A novel meiotic protein required for homolog pairing and regulation of synapsis in *C. elegans*

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Interactions between chromosomes and LINC (Linker of Nucleoskeleton and Cytoskeleton) complexes in the nuclear envelope (NE) promote homolog pairing and synapsis during meiosis. By tethering chromosomes to cytoskeletal motors, these connections lead to rapid, processive chromosome movements along the NE. This activity is usually mediated by telomeres, but in the nematode *Caenorhabditis elegans* special chromosome regions called “Pairing Centers” (PCs) have acquired this meiotic function. Through a genetic screen for mutations that cause meiotic nondisjunction, we discovered an uncharacterized meiosis-specific NE protein, MJL-1 (MAJIN-Like-1) that is essential for interactions between PCs and LINC complexes. MJL-1 colocalizes with PCs and LINC complexes during pairing and synapsis. Mutations in MJL-1 disrupt these interactions and eliminate active chromosome movements. *mjl-1* mutants display promiscuous nonhomologous synapsis, reduced clustering of PCs, and severely impaired homolog pairing. MJL-1 likely interacts directly with SUN-1 and DNA-binding proteins to connect PCs to the LINC complex. Similarities in the molecular architecture of chromosome-LINC complex attachments between *C. elegans* and other organisms suggest that these connections may play previously unrecognized roles during meiois across eukaryotes.

Introduction

Sexual reproduction relies on meiosis, the specialized cell division program that produces haploid gametes. During meiosis, homologous chromosomes must pair, synapse, and undergo crossover recombination to segregate accurately. Upon meiotic entry, each replicated chromosome is assembled into an array of loops anchored to a linear structure known as the chromosome axis (Zickler and Kleckner, 1999; Blat et al., 2002). Pairing of homologs is gradually stabilized by assembly of a protein matrix, the synaptonemal complex (SC), between axes (Zickler and Kleckner, 1999; Page and Hawley, 2004; Cahoon and Hawley, 2016). SCs promote and regulate crossover recombination, which results in chiasmata, physical linkages between homologous chromosomes that persist until segregation and mediate bipolar alignment on the spindle (Zickler and Kleckner, 2015; Page and Hawley, 2004; Kleckner 2006).

Chromosome pairing, synapsis, and recombination are promoted by nuclear envelope (NE)-associated chromosome dynamics during meiosis (Conrad et al., 1997; Chua et al., 1997; Cooper et al., 1998; Ding et al., 2004; Ding et al., 2007; Conrad et al., 2008). At meiotic entry, chromosomes become tethered to LINC (Linker of Nucleoskeleton and Cytoskeleton) complexes comprised of SUN and KASH domain proteins that span the two membranes of the NE (Hiraoka and Dernburg, 2009; Link and Jantsch, 2019). Cytoskeletal motors interact with LINC complexes on the cytoplasmic face of the NE, resulting in dramatic chromosome movements during early meiotic prophase (Hiraoka, 1998; Chikashige et al., 1994; Scherthan et al., 1996). This often leads to clustering of chromosome ends near cytoplasmic microtubule organizing centers to form a chromosome configuration known as the “meiotic bouquet” (Zickler and Kleckner, 1998; Scherthan, 2001; Hiraoka, 1998).

In the nematode *Caenorhabditis elegans*, specialized regions on each chromosome known as “Pairing Centers” (PCs) mediate homolog pairing and synapsis (Rog and Dernburg, 2013; Rosenbluth and Baillie 1981; McKim et al., 1988; Villeneuve 1994; MacQueen et al., 2005). Each PC recruits one of four meiosis-specific zinc finger proteins, ZIM-1, ZIM-2, ZIM-3, or HIM-8, through DNA binding sites present in clusters throughout the PC regions (Phillips et al., 2005; Phillips and Dernburg 2006, Phillips et al., 2009). During early meiotic prophase, PCs associate with LINC complexes comprised of SUN-1 and ZYG-12 (Penkner et al., 2007; Sato et al., 2009). ZYG-12 interacts with cytoplasmic dynein and perhaps other microtubule motors to drive processive movement of chromosomes that promote pairing and synapsis (Sato et al., 2009; Penkner et al., 2009; Wynne et al., 2012; Baudrimont et al., 2010).
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In addition to the LINC complexes, additional meiosis-specific inner NE proteins are required for telomere-led chromosome movements in fission yeast and mice (Chikashige et al., 2009; Shibuya et al., 2015). Homologs of the mouse protein MAJIN (Membrane-Anchored Junction Protein) have been identified by sequence homology in many metazoans, but not in nematodes (da Cruz et al., 2020). Here we report the identification and characterization of a novel meiosis-specific NE protein that is essential for homolog pairing and synapsis in C. elegans. Based on the functions we have characterized, we named it MJL-1 (MAJIN-Like-1).

