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Principles of Meiotic Chromosome Assembly

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16 Abstract

During meiotic prophase, chromosomes organise into a series of chromatin loops emanating from 17 a proteinaceous axis, but the mechanisms of assembly remain unclear. Here we elucidate how 18 this elaborate three-dimensional chromosome organisation is underpinned by genomic sequence 19 in Saccharomyces cerevisiae. Entering meiosis, strong cohesin-dependent grid-like Hi-C 20 interaction patterns emerge, reminiscent of mammalian interphase organisation, but with distinct 21 regulation. Meiotic patterns agree with simulations of loop extrusion limited by barriers, yet are 22 patterned by convergent transcription rather than binding of the mammalian interphase factor, 23 CTCF, which is absent in S. cerevisiae-thereby both challenging and extending current 24 paradigms of local chromosome organisation. While grid-like interactions emerge independently 25 of meiotic chromosome synapsis, synapsis itself generates additional compaction that matures 26 differentially according to telomere proximity and chromosome size. Collectively, our results 27 elucidate fundamental principles of chromosome assembly and demonstrate the essential role of 28 cohesin within this evolutionarily conserved process. 29

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30 Introduction and Results

During meiosis, eukaryotic chromosomes are broken, repaired and paired with their homologs 31 followed by two rounds of segregation-a series of events accompanied by dynamic structural 32 changes of the chromosomes. Most prominent is the paired arrangement of pachytene 33 chromosomes into a dense array of chromatin loops emanating from proteinaceous axes linked 34 by a central core, the synaptonemal complex (SC), which is highly conserved across 35 eukarvotes^{1,2}. In S. cerevisiae, structural components include the meiotic cohesin kleisin subunit, 36 Rec8³, the transverse filament, Zip1⁴, the axial/lateral elements, Hop1 and Red1^{5,6}, and the pro-37 DSB factors Rec114-Mei4-Mer2 (RMM)^{7,8}. Much of our understanding of meiotic chromosome 38 structure has been deduced from a combination of electron microscopy, immunofluorescence 39 40 microscopy, and the genome-wide patterns of protein localisation determined by ChIP. However, the link between key meiotic protein complexes, chromosome conformation, and genomic 41 sequence remains uncharacterized. 42

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Chromosome conformation capture (3C) techniques generate maps of pairwise contact 44 frequencies that are snapshots of chromosome organisation. 3C methods were originally applied 45 to assay chromosome conformation in S. cerevisiae, including during meiosis⁹. Now they are 46 widely used across a range of organisms and cellular contexts to link 3D organisation directly with 47 genomic sequence¹⁰, revealing important roles of the Structural Maintenance of Chromosomes 48 (SMCs) cohesin and condensin in genomic organization^{11,12}, where they likely mediate 49 chromosome compaction via the process of loop extrusion¹³. Here we return to yeast meiosis to 50 interrogate genome-wide chromosome organisation by Hi-C, elucidate mechanisms of 51 chromosome assembly, and define the role of key meiotic chromosome components, including 52 cohesin and the SC. 53

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Starting with a synchronized G1 population we analysed timepoints encompassing DNA replication, meiotic prophase and both meiotic divisions (**Fig. 1a,b,c, Extended Data Fig. 1a,b,c**). In G1, we detect strong centromere clustering (**Fig. 1a,d**) and folding back of the arms at the centromeres (**Fig. 1a, Supplementary Fig. 2**), characteristic of a Rabl conformation^{9,14}. During meiosis, centromere clustering is transiently dissolved (3-5h, **Fig. 1a,d, Extended Data Fig. 1a**); this coincides with a global decrease in inter-chromosomal contact frequency at mid-prophase, reflecting chromosome individualisation. Subtelomeric clustering also decreases during meiotic

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prophase (Fig. 1a,d, Extended Data Fig. 1a), with no evidence for the transient telomeric
 bouquet conformation, consistent with prior microscopic analyses¹⁵.

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Entering meiosis, contact frequency versus distance, P(s), curves display a shoulder, consistent with the linear compaction of chromosome arms increasing due to *cis*-loop formation (2-4h, **Fig. 1e, Extended Data Fig. 1d**, e.g. as defined¹⁶; for review¹³). This change in P(s) is reminiscent of the SMC-dependent changes observed via Hi-C during mitosis across species^{17–21}. Compaction coincides with meiotic prophase I and the formation of the SC at pachytene, and is lost at later stages (**Fig. 1e, Extended Data Fig. 1d**).

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To study meiotic chromosome conformation in more detail, and to eliminate cell-to-cell 72 heterogeneity (Fig. 1b,c), we enriched for pachytene cells in subsequent experiments by 73 inactivating Ndt80, a transcription factor required for exit from meiotic prophase²². ndt80^Δ cells 74 entered meiosis synchronously, assessed by bulk DNA replication (Fig. 1f), but do not initiate 75 the first nuclear division²². Similar to the wild type prophase population (3-5h), but likely 76 accentuated by the increased homogeneity, Hi-C maps of pachytene-enriched cells displayed 77 total loss of centromere clustering (Extended Data Fig. 2) and dramatic chromosome arm 78 compaction (Fig. 1e). Analysing compaction in more detail, shorter chromosomes (Extended 79 Data Fig. 1e) and, in particular, shorter chromosome arms (Fig. 1h, Extended Data Fig. 1f), 80 displayed elevated contact frequency at short genomic separations, and an earlier shoulder, 81 apparently arising from distinct behavior of subtelomeric and subcentromeric regions (Fig. 1h, 82 **Extended Data Fig. 1g**). Moreover, distinct *P*(*s*) for chromosomes with different length arms 83 (Extended Data Fig. 1h) suggests that the centromere can insulate the process that leads to 84 differences between arms. In agreement with this, compaction is interrupted at centromeres in Hi-85 C maps (Fig. 1a, Extended Data Fig. 2b). 86

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Zooming in to consider within-arm organization revealed punctate grid-like Hi-C interactions between pairs of loci during prophase (**Fig. 2a**), particularly prominent in $ndt80\Delta$ (**Fig. 2a,b**). Such focal meiotic patterns are more prominent than reported previously²³—resembling peaks between CTCF sites²⁴ rather than topological domains^{25,26} detected in mammalian interphase Hi-C maps and likely arise from a heterogeneous mixture of 'transitive' interactions and 'skipping' of peak bases (**Fig. 2c**).

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Genomic regions underlying the punctate Hi-C interactions display a remarkable visual (Fig. 95 2a,b), and quantitative (Fig. 2d-g), correspondence with previously characterized sites of high 96 Rec8 occupancy²⁷. At pachytene, Rec8 sites display elevated cis/total contact frequencies (Fig. 97 2c), enriched contact frequency (Fig. 2e,f), and evidence of insulation (Fig. 2g)-features that 98 correlate with Rec8 occupancy measured by ChIP (Fig. 2a, lower). In wild type cells, Rec8-Rec8 99 interactions became visible in early prophase (2h), peaked at mid prophase (4h), and were 100 especially prominent in the homogenous *ndt80*∆ cell population (Fig. 2a-g, Extended Data Fig. 101 4b,c). Importantly, Rec8-Rec8 enrichments are strongest between adjacent sites, decrease 102

between non-adjacent sites with increasing genomic separation, and are absent in *trans* (Extended Data Fig. 4b,c). As for enrichments between CTCF sites in mammalian interphase²⁸,
 these observations argue that a cis-acting process generates such focal interactions in meiosis.

Rec8 is a central component of the meiotic chromosome axis³. Assaying a rec8^Δ mutant enabled 107 us to determine that Rec8 is absolutely required for the emergence of the grid-like Hi-C patterns 108 present in meiosis (Fig. 2a,b). Moreover, rec8 Δ cells completely lose the shoulder in P(s), 109 indicative of a dramatic loss of arm compaction (Fig. 2b, Extended Data Fig. 4a), similar to that 110 caused by depletion of SMCs in diverse contexts^{17,19,21,29-32}. Instead of assembling an axis of 111 loops, rec8Δ cells appear to be caught in a state with highly clustered telomeres (Extended Data 112 Fig. 4d, Extended Data Fig. 3), consistent with previous observations by microscopy^{33,34}. 113 Moreover, in rec8^Δ cells cis contact frequency is reduced (Fig. 2d), similar to G1 cells, and 114 cis/total no longer correlates with Rec8 occupancy. Instead, rec8∆ cis/total displays a decreasing 115 trend along chromosome arms, likely due to persistent telomere clustering (Extended Data Fig. 116 117 **4d**).

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To test how compaction and grid-like interaction patterns could jointly emerge in meiosis, we 119 developed polymer simulations (Fig. 3a, Methods) similar to those used to successfully describe 120 the assembly of TADs in mammalian interphase chromosomes¹³. Importantly, these simulations 121 employ the *cis*-acting process of loop extrusion, where extruders form progressively larger 122 chromatin loops, unless impeded by adjacent extruders or barrier elements (Fig. 3a). Extrusion 123 dynamics are controlled by parameters dictating the processivity (average loop size) and 124 separation (number of active extruders), as well as the strength of barriers (Methods). Because 125 the accumulation of Rec8 at ChIP-seq sites²⁷ concomitant with convergent transcription³⁵ is 126 indicative of barriers to extrusion²⁸, we positioned bi-directional barriers at Rec8 sites. 127

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Simulations were used to explore variations in loop extrusion dynamics to determine whether 129 specific parameter combinations are able to generate Hi-C maps that agree with experimental 130 observation (Fig. 3, Methods). Models with excellent fits were identified in which ~65% of the 131 genome is covered by extruded loops (Fig. 3b,c, Extended Data Fig. 5)—a far denser array than 132 present in S. cerevisiae mitosis²¹, but still less compact than human mitotic cells²⁰. Even though 133 extrusion can generate compaction independently of barriers (Fig. 3d), an intermediate barrier 134 strength is essential to match the grid-like patterns observed experimentally (Fig. 3b). Despite 135 the simplifying assumptions, simulated chromosomes displayed many features observed 136 experimentally: (i) chromosomes fold into a loose polymer brush^{3,36,37}, with a Rec8-rich core³ (Fig. 137 3f, Extended Data Fig. 5 a); (ii) a grid-like interaction pattern naturally emerges in simulated Hi-138 C maps (Fig. 3d); (iii) importantly, because loop extrusion is a *cis*-acting process, pairs of Rec8 139 sites at increasing separations naturally have lower contact frequency (Fig. 3e). 140

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Simulations also highlight the stochasticity of loop positions across the cell population, with most 142 barriers (73%) unoccupied by an extruder, and extruders paused with barrier elements on both 143 sides only a minority of the time (15%) in the best fitting models (Extended Data Fig. 6 c). 144 Because of this, the majority (65%) of extruded loops cross over Rec8 sites, consistent with an 145 average loop size roughly twice the average distance between Rec8 ChIP peaks (26 kb versus 146 12 kb, Extended Data Fig. 6 d), and remarkably consistent with estimates made using EM 147 (~20kb³⁶). Most strikingly—despite the prominence of Rec8-dependent grid-like features in the 148 experimental data (Fig. 2c)—our simulations indicate that Rec8 sites are not always occupied by 149 extruding cohesins and thus are present at the meiotic chromosome core in only a subset of cells, 150 as inferred previously³⁸. 151

