

The *Caenorhabditis elegans* SUN protein UNC-84 interacts with lamin to transfer forces from the cytoplasm to the nucleoskeleton during nuclear migration

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ABSTRACT Nuclear migration is a critical component of many cellular and developmental processes. The nuclear envelope forms a barrier between the cytoplasm, where mechanical forces are generated, and the nucleoskeleton. The LINC complex consists of KASH proteins in the outer nuclear membrane and SUN proteins in the inner nuclear membrane that bridge the nuclear envelope. How forces are transferred from the LINC complex to the nucleoskeleton is poorly understood. The *Caenorhabditis elegans* lamin, LMN-1, is required for nuclear migration and interacts with the nucleoplasmic domain of the SUN protein UNC-84. This interaction is weakened by the *unc-84(P915)* missense mutation. These mutant nuclei have an intermediate nuclear migration defect—live imaging of nuclei or LMN-1::GFP shows that many nuclei migrate normally, others initiate migration before subsequently failing, and others fail to begin migration. At least one other component of the nucleoskeleton, the NET5/Samp1/lma1 homologue SAMP-1, plays a role in nuclear migration. We propose a nut-and-bolt model to explain how forces are dissipated across the nuclear envelope during nuclear migration. In this model, SUN/KASH bridges serve as bolts through the nuclear envelope, and nucleoskeleton components LMN-1 and SAMP-1 act as both nuts and washers on the inside of the nucleus.

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INTRODUCTION

Nuclear migration is essential for a wide variety of cellular processes conserved across eukaryotes, including cell migration, cell division, and polarity establishment (Morris, 2000; Burke and Roux, 2009; Starr and Fridolfsson, 2010; Gundersen and Worman, 2013). Nuclear migrations are also central to many developmental processes, including fertilization (Reinsch and Gonczy, 1998), neurogenesis (Tsai *et al.*, 2007; Zhang *et al.*, 2009), and muscle development

(Bruusgaard *et al.*, 2006). In addition, nuclear migration is an important process in metastasizing cancer cells (Chow *et al.*, 2012; Fu *et al.*, 2012).

Nuclear migration requires mechanical forces generated by the cytoskeleton to be transferred to the nucleoskeleton—the structural network of the nucleus consisting of lamins, actin, and inner nuclear membrane components (Simon and Wilson, 2011). The nuclear envelope is a specialized extension of the endoplasmic reticulum consisting of an outer nuclear membrane, an inner nuclear membrane, and the lumen between the membranes, called the perinuclear space (Franke *et al.*, 1981). The inner nuclear membrane is tightly associated with the underlying nucleoskeleton. The unique architecture of the nuclear envelope presents a special challenge to force transfer from the cytoskeleton to the nucleoskeleton.

A nuclear membrane complex of Sad1 and UNC-84 (SUN) and Klarsicht, ANC-1, and Syne homology (KASH) proteins, termed the linker of nucleoskeleton and cytoskeleton (LINC) complex, traverses the barrier created by the nuclear envelope and allows for forces generated in the cytoplasm to be transduced into the nucleus

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Abbreviations used: KASH, Klarsicht, ANC-1, and Syne homology; LINC, linker of nucleoskeleton and the cytoskeleton; SUN, Sad1 and UNC-84.

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(Starr and Fridolfsson, 2010; Tapley and Starr, 2013). SUN proteins are single-pass transmembrane proteins specifically localized to the inner nuclear membrane. They consist of an N-terminal nucleoplasmic domain and a C-terminal domain in the perinuclear space containing the conserved SUN domain (Turgay et al., 2010; Tapley et al., 2011; Tapley and Starr, 2013). The SUN domain functions to recruit KASH proteins to the outer nuclear membrane through a direct interaction between conserved SUN and KASH domains in the perinuclear space (Crisp et al., 2006; McGee et al., 2006; Sosa et al., 2012; Tapley and Starr, 2013). KASH proteins are the only known integral membrane proteins that are specifically localized to the cytoplasmic surface of the nucleus. They are classified by a small conserved KASH peptide at the C-terminus of the protein (Starr and Han, 2002; Starr and Fridolfsson, 2010). The large cytoplasmic domains of KASH proteins interact with a variety of cytoskeletal elements, including microtubule motors, actin, and intermediate filaments (Luxton and Starr, 2014). Thus KASH proteins interact with the cytoskeleton and then partner with SUN proteins to form a bridge across both membranes of the nuclear envelope, allowing the transfer of force to position nuclei.

Interactions between the cytoskeleton and KASH proteins and between SUN and KASH proteins are relatively well understood (Tapley and Starr, 2013; Luxton and Starr, 2014). However, it is much less clear how SUN proteins interact with the nucleoskeleton. The major component of the nucleoskeleton is the intermediate filament lamin, which provides structure and strength to the nuclear envelope. Vertebrates have two types of lamin proteins; B-type lamins are broadly expressed, and A/C-type lamins are expressed in differentiated tissues (Gruenbaum et al., 2005; Dittmer and Misteli, 2011; Simon and Wilson, 2011). A large class of diseases, called laminopathies, has been linked to mutations primarily in lamin A/C (Worman, 2012). Because lamin A/C is involved in disease, most studies on interactions between lamins and SUN proteins have focused on lamin A/C rather than the more broadly expressed lamin B. Therefore how SUN proteins interact with the nuclear lamina and especially lamin B remains an open question. Here we test the hypothesis that SUN proteins interact with lamin B during nuclear migration.

Reports of interactions between SUN proteins and lamin A/C are limited to in vitro glutathione S-transferase (GST) pull-down assays and fluorescence recovery after photobleaching and fluorescence resonance energy transfer assays in transfected tissue culture cells. These data show that SUNs interact with lamin A/C, but conflict as to whether mammalian SUN1 or SUN2 binds more tightly (Crisp et al., 2006; Ostlund et al., 2009). Other studies show that some lamin A disease mutations disrupt the ability of lamin A to bind SUN proteins, whereas other mutations increase the interaction between lamin A and SUN1 (Haque et al., 2010). Nonetheless, SUN proteins properly localize to the nuclear envelope in lamin A mutant cells (Crisp et al., 2006; Haque et al., 2010; Chen et al., 2012). Lamin A is also required for nuclear migrations in polarizing fibroblasts (Folker et al., 2011). Depletion of SUN1 in lamin A-knockout mice increases longevity and the health of the mice, suggesting that the interaction between lamin A and SUN1 is important in the progression of laminopathies (Chen et al., 2012; Chen et al., 2014). One possible explanation for this finding is that by removing SUN1, the forces transferred to the weakened nucleoskeleton are reduced, which may lead to less mechanical damage to already fragile nuclei (Starr, 2012). Such a model fits well with our hypothesis that SUN proteins interact with lamins to move nuclei.

SUN protein interactions with lamin B are less well understood at a biochemical level than their lamin A/C counterparts. In *Caenorhabditis elegans* and *Drosophila*, lamin B is required for the localization

of SUN proteins (Lee et al., 2002; Kracklauer et al., 2007). However, the extent to which SUN localization to the nuclear envelope requires direct interaction with lamin B is not clear. There is conflicting evidence from in vitro pull-down assays as to whether lamin B interacts with mammalian SUN1 or SUN2 (Crisp et al., 2006; Haque et al., 2006). However, two important developmental genetic experiments suggest that lamin B functions in some of the same nuclear migration events as SUN and KASH proteins. Mice with knockout mutations in lamin B2 have nuclear migration defects in the CNS similar to SUN- and KASH-knockout defects (Zhang et al., 2009; Coffinier et al., 2010a,b, 2011). Similarly, null mutations in the *Drosophila* lamin B gene *Lam Dm_o* cause a nuclear migration defect in the developing eye disk very similar to that in SUN and KASH mutants (Patterson et al., 2004; Kracklauer et al., 2007). Taken together, these data are consistent with a model in which SUN proteins interact with lamin B to mediate nuclear migration.

Here we used nuclear migration in *C. elegans* embryonic hypodermal cells (Starr and Han, 2005; Zhou and Hanna-Rose, 2010) as a model for studying the interaction between SUN proteins and lamins. *C. elegans* has a single lamin gene, as compared with three to four lamins in vertebrate systems. Invertebrate lamins are widely considered as B-type lamins, but unrooted phylogenetic trees place invertebrate lamins in their own clade nearly equal distant from vertebrate lamin As and Bs (Liu et al., 2000; Dittmer and Misteli, 2011). Having a single lamin gene is both an advantage and a disadvantage for this study. It makes the study feasible but complicates the significance of the study when thinking about vertebrate cells. The *C. elegans* lamin protein LMN-1, also known as Ce-lamin and Ce-Lam-1, is broadly expressed and required for early embryonic cell divisions; *lmn-1(RNAi)* embryos die at around the 100-cell stage with multiple mitotic defects (Liu et al., 2000). Furthermore, only one SUN protein, UNC-84, is present in the cell at the time of hyp7 nuclear migration (Fridkin et al., 2004; Minn et al., 2009; Wang et al., 2009). Finally, *C. elegans* hyp7 nuclear migration is amenable to the use of many genetic and live-imaging tools (Starr et al., 2001; Fridolfsson et al., 2010; Fridolfsson and Starr, 2010). Here we combine *C. elegans* genetics and yeast two-hybrid assays to test our hypothesis that the SUN protein UNC-84 binds to the lamin B protein LMN-1. Furthermore, we use live imaging to carefully describe the nuclear migration phenotypes of *unc-84* mutants that disrupt the interaction with lamin B. Our data strongly support that SUN proteins bind directly to lamin B to transfer forces generated in the cytoskeleton to the nucleoskeleton, thus facilitating nuclear migration.

