Morphogenesis of *Drosophila* ovarian ring canals

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**SUMMARY**

We analyzed the structure of cytoplasmic bridges called ring canals in *Drosophila* egg chambers. Two mutations, *hu-li tai shao* (*hts*) and *kelch*, disrupt normal ring canal development. We raised antibodies against the carboxy-terminal tail of hts and found that they recognize a protein that localizes specifically to ring canals very early in ring canal assembly. Accumulation of filamentous actin on ring canals coincides with the appearance of the hts protein. *kelch*, which is localized to the ring canal hours after hts and actin, is necessary for maintaining a highly ordered ring canal rim since *kelch* mutant egg chambers have ring canals that are obstructed by disordered actin and hts. Anti-phosphotyrosine antibodies immunostain ring canals beginning early in the gerarium before hts and actin and throughout egg chamber development. The use of antibody reagents to analyze the structure of wild-type and mutant ring canals has shown that ring canal development is a dynamic process of cytoskeletal protein assembly, possibly regulated by tyrosine phosphorylation of some ring canal components.

Key words: *Drosophila*, oogenesis, ring canals, *kelch*, *hu-li tai shao*

**INTRODUCTION**

Cells use a number of mechanisms to communicate with one another during development. These mechanisms include secretion of diffusible ligands, secretion of extracellular matrix, and the formation of adherens and gap junctions upon contact with each other. During gametogenesis of several vertebrate and invertebrate organisms, an unusual type of intercellular communication is used. As a result of incomplete cytokinesis during mitotic divisions, gametes of males and females develop as syncytia connected by large cytoplasmic bridges called ring canals (Burgos and Fawcett, 1955; Brown and King, 1964). In mammals, syncytial development may be required to synchronize mitotic divisions and entry into meiosis and possibly to synchronize atresia or selective degeneration of oocytes (Franchi and Mandl, 1962; Weakley, 1967; Zamboni and Gondos, 1968; Dyer et al., 1968; reviewed by Gondos, 1973). During oogenesis of many insects, the cytoplasmic bridges allow the flow of nutrients between cells in a syncytium. Only one cell develops into a mature oocyte while the other cells retract and die after contributing their cytoplasmic contents to the oocyte (King and Mills, 1962; Mahowald and Kambsellis, 1980; Anderson and Huebner, 1968). In *Drosophila* oogenesis, cytoplasmic transport between nurse cells and the oocyte is essential for the development of a normal oocyte (Mahowald and Kambsellis, 1980; Spradling, 1993 for reviews of *Drosophila* oogenesis). The cytoskeleton plays an integral role in cytoplasm transport because disruption of the cytoskeleton by mutation (Storto and King, 1988; Cant et al., 1994; Cooley et al., 1992) or by pharmacological agents (Gutzeit, 1986a,b; Theurkauf et al., 1993) causes defective transport. Cytoplasm transport also depends on a class of specialized cytoskeletal proteins that are required for development of a functional ring canal during *Drosophila* oogenesis.

The *Drosophila* ovary consists of 15-18 discrete tubular ovarioles (Fig. 1A). The germline stem cells reside at the anterior of each ovariole in the gerarium (Fig. 1B). In region 1 of the gerarium, stem cell daughters undergo four rounds of mitosis with incomplete cytokinesis. This gives rise to 16-cell clusters in which two cells have 4 intercellular bridges, two cells have three, four cells have two, and eight cells have one. One of the 16 cells differentiates into the oocyte while the other fifteen differentiate into the nurse cells (Fig. 1C). In gerarium region 2a, newly formed 16-cell clusters are enclosed by migrating follicle cells forming the egg chamber, and in region 2b the egg chambers become lens-shaped. Region 3 contains stage 1 spheroidal egg chambers that are ready to enter the vitellarium for maturation (Brown and King, 1964; Koch and King, 1966; Mahowald and Strassheim, 1970). In the vitellarium, progressively more mature egg chambers extend from the anterior to the posterior end of the ovariole. These egg chambers have been divided into stages 2-14 where a stage 14 oocyte is ready to be fertilized and oviposited.

