Spindle assembly checkpoint proteins regulate and monitor meiotic synopsis in C. elegans

Tisha Bohr, Christian R. Nelson, Erin Klee, and Needhi Bhalla

Department of Molecular, Cell, and Developmental Biology, University of California, Santa Cruz, Santa Cruz, CA 95064

Homologue synopsis is required for meiotic chromosome segregation, but how synopsis is initiated between chromosomes is poorly understood. In Caenorhabditis elegans, synopsis and a checkpoint that monitors synopsis depend on pairing centers (PCs), cis-acting loci that interact with nuclear envelope proteins, such as SUN-1, to access cytoplasmic microtubules. Here, we report that spindle assembly checkpoint (SAC) components MAD-1, MAD-2, and BUB-3 are required to negatively regulate synopsis and promote the synopsis checkpoint response. Both of these roles are independent of a conserved component of the anaphase-promoting complex, indicating a unique role for these proteins in meiotic prophase. MAD-1 and MAD-2 localize to the periphery of meiotic nuclei and interact with SUN-1, suggesting a role at PCs. Consistent with this idea, MAD-1 and BUB-3 require full PC function to inhibit synopsis. We propose that SAC proteins monitor the stability of pairing, or tension, between homologues to regulate synopsis and elicit a checkpoint response.

Introduction

Cell cycle checkpoints ensure accurate chromosome segregation by monitoring the progression of critical events (Murray, 1992). When errors occur, checkpoints prevent the production of aneuploid daughter cells by either arresting the cell cycle to promote error correction or targeting the cell for apoptosis. Aneuploidy is a hallmark of tumor cells undergoing mitosis (Kops et al., 1992). When errors occur, checkpoints prevent the production of aneuploid daughter cells by either arresting the cell cycle to promote error correction or targeting the cell for apoptosis. Aneuploidy is a hallmark of tumor cells undergoing mitosis (Kops et al., 1992) and is associated with birth defects and infertility during sexual reproduction (Hassold and Hunt, 2001).

Sexual reproduction requires meiosis, a specialized cell division that produces gametes such as eggs and sperm. During meiotic prophase, homologous chromosomes pair and synapse to undergo crossover recombination, a prerequisite for proper meiotic chromosome segregation (Bhalla and Dernburg, 2008). In Caenorhabditis elegans, the synopsis checkpoint induces apoptosis to remove nuclei with unsynapsed chromosomes (Bhalla and Dernburg, 2005). This checkpoint depends on cis-acting sites near one end of each chromosome termed pairing centers (PCs; Bhalla and Dernburg, 2005), which are also essential for pairing and synopsis (MacQueen et al., 2005). Early in meiotic prophase PCs recruit factors, such as HIM-8, ZIM-1, ZIM-2, and ZIM-3 (Phillips et al., 2005; Phillips and Dernburg, 2006), to assemble a transient regulatory platform that interacts with the conserved nuclear envelope proteins SUN-1 and ZYG-12. This interaction allows PCs access to the cytoplasmic microtubule network and the microtubule-associated motor, dynein (Penkner et al., 2007; Sato et al., 2009; Labrador et al., 2013). The mobilization of chromosomes by cytoskeletal forces is a conserved feature of meiotic prophase (Bhalla and Dernburg, 2008) that facilitates homologue pairing and synopsis (Sato et al., 2009; Labrador et al., 2013).

When dynein function is abolished, chromosomes pair but fail to synapse (Sato et al., 2009). These data have led to a working model in which dynein is dispensable for homologue pairing but licenses synopsis through a tension-sensing mechanism (Sato et al., 2009; Wynne et al., 2012). This model proposes that when a chromosome identifies its homologue and remains stably paired, it resists the pulling forces of dynein. This resistance, or tension, is thought to initiate synopsis at PCs. However, if nonhomologous chromosomes interact, they cannot resist dynein’s pulling forces and restart the homology search. How tension between PCs could be monitored to regulate synopsis is unknown.

