Research Article

Changes in γ-Tubulin Protein Distribution in Zebrafish (Danio rerio) Oocytes and the Early Cleavage-Stage Embryo

Jianxiong Liu¹,² and Charles A. Lessman¹,³

¹Department of Biology, The University of Memphis, Memphis, TN 38152, USA  
²Department of Physiology, University of Tennessee-Memphis, Memphis, TN 38163, USA  
³Department of Biological Sciences, The University of Memphis, 223 Life Science Building, Memphis, TN 38152-3560, USA

Correspondence should be addressed to Charles A. Lessman; clessman@memphis.edu

Received 1 October 2012; Accepted 23 October 2012

Copyright © 2013 J. Liu and C. A. Lessman. This is an open access article distributed under the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

We investigated the distribution of γ-tubulin in zebrafish oocytes and embryos using epifluorescent or confocal microscopy and γ-tubulin antibodies. During meiotic maturation of zebrafish oocytes, γ-tubulin begins redistribution from oocyte ooplasm and cortex to the future blastodisc region at the animal pole. In activated eggs, γ-tubulin was uniformly distributed in the enlarging blastodisc with label emanating from the yolk cell. In newly fertilized eggs, γ-tubulin was evenly distributed in blastomere cytoplasm, with the presence of pronuclei but initially lacking discernable centrosomes. During early cleavage, especially at the eight-cell stage, striking arc-shaped/rings (A/R) of putative centrosomes were detected. Decreasing γ-tubulin was seen in yolk cells while early cleavage blastomeres had strong cytoplasmic label along with obvious A/R arrays. In addition, we found the orientation of the A/R array and nuclear division alternated by about 90 degrees for each cell cycle along with alternation of punctate and A/R arrays.

1. Introduction

The microtubule cytoskeleton is essential for a variety of cellular processes, including cell movement, organelle transport, and cell division. Moreover, in oocytes and early embryos, microtubules have been implicated in localization of important embryonic determinants such as bicoid mRNA in Drosophila [1] and Vg1 mRNA in Xenopus [2] as well as trafficking cell components such as β-catenin in cleaving zebrafish embryos [3]. Recently, imaging of cytoskeleton in live zebrafish embryos has been described and the field reviewed [4]. The highly ordered microtubular array found in typical eukaryotic cells is organized by the microtubule-organizing center (MTOC). MTOCs organize microtubules by initiating noncovalent assembly of α-/β-tubulin heterodimers, anchoring them at their minus ends, and facilitating microtubule extension at the rapidly growing plus ends [5]. The morphology, subcellular localization, and molecular makeup of MTOCs vary across different species and different cell types within single species. The major MTOC in proliferating animal cells is the centrosome. Because of its central role in many essential aspects of cell physiology, both in interphase and during cell division, intense research activity has been carried out to characterize centrosomes [6–8]. Recently a review, describing centrosome and basal body research in zebrafish, has been published that enumerates the advantages of the zebrafish model in the study of MTOCs [9].

In most animal cells, centrosomes are composed of a pair of centrioles, surrounded by an amorphous cloud of electron-dense material, the pericentriolar material (PCM) [10]. An extensively characterized PCM protein required for MT nucleation is γ-tubulin. In a number of species, including Drosophila and Xenopus, γ-tubulin forms a macromolecular complex (called the γ-tubulin ring complex, γTuRC), with a characteristic ring structure (~25 nm in diameter), together with a variety of associated proteins [11–15]. Hundreds of γTuRCs tether to the centrosome scaffold, to serve as
the site of origin for MTs [15], and are required for spindle assembly and progression through mitosis [16–20]. Thus, the centrosome is the primary site for microtubule nucleation, and the centrosomal γTuRC is thought to be necessary for anchoring of MTs minus ends to centrosomes [21–24]. Using immunofluorescence and GFP reporter construct techniques, γ-tubulin was shown to be associated with centrosomes dynamically in mitotic cells, being massively recruited at prophase and released at anaphase-telophase. This accumulation in mitotic centrosomes is dramatic during the first embryonic divisions [25]. Moreover, fluorescence measurements suggest that the amount of antigenic γ-tubulin increases during mitosis and that the total amount of γ-tubulin in the spindle is larger than the amount of γ-tubulin in the spindle poles [26]. The cell stage distribution of γ-tubulin varies between animal cells of different species and between cells of different tissues within the same species. The function of the soluble cytoplasmic γ-tubulin remains unclear, and very little is known about how the γTuRC is assembled and tethered in the centrosome. Moreover, the redistribution of centrosomal proteins to specific sites of the cell is poorly understood, as are the mechanisms controlling microtubule nucleation within the living cell.

