sup>-specific loops largely  
239 remained in the *A-myb* mutant PS (Fig. 4c: right). Further, KIT-specific TAD boundaries retained high  
240 contact strength in the *A-myb* mutant PS (Fig. 4d). These results demonstrate that A-MYB is required for  
241 the establishment of meiosis-type 3D chromatin, and that its loss leads to the retention of spermatogonia-  
242 type 3D chromatin.

243

244 We next examined whether A-MYB is required to establish meiosis-specific chromatin loops.  
245 Although the total number of chromatin loops increased in the *A-myb* mutant PS compared to wild-type



246 PS, only 29 % of wild-type PS-specific chromatin loops (357 out of 1,223) were detected in the *A-myb*  
247 mutant PS. Pile-up analyses show that *A-myb* mutant PS retained the contact strength of KIT<sup>+</sup> chromatin  
248 loops (Fig. 4f, left), while *A-myb* mutant PS did not show high contact strength for PS chromatin loops  
249 (Fig. 4f). To test whether A-MYB directly mediates the formation of chromatin loops in PS, we  
250 reanalyzed previous ChIP-seq data of A-MYB using whole testis<sup>25</sup>. A-MYB is enriched at the anchor sites  
251 of PS chromatin loops (Fig. 4g), and A-MYB binds to 41% of genes at anchor sites of PS (849 out of  
252 2,092; Fig. 4h). This association is statistically significant when compared to the ratio of all A-MYB  
253 binding genes to all RefSeq genes (5,929/22,661;  $P = 8.5 \times 10^{-59}$ , Hypergeometric test). These results  
254 indicate that A-MYB mediates the formation of a large part of chromatin loops in PS (Fig. 4i).

255

### 256 **A-MYB-dependent 3D chromatin is associated with the production of pachytene piRNAs.**

257 Another major function of A-MYB is the production of pachytene piRNAs<sup>40</sup>, which are involved  
258 in the maintenance of genome integrity and gene regulation in late spermatogenesis<sup>41,42</sup>. A-MYB drives  
259 the production of pachytene piRNAs in a parallel mechanism with its regulation of enhancers through the  
260 induction of H3K27ac at pachytene piRNA clusters<sup>17</sup>. The loci of pachytene piRNA clusters switch from  
261 the B compartment to the A compartment during the mitosis-to-meiosis transition<sup>22</sup>. Our Hi-C data  
262 showed that 3D chromatin contacts were specifically detected at pachytene piRNA clusters in PS and  
263 were retained in RS, and that the formation of 3D chromatin is A-MYB dependent (Extended Data Fig.  
264 4b). Thus, the A-MYB-mediated formation of 3D chromatin is associated with the production of  
265 pachytene piRNAs.

266

### 267 **Meiotic super-enhancers are poised within 3D chromatin.**

268 We next sought to determine how the global transcriptional changes that occur during  
269 spermatogenesis are regulated in the context of the 3D genome. To address this question, we focused on  
270 super-enhancers (SEs), which are long stretches of enhancers that play a central role in driving cell-type-  
271 specific gene expression and determining cellular identities<sup>43-45</sup>. In PS, A-MYB activates meiosis-specific  
272 SEs (meiotic SEs) through the establishment of H3K27ac to drive the expression of late spermatogenesis-  
273 specific genes<sup>17</sup>. Based on the specific enrichment of H3K27ac, we defined 399 meiotic SEs on  
274 autosomes, which are specific to PS. Among these, 270 (67.7%) are associated with PS chromatin loops  
275 (Extended Data Fig. 5a). A chromosome-wide track view confirms that chromatin loops (detected as  
276 stable chromatin loops in this study) are largely associated with SEs and the active genic loci that are in  
277 context with SEs (Fig. 5a). Therefore, these stable chromatin loops are associated with gene regulation  
278 and are distinct from meiotic chromatin loop arrays that are formed along the chromosome axes during  
279 meiotic prophase<sup>46</sup>.

280

281 We also defined 107 “mitotic” SE that are specific to THY1<sup>+</sup> and KIT<sup>+</sup> spermatogonia and found  
282 that mitotic SEs and meiotic SEs exhibited distinct 3D chromatin dynamics during spermatogenesis. At  
283 the mitotic SE loci, strong 3D contacts were detected in THY1<sup>+</sup> and KIT<sup>+</sup> spermatogonia, which resolved  
284 together with the resolution of mitotic SEs in PS and RS (Fig. 5b). On the other hand, at meiotic SE loci,  
285 3D chromatin contacts were detected in THY1<sup>+</sup> and KIT<sup>+</sup> spermatogonia prior to the establishment of  
286 meiotic SEs and increased upon activation of meiotic SEs in PS (Fig. 5c). Pre-establishment of 3D  
287 contacts of meiotic SEs in KIT<sup>+</sup> was also detected in a representative Hi-C map (Extended Data Fig. 5b).  
288 These 3D contacts cover relatively large regions and are distinct from chromatin loops that are detected as  
289 local point interactions. To further analyze the 3D chromatin structure around the meiotic SEs, we  
290 detected Hi-C interacting loci centered around the meiotic SEs (Extended Data Fig. 5c). A pile-up  
291 analysis shows that, consistent with the SE-SE interactions, contact strength of Hi-C interacting loci  
292 increase in PS, while modest contacts are already present in mitotic spermatogonia (Extended Data Fig.  
293 5d). These results suggest that 3D contacts at meiotic SEs are preprogrammed in spermatogonia, raising  
294 the possibility that meiotic SEs are poised for later activation through 3D chromatin.

295  
296 Because SEs determine cell type-specific gene expression programs, we next sought to determine  
297 how meiotic SEs regulate target genes via 3D chromatin. We detected 611 genes that are overlapping  
298 with the genomic region interacting with meiotic SEs. Among these genes, 26 genes are associated with  
299 the GO term spermatogenesis, and were largely upregulated in PSs and RSs (Extended Data Fig. 6a and  
300 b). In our previous study, we showed that spermatogenesis-related genes adjacent to the meiotic SEs are  
301 upregulated during late spermatogenesis<sup>17</sup>. Here, we extend this observation by demonstrating that  
302 meiotic SEs also upregulate SE-interacting genes via 3D contacts.

303  
304 Next, to investigate how meiotic SEs regulate gene expression, we examined the epigenetic states  
305 of genes adjacent to meiotic SE and SE-interacting loci. H3K4me2, which is implicated in the poised  
306 chromatin state<sup>15</sup> and associated with poised meiotic SEs<sup>17</sup>, accumulated highly at these genes in KIT<sup>+</sup>  
307 spermatogonia, but decreased in PS upon activation of meiotic SEs (Extended Data Fig. 6c, d). In PS,  
308 H3K4me3 and H3K27ac, markers for active promoters and enhancers, are enriched at these loci instead.  
309 Together, these results suggest that meiotic SE pre-establish H3K4me2-enriched 3D contacts with target  
310 genes in mitotic spermatogonia, and this epigenetic state is reprogrammed to an H3K4me3/H3K27ac-  
311 enriched state upon activation of meiotic SEs.

### 312 313 **CTCF predetermines the 3D contacts of the meiotic SEs in spermatogonia**

314 To determine how the 3D contacts of the meiotic SE are predetermined in spermatogonia, we  
315 focused on CTCF, which is involved in mediating 3D chromatin contacts via CTCF-CTCF

316 interactions<sup>47,48</sup>. To examine the relationship between CTCF and meiotic SEs, we extracted 844 CTCF-  
317 binding sites that overlapped with meiotic SEs and their interacting genomic regions in PS (Fig. 5d). We  
318 also identified 13,690 sites that did not overlapped with those genomic regions (Extended Data Fig. 7a).  
319 CTCF enrichment was largely maintained at the 844 CTCF binding sites that overlapped with meiotic  
320 SEs from THY1<sup>+</sup> spermatogonia to PS (Extended Data Fig. 7b), raising the possibility that CTCF-  
321 mediated 3D chromatin contacts persist from mitotic spermatogonia to PS. Indeed, a representative Hi-C  
322 heatmap shows that 3D chromatin contacts at the meiotic SE loci are pre-established in THY1<sup>+</sup>  
323 spermatogonia and that CTCF is highly enriched at these sites (Fig. 5e). A pile-up analysis confirmed that  
324 strong 3D contacts are maintained from THY1<sup>+</sup> spermatogonia to RS at CTCF binding sites that overlap  
325 with meiotic SEs (Fig. 5f). Of note, this feature is specific to SE loci because CTCF-CTCF chromatin  
326 loops are largely reprogrammed from THY1<sup>+</sup> spermatogonia to PS (Fig. 2e). These results demonstrate  
327 that CTCF predetermines the 3D contacts of meiotic SEs in spermatogonia, poising them for later  
328 activation.

329

### 330 **A-MYB strengthens 3D contacts of meiotic SEs on autosomes**

331 Because A-MYB establishes meiotic SEs<sup>17</sup>, we reasoned that A-MYB strengthens 3D contacts of  
332 meiotic SEs in PS. Indeed, a representative track-view shows that A-MYB binds to meiotic SE-  
333 interacting loci (Fig. 5g). Specifically, it binds to the promoter regions of 294 of the 611 genes that  
334 interact with meiotic SE-interacting loci (48.1%; Extended Data Fig. 7c). To examine the role of A-MYB  
335 in the regulation of 3D chromatin at meiotic SEs, we analyzed the Hi-C data of *A-myb* mutant PS. 3D  
336 contacts between meiotic SE and the interacting loci were attenuated in the *A-myb* mutant PS compared to  
337 wild-type PS, although modest contacts were still observed (Fig. 5h, i). Together with the CTCF analysis,  
338 we conclude that there are two regulatory mechanisms for the establishment and maintenance of meiotic  
339 SEs: CTCF predetermines the overall 3D contacts of meiotic SEs in mitotic spermatogonia, and A-MYB  
340 strengthens these 3D contacts upon activation of meiotic SEs in meiotic spermatocytes (Fig. 5j).