Cleavage furrow arrest and ring canal development are not well understood. Early studies using electron microscopy revealed a highly vesiculated structure called fusome extending through the cytoplasmic bridges, possibly functioning to stabilize the cleavage furrow until a ring canal is established (Meyer, 1961; Koch and King, 1966; King and Storto, 1988; Lin et al., 1994). Similar studies also revealed that when a ring canal is formed, two components are added sequentially. First, an electron opaque backbone that is a part of or adjacent to the plasma membrane is added upon arrest of the cleavage furrow. Second, an electron dense inner rim accumulates after the final mitotic division of each germline syncytia (Fig. 1D;
The inner rim is established coordinately with the degeneration of the fusome (Koch and King, 1969; Lin et al., 1994) and the addition of actin to the ring canal (Warn et al., 1985; Theurkauf et al., 1993).

The genes for two mutations, *hu-li tai shao* and *kelch*, which either disrupt ring canal structure or cytoplasm transport have been cloned. Mutants for *hu-li tai shao* (*hts*) (Yue and Spradling, 1992) produce degenerate ovaries that contain egg chambers with less than a full complement of germline cells. Frequently, an oocyte fails to differentiate in these mutant egg chambers. In addition, mutant egg chambers do not accumulate the inner rim of ring canals and have no actin on their ring canals (Yue and Spradling, 1992). The known *hts* cDNA predicts a protein that is 37% identical to adducin in the amino-terminal globular region, hinge region and the first part of the adducin tail domain. *hts* protein and adducin share no homology in the final 160 amino acids of adducin, and *hts* contains an additional 400 amino acids (carboxy-terminal tail) not found in adducin (Fig. 2; Yue and Spradling, 1992; Ding et al., 1993).

Mutants for *kelch* fail to complete cytoplasm transport from the nurse cells into the oocyte resulting in a female sterile phenotype. Cytoplasm transport is affected beginning early in the vitellarium where oocytes in mutant egg chambers are smaller than wild-type oocytes of the same stage (Xue and Cooley, 1993). The *kelch* gene was cloned and found to encode a protein specifically localized to ring canals (Xue and Cooley, 1993). The kelch protein contains two motifs found in several other proteins (Koonin et al., 1992). The first motif is a 100 amino acid domain called the BTB box (Godt et al., 1993) that is positioned approximately 150 amino acids from the amino terminus. The BTB box is found at the amino terminus of several *Drosophila* transcription factors including Broad-Complex (DiBello et al., 1991), Tramtrack (Harrison and Travers, 1990), and Adf2 (Benyajati, personal communication). Several open reading frames (ORFs) in a number of Pox viruses (Koonin et al., 1992; Senkevich et al., 1993) also contain the BTB box. The function of the BTB box is unknown. The second motif consists of six 50 amino acid repeats called kelch repeats. The Pox virus ORFs (Koonin et al., 1992; Senkevich et al., 1993) mouse MIPP (Chang-Yeh et al., 1991), and scrnin protein (Tilney, 1975) from *Limulus* contain several kelch repeats (Way and Matsudaira, personal communication).

We present studies on the formation of functional ring canals...
and ring canal structure in hts and kelch mutants. This work has relied on several antibody reagents including new antibodies raised against the carboxy-terminal tail of hts that specifically recognize ring canals and anti-phosphotyrosine antibodies that recognize one or more additional components of a ring canal. The proteins are added sequentially to the arrested cleavage furrow during ring canal ontogeny. These results have allowed us to formulate a model of ring canal structure and to speculate on the early events necessary to stabilize and transform an arrested cleavage furrow into a ring canal.

MATERIALS AND METHODS

Fly strains

w^{118} or cn; ry (Lindsay and Zimm, 1992) flies were used as wild type in these experiments. hts1 mutant flies have been described previously by Yue and Spradling (1992). kelneo mutant flies were described previously by Xue and Cooley (1993). Fly stocks were maintained under standard conditions.

Antibody production

Monoclonal antibodies to the carboxy-terminal tail of hts were generated using standard procedures. Recombinant fusion proteins were made using the pGEX system (Smith and Johnson, 1988). A BgIII-EcoRI fragment from hts5 (Yue and Spradling, 1992) encoding the carboxy-terminal 351 amino acids (Fig. 2) was cloned into pGEX-2T. Protein was produced as a fusion to the carboxy terminus of Schistosoma japonicum glutathione-S-transferase (Sj26) and purified from inclusion bodies. Purified protein was injected into mice and hybridoma cell lines were generated according to Harlow and Lane (1988). Antiserum and hybridoma cell line supernatants were screened for immunoreactivity to the fusion protein and to wild-type but not hts1 mutant ovaries.

