Spindle Dynamics during Meiosis in Drosophila Oocytes

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Abstract. Mature oocytes of Drosophila are arrested in metaphase of meiosis I. Upon activation by ovulation or fertilization, oocytes undergo a series of rapid changes that have not been directly visualized previously. We report here the use of the Nonclaret disjunctional (Ncd) microtubule motor protein fused to the green fluorescent protein (GFP) to monitor changes in the meiotic spindle of live oocytes after activation in vitro. Meiotic spindles of metaphase-arrested oocytes are relatively stable, however, meiotic spindles of in vitro–activated oocytes are highly dynamic: the spindles elongate, rotate around their long axis, and undergo an acute pivoting movement to reorient perpendicular to the oocyte surface. Many oocytes spontaneously complete the meiotic divisions, permitting visualization of progression from meiosis I to II. The movements of the spindle after oocyte activation provide new information about the dynamic changes in the spindle that occur upon re-entry into meiosis and completion of the meiotic divisions. Spindles in live oocytes mutant for a loss-of-function ncd allele fused to gfp were also imaged. The genesis of spindle defects in the live mutant oocytes provides new insights into the mechanism of Ncd function in the spindle during the meiotic divisions.

Meiosis is a specialized cell division that results in the formation of haploid gametes instead of diploid daughter cells, as in mitosis. Special features of meiosis that differ from mitosis include pairing of homologous chromosomes, recombination between homologues, and the occurrence of two successive divisions that reduce the chromosomes to the haploid number. Oocytes of most organisms are arrested at specific stages of meiosis that differ depending on the organism and can be activated by ovulation or fertilization to re-enter the cell cycle and complete the meiotic divisions.

Insights into the regulation of meiosis are anticipated to come from an understanding of the assembly, arrest, and reactivation of the meiotic spindle. Several structural differences exist between meiotic spindles of some oocytes and mitotic spindles of most animal cells. A striking difference is the absence of centrosomes at spindle poles of oocytes of several organisms that have been examined, including Drosophila (Sonnenblick, 1950), Xenopus (Gerhart, 1980), and the mouse (Szollosi et al., 1972). The absence of centrosomes raises the question of how microtubules are nucleated and organized for assembly of these spindles.

Thin-section electron microscopy has demonstrated that meiotic spindles of mouse oocytes lack centrosomes and centrioles, but electron-dense foci of microtubules are present at the broad poles of the oocyte spindles (Szollosi et al., 1972). A centrosomal protein, pericentrin, has been localized to multiple foci at the poles of the mouse oocyte spindle (Doxsey et al., 1994), consistent with the interpretation that the foci constitute dispersed microtubule organizing centers. Assembly of the meiotic spindle could occur from these dispersed centers in a manner similar to that of a single center. In contrast, known centrosomal components have not been identified in the Drosophila meiosis I spindle and are thought to be absent. The centrosomal protein, DMAP190 (CP190), was not found at the poles of the meiosis I spindle (Theurkauf and Hawley, 1992), but it has recently been reported to localize to a ring- or disk-shaped structure between the two tandem meiosis II spindles of Drosophila oocytes (Riparbelli and Callaini, 1996). The ring-shaped structure in the central pole region of the Drosophila meiosis II spindles was observed previously (Puro, 1991) and was suggested to function in pole organization for the second meiotic division. Nucleation and organization of microtubules for assembly of the Drosophila meiosis I and II spindles could therefore differ from one another and from anastral spindles of other organisms.

The discovery that several of the recently identified kinesin microtubule motor proteins are spindle-associated has led to the hypotheses that the kinesin motors function to generate the forces required for spindle assembly and maintenance of spindle bipolarity (for review see Walczak and Mitchison, 1996). Drosophila Nonclaret disjunctional (Ncd1, Yamamoto et al., 1989; Endow et al., 1990; McDonald and

1. Abbreviations used in this paper: blo, bloated; ca11, claret nondisjunctional; GFP, green fluorescent protein; Ncd, Nonclaret disjunctional; so, sine oculis; y, yellow.
Goldstein, 1990) is a minus-end directed kinesin motor protein (McDonald et al., 1990; Walker et al., 1990) that, when mutant, causes the formation of abnormal meiotic spindles in oocytes (Wald, 1936; Kimble and Church, 1983; Hatsumi and Endow, 1992b; Matthies et al., 1996). The wild-type Ncd motor protein has been proposed to function in establishing poles for assembly of bipolar meiosis I spindles (Kimble and Church, 1983; Hatsumi and Endow, 1992a,b; Matthies et al., 1996), replacing centrosomes in the spindle. A role in establishing bipolarity during assembly of the meiosis I spindle is consistent with the effects of null mutants on chromosome segregation (Sturtevant, 1929; Lewis and Gencarella, 1952), the abnormal spindles that have been observed in oocytes of the claret nondisjunctional (ca^{nd}) null mutant, and the association of the Ncd motor with meiotic spindle fibers (Hatsumi and Endow, 1992a; Matthies et al., 1996).

Assembly of the meiosis I spindle in wild-type oocytes and oocytes of the ca^{nd} null mutant has been examined by injection of rhodamine-conjugated tubulin to visualize spindle microtubules (Matthies et al., 1996). The results indicate that Ncd is required for normal kinetics of spindle assembly, as well as stabilization of the meiosis I spindle after assembly. Spindles were followed in this study from nuclear envelope breakdown through meiosis I spindle assembly and arrest of the meiosis I spindle.

Despite the emerging information regarding anastral spindle assembly in oocytes of Drosophila, the events of meiosis that occur after release from arrest and during progression through the meiotic divisions have not been adequately described. The classical account by Sonnenblick (1950) begins well into oocyte activation, with the spindle perpendicular to the oocyte surface. Spindles in mature arrested oocytes, however, are positioned parallel to the cortex (Theurkauf and Hawley, 1992; White-Cooper et al., 1993; Matthies et al., 1996). This raises the questions of how the spindle becomes oriented vertical to the cortex and when this occurs. There also exists a gap in our knowledge of spindle dynamics during re-entry into the meiotic cell cycle and completion of the meiotic divisions, which have not been directly visualized because of their rapid occurrence. The role of the Ncd motor during the meiotic divisions is also not known, although both the meiosis I and II spindles have been reported to be abnormal in ncd mutant oocytes (Wald, 1936; Hatsumi and Endow, 1992b).

To provide further information about meiotic spindle dynamics and Ncd function in the spindle, we fused the Ncd motor to the green fluorescent protein (GFP) of the jellyfish, Aequorea victoria (Prasher et al., 1992; Chalfie et al., 1994). An initial report of Ncd–GFP in mitosis in early embryos, including the rescue of an ncd null mutant by one of the first ncd–gfp gene fusions, has been published elsewhere (Endow and Komma, 1996). Here we show that Ncd–GFP is spindle-associated during meiosis in oocytes, like wild-type Ncd. The binding of Ncd–GFP to spindle microtubules provides a highly effective method for visualizing the meiotic spindle in live oocytes, resulting in new information regarding spindle dynamics during re-entry into meiosis and completion of the meiotic divisions. A mutant form of Ncd fused to GFP shows loss of function but binds to oocyte meiotic spindles, permitting spindle dynamics and the genesis of abnormal spindles in the presence of a loss-of-function Ncd motor to be observed. The results provide evidence that the Ncd microtubule motor is required for spindle elongation and, unexpectedly, stability of spindle fibers during the meiotic divisions. The Ncd motor probably also functions in polar body formation after the meiotic divisions, which are needed to prevent continued spindle-associated divisions of the maternal chromosomes after the initial two divisions.

### Materials and Methods

#### Drosophila Stocks

Drosophila mutants and balancer chromosomes used in this study are described in Lindsley and Zimm (1992). The classical ncd mutant allele (O’Toole and Szauter, 1980) is designated ncd by Lindsley and Zimm (1992). ncd^2 was obtained from J. Kennison (National Institutes of Health, Bethesda, MD) in 1985 and has been maintained in our stock collection. For the present studies, chromosome 3 proximal to ebony (e, 3-70.7) in the ncd^2 stock was replaced by recombination, and the X chromosome and chromosome 2 were replaced using balancer chromosomes to remove any modifiers that may have accumulated in the stock. The re-cloned e ncd^2 chromosome was used as a control in genetic tests of the ncd^2–gfp transgene.

