**Myosin II Controls Junction Fluctuations to Guide Epithelial Tissue Ordering**

**Highlights**
- Fluctuations in junction length cause neighbor exchange events without morphogenesis
- The variance in junction tension determines how actomyosin influences T1 rates
- Globally increasing junctional actomyosin levels inhibits neighbor exchange
- A developmental increase in isotropic junction tension refines cellular packing

**Graphical Abstract**

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**In Brief**
Curran et al. investigate adherens junction remodeling in the fly notum, where differences in actomyosin levels drive fluctuations in junction length and neighbor exchange, but not morphogenesis. A developmental increase in global junctional tension reduces, rather than drives, neighbor exchange to promote tissue ordering.

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Myosin II Controls Junction Fluctuations to Guide Epithelial Tissue Ordering

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SUMMARY

Under conditions of homeostasis, dynamic changes in the length of individual adherens junctions (AJs) provide epithelia with the fluidity required to maintain tissue integrity in the face of intrinsic and extrinsic forces. While the contribution of AJ remodeling to developmental morphogenesis has been intensively studied, less is known about AJ dynamics in other circumstances. Here, we study AJ dynamics in an epithelium that undergoes a gradual increase in packing order, without concomitant large-scale changes in tissue size or shape. We find that neighbor exchange events are driven by stochastic fluctuations in junction length, regulated in part by junctional actomyosin. In this context, the developmental increase of isotropic junctional actomyosin reduces the rate of neighbor exchange, contributing to tissue order. We propose a model in which the local variance in tension between junctions determines whether actomyosin-based forces will inhibit or drive the topological transitions that either refine or deform a tissue.

INTRODUCTION

Epithelia play an important role as selective barriers that separate animal tissues from the external environment. This depends upon the presence of linear adhesive contacts, called adherens junctions (AJs), which bind neighboring cells to one another (Harris and Tepass, 2010; van Roy and Berx, 2008; Tepass et al., 2001). Since epithelia must tolerate changes in cell packing, even during periods of homeostasis, it is important that the loss and gain of AJs occurs without compromising tissue integrity. This requires that AJs be dynamic structures.

In a monolayer epithelium, the loss and birth of AJs follows a characteristic trajectory as cells change neighbors. First, an adhesive contact connecting two neighboring epithelial cells is lost, leading to the formation of a four-way vertex. This is then resolved by the birth and elongation of a new AJ interface at ~90° to the first. This simple process, often called a T1 transition, connects the two cells in a quartet that were previously separate from one another (Guillot and Lecuit, 2013; Nishimura and Takeichi, 2009; Takeichi, 2014). Such topological transitions provide epithelial monolayers with the fluidity necessary to preserve tissue integrity in the face of the disruptive influence of epithelial cell division (Gibson et al., 2006) and cell delamination (Marinari et al., 2012); while also allowing packing irregularities and defects in the tissue to be resolved (Classen et al., 2005; Farhadifar et al., 2007). Moreover, when accompanied by a redistribution of cell mass, directed neighbor exchange events can be used to drive large-scale morphogenetic movements (Fristrom, 1988; Heisenberg and Bellaiche, 2013).

In many systems, the forces required to drive T1 transitions are generated by the molecular motor, non-muscle Myosin II, as it acts on AJ-associated actin filaments (Desai et al., 2013; Lecuit and Lenne, 2007). Through the action of Myosin II, the sliding of anti-parallel filaments, coupled to the AJ, generates localized mechanical tension that causes AJs to shorten, thereby triggering neighbor exchange. This has been especially well studied in the developing Drosophila germband, where polarized junctional actomyosin (Simoes Sde et al., 2010), actomyosin flows (Bertet et al., 2004; Rauzi et al., 2010), together with the destabilization of E-cadherin at dorsal-ventral AJs (Tamada et al., 2012) drives tissue elongation (Blankenship et al., 2006; Irvine and Wieschaus, 1994; Simoes Sde et al., 2010). Nevertheless, the impact of actomyosin-based forces on individual AJs and on the tissue as a whole critically depends on the precise localization and polarity of the actomyosin network. Thus, while a pulsed polarized actomyosin network drives neighbor exchange (Zallen and Blankenship, 2008; Zallen and Wieschaus, 2004), medial actomyosin pulses tend to induce apical cell constriction, as seen during ventral furrow invagination (Martin et al., 2009; Mason et al., 2013; Vasquez et al., 2014) and dorsal closure (Solon et al., 2009).

Neighbor exchange events have also been suggested to play a much more general role in maintaining the balance between order and disorder in epithelia (Farhadifar et al., 2007; Marinari et al., 2012). However, under conditions of balanced growth or stasis, it is not yet known whether or not actomyosin plays a
direct role in the process of neighbor exchange. To address this question, here we utilize the Drosophila pupal notum to explore the regulation and function of junction dynamics in an epithelium during a period in which it remains relatively stable in size and shape (Bosveld et al., 2012; Guirao et al., 2015). Strikingly, in this context, the impact of Myosin-dependent tension on neighbor exchange is different from that described previously. Instead of driving topological rearrangements, junctional acto-myosin limits the number of neighbor exchanges in this tissue. This is explained, at least in part, by a computational model, which shows how the variance in actomyosin-based tension across cell-cell junctions over time can determine the impact of junctional actomyosin on tissue topology. Thus, as the levels of junctional Myosin II rise across the tissue over the course of development, the rate of neighbor exchange events declines, aiding the gradual refinement of tissue packing as metamorphosis reaches an end.

RESULTS

For this analysis, we began by examining apical junction dynamics in flies expressing endogenous levels of E-cadherin-GFP (Huang et al., 2009) in cells outside of the midline (Figure 1A) (Marinari et al., 2012). To facilitate the analysis of cellular dynamics, images were taken at a high frame rate (30 s intervals), between 12 and 13.5 hr after puparium formation (APF), prior to the onset of cell division, thereby avoiding the impact of cell rounding upon fluctuations in length of apical cell-cell contacts (Bosveld et al., 2012; Guirao et al., 2015) (Movie S1). We observed large numbers of neighbor exchange events throughout this period (Figures 1B–1D, 1F, and 1G and Movie S2), at a rate of 8.5 ± 1.5 × 10^{-4} T1 events per minute per junction (Figure S5J).

One of the initial goals of our analysis was to compare the junction dynamics of this system with those described for germ band extension (GBE) in the fly embryo (Irvine and Wieschaus, 1994), where cell intercalation has been proposed to proceed via relatively discrete steps that occur in sequence to make the process irreversible (Bertet et al., 2004). During GBE, dorsal-ventral oriented junctions are lost, leading to the formation of four-way vertices and rosettes (Blankenship et al., 2006). New junctions are then formed and expand perpendicularly to elongate the embryo along the anterior-posterior axis (Irvine and Wieschaus, 1994). In the notum, by contrast, neighbor exchange was found to be a reversible process (Figure 1B). In many instances, quartets of cells underwent multiple rounds of neighbor exchange within the imaging period (75–180 min) (Figures 1B, 1C, S1A, and S1B). At the same time, many cells reaching a four-way vertex failed to undergo a neighbor exchange event, so that the “lost” junction was later reformed, restoring the original local tissue topology (not shown).

There was no apparent pattern to the timing or position of these neighbor exchange events (Figure 1D). Further, neighbor exchange in the notum was not accompanied by a significant global change in tissue area (Figure 1E) or aspect (width/length) ratio, which remained nearly constant over the 90-min imaging period (Figure 1F). This was also true at the local level. There was no correlation between the length of AJs and their orientation (Figure S1D), and junctions lost or gained during neighbor exchange did not have an orientation bias, as observed in GBE (Figure 1D).

This revealed a pool of Myosin II at apical cell-cell junctions (Figures 3A and 4A). Importantly, this localization is dependent upon the presence of AJs and was lost following RNAi-mediated
silencing of β-catenin or DE-cadherin (Figures 3A and 3B), implying that actomyosin is physically associated with the adherens junction itself.

