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| 3 | Structural basis of meiotic chromosome synapsis through SYCP1 self-assembly | | | | |
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22 Abstract

23 Meiotic chromosomes adopt unique structures in which linear arrays of chromatin loops are bound 24 together in homologous chromosome pairs by a supramolecular protein assembly, the 25 synaptonemal complex. This three-dimensional scaffold provides the essential structural framework 26 for genetic exchange by crossing over and subsequent homologue segregation. The core 27 architecture of the synaptonemal complex is provided by SYCP1. Here, we report the structure and 28 self-assembly mechanism of human SYCP1 through X-ray crystallographic and biophysical studies. 29 SYCP1 has an obligate tetrameric structure in which an N-terminal four-helical bundle bifurcates into 30 two elongated C-terminal dimeric coiled-coils. This building-block assembles into a zipper-like lattice 31 through two self-assembly sites. N-terminal sites undergo cooperative head-to-head assembly in the 32 midline, whilst C-terminal sites interact back-to-back on the chromosome axis. Our work reveals the 33 underlying molecular structure of the synaptonemal complex in which SYCP1 self-assembly 34 generates a supramolecular lattice that mediates meiotic chromosome synapsis.

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44 Keywords

45 Meiosis, chromosome structure, double-strand break, chiasmata, synaptonemal complex, SYCP1,

46 self-assembly

48 Introduction

49 The reduction in chromosome number during meiosis requires a unique programme of intricate 50 molecular processes including the synapsis of homologous chromosome pairs, their exchange of 51 genetic material by crossing over, and ultimately their segregation into haploid cells. At the centre of 52 these processes is a supramolecular protein assembly, the synaptonemal complex (SC). The SC binds 53 together homologous chromosome pairs, structured as linear arrays of chromatin loops, in a single 54 continuous synapsis along their entire length^{1,2}. SC assembly occurs in a spatiotemporal manner, 55 dependent on the prior establishment of inter-homologue recombination intermediates through 56 double-strand break induction, which act as guides to ensure the synapsis of perfectly aligned 57 homologues^{3,4}. The three-dimensional structure of the SC provides the essential architectural 58 framework for the resolution of recombination intermediates, which includes the generation of one genetic crossover per chromosome arm^{5,6}. Crossovers are essential for correct segregation of 59 homologues at anaphase I, and additionally contribute to genetic diversity. The defective assembly 60 of the SC is associated with human infertility, miscarriage and aneuploidy^{7,8}. However, despite its 61 62 discovery more than half a century ago, the molecular structure and function of the SC have 63 remained unknown.

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Electron micrographs of the SC reveal a characteristic tripartite structure that is conserved across eukaryotes⁹. This consists of two lateral elements, each coating a chromosome axis, separated by a 100 nm central region that contains a midline 20-40 nm wide central element (Fig. 1a). The central and lateral elements are connected together by a network of angled transverse filaments, which in hamster have a diameter of approximately 16 Å and are spaced at a density of 50-80 per 1 μm of chromosome axis¹⁰. In addition to its 100 nm width, the SC central region has a depth of up to 100 nm, so is a truly three-dimensional protein assembly^{11,12}.

In mammals, SC transverse filaments are formed by SYCP1¹³. This 976 amino acid protein contains a 73 74 central α -helical core flanked by unstructured N- and C-terminal tails (Fig. 1b). SYCP1 N- and Ctermini are localised within SC central and lateral elements respectively, and so is bioriented with 75 76 juxtaposed SYCP1 molecules providing a 150 nm separation between opposing C-termini in mice^{12,14,15} (Fig. 1a). The SC contains at least two layers of SYCP1 molecules; N-termini are detected 77 78 in two vertically separated chains within the central element, whereas C-termini are present in a 79 single chain within the lateral element^{12,16}. SC lateral elements also contain SYCP2 and SYCP3^{17,18}, the 80 latter contributing to chromosome compaction through stabilisation of chromatin loop structures¹⁹⁻ ²¹. The SC central element contains initiation factors SYCE3, SYCE1 and SIX6OS1 that stabilise initial 81 tripartite structures²²⁻²⁵, and elongation complex SYCE2-TEX12 that stabilises the long-range 82 83 extension of the tripartite structure²⁶⁻²⁹.

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85 SYCP1 disruption leads to a complete failure of synapsis; recombination intermediates are formed 86 but fail to resolve, crossovers fail to form, cells undergo meiotic arrest and there is a resultant 87 complete infertility⁵. Whilst SC central and lateral element components are essential for the 88 structure and function of the mature SC, SYCP1 is recruited to meiotic chromosomes in the absence 89 of other SC central and lateral element components, albeit at reduced levels, and is essential for the recruitment of all SC central element proteins^{5,17,18,22,23,25-28}. Furthermore, SYCP1 in isolation has an 90 intrinsic capacity for self-assembly into rudimentary SC-like structures³⁰. Thus, SYCP1 self-assembly 91 92 seemingly provides the underlying architectural framework of the SC.

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Here, we report the structure and self-assembly mechanism of SYCP1. The obligate unassembled structure of SYCP1 is an N-terminal tetramer that bifurcates into two elongated C-terminal dimeric coiled-coils. This building-block self-assembles into a supramolecular lattice that defines the SC structure through sites within its N- and C-termini. Whilst N-terminal sites undergo cooperative head-to-head assembly, C-terminal sites interact back-to-back in a protonation-dependent manner

- that relies upon chromosomal recruitment by unstructured C-terminal tails. Together, our data lead
 to a complete molecular model for the structure of SYCP1 in which recursive self-assembly at N- and
 C-terminal sites leads to the formation of a continuous and cooperative supramolecular lattice.
 Through this, we reveal the underlying structure of the synaptonemal complex and the molecular
 basis of meiotic chromosome synapsis by SYCP1.
- 105

106 Results

107 The obligate structure of SYCP1

108 Human SYCP1 contains a large α -helical core (α Core) of amino acids 101-783, flanked by 109 unstructured N- and C-terminal tails (Fig. 1b and Supplementary Fig. 1a). Size-exclusion 110 chromatography multi-angle light scattering (SEC-MALS) analysis of purified recombinant SYCP1 111 α Core revealed heterogeneous 1-12 MDa species (Fig. 1c and Supplementary Fig. 1b), indicating an 112 intrinsic capacity to self-assemble in vitro. Self-assembly of large molecular weight species is 113 completely abrogated by deletion of the first 11 amino acids at its N-terminal tip (α N-tip), with 114 α Core- Δ Ntip (residues 112-783) forming a stable tetramer (Fig. 1c). Circular dichroism (CD) 115 spectroscopy confirms that α Core- Δ Ntip is almost entirely α -helical (Supplementary Fig. 2a,b). Size 116 exclusion chromatography small-angle X-ray scattering (SEC-SAXS) analysis reveals scattering profiles 117 and real space pair-distance distribution functions (P(r) distributions) corresponding to an elongated 118 molecule of 900 Å length (Fig. 1d and Supplementary Fig. 2c). This matches its theoretical α -helical coiled-coil length and is sufficient to span just over half of the inter-chromosomal distance, in 119 120 keeping with SYCP1 biorientation within the SC. We conclude that α Core- Δ Ntip is an extended α -121 helical coiled-coil tetramer that represents the obligate structure of SYCP1, and self-assembly of this 122 minimum building-block into higher molecular weight species is dependent on the N-terminal tip of 123 SYCP1 αCore.

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The obligate α Core- Δ Ntip is composed of two distinct structural units, an N-terminal tetramer (residues 206-362) and C-terminal dimer (residues 358-783) (Fig. 1c). These boundaries were identified through exhaustive screening to define clearly demarcated structural regions of maximal stability; nevertheless, oligomer states and structures of these and other constructs described herein are robust across a range of sequence boundaries (Supplementary Fig. 1a and Supplementary Table 1). The α N-tetramer and α C-dimer are almost entirely α -helical (Supplementary Fig. 2a,b); SEC-SAXS analysis reveals elongated structures of respective lengths 260 Å and 645 Å (Fig. 1d and Supplementary Fig. 2c-e), matching their theoretical coiled-coil lengths. The cross-sectional radius of gyration (Rc) was determined as 10.3 Å and 8.9 Å for α N-tetramer and α C-dimer (Supplementary Fig. 2f), corresponding to the known dimensions of four-helical and dimeric coiled-coils respectively. The Rc of α C-dimer (8.9 Å) indicates a diameter of 17.8 Å that closely matches the measured 16 Å diameter of transverse filaments in the hamster SC¹⁰, suggesting that α C-dimers constitute the individual structures visualised spanning between SC central and lateral elements.