The zebrafish (Danio rerio), a small tropical freshwater teleost, has emerged as a model for cell and developmental biology because of its high fecundity, short generation time, and rapid development of the externally fertilized and translucent embryos [27]. We previously revealed the presence of soluble tubulin pools in zebrafish oocytes, which were associated with large-molecular-weight complexes, including α-, γ-tubulin and other proteins. These ovarian tubulin complexes were sequestered and maintained in a temporary “oligomeric” state and appear to be relatively stable storage forms of soluble tubulins, which subsequently may play important roles in microtubule dynamics in zebrafish oogenesis and embryogenesis [28].

Early embryonic, maternally regulated vertebrate cell cycles (i.e., cleavage) differ from somatic cell cycles. Early embryonic cells have rapid synchronous cell cycles with alternating M (mitosis) and S (DNA synthesis) phases and lack G (gap/growth) phases. The rapid repeated cleavages result in cells with successively smaller volumes. Zebrafish embryogenesis begins with 10 metasynchronous mitotic cycles [29], and the genome is silenced until the ninth cleavage cycle [30]. These early divisions are remarkably fast, with 15 minutes per cycle [31]. Subsequently, the cycles slow gradually, leading to a transition from maternal to zygotic control of the cell cycle in cycle 10, indicating that meiosis, mitosis, and many other essential processes required for embryonic development are regulated by maternal factors until this stage [32]. Since only 15 minutes are allotted for (1) the replication of the entire genome, (2) the assembly of the spindles with centrosomes, and (3) the segregation of chromosomes for each early cell cycle, it is remarkable that about two thousand centrosomes are calculated (i.e., a pair per dividing cell) to be assembled in less than 3 hours. Moreover, this stage of development relies primarily on maternally supplied materials and does not depend on zygotic gene expression until after the midblastula transition [30]. So all the components needed to build thousands of centrosomes should be preformed and laid down in the oocyte by the mother. Thus maternal regulation is a key feature of early development with many components necessary for embryogenesis already stored in the oocyte. This suggests that the oocyte, egg, and early embryo should be excellent sources of centrosome and spindle components for biochemical analysis. In a previous paper, using biochemical techniques, we demonstrated the presence of γ-tubulin along with α-tubulin in large, soluble complexes (~2 MDa) in zebrafish oocytes, eggs, and early embryos [28]. In this present study, using epifluorescent and confocal microscopy with monoclonal antibody against γ-tubulin (GTU-88), we aimed to reveal the cellular distribution of zebrafish γ-tubulin in oocytes and during early embryogenesis.