#### Sequence Analysis of ncd^2

The ncd^2 mutant allele was sequenced using as template DNA fragments amplified by PCR from genomic DNA of ncd^2 adult Drosophila, and cloned into pBlueScript KS(+) (Stratagene, La Jolla, CA). The overlapping DNA fragments spanned the length of the ncd^2 coding region, starting in intron of the first in-frame AUG and continuing past the UAA stop codon (Endow et al., 1990) and included the two introns present in the ncd coding region. Changes in the DNA sequence that resulted in amino acid changes relative to wild-type Ncd were confirmed by sequence analysis of both DNA strands of two independent PCR clones.

#### Construction of ncd–gfp Plasmids

An ncd–gfp gene fusion was constructed in the P element vector, pCaSpeR3 (Pirrotta, 1988; Thummel et al., 1988), by insertion into BglIII plus EcoRI-digested plasmid of a 3.1-kb BamHI–EcoRI fragment containing the promoter and 5' region of ncd (Yamamoto et al., 1989), and a 2.1-kb EcoRI cDNA fragment encoding the remainder of Ncd (Endow et al., 1990). The XbaI site in the plasmid at the 5' end of the insertion was removed by digestion with NotI and StuI, followed by repair and religation. The plasmid was then digested with BamHI and XbaI, and the 3' end of the ncd gene in the plasmid was replaced with a BamHI–AIII ncd fragment and an AIII–XbaI gfp fragment. The ncd fragment was synthesized by PCR using an ncd cDNA plasmid as template and the primers, 5' ACA ATG GAC GGA GTG 3' and 5' TCA TCT TAA GGA AG 3', followed by digestion with BamHI and AIII, and gel purification. The gfp fragment was synthesized by PCR using a gfp-csa plasmid (Wang and Hazellrigg, 1994) as template and the primers, 5' GTG CTT AAG ATG AGT AAA 3' and 5' CTT CTA GAA TTC TTT GTA TAG TTC 3', followed by digestion with BamHI and XbaI, and gel purification. The ncd PCR insertion in the recombinant plasmid and all but the last 50 bp of the gfp insertion were sequenced to confirm the absence of PCR synthesis errors. The plasmid, pCaSpeR/ncd–gfp, consists of a wild-type ncd cDNA fused in-frame to wild-type gfp and encodes residues 1–700 of Ncd (Endow et al., 1990) followed by residues 1–238 of GFP (Prasher et al., 1992), with a change of D_{500} → L in Ncd. Transcription of ncd–gfp is under the regulation of the ncd promoter.

A plasmid encoding wild-type Ncd fused to the S_{65} → T mutant GFP (Heim et al., 1995), denoted GFP*, was constructed by replacing the AIII-XbaI gfp fragment in pCaSpeR/ncd–gfp with an AIII-XbaI PCR fragment encoding the mutant gfp. The S_{65} → T mutation in the pCaSpeR/ncd–gfp* plasmid was confirmed by DNA sequence analysis.

A plasmid containing ncd with the G_{65} → R mutation of the classical ncd^2 mutant allele fused to the S_{65} → T mutant gfp, ncd^2–gfp*, was constructed by replacing the Sphi–BamHI fragment of pCaSpe/R/ncd–gfp*
with an Sph–BamHI fragment encoding the G\text{ua} \rightarrow R mutation. The Sph–BamHI fragment of the new plasmid, \textit{pCasSpcRned–gfp}, was sequenced to confirm the presence of the G\text{ua} \rightarrow R mutation and the absence of PCR synthesis mutations.

**Germine Transformation and Genetic Tests for Null Mutant Rescue**

For P element–mediated germline transformation, the \textit{pCasSpcRned–gfp}, \textit{pCasSpcRned–gfp*}, and \textit{pCasSpcRned–gfp+} plasmids were injected into embryos of \textit{w}\textsuperscript{1185} females crossed to \textit{y w} \textit{Y} \textit{Sb} \textit{Tm6} Ub\text{x} males. Transplanting lines were established and made homozygous in \textit{ca}\textsuperscript{n} mutant females for cytological analysis of live or fixed oocytes and embryos. Females from the lines were tested genetically for rescue of \textit{ca}\textsuperscript{nl}.

Tests of transgenes for rescue of \textit{ca}\textsuperscript{nl} and tests of \textit{ncd}\textsuperscript{mut} effects on embryo viability and \textit{X} chromosome segregation were carried out as described previously (Komma et al., 1991; Endow et al., 1994). Parental females were mated to tester \textit{+}/\textit{Y} or \textit{y} \textit{w}\textsuperscript{1185}/\textit{Y} males. Regular offspring of these matings are + females and \textit{B} males. Meiotic nondisjunction of the \textit{X} chromosome in oocytes results in \textit{B}\textsuperscript{+}(\textit{X}/\textit{X} or \textit{Y}) females and \textit{X}/\textit{0} males; meiotic loss of the \textit{X} in early embryos gives rise to gynandromorphs (\textit{XX}/\textit{X0} mosaics). Minute offspring, haploid for \textit{X} chromosome, were scored but are excluded from calculations of total chromosome missegregation because of their highly variable recovery (Lindsley and Zimm, 1992). Calculations of meiotic nondisjunction and chromosome loss were performed as described (Komma et al., 1991; Endow et al., 1994). Statistical tests of significance were carried out using standard \textit{\chi}\textsuperscript{2} and Poisson distribution tests, assuming the null hypothesis that the offspring being compared were from the same population (Komma et al., 1991).

**Localization of Transgene Insertions**

The cytological sites of transgene insertions were determined by in situ hybridization using a biotin-11-DUTP–labeled \textit{cd} \textit{cDNA} probe, as described (Yamamoto et al., 1989). The \textit{F24M1} and \textit{F24M3} \textit{ncd–gfp} insertions were localized to 44B.C on \textit{2RA} and 2A1.2 on the \textit{X} chromosome, respectively. Recombination analysis of \textit{so} (\textit{so} \textit{257.1}, 43B1-2) and \textit{blo} (\textit{blo} \textit{2-855}, 44F1-2 to 45E1-2) with \textit{F24M1} and \textit{yellow} (\textit{y} \textit{1-0.0}, \textit{1B1}) and \textit{acute} (\textit{ac} \textit{1-0.0}, \textit{1B3}) with \textit{F24M3} was carried out to confirm the cytological map positions. The \textit{ncd–gfp*} transgene insertions, \textit{M3M1} and \textit{M9M1}, were localized by in situ hybridization to 42A.B at the base of \textit{2R} and 75C.D on \textit{3L}, respectively. The map positions were confirmed by recombination analysis of \textit{so} and \textit{blo} with \textit{M3M1} and \textit{blurry} (\textit{by} \textit{3-48.7}, \textit{8SD1–E3}) with \textit{M9M1}.

**Visualization of GFP in Live Oocytes**

Nonactivated live mature stage 14 oocytes were dissected from ovaries of \textit{ncd}\textsuperscript{2}–\textit{gfp} and \textit{ncd}\textsuperscript{2}–\textit{gfp*} females and transferred to a drop of light halocarbon oil on a glass slide. The chorion was partially removed under oil by pulling on the dorsal appendages with fine-tipped watchmaker’s forceps, and a coverslip fragment was mounted over the oocytes, as described above for nonactivated oocytes. In some cases the halocarbon oil was bubbled briefly with \textit{O}2 before use. The meiotic spindles of activated live oocytes could be effectively visualized without removal of the chorion, although sharper images were usually obtained by removing the chorion over the spindle. Laser scanning confocal microscopy was carried out using the 488-nm line of the laser, the \textit{GR2} or custom GFP filter block, and a neutral density filter that transmitted 1 or 3% of the laser light. Images were collected into stacks at 16.5 or 23 s intervals with 3 or 5 Kalman-averaged slow scans for each image.

