Arf-GEF localization and function at myosin-rich adherens junctions via coiled-coil heterodimerization with an adaptor protein

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ABSTRACT Tissue dynamics require regulated interactions between adherens junctions and cytoskeletal networks. For example, myosin-rich adherens junctions recruit the cytoskeleton Arf-GEF Steppke, which down-regulates junctional tension and facilitates tissue stretching. We dissected this recruitment mechanism with structure–function and other analyses of Steppke and Stepping stone, an implicated adaptor protein. During Drosophila dorsal closure, Steppke’s coiled-coil domain was necessary and sufficient for junctional recruitment. Purified coiled-coil domains of Steppke and Stepping stone heterodimerized through a hydrophobic surface of the Steppke domain. This mapped surface was required for Steppke’s junctional localization and tissue regulation. Stepping stone colocalized with Steppke at junctions, and was required for junctional Steppke localization and proper tissue stretching. A second conserved region of Stepping stone was necessary and largely sufficient for junctional localization. Remarkably, this region could substitute for the Steppke coiled-coil domain for junctional localization and regulation, suggesting the main role of the Steppke coiled-coil domain is linkage to the junctional targeting region of Stepping stone. Thus, coiled-coil heterodimerization with Stepping stone normally recruits Step to junctions. Intriguingly, Stepping stone’s junctional localization also seems partly dependent on Steppke.

INTRODUCTION
Epithelia provide regulated barriers for organ structure and function. These multicellular sheets also undergo amazing dynamics. Epithelial cells can change shape and interactions while assembling, disassembling, and remodeling the adherens junctions (AJs) that connect them. The classical cadherins of AJs (such as Drosophila E-cadherin [DE-cad]) mediate homophilic cell–cell adhesion and link cortical actin networks through the adaptor proteins β-catenin and α-catenin (Harris and Tepass, 2010; Takeichi, 2014). These cadherin–catenin complexes also transduce signals for AJ growth and regulation. Such signaling can be biochemical, such as the local induction of Arp2/3-based actin networks for junction growth (Ratheesh et al., 2013), or the regulated recruitment of endocytic machinery for junction removal (Cadwell et al., 2016). Additionally, mechanical signal transduction occurs. For example, actomyosin-based tension can unfurl α-catenin to expose binding sites for proteins such as Vinculin, which then reinforce AJ–actin interactions (Leckband and de Rooij, 2014; Pinheiro and Bellaïche, 2018; Yap et al., 2018). AJ regulatory signaling is an area of intense research, and many mechanisms remain incompletely understood.

Actomyosin activity at AJs has been shown to recruit the cytohesin Arf-GEF Steppke (Step) during Drosophila dorsal closure (West et al., 2017) and in Drosophila wing discs (Rauskolb et al., 2019). This recruitment is part of a negative feedback loop that antagonizes junctional actomyosin contractility through Step Arf-GEF activity. During dorsal closure, the negative feedback allows orderly stretching of the epidermis. Without Step, heightened junctional actomyosin activity holds local cell groups in distorted configurations (multicellular rosettes). Cytohesin Arf-GEFs also down-regulate actomyosin for zebrafish epiboly (West et al., 2017), for the partial cleavage of the syncytial Drosophila embryo (Lee and Harris, 2013), for proper organization of the Drosophila wing disc epithelium
(Rauskolb et al., 2019), and for mammalian cell podosome formation (Ratliff et al., 2017).

Cytohesins contain a central Sec7 domain that conveys Arf-GEF activity, as well as two domains implicated in plasma membrane recruitment: an N-terminal coiled-coil (CC) domain and a C-terminal pleckstrin homology (PH) domain (Gillingham and Munro, 2007; Donaldson and Jackson, 2011). Cytohesin PH domains bind either PIP2 or PIP3 (Klarlund et al., 2000)—the Step PH domain preferentially binds PIP3 (Britton et al., 2002; Lee et al., 2015)—and separate sites of the PH domain can also bind Arf6 and Arf4 (Cohen et al., 2007; Hofmann et al., 2007). The CC domains of cytohesins can bind to a variety of adaptor proteins, including CNK1, GRASP/Tamalin, FRMD4A, and GRSP-1 (Nevrivy et al., 2000; Ikenouchi and Umeda, 2010; Lim et al., 2010; Hahn et al., 2013). Relevant to AJs, FRMD4A and GRSP-1 both bind cytohesin-1 via CC domain heterodimerization and to the AJ-associated protein Par-3 through a separate site, and these interactions recruit cytohesin-1 to nascent AJs for their maturation in mammalian cell culture (Ikenouchi and Umeda, 2010). Of note, cytohesin CC domains can also undergo homodimerization (DiNitto et al., 2007; Liu et al., 2015), and have additionally been implicated in cytohesin autoinhibition through interaction with the PH domain (Li et al., 2012; Hiester and Santy, 2013).

From cleavage-stage Drosophila embryos, Stepping stone (Sstn) was identified as the main protein to communoprecipitate with Step (Liu et al., 2015). The closest mammalian homologue of Sstn was found to be FRMD4A. In vitro, Sstn and Step were shown to bind directly, and this interaction required each protein’s CC domain. In vivo, embryos depleted of Sstn or Step displayed similar cleavage-stage actomyosin misregulation defects, and overexpressed versions of the proteins displayed striking colocalization at myosin-rich domains of pseudocleavage furrows. Step localization to these domains required its CC domain, whereas the Sstn localization required a separate region of the protein conserved in insects (termed conserved region [CR]; Liu et al., 2015). Liu et al. (2015) studied the syncytial embryo before cells and AJs had formed. More recently, overexpressed versions of Step and Sstn were also found to colocalize at AJs of Drosophila wing discs, and the AJ localization of overexpressed Step was diminished by depletion of Sstn (Rauskolb et al., 2019). Moreover, depletions of Step or Sstn resulted in excessive junctional actomyosin in the wing disc (Rauskolb et al., 2019). The studies of Liu et al. (2015) and Rauskolb et al. (2019) suggest a model of how Step–Sstn interaction could mediate Step recruitment to myosin-rich AJs. We sought to test this model by addressing the following questions. Step and Sstn are recruited to both myosin-rich domains of pseudocleavage furrows and to myosin-rich AJs, but do their protein domains play similar roles in both contexts? The CC domain of Step is necessary and sufficient for LE localization, and intriguingly, the junctional localization of endogenous Sstn also relied substantially on Step. Thus, Step and Sstn may support each other’s localization as an interacting pair. Additionally, the Sstn CR was found to be necessary and largely sufficient for localization to myosin-rich junctions, and could functionally substitute for the Step CC domain, implicating Sstn function as an adaptor between Step and junctions.

