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Human Male Meiotic Sex Chromosome Inactivation

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Abstract

In mammalian male gametogenesis the sex chromosomes are distinctive in both gene activity and epigenetic strategy. At first meiotic prophase the heteromorphic X and Y chromosomes are placed in a separate chromatin domain called the XY body. In this process, X,Y chromatin becomes highly phosphorylated at S139 of H2AX leading to the repression of gonosomal genes, a process known as meiotic sex chromosome inactivation (MSCI), which has been studied best in mice. Post-meiotically this repression is largely maintained. Disturbance of MSCI in mice leads to harmful X,Y gene expression, eventuating in spermatocyte death and sperm heterogeneity. Sperm heterogeneity is a characteristic of the human male. For this reason we were interested in the efficiency of MSCI in human primary spermatocytes. We investigated MSCI in pachytene spermatocytes of seven probands: four infertile men and three fertile controls, using direct and indirect in situ methods. A considerable degree of variation in the degree of MSCI was detected, both between and within probands. Moreover, in post-meiotic stages this variation was observed as well, indicating survival of spermatocytes with incompletely inactivated sex chromosomes. Furthermore, we investigated the presence of H3K9me3 posttranslational modifications on the X and Y chromatin. Contrary to constitutive centromeric heterochromatin, this heterochromatin marker did not specifically accumulate on the XY body, with the exception of the heterochromatic part of the Y chromosome. This may reflect the lower degree of MSCI in man compared to mouse. These results point at relaxation of MSCI, which can be explained by genetic changes in sex chromosome composition during evolution and candidates as a mechanism behind human sperm heterogeneity.


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Introduction

During mammalian male meiosis, the heteromorphic sex chromosomes (X and Y) condense in a separate chromatin domain known as the XY or sex body (reviewed in [1]). XY body formation starts at the zygotene to pachytene transition of prophase I [1] and persisting DNA double strand breaks (DSBs), made by SPO11 at the onset of meiosis, contribute to its formation [2]. Autosomal chromosomes are able to repair these breaks by searching for their homolog, subsequently forming a so-called bivalent in which paternally and maternally derived chromatids are closely aligned, intermediated by the synaptonemal complex (SC). However, the X and Y chromosomes are largely heterologous and show homologous synopsis only at the small pseudoautosomal region (PAR). Non-synapsis of chromosomal axial elements is detected by BRCA1 which localises ATR, a member of the PI3-like kinase family, to the X,Y chromosomes, followed by phosphorylation of H2AX at serine 139 [3]. Hence, from its formation on, the XY body is positive for γH2AX [4].

Phosphorylation of H2AX initiation represses transcription of genes on the sex chromosomes [5], a process called meiotic sex chromosome inactivation (MSCI). Transcriptional repression of the XY body was first identified by 3H-uridine incorporation [6,7], and further explored by (Cot-1) RNA FISH [8] and microarray analysis [5,9,10]. MSCI possibly occurs to prevent activation of the pachytene checkpoint at stage IV/V of the seminiferous epithelium in the mouse [11,12]. MSCI is supposed to be a general phenomenon among mammals (mouse, hamster [19]) including the human [14,15], and insects [13] and is observed in bird species as well [16]. A similar silencing process can take place at autosomal chromatin, referred to as meiotic silencing of unsynapsed chromatin (MSUC) and functions in the elimination of aberrant meiocytes bearing asynaptic autosomes [17]. MSCI is thought to be an evolutionary derivation of the more general process of MSUC [8].

Probably as a result of H2AX phosphorylation, many DNA repair proteins are attracted [18,19] and major changes take place at XY body chromatin, including histone posttranslational modifications (PTMs) and incorporation of histone variants [20,21]. In a previous study, we have shown for mice that specifically at XY chromatin and starting in early pachytene, nucleosomes containing H3.1/3.2 are replaced by nucleosomes containing H3.3 [22]. Nucleosome eviction is complete at the end of mid pachytene (stage V) [22]. Existing Histone 3 and 4 N-tail PTMs disappeared in this process and were only selectively placed back [22], allegedly to create a sex chromosome-specific chromatin composition that most likely has a function in MSCI and, by transmission over the meiotic divisions to round
spermatids [22], in post-meiotic sex chromosome repression (PSCR [20]) [9]. Microarray analysis, followed by RT-PCR, demonstrated that 87% of X genes repressed during MSCI remain repressed in PMSC (post meiotic sex chromatins) [9]. The fact that in the mouse MSCI and PSCR share several features, like the same repressive histone markers and the presence of H3.3 containing nucleosomes, indicates that PSCR is a downstream consequence of MSCI.

In the human XY body, features such as a condensed chromatin domain [23] and staining for H2AX, BRCA1 [24] and ATR [25], were shown to be present as well. Also an indication for H3.1/3.2 nucleosome eviction was obtained [22]. Cytotoxic tests (acidin-orange, methylgreen pyronin) [23] and autoradiography with 3H-uridine incorporation [14,15] show the absence of RNA in the human XY body thereby indicating MSCI to be present in the human. However, in a recent whole testis cDNA array analysis, MSCI could be demonstrated for the chimpanzee, but not for the human [26].

In the mouse there are extensive data documenting the association of unsynapsed autosomal segments with the XY body at first meiotic prophase. In these chromosome mutants, prophase progression, the meiotic divisions, and spermiogenesis are affected, resulting in a reduction of the number of round spermatids and a deviant heterogeneous sperm population as to morphology and motility [10,27–29]. A number of observations in these mutants suggest MSCI to be compromised by the MSUC that now takes place in association with MSCI (RNA autoradiography [30] gene specific RNA FISH, [29] RNA chip analysis [10] and incomplete eviction of H3.1/H3.2 containing nucleosomes [22]). These data indicate that in mouse proper MSCI is essential for the progression of pachytene, first meiotic prophase/metaphase and influences spermiogenesis. In mice with complete avoidance of MSCI, all spermatocytes are removed by the pachytene checkpoint at the mid to late pachytene transition [12]. In human carriers of translocations a similar association of asynaptic autosomal chromosome segments to the XY body at pachytene is observed. In all cases, these men were infertile [24] (reviewed in [31]).

Additional support for the effect of disrupted XY expression on spermiogenesis in mouse comes from two sources: a) In the H/δδ knockout model which is characterised by poor spermatid elongation, hence male sterility, there is a general increase in post-meiotic transcription for X-linked genes [32,33], b) In a series of Y long arm deletions, from two third to complete, post-meiotic sex chromatin showed more deviant histone and chromatin parameters when the deletion was bigger [34], in line with upregulation of YX genes in round spermatids [35]. Spermatid nucleus chromatin condensation is affected and so is, for the more severe deletions, the frequency of karyotype abnormalities in embryos generated by intra cytoplasmic sperm injection (ICSI) [36].

Here we have studied cytological parameters of MSCI in man over a number of probands. The main reason for this study is that human sperm samples are known to be variable between and within men [37,38] a variability that is increasing when sperm numbers are decreasing [39,40]. Hence, the spermiogram (number, morphology, and motility of spermatzoa) of many human probands resembles that of mouse chromosome mutants in which impaired MSCI is indicated, a variability never seen in chromosomally normal mice. A second reason is that the spermatogenic phenotypes of men with AZFc deletions suggest the Y chromosome to be involved in spermiocyte development: If MSCI is systematically operative in our species, one would not expect the deletion to be of consequence for meioyte survival.

