Kinesin-1 mediates translocation of the meiotic spindle to the oocyte cortex through KCA-1, a novel cargo adapter

Hsin-ya Yang, 1 Paul E. Mains, 2 and Francis J. McNally 1

1 Section of Molecular and Cellular Biology, University of California, Davis, Davis, CA 95616
2 Department of Biochemistry and Molecular Biology, Genes and Development Research Group, University of Calgary, Calgary, Alberta T2N 4N1 Canada

In animals, female meiotic spindles are attached to the egg cortex in a perpendicular orientation at anaphase to allow the selective disposal of three haploid chromosome sets into polar bodies. We have identified a complex of interacting Caenorhabditis elegans proteins that are involved in the earliest step in asymmetric positioning of anastral meiotic spindles, translocation to the cortex. This complex is composed of the kinesin-1 heavy chain orthologue, UNC-116, the kinesin light chain orthologues, KLC-1 and -2, and a novel cargo adaptor, KCA-1.

Depletion of any of these subunits by RNA interference resulted in meiosis I metaphase spindles that remained stationary at a position several micrometers from the cell cortex during the time when wild-type spindles translocated to the cortex. After this prolonged stationary period, unc-116(RNAi) spindles moved to the cortex through a partially redundant mechanism that is dependent on the anaphase-promoting complex. This study thus reveals two sequential mechanisms for translocating anastral spindles to the oocyte cortex.

Introduction

Meiosis is essential for sexual reproduction in all eukaryotes. During meiosis, a replicated diploid genome is reduced to four haploid genomes through two consecutive rounds of chromosome segregation. In females of both plants and animals, three haploid genomes are discarded and only one haploid genome is inherited by a female gamete. In animals, this asymmetric inheritance is mediated by spindles that are attached by one pole to the oocyte cortex during anaphase. In most animals, this perpendicular attachment to the cortex allows segregation of three haploid genomes into tiny cells called polar bodies and segregation of one haploid genome into a large egg, thus reserving almost all of the oocyte’s cytoplasm for embryo development (Selman, 1966; Maro and Verlhac, 2002).

Detailed studies of meiotic spindle movements in mouse (Maro et al., 1984, 1986; Verlhac et al., 2000) and Caenorhabditis elegans (Yang et al., 2003) have revealed a conserved series of movements that include translocation of the spindle to the cortex and rotation of the spindle from a parallel to a perpendicular orientation to allow chromosome segregation into a polar body. Movement and orientation of mitotic spindles in animals and fungi is thought to occur through astral microtubules that emanate from centriole-containing centrosomes or spindle pole bodies (Gonczy, 2002; Sheeman et al., 2003). However, the female meiotic spindles of humans (Sathananthan, 1997), cows (Navara et al., 1994), mice (Gueth-Hallonet et al., 1993), Drosophila melanogaster (Theurkauf and Hawley, 1992), and C. elegans (Albertson and Thomson, 1993) do not have centrioles or their associated astral microtubule arrays.

Mice and C. elegans have evolved different mechanisms for translocating their meiotic spindles to the oocyte cortex in the absence of astral microtubule arrays. Translocation of the mouse meiosis I spindle to the cortex is dependent on F-actin and c-mos but does not require microtubules (Verlhac et al., 2000). In contrast, we have previously shown that translocation of the C. elegans meiosis I spindle is dependent on microtubules and the microtubule-severing enzyme MEI-1 but is not dependent on F-actin (Yang et al., 2003). In both cases, the mechanism that polarizes cytoskeletal filaments toward the cortex and the mechanism of movement are unknown.

The microtubule dependence of C. elegans meiotic spindle translocation suggested that one or more microtubule motor proteins would be required either to establish bipolarity of spindle microtubules or to directly transport the spindle on the cytoplasmic microtubule array. To identify this motor (or motors), we initiated an RNA interference (RNAi) screen of the 23
microtubule motor subunits encoded in the *C. elegans* genome. We identified UNC-116, the kinesin-1 heavy chain (Patel et al., 1993), as essential for normal translocation of the meiotic spindle to the cortex. Because meiotic spindle structure appears normal in UNC-116–depleted embryos, this result suggests that the spindle is translocated on the acentrosomal cytoplasmic microtubule array. Such directional transport on an acentrosomal microtubule array is also observed during mRNA localization in *D. melanogaster* oocytes (Cha et al., 2001) and vesicle transport in plant cells (Gunning and Steer, 1996).

### Results

In *unc-116* (RNAi) embryos, the meiotic spindle remains stationary during the period when wild-type spindles translocate to the cortex

To determine whether or not microtubule motor proteins are involved in the translocation of the meiotic spindle to the oocyte cortex in *C. elegans*, we analyzed meiotic spindle translocation in worms depleted of different motor subunits by RNAi. There are 21 kinesin motor-domain homologues and two dynein heavy chain subunits encoded in the *C. elegans* genome (Wormbase). We recorded time-lapse sequences of meiotic spindle movements in worms expressing GFP-tubulin and that were treated with double-stranded RNA corresponding to seven different kinesin motor-domain homologues (UNC-116/R05D3.7, KLP-3/T09A5.2, KLP-7/K11D9.1, BMK-1/F23B12.8, KLP-15/M01E11.6, KLP-18/C06G3.2, and KLP-20/Y50D7A.6). Defective meiotic spindle translocation was observed only in *unc-116* (RNAi) worms. UNC-116 is the *C. elegans* orthologue of kinesin-1 heavy chain (Patel et al., 1993; Lawrence et al., 2004). Maturing *C. elegans* oocytes move into a somatic structure called the spermatheca after germinal vesicle breakdown, and then squeeze out of the other side of the spermatheca into the uterus. In the examples shown in Fig. 1 (A and B), the wild-type spindle contacted the cortex 42 s after exit from the spermatheca, whereas the *unc-116* (RNAi) spindle contacted the cortex 8.2 min after exit from the spermatheca. Similar results were obtained from 27/27 time-lapse sequences of wild-type worms and 15/19 time-lapse sequences of *unc-116* (RNAi) worms (Table I).