Despite their functional differences, PCs have been compared with centromeres (Dernburg, 2001; Labella et al., 2011). Both are cis-acting chromosomal sites that nucleate transient structures to mediate microtubule binding, promote specific chromosome behavior, and generate a checkpoint response. In addition, centromeres can act as sites for meiotic synopsis initiation in budding yeast (Tsoubouchi and Roeder, 2005; Tsoubouchi et al., 2008) and Drosophila (Takeo et al., 2011; Tanneti et al., 2011). Centromeres assemble kinetochores to orchestrate chromosome segregation (CheezeMAN and Desai, 2008). Kinetochores also provide a platform for the spindle assembly checkpoint (SAC), which inhibits the anaphase-promoting...
complex (APC) and halts cell cycle progression until all kinetochores are successfully bioriented (Foley and Kapoor, 2013). Because of the similarities between PCs and centromeres, we hypothesized that components of the SAC might act at PCs during meiotic prophase. We report that MAD-1, MAD-2, and BUB-3 are required for the synapsis checkpoint and negatively regulate synapsis in *C. elegans*. Mutation of mad-1, mad-2, or bub-3 suppresses synapsis defects in dynein mutants, implicating SAC components in the tension-sensing mechanism that is thought to license meiotic synapsis. These roles in monitoring and regulating synapsis are independent of a conserved APC component, indicating MAD-1, MAD-2, and BUB-3 are performing a role aside from inhibiting the APC. MAD-1 and MAD-2 localize to the nuclear periphery and coimmunoprecipitate with SUN-1. Furthermore, MAD-1 and BUB-3 require full PC function to inhibit synapsis. Altogether, these data strongly suggest that SAC proteins function at PCs. Therefore, we propose that the ability of some SAC components to monitor tension is conserved and may have been coopted in a variety of biological contexts to maintain genomic integrity.

**Results and discussion**

**MAD-1, MAD-2, and BUB-3 are required for the synapsis checkpoint**

To determine whether SAC proteins are required for the synapsis checkpoint, we used a hypomorphic allele of *mdf-1* (*mdf-1[av19]*) defective in SAC function (Stein et al., 2007; Yamamoto et al., 2008) and null mutations of three core but nonessential SAC components, *mdf-2*Δ and *mdf-3/san-1*Δ (known as *mdf-2* and *mdf-3/san-1*, respectively, in *C. elegans*), and *bub-3*Δ (Kitagawa and Rose, 1999; Essex et al., 2009). We refer to the *mdf-1[av19]* allele as *mdf-1(cd)* for checkpoint deficient. *meDf2* is a deficiency that removes the X chromosome PC (Villemuneve, 1994). Animals heterozygous for *meDf2/*+ have unsynapsed X chromosomes in a subset of meiotic nuclei because synapsis cannot initiate efficiently (MacQueen et al., 2005). We introduced SAC mutations into *meDf2/+* and found that loss of *mdf-1*, *mdf-2*, or *bub-3*, but not *mdf-3*, decreased apoptosis in *meDf2/+* to wild-type levels (Fig. 1 B), illustrating that MAD-1, MAD-2, and BUB-3 are required for the synapsis checkpoint when X chromosomes are unsynapsed.
meDf2 homozygotes exhibit asynapsis in almost all meiotic nuclei (MacQueen et al., 2005). However, these mutant worms exhibit elevated germline apoptosis as the result of the DNA damage checkpoint (Fig. S1, A and B) because functional PCs are required for the synapsis checkpoint (Bhalla and Dernburg, 2005). Mutation of mad-1, mad-2, mad-3, or bub-3 did not reduce apoptosis in meDf2 homozygotes (Fig. S1 B). Therefore, MAD-1, MAD-2, and BUB-3 are specifically required to induce germline apoptosis in response to the synapsis checkpoint.

Next, we tested the requirement for SAC components in the synapsis checkpoint when all chromosomes are unsynapsed. Synapsis requires the assembly of the synaptonemal complex (SC) between homologous chromosomes (Bhalla and Dernburg, 2008). syp-1 mutants do not load SCs between homologues (MacQueen et al., 2002), leading to high levels of checkpoint-induced germline apoptosis as a result of both the synapsis and DNA damage checkpoints (Fig. 1, A, C, and D; Bhalla and Dernburg, 2005). Mutation of mad-1, mad-2, mad-3, or bub-3 in the syp-1 mutant background reduced apoptosis to intermediate levels, indicating loss of one checkpoint but not both (Fig. 1, C and D).

To determine which checkpoint these genes are required for, we prevented the DNA damage checkpoint response in syp-1 mutants by mutating either spo-11 or cep-1 (Fig. 1 A; Bhalla and Dernburg, 2005). SPO-11 generates double-strand breaks that initiate meiotic recombination (Dernburg et al., 1998), and CEP-1 (the C. elegans p53 orthologue) promotes germline apoptosis in response to DNA damage (Derry et al., 2001; Schumacher et al., 2001). Therefore, the elevated apoptosis in spo-11;syp-1 and cep-1;syp-1 double mutants is solely a consequence of the synapsis checkpoint (Fig. 1 A; Bhalla and Dernburg, 2005). Mutation of mad-1 or bub-3 in the spo-11;syp-1 background or mad-2 in the cep-1;syp-1 background produced wild-type levels of apoptosis (Fig. 1, C and D). However, mutation of mad-3 in spo-11;syp-1 mutants did not further decrease apoptosis (Fig. 1 C), suggesting that MAD-3 is required for the DNA damage checkpoint in syp-1 mutants. Because this differs from our results with meDf2;mad-3Δ double mutants (Fig. S1 B), we infer that the DNA damage checkpoint responds differently if all chromosomes are unsynapsed (syp-1 mutants) versus if one pair of chromosomes is unsynapsed (meDf2 mutants). More importantly, these data establish that MAD-1 and MAD-2 are specifically required for the synapsis checkpoint when all chromosomes are unsynapsed.

