Meiotic Cohesin SMC1β Provides Prophase I Centromeric Cohesion and Is Required for Multiple Synapsis-Associated Functions

Uddipta Biswas1, Cornelia Wetzker1, Julian Lange2, Eleni G. Christodoulou3, Michael Seifert3, Andreas Beyer3,4*, Rolf Jessberger1*

1 Institute of Physiological Chemistry, Medical Faculty Carl Gustav Carus, Technische Universität Dresden, Dresden, Germany, 2 Molecular Biology Program, Memorial Sloan-Kettering Cancer Center, New York, New York, United States of America, 3 Biotechnology Center, TU Dresden, Dresden, Germany, 4 Center for Regenerative Therapies Dresden, Dresden, Germany

Abstract

Cohesin subunit SMC1β is specific and essential for meiosis. Previous studies showed functions of SMC1β in determining the axis-loop structure of synaptonemal complexes (SCs), in providing sister chromatid cohesion (SCC) in metaphase I and thereafter, in protecting telomere structure, and in synapsis. However, several central questions remained unanswered and concern roles of SMC1β in SCC and synapsis and processes related to these two processes. Here we show that SMC1β substantially supports prophase I SCC at centromeres but not along chromosome arms. Arm cohesion and some of centromeric cohesion in prophase I are provided by non-phosphorylated SMC1α. Besides supporting synapsis of autosomes, SMC1β is also required for synapsis and silencing of sex chromosomes. In absence of SMC1β, the silencing factor H2AX remains associated with asynapsed autosomes and fails to localize to sex chromosomes. Microarray expression studies revealed up-regulated sex chromosome genes and many down-regulated autosomal genes. SMC1β is further required for non-homologous chromosome associations observed in absence of SPO11 and thus of programmed double-strand breaks. These breaks are properly generated in Smc1β−/− spermatocytes, but their repair is delayed on asynapsed chromosomes. SMC1α alone cannot support non-homologous associations. Together with previous knowledge, three main functions of SMC1β have emerged, which have multiple consequences for spermatocyte biology: generation of the loop-axis architecture of SCs, homologous and non-homologous synapsis, and SCC starting in early prophase I.

Introduction

Meiosis requires unique chromosome structures and dynamics, which are most prominent in the first of the two meiotic divisions (for reviews on aspects of meiosis relevant to this study see [1,2,3,4,5,6,7,8,9]). During premeiotic replication, two pairs of sister chromatids are formed from the two homologous chromosomes. Within each pair, the two sister chromatids are linked through sister chromatid cohesion (SCC). In early meiotic prophase I the two pairs of sister chromatids form axial elements (AEs) through association with proteins like SYCP2 and SYCP3. The AEs start to pair and synapse, and full synapsis is reached in pachynema. The synaptonemal complex (SC) is generated, which in addition to AEs includes transverse elements made of SYCP4, SYCE1 and other proteins. Consequently, SYCP1 serves as a marker for synapsis. Other proteins such as HORMAD1 are displaced from chromosomes upon synapsis and thus their association with chromosomes indicates asynapsed chromosomes or chromosomal regions [10,11]. We use the terms “asynapsed” for never synapsed, and the term “desynapsed” for lost synapsis.

Homologous pairing requires programmed DNA double-strand breaks (DSBs), generated by the type II topoisomerase-like enzyme SPO11. SPO11-deficient mice of both sexes are infertile and Spo11−/− spermatocytes die in late zygonema/early pachynema. In Spo11−/− meiocytes, these DSBs are not introduced into DNA, and homologous synopsis of the AEs, which still form, is defective. Non-homologous associations between multiple pairs of sister chromatids are observed [12,13]. DSBs may give rise to crossovers and chiasmata. The processing of DSBs requires recombinases RAD51 and the meiosis-specific DMC1. Characteristic RAD51 and DMC1 foci are formed during leptonema and early zygonema at sites of DSBs and disappear progressively during zygonema and early pachynema as repair proceeds.

In spermatocytes, the X and Y sex chromosomes behave uniquely, for they are mostly non-homologous and only synapse at a short, about 700 kbp homologous region called the pseudoautosomal region (PAR) [14,15]. Within the PAR, meiotic recombination and chiasma formation take place [16], although repair foci appear later on PAR than on autosomes [17]. Compared to
The generation of mammalian gametes through meiosis comprises two subsequent cell divisions. The first division, meiosis I, features highly specific chromosome structures, and behavior, and requires distinct sets of chromosome-associated proteins. Cohesin proteins, of which some are meiosis-specific, are essential for meiosis, but their particular roles in meiosis are incompletely understood.

We show here that SMC1b, a meiosis-specific cohesin, serves key functions already in prophase of meiosis I: SMC1b contributes to keeping sister chromatids in cohesion at their centromeres and supports synopsis of the four sister chromatids present in these cells. SMC1b is required for the synopsis of the X and Y sex chromosomes. The failure of autosomes to properly synapse in absence of SMC1b causes extensive alterations in gene expression. This leads to expression of sex chromosome-linked genes, which are lethal at this stage, explaining the death of spermatocytes in mid-prophase I. Together with the analyses of other cohesin proteins and of phosphorylated forms of SMC3 and SMC1z, this paper describes hitherto undescribed properties and functions of meiotic cohesin in sister chromatid cohesion and synopsis.

Cohesin executes additional functions besides in SCC. In somatic cells cohesin is required for efficient DSB repair through homologous recombination. Cohesin is recruited to sites of DNA damage, and its SMC proteins become phosphorylated in response to DNA damage (reviewed in [38,39,40,41]). Cohesin also regulates gene expression (reviewed in [42,43,44,45]) and is thought to promote chromosome looping to support promoter-enhancer interactions. Cohesin may assist the CTCF-mediated insulator function, and often binds to sites that overlap with CTCF binding sites, or to binding sites of other factors.

Together, it remained elusive whether SMC1b contributes to SCC in early meiosis, whether SMC1b plays a particular role in sex body formation and MSCI, whether it acts in induction and processing of DNA DSB repair foci, and whether it affects meiotic gene expression. Our analyses of these features elucidate the relationship between SMC1b and SMC1z, and reveal a consistent concept of a major function of SMC1b in two fundamental processes, SCC and synopsis.
primary antibody incubation in PBSTG for 90 min at 37°C or overnight at 4°C. Antibodies were used as follows: mouse anti-SYCP3 (hybridoma supernatant), guinea-pig anti-HORMAD1 (1:100, a kind gift from Dr. A. Tōth, TUD, Dresden, Germany), anti RAD21L (a kind gift from Dr. T. Hiranò, Riken, Japan; [49]), yH2AX (1:100, Upstate, 05-636), SUMO-1 (1:200 Cell Signaling Technology; #4930), rabbit anti-SMC3 (1:100), rabbit anti-STAG3 (1:100), rabbit anti SMC1α (1:20) raised against the C-terminus, rabbit and rabbit or mouse monclonal anti SMC1β as described [36,50].

Fluorescence In Situ Hybridization

FISH was performed according to the manufacturer’s protocol (MetaSystems GmbH). Ten μl of probe mixture was put on slides and covered with a coverslip. Both sample and probe were denatured simultaneously by heating on a hotplate at 75°C for 2 min, followed by incubation of slides in a humidified chamber at 37°C overnight for hybridization. Post hybridization washes were performed with 0.5× SSC at 72°C for 2 min and 2× SSC, 0.05% Tween-20 at room temperature for 30 sec. Slides were incubated with DAPI and signals were analyzed. Cells were identified based on their DAPI staining and 5–7 pericentric heterochromatin domains. Wash and Smc1β−/− pachytene cells were characterized by 5–7 pericentric heterochromatin domains [51,52].

Measurement of SPO11–oligonucleotide complexes and western blot analysis

For measurement of SPO11–oligonucleotide complexes, both tests from each mouse were used per experiment, that is, littermate comparisons were made on a per-testis basis. Testis extract preparation, immunoprecipitation and western blot analysis were performed essentially as described [33]. Testes were decapsulated, then lysed in 800 μl lysis buffer (1% Triton X-100, 400 mM NaCl, 25 mM HEPES-NaOH at pH 7.4, 5 mM EDTA). Lysates were centrifuged at 100,000 rpm (353,040 × g) for 25 min in a TLA100.2 rotor. Supernatants were incubated with anti-mouse SPO11 antibody 180 (5 μg per pair of testes) at 4°C for 1 h, followed by addition of 40 μl protein-A–agarose beads (Roche) and incubation overnight. Beads were washed three times with IP buffer (1% Triton X-100, 150 mM NaCl, 15 mM Tris-HCl at pH 8.0). Immunoprecipitates were eluted with Laemmli sample buffer and diluted six-fold in IP buffer. Eluates were incubated with an additional 5 μg anti-mouse SPO11 antibody 180 at 4°C for 1 h, followed by addition of 40 μl protein-A–agarose beads and incubation overnight. Beads were washed three times with IP buffer and twice with buffer NEB4 (New England BioLabs). SPO11-oligonucleotide were radio-labelled at 37°C for 1 h using terminal deoxynucleotidyl transferase (Fermentas) and [α-32P]dCTP. Beads were washed three times with IP buffer, boiled in Laemmli sample buffer, and fractionated on 8% SDS–PAGE. Immunoprecipitates were transferred to a PDVF membrane by semi-dry transfer (Bio-Rad). Radiolabelled species were detected and quantified with Fuji phosphor screens and ImageGage software. For western blot analysis, membranes were probed with anti-mouse SPO11 antibody 180 (1:2,000 in PBS containing 0.1% Tween 20 and 5% non-fat dry milk), then horseradish-peroxidase-conjugated protein A (Abcam; 1:10,000 in PBS containing 0.1% Tween 20 and 5% nonfat dry milk), and detected using the ECL+ reagent (GE Healthcare).