Western analysis

Drosophila ovaries were dissected in Drosophila Ringer’s solution and ground in 1x Laemmli’s buffer under reducing conditions (Laemmli, 1970). Protein concentrations of the ovary extracts were measured using the BioRad Protein Assay. Proteins separated by SDS-polyacrylamide gel electrophoresis (Laemmli, 1970) were transferred to Hybond-ECL nitrocellulose (Towbin et al., 1979; Amersham). Following transfer, membranes were blocked in Blotto-Tween (5% powdered milk, 0.2% Tween, and PBS) for 2 hours, then incubated in hybridoma cell supernatant (1:10 in Blotto-Tween) for 1 hour to overnight. Membranes were washed four times for 10 minutes each in 0.2% Tween in PBS at room temperature. Membranes were incubated for 30 minutes at room temperature in goat anti-mouse IgG secondary antibody conjugated to horseshadish peroxidase (Pierce Chemical Co.) in Blotto-Tween (1:10,000). After four washes, signals were detected using ECL western blotting detection reagents (Amersham) according to manufacturer’s specifications.

Immunolocalization and confocal imaging

Whole ovaries were dissected in Drosophila Ringer’s, fixed in 1% formaldehyd saturated with heptane (Cooley et al., 1992), washed three times for 10 minutes in PBS and once for 10 minutes in PBT (0.3% Triton X-100 Sigma), 0.5% BSA (Sigma) in PBS) at room temperature. For actin visualization, ovaries were incubated in 2-4 units of rhodamine-conjugated phalloidin (Molecular Bioprobes) for 20 minutes at room temperature. For antibody staining, ovaries were incubated in hts carboxy-terminal antibody hybridoma supernatant (1:1), kelch hybridoma supernatant (Xue and Cooley, 1993) (1:1), or anti-phosphotyrosine antibody (UBI, 4G10 or ICN Biochemicals, P320) (1:1000) diluted in PBT. The ovaries were incubated with rocking for 2 hours at room temperature then overnight at 4°C. After washing four times for 15 minutes in PBT, bound antibodies were detected using a goat anti-mouse IgG secondary antibody conjugated to fluorescein isothiocyanate (Jackson Laboratories). Immunolocalizations were visualized by collecting 1-2 µm optical sections on a laser-scanning confocal microscope (BioRad MRC 600) and counterstaining and displaying optical sections using the CoMOS software package (BioRad). Either a 50× (0.95 NA) or a 100× (1.30 NA) objective was used in each case.

RESULTS

Temporal order of hts, actin, and kelch addition to ring canals

To study the role of the hts gene in the development of Drosophila egg chambers, we generated mouse monoclonal antibodies against the carboxy-terminal 351 amino acids (carboxy-terminal domain) of hts (Fig. 2). By western analysis, the monoclonal antibodies recognized a doublet of proteins with apparent relative molecular masses of approximately 60x10^3 M_r (Fig. 3A). Both bands were present in wild-type and kelch neo ovary extracts but were absent in hts1 mutant extracts. The monoclonal antibodies specifically immunostained ring canals (Fig. 3B). Ring canal staining was first apparent in region 2a of the germarium (see Fig. 4) and persisted on the ring canals until stage 13 (not shown). Antibody staining was reduced or absent in hts mutants depending on the severity of the hts allele examined (not shown). When there was protein present in weak hts mutants, it localized to abnormal ring canals.

