Wounded cells drive rapid epidermal repair in the early *Drosophila* embryo

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**ABSTRACT**  Epithelial tissues are protective barriers that display a remarkable ability to repair wounds. Wound repair is often associated with an accumulation of actin and nonmuscle myosin II around the wound, forming a purse string. The role of actomyosin networks in generating mechanical force during wound repair is not well understood. Here we investigate the mechanisms of force generation during wound repair in the epidermis of early and late *Drosophila* embryos. We find that wound closure is faster in early embryos, where, in addition to a purse string around the wound, actomyosin networks at the medial cortex of the wounded cells contribute to rapid wound repair. Laser ablation demonstrates that both medial and purse-string actomyosin networks generate contractile force. Quantitative analysis of protein localization dynamics during wound closure indicates that the rapid contraction of medial actomyosin structures during wound repair in early embryos involves disassembly of the actomyosin network. By contrast, actomyosin purse strings in late embryos contract more slowly in a mechanism that involves network condensation. We propose that the combined action of two force-generating structures—a medial actomyosin network and an actomyosin purse string—contributes to the increased efficiency of wound repair in the early embryo.

**INTRODUCTION**

Epithelial tissues serve as protective barriers that respond rapidly to wounds. It has long been recognized that wound repair is often faster in embryos than in adults (Longaker et al., 1990; Whitby and Ferguson, 1991; McCluskey and Martin, 1995). The basis for this difference is not well understood. Epithelial repair of multicellular wounds in fly, frog, chick, and mouse embryos is accompanied by an accumulation of actin and myosin II around the wound, forming a multicellular cable that has been proposed to act as a purse string that draws the wound closed (Martin and Lewis, 1992; McCluskey and Martin, 1995; Kiehart et al., 2000; Davidson et al., 2002; Wood et al., 2002). Actomyosin purse strings are also present during wound closure by cells in culture (Bement et al., 1993) and in adult epithelia such as the cornea (Danjo and Gipson, 1998) and gut (Russo et al., 2005), where they are important for the repair of incisional wounds and ulcers. The actomyosin purse strings that form around multicellular wounds in the *Xenopus* embryo, however, do not generate mechanical force (Davidson et al., 2002), raising the question of whether they contribute functionally to wound repair. The cell behaviors required to generate the mechanical forces that drive epithelial wound repair remain unclear.

Force-generating actomyosin networks display a range of architectures and contract through distinct mechanisms. For example, apical constriction can be driven by an actomyosin band around the apical circumference of the cell (Burnside, 1971; Hildebrand, 2005) or a medial meshwork of actin and myosin filaments that span the apical cell surface (Martin et al., 2009). Actomyosin networks in some cases actively disassemble their component actin and myosin filaments as they contract (network disassembly). Other networks contract by concentrating a constant number of actin and myosin filaments into a smaller volume (network condensation). Contraction...
of the cytokinetic ring during early cell divisions in Caenorhabditis elegans (Carvalho et al., 2009), medial actomyosin networks in intercalating cells (Fernandez-Gonzalez and Zallen, 2011), and mesoderm precursors in Drosophila (Mason et al., 2013) is associated with network disassembly. By contrast, cytokinetic rings in Dictyostelium (Robinson et al., 2002; Reichl et al., 2008), Schizosaccharomyces pombe (Wu et al., 2003), and Neurospora crassa (Calvert et al., 2011) contract through network condensation. Different mechanisms of contraction may be associated with differences in the magnitude, dynamics, or regulation of the mechanical forces generated during contraction. It is unknown which mechanisms of actomyosin contraction operate during wound repair.

In this study we investigate epidermal wound repair in the Drosophila embryo and find that multicellular wounds heal faster at early stages of embryonic development. Wound repair in early embryos is associated with actomyosin networks at the medial cortex of the wounded cells, whose contraction drives rapid wound closure in parallel with a loss of actomyosin filaments. By contrast, wound repair in late embryos is driven by the assembly of an actomyosin purse string at the wound margin, which contracts more slowly in a mechanism that involves actomyosin condensation. Both structures generate mechanical force, and medial actomyosin networks in early embryos generate greater contractile force than purse strings. We propose that the combined action of these two actomyosin networks contributes to the increased efficiency of wound repair in early embryos.

RESULTS
Quantitative imaging reveals rapid wound closure in early Drosophila embryos
To investigate how wounds heal in developing epithelial tissues undergoing active morphogenetic movements, we compared multicellular wound closure and cytoskeletal dynamics in the epidermis of early Drosophila embryos during axis elongation (stages 7–8) with Drosophila embryos at later stages of epithelial maturation (stages 14–15). We wounded the epidermis using an ultraviolet laser to generate a 7- to 8-μm linear lesion across a single cell interface (Figure 1A and B). We analyzed the dynamics of wound closure using the LiveWire image segmentation algorithm (Fernandez-Gonzalez and Zallen, 2011) to rapidly delineate wound margins in time-lapse movies (Figure 1C).