Mad2, Bub3, and Mad3 form the mitotic checkpoint complex (MCC), which inhibits the APC activator Cdc20 and entry into anaphase (Sudakin et al., 2001). MAD-3’s primary role during the SAC response may be inhibition of the APC (Shonn et al., 2003), suggesting that the APC might also not be involved in the synapsis checkpoint. To test this, we used a temperature-sensitive allele of mat-3, the orthologue of Cdc23/Apc8 and an essential subunit of the APC (Golden et al., 2000). We predicted that if SAC components were acting through the APC, loss of APC activity would elevate germline apoptosis as the result of an inappropriate checkpoint response (Fig. S1 C). However, germline apoptosis in mat-3, syp-1;mat-3;mad-1(cd), or syp-1;mat-3;bub-3Δ mutants was unaffected in comparison with wild-type, syp-1;mad-1(cd), and syp-1;bub-3Δ backgrounds, respectively (Fig. S1 D). We also evaluated whether Cdc20 (FZY-1 in C. elegans) was involved in the synapsis checkpoint using a loss of function allele (Kitagawa et al., 2002) but detected no change in apoptosis in fzy-1 or syp-1;fzy-1;mad-1(cd) mutants when compared with wild-type or syp-1;mad-1(cd) worms, respectively (Fig. S1 E). Together these data indicate that the APC is unlikely to be the target of SAC components in the synapsis checkpoint. Intriguingly, orthologues of some SAC components, but not APC components, have been identified in the Giardia genome (Gourguechon et al., 2013). Loss of these conserved SAC components produces chromosome segregation errors (Vicente and Cande, 2014), suggesting that they may have additional roles in regulating chromosome behavior.

In addition to the synapsis checkpoint, asynapsis is associated with a delay in meiotic progression in which chromosomes remain asymmetrically localized (clustered) in meiotic nuclei (Fig. S2 A; MacQueen et al., 2002). To evaluate whether SAC proteins affect meiotic progression, we quantified the percentage of nuclei in which chromosomes appeared clustered (Fig. S2, B and C). mad-1(cd) and bub-3Δ mutants had slightly more nuclei with clustered chromosomes than wild-type germ-lines, suggesting defects in synapsis or recombination (Fig. S2, B and C). However, neither of the single mutants exhibited any achiasmate chromosomes (not depicted), indicating that cross-over recombination is not disrupted. More importantly, we did not detect any difference in the percentage of nuclei with clustered chromosomes between syp-1;mad-1(cd) or syp-1;bub-3Δ double mutants and syp-1 single mutants (Fig. S2, B and C). Therefore, MAD-1 and BUB-3 are specifically required for the synapsis checkpoint and not for the delay in meiotic progression that accompanies asynapsis.

**MAD-1 and MAD-2 interact with SUN-1 and localize to the periphery of meiotic nuclei**

SAC components localize to unattached kinetochores to initiate checkpoint signaling (Foley and Kapoor, 2013). In meiotic prophase, SUN-1 is present at the nuclear periphery and colocalizes with PCs during pairing and synapsis (Penkner et al., 2007; Sato et al., 2009). To test whether SAC proteins also interact with PC-associated proteins to promote synapsis checkpoint signaling, we performed coimmunoprecipitations (co-IPs) of SUN-1::GFP and probed the immunoprecipitates with antibodies against MAD-1 and MAD-2 (Fig. 2 A). Both MAD-1 and MAD-2 coimmunoprecipitated with SUN-1::GFP but not with our untagged control (Fig. 2 A). We also assessed whether BUB-3 coimmunoprecipitated with SUN-1::GFP but were unable to detect an interaction (not depicted). Therefore, both MAD-1 and MAD-2 interact with the PC-associated protein SUN-1.

We evaluated whether SAC proteins are at the nuclear periphery by staining germlines with antibodies against nuclear pore complexes (NPCs) and MAD-1::GFP or MAD-2. MAD-1::GFP and MAD-2 localized to the nuclear periphery in a punctate pattern (Fig. 2 B), and colocalization with SUN-1::GFP confirmed that both proteins were inside the nucleus (not depicted). MAD-1 localization to the nuclear periphery in embryos is dependent on the nonessential NPC component NPP-5 (Ródenas et al., 2012). However, in meiotic nuclei, MAD-1 or MAD-2 localization is not disrupted in npp-5Δ, sun-1Δ, or npp-5Δ,sun-1Δ mutants (not depicted). We stained for BUB-3 but were unable to localize it in meiotic nuclei (not depicted).

