

RESEARCH ARTICLE

Oriented basement membrane fibrils provide a memory for F-actin planar polarization via the Dystrophin-Dystroglycan complex during tissue elongation

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ABSTRACT

How extracellular matrix contributes to tissue morphogenesis is still an open question. In the *Drosophila* ovarian follicle, it has been proposed that after Fat2-dependent planar polarization of the follicle cell basal domain, oriented basement membrane (BM) fibrils and F-actin stress fibers constrain follicle growth, promoting its axial elongation. However, the relationship between BM fibrils and stress fibers and their respective impact on elongation are unclear. We found that Dystroglycan (Dg) and Dystrophin (Dys) are involved in BM fibril deposition. Moreover, they also orient stress fibers, by acting locally and in parallel to Fat2. Importantly, Dg-Dys complex-mediated cell-autonomous control of F-actin fiber orientation relies on the preceding BM fibril deposition, indicating two distinct but interdependent functions. Thus, the Dg-Dys complex works as a crucial organizer of the epithelial basal domain, regulating both F-actin and BM. Furthermore, BM fibrils act as a persistent cue for the orientation of stress fibers that are the main effector of elongation.

KEY WORDS: Morphogenesis, Extracellular matrix, *Drosophila*, Tissue elongation, Planar cell polarity, Dystrophin, Dystroglycan, Basement membrane

INTRODUCTION

Deciphering the mechanisms underlying tissue morphogenesis is crucial for our fundamental understanding of development and also for regenerative medicine. Building organs generally requires the precise modeling of a basement membrane extracellular matrix (ECM), which in turn can influence tissue shape (Dzamba and DeSimone, 2018; Sekiguchi and Yamada, 2018; Isabella and Horne-Badovinac, 2015a,b). However, the mechanisms driving the assembly of a specific basement membrane (BM) and how this BM then feeds forward on morphogenesis are still poorly understood.

Drosophila oogenesis offers one of the best tractable examples in which such a morphogenetic process can be studied. Each ovarian follicle, which is composed of a germline cyst surrounded by the somatic follicular epithelium, undergoes a dramatic growth, associated with tissue elongation, starting from a little sphere and ending with an egg in which the anteroposterior (AP) axis is 3-fold longer than the mediolateral (ML) axis (Fig. 1A). This elongation is roughly linear from the early to the late stages, but can be separated in at least two mechanistically distinct phases (Alégot et al., 2018; Aurich and Dahmann, 2016). The first phase (from stage 3 to stage 8; hereby ‘early stages’) requires a double gradient of JAK-STAT pathway activity that emanates from each pole and that controls myosin II-dependent apical pulsations (Alégot et al., 2018). In the second phase, from stage 7-8, elongation depends on the atypical cadherin Fat2 that is part of a planar cell polarity (PCP) pathway orienting the basal domain of epithelial follicle cells (Gutzeit et al., 1991; Barlan et al., 2017; Chen et al., 2016; Viktorinová et al., 2009). Earlier during oogenesis, Fat2 gives a chirality to the basal domain cytoskeleton in the germarium, the structure from which new follicles bud (Chen et al., 2016). This chirality is required to set up a process of oriented collective cell migration perpendicularly to the elongation axis that induces follicle revolutions from stage 1 to stage 8 (Haigo and Bilder, 2011; Chen et al., 2016; Viktorinová and Dahmann, 2013). From each migrating cell, Fat2 also induces, in the rear adjacent cell, the formation of planar-polarized protrusions that are required for rotation (Cetera et al., 2014; Barlan et al., 2017). These rotations allow the polarized deposition of BM fibrils, which involves a Rab10-dependent secretion route targeted to the lateral domain of the cells. These BM fibrils are detectable from stage 4 onwards and persist until late developmental stages (Haigo and Bilder, 2011; Isabella and Horne-Badovinac, 2016). Follicle rotation also participates in the planar cell polarization of integrin-dependent basal stress fibers that are oriented perpendicularly to the AP axis (Cetera et al., 2014). Moreover, at stage 7-8, a gradient of matrix stiffness controlled by the JAK-STAT pathway and Fat2 contributes to elongation (Crest et al., 2017). Then, from stage 9, the epithelial cell basal domain undergoes anisotropic oscillations, as a result of periodic contraction of the oriented stress fibers, which also promotes follicle elongation (He et al., 2010; Qin et al., 2017). To explain the impact of *fat2* mutations on tissue elongation, it is generally accepted that oriented stress fibers and BM fibrils act as a molecular corset that constrains follicle growth in the ML axis and promotes its elongation along the AP axis. However, the exact contribution of F-actin versus BM to this corset is still unclear, as is whether the orientations of stress fibers and of BM fibrils are causally linked.

Here, we analyzed the function of Dystrophin (Dys) and Dystroglycan (Dg) during follicle elongation. Dys and Dg are the two main components of the Dystrophin-associated protein

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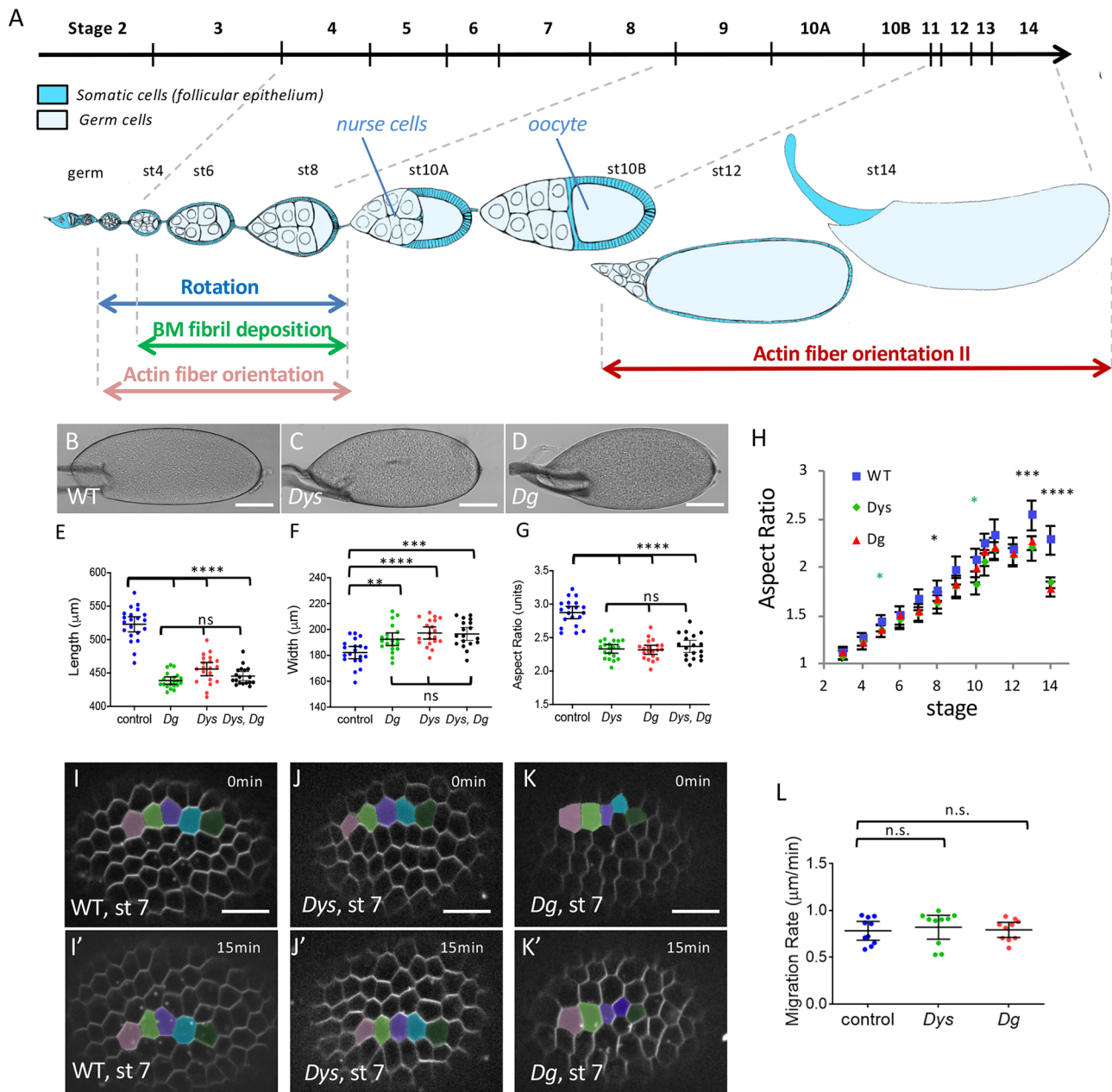


Fig. 1. The DAPC is involved in follicle elongation but not rotation. (A) Scheme of an ovariole with the main events involved in follicle elongation. The top line indicates the time scale of the different developmental stages. The ovariole is oriented anterior to posterior (germ, germarium). Each follicle is composed of a germline cyst surrounded by the follicular epithelium. Rotation occurs from very early to stage 8. It promotes F-actin fiber orientation and allows polarized BM fibril deposition, both perpendicularly to the elongation axis. At stage 11, actin fibers lose their orientation and then progressively reorient (orientation II). (B-D) Representative mature eggs from WT (B), *Dys*^{E171/Exel6184} (C) and *Dg*^{O46/O83} females (D). Scale bars: 100 μ m. (E-G) Quantification of the length (E), the width (F) and the aspect-ratio (G) for mature eggs from WT, *Dys*^{E171/Exel6184}, *Dg*^{O46/O83} mutant and *Dys*^{E171/Exel6184}, *Dg*^{O46/O83} double-mutant females ($n > 18$ eggs). (H) Elongation kinetics of WT, *Dys*^{E171/Exel6184} and *Dg*^{O46/O83} mutant follicles ($n > 6$ follicles for each point, green asterisks for *Dys*, black asterisks for *Dys* and *Dg*). There are two series of value for stage 10, corresponding to stage 10A and 10B. (I-K') Images of rotation movies of stage 7 WT (I,I'), *Dys*^{E171/Exel6184} (J,J') and *Dg*^{O46/O83} (K,K') mutant follicles. Scale bars: 10 μ m. (L) Rotation velocity of WT, *Dys*^{E171/Exel6184} and *Dg*^{O46/O83} mutant stage 7 follicles ($n > 8$ follicles). For all panels, error bars represent s.d.; * $P < 0.01$, ** $P < 0.005$, *** $P < 0.001$, **** $P < 0.0001$ (unpaired *t*-test). n.s., not significant.

complex (DAPC), an evolutionarily conserved transmembrane complex that links the ECM (via Dg) to the F-actin cytoskeleton (via Dys) (Barresi and Campbell, 2006). This complex is expressed in a large variety of tissues and is implicated in many congenital degenerative disorders. Loss-of-function studies in model organisms have revealed an important morphogenetic role for Dg during development, usually linked to defects in ECM secretion, assembly or remodeling (Clements et al., 2017; Henry and

Campbell, 1998; Satz et al., 2010; Bello et al., 2008; Buisson et al., 2014; Naegeli et al., 2017; Yatsenko and Shcherbata, 2014). A developmental role for Dys is less clear, possibly because of the existence of several paralogs in vertebrates. As *Drosophila* has only one *Dg* and one *Dys* gene, it is a promising model for their functional study during development and morphogenesis.

We found that *Dg* and *Dys* are required for follicle elongation and proper BM fibril formation early in fly oogenesis. During these

early stages, DAPC loss and hypomorphic *fat2* conditions similarly delay stress fiber orientation. However, DAPC promotes this alignment more locally than Fat2. Moreover, DAPC genetically interacts with *fat2* in different tissues, suggesting that they belong to a common morphogenetic network. Later in oogenesis, *Dg* and *Dys* are required for stress fiber orientation in a cell-autonomous manner. This is the period when the main elongation defect is seen in these mutants, arguing for a more determinant role for stress fibers compared with BM fibrils in the elongation process. Nonetheless, this latter function depends on the earlier DAPC function in BM fibril deposition. We propose that BM fibrils serve as a PCP memory for the late stages that are used as a template by the DAPC for F-actin stress fiber alignment.

RESULTS

Dys and *Dg* mutants reveal a new phase of follicle elongation

As *Drosophila* follicle elongation requires many proteins that function at the basal side of the follicular epithelium, we checked whether the DAPC could be involved in this process. Both *Drosophila Dg* and *Dys* null mutants are viable and we used combinations of two different null alleles for each of them (Christoforou et al., 2008). Mature eggs from mutant females are significantly shorter and wider (Fig. 1B-F), with an aspect ratio (AR) around 2.3 compared with 2.9 in wild-type (WT) follicles (Fig. 1G). Importantly, such a phenotype can be rescued by restoring expression in the follicle cells, confirming that it is associated with these mutations, and indicating that the DAPC is required for elongation in the follicle cells (see below). Moreover, double-mutant eggs for *Dys* and *Dg* show the same defect as single mutants, confirming that these genes work together for follicle elongation (Fig. 1E-G). Comparison of the elongation kinetics in mutant and WT follicles shows that such morphological differences appear mainly during the later stages of oogenesis (stages 12 to 14), although a slight, but significant defect is present around stage 8 (Fig. 1H). The *Dg* and *Dys* null mutants' elongation kinetic curves are clearly different from those of *fat2* mutant in which elongation is strongly affected from stages 7-8 (Alégot et al., 2018; Aurich and Dahmann, 2016). Thus, the DAPC is involved in follicle morphogenesis, and its specific temporal requirement suggests the existence of an elongation phase that is mechanistically different from the Fat2-dependent one.

Next, to determine how the DAPC contributes to follicle elongation, we first checked the collective cell migration that occurs during the early stages because it is upstream of many subsequent events that are potentially important for elongation. Both *Dys* and *Dg* null mutant follicles rotate and the migration velocity is not significantly different from that of WT follicles (Fig. 1I-L). Thus, the DAPC is not involved in the collective cell migration and in the initial PCP established by Fat2, confirming that it acts through a distinct mechanism. We then focused our phenotypic analysis on the two events induced by follicle rotation: BM fibril deposition and stress fiber orientation.

