Multiple Pools of Nuclear Actin

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ABSTRACT

While nuclear actin was reported ~50 years ago, its in vivo prevalence and structure remain largely unknown. Here, we use Drosophila oogenesis, that is, follicle development, to characterize nuclear actin. We find that three different reagents—DNase I, anti-actin C4, and anti-actin AC15—recognize distinct pools of nuclear actin. DNase I labels monomeric or G-actin, and, during follicle development, G-actin is present in the nucleus of every cell. Some G-actin is recognized by the C4 antibody. In particular, C4 nuclear actin colocalizes with DNase I to the nucleolus in anterior escort cells, follicle stem cells, some mitotic follicle cells, and a subset of nurse cells during early oogenesis. C4 also labels polymeric nuclear actin in the nucleoplasm of the germline stem cells, early cystoblasts, and oocytes. The AC15 antibody labels a completely distinct pool of nuclear actin from that of DNase I and C4. Specifically, AC15 nuclear actin localizes to the chromatin in the nurse and follicle cells during mid-to-late oogenesis. Within the oocyte, AC15 nuclear actin progresses from localizing to puncta surrounding the DNA, to forming a filamentous cage around the chromosomes. Together these findings reveal that nuclear actin is highly prevalent in vivo, and multiple pools of nuclear actin exist and can be recognized using different reagents. Additionally, our localization studies suggest that nuclear actin may regulate stemness, nucleolar structure and function, transcription, and nuclear structure. Such findings call for further studies to explore the prevalence, diversity, and functions of nuclear actin across tissues and organisms. Anat Rec, 2018. © 2018 Wiley Periodicals, Inc.

Key words: nuclear actin; oogenesis; Drosophila; oocyte; nucleolus

In addition to its well-established cytoskeletal roles, actin also localizes to and functions within the nucleus. Nuclear actin was first reported over 50 years ago (Ohnishi et al., 1963, 1964; Lane, 1969). These findings were met with skepticism. These studies largely relied on subcellular fractionation, raising concerns about the purity of the isolated nuclei. The levels of actin in the nucleus were also low compared to that in the cytoplasm. Additionally, nuclear filamentous actin (F-actin) could not be visualized, and functions for nuclear actin had not been identified. More recent studies have overcome these issues. Indeed, the active mechanisms regulating the nuclear localization of actin have been identified (Wada et al., 1998; Stuven et al., 2003; Dopie et al., 2012; Mun-sie et al., 2012). Additionally, functional studies have revealed nuclear actin regulates transcription (Pomproix and Percipalle, 2004; Hofmann et al., 2004; Hu et al., 2004; Philimonenko et al., 2004), chromatin remodeling...
(reviewed in Venit et al., 2018), nuclear organization and structure (Sasseville and Langelier, 1998; Wasser and Chia, 2000; Krauss et al., 2003; Holaska et al., 2004), DNA damage repair (Andrin et al., 2012; Belin et al., 2015; Serebryannyy et al., 2017; Wang et al., 2017), cell cycle (Goyal et al., 2011; Baarlink et al., 2017), and meiosis (Bogolyubova and Ginzburg, 2013; Mogessie and Schuh, 2017). Nuclear actin has also been implicated in diseases, including cancer (Spencer et al., 2011; Fiore et al., 2017), neurodegeneration (Minamide et al., 2000; Maloney et al., 2005; Lim et al., 2007; Maloney and Bamburg, 2007; Munsie et al., 2011), and myopathies (Goebel and Warlo, 2001; Domazetovska et al., 2007a, 2007b; Serebryannyy et al., 2016c). Thus, actin has many critical functions within the nucleus that are important not only for key cellular functions but also for human health.

As the functions of actin within the cytoplasm depend upon its structure, it is likely that the structure of nuclear actin similarly impacts its activities. However, insight into the structures of nuclear actin remains limited. This limitation is due to the complexities involved in visualizing actin. Actin can exist as monomers (G-actin), short polymers, filaments (F-actin), and networks of filaments. It also binds to a large number of proteins. These different structures and complexes make it unlikely that any tool or reagent, such as an antibody, will recognize all of the actin within the cell. Indeed, a study using fluorescent recovery after photobleaching and fluorescence correlation spectroscopy of green fluorescent protein (GFP)-tagged actin indicates there are at least two pools of nuclear actin with different dynamics, suggesting nuclear actin exists as both monomers and polymers (McDonald et al., 2006). Another study generated nuclear actin probes using actin binding domains from different proteins, and identified probes that differentially label monomeric versus polymeric nuclear actin. Monomeric actin was observed in nuclear speckles and sites of RNA processing, while polymeric actin seemed to generate a viscoelastic structure (Belin et al., 2013). Polymeric nuclear actin visualized by this same tool has recently been implicated in DNA damage repair (Belin et al., 2015; Wang et al., 2017). Numerous antibodies have also been used to examine nuclear actin. Gonsior et al. (1999) generated an antibody (2G2) that labels nuclear dots in myogenic cells and fibrils in the nuclei of Xenopus oocytes (Gonsior et al., 1999). Similarly, another study generated an additional antibody (1C7) and found that it labeled distinct nuclear structures when compared to 2G2 (Schoenenberger et al., 2005). The finding that different antibodies label distinct subsets or pools of nuclear actin has been widely observed across cell types and organisms (Grenklo et al., 2004; Jockusch et al., 2006; Cruz and Moreno Diaz de la Espina, 2009; Asumda and Chase, 2012). It is important to note that fixation conditions dramatically affect the ability of actin antibodies to bind to the different actin structures; for example, methanol fixation denatures actin filaments, making them accessible to antibodies, but prevents phalloidin labeling. While the last two decades have significantly increased our understanding of the structures of nuclear actin, much remains to be determined, including how the structure of nuclear actin impacts its different functions.

Both the research into the structures of nuclear actin and its functions have largely been restricted to cultured cells, oocytes, or unicellular organisms; thus, the prevalence, structures, and functions of nuclear actin in vivo, multicellular, and developmental contexts remain largely unknown. To address this knowledge gap and advance our understanding of nuclear actin, here we utilize the robust and well-characterized process of Drosophila oogenesis, that is, follicle development. Oogenesis is an excellent system to explore the developmental functions of nuclear actin, as the whole process, from germline and somatic stem cells to a mature ready to ovulate eggs, is easily observed (Fig. 1). See below for a detailed description of follicle development. Briefly, each developing follicle is composed of multiple cells types, including 16-germline-derived cells, 15 nurse or support cells and one oocyte, and a somatic epithelium of follicle cells (Fig. 1B,D). The temporal progression of both cellular events and morphogenesis occurring within these different cell types is well-established. Thus, this knowledge can be used to provide insight into the potential functions of nuclear actin.