Localization of actin, hts, and kelch showed a distinct temporal order of their recruitment to ring canals (Fig. 4). Actin was first seen on ring canals beginning in region 2a (Fig. 4A; Theurkauf et al., 1993). hts also began accumulating on ring canals at this time (Fig. 4B). To verify that actin and hts appear simultaneously, double labeling with anti-hts antibodies and rhodamine-conjugated phalloidin was performed (Fig. 4C). All of the ring canals appeared yellow indicating that hts (green) and actin (red) arrived on the ring canals at the same time. Finally, kelch localization to ring canals began in region 3 of the germarium. Initially, the kelch staining pattern was on a subset of ring canals, and all ring canals were labeled by stage 3 (Fig. 4D).
Anti-phosphotyrosine antibodies identify another component of ring canals

A number of colleagues informed us that anti-phosphotyrosine antibodies immunostain ring canals. Indeed, immunostaining of ovaries with anti-phosphotyrosine antibodies showed strong staining of all 15 ring canals as well as a lower level staining on membranes and in cytoplasm (Fig. 5A). We refer to the protein or proteins identified by the anti-phosphotyrosine antibodies as phosphotyrosine protein. The anti-phosphotyrosine immunostaining of ring canals began in the gerarium (Fig. 5B,C). Ring canal staining was observed as early as late region 1 when the germline syncytia are undergoing their final round of mitosis (Fig. 5C). The phosphotyrosine staining increased in amount on the ring canal by region 2a when actin and hts were first seen on the ring canals (Fig. 5B, arrow; Fig. 4). hts<sup>1</sup> mutant ring canals that lack the electron dense inner rim (Yue and Spradling, 1992) still stained with the anti-phosphotyrosine antibodies (Fig. 5D). Anti-phosphotyrosine antibodies recognized several bands on western blots of wild-type and hts<sup>1</sup> mutant ovary protein extracts (not shown) indicating that there may be several phosphotyrosine proteins involved in ring canal development.

kelch is a structural component of ring canals

The localization of kelch protein to the ring canal beginning in region 3 of the gerarium (Fig. 4D) suggested that kelch is not involved in the initial establishment of the ring canal. To investigate its role in ring canal structure, we used anti-hts antibodies and actin staining to examine ring canal structure in kelch mutants. Comparison of hts protein localization in wild-type and kelch mutant germaria suggested that the kelch mutant ring canals were morphologically normal in these early stages. The wild-type and mutant hts protein pattern resembled a tire rim with a concave outer face (Fig. 6A,B). The canal had a diameter of approximately 0.5 μm whereas the outer diameter of the ring was 1-2 μm. In wild-type stage 4 egg chambers, hts was localized in very thin bands with concave outer faces similar to the pattern found in germinaria (Fig. 6C). The canal pore was completely devoid of hts protein and had a diameter ranging between 2-3 μm depending on position of the ring canal in the egg chamber. The rings visualized by anti-hts anti-bodies in kelch mutants also had a concave outer face, but the organization was more diffuse. Many canals appeared to be occluded by diffusely distributed hts protein reducing the diameter to 0.5-2 μm (Fig. 6D). By stage 10 in wild-type egg chambers, the anti-hts staining pattern revealed rings with a smooth appearance forming a highly organized band that was slightly concave on the outer face (Fig. 6E). The inner diameter of the wild-type ring was approximately 6-7 μm and the outer diameter ranged from 7-8 μm. In stage 10 kelch mutant egg chambers, hts antibodies revealed rings that were severely disorganized with hts protein diffusely distributed through the canal (Fig. 6F). The concavity of the outer face of the rim was completely absent, and the inner diameter of the canal was reduced to approximately 2 μm. The anti-hts immunofluorescence pattern in these late stage kelch mutant ring canals appeared to be more intense either due to greater abundance of hts protein on the ring canal or due to greater availability of antibody epitopes.

The organization of actin in the ring canals was also disrupted in kelch mutants. In wild-type stage 7-8 egg chambers, the actin localized to a tight band of protein embedded in the subcortical actin meshwork (Fig. 6G). Wild-type actin rings at this stage had an outer diameter of 5-6 μm and an inner diameter of approximately 4 μm. In kelch stage 7-8 egg chambers, the actin ring had an outer diameter of 5-6 μm but an inner diameter of only 1 μm (Fig. 6H). In some kelch mutant egg chambers, the actin appeared to form a smaller ring suspended within the lumen of the canal by bundles of actin extending from the thick outer ring of actin (Fig. 6H). In wild-type stage 10 egg chambers, the actin ring formed a tightly ordered band or ring of protein with dimensions similar to the wild-type stage 10 hts protein ring (compare Fig. 6E, I). In stage 10 kelch mutant egg chambers, the actin ring was diffuse and extended into the canal (Fig. 6I). Actin staining also appeared to be more intense in kelch mutant ring canals than in the wild-type counterparts. Like the anti-hts staining, this may signify an increase in the amount of filamentous actin on the kelch mutant ring canal or a greater availability of phalloidin binding sites. Despite the altered actin organization in ring canals, cytoplasmic actin bundles still formed in stage 10 kelch mutant egg chambers (Fig. 6J arrowhead; Xue and
It appears that kelch functions to organize only the actin in ring canals and does not play a role in actin organization in other parts of the egg chamber.