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The range of loop extrusion parameters we explored encompasses the situation where Rec8 sites 153 always halt extrusion and cis-loops are formed between each consecutive Rec8 site. However, 154 simulations with these parameters have quantitatively poor fits with experimental maps (Fig. 3d-155 e, ii): the bend in P(s) comes too early to recapitulate experimental P(s), and Rec8-Rec8 contacts 156 are much too strong. The poor fit of such 'direct-bridging' simulations underscores the conclusion 157 that only a fraction of Rec8 sites are occupied in a given cell, and argues that cohesin-dependent 158 cis-loops must link regions that are not primary Rec8 binding sites in order to provide compaction 159 without making Rec8-Rec8 enrichments overly strong. 160

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A crucial prediction of our loop extrusion simulations is that depletion of extruders in meiosis would 162 lead to both decompaction and loss of the grid-like pattern of Hi-C interactions. When we repeated 163 our fitting procedure for rec8A, the best fits were for simulations with either no, or very few, 164 extruded loops (Extended Data Fig. 5e). The lack of compaction in these simulations (Extended 165 **Data Fig. 5a**) is consistent with previous EM showing decompacted chromatids in $rec8\Delta^3$. Such 166 joint consistency between Hi-C and imaging data further supports loop extrusion as a mechanism 167 underlying assembly of the cohesin-rich core and contributing to chromosomal compaction in 168 meiosis. Our simulations also open the possibility that overly-shortened axes observed upon 169 Wapl^{39,40} and Pds5⁴¹ depletion may reflect heightened extruder processivity⁴² upon which 170 shortened SCs are assembled, and predict that such perturbations would cause a rightward shift 171 in the P(s) shoulder measured via Hi-C (Extended Data Fig. 5c). 172

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To investigate how homologue synapsis affects chromosome conformation, we assayed 174 pachytene cells in the absence of Zip1, the transverse filament of the SC⁴, and Hop1, an axial 175 element required for Zip1 loading⁶ (Fig. 4a,b). Both $zip1\Delta$ and $hop1\Delta$ Hi-C maps retained the 176 Rec8-dependent punctate interactions (Fig. 4b, Extended Data Fig. 2b,c), and displayed 177 compaction relative to G1 or rec8 Δ , but with the P(s) shoulder shifted left relative to ndt80 Δ (Fig. 178 **4c**). Attempts to model the known $zip1\Delta$ and $hop1\Delta$ defects in chromosome synapsis simply by 179 removing interhomologue crosslinks from best-fitting ndt80Δ simulations did not recapitulate the 180 P(s) shift observed experimentally (Extended Data Fig. 5 f), consistent with the suggestion that 181 interhomologue contacts make only a minor contribution within meiotic Hi-C maps²³. Instead, 182 best-fitting simulations had shifts towards slightly lower processivity and larger separation, 183 consistent with less axial compaction relative to the $ndt80\Delta$ control (Fig. 4e). Interestingly, 184 subtelomeric regions no longer displayed a distinct P(s) in $zip1\Delta$ and $hop1\Delta$ (Fig. 4d), suggesting 185 that chromosome compaction at chromosome termini is regulated differentially. 186

187 Discussion

Our analysis of meiotic chromosome organisation via Hi-C reconciles the function and localisation of factors thought to shape meiotic chromosomes with their 3D organisation, revealing the emergence of a punctate grid of interactions concomitant with initial stages of meiotic chromosome compaction. Crucially, we formally demonstrate the link between preferential positioning of meiotic cohesin along the genome^{27,35} and the inference that these loci come into close proximity based on the localization of Rec8 to the chromosomal axes³. Remarkably, the

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punctate cohesin-dependent interactions in yeast meiosis emerge despite the absence of CTCF
 in this organism; this challenges previous models where focal Hi-C peaks are strictly dependent
 on CTCF^{24,31,43}, and indicates that alternative mechanisms of loop positioning must exist.

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Notably, whilst much less prominent, locus-specific folding is evident in equivalent high-resolution 198 Hi-C maps of mitotic cells (Extended Data Fig. 7)-something that was hidden within lower-199 resolution analyses^{19,21}. The correspondence between cohesin positioning and convergent 200 transcription in both meiosis³⁵ and mitosis^{44,45} argues that transcription may be a fundamental and 201 ubiquitous process capable of broadly patterning locus-specific chromosome organisation by 202 modulation of cohesin dynamics⁴⁶. Indeed, the stronger meiotic patterns are particularly 203 reminiscent of the extended grid-like Hi-C patterns observed in interphase mammalian cells upon 204 depletion of the cohesion unloader, Wapl^{31,47}, wherein "vermicelli"-like chromatids arise with a 205 cohesin-rich backbone⁴⁸, emphasising the influence of cohesin dynamics on loop extrusion. We 206 favour the view that transcription acts as a barrier to cohesin-dependent loop extrusion, rather 207 than as a motive force as previously proposed^{35,49,50}, consistent with transcription-independent 208 compaction by cohesin in mammalian interphase⁴⁷ and direct observation of extrusion by the 209 related SMC condensin in vitro⁵¹. 210

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Consistent with the idea that large DNA-protein complexes, like a kinetochore, can impede 212 extrusion, we observe a paucity of Rec8-dependent loops spanning the centromere. 213 Nevertheless, whether barriers arise directly from Pol2 binding, or indirectly via other axis 214 proteins, remains to be determined. Indeed, the reason for the why loops are more strictly 215 positioned in meiosis compared to mitosis is intriguing. However, our observations enable us to 216 rule out the axial element, Hop1, the SC lateral element, Zip1, and the process of homologous 217 recombination mediated by Spo11, Sae2, and Dmc1 (unpub. obs.) as important for the generation 218 of such patterns. 219

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Our simulations also reveal a nuanced picture of meiotic chromosome assembly: loops are, on average, larger than the inter-Rec8 peak distance, and more than half of the loop bases are not associated with preferred sites of Rec8 binding. Moreover, it is likely that loop sizes and positions vary widely from one cell to another, making classifications of genomic regions as 'axis' or 'loop' a great oversimplification. The agreement between our simulations and experimental data furthers the case for loop extrusion as a general mechanism^{19–21,23,28,52–55} that is flexibly employed and regulated in interphase, mitosis, and meiosis.

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Our results also reveal how the interplay between the synapsis components, Hop1 and Zip1, 229 influences chromosome morphology. That Hop1 and Zip1 are both required to increase 230 chromosome compaction at pachytene likely points at their joint role in promoting synapsis^{4,6}, and 231 supports the view that synapsis itself modulates axial compaction. Interestingly, whilst Zip1 binds 232 largely uniformly along the arms of pachytene chromosomes⁵⁶, subtelomeres and short 233 chromosomes display an increase in short-range contacts and an earlier shoulder in P(s), 234 consistent with smaller loops or less compression of spacers between loops in these regions, and 235 therefore less axial compaction. Because such differences correlate with disproportionate 236 retention of Hop1 in these regions⁵⁶ and diminished efficiency of synapsis⁵⁷, it is possible that 237 Hop1 impedes the pathway whereby Zip1 imposes additional compaction upon synapsis. 238 Nevertheless, it is unclear whether Zip1 mediates this effect by modifying loop extrusion 239 dynamics, or via a distinct process of axial compression, as has been argued for higher eukaryote 240 mitotic chromosome compaction²⁰. Given the influence that chromosome structure has over so 241 many aspects of meiosis, teasing apart these mechanisms is of great future interest. 242 243

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SAS and MJN planned the study, performed wet-lab work and data analysis. GF and KSP developed polymer simulations and performed data analysis. SAS, GF, JB, KSP and MJN discussed results and wrote the manuscript. We thank Tim Cooper for deploying the HiC Pro installation, Scott Keeney and Franz Klein for sharing *S. cerevisiae* strains, Svetlana Lyalina for assistance with QB3 GPUs, Anton Goloborodko for suggesting the use of *looplib*, and Nezar Abdennur for feedback.

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²⁵² Data & Code Availability

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All processed Hi-C matrices are publicly accessible via the interactive HiGlass viewer⁵⁸. Live web links are provided with each figure. Raw sequence reads will be made publicly accessible via the SRA repository upon final publication. Code for data analysis is either already publicly available online or will be made available on request or via Github upon final publication.

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²⁶⁵ Figure Legends

Figure 1. Chromosome conformation during yeast meiosis.

a. Cells were collected during meiosis at indicated timepoints and analysed by Hi-C. At 0h the cells are in G1. Representative Hi-C contact maps of chromosomes 6, 11, and 7 plotted at 5 kb resolution. Centromeres, telomeres and arm fold-back at the centromere are indicated by blue, red and grey arrows, respectively, and axial compaction by the width of the main diagonal relative

to the fixed-width black clamp. For interactive HiGlass⁵⁸ views see:
 http://higlass.pollard.gladstone.org/app/?config=Z5iwKpjzQpePCXXyvuYGeQ

b. Meiotic entry assessed by FACS; at 4h, the majority of cells show a 4C peak indicating completion of DNA replication.

c. Meiotic progression was monitored by quantification of nuclear divisions determined by DAPI

staining. Around 4h, cells start to undergo meiotic divisions I and II. The majority of cells undergo
 meiotic divisions between 4h and 8h, indicating the degree of heterogeneity within the cell
 population.

d. *Upper panels*: Average *trans* centromere-centromere contact maps. *Lower panels*: *trans* telomere-telomere contact maps. *Right*: ratio of *cis* to total contact frequency.

e. Intra-arm contact probability versus genomic distance, P(s), indicating the emergence (*left*) and

disappearance (*right*) of chromosome arm compaction during meiosis. Shaded area bounded above and below by the two $ndt80\Delta$ 8h replicates.

f. Meiosis was induced in $ndt80\Delta$ cells for 8h and meiotic entry was checked by monitoring DNA

replication by FACS.

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- 286 g. ndt80Δ cells were grown for 8h in sporulation media and analysed by Hi-C (left). Log2 ratio of
- ndt80Δ cells 8h over G1 (right). Centromeres and telomeres are indicated by blue and red arrows,
- respectively, and axial compaction by a black clamp.
- h. Left: Contact probability of individual chromosome arms stratified by length. Right: Contact
 probability stratified by the distance from the telomere.

291 Figure 2. Emergence of a Rec8-dependent grid of punctate interactions in meiosis

- **a.** Hi-C contact maps of chromosome 11 for the indicated genotypes plotted at 2 kb bin resolution, showing near-diagonal interactions. Wild type timepoints as in **Fig. 1a. Lower panels**: log2(insulation); cis/total ratio, Rec8 ChIP-seq²⁷, all binned at 2kb. Insulation and cis/total calculated from *ndt80* Δ maps. Positions of Rec8 sites indicated as green circles. Genome-wide cis/total (Spearman's R=0.62, P<1e-10) and insulation (R= -0.23, P<1e-10, insulation window = 20 kb) profiles are correlated with Rec8 occupancy.
- **b.** Zoom-in into contact maps on chromosome 11 (0-200kb) of wt-4h and $ndt80\Delta$ (top) and $rec8\Delta$
- (bottom left). Contact probability versus genomic distance, P(s), for G1(*ndt80* Δ -0*h*) and *ndt80* Δ
- and $rec8\Delta$ (bottom right). Data shown is the average (n=2) except for wt-4h. Rec8 peak sites called from ChIP-seq data²⁷ are indicated in green. For an interactive view see: <u>http://higlass.pollard.gladstone.org/app/?config=Twrh61jGT4SlxotaguTIJg</u>
- c. Simplified illustration of how a grid of peaks on a Hi-C map can emerge between Rec8 sites
 either by transitive contacts between adjacent loops, or by loops that skip over adjacent sites.
- 305 Experimentally observed grids extend much further than separation=2 (Extended Data Fig. 4c)
- d. Cis/total ratios for Rec8 (green) and nonRec8 (grey) sites for indicated datasets.
- e. Contact frequency versus distance between Rec8-Rec8 sites (green), Rec8-nonRec8 sites
 (light green) and nonRec8-nonRec8 sites (green).
- f. Log2 ratio of contact frequency between adjacent Rec8-sites (separation=1) compared to
 average cis interactions.
- **g.** Log2 ratio of contact frequency centered at Rec8 sites compared to average cis interactions.
- In *ndt80*Δ, Rec8 sites show: elevated cis/total frequency (0.85 versus 0.77), elevated pairwise
- contact frequency (~2-fold at 20 kb), and mild insulation (f-g). These distinctions are similar in
- wild type pachytene (4h) yet absent in G1 ($ndt80\Delta$ -0h) or in $rec8\Delta$.