To visualize nuclear actin during Drosophila oogenesis, we utilize three reagents—fluorescently conjugated-DNase I, anti-actin C4, and anti-actin AC15. DNase I binds to and, thus, can be used to label monomeric or G-actin (Hitchcock, 1980). Anti-actin C4 is a broad specificity actin antibody (Lessaard, 1988) that has been used to examine nuclear actin in other systems (Parkenov et al., 1995; Hofmann et al., 2004; Gedge et al., 2005; Lenart et al., 2005; Spencer et al., 2011; Maslova and Krasikova, 2012). Additionally, we recently uncovered that anti-actin C4 labels some nuclear actin during Drosophila oogenesis (Kelpsch et al., 2016). Anti-actin AC15 (Gimona et al., 1994) also recognizes nuclear actin in other systems, including cultured Drosophila cells (Hofmann et al., 2004; Cruz and Moreno Diaz de la Espina, 2009; Miyamoto et al., 2011; Dopie et al., 2012). Here, we define the cellular and subnuclear localization of the nuclear actin recognized by the three reagents. By comparing the labeling of nuclear actin by the two antibodies to that of DNase I, we can gain insight into the structure of the nuclear actin recognized by the different antibodies. Additionally, the subnuclear, cell-type, and developmental/temporal localization of the reagents provides insight into potential functions of nuclear actin.

We find that while nuclear G-actin is ubiquitous throughout follicle development, as DNase I labels a nuclear structure in every cell, the two antibodies recognize more restricted and distinct pools of nuclear actin. Indeed, C4 nuclear actin is observed in a subset of cells through follicle development, as DNase I labels distinct nuclear regions. This finding suggests that anti-actin C4 recognizes both monomeric and polymeric actin. The G-actin labeled by DNase I and anti-actin C4 is primarily within the nucleolus, suggesting roles of nuclear actin in the function of this nuclear body. Furthermore, the cell-type and pattern of polymeric C4 nuclear actin suggests it may regulate differentiation and nuclear structure. The nuclear actin recognized by anti-actin AC15 is different from that labeled with either of the other reagents. AC15 nuclear actin is observed in all cells from mid to late oogenesis, largely associates with the chromatin, and does not colocalize with DNase I. These findings indicate AC15 nuclear actin is likely polymeric and suggests it may have roles in transcription. Together our study reveals that multiple pools of nuclear actin are observed during an in vivo, developmental
process, and provides insight into the potential functions of nuclear actin. Such findings are likely to be applicable to other developmental processes and systems.

MATERIALS AND METHODS

Fly Stocks

Fly stocks were maintained on cornmeal-agar-yeast food at 21°C. Prior to immunofluorescence analysis, flies were fed wet yeast paste daily for 3–4 days. yw was used as the wild-type background. Fly lines expressing CRISPR-mediated enhanced GFP (eGFP)-tagged nuclear proteins, Fibrillarin and Nopp-140, were a generous gift from Eric Wieschaus (Falahati and Wieschaus, 2017).

Immunofluorescence

Whole-mount Drosophila ovary samples were dissected into Grace’s insect media (Lonza, Walkersville, MD) and were fixed for 10 min at room temperature in 4% paraformaldehyde in Grace’s insect media. Briefly, samples were blocked by washing in Triton antibody wash (1X phosphate-buffered saline (PBS), 0.1% Triton X-100, and 0.1% bovine serum albumin) six times for 10 min each at room temperature. Primary antibodies were incubated for a minimum of 20 hr at 4°C. The following antibodies and concentrations were used: mouse anti-actin C4 1:50 (EMB Millipore, Billerica, MA); mouse anti-actin AC15 1:50–1:100 (Sigma-Aldrich, St. Louis, MO); rabbit anti-GFP 1:5000 (pre-absorbed on yw ovaries at 1:20 and used at 1:500; Torrey Pines Biolabs, Inc., Secaucus, NJ). After six washes in Triton antibody wash (10 min each), secondary antibodies were incubated from 12–36 hr at 4°C. The following secondary antibodies were used at 1:500: AF488::goat anti-mouse, AF568::goat anti-mouse, or AF488::goat anti-rabbit (Life Technologies, Grand Island, NY). For DNase I staining, AF488-DNase I 1:500 (Life Technologies) was included in both the primary and
RESULTS

Overview of Drosophila Oogenesis

The prevalence of nuclear actin, its structure, and its subnuclear localization during Drosophila oogenesis are largely unknown. Here, we utilize three reagents—DNase I, anti-actin C4, and anti-actin AC15—to characterize nuclear actin throughout follicle development. DNase I binds to and labels monomeric or G-actin (Hitchcock, 1980), while the two antibodies are reported to have the ability to label both monomeric and filamentous actin (F-actin). Here, we define the developmental, cell-type, and subnuclear distribution of the pools of nuclear actin recognized by the different reagents. This information, in conjunction with the vast knowledge of the processes occurring during oogenesis, provides insight into the potential functions of nuclear actin.

Drosophila oogenesis is a well-characterized and organized process of development (Spradling, 1993). Each adult female has a pair of ovaries (Fig. 1A). These ovaries are comprised of ~15 ovarioles, chains of sequentially developing follicles (Fig. 1A,B). This means every stage of follicle development is seen multiple times from a single female. At the anterior tip of the ovariole is a structure called the germarium (Fig. 1B,C). The germarium houses the stem cells that give rise to the individual follicles. The germarium is broken into four regions—1, 2a, 2b, and 3. In region 1, at the anterior of the germarium, there are two to three germline stem cells (GSCs, Fig. 1C, bright cyan) that are supported by a somatic niche including the terminal filament (dark gray), cap (light gray), and escort cells (red). The GSCs give rise to a cystoblast (light cyan) that goes on to have four incomplete cell divisions to generate an interconnected 16-cell cyst (light blue, Fig. 1C, regions 2a/b and 3); the cells are connected by ring canals which are remnants of the incomplete cytokinetic furrows. As the cysts develop they are wrapped by the escort cells (red) through region 2a (Fig. 1C). At the 2a-2b boundary, two follicle stem cells (FSCs, dark green) reside (Fig. 1C). The FSCs give rise to all the follicle cells (light green) that go on to surround and encase the 16-cell cysts (Fig. 1C). Within the 16-cell cyst, one cell will become the oocyte (dark blue nucleus), while the other 15 cells become nurse cells (light blue nuclei), which support the growing oocyte; this specification of cell fate occurs in region 3 of the germarium (Fig. 1C). The 16-cell cyst covered in follicle cells (light green) then buds off the germarium to generate a Stage 3 (S3) follicle (Fig. 1B).