2. Results

2.1. Cellular Distribution of γ-Tubulin in Zebrafish Oocytes. To visualize the cellular distribution of γ-tubulin in zebrafish oocytes, the γ-tubulin proteins were detected specifically with immunofluorescence staining using monoclonal antibody GTU-88, followed by Alexa green-conjugated secondary antibody. Figure 1 illustrates the localization of γ-tubulin in fully grown oocytes during the time course of oocyte maturation in the latter induced with 17α-20β-dihydroxyprogesterone (DHP) in vitro. In fully grown, immature oocytes (i.e., prophase I of meiosis), under lower magnification of confocal microscopy, γ-tubulin signals were detected uniformly in the ooplasm and around the cortex area (Figure 1(a)). Upon maturation represented by the oocytes treated with DHP, much of the γ-tubulin remains uniformly distributed, but some signal starts to accumulate in the future blastodisc region as the oocyte proceeds to metaphase II of meiosis (Figures 1(b) and 1(c), representing 2 hours and 4 hours of DHP treatment, resp.). Furthermore, naturally matured and oviposited eggs activated artificially with water to release the cells from the metaphase II block revealed that γ-tubulin proteins were found uniformly distributed in the enlarged blastodisc area, with decreased staining in the yolk cell (Figure 1(d)). The images obtained from higher magnification revealed discreet γ-tubulin foci and diffuse labeling in cytoplasm of immature oocytes (Figure 1(e), arrow heads). Formation of the animal pole blastodisc (Figures 1(f) and 1(g)) and clearing of the opaque ooplasm are markers for meiotic maturation in the live cells, as is the dissolution of the oocyte nucleus or germinal vesicle (GV; Figures 1(j) and 1(k)) [33]. The γ-tubulin continued to stream from the yolk cell and into the blastomeres at the 4–8 cell stage as seen in confocal optical section (Figure 1(l)). Also apparent is the presence of diffuse γ-tubulin in the cytoplasm of the cleaving blastomeres along with intensely labeled putative centrosomes, while the yolk cell has markedly reduced label (Figure 1(l)). These data were strongly consistent with our in vivo hybridization results [34], which revealed that γ-tubulin mRNA was diffusely distributed along the oocyte cortex area, but became localized to the blastodisc of the animal pole in mature oocytes (i.e., eggs) and blastomeres in dividing embryos.
2.2. Cellular Distribution of γ-Tubulin in Zebrafish Embryos.

Figure 2 demonstrates the cellular distribution of γ-tubulin in early cleavage embryos. In newly fertilized 1-cell embryos, γ-tubulin was found to be uniformly distributed (stained green with GTU-88) in embryonic cytoplasm, with distinguishable 4′,6-diamidino-2-phenylindole (DAPI) stained pronuclei (arrow head, pseudocolored red). No obvious centrosomal structure was detected at this stage (Figure 2(a)). After approximately 30 minutes, when the embryo finished the first mitotic cycle and became a 2-cell embryo, clusters of putative centrosomal material with intense GTU-88 label were assembled around nuclei (Figure 2(b), arrow heads mark nuclei). The blastomere cytoplasm was also diffusely stained with GTU-88 around the brightly stained clusters of
\[\gamma\text{-tubulin}\]

\[\gamma\text{-tubulin}\] in putative centrosomal structures (Figure 2(b)). In most 4 to 8-cell embryos, centrosomes appeared with prominent curvilinear-array shapes (Figure 2(c)) associated with later mitotic phases (i.e., telophase). These interesting arc-shaped accumulations of \[\gamma\text{-tubulin}\] varied in length from about 10 to 25 \(\mu\text{m}\) or more.

In cleavage and blastula staged embryos, the \[\gamma\text{-tubulin}\] positive structures are varied in morphology ranging from punctuate foci (P) to “arc-shaped/ring” (A/R) structures (Figure 3). Interphase nuclei tended to have arc-shaped/ring clusters of \[\gamma\text{-tubulin}\] positive structures (Figures 3(a)–3(c)). Figure 3(d) shows an early blastula with both foci and arc-shaped/rings in different blastomeres. When double-labeled with DAPI and GTU-88, the mitotic blastomeres tended to have foci at the spindle poles enriched in \[\gamma\text{-tubulin}\] (i.e., conventional centrosomes; Figures 3(e) and 3(f)). In addition, the blastomeres all contained significant levels of diffuse \[\gamma\text{-tubulin}\] compared to the yolk cell suggesting continued partitioning of this protein.