341

### 342 **SCML2 and A-MYB establish the unique 3D chromatin architecture of the meiotic sex** 343 **chromosomes**

344 During meiosis, sex chromosomes undergo epigenetic programming that is different from  
345 autosomes. They are subject to MSCI and form a distinct nuclear compartment called the XY body (also  
346 known as the sex body)<sup>11,18</sup> (Fig. 1a). After meiosis, the silent XY-chromosomal structure, called  
347 postmeiotic sex chromatin (PMSC), persists in haploid spermatids<sup>49</sup>. Previous Hi-C studies demonstrated  
348 that meiotic sex chromosomes do not show specific 3D chromatin features<sup>20-24</sup>, supporting the notion that  
349 3D chromatin structures of the sex chromosomes are random throughout a cell population. In our Hi-C  
350 data, we confirmed that spermatogonia-type far-cis interactions disappeared from the X chromosome in

351 PS (Fig. 6a). Although there are 26 meiotic SEs on the X chromosome in wild-type PS (Extended Data  
352 Fig. 8a), we did not detect loci that interacted with these meiotic SEs (Extended Data Fig. 8b). We did,  
353 however, detect an enrichment of short-range interactions (less than 1.5 Mb) on the X chromosome  
354 specifically in PS (Extended Data Fig. 8c-e), which might be related to fact that the X chromosome  
355 remains unsynapsed during meiotic prophase I.

356  
357 To determine the mechanisms underlying the unique 3D chromatin organization of the X  
358 chromosome, we focused on SCML2 and A-MYB. SCML2 is known to accumulate and function on  
359 meiotic sex chromosomes, independently and via a distinct mechanism compared to autosomes<sup>14</sup>. A-  
360 MYB is required to establish chromosome-wide accumulation of H3K27ac on the sex chromosome<sup>17</sup>,  
361 which facilitates the activation of sex-linked genes in postmeiotic RS<sup>38</sup>. These functions of SCML2 and  
362 A-MYB are regulated downstream of the DNA damage response pathways centered on  $\gamma$ H2AX and its  
363 binding protein MDC1, which initiate MSCI at the onset of the early pachytene stage<sup>14,17,50</sup>. In *Scml2*-KO  
364 PS, spermatogonia-type far-cis interactions remain on the X chromosome (Fig. 6b, Extended Data Fig.  
365 8f), and this feature persists in *Scml2*-KO RS (Extended Data Fig. 8h). In *A-myb* mutant PS,  
366 spermatogonia-type far-cis interactions also remain on the X chromosome and we observe a plaid pattern  
367 of Hi-C signals, which represents the maintenance of spermatogonia-type compartment strengths (Fig. 6c,  
368 Extended Data Fig. 8g). These results indicate that both SCML2 and A-MYB are necessary to establish a  
369 unique 3D chromatin organization of the X chromosome.

370  
371 Since meiotic sex chromosome are segregated from autosomes through the formation of the XY  
372 body, we examined interchromosomal interactions between the X and autosomes. In wild-type PS, these  
373 interchromosomal interactions decreased during spermatogenesis (Fig. 6d, Extended Data Fig. 9). In  
374 contrast, in *Scml2*-KO and *A-myb* mutant PS, interchromosomal interactions remained (Fig. 6d, Extended  
375 Data Fig. 9), indicating that the segregation of the sex chromosomes from autosomes is dependent on  
376 SCML2 and A-MYB (Fig. 6f). Interchromosomal interactions between autosomes remain intact in *Scml2*-  
377 KO and *A-myb* mutant PS, including the association of pericentromeric heterochromatin and telomeres as  
378 well as the overall association of autosomes (represented by “X” shape signals on Hi-C maps as described  
379 previously<sup>51</sup>) (Fig. 6e). Therefore, the role of SCML2 and A-MYB in regulating interchromosomal  
380 interactions of meiotic chromosomes is specific to the interaction between the X chromosome and  
381 autosomes (Fig. 6g).

382  
383 In summary, we conclude that SCML2 and A-MYB are required for the establishment of the  
384 unique 3D chromatin architecture of the sex chromosomes (Fig. 6h) and the formation of the segregated  
385 XY body.

386

## 387 **Discussion**

388 In this study, we determined the high-resolution 3D genome architecture of cell types  
389 representative of different stages of spermatogenesis and defined regulatory mechanisms underlying the  
390 transition from mitotic spermatogonia to meiotic spermatocytes. We demonstrated that, in spermatogonia,  
391 CTCF-mediated 3D contacts at meiotic SE are pre-established. Since meiotic SEs instruct the burst of  
392 meiotic gene expression, these poised 3D contacts represent a mechanism to maintain the cellular identity  
393 of male germ cells during spermatogenesis. Thus, we show that pre-programming through 3D contacts  
394 represents a novel feature of epigenetic priming. Of note, the poised 3D contacts in juvenile  
395 spermatogonia reflect the gene expression program of meiotic spermatocytes in adult testis, therefore they  
396 are maintained for a long time. Such 3D chromatin-based memories are likely to be prevalent in the  
397 germline, as sperm 3D chromatin is preset through histone modifications in late spermatogenesis as  
398 well<sup>20</sup>.

399

400 Epigenetic priming enables a rapid change in gene expression upon a signaling cue based on a  
401 pre-established chromatin state. It has been observed in various biological contexts, including immune  
402 cells<sup>52</sup>, neuronal<sup>53</sup> and cancer development<sup>54</sup>, as well as during spermatogenesis to instruct the gene  
403 expression program in late spermatogenesis<sup>15</sup>. Mechanistically, the pre-formation of enhancer-promoter  
404 pairs drives transcriptional changes upon differentiation in a variety of organisms and cell types<sup>55-57</sup>. Pre-  
405 formed enhancer-promoter pairs are associated with paused RNA polymerase<sup>56</sup> and a recent study showed  
406 that meiotic transcription bursts in the male germline are associated with the release of paused RNA  
407 polymerase, which is mediated by A-MYB and the testis-specific bromodomain protein BRDT<sup>59</sup>. We  
408 propose that at meiotic SEs these mechanisms operate in the context of 3D chromatin. In support of this  
409 hypothesis, in somatic cells, SEs are driven by another bromodomain protein, BRD4, as well as Mediator  
410 to form liquid-like condensates<sup>60</sup>, thereby providing spatial SE organization. In meiotic spermatocytes,  
411 BRDT is expressed in lieu of BRD4<sup>61</sup>. Thus, it is conceivable that preestablished 3D contacts provide  
412 venues for A-MYB and BRDT-driven spatial organization of meiotic SEs via phase separation to instruct  
413 the burst of meiotic gene expression (Fig. 7a).

414

415 A key question that remains is the timepoint at which the pre-programmed 3D contacts are  
416 established during spermatogenesis. Male germ cells acquire the androgenic epigenome in  
417 prospermatogonia (also known as gonocytes) prior to birth<sup>62</sup>. Prospermatogonia are arrested at the G<sub>0</sub>/G<sub>1</sub>  
418 phase of the cell cycle and genome-wide de novo DNA methylation takes place<sup>63,64</sup>. A recent study has  
419 shown that 3D chromatin reprogramming occurs in prospermatogonia<sup>65</sup>. Therefore, it is possible that the  
420 3D contacts necessary for the spermatogenic gene expression program are established in

421 prospermatogonia. Notably, the number of CTCF binding sites is reduced during in vitro differentiation  
422 of primordial germ cell-like cells to germline stem cell-like cells<sup>66</sup>, raising the possibility that the  
423 androgenic pattern of CTCF binding sites and 3D contacts are reprogrammed in prospermatogonia.  
424 CTCFL (also known as BORIS), a paralog of CTCF<sup>67,68</sup>, is expressed during spermatogenesis, along with  
425 CTCF, and some CTCF-binding sites overlap with CTCFL-binding sites<sup>69</sup>. Hence, CTCFL may be  
426 involved in the regulation of CTCF and CTCF-mediated 3D chromatin in spermatogenesis.

427

428 When germ cells enter meiosis, cohesin-mediated axial loops are formed along the chromosome  
429 axes to promote homolog pairing and recombination. Cytological analyses suggest that an average axial  
430 loop length is ~ hundred(s) kb<sup>46,70</sup> in mice, but the average axial loop lengths estimated from contact  
431 probability analyses of previous Hi-C studies<sup>24,51</sup> is larger (~ 1 Mb). The average length of stable meiotic  
432 chromatin loops detected by our Hi-C analysis is also ~ 1 Mb (Fig. 1f). Yet, we show that these stable  
433 chromatin loops are associated with transcription and are formed around super-enhancers (Fig. 5a),  
434 suggesting that they are distinct from axial loops. Further studies using independent approaches are  
435 needed to clarify if axial loop structures are distinct from stable chromatin loops.