During the early stages of oogenesis, the follicles grow in length and in somatic or follicle cell number (Fig. 1B). The follicle cells proliferate through S6 and then endocycle from S7-9, replicating their DNA without dividing (Claycomb and Orr-Weaver, 2005). The follicle cells differentiate into six major subtypes: stalk, polar, border, stretch follicle, centripetal follicle, and main body follicle cells (Dobens and Raftery, 2000). The stalk cells are the cells that connect the developing follicles and are not shown in Figure 1 or examined in this study. The polar cells are four cells, two at the anterior and two at the posterior tip of each follicle (purple, Fig. 1B,D). During S9, the posterior polar cells specify a group of 6–8 surrounding cells to be border cells (yellow border cells, Fig. 1D). The border cells then migrate posteriorly between the nurse cells to the nurse cell-oocyte boundary (see yellow border cell cluster in S10B in Fig. 1B,D); the border cells ultimately move dorsally and will secrete the eggshell structure called the micropyle, which is where the sperm enters (Montell, 2003). During S9, while the border cells are migrating, the majority of the other follicle cells will become the main body follicle cells (light green), which ultimately cover the oocyte (Fig. 1D). The remaining ~50 cells become the squamous stretch follicle cells (dark orange) that cover the nurse cells (Fig. 1B,D). During S10B, the anterior main body follicle cells differentiate into centripetal follicle cells (light orange) and migrate to enclose the anterior edge of the oocyte (Fig. 1B,D). During S10B-12, the border, centripetal follicle and main body follicle cells undergo gene amplification to over-replicate the regions of the genome encoding eggshell genes and begin to secrete the eggshell toward the oocyte (Claycomb and Orr-Weaver, 2005).

The germline nurse cells do not divide, but undergo endocycling during S3–10 (Spradling, 1993). The nurse cells are responsible for producing mRNA, proteins, and organelles that will be supplied to the oocyte in S10B-11, in a process termed nurse cell dumping. During nurse cell dumping (S11), the nurse cells squeeze their cytoplasmic contents into the oocyte. This process results in the nurse cells decreasing in size and the oocyte growing in length (see S10B–S12, Fig. 1B). During the subsequent S12–13, the nurse cells undergo a complex process of cell death and are phagocytized by the stretch follicle cells (dark orange), while the centripetal (light orange), border (not illustrated), and main body (green) follicle cells secrete the eggshell structures (Fig. 1B). It is important to note that the process of germline cells supplying their contents into the surviving oocyte is conserved to mammals (Lei and Spradling, 2016). The final stage of oogenesis, S14, is the mature, ready to ovulate egg, which is comprised of the oocyte, eggshell, and surrounding follicle cells (Fig. 1B). During ovulation, these follicle cells will be sluffed off and remain at the base of the ovariole, where...
they serve as an endocrine organ, much like the corpus luteum in mammals (Deadly et al., 2015). Using this knowledge, we can gain significant insight into the potential functions of nuclear actin by simply assessing which cells and when during follicle development the different reagents label nuclear actin.

**Nuclear G-Actin Is Ubiquitous**

DNase I labeling reveals that while high levels of G-actin are seen in the cytoplasm, lower levels of G-actin are present in the nuclei of all cells throughout follicle development (Figs. 2A–A’ and 3). Note that while all the cells exhibit nuclear G-actin, marks are used to point out specific cells in which the DNase I-labeled structure is readily apparent in the focal plane shown. In the germarium (Fig. 2B,B’), DNase I labels a single nuclear spot in all the early germ cells from the GSCs (yellow dashed circles) to the 16-cell cysts (yellow arrows). These 16-cell cysts will differentiate into one oocyte and 15 nurse cells. Upon differentiation, in region 3 of the germarium, the nuclear structure labeled within the nurse cells becomes more irregular in shape (Fig. 2B,B’, yellow arrow). This irregular appearance gets more pronounced and takes on a tubular structure during S3–10 (Fig. 2A–A’,C–D’, yellow arrows), and then becomes progressively more diffuse in the final stages of oogenesis, S11–13 (Fig. 3, yellow arrows). Notably, this nuclear G-actin does not overlap with the chromatin.

The DNase I labeling in the oocyte is distinct from that of the nurse cells. From region 3 of the germarium to S5, there is a single spot of DNase I staining directly adjacent to the DNA (Fig. 2E–E”, red arrowhead and data not shown). From S6 onward, this spot is no longer observed and only a nuclear G-actin haze is present (Fig. 2F–F’); this nuclear actin haze does not overlap with the chromatin.

DNase I also labels nuclear G-actin within the somatic cells during oogenesis. A single spot is labeled within the escort cells in the anterior of the germarium (data not shown and see Fig. 7K’, blue arrowheads), and in the follicle cells, from the stem cells (data not shown and see Fig. 7L’, blue dashed oval) throughout development (Fig. 2 and insets in Fig. 3A’–D’, blue arrows). Again, these foci do not overlap with the DNA.

Together these findings indicate that during *Drosophila* follicle development, all cells have nuclear G-actin. Given that this G-actin is within a single nuclear structure within each cell and does not overlap with the chromatin, it suggests G-actin is within a nuclear body.

**One Actin Antibody (C4) Labels a Subset of Nuclear Actin during Early Oogenesis**

The C4 actin antibody has broad specificity. While it was raised against chicken gizzard muscle actin, it reacts with all isoforms of vertebrate actin and actins from lower eukaryotes including *Dictyostelium* and slime mold (Lassard, 1988). The antibody can label both G- and F-actin. The epitope recognized by the C4 antibody is within the N-terminus of actin, and has been reported to be near amino acids 50–70 (Lassard, 1988; Gedge et al., 2005). Importantly, numerous studies have utilized the C4 antibody to examine nuclear actin (Parfenov et al., 1995; Hofmann et al., 2004; Gedge et al., 2005; Lenart et al., 2005; Spencer et al., 2011; Maslova and Krasikova, 2012). Furthermore, we previously found that it labels nuclear actin during the early stages of *Drosophila* oogenesis (Kelsch et al., 2016). Thus, the C4 actin antibody is well-suited to study nuclear actin.

During follicle development, anti-actin C4 weakly labels the cytoplasm and strongly labels a subset of the nuclear actin, subsequently referred to as C4 nuclear actin (Figs. 4 and 5). In region 1 of the germarium (Fig. 4A–B’), high C4 nuclear actin levels are observed in the GSCs (yellow dashed circles) and the early cystoblasts (white arrows). Conversely, C4 nuclear actin is not observed in the developing cysts in regions 2A and 2B of the germarium. This finding leads us to speculate that C4 nuclear actin may help maintain an undifferentiated state. Upon differentiation of the 16-cell cyst into an oocyte and 15 nurse cells, C4 nuclear actin labels an irregularly shaped structure within a subset of the nurse cells (Fig. 4A–A’ and S3 in C–C’, yellow arrows). This C4 nuclear actin is also observed in some of the nurse cells through S10 and exhibits a more tubular appearance as development progresses (Figs. 4C–D” and 5A–A’, yellow arrows). We previously observed that the frequency of nurse cells exhibiting C4 nuclear actin decreases with developmental stage (Kelsch et al., 2016). Notably, like the DNase I-labeled nuclear G-actin, the C4 nuclear actin does not overlap with the chromatin. During S11 and 12, a ring of C4 nuclear actin is observed just inside the nuclear envelope (Fig. 5B–C’, magenta arrows). Conversely, no C4 nuclear actin is observed during S13, and instead C4 labels puncta either within the nurse cell cytoplasm or the extensions of the stretch follicle cells surrounding them (Fig. 5D,D’, magenta arrowheads).