**RESULTS**

The CC domain of Step is necessary and sufficient for LE and tricellular junction localization

To investigate how deletion of the CC domain or the PH domain of Step affects the recruitment of Step to AJs at dorsal closure, we compared three previously generated constructs: GFP-tagged full-length Step, StepΔCC, and StepΔPH (Figure 1A shows all Step and Sstn constructs used for the in vivo structure–function analyses of this article; Lee and Harris, 2013; Lee et al., 2015; Liu et al., 2015). Expression of the UAS constructs was driven with daughterless-Gal4 from the same genomic site (the constructs were each inserted at the attp40 recombination site on chromosome 2 to avoid positional effects on transcription) and imaged live during dorsal closure with the same microscope settings (to allow comparisons of signal intensities between constructs). Full-length Step was enriched most strongly at epidermal LE and tricellular junctions (Figure 1B; white arrow indicates the LE and yellow arrows indicate tricellular junctions of the lateral epidermis), and less so at bicellular junctions of the lateral epidermis (Figure 1, B and E; West et al., 2017). GFP-StepΔPH showed a similar localization pattern as full-length GFP-Step, with statistically indistinguishable distributions among the junction types (Figure 1, B and E). In contrast, removal of the CC domain severely perturbed the localization pattern. GFP-StepΔCC displayed a relatively low level of general plasma membrane localization (observed consistently in comparisons of eight embryos expressing full-length GFP-Step and seven embryos expressing GFP-StepΔCC), without substantial enrichment at epidermal LE or tricellular junctions (Figure 1, B and E).
Deletion of the CC domain could affect the location of Step because a specific interaction site was eliminated or because overall Step structure was destabilized. Thus, we tested whether the Step CC domain alone was sufficient for conveying the full-length Step localization pattern. UAS-StepCCalone-GFP was generated and expressed using daughterless-Gal4. Remarkably, StepCCalone-GFP was enriched at epidermal LE and tricellular junctions in a pattern resembling that of full-length GFP-Step (Figure 1, C and E). To test the localization capability of the Step PH domain, we imaged a GFP-tagged construct of the Step PH domain together with its C-terminal polybasic region constitutively driven by a tubulin promoter, a construct called "tGPH," which was developed as a PIP3 marker in...
Coiled coils form from the interactions of CC domains. CC domains consist of an α-helix with a hydrophobic face flanked by polar amino acid residues. Interactions between the hydrophobic residues form the core interface of a coiled coil, and interactions of hydrophobic and polar residues additionally determine the orientation and specificity of the interaction (Mason and Arndt, 2004; Grigoryan and Keating, 2008). Using the Phyre2 web portal, the structure of the CC domain of Step was predicted based on sequence similarities to CC domains with known structures (Kelley et al., 2015). The predicted structure was then analyzed with CN3D software to localize hydrophobic residues (Wang et al., 2000). A hydrophobic face was detected along the full length of the predicted structure of the Step CC domain (Figure 3A).

MBP- and GST-tagged forms of the CC domains of Step and Sstn were generated and used in blot overlay assays to test for direct interaction. GST-StepCC showed strong interaction with MBP-SstnCC and vice versa (Figure 3C, asterisks), but no homodimerization of either protein was detected (Figure 3C). To test whether the hydrophobic residues of the Step CC domain were necessary for this interaction, three mutated constructs were tested: the hydrophobic residues were mutated to polar serine residues along the full length of the CC domain, or only along its N-terminal half or its C-terminal half (Figure 3B shows the wild-type and altered Step CC domains as well as the wild-type Sstn CC domain used in the in vitro binding assays). Strikingly, none of the mutated Step CC domains showed any interaction with the Sstn CC domain (Figure 3D), in contrast with the strong binding of the control wild-type Step CC domain (Figure 3D, asterisk). Thus, the CC domains of Step and Sstn interact directly and this interaction requires the full hydrophobic region of the Step CC domain.

**The Step-Sstn interaction site is necessary for Step localization, and Step homodimerization seems irrelevant.**
To test the role of the Step-Sstn binding site in vivo, we inserted the sequence for StepCC[N*] into GFP-Step to replace the wild-type CC domain with the mutated CC domain (creating GFP-StepΔCC+CC[N*]). Because all three mutated Step CC domains abrogated binding to the Sstn CC domain in vitro, we selected StepCC[N*] for the in vivo analysis because we expected it to have less chance of disrupting full-length Step structure—it contains fewer mutations than StepCC[N*+C*], and compared with StepCC[C*] the StepCC[N*] mutations are farther from the Sec7 and PH domains. GFP-StepΔCC+CC[N*] was expressed in vivo using daughterless-Gal4. In contrast to full-length GFP-Step, GFP-StepΔCC+CC[N*] failed to enrich at epidermal LE or tricellular junctions, and instead displayed a more general plasma membrane localization pattern which included enrichment at protrusions of the LE and of epidermal cells to the rear (Figure 4, cyan arrows). We noted that the general
plasma membrane localization of GFP-StepΔCC+CC[N*] was more easily detected than that of GFP-StepΔCC, suggesting better general stability of GFP-StepΔCC+CC[N*].