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Figure 3. Modelling meiotic chromosome compaction.

- a. In simulations, yeast chr13 was represented as a polymer fiber confined to the nucleus subject 316 to additional meiosis-specific constraints. These include: extruded loops, sister crosslinks, and 317 homolog crosslinks (Methods). Barriers to extruded loops were placed at Rec8 sites²⁷. We 318 imposed inter-sister and inter-homologue crosslinks at sites of extruded loop bases in order to 319 approximate the paired arrangement of homologues at pachytene (Extended Data Fig. 6). For 320 each set of extruded loop parameters (processivity, separation, and barrier strength), 321 322 conformations were collected and used to generate simulated contact maps. Roughly, processivity dictates the size of an extruded loop unimpeded by collisions, separation controls the 323 number of active extruders on the chromosome, and barrier strength controls the probability that 324 an extruder gets paused when attempting to step past a barrier. Simulated and experimental 325 contact maps were then compared via the combined average fold discrepancy between P(s)326 curves for Rec8-Rec8, Rec8-non, and non-non bin pairs at 2kb resolution. 327
- b. Goodness-of-fit for indicated barrier strengths over coarse grids of processivity and separation
 demonstrate that intermediate barrier strengths are required to agree with experimental *ndt80*Δ
 Hi-C maps.
- **c.** Goodness-of-fit for a fine grid of processivity versus separation at barrier strength 0.95. Best-
- fitting models had separation ~32kb and processivity ~76kb, corresponding to ~60% coverage of the genome by extruded loops of average length 26kb.
- **d.** From *left* to *right*: contact maps for chr13 for *ndt80*Δ, and simulations with (i) best-fitting parameters, (ii) relatively stable loops between neighboring Rec8 sites, and (iii) no barriers.
- **e.** P(s) split by Rec8-Rec8, Rec8-non, and non-non, as in Fig. 2D.
- **f.** Conformations for best-fitting simulations, which highlight: (*left*) one chromatid colored from start (red) to end (blue); (*right*) extruders (yellow), extrusion barriers (red), and extruders paused at barriers (orange).

³⁴⁰ Figure 4. Hop1 and Zip1-dependent compaction of Rec8-dependent loops.

- **a.** Top: Hi-C maps for hop 1Δ and zip 1Δ (plotted as in Fig. 1a). Bottom: Log2 ratio of hop 1Δ over 341 For ndt80∆ (as in Fig. **1g)**. interactive views of the full genome, 342 see http://higlass.pollard.gladstone.org/app/?config=TTBGu5DDR0SHAa09zrjTXA 343
- **b.** Hi-C contact maps of chromosome 11 for $hop1\Delta$ and $zip1\Delta$ plotted at 2kb bin resolution,
- showing near-diagonal interactions, as in **Fig. 2a**.

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c. Contact probability versus genomic distance for G1, *ndt80*Δ, *hop1*Δ, *zip1*Δ. Shaded area

bounded above and below by *ndt80* Δ replicas. Average between two replicas for *zip1* Δ and one

sample for G1 and $hop1\Delta$ are shown.

d. Contact probability over genomic distance averaged over all chromosome arms stratified by
 distance from the telomere.

e. Goodness-of-fit for simulations without homolog crosslinks with a fine grid of processivity versus separation at barrier strength 0.95 $zip1\Delta$ and $hop1\Delta$.

f. Model of meiotic chromosome compaction: Rec8-dependent loop formation leads to initial chromosome arm compaction and emergence of a grid-like pattern of Hi-C interactions that jointly agrees with a mechanism of loop extrusion including barrier elements. We suggest that transcription could impose such barriers. Hop1 and Zip1 are dispensable for this step, but are required for synapsis, where additional compaction occurs differentially along chromosome arms.

359 Extended Data Fig. 1

Temporal and chromosome length-specific analysis of meiotic chromatin conformation

a-d. Results from a replicate timecourse, collected and characterized independently of the timecourse in Fig. 1.

- 362 **a.** Hi-C maps, plotted as in Fig. 1a.
- 363 **b.** FACS as in Fig. 1b.
- 364 **c.** DAPI as in Fig. 1c.
- 365 **d.** *P*(*s*) as in Fig. 1e.
- **e.** P(s) for chromosomes stratified by size for *ndt80* Δ -0h, *ndt80* Δ -8h. Short chromosomes display
- relatively elevated P(s) at short distances, and an earlier shoulder.

368 f. Left: P(s) for individual chromosome arms, stratified by size for wt-4h. Short arms display

- relatively elevated *P*(s) at short distances, and an earlier roll-over. *Right*: Intra-arm *P*(s) stratified
- by the distance from the telomere for wt-4h, averaged across all chromosomes. Telomere-
- proximal regions display elevated P(s) at short distances.
- **g.** Intra-arm P(s) stratified by the distance from the centromere for G1 (*ndt80* Δ -0h), *wt*-4h, *ndt80* Δ -
- ³⁷³ 8h, averaged across all chromosomes.
- **h.** Contact probability of single chromosome arms for *ndt80*Δ-8h.

375 Extended Data Fig. 2

Aggregate analysis of centromeric interactions in meiosis

a. Average *trans* centromere-centromere contact maps for indicated data sets.

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- **b.** Average *cis* centromere-centromere contact maps for indicated data sets. Note the loss of the
- folding back in meiosis, and how the intra-arm enrichment is insulated at centromeres in meiosis.

379 Extended Data Fig. 3

Aggregate analysis of telomeric interactions in meiosis

- **a.** Average *trans* telomere-telomere contact maps for indicated datasets.
- **b.** Average telomere-telomere contact maps between the two telomeres of the same chromosome.
- 383 c. Average contact map around each telomere in *cis*.

384 Extended Data Fig. 4

Preferred sites of Rec8 occupancy define sites of locus-specific interaction

- a. Left: Hi-C contact maps of $rec8\Delta$ ndt80 Δ . Chromosomes 6, 11 and 7 are shown as
- representatives for the whole genome. *Right*: Log2 Hi-C ratio maps of $rec8\Delta$ ndt80 Δ / ndt80 Δ .
- 387 Plotted as in Fig. 1g.
- b. Log2 observed over expected contact frequency at Rec8-Rec8 peak pairs as a function of
 separation across datasets.
- c. Log2 observed over expected contact frequency +/-8kb around Rec8-Rec8 peak pairs at the
 indicated separations.
- ³⁹² Together, **b-c** demonstrate that Rec8-Rec8 enrichments are strongest between adjacent sites,
- decrease between non-adjacent sites with increasing genomic separation, and are absent in
- trans. Equally important, these meiotic features are lost in $rec8\Delta$. As for mammalian interphase,
- this observation in meiosis argues for a *cis*-acting process underlying the formation of focal interactions between Rec8 sites.
- 397 **d.** cis/total as a function of distance along the chromosomal arm, Rec8 sites marked in green.
- 398

399 Extended Data Fig. 5

- Polymer simulations of loop extrusion reveal best fitting parameters and conformations
- **a.** Representative conformation for the indicated parameter sets. As in Fig. 3F, one chromatid
- from a homologous quartet of chromatids colored from start to end according to the spectrum;
- 402 other three colored in grey.
- **b.** For the same three conformations, positions of Rec8 sites indicated with red spheres, positions
- ⁴⁰⁴ of extruded loop bases in yellow, and extruders overlapping a Rec8 site in orange. Note the stable
- loops between neighboring Rec8 sites creates a very elongated chromatid (ii). Also note the

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⁴⁰⁶ majority of Rec8 sites are unoccupied in (iii), despite the self-assembly of two axial cores and a ⁴⁰⁷ strong brush. Finally, note very dispersed chromosomes in (iv), consistent with EM³ for *rec8* Δ .

408 **c.** Contact frequency versus distance, P(s), for indicated simulations. Note that the loss of the

shoulder in *P*(*s*) in the case of full extruder depletion mirrors the difference between experimental

 $ndt80\Delta$ and $rec8\Delta$ Hi-C maps. Simulations with increased processivity predict that P(s) would shift

- rightward if unloading was impaired, as could happen in $wapl\Delta$. Conversely, if unloading was enhanced, simulations with decreased processivity indicate a leftward shift in P(s), until the
- 413 absence of extruders.
- d. Goodness-of-fit for a fine grid of processivity versus separation at barrier strength 0.90. The
- ⁴¹⁵ best-fit occurs at similar processivity and separation as for barrier strength 0.95 shown in Fig. 3c,
- 416 but with slightly lower goodness-of-fit.
- **e.** Goodness-of-fit to $rec8\Delta$ data for simulations with the indicated barrier strengths (in grey: 0.00,

0.75, 0.90, 0.95, 0.99, 1.00) over coarse grids of processivity and separation demonstrates that
the best fits have few if any extruded loops, regardless of barrier strength.

- **f.** P(s) curves for simulations with sisters and homologs with the best-fitting parameters for *ndt80* Δ -*8h* maps compared to P(s) for simulations with sisters only show that simply removing
- homolog tethering does not recapitulate the sort of shifted P(s) seen experimentally in $zip1\Delta$ Hi-C.
- 424

425 Extended Data Fig. 6

Polymer simulations of loop extrusion elucidate meiotic barrier strength

a. Simulated contact maps for the indicated region of chr13 for: (i) best-fitting simulations, (ii)
 simulations with relatively stable loops between neighboring Rec8 sites (barrier strength=1 and
 high processivity), and (iii) no barriers, as in Fig. 2d.

- b. Simulated ChIP-seq profiles for the indicated region of chr13. Best-fitting simulations (i) display
 occupancy well below 100% at Rec8 sites. Simulations with stable loops (ii) display highly
 occupied Rec8 sites. Simulations without barriers (iii) have homogenous Rec8 occupancy across
 the genome.
- **c.** Positions of extruded loops (arcs) sister crosslinks (solid black lines) and homolog crosslinks
- (dashed lines) for four chromatids in two separate cells, showing how the simulated Hi-C maps
- and ChIP-seq profiles emerge from the stochastic positioning of extruded loops from cell-to-cell.
- 436 For statistics, see **Supplementary Table S1**.
- 437 **d.** Histogram of extruded loop lengths for indicated parameters (i, ii, iii).
- 438

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439 Extended Data Fig. 7

Cohesin and transcription patterns loops in meiosis and mitosis

a. Hi-C contact maps of chromosome 11 for meiotic ($ndt80\Delta$, pachytene - top) and mitotic (wild type, nocodazole arrest - bottom) plotted at 2 kb bin resolution, showing near-diagonal interactions. Data shown is the average (n=2).