RESULTS

Mutations in the nucleoplasmic domain of UNC-84 lead to an intermediate nuclear migration defect

Null mutations in *unc-84* cause a complete block of nuclear migration in hyp7 precursors, resulting in ~14 nuclei residing abnormally in the dorsal cord of larvae (Figure 1; Malone et al., 1999; Fridolfsson and Starr, 2010). However, three alleles—*unc-84(e1411, e1174, and n322)*—were originally reported to cause an intermediate hyp7 precursor nuclear migration defect (Malone et al., 1999). Normally, nuclei migrate across the length of the dorsal hyp7 precursors during *C. elegans* embryogenesis. After the bulk of embryonic cell division and just before the initiation of morphogenesis, 15 dorsal epithelial cells intercalate, and their nuclei migrate across the dorsal midline to the contralateral side of the embryo (Figure 1A; Sulston et al., 1983). Nuclei are pulled along polarized microtubules by kinesin-1 and dynein, which are recruited to the surface of the nuclear envelope by the KASH protein UNC-83 (Meyerzon et al., 2009a; Fridolfsson et al., 2010; Fridolfsson and Starr, 2010). These cells

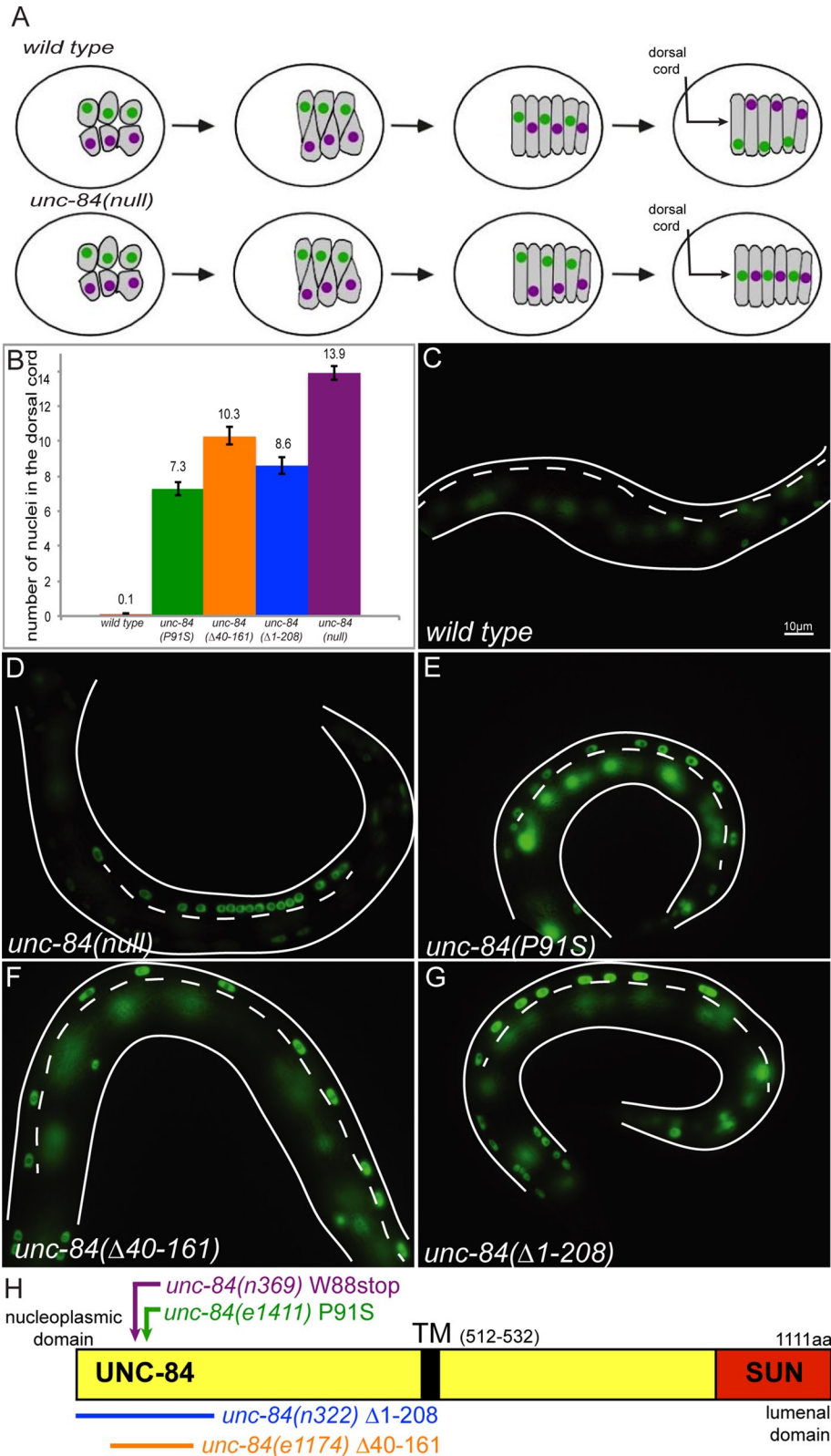


FIGURE 1: Mutations in the nucleoplasmic domain of UNC-84 lead to an intermediate nuclear migration defect. (A) Cartoon describing hyp7 precursor nuclear migration on the dorsal surface of the pre-comma-stage embryo. In wild-type embryos (top), two rows of hyp7 precursors (gray) intercalate to form a row of column-shaped cells. Nuclei then migrate from right to left (green) or left to right (purple). In *unc-84(null)* mutant embryos, intercalation occurs normally, but the nuclei fail to migrate. Instead, underlying body wall muscle migrations push *unc-84* nuclei to the dorsal cord (arrow). The dorsal surface is shown; anterior is left. (B) Average number of nuclei present in the dorsal cord of L1 larvae, which approximates the number of failed nuclear migrations. Error

subsequently fuse to form the embryonic dorsal hyp7 syncytium (Sulston *et al.*, 1983; Altun, 2009). Mutations that block nuclear migration in hyp7 precursors result in nuclei abnormally residing in the dorsal cord of newly hatched L1 larvae, having been pushed there by underlying body wall muscles (Figure 1A; Sulston and Horvitz, 1981; Malone *et al.*, 1999). About 90% of the nuclei that fail to migrate end up in the dorsal cord (Fridolfsson and Starr, 2010).

The *unc-84* alleles *e1411*, *e1174*, and *n322* resulting in an intermediate hyp7 precursor nuclear migration defect all disrupt the N-terminal nucleoplasmic domain of UNC-84. *unc-84(e1411)* is a P91S missense mutation, *unc-84(e1174)* is a deletion removing residues 40–161 of UNC-84, and *unc-84(n322)* is a small deletion of the ATG and is predicted to use the ATG at residue 209 (Figure 1H; Malone *et al.*, 1999). We henceforth refer to these alleles as *unc-84(P91S)*, *unc-84(Δ40-161)*, and *unc-84(Δ1-208)*.

To quantify partial nuclear migration defects, we created a transgenic line expressing nuclear GFP specifically in hypodermal nuclei (*ycls10[p_{col-10}nls::gfp::lacZ]*). The *unc-84* alleles *P91S*, *Δ40-161*, and *Δ1-208* caused nuclear migration defects in which 7.3 ± 0.4 , 10.3 ± 0.5 , and 8.6 ± 0.5 (mean \pm 95% confidence interval [CI]; Figure 1, B and E–G) nuclei fail to migrate, respectively. This intermediate phenotype is significantly different from either the null allele *unc-84(n369)*, for which 13.9 ± 0.4 nuclei failed to migrate, or wild-type animals, for which only 0.1 ± 0.1 nucleus was mispositioned to the dorsal cord (Figure 1). The intermediate nuclear migration defect of at least *unc-84(P91S)* is unlikely due to a reduction in the levels of the mutant UNC-84 protein as compared with wild type. Quantification of immunofluorescence intensity showed that approximately equal levels of UNC-83 protein were found at the nuclear envelope in both the *unc-84(P91S)* mutant and wild-type embryos (Supplemental

bars, 95% CI. (C–G) The number of nuclei in the larval dorsal cord was counted following hypodermal nuclei that express a nucleoplasmic GFP from *ycls10[p_{col-10}nls::gfp::lacZ]*. Lateral views of L1 larvae. Dorsal is up, and anterior is left; the dorsal cord (arrow in A) is demarcated by the white dotted line. Scale bar, 10 μ m. Representative images of (C) wild type, (D) *unc-84(null)*, (E) *unc-84(P91S)*, (F) *unc-84(Δ40-161)*, and (G) *unc-84(Δ1-208)*. (H) Schematic of the domain structure of UNC-84. The conserved SUN domain is red, and the transmembrane span is black. The mutants discussed in the text are indicated.