The DAPC is important for BM fibril deposition

The follicle cell BM contains two ECM types: a planar BM without a specific organization visible by confocal microscopy, and BM fibrils that are oriented perpendicular to the follicle AP axis. BM fibrils are deposited during follicle rotation via a secretory route targeted to the cell lateral domain, and can be observed by monitoring the expression of the main ECM components: Collagen IV (ColIV), Perlecan (Pcan; also known as Trol) and Laminin A (LanA), the two latter being direct *Dg* ligands (Gutzeit et al., 1991; Haigo and Bilder, 2011; Isabella and Horne-Badovinac, 2016). ColIV and Pcan can be visualized using the

Viking- and Trol-GFP protein traps, respectively (Morin et al., 2001). Moreover, a fosmid containing the whole *Laminin A* gene tagged with GFP has also been generated (LanA-GFP) (Sarov et al., 2016). We also employed the MiMIC system to generate RFP-tagged Pcan (Venken et al., 2011). This line gives an identical pattern to Pcan-GFP and is homozygous viable and fertile, indicating that it does not disrupt Pcan function (Fig. S1A). Comparison of the BM fibril composition using these different tagged proteins (Fig. 2A-B'') shows that in WT follicles, Pcan-RFP colocalizes exactly with ColIV-GFP and LanA-GFP. This indicates that BM fibrils are uniformly composed of these three proteins, as has been previously shown by immunostaining (Isabella and Horne-Badovinac, 2016).

We then wondered whether BM fibril deposition is affected in *Dys* and *Dg* mutants. In these two mutants, the general targeting of BM secretion towards the basal domain is normal, with no visible apical secretion as has been observed in different mutants affecting BM secretion (Fig. S1B-D) (Denef et al., 2008; Devergne et al., 2014, 2017; Lerner et al., 2013). However, images in the focal plane of the BM reveal a very strong decrease of the ColIV⁺ fibrils in the *Dys* null mutant (Fig. 2D). This observation is also true for LanA and Pcan (Fig. 2C,E). Quantification of the BM fibril fraction on stage 8 follicles, which corresponds to the fluorescence fraction present in the fibrils, indicates that it is about 15% in the WT situation, but drops to 2% in *Dys* mutants, and that a similar phenotype is observed in *Dg* mutants (Fig. 2F). This defect is specific to the fibrils and does not affect unpolarized matrix because the total basal fluorescence intensity tends to increase in these mutants, although it is variable from one follicle to another (Fig. S1E). Loss of the DAPC affects both BM fibril number and length (Fig. 2G,H). As a result of this defect, some fibrils are still visible in stage 12 WT follicles but hardly detectable in *Dys* and *Dg* mutant ones (Fig. S1F-I). This shows that the DAPC is involved in BM fibril deposition.

This deposition depends on the lateral secretion of ECM proteins while the rotation facilitates both the deposition and orientation of the fibrils in the BM (Isabella and Horne-Badovinac, 2016). The lateral secretion of BM proteins can be visualized by immunostaining on non-permeabilized follicle cells to detect only the proteins localized between the cells. Such an approach on *Dys* mutant clones shows that some ColIV is detected, as observed between WT cells (Fig. 2I-J). This suggests that *Dys* does not directly interfere with the lateral targeting of BM fibril secretion. In a mutant blocking rotation, such as *fat2*, BM proteins accumulate in the lateral space between cells, whereas no massive accumulation is observed in a *Dys* mutant, suggesting that the BM fibril defect associated with DAPC loss of function cannot be explained by a failure of the latest step of BM fibril deposition (Fig. 2K) (Isabella and Horne-Badovinac, 2016). Moreover, in a *fat2*, *Dys* double mutant we observed the same lateral accumulation of ECM proteins as in the *fat2* mutant, confirming that lateral secretion is not affected by DAPC loss of function (Fig. 2K,L). It is therefore unclear by which mechanism the DAPC contributes to BM fibril formation. Nonetheless, these mutants are the first examples of mutations that specifically reduce BM fibril formation without any defect in the general targeting of basal secretion or in rotation, offering a unique opportunity to address the functional relevance of the fibrils.

The DAPC contributes to stress fiber orientation during early stages

In parallel with the analysis of BM fibrils, we also studied the impact of the loss of the DAPC on basal F-actin organization. This alignment is still weak at stage 4 and then increases to reach a maximum around stage 8 (Fig. 3A-F) (Cetera et al., 2014). In both

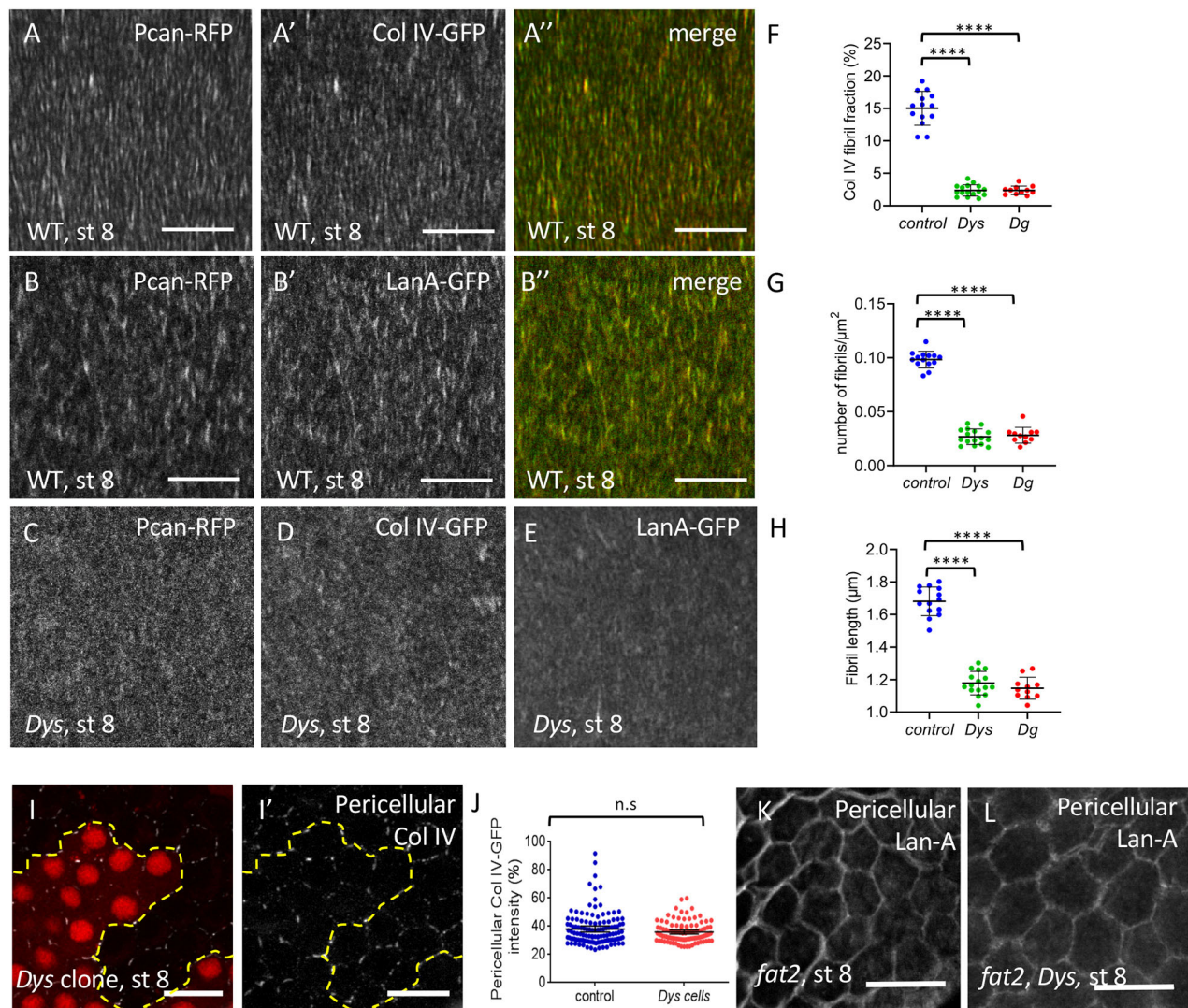


Fig. 2. The DAPC is important for BM fibril deposition. (A–B'') Basal view of WT (w^{1118}) follicle ECM at stage 8 visualized with Pcan-RFP and ColIV-GFP (A) or Pcan-RFP and LanA-GFP (B). (C–E) Basal view of *Dys*^{E17/Exel6184} follicles expressing Pcan-RFP (C), ColIV-GFP (D) and LanA-GFP (E). (F–H) Quantification of the BM fibril fraction (F), fibril numbers (G) and fibril length (H) at stage 8 in WT, *Dys*^{E17/Exel6184} and *Dg*^{O86/O43} mutant follicles ($n > 10$ follicles). (I, I') Mutant clones for *Dys*^{Exel6184} marked by the absence of RFP and stained to detect pericellular ColIV. Dashed line indicates the border between WT and mutant cells. (J) Quantification of pericellular ColIV intensity around WT and *Dys* mutant cells ($n > 20$ cells). (K, L) Accumulation of pericellular LanA in *fat2* (K) and *fat2*, *Dys* (L) mutant follicles. For all panels, error bars represent s.d.; **** $P < 0.0001$ (unpaired t -test). n.s., not significant. Scale bars: 10 μ m.

Dys and *Dg* null mutants, although the stress fibers are globally oriented perpendicular to the AP axis in mutant follicles, quantification of the angular distribution reveals a delay in their proper orientation at stage 4 and 6 to reach a normal level at stage 8 (Fig. 3A–R). Stress fiber orientation can be also analyzed by the determination of the tissue order parameter (Tissue OP) based on a method previously described (Cetera et al., 2014). A complete random orientation Tissue OP value will tend to 0, whereas a perfect alignment between the cells will tend to 1 (Fig. 3S). This method confirms that *Dys* and *Dg* mutants behave similarly, with a delay in the proper stress fiber alignment (Fig. 3T). Although rotation is required for stress fiber orientation, this motion is not affected by DAPC loss of function. Thus, this complex likely acts downstream of or in parallel with rotation to orient stress fibers.

We also addressed whether *Dys* is required cell-autonomously to promote stress fiber orientation by generating mutant clones and we focused on stage 5, when the defect is obvious in *Dys* mutant follicles. Comparing the mutant cells versus the WT ones in a same

follicle, we observed no apparent effect on F-actin density but a cell-autonomous effect on their orientation that can be quantified by the Tissue OP (Fig. 3U–V). Thus, *Dys* promotes stress fiber orientation in each cell.

The DAPC and *Fat2* have partially redundant functions

We found that the DAPC has only a minor impact on the early phase of stress fiber orientation, with a delayed orientation of actin fibers that is achieved by stage 8, suggesting alternative mechanisms to orient fibers at these stages (4–8). Similarly, deletion of the intracellular part of *Fat2* (*fat2-DeltaICD*) induces a hypomorphic defect with only a slight delay of stress fiber orientation, despite a slower rotation (Aurich and Dahmann, 2016; Barlan et al., 2017; Chen et al., 2017). We therefore wondered what could be the relationship between these two genetic backgrounds. We generated a double-mutant background for *fat2/fat2-DeltaICD* and *Dys* null alleles (*Dys*, *fat2-DeltaICD/Dys*, *fat2*) and compared stress fiber alignment for these different genotypes (Fig. 4A–D). Tissue OP