As a marker for MSCI we studied H3.1/3.2 nucleosome eviction. Variation was observed in the degree of nucleosome eviction in late pachytene spermatocytes. Complementary to nucleosome eviction and accompanying histone X-tail post translational modifications, we used Cot-1 RNA FISH and incorporation of the uridine analog 5-ethynyluridine into newly transcribed RNA as direct transcription markers. Our results point at both intra- and inter-proband heterogeneity in MSCI and at the absence of meiotic selection steps in the generation of round spermatids. In the discussion, we relate our observations to the roles of the sex chromosomes in spermatogenesis and to the influence of mating systems on sperm heterogeneity.

**Results**

**Variation in H3.1/3.2 nucleosome eviction in XY bodies of human pachytene spermatocytes**

We examined primary spermatogenic cell samples of 3 patients and 3 controls which were obtained from testicular biopsies (the controls were of proven fertility, see Materials and Methods; Table S1). To study nucleosome eviction in human XY bodies, we stained pachytene spermatocytes for histone isoforms H3.1/3.2 and SYCP3 (as a marker for the SC and X,Y axial elements). Staining patterns were recorded by immunofluorescence (IF) microscopy. A first sign of H3.1/H3.2 loss was observed at mid pachytene when the XY axial elements locally start to unravel. (type III, [25,41] (Figure 1A). In late pachytene XY bodies (types IV and V [25,41]), when the XY axial elements appeared as a tangled knot of fibrils and DAPI was most intense, a highly variable H3.1/3.2 signal remained (Figure 1B, Table 1). We categorized the signals in four classes based on increasing intensity and coverage (see Materials and Methods). A minority of nuclei with a complete loss of signal for H3.1/3.2 containing nucleosomes were found (Figure 1B, Table 1), indicating a high degree of H3.1/3.2 remodelling. Remarkably, in these nuclei the border of the XY body was not as sharply defined as in our mouse samples [22]. Also between individuals in the patient group and in the control group, variation in H3.1/H3.2 staining was observed. When group data were pooled, no difference was observed between the degree of nucleosome eviction in controls versus patients (Table 1). The nuclear localization of the XY body was determined for all pachytene spermatocytes analyzed in Table 1 to investigate if there is a relation between nuclear localization and H3.1/3.2 nucleosome eviction. Figure S1 shows that XY bodies of nuclei belonging to classes ‘low’ and ‘mid 1’ nucleosome eviction are less often localized at the border of the nucleus compared to XY bodies of nuclei with a higher degree of nucleosome eviction (classes ‘mid 2’ and ‘high’). However, in every nucleosome eviction class, all possible XY body locations are represented. This indicates that our results are not a reflection of a technical artefact, such as overlaying autosomal chromatin that by DAPI and SYCP3 staining was judged to be rare: all chromatin domains are in one focal plane.

To check for enrichment of H3.3 containing nucleosomes, we probed primary spermatocyte nuclei for H3.3S31ph (due to lack of an appropriate antibody detecting unmodified H3.3) and H3.1/ H3.2. The high signal for H3.3S31ph in the late pachytene XY body showed that nucleosomes containing H3.1/3.2 were indeed replaced by nucleosomes containing H3.3 (Figure 1C). Variation in signal intensity was observed for H3.3S31ph (n = 20, P2, Table S1). However, this showed no relation to the loss of H3.1/3.2 (not shown).

To conclude, we observed a considerable variation in gonosomal chromatin remodeling during pachytene, both within and between probands, contrasting strongly with our previous observations in mice [22].
Human Male Meiotic Sex Chromosome Inactivation

A early pachytene  mid pachytene  C late pachytene

B four groups of nucleosome eviction in late pachytene spermatocytes

low  mid 1  mid 2  high
No H3.1/3.2 nucleosome replacement at Yq heterochromatin

In about 50% of the XY bodies with the highest H3.1/H3.2 nucleosome replacement (n = 41, patients and controls, Table 1), we noticed a protrusion of H3.1/3.2 signal, probably of sex chromatin origin (Figure 1B). An axial element loop emanating from the SYCP3-stained tangled knot reached towards or into this protrusion. The loop was observed in 65% of all late pachytene spermatocytes (n = 526, Table 1; patients: 62%; controls: 70%). We performed FISH with an X chromosome painting probe and a Yq heterochromatin (Yqh) probe followed by IF analysis to determine whether there could be a systematic overlap between X and/or Y chromatin signals and H3.1/3.2 signal (Figure 2). From individual C3, 33 late pachytene nuclei were scored of which five showed highest nucleosome eviction. In four of these a clear H3.1/3.2 protrusion was observed, with a co-localizing Yqh probe signal (Figure 2A). In the other 28 nuclei with an intermediate to low degree of H3.1/3.2 nucleosome eviction, overlapping signals were difficult to analyse because of remaining H3.1/3.2 nucleosomes in the XY body. However, Yqh signal was always found together with H3.1/3.2 signal (Figure 2B, C). In some of these nuclei a protrusion could be observed as well (Figure 2B). As expected, the X chromosome showed variable H3.1/3.2 nucleosome eviction (Figure 2A–C). Overall, in nuclei with highest nucleosome eviction, H3.1/3.2 signal was only left on Yqh and at the edges (Figure 2C).

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<th>Distribution of late pachytene spermatocytes among low to high nucleosome eviction classes (%)</th>
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Patients (P), Controls (C).

1Heterogeneity between patients \( \chi^2 \text{ df. } 6: 13.6, p = 0.035 \).
2Heterogeneity between controls \( \chi^2 \text{ df. } 6: 31.2, p < 0.001 \).

Table 1. Degree of sex chromatin H3.1/3.2 nucleosome eviction in patients and control men.

H3.1/3.2 nucleosome replacement and H2AX phosphorylation at S139

\( \gamma \text{H2AX} \) is a marker of sex chromatin during pachytene, indicating chromatin remodelling leading to meiotic sex chromosome inactivation (MSCI) [5]. We were interested to study whether there is a relation between the degree of sex chromatin nucleosome eviction and the intensity of the \( \gamma \text{H2AX} \) signal. In early and mid pachytene a strong \( \gamma \text{H2AX} \) signal was consistently found in the XY body (n = 74, Figure 3A). In the majority of late pachytene nuclei with low XY nucleosome eviction, a strong to mid \( \gamma \text{H2AX} \) signal remained, extending into H3.1/3.2 positive sex chromatin (Figure 3C,D,E). In nuclei with higher nucleosome eviction the \( \gamma \text{H2AX} \) signal became weaker (Figure 3B,E). Also the area of XY chromatin which was stained by \( \gamma \text{H2AX} \) became smaller and usually was in the centre of the XY body (Figure 3B,E). When a H3.1/3.2 Yq heterochromatin protrusion was identified in highly remodelled XY bodies (44%, n = 41) the overlaying \( \gamma \text{H2AX} \) signal was often weaker (72%, n = 18, Figure 3B). Complete absence of the \( \gamma \text{H2AX} \) signal was never observed in late pachytene and as human meiosis has no significant diplotene stage [25,41,42] we investigated metaphase I nuclei. This is a rare

Figure 1. H3.1/3.2 nucleosome eviction in human pachytene spermatocytes. A H3.1/3.2 nucleosome eviction was never observed in early pachytene spermatocytes (Type I) [23,41]. First signs of H3.1/3.2 loss were detected in mid pachytene spermatocytes. B Illustration of four degrees of nucleosome eviction distinguished in late pachytene XY bodies. Arrows indicate the bulge of H3.1/3.2 invading the XY body. Arrowheads indicate the axial element loop reaching towards the H3.1/3.2 bulge. C Simultaneous loss of H3.1/3.2-containing nucleosomes and formation of nucleosomes containing H3.3S31ph at meiotic sex chromatin. Arrow indicates the H3.1/3.2 bulge, arrowhead points at the absence of enrichment for H3.3S31ph at the H3.1/3.2 bulge signal.