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139 We determined the orientation of helices within α Core- Δ Ntip, α N-tetramer and α C-dimer through 140 SEC-SAXS P(r) analysis of N-terminal MBP fusion proteins, exploiting the strong scattering of globular 141 proteins in comparison to coiled-coils to identify the relative positions of globular tags. In all cases, 142 P(r) distributions demonstrate strong inter-MBP peaks at short distances, compatible with their 143 parallel orientation, but lack inter-MBP peaks at long distances that would occur in anti-parallel 144 structures (Fig. 1e and Supplementary Fig. 2d-h). Similarly, an N-terminal GST fusion of α N-tetramer 145 shows only short distance inter-GST peaks (Supplementary Fig.2 j-n). Finally, the α N-tetramer and 146 α C-dimer structures are compatible with their N-terminal fusion to a constitutive tetramer (RecE) 147 and dimer (GST) respectively (Fig. 1f and Supplementary Fig. 2i-n), confirming their parallel 148 orientation. Thus, αCore-ΔNtip, αN-tetramer and αC-dimer are parallel coiled-coils, in keeping with 149 the biorientation of SYCP1 molecules within the SC. We conclude that the obligate structure of 150 SYCP1, which provides the minimal building block for self-assembly, is an N-terminal four-helical 151 bundle that bifurcates into C-terminal dimeric coiled-coils of sufficient length to span between SC 152 central and lateral elements (Fig. 1g).

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154 SYCP1 N-terminal self-assembly

155 The α N-tip (residues 101-111) is essential for self-assembly of α Core into large molecular weight 156 species *in vitro* and is part of a short α N-end region (residues 101-206), immediately preceding the 157 α N-tetramer, that is the most highly conserved portion of SYCP1 (Fig. 1b). The X-ray crystal structures of two α N-end constructs (residues 101-206 and 101-175) reveal tetrameric assemblies in which two parallel dimeric coiled-coils interact head-to-head (Fig. 2a,b, Supplementary Fig. 3 and Table 1). The head-to-head interface is mediated entirely by the α N-tip (Fig. 2a,b), suggesting that this 'dimer of dimers' structure may be responsible for SYCP1 N-terminal self-assembly into higher order structures.

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164 The two α N-end crystal structures demonstrate a common fold in which parallel dimeric coiled-coils 165 splay apart through a wedge formed of W119 and I116 to allow the α N-tips of opposing molecules to 166 interact head-to-head (Figs. 2a,b and 3a,d). The head-to-head interface shows distinct but highly 167 related conformations in the two structures, indicating conformational plasticity. The open 168 conformation of αN -end is asymmetrical and crescent-shaped, formed of midline and lateral anti-169 parallel coiled-coil interactions (Figs. 2a and 3a-c,g). The closed conformation of truncated α N-end is 170 a symmetrical four-helical bundle, consisting of a hydrophobic core and analogous midline and 171 lateral helical interfaces (Figs. 2b and 3d-f,h). The two conformations are formed of identical amino 172 acids undergoing largely similar coiled-coil and aromatic stacking interactions (Fig. 3b-c,e-h), and 173 likely exist in equilibrium, undergoing conformational change through a rotamer flip of central Y106 174 residues (Fig. 3g,h and Supplementary Fig. 3e). This structural plasticity may be important in 175 enforcing synapsis whilst accommodating large-scale twisting and bending of synapsed meiotic 176 chromosomes, with the open conformation permitting wider angulation between opposing SYCP1 177 molecules than the more rigid closed conformation.

178

SYCP1 αCore self-assembly is recapitulated by construct αN (residues 101-362) that includes both αN-end and αN-tetramer (Fig. 4a and Supplementary Fig. 4a,b). Its self-assembly into large molecular weight species is blocked by removal of either sequence, and is retained in the presence of the unstructured N-terminal tail (Fig. 4a,b and Supplementary Fig. 4c-e). Thus, the presence of αN-end and αN-tetramer is necessary and sufficient for SYCP1 N-terminal self-assembly *in vitro*. 184 Mutation of head-to-head interacting residues V105 and L109 to glutamate completely abrogates 185 αN self-assembly into large molecular weight species, leaving a stable obligate tetramer (Fig. 4a). 186 Thus, the α N-end head-to-head interaction is likely responsible for SYCP1 N-terminal self-assembly. 187 We propose that the α N-tetramer provides a structural scaffold from which two α N-end dimensions 188 splay apart, with their α N-tips interacting head-to-head with opposing SYCP1 molecules. A staggered 189 configuration provides a simple model for the cooperative assembly of a continuous lattice structure 190 of potentially limitless length, which we propose defines the structural basis of midline SYCP1 N-191 terminal self-assembly (Fig. 4c).

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193 Isolated α N-end is monomeric (Fig. 4b), indicating that individual head-to-head interactions are 194 weak and only form when the α N-tetramer mediates lattice formation. This requirement for 195 cooperativity favours the self-assembly of a single continuous lattice between appropriately aligned 196 meiotic chromosomes rather than forming heavily branched unproductive cellular assemblies (Fig. 197 4d).

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199 SYCP1 C-terminal self-assembly

A highly conserved sequence at the C-terminal end of SYCP α Core caps off the α C-dimer parallel coiled-coil (Fig. 1b). The X-ray crystal structure of α C-end (residues 676-770) reveals an anti-parallel tetramer in which two α C-end parallel dimers interact back-to-back in an intertwined α -helical assembly (Fig. 5, Supplementary Fig. 5 and Table 1). We suggest that this α C-end tetrameric assembly provides the structural basis for SYCP1 C-terminal self-assembly on the chromosome axis.

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In solution, αC-end is dimeric at pH 8.0 and tetrameric at pH 5.5 (Fig. 6a and Supplementary Fig.
6a,b). SEC-SAXS reveals that both species have similar length, but the cross-sectional radius
increases from 7.8 Å to 10.1 Å at pH 5.5, consistent with a transition from dimeric to four-helical
coiled-coil (Fig. 6b and Supplementary Fig. 6c-e). SAXS *ab initio* envelopes of the pH 8.0 and pH 5.5

210 species match the dimensions of a dimeric coiled-coil and the α C-end tetramer structure 211 respectively (Fig. 6c,d). SEC-SAXS P(r) distributions of MBP fusions of α C-end at pH 8.0 show inter-212 MBP peaks at short distances, compatible with their parallel orientation; peaks at long anti-parallel 213 distances were observed only upon MBP fusion at both termini, and for MBP- α C-end upon 214 tetrameric assembly at pH 5.5 (Fig. 6e and Supplementary Fig. 6f-j). Similarly, GST- α C-end forms a 215 stable dimer at pH 8.0 (Supplementary Fig. 6h-k). Finally, a tethered dimer of two consecutive α C-216 end sequences joined by a flexible linker is dimeric at pH 8.0, with length 241 Å and cross-sectional 217 radius 8.8 Å, consistent with it forming two consecutive dimeric coiled-coils (Figs. 6f,g and 218 Supplementary Fig. 6c-e). It remains dimeric at pH 5.5, but becomes a compact molecule of length 219 156 Å and cross-sectional radius 10.7 Å, indicating the folding back of α C-end sequences into an anti-220 parallel tetramer (Figs. 6f,g and Supplementary Fig. 6c-e). We conclude that α C-end is a parallel 221 dimeric coiled-coil that undergoes pH-induced back-to-back assembly into the anti-parallel tetramer 222 observed in the crystal structure.

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224 The α C-end crystal structure has a highly conserved central tetrameric interface in which H717 and 225 Y721 residues (invariant throughout vertebrates) form a hydrophobic core and engage in hydrogen 226 bonding networks with Q720 residues (Figs. 1b and 5a,b, Supplementary Figs. 1a and 5g). The 227 position of H717 residues suggested that their protonation may mediate pH-induced assembly. We 228 introduced mutation H717W Y721F, designed to stabilise the core whilst eliminating pH-sensitivity 229 (Supplementary Fig. 5h), into an extended α C-end construct (residues 676-783) that accentuates the 230 elution difference between dimer and tetramer (Supplementary Fig. 7). H717W Y721F is tetrameric 231 at pH 8.0 (Fig. 6h and Supplementary Fig. 7j), suggesting that pH-induced assembly in wild type 232 involves stabilisation of the core through H717 protonation. Accordingly, mutation H717E blocked 233 pH-induced tetrameric assembly (Supplementary Fig. 7k).