Cross-sectional view of ring canals
Immunofluorescence of hts protein in early to mid-stage wild-type and kelch mutant egg chambers (Fig. 6) showed that hts forms a tire rim-shaped ring of protein. To assess the relationship between hts and actin, double labeling of wild-type ring canals for hts and actin was performed. A cross-sectional view of labeled stage 4-5 ring canals showed different localization patterns for hts and actin. A cross-sectional view of a tire rim appears as two horseshoes with the concave sides facing away from each other. hts staining appeared as horseshoes having a concave outer face (Fig. 7A, arrows). The cross-sectional view of actin in the same ring canal rim was globular rather than horseshoe-shaped (Fig. 7B, arrowheads). The merged image of the two immunofluorescence patterns showed that hts and actin largely co-localized but that the actin filled in the horseshoe (Fig. 7C).

Double-labeling experiments with combinations of anti-phosphotyrosine, anti-hts, anti-kelch, and rhodamine-conjugated phalloidin provided further insight into the structure of the ring canal. Since the phosphotyrosine protein was found on ring canals before hts and actin and was present in hts1 mutant egg chambers, we predicted that phosphotyrosine protein might be found exclusively at the outer-most rim of wild-type ring canals. Instead, double-labeling for phosphotyrosine protein and actin showed that the proteins appeared to co-localize throughout the inner rim (Fig. 8A-B). The phosphotyrosine protein at the outer rim may also be associated throughout the actin layer or there may be additional phosphotyrosine-containing proteins distributed in the inner rim. Double-labeling with anti-hts antibodies and rhodamine-conjugated phalloidin showed nearly complete overlap of actin and hts (yellow) with an enrichment of hts (green) to the inner surface of the actin (red) in ring canals in stage 3-6 egg chambers (Fig. 8C-E). Double-labeling for kelch and actin also showed an enrichment of kelch toward the inner surface of stage 3-6 ring canals with nearly complete overlap between kelch and actin (Fig. 8F). In older stage 9 and 10 egg chambers, the phosphotyrosine, actin, hts, and kelch proteins compacted so that total overlap was observed (Fig. 8G).

**DISCUSSION**
Formation of a ring canal involves sequential recruitment of type and kelch.
several proteins during germarial stages of development and dynamic rearrangement of overall ring canal rim structure during ring canal growth. By late germarium region 1 where 8-cell clusters are present, at least one phosphotyrosine protein can be detected on the ring canals. hts and actin are recruited to the ring canal shortly after the mitotic divisions are complete in region 2a. Finally, kelch is organized in ring canals beginning in region 3 of the germarium and is necessary to maintain the compaction of hts and actin in the inner rim of the ring canal. The sequential recruitment of components to the ring canal is consistent with the ultrastructure of developing ring canals. The electron opaque outer rim is established as the cleavage furrows are arrested, and the electron dense inner rim is added after the completion of the fourth mitotic division in region 2 of the germarium (Yue and Spradling, 1992).

