Fine structure of *Drosophila* larval salivary gland ducts as revealed by laser confocal microscopy and SEM

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**Key words.** Diptera, Drosophilidae, *Drosophila*, larval salivary glands, secretory glue, salivary duct, taenidia, ultrastructure

**Abstract.** The functions of the larval salivary glands (SGs) of *Drosophila* are traditionally associated with the production of a massive secretion during puparium formation; it is exocytosed into a centrally located lumen and subsequently expectorated via ducts, the pharynx and mouth. This so-called proteinaceous glue serves as an adhesive to attach the puparial case to a solid substrate. Great attention has been paid to the secretory cells of SGs, which are famous for their giant polytene chromosomes. However, substantially less attention has been devoted to individual or common ducts that form the most proximal portion of the SG organ via which the glue is released into the pharynx. In the present paper, we describe the organization and fine structure of the taenidia, highly specialized circumferential ring-like extracellular (cuticular) components on the internal side of these tubes. Two chitin-specific probes that have previously been used to recognize taenidia in *Drosophila* tracheae, Calcofluor White M2R (also known as Fluorescent Brightener 28) and the novel vital fluorescent dye SiR-COOH, show positively stained ductal taenidia in late larval SGs. As seen using scanning electron microscopy (SEM), the interior of the ductal tube contains regular and densely-arranged ridge-like circumferential rings which represent local thickenings of the cuticle in various geometries. The microtubular arrays that optically colocalize with taenidia in both the trachea and SG ducts are specifically and strongly recognized by fluorescently-conjugated colchicine as well as anti-tubulin antibody. In contrast to taenidia in the tracheae, the analogous structures in SG ducts cannot be detected by fluorescently-labeled phalloidin or even actin-GFP fusion protein, suggesting that the ducts lack a cortical network made of filamentous actin. We speculate that these taenidia may serve to reinforce the duct during the secretory processes that SGs undergo during late larval and late pupal stages.

**INTRODUCTION**

The larval salivary glands (SGs) of *Drosophila melanogaster* represent a unique model organ. They are best known for their giant polytene chromosomes, the detailed analysis of which led to several important advances, including establishment of the first cytogenetic maps and the elaboration of chromomere theory (Painter, 1933; Bridges, 1935; Pelling, 1966; Ashburner, 1970; Lefevre, 1976; Sorsa, 1989; Lindsley & Zimm, 1992). Furthermore, the degree of polyteny of SG chromosomes has made it possible to study the puffs which represent active gene sites. The main function of the larval SG appears to be the production of a mucinous glue, which serves as an adhesive during puparium formation (Fraenkel & Brookes, 1953). In anticipation of this function, at about the middle of the third-instar, globular, highly refractile PAS-positive granules appear in the cytoplasm of SG cells (Berendes, 1965; Kolesnikov & Zhimulev, 1975). The hormone ec dysone plays a dual role in this process, by both starting and then terminating a specific developmental program of protein synthesis required for the production of these secretory granules. Upon a dramatic increase in the ec dysone titer a few hours prior to pupariation, the contents of the granules are released by exocytosis into the SG lumen, which is then emptied by expectoration during puparium formation (Lane et al., 1972; von Gaudecker, 1972; Berendes & Ashburner, 1978; Farkaš & Šuťaková, 1998) where it serves as a glue to attach the puparial case to a solid substrate (e.g. vial glass wall). Thus, these granules constitute the components of the salivary glue secretion (Sgs). The Sgs represents a highly specialized and unique extracellular composite bioadhesive secretion, the individual protein components of which are encoded by a series of eight Sgs genes (Korge, 1975, 1977a, b; Ashburner & Berendes, 1978; Lehmann, 1996; Farkaš, 2016) that are known to form a set of so-called interec dysial chromosomal puffs.
Individual Sgs-genes at these cytological bands have been subsequently cloned and molecularly characterized (reviewed in Farkaš, 2016; Da Lage et al., 2019).

