Stable intercellular bridges are a conserved feature of gametogenesis in multicellular animals observed more than 100 years ago, but their function was unknown. Many of the components necessary for this structure have been identified through the study of cytokinesis in Drosophila; however, mammalian intercellular bridges have distinct properties from those of insects. Mammalian germ cell intercellular bridges are composed of general cytokinesis components with additional germ cell–specific factors including TEX14. TEX14 is an inactive kinase essential for the maintenance of stable intercellular bridges in gametes of both sexes but whose loss specifically impairs male meiosis. TEX14 acts to impede the terminal steps of abscission by competing for essential component CEP55, blocking its interaction in nongerm cells with ALIX and TSG101. Additionally, TEX14-interacting protein RBM44, whose localization in stable intercellular bridges is limited to pachytene and secondary spermatocytes, may participate in processes such as RNA transport but is nonessential to the maintenance of intercellular bridge stability.
Figure 1. Intercellular bridge formation in Drosophila. Both male and female germ cells initially go through four divisions to form a 16-cell cyst (A,B). In males, meiotic divisions increase the number of germ cells in the cyst to 64. These cells remain connected through the intercellular bridges by the fusome (A). In females, the fusome initially connects all 16 cells through intercellular bridges, with two cells containing the greater fraction of the fusions (B, cells 1 and 2). The fusome then breaks down after oocyte selection. (A, Adapted from Hime et al. 1996; reprinted with permission from The Company of Biologists © 1996. B, Adapted from Huynh and St Johnston 2004; reprinted with permission from Elsevier © 2004.) Both male (C) and female (D) Drosophila intercellular bridge formation begins in both sexes with an increase in phosphotyrosine epitopes, but subsequent processes are distinct. (See facing page for legend.)
detail by microscopy for more than half a century, we have only recently begun to learn how mammalian germ cell intercellular bridges form and function at the molecular level. Inroads have been made to determine the role of this unique mode of intercellular communication. Many molecules and organelles have been observed to pass through intercellular bridges, but their role in fertility is not fully understood. Although the bridges themselves have been conserved, the way they form and their function appears to vary between gender and species.

This article will review our current understanding of how intercellular bridges form and the role they play in gametogenesis. Examples will primarily be drawn from mammalian and Drosophila models where the most study has been completed. We will first provide a general description of somatic cell cytokinesis, because modification of this process forms intercellular bridges. Next, we will review Drosophila melanogaster ring canals, the name given to intercellular bridges in that species. We will conclude with what is known about formation and function of mammalian intercellular bridges.

INTERCELLULAR BRIDGES (RING CANALS) IN DROSOPHILA MELANOGASTER

Syncytial development of germ cell cysts is conserved from fruit flies to mammals in both males (Burgos and Fawcett 1955; Fawcett et al. 1959; Dym and Fawcett 1971; Moens and Hugenholtz 1975; Huckins and Oakberg 1978; Weber and Russell 1987; Ren and Russell 1991; Ventela et al. 2003) and females (Gondos et al. 1971; Pepling and Spradling 1998, 2001; Pepling et al. 1999). Drosophila male and female gametes develop clonally (Rasmussen 1973; Spradling et al. 2001; Gonzalez-Reyes 2003) from a single founder cell, the gonial cell in males and the cystoblast in females. In both genders, the founder cell initially divides four times. “Incomplete” cytokinesis during these divisions results in the formation of a 16-cell cyst (Fig. 1A) (Hime et al. 1996; Huynh and St Johnston 2004). The cells of the cyst are connected by intercellular bridges, also called ring canals in Drosophila. The number of intercellular bridges borne by each germ cell in the cyst directly correlates to the round of division from which it was derived. Thus, the founder cell and its first daughter cell will each have four intercellular bridges, their two daughter cells will have three bridges, the next four cells will have two bridges, and the final eight cells have only one bridge, for a total of 15 shared intercellular bridges (Fig. 1B) (intercellular bridges in cysts are in green).

Sixteen cells is the final number of germ cells in the ovarian cyst. Once this number is reached, the cyst migrates posteriorly in the ovary as it matures, and one germ cell will be selected as the oocyte and supported by the other 15 “nurse” cells (Huynh and St Johnston 2004). When male cysts reach 16 cells, two meiotic divisions occur to form a total of 63 shared intercellular bridges. This pattern of division is highly reproducible and, at least in females, is dependent on a structure called the fusome. By comparison, the number of cells in a mammalian germ cell cyst is much greater and less regulated (see below). Furthermore, no
equivalent of the fusome has been identified in mammals.

The fusome (Fig. 1A,B, red) is a structure containing spectrin and hu-li tai shao (Hts), an adducin homolog (Yue and Spradling 1992) that connects the germ cells of the cyst through their intercellular bridges. The fusome is derived from a spherical structure in the germline stem cell called the spectrosome (Lin et al. 1994; de Cuevas et al. 1996). In females, the cystoblast inherits one third of the fusome from the germline stem cell (Deng and Lin 1997). As the germ cells divide to form the cyst the fusome grows and is distributed, but a larger fraction of the fusome is kept in one of the two cells formed by the first division. This asymmetric division is repeated three times. At the end, the cell with the most fusome material (one of the first two cells) will become the oocyte (de Cuevas and Spradling 1998), and the fusome breaks down (Fig. 1B). Hts and α-spectrin mutants lack a fusome and fail to specify an oocyte. Additionally, the divisions become randomly oriented and result in a variable number of cells (Yue and Spradling 1992; Lin et al. 1994; de Cuevas et al. 1996).