To further address the possible influence of Step CC domain homodimerization, we sought to replace the CC domain of Step with a CC domain capable of homodimerization but that would not interact with Sstn. Two CC domains were selected from the literature: APH and E/K42. Both are synthetic sequences optimized for CC homodimerization in vitro. The APH sequence does so in an antiparallel orientation (Gurnon et al., 2003), and the E/K42 sequence does so in a parallel orientation (Graddis et al., 1993). Two constructs were generated to replace the wild-type Step CC domain
GFP-StepΔCC+CC[N*] displayed a diffuse plasma membrane distribution with some enrichment to cell protrusions at the LE and within the epidermis (cyan arrows). A relatively high cytosolic pool was also apparent. GFP-StepΔCC+APH was very weakly detected and appeared to localize diffusely over the plasma membrane. GFP-StepΔCC-EK42 was detected evenly over the plasma membrane with some enrichment to cell protrusions (cyan arrow). Also, a number of very bright puncta appeared to be cytosolic. Embryo numbers combined from two experiments are shown at the bottom of each image.

FIGURE 4: Mutating or replacing the CC domain of Step disrupts Step localization. All images were taken live with the same settings at dorsal closure (the amnioserosa is at the top of each image). Construct expression was driven with daughterless-Gal4. Full-length GFP-Step localized to the LE and tricellular junctions of the epidermis. GFP-StepΔCC+CC[N*] displayed a diffuse plasma membrane distribution with some enrichment to cell protrusions at the LE and within the epidermis (cyan arrows). A relatively high cytosolic pool was also apparent. GFP-StepΔCC+APH was very weakly detected and appeared to localize diffusely over the plasma membrane. GFP-StepΔCC-EK42 was detected evenly over the plasma membrane with some enrichment to cell protrusions (cyan arrow). Also, a number of very bright puncta appeared to be cytosolic. Embryo numbers combined from two experiments are shown at the bottom of each image.

with either synthetic CC domain, and the constructs were expressed in vivo using daughterless-Gal4. Neither protein displayed the epidermal LE and tricellular junction enrichment of full-length GFP-Step. GFP-StepΔCC+APH was weakly expressed (Figure 4). GFP-StepΔCC+EK42 was more strongly expressed but localized generally to the plasma membrane with enrichment at LE protrusions (Figure 4, cyan arrow). Taken together, these results suggest the specific importance of CC heterodimerization with Sstn for the junctional localization of overexpressed Step constructs.

The Sstn CR is necessary and sufficient for Sstn localization

Next, we performed a structure–function analysis to investigate how Sstn is localized. As Sstn only has two distinctive regions, the CC domain and the region conserved across insects (Liu et al., 2015), we compared full-length Sstn with SstnΔCC and SstnΔCR, each tagged with GFP and inserted at the same genomic site (Liu et al., 2015), after expression with daughterless-Gal4. Full-length GFP-Sstn localized in a similar pattern as Step, with highest enrichment at epidermal LE and tricellular junctions (Figure 5A). GFP-SstnΔCC had similar junctional levels compared with GFP-Sstn (Figure 5A; observed for 6/6 GFP-SstnΔCC embryos compared with 8/8 GFP-Sstn embryos), and its distribution resembled that of GFP-Sstn, except for less accumulation at LE junctions relative to other epidermal junctions (Figure 5, A and B). More strikingly, GFP-SstnΔCR showed a drastic loss of signal compared with full-length GFP-Sstn at all junctions (Figure 5A; observed for 8/8 GFP-SstnΔCR embryos compared with 8/8 GFP-Sstn embryos). Despite this loss, the residual cortical SstnΔCR did show some, although reduced, enrichment at epidermal LE and tricellular junctions (Figure 5B).

Because loss of the CR might destabilize Sstn, we tested whether the CR is sufficient for mediating junctional localization. The Sstn CR was tagged with GFP, driven by daughterless-Gal4, and imaged live at dorsal closure. Strikingly, the Sstn CR was sufficient for enrichment to epidermal LE and tricellular junctions (Figure 5, A and B), although its enrichment to LE junctions was less than that of full-length Sstn (Figure 5, A and B). Thus, the CR of Sstn is necessary and sufficient for mediating much of the localization observed for full-length Sstn. In addition, the CC domain of Sstn contributes to Sstn recruitment to LE junctions.

Replacement of the Step CC domain with the Sstn CR conveys Step localization

Because Step and Sstn were shown to interact via their respective CC domains and the Sstn CR was shown to be sufficient for substantial junctional localization, we hypothesized that the main role of the Step CC domain is to link Step to the Sstn CR through heterodimerization. A prediction from this hypothesis is that the interaction of Step and Sstn could be bypassed by replacing the Step CC domain with the Sstn CR. Thus, we generated a Step construct that substituted the Step CC domain with the Sstn CR, GFP-StepΔCC+CR, and expressed it with daughterless-Gal4 for live imaging at dorsal closure. GFP-StepΔCC was imaged in parallel as a negative control, and as discussed, it displayed weak, general plasma membrane localization (Figure 5C). When the Sstn CR replaced the Step CC domain, however, enrichment to epidermal LE and tricellular junctions occurred (Figure 5C; seen for comparisons of 8/8 embryos each). Thus, the Sstn CR can convey substantial localization activity to a Step construct lacking its CC domain.

Step and Sstn colocalize at tricellular junctions

Overexpressed Step and Sstn constructs can colocalize at myosin-rich cortical domains of the syncitial Drosophila embryo (Liu et al., 2015), and overexpressed Step and Sstn constructs have each been shown to localize to AJs of Drosophila wing disc (Rauskolb et al., 2019). To assess whether the overexpressed proteins colocalize at dorsal closure, daughterless-Gal4 was used to coexpress full-length GFP-Step with an mCherry-tagged construct of Sstn (Liu et al., 2015) and both were coimaged live. GFP-Step and mCherry-Sstn displayed close colocalization with both enriched most strongly at epidermal LE and tricellular junctions (Figure 6A). Side views showed the proteins enriched at the apicolateral domain (Figure 6A), where AJs are found.