The overall distribution of this pool of junctional Myosin was not polarized, nor was it associated with junctions with particular orientations (Figure 4B). Similarly, Bazooka was not polarized (Figures S3A and S3B), nor was it preferentially associated with junctions that had low levels of Myosin II (Figure S3C), as has been reported for other tissues (Nakayama et al., 2008; Simoes Sde et al., 2010). In this, both Myosin II and Bazooka reinforce the view suggested above, that this region of the notum lacks a strong axis of junctional polarity. Nevertheless, it is still possible for junctionally associated actomyosin to drive changes in junctional length in this system. To address whether or not this might be case, we compared changes in junctional length (Figure 4D) with the time evolution of Myosin II-GFP intensity at cell-cell junctions, which fluctuates by ~24% around the mean (Figure 5 B, middle). Strikingly, a cross-correlation analysis showed that Myosin II levels are negatively correlated with junction length. Moreover, the accumulation of non-muscle Myosin II...

Figure 1. Neighbor Exchange Events Do Not Contribute to Tissue Morphogenesis

(A) Apical surface projection of a live Drosophila notum labeled with DE-cadherin-GFP at 12 hr APF. Image regions are indicated by dashed boxes. Scale bar, 50 μm.

(B) Neighbor exchange events are reversible. Top: Example of a uni-directional neighbor exchange event. Bottom: A multi-directional neighbor exchange event. Scale bar, 5 μm.

(C) Bar graph showing the fraction of multi-directional transition events for a representative fly. See Figure S1A for further n.

(D) Representative region of the notum, outside the midline, at 12.0 and 13.5 hr APF. Yellow arrows at 12 hr label junctions that are lost, and at 13.5 hr label junctions that have been gained through neighbor exchange events. Scale bar, 10 μm.

(E and F) Line plots showing the area (E) and aspect ratio (F) of the virtual clone of cells shown within the red line in (D), measured over 90 min at 30 s intervals. Crosses in (F) mark the temporal position of neighbor exchange events during this time (each cross represents the time at which a four-way vertex is formed).

(G) Cumulative frequency, for four individuals, of neighbor events over a 90-min period, normalized to the number of junctions within the frame at 12 hr APF.

(H) Diagram of a neighbor exchange event at the center of a four-cell cluster. The center of area of each cell (CoA, marked with a dot) is calculated. The internal aspect ratio is the distance between the CoAs of the cells losing a junction (red) divided by the distance between the CoAs of cells gaining a junction (green). For the external aspect ratio, the axis between the CoAs is extended out and the distance between the perimeter intersections is calculated.

(I) Mean aspect ratio (left, internal; right, external) for the four-cell cluster neighbor exchange event. Error bars represent SEM.

(J) Mean change in cell area for the four-cell cluster involved in neighbor exchange. The cell area was measured from 15 min before to 15 min after the transition and, for each four-cell cluster, the mean was subtracted from the time series. Mean area for four-cell cluster = 222.3 μm². Error bars represent SD n for (I) and (J), 33 exchange events from four flies.
at AJs preceded junction shortening by 30–60 s (Figure 4E). In line with the data suggesting that Myosin II actively shortens junctions, Myosin II levels tended to be higher at shorter junctions, as well as at three- and four-way vertices (Figure 4C).

These data suggest that the junctional pool of Myosin II generates tension that reduces the length of AJs in the notum. It should be noted that we did not observe pulsatile changes in cell areas or medial Myosin intensity, and changes in the intensity of the medial Myosin pool did not precede changes in cell area (Figures S3D–S3G). Moreover, Myosin II was present at the junctions lost as well as the junctions gained following a neighbor exchange event (Figure 4F).

Using Computational Modeling to Determine How Myosin-Dependent Junction Tension Likely Influences Neighbor Exchange

In order to better understand how stochastic fluctuations in junctional Myosin II might influence neighbor exchange events in this system, we developed a stochastic vertex model. In this model, forces acting on vertices arise from line tensions acting on cell interfaces and from a cell area elasticity term constraining the apical cell area to a target area. (Farhadifar et al., 2007; Marinari et al., 2012). The model also assumes that vertices are subjected to a dynamic friction force, such that the equation of motion of the position of a vertex $x_i$ is given by

$$\frac{dx_i}{dt} = f_i$$

(Equation 1)

with $\alpha$ a friction coefficient, and the force $f_i$ dependent in particular on the line tensions along cellular junctions, $\gamma_j$ (Figure 5A). To account for the fluctuations in junction length observed in vivo, we introduced stochastic fluctuations in line tensions into the model (Equation 2). Under this simple assumption, the dynamic evolution of each vertex now depends both on the fluctuating forces to which they are subjected and on an effective friction coefficient that determines how quickly they respond to external forces.

Fluctuations in force in the system were implemented so as to mirror observed changes in Myosin II levels at individual junctions (measured using Spaghetti Squash-GFP; Figure 4A). In
Figure 3. E-Cadherin Couples the Actomyosin Cytoskeleton to the Apical Adherens Junction

(A) Apical surface projection from a live pupa expressing ubi-E-cadherin-GFP (green) and MRLC-mCherry (magenta) for control and UAS-β-catenin (armadillo) RNAi. Scale bar, 25 μm.

(B) Representative nota of control, UAS-Shotgun (DE-cadherin) RNAi and UAS-β-catenin RNAi, driven by pnr-GAL4. Tissues were fixed and stained for E-cadherin (anti-GFP against DE-cadherin-GFP), F-actin (phalloidin), and phospho-Myosin II (S19). Scale bar, 5 μm.

Introducing these terms, we were careful to distinguish between different sources of variation in levels of active Myosin II. As a measure of extrinsic junction-to-junction variation, we characterized the spatial variation in junctional Myosin II density across different junctions in the tissue by averaging the Myosin II intensity for each bond over the course of the 1.5–2 hr of observation. The resulting averages followed a near Gaussian distribution for each bond over the course of the 1.5–2 hr of observation.

We also enforced line tensions to stay positive, and introduced a normal distribution of preferred cell areas with SD σA (Methods S1). We then simulated realizations of the stochastic vertex model described by Equations 1 and 2 (Figure 5E). A topological T1 transition is induced in simulations when, as a result of fluctuations in line tensions, the length of cellular interfaces falls below a defined threshold length.

Using these simulations, we first asked how cell packing is affected by initial conditions. To test this, we ran simulations starting either from a Voronoi tessellation of randomly distributed points in the plane or from a regular honeycomb packing of hexagons (Figure 5F, left). Relaxing both configurations without line tension fluctuations led to different packing states (Figure 5F, right). Relaxing both configurations without line tension fluctuations led to different packing states (Figure 5F, right). Relaxing both configurations without line tension fluctuations led to different packing states (Figure 5F, right). Relaxing both configurations without line tension fluctuations led to different packing states (Figure 5F, right).

The rate of neighbor exchange over time was relatively constant under this model, as observed in the notum (Figure 5G, related to Figure 1G). In order to ensure that the parameters used in this fluctuating vertex model are close to those observed in experiments, we used the measured variations in Myosin II intensity in the tissue to set the ratio of extrinsic and intrinsic fluctuations in line tensions σE/σI and persistence time τp. We then adjusted the magnitude of fluctuations, the area elasticity and the characteristic packaging time, τp = α/Lγ, with α a characteristic cell length. For τp ≈ 4.4 min, line tension to area elasticity ratio γ/K02 = 0.025, preferred area SD σA = 0.192, and σE ≈ 1.07γ, the fluctuating vertex model accounted for the observed rate of T1 transitions and exhibited junction length and cell area fluctuations with a strength close to experimental values (S4A-P, parameters are reported in Table 1 in Methods S1).

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line tension fluctuations were negatively correlated, with a lag time of ~1 min, consistent with experiments (Figure 5H). When comparing the simulated and experimental distributions of edge lengths, we found that the distribution of junctions in the fly notum was skewed toward short junctions (Figure S4Q), while simulated tissues were more ordered than their experimental counterparts (Figure S4R). Overall, this analysis supports the idea that T1 transitions in the tissue are driven, at least in part, by the stochastic changes in the length of cell-cell contacts that result from intrinsic and extrinsic fluctuations in Myosin-dependent tension across AJs.

