Apposition of iroquois expressing and non-expressing cells leads to cell sorting and fold formation in the Drosophila imaginal wing disc

Eugenia Villa-Cuesta†1,2, Esther González-Pérez†1 and Juan Modolell*1

Address: 1Centro de Biología Molecular Severo Ochoa, CSIC and UAM, Cantoblanco, 28049 Madrid, Spain and 2Department of Ecology and Evolutionary Biology, Bow G-W, Brown University, Providence, RI 02912, USA

Email: Eugenia Villa-Cuesta - Eugenia.Villa-Cuesta@brown.edu; Esther González-Pérez - egperez@cbm.uam.es; Juan Modolell* - jmodol@cbm.uam.es

* Corresponding author    †Equal contributors

Abstract

Background: The organization of the different tissues of an animal requires mechanisms that regulate cell-cell adhesion to promote and maintain the physical separation of adjacent cell populations. In the Drosophila imaginal wing disc the iroquois homeobox genes are expressed in the notum anlage and contribute to the specification of notum identity. These genes are not expressed in the adjacent wing hinge territory. These territories are separated by an approximately straight boundary that in the mature disc is associated with an epithelial fold. The mechanism by which these two cell populations are kept separate is unclear.

Results: Here we show that the Iro-C genes participate in keeping the notum and wing cell populations separate. Indeed, within the notum anlage, cells not expressing Iro-C tend to join together and sort out from their Iro-C expressing neighbours. We also show that apposition of Iro-C expressing and non-expressing cells induces invagination and apico-basal shortening of the Iro-C cells. This effect probably underlies formation of the fold that separates the notum and wing hinge territories. In addition, cells overexpressing a member of the Iro-C contact one another and become organized in a network of thin strings that surrounds and isolates large groups of non-overexpressing cells. The strings appear to exert a pulling force along their longitudinal axis.

Conclusion: Apposition of cells expressing and non-expressing the Iro-C, as it occurs in the notum-wing hinge border of the Drosophila wing disc, influences cell behaviour. It leads to cell sorting, and cellular invagination and apico-basal shortening. These effects probably account for keeping the prospective notum and wing hinge cell populations separate and underlie epithelial fold formation. Cells that overexpress a member of the Iro-C and that confront non-expressing cells establish contacts between themselves and become organized in a network of thin strings. This is a complex and unique phenotype that might be important for the generation of a straight notum-wing hinge border.
mented to prevent cells from intermingling along boundaries between tissues and sometimes even within a tissue. In *Drosophila*, developmental boundaries were discovered in the wings, by means of genetically marked recombination clones, as straight lines that proliferating cells did not trespass [1] (reviewed by [2-5]). One of these boundaries corresponded to an invisible line that subdivided the wing into anterior (A) and posterior (P) compartments. Another boundary separated the dorsal (D) from the ventral (V) compartment, and corresponded to the wing margin. Subsequent work showed that the inheritable on or off state of the selector homeobox genes *engrailed* and *invected* (on in P and off in A) and *apterous* (on in D and off in V) defined the four compartments of the wing. Later work showed that short-range cell-signalling occurred from P to A and between D and V cells. This was mediated by Hedgehog and Notch, respectively, and it was essential to maintain the A/P and D/V boundaries. Moreover, the short-range signalling gave rise to long-range signalling that organized the growth and patterning of the entire wing.

Differential cell-cell affinities were invoked to explain the presence of these compartment boundaries and their straight and smooth shape [6] (reviewed in [5]). Cells of a compartment would have an affinity for each other higher than for the cells of the neighbouring compartment. At the interface, cell-cell contacts would be minimized leading to straight and smooth boundaries. Aggregation experiments with mixtures of cells expressing either different amounts or different classes of cadherin adhesion molecules showed that cells could indeed sort out [7]. Moreover, overexpression of a single DE-cadherin in cell clones in the wing disc caused the clones to segregate from non-overexpressing cells and fuse together [8]. Thus, quantitative differences in the accumulation of adhesion molecules seem sufficient for cell segregation and may be critical for compartment boundaries to be formed and maintained. Still, the adhesion molecules involved in the compartment boundaries of *Drosophila* remain to be identified.

