Dynamic myosin phosphorylation regulates contractile pulses and tissue integrity during epithelial morphogenesis

Claudia G. Vasquez, Mike Tworoger, and Adam C. Martin

Department of Biology, Massachusetts Institute of Technology, Cambridge, MA 02142

A

pical constriction is a cell shape change that promotes epithelial bending. Activation of non-muscle myosin II (Myo-II) by kinases such as Rho-associated kinase (Rok) is important to generate contractile force during apical constriction. Cycles of Myo-II assembly and disassembly, or pulses, are associated with apical constriction during Drosophila melanogaster gastrulation. It is not understood whether Myo-II phosphorylation organizes contractile pulses or whether pulses are important for tissue morphogenesis. Here, we show that Myo-II pulses are associated with pulses of apical Rok. Mutants that mimic Myo-II light chain phosphorylation or depletion of myosin phosphatase inhibit Myo-II contractile pulses, disrupting both actomyosin coalescence into apical foci and cycles of Myo-II assembly/disassembly. Thus, coupling dynamic Myo-II phosphorylation to upstream signals organizes contractile Myo-II pulses in both space and time. Mutants that mimic Myo-II phosphorylation undergo continuous, rather than incremental, apical constriction. These mutants fail to maintain intercellular actomyosin network connections during tissue invagination, suggesting that Myo-II pulses are required for tissue integrity during morphogenesis.

Introduction

Epithelial morphogenesis is critical for organs and embryos to change shape during development (Leptin, 2005; Heisenberg and Bellaïche, 2013). A cell shape change that often accompanies epithelial remodeling is apical constriction, which promotes epithelial sheet bending and cell invagination (Sawyer et al., 2010; Martin and Goldstein, 2014). One example of apical constriction–induced tissue remodeling occurs in Drosophila melanogaster gastrulation. It is not understood whether Myo-II phosphorylation organizes contractile pulses or whether pulses are important for tissue morphogenesis. Here, we show that Myo-II pulses are associated with pulses of apical Rok. Mutants that mimic Myo-II light chain phosphorylation or depletion of myosin phosphatase inhibit Myo-II contractile pulses, disrupting both actomyosin coalescence into apical foci and cycles of Myo-II assembly/disassembly. Thus, coupling dynamic Myo-II phosphorylation to upstream signals organizes contractile Myo-II pulses in both space and time. Mutants that mimic Myo-II phosphorylation undergo continuous, rather than incremental, apical constriction. These mutants fail to maintain intercellular actomyosin network connections during tissue invagination, suggesting that Myo-II pulses are required for tissue integrity during morphogenesis.

Myo-II is clearly involved in apical constriction, the mechanism of apical actomyosin contraction remains poorly understood. Myo-II activity is regulated by phosphorylation of its regulatory light chain (RLC; sqh in Drosophila), which promotes Myo-II oligomerization into minifilaments and activation of the Myo-II motor ATPase activity (Karess et al., 1991; Sellers, 1991; Jordan and Karess, 1997). An effector of the RhoA GTPase (Rho1 in Drosophila), Rho-associated coiled-coil kinase (or Rho-associated kinase [Rok] in Drosophila), can phosphorylate and activate Myo-II via direct RLC phosphorylation or by inhibiting the phosphatase that dephosphorylates the RLC (myosin phosphatase; Amano et al., 1996; Kimura et al., 1996; Kawano et al., 1999; Winter et al., 2001). In the Drosophila mesoderm, the transcription factors Twist and Snail are thought to promote apical constriction and apical Myo-II accumulation by activating Rho1 and its effectors, including Rok (Barrett et al., 1997; Hacker and Perrimon, 1998; Dawes-Hoang et al., 2005; Fig. S1 A). Mutants or chemical inhibition of Rok results in loss of apical

© 2014 Vasquez et al. This article is distributed under the terms of an Attribution-Noncommercial-Share Alike-No Mirror Sites license for the first six months after the publication date (see http://www.rupress.org/terms). After six months it is available under a Creative Commons License Attribution-Noncommercial-Share Alike 3.0 Unported license, as described at http://creativecommons.org/licenses/by-nc-sa/3.0/
after a pulse is not stabilized, resulting in cell shape fluctuations and inefficient constriction (Martin et al., 2009, 2010). Thus, one model for ventral furrow formation is that cells undergo ratcheted apical constriction, whereby pulses drive constrictions that are stabilized via a Twist-dependent mechanism for efficient apical constriction. We proposed that the polarized localization of Rok to medioapical foci results in the persistence of medioapical actomyosin fibers that stabilize cell shape between pulses; we have named this cellular organization radial cell polarity (Mason et al., 2013). The persistence of medioapical Myo-II fibers is important for the formation of a supracellular Myo-II meshwork that transmits tension across the ventral furrow tissue (Martin et al., 2010). However, the roles of Rok and Myo-II phosphorylation during pulsatile constriction are still unclear. Furthermore, it is not known why cells undergo pulsatile rather than continuous contraction to promote tissue morphogenesis.

Myo-II and lack of apical constriction, demonstrating that Rok is necessary for cortical Myo-II localization in the ventral furrow (Dawes-Hoang et al., 2005; Mason et al., 2013). In addition, mutants that mimic Myo-II RLC phosphorylation progress through development and suppress rok mutants, suggesting that Rok increases levels of active Myo-II to promote actomyosin contractility and morphogenesis (Jordan and Karess, 1997; Winter et al., 2001; Royou et al., 2002). It is not clear, however, whether coupling of Rok activity to Myo-II activation plays a role in organizing apical actin network contraction.

We have developed a system to visualize the dynamics of actomyosin contraction and apical constriction during Drosophila gastrulation. In ventral furrow cells, Myo-II undergoes cycles of assembly in the center of the apical surface (medioapical cortex) followed by disassembly or remodeling, which we refer to as pulses (Martin et al., 2009; Fig. 1A). Phases of rapid apical constriction are associated with pulses of Myo-II assembly and the coalescence of Myo-II and F-actin structures into medioapical foci, which possibly represent contractions of the actin cortex (Martin et al., 2009; Roh-Johnson et al., 2012). After Myo-II coalescence, Myo-II structures remodel by either disassembling or changing in morphology. Despite Myo-II remodeling, cell shape is often stabilized between pulses, resulting in incremental apical constriction. Although Myo-II still undergoes pulsing in Twist knockdowns, the contracted cell shape after a pulse is not stabilized, resulting in cell shape fluctuations and inefficient constriction (Martin et al., 2009, 2010). Thus, one model for ventral furrow formation is that cells undergo ratcheted apical constriction, whereby pulses drive constrictions that are stabilized via a Twist-dependent mechanism for efficient apical constriction. We proposed that the polarized localization of Rok to medioapical foci results in the persistence of medioapical actomyosin fibers that stabilize cell shape between pulses; we have named this cellular organization radial cell polarity (Mason et al., 2013). The persistence of medioapical Myo-II fibers is important for the formation of a supracellular Myo-II meshwork that transmits tension across the ventral furrow tissue (Martin et al., 2010). However, the roles of Rok and Myo-II phosphorylation during pulsatile constriction are still unclear. Furthermore, it is not known why cells undergo pulsatile rather than continuous contraction to promote tissue morphogenesis.

Here, we combined live imaging and quantitative analysis of Myo-II regulators and GFP-tagged Myo-II RLC phosphomutants to define the role of coupling Myo-II activation to upstream signals during contractile pulses. We show that temporal and spatial control of Myo-II phosphorylation organizes contractile pulses. We suggest that the pulsatile nature of apical constriction is required for the stable transmission of intercellular forces during tissue morphogenesis.
Results

Myo-II pulses correlate with fluctuations in apical Rok localization
Contractile pulses during ventral furrow cell apical constriction are associated with the coalescence and increase in apical Myo-II intensity, after which apical Myo-II structures are remodeled (Fig. 1 A; Martin et al., 2009). Because Rok is required for cortical Myo-II localization in the Drosophila embryo (Royou et al., 2002; Dawes-Hoang et al., 2005), we examined whether dynamic Myo-II localization was correlated with changes in Rok localization. Before gastrulation, Rok localizes to basal furrow canals and is relocalized to foci in the center of the apical domain during apical constriction (Mason et al., 2013; Fig. 1 B). At the onset of apical constriction, we found that Rok intensity exhibited clear fluctuations, with medioapical Rok foci appearing and disappearing in association with Myo-II coalescence (Fig. 1, C and D; and Video 1). Analysis of the time-resolved cross-correlation between Myo-II and Rok intensity during individual pulses demonstrated a significant correlation that peaked at 0 s offset (Fig. 1 E). Thus, Myo-II foci appear at the same time as Rok foci. Additionally, apical Rok intensity increased during apical constriction, often in bursts that corresponded with phases of rapid constriction (Fig. S1 B). The pulsatile behavior of Rok during ventral furrow formation suggests that dynamic changes in Myo-II phosphorylation by Rok direct contraction pulses that result in incremental apical constriction.