C4 nuclear actin is also in the oocyte nucleus, that is, germinal vesicle. During early oogenesis, very high levels of C4 nuclear actin are observed throughout the nuclear-plasm of the oocyte (Fig. 4A–A”,C–D”, orange arrowheads). This labeling is weaker in the later stages (Fig. 5, orange arrowheads). Notably, the nuclear actin within...
the oocyte is largely excluded from the chromatin (Fig. 4C–C" and see Fig. 7H–I", orange dashed circles).

C4 nuclear actin is also observed as a single spot within a subset of the somatic cells during early oogenesis. Specifically, in the germarium, high C4 nuclear actin levels are seen in the escort cells in region 1, but not those in region 2a (Fig. 4B–B", blue arrowheads). C4 nuclear actin is also observed in the FSCs at region 2a/2b boundary (Fig. 4B–B", blue dashed ovals). The findings that all the stem cell populations in the Drosophila ovary have high C4 nuclear actin suggests that nuclear actin may play a role in maintaining stemness. Subsequently, C4 nuclear actin is seen in a subset of the follicle cells from region 3 (Fig. 4B–B", blue arrows) through S6 (Fig. 4C,C", blue arrows). The frequency of follicle cells with C4 nuclear actin is high through S5, and only a few cells are labeled in S6 (Fig. 4C,C", blue arrows). No follicle cells are labeled in the later stages (Figs. 4D–D" and 5).

Fig. 3. DNase I labels nuclear actin during late stage follicle development. (A–D') Maximum projections of 2–4 slices of confocal stacks of the indicated stages of wild-type (yw) follicles. (A–D) Magenta = DAPI, Green = DNase I, (A'–D') White = DNase I. Insets are a zoom in of a different focal plane of the same follicle revealing the localization of DNase I within the follicle cells. While all the cells exhibit nuclear G-actin, marks are used to point out specific cells in which the DNase I-labeled structure is readily apparent in the focal planes shown. During the late stages of follicle development, DNase I staining in the nurse cells (yellow arrows) goes from tubular (S10B, A–A') to diffuse (S11–13, B–D'). Conversely, DNase I remains as a single spot within the follicle cells (blue arrows and insets in A–D') throughout the end of oogenesis. Scale bars = 50 μm in (A–D') and 10 μm in insets in (A'–D').
Interestingly, the follicle cells from region 3 to S5 are mitotic, in S6 the cells transition from mitotic to endocycling, and then endocycle through S9 (Claycomb and Orr-Weaver, 2005). Thus, the developmental pattern of C4 nuclear actin within the follicle cells suggests it may regulate or be regulated by the cell cycle. Additionally, like the DNase I labeled nuclear G-actin, the single puncta within all the somatic cells does not colocalize with the DNA.

**C4 Nuclear Actin Localizes to the Spindle during Mitosis**

To examine the relationship of C4 nuclear actin within the follicle cells and the cell cycle, we assessed C4 localization during the different stages of the cell cycle. In interphase cells, C4 nuclear actin labels a single spot in a subset of the cells (Fig. 6A–A’, blue arrows). This subnuclear localization pattern changes with the cell cycle. During early metaphase, C4 nuclear actin forms a double bar structure, surrounding the forming metaphase plate (Fig. 6B–C’). Later in metaphase, C4 nuclear actin labels the spindle (Fig. 6D–E’); notably, this nuclear actin appears filamentous. During anaphase, C4 nuclear actin initially forms a ring around each group of segregating chromosomes (Fig. 6F–F’), and later becomes restricted to the region between the chromosomes (Fig. 6G–G’). In early telophase, C4 nuclear actin is seen stretched between the chromosomes (Fig. 6H–H’), while in late telophase only a single vertical bar with low levels of C4 nuclear actin is observed (Fig. 6I–I’). These data suggest that C4 nuclear actin may regulate chromosome segregation during mitosis.

**Some C4 Nuclear Actin Colocalizes with DNase I**

As the C4 antibody is reported to recognize both monomeric and polymeric actin, we next assessed the structure of C4 nuclear actin by examining its localization in relation to DNase I, which recognizes monomeric or G-actin (Hitchcock, 1980). Interestingly, during oogenesis, C4 nuclear actin significantly colocalizes with DNase I in many cells, but is distinct in others (Fig. 7). Specifically, in the region 1 of the germarium, C4 nuclear actin within the GSCs (yellow dashed circles) and early cysts (white arrows) overlaps, but extends significantly beyond the DNase I foci (Fig. 7A–A’,C–C’). This finding suggests that C4 recognizes both monomeric and polymeric actin in the early, undifferentiated germ line cells. Note that we define polymeric nuclear actin as nuclear actin that does not co-label with DNase I and also fails to robustly label with phalloidin, a marker of F-actin; this suggests that the polymeric actin may be short filaments or have an altered structure compared to canonical F-actin. In the nurse cells during S3–10, C4 nuclear actin colocalizes with DNase I (Fig. 7B–B’,D–D’, yellow arrows). It is important to note that C4 nuclear actin is only in some of, while DNase I labels the nuclear, tubular structure in all the nurse cells. This finding suggests that C4 nuclear actin labels a subset of the G-actin in the nurse cells.

The localization of C4 nuclear actin and DNase I changes with oocyte development. In the newly specified oocyte in region 3 of the germarium, C4 nuclear actin and DNase I colocalize (Fig. 7A–A’, orange arrowhead and H–H’, red arrowhead). This colocalization suggests C4 nuclear actin is monomeric in the early oocyte. However, in S4 through early S6, C4 nuclear actin is high throughout the oocyte nucleoplasm and is enriched in a sphere encasing, but not overlapping the single DNase I puncta (Fig. 7J–J’), red arrowhead). In late S6 onward, C4 nuclear actin is throughout the nucleoplasm while DNase I appears as a weak haze (Fig. 7d–j” and data not shown). These data indicate that C4 nuclear actin is likely polymeric within the oocyte nucleus from S4 onward.

C4 nuclear actin largely colocalizes with DNase I in interphase somatic cells (Fig. 7, blue arrows, arrowheads, and dashed ovals). Specifically, C4 nuclear actin and DNase I label a single spot within the anterior escort cells in region 1 of the germarium (Fig. 7A–A’,K–K’, blue arrowheads) and the FSCs (Fig. 7A–A’,L–L’, blue dashed oval); note that the C4 nuclear actin labeling extends slightly beyond that of DNase I. When C4 nuclear actin is observed in the interphase follicle cells from region 3 of the germarium through S6, it largely colocalizes with DNase I (Fig. 7A–B’,D–D’,M–M’, blue arrowheads). These data suggest that C4 nuclear actin labels a subset of the G-actin within interphase somatic cells. In dividing cells, C4 nuclear actin is dynamic (see Fig. 6), while DNase I is uniform throughout the nucleoplasm but is excluded from the chromatin region (Fig. 7N–O”, white dashed circles). This finding indicates that the C4 nuclear actin observed from metaphase to telophase is likely polymeric.