**Analysis of Time Lapse Images**

Stack files of time lapse images were opened, converted to single images, and saved as PICT or TIFF files using the public domain program, NIH Image v 1.59. Image contrast and size was adjusted as necessary using Adobe Photoshop v 3.0.4. Measurements of spindle length were carried out with NIH Image calibrated with a 10 \textmu m bar superimposed on the image using COMOS (BioRad). The montages shown in Figs. 6 and 7 were made using the Crop and Make Montage stack macros of NIH Image. Image stacks were animated using NIH Image and recorded to videotape using NIH Image or Adobe Premiere v 4.0. Methods for making videotape recordings and QuickTime movies from GFP time lapse images are described in detail elsewhere (Endow and Piston, 1997). Video sequences of the figures can be viewed at http://abacus.mc.duke.edu

**Indirect Immunofluorescence Labeling of Fixed Oocytes and Embryos**

Nonactivated fixed oocytes were prepared by dissecting ovaries from wild-type (Oregon \textit{R}), \textit{ncd–gfp*}, or \textit{ncd–gfp*} females, partially submerged in modified Robb’s medium (Theurkauf and Hawley, 1992) containing 4.7% (vol/vol) formaldehyde. Mature oocytes were teased from the ovaries, fixed for a total of 1.5 min at room temperature in formaldehyde/saline solution, and then transferred into PBS. The chorion and vitelline membranes were removed from the fixed oocytes manually using fine-tipped watchmaker’s forceps. Fixed oocytes were then incubated overnight at room temperature in TBST (PBS plus 0.3% Triton X-100; Williams et al., 1992) before blocking in TBST plus 10% fetal calf serum for 1–2 h. Oocytes were incubated in rhodamine-conjugated anti-\textit{a}-tubulin monoclonal antibody (a gift of W. Sullivan; University of California, Santa Cruz, CA), as described (Endow et al., 1994). 5 \mu g/ml of DAPI (Boehringer Mannheim Corp., Indianapolis, IN) was added to one of the washes after antibody incubation to stain the chromosomes.

Normally fixed eggs at various stages of the meiotic divisions were obtained by collecting embryos at 12–15-min intervals from wild-type females or \textit{ncd}\textsuperscript{2–gfp*} mutant females, followed by dechorionation, removal of the vitelline membrane, and fixation in MEOH/EGTA without taxol, as described previously (Hatsumi and Endow, 1992b). Some collections of wild-type or \textit{ncd–gfp*} eggs were dechorionated manually before removal of the vitelline membrane rather than by treatment with Clorex, to save time. Rehydration of embryos, antibody staining, and subsequent washes were performed as previously described (Hatsumi and Endow, 1992b), except that TBST (PBS plus 0.1% Triton X-100) was used throughout instead of PBST (PBS plus 0.05% Triton X-100).

Antibody-stained oocytes and embryos were mounted in anti-fade solution consisting of 90% glycerol plus 0.1 vol of 10 mg/ml p-phenylenediamine in PBS, pH 9 (Stearns et al., 1991) for visualization of fluorescence.

**Laser Scanning Confocal and Fluorescence Microscopy**

Laser scanning confocal microscopy of fixed oocytes and embryos stained with rhodamine-conjugated anti-\textit{a}-tubulin antibody was carried out using a scanning confocal detector (MRC 600; BioRad) equipped with a krypton/argon laser and attached to a microscope (Axioptoph; Zeiss, Inc.). A 63X/1.4 NA Planapochromat objective and COMOS software (BioRad) were used to collect images of rhodamine-labeled spindles. Images of DAPI-stained chromosomes were collected using a cooled CCD camera (Pen-
Table I. Rescue of the cand Null Mutant by the ncd–gfp Transgene

| Female parent                  | Copies of ncd | Total gametes | X nd | X loss | Zygo X loss | Total* adults | Total embryos | Total missegregation | Embryo viability |
|--------------------------------|---------------|---------------|------|--------|-------------|---------------|---------------|---------------------|-----------------|
| 1. cd<sup>ad</sup>/cd<sup>ad</sup> | 0             | 143           | 0.238| <0.007 | 0.056       | 144           | 1,277         | 0.294               | 0.113           |
| 2. F24M1/F24M1: cd<sup>ad</sup>/cd<sup>ad</sup> | 2             | 1,515         | <0.001| 0.003  | 0.001       | 1,513         | 2,454         | 0.004               | 0.617           |
| 3. F24M3/F24M3: cd<sup>ad</sup>/cd<sup>ad</sup> | 2             | 664           | 0.012| 0.021  | 0.011       | 670           | 1,217         | 0.044               | 0.551           |
| 4. F24M3/F24M3: F24M1/F24M1:cd<sup>ad</sup>/cd<sup>ad</sup> | 4             | 1,767         | 0.002| <0.001 | <0.001     | 1,765         | 1,891         | 0.002               | 0.933           |
| 5. +/+                          | 2             | 1,883         | <0.001| <0.001 | <0.001     | 1,883         | 1,995         | <0.001              | 0.944           |

Females of the indicated genotypes were tested for embryo viability and segregation of the X chromosome. Females carrying the ncd–gfp F24M1 or F24M3 transgene were y<sup>1</sup> w<sup>1118</sup> scute<sup>3</sup> M9F1. Tests of function showed almost complete rescue of cd<sup>ad</sup> for meiotic and mitotic chromosome segregation by two copies of the ncd–gfp F24M1 transgene, resulting in a frequency of 0.004 X chromosome missegregation compared to 0.294 for cd<sup>ad</sup> and <0.001 for wild type (Table I). Embryo viability (0.617) was partially rescued compared to 0.113 for cd<sup>ad</sup> and 0.944 for wild type. Two copies of the F24M3 transgene partially rescued cd<sup>ad</sup> both for chromosome segregation and embryo viability (Table I). The incomplete rescue of cd<sup>ad</sup> by F24M1 and F24M3 indicates a position effect on ncd–gfp expression or interference by GFP with Ncd function. Drosophila with four copies of the ncd–gfp transgene (two copies each of F24M1 and F24M3) were also tested for rescue of cd<sup>ad</sup>. These females showed no significant differences from wild type with respect to chromosome segregation (0.002 missegregation compared with <0.001 for wild type; P = 0.06) or embryo viability (0.933 compared with 0.944 for wild type; χ<sup>2</sup> = 1.79, 1 degree of freedom, 0.5 > P > 0.1; Table I). The results demonstrate that the ncd–gfp F24M1 and F24M3 insertions together can rescue cd<sup>ad</sup> and replace the function of the wild-type Ncd microtubule motor protein.

Association of Ncd–GFP with Oocyte Meiotic Spindles

Initial observations using a cooled CCD camera or laser scanning confocal microscopy to visualize GFP fluorescence in live ncd–gfp oocytes revealed green fluorescent spindles, demonstrating that the ncd–gfp transgene is expressed in meiotic spindles.

Females of the indicated genotypes were tested for embryo viability and segregation of the X chromosome. Females carrying the ncd–gfp F24M1 or F24M3 transgene were y<sup>1</sup> w<sup>1118</sup> scute<sup>3</sup> M9F1. Tests of function showed almost complete rescue of cd<sup>ad</sup> for meiotic and mitotic chromosome segregation by two copies of the ncd–gfp F24M1 transgene, resulting in a frequency of 0.004 X chromosome missegregation compared to 0.294 for cd<sup>ad</sup> and <0.001 for wild type (Table I). Embryo viability (0.617) was partially rescued compared to 0.113 for cd<sup>ad</sup> and 0.944 for wild type. Two copies of the F24M3 transgene partially rescued cd<sup>ad</sup> both for chromosome segregation and embryo viability (Table I). The incomplete rescue of cd<sup>ad</sup> by F24M1 and F24M3 indicates a position effect on ncd–gfp expression or interference by GFP with Ncd function. Drosophila with four copies of the ncd–gfp transgene (two copies each of F24M1 and F24M3) were also tested for rescue of cd<sup>ad</sup>. These females showed no significant differences from wild type with respect to chromosome segregation (0.002 missegregation compared with <0.001 for wild type; P = 0.06) or embryo viability (0.933 compared with 0.944 for wild type; χ<sup>2</sup> = 1.79, 1 degree of freedom, 0.5 > P > 0.1; Table I). The results demonstrate that the ncd–gfp F24M1 and F24M3 insertions together can rescue cd<sup>ad</sup> and replace the function of the wild-type Ncd microtubule motor protein.