235 The central interface leads to pinch points, where N-terminal parallel coiled-coil dimers are flanked 236 by angulated C-terminal chains (Fig. 5a,c,d). The coiled-coil includes C703 heptad interactions that 237 are disulphide and non-disulphide at the respective smoothly and sharply angulated ends of the 238 molecule. An alternative α C-end crystal form contains symmetry-related pinch points with C703 239 partial disulphide bonds and smoothly angulated flanking chains (Supplementary Fig. 5). Whilst 240 disulphide bond formation may be a crystallisation artefact, it may also provide an intriguing means 241 for stabilising assembly in vivo; notably, the α N-end head-to-head assembly includes similar heptad 242 interactions between pairs of C183 and C190 residues.

243

244 The ends of the tetrameric structure are formed of four-helical bundles, consisting of a hydrophobic 245 core and anti-parallel coiled-coil interfaces (Fig. 5a,e-g). Hydrophobic core residues outline heptad 246 repeats within N- and C-terminal chains, with the latter constituting a three-heptad leucine zipper 247 (Fig. 5g). These residues likely also mediate parallel coiled-coil interactions in the dimeric 248 conformation. Amino acids L679 and I688 mediate anti-parallel interactions but lie outwith the 249 hydrophobic core heptads, so may be specific for the tetramer. The mutation L679A I688A 250 eliminated tetramer assembly but retained dimer formation (Fig. 6h and Supplementary Fig. 7l). We 251 conclude that heptad residues of the α C-end termini are bifunctional in mediating parallel dimeric 252 and anti-parallel tetrameric interactions, with the conformational change triggered by structural 253 alteration of the protonation-sensitive central interface.

254

In the cell, back-to-back assembly of α C-end may be triggered by its concentration on the chromosome axis, through local protonation induced by the high proton density in the close proximity of DNA³¹ or by specific interactions with chromosome axis proteins. Thus, protonationdependent conformational change of α C-end provides an elegant mechanism for triggering SYCP1 Cterminal self-assembly upon chromosomal recruitment.

261 DNA binding by SYCP1

The α C-end tetrameric structure contains a series of surface basic patches separated by ~30 Å (Fig. 7a), suggesting a direct interaction with the DNA backbone. Analysis by electrophoretic mobility shift assay (EMSA) revealed strong double-stranded DNA binding of tetrameric α C-end at pH 5.5, but not of the dimer at pH 8.0 (Fig. 7b). The presence of DNA-binding interfaces on both surfaces of the α Cend tetramer could mediate the formation of large protein-DNA assemblies, possibly accounting for the range of species observed. The α C-end tetrameric conformation is likely stabilised by interaction with DNA, and so SYCP1 C-terminal self-assembly and DNA-binding may be mutually reinforcing.

269

270 How is the SYCP1 C-terminus first recruited to chromosomes? The SYCP1 C-terminal tail contains 271 basic patches that could be obligate DNA-binding sites. A C-terminal construct including both α C-end 272 and Ctail (residues 640-976) interacts with DNA at neutral pH; DNA-binding is dependent on the Ctail 273 and is diminished upon deletion of α C-end (Fig. 7c and Supplementary Fig. 8a-d). We observe slightly 274 enhanced DNA-binding at neutral pH by SYCP1 constructs extended N-terminally to include the α C-275 dimer and α Core (residues 358-976 and 101-976) (Fig. 7c and Supplementary Fig. 8e-h). Electron 276 microscopy reveals the formation of ~10 nm wide protein-DNA complexes by α C-end~Ctail, which 277 develop a wider frayed appearance upon inclusion of the α C-dimer, consistent with transverse 278 filaments emanating from a core protein-DNA structure (Fig. 7d). Finally, we tested DNA-binding of 279 full length SYCP1 using refolded protein that demonstrates α -helical structure and higher order 280 assembly consistent with our findings for α Core (Supplementary Fig. 9). Full length SYCP1 interacts 281 with DNA at neutral pH, and the interaction is disrupted by deletion of the Ctail (Fig. 7e). Together, 282 these data demonstrate that SYCP1 binds DNA through its Ctail and the interaction is enhanced by 283 the α C-end and wider SYCP1 structure.

284

We propose that SYCP1 molecules are recruited to meiotic chromosomes through sites within Ctails,
leading to their concentration on chromatin. The close proximity of DNA and/or interactions with

axis proteins then triggers protonation-induced assembly of α C-end into tetramers that bind DNA and strengthen axis associations. The anti-parallel α C-end tetramers also mediate back-to-back interactions between SYCP1 molecules, which given their known orientation within the SC, likely result in looped U-shaped linkages between adjacent α C-dimer transverse filaments (Fig. 7f). Thus, SYCP1 C-terminal self-assembly integrates DNA-binding and interactions between adjacent transverse filaments to achieve SYCP1 coating of chromosome axes.

294 Discussion

295 We integrate our crystallographic and biophysical findings into a molecular model for meiotic 296 chromosome synapsis by SYCP1. The SYCP1 core consists of an α N-tetramer that bifurcates into two 297 α C-dimers (Fig. 8a). This tetrameric building-block self-assembles into a supramolecular lattice 298 through its N- and C-terminal ends. In the midline, α N-end dimers splay from α N-tetramer scaffolds 299 and interact head-to-head in a highly cooperative lattice. In the lateral element, α C-end dimers 300 assemble back-to-back as discrete intertwined tetramers that tether together adjacent α C-dimer 301 transverse filaments and reinforce chromosomal associations of C-terminal tails. Together, N- and C-302 terminal self-assembly collaborate to generate a cooperative zipper-like supramolecular lattice of 303 SYCP1 molecules capable of mediating continuous synapsis between homologous chromosomes (Fig. 304 8b). During SC assembly, midline lattice formation and chromosomal recruitment likely occur 305 concomitantly in one dynamic process of progressive chromosome synapsis. Whilst we cannot 306 exclude additional roles for N- and C-terminal tails in SYCP1 assembly in vivo, these regions are 307 largely unstructured and have no effect on oligomer states in vitro. Conserved amino acid sequences 308 within C-terminal tails may mediate currently unidentified interactions with chromosome axis 309 proteins, which act in concert with direct DNA-binding to achieve meiotic chromosome recruitment 310 of SYCP1.

311

312 Our model for SYCP1 self-assembly is consistent with the dimensions of the native SC. The SYCP1 313 tetrameric core has a length of 900 Å, sufficient to span just over half of the inter-chromosomal distance. The α C-dimer has an 8.9 Å cross-sectional radius and 645 Å length, matching the 314 315 dimensions of individual transverse filaments measured by electron microscopy in the hamster SC¹⁰. 316 We propose that α C-dimers constitute the transverse filaments visualised spanning between central 317 and lateral elements, with α N-tetramers buried within the central element. Importantly, anti-parallel 318 tetramer formation by α C-end explains how parallel SYCP1 molecules interact back-to-back to achieve the well-established biorientation of SYCP1 N- and C-termini within the SC^{12,14,15}. A recent 319

study reported that a region similar to α C-end is an anti-parallel dimer³², incompatible with established localisation patterns. Examination of their structural data (pdb 4YTO) reveals the presence of an anti-parallel tetramer within the crystal lattice, indicating that the anti-parallel dimer of the asymmetric unit was incorrectly attributed as the biological molecule (Supplementary Figure 5i).

325

The three-dimensional SC assembly contains at least two layers of transverse filament proteins^{11,12,16}, which is compatible with the SYCP1 supramolecular assembly that we describe. We propose that two parallel head-to-head SYCP1 lattices are connected by vertically (or obliquely) orientated backto-back assemblies within lateral elements (Supplementary Fig. 10a). This model is consistent with the observed vertical separation of SYCP1 N-termini by up to 100 nm, and the presence of single tracks of SYCP1 C-termini within lateral elements^{12,16}.

332

333 How is SYCP1 self-assembly directed to occur predominantly between aligned chromosomes? Whilst SYCP1 can form chromatin-free polycomplexes in meiotic tissue³³, assembly into SCs is heavily 334 335 favoured. Two distinct mechanisms cooperate to favour timely SYCP1 self-assembly between aligned 336 chromosomes. Firstly, αN -end head-to-head interactions are individually weak and thus the prior 337 accumulation of juxtaposed SYCP1 molecules between aligned chromosomes may nucleate its 338 cooperative supramolecular assembly. Secondly, α C-end self-assembly occurs through a 339 protonation-induced conformational change triggered by the proton density in the immediate vicinity of DNA³¹ and/or axis protein interactions, thereby coupling assembly to chromosomal 340 341 recruitment.