Having established a model based on the wild-type tissue at 12–13.5 hr APF, we wanted to determine the influence of the mean line tension on tissue dynamics. To do so, we altered the mean line tension $g$, while keeping all other parameters constant. Under these conditions, in which the strength of fluctuations does not vary with Myosin II intensity, we observed a steady decrease in length fluctuations with increasing active Myosin II (Figure 5I and Movie S3). We then used simulations to determine how neighbor exchange frequencies and topological order changed with average line tension. An increasing line tension led both to a decrease in the number of T1 transitions (Figure 5J).
Figure 5. A 2D Vertex Model, with Fluctuating Line Tensions, Recapitulates Experimental Rates of Topological Transitions

(A) Schematic of the vertex model: each vertex $i$ is subjected to a mechanical force $f_i$, that depends on line tensions across cellular interfaces $\gamma_{ij}$.

(B) Left: Histogram for the distribution of time-averaged junctional Myosin II intensities for different junctions in the notum (extrinsic variation), $n = 333$ junctions/3 nota, coefficient of variation (CoV) = 0.13, 60 min = 120 time points. Middle: Distribution of the Myosin II intensity over time (intrinsic variation) for a single junction with CoV = 0.08. The average CoV for Myosin II intensity over time is 0.08. $n = 333$ junctions/3 nota, 60 min = 120 time points. Right: Histogram showing the spatial distribution of Myosin II intensities for a single time frame at 12.5 hr APF (total variation), $n = 333$ junctions/3 nota, CoV = 0.15.

(C) Autocorrelation for Myosin II intensity variation on single junctions $(I(t)I(t+\Delta t))/C_0$, with $I$ the mean intensity of the junction, as a function of lag time $\Delta t$ (blue). Red curve: best fit for an exponential of the form $y = a \exp(-b x)$. The coefficients are: $a = 0.0064 \pm 0.00012$ $b = -0.451 \pm 0.0130$. $n = 315$ junctions/3 nota.

(D) In the stochastic vertex model, line tensions fluctuate over time according to an Ornstein-Uhlenbeck process, with mean tension varying over different junctions. 1–3 are realizations of line tensions over time. The mean of each junction’s fluctuations $\gamma_{ij}$ (dotted lines) is taken from a probability distribution parameterized by the line tension $\gamma$. Preferred cell areas are taken from a normal distribution.

(legend continued on next page)
and to a more ordered tissue, as measured by a larger fraction of cells with six sides (Figure 5K). Thus, the 2D vertex model suggests that under a model in which local Myosin II can drive a neighbor exchange event, an increase in global junctional Myosin II and line tension across the tissue will tend to drive the tissue toward an ordered, hexagonally packed state.

Consequences of Developmental Changes in Myosin II Organization

Since levels of junction tension change with developmental time in this tissue (Bosveld et al., 2012; Marinari et al., 2012), we wanted to determine if the relationship seen in the model between increasing line tension and increasing rate of topological transitions was born out during the course of notum development. To explore this question in detail, we imaged the notum over a much longer period from 20 hr APF. This starting point was chosen to exclude the period in which a global wave of cell division disorders the tissue (14–20 hr APF). Moreover, from 20 hr APF, development in the tissue is accompanied by a gradual shift in Myosin II localization, as apico-medial Myosin is lost and prominent, relatively uniform actomyosin cables are formed around apical cell-cell junctions (Figure 6A). Using the junctional recoil induced by laser ablation as a measure of tension across a junction, we were able to confirm that this visible rise in the level of junctional Myosin II is associated with a significant increase in line tension (Figure 6B and Movie S4); as suggested by previous studies (Guiroa et al., 2015; Marinari et al., 2012).

To test whether this increase in tension is accompanied by changes in tissue order, we examined tissue packing in flies imaged from 20 hr through to 30 hr APF. As seen in the model, the observed increase in the level of junctional Myosin II over time was accompanied by a significant (~15%) increase in the proportion of hexagonally packed cells (Figures 6C and 6D) and by a reduction in the variance of junction lengths across the tissue (Figure 6E). Furthermore, increasing line tensions were associated with an overall reduction in the rate of neighbor exchange from 20 to 30 hr APF (Figures 6F and 6G). Although we observed a temporary rise in the neighbor exchange rate between 24 and 28 hr, when examined closely this appeared to be the result of a change in the relative growth rate of sensory organ precursor cells at this time, which was associated with an increase in the neighbor exchange rate of cells in their vicinity (Figure S5). Thus, overall, while the increase in junctional Myosin II is associated with a decrease in the rate of neighbor exchange events, the T1 transitions that do occur aid the approach to optimal cell packing.

Finally, it was important to test whether the observed increase in the level of junctional non-muscle Myosin II across the tissue is sufficient to explain observed changes in the level of neighbor exchange events that were associated with developmental progression. To do so, building upon previously published work, we perturbed the function of the Myosin activator Rho kinase (Rok) in early development (12 hr APF) (Simoës Sde et al., 2010; Verdier et al., 2006). Manipulating levels of Rok seemed a good choice for this perturbation analysis since Rok is present at AJs and at four-way vertices (Figure 7A), is required for Myosin II phosphorylation and activation (tested using Rok RNAi, Figure 7B), is sufficient to increase levels of junctional Myosin and p-Myosin when expressed as a constitutively active kinase (Rok-CAT expressed under the pnr-GAL4 driver, Figure 7B), and can increase the tension across AJs in the tissue (Figure 7C). Having established this, we were in a position to use Rok-CAT to artificially elevate junctional Myosin II in early development. In line with the model, the increase in junctional Myosin induced by the overexpression of Rok-CAT led to a decrease in the frequency of neighbor exchange events, mirroring the late notum. It also reduced the numbers of reversible T1 events (Figure 7E). Conversely, Rok RNAi led to a corresponding increase in neighbor exchange events (Figure 7D) and to an increase in the chances of a T1 being reversed (Figure 7E). Since the small pool of medial Myosin is also lost following Rok RNAi (Figure 7B), one can rule out an essential role for medial Myosin II in driving neighbor exchanges. Moreover, treatments that used perturbations in Moesin function to change the relevant levels of junctional and medial Myosin suggest that it is the junctional pool of Myosin that functions to damp neighbor exchange (Figure S6). Importantly, these results were confirmed using phospho-null (sqh-AA) and phospho-active (sqh-EE) forms of Myosin regulatory light chain to decrease or increase Myosin II activity, respectively (Figures S6A and S6G).

Previously, we proposed a role for neighbor exchange in midline cell delamination, an event that begins at ~14 hr APF.
This led us to explore how perturbations in the levels of junctional Myosin affect delamination in the tissue. In line with the data above, increased Myosin levels, induced by Rok-CAT expression, were found to block cell delamination in the midline. Conversely, Rok RNAi increased rates of midline delamination and caused cells to delaminate outside of the midline, something never observed in the wild-type (Figure S7). These observations confirm a role for junctional Myosin in the inhibition of neighbor exchange and suggest an important role for neighbor exchange in crowding-induced epithelial cell delamination (Marinari et al., 2012).

In sum, these data suggest that increasing the average levels of Myosin does not promote neighbor exchange in the notum, as might have been expected based upon work observed in other tissues and systems. Instead, in this relatively isotropic tissue, increasing levels of junctionally associated Myosin II actively limits neighbor exchange.

**DISCUSSION**

In the context of a planar polarized tissue, the localization of Myosin II along cell-cell contacts, with a specific orientation,
can generate contractile forces that contribute to tissue remodelling on a macroscopic scale, as individual cell-cell contacts are lost and new contacts form. This has been studied in detail by many groups in the context of the developing embryo. How though does Myosin-dependent tension contribute to neighbor exchange in epithelia at steady state—like the adult gut or skin?

In this paper, we explore this question by investigating the role for Myosin II in a stable epithelium, the fly pupal notum, during a period of developmental time in which there are no cell divisions, no cell delamination, and no overt changes in tissue shape or size. Strikingly, in this tissue, Myosin II limits fluctuations in AJ length and, as a consequence, neighbor exchange. Therefore, with developmental time, as levels of junctional Myosin II increase in a relatively isotropic manner, there is a corresponding decrease in the rates of neighbor exchange and an increase in tissue order, as seen by a decrease in junction length variance and by an increased proportion of hexagonal cells, both measures of improved cell packing. In this way, changes in Myosin II activity levels and localization contribute to the refinement of the tissue observed at the end of development.