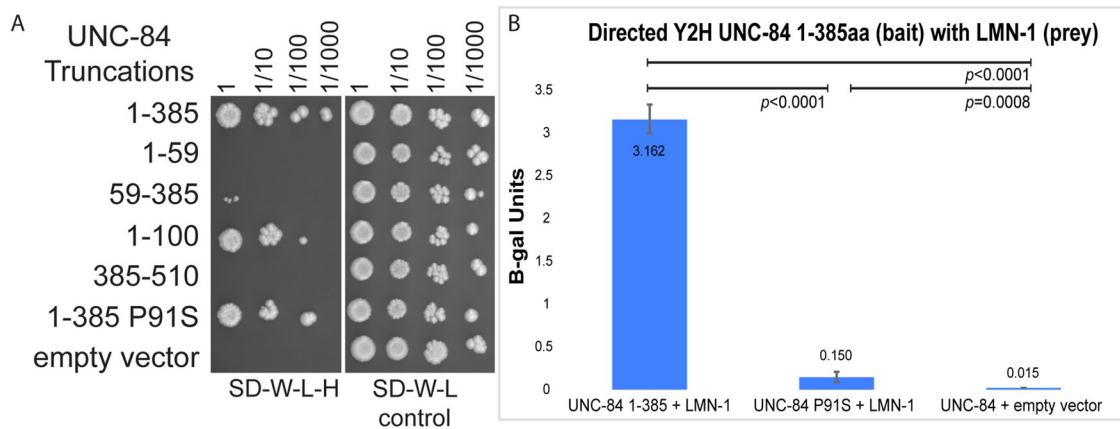


FIGURE 2: UNC-84 and LMN-1 interact in a yeast two-hybrid assay. (A) Yeast growing in a directed yeast two-hybrid assay. All yeast express the LMN-1::Gal4AD prey construct and the UNC-84::Gal4BD bait construct indicated on the left. Yeast were grown to the same concentration, serially diluted (as indicated at the top), and plated on SD–Trp–Leu–His medium, which requires an interaction to grow (left), or SD–Trp–Leu medium as control (right). (B) Activity of the *lacZ* gene as activated by a liquid *o*-nitrophenyl- β -galactosidase assay that represents a two-hybrid interaction. Average β -galactosidase units ($\Delta OD_{420}/\text{min/ml}$ of cells) from three different experiments, each done in triplicate, and the associated 95% CI error bars. Significant statistical differences as determined by Student's *t* test are noted at the top.

Figure S1). Because *unc-84(n369)*-null mutations disrupt both migration and anchorage (Malone *et al.*, 1999), we next asked about the extent to which these three mutant lines caused any anchorage defects. The nuclei that failed to migrate and are abnormally found in the dorsal cord of the *hyp7* syncytium are usually clumped together in *unc-84(n369)* mutant larvae (Figure 1D). We classified a nuclear anchorage defect (Anc^-) if an L1 larva had a row of at least three nuclei touching each other. In the null *unc-84(n369)* allele, 43% ($n = 14$) of larvae were Anc^- . In contrast, 0% of *unc-84(P91S)*, 6% of *unc-84(\Delta 40-161)*, and 0% of *unc-84(\Delta 1-208)* L1 larvae were Anc^- ($n \geq 30$). Our data therefore suggest that disruption of the nucleoplasmic domain of UNC-84 results in partial nuclear migration, but not nuclear anchorage, defects.

The nucleoplasmic domain of UNC-84 binds to lamin

We hypothesized that the P91S mutation in the nucleoplasmic domain of UNC-84 disrupted an interaction between UNC-84 and some unknown component of the nucleoskeleton. A yeast two-hybrid screen of a *C. elegans* mixed-stage cDNA library was conducted to identify proteins interacting with the nucleoplasmic domain of UNC-84. As bait we used the first 385 amino acids of UNC-84 fused to the GAL4 DNA-binding domain. This construct incorporates the majority of the nucleoplasmic domain of UNC-84 upstream of the transmembrane domain located at residues 512–532 (Figure 1H; Tapley *et al.*, 2011). Approximately 4×10^6 yeast clones were screened, and the prey inserts of 106 positive colonies were sequenced. Sixteen different proteins were identified as potential interacting partners of UNC-84. LMN-1, the sole *C. elegans* lamin protein (Liu *et al.*, 2000), was found in 16 independent clones. No other known component of the nucleoskeleton was identified.

We used the yeast two-hybrid assay to further map the LMN-1 interaction domain of UNC-84 (Figure 2A). The assay was repeated five times with UNC-84(1-385) and the empty vector to verify the interaction. The other constructs containing smaller regions of UNC-84 were examined at least twice. The original bait used for the screen, UNC-84(1-385), strongly interacted with the LMN-1 prey. A smaller bait, UNC-84(1-100), also interacted with LMN-1. However, UNC-84(1-59), UNC-84(59-385), and UNC-84(385-510) did not interact with LMN-1. These data suggest that the minimal interaction

domain is somewhere in the first 100 amino acids of UNC-84. Of interest, all three *unc-84* alleles with the intermediate *hyp7* nuclear migration phenotype disrupt this portion of UNC-84 (Figure 1H). We therefore tested the hypothesis that the *unc-84(P91S)* mutation disrupted the two-hybrid interaction with LMN-1. We used quantitative β -galactosidase liquid assays to measure the yeast two-hybrid interaction between LMN-1 and wild-type or P91S mutant UNC-84. The P91S mutation significantly reduced the strength of the interaction between LMN-1 and UNC-84, as determined by Student's *t* tests (Figure 2B).

lmn-1(RNAi) leads to a nuclear migration defect

The yeast two-hybrid data are consistent with a hypothesis that the *unc-84(P91S)* intermediate nuclear migration defect is due to a reduced interaction between UNC-84 and LMN-1. One prediction of this model is that disruptions of *lmn-1* should lead to similar nuclear migration defects. *lmn-1* is an essential gene required for the earliest embryonic cell divisions. Adults fed double-stranded RNA (dsRNA) against *lmn-1* for >24 h produce embryos that have small pronuclei and chromosomal segregation defects, leading to embryonic lethality before the 100-cell stage (Liu *et al.*, 2000; Meyerzon *et al.*, 2009b). To study the effect of *lmn-1(RNAi)* later in embryogenesis, at the time of nuclear migration in *hyp7* precursors, we fed young adults dsRNA against *lmn-1* over shorter windows, which allowed for the survival of 10–40 larvae per mother. These larvae demonstrated a nuclear migration defect in which screening a total of 121 larvae from four different experiments resulted in an average of 2.4 ± 0.5 (mean \pm 95% CI) *hyp7* nuclei in the dorsal cord (Figure 3). An example of an animal with ~50% *hyp7* nuclear migration failure is depicted in Figure 3B. The *lmn-1(RNAi)* *hyp7* nuclear migration failure is statistically more severe than in wild type ($p < 0.0001$ when using an unpaired *t* test with Welch's correction). The number of nuclei in the dorsal cord per animal ranges from 0 to 10. The range is large because individuals with no nuclei in the dorsal cord were likely subjected to little or no dsRNA, leading to incomplete knock-down of *lmn-1*. Finally, *lmn-1(RNAi)* treatment of the three UNC-84 N-terminal mutant lines resulted in minor enhancement. Given the hypomorphic nature of both the N-terminal mutations and *lmn-1(RNAi)*, this is consistent with our model that UNC-84 and LMN-1

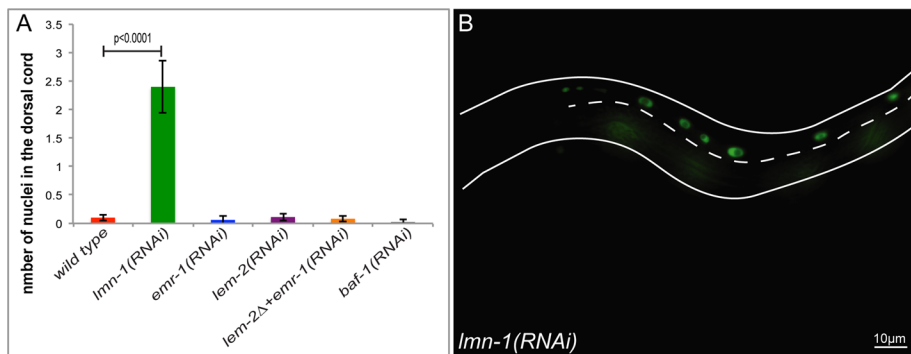


FIGURE 3: *Imn-1(RNAi)* animals have a nuclear migration defect. (A) Mean number of nuclei present in the dorsal cord of wild type, *Imn-1(RNAi)*, *emr-1(RNAi)*, *lem-2(RNAi)*, *lem-2(tm1582)*; *emr-1(RNAi)*, and *baf-1(RNAi)*. Error bars, 95% CI. (B) A representative *Imn-1(RNAi)* L1 larva depicting nuclei in the dorsal cord that have failed to migrate. The dashed line marks the dorsal cord, anterior is to the left, and dorsal is up. Scale bar, 10 μm.

function in the same pathway. This enhancement was not significant for either of the two deletion alleles: *unc-84(Δ40-161)* increased by an average of 0.35 nuclei, and *unc-84(Δ1-208)* increased by an average of 0.45 nuclei. The average enhancement in *unc-84(P91S)* of 1.9 nuclei was significant ($p < 0.0001$ with Student's *t* test) but did not increase to anywhere near the levels observed in *unc-84(null)* animals.