In the course of our SEM studies on the surface properties and internal substructure of exoyctosed and expectorated glue (Beňová-Liszková et al., 2019b), we found that the internal lining of the tubular, non-secretory structure of the SG, i.e., the duct that develops during embryogenesis by the coordinated action of a differently expressed set of genes, in contrast to the SG proper (Kuo et al., 1996; Jones et al., 1998; Yao & Sun, 2005), shows peculiar structures resembling tracheal taeniaid. Using classical light microscopy, the potential presence of taenial-like chitinous circumferential rings in Drosophila SG duct was noticed by Robertson (1936) and Ross (1939). Their work was cited by Bodenstein (1950) and by Berendes & Ashburner (1978). Then, there was an informational gap on taeniaid until Haberman et al. (2003) studied the embryonic specification of cell fates within the salivary gland primordium, including salivary ducts, where they observed the formation of taeniaid-like structures during the embryogenesis of D. melanogaster; however, no additional attention was paid to the SG ducts. The taeniaid are distinctive to the insect tracheal system, and are defined as circumferential thickening of the cuticle inside a trachea or tracheole where they show various geometries such as loops or spirals (Snodgrass, 1935; Richards, 1951; Eidmann & Kühllhorn, 1970; Wigglesworth, 1972; Mill, 1972, 1998; Noirot & Noirot-Timothée, 1982; Manning & Krasnow, 1993). Using laser confocal microscopy and SEM we show herein that the larval SGs of Drosophila contain microtubule-lined taeniidial structures inside the ductal tubes, similar to those present in the tracheae, but in which filamentous actin is conspicuously absent.

**MATERIALS AND METHODS**

**Abbreviations**

FITC – fluorescein isothiocyanate; GFP – green fluorescent protein; HMDS – hexamethyldisilazane; PAS – periodic acid-Schiff stain; PBS – phosphate-buffered saline; (PBT) – PT + bovine serum albumin; HMDS – hexamethyldisilazane; PAS – periodic acid-Schiff stain; PBS – phosphate-buffered saline; (PBT) – PT + bovine serum albumin (PBT; Serva) and 2% goat serum (PBTS; Sigma). To detect microtubules, primary rabbit anti-β-Tubulin polyclonal antibody (Sigma # T-3526) diluted 1:200 was added, before incubating overnight at 4°C in the presence of PBTS. To detect the borders (membranes) of individual ductal cells, primary mouse anti-Fasciclin III monoclonal antibody (DSHB # 7G10) diluted 1:10 was used. After intensive washing in PT, goat affinity purified and preadsorbed Cy5-anti-mouse F(ab)_2-specific and Cy3-anti-rabbit F(ab)_2-specific secondary antibodies (Jackson ImmunoResearch Labs.) was added before incubating for 2 h at room temperature (23–25°C). As an alternative protocol to detect microtubules, FITC-labeled colchicine (Molecular Probes/Invitrogen) was diluted 1:200 before incubating for 2 h at room temperature. After intensive washing in PT, the salivary glands were mounted in Dabco-supplemented (Sigma) Elvanol under a Schott high-performance No 1.5H coverslip, sealed with Picodent nontoxic duplicating silicone or nail polish, and immediately viewed under a Zeiss LSM-510 META laser confocal microscope using a 40× (NA 1.3) oil objective lens. Collected Z-stack images were deconvolved using Zeiss Axiovision 4.8 or SVI Huygens Professional 4.3 software, rendered at maximum intensity, and further processed in Adobe Photoshop.

To detect chitin (a major component of taeniaid) in fixed tissue, SGs (or tracheae) were stained with Calcofluor White M2R (also known as Fluorescence Brightener 28, FB28; Sigma, # F-3543) at pH 9 in 0.2 M Tris buffer containing 0.1% Triton X-100 (Serva) for 2–3 h at room temperature. Whenever necessary, samples were also counterstained with Alexa Fluor 488 or Alexa Fluor 546 phallodin (Molecular Probes/Invitrogen) to detect filamentous actin, as previously described (Farkaš et al., 2015, 2018).