Cell lineage studies in the adult mesothorax suggested the presence of another cell restriction boundary between the notum and wing hinge [9]. However, the anatomical complexity of the wing hinge prevented its direct visualization by genetic marking in the adult fly. Moreover, analyses performed in the imaginal wing discs, the precursors of most of the mesothorax and the wings, indicated that the notum/wing hinge boundary is not a classical compartment boundary [10] (reviewed in [5]). Here, the notum region is defined by the expression of the homeodomain proteins of the *iroquois* complex (*Iro-C*, [11,12]), namely *Araucan* (Ara), *Caupolican* (Caup) and *Mirror* (Mirr), in the notum territory. The loss of these proteins from prospective notum cells transforms them into wing hinge cells [10]. Hence, the border of the Iro-C domain establishes the notum-wing hinge boundary. Mosaic analyses showed that during proliferation cells can trespass this boundary [10], which indicates that they can change the state of activation of Iro-C and consequently their notum or wing hinge identity. However, the change of identity is only possible during the second and early third instar. Later loss of Iro-C activity does not abolish notum commitment.

Differential cell adhesion is probably involved in formation and/or maintenance of the notum-wing hinge border. Indeed, dissociated notum and wing hinge cells sort out during aggregation [13]. Moreover, the borders of the Iro-C clones within the notum territory are smooth and rounded [10], which suggests that they tend to minimize contacts with the surrounding Iro-C+ cells and possess a differential cell-cell affinity. However, this interpretation is complicated by the fact that the Iro-C clones are surrounded by a fold that spans several cell diameters and seems identical to that which separates the notum and wing hinge territories in the late third instar disc. Conceivably, formation of the fold might promote the roundness and smoothness of the clones.

Here we report that, within the notum anlage, cells not expressing Iro-C tend to join together and sort out from their Iro-C expressing neighbours. This supports a role for differential cell adhesion in formation and/or maintenance of the notum-wing hinge border. We also show that apposition of Iro-C expressing and non-expressing cells induces invagination and apico-basal shortening of the latter. This effect appears to be limited to a few cell diameters from the interface and probably underlies formation of the fold that separates the notum and wing hinge territories of the wing disc. In addition, we analyze in detail the previous finding [10] that cells overexpressing a member of the Iro-C arrange themselves in strings that can give rise to a bidimensional lattice that surrounds and isolates large groups of non-overexpressing cells. The data suggest that cells from different overexpressing clones have a tendency to establish contacts and exert a pulling force parallel to the direction of the strings.

**Results**

**Cell clones lacking Iro-C tend to join together**

Third instar wing discs bearing mitotic recombination clones lacking the Iro-C (*iroDFM3* clones) are noticeable in that they usually show only a single, relatively large clone within the notum territory, even when many clones are present in the hinge and wing regions (Figs. 1A*, C; 43 out of 48 discs examined). Moreover, within the prospective notum territory, the *iro+*/+ “twin spots” often lack their associated *iroDFM3* clones (Fig. 1A*). Sometimes, a large
twin spot is associated with the single, large \textit{iroDFM3} clone (Fig. 1C). We examined the reason for this scarcity of notum \textit{iro} mutant clones by simultaneously inducing \textit{iroDFM3} clones and clones lacking the neutral marker \textit{arm-lacZ}. The distribution of the latter showed that the frequency and size of the clones was similar in the notum and in the wing regions (Fig. 1A'). Hence, the scarcity of notum \textit{iroDFM3} clones could be due to either low viability, clone fusion, or both. Their relatively large size, as observed in the notum region of the late third instar disc, were still roundish (Fig. 2A, E), but lacked a fold around them (Fig. 2B–D, G) and had a peculiar morphology. The apical regions of the mutant cells, as visualized in z optical sections by the adherens junction marker Echinoid (Ed, [14]), were recessed into the tissue (Fig. 2B, arrowhead). This invaginated apical region appeared to be mostly or exclusively formed by \textit{Iro-C} mutant cells (Fig. 2A, B). This was clearer in larger, and presumably, older clones (Fig. 2C). Often, the z sections of these clones had the shape of a pouch, with the opposite apical regions of the cells not apposed to one another (Fig. 2D). The invaginations suggested that the mutant cells underwent an apical-basal shortening. This was clear in very small clones (Additional File 1A–C), which indicated that it was one of the earliest morphological changes undergone by their cells. The larger, pouch-shaped clones often sunk deeply into the adephelial layer (Additional File 1D), although their cells never lost the continuity of their apical regions (Ed or Actin markers) with those of the adjacent non-mutant cells. In this large clones, it is uncertain whether cells at the clone interface also maintain the apico-basal shortening evident in cells located deep into the pouch.