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Figure 2. Differential H3.1/3.2 nucleosome remodelling between Yq heterochromatin and the X chromosome in late pachytene spermatocytes. A Late pachytene nucleus showing a high degree of nucleosome remodelling for the X chromosome, but no remodelling for Yqh. B Late pachytene nucleus showing an intermediate degree of nucleosome remodelling for the X chromosome, but no remodelling for Yqh. C Late pachytene nucleus showing a low degree of nucleosome remodelling for the X chromosome, but no remodelling for the Yqh.

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Table 1. Degree of sex chromatin H3.1/3.2 nucleosome eviction in patients and control men.
stage in spreading preparations of human spermatogenic cell samples as well. In those found, we did not observe any γH2AX signal (n = 4, not shown). To conclude, in human late pachytene nuclei the disappearance of γH2AX is related to the degree of nucleosome eviction, which constitutes another difference from mice, in which the sex bivalent is characterized by heavy γH2AX staining up to the diplotene stage [4].

Meiotic sex chromosome silencing

In the mouse, we suggested H3.1/3.2 nucleosome replacement to be functional in MSCI [22]. If this supposition is right, then MSCI would constitute a much more variable phenomenon in the human. To investigate MSCI in man, we first stained patient and control nuclei for SYCP3 or H3.1/3.2 in combination with RNA polymerase II (RNA pol II), an indirect marker of transcription. In mouse it was shown that RNA pol II is largely absent at sex chromatin of late pachytene nuclei, while it is strongly present on autosomal bivalents [43,44]. Our results on human primary spermatocytes, illustrated and graphically represented in Figure 4, show variation in the intensity/presence of the RNA pol II signal on the XY body in late pachytene spermatocytes (compare Figure 4A,B) and suggest a relation between the degree of H3.1/3.2 nucleosome eviction and the amount of RNA pol II demonstrable (Figure 4C). Nuclei with low nucleosome replacement show more RNA pol II than nuclei with high nucleosome replacement. However, in nuclei with high nucleosome eviction we never saw complete absence of RNA pol II signal. Notably, in nuclei with high H3.1/3.2 nucleosome loss no distinction in RNA pol II signal intensity was observed between Yqh (the H3.1/3.2 protrusion) and remaining XY chromatin.

Second, we performed RNA FISH using Cot-1 DNA as a probe. The Cot-1 probe is used to visualize repetitive sequences in intronic and 3′ untranslated regions which are present throughout the genome and emerge upon gene transcription into pre-spliced RNA [8]. To demonstrate genomic coverage, we performed DNA FISH with the Cot-1 probe on human lymphocyte metaphase nuclei and pachytene spermatocytes. In both types of nuclei the complete chromatin, including X and Y, was covered by Cot-1 DNA signal (Figure S2A,C), the signal being stronger for constitutive centric heterochromatin.

Figure 3. γH2AX in early and late pachytene XY bodies. A Early pachytene nucleus. The complete X and Y chromatin is covered by γH2AX. B Late pachytene XY body with a high degree of H3.1/3.2 nucleosome remodelling and a decreased γH2AX signal (very weak, <H3.1/3.2 depleted area). A clear bulge of H3.1/3.2 signal is present (arrow) with almost no overlaying γH2AX signal. H3.1/3.2 bulge indicated by the dotted lines. C Late pachytene XY body with a low degree of H3.1/3.2 nucleosome remodelling and a strong γH2AX signal (> H3.1/3.2 depleted area). D Histogram displaying the relation between the degree of H3.1/3.2 nucleosome remodelling and the intensity of the γH2AX signal, classified into four categories (bars) (n = 203, P1-3, C3). E Histogram displaying the relation between the degree of H3.1/3.2 nucleosome remodelling and the area of XY chromatin covered by γH2AX, classified into four categories (bars) (n = 203, P1-3, C3).

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To detect transcripts in the XY body, we combined RNA FISH with IF for SYCP3 in P2,3 and C3 (see Table S1). Sertoli cell nuclei were the most heavily stained, corresponding to their active state, while at meiotic metaphase no signal was present (not shown). As expected, very weak RNA signals were seen in the early meiotic prophase stages (leptotene, zygotene). In early pachytene nuclei, the signal intensity increases for the autosomal bivalents (not shown). Over the late pachytene XY bodies of all three men, there was a clear reduction of Cot-1 RNA signal, contrasting to the autosomes. However, variation was observed (Figure 5A: low; B: high). Of all late pachytene nuclei scored (n = 117), 4% showed high Cot-1 RNA staining over the XY body, 54% an intermediate (mid) staining and 42% a low Cot-1 RNA signal. In all early, mid pachytene nuclei in which the XY axial elements were well recognizable, a decrease in Cot-1 RNA was observed (n = 38, Figure 5F).

Third, we cultured spermatogenic cells derived from a fresh TESE biopsy (P4, Table S1, Materials and Methods) with the uridine analog EU. Pilot experiments indicated acid fixation and spreading to better preserve the RNA signal than did basic (PFA) fixation and spreading. Identification of late pachytene nuclei could still be based on the prominent DAPI intensive XY body (Figure 6A,B). Consistent with the results from Cot-1 RNA FISH, Sertoli cells displayed the strongest EU signal and meiotic metaphase nuclei displayed no signal (not shown). In meiotic nuclei without a DAPI intense XY body, autosomal EU signals were low (not shown). In nuclei with an XY body, the autosomal signal was, in most cases, stronger. Late pachytene nuclei were variable in RNA transcriptional activity over the XY body (Figure 6, compare A with B). About 5% of nuclei showed a relatively high production of RNA on the XY body (Figure 6B), 42% an intermediate (mid) expression and 53% of the nuclei displayed only low RNA expression (Figure 6A) (n = 60, P4, Table S1). Complete absence of EU signal was never observed. Taken together, both indirect (by RNA Pol II IF) and direct demonstrations of transcription (by Cot1 RNA FISH and EU incorporation) do show MSCI to be variable between human primary spermatocytes. Double labelling for H3.1/3.2 showed a correlation between higher gonosomal transcription and lower nucleosome remodelling, supporting our original supposition of a mechanistic link between this phenomenon and MSCI [22].