**G-Actin Localizes to the Nucleolus**

We next wanted to determine where the monomeric or G-actin that is labeled by DNase I and, in some cases, C4, is localizing within the nucleus. As the G-actin was excluded from the DNA, appeared as a single spot in the early germ cells and the somatic cells, and exhibited a tubular appearance within the nurse cells, we...
hypothesized that the G-actin was within the nucleolus. To test this hypothesis, we compared the localization of C4 nuclear actin to eGFP-tagged nucleolar components, fibrillarin (Fig. 8) and Nopp-140 (data not shown). In the GSCs and cystoblasts (region 1 of germarium), the nucleolus is a single spot; C4 nuclear actin is enriched in the nucleolus, but extends throughout the nucleoplasm (Fig. 8A–A', yellow dashed circles, and C–C', red arrowheads mark the nucleolus). Conversely, in the nurse cells, C4 nuclear actin completely colocalizes with the nucleolus (Fig. 8A–B' and D–E', yellow arrows). In the oocyte, C4 nuclear actin is initially enriched in the nucleolus (Fig. 8F–F', red arrowhead). In S4 through early S6, C4 nuclear actin is throughout the nucleoplasm, and is also enriched in the immediate vicinity of the nucleolus (S4 Fig. 8G–G', and early S6, B–B', red arrowheads). Starting in late S6, no strong nucleolar signal is observed in the oocyte nucleus and C4 nuclear actin is throughout the nucleoplasm (Fig. 8H–H').

In the somatic cells during *Drosophila* oogenesis, C4 nuclear actin largely localizes to the nucleolus (Fig. 8, blue arrowheads, arrows and dashed ovals). In region 1 of

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**Fig. 5.** C4 nuclear actin is significantly reduced and relocates during late oogenesis. (A–D') Maximum projections of 2–4 slices of confocal stacks of the indicated stages of wild-type (yw) follicles. (A–D) Magenta = DAPI, Green = C4. (A'–D') White = C4. During the late stages of follicle development, C4 nuclear actin goes from a tubular structure observed at a low frequency in S10B (A–A', yellow arrows), to localizing to a ring just underneath the nuclear envelope in S11–12 (B–C', magenta arrows). Finally, during S13, C4 labels puncta that appear to be in either the cytoplasm of the nurse cells or the stretch follicle cells enveloping them (D–D', magenta arrowheads). C4 nuclear actin is observed throughout the nucleoplasm of the oocyte (A–D', orange arrowheads). Note that not all cells with C4 nuclear actin are marked; marks are used to point out specific cells in which the C4 nuclear actin is readily apparent in the focal planes shown. Scale bars = 50 μm.
the germarium, C4 nuclear actin within the escort cells overlaps and extends beyond the nucleolus to form a larger spot (Fig. 8A–A”, blue arrowheads). Within the FSCs, C4 nuclear actin completely localizes to the nucleolus (Fig. 8A–A”, blue dashed ovals). When C4 nuclear actin is observed in the follicle cells from region 3 of the germarium to S6, it localizes to and extends slightly beyond the nucleolus (Fig. 8A–A”, blue arrows).

Similar to monomeric C4 nuclear actin, DNase I-labeled G-actin localizes to the nucleolus as visualized using either eGFP-tagged nucleolar components or an antibody to fibrillarin (data not shown). These data, in conjunction with the colocalization data between C4 nuclear actin and DNase I, suggest that monomeric nuclear actin is largely found within the nucleolus in both somatic and germline cells.

One Actin Antibody (AC15) Recognizes a Pool of Nuclear Actin during Mid and Late Oogenesis

The AC15 actin antibody specifically recognizes β-, but not α- or γ-actin (Gimona et al., 1994). It was raised...
against a modified N-terminal peptide from β-actin (amino acids 2–16) conjugated to a carrier protein (Keyhole Limpet Hemocyanin). This peptide is 100% similar to the N-termini of the two ubiquitous Drosophila actins, Actin 5C and 42A. Notably, these two actins are highly expressed during oogenesis (ModENCODE; Tootle et al., 2011), and we previously found that expression of GFP-Actin 5C or 42A within the germline results in
nuclear actin rod formation (Kelsch et al., 2016). The AC15 antibody has been widely used to examine nuclear actin, including in plants (Cruz and Moreno Diaz de la Espina, 2009), Drosophila cultured cells (Dopie et al., 2012), amphibian oocytes (Miyamoto et al., 2011), and mammalian cells (Hofmann et al., 2004). Thus, AC15 antibody is suitable for examining nuclear actin.

During oogenesis, anti-actin AC15 weakly labels the cytoplasm and recognizes a distinct pool of nuclear actin from mid-oogenesis onward, subsequently referred to as AC15 nuclear actin (Figs. 9 and 10). During early oogenesis (gerarium through S5), there are very low levels of AC15 nuclear actin (Fig. 9A–A'). At S6, weak AC15 nuclear actin is observed in the nurse and follicle cells (Fig. 9A–A'), and the level of AC15 nuclear actin in these cells continues to increase through S10 (Fig. 9A–C'). The AC15 nuclear actin in the nurse cells is structured, with enrichment at the nuclear periphery (Fig. 9B–C). This peripheral localization increases with follicle development (S10B–12, Fig. 9D–F'). At S13, AC15 nuclear actin is largely absent and AC15 labels a cytoplasmic cytoskeletal structure (Fig. 9G–G'). In the follicle cells, AC15 nuclear actin is observed in the early migrating border cells during S9 (Fig. 9B–B', white dashed oval), the main body follicle cells from S6 onward (Fig. 9A–C',D'–G'), and the stretch follicle cells over the nurse cells (Fig. 9D'–G'). Notably, the nuclear actin structures recognized by AC15 in both the nurse and follicle cells is distinct from that of DNase I and C4, as it does not colocalize with DNase I (data not shown). This finding suggests that AC15 nuclear actin is likely polymeric. As phalloidin does not label the AC15 nuclear actin, we speculate that the antibody recognizes short polymers of actin and not canonical F-actin. Based on the distinct localization pattern, we next examined where AC15 nuclear actin is compared to the DNA. In the nurse cells, AC15 nuclear actin is enriched on the chromatin (Fig. 10A–D'), but also labels puncta in regions that are devoid of DNA (yellow arrowheads).

In the oocyte, AC15 nuclear actin changes with development. In S3–4, AC15 nuclear actin exhibits a speckled pattern throughout the oocyte nucleoplasm, with the brightest spots at the edge of the chromatin (Fig. 10E–E'). As the oocyte develops (S6–8), AC15 nuclear actin becomes restricted to puncta adjacent to the DNA (Fig. 10F–G'). These puncta appear to transition to filamentous structures that surround the DNA starting at S9 and continuing through S11 (Fig. 10H–I' and data not shown). Finally, in S12–13, AC15 nuclear actin is more diffuse but still encircles the chromatin (Fig. 10J–J').