We reexamined in optical z sections the notum-to-hinge transformed \textit{iroDFM3} clones induced during the first/second instar. They were not pouch-shaped but cylindrical (Fig. 2H, I) and the fold surrounding them was formed by wild-type cells in the outer side and mostly, but not exclusively, \textit{iroDFM3} cells in the inner side. We asked whether this morphology, more complex than that of the clones induced at 72 to 96 h AEL, represented a more evolved
stage of the iroDFM3 clones that could not be attained by the late-induced clones, possibly because cell proliferation did not last long enough. Some rare late-induced clones suggested that this might be the case (Fig. 2E–G).

The apical zones of the cells deep in the pouch (Ed staining) of these clones were not disposed as in a concave bowl, but delineated a twisted form in a roughly “w” shape (Fig. 2F, arrowhead). This suggested that the cells at the bottom of the pouch might be recovering their normal apical-basal length, and consequently that, in older clones, the progeny of these cells might reach up to the apical plane of the epithelium (see model in Fig. 2J), producing an “eversion” of the clone. The lengthening would occur only in the internal cells of the clone, namely, those further removed from the border of the clone. Note also that the upper regions of the pouch of that rare clone were formed by wild-type cells (Fig. 2G, arrowhead). This could facilitate, upon eversion, formation of a fold with wild-type cells at the external side and mostly mutant cells at the internal side. According to this model, eversion of the clone would occur while the clone grew, although the rarity of the “w” stage suggests that eversion would be a relatively rapid process. In summary, the data above suggest that in the notum territory the apposition of Iro-C expressing and non-expressing cells causes the latter to undergo complex morphological changes. These would start with an apico-basal contraction and invagination of the mutant cells and, as the clone grew in size, would continue with a lengthening of the more internal cells of the clone (Fig. 2J). A fold at the interface between wild-type and mutant cells would be the end result of this process. The fold would probably help to smooth this interface.

Apical-basal cell shortening and invagination caused by apposition of Iro-C expressing and non-expressing cells probably induces formation of the fold between the notum and hinge territories during the third larval instar. This is supported by the observations that the fold arises at or very close to the border of the Iro-C expressing cells (Additional File 2A, B), the z sections of the fold in the late third instar disc (Additional File 2C) is most similar to those of the early-induced iroDFM3 clones (Fig. 2H, I), clones near the notum/hinge border can display folds continuous with that of the border [10], and removal of the Iro-C+/− interface, removes the notum/hinge fold [15].

**UAS-ara-expressing cells from different clones establish connections**

We further examined the behaviour of apposed cells containing different amounts of Iro-C products by overexpressing araucan, one of the Iro-C homeodomain genes [11], in cell clones. Previous work reported, but did not examine in detail, that in late third instar discs these clones appeared to contact each other and that their cells disposed themselves in thin, linear arrangements separating large, roundish islands of nonexpressing cells [10] (and Fig. 3C). We examined these clones in earlier discs and compared them with clones expressing only the neutral marker UAS-GFP. In young third instar discs, connec-
Figure 3

UAS-ara-expressing clones interconnect with one another. Clones (green) were detected by expression of UAS-GFP, except in H where anti Ara/Caup was used. Counterstaining (in red) was Actin (A-E, K, K’), anti RhoGEF2 (G), anti Ara/Caup (I, J) and (in purple) anti Engrailed (H). (A, B) Mid third instar wing discs with clones overexpressing UAS-ara and UAS-GFP or UAS-GFP alone, respectively, induced 48 hours earlier. Magnified views of the areas within squares are shown in A’ and B’. Interconnections between the clones noticeable in A and A’ (arrowheads) are absent in B, B’. In more mature third instar discs most clones overexpressing UAS-ara (C) are arranged in an interconnected net, but not so the clones not overexpressing the Iro-C gene (D). (E) xy confocal section of the wing pouch region of a disc bearing interconnected UAS-ara clones. (F) A stack of xy images of the same disc is shown in a stereo projection selecting only the UAS-ara expressing cells. Asterisks identify same areas in E and F. UAS-ara expressing cells tend to form “walls” separating large territories of non-expressing cells. In x, y sections these walls appear as thin strings. (G) A very long string (arrow) connecting two large separate clones. (H) An anterior compartment clone (green) and a posterior compartment clone (white) displaying connecting cells (arrow). Cell nuclei were stained with anti Ara/Caup antibody. Purple: Engrailed posterior marker. Picture modified from [16] (data courtesy of R. Diez del Corral). (I, J) xy and z optical sections, respectively, of a connecting string. Cell nuclei of overexpressing cells are shown in yellow or orange. Note that the string can be only one cell thick (arrow). (K, K’) z optical section across a single cell-wide wall. The green channel brightness has been strongly enhanced in K’ to show the body of the cell(s) from apical to basal.