**Figure 4. Variation in presence of RNA polymerase II in late pachytene XY bodies.** A Late pachytene nucleus, showing no decrease in RNA pol II signal intensity at the XY body compared to the autosomal chromatin. Nuclei were stained with the mouse monoclonal antibody. B Late pachytene nucleus, showing a strong decrease in RNA pol II signal intensity at the XY body compared to the autosomal chromatin. Nuclei were stained with the mouse monoclonal antibody. C Histogram displaying the relation between the degree of nucleosome eviction and the decrease of RNA pol II signal at the XY body, classified into four categories (bars). Total absence of RNA Pol II signal was never observed. Nuclei were stained with the rabbit polyclonal antibody (n = 106, P2, C1-3, Table S1).

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Human Male Meiotic Sex Chromosome Inactivation

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Figure 5. Cot-1 RNA expression in pachytene spermatocytes. A Late pachytene nucleus displaying a low Cot-1 RNA signal on XY. B Late pachytene nucleus displaying a higher Cot-1 RNA signal on XY. C Late pachytene nucleus showing a high degree of H3.1/3.2 nucleosome eviction and a low Cot-1 RNA signal on XY. D Late pachytene nucleus showing an intermediate (mid) degree of H3.1/3.2 nucleosome eviction and intermediate (mid) Cot-1 RNA signal on XY. E Late pachytene nucleus showing a low degree of nucleosome eviction and high Cot-1 RNA signal on XY. F Early pachytene nucleus displaying low Cot-1 RNA expression on XY. G Early or mid pachytene nucleus before H3.1/3.2 nucleosome eviction but with an intermediate (mid) Cot-1 RNA signal. H Histogram showing the relation between the degree of H3.1/3.2 nucleosome eviction and the level of Cot-1 RNA signal (bars) (n = 134, P2, C2,3).

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Transmission of incompletely H3.1/3.2-evicted sex chromosomes to the round spermatid stage

In the mouse, disturbances in MSCI have been associated with the pachytene checkpoint and spermatocyte death (see introduction). Therefore we were interested to discover if at the meiotic divisions there has been selection for spermatocytes with more fully remodelled XY bodies to reach the round spermatid stage. Therefore, we first studied meiotic metaphase I/II nuclei, of which we found 32. In these, some gonosomal H3.1/3.2 remnants were always observed, concordant with the varying degree of replacement found in late pachytene (Figure 7B). The best XY remodelled nuclei showed a little cloud of H3.1/3.2 on a more DAPI intense sex chromatin domain, reminiscent of the absence of nucleosome eviction for Yqh (Figure 7A). These results suggest that incomplete XY H3.1/3.2 replacement does not influence the efficiency of the meiotic divisions, as it does in mice. Consequently, round spermatids with incompletely remodelled sex chromatin must be present.

We utilized IF/FISH analysis (X chromosome painting probe, Yqh probe combined with IF staining of H3.1/3.2) on proband C3 to analyze round spermatids, that were identified by nuclear morphology and by the presence of only one FISH signal (X or Y, Figure 7C,D). Yqh FISH confirmed the lack of nucleosome replacement for this region (Figure 7C). The X chromosome painting probe allowed the study of nucleosome eviction over the whole chromosome (n = 54) (Figure 7D,E). In 40% of spermatid nuclei, the degree of evicted H3.1/3.2 containing nucleosomes was low. In 43% of nuclei there was an intermediate degree of H3.1/3.2 nucleosome eviction (example in Figure 7D) and in the remaining nuclei (17%) the degree of nucleosome eviction was high. These findings were compatible with the interpretation of nucleosome eviction in late pachytene spermatocytes of this proband (Table 1). In all, the data support the concept that meiocytes metamorphose into spermatids irrespective of the level of gonosomal nucleosome eviction.

Histone characteristics of human meiotic and postmeiotic sex chromatin

Because of the contracted nature and heterochromatic appearance (judged by DAPI staining) of late pachytene spermatocyte XY bodies in the human and mouse and the prominent presence of H3K9me3 in late pachytene XY bodies in the mouse [22], we studied several histone modifications in combination with H3.1/H3.2 staining (Figure 8, Table 2, Figure S3). For both H3K9me3 and H3K4me2 detection, two independent antibodies (see Materials and Methods) were used which did not differ in staining pattern (data not shown). Based on our results and on IF staining patterns that are commonly observed in mouse [9,22], the histone 3 N-tail methyl PTMs tested for could be divided in two groups: 1) H3K9me2,3 and H4K20me3, which on autosomes at late pachytene yielded a heterochromatin staining pattern (determined by co-staining with the centromere-specific anti-sperm CREST, data not shown) (Figure 8 A–C); 2) Histone modifications that showed an even autosomal staining, such as H3K4me2 and H3K27me2,3. The latter marks disappeared from the sex chromatin during nucleosome eviction and were also lost from Yqh while H3.1/3.2 was still present (Figure 8D–F, Table 2, Figure S3C–E).

H3K9me3 decreased over the XY body in conjunctive with H3.1/H3.2 loss as well, but more selectively as often clearly stained domains remained (Figure 8B; Table 2, Figure S3B). An identical pattern was observed for H4K20me3 (Figure 8C). These domains (for H4K20me3 often more than one per XY body) were located in the periphery of the DAPI-intense sex chromatin. When a H3.1/3.2 proportion (i.e. Yqh) was present (n = 37), colocalization with H4K20me3 was observed in 95% of cases (Figure 8C). In only 12% of the H4K20me3-stained nuclei (n = 115) the signal was absent from the XY body. Similarly, H3K9me3 was present in all observed Yqh protrusions (Figure 8B) (11%, n = 146) and in seven cases exclusively so.

The staining pattern of H3K9me2 differed from the H3K9me3 and H4K20me3 heterochromatin markers, as no specific domains were associated with the XY body. We observed three types of H3K9me2 staining patterns in late pachytene spermatocytes (Figure S3A) without a clear link to H3.1/3.2 nucleosome eviction. However, a trend is visible towards more H3K9me2 staining on XY in higher H3.1/3.2 remodelled nuclei (Figure S3A, Table 2).

In early round spermatids (Table 2), the sex chromosome signals for histone modifications resembled those observed in late pachytene XY bodies. Least variability was found for modifications that tended to be low at XY during/after nucleosome eviction such as H3K4me2, H3K27me2 and H3K27me3. H3K9me2 reached higher coverage over the sex chromosomes than did H3K9me3, which was faintly present and showed stronger domains next to the H3.1/3.2-evicted PMSC, as in late pachytene. In the majority of spermatid nuclei H4K20me3 seemed to be faintly present over the X or Y, while this was not observed at late pachytene. Furthermore, similar domains were associated with the PMSC as found for H3K9me3 staining. In most cases H3.3S31ph was stronger on the sex chromatin compared to autosomal chromatin. We never observed H2AX in round spermatids.