To compare the rates of wound closure in early and late embryos, we measured the time necessary for wound area to reach 50 μm² (15–25% of the maximum wound size). We found that wound closure was faster on average in early embryos (Figure 1D). Wound area reached 50 μm² in 11.6 ± 2.1 min (mean ± SEM) in early embryos, less than half the time necessary to reach this size in late embryos (25.9 ± 3.1 min, p = 0.0015). There was no significant difference in maximum wound size in early (266 ± 41 μm²) and late (320 ± 42 μm²) embryos, and wounds with a similar maximum size closed faster in early embryos (Figure 1E, F and G), demonstrating that faster wound closure in early embryos is not due to differences in wound size. Wound repair occurred in two phases: an initial fast phase and a subsequent slow phase (Figure 1G). Although both phases were observed in early and late embryos, the fast phase was shorter (Figure 1H) and faster (Figure 1I) in the early embryo, resulting in a more dramatic transition between phases. In early embryos, wounds initially contracted at a rate of 37.8 ± 5.0 μm²/min in the fast phase and a significantly slower rate of 3.1 ± 0.8 μm²/min in the slow phase (p = 8 × 10⁻⁴). The fast phase was significantly slower in late embryos (15.7 ± 3.2 μm²/min, p = 0.002), but the rate of contraction during the slow phase was similar to that in early embryos (3.3 ± 0.8 μm²/min in late embryos).

These results demonstrate that the rate of wound repair decreases as Drosophila embryos develop.

Wounded cells in early embryos assemble medial actomyosin networks
To maintain epithelial continuity, wounded cells must be extruded from the epidermis. Cells can leave an epithelium via extrusion from the apical or the basal surface of the tissue. The direction of extrusion is determined by the position of the contractile machinery (Slattum et al., 2009). To determine how wounded cells exit the epidermis during wound repair in Drosophila, we imaged wound closure in three dimensions (Supplemental Figure S1). Wounded cells in early embryos constricted at their apical surface, rounded up, and shifted their volume basally (Supplemental Figure S1, 7.5 min), disappearing from the apical surface of the epithelium and moving basally at a speed of –0.5 μm/min (Supplemental Figure S1, 15 min). Extruding cells were not detected after wounding in late embryos, possibly due to the rapid phagocytosis of wounded cells by hemocytes (Supplemental Figure S2; Stramer et al., 2005; Wood et al., 2006; Moreira et al., 2010). Wounded cells in the early embryo are therefore extruded basally through apical constriction.

The forces that drive the apical constriction of wounded cells in the early embryo could be provided by the wounded cells themselves or by the surrounding tissue. To distinguish between these possibilities, we investigated myosin localization and dynamics during wound repair in embryos expressing myosin:mCherry and the green fluorescent protein (GFP) fused to E-cadherin. In early and late embryos, cells adjacent to the wound assembled a myosin purse string around the wound (Figure 2, A and B). The myosin purse string formed faster in early embryos, reaching 50% of its maximum intensity at 299 ± 58 s after wounding, compared with 577 ± 77 s in late embryos (p = 0.012, n = 10–13 wounds). The myosin purse string appeared more fragmented in early embryos, failing to form a smooth, continuous boundary around the wound (Figure 2A). Of note, myosin also accumulated at the medial surface of wounded cells during the fast phase of wound closure in early embryos (Figure 2A and Supplemental Movie S1). Medial myosin puncta converged from the periphery of the wounded cell toward the center (Figure 2A and Supplemental Movie S1), suggesting that this pool of protein does not contribute to the purse string. Medial myosin was not observed inside wounds in late embryos (Figure 2B and Supplemental Movie S2). These results suggest that two populations of myosin—a myosin purse string around the wound and medial myosin networks in the wounded cells—contribute to wound closure in early embryos.