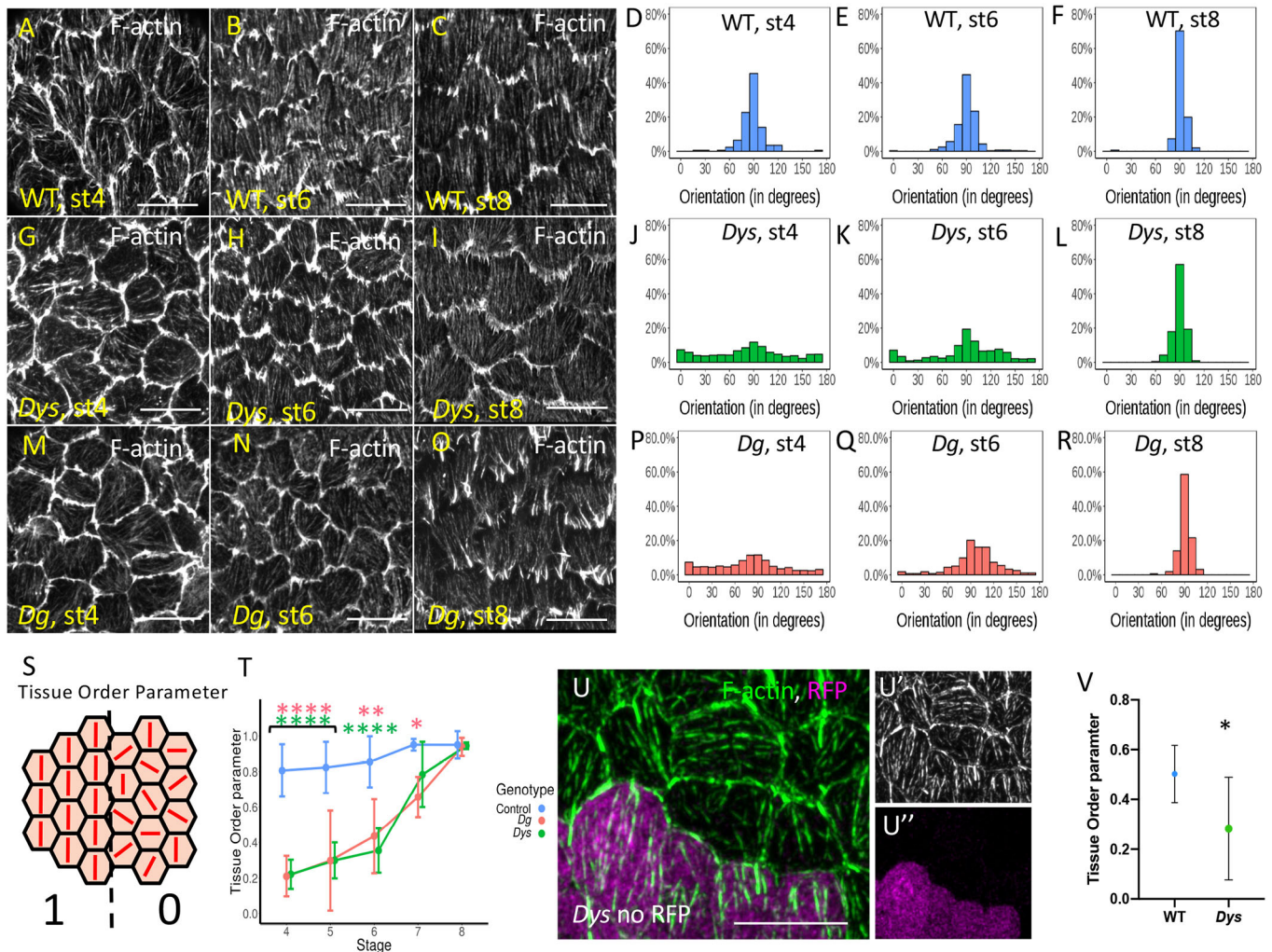


Fig. 3. Stress fiber orientation delay in DAPC mutants during early stages. (A–R) Representative images of basal F-actin in WT (w^{1118}) (A–C), $Dys^{E17/Exel6184}$ (G–I) and $Dg^{O46/O43}$ (M–O) follicles at stage 4, 6 and 8 and quantification of the corresponding angular distribution (D–F, J–L, P–R). (S) Illustration of Tissue OP: if all the cells have the same orientation Tissue OP is equal to one, if orientation is random Tissue OP is equal to zero. (T) Tissue OP analysis of control, $Dys^{E17/Exel6184}$ and $Dg^{O46/O43}$ follicles. A two-way ANOVA was performed, followed by applicable post-hoc pairwise comparisons (Tukey). (U–U'') F-actin in a mutant clone for $Dys^{Exel6184}$ marked by the absence of RFP (purple) at stage 5. Note the cell-autonomous effect on stress fiber orientation. (V) Tissue OP quantification $Dys^{Exel6184}$ mutant clones and surrounding WT cells at stage 5. Error bars represent s.d. ($n > 7$ follicles for each stage and each genotype). * $P < 0.01$, ** $P < 0.005$, **** $P < 0.0001$. Scale bars: 10 μ m.

analysis shows that single mutants are similar and that the double mutants display an additive effect (Fig. 4E,I), an additivity that is also observed on follicle elongation (Fig. 4Q). Thus, *Dys* and *Fat2* probably work in parallel for stress fiber orientation once *Fat2* has initiated follicle rotation.

To get a better understanding of this additivity, we performed a deeper analysis of stress fiber orientation. First, we wondered whether the alignment robustness was the same in each cell and we therefore analyzed the cell order parameter (Cell OP) (Fig. 4F). This clearly shows that the mutants behave in a different manner, with *fat2-DeltaICD* behaving similarly to WT and with *Dys* and *Dys, fat2-DeltaICD* giving an identical weaker Cell OP (Fig. 4J). This effect on Cell OP of *Dys* can be confirmed by its analysis on *Dys* mutant clones (Fig. S2E). Moreover, a close observation of *Dys* single cells reveals more divergent stress fibers than we have observed in *fat2-DeltaICD* cells (Fig. S2C,D). This indicates that *Dys* promotes stress fiber alignment in each cell, a process that does not seem to involve *Fat2*. We also calculated a local order parameter (Local OP), determined for each cell on a cluster including this cell and its neighbors within a

given radius (Fig. 4G). Globally, for all mutant genotypes that we considered, Local OP values are initially higher than the respective Tissue OP ones, but they tend to be equal to the Tissue OP values when the analyzed radius is increased (Fig. S2F). The additivity of *Dys* and *fat2* can be also observed on the Local OP (Fig. 4K, Fig. S2F). Finally, we looked at the difference between Local OP, for a radius corresponding to one row of neighbors, and Tissue OP (Fig. 4H). Interestingly, we noticed a stronger Local/Tissue OP difference for *fat2-DeltaICD* than for *Dys* mutants, a difference culminating at stage 6, indicating that stress fiber orientation is less affected at a small scale in *fat2* than in *Dys* mutants (Fig. 4L, Fig. S2G). This difference can be also directly observed on images of F-actin orientation (Fig. S2A,B). For *fat2-DeltaICD*, several patches of neighboring cells with the same orientation but changing from one patch to another can be observed. For *Dys*, we do not detect these patches but do see changes in orientation from one cell to another. Altogether, these data indicate that *Dys* promotes stress fiber planar polarization at a cellular and local scale whereas *Fat2*, or at least its intracellular domain, acts at a more global scale.

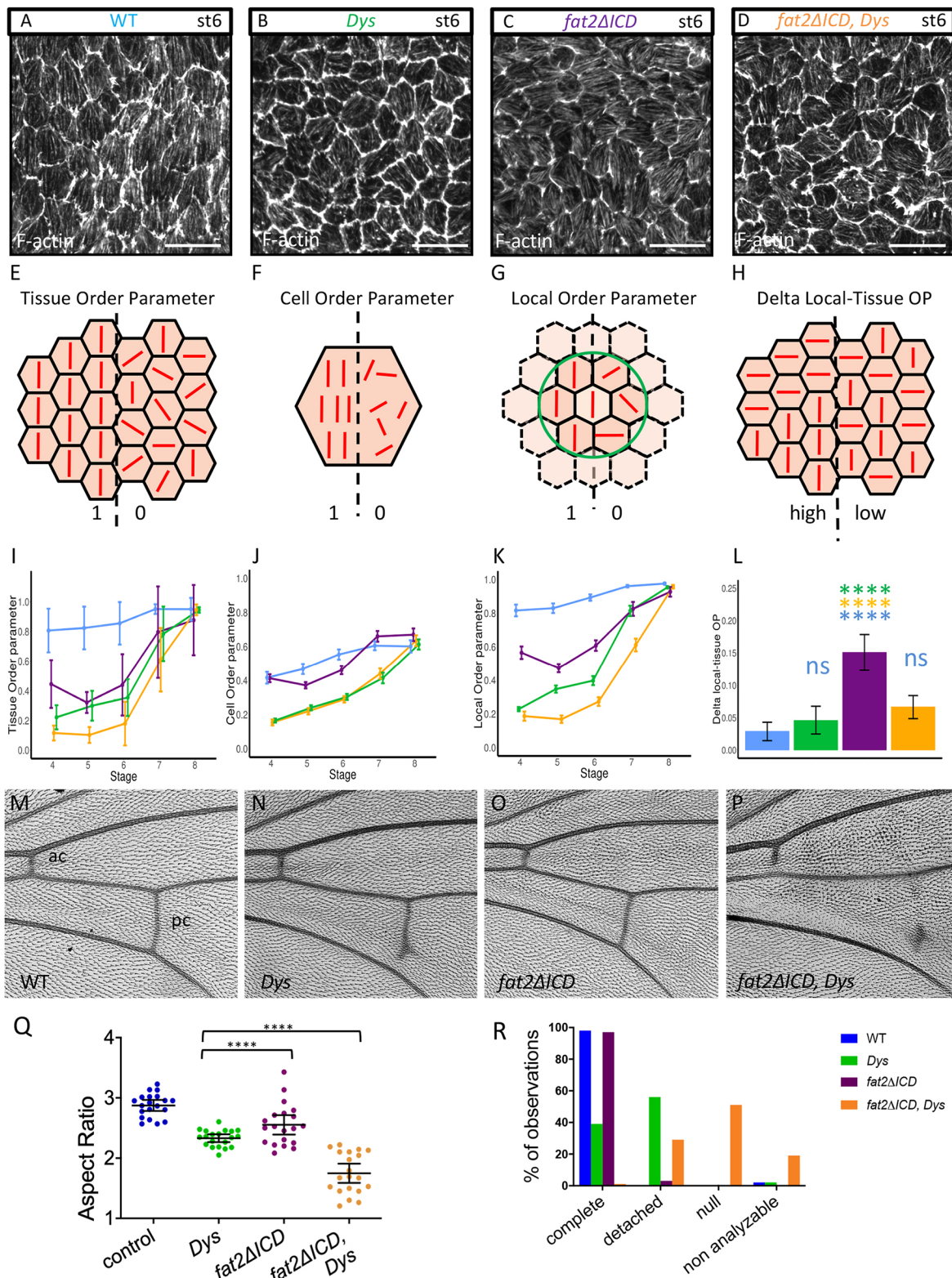


Fig. 4. Genetic interaction between *fat2* and *Dys*. (A-D) Representative images of stress fibers at stage 6 of WT (A), *Dys*^{E171/Exel6184} mutant (B), *fat2*-*Delta**ICD* mutant (C) and *Dys*, *fat2*-*Delta**ICD* double mutant (D). Genotype color code is conserved on all panels. (E-H) Schemes explaining the different parameters that are analyzed in I-L: (E,I) Tissue OP ($n > 7$ follicles for each stage and each genotype); (F,J) Cell OP; (G,K) Local OP; (H,L) difference between Local and Tissue OPs ($n > 171$ cells for each stage and each genotype). (I-L) A two-way ANOVA was performed, followed by applicable post-hoc pairwise comparisons (Tukey). *P*-values between the different genotypes for I,J,K are indicated in Table S3. (M-P) Representative images of wing veins of WT (M), *Dys*^{E171/Exel6184} mutant (N), *fat2*-*Delta**ICD* mutant (O) and *Dys*, *fat2*-*Delta**ICD* double mutant (P). ac, anterior crossvein; pc, posterior crossvein. (Q) Quantification of egg elongation for the indicated genotypes ($n > 20$ eggs for each genotype, unpaired *t*-test). (R) Quantification of wing defects observed in the different genotypes on the pc. 'Non analyzable' corresponds to wrinkled or misfolded wings for which crossveins cannot be observed ($n = 40$ wings). For all panels, error bars represent s.d.; **** $P < 0.0001$. Scale bars: 10 μ m.

These additive effects revealed by these experiments indicate that the DAPC and *fat2* act partially redundantly to orient stress fibers during early stages. They further suggest that Fat2 and the DAPC are part of a same functional network during follicle elongation, although they probably work in parallel once BM fibrils are deposited.

Intriguingly, *Dys*, *Dg* and *fat2* mutants show a very specific defect in the wing, with a partial absence of the posterior crossvein, although the cellular basis for this defect is unknown (Christoforou et al., 2008; Viktorinová et al., 2009) (Fig. 4M,N,R). However, this defect is not observed in a *fat2/fat2-DeltaICD* hypomorphic background (Fig. 4O,R). In the double-mutant *Dys, fat2-DeltaICD* flies, the posterior crossvein is usually completely absent or limited to a dot, and the anterior crossvein, which is never affected in *Dys* mutant flies, is incomplete (Fig. 4P,R). These defects indicate a strong genetic interaction in the wings, as observed in follicles.

Thus, these results tend to confirm that Fat2 and the DAPC work in the same network, which might be involved in many contexts.

The DAPC is required for stress fiber orientation during late stages

Elongation curves of *Dys* and *Dg* mutants indicate that the major elongation defect appears after stage 12 (Fig. 1H). Of note, for unknown reasons, stress fibers lose their planar polarization at stage 10B/11 but regain their proper orientation around stage 12 (Delon and Brown, 2009; Wahlström et al., 2006) (Fig. 5A,B). In WT follicles, this transition can be observed by the change in angular distribution between stage 11 and stage 13 (Fig. 5C,D). F-actin staining of *Dys* and *Dg* null mutant follicles indicates that the stress fibers lose their orientation at stage 11 as in WT, but fail to regain their proper orientation at stage 13 (Fig. 5E,F,I,J). Angular distribution quantification even indicates a tendency of the stress fibers to be