Results

Identification of MJL-1, a meiosis-specific NE protein

In C. elegans, defects in meiosis result in a High incidence of males (Him) phenotype due to nondisjunction of the X chromosome (Hodgkin et al., 1979); most meiotic mutants also produce many inviable embryos due to autosomal aneuploidy. We used a “Green eggs and Him” screen, based on a xol-1p::gfp reporter expressed in XO (male) embryos, to identify hermaphrodites with elevated meiotic nondisjunction (Kelly et al., 2000). Molecular lesions in the mutants were identified by outcrossing to the CB4856 Hawaiian strain, reisolating homozygous mutants, genome sequencing of their progeny, and computational analysis to identify likely causal mutations (Doitsidou et al., 2010; Wicks et al., 2001; Swan et al., 2002). Most of the mutations we identified (50/52) were in genes previously shown to be important for meiosis, indicating that such Him screens are nearing saturation. Two mutations mapped to previously uncharacterized genes. One of these resulted in a premature stop codon in C17E4.4, which encodes a small protein with a single predicted transmembrane domain. Based on previous transcriptome data, C17E4.4 is specifically expressed in germline and arcade cells (Spencer et al., 2011; Han et al., 2017; Grun et al., 2014).

Using Cas9/CRISPR-based editing, we inserted an HA epitope tag at the N-terminus of the C17E4.4 protein. Hermaphrodites homozygous for this insertion produced a normal number of embryos and a slightly elevated frequency of male self-progeny (~1%, compared to 0.2% in wild-type broods). Immunofluorescence of the HA-tagged protein showed NE-specific localization throughout the meiotic region of the germline. The protein was undetectable in proliferating germline stem cells (GSCs) but was clearly observed at the NE upon meiotic entry and concentrated to form NE “patches” in transition zone (leptotene-zygotene) nuclei. Following synapsis, the protein redistributed throughout the NE and persisted until late pachytene (Fig. 1a, 1b).

During pairing and synapsis, the zinc finger proteins HIM-8, ZIM-1, -2, and -3 bind to PCs and interact with the LINC complex proteins SUN-1 and ZYG-12, which concentrate within the NE to form multiple patches. Immunofluorescence revealed that HA-tagged C17E4.4 colocalized with all four PC proteins during this transient stage (Fig. 1c).

Based on structural and functional similarities between C17E4.4 and mouse MAJIN (Shibuya et al., 2015), we named the gene mj-1. Although MJL-1 shares no discernible sequence homology with MAJIN, both are small, single-pass transmembrane proteins with similar meiosis-specific functions (below) (Fig. 2a). MJL-1 is only weakly conserved within the genus Caenorhabditis, and we have not yet identified homologs in other nematode genera, including those that express homologs of the PC proteins (Fig. S1) (Rillo-Bohn et al., 2021).
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MJL-1 is required for association of PCs with LINC complexes

The allele identified in our screen, mjl-1(ie142), is likely a null mutation since it contains a stop codon before its transmembrane domain. We also obtained a deletion allele, mjl-1(tm1651), from the Japanese National BioResource Project (NBRP) (Fig. 2a). Hermaphrodites homozygous for either mutant allele produced very few viable self-progeny (<1%), and among these, many were males (29% and 33%, statistically indistinguishable), indicative of extensive meiotic nondisjunction. At diakinesis, most oocytes in mjl-1(ie142) and mjl-1(tm1651) hermaphrodites displayed 10-12 DAPI-staining bodies (Fig. 2b), indicating that chromosomes failed to undergo crossing-over.

Based on its localization, we hypothesized that MJL-1 might mediate interaction between PCs and LINC complexes. Immunofluorescence confirmed that in mjl-1 mutants, SUN-1 did not colocalize with PC proteins or form NE patches in transition zone nuclei (Fig. 2c). However, HIM-8 still appeared to associate with the NE, suggesting that HIM-8 may interact directly with the membrane or another NE protein (Fig. S2a).