For live cell in vivo imaging of salivary ducts, freshly dissected salivary glands were uploaded for 1 h in 1 μM SiR-COOH or 1 μM 580CP-LTX at room temperature. SiR-COOH and 580CP-LTX are novel silicon-rhodamine and carbophyronine-based fluorescent dyes (Lukinavičius et al., 2018). After incubation, the SGs were briefly rinsed in three volumes of Drosophila saline solution, mounted in Drosophila Ringer containing 20% glycerol under a Schott high-performance No 1.3H coverslip, sealed with Picodent nontoxic duplicating silicone or nail polish, and immediately viewed under a Zeiss LSM-510 META or Leica TCS SP8 STED-3X laser confocal microscope using 40× (NA 1.3) and HCX-PL-AP0 100× (NA 1.4) oil objective lenses, respectively. Collected Z-stack images were deconvolved using Zeiss Axiovision 4.8 or SVI Huygens Professional 4.3 software, rendered at maximum intensity, and further processed in Adobe Photoshop.

Image fusion of widefield light or phase contrast microscopic pictures was performed by focal plane merging of selected Z-stacks via edge detection and Fourier analysis in order to obtain a greater depth of field focus in the final blended image.

**Light and phase contrast microscopy**

Salivary gland ducts and tracheae were also imaged under Leica DMRE and Leica DM6000B microscopes equipped with Leica DFC320 digital color cameras controlled by Leica LAS 2.6.0 software. Image fusion of widefield light or phase contrast...
microscopic pictures was performed by focal plane merging of selected Z-stacks via edge detection and Fourier analysis in order to obtain a greater depth of field in the final blended image.

**Sample processing for SEM**

Whole salivary glands were fixed in 4% paraformaldehyde (Polysciences) + 2% glutaraldehyde (Ted Pella) in 0.1 M sodium cacodylate (pH 7.2; Serva) for 1 h at 23–25°C; the samples were placed in inverted caps from 1.5 ml eppendorf tubes and approximately 100 μl of fixative was used per 10 pairs of glands to ensure that they were fully covered. The SGs were then rinsed five times (5–10 min per rinse) in 100–200 μl of clean 0.1 M sodium cacodylate (pH 7.2) at 23–25°C. Samples were postfixed in 0.5% osmium tetroxide (EMS) in ddH2O for 30 min at 23–25°C; and then extensively rinsed in ddH2O (minimum 6 times for 10 min each). SGs were dehydrated in an ascending series of ethanol (30%, 50%, 70%, 96% and 100%; Merck); the dehydration step with 100% ethanol was repeated twice before applying a 1 : 1 mixture of 100% ethanol + 100% acetone (Merck) twice, followed by three changes of 100% acetone. At this step it was important that sample did not dry completely during the exchanges of absolute ethanol or acetone, or at any subsequent time before critical point drying. Therefore, the addition of hexamethyldisilazane (HMDS; Sigma-Aldrich) in place of Peldri II to facilitate critical point drying (Běhová et al., 2007; Běhová-Liszko et al., 2019a), was performed carefully in several steps: the first volume of HMDS was applied in the presence of small remnants of acetone; then, after 5–10 min, the HMDS was quickly removed and fresh HMDS added, taking care that the previous HMDS did not dry up; finally, SGs were kept in HMDS for 30 min, and then the remnants of the HDMS were allowed to evaporate completely. The process of HMDS evaporation has a tendency to generate a slight electrostatic potential on the surface of samples, thus the last step was performed in a very clean air environment (e.g. under a desiccator) to minimize the attraction of dust particles.

Dried SGs were mounted on pieces of double-sided tape (Scotch) on 16 or 24 mm aluminum SEM stubs and then sputtered for 2.5 min with gold-palladium using a Balzers sputter coater device SCD-030 at 35–40 mA per stub and at a pressure of 0.05 to 0.1 mbar. This produced a 40–50 nm continuous alloy layer. Samples were viewed and photographed using an FEI Quanta FEG250 scanning electron microscope with the emission field cathode at an accelerating voltage of 10 kV. The bitmap images obtained were processed and labeled using Adobe Photoshop or Corel Draw software and assembled into figures using Adobe Photoshop.

