Septate Junction Proteins Play Essential Roles in Morphogenesis Throughout Embryonic Development in Drosophila

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ABSTRACT The septate junction (SJ) is the occluding junction found in the ectodermal epithelia of invertebrate organisms, and is essential to maintain chemically distinct compartments in epithelial organs, to provide the blood–brain barrier in the nervous system, and to provide an important line of defense against invading pathogens. More than 20 genes have been identified to function in the establishment or maintenance of SJs in Drosophila melanogaster. Numerous studies have demonstrated the cell biological function of these proteins in establishing the occluding junction, whereas very few studies have examined further developmental roles for them. Here we examined embryos with mutations in nine different core SJ genes and found that all nine result in defects in embryonic development as early as germ band retraction, with the most penetrant defect observed in head involution. SJ genes are also required for cell shape changes and cell rearrangements that drive the elongation of the salivary gland during midembryogenesis. Interestingly, these developmental events occur at a time prior to the formation of the occluding junction, when SJ proteins localize along the lateral membrane and have not yet coalesced into the region of the SJ. Together, these observations reveal an underappreciated role for a large group of SJ genes in essential developmental events during embryogenesis, and suggest that the function of these proteins in facilitating cell shape changes and rearrangements is independent of their role in the occluding junction.

KEYWORDS septate junction morphogenesis head involution dorsal closure salivary glands

Pleated septate junctions (hereafter referred to as SJs) are found in all ectodermally derived epithelia (e.g., epidermis, salivary glands, hindguts, and tracheae) in invertebrates, where they localize basal to the adherens junction (Noirot-Timothee et al. 1978), and function to prevent paracellular flow between the apical and basal sides of an epithelium, much as tight junctions provide a barrier function in vertebrate epithelia (Urakabe et al. 1970; Lord and DiBona 1976). More than 20 genes have been implicated in the establishment or maintenance of SJs in Drosophila (reviewed in Izumi and Furuse 2014). Many of these genes encode membrane proteins with extracellular motifs suggesting a role in cell adhesion. Early in development (stage 12 of embryogenesis, or about 8 hr into the 24 hr embryonic period) most SJ proteins are membrane associated and line the length of the lateral domain. In stage 13 embryos, some SJ protein can be observed in intracellular puncta that colocalize with early and recycling endosomal markers, while the majority of the protein remains localized to the lateral membrane (Tiklova et al. 2010). In stages 14 and 15 SJ proteins are gradually enriched at the apical lateral region, although considerable protein can still be detected along the lateral membrane. At stage 16 (about 14 hr after egg laying), SJ proteins are finally tightly localized to the apical lateral region that defines the SJ. Electron microscopic studies revealed the presence of dispersed electron-dense intercellular septae in wild-type embryos beginning at stage 14, and increasing in number and regularity until an ultrastructurally mature SJ is established in stage 17 (Tepass and Hartenstein 1994). Functional studies revealed that the paracellular barrier is not established until late-stage 15 in wild-type embryos (Paul et al. 2003). The correct organization and function of epithelial SJs requires that each member of the complex is present, suggesting that the SJ is a large, stable, and highly cross-linked protein complex (e.g., Ward et al. 1998; Genova and Fehon 2003).

Mutations that result in the loss of SJs in embryonic epithelia and glia are embryonic lethal, with paralysis due to the loss of the occluding
function at the blood–brain barrier in glia (Baumgartner et al. 1996). Most of the studies characterizing SJ genes note this embryonic lethality, but focus on the cell biological role of the SJ protein in the organization and function of the occluding junction. A few studies, however, have characterized defects in developmental events associated with these mutations. For example, we initially identified Macroglobulin complement-related (Mcr) in a screen for genes involved in imaginal disc morphogenesis during metamorphosis (Ward et al. 2003; Hall et al. 2014). Additionally, zygotic loss of function alleles of coracle (cor) and Neurexin-IV (Nrx-IV), as well as maternal/zygotic mutations in Discs large (Dlg), result in defective dorsal closure (DC) during stages 13–14 of embryogenesis (Perrimon 1988; Fehon et al. 1994; Baumgartner et al. 1996). Furthermore, loss of Fasciclin III (FasIII) results in defective hindgut morphogenesis that normally occurs in stage 13 of embryogenesis (Wells et al. 2013). Finally, mutations in many SJ genes were initially identified as having highly convoluted embryonic trachea, suggesting a requirement for SJ genes in tracheal morphogenesis (e.g., Behr et al. 2003; Wu et al. 2004, 2007; Batz et al. 2014).

Here we set out to determine whether an essential role in embryonic morphogenesis is a common function of all the SJ proteins, or if it rather reflects a pleiotropic effect of a small number of proteins that also happen to function in the establishment or maintenance of SJs. To address this question we undertook a reevaluation of mutations in nine core SJ genes for their role in embryonic morphogenesis. We chose alleles that had been previously characterized for their cell biological role in the SJ, and found that all nine are essential for numerous developmental events during embryogenesis, suggesting that at least a large collection of SJ proteins participates in morphogenetic events shaping the body plan and organs in Drosophila. Detailed examination of salivary gland (SG) organogenesis reveals that SJ proteins are required for both cell rearrangements and cell shape changes that drive the morphogenesis of this tissue. Interestingly, the timing of the developmental events that are perturbed by loss of function SJ mutations precedes the biogenesis of the occluding junction, suggesting distinct roles for these proteins in morphogenesis and in forming the SJ.