LMN-1 is part of a network of *C. elegans* proteins at the nucleoskeleton, including the Baf homologue BAF-1 and the LEM proteins *emr-1* and *lem-2*, that are believed to function in a mutually dependent network. Mutants with *baf-1* or *emr-1*; *lem-2* double mutations have identical early embryonic phenotypes as *Imn-1(RNAi)* (Liu et al., 2003; Margalit et al., 2005; Meyerzon et al., 2009b). We therefore examined the extent to which *baf-1*, *emr-1*, and *lem-2* functioned similarly to *Imn-1* during nuclear migration in hyp7 precursors. We observed the previously reported early embryonic lethality in *baf-1(RNAi)* and *lem-2(tm1582)*; *emr-1(RNAi)* embryos (unpublished data). However, when fed dsRNA for shorter periods of time as in our *Imn-1(RNAi)* experiment, we did not observe any nuclear migration defects in *baf-1(RNAi)* or *lem-2(tm1582)*; *emr-1(RNAi)* larvae (Figure 3A). We therefore conclude that LMN-1 functions independently of BAF-1 and the LEM proteins during embryonic hyp7 nuclear migration.

Live imaging of the partial nuclear migration defect in *unc-84(P91S)* embryos

We envisioned three possible ways that the *unc-84(P91S)* mutation could partially disrupt nuclear migration. First, half of the nuclei could migrate normally, whereas the other half failed to even initiate nuclear migration. Alternatively, *unc-84(P91S)* mutant nuclei could migrate at a slower velocity than wild-type nuclei, such that only half finish migration in the available period of time. Finally, *unc-84(P91S)* mutant nuclei might initiate nuclear migration at the normal time and velocity and then fail at some point later in nuclear migration. To distinguish among these possibilities, we observed *unc-84(P91S)* mutant nuclei by time-lapse imaging during hyp7 precursor development. Time-lapse imaging of hyp7 nuclear migration in wild-type, *unc-84(null)*, or microtubule motor mutant embryos previously led to great insights into the mechanisms of nuclear migration (Fridolfsson and Starr, 2010; Zhang and Han, 2010; Starr, 2011). For example, see our previous studies for details on wild-type and *unc-84(null)* nuclear migrations (Fridolfsson and Starr, 2010). Here we used both differential interference contrast (DIC)

microscopy and fluorescence imaging of LMN-1::green fluorescent protein (GFP) to follow nuclear migration in a subset of hyp7 precursor cells on the dorsal surface of the embryo (Figures 1A and 4, A–D; hyp7 precursors 11–16, following designations of Sulston et al., 1983). Nuclear migration defects in wild-type, *unc-84(null)*, and *unc-84(P91S)* embryos were quantified by four different assays. For the first three assays, DIC time-lapse microscopy was used to characterize the complete nuclear migration in embryonic hyp7 precursors (Figure 4, A–D). Images were taken at 15-s intervals over 35–50 min. Representative time-lapse movies of wild-type, *unc-84(null)*, and *unc-84(P91S)* embryos at the time of nuclear migration are shown in Supplemental Movies S1–S3.

Nuclear migration events were scored as normal if full contralateral movement was completed within 30 min, partial if migration initiated but failed to complete, or static if initiation did not occur (Figure 4E). See Supplemental Movie S3 for an example of a static nucleus (white arrowhead) and a nucleus that partially migrated (black arrowhead) in an *unc-84(P91S)* embryo. One hundred percent of wild-type nuclei completed their migrations normally. Eighty percent of the *unc-84(null)* nuclei were static, whereas 20% had partial migrations. The *unc-84(P91S)* nuclei had an intermediate phenotype for which 63.8% of nuclei migrated normally, 20.7% initiated but failed to complete migration, and 15.5% were static.

The time it took a nucleus to reach the midline was also analyzed (Figure 4F). We previously showed that wild-type nuclei reached the midline ~11 min after intercalation, whereas most *unc-84(null)* nuclei failed to reach the midline, and those that did took an average of 37 min (Fridolfsson and Starr, 2010). Here, the time a migrating nucleus took to reach the midline was binned into <10, 10–30, or >30 min or not within the length of the film (films were 35–50 min in length). Late during filming, embryos began to rotate in the egg shell. This often resulted in the midline moving to or past a subset of nuclei, thus making the *unc-84(null)* phenotype described in Figure 4F appear less severe than the phenotype presented in Figure 4E. Consistent with our previous results, the *unc-84(P91S)* mutant nuclei had an intermediate defect; 27% migrated to the midline in <10 min, but 25% took >30 min or never reached the midline. This intermediate phenotype was significantly different from wild-type or *unc-84(null)* nuclei as determined by χ^2 contingency tests (Figure 4F).

As a third assay, the distance nuclei moved in the first 10 min of migration was binned into half-micrometer increments and plotted (Figure 4G). Wild-type nuclei moved an average of 2.7 ± 0.4 μm and had the highest frequency of nuclei that migrated 2.6–3.0 μm in the first 10 min of migration (blue in Figure 4G). *unc-84(null)* nuclei migrated an average of 1.2 ± 0.3 μm and were shifted greatly toward the three lowest categories that migrated <1.5 μm in 10 min (green in Figure 4G). *unc-84(P91S)* nuclei migrated an average of 1.9 ± 0.4 μm, which is statistically less than wild-type movements ($p = 0.005$ in a *t* test) and more than *unc-84(null)* nuclei move ($p = 0.016$).

To visualize changes in nuclear envelope morphology over shorter time frames, we performed time-lapse fluorescence imaging to follow LMN-1::GFP in migrating nuclei. A transgenic line expressing *Imn-1::gfp* under the control of a hypodermal-specific promoter

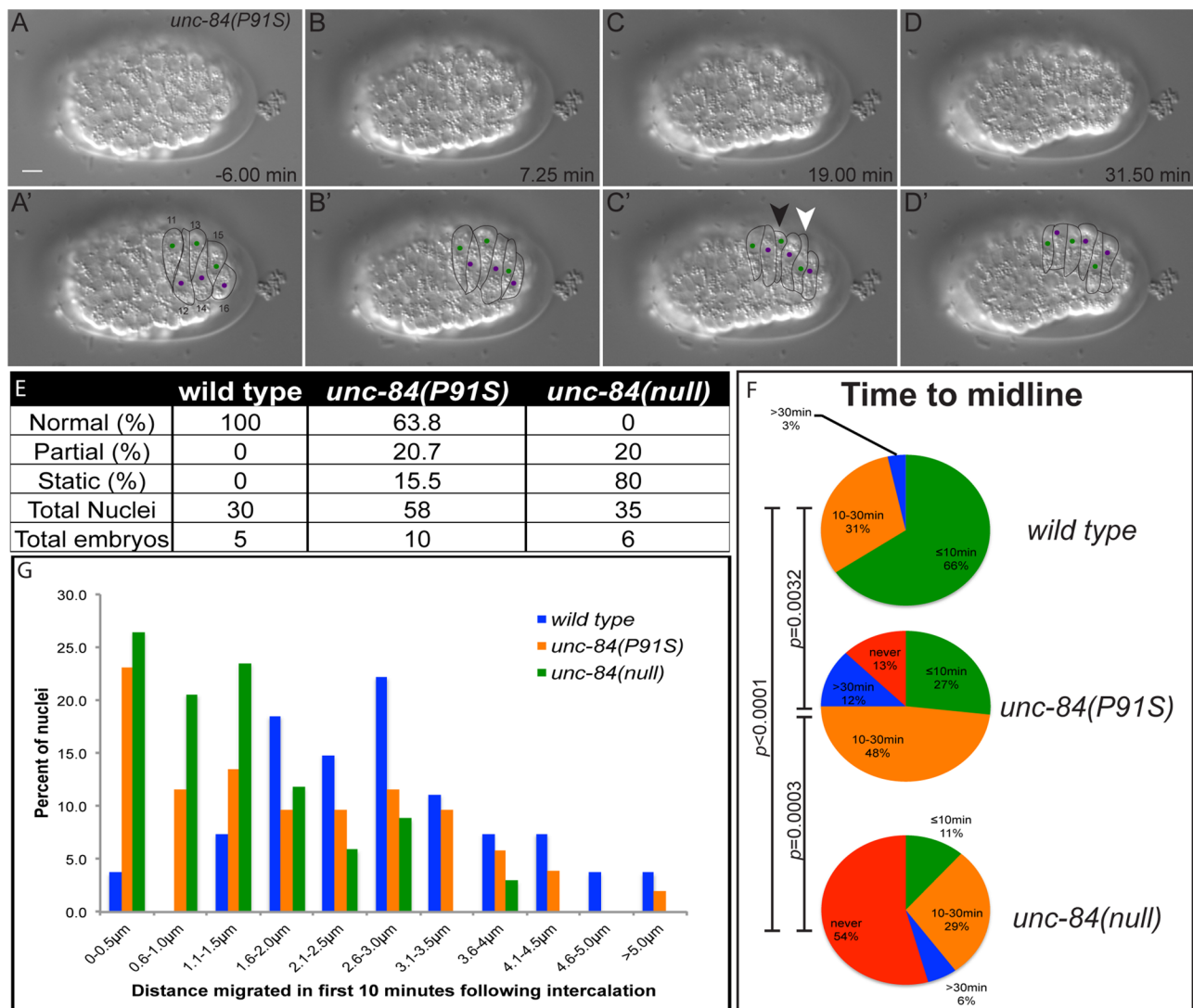


FIGURE 4: Time-lapse imaging of *unc-84* mutant nuclear migration events. (A–D) A time-lapse series showing nuclear migration in an *unc-84(P91S)* mutant embryo. Dorsal views with anterior on the left. Top row, raw DIC images; bottom row, marked cell–cell boundaries and nuclei. The green nuclei in cells 11, 13, and 15 migrate from right to left, and the purple nuclei in cells 12, 14, and 16 migrate left to right. The nucleus of cell 13 (white arrowhead) fails to migrate, whereas the nucleus in cell 15 (black arrowhead) migrated halfway across and then stopped. Scale bar, 5 μm . This is the same embryo as in Supplemental Movie S3. (E) Quantification of percentage of nuclei that migrated normally, initiated nuclear migration but failed to complete it (partial), or failed to move at all (static). (F) Quantification of the time it took nuclei to reach the dorsal midline of the embryo. Nuclei were categorized into those that reached the midline within 10 min of the completion of intercalation (green), at 10–30 min (orange), at >30 min (blue), or never (red). Significant statistical differences as determined by χ^2 contingency tests are noted on the left. (G) The distance a nucleus traveled in the first 10 min after completion of intercalation plotted in a histogram. Each individual nucleus was binned into 0.5- μm increments.