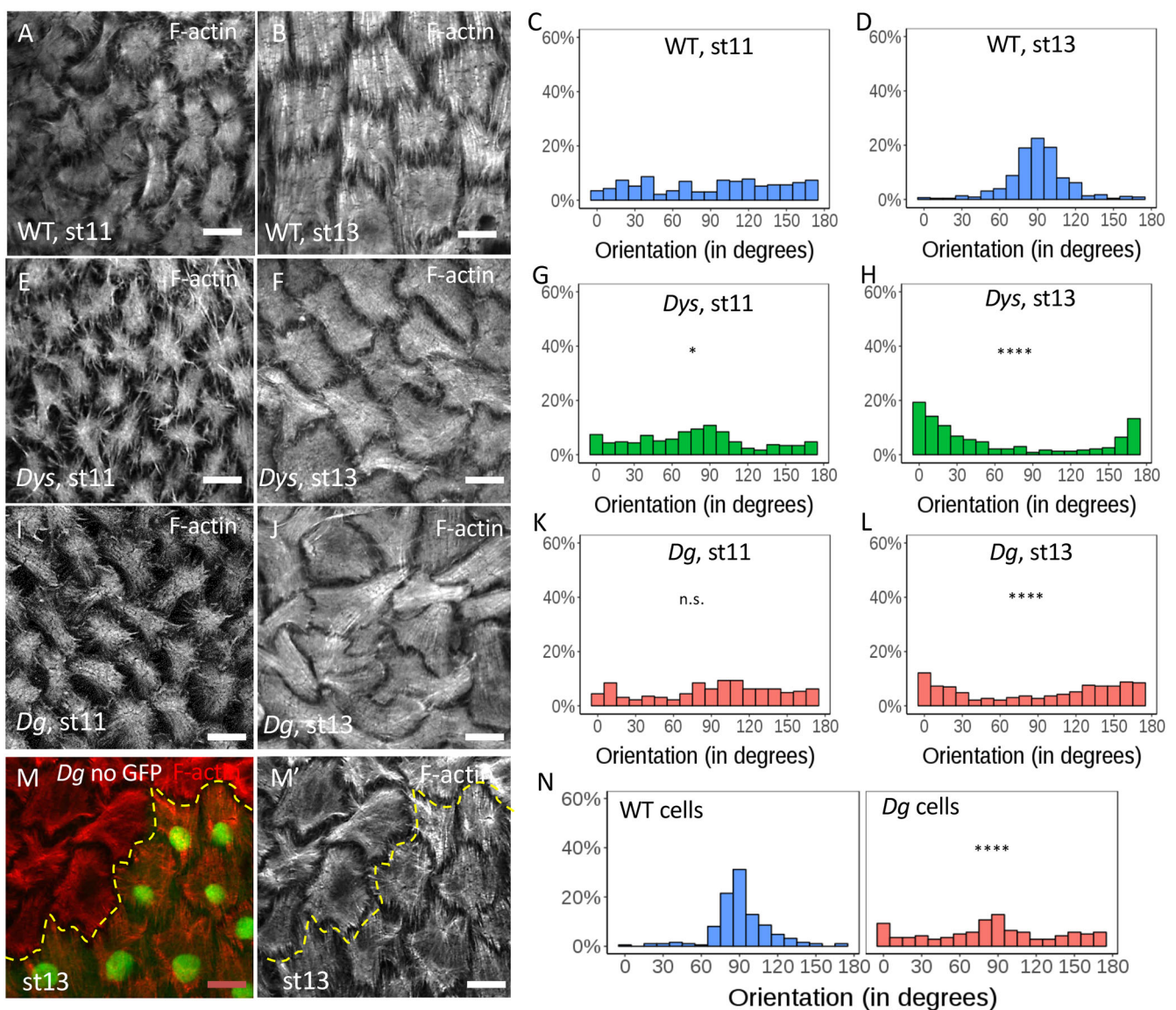


Fig. 5. Stress fiber orientation defect in DAPC mutants during late stages. (A-L) Representative images of basal F-actin in WT (A,B), *Dys*^{E17/Exe6184} (E,F) and *Dg*^{O86/O43} (I,J) follicles at stages 11 and 13 and quantification of the corresponding angular distribution (C,D,G,H,K,L) ($n > 5$ follicles). (M-N) Representative image of basal F-actin in *Dg* mutant clone marked by the absence of GFP (M,M') and quantification of the corresponding angular distribution of stress fibers in the mutant cells and the neighboring WT cells (N) ($n > 5$ clones). Angular distribution was compared with that of WT at same stage (and statistically analyzed as described in the Materials and Methods); $*P < 0.01$, $****P < 0.0001$. n.s., not significant. Scale bars: 10 μ m.

aligned parallel to the AP axis at stage 13 (Fig. 5G,H,K,L). Despite their misorientation, these fibers appear normal in structure with, for instance, a correct localization of integrins at their extremities (Fig. S3A-C). Importantly, mitotic recombination to generate mosaic tissues indicates that this function is cell-autonomous for *Dg* with misaligned stress fibers only in the mutant cells (Fig. 5M-N), as previously observed in *Dys* clones in early stages. The defect is strictly cell-autonomous as it is also observed in mutant cells that are adjacent to WT cells, whereas WT cells that are adjacent to mutant cells have stress fibers properly oriented (Fig. 5M). Importantly, this cannot be a consequence of a primary effect on BM fibril deposition because follicle rotation has moved the position of the clone with respect to the BM. It reveals, therefore, another function of the DAPC during follicle development: given the known link of the DAPC with F-actin, it suggests a direct role in stress fiber orientation, sensing and transmitting a cue defining this planar polarization.

Late DAPC function relies on both early and late activities

Because *Dys* and *Dg* mutants affect BM fibril deposition and stress fiber orientation, we aimed to determine which of these defects is the main contributor to the follicle elongation phenotype. To address this question, we took advantage of the temporality of the DAPC function during oogenesis, being required for BM fibril deposition during early stages and for stress fiber orientation mainly during later stages. Importantly, these three defects – follicle elongation, BM fibril deposition and stress fiber orientation – are recapitulated when RNAi against *Dg* is induced in follicle cells during all stages using a *traffic jam:Gal4* (*tj:Gal4*) driver, expressed in follicle cells throughout oogenesis (Fig. 6F-I). We also used *Cy2:Gal4*, which is only expressed in follicle cells from stage 9 (Fig. S4N). The expression of *Dg* RNAi with this driver knocks it down after the correct BM fibril deposition (Fig. S4A,B), but before the second phase of stress fiber orientation. As expected, the stress fiber orientation is disrupted at stage 13, confirming that this DAPC function is distinct from the one influencing the BM (Fig. S4C,D). Importantly, egg elongation is also affected (Fig. S4E). Thus, BM fibrils are not sufficient to drive late follicle elongation, and stress fiber orientation is necessary for it.

We also performed the reverse experiment by trying to rescue the late stress fiber orientation defect in a *Dg* mutant using a UAS:*Dg* transgene. Expression of *Dg* during the whole oogenesis in the follicle cells, using the *tj:Gal4* driver in a mutant background, rescues the elongation defect, the BM fibril deposition and stress fiber orientation (Fig. S4F,H,I,J,M). We then tried to rescue the *Dg* mutant using the *Cy2:Gal4* driver, which clearly allows *Dg* detection from stage 9 with an antibody failing to detect endogenous level of the protein (Fig. S4N-P'). As expected, such follicles show BM defects associated with *Dg* mutation (Fig. S6G,H). However, surprisingly, this genetic combination also fails to rescue stress fiber orientation (Fig. S4K,L). Moreover, egg elongation is only weakly rescued (Fig. S4M). Together, these results indicate that late expression of *Dg* is not sufficient to restore its late functions, and, thus, that these late functions also rely on a previous role of *Dg* during oogenesis.

To confirm these important observations, we set up an alternative method to control the timing of *Dg* inhibition by RNAi, using *tj:Gal4* and *Tub:Gal80^{ts}*, which is a thermosensitive *Gal4* inhibitor allowing a temporal control of its activity. We defined two conditions allowing either *Gal4* activity during early stages (switch from 30 to 18°C, 'Early *Dg* RNAi') or late stages (switch from 18 to 30°C, 'Late *Dg* RNAi') (Fig. 6A). Moreover, positive and negative controls were performed at 18°C and 30°C (Fig. 6). For

each condition, an internal negative control (driver line) with the same temperature was performed. *Dg* RNAi at 18°C is similar to control (Fig. 6B-E) whereas at 30°C it recapitulates all the defects observed in *Dg* mutants, as previously mentioned (Fig. 6F-I). The best timing for the temperature switches was adjusted by checking their effect on fibril in late stages, taking into account that these fibrils are formed during early stages. Thus, fibrils are normal in 18–30°C conditions and affected in 30–18°C conditions (Fig. 6J,N). We observed that F-actin orientation at late stages and follicle elongation are affected in both late (Fig. 6K-M) and early (Fig. 6O-Q) *Dg* RNAi conditions, confirming our previous results. Thus, the DAPC is required at two different time windows and its second function is dependent on the first one.

BM fibril deposition provides a persistent cue for F-actin orientation

Overexpression of *Rab10* increases the BM fibril fraction and enhances follicle elongation, a phenotype opposite to the loss of function of the DAPC (Isabella and Horne-Badovinac, 2016). We therefore performed an epistasis test between these two conditions. Concomitant overexpression of *Rab10* and knockdown of *Dys* in follicle cells using *tj:Gal4* increases the amount of BM fibrils to a higher level than that of the WT control and thus fully compensates for the absence of *Dys* (Fig. S5A-D,E). This rescue is not due to titration of the *Gal4* protein by the second UAS line when we used both *Dys* RNAi and UAS:*Rab10* constructs as the BM fibril fraction in a UAS:tomato; *Dys* RNAi control is not increased (Fig. S5I). Classical interpretation of such a test would indicate that *Rab10* acts downstream (or in parallel) of the DAPC. These experiments were performed at optimal temperature for *Gal4*-induced expression (30°C). However, in this context *Rab10* overexpression disrupts follicle elongation (Isabella and Horne-Badovinac, 2016), and it is also associated with the abnormal targeting of *ColIV*-containing vesicles to the apical side of the cells (Fig. S5H). Therefore, these conditions did not allow us to determine whether it could also rescue follicle elongation. As a consequence, we repeated these experiments at 25°C and observed an almost complete rescue of BM fibril fraction (Fig. 7A). However, mature egg elongation was still disrupted, as in DAPC loss of function (Fig. 7B). Thus, for elongation, *Dys* should be placed downstream of *Rab10*. Taken together, these results are consistent with the proposal that the DAPC has two separate functions, one upstream and one downstream of *Rab10*. Moreover, it also confirms that, in absence of the DAPC, BM fibrils are not sufficient for elongation.

We also analyzed stress fiber orientation in late stages in this context. Angular distribution for *Dys* RNAi (Fig. S5J) compared with that of null mutants (Fig. 5H), indicates that it corresponds to hypomorphic conditions. This result prompted us to also analyze the different order parameters, as previously done at early stages, to get a more precise evaluation of stress fiber orientation. The three parameters are strongly affected in *Dys* RNAi (Fig. 7C-E). When *Rab10* is overexpressed at the same time we observed a partial rescue of this defect on the angular distribution, the Tissue OP and the Local OP but not the Cell OP (Fig. S5J, Fig. 7C-E,I). These observations could be explained by the rescue of the early function of the DAPC by *Rab10* and the hypomorphic nature of *Dys* RNAi. Nonetheless, this approach offers a unique opportunity to test whether the DAPC's first function in BM fibril deposition is required for its later function in stress fiber orientation. We repeated the experiments but allowing *Dys* expression during late stages by a temperature switch (25°C to 18°C). Strikingly, the late expression of *Dys* promotes correct stress fiber orientation on all the parameters

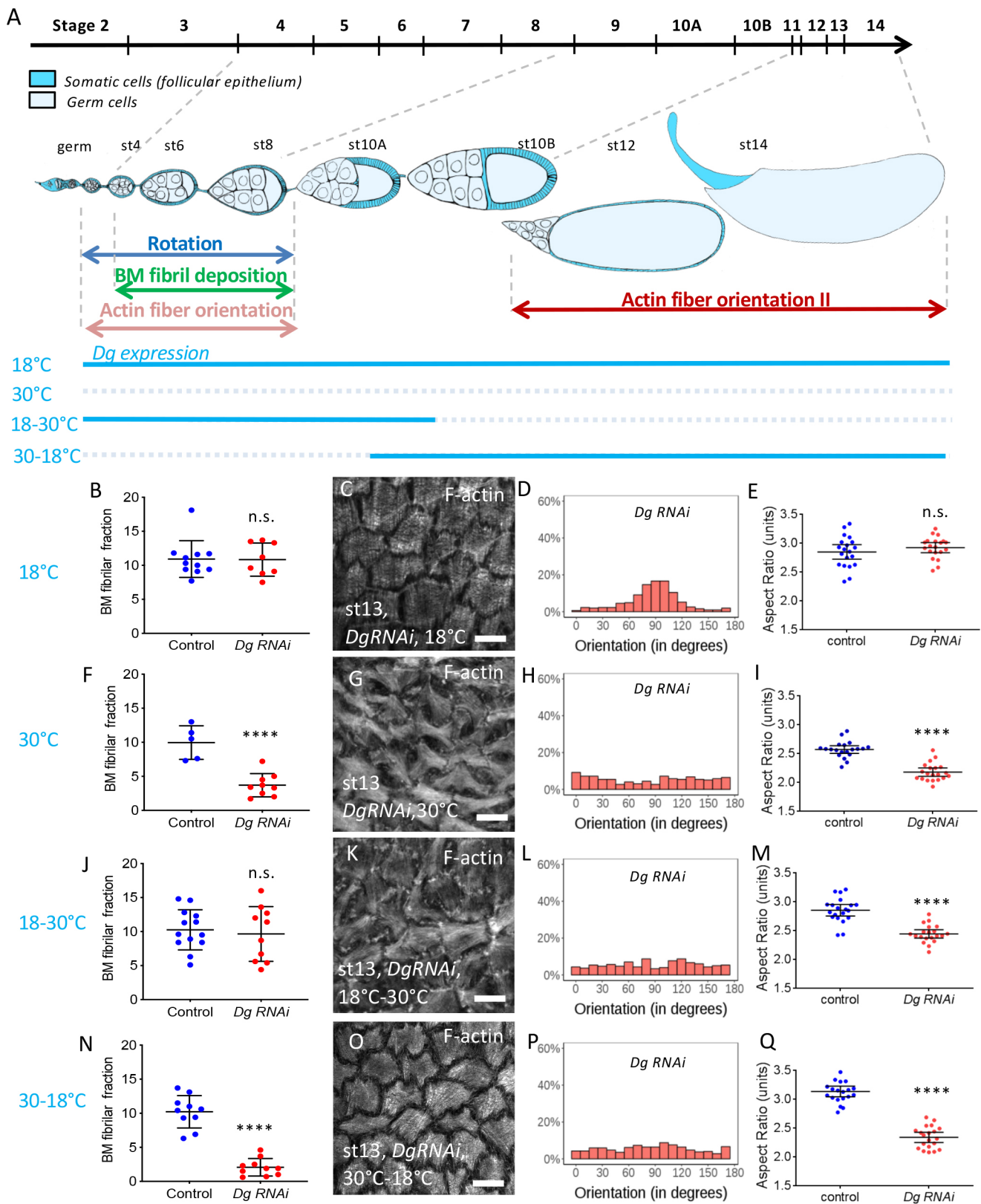


Fig. 6. The DAPC is required at two time frames for follicle elongation. (A) Scheme of an ovariole with the main events involved in follicle elongation. The top line indicates the time scale of the different developmental stages. The ovariole is oriented anterior to posterior (germ, germarium). The level of endogenous Dg is temporally controlled by the temperature (blue lines). (B, F, J, N) Quantification of the BM fibril fraction of *Dg RNAi* stage 12 follicles expressing ColIV-GFP in the indicated condition compared with control performed in the same temperature condition ($n > 5$ follicles). (C, D, G, H, K, L, O, P) Representative images of basal F-actin of *Dg RNAi* stage 13 follicles in the indicated condition (C, G, K, O) and quantification of the corresponding angular distribution (D, H, L, P) ($n > 5$ follicles). (E, I, M, Q) Aspect-ratio quantification of temporally controlled *Dg RNAi* mature compared with control performed in the same temperature condition ($n > 20$ eggs). For all panels, error bars represent s.d.; **** $P < 0.0001$ (unpaired t -test). n.s., not significant. Scale bars: 10 μ m.