- **b.** Zoom-in into contact maps on chromosome 11 (0-200kb) of *ndt80*Δ (top) and mitotic (*bottom*).
- Data shown is the average (n=2). Rec8 peak sites called from ChIP-seq data²⁷ are indicated in
- green. Arrowheads indicate sites of prominent focal interaction.

446

447 Tables

448

449 Table 1. S. cerevisiae strains used in this study

Strain name	genotype
MJ6	ho::LYS2/", lys2/", ura3/", arg4-nsp/", leu2::hisG/", his4X::LEU2/", nuc1::LEU2/"
SSY14	ho::LYS2/", lys2/", ura3/", arg4-nsp/", leu2::hisG/", his4X::LEU2/", nuc1::LEU2/", ndt80∆::LEU2/"
SSY20	ho::LYS2/", lys2/", ura3/", arg4-nsp/", leu2::hisG/", rec8∆::KanMX4/", ndt80∆::LEU2/"
SSY25	ho::LYS2/", lys2/", ura3/', arg4-nsp/", leu2::hisG/", his4X::LEU2/", nuc1::LEU2/", zip1::LEU2/", ndt80∆::LEU2/"
SSY49	ho::LYS2/", lys2/", ura3/", arg4-nsp/", leu2::hisG/", nuc1::LEU2/", his4X::LEU2/", hop1::LEU2/", ndt80∆::LEU2/"
SSY58	ho::hisG/", lys2/", ura3/", leu2::hisG/", nuc1::LEU2/", arg4-nsp/", rec8::KanMX/", ndt80∆::LEU2/"

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450

451 Table 2. Hi-C Libraries

Name	mutations	Sample name	valid pairs (M)
Main figures			
wt-0h/G1		HiC_MJ6_wt_2A_0h	14.5
wt-2h		HiC_MJ6_wt_2A1_2h	27.6
wt-3h		HiC_MJ6_wt_2A_3h	24.1
wt-4h		HiC_MJ6_wt_2A_4h	28
wt-5h		HiC_MJ6_wt_2A1_5h	27.6
wt-6h		HiC_MJ6_wt_2A1_6h	27.6
wt-8h		HiC_MJ6_wt_2A3_8h	19
rec8∆	rec8∆ ndt80∆	average	
<i>rec8</i> ∆ replica 1	rec8∆ ndt80∆	HiC_SSY20_ndt80Drec8D_1A2_8h	39.3
<i>rec8</i> ∆ replica 2	rec8∆ ndt80∆	HiC_SSY58_ndt80Drec8D_2A_8h	20.2
ndt80∆	ndt80∆	average, 8h	
G1	ndt80∆	HiC_SSY14_ndt80D_1A2_0h	36
<i>ndt80</i> ∆-4h	ndt80∆	HiC_SSY14_ndt80D_1A_4h	11.9
<i>ndt80</i> ∆ replica 1	ndt80∆	HiC_SSY14_ndt80D_1A1_8h	22.9
<i>ndt80</i> ∆ replica 2	ndt80∆	HiC_SSY14_ndt80D_2A2_8h	37
zip1∆	$zip1\Delta$ ndt80 Δ	average	
<i>zip1</i> ∆ replica 1	$zip1\Delta$ ndt80 Δ	HiC_SSY25_ndt80Dzip1D_1B2_8h	22.7
<i>zip1</i> Δ replica 2	zip1 Δ ndt80 Δ	HiC_SSY25_ndt80Dzip1D_2A_8h	28.6

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hop1∆	hop1 Δ ndt80 Δ	hop1 ndt80	
<i>hop1</i> ∆ replica1	hop1 Δ ndt80 Δ	HiC_SSY49_ndt80Dhop1D_1A_8h	32.8
Supplementary figures			
wt-2h		HiC_MJ6_wt_3A_2h	22.5
wt-3h		HiC_MJ6_wt_3A_3h	19.8
wt-4h		HiC_MJ6_wt_3A_4h	16.7
wt-6h		HiC_MJ6_wt_3A_6h	37.6

452

453 Table 3. Overview of proteins described in this study

Protein	Description
Ndt80	Transcription factor required for exit from pachytene
Rec8	Meiosis-specific kleisin subunit of cohesin
Hop1	Axial element of the synaptonemal complex
Zip1	Transverse filament of the synaptonemal complex

454

455

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457 Methods

458 Yeast strains and cell culture growth.

459 Strains used in this study were derived from SK1 and are listed in Table 1.

460 Monitoring meiotic progression by flow cytometry and quantification of nuclear divisions

Cells were fixed in 70% EtOH, digested with 1 mg/ml RNAse (10 mM Tris-HCl pH 8.0, 15 mM 461 NaCl, 10 mM EDTA pH 8.0) for 2 h at 37 °C, 800 rpm and subsequently treated with 1 mg/ml 462 Proteinase K in 50 mM Tris-HCl pH 8.0 at 50 °C, 800 rpm for 30 min for analysis by FACS. Cells 463 were then washed in 50 mM Tris-HCl pH 8.0 and stained in the same buffer with 1 uM Sytox 464 green overnight in the fridge. FACS profiles were plotted with R using the library hwglabr2 465 (https://github.com/hochwagenlab/hwglabr2). Fixed cells were also used for quantification of 466 nuclear divisions by spreading onto a microscope slide, mounting with Fluoroshield containing 467 DAPI followed by analysis with a Zeiss Scope.A1 microscope. 468

469 Hi-C library preparation

The Hi-C protocol used was amended from⁵⁹ by ~5-fold reduction in all materials and volumes. 470 Briefly, S. cerevisiae diploid cells were synchronised in G1 by growth at 30 °C for ~15 h in 30 ml 471 YPA (1% Yeast extract, 2% Peptone, 1% K-acetate) to OD600 of ~4, harvested, washed, and 472 resuspended in prewarmed sporulation medium (2% KAc with 0.2x nutritional supplements) 473 before fixing 5 ml aliquots (20-30 ODs) of relevant timepoints with formaldehyde at 3% final 474 concentration for 20 min at 30 °C, 250 rpm, then guenched by incubating with a final concentration 475 of 0.35 M Glycine (2x the volume of Formaldehyde added) for an additional 5 minutes. Cells were 476 washed with water split into two samples and stored at -80°C ready for library preparation. Cells 477 were thawed, washed in spheroplasting buffer (SB, 1 M Sorbitol, 50 mM Tris pH 7.5) and digested 478 with 100 ug/ml 100T Zymolyase in SB containing 1% beta-Mercaptoethanol for 15-20 min at 35 479 °C. Cells were washed in restriction enzyme buffer, chromatin was solubilised by adding SDS to 480 0.1% and incubating at 65 °C for 10 minutes. Excess SDS was quenched by addition of Triton 481 X100 to 1%, and chromatin was incubated with 2.07U/ul of DpnII overnight at 37 °C. DNA ends 482 were filled in with nucleotides, substituting dCTP for biotin-14-dCTP using Klenow fragment DNA 483 polymerase I at 37 °C for 2 h followed by addition of SDS to 1.5% and incubation at 65 °C for 20 484 min to inactivate Klenow and further solubilise the chromatin. The sample volume was diluted 15-485

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fold, crosslinked DNA ends ligated at 16 °C for 8 h using 0.024U/ul of T4 DNA ligase, and 486 crosslinks reversed by overnight incubation at 65 °C in the presence of proteinase K. DNA was 487 precipitated with ethanol, dissolved in TE and passed through an Amicon 30 kDa column. DNA 488 was further purified by phenol:chloroform:isoamylalcohol extraction and precipitated again before 489 treating with RNaseA at 37 °C for 1 h. Biotin was removed from unligated ends by incubation with 490 T4 DNA polymerase at 20°C for 4 h and at 75 °C for 20 min for inactivation of the enzyme. DNA 491 was subsequently fragmented using a Covaris M220 (Duty factor 20%, 200 cycles/burst, 350s, 492 20 °C), and DNA ends were repaired and A-tailed using T4 DNA polymerase, T4 Polynucleotide 493 Kinase and Klenow fragment DNA polymerase I before isolating fragments of 100-250 bp using 494 a Blue Pippin (Sage). Biotinylated fragments were enriched using streptavidin magnetic beads 495 (C1) and NextFlex (Bioo Scientific) barcoded adapters were ligated while the DNA was on the 496 beads. Resulting libraries were minimally amplified by PCR and sequenced using paired end 42 497 bp reads on a NextSeq500 (Ilumina; Brighton Genomics). 498

499 Hi-C data processing and analysis

Hi-C sparse matrices were generated at varying spatial resolutions using the Hi-C-pro pipeline⁶⁰, 500 using a customised S288c reference genome (SK1Mod, in which high confidence SK1-specific 501 polymorphisms were inserted in order to improve read alignment rates; manuscript in preparation) 502 and plotted using R Studio (version 1.0.44) after correcting for read depth differences between 503 samples. Raw read statistics are presented in Table 2. Repeat biological samples gave broadly 504 similar matrices and, unless indicated otherwise, were averaged to improve their expected 505 guantitative accuracy. As visual inspection indicated a number of potential translocations in the 506 SK1 strain as compared with the S288c reference genome, for conservative downstream 507 analyses, additional bins were masked if they contained potential translocations. Such bins were 508 identified if they either had values in trans at the level of the median of the third diagonal in cis, or 509 the maximum value in trans exceeded the maximum value in cis for SSY14 for bins displaying 510 these properties in either ndt80D-0h or in ndt80D-8h and for MJ6 in wt-0h or wt-4h. chr1 was 511 excluded from downstream analysis as few informative bins remained after filtering potential 512 translocations. 513

514

Average maps centered at centromeres and telomeres were calculated as in⁶¹, ensuring that collected patches for average centromere maps did not extend inter-chromosomally, and collected patches for average telomere maps did not extend beyond centromeres or interchromosomally. Contact frequency versus distance curves, P(s), were calculated from 2 kb

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binned maps, with logarithmically-spaced bins in s (numutils.logbins, https://bitbucket.org / 519 mirnylab/mirnylib, start =2, end = max(binned arm lengths), N=50), and restricting the calculation 520 to bin pairs within chromosomal arms and excluding bins less than 20 kb from centromeres or 521 telomere (as in⁶¹), and normalized to the average value at 4 kb. P(s) stratified by distance to 522 telomeres was calculated using the combined distance to telomeres for each bin-pair (as in⁶²), 523 and excluded bins-pairs where one bin was closer to a centromere than telomere along that arm. 524 Distance to centromeres, and P(s) stratified by this distance, was calculated similarly. Log2 525 insulation profiles were calculated using a sliding diamond window (as in⁶³) with a +/-20 kb (+/-10 526 bins) extent; as in⁴³ downstream analyses were restricted to when there were zero or one filtered 527 bins in the sliding window. To calculate histograms of cis/total (Fig. 2D), bins were defined as 528 either Rec8 or non-Rec8. To calculate P(s) split by Rec8 bin-pair status, each bin-pair (i.e. entry 529 of the heatmap) was assigned as either Rec8-Rec8, Rec8-nonRec8, or non-non (e.g. Fig. 2E). 530 P(s) was then aggregated separately across chromosomes for these three categories, similar to 531 calculation of P(s) within and between TADs²⁸. Average log2 observed/expected maps were 532 calculated by first dividing by intra-arm P(s) and then averaging together appropriate patches of 533 Hi-C maps. Correlations between Rec8 occupancy from²⁷ and insulation or cis/total profiles 534 excluded chromosome 12 because the rDNA locus greatly alters the insulation profile within the 535 right arm of the chromosome. 536