**MATERIALS AND METHODS**

**Drosophila stocks**

All Drosophila stocks were maintained on media consisting of corn meal, sugar, yeast, and agar in a room that typically fluctuated between 21.5°C and 23°C. Genetic experiments were conducted in incubators controlled at a constant temperature of 25°C. The following SJ mutant strains were used: Contflr56 (Faire-v-Sarrailh et al. 2004), cor1 (Ward et al. 1998), cor2 (Ward et al. 2001), kunec309 (Nelson et al. 2010), LacG00044 (Llimargas et al. 2004), Mcr1 and McrEY07421 (Hall et al. 2014), Nrg14 (Hall and Bieber 1997), Nrg17 (Paul et al. 2003), nrv2Kc12649 (Buszczak et al. 2007; Hall et al. 2014), Nrx-IVfl94 (Baumgartner et al. 1996), and Tsj2kG01571 (Tiklova et al. 2010). w1118, actin (act)-Gal4, daughterless (da)-Gal4, Df(3R)BSC146, Df(2R)BSC696, Df(2R)BSC305, cor1, FasIII25, kunec309, LacG00044, Nrg14, Nrg17, nrv2Kc12649, Nrx-IVj4304, and Tsj2kG01571 were obtained from the Bloomington Drosophila Stock Center (BDSC, Bloomington, IN). UAS-kun-RNAi (stock 3962), UAS-Mcr-RNAi (stock 100197), and UAS-Nrg-RNAi (stock 107991) strains were obtained from the Vienna Drosophila RNAi Center (VDRC, Vienna, Austria; Dietzl et al. 2007). The Nrg-GFP Fly Trap line G00305 (Morin et al. 2001) and the Nrx-IV-GFP fly trap line CA06597 (Buszczak et al. 2007) were obtained from the FlyTrap consortium (Yale University School of Medicine, New Haven, CT). yk-h-GAL4, UAS-GFP (Wang et al. 2008) was obtained from Arash Bashirullah (University of Wisconsin, Madison, WI). Contfsr56 was obtained from Manzoor Bhat (University of Texas Health Science Center, San Antonio, TX), cor1, cor2, cor3, Mcr1, McrEY07421, nrv2Kc12649, act-Gal4 were balanced with CyO, P[w+, Df(2)EYFP], Nrg14 and Nrg17 were balanced with FM7c, P[w+, Df(2)EYFP], and Contfsr56, kunec309, LacG00044, Nrx-IVj4304, and Tsj2kG01571 were balanced with TM6B, P[w+, Df(2)EYFP] to allow for unambiguous identification of embryos (Le et al. 2006).
in 6% sodium hypochloride, mounted in Hoyer (YFP). Nonhatched embryos 48 hr after egg laying were dechorionated and then selected based upon the absence of Yellow Fluorescent Protein scored on a Nikon Eclipse 80 compound microscope slides, and cleared for 24 hr at 50°C. All cuticular phenotypes were scored on a Nikon Eclipse 80i compound microscope. For the devitellinized cuticles shown in Figure 1, mutant nonhatched embryos 48 hr after egg laying were dechorionated in 6% sodium hypochloride, devitellinized by shaking in a 1:1 mixture of methanol and heptane, expanded by heating at 37°C in 1X phosphate buffered saline plus 0.1% Triton X-100, mounted in CMCP-10 (Polysciences, Inc, Warrington, PA), and cleared overnight at 50°C on microscope slides. The cuticles were imaged using differential interference contrast microscopy (DIC) on a Nikon Eclipse 80i compound microscope equipped with a Photometrics CoolSNAP ES high-performance digital CCD camera. The w<sup>1118</sup> control cuticle was from a first instar larva <30 min after hatch. Then stained with antibodies against α-Spectrin to show the general morphology of the embryo. Anterior is to the left. Note that the Cont embryo has arrested prior to the initiation of germ band retraction, the kune embryo is at late stage 14/early stage 15, and the Mcr mutant embryo is generally disorganized. (E) Quantitation of developmental stages of w<sup>1118</sup> and SJ mutant embryos from a 1 hr embryo collection, subsequently aged 17 hr and then stained with antibodies against α-Spectrin. Note that all 10 SJ mutants had a significantly higher percentage of embryos arrested prior to stage 15 than the w<sup>1118</sup> control (Fisher’s exact test). n, total numbers of embryos scored for each genotype. * P < 0.05, ** P < 0.0001. Scale bars = 100 μm.

Cuticle analysis

Embryos from SJ mutant strains and deficiencies balanced with FM6, P[wm, Dfd-EYFP], CyO, P[wm, Dfd-EYFP], or TM6B, P[wm, Dfd-EYFP] were collected on apple juice agar plates for 1 hr at 25°C, aged for 17 hr and then selected based upon the absence of Yellow Fluorescent Protein (YFP). Nonhatched embryos 48 hr after egg laying were dechorionated in 6% sodium hypochloride, mounted in Hoyer’s medium on microscope slides, and cleared for 24 hr at 50°C. All cuticular phenotypes were scored on a Nikon Eclipse 80i compound microscope. For the devitellinized cuticles shown in Figure 1, mutant nonhatched embryos 48 hr after egg laying were dechorionated in 6% sodium hypochloride, devitellinized by shaking in a 1:1 mixture of methanol and heptane, expanded by heating at 37°C in 1X phosphate buffered saline plus 0.1% Triton X-100, mounted in CMCP-10 (Polysciences, Inc, Warrington, PA), and cleared overnight at 50°C on microscope slides. The cuticles were imaged using differential interference contrast microscopy (DIC) on a Nikon Eclipse 80i compound microscope equipped with a Photometrics CoolSNAP ES high-performance digital CCD camera. The w<sup>1118</sup> control cuticle was from a first instar larva <30 min after hatch. Then stained with antibodies against α-Spectrin to show the general morphology of the embryo. Anterior is to the left. Note that the Cont embryo has arrested prior to the initiation of germ band retraction, the kune embryo is at late stage 14/early stage 15, and the Mcr mutant embryo is generally disorganized. (E) Quantitation of developmental stages of w<sup>1118</sup> and SJ mutant embryos from a 1 hr embryo collection, subsequently aged 17 hr and then stained with antibodies against α-Spectrin. Note that all 10 SJ mutants had a significantly higher percentage of embryos arrested prior to stage 15 than the w<sup>1118</sup> control (Fisher’s exact test). n, total numbers of embryos scored for each genotype. * P < 0.05, ** P < 0.0001. Scale bars = 100 μm.