The strings were either straight or smoothly curved around the masses of non-expressing cells. This suggested that they exerted a force in the longitudinal direction, like that of a thread wrapping around a stick. This interpretation was reinforced by the finding that the clones did pull on the dorsal/ventral compartment border, distorting it (Figs. 4A, A’), and that the contour of the cells of the strings, most easily observed in the apical sections, was stretched in a direction parallel to the length of the string (Figs. 4B, B’). Clones overexpressing caupolican, another member of the Iro-C, had the same morphology (not shown). Interconnecting clones were also observed in the eye-antenna disc (not shown). Flies with UAS-ara expressing clones did not survive to adulthood.

When the UAS-ara overexpressing cells surrounded a small number of wild-type cells, these could undergo an apical/basal shortening similar to those of the notum iroDFM3 cells surrounded by wild-type cells (Figs. 4C, D, compare with Figs. 1E, F). Groups of wild-type cells with a surrounding fold at the interface with the overexpressing cells, like that in between Iro-C+ and Iro-C- cells, were also observed (Fig. 4E). These results again suggested that the apposition of cells containing Iro-C proteins with those devoid of them (or with only low levels of them) caused the latter to undergo apical/basal contraction and the subsequent formation of a fold. Often, strings or zones harbouring many expressing cells were adjacent to a fold or even were associated with an ectopic fold that would join with an extant fold of the disc (Figs. 4F, G and [10]). Still,
Figure 4
Some distinct features of UAS-ara overexpressing cells. UAS-ara expressing cells (green) were detected by coexpression of UAS-GFP, except in F and G where they were labelled with an anti Ara/Caup antibody. Counterstaining (in red) was Actin (B-D, F, G) or E-Cadherin (E). (A, A') UAS-ara-expressing cells distort the prospective wing margin (anti Cut staining, purple) by apparently pulling on it (arrowheads). (B, B') Apical x, y section of a connecting stripe between larger masses of UAS-ara expressing cells. Cell contours (arrowheads) are stretched parallel to the longitudinal axis of the stripe. (C, D) xy and z optical sections, respectively, of a small group of wild-type cells surrounded by UAS-ara-expressing cells. The former undergo apical-basal shortening (arrowheads). (E) z section of a group of wild-type cells surrounded by UAS-ara expressing cells. The apical marker DE-Cadherin shows that a fold has been formed at the interface (arrowheads). (F, G) xy and z optical sections, respectively, of UAS-ara expressing clones that have induced folds of the wing pouch epithelium (arrowheads) that comprise overexpressing and non-expressing cells. The folds can join with the extant folds of the disc (arrow). Dashes mark the approximate plane of the reciprocal view. For orientation, blue arrows in D, E and G point towards the apical part of cells.

Discussion
Little is known of the molecules that prevent the mixing of cells between the subregions of the wing imaginal disc. Our data argue for a role of the Iro-C homeoproteins, which accumulate in the notum region of the disc, in promoting the differential affinity of notum and wing hinge cells. This differential affinity was discovered by the sorting out during aggregation of dissociated notum and wing hinge cells [13]. We have now shown that cell sorting behaviour indeed occurs in the intact wing disc: Iro-C+ cells located within the notum, which will develop as wing hinge cells, have a tendency to join together and minimize contacts with wild-type cells. Moreover, this behaviour seems to be induced by the different levels of Iro-C homeoproteins that occur in the apposing cells, rather than by the transformation of notum cells to hinge cells, since homozygous Iro-C- cells also tend to join together when located in an heterozygous Iro-C+/ background of non-transformed cells. Moreover, wing hinge cells forced to express Iro-C by local activation of the EGFR pathway arrange themselves in roundish clones [17] suggesting a differential affinity with their non-expressing neighbours.

