Regulators of cell movement during development and regeneration in *Drosophila*

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Cell migration is a fundamental cellular biological process essential both for normal development and tissue regeneration after damage. Cells can migrate individually or as a collective. To better understand the genetic requirements for collective migration, we expressed RNA interference (RNAi) against 30 genes in the *Drosophila* embryonic salivary gland cells that are known to migrate collectively. The genes were selected based on their effect on cell and membrane morphology, cytoskeleton and cell adhesion in cell culture-based screens or in *Drosophila* tissues other than salivary glands. Of these, eight disrupted salivary gland migration, targeting: Rac2, Rab35 and Rab40 GTPases, MAP kinase-activated kinase-2 (MAPk-AK2), RdgA diacylglycerol kinase, Cdk9, the PDSW subunit of NADH dehydrogenase (ND-PDSW) and actin regulator Enabled (Ena). The same RNAi lines were used to determine their effect during regeneration of X-ray-damaged larval wing discs. Cells translocate during this process, but it remained unknown whether they do so by directed cell divisions, by cell migration or both. We found that RNAi targeting Rac2, MAPk-AK2 and RdgA disrupted cell translocation during wing disc regeneration, but RNAi against Ena and ND-PDSW had little effect. We conclude that, in *Drosophila*, cell movements in development and regeneration have common as well as distinct genetic requirements.

1. Introduction

Collective cell migration is important for forming the complex architecture of tissues and organs during embryogenesis and also plays an important role in cancer progression. Examples include neural crest cells that migrate as a group and sheets of epithelial cells that migrate collectively during gastrulation [1]. Unlike single migrating cells, collectively migrating cells face the additional challenge of having to coordinate the activities of all cells in the group to achieve directed movement. Studies in vertebrate and invertebrate model organisms have identified a number of molecular features of collective cell migration. These include the identification of leader and follower cells, the molecular basis for adhesive contacts among cells and between cells and the surrounding environment, and the collective response to guidance cues. How these are regulated to allow sufficient plasticity for collective migration while maintaining tissue integrity remains an active area of research.

The *Drosophila* embryonic salivary gland is a well-established experimental system for studying collective cell migration [2–4]. The gland consists of a pair of elongated secretory tubes that are connected to the larval mouth by fine duct...
tubes. Salivary gland development begins with invagination of primordial cells from the embryo surface followed by collective migration of the gland as an intact organ. There is no cell death or cell proliferation throughout gland development and the gland cells retain their epithelial characteristics during morphogenesis. Collective migration of the salivary gland occurs through coordinated migration of the distal and proximal gland cells. While distal gland cells elongate and extend actin-rich basal membrane protrusions in a process dependent on Rac GTPases, proximal gland cells change shape from columnar to cuboidal and rearrange in a Rho1 GTPase-dependent manner [5–8].

Rac/Rho-dependent intracellular changes are governed by the activity of integrin receptors at the sites of contact between salivary gland cells and between these cells and the substratum. Stable microtubules and the KASH-domain containing protein Klarsicht are responsible at least partially for localizing integrins at the contact sites [9]. Integrin adhesion receptors, αPS1βPS (expressed in the salivary gland) and αPS2βPS (expressed in the surrounding mesoderm) concentrate at sites of contact between the migrating gland cells and the surrounding mesoderm-derived tissues [10]. Loss of integrin expression or alteration of integrin localization results in gland migration defects with gland cells being unable to initiate or complete posterior migration, respectively [2,9]. One mechanism by which αPS1βPS integrin controls salivary gland migration is by downregulating E-cadherin and promoting basal membrane protrusions through Rac1 in the distal gland cells [6]. Dynamin GTPase, which mediates endocytosis, is also required for E-cadherin downregulation in migrating gland cells [5], suggesting that Rac1 may act through endocytosis to downregulate E-cadherin. E-cadherin downregulation is likely to be temporally and spatially regulated in the migrating salivary gland such that cells acquire sufficient plasticity for collective migration while maintaining enough cell–cell adhesion to ensure tissue integrity.