The results support our previous hypothesis [28] that maternal, soluble tubulin, including \[\gamma\text{-tubulin}\], is accumulated and maintained in a temporary “oligomeric” form that is composed of large molecular weight complexes residing in the 40,000 \(\times\)g supernatant pool during zebrafish oogenesis. In addition, this pool of maternal tubulin complexes would be subsequently sequestered and reallocated into many structures (e.g., centrosomes) and incorporated into embryonic MT networks in ensuing embryogenesis [28].

2.3. Nuclei and Centrosomal Orientation during Zebrafish Early Embryogenesis. Since embryonic centrosomes were undetectable in the 1-cell stage then suddenly appeared during the first mitotic division, we wanted to further investigate their cellular distribution in dividing blastomeres in later embryonic stages. The image data demonstrated that the centrosome arrays alternated from curvilinear or arc-shaped to punctate in one direction, then from punctate to arc-shaped or curvilinear in another direction, while the nuclear division orientation alternated after each cell division (Figure 4). The striking curvilinear arrays of \[\gamma\text{-tubulin}\] appeared in blastomeres in late mitosis (i.e., telophase) during early cleavage (Figures 4(a), 4(b), and 4(c)). The schematic diagrams, indicating nuclei (red dots) and centrosomes (green lines or dots) for each embryonic stage, were derived from representative micrographs.
3. Discussion

3.1. γ-Tubulin and Centrosomes during Oogenesis and Embryogenesis. The results presented here, demonstrating a lack of typical γ-tubulin-enriched centrosomes in zebrafish oocytes, unfertilized eggs, and water-activated eggs, are consistent with findings for other species [35, 36]. These findings further suggest that eccentric spindle placement for polar body formation in zebrafish does not require γ-tubulin enriched centrosomes, since these were not seen in oocytes or eggs. Typical centrosomes consist of a centriole pair surrounded by pericentriolar material (PCM). Previous work suggested that centrioles are required to organize PCM to form a structurally stable organelle. The PCM contains a matrix of 12–15 nm fibers termed the “centromatrix” [37]. The “centromatrix” is tightly associated with the centrioles and directs the recruitment of other centrosomal components including ring-shaped complexes containing γ-tubulin [23, 37]. Thus, its centriolar stoichiometry may affect the rate of PCM recruitment or the retention of PCM at the centrosome by activating components for assembly or by affecting their turnover. In addition to control by a centriolar structural element, limiting levels of PCM components also influence centrosome size. SAS-4 is a centriole-associated component whose amount dictates centrosome size [38]. Such a regulatory agent may be responsible for the increased size of the γ-tubulin arrays seen here in early cleaving zebrafish embryos. An effect of PCM component limitation is expected because centrosomes get smaller when the ratio of cytoplasmic volume to centriole number is reduced during embryonic development [38].

Xenopus oocytes and eggs lack recognizable centrioles or centrosomes [39, 40]. The maternal centrosomes of Xenopus oocytes are inactivated early in stage I of oogenesis [40]. Despite lacking centrioles or definitive centrosomes, Xenopus eggs contain a substantial pool of centrosome components [41, 42] including γ-tubulin [19]. Using confocal immunofluorescence microscopy, γ-tubulin is found to be apparent surrounding the germinal vesicle (GV) of stage VI Xenopus oocytes [43]. It is also concentrated in the cortex of stage VI oocytes and it is bound to spindle microtubules only after full elongation of the meiotic spindle then γ-tubulin became heavily concentrated at the spindle poles [44]. In the animal hemisphere, γ-tubulin is evenly distributed as small cortical foci.

Centrosomes are not an essential component in the formation of the metaphase spindle during meiotic maturation of horse oocytes, but they can be introduced from the spermatozoon or donor cell and are necessary for the organization of normal embryonic development [45]. Centrosomes of the horse oocyte reorganize themselves from the beginning of GV stage to leave only PCM of γ-tubulin surrounding both poles of the MI and MII stage spindles [45].