436

437 Finally, our study showed that 3D chromatin organization of the sex chromosomes is regulated by  
438 SCML2 and A-MYB. After the initiation of MSCI directed by the DDR pathway at the early pachytene  
439 stage, SCML2 localizes on the sex chromosome after the mid-pachytene stage<sup>14</sup>. At that time, A-MYB  
440 regulates chromosome-wide spreading of H3K27ac on the sex chromosome<sup>17</sup>, downstream of the DDR  
441 factor RNF8<sup>71</sup>. RNF8 interacts with the MSCI initiator, MDCI, which functions as a  $\gamma$ H2AX binding  
442 protein<sup>50</sup>. Therefore, the DDR pathway coordinates both SCML2 and A-MYB-dependent processes on the  
443 sex chromosomes (Fig. 6h). Spermatogonia-type 3D chromatin features are retained in the *Scml2*-KO and  
444 *A-myb* mutant PS, suggesting that SCML2 and A-MYB are both required for the establishment of 3D  
445 chromatin features of the XY body (Fig. 7b). We suggest that SCML2 and A-MYB may work in concert  
446 on the sex chromosomes because SCML2 and RNF8 function together in the regulation of histone  
447 ubiquitination on meiotic sex chromosomes<sup>38</sup>. Similar to A-MYB's function on autosomes, A-MYB could  
448 drive a phased separated compartment of the sex chromosomes. Of note, this 3D chromatin feature of the  
449 male X chromosome in MSCI is distinct from that of the female inactive X chromosome. The inactivated  
450 female X chromosome splits into two mega domains bounded by the *Dxz4* locus and forms long-range  
451 loop structures called super loops (>7 Mb)<sup>27</sup>. CTCF binds around the *Dxz4* locus, and this structure is  
452 essential for the formation of mega domains and super loops in female cells<sup>72</sup>. This is quite different from  
453 the male X chromosome in meiosis, where short-range interactions are increased, and a domain structure  
454 is not clearly visible.

455 Overall, our results uncover the mechanisms underlying the organization of the meiotic chromatin  
456 structure on both autosomes and sex chromosomes and establish that CTCF-mediated pre-programming  
457 drives the burst of autosomal gene expression during male meiosis.

458

## 459 **Methods**

### 460 **Animals and germ cell isolation.**

461 Mice were maintained and used according to the guidelines of the Institutional Animal Care and  
462 Use Committee (protocol no. IACUC2018-0040) at Cincinnati Children's Hospital Medical Center. Wild-  
463 type C57BL/6J mice, *Scml2*-KO mice<sup>14</sup> on the C57BL/6J background, and *A-myb*<sup>mut/mut</sup> (*Mybl1*<sup>repro9</sup>)<sup>25</sup> on  
464 the C57BL/6J background were used for Hi-C analyses. Spermatogonia were isolated from C57BL/6J  
465 wild-type aged 6-8 days through magnetic cell-sorting (MACS) as described previously<sup>26</sup>. Pachytene  
466 spermatocytes and round spermatids, including *Scml2*-KO PS, *Scml2*-RS, and *A-myb* mutant PS, were  
467 isolated from adult testes through sedimentation velocity at unit gravity as described previously<sup>26,39,73</sup>.

468

### 469 **Hi-C library generation and sequencing.**

470 To generate and sequence Hi-C libraries, Hi-C was used using the Arima Hi-C kit, according to  
471 the manufacturer's instructions. We used the Arima-Hi-C kit, which enables high-resolution detection of  
472 3D chromatin by using a combination of multiple restriction enzymes.  $4 \times 10^5$  to  $1 \times 10^6$  cells were used for  
473 THY1<sup>+</sup> and KIT<sup>+</sup>,  $3 \times 10^6$  cells were used for WT PS, *Scml2*-KO PS, and *A-myb* mutant PS, and  $4 \times 10^6$   
474 cells were used for WT RS and *Scml2*-KO RS for crosslinking. For library preparation, Accel-NGS<sup>®</sup> 2S  
475 Plus DNA Library Kit (Swift Biosciences, Inc. Ann Arbor, MI) was used. All libraries were sequenced on  
476 Illumina HiSeq4000 sequencers according to the manufacturer's instructions.

477

### 478 **Hi-C data mapping.**

479 Paired-end .fastq files of Hi-C libraries were aligned and processed using the Juicer package<sup>74</sup>  
480 (version 1.5). In brief, each end of the raw reads was mapped separately to the *Mus musculus* mm10  
481 reference genome, and Hi-C pairs files were created using BWA<sup>75</sup> (version 0.7.3a). Mapping statistics are  
482 summarized in Supplementary Table 1. .hic files, a highly compressed binary file was created by Juicer  
483 tools pre. Matrix balancing was performed with the cooler software package (version 0.8.11) and  
484 visualized using the HiCEXplorer<sup>76</sup> (version 3.6) for use with the application hicPlotMatrix. To generate  
485 and visualize interaction frequency heat maps of whole chromosomes, Hi-C matrices at 250-kb resolution  
486 were imported to the software package HiCEXplorer for use with the application hicPlotMatrix. To aid  
487 visual comparisons between the datasets, matrices were natural log transformed. To analyze differential  
488 interaction frequencies between samples, the HiCEXplorer application hicCompareMatrices was used to

489 generate log<sub>2</sub> ratios of interaction frequency matrices between two separate datasets and then visualized  
490 by hicPlotMatrix.

491

### 492 **Hi-C: Evaluation of Hi-C biological replicates.**

493 Pearson correlation coefficients between Hi-C biological replicates at each stage were obtained  
494 by using the hicCorrelate included in HiCEXplorer using the cool files binned at 10kb with the parameter  
495 ‘--log1p --method pearson’. A range from 10kb to 5Mb was used in the calculations. The reproducibility  
496 of the results was confirmed with biological replicates (Extended Data Fig. 1c).

497

### 498 **Hi-C: Contact frequency.**

499 Enrichment of Hi-C counts at different genomic ranges/distances to whole chromosomes was  
500 calculated for autosomes and X chromosomes respectively using hicPlotDistVsCounts including  
501 HiCEXplorer with the option ‘--maxdepth 300000000’.

502

### 503 **Hi-C: Identification of chromatin loops.**

504 Chromatin loops were called by using the HiCEXplorer for the use with the application  
505 hicDetectLoops using cool files binned at 5kb, 10kb and 25kb respectively with the parameter ‘--  
506 maxLoopDistance 2000000’. The cool files were generated from hic files created using only reads with a  
507 MAPQ score of 30 or higher using the -q 30 option during Juicer tools pre procedure and converted to  
508 cool files using the hicConvertFormat included in HiCEXplorer and the cooler balance included in cooler.  
509 After detecting chromatin loops at each resolution, the loops from each resolution were merged by using  
510 the hicMergeLoops included in HiCEXplorer with the ‘-r 25000’ option. CTCF, H3K4me3/H3K27ac and  
511 H3K27me3-dependent loops were detected using bed files of CTCF (this study),  
512 H3K4me3<sup>39</sup>/H3K27ac<sup>17,38</sup> or H3K27me3<sup>39</sup> ChIP-seq datas by the hicValidateLocations with the ‘--method  
513 loops --resolution 25000’ option. Juicer tools compare was used to detect common loops between the two  
514 types of chromatin loops, with the option ‘-m 25000 0 mm10’. The loop listed as “Common”, “A” or “B”  
515 in parent\_list of the output data was used as the common loops between the two loops. IGV<sup>77</sup> (version  
516 2.8.3) was used to visualize chromatin loops in the genomic track view. Genes where the anchor site of  
517 the loop overlaps with the TSS region using refTSS\_v3.1\_mouse\_annotation.txt  
518 ([https://refTSS.riken.jp/datafiles/3.1/mouse/gene\\_annotation/](https://refTSS.riken.jp/datafiles/3.1/mouse/gene_annotation/)) were identified as a group of genes  
519 associated with a specific loop.

520

### 521 **Hi-C: Identification of Topologically Associated Domains (TADs).**

522 TADs were detected by the hicFindTads including HiCEXplorer using the cool files binned at  
523 25kb with the parameter ‘--correctForMultipleTesting fdr --minDepth 80000 --maxDepth 800000 --step



524 40000 minBoundaryDistance 80000 --thresholdComparisons 0.01 --delta 0.01'. TAD boundaries were  
525 analyzed by extending 25 kb each upstream and downstream from the TAD boundaries detected by  
526 hicFindTads for downstream analysis. CTCF, H3K4me3/H3K27ac and H3K27me3-dependent TAD  
527 boundaries were detected using bed files of CTCF, H3K4me3/H3K27ac or H3K27me3 ChIP-seq datas by  
528 the hicValidateLocations with the '--method tad --resolution 25000' option. Common TAD boundaries  
529 between the two data sets were detected using bedtools intersect, and were assumed to be common if they  
530 covered even 1 bp.

531

### 532 **Hi-C: Identification of mitotic and meiotic specific SEs and genomic sites interacting with meiotic** 533 **SEs.**

534 The SE files downloaded from Maezawa et al., 2020<sup>17</sup> was used with a modified version of the  
535 SE file. The SEs detected in THY1<sup>+</sup> and KIT<sup>+</sup>, excluding those overlapping with the SEs detected in PS  
536 and RS, were used as mitotic specific SEs, and conversely, the SEs detected in PS and RS, excluding  
537 those detected in THY1<sup>+</sup> and KIT<sup>+</sup>, were used as meiotic specific SEs in this study. To determine whether  
538 PS chromatin loops overlapped with meiotic SE, the anchor sites of PS chromatin loops were added +0.4  
539 Mb upstream and downstream, and it was determined whether this region overlapped with meiotic SE.