In conclusion, with the exception of Yqh, the sex chromosomes at late pachytene miss an overt heterochromatic signature as based on H3 and H4 N-tail modifications. In early round spermatids, this pattern is maintained with the addition of H4K20me3 that might signal heterochromatinization.
In the present study we show by indirect and direct cytological methods (H3.1/3.2 nucleosome eviction and presence of RNA polymerase II, EU incorporation and Cot-1 RNA FISH) MSCI to occur in man. In late pachytene, H3.1/3.2 nucleosome eviction is correlated with the repression of Cot-1 RNA expression. Using the loss of H3.1/3.2 as an indirect marker for MSCI, variation was detected between and within patients and controls. Cot I RNA FISH and EU incorporation confirmed this observation. This variability did not occur for H3.1/3.2 nucleosome eviction in spermatocytes from Swiss random bred mice [22]. To follow up and confirm this observation, we checked H3.1/3.2 nucleosome eviction in spermatocytes from a Swiss derived outbred NMRI mouse and a Swiss derived inbred FVB mouse. Similarly, both mice had homogeneously H3.1/3.2 evicted late pachytene XY bodies that by this characteristic usually were sharply demarcated (data not shown). Variation in MSCI has neither been specifically noticed in 3H-uridine uptake studies using inbred C3H mice and random bred Swiss mice [6,30].

In the human, variation was most apparent at the circumference of the XY body that is much more fuzzy, as observed by DAPI staining, compared to mouse. Also stainings by H3,H4 PTMs that mark heterochromatin failed to uniformly label gonosomal chromatin at late first meiotic prophase, with the exception of Yqh. Our notion that nucleosome remodelling is linked to MSCI is strengthened by its absence in Yqh. Apparently, this domain is

**Discussion**

**MSCI in the human male**

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Figure 8. Histone characteristics of the XY body. A–F Late pachytene nuclei are shown with a mid-2 to high degree of H3.1/3.2 nucleosome eviction. In the merge panel DAPI, H3.1/3.2 and the PTM are depicted. **A** H3K9me2 was strongly present on XY chromatin including Yqh, as indicated by the brighter DAPI domain (arrowhead). **B** H3K9me3 was present on the Yqh H3.1/3.2 bulge (arrow) and irregular on other XY chromatin. **C** Two H4K20me3 stained domains were found in and associated with X,Y chromatin and were co-staining with H3.1/3.2. Part of the upper domain was located in brighter DAPI XY body chromatin (arrowhead) reminiscent to Yqh. **D–F** H3K4me2, H3K27me2 and H3K27me3 were strongly decreased in XY chromatin, including Yqh as indicated by the dotted lines.

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sufficiently made incapable of mechanistic transcription by its intrinsic chromatin structure, illustrated by the histone PTM stainings. In the mouse, pachytene XY body localization is related to the pachytene substage [15]. The later in pachytene, and clearly from stage IV on, the more the XY body was found in the periphery of the nucleus. For chromatin remodelling, we studied

Table 2. Histone variant- and histone N-tail-modification dynamics at meiotic and post-meiotic X,Y chromatin.

<table>
<thead>
<tr>
<th>Histone modification</th>
<th>General dynamics on X,Y</th>
<th>Appearance on Yqh</th>
<th>Early round spermatids</th>
</tr>
</thead>
<tbody>
<tr>
<td>H3K9me2&lt;sup&gt;1&lt;/sup&gt;</td>
<td>increase (n = 176)</td>
<td>present</td>
<td>+/− or + (n = 82)</td>
</tr>
<tr>
<td>H3K9me3&lt;sup&gt;1&lt;/sup&gt;</td>
<td>decrease, domains (n = 146)</td>
<td>present</td>
<td>− or +/−, domains (n = 95)</td>
</tr>
<tr>
<td>H4K20me3&lt;sup&gt;1&lt;/sup&gt;</td>
<td>decrease, domains (n = 115)</td>
<td>present</td>
<td>− or +/−, domains (n = 115)</td>
</tr>
<tr>
<td>H3K27me2&lt;sup&gt;2&lt;/sup&gt;</td>
<td>disappears (n = 182)</td>
<td>disappears</td>
<td>− (n = 107)</td>
</tr>
<tr>
<td>H3K27me3&lt;sup&gt;2&lt;/sup&gt;</td>
<td>strong decrease (n = 171)</td>
<td>strong decrease</td>
<td>− or +/− (n = 106)</td>
</tr>
<tr>
<td>H3K4me2&lt;sup&gt;2&lt;/sup&gt;</td>
<td>disappears (n = 75)</td>
<td>disappears</td>
<td>− or +/− (n = 27)</td>
</tr>
</tbody>
</table>

Histone variant modification

| H3.3S31ph | strongly present (n = 10) | present | + (n = 17) |
| γH2AX | decreases (n = 203) | decreases | − * |

<sup>1</sup>group 1: histone lysine methyl modifications displaying autosomal heterochromatin staining.
<sup>2</sup>group 2: histone lysine methyl modifications displaying autosomal even staining.

In parenthesis: number of nuclei.

*No quantitative analysis was performed on γH2AX staining in early round spermatids as this modification was never observed.

Modifications were studied in P1-3 and C3, except for H3K4me2 which was studied in P2 and C3, and H3.3S31ph which was studied in P2.

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late pachytene spermatocyte XY bodies only, which according to the available mouse data [45] should have a preference for the nuclear periphery. However, we found XY bodies to be located over the whole spread area. Low H3.1/3.2 remodelled late pachytene XY bodies tended to be located more towards the centre compared to highly remodelled late pachytene XY bodies that were more often found near the periphery. One explanation for this phenomenon could be that for maintenance of MSCI a peripheral localization is advantageous.

In the mouse, MSCI commences at early pachytene [6]. Cot-1 RNA FISH in combination with staining for SYCP3 revealed that from early to late pachytene, Cot-1 RNA was reduced at the human XY body. The first sign of gonosomal H3.1/3.2 loss in human spermatocytes was at mid pachytene, which is different from the mouse, where H3.1/3.2 nucleosome eviction starts at early pachytene [22]. These results indicate that also in the human, H3.1/3.2 nucleosome eviction follows X,Y gene silencing, though with a greater time lapse.

The general idea is that MSCI is initiated by the cascade of events following H2AX S139 phosphorylation [5] the spreading of which is assisted by MDC1 [19]. Strong γH2AX staining is present in both human and mouse early pachytene spermatocytes, indicating that initiation of MSCI is similar in human and mouse. This is reflected by an inactive XY body in early pachytene for both human (this study) and mouse [6]. The variation in MSCI we observed in late pachytene might therefore be caused by variation in the maintenance of silencing in which H3.1/3.2 nucleosome remodelling (and resulting histone PTM changes) might have a role, concomitant with the chromatin domain moving away from the transcriptionally active autosomal bivalents.

Could the decrease in γH2AX staining we observed in human late pachytene be involved in maintenance variation of MSCI? In the mouse, staining continues into diplotene and is lost at first meiotic metaphase, which is different in the human (who are missing a substantial diplotene stage [25,41,42]). Here this mark is reduced up to a very low level when nucleosome remodelling is maximal (late pachytene). For the mouse, an interpretation could be that this modification helps in maintaining MSCI. In the human our results do not support this view, as the human XY bodies showing low H2AX S139 phosphorylation were better nucleosome remodelled and on average showed a lower Cot-1 RNA signal.