In males, all 16 germ cells of the cyst are equivalent, and the fusome is maintained although meiosis and connects all 64 haploid spermatids (Fig. 1B) (Hime et al. 1996). The fusome plays a role in centrosome inheritance in spermatogenesis. A male sterile allele of hts eliminates the fusome. In this mutant dividing spermatogonia all have two centrosomes and a normal mitotic spindle, but spermatocytes have a variable number of centrosomes and defective spindles.

Passage of the fusome through intercellular bridges is a common feature of male and female Drosophila gametogenesis, and fusome communication through intercellular bridges is essential for fertility. Although intercellular bridges play a similar role in both genders in this respect, the mechanism of intercellular bridge formation differs in male and female Drosophila.

Stable intercellular bridges between male germ cells are composed of proteins involved in cytokinesis (Hime et al. 1996), namely, anillin and three Drosophila septins, Peanut (Pnut), Sep1, and Sep2. Septins are required for somatic cell cytokinesis in some, but not all, cell types (Glotzer 2005), and are capable of self-assembly into rings (Kinoshita et al. 2002). Anillin, which localizes early to the cleavage furrow (Field and Alberts 1995), directly interacts with septins (Field and Alberts 1995; Kinoshita et al. 2002), and was initially identified as an actin-binding protein in Drosophila embryos (Miller et al. 1989).

The earliest sign of intercellular bridge formation in both male and female Drosophila is the appearance of phosphotyrosine epitopes in late telophase (Fig. 1C,D) (Robinson and Cooley 1996). These “rings” of phosphotyrosine epitopes remain in the intercellular bridge even after it matures, growing in diameter with the bridge as the germ cells develop. After the appearance of phosphotyrosine epitopes, myosin II and actin from the contractile ring are lost from the mature intercellular bridge in male Drosophila (Fig. 1C). In contrast, actin remains a major component of the mature female intercellular bridge (Fig. 1D), but anillin and septins are no longer present. These are not the only compositional differences between male and female Drosophila intercellular bridges. Hts and kelch, essential proteins in the female intercellular bridge, are absent from male bridges. In fact, only one protein has been identified as a component of both male and female intercellular bridges, Pavarotti/MKLP1 (Carmena et al. 1998; Minestrini et al. 2002). Interestingly, although male germ cell intercellular bridges have different components from female germ cell intercellular bridges, they may be similar to intercellular bridges that have been reported connecting somatic cells of the follicular epithelium in ovary, imaginal disc cells, and larval brain in Drosophila (Kramerova and Kramerov 1999). There is also a single report of somatic intercellular bridges seen by electron microscopy in mammals, but the authors caution that it may be the result of a tissue preparation artifact (Witkin et al. 1995).

Brill et al. (2000) used Drosophila spermatocytes to screen for cytokinesis-defective mutations, because mutants with failure of cytokinesis in meiosis are easily identified by their sterility and multinucleated spermatids.
Nineteen genes were identified, including 16 novel loci and three known cytokinesis genes. The mammalian homolog is given, if known. However, detailed mapping of the mutated genes was not performed, so the actual gene products were not all identified. The mutants blocked cytokinesis at several stages (Fig. 1B) (Brill et al. 2000). Two mutants, *diaphanous* (*dia*, *Diaphanous* homolog) and *pebble* (*pb1*, *ECT2/RhoGEF* homolog), prevent formation of the actin and anillin rings in early telophase. Five of the mutants, *scapolo* (*scpo*), *celibe* (*cbe*), *james bond* (*bond*), *sauron* (*sau*), and *smeagol* (*sgo*), affect the central spindle and the F-actin ring, but do not prevent anillin accumulation at the equator. Ten genes, *four score* (*fsc0*), *brunelleschi* (*bru*), *ciamella* (*cia*), *four way stop* (*fws*, *Cog5* homolog), *funnel cakes* (*fun*), *omelette* (*onit*), *onion rings* (*onr*), *four wheel drive* (*fwd*, a phosphatidylinositol 4-kinase), and *giotto* (*gio*), are primarily required for constriction of the F-actin ring. Lastly, *bird’s nest soup* (*bns*) was required for actin ring disassembly. It is unclear if any of these genes are specifically involved in intercellular bridge formation. The effect on spermatogonial intercellular bridges in these mutants was not reported, but it can be inferred that the phenotypes were specific for spermatocytes. The fact that the known genes, *dia*, *pb1*, *fwd*, and *fws*, have previously been implicated in cytokinesis further supports the model that male *Drosophila* intercellular bridges are composed of cytokinesis proteins.

Female *Drosophila* intercellular bridges are required even after the oocyte is specified. Intercellular bridges connecting the 15 nurse cells to the oocyte grow dramatically in size from <1 μm to ~10 μm as the egg chamber matures (Cooley 1998). The asymmetric division of the fusome required for oocyte specification also causes microtubules to become polarized with their minus ends directed at the oocyte (Grieder et al. 2000). These modifications allow the ring canals to transfer several cytoplasmic components from the nurse cells to the oocyte, increasing its size. For example, oocyte-specific proteins and mRNAs (e.g., *Bicoid D* [Mische et al. 2007] and *oskar* [Huynh and St Johnston 2004]) are concentrated in the oocyte. The centrosomes from the nurse cells are inactivated and transferred to the oocyte (Bolivar et al. 2001), and mitochondria are also concentrated in the oocyte (Cox and Spradling 2003).