Careful analysis of the distance of the meiotic spindle from the cortex over time (Fig. 1, C and D) revealed that *unc-116* (RNAi) spindles remain completely stationary for up to 8 min after exit from the spermatheca, whereas wild-type spindles begin movement toward the cortex before the zygote exits from the spermatheca. An example of this prolonged stationary...
period is clearly seen in Fig. 1 B from 0 to 6.0 min. These observations suggested that an early translocation mechanism that is activated while the zygote is in the spermatheca of wild-type worms is completely absent in unc-116(RNAi) worms and that a distinct mechanism is responsible for the late movement of unc-116(RNAi) spindles to the cortex.

Consistent with this hypothesis, the orientation of spindles moving toward the cortex was different between unc-116(RNAi) and wild-type spindles. In wild-type worms, 14/17 (82.4%) of the meiotic spindles approached the cortex in a sideways orientation (e.g., Fig. 1 A). After the prolonged stationary period, 10/12 (83.3%) unc-116(RNAi) spindles moved toward the cortex with one pole leading (e.g., Fig. 1 B). The velocity of late translocation in unc-116(RNAi) was also different than the velocity of early translocation in wild-type worms (Table I). Thus, unc-116(RNAi) worms appear to be completely defective in an early translocation mechanism but retain a distinct, late translocation mechanism.

The movement of unc-116(RNAi) spindles is mediated by a distinct, anaphase-promoting complex (APC)-dependent mechanism

If movement of unc-116(RNAi) spindles to the cortex is indeed due to a discrete, late mechanism, this should be revealed by distinct genetic requirements for the early and late mechanisms. In a previous study (Yang et al., 2003), we showed that an early translocation mechanism that is activated while the zygote is in the spermatheca of wild-type worms is completely absent in unc-116(RNAi) worms and that a distinct mechanism is responsible for the late movement of unc-116(RNAi) spindles to the cortex.

However, in 9/9 unc-116(RNAi) time-lapse sequences, where the spindle was oriented such that both translocation and shortening could be measured, translocation initiated at the same time that spindle shortening initiated and spindles arrived at the cortex 3.6 ± 1.1 min after shortening started (Fig. 1 D). As previously reported for worms depleted of the FZY-1 subunit of the APC (Yang et al., 2003), spindle shortening was blocked in mat-2(ts) worms at nonpermissive temperature but spindle translocation occurred with wild-type kinetics. (Fig. 1 E, n = 5). In 5/5 of mat-2(ts); unc-116(RNAi) double mutant worms, spindle length did not change and spindle translocation to the cortex was completed blocked (Fig. 1 F). The complete block to spindle translocation in mat-2(ts); unc-116(RNAi) double mutant worms was confirmed by single time point analysis (Fig. 1 H) to eliminate any contribution by photodamage. These results dem-