**RESULTS**

**Tubular ducts of salivary glands**

The *Drosophila* larval SG is a paired organ which is split into the SG proper and the tubular duct (Fig. 1). In contrast to the widely studied SG proper, little attention has been paid to duct morphology; to our knowledge its histochemical properties and SEM ultrastructure have not been documented. The tubular duct is the last portion of the salivary gland system through which Sgs-glue must pass to reach the pharynx and mouth, where it is expectorated into the environment. The larval duct is composed of about 55 cuboidal epithelial cells with polytene nuclei that form simple tubes. These tubes connect the secretory cells to the larval mouth by a Y-shaped tubular conduit (Berendes & Ashburner, 1978; Abrams et al., 2003). The ducts of the left and right gland unite to form a common duct opening into the floor of the pharynx. This is the only structure of *Drosophila* SGs which has not been studied in more detail throughout the long history of SG research.

While investigating SGs for another purpose under SEM, we found that the interior of the tubular duct contains regular and densely-arranged ridge-like circumferential rings or taenidia. These represent local thickenings of the cuticle in various geometries such as loops, rings, or helices, as would normally be found in the tracheal system (Fig. 2a–d). The thickness of the individual taenidia in SG ducts is between 280 and 690 nm (Fig. 2e, f), similar albeit not identical to that of tracheal taenidia (400 to 760 nm) from identically aged third-instar larvae, with the distance between taenidia also showing some level of similarity (Fig. 2g, h). However, from these images it is also clear that, unlike in tracheae, taenidia in SG ducts are rough and ladder-like with irregularly-sized randomly distributed side ramifications protruding into the spaces between them. This is in contrast to the smoother surface of tracheal taenidia which have spaces filled with trabecular structure resembling a dense arterial network. Interestingly, on the basal surface of the SG duct (under basal lamina) we observed circular, rope-like surrounding fascicles 250 to 350 nm in diameter (Fig. 2i, j) which we did not find in tracheae. In this study we were not able to discern their nature and composition.
Distribution of cytoskeletal and chitin probes in SG ducts

After initial inspection under wide field light and phase contrast microscopes, which also revealed dense taenidial-like rings inside SG ducts of late third instar larva (Fig. 3a), we tested two chitin-specific probes, that have previously been used to recognize taenidia in Drosophila tracheae, Calcofluor White M2R (also known as Fluostain or Fluorescent Brightener 28) and the novel vital fluorescent dye SiR-COOH (Lukinavičius et al., 2018). Fig. 3b shows positively stained ductal taenidia in late larval SGs of Drosophila using SiR-COOH. The chitinous taenidial character of these structures was further confirmed by their strong/specific fluorescent staining using Calcofluor White M2R (Fig. 3c). In both cases, tracheal tubules stained in parallel with SGs under the same conditions were used as positive controls (Fig. 3d, e). In contrast to tracheae, the intracellular actin lining parallel to taenidia in SG ducts is absent (Fig. 3f), but the chitin signal overlaps strongly with intracellular tubulin recognized by both fluorescently-labeled colchicine and anti-tubulin antibody (Fig. 3h, i).
The distribution of fasciclin-III demarcates the limits of individual ductal cells (Fig. 3g). Identical parallel staining of tracheae (Fig. 3k–n) revealed that, in contrast to SG ducts, tracheal phalloidin-bound F-actin bundles are highly visible (Fig. 3k) and their signal clearly overlaps with the position of microtubules (Fig. 3m) and Fluostain-positive chitin (Fig. 3n). The merge of all four channels (Fig. 3j and 3o) highlights the fact that microtubular and chitinous signals overlap significantly in both tissues. In addition, in Fig. 3o the overlap of filamentar, microtubular and chitinous signals in the tracheal trunk is highly apparent.