from an extrachromosomal array (Fridolfsson and Starr, 2010) was crossed to *unc-84(P91S)* and *unc-84(null)* animals. Embryos at the stage at which *hyp7* nuclear migration would normally occur were identified using DIC microscopy. LMN-1::GFP was then imaged in these embryos at 1-s intervals for 3–9 min to follow changes in nuclear envelope morphology during nuclear migration. Movies of LMN-1::GFP in wild-type, *unc-84(null)*, and *unc-84(P91S)* embryos were visually different (Supplemental Movies S4–S6). Nuclei in wild-type embryos underwent large movements—greater than half the width of a nucleus—and were almost constantly moving (Figure 5A and Supplemental Movie S4). In contrast, *unc-84(null)* nuclei tended to remain in place over several minutes of filming (Figure 5B and

Supplemental Movie S5); most movements were due to the drift of the entire embryo within its eggshell. Of interest, in *unc-84(P91S)* nuclei, both phenotypes were visualized. Some nuclei were observed undergoing large directional movements of up to 1 $\mu\text{m}/\text{min}$, whereas other nuclei did not move at all.

To categorize the movements of LMN-1::GFP during nuclear migration, we made projections combining each frame of an 8 min, 20 s time-lapse series (Figure 5, A'–C'). The projections were split into three colors to show the direction of movement. Magenta signifies the first third of the series, yellow the second, and cyan the final third. Using the time-lapse projections of LMN-1::GFP, we binned nuclei into three categories based on the size of an individual

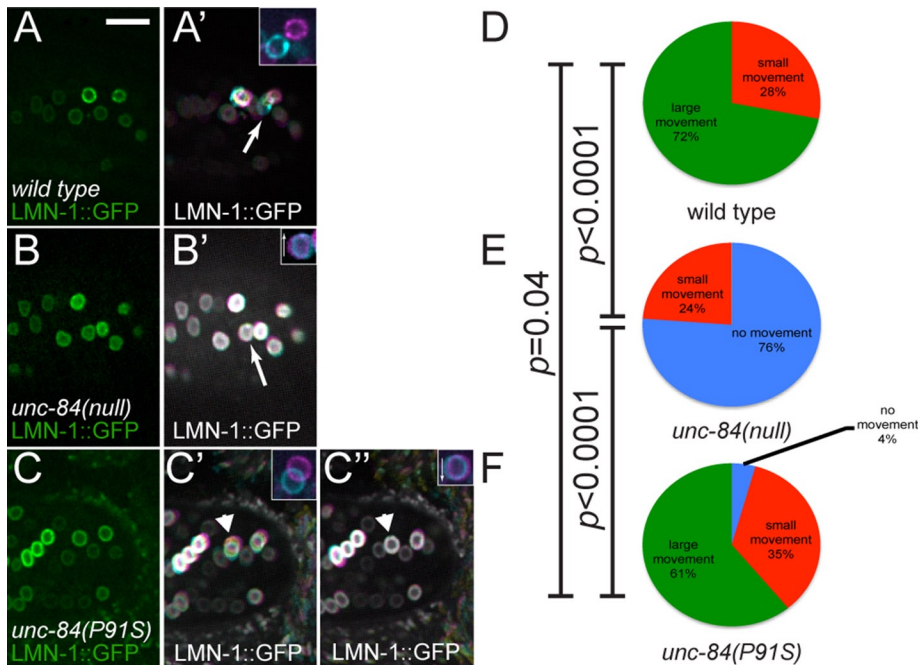


FIGURE 5: LMN-1::GFP shows dynamic nuclear morphology during nuclear migration. (A–C) Images of embryos expressing LMN-1::GFP specifically in hypodermal cells at the start of time-lapse imaging. Dorsal views; anterior is left. Insets show the identified nucleus at the beginning (magenta) and end (cyan) of the 8 min, 20 s film. Arrows in insets show the direction the nucleus is supposed to be moving. (A) Wild-type, (B) *unc-84(null)*, and (C) *unc-84(P91S)* embryos. (A'–C') Time projections of 500 frames taken at 1-s intervals. In these projections, frames 1–166 are colored magenta, 167–333 are yellow, and 334–500 are cyan to show the direction of movement (A'–C'). A second time-lapse projection of the same embryo for *unc-84(P91S)* (C''). The arrowheads in C' and C'' mark a *unc-84(P91S)* nucleus that was migrating normally in time-lapse 1 (C') but then failed to continue migration in time lapse 2 (C''). Scale bar, 10 μ m. (D–F) Nuclei were classified into three categories: no movement, small movement, and large movement. The percentage in each category is depicted. Significant statistical differences as determined by χ^2 contingency tests are noted on the left. The arrow in A' is an example of a large movement, and the arrow in B' demonstrates no movement.

nuclear projection (Figure 5, D–F). To better visualize movement, insets show the nuclei identified in the projections in the first frame (magenta) and the last frame (cyan) of the film. Many nuclei had large directional movements over the course of imaging, as visualized by lack of overlap between the initial and final positions of the nucleus of at least half the width of the nucleus (arrow and inset in Figure 5A'; green in Figure 5, D–F). Other nuclei that moved small amounts but the projections of which remained mostly circular were classified as small movements. Finally, nuclei that did not move in up to 9 min of imaging were scored as static when the time-lapse projection remained circular, and when the projection was split into thirds, the colors were merged to white (arrow in Figure 5B'). The same identified nucleus is shown in the inset, which demonstrates slight embryo drift, as the first and last images are not directly superimposed (inset in Figure 5B'). In summary of these data, 72% of wild-type nuclei moved large distances, whereas 28% had small movements (Figure 5D). Seventy-six percent of *unc-84(null)* nuclei did not move, whereas the remaining 24% had only small movements (Figure 5E). In *unc-84(P91S)* animals, large movements were seen 61% of the time, and small movements were seen in 35% of nuclei; the remaining 4% of nuclei did not move (Figure 5F). Our LMN-1::GFP movement assay demonstrated statistically significant differences when comparing *unc-84(null)* nuclear migrations to both wild-type and *unc-84(P91S)* embryos ($p < 0.0001$ using a χ^2 contingency test). However, wild type and *unc-84(P91S)* were not significantly

different ($p = 0.4$). The lack of statistical significance may result from the relatively short duration of the time-lapse series, such that only a snapshot of nuclear migration was visualized as compared with the longer analyses in Figure 4. Nonetheless, the *unc-84(P91S)* phenotype followed the trend of intermediate nuclear migration phenotypes. Multiple time-lapse series were taken of some embryos. Occasionally *unc-84(P91S)* nuclei were observed to move in one series but then failed to migrate in the subsequent series (arrowhead and insets in Figure 4, C' and C''). In another *unc-84(P91S)* time-lapse movie, a nucleus was observed in which a large and rapid invagination appeared to push the nucleus just before the time of nuclear migration initiation (Supplemental Movie S7). This rapid change may have resulted from abrupt microtubule motor activity acting against a weakened UNC-84–LMN-1 interaction. Together these data are consistent with our hypothesis that a weakened connection between UNC-84 and LMN-1 could lead to a nucleus that initiates migration normally but then fails to complete its migration.