Table I. Velocities of meiotic spindle movements

<table>
<thead>
<tr>
<th></th>
<th>Wild type</th>
<th>unc-116 (RNAi)</th>
<th>klc-1 (RNAi); klc-2 (RNAi)</th>
<th>C10H11.10</th>
<th>unc-116 (sb79r;h24)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Movies</td>
<td>27</td>
<td>15</td>
<td>4</td>
<td>11</td>
<td>5</td>
</tr>
<tr>
<td>Time from GVBD to pronucleus formation (min)</td>
<td>42.2 ± 5.1</td>
<td>38.5 ± 5.0</td>
<td>39.4 ± 0.8</td>
<td>42.0 ± 5.7</td>
<td>39.0 ± 1.3</td>
</tr>
<tr>
<td>Diakinesis nuclear positioning (dorsal-ventral axis)</td>
<td>46%</td>
<td>47%</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>Meiosis I</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Maximum spindle length (µm)</td>
<td>8.0 ± 0.7</td>
<td>8.2 ± 0.7</td>
<td>8.5 ± 0.3</td>
<td>8.0 ± 0.4</td>
<td>8.1 ± 0.6</td>
</tr>
<tr>
<td>Maximum spindle width (µm)</td>
<td>6.3 ± 0.5</td>
<td>6.4 ± 0.5</td>
<td>7.2 ± 0.6</td>
<td>5.9 ± 0.4</td>
<td>6.1 ± 0.3</td>
</tr>
<tr>
<td>Distance from spindle to cortex after exit from spermatheca (µm)</td>
<td>2.3 ± 2.1</td>
<td>7.2 ± 0.1</td>
<td>5.4 ± 1.6</td>
<td>6.8 ± 1.5</td>
<td>5.3 ± 1.8</td>
</tr>
<tr>
<td>Timing of cortical contact after exit from spermatheca (min)</td>
<td>1.0 ± 0.7</td>
<td>8.3 ± 2.2</td>
<td>10.9 ± 4.5</td>
<td>10.0 ± 2.0</td>
<td>9.2 ± 1.5</td>
</tr>
<tr>
<td>Translocation rate (µm/min)</td>
<td>0.9 ± 0.4</td>
<td>3.5 ± 1.2</td>
<td>3.2 ± 0.6</td>
<td>4.6 ± 1.8</td>
<td>2.0 ± 0.8</td>
</tr>
<tr>
<td>Time from GVBD to first spindle shortening (min)</td>
<td>18.7 ± 3.3</td>
<td>13.3 ± 1.9</td>
<td>17.8 ± 6.1</td>
<td>16.9 ± 2.0</td>
<td>15.2 ± 4.2</td>
</tr>
<tr>
<td>Shortening rate (µm/min)</td>
<td>0.8 ± 0.2</td>
<td>1.2 ± 0.3</td>
<td>0.9 ± 0.2</td>
<td>1.2 ± 0.4</td>
<td>0.7 ± 0.2</td>
</tr>
<tr>
<td>Chromosome segregation rate (µm/min)</td>
<td>0.7 ± 0.1</td>
<td>1.1 ± 0.5</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>(n = 11)</td>
<td>(n = 5)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Meiosis II</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Maximum spindle length (µm)</td>
<td>6.3 ± 0.5</td>
<td>6.6 ± 0.4</td>
<td>6.7 ± 0.5</td>
<td>6.7 ± 0.9</td>
<td>6.5 ± 0.5</td>
</tr>
<tr>
<td>Maximum spindle width (µm)</td>
<td>4.9 ± 0.5</td>
<td>5.0 ± 0.6</td>
<td>4.0 ± 0.5</td>
<td>4.7 ± 0.5</td>
<td>5.0 ± 0.4</td>
</tr>
<tr>
<td>Distance to cortex after spindle assembly (µm)</td>
<td>0.9 ± 0.4</td>
<td>3.4 ± 1.2</td>
<td>2.1 ± 0.5</td>
<td>2.9 ± 1.3</td>
<td>4.0 ± 2.6</td>
</tr>
<tr>
<td>Timing of cortical contact after spindle assembly (min)</td>
<td>0.7 ± 0.4</td>
<td>6.0 ± 0.9</td>
<td>7.2 ± 1.8</td>
<td>8.5 ± 2.5</td>
<td>7.1 ± 1.0</td>
</tr>
<tr>
<td>Translocation rate (µm/min)</td>
<td>1.1 ± 0.6</td>
<td>4.6 ± 1.5</td>
<td>2.7 ± 2.0</td>
<td>4.7 ± 1.1</td>
<td>4.8 ± 2.5</td>
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<tr>
<td>Time from GVBD to second spindle shortening (min)</td>
<td>33.5 ± 2.8</td>
<td>27.6 ± 3.1</td>
<td>30.1 ± 1.1</td>
<td>32.1 ± 4.5</td>
<td>28.3 ± 1.5</td>
</tr>
<tr>
<td>Shortening rate (µm/min)</td>
<td>0.8 ± 0.2</td>
<td>1.5 ± 0.4</td>
<td>1.5 ± 0.6</td>
<td>1.3 ± 0.4</td>
<td>1.3 ± 0.4</td>
</tr>
<tr>
<td>Chromosome segregation rate (µm/min)</td>
<td>1.1 ± 0.6</td>
<td>1.2 ± 0.6</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>(n = 11)</td>
<td>(n = 5)</td>
<td></td>
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</tr>
</tbody>
</table>

aGVBD denotes germinal vesicle breakdown or nuclear envelope breakdown.

bThe change in pole–pole spindle length over time.

cThe change in distance between two sets of GFP-histone–labeled chromosomes over time.
demonstrated that the late spindle translocation observed in unc-116(RNAi) worms is APC dependent and kinesin-1 independent whereas the early translocation mechanism observed in wild-type worms is APC independent and kinesin-1 dependent.

**UNC-116 is not required for other aspects of meiosis**

We previously demonstrated that both tubulin and the katanin orthologue MEI-1 are required for meiotic spindle translocation (Yang et al., 2003). In both of these cases, however, spindle structure was severely perturbed, making it impossible to discern if MEI-1 is directly involved in spindle translocation or if normal spindle architecture is required for translocation. In contrast, unc-116(RNAi) meiotic spindles observed by GFP-tubulin fluorescence had a wild-type structure (Fig. 1, B and H). These spindles had wild-type length and width (Table I), exhibited normal anaphase chromosome segregation (Fig. 2 and Table I), and shortened with normal kinetics (Table I and Fig. 1 D). These results indicate that UNC-116 is primarily (if not exclusively) required in the early embryo for meiotic spindle translocation.

**UNC-116 is also essential for cortical positioning of the meiosis II spindle**

The early, APC-independent and UNC-116–dependent translocation mechanism might be activated only transiently or it might be active throughout meiosis. Careful observation of meiosis II spindles in unc-116(RNAi) worms indicated that the UNC-116–dependent mechanism continues to be active during meiosis II. Whereas the meiosis II spindle in a wild-type embryo is already extremely close to the cortex as it assembles (Fig. 1 C and Video 1, available at http://www.jcb.org/cgi/content/full/jcb.200411132/DC1), the meiosis II spindle in an unc-116 (RNAi) embryo remains stationary several micrometers from the cortex (Fig. 1 D, Table I, and Video 2, available at http://www.jcb.org/cgi/content/full/jcb.200411132/DC1) until the meiosis II spindle begins to shorten. Meiosis II spindles in unc-116(RNAi) embryos also move in a pole-first orientation with a high velocity (Table I). These results indicate that the force generated on the spindle by wild-type UNC-116 is active throughout meiosis I and meiosis II. It is likely that the same APC-dependent mechanism that mediates movement of meiosis I spindles to the cortex in the absence of UNC-116 also mediates the delayed translocation of meiosis II spindles to the cortex.

