Inter-Cellular Forces Orchestrate Contact Inhibition of Locomotion

Graphical Abstract

Highlights

- CIL in embryonic dispersing hemocytes requires choreographed changes in motility
- Tracking actin flow in vivo reveals synchronous changes in colliding actin networks
- An inter-cellular actin-clutch between colliding cells induces lamellar tension
- Tension buildup and release orchestrate the CIL response

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In Brief

Contact inhibition of locomotion in Drosophila macrophages involves mechanical coupling of colliding actin networks. The resultant buildup in lamellar tension orchestrates the response and choreographs cell repulsion.

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**Inter-Cellular Forces Orchestrate Contact Inhibition of Locomotion**

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**SUMMARY**

Contact inhibition of locomotion (CIL) is a multifaceted process that causes many cell types to repel each other upon collision. During development, this seemingly uncoordinated reaction is a critical driver of cellular dispersion within embryonic tissues. Here, we show that *Drosophila* hemocytes require a precisely orchestrated CIL response for their developmental dispersal. Hemocyte collision and subsequent repulsion involves a stereotyped sequence of kinematic stages that are modulated by global changes in cytoskeletal dynamics. Tracking actin retrograde flow within hemocytes in vivo reveals synchronous reorganization of colliding actin networks through engagement of an inter-cellular adhesion. This inter-cellular actin-clutch leads to a subsequent build-up in lamellar tension, triggering the development of a transient stress fiber, which orchestrates cellular repulsion. Our findings reveal that the physical coupling of the flowing actin networks during CIL acts as a mechanotransducer, allowing cells to haptically sense each other and coordinate their behaviors.

**INTRODUCTION**

Contact inhibition of locomotion (CIL), which is a cessation of forward movement upon migratory collision, is a process common to many cell types (Abercrombie and Heaysman, 1953; Astin et al., 2010; Dunn and Paddock, 1982; Gloushankova et al., 1998) that has recently been revealed to behave as a migratory cue for developmentally dispersing populations of cells during embryogenesis (Carmona-Fontaine et al., 2008; Davis et al., 2012; Stramer et al., 2010; Villar-Cerviño et al., 2013). This multifaceted phenomenon requires cells to specifically recognize each other, modulate their migratory capacity, and depending on the cell-type, subsequently repolarize. As a result of this complexity, the mechanisms behind CIL are largely unknown, and it is additionally unclear how these various behaviors during the process are integrated to induce a seamless response.

A range of inter-cellular adhesions and intracellular signaling pathways are postulated to be involved in CIL (e.g., Eph-ephrin [Astin et al., 2010], small GTPases [Carmona-Fontaine et al., 2008], planar cell polarity pathway [Carmona-Fontaine et al., 2008], and cell-cell adhesion [Gloushankova et al., 1998]). However, it is unclear exactly how these various signals feed into the cytoskeletal machinery to control the response. More crucially, nothing is known about the actin dynamics involved in CIL. As a central aspect of CIL is a rapid change in migration, it is clear that to understand the mechanisms behind this phenomenon it will be crucial to elucidate the dynamics of the actin network during the response.

During cell migration, the actin network provides the propulsion that allows a cell to generate movement. The actin cytoskeleton within the lamella of a migrating cell is in a constant state of retrograde flow. Actin polymerizes at the leading edge, which pushes the cell membrane forward. Subsequently, the force of polymerization against the membrane along with Myosin II driven contraction drives retrograde movement of the actin network; it is this treadmill that generates the forces behind cell motility. When a cell moves, cell-matrix receptors, such as integrins, become engaged and bind to the extracellular matrix. Integrin activation leads to a slowing of the actin flow at this integrin-based point of friction, and the force of the moving actin network is then transformed into extracellular traction stresses (Gardel et al., 2008). This integrin-dependent actin-clutch, and the resultant inverse correlation between actin flow and traction force, is hypothesized to be involved in the movement of numerous cell types.

We have been exploiting the embryonic migration of *Drosophila* macrophages (hemocytes) to understand the regulatory mechanisms of CIL and the function of this process during embryogenesis (Davis et al., 2012; Stramer et al., 2010). These cells develop from the head mesoderm and disperse throughout the *Drosophila* embryo taking defined migratory routes. One of these routes occurs just beneath the epithelium along the ventral surface where their superficial location in the embryo allows them to be imaged live at high spatio-temporal resolution approaching what can be achieved from cells in culture. This has revealed that hemocytes spread out to form an evenly
distributed pattern beneath the ventral surface within a thin acellular cavity (the hemocoel) (Stramer et al., 2010). We previously developed a mathematical model of hemocyte dispersal, and computer simulations revealed that this uniform cell spacing may be driven by contact inhibition (Davis et al., 2012). Indeed, a similar analysis of Cajal Retzius cell migration in the cerebral cortex showed an identical requirement for CIL in their dispersion (Villar-Cervero et al., 2013), suggesting that CIL is a conserved mechanism capable of generating tiled cellular arrays.