342

The nascent synapsis generated by SYCP1 self-assembly is stabilised and matured into a full SC through assembly of central element proteins SYCE3, SYCE1, SIX6OS1 and SYCE2-TEX12^{16,22,23,25-29}. Their recruitment is dependent on SYCP1 and is essential for the tripartite structure and meiotic

function of the SC^{22,23,25-28}. Central element assembly likely occurs concomitantly with SYCP1 self-346 347 assembly, rapidly converting the underpinning SYCP1 structural framework into a mature SC. Initial 348 SYCP1 assemblies recruit central element proteins to stabilise the nascent lattice, enabling its 349 growth, and providing a mutually reinforcing cycle that results in full synapsis (Fig. 8c). Central 350 element proteins may provide vertical and longitudinal supports between αN -tetramers that rigidify 351 SYCP1 hemi-lattices and orientate α N-end sites for long-range cooperative head-to-head assembly 352 (Fig. 8d and Supplementary Fig. 10b). They may further act as transverse bridges that connect hemi-353 lattices across the midline to directly reinforce α N-end head-to-head interactions. Initiation factors SYCE3, SYCE1 and SIX6OS1 may act as transverse bridges and vertical supports²²⁻²⁵, whilst SYCE2-354 TEX12 may provide longitudinal supports that enable SC elongation²⁶⁻²⁹. This results in a mature SC in 355 356 which an underlying SYCP1 lattice is structurally supported by the central element. The true 357 molecular roles of SC central element proteins will be revealed upon their structure elucidation, and 358 it will be intriguing to see whether they simply dock onto the SYCP1 lattice or induce structural 359 remodelling upon recruitment.

360

361 Whilst SYCP1 chromosome axis recruitment is retained upon disruption of SC lateral element 362 proteins, synapsis is discontinuous, indicating that chromosome axis structure facilitates the efficient 363 loading of SYCP1 necessary for continuous synapsis^{17,18,20}. This may occur through positioning 364 chromatin loops to achieve a regular spacing of SYCP1 molecules that is compatible with long-range 365 lattice formation. SYCP1 loading may similarly be regulated by the underlying chromatin structure. 366 For example, if both surfaces of α C-end tetramers interact with DNA, they may sit between adjacent 367 nucleosomes and would by spaced apart by the 11 nm nucleosome diameter.

368

How is the SC supramolecular structure efficiently disassembled following its function in meiosis?
 SYCP1 self-assembly is intrinsic to the protein sequence and hence independent of post-translational
 modifications, but phosphorylation has been implicated in SC disassembly³⁴. Whilst there are no

372 clear candidate sites within SYCP1 α N-end or α C-end, phosphorylation of the numerous predicted 373 sites within the C-terminal tail could destabilise axis assembly. Similarly, central element protein 374 phosphorylation could destabilise SYCP1 midline lattice assembly. The molecular features of α N-end 375 and α C-end that achieve cooperative assembly may facilitate the continuous turnover of SYCP1 376 molecules within the SC. Whilst dynamic interchange will normally lead to continual renewal of the 377 SYCP1 lattice, phosphorylation-induced destabilisation of self-assembly sites would shift the balance 378 towards a net loss of molecules and ultimately disassembly.

379

380 SYCP1 fulfils the classic functions of coiled-coil proteins in acting as molecular spacers that scaffold supramolecular assemblies and separate functional units³⁵. SYCP1 imposes a 100 nm synapsis 381 382 between homologous chromosomes, raising the question of why it is necessary to impose an 383 evolutionarily conserved separation between homologues? This distance may be optimal for the 384 maintenance and resolution of meiotic recombination intermediates, and so an answer may lie in differences in recombination in the few meiotic organisms that lack an SC¹. Interestingly, the SC 385 386 central region and central element are approximately 10% narrower in female mice than in males³⁶. 387 This variation can be accommodated by the SYCP1 lattice that we propose through alteration in 388 angulation of α N-end assemblies and α C-dimers (Supplementary Fig. 10c). Furthermore, midline 389 angulation and SC central region width are determined by the frequency of α C-end binding to 390 chromosomes; thus, midline variation could originate from differences in chromosomal axis 391 structure between sexes.

392

Despite evolutionary conservation of the SC ultrastructure, its constituent protein sequences are divergent between vertebrates and lower eukaryotes². Nevertheless, yeast transverse filament protein Zip1 is approximately the same size as SYCP1 and displays similar patterns of conservation and structure prediction. Thus, it is possible that Zip1 adopts a similar structure and self-assembly mechanism through underlying structural conservation. The molecular functions of the SC in recombination, crossover formation and interference remain unknown. Nevertheless, we speculate that its three-dimensional structure may direct these processes by regulating enzymatic access to recombination sites. The ability of coiled-coil proteins to transmit conformational changes recursively may further enable the SC to communicate signals along synapsed chromosomes. As our understanding of the SC structure deepens, its molecular functions will gradually be uncovered, ultimately leading to a complete mechanistic understanding of recombination and crossover formation within the functional architecture of the SC.

406

407 Accession codes

408 Crystallographic structure factors and atomic co-ordinates have been deposited in the Protein Data
409 Bank (PDB) under accession numbers 6F5X, 6F62, 6F63 and 6F64.

410

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422 Author contributions

423 J.M.D. performed the majority of biochemical, biophysical and crystallographic experiments. O.M.D. 424 performed SAXS experiments and analyses. M.R. crystallised truncated α N-end and analysed α N 425 constructs. C.M. and I.U. solved the α C-end crystal structures. O.R.D. solved the α N-end crystal 426 structures and built/refined all structures. S.M. assisted with initial experiments. J.M.D. and O.R.D. 427 designed experiments; O.R.D. wrote the manuscript.

428

429 **Competing financial interests**

430 The authors declare no competing interests.

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- 512

513 Figure Legends

514 Figure 1

515 The obligate tetrameric structure of SYCP1.

516 (a) SYCP1 molecules are bioriented within the synaptonemal complex (SC), with midline N-termini 517 and chromosome-bound C-termini, providing a 100 nm separation between chromosome axes. (b) 518 Sequence analysis of SYCP1 demonstrating the presence of an α -helical core (amino acids 101-783) 519 that is highly conserved at both ends, flanked by unstructured N- and C-terminal tails. Amino acid 520 conservation was calculated amongst vertebrate sequences. The principal protein constructs 521 analysed in this study are indicated along with their amino acid boundaries. An extensive summary 522 of SYCP1 constructs is provided in Supplementary Fig. 1a and biophysical data are compiled in 523 Supplementary Table 1. (c) SEC-MALS analysis; light scattering (LS) and differential refractive index 524 (dRI) are shown as solid and dashed lines respectively, with fitted molecular weights (Mw) plotted as 525 diamonds across elution peaks. SYCP1 α Core (101-783) forms large molecular species of 1-12 MDa, 526 whereas α Core- Δ Ntip (112-783) is a 306 kDa tetramer (theoretical tetramer – 320 kDa) consisting of 527 a 68 kDa α N-tetramer (theoretical tetramer – 76 kDa) and 97 kDa α C-dimer (theoretical dimer – 101 528 kDa). (d) SEC-SAXS P(r) distributions of α Core- Δ Ntip, α N-tetramer and α C-dimer; maximum 529 dimensions (Dmax) and cross-sectional radii (Rc) are indicated. (e) SEC-SAXS P(r) distributions of 530 MBP- α Core- Δ Ntip, MBP- α N-tetramer, MBP- α C-dimer and MBP; intra-MBP and inter-MBP peaks are 531 indicated. (f) SEC-MALS analysis showing that RecE- α N-tetramer is a 208 kDa tetramer (theoretical 532 tetramer- 214 kDa) and GST- α C-dimer is a 157 kDa dimer (theoretical dimer - 160 kDa). (g) Model of 533 the SYCP1 obligate unassembled structure. The SYCP1 α -helical core has a parallel organisation and 534 consists of a 260 Å α N-tetramer that bifurcates into two 645 Å α C-dimer coiled-coils.

535

536 Figure 2

537 Crystal structures of the SYCP1 αN-end head-to-head assembly in open and closed conformations.

538 (a) Crystal structure of SYCP1 αN-end (101-206) demonstrating head-to-head 'dimer of dimers' 539 assembly of two eleven heptad parallel coiled-coils, spanning a total length of 288 Å. The long 540 dimeric coiled-coils are interrupted by a wedge-like structure that splays apart the two α -helices to 541 enable their αN -tip sites to mediate midline head-to-head assembly in an open conformation. The head-to-head interface provides 1,990 Å² buried surface area in addition to 4,520 Å² for each coiled-542 543 coil dimer alone. (b) Crystal structure of truncated SYCP1 α N-end (101-175) demonstrating a similar 544 head-to-head 'dimer of dimers' assembly of two seven heptad parallel coiled-coils, spanning 194 Å, 545 with αN -tips undergoing head-to-head assembly in a closed conformation. The head-to-head interface provides 2,950 Å² buried surface area in addition to the 2,210 Å² for each coiled-coil dimer 546 547 alone. C-terminal interactions of α N-end-truncated chains within the crystal lattice were determined 548 to be artefactual owing to their absence in the αN -end structure and through *in vitro* mutagenesis 549 experiments (M.R. and O.R.D., unpublished data).