Since a ring canal consists of two domains with different electron densities when viewed by electron microscopy, one would predict that the chemical makeup of the two domains would be different. Koch and King (1969) determined that the outer rim contained glycoproteins by staining with periodic acid/Schiff reagent, whereas the inner rim stained only with fast green at pH 2 indicating that it is made of polysaccharide-free proteins. Our immunocytochemistry experiments allow us to define further the nature of the proteins of the inner and outer rims (Fig. 9). In our model, at least one phosphotyrosine protein is adjacent to or associated with the membrane and is part of the outer rim (Fig. 9A). This or other phosphotyrosine proteins are also distributed through the actin domain of the ring canal. Actin is a component of the inner rim since its absence from the ring canal corresponds with the absence of the inner rim (Yue and Spradling, 1992). Hts and kelch are distributed in the actin domain being enriched toward the inner most face of the ring canal rim in young egg chambers (Fig. 9B). By later stages, the ring canal rim is compacted, and actin, phosphotyrosine protein, hts, and kelch are all co-localized (Fig. 9C). Compaction of the inner rim between early stage ring canals and stage 9-10 ring canals can also be observed by electron microscopy (Yue and Spradling, 1992). Proper compaction depends on the presence of the kelch protein since in its absence, the ring canal is disorganized and nearly occluded by hts and actin. In addition to compacting during growth of the egg chamber, the ring canal grows from 1 µm in diameter in region 1 of the germarium (Fig. 9A) to 7-8 µm in diameter in a stage 10 egg chamber (Fig. 9C). New proteins are added to the ring canal as it develops the inner rim from region 1 of the germarium to stage 3 in the vitellarium. It is not clear whether new addition of protein to the ring canal is required as the ring canal matures from stage 3 to stage 10. Growth might be accomplished by simply rearranging the proteins that are already present on the canal. For example, the actin filaments may slide with respect to one another in order to increase the circumference of the ring canal. However, it is also possible that ring canal growth requires additional actin polymerization and an increase in the other ring canal protein levels.

Fig. 5. A phosphotyrosine-containing protein is found on ring canals. (A) Anti-phosphotyrosine antibodies intensely immunostain ring canals in egg chambers of all stages (stage 5-6 egg chamber shown). Less intense staining is present elsewhere in the egg chamber including the anterior and posterior follicle cells (arrows). (B,C) Anti-phosphotyrosine antibodies stain ring canals very early in the germarium. Ring canals are readily identified in region 2a (large arrow), 2b, and 3. Tiny rings are visible in late region 1, just prior to when actin and hts begin to be localized to ring canals. The image in C is a magnified view (4×) of the boxed region in region 1 in B. Arrow in C identifies a late region 1 ring canal. (D) In hts1 mutant egg chambers, the cytoplasmic bridges are immunostained by anti-phosphotyrosine antibodies. In this stage 4 egg chamber, there were only three ring canals. Scale bars, 10 µm.
Fig. 6. kelch protein maintains the order of hts and actin in a growing ring canal. Wild type (A) and kelch mutant (B) germaria have a similar hts protein immunofluorescence pattern. In wild-type and in kelch mutants, the rings stained with hts antibodies are highly organized, forming a band that is concave on the outer face (arrowheads) giving a tire rim appearance. An open canal pore of 0.5 µm is apparent in wild-type and in kelch mutant ring canals (arrows). (C) By stage 4, the wild-type rings stained with hts antibodies are highly organized, appearing as thin bands that are concave on the outer face (arrowhead) and that define a wide passageway (2-3 µm) between cells (arrow). (D) In a stage 4 kelch mutant egg chamber, the hts protein is more diffuse on the ring canal rim but still forms a concave outer face (arrowhead). The center of the canal is nearly occluded by hts protein (arrow). (E) In wild-type stage 10 egg chambers, the hts protein forms a tightly organized band of protein that is slightly concave. (F) In the kelch mutant, the hts protein is localized but very diffuse. It extends into the canal nearly completely obstructing it. The increased brightness of the hts rings of the mutant egg chamber may have been due to increased amount of protein or due to increased epitope accessibility. (G) In wild-type stage 7-8 ring canals, actin forms a tight band of protein with a lumen devoid of filaments. (H) In stage 7-8 kelch mutant egg chambers, the shape of actin staining is toroidal rather than in a band as in wild type. In addition, actin is distributed into the canal forming a diffuse array. Sometimes the actin forms a smaller ring within the canal that is connected by actin bundles extending from the outer actin rim. The amount of actin appears to be increased compared to wild type. (I) In wild-type stage 10 ring canals, actin forms a highly organized band of protein similar to the hts protein band at this stage (E). (J) In the kelch mutant, actin is diffuse and extends into the lumen of the canal. The defect in actin organization is not general since normal cytoplasmic actin bundles form in stage 10 kelch mutant egg chambers (arrowhead). Scale bar, 10 µm for all panels.
The role of tyrosine phosphorylation of ring canals proteins remains unknown. Phosphotyrosine proteins have been found in other types of cell-cell and cell-substrate contacts such as adherens junctions and focal adhesions (Volberg et al., 1991, 1992). Treatment of cells with phosphatase inhibitors like H2O2/orthovanadate or tyrphostins can cause adherens junction proteins to disassemble and reassemble in focal adhesions suggesting that tyrosine phosphorylation may be an important regulatory event for the formation of these junctions (Volberg et al., 1992). While the relevant targets of these phosphorylation events remain in question, the junctions have proteins in common with cleavage furrows. For example, actin (reviewed by Burridge et al., 1988; Satterwhite and Pollard, 1992) and α-actinin (Fujiwara et al., 1978; Sanger et al., 1987) have been identified in adherens junctions, focal adhesions, and cleavage furrows (reviewed by Satterwhite and Pollard, 1992). Radixin (Tsukita et al., 1989; Sato et al., 1991) has also been identified in adherens junctions and cleavage furrows. In Drosophila egg chambers, phosphotyrosine proteins are present in cleavage furrows just prior to the arrival of actin and hts and the accumulation of the inner rim. We speculate that...
tyrosine phosphorylation of proteins found in cleavage furrows may be necessary to arrest cytokinesis or to initiate the formation of a ring canal. Anti-phosphotyrosine antibodies recognize several protein bands on western blots of whole ovary protein extracts (D. N. R., unpublished data) indicating that there are several candidates for ring canal proteins. Clearly, we need to identify the ring canal phosphotyrosine protein(s) in order to elucidate the role of tyrosine phosphorylation in ring canal development.