Here, we show that hemocyte developmental dispersal requires precise contact inhibition dynamics. Quantification of changes in speed and direction during cellular collisions reveals that their CIL response is not stochastic but involves distinct kinematic stages that are synchronized between colliding partners. We also show that this choreographed movement involves a coordinated change in actin dynamics. Tracking actin flow within hemocytes in vivo reveals a physical coupling of the colliding actin networks through engagement of a transient inter-cellular adhesion. It is this “inter-cellular actin-clutch” and the coordinated build-up and release of lamellar tension in colliding cells that orchestrates their behaviors, allowing CIL to behave as an instructive migratory cue.

RESULTS

The Kinematic Steps of the CIL Response Are Synchronized in Colliding Hemocytes

Hemocytes disperse evenly within the ventral hemocoel during Drosophila embryogenesis and their CIL dynamics can be precisely analyzed during this process (Figure 1A; Movie S1) (Davis et al., 2012; Stramer et al., 2010). To elucidate the migratory phases of CIL, we first analyzed the changes in acceleration throughout the response with reference to the time of microtubule alignment between colliding hemocytes (Figure 1B; Movie S1), which we previously revealed is a hallmark of CIL that is associated with a change in hemocyte motility (Stramer et al., 2010). Our data revealed that there is a back acceleration upon microtubule alignment, signifying that cells were slowing down and/or changing direction (Figure 1C) (Davis et al., 2012). This was significant when calculated at either 60- or 20-s intervals (Figures 1C and S1A) highlighting that the time of microtubule alignment is correlated with a sudden change of motion during CIL.

The time of microtubule alignment allowed us to temporally register collisions and extend the time course of the acceleration analysis. We observed that 120 s before microtubule alignment there was a sudden forward acceleration, and 180 s after, an additional back acceleration event (Figure 1C). To determine whether these accelerations were due to changes in cell speed and/or direction, we quantified the internuclear distance of colliding cells during the CIL time course. Two minutes prior to microtubule alignment, the graph of internuclear distance over time revealed a sudden increase in slope (Figure 1D). This suggested that the cell speed increased, which was confirmed by analyzing the nuclear displacement rates (Figure 1E). Immediately upon microtubule alignment, the speed reduced (Figure 1E), which explained the sudden back acceleration (Figure S1A), and ~120 s later the nuclei moved apart (Figure 1D). Analysis of the SD of the internuclear distance over time also highlighted these stages by showing an abrupt decrease in variance as cells progressed from one phase to the next, suggesting that these stages were differentially regulated (Figure S1B). These distinct phases were also visualized by calculating the average velocity vector of left and right colliding cells (Movie S1), which additionally revealed the coordinated behavior of hemocytes during CIL.

As we previously revealed that the lamellae of colliding cells overlap before microtubule alignment (Stramer et al., 2010), we hypothesized that this initial interaction was instigating the kinematic changes. Indeed, lamellae of colliding cells made contact 105 ± 22 s prior to microtubule alignment (Figures 1F and 1G; Movie S1), coinciding with the forward acceleration phase of CIL. Furthermore, an actin fiber developed after lamellae contact that connected the colliding cells, which microtubules subsequently utilized as guides during alignment (Figure 1G). It is important to note that the formation of this actin fiber, and the subsequent repulsion, was not observed when a hemocyte contacted the rear of another cell (i.e., not the lamella), or collided with the lamella of a static cell (Movie S1). This analysis highlights that an interaction between lamellar actin networks of migrating cells is initiating the CIL response.

Analysis of the separation phase of CIL also revealed a synchronous response between colliding cells. Kymography of lamellar retraction revealed that colliding partners simultaneously retracted their lamellae at two to three times the speed of retraction events of freely moving cells (Figures 1H–1J). This retraction event initiated 32 ± 22 s after microtubule alignment, which coincided with the initiation of movement away from the colliding partner. The retraction of lamellae occurred prior to the development of new protrusions away from the colliding partner, suggesting that rapid lamellar retraction initiates cell repolarization (Figures S1C and S1D). Different cell-types exhibit distinct CIL behaviors, which are classified as either type 1 (involving contact-induced lamellar contraction) or type 2 (inhibition of locomotion without contraction) responses (Stramer et al., 2013). Our analysis suggests that hemocytes undergo a classical type 1 response similar in description to chick heart fibroblasts, which were also observed to exhibit sudden lamellar recoil (Abercrombie and Heaysman, 1953).