550

551 Figure 3

552 Head-to-head assembly interfaces of SYCP1 αN-end.

553 (a-c) Crystal structure of SYCP1 α N-end (101-206). (a) A wedge structure formed of residues I116 and 554 W119 splays apart coiled-coil α -helices to enable their head-to-head assembly. (b-c) The open 555 assembly is formed of one midline and two lateral interfaces. (b) The midline interface (open) is an 556 anti-parallel coiled-coil between symmetry-related chain A copies, with heptad residues L102, V105, 557 L109 and E112. (c) The lateral interface (open) is an anti-parallel association of unique chains A and 558 B, formed of coiled-coil and aromatic stacking interactions of residues L102, L109, Y106 and Y110. 559 (d-f) Crystal structure of truncated SYCP1 α N-end (101-175). (d) Similar to α N-end, a wedge 560 structure of residues 1116 and W119 splays apart α -helices to enable their head-to-head assembly. 561 (e-f) The closed assembly is formed of a hydrophobic core and interfaces that are analogous to the 562 midline and lateral interfaces of the open conformation. (e) The midline interface (closed) is an anti-563 parallel coiled-coil of heptad residues G101, V105, K108 and E112. (f) The lateral interface (closed) is anti-parallel, consisting of coiled-coil and interlaced aromatic stacking interactions of residues L102, L109, Y106 and Y110. (g-h) Cross-sections through the αN-end head-to-head open and closed conformations. (g) The open conformation contains no hydrophobic core and is asymmetrical in nature, with midline chain A copies flanked by two copies of chain B. (h) The closed conformation is formed of symmetry-related chains and contains a hydrophobic core of residues L102, L109 and 1116.

570

571 Figure 4

572 SYCP1 N-terminal self-assembly into higher order structures is mediated by αN-end head-to-head 573 interactions.

574 (a-b) SEC-MALS analysis. (a) SYCP1 α N (101-362) (black, left) forms large molecular species of 3-25 575 MDa, whereas αN-ΔNtip (112-362) (grey) and αN (101-362) V105E L109E (black, right) form 576 tetramers of 118 kDa and 121 kDa respectively (theoretical tetramers – 121 kDa and 126 kDa). (b) 577 SYCP1 α N-end, truncated α N-end and α N-end \sim Ntail are monomeric species of 15 kDa, 10 kDa and 578 23 kDa respectively (theoretical monomers - 13 kDa, 9 kDa and 20 kDa). (c) Model of SYCP1 N-579 terminal self-assembly. SYCP1 α N-ends splay from α N-tetramers and interact head-to-head in the 580 midline to create a continuous lattice-like assembly. (d) SYCP1 N-terminal self-assembly is predicted 581 to be highly cooperative, enabling stable structure formation through a series of individually weak 582 head-to-head associations. This allows transient chromosome associations to be formed and 583 remodelled rapidly, ultimately favouring a single continuous assembly between aligned chromosome 584 axes.

585

586 Figure 5

587 **Crystal structure of the SYCP1 C-terminal tetrameric assembly.**

588 (a) Crystal structure of SYCP1 α C-end (676-770) in crystal form 1, demonstrating an anti-parallel 589 tetrameric assembly of length 142 Å. The structure includes a central tetrameric interface flanked by 590 C703 pinch points that lead to lateral four-helical bundles. N- and C-termini are coloured in green 591 and red respectively. (b) The central tetrameric interface consists of two stacked layers each 592 containing a hydrogen bonding network of pairs of H717, Q720 and Y721 residues. (c-d) The C703 593 pinch point consists of a parallel dimeric coiled-coil (containing C703) flanked by surrounding anti-594 parallel chains. (c) The parallel dimeric coiled-coil is formed of heptad residues D700, C703, I707, 595 M710 and M714 (d) The flanking chains have a distinct angulation at E731 and provide pseudo-cores 596 of loose anti-parallel interactions. (e-g) The lateral four-helical bundle (4HB) is formed of a 597 hydrophobic core and anti-parallel interfaces. (e) The lateral 4HB type 1 interface is an anti-parallel 598 coiled-coil of heptad residues L679, V682 and K686, L753, K757, L760 and K764. (f) The lateral 4HB 599 type 2 interface is an anti-parallel coiled-coil of heptad residues L678, E681, A685, I688 and A692, 600 L745, L749, E752, L756 and E759. (g) Cross-section through the lateral 4HB assembly. A hydrophobic 601 core is formed from residues that also contribute to 4HB anti-parallel interfaces and are predicted to 602 mediate the formation of N- and C-terminal parallel dimeric coiled-coils in the non-assembled 603 conformation. L679 and I688 are the only hydrophobic 4HB residues not also implicated in the 604 putative parallel dimeric coiled-coil structure.

605

606 Figure 6

607 SYCP1 αC-end undergoes pH-induced assembly into an anti-parallel tetramer.

608 (a) SEC-MALS analysis. SYCP1 α C-end (676-770) is a 22 kDa dimer at pH 8.0 (grey) (theoretical dimer 609 - 23 kDa) and a 43 kDa tetramer at pH 5.5 (black) (theoretical tetramer - 46 kDa). (b) SEC-SAXS P(r) 610 distributions of α C-end at pH 5.5 (black) and pH 8.0 (grey). (c-d) SAXS ab initio models of the 611 tetrameric and dimeric conformations of SYCP1 α C-end (676-770) at (c) pH 5.5 and (d) pH 8.0. 612 Averaged models were generated from 20 independent DAMMIF runs with NSD values 0.527 (± 0.014) and 0.513 (± 0.014), and reference model χ^2 values 1.81 and 1.49. The α C-end tetrameric 613 614 crystal structure and a theoretical dimeric coiled-coil were docked into the respective envelopes. (e) 615 SEC-SAXS P(r) distributions of N-terminal, C-terminal and both N- and C-terminal MBP fusions of α C-

616 end at pH 8.0, alongside MBP- α C-end at pH 5.5. (f) SEC-MALS analysis reveals that α C-end tethered 617 dimer forms dimers of 44 kDa and 43 kDa (theoretical dimer of dimers – 47 kDa) at pH 5.5 (black, 618 right) and pH 8.0 (black, left), with an increase in elution volume at pH 5.5. The α C-end (single chain) 619 tetramer at pH 5.5 is shown in grey. (g) SEC-SAXS P(r) distributions of the α C-end tethered dimer at 620 pH 5.5 (black) and pH 8.0 (grey). (h) SEC-MALS analysis of SYCP1 αC-end extended (676-783) point 621 mutants. H717W Y721F (black) forms 40 kDa tetramers at pH 8.0 (theoretical tetramer – 52 kDa). 622 L679A I688A (grey) fails to undergo pH-induced assembly and remains mostly as a 26 kDa dimer 623 (theoretical dimer – 26 kDa) at pH 4.6.

624

625 Figure 7

626 **DNA binding by SYCP1.**

627 (a) Surface electrostatic potential of the SYCP1 α C-end crystal structure (red – electronegative; blue 628 - electropositive). The flat surface of the α C-end structure contains five demarcated basic patches 629 that are separated by approximately 30 Å. (b) EMSA analysing the ability of SYCP1 α C-end (676-770) 630 to interact with linear double-stranded DNA (dsDNA) at pH 8.0 (top) and pH 5.5 (bottom). 631 Uncropped gel images are shown in Supplementary Data Set 1 (c) EMSA of MBP fusions of αC -632 end~Ctail (640-976), Ctail (784-976), αC-dimer~Ctail (358-976) and αCore~Ctail (101-976) with linear 633 dsDNA at pH 8.0. (d) Electron microscopy (EM) analysis of MBP fusions of αC-dimer~Ctail (358-976) 634 and α C-end~Ctail (640-976) in complex with plasmid dsDNA. Scale bars, 50 nm. (e) EMSA of refolded 635 full length SYCP1 (1-976) and Δ Ctail (1-783) with linear dsDNA at pH 8.0. (f) Model of SYCP1 636 chromosomal axis assembly. SYCP1 molecules are initially recruited to chromosomes through Ctail 637 DNA-binding sites. The close proximity of DNA and/or interactions with chromosome axis proteins 638 then triggers protonation-induced assembly of α C-ends into anti-parallel tetramers that bind DNA 639 and thereby reinforce Ctail interactions. This results in the complete coating of the chromosome axis 640 with SYCP1 molecules linked together through U-shaped assemblies that are anchored to 641 chromosomal DNA.