In unfertilized mouse oocytes, meiosis is arrested in the second meiotic metaphase. γ-tubulin is found to be evenly distributed in both the cytoplasm and the germinal vesicle in the fully grown germinal vesicle (GV) stage oocytes. After GVBD, γ-tubulin dots were localized in both the cytoplasm...
and the vicinity of the condensed chromosomes aligned at both poles of the meiotic spindle at prometaphase I and metaphase I. At anaphase I and telophase I, γ-tubulin is detected between the separated chromosomes, while it is absent in the midbody. At the MII stage, γ-tubulin was again accumulated at the spindle poles. At an early stage of mouse fertilization, there was no γ-tubulin foci observed around the sperm chromatin or early male pronucleus [46]. At this stage, γ-tubulin is concentrated in the broad spindle poles of the meiotic spindle and at the distinct foci which form the centers of the cytoplasmic microtubule asters [47].

Besides Xenopus, oocytes lose centrioles during oogenesis in humans [48, 49], rhesus monkeys [50], cows [36], and many other mammals except rodents [35]. They also retain a stockpile of pericentriolar material (PCM), and microtubules are nucleated in the absence of centrioles in mammalian oocytes and early embryos [35]. This pool of γ-tubulin and centrosome components may play an important role in regulating MT organization during oocyte differentiation and animal-vegetal axis establishment [51]. In contrast, the spermatozoa lose most centrosomal proteins but retain centrioles during spermiogenesis [52]. The low abundance of endogenous γ-tubulin in most cells and the complexity of the seminiferous epithelial architecture make such evaluations in sperm cells extremely difficult [53]. In fact, we tried to visualize any possible centrosome structure by GTU-88 staining in zebrafish sperm but none was detected (Liu and Lessman, unpublished data). So the turnover of γ-tubulin or other centrosomal proteins during zebrafish spermatogenesis is still unclear.

Centrosomes are assembled at fertilization, in which the centriole pair, inherited from the male gamete, binds nucleating components from the female gamete [18] and matures by acquiring maternally derived pericentriolar material components [52]. During fertilization, a functional centrosome is likely a composite structure blending both paternal and maternal centrosomal components. The mixing of the maternal and paternal chromosomes occurs at the first mitosis. In species such as sea urchin, C. elegans, Xenopus, and zebrafish, a diploid nucleus forms through the fusion of two haploid pronuclei before the first mitosis [32, 54–56]. The process of pronuclear fusion occurs in zebrafish zygotes between 14 and 20 minutes postfertilization (mpf) [32]. In another cyprinid species, the common white sucker (Catostomus commersoni), an aster begins forming by 30 mpf at one side of the fertilizing sperm cell within the ooplasm of the blastodisc. Formation of the male pronucleus, with growing aster, and an asterless female pronucleus is complete by 60 mpf in this cool-water species (16–18 °C) [57].

In Caenorhabditis elegans, γ-tubulin is recruited around sperm centrioles after fertilization and accumulates massively at the centrosome during prophase of the first embryonic
division. The centrosome undergoes striking morphological changes from metaphase to interphase. The compact structure visualized in metaphase changes into a spindle-like shape in anaphase before losing most of the \( \gamma \)-tubulin localization in telophase. In interphase, the \( \gamma \)-tubulin signal is similar in size to that of \( \alpha \)-tubulin, suggesting that most of the \( \gamma \)-tubulin protein left resides inside, or very near, the centriole. Concomitantly with these events, the number of microtubules nucleated at the centrosomes decreases considerably from telophase [25]. This “catastrophic” disassembly of \( \gamma \)-tubulin from the centrosome as cells exit mitosis has been reported in mammalian cells and is likely to be due to the release of most of the pericentriolar material into the cytoplasm [58]. The results suggest that the origin of mitotic spindle poles is associated with MTOCs near the condensed chromosomes and that spindle \( \gamma \)-tubulin is recruited from the cytoplasm during the process of spindle assembly.