The Actin Cytoskeleton Rapidly Reorganizes in Colliding Partners during CIL

To understand how the actin networks were mediating the response, we analyzed the actin retrograde flow dynamics during CIL. Time-lapse movies of freely moving hemocytes, labeled with the actin probe, LifeAct-GFP, during their developmental dispersal revealed a highly dynamic actin network within their lamellae (Movie S2). We adapted a fluorescent pseudo-speckle tracking technique (Betz et al., 2009) to quantify the precise speed and direction changes of actin flow within hemocytes in vivo (Figure S2A; Movie S2). This revealed that freely moving hemocytes in vivo have a mean actin flow rate of 3.2 ± 1.8 μm/ min, which is similar to growth cones in vitro (Betz et al., 2009).

Live imaging of collisions revealed significant reorganization of the actin networks during the response (Figure S2A; Movie S2). This reorganization coincided with the development of an actin fiber, which ran perpendicular to the leading edge, linking the
lamellae of colliding cells (Figures 2A and S2B). Pseudo-speckle microscopy of collisions highlighted a slowing of the actin flow within a corridor that colocalized with the actin fiber (Figure 2A; Movie S2) and the aligned microtubule bundle (Figure 2B; Movie S2). Quantification of the actin flow rate within the region surrounding the actin fiber revealed a decrease in magnitude during the response, which suddenly increased upon lamellae separation (Figures 2C, 2D, and S2C; Movie S2). It is interesting to note that these analyses highlight that the increase in actin flow speed occurred in two phases; there was an abrupt spike immediately upon lamellae separation lasting ~20 s (that coincided with the duration of lamella recoil) (Figure 1I), followed by an additional increase during cellular repolarization (Figures 2D and S2C). Analysis of instantaneous changes in flow direction also revealed an increase in rotation after lamellae contact (Figure 2E), which rapidly returned to levels observed in freely moving cells after lamellae separation (Figure 2F; Movie S2). The change in flow direction coincided with a movement of actin fibers within the lamella toward the nascent actin cable, which contributed to its formation (Figure S2D). Upon lamellae...
separation, the actin fiber deformed and was subsequently lost as the actin flow rapidly returned to its normal retrograde direction (Figure 2A; Movie S2). These data highlight that CIL involves a dramatic reorganization of the lamellar actin network.

**Development of a Transient Cell-Cell Adhesion during CIL Coincides with a Coordinated Reorganization of the Actin Network**

As a number of cell types have been reported to form transient inter-cellular adhesions during CIL (Gloushankova et al., 1998; Theveneau et al., 2010), we wanted to determine whether a cell-cell adhesion was responsible for the rapid and seemingly coordinated actin network changes in colliding hemocytes. As Zyxin is known to be a marker of both cell-matrix and cell-cell adhesions (Hirata et al., 2008), we expressed mCherry-Zyxin in hemocytes and colocalized Zyxin-labeled adhesions with actin during CIL. Immediately upon lamellae overlap, a punctum of Zyxin developed at the site of cell-cell contact and persisted for the duration of the response (Figures 3A and 3B; Movie S3). Subsequently, the actin fiber formed immediately behind this concentration of Zyxin (Figures 3A and 3B; Movie S3). We hypothesized that Zyxin foci represented transient cell-cell adhesions that modulate actin retrograde flow in a process analogous to the integrin-based actin-clutch reported in migrating cells in vitro (Gardel et al., 2008). Indeed, visualization of Zyxin while analyzing actin flow revealed that after development of the Zyxin puncta the retrograde flow rate decreased within a corridor immediately behind (Figures 3C and 3D; Movie S3). Furthermore, microtubules polymerized toward this site of adhesion (Figure 3E; Movie S3). Immediately upon microtubule targeting of this adhesion, Zyxin levels decreased (Figure 3F).