RLC mutants that mimic mono- or diphosphorylation result in constitutive Myo-II oligomerization independent of Rok
To test whether dynamic Myo-II phosphorylation, and thus changes in Myo-II minifilament assembly and motor activity, is required for Myo-II pulses, we generated GFP-tagged sqh mutants affecting threonine-20 and serine-21 (Fig. 2 A; Jordan and Karess, 1997). Alanine substitutions in these residues decrease Myo-II motor ATPase activity in vitro (Kamisoyama et al., 1994), and the sqh-AA mutant resembles a sqh null allele in vivo (Jordan and Karess, 1997). Conversely, glutamate substitutions activate Myo-II minifilament assembly and ATPase activity in the absence of phosphorylation in vitro, although the ATPase activity of phosphomimetic myosin mutants is lower than that of phosphorylated Myo-II (Kamisoyama et al., 1994). We differentiated between possible roles of mono- and diphosphorylation by generating mutants that trap the Myo-II motor in all possible combinations of RLC phosphorylation states that are likely present in vivo (Fig. 2 B). The sqh phosphomutants were expressed at similar levels as endogenous sqh and did not exhibit gastrulation phenotypes in the presence of the endogenous Sqh (Figs. S1 C and S2 A). We reduced endogenous Sqh protein levels ~90% by making germline clones with the hypomorphic sqh’ allele (Fig. S1, C and D). The wild-type sqh transgene (sqh-TS::GFP), but not the inactive sqh-AA::GFP allele, rescued nuclear migration and the uneven celluarization phenotypes of sqh’ mutants (Fig. 2 C; Wheatley et al., 1995; Royou et al., 2002, 2004). In addition, we obtained gastrulating embryos from all mutants, including the sqh-AA allele. Thus, the partial activity of the sqh’ allele enabled us to examine the function of sqh phosphomutants during gastrulation.

We first determined whether the different sqh phosphomimetic mutants activate Myo-II minifilament assembly in vivo. During cellularization, Myo-II localizes to basal furrow canals and cytoplasmic aggregates (Royou et al., 2004). Although we did not observe ectopic cortical Myo-II assembly, phosphomimetic sqh mutants, particularly sqh-AE and sqh-EE, accumulated Myo-II in aggregates that are immediately basal to the furrow canals (Fig. 2 C and Fig. S1 E). Myo-II aggregates were reduced in size when wild-type levels of endogenous Sqh were present and Myo-II aggregates also contained the myosin heavy chain (Drosophila zipper), suggesting that these aggregates are not denatured protein (Fig. 2, C and D). Importantly, Sqh aggregates do not contain F-actin, suggesting that the Sqh aggregates do not form by actomyosin contraction, but could result from constitutive Myo-II minifilament assembly (Fig. 2 E). The phosphomimetic mutants resulted in a decrease in cytoplasmic Myo-II levels relative to wild-type embryos, suggesting that RLC mutants that mimic mono- or diphosphorylation shift the Myo-II complex toward the active conformation in vivo, promoting constitutive oligomerization (Fig. 2 F). Thus, our GFP-tagged phosphomimetic RLC mutants appear to uncouple activation of Myo-II oligomerization from phosphorylation by Rok.

RLC phosphorylation dynamics do not trigger changes in Myo-II apical–basal localization
Apical activation of Rok has been proposed to localize Myo-II assembly during apical constriction (Dawes-Hoang et al., 2005). Thus, mutants that uncouple Myo-II activity from phosphorylation by Rok might disrupt the apical–basal localization of Myo-II. Therefore, we tested whether the apical–basal polarity of Myo-II localization is controlled by RLC phosphorylation and oligomerization. Mutants that mimic mono- or diphosphorylation localized specifically to the apical surface of ventral furrow cells, despite the presence of Myo-II aggregates throughout the entire embryo during cellularization (Fig. 3, A and B; Fig. S2 B; and Video 2). Furthermore, Myo-II was reduced in basal furrow canals during ventral furrow invagination in phosphomimetic mutants, similar to wild-type Myo-II, suggesting that diphosphorylation of the RLC is not the mechanism of basal Myo-II loss (Fig. 3, A and B). Inactivation of the RLC phosphorylation sites similarly did not disrupt apical Myo-II localization (Fig. 3 C, Fig. S2 B, and Video 2). These results demonstrate that despite the tissue-specific apical recruitment of Rok in ventral furrow cells, regulation of oligomerization does not instruct apical Myo-II localization at the onset of gastrulation.

Phosphomimetic RLC mutants disrupt polarized actomyosin condensation
Mediaopical Myo-II pulses involve the coalescence or condensation of both Myo-II and F-actin into medioapical foci (Martin et al., 2009; Mason et al., 2013). Because Rho1 and Rok are also enriched in medioapical foci, we have proposed that the apical domain of ventral furrow cells exhibits radial cell polarity, with protein localization differing between the apical center
Figure 2. Characterization of sqh phosphomutant oligomerization. (A) Phosphorylation of the Drosophila Myo-II RLC sqh on threonine-20 and serine-21 is predicted to activate motor activity and promote bipolar minifilament assembly. (B) Chart summarizing phosphorylation states available to different Sqh phosphomutants. (C) Sqh mutants that mimic phosphorylation result in cytoplasmic Myo-II aggregates. Representative images of XY semisagittal sections and YZ cross sections for live cellularizing embryos. Dashed lines highlight evenness (sqh-TS) or unevenness (sqh-AA) of cellularization front. Arrowheads indicate examples of cytoplasmic Myo-II aggregates. Bar, 20 µm. (D) Basal Sqh aggregates colocalize with Zipper aggregates. Images are fixed embryos.
to the junctions (Mason et al., 2013). To test whether dynamic regulation of RLC phosphorylation polarizes actomyosin within the apical cortex, we combined sqh::GFP transgenes with a membrane marker fused to mCherry (Memb::Cherry) to visualize and quantify the apical organization of Myo-II. Because we have been unable to obtain viable embryos expressing Memb::Cherry and sqh-EE::GFP, which exhibits additional oogenesis defects associated with constitutive Myo-II activation, we focused our quantitative analysis on the sqh-AE mutant. We observed similar Myo-II localization and mutant phenotypes between sqh-AE and sqh-EE, consistent with these mutants similarly activating Myo-II oligomerization. The sqh-TS::GFP transgene rescued Myo-II coalescence in sqh1 germline clone mutants, forming medioapical foci during pulses and thus exhibiting radial cell polarity in Myo-II accumulation (Fig. 4 A). In contrast, Myo-II in the sqh-AE mutant did not exhibit this medioapical enrichment, indicating a defect in medioapical Myo-II coalescence (Fig. 4 D). The defect in the medioapical polarization of Myo-II was specific to phosphomimetic mutants because Myo-II in both the sqh-AA and sqh-TA mutants accumulated as concentrated foci and the sqh-TA mutant exhibited medioapical F-actin condensation (Fig. 4, C, D, and G; and Videos 2 and 3). Thus, the defects in actomyosin coalescence observed in the sqh-AE mutants suggest that coupling of Myo-II activation to Rok phosphorylation is critical to polarize actomyosin contractile activity within the apical domain of ventral cells.

Because Myo-II localization in the phosphomimetic mutants could be influenced by the low (10%) levels of endogenous Sqh, we sought to fully uncouple Myo-II activation from upstream signals by activating Myo-II in a rok mutant background. Either

stained for Zipper (myosin heavy chain) and endogenous GFP. Bar, 10 µm. (E) Basal Sqh aggregates do not colocalize with F-actin. Images are fixed embryo cross sections during sequential stages of apical constriction stained for Zipper (myosin heavy chain) and neurotactin (membrane). (B and C) Sqh phosphomutants do not alter the basal–apical Myo-II redistribution. Embryo preparation was the same as in A, except germline clones expressing sqh-AE (B) and sqh-TA (C) were fixed. Bars, 20 µm.