642

643 Figure 8

644 Meiotic chromosome synapsis through SYCP1 self-assembly.

645 (a) Model of the SYCP1 obligate unassembled structure. The α Core consists of a parallel α N-tetramer 646 that splays into two α C-dimers. The α N-tetramer splays at its N-terminus into α N-end self-assembly 647 sites that lead to unstructured Ntails. The α C-dimers terminate as α C-end self-assembly sites, 648 leading to unstructured Ctails that contain DNA-binding sequences. (b) Model of chromosome 649 synapsis by SYCP1. The bifurcating SYCP1 α Core presents pairs of α N-end and α C-end self-assembly 650 sites in the midline and chromosome axis respectively. αN -end sites undergo head-to-head assembly 651 through their α N-tips to provide zipper-like associations that mediate synapsis of SYCP1-coated 652 homologous chromosomes. αC-end sites undergo back-to-back assembly into tetrameric structures 653 that bind directly to DNA within the lateral element and reinforce axis associations of Ctails. 654 Together, these distinct mechanisms of SYCP1 self-assembly generate a supramolecular lattice 655 between meiotic chromosome pairs. (c) Concomitant and mutually reinforcing assembly of SYCP1 656 and central element proteins in SC formation. Initial SYCP1 contacts trigger central element 657 recruitment, enabling growth of the SYCP1 assembly, extending the central element and thereby 658 enabling further SYCP1 growth. (d) Model of the mature SC. The central element may provide 659 vertical and longitudinal supports between SYCP1 α N-tetramers to rigidify SYCP1 hemi-lattices and 660 orientate αN -ends for long range cooperative head-to-head assembly. They may also act as 661 transverse bridges that provide direct connections across the midline to reinforce SYCP1 head-to-662 head interactions. This leads to the formation of a mature SC in which an underlying SYCP1 lattice is 663 stabilised and extended across long distances through central element assembly.

664

Table 1 Data collection and refinement statistics

| | SYCP1 aN-end | SYCP1 αN-end truncated | SYCP1 αC-end | SYCP1 aC-end |
|------------------------------------|---------------------------------------|-------------------------------------|--------------------------------------|-------------------------------------|
| | 101-206 | 101-175 | 676-770 | 676-770 |
| | Open conformation | Closed conformation | Crystal form 1 | Crystal form 2 |
| | (PDB 6F62) | (PDB 6F5X) | (PDB 6F63) | (PDB 6F64) |
| Data collection | i i i i i i i i i i i i i i i i i i i | · · · · · | | · · · · · · |
| Space group | I2 | 1222 | C2 | I4 ₁ 22 |
| Cell dimensions | | | | |
| <i>a</i> , <i>b</i> , <i>c</i> (Å) | 65.67, 37.31, 108.52 | 28.64, 39.38, 165.77 | 233.42, 42.85, 43.69 | 43.38, 43.38, 292.18 |
| α, β, γ (°) | 90.00, 106.66, 90.00 | 90.00, 90.00, 90.00 | 90.00, 93.61, 90.00 | 90.00, 90.00, 90.00 |
| Wavelength (Å) | 0.9282 | 1.7712 | 0.9795 | 0.9795 |
| Resolution (Å) | 34.87-2.06 (2.12-2.06) ^a | 41.44–1.91 (1.95–1.91) ^a | 116.48-2.15 (2.27-2.15) ^a | 42.91-2.48 (2.58-2.48) ^a |
| R _{merge} | 0.071 (0.919) | 0.028 (0.678) | 0.052 (0.695) | 0.080 (2.567) |
| $R_{\rm pim}$ | 0.023 (0.286) | 0.017 (0.541) | 0.032 (0.429) | 0.023 (0.727) |
| $I / \sigma(I)$ | 15.0 (1.8) | 27.9 (1.8) | 12.4 (1.9) | 14.8 (1.5) |
| $CC_{1/2}$ | 0.999 (0.969) | 1.000 (0.839) | 0.998 (0.872) | 1.000 (0.935) |
| Completeness (%) | 99.9 (100.0) | 99.3 (92.1) | 97.4 (88.0) | 99.8 (99.7) |
| Redundancy | 11.0 (11.4) | 5.9 (3.7) | 3.6 (3.5) | 13.2 (13.2) |
| Refinement | | | | |
| Resolution (Å) | 27.23-2.07 | 41.44-1.91 | 58.26-2.15 | 39.63-2.49 |
| UCLA anisotropy (Å) | 2.1, 2.1, 2.6 | 1.9, 2.0, 2.1 | 2.2, 2.3, 2.2 | 2.9, 2.9, 2.5 |
| No. reflections | 12467 | 6754 | 21416 | 4138 |
| $R_{\rm work}$ / $R_{\rm free}$ | 0.2264/0.2441 | 0.2272/0.2392 | 0.2186/0.2526 | 0.2251/0.2517 |
| No. atoms | 1866 | 677 | 3318 | 806 |
| Protein | 1744 | 633 | 3143 | 786 |
| Ligand/ion | 18 | 12 | 0 | 4 |
| Water | 104 | 32 | 175 | 16 |
| B-factors | 42.79 | 58.4 | 46.97 | 60.86 |
| Protein | 42.51 | 57.7 | 47.30 | 60.80 |
| Ligand/ion | 62.91 | 83.9 | N/A | 81.59 |
| Water | 44.02 | 61.7 | 41.20 | 58.92 |
| R.m.s. deviations | | | | |
| Bond lengths (Å) | 0.002 | 0.009 | 0.004 | 0.004 |
| Bond angles (°) | 0.334 | 1.020 | 0.511 | 0.575 |

^aValues in parentheses are for highest-resolution shell.

666 Online Methods

667 Recombinant protein expression and purification

668 Sequences corresponding to regions of human SYCP1 were cloned into pHAT4, pGAT3 or pMAT11 669 vectors for expression as TEV-cleavable N-terminal His₆-, His₆-GST or His₆-MBP fusion proteins 670 respectively. A list of protein constructs, including sequence boundaries, is provided in 671 Supplementary Fig. 1a and Supplementary Table 1. Constructs were expressed in BL21 (DE3) cells (Novagen®), in 2xYT media, induced with 0.5 mM IPTG for 16 hours at 25°C. Cells were lysed by 672 673 sonication in 20 mM Tris pH 8.0, 500 mM KCl, and fusion proteins were purified from clarified lysate 674 through consecutive Ni-NTA (Qiagen), amylose (NEB) or glutathione sepharose (GE Healthcare), and 675 HiTrap Q HP (GE Healthcare) ion exchange chromatography. Affinity tags were removed by 676 incubation with TEV protease and cleaved samples were purified by HiTrap Q HP ion exchange chromatography and size exclusion chromatography (HiLoadTM 16/600 Superdex 200, GE Healthcare) 677 678 in 20 mM Tris pH 8.0, 150 mM KCl, 2 mM DTT. Protein samples were concentrated using Amicon 679 Ultra® 10,000 MWCO centrifugal filter units (Millipore), and were stored at -80°C following flash-680 freezing in liquid nitrogen. Protein samples were analysed by SDS-PAGE with Coomassie staining, 681 and concentrations were determined by UV spectroscopy using a Cary 60 UV spectrophotometer 682 (Agilent) with extinction coefficients and molecular weights calculated by ProtParam 683 (http://web.expasy.org/protparam/).

684

685 Purification of refolded full length SYCP1

Full-length human SYCP1 (amino acids 1-976) was expressed using a pHAT4 vector in Rosetta (DE3) cells, grown in 2xYT media and induced with 0.5 mM IPTG for 3 hours at 37°C. Cells were lysed by sonication in 20 mM Tris pH 8.0, 500 mM NaCl and the insoluble fraction pelleted through centrifugation at 40,000 g for 30 minutes. The resultant pellet was washed in 20 mM Tris pH 8.0, 500 mM NaCl prior to solubilisation in 20 mM Tris pH 8.0, 500 mM NaCl, 8M urea pH 8.0. DNA-containing hydrogels were formed by consecutive dialysis into 20 mM Tris pH 8.0, 500 mM NaCl, 500 mM L- arginine pH 8.0, followed by 20 mM Tris pH 8.0, 500 mM NaCl. Soluble SYCP1 was produced by removal of DNA from the denatured material through ion exchange chromatography, prior to the refolding protocol through dialysis, as described above.