Drosophila carrying the ncd–gfp<sup>+</sup> transgene, with wild-type gfp replaced by the S<sup>65</sup> T mutant gfp (Heim et al., 1995), were also recovered. The ncd–gfp<sup>+</sup> transgenes, M3M1 and M9F1, were localized by in situ hybridization to 42A,B at the base of 2R and 75C,D on 3L, respectively. The cytological map positions were confirmed by recombination analysis of so and blo with M3M1 and blistery with M9F1. Tests of homozygous M3M1 or M9F1 showed partial rescue of cd<sup>ad</sup>, while four copies of ncd–gfp<sup>+</sup>, two each of M3M1 and M9F1, showed essentially complete rescue of cd<sup>ad</sup> both for chromosome segregation and embryo viability (Endow and Komma, 1996).
pressed and can be detected in live oocytes. The spindle fluorescence indicated that the Ncd–GFP fusion protein is associated with the oocyte meiotic spindle, as observed previously for wild-type Ncd using antibody staining (Hatsumi and Endow, 1992a; Matthis et al., 1996). Association of Ncd–GFP with meiotic spindles permits analysis of meiotic spindle dynamics in live oocytes of Drosophila.

Live ncd–gfp oocytes were prepared for observation by dissection from ovaries either under halocarbon oil or into Drosophila PBS (Robb, 1969), followed by mounting under oil. Oocytes dissected under oil were interpreted to be nonactivated, by comparison with nonactivated fixed oocytes prepared by dissecting ovaries directly into fixative. Oocytes dissected into Drosophila PBS showed dramatic changes in the spindle, including meiosis I spindle elongation and assembly of tandem meiosis II spindles. These oocytes are interpreted to be activated since the cytological changes in the spindles are consistent with oocyte activation, as evidenced by analysis of normally activated fixed eggs. These results are described below.

**Meiotic Spindle Dynamics in Nonactivated Live Oocytes**

Spindles of nonactivated live ncd–gfp oocytes, prepared by dissection from ovaries of females under halocarbon oil, could be observed through the chorion by laser scanning confocal microscopy. The tapered bipolar spindles, located near the base of the dorsal appendages, were brightly fluorescent with Ncd–GFP (Fig. 1). The dark region in the center of the fluorescent spindles was presumed to correspond to the chromosomes, which excluded Ncd–GFP. This was confirmed by antibody staining of nonactivated fixed oocytes. Spindles of nonactivated live oocytes imaged without removal of the chorion were relatively stable, remaining in the same position throughout the 15–30 min periods of image collection with little or no net change in length or orientation. The spindle shown in Fig. 1 was from a ca	extsuperscript{nd} oocyte with four copies of the ncd–gfp	extsuperscript{*} transgene. Stable metaphase I meiotic spindles were also observed in nonactivated live oocytes with one copy of the ncd–gfp	extsuperscript{*} transgene or four copies of the ncd–gfp transgene.

Despite their relatively stable positions, dynamic changes in the spindles of nonactivated live oocytes were occurring that could be detected over time. These changes comprised small lengthwise extensions and contractions of the spindles and alternating clockwise and counter clockwise rotational movements around their long axis, as diagrammed in Table II. These slight movements of the spindle were observed in all of the nonactivated live oocytes that were imaged (n = 13) and were best analyzed by animating the time lapse images and determining differences in orientation and position of the spindles over time. The spindles of most nonactivated live oocytes showed both extensions/contractions and rotational movements, but no net change in length or position over time (Table II). Spindles of several oocytes showed more extensive movements: one spindle rotated two to three revolutions around its long axis during the first 4 min of image collection before assuming a more stable position, and one spindle rotated approximately a quarter turn over the 16.5-min observation time. Another spindle moved lengthwise a quarter of its length and then returned to its original position, and one spindle elongated ~75% of its length. The oocyte containing this last spindle may have been unintentionally activated by the continuous laser irradiation. With the exception of this oocyte, none of the nonactivated live oocytes with intact chorions showed changes in the spindle that resembled completion of meiosis I and progression into meiosis II. These oocytes were therefore interpreted as arrested in meiosis I.

The spindles of nonactivated live oocytes were typically observed through the chorion. Removal of the chorion from the live oocytes resulted in changes that included elongation of the spindle followed by the formation of two tandem meiosis II spindles. These changes are consistent with oocyte activation and occurred even though the oocytes were dissected under oil and the chorion was removed under oil. These changes were observed in 3/3 oocytes that were prepared in this manner and observed. Perturbation of nonhydrated Drosophila oocytes can therefore cause oocyte activation, resulting in re-entry into meiosis and completion of the meiotic divisions.

**Table II. Spindle Movement in Nonactivated Live Oocytes**

| No net change over time | Alternating CW and CCW rotations | Lengthwise movement | Elongation | Rotation |
|-------------------------|----------------------------------|---------------------|------------|----------|
|                         | Extension/contraction             |                     |            |          |
| 12                      | 7                                | 1                   | 1          | (2–3 revolutions) |
|                         |                                   |                     | 1          | (1/4 turn) |

* Barely detectable rotational movement.

CW, clockwise; CCW, counter clockwise.
Chromosomes associated with the meiotic spindles could not be visualized in nonactivated live ncd–gfp oocytes in the absence of further perturbing treatments, such as injection of fluorescent DNA dyes, that could cause oocyte activation. Nonactivated fixed ncd–gfp and wild-type oocytes were therefore prepared and stained with anti–α-tubulin antibody and DAPI for comparison with spindle images from nonactivated live ncd–gfp oocytes and determination of the chromosome configurations. Metaphase I–arrested spindles in nonactivated fixed wild-type oocytes, merged with DAPI-stained chromosomes, are shown in Fig. 2. The fluorescent spindles visualized by tubulin antibody staining are similar in appearance to the Ncd–GFP fluorescent spindles. The highly condensed metaphase I chromosomes are present in the center of the spindle, in a position corresponding to the dark regions in the spindles visualized with Ncd–GFP. The spindles stained with anti-tubulin antibody also showed a dark region in the center corresponding to the chromosomes. The small dotlike chromosome 4 could be observed either associated with the remainder of the chromosomes and identified as a small protrusion at the ends of the chromosome mass (Fig. 2 A, arrows), or separated from the rest of the chromosomes and positioned closer to the poles (Fig. 2 B, arrows). The spindle and chromosome configurations of nonactivated fixed ncd–gfp oocytes, prepared using the same methods as nonactivated fixed wild-type oocytes and stained with anti-tubulin antibody and DAPI, were similar in appearance to the spindles and chromosomes shown in Fig. 2.

**Rotation of the Meiotic Spindle in Activated Live Oocytes**

Oocytes were also prepared by dissection from ovaries of ncd–gfp or ncd–gfp* females into Drosophila PBS (Robb, 1969). Mature oocytes, visibly swollen by the brief (<3 min) immersion in the undiluted saline solution, were transferred to a drop of halocarbon oil on a slide, the chorion was removed from the anterior of the oocyte, and a coverslip was mounted over the oocytes. Time lapse images collected for periods of 15–50 min by laser scanning confocal microscopy showed that the meiotic spindles of live oocytes prepared by immersion in Drosophila PBS exist in a highly dynamic state. As an example, the spindle shown in Fig. 3 was first observed parallel to the oocyte surface (Fig. 3 A). The spindle elongated (Fig. 3 B), contracted, and then underwent an acute pivoting movement (Fig. 3 C) that reoriented the spindle from its initial position parallel to the oocyte surface into a position perpendicular to the oocyte cortex (Fig. 3 D). Fig. 3 D shows an image looking down the long axis of the spindle formed by the spindle poles, with the dark “holes” corresponding to the chromosomes, based on fixed oocytes and embryos stained for tubulin and DNA. The spindle remained in this position for at least 9.2 min, rotating around its long axis.

Rotations of the spindle around its long axis, after pivoting vertical to the cortex, were either clockwise or counter clockwise, or rapidly alternating clockwise and counter clockwise rotations. Spindles of 11 oocytes were analyzed after pivoting. Six spindles showed only rapidly alternating rotations, three rotated clockwise accompanied by rapidly alternating rotations, one only rotated clockwise, and two rotated counter clockwise and also showed rapidly alternating rotations.

Spindles of oocytes prepared by immersion in Drosophila PBS could rotate around their long axis before or after pivoting vertical to the oocyte surface, and several spindles elongated or contracted, as shown in Fig. 3. Microtubules, either single fibers or bundles, could be seen in images in which the spindles were cross-sectioned, projecting out from the spindles. Comparison of successive images in the time lapse sequences gave the impression that rapid shortening and elongation of the spindle microtubules was occurring and that the spindle was in a highly dynamic state.