The development of an inter-cellular adhesion during CIL suggested that the colliding actin networks were becoming physically coupled. We therefore examined whether this coupling could lead to synchronous changes in actin retrograde...
flow in colliding cells. Investigation of the correlation between instantaneous changes in flow speed in colliding partners revealed an increase immediately upon lamellae overlap, which slowly diminished as cells remained in contact (Figure 3G). Subsequently, at the time of lamellae separation there was an additional abrupt increase in the correlation of instantaneous changes in flow speed (Figure 3H; Movie S3). Similarly, correlation between the instantaneous changes in flow direction of colliding partners showed an increase ~20 s after lamellae contact, which remained high until the time of separation (Figures 3I and 3J). These data reveal that, similar to the orchestrated motion of colliding cells, the actin networks are behaving in a coordinated fashion during CIL.

Increase and Redistribution of Actin Network Stress during Cell Collision

The sudden and synchronous retraction event that occurred upon cell separation (Figure 1H) suggested that tension is developing within the actin network during CIL. We therefore analyzed lamellar tension by laser abscission of the actin cytoskeleton as the recoil rate of the network is indicative of tension within the lamella. Ablating the leading edge or an actin fiber within the lamellae of a freely moving cell led to an initial recoil rate of 28.6 ± 6.8 and 33.2 ± 4.3 μm/min, respectively. Interestingly, the recoil was unidirectional toward the cell body when the ablation was performed within the lamella (Figures 4A and 4B; Movie S4), which may be explained by a bias of myosin contraction toward the rear of the lamella (Svitkina et al., 1997; Yam et al., 2007). In contrast, cutting the region of lamella overlap across the actin fiber linking colliding
cells led to a significantly enhanced retraction rate of 65.8 ± 11.1 µm/min suggesting increased tension was stored within the actin network during CIL (Figures 4A and 4B; Movie S4). We subsequently modeled the amount of force present within the actin network by assuming that the actin cytoskeleton behaves elastically over short time scales (Gardel et al., 2004). Tracking edge displacement of the lamellae upon laser abscission allowed us to measure the changes in strain of the lamellar network as it retracts. This revealed that laser abscission of the lamellae leads to an initial rapid exponential decay that is caused by the sudden release of lamellar tension, followed by a slower linear phase as the retrograde actin flow continuously pulls in the network (Figure 4C). Assuming a lamellar stiffness similar to growth cones, the measured strain upon the release of the tension represents the end of the exponential decay. Assuming mechanical properties similar to previously published lamellae we can estimate the tension. Inset: Sketch illustrating the mechanical model of an elastic and dissipative element. The strain $\epsilon$ is calculated by the ratio $\Delta l / l$. Error bars represent SEM. **p < 0.01.

(D) Hemocyte velocities in freely moving and colliding cells 60 s after laser abscission with respect to the ablation site (red arrow). Magenta arrow is the average direction of the population. Note that after mock ablation there was a significant forward movement of cells, while ablation of the fiber during cell collision led to a significant rearward movement. *p < 0.05.

(E) Localization of actin network stress during cell collision. Top panels: a time-lapse series of a hemocyte containing labeled F-actin undergoing a collision (adapted from Figure 2A). Bottom panels: modeled intracellular actin stresses. Note that stresses were only measured for regions of the lamella that persisted for a 40-s period as deformation history is required in the analysis. Arrows highlight region of lamellae overlap. Dotted line highlights the redistribution of stresses around the cell body and asterisks the regions of high stress that colocalize with the actin fiber.

(F) Kymograph of lamellar stresses over the region that colocalized with the actin fiber. Note the redistribution of stress from the back of the network to the front.

(G) Kymograph of the instantaneous changes in actin flow direction in the region colocalizing with the actin fiber.

(H) Quantification of the mean change in flow direction of the actin network in three regions corresponding to the back, middle, and front of the actin fiber. Note that the changes initially increase in the rear of the network.

See also Movie S4.
lamellar tension allowed us to infer the forces present within the actin network in freely moving and colliding cells. This revealed that forces in colliding lamellae were ~3-fold higher than freely moving cells (Figure 4C). Furthermore, release of this tension by laser abscission during cell collision was sufficient to induce a rearward movement of the cell body away from the ablation site (Figure 4D; Movie S4).