We crossed mjl-1(ie142) mutants to a strain expressing GFP-tagged HIM-8 (Wynne et al., 2012) to analyze chromosome movement. The average speed of HIM-8 foci in early meiotic nuclei was greatly reduced in the absence of MJL-1, from 59.8 nm/s in wild-type oocytes to 22.7 nm/s in mjl-1(ie142) (p<0.0001), similar to our measurements for sun-1(jf18) (18.7 nm/s) (Fig. 2d, 2e). Previous analysis has shown that sun-1(jf18), which results a missense mutation (G311V) in the SUN domain, abrogates active chromosome movement (Penkner et al., 2009; Baudrimont et al., 2010). The residual movement in mjl-1(ie142) and sun-1(jf18) mutants is likely due to diffusion rather than active motility (Wynne et al., 2012; Baudrimont et al., 2010; Woglar et al., 2013).

MJL-1 is required to regulate synopsis

We observed extensive SC assembly despite very low levels of homolog pairing in mjl-1 mutant nuclei (Fig. 3a, 3b). Similar promiscuous nonhomologous synopsis is observed in sun-1(jf18) mutants (Penkner et al., 2007; Sato et al., 2009). This effect of MJL-1 loss-of-function is consistent with prior

Figure 2. Loss of MJL-1 disrupts PC function.
(a) Diagram of the mjl-1 gene, indicating the mutations described in this work (top). Primary structure of MJL-1 in C. elegans and MAJIN in M. musculus (bottom) (TM: Transmembrane; DB: DNA binding domain).
(b) Loss of MJL-1 disrupts the connection between PC proteins and LINC complex. Transition zone nuclei were stained with antibodies against HIM-8 (green) and SUN-1 (magenta). Scale bars, 5 μm.
(c) Average speed of GFP::HIM-8 foci in transition zone nuclei in wild-type, mjl-1(ie142) (p<0.0001), and sun-1(jf18) (p<0.0001) hermaphrodites. Each point represents a single nucleus. p-values were computed using Student’s t-test with Bonferroni post-hoc correction.
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evidence that the interaction between PCs and LINC complexes regulates synapsis so that it occurs only between homologous chromosomes. Nonhomologous synapsis in mj-l-1 and sun-1(jf18) mutants is more extensive than that caused by disruption of ZYG-12 or dynein (Sato et al., 2009; Zhang et al., 2015), suggesting a more direct role for these proteins in regulating synapsis initiation.

MJL-1 depends on SUN-1 for its NE localization

Unexpectedly, MJL-1 was not detected by immunofluorescence in sun-1(ok1282) null mutants or following auxin-induced degradation of SUN-1 (Fig. 4a). The abundance of MJL-1 detected on western blots was also strongly reduced following auxin-induced degradation of SUN-1 (Liu et al., 2021) (Fig. S2b). These findings indicate that SUN-1 is required for subcellular localization and/or stabilization of MJL-1. In contrast, neither the sun-1(jf18) missense mutation nor auxin-induced degradation of the KASH domain protein ZYG-12 (Liu et al., 2021) (Fig. 4a), suggesting that ZYG-12 is not required for association between MJL-1 and SUN-1.

Our screen also identified a novel meiosis-specific separation-of-function mutation in sun-1 that resulted in phenotypes similar to sun-1(jf18). This new missense mutation, sun-1(ie143), changes tyrosine 128 within the predicted transmembrane domain of SUN-1, very close to the perinuclear domain, to phenylalanine (Y128F). In contrast to sun-1(jf18), sun-1(ie143) resulted in loss of MJL-1 protein in the germline, indicating that it disrupts the interaction between MJL-1 and SUN-1 (Fig. 4b, 4c), and thus suggesting that these proteins interact through their transmembrane and/or perinuclear domains. These regions in MJL-1 are conserved within Caenorhabditis (Fig. S3).

MJL-1 was also detected in apoptotic nuclei in the loop region of the germline. These nuclei retain SUN-1 at their NE but not ZYG-12 (Fig. S4), which may reflect disruption of the outer nuclear membrane during apoptosis, although this has not been directly demonstrated. The persistence of MJL-1 in these nuclei, together with our evidence that the protein requires SUN-1 for its localization to the NE (above) and may interact directly with PCs (below), indicates that MJL-1 probably resides within the inner nuclear membrane with its N-terminal domain in the nucleoplasm, similar to mouse MAJIN (Shibuya et al., 2015).