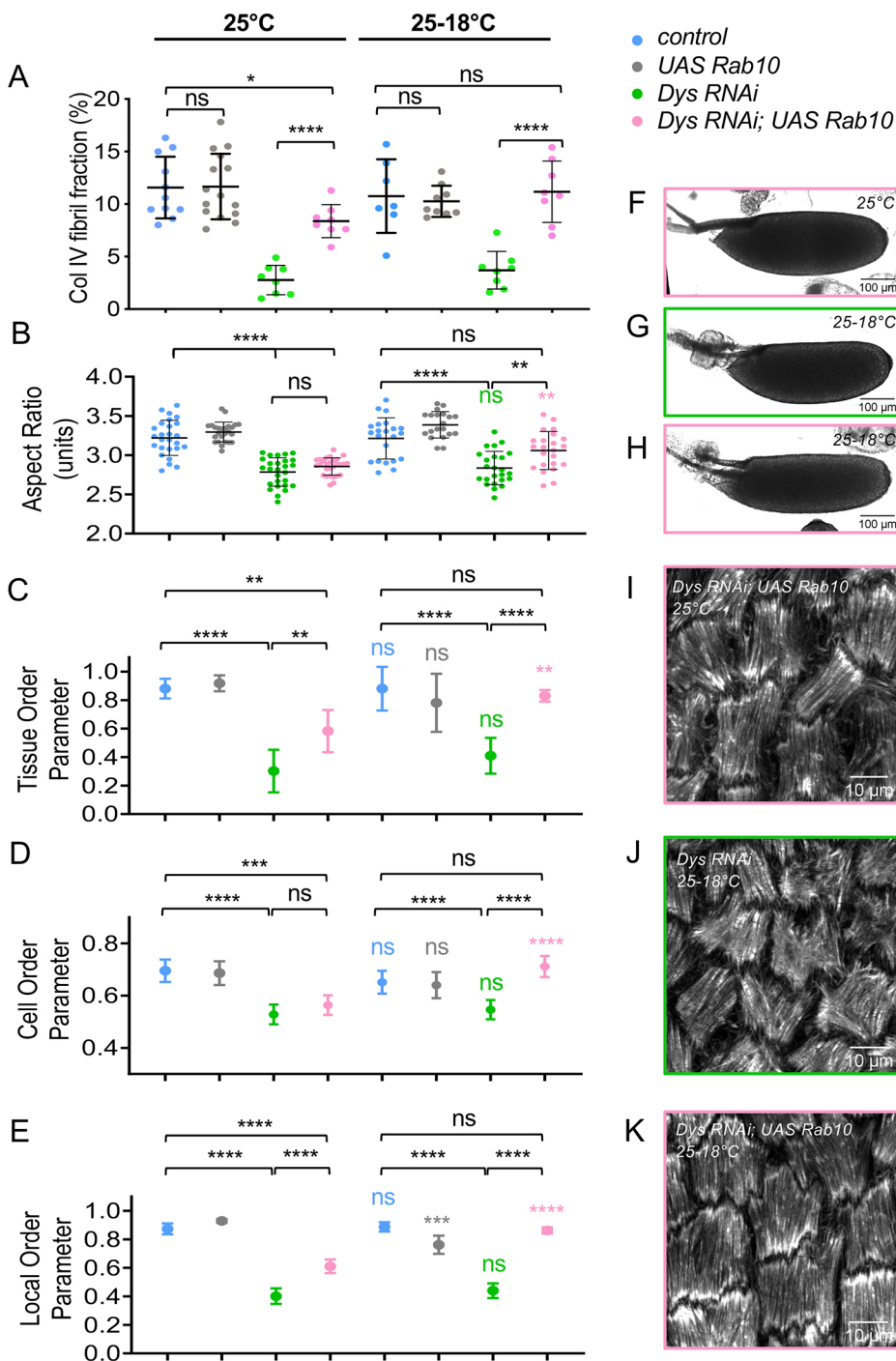


Fig. 7. BM fibrils provide a cue for DAPC-dependent F-actin orientation.

(A) Quantification of the BM fibril fraction at stage 8 for the indicated genotypes at 25°C ($n > 8$). (B-E) For the indicated genotypes, without (25°C) or with (25-18°C) a switch to non-permissive temperature, quantification of the aspect-ratio of mature eggs (B), the Tissue-OP ($n > 7$ follicles) (C), the Cell-OP (D), and the Local-OP at stage 13 ($n > 45$ cells) (E). (A,B) unpaired t -test; (C-E) two-way ANOVA performed, followed by applicable post-hoc pairwise comparisons (Tukey). (F-K) Representative images of mature eggs (F-H) and of stress fibers (I-K) at stage 13 for the indicated genotypes without (25°C) or with (25-18°C) a switch to non-permissive temperature. For all panels, error bars represent s.d.; * $P < 0.01$, ** $P < 0.005$, *** $P < 0.001$, **** $P < 0.0001$. Colored asterisks (or ns) correspond to the statistical test for the same genotype between 25°C and 25-18°C conditions. ns, not significant.

that we analyzed only if the BM fibrils have been restored by *Rab10* overexpression (Fig. 7C-E,J,K, Fig. S5J). Moreover, this condition also restores egg elongation (Fig. 7B,F-H). These experiments demonstrate the double function of the DAPC, with the latter in F-actin organization depending on the first one in BM structure. It also shows that the BM fibrils are required for F-actin orientation long after their deposition and that this function explains their contribution to follicle elongation.

DISCUSSION

In this article, we show that the DAPC contributes to follicle elongation by two distinct functions in which the second relies on the output of the first one. These results are important for understanding

the activity of this key complex and the morphogenetic mechanisms involving the ECM.

Follicle elongation relies on multiple mechanisms

Genetic data has already demonstrated that follicle elongation relies on at least two different and successive mechanisms. The first is controlled by JAK-STAT and involves the follicle cell apical domain, whereas the second is controlled by Fat2 and involves the basal domain and the BM (Alégot et al., 2018; Aurich and Dahmann, 2016). Between these phases, around stage 7-8, JAK-STAT and Fat2 seem to be integrated in a third mechanism based on a BM stiffness gradient (Crest et al., 2017). Interestingly, a very recent report suggests that this gradient may not directly influence

tissue shape but rather do so by modifying the properties of the follicle cells underneath (Chen et al., 2019). Here, we show that the DAPC influences elongation mainly at very late stages, suggesting the existence of a fourth mechanistic elongation phase. Of note, elongation at these late stages is also defective in *fat2* mutants (Alégot et al., 2018; Aurich and Dahmann, 2016). This is consistent with the fact that rotation is required for polarized BM fibril deposition, and that this deposition depends on and is required for DAPC function. The existence of multiple and interconnected mechanisms to induce elongation, a process that initially appeared to be very simple, highlights the true complexity of morphogenesis, and the necessity to explore it in simple models.

BM fibrils act as a planar polarity cue and F-actin as a molecular corset

Fat2 is clearly part of the upstream signal governing the basal planar polarization. However, how this polarization leads to tissue elongation is still debated. It has been proposed that elongation relies on a molecular corset that could be formed, non-exclusively, by BM fibrils or F-actin stress fibers (Gutzeit, 1990; Gutzeit et al., 1991; Haigo and Bilder, 2011; Frydman and Spradling, 2001; Bateman et al., 2001; Viktorinová et al., 2009; Cetera et al., 2014). The initial observation that rotation is required for both elongation and BM fibrils favored a direct mechanical role for these structures (Haigo and Bilder, 2011). Recent data showing that BM fibrils are stiffer than the surrounding ECM supports this view (Chlasta et al., 2017). Moreover, increasing the BM fibril number and size can lead to hyper-elongation (Isabella and Horne-Badovinac, 2016). Finally, addition of collagenase induces follicle rounding, at least at some stages, and genetic manipulation of the ECM protein levels also influences elongation (Haigo and Bilder, 2011; Isabella and Horne-Badovinac, 2015a,b; Chlasta et al., 2017). However, these experiments did not discriminate between the function of the fibril fraction and a general BM effect. Moreover, they do not demonstrate whether their impact on elongation is direct and mechanic or, indirect by a specific response of the epithelial cells. Fat2 and rotation are also required for the proper orientation of the stress fibers. The F-actin molecular corset is dynamic with follicle cells undergoing basal pulsations, and perturbation of both these oscillations and of the stress fiber structure affect elongation (Bateman et al., 2001; He et al., 2010). In the DAPC mutants, we observed a faint but significant elongation defect during mid-oogenesis and a stronger one after stage 12. These defects are clearly correlated with the stress fiber orientation defects observed in the same mutants, both temporally and in terms of

intensity. Moreover, although overexpression of *Rab10* in a *Dys* loss-of-function mutant restores BM fibrils, it does not rescue elongation, indicating that stress fiber orientation is instrumental.

Thus, if the role of the BM fibrils as a direct mechanical corset appears limited, what is their function? One possibility could have been that they promote rotation, acting by positive feedback and explaining the speed increase over time. However, the rotation reaches the same speed in WT and DAPC mutants, excluding this possibility. Similarly, increasing the fibril fraction also has no effect on rotation speed (Isabella and Horne-Badovinac, 2016).

Our results strongly argue that BM fibrils act as a cue for the orientation of stress fibers, which then generate the mechanical strain for elongation. This appears clear in late stages when the function of the DAPC for stress fiber orientation is dependent on the previous BM fibril deposition. Although it is unknown why the cells lose their orientation from stages 10 to 12, the BM fibrils provide the long-term memory of the initial PCP of the tissue, allowing stress fiber reorientation (Fig. 8). Such a mechanism appears to be a very efficient way to memorize positional cues, and could represent a general BM function in many developmental processes.

How BM fibrils contribute to stress fiber orientation remains to be elucidated. Nevertheless, we can hypothesize that it involves their higher stiffness or a higher density in binding sites for ECM receptors. Our data indicate that the DAPC might be such a receptor. This hypothesis is supported by our findings that the DAPC acts in a cell-autonomous manner and promotes stress fiber orientation in each cell and acts locally rather than at a tissue scale. Of note, this kind of PCP is unusual because it does not require signal transmission between neighboring cells, usually through heterophilic protein-protein interactions. However, an interesting parallel with the PCP established by Fat/Dachsous and the core PCP pathways could be drawn. In many tissues, these two pathways work in parallel and their cooperation is required for proper PCP establishment at the apical domain of epithelial cells (Hale and Strutt, 2015). The genetic interaction that we revealed between Fat2 and the DAPC suggests that robust basal PCP establishment also requires the integration of two independent mechanisms. Importantly, this genetic interaction applies to at least two tissues, suggesting a general functional link between them.

The DAPC acts as an organizer of the basal domain of epithelial cells

We found that the DAPC impacts the two key actors at the basal domain of the follicle cells: the BM and the stress fibers linked to the

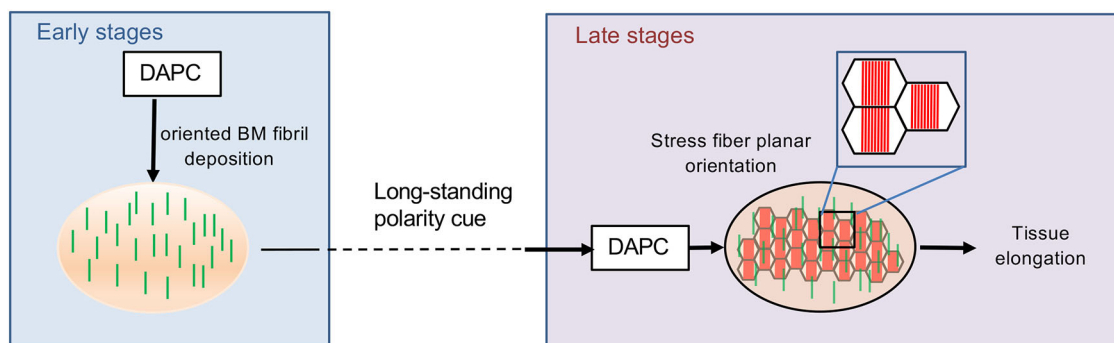


Fig. 8. Dual function of the DAPC during elongation revealing that BM fibrils work as a persistent planar polarity cue. During the early stages (4–8) the DAPC participates in the formation of oriented BM fibrils perpendicular to the follicle elongation axis. These fibrils are maintained during follicle development including stages with intense morphogenetic processes unrelated to elongation (stages 9–11). At late stages, cells use the persistent BM fibrils as a signal to guide the planar orientation of the stress fibers. This event depends on a distinct second function of the DAPC. The stress fibers are the main mechanical effector of follicle elongation.

BM. All the defects of *Dg* null mutants were also observed in *Dys* mutants, demonstrating a developmental and morphogenetic role for this gene. In vertebrates, at least in some tissues, *Dg* presence appears to be essential for BM assembly (Henry and Campbell, 1998; Satz et al., 2010; Bello et al., 2008; Buisson et al., 2014). However, BM formation on the follicular epithelium does not require *Dg*, suggesting the existence of alternative platforms for its general assembly. Our genetic data suggest that *Rab10* is epistatic to *Dg* for BM fibril deposition. The usual interpretation of such a result would be that *Dg* is involved in the targeting of ECM secretion upstream of *Rab10* rather than in ECM assembly in the extracellular space. In *Caenorhabditis elegans*, *Dg* acts as a diffusion barrier to define a precise subcellular domain for ECM remodeling (Naegeli et al., 2017). One could imagine that the DAPC has a similar function in follicle cells, by defining the position where the *Rab10* secretory route is targeted. However, in DAPC loss of function, some ECM is still secreted between cells, suggesting that the lateral *Rab10* route is not affected. Moreover, ECM proteins do not abnormally accumulate between cells in such mutants, suggesting that they are able to leave this localization but without forming BM fibrils. Therefore, the functional interplay between *Rab10* and the DAPC is still unclear.