As in males, the first evidence of female *Drosophila* intercellular bridge formation is an increase in phosphotyrosine epitopes (Fig. 1D) (Robinson and Cooley 1996). The phosphotyrosines are seen as 0.5–1-μm rings in the stalled cleavage furrows at the end of the third mitosis (Robinson et al. 1994). The development of female intercellular bridges is known to require the convergence of cytoskeletal and kinase signaling pathways not regularly involved in cytokinesis (Fig. 1F) (Cooley 1998). First, consider the cytoskeletal pathway (Fig. 1F). By the fourth division *hu-li tai shao* (*Hts*) (Robinson et al. 1994) and additional filamentous actin (*Tilney et al. 1996*) localize to the inner rim of the intercellular bridge (Fig. 1D). Over the next few hours Kelch is added to the inner rim (Robinson et al. 1994), while anillin is removed from the outer rim of the intercellular bridge, and the bridge’s diameter increases to 3–4 μm (Robinson and Cooley 1997). At this point, the bridges are considered “mature,” but will continue to increase in diameter. Incorporation of Hts, Kelch, and filamentous actin all depend on *cheerio* (the mammalian *filamin* homolog), an actin cross-linking protein. Other proteins involved in actin dynamics such as the Arp2/3 complex (Hudson and Cooley 2002), Cortactin (Somogyi and Rorth 2004), and *Akap200* (Jackson and Berg 2002) have also been shown to regulate the intercellular bridge cytoskeleton.

Kelch is able to dimerize through its BTB domain and bind to actin with its Kelch repeat domain (Robinson and Cooley 1997). It may therefore serve as an actin cross-linking protein in the growing intercellular bridge (Robinson and Cooley 1997). *Kelch* mutant females are sterile, but otherwise the flies are completely normal (Robinson et al. 1994). Loss of Kelch causes disorganization of the actin filaments of the intercellular bridge preventing cytoplasm transport from the nurse cells to the oocyte (Robinson et al. 1994). The *kelch* gene produces a single transcript separated into two open reading frames by a stop codon. Although both the
truncated, ORF1, and full-length versions of the protein are expressed by tRNA suppression of the stop codon, and both proteins are in the intercellular bridge. Oddly, ORF1 is sufficient to fully rescue the *kelch* intercellular bridge phenotype but full length is not.

*Kelch* is one protein in which the kinase regulatory pathway converges with the cytoskeletal pathway (Fig. 1F). The ability for *Kelch* to organize actin is regulated by the *Drosophila* Src nonreceptor tyrosine kinase homolog *Src64*. *Src64* is does not effect the localization of Hts of *Kelch*, but its kinase activity is required for expansion of the intercellular bridge diameter (Dodson et al. 1998). *Src64* signaling is required for phosphorylation of *Kelch* at tyrosine 627 (Kelso et al. 2002). Using site-directed mutagenesis, *KelchY627A* is able to rescue the actin disorganization phenotype of *kelch* mutant intercellular bridges, but the bridges fail to expand, phenocopying the *src64* mutant phenotype. Actin monomer exchange is also greatly reduced with *KelchY627A*. These results suggest that a major function of *Src64* signaling is negative regulation of actin cross-linking by *Kelch* (Kelso et al. 2002).

Although *Src64* is required for the phosphorylation of *Kelch*, it may not be the kinase that actually phosphorylates the protein. Mutations of another nonreceptor tyrosine kinase, Tec29, phenocopies the *Src64* intercellular bridge phenotype (Roulier et al. 1998). Tec29 localizes to the intercellular bridge. Although *Src64* does not localize to the intercellular bridge, it interacts with Tec29, and *Src64* activity is required for Tec29 localization (Roulier et al. 1998; Lu 2004; Lu et al. 2004). The loss of either kinase results in a dramatic decrease in phosphotyrosine epitopes in the intercellular bridge. Although Tec29 may be the kinase that regulates *Kelch*, the only function currently known for Tec29 kinase activity is the creation of phosphotyrosine binding sites on unidentified intercellular bridge substrates for its own SH2 domain (Lu et al. 2004).

This overview of male and female *Drosophila* intercellular bridges shows that, although essential for fertility in both genders, they form by very different mechanisms. Mammalian intercellular bridges were also initially thought to have an essential role in fertility for both genders. The next section will review the current understanding of mammalian germ cell cysts.

**MAMMALIAN INTERCELLULAR BRIDGES**

Until 1955, the term “intercellular bridge” referred to short processes that appeared to connect squamous epithelial cells (Fawcett 1961). Electron microscopy has since shown that these structures are actually desmosomes, not true cytoplasmic connections between cells. As reviewed by Dym and Fawcett (1971), as early as 1865 LaValette St. George teased apart fresh testis and noticed that cells were able to divide, yet remain connected in a chain. Later observations by Sertoli (1877) (Fig. 2A) and von Ebner (1888) (Fig. 2B) described spermatogenesis connected by processes. It was not until 1955 when electron microscopy studies examining the fine structure of cat spermatids defined the intercellular bridge as a stable cytoplasmic channel connecting cells (Fawcett et al. 1959). Shortly after, intercellular bridges were found in many animals including Hydra, fruit fly, opossum, pigeon, rat, hamster, guinea pig, rabbit, monkey, and man (Fawcett et al. 1959).

Intercellular bridges have become the basis for describing early stages of spermatogenesis. A widely used model of spermatogenesis is based on the work of Huckins (1971) and Oakberg (1971). Male germ cells start as diploid spermatogonia, proceed through meiosis as spermatocytes, and complete development as haploid spermatids. The complete process involves nine to 11 spermatogonial divisions (Huckins 1971; de Rooij and Russell 2000; Chiarini-Garcia and Russell 2002) plus two meiotic divisions. Each division results in the formation of an intercellular bridge (Weber and Russell 1987), and spermatogonia are initially classified by their intercellular bridges. The nomenclature for spermatogonial development is as follows:

\[ A_3 \rightarrow A_{pr} \rightarrow A_d (\text{multiple rounds}) \rightarrow A_1 \]  
\[ \rightarrow A_2 \rightarrow A_3 \rightarrow A_4 \rightarrow In \rightarrow B. \]
The A type single (A<sub>s</sub>) spermatogonium is the stem cell for spermatogenesis and is the only dividing male germ cell without intercellular bridges. A single mouse testis is thought to contain about 35,000 A<sub>s</sub> spermatogonia (Tegelenbosch and de Rooij 1993). The A<sub>s</sub> cell differentiates by dividing into A type paired (A<sub>pr</sub>) spermatogonia. A<sub>pr</sub> spermatogonia divide to form a chain of four A type aligned (A<sub>al</sub>) spermatogonia, and A<sub>al</sub> spermatogonia can go through multiple rounds of division before differentiating into A<sub>1</sub> spermatogonia (Huckins...
The remaining stages through intermediate and B type spermatogonia link mitotic division with progression to the next stage. This process theoretically leads to syncytia joining thousands of cells (Moen and Hugenholtz 1975; de Rooij and Russell 2000). In practice, up to 650 cells have been observed in a single syncytium (Ren and Russell 1991).

The different stages of spermatogonia can be defined by a skilled examiner by their appearance and morphology in light (Huckins 1971; Chiarini-Garcia et al. 2001) and electron microscopy (Chiarini-Garcia and Russell 2002). Early stages of spermatogonia, such as Aα, are most easily identified in whole mounted tubules. The examiner focuses on the basal plane, where the least differentiated cells lie (Clermont and Bustos-Obregon 1968). Aα cells are identified when no other A type spermatogonia is within 25 μm of the cell. Aα, Aγ, and Aαl spermatogonia can also be identified with the molecular markers such as promyelocytic leukemia zinc finger (PLZF), a protein required for spermatogonial stem cell self-renewal (Buaas et al. 2004).

The morphology of intercellular bridges has been studied exhaustively by microscopy (e.g., Burgos and Fawcett 1955; Fawcett et al. 1959; Dym and Fawcett 1971; Moens and Hugenholtz 1975; Huckins and Oakberg 1978; Weber and Russell 1987; Ren and Russell 1991; Ventela et al. 2003). The diameter of the bridge ranges in size starting at under 1 μm in spermatogonia, 1–1.3 μm in intermediate and B type spermatogonia, 1.4–1.7 μm in spermatocytes, 1.8 μm in step 1 spermatids, 3 μm in step 18 spermatids, and then shrinking slightly to 2 μm in step 19 spermatids (Weber and Russell 1987). By electron microscopy, a prominent electron dense “bridge density” ranging from 30–60 nm in thickness lines the walls of the intercellular bridge. In cytokinesis, the bridge density begins to form on the inner edge of the cleavage furrow during ingression. A midbody with normal appearance forms and then breaks down with the central spindle, leaving the intercellular bridge and the lining bridge density (Weber and Russell 1987).

Several roles for mammalian intercellular bridges in spermatogenesis have been proposed (Fig. 2C) (Guo and Zheng 2004). Two hypotheses have the most support. The first hypothesis is that intercellular bridges permit “cytoplasmic sharing” of essential signals for the synchronous cell divisions seen in longitudinal segments of seminiferous tubules (Fawcett et al. 1959; Huckins and Oakberg 1978; Ren and Russell 1991). In this untested model, synchronization is proposed to play an essential role in progression through spermatogenesis. A second hypothesis points out that intercellular bridges may be necessary for haploid germ cells to remain “phenotypically diploid” after meiosis (Fawcett et al. 1959; Erickson 1973; Braun et al. 1989). The most extreme example of this would be for genes with only one copy (e.g., genes on the sex chromosomes). Supporting this second hypothesis, mRNA expressed from a single copy transgene encoding human growth hormone (hGH) under the control of the protamine 1 regulatory sequences was observed to distribute evenly in haploid spermatids (Braun et al. 1989). The protein product was also found equally distributed among the spermatids. In a nontransgenic example, the testis brain RNA-binding protein (TB-RBP) functions as an RNA-binding protein in the testis and binds to Akap4 mRNA. Akap4 is an X-linked gene essential for sperm motility and fertility (Miki et al. 2002). Akap4 mRNA forms a complex with TB-RBP and the Ter ATPase and passes through intercellular bridges between spermatids (Morales et al. 2002). Additionally, because of the large size of intercellular bridges, even organelles (Ventela et al. 2003) have been observed to move between haploid spermatids.

A second hypothesis proposes that intercellular communication between germ cells occurs at “critical stages,” such as coordinating the entry into meiosis (Stanley et al. 1972; Robinson and Cooley 1996). In this model, the event is coordinated by the “rapid” transfer of an unidentified signal/substance through the syncytium. A third hypothesis is that intercellular bridges create “gamete equivalence” and would thus result in a more uniform population of sperm which, overall, would be of higher quality (Guo and Zheng 2004). A final hypothesis is that intercellular bridges allow for the
elimination of cells with “self-promoting genes” (LeGrand 2001). In this model, a mutation exists which would give a selective advantage to the mutated gamete but would be detrimental to future progeny. For example, the cell could have uncontrolled and greatly increased cell proliferation and have a mutation in a gene required for apoptosis. Normally functioning cells in the syncytium detect something is wrong through the intercellular bridges and the problem is eliminated by apoptosis of the entire cyst. Of course, because all the cells are clonally related, the mutation would have to be acquired somehow after the initial A1 cell division.

To test these hypotheses requires manipulation of the intercellular bridge. The molecular mechanism of mammalian intercellular bridge formation is a recent area of study. The protein TEX14 is expressed specifically in germ cells and localizes to the intercellular bridge in mice (Greenbaum et al. 2006) and humans (Fig. 3A) (Greenbaum et al. 2007). The protein contains three amino-terminal ankyrin repeats, centrally is a dead-kinase domain, and the remainder contains coiled-coil motifs. Interestingly, when compared by BLAST analysis to the Drosophila genome, the dead-kinase domain bears more resemblance to src64 and tec29 (see above) than any other kinases.