The hts gene encodes a component of ring canals that is recruited to ring canals at the same time as actin in region 2a of the germarium. Strong mutants for the hts gene fail to establish actin on the ring canals and fail to accumulate an inner rim (Yue and Spradling, 1992). Weak mutant alleles for the hts gene have a lower level of hts, actin, and kelch proteins on abnormal ring canals (D. N. R. and L. C., unpublished data). The mutant phenotype and the immunolocalization of the hts product suggest that hts plays a direct role in the establishment of actin on the inner rim on the ring canal. The homology of the amino-half of the predicted product of the hts cDNA to vertebrate adducin (Yue and Spradling, 1992; Ding et al., 1993) makes it tempting to speculate that hts interacts with ring canal actin through the adducin domain. However, there are several reasons to conclude that the hts ring canal protein does not contain the adducin domain. The ring canal staining antibodies were generated against the carboxy-half of the hts protein that does not share homology with any known protein. These antibodies recognize a doublet of proteins at 60×10³ M_r by western blot, which is much smaller than the 128×10³ M_r predicted by the hts cDNA (Yue and Spradling, 1992; Ding et al., 1993). Polyclonal antibodies raised to the adducin domain fail to recognize the 60×10³ M_r doublet and instead recognize a 140×10³ M_r protein (D. N. R. and L. C., unpublished data).

The 60×10³ M_r proteins and the 140×10³ M_r protein are likely to be encoded by distinct mRNAs. DNA probes that include the adducin domain hybridize to three transcripts (4.4 kb, 3.8 kb and 3.7 kb) that are expressed throughout development, including in the ovary (Yue and Spradling, 1992; D. N. R. unpublished data). Probes to the carboxy (ring canal) domain hybridize to a single ovary-specific 4.4 kb transcript (D. N. R., unpublished data), the cDNA of which has been sequenced (Yue and Spradling, 1992; Ding et al., 1993). Since the carboxy domain DNA probe recognizes a single transcript, the 60×10³ M_r ring canal proteins must be produced from this known hts cDNA (Yue and Spradling, 1992; Ding et al., 1993). Therefore, it is possible that the 60×10³ M_r ring canal protein arises from proteolysis of a precursor protein encoded by the known hts cDNA. Peptide sequences of the 60×10³ M_r proteins are required to confirm the origin and nature of the hts ring canal protein. Actin binding assays with the hts ring canal protein might confirm that it is a new type of actin binding protein.