**Meiosis I to II Progression in Activated Live Oocytes**

Spindles of oocytes that were prepared by immersion in Drosophila PBS could spontaneously complete the meiosis I and II divisions, permitting, for the first time, visualization of spindle dynamics during the transition from meiosis I to II. The meiosis I spindle shown in Fig. 4 A was

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**Figure 2.** Metaphase I spindles in nonactivated fixed wild-type oocytes. Fixed oocytes were stained with anti–α-tubulin antibody (red) and DAPI (green) to visualize meiotic spindles and chromosomes. The position of the chromosomes in the spindle corresponds to the dark hollow in the center of the spindle in the live ncd–gfp* oocyte (Fig. 1). The dot-like fourth chromosomes (arrows) were found either (A) associated with the condensed mass of chromosomes or (B) positioned closer to the poles in the metaphase I spindles. Bar, 5 μm.
positioned obliquely to the oocyte surface and was viewed from an angle that allowed imaging of the entire spindle. The spindle extended into an elongated, tapered meiosis I spindle (Fig. 4 B), measuring ~30 μm from pole to pole, and then rapidly reassembled into two tandemly arrayed meiosis II spindles (Fig. 4, C and D). Assembly of the tandem meiosis II spindles occurred by the formation of a bright focus of Ncd–GFP in the center of the elongated meiosis I spindle (Fig. 4 B, arrow), followed by the establishment of two central spindle poles. The tandem meiosis II spindles were interpreted to assemble by rapid sliding of the spindle microtubules against one another, based on the movements of the spindle observed in the time lapse sequence. The transition into meiosis II occurred without disassembly of the meiosis I spindle and involved reorganization of the spindle fibers into the two tandem meiosis II spindles.

The time required to complete the meiotic divisions was determined by analysis of oocytes for which a complete sequence of meiosis I and II had been recorded. Meiosis I and II in time lapse sequences of three oocytes with four copies of the ncd–gfp transgene each lasted ~2.5–5 min, giving an estimate of ~5–10 min for completion of both divisions. In a previous study by other workers, eggs were collected for 5 min and fixed after an additional 5 min for a total development time of 5–10 min, and then stained and analyzed (Riparbelli and Callaini, 1996). 37% of the eggs were in meiosis I, 51% were in meiosis II, and 12% were in mitosis. This distribution is consistent with our estimate of ~5–10 min for completion of both meiotic divisions.

To determine if the spindles observed in activated live ncd–gfp oocytes resembled those of normally activated eggs, wild-type embryos collected at 0 to 12–15-min intervals were fixed and stained with anti-tubulin antibody and DAPI. The meiosis II spindles of normally activated fixed eggs were typically perpendicular to the cortex and aligned in tandem (Fig. 5). A faint ring-like central spindle pole body consisting of foci of tubulin with radiating microtubules could be observed between the two meiosis II spindles (Fig. 5 A, arrow). Other spindles showed only diffuse tubulin staining in the region between the two meiosis II spindles (Fig. 5 B). The central pole body of meiosis II spindles of normally activated fixed eggs was also stained by anti-Ncd antibody (not shown).

A Loss-of-function ncd Mutant and ncd–gfp Transgene

ncd2 is a classical mutant of ncd that was originally isolated after EMS mutagenesis. The mutant was shown to fail to complement cand for chromosome missegregation (O’Tousa and Szauter, 1980), but a large-scale genetic analysis of the effects of ncd2 has not been reported previously. Tests of ncd2 mutant females showed markedly reduced embryo vi-

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**Figure 3.** Meiotic spindle dynamics in an activated live ncd–gfp oocyte. Images from a time lapse series showing, from top to bottom, the spindle initially positioned parallel to the surface of the oocyte (A) and elongation of the spindle (B), followed by contraction and pivoting (C) into a vertical position with respect to the oocyte surface (D). The spindle remained in this position at least 9.2 min after pivoting, rotating around its long axis. Time in minutes and seconds is shown on each image. Bar, 5 μm.
ability and elevated chromosome nondisjunction and loss compared to wild type (Table III). ncd² shows a slightly more severe effect on embryo viability (0.098) than ca² (0.113), but there is no significant difference in total X chromosome missegregation between ncd² and ca² (χ² = 0.371, 1 degree of freedom, 0.5 < P < 0.9). Chromosome missegregation is fully rescued in heterozygous ncd²/+ females, but embryo viability (0.810 compared to 0.944 for wild type) is only partially rescued (Table III). ncd² therefore behaves like a loss-of-function mutant of ncd with respect to chromosome segregation and shows a small semidominant effect on embryo viability. The mutant ncd² gene is transcribed (Yamamoto et al., 1989), and the mutant protein is spindle-associated (Hatsumi and Endow, 1992a). The cytological effects of ncd² on meiotic spindles were found previously to be the same as those of ca² (Hatsumi and Endow, 1992b).

Sequence analysis of the coding region of ncd² revealed four amino acid changes, G446 → R, A566 → S, G696 → A, and N697 → S, compared to wild-type Ncd. The G446 → R missense mutation affects a glycine residue in the ATP-binding motif of Ncd, IFAYGQTGSGLKYTMGDG, that is highly conserved among the kinesin proteins. This amino acid change is the likely cause of the loss of function of ncd². Of the other changes in ncd², A566 → S is a conservative amino acid change, G696 → A lies outside the motor domain, and N697 → S is a polymorphism found in some presumed wild-type strains. No other nucleotide substitutions were found in ncd² that result in missense mutations, or changes in the introns or intron/exon boundaries compared to ncd¹.

The G446 → R missense mutation, but not the other 3 amino acid changes in ncd², was introduced into an ncd–gfp* plasmid containing the S65 → T mutant gfp, and the plasmid was injected into embryos for germline transformation. Transformants carrying the ncd²–gfp* transgene were recovered, and the transgenes were made homozygous in ca² females. The ncd²–gfp* transgene that was analyzed in this study is designated M4F1. The ncd²–gfp* ca² females showed poor fertility, high egg inviability, and elevated chromosome missegregation that paralleled the effects of ncd² (Table III). The effects of the M4F1 ncd²–gfp* transgene on chromosome segregation were not significantly different from those of ncd² (χ² = 0.944, 1 d.f., 0.5 < P < 0.9), although M4F1 showed a slightly more severe effect on embryo viability (0.085) than ncd² (0.098). Females carrying one copy of M4F1 together with one ncd¹ allele (M4F1; ca²/¹ females) showed complete rescue for chromosome segregation, but embryo viability (0.886 compared with 0.944 for wild type) was only par-

Figure 4. Meiosis I to meiosis II progression in an activated live ncd–gfp oocyte. Time lapse images showing, from top to bottom, the meiosis I ncd–gfp oocyte spindle (A), elongation of the meiosis I spindle (B), reassembly into two tandem meiosis II spindles, and progression into meiosis II (C and D). Time in minutes and seconds is shown on each image. The arrow (B) indicates a bright focus of Ncd–GFP where the central spindle poles form. The spindles are positioned obliquely to the oocyte cortex, permitting the meiosis I and II spindles to be completely imaged. The interior of the oocyte is to the left. Bar, 10 μm.
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Table III. The ncd-2–gfp* Transgene Fails to Rescue the cd-Null Mutant

| Female parent | Copies of ncd-2 | Total gametes | Gametic | Zygotic | Total* | Total embryos | Total missegregation | Embryo viability |
|---------------|----------------|---------------|---------|---------|--------|---------------|---------------------|-----------------|
|               |                |               | X nd    | X loss  |        |               |                     |                 |
| 1. cd-0/cd-0  | 0              | 143           | 0.238   | <0.007 | 0.056  | 144           | 1,277               | 0.294           |
| 2. cd-0/+     | 0              | 1,569         | 0.003   | <0.001 | <0.001 | 1,567         | 1,748               | 0.003           |
| 3. ncd-2/ncd-2| 2              | 79            | 0.101   | 0.127  | 0.127  | 155           | 1,582               | 0.355           |
| 4. ncd-2/0    | 1              | 1,829         | <0.001  | <0.001 | <0.001 | 1,830         | 2,258               | <0.001          |
| 5. M4F1/M4F1; cd-0/cd-0 | 2 | 86  | <0.012 | 0.326  | 0.070  | 98            | 1,153               | 0.396           |
| 6. M4F1;cd-0/+ | 1            | 3,322         | <0.001  | 0.001  | <0.001 | 3,321         | 3,747               | 0.001           |
| 7. +/-        | 0              | 1,883         | <0.001  | <0.001 | <0.001 | 1,883         | 1,995               | <0.001          |

*Including M offspring.