Isolation of RNA

Total testis RNA was extracted from four pairs of 12-day-old and 16-day-old Smc1β−/− and wild-type control littermate mice with TRIzol reagent (Invitrogen Inc.) according to the manufacturer’s protocol. Briefly, testes were surgically removed and the Tunicat albignascia was detached. Testes were dounce-homogenized in TRIzol reagent prior to phenol-chloroform extraction of RNA. The integrity of the RNA solubilized in water was confirmed by use of the BioAnalyzer (Bio-Rad).

Microarray procedures

Microarray experiments were performed in quadruplicates with littermate mouse pairs. 800 ng of RNA were applied to microarray analysis on the One-Color Microarray-Based Gene Expression Analysis System (Agilent Technologies Inc.) according to the manufacturer's protocols. Briefly, cDNA was synthesized using M-MLV Reverse Transcriptase followed by labeling with the fluorescent dye Cy3. Amplified cRNA was purified with RNaseasy mini spin columns (Qiagen Inc.) followed by hybridization to 4×44K mouse whole genome microarrays (Agilent Technologies). Microarray experiments were performed in quadruplicates with littermate mouse pairs. Scanning of the chips was performed using the Agilent Microarray Scanner. Normalization of the data was performed using the software GeneSpring GX 7.3 (Agilent Technologies). The normalization algorithm was scaling to the median of all samples. No baseline transformation was performed.

Quantitative reverse-transcription PCR

Total testis RNA of Smc1β−/− and wt mice was DNase-treated (1.25 U DNase/μg RNA) for 30 min at 37°C and reversely transcribed using Superscript II Reverse Transcriptase (Invitrogen). For PCR amplification, full-length mRNA sequence primer pairs were designed using the web-based tool Primer3 version 3.0 [54] to yield 120 to 180 bp long intron-spanning PCR products. For each reaction, 1 μl of diluted cDNA generated from 40 ng of RNA was amplified in a 10 μl reaction volume using the QuantiTect SYBR Green PCR kit (QIAGEN) in a Rotorgene 3000 thermal cycler (Corbett Research Inc.). Reactions were performed in duplicates and two or three mouse pairs were analyzed for each gene. Average mRNA levels of beta-actin (Actb) and glyceraldehyde-3-phosphate dehydrogenase (Gapdh) were used for normalization. PCR primer sequences are shown in Supplementary Table S1.

Spatial clustering of differentially regulated genes

1. Using the Wilcoxon test.

Chromosomal regions enriched for up- or down-regulated genes were identified using the two-sided Wilcoxon rank sum test [55], as implemented in R (function wilcox.test.R). A 10-gene sliding window approach was used, testing if genes inside the given window are more strongly up- (or down-) regulated than expected by chance. Unlike the t-test, the non-parametric Wilcoxon test makes no assumptions about the distribution of the data. In order to quantify the global significance (accounting for multiple testing) fold-changes were permuted 10 times across the whole genome and the same approach was applied again. Observed Wilcoxon scores were compared to the permutation-based scores in order to determine False Discovery Rates (FDR). Reported clusters have a FDR less than 0.2.

2. Using Hidden Markov Models.

Average log-ratio profiles of gene expression levels for Smc1β−/− compared to wt testes have also been analyzed using a three-state Hidden Markov Model (HMM) with Gaussian emission distributions as described [56]. Hidden states were determined using a Bayesian Baum-Welch algorithm: decoding of the most likely expression state of each gene (under-expressed, unchanged, or over-expressed in Smc1β−/−) has been done using state-posterior decoding assigning each gene to its most probable underlying expression state. Groups
of contiguous genes identified as underexpressed or groups of contiguous genes identified as being overexpressed where considered as domains. Applications of the HNM to permuted log-ratio profiles (100 permutations per chromosome) identified that domains of length greater or equal than two have been observed significantly more frequently in the original data than in the permuted profiles.

**MiRNA expression profiling**

MiRNA expression analysis was performed using the miR-plex microarrays (Milenyi Biotec GmbH) according to the manufacturer’s protocol. Microarrays carried DNA oligonucleotides with a reverse-complementary sequence of mature mouse-miRNAs. The 50th percentile of background intensity values was applied for data normalization and dye bias was corrected by Lowess normalization. The normalized mean ratios of Smc1β−/− versus wt were calculated.

The microarray mutant versus wild-type log fold changes were normalized according to the more stable RT-qPCR measurements of 20 selected genes. The respective regression model, which was applied to all the genes, is the following, Y corresponds to the RT-qPCR measurement and x to the microarray fold change. The normalized data have been subjected to statistical clustering testing by applying a two-sided Wilcoxon rank sum test [55] as implemented in R (function wilcox.test.R). A 10 gene sliding window approach was used, testing the null hypothesis that the fold changes of the genes in each window cluster better than in the rest of the genome. Application of the same approach to permuted log-ratio profiles (10 whole genome permutations) identified that the detected clusters have been observed significantly more frequently in the original data than in the permuted ones. The respective false discovery rates (FDR) are plotted in dots across chromosomes 1-X.

**Fluorescence-activated cell sorting (FACS)**

Single-cell suspensions of testes were resuspended in FACS buffer (HBSS supplemented with 20 mM HEPES (pH 7.4), 1.2 mM MgSO4, 1.3 mM CaCl2, 6.6 mM sodium pyruvate, 0.05% lactate, glutamine, and 1% fetal calf serum) at a density of 2 million cells/ml. Bis-benzamide Hoechst33342 (5 µg/ml, Hoechst) was added before incubation of the cells for 1 h at 32°C. For exclusion of dead cells, propidium iodide (PI, 2 µg/ml) was added before FACS analysis. Cells were analyzed using a LSRII flow cytometer (BD Biosciences) using the FACS Diva software version (BD Biosciences, version 6.1.3). Hoechst was excited with the 355 nm UV laser and emission filters Emerald (585/42 nm, LP) and Alexa 350 (505 nm) were applied. PI was excited with the 405 nm blue laser and emission was filtered using PE (685 nm).

**Results**

**Autosomal asynapsis and defective sex body formation in Smc1β−/− spermatocytes**

The current study aimed at elucidating in detail the role of Smc1β in SCC and synapsis, and therefore a number of meiotic processes and features were analyzed, all of which relate to these two processes.

Previously, we showed that loss of Smc1β causes partial asynapsis of many autosomes and total asynapsis of a few [36,57]. In wild-type (wt) pachytene cells only the sex body with its large unpaired regions stains intensely for γH2AX. In Smc1β−/− spermatocytes that we classified as early/mid pachytene small clouds of γH2AX staining were observed on many chromosomes [36]. The most advanced Smc1β−/− spermatocytes show widespread SYCP1 staining, little HORMAD1 staining, and a very low percentage of cells express H1t, a mid-pachytene marker. We classified the early/mid pachytene Smc1β−/− spermatocytes based on the highest degree of axes compaction that can be observed. The sex body was mostly absent Smc1β−/− spermatocytes, and if a prominent γH2AX cloud was seen, it was one among several γH2AX clouds, and thus the occurrence of a sex body remained uncertain.

To initiate the present study, a thorough analysis of spermatocyte asynapsis was performed using various combinations of staining for SYCP1, SYCP3, γH2AX, HORMAD1, and SUMO-1 (Fig. 1, Suppl. Fig. S1). SYCP1 localizes to synapsed regions of prophase I chromosomes, SYCP3 stains asynapsed and synapsed axial elements (AEs), and γH2AX localizes to synapsed regions, which are also transcriptionally silenced regions, and to DNA DSBs. HORMAD1 specifically stains asynapsed chromosomes or asynapsed regions of chromosomes and disappears from chromosomes upon formation of the SC [10,11], and SUMO-1 localizes preferentially to asynapsed regions including the sex body [58,59,60]. The level of asynapsis varies considerably between individual Smc1β−/− cells as illustrated in Fig. 1A, but there were virtually no early to mid pachytene Smc1β−/− spermatocytes without at least partial asynapsis. The average number of partially asynapsed chromosomes was 1.8+/−0.9 per cell, the average number of entirely asynapsed chromosomes including the sex chromosomes was 6.8+/−1.7 per cell. Except the unsynapsed regions of the X and Y chromosomes, there are no asynapsed chromosomes in wt pachytene cells.