**UNC-116 is essential maternally because it is required for polar body formation**

To determine the consequences of delayed spindle translocation, we filmed GFP-histone in unc-116(RNAi) worms to track chromosome movements during meiosis. In 5/5 cases where delayed spindle translocation was observed, anaphase chromosome segregation occurred normally with one set of chromosomes being pushed into the cortex during both anaphase I and II (Fig. 2). However, failures in polar body formation during meiosis I, meiosis II, or both were frequently observed. In the example shown in Fig. 2, chromosomes that segregated into the cortex at meiosis I (21 min) collapsed back into the embryo so that 12 rather than 6 chromosomes were present in the meiosis II spindle (24.3 min). In this example, meiosis II polar body formation was successful (36.3 min). In 60% of GFP-histone unc-116(RNAi) movies, one or both polar bodies failed to form, even though anaphase occurred at the cortex. In contrast, both polar bodies formed successfully in 100% of GFP-histone wild-type time-lapse sequences (n = 11). To eliminate any contribution from photodamage, polar body number was also scored by fixed time point analysis. A failure to form one or both polar bodies was observed in 67% of unc-116(RNAi) embryos compared with 5% of wild-type embryos (see Table III). A failure to form one or both polar bodies should result in embryos that are triploid or pentaploid. Whereas triploid worms can be viable (Madl and Herman, 1979), pentaploidy would be expected to cause embryonic lethality later in development. Indeed, 67% embryonic lethality was observed among embryos laid by unc-116(RNAi) worms (Table II). These results indicate that UNC-116 and the early spindle translocation pathway are required for meiotic cytokinesis, even though the spindles always achieve perpendicular cortical contact eventually.

Because kinesin-1 has never been found to be essential for cell division in other species (Wright et al., 1993; Brendza...

19 for www.jcb.org/cgi/content/full/jcb.200411132/DC1) versus 15/ location in only 7/14 worms (Fig. S2, available at http://

worms revealed a block in preanaphase meiotic spindle trans-

tion) of UNC-116 activity. Therefore, we analyzed a recessive, loss-of-function methods). Homozygous unc-116(RNAi) and because RNAi can result in depletion of homologous mRNAs, we wished to confirm that the spindle translocation defect and the maternal-effect embryonic lethality were actually due to a reduction in the amount of UNC-116 activity. Therefore, we analyzed a recessive, loss-of-function allele of unc-116, unc-116(rh24sb79) (see Materials and methods). Homozygous unc-116(rh24sb79) worms are extremely uncoordinated and exhibited 28% maternal-effect embryonic lethality at 25°C compared with 67% for unc-116(RNAi) (Table II), indicating that unc-116(rh24sb79) worms retain more UNC-116 activity than unc-116(RNAi) worms. Western blotting with an UNC-116–specific antibody revealed more intact UNC-116 polypeptide in unc-116(rh24sb79) adult hermaphrodites (Fig. S1, lane 8, available at http://www.jcb.org/cgi/content/full/jcb.2004111132/DC1) than in unc-116(RNAi) adults (Fig. S1, lane 2), confirming the hypomorphic classification of this allele.

Consistent with the hypomorphic nature of the rh24sb79 allele, time-lapse imaging of GFP-tubulin unc-116(rh24sb79) worms revealed a block in preanaphase meiotic spindle translocation in only 7/14 worms (Fig. S2, available at http://www.jcb.org/cgi/content/full/jcb.2004111132/DC1) versus 15/19 for unc-116(RNAi). Significantly, the combination of RNAi and this loss-of-function allele did not increase the level of embryonic lethality beyond that of unc-116(RNAi) alone (Table II). This finding strongly suggests that unc-116(RNAi) indeed represents the unc-116(null) embryonic phenotype. We conclude that both the spindle translocation defect and the maternal-effect embryonic lethality observed in unc-116(RNAi) worms is in fact due to a severe reduction (if not total elimination) of UNC-116 activity.

**Table II. Embryonic lethality due to depletion of UNC-116 and associated proteins**

<table>
<thead>
<tr>
<th>Genotype</th>
<th>Embryonic lethality</th>
<th>Brood size</th>
</tr>
</thead>
<tbody>
<tr>
<td>wild type</td>
<td>3.3 ± 4 (9,999/95)</td>
<td>174.4</td>
</tr>
<tr>
<td>wild type + males</td>
<td>2.6 ± 0.4 (4,509/28)</td>
<td>241.6</td>
</tr>
<tr>
<td>unc-116(RNAi)</td>
<td>66.9 ± 19 (2,230/45)</td>
<td>71.8</td>
</tr>
<tr>
<td>unc-116(RNAi) + males</td>
<td>65.3 ± 26 (1,904/16)</td>
<td>161.6</td>
</tr>
<tr>
<td>unc-116(rh24, sb79)</td>
<td>27.8 ± 9 (1,099/23)</td>
<td>57.5</td>
</tr>
<tr>
<td>unc-116(rh24, sb79) + males</td>
<td>24.0 ± 9 (800/24)</td>
<td>40.4</td>
</tr>
<tr>
<td>unc-116(rh24, sb79, RNAi)</td>
<td>72.5 ± 6 (560/24)</td>
<td>28.5</td>
</tr>
<tr>
<td>unc-116(rh24, sb79, RNAi) + males</td>
<td>58.2 ± 11 (756/23)</td>
<td>39.2</td>
</tr>
<tr>
<td>klc-1(RNAi)</td>
<td>16.4 (785/10)</td>
<td>156.1</td>
</tr>
<tr>
<td>klc-1(RNAi) + males</td>
<td>11.7 (725/9)</td>
<td>132.2</td>
</tr>
<tr>
<td>klc-2(RNAi)</td>
<td>1.7 (879/9)</td>
<td>166.1</td>
</tr>
<tr>
<td>klc-2(RNAi) + males</td>
<td>4.3 (836/7)</td>
<td>155.3</td>
</tr>
<tr>
<td>klc-1(RNAi); klc-2(RNAi)</td>
<td>65.8 (912/17)</td>
<td>113.4</td>
</tr>
<tr>
<td>klc-1(RNAi); klc-2(RNAi) + males</td>
<td>64.6 (1,584/10)</td>
<td>221.6</td>
</tr>
<tr>
<td>C10H11.10/kc-1(RNAi)</td>
<td>89.2 ± 9 (1,719/54)</td>
<td>73.4</td>
</tr>
<tr>
<td>C10H11.10/kc-1(RNAi) + males</td>
<td>69.0 ± 20 (2,660/30)</td>
<td>121.7</td>
</tr>
</tbody>
</table>