As mentioned before, *Dg* has often been proposed to act as a scaffold to promote BM assembly in mice. Deletion of the *Dg* intracellular domain is only sub-lethal in mice, whereas complete loss of this protein is lethal very early during development, indicating that the abolishment of *Dg*'s interaction with *Dys* affects its function only partially (Henry and Campbell, 1998; Satz et al., 2009). In these mice, laminin assembly can still be observed, for instance in the brain and retina (Satz et al., 2010; Clements et al., 2017). Similar results were also obtained in cultured mammary epithelial cells (Weir et al., 2006). Thus, despite the existence of *Dys* paralogs that could mask some effects on ECM and the fact that we observed the same ECM alteration in *Dg* or *Dys* mutant fly follicles, not all the *Dg* functions related to ECM assembly or secretion involve *Dys*. It is possible that *Dys* is required when *Dg* needs a very specific subcellular targeting for its function, whereas a more general role in ECM assembly would be independent of *Dys*. Our results suggest that some specific effects of *Dys* on ECM could have been underestimated and this could help to explain the impact of its loss of function on tissue integrity maintenance. For instance, as it has been reported that *Dg* influences ECM organization in fly embryonic muscles, it would be interesting to determine whether this also involves *Dys* (Yatsenko and Shcherbata, 2014).

The DAPC is involved in planar polarization of the basal stress fibers and its ability to read ECM structure to orchestrate integrin-dependent adhesion could play a role in many developmental and physiological contexts. The link between the ECM and F-actin provided by this complex is likely required for this function, although this remains to be formally demonstrated. BM fibrils could provide local and oriented higher density of binding sites for *Dg*, and the alignment could then be transmitted to the actin cytoskeleton. Alternatively, DAPC function could rely on sensing the mechanical ECM properties. The hypothesis that the DAPC could act as a mechanosensor is a long-standing proposal, partly due to the presence of spectrin repeats in *Dys* (Brown and Lucy, 1993; Garbincius and Michele, 2015). The basal domain of the follicle cells may offer an amenable model to combine genetics and cell biology approaches to decipher such function.

Altogether, this work provides important insights on the role of the BM during morphogenesis, by acting as a static PCP cue retaining spatial information while cells are highly dynamic. It also

reveals important functions of the DAPC, including *Dys*, that may be broadly involved during animal development and physiology.

MATERIALS AND METHODS

Genetics

The detailed genotypes, temperature and heat-shock conditions are given in Table S1.

Live imaging and rotation speed analysis

For live imaging, ovaries were dissected, as described previously (Alégot et al., 2018), and stained with the membrane dye FM4-64. Samples were cultured for less than 1 h before imaging. Damaged follicles (identified on the basis of abnormal FM4-64 incorporation) were not analyzed. Rotation speed analysis was performed using a Fiji Manual Tracking plug-in. For each follicle, speed was measured at three independent positions during 30 min.

Dissection and immunostaining

Resources and reagents are listed in Table S2. Dissection and immunostaining were performed as described previously (Vachias et al., 2014) with the following changes: ovaries were dissected in supplemented Schneider medium, and ovarioles were separated and the muscle sheath was removed before fixation to obtain undistorted follicles. Stage was determined using unambiguous reference criteria that are independent of follicle shape (Spradling et al., 1993). For quantification of stress fiber orientation, ovaries were fixed with 8% paraformaldehyde (PFA) containing phalloidin-TRITC or phalloidin Atto-488. They were then permeabilized with 0.02% Triton X-100 in PBS at 0.02% and incubated again with phalloidin-TRITC (or Atto-488). Images were taken using a Leica SP8 or a Zeiss LSM800 Airyscan confocal microscope except for stage 12 BM images, which were acquired on an inverted Zeiss Spinning Disc (Yokogawa CSU-22). Immunostaining of pericellular ECM was performed as described by Isabella and Horne-Badovinac (2016), using antibodies against GFP (ColIV-GFP-expressing flies) or against LanA (a gift from T. Volk, Weizmann Institute, Rehovot, Israel).

Aspect ratio determination

The length of the long and short axes of each follicle were directly measured by transmitted light microscopy and used to determine the aspect ratio.

BM fraction, pericellular matrix and F-actin orientation quantification

BM fibril fraction was determined using a homemade Fiji macro, using the principles developed by Isabella and Horne-Badovinac (2016). Threshold for fibril fraction determination was manually adjusted on each control experiment to obtain accurate fibril detection, taking into account the slight variation of the fluorescence depending on the temperature, the developmental stage and the microscope used. The same threshold was then assigned for all the genotypes of the same experiment. The pericellular matrix was quantified using the Fiji tool to measure the 'meangray value' of a 1.32- μm -wide line drawn on the cell walls (about 100 lines for each genotype, 20 cells).

Stress fiber orientation and Tissue OP analysis were performed as described by Cetera et al. (2014), after manual segmentation. Analyses were performed on 40 and 96 μm^2 images for the early and late stages, respectively. The Cell OP was determined in the same way, by analyzing the alignment robustness at the cellular scale. For Local OP, the average cell radius of each image was determined. Then, we analyzed the cells with their barycenter being in 3.5 radiuses from the considered cell, this value fitting with an average of six neighbors. Clusters with fewer than four cells were excluded. Then, the radius was progressively increased from 3.5 to 25. At 25, all the cells of the image were included, and thus the Local OP value corresponded to the Tissue OP value. Therefore, this approach was used to observe the tendency from Local OP to Tissue OP and to calculate for each cell the difference between them. As the orientation of the fibers is only between 0° and 180°, the OP, as it has been defined, is a measure of the variability among corresponding mean

directors, independently of the scale at which it is used: local directors were used for the Cell OP whereas the mean cell orientation was used to compute the OP for larger scales.

Statistical analyses

For all experiments, the minimal sample size is indicated in the figure legends. Results were obtained from at least two independent experiments, and for each experiment multiple females were dissected. Randomization or blinding was not performed. The normality of the samples was calculated using the D'Agostino and Pearson normality test. The unpaired *t*-test was used to compare samples with normal distribution, and the unpaired Mann–Whitney test for samples without normal distribution.

Concerning the stress fiber angular distribution, the circular mean orientation of the fibers was computed for each cell and then used for the Rao's test of homogeneity. For each stage, a pairwise comparison of genotypes was performed to determine whether the polar vectors or the dispersions were equal. These tests were performed in R using the 'circular' package.

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Competing interests

The authors declare no competing or financial interests.

Author contributions

Conceptualization: F.C.C., H.A., C.F., A.I., S.H.-B., V.M.; Methodology: F.C.C., C.D., H.A., O.B., V.M.; Software: P.P.; Validation: F.C.C., C.D., V.M.; Formal analysis: F.C.C., C.D., H.A., P.P., V.M.; Investigation: F.C.C., C.D., H.A., C.F., A.I.; Data curation: H.A.; Writing - original draft: F.C.C., S.H.-B., V.M.; Visualization: F.C.C., C.D., P.P., V.M.; Supervision: S.H.-B., V.M.; Project administration: V.M.; Funding acquisition: S.H.-B., V.M.

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Supplementary information

Supplementary information available online at <http://dev.biologists.org/lookup/doi/10.1242/dev.186957.supplemental>

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Supplemental data

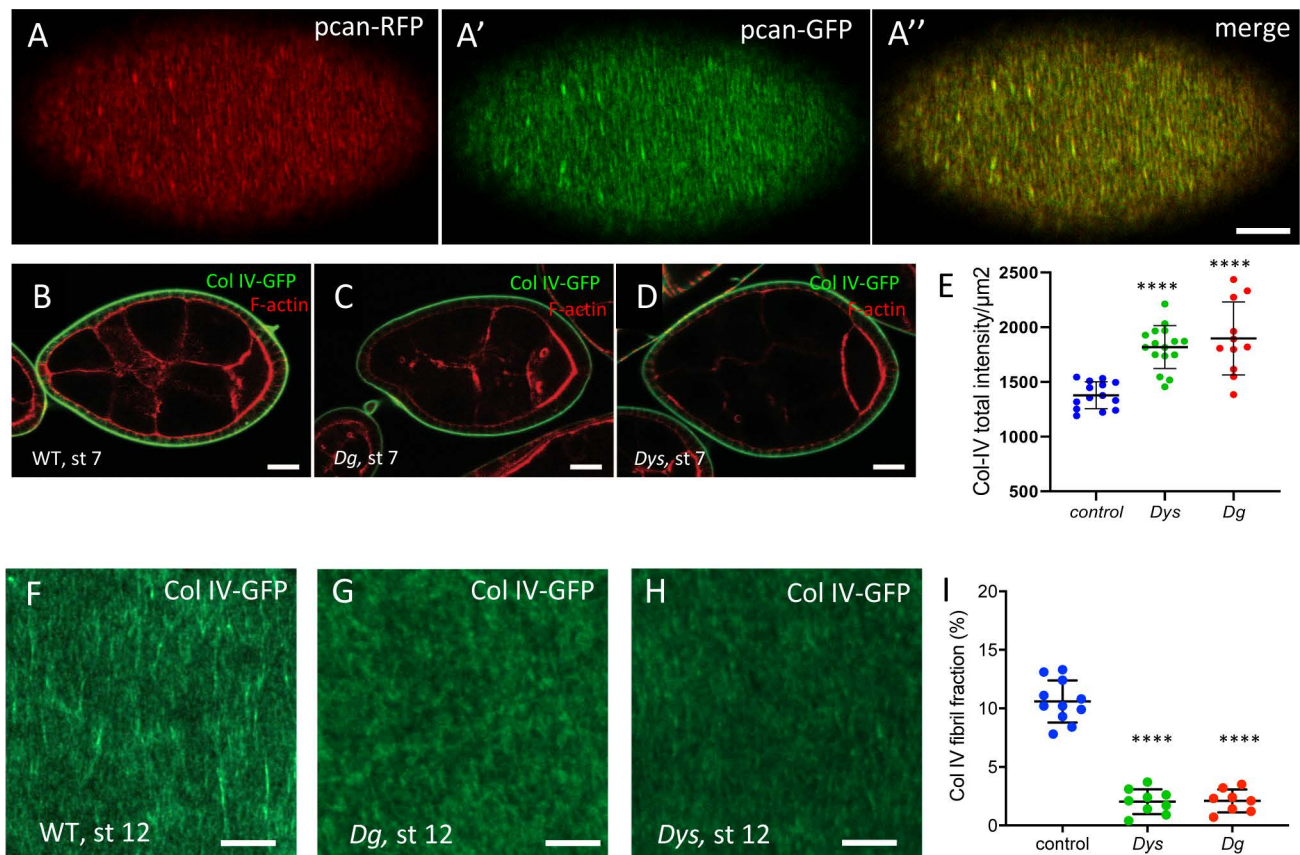


Figure S1

A) Basal view of the ECM of stage 7 from fly trans-heterozygous flies for *Pcan-RFP* and *Pcan-GFP*

B-D) Optical cross-section of stage 7 WT, *Dys*^{E17/Exel6184} and *Dg*^{O46/O83} mutant follicles that express ColIV-GFP and stained for F-actin.

E) Quantification of the total fluorescence intensity per area unit in stage 8 WT, *Dys*^{E17/Exel6184} and *Dg*^{O46/O83} null mutant follicles.

F-H) Basal view of the ECM of stage 12 WT, *Dys*^{E17/Exel6184} and *Dg*^{O46/O83} null mutant follicles that express ColIV-GFP.

I) Quantification of the BM fibril fraction in stage 12 WT, *Dys*^{E17/Exel6184} and *Dg*^{O46/O83} null mutant follicles. Scalebar 10μm.

For all panels error bars represent s.d.; p * < 0.01, ** < 0.005, *** < 0.001, **** < 0.0001.

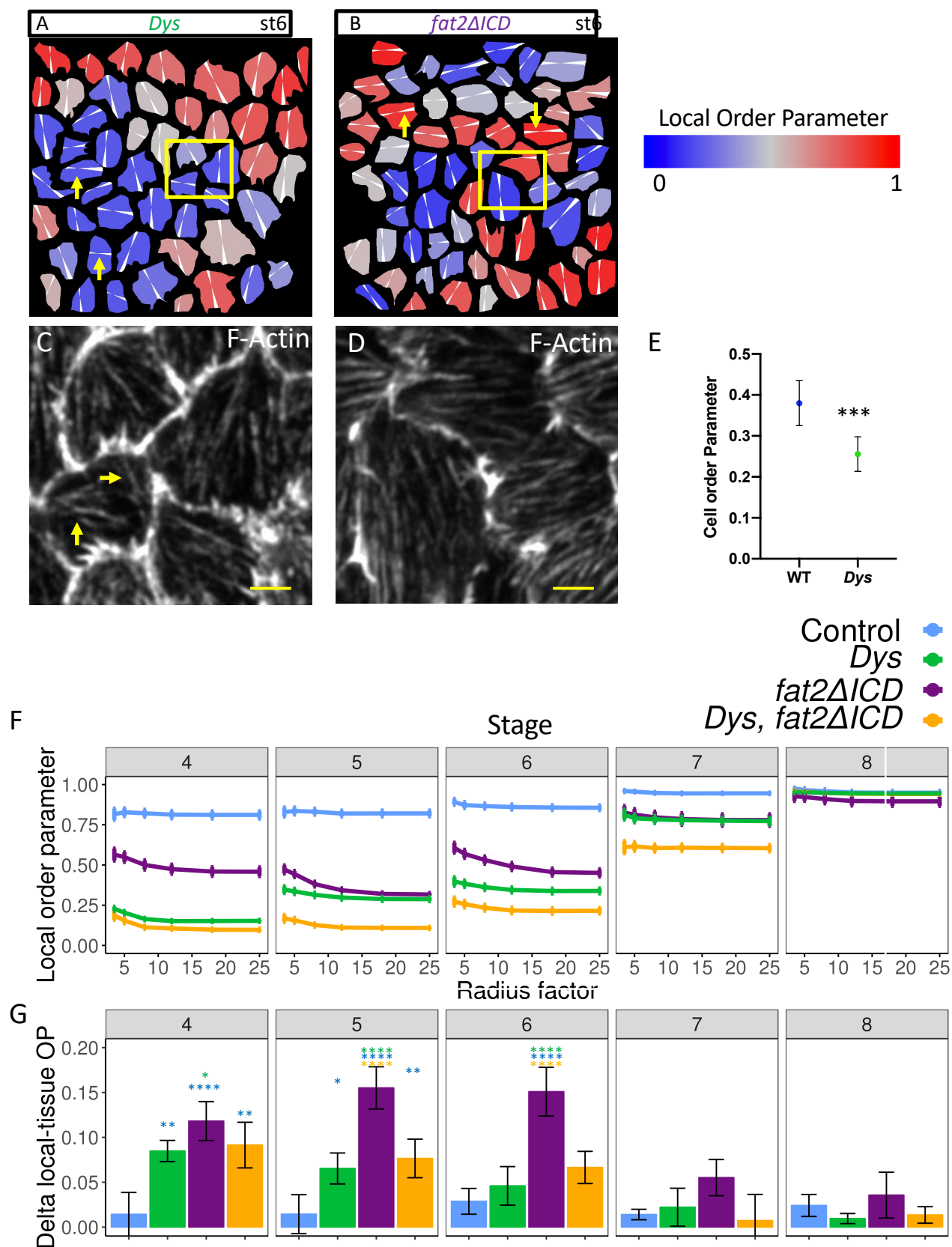


Figure S2:

A-B) Color coding of the Local OP by cell on representative images (B and C of Fig. 4) of A) *Dys*^{E17/Exel6184} and B) *fat2deltaICD* mutants. Main orientation of the stress fibers is indicated in white. Note that cells with a wrong orientation (yellow arrows) can have a high Local OP in *fat2deltaICD* mutant (but not in *Dys* mutant) indicating that they are surrounded by cells with a similar orientation. C-D) Zoom-in corresponding to the yellow squares in A and B. Yellow arrows highlight divergent stress fibers in a same cell in *Dys* mutant.