Knockout of mouse Tex14 eliminates the formation of intercellular bridges (Greenbaum et al. 2006) and results in male infertility. It was found that without intercellular bridges spermatogenesis fails to complete meiosis. In fact, molecular studies showed that only a very small number of spermatocytes make it to pachytene. Importantly, this result proved that intercellular bridges are not required for the transit amplification of germ cells from the A1 stem cells spermatogonia through the initiation of meiosis.

Although it is clear completion of meiosis during spermatogenesis requires intercellular bridges, the exact role of the intercellular bridge remains unknown. Many of the proposed roles listed above may be relevant. Perhaps intercellular bridges do allow passage of a signal for entering the “critical stage” of meiosis through an unknown mechanism. Interestingly, mammalian somatic cells die through a p53-dependent mechanism if they fail to complete cytokinesis (Postiglione et al. 2009). For germ cells, incomplete cytokinesis is the norm and it is completion that results in increased apoptosis (Greenbaum et al. 2006). The one function intercellular bridges are known to play, passage of RNA material in haploid cells, could not be evaluated because no germ cells progressed through meiosis. Thus, intercellular bridges are required for spermatogenesis but their essential role remains undiscovered.

Embryonic mammalian male (Gondos and Hobel 1971; Fukuda et al. 1975) and female

Figure 3. TEX14 is a critical component of mammalian intracellular bridges. (A) Immunohistochemical staining of control testes using anti-TEX14 antibody labels ring-shaped structures (arrows) that are absent from TEX14 null germ cells. (B–E) Electron microscopy reveals electron dense material lining cytoplasmic channels between newborn ovarian germ cells. No intercellular bridges were found in knockout ovaries. ICB, intercellular bridge; M, mitochondria; N, nucleus.
(Gondos et al. 1971; Pepling and Spradling 1998, 2001; Pepling et al. 1999) germ cells are also connected by intercellular bridges. Mitotic division of germ cells occurs for a finite period during embryogenesis in mammals. Germ cell division in the embryonic ovary is synchronized, presumably through intercellular bridges (Pepling and Spradling 1998). In female mice, germ cells enter meiosis around E14.5 (Borum 1961; Pepling and Spradling 1998). No divisions occur beyond meiosis, so no new intercellular bridges form beyond embryogenesis. Female germ cell cysts break down over the second through fourth day after birth (Pepling and Spradling 2001) leaving single oocytes to form primary follicles. Similar to Drosophila, intercellular bridges have been proposed to allow “nurse cells” to promote oocyte development (Pepling and Spradling 2001). Supporting this idea in mice, mitochondria have been shown to reorganize and increase in number around the time of birth (Pepling et al. 1999; Pepling and Spradling 2001).

The Tex14 knockout mice were used to evaluate the role of embryonic intercellular bridges. Tex14 was found to localize to embryonic bridges and intercellular bridges seemed to be absent in Tex14 knockouts by electron microscopy (Fig. 3B). Still, a second method of confirmation was desired to show the bridges were truly not present. Immunofluorescence is the best way to look at a large amount of tissue for intercellular bridges, but there were no other known embryonic bridge components. A biochemical approach was taken to find more components of the intercellular bridge (Greenbaum et al. 2007). Testis bridges were used because they were more abundant and easier to obtain. Several new components were identified, and the order in which they joined and left the intercellular bridge could be followed in newborn testis. Using one of these new intercellular bridge markers it was confirmed by immunofluorescence that the embryonic intercellular bridges were gone.

Surprisingly, Tex14 knockout females remained fertile without intercellular bridges! This was not the case in Drosophila in which intercellular bridges play a well-described critical role in fertility (see above). Apparently, despite a billion years of conservation, female intercellular bridges are not essential. That is not to say they do not play an important function. It is obvious that selective pressures exist in nature that are absent in a controlled laboratory environment. However, there is a possibility that the female bridges exist as a vestigial structure purely because of the essential role intercellular bridges play in spermatogenesis.

A SPECIAL FORM OF CYTOKINESIS

A typical description of cytokinesis is presented as, “Cytokinesis is the final process in cell division through which one cell is separated into two daughter cells.” Clearly, this is not the best definition for a section whose central topic is intercellular bridges, the stable cytoplasmic connection joining the daughter cells. Nonetheless, it is important to describe the basic principles of cytokinesis, as intercellular bridge formation is a modification of how the vast majority of cells divide. Although the most dramatic event of cytokinesis in mammalian cells is ingestion of the cleavage furrow, this single event is not the whole picture. The process of cytokinesis can be divided into at least three broad stages as follows: (1) selecting the site of cell division (Burgess and Chang 2005), (2) furrow ingression and formation of a contractile ring (Matsumura 2005; Wang 2005), and (3) abscission of the midbody (von Dassow and Bement 2005). Although the end result of this process, cell separation, is the same in all eukaryotic organisms, there are several variations on the theme (Guertin et al. 2002), with plants even replacing the processes of furrow formation and ingestion with a cytoskeletal vesicular framework called the phragmoplast (Otegui et al. 2005). For our purpose, the
following section, when possible, will review the mechanism of cytokinesis as it relates to mammalian cells. Furthermore, although early events in cytokinesis will be addressed briefly, the greatest focus will be on the late events of cytokinesis, as these directly relate to intercellular bridge formation.