To assess the localization of Step and Sstn expressed at endogenous levels, we compared proteins expressed from a GFP-tagged step allele and from a GFP-tagged sstn allele. These alleles were produced by a CRISPR/Cas9 protocol that added the sequence for GFP at the 3′ end of the sequence for each endogenous gene’s coding region (see Materials and Methods). Indicating normal expression and function of the tagged proteins, each line was homozygous viable and fertile. Live imaging at dorsal closure with the same microscope settings revealed strong...
enrichment of endogenous Step-GFP and endogenous Sstn-GFP at epidermal LE and tricellular junctions (Figure 6B). Fixation and staining for DE-cad showed that both endogenously expressed proteins localized at the same apicolateral position as AJs with enrichment to epidermal LE and tricellular junctions (Figure 6C). Thus, endogenous Step and Sstn appear to colocalize at myosin-rich AJs, and junctional levels of Sstn seem to be somewhat lower than those of Step.

Junctional Step requires Sstn, and junctional Sstn depends substantially on Step

With cooverexpression Step and Sstn can increase each other's accumulation at myosin-rich cortical domains of the syncytial Drosophila embryo (Liu et al., 2015). Moreover, depletion of Sstn reduces the localization of overexpressed Step to AJs of the Drosophila wing disc (Rauskolb et al., 2019). To test how Sstn affects the localization of endogenous levels of Step, we probed GFP-tagged endogenous Step and antibody staining of endogenous Step in sstn mutants versus controls at dorsal closure. In control embryos, endogenous Step-GFP localized to epidermal LE and tricellular junctions, as discussed above (Figures 6B and 7A). Similarly, antibody staining for Step revealed clear enrichment at LE junctions, although other junctions were more difficult to discern (Figure 7B). For both Step probes, junctional localization was almost undetectable in sstn mutants (Figure 7, A and B). Live imaging could not detect Step-GFP expressed in the sstn mutants (unpublished data), but fixation revealed marginal Step-GFP enrichment along the LE (Figure 7A, arrows). Staining for DE-cad confirmed the localization of AJs and also showed an abnormally high frequency of multicellular rosettes in the sstn mutants (Figure 7, A and B; highlighted in yellow and quantified to the right). Thus, Sstn is required for recruiting Step to myosin-rich AJs, and sstn mutants display tissue distortions similar to those of step mutants (West et al., 2017).

Next, we analyzed the localization of GFP-tagged endogenous Sstn expressed in step mutant or control embryos. Live imaging was able to detect the junctional localization of Sstn-GFP in both control and step mutant embryos, but the signal was substantially reduced in the step mutants (Figure 7C, arrows). Fixation and staining for DE-cad confirmed that junctional Sstn-GFP was substantially reduced in step mutants (Figure 7D). Together, these results show that the relationship between Sstn and Step is not simply linear, and that it is not equally reciprocal either. Sstn seems to have a greater effect on Step localization than Step does on Sstn localization.
FIGURE 6: Step and Sstn colocalization at dorsal closure. (A) Side and surface views of GFP-Step and mCherry-Sstn expressed with daughterless-Gal4 in the same embryo and coimaged live. Both constructs localized to apicalateral epidermal junctions (side views) and showed strongest enrichment and colocalization to epidermal LE and tricellular junctions (surface views). (B) Surface views of Step-GFP and Sstn-GFP expressed from endogenous GFP insertion alleles and imaged live at dorsal closure with the same settings. The junctional distributions of the proteins were similar, but the junctional levels of Step-GFP were higher than those of Sstn-GFP. LE junction levels are quantified at right. Each replicate comparison is shown in a different color, and for each replicate the data were normalized to the average embryo values for the Step-GFP signal. Total embryo numbers are indicated in brackets. (C) Both Step-GFP and Sstn-GFP were enriched at epidermal LE junctions and tricellular junctions at the apicalateral position of DE-cad. Fixed and stained embryos shown.

The Step CC domain and the Step PH domain are each needed for proper tissue elongation

A step mutant rescue assay was previously used to show the requirement of Step Arf-GEF activity for proper epidermal organization during dorsal closure (West et al., 2017). We used the same assay to test the roles of the Step CC and PH domains. With paired-Gal4, constructs were expressed in stripes in step mutant embryos, and multicellular rosette numbers were compared between construct-expressing and nonexpressing tissue. Full-length Step-GFP substantially rescued the tissue distortions (Figure 8, top left). In contrast, expression of GFP-StepΔCC+CC[N*] failed to reduce the frequency of abnormal multicellular rosettes (Figure 8, bottom left). To test whether the CC domain’s role in tissue organization was related to Sstn, we tested the ability of GFP-StepΔCC+CR to rescue the abnormal rosette numbers in step mutant tissue, and discovered that it had substantial rescue activity (Figure 8, bottom right). Thus, the CC domain of Step is needed for its function, and the CR of Sstn can substitute for this role.

GFP-StepΔPH also failed to reduce multicellular rosette numbers (Figure 8, top right). This lack of function was surprising because GFP-StepΔPH was effectively recruited to myosin-rich junctions of wild-type or step mutant embryos (Figures 1B and 2, A and B). Together, these data indicate that the PH domain is dispensable for the junctional localization of Step but is needed for Step to regulate epidermal organization. This relationship was observed previously for GEF-dead Step (West et al., 2017), and thus it seems that both GEF activity and the PH domain have roles downstream from junctional recruitment.

DISCUSSION

Our data show that the CC domain of Step is a key determinant of its localization to myosin-rich AJs. This localization mechanism seems to depend on direct heterodimerization with the CC domain of Sstn, an interaction mediated by a hydrophobic face of the Step CC domain. For Sstn, a small region conserved across insects is critical for its localization to myosin-rich AJs, suggesting Sstn acts as an adaptor protein between Step and an unknown component of the junctions. Consistent with this model, Step and Sstn display strong colocalization at myosin-rich AJs. Moreover, Sstn is required for Step localization to these sites. However, the localization of Sstn also depends substantially on Step. Taken together, these data show how Sstn can act as an adaptor for recruiting Step to myosin-rich AJs, although the proteins also display partial codependence for localization to these sites. Downstream, the Sstn-Step interaction appears to aid the resolution of multicellular rosettes during dorsal closure.