Immunostaining

Embryos were fixed and processed for antibody staining as described in Fehon et al. (1991). Detailed protocols are available at https://www.protocols.io/view/Immunohistochemistry-Drosophila-Embryocutwwm. Embryos were collected for 1–2 hr or overnight depending on the experiment and aged to the appropriate developmental stage as determined by gut and head morphology. Antibody concentrations for all the primary antibodies used in this study can be found at protocols.io at https://www.protocols.io/view/Optimized-Concentrations-for-Developmental-Studies-daq2dv?guidelines. Antibodies against α-Spectrin, ATPα, Coracle, and DE-Cadherin were obtained from the Developmental Studies Hybridoma Bank (University of Iowa, Iowa City, IA), whereas antibodies against Nrx-IV and Contactin were obtained from Mazoor Bhat (University of Texas Health Science Center, San Antonio, TX), antibodies against Nrg were obtained from Nipam Patel (University of California, Berkeley, CA), and antibodies against Kune were obtained from Mikio Furuse (Kobe University School of Medicine, Kobe, Japan). FITC-labeled antiphosphotyrosine antibodies were obtained from Sigma-Aldrich (St. Louis, MO) and used at 1:800. Secondary antibodies (Jackson ImmunoResearch Laboratories, West Grove, PA) were used at 1:800. Confocal images were acquired on an Olympus FV1000 confocal microscope (Olympus America, Inc., Center Valley, PA) equipped with Fluoview software, a Zeiss LSM510 Meta Laser Scanning confocal microscope (Carl Zeiss Inc, Thornwood, NY), or a Zeiss LSM700 Axiolamager.Z2 confocal microscope. Photomicrographs were cropped, rotated, and adjusted for brightness and contrast in ImageJ. Figures were compiled in Adobe Illustrator (version CS6). The statistical significance for the experiment to address early arrested development (Figure 2) was calculated using a Fisher’s exact test.

Morphometric analysis and nuclear quantification of SGs

Confocal z-series stacks were collected on wild-type and SJ mutant embryos mounted such that lateral views of SGs could be imaged
immediately under the epidermis using a Zeiss LSM700 AxioImager.Z2 with a Plan-APOCROMAT 63X/1.4 Oil DIC lens using a step size of 0.39 μm. Individual images were stacked in Image J to generate a 3-dimensional reconstruction. We used these reconstructions to generate cross-sectional views at three positions roughly equally spaced along the long axis of the gland, and did this for 10 glands per genotype. The number of SG cells per cross-section was determined by counting the number of Hoescht-stained nuclei in three independent cross-sections per gland as described (Chung and Andrew 2014). We also measured the length of two lateral membranes per cell and the intervening apical membrane for a total 10 cells per gland. Statistical significance was calculated using a Student’s t-test.

Data availability
Drosophila strains and antibodies generated in our laboratory (anti-Mcr and anti-Uif) are available upon request. Raw confocal z-series stacks used to generate cross-sectional views used in the morphogenetic analyses described in Figure 5, Table 2, and Table 3 are available upon request. stacks used to generate cross-sectional views used in the morphogenetic analyses described in Figure 5, Table 2, and Table 3 are available upon request.

RESULTS
Loss of zygotic expression of SJ genes results in head involution and dorsal closure defects
We initially identified mutations in Mcr in a screen for genes required for imaginal disc morphogenesis during metamorphosis (Ward et al. 2003; Hall et al. 2014). Coupled with previous studies showing a requirement for the core SJ genes cor and Nrx-IV during dorsal closure (DC) (Fehon et al. 1994; Baumgartner et al. 1996), we wondered whether other SJ genes were required for morphogenesis, and for which developmental events they were required. In this way we could address whether an essential role in morphogenesis was a general feature of SJ developmental events. We initially identifi
can be used to assess the major developmental events shaping the body plan including germ band retraction, head involution, and dorsal closure. The most obvious defect observed for all of these mutations was an inability to complete head involution (Figure 1). We noted a large variation in the expressivity of this phenotype, from a complete inability to invert the head segments (Figure 1, E and F) to minor defects in head skeleton positioning and morphology (Figure 1, K and L). The penetrance of head involution defects in embryos that secreted a cuticle varied from about 46% of cres956/Df(3R)BSC146 to 100% of TsfKG01571 embryos (Table 1). Using the same set of cuticle preparations, we observed similarly variable defects in DC. These phenotypes ranged from puckering of the dorsal epidermis and small cuticular scabs (Figure 1B), to small (Figure 1C) and large dorsal holes (Figure 1D). DC phenotypes were less penetrant than head involution defects, but were nonetheless present at some level in every line examined (Table 1). By observing cuticles from devitellinized embryos, we also noticed that the cuticle had separated from the epidermis in most of the mutant animals examined (small black arrows in Figure 1), but had not done so in the wild-type (Figure 1, A and J). This phenotype had been previously reported in cor mutant embryos (Lamb et al. 1998), but not for other SJ mutants. Also consistent with the previous report about cor mutant phenotypes, we observed fainter denticle belts (that were nevertheless correctly patterned), and deposits in the areas of the SGs in nearly every mutant cuticle examined (data not shown).

