Transition from a meiotic to a somatic-like DNA damage response during the pachytene stage in mouse meiosis

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Abstract

Homologous recombination (HR) is the principal mechanism of DNA repair acting during meiosis and is fundamental for the segregation of chromosomes and the increase of genetic diversity. Nevertheless, non-homologous end joining (NHEJ) mechanisms can also act during meiosis, mainly in response to exogenously-induced DNA damage in late stages of first meiotic prophase. In order to better understand the relationship between these two repair pathways, we studied the response to DNA damage during male mouse meiosis after gamma radiation. We clearly discerned two types of responses immediately after treatment. From leptotene to early pachytene, exogenous damage triggered the massive presence of γH2AX throughout the nucleus, which was associated with DNA repair mediated by HR components (DMC1 and RAD51). This early pathway finished with the sequential removal of DMC1 and RAD51 and was no longer inducible at mid pachytene. However, from mid-pachytene to diplotene, γH2AX appeared as large discrete foci. This late repair pattern was mediated initially by NHEJ, involving Ku70 and XRCC4, which were constitutively present, and 53BP1, which appeared at sites of damage soon after irradiation. Nevertheless, 24 hours after irradiation, a HR pathway involving RAD51 but not DMC1 mostly replaced NHEJ. Additionally, we observed the occurrence of synaptonemal complex bridges between bivalents, most likely representing chromosome translocation events that may involve DMC1, RAD51 or 53BP1. Our results reinforce the idea that the early “meiotic” repair pathway that acts by default at the beginning of meiosis is replaced from mid-pachytene onwards by a “somatic-like” repair pattern. This shift might be important to resolve DNA damage (either endogenous or exogenous) that could not be repaired by the early meiotic mechanisms, for instance those in the sex chromosomes, which lack a homologous chromosome to repair with. This transition represents another layer of functional changes that occur in meiotic cells during mid pachytene, in addition to epigenetic reprogramming, reactivation of transcription, changes in the gene expression profile and acquisition of competence to proceed to metaphase.
DNA repair is critical for both somatic and meiotic cells. During meiosis, hundreds of DNA double strand breaks (DSBs) are introduced endogenously. To repair this damage, meiotic cells use a specialized version of the homologous recombination (HR) pathway that uses specific meiotic recombinases, such as DMC1, to promote repair with the homologous chromosome instead of the sister chromatid. This process is important to ensure chromosome segregation during meiosis and, as a side consequence, increases the genetic diversity of offspring. Nevertheless, under specific circumstances, meiotic cells can use other DNA repair mechanisms such as non-homologous end joining (NHEJ), which is error-prone. We investigated the response of mouse spermatocytes to increased DNA damage caused by gamma radiation, which is commonly used in cancer therapy. We found that the excess of DSBs produced by irradiation is processed by the meiotic HR recombination pathway in spermatocytes at the early stages of first meiotic prophase. However, this response is not inducible from the mid-pachytene stage onwards. From this point on, spermatocytes rely on a response that shares many features with that of somatic cells. In this response, the NHEJ pathway is first used to repair DNA damage but is subsequently replaced by a HR mechanism that does not use DMC1. Instead, it relies only on RAD51, which is known to function in both somatic and meiosis cells and, contrary to DMC1, has a preference for the sister chromatid. This switch from a meiotic to a somatic-like response is accompanied by a conspicuous change in the epigenetic response to DNA damage, reinforcing the idea that a functional transition occurs in meiotic cells during the mid-pachytene stage.

Introduction

DNA damage response is one of the most critical processes for cell survival and proliferation. Of the different forms of DNA damage, double-strand breaks (DSBs) are by far the most harmful. DSBs can arise spontaneously as a consequence of exposure to physical and chemical agents or following replication errors. In somatic cells, two main mechanisms, non-homologous end joining (NHEJ) and homologous recombination (HR) operate to repair DSBs [1]. NHEJ is the most common mechanism, working in all phases of the cell cycle [2] and its action is apparently simple. Classical NHEJ relies on the recruitment of the Ku70/80 complex and other regulatory factors, such as 53BP1, to the site of breaks to prevent DNA resection. This is followed by the incorporation of DNA-PK and DNA ligase IV, which reseals the break with the help of accessory factors such as XRCC4 [3,4]. In recent years, in addition to the classical NHEJ, a variety of alternative end-joining pathways, which use additional biochemical components, have been uncovered [4,5].

Although NHEJ pathways are quite efficient, they are also error-prone as they do not discriminate whether the two rejoined ends were the correct ones and resection and/or exonuclease activity may have removed sequences from the broken ends. In contrast, HR uses an intact DNA molecule as a template for repair, ensuring high fidelity of repair. However, this mechanism only acts when a DNA copy, usually the sister chromatid, is available, which only happens during the S/G2 phases of the cell cycle. The critical difference of HR repair versus NHEJ is that DNA is resected around the break point [4], generating single-stranded DNA (ssDNA) fragments that are used to search for the template sequence. The ATM kinase and the MRN protein complex (comprised of MRE11, RAD50 and NBS1) function as damage sensors by...
recognizing DSBs \([3,6,7]\). The MRN complex, together with other proteins (e.g. CtIP, BRCA1, BLM, EXO1, DNA2), then performs a 5’ to 3’ resection of DNA on either side of the break, which forms 3’-protruding ends of ssDNA \([4]\). The newly produced ssDNA is covered by RPA, which protects it from degradation \([3]\). Then, the ATR-ATRIP (Ataxia Telangiectasia and Rad3-related and ATR-Interacting Protein) complex binds directly to the RPA-coated ssDNA, thus localizing the kinase ATR to DSBs \([6]\). After DNA resection, the recombinase protein RAD51 replaces RPA and forms nucleoprotein filaments, allowing the ssDNA to invade the DNA double helix of the template DNA and further proceed with the repair of the DSB \([4,8]\).

HR may operate in both somatic and meiotic cells. During meiosis, homologous chromosomes undergo a series of complex processes, including pairing and synapsis, recombination and segregation. Meiotic recombination is in essence a HR repair mechanism that ensures the proper segregation of chromosomes during the first meiotic division and increases genetic diversity \([8,9]\). Although the molecular mechanisms mediating HR in somatic and meiotic cells are similar, there are several differences. The first one is the way DSBs are produced. In somatic cells, DSBs are usually generated by spontaneous events while in meiosis hundreds of DSBs are endogenously induced by SPO11 endonuclease during the leptotene stage of the first meiotic prophase \([9–11]\). Template choice for DSB repair is another important difference between somatic and meiotic HR. Sister chromatid is the common choice for DSB repair in somatic cells. However, in meiosis HR is tightly regulated to favor recombination with the homologous chromosome, although the sister chromatic can still be used. Repair with the homologous chromosome promotes crossing-over formation, which ensures coordinated chromosomal disjunction at the first meiotic anaphase \([11–14]\). Finally, during meiosis a specific recombinase, DMC1, is expressed in addition to RAD51. The coordinated action of DMC1, RAD51 and other regulatory factors drives repair to favor non-sister chromatid donors during meiosis \([15]\). In meiosis, DNA contacts between homologous chromosomes can ultimately resolve as reciprocal or non-reciprocal recombination events, which lead to crossovers or gene conversion events, respectively.

Although HR is the main DNA repair pathway acting during meiosis, NHEJ can also be used \([16]\). Components of the classical NHEJ pathway such as Ku70/80 and 53BP1 have been detected in mouse meiotic cells, both in the course of normal meiosis \([17]\) and after the exogenous induction of DNA damage \([16,18]\). Not surprisingly, this mechanism seems to be triggered only in the late stages of first meiotic prophase. This may be a consequence of the upregulation of HR repair during the early stages of meiosis following the endogenous production of DSBs by SPO11 and the resection of DNA that is concomitant with SPO11 removal \([19]\). However, coexistence of HR and NHEJ is possible during the late stages of meiotic prophase to repair DNA damage that was induced by either endogenous or exogenous mechanisms. Radiation exposure experiments, for instance, have reported an increase of both 53BP1 and RAD51 levels in pachytene and diplotene spermatocytes \([16,18,20,21]\).

As mentioned above, the key event for the choice between HR and NHEJ relies on the resection of DNA around the break \([2]\). The production of ssDNA overhangs hampers the action of NHEJ mechanisms, which require intact ends. Although the regulation of DNA resection at DSBs is not completely clear, a reciprocal regulation of factors promoting and inhibiting resection has been reported \([22]\). 53BP1, which plays several roles in the regulation of DNA repair \([2]\), has been proposed to play a key function in inhibiting resection by hampering the loading of the CtIP-BRCA1 complex to the DNA, thus diverting repair to the NHEJ pathway \([4,23,24]\). CtIP-BRCA1, in turn, is thought to negatively regulate 53BP1 by inducing displacement of both 53BP1 and Ku70/80 from the break point and stimulating DNA resection by the MRN-(EXO1-DNA2-BLM) complex \([2,25]\). Interestingly, both 53BP1 and BRCA1
seem to rely on ATM kinase for phosphorylation, which is necessary for their function. Additional factors, such as the action of specific CDK-cyclin complexes and the epigenetic landscape around the break point also contribute to the regulation of DNA end resection [22].