E) Cell Order Parameter for *Dys* mutant cells and WT neighboring cells at stage 5 (n >138 cells).

F) Representation for stages 4 to 8 of the Local OP as a function of the distance (measured in cell radius), showing the tendency from Local OP to Tissue OP. G) Difference between Local OP (minimal value for radius in E) and Tissue OP (maximal value for radius in E) (n > 171 cells for each stage and each genotype).

For all panels error bars represent s.d.; p * < 0.01, ** < 0.005, *** < 0.001, **** < 0.0001. Scale bar 2µm

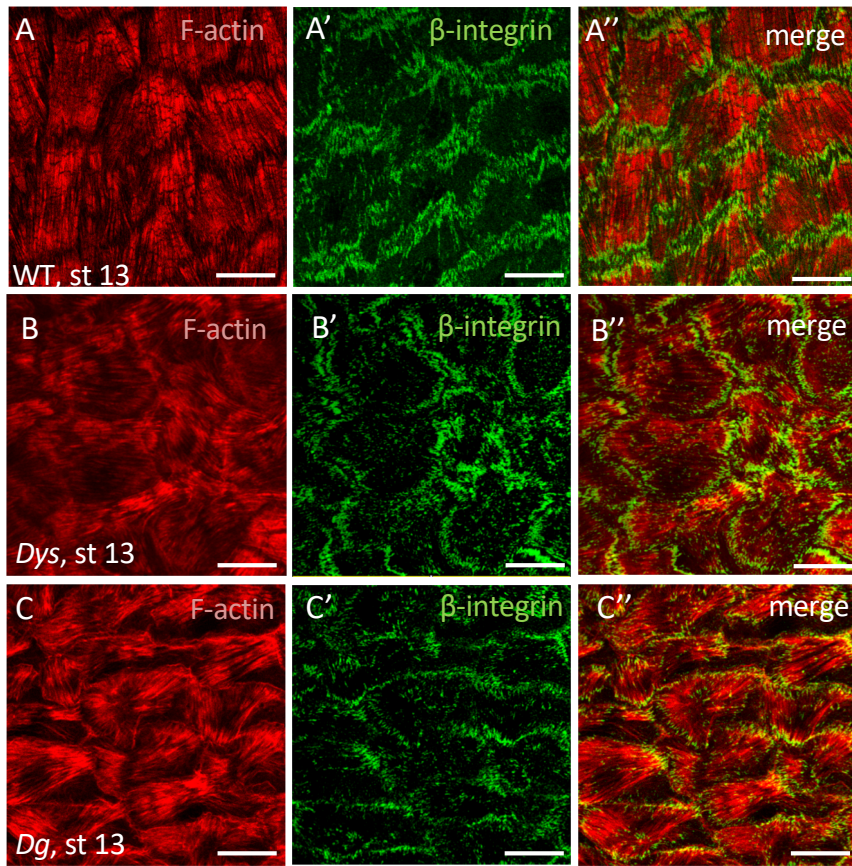


Figure S3:

F-actin and Beta-integrin staining in A) WT, B) *Dys*^{E17/Exel6184} and C) *Dg*^{O46/O83} mutant follicles at stage 13.

Scale bar 10μm

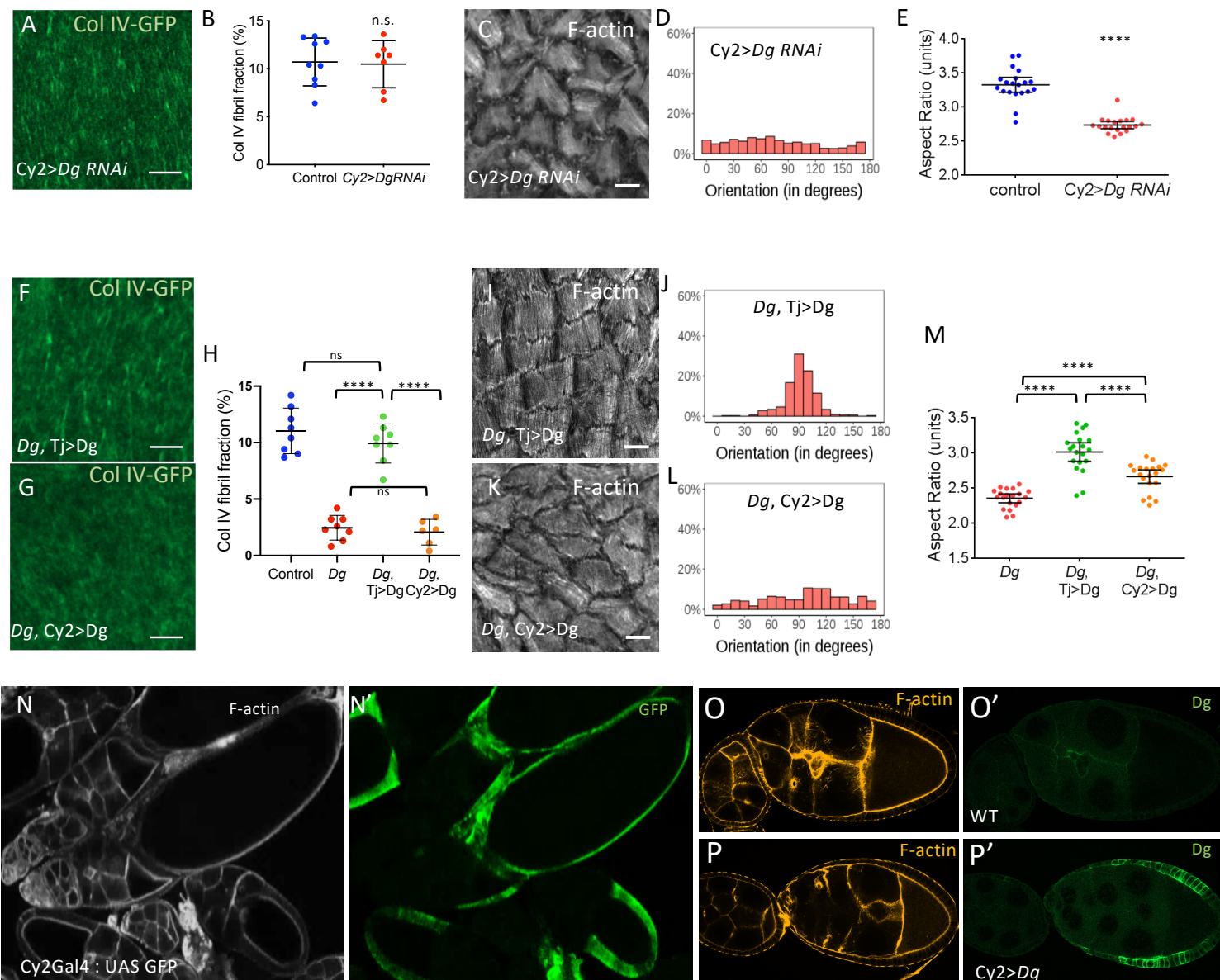


Figure S4:

A,F,G) Basal view of the ECM at stage 12 of A) Dg RNAi driven with Cy2:Gal4, F) *Dg*^{O46/O83} mutant expressing UAS:Dg driven with tj:Gal4, G) Dg mutant expressing UAS:Dg driven with Cy2:Gal4

B,H) Quantification of the BM fibril fraction for the indicated genotypes at stages 12.

C,I,K) Representative images of basal F-actin at stage 13 of C) Dg RNAi driven with Cy2:Gal4,

I) Dg mutant expressing UAS:Dg RNAi driven with tj:gal4, K) Dg mutant expressing UAS:Dg RNAi driven with Cy2:Gal4

D,J, L) Quantification of stress fiber angular distribution of the same genotypes than in C,J,K).

E, M) Aspect-Ratio quantification of mature eggs of the indicated genotypes.

N) Expression profile of Cy2:Gal4. GFP can be detected from stage 9.

O-P) Dg immunostaining in WT and Cy2:Gal4 UAS:Dg flies. Endogenous Dg is not detected but induced expression is clearly detectable at stage 9.

(For all panels error bars represent s.d.; p < 0.01, **<0.005, ***<0.001, ****<0.0001) Scale bar 10µm.

30°C

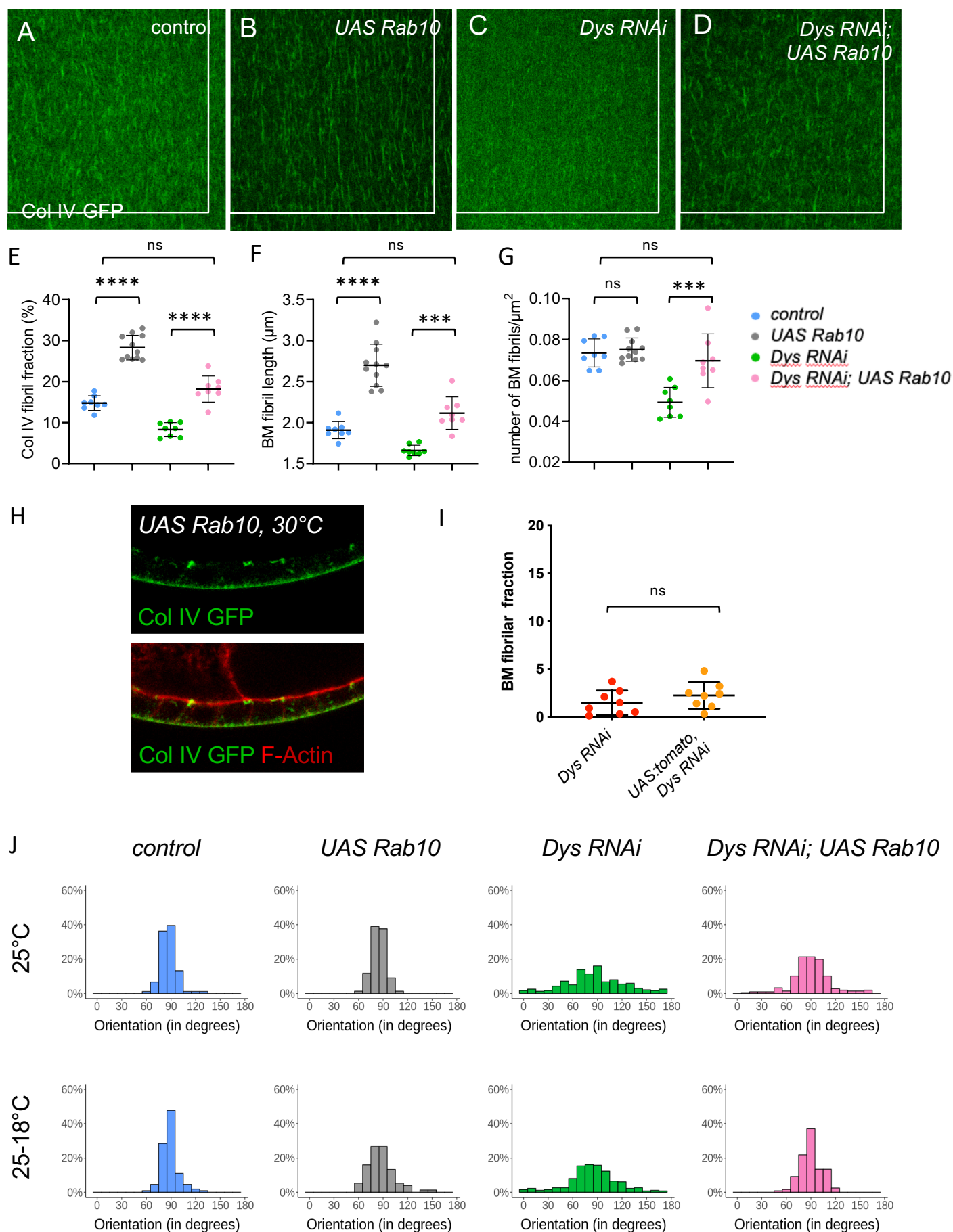


Figure S5:

A-D) Basal view of the ECM at stage 8 of A) WT, B) *tj:Gal4>UAS:Rab10*, C) *tj:Gal4>UAS Dys RNAi*, D) *tj:Gal4> UAS Dys RNAi, UAS Rab10*.

E-G) Quantification of E) BM fibril fraction F) fibril length and G) fibril numbers per μm^2 at stage 8 for the indicated genotypes at 30°C (n>8). (error bars represent s.d.; p *< 0.01, **<0.005, ***<0.001, ****<0.0001)

H) Section view of the follicular epithelium expressing CollIV-GFP and overexpressing Rab10 at 30°C.