To verify that these observed phenotypes were due to the loss of the SJ genes and not due to second site mutations on the mutant chromosomes, we examined animals hemizygous for a subset of these mutations, as well as animals ubiquitously expressing RNAi against SJ genes, and observed similar defects in DC and head involution (Table 1). These observations demonstrate that it is the zygotic loss of SJ gene function that is responsible for defective head involution and DC, thus indicating an essential role for multiple SJ genes in these morphogenetic developmental events.

During the analysis of cuticle phenotypes in the SJ mutants we observed a number of embryos that did not produce any observable cuticle (Figure 1G). This phenotype varied from having a thin cuticle that has some recognizable features (Figure 1H), to essentially no cuticle, resulting in a disorganized tissue mass (Figure 1I). To characterize developmental defects associated with

| Genotype                  | % Empty Cuticle Phenotype* (n) | % Head Involution Defectivec | % Dorsal Closure Defectivec |
|---------------------------|-------------------------------|-------------------------------|----------------------------|
| Cor956                    | 22.4 ± 8.6 (245)              | 68.7 ± 27.1                  | 5.9 ± 1.4                  |
| Df(3R)BSC146 / Cont*956   | 21.3 ± 13.8 (337)             | 46.2 ± 14.6                  | 6.2 ± 1.9                  |
| cora                       | 53.5 ± 4.5 (314)              | 80.5 ± 17.3                  | 27.6 ± 2.9                 |
| kune309                    | 50.7 ± 4.5 (417)              | 87.2 ± 12.8                  | 5.1 ± 5.7                  |
| Df(2R)BSC696 / kune309    | 31.4 ± 8.0 (358)              | 55.8 ± 20.0                  | 6.3 ± 2.6                  |
| LacG00044                 | 61.5 ± 12.0 (356)             | 61.4 ± 25.4                  | 2.6 ± 2.5                  |
| Df(2R)BSC305 / LacG00044 | 42.9 ± 10.7 (358)             | 78.1 ± 5.1                   | 5.5 ± 2.7                  |
| Mcr1                      | 47.3 ± 19.3 (394)             | 97.5 ± 4.2                   | 19.5 ± 11.2                |
| Mcr2ydro421               | 28.9 ± 6.5 (467)              | 74.7 ± 15.9                  | 1.1 ± 1.2                  |
| Nrg17                     | 64.6 ± 26.8 (490)             | 99.6 ± 0.6                   | 1.6 ± 1.4                  |
| Act5c > Nrg-RNAi          | 73.3 ± 33.0 (408)             | 72.6 ± 15.1                  | 0.9 ± 1.3                  |
| nrv22CC1649               | 26.4 ± 16.9 (201)             | 96.9 ± 6.3                   | 3.0 ± 2.1                  |
| Nrx-IV4304                | 25.4 ± 16.1 (474)             | 95.8 ± 3.8                   | 41.4 ± 16.9                |
| Tsf2KG01571               | 26.2 ± 27.3 (336)             | 100 ± 0                      | 5.2 ± 3.6                  |

* Mean plus standard deviation from at least three independent experiments.
* Total number of embryos examined in all experiments.
* Mean plus standard deviation of all the embryos that produced a cuticle from all the experiments.

Table 1 Cuticle phenotypes of SJ mutant embryos

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these embryos, we collected embryos from a 1 hr egg lay, aged them for 17 hr, and stained them with antibodies against α-Spectrin and with 4',6-diamidino-2-phenylindole (DAPI). At this stage, wild-type animals have secreted a cuticle that prevents their examination by indirect immunofluorescence, whereas many mutant animals (identified by a lack of Yellow Fluorescent Protein expressed from balancer chromosomes) could be examined in this way (Figure 2A). Notably, all 10 of the SJ mutant alleles we tested had a significantly higher percentage of embryos arrested prior to stage 16 (as judged by gut morphology and degree of head involution) than w^{1118} control embryos (Fisher’s exact test; Figure 2E). Examples of arrested mutant embryos are shown in Figure 2, B–D, demonstrating a range of terminal phenotypes from those arrested as germ band extended embryos (Figure 2B), to ones showing severely disorganized body plans (Figure 2D).

**SJ proteins are expressed in early embryos coincident with major morphogenetic events**

The finding that SJ genes are required for morphogenetic events that happen as early as germ band retraction raised questions about the protein expression of these genes during early stages of development. Typically, studies of SJ genes have monitored the expression of these proteins beginning in stage 12–13 embryos where they localize all along the lateral membrane prior to assembling into the region of the SJ beginning in stage 15 (e.g., Tikova et al. 2010). In order to gain a better understanding of their expression prior to these stages, we used immunohistochemistry to examine the expression and subcellular localization of eight core SJ proteins. These experiments revealed that all eight SJ proteins (Cor, Cont, Kune, Mcr, Mtx, Nrg, Nrv, and Nrx-IV) are expressed as early as stage 10 of embryogenesis where they are associated with the membrane (Figure 3 and data not shown).