To explore the role of cells around the wound in wound closure, we used watershed image segmentation (Fernandez-Gonzalez and Zallen, 2011) to quantify apical area changes in cells adjacent to the wound border (adjacent cells) and cells one to five diameters away (distant cells). Immediately after wounding, the apical area of adjacent cells increased by 7.3 ± 4% and the apical area of distant cells decreased by 3.5 ± 1%, resulting in a net decrease in the area of the epithelium surrounding the wound (p = 0.01; Supplemental Figure S3A). The apical areas of adjacent cells decreased by 20.5 ± 6% of their initial value within 3 min of wounding, and the apical areas of distant cells decreased by 9.6 ± 2% within 5 min of wounding (Supplemental Figure S3A). Of note, wound closure did not start until the apical areas of both adjacent and distant cells had stabilized, indicating that changes in the apical area of cells around the wound do not contribute directly to wound repair (Supplemental Figure S3B).
but medial networks were not observed in the wounded cells, suggesting that the absence of a medial network is not due to increased damage to these cells. Wound healing in early embryos could also involve a distinct, single-cell wound response (Bement et al., 1999; Abreu-Blanco et al., 2011). To test this possibility, we induced wounds within single cells in late embryos, preserving the cell boundary. Single-cell wounds resulted in the assembly of a myosin purse string around the wounded cell, but medial networks did not form in late embryos due to increased laser damage, as cells at this stage have a smaller apical area (Supplemental Figure S4, A–C). To address the possibility that medial myosin networks do not form in late embryos due to increased laser damage, we wounded late embryos using the minimum laser power that elicited a response (Supplemental Movie S3). Reduced-intensity wounds in late embryos still resulted in the assembly of a myosin purse string, but medial networks were not observed in the wounded cells, suggesting that the absence of a medial network is not due to increased damage to these cells. Wound healing in early embryos could also involve a distinct, single-cell wound response (Bement et al., 1999; Abreu-Blanco et al., 2011). To test this possibility, we induced wounds within single cells in late embryos, preserving the cell boundaries. Single-cell wounds resulted in the assembly of a myosin purse string around the wounded cell, but medial networks did not form in late embryos due to increased laser damage, as cells at this stage have a smaller apical area (Supplemental Figure S4, A–C). To address the possibility that medial myosin networks do not form in late embryos due to increased laser damage, we wounded late embryos using the minimum laser power that elicited a response (Supplemental Movie S3). Reduced-intensity wounds in late embryos still resulted in the assembly of a myosin purse string, but medial networks were not observed in the wounded cells, suggesting that the absence of a medial network is not due to increased damage to these cells. Wound healing in early embryos could also involve a distinct, single-cell wound response (Bement et al., 1999; Abreu-Blanco et al., 2011). To test this possibility, we induced wounds within single cells in late embryos, preserving the cell boundaries. Single-cell wounds resulted in the assembly of a myosin purse string around the wounded cell, but medial networks did not form in late embryos due to increased laser damage, as cells at this stage have a smaller apical area (Supplemental Figure S4, A–C). To address the possibility that medial myosin networks do not form in late embryos due to increased laser damage, we wounded late embryos using the minimum laser power that elicited a response (Supplemental Movie S3). Reduced-intensity wounds in late embryos still resulted in the assembly of a myosin purse string, but medial networks were not observed in the wounded cells, suggesting that the absence of a medial network is not due to increased damage to these cells. Wound healing in early embryos could also involve a distinct, single-cell wound response (Bement et al., 1999; Abreu-Blanco et al., 2011). To test this possibility, we induced wounds within single cells in late embryos, preserving the cell boundaries. Single-cell wounds resulted in the assembly of a myosin purse string around the wounded cell, but medial networks did not form in late embryos due to increased laser damage, as cells at this stage have a smaller apical area (Supplemental Figure S4, A–C). To address the possibility that medial myosin networks do not form in late embryos due to increased laser damage, we wounded late embryos using the minimum laser power that elicited a response (Supplemental Movie S3). Reduced-intensity wounds in late embryos still resulted in the assembly of a myosin purse string, but medial networks were not observed in the wounded cells, suggesting that the absence of a medial network is not due to increased damage to these cells. Wound healing in early embryos could also involve a distinct, single-cell wound response (Bement et al., 1999; Abreu-Blanco et al., 2011). To test this possibility, we induced wounds within single cells in late embryos, preserving the cell boundaries. Single-cell wounds resulted in the assembly of a myosin purse string around the wounded cell, but medial networks did not
FIGURE 2: Distinct myosin structures form during epidermal wound closure in early and late Drosophila embryos. (A, B) Epidermal cells expressing E-cadherin:GFP and myosin:mCherry in a stage 7 (A) and a stage 14 (B) embryo. The time after wounding is indicated. Yellow target indicates the site of wounding, dots delineate the wound margin. White arrows track the position of a denticle precursor. Anterior, left; dorsal, up. Bars, 5 μm. (A′, B′) The myosin:mCherry signal in A and B. (C) Correlation coefficient between wound area and mean myosin fluorescence in the wounded cells or the purse string in early (n = 10, red bars) and late (n = 8, blue bars) embryos. Error bars, SEM. (D) Rate of change of wound area, perimeter, or mean myosin fluorescence in the medial region of wounded cells or at the purse string in early (n = 10, red bars) and late (n = 8, blue bars) embryos. The rate of wound area contraction is the average ratio of area(t – 1 min)/area(t). Perimeter reduction was similarly quantified. The rate of increase in mean myosin fluorescence was quantified with the ratio myosin(t)/myosin(t – 1 min) to account for the fact that area and mean fluorescence are inversely correlated. Error bars, SEM.
Actin structures also displayed different dynamics in early and late embryos. We investigated the dynamics of filamentous actin (F-actin) during wound closure using a GFP fusion to the F-actin–binding domain of moesin (Kiehart et al., 2000). Like myosin, F-actin accumulated at the wound margin in early and late embryos (Figure 3, A and B, arrows) and at the apical surface of wounded cells and some cells adjacent to the wound in early embryos (Figure 3A′′). In late embryos, cells around the wound assembled filopodia that extended into the wound (Figure 3, B–B′′, arrowheads; Wood et al., 2002; Abreu-Blanco et al., 2012; Pickering et al., 2013). In early embryos, wounded cells appeared to assemble actin protrusions pointing outward from the wound (Figure 3, A–A′′, arrowheads) and blebs (Supplemental Movie S5). These experiments reveal complex actin dynamics during wound repair and provide further evidence that wounded cells in the early embryo may actively participate in wound closure.