Cell movements occur not only during normal development but also during tissue regeneration. Compared with normal development, cell movements during regeneration are even less well understood. Drosophila larval wing discs have been established as a system to study regeneration after a variety of damage including surgical ablation, genetic ablation through the expression of apoptotic genes, and by ionizing radiation (IR) [11]. We reported before that cells change position during regeneration of larval wing discs damaged by IR. IR induces apoptosis in the single layer epithelium of the larval wing disc. IR-induced apoptosis is scattered but not random and occurs instead in an invariant pattern [12]. We found previously that cells of the future hinge are protected from IR-induced apoptosis [12]. Some hinge cells then lose their hinge fate and translocate to the wing pouch area that suffers more IR-induced apoptosis, where the hinge cells convert to the pouch fate and participate in regeneration of the pouch. Signalling through Drosophila STAT92E (homologue of STAT3/5) and Wingless (homologue of Wnt1) are required cell autonomously for IR-induced regenerative behaviour; knocking down each with RNAi or genetic inhibitors (e.g. Axin against Wg) prevented the translocation and fate change by the hinge cells [12]. Using this model, we have uncovered the requirement for epigenetic regulators of IR-induced fate change and translocation [13]. But cell biological mechanisms by which former hinge cells translocate from the hinge into the pouch remained completely unknown. We do not know even whether the hinge cells migrate as opposed to use directed cell divisions that ‘push’ daughter cells towards the pouch.

To better understand how cellular adhesion and the cytoskeleton control collective cell migration, we used RNA interference (RNAi) against a focused group of 30 genes known or predicted to affect cell or membrane morphology, adhesion and cytoskeleton. We identified eight lines that, when expressed specifically in the salivary gland, disrupt gland migration. These include four genes with previously unknown roles in collective migration of the salivary gland. To address whether common genetic requirements contribute to cell migration in salivary glands and cell position changes during regeneration in the wing disc, we tested a subset of the RNAi lines also in the wing disc. The results identified genes with previously unknown roles in regeneration and suggest that epithelial cell movements during development and regeneration have overlapping as well as distinct genetic requirements.

2. Results

To test for a cell-autonomous requirement in the salivary glands, we generated a recombinant Drosophila line with UAS-GFP-NLS (green fluorescent protein (GFP) tagged with 14 nuclear localization signals) and the fkh-GAL4 driver (for gland-specific expression) on the third chromosome. This recombinant line was then crossed with each of 30 RNAi lines (30 genes). We chose the genes based on previous reports that depletion by RNAi of these genes led to defects in cell or membrane morphology, cytoskeleton or cell adhesion in Drosophila cell culture or in tissues other than the salivary gland [14–17] (electronic supplementary material, table S1). Another selection criterion was for genes for which transgenic lines were available at the time the project was initiated and had RNAi constructs inserted on the second chromosome. This was in case we needed to generate a stock that expressed both RNAi (chromosome II) and UAS-GFP-NLS fkh-GAL4 (chromosome III) in the future. For each RNAi line tested, we analysed the shape of GFP-expressing glands in a minimum of 20 live embryos. If at least five GFP-expressing glands showed abnormal morphology indicative of a possible migration defect, then the phenotype was confirmed in fixed embryos immunostained for the salivary gland marker protein dCREB-A. Approximately 20 stage 14 glands each were analysed in fixed embryos and the penetrance ranged from 5 to 10 defective glands per sample.

Of the 30 RNAi lines, we identified eight that inhibited gland migration when expressed cell autonomously, targeting: enabled, which encodes an actin-binding protein; rdgA, which encodes a diacylglycerol kinase; MAP kinase-activated kinase 2 (MAPk-AK2); ND-PDSW, which encodes a subunit of NADH dehydrogenase (to be called PDSW hereafter); cyclin-dependent kinase 9 (Cdk9); and Rac2, Rab35 and Rab40, which encode small GTPases. Rac and Rab GTPases are already known to be required for salivary gland migration (for example, [5,6]). Cdk9 has a well-documented role in general transcription [18] so defects due to its depletion may be indirect. Therefore, we focused our analysis on PDSW, RdgA, Enabled (Ena) and MAPk-AK2 to understand their roles in collective migration of the salivary gland.
2.1. Salivary glands failed to complete posterior migration in enabled, rdgA, MAPk-AK2 and PDSW mutant embryos