Although these observations might appear to conflict with studies of neighbor exchange in other tissues where Myosin II has been shown to drive neighbor exchange, the function of the molecules involved seems to be identical. Thus, in the notum, DE-cadherin and β-catenin couple Myosin II to cell-cell interfaces enabling Myosin to influence tissue packing. In addition, increases in the level of Myosin II at junctions are associated with increased junction tension. As a result of this, junctions with high levels of Myosin II tend to be shorter than those with low Myosin II levels. Moreover, a cross-correlation analysis shows that changes in Myosin II levels precede, and therefore likely drive, changes in junction length, as one would...
expect if the recruitment of Myosin II to a junction led to its contraction.

Why then is the impact of Myosin action at the level of the tissue so different in different systems? Our model suggests that the answer may lie in the spatial organization and temporal control of Myosin II. The ability of junctional Myosin II to drive the loss of a specific junction will depend on the tension acting at neighboring cell-cell junctions that resist its contraction. Thus, neighbor exchange will be favored in tissues where there is a high variance in Myosin II levels between neighboring junctions (see Figure 7A), as exemplified by the early fly embryo, where planar polarization in the developing germband generates extreme differences in the levels of Myosin II at perpendicular junctions (Pare et al., 2014; Simoes Sde et al., 2010), driving efficient and directed neighbor exchange. Conversely, in a tissue like the notum, where the distribution of Myosin II is relatively isotropic but fluctuates in time and space (see Figure 7C), the impact of Myosin on the rate of topological transitions will depend on the balance between the average force generated on cell edges and the spatial and temporal fluctuations in these forces. Thus, the impact of an increase in the average levels of Myosin across a tissue on tissue order will depend on the change in variance. If, as observed in the notum, the increase in junctional Myosin and line tension is accompanied by a visible decrease in the spatial variation of Myosin II (Figure 6A), neighbor exchange rates will slow and the tissue will tend to become more ordered with time. Our analysis therefore suggests that epithelia can finely tune their behavior by controlling the average levels of junctional Myosin II and the temporal and spatial variation in its localization at individual junctions.

We note here that while the impact of Myosin II on junctional length fluctuations can explain the observed changes in tissue organization, Myosin II is also likely to play additional roles in the process of neighbor exchange in the tissue. This is suggested by the observed accumulations of Myosin II and Rok at tri-cellular junctions (Figures 7A and 7B). Thus, in limiting neighbor exchange, Myosin II may also limit the ability of fluctuations in junction length to induce smooth passage through a four-way vertex. This may help explain the reduction in junctional reversals seen with increasing levels of Myosin II (Figure 7E), and may explain why a subset of junctions pause at four-way vertices (not shown).

Finally, this study shows how local fluctuations in the activity, localization, and levels of a molecule, in this case Myosin II, can drive local changes in cell shape, to produce larger changes in tissue order. In this way, our analysis of length fluctuations bridges the molecular, cellular, and tissue scales. While this type of analysis remains in its infancy, it is likely to be important for coming to a mechanistic understanding of a wide range of biological processes. Moreover, our analysis shows how the emergence of tissue order can be driven by apparently stochastic fluctuations (Cohen et al., 2011) that are the inevitable consequence of the action of small numbers of molecules, rather than by a directed developmental program. While it may not be possible to use stochastic processes to aid rapid morphogenetic events, like those that accompany early embryonic development, organizing a tissue in this way has its advantages. The use of this type of system can help ensure that an epithelium is robust to both intrinsic (e.g., cell division and delamination) and extrinsic (e.g., forced deformation) perturbations. Thus, fluctuation-induced changes in cell packing, of the type we see here, would seem to be a good way of maintaining integrity in a dynamic, living epithelium. We therefore expect to see similar processes throughout the animal world.

STAR METHODS

Detailed methods are provided in the online version of this paper and include the following:

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SUPPLEMENTAL INFORMATION

Supplemental Information includes seven figures, one table, supplemental text, and four movies and can be found with this article online at https://doi.org/10.1016/j.devcel.2017.09.018.

AUTHOR CONTRIBUTIONS

S.C., C.S., and B.B. initially conceived the project. S.C., G.S., and B.B. wrote the manuscript. S.C. designed, performed, and analyzed fly experiments under the guidance of B.B. C.S. designed the data analysis pipeline and wrote the analysis software package, under the guidance of A.K. and B.B., performed data analysis (along with S.C.), and helped J.B. with simulation analysis. J.B. and M.d.G. developed, designed, and ran simulations of the 2D vertex model, under the guidance of G.S. and B.B., and G.S., and B.B. jointly developed the ideas proposed in this manuscript.

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EPSRC - CoMPLEX PhD program UCL (C.S.); Cancer Research UK (C1529/A9786; C1529/A17343 to B.B.); the MRC-LMB, University College London (MC_U12266B to S.C., C.S., and B.B.); the Francis Crick Institute, which receives its core funding from Cancer Research UK (FC001317), the UK Medical Research Council (FC001317), and the Wellcome Trust (FC001317) (G.S., J.B., and M.d.G); and ERASMUS+ (J.B.).

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REFERENCES


STAR METHODS

KEY RESOURCES TABLE

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CONTACT FOR REAGENT AND RESOURCE SHARING

Further information and requests for reagents should be directed to and will be fulfilled by the Lead Contact, Buzz Baum (b.baum@ucl.ac.uk).

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Drosophila Genetics

The following stocks were used: w^{118} (Bloomington: 3605), sqh::mCherry (Martin et al., 2009), sqh^{A3}; sqh-GFP42 (Royou et al., 2004), DE-cadherin-GFP (Huang et al., 2009), ubi-E-cadherin-GFP (Oda and Tsukita, 2001), ubi-Bazooka-mCherry (Bosveld et al., 2012), sqh-Rok^{A/16A-Venus (Simoes Sde et al., 2014), tubP-GAL80[ts] (BL7018) dsRNA interference (RNAi) and overexpression constructs were expressed using the UAS/GAL4 system (Brand and Perrimon, 1993; Busson and Pret, 2007). Notum-specific promoter: pannier-GAL4 (Bloomington: 3039). Rok overexpression: UAS-Rok^{CAT} (Verdier et al., 2006). The following RNAi lines were used to
silence gene expression: armadillo (VDRC: KK107344), Shotgun (NIG: 3722R-1), Rok (VDRC: KK104675), Slik (VDRC:GD43783), sds22 (VDRC: GD42051), MYPT75-D (NIG: 68976R-1).

Adult fly crosses were kept at room temperature (21°C). Fly food recipe: 39 l dH2O, 675 g yeast, 390 g soy flour, 2.85 kg yellow cornmeal, 224g agar, 3 l light corn syrup, 188 ml propionic acid. After 2-3 days of egg laying, stocks were flipped and the tube, containing the eggs, was transferred to 18, 25, or 29°C for larval development. The incubation temperature used is stated with the experimental genotypes. AP age develops in real-time at 25°C, and at approximately half pace at 18°C. For the majority of imaging, pupae were staged at 12 h AP (at 25°C) for dissection and imaging, so at 0 h they were transferred to 18°C overnight for imaging the next day. For later developmental stages pupae were moved between 18 and 25°C accordingly.

**Experimental Genotypes**

<table>
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<tr>
<th>FIGURE</th>
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<th>INCUBATION (°C)</th>
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METHOD DETAILS

Notum Dissection for Live Imaging
For live imaging, flies were raised to 11.5 to 12 h AP (at 18, 25, or 29°C) and fixed to a microscope slide with double-sided Sellotape, as (Georgiou et al., 2008). The pupal case was removed to the abdomen to expose the notum. This is achieved by removing the operculum with forceps, cutting down the mediolateral side of the fly with dissection scissors and pealing the case off (Zitserman and Roegiers, 2011). A stack of 18 x 18mm slides was glued to the slide, with clear nail varnish, anteriorly (5 slides) and posteriorly (4 slides). A coverslip was spread with halocarbon oil 700 (Sigma – H8898) and rested onto the stacks so that it is in contact, but not crushing, the notum. The coverslip was glued in place with clear nail varnish at the posterior end. Flies were allowed to settle for 10 mins prior to imaging.