Figure 3. Dynamic RLC phosphorylation is not required to trigger basal to apical relocalization of Myo-II. (A) Wild-type Myo-II localizes to the apical surface and is reduced basally during ventral furrow formation. Images are fixed embryo cross sections during sequential stages of apical constriction stained for Zipper (myosin heavy chain) and neurotactin (membrane). [B and C] Sqh phosphomutants do not alter the basal–apical Myo-II redistribution. Embryo preparation was the same as in A, except germline clones expressing sqh-AE (B) and sqh-TA (C) were fixed. Bars, 20 µm.
Figure 4. Polarized actomyosin condensation requires dynamic Myo-II phosphorylation. (A–C) Uncoupling Myo-II activation from Rok phosphorylation disrupts medioapical Myo-II coalescence. Time-lapse images are representative cells from sqh1 germline clone embryos expressing the indicated phosphomutants and Memb::Cherry (membrane). Bars, 5 µm. (D) Quantification of Myo-II medioapical polarity, which is the difference between medioapical Myo-II (light green in schematic) and junctional Myo-II (green in schematic) intensities normalized by the total Myo-II intensity in a cell. Error bars are SEM (n = 120 sqh-TS cells, two embryos; n = 100 sqh-AE cells, two embryos; n = 118 sqh-TA cells, two embryos). (E–G) The sqh-AE mutant fails to condense apical F-actin. Time-lapse images are of representative cells from sqh1 germline clone embryos expressing the indicated phosphomutants and Utr::mCherry (F-actin). Arrowheads indicate Myo-II and F-actin condensation. Cell outlines in yellow were made by manually segmenting the subapical Utr::mCherry signal 2 µm below the apical meshwork. Bars, 5 µm. *, P < 0.05; **, P < 0.01.
rok1 or rok2 mutant germline clones result in a loss of Myo-II from the cortex, consistent with Rok and Myo-II phosphorylation being required for Myo-II activation (Fig. S3 and Video 4; Royou et al., 2002; Dawes-Hoang et al., 2005; Mason et al., 2013). Previous studies suggested that sqh-EE can suppress loss of Rok function (Winter et al., 2001; Bertet et al., 2004), suggesting that Rok activates Myo-II minifilament assembly, whereas minifilament localization is dependent on other cortical cues. Consistent with these results, we found that sqh-EE rescued cortical Myo-II localization in rok1 and rok2 mutant germline clones (Fig. S3). However, despite rescuing cortical localization in rok germline clones, sqh-EE Myo-II failed to undergo medioapical coalescence and generate cell contractions (Video 4). Instead, Myo-II accumulated uniformly across the apical domain (Fig. S3). Furthermore, the rok sqh-EE double mutants failed to undergo mesoderm invagination (Video 4). Although we cannot rule out the possibility that other Rok substrates are required for Myo-II coalescence, our data suggest that simply activating apical Myo-II is not sufficient for medioapical actomyosin network condensation and radial cell polarity. We favor a model whereby spatial regulation of Myo-II by Rok within the apical domain is required for actomyosin network contraction during a pulse.

**Myo-II phosphomutants exhibit defects in Myo-II assembly/disassembly cycles**

We next determined whether dynamic Myo-II phosphorylation is critical for temporal organization of Myo-II pulses. Similar to Myo-II in wild-type embryos, sqh-TS::GFP displayed Myo-II contraction pulses in sqh germline clones, with the appearance of Myo-II structures lasting 20 to 30 s before being disassembled or remodeled (Fig. 5 A). In contrast, phosphomimetic mutants caused a gradual increase in Myo-II accumulation (Figs. 5 B and S2 C). Myo-II persisted long after it initially appeared, failing to undergo disassembly over the course of constriction (Fig. 5 B and Video 5). Based on the frequency of events involving rapid Myo-II intensity increase, we found that the sqh-AE mutant had significantly fewer Myo-II pulses than wild-type embryos (Fig. 5 D). Correspondingly, apical constriction in the sqh-AE mutant appeared more continuous, with fewer rapid phases of constriction that were lower in magnitude than those observed in wild-type (Fig. 5, B, E, and F; and Fig. S4 A). During pulsatile constriction, phases of rapid Myo-II accumulation correlate with increased constriction rate (Martin et al., 2009). However, the sqh-AE mutant displayed a significantly weaker cross-correlation between Myo-II accumulation and the constriction rate, confirming our observation that mimicking Sqh phosphorylation disrupts contractile pulses and results in more continuous apical constriction (Fig. 5 G).

Mutants that decrease Myo-II phosphorylation resulted in Myo-II dynamics that were distinct from the phosphomimetic alleles. The sqh-A5 mutant displayed contractile oscillations and Myo-II disassembly similar to wild-type embryos (Fig. S2 C), suggesting that dynamic phosphorylation of serine-21 is sufficient for Myo-II remodeling. The sqh-TA mutant exhibited phases of rapid Myo-II accumulation that were temporally correlated with constriction (Fig. 5, C and G), which possibly results from Rok phosphorylating threonine-20. However, the sqh-TA and sqh-AA mutants did not exhibit Myo-II disassembly after accumulation, demonstrating that dynamic phosphorylation of threonine-20 is insufficient for Myo-II remodeling (Fig. 5, C–G; and Figs. S2 C and S4 A). These data are consistent with in vitro experiments that show phosphorylation of either threonine-20 or serine-21 can promote minifilament formation, but that serine-21 is most important for Myo-II ATPase activity (Kamisoyama et al., 1994; Bresnick et al., 1995), suggesting that high levels of Myo-II motor activity are required to remodel the actomyosin network after a pulse. Thus, cycling between high and low Myo-II activity states is required for contractile oscillations.

**Myosin phosphatase localizes to the Myo-II contractile network and is required for contractile pulses**

Continuous Myo-II accumulation in the phosphomimetic mutants suggested that Myo-II dephosphorylation plays a critical role in remodeling Myo-II pulses. Myosin phosphatase is a multisubunit enzyme that includes a catalytic protein phosphatase 1cβ (PP1cβ; also referred to as PP1cβ in *Drosophila*) and a myosin-binding subunit (MBS) that targets PP1cβ to Myo-II (Hartshorne et al., 1998). Previous studies showed that the *Drosophila* homologues of MBS and PP1cβ regulate the phosphorylation state of Myo-II during multiple developmental stages (Tan et al., 2003; Ong et al., 2010; Sun et al., 2011; Majumder et al., 2012). Therefore, we determined whether MBS could catalyze the dynamic remodeling of Myo-II that occurs during pulsing. During cellularization, MBS localizes to Rok/Myo-II–containing furrow canals (Figs. 1 B and S1 F). In addition, MBS localizes to basal Myo-II particles in sqh-TS mutants and to basal aggregates in sqh-AE mutants, providing further evidence that phosphomimetic mutants convert Myo-II into an active conformation that undergoes oligomerization (Fig. S1 F). During ventral furrow formation, MBS is present in the apical cytoplasm of all cells and is enriched in the apical cortex of ventral furrow cells (Fig. 6 A). MBS colocalized with both apical Myo-II and Rok foci throughout the process of ventral furrow formation (Fig. 6, B and C). To determine whether MBS is required for Myo-II pulses, we knocked down MBS using maternal expression of a shRNA that targets MBS (Ni et al., 2011; Fig. S1 G). MBS knockdown resulted in apical Myo-II assembly that exhibited a defect in medioapical coalescence, reminiscent of Myo-II organization in phosphomimetic mutants (Fig. 6, D–F; and Video 6). Although periods of rapid Myo-II accumulation did occur in MBS knockdowns (Fig. 6 G), changes in Myo-II signal were not well correlated with constriction (Fig. 6 H). These data suggest that Myo-II dynamics in MBS knockdown cells is uncoupled from area constriction, indicating that there is a defect in the pulsing behavior compared with wild-type cells. Consistent with this atypical Myo-II behavior, cells in MBS knockdown embryos exhibited more persistent Myo-II accumulation and constriction (Fig. 6 E, F, S4 B, and Video 6). Thus, MBS is recruited to apical Myo-II where it is required to polarize Myo-II accumulation and to generate cycles of assembly and disassembly.
Fig. S5 B). However, phosphomimetic mutations in other Myo-II RLCs have been shown to not fully recapitulate the ATPase activity of phosphorylated Myo-II in vitro (Kamisoyama et al., 1994). Importantly, MBS depletion, which also appears to trap Myo-II in the phosphorylated state, resulted in apical constriction with similar rates to control cells, despite greater heterogeneity in cell size (Fig. 7, D–F). Thus, although contractile pulses may enhance constriction rate, they do not appear to be required for individual cell apical constriction.

Although apical constriction and tissue invagination occur in Sqh phosphomutants, the coordination of tissue invagination was perturbed. In wild-type ventral furrows, Myo-II foci in neighboring cells come together steadily to form a dense supracellular Myo-II meshwork across the entire tissue (Fig. 7 G and Video 7). Furrows in Myo-II phosphomimetic mutants underwent

Figure 5. Cycles of Myo-II assembly and disassembly require dynamic RLC phosphorylation. (A–C) The sqh-AE and sqh-TA mutants inhibit Myo-II remodeling associated with pulses. Time-lapse images are representative cells from sqh+ germ line clone embryos expressing the indicated phosphomutants and Mem::Cherry (membrane). Plots represent apical area and Myo-II intensity as a function of time for the same cells; the time highlighted in gray corresponds to the time-lapse shown. Bars, 5 µm. (D) Quantification of frequency of instances where there is a rapid increase in Myo-II intensity [n = 94 sqh-TS cells, two embryos; n = 131 sqh-AE cells, three embryos; n = 92 sqh-TA cells, two embryos]. (E) Quantification of frequency of instances of rapid apical area reduction [n = 94 sqh-TS cells, two embryos; n = 131 sqh-AE cells, three embryos; n = 92 sqh-TA cells, two embryos]. (F) Quantification of maximal instantaneous constriction rate achieved in control or mutant cells [n = 138 sqh-TS cells, two embryos; n = 187 sqh-AE cells, three embryos; n = 172 sqh-TA cells, two embryos]. (D–F) Error bars are SEM. (G) Mean cross-correlation between constriction rate and change in Myo-II intensity [n = 123 sqh-TS cells, two embryos; n = 143 sqh-AE cells, three embryos; n = 88 sqh-TA cells, two embryos]. P-values were calculated at 0 time offset. **, P < 0.01; n.s., not significant.