540 The regions interacting with meiotic SEs followed the method described in '3.4.6 Identification  
541 of super-enhancer-promoter interactions' by Sakashita et al., 2023<sup>78</sup>. The first step in the analysis is to  
542 calculate the quality of each viewpoint (SE locus) using the chicQualityControl program including in  
543 HiCExplorer, which considers the sparsity of the Hi-C contact frequency with the '--sparsity 0.3' option.  
544 Next, using the bed file containing the filtered viewpoints and the program  
545 chicViewpointBackgroundModel, a background model of all given viewpoints is calculated based on the  
546 Hi-C contact matrix, with the option to set the range of the background model to 500 kb with the '--  
547 fixateRange 500000' option. Using the chicViewpoint program, all interaction points in physical contact  
548 with the SE locus are detected based on the background model, ranging up to 500 kb (--range 500000  
549 500000)). Finally, using the chicSignificantInteractions program with the '-p 0.05 --range 500000 500000  
550 --loosePValue 0.1', only significant interaction points (P<0.05 (-p 0.05)) were extracted, which were  
551 designated as genomic regions interacting with meiotic specific SEs.

552

### 553 **Hi-C: Pile-up analysis.**

554 Pile-up analysis was performed using coolup.py<sup>36</sup> (version 0.9.5) to visualize the average  
555 interaction strength of chromatin loops and TADs. Chromatin loops were analyzed using the cool files  
556 binned at 10kb resolution and a bedpe file of the corresponding chromatin loops. TAD domains were  
557 analyzed using the cool files binned at 25kb resolution and bed file showing TAD domains with the '--  
558 rescale --local' option. TAD boundaries were analyzed using the cool files binned at 25kb resolution and

559 bed file showing TAD boundaries with the ‘--pad 500 --local’ option. For the analysis of interactions  
560 between mitotic or meiotic SEs, bed files showing mitotic or meiotic SEs were used. For the pile-up  
561 analysis showing the interaction among mitotic or mitosis-specific SEs, each SEs were analyzed using the  
562 cool files binned at 10kb resolution and bed file showing TAD boundaries with the ‘--pad 500’ option.  
563 For the interaction between CTCFs on meiotic SEs, the overlap regions between meiotic SEs and CTCF  
564 binding sites in PS were used for analysis. CTCFs on meiotic SEs were analyzed using the cool files  
565 binned at 10kb resolution and bed files showing TAD boundaries with the ‘--pad 500’ option. Piled-up  
566 data were visualized by performing plotup.py.

567

### 568 **RNA-seq data analysis.**

569 Row RNA-seq reads after trimming by Sickle (<https://github.com/najoshi/sickle>) (version 1.33)  
570 trimmed regions with quality less than 30 and excluded reads that were less than 20 bp. Trimmed  
571 sequencing reads were aligned to the *Mus musculus* mm10 reference genome using HISAT2<sup>79</sup> (version  
572 2.2.1) with default parameters. All unmapped reads and non-uniquely mapped reads were filtered out and  
573 then sorted by samtools<sup>80</sup> (version 1.14) with default parameters. The output bam file was assembled and  
574 quantified using StringTie<sup>81</sup> (version 2.2.1) based on the mouse gene annotation  
575 ([gencode.vM25.annotation.gtf](https://www.encodeproject.org/annotation/gtf)). Transcripts per million (TPM) value was used for downstream analyses.

576 Genes associated with *Scml2*-KO PS-specific loops were defined with genes whose TSS regions  
577 overlapped with the anchor sites of these loops. Then, only genes with TPM values greater than 1 in any  
578 of the cells were extracted and used for analysis. Violinplot was drawn using the R package ggplot2. The  
579 log10-transformed values of TPM values+1 were used for statistical analysis and plotting. GO term  
580 analysis was performed using the website tool DAVID (<https://david.ncifcrf.gov/home.jsp>). GO term was  
581 visualized by ggplot2 of the R package based on gene number, fold enrichment, and *P* value.

582

### 583 **ChIP-seq data analysis.**

584 Cross-linking ChIP-seq was performed for CTCFs of THY1<sup>+</sup>, WT PS, and *Scml2*-KO PS, using  
585 the same methods as previously reported<sup>17</sup>. The reproducibility of the results was confirmed with  
586 biological replicates (Extended Data Fig. 2a, 3b). Row ChIP-seq reads after trimming by Sickle trimmed  
587 regions with quality less than 20 and excluded reads that were less than 20 bp. Trimmed sequencing reads  
588 were aligned to the *Mus musculus* mm10 reference genome using Bowtie2<sup>75</sup> (version 2.4.5) with default  
589 parameters. All unmapped reads and non-uniquely mapped reads were filtered out and then sorted by  
590 samtools (version 1.14) with default parameters. All unmapped and uniquely mapped reads were filtered  
591 out, and sorted by default parameters using samtools, and then ‘MarkDuplicates’ command in Picard tools  
592 (version 2.26.9; <https://broadinstitute.github.io/picard/>) was used to remove PCR duplicates by using the  
593 option ‘VALIDATION\_STRINGENCY=LENIENT ASSUME\_SORTED=true

594 REMOVE\_DUPLICATES=true'. After this process, the bam files sorted by samtools again were used for  
595 downstream analysis.

596 To compare biological replicates, Pearson correlation coefficients were calculated and plotted by  
597 multiBamSummary bins and plot correlation from deepTools<sup>82</sup> (version 3.5.1). For visualization of ChIP-  
598 seq using IGC, normalized genome coverage tracks based on counts per million mapped reads were  
599 generated as bigwig files using bamCoverage function of deepTools with '--binSize=5 --normalization  
600 CPM' parameter. Bigwig files were also used for visualization of ChIP-seq data using IGV. Peak calls  
601 were identified using MACS2<sup>83</sup> (version 2.2.7.1). The ngs.plot was used to draw tag density and heat  
602 maps for read enrichment within  $\pm$  2kb for CTCF and histone modification,  $\pm$ 3kb for A-MYB analysis at  
603 meiotic SEs interacting sites, and  $\pm$  5kb for A-MYB analysis at anchor sites of PS chromatin loops<sup>84</sup>. A-  
604 MYB binding genes were extracted using the online website GREAT (version 4.0.4;  
605 <http://great.stanford.edu/public/html/>) for genes with TSS in the peak  $\pm$ 2 kb region of A-MYB ChIP-seq.  
606

## 607 **Statistics.**

608 Statistical methods and *P* values for each plot are listed in the figure legends and/or in the  
609 Methods. For all experiments, no statistical methods were used to predetermine sample size. Experiments  
610 were not randomized, and investigators were not blinded to allocation during experiments and outcome  
611 assessments.

612

## 613 **Data availability**

614 Hi-C and CTCF ChIP-seq datasets were deposited in the Gene Expression Omnibus under  
615 accession no. GSE244681. All other next-generation sequencing datasets used in this study are publicly  
616 available. RNA-seq data from THY1<sup>+</sup> spermatogonia, PS and RS were downloaded from the GEO  
617 (accession no. GSE55060). ChIP-seq data for H3K4me2, H3K4me2 and H3K27me3 and RNA-seq data  
618 from KIT<sup>+</sup> spermatogonia were downloaded from the GEO (GSE89502). ChIP-seq data for H3K27ac in  
619 WT PS were downloaded from the GEO (GSE107398). H3K27ac in THY1<sup>+</sup> and KIT<sup>+</sup> spermatogonia and  
620 input for CTCF ChIP-seq were downloaded from the GEO (GSE130652). A-MYB ChIP seq in whole  
621 testis was downloaded from GEO (GSE44588). Source data are provided in this paper.  
622

622

## 623 **Code availability**

624 Source code for all software and tools used in this study, with documentation, examples, and  
625 additional information, is available at the URLs listed above.  
626