695

696 Circular dichroism (CD) spectroscopy

697 Far UV circular dichroism (CD) spectroscopy data were collected on a Jasco J-810 spectropolarimeter 698 (Institute for Cell and Molecular Biosciences, Newcastle University). CD spectra were recorded in 699 10 mM Na₂HPO₄/NaH₂PO₄ pH 7.5, at protein concentrations between 0.1-0.5 mg/ml, using a 0.2 mm 700 pathlength quartz cuvette (Hellma), at 0.2 nm intervals between 260 and 185 nm at 4°C. Spectra 701 were averaged across nine accumulations, corrected for buffer signal, smoothed and converted to mean residue ellipticity ([0]) (x1000 deg.cm².dmol⁻¹.residue⁻¹). Deconvolution was performed using 702 the CDSSTR algorithm of the Dichroweb server (http://dichroweb.cryst.bbk.ac.uk)³⁷. CD thermal 703 704 denaturation was performed in 20 mM Tris pH 8.0, 150 mM KCl, 2 mM DTT, at protein 705 concentrations between 0.1-0.4 mg/ml, using a 1 mm pathlength quartz cuvette (Hellma). Data were 706 recorded at 222 nm, between 5°C and 95°C, at 0.5°C intervals with ramping rate of 2°C per minute, 707 and were converted to mean residue ellipticity ($[\theta_{222}]$) and plotted as % unfolded ($[\theta]_{222,x^-}$ 708 $[\theta]_{222,5}/([\theta]_{222,5})$. Melting temperatures (Tm) were estimated as the points at which samples 709 are 50% unfolded. SYCP1 α C-end constructs were also analysed in 50 mM NaOAc pH 5.5 or 4.6, 150 710 mM KCl.

711

712 Size-exclusion chromatography multi-angle light scattering (SEC-MALS)

The absolute molar masses of SYCP1 constructs were determined by size-exclusion chromatography multi-angle light scattering (SEC-MALS). Protein samples at >1 mg/ml were loaded onto a Superdex[™] 200 Increase 10/300 GL size exclusion chromatography column (GE Healthcare) in 20 mM Tris pH 8.0, 150 mM KCl, 2 mM DTT, at 0.5 ml/min using an ÄKTA[™] Pure (GE Healthcare). SYCP1 αC-end constructs were also analysed in 50 mM NaOAc pH 5.5 or 4.6, 150 mM KCl, 2 mM DTT. The column outlet was fed into a DAWN[®] HELEOS[™] II MALS detector (Wyatt Technology), followed by an
Optilab[®] T-rEX[™] differential refractometer (Wyatt Technology). Light scattering and differential
refractive index data were collected and analysed using ASTRA[®] 6 software (Wyatt Technology).
Molecular weights and estimated errors were calculated across eluted peaks by extrapolation from
Zimm plots using a dn/dc value of 0.1850 ml/g. SEC-MALS data are presented with light scattering
(LS) and differential refractive index (dRI) profiles, with fitted molecular weights (M_w) plotted across

725

726 Electrophoretic mobility shift assay (EMSA)

SYCP1 protein constructs were incubated with 25 or 32 µM (per base pair) 470 or 75 bp linear dsDNA
substrate at concentrations indicated, in 20 mM Tris pH 8.0, 150 mM KCl or 50 mM NaOAc pH 5.5,
150 mM KCl, for 5 minutes at 4°C. Glycerol was added at a final concentration of 3% and samples
were analysed by electrophoresis on a 0.5% (w/v) agarose gel in 0.5x TBE pH 8.0 or 25 mM GABA pH
5.5 at 20V for 4 hours at 4°C. DNA was detected by SYBR™ safe (ThermoFisher).

732

733 Size-exclusion chromatography small-angle X-ray scattering (SEC-SAXS)

734 SEC-SAXS experiments were performed at beamline B21 of the Diamond Light Source synchrotron 735 facility (Oxfordshire, UK). Protein samples at concentrations >10 mg/ml were loaded onto a 736 Superdex[™] 200 Increase 10/300 GL size exclusion chromatography column (GE Healthcare) in 20 737 mM Tris pH 8.0, 150 mM KCl at 0.5 ml/min using an Agilent 1200 HPLC system. SYCP1 α C-end 738 constructs were also analysed in 50 mM NaOAc pH 5.5 or 4.6, 150 mM KCl. The column outlet was 739 fed into the experimental cell, and SAXS data were recorded at 12.4 keV, detector distance 4.014 m, 740 in 3.0 s frames. Data were subtracted and averaged, and analysed for Guinier region Rg and cross-741 sectional Rg (Rc) using ScAtter 3.0 (http://www.bioisis.net). Approximate parameters for real space 742 analysis were determined using the server www.bayesapp.org, and P(r) distributions fitted using 743 PRIMUS³⁸. Ab initio modelling was performed using DAMMIF³⁹ run in interactive mode with random chain selected as expected shape. 10-20 independent runs were performed and averaged. Crystal

745 structures and models were docked into DAMAVER molecular envelopes using SUPCOMB⁴⁰.

746

747 Electron Microscopy

Electron microscopy (EM) was performed using an FEI Philips CM100 transmission electron microscope at the Electron Microscopy Research Services, Newcastle University. MBP fusion SYCP1 samples at 10 μ M were incubated with 100 μ M (per base pair) plasmid double-stranded DNA in 20 mM Tris pH 8.0, 250 mM KCl for 10 minutes, and applied to carbon-coated EM grids. Negative staining was performed using 2% (wt/vol) uranyl acetate.

753

754 Protein crystallisation and X-ray structure solution of SYCP1 αN-end (101-206)

755 SYCP1 α N-end (101-206) protein crystals were obtained through vapour diffusion in hanging drops, 756 by mixing 200 nl of protein at 10 mg/ml with 100 nl of crystallisation solution (100 mM MES pH 6.0, 757 40% (v/v) MPD) and equilibrating at 4°C for 4-9 days. Crystals were flash frozen in liquid nitrogen. X-758 ray diffraction data were collected at 0.9282 Å, 100 K, as 2000 consecutive 0.10° frames of 0.050 s 759 exposure on a Pilatus 6M detector at beamline IO4-1 of the Diamond Light Source synchrotron facility (Oxfordshire, UK). Data were indexed and integrated in XDS⁴¹ using AutoPROC⁴²; datasets 760 from three crystals were scaled together using XSCALE⁴³ and then merged in Aimless⁴⁴. Crystals 761 belong to monoclinic spacegroup I2 (cell dimensions a = 65.67 Å, b = 37.31 Å, c = 108.52 Å, α = 90°, β 762 763 = 106.66°, γ = 90°), with two SYCP1 chains per asymmetric unit. Data were corrected for anisotropy using the UCLA diffraction anisotropy server (https://services.mbi.ucla.edu/anisoscale/)⁴⁵, imposing 764 765 anisotropic limits of 2.1 Å, 2.1 Å, 2.6 Å, with principal components of 24.09 Å², 6.01 Å² and -20.19 Å². 766 Structure solution was achieved using AMPLE⁴⁶ on the CCP4 online web server (https://www.ccp4.ac.uk/ccp4online/), through molecular replacement of Quark ab initio model 767 decoys⁴⁷, with auto-tracing and rebuilding in SHELX E and ARP/wARP. Phase improvement was 768 achieved through iterative re-building by PHENIX Autobuild⁴⁸. The structure was completed through 769

manual model building in Coot and refinement using PHENIX refine⁴⁸, with the addition of two 2methyl-2,4-pentanediol (MPD) ligands and two chloride ions. Refinement was performed using
isotropic atomic displacement parameters with riding hydrogens. The structure was refined against
anisotropy corrected 2.07 Å data to R and R_{free} values of 0.2264 and 0.2441 respectively, with 100%
of residues within the favoured regions of the Ramachandran plot, clashscore of 3.05 and overall
MolProbity score of 1.10.