In addition to the biochemical interactions described above, the morphological, temporal and epigenetic scenario in which DNA repair occurs during meiosis must be considered. Synapsis, the intimate association of homologues, is mediated by a highly specialized structure called the synaptonemal complex (SC) [26]. Assembly and disassembly of SC components during the first meiotic prophase is a tightly regulated process crucial for proper chromosome recombination and segregation [27], as evidenced by the number of synopsis mutants in which recombination is disturbed, and vice versa [28–32]. Furthermore, during first meiotic prophase, the complex regulation of transcription and chromatin modifications can influence the response to DNA damage [33–35]. Most conspicuously, histone H2AX is phosphorylated to give rise to γH2AX, which localizes throughout the chromatin during the leptotene stage in response to DSBs [36]. This contrasts with the pattern of γH2AX in somatic cells, where it usually forms small and discrete foci after DNA damage [37]. γH2AX is involved in recruiting many DNA repair factors [4,10,36,38,39] and in the transcriptional silencing that is characteristic of the beginning of meiosis and the sex chromosomes [34,36,40]. Notably, ATM, ATR and DNA-PK can all phosphorylate H2AX [4,41,42]; therefore, γH2AX is a marker of both the HR and NHEJ pathways. Upon DNA repair, γH2AX seems to be displaced from the chromatin and/or dephosphorylated by protein phosphatases [43–45].

To shed light on the complex relationships between HR and NHEJ repair mechanisms acting during meiosis, we assessed DNA repair responses during mammalian male meiosis after the exogenous production of DSBs. We irradiated mice with gamma rays and then analyzed the localization and dynamics of various markers of DNA repair response, including γH2AX, DMC1, RAD51, 53BP1, Ku70 and XRCC4, at different times of recovery. We have uncovered two distinct epigenetic patterns in response to DNA damage in early and late prophase-I spermatocytes: a typical meiotic one and a somatic-like one acting at early and late stages, respectively. The transition to a somatic-like response during mid-pachytene coincides with the sequential cessation of the meiotic HR response at mid-pachytene and the consecutive activation of NHEJ and somatic-like HR repair mechanisms. In addition, we report the formation of chromosome bridges between non-homologous chromosomes associated with either HR or NHEJ markers.

Results

γH2AX dynamics after irradiation

We firstly analyzed the distribution pattern of γH2AX (H2AX phosphorylated at serine 139) in response to DNA damage. Phosphorylation of this histone is one of the first cytological events detected after DNA damage and has been used extensively to localize DSBs in both somatic and meiotic cells [4,10,36,37,39]. Staging of spermatocytes during first meiotic prophase was made on the basis of the degree of chromosome synapsis between autosomes, the morphology of SC and the morphology of the sex chromosomes following SYCP3 immunolabeling, as previously characterized [34].

In control spermatocytes, γH2AX is first detectable at the beginning of leptotene, when short threads of SYCP3 mark the initial assembly of axial elements (AEs) along the chromosomes. At this early stage, only a few discrete γH2AX foci are observed scattered throughout the nucleus (Fig 1A). During mid to late leptotene, when AEs form longer filaments, γH2AX is broadly localized throughout most of the nucleus (Fig 1B). This broad nuclear distribution is maintained during early zygotene (Fig 1C), when AEs start to synapse. From mid-zygotene
Fig 1. Pattern of γH2AX after irradiation in early prophase mouse spermatocytes. SYCP3 (green) and γH2AX (red) at different stages of prophase I by recovery time after irradiation. (A-E) Control. (A) Early leptotene. γH2AX localizes as small scattered foci over the short threads of forming AEs. (B) Mid-late leptotene. AEs are more extended and now a massive γH2AX signal covers the nucleus. This pattern is also found in early zygotene (C), when AEs are completely formed and homologues start to synapse. (D) Late zygotene. Homologous chromosomes have
onwards, γH2AX signal decreases and, by the end of zygotene, is mainly associated with unsynapsed regions (Fig 1D). During pachytene, when homologous chromosomes are fully synapsed, γH2AX localizes almost exclusively on the sex chromosomes, which have extensive unsynapsed regions (Figs 1E, 2A and 2B). Nevertheless, large γH2AX foci are sometimes observed associated to the SCs of some autosomal bivalents (Fig 2B, arrowhead). These foci have been previously described [10,34] and interpreted as unrepaired DSBs that tend to disappear with pachytene progression or, alternatively, as regions of transcriptional silencing [46]. During diplotene, when homologues desynapse, γH2AX remains present only on the sex chromosomes (Fig 2C).

In gamma-irradiated spermatocytes, visible changes in the pattern of γH2AX localization are observed one hour after irradiation. In early leptotene cells, γH2AX is seen throughout the nucleus, in contrast to the small scattered foci seen in controls, indicative of a massive broadly distributed DNA repair response (Fig 1F). This pattern is also observed in late leptotene, zygotene and early pachytene spermatocytes (Fig 1G–1J). Changes at late leptotene and zygotene stages are less evident as γH2AX is already broadly localized throughout the nucleus in control cells at these stages. This pattern indicates that cells at the beginning of meiosis up to early pachytene respond to the induction of DNA damage similarly. In contrast, the response of spermatocytes from mid-pachytene onwards is rather localized. Large γH2AX foci are observed emerging from the SCs (Fig 2D–2F). This kind of signals have been called large foci [10], flares [46] or eruptions [47] and their morphology resembles that found in control spermatocytes (Fig 2B) and somatic cells [37]. These results reveal the existence of morphological differences in the response to DNA damage between early and late meiotic prophase spermatocytes.

Irradiated spermatocytes show a clear diminution of γH2AX in most stages 24 hours after treatment. Similar to control cells, early leptotene cells have a few scattered γH2AX foci (Fig 1K). Provided that meiotic progression is not greatly affected by irradiation, then cells should progress to further stages during the recovery time. In order to test this possibility, we followed the progression of meiosis in control and irradiated mice after incorporation of EdU. Comparison spermatocyte population progression at different time points (24 and 72 hours) show that the advance of meiosis is not affected after irradiation (S1 Fig and S1 Table), in agreement with previous reports [48]. Therefore, early leptotene cells at 24 hours post treatment could have been at pre-leptotene when irradiated (see S1 Fig for an estimation of the length of each meiotic stage). In late leptotene and early-mid zygotene spermatocytes, γH2AX is distributed throughout the nucleus, similar to control cells (Fig 1L and 1M). Likewise, the pattern of γH2AX at late zygotene and early pachytene is comparable to that of the controls (Fig 1N and 1O), in which γH2AX appears to label the unsynapsed chromosomal regions and some foci in a few chromosomes. These cells were likely irradiated at leptotene and zygotene stages, respectively, indicating that cells irradiated at early meiotic stages are able to achieve a control pattern corresponding to their stage 24 hours after irradiation. In contrast, cells from mid-pachytene to diplotene retain several foci associated with SCs (Fig 2G–2I). These differences are also observed 72 hours after irradiation (Figs 1P–1T and 2J–2L). In this case though, cells at mid-
Fig 2. Pattern of γH2AX after irradiation in late prophase mouse spermatocytes. SYCP3 (green) and γH2AX (red) at different stages of prophase-I by recovery time after irradiation. (A–C) Control. γH2AX appears over the sex chromosomes (X and Y) in mid-pachytene (A), late pachytene (B) and diplotene (C). Occasionally, some small foci remain associated with autosomes (arrowheads). (D–F) 1 hour of recovery. In addition to the sex chromosomes, γH2AX localizes on the autosomes as large foci that emerge from the SCs (arrowheads) at all three stages. (G–I) 24 hours of recovery. All stages show a visible decrease in the amount of γH2AX. In mid-pachytene cells (G), γH2AX foci are almost absent yet many foci are still present in late pachytene (H) and diplotene (I) cells. (J–L) 72 hours of recovery. The pattern is similar to 24 hours; some foci (arrowheads) remain present in late pachytene (K) and diplotene (L) cells. Some chromosomal connections are visible and appear to involve γH2AX signals (arrows). (M) Dotplot of the number of γH2AX foci in spermatocytes grouped by recovery time. The increase in the number of foci is evident 1 hour after irradiation. ANOVA analysis showed no statistical differences at this time between the three stages analyzed (p = 0.22). Tukey’s multiple comparisons test for individual comparisons between different stages showed no statistical differences. 24 hours after irradiation, a reduction in the number of foci is observed at all stages, but now statistical differences between stages are observed (ANOVA p < 0.0001). Individual comparisons indicate the existence of differences between all stages. An analogous situation is found 72 hours after irradiation (ANOVA p < 0.0001). (N) Dotplot of the number of γH2AX foci in spermatocytes grouped by stage. While mid-pachytene cells...
pachytene 72 hours after irradiation were at an earlier pachytene stage at the time of irradiation. These cells likely had widespread localization of γH2AX at an earlier stage in response to DNA damage but their γH2AX pattern changes as they progress, very much like under endogenous production of DSBs. The persistence of γH2AX foci, however, indicates incomplete DNA repair.

Notably, leptotene cells were very scarce 72 hours after irradiation (see S1 Fig). Previous reports indicated that spermatogonia are particularly sensitive to radiation [18,20,49,50]. In order to confirm apoptosis of these cells, we performed a TUNEL assay on testicular sections (S2 Fig) and observed an increase of apoptosis in specific cell populations at different recovery times. Specifically, 24 hours post irradiation, a noticeable, but not massive, increase of apoptosis is observed in spermatogonia and prophase-I spermatocytes, while at 72 hours apoptosis is mainly observed in metaphase cells. This leads us to infer that irradiation may partially ablate spermatogonia population, but probably also interrupts the normal entrance of these cells in meiosis, which explains the scarcity of leptotene cells.