Myo-II pulses are important for maintaining tissue integrity during morphogenesis

It is unknown why cells undergo pulsatile, rather than continuous, contraction during tissue morphogenesis. Because we were able to disrupt contractile pulses and promote more persistent apical Myo-II accumulation using both Myo-II phosphomutants and MBS depletion, we next investigated the consequences of continuous apical constriction on tissue invagination. Surprisingly, many of the phosphomutants that disrupted contractile pulses (with the exception of the sqh-AA mutant) underwent apical constriction and tissue invagination (Fig. 7, A–C; Fig. S5, A–C; and Video 2). On average, the sqh-AE mutant cells constricted more slowly than wild-type sqh-TS control cells, suggesting that unregulated Myo-II may decrease the rate of apical constriction and tissue invagination (Fig. 7, A and C; and Fig. S5 B). However, phosphomimetic mutations in other Myo-II RLCs have been shown to not fully recapitulate the ATPase activity of phosphorylated Myo-II in vitro (Kamisoyama et al., 1994). Importantly, MBS depletion, which also appears to trap Myo-II in the phosphorylated state, resulted in apical constriction with similar rates to control cells, despite greater heterogeneity in cell size (Fig. 7, D–F). Thus, although contractile pulses may enhance constriction rate, they do not appear to be required for individual cell apical constriction.

Although apical constriction and tissue invagination occur in Sqh phosphomutants, the coordination of tissue invagination was perturbed. In wild-type ventral furrows, Myo-II foci in neighboring cells come together steadily to form a dense supracellular Myo-II meshwork across the entire tissue (Fig. 7 G and Video 7). Furrows in Myo-II phosphomimetic mutants underwent
Figure 6. Myo-II contractile pulses require MBS. (A) MBS is apically enriched in ventral furrow cells. Images of XY semisagittal and YZ cross sections for fixed sqh::GFP embryos stained for MBS. Bar, 10 µm. (B) MBS structures colocalize with Myo-II foci and the supracellular Myo-II meshwork (arrowheads). Images are from fixed sqh::GFP embryos stained for MBS. Bars, 5 µm. Segmented cell outlines are from subapical sqh::GFP signal. (C) MBS colocalizes with Rok foci (arrowheads). Images are from fixed GFP::Rok embryos stained for MBS. Bars, 5 µm. Segmented cell outlines are from subapical Rok signal. (D and E) MBS knockdown inhibits Myo-II pulses. Time-lapse images are cells from embryos expressing control-shRNA (ctl-shRNA) (D) or MBS-shRNA (E), sqh::GFP (Myo-II), and Memb::Cherry (membrane). (F) Quantification of medioapical polarity in Myo-II organization (n = 115 ctl-shRNA cells, two embryos; n = 123 MBS-shRNA cells, two embryos). Error bars are SEM. (G) Frequency of rapid increases of Myo-II intensity (n = 82 ctl-shRNA cells, two embryos; n = 70 MBS-shRNA cells, three embryos). Error bars are SEM. (H) Mean cross-correlation between constriction rate and change in Myo-II intensity for cells as a function of time offset (n = 118 ctl-shRNA cells, two embryos; n = 84 MBS-shRNA cells, three embryos). P-values were calculated at 0 time offset. **, P < 0.01; n.s, not significant.
apical constriction and the contractile force balance between cells to maintain intercellular cytoskeletal connections in the supracellular actomyosin meshwork that transmits tension across the ventral furrow tissue.

Discussion

Recent studies demonstrated that pulsatile Myo-II contractions drive diverse morphogenetic processes, including Caenorhabditis elegans embryo polarization (Munro et al., 2004), Drosophila gastrulation (Martin et al., 2009; Roh-Johnson et al., 2012; He et al., 2014), dorsal closure (Blanchard et al., 2010; David et al., 2010; Azevedo et al., 2011), germband extension (Rauzi et al., 2010; Fernandez-Gonzalez and Zallen, 2011; Sawyer et al., 2011), oocyte elongation (He et al., 2010), and Xenopus laevis convergent extension (Skoglund et al., 2008; Kim and Davidson, 2011; Shindo and Wallingford, 2014). Although Rok, and likely Myo-II activation via Rok phosphorylation, is required for contraction (Dawes-Hoang et al., 2005; He et al., 2010; Kim and Davidson, 2011; Mason et al., 2013), it was not clear whether Myo-II activation simply regulates cortical Myo-II levels or global contraction, but exhibited abnormal stretching of Myo-II networks and separation of Myo-II foci at late stages of furrow invagination (Fig. 7 H and Video 7; separations observed in 16/21 sqh-AE and 14/14 sqh-EE embryos). These separations involved recoil between Myo-II structures in adjacent cells, resulting in gaps in the supracellular Myo-II meshwork (Fig. 7, H and J). Furthermore, the phenotype of MBS knockdown resembled phosphomimetic Myo-II mutants, also displaying Myo-II separation between cells. Time-lapse images of representative cells from sqh-1 germline clone embryos with the indicated phosphomutants or from embryos expressing the indicated shRNA. Arrowheads indicate contraction of an intact supracellular Myo-II meshwork in control embryos (G and K) or instances of intercellular Myo-II network separation in mutants or knockdowns (H–J). Bars, 10 µm. Box-and-whisker plots display the median (central line), 25th and 75th percentiles (box edges), the most extreme data points not considered outliers (whiskers), and outliers (plotted individually). **, P < 0.01; n.s., not significant.

Figure 7. Defects in Myo-II pulsing disrupt the integrity of the supracellular actomyosin meshwork. (A and B) Mean apical area (A) and normalized mean cellular Myo-II intensity (B) for representative sqh-1 germline clone embryos expressing the indicated phosphomutants (n = 50 sqh-TS cells; n = 49 sqh-TA cells, and n = 50 sqh-AE cells). Shaded area is ± SD. (C) Box-and-whisker plot of cell constriction rates for multiple sqh-1 germline clone embryos expressing the indicated phosphomutants (n = 102 sqh-TS cells, two embryos; n = 169 sqh-AE cells, three embryos; and n = 189 sqh-TA cells, three embryos). (D and E) Mean apical area (D) and normalized mean cellular Myo-II intensity (E) for representative embryos expressing the indicated shRNA (n = 78 control-shRNA cells; n = 31 MBS-shRNA cells). Apical area of cells in MBS-shRNA knockdown embryos were more heterogeneous and larger than that of ctl-shRNA embryos; however, they undergo apical constriction. Shaded area is ± SD. (F) Box-and-whisker plot of constriction rates for multiple embryos expressing the indicated shRNA (n = 104 control-shRNA cells, two embryos; n = 78 MBS-shRNA cells, three embryos). (G–I) Myo-II phosphomutants and MBS knockdown cause separation of Myo-II networks between cells. Time-lapse images of representative cells from sqh-1 germline clone embryos with the indicated phosphomutants or from embryos expressing the indicated shRNA. Arrowheads indicate contraction of an intact supracellular Myo-II meshwork in control embryos (G and K) or instances of intercellular Myo-II network separation in mutants or knockdowns (H–J). Bars, 10 µm. Box-and-whisker plots display the median (central line), 25th and 75th percentiles (box edges), the most extreme data points not considered outliers (whiskers), and outliers (plotted individually). **, P < 0.01; n.s., not significant.
whether coupling between Myo-II activity and its regulators organizes contractile pulses in space and time. Furthermore, why cells undergo pulsatile, rather than continuous, contraction to drive tissue morphogenesis was unknown. We were able to answer these questions by visualizing the consequences of uncoupling Myo-II activation from upstream signaling pathways on cell and tissue dynamics.

**Dynamic Myo-II phosphoryregulation organizes contractile pulses**

We identified dynamic Myo-II phosphorylation as a key mechanism that regulates contractile pulses. We found that Myo-II pulses are associated with dynamic medioapical Rok foci and myosin phosphatase. In addition, the phosphomimetic sqh-AE and sqh-EE mutants, which exhibited constitutive cytoplasmic Myo-II assembly in vivo, exhibited defects in two properties of contractile pulses. First, phosphomimetic mutants did not initially condense apical Myo-II or F-actin into medioapical foci, resulting in Myo-II accumulation across the apical domain and thus a defect in Myo-II radial cell polarity. Second, phosphomimetic mutants continuously accumulated Myo-II in the apical cortex, lacking clear cycles of Myo-II remodeling that are observed in wild-type embryos. Although the phosphomimetic alleles are predicted to partially activate the Myo-II motor’s ATPase activity compared with normal phosphorylation (Kamisoyama et al., 1994), the similarity of the MBS knockdown phenotype suggests that the changes in Myo-II organization and dynamics in phosphomimetic mutants reflect defects in the control over Myo-II dynamics rather than a reduction in motor activity. The consequence of persistent Myo-II assembly across the apical surface in phosphomimetic mutants and MBS knockdown is a more continuous, rather than incremental, apical constriction, demonstrating that pulsatile cell shape change results from temporal and spatial regulation of Myo-II activity via a balance between kinase (Rok) and phosphatase (myosin phosphatase) activity.