AC15 nuclear actin within the follicle cells is also on the DNA. Interestingly, AC15 nuclear actin is dynamic in the border cells during their migration. Early in border cell migration, AC15 nuclear actin is weakly observed in all the cells of the cluster (Fig. 9B–B', white dashed oval and Fig. 10K–K', blue arrows). Later during both the posterior (Fig. 10L–L') and dorsal migration (Fig. 10M–M') of the border cells, AC15 nuclear actin labels one to two cells. Given the location of the labeled cells during the late stages of migration, we speculate that these are the polar cells. Compared to the border cells, the level of AC15 nuclear actin is significantly higher in the other follicle cells (see Fig. 10N–P'). In the main body follicle cells, AC15 nuclear actin exhibits a speckled appearance and the speckles localize to the DNA (Fig. 10N–N'). AC15 nuclear actin is also enriched in the centripetal follicle cells, compared to the adjacent main body follicle cells (Fig. 10O–O', blue dashed oval compared to blue arrows), as they begin their migration between the nurse cells and oocyte. In the stretch follicle cells, AC15 nuclear actin also localizes to the chromatin (Fig. 10P–P').

In summary, AC15 nuclear actin generally localizes to the chromatin during mid to late oogenesis in both the germline and somatic cells. These findings lead us to speculate that this pool of nuclear actin may play a role in transcription.

**DISCUSSION**

Using Drosophila oogenesis as a model to study nuclear actin reveals that multiple pools of nuclear actin exist and provides insight into the potential functions of nuclear actin (Table 1). Thus, while all statements made about the functions of nuclear actin during Drosophila oogenesis are based on both our findings and prior studies, they are speculative and remain to be tested. We find that not all cells contain G-actin, recognized by DNase I, in a...
Nuclear Actin and Stemness

Our findings suggest nuclear actin may play a critical role in maintaining an undifferentiated state. Specifically, high levels of C4 nuclear actin are observed in the GSCs and the undifferentiated cystoblasts, but not in the later cysts within the germarium. This C4 nuclear actin is enriched at and overlaps the small DNase I spot that localizes to the nucleolus, but also extends throughout the nucleoplasm. These data lead us to hypothesize that polymeric C4 nuclear actin regulates germline differentiation (Table 1). Supporting this idea, nuclear actin has been implicated in controlling both cell quiescence, transcription, nuclear structure/function, cell cycle, and nuclear structure.

Nuclear Actin and Transcription

The best characterized function of nuclear actin across all systems is its role in mediating transcription. Indeed, nuclear actin plays a general role within every cell. Intriguingly, one actin antibody, C4, exhibits a more restricted pattern during early oogenesis, but when it is present, it often colocalizes with DNase I. This finding supports the model that C4 nuclear actin recognizes a specific subset or pool of G-actin. This G-actin may be post-translationally modified or in a particular protein complex that results in the antigen on being available. Future proteomic studies are needed to confirm these possibilities. C4 nuclear actin also colocalizes with the FSCs, C4 nuclear actin that colocalizes with DNase I is not monomeric but polymeric actin at the same subnuclear localization. Indeed, polymeric C4 nuclear actin, defined by the lack of colocalization with DNase I, is also observed in specific cells, including the GSCs and oocytes. Such findings support the idea that the structure of nuclear actin impacts its functions. Interestingly, another actin antibody, AC15, labels a distinct pool of polymeric actin. AC15 nuclear actin is observed from mid-to-late oogenesis in almost every cell, and does not colocalize with DNase I. These findings suggest that AC15 nuclear actin is polymeric and mediates completely different processes from DNase I-labeled and C4 nuclear actin. The cell-type, developmental, and subnuclear pattern of the different pools of nuclear actin leads us to speculate that nuclear actin contributes to stemness, transcription, nuclear structure/function, cell cycle, and nuclear structure.
actin binds to and regulates all three RNA polymerases (RNAP) (Smith et al., 1979; Fomproix and Percipalle, 2004; Hofmann et al., 2004; Hu et al., 2004; Philimonenko et al., 2004). Nuclear actin promotes transcription as it was purified as an activator of transcription (Egly et al., 1984), and microinjection of antibodies to actin and actin binding proteins inhibits transcription (Scheer et al., 1984). More recent studies have shown that actin is required for RNAPI activity (Fomproix and Percipalle, 2004; Philimonenko et al., 2004; Almuzzaini et al., 2016) and transcriptional initiation and elongation by RNAPII (Hofmann et al., 2004). This actin may be polymeric as mutations in actin and drugs that prevent actin polymerization block in vitro transcription by RNAPI (Ye et al., 2008), and nuclear myosin I is a well-established regulator of transcription (Hofmann et al., 2004; Philimonenko et al., 2004, 2010; Ye et al., 2008). However, monomeric actin may also contribute to transcription, as monomers mediate chromatin remodeling complex recruitment and formation (reviewed in Grosse and Vartiainen, 2013), and sequestering monomeric nuclear actin by inducing nuclear actin rod formation inhibits transcription.
Fig. 10. AC15 nuclear actin localizes to the DNA. (A–P") Zoomed in images of different cell types from maximum projections of 2–4 slices of confocal stacks of the indicated stages of wild-type (yw) follicles. (A–D, K–P) Magenta = DAPI, Green = AC15. (E–J) Magenta = DAPI, Green = AC15, and White = Wheat germ agglutinin (WGA, marks nuclear envelope). (A’–P’) White = AC15. (A’–P”) White = DAPI. (A–D”) Nurse cells. (E’–J”) Oocytes. (K–M”) Border cells. (N–N”) Main body follicle cells. (O–O”) Centripetal follicle cells. (P–P”) Stretch follicle cells. Within the nurse cells, AC15 largely colocalizes with the DNA (A–D”), however, it also labels puncta within regions devoid of chromatin in S9–11 (yellow arrowheads). In the oocyte (nuclei circled by orange dashed line), AC15 nuclear actin structure changes with development (E–J). In S4 oocytes, AC15 nuclear actin exhibits a speckled appearance throughout the nucleoplasm, with bright puncta adjacent around the edge of the chromatin (E–E’). This enrichment of AC15 puncta around the DNA in the oocyte increases in S4–6 (F–G’), and then appears to form filaments that encircle the chromatin in S9–11 (H–I’). In S12 oocytes, AC15 nuclear actin becomes more diffuse, but still surrounds the DNA (J–J”). AC15 nuclear actin is also observed in the somatic cells. In the migrating border cells, AC15 nuclear actin is weakly observed in all of the cells (both border and polar cells) in the early stage of migration (K–K”), blue arrows). Toward the end of the border cell migration (late S9 and S10A), AC15 nuclear actin appears restricted to a subset of the cells (L–M”, blue arrows); given the localization of the AC15 positive nuclei, we hypothesize that they are the polar cells. AC15 nuclear actin is observed in all of the other follicle cell populations, including the main body follicle cells (N–N” and blue arrows in O–O”) and the stretch follicle cells (P–P”, blue arrows). Scale bars = 10 μm.
Thus, it remains unclear what form or forms of nuclear actin contribute to transcription. Our findings suggest that during *Drosophila* oogenesis multiple pools of polymeric actin may regulate RNAPII transcription (Table 1). Indeed, polymeric C4 nuclear actin overlaps with the DNA in the GSCs and early cystoblasts. This finding leads us to speculate that this polymeric C4 nuclear actin mediates RNAPII transcription to drive the transcriptional program necessary for maintaining an undifferentiated state. Additionally, polymeric AC15 nuclear actin localizes to the chromatin within the nurse cells and follicle cells from mid-to-late oogenesis. During this period of development, the nurse cells massively upregulate transcription to supply the oocyte with mRNA, protein, and organelles necessary for it to complete embryogenesis (Spradling, 1993). The follicle cells undergo gene amplification and express high levels of genes encoding the eggshell (Claycomb and Orr-Weaver, 2005). Thus, we hypothesize that the chromatin localized AC15 nuclear actin mediates the upregulation of transcription in these cells. These hypotheses can be explored by defining the transcriptional changes occurring when nuclear actin levels or dynamics (i.e., actin rod formation) are perturbed.