Products of the hts gene are important at other stages of development. In addition to containing defective ring canals,
hts mutants also disrupt the formation of the fusome normally present in mitotic germline cells in the germarium (Lin et al., 1994). Antibodies to the adducin portion of the hts cDNA immunostain the fusome as well as the membranes of follicle cells surrounding egg chambers (D. N. R. and L. C., unpublished data; Lin et al., 1994). The hts protein in fusomes and cell membranes is probably functionally similar to vertebrate adducin, serving to stabilize the cortical cytoskeleton. Finally, some alleles of hts have a maternal effect lethal phenotype (Yue, 1992) that may be related to the unusual anterior localization of hts mRNA in oocytes late in oogenesis (Yue and Spradling, 1992; Ding et al., 1993).

The last protein we have found to be recruited to ring canals is kelch. It accumulates after the ring canal is established and is essential for maintenance of ring canal rim structure. In a kelch mutant ring canal, both hts and actin protein distributions are disorganized, and the proteins fail to compact (Fig. 6). This suggests that kelch interacts with actin or hts directly to organize or bundle actin into the neatly packed inner rim of the ring canal. An interaction with actin may be mediated by the carboxy-terminal kelch repeats. The gene for the Limulus actin binding protein, scruin (Tilney, 1975), was cloned and found to encode two sets of six kelch repeats, one in the amino-terminal half and the second in the carboxy-terminal half (Way and Matsudaira, personal communication). Ultrastructural analysis of Limulus sperm actin bundles (Owen and DeRosier, 1993; Schmid et al., 1994) shows that the repeat domains bind adjacent actin monomers in a filament. The carboxy-terminal repeat domain may be responsible for bundling since it appears to associate with a carboxy-terminal repeat domain of a scruin molecule in a neighboring actin filament (Bullitt et al., 1988). By analogy, it is probable that kelch binds to an actin filament through its repeat domain and could also provide a bundling function. The morphology of kelch mutant ring canals supports the idea that kelch is providing a bundling function. Actin filaments in kelch mutant ring canals are distributed in the lumen of the canal, sometimes appearing to form a smaller ring suspended within the canal lumen. In the absence of kelch, any remaining actin filament bundling proteins might bundle actin filaments inappropriately in a loose meshwork. The absence of kelch may even allow the binding of actin binding proteins not normally found in ring canals. This could result in a substantially different filament crosslinking pattern that accounts for the diffuse appearance of actin in kelch mutants.

In kelch mutant ring canals, both hts and actin appear to be present in increased amounts (Fig. 6) suggesting another possible function for kelch. kelch may limit actin polymerization on the ring canal rim by capping growing filaments or by sterically limiting actin polymerization. However, we cannot rule out the possibility that ring canal assembly and growth require a carefully regulated balance between the different ring canal protein levels. Loss of one of these (for example, kelch) may perturb this balance resulting in aberrant ring canal morphology.

The failure to complete cytoplasmic transport to the oocyte in kelch mutant females (Xue and Cooley, 1993) can now be attributed to occluded ring canals. The transport defect appears to begin early in the vitellarium at the same time that the disorganization of hts and actin on the inner rims of ring canals first becomes apparent. The partially occluded ring canals do not block all transport because some mRNAs and proteins that are transported at specific times in egg chamber development still arrive on time in kelch mutants (D. N. R. and L. C., unpublished data). Oocyte growth in kelch mutant egg chambers is retarded perhaps because the mutant ring canal cannot permit passage of larger cytoplasmic components such as mitochondria or because the mutant ring canal cannot accommodate the enormous volume of cytoplasm necessary for oocyte maturation.

Conclusion

Cytoplasmic bridges provide important intercellular connections between developing gametes of many invertebrate and vertebrate organisms. We have gained insight into the development and structure of germline ring canals in Drosophila females. We expect ring canal structure is somewhat different in males since both hts and kelch are absent from the testes (D. N. R., unpublished data). Perhaps the transport of cytoplasm to a selected cell in female syncytia places a special structural burden on the intercellular junctions. Ultrastructural similarities between female ring canals of insects and mammals, in addition to the presence of a possible kelch homolog in mouse (MIPP), suggest that mammalian ring canals might be similar to Drosophila ring canals. It is possible that the sequential addition of proteins to a stabilized cleavage furrow during the formation of a ring canal may be a recurring theme found in other systems. Additional genetic and biochemical analyses of Drosophila ring canal proteins and a study of homologous proteins in the mouse are needed to determine the degree of conservation of this fascinating structure.

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REFERENCES


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