626

627

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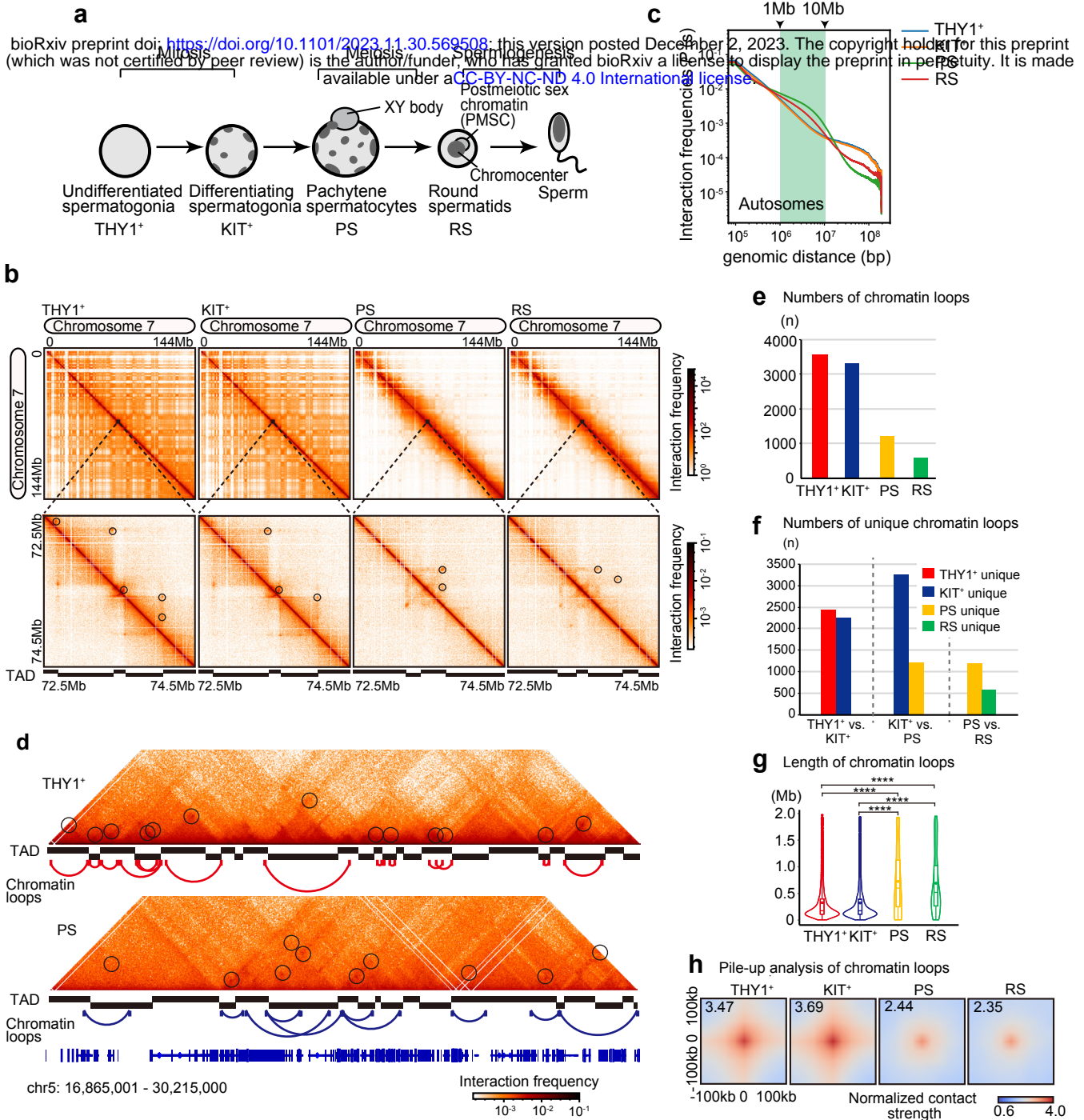
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## 806 **Author contributions**

807 Y.K. and S.H.N. designed the study. K.T. and S.M performed experiments. Y.K. performed the  
808 computational analyses. Y.M. contributed to the computational analyses. A.S. contributed to developing a  
809 computational tool. Y.K., N.K, and S.H.N interpreted the computational analyses. Y.K. and S.H.N. wrote  
810 the manuscript with critical feedback from all other authors. S.H.N. supervised the project.

## 811 **Competing interest statement**

812 The authors declare no competing interests.



**Figure 1. 3D chromatin reprogramming and inter-TAD chromatin loop formation in meiosis.**

**a**, Schematic of the stages of mouse spermatogenesis analyzed in this study. THY1<sup>+</sup>: undifferentiated spermatogonia; KIT<sup>+</sup>: differentiating spermatogonia; PS: pachytene spermatocytes; RS: round spermatids.

**b**, Hi-C maps showing normalized Hi-C interaction frequencies (100kb bins, chromosome 7) in THY1<sup>+</sup>, KIT<sup>+</sup>, PS, and RS. 10kb bins normalized Hi-C matrices were used for the zoom-in. Black circles in the Hi-C map indicate chromatin loops.

**c**, Hi-C interaction frequency probabilities  $P$  stratified by genomic distance  $s$  for each cell type shown (100kb bins). All autosomes were analyzed.

**d**, Hi-C interaction heat maps (25kb bins, chromosome 5, 16, 865, 001-30, 215, 000bp) in THY1<sup>+</sup> and PS. Chromatin loops are indicated by black circles, red lines in THY1<sup>+</sup>, and blue lines in PS.

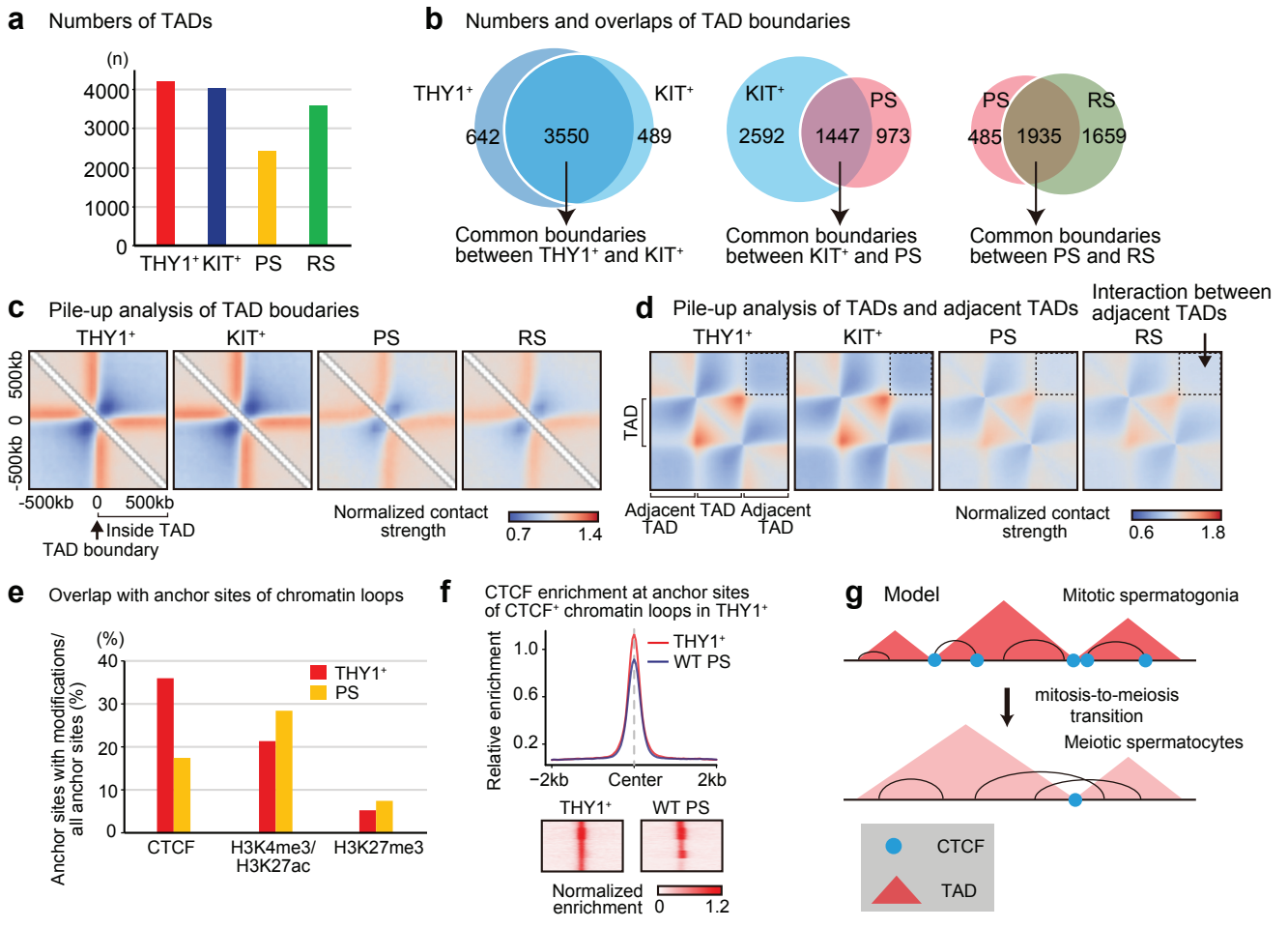
**e**, Numbers of chromatin loops (n) detected from each Hi-C data set (merged results for each using 5kb, 10kb, and 25kb bin data).

**f**, Numbers of unique chromatin loops comparing each pairwise developmental stage.

**g**, Chromatin loop length (Mb) from each Hi-C data set (merged results for each using 5kb, 10kb, and 25kb bin data). The number of loops used in the analysis was equal to the number shown in e (THY1<sup>+</sup>: n=3,562, KIT<sup>+</sup>: n=3,336, PS: n=1,223, RS: n=609). The box indicates the 25th, median and 75th percentiles, and the dot in the box indicates mean. Statistical analysis is based on Bonferroni correction. \*\*\*\* indicates  $p < 2e^{-16}$ .

**h**, Chromatin loop pile-up in each cell type with 100kb padding. Color represents normalized contact strength in the log scale. The normalized contact strength values in the central pixel are shown on the top left.