The inner nuclear membrane component SAMP-1 functions during nuclear migration

In our working model, forces generated in the cytoplasm are transmitted across the nuclear envelope by SUN/KASH bridges and then dissipated across the nucleoskeleton by lamins. The nucleoskeleton consists of lamins, scores of inner nuclear membrane proteins, and other proteins that mediate interactions between the nuclear envelope and chromatin (Simon and Wilson, 2011). We therefore hypothesized that other components of the nucleoskeleton play roles in connecting the nucleus to the nuclear envelope to allow for force dissipation during nuclear migration. An attractive candidate to play such a role is the Samp1/NET5/lma1 *C. elegans* homologue T24F1.2, which we named SAMP-1. The mammalian putative orthologue was originally found in a proteomic screen for integral components of the inner nuclear membrane and named NET5 (Schirmer *et al.*, 2003). NET5 was subsequently named Samp1 and shown to play a role in positioning nuclei in polarizing NIH 3T3 cells. Nuclear migration in polarizing mouse NIH3T3 cells relies on SUN-KASH bridges to couple moving actin arrays in the cytoplasm to the nucleoskeleton (Luxton *et al.*, 2010; Folker *et al.*, 2011). This nuclear migration also requires Samp1, which partially colocalizes and coimmunoprecipitates with SUN proteins in transmembrane actin-associated nuclear (TAN) lines (Borrego-Pinto *et al.*, 2012). The homologous protein in *Schizosaccharomyces pombe*, lma1, interacts in yeast two-hybrid assays with the SUN protein Sad1 and has been implicated in the maintenance of nuclear morphology (Hiraoka *et al.*, 2011). Previously a broad bioinformatics study predicted that *C. elegans* SAMP-1 would be a component of the nuclear envelope and confirmed this localization in the early embryo using a transgenic SAMP-1::GFP fusion protein (Gunsalus *et al.*, 2005). However, nothing else is known about the function of *C. elegans* SAMP-1. We

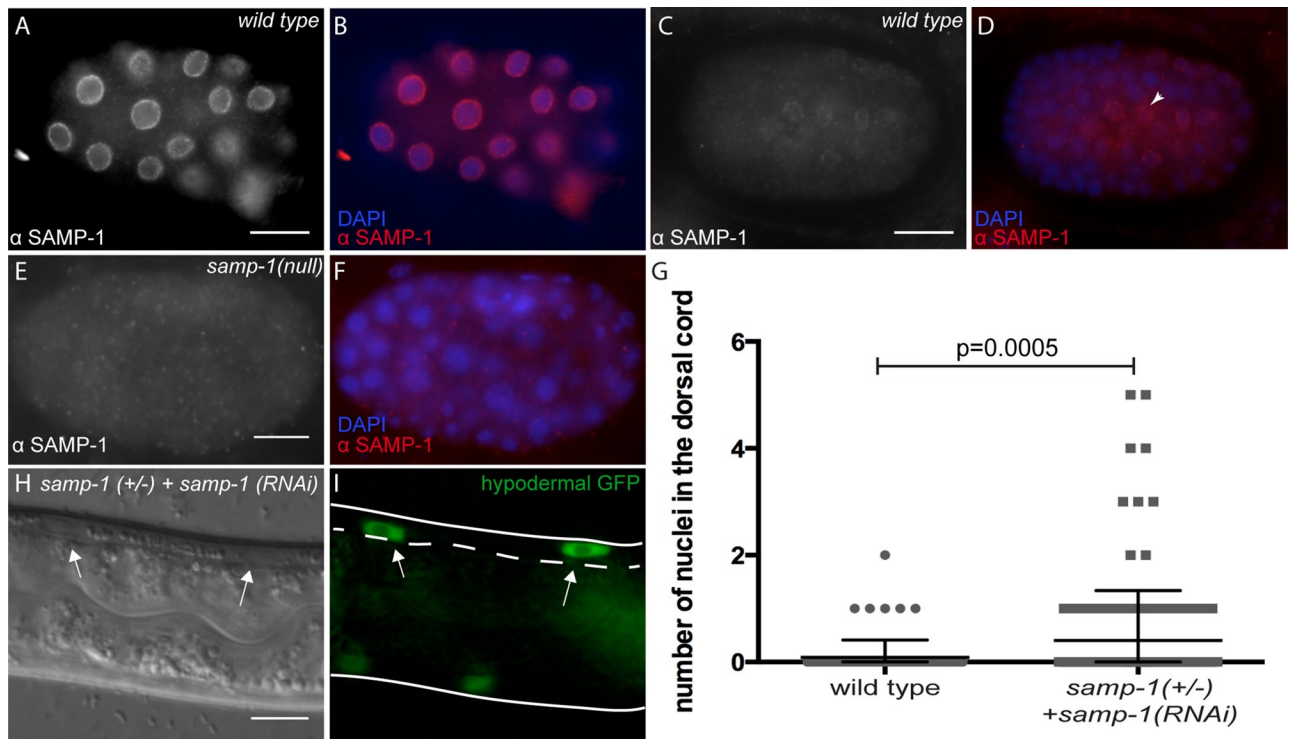


FIGURE 6: *samp-1(RNAi)* animals have a weak nuclear migration defect. (A–F) Embryos were stained for SAMP-1 localization. Lateral views, with anterior left and dorsal up. Scale bars, 10 μ m. For each pair of images, SAMP-1 immunostaining is shown in white on the left and in red on the right when it is merged with DAPI staining of nuclei in blue. (A, B) An early wild-type embryo. (C, D) A later, pre-comma-stage embryo. Arrowhead points to a hyp7 precursor nucleus. (E, F) A *samp-1(tm2710)*-null embryo is shown to demonstrate specificity of the antibody. (G) Numbers of nuclei in the dorsal cords of wild-type or *samp-1(tm2710)/+; samp-1(RNAi)* L1 larvae. Each gray dot represents an individual animal. The mean and 95% CI error bars are shown. (H, I) DIC and GFP images showing two hyp7 nuclei abnormally in the dorsal cord (arrows) of a *samp-1(tm2710)/+; samp-1(RNAi)* L1 larva. The dorsal cord is up and is demarcated by the dotted line. Scale bar, 10 μ m.

therefore set out to examine the role of *C. elegans* SAMP-1 in nuclear migration.

We first characterized the intracellular localization pattern of endogenous SAMP-1 to see whether it was plausible that SAMP-1 functions at the nuclear envelope during nuclear migration in embryonic hyp7 precursor. Antibodies were raised against the C-terminus of SAMP-1. Anti-SAMP-1 antibodies recognized a band of the predicted size on a Western blot. The band intensity was greatly reduced in *samp-1(RNAi)* extracts (Supplemental Figure S2). SAMP-1 antibodies localized strongly to a ring around 4',6-diamidino-2-phenylindole (DAPI)-stained nuclei, consistent with nuclear envelope staining, in all cells of wild-type early embryos but not in *samp-1(tm2710)* likely null embryos (Figure 6 and Supplemental Figure S2). Therefore the antibody is specific for SAMP-1, with a localization pattern expected for a nuclear membrane protein. Although we did not test the specific localization within the nuclear envelope, we hypothesize that SAMP-1 is an inner nuclear membrane protein based on the published localization of the mouse orthologue, Samp1 (Buch et al., 2009). In later embryos at the time of hyp7 nuclear migration, SAMP-1 localization at the nuclear envelope was less strong and limited to a subset of cells that included hyp7 precursors (Figure 6, C and D).

In mammalian tissue culture, Samp1 requires lamin A/C for localization to the nuclear envelope (Borrego-Pinto et al., 2012). It has also been demonstrated that *C. elegans* UNC-84 requires LMN-1 for nuclear envelope localization (Lee et al., 2002). Surprisingly, SAMP-1 localized to the nuclear envelope in *lmn-1(RNAi)* embryos

(Figure 7). In both early embryos (Figure 7, A and B) and embryos around the time of migration (Figure 7, C and D), SAMP-1 was able to localize in lamin-knockdown animals, whereas UNC-84 was not. LMN-1 staining was used as a control to confirm that the *lmn-1(RNAi)* knockdown was efficient.

After showing that SAMP-1 localizes to the nuclear envelope in migrating nuclei, we tested the extent to which SAMP-1 functions to move nuclei. Homozygous *samp-1(tm2710)* was embryonic lethal. We therefore fed *samp-1(tm2710)/+* adults dsRNA against *samp-1* for 48–72 h and examined their offspring. Nuclei abnormally located in the dorsal cord were counted in 266 *samp-1(tm2710)/+; samp-1(RNAi)* L1 larvae. On average, 0.4 ± 0.1 nuclei (mean \pm 95% CI) were observed in the dorsal cord (Figure 6, G–I), which is statistically significantly when compared with wild type ($p = 0.005$ by unpaired t test with Welch's correction). Occasionally, *samp-1(RNAi)* L1 larvae had up to five nuclei/worm that failed to migrate (Figure 6G). We therefore concluded that *samp-1* plays a small but significant role in nuclear migration.

DISCUSSION

The results presented here combine genetic analyses, time-lapse imaging of nuclear migration, and a yeast two-hybrid screen. Together the data provide mechanistic insights into both the molecular interaction between the SUN protein UNC-84 and lamin and the functional implications of disruption of this interaction during nuclear migration. We showed that alleles disrupting the N-terminal nucleoplasmic domain of UNC-84 led to an intermediate nuclear