No selection for incomplete H3.1/3.2 evicted pachytene spermatocytes

In mouse chromosome mutants in which the association of asynapsed chromosome segments with the XY body interferes with MSCI, pachytene and metaphase I death usually follows [10,28]. Assuming that this typical pattern of spermatocyte death is related to expression of gonosomal genes, one could ask if such a selection step does occur in human spermatogenesis. In the light of the variation in MSCI indicated by our data, this would not seem likely. We observed variation in X chromosomal H3.1/3.2 depletion in round spermatids, reproducing the variegated pattern found in late pachytene. Also around the Yqh, variation in an adjacent H3.1/3.2 depleted area was witnessed. Hence, there does not seem to be a clear selection against primary spermatocytes with respect to chromatin remodelling, i.e. the global transcriptional status of the sex chromosomes. Observations on XY chromatin remodelling at first and second meiotic metaphase on average yielded the impression of a more fully executed remodelling process, but traces of H3.1/3.2 outside the Yqh area were found. The more fully remodelled impression of the metaphase sex chromosomes may well be caused by the more compacted nature of the chromatin in which fainter histone signals are not picked up. To conclude, these data indicate that in man a variation in meiotic gonosomal expression pattern is tolerated before, and likely also after the meiotic divisions.

Also in the human, the aggregation of unsynapsed autosomal chromatin with the sex chromosomes intervenes with spermatogenesis [31]. As in mouse this increases heterogeneity of human sperm even further and generally results in the oligo-astenoteratozoospermia (OAT) syndrome as well [46], suggesting that a link between X,Y gene activity and spermatogenic cell death/spermatid differentiation is maintained [24,47]. Transcriptional inactivation at the unsynapsed autosomal segments might be a contributing factor [47], which is omitted when studying mouse and human males with an XYY karyotype. In the mouse, Y activity during pachytene is detrimental as evidenced by strong selection during first prophase against spermatocytes with a Y bivalent that escapes MSCI [12]. This selection is less strong in the human XYY spermatocytes as is demonstrated in patients in which the habitual loss of one Y chromosome during the spermatogonial divisions did not occur and the Y chromosomes could partly or wholly synapse [48–51]. Loss could occur at the meiotic divisions or during spermiogenesis, and to a much lesser extend at pachytene. These observations support our interpretation that Y (and X) expression during first meiotic prophase in the human is less strictly controlled as compared to mouse, leading to the greater between gamete variation as observed in our species [37,38].

An evolutionary explanation for variable MSCI in the human

In human, the phenotypes of the AZFb and c deletions allow an indirect genetic analysis into Y gene activity during spermatogenesis. AZFb presents with a uniform early meiotic prophase maturation arrest [52] suggesting transcription from the Y being required before MSCI. In the AZFc deletion hypospermatogenesis is found, leading to ejaculated sperm in 81% of men [53,54] suggesting that absence of AZFc gene transcripts leads to variation in cell fate. Table S2 lists several genes from the male specific region of the Y (MSY), mostly of the AZFc, b and c regions, from which only scarce in situ gene expression data (RNA and protein) were available in the literature. Although gene specific RNA FISH is needed to confirm the pattern presented in this table, the suggestion is that some genes do display MSCI (DAZ, SRY, TSPY) while others do not (DDX3Y, RBMY, SMCY, RP5H1Y). DDX3Y and RBMY are also present at the mouse Y where they are subjected to MSCI [12].

When we compare the MSY gene content in human [55] and mouse [12], several more differences can be observed: first, the number of protein-coding genes is lower in mouse (15) [human: 27] and second, only 40% of the mouse Y genes are preserved on the human Y illustrating the high rate of evolution, despite the fact that the Y specialises in spermatogenesis in both species. The recent finding of remarkable differences in MSY gene content between human and chimpanzee stress the rate of evolutionary change even further [56]. These authors observed that compared with man, the chimpanzee Y chromosome had lost 4 out of 16 genes from the X degenerate regions that do not have a spermatogenesis specific function. Interestingly, a recent in silico analysis on array hybridisation expression data of whole testes from the chimpanzee and man revealed signs of MSCI for the chimpanzee but could not demonstrate decreased expression of human XY genes [26].

Among the evolutionary forces that shape the Y chromosome, as summarized by Hughes [56], there is first of all specialization of
gene function towards spermatogenesis, secondly the high frequency of intrachromosomal ectopic recombination, and thirdly the absence of homologous interchromosomal recombination (for the MSY region). Furthermore, sperm competition within the female reproductive tract likely speeds up selection and hence evolution of the Y chromosome [56]. From human population genetics [57–59] and physiological data [60] it can be deduced that in the human polygyny has not been the norm over our recent evolutionary history though it likely prevailed over earlier times. Hence, multiformal-multiformal mating systems are assumed to be rare [60]. This may have caused a relaxation of MSCI. This relaxation might also be demonstrated by the fact that some AZFb, c genes probably are subjected to MSCI and others not (Table S2). Thereby, systematic MSCI is alleviated to allow Y gene activity in pachytene and most likely the post meiotic stages. As a by product, the heterochromatinization of the meiotic sex chromosomes in late pachytene as witnessed in mouse is much less pronounced in man. Also, the selective pressure on morphological uniformity in gamete quality, that seems better guarded by complete MSCI and is here assumed to be functional in sperm competition between males, has been given up. Relaxation of MSCI might also involve a difference in activation of the pachytene checkpoint which in mouse was found to be induced by miss expression of Zfy1/2 [12]. When MSCI is less effective in human, human ZFY transcripts might not activate this checkpoint, as also is indicated by the occurrence of meiotic divisions in XYY men, who escape MSCI when a Y bivalent is formed [51]. However, of the ZFY/ZFX gene family which is present in both mouse and man, mouse Zfy1/2 are the least conserved, enabling evolution of function [61]. Alternatively, the human pachytene checkpoint might be placed before the time of relaxation of MSCI. However, this theory would not hold for XYY spermatocytes with a Y bivalent. Our observations need follow up by determining gonosomal gene expression in the spermatocytes and round spermatids, especially as also in the mouse, MSCI is not systematically executed: The majority of X-linked miRNAs (86%) escapes MSCI [62].

Conclusively, we postulate variation in meiotic and post-meiotic X,Y chromatin repression to contribute to morphological sperm heterogeneity, the consequences of which are alleviated by alternative selection mechanisms like within proband sperm selection and early embryonic death [63], which does not lead to a reproductive disadvantage for the couple.

**Materials and Methods**

**Ethics Statement**

On October 18th, 2006, the CCMO (Central Committee on Research involving Human Subjects) approved of the research protocol entitled 'Intracytoplasmatic Sperm Injection using testicular spermatooza in men with azoospermia: an observational study'.