In order to provide a more detailed view of the SG ducts, some of these vital-stained as well as fixed and FB28-labelled taenidia are shown at a higher magnification and with a monochrome regime in Fig. 4. This finding was intriguing, since previous reports described the presence of F-actin bundles parallel to taenidia not only in embryonic SG ducts, but also in ducts of second-instar larvae (Haberman et al., 2003). Their absence in late third-instar larvae was therefore unexpected. To clarify this observation, we decided to verify the binding of the fluorescent probe, phalloidin, to typical locations for filamentous actin such as the cortical cytoskeleton and exocytotic vesicles within the corpuscular secretory cells of SGs. The staining protocol was identical to that of the duct cells, and several observations of both secretory and duct cells were performed from the same microscope slides. As documented in Fig. 5a–d, both Alexa Fluor 488 phalloidin and Alexa Fluor 546 phalloidin bind specifically and strongly to both the cortical cytoskeleton and exocytotic vesicles. Thus, the absence of visible F-actin rings that encircle the entire SG duct was not an artifact or error in the protocol.

Detection of actin-GFP fusion protein in SG ducts

In the next step we used act-GFP transgenic flies expressing actin-GFP fusion protein ubiquitously in most if not all tissues during development. Out of 5 tested actin-GFP fusion lines, P[w/+mC] = ActGFP]JMR3 expresses GFP the most intensely in the SGs of late third-instar larvae and fluorescence could be detected not only in vivo but also after paraformaldehyde fixation. Therefore, this line was used to observe the GFP signal in SG ducts. We found that GFP, either in living SGs or in fixed organs, produces barely visible fluorescence without F-actin rings inside the duct (Fig. 5e, f). This indicates that even the natural expression and distribution of actin-GFP fusion protein does not occupy the positions of the F-actin rings visible in younger stages.

In addition, we also used the UAS-Act5C-GFP line, the expression of which was under Sgg3-Gal4, dri-Gal4 or hs-Gal4 drivers, to enhance the abundance of Actin-5C-GFP.
fusion protein in salivary gland cells. To obtain the most reliable data, we collected a series of at least 50 Z-stack optical sections (300 nm each); after deconvolution we performed 3D reconstruction rendering using transparency mode of Zeiss AxioVision software. As shown in Fig. 5g and 5h, neither the expression of Sgs3-Gal4 nor dri-Gal4 led to any change in the Actin-5C-GFP signal for the SG ducts of late third-instar larvae. Due to certain properties of Sgs3-Gal4 and dri-Gal4 discussed below, we decided to also use a hs-Gal4 construct to drive the expression of UAS-Act5C-GFP ubiquitously. This was achieved by performing a 1-h heat shock of 30°C at the early wandering stage of third-instar larvae, and checking for a GFP signal 2 h later. Under this condition we were able to detect a faint F-actin signal in the SG duct, which resembled bundles of filaments parallel to taenidial rings (Fig. 5i). However, it
should be noted that the structures shown in Fig. 5i became visible only after deconvolution and 3D rendering.

**DISCUSSION**

Since their first mention by Robertson (1936) and slightly more detailed description by Ross in 1939 and her camera obscura drawings of the SG duct, there were very few references to or documentation of the circumferential chitinous lining (taenidia) in larval SGs of *Drosophila melanogaster* for decades. In fact, the presence of duct taenidia was not documented until the late 1970s by Berendes & Ashburner (1978) who used light microscopy on the SGs of larval *D. hydei*. Several decades later, Haberman et al. (2003) studied the embryonic specification of cell fates within the salivary gland primordium, including salivary ducts, where they observed the formation of taenidia-like structures during the embryogenesis of *D. melanogaster*. The existence of taenidia inside the ducts of adult *Drosophila* salivary glands was only noticed by Miller (1950), although adult glands were never the subject of such intense studies as was the case for the larval organ. Furthermore, adult SGs are a completely different organ to larval glands whose polytene cells histolyse during metamorphosis. The taenidia are distinctive of the tracheal system, and are defined as circumferential annular thickenings of the cuticle inside a trachea or tracheole where they show various geometries such as loops or helices (Snodgrass, 1935; Richards, 1951; Eidmann & Kühlhorn, 1970; Wigglesworth, 1972; Mill, 1972, 1998; Noirot & Noirot-Timothee, 1982; Manning & Krasnow, 1993). From our confocal and SEM observations it is apparent that larval SGs of *Drosophila* contain highly similar, if not identically composed taenidial structures inside the ductal tubes, and most probably serve as the primary source of its scaffolding structural integrity and reinforcement as in trachea.