We also adapted previously developed techniques to infer the localization of stresses that drive the retrograde flow using an estimation of actin network deformation (Betz et al., 2011; Ji et al., 2008). As previous correlation of flow changes suggested that the colliding actin networks were becoming coupled (Figures 3G–3J), the model assumed that the colliding networks were behaving as a single visco-elastic material. Interestingly, this analysis revealed that there was a redistribution of stresses during the response. Prior to cell-cell contact, most of the stress was localized around the cell body similar to freely moving hemocytes. However, upon collision the stress redistributed from around the cell body to the base of the actin fiber (Figure 4E; Movie S4) where it subsequently propagated to more distal regions of the fiber (Figure 4F). After cell separation, the stress became localized again to the region around the cell body as in freely moving cells (Movie S4). This stress redistribution indicated that actin cytoskeletal changes were spreading, not from the site of lamellae contact, but from the rear of the network. Indeed, when we examined the distribution of instantaneous changes in the direction of actin flow, we observed a similar propagation from the cell rear (Figures 4G and 4H). These data confirm that lamellar tension increases during CIL and propagates from the rear of the network, suggesting that these cytoskeletal changes are not initiated by a local signal released from the site of cell contact.

**Coupling of Colliding Actin Networks Leads to the Development of a Transient Stress Fiber**

As Myosin II is the major motor responsible for generating contraction within the actin cytoskeleton, we examined its dynamics during hemocyte migration. In freely moving cells, both actin and Myosin II flow in a similar retrograde fashion (Figures 5A, 5B, S3A, and S3B). However, comparable to our previously observed changes in actin flow direction during CIL (Movie S2), Myosin II flow reoriented perpendicularly toward the actin fiber (Figures 5C, 5D, and S3C; Movie S5). The actin fiber developed coincidently with Myosin II accumulation along its length (Figures 5E and 5F) and subsequently became decorated with repeating puncta of Myosin II (Figures S3D). Expression of a GFP-tagged Myosin II specifically in hemocytes within zip1 mutants rescued their developmental dispersal (Figure S4A) and retrograde flow rates (Figures S4B and S4C), showing that Myosin II is critical for actin flow in vivo. These data suggest that myosin II mutant hemocytes have a reduction in the contractility of their actin networks.

We subsequently examined the lamellar dynamics of myosin II mutant hemocytes during collisions. Time-lapse movies of zip1 hemocyte collisions revealed that colliding cells failed to reorganize their actin networks (Figure 6A), or increase their lamellar retraction rates upon separation (Figures 6B and 6C; Movie S6). They also did not develop a prominent actin fiber during cell collision (Figure 6D). These defects were accompanied by a failure to cease their forward motion upon collision and an increased time in contact during the CIL response (Figures 6E and 6F). However, similar to fibroblasts in vitro (Giannone et al., 2007), loss of Myosin II in freely moving hemocytes also reduced their rate of lamellar retraction, showing that they had general defects in lamellar tension (Figure 6C).

We therefore specifically analyzed the role of the stress fiber during CIL by examining the response in diaphanous (dia) mutants. In contrast to zip1 hemocytes, freely moving dia5 mutant cells showed no aberration in actin retrograde flow (Figures SSA–SSC; Movie S6), cell migration (Figures SSD and S5E), or rate of lamellar retraction (Figure 6I). However, colliding cells...
showed a highly variable response (Movie S6) and, on average, failed to generate an actin fiber (Figure 6D) or coordinate their actin dynamics (Figures 6G and S5F–S5I). Additionally, their lamellar retraction rates upon collision were no different from freely moving cells (Figures 6H and 6I). Furthermore, similar to zip^3 hemocytes, dia^5 mutant cells showed a reduced capacity to cease their forward movement upon collision (Figure 6E) and persisted in contact for a longer duration (Figure 6F). Despite these defects, dia^5 hemocytes, similar to wild-type cells, formed a Zyxin puncta and showed some microtubule alignment during collision (Figure S6). These data reveal that preventing stress fiber formation results in an aberrant response whereby cells eventually separate, but in the absence of the sudden lamellar retraction characteristic of type 1 CIL.

A Coordinated CIL Response Is Essential for Hemocyte Dispersal

As dia^5 mutant hemocytes showed uncoordinated cytoskeletal dynamics during collisions (Figures S5G and S5I), we wanted to determine whether this defect also led to an uncoordinated kinematic response during CIL. Analysis of the acceleration time course revealed that, in contrast to the three acceleration changes observed in wild-type hemocytes (Figure 1C), dia^5 mutant cells only showed the back acceleration upon microtubule alignment (Figure 7A). However, this back acceleration was significant only when calculated at 60-s
intervals (Figure S7A) suggesting their response was not as tightly coordinated as controls (Figure S1A). Furthermore, after microtubule alignment the dia5 mutant cells showed no obvious movement away from their colliding partners (Figures 7B and S7B). We also quantified the velocities 240 s after microtubule alignment, which corresponds to the time when both control and dia5 mutant cells have separated (Figure 6F). This revealed that while control cells migrated away from the collision, dia5 mutants showed no significant directional preference with respect to their colliding partners (Figure 7C). These data suggest that dia5 mutant hemocytes fail to actively repel each other.