Interactions between MJL-1 and PC proteins

Sequence alignment of Caenorhabditis MJL-1 homologs revealed a short region of relatively high conservation enriched for acidic residues (Fig. 5a). Using two crRNAs flanking this region, we generated an in-frame deletion of amino acids 34-49. The resulting MJL-1Δacidic protein was expressed and localized to the NE but did not interact with PCs (Fig. 5a). Together with evidence that SUN-1 is essential for localization and stability of MJL-1 (above), this suggests that the mutant protein retains the ability to interact with SUN-1, but not with PC proteins. Homozygotes showed extensive nonhomologous synapsis, similar to mj-l-1 null mutants (Fig. S5). In contrast, an in-frame deletion of the sequence encoding amino acids 9-26, which are also relatively well conserved within Caenorhabditis, led to no apparent defects (data not shown).
The Polo-like kinase PLK-2 is recruited to PCs through Polo-box binding motifs in the zinc finger proteins and is required for interaction between PCs and the LINC complexes (Harper et al., 2011, Labella et al., 2011). Therefore, we asked whether PLK-2 activity at PCs is essential for the association of PC proteins with MJL-1. In a kinase-dead \textit{plk-2}(K65M) mutant (Link et al., 2018; Brandt et al., 2020), MJL-1 localized throughout the NE and did not associate with PC proteins, suggesting that PLK-2 activity is required for the interaction between MJL-1 and PC proteins, but not for association between MJL-1 and SUN-1 (Fig. 5b), which is important for MJL-1 localization. Recruitment of PLK-2 to PCs requires phosphorylation of PC proteins by CHK-2 (Kim et al., 2015). We found that auxin-induced degradation of CHK-2 also abrogated the association between MJL-1 and PCs (Fig. 5c).

We examined the association of MJL-1 with two mutant versions of HIM-8, the PC protein specific for the X chromosome. HIM-8\textit{T64A}, which does not recruit PLK-2 due to a point mutation in its Polo-box binding motif (Harper et al., 2011), and HIM-8\textit{S85F} (encoded by \textit{him-8}(me4) allele), which fails to recruit both CHK-2 and PLK-2 (Kim et al., 2015), did not associate with patches of MJL-1 (Fig. 5c). Thus, PLK-2 activity at PCs is required for the interaction between PC proteins and MJL-1. We did not detect interactions between the putative nucleoplasmic domain of MJL-1 and PC proteins in a yeast two-hybrid assay (data not shown) but this may reflect a requirement for meiosis-specific post-translational modifications of these proteins.

**MJL-1 promotes homolog pairing**

Inhibition of SC assembly prevents nonhomologous synapsis and partially restores X chromosome pairing in \textit{sun-1}(f18) mutants (Sato et al., 2009). To assess the role of MJL-1 in pairing, we compared the extent of X chromosome pairing in \textit{zyg-12::aid} following auxin treatment, \textit{sun-1}(f18), and \textit{mjl-1}(ie142), all in the absence of synapsis. Loss of MJL-1 resulted in more severe pairing defects than the absence of ZYG-12 or the \textit{sun-1}(f18) mutation, indicating that connection of chromosomes to MJL-1 and LINC complexes facilitates pairing even in the absence of rapid chromosome movements, although some “baseline” pairing (~25%) was still detected in mutants lacking MJL-1 (Fig. 6a, 6b). We also observed a reduction in nonhomologous associations between HIM-8 and other PCs in the absence of MJL-1 compared to \textit{sun-1}(f18) mutants, suggesting that MJL-1 promotes clustering of PCs.
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Conservation activity at PCs contributes to pairing through a lamin-between LMN-1 proteins (Link et al., 2018). We found that along the NE, likely by reducing homotypic interactions the nuclear lamina by PLK-2 promotes chromosome mobility chromosome pairing in the absence of synapsis. This indicates C.briggsae C.remanei of HIM-8 with the MJL-1 (magenta). Scale bar, 5 μm.

**Figure 5.** MJL-1 requires a small domain enriched in acidic amino acids to interact with PC proteins. (a) Sequence alignment of the N-terminal region of MJL-1 homologs within Caenorhabditis, generated using MAFFT. (b) Maximum intensity projection images showing zone nuclei stained with antibodies against HIM-8 (green in merged images) and HA (magenta) from hermaphrodites expressing HA::MJL-1WT (top) or HA::MJL-1K65M (bottom). Scale bar, 5 μm. (c) PLK-2 activity is required for interaction between MJL-1 (magenta) and PC proteins (green). Scale bar, 5 μm. (d) Recruitment of PLK-2 by HIM-8 (green) is required for the association of HIM-8 with the MJL-1 (magenta). Scale bar, 5 μm.