We attempted to localize SAC proteins with PCs. However, despite the biochemical interaction with SUN-1, neither MAD-1 nor MAD-2 colocalized with PC proteins in wild-type, syp-1, or meDf2/+ germlines (not depicted). We
provide two potential explanations for the inconsistency between our biochemical and cytological experiments: (1) the interaction of SAC proteins at PCs is transient, and/or (2) the pool of MAD-1 and MAD-2 that interacts with PCs is a small fraction of the total protein present in meiotic nuclei. Similar explanations have been made to argue that Mad1 and Mad2 sense tension during mitosis despite an inability to localize these proteins to tensionless kinetochores (Maresca and Salmon, 2010).

**MAD-1 and BUB-3 inhibit synapsis in a PC-dependent manner**

Given that the synapsis checkpoint component PCH-2 inhibits synapsis (Deshong et al., 2014), we hypothesized that SAC pro-
teins might also regulate this process. We were concerned that defects in germline organization might complicate our analysis (Kitagawa and Rose, 1999; Stein et al., 2007), so we first analyzed meiotic entry in SAC mutants. In C. elegans, meiotic nuclei undergo mitotic divisions at the distal end of the germ-line until they enter meiotic prophase (Fig. S2 D). Meiotic entry is accompanied by the temporary clustering of chromosomes within nuclei and the appearance of phosphorylated SUN-1 (SUN-1pSer8) at the nuclear envelope (Penkner et al., 2009; Woglar et al., 2013). We assessed whether meiotic entry was affected in SAC mutants by quantifying the number of rows of mitotic germline nuclei from the distal tip to the appearance of SUN-1pSer8 in nuclei with clustered chromosomes (Fig. S3 E). Although mad-2Δ mutants delayed the onset of meiosis, as indicated by an increased number of rows of mitotic germline nuclei compared with wild-type worms, we did not detect any significant difference in the number of rows of mitotic germline nuclei in wild-type, mad-1(cd), and bub-3Δ mutants (Fig. S2, E and F). For this reason, additional analysis of meiotic prophase events was performed in mad-1(cd) and bub-3Δ mutants.

To test whether MAD-1 and BUB-3 negatively regulate synapsis, we assayed SC assembly by staining for HTP-3, an axial element protein that assembles on chromosomes before synapsis (MacQueen et al., 2005), and SYP-1, a central element component whose addition to the SC is concomitant with synapsis (MacQueen et al., 2002). Extensive stretches of HTP-3 without SYP-1 indicate the presence of unsynapsed chromosomes (arrows in Fig. 2 C and Fig. S4 A), and colocalization of HTP-3 with SYP-1 indicates complete synapsis (Fig. 2 C and Fig. S4 A). Because nuclei in the germline are arrayed in a spatiotemporal gradient, we divided germlines into six equivalent zones and calculated the percentage of nuclei that had completed synapsis (Fig. 2 D). mad-1(cd) and bub-3Δ mutants accelerated synapsis, exhibiting significantly more nuclei with complete synapsis in zones 2 and 3 than wild-type germlines (Fig. 2, C and D; and Fig. S4 A). Null mad-1 (mad-1Δ) mutants also exhibited normal meiotic entry (Fig. S2, E and F) and similar acceleration of synapsis as bub-3Δ and mad-1(cd) mutants (not depicted). We tested whether accelerated synapsis in mad-1(cd) and bub-3Δ mutants produced nonhomologous synapsis by monitoring pairing at a PC locus (HIM-8; Phillips et al., 2005; Phillips and Dernburg, 2006). In addition, mad-1(cd);bub-3Δ;dhc-1;dlc-1 RNAi mutants resembled bub-3Δ;dhc-1;dlc-1 RNAi mutants with regard to percentage of synapsed germlines (not depicted), indicating that MAD-1 and BUB-3 act in the same pathway. Only 5% of pch-2;dhc-1;dlc-1 RNAi triple mutants exhibited normal synapsis, similar to dhc-1;dlc-1 RNAi double mutants (Fig. 3 B), suggesting that PCH-2’s effect on regulating synapsis is independent of the role that dynein and SAC components play in this process.

Similar to a meiosis-specific mutation of sun-1, sun-1(j18) (Sato et al., 2009), mad-1(cd), mad-2Δ, and bub-3Δ mutations suppressed defects in synapsis when only dhc-1 was knocked down by RNAi (Fig. 3, A and D). Loss of pch-2 also did not suppress the asynapsis phenotype of dhc-1 RNAi (Fig. 3 D). Furthermore, mad-3Δ;dlc-1 RNAi worms had extensive asynapsis (Fig. 3 D), consistent with our finding that mad-3 is not required for the synapsis checkpoint (Fig. 1, B and C).