In the mouse model, after fertilization, \( \gamma \)-tubulin foci in the egg cytoplasm and spindle poles were transformed to amorphous \( \gamma \)-tubulin and translocated from spindle poles to the area between the separating chromatids. When the pronuclei became enlarged and got close to each other, many \( \gamma \)-tubulin dots in the vicinity of the pronuclei were observed. The accumulation of \( \gamma \)-tubulin around the pronuclei may be responsible for extensive microtubule assembly in this region. It aggregated into some dots in interphase but was distributed on the mitotic spindle poles in early embryos [46]. The integrity of these \( \gamma \)-tubulin foci and their cytoplasmic location is maintained during drug- or cold-induced depolymerization of microtubules [59].

In unfertilized bovine oocytes, \( \gamma \)-tubulin was identified in the cytoplasm, mainly in the cortex and concentrated in the meiotic spindle. Following sperm penetration, \( \gamma \)-tubulin in the cytoplasm was recruited by a sperm component to reconstitute the zygotic centrosome. During pronuclear apposition, \( \gamma \)-tubulin was localized as spots at the spindle poles. \( \gamma \)-tubulin spots were also observed in blastomeres of embryos cleaved in vitro. In the later pronuclear stage, considerably less \( \gamma \)-tubulin and fewer microtubules were detected in the cytoplasm. At the mitotic metaphase of parthenotes, \( \gamma \)-tubulin was recruited to the condensed chromatin and concentrated in the spindle. In the absence of sperm components, the cell cycle-related assembly of \( \gamma \)-tubulin organizes microtubule nucleation for positioning the pronucleus and spindle protein of mitotic metaphase during the first cycle of bovine parthenotes [60].

When the \textit{Xenopus} egg is fertilized, the centrosome is assembled from both paternal components (the centriole itself) and maternal components (\( \gamma \)-tubulin; [61]). The latter are stored in the oocyte cytoplasm during oogenesis in sufficient quantities to contribute to the formation of thousands of centrosomes until the onset of massive zygotic gene expression, at the midblastula transition [41, 62]. Extracts prepared from embryos taken later in development actually cannot assemble centrosomes into sperm centrioles. Our current data suggest that the assembly of zebrafish early embryo may adopt the similar pathway as \textit{Xenopus}.

In summary, the distribution of \( \gamma \)-tubulin in animal oocytes and early embryos is more complex than initially assumed. Although the mechanism of recruitment of \( \gamma \)-tubulin to the sperm component is currently unclear, it appears to be true that a model of blended zygotic centrosome composed of maternal constituents is introduced to a paternal template after insemination [63]. The present study revealed no centrosomal structures assembling after egg activation with water alone, suggesting the requirement of the sperm for centrosome assembly in zebrafish. Our data are consistent with the idea that maternal \( \gamma \)-tubulin is recruited around sperm centrioles after fertilization and accumulates at the centrosomes during the first few embryonic divisions forming a variety of \( \gamma \)-tubulin-containing structures including punctate foci, arc-like/rings, and linear arrays. The sperm centrosome is required to organize the radial array of microtubules, called the sperm aster, within the inseminated oocyte of nonrodent mammals. Thus, specific regulatory events must take place at fertilization to turn on the nucleating activity of the sperm centrosome.

3.2. \( \gamma \)-Tubulin and Cell Cycle Regulation in Oocytes and Embryos. The colocalization of \( \gamma \)-tubulin and phosphorylated MAP kinase with microtubule assembly in both control and taxol-treated pig oocytes suggests that emanation of microtubules from the chromosomes may be regulated/directed by microtubule-organizing material including \( \gamma \)-tubulin and phosphorylated MAP kinase in pig oocytes [64].

In newt embryos, cyclin B1 begins to accumulate in the nucleus during interphase in synchronous cleavage, and its greatest expression is in the centrosomes and the nucleus at prometaphase. The centrosomes of the principle sperm nucleus and the zygote nucleus have greater accumulation of both \( \gamma \)-tubulin and cyclin B1 compared with the centrosomes of the accessory sperm nuclei [65]. These data suggest that cyclin B1 may play roles in regulating \( \gamma \)-tubulin distribution in dividing embryos.