The two patterns of response to DSBs, early and late, also appear to differ in terms of γH2AX removal. Spermatocytes irradiated at late pachytene or diplotene, or those that reach these stages during recovery, remove γH2AX more slowly than those irradiated at earlier stages. In order to ascertain the efficiency of DNA repair, we recorded the number of γH2AX foci from mid-pachytene to diplotene (S1 Table) and analyzed the progression of repair by recovery time (Fig 2M) and cell stage (Fig 2N). One hour after irradiation, the number of foci increases in the three stages. The ANOVA test showed no significant differences between stages at this time. However, at 24 hours, the number of foci returns to control levels in mid-pachytene spermatocytes. In contrast, late pachytene and diplotene spermatocytes still show an increased number of foci, which is maintained even 72 hours after treatment. These results support the idea that γH2AX removal is less efficient as cells progress to later stages of prophase-I and that the number of foci seems to reach a steady state with no significant reduction.

Formation of chromosomal bridges

One striking feature observed after irradiation is the formation of connections between non-homologous chromosomes, which can be visualized by SYCP3 immunostaining (Figs 1T, 2J and 3). Connections are observed at all post-treatment times (1, 24 and 72 hours) and could be clearly identified in zygotene to diplotene spermatocytes. On the basis of their morphological appearance, we classified connections in three categories (Fig 3): 1) distal contacts, in which chromosomes interact end-to-end (Fig 3A and 3B); 2) interstitial contacts, in which a filament emerges from one bivalent and contacts one or more bivalents laterally (Fig 3C–3F) and 3) intrachromosomal contacts, in which the connection is observed within the same bivalent (Fig 3G–3I). In some cases, the SYCP3-positive filament of a bivalent seems to split into two with a thin filament, probably involving a single chromatid, providing the connection (Fig 3C). In other cases, the filament appears thicker (Fig 3D).

Chromosome connections can be observed between autosomal bivalents, between autosomes and sex chromosomes or between sex chromosomes. The presence of these bridges is likely not an artifact of the spreading technique as they are also observed in squashed spermatocytes (Fig 3I). Furthermore, chromosome fragments and bridges are observed during anaphase- and telophase-I (Fig 3K and 3L), indicating that these connections may represent chromosomal translocations. While connections between bivalents can result in a non-
Fig 3. Types of chromosomal bridges found after irradiation. SYCP3 protein in green. (A–I) Spread spermatocytes at pachytene, except (G), which is at zygotene. (A) Distal junction between two autosomes. (B) Distal junction between an autosome and a sex chromosome, in this case the X. (C–D) Interstitial junctions between autosomal bivalents. In (C) a bivalent with two bridges, each contacting a different bivalent, is shown. In the inset, a higher power view of one of the bridges is shown. The lateral element of the homologue involved in the bridge is split into two filaments. One filament remains associated with the homologous chromosome and the other is linked to the chromosome of the other bivalent. In (D) two bivalents are sharing a bridge. In this case, the bridge is a whole counterpart, which has invaded the other bivalent. A
homologous chromosomal translocation, bridges within bivalents can potentially link the two homologous chromosomes or different parts of the same chromosome. The presence of these chromosomal aberrations at metaphase-I, which are rarely detected in the control cells, might account for the increased apoptosis observed at this stage 24 and 72 hours after treatment (S2 Fig).

In order to understand the dynamics of chromosome bridge formation, we quantified the number of cells showing at least one of these chromosomal connections during pachytene and diplotene (Fig 3M) (connections were more difficult to discern from chromosome tangles in earlier stages). No bridges were found in 316 control cells analyzed. However, after irradiation, the frequency of spermatocytes bearing bridges increases from 6.48% at one hour to 9.15% at 24 hours and 24.13% at 72 hours, indicating a clear rise in the number of bridges with time. Regarding the distribution of bridges by stage and time, at 24 hours, most of the cells with bridges are at early pachytene; however, by 72 hours, the distribution is more uniform among stages.

**Localization pattern and dynamics of DMC1**

In order to investigate the action of HR mechanisms, we first examined the spatial and temporal localization pattern of DMC1, which is exclusively present in meiosis and acts together with RAD51 [51,52]. To compare DMC1 distribution with the γH2AX pattern just described, we performed triple immunostaining of SYCP3, DMC1 and γH2AX. In control spermatocytes, a few DMC1 foci are seen scattered throughout the nucleus at early leptotene (Fig 4A). These foci are not specifically associated with either the short SYCP3 fragments or the small γH2AX foci already present. The presence of DMC1 foci at the beginning of leptotene suggested that they might be responding to DSBs produced by a SPO11-independent mechanism. However, their absence in SPO11 null mutants (S3 Fig) rules out this possibility. During late leptotene (Fig 4B) and early zygotene (Fig 4C), many more DMC1 foci are observed. At late zygotene, the number of DMC1 foci decreases (Fig 4D). Some foci remain associated with autosomes but they are clearly more abundant on the unsynapsed AE of the X chromosome. During early pachytene (Fig 4E), fewer foci are visible. Although DMC1 and γH2AX are co-localized on some autosomes, in many cases, DMC1 and γH2AX foci are not associated with one another (see detail in Fig 4E). At mid-pachytene, the number of autosomal DMC1 foci still decreases, though the sex chromosomes still have a high number of foci (Fig 4F). DMC1 is no longer detectable at a cytological level after mid-pachytene.

After irradiation, a notable increase in DMC1 foci is observed (Fig 4G–4Z). As in control mice, these foci appear associated with unsynapsed AEs during leptotene, with synapsed and unsynapsed regions during zygotene and with synapsed autosomes and the AE of the X chromosome from pachytene onwards. Interestingly, DMC1 is not detected beyond mid-pachytene, indicating that radiation exposure is not able to trigger the appearance of DMC1 foci after this stage. Similar to control cells, some co-localization of DMC1 and γH2AX is observed in irradiated pachytene cells (see details in Fig 4K and 4L). We also observed DMC1-positive filaments connecting different chromosomes (Fig 4N, 4W and 4Z). These filaments are mainly...
Fig 4. Pattern of DMC1 at different stages of prophase-I by recovery time after irradiation. SYCP3 (red), γH2AX (blue) and DMC1 (green). (A-F) Control. (A) Early leptotene. A few small foci of DMC1 appear distributed throughout the nucleus; however, they do not seem to specifically co-localize with SYCP3 or γH2AX. (B) Mid-late leptotene. DMC1 foci are more abundant and now mostly associated with short SYCP3 filaments. (C) Early-mid zygotene. DMC1 foci are very abundant over the formed AEs, some of which are undergoing synopsis. (D) Late zygotene. DMC1 foci are located over both synapsed and unsynapsed chromosomes. Some signal co-localizes with remaining clouds of γH2AX, while others do not. The X chromosome (X) appears coated with many foci, while only a single focus is seen on the Y chromosome (Y). Gradually, DMC1 foci disappear during early pachytene (E) and mid-pachytene (F), but remain on the sex chromosomes and some autosomes. Only occasionally do some of these DMC1 foci co-localize with γH2AX (see enlarged detail of the bivalent indicated by an arrow in E). (G-L) 1 hour of recovery. The number of DMC1 foci increases at all stages from leptotene to early pachytene. At early leptotene (G), DMC1 coincides with the increase and spread of γH2AX to the whole nucleus. The number of DMC1 foci is clearly higher than in the control shown in A. No conspicuous differences in the pattern of DMC1 are observed at late leptotene (H) or zygotene (I-J). The Y chromosome still shows a single DMC1 focus. In early (K) and mid-pachytene (L) spermatocytes, DMC1 is observed on autosomes and sex chromosomes. Again, DMC1 foci may co-localize or not with γH2AX. Enlarged views of some bivalents (arrows) are shown as insets in panels K and L. Chromosomal bridges are also found (arrowheads). (M-R) 24 hours of recovery. The distribution of γH2AX resembles that of control cells at all stages but the number of DMC1 foci seems reduced compared with cells 1 hour after irradiation. In some cells DMC1 appears to form filaments, sometimes joining AEs together (see arrowheads and detail in N). (S-Z) 72 hours of recovery. Leptotene cells (S-T) are found at a very low frequency and usually include morphological distortions. The morphological features of cells from zygotene to mid-pachytene are similar to those found at 24 hours. Again, small DMC1 filaments appear on the chromosomes. These filaments seem to occasionally mediate the formation of bridges between two bivalents (arrowheads and details in W and Z); in some cases, γH2AX signal is associated with bridges (Z). A’–F’) Dotplot representation of DMC1 foci distribution by cell stage. ANOVA analysis showed that the number of DMC1 foci increased at all stages after irradiation (p<0.0001) except mid-pachytene (p = 0.93). The increase of DMC1 foci observed 1 hour after irradiation compared to control is lower as cells are in more advanced stages, and no increase is found at mid pachytene. Likewise, differences in the number of foci between 1 and 24 hours, or between 24 and 72 hours, become less or not significant as cells are in later stages. Nevertheless, control levels in terms of number of foci were not observed for any of the stages, even after 72 hours of recovery, except obviously mid pachytene. Tukey’s multiple comparisons test for individual comparisons (ns: non-significant; *: p<0.05; **: p<0.01; ***: p<0.001; ****: p<0.0001).