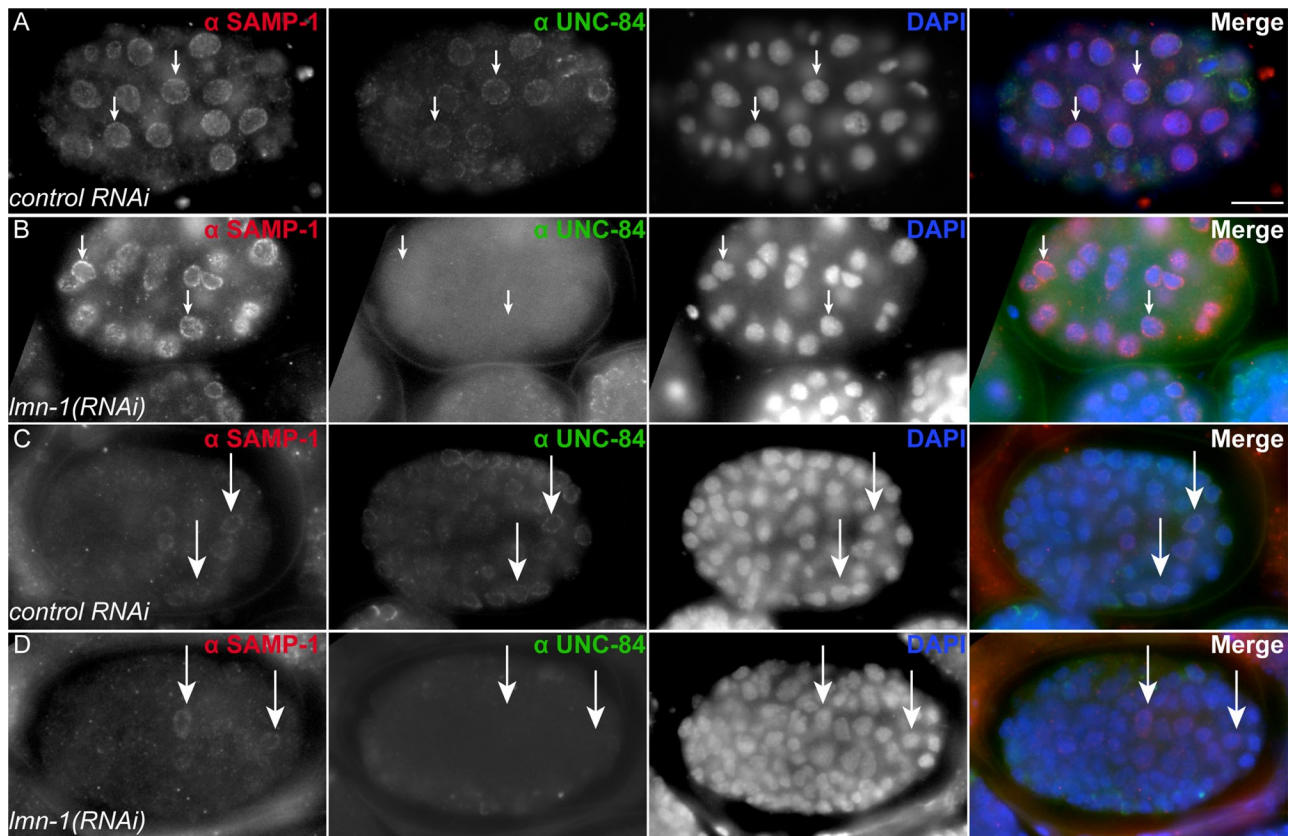


FIGURE 7: SAMP-1 localizes independently of LMN-1. (A–D) Embryos were stained for SAMP-1 and UNC-84 localization. Lateral views, with anterior left and dorsal up. For each row, SAMP-1 immunostaining is shown in white in the left column and in red on the right when all channels are merged. UNC-84 is shown in white in the second column and in green when merged. DAPI staining of nuclei is shown in white in the third column and in blue when merged. (A) An early embryo fed bacteria containing the empty L4440 vector as control. (B) An early embryo fed *lmn-1(RNAi)*. (C) A later, pre-comma-stage embryo fed bacteria containing the empty L4440 vector as control. (D) A later, pre-comma-stage embryo fed *lmn-1(RNAi)*. Arrows highlight specific nuclei to provide reference points in all four columns. Scale bar, 10 μ m.

migration defect. We then performed a yeast two-hybrid screen to find candidate interacting partners of the nucleoplasmic domain of UNC-84. Of interest, we identified an interaction between UNC-84 and the *C. elegans* lamin protein LMN-1. Furthermore, the point mutation UNC-84(P91S) that led to an intermediate nuclear migration phenotype also disrupted the interaction between UNC-84 and LMN-1. As predicted from these data, *lmn-1(RNAi)* led to a similar nuclear migration defect. Knockdown of another member of the nucleoskeleton, *samp-1*, led to a weak nuclear migration phenotype. Nuclear migrations in *unc-84(P91S)* embryos were carefully analyzed by time-lapse imaging to provide insight into the intermediate nuclear migration defect. Some nuclei in the *unc-84(P91S)* mutant background migrated normally, whereas others failed. Of the failed nuclei, many initiated migration normally before stopping part way through, whereas others failed to move at all. Of interest, we did not see slow-moving nuclei in *unc-84(P91S)* embryos; the nuclei that did move did so similarly to wild-type nuclei.

Our proposed model is shown in Figure 8. In this model, KASH/SUN bridges serve as molecular bolts through the nuclear envelope to transfer forces generated in the cytoplasm across the nuclear envelope. UNC-83 on the cytoplasmic surface of the nuclear envelope interacts with microtubule motors kinesin-1 and dynein (Meyerzon *et al.*, 2009a; Fridolfsson *et al.*, 2010). The forces generated by the motors are transmitted across the outer nuclear membrane by UNC-

83 and then to the SUN protein UNC-84 through an interaction between KASH and SUN domains (Sosa *et al.*, 2012; Tapley and Starr, 2013). Finally, UNC-84 spans the inner nuclear membrane (Tapley *et al.*, 2011) and interacts with lamin (Figure 2) to complete the connection between the cytoskeleton to the nucleoskeleton (Figure 8). However, the amount of force that can be transferred across the nuclear envelope by the KASH/SUN molecular bolt without stabilization by an interaction with the nucleoskeleton is limited. It is useful to think of this complex as analogous to a nut and bolt in drywall, where the interaction of UNC-84 with nucleoskeletal components dissipates forces across a larger area along the inside of the wall, reducing the likelihood of failures under tension. In our model, LMN-1 functions as a key component of the molecular nut and washer to dissipate the forces transmitted across the KASH/SUN bolt to structural elements inside the nucleus. The LMN-1 nut and washer are only as strong as the interaction with the UNC-84 nucleoplasmic domain of the bolt. If the UNC-84–LMN-1 interaction fails, as in the UNC-84(P91S) mutant, the nucleus could continue to migrate for some time without lamin serving as a nut and washer. However, at some critical point the forces are too strong, and the KASH/SUN bolt is catastrophically detached, leading to a failure in nuclear migration. The stability of the bolt depends on many factors. It is therefore difficult to predict when the KASH/SUN bridge will fail, explaining why some UNC-84(P91S) nuclei fail at the initiation of

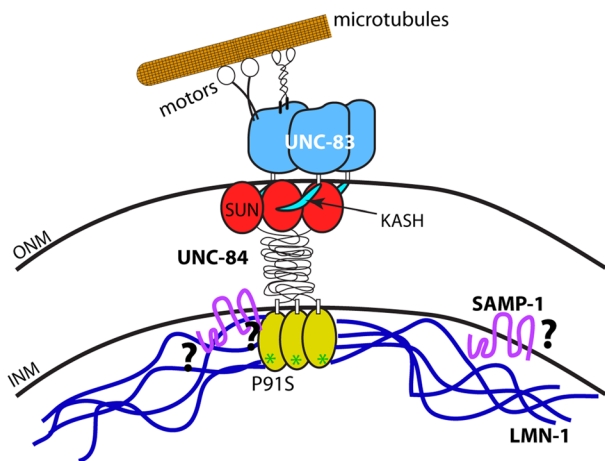


FIGURE 8: Nut-and-bolt model for nuclear migration. Cartoon of the KASH/SUN nuclear envelope bridge during nuclear migration. UNC-83 is shown in blue, with the KASH peptide in teal. UNC-84 is shown with the SUN domain in red, the domain spanning the perinuclear space in black, and the nucleoplasmic domain in yellow. The green asterisks indicate the P91S mutation in UNC-84. LMN-1 is shown in dark blue and SAMP-1 in fuchsia. Microtubule motors are shown in black and white interacting with a single microtubule in orange. Question marks symbolize open questions regarding protein interactions.

migration, whereas others migrate part way before stalling and many successfully complete migration. Although the LMN-1 nut is required to ensure normal nuclear migration, it does not appear necessary for nuclear anchorage, because *unc-84(P91S)* nuclei are normally anchored (Figure 1; Malone *et al.*, 1999).

It is difficult to translate our findings from the *C. elegans* system, in which there is a single lamin, to vertebrate cells, in which there are three to four lamins. However, mutations in lamins in other developmental systems lead to nuclear migration phenotypes consistent with our model. Mice carrying knockout mutations in lamin B2 have nuclear migration defects in the developing CNS similar to defects in SUN or KASH mutant mice (Coffinier *et al.*, 2010b). Similarly, mutations in *Drosophila Lam Dm_o* have nuclear migration defects in the developing eye disk similar to SUN and KASH mutants (Patterson *et al.*, 2004). Furthermore, heterozygous, dominant loss-of-function mutations in *Drosophila Lam Dm_o* were identified as enhancers of phenotypes caused by overexpression of the KASH protein in the developing eye disk, suggesting that the function of SUN-KASH bridges is sensitive to the levels of lamin (Patterson *et al.* 2004). These nuclear migration defects are consistent with the model that lamin serves an evolutionarily conserved role as a molecular nut and washer during many nuclear migration events in developing tissues. Defects in human lamin B have been associated with disease, although nowhere near to the extent of lamin A mutations (Coffinier *et al.*, 2010b; Worman, 2012). A duplication of *LMNB1* is associated with a leukodystrophy, in which myelin is progressively lost (Padiath *et al.*, 2006). Mutations in the *LMNB2* gene are linked to a lipodystrophy (Hegele *et al.*, 2006). Given the mouse knockout phenotypes, it is likely that additional mutations in *LMNB1* and *LMNB2* will be found to be associated with neuronal diseases (Coffinier *et al.*, 2010b).