To further test the involvement of the APC in synapsis, we assessed synapsis in mat-3;dhc-1;dlc-1 RNAi mutants, as well as mat-3;mad-1Δ;dhc-1;dlc-1 RNAi and mat-3;mad-2Δ;dhc-1;dlc-1 RNAi mutants. Mutation of mat-3 had no effect on the percentage of germlines with synapsed chromosomes (Fig. 3 C). These data indicate that when specific SAC proteins are absent, the reliance on dynein to license synapsis is lost and this is independent of the APC.

MAD-1 and BUB-3 regulate synapsis by a mechanism redundant with PCH-2. Our experiments in meDf2/+ (Fig. 2, E and F) and dynein mutants (Fig. 3, C and D) suggest that SAC components and PCH-2 regulate synapsis by independent mechanisms. If so, loss of both of these mechanisms should affect synapsis more
severely than loss of only one. First, we verified that meiotic entry in pch-2;mad-1(cd) and pch-2;bub-3 Δ double mutants was unaffected (Fig. S2 F). We then assessed synapsis in pch-2;mad-1(cd) and pch-2;bub-3 Δ double mutants and detected a similar acceleration of synapsis as single mutants, indicating that loss of both of these mechanisms does not further hasten synapsis (Fig. 4 A, zones 2 and 3). However, these double mutants, particularly pch-2;bub-3 Δ, had significantly more nuclei with unsynapsed chromosomes throughout the germline (Fig. 4, A [zones 4 and 5] and B; and Fig. S4 C). Therefore, loss of both PCH-2 and SAC components produces defects in synapsis that are more severe than any of the single mutants, indicating that the regulation of synapsis by SAC components and PCH-2 are distinct, biologically parallel processes.

The loss of both SAC components and PCH-2 does not result in nonhomologous synapsis (not depicted), suggesting that the mechanisms controlling synapsis are distinct from those that assess homology between chromosomes. Given that synapsis initiation, not homologous interactions, is the rate-limiting step for synapsis (Rog and Dernburg, 2015) and SC assembly on meiotic chromosomes is highly processive, even when confronted with extensive regions of nonhomologous sequence (MacQueen et al., 2005), it seems likely that synapsis between homologous PCs must overcome multiple barriers, such as those enforced by PCH-2 and SAC components. Why these multiple barriers exist and how they contribute to accurate meiotic chromosome segregation are currently unknown.

We previously proposed that highly stable PC pairing, whether accomplished normally through synapsis or via the inappropriate stabilization of paired PCs, as in pch-2 mutants, satisfies the synapsis checkpoint (Deshong et al., 2014). Our analysis of mad-1, mad-2, and bub-3 mutants introduces another layer of complexity to the mechanisms that control synapsis: tension, potentially generated by stably paired PCs, may be translated into a molecular signal that silences the checkpoint and initiates synapsis. Because we cannot cytologically detect SAC components at PCs (not depicted), an alternate interpretation is that SAC components perform some other role at the nuclear envelope that affects the checkpoint and synapsis. However, our data support a model in which once stable PC pairing has generated sufficient tension to resist the pulling forces of the microtubule motor dynein, SAC proteins are either inactivated or removed from PCs to initiate synapsis and the synapsis checkpoint is silenced (Fig. 5).