In wild-type embryos, cells of the salivary gland form a tube that first elongates dorsally before starting to migrate posteriorly. Posterior migration begins with cells at the distal end of the salivary gland but by embryonic stage 14 both the distal and proximal parts of the salivary glands have turned and migrated posteriorly (figure 1a; [19]). By contrast, in embryos of the same stage where dsRNA against ena was expressed specifically in the salivary gland, the distal gland turned and migrated posteriorly whereas the proximal gland did not (figure 1b). To confirm the gland migration defect observed with RNAi knockdown, we analysed ena zygotic loss of function mutant embryos for gland migration defects. ena\(^{210}\) is a single C to T change resulting in an A97 V change in the conserved EVH1 (enabled/VASP homology 1) domain; ena\(^{25}\) is an A to G change resulting in an N379F change in the proline-rich domain as well as an A to T change resulting in a stop codon to delete the EVH2 domain; ena\(^{2029}\) is a transposition insertion in the 5'-UTR [20]. In embryos homozygous mutant for ena\(^{210}\) or ena\(^{25}\), salivary glands failed to migrate completely, with the distal gland turned posteriorly but not the proximal gland (figure 1c and data not shown). This phenotype is similar to glands where ena was knocked down with RNAi. During posterior migration, proximal gland cells rearrange to form a narrower and more elongated tube [7]. Proximal glands of ena\(^{25}\) and ena\(^{2029}\) mutant embryos showed an average of 10.0 ± 0.8 nuclei per cross section (\(n = 7\) glands) compared with 8.0 ± 1.1 in wild-type (\(n = 3\) glands; for examples of nuclear density changes, see fig. 2 of [21] and electronic supplementary material, figure S1). The difference was significant (\(p = 0.009\), suggesting that proximal gland cells failed to rearrange in ena mutant embryos. We also analysed salivary gland migration defects in embryos homozygous for MAPk-AK2, PDSW and rdgA mutations and wild-type embryos where MAPk-AK2 or rdgA have been knocked down specifically in the gland. PDSW\(^{10501}\) was generated by P-element insertional mutagenesis (Flybase), and rdgA\(^{3}\) by EMS mutagenesis [22]. Similar to knockdown and loss of function of ena, depletion of MAPk-AK2, PDSW and rdgA resulted in gland migration defects where the distal gland turned posteriorly but the proximal gland did not (figure 1d–h). However, unlike in ena mutants, quantification of the number of nuclei in PDSW mutant embryos and glands with rdgA knocked down showed no statistically significant differences (data not shown).

2.2. Ena localization and salivary gland lumen defects

To better understand how ena contributes to cell migration during salivary gland development, we determined the subcellular localization of Ena by immunostaining migrating salivary glands of wild-type embryos. At embryonic stage 12 when the gland is beginning to turn posteriorly, Ena was found in discrete foci sub-apical to the apical domain marker protein, DaPKC (figure 2a). At stage 14, when the gland had turned posteriorly, Ena localized in a zonular manner sub-apical to DaPKC (figure 2b). Ena also localized to the basal domain of migrating gland cells (figure 2b, b'). Because Ena localized to the apical domain, we determined if the loss of ena affected the lumen morphology and/or the localization of apical proteins. Analysis of the gland lumen with F-actin labelling showed that the lumen morphology of ena\(^{210}\) mutant embryos was irregularly shaped in the distal and medial parts of the gland compared with heterozygous siblings (figure 3a–c). Immunostaining for the apical protein DaPKC (figure 3d,e) and the apical–lateral protein DE-cadherin (DE-cad; figure 4a,b) showed that loss of ena did not affect the apical localization of these proteins. Measurements of the apical domain area showed no difference between ena mutant gland cells and wild-type cells (data not shown). However, apical domains of ena mutant gland cells did not elongate in the proximal–distal direction to the same extent as in wild-type gland cells (figure 4d, quantified from confocal images). Similar to ena mutant gland cells, the apical domain area in PDSW mutant gland cells was not different from that in wild-type cells (data not shown); however, the apical domain failed to elongate in the proximal–distal direction (figure 4c,d) and irregularities in the lumen shape at the distal tip were also observed (figure 4c). RNAi against MAPk-AK2 or rdgA did not produce apparent lumen defects; therefore, the apical domain was not analysed in these embryos.
2.3. Cell division makes a partial contribution to cell translocation during regeneration