(Fig. 6c, 6d), which may be required for efficient pairing of homologous PCs (Fig. 6e).

**PLK-2 activity is required for homolog pairing**

In contrast to mjl-1 mutants, which showed a low level of X-chromosome pairing, hermaphrodites expressing kinase-dead PLK-2K65M or HIM-8T64A displayed almost no detectable X chromosome pairing in the absence of synapsis. This indicates that PLK-2 contributes to pairing beyond tethering PCs to MJL-1 during homolog pairing (Fig. 6a, 6b). Phosphorylation of the nuclear lamina by PLK-2 promotes chromosome mobility along the NE, likely by reducing homotypic interactions between LMN-1 proteins (Link et al., 2018). We found that depletion of LMN-1 did not rescue X chromosome pairing in syp-3(ok758); him-8(T64A) mutant, indicating that PLK-2 activity at PCs contributes to pairing through a lamin-independent mechanism (Fig. S6). In SC-deficient mutants, paired HIM-8 foci dissociated during late prophase, concomitant with the disappearance of PLK-2, indicating that PLK-2 activity increases affinity between homologous PCs (Fig. S7).

**Discussion**

**Similarities and differences between MJL-1 and MAJIN**

Homologs of MJL-1 are only detected within Caenorhabditis. PC proteins have been detected in related nematode genera (Rillo-Bohn et al., 2021), but show rapid divergence, particularly in their N-terminal domains, which act as scaffolds to recruit kinases and may also directly interact with NE proteins. If PC proteins interact with MJL-1, as suggested by our findings, rapid coevolution of these proteins may account for our inability to detect MJL-1 homologs in other nematode species.

Fission yeast Bqt3 and Bqt4 were the first NE proteins shown to be required for tethering of telomeres to LINC complexes during meiosis (Chikashige et al., 2009). MAJIN is likely to play a similar role in most metazoans (Shibuya et al., 2015; da Cruz et al., 2020). Bqt4, MAJIN, and MJL-1 share a similar structure, with a single transmembrane domain near their C termini, and may represent orthologous proteins despite their lack of sequence conservation (Hu et al., 2019; Shibuya et al., 2015).
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**Figure 6.** MJL-1 promotes pairing even in the absence of chromosome movements.  
(a) Blocking synapsis does not restore pairing of HIM-8 in *mjl-1(e142)* mutants, in contrast to *sun-1(j18)* or depletion of ZYG-12::AID. AID::SYP-3 was depleted by treatment with auxin for 24 h. Nuclei display polarized morphology due to lack of synapsis. Scale bar, 5 μm.  
(b) Quantification of X chromosome pairing. The extended transition zone was divided into three equal regions (zones 2-4) by length (zone 1: pre-meiotic). Each point represents a single gonad. *p*-values were calculated by one-way ANOVA with pairwise post-hoc Bonferroni correction (** *p* < 0.01; *** *p* < 0.001; **** *p* < 0.0001).  
(c) In the absence of synapsis, clustering of HIM-8 with other PC proteins is lower in *mjl-1(e142)* than in *sun-1(j18)*. Scale bar, 5 μm.  
(d) Quantification of clustering between HIM-8 and other PC proteins in various mutants. Only nuclei in zone 2 were analyzed, since pHIM-8/ZIM staining decreases in zone 3-4. *p*-values were calculated by one-way ANOVA with post-hoc pairwise Bonferroni correction (* *p* < 0.05; ** *p* < 0.01; *** *p* < 0.001; **** *p* < 0.0001)
Our evidence indicates that MJL-1 connects PC proteins to LINC complexes in *C. elegans*. Similarly, in mouse spermatocytes, MAJIN connects the meiosis-specific shelterin-binding proteins TERB1 and TERB2 to LINC complexes (Wang et al., 2020). MJL-1 requires PLK-2 activity to interact with PC proteins, while CDK2 activity is required for interaction between MAJIN and SUN1 in mice (Wang et al., 2020; Mikolcevic et al., 2016; Tu et al., 2017). In fission yeast, a direct interaction between NE proteins Bqt3 and -4 and LINC complexes has not been detected.