Nuclear actin also regulates specific transcription factors. For example, myocardin-related transcription factor A (also referred to as megakaryocytic acute leukemia (MAL)/serum response factor (SRF) localization and activity is negatively regulated by G-actin in the cytoplasm and the nucleus (Vartiainen et al., 2007). MAL/ SRF transcription plays important roles in cell migration, including during cancer metastasis (Medjkane et al., 2009) and *Drosophila* border cell migration (Somogyi and Rorth, 2004). Interestingly, during early border cell migration, AC15 nuclear actin is observed in all cells within the cluster, both border and polar cells. But later in migration, and when the cluster has reached the oocyte, only one to two cells exhibit AC15 nuclear actin; we presume these are the polar cells. These findings lead us to hypothesize that AC15 nuclear actin may regulate MAL/SRF activity to control the collective and invasive migration of the border cells (Table 1). As these cells are a good model for cancer metastasis (Montell, 2003), examination of the prevalence and level of nuclear actin in the different stages of cancer is warranted. Additionally, we find that AC15 nuclear actin levels are higher in the migrating centripetal follicle cells compared to the neighboring main body follicle cells. Thus, nuclear actin may play a more general role in mediating cell migration (Table 1). Indeed, the level of

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It indicates the cell-types where C4 nuclear actin localizes to the nucleolus, but it also extends slightly beyond it.
nuclear actin must be tightly controlled to mediate the transcriptional program necessary for cell motility in keratinocytes (Sharill et al., 2016).

During *Drosophila* oogenesis, nuclear actin may also regulate RNAPI- and/or RNAPIII-dependent transcription (Table 1). The primary site of RNAPI and RNAPIII activity is the nucleolus, where they mediate the transcription of ribosomal genes. G-actin, recognized by DNase I, is observed in the nucleolus of every cell throughout follicle development. This finding suggests monomeric actin may have a general role in nucleolar function, such as ribosomal gene expression (Table 1). Some C4 nuclear actin also localizes to the nucleolus. Specifically, every GSC, early cystoblast, and FSC has nucleolar C4 nuclear actin. Interestingly, this is the only C4 nuclear actin observed in the FSCs. This finding suggests that in undifferentiated cells, including stem cells, the protein production needs are higher (Table 1). Indeed, the mouse β-actin knockout studies implicate nuclear actin’s role in RNAPI activity and ribosomal RNA production as a key factor in cell growth and proliferation (Almuzzaini et al., 2015), properties that are particularly important for the rapidly dividing *Drosophila* GSCs and FSCs. C4 nuclear actin is also observed in the nucleoli of a subset of mitotic follicle cells and the nurse cells during S3–10B. We speculate that this nuclear actin might reflect that the cells are in a period where higher RNAPI and RNAPIII activity is needed (Table 1). While AC15 nuclear actin is primarily on the chromatin in the nurse cells, small puncta are observed within the areas devoid of DNA that, based on our prior work (Groen et al., 2015), are likely in the nucleolus. Interestingly, these nuclear AC15 puncta are most frequently observed during S9–10B. During this period of development, the nurse cells are producing vast quantities of ribosomes, and proteins to be transported into the oocyte. Additionally, the puncta strikingly resemble what is seen for nuclear myosin I, which is required for RNAPI transcription (Fumproix and Percipalle, 2004). Thus, we hypothesize that polymeric AC15 nuclear actin interacts with a nuclear myosin to drive RNAPI transcription in the nurse cells during this critical period of development (Table 1). In the future, it will be important to define the specific roles of the different nuclear actin pools in RNAPI and RNAPIII activity.

**Nuclear Actin and the Nucleolus**

Nuclear actin may play other roles in the nucleolus. Actin has been observed in the nucleolus in numerous species, including plants (Cruz and Moreno Diaz de la Espina, 2009), slime mold (Jockusch et al., 1971), *Drosophila* (this study), and human cells (Belin et al., 2013). Additionally, numerous actin binding proteins are also found in the nucleolus (Hubert et al., 2008; Deng et al., 2012; Kitamura et al., 2015). While it is possible that nuclear actin and its regulators are simply sequestered in the nucleolus until they are needed for their nuclear functions, in a similar manner to cell cycle factors (Visintin and Amon, 2000; Boisvert et al., 2007), it is likely that these proteins also have roles in this nuclear body. Indeed, one of these proteins, ARP6, mediates both ribosomal gene expression and repression, depending upon the cellular environment (Kitamura et al., 2015). This finding leads us to speculate that the dynamic localization of C4 and AC15 nuclear actin to the nucleolus during *Drosophila* oogenesis may reflect periods of inhibiting and driving ribosome biogenesis. ARP6 also regulates the structure of the nucleolus, as knockdown in HeLA cells results in striking morphological changes in the nucleolus (Kitamura et al., 2015). We find this particularly intriguing, as the actin bundling protein Fascin localizes to the nucleus, regulates nucleolar structure (Groen et al., 2015), and positively influences C4 nuclear actin (Kelsch et al., 2018). In *Drosophila*, Fascin localization and function are regulated by lipid signals termed prostaglandins (Groen et al., 2012, 2015). Together these data lead us to hypothesize that prostaglandins tightly control the levels of nuclear Fascin to modulate nuclear actin to ultimately regulate nucleolar structure. Supporting this, preliminary studies indicate that loss of prostaglandin synthesis, which causes an aberrant nucleolar structure (Groen et al., 2015), increases C4 nuclear actin within the nurse cells (Wineland, Kelsch, and Tootle; unpublished observation).