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Our quantitative analysis revealed some interesting features (Fig 4A’–4F’, S4 Fig and S1 Table). First, the early leptotene cell population of control spermatocytes has a low number of DMC1 foci and very low standard deviation. As described above, this population is also characterized by a few small γH2AX foci. In contrast, mid-late leptotene cells show an increase in the number and standard deviation of DMC1 foci, in agreement with a previous report [53]. This stage is also associated with broad γH2AX labeling, as pointed above. Peak abundance of DMC1 foci occurs during early-mid zygotene and decreases thereafter. According to the ANOVA and Tukey’s multiple comparisons tests, differences between each stage and the next one are significant (S4 Fig), indicating that DMC1 distribution can be used to distinguish the cell populations of the six substages.

Second, as expected, the number of foci increases one hour after irradiation in most phases. As in the control, peak abundance of DMC1 foci is observed in early-mid zygotene spermatocytes, and each stage differs significantly from the following one, excepting mid-late leptotene and early-mid zygotene. However, the number of DMC1 foci induced by irradiation differs greatly among the different meiotic stages. The increase of foci compared to control is on
average 79, 125, 73, 44, 14 and 1 for each of the six substages, respectively (see S1 Table). This striking result indicates that the cell stages are not equally sensitive to irradiation or that DMC1 localization to DSBs may be differentially regulated at the different stages due to the availability of this protein or other DNA repair factors. Furthermore, in irradiated mid-pachytene spermatocytes, the number of DMC1 foci did not increase significantly regardless of recovery time, indicating that DMC1 is no longer inducible at this or later stages. These results can be easily discerned when data are grouped by cell stage (Fig 4A’–4F’).

Third, after the increase of DMC1 foci immediately after irradiation, a slow diminution is observed with recovery time for most stages; however, most did not reach control levels even after 72 hours of recovery time (Fig 4A’–4F’). Nevertheless, we observed two main stage-specific features: 1) early leptotene cells show almost control levels 24 hours later. Moreover, while the number of DMC1 foci is quite variable one hour after treatment, 24 hours later, the range of foci narrows, similar to the control. This finding may reflect the presence of newly formed leptotene cells that had just entered meiosis. Unfortunately, we could not record the number of DMC1 foci in early leptotene spermatocytes 72 hours after irradiation owing to the scarcity of this stage; and 2) in early pachytene, the number of DMC1 foci does not decrease but maintains constant in time. Since irradiation does not greatly disrupt meiotic progression, cells irradiated at a particular stage would continue to advance through meiosis and be at later stages when observed 24 or 72 hours later. Therefore, we arranged the quantitative data following a putative duration of 24 hours for leptotene, zygotene and early pachytene [35,50,54] (S1 and S4 Figs). This means that a cell irradiated at early leptotene would be at early zygotene 24 hours later and at mid-pachytene 72 hours later, and so on (S4 Fig). Considering four initial cell populations (early leptotene, late leptotene, early zygotene and late zygotene), we observed that, one hour after irradiation, the number of DMC1 foci increases in all cases and, in most cases, decreases 24 and 72 hours later, indicating efficient DNA repair in all cell populations. Control levels of DMC1 foci are reached after 72 hours of recovery, since all these cell populations are expected to reach mid-pachytene within the 72-hour period following irradiation. This suggests that DNA repair has been successfully completed in all cells. Alternatively, DMC1 might have been released from chromosomes at mid-pachytene, regardless of whether repair had been completed or not.

**Localization pattern and dynamics of RAD51**

We then analyzed the distribution of RAD51, which acts with DMC1 in the HR pathway, in control and irradiated spermatocytes. In agreement with previous reports [51,52], we found that RAD51 has a similar, albeit not identical, distribution pattern as DMC1 during first meiotic prophase (Fig 5). During zygotene stage (Fig 5A, S5 and S6 Figs), RAD51 localizes to the AEs of chromosomes, with a peak number of foci observed mainly in early-mid zygotene, decreasing continuously thereafter. At early pachytene and later stages, RAD51 foci remain associated with some autosomal SCs but are mainly found on the unsynapsed AE of the X chromosome (Fig 5B and 5C). Most of these foci are not associated with γH2AX, which at this stage is restricted to the sex chromosomes and a few foci over the autosomes. RAD51 disappears during late pachytene (Fig 5D) and is absent at diplotene (Fig 5E). This pattern is very similar to that of DMC1; however, we observed that RAD51 remains associated with chromosomes up to later pachytene stages. In order to observe this more clearly, we performed double immunostaining for both proteins (S5 and S6 Figs). During early stages of prophase-I, the localization of both proteins is almost, but not completely, identical. We noticed that not all DMC1 foci are associated with RAD51 foci and vice versa. More importantly, these two proteins are removed from chromosomes sequentially since the number of RAD51 foci on both
Fig 5. Pattern of RAD51 at different stages of prophase-I by different recovery time after irradiation. SYCP3 (red), γH2AX (blue) and RAD51 (green). (A-F) Control. (A) Late Zygotene. RAD51 foci are found in the non-synaptic AEs, which are also labeled with γH2AX, some synapsed autosomes and the sex chromosomes (X and Y). During early (B) and mid (C) pachytene, fewer RAD51 foci are observed. Some remain on the autosomes, but most are in the non-synapsed region of the X chromosome. RAD51 does not appear during the late pachytene
autosomes and sex chromosomes clearly exceeds that of DMC1 at the mid to late pachytene transition (S5E–S5E’ Fig). Thus, while the recruitment of RAD51 and DMC1 can be simultaneous upon the production of DSBs at the beginning of meiosis, persistent DSBs at the last stages of repair may lose DMC1 but maintain RAD51, which may reflect a role in promoting inter-sister versus inter-homolog interactions for the repair of DSBs [15]. Additionally, this persistence could be explained by the late production of DNA damage, which would not incorporate DMC1.

Similar to the results with DMC1, we observed an increase in the number of RAD51 foci after irradiation (Fig 5F–5T). Although we did not quantify the distribution of RAD51 at early meiotic stages, the broad co-localization of RAD51 and DMC1 up to mid-pachytene (S5 and S6 Figs) suggests that both proteins follow a very similar pattern, i.e., peaking one hour after treatment then decreasing with recovery time. This pattern agrees with that reported in previous studies [18,20]. However, the localization patterns of these proteins are not identical. For instance, though RAD51-positive filaments bridging chromosomes are also observed (S6 Fig), they are thinner and scarcer than DMC1 ones.

Strikingly, after irradiation, RAD51 is observed in late pachytene and diplotene spermatocytes (Fig 5I, 5J, 5N, 5O, 5S and 5T). Given that RAD51 is not observed at these stages in control spermatocytes, these foci must represent newly localized protein induced after irradiation. Indeed, this RAD51 population differs with the one observed at earlier stages. First, the signal strength of RAD51 on the sex chromosomes is similar to that of autosomes. Second, foci tend to be larger and sometimes irregularly shaped. Finally, while virtually all RAD51 foci observed at early stages (up to mid-pachytene) are associated with the AEs or SCs, during late prophase, a significant proportion of RAD51 is detached from the SCs (Fig 5J, 5N, 5O). Most of these foci co-localize with γH2AX one hour after irradiation, indicating they correspond to regions of DNA damage (Fig 5J and 5O).

To examine the dynamics of this late-appearing population of RAD51, we scored the number of foci present in mid-pachytene to late diplotene spermatocytes (Fig 6A and 6B; S1 Table). This analysis uncovered some interesting features. First, one hour after irradiation, RAD51 increases significantly in all stages analyzed except mid-pachytene. This striking result parallels the behavior of DMC1, which is also not inducible at mid-pachytene, suggesting that HR repair can be compromised at this stage immediately after irradiation. The increase of RAD51 observed in late pachytene, early and late diplotene stages is quite similar. Second, after 24 hours of recovery, the number of RAD51 foci is significantly higher in all cell populations, though the increase is more pronounced in late pachytene cells and conspicuously lower in diplotene cells. After 72 hours of recovery, RAD51 levels remain high, though a slight decrease is observed in all stages, except mid-pachytene. The unexpected behavior of RAD51 during mid-late pachytene and diplotene stages suggests that the HR response to induction of exogenous DSBs initially may be absent or weak but increases with time, at least until 24 hours after irradiation.

We were intrigued by the presence of RAD51 foci that were not associated with SCs. We analyzed the dynamics of RAD51 foci during diplotene (Fig 6C and S1 Table) and found that
Fig 6. Dotplot representation of RAD51 foci distribution. (A) Analysis of RAD51 distribution by recovery time. Four substages were considered (MP: mid-pachytene; LP: late pachytene; ED: early diplotene; LD: late diplotene). ANOVA analysis showed statistical differences ($p<0.0001$) for the control and the three recovery times. In the control, MP cells have a high number of RAD51 foci but later-staged cells have little to none. A similar increase in the number of RAD51 foci is observed from LP to LD 1 hour after irradiation. Tukey’s multiple comparisons test for individual comparisons showed statistical differences between MP and the rest of the stages, and also between LP and LD. Twenty-four hour after irradiation, the increase in the number of RAD51 is more obvious at all stages, while 72 hours after irradiation, the number of foci decreases from LP to LD. (B) Analysis of RAD51 distribution by cell stage. The number of RAD51 foci increases significantly in cells at all stages after irradiation ($p<0.0001$). However, according to Tukey’s test, the number of foci in irradiated mid-pachytene cells 1 hour after irradiation is not significantly different from control cells. At 24 hours, the number of foci increases in mid-pachytene cells and remains stable at 72 hours. This distribution departs from the pattern observed in cells at other stages, in which RAD51 increases slightly at 1 hour, peaks at 24 hours and then decreases at 72 hours. (ns: non-significant; *: $p<0.05$; **: $p<0.01$; ***: $p<0.001$; ****: $p<0.0001$). (C) Analysis of RAD51 foci associated (ON) or not associated (OFF) with SCs at early (blue) and late diplotene (brown). The distribution of both kinds of foci is similar, increasing at 1 hour, peaking at 24 hours and decreasing at 72 hours. Notably, the proportion of foci not associated with SCs (OFF) is higher in late diplotene spermatocytes.