Females carrying the ncd-2 mutant allele of ncd or the ncd-2–gfp* M4F1 transgene were tested for embryo viability and segregation of the X chromosome. Females carrying the ncd-2–gfp* M4F1 transgene were y w; M. Total missegregation is that of the X chromosome. The data shown for ncd-2/ncd-2 were obtained using a newly recloned ncd-2 allele. Data from Table I for cd-0 and + females are shown for comparison.

Figure 5. Meiosis II spindles in normally activated fixed wild-type eggs. Normally oviposited wild-type eggs were fixed and stained with anti-α-tubulin antibody (red) and DAPI (green) to visualize the spindles and chromosomes. The meiosis II spindles are perpendicular to the cortex and are viewed down their long axis. The bottom spindle in each image, which is closer to the surface of the embryo, is slightly delayed relative to the more internal spindle; this is probably an artifact caused by the time required for penetration of the fixative. (A) Meiosis II spindles in early (bottom) and mid- (top) anaphase. The faint central spindle pole body (arrow), which consists of foci of tubulin in a ring-like array with radiating microtubules, can be observed in the region between the two spindles. (B) Meiosis II spindles in mid- (bottom) and late (top) anaphase. The central spindle pole body appears as a diffuse array of microtubules between the two spindles. Bar, 10 μm.

Metaphase I spindles were detected in the ncd-2–gfp* M4F1 transgenic line, which contains an intact X chromosome (Fig. 5 A, bottom). The ncd-2–gfp* transgene therefore closely parallels the ncd-2 mutant allele in its effects on chromosome segregation and embryo viability.

Meiotic Divisions in Loss-of-Function Mutant Oocytes

Green fluorescent spindles were detected in nonactivated live ncd-2–gfp* oocytes prepared by dissection under halocarbon oil and observed by laser scanning confocal microscopy. Nonactivated fixed wild-type mature oocytes contained long or short, unconnected spindles, or partially formed abnormal spindles (not shown). Fully formed meiosis I spindles were not observed in the nonactivated fixed mutant oocytes (n = 104), even though mature oocytes were selected for the cytological analysis. The condensed meiotic chromosomes associated with the spindles also appeared to be at an earlier stage than chromosomes in nonactivated fixed wild-type mature oocytes. It is therefore probable that spindle assembly and maturation of the ncd-2–gfp* mutant oocytes is delayed relative to wild type.

Live ncd-2–gfp* oocytes, prepared by dissection into Drosophila PBS, contained green fluorescent spindles, provid-
ing evidence that the ncd2–gfp* transgene is expressed and the Ncd2–GFP* protein is associated with meiotic spindles. The basis of the ability to visualize spindles in activated live oocytes but not nonactivated live oocytes is not certain, but is likely to be due to changes in the oocyte that occur upon activation. The spindles in the live mutant oocytes were activated to complete the meiotic divisions, but the divisions were highly abnormal. In the time lapse series shown in Fig. 6, a spindle was initially observed that appeared to be formed of closely apposed spindles (A and B). The spindle was interpreted to be in meiosis I, based on the subsequent events in the time lapse sequence. Multiple poles arose by separation of the spindles at the poles (Fig. 6 C, arrows). The spindles failed to extend into the highly elongated meiosis I spindles typical of wild-type oocytes. A small focus of Ncd2–GFP* formed in part of the spindle (Fig. 6 C, arrowhead), marking the position of the meiosis II-like central spindle poles, but normal meiosis II spindles failed to form. Instead, a short multipolar spindle arose by movement of the newly formed central spindle poles away from the rest of the spindle, pulling on microtubules attached to the original poles (Fig. 6, D–H).

Although spindles in most of the activated live mutant oocytes failed to elongate into typical meiosis I spindles, spindles in 2 of the 23 mutant oocytes that were recorded did show elongation typical of wild-type meiosis I spindles. Elongation of spindles in these two oocytes was followed by the formation of two independent pairs of central spindle poles in the closely apposed spindles that comprised the spindle; movement of the new poles in opposite directions resulted in the formation of cruciform spindles.

Multiple spindles were present in some activated live mutant oocytes that arose by separation of meiotic spindles into several spindle-like components. As an example, the time lapse series in Fig. 7 shows three spindles that were separated from one another when initially observed. Movement of the spindles in different directions caused them to become somewhat more widely separated (Fig. 7, A–C). The spindle fibers in the centers of the spindles then began fragmenting (Fig. 7, D–H), releasing the poles from the spindles (G and H). Fluorescent particles, presumed to be microtubule fragments bound to Ncd2–GFP*, were observed in the cytoplasm and formed a hazy network in the center of the spindles (Fig. 7, E–H). The released poles of the spindles moved far from the original site of the spindles and could be observed in the oocyte after the time lapse imaging, separated from one another by as much as 75–80 μm.

Progression of the meiotic divisions was slow in the activated live mutant oocytes compared with wild-type oocytes, requiring 30 min or longer for completion of two divisions, and spindles in some mutant oocytes continued dividing after the initial two divisions without undergoing disassembly.

Spindles in normally activated fixed mutant eggs were examined for comparison with spindles in activated live mutant oocytes. Mutant eggs in various stages of the meiotic divisions were collected at 0–15 min intervals, fixed,
and stained with α-tubulin antibody and DAPI to visualize the meiotic spindles and chromosomes. Anaphase I and metaphase II spindles in normally activated fixed ncd2–gfp* mutant eggs are shown in Fig. 8. Several of the normally activated fixed mutant eggs exhibited a haze of microtubules associated with the meiotic spindles (Fig. 8 A), as observed in activated live mutant oocytes. Multiple spindles were present, both in meiosis I and II, with each of the separated chromosomes, including the small chromosome 4 (Fig. 8 A, arrows), associated with a spindle spur or spindle. The spindles were oriented obliquely or vertically with respect to the embryo cortex. The arrow in Fig. 8 B indicates a cross-sectioned spindle that is vertical, associated with a spindle spur, indicating that the mutant Ncd–GFP does not prevent reorientation of the spindles perpendicular to the cortex. One or more of the spindle-associated meiotic chromosomes in some eggs was widely separated from the others. For example, one of the 8 half-bivalent chromosomes was missing from the cluster of spindle-associated metaphase II chromosomes in one egg. The missing chromosome was found 85–90 μm from the nearest chromosome in the cluster, associated with a small spindle remnant.

Normally activated fixed early eggs of ncd2–gfp* mutant females frequently contained multiple spindle-associated chromosomes that exceeded the 4N = 16 number expected upon completion of the two meiotic divisions. The spindles lacked centrosomes, like the oocyte meiotic spindles, and, in many eggs, were abnormal (either multipolar, branched, or spurred). The fixed mutant egg shown in Fig. 8 C contained 18 chromosomes or pairs of nondisjoined sister chromatids. Many of the chromosomes were separated from one another and associated with separate spindles, and some of the chromosomes showed anaphase configurations, as if they were in the process of dividing. One of these is indicated by an arrow. The spindles were located near the cortex of the egg, where the polar bodies are found in wild-type embryos. The egg was unfertilized, and no mitotic spindles were observed. The >4N chromosomes in this and other eggs probably arose by continued meiotic divisions of the maternal chromosomes. This interpretation is consistent with the more than two meiotic divisions that were observed in activated live mutant oocytes.

Discussion

GFP as a Fluorescent Tag to Monitor Meiotic Spindle Dynamics

The meiotic divisions in oocytes of most organisms, including Drosophila, have not been directly visualized previously. Several steps after oocyte activation occur very rapidly and have not been observed even in rapid collections of eggs analyzed by fixation and antibody staining. These
steps include the expansion/contraction of the spindle, spindle rotations, and the acute pivoting movement that reorients the spindle into a position perpendicular to the cortex before completion of the meiotic divisions.