Synapsis of X and Y chromosomes is restricted to the short PAR. In the vast majority of Smc1β−/− spermatocytes neither the PAR-synapsed X and Y chromosomes nor the typical sex body was detected by either HORMAD1, γH2AX, or SUMO-1 staining combined with labeling the SC (SYCP1) or the AEs (SYCP3) (Fig. 1B, 1C). HORMAD1, γH2AX or SUMO-1 localized to several asynapsed chromosomes. Quantification of γH2AX staining revealed that a typical sex body could not be observed. Rather, two (app. 25% of the cells) or more (app. 72% of the cells) γH2AX clouds or multiple patches, of which occasionally (app. 3% of cells) one was more prominent, were found. In addition, we also evaluated X-Y pairing using fluorescence in situ hybridization (FISH) (Fig. 1D). Pachytene cells were identified by light microscopy based on their characteristic DAPI-stained pattern of heterochromatin [51,52] (Suppl. Fig. S1D). In 98% of the Smc1β−/− cells (59 of 60), the two FISH signals were clearly apart, indicating that the X and Y were not synapsed within the PAR, consistent with the absence of a sex body in which both chromosomes ought to be present. In one of the cells the signals were very near each other, but FISH did not allow us to determine if PAR synapsis had occurred. In wt, 90% of the cells (36 of 40) showed partially overlapping signals, and in the remaining four cells, the signals did not visibly overlap but were very close to each other.

This indicates that X/Y synapsis and sex body formation fails in the absence of Smc1β, and that the remaining Smc1β-type cohesin complexes alone are not sufficient for either process. This also implies that Smc1β−/− spermatocytes contain at least 21 microscopically visible chromosomes, i.e., at least 19 fully or partially formed SCs and the two separate sex chromosome AEs. Thus every cell has at least one pair of fully asynapsed chromosomes – the X and Y chromosomes –, and often many more, in addition to some partial asynapsis.

**Disturbed patterns of gene expression in Smc1β−/− spermatocytes**

The previous section showed significant levels of autosomal asynapsis and a failure to synapse the X and Y chromosomes in
Figure 1. Synapsis defects of autosomes and sex chromosomes of Smc1b−/− spermatocytes. A. Wt and Smc1b−/− spermatocyte chromosome spreads stained with anti SYCP1 for synapsed chromosomes and anti HORMAD1 for unsynapsed chromosomes or chromosomal regions. Three examples of Smc1b−/− cells illustrate different degrees of asynapsis (n = 60). B. Staining of the sex body in wt and Smc1b−/− spermatocyte chromosome spreads using anti SYCP3 for AEs, anti γH2AX for the sex body or unsynapsed chromatin domains, and anti HORMAD1 for the unsynapsed sex chromosome axes or unsynapsed autosomes. C. Staining of the sex body in wt and Smc1b−/− spermatocyte chromosome spreads using anti SYCP3 for AEs, and anti SUMO-1 for the sex body or unsynapsed chromatin domains. D. FISH of wt (n = 40) and Smc1b−/− (n = 60) spermatocytes using a green (X) and red (Y) sex chromosome-specific probe.

Cohesin SMC1β in Synapsis and Prophase I Cohesion

absence of SMC1β. High levels of autosomal asynapsis correlate with decreased expression of autosomal genes, i.e. MSUC, and with decreased MSCI. In male mice with mutations in genes essential for establishment of synapsis (Spo11, Dmc1, Atm5, Dmc1), regions of asynapsis are transcriptionally silenced through sequestration of silencing factors such as γH2AX that are no longer available to establish MSCI [24]. The deficiency to form sex bodies in Smc1β−/− spermatocytes, and thus the failure to see γH2AX accumulation on the sex chromosomes, suggests failing MSCI. In addition, the presence of autosomal asynapsis in Smc1β−/− spermatocytes renders an MSUC response likely. However, a direct contribution of cohesin to regulation of gene expression in meiotic cells appears also possible considering this role of cohesin in somatic cells. To test this, we performed microarray analysis of gene expression of wt and Smc1β−/− spermatocytes. We chose to compare mRNA expression in testes from 16 day-old mice when the first, synchronized wave of meiosis proceeds. At this stage the cells in our wt and Smc1β−/− littermate mice have just reached pachynema. There is no apoptosis yet in Smc1β−/− mice, where apoptosis commences at day 17 at the earliest [36]. Minor differences were observed in the cellularity of wt and Smc1β−/− testes at that age (Suppl. Fig. S2).

Gene expression patterns of testes of four pairs of wt and Smc1β−/− littermates were analyzed. We performed eight independent gene expression measurements using the One-Color 4 × 44K Whole Mouse Genome Array system (Agilent). The data yielded a highly significant pattern of changes in gene expression (Fig. 2A). Of all 347 genes whose expression was altered in Smc1β−/− spermatocytes by at least two-fold with a stringent p-value threshold of 5×10−5, 85% were down-regulated and 15% up-regulated. Of the aberrantly regulated autosomal genes, 91.2% were down-regulated, between 5 and 25 genes per chromosome (Fig. 2). In striking contrast, a large number of up-regulated genes are found on the X chromosome. Nineteen X chromosomal genes were up-regulated, while only 2 X-linked genes were down-regulated (Fig. 2B). On the Y chromosome, 2 genes were up-regulated, none were down-regulated. Thus, early pachytene spermatocyte gene expression on autosomes and sex chromosomes reacts inversely to loss of SMC1β.

For the validation of microarray data, reverse transcription quantitative PCR (RT-qPCR) was performed for top hits of several categories of mis-regulated genes. The genes were grouped into autosomal down-regulated genes, autosomal up-regulated genes and mis-regulated genes on the X- and Y-chromosome. For 21 of the genes of these groups, expression ratios from microarrays and RT-qPCRs from two or three mouse pairs are depicted in Figure 2D. Respective ratios of data from the microarray and RT-qPCR analyses correlated with a high correlation coefficient (R² = 0.85) (Fig. 2E).

Interestingly, the Y-linked Zfy1/2 genes, which were found up-regulated in Smc1β−/− spermatocytes, are known to strongly contribute to the pachynema, stage IV, quality surveillance mechanisms [26]. Expression of the transcription factor ZFY1/2 at that stage causes cell death. Up-regulation of Zfy1/2 gene expression was also confirmed by RT-qPCR (Fig. 2D) and suggests that this contributes to apoptosis of stage IV Smc1β−/− spermatocytes.

To test if certain genomic regions in Smc1β−/− pachytenye cells show specific clustering of up- or down-regulated genes we performed two types of analysis, using (1) a two-sided Wilcoxon rank sum test using a sliding window of 10 genes, and (2) a three-state Hidden Markov model (Fig. 2C; Suppl. Fig. S3, S4). Both methods identified focal clusters of predominantly down-regulated genes on the autosomes, whereas clusters consisting of primarily up-regulated genes were identified on the X-chromosome (the Y-chromosome was excluded from this analysis). However, after correcting for multiple hypothesis testing, the remaining number of significant clusters on the autosomes was relatively small.
Figure 2. Gene expression analysis of testes of juvenile Smc1b−/− mice. A–E. Analysis of total testis gene expression patterns of Smc1b−/− mice aged 16 dpp compared to wt littermates. A. In testes of Smc1b−/− mice, 86% of genes changed at least two-fold in expression are down-regulated (grey) and 14% are up-regulated (black, left panel). Mis-expressed genes are predominantly down-regulated on the autosomes and mostly up-regulated on the X and Y chromosomes (right panel). B. Genes on X and Y chromosomes with highest fold changes in microarray-based profiles. C. Idiogram of chromosomal location of mis-expressed genes. Down-regulated (grey) and up-regulated (black) genes are represented by circles and gene density is depicted by grey bars. D. Microarray-based expression of 21 selected genes was validated by quantitative reverse transcriptase PCR (RT-PCR). Genes were grouped according to chromosomal location and the mode of regulation. E. Microarray-based fold changes of the 21 selected genes correlate with RT-PCR data. The coefficient of determination $R^2 = 0.888$ confirms high correlation. F. In testes of Smc1b−/− mice aged 12 dpp, 66% of genes changed in expression by at least 1.5-fold are down-regulated (grey) and 34% are up-regulated (black, left panel). Mis-expressed genes are broadly distributed amongst chromosomes and moderately enriched on chromosome 15 (right panel). G. Numbers of mis-expressed genes in testes of Smc1b−/− mice aged 12 dpp (left) and 16 dpp (right) with different statistical significance and fold regulation thresholds.