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both SC-associated and non-associated RAD51 foci follow the same pattern, increasing one hour after irradiation, peaking 24 hours later and decreasing thereafter. We found that the number of foci associated with SCs is clearly higher in cells at early and late diplotene but that the proportion of non-associated RAD51 foci increases in late diplotene cells 24 and 72 hours after irradiation.

Localization of NHEJ markers

In order to ascertain the action of the NHEJ repair mechanism, we studied the temporal and spatial localization of XRCC4 and Ku70 components of this pathway. We first examined the localization XRCC4 (Fig 7), which is a ligase-IV co-factor. No protein is observed at a cytological level during early meiotic stages in control spermatocytes (Fig 7A). At late pachytene, however, a weak signal appears throughout the nucleus (Fig 7B) and becomes more intense at diplotene (Fig 7C). At this stage, the signal appears slightly more intense over the sex chromosomes. In order to rule out the absence of XRCC4 labeling in early spermatocytes as an artifact of the spreading technique, we also immunostained testicular sections (S7 Fig). XRCC4 is

![Fig 7. Distribution of NHEJ markers at different stages of prophase-I by recovery time after irradiation. SYCP3 (green) and XRCC4 (red) in late prophase-I spermatocytes. (A-C) Control. XRCC4 is absent up to mid-pachytene (A). At late pachytene (B), a faint signal is observed in the nucleus, which becomes more intense at diplotene (C). The signal appears more concentrated on the sex chromosomes (XY). (D-F) 1 hour, (G-I) 24 hours and (J-L) 72 hours after irradiation. The localization pattern of XRCC4 at each stage is almost identical. Foci do not form at any stage or recovery time.](https://doi.org/10.1371/journal.pgen.1007439.g007)
absent in spermatogonia and basal spermatocytes of the seminiferous tubules and is only detectable in spermatocytes located in the middle of the epithelium, corresponding to late pachytene-diplotene cells. We also studied the localization of Ku70, which is involved in the protection of broken DNA ends. Immunostaining of this protein yielded nearly identical results to XRCC4 (S8 Fig). These results indicate that these components are present by default during the normal course of meiosis, in agreement with previous reports [16,17]. We observed a very similar pattern in irradiated spermatocytes: no signal is detected prior to late pachytene and, from this stage onwards, the proteins are distributed homogeneously throughout the nucleus (Fig 7D–7L; S8 Fig). No marked differences in the signal strengths of these proteins were observed between control and irradiated cells. Moreover, neither XRCC4 nor Ku70 accumulates at putative DSB sites after irradiation (e.g. in a pattern resembling that of γH2AX). Therefore, induction of DNA damage has little to no effect on the spatial and temporal localization of XRCC4 and Ku70, consistent with these proteins being present by default at these stages.

We also analyzed the localization of 53BP1, which has a main role in protecting broken DNA ends from resection during NHEJ repair. In control cells, 53BP1 is absent during leptotene, zygotene and early pachytene (Fig 8A) but present by mid-pachytene (Fig 8B), accumulating over the chromatin of the sex chromosomes, in a similar way to γH2AX. 53BP1 signal is maintained during late pachytene (Fig 8C) and early diplotene (Fig 8D) but becomes weak by late diplotene.

After irradiation, in addition to sex chromosomes, 53BP1 localizes to the autosomes from mid-pachytene up to the end of diplotene. One hour after treatment (Fig 8E–8H), a large number of irregularly shaped foci are observed on the autosomes, very similar to the γH2AX eruptions. Indeed, most 53BP1 foci on the autosomes co-localize with γH2AX, although unassociated foci of both proteins are also observed. The same pattern is found at both 24 (Fig 8I–8L) and 72 hours after treatment (Fig 8M–8P). We observed that some chromosomal bridges, which are frequent in cells after treatment, are associated with 53BP1 (Fig 8N), indicating the involvement of NHEJ repair pathway proteins in this type of chromosome interactions.

The quantitative analysis of 53BP1 after treatment (Fig 8Q and 8R, S1 Table) shows the dramatic increase in the number of foci one hour after irradiation in mid- and late pachytene and early diplotene spermatocytes and its sharp decline 24 and 72 hours later. This pattern clearly contrasts with and seems antagonistic to that of RAD51, with NHEJ proteins acting as a fast response and HR proteins acting in two phases, weakly immediately after DNA damage and strongly 24 hours later. We also observed that late stages tend to have more 53BP1 foci, indicating stage-specific differences in the response. Nevertheless, by 72 hours after irradiation, all stages show control levels of 53BP1.

Discussion

The accurate repair of DNA damage is critical for the survival of cells. Meiosis is an excellent model to investigate the response of cells to genomic damage owing to the occurrence of programmed DNA DSBs. However, the response to this endogenous damage must coexist with the sporadic occurrence of exogenous DNA damage, for instance, that caused by exposure to ionizing radiation. The specific processes that occur during meiosis, with the assembly of the SC being the most relevant, work in combination with the endogenous program to bias DSB repair towards the HR pathway [16,17]. Nevertheless, at the end of first meiotic prophase, some constraints might be relaxed, allowing the operation of somatic-like mechanisms. The results presented here offer new ways to understand the interplay of these two responses including how and when this transition occurs during male mouse meiosis.
Fig 8. Pattern of 53BP1 at different stages of prophase-I by recovery time after irradiation. SYCP3 (red), γH2AX (blue) and 53BP1 (green).

(A-D) Control. 53BP1 is first detected at mid-pachytene (B) over the sex chromosomes and is maintained during late pachytene (C) and...
Differential stages of prophase-I have different responses to DNA damage

Phosphorylation of histone H2AX is one of the first key events to occur in response to DNA induced damage [21]. Two types of responses can be clearly distinguished in prophase-I according to the cellular phase: a massive response, characterizing the early stages, and a more focused response from mid-pachytene to diplotene. This focused response is typically found in somatic cells [37] even though, under irradiation overexposure, both somatic and meiotic cells can show a pan-nuclear response [21,55]. However, in our case, all cells were exposed to the same dose of irradiation; therefore, the differences in response are not due to dosage-dependent effects. The origin of these differences could be related, in part, to changes in chromatin configuration and transcriptional activity, as previously suggested for somatic cells [36]. Highly dynamic replacement and modification of histones and proteins associated with chromatin are known to occur during prophase-I [34,35,57–59]. Mouse spermatocytes in early prophase-I are characterized by a widespread distribution of histone H3 monomethylated at lysine 4 and trimethylated at lysine 9, which are both related to chromatin compaction and transcriptional repression [34,35,60]. These modifications are lost or re-localized between early and mid-pachytene, concomitant with other relevant epigenetic changes, such as the incorporation of histone H1t, which is related to the competency of cells to proceed to chromatin condensation stages [57], and a general reactivation of transcriptional activity, which is accompanied by the acetylation of histone H3 and other associated factors [34,35,61,62]. Therefore, the epigenetic changes occurring in meiotic cells at this stage likely act as regulatory factors modulating the DNA damage response.

Changes in chromosome organization may also play a role in the shift in the DNA damage response. In C. elegans, changes in both chromatin conformation and organization of the SC central element are proposed to be involved in the change in the DNA damage response during the mid to late pachytene transition [63,64]. Indeed, exogenous damage can lead to desynapsis of homologous chromosomes [64]. Although no dramatic remodeling of the SC occurs in mouse spermatocytes during this transition, the gradual lengthening of the SC during pachytene [65] is a feature that potentially resembles such reorganization and thus may change the framework in which DNA repair proteins function.

An additional cause of this change may be related to the different kinases that promote H2AX phosphorylation. At least two rounds of H2AX phosphorylation dependent on two different kinases have been proposed to occur in mouse meiosis: the first during leptotene involving ATM and the second at the end of zygotene involving ATR [40,47]. Our efforts to corroborate this hypothesis by immunostaining for kinases, including ATM, ATR and DNA-PK, were unsuccessful. However, indirect proof can be inferred. In this sense, ATM
kinase activity seems to produce an amplification loop in the phosphorylation of H2AX that extends up to several megabases beyond the DSBs [42], whereas the phosphorylation produced by ATR and DNA-PK entails a more focused response in which the signal is limited to areas close to DSBs [66]. Therefore, the two responses we observed with γH2AX may reflect a main role of ATM at the beginning of prophase-I and a higher activity of ATR and DNA-PK at later stages [47]. Interestingly, the response of somatic cells to irradiation, in which most DSBs are repaired by NHEJ [67,68], usually produces discrete foci of γH2AX in the nucleus [37,42], similar to those found in pachytene and diplotene spermatocytes. Therefore, it seems that early stages have a meiotic-specific γH2AX response, while late stages have a repair response more similar to somatic cells (Fig 9).