We next determined how the alteration in cell repulsion in dia5 mutant hemocytes affected their ability to form an evenly spaced pattern during their developmental dispersal. Analysis of the average regions occupied by hemocytes during their dispersal revealed that control cells migrated within defined domains along the ventral surface of the embryo (Figure 7D). In contrast, dia5 mutant hemocytes showed an aberrant domain pattern (Figure 7D) and cells patrolled for greater distances within the embryo (Figures 7E and 7F), suggesting that they were less confined in their motility. dia5 mutant hemocytes also had a slight reduction in cell spacing as shown by their decreased nearest neighbor distances (median = 17.5 μm for controls and 15.8 μm for dia5, p < 0.05) (Figures S7C and S7D; Movie S7). This suggests that precise repulsion dynamics are helping confine the migration of cells to defined regions within the embryo. Previous mathematical modeling of hemocyte dispersal suggested that tightly controlled CIL behavior was
essential for their normal dispersal (Davis et al., 2012). As hemocytes in dia\(^{5}\) mutants showed no directional preference with respect to their colliding partners upon cell separation (Figure 7C), we wanted to determine how this reflected on the overall cell distribution in the simulation. Indeed, randomizing the sensitivity of simulated cells to the direction of their colliding partners (which in control simulations is a fixed parameter) led to a similar acquisition of aberrant domains (Figures S7E and S7F; Movie S7) and a slight reduction in cell spacing (median = 24.2 \(\mu\)m for wild-type parameters and 23.3 \(\mu\)m for randomized repulsion, \(p < 0.001\)). These data show that a precisely orchestrated CIL response in hemocytes is essential for it to behave as an efficient patterning cue.

**DISCUSSION**

Here, we show that hemocyte dispersal requires a precisely orchestrated CIL response; cells are not stochastically repelling each other. Hemocyte collision and subsequent repulsion involves a stereotyped sequence of kinematic stages that are modulated by synchronized changes in actin and microtubule dynamics. These integrated cytoskeletal changes are regulated by a transient inter-cellular adhesion, which physically couples the actin networks of colliding cells and builds up lamellar tension. It is this inter-cellular actin-clutch and the mechanical response to the collision that allows for a precise orchestration of cellular behaviors during CIL.
In recent years, there has been speculation that clutch-like mechanisms also exist at cell-cell junctions (Giannone et al., 2009). Indeed, engagement of cell-cell adhesion molecules in neuronal growth cones leads to similar slowing of actin retrograde flow (Schaefer et al., 2008). Furthermore, apical constriction during gastrulation, which is driven by acto-myoosin contraction, is hypothesized to be induced by a “clutch-like” adhesion (Roh-Johnson et al., 2012). During CIL in hemocytes, this cell-cell adhesion is critical to orchestrate both intracellular responses and inter-cellular behaviors, which allows the response to be coordinated in colliding cells. As apical constriction during gastrulation is also coordinated in space and time across numerous cells within an epithelial sheet (Martin et al., 2010), it is possible that a similar clutch-like mechanism is allowing such inter-cellular integration of forces.

The engagement of a transient inter-cellular adhesion is characteristic of CIL in a number of cell-types (Gloushankova et al., 1998; Heaysman and Pegrum, 1973; Theveneau et al., 2010). This transient adhesion is very similar to the initial punctate adhesion between epithelial cells prior to their formation of a mature cell-cell junction, which also involves a radial actin bundle running perpendicular to the leading edge (Adams et al., 1998; Gloushankova et al., 1998). These transient cadherin-based adhesions, which have been called focal adhesion junctions (Huveneers and de Rooij, 2013), depend on the development of tension (Huveneers et al., 2012). Indeed, cadherins in epithelial cells, astrocytes, and fibroblasts are observed to flow in a retrograde fashion with the actin network (Peglion et al., 2014), which has led to speculation that cadherin-based cell-cell adhesions could lead to an analogous clutch-like mechanism during maturation of adherens junctions (Giannone et al., 2009). It will be interesting to investigate why epithelial cells transform a focal adherens junction into a stable cell adhesion, whereas in fibroblasts and hemocytes the adhesion results in active repulsion.