The observation that dynamic actomyosin networks form during wound healing raises the question of whether actomyosin contractility could drive wound repair. In late Drosophila embryos, contractile actomyosin networks accelerate wound repair but are not essential for wounds to close (Wood et al., 2002; Abreu-Blanco et al., 2012). Furthermore, myosin could contribute to wound healing through alternative mechanisms, such as the regulation of actin organization and cross-linking (Burkel et al., 2012; Ma et al., 2012) or actin-based protrusions (Wood et al., 2002; Abreu-Blanco et al., 2012).
Rho-kinase activates myosin directly by phosphorylating the myosin light chain (Amano et al., 1996) and indirectly by inhibiting the myosin phosphatase (Kimura et al., 1996). To determine whether myosin contractile activity is necessary for rapid wound repair in early Drosophila embryos, we injected the Rho-kinase inhibitor Y-27632 into early embryos 1–3 min after wounding to specifically inhibit myosin activity during wound closure. In contrast with water-injected embryos, inhibition of Rho-kinase led to an immediate loss of junctional and medial myosin in epidermal cells (Figure 4, A and B) and a cessation of wound closure (Figure 4C). Small changes in wound area were observed due to relaxation of the cells around the wound (Figure 4B). These results demonstrate that contraction of actomyosin networks is necessary for wound repair in the early embryo and provide further evidence that the mechanisms of wound repair change as embryos develop.

**Wounded cells exert contractile forces that promote wound repair in early embryos**

The dynamic myosin and actin networks in wounded cells are consistent with a model in which medial and purse-string networks actively participate in wound closure in early embryos. Not all actomyosin structures during wound closure, however, are contractile (Davidson et al., 2002). To determine which myosin networks are functionally important for wound closure, we used laser ablation to investigate whether myosin structures generate force during wound healing. We previously showed that cells in the early embryo display uniform viscoelastic properties (Fernandez-Gonzalez et al., 2009). Under these conditions, the initial recoil velocity after laser ablation is proportional to the tension acting on the network (Hutson et al., 2003). To estimate the tension acting on medial myosin networks in wounded cells in early embryos, we ablated these networks ∼2 min after the onset of medial myosin accumulation (Figure 5A). Using particle image velocimetry (Raffel et al., 2007), we observed radial recoil velocities of 160 ± 13 nm/s (Figure 5, A, C, and E). When medial myosin networks were ablated, wounds stopped healing and expanded at an initial rate of 6.1 ± 0.7 μm2/s (Figure 5D). These results demonstrate that the medial myosin networks in wounded cells exert forces on the wound margin and that these forces actively contribute to tissue repair.

To evaluate the contributions of different myosin structures to wound closure, we compared the recoil velocities after ablation of the medial surface of wounded cells or the purse strings around wounds. In early embryos, ablation of medial myosin networks in wounded cells resulted in recoil velocities that were 1.5 times greater than the recoil velocities after ablation of the purse string (p = 0.009; Figure 5, A and E), indicating that wounded cells generate greater local forces. By contrast, ablation of purse strings in late embryos resulted in recoil velocities that were 1.8 times greater than the recoil velocities at the medial surface of the wounded cells (p = 0.070; Figure 5, B and E), consistent with the absence of detectable myosin in wounded cells. Together these results demonstrate that contractile medial actomyosin networks in wounded cells contribute to rapid wound healing in early embryos, and this contribution is greater than that of the purse string around the wound.