**Mutations in SJ genes show defective morphogenesis of embryonic SGs**

Given our results showing that SJ genes are required for morphogenetic events during midembryogenesis, we set out to examine if these genes are also required for morphogenetic processes during organogenesis that occur at the same time. The salivary gland is an epithelial tube that begins to form at stage 10 when cells invaginate from the ventral surface (Myat et al. 2005). The SG elongates through a process of coordinated cell shape changes and collective cell migration where the distal tips cells extend protrusions and drive the migration of the gland over the visceral mesoderm (Bradley et al. 2003; Vining et al. 2005). Since the SG reaches its full extension by stage 15 of embryogenesis, we examined SGs (immunostained with antibodies against ATPα and Uif) from stage 16 animals mutant for Cont^{EX56}, kune^{C309}, Mcr^{1}, and Nrg^{14} (Figure 4). In all cases the SGs invaginated and migrated, giving rise to organs that could be unambiguously identified as SGs. Nevertheless, the SGs from each mutant line exhibited abnormal morphology characterized by short and fat glands. In kune^{C309} (Figure 4C) and Mcr^{1} (Figure 4D) mutant animals, the most notable phenotype was an abnormal broad morphology of the lumen of the SG. In addition to abnormal lumen morphology, Cont^{EX56}, Mcr^{1}, and kune^{C309} glands often displayed a bent appearance suggestive of aberrant migration. Of these, Cont^{EX56} glands are the most abnormal, often folding back upon themselves (Figure 4B). We knocked down Mcr and Kune expression specifically in the SG using RNAi driven by fkh-GAL4, and observed identical phenotypes to those produced in the loss of function alleles (Figure 4F and data not shown). These results indicate that the SG defects in the loss of function SJ mutations are not due to second site mutations, and further suggest that the requirement for these genes in morphogenesis is tissue autonomous.

**SGs from SJ mutant embryos exhibit defective cellular rearrangements and cell shape changes**

To examine the cellular defects associated with the aberrant SG phenotypes in SJ mutant animals, we fixed and stained w^{1118}, Cont^{EX56}, Mcr^{1}, Nrg^{14}, kune^{C309}, fkh-GAL4 > Mcr-RNAi, and fkh-GAL4 > kune-RNAi stage 16 embryos with Hoescht and antibodies against Unif and spectrin (Figure 4D) mutant animals, the most notable phenotype was an abnormal broad morphology of the lumen of the SG. In addition to abnormal lumen morphology, Cont^{EX56}, Mcr^{1}, and kune^{C309} glands often displayed a bent appearance suggestive of aberrant migration. Of these, Cont^{EX56} glands are the most abnormal, often folding back upon themselves (Figure 4B). We knocked down Mcr and Kune expression specifically in the SG using RNAi driven by fkh-GAL4, and observed identical phenotypes to those produced in the loss of function alleles (Figure 4F and data not shown). These results indicate that the SG defects in the loss of function SJ mutations are not due to second site mutations, and further suggest that the requirement for these genes in morphogenesis is tissue autonomous.

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or were strongly reduced in their rearrangements relative to wild type (Table 2).

The analysis of z-series reconstructions also allowed us to examine the lumen morphology that we noted to be aberrant in the SJ mutant stage 16 glands (Figure 4). Interestingly, the broad lumen phenotype frequently observed in SJ mutant glands was not due to an overall increase in lumen diameter, but rather was associated with a substantial asymmetry of the lumen that often appeared flat or collapsed (Figure 5L). In addition to the flat lumen phenotype, we observed variations in diameter along the length of the lumen from

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**Figure 4** SJ mutant SGs exhibit abnormal morphology. Single confocal optical section selected from z-stack that revealed the largest number of cells in cross-section from stage 16 w^{1118} (A) Contex^{956} (B), kune^{309} (C), Mcr^{1} (D), Nrg^{14} (E), and fkh-GAL4 > Mcr-RNAi (F) SGs stained with antibodies against ATPa (red, and in individual channels in center panels) to outline cells and against Uif (green, and in individual channels at far right) to mark the apical membrane, and with DAPI (blue) to mark the nuclei. Note that many of the SJ mutant SGs are shorter and fatter than w^{1118}, and have a broad lumen. The Nrg^{14} SG (E) is most similar to wild type. Scale bars = 20 μm.

**Figure 5** Developmental time course of SG organogenesis from wild-type, Contex^{956}, and Mcr^{1} mutant embryos reveals defective cell rearrangements in the mutant glands. Confocal z-series rendered in xz transverse cross-section of stage 11–16 w^{1118} (A–D), Contex^{956} (E–H), and Mcr^{1} (I–L) SGs stained with antibodies against Uif to mark the apical membrane and Hoescht to label nuclei. The basal surface of each SG is indicated by the yellow line. Note that the number of nuclei surrounding the lumen in w^{1118} SGs decreases from stage 11 to stage 16, whereas the number of nuclei surrounding the lumen of SJ glands does not. These views also highlight the broad, unexpanded or asymmetric lumen found in the mutant glands (arrows in G, H, K and L). Scale bars = 100 pixels.
very narrow to wide, resulting in a convoluted appearance that bends and turns through focal planes (data not shown), explaining the cystic-like lumen phenotype observed in the Contex956 mutant by single confocal section of a lateral view (Figure 4B). Interestingly, we did not observe any noticeable variation in lumen diameter in stage 13, and stage 16 intervening apical membrane, for a total of 10 cells per gland in stage 11, measured the length of two lateral membranes per cell, and the in-mutant SG cells have altered membrane lengths. To accomplish this, we explored whether SJ protein function during embryogenesis. Although we were not the first to notice that some SJ mutants through these z-series reconstructions (data not shown), consistent with what we observed by single confocal section (Figure 4E).