UAS-ara or UAS-caup overexpressing cells in a background of non-expressing cells also appear to contact one another. However, in this case the interaction between overexpressing cells is complicated by the fact that many of them arrange themselves in strings that can be as thin as a single cell and interconnect larger masses of overexpressing cells. The strings tend to surround areas with large numbers of cells not expressing Iro-C products or doing so at low levels. Conceivably, formation of the strings might be a passive process resulting from chance encounters by cells from different clones. If the overexpressing cells established strong adhesive interactions, the linkages between clones might be maintained and stretched into thin strings during disc growth. However, several observations support the alternative possibility that arrangements of the cells in strings is an active process. First, in young third instar discs bridges composed of a few cells arranged in chains that interconnect different overexpressing clones are quite abundant, while they are quite rare in control clones. Second, very long thin strings spanning many dozens of cell diameters can be observed in discs harbouring few clones. Third, pairs of clones born in the A and P compartments and whose main masses of cells are well separated from the compartment boundary can display interconnecting strings. Fourth, previous work [18] has shown that during the growth of the wing discs the cells of a clone normally remain together and usually separated from those of other clones. And fifth, control clones in the wing pouch grow mainly in a direction roughly perpendicular to the prospective wing margin [19] while strings between overexpressing clones take any
possible direction. If indeed cells from different clones actively search for one another and establish contacts, the mechanisms involved are unknown. Other observations pertaining to clones overexpressing UAS-ara, like the distortion of the D/V boundary by clones contacting it (Fig. 4A, A’), the stretching of the cells in a direction parallel to the plane of the string (Fig. 4B, B’), and the relative roundness of the domains of wild-type cells surrounded by strings, suggest that the threads of overexpressing cells exert a pulling force along its longitudinal axis. This force could be actively generated by these cells or result from a restraining action of the strings on the growth of the encircled territories of nonexpressing cells. Regardless of the mechanism, it seems most likely that the overexpressing cells display a strong adhesion between themselves. Since clones of cells with a high differential affinity normally have roundish and smooth contours to minimize cell-cell contacts along the interface of the clone, it is of interest that the contour of the UAS-ara overexpressing clones, excepting for the stretched interconnecting strings, generally appear as wiggly as that of the wild type clones (Figs. 3A–D). This suggest the presence of a polarized affinity between the overexpressing cells that permits their arrangement in strings or threads a few cells thick, but does not tend to minimize their interface with non-overexpressing cells. To our knowledge, this phenotype of clones interconnected by strings is so far unique. It appears to be difficult to explain by simple differential affinity models.

We have shown that the apposition of cells expressing and not expressing Iro-C causes the non expressing cells to undergo apical-basal shortening and invagination. Our observations suggest that this effect has only a short range and that, as cells proliferate, those that are further removed from the interface recover a normal apical-basal length. This would provide a mechanism for the formation of the fold that surrounds the older clones or of that which separates the notum and wing domains of the imaginal disc. Since both these folds appear to be formed in an approximately symmetric way, at one side by Iro-C expressing cells and at the other by non-expressing cells, the apical-basal shortening effect may gradually and actively extend to the Iro-C expressing cells close to the interface. Alternatively, these may passively accommodate to the shortening of the non expressing cells. It should be stressed that previous evidence has already disclosed non-autonomous patterning effects of Iro-C clones located in the notum region on the surrounding Iro-C+ cells [10]. Moreover, the suppression of the notum-wing hinge Iro-C border of expression negatively affects the growth of the wing disc [15]. Taken together, these lines of evidence suggest that the notum-wing hinge boundary is a source of signals that affect the growth and patterning of the surrounding tissue, an activity reminiscent of the signals that emerge from the apposition of cells at the A/P and D/V boundaries of the disc ([5], review).

The molecules responsible for the communication between Iro-C expressing and non-expressing cells are unknown. We have tested likely candidates, but the results have been negative. For instance, concerning fold formation, the role of the Rho GTPase pathway has been evaluated by producing dRhoGEF24.1 clones. dRhoGEF24.1 is activated by folded gastrulation and reiteratively required for epithelial folding and mesoderm invagination, but not for other processes regulated by Rho1 [20]. Contrary to previous evidence [20], these clones did not interfere in our hands with the notum/hinge fold, whose formation is dependent on the apposition of Iro-C expressing and non-expressing cells, or with the other extant folds of the disc. Overexpression of UAS-RhoGEF24 in clones did induce apical-basal contraction of cells. However, these clones had low viability and the apical contraction might be due to a basal extrusion of the clones from the epithelium [21,22]. Overexpression of either folded gastrulation or a dominant negative form of Rho strongly disrupted the epithelium of the disc, so no conclusions could be reached. A change in Myosin II localization is needed for the apical constriction that precedes mesodermal invagination [23]. However, Myosin II accumulation was apparently unaffected in either Iro-C or UAS-ara overexpression clones (data not shown).