**Rapid contraction is associated with disassembly of actomyosin networks**

Our data demonstrate that actomyosin contractility drives rapid wound repair in the early Drosophila embryo. But the mechanisms of actomyosin network contraction during wound repair are unclear. To investigate how actomyosin networks contract during wound repair, we quantified changes in myosin and F-actin fluorescence during
Actomyosin networks that contract by condensation should display mean myosin fluorescence that increases at a similar rate to the rate of wound closure. By contrast, actomyosin networks that contract by disassembly should have a mean myosin intensity that does not change or increases slowly compared with the rate of wound closure. In early embryos, mean myosin fluorescence in wounded cells increased at a rate of $1.15 \pm 0.05$ fold/min during closure, slower than the decrease in wound area ($1.40 \pm 0.08$ fold/min, $p = 0.009$; Figures 2D and 6, A and F). Mean myosin fluorescence in the purse string in early embryos also increased more slowly than the decrease in wound perimeter ($1.08 \pm 0.02$ vs. $1.16 \pm 0.03$ fold/min, respectively, $p = 0.006$; Figure 2D). These data suggest that medial and purse-string actomyosin networks are disassembled during contraction. These differences were observed for both myosin and F-actin throughout wound closure in early embryos (Figure 6, F and G, insets). By contrast, in late embryos mean myosin fluorescence in wounded cells increased at a rate of $1.10 \pm 0.02$ fold/min, similar to the rate of change of wound area ($1.14 \pm 0.02$ fold/min, $p = 0.16$; Figure 2D), and mean myosin fluorescence in the purse string increased at $1.10 \pm 0.02$ fold/min, similar to the rate at which wound perimeter decreased ($1.07 \pm 0.01$ fold/min, $p = 0.10$; Figures 2D and 6, B and J). Wound perimeter and myosin and F-actin fluorescence at the purse string changed at a similar rate during most of wound closure (Figure 6, J and K, insets). These results suggest that medial actomyosin networks in wounded cells in early embryos contract through disassembly, whereas purse-string contraction in late embryos occurs primarily through condensation (Figure 6C). These data indicate that embryonic development brings about changes in the mechanisms of actomyosin network contraction during wound repair.

**FIGURE 5:** Contractile actomyosin networks in wounded cells drive epidermal repair in early *Drosophila* embryos. (A–A′′, B–B′′) Epidermal cells expressing myosin:GFP in a wounded stage 7 (A) or stage 14 (B) embryo. Yellow dots delineate the wound margin, red targets indicate the site of ablation. Myosin networks are shown immediately before ablation (A, B) and 3.3 s after ablation (A′, B′). (A′′, B′′) Overlay of images before (cyan) and after (red) ablation of myosin networks. Anterior, left; dorsal, up. Bars, 5 μm. (C) Particle image velocimetry analysis of the response to ablation in A. Red target indicates the site of ablation, red arrows indicate the direction and magnitude of recoil. Bar, 2 μm. (D) Wound area over time in early embryos. Medial actomyosin networks that form after wounding were ablated at the time indicated by the gray line. (E) Radial recoil velocity after spot ablation of distinct myosin networks during wound repair. Red bars correspond to medial networks in wounded cells (medial, $n = 8$) and purse strings around wounds in early embryos (ps, $n = 6$); blue bars correspond to the medial surface of the wounded cells (medial, $n = 6$) and purse strings around wounds in late embryos (ps, $n = 8$). Error bars, SEM.

with the reduction in wound area (Figure 6, A, D, and E). By contrast, in late embryos, myosin and F-actin fluorescence in the purse string remained relatively constant or decreased slowly during closure (Figure 6, B, H and I). The fold change in total myosin fluorescence during medial network contraction in early embryo was $24.4 \pm 10.4$, significantly greater than the fold change in total myosin fluorescence during purse-string contraction in late embryos, which was $2.6 \pm 0.04$ (Figure 6L; $p = 0.01$). Similar results were obtained for F-actin, which decreased $14.3 \pm 4.4$ fold in medial networks in early embryos and $2.3 \pm 0.4$ fold in purse strings in late embryos (Figure 6L; $p = 0.01$). Medial networks decreased in area and became dimmer as they contracted (Figure 6M), whereas purse strings became thicker and brighter (Figure 6N). Together these results suggest that medial and purse-string networks during wound healing contract through different mechanisms.