**Materials and methods**

**Genetics and worm strains**

The wild-type *C. elegans* strain background was Bristol N2 (Brenner, 1974). All experiments were performed on adult hermaphrodites at 20°C under standard conditions unless otherwise stated. Mutations and rearrangements used were as follows: LG I: *mmDp66*, *dgc-1*(or195), *san-1/mdf-3*(ok1580), *cep-1*(gk138); LG II: *fyz-1*(h1983), *mat-3*(cd), *mad-1*, *mad-2*, *bub-3*, *sun-1*; and LG III: *fzy-1*(h1983), *mad-1*, *mad-2*, *bub-3*; and LG IV: *fzymd-3*(ok1580), *cep-1*(h1983). We previously proposed that highly stable PC pairing, whether accomplished normally through synapsis or via the inappropriate stabilization of paired PCs, as in pch-2 mutants, satisfies the synapsis checkpoint (Deshong et al., 2014). Our analysis of mad-1, mad-2, and bub-3 mutants introduces another layer of complexity to the mechanisms that control synapsis: tension, potentially generated by stably paired PCs, may be translated into a molecular signal that silences the checkpoint and initiates synapsis. Because we cannot cytologically detect SAC components at PCs (not depicted), an alternate interpretation is that SAC components perform some other role at the nuclear envelope that affects the checkpoint and synapsis. However, our data support a model in which once stable PC pairing has generated sufficient tension to resist the pulling forces of the microtubule motor dynein, SAC proteins are either inactivated or removed from PCs to initiate synapsis and the synapsis checkpoint is silenced (Fig. 5).
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Figure 4. Loss of both PCH-2 and MAD-1 or BUB-3 results in more severe defects in synapsis. (A) pch-2;mad-1(cd) and pch-2;bub-3Δ double mutants exhibit more severe synapsis defects than pch-2, mad-1(cd), and bub-3Δ single mutants (mad-1(cd) and bub-3Δ data from Fig. 2 D). Error bars represent 95% confidence intervals. *, P < 0.01; **, P < 0.0001. (B) Images of nuclei in wild-type worms and pch-2, mad-1(cd), bub-3Δ, pch-2;mad-1(cd), and pch-2;bub-3Δ mutants stained to visualize SYP-1 and HTP-3. Arrows indicate unsynapsed chromosomes. Bar, 5 µm.

Figure 5. Model for synapsis initiation and checkpoint satisfaction in C. elegans. A pair of chromosomes with PCs interact with proteins at the nuclear envelope, including SUN-1 and ZYG-12, to gain access to the cytoplasmic microtubule network and dynein. SAC components are presumed to function at PCs despite our inability to colocalize them. When a chromosome encounters another chromosome, homology is assessed by unknown mechanisms. If chromosomes are homologous and remain stably paired, they resist the pulling forces of the microtubule motor dynein, generating tension (black arrows between PCs) that is monitored by SAC components. Once sufficient tension has been generated (YES!), SAC components are removed, synapsis is initiated, and the checkpoint is silenced. If chromosomes are not homologous, they cannot resist the pulling forces of dynein, are pulled apart, and do not generate tension (NO!). Unsynapsed PCs initiate the synapsis checkpoint response. If unsynapsed chromosomes persist, these nuclei are removed by apoptosis. mad-2 is a terminal deficiency of the left end of the X chromosome that removes the X chromosome PC as well as numerous essential genes (Villeneuve, 1994). For this reason, homo- and hemizygous meDf2 animals also carry a duplication (mnDp66) that includes these essential genes but does not interfere with normal X chromosome segregation (Herman and Kari, 1989) or the synapsis checkpoint (Bhalla and Dernburg, 2005). For clarity, it has been omitted from the text. Because the mad-2 gene is closely linked to spo-11, we used cep-1 to prevent DNA damage checkpoint–induced apoptosis in mad2Δ mutants.

Quantification of germline apoptosis

Scoring of germline apoptosis was performed as previously described in Bhalla and Dernburg (2005). L4 hermaphrodites were allowed to age for 22 h at 20°C, except for the mat-3 temperature-sensitive mutation, which was aged for 18 h at the restrictive temperature of 25°C. We verified that MAT-3 function had been reduced by the increase in the number of germline nuclei positive for phosphorylation of histone H3 serine 10 (Golden et al., 2000). Because nuclei progress from mitosis to meiosis as they travel through the germline, this incubation period guaranteed that early meiotic prophase nuclei, but not mitotic nuclei at the start of the germline, had sufficient time to progress to where checkpoint-induced apoptosis occurs in late meiotic prophase (Jaramil-lo-Lambert et al., 2007). Live worms were mounted under coverslips on 1.5% agarose pads containing 0.2 mM levamisole for wild-type moving strains or 0.1 mM levamisole for dpy-11 strains. A minimum of 25 germlines were analyzed for each genotype by performing live fluorescence microscopy and counting the number of cells fully surrounded by CED-1::GFP. Significance was assessed using a paired t test. All experiments were performed at least twice.
Antibodies, immunostaining, fluorescence in situ hybridization, and microscopy

Immunostaining was performed on worms 20–24 h after L4 stage. Gonad dissections were performed in 1× EBT (250 mM Hepes-Cl, pH 7.4, 1.18 M NaCl, 480 mM KCl, 20 mM EDTA, and 5 mM EGTA) + 0.1% Tween 20 and 20 mM sodium azide. An equal volume of 2% formaldehyde in EBT (final concentration was 1% formaldehyde) was added and allowed to incubate under a coverslip for 5 min. The sample was mounted on Histobond slides (75 × 25 × 1 mm from Lamb), freeze-cracked, and incubated in methanol at −20°C for slightly more than 1 min and transferred to PBST (PBS with Tween 20). After several washes of PBST, the samples were incubated for 30 min in 1% bovine serum albumin diluted in PBST. A hand-cut paraffin square was used to cover the tissue with 50 µl of antibody solution. Incubation was conducted in a humid chamber overnight at 4°C. Slides were rinsed in PBST and then incubated for 2 h at room temperature with fluorescence-conjugated secondary antibody at a dilution of 1:500. Samples were rinsed several times and DAPI stained in PBST, then mounted in 13 µl of mounting media (20 µM N-propyl gallate [Sigma-Aldrich] and 0.14 M Tris in glycerol) with a no. 1 1/2 (22 mm²) coverslip, and sealed with nail polish.