Multiple results indicate that the junctional localization of Step relies on CC heterodimerization with Sstn. Blot overlay assays showed that the purified CC domains of each protein can interact directly, and allowed mapping of an interaction interface. Deletion of the Step CC domain, or mutation of its interaction interface with Sstn, eliminated junctional localization of overexpressed Step constructs in vivo. Highlighting that it is specifically Step-Sstn CC domain heterodimerization that is critical for the localization of Step to myosin-rich junctions, the Sstn CR was able...
to substitute for the Step CC domain to mediate Step localization and function. Finally, endogenous Step was lost from AJs of sstn mutants. Arguing against a role for Step CC homodimerization, replacement of the native CC domain with artificial CC domains optimized for parallel or antiparallel homodimerization failed to restore junctional localization. Arguing against a nonspecific effect of CC domain manipulations on Step protein stability, junctional localization was disrupted for four constructs each with a distinct CC domain manipulation, and a GFP-tagged construct of the CC domain alone was sufficient for localization to myosin-rich junctions. Thus, CC heterodimerization with Sstn seems critical for the junctional localization of Step.

We also found that a conserved region of Sstn mediates its localization to myosin-rich junctions. Similar to the CC domain of Step, the Sstn CR was necessary and sufficient for localization to such junctions, whereas the Sstn CC domain was relatively dispensable (except for full recruitment to LE junctions). What then is the localization mechanism of the CR? Presumably it binds a partner that is recruited by the junctional actomyosin network and/or opened for binding by actomyosin-based tension. It is important to note that the CR of Sstn is mainly conserved among insects (Liu et al., 2015).

Interestingly though, the closest mammalian homologue of Sstn, the protein FRMD4A, localizes to forming AJs and contains a CC domain that binds to the CC domain of cytohesin-1 (Ikenouchi and Umeda, 2010). In addition, FRMD4A contains an N-terminal FERM domain not found in Sstn and a direct binding site for Par-3 not conserved in Sstn (Ikenouchi and Umeda, 2010; Liu et al., 2015). It is possible that one of these sites, or a separate site, in FRMD4A plays a role similar to the CR of Sstn. For example, Par-3 functions in the junctional recruitment of FRMD4A (Ikenouchi and Umeda, 2010). However, Drosophila Par-3 typically localizes to junctions depleted of actomyosin, as shown in the embryo germ band (Zallen and Wieschaus, 2004), for example.

Our discovery that the junctional localization of Sstn has partial dependence on Step indicates that their relationship is more complicated than the recruitment of an enzyme by a localized adaptor. This additional complexity may also explain the requirement of the Sstn CC domain for full recruitment of Sstn to LE junctions, and the substoichiometric junctional levels of Sstn relative to Step. We suggest two additional effects. First, Step and Sstn may both be more stable as Step-Sstn heterodimers than as individual proteins. Although isolated pieces of Step and Sstn

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**FIGURE 7:** Step and Sstn promote each other’s junctional localization. (A) sstn mutants were compared with control embryos both expressing the same dosage of step^GFP^ The embryos were fixed and imaged at dorsal closure. A strong reduction of junctional Step-GFP was observed in the mutants, although a slight signal was detected along the epidermal LE (arrows). All images were taken at the same settings, and the observed pattern was consistent for comparisons of 9/9 mutant embryos and 17/17 control embryos from two experiments. DE-cad staining revealed AJ positioning and also showed distorted organization of the sstn mutant epidermis, including the abnormally frequent presence of multicellular rosettes (shaded in yellow and quantified to the right with embryo numbers in brackets).

(B) Antibody staining of endogenous Step also revealed its loss from the epidermal LE (arrows) of sstn mutants vs. sibling controls in fixed dorsal closure embryos. All images were taken with the same settings, and the observed pattern was consistent for comparisons of 8/8 mutant embryos and 16/16 control embryos from two experiments. DE-cad staining revealed AJ positioning and also showed distorted organization of the sstn mutant epidermis (multicellular rosettes shaded in yellow and quantified to the right with embryo numbers in brackets).

(C) Live imaging of Sstn-GFP revealed substantial loss from LE (arrows) and other junctions of the epidermis of step mutant embryos vs. sibling controls. LE junction levels are quantified at right. Two replicate comparisons are shown in different colors, and for each replicate the data were normalized to the average embryo values for the Sstn-GFP signal at LE junctions of the sibling controls. Embryo numbers are indicated in brackets.

(D) Sstn-GFP was also observed to be lost from DE-cad–positive AJs of step mutants of a distinct genotype, in contrast to costained and comounted sibling controls.
(the Step CC and the Sstn CR) can localize to junctions when overexpressed, stabilized complexes might promote and/or maintain full junctional localization of Step and Sstn when expressed at endogenous levels. Endogenous Sstn would presumably provide adaptor function as part of such stabilized heterodimers. Second, additional proteins may contribute to Step localization. Indeed, it was recently reported that Ajuba, a tension-induced partner of junctional α-catenin (Yap et al., 2018), 1) colocalizes with junctional Step and Sstn, 2) is required for recruitment of overexpressed Step to AJs, and 3) affects junctional myosin and tissue organization in ways very similar to Step and Sstn (Rauskolb et al., 2019). Because no other portion of Step was sufficient for its AJ localization in the absence of its CC domain, interactions with additional proteins may involve the CC domain, an idea consistent with the ability of CC-mediated interactions to support trimers, tetramers, or even heptamers (Grigoryan and Keating, 2008). It is important to note that these two proposals are not mutually exclusive. During Drosophila embryo development, it is tempting to speculate that the first mechanism might dominate for actomyosin regulation in the syncytial embryo, when epithelial cells, and thus AJs, are absent. The second mechanism may then arise during gastrulation when actomyosin-associated AJs are subjected to tension.