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because of the global down-regulation on the autosomes. Thus, due to the large number of down-regulated genes on the autosomes such clusters could theoretically emerge by chance.

The patterns of changes in gene expression between leptonema and mid-pachynema are complex and not yet understood at high resolution for individual, defined stages or cell populations. Generally, it is thought that with completion of synopsis in pachynema, autosomal gene expression increases as the silencer proteins such as H2AX retract from synapsed autosomes and assembles on the sex chromosomes. In early/mid pachytenic Smc1β−/− spermatocytes a significant number of autosomal genes fails to be expressed. To determine whether the down-regulation of autosomal gene expression seen in Smc1β−/− spermatocytes correlates with synopsis, i.e. likely originates from an MSUc response rather than a direct effect of SMC1β on transcription, we prepared mRNA from testes of 12 day-old mice (Fig. 2F). At that stage, spermatocytes of our mouse strains are in early pachynema and thus there is still little synopsis. If SMC1β has a direct, synopsis-independent effect on transcription it should be measurable at this stage, similar to the pachytene effect, albeit limited to those relatively few genes that are normally expressed in early zygonyema [22]. Assaying the samples by the same microarray technology revealed, however, not a single down- or up-regulated gene at a p-value below 0.0001 and a minimum 2-fold change (Fig. 2F, 2G). At a less stringent p-value of 0.001 eight genes were identified of which half were down-regulated, half up-regulated. In contrast, at this p-value 1080 genes, almost all down-regulated, were identified for the pachytene samples. This strongly suggests a direct correlation of autosomal down-regulation with the failure to completely synapse. Chromosomal distribution analysis of 160 mis-regulated genes (day 12) identified at low stringency (p<0.05; f.c>1.5) shows them on all chromosomes (Fig. 2F). Many of the mis-regulated genes localize to chromosome 15 with an enrichment in the proximity to the Smc1β locus, indicative of an artifact generated through alterations of the locus environment by gene targeting. We therefore disregarded these genes. No significant changes were detected in mRNA microarrays (Suppl. Fig. S5).

Thus the vast majority of down-regulation in Smc1β−/− pachytene cells is very likely not caused by a direct effect of SMC1β on transcription, but rather by an MSUC response.

Phosphorylated SMC1 and SMC3 distinctly localize to the sex body

Recently it was shown that synopsis affects localization of phosphorylated SMC3 [61]. In mitotic cells, SMC1 and SMC3 become phosphorylated in an ATM- or ATR-dependent manner within the DNA damage response [62,63]. SMC1 and SMC3 localize to DSBs in somatic cells (reviewed in [40,64]), and in this respect behave similar to H2AX. ATR localizes to asynapsed regions in meiocytes [9], and recently it was reported that pSer1083-SMC3 also localizes preferentially to asynapsed chromosomes, including the sex chromosomes in pachynema and diplonema [61]. We extended this observation, and asked whether pSer966-SMC1 also localizes to asynapsed regions including the sex body, and how phosphorylated SMC1β and SMC3 behave in mutant backgrounds.

In leptonema and early zygonyema, pSer1083-SMC3 localizes along chromosomes in a punctuated pattern, in pachynema and diplonema pSer1083-SMC3 marks the sex body (Fig. 3A). Unlike HORMAD1, which localizes along the sex chromosome axes (Fig. 3B), pSer1083-SMC3 appears as a cloud similar to, but not as expanded as the H2AX cloud (Fig. 3A, B and Fig. 1B). In wt pSer1083-SMC3 is found exclusively at the sex body from late zygonyema to diplonema (Fig. 3A). Controls showed that the phospho-SMC1 peptide did not block anti pSer1083-SMC3 reactions, but the phosphorylated SMC3 epitope peptide eliminated the signal (Suppl. Fig. S6A). No X/Y axes staining appeared even at low concentrations of the pSMC3 peptide, i.e. the pSer1083-SMC3 always appeared as a cloud. In metaphase I, SMC3 was observed at centromeres, a prominent structure stained by SYCP3 at this stage (Suppl. Fig. S6B).

In Smc1β−/− spermatocytes (Fig. 3C, D) pSer1083-SMC3 localizes to unsynapsed chromosomes in late zygonyema. In early/mid pachytene the pSer1083-SMC3 is not detected at sex bodies, which as described above do not form. Occasionally, deposits of pSer1083-SMC3 are seen that associate with part of a chromosome. On Smc1β−/− spermatocytes the pSer1083-SMC3 is observed at the HORMAD1-stained asynapsed chromosomes, regardless of their number (Fig. 3D). Since SMC1β is present [36] and SMC1β-deficient cells were analyzed, this indicates that an SMC1β-pSer1083SMC3 complex marks asynapsed regions, at least in the absence of SMC1β.

The anti pSer966-SMC1 antibody recognizes only phosphorylated SMC1β since the Ser966 is not conserved between SMC1α and SMC1β. In wt leptonema spermatocytes, no pSer966-SMC1 was detected. In late zygonyema, pSer966-SMC1 is preferentially seen at the asynapsed axes. In pachynema and early diplonema pSer966-SMC1 localizes only to the sex body, and in late diplonema pSer966-SMC1 disappears (Fig. 4A). pSer966-SMC1 localizes almost exclusively to the chromosome axes and does not form clouds like pSer1083-SMC3. This suggests that non-phosphorylated SMC1β/pSMC3 complexes localize to the sex chromosome loops and show as clouds, and that pSMC1α/SMC3 complexes associate with the X/Y axes. It is rather unlikely that an alternative SMC1β/pSMC3 complex localizes to the sex body cloud, since SMC1β was only seen at the sex chromosome axis [35]. In Smc1β−/− spermatocytes, the pSer966-SMC1 localizes along the asynapsed chromosomes (Fig. 4B).

**SPO11-generated DSBs are not required for cohesin loading but for cohesin phosphorylation**

Cohesin can be loaded onto mitotic cell chromosomes upon introduction of DNA DSBs [65]. The use of Spo11−/− mice lacking programmed DSBs [12,13] allowed us to analyze the recruitment of cohesins, including phosphorylated SMC1 and SMC3, to meiotic chromosomes. Earlier, SMC1α and SMC3 were found to localize to chromosomes of Spo11−/− spermatocytes [66]. We observed all of the cohesin proteins tested here to localize to chromosomes of Spo11−/− cells, including SMC1α, SMC1β, SMC3, STAG3, RAD21 and RAD21L (Suppl. Fig. S7A–G). The patterns of association differed between individual cohesins. Coating of SMC1α with HORMAD1 revealed that in Spo11−/− spermatocytes SMC1α localizes along AEs preferentially where no HORMAD1 associates, i.e. more SMC1α localizes to regions of non-homologous associations. There is little SMC1α on non-associated chromosomes (Suppl. Fig. S7A; quantification in Suppl. Fig. S8). As published before [50], in wt cells SMC1α localizes along every chromosome, including asynapsed chromosomes in zygonyema, and stains in comparable intensities whether synapsed or not. In Spo11−/− non-associated chromosomes (Suppl. Fig. S7A; quantification in Suppl. Fig. S8). In wt, SMC3 also localizes to asynapsed and synapsed regions with equal intensity [50]. SMC1β localizes along chromosomes in wt and Spo11−/− spermatocytes. Together, this suggests that SMC1β and not
Figure 3. Phosphorylated SMC3 distinctly localizes to asynaptic regions in wt, Smc1β/−, and Smc1β/−Spo11−/− spermatocytes. A. Wt spermatocyte chromosome spreads stained with anti SYCP3 for AEs and anti phosphorylated SMC3(S1083). B. Staining of unsynapsed chromosome axes and of sex chromosomes axes in wt spermatocyte spreads with anti HORMAD1, combined with staining by anti phosphorylated SMC3(S1083). C. Staining of Smc1β/− spermatocyte chromosome spreads using anti SYCP3 for AEs and anti phosphorylated SMC3(S1083). D. Staining of unsynapsed sex chromosome axes or unsynapsed autosomes in Smc1β/− spermatocyte chromosome spreads with anti HORMAD1, combined with staining by anti phosphorylated SMC3. E. Staining of Spo11−/− spermatocyte chromosome spreads using anti phosphorylated SMC3(S1083) combined with either anti SYCP3 and DAPI (left, top), or with anti γH2AX (n = 40). F. Staining of Smc1β/− Spo11−/− spermatocyte chromosome spreads with anti SYCP3 and anti phosphorylated SMC3(S1083) (n = 190).