**DISCUSSION**

Here we provide evidence that the closure of multicellular wounds in the *Drosophila* embryo is significantly faster at early stages of development and is associated with distinct cellular structures and actomyosin network properties. In late embryos, cells adjacent to the wound form an actomyosin purse string that contracts by network condensation. By contrast, early embryos form a purse string around the wound and medial actomyosin networks in the wounded cells that contract by network disassembly. Laser ablation experiments demonstrate that medial networks and purse strings both generate force. By comparing the magnitude of the forces generated by these structures, we demonstrate that medial actomyosin networks in wounded cells make the dominant contribution to wound closure in early embryos, indicating that wounded cells can actively drive wound repair. This is in contrast with the late embryo,
first demonstration that wounded cells actively participate in wound repair. These results are reminiscent of mechanisms of epithelial closure during development, such as the apical constriction of amnio-serosa cells at the dorsal surface of the Drosophila embryo, which promotes dorsal closure (Toyama et al., 2008; Solon et al., 2009; Blanchard et al., 2010; David et al., 2010). In addition, although actomyosin purse strings are present during wound healing in many systems (Martin and Lewis, 1992; Bement et al., 1993; Brock et al.,

FIGURE 6: Distinct F-actin and myosin dynamics during embryonic wound closure in early and late embryos. (A, B) Wound area over time in early embryos (A) and wound perimeter over time in late embryos (B). T = 0 min is the time of wounding. (C) Model of wound closure in early and late embryos. (D, E) Total myosin (D) and F-actin (E) fluorescence in the medial cortex of wounded cells in early embryos decreased rapidly during wound closure. (F, G) Mean myosin (F) and F-actin (G) fluorescence in the medial cortex of wounded cells in early embryos increased during wound closure. Insets, ratio of the rates of change of wound area to the rate of change of mean myosin (F) or F-actin (G) fluorescence. The black dashed line indicates the point where wound area and mean fluorescence change at the same rate. Error bars, SEM. (H, I) Total myosin (H) and F-actin (I) fluorescence in the purse string in late embryos decreased only slightly during wound closure. Note the different scales in the x- and y-axes with respect to D and E. (J, K) Mean myosin (J) and F-actin (K) fluorescence in the purse string in late embryos increased during wound closure. Insets, ratio of the rates of change of wound perimeter to the rate of change of mean myosin (J) or F-actin (K) fluorescence. The black dashed line indicates the point where wound perimeter and mean fluorescence change at the same rate. Error bars, SEM. (L) Fold change in total myosin (red) and F-actin (blue) levels in medial networks in early embryos or purse strings in late embryos during wound repair. Error bars, SEM. (M, N) myosin:mCherry in a medial network in an early embryo (M) and a purse string in a late embryo (N). Times shown in minutes after wounding. Anterior, left; ventral, down. Bar, 2 μm.
1996; Danjo and Gipson, 1998; Kiehart et al., 2000; Russo et al., 2005), they were shown to generate force only during the repair of subcellular wounds in Xenopus oocytes (Mandato and Bement, 2001), and the mechanisms that drive force generation during multicellular wound healing were largely unknown. Our findings reveal that both purse-string contractility and medial actomyosin networks in wounded cells generate mechanical forces that promote wound closure and show that the apical constriction of wounded cells accelerates tissue repair in the early Drosophila embryo.

Actomyosin contractility is essential for wound repair in early embryos. This is in contrast with late embryos, in which inhibiting myosin activity slows wound repair but does not prevent it (Wood et al., 2002; Abreu-Blanco et al., 2012). In the absence of myosin contractility in the late embryo, wound repair is driven by the zippering of actin-based protrusions that interdigitate at the corners of the wound (Wood et al., 2002; Abreu-Blanco et al., 2012). By contrast, protrusive activity is not sufficient to drive wound repair in the absence of actomyosin contractility in early embryos. The distinct requirements for actomyosin contractility at different stages could be due to differences in the organization and function of the actomyosin machinery, differences in the compensatory mechanisms that contribute to wound closure in the absence of actomyosin contractility, or differences in the extent of actomyosin inhibition in different studies. Of note, wound repair in single-cell Drosophila embryos also requires actomyosin contractility (Abreu-Blanco et al., 2011), suggesting that the requirement for contractile networks is a general feature of wound repair early in development.

Our results indicate that the mechanisms of actomyosin contraction change as embryos develop, with network disassembly playing a dominant role in the early embryo and network condensation driving contraction in the late embryo. These results may reflect changes in the molecular machinery that promotes actomyosin contraction. For instance, the actin depolymerizing and severing factor coflin (Elam et al., 2013) could be more important for wound repair in early embryos, whereas actin cross-linkers such as α-actinin or fimbrin, which stabilize actin filaments (Biron and Moses, 2004; Laporte et al., 2012), may be more important for wound repair in the late embryo. It will be informative to determine the role of actin and myosin regulators in wound repair at different stages of development.