537 **Polymer simulations**

Meiotic loop extrusion simulations begin with a generic polymer representation of the yeast 538 chromatin fiber similar to that used in previous models of yeast mitotic chromosomes²¹, where 539 each 20 nm monomer represents 640 bp (~4 nucleosomes). We simulated the chromatin fiber 540 with excluded volume interactions and without topological constraints, using Langevin dynamics 541 in OpenMM, as in^{64,65}. Importantly, meiotic simulations remove the geometric constraints specific 542 to the Rabl conformation^{66,67} because this is not visible in meiotic pachytene *ndt80* Hi-C maps. 543 As our focus was to characterize the grids of intra-chromosomal interactions, we considered a 544 system with multiple copies of chromosome 13, equivalent to four copies of the haploid genome 545 in terms of total genomic content (4 x 13 copies of chromosome 13), to enable efficient 546 computational averaging of simulated Hi-C maps. Extruded loops were generated according to 547 parameters that describe the dynamics of loop extruders, using the simulation engine described 548 in⁶⁸: extruder separation, extruder processivity, chromatin fiber relaxation time relative to extruder 549 velocity, and barrier strength. Because yeast chromosomes are short compared to higher 550 eukaryote chromosomes, relaxation time is relatively rapid and we focused on separation, 551

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processivity, and barrier strength. At every given timepoint an extruded loop is realized as a bond
 between monomers at the two bases of the loop (see ./src/examples/loopExtrusion in
 https://bitbucket.org/mirrylab/openmm-polymer/).

555

Upon encountering a barrier, a loop extruder is paused with probability according to the barrier 556 strength; barrier strength =1 indicates an impermeable barrier, barrier strength =0 indicates no 557 impediment to extrusion. We assume loop extrusion occurs independently on each chromatid, 558 and simulate loop extrusion dynamics on a 1D lattice (as in²⁸) where the number of lattice sites 559 equals the total number of monomers (75,140). Bi-directional barriers were placed at monomers 560 with positions corresponding to Rec8 ChIP-seq sites²⁷, and pause extruders according the barrier 561 strength parameter. We assume a uniform birth probability, constant death probability, and that 562 all barriers have an equal strength; as additional data becomes available, these assumptions can 563 be relaxed and more detailed models can be built. 564

565

We investigated scenarios where chromatids are then either left individualized (52 copies), 566 crosslinked to sisters (26 pairs), or additionally paired with homologs (13 pairs-of-pairs). For 567 simulations with sister crosslinks, these were added (following⁵⁴) when extruded loop bases were 568 present at cognate positions ±30 monomers (~20kb) on both chromatids (distance=20nm); 569 homolog crosslinks were added similarly when sister crosslinks were present on both chromatids 570 (distance=100nm); centromeres and telomeres were always paired, and both presented 571 impermeable (strength=1) boundaries to extruders. To avoid introducing pseudo-knots, if 572 extruded loops were nested only the outer cohesins were considered as possible bases for sister 573 crosslinks, sister crosslinks were only allowed between the same side of loop bases (i.e. left-to-574 left arm or right-to-right arm), and sister crosslinks were only added between bases at the 575 reciprocal minimum distance. 576

577

For calculation of simulated Hi-C maps, contacts were recorded from conformations of the full 578 system, which includes intra- and inter-sister, and interhomologue contacts. Because 579 experimental Hi-C here does not distinguish either sisters or homologs, contacts were then 580 aggregated into one simulated map. For each model and parameter set we investigated, we 581 collected an ensemble of conformations, generated simulated chr13 Hi-C maps, and compared 582 their features and P(s) with those from experimental Hi-C maps. Each simulated chr13 map 583 represented an average over 5400 conformations. P(s) for chr13 was calculated from 2kb binned 584 simulated maps exactly as for experimental maps. Maps of goodness-of-fit between simulations 585

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586	and experimental data (e.g. Fig. 3b,c) were computed as the geometric standard deviation of the
587	ratio of simulated to experimental $P(s)$ combined across $P^{Rec8-Rec8}(s)$, $P^{Rec8-non}(s)$, and $P^{non-non}(s)$,
588	as was previously done for P(s) within TADs of multiple sizes and between TADs ²⁸ , for s from
589	10kb to 300kb. This measure reflects the typical fold-deviation for $P(s)$.

590

Simulated ChIP-seq profiles (Extended Data Fig. 6b) for Rec8 were generated by aggregating the position of extruded loop bases (two per extruded loop) across conformations. Statistics of extruded loop positioning relative to Rec8 sites was calculated with *loopstats.py* in *looplib* (<u>https://github.com/golobor/looplib</u>), and arc diagrams (Extended Data Fig. 6c) with *loopviz.py*. Conformations showing chromatids or positions of extruded loop bases were rendered in PyMOL (https://pymol.org/sites/default/files/pymol.bib).

597

598 References

- Zickler, D. & Kleckner, N. Meiotic chromosomes: integrating structure and function. *Annu Rev Genet* 33, 603-754 (1999).
- Gray, S. & Cohen, P. E. Control of Meiotic Crossovers: From Double-Strand Break Formation to
 Designation. *Annu Rev Genet* (2016).
- Klein, F. et al. A central role for cohesins in sister chromatid cohesion, formation of axial elements, and
 recombination during yeast meiosis. *Cell* **98**, 91-103 (1999).
- Sym, M., Engebrecht, J. A. & Roeder, G. S. ZIP1 is a synaptonemal complex protein required for meiotic
 chromosome synapsis. *Cell* 72, 365-378 (1993).
- 5. Smith, A. V. & Roeder, G. S. The yeast Red1 protein localizes to the cores of meiotic chromosomes. *J Cell Biol* **136**, 957-967 (1997).
- Hollingsworth, N. M., Goetsch, L. & Byers, B. The HOP1 gene encodes a meiosis-specific component of
 yeast chromosomes. *Cell* 61, 73-84 (1990).
- Maleki, S., Neale, M. J., Arora, C., Henderson, K. A. & Keeney, S. Interactions between Mei4, Rec114,
 and other proteins required for meiotic DNA double-strand break formation in Saccharomyces cerevisiae.
 Chromosoma **116**, 471-486 (2007).
- 8. Li, J., Hooker, G. W. & Roeder, G. S. Saccharomyces cerevisiae Mer2, Mei4 and Rec114 form a complex required for meiotic double-strand break formation. *Genetics* **173**, 1969-1981 (2006).
- 9. Dekker, J., Rippe, K., Dekker, M. & Kleckner, N. Capturing chromosome conformation. *Science* 295, 1306-1311 (2002).
- 10. Bonev, B. & Cavalli, G. Organization and function of the 3D genome. Nat Rev Genet 17, 661-678 (2016).
- Merkenschlager, M. & Nora, E. P. CTCF and Cohesin in Genome Folding and Transcriptional Gene
 Regulation. *Annu Rev Genomics Hum Genet* 17, 17-43 (2016).

Schalbetter, Fudenberg et al. 2018

621	12.	Dekker, J. & Mirny, L. The 3D Genome as Moderator of Chromosomal Communication. <i>Cell</i> 164 , 1110-
622	10	1121 (2016).
623	13.	Fudenberg, G., Abdennur, N., Imakaev, M., Goloborodko, A. & Mirny, L. A. Emerging Evidence of
624		Chromosome Folding by Loop Extrusion. <i>Cold Spring Harb Symp Quant Biol</i> 82 , 45-55 (2017).
625	14.	Jin, Q., Trelles-Sticken, E., Scherthan, H. & Loidl, J. Yeast nuclei display prominent centromere
626		clustering that is reduced in nondividing cells and in meiotic prophase. <i>J Cell Biol</i> 141 , 21-29 (1998).
627	15.	Hayashi, A., Ogawa, H., Kohno, K., Gasser, S. M. & Hiraoka, Y. Meiotic behaviours of chromosomes and
628		microtubules in budding yeast: relocalization of centromeres and telomeres during meiotic prophase.
629		Genes Cells 3 , 587-601 (1998).
630	16.	Goloborodko, A., Marko, J. F. & Mirny, L. A. Chromosome Compaction by Active Loop Extrusion.
631		<i>Biophys J</i> 110 , 2162-2168 (2016).
632	17.	Gibcus, J. H. et al. A pathway for mitotic chromosome formation. Science 359, (2018).
633	18.	Kakui, Y., Rabinowitz, A., Barry, D. J. & Uhlmann, F. Condensin-mediated remodeling of the mitotic
634		chromatin landscape in fission yeast. Nat Genet 49, 1553-1557 (2017).
635	19.	Lazar-Stefanita, L. et al. Cohesins and condensins orchestrate the 4D dynamics of yeast chromosomes
636		during the cell cycle. <i>EMBO J</i> 36 , 2684-2697 (2017).
637	20.	Naumova, N. et al. Organization of the mitotic chromosome. Science 342, 948-953 (2013).
638	21.	Schalbetter, S. A. et al. SMC complexes differentially compact mitotic chromosomes according to
639		genomic context. Nat Cell Biol 19, 1071-1080 (2017).
640	22.	Xu, L., Ajimura, M., Padmore, R., Klein, C. & Kleckner, N. NDT80, a meiosis-specific gene required for
641		exit from pachytene in Saccharomyces cerevisiae. Mol Cell Biol 15, 6572-6581 (1995).
642	23.	Muller, H. et al. Characterizing meiotic chromosomes' structure and pairing using a designer sequence
643		optimized for Hi-C. Mol Syst Biol 14, e8293 (2018).
644	24.	Rao, S. S. P. et al. A 3D map of the human genome at kilobase resolution reveals principles of chromatin
645		looping. Cell 159, 1665-1680 (2014).
646	25.	Nora, E. P. et al. Spatial partitioning of the regulatory landscape of the X-inactivation centre. Nature 485,
647		381-385 (2012).
648	26.	Dixon, J. R. et al. Topological domains in mammalian genomes identified by analysis of chromatin
649		interactions. Nature 485, 376-380 (2012).
650	27.	Ito, M. et al. Meiotic recombination cold spots in chromosomal cohesion sites. Genes Cells 19, 359-373
651		(2014).
652	28.	Fudenberg, G. et al. Formation of Chromosomal Domains by Loop Extrusion. Cell Rep 15, 2038-2049
653		(2016).
654	29.	Schwarzer, W. et al. Two independent modes of chromatin organization revealed by cohesin removal.
655		Nature 551 , 51-56 (2017).
656	30.	Rao, S. S. P. et al. Cohesin Loss Eliminates All Loop Domains. Cell 171, 305-320.e24 (2017).
657	31.	Wutz, G. et al. Topologically associating domains and chromatin loops depend on cohesin and are
658		regulated by CTCF, WAPL, and PDS5 proteins. EMBO J 36, 3573-3599 (2017).