Figure 8. Meiotic divisions in normally activated fixed ncd2–gfp* mutant eggs. Normally oviposited mutant ncd2–gfp* eggs were fixed and stained with anti-α-tubulin antibody (red) and DAPI (green) to visualize the spindle microtubules and meiotic chromosomes. The cortex of the egg is nearer to the bottom of each of the images. (A) Late anaphase I half bivalents are segregating on separate spindles. The tiny fourth chromosomes (arrows) are associated with spindle spurs or small spindles. A hazy mass of tubulin-positive material is present in the central region of each spindle and is interpreted to be fragmented microtubules. The spindles at the top and bottom are bent or skewed, and the spindle in the middle is positioned obliquely with respect to the cortex. (B) Metaphase II chromosomes are associated with separate spindles or spindle spurs. The spindles are positioned obliquely or vertically (arrow) with respect to the cortex. The spindle indicated by the arrow is cross-sectioned and joined to a spindle spur with an associated chromosome. (C) Continued spindle-associated divisions of the maternal chromosomes after the initial two divisions. The egg contained 18 spindle-associated chromosomes or nondisjoined sister chromatids, more than the number of chromosomes expected after completion of the two meiotic divisions. Several of the chromosomes appear to be dividing. An anaphase chromosome configuration is indicated by the arrow. The spindles lack centrosomes and are present near the cortex, where the polar bodies are found in wild-type embryos. The egg was unfertilized and contained no mitotic spindles. Bars, 10 μm.

Fusion of ncd to gfp and expression of the gene fusion in Drosophila under the control of the wild-type ncd promoter targets the fluorescent motor–GFP fusion protein to the meiotic spindle, permitting visualization of meiotic
spindle dynamics in live oocytes. The Ncd–GFP fusion protein can replace the function of the wild-type Ncd motor protein, rescuing the ca16 null mutant both for chromosome segregation and embryo viability. Visualization of meiotic spindles using a microtubule-associated motor protein fused to GFP is a highly effective method for monitoring changes that occur in the oocyte spindle. The meiotic spindles can be visualized through the chorion of nonactivated or activated live oocytes, minimally perturbing the oocyte.

**Meiosis I and II Spindle Dynamics in Wild-Type Oocytes**

Spindles of nonactivated live oocytes were observed after dissection under oil, conditions that prevent oocyte hydration and consequent activation (Theurkauf, 1994). Most of the spindles were stable, showing only slight movements or changes in position over periods of 15–30 min observation. The spindles appeared to be undergoing dynamic changes, as evidenced by the detection of slight extensions/contractions in length and alternating clockwise and counter clockwise rotational movements. The spindles of nonactivated live oocytes were positioned parallel to the oocyte cortex and could be observed as early as stage 13 (King et al., 1956; Spradling, 1993) in oocytes that were still associated with degenerating nurse cells.

In contrast to the nonactivated live oocytes, live oocytes that were observed after brief immersion in *Drosophila* PBS contained spindles that were highly dynamic. The spindles extended and contracted, rotated around their long axis, and pivoted from their initial position parallel to the oocyte cortex into a vertical position, where they continued to rotate. We interpret the highly dynamic state of the meiotic spindle that we observe upon hydration of oocytes as representing the state of the spindle immediately after activation. The dynamic changes in the meiotic spindle after oocyte activation are shown schematically in the model in Fig. 9 and can be summarized as follows: (a) rotation of the spindle, (b) pivoting of the spindle vertical to the cortex, followed by rotations, (c) completion of meiosis I, and (d) assembly of meiosis II spindles and completion of meiosis II. Several of these changes can also be detected or observed in normally activated fixed eggs.

Rotation and pivoting of the *Drosophila* meiotic spindle have not been observed previously, although reorientation of the meiosis I spindle has been inferred to occur based on positional differences in spindles in metaphase-arrested and activated fixed oocytes (Riparbelli and Callaini, 1996). The rotations and acute pivoting movement of the spindle represent two of the initial changes that occur in the spindle after oocyte activation. The ability of the spindle to rotate and reorient before completing the meiotic divisions explains why the spindle in nonactivated live (Fig. 1) and fixed oocytes (Fig. 2; Theurkauf and Hawley, 1992; White-Cooper et al., 1993) can be observed parallel to the cortex, while the meiotic divisions occur with the spindle perpendicular to the cortex (Sonnenblick, 1950; Figs. 4 and 5). Our observations indicate that the meiotic spindle reorients soon after oocyte activation by an acute pivoting movement, accompanied by spindle rotations.

Upon reorientation perpendicular to the oocyte cortex, the spindles of in vitro–activated *Drosophila* oocytes frequently elongated and completed the meiosis I and then the meiosis II division. The transition into meiosis II after the completion of meiosis I, including assembly of the tandem meiosis II spindles, has not been observed previously, although the second meiotic division is known to occur with the two spindles tandemly aligned (Sonnenblick, 1950). The observations reported here indicate that assembly of the meiosis II spindles occurs by the formation of new spindle poles in the center of the anaphase I spindle, as anticipated by the work of others (Puro, 1991; Riparbelli and Callaini, 1996). The newly formed spindle poles serve as the central poles for the two meiosis II spindles. Assembly of the tandem meiosis II spindles appears to involve reorganization of the meiosis I spindle fibers, possibly by rapid sliding of the spindle microtubules, without disassembly of the meiosis I spindle.

The mechanism by which the central meiosis II spindle poles form is not known, although the ring- or disk-shaped central spindle pole body that lies between the two meiosis II spindles has been suggested to organize the poles (Puro, 1991). The recent observation that a centrosomal protein, CP190 (DMAP190), localizes to the central spindle pole body (Riparbelli and Callaini, 1996) suggests that centrosomal proteins are involved. In contrast, centrosomal proteins such as γ-tubulin, DMAP60, and DMAP190 have not been found associated with meiosis I spindles of *Drosophila* oocytes (Theurkauf and Hawley, 1992; Matthies et al., 1996). These observations raise the possibility that the mechanism by which spindle poles are formed for meiosis I and II spindle assembly in *Drosophila* oocytes differs, and that assembly of meiosis II spindles requires centrosomal proteins that are found in mitotically dividing cells.

**Rotation of Meiotic Spindles in Xenopus Oocytes**

In addition to the present work with *Drosophila*, meiotic spindle dynamics have been followed in activated live oocytes of *Xenopus* after injection of FITC-conjugated tubulin (Gard, 1992). The *Xenopus* meiosis I and II spindles as-

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**Figure 9. Model of spindle dynamics after oocyte activation.** Meiosis I spindles in *Drosophila* oocytes are assembled parallel to the cortex of the oocyte (represented by the line) and remain in this position during metaphase arrest. After normal activation by ovulation, the oocyte swells. The spindle rotates around its long axis and pivots into a vertical position with respect to the cortex, followed by further rotations. Completion of meiosis I occurs with the spindle perpendicular to the cortex and is immediately followed by reorganization of the meiosis I spindle into two tandem meiosis II spindles and progression into meiosis II.
sembled with the long axis parallel to the oocyte surface and then rotated into a vertical position before undergoing division. Reorientation of the meiotic spindle in *Xenopus* oocytes was inhibited by treatment of oocytes with cytochalasin B during maturation (Gard et al., 1995), indicating the requirement for an intact actin cytoskeleton. Based on our present observations, meiosis I spindles of *Drosophila* and *Xenopus* oocytes undergo an analogous acute pivoting rotation before division. Division with the spindle perpendicular to the cortex positions one of the daughter nuclei more internally and the other closer to the surface. After completion of the second meiotic division, the innermost nucleus in *Drosophila* or the more internal nucleus in *Xenopus* becomes the female pronucleus, and the nuclei or nucleus near the cortex undergo polar body formation. The basis for the initial assembly of spindles parallel to the cortex is not certain, but may have to do with the anchoring or attachment of the spindle to the cortical actin cytoskeleton.

Despite their overall similarity, some aspects of spindle dynamics in *Xenopus* and *Drosophila* oocytes differ. For example, *Xenopus* meiotic spindles were not reported to rotate around their long axis before the acute pivoting rotation that reorients the spindle perpendicular to the cortex. The rotations of the *Drosophila* oocyte spindles may cause the pivoting and, after reorientation, lead to spindle elongation and completion of the meiotic divisions. The spindle pivots in *Xenopus* and *Drosophila* could therefore be similar in effect, but differ in basis.