Embryonic lethality is the percentage of eggs that did not hatch within 12 h. Numbers in parentheses indicate the number of embryos counted/number of worms tested by RNAi soaking. + males indicates that male worms were added to provide wild-type sperm. Brood size is the average number of eggs laid by a single worm in 48 h. Wild-type is the GFP-tubulin–expressing strain Wh204.

The kinesin light chains KLC-1 and -2 are required for normal translocation of the meiotic spindle to the cortex. Images of GFP-tubulin fluorescence are shown from representative time-lapse sequences of a meiotic embryo within a klc-

1(RNAi) worm [A] and a klc-1(RNAi); klc-2(RNAi) worm [B]. The cell cortex was highlighted in each image for clarity. In both cases, the meiosis I and II spindles do not move toward the cortex until after spindle shortening has initiated. Asterisks indicate exit from the spermatheca. (C) Fixed time point image of a mat-2(ts); klc-1(RNAi); klc-2(RNAi) triple mutant worm shows a meiotic spindle arrested far from the cortex. Bars, 10 μm.

Kinesin-1 purified from a variety of species, including C. elegans, consists of a tetramer with two heavy chain and two light chain subunits (Vale et al., 1985; Saxton et al., 1988; Signor et al., 1999). To test whether or not early meiotic spindle translocation depends on a kinesin-1 heavy chain/light chain complex, we acquired GFP-tubulin time-lapse sequences from worms treated with dsRNA corresponding to either of the C. elegans kinesin light chain homologues, KLC-1 or KLC-2. A block to early spindle translocation was observed in 3/6 klc-1(RNAi) worms, 2/7 klc-2(RNAi) worms, and 4/5 doubly treated klc-

1(RNAi); klc-2(RNAi) worms (Fig. 3, A and B; and Table I). In addition, depletion of kinesin light chains in metaphase-arrested mat-2(ts) embryos resulted in spindles arrested far from the cortex (Fig. 3 C). Thus, the phenotype of klc-1(RNAi) or klc-2(RNAi) worms was qualitatively the same, but quantitatively weaker, than the phenotype of unc-116(RNAi) embryos. Western blots of klc-2(RNAi) worms probed with a KLC-2–specific antibody revealed that considerable KLC-2 protein
product remains in these worms (Fig. S1). Thus the weaker phenotypes observed were at least in part due to the incomplete effectiveness of the RNAi. Simultaneous treatment of worms with dsRNAs corresponding to both light chains (klc-1(RNAi); klc-2(RNAi)) resulted in embryonic lethality (Table II), polar body defects (Table III), and translocation defects that were quantitatively similar to unc-116(RNAi) worms. These results indicate that the two kinesin light chains act redundantly and that a complex of kinesin heavy and light chains is essential for the preanaphase translocation of the meiotic spindle.

KCA-1 mediates UNC-116-dependent spindle translocation

In neurons, a membrane-associated cargo-adaptor protein called sunday driver forms a bridge between kinesin light chains and vesicles transported during fast axonal transport (Bowman et al., 2000). To identify a possible cargo-adaptor protein linking KLC-1 or KLC-2 to the meiotic spindle, we used RNAi to screen putative kinesin light chain interacting proteins identified in a large scale yeast two hybrid study (Li et al., 2004). Treatment of mat-2(ts), GFP-tubulin worms with dsRNA corresponding to one of these interacting gene products (C10H11.10), resulted in meiotic spindles that were far from the cortex of metaphase-arrested embryos (Fig. 4 B). In contrast, all meiotic spindles were at the cortex of metaphase-arrested embryos of mat-2(ts), GFP-tubulin worms treated with other dsRNAs (unpublished data). Time-lapse imaging of GFP-tubulin in 11/11 C10H11.10(RNAi) single mutant worms revealed a block in preanaphase spindle translocation identical to that observed in unc-116(RNAi) or klc-1(RNAi); klc-2(RNAi) worms (Fig. 4 A and Table I). C10H11.10(RNAi) worms also exhibited embryonic lethality (Table II) and polar body defects (Table III) similar to those seen in unc-116(RNAi) and klc-1(RNAi); klc-2(RNAi) worms. These data are consistent with a mechanism in which the C10H11.10 gene product forms a bridge between the UNC-116–KLC-1,2 kinesin and the meiotic spindle. We have therefore given C10H11.10 the name kca-1, for kinesin cargo adaptor.

To directly demonstrate the existence of a protein complex containing UNC-116, KLC-2, and KCA-1, all three proteins were expressed as fusion proteins in E. coli and interactions were analyzed by glutathione Sepharose chromatography. As shown in Fig. 5 A (lane 10), a chitin-binding domain fusion to the UNC-116 stalk-tail domains stochiometrically copurified with a glutathione S–transferase fusion to KCA-1 only in the presence of 6his-KLC-2b. This experiment demonstrated that KLC-2b can either bind simultaneously to both UNC-116 and KCA-1 or that KLC-2b induces a conformational change that increases the affinity between UNC-116 and KCA-1. Similar experiments with KCA-1 deletion derivatives (Fig. 5 B) indicated that the NH2-terminal 155 amino acids of KCA-1 contain the KLC-2 binding domain. This result implies that the COOH-terminal part of KCA-1 may bind to some spindle component to generate a bridge between UNC-116 and the meiotic spindle. KCA-1 has no homology with sunday driver or other known kinesin-associated proteins and thus represents a novel kinesin cargo adaptor.