The shape, behavior, and junctional composition of epithelial cells change during development, and these changes may influence the wound repair response. In early embryos, epithelial cells undergo intercalary cell rearrangements that promote axis elongation (Irvine and Wieschaus, 1994), whereas in late embryos cells are relatively static (Simone and DiNardo, 2010; Marcinkevicius and Zallen, 2013). Adherers junctions are more dynamic in early embryos (Huang et al., 2011), which could accelerate junctional disassembly and wound closure. Adherers junctions in early embryos form “spot” complexes (Tepass and Hartenstein, 1994) that could serve as functional anchor points for medial contractile networks (Martin et al., 2010). Spot junctions are rarely observed in the epidermis of late embryos, and this may cause a less efficient engagement of medial networks and adherers junctions at this stage. In addition, cells in the late embryo have septate junctions (Tepass and Hartenstein, 1994), which are associated with proteins such as Discs large and Lethal giant larvae that may negatively regulate myosin activity (Ohshiro et al., 2000; Peng et al., 2000; Simone and DiNardo, 2010). Furthermore, cells in the late embryo contain extracellular matrix proteins that could slow down wound repair (Leptin et al., 1989; Fox et al., 1999). Comparing the role of adhesion molecules in the assembly and contraction of actomyosin networks during wound repair in early and late embryos will determine whether cell–cell and cell–matrix adhesion influence the mechanisms of wound closure.

A striking feature of epidermal wound healing is that its efficiency decays with age. In humans, epidermal wounds in embryos are rapidly repaired with no inflammatory response or scar formation (Rowlatt, 1979). By contrast, wound closure in adults is a slow process that involves inflammation and scarring. We demonstrate that rapid contraction and disassembly of medial myosin networks are associated with faster rates of wound healing than purse-string condensation, and both medial networks and purse strings contribute to rapid wound repair in the early Drosophila embryo. Elucidating the molecular and cellular mechanisms of embryonic wound repair may have implications for the development of therapies to accelerate wound healing in adults through the manipulation of signaling pathways that govern actomyosin localization and dynamics.

**MATERIALS AND METHODS**

**Fly stocks**

We used the following markers for live imaging: *ubi-E-cadherin:GFP* (Oda and Tsukita, 2001), *sqh-sqh:mCherry* (Martin et al., 2009), *sqh-GFP:Moesin* (*spaghetti squash* driven, GFP, moesin-α-helical-coiled, and actin-binding site, or sGMCA; Kiehart et al., 2000), and *sqh-GFP* (Royou et al., 2004). Embryos expressing UASp-GFP:Rho1 (#9392; Bloomington Drosophila Stock Center, Bloomington, IN) were the F2 progeny of UASp-GFP:Rho1 males by mattxtb67;15 females (gift of D. St. Johnston, University of Cambridge).

**Time-lapse imaging**

Embryos were dechorionated in 50% bleach for 2 min and transferred to a drop of 1:1 halocarbin oil 27:700 (Sigma-Aldrich, St. Louis, MO) on a coverslip. Embryos were oriented with their ventral-lateral side down and mounted on an oxygen-permeable membrane (YSI, Yellow Springs, OH). Similar results were obtained when embryos were glued to a coverslip using heptane glue and imaged without additional flattening. GFP was excited with an argon line (488 nm, 100 mW; Melles Griot, Albuquerque, NM). mCherry was excited with a krypton laser (568 nm, 1 mW; Melles Griot). A 63× oil immersion lens (numerical aperture [NA] 1.4; Carl Zeiss, Jena, Germany) mounted on an UltraView RS5 spinning disk system (PerkinElmer, Waltham, MA) equipped with an Orca ER camera (Hamamatsu, Hamamatsu, Japan) was used for imaging. Sixteen-bit, two-color Z-stacks were acquired at 0.5-μm steps every 15 s and projected for analysis (13 slices/stack).

**Embryo injections**

Embryos were dechorionated in 50% bleach for 2 min and aligned on an apple juice–agar pad, transferred to a coverslip coated with heptane glue, and dehydrated for 10–12 min. Embryos were covered in 1:1 halocarbon oil 27:700 (Sigma-Aldrich) and imaged in a Revolution XD spinning disk confocal (Andor, Belfast, United Kingdom) equipped with an iXon Ultra 897 camera (Andor), a Transfer- man NK2 micromanipulator (Eppendorf, Hauppauge, NY), and a PBV820 microinjector (WPI, Sarasota, FL). A 60× oil immersion lens (NA 1.35; Olympus, Tokyo, Japan) was used for imaging. Sixteen-bit, two-color Z-stacks were acquired at 0.3-μm steps every 15 s and projected for analysis (15 slices/stack). Embryos were injected in the perivitelline space with 100–200 pl of 100 mM Y-27632 (Tocris, Ellisville, MO) in water. Control embryos were injected with water. Drugs are predicted to be diluted 50-fold in the embryo (Foe and Alberts, 1983).
Wound delineation and quantification