Based on γH2AX removal, the early response seems to be more efficient as cells irradiated at early stages return to control levels 24 after treatment. In contrast, cells irradiated at later meiotic stages retain a number of γH2AX foci for the duration of recovery. This contradicts findings that, on the basis of the removal dynamics of several repair proteins, suggested repair of DSBs induced at early stages of meiosis is slower than those occurring at later stages [20]. Therefore, it is important to be cautious with these interpretations. We found that many of the γH2AX foci observed at the different stages and recovery times are not associated with DMC1, RAD51 or 53BP1. The persistence of γH2AX in late stages may not be completely related to a delay in the completion of DNA repair but instead to delayed dephosphorylation or turnover of the histone [37]. A similar persistence of γH2AX foci has been also reported after etoposide-induced damage [21]. On the other hand, a substantial number of DMC1, RAD51 or 53BP1 foci are associated with chromosomes long after γH2AX has been displaced, indicating the persistence of unrepaired events. Therefore, it seems that the production of exogenous DSBs challenges both early and late repair pathways during meiosis, resulting in an overall lower efficiency of meiotic repair of exogenous damage compared to somatic cells, as has been previously suggested [18,20].

**Response to endogenous DNA damage involves similar, but not identical, responses of DMC1 and RAD51**

Our analysis of DSB repair pathways clearly indicates that HR is preeminent or exclusive at early meiotic stages, up to mid-pachytene, and reveals interesting clues about the pattern of HR response under both normal and experimental situations.

We identified a population of early leptotene cells that in control mice is characterized by a low number of γH2AX and DMC1 foci. Then, a burst of these proteins is detected in late leptotene. Although we cannot rule out the possibility that these two patterns are just the two extremes of a linear rise of DSBs during leptotene [53], it is also possible that they represent two different physiological stages. While early DSBs are clearly SPO11-dependent, the rate in which they arise is limited, probably owing to the action of a limited number of SPO11 complexes [53,69] or to restrictions imposed by associated factors. Some proteins that stimulate SPO11 activity, like IHO1, are associated with the AEs [70]. Therefore, DSB production could be limited in a chromosomal context in which AEs have not yet formed. Given this context, these DSBs would be able to only trigger a focus-limited (somatic-like?) γH2AX response. The extensive γH2AX labeling of this early leptotene population after irradiation indicates that these cells are competent to display a broad (meiotic) γH2AX reaction. Nonetheless, this bona fide meiotic response is only detected later in leptotene, once AEs have become more extended. This interpretation poses interesting questions about the transition from spermatogonia to meiosis, which includes other puzzling features like the premeiotic pairing of homologous chromosomes [71].
An intriguing issue arose when we compared the distribution of DMC1 and RAD51. We noticed that these two proteins tend to form mixed foci but that their localization patterns are not identical. Several studies reporting similar findings in budding yeast, plants and female mouse meiosis have suggested that these proteins occupy different positions along the nucleoprotein filaments [15,72,73], perhaps performing complementary functions. More strikingly, we found that the temporal pattern of DMC1 and RAD51 do not coincide, particularly in the transition from mid to late pachytene. Temporal displacement between DMC1 and RAD51 loading and unloading has been also observed in plant meiosis [74]. This result may provide
insight on the last stages of meiotic DNA repair pattern, particularly on the sex chromosomes. Several studies have hypothesized that DSBs on the X and Y chromosomes do not have homologous templates which can be used for repair, except obviously the pseudoautosomal region, and that repair can only be accomplished with the sister chromatid [33,36,75,76]. DMC1 may play a key role in interhomolog bias [15], such that its persistence on the sex chromosomes may explain why unresolved DSBs remain on these chromosomes long after most breaks have been repaired on the autosomes. The removal of DMC1 from sex chromosomes, and autosomes, at the mid-late pachytene stage may be related to a relaxation of the interhomolog bias, allowing the repair with the sister chromatid.

**Response to irradiation in early meiosis is characterized by the action of HR proteins that end at mid-pachytene**

Irradiation clearly stimulates the increase in the number of DMC1 and RAD51 foci from lepto-tene up to early pachytene. We did not find any markers for NHEJ at these stages. Although we cannot rule out that alternative NHEJ pathways might be present, it seems that the early response is mostly mediated by HR mechanisms, consistent with the findings of other studies [17,18,20]. It is reasonable to assume that induction of additional breaks cells will simply use the repair machinery that is already present. Therefore, endogenous and exogenous DSBs can enter the same repair pathway. Consequences of this include, for instance, an increased number of chiasmata, as previously reported [20,77]. Nevertheless, this response is limited as prophase-I proceeds. The net increase of DMC1/RAD51 foci at each stage is lower as spermatocytes progress to more advanced stages. Most strikingly, a meiotic HR response to exogenous DNA damage is very weak or not detected at mid-pachytene, thus providing additional evidence of the functional shift of spermatocytes at this stage (Fig 9). This transition likely involves the cessation of expression of some meiotic-specific genes (like DMC1) and the initiation of a new gene expression profile [78–80]. This is interesting not only in terms of DNA repair but also in relation to the regulation of meiotic progression. Several studies have provided evidence of a pachytene checkpoint that monitors DNA repair, chromosome synopsis and other physiological processes such as sex chromosome inactivation [40,54,81–83]. Although the mechanisms that drive this checkpoint in mouse have not been completely elucidated, defective spermatocytes appear to be largely eliminated at a specific point of meiotic progression, identified as stage IV of the seminiferous epithelium in mouse, which most likely corresponds to the mid-pachytene stage [54,81,84]. Once cells have cleared this checkpoint, inactivation of these surveillance mechanisms would be necessary to allow the progression of spermatocytes to later stages, allowing for instance desynapsis of chromosomes during diplote-nne without triggering meiotic arrest or inactivation of desynapsing regions. Likewise, inactivation of the early meiotic DNA damage response would be necessary after passing the checkpoint at mid-pachytene, with new events of DNA damage that occur from this stage onwards being subject to new control mechanisms, as previously suggested [21]. These new mechanisms would rely on different checkpoints, as exemplified by the elimination of spermatocytes in mutant mice for late recombination proteins such as MLH1 or MLH3 at metaphase-I [85,86].

**NHEJ response is quickly stimulated upon irradiation from mid-pachytene onwards**

One may expect that, in the absence of a DMC1 response at mid-pachytene, RAD51 takes the role of driving HR repair at this stage. As discussed above, RAD51 remains associated with chromosomes after DMC1 has detached, and previous reports have indicated that RAD51 is
inducible in late prophase-I after irradiation treatments [18,20,21]. Consistent with this, one hour after irradiation, we found a modest but significant increase in RAD51 from late pachytene onwards; however, the increase was not significant at mid-pachytene. Instead, we observed increased levels of 53BP1 at all stages from mid-pachytene to late diplotene, indicating a faster response of the NHEJ pathway at these stages. This contrasts with a previous study that reported the presence of 53BP1 only after longer periods of recovery [18]. Differences in methodological approaches used to determine 53BP1 localization might account for these discrepancies.

The quick trigger of NHEJ in late prophase-I may be due to a change in the choice of the default mechanism for DSB repair (Fig 9). Somatic cells first attempt to use NHEJ to repair DSBs, even in the G\textsubscript{2} phase of the cell cycle when a sister chromatid is available to carry out more reliable repair by HR [2,5,87]. The choice of NHEJ as the default mechanism from late pachytene onwards is illustrated by the constitutive presence of Ku70 and XRCC4 in the nucleus and the location of 53BP1 on the sex chromosomes. Therefore, as soon as new exogenous or even endogenous DNA breaks appear, this mechanism would quickly respond. This proposal is plausible in terms of the biochemistry of DNA repair. Given that the choice between NHEJ or HR relies on the regulation of DNA resection around the break point [4], for which 53BP1 has an inhibitory role [23,24], it is clear that NHEJ must be a first option. Otherwise, once resection has been performed, repair by this mechanism would be no longer possible. Nevertheless, it is also clear that both mechanisms are acting at the same time, raising the possibility that NHEJ and HR proteins are competing for DNA repair, especially from late pachytene onwards. In any case, after 24 hours of recovery, few 53BP1 loci remain, which may be due to rapid repair by NHEJ [2], but also to competition with HR. In somatic cells, HR seems to be preeminent in regions of high transcrip tional activity [56], which is indeed the case of late pachytene and diplotene cells during meiosis. The increased presence of RAD51 foci 24 hours after irradiation indicates that HR repair mechanisms prevail again at that time.

**Late HR response involves RAD51 only**

The late HR response has many differences with the early one. The most relevant is that it only involves RAD51. In the transition to a somatic-like DNA damage response, DMC1 is clearly no longer inducible, likely related to the change in gene expression pattern during pachytene [78–80]. The activity of RAD51 alone means that some of the constraints introduced by DMC1 in relation to DNA repair, such as homologous bias [11–13,15], would be relaxed. Therefore, this late repair could favor repair with the sister chromatid, which would be advantageous at the diplotene stage as desynapsis of chromosomes potentially hinders repair with the homologous chromosome.

The RAD51 foci present during the late HR response are larger. Although we do not have a clear explanation for the morphological change of RAD51 foci, differential organization of the repair machinery around the break point involving, for instance, the accumulation of several DSBs in each foci, or comprising the resection of longer DNA stretches, may account for this change, as previously suggested in *C. elegans* [63]. These foci are also correlated with the formation of smaller discrete γH2AX foci, which, as mentioned above, may be due to the action of ATR or DNA-PK over ATM in the phosphorylation of H2AX, leading to a different architecture of the repair foci [42,66].