Defective apical membrane expansion along the P/D axis of the developing gland has been shown to contribute to defects in lumen width (Pirraglia et al. 2010). In addition, Maruyama and Andrew demonstrated that changes in apical membrane length also contribute to cell shape changes that are critical for coordinated migration (Maruyama and Andrew 2012). Since we observed defects in both SG lumen morphology and cellular rearrangements, we sought to explore whether SJ mutant SG cells have altered membrane lengths. To accomplish this, we measured the length of two lateral membranes per cell, and the intervening apical membrane, for a total of 10 cells per gland in stage 11, stage 13, and stage 16 w1118, Mcr1, and Contex956 animals (Table 3). Mcr1 and Contex956 SG cells have apical membranes that are significantly shorter in length than in w1118 SG cells in stage 11 embryos, yet are equivalent to w1118 in length by stage 13. Interestingly, the apical membrane span decreases in the Mcr1 mutant glands, while the apical membrane of Contex956 and w1118 glands remain relatively constant from stages 13 to 16. On the lateral membrane, we found that both Mcr1 and Contex956 mutant glands had similar lengths to w1118 at stage 11 and 13 SGs, whereas Mcr1 mutant SG cells had significantly longer lateral membrane length compared to w1118 at stage 16. Since this phenotype was not observed in Contex956, we examined stage 16 SGs from two additional SJ mutants, kune309 and Nrg14, and found that only the lateral membrane length in kune309 was significantly different from w1118 (Table 3). We also used Forkhead-GAL4 to specifically express RNAi against Mcr and kune in the SG. At stage 16, Fkh > Mcr-RNAi recapitulated both the shorter apical membrane and longer lateral membrane found in Mcr1 mutant animals at stage 16 (Table 3), and Fkh > kune-RNAi recapitulated the longer lateral membrane found in kune309 mutant animals. Together these results reveal that as wild-type SG cells undergo a modest reduction in apical basal length between stages 11 and 16, the SG cells in some, but not all, SJ mutant animals fail to undergo this change. Furthermore, the similarity in phenotypes between the loss of function alleles and the SG-specific RNAi suggests a tissue autonomous role for these genes in normal morphogenesis.

**DISCUSSION**

The purpose of this study was to reinvestigate a representative collection of SJ genes for a developmental role during essential morphogenetic events during embryogenesis in Drosophila. For this study we chose nine different core SJ genes including those that encode transmembrane proteins such as Nrx-IV, Contactin, the homophilic adhesion protein Nrg, a component of the ATPase complex, a claudin, as well as the cytoplasmic SJ-organizing protein Cor. Somewhat surprisingly, we found that all nine of these genes are required for embryonic developmental events as early as germ band retraction, with highly penetrant defects in head involution and SG organogenesis. Two important implications follow from these new observations. First, a large collection of SJ proteins plays essential roles in morphogenesis throughout embryonic development. Although we were not the first to notice that some SJ genes are required for DC (Feohan et al. 1994; Baumgartner et al. 1996), this study demonstrates how prevalent the requirement for SJ genes is throughout embryonic development. Second, and more importantly, the timing of developmental events requiring SJ gene function indicates that these proteins are likely required for a function that is independent of their role in forming an occluding junction in the lateral membrane.

**SJ genes are required for embryonic morphogenesis**

While examining late-stage mutant embryos by immunofluorescence, we noticed a substantial number of embryos that appeared to arrest development prior to stage 16, with some failing to complete germ band retraction (Figure 2). Because these embryos failed to secrete cuticle they likely represent the high percentage of “Empty Cuticle” embryos we observed in our cuticle analyses of SJ mutants (Figure 1 and Table 1), suggesting a role for SJ genes in early developmental events. Of those mutant embryos that progressed far enough to secrete a cuticle, the most penetrant defect we observed was in head involution (Figure 1 and Table 1). We also found clear evidence for defects in dorsal closure, although scoring this phenotype by obvious holes or scabs in cuticle preparations with their vitelline envelope intact almost certainly underestimated the penetration of these defects. Several other genes that are required for the coordination of DC lead to subtle defects that are not obvious by cuticle preparations, but clearly play important roles. For example, mutations in wingless perturb the organization of the dorsal most epithelial cells and result in a slower and uncoordinated closure that nevertheless goes on to completion (Kalschmidt et al. 2002). Thus, to better examine the role of SJ genes in DC it will be important to examine mutant embryos with a variety of cellular markers and by live imaging.
Since most SJ genes have been studied for their role in establishing the occluding junction, there is incomplete knowledge about the expression and localization of these proteins in earlier embryos. RNA sequencing data from the modENCODE project indicate that RNA for many SJ genes is present in 0–2 hr embryos (Graveley et al. 2011), suggesting that they are expressed maternally. We found that at least the eight SJ proteins we examined are expressed as early as germ band extended embryos, where they are associated with the lateral membrane (Figure 3). Although we did not examine them fully, we noticed expression of some SJ proteins in even earlier embryos raising the question of whether they could be required for earlier developmental events (S.H. and R.W., unpublished observation). Since our analysis was limited to zygotic loss of function mutations, we would have missed any earlier developmental requirements that were rescued by maternal contributions. A requirement for some SJ genes in oogenesis (Batz et al. 2014; R. W., unpublished observation), however, precludes our ability to examine embryos with maternal/zygotic loss of function mutations in these genes.