Mutants that decrease Myo-II phosphorylation affected contractile pulses in a manner that was distinct from the phosphomimetic alleles. Both the sqh-AA and the sqh-TA mutants exhibited Myo-II assembly into apical foci, potentially mediated by phosphorylation of low levels of endogenous Sqh or phosphorylation of threonine-20, respectively. For the sqh-TA mutant, Myo-II assembly was correlated with constriction, suggesting that Myo-II motor activity is not rate limiting to initiate a contractile pulse. However, Myo-II foci in sqh-TA and sqh-AA mutants were not efficiently remodeled after assembly and coalescence. The persistence of cortical Myo-II foci in sqh-AA and sqh-TA mutants was surprising given that rok mutants and injection of Rok inhibitor reduce cortical localization of Myo-II (Royou et al., 2002, 2004; Dawes-Hoang et al., 2005; Mason et al., 2013). One explanation is that high levels of Myo-II activity induce actomyosin turnover and thus could be required to remodel the actomyosin network after contraction (Murthy and Wadsworth, 2005; Wilson et al., 2010). Alternatively, apical recruitment of myosin phosphatase or proteins that negatively regulate Rok could depend on Myo-II phosphorylation or actomyosin contraction (David et al., 2013). Although future work is needed to address the role of Myo-II motor activity in contractile pulses, the phenotypes of alleles that constitutively reduce phosphorylation further suggest that cycling between high and low phosphorylation states is required for proper Myo-II pulses.

We propose a model for contractile pulses in the ventral furrow where, in combination with unknown cortical cues that apically localize Myo-II, local pulses of apical Rok activity within the medioapical cortex polarize Myo-II assembly and coalescence (Fig. 8 A). Rok foci could polarize actomyosin condensation by generating an intracellular gradient of minifilament assembly and tension that results in inward centripetal actomyosin network flow (Fig. 8 A, red arrows; Bray and White, 1988; Munro et al., 2004; Mayer et al., 2010). In addition, local Myo-II activation by Rok foci combined with broader myosin phosphatase activity throughout the apical cytoplasm could generate a gradient of Myo-II turnover that will concentrate Myo-II into medioapical foci (Fig. 8 A). MBS is required to restrict phosphorylated Myo-II to specific cell–cell interfaces during dorsal closure, demonstrating that the balance between Myo-II kinases and phosphatase can generate spatial patterns of Myo-II activation in epithelial cells (Mizuno et al., 2002). Myo-II remodeling after coalescence could result from local decreases in Rok activity and enrichment of apical myosin phosphatase with Myo-II structures (Fig. 8 A). Thus, coupling Myo-II activation to dynamic signals that regulate Myo-II phosphorylation organizes contractile pulses in space and time to drive incremental apical constriction.

Polarized actomyosin contraction, pulses, and flows generate force and organize the actin cortex in a variety of cellular and developmental contexts (Lecuit et al., 2011; Salbreux et al., 2012). In contrast to the ratchet-like constriction of ventral furrow cells, some cell types undergo extended periods of actomyosin pulsing and area fluctuations without net reduction in area (Solon et al., 2009; Blanchard et al., 2010; He et al., 2010). Furthermore, directional rearrangement of cell contacts, such as during convergent extension in the *Drosophila* germband, can be achieved through planar polarized accumulation of junctional Rok and Myo-II (Bertet et al., 2004; Blankenship et al., 2006; Simões et al., 2010) in conjunction with planar polarized medioapical actomyosin flows (Rauzi et al., 2010; Levayer and Lecuit, 2013). Modulating the spatial and temporal regulation of Myo-II phosphorylation and dephosphorylation provides a possible mechanism to tune contractile dynamics and organization to generate diverse cell shape changes (Mason and Martin, 2011). Consistent with this organizational role, phosphomimetic RLC mutants also disrupt the planar polarized localization of junctional Myo-II in the *Drosophila* germband (Kasza et al., 2014). Thus, it will be important to define the principles that control Myo-II activity and dynamics and how tuning Myo-II dynamics impacts force generation and tissue movement.

**Role of pulsatile Myo-II contractions during tissue morphogenesis**

Myo-II phosphomutants resulted in a more continuous apical Myo-II assembly and apical constriction, enabling us to investigate the role of pulsation during tissue morphogenesis. Continuous Myo-II assembly and contraction in the sqh-AE mutant resulted in a slower mean rate of apical constriction and thus...
dynamics during apical constriction appear to sensitize the tissue to loss of intercellular cytoskeletal integrity during morphogenesis (Fig. 8 B). Although loss of cytoskeletal continuity in phosphomimetic mutants does not block tissue invagination, we speculate that dynamic Myo-II pulses are important to make tissue invagination robust to changes in tensile stress. One possible function of Myo-II pulses is to attenuate tissue tension or stiffness during morphogenetic movements (Fischer et al., 2014). Because pulsated Myo-II contractions are staggered between neighboring cells, pulsation could serve as a mechanism to coordinate contractile force generation across the tissue such that intercellular connections are buffered from high levels of tension (Fig. 8 B).

Indeed, reducing adherens junction proteins sensitizes the intercellular connections between cytoskeletal networks to tensile forces generated in ventral furrow cells (Martin et al., 2010; Sawyer et al., 2011; Spahn et al., 2012). Alternatively, remodeling of actomyosin networks that occurs during pulses could be required to adapt the cytoskeletal organization such that forces transmitted between cells accommodate the changing pattern of tissue-scale forces during the course of morphogenesis. In either case, our data suggest that Myo-II pulsing and remodeling are important for collective cell behavior by ensuring proper force transmission between cells in a tissue undergoing morphogenesis.

Figure 8. Model for mechanism and function of contractile pulses. (A) Myo-II dynamics during a contractile pulse. The contractile pulse is initiated by a local increase in Rok activity that elevates Myo-II phosphorylation and activity. Myo-II coalescence into medioapical foci results from cortical flow (red arrows) resulting from a gradient in cortical tension and net Myo-II minifilament assembly (dark green) at Rok foci with net minifilament disassembly (light blue) in regions of low Rok by myosin phosphatase (purple). Decreased Rok activity after Myo-II coalescence results in Myo-II dephosphorylation followed by remodeling of the contracted cortex. (B) Contractile pulses are required to maintain tissue integrity. Pulsatile Myo-II contraction occurs asynchronously in adjacent cells, which reduces stress at adherens junctions. In addition, Myo-II remodeling allows cells to adjust contacts to maintain stable intercellular cytoskeletal connections. Continuous Myo-II assembly and apical constriction decreases ability of actomyosin networks to dynamically adjust to changes in tissue mechanics, resulting in stretching of Myo-II structures and loss of intercellular connections (red arrows).

delayed tissue invagination. This delay suggested that pulsing might be important for the efficiency of apical constriction. However, phosphomimetic mutants might not fully recapitulate the ATPase activity of phosphorylated Myo-II. The sqh-TA mutant, which also perturbs Myo-II remodeling, constricted ventral furrow cells at a rate that is only slightly slower than wild type. In addition, MBS knockdown, which disrupted Myo-II pulses, exhibited a more variable constriction rate, but with a mean rate comparable to control embryos. Our finding is distinct from studies in other cell types where loss of MBS results in excessive phosphorylated Myo-II accumulation and cell invagination (Lee and Treisman, 2004; Corrigall et al., 2007). Thus, MBS can regulate Myo-II organization and dynamics without causing a significant increase in apical Myo-II levels. We conclude that Myo-II pulses are not absolutely required for individual cell apical constriction.

Although phosphomimetic mutant cells constrict and undergo tissue invagination, the coordination of invagination and the stability of the supracellular actomyosin meshwork were perturbed. Continuous apical constriction was associated with abnormal separation events between Myo-II structures in adjacent cells, resulting in gaps or holes in the supracellular Myo-II meshwork. Thus, continuous Myo-II assembly and a lack of Myo-II dynamics during apical constriction appear to sensitize the tissue to loss of intercellular cytoskeletal integrity during morphogenesis (Fig. 8 B). Although loss of cytoskeletal continuity in phosphomimetic mutants does not block tissue invagination, we speculate that dynamic Myo-II pulses are important to make tissue invagination robust to changes in tensile stress. One possible function of Myo-II pulses is to attenuate tissue tension or stiffness during morphogenetic movements (Fischer et al., 2014). Because pulsed Myo-II contractions are staggered between neighboring cells, pulsation could serve as a mechanism to coordinate contractile force generation across the tissue such that intercellular connections are buffered from high levels of tension (Fig. 8 B). Indeed, reducing adherens junction proteins sensitizes the intercellular connections between cytoskeletal networks to tensile forces generated in ventral furrow cells (Martin et al., 2010; Sawyer et al., 2011; Spahn et al., 2012). Alternatively, remodeling of actomyosin networks that occurs during pulses could be required to adapt the cytoskeletal organization such that forces transmitted between cells accommodate the changing pattern of tissue-scale forces during the course of morphogenesis. In either case, our data suggest that Myo-II pulsing and remodeling are important for collective cell behavior by ensuring proper force transmission between cells in a tissue undergoing morphogenesis.
Materials and methods