Primary antibodies were as follows (dilutions are indicated in parentheses): rabbit anti-SYP-1 (1:500; MacQueen et al., 2002), chicken anti-HTP-3 (1:250; MacQueen et al., 2005), rabbit anti-MAD-2 (1:10,000; Essex et al., 2009), mouse anti-NPC MAb414 (1:5,000; Covance; Davis and Blobel, 1986), guinea pig anti-HIM-8 (1:250; Phillips et al., 2005), rat anti-HIM-8, guinea pig anti-ZIM-2 (1:2,500; Phillips and Dernburg, 2006), guinea pig anti-SUN-1pSer8 (1:500; Penkner et al., 2009), and goat anti-GFP (1:10,000; Hua et al., 2009). Secondary antibodies were Cy3 anti-mouse, anti-rabbit, anti-guinea pig, anti-rat, and anti-chicken (Jackson ImmunoResearch Laboratories, Inc.) and Alexa Fluor 488 anti-goat, anti-guinea pig, and anti-rabbit (Invitrogen). Antibodies against SYP-1 were provided by A. Villeneuve (Stanford University, Palo Alto, CA). Antibodies against HTP-3, HIM-8, and ZIM-2 were provided by A. Dernburg (University of California, Berkeley/E.O. Lawrence Berkeley National Lab, Berkeley, CA). Antibodies against MAD-2 were provided by A. Desai (Ludwig Institute/University of California, San Diego, La Jolla, CA). Antibodies against SUN-1pSer8 were provided by V. Jantsch (Max F. Perutz Laboratories, Vienna, Austria). Antibodies against GFP were provided by S. Strome (University of California, Santa Cruz, Santa Cruz, CA).

Fluorescence in situ hybridization was performed as described in Phillips et al. (2005). 5S rDNA probe was generated using genomic DNA as a template by PCR and gel purified. The PCR product was digested with the TasI restriction enzyme and ethanol precipitated. 10 µg of digested DNA was diluted into 50 µl water, denatured for 2 min at 95°C, chilled on ice, and spun briefly. At room temperature, 20 µl Roche 5x Taq reaction buffer (Tris-HCl, pH 7.2, potassium cacodylate, and BSA), 20 µl of 25 mM CoCl2 solution, 3.3 ml of 1 mM aa-dUTP, 6.6 ml of 1 mM unlabelled dTTP, and 2 µl (800 U) recombinant terminal deoxynucleotidyl transferase were added. This solution was incubated for 1 h at 37°C. EDTA was added to 5 mM, and the DNA was ethanol precipitated. The probe was conjugated with Cy3 dye (Life Technologies) by adding 5 µl of 1 mg/ml resuspended probe and 3 µl of 1 M bicarbonate buffer to one aliquot of dry dye. The reaction was mixed, shielded from light, and incubated for 1 h at room temperature and ethanol precipitated.

Worms were dissected 24 h after L4 stage in 30 µl EBT (1× egg buffer, 0.1% Tween 20, and 20 mM sodium azide). We added 30 µl of 1× egg buffer and 0.5% EGS (ethylene glycol bis[succinimidylsuccinate] in dimethyl formamide) and pipetted to extrude gonads. 30 µl of this liquid was removed, and the sample was mounted on Histobond slides (75 × 25 × 1 mm from Lamb) and allowed to incubate in a humid chamber for 30 min at room temperature. Samples were freeze-cracked and incubated in methanol at −20°C for slightly more than 1 min and transferred to 2× SSCT (2× SSC and 0.1× Tween 20) at room temperature. Samples were placed in 3.7% formaldehyde in 1× egg buffer for 5 min, rinsed briefly in 2× SSCT, and washed twice in 2× SSCT for 5 min. The samples were incubated in 50% formamide in 2× SSCT for 5 min, transferred to fresh 50% formamide in 2× SSCT, and incubated at 37°C overnight. Samples were cooled to room temperature, and 20 ng of 5S rDNA probe in hybridization solution (50% formamide, 3% SSC, 10% dextran sulphate) was added and sealed onto the sample with a coverslip and nail polish. Slides were denatured on a hot block at 95°C for 3 min and placed in a humid chamber at 37°C overnight. Coverslips were removed, and the samples were washed twice in 50% formamide in 2× SSCT for 30 min each. Samples were rinsed several times and DAPI stained in 2× SSCT. Samples were mounted in 13 µl of mounting media (20 M N-propyl gallate [Sigma-Aldrich] and 0.14 M Tris in glycerol) with a no. 1 1/2 (22 mm²) coverslip and sealed with nail polish.