Early in cytokinesis, it is necessary to select the site of cell division. In animal cells, the region where the cleavage furrow will form is specified by the microtubules of the mitotic apparatus at the onset of anaphase. This linkage to the mitotic apparatus provides the necessary coordination of chromosomal segregation with cell division. The cleavage site forms at the former metaphase plate, bisecting the two centrosomes. Microtubule depolymerization studies and manipulation of the spindle apparatus (Rappaport 1986; Rieder et al. 1997) show that the mitotic spindle somehow signals to the cortex to determine the region where the cleavage furrow will form. Despite more than a hundred years of study pioneered by Ray Rappaport (Canman and Wells 2004), most aspects of the signaling process remain unknown (Burgess and Chang 2005). Both the signal and the region of the mitotic apparatus that emits the signal remain to be identified. Although many models have been proposed (Canman and Wells 2004; Burgess and Chang 2005), they each have drawbacks. The mechanism of cleavage furrow site selection seems distinct from intercellular bridge formation because it is an early event in cytokinesis that occurs normally in mammalian germ cells. For this reason, these models will not be discussed in further detail.

Furrow ingression follows site selection. The predominant model for the process of furrow formation is the contractile ring hypothesis (Satterwhite and Pollard 1992). This process is very similar in yeast and animal cells, which both divide using an actomyosin contractile ring. Actin and myosin II assemble in a ring around the perimeter of the putative cleavage site. Like a contracting myofiber, antiparallel actin filaments slide together through the action of myosin II resulting in contraction of the ring. The process often draws comparison to a “purse string” as it squeezes the plasma membrane (Wang 2005). The contraction of the equatorial membrane generates enough force to bend a flexible microneedle (Rappaport 1967). As in cleavage site selection (above), the mitotic spindle (now called the central spindle or spindle midzone during furrowing) is generally accepted to regulate cleavage furrowing (Matsumura 2005).

Several evolutionarily conserved complexes essential for cytokinesis have been identified in the central spindle (Glotzer 2005). These are the Aurora B-INCENP-CSC1-Survivin complex, the PRC1-KIF4 complex, the ECT2/RhoGEF (Glotzer 2005), and the mitotic kinesin-like protein 1 (MKLP1)-Male germ cell Rac GTPase-activating protein (MgcRacGAP) complex (also called centralspindlin) (Mishima et al. 2002). Loss of any of these complexes causes cytokinesis failure. It should be pointed out that the name MgcRacGAP (Male germ cell Rac GTPase-activating protein) is somewhat misleading. Although originally described as having predominantly male germ cell expression (Toure et al. 1998), like MKLP1, it is ubiquitously expressed and required for cytokinesis in all cells. Accordingly, MgcRacGAP homozygous gene trap mice die during preimplantation development with multinucleated blastomeres and a dramatic decrease in cell number at day E3.5 because of cytokinesis failure (Van de Putte et al. 2001).

The completion of furrow ingression forms a transient intercellular bridge called the midbody, which connects the daughter cells at the end of cytokinesis. Germ cells form intercellular bridges by altering the fate of the midbody. Rather than discard the transient intercellular bridge by abscission, it is modified into a permanent structure. The composition of the midbody matrix contains several of the conserved complexes mentioned previously (Glotzer 2005). In particular, the centralspindlin complex (Mishima et al. 2002), MKLP1 (Matuliene and Kuriyama 2004), and MgcRacGAP (Arar et al. 1999), is essential for forming the midbody matrix. RNA interference (RNAi) against MKLP1 in mammalian cells prevents formation of a normal midbody matrix resulting in
unstable midbodies and failure of cytokinesis (Matuliene and Kuriyama 2004). MKLP1 in the midbody exists in a complex with 1-to-1 stoichiometry with MgcRacGAP, and this complex, centralspindlin, has been reconstituted and shown to bind microtubules in vitro (Mishima et al. 2002). It also indirectly recruits components needed for abscission.

Abscission is the final process in cytokinesis by which the midbody is removed to achieve truly individual cells. Compared to cleavage furrow ingression, which takes \( \approx 20 \) min, abscission is a relatively slow process, requiring 1–3 h to complete (von Dassow and Bement 2005). At least two processes are necessary to achieve abscission. First, the organization of the matrix in the middle of the midbody forms a central ring. By light microscopy, this structure is called the Flemming body, a slightly bulging region right in the middle of the midbody. The formation of the midbody ring is followed by membrane fusion events to cleave the midbody (von Dassow and Bement 2005).

Many of the proteins required in these processes have been identified (Gromley et al. 2003, 2005; Low et al. 2003).

There are several vesicle-associated proteins whose essential role in abscission has been identified. The first identified proteins of this class were syntaxin 2 and endobrevin (Low et al. 2003), members of the soluble NSF attachment receptor (SNARE) complex required for SNARE-mediated membrane fusion events (Hanson et al. 1997). Syntaxin 2 and endobrevin colocalize laterally to the midbody matrix. RNAi showed that both proteins are essential for abscission of the midbody in HeLa cells (Low et al. 2003).

The identification of other trafficking proteins came, surprisingly, from studying centrosomal proteins. Centriolin, initially known only as a centrosome component, was found to localize to the midbody of HeLa cells (Gromley et al. 2003). RNAi of centriolin did not lead to a microtubule defect but resulted in a novel cytokinesis phenotype (Gromley et al. 2003). Rather than form multinucleated cells, as seen in previous cytokinesis defects, loss of centriolin caused long, thin intercellular bridges to remain connecting the daughter cells. Further studies showed that the SNARE proteins, syntaxin 2, endobrevin, and snapin, require centriolin for localization (Gromley et al. 2005). The exocyst complex (Hsu et al. 2004) is required for vesicle mediated secretion and also plays a role in abscission. Centriolin is also required for localization of the exocyst complex to the midbody. Exocyst complex proteins sec3, sec5, sec8, sec15, and exo84 failed to localize to the midbody in centriolin RNAi knockdown experiments (Gromley et al. 2005). Another centrosomal protein, centrosome protein 55 kDa (CEP55), was found to localize to the midbody matrix in somatic cells (Fabbro et al. 2005; Martinez-Garay et al. 2006). CEP55 was found to bind to the centralspindlin complex (Zhao et al. 2006). Deficiency of CEP55 leads to arrest at the midbody stage in somatic cells and formation of multinucleated cells (Carlton and Martin-Serrano 2007; Morita et al. 2007). In addition, knockdown of the direct downstream interacting partners of CEP55 such as ESCORT-1 (endosomal sorting complex required for transport-1) or ALIX (ALG-2 interacting protein X, also known as programmed cell death 6 interacting protein or PCD6IP), leads to a similar phenotype (Carlton and Martin-Serrano 2007; Morita et al. 2007). These studies suggest that these interactions are essential for somatic cell abscission (Carlton and Martin-Serrano 2007; Morita et al. 2007; Carlton et al. 2008).