Finally, the finding of RAD51 foci not associated with the AEs/SCs of chromosomes is an intriguing feature. Whether RAD51 foci are always associated with AEs/SCs has been a matter of long debate [88]. Current models propose that endogenous DSBs in early stages are produced either in the context of the AE or rapidly taken there by the action of regulatory factors,
including MEI4, IHO1 and HORMAD1, among others [70,89]. This is probably provided by their ability to interact with SPO11 before or at the time of DSB production. We found that most DSBs induced after radiation also localize at the AEs/SCs, as revealed by the pattern of DMC1/RAD51 foci and as reported in a previous study [50]. Thus, at early meiotic stages, both endogenous and exogenously induced DSBs likely rely on similar mechanisms to be taken to the chromosomal axis. However, at late stages, the situation might be different. The presence of IHO1 and HORMAD1 has been reported in diplotene cells [70,90]; however, it remains unclear whether these proteins, or others required for DSB localization at the axes, are completely functional at these stages. Partial failure of this process might explain the fraction of DSBs located far from AEs/SCs. Likewise, the progressive loss of these proteins as prophase-I proceeds may account for the increased frequency of non-associated foci in late diplotene spermatocytes compared with previous stages.

**Homologous recombination homeostasis and chromosomal bridges**

The appearance of chromosomal bridges involving SYCP3-positive filaments is an intriguing feature that poses a number of questions about the nature of DNA repair. The presence of chromosomal connections and fragments is commonly found in irradiation experiments [91,92]; however, they are usually observed in metaphase and do not involve SC connections. To our knowledge, our study is the first to report that interactions between non-homologous chromosomes may involve not only the DNA contacts but also the axial structures they are attached to. This contrasts with the normal interactions of endogenous DSBs, which do not involve the formation of connections between the AEs of homologous chromosomes. A more in-depth characterization of these connections is needed to better understand the organization of the SC around break points and their role in promoting, facilitating or stabilizing chromosomal links.

Bridges appear soon after irradiation, indicating they are part of a very fast response, and at increased frequencies with recovery times. Several possible interpretations can be drawn from these results. Given that bridges are observed between non-homologous chromosomes, or between non-homologous sequences of the same chromosome (intrachromosomal junctions), one would expect that they correspond to DSBs repaired by NHEJ. However, the appearance of bridges at early stages such as zygotene and early pachytene, in which the main repair mechanism is HR, challenges this interpretation. Our results and previous reports [16,17] suggest that NHEJ does not operate at early stages; however, alternative NHEJ mechanisms may potentially be present. Involvement of the NHEJ repair pathway in the formation of chromosomal bridges is doubtless only from mid-pachytene onwards.

The presence of chromosomal bridges at early stages may be a consequence of the action of the HR pathway over homologous regions. This idea may seem completely counterintuitive but would be supported by the observation of DMC1/RAD51 filaments bridging non-homologous chromosomes. For HR to efficiently start, a minimum length of perfect homology is needed, which in mammals is 200–250 base pairs [4]. For this reason, repair templates are usually the sister chromatid or the homologous chromosome. However, the multitude of repeated sequences in the genome could provide sufficient homology to induce repair by HR. In the normal course of meiosis, endogenous DSBs are prevented over repeated DNA sequences. Moreover, mismatch repair mechanisms are in place to avoid recombination between highly homologous sequences of non-homologous chromosomes [93]. In addition, the number of DSBs is tightly regulated to not exceed a certain number, thus allowing mismatch repair mechanisms to function effectively [94]. Proteins like MEI4 and IHO1 seem to be involved in this limitation [70,89]. However, the excess of DSBs produced by radiation may have a
deregulatory effect on the control of repair mechanisms such that homology requirements may be bypassed, thus allowing repair between non-homologous chromosomes.

Regardless of the mechanisms used to form bridges, the final output is the production of chromosome connections that likely lead to the occurrence of translocations and fragmentation. This may cause cells to be compromised in the faithful distribution of chromosomes during the first meiotic division. Indeed, a noticeable increase of apoptosis is observed in metaphase/anaphase cells 24 and 72 hours after irradiation.

In conclusion, the results presented here provide new insights on the transition between different programs of DNA repair during meiosis that act in a stage-dependent manner. A switch in DNA damage repair responses during meiosis has been also reported in other animal models such as *C. elegans* [63,64] and *Drosophila* [95]. Strikingly, these transitions also occur at the mid or late pachytene stages, indicating they may represent a conserved feature of meiosis. However, since both *Drosophila* and *C. elegans* control synopsis and DNA repair differently than mammals, particularly as they lack a DMC1 orthologue, the regulation of this transition might be different.

The evidence presented also offers new clues about the location and dynamics of DNA repair mechanisms during meiosis and raises new questions about the differential functions performed by DMC1 and RAD51. The late HR pathway very much resembles the somatic response, presenting focalized γH2AX and involving only RAD51. This somatic-like response likely acts to repair DSBs that were not properly repaired by the meiotic default pathway (e.g. those on the sex chromosomes) or the occasional DNA damage that occurs after the primary phase of meiotic repair has concluded until cells start to condense chromatin and prepare for cell division. At this point, re-triggering a complex repair mechanism leading to the production of crossovers (provided chiasmata formation has been properly accomplished) would not be necessary. A simpler response using NHEJ or somatic HR would be sufficient. As previously suggested [63], this shift could simply be contributing to the maintenance of genome integrity before spermatocytes are engaged in segregating chromosomes to daughter cells.

**Materials and methods**

Adult CD1 male mice were used in this study. Animals were kept at the animal facility of the Universidad Autónoma de Madrid, following the animal care standards of the institution. All experiments were approved by the UAM Ethics Committee (certificate CEI 55-999-A045). Males were exposed to 5Gy gamma radiation by whole body irradiation in a CIS Bio International irradiator, equipped with a Cesium¹³⁷ source. Mice were sacrificed by cervical dislocation 1, 24 and 72 hours after irradiation and the seminiferous tubules processed as described below. Testicular samples of Spo11 knockout mice [32] were kindly shared by Dan Camerini-Otero (NIDDK, NIH, Bethesda, MD).

**Cell spreads and squashes**

For spermatocyte spreads, we used the procedure described by Peters and coworkers [96]. Seminiferous tubules were disaggregated with forceps in a petri dish and a cell suspension was collected in phosphate buffered saline (PBS: 137 mM NaCl, 2.7 mM KCl, 10.1 mM Na₂HPO₄, 1.7 mM KH₂PO₄, pH 7.4). After tubule fragments settled to the bottom of the dish, the cell suspension was transferred to a tube and centrifuged. The pellet was then resuspended in 400 μl of 100 mM sucrose. Cells were spread onto a slide submerged in 1% formaldehyde in distilled water containing 50 mM Na₂B₄O₇ and 0.15% Triton X-100 and then left to dry for two hours. Slides were subsequently washed with 0.04% Photo-Flo (Kodak) in distilled water and air-dried before being used for immunofluorescence or stored at -80°C.
Spermatocyte squashes were prepared as previously described [97]. Seminiferous tubules were fixed for 10 minutes in 2% formaldehyde in PBS containing 0.1% Triton X-100. Fragments of tubules were placed on a slide coated with 1 mg/ml poly-L-lysine (Sigma) with two drops of fixative. A coverslip was put on top of the tubules and the cells were released by gently pressing the coverslip with a pencil. Finally, tubules were squashed, the slide was frozen in liquid nitrogen and the coverslip removed with a blade. Slides were immediately placed in PBS for further use.

**Immunofluorescence**

Spread and squashed slides were rinsed three times for 5 min each in PBS and incubated overnight at room temperature with primary antibodies diluted in PBS. The following primary antibodies and dilutions were used: mouse monoclonal anti-SYCP3 (Abcam, 97672) at 1:200; rabbit anti-SYCP3 (Abcam, 15093) at 1:100; mouse monoclonal against histone H2AX phosphorylated at serine 139 (γ-H2AX) (Upstate, 05–636) at 1:1000; rabbit anti-DMC1 (Santa Cruz, SC-22768) at 1:50; rabbit anti-RAD51 (Santa Cruz SC-8349) at 1:50; rabbit anti-53BP1 (Abcam 36823) at 1:100; goat anti-XRCC4 (Santa Cruz, SC-8285) at 1:100; goat anti-Ku70 (Santa Cruz, SC-1486) at 1:50. After incubation, slides were rinsed in PBS three times for 5 minutes each and subsequently incubated with the appropriate secondary antibodies in a moist chamber at room temperature for 1 h. We used anti-rabbit, anti-mouse and anti-goat secondary antibodies raised in donkey and conjugated with either Alexa 350, Alexa 488, Alexa 594 (Invitrogen), DyLight 549 or DyLight 649 (Jackson ImmunoResearch). Slides were subsequently rinsed in PBS three times for 5 min each and mounted with Vectashield (Vector). For double detection of two antibodies raised in the same species, we used Fab secondary antibodies as previously described [98].

Observations were made on an Olympus BX61 microscope equipped with a motorized Z axis. Images were captured with an Olympus DP72 digital camera using the Cell-F software (Olympus, Hamburg, Germany) and processed using the public domain software ImageJ (National Institutes of Health, USA; http://rsb.info.nih.gov/ij) and Adobe Photoshop 7.0.

**Testicular sections and TUNEL assay**

Testicles were fixed in cold 1% formaldehyde in PBS for 6 hours and then dehydrated and embedded in paraffin. Transverse sections (7 μm) were cut and mounted onto slides. Slides were then deparaffinized and treated with 0.1% sodium citrate buffer containing 0.1% Triton-X 100 for 10 min at 37˚C. Sections were subsequently processed for immunofluorescence as described above or for TUNEL (Roche) following manufacturer instructions. Slides were counterstained with DAPI and mounted with Vectashield.