To test the universality of the above findings, we turned to another experimental model where cells change their location. Larval imaginal discs develop into adult structures during metamorphosis. The larval wing disc is a single layer of columnar epithelium covered with a layer of squamous cells. The wing disc can regenerate and develop into a normal wing after many types of damage including genetic and surgical ablation of parts of the disc or doses of IR that kill about half of the cells [23–25]. This regeneration occurs without a dedicated stem cell pool. We reported previously that the future wing hinge region of the wing disc shows regenerative properties. Specifically, hinge cells are resistant to killing by IR and can translocate to the wing pouch area that suffers more IR-induced apoptosis, where they convert from hinge to pouch fate and participate in regeneration [12]. This behaviour of the hinge cells was also seen after genetic ablation of the pouch [26].
In our published studies, we monitored the translocation of the hinge cells into the pouch and fate change using a G-trace lineage tracing system [27]. Here, GAL4 drives UAS-RFP (real time marker) and UAS-FLP that catalyses a recombination event to result in stable GFP expression (lineage marker). We used the 30A-GAL4 driver to express G-trace in the hinge as we have done in previous studies (figure 5a–c) [12,13,28]. In un-irradiated discs, red fluorescent protein (RFP) (figure 5a) and GFP (figure 5b) were mostly overlapping (figure 5c), indicating that cell fates are stable. We have shown before that at 72 h after irradiation with 4000R (40 Gy) of X-rays, GFP+RFP− cells are found in the pouch area (figure 5d, enclosed by yellow dashed line) [12]. These are former hinge cells that are expressing the GFP lineage tracer but have lost the hinge fate (became RFP−) and translocated towards the pouch. These cells express the pucker marker VgQ-lacZ [28].

The RFP−GFP+ area enclosed by the yellow line is quantified in Image J [29] and normalized to the RFP+GFP+ area, as a quantitative measure of cell translocation (figure 6a, ‘GAL4 only’). Expression of UAS-Axin (Wg inhibitor) reveals a cell-autonomous requirement for Wg signalling in the translocating cells ([12]; reproduced in new experiments in figure 5e,f, quantified in figure 6e). Because Wg and other genes we study are essential for development, we used GAL80th to repress GAL4 and allow disc development to proceed until mid-third instar larva stage (figure 5m and Material and methods; see also [12]). GAL4 was de-repressed by a shift to 29°C for 24 h before irradiation and irradiated wing discs analysed 72 h after irradiation. This system has been used successfully to identify genes that regulate regenerative behaviour in the hinge cells including Wg, STAT92E, STAT effector and transcription factor Zfh2, and a member of the nucleosome remodelling complex Nurf-38 [12,13,28]. We used this published assay to investigate the mechanism(s) responsible for cell translocation during regeneration.

Directed cell divisions can play a role in the final placement of cells within a tissue. During Drosophila wing development, oriented cell divisions are thought to contribute to the oblong shape of the wing discs and the adult wing because randomizing of division orientation produces a rounded wing [30], although compensatory mechanisms such as cell rearrangements also operate [31]. During cell competition in the wing disc when two groups of cells with different growth characteristics become juxtaposed, dividing ‘winner’ cells orient their mitotic spindle in such a way that their daughter cells end up among the ‘losers’ [32]. The hinge cells could likewise direct their daughter cells towards the pouch, and it may be these directed cell divisions that cause cell translocation during regeneration.

To address this possibility, we blocked cell divisions in the hinge, by expressing Rux, an inhibitor of mitotic Cdk1 activity. We have used this approach before for a different purpose: to keep the number of hinge cells constant while we monitor an abnormal mode of regeneration that produces an ectopic wing disc [13]. Expression of Rux with the same 30A-GAL4 driver, we showed, kept the cell number constant, i.e. inhibited mitotic divisions. Here, we asked whether the same experimental manipulation prevents the translocation of hinge cells into the pouch. Discs expressing 30A-GAL4 > UAS-Rux in the hinge cells show RFP+ cells with enlarged nuclei (figure 5g, compare arrow and arrowhead); this is expected as inhibition of Cdk1 blocks mitosis but still allows repeated rounds of S phase [13,33,34]. We found that the hinge cells still translocated into the pouch after irradiation, albeit less efficiently (figure 5g; ‘Rux’ in figure 6a,b). Rux expression reduced translocation by about threefold (15 ± 10% from 53 ± 18% in GAL4 only controls). Translocated GFP+RFP− cells showed reduced nuclear size, which is expected as cells lose their hinge identity and GAL4/Rux expression (arrow in figure 5g). The finding that Rux reduced but did not eliminate cell translocation suggested that we cannot explain all cell translocation by directed divisions. Instead, cell migration may contribute.