Establishment of polarity in \textit{C. elegans} embryos is dependent on the centrosome. The sperm contributes a pair of centrioles to the egg, and these centrioles remain incapable of polarizing the cortex while the egg completes meiosis. Coincident with the establishment of polarity, the centrioles recruit centrosomal proteins, several of which are required for polarity, suggesting that temporal regulation of centrosome assembly may control the initiation of polarization. They found that cyclin E-Cdk2 controls the recruitment of centrosomal proteins specifically at the time of polarity establishment. Cyclin E is required for several examples of asymmetric cell division and fate determination in \textit{C. elegans} and \textit{Drosophila} [66].

The cycles of centriole and centrosome duplication are intimately coupled. Centrioles initiate duplication at the onset of S phase in a process that requires Cdk2 (reviewed in [67]). At this time a generative disc, a precursor of the daughter centriole, forms at right angles to the proximal end of each mother centriole. Daughter centrioles elongate as the cell cycle progresses, reaching full length by G2/M. During the G2/M transition, the centrosome also matures, accumulating \( \gamma \)-tubulin and other centrosomal components, and...
increasing in size and nucleating capacity (reviewed in [68]). Cdk2 is not required for normal centrosome duplication, maturation, and bipolar mitotic spindle formation in mouse embryonic fibroblast cell. In contrast, Cdk2 deficiency completely abrogates aberrant centrosome duplication induced by a viral oncogene. These results indicate that normal and abnormal centrosome duplications have significantly different requirements for Cdk2 activity and point to a role of Cdk2 in licensing centrosomes for aberrant duplication [69].

4. Methods

4.1. Zebrafish. Wild-type zebrafish were obtained from a local distributor and housed in a dedicated, temperature (28°C) and photoperiod (14L) controlled fish room in racks of food-quality plastic containers (6 and 12 l) with flow-through dechlorinated water. Fish were fed 1-2 times daily a diet of TetraMin and dried brine shrimp flakes. This research was carried out under an IACUC approved protocol.

4.2. Fully Grown Oocyte Collection. Gravid adult female zebrafish were decapitated and the spinal cords pithed, the whole ovary was removed and placed in a Petri dish half filled with Cortland’s solution (NaCl 7.25 g, CaCl₂ 2H₂O 0.23 g, KCl 0.38 g, MgSO₄·7H₂O 0.23 g, NaHCO₃ 1.0 g, penicillin 30 mg, and streptomycin 50 mg made up to 1000 mL with double distilled water). Then the opaque fully grown oocytes were dissected from the ovary with watchmaker forceps and pooled into a 1.5 mL microcentrifuge tube.

4.3. Oocyte Maturation In Vitro. The opaque fully grown oocytes were separated from the others following the above procedure, pooled into a 60 × 15 mm Petri dish with 10 mL fresh Cortland’s solution with 1 μg/mL 17α- and 20β-dihydroxyprogesterone (DHP) dispensed in steroid vehicle (EtOH:propylene glycol, 1:1), and incubated at 28°C for 6 hr. Aliquots of these in vitro maturing oocytes were then collected at different time points and pooled separately into a 1.5 mL microcentrifuge tube. The oocytes were then fixed in freshly made 4% paraformaldehyde at 4°C overnight. On the following day, the methanol was gradually added until they were brought to 100% methanol. Then the oocytes were stored in 100% methanol at −20°C until used for the next step in the analysis.

4.4. Embryo Collection. Male and female fish were placed in separate compartments within spawning tanks, and the separation barriers were removed just prior to the desired mating time. This allowed collection of embryos at precise times after fertilization. Crosses were set up in the afternoon using 3 females and 4-5 males per breeding tank. After mating, males and females were separated again and allowed to rest for at least one week before the next cross. Embryos were collected next morning and were washed to remove any debris [70]. Standard fish water [31] supplemented with methylene blue (final concentration was 2 ppm) was used for the washes and maintained at 28°C. Washed embryos were grouped according to the zebrafish stage tables [31] and collected into separate Eppendorf tubes. For further drug treatment, the embryos were grouped into Petri dishes and treated with different drugs. All the sample embryos were then fixed and dehydrated following the above procedures outlined for oocytes.