Our study is significant for defining a molecular mechanism that recruits Step to myosin-rich junctions for an established role in regulating actomyosin activity at these sites (West et al., 2017). Tension-based conformational change of α-catenin also recruits actin-binding proteins to reinforce junction–actomyosin connections (Leckband and de Rooij, 2014; Pinheiro and Bellaïche, 2018; Yap et al., 2018). These mechanisms may work together to modify both the generation of junctional tension and resistance to it.

**MATERIALS AND METHODS**

**Previously established Drosophila lines**

The following alleles were used: sstrn^{Df(3R)USBC411} (Bloomington Drosophila Stock Center [BDSC] #24945); sstrn^{Df(3R)USBC415} (Exelixis; Stock #c04515); step^{Df(3L)USBC150} (BDSC; #9509); step^{Df(3R)USBC108710} (gifts from M. Hoch, Life and Medical Science Institute of Bonn, Germany); zip^{GFP} (FlyTrap; Stock #CC01626; BDSC; #51564).

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**FIGURE 8:** Abilities of Step constructs to rescue the tissue distortions of step mutant embryos. In a step mutant background, the indicated UAS constructs were expressed in epidermal stripes using paired-GAL4. The constructs and the AJ marker Canoe were imaged at dorsal closure, and quantifications of multicellular rosettes were compared between construct-expressing and –nonexpressing tissue. Example rosettes are labeled in cyan. Quantifications to the right of each pair of images are from two separate replicates, and rosette quantifications of construct-expressing and –nonexpressing tissue of the same embryo are connected with a line. Rescue of the abnormal rosettes was evident for expression of GFP-Step (top left) and of GFP-StepCC+CR (bottom right), but not for expression of GFP-StepΔCC+CC[N*] (bottom left) or of GFP-StepΔPH (top right).
The following UAS constructs were used: UAS-GFP at attP2 (gift of U. Tepass, University of Toronto, Canada); UAS-GFP-Sttn at attP40 (Liu et al., 2015); UAS-GFP-SttnΔCR at attP40 (Liu et al., 2015); UAS-mCherry-Sttn at attP2 (Liu et al., 2015); UAS-GFP-Step (RNA interference [RNAi]-resistant) at attP40 (Lee and Harris, 2013); UAS-GFP-StepΔCC at attP40 (Liu et al., 2015); UAS-GFP-StepΔPH (RNAi-resistant) at attP40 (Lee et al., 2015).

UAS constructs were driven using either daughterless-Gal4 (gift of U. Tepass) or paired-Gal4 (BDSC, #1947).

The following construct was also used: tGPH (BDSC #8163). It was described as GFP-StepPH+PBalone.

**Generated UAS constructs**

For StepCCalone-GFP, the Step CC domain (amino acids [aa] 25–70) was PCR amplified from the step isoform A cDNA (RE34385; Canadian Drosophila Microarray Centre) using 5′-CCGGAAC-CAATTCGAGCTGACATGCCAGAATCTACCAAGG-3′ (forward) and 5′-AAAGCTGGTCTAGATCTGAGTGGGGAACATCCAG-3′ (reverse) primers (all primers from Life Technologies), and cloned into the pENTR vector using Sall and Xhol restriction sites. Gateway cloning (Life Technologies) recombined the construct into pPWG downstream of a UASP promoter for C-terminal enhanced GFP tagging. An attB recombination site cloned into the pPWG NsiI site targeted the vector to the attP40 recombination site on chromosome 2 (BestGene).

For GFP-SttnCRalone, the Sttn CR (aa 576–635) was PCR amplified from the sttn cDNA (RE36140; Drosophila Genomics Resource Center) using 5′-CAGTCGACTGATTTGAACCATCCAC-3′ (forward) and 5′-GGGTCTAGATCTGAGGAGAAGGG-CCATGGCATCCAG-3′ (reverse) primers (all primers from Life Technologies), and cloned into the pENTR vector using Sall and Xhol restriction sites. Gateway cloning (Life Technologies) recombined the construct into pPwg and then the hybrid construct was recombined into pPGW. These final constructs were each targeted to the attp40 recombination site in transgenic flies (BestGene). For GFP-StepΔCC+CR, the Sttn CR was altered to change its first seven leucines and isoleucines to serine residues (as in Figure 3B). For GFP-StepΔCC+APH, the sequence encoded the APH synthetic CC domain (MKQELKQKELKQKQAEKOLQWQLKWQARKKKQLQKQLKAQ) with a 3′ proline as a linker; Gurnon et al., 2003). For GFP-StepΔCC+APH/K42, the sequence encoded VSSLESK repeated six times (Graddis et al., 1993). For GFP-StepΔCC+CR, the sequence encoded the Sttn CR (aa 576–635).

**Generated GFP-tagged sttn allele**

The CRISPR-Cas9 tagging of Sttn with GFP was based on an existing protocol (Harris et al., 2016).

The NEBuilder HiFi DNA Assembly Cloning Kit (New England Biolabs) connected three pieces: 1) a Phusion high-fidelity DNA polymerase (New England Biolabs) PCR amplification of a portion of the sttn genomic DNA Bac clone (BACPAC resources center; #BACR18G16) covering sequence 5′ of the stop codon and replacing the stop codon with a short sequence overlapping with the 5′ of the GFP sequence (forward primer, 5′-CGCGGACATATG-CACCTGCTAAATATAGCTAG-3′; reverse primer, 5′-CCTTGCT-CACCATTTAGATCTGCCTTCT-3′); 2) a Phusion high-fidelity DNA polymerase PCR amplimer of the sequence of GFP plus overlapping ends with the sttn construct and the pDSRed vector sequence (forward primer, 5′-AGGGCGGAATCTCTAT-ATATGAGGAACATTGCCC-3′; reverse primer, 5′-AGGTTGGCCAC-TAGCATTCATTTGCCTTCT-3′; template, PPWG vector); and 3) the pDSRed vector (gift of K. O’Connor Giles, University of Wisconsin; forward primer CATGGACAGCTGTAACATGATCTGACTGATTGCCC; reverse primer 5′-TTAGCAGGTTGTCAATGCTCGCCG-GCG-3′).