Microscopes Used for Live Imaging
The microscope used for each experiment is stated with the experimental genotypes:

Leica SPE2 scanning confocal with a 40x 1.3 NA oil objective.
For 30 s junction dynamics imaging. z-slices were 1 μm, spanning 5 μm and for long-term time-lapse at 5 min intervals (1 μm x 20 z-slices).
Leica SP5 inverted confocal with a 40x 1.3 NA oil objective and 63x 1.3 NA oil objective.
For imaging of fixed samples.

Nikon Eclipse Ti-E inverted system with a Yokogawa CSU-X confocal spinning disk unit fitted with an Andor EMCCD camera using a 100x 1.4 NA objective.
For two-color time-lapse (30 s interval) imaging.
Z-slices = 0.5 μm covering 4 μm. Maximum intensity projections were made of three slices in the focal plane for each time-point. 488 laser for GFP imaging at 5% power for 50 ms exposure. 561 laser for mCherry imaging at 20% power for 150 ms exposure.
Carl Zeiss LSM 200 M with a Yokogawa CSU-X confocal spinning disk unit and Andor Zyla cSMOS 5.5 camera with a 63x 1.3 NA water objective.
For two-color single time-point imaging at different stages of pupal development.
Z slices = 0.5 μm covering 4 μm. Maximum intensity projections were made of three slices in the focal plane.
Carl Zeiss LSM510 Meta upright confocal microscope with a Plan-Neofluor 40x/1.3 Oil DIC objective.
For laser ablation of adherens junctions.

Laser Ablation
Laser ablations were undertaken as (Marinari et al., 2012). Flies were prepared on a microscope slide for live imaging as described above. DE-cadherin-GFP was visualised using 488 nm light from an Ar-Kr laser with a Plan-Neofluor 40x/1.3 Oil DIC objective, coupled to a Zeiss LSM510 Meta upright confocal microscope. Image acquisition prior to, and after, ablation was at 1 s intervals. Junctions were ablated with 720 nm multiphoton excitation from a Chameleon-XR Ti-Sapphire laser (AIM, Zeiss). Junctions were ablated by scanning over a 3x3 pixel region of interest (0.009 μm²) at 25% laser power, for 1 iteration, with a dwell time of 2.56 μs / pixel.

Notum Dissection for Immunohistochemistry
For p-Myosin II staining 12-13 h AP nota were dissected in PBS before fixation in 4% paraformaldehyde in PBS for 20 min at RT. Samples were washed and permeabilized for 3x 10 min in PBT (PBS with 0.1% Triton X-100) at room temperature (RT). Nota were incubated with 1:1 PBT: blocking buffer (5% BSA, 3% FBS in PBS). Incubation with primary antibodies was undertaken at 4°C overnight (O/N) prior to 4x PBT 10 min washes. Incubation with secondary antibodies was carried out for 1-2 h at RT in PBT, with gentle shaking. Nota were 3x washed with PBS for 10 mins and kept in 50% glycerol in PBS at 4°C O/N before mounting. TritC-Phalloidin (1:500) and DAPI (1:2000) staining was undertaken during the second wash. Nota were mounted with 50% glycerol in PBS and imaged within days. The following primary and secondary antibodies were used: Phospho-Myosin Light Chain Ser19 (rabbit, 1:30 dilution, Cell Signalling 3671), anti-GFP (chicken, 1:500), goat anti-chicken Alexa 488 (1:250), goat anti-rabbit Alexa 647 (1:250).

Theoretical and Computational Models
Theoretical and computational models used in this study are described in Methods S1.
QUANTIFICATION AND STATISTICAL ANALYSIS

Image Processing and Statistics
All images presented were processed with FIJI software (http://fiji.sc/Fiji) (Schindelin et al., 2012) and Adobe Illustrator CS5.1 (Adobe Systems, Inc.). Graphs were produced using GraphPad Prism 6 (GraphPad Software, Inc). Statistical analyses were performed in Prism. D’Agostino and Pearson omnibus normality tests were used to determine if data were Gaussian. Nonparametric datasets were compared using a Kolmogorov-Smirnov test, or a Mann-Whitney test, where appropriate and stated in legend. Data from a Gaussian distribution were compared using unpaired Student’s t-tests.

Laser Ablation Measurements
Kymographs were used in order to calculate vertex displacement velocities. In FIJI, straight lines (1 pixel thick) were drawn across the pre-ablated junction and the ‘Dynamic Reslice’ function was used to produce a kymograph (X = length, Y = time) of each ablation event. The ‘segmented line’ tool was used to plot the progression of each of the two junction vertices over time. In Matlab, the distance between the two X,Y vertex coordinates at each time point was calculated (function supplied below). Total displacement between vertices was then plotted in Prism at 10 sec after ablation. The first time frame after ablation was not used for this study, as the initial vertex displacement measure, because unlike in other tissues, such as the wing disc (Mao et al., 2013), measurable and significant displacement was not observed in the notum in the first frame after ablation.

In Excel, the laser ablation data should be saved as: Column A: Vertex 1 coordinate; Column B: Time Frame of V1; Column C: Vertex 2 coordinate; Column D: Time Frame of V2; Column E: Ablation Frame; Column F: Distance Scale; Column G: Time interval.

MATLAB function for quantifying distance between XY coordinates:

function magic1
    clear all
    close all
    clc
    [filename, url] = uigetfile(‘*.xlsx’, ‘select data file’);
    data = importdata([url, filename]);
    vertex1 = data.data(:,1);
    vertex2 = data.data(:,3);
    time1 = data.data(:,2);
    time2 = data.data(:,4);
    vertex1(isnan(vertex1)) = [];
    vertex2(isnan(vertex2)) = [];
    time1(isnan(time1)) = [];
    time2(isnan(time2)) = [];
    startTime = max(min(time1),min(time2));%seconds
    endTime = min(max(time1),max(time2));
    timeInterval = 1; %seconds
    time = (startTime:timeInterval:endTime)’;
    V1 = interp1(time1,vertex1,time);
    V2 = interp1(time2,vertex2,time);
    length = abs(V2-V1);
    ablationTime = 0; %CHANGE
    output = [time-ablationTime,V1,V2,length];
    xlswrite(‘result.xls’ ,output);
    end

Aspect Ratio Quantification
For Figure 1F, the aspect ratio measure was made using FIJI, from an ellipse fit to a selection. Fit ellipse is found within FIJI at Analyze – Measure – Fit Ellipse. The macro used to fit an ellipse can be found at: http://imagej.nih.gov/ij/macros/DrawEllipse.txt. The area, centroid and orientation of the original selection are retained. The aspect ratio is calculated by dividing the major axis of the ellipse with the minor axis.

For Figures 1H–1J, the aspect-ratio for the four-cell clusters involved in a neighbor exchange was measured. The center of area (CoA) for each cell was calculated, and the internal aspect ratio defined as the distance between the CoA of cells losing an edge divided by that the cells gaining an edge. For the external aspect ratio, the axis between the CoAs was extended to the point it intersected with the perimeter. The external ratio was then calculated as the distance between perimeters of cells losing an edge, divided by the distance between perimeters of cells gaining an edge. This definition of the aspect ratio, would mean that an elongation of the four-cell cluster along the same axis as the expansion of the T1 junction, would give an increase in aspect ratio.
Image Segmentation

Images were segmented using Packing Analyzer V2.0. Tutorials are available at: https://idisk-srv1.mpi-cbg.de/~eaton/ (Aigouy et al., 2010).