The nucleoskeleton is a complex network consisting of lamins, inner nuclear membrane proteins, and small proteins that link the

nuclear envelope to chromatin (Simon and Wilson, 2011). It is therefore unlikely that lamin B is the only component of the molecular nut and washer that dissipates forces throughout the nucleoskeleton during nuclear migration. Obvious candidates to participate with lamins during nuclear migration include the LEM proteins, Baf, and Ima1/NET5/Samp1. In *C. elegans*, mutations in *baf-1* or double mutations in *emr-1* and *lem-2* phenocopy all previously described *lmn-1* phenotypes (Liu *et al.*, 2003; Margalit *et al.*, 2005; Meyerzon *et al.*, 2009b). We were therefore surprised that disrupting *baf-1* or *emr-1* and *lem-2* did not lead to a nuclear migration phenotype. The inner nuclear membrane protein Ima1/NET5/Samp1 was previously shown to play a role in nuclear migration in mammalian tissue culture cells (Borrego-Pinto *et al.*, 2012). It was also shown to interact with a SUN protein in both yeast and mammalian tissue culture (Hiraoka *et al.*, 2011; Borrego-Pinto *et al.*, 2012). Here we showed that the *C. elegans* homologue SAMP-1 plays a minor role in nuclear migration. Of interest, we found despite the requirement of lamin A/C in tissue culture (Borrego-Pinto *et al.*, 2012), *C. elegans* SAMP-1 localized independently of LMN-1. The SAMP-1 binding partners in the *C. elegans* nuclear envelope remain to be determined (question marks in Figure 8). We suspect that *baf-1*, *emr-1*, *lem-2*, *samp-1*, and other components of the nucleoskeleton play partially redundant roles with lamin during nuclear migration. Therefore characterization of how the network of proteins forming the nucleoskeleton functions as a unit during nuclear migration requires further investigation.

MATERIALS AND METHODS

C. elegans strains and RNA interference

C. elegans were cultured using standard conditions, and N2 was used as wild type (Brenner, 1974). Some nematode strains used in this work were provided by the *Caenorhabditis* Genetics Center (University of Minnesota, Minneapolis, MN), which is funded by the National Institutes of Health National Center for Research Resources. The *unc-84(n322)*, *unc-84(e1174)*, *unc-84(e1411)*, and *unc-84(n369)* alleles, from strains MT322, CB1174, CB1411, and MT369, respectively, were previously described (Malone *et al.*, 1999). The construct pSL589 was created by cloning the *XmaI/SphI* fragment of the *col-10* promoter from pOS12 (Spencer *et al.*, 2001) into pPD96.04 to specifically drive expression of NLS::GFP::LacZ in embryonic hypodermal cells. This construct, pSL589, was injected into N2 worms and then integrated (Kage-Nakadai *et al.*, 2012) to create strain UD469 (*ycls10[p_{col-10}NLS::gfp::lacZ*]). This outcrossed line was then crossed to *unc-84(n322)*, *unc-84(e1174)*, *unc-84(e1411)*, and *unc-84(n369)* strains to create UD414, UD412, UD396, and UD399, respectively. UD87, expressing the full-length UNC-84 rescue construct, was used as the starting strain for SAMP-1 localization studies in *lmn-1(RNAi)* (McGee *et al.*, 2006; Chang *et al.*, 2013). The *lem-2(tm1582)* deletion strain FX1582 and the *samp-1(tm2710)* deletion strain FX2710 were kindly supplied by Shohei Mitani (National Bioresource Project at the Tokyo Women's Medical University, Tokyo, Japan). *samp-1(tm2710)* was balanced with *mIn1 [dpy-10(e128) mIs14] II* from strain BS3493 to make strain UD470. The *p_{lbp-1} lmn-1::gfp* strain UD324 was previously described (Fridolfsson and Starr, 2010). UD324 was crossed to the *unc-84(n369)* and *unc-84(e1411)* strains to create UD437 and UD436, respectively.

For the *lmn-1(RNAi)* experiments, *ycls10* animals in an otherwise wild-type background were staged for 24 h post L4 at 20°C and fed bacteria expressing dsRNA against *lmn-1* from the Ahringer library (Fraser *et al.*, 2000). Worms were transferred after 24 h, and the plate from the first 24 h was counted at 48 h. For *emr-1(RNAi)*; *lem-2(1582)* treatment the first 24 h was discarded

and the second day was counted at 72 h (Meyerzon *et al.*, 2009b). Empty vector L4440 was used as control RNA interference (RNAi) for all feeding experiments. For *baf-1(RNAi)*, dsRNA was transcribed in vitro from the EST yk333d11 and subsequently injected into young adults (Meyerzon *et al.*, 2009b). Progeny laid from 12 to 24 h postinjection were screened. RNAi efficiency was assessed by progeny inviability.

Yeast two-hybrid assay

Constructs expressing GAL-4 DNA-binding domain::UNC-84 fusion proteins for yeast two-hybrid baits were created by amplifying inserts with PCR from the *unc-84* cDNA, yk402g1 (Kohara, 1996; McGee *et al.*, 2006), and cloning the inserts into pDEST32 using Gateway Technology (Invitrogen, Grand Island, NY). pSL242 expresses residues 1–385 of UNC-84, pSL244 has 59–385, pSL593 has 1–100, pSL592 has 1–59, and pSL595 has 385–510. The P91S mutation was introduced into pSL242 using PCR SOEing to create the mutant bait construct pSL596. The ProQuest *C. elegans* mixed-stage cDNA library (Invitrogen) was screened using the UNC-84(1-385) as a bait as previously described (Fridolfsson *et al.*, 2010). Positives with candidate interacting partners were selected on SD-Trp-Leu-His. To map the LMN-1 interaction domain of UNC-84, full-length LMN-1 prey, pSL719, obtained from the screen, was transformed into yeast strain Y187 (Clontech Laboratories, Mountain View, CA). The various UNC-84 baits were transformed into yeast strain Y2HGold (Clontech Laboratories). The bait strains were then mated to the prey-containing Y187 strains. Spot assays were conducted by spotting 2 μ l of yeast serial dilutions; growth was then imaged with an Alphamager 3400 (Alpha Innotech Corporation, San Leandro, CA). Liquid β -galactosidase assays were conducted following Clontech protocol PT1020-1 (Schneider *et al.*, 1996).

Nuclear migration assays

To count abnormally positioned hyp7 nuclei in the dorsal cord, L1 animals carrying *ycls10[nls::gfp::lacZ]* were examined on a 2% agar pad in M9 solution with 1 mM tetramisole (Starr *et al.*, 2001). DIC time-lapse imaging of migrating embryonic hypodermal nuclei was performed as previously reported (Fridolfsson and Starr, 2010). Images were taken every 15 s; time 0 was defined as the time that cell 12 hit the opposite seam cell boundary (Fridolfsson and Starr, 2010). Nuclear position was determined by measuring the leading edge, lagging edge, and midpoint of migrating nuclei using the manual tracking plug-in for ImageJ (National Institutes of Health, Bethesda, MD). The distance a nucleus migrated in the first 10 min determined in a cell-specific manner such that time 0 was when the respective cell's leading edge reached the opposite seam cell boundary. Images were obtained using DIC optics and a 63 \times Plan Apo 1.40 numerical aperture (NA) objective on a Leica DM6000 compound microscope equipped with a Leica DC350 camera operated by Leica LAS AF software (Leica Microsystems, Wetzlar, Germany). Images were processed using ImageJ.

LMN-1::GFP time-lapse images were captured at a frame rate of 1/s using the 488-nm laser on a spinning disk confocal microscope (Marianas Real Time Confocal SDC Workstation; Intelligent Imaging Innovations, Denver, CO), using a scan head (CSU-X1; Yokogawa, Tokyo, Japan), an autofocus laser, Definite Focus (Zeiss, Oberkochen, Germany), and a modified electron multiplying charge-coupled device camera (Cascade QuantEM 512SC; Photometrics). A 63 \times /NA 1.4 oil immersion objective was used (Zeiss). Images were acquired using SlideBook software (version 5.5; Intelligent Imaging Innovations).

Antibodies and immunofluorescence

Polyclonal antibodies (HJA2) against the *C. elegans* SAMP-1 were raised in rabbits injected with a histidine-NusA-tagged C-terminal fragment of SAMP-1 comprising residues 338–555. SAMP-1 antibodies were affinity purified using the same C-terminal SAMP-1 fragment tagged with GST. For immunofluorescence, embryos were extruded from hermaphrodites, permeabilized by the freeze-crack method, fixed for 10 min in -20°C methanol, blocked in PBST (phosphate-buffered saline [PBS] and 0.1% Triton X-100) with 5% milk, and stained as previously described (Miller and Shakes, 1995). The rabbit antibody against SAMP-1 was diluted 1:200 in PBS. UNC-83 monoclonal 1209D7D5 was used undiluted (Starr *et al.* 2001). UNC-84 IgM monoclonal L 72 6 was diluted 1:100 in PBS (Cain *et al.*, 2014). Alexa Fluor antibodies 594 donkey anti-rabbit immunoglobulin G (IgG), goat 488 anti-mouse IgM, and 488 goat anti-mouse IgG diluted 1:500 (Life Technologies, Carlsbad, CA) were used as secondary antibodies. DNA was visualized by a 5-min stain in 1 $\mu\text{g}/\text{ml}$ DAPI in PBS.

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