Some of the mechanical properties of ducts, and notably taenidia, could be more appropriately assessed if we know, for example, the composition and distribution of resilin and sclerotin. Resilin, in particular, is known to be responsible for the deformability and flexibility in the membrane and joint systems of insects (Michels, 2007) and could therefore provide valuable information. Unfortunately, resilin antibody (Burrows et al., 2011; Wong et al., 2012) has not been available for at least the last four to five years, and sclerotin is a biochemically variable multiprotein complex against which no antibody can be generated since it incorporates different proteins in varying proportions in different body structures. We considered the autofluorescent technique described by Zill et al. (2000) to detect resilin or sclerotin, which has been subsequently applied to insects and some other arthropod exoskeletons (Klaus et al., 2003; Schawaroch et al., 2005; Klaus & Schawaroch, 2006; Michels & Gorb, 2012; Klaus et al., 2014); however, despite our repeated trials we found that this does not
work for ductal structures. This can be explained not only by the much softer nature of the SG ducts, which contain less resilin and sclerotin than the extremely hard and thick exoskeleton of adult beetles, crickets and grasshoppers, but mainly by the fact that sclerotin protein composition can be unique to these exoskeletal structures, which allows for bright or at least usable autofluorescence.

Fig. 4. Monochromatic and detailed (large-scale) presentation of salivary gland ducts and tracheae of Drosophila larvae. (a) Monochromatic presentation of the vital staining of the SG ducts with novel fluorescent probe SiR-COOH. (b) Monochromatic presentation of the vital staining of a segment of the main thoracic left dorsal tracheal tube from the same larva as in Fig. 3d with SiR-COOH. (c) Monochromatic presentation of taenidial structures in SG duct detected specifically by Fluostain (FB28) in a paraformaldehyde-fixed sample. (d) Monochromatic view of the segment of the main thoracic left dorsal tracheal tube stained by Fluostain (FB28) in a paraformaldehyde-fixed sample. (e) Detailed view of diffused signal of the filamentous actin in SG duct detected using Alexa Fluor 488 phalloidin. (f) Detailed view of apparent taenidial ridges in the same SG duct detected by anti-β-Tubulin polyclonal antibody. (g) Detailed view of clear taenia inside SG duct as detected by chitin-specific probe Fluostain (FB28). (h) Detailed view of the pattern of filamentous actin in main dorsal left tracheal trunk corresponding with the arrangement of taenidial circumferential rings. (i) Detailed view of the pattern of microtubules in main dorsal left tracheal trunk, detected by anti-β-Tubulin polyclonal antibody, and highlighting circumferential rings of taenia. (j) Detailed view of tracheal taenia detected directly as chitin recognized by the Fluostain (FB28) probe. Scale bar = 20 μm for (a) to (g), and scale bar = 7 μm for (h) to (j).
Although the fluorescent signal emitted from the binding of SiR-COOH or Fluostain to ductal taenidia is weaker than the signal from tracheal tubules, meaning that they may contain less chitin, these ductal taenidia can provide sufficient strength and flexibility to the duct. Importantly, the taenidia may prevent collapse of the ductal tubule during ecdysis in a similar way as for tracheae (Matusek et al., 2006; Hannezo et al., 2015), at least during the larval stages. The common duct of the SGs opens in the floor of the muscular pharynx, the portion of anterior armature chiefly involved in foraging (Sokolowski et al., 1997), which is the permanent locomotor activity of the larval foregut since hatching of the first-instar larva until the prepupa. This continuous muscular activity requires highly elastic connections that are able to resist mechanical tension; duct taenidia meet this specification. Moreover, there is a good reason to believe that taenidia also prevent collapse of the duct during glue expectoration. This is a time when the larva undergoes very strong extracardiac pulsations executed by the abdominal musculature, generating high internal hemolymph pressure to facilitate body contractions at the onset of pupariation (Žďárek & Fraenkel, 1972; Žďárek, 1985). These hemolymph pressure pulsations in Sarcophaga flies and some other holometabolic insects reach 100 to 300 mm Hg (10 to 40 kPa) of internal pressure during pupariation (Sláma, 1976, 1984, 2000; Žďárek et al., 1979; Žďárek, 1985). We estimate that in the much smaller Drosophila, the immobilization and contraction phase of the pupariation would result in 2 to 5 kPa of internal hemolymph pressure, still significantly above atmospheric level. Many internal organs, including salivary glands and their duct, must therefore possess the ability to resist this distort-