**Human testis material**

Testis material was obtained from seven men (patients and controls) with non-obstructive azoospermia (NOA) or obstructive azoospermia (OA) willing to conceive, who underwent a testicular biopsy for sperm retrieval (TESE: testicular sperm extraction). One large biopsy was taken following the procedure of Silber [64]. From all probands a drop of spermatogenic cell suspension was smeared on a microscope slide prior to sperm retrieval. Cells were Giemsa stained and pachytene spermatocytes, spermatooza and Sertoli cells were counted. In Table S1, ratios between pachytene nuclei and mature elongated spermatids and Sertoli cells are given. The latter ratio indicates the spermatogenic activity of the tissue sampled, the former ratio the efficiency of the production of mature spermatids per meiotic cell. Although not strictly comparable with histological studies, our data are in line with those published [65–67]. Spermatogenic cells were available for research, after sperm retrieval had been successful. Patients were selected that at first inspection had relatively high numbers of spermatooza in the biopsy. The diagnosis of NOA was established after repeated absence of spermatooza in the ejaculate in combination with an elevated FSH level (>15 IU/ml) and no indications for an obstruction (Table S1). In cases of OA, repeatedly no spermatooza were found in combination with a medical history of obstruction and/or normal FSH levels (Table S1). Testis material from men of proven fertility, with a history of a vasectomy and absence of spermatooza in the caput epididymis thereafter (OA), was used as control material (C1/2/3 Table S1). Vasoscopytomy testis material is often used to represent normal spermatogenesis [25,68–70]. For control men 1 and 2, histology of the biopsy was evaluated by the Johnsen score [71]. Normal spermatogenesis was indicated by scores of 9.1 and 9.5 (with tubule scores of only 9 and 10). All men signed an informed consent for participation in this project. Testis material was used from one outbred NMRI and one inbred FVB mouse, both Swiss random bred derived. The procedure involving these animals is approved by the animal ethics committees of the Radboud University Nijmegen Medical Centre and Wageningen University and Research Centre, in conformance with the Dutch law on the use of experimental animals.

**Surface spread preparations**

Nucleus spreads were made as described by Peters [72], with minor modifications. Briefly, a suspension of spermatogenic cells was made by crushing the remnant seminiferous tubuli with two ribbed forceps in a drop of SIM (spermatocyte isolation medium [73]). Remnants were separated from the cell suspension by a quick spin (25 g). The supernatant was transferred to a clean tube, centrifuged for 7 minutes (159 g), and the pellet was resuspended in 1 ml SIM. An equal volume of a hypotonic solution (17 mM sodium citrate, 50 mM sucrose, 30 mM Tris-HCl pH 8.2) was added for 7 minutes. Cells were centrifuged again (7 minutes, 159 g) and resuspended in 100 mM sucrose (pH 8.2) at a concentration of 10–15 × 10⁶/ml. Two μl drops were pipetted onto a paraformaldehyde (1% PFA, 0.15% Triton-X-100 pH 7.4) fluid coated microscope slide, placed in a levelled humid box for about 75 minutes, rinsed twice in 0.08% photoflow (Koak), and air dried. Slides were stored at −80°C until use.

**Immunofluorescence (IF)**

Surface spread preparations were washed twice in PBS containing 0.05% Triton-X-100 and blocked for 1 hour at 37°C (blocking buffer: 1% bovine serum albumin, 10% normal goat serum or normal donkey serum in PBS containing 0.05% Triton-X-100). Primary antibodies were diluted in blocking buffer and slides were incubated for 20 minutes at 37°C, followed by overnight incubation at 4°C (when using the RNA polymerase II antibodies, we incubated 2 nights at 4°C) extended by 20 minutes at 37°C. Then slides were rinsed and washed once in PBS containing 0.05% Triton-X-100 and afterwards rinsed and washed once in PBS. A second 30 minute blocking step was applied in blocking buffer without Triton-X-100, followed by a 2 hour incubation with the secondary antibodies diluted in blocking buffer without Triton-X-100. After rinsing and washing once in PBS, nuclei were stained with DAPI (10 min, 0.3 μg/ml in PBS) and mounted with Vectashield (Vector). When performing
FISH after IF, nuclei were dehydrated in an 70–80–90–100% ethanol series after the last PBS wash.

DNA fluorescent in situ hybridization (FISH) with directly labelled probes

FISH analysis was performed with directly labelled whole chromosome X (LPP0X/R Aquarius, Cytocell) and Yq12 (32-131024 CEP Y (satellite III) Spectrum/Aqua, Abbott Molecular Inc.) probes. DNA was denatured for 5 minutes at 72°C and probes were able to hybridize overnight at 37°C. Slides were washed twice (1 minute) in 0.4× SSC/0.1% Tween-20 at 68°C and subsequently once in 2× SSC, once in 4× SSC and twice in PBS at room temperature. Nuclei were dehydrated in an ethanol series and mounted with Vectashield containing DAPI. For the DNA FISH protocol with bioin labelled Cot-1 probe see Methods S1.

Cot-1 RNA FISH

Cot-1 RNA FISH was performed as described [8] with modifications to enhance the signal (see Methods S1 and Figure S2A-H for rationales of this RNA FISH protocol). Briefly, human Cot-1 DNA (Invitrogen 15279-011) was labelled with bioin using the BioNick Labeling System (Invitrogen 18247-015). Probes were directly dissolved in hybridization buffer containing 50% formamide (Vysis). Probe was mixed (1:1) with preheated (37°C) hybridization buffer (4× SSC, 20% dextran sulphate, 2 mg/ml BSA, 2 mM vanadyl ribonucleoside), put onto a surface spread preparation and covered with a 18×18 mm coverslip. Nuclei and probe were heat denatured at 72°C (3 min) and hybridized overnight at 37°C in a humid box. The next day, slides were subjected to several stringency washes specific for RNA FISH starting with 3×5 minutes in 1× SSC/50% formamide at 42°C followed by 3×5 minutes in 2× SSC and subsequently placed in 4× SSC containing 0.1% Tween-20. Preparations were blocked (4× SSC, 0.1% Tween-20, 4 mg/ml BSA) for 30 minutes in a humid box at 37°C. Probe detection was carried out using avidine-FITC followed by goat anti-avidine FITC. Both reagents were incubated for 30 minutes at 37°C. Subsequently, slides were washed in PBS and directly used for primary antibody staining by IF. Primary antibody was diluted in PBS and incubated for 1 hour at 37°C followed by three PBS washes. Secondary antibody was incubated for 30 minutes at 37°C followed by 3 PBS washes. DAPI staining was performed and slides were mounted with Vectashield.

RNA imaging by 5-ethyluridine (EU) incorporation

Newly synthesized RNA was visualized by incorporation of the uridine analog 5-ethyluridine (EU) [74]. Incorporated EU can be detected by a copper-catalyzed cyclo addition reaction with a fluorescent azide (aka 'click reaction'). From a fresh biopsy (P4, Table S1) a cell suspension was made. After the first centrifuge step (7 minutes, 159 g) cells were resuspended in MEM alpha-medium [25]. Small cultures (200 μl) containing about 15×10⁶ cells/ml were made and EU (Invitrogen C10929) was added in a concentration of 4 mM. Cells were cultured for 3 hours at 32°C in an atmosphere containing 5% CO₂ [25]. Acid-fixed air-dried nuclei were prepared as described by Evans [75]. EU was detected using the Click-iT® RNA imaging kit (Invitrogen C10929) followed by DAPI staining and mounting in Vectashield.