Subsequent to adhesion engagement during CIL, we observe a sudden and synchronous reorganization of the colliding actin networks. The engagement of this actin-clutch develops tension between colliding cells, which we hypothesize causes the sudden forward acceleration that we observed immediately upon lamellar overlap. It is intriguing to note that chick heart fibroblasts similarly have a momentary acceleration toward each other during their CIL response (Abercrombie and Heaysman, 1953). This intracellular tension may also help form the transient stress fiber that couples the colliding cells, as stress fiber formation is a tension sensitive process (Burridge and Wittchen, 2013). This creates a mechanism of haptic feedback whereby the cells “pull” on each other with the stress fiber acting as a mechanosensor during collisions, similar to its hypothesized role in sensing substrate stiffness (Trichet et al., 2012). The contraction of this actin fiber, which must be embedded within the lamellar actin network, would also explain the network-wide reorganization of actin flows in a process analogous to the “contractile treadmilling” observed toward regions of actomyosin contraction in fibroblasts (Rossier et al., 2010).

As the actin networks reorganize in colliding hemocytes, microtubules polymerize into the region of low retrograde flow (that also correlates with the region of stress fiber development). Microtubules polymerize toward the leading edge in a number of cell-types and undergo frequent catastrophes as they fight against the flowing actin network (Waterman-Storer and Salmon, 1997). In growth cones, upon adhesion engagement, the actin retrograde flow is slowed, allowing microtubules to polymerize toward the contact site (Schaefer et al., 2008) analogous to the initial stages of hemocyte CIL. We therefore speculate that the microtubule alignment observed between colliding cells during CIL is the result of microtubules following a path of least resistance within the actin network. However, it is also possible that microtubules are coupled to the stress fiber through an actin-microtubule crosslinker. Nevertheless, there must be tight coordination of both actin and microtubules during the CIL process.

While it is clear that microtubules are critical for CIL in a number of cell-types (Kadir et al., 2011; Stramer et al., 2010), their exact role during the response is currently unknown. The synchronous back acceleration in colliding partners is strongly correlated with the time when microtubule alignment occurs, suggesting that microtubule bundles are playing a role in stopping forward movement. The microtubules also appear to be critical for the generation of the precise internuclear spacing of the cells during CIL—that is critical for the emergence of the even dispersal pattern (Davis et al., 2012). Part of this spacing is governed by the regular size of hemocyte cell bodies; we hypothesize that the remainder of the nuclear spacing is controlled by the physical and dynamic properties of the microtubules themselves. As microtubules are rigid and capable of bearing compressive loads, the internuclear spacing may be controlled by a combination of tensional elements (i.e., the acto-myoosin network) and elements that resist compression (i.e., microtubules). It may be the sum of these mechanical components—in a process analogous to cellular tensegrity (Ingber, 2003)—that allows the cells to precisely define their separation distance. Another possible role for microtubules may be in regulating the cell-cell adhesion. Microtubules target cell-matrix adhesions to promote their dissolution (Kaverina et al., 1999), and an analogous process may be occurring at the cell-cell adhesion during CIL.

Finally, the retraction phase of CIL involves a seemingly synchronous lamellar response in colliding partners. It is possible that the tension generated by engagement of the actin-clutch is suddenly released due to a breaking of the cell-cell adhesion. Alternatively, the tension may cause cytoskeletal components, such as the microtubule bundle or the actin stress fiber, to undergo a sudden catastrophe. Either way, it is this sudden release of lamellar tension that causes the synchronous retraction of colliding partners, which is essential to generate a choreographed CIL response. Lamellar retraction occurs prior to the generation of new protrusions away from the contact site, and we hypothesize that it is this sudden contraction that drives the subsequent repolarization phase of the response. Indeed, acto-myoosin contractility initiates symmetry breaking and polarization in a number of cell-types (Cramer, 2010; Yam et al., 2007). It is also interesting to note that during initiation of polarized cell motility there is a propagation of actin network changes from the lamellar rear to the front (Yam et al., 2007), which we also observed during CIL initiation, although the mechanics behind this redistribution are currently unclear.
Here, we reveal that hemocyte CIL involves distinct stages, leading to both cells retracting from one another and subsequently repolarizing. It should be made clear that not all contact inhibitory cell-types undergo a similar sequence of events as CIL is not a homogenous response (Stramer et al., 2013). We favor the idea of broadly separating CIL into two types, first envisaged by Abercrombie (1970) and Vesely and Weiss (1973): type 1, which involves active retraction (e.g., hemocyte and fibroblast CIL), and type 2 in which forward movement is randomly deflected or stopped altogether (e.g., dia² mutant hemocytes, epithelial wound closure). While inhibition of cellular protrusions is sometimes thought to be the hallmark of CIL (Mayor and Carmona-Fontaine, 2010), Abercrombie (1970) believed that the predominant response in colliding cells is “a spasm of contraction” that “obliterates the process of ruffling.” It is also interesting to note the initial description of CIL by Abercrombie and Heaysman (1953): “As a result of the adhesion…. they [cells] push or pull against each other to some extent; some of the energy which would normally go into movement is thereby dissipated or becomes potential energy of elastic tension between the cells. When an adhesion breaks, the release of potential energy stored as elastic tension produces the sudden acceleration.” An inter-cellular actin-clutch is an ideal candidate to be responsible for such a response.