776

777 Protein crystallisation and X-ray structure solution of truncated SYCP1-αN-end (101-175)

778 SYCP1 α N-end-tr (101-175) protein crystals were obtained through vapour diffusion in hanging 779 drops, by mixing 1 μ l of protein at 10 mg/ml with 1 μ l of crystallisation solution (140 mM NaCl, 70 780 mM Na/K phosphate pH 6.2, 35% (v/v) PEG200) and equilibrating at 20°C for 4-9 days. Crystals were 781 soaked for 30 minutes in crystallisation solution containing 40% (v/v) PEG200 and 100 mM NaI, prior 782 to flash freezing in liquid nitrogen. X-ray diffraction data were collected at 1.7712 Å, 100 K, as 2000 783 consecutive 0.10° frames of 0.050 s exposure on a Pilatus 6M detector at beamline IO2 of the 784 Diamond Light Source synchrotron facility (Oxfordshire, UK). Data were indexed, integrated and scaled in XDS⁴¹ and XSCALE⁴³, and merged in Aimless⁴⁴. Crystals belong to orthorhombic spacegroup 785 786 1222 (cell dimensions a = 28.64 Å, b = 39.38 Å, c = 165.77 Å, α = 90°, β = 90°, γ = 90°), with one SYCP1 787 chain per asymmetric unit. SAD structure solution was achieved through identification of five 788 putative iodide sites and secondary structure auto-tracing by SHELX C/D/E, utilising the HKL2MAP interface⁴⁹. Phase improvement was achieved through iterative re-building by PHENIX Autobuild⁴⁸. 789 790 Data were corrected for anisotropy using the UCLA diffraction anisotropy server 791 (https://services.mbi.ucla.edu/anisoscale/)⁴⁵, imposing anisotropic limits of 1.9 Å, 2.0 Å, 2.1 Å, with principal components of 13.25 $Å^2$, 0.78 $Å^2$ and -14.08 $Å^2$. The structure was completed through 792 manual model building in Coot and refinement using PHENIX refine⁴⁸, with the truncation to two 793 iodide sites (based on anomalous difference map peaks) and the addition of a triethylene glycol 794 795 ligand (PGE). Refinement was performed using isotropic atomic displacement parameters with five

TLS groups. The structure was refined against anisotropy corrected 1.91 Å data to R and R_{free} values
 of 0.2272 and 0.2392 respectively, with 100% of residues within the favoured regions of the
 Ramachandran plot, clashscore of 6.77 and overall MolProbity score of 1.37.

799

800 Protein crystallisation and X-ray structure solution of SYCP1 αC-end (676-770) crystal form 1

801 SYCP1 α C-end (676-770) protein crystals were obtained through vapour diffusion in hanging drops, 802 by mixing 100 nl of protein at 31 mg/ml with 100 nl of crystallisation solution (3.5 M sodium formate 803 pH 7.0) and equilibrating at 20°C for 2 months. Crystals were soaked in a cryoprotectant solution of 6 804 M sodium formate pH 7.0 and flash frozen in liquid nitrogen. X-ray diffraction data were collected at 805 0.9795 Å, 100 K, as 2000 consecutive 0.10° frames of 0.080 s exposure on a Pilatus 6M detector at 806 beamline IO2 of the Diamond Light Source synchrotron facility (Oxfordshire, UK). Data were indexed and integrated in XDS⁴¹, and scaled and merged in Aimless⁴⁴, using AutoPROC⁴². Crystals belong to 807 monoclinic spacegroup C2 (cell dimensions a = 233.42 Å, b = 42.85 Å, c = 43.69 Å, α = 90°, β = 93.61°, 808 809 $y = 90^{\circ}$), with four SYCP1 chains per asymmetric unit. Structure solution was achieved through fragment-based molecular replacement using ARCIMBOLDO SHREDDER⁵⁰, a program that derives 810 small models from distant homologs, decomposes and refines the fragments against PHASER's⁵¹ gyre 811 and gimble functions⁵², and combines partial solutions⁵³ for expansion through density modification 812 and main chain tracing with SHELXE⁵⁴ to generate the full structure. The SYCP1- α C-end I4₁22 813 814 structure (crystal form 2) was used as a starting template for generating 74 models containing 99 815 amino acids each. A phase set combining 25 partial solutions was expanded into a full solution, 816 recognisable by a correlation coefficient of 48.2%. Phase improvement was achieved through 817 iterative re-building by PHENIX Autobuild⁴⁸. Data were corrected for anisotropy using the UCLA diffraction anisotropy server (https://services.mbi.ucla.edu/anisoscale/)⁴⁵, imposing anisotropic 818 limits of 2.2 Å, 2.3 Å, 2.2 Å, with principal components of 18.46 $Å^2$, 3.44 $Å^2$ and -21.90 $Å^2$. The 819 820 structure was completed through manual model building in Coot and refinement using PHENIX 821 refine⁴⁸. Refinement was performed using isotropic atomic displacement parameters with seven TLS groups per chain. The structure was refined against anisotropy corrected 2.15 Å data to R and R_{free}
values of 0.2186 and 0.2526 respectively, with 100% of residues within the favoured regions of the
Ramachandran plot, clashscore of 6.86 and overall MolProbity score of 1.38.

825

826 Protein crystallisation and X-ray structure solution of SYCP1 αC-end (676-770) crystal form 2

827 SYCP1 α C-end (676-770) protein crystals were obtained through vapour diffusion in hanging drops, 828 by mixing 100 nl of protein at 15 mg/ml with 100 nl of crystallisation solution (0.1 M sodium 829 cacodylate pH 6.5, 1.4M sodium acetate) and equilibrating at 20°C for 2 months. Crystals were 830 soaked in a cryoprotectant solution of 0.1 M sodium cacodylate pH 6.5, 1.4 M sodium acetate, 20% 831 PEG400 and flash frozen in liquid nitrogen. X-ray diffraction data were collected at 0.9795 Å, 100 K, 832 as 2000 consecutive 0.10° frames of 0.080 s exposure on a Pilatus 6M detector at beamline IO2 of 833 the Diamond Light Source synchrotron facility (Oxfordshire, UK). Data were indexed and integrated in XDS^{41} using AutoPROC⁴², scaled in $XSCALE^{43}$ and scaled merged in Aimless⁴⁴. Crystals belong to 834 tetragonal spacegroup $14_{1}22$ (cell dimensions a = 43.38 Å, b = 43.38 Å, c = 292.18 Å, α = 90°, β = 90°, 835 836 $y = 90^{\circ}$), with one SYCP1 chain per asymmetric unit. Structure solution was achieved through fragment-based molecular replacement using ARCIMBOLDO LITE⁵⁵. Substructures made up of two 837 838 ideal polyalanine helices of 30 residues each were located with PHASER, profiting from its features 839 for small fragments, and were extended with SHELXE within the Arcimboldo mode for coiled coil 840 structures⁵⁶. A correct solution was identified by a SHELXE Correlation Coefficient of 40.5%. Phase improvement was achieved through iterative re-building by PHENIX Autobuild⁴⁸. Data were 841 842 corrected for UCLA diffraction anisotropy using the anisotropy server (https://services.mbi.ucla.edu/anisoscale/)⁴⁵, imposing anisotropic limits of 2.9 Å, 2.9 Å, 2.5 Å, with 843 principal components of 16.05 $Å^2$, 16.05 $Å^2$ and -32.09 $Å^2$. The structure was completed through 844 manual model building in Coot and refinement using PHENIX refine⁴⁸, with the addition of one 845 846 acetate ligand. Refinement was performed using isotropic atomic displacement parameters with riding hydrogens. The structure was refined against anisotropy corrected 2.49 Å data to R and R_{tree} 847

values of 0.2251 and 0.2517 respectively, with 100% of residues within the favoured regions of the

849 Ramachandran plot, clashscore of 1.24 and overall MolProbity score of 0.84.

850

851 **Protein sequence and structure analysis**

Conservation of SYCP1 amongst vertebrate sequences was calculated as per residue scores for the
full SYCP1 sequence and the αC-end structure by ConSurf (http://consurf.tau.ac.il/), and secondary
structure prediction was performed by JNet (http://www.compbio.dundee.ac.uk/www-jpred/).
Protein structures were superposed and rsmd values calculated by PHENIX superpose⁴⁸. Structural
assemblies were analysed by PISA. Molecular structure images were generated using the PyMOL
Molecular Graphics System, Version 1.3 Schrödinger, LLC.

858

859 Data availability

860 Crystallographic structure factors and atomic co-ordinates have been deposited in the Protein Data

861 Bank (PDB) under accession numbers 6F5X, 6F62, 6F63 and 6F64. Uncropped gel images are shown

in Supplementary Data Set 1. All other data are available from the corresponding author upon

reasonable request. A Life Sciences Reporting Summary for this article is available.

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d





С





L749, L753, L756, L760, V763, L767



20

0

13.5

14.0

14.5

Elution volume (ml)

26 kDa

Dimer

15.0

15.5

0.4 0.2 0 0 50 100 150 200 250 Interatomic distance (Å)

g





Meiotic

(paternal)

Ň C chromatin loops