<table>
<thead>
<tr>
<th>Genotype</th>
<th>Fraction of embryos with one or no polar bodies</th>
<th>Number of embryos scored</th>
</tr>
</thead>
<tbody>
<tr>
<td>wild type</td>
<td>5.6</td>
<td>54</td>
</tr>
<tr>
<td>unc-116(RNAi)</td>
<td>67.7</td>
<td>31</td>
</tr>
<tr>
<td>klc-1(RNAi)</td>
<td>38.5</td>
<td>26</td>
</tr>
<tr>
<td>klc-2(RNAi)</td>
<td>28.1</td>
<td>31</td>
</tr>
<tr>
<td>klc-1(RNAi); klc-2(RNAi)</td>
<td>55.6</td>
<td>27</td>
</tr>
<tr>
<td>C10H11.10/kca-1(RNAi)</td>
<td>72.7</td>
<td>44</td>
</tr>
</tbody>
</table>

Hermaphrodites expressing GFP-histone were soaked in the indicated dsRNAs for 16 h at 25°C, and polar bodies were counted in pronuclear stage embryos 24 h after returning the worms to bacteria.

Figure 4. C10H11.10/KCA-1 is required for normal translocation of the meiotic spindle to the cortex. (A) Images of GFP-tubulin fluorescence are shown from a representative time-lapse sequence from a meiotic embryo within a kca-1(RNAi) worm. The cell cortex was highlighted in each image for clarity. The meiosis I and meiosis II spindles do not move toward the cortex until after spindle shortening has initiated. The asterisk indicates exit from the spermatheca. (B) Fixed time point image of mat-2(ts); kca-1(RNAi) worms shows a meiotic spindle arrested far from the cortex. Bars, 10 μm.
Cytoplasmic dynein is not required for preanaphase spindle translocation

One hypothesis consistent with our results is that kinesin-1 transports the spindle on a subset of cytoplasmic microtubules that have plus ends oriented toward the cortex. An alternative model, suggested by work in *Aspergillus nidulans* (Zhang et al., 2003) and *D. melanogaster* (Brendza et al., 2002), is that UNC-116 is only required to localize the minus end–directed motor, cytoplasmic dynein. In this model, cytoplasmic dynein would be required to move the spindle on a subset of cytoplasmic microtubules that have minus ends oriented toward the cortex. Therefore, we tested if early meiotic spindle translocation is blocked when cytoplasmic dynein heavy chain, DHC-1, is depleted by RNAi. As previously reported (Gonczy et al., 1999), the strongest phenotype observed after prolonged treatment with dhc-1 dsRNA was failure to ovulate mature oocytes. Shorter treatment of worms with dhc-1(RNAi) resulted in meiosis I spindles with a variety of structural defects. Some spindles were longer than wild-type spindles and had extremely pointed poles, whereas others were extremely disorganized. Time-lapse imaging of GFP-tubulin–labeled spindles under these weak dhc-1(RNAi) conditions revealed that 7/7 spindles arrived at the cortex either before or within 1.0 ± 0.4 min (1.0 ± 0.7 min in wild type) after exit from spermatheca (Fig. 6 A). At longer time points of dhc-1 dsRNA treatment, before worms ceased to produce mature oocytes, multiple small diakinesis nuclei were observed in immature oocytes (Fig. 6 B). Time-lapse imaging of GFP-tubulin revealed that each of these nuclei gave rise to a spindle-like structure at germinal vesicle breakdown and that all of these structures were eventually associated with the cortex (Fig. 6 B, 30.5 min). Tracking the early movements of these spindles to determine the time of cortical contact, however, proved difficult and was not pursued. To determine whether or not the early, APC-independent translocation mechanism was functional in these dhc-1(RNAi) embryos, mat-2(ts), GFP-tubulin worms, were treated with dhc-1 dsRNA at 25°C so that metaphase-arrested meiotic embryos that were derived from oocytes with multiple germinal vesicles could be observed. The majority of meiotic spindles in these mat-2(ts); dhc-1(RNAi) double mutant embryos were associated with the cortex just as in mat-2(ts) single mutants (Fig. 6 C). These results provide no indication of a role for cytoplasmic dynein in the early, APC-independent spindle translocation. Thus, UNC-116 appears to play a more direct role in early spindle translocation, possibly by transporting the spindle on cytoplasmic microtubules that are oriented with their plus ends toward the cortex. An obvious challenge is elucidating how the actinomysyl cytoplasmic microtubule array shown in Fig. 6 D can be organized to allow directional transport.