Protocol:

- Save movie in FIJI as an 8-bit Image Sequence TIFF.
- Open Packing Analyzer, drag and drop the Image Sequence into the List.
- UnderInit, change the parameters to 3.8 (top) and 1.9 (bottom), or change these parameters as you see fit.
- Click Detect Bonds. A notum image with the above parameters is likely to be over segmented. Save.
- Under Correction, correct the mistakes. Right click to delete junctions. Left click and hold to draw new junctions. Change the pencil size, on the left above the tabs. Use a large pencil size to delete lots of junctions at once. Click second green tick to apply changes. Save.
- To apply changes to the next image in the sequence click Seed Next.
- Once finished, Post Process, change 4 way vertices cutoff to 3. Click Finish All. (large datasets will take 10-15 mins).
- Rectenter tab, click autocenter based on 2D correlation (10-15 mins).
- Tracking, click Track cells (10 mins).
- Tracked bonds, click track bonds (10 mins).
- Plots, can plot all bonds or plot all cells. To plot individual cells/junctions draw lines through cells/junction of interest in Current Image then click plot selected cells / plot selected bonds. Always exclude border bonds or cells.
- Virtual cloning, draw a line through cells of interest in Current Image, click track clones. Can go back to Plots and plot clone info.
- Under the Viewer tab you can see a range of segmentations for each image.
- In the folder where the original Image Sequence is saved, each image will have a folder containing all the files produced.
- Number the bd_fate files and create an Image stack in FIJI.
- Plots will be saved as .csv files in first image folder ....000.
- Open in Excel, highlight first column. Go to Data, and click Text to columns. Highlight the whole dataset and go to Data, Sort. Sort by ID then Frame Number. Sizes (junction lengths and areas) are in pixels.

Analyzing Junction Fluctuation Behavior

In order to extract data for the behavior of junctions and cells in a format amenable to the type of analysis we wanted to do, we developed a custom software package. The code is written in Matlab using class-based object-oriented programming. It detects junctions and cells from segmented images, corrects for drift, tracks junctions and cells between frames, and calculates connectivity within the tissue. This makes it possible to extract time series data for various properties, analyze spatial correlations, and detect when cells change neighbors.

The flow of data within the code is as follows:

- Input segmented time-lapse images.
- In case of microscope drift, generate set of stabilized images by subtracting net translation.
- Identify individual junctions in each image and store these as objects.
- Track junctions between frames and assign a unique ID to each junction.
- For each junction, find the IDs of neighboring junctions.
- Detect individual cells in each image and store these as objects.
- Track cells between frames and assign a unique ID to each cell.

The input for the code are segmented 8-bit image sequences of the Drosophila notum produced with Packing Analyzer V2.0 (Aigouy et al., 2010), tutorials are available at: https://idisk srv1.mpi-cbg.de/~eaton/. Segmentation results in skeletonized images where the width of a cell-cell interface (the junction) is 1 pixel.

In some cases, the tissue drifts relative to the microscope during imaging. The algorithm for tracking junctions and cells cannot deal with large-scale deformation or significant displacement of the tissue between frames. To deal with this, we used Optical Flow Analysis (modified from the OFA algorithm available at http://cs.brown.edu/people/black/code.html and described in Sun et al., 2010) to calculate the flow field for each consecutive pair of frames. Taking the average of the flow field gives the direction and magnitude of the net translation of the tissue. Mapping the images into a larger space, by subtracting the cumulative net translation for each time point, yields a set of stabilized images that can be input into the code. The first part of the code identifies individual junctions in each image. Specifically, the code initiates a junction and ‘walks’ along the bright pixels in the image, storing the coordinates along the way, then terminating the junction when a vertex is reached. This is then repeated until all pixels in the image have been visited. Within the code, each junction is an object with associated properties. At this stage, only the fields for the vertex coordinates and junction coordinates are filled.

The code then calculates various properties of junctions, including: vertex 1 and 2 positions, junction coordinates, length, midpoint, angle and neighbor IDs. Since junctions can be curved, the vertex-vertex distance is not an accurate measure of junction length. In
addition, discretisation has the consequence that calculating the length by summing the distance between pixels along the segmented junction would slightly overestimate the length. Instead, junction length is calculated as the diagonal distance connecting consecutive blocks of pixels.

The next step involves tracking junctions between frames and assigning a unique ID to each junction, to make it possible to extract time series for various properties and detect changes in connectivity. The tracking of junctions is done by finding the midpoints of junctions and using these coordinates as the input for a particle tracking algorithm (based on code available from the Mathworks repository, written by John C. Crocker (Crocker et al., 1996)). The tracking algorithm takes the coordinates of the midpoints at time t and considers all possible matches with the midpoints at time t + 1 to choose the pairings that minimize the total squared displacement. This is then used to assign each junction a unique ID that identifies it across frames. The results were validated by visual inspection of the assigned IDs – specifically by creating a movie of junctions colored according to their ID, making errors in continuity easy to spot. Having assigned a unique ID to each junction, the connectivity of the tissue is found and the IDs of neighboring junctions are stored. In addition to junctions, the code also detects and tracks cells in the time-lapse images and stores each as an object. The properties for this class include: Cell ID, junction IDs, midpoint, area, perimeter, vertices and angles. Cells are detected using the junction objects. Specifically, the code starts at one junction and moves to neighboring junctions in a clockwise fashion until getting back to the first one, thereby identifying the junctions that make up a cell. This is repeated in a counterclockwise fashion for the same junction. To avoid storing the same cell multiple times, the code loops over junctions starting from j = 1 and requires that only neighboring junctions with larger values of j can be traversed - if that is not available, the code breaks and starts from a different junction instead. Next, the area, perimeter length and midpoint of each cell is calculated. The midpoints are used to track cells between frames and assign unique IDs in the same way as described for junctions.

**Detecting Neighbor Exchange Events**

Neighbor exchange events are difficult to quantify manually. Therefore, we wrote an algorithm to detect transitions and make it possible to extract quantitative data related to the junctions and cells involved. The code detects all junctions that contract to a four-way vertex and expand back out, and determines whether they change neighbors in the process. For the DE-cadherin-GFP imaging at 30 s intervals on the Leica SP2 scanning confocal, the diameter of four-way vertices is ~6 pixels, corresponding to 538 nm. Segmentation of very short junctions, and especially four-way vertices, is difficult and error-prone. In particular, segmented short junctions tend to ‘flip’, changing orientation and neighbors. If not corrected, such false neighbor exchange events would bias subsequent analysis. To ensure the quality of the data that form the basis of this study, we manually checked every computationally detected event by looking at the corresponding junction in the fluorescent time-lapse images. We used the criteria that the extension of a junction, coming from a four-way vertex configuration, should be stable and the change in cell neighbors should be clearly visible in the fluorescent images. Events that did not fit these criteria were excluded from subsequent analysis. In addition, we checked that the time point for the event, as identified by the algorithm, was consistent with when a four-way vertex was reached in the fluorescent images.

**Fluorescence Intensity Measurements**

We concurrently imaged Myosin II and junction dynamics using transgenic fly stocks expressing both Sqh-GFP (Myosin) and Bazooka-mCherry (adherens junction marker). To correlate Myosin II intensities with junction dynamics (Figure 4), we developed code to extract the time series data for Myosin intensities on individual junctions. For each junction, we used the pixel coordinates from the segmented images to identify the junction in the fluorescent images. To include the fluorescence intensity across the width of the junction, we performed a morphological dilation to give each junction an average width of 7 pixels. This corresponds to a width of 488 nm, for the time-lapse imaging taken at 30s intervals with a resolution of 0.06974 mm/pixel. The vertices tend to be the bright regions in the image and including them would give rise to artificial artifacts in the correlation functions - e.g. as junctions contract to a four-way vertex and expand back out, and determines whether they change neighbors in the process. For the DE-cadherin-GFP imaging at 30 s intervals on the Leica SP2 scanning confocal, the diameter of four-way vertices is ~6 pixels, corresponding to 538 nm. Segmentation of very short junctions, and especially four-way vertices, is difficult and error-prone. In particular, segmented short junctions tend to ‘flip’, changing orientation and neighbors. If not corrected, such false neighbor exchange events would bias subsequent analysis. To ensure the quality of the data that form the basis of this study, we manually checked every computationally detected event by looking at the corresponding junction in the fluorescent time-lapse images. We used the criteria that the extension of a junction, coming from a four-way vertex configuration, should be stable and the change in cell neighbors should be clearly visible in the fluorescent images. Events that did not fit these criteria were excluded from subsequent analysis. In addition, we checked that the time point for the event, as identified by the algorithm, was consistent with when a four-way vertex was reached in the fluorescent images.