FORMATION OF THE STABLE INTERCELLULAR BRIDGE

Knockout of mouse Tex14 showed that the formation of stable intercellular bridges is essential for spermatogenesis and fertility (Greenbaum et al. 2006). However, until quite recently, it remained unclear how Tex14 participated in stable intercellular bridge formation to prevent abscission and the completion of cytokinesis in male germ cells.

CEP55 was identified as a component of stable intercellular bridges, and is perfectly colocalized with Tex14 as ring-shaped intercellular bridge structures in germ cells throughout
the seminiferous tubules at all stages of spermatogenesis. The bridge diameter expands from the juvenile to the adult stage, suggesting that additional factors such as CEP55 and TEX14 are added to the stable intercellular bridge during spermatogenesis. In the female, CEP55 is expressed and colocalized with TEX14 in the embryonic 18.5-day-old mouse ovary.

In somatic cells, abscission requires CEP55, ALIX, and TSG101 (a component of the ESCRT complex). CEP55 is recruited from the centrosome to the midbody and interacts with MKLP1 (Fabbro et al. 2005; Martinez-Garay et al. 2006; Zhao et al. 2006). Furthermore, CEP55 forms homodimers through coiled-coil interactions, and subsequently, ALIX and TSG101 are recruited to the midbody by binding the “hinge” region of CEP55 (Fig. 4, lower panel) (Martinez-Garay et al. 2006; Carlton and Martin-Serrano 2007; Morita et al. 2007). Glycine (G)-proline (P)-proline (P)-X-X-X-tyrosine (Y) (GPPX3Y) motifs in ALIX and TSG101 are critical for this interaction with the “hinge” region of CEP55 (Morita et al. 2007; Lee et al. 2008). CEP55 was also identified as a TEX14 interactor (Iwamori et al. 2010). Nineteen TEX14 orthologs contain a GPPX3Y, which mimic the ALIX and TSG101 interactions with CEP55 (Iwamori et al. 2010; Wang et al. 2001; Wu et al. 2003). Although there is significant divergence of these orthologs in their carboxy-terminal regions compared to the
ankyrin repeats (ANK) and kinase-like (KL) domains in their amino termini, 17 of the 19 proteins share a similar GPPX3Y motif in their carboxy termini, with the exceptions being the *Xenopus laevis* and dolphin orthologs. In addition, several additional amino acids are conserved between TEX14 and other GPPX3Y-containing proteins including an alanine upstream of the GPPX3Y motif (also present in human ALIX and TSG101), serine/threonine/alanine within the motif, and two prolines downstream (one also present in TSG101). We hypothesize the GPPX3Y motif of TEX14 binds strongly to CEP55 to block similar GPPX3Y motifs of ALIX and TSG101 preventing their interaction with CEP55 and localization to the midbody (Fig. 4, upper panel). In particular, the first proline and the tyrosine of the GPPX3Y motif were sufficient for this portion of TEX14 to interact with CEP55. The exogenous expression of TEX14 in cell culture precludes ALIX from the midbody and produces interconnected somatic cells. By binding endogenous CEP55 and blocking ALIX and TSG101 from interacting with CEP55, TEX14 inhibits the completion of cytokinesis and stabilizes the transient intercellular bridge. TEX14:CEP55 complexes are critical for the formation of stable intercellular bridges.

In general, abscission can be described as MKLP1 → CEP55 → ALIX/TSG101 → → → abscission; however, in differentiating male (and presumably female) germ cells, the steps involved appear to be MKLP1 → TEX14 → CEP55 → intercellular bridge. The fact that TEX14 appears early in the process of cytokinesis bound to MKLP1 and filling the intercellular bridge (Greenbaum et al. 2007) may allow TEX14 to compete for CEP55 more effectively. In testes of TEX14 null mice, CEP55 localizes to the midbody by binding MKLP1 and, in the absence of local TEX14, its interaction with ALIX and TSG101 is permitted. Complete cytokinesis occurs, disrupting the transient intercellular bridge, and resulting in an eventual failure of spermatogenesis during meiotic prophase (Greenbaum et al. 2006).