**Figure 2. TAD and chromatin loop reorganization during spermatogenesis.**

**a**, Numbers of TADs (n) detected from each Hi-C data set (25kb bins).

**b**, Venn diagram showing numbers and overlaps of TAD boundaries in each developmental stage.

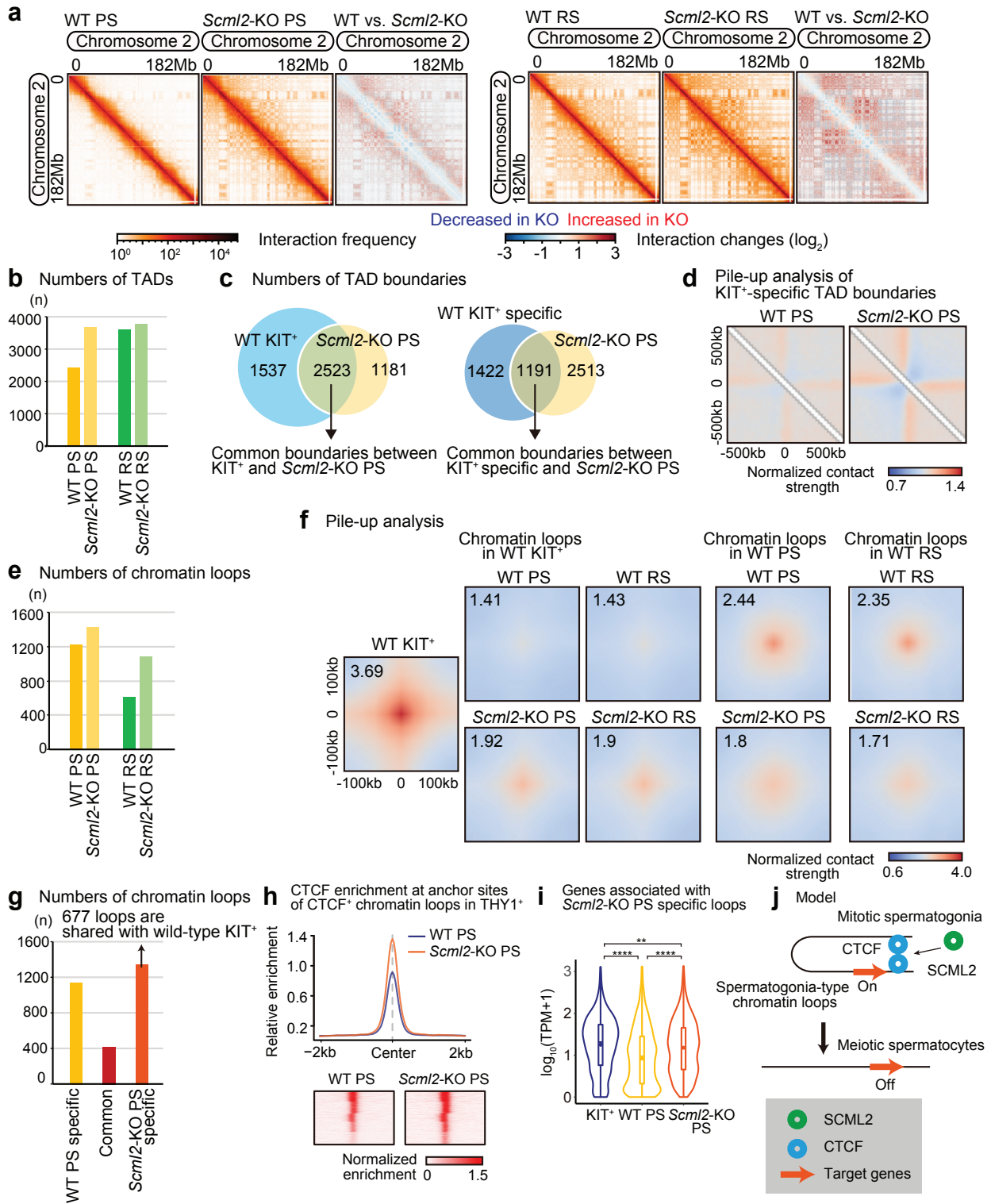
**c**, Local pile-up analysis of TAD boundaries in each cell type. 10kb bins Hi-C data with 500 kb padding around the central pixel. Color represents normalized contact strength in the log scale.

**d**, Local rescaled pile-ups of TADs from 10kb bin Hi-C data in each cell type. The dotted regions represent interactions between adjacent TADs.

**e**, Ratio of accumulation of CTCF, H3K4me3/H3K27ac, or H3K27me3 at the anchor sites of chromatin loops in THY1+ and PS.

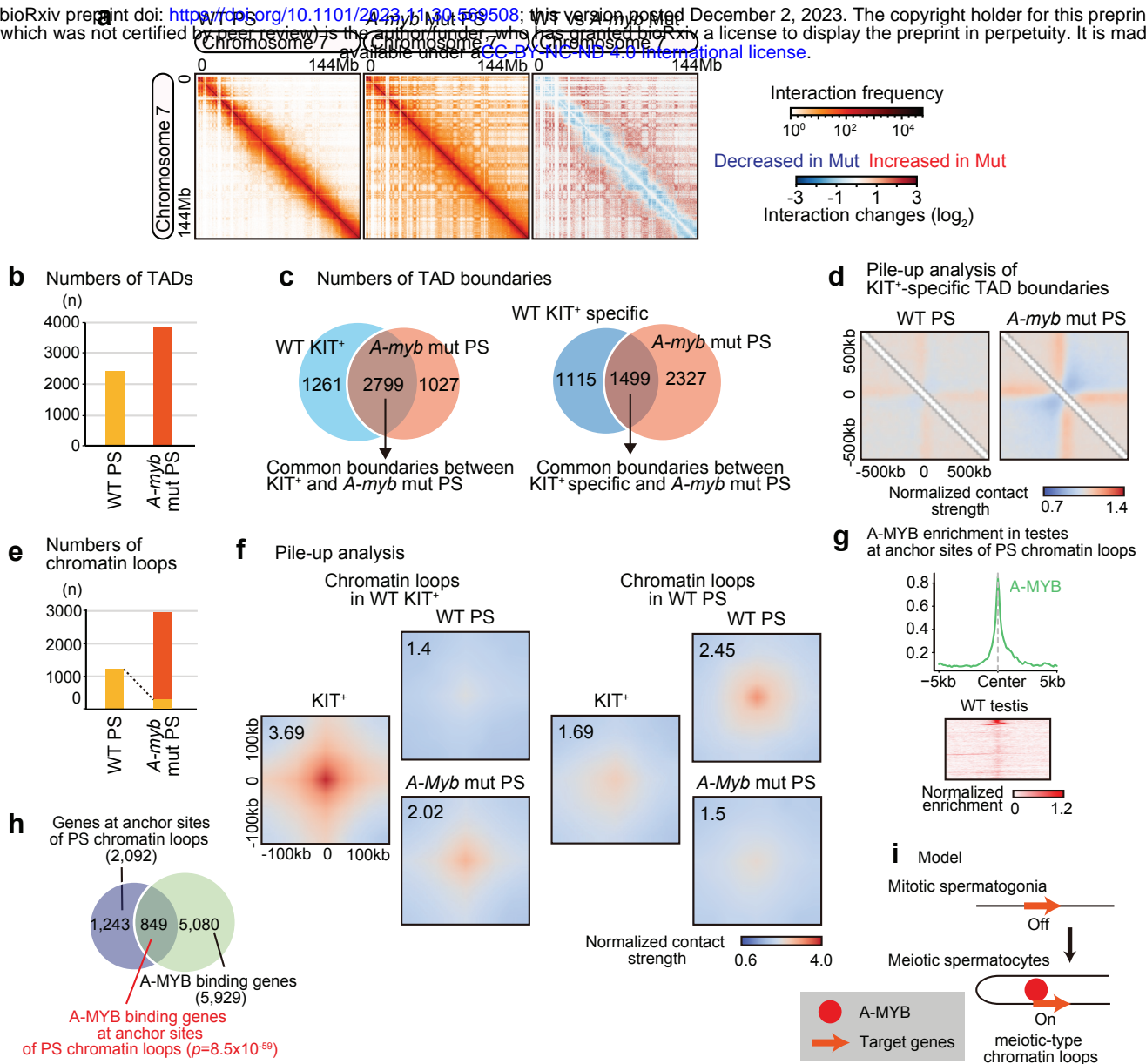
**f**, CTCF enrichment at anchor sites of CTCF-dependent chromatin loops in THY1+ (detected in panel e, 2,532 sites). Heat maps for each locus are shown at the bottom.