Fig. 5. Filamentous actin distribution in larval salivary glands and SG ducts of Drosophila. (a) Alexa Fluor 488 phalloidin and (b) Alexa Fluor 546 phalloidin detection of cortical actin in pre-wandering third-instar larval SGs. Arrows point to typical membrane-associated cortical locations, and the white arrowhead indicates a strong signal of filamentous actin on the apical membrane lining of the SG lumen. (c) Alexa Fluor 488 phalloidin and (d) Alexa Fluor 546 phalloidin detection of secretory vesicles in wandering stage third-instar larvae during active exocytosis of Sgs-glue granules into the lumen. Process of vesicle docking and priming to apical membrane prior to exocytic event is associated with a strong F-actin signal. White arrows point to docked secretory granules. (e) Actin-GFP signal in living SG ducts mounted in PBS with 20% glycerol under coverslip, and (f) actin-GFP signal in paraformaldehyde-fixed prep mounted under Elvanol. In both cases the signal is relatively weak and does not show structures of the circumferential ring lining. (g) GFP signal of UAS-Act5C-GFP transgenic construct driven by Sgs3-Gal4, while (h) is the same construct expressed under the control of dir-Gal4. (i) GFP signal of UAS-Act5C-GFP transgenic construct induced temporarily by a short heat shock (30°C) via hs-Gal4 driver where a faint signal of F-actin in the SG duct resembling bundles of filaments parallel to taenidial rings can be seen. (j) Grayscale image of (i) to provide better visibility of the ring bundles of F-actin indicated by white arrows on the both individual and common ducts. Scale bar in (c) and (d) = 10 μm, scale bar elsewhere = 20 μm.
Having provided structural and histochemical information on the identity of taenidia inside SG ducts, the challenge now is to analyze their development and formation using the series of genetic tools available for *Drosophila*. In the SGs of late third instar *Drosophila* larvae we were unable to obtain a clear pattern of the filamentous actin rings that corkscrew around the lumen of the ducts and line or colocalize with the taenidia as described for embryonic SG ducts by Haberman et al. (2003). It has been clearly established in *Drosophila* trachea that F-actin bundle organization is important for embryonic taenidium formation (Chihara et al., 2003; Matuske et al., 2006); the situation in SG ducts may be very similar. Our observations indicate that this early morphogenetic function of F-actin in SG duct tubules may be limited to prefiguring the cuticular ridges during embryogenesis. There are at least two plausible explanations for this. First, actin that is primarily required for prefiguring the cuticular ridges in embryos is subsequently diluted during postembryonic development by simple mechanical expansion of the ductal tube. The tube grows only by volumetric expansion, gradually making filamentous bundles less visible, although they still may persist from the embryonic period. It should be emphasized that the most rapid expansion of the SG duct takes place during the second half of the late third instar, when the SG proper grows significantly due to the huge production of Sgs-secretory granules. At this point, the SGs increase in size more than 20-fold. This could explain why much-less diluted F-actin bundles are detectable in second-instar larvae, as observed by Haberman et al. (2003). Secondly, during larval development and towards metamorphosis, the originally embryonic prefiguring function of F-actin is no longer required and is exchanged or replaced by microtubular scaffold. This is easy to detect by both anti-tubulin specific antibodies and by fluorescently-conjugated microtubule-binding probes (colchicine or paclitaxel). In fact, it is plausible that both of the abovementioned explanations are true and are complementary. The absence of F-actin bundles in third-instar larval SG ducts is in striking contrast to that of third-instar larval tracheae where the optical pattern of actin microfilaments corresponds to the taenidial chitinous ridge pattern or is closely co-located. It needs to be stressed that both actin and tubulin are localized intracellularly, whereas chitinous ridge taenidia are an extracellular matrix (cuticle) forming a supra-cellular structure.