The arrangement of UAS-ara expressing cells in interconnecting clones was not disturbed by reduction of the MAP kinase pathway (coexpression with UAS-ratPNI), which is active in the presumptive notum [17,24] and regulates cell adhesion [25], by the loss-of-function of DaPKC (coexpression with UAS-DaPKCDN), a protein required for apical/basal cell polarity [26], or by the reduction of function of Ephrin (coexpression with UAS-DaEphDN), a molecule involved in cell attraction/repulsion, adhesion/de-adhesion and migration in vertebrates (reviewed in [27]).

Conclusion

The genes of the Iro-C, whose activity at the proximal part of the Drosophila wing disc is essential to define this territory as the notum anlage [10], also appear to participate in keeping the prospective notum cells separate from those of the adjacent wing territory, which do not express the Iro-C genes. The available evidence suggests that the cell sorting behaviour between notum and hinge cells is induced by the different levels of Iro-C homeoproteins that occur in the apposing cells. Thus, the Iro-C homeoproteins appear instrumental in establishing not only the identity of the notum cells, but also the differential affinity of the notum versus the wing hinge cells, a property first recognized by the sorting behaviour during aggregation of dissociated notum and hinge cells [13].
These roles of the Iro-C seem similar to those of the homeobox selector genes *engrailed*/*invected* and *apterous* [2-5]. These genes are ultimately responsible, on the one hand, to impart "posterior" or "dorsal" identities to the cells of their respective expression compartments and, on the other hand, to promote the differential affinities that allow them to keep separate from the adjacent "anterior" or "ventral" cells. However, in contrast to these selector genes whose expression is inheritable maintained, this is not so for the expression of the Iro-C [10]. Hence, the border between the notum and wing hinge can be trespassed by cells when they change the state of Iro-C activation. Another difference between the borders at the anterior/posterior or dorsal/ventral compartments and at the notum/hinge territories is that only the last one is associated with a readily distinguishable morphological feature, namely, a fold of the epithelium. This fold appears to be induced by the apposition of the Iro-C expressing and non-expressing cells, which promotes an apical-basal shortening and invagination of the non-expressing cells. We propose a model that may explain how this change in cell configuration underlies formation of the notum/hinge epithelial fold.

Finally, we reexamine and extend the description of the unique and complex phenotype of disc cells that overexpress a member of the Iro-C [10]. The cells from different overexpressing clones establish contacts and become organized in a network composed of thin strings that surround and isolate large groups of non-overexpressing cells. The strings appear to exert a pulling force along their longitudinal dimension. Hitherto, we do not know of any similar phenotype.

**Methods**

**Drosophila stocks**

*Drosophila* stocks used were: Df(3L)iroDFM3 (= *iroDFM3*), UAS-ara [11]. Other stocks are described in FlyBase [28].

**Mosaic analyses**

Mitotic recombination clones homozygous for *iroDFM3* were induced by the FLP-FRT technique [29]. Flies *hsFLP1.22; mwh* *iroDFM3* FRT80B/TM6B were crossed with *hsFLP1.22; ubi-GFP* FRT80B individuals. To obtain *iroDFM3* mitotic recombination clones simultaneously with recombination clones for the *arm-lacZ* marker, *w hsFLP1.22; P[arm-lacZ] FRT40A/CyO; mwh Df(3L)iroDFM3 FRT80B/TM6B* flies were crossed with *FRT40A; ubi-GFP* FRT80B flies. Clones were induced by incubation of larvae at 37°C for 1 hour. Clones overexpressing UAS-ara were obtained by crossing flies carrying this transgene with *y w hs-FLP1.22; act-FRT* γ* FRT Gal4 UASGFP/Cyo* [30] flies. Clones were induced by incubation of larvae at 37°C for 15 minutes at 24–48 hours after egg laying (AEL) and 48–72 hours AEL.

**Antibody staining**

Imaginal discs were fixed and stained as described [29]. Primary antibodies were: rat anti-Ara, which reacts with Ara and Caup proteins [10], guinea pig anti-Echinoid provided by Jui-Chou Hsu, rat anti-DE-Cadherin (DCAD2), rabbit anti-RhoGEF2 provided by Ronald D. Vale and mouse anti-Cut(2B10) from Developmental Studies Hybridoma Bank (DSHB), mouse anti-β-galactosidase from Promega, TO-PRO-3 from Invitrogen. Secondary antibodies were from Molecular probes. Rhodamine phalloidin was from Molecular Probes. Images were collected in an LSM510 META (Zeiss) confocal microscope.

**Competing interests**

The author(s) declares that there are no competing interests.