All the algorithms were developed using Matlab (MathWorks, Natick, MA) and DIPimage (Delft University of Technology, Delft, Netherlands) and were integrated in our custom Scientific Image Segmentation and Analysis (SIESTA) software (Fernandez-Gonzalez and Zallen, 2011). We used the LiveWire algorithm implemented in SIESTA (Fernandez-Gonzalez and Zallen, 2011) for semi-automated tracing of wounds. Briefly, the user moves the mouse pointer around the wound, and the algorithm automatically identifies, in real time, the brightest pixels that follow the trajectory of the mouse (Supplemental Movie S6). The user can approve sections of the contour by clicking with the mouse, which accelerates computational processing by restricting the area of the image under consideration to that bounded by the coordinates of the previous and current positions of the mouse.

Cell areas were quantified using the watershed algorithm in SIESTA (Fernandez-Gonzalez and Zallen, 2011). To quantify protein levels, we divided wounds into two compartments. The pure-string compartment was determined by a three-pixel-wide (0.6 μm) dilation of the wound outline. The medial compartment was obtained by inverting a binary image representing the pure-string compartment and filling the hole in the resulting image using binary propagation. Mean fluorescence was the mean pixel value in each compartment background-subtracted using the image mode as the background and corrected for photobleaching by dividing by the mean pixel value of the entire image. Intensities were normalized to the mean value of each structure in the time-lapse sequence to compare wounds from different embryos. Total fluorescence was the product of mean fluorescence and area or perimeter of the compartment under consideration. Changes in total F-actin and myosin fluorescence were outside the photobleaching range calculated as the change in mean image intensity during image acquisition (1.10 ± 0.02 for F-actin and 1.02 ± 0.01 for myosin).

To distinguish between the fast and slow phases of wound closure, we manually identified the time in which wound closure began. A line was automatically fit to the wound area curve starting from the initial wound closure time. The end of the fast phase was determined as the time in which the correlation coefficient between the wound area curve and the fit line was <0.98. To quantify the rate of wound area contraction in Figure 2D, we took the ratio area(t−1 min)/area(t) and averaged all the values for each wound at all time points from 3 min after ablation until the end of wound closure at 15-s intervals. Perimeter change was similarly measured. To quantify the increase in mean myosin fluorescence, we used the ratio myosin(t)/myosin(t−1 min) to account for the fact that area and mean fluorescence are inversely correlated.

Laser ablation

An N₂ MicroPoint laser (Photonic Instruments, St. Charles, IL) tuned to 365 nm was used to wound cells and cytoskeletal networks as described (Fernandez-Gonzalez et al., 2009). The laser delivers 120-μJ pulses with a duration of 2–6 ns each, producing wounds of ~1 μm in diameter. To wound cells, 10 laser pulses were delivered at three or four spots 2 μm apart along a line that crosses a cell interface parallel to the dorsal–ventral axis of the embryo. This generated a 7- to 8-μm linear damage region. To measure the contractile force generated by medial networks and purse strings, we ablated a single spot in the structure in question. Low-power ablations (Supplemental Movies S4 and S5) were done by a single laser pulse at three spots. These were the minimum conditions that caused ablation of cell interfaces as determined by a visible retraction of the severed ends of the interface and the connected vertices.

The initial velocity of displacement after laser ablation can be used to quantify the amount of tension released (Hutson et al., 2003; Fernandez-Gonzalez et al., 2009). We implemented a custom particle image velocimetry algorithm in SIESTA to quantify the velocity of myosin particles around the wound within the first 4 s after ablation (670-ms laser ablation + 3.3-s image acquisition). Pixels 3–5 μm from the ablation site were considered for velocimetry analysis to account for the material destroyed by ablation and the local effect of force relaxation. The velocity field calculated had a spatial resolution of 0.2 μm × 0.2 μm, and >1250 velocity vectors were measured per experiment. For each experiment, the average magnitude of the radial component of the velocity vectors was considered the initial velocity of displacement induced by ablation.

Statistical analysis

To compare time curves, we used the area under each curve as the test statistic. Sample variances were compared using the F test (Glantz, 2002). Sample means were contrasted using Student’s t test, applying Holm’s correction when three or more groups were considered. The relative correlations and rates of change of wound area or perimeter and medial or purse-string myosin fluorescence were compared using a paired t test. The fold changes of actin and myosin during network contraction were compared between early and late embryos using the Mann–Whitney rank-sum test. We compared the wound size distribution in early and late embryos using the Kolmogorov–Smirnov test. Error bars indicate SEM.

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