32. Gassler, J. et al. A mechanism of cohesin-dependent loop extrusion organizes zygotic genome

659

Schalbetter, Fudenberg et al. 2018

660		architecture. <i>EMBO J</i> 36 , 3600-3618 (2017).
661	33.	Conrad, M. N., Lee, C. Y., Wilkerson, J. L. & Dresser, M. E. MPS3 mediates meiotic bouquet formation in
662		Saccharomyces cerevisiae. Proc Natl Acad Sci U S A 104 , 8863-8868 (2007).
663	34.	Trelles-Sticken, E., Adelfalk, C., Loidl, J. & Scherthan, H. Meiotic telomere clustering requires actin for its
664		formation and cohesin for its resolution. J Cell Biol 170, 213-223 (2005).
665	35.	Sun, X. et al. Transcription dynamically patterns the meiotic chromosome-axis interface. <i>Elife</i> 4, (2015).
666	36.	Møens, P. B. & Pearlman, R. E. Chromatin organization at meiosis. <i>Bioessays</i> 9, 151-153 (1988).
667	37.	Marko, J. F. & Siggia, E. D. Polymer models of meiotic and mitotic chromosomes. Mol Biol Cell 8, 2217-
668		2231 (1997).
669	38.	Blat, Y. & Kleckner, N. Cohesins bind to preferential sites along yeast chromosome III, with differential
670		regulation along arms versus the centric region. Cell 98, 249-259 (1999).
671	39.	Crawley, O. et al. Cohesin-interacting protein WAPL-1 regulates meiotic chromosome structure and
672		cohesion by antagonizing specific cohesin complexes. <i>Elife</i> 5 , e10851 (2016).
673	40.	Challa, K., Lee, MS., Shinohara, M., Kim, K. P. & Shinohara, A. Rad61/Wpl1 (Wapl), a cohesin
674		regulator, controls chromosome compaction during meiosis. Nucleic Acids Res 44, 3190-3203 (2016).
675	41.	Jin, H., Guacci, V. & Yu, HG. Pds5 is required for homologue pairing and inhibits synapsis of sister
676		chromatids during yeast meiosis. J Cell Biol 186, 713-725 (2009).
677	42.	Petela, N. J. et al. Scc2 Is a Potent Activator of Cohesin's ATPase that Promotes Loading by Binding
678		Scc1 without Pds5. <i>Mol Cell</i> 70 , 1134-1148.e7 (2018).
679	43.	Nora, E. P. et al. Targeted Degradation of CTCF Decouples Local Insulation of Chromosome Domains
680		from Genomic Compartmentalization. Cell 169, 930-944.e22 (2017).
681	44.	Glynn, E. F. et al. Genome-wide mapping of the cohesin complex in the yeast Saccharomyces
682		cerevisiae. <i>PLoS Biol</i> 2 , E259 (2004).
683	45.	Lengronne, A. et al. Cohesin relocation from sites of chromosomal loading to places of convergent
684		transcription. <i>Nature</i> 430 , 573-578 (2004).
685	46.	Heinz, S. et al. Transcription Elongation Can Affect Genome 3D Structure. Cell 174, 1522-1536.e22
686		(2018).
687	47.	Haarhuis, J. H. I. et al. The Cohesin Release Factor WAPL Restricts Chromatin Loop Extension. Cell
688		169 , 693-707.e14 (2017).
689	48.	Tedeschi, A. et al. Wapl is an essential regulator of chromatin structure and chromosome segregation.
690		<i>Nature</i> 501 , 564-568 (2013).
691	49.	Busslinger, G. A. et al. Cohesin is positioned in mammalian genomes by transcription, CTCF and Wapl.
692		Nature 544 , 503-507 (2017).
693	50.	Ocampo-Hafalla, M., Muñoz, S., Samora, C. P. & Uhlmann, F. Evidence for cohesin sliding along
694		budding yeast chromosomes. <i>Open Biol</i> 6 , (2016).
695	51.	Ganji, M. et al. Real-time imaging of DNA loop extrusion by condensin. Science (2018).
696	52.	Sanborn, A. L. et al. Chromatin extrusion explains key features of loop and domain formation in wild-type
697		and engineered genomes. Proc Natl Acad Sci U S A 112, E6456-65 (2015).

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698	53.	Nasmyth, K. Disseminating the genome: joining, resolving, and separating sister chromatids during
699		mitosis and meiosis. Annu Rev Genet 35 , 673-745 (2001).
700	54.	Goloborodko, A., Imakaev, M. V., Marko, J. F. & Mirny, L. Compaction and segregation of sister
701		chromatids via active loop extrusion. <i>Elife</i> 5 , (2016).
702	55.	Alipour, E. & Marko, J. F. Self-organization of domain structures by DNA-loop-extruding enzymes.
703		Nucleic Acids Res 40 , 11202-11212 (2012).
704	56.	Subramanian, V. V. et al. Persistent DNA-break potential near telomeres increases initiation of meiotic
705		recombination on short chromosomes. Biorxiv 201889 (2017).
706	57.	Thacker, D., Mohibullah, N., Zhu, X. & Keeney, S. Homologue engagement controls meiotic DNA break
707		number and distribution. Nature 510, 241-246 (2014).
708	58.	Kerpedjiev, P. et al. HiGlass: web-based visual exploration and analysis of genome interaction maps.
709		Genome Biol 19 , 125 (2018).
710	59.	Belton, J. M. & Dekker, J. Hi-C in Budding Yeast. Cold Spring Harb Protoc 2015, 649-661 (2015).
711	60.	Servant, N. et al. HiC-Pro: an optimized and flexible pipeline for Hi-C data processing. Genome Biol 16,
712		259 (2015).
713	61.	Hsieh, T. S., Fudenberg, G., Goloborodko, A. & Rando, O. J. Micro-C XL: assaying chromosome
714		conformation from the nucleosome to the entire genome. Nat Methods 13, 1009-1011 (2016).
715	62.	Mizuguchi, T. et al. Cohesin-dependent globules and heterochromatin shape 3D genome architecture in
716		S. pombe. Nature 516, 432-435 (2014).
717	63.	Crane, E. et al. Condensin-driven remodelling of X chromosome topology during dosage compensation.
718		Nature 523, 240-244 (2015).
719	64.	Eastman, P. et al. OpenMM 7: Rapid development of high performance algorithms for molecular
720		dynamics. PLoS Comput Biol 13, e1005659 (2017).
721	65.	Eastman, P. & Pande, V. S. Efficient nonbonded interactions for molecular dynamics on a graphics
722		processing unit. J Comput Chem 31 , 1268-1272 (2010).
723	66.	Wong, H. et al. A predictive computational model of the dynamic 3D interphase yeast nucleus. Curr Biol
724		22 , 1881-1890 (2012).
725	67.	Tjong, H., Gong, K., Chen, L. & Alber, F. Physical tethering and volume exclusion determine higher-order
726		genome organization in budding yeast. Genome Res 22, 1295-1305 (2012).
727	68.	Fudenberg, G. & Imakaev, M. FISH-ing for captured contacts: towards reconciling FISH and 3C. Nat
728		Methods 14, 673-678 (2017).
729		

Figure 1

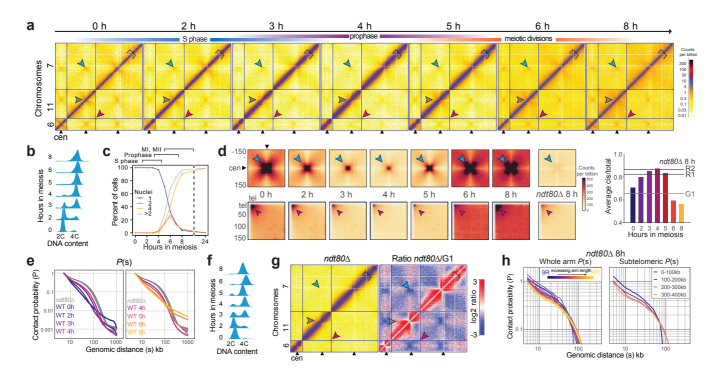


Figure 1. Chromosome conformation during yeast meiosis.

a. Cells were collected during meiosis at indicated timepoints and analysed by Hi-C. At 0h the cells are in G1. Representative Hi-C contact maps of chromosomes 6, 11, and 7 plotted at 5 kb resolution. Centromeres, telomeres and arm fold-back at the centromere are indicated by blue, red and grey arrows, respectively, and axial compaction by the width of the main diagonal relative to the fixed-width black clamp. For interactive HiGlass⁵² views see: <u>http://higlass.pollard.gladstone.org/app/?</u> config=Z5iwKpjzQpePCXXyvuYGeQ

b. Meiotic entry assessed by FACS; at 4 h, the majority of cells show a 4C peak indicating completion of DNA replication.
 c. Meiotic progression was monitored by quantification of nuclear divisions determined by DAPI staining. Around 4 h, cells start to undergo meiotic divisions I and II. The majority of cells undergo meiotic divisions between 4 and 8 h, indicating the degree of heterogeneity within the cell population.

d. Upper panels: Average trans centromere-centromere contact maps. Lower panels: trans telomere-telomere contact maps. Right: ratio of cis to total contact frequency.

e. Intra-arm contact probability versus genomic distance, P(s), indicating the emergence (*left*) and disappearance (*right*) of chromosome arm compaction during meiosis. Shaded area bounded above and below by the two *ndt80* Δ 8h replicates.

f. Meiosis was induced in *ndt80*Δ cells for 8h and meiotic entry was checked by monitoring DNA replication by FACS.
 g. *ndt80*Δ cells were grown for 8h in sporulation media and analysed by Hi-C (*left*). Log2 ratio of *ndt80*Δ cells 8h over G1 (*right*). Centromeres and telomeres are indicated by blue and red arrows, respectively, and axial compaction by a black clamp.
 h. *Left*: Contact probability of individual chromosome arms stratified by length. *Right*: Contact probability stratified by the distance from the telomere.

Figure 2

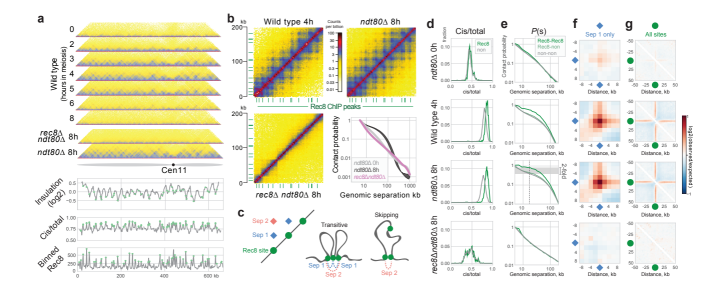


Figure 2. Emergence of a Rec8-dependent grid of punctate interactions in meiosis

a. Hi-C contact maps of chromosome 11 for the indicated genotypes plotted at 2 kb bin resolution, showing near-diagonal interactions. Wild type timepoints as in **Fig. 1a**. **Lower panels**: log2(insulation); cis/total ratio, Rec8 ChIP-seq²⁸, all binned at 2kb. Insulation and cis/total calculated from *ndt80*Δ maps. Positions of Rec8 sites indicated as green circles. Genome-wide cis/total (Spearman's R=0.62, P<1e-10) and insulation (R= -0.23, P<1e-10, insulation window = 20 kb) profiles are correlated with Rec8 occupancy.

b. Zoom-in into contact maps on chromosome 11 (0-200kb) of wt-4h and $ndt80\Delta$ (top) and $rec8\Delta$ (bottom left). Contact probability versus genomic distance, P(s), for G1($ndt80\Delta$ -0h) and $ndt80\Delta$ and $rec8\Delta$ (bottom right). Data shown is the average (n=2) except for wt-4h. Rec8 peak sites called from ChIP-seq data²⁸ are indicated in green. For an interactiveview see: <u>http://</u>higlass.pollard.gladstone.org/app/?config=Twrh61jGT4SIxotaguTIJg

c. Simplified illustration of how a grid of peaks on a Hi-C map can emerge between Rec8 sites either by transitive contacts between adjacent loops, or by loops that skip over adjacent sites. Experimentally observed grids extend much further than separation=2 (Extended Data Fig. 4c)

d. Cis/total ratios for Rec8 (green) and nonRec8 (grey) sites for indicated datasets.

e. Contact frequency versus distance between Rec8-Rec8 sites (green), Rec8-nonRec8 sites (light green) and nonRec8-nonRec8 sites (green).

f. Log2 ratio of contact frequency between adjacent Rec8-sites (separation=1) compared to average cis interactions.

g. Log2 ratio of contact frequency centered at Rec8 sites compared to average cis interactions.