SJ genes are also required for the organogenesis of epithelial organs that is occurring coincident with the major developmental events of midembryogenesis. We noticed that the SGs of SJ mutant embryos appeared short and fat compared to wild-type glands at stage 16 (Figure 3). Although we did not examine them fully, we noticed expression of some SJ proteins in even earlier embryos raising the question of whether they could be required for earlier developmental events (S.H. and R.W., unpublished observation). Since our analysis was limited to zygotic loss of function mutations, we would have missed any earlier developmental requirements that were rescued by maternal contributions. A requirement for some SJ genes in oogenesis (Batz et al. 2014; R. W., unpublished observation), however, precludes our ability to examine embryos with maternal/zygotic loss of function mutations in these genes.

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This phenotype results primarily from defects in cell rearrangements that normally occur between stages 11 and 16 (Figure 5 and Table 2), although we also found evidence for defects in cell shape changes for two of the four SJ mutants we examined (Table 3).

How might the loss of SJ gene function affect morphogenesis of the SGs? Chung and Andrew found that loss of function mutations in Cadherin 99C (Cad99C) result in glands that are longer and thinner with fewer cells around the lumen of the gland, whereas overexpression of Cad99C had the opposite effect, and was similar to what we observed with the SJ mutations (Chung and Andrew 2014). They went on to show that Cad99C localizes to the apical domain of SG cells and promotes apical character in the cells. Laprise and colleagues showed that a subset of SJ genes acts in a redundant pathway with yurt to maintain apical basal polarity during midembryogenesis (Laprise et al. 2009). Although the authors indicated that there is no strong loss of apical/basal polarity in single SJ mutants, it is possible that loss of SJ gene function may result in a subtle defect in polarity, perhaps by playing a role in vesicle trafficking. In this regard, it is noteworthy that mutations in all of these SJ genes lead to a phenotype in which the epidermis secretes a thin and undeveloped cuticle (Figure 1; Lamb et al. 1998), and tracheal cells of most SJ mutants fail to secrete Serpentine or Vermiform, two chitin deacetylases required for tracheal tube length (Luschnig et al. 2006; Wang et al. 2006). In further support of the notion that single SJ mutations may have a subtle effect on apical/basal polarity, the convoluted tracheal phenotype found in cor mutant animals can be partially suppressed by loss of one copy of crumbs (Laprise et al. 2010).

The role of SJ proteins in morphogenesis is likely independent of its role in forming an occluding junction

It is instructive to compare the timing of embryonic developmental events with the biogenesis of the SJ. Germ band retraction is complete by stage 12 of embryogenesis, whereas head involution and DC occur during stages 13 and 14 (Figure 6). Similarly, SG and hindgut morphogenesis occur between stages 11 and 15. The biogenesis of the SJ is an extended process, but the refinement of SJ proteins to the region of the SJ occurs primarily during stages 15 and 16 (Figure 6). This correlates well with ultrastructural analysis showing the formation of extracellular septa in the epidermis starting in stage 14 embryos, but not showing a mature junction until stage 17 (Tepass and Hartenstein 1994). Functionally, the SG and tracheal epithelia are not physiologically tight until late stage 15 (Paul et al. 2003). Together these observations suggest that a physiologically “tight” junction is not required for the major body plan morphogenetic events, nor for organogenesis in Drosophila. Interestingly, many SJ proteins are expressed very early in embryonic development and are only re-targeted to the region of the SJ relatively late in embryogenesis (Figure 3 and Figure 6; Tiklova et al. 2010). We therefore propose that SJ proteins play an essential role in morphogenesis during early and midembryogenesis, and then are redeployed for a role as an occluding junction later in development. There is an interesting evolutionary implication of this model. If these functions of SJ proteins are truly independent, then the selective pressure for each function may have distinct evolutionary trajectories. A recent paper by Ganot examined the evolution of occluding junction proteins (Ganot et al. 2015), and concluded that the ancestral occluding junction was a septate-like junction composed of proteins closely related to the SJ proteins of Drosophila. At least some of these SJ genes are present in unicellular choanoflagellates (one of the last unicellular ancestors to metazoans). Core tight junction proteins, on the other hand, are not found in basal metazoan lineages, and appear to originate in chordates. As ancestral organisms with SJs evolved into chordates in which the SJ was supplanted by a tight junction, what became of the core SJ proteins? It will be interesting to identify the true orthologs of SJ genes and characterize their subcellular localization and function. We would predict that the proteins encoded by these genes will have lost their role in the occluding junction, but may still be present and function in essential morphogenetic processes in development.

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LITERATURE CITED

Batz, T., D. Forster, and S. Luschnig. 2014 The transmembrane protein Macroglobulin complement-related is essential for septate junction formation and epithelial barrier function in Drosophila. Development 141: 899–908.

Baumgartner, S., J. T. Littleton, K. Broadie, M. A. Bhat, R. Harbecke et al. 1996 A Drosophila neurexin is required for septate junction and blood-nerve barrier formation and function. Cell 87: 1059–1068.

Behr, M., D. Riedel, and R. Schuh. 2003 The claudin-like megatraceha is essential in septate junctions for the epithelial barrier function in Drosophila. Dev. Cell 5: 611–620.

Bradley, P. L., M. M. Myat, C. A. Comeaux, and D. J. Andrew. 2003 Posterior migration of the salivary gland requires an intact visceral mesoderm and integrin function. Dev. Biol. 257: 249–262.