4.5. Antibody. Anti-γ-tubulin mouse monoclonal antibody GTU-88 (mouse IgG1 isotype; Sigma Chem. Co., St. Louis, MO) recognizes EEFATEGTDRKDVFFY located in position 38–53 of the amino acid sequence of γ-tubulin near N-terminal end. GTU-88 reacts well to zebrafish, [28, 32] human, bovine, dog, hamster, rat, mouse, chicken, and Xenopus γ-tubulin [71].

4.6. Immunofluorescence Microscopy. Before incubation in primary antibody, the fixed oocytes or embryos were rehydrated into 100% PBS in a graded series. The chorions of embryos were manually removed with watchmaker forceps after rehydration. Then the sample oocytes or embryos were incubated in 5% nonfat milk (Carnation) in PBS solution for 4–6 hours at room temperature with gentle rotation. After the samples were washed several times with PBS, they were incubated with 1 mL 1: 200 primary antibody GTU-88 overnight at 4°C. Then the oocytes or embryos were washed in PBS for 8 hours at room temperature, changing buffer at 1-hour intervals. The embryos were then incubated with 750 μL 1: 200 secondary Alexa green-conjugated-anti-mouse antibody overnight at 4°C. After the oocytes or embryos were washed several times with PBS (for 8 hours at room temperature, changing buffer at 1-hour intervals), the samples were incubated with 1 mL 4’,6-diamidino-2-phenylindole (DAPI) solution (final concentration is 2.5 μg/mL) for 5 minutes to stain DNA, with gentle rotation at room temperature. Then the staining solution was decanted, and the embryos were washed with PBS for 8 hours to overnight at room temperature. The oocytes or embryos were then dehydrated gradually with 100% methanol, three or four changes over 60–90 minutes at room temperature. After the methanol was removed, the samples were cleared with Benzyl Benzoate : Benzyl alcohol (BB : BA = 2 : 1) solution and mounted on slides in BB : BA solution. Then the oocytes or embryos were examined using either a Nikon laser scanning confocal microscope or Nikon eclipse 400 epifluorescence microscope. Digital RGB images captured with cameras mounted on the microscopes were processed (i.e., RGB split/merge), and iterative deconvolution was performed on images to remove unfocused light using the ImageJ program (http://rsb.info.nih.gov/ij/). Generally, for epifluorescence images, 3 iterations were computed using the 2D deconvolution and diffraction limit point-spread-function (psf) ImageJ plug-ins available at http://www.optinav.com/imagej.html.

4.7. Confocal Microscopy. Embryos, oocytes, and eggs were examined on a Nikon C1 laser confocal microscope equipped with Nikon triple-laser scanning assembly using Argon (ex.488 nm), green helium-neon (ex.543 nm), and red
helium-neon (ex.633 nm) lasers. Objectives of 10X and 20X were used. Some images presented are projections of up to three optical sections, providing increased depth of focus.

**Abbreviations**

MT: Microtubule  
DHP: 17α,20β-dihydroxyprogesterone  
MTOC: Microtubule organizing center  
γ-TuRC: γ-Tubulin ring complex  
DAPI: 4′,6-Diamidino-2-phenylindole  
GV: Germinal vesicle  
Bl: Blastodisc.

**Conflict of Interests**

The authors have no financial or other conflict of interests regarding this work.

**Acknowledgments**

This work was supported in part by the Department of Biology, the University of Memphis Graduate School, a Faculty Research Grant from the University of Memphis, and the American Society for Cell Biology Minority Affairs Council. The authors thank the Integrated Microscopy Center, the University of Memphis, for their support in confocal microscopy analysis.

**References**


[57] C. A. Lessman and C. W. Huver, "Quantification of fertilization-induced gamete changes and sperm entry without egg

---

*ISRN Developmental Biology*