**g**, Model showing TAD and chromatin loop reorganization at the mitosis-to-meiosis transition.



**Figure 3. SCML2 is required for the resolution of spermatogonia-type 3D chromatin.**

- a**, Heat maps showing normalized Hi-C interaction frequencies (100kb bins, chromosome 2) in wildtype (WT) PS, *Scml2*-KO PS (left), and WT RS and *Scml2*-KO RS (right). Red and blue Hi-C maps represent a log<sub>2</sub> ratio comparison of Hi-C interaction frequencies between WT and *Scml2*-KO.
- b**, Numbers of TADs (n) detected from each Hi-C data set (25kb bins) in WT PS, *Scml2*-KO PS, WT RS, and *Scml2*-KO RS.
- c**, Venn diagram showing the overlap between all KIT<sup>+</sup> TAD boundaries and *Scml2*-KO PS TAD boundaries (left), and the overlap between KIT<sup>+</sup>-specific TAD boundaries and *Scml2*-KO PS TAD boundaries (right). KIT<sup>+</sup>-specific boundaries are defined by excluding TAD boundaries detected in WT PS.
- d**, Local pile-up analysis of KIT<sup>+</sup> specific TAD boundaries in WT PS and *Scml2*-KO PS.
- e**, Numbers of chromatin loops (n) detected in each Hi-C data set (merged results for each using 5kb, 10kb, and 25kb bin data) in WT PS, *Scml2*-KO PS, WT RS, and *Scml2*-KO RS.
- f**, Chromatin loop pile-up analysis in each cell type with 100kb padding. The normalized contact strength in the central pixel is displayed on the top left.
- g**, Numbers of specific and common chromatin loops between WT PS and *Scml2*-KO PS. 677 *Scml2*-KO PS-specific loops overlapped with loops detected in KIT<sup>+</sup>. Overlapping loops were detected by Juicer.
- h**, CTCF enrichment in WT PS and *Scml2*-KO PS at the anchor site of CTCF-chromatin loops in THY1<sup>+</sup> spermatogonia.
- i**, Violin plots of RNA-seq reads converted to log<sub>10</sub> (TPM+1) value for genes associated with *Scml2*-KO PS specific loops in KIT<sup>+</sup>, WT PS and *Scml2*-KO PS. 1,243 genes were identified by extracting genes present in the anchor site of *Scml2*-KO PS-specific loops. The box indicates the 25th, median and 75th percentiles, and the dot in the box indicates mean. Statistical analysis is based on Bonferroni correction. \*\*\*\*:  $p < 2e^{-16}$ , \*\*:  $p < 0.005$ .
- j**, Model of resolution of spermatogonia-type 3D chromatin by SCML2.



**Figure 4. A-MYB is required for the formation of meiotic-type 3D chromatin.**

**a**, Heat maps showing normalized Hi-C interaction frequencies (100kb bins, chromosome 7) in WT PS, *A-myb* mutant PS (left). Red and blue Hi-C maps represent a log<sub>2</sub> ratio comparison of Hi-C interaction frequencies between wild-type and *A-myb* mutant PS.

**b**, Number of TADs (n) detected from each Hi-C data set (25kb bins) in WT PS and *A-myb* mutant PS.

**c**, Venn diagram showing the overlap between all KIT<sup>+</sup> TAD boundaries and *A-myb* mutant PS TAD boundaries (left), and the overlap between KIT<sup>+</sup>-specific TAD boundaries and *A-myb* mutant PS TAD boundaries (right). KIT<sup>+</sup>-specific boundaries are defined by excluding boundaries detected in WT PS.

**d**, Local pile-up analysis of KIT<sup>+</sup>-specific TAD boundaries in WT PS and *A-myb* mutant PS.

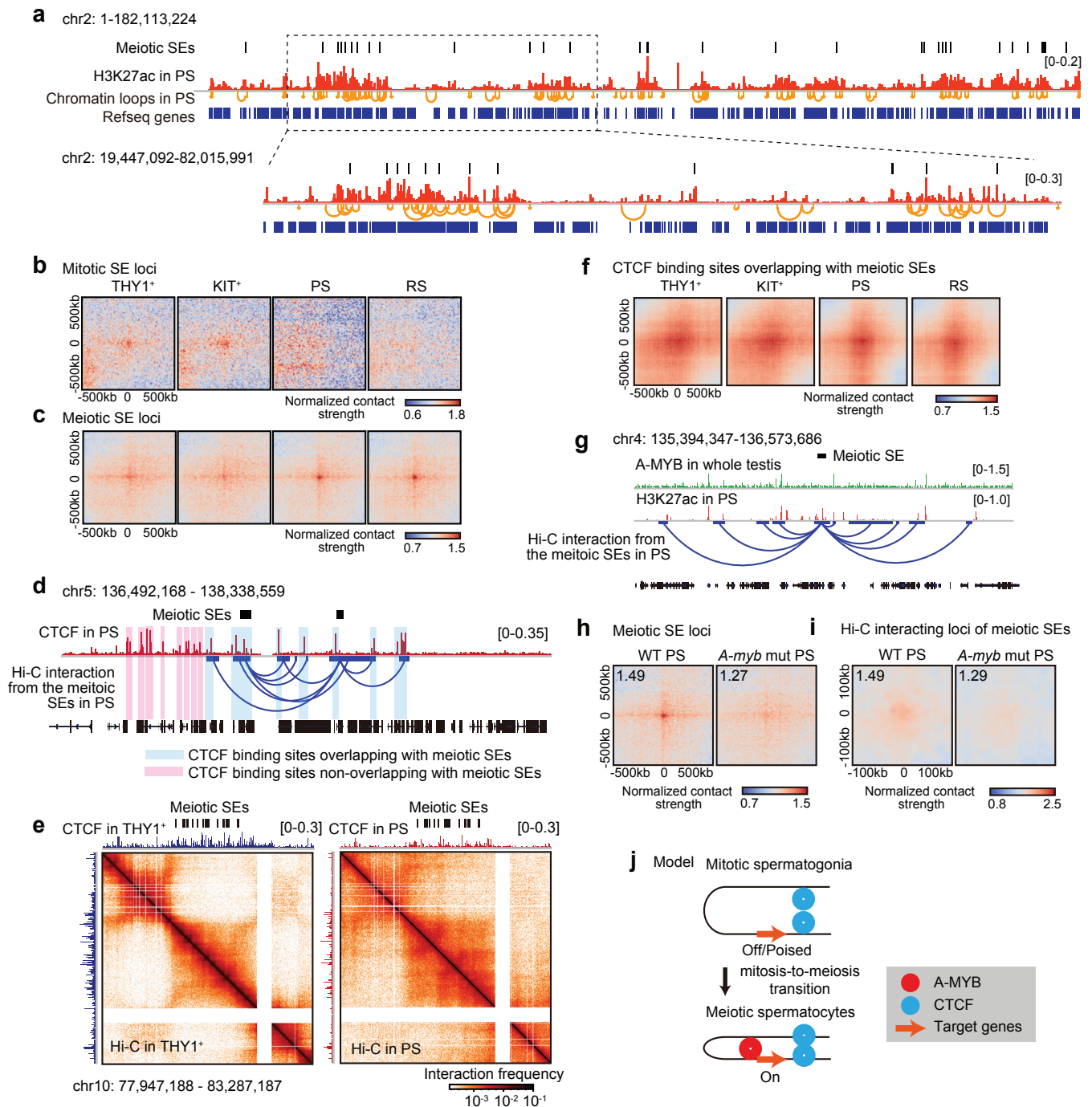
**e**, Number of chromatin loops (n) detected in each Hi-C data set (merged results for each using 5kb, 10kb, and 25kb bin data) in WT PS and *A-myb* mutant PS. Yellow area in the graph of *A-myb* mutant PS indicate that the same loops are detected in WT PS (357 loops).

**f**, Chromatin loop pile-up analysis in each cell type with 100kb padding. The normalized contact strength in the central pixel is displayed on the top left.

**g**, ChIP-seq data for A-MYB using whole testis at the regions adjacent to TSS of 849 genes that overlap with anchor sites of chromatin loops in PS.

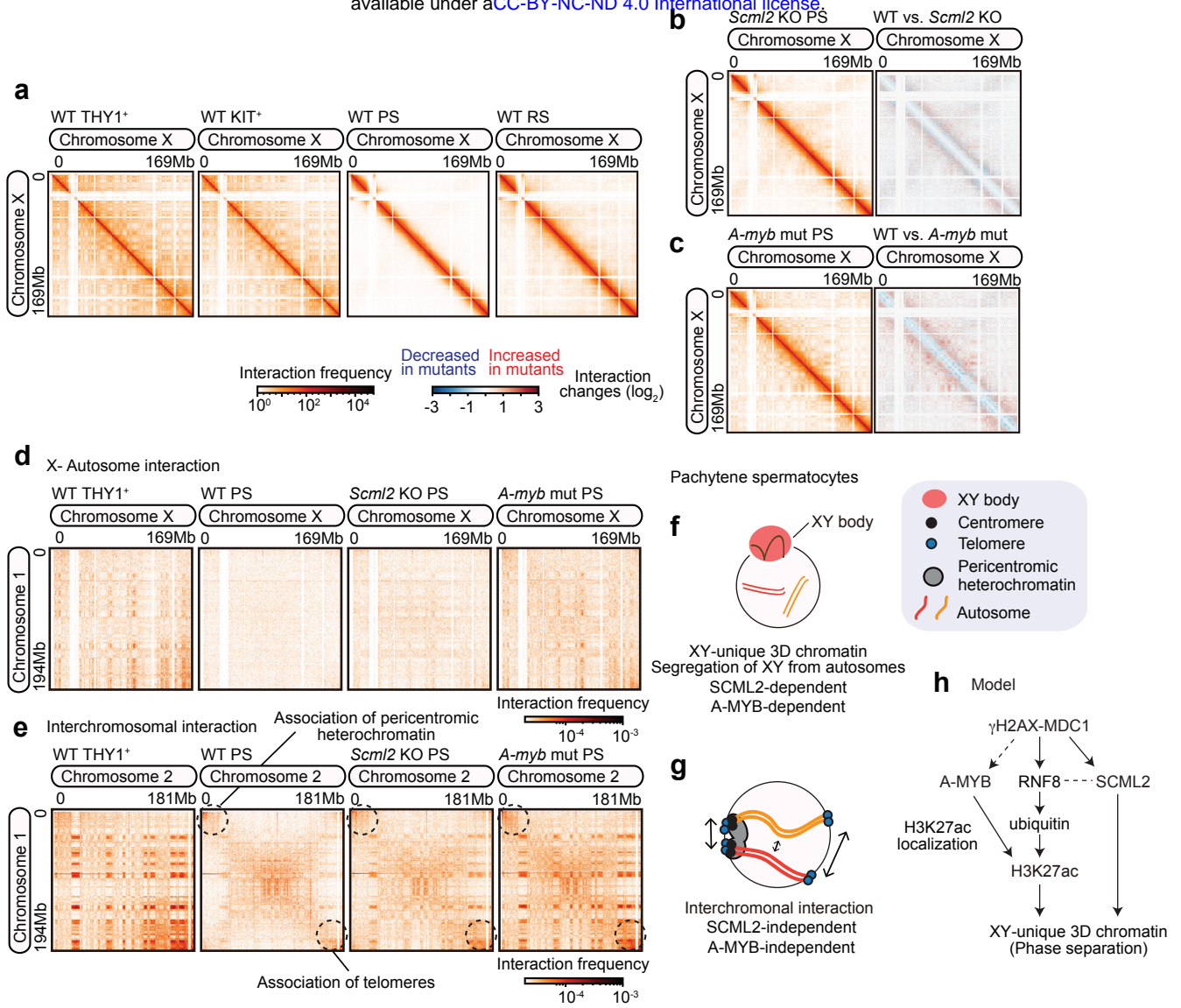
**h**, Venn diagram showing the intersection of genes located at anchor sites of chromatin loops in PS (blue) and all A-MYB bound genes (green). The overlap is statistically significant ( $p=8.5 \times 10^{-59}$ ) compared to the proportion of all A-MYB bound genes to all RefSeq genes based on the hypergeometric test.