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For each junction, we sum over the intensity of pixels within the region covered by the morphological dilation. There is a slight bleaching of the tissue over time, leading to a gradual decrease in intensity. We remove the trend associated with bleaching in the following way: for each time frame, we sum the total intensity for all pixels within the dilated junctions ($I_{tot}$) and calculate the total number of pixels ($P_{tot}$). For each junction, the total intensity $I_j$ is normalized by $I_j/P_{tot}$, such that the average intensity per pixel is equal to 1 for every time frame $\sum_j I_j/\sum P_j = 1$. For each junction, we calculate the ‘normalized average intensity’ by taking the total intensity for a junction, normalizing it as described, then dividing by the number of pixels in the junction. Qualitatively, dividing by the number of pixels gives the same results as dividing by junction length.
Supplemental Information

Myosin II Controls Junction Fluctuations to Guide Epithelial Tissue Ordering

Scott Curran, Charlotte Strandkvist, Jasper Bathmann, Marc de Gennes, Alexandre Kabla, Guillaume Salbreux, and Buzz Baum
Supplemental Inventory

**Supplemental Figures**

Figure S1: Related to Figure 1.
Figure S2: Related to Figure 2.
Figure S3: Related to Figure 4.
Figure S4: Related to Figure 5.
Figure S5: Related to Figure 6.
Figure S6: Related to Figure 7.
Figure S7: Related to Figure 7.

**Supplemental Tables**

Table S1: Related to Figure 2A-D and Figure S2.

**Supplemental Movies**

Movie S1: Related to Figure 1.
Movie S2: Related to Figure 1.
Movie S3: Related to Figure 5.
Movie S4: Related to Figure 6.

**Supplemental Theory**

Methods S1: Related to STAR Methods
Figure S1. Related to Figure 1. Neighbour exchange events are reversible

(A), Bar charts showing the proportion of junctions that undergo unidirectional (one), bidirectional (two), and multidirectional (three or more) neighbour exchange events. n = 3 nota (not including nota from Figure 1C) over 75, 80, and 115 mins, respectively, labelled with DE-cadherin-GFP and imaged at 30 s intervals. (B), Kaplan-Meier survival curve showing the probability that a neighbour exchange configuration persists for a given length of time. Events are 'censored' if the reverse transition has not occurred by the end of the movie.
The 95% confidence intervals are estimated using Greenwood’s Formula. The probability of a configuration persisting for at least 150 min, along with the 95% confidence interval = 0.687 [0.5175, 0.8567].

(C), Histogram of junction lengths in a single frame at 12h AP. n = 771 junctions / 3 nota. Average junction length = 4.69 ± 2.315μm.

(D), Junction lengths in a single frame at 12 h AP versus angle, with respect to the AP midline (0°). n = 771 junctions / 3 nota.

(E), Related to Figure 1H. Change in aspect ratio from t = -15 mins to t = +15 mins. A paired t-test was used to compare the aspect ratios at t = -15 mins and t= + 15 mins. Internally, the aspect ratio changes during a T1 event (p < 0.0001). Externally there is no change in aspect ratio (p = 0.0502). n = 33 exchange events from 4 flies.

(F), Diagram of a neighbour exchange event. The red junction represents the transition junction that is lost and gained. An ellipse (black) is fit to the four cells involved in the transition, and the feret angle of the ellipse is measured with respect to the midline (0°).

(G), The feret angle of fit ellipses are plotted at t=-15, and t=+15 mins, with paired results connected by a line. n = 33 exchange events from 4 flies.
Figure S2. Related to Figure 2. Persistence length analyses results are consistent for a range of filter settings

(A), Time series line plot for a junction fluctuating in length. The data have been filtered using a moving average Hanning window, with a filter setting of 5, and split into segments where the junction is contracting or expanding. The persistence time (PT) and persistence length (PL) are defined as the duration of a segment and the change in junction length, respectively. (B), Histogram of persistence lengths for a filter setting of 5. n = 60737 segments, 4 nota (green line). Best fit for an exponential function of the form $1/\tau \exp(-t/\tau)$. The parameter is $\tau = 0.197$ mm, with confidence bounds [0.1958 0.1989]. Inset, Number of segments in each bin of the histogram plotted on semi-log along with the exponential fit. (C-E), Line plots of the length of a representative junction (blue) over time, for a junction that undergoes a neighbour exchange event at 40 min (marked by a red star). The time series has been split into segments where the junction length is monotonically increasing or decreasing. This was achieved by filtering the data using a moving average Hanning window
(green) - if the first derivative changes sign or is zero with opposite signs on either side of that point, a segment boundary is placed. Using a higher setting for the filter, results in fewer segments. The filter settings used are (C) 10, (D) 20 and (E) 40. The analysis is done using the raw data - the filter is only used to establish the position of the segment boundaries. The segments on either side of a T1 event (marked with red dotted lines) are defined as T1-segments. (F-H), Persistence length distributions for (left) non-T1 and (right) T1 segments. The filter settings used are (F) 10, (G) 20, (H) 40, as shown in C-E. The number of segments included in each histogram are stated in the top right of each graph. Higher levels of filtering result in longer, and therefore fewer segments. The first and last segment of each time series is excluded from the analysis, regardless of whether a T1 event occurs (therefore the number of T1 segments included can vary with filter level). The distributions for T1 and non-T1 segments are compared used a two-sample Kolmogorov-Smirnov test. The p-values are listed in Table S1 and in every case the statistical test, at the 0.05 significance level, supports the null hypothesis that the samples are drawn from the same underlying distribution.
Figure S3. Related to Figure 4. Myosin and Bazooka intensities are uncorrelated and do not exhibit tissue-wide polarity. Cell areas and medial Myosin intensity do not display pulsatile behaviour and changes in medial Myosin intensity do not precede changes in cell area.

(A), Normalized junction intensities of Baz-mCh versus junction angle, as measured with respect to the AP midline (0°). Slope (with 95% confidence bounds) = -0.0003 (-0.00061 0.00011). Spearman’s Rank = -0.048. n = 688 junctions / 3 nota. (B), Normalized junction intensities of Baz-mCh versus junction length. Slope (with 95% confidence bounds) = 0.003 (-0.0019 0.0070). Spearman’s Rank = 0.058. n = 688 junctions / 3 nota. (C), Normalised (to mean intensity) paired junctional intensities of Baz and Myo-II show no correlation. Spearman’s Rank = 0.024. n = 688 junctions / 3 nota. (D), Apical surface projection of a live nota imaged with Bazooka-mCherry (left panel) and Spaghetti-Squash-GFP (right panel). Scale bar = 5μm. The medial region included in the analysis is outlined in green. (E), A line plot showing a representative example of medial Myo-II-GFP (green) and cell area (blue) plotted as a function of time. Myo-II-GFP intensity is normalised to mean tissue intensity and the line plot shows the average Myo-II-GFP intensity per pixel. (F), Mean autocorrelation for cell area variation of individual cells $\Delta A(t) \Delta A(t + \Delta t)/(\langle A \rangle^2)$ as a function of lag time $\Delta t$. The autocorrelation is normalised by the time-averaged cell area $A$ for each cell. n = 115 cells / 3 nota. (G), Mean autocorrelation for medial Myo-II-GFP intensity in individual cells $\Delta I(t) \Delta I(t + \Delta t)/(\langle I \rangle^2)$ as a function of lag time $\Delta t$. The autocorrelation is normalised by the time-averaged medial intensity for each cell. n = 115 cells / 3 nota. (H),
Mean normalised crosscorrelation for medial Myosin intensity and cell area \( \frac{\Delta I(t) \Delta A(t + \Delta t)}{\sigma_I \sigma_A} \) as a function of lag time \( \Delta t \), with \( \sigma_I \) and \( \sigma_A \) the intensity and cell area S.D. The minimum occurs at zero lag. The normalised crosscorrelation is calculated for each cell and then averaged over all cells in the analysis. \( n = 115 \) junctions / 3 nota, imaged at 30 s intervals for 60 mins.
Figure S4. Related to Figure 5. 2D vertex model fitting parameters