**EXPERIMENTAL PROCEDURES**

**Microscopy**

Embryos were mounted as previously described (Davis et al., 2012) and time-lapse images acquired every 5 s (for retrograde flow analysis) or 10 s (for kinetic and co-localization analyses) with a PerkinElmer Ultraview spinning disk microscope during developmental dispersal (stages 14–16). See Extended Experimental Procedures for a list of fly lines and a more detailed description of microscopy.

**Kinematics Analysis and Modeling**

For kinetic analysis, hemocytes were labeled with nuclear and microtubule markers and time-lapse movies were acquired at 10 s/frame. Nuclei were automatically tracked using Volocity software (PerkinElmer). Microtubule markers and time-lapse movies were acquired at 10 s/frame. Nuclei were labeled with either Lifeact-GFP or Moesin-cherry were acquired at 5 s/frame (when imaging actin alone) or at 10 s/frame (when imaging actin with other fluorescent probes). For collision analysis, cells were chosen such that the cells collided once over the duration of the time course. Cells were manually segmented prior to analysis. To quantify retrograde flow rates in lamellae, the cell body was manually segmented and data points within this region discarded. Pseudo-speckle analysis was performed as previously described (Betz et al., 2009). See Extended Experimental Procedures for a more detailed description of retrograde flow analysis.

**Retrograde Flow Analysis**

Time-lapse images of freely moving or colliding hemocytes containing actin labeled with either Lifeact-GFP or Moesin-cherry were acquired at 5 s/frame (when imaging actin alone) or at 10 s/frame (when imaging actin with other fluorescent probes). For collision analysis, cells were chosen such that the cells collided once over the duration of the time course. Cells were manually segmented prior to analysis. To quantify retrograde flow rates in lamellae, the cell body was manually segmented and data points within this region discarded. Pseudo-speckle analysis was performed as described previously (Betz et al., 2009). See Extended Experimental Procedures for a more detailed description of retrograde flow analysis.

**Laser Abscission**

Hemocytes were labeled with UAS-LifeAct-GFP and UAS-RedStinger and imaged on an inverted 780 Zeiss LSM multi-photon confocal with a time interval of 1 s. Cells were imaged for 5–10 s and then ablated with a two-photon laser tuned to 730 nm and focused in a 0.4 μm x 1.5 μm region, either at the edge or within the actin network for freely moving cells or at the region of lamella overlap along the actin fiber for colliding cells. Hemocytes were then imaged for 60 s with only 1.1 s elapsing between frames surrounding the ablation. For mock ablation of colliding cells, the same protocol was performed as mentioned except the laser was switched off. See Extended Experimental Procedures for a more detailed description of modeling lamellar forces.

**Modeling Cytoskeletal Stresses**

To compute the stresses inside the actin cytoskeleton in both freely moving and colliding hemocytes, the actin network was assumed to behave as a linear viscoelastic material. This work used the same model as Betz et al. (2011) to calculate the cytoskeletal forces developed by growth cones in vitro. See Extended Experimental Procedures for a more detailed description of modeling cytoskeletal stresses.

**SUPPLEMENTAL INFORMATION**

Supplemental Information includes Extended Experimental Procedures, seven figures, and seven movies and can be found with this article online at http://dx.doi.org/10.1016/j.cell.2015.02.015.

**AUTHOR CONTRIBUTIONS**

J.R.D., A.L., G.A.D., T.B., M.M., and B.M.S. contributed to the study concept and design, the analysis and interpretation of data, and the writing and revision of the manuscript. F.M., J.T., J.A.S., and Y.M. contributed to the acquisition and interpretation of data and review of the manuscript.

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