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SMC1\(\alpha\) complexes localize preferentially to non-associated regions (Suppl. Fig. S7G). The data also suggest that localization of SMC1\(\alpha\) to asynapsed regions depends on SPO11. STAG3 decorates all axes in a punctuate pattern in wt and Spo11\(^{-}\) cells (Suppl. Fig. S7C). RAD21L localizes to the asynapsed axes in Smc1\(^{+/}\) spermatocytes (Suppl. Fig. S7D). RAD21L also localizes in a somewhat punctuated fashion all along the Spo11\(^{-}\) axes, and in wt pachytene cells preferentially accumulates at asynapsed regions, particularly the sex chromosomes (Suppl. Fig. S7E). In wt, RAD21 co-localizes in many dots with SYCP3 along autosomes, but there is little overlap with SYCP3 on the sex chromosomes (Suppl. Fig. S7F), where the RAD21 signals accumulate next to the axes. In Spo11\(^{-}\) chromosome spreads, very little of RAD21, which appears as many dots throughout the chromatin, co-localizes with SYCP3 on the chromosome axes.

In spermatocytes lacking SPO11 we detected no pSer1083-SMC3 on autosomes, whether unsynapsed or non-homologously associated, consistent with the report by Fukuda et al. (2012) (Fig. 3E). However, we found pSer1083-SMC3 to form some clusters within the nuclei reminiscent of pseudo sex bodies, and to localize in dispersed spots throughout the nuclei in Spo11\(^{-}\) spermatocytes (Fig. 3E). In many cells there was overlap of pSer1083-SMC3 with staining for \(\gamma H2AX\) at the pseudo sex bodies. In Smc1\(^{+/}\) Spo11\(^{-}\) spermatocytes, the localization of pSer1083-SMC3 in many cells is similar to that seen in Spo11\(^{-}\) spermatocytes, but in about 40% of the cells, no staining for pSer1083-SMC3 was observed (Fig. 3F). Together this indicates that SMC1\(\alpha\) forms complexes with pSer1083-SMC3 in absence of SPO11 and SMCM1\(\beta\). We also noted moderately longer chromosome axes in Smc1\(^{+/}\) Spo11\(^{-}\) spermatocyte spreads as compared to Smc1\(^{+/}\) spreads (Fig. 4B, 4D; quantification in Suppl. Fig. S9), although this varied and in some cells this difference was small. In Smc1\(^{+/}\) spermatocytes the pSer966-SMC1 signal is seen on asynapsed autosomes (Fig. 4B). This supports the above hypothesis of pSMC1\(\alpha\)/SMC3 complexes to localize to asynaptic regions. No pSer966-SMC1 was detected in Spo11\(^{-}\) or Smc1\(^{+/}\) Spo11\(^{-}\) spermatocytes (Fig. 4C, D). These data indicate that SPO11, and presumably SPO11-generated DSBs, are required for SMC1\(\alpha\) Ser966 phosphorylation, consistent with its kinetics of appearance in early zygona.

**Delayed disappearance of repair foci in absence of SMC1\(\beta\)**

Asynapsis may have consequences for the processing of meiotic double-strand breaks (DSBs) as suggested by the correlation of synapsis and disappearance of repair foci, which is delayed on the largely unpaired X/Y chromosomes. To address formation and processing of DSBs in Smc1\(^{+/}\) spermatocytes, we first determined whether programmed DSBs are properly generated in absence of SMC1\(\beta\). Thus, SPO11-dependent DSBs were measured by visualizing the SPO11-oligonucleotide intermediate product of the reaction [67].

Fig. 5A demonstrates that SPO11 protein is produced in adult wt and Smc1\(^{+/}\) cells, although in the latter the SPO11\(\alpha\) splice variant is absent as it is expressed later and may not contribute much to leptotene spermatocyte DSB formation [68]. The alternative splice isoform SPO11\(\beta\) is present although in much smaller quantities as in wt, probably because late pachytene cells, which contribute significantly to total SPO11 in spermatocytes, are absent. The early SPO11 enzyme is sufficient to produce substantial levels of SPO11-oligonucleotide complexes, which is also seen in the control Dmc1\(^{-}\) samples that contain even less SPO11 enzyme, and whose spermatocytes arrest in late zygona. Smc1\(^{+/}\) spermatocytes develop further into pachynema and probably therefore synthesize more SPO11.

We assessed steady-state levels of SPO11-oligonucleotide complexes in whole-testis extracts from wt and mutant mice. We found a very mild reduction in adult Smc1\(^{+/}\) testes (0.84±0.20-fold relative to wt, mean and s.d., \(n=2\) littermate pairs). In contrast, adult Dmc1\(^{-}\) testes, which are similarly reduced in size as Smc1\(^{+/}\) testes, displayed a more substantial reduction in SPO11-oligonucleotide complex levels (0.42±0.07-fold, \(n=2\)), as previously reported [53]. To account for different sizes and cellularity in wt and mutant testes, we also examined SPO11-oligonucleotide levels in juvenile mice at day 14.5 post partum, when the majority of spermatocytes have not yet progressed far enough into meiotic prophase to be affected by the mid-pachynema checkpoint. At this age, SPO11\(\beta\) is only weakly expressed, but SPO11-oligonucleotide complexes are readily detectable. Complex levels were moderately reduced in Smc1\(^{+/}\) testes (0.68±0.14-fold, \(n=3\)) and in Dmc1\(^{-}\) testes (0.77±0.03-fold, \(n=2\)). This does not argue for an impairment in SPO11 DSB formation, since minor variations such as slightly delayed initial spermatogenesis in the Smc1\(^{+/}\) mutant may cause this phenotype.

Efficient production of DSBs in the adult was confirmed by quantifying the number of initial repair foci. Repair proteins assemble at DSBs and form foci whose appearance is indicative of DSBs, and whose processing, i.e. disappearance, is indicative of proper repair. If analysis of RAD51 and DMC1 foci revealed that the same number of foci are generated in wt and Smc1\(^{+/}\) spermatocytes at the early zygotene stage (Fig. 5B, Suppl. Fig. S10). In late zygotene and early pachytene, these foci are processed and disappear in wt cells, but fail to do so efficiently in Smc1\(^{+/}\) spermatocytes. Detailed analysis of the persistent foci by co-staining with HORMAD1 revealed that they localize predominantly to asynapsed regions and to sex chromosomes (Fig. 5B, C). As expected, no foci were detected in Spo11\(^{-}\) [not shown] and in Smc1\(^{+/}\) Spo11\(^{-}\) spermatocytes (Fig. 5B).

**Non-homologous synopsis requires SMC1\(\beta\)**

Staining of Smc1\(^{+/}\) Spo11\(^{-}\) spermatocyte spreads (Fig. 4D) unexpectedly showed 40 clearly separate, SYCP3-stained chromosome cores. Therefore, we further analyzed whether non-homologous associations in Spo11\(^{-}\) spermatocytes depend on SMC1\(\beta\). Staining of Smc1\(^{+/}\) Spo11\(^{-}\) spermatocyte spreads for HORMAD1 and SYCP3 showed 40 separate, short chromosome cores, i.e. univalents without any non-homologous association (Fig. 6A). Axes shortening is a known, prominent phenotype of Smc1\(^{+/}\) spermatocytes [36]. Since Smc1\(^{+/}\) Spo11\(^{-}\) spermatocytes die slightly earlier, in late zygona, than Smc1\(^{+/}\) spermatocytes, the axes are not yet as fully compacted and are slightly longer than in the latest stages of Smc1\(^{+/}\) spermatocytes (Suppl. Fig. S9; Fig. 4).

Quantification showed that 58% (22 of 38) of Smc1\(^{+/}\) Spo11\(^{-}\) spermatocytes carry 40 chromosome cores without non-homologous
Spermatocyte chromosome spreads with anti HORMAD1, anti DMC1 and anti SYCP3, and DAPI.

The number of DSBs depends on SPO11 dosage [69]. Would the introduction of fewer DSBs still allow homologous synopsis in the suppression-weakened Snc1β−/− spermatocytes? Snc1β−/− Spo11−/− spermatocytes appeared phenotypically not different from Snc1β−/− cells, i.e. showed about the same level of asynapsis when stained for SYCP3 and SYCP1 (Fig. 6D) or pSer966-SMC1α (Fig. 6E). This indicates that wild-type levels of DSBs are not required for the level of synopsis seen in the absence of SMC1β. As expected, the number of DMC1 foci was slightly decreased in Snc1β−/− Spo11−/− spermatocytes compared to wt and persisted longer than in wt as described for the Snc1β−/− cells [not shown].

Further, we observed in some Snc1β−/− Spo11−/− spermatocytes but not in wt or single “knockout” cells what appears to be end-fusions of chromosomes, i.e. SYCP3-stained axes with a centromere signal at both ends (Fig. 6B shows 2 examples). Mostly two, but also up to 6, of the 40 or more chromosomes of about half of the cells (n = 30) were seemingly connected or at least associated at one of their ends. It appeared as if mostly the centromere-distal ends of the two chromosomes were fused.