Fly stocks and genetics

Stocks used in this investigation are listed in Table S1. Germline clones were generated using the FLP-DHS technique by heat shocking mutant/ovo clones for 2 h at 37°C for 3–4 d (Chou and Perrimon, 1992). For two-channel imaging of Venus::Rok (UAS::Rok; Simões et al., 2010; gift from J. Zallen, Sloan Kettering Institute, New York, NY) and sqh::mCherry (sqh promoter, rok1::FKTPM7; mat15 sqh::mCherry/TM3 females were crossed to ovo/FTY/; hsFlp UAS-Venus::Rok/CyO males, and embryos from heat shocked rok FRT/ovo::FTY; hsFlp UAS-Venus::Rok/+; mat15 sqh::mCherry/+ females were collected for live imaging. For sqh::GFP rescue of sqh1 germline clones, sqh FRT/FM7; sqh::GFP/CyO females were crossed to ovo/FTY/; hsFlp males, the resulting larvae were heat shocked, and sqh FRT/ovo::FTY; sqh::GFP/hsFlp females were crossed to OreR to collect embryos that resulted from germline clones. All sqh::GFP transgenes were expressed via the endogenous sqh promoter. Sqh phosphomutants were recombined with Gap43-mCherry (Memb;::Cherry, plasma membrane marker, driven by sqh promotor; Martin et al., 2010) or with Utr::mCherry (F-actin marker, driven by sqh promotor; Rauzi et al., 2010; gift from T. Lecuit, Institut de Biologie du Développement de Marseille, Marseille, France). For sqh::EE::GFP suppression of rok mutant germline clones, rok1 FRT/FM7; sqh::GFP/CyO females were crossed to ovo/FTY/; hsFlp males and heat shocked. rok1 FRT/ovo::FTY; sqh::EE::GFP/hsFlp females were crossed to OreR males to collect embryos. In all cases, control crosses lacking heat shock were performed to verify the presence of the ovo allele. To generate MBS knockdown embryos, females containing a UAS-driven shRNA against MBS (MBS-shRNA) or the white gene (control) were crossed to males containing a double maternally driver line with both sqh::GFP and Gap43-mCherry. Both UAS stocks were gifts from N. Perrimon, L. Perkins, and the Transgenic RNAi Project (Harvard Medical School, Boston, MA). The resulting larvae were raised at 25°C and females that maternally expressed MBS-shRNA or cfl-shRNA and both fluorescent markers were crossed to OreR males, and the resulting embryos were imaged.

Construction of GFP-tagged sqh phosphomutants

Substitution of threonine-20 and serine-21 with alanine or glutamate prevents or mimicked phosphorylation, respectively (Jordan and Karess, 1997; Substitution of threonine-20 and serine-21 with alanine or glutamate pre- vented or mimicked phosphorylation, respectively (Jordan and Karess, 1997; Jordan and Karess, 1997; Jordan and Karess, 1997; Jordan and Karess, 1997; Jordan and Karess, 1997) (sqh-EE::GFP). For fixed imaging, all embryos were first dechorionated in 80% glycerol and sliced using a syringe blade to create cross sections. PFA-fixed embryos were mounted in AquaPolyMount (Polysciences, Inc.). Images were acquired on a confocal microscope (LSM 710; Carl Zeiss) with a 40×/1.2 Apochromat water objective (Carl Zeiss), using argon ion, 561-nm diode, 594-nm HeNe, and 633-nm HeNe lasers. In Figs. 1 and 6, endogenous GFP was used to visualize Myo-II and Rok, respectively.

Time-lapse imaging

Embryos were dechorionated in 50% bleach and mounted ventral side up on a slide coated with “embryo glue” (double-sided tape soaked in heptane). No. 1.5 coverslips were used as spacers and to create a chamber for the mounted embryo. A No. 1 coverslip completes the top of the chamber, and the chamber was filled with Halocarbon 27 oil. All imaging occurred at room temperature (−23°C) on a confocal microscope with a 40×/1.2 Apochromat water objective, argon ion and 561-nm diode lasers, and a pinhole setting between 1 and 2 Airy units. Simultaneous excitation was used for live two-channel imaging of GFP/mCherry or Venus/mCherry. The band-pass filters for Venus/mCherry were set at 519–578 nm and 599–696 nm, respectively. For Venus/mCherry two-channel imaging, we confirmed that there was minimal spectral bleedthrough between channels. The band-pass selected for GFP was −488–558 nm and for mCherry −573–696 nm. All images were acquired using Zen software (Carl Zeiss).

Image processing and analysis

Images were processed using Fiji (http://fiji.sc/wiki/index.php/Fiji) and MATLAB (MathWorks).

A Gaussian filter (σ = 0.5–0.7) was applied to images. Apical images are maximum intensity projections of −2–5 µm. Subapical image and membrane images are single sections −1–2 µm below the apical projection.

To quantify Rok and Myo-II intensity over time, we generated kymographs of pulses and acquired intensity values along an internal trace through the kymographs. To determine the phase relationship between Rok and Myo-II signals, a Pearson correlation was calculated for various time offsets where the Rok and Myo-II signals were shifted relative to each other.

We used custom MATLAB software, Embryo Development Geometry Explorer (EDGE, Gelbart et al., 2012), to segment images for quantification of apical area and Myo-II intensities. EDGE automatically segmented cell membranes; however, we manually corrected cells with errors in segmentation. Segmented cell membranes were subapical to the Myo-II signal. Embryos were aligned in time using the mean apical area signals of each embryo by choosing the time where the tissue begins to constrict. For quantification of junctional and medioapical Myo-II intensities in Fig. 4, we used a maximum intensity projection of the raw apical Myo-II signal and applied EDGE to segment images. The medioapical domain of a cell is defined by shrinking the cell’s segmented contour by 2 pixels wide; the pixel intensity in this area was used to define medioapical Myo-II, and junctional Myo-II was defined as the difference between the Myo-II intensity in the entire cell and the intensity in the medioapical area [see equations]. We then calculated medioapical polarity using the following equations:

\[
\text{total Myo-II} = \sum \frac{P}{I} - \frac{1}{I} \sum \frac{P}{I}
\]

\[
\text{medial Myo-II} = \frac{\sum \frac{P}{I} - \frac{1}{I} \sum \frac{P}{I}}{P_m}
\]

\[
\text{junctional Myo-II} = \frac{\sum \frac{P}{I} - \frac{1}{I} \sum \frac{P}{I}}{P_m}
\]

\[
p = \frac{\text{medial Myo-II} - \text{junctional Myo-II}}{\text{total Myo-II}}
\]

Generation of full-length Sqh antibody and Western blotting

Full-length Sqh cDNA was cloned and purified from E. coli by GenScript (GenScript USA Inc.). Rabbit sera were obtained using the full-length Sqh as antigen (Panigen, Inc.). Polyclonal anti-Sqh antibody was affinity purified using standard biochemical procedures. In brief, purified Sqh was coupled to CNBr-activated Sepharose 4B (GE Healthcare); the Sqh-coupled resin was incubated with sera, the resin was washed, and the antibody was eluted with glycine, pH 2.5.

Western blotting was performed by grinding embryos directly in sample buffer and running samples on 12% SDS-PAGE gels. Protein was transferred to 0.45-µm nitrocellulose membrane [Bio-Rad Laboratories] and the indicated primary antibodies were then added using horseradish peroxidase-conjugated secondary antibodies [Jackson ImmunoResearch Laboratories, Inc.].

Immunohistochemistry

Antibodies and corresponding concentrations used in this investigation are listed in Table S2. For fixed imaging, all embryos were first dechorionated in 50% bleach and then either methanol/heat fixed or PFA fixed. Embryos stained with a zipper (gift from L. Wieschaus, Princeton University, Princeton, NJ) and α-neurotactin were methanol/heat fixed [Fig. 3]; all other immunostainings used PFA-fixed embryos. For methanol/heat fixations, embryos were placed in boiling Triton salt solution (0.03% Triton X-100 and 0.4% NaCl in water), cooled on ice, and then devitellinized in a 1:1 heptane/methanol solution. PFA-fixed embryos were fixed in a 1:1 solution of 8% PFA in 0.1 M phosphate buffer, pH 7.4, and heptane for 30 min, transferred to a Petri dish, and manually devitellinized using a syringe needle. After immunostaining, heat-fixed embryos were placed on a slide in mounting medium (4% N-propargyl-p-aminobenzoic acid in 80% glycerol) and the chamber was filled with Halocarbon 27 oil. All imaging occurred at room temperature (−23°C) on a confocal microscope with a 40×/1.2 Apochromat water objective, argon ion and 561-nm diode lasers, and a pinhole setting between 1 and 2 Airy units. Simultaneous excitation was used for live two-channel imaging of GFP/mCherry or Venus/mCherry. The band-pass filters for Venus/mCherry were set at 519–578 nm and 599–696 nm, respectively. For Venus/mCherry two-channel imaging, we confirmed that there was minimal spectral bleedthrough between channels. The band-pass selected for GFP was −488–558 nm and for mCherry −573–696 nm. All images were acquired using Zen software (Carl Zeiss).