**EdU incorporation**

Mice were injected intraperitoneal with 100 μl 10 mM 5-ethynyl-2-deoxyuridine (EdU) (Thermo Fisher Scientific) diluted in PBS. Since first meiotic prophase time length is longer than 72 hours (the longest period of recovery we analyzed), it was necessary to irradiate mice at two different times after EdU injection in order to evaluate the interference of meiotic progression at different sub-stages. Thus, mice were irradiated with 5Gy of gamma radiation one or six days after EdU administration and sacrificed 0, 24 and 72 hours after irradiation (see [S1 Fig](#) for a detailed description of the EdU administration, mice irradiation and sample collection). This allowed us to cover the most relevant periods under analysis: the transition from leptotene to mid-pachytene and from mid-pachytene to diplotene. Testicular samples were processed as described above and spermatocytes were labelled with anti-SYCP3. EdU was revealed using the Click-iT as recommended by the supplier.
Statistical analysis

γ-H2AX, DMC1, RAD51 and 53BP1 foci were scored manually in three individuals per each recovery time and the results were compared between different cell stages and times after irradiation. Foci were counted in at least 15 cells for each protein and cell stage per individual. Data were analyzed using ANOVA and Tukey’s multiple comparison tests for individual comparisons between different stages. Chromosomal bridges, apoptotic cells and EdU labelled spermatocytes were scored in two individuals for each recovery time after irradiation. For chromosomal bridges, spermatocytes containing at least one bridge were considered as positive. At least 150 cell were analyzed per individual. For the TUNEL assay, 300 tubules (150 per individual) were analyzed for each treatment, recording the proportion of tubules with apoptotic cells and the total number of positive cells, which were classified as spermatogonia, prophase-I and division spermatocytes owing to their size, position on the seminiferous epithelium and chromosome condensation. For EdU labeling, at least 200 cells were scored per individual and the proportion of labeled cells at different meiotic stages was analyzed using Chi squared test. Statistics and graphics were made using GraphPad Prism 6 or Excel software.

Supporting information

S1 Table. Quantitative data for all proteins analyzed and chromosomal bridges organized by cell stage and recovery time.

S1 Fig. Spermatogenesis progression in CD1 mice. A) Schedule for EdU labeling and analysis. Mice were injected with EdU at day 0. Then, control samples were taken at day 1, day 2 and day 4. A subset of mice were irradiated and samples collected 24 (day 2) and 72 hours (day 4) later. In an additional experiment, control samples were taken at day 6, day 7 and day 9. A subset of mice were irradiated at day 6 and samples collected 24 (day 7) and 72 hours (day 9) later. B) Analysis of the proportion of spermatocytes labelled with EdU at different stages. Bars in the graph indicate the proportional contribution of each stage at the different time points analyzed. Percentages and the total number of cells analyzed are indicated in the table below. The results of day 2, day 4, day 7 and day 9 were compared between control and irradiated cells. Chi squared test showed no statistical differences in any of the days analyzed (X² = 0.78, 2 degrees of freedom (df) for day 2; X² = 8.01, 4 df for day 7; X² = 6.92, 3 df for day 9), excepting day 4 (X² = 32.08, 3 df). This is due to the conspicuous absence of leptotene cells 72 hours after irradiation. When this cell population was not considered, differences were not significant (X² = 0.48, 2 df). C) EdU labeling (red) and SYCP3 staining (green) in spermatocytes and estimated length of meiotic stages. The duration of each stage was estimated on the basis of the day after EdU injection in which each stage is detected for the first time. The approximate duration of 1 day for leptotene, zygotene and early pachytene is consistent with previous reports [54,99].

S2 Fig. Apoptosis induction after irradiation. TUNEL (green) and DAPI (blue). (A-C) Section of a seminiferous tubule 24 hours after treatment. Apoptotic cells (arrows) are found close to the basal stratum of the seminiferous epithelium. (D-F) Section of a seminiferous tubule 72 hours after treatment showing apoptotic cells at metaphase or telophase (arrows). Enlarged images from D-F showing apoptotic cells at metaphase (G-I) and telophase (J-L). (M). Quantitative distribution of apoptotic cells. Total number of apoptotic cells were recorded in 300 seminiferous tubules. Peak apoptosis is observed 24 after irradiation with 58.6% of tubules.
showing at least one apoptotic cell. At this time, spermatogonia are the most affected population, followed by spermatocytes and cells undergoing division. After 72 hours of recovery, the total number of apoptotic cells decreases with only 27.6% of tubules showing apoptotic cells. At this recovery time, the majority of cells undergoing apoptosis are at metaphase or anaphase.

**S3 Fig. DMC1 localization in SPO11 knockout mice spermatocytes.** (A-C) SYCP3 (green) and (A'-C') DMC1 (red) at early leptotene (A, A'), mid leptotene (B, B') and zygotene-like (C, C'). No specific signal of DMC1 is detected at any of the meiotic stages in this knockout model.

**S4 Fig. Quantitative analysis of DMC1 dynamics during early prophase-I.** A) Analysis of DMC1 distribution by time of recovery. Six substages were considered (EL: early leptotene; LL: mid-late leptotene; EZ: early-mid zygotene; LZ: late zygotene; EP: early pachytene; MP: mid pachytene). The six populations, including early leptotene, are clearly distinguishable in the control. A low number of foci is found in EL cells but numbers increase in LL, peak in EZ and then gradually decrease in LZ, EP and MP cells. ANOVA analysis showed statistical differences (p\(\leq\)0.0001) for the control and the three recovery times, and Tukey’s multiple comparisons test for individual comparisons between different stages showed statistical differences in all cases (*: p\(\leq\)0.05; **: p\(\leq\)0.01; ***: p\(\leq\)0.001; ****: p\(\leq\)0.0001). B) Distribution of DMC1 foci arranged according to the recovery time after irradiation and the putative stages that cells should have reached at that time. We arranged four cell populations: early leptotene, mid-late leptotene, early-mid zygotene and late zygotene. For each case, irradiated cells were compared with their respective control counterparts and statistical differences indicated (ANOVA and Tukey’s multiple comparisons test). All cells advanced to mid-pachytene after 72 hours of recovery. Since DMC1 is not inducible at this stage, all cells reached DMC1 control levels, regardless of whether repair had been completed or not. EL: early leptotene; LL: mid-late leptotene; EZ: early-mid zygotene; LZ: late zygote; EP: early pachytene; MP: mid pachytene; ns: non-significant; *: p\(\leq\)0.05; **: p\(\leq\)0.01; ***: p\(\leq\)0.001; ****: p\(\leq\)0.0001.

**S5 Fig. Localization of DMC1 and RAD51 in control spermatocytes.** SYCP3 (blue), DMC1 (green) and RAD51 (red). Merge of the (A-F) SYCP3 and DMC1 channels, (A'-E') SYCP3 and RAD51 channels and (A"-F") SYCP3, DMC1 and RAD51 channels. DMC1 and RAD51 are largely co-localized in foci observed from early leptotene to mid-pachytene (A-D). However, DMC1 and RAD51 signal on these foci are usually not identical in size or shape. Moreover, there are some instances in which either of the two proteins seem to form single foci. At mid-late pachytene (E-E"), DMC1 is no longer present on the chromosomes, but RAD51 is still abundantly observed on both autosomes and sex chromosomes (X and Y). By late pachytene (F-F"), neither DMC1 nor RAD51 are observed.

**S6 Fig. Localization of DMC1 and RAD51 at different stages of prophase-I by different recovery time after irradiation.** SYCP3 (blue), DMC1 (green) and RAD51 (red). (A-F) Control. The cells shown in A-D are the same as those shown in S5 Fig. (G-L) 1 hour after irradiation. As shown in Figs 4 and 6, both DMC1 and RAD51 become more abundant after irradiation with both proteins being present in the same foci in most cases; however, the overlap in signals is not identical in many instances as the sizes and shapes of foci of the individual proteins differ. From late pachytene onwards, only RAD51 foci are detectable. (M-R) An analogous result is found 24 after irradiation. Filaments containing both DMC1 and RAD are
observed in some spermatocytes (arrows in M).

**(TIF)**

**S7 Fig. Localization of XRCC4 in testicular sections.** γH2AX (green), XRCC4 (red) and DAPI (blue) in seminiferous tubules at equivalent developmental stages. (A-D) Control. Basal layers of spermatocytes, corresponding to leptotene and zygotene, are broadly stained with γH2AX and devoid of XRCC4. Spermatocytes in the interstitial strata of the epithelium show an almost inverse labeling pattern, with abundant XRCC4 and nearly no γH2AX staining. (E-H). 1 hour after irradiation. Cells showing broad γH2AX labeling in the basal strata are again devoid of XRCC4. In contrast, spermatocytes stained with XRCC4 now also have an abundance of γH2AX localized foci, corresponding to the late γH2AX response. No noticeable increase in the intensity of XRCC4 labeling is observed. (I-L) 24 hours and (M-P) 72 hours after irradiation γH2AX tends to return to control levels. No variation of XRCC4 labeling is observed after longer periods of recovery.

**(TIF)**

**S8 Fig. Localization of Ku70 in testicular squashes.** Ku70 (red), SYCP3 (green) and DAPI (blue) in squashed spermatocytes at different meiotic stages. Z-projections of image stacks across spermatocytes. Ku70 seems to be absent in early or mid-pachytene cells in both control (A-B) and irradiated spermatocytes (C-D). At late pachytene, a clear labeling is observed in the nucleus, covering most of the chromosomes. The labeling is more intense in diplotene spermatocytes. No evident differences in the intensity of labeling are observed between control and irradiated cells.

**(TIF)**

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