Antibodies

To localize H3.1/3.2, monoclonal antibody #34 [76] provided by J. van der Vlag was used at a dilution of 1:1500 for IF, at 1:1000 before DNA FISH and at 1:500 after RNA FISH. A rabbit polyclonal antibody against SYCP3 (Abcam, ab15092) was used at 1:200 for IF and at 1:50 after Cot-1 RNA FISH. A rabbit polyclonal antibody against γH2AX [Upstate, 07-164] was used at 1:100. A polyclonal antibody against H3.3 phosphorylated at serine 31 (Abcam, 2009) was used at 1:100. A rabbit polyclonal antibody against H3K9me2, provided by T. Jenuwein was used at 1:150. To detect H3K9me3 two rabbit polyclonal antibodies were used, Abcam (ab8898) at 1:1000, and one generated by A.H.F.M. Peters at 1:750. Polyclonal rabbit antibodies against H3K27me2,3, provided by T. Jenuwein were used at 1:350. Two rabbit polyclonal antibody against H3K4me2 were used, Abcam (ab27671) at a dilution of 1:20 and a mouse monoclonal antibody (Abcam, ab817) at 1:50. Centromeres were detected by a human autoantibody (Crest) at a dilution of 1:500 (ImmunoVision, Springdale AR). Primary antibodies were detected by using goat or donkey anti-mouse and goat or donkey anti-rabbit secondaries with respectively a red or a green fluorochrome at 1:500 dilution (Invitrogen Alexa 488; A10001, A21292, Alexa 594; A11012, A21207). To detect the biotinylated Cot-1 RNA FISH probe, avidine FITC (Vector, SA-5001) was used in a 1:500 dilution enhancing the signal with a goat anti-avidine FITC antibody (Vector, SP-2040, 1:200).

Analysis and image capture

Staging of pachytene nuclei was based on staining of SYCP3 to visualize the synaptonemal complexes (SCs) [25], in combination with nuclear morphology by DAPI. To determine the degree of H3.1/3.2 nucleosome eviction, the level of Cot-1 RNA hybridisation and the level of EU incorporation of the late pachytene XY body, SYCP3 staining and/or its prominent and intense DAPI signal was used for localization and determination of its area. We subjectively determined the coverage of this area with H3.1/3.2, Cot-1 and EU signal. For H3.1/3.2 staining, late pachytene nuclei were categorized into four groups ranging from low nucleosome eviction to high nucleosome eviction (low > mid1 > mid2 > high) (Figure 1B). Cot-1 (Figure 5A-E) and EU RNA (Figure 6) staining was categorized into three groups (low > mid > high) in which ‘low’ indicates a strong decrease in signal and ‘high’ a modest decrease compared to autosomal staining (see Figures 5, 6). Categorization into three groups instead of four was carried out when fewer nuclei were included in the experiment. For the combination of H3.1/3.2 with Cot-1 RNA FISH, H3.1/3.2-stained late pachytene nuclei were categorized into three groups as well (see Figure 5 C,D,E). To determine the degree of H3.1/3.2 nucleosome eviction in round spermatids, nuclei were categorized into three groups: H3.1/3.2 was reduced on the X chromatin (detected by FISH) but signal was still present (low), H3.1/3.2 was strongly reduced and parts of the chromatin were free of H3.1/3.2 (mid), in a large part of the sex chromatin H3.1/3.2 was completely evicted (high) (Figure 7D).

Staining intensities and patterns of γH2AX, RNA polymerase II and the H3, H4 N-tail PTMs were subjectively determined as well and classified as described in the figure legends. Nuclei were captured by a Zeiss AxiosCam MR camera on a Zeiss Axiosplan fluorescence microscope using Axiosvision 3.1 software (Carl Zeiss). Around 30 late pachytene and 25 round spermatid nuclei were captured per patient/control man per immunostaining at an exposure time reflecting the microscopic image.

Statistics

Deviations from random class distributions have been tested by Chi square analysis.
Supporting Information

Figure S1 XY body localization. The degree of H3.1/3.2 nucleosome eviction of late pachytene spermatocytes from patients and controls (Table 1) is shown with their accompanying XY body localization. Bars indicate the location of the XY body: in or close to the centre (centre), halfway the centre and the edge (1/2), at two-third from the centre to the edge (2/3), and at the edge (edge). In grey the number of nuclei. Statistical analysis by Chi-square: overall $\chi^2$: 30.26 df 9 $p<0.001$, low vs. mid 2 $\chi^2$: 12.68 df 3, $p<0.01$, low vs. high $\chi^2$: 24.09 df 3, $p<0.001$, mid1 vs high $\chi^2$: 13.35 df 3, $p<0.01$, low + mid1 vs mid2 + high $\chi^2$:24.69 df 3, $p<0.001$.

Figure S2 Validation of RNA FISH signals at different denaturation temperatures. B–H The dotted circle indicates the XY body. A Lymphocyte metaphase. After DNA FISH the Cot-1 probe hybridizes to all chromosomes and showed more intense staining at the centromeric heterochromatin. B Pachytene spermatocyte nucleus. After DNA FISH the Cot-1 probe hybridized to all autosomal chromosomes and showed decreased hybridization signal for XY chromatin. C RNase-treated pachytene spermatocyte nucleus. After DNA FISH the Cot-1 probe hybridized to all chromosomes with more intense staining at the centromeric heterochromatin. D Pachytene spermatocyte nucleus. After RNA FISH at 72°C the Cot-1 probe hybridized to all autosomal chromosomes and showed decreased hybridization signals for XY chromatin. E RNase-treated pachytene spermatocyte nucleus. After DNA FISH at 62°C the Cot-1 probe was not able to hybridize. F–H Pachytene spermatocyte nucleus. After RNA FISH at 62°C, 52°C and 37°C respectively, the Cot-1 probe hybridized to all autosomal chromosomes and showed decreased hybridization signals for the XY chromatin.

Figure S3 Graphical representation of histone dynamics at the XY chromatin in late pachytene spermatocytes. A–E Late pachytene nuclei were assigned to four groups, ranging from a low to a high degree of nucleosome eviction (see Figure 1B) (for probands included and number of nuclei see Table 2). Histone characterstics and histone N-tail modifications were subjectively determined for the XY body and in A for autosomal chromatin as well: A Nuclei were categorized into three groups (bars). ‘Low staining, higher at XY’ indicates at the observation of low, even autosomal staining with more intense staining at the XY body. ‘General low staining’ indicates an overall low staining including the XY body. ‘Autosomal domains, strong XY’ points at the observation of a heterochromatin staining pattern on autosomal chromosomes and overall staining of the XY body. B The part of the XY chromatin stained was determined and arranged into four groups (bars). C–E The decrease in signal for the XY chromatin, compared to the autosomal chromatin, was determined and categorized into classes (bars). Statistical analysis by Chi-square: A: $\chi^2 = 21.90$ df 6, $p<0.01$, B: $\chi^2 = 47.87$ df 9, $p<0.001$ (Abcam antibody), C: $\chi^2 = 29.2$ df 6, $p<0.001$, D: $\chi^2 = 40.2$ df 9, $p<0.001$, E: $\chi^2 = 10.9$ df 6, $p = 0.21$ (Abcam antibody).

Table S1 Proband details.

Table S2 Literature data on meiotic expression of several MSY/AZFa,b,c genes.

Methods S1 Supporting methods.

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Author Contributions

Conceived and designed the experiments: PdB MdV LR. Performed the experiments: MdV SV GM. Analyzed the data: PdB MdV SV DGW. Contributed reagents/materials/analysis tools: GM KD LR MdV PdB. Wrote the paper: MdV PdB LR DGW.

References