MAJIN and Bqt4 contain N-terminal DNA-binding motifs that are required for recruitment of telomeres to the NE (Hu et al., 2019; Shibuya et al., 2015). The DNA binding motif of MAJIN has been implicated in “telomere cap exchange,” whereby telomeres release shelterin and directly interact with TERB1, -2 and MAJIN (Shibuya et al., 2015). In contrast, MJL-1 lacks an apparent DNA-binding motif, and PCs in *C. elegans* associate with the NE even in the absence of MJL-1 or SUN-1.

**Roles of MJL-1 and LINC complexes in regulation of pairing and synopsis**

Association of PCs with MJL-1 and LINC complexes inhibits inappropriate synopsis in *C. elegans* (Penkner et al., 2007; Sato et al., 2009). This is consistent with observations that these associations and resulting chromosome movements persist after pairing of homologs, which occurs soon after entry into meiosis. In contrast, loss of SUN-1 or MAJIN does not result in inappropriate synopsis in mice (Ding et al., 2007; Shibuya et al., 2015; Wang et al., 2019), which suggests that inhibition of nonhomologous synopsis by MJL-1 and LINC complexes may be unique to *C. elegans*. Nevertheless, CDK2, which inhibits inappropriate synopsis in mice, is also recruited to LINC complexes (Mikolcevic et al., 2016; Viera et al., 2009; Tu et al., 2017; Chen et al., 2021), suggesting that regulation of synopsis may be a general role of chromosome-LINC complex attachments, despite some differences in the details of this regulation between organisms.

Intriguingly, telomeric attachments in mammals and PC attachments in *C. elegans* each require a kinase (CDK2 and PLK-2, respectively) that is also involved in crossover regulation, suggesting that coordination between chromosome attachments sites and CO intermediates may be a conserved feature of meiosis (Woglar and Villeneuve, 2018; Zhang et al., 2021; Ashley and Rooij, 2001; Palmer et al., 2020).

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Author contributions

Conceptualization: HJK, AFD
Methodology: HJK, AFD
Investigation: HJK
Supervision: AFD
Writing—original draft: HJK, AFD

Competing interest statement

The authors declare no competing financial interests.

Materials and Methods

C. elegans strains

N2 Bristol was used as the wild-type C. elegans strain; all mutations described here were generated in this background. The Hawaiian isolate CB4856 was used for genetic mapping. All strains were maintained at 20°C under standard laboratory conditions. The following mutations and balancers were used: mjll-1[ie142], mjll-1(tm1651), sun-1(jf18), sun-1(ok1282), syp-2(ok307), syp-3(ok758), him-8(me4), plk-2(K65M):3xflag (Brandt et al., 2020), nt1[qIs51] (IV;V), hT2[bli-4(e937) let-?(q782) qIs48] (I;III). The following constructs were used for auxin-inducible degradation: zyg-12::ha::aids::v5::sun-1 (Liu et al., 2021), ha::aids::chk-2 (Zhang et al., 2021) where “aids” designates a 44-amino acid degron sequence (Zhang et al., 2015).

To generate ha::mjll-1 strains, single-stranded (ss) DNA templates were designed to insert one or two copies of the HA tag at the N-terminus of MJL-1, separated by a flexible linker (GGGGS). These were co-injected with Cas9-NLS prebound to duplexed gRNAs, as well as a gRNA and ssDNA template for co-CRISPR of dpy-10 (Arribere et al., 2014). (Final concentrations: dpy-10 crRNA, 20 µM; mjll-1 crRNA, 50 µM; trRNA, 40 µM; Cas9-NLS protein, 20 µM; dpy-10 repair template, 1 µM, mjll-1 repair template, 1 µM). To label MJL-1 using the split-GFP system and V5 tag, a template to insert GFP11 and V5 was co-injected with the Cas9-gRNA RNP complex into DUP223 glhl-1[sa129][glhl-1:2x:a:sGF2P2(1-10)] (Goudeau et al., 2021). Essentially the same procedure was used to generate in-frame deletions in mjll-1, except that two gRNAs were used.

Auxin-induced degradation

A stock solution containing 250 mM indole acetic acid (IAA, auxin) in EtOH was diluted to 1 mM in NGM agar just before pouring plates. After drying overnight, plates were seeded with E. coli. OP50 freshly cultured overnight to saturation in 50mL of LB was pelleted by centrifugation at 3,000xg for 5 min and resuspended in 500 µl of M9 buffer containing 1 mM auxin. This concentrated bacteria + auxin was spread on the plates and allowed to grow at room temperature for 1-2 days. To deplete degron-tagged proteins, young adult animals aged 24-48 h from L4 were picked onto these plates and analyzed 4-24 h later.