Downloaded from on June 27, 2017

Myosin phosphorylation regulates contractile pulses • Vasquez et al. 447

Published August 4, 2014
Published August 4, 2014

Submitted: 3 February 2014
Accepted: 26 June 2014

This work was supported by grants R00GM089826 and R01GM105984 to A.C. Martin from the National Institute of General Medical Sciences and by National Institutes of Health predoctoral training grant T32GM007287.

The authors declare no competing financial interests.

References


This online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.201402004/DC1.

We thank E. Wieschaus, J. Zallen, S. Simões, R. Fernandez-Gonzalez, T. Lecuit, Y. Bellaiche, and the Bloomington Stock Center for fly stocks and antibodies used in this study. Additionally, we thank N. Perrimon, L. Perkins, and the Transgenic RNAi Project at Harvard Medical School (National Institutes of Health/National Institutes of General Medical Sciences RO1-GM084947) for providing transgenic RNAi fly stocks used in this study. The neurotactin and E-cadherin antibodies, developed by C. Goodman and T. Uemura, respectively, were obtained from the Developmental Studies Hybridoma Bank, created by the National Institute of Child Health and Human Development of the National Institutes of Health, and maintained at the Department of Biology, University of Iowa, Iowa City, IA 52242. Finally, we thank A. Sokac, I. Cheeseman, J. Zallen, K. Kazaa, and members of the Martin laboratory for their helpful comments and discussion on this manuscript.

This work was supported by grants R00GM089826 and R01GM105984 to A.C. Martin from the National Institute of General Medical Sciences and by National Institutes of Health predoctoral training grant T32GM007287.

The authors declare no competing financial interests.

Online supplemental material

Table S1 lists the genotypes of all of the fly stocks used in this study. Table S2 lists the antibodies and the concentration used for Western blot or immunostaining. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.201402004/DC1.

This work was supported by grants R00GM089826 and R01GM105984 to A.C. Martin from the National Institute of General Medical Sciences and by National Institutes of Health predoctoral training grant T32GM007287.

The authors declare no competing financial interests.

Downloaded from on June 27, 2017

Electronic supplementary material

Fig. S1 shows the characterization of the sqh phosphomutant germline clones before gastrulation and control staining for the MBS knockdown. Fig. S2 shows the characterization of the clones before gastrulation and control staining for the MBS knockdown. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.201402004/DC1.

Table S1 lists the genotypes of all of the fly stocks used in this study. Table S2 lists the antibodies and the concentration used for Western blot or immunostaining. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.201402004/DC1.


Figure S1. Characterization of Myo-II phosphomutants. (A) Schematic of changes in Myo-II localization in the ventral furrow. During a process called cellularization, Myo-II is localized to the basal tips of invaginating plasma membranes, called furrow canals. The invaginating membranes compartmentalize nuclei, forming epithelial cells immediately before gastrulation. At the onset of gastrulation, the Twist transcription factor activates signaling through the small GTPase Rho1 in ventral furrow cells. Rho1 signaling leads to Rok activation, which is thought to phosphorylate and consequently activate Myo-II contractility. (B) Rok intensity increases throughout apical constriction. Mean apical area (magenta) decreases, whereas mean Rok intensity (green) increases (top; n = 67 cells; shaded area is ±SD from the mean [solid line]). Plot of apical area (magenta) and Rok intensity (green) for an individual cell (bottom). (C) GFP-tagged sqh mutants that prevent (mutated to Alanine) or mimic (mutated to Glutamate) phosphorylation are expressed at the same level as endogenous sqh. Western blot of embryo lysates prepared from cellularizing blastoderms probed with α-Sqh, α-GFP, or α-tubulin (loading control). Green arrowhead indicates GFP-tagged Sqh and black arrowhead indicates endogenous Sqh. Germline clones of the sqh1 allele (indicated by asterisk) were estimated to express one-tenth of the endogenous amount of Sqh of wild-type (OreR) embryos. (D) Endogenous sqh expression in sqh1 germline clone embryos is 10% of endogenous sqh expression in OreR embryos. Red arrowhead indicates endogenous Sqh. (E) Representative images of XY semisagittal sections and YZ cross sections for cellularizing live embryos. Bar, 20 µm. (F) Images represent fixed cellularizing embryos expressing sqh::GFP (top) or sqh-AE::GFP (bottom) and stained for MBS. Note that MBS localizes to basal furrow canals and to small or large cytoplasmic aggregates in sqh-TS and sqh-AE, respectively. Bar, 5 µm. (G) Representative images of fixed cellularizing ctl-shRNA (top) and MBS-shRNA (bottom) embryos expressing sqh::GFP and stained for MBS. Note that in ctl-shRNA, MBS localized at the cellularization front and in the cytoplasm; however, in the MBS knockdown there is little detectable MBS signal. Bars, 5 µm.
Figure S2. Ventral furrow phenotypes for sqh phosphomutants. (A) Apical Myo-II localization in sqh mutants expressed with wild-type levels of endogenous Sqh. Images represent live embryos expressing the indicated sqh allele. Bar, 20 µm. (B) Sqh phosphomutants exhibit apical localization specifically in ventral furrow cells. Images represent live embryos expressing the indicated sqh phosphomutant allele expressed in sqh1 germline clones. Note that sqh-AA::GFP localizes to the apical surface of ventral cells, but the tissue fails to form a tissue-wide furrow. Bar, 20 µm. (C) Dynamic phosphorylation of serine-21 is critical for contraction pulses. Time-lapse images show sqh::GFP structures in sqh1 germline clone embryos expressing the indicated allele. Note the shorter time scale for the sqh-AS mutant. Mutants that inactivate or activate serine-21 exhibit continuous Myo-II assembly without apparent disassembly/remodeling. Yellow arrowheads indicate instances of Myo-II assembly. Bar, 5 µm.
Phosphomimetic sqh fails to suppress ventral furrow defects of rok mutants. (A–D) The sqh-EE mutant rescues apical Myo-II localization but not polarized contraction in rok germline clones. Images represent endogenous GFP fluorescence from live rok^1 (A and B) and rok^2 (C and D) germline clone embryos expressing either sqh-TS::GFP or sqh-EE::GFP. Representative images of YZ cross sections (A and C; a, apical; b, basal) or surface views (B and D) are shown. The rok mutants disrupt cortical myosin localization, which is partially suppressed by sqh-EE. However, sqh-EE is present across the entire apical surface in rok mutants.
Figure S4. Phosphomimetic sqh mutants and knockdown of MBS result in more continuous Myo-II accumulation and apical constriction. Examples of individual cell apical constriction in sqh phosphomutants (A) and csl-shRNA and MBS-shRNA mutants (B). Data plots show apical area and Myo-II intensity as a function of time for representative cells from sqh germline clones expressing the indicated Sqh phosphomutants or embryos expressing the indicated shRNAs. Starting time corresponds to 1–2 min before the onset of mean apical constriction in the corresponding embryo.
Figure S5. Tissue invagination and E-cadherin localization in Myo-II phosphomutants. (A–C) Activating and inactivating sqh serine-21 phosphomutants rescue ventral furrow invagination. The sqh-AE allele exhibits delayed apical constriction and tissue invagination. Representative images of XY semisagittal sections and YZ cross sections of live embryos from sqh germline clones expressing sqh-TS::GFP (A), sqh-AE::GFP (B), and sqh-TA::GFP (C). Embryos also express Gap43::mCherry as a membrane marker (magenta). a-b indicates apical–basal polarity. Bars, 10 µm. (D–F) Representative images of furrows of fixed sqh germline clone embryos expressing sqh-TS::GFP (D), sqh-AE::GFP (E), and sqh-TA::GFP (F) and stained for E-cadherin. Bars, 5 µm.
**Video 1.** Rok pulses coincide with Myo-II pulses. Time-lapse images represent maximum intensity Z projections of the apical surface of rok2 germline clones expressing Venus::Rok (wild type) and Sqh::mCherry. Panels are split and merged images from the same movie (Rok, green; Myo-II, magenta). Image stacks were acquired on a laser-scanning confocal microscope (LSM 710; Carl Zeiss) at a time interval of 8.1 s per stack.

**Video 2.** Phenotypes of sqh phosphorylation mutants during gastrulation. Time-lapse images represent ventral furrow formation in sqh1 germline clone embryos expressing the indicated sqh::GFP phosphomutant transgene. Image stacks were acquired on a laser-scanning confocal microscope (LSM 710; Carl Zeiss) at a time interval of ~4.4 s per stack. Images represent maximum intensity Z-projections of sqh::GFP signal.

**Video 3.** Phosphomimetic sqh-AE fails to condense apical F-actin. Time-lapse images represent maximum intensity Z projections of the apical surface of sqh1 germline clone embryos expressing Utr::mCherry (labels F-actin) and the indicated sqh::GFP alleles. Panels are Utr::mCherry/F-actin channel (top) and merged images (bottom; Myo-II, green; F-actin, magenta). Image stacks were acquired on a laser-scanning confocal microscope (LSM 710; Carl Zeiss) at a time interval of 6.7 to 7 s per stack.