**Oocyte Activation in Drosophila**

The initial report of in vitro activation of *Drosophila* oocytes showed that hydration in hypotonic (diluted) Robb’s medium (Robb, 1969) induces physiological changes in oocytes that are consistent with normal activation by ovulation (Mahowald et al., 1983). Tests of undiluted Robb’s medium did not result in the ultrastructural changes typical of normally ovulated eggs that were also observed in oocytes treated with hypotonic medium. These results have led to the belief that hypotonic solution is required to cause activation, since 3/3 oocytes that were dissected under oil completed the meiotic divisions after their chorions were partially removed. Hydration of oocytes is therefore not required for activation. Activation of *ncd–gfp* oocytes using methods recently developed for efficient mass activation of *Drosophila* oocytes (Page and Orr-Weaver, 1997) may prove of value in future studies.

**Effects of Loss of Ncd Function on Oocyte Meiotic Divisions**

The classical mutant allele, *ncd*2, shows loss of function based on its failure to rescue the *cad* null mutant. The basis of the loss of function of *ncd*2 is a missense mutation in a residue that forms part of the ATP-binding motif of Ncd: an *ncd*2–*gfp* transgene containing the missense mutation but not the other three amino acid changes of *ncd*2 causes mutant effects that parallel those of *ncd*2. The G446→R missense mutation of *ncd*2 affects a glycine residue that is highly conserved among the kinesin proteins and located at the base of a loop, L5, on the surface of the Ncd motor domain in the crystal structure (Sablin et al., 1996). Together with two other surface loops, L5 forms the entry to the nucleotide-binding pocket of Ncd. Replacement of G446 with a positively charged arginine is likely to affect nucleotide binding or release from the motor, and alter the ability of the motor to bind to or dissociate from microtubules. This is expected to impair the ability of the motor to function in the spindle by causing defective movement on microtubules and/or aberrant crosslinking activity.

The Ncd2–*gfp* fusion protein is associated with meiotic spindle fibers in *ncd*2–*gfp* mutant oocytes, as reported previously for Ncd (Hatsumi and Endow, 1992a), but the meiotic spindles are abnormal. Previous reports (Hatsumi and Endow, 1992b; Matthies et al., 1996) have focused on Ncd and the assembly and stability of the meiosis I spindle in *Drosophila* oocytes. Based on the observation of multiple or multipolar spindles in mutant oocytes, the Ncd motor has been proposed to crosslink and move on the microtubules associated with the meiosis I chromosomes, forming a single bipolar spindle (Hatsumi and Endow, 1992a,b; Matthies et al., 1996). Analysis of live oocytes injected with rhodamine–tubulin to visualize meiotic spindles indicates that the Ncd motor is required for maintenance, as well as assembly, of bipolar meiosis I spindles (Matthies et al., 1996).

The present study focuses on spindle dynamics after oocyte activation. Our observations demonstrate that multipolar spindles are formed by separation of spindle poles upon oocyte activation in the presence of a loss-of-function Ncd motor. Wild-type Ncd is therefore required to maintain spindle pole integrity during the meiotic divisions, probably by crosslinking and moving on microtubules, preventing sliding forces from disrupting the spindle and separating the microtubule-associated chromosomes from one another. Normal elongation of the meiosis I and II spindles was infrequently observed in *ncd* mutant oocytes, implying that the Ncd motor facilitates the microtubule sliding that is probably needed for spindle elongation and the rapid assembly of the meiosis II spindles in wild-type oocytes. Loss of Ncd function does not eliminate the ability of the central meiosis II spindle poles to form, but it is
probably the impaired ability of the spindle fibers to slide against one another that causes failure of the meiosis II spindles to assemble, resulting instead in the formation of short tripolar or multipolar spindles.

Sliding of spindle microtubules against one another has been proposed previously to contribute to poleward translocation, or flux, of microtubules in mitosis that may underlie poleward movement of chromosomes (Mitchison, 1989). Microtubule polymerization/depolymerization and microtubule motors are both thought to produce forces that result in poleward microtubule flux (Mitchison and Salmon, 1992). The observations reported here indicate that microtubule sliding is an important aspect of meiotic spindle dynamics as well as the dynamics of mitotic spindles, and is probably facilitated by the Ncd microtubule motor.

In addition to defective spindle elongation and meiosis II spindle assembly, the spindle microtubules appear to be destabilized in ncd−gfp* mutant oocytes. Fragmentation of microtubules was observed in activated live ncd−gfp* mutant oocytes, but not in activated live ncd−gfp or ncd−gfp* oocytes. Fluorescent particles, probably microtubule fragments bound to Ncd−GFP*, were associated with depolymerizing spindle fibers of ncd−gfp*, forming a hazy network in the center of the meiotic spindles, where the more unstable microtubule plus ends are expected to lie. Normally activated fixed ncd2−gfp* mutant eggs, stained with α-tubulin antibody to visualize microtubules, also showed a hazy mass of tubulin-positive fragments in the central regions of the meiosis I or II spindles, which was not observed in normally activated wild-type eggs fixed and stained with α-tubulin antibody. These observations indicate that the mutant Ncd2−GFP* motor causes destabilization of spindle fibers and imply that wild-type Ncd functions to stabilize spindle fibers during completion of the meiotic divisions. Destabilization of microtubules by Ncd2−GFP* could be due to altered ability of the mutant motor to bind to and move on microtubules, or to crosslink or bundle spindle fibers, as a consequence of the mutational change in its nucleotide binding site. It is also possible that Ncd2−GFP* exhibits gain-of-function effects, based on its small semidominant effect on embryo viability. The implied role of Ncd in stabilizing microtubules therefore warrants substantiation by further evidence.

Dynamics of Spindle Pole Formation and the Ncd Microtubule Motor

A recent model for microtubule motor protein function in spindle pole formation invokes the ability of the motor both to crosslink microtubules and move along the crosslinked microtubules (Vernos and Karsenti, 1995). The proposed requirement for movement of the motor on the crosslinked microtubules is consistent with the cytological effects of the mutant Ncd2−GFP* motor. Ncd2 can bind to spindle microtubules and could also bundle them by interactions with its highly charged tail region (Chandra et al., 1993), but movement of the mutant motor on microtubules is probably impaired, as evidenced by its apparent inability to mediate microtubule sliding needed for spindle elongation. The proposed impaired ability of Ncd2−GFP* to move on microtubules is correlated with the inability of the motor to maintain spindle poles, thus movement of the Ncd crosslinking activity along spindle microtubules is likely to be required to maintain spindle bipolarity.

Loss of Ncd function in ncd2−gfp* mutant oocytes causes misregulation of the meiotic divisions, resulting in continued divisions after the initial two divisions in in vitro–activated live mutant oocytes. Maternal chromosomes are also associated with spindles instead of the normal polar bodies in normally activated fixed mutant eggs. In wild-type embryos, the formation of polar bodies could sequester the maternal chromosomes from cytoplasmic factors, preventing further spindle-associated divisions. Loss of regulation over the number of meiotic divisions could be due to the inability of the mutant Ncd2 motor to focus the meiotic chromosomes into polar bodies, which consist of chromosomes with centromeres oriented inward, surrounded by an array of short microtubules (Hatsumi and Endow, 1992b; Komma and Endow, 1997; Page and Orr-Weaver, 1997). The polar bodies of Drosophila resemble monopolar spindles and may undergo assembly in a manner similar to that of meiotic spindle poles. The apparent requirement for Ncd to focus the microtubule-associated chromosomes into polar bodies reinforces the idea that both crosslinking activity and movement of the crosslinking activity along microtubules are required to maintain focused arrays of microtubules such as spindle poles and polar bodies. Spindle-associated maternal chromosomes are also observed in embryos of the cad null mutant (Hatsumi and Endow, 1992b), evidence that the absence of Ncd, as well as the loss of Ncd function, results in continued divisions of the oocyte chromosomes.

Several effects of the loss of Ncd function that are reported here were not detected in previous studies of antibody-stained fixed oocytes and embryos. These include failure of meiosis I spindle elongation, failure of meiosis II spindle assembly, and destabilization of spindle fibers. The ability to monitor meiotic spindle dynamics in live oocytes provides an important means of determining the effect of mutants on spindle assembly and dynamics. Further studies should lead to a complete picture of the role of microtubule motors like Ncd, and microtubule dynamics, in meiotic spindle assembly and function.

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