RNA interference

Carbenicillin and IPTG were added to NGM agar to 200 µg/mL and 1 mM final just before pouring plates, respectively. Clones from the Ahringer laboratory (Fraser et al., 2000) were freshly cultured overnight to saturation in 10mL of LB containing 200 µg/mL carbenicillin. Then the culture was pelleted by centrifugation at 3,000xg for 5 min and resuspended in 50 µl of M9 buffer. Concentrated E. coli was spread on the plates and allowed to grow at 37°C for 1 day. For feeding, young adult animals aged 24-48 h from L4 were picked onto these plates and analyzed 24-48 h later. LMN-1 depletion was confirmed by shrunken nuclei in late pachytene-diplotene.

Cytological Methods

Immunofluorescence and in situ hybridization were performed essentially as described previously (Dernburg et al., 1998). In brief: young adult worms were cut with a scalpel blade in egg buffer containing 0.05% tetramisole to release their gonads on slides, fixed in 1% formaldehyde in egg buffer for 2 min, transferred to tubes, and incubated with methanol pre-chilled to ~30°C for 5 min. The tissue was then washed 3x in PBS containing 0.1% Tween 20 (PBST) at room temperature. Tissues were blocked using 1x Roche Blocking Reagent in PBST for 20 min. Primary antibodies were diluted into the same blocking solution and incubated with the tissues overnight at 4°C. Secondary antibodies were prepared in the blocking solution (1:200), mixed with samples, and incubated 1-2 h at room temperature. Samples were mounted in Prolong Diamond mounting medium containing DAPI (Invitrogen).

For fluorescence in situ hybridization, dissected gonads were fixed in 2% formaldehyde in egg buffer for 5 min, incubated with methanol pre-chilled to -30°C for 5 min, and washed 3x in 2x SSC containing 0.1% Tween 20 (2xSSCT) at room temperature. The tissue was then incubated in 50% formamide in 2x SSCT overnight at 37°C. 0.5-1 µl of 100µM fluorophore-conjugated oligonucleotide probes (IDT) “lV-2” (Adilardi and Dernburg, 2022) were added to 100 µl of hybridization buffer (50% formamide and 10% dextran sulfate in 2x SSCT) and tissues were moved into this mix, incubated 2-3 min at 91°C and then overnight at 37°C. The tissues were washed 3x in 2x SSCT at room temperature and mounted as for immunofluorescence (above).

Images were acquired using a DeltaVision Elite wide-field microscope system (GE) with a 100X 1.40 or 1.45 NA oil-immersion objective, or CSU-W1 SoRa confocal microscope system equipped with a 100X, 1.49 NA oil-immersion objective (Intelligent Imaging Innovations, Inc. [3i]). Deconvolution, projection, and analysis were performed using the softWoRx package or Slidebook 6.3.

Antibodies

All antibodies used in this study were obtained from commercial sources or have been previously described. They include the following antibodies and dilutions: mouse monoclonal anti-HA (Invitrogen 2-22.2.14) (1:250), mouse monoclonal anti-V5 (Invitrogen 46-0705) (1:250), and rabbit polyclonal anti-V5 (Sigma V8137) (1:250). Custom polyclonal antibodies included rabbit anti-SUN-1 (1:250) (Sato et al., 2009), rat anti-HIM-8 (1:500) (Phillips et al., 2005), rabbit anti-SYP-2
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(1:500) Colaiácovo et al., 2003), rabbit anti-phospho-HIM-8/ZIMs
(1:1000) Kim et al., 2015). Fluorophore-conjugated secondary
antibodies were purchased from Jackson ImmunoResearch and used
at 1:200 dilution.

Quantification of homolog pairing

3D distances between HIM-8 or FISH signals were measured using
the “Measure Distance” tool in softWoRx. Foci closer than 0.6 µm were
considered to be paired. We defined this threshold based on the
maximum width of PC protein patches associated with paired
chromosomes in wild-type oocytes. For analysis of pairing of FISH
signals, we included only nuclei displaying two clear foci.