Recently, we reported on gene dosage effects in Snc1β−/+ and Rec8−/+ mice [70]. A gene dosage effect of SMC1β was also observed for Snc1β−/+ Spo11−/− spermatocytes. In a small subset of these cells (2%), no synopsis – whether homologous or non-homologous – was observed. In the remaining 98% of the cells, partial non-homologous synopsis typical for Spo11−/− spermatocytes was seen (n = 30; not shown).

SMC1β is necessary for sister chromatid cohesion in prophase I

Previously, we reported a requirement for SMC1β in centromeric SCC in an okadac acid induced metaphase I-like stage in spermatocytes and in later stages of oocyte meiosis [36,37]. However, it remained unclear whether SMC1β is required for SCC in prophase I and whether it acts in arm and/or centromeric cohesion in early meiosis. In almost half of the Snc1β−/− Spo11−/− spermatocytes, such as the one shown in Figure 6B, the increased number of axes indicates loss of cohesion in some pairs of sister chromatids. However, 58% of the cells carry 40 axes, which show no signs of split sister chromatids, i.e. feature cohesion all along the chromosome arms. Thus, the remaining SMC1α complexes are sufficient for this fraction of arm cohesion at this stage. Since there is no pSer966-SMC1α in Spo11−/− cells, the non-phosphorylated form of SMC1α provides this sister chromatid cohesion. Non-phosphorylated SMC1α likely acts together with non-phosphorylated SMC3, since the pS3MC3 is dispersed, not associated with the chromosome axes. Thus, both SMC1α and SMC1β can contribute to arm cohesion in early prophase I.

Centromeric cohesion was analyzed in Snc1β−/− Spo11−/− and in Snc1β−/+ spermatocytes (Fig. 7). The axes of Snc1β−/− Spo11−/− cells were investigated for split centromere signals indicative of loss of centromeric cohesion. Clearly split centromeres were observed in 35% of the axes (n = 60 cells) (Fig. 7A). Chromosome surface spreading does not always resolve separate structures, and thus centromeres that were of large, unusual shape were distinctly

![Figure 5. Normal formation but delayed processing of DNA DSBs in Snc1β−/+ spermatocytes.](image-url)

**Figure 5. Normal formation but delayed processing of DNA DSBs in Snc1β−/+ spermatocytes.** A: Top: Immuno blot of nuclear protein extracts from testis of the indicated mouse strains, probed with anti SPO11 antibody. The two SPO11 variants are indicated. Bottom: SPO11-DNA complexes assayed as described in Materials and Methods, analyzed in tests from adult or juvenile (day 14.5 pp) mice of the indicated mouse strains. B: Staining of wt (n = 120), Snc1β−/− (n = 160) or Snc1β−/− Spo11−/− (n = 80) spermatocyte chromosome spreads with anti HORMAD1, anti DMC1 and anti SYCP3, and DAPI. C: Graphical representation of DSB processing in wt and Snc1β−/+ spermatocytes at the indicated stages of meiosis. Numbers of DMC1 foci in synapsed and asynapsed regions are provided. doi:10.1371/journal.pgen.1003985.g005
quantified and found in 45% of the axes, excluding those with clearly separate centromeres. Perfectly overlapping centromeres, suggesting presumably conserved centromeric SCC, were present in the remaining 20%. This suggests a very significant contribution of SMC1β to centromeric cohesion. To confirm this, Smc1β−/− spermatocytes were similarly analyzed, although here the analysis is more complex, since the two pairs of sister chromatids with their four sister chromatids can be partially asynapsed, and may show two centromere signals because either of asynapsis or of loss of cohesion, which cannot be distinguished. We can conclude with certainty that centromeric SCC was lost only if there are three or four centromere signals. Some of the SCs showing two centromere signals may also have lost centromeric SCC. We observed 3 or 4 centromeres, i.e. loss of centromeric cohesion, in at least 3% of SCs. Another 5% of SCs showed 2 centromere signals and an unknown fraction of those may be derived from loss of SCC instead of asynapsis. Some SCs showed clearly asynapsis at the centromeric end with (30%) or without (70%) loss of centromeric cohesion. Centromeric asynapsis was as frequent as asynapsis within the chromosome arms or at the centromere-distal end. No split centromeres were observed in wt or Spo11−/− spermatocyte chromosome spreads.

**Discussion**

This report addresses two major roles of cohesin in meiosis: sister chromatid cohesion and synopsis. All data and phenotypes reported here relate to these functions, either directly or indirectly. Previous reports implicated the meiosis-specific SMC1β cohesin in synopsis, post-prophase I SCC, chiasma maintenance and telomere protection [36,37,46]. However, important questions, which concern fundamental processes of male meiosis, remained to be resolved: (1) is SMC1β required for synapsis of sex chromosomes and for non-homologous synaptic associations; (2) does SMC1β contribute to SCC during prophase I of meiosis; (2) is SMC1β necessary for generation and processing of meiotic, programmed DSBs? Other, more complex questions are step-wise to be resolved as new mouse models become available, such as for the direct roles of SMC1α and SMC1β complexes in meiosis, though this study provides several insights into the functions of these two complexes. Conclusions about the role of SMC1α complexes remain indirect, since these conclusions are based on functions of SMC1α that is present in SMC1β deficient spermatocytes. Whether SMC1α fulfills the same roles in wt cells cannot be determined with certainty.

Sister chromatid cohesion as the prominent function of cohesin may be provided early in meiosis either by SMC1α- or SMC1β-based complexes. No evidence had previously been presented for a contribution of SMC1β to prophase I SCC, which did not appear to be grossly dysfunctional in Smc1β−/− prophase I cells. In addition, expression of the Smc1β gene starts only right after cells have entered meiosis, i.e. in early leptotene [35,71]. Thus the previous assumption was that SMC1α complexes provide most if not all SCC before metaphase I. A contribution of SMC1β to centromeric cohesion was suggested by published data, based on experimentally driving Smc1β−/− spermatocytes into a metaphase I-like state by okadaic acid treatment [36]. In this rather artificial set-up, 80 separate centromeres were observed in many Smc1β−/− cells indicative of complete loss of centromeric SCC of the 4×20 mouse chromatids [36]. However, some of the cells only partially lost SCC (average of 40 centromere signals), and some showed app. 20 signals for intact centromeric SCC. We speculated that those cells suffering complete loss of SCC were of pachytene origin, while those that showed only partial loss of SCC may have originated from earlier stages. Together, it remained unclear whether SMC1β contributes to SCC earlier than metaphase I, i.e. in prophase I.

The analyses of wt versus Smc1β−/− or Smc1β−/− Spo11−/− spermatocytes demonstrate an important contribution of SMC1β to zygotene/pachytene SCC. At least one-third of centromeric SCC in Smc1β−/− Spo11−/− spermatocytes is impaired. At least 9% of SCs in Smc1β−/− spermatocytes lose centromeric cohesion, but this number, which includes chromosomes with three centromere signals only, is probably a significant underestimation. The many centromeres that show two signals cannot be assigned to either asynaptic homologs with intact centromeric cohesion or to synapsed homologs with defective centromeric cohesion. SCC of chromosome arms appeared intact in zygotene and pachytene Smc1β−/− spermatocytes, but much higher microscopic resolution may be required to determine this. The Smc1β−/− Spo11−/− strain with its complete absence of homolog associations revealed in almost half of the cells more than 40 axes, mostly between 42 and 48 axes, and thus showed individual sister chromatids. Thus, about 5 to 10% of the homologs in about half of the cells lost SCC. Thus, SMC1β also contributes to arm cohesion in early prophase I. However, arm cohesion is primarily supported by SMC1α cohesin under these conditions. We did not observe partial loss of arm cohesion, which would show as bubbles, i.e. separate strands, within homologs. This “all or none” response may indicate cooperative behavior of cohesin. It shall be explored in the future why SMC1α does not provide full centromeric cohesion, and why SMC1β does relatively little for arm cohesion, i.e. how and why this functional specialization occurs. SMC1β remains associated with centromeres until metaphase II, consistent with the hypothesis that SMC1β provides the essential centromeric cohesion beyond prophase I. Formally, the synergistic effect of deficiencies in SMC1β and SPO11 renders it possible that SPO11 itself also contributes to SCC. Given the prior knowledge on SPO11 we deem this unlikely, and rather think that the much more likely interpretation is that the deficiency in SPO11 and thus absence of homologous synopsis allowed us to reveal the SPO11-independent role of SMC1β in SCC.