**i**, Model of the establishment of meiotic-type chromatin loops by A-MYB.



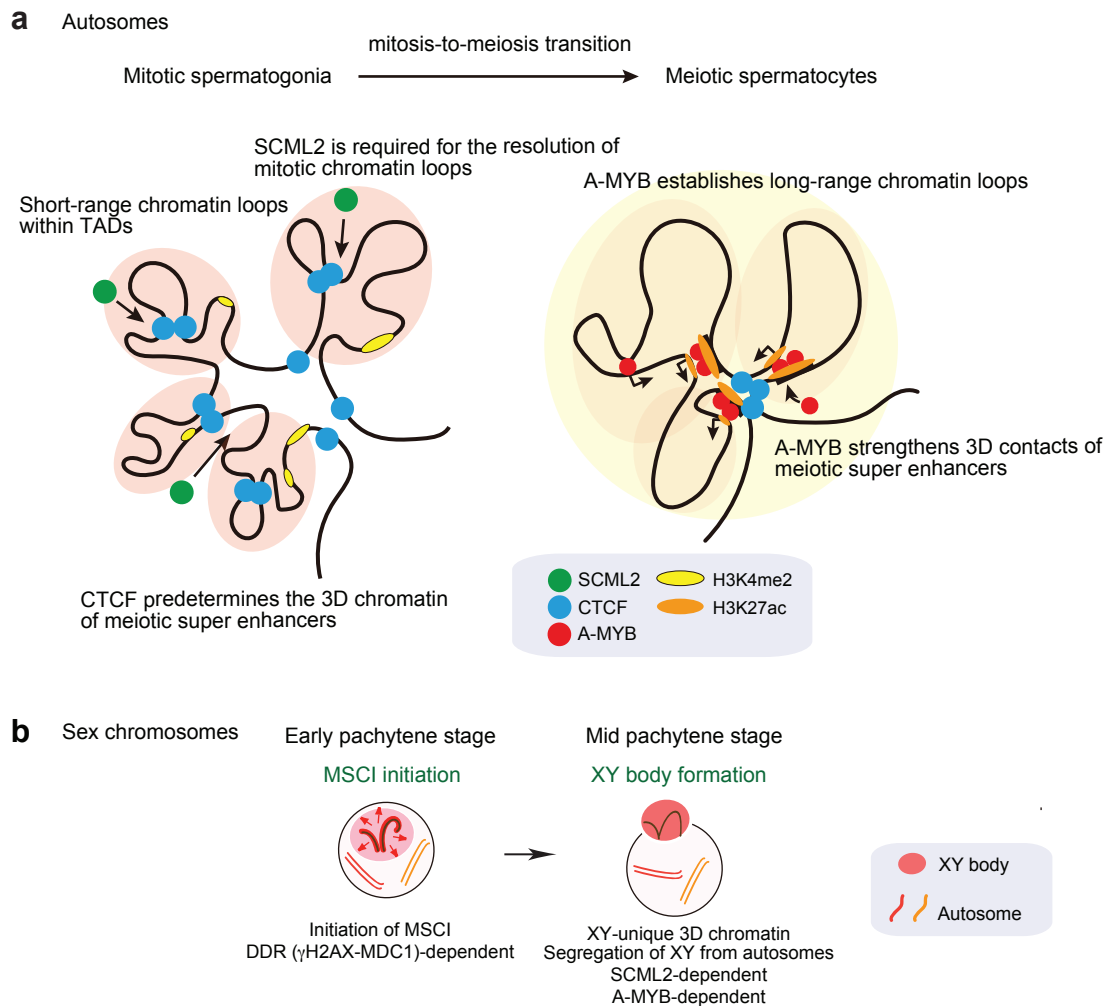
**Figure 5. Meiotic super-enhancers are poised with 3D chromatin.**

- a**, Track view showing meiotic SEs, H3K27ac, and chromatin loops in PS on the entire chromosome 2 (top). Enlargement of the boxed area is shown below.
- b**, Pile-up analysis of averaged intersections of mitotic SEs with 500kb paddles.
- c**, Pile-up analysis of averaged intersections of meiotic SEs with 500kb paddles.
- d**, Track view showing CTCF distribution and Hi-C interactions of the meiotic SEs in PS on a region of chromosome 5. Pink highlights indicate CTCF binding sites that do not overlap with meiotic SEs; blue highlights indicate CTCF binding sites that overlap with meiotic SEs and their loops.
- e**, CTCF binding and Hi-C maps of THY1<sup>+</sup> spermatogonia and PS around meiotic SEs (25kb bins, chr10: 77,947,188-83,287,187).
- f**, Pile-up analysis showing average interactions of CTCF binding sites overlapping with meiotic SEs. The pile-up analysis in THY1<sup>+</sup>, KIT<sup>+</sup>, PS, and RS is based on the Hi-C data from each developmental stage and the genomic coordinates of the CTCF binding sites that overlapped with meiotic SEs or their interacting genomic regions.
- g**, Track view showing the distributions of A-MYB binding and H3K27ac around meiotic SEs. Hi-C interaction from the meiotic SEs is also shown.
- h**, Pile-up analysis showing average interactions of meiotic SEs with 500kb paddles in WT PS and *A-myb* mutant PS. The normalized contact strength in the central pixel is displayed on the top left.
- i**, Pile-up analysis showing average interactions of loci that interacted with meiotic SEs based on Hi-C data with 100kb paddles in WT PS and *A-myb* mutant PS.
- j**, Model of the predetermination of 3D chromatin at meiotic SE loci via CTCF in mitotic spermatogonia. A-MYB strengthens these 3D contacts in meiotic spermatocytes.



**Figure 6. SCML2 and A-MYB establish unique 3D chromatin of the meiotic sex chromosomes**

- a**, Hi-C maps of the X chromosome showing normalized Hi-C interaction frequencies (100kb bins) in WT THY1<sup>+</sup>, KIT<sup>+</sup>, PS, and RS.
- b**, Heat maps showing normalized Hi-C interaction frequencies (100kb bins, chromosome X) in *Scml2*-KO PS (left). Red and blue Hi-C maps represent a log<sub>2</sub> ratio comparison of Hi-C interaction frequencies between wild-type and *Scml2*-KO PS (right).
- c**, Heat maps showing normalized Hi-C interaction frequencies (100kb bins, chromosome X) in *A-myb* mutant PS (left). Red and blue Hi-C maps represent a log<sub>2</sub> ratio comparison of Hi-C interaction frequencies between wild-type and *A-myb* mutant PS (right).
- d**, Heat maps showing normalized Hi-C interchromosomal interactions (250-kb bins, chromosomes 1 and X) for WT THY1<sup>+</sup>, WT PS, *Scml2*-KO PS and *A-myb* mutant PS.
- e**, Heat maps showing normalized Hi-C interchromosomal interactions (250-kb bins, chromosomes 1 and 2) for WT THY1<sup>+</sup>, WT PS, *Scml2*-KO PS and *A-myb* mutant PS.
- f**, Model for the establishment of a unique 3D chromatin in the XY body and segregation of XY from autosomes in PS.
- g**, Model of interchromosomal interactions in pachytene spermatocytes.
- h**, Schematic of the molecular pathway that establishes a XY-unique 3D chromatin in pachytene spermatocytes.



**Figure 7. Models of 3D chromatin dynamics and gene regulation on autosomes and sex chromosomes during spermatogenesis.**

**a**, Model showing the changes in chromosome interactions from mitotic spermatogonia to meiotic spermatocytes on autosomes.

**b**, Model of 3D chromatin dynamic on the sex chromosomes. At the onset of MSCI at the early-pachytene stage, DDR initiated MSCI and, subsequently, SCML2 and A-MYB establish unique 3D chromatin of the sex chromosomes and segregate the sex chromosomes from autosomes at the mid-pachytene stage.