The potential connection between Fascin, prostaglandins, nuclear actin, and nucleolar structure has interesting implications in the context of cancer, as the nucleolus is a new target for anticancer therapies (reviewed in Ruggiero, 2012; Hein et al., 2013; Quin et al., 2014). Indeed, pathologists diagnose cancer by alterations in nucleolar morphology (reviewed in Ruggiero, 2012; Hein et al., 2013; Quin et al., 2014). The changes in the number and structure of nucleoli may affect cell proliferation, differentiation, senescence, apoptosis, and coordination of response to stress. Additionally, the severity the nucleolar morphological changes is used as a marker of cancer aggressiveness (reviewed in Ruggiero, 2012; Hein et al., 2013; Quin et al., 2014). Notably, increased Fascin expression (Hashimoto et al., 2004; Yoder et al., 2005; Lee et al., 2007; Okada et al., 2007; Li et al., 2008; Chan et al., 2010) and prostaglandin production (Rolland et al., 1980; Chen et al., 2001; Khuri et al., 2001; Denkert et al., 2003) contribute to tumorigenesis and metastasis, and are associated with poor patient prognosis. Finally, recent studies indicate cancer cells have misregulated and higher nuclear actin levels than the normal epithelium (Fiore et al., 2017). Additionally, many actin binding proteins exhibit increased nuclear localization during tumorigenesis and progression (Yang and Lin, 2018). However, the nucleolar versus other nuclear function of actin in cancer development and progression remain to be elucidated.

**Nuclear Actin and Mitosis**

Nuclear actin also plays roles in the cell cycle and chromosome segregation. While studies have implicated nuclear actin in mitosis (Jockusch et al., 1971), and even observed it localizing in the vicinity of the mitotic spindle (Woolner et al., 2008; Hubert et al., 2011), clear evidence of its presence and function have remained elusive. Here, we find that during mitosis in the follicle cells, C4 nuclear actin is dynamic, forming a double bar around the forming metaphase plate, localizing to the spindle during metaphase, rearranging to ultimately be restricted to the area between the segregating chromosomes in anaphase, and this structure elongating and thinning as the segregation continues in telophase (Table 1). Notably, DNase I is diffuse throughout the extrachromosomal nucleoplasm, indicating that the C4 nuclear actin is likely polymeric.
Interestingly, the *Drosophila* protein EAST is known to bind to nuclear actin (Wasser and Chia, 2000) and exhibits a similar, dynamic localization during mitosis in the embryo (Wasser and Chia, 2000). As EAST is highly expressed during oogenesis, it is tempting to speculate it may regulate nuclear actin localization or function during follicle cell mitosis (Brown et al., 2014). Further studies are needed to determine the role(s) of nuclear actin in cell cycle progression and chromosome segregation.

While actin has not been previously shown to be on the spindle or regulate chromosome segregation in mitosis, it has been widely implicated in these processes during meiosis. Indeed, over 30 years ago, actin was observed on the meiotic spindles in plants (Forer et al., 1979; Forer and Jackson, 1979). Subsequently, it was observed on the meiotic spindles of insects (Silverman-Gavrila and Forer, 2000; Fabian and Forer, 2007) and mice (Bogolyubova and Ginzburg, 2013). A recent study extended these findings from observational to functional, showing that in mammalian oocytes, including humans, spindle actin is required to align and segregate the chromosomes (Mogessie and Schuh, 2017). These studies, in conjunction with our findings, lead us to hypothesize that nuclear actin similarly regulates chromosome dynamics during meiosis (Table 1).

**Nuclear Actin and Nuclear Structure**

Although we did not examine whether nuclear actin regulates meiosis, our data suggest that nuclear actin may play a structural role in the developing *Drosophila* oocyte. We find that high levels of C4 nuclear actin are present in the nucleoplasm of the oocyte (Table 1). Given that this nuclear actin does not overlap with DNase I staining, we hypothesize this is polymeric actin. Supporting that polymeric or filamentous actin is important for nuclear structure within the oocyte, numerous studies have shown the presence of a filamentous nuclear actin network within the germinal vesicles of organisms with large oocytes. This network is thought to mediate nuclear organization, including chromosome and nuclear body dispersal (Clark and Merriam, 1977; Parfenov et al., 1995; Gonsior et al., 1999; Kiseleva et al., 2004; Maslova and Krasikova, 2012). For example, *Xenopus* oocytes accumulate large amounts of nuclear actin because they do not express the specific export factor, Exportin 6 (Stuven et al., 2003), that translocates actin from the nucleus. Ectopic expression of Exportin 6 results in the loss of the intranuclear F-actin scaffold and a fragile nucleus (Bohsack et al., 2006). This nuclear F-actin scaffold forms a weak viscoelastic network that helps to stabilize the intranuclear localization of the chromatin and nuclear bodies against gravitational forces that affect large cells (Feric and Brangwynne, 2013; Feric et al., 2015). Indeed, if the nuclear actin network is disrupted, the nuclear bodies sediment and fuse (Feric and Brangwynne, 2013). These data lead us to speculate that polymeric C4 nuclear actin within the oocyte nucleus plays a similar role (Table 1).

In addition to C4 nuclear actin, AC15 nuclear actin is also found in the oocyte. During S3–8, AC15 nuclear actin is seen as puncta that are enriched around the chromosomes. While during S9–11, filamentous structures are observed surrounding the DNA, and in S12–13, AC15 nuclear actin forms a more diffuse cage around the chromosomes. Given the association of AC15 nuclear actin with the chromatin in the other cells during *Drosophila* oogenesis, the well-established role of nuclear actin in transcription (reviewed in Venit et al., 2018), and the finding that inducing stable nuclear actin filaments blocks transcription (Serebryannyy et al., 2016b), we hypothesize that in the oocyte, AC15 nuclear actin is sequestered into filaments to block transcription (Table 1). These filaments can then be depolymerized during embryogenesis to allow for transcription to resume. If this hypothesis is correct, then altering nuclear actin level or dynamics will have severe effects on embryonic development. Supporting this idea, mutations in the gene encoding the importin (Importin 9) responsible for taking actin into the nucleus results in lethality in *Drosophila* (Kelpsich, Jaime, and Tootle, unpublished observation) and mice (Blake et al., 2017). Future studies are needed to determine the cause of the lethality.

**CONCLUSIONS**

This study, in conjunction with the last 50 years of research on nuclear actin, strongly supports the ideas that the prevalence of nuclear actin is widespread, and actin within the nucleus likely has a variety of functions that are impacted by its structure and binding partners. Our data lead us to speculate that nuclear actin regulates transcription, nucleolar structure and function, cell cycle, and nuclear structure (Table 1). Such activities are important for regulating cell fate, growth, proliferation, and function. Thus, nuclear actin is predicted to have widespread roles in mediating development, and its misregulation likely contributes to diseases. Future studies are needed to determine nuclear actin prevalence and structure across tissues and organisms. Additional mechanistic studies are required to elucidate the activities of the different forms or pools of nuclear actin.

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**LITERATURE CITED**


MULTIPLE POOLS OF NUCLEAR ACTIN