Buszczak, M., S. Paterno, D. Lighthouse, J. Bachman, J. Planck et al. 2007 The Carnegie protein trap library: a versatile tool for Drosophila developmental studies. Genetics 175: 1505–1531.

Chung, S., and D. J. Andrew. 2014 Cadherin 99C regulates apical expansion and cell rearrangement during epithelial tube elongation. Development 141: 1950–1960.

Dietzl, G., D. Chen, F. Schnorrer, K. C. Su, Y. Barinova et al. 2007 A genome-wide transgenic RNAi library for conditional gene inactivation in Drosophila. Nature 448: 151–156.

Faireva-Sarraill, C., S. Banerjee, J. Li, M. Hortsch, M. Laval et al. 2004 Drosophila contactin, a homolog of vertebrate contactin, is required for septate junction organization and paracellular barrier function. Development 131: 4931–4942.

Fehon, R. G., K. Johansen, I. Rebay, and S. Artavanis-Tsakonas. 1991 Complex cellular and subcellular regulation of notch expression during embryonic and imaginal development of Drosophila: implications for notch function. J. Cell Biol. 113: 657–669.

Fehon, R. G., I. A. Dawson, and S. Artavanis-Tsakonas. 1994 A Drosophila homologue of membrane-skeleton protein 4.1 is associated with septate junctions and is encoded by the coracle gene. Development 120: 545–557.

Ganot, P., D. Zoccola, E. Tambutte, C. R. Voolstra, M. Aranda et al. 2015 Structural molecular components of septate junctions in cnidarians point to the origin of epithelial junctions in eukaryotes. Mol. Biol. Evol. 32: 44–62.

Genova, J. L., and R. G. Fehon. 2003 Neuroglian, Gliotactin, and the Na+ K+ ATPase are essential for septate junction function in Drosophila. J. Cell Biol. 161: 979–989.
Perrimon, N., 1988 The maternal effect of lethal[1] discs-large-1: a recessive oncogene of Drosophila melanogaster. Dev. Biol. 127: 392–407.

Pirraglia, C., J. Walters, and M. M. Myat, 2010 Pak1 control of E-cadherin endocytosis regulates salivary gland lumen size and shape. Development 137: 4177–4189.

Schneider, C. A., W. S. Rasband, and K. W. Eliceiri, 2012 NIH Image to ImageJ: 25 years of image analysis. Nat. Methods 9: 671–675.

TePas, U., and V. Hartenstein, 1994 The development of cellular junctions in the Drosophila embryo. Dev. Biol. 161: 563–596.

Tiklova, K., K. A. Senti, S. Wang, A. Graslund, and C. Samakovlis, 2010 Epithelial septate junction assembly relies on melanotransferrin iron binding and endocytosis in Drosophila. Nat. Cell Biol. 12: 1071–1077.

Uurakabe, S., J. S. Handler, and J. Orloff, 1970 Effect of hypertonicity on permeability properties of the toad bladder. Am. J. Physiol. 218: 1179–1187.

Vining, M. S., P. L. Bradley, C. A. Comeaux, and D. J. Andrew, 2005 Organ positioning in Drosophila epithelia requires complex tissue–tissue interactions. Dev. Biol. 287: 19–34.

Wang, L., J. Evans, H. K. Andrews, R. B. Beckstead, C. S. Thummel et al., 2008 A genetic screen identifies new regulators of steroid-triggered programmed cell death in Drosophila. Genetics 180: 269–281.

Wang, S., S. A. Jayaram, J. Hemphala, K. A. Senti, V. Tsarouhas et al., 2006 Septate-junction-dependent luminal deposition of chitin deacetylases restricts tube elongation in the Drosophila trachea. Curr. Biol. 16: 180–185.

Ward, R. E., R. S. Lamb, and R. G. Fehon, 1998 A conserved functional domain of Drosophila coracle is required for localization at the septate junction and has membrane-organizing activity. J. Cell Biol. 140: 1463–1473.

Ward, R. E., L. Schweizer, R. S. Lamb, and R. G. Fehon, 2001 The protein 4.1, ezrin, radixin, moesin (FERM) domain of Drosophila Coracle, a cytoplasmic component of the septate junction, provides functions essential for embryonic development and imaginal cell proliferation. Genetics 159: 219–228.

Ward, R. E., J. Evans, and C. S. Thummel, 2003 Genetic modifier screens in Drosophila demonstrate a role for Rbo1 signaling in edcsyne-triggered imaginal disc morphogenesis. Genetics 165: 1397–1415.

Wells, E. R., J. D. Barry, S. J. Warrington, S. Cuhlmann, P. Evans et al., 2013 Control of tissue morphology by Fasciclin III-mediated intercellular adhesion. Development 140: 3858–3868.

Wu, V. M., J. Schulte, A. Hirschi, U. Tepass, and G. J. Beitel, 2004 Sinuous GTPase controls epithelial tube size control. J. Cell Biol. 164: 313–323.

Wu, V. M., M. H. Yu, R. Paik, S. Banerjee, Z. Liang et al., 2007 Drosophila Varicose, a member of a new subgroup of basolateral MAGUKs, is required for septate junctions and tracheal morphogenesis. Development 134: 999–1009.

Xu, N., G. Bagumian, M. Galiano, and M. M. Myat, 2011 Rho GTPase controls Drosophila salivary gland lumen size through regulation of the actin cytoskeleton and Moesin. Development 138: 5415–5427.

Zhang, L., and R. E. Ward, 2009 uninsflatable encodes a novel ectodermal apical surface protein required for tracheal inflation in Drosophila. Dev. Biol. 336: 201–212.