In vivo imaging and quantification of chromosome movement

3D confocal image acquisition was performed essentially as
described (Wynne et al., 2012) using Marianas spinning-disc confocal
microscope system equipped with a 100X, 1.46 NA oil-immersion
objective (Intelligent Imaging Innovations, Inc. [3i]). Exposure time was
set to 100-150 ms depending on the brightness of foci. Stacks of 10–
20 optical sections at 0.5 µm z-spacing were acquired every 5 s for a
total of 5-10 min. 3D time-lapse images were analyzed using Imaris
9.2.0 (Bitplane). Background drift was corrected using the “Reference
Frame” tool. Foci were detected using the “Spots” tool with an
estimated XY diameter of 1.33 µm and filtered with “Quality” and
“Intensity Sum”. Tracks were obtained with max distance of 1.75 µm
and max gap of 3. Tracks from the background noise were manually
removed. p-values were calculated using the pairwise t-test with post-
hoc Bonferroni correction.

Western blots

200 young adult animals aged 48 h from the L4 stage were picked
into M9 buffer, washed three times, and then incubated in 1x SDS
sample buffer at 50°C for 10 min. Samples were vortexed for 2–3 min
until no visible solid material remained. Proteins were separated using
SDS-PAGE gradient gels (Invitrogen NuPAGE™ 4-12%, Bis-Tris, 1.0
mm, 10-well) and transferred to Amersham Protran Nitrocellulose
Membranes. The membrane was cut into slices and probed with mouse
anti-HA (Invitrogen 2-2.2.14) (1:1,000) and mouse anti-α-tubulin (Sigma
DM1A) (1:2,000) overnight at 4°C. HRP-conjugated donkey anti-mouse
secondary antibody (Jackson ImmunoResearch) (1:10,000) was
incubated with membranes 1-2 h at room temperature. Pierce ECL
Western Blotting Substrate (Thermo) and SuperSignal West Femto
Maximum Sensitivity Substrate (Thermo) were used as HRP substrates
for detection of tubulin and 2xHA::MJL-1, respectively.
**Figure S1.** Phylogenetic tree of MJL-1 homologs in *Caenorhabditis*.
A maximum-likelihood phylogenetic tree of MJL-1 homologs from representative *Caenorhabditis* species. Numbers on each node are Bootstrap values. Scale bar, 0.5 substitutions per site.

**Figure S2.** Interdependence among meiotic nuclear envelope proteins for localization and function.
(a) Pairing Centers (PCs) interact with the nuclear envelope in the absence of MJL-1. A cross-section of meiotic nuclei showing immunofluorescence of SUN-1 (left), which marks the nuclear envelope in meiotic cells; HIM-8 (center) marks X chromosome PCs. Although pairing is severely reduced in *mjll-1(tm1651)*, PCs are still associated with the nuclear envelope.
(b) MJL-1 abundance is greatly reduced in the absence of SUN-1. Western blot of proteins in strains expressing 2xHA::MJL-1 and (degron-tagged) AID::SUN-1, showing the abundance of MJL-1 detected with anti-HA antibodies, either in the absence of auxin treatment or following depletion with auxin for 24 hours.
C. brenneri
C. remanei
C. nigoni

Figure S3. Sequence alignment of transmembrane and perinuclear regions in Caenorhabditis MJL-1 homologs. Alignment was generated using MAFFT.

Figure S4. MJL-1 and SUN-1 are detected at the NE of oocyte nuclei undergoing apoptosis, while ZYG-12 is absent. Maximum-intensity projection images showing late pachynem nuclei. Arrowheads indicate apoptotic nuclei. Scale bar, 5 μm.

Figure S5. Deletion of a small acidic region in MJL-1 results in nonhomologous synopsis. Maximum-intensity projection images of pachynem nuclei stained with antibodies against HIM-8 (green in merged image) and SYP-2 (red). Scale Bar, 5 μm.
Figure S6. Depletion of LMN-1 by RNAi fails to rescue pairing in HIM-8^{T64A}.
Quantification of X chromosome pairing following depletion of the nuclear lamin protein LMN-1 by RNAi. HIM-8^{T64A} lacks a recruitment motif for PLK-2. The extended transition zone was divided into zones 2-4, with zone 1 corresponding to the pre-meiotic region. Each point represents a single gonad. P-values were calculated by one-way ANOVA with pairwise Bonferroni post-hoc correction (**p<0.0001).

Figure S7. Loss of PLK-2 correlates with dissociation of synopsis-independent pairing.
Images show maximum-intensity projections of the proximal region of the gonad, corresponding to the end of the extended transition zone, in syp-2(ok307) hermaphrodites. Gonads were stained with antibodies against phosphorylated PC proteins (green) and HIM-8 (magenta). Separation of HIM-8 foci correlates with a loss phosphorylation of HIM-8, indicative of loss of PLK-2 from the X chromosome PC.