The analysis of Smc1β−/−, Spo11−/−, and Smc1β−/− Spo11−/− spermatocytes revealed several other important aspects of SMC1β function. It confirmed that SPO11 and thus SPO11-generated DSBs are not required for loading of cohesin proteins. The localization of RAD21 in Spo11−/− spermatocyte chromosome spreads differed from wt since much less RAD21 co-localized with the SYCP3-stained axes. More importantly, the non-homologous synaptic associations observed in Spo11−/− spermatocytes [12,13] entirely depend on SMC1β. SMC1α alone cannot sufficiently

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**Figure 6. Non-homologous association of axial elements depends on SMC1β.** A Smc1β−/− Spo11−/− spermatocyte chromosome spreads were stained with anti SYCP3 for AEs and anti HORMAD1 for unsynapsed chromosomes or chromosomal regions (n=70). B Smc1β−/− Spo11−/− spermatocyte chromosome spreads stained with anti SYCP3 for AEs and anti ACA for centromere staining. Two examples where chromosomes apparently show fusion at the centromere-distal ends are shown. C Localization of different cohesin proteins in Smc1β−/− Spo11−/− spermatocyte chromosome spreads, which were stained with anti SYCP3 for AEs, Smc3 (n=40), STA3G (n=30), and RAD21L (n=30) for different cohesin proteins. D Staining of Smc1β−/− Spo11−/− spermatocytes for SYCP1 and SYCP3. E Staining of Smc1β−/− Spo11−/− spermatocytes for pSer966-SMC1α and SYCP3.
A

**Smc1β**<sup>−/−</sup> **Spo11**<sup>−/−</sup>

**Smc1β**<sup>−/−</sup> **Spo11**<sup>+/−</sup>

2 1 2 2

**SYCP3**  **ACA**

| % of chromosomes (n=60) | 2 split centromere | potential split centromere | 1 centromere |
|-------------------------|--------------------|----------------------------|--------------|
|                         | 35                 | 45                         | 20           |

B

**Smc1β**<sup>−/−</sup>

**Smc1β**<sup>−/−</sup>

**Smc1β**<sup>−/−</sup>

**Smc1β**<sup>−/−</sup>

2 2 3 3

**ACA**  **SYCP1**  **SYCP3**
Figure 7. Loss of centromeric sister chromatid cohesion in Smc1β−/− and Smc1β−/−Spo11−/− early pachytene spermatocytes. A Smc1β−/−Spo11−/− spermatocyte chromosome spreads were stained with anti SYCP3 for AEs and anti ACA for centromeres (n = 60). The percentages of univalents showing 2 centromeres (clearly split signals), potentially split centromeres, and one centromere (non split) are provided. B Smc1β−/− spermatocyte chromosome spreads were stained with anti SYCP3 for AEs, SYCP1 for SCs, and anti ACA for centromeres (n = 50).

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support these associations. Thus, a second instance of functional specialization between the two SM1 variants has been identified. Non-homologous associations might reflect an early, transient stage in the search of the homologs to find each other even before DSBs are generated. We suggest that cohesin is involved in this early stage of partner search and may provide a platform or code that allows transient, weak “trial-and-error” interactions to enhance the chance of finding the homolog.

In many Smc1β−/−SPO11−/− spermatocytes, but not in either single mutant or wt, several seemingly fused chromosomes were observed, where often the centromere-distal ends were fused. Earlier, we described telomere defects in Smc1β−/− meioocytes such as loss of telomeric sequences, SCs without telomeres, telomere stretches and extended bridges between telomeres consisting of stretched telomere DNA and telomeric proteins [46]. About 20 to 30% of the telomeres showed such aberrations, which we interpreted as loss of telomere protection in absence of Smc1β. The absence of protection of chromosome ends and the failure of the chromosomes to homologously synapse may contribute to end-fusions in the Smc1β−/− Spo11−/− spermatocytes. Homologously synapsed chromosomes are highly compacted and may have acquired a chromosome configuration prohibitive of end-fusions. This configuration does not exist in Smc1β−/− Spo11−/− spermatocytes. Telomere aberrations also cause failures in synapsis and may contribute to the asynapsis seen in SM1β deficient cells.

Average chromosome length was moderately increased in Smc1β−/− Spo11−/− spermatocytes compared to wt cells, but variation was high. This may either reflect a slightly different developmental stage of the double-deficient cells or reduced compaction in absence of SPO11-dependent synapsis.

Why does SM1 interact efficiently to asynapsed, non-associated regions in Spo11−/− spermatocytes but binds to the largely unpaired sex chromosomes in wild-type cells? One may speculate that there is a specific chromatin mark or other quality in wt cells that supports SM1 binding to the sex chromosomes. This study further demonstrates that sex body formation depends on SM1β and that in the absence of SM1β, SM1 alone cannot support PAR synapsis. The involvement of SM1β may be direct in supporting synapsis for example through interaction of cohesin rings between the two pairs of sister chromatids. It can also be indirect if a certain axis length of the PAR is required for sufficient synapsis, and that length may not be available with the shortened Smc1β−/− axes. Our data also provides a likely explanation for the apoptosis of Smc1β−/− spermatocytes observed in mid-pachynema. While asynapsed autosomes are silenced, genes on the sex chromosome are transcriptionally activated. Thus, SM1 fails. This correlation between an MSUC reaction of autosomes and a failure of SM1 has been described and is attributed to the relocalisation of silencing factors such as γH2AX from sex chromosomes to asynaptic autosomes [9]. Made possible through the MS1 failure, the Y chromosome “spermatocyte killer gene” Zfy1/2 is expressed and probably substantially contributes to eliminating the mutant spermatocytes within the pachytenocyte quality surveillance mechanism [26] reviewed in [72].

Our analysis of phosphorylated forms of SM1 and SM3 revealed a distinct pattern of localization, particularly at the sex chromosomes. While it is difficult to entirely exclude antibody staining artifacts, the measures taken such as peptide blocking experiments, single antibody staining, secondary antibody controls, and the use of well-established anti phospho SM1 antibodies, as well as the specific staining patterns like staining of the axis only of unsynapsed chromosomes (pSM1) or staining only of the sex body chromatin (pSM3) render artifacts unlikely.

Phosphorylation of SM1 or SM3 may serve as a mechanism to attribute functional properties to these cohesins at different localizations. The pSer1083-SM1β may also serve as a marker for sex bodies, and the pSer966-SM1α as a marker for the sex chromosome axes or asynapsed axes in mutants like Smc1β−/−. Since in about 40% of Smc1β−/− Spo11−/− spermatocytes no pSer1083-SM1β was detected, but all Spo11−/− spermatocytes showed pSer1083-SM1β, the phospho-form may either be unstable or these 40% of cells may have been at slightly distinct stages of spermatogenesis as their counterparts in the same mouse. The absence of pSer966-SM1α from Spo11−/− spermatocytes indicates distinct requirements for phosphorylation of SM1α and SM1β. The pSer966-SM1α is observed at asynapsed chromosomes in Smc1β−/− spermatocytes but not in Spo11−/− spermatocytes that carry extensively asynapsed chromosomes. Thus, it appears as if SPO11-generated DSBs are required for SM1α phosphorylation. SM1α phosphorylation in somatic cells is catalyzed by ATM, whose activation in meiocytes depends on DSBs, which could explain the failure to see pSer966-SM1α in Spo11−/− spermatocytes.

Spo11 heterozygosity rescues the prophase I arrest of Abt−/− spermatocytes, and revealed that sex body formation and H2AX phosphorylation are supported by ATR [73,74]. In one explanation, the reduced number of DSBs and their slower generation allows ATR to complement the ATM deficiency, particularly at zygonema and pachynema, when ATR is strongly expressed. Recent work showed, however, that ATM restricts the number of SPO11-generated DSBs and that in the absence of ATM, expression levels of SPO11 become critical [53]. Too many DSBs in an ATM-deficient, SPO11-proficient situation are deleterious and the reduced levels seen in Abt−/− Spo11−/− spermatocytes are more permissive for meiotic progression. Despite this ATM-dependent DSB-triggered feedback mechanisms, the number of DSBs depends on SPO11 dosage, since Spo11−/− spermatocytes show reduced levels of DSBs [69]. Our analysis shows that wt levels of SPO11-generated DSBs are not required for SM1β-independent non-homologous synapsis in Smc1β−/− Spo11−/− cells.