In $ndt80\Delta$, Rec8 sites show: elevated cis/total frequency (0.85 versus 0.77), elevated pairwise contact frequency (~2-fold at 20 kb), and mild insulation. These distinctions are similar in wild type pachytene (4h) yet absent in G1 ($ndt80\Delta$ -0h) or in $rec8\Delta$.

Figure 3

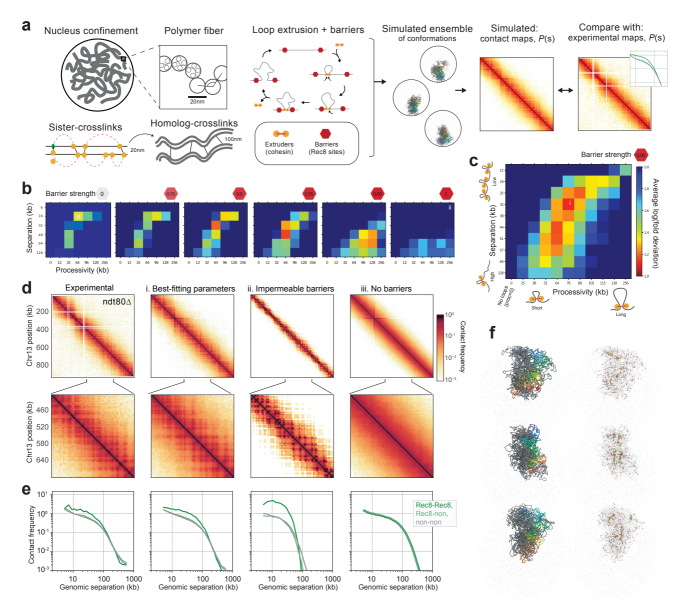


Figure 3. Model for meiotic chromosome compaction in prophase.

a. In simulations, yeast chr13 was represented as a polymer fiber confined to the nucleus subject to additional meiosisspecific constraints. These include: extruded loops, sister crosslinks, and homolog crosslinks (**Methods**). Barriers to extruded loops were placed at Rec8 sites²⁸. We imposed inter-sister and inter-homologue crosslinks at sites of extruded loop bases in order to approximate the paired arrangement of homologues at pachytene (**Extended Data Fig. 6**). For each set of extruded loop parameters (*processivity, separation,* and *barrier strength*), conformations were collected and used to generate simulated contact maps. These were then compared with experimental contact maps via the combined average fold discrepancy with P(s) curves for Rec8-Rec8, Rec8-non, and non-non bin pairs at 2 kb resolution.

b. Goodness-of-fit for indicated barrier strengths over coarse grids of processivity and separation demonstrate that intermediate barrier strengths are required to agree with experimental *ndt80* Hi-C maps.

c. Goodness-of-fit for a fine grid of processivity versus separation at barrier strength 0.95. Best-fitting models had separation ~32kb and processivity ~76kb, corresponding to ~60% coverage of the genome by extruded loops of average length 26kb. **d.** From *left* to *right*: contact maps for chr13 for *ndt80*Δ, and simulations with (i) best-fitting parameters, (ii) relatively stable loops between neighboring Rec8 sites, and (iii) no barriers.

e. P(s) split by Rec8-Rec8, Rec8-non, and non-non, as in Fig. 2d.

f. Conformations for best-fitting simulations, which highlight: *(left)* one chromatid colored from start (red) to end (blue); *(right)* extruders (yellow), extrusion barriers (red), and extruders paused at barriers (orange).

Figure 4

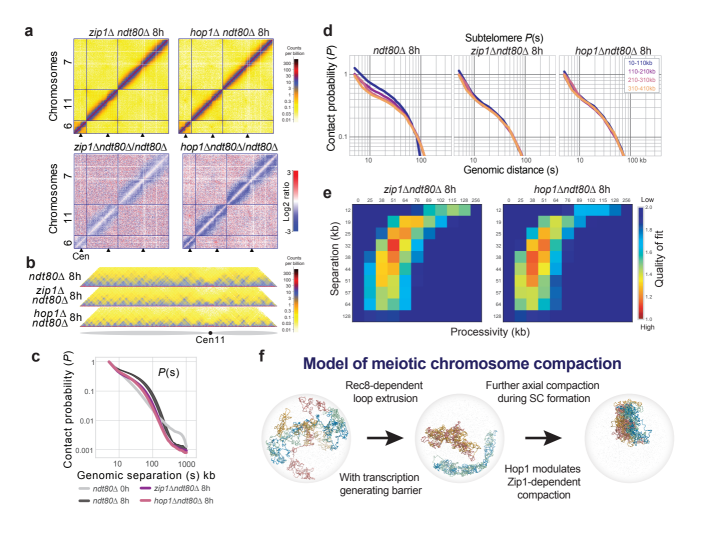


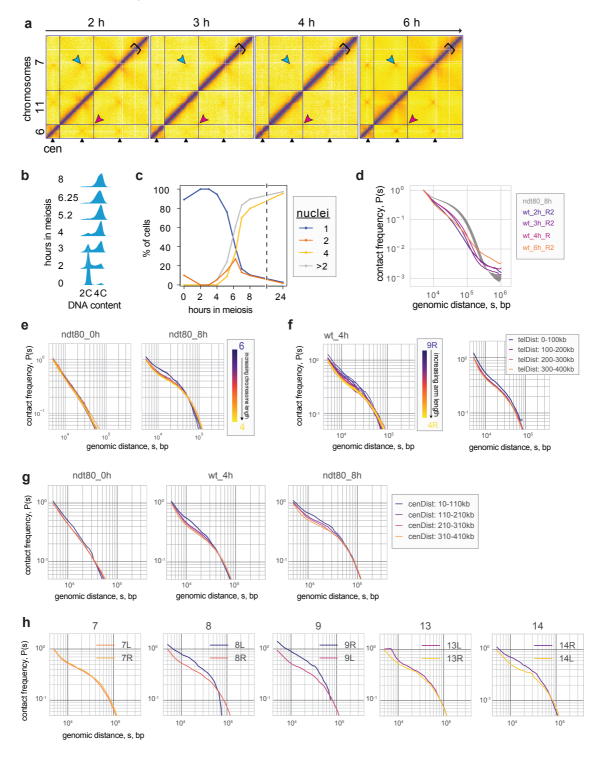
Figure 4. Hop1 and Zip1-dependent compaction of Rec8-dependent loops.

a. Top: Hi-C maps for $hop1\Delta$ and $zip1\Delta$ (plotted as in **Fig. 1a**). Bottom: Log2 ratio of hop1 Δ over *ndt80* Δ (as in **Fig. 1g**). For interactive views of the full genome, see <u>http://higlass.pollard.gladstone.org/app/?config=TTBGu5DDR0SHAa09zrjTXA</u> **b.** Hi-C contact maps of chromosome 11 for $hop1\Delta$ and $zip1\Delta$ plotted at 2kb bin resolution, showing near-diagonal interactions, as in **Fig. 2a**.

c. Contact probability versus genomic distance for G1, *ndt80* Δ , *hop1* Δ , *zip1* Δ . Shaded area bounded above and below by *ndt80* Δ replicas. Average between two replicas for *zip1* Δ and one sample for G1 and *hop1* Δ are shown. **d.** Contact probability over genomic distance averaged over all chromosome arms stratified by distance from the telomere.

e. Goodness-of-fit for simulations without homolog crosslinks with a fine grid of processivity versus separation at barrier strength 0.95 $zip1\Delta$ and $hop1\Delta$.

f. Model of meiotic chromosome compaction: Rec8-dependent loop formation leads to initial chromosome arm compaction and emergence of a grid-like pattern of Hi-C interactions that jointly agrees with a mechanism of loop extrusion including barrier elements. We suggest that transcription could impose such barriers. Hop1 and Zip1 are dispensable for this step, but are required for synapsis, where additional compaction occurs differentially along chromosome arms.



Extended Data Figure 1

Extended Data Fig. 1 Temporal and chromosome length-specific analysis of meiotic chromatin conformation a-d. Results from a replicate timecourse, collected and characterized independently of the timecourse in Fig. 1.

a. Hi-C maps, plotted as in Fig. 1a.

b. FACS as in Fig. 1b.

c. DAPI as in Fig. 1c.

d. P(s) as in Fig. 1e.

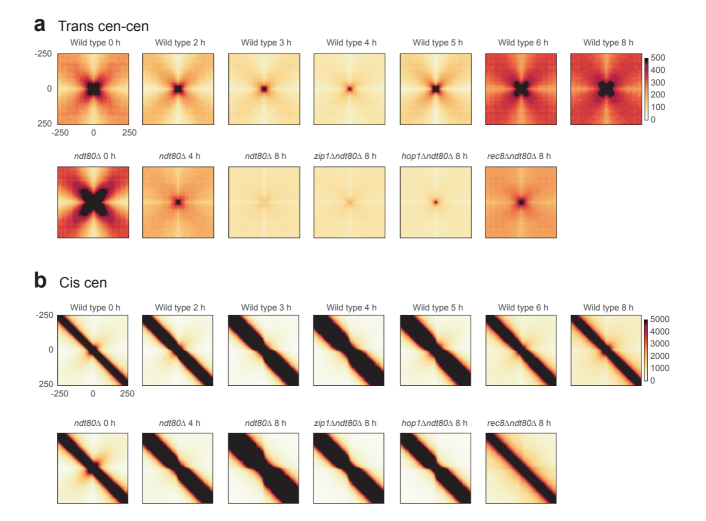
e. P(s) for chromosomes stratified by size for *ndt80* Δ -0h, *ndt80* Δ -8h. Short chromosomes display relatively elevated P(s) at short distances, and an earlier shoulder.

f. *Left*: P(s) for individual chromosome arms, stratified by size for wt-4h. Short arms display relatively elevated P(s) at short distances, and an earlier roll-over. *Right*: Intra-arm P(s) stratified by the distance from the telomere for wt-4h, averaged across all chromosomes. Telomere-proximal regions display elevated P(s) at short distances.

g. Intra-arm P(s) stratified by the distance from the centromere for G1 (*ndt80* Δ -0h), *wt*-4h, *ndt80* Δ -8h, averaged across all chromosomes.

h. Contact probability of single chromosome arms for *ndt80*Δ-8h.