For the second construction step, a 1 kb sequence of sttn 3′ to the stop codon was synthesized (GenScript) based on the sttn genomic DNA Bac clone (BACR18G16) with restriction enzyme sites of BglII and Xhol added at the ends. This synthesized DNA was ligated with the first assembled construct, which was cut with BglII and Xhol.

The g-RNA sequence was selected with an online tool (http://tools.flycrispr/molbio.wisc.edu/targetFinder/) and contained a PAM site close to the sttn stop codon (g-RNA sequence, 5′-GCCACTTTCCAGCCGTACAAAAGGTACGAAAACC-3′). To generate the g-RNA construct, the g-RNA sequence was ligated into the pCDf5 vector (gift of K. O’Connor Giles). The corresponding PAM site in the donor construct was mutated from 5′-GGCCACTTTCCAGCCGTACAAAAGGTACGAAAACC-3′ to 5′-GGCCACTTTCCAGCCGTACAAAAGGTACGAAAACC-3′. The complete donor construct was sequenced for confirmation.

Both the g-RNA and the donor construct were injected into yw; nos-Cas9 III-attP2 flies. The injection and subsequent screening, Cas9 removal, and genetic balancing were done by Bestgene (Plan RI). Viable and fertile homozygous flies were utilized for the study with the insertion confirmed by PCR and sequencing.

**Generated GFP-tagged step allele**

The CRISPR-Cas9 tagging of Step with GFP was based on an existing protocol (Harris et al., 2016).

The NEBuilder Hifi DNA Assembly Cloning Kit (New England Biolabs) was used to generate the donor construct in two steps. The first step included three pieces: 1) 1 kb upstream of the stop codon was amplified (forward primer, 5′-GGGCACATATGCA-CACCTGCACACTATGCTAT-3′; reverse primer, 5′-CCCTGCTCACCATACTTTGCTGAGTGCCTTT-3′) with the template of step genomic DNA (BAC09H20, BACPAC resources center); 2) the g-RNA sequence with a new stop codon incorporated (forward primer, 5′-GGCACTCGCAAGATGTTGTGACAAAGGC-3′; reverse primer, 5′-GGGCACATATGCA-CACCTGCACACTATGCTAT-3′) with the template of the PPWG vector; and 3) the pDSRed vector (gift of K. O’Connor Giles, University of Wisconsin; forward primer CATGGACAGCTGTAACATGATCTGACTGATTGCCC; reverse primer 5′-CATCATGTATGCAATGCTCGCCG-3′).

The second step for assembling the donor construct amplified 1 kb downstream of the stop codon (forward primer, 5′-GCTTAGGCCTTCTGCAGAATCAACAGCAGC-3′; reverse primer, 5′-AGCCTGACGGGCCAAATTATACATCTTCT-3′; from the step genomic DNA). This downstream sequence was then assembled into the vector from the first step (forward primer 5′-GTATA-AATTGGCACGCTGAGGCTTCTTC-3′, reverse primer 5′-GATTCTGATTACTGCGAAGGCGTAC-3′).
The g-RNA was found using an online tool (http://tools.flycrispr .molbio.wisc.edu/targetFinder/) with a PAM site near the step stop codon (g-RNA sequence: 5′-CGGAGGAAGATCAGCAGGAG-3′). To generate the g-RNA, the sequence was ligated into the BbsI site of the pCDf3 vector (gift of K. O’Connor Giles, University of Wisconsin). The corresponding PAM site in the donor construct was mutated from 5′-GCAATCCGCACTTTACGAGGAAGATCAGC-3′ to 5′-GCTGATCTCCTCCGTAACAGCGCGACATGC-3′. The complete donor construct was sequenced for confirmation.

Both the g-RNA and the donor construct were injected into yw;; nos-Cas9 III-attp2 flies. The injection and subsequent screening, Cas9 removal, and genetic balancing were done by Bestgene (Plan RI). Viable and fertile homozygous flies were utilized for the study with the insertion confirmed by PCR and sequencing.

**Synthesized lines**

For complex genotypes with alleles on separate chromosomes, standard *Drosophila* genetics was used for synthesis. For the recombinant chromosomes of step*Df(2L)BSC150* with UAS constructs, PCR was used to confirm the presence of step*Df(2L)BSC150* by amplifying the opposite flanking regions of the deletion, and the UAS constructs were detected using a linked minixwhite cassette.

**Antibody staining of embryos**

Embryos were collected from apple juice agar plates (25 g agar; 250 ml store-bought apple juice; 12.5 g store-bought granulated white sugar; 10 ml, 10% Tegosept [in ethanol]) using a brush into 0.1% Triton X-100, and washed three times. Dechorionation was performed with 50% bleach for 4 min, and the embryos were then washed another three times with 0.1% Triton X-100. Fixation was performed using 1:1 3.7% formaldehyde in phosphate-buffered saline (PBS):heptane for 20 min. Embryos were devitel-linized by shaking in methanol and rinsed three times afterward with methanol. Blocking and staining were conducted with PBS with 1% goat serum, 1% sodium azide, and 0.1% Triton X-100. Primary antibodies were guinea pig anti-Step (1:350; gift of M. Hoch), rabbit anti-Canoe (1:1000; gift of M. Peifer, University of North Carolina, Chapel Hill), and rat anti–DE-cadherin (1:100; Developmental Studies Hybridoma Bank). Secondary antibodies were conjugated to Alexa Fluor 546 and 647 (Life Technologies). Mounting of fixed and immunostained embryos was with Aqua Polymount (Polysciences).

**Live imaging preparation**

Embryos for live imaging were collected from apple juice agar plates using a brush into 0.1% Triton X-100, and washed three times. Dechorionation was performed with 50% bleach for 4 min, and the embryos were then washed another three times with 0.1% Triton X-100. Embryos were then mounted on a glass coverslip using tape glue dissolved in heptane and overlaid with halocarbon oil (series 700; Halocarbon Products). The coverslip, with the embryos facing up, was set into the bottom of a glass-bottom culture dish with its original coverslip removed.