Quantification of synthesis and pairing was performed with a minimum of three whole germlines per genotype as in Phillips et al. (2005) on animals 24 h after L4 stage. The gonads were divided into six equal-sized regions, beginning at the distal tip of the gonad and progressing through the end of pachytenes. Significance was assessed by performing Fisher’s exact test.

Quantification of rows of mitotic nuclei was performed as in Stevens et al. (2013), and a minimum of 18 germlines were analyzed on animals 24 h after L4 stage. Significance was assessed by performing a paired t test.

Quantification of meiotic progression was performed with a minimum of three whole germlines per genotype on animals 24 h after L4 stage by quantifying the percentage of nuclei with clustered chromosomes. Significance was assessed by performing Fisher’s exact test.

All images were acquired at room temperature using a DeltaVision Personal DV system (GE Healthcare) equipped with a 100× NA 1.40 oil immersion objective (Olympus), resulting in an effective xy pixel spacing of 0.064 or 0.040 µm. Images were captured using a charge-coupled device camera (Cool-SCAN HQ; Photometrics). Three-dimensional image stacks were collected at 0.2-µm z-spacing and processed by constrained, iterative deconvolution. Imaging, image scaling, and analysis were performed using functions in the softWoRx software package. Projections were calculated by a maximum intensity algorithm. Composite images were assembled, and some false coloring was performed with Photoshop software (Adobe).

IPs

Asynchronous liquid worm cultures were grown at 20°C for 4 d in S medium supplemented with concentrated HB101 bacteria, and embryos were extracted in a sodium hypochlorite solution (25% [vol/vol] NaClO and 0.25% [vol/vol] 10N NaOH) and allowed to hatch overnight on unseeded NGM plates. Hatched L1s were washed off NGM plates and grown at 19°C for 66–68 h or until the majority of animals reached adulthood. Adult worms were harvested, washed twice in sterile water and once in buffer H0.15 (50 mM Hepes, pH 8.0, 2 mM MgCl2, 0.1 mM EDTA, pH 8.0, 0.5 mM EGTA-KOH, pH 8.0, 15% glycerol, 0.1% NP-40, and 150 mM KCl), and frozen into “popcorn” by dripping into liquid nitrogen. Popcorn was then pulverized three times for 2 min at 25 Hz in a MM-400 mixer mill (Retsch Technology) with liquid nitrogen immersion between milling sessions. Worms were lysed by adding 5 ml ice-cold buffer H0.15 supplemented with protease and phosphatase inhibitors (0.1 mM AEBSF, 5 mM benzamidine, 1:200 aprotinin, Roche Complete Mini tablets w/o EGTA, 1 mM Na4P2O7, 2 mM Na-b-glycerophosphate, 0.1 mM Na3VO4, and 5 mM...
For immunoblotting, samples were run on SDS-PAGE gels, transferred to nitrocellulose, blocked in a PBST + 5% (wt/vol) non-fat milk solution, and then probed with mouse anti-GFP (1:1,000; Roche), rabbit anti–MAD-1 (1:2,000; Yamamoto et al., 2008), or rabbit anti–MAD-2 (1:5,000; Essex et al., 2009) overnight at 4°C. Blots were washed three times for 10 min in PBST, probed for 1 h using an HRP-conjugated secondary antibody (rabbit or mouse; GE Healthcare), washed three times for 10 min in PBST, and then analyzed using a chemiluminescent substrate (Thermo Fisher Scientific). 0.1% of starting material is shown for all input samples, 10% of the IP elution is shown for anti-GFP Western blots, and 30% of IPs are shown for anti–MAD-1 and anti–MAD-2 Western blots. IP samples were analyzed with Pico chemiluminescent substrate (Thermo Fisher Scientific), and input samples were analyzed using Dura enhanced chemiluminescent substrate (Thermo Fisher Scientific). We estimate IP to have purified between 20 and 30% of SUN-1::GFP present in the input.

Feeding RNAi

For RNAi, dlc-1RNAi and empty vector (L4440) clones from the Ahringer laboratory (Fraser et al., 2000) were used. Bacteria strains containing Caenorhabditis Genetics Center, which is funded by NIH Office of R01GM097144 [to N. Bhalla]. Some strains were provided by the A. Villeneuve, and Verena Jantsch for valuable strains and reagents.

Spindle checkpoint proteins regulate synapsis

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