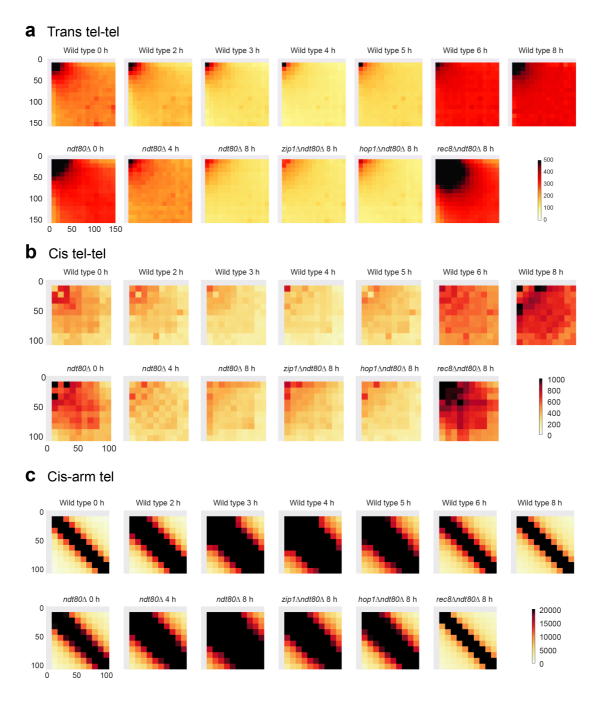
Extended Data Figure 2



Extended Data Fig. 2 Aggregate analysis of centromeric interactions in meiosis

a. Average trans centromere-centromere contact maps for indicated data sets.

b. Average *cis* centromere-centromere contact maps for indicated data sets. Note the loss of the folding back in meiosis, and how the intra-arm enrichment is insulated at centromeres in meiosis.



Extended Data Figure 3

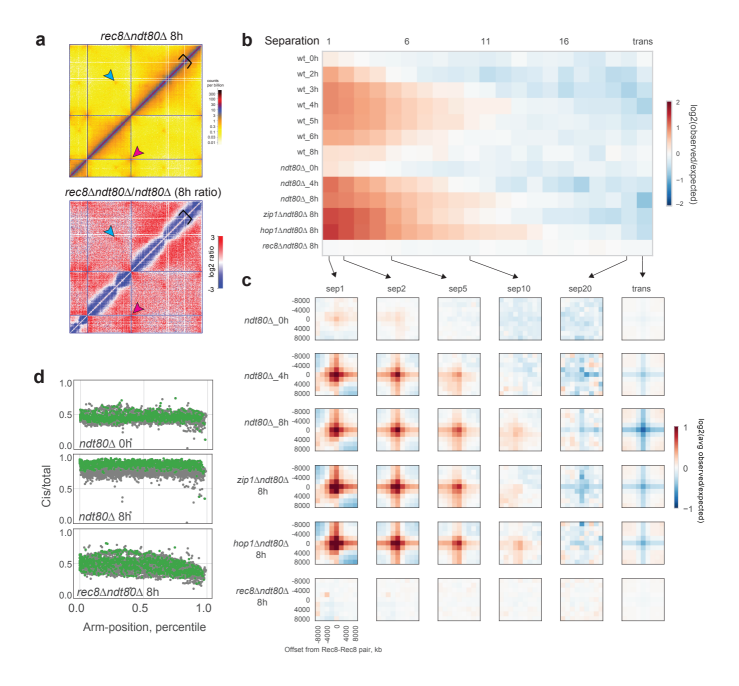
Extended Data Fig. 3 Aggregate analysis of telomeric interactions in meiosis

a. Average trans telomere-telomere contact maps for indicated datasets.

b. Average telomere-telomere contact maps between the two telomeres of the same chromosome.

c. Average contact map around each telomere in cis.

Extended Data Figure 4



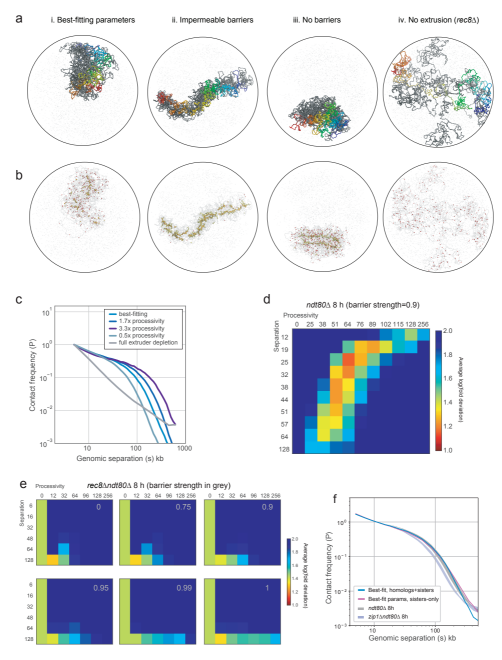
Extended Data Fig. 4 Preferred sites of Rec8 occupancy define sites of locus-specific interaction

a. *Left*: Hi-C contact maps of *rec8 ndt80 A*. Chromosomes 6, 11 and 7 are shown as representatives for the whole genome. *Right*: Log2 Hi-C ratio maps of *rec8 ndt80 A ndt80 A*. Plotted as in Fig. 1g.

 b. Log2 observed over expected contact frequency at Rec8-Rec8 peak pairs as a function of separation across datasets.
 c. Log2 observed over expected contact frequency +/-8 kb around Rec8-Rec8 peak pairs at the indicated separations. Together, b-c demonstrate that Rec8-Rec8 enrichments are strongest between adjacent sites, decrease between non-adjacent sites with increasing genomic separation, and are absent in *trans*. Equally important, these meiotic features are lost in *rec8*Δ. As for mammalian interphase, this observation in meiosis argues for a *cis*-acting process underlying the formation of focal interactions between Rec8 sites.

d. cis/total as a function of distance along the chromosomal arm, Rec8 sites marked in green.

Extended Data Figure 5



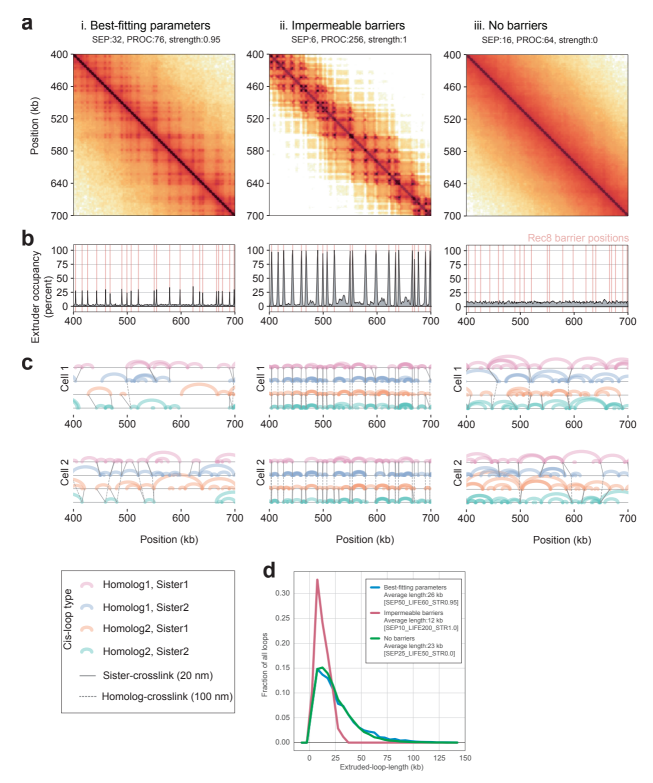
Extended Data Fig. 5 Polymer simulations of loop extrusion reveal best fitting parameters and conformations a. Representative conformation for the indicated parameter sets. As in **Fig. 3F**, one chromatid from a homologous quartet of chromatids coloured from start to end according to the spectrum; other three coloured in grey.

b. For the same three conformations, positions of Rec8 sites indicated with red spheres, positions of extruded loop bases in yellow, and extruders overlapping a Rec8 site in orange. Note the stable loops between neighbouring Rec8 sites creates a very elongated chromatid (ii). Also note the majority of Rec8 sites are unoccupied in (iii), despite the self-assembly of two axial cores and a strong brush. Finally, note very dispersed chromosomes in (iv), consistent with EM³ for *rec8*\Delta. **c.** Contact frequency versus distance, P(s), for indicated simulations. Note that the loss of the shoulder in P(s) in the case of full extruder depletion mirrors the difference between experimental $ndt80\Delta$ and $rec8\Delta$ Hi-C maps. Simulations with increased processivity predict that P(s) would shift rightward if unloading was impaired, as could happen in *wapl*\Delta. Conversely, if unloading was enhanced, simulations with decreased processivity indicate a leftward shift in P(s), until the absence of extruders.

d. Goodness-of-fit for a fine grid of processivity versus separation at barrier strength 0.90. The best-fit occurs at similar processivity and separation as for barrier strength 0.95 shown in Fig. 3c, but with slightly lower goodness-of-fit.
e. Goodness-of-fit to *rec8*∆ data for simulations with the indicated barrier strengths (in grey: 0.00, 0.75, 0.90, 0.95, 0.99, 1.00) over coarse grids of processivity and separation demonstrates that the best fits have few if any extruded loops, regardless of barrier strength.

f. P(s) curves for simulations with sisters and homologs with the best-fitting parameters for *ndt80* Δ -8*h* maps compared to P(s) for simulations with sisters only show that simply removing homolog tethering does not recapitulate the sort of shifted P(s) seen experimentally in *zip1* Δ Hi-C.

Extended Data Figure 6



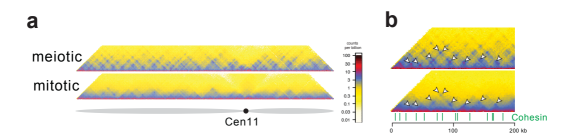
Extended Data Fig. 6 Polymer simulations of loop extrusion elucidate meiotic barrier strength

a. Simulated contact maps for the indicated region of chr13 for: (i) best-fitting simulations, (ii) simulations with relatively stable loops between neighboring Rec8 sites (barrier strength=1 and high processivity), and (iii) no barriers, as in **Fig. 2d.** b. Simulated ChIP-seq profiles for the indicated region of chr13. Best-fitting simulations (i) display occupancy well below 100% at Rec8 sites. Simulations with stable loops (ii) display highly occupied Rec8 sites. Simulations without barriers (iii) have homogenous Rec8 occupancy across the genome.

c. Positions of extruded loops (arcs) sister crosslinks (solid black lines) and homolog crosslinks (dashed lines) for four chromatids in two separate cells, showing how the simulated Hi-C maps and ChIP-seq profiles emerge from the stochastic positioning of extruded loops from cell-to-cell. For statistics, see Supplementary Table S1.

d. Histogram of extruded loop lengths for indicated parameters (i, ii, iii).

Extended Figure 7



Extended Data Figure 7. Cohesin and transcription patterns loops in meiosis and mitosis a. Hi-C contact maps of chromosome 11 for meiotic (*ndt80*Δ, pachytene - top) and mitotic (wild

a. HI-C contact maps of chromosome 11 for melotic ($nat80\Delta$, pachytene - top) and mitotic (wild type, nocodazole arrest - bottom) plotted at 2 kb bin resolution, showing near-diagonal interactions. Data shown is the average (n=2).

b. Zoom-in into contact maps on chromosome 11 (0-200kb) of *ndt80* Δ (top) and mitotic (*bottom*). Data shown is the average (n=2). Rec8 peak sites called from ChIP-seq data²⁷ are indicated in green. Arrowheads indicate sites of prominent focal interaction.