**Authors’ contributions**

EVC and EGP carried out the experiments. EVC and JM designed the project and wrote the manuscript. All authors read and approved the final manuscript.

**Additional material**

**Additional file 1**

Apical-basal cell shortening and disposition within the notal epithelium of *iroDFM3* clones induced 72 to 96 h AEL. *iroDFM3* cells: absence of green. (A) Conventional xz view of two small *iroDFM3* clones. Red: Actin staining. In clone b, the optical section shows only three nuclei (unstained material), but at least an additional one is visible in a more basal focal plane (not shown). (B, C) Optical x section views of clones b and c. Actin strongly labels the apical (arrowheads) and basal (asterisks) regions of the cells, and more weakly cell contours. Note that the apical regions of the cells of these small clones are already recessed into the epithelium (arrows), indicating an apical-basal shortening of the cells. Red channels are also shown in white. (D) Optical z section view of a relatively large clone in a disc stained with anti Laminin alpha (Kumagai et al. 1997 FERS Lett 412, 211–216) (red or white) and anti Integrin betaPS (DS Hybridoma Bank; Brower et al. 1984 Proc Nat Acad Sci USA 81, 7485–7489) (blue) antibodies. The extracellular matrix Laminin alpha delineates the invaginated apical region of the cells of the clone (arrowhead) and the continuous basal region of the cells (arrow). The integrin betaPS staining shows that the clone is deeply sunk into the adip epithelial cell layer, e, epithelial cell layer; ad, adip epithelial cell layer; yellow arrowhead, border between epithelial and adip epithelial cells. Interestingly, levels of Integrin betaPS appear diminished in *iroDFM3* cells.

Click here for file [http://www.biomedcentral.com/content/supplementary/1471-213X-7-106-S1.jpeg](http://www.biomedcentral.com/content/supplementary/1471-213X-7-106-S1.jpeg)
Additional file 2

An epithelial fold develops during the third instar developmental stage between the notum and wing hinge territories of the wing disc. The fold is located at the distal border of the Iro-C expression domain. Green: Actin; red: ARA/Caup. (A, B) Early and mid third instar wing discs showing the epithelial fold arising at the distal border of the ara/caup expressing domain (arrowheads). (C, D) Conventional and optical z section views, respectively, of the notum (n) – hinge (h) interface of a late third instar wing disc. ara/caup-expressing cells reach to the bottom of the fold (arrowhead). Dashes mark the approximate plane of the reciprocal view. Images A and B courtesy of Ruth Diez del Corral. 

Click here for file: [http://www.biomedcentral.com/content/supplementary/1471-213X-7-106-S2.png]

Acknowledgements

We are grateful to S. Campuzano, J. Castelli-Gair, J. Culi, J.F. de Celis, J.L. Gómez-Skarmeta, M. Ruiz-Gómez, S. Sotillos and colleagues of the JM laboratory for advice on the work and constructive criticism of the manuscript; to Ruth Diez del Corral for images shown in Fig. S2; to K. Barrett, J. Culi, J.F. de Celis, J.-C. Hsu, R. Karess, F. Matsumura, M. Mlodzic, Y. Nishida, E. Wieschaus, and to R.D. Vale, Developmental Studies Hybridoma Bank, and the Bloomington stock centers for providing reagents and stocks. Predoctoral fellowship from Ministerio de Educacion y Ciencia to EGP is acknowledged. Work was supported by grants from Dirección General de Investigación Científica y Técnica (BMC2002-411, BFU2005-02888) and an institutional grant from Fundación Ramón Areces to the Centro de Biología Molecular Severo Ochoa.