**Discussion**

Two redundant pathways can move *C. elegans* meiotic spindles to the cortex

Our results indicate that in *C. elegans* female meiotic spindles are positioned at the cell cortex by two sequential mechanisms, an early, kinesin-1–dependent pathway that moves spindles in a sideways orientation and a late, APC-dependent pathway that pulls the spindle toward the cortex with one pole leading (Fig. 7). Wild-type meiotic spindles begin kinesin-1–dependent movement toward the cortex as soon as they have assembled into bipolar structures, while the embryo is still inside the spermatheca. In the absence of kinesin-1, the meiosis I spindle is stationary in the cytoplasm during the 7-min period before spindle shortening begins. The spindle then suddenly moves to the cortex with one pole leading just after spindle shortening initiates. We propose that the sudden movement of unc-116(RNAi) spindles to the cortex is mediated by the same mechanism as wild-type spindle rotation and that it is mediated
by microtubules extending from one spindle pole to the cortex. In wild-type spindles, pulling a spindle pole for only 2 μm would generate a rotation, whereas in unc-116(RNAi) spindles, the same pulling force would generate a pole-first translocation over a distance of 7 μm (Fig. 7). This model is supported by the finding that both wild-type spindle rotation and UNC-116–independent translocation are blocked when the APC is inhibited. The model is also supported by the finding that both wild-type rotation and UNC-116–independent translocation occur just after initiation of APC-dependent spindle shortening. Finally, UNC-116–dependent translocation seen in wild-type worms occurs with the spindle oriented parallel to the cortex, whereas UNC-116–independent translocation occurs with one pole leading. Direct visualization of microtubules extending from one pole to the cortex has thus far been stymied by the presence of a microtubule meshwork that fills the entire meiotic embryo. Two successive and partially redundant pathways also mediate mitotic spindle positioning in budding yeast (Schuyler and Pellman, 2001).

**The relationship between spindle translocation and polar body formation**

Both time-lapse imaging and fixed time point analysis revealed that one or both polar bodies failed to form in 67% of unc-116(RNAi) embryos. This finding could indicate that UNC-116 has a separate role in polar body formation. Alternatively, the delayed arrival of unc-116(RNAi) spindles at the cortex might sometimes be “just in time” and sometimes be “just too late” to induce polar body formation. We favor the latter explanation. The finding that treatment of unc-116(rh24sb79) mutant worms with unc-116 dsRNA did not increase embryonic lethality beyond the 67% seen in unc-116(RNAi) alone indicates that 67% embryonic lethality is the null embryonic phenotype for unc-116. Thus, in 33% of unc-116(RNAi) embryos, both polar bodies form in the absence of UNC-116 activity. Examples of this successful polar body formation were observed in time-lapse sequences where early spindle translocation was completely blocked, and the spindle reached the cortex by the late, pole-first mechanism. These results suggest that even in the complete absence of maternal UNC-116, the late, APC-dependent mechanism can sometimes move the spindle to the cortex just in time for polar body induction.

**KCA-1 as a novel cargo adaptor for the meiotic spindle**

We showed that the previously uncharacterized gene product C10H11.10/KCA-1 can form a stoichiometric protein complex with the kinesin light chain KLC-2b and the kinesin heavy chain UNC-116 in vitro. Furthermore, kca-1(RNAi) yields the same phenotype as depletion of heavy chain or depletion of
both light chains. These results indicate that KCA-1 may be the cargo adaptor that allows UNC-116 to form transient attachments to the meiotic spindle. The Caenorhabditis briggsae predicted gene CBG03328 (Wormbase) exhibits 40% amino acid identity over its entire length with KCA-1, but no homologues outside of nematodes or obvious sequence motifs are apparent. In the C. elegans global yeast two-hybrid analysis of Li et al. (2004), KCA-1 interacted only with the kinesin light chains KLC-1 and KLC-2 and with HPL-2. HPL-2 is one of two C. elegans orthologues of heterochromatin binding protein 1, a highly conserved component of pericentric heterochromatin (Couteau et al., 2002). If KCA-1 and HPL-2 interact in vivo, this interaction would complete a physical bridge between UNC-116 and the meiotic chromosomes. In this model, UNC-116 would transport meiotic chromosomes toward the cell cortex along cytoplasmic microtubules and the spindle would be dragged along with the chromosomes. This hypothetical mechanism would be consistent with the sideways orientation of spindles during wild-type spindle translocation.

Although our data support a model in which kinesin-1 transports the spindle along cytoplasmic microtubules, there is a 120-fold difference between the gliding velocity of purified C. elegans kinesin-1 (2 μm/s; Signor et al., 1999) and spindle translocation velocity (1 μm/min). The unexpectedly slow translocation might be explained if the UNC-116–KLC–KCA-1 complex actually acts indirectly, perhaps by organizing the cytoplasmic microtubule array. However, we favor a model in which translocation is slowed by opposing forces from kinesin-1 molecules that attempt to transport the spindle on microtubules with plus ends oriented away from the cortex.

Acentriolar spindles are positioned by unique mechanisms

Positioning of female meiotic spindles at the oocyte cortex is observed in all animal species, resulting in highly asymmetric divisions giving rise to small polar bodies and one large oocyte in most species. In some organisms such as Chaetopterus variopedatus (Lutz et al., 1988), Spisula solidissima (Palazzo et al., 1992), and starfish (Hamaguchi, 2001; Zhang et al., 2004), female meiotic spindles have robust astral microtubule arrays nucleated by centriole-containing centrosomes. In these species, it is likely that motor proteins associated with the cortex generate pulling forces on astral microtubules by the same mechanisms proposed for mitotic spindles in many species (Gonczy, 1992), and starfish (Hamaguchi, 2001; Zhang et al., 2004), female meiotic spindles require F-actin (Yang et al., 2003). If the interaction between the C. elegans heterochromatin protein HPL-2 and the kinesin cargo adaptor KCA-1 occurs in vivo, then kinesin-1 may transport meiotic chromosomes, dragging the spindle along with the chromosomes. In this case, mice and worms would be using identical mechanisms mediated by different molecules.

Materials and methods

C. elegans strains

In this study, wild type indicates the integrated GFP-tubulin strain WH204 (Strome et al., 2001) or the integrated GFP-histone H2b strain A2212 (Fraitis et al., 2001), both of which are derived from N2. The integrated GFP-tubulin from WH204 was introduced into strains carrying mat-2(ax76), mat-2(ax102), or unc-116(h24sb79) through crosses. mat-2 strains were obtained from D. Shakes [College of William and Mary, Williamsburg, VA].