**OTHER MAMMALIAN INTERCELLULAR BRIDGE FUNCTIONS**

Recently our group identified a second testis-abundant component of the mammalian intercellular bridge, RBM44 (RNA binding motif 44), through its association with TEX14 (Iwamori et al. 2011). RBM44 is not, however, a known component of the midbody, but RBM44 orthologs are present in placental and marsupial mammals, birds, and fish. Distinct from TEX14, RBM44 does not localize to all intercellular bridges, but only to those in meiotic germ cells. Based on the presence of an RRM (RNA recognition motif) domain in RBM44 and robust immunostaining of spermatocyte cytoplasm in addition to the bridge, the protein may serve to transport mRNA, including across intercellular bridges. Furthermore, targeted disruption of *Rbm44* results in an unexpected increase in sperm number without the loss of intercellular bridges. Therefore, the mechanistic role of RBM44, unlike the CEP55-TEX14 interaction, is not to physically stabilize the bridge but may be to eliminate select meiotic germ cells, perhaps as a quality checkpoint. Because physiologic apoptosis of spermatocytes predominantly occurs during the meiotic divisions, potentially by activation of the spindle checkpoint, it is notable that the peak of RBM44 staining occurs during stage XII of spermatogenesis when the meiotic divisions occur. Alternatively, the lack of infertility in RBM44 null mice may be due at least partially from redundancy with the more than 100 RRM domain-containing proteins; however, these possess little similarity to RBM44 outside the RRM domain. The RRM domain of RBM22 has the highest identity to that of RBM44, with those of RBMX, RBMXL2, RBMXRT, RBMY1A1, RBM4, RBM4B, RBM14, RBM15B, BRUNOL6, and DAZAP1 only slightly less identical. Tissue microarray data suggest that some of this group may be testis-expressed (e.g., RBM4B, RBM14) or testis-specific (e.g., RBMXL2) (UCSC genes http://genome.ucsc.edu/). RBM22 mRNA appears to
increase over ovarian follicular development, and RBM4 and RBM14 are expressed in male germ cells, with distinct kinetics (Mammalian Reproductive Genetics database http://mrg.genetics.washington.edu/). Intriguingly, RBM22 interacts with ALG-2 (Montaville et al. 2006), a binding partner of midbody protein ALIX, so TEX14 interactions with RRM domain-containing proteins could act to redundantly block ALIX entry into the midbody by a CEP55-independent mechanism.

CONCLUDING REMARKS

Additional questions remain regarding mammalian stable intercellular bridges. TEX14 knockout studies strongly support the “critical stage” hypothesis (Fig. 2C), but do not exclude the “synchronization” or “phenotypically diploid” hypotheses. The particular functions contributed by intercellular bridges essential to completion of meiotic prophase in spermatocytes have yet to be elucidated. Intercellular bridges are not essential to ovarian folliculogenesis, but whether they contribute to oocyte function or act redundantly with oocyte-specific pathways is unclear. To ensure that germ cell intercellular bridges remain stable in the face of environmental insults, additional protein interactions may be required in addition to the TEX14:CEP55 interaction, such as between TEX14 and other known or novel bridge components. Exclusion of abscission components from the midbody by the TEX14:CEP55 interaction may also indirectly affect other abscission events, such as those involving the ESCRT complex. Because TEX14 exerts a large role in the formation of stable intercellular bridges and interacts with general abscission components, we believe that characterization of the key molecular mechanisms underpinning this germ cell–specific variation on abscission will facilitate drug therapy targeted at disrupting stable intercellular bridges in germ cells for male contraception. Conversely, a small molecule mimic of TEX14 might be applicable as a cancer therapeutic targeted at blocking abscission in continuously proliferating cancer cells.

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REFERENCES


M.P. Greenbaum et al.


deo Rj DG, Russell LD. 2000. All you wanted to know about spermatogonia but were afraid to ask. J Androl 21: 776–798.


Robinson DN, Cooley L. 1997. *Drosophila* kelch is an oligomer- 
Robinson DN, Cant K, Cooley L. 1994. Morphogenesis 
of *Drosophila* ovarian ring canals. *Development* 120: 
2015–2025. 
Roulier EM, Panzer S, Beckendorf SK. 1998. The Tec29 tyro-
sine kinase is required during *Drosophila* embryogenesis 
and interacts with Src64 in ring canal development. *Mol 
Saez JM, Perrard-Sapori MH, Chatelain PG, Tabone E, 
Rivarola MA. 1987. Paracrine regulation of testicular 
Sertoli E. 1877. Sulla struttura dei canalicoli seminiferi dei 
testicoli studiata in rapport allo sviluppo dei nemas-
Somogyi K, Rorth P. 2004. Cortactin modulates cell migra-
tion and ring canal morphogenesis during *Drosophila* 
Stanley HP, Bowman JT, Romrell LJ, Reed SC, Wilkinson RE. 
1972. Fine structure of normal spermatid differentiation 
Tegelenbosch RA, de Rooij DG. 1993. A quantitative study of 
spematogonial multiplication and stem cell renewal in the 
Tilney LG, Tilney MS, Guild GM. 1996. Formation of actin 
filament bundles in the ring canals of developing *Dro-
Touré A, Dorseuil O, Morin L, Timmons P, Jegou B, Reibel L, 
Gacon G. 1998. MgcRacGAP, a new human GTPase-acti-
vating protein for Rac and Cdc42 similar to *Drosophila* 
rotundRacGAP gene product, is expressed in male 
Van de Putte T, Zweijen A, Lonnøy O, Rybin V, Cozijnens M, 
trap vector insertion in mgcRacGAP die during pre-
Ventela S, Toppari J, Parvinen M. 2003. Intercellular organ-
elle traffic through cytoplasmic bridges in early sperma-
tids of the rat: Mechanisms of haploid gene product 
von Ebner V. 1888. Zur Spermatogenese bei den Saugethi-
Witkin JW, O’Sullivan H, Silverman AJ. 1995. Novel associ-
ations among gonadotropin-releasing hormone neurons. 
*Endocrinology* 136: 4323–4330. 
gene 14 (Test14): A gene encoding a protein kinase pref-
erentially expressed during spermatogenesis. *Gene Expr 
Patterns* 3: 231–236. 
ring canal formation during *Drosophila* oogenesis, en-
Zhao WM, Seki A, Fang G. 2006. Cep55, a microtubule- 
bundling protein, associates with centralspindlin to con-
trol the midbody integrity and cell abscission during 
Germ Cell Intercellular Bridges

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