References

1. García-Bellido A, Ripoll P, Morata G: Developmental compartmentalisation of the wing disc of Drosophila. Nature New Biol 1973, 245:251-253.
2. Dahmann C, Basler K: Compartment boundaries: at the edge of development. Trends Genet 1999, 15:320-326.
3. Mann RS, Morata G: The developmental and molecular biology of genes that subdivide the body of Drosophila. Annu Rev Cell Dev Biol 2000, 16:243-271.
4. Irvine KD, Rauskolb C: Boundaries in development: formation and function. Annu Rev Cell Dev Biol 2001, 17:189-214.
5. Teppu T, Göt z D, Winklbauer R: Cell sorting in animal development: signalling and adhesive mechanisms in the formation of tissue boundaries. Current Op Genet Dev 2002, 12:572-582.
6. García-Bellido A, Santamaria P: Developmental analysis of the wing disc in the mutant engrailed of Drosophila melanogaster. Genetics 1972, 72:87-104.
7. Steinberg MS, Takeichi M: Experimental specification of cell sorting, tissue spreading, and specific spatial patterning by quantitative differences in cadherin expression. Proc Natl Acad Sci USA 1994, 91:206-209.
8. Dahmann C, Basler K: Opposing transcriptional outputs of Hedgehog signalling and Engrailed control compartmental cell sorting at the Drosophila A/P boundary. Cell 2000, 100:411-422.
9. García-Bellido A, Morata G, Ripoll P: Developmental compartmentalization in the dorsal mesothoracic disc of Drosophila. Dev Biol 1976, 48:132-147.
10. Diez del Corral R, Aroca P, Gómez-Skarmeta JL, Cavodeassi F, Modolell J: The Iroquois homeodomain proteins are required to specify body wall identity in Drosophila. Genes Dev 1999, 13:1754-1761.
11. Gómez-Skarmeta JL, Diez del Corral R, de la Calle-Mustienes E, Ferrés-Marcó D, Modolell J: araucan and caupolican, two members of the novel Iroquois complex, encode homeoproteins that control proneural and vein forming genes. Cell 1996, 85:95-105.
12. McNeill H, Yang CH, Brodsky M, Ungos J, Simon MA: mirror encodes a novel PBX-class homeoprotein that functions in the definition of the dorso-ventral border of the Drosophila wing disc. Genes Dev 1997, 11:1073-1082.
13. Fausto-Sterling A, Hsieh L: In vitro culture of Drosophila imaginal disc cells: aggregation, sorting out, and differentiative abilities. Dev Biol 1987, 120:284-293.
14. Wei SY, Escudero LM, Yu F, Chang LH, Chen LY, Ho YH, Lin CM, Chou CS, Chia W, Modolell J, Hsu JC: Echinoid is a component of adherens junctions that cooperates with DE-Cadherin to mediate cell adhesion. Dev Cell 2005, 8:493-504.
15. Villa-Cuesta E, Modolell J: Mutual repression between msh and Iro-C is an essential component of the boundary between body wall and wing in Drosophila. Development 2005, 132:4087-4096.
16. Cavodeassi F, Modolell J, Gómez-Skarmeta JL: The Iroquois family of genes: from body building to neural patterning. Development 2001, 128:2847-2855.
17. Zecca M, Struhl G: Control of growth and patterning of the Drosophila wing imaginal disc by EGF-mediated signaling. Development 2002, 129:1369-1376.
18. González-Gaitán M, Capdevila MP, García-Bellido A: Cell proliferation patterns in the wing imaginal disc of Drosophila. Mech Dev 1994, 40:183-200.
19. Baena-Lopez LA, Baonza A, García-Bellido A: The orientation of cell divisions determines the shape of Drosophila organs.Curr Biol 2005, 15:1640-1644.
20. Nikolaidou KK, Barrett K: A Rho GTPase signaling pathway is used reiteratively in epithelial folding and potentially selects the outcome of Rho activation. Curr Biol 2004, 14:1822-1826.
21. Gibson MC, Perrimon N: Extrusion and death of DPP/BMP-compromised epithelial cells in the developing Drosophila wing. Science 2000, 287:1785-1789.
22. Shen J, Dahmann C: Extrusion of cells with inappropriate Dpp signaling from Drosophila wing disc epidermis. Science 2005, 307:1789-1790.
23. Dawes-Hoang RE, Parmar KM, Christiansen AE, Phelps CB, Brand A, Wieschaus EF: folded gastrulation, cell shape change and the control of myosin localization. Development 2005, 132:4165-4178.
24. Simcox AA, Grumbling G, Schnepf B, Bennington-Mathias C, Her-sperger E, Shearn A: Molecular, phenotypic, and expression analysis of vein, a gene required for growth of the Drosophila wing disc. Dev Biol 1996, 177:475-489.
25. Islam R, Kristiansen LV, Romani S, Garcia-Alonso L, Hortsch M: Activation of EGF receptor kinase by L1-mediated homophilic cell interactions. Mol Biol Cell 2004, 15:2003-2012.
26. Sotillos S, Diaz-Meco MT, Caminero E, Moscat J, Campuzano S: DaPKC-dependent phosphorylation of Crumbs is required for epithelial cell polarity in Drosophila. J Cell Biol 2004, 164:549-557.
27. Pascaule EB: Eph receptor signalling casts a wide net on cell behaviour. Nat Rev Mol Cell Biol 2005, 6:462-475.
28. FlyBase [http://flybase.org].