I) Quantification of fibril fraction at stage 12 for *tj>Dys RNAi* and *tj>UAS:tomato; Dys RNAi*

J) Angular distribution for the indicated genotypes without (25°C) or with (25-18°C) a switch to non-permissive temperature.

Table S1: Genotypes and specific conditions

Figure		Genotype	Conditions
Figure 1	B, E, F, G, H, I, L C, E, F, G, H, J, L D, E, F, G, H, K, L E, F, G	WT (<i>w¹¹¹⁸</i>) <i>Dys^{E17}/Dys^{Exel6184}</i> <i>Dg^{O43}/Dg^{O86}</i> <i>Dg^{O43}/Dg^{O86}; Dys^{E17}/Dys^{Exel6184}</i>	25°C
Figure 2	A B C D E F, G, H I, J K L	Trol-RFP/+ ; ColIV-GFP/+ Trol-RFP/+ ; ; LanA-GFP/+ <i>Trol-RFP/+ ; ; Dys^{E17}/Dys^{Exel6184}</i> <i>ColIV-GFP/+ ; Dys^{E17}/Dys^{Exel6184}</i> <i>Dys^{E17}/LanA-GFP, Dys^{Exel6184}</i> WT <i>Dys^{E17}/Dys^{Exel6184}</i> <i>Dg^{O43}/Dg^{O86}</i> <i>y, w, HS:flp122/+;;FRT82B, Ubi:RFP^{nls}/FRT82B, Dys^{Exel6184}</i> <i>fat2^{N103-2}/fat2^{58D}</i> <i>fat2^{N103-2}, Dys^{E17}/fat2^{58D}, Dys^{Exel6184}</i>	25°C 25°C, 2HS 1H, dissection 5 days after HS 25°C 25°C
FigureS1	A B, E, F, I C, G, E, I D, H, E, I	Pcan-RFP / Pcan-GFP ColIV-GFP/+ ColIV-GFP, <i>Dg^{O86}/Dg^{O43}</i> ColIV-GFP/+ ; <i>Dys^{E17}/Dys^{Exel6184}</i>	25°C
Figure 3	A-F, T G-L, T M-R, T U, V	WT <i>Dys^{E17}/Dys^{Exel6184}</i> <i>Dg^{O43}/Dg^{O86}</i> <i>y, w, HS:flp122/+ ; ; FRT82B, Ubi:RFP^{nls} / FRT82B, Dys^{Exel6184}</i>	25°C 25°C, 2HS 1H, dissection 5 days after HS
Figure 4	A, I, J, K, L, M, Q, R B, I, J, K, L, N, Q, R C, I, J, K, L, O, Q, R D, I, J, K, L, P, Q, R	WT <i>Dys^{E17}/Dys^{Exel6184}</i> <i>fat2^{N103-2}/fat2DeltaICD-3xGFP</i> <i>fat2^{N103-2}, Dys^{E17}/fat2DeltaICD-3xGFP, Dys^{Exel6184}</i>	25°C
Figure S2	F, G A, C, F, G E B, D, F, G F, G	WT <i>Dys^{E17}/Dys^{Exel6184}</i> <i>y, w, HS:flp122/+ ; ; FRT82B, Ubi:RFP^{nls} / FRT82B, Dys^{Exel6184}</i> <i>fat2^{N103-2}/fat2DeltaICD-3xGFP</i> <i>fat2^{N103-2}, Dys^{E17}/fat2DeltaICD-3xGFP, Dys^{Exel6184}</i>	25°C
Figure 5	A-D E-H I-L M, N	WT <i>Dys^{E17}/Dys^{Exel6184}</i> <i>Dg^{O86}/Dg^{O43}</i> <i>y, w, HS:flp122/+;FRTG13, Ubi:GFP^{nls}/FRTG13, Dg</i>	25°C
Figure S3	A B C	WT <i>Dys^{E17}/Dys^{Exel6184}</i> <i>Dg^{O86}/Dg^{O43}</i>	25°C
Figure 6	B-E F-I	Tj:Gal4, ColIV-GFP; tub:Gal80 ^{ts} /+ Tj:Gal4, ColIV-GFP / UAS: Dg-RNAi ; tub:Gal80 ^{ts} /+ Tj:Gal4, ColIV-GFP; tub:Gal80 ^{ts} /+ Tj:Gal4, ColIV-GFP / UAS: Dg-RNAi ; tub:Gal80 ^{ts} /+	18°C 30°C

	L-O	Tj:Gal4, ColIV-GFP; tub:Gal80 ^{ts} /+	18°C switch at 30°C during 30 hours
	P-S	Tj:Gal4,ColIV-GFP / UAS: Dg-RNAi ; tub:Gal80 ^{ts} /+ Tj:Gal4, ColIV-GFP; tub:Gal80 ^{ts} /+ Tj:Gal4,ColIV-GFP / UAS: Dg-RNAi ; tub:Gal80 ^{ts} /+	30°C switch at 18°C during 72 hours
Figure S4	A-E	Cy2:Gal4, ColIV-GFP/ + Cy2:Gal4, ColIV-GFP/UAS Dg RNAi	Cross at 25°C ,2days at 30°C
	F,H,I,J,M	Tj:Gal4,ColIV-GFP, <i>Dg</i> ^{O86} / UAS: Dg-C, <i>Dg</i> ^{O43}	25°C
	G,H,K,L,M	Cy2:Gal4,ColIV-GFP, <i>Dg</i> ^{O86} / UAS: Dg-C, <i>Dg</i> ^{O43}	25°C
	H,M	ColIV-GFP, <i>Dg</i> ^{O86} / <i>Dg</i> ^{O43}	25°C
	N	Cy2:Gal4/UAS-GFP	18°C
	O	WT	
	P	Cy2:Gal4 /UAS-DG	25°C
Figure 7	A	Tj:Gal4, ColIV-GFP Tj:Gal4, ColIV-GFP /UAS <i>Dys</i> RNAi Tj:Gal4, ColIV-GFP; UAS:Rab10-RFP Tj:Gal4, ColIV-GFP /UAS <i>Dys</i> RNAi ; UAS:Rab10-RFP	25°C
	B-K	Tj:Gal4; tub:Gal80 ^{ts} Tj:Gal4; tub:Gal80 ^{ts} /UAS <i>Dys</i> RNAi Tj:Gal4; tub:Gal80 ^{ts} ; UAS:Rab10-RFP Tj:Gal4; tub:Gal80 ^{ts} /UAS <i>Dys</i> RNAi ; UAS:Rab10-RFP	25°C with or w/o switch at 18°C during 72 hours
Figure S5	A,E,F,G B,E,F,G,H C,E,F,G D,E,F,G I	Tj:Gal4, ColIV-GFP Tj:Gal4, ColIV-GFP; UAS:Rab10-RFP Tj:Gal4, ColIV-GFP /UAS <i>Dys</i> RNAi Tj:Gal4, ColIV-GFP /UAS <i>Dys</i> RNAi ; UAS:Rab10-RFP Tj:Gal4, ColIV-GFP /UAS <i>Dys</i> RNAi Tj:Gal4, ColIV-GFP /UAS <i>Dys</i> RNAi ; UAS:tomato	30°C
	J	Tj:Gal4; tub:Gal80 ^{ts} Tj:Gal4 ; tub:Gal80 ^{ts} ; UAS:Rab10-RFP Tj:Gal4 ; tub:Gal80 ^{ts} /UAS <i>Dys</i> RNAi Tj:Gal4; tub:Gal80 ^{ts} /UAS <i>Dys</i> RNAi ; UAS:Rab10-RFP	25°C with or w/o switch at 18°C during 72 hours

Table S2 : RESOURCES and REAGENT TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
Anti-GFP, goat	Abcam	5450
Laminin, Guinea Pig	(Harpaz and Volk 2012) (Volk lab)	N/A
Beta integrin, mouse	Developmental Studies Hybridoma Bank	CF.6G11
Chemicals, Peptides, and Recombinant Proteins		
FM4-64	Invitrogen	T13320
Phalloidin Alexa568 or Alexa 488	invitrogen	A12380 or A12379
Experimental Models: Organisms/Strains		
<i>w¹¹¹⁸</i>	Bloomington Drosophila Center Stock	FlyBase ID: FBal0018186 BDSC:3605
<i>Dg^{O86}</i>	{Christoforou 2008} (Rob Ray)	FlyBase ID: FBal0218114
<i>Dg^{O43}</i>	Christoforou 2008} (Rob Ray)	FlyBase ID: FBal0218112
<i>Dys^{E17}</i>	Christoforou 2008} (Rob Ray)	FlyBase ID: FBal0241311
<i>Dys^{Exel6184}</i> ; <i>FRT 82B</i> , <i>Df(3R)Exel6184</i>	Bloomington Drosophila Center Stock	BDSC: 7663; FlyBase ID: FBab0038239
RNAi Dg: UAS:Dg(KK100828)	Vienna Drosophila Resource Center	FBal0230939 VDRC: v107029
RNAi Dys : UAS :Dys(KK101285)	Vienna Drosophila Resource Center	FBal0231174 VDRC: v108006
Pcan-GFP : <u>P{PTT-un1}trol¹⁹⁷³</u>	Kyoto Stock Center (DGRC)	FlyBase ID: FBal0191079 : DGRC : 110836
collIV-GFP : P{PTT-un1}CollIV ^{G205}	Kyoto Stock Center (DGRC)	FlyBase ID: FBti0074199; DGRC: 110626
LanA-GFP : LanA(fTRG00574.sfGFP-TVPTBF)	Vienna Drosophila Resource Center	VDRC: v318155 FlyBase ID: FBal0339089
Pcan-mCherry	This paper	based on MI03800
UAS:Rab10-RFP	Isabella and Horne-Badovinac, 2016	
UAS: tomato	Bloomington Drosophila Center Stock	BDSC: 30124
Tj:Gal4 : P{w+mW.hs=GawB}NP1624	Kyoto Stock Center	DGRC: 104055 FlyBase ID: FBti0034540
Cy2:Gal4 : w; P{GawB}CY2	{Queenan 1997} (St Johnston lab)	FlyBase ID: FBti0007266

Tub:Gal80ts : w[*]; sna[Sco]/CyO; P{w[+mC]=tubP-GAL80[ts]}7	Bloomington Drosophila Stock Center	BDSC: 67071; FlyBase ID: FBti0027798
<i>Fat2</i> ^{N103-2}	{HorneBadovinac 2012}	FlyBase ID: FBal0267777
<i>Fat2</i> ^{58D}	{Viktorinová 2009} (Dhamann)	FlyBase ID: FBal0241216
UAS:Dg-C	(Deng et al, 2003) (Ruhola-Baker lab)	FlyBase ID: FBal0145088
FRTG13, Ubi:GFP ^{nls} : w[1118]; P{w[+mW.hs]=FRT(w[hs])}G13 P{w[+mC]=Ubi-GFP.nls}2R1 P{Ubi-GFP.nls}2R2	Bloomington Drosophila Stock Center	BDSC: 5826
Fat2-DeltaICD-3xGFP	(Barlan et al, 2017)	FlyBase ID: FBal0326665
y,w, hs:flp : P{ry[+t7.2]=hsFLP}22, w[*]	Bloomington Drosophila Stock Center	BDSC: 8862; FlyBase ID: FBst0008862
Software and Algorithms		
Fiji	(Schindelin et al, 2012)	
R		
Prism		

Table S3: Results of the statistical analysis for Fig 7I-L

test genotype 1- genotype 2	LOP p value	COP p value	TOP p value	Delta LOP -TOP p value
Stage 4:				
WT-Fat2ΔICD	<.0001	0.9990	0.0008	<.0001
WT-Dys,Fat2ΔICD	<.0001	<.0001	<.0001	0.0038
WT-Dys	<.0001	<.0001	<.0001	0.0023
Fat2ΔICD-Dys,Fat2ΔICD	<.0001	<.0001	<.0001	0.3807
Fat2ΔICD-Dys	<.0001	<.0001	0.0055	0.0452
Dys,Fat2ΔICD-Dys	0.0613	0.8408	0.4865	0.9629
Stage 5:				
WT-Fat2ΔICD	<.0001	<.0001	<.0001	<.0001
WT-Dys,Fat2ΔICD	<.0001	<.0001	<.0001	0.0059
WT-Dys	<.0001	<.0001	<.0001	0.0166
Fat2ΔICD-Dys,Fat2ΔICD	<.0001	<.0001	0.0250	<.0001
Fat2ΔICD-Dys	<.0001	<.0001	0.9973	<.0001
Dys,Fat2ΔICD-Dys	<.0001	0.3865	0.0413	0.8432
Stage 6:				
WT-Fat2ΔICD	<.0001	<.0001	0.0004	<.0001
WT-Dys,Fat2ΔICD	<.0001	<.0001	<.0001	0.1535
WT-Dys	<.0001	<.0001	<.0001	0.7643
Fat2ΔICD-Dys,Fat2ΔICD	<.0001	<.0001	0.0209	<.0001
Fat2ΔICD-Dys	<.0001	<.0001	0.8242	<.0001
Dys,Fat2ΔICD-Dys	<.0001	0.6676	0.1256	0.3830
Stage 7:				
WT-Fat2ΔICD	<.0001	0.0549	0.5779	0.2422
WT-Dys,Fat2ΔICD	<.0001	<.0001	0.0079	0.9879
WT-Dys	<.0001	<.0001	0.5052	0.9830
Fat2ΔICD-Dys,Fat2ΔICD	<.0001	<.0001	0.3924	0.0844
Fat2ΔICD-Dys	0.9693	<.0001	1.0000	0.4280
Dys,Fat2ΔICD-Dys	<.0001	0.5118	0.4610	0.8844
Stage 8:				
WT-Fat2ΔICD	0.4559	0.0574	0.9600	0.9742
WT-Dys,Fat2ΔICD	0.9388	0.5843	1.0000	0.9784
WT-Dys	0.9164	0.9753	1.0000	0.9538
Fat2ΔICD-Dys,Fat2ΔICD	0.7539	0.4490	0.9685	0.8093
Fat2ΔICD-Dys	0.8422	0.0979	0.9620	0.7549
Dys,Fat2ΔICD-Dys	0.9995	0.7944	1.0000	0.9986