Several studies investigated gene expression patterns during male gametogenesis [75,76,77,78]. Most of these studies compared two or more cell populations, e.g. spermatogonia with spermatocytes, spermatocytes with spermatids, or germ cells with Sertoli cells, but did not venture into analyzing meiotic sub-populations, which, for example, represent leptotene, zygotene or pachytene cells. These are very difficult to separate and to obtain in high purity and new mouse models expressing specific markers may be needed to achieve this. Therefore, juvenile male mice were used in some studies to exploit the synchrony of the first wave of meiosis during mouse puberty. However, this first wave of meiosis may not faithfully represent adult meiosis in all of its aspects. Nevertheless, altered patterns of gene expression were observed in pachytene...
spermatocytes compared to spermatagonia or spermatids. Some studies used mutant mouse strains such as Spo11−/− mice [79] to reveal gene expression patterns that depend on this gene and may be indicative of genes required for meiotic progression, but a clear assignment to this function requires additional analysis. In our study, we compared wt and Smc1b−/− spermatocytes at day 12 or day 16 post partum, at which spermatocytes have reached zygotene or early pachytene, respectively. FACS analysis of testis cells at day 16 revealed no major difference between wt and Smc1b−/− spermatocytes, but in Smc1b−/− samples some cells accumulated at a different position in the analysis window, indicating altered morphology. We cannot exclude a minor effect of slightly changed cellularity of the testsis on the gene expression patterns. However, we assume that such an effect would be small given the absence of increased apoptosis at day 16 dpf in wt and Smc1b−/− tests, the use of littermates, and the similarity of testis tubule structure at this age. At the p-value of <0.001 more than a thousand genes differed by at least 2-fold between day 16 spermatocytes from wt and Smc1b−/− spermatocytes. A large majority (92%) of the differently regulated autosomal genes, were down-regulated. In zygotene, however, only 8 genes differed at that p-value and they were down- or up-regulated. At the least stringent p-value of <0.05, only 64 genes changed at least 2-fold, and down- and up-regulation was equally distributed among them. Therefore, a strong correlation was revealed between down-regulation of many genes in Smc1b−/− pachytene spermatocytes, and their significant asynapsis. As asynapsis is known to cause the MSUC response, we suggest that most if not all down-regulation of gene expression seen in Smc1b−/− spermatocytes is caused by asynapsis-dependent MSUC. One may also describe the down-regulation as a failure to up-regulate the genes upon synapsis in early pachynema. Provided that this view is correct, one may speculate that there may be regions on autosomes that are particularly prone to asynapsis. The presence of specific genes that are very significantly down-regulated may indicate “asynapsis-sensitive” regions, and thus may suggest that cohesin-dependent synapsis happens with different efficiency and perhaps timing on distinct regions of meiotic chromosomes. Our search for chromosomal clusters of aberrantly expressed genes by two different methods revealed the presence of such clusters, although only few remained significant after correcting for multiple hypothesis testing. Hence, we observed a global reduction of expression on autosomes, but we could not identify a substantial number of specific genomic regions that are particularly affected by asynapsis in Smc1b−/− pachytene cells. Genes that are not expressed before and after synapsis, very low absolute expression levels, and possible feedback mechanisms may limit the analysis and thereby may prevent the identification of putative “asynapsis-sensitive” regions.

Considering the few genes expressed in zygotene that are misregulated in Smc1b−/− spermatocytes and are either up- or down-regulated, and considering the few pachytene genes that are up-regulated in Smc1b−/− spermatocytes, we cannot exclude, however, a small direct effect of SMC1β cohesin on gene expression, possibly through interaction with transcription factors. Alternatively, these changes in gene expression may be caused by the altered loop-axis structure seen in absence of SMC1β. For example, certain chromosome regions may be aberrantly localized in chromatin loops or packed into the axis, which may affect gene expression.

Yet another phenotype of Smc1b−/− spermatocytes correlates with asynapsis: the failure to timely process DNA DSBs. While the DSBs are formed properly as indicated by analyzing SPO11-oligonucleotide complexes and by staining for RAD51 and DMC1 foci in early zygote, these foci persist longer in Smc1b−/− spermatocytes. The repair foci are processed more slowly on asynapsed chromosomes such as sex chromosomes, and in mutants that show asynapsis. Our analysis showed that persistent repair foci localize predominantly to asynapsed chromosomes, lending support to the hypothesis that the delay in processing the repair foci is caused by asynapsis, where recombinational repair using the homolog cannot take place. The alternative pathway to repair such foci, that is the repair through recombination between sister chromatids of one homolog, is apparently not enhanced in Smc1b−/− spermatocytes, at least not to an extent that would rescue the delay in foci processing. Alternatively or additionally, more DSBs may be introduced where a chromosome or chromosomal region fails to synapse in order to further promote synapsis. The axes in the absence of SMC1β are only about half as long as those in wild-type [36]; yet the number of DSBs is only little reduced in Smc1b−/− spermatocytes. Therefore, axes length per se is unlikely to be a prime determinant of DSB numbers.

Together, this study revealed important insights into the functions of SMC1β cohesin in prophase I SCC, in autosome and sex chromosome synapsis, and it provides evidence for specific, distinct functions of SMC1β and SMC1α cohesin complexes.

Taking previous reports into account, three main functions of SMC1β have emerged that have multiple biological consequences for meiocytes: generation of the loop-axis architecture of SCs, homologous and non-homologous synapsis, as well as sister chromatid cohesion starting in early prophase I.

Supporting Information

Figure S1 Single color images and merged color images for Figure 1A–C. A. Wt and Smc1b−/− spermatocyte chromosome spreads stained with anti SYCP1 for synapsed chromosomes and anti HORMAD1 for unsynapsed chromosomes or chromosomal regions. B. Staining of the sex body in wt and Smc1b−/− spermatocyte chromosome spreads using anti SYCP1 for ASCs, anti γH2AX for the sex body or unsynapsed chromatin domains, and anti HORMAD1 for the unsynapsed sex chromosome axes or unsynapsed autosomes. C. Staining of the sex body in wt and Smc1b−/− spermatocyte chromosome spreads using anti SYCP3 for AEs, and anti SUMO-1 for the sex body or unsynapsed chromatin domains. D. Example of DAPI-staining and analysis by light microscopy for identification of pachytene cells in the FISH experiment shown in Fig. 1D. (TIFF)

Figure S2 Flow cytometry analysis of Hoechst 33342-stained testicular cells of mice aged 16 dpf. A. Scheme of the distribution of spermatogenic subpopulations displays several groups of cells during the process of spermatogenesis. Cell populations are divided into spermatogonia (Spg), cells in premeiotic S phase and preleptotene (Pre-L), spermatocytes I (Spc I) and spermatocytes II (Spc II). B–D Total testis cells of an adult WT mouse (B) and juvenile WT and Smc1b−/− mice aged 16 dpf (C and D) were stained with Hoechst 33342 (Hoechst) and analyzed by flow cytometry. Propidium iodide allowed exclusion of dead cells (not shown). Detection of Hoechst emission using two different optical filters (Alexa Fluor 350 and Emerald 360) facilitated the distinction of cell populations of the testsis. An uncharacterized population is enclosed by a dashed line and likely represents zygotene/pachytene cells present in the Smc1b−/− testsis. Percentages of cell populations are represented with exclusion of the smear of events in the lower left corner. Scheme in A adapted from [80]. (TIFF)
Figure S3 Spatial clustering of differentially expressed genes. Genomic domains significantly enriched for up- or down-regulated genes. Red dots indicate regions identified through conducting a two-sided Wilcoxon rank sum test inside sliding windows (window size: 10 genes). Green dots indicate regions identified through a three-state Hidden Markov Model. Significance of both types of regions was established via permutation testing resulting in False Discovery Rates (FDR). Regions with FDR<0.2 are shown for the Wilcoxon test. Dots above (below) the x-axis indicate significant clustering of up-regulated (down-regulated) genes in the respective region. Vertical black bars show fold changes of individual genes. The scale of the y-axis represents -log10(FDR) values for the genomic domains and log fold changes in case of differential gene expression. Chromosome numbers are displayed on the right side of each x-axis.

Figure S4 Gene expression fold changes in testes of 16 dpp Smc1b/-/- mice. Log2 transformed gene expression fold changes are displayed for all chromosomes (A) and as examples for chromosomes 1, 19 and X individually (B-D).

Figure S5 Comparative expression analysis of miRNAs. Levels of the displayed mature miRNAs are at least 1.5-fold increased or decreased in the testes of at least one of three 16 dpp Smc1b/-/- mice compared to wt littermates. The graph depicts miRNA ratios (n = 3, mean±s.e.m.). Dashed lines correlate with 1.5-fold increase or reduction.

Figure S6 Anti pSMC3 staining control. Since the anti pSMC3 staining showed only a cloud-like pattern and not a specific axis staining as the pSMC1, we controlled for the specificity of the antibody, although both were described before, by blocking experiments using either phosphorylated epitope peptide or the non-phosphorylated peptide. Wt spermatocyte chromosome spreads were stained with anti SYCP3, anti pSMC3, and DAPI. A the antibodies were co-incubated with a phospho-SMC1 peptide or a phospho-SMC3 peptide. B. Staining of metaphase I chromosome spreads with anti SYCP3, anti pSMC3, and DAPI.

Figure S7 Cohesin loading does not depend on SPO11. A. SMC1α localization in wt and Spo11/-/- spermatocytes. Spermatocyte chromosome spreads stained with anti SYCP3 for AEs and anti HORMAD1 for unsynapsed sex chromosome axes or unsynapsed autosomes and with anti SMC1α. Inset shows preferential localization of SMC1α to synapsed (HORMAD1-negative) regions. B. SMC3 localization in Spo11/-/- spermatocytes. Spermatocyte chromosome spreads were stained with anti SYCP3 for AEs and anti HORMAD1 for unsynapsed sex chromosome axes or unsynapsed autosomes and with anti SMC3.

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