(A-P) Plots showing the dependency of tissue properties on the fitting parameters. (A, E, I, M) T1 transition rate, (B, F, J, N) extrinsic area fluctuations ($CV_A$), (C, G, K, O) junction length intrinsic fluctuations ($CV_L$), and (D, H, L, P) relative perimeter fluctuations ($CV_p$), (dependency on the dimensionless model parameters: (A-D) dimensionless ratio of mean line tension over cell bulk elastic modulus ($\gamma/Kl^2$), (E-H) normalized intrinsic...
noise intensity \((\sigma_l/\gamma)\), (I-L) characteristic packing time \((\tau_p/\tau_m)\) and (M-P) normalized standard deviation of preferred cell areas \((\sigma_h/l^2)\). Dotted horizontal lines: experimental measurement, with shaded area indicating standard deviation. Dotted vertical lines: parameters used in wild-type simulations. (Q) Junction length distribution of simulations with wild type parameter settings to data obtained from experiments. (R) Polygon distribution of simulations with wild type parameter settings and data obtained from experiments.
Supplementary Figure 5

Figure S5. Related to Figure 6. Increasing neighbour exchange rates observed between 24 and 28 h AP are due to sensory organ precursor cell growth.

(A) Time-lapse montage of a representative SOP cell (circled by green dotted line), labeled with DE-cadherin-GFP, imaged from 20 to 30 h AP. Yellow arrows, label junctions that have been gained via neighbour exchange events. Scale bar, 5 µm. (B-C) Line plots showing the mean (with S.D) (B) SOP cell area and (C) the mean number of neighbours (polygon number) for SOP cells, from 20 – 30 h AP. Dotted box from 24-28 h AP indicates the time at which the T1 transition rate, as shown in Fig 6G, increases. n = 30 SOP cells / 3 nota.
Figure S6. Related to Figure 7. Decreased Moesin activity causes a loss of junctional Myosin II and increases the medial pool. Phospho-null and phospho-active Myosin increase and decrease neighbour exchange rates, respectively.

(A) Apical surface projections of DE-cadherin-GFP labelled nota for control, UAS-sqhAA (Phospho-null Myosin II), UAS-sqhEE (Phospho-active Myosin II), UAS-Slik RNAi (decreased Moesin activity – note wiggly junction phenotype caused by pulling of medial Myosin on the junction), UAS-Moesin-TD (increased Moesin activity) and UAS-sds22 (increased Myosin and Moesin activity) driven by pnr-GAL4. Scale bar = 5µm. (B) Quantification of total vertex displacement at 10 s after laser dissection of single junctions in 12-13.5 h AP

(C) (D) (E) (F) (G)
pupae expressing Slik RNAi and sds22 RNAi. Dots indicate individual experiments, line represents median. 5-7 flies/condition. P-values calculated from unpaired t-tests. (C) Maximum surface projection images of nota live-imaged with ubi-DE-cadherin-GFP and Spaghetti-Squash-mCherry (MRLC) labelling total Myosin levels for control, Slik RNAi (note lack of Myosin on junctions) and sds22 RNAi (increased junctional Myosin). (D) Fixed-stain images for DE-cadherin-GFP (anti-GFP), F-actin (Phalloidin) and p-Myosin II (S19) for control, reduced Moesin activity (Slik RNAi) and increased Myosin and Moesin activity (sds22 RNAi). Scale bar = 5µm. (E) Increased zoom of vertices in D. Arrows for Slik RNAi highlight DE-cadherin junction, and absence of p-Myosin. Arrows for sds22 RNAi label junction break at 3-way vertex. Scale bar = 1µm. (F) Apical surface maximum projection montages of DE-cadherin-GFP labelled nota showing representative neighbour exchange events for control, sds22 RNAi and Slik RNAi. White arrows label junction breaks at the vertex during the sds22 RNAi transition. Slik RNAi nota can still undergo neighbour exchange in the absence of junctional Myosin – akin to Rok RNAi. (G) Quantification of normalised T1 transition rates for altered levels of Myosin activity. Dot indicates mean, tails show the data range. n = 3-4 flies / condition. P-values calculated from Kolmogorov-Smirnov tests.
Figure S7. Related to Figure 7. A tissue-wide reduction in Myosin activity increases midline delamination levels and causes cells to delamate outside of the midline.

(A) Apical surface maximum intensity projections of representative nota, labelled with ubi-E-cadherin-GFP, for altered Myosin II activity, showing the midline (dotted green line) with cells that delaminate (yellow) up to 18 h AP. Scale bar = 5µm. (B) Bar chart of percentage midline delamination for decreased (Rok RNAi) and increased (MYPT75-D RNAi and sds22 RNAi) Myosin II activity. Bar values indicate mean. P-values calculated from unpaired t-tests vs control. N = 4-6 nota per condition. (C) A representative notum with cells labelled outside of the midline that undergo basal delamination (blue). Midline region shaded red. Half blue cell indicates a daughter cell that delaminates after division. (D) Time-lapse montage of a cell (star labelled in C) that progressively loses junctions, and area, through time before extrusion. (E) Delamination levels were measured outside of the midline up to 18 h AP for control and Rok RNAi. Delamination levels were measured as a percentage of the total number of outer midline cells at 12 h AP. N = 6-7 flies with 110-237 (mean = 156) cells at 12 h AP. p-value calculated from a Mann-Whitney statistical test.
**Supplementary Tables**

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**Table S1. Related to Figure 2A-D and Figure S2.**

The persistence length distributions for T1 and non-T1 segments are compared using a two-sample Kolmogorov-Smirnov test. The null hypothesis, $H_0$, is that the samples are drawn from the same underlying distribution. The analysis was carried out for seven different levels of filtering, between 10 and 40, and in every case the statistical test supports the null hypothesis at the 0.05 significance level. Note that, since we are testing a set of statistical inferences simultaneously, the appropriate significance level for the individual hypothesis tests is lower than the significance level for the set as a whole. Using the Bonferroni correction, we would get $\alpha = \frac{\alpha}{k} = 0.05/7 = 0.007$. P-values are all above 0.05.
**Supplementary Movie Legends**

**Movie S1. Related to Figure 1.**
This movie shows a region outside of the midline, of a wild-type developing pupal notum, visualised with DE-cadherin-GFP. The movie covers the period 12-13.5 h AP, prior to the onset of cell division and delamination, during which time cells undergo neighbour exchange (indicated in Figure 1), caused by fluctuations in junction length (indicated in Figure 2). Time interval between frames is 30 s and the video length is 1 h 30 mins. Scale bar, 10 µm.

**Movie S2. Related to Figure 1.**
This movie shows a four-cell cluster undergoing a uni-directional neighbour exchange event (indicated in Figure 1), visualised with DE-Cadherin-GFP. This is an event that occurs in Movie S1. Successive yellow arrows throughout the movie label the junction that is lost, the four-way vertex generated through junction loss, and the subsequent newly formed junction. Note neighbour exchange events also occur on the edge of the cluster throughout the movie. Time interval between frames is 30 s and the video length is 1 h 30 mins covering 12 - 13.5 h AP. Scale bar, 10 µm.

**Movie S3. Related to Figure 5.**
This movie shows three simulations of the 2D vertex model at low (0.8), control (1.0) and high (1.2) mean line tensions ($\gamma/\gamma_0$) (indicated in Figure 5). The edge colour, of each interface, corresponds to the level of line tension relative to it’s mean (blue = low, red = high). Virtual time interval between frames is 15 s, with the video length 83 min 15 s.

**Movie S4. Related to Figure 6.**
This movie shows an example of a control laser ablation experiment (indicated in Figure 6B and 7C). Junctions outside of the midline are visualised with DE-cadherin-GFP between 12 and 13.5 h AP. The time interval between frames is 1 s with the laser ablation occurring at t = 0, indicated by a yellow star. The video length runs from 16 s pre-ablation to 120 s post-ablation. Scale bar, 5 µm.