In utero filming

Adult hermaphrodites were anesthetized with tricaine/tetramisole and mounted between a coverslip and a thin agarose pad as described previously (Yang et al., 2003). Stage temperature was 22–24°C. Images were acquired at 15-s intervals on an upright microscope (model Microphot SA; Nikon) using a 60× PlanApo 1.4 objective and a Photometrics Quantix/KAF1400 camera [Roper Scientific]. Illumination from an HBO100 light source was shuttered with a Univiblitz shutter. Time-lapse acquisition was controlled with IP Lab Spectrum software [Scanalytics]. Quantitative analysis of translocation velocities was also performed with IP Lab Spectrum.

RNAi by soaking

The cDNA inserts of plasmid cDNA clones were amplified by PCR using one primer with a 5′ T7 promoter extension and a second primer with a 5′ T3 promoter extension. Linear PCR products were concentrated using spin columns (Qiagen) and used as templates in separate T3 and T7 RNA polymerase transcription reactions (MegaScript T7 or T3 Kit; Ambion). After treatment with DNase I, RNAs were purified by LIC precipitation and annealed. 40–50 L4 worms were soaked in 10 μl of 1 mg/ml dsRNA dissolved in 66 mM K-PO4 and 10 mM K-citrate, pH 7.5, in a humidified chamber for 16 h at 25°C and then transferred to OP50-seeded plates to recover for 24 h before filming or lethality scoring. The following cDNA clones were used: KLC-1-yk1256g06, KLC-2-yk1323f03, C10H11.10-yk442h9, and UNC-116-yk255a4 (all obtained from Y. Kuhara, National Institute of Genetics, Mishima, Japan). C10H11.10 was initially identified using RNAi by feeding with clone I-2I09 from the genomic RNAi feeding library [MRC Gene Service; Kamath et al., 2003].

Antibody generation

Rabbits were immunized with inclusion bodies composed of 6his-UNC-116 stalk tail or 6his-KLC-2b. Antibodies were bound to 6his-UNC-116 or 6his-KLC-2b immobilized on nitrocellulose strips and eluted with 200 mM glycine, pH 2.4, before neutralization with Tris-Cl, pH 8.0.

Protein interaction experiments

Different fragments of KA1 were PCR amplified from the yak624e12 cDNA clone. In the amino acid numbering shown in Fig. 5, C corresponds to a methionine that is 43 aa downstream from the predicted start codon in Wormbase. These fragments were cloned into pET41a (Novagen), expressed in BL21(DE3) E. coli, and purified by Ni2+ chelate chromatography via the 6his tag on the GST encoded by pET41a. Full-length KLC-2b was PCR amplified from yak886b10 and cloned into pET28a (Novagen) for expression of 6his–KLC-2b and into pTYB12 (New England Biolabs, Inc.) for expression of CBD–KLC-2b. The COOH-terminal 442 aa of UNC-116 were PCR amplified from yak859d09 and cloned into pTYB12.
length MEI-1 was also PCR amplified and cloned into pTYB12. 6his-KLC-2b was purified by Ni²⁺ chelate chromatography after expression in E. coli. CBD-KLC-2b, CBD-UNC-116, and CBD-MEI-1 were expressed in E. coli, and high speed supernatants from microfluidized extracts were used directly in the binding assays without further purification after normalizing to equal concentrations of CBD fusion protein. 50 µL of glutathione Sepharose-4B beads were incubated with an excess of GST-fusion protein for 1 h, and then washed extensively. Coated beads were incubated with 1 ml of E. coli lysate containing the appropriate CBD fusion and 25 mM Tris 8.0, 1 M NaCl, 0.1% Triton X-100, and 5% glycerol for 2 h at 22°C with constant rocking. After extensive washing with the same buffer, bound complexes were eluted with SDS-Laemmli buffer.

Isolation of unc-116(rh245b79)

To identify a loss-of-function allele of unc-116(rh24), these were isolated as intragenic revertants that blocked the temperature-sensitive enhancement between rh24/+ and mei-1(ct46ds)/+. Both unc-116(rh24) and mei-1(ct46ds)/+ result in small, misoriented first cleavage spindles. At 20°C, rh24/+ hermaphrodites segregate 12% dead embryos, 78% are lethal from the segregation of holocentric chromatin. At 25°C, these are 46% dead embryos, 47% are lethal from the segregation of holocentric chromatin. At 20°C, mei-1(ct46ds)/+ fail to hatch. Replacing rh24 with a chromosomal deficiency of the region in rh24/C. A total of 7,200 F1 progeny were shifted, of which 1,418 [1,800] were the relevant rh24/++; mei-1(ct46ds)/+ genotype. Six plates produced an F2 generation, indicating the presence of an F1 animal with a suppressor. One of these contained sb79, which is tightly linked to the original rh24 mutation. Three-factor mapping indicated that sb79 is <0.12 cm from rh24, consistent with an intragenic event (as confirmed by sequencing). The other five suppressors were linked to chromosome I and are likely alleles of mei-1 and/or mei-2, which mutate to dominant suppressors of rh24 at a high rate (Maine et al., 1990).

Online supplemental material

Video 1 corresponds to Fig. 1 A and Video 2 corresponds to Fig. 1 B. Fig. S1 shows anti-UNC-116 and anti-KLC-2 immunoblots of dsRNA-treated worms. Fig. S2 shows the spindle translocation defect in unc-116(rh245b79). Online supplemental material is available at http://www.icg.org/cgi/content/full/icb/200411132/DC1.

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