Experiments with *actin-GFP* fusion constructs have shown that expression, either under the control of an endogenous promoter or ectopic drivers (*Sgs3-Gal4* or *dri-Gal4*) does not enhance the detection of ductal F-actin bundles in late third-instar larvae. Despite the fact the *Sgs3-Gal4* driver is very strong and specific to the second half of the third instar, its expression appears to be restricted only to SG proper cells and is absent from ductal cells. The *dri-Gal4* driver has not been used in similar experiments to our knowledge and is derived from the dead-ringer (also known as retained) promoter. Strong expression of the *dri* gene is required for determining SG ducts during *Drosophila* embryogenesis (Bradley et al., 2001; Vining et al., 2005; Kerman et al., 2006), and it is not known if its duct-specific expression is also maintained during postembryonic development. It may be the case that after the determination and full differentiation of embryonic SG ducts is complete, expression of *dri* is downregulated. Interestingly, ectopic overexpression of *actin-GFP* under the control of a *hs-Gal4* driver slightly increased the detectability of circumferential rings of F-actin in the duct cells. The heat shock promoter in the *hs-Gal4* construct is most inducible at 37°C; however, we decided to use a lower temperature (30°C) and for a limited time (60 min) to keep ectopic overexpression at moderate levels. Despite this, it was still sufficient to produce enough ectopic actin-GFP protein to make it visible in ductal cells at locations observed during embryogenesis and early instars. This result provides strong support to our above mentioned conclusion that F-actin in the ductal cells of late third-instar larvae could become diluted during postembryonic development by mechanical expansion of the ductal tube. However, its modest ectopic overexpression can restore it to locations typical of much earlier stages. We purposely did not use a stronger and longer heat shock to avoid ectopic mislocalization of F-actin in inappropriate subcellular locations.

Several genes that are involved in determining tubulogenesis during embryonic development are identical for both the tracheal system and for SGs (Kuo et al., 1996; Henderson & Andrew, 2000; Jin et al., 2001; Kerman et al., 2006). Based on our observations, one can anticipate that at least some of the genes which affect taenidium formation in a tracheal system (Öztürk-Colak et al., 2016) may be different from the genes required to determine and maintain taenia in SG ducts. In addition, the presented observations may indicate a crucial difference between the fate of SGs and the tracheal system: SGs (including their ducts) are ultimately doomed for histolysis, whereas the tracheal system remains functional during metamorphosis, and undergoes extensive remodeling (Whitten, 1980; Manning & Krasnow, 1993; Weaver & Krasnow, 2008; Pitsouli & Perrimon, 2010; Hayashi & Kondo, 2018). The different distribution of F-actin can be an indicator forecasting this fate.

**ACKNOWLEDGEMENTS.** We acknowledge K. Sláma’s advice and helpful discussions on hemolymph pressure pulsations. We are indebted to G. Lukinavičius and S.W. Hell of the Max Planck Institute for Biophysical Chemistry (Göttingen, Germany) for their generous gift of novel fluorescent probes SiR-COOH and S80CP-LTX. We also thank A.J. Andres (University of Nevada, Las Vegas, NV, USA) and Bloomington *Drosophila* Stock Center (Indiana University, Bloomington, IN, USA) for fly
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