**Embryo imaging**

Most imaging was done using a spinning disk confocal microscope (Quorum Technologies) with a Hamamatsu EM CCD camera. Image capture was performed using Velocity software (PerkinElmer). The imaging took place at room temperature at 63x magnification (Plan-Apochromat; NA 1.4). Samples were mounted on a piezo stage and z-stacks were imaged with 0.3 µm spacing.

Whole embryo imaging of Figure 8 was performed with a Leica TCS SP8 confocal system at RT with a 40× NA 1.4 objective (Leica), and 300 nm step sizes.

**Image quantification and manipulation**

For local protein levels within each embryo, there were three junctions of interest: 1) LE junctions, 2) tricellular junctions within the lateral epidermal sheet, and 3) A-P bicellular junctions within the lateral epidermal sheet. Any cell at least one cell away from the LE cells was considered part of the lateral epidermis. Per embryo, each junction type was measured from five separate cells showing the highest signal intensity. For each measurement, a small circle with was drawn over the center of the junction of interest and the mean fluorescence was acquired using Fiji (Schindelin et al., 2012). To correct the measurement for background, signal in the cytoplasm of the cell was also measured using the same method and the mean fluorescence was subtracted from the junction measurement. Means of the five background-corrected measurements of each junction type were calculated for each embryo, resulting in one value per junction type per embryo. These single junction values were then compared to calculate junction:junction ratios within each embryo.

For epithelial rosette identification and quantification, the methodology from West et al. (2017) was used. Rosette structures were considered as five or more cells with a common vertex within a square area of 1.6 µm².

Statistical significance for pairwise comparisons was calculated using Student’s t tests in Excel (Microsoft).

**Blot overlays**

For glutathione S-transferase (GST)-StepCC, the Step CC domain was PCR amplified using 5′-CCCTGAGATCCCAGGAGC-CAAAAAAC-3′ (forward) and 5′-ATGCGGCCGCTCGAGTAAAGG-CACTCCAG-3′ (reverse) primers, and cloned into the pGEX-6p2 vector using BamHI and Xhol restriction sites. For GST-StnCC, the Stsn CC domain (aa 60–87) was PCR amplified using 5′-GGCCCCCTGGATCCTCCAAGGTTAGAAG-3′ (forward) and 5′-GGCCGTCGATGTTAGGTTAGTAATTG-3′ (reverse) primers, and cloned into the pGEX-6p2 vector using BamHI and Xhol restriction sites.

For maltose-binding protein (MBP)-StepCC, the Step CC domain was PCR amplified using 5′-CAGAATTCCGATCCCAGGAG-CAGCAAAAAC-3′ (forward) and 5′-CAGTCCCAAGCTTTAGGGCAC- CATCC-3′ (reverse) primers, and cloned into the pMal-C2X vector using BamHI and HindIII restriction sites. For MBP-StnCC, the Stsn CC domain was PCR amplified using 5′-CAGAATTCCGATCCCAGGAG-CAGCAAAAAC-3′ (forward) and 5′-CAGTCCCAAGCTTTAGGGCAC- CATCC-3′ (reverse) primers, and cloned into the pMal-C2X vector using BamHI and HindIII restriction sites.

For MBP-tagged Step mutant CC domains, the specific mutant fragments were synthesized (GenScript) and inserted into pMalC2X with BamHI N-terminal sites and HindIII C-terminal sites.

Using the supplier’s instructions, the MBP and GST fusion protein constructs were inoculated in BL21 competent cells and induced using 0.1 mM isopropyl-β-D-thiogalactoside (IPTG) (final concentration). The cells were pelleted at 3500 × g and 4°C and lysed with 20 ml PBS-T (PBS with 1% Triton X-100) with 2 µg/ml aprotinin, 1 µg/ml leupeptin, 25 µg/ml phenylmethylsulfonyl fluoride, and 1 mg/ml lysozyme. The mixture was incubated at room temperature for 40 min and centrifuged at 12,000 × g for 20 min at 4°C. The supernatant was run through the purification column with either Pierce glutathione agarose (Thermo Scientific) for GST fusion
proteins or amylose resin (New England Biolabs) for MBP fusion proteins. Following washes, the proteins were eluted with either GST elution buffer (10 mM glutathione, 50 mM Tris-HCl, pH 9.0, and 15% glycerol) or MBP elution buffer (10 mM maltose, 200 mM NaCl, 20 mM Tris-HCl, 1 mM EDTA, 1 mM dithiothreitol [DTT], pH 7.4).

Purified GST and MBP fusion proteins were first individually tested using GST and MBP antibodies (New England Biolabs) on diagnostic Western blots. For blot overlays, MBP fusion proteins (1–3 µg) were separated with 10% SDS–PAGE and individually blotted on nitrocellulose. Blocking was done with 5% milk in TBST buffer (8.0 g of NaCl, 2.42 g of Tris, and 1 ml of Tween-20 in 1 l of double-distilled H2O, pH adjusted to 8.0). Blots were incubated with each GST fusion protein (–7 µg) in TBST buffer and 5% milk, washed, incubated with GST antibodies (raised in rabbit–lab generated) in TBST buffer and 5% milk, washed, incubated with horseradish peroxidase–conjugated goat anti-rabbit antibodies, washed, and incubated with chemiluminescent reagents (Thermo Fischer Scientific). Imaging of the blots was done on a ChemiDoc XR (Bio-Rad). For each binding assay, control and noninteracting proteins were adjusted so they were present at higher molar levels than interacting proteins (levels were evaluated based on diagnostic Western blots).

Protein structure prediction and visualization
The Step CC structure was predicted with the Phyre2 server (Kelley et al., 2015). After conversion to .c3d file format with the National Center for Biotechnology Vast Search Tool (Madej et al., 2014), the file was analyzed with the CN3D viewer to identify hydrophobic residues in the 3D structure (Wang et al., 2000).

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