**Video 4.** sqh-EE mutant suppresses apical Myo-II localization, but not polarized contraction in rok mutants. Time-lapse images represent maximum intensity Z projections of rok1 germline clone embryos expressing the indicated sqh::GFP allele. Images stacks were acquired on a laser-scanning confocal microscope (LSM 710; Carl Zeiss) at a time interval of ~12.6 s per stack. Note that cells become round and undergo cytokinesis at the end of the movie.

**Video 5.** Contractile pulses are inhibited in sqh phosphorylation mutants. Time-lapse images are from sqh1 germline clone embryos expressing Memb::Cherry and either sqh-TS::GFP (TS), sqh-AE::GFP (AE), or sqh-TA::GFP (TA). Top image for each genotype is sqh::GFP signal and bottom image is a merge of sqh::GFP (green) and Memb::Cherry (magenta) channels. Image stacks were acquired on a laser-scanning confocal microscope (LSM 710; Carl Zeiss) at a time interval of ~6 to 6.7 s per stack. Images represent maximum intensity Z projections of apical sqh::GFP and Memb::Cherry signals.

**Video 6.** Knockdown of MBS inhibits contractile pulses. Time-lapse images are from a MBS-shRNA embryo expressing Memb::Cherry (magenta) and sqh::GFP (green). Images represent a maximum intensity Z projection of apical sqh::GFP and a single apical slice of Gap43::mCherry. Image stacks were acquired on a laser-scanning confocal microscope (LSM 710; Carl Zeiss) at a time interval of 6.8 s per stack.

**Video 7.** Phosphomutant Myo-II and MBS knockdown cause the supracellular Myo-II meshwork to stretch and tear. Time-lapse images represent maximum intensity Z projections of sqh1 germline clones expressing the indicated sqh::GFP transgene and an embryo expressing the MBS-shRNA with sqh::GFP. Fluorescent intensity shown is from GFP signal. Image stacks were acquired on a laser-scanning confocal microscope (LSM 710; Carl Zeiss) at time intervals of 4.5–9.0 s per stack.
Table S1. **Fly stocks**

<table>
<thead>
<tr>
<th>Genotype</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>OreR</td>
<td>2</td>
</tr>
<tr>
<td>sqh^{A0}; P[w+; sqh::GFP]42</td>
<td>2</td>
</tr>
<tr>
<td>ovo^{D3}; hsFLP UAS-Venus::Rok/CyO</td>
<td>4, 5</td>
</tr>
<tr>
<td>rok^{D4}; mat1.5 sqh::mCherry^{411}</td>
<td>1</td>
</tr>
<tr>
<td>y w; P[w+; sqh-TS::GFP]attP1</td>
<td>1</td>
</tr>
<tr>
<td>y w; P[w+; sqh-AS::GFP]attP1</td>
<td>1</td>
</tr>
<tr>
<td>y w; P[w+; sqh-TE::GFP]attP1</td>
<td>1</td>
</tr>
<tr>
<td>y w; P[w+; sqh-TA::GFP]attP1</td>
<td>1</td>
</tr>
<tr>
<td>y w; P[w+; sqh-AE::GFP]attP1</td>
<td>1</td>
</tr>
<tr>
<td>y w; P[w+; sqh=localhost:EE::GFP]attP1</td>
<td>1</td>
</tr>
<tr>
<td>y w; P[w+; sqh-TS::GFP]attP40</td>
<td>1</td>
</tr>
<tr>
<td>y w; P[w+; sqh-TA::GFP]attP1 P{w+; Gap43::mCherry}attP40/CyO</td>
<td>1</td>
</tr>
<tr>
<td>sqh^{A11} FRT101; P{w+; sqh-TS::GFP}attP1</td>
<td>1</td>
</tr>
<tr>
<td>sqh^{A11} FRT101; P{w+; sqh-AS::GFP}attP1</td>
<td>1</td>
</tr>
<tr>
<td>sqh^{A11} FRT101; P{w+; sqh-TE::GFP}attP1</td>
<td>1</td>
</tr>
<tr>
<td>sqh^{A11} FRT101; P{w+; sqh-TA::GFP}attP1</td>
<td>1</td>
</tr>
<tr>
<td>sqh^{A11} FRT101; P{w+; sqh-AE::GFP}attP1</td>
<td>1</td>
</tr>
<tr>
<td>sqh^{A11} FRT101; P{w+; sqh-EE::GFP}attP1</td>
<td>1</td>
</tr>
<tr>
<td>sqh^{A11} FRT101; P{w+; sqh-TS::GFP}attP1 P{w+; Gap43::mCherry}attP40/CyO</td>
<td>1</td>
</tr>
<tr>
<td>Utr::mCherry</td>
<td>3</td>
</tr>
<tr>
<td>sqh^{A10} FRT101; P{w+; sqh-TS::GFP}attP1</td>
<td>1</td>
</tr>
<tr>
<td>sqh^{A10} FRT101; P{w+; sqh-AS::GFP}attP1</td>
<td>1</td>
</tr>
<tr>
<td>sqh^{A10} FRT101; P{w+; sqh-TE::GFP}attP1</td>
<td>1</td>
</tr>
<tr>
<td>sqh^{A10} FRT101; P{w+; sqh-TA::GFP}attP1</td>
<td>1</td>
</tr>
<tr>
<td>rok^{D4}; y^{A8} gal4, hsFLP-39/FM7; P{w+; sqh-TS::GFP}attP1</td>
<td>1</td>
</tr>
<tr>
<td>rok^{D4}; y^{A8} gal4, hsFLP-39/FM7; P{w+; sqh-TA::GFP}attP1</td>
<td>1</td>
</tr>
<tr>
<td>rok^{D4}; y^{A8} gal4, hsFLP-39/FM7; P{w+; sqh-AE::GFP}attP1</td>
<td>1</td>
</tr>
<tr>
<td>P{Ubi-GFP::Rok}</td>
<td>6</td>
</tr>
<tr>
<td>y{[1] sc/+} y{[1]}; P{y^{+}; P{TRiP.GL00094}attP2}ctl-shRNA</td>
<td>7</td>
</tr>
<tr>
<td>y{[1] sc/+} y{[1]}; P{y^{+}; P{TRiP.GL01207}attP2}MBS-shRNA</td>
<td>7</td>
</tr>
<tr>
<td>y{[1] sc/+} y{[1]}; P{w+; sqh::GFP}42, mat67; P{G43::mCherry}, mat15/TM3</td>
<td>1</td>
</tr>
<tr>
<td>y{[1] sc/+} y{[1]}; P{w+; sqh::GFP}42; P{G43::mCherry}, mat15/TM3</td>
<td>1</td>
</tr>
</tbody>
</table>

sqh-XX = sqh promoter and ORF with site-directed mutagenesis at T20 and S21 as noted here. Gap43 = sqh promoter with N-terminal 20 amino acids of rat Gap43 gene, which contains a myristylation sequence. Utr = sqh promoter with Utrophin actin-binding domain. Name and mutations: TS, T20, S21; TA, T20, S21A; AS, T20A, S21; TE, T20, S21E; AE, T20A, S21E; EE, T20E, S21E. Sources: 1, this study; 2, Bloomington Drosophila Stock Center; 3, Rauzi et al. (2010), gift from T. Lecuit; 4, Mason et al. (2013); 5, gifts from J. Zallen, S. Simões [Sloan Kettering Institute, New York, NY], and R. Fernandez-Gonzalez [University of Toronto, Toronto, Canada]; 6, Bardel et al. (2013), gift from Y. Bellaiche [Institut Curie, Paris, France]; 7, gifts from N. Perrimon, L. Perkins, and the Transgenic RNAi Project.

Table S2. **Antibodies used and concentrations**

<table>
<thead>
<tr>
<th>Antibody</th>
<th>Use</th>
<th>Concentration</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rabbit anti-Zipper</td>
<td>HF and PFA</td>
<td>1:500</td>
<td>Wieschaus Laboratory</td>
</tr>
<tr>
<td>Rabbit anti-MBS</td>
<td>PFA</td>
<td>1:500</td>
<td>Tan Laboratory, University of Missouri, Columbia, MO</td>
</tr>
<tr>
<td>Rabbit anti-E-Cad2</td>
<td>PFA</td>
<td>1:50</td>
<td>Developmental Studies Hybridoma Bank</td>
</tr>
<tr>
<td>Mouse anti-neurotactin</td>
<td>HF</td>
<td>1:100</td>
<td>Developmental Studies Hybridoma Bank</td>
</tr>
<tr>
<td>Rabbit anti-Sqh</td>
<td>WB</td>
<td>1:5,000</td>
<td>This study</td>
</tr>
<tr>
<td>Mouse anti-GFP</td>
<td>WB</td>
<td>1:1,000</td>
<td>Roche</td>
</tr>
<tr>
<td>Mouse anti-tubulin</td>
<td>WB</td>
<td>1:5,000</td>
<td>Sigma-Aldrich</td>
</tr>
</tbody>
</table>

HF, heat fixation; WB, Western blot.