**Figure 4.** Ena and PDSW control apical domain elongation but not size. (a–c) Immunostaining for DE-cadherin detects the apical domain in ena+/+ heterozygous (a) and homozygous embryos (b), and PDSW+/+ homozygous embryos (c). (d) Graph depicting the apical domain elongation ratio in wild-type, ena+/+ and PDSW+/+ salivary gland cells. The number of salivary gland cells analysed for apical domain elongation for each genotype is indicated. p-values against wild-type controls were calculated using a two-tailed t-test. All embryos shown are at stage 14. ena+/+ embryos were immunostained for β-galactosidase (not shown) to distinguish heterozygous from homozygous embryos. Scale bar in (a) represents 5 μm. Images are shown dorsal up and posterior right.
2.4. RNAi against Rac2, MAPk-AK2 and RdgA disrupt cell translocation during regeneration

To address the possibility that cell migration contributes to hinge-to-pouch translocation during regeneration, we tested the effect of some of the RNAi lines used in the salivary gland assay.

From 30 RNAi lines tested in the salivary gland assay, we randomly selected 14 lines to test in the wing discs (three examples shown in figure 5h–j). Ena and Rab35 were not among the 14 randomly selected. But because we detect a role for these genes in the salivary gland assay (this report), we analysed RNAi against Ena and Rab35 in the wing disc assay. Of the total 16 thus analysed in the wing disc, one produced deformed discs even without irradiation and was not considered further (Clatherin heavy chain; figure 5)). We also analysed two empty vector controls (KK and GD) and found that the extent of translocation was lower in the GD discs relative to the KK discs although the difference was not statistically significant \( p = 0.153, \) two-tailed \( t \)-test). Nonetheless, for statistical analysis, we compared each RNAi line with its corresponding vector control, KK or GD. Most \( 14/16 \) RNAi lines thus analysed showed statistically significant \( p < 0.05 \) deviation from the vector controls in the translocation of hinge cells into the pouch but differed in the magnitude of the defect (figure 6). The three strongest hits targeted Ankyrin repeat and LEM domain containing 2 \((\text{Ankle2, CG8465, past1 (CG6148)}\) and an uncharacterized gene \((\text{CG2794})\). Ankle2 is required for \textit{Drosophila} S2 cell spreading [14]. It shows high expression in the \textit{Drosophila} larval central nervous system and \textit{Ankle2} mutant larvae show smaller brains than control larvae [35]. Past1 \((\text{Putative Achaete Scute Target 1})\) is a plasma membrane-associated protein that is implicated in endocytosis and genetically interacts with Notch [36–38]. CG2794 encodes an essential gene of unknown function \((\text{FBgn}0031265\) from FB2018_02, released 3 April 2018). Ankle2, \textit{past1} and CG2794 have not been implicated in cell movement or regeneration, thus representing novel regulators of these processes.

**Figure 5.** Lineage tracing reveals cell translocation during regeneration in irradiated wing discs. Wing discs were fixed after irradiation with 0R \((-\text{IR})\) or 4000R \((+\text{IR})\) of X-rays and visualized for RFP/GFP. Scale bar in \((a)\) represents 100 μm in \((a–j)\), 10 μm in \((k,l)\) and 25 μm in \((g)\). Images are shown dorsal up and posterior right. \((a–d)\) 30A-GAL4 > UAS-G-trace controls show translocated GFP + RFP + cells after irradiation (within the yellow circle in \((d)\)). \((e,f)\) Co-expression of Wg inhibitor Axin has been shown before to disrupt cell translocation; 30A-GAL4 > UAS-Axin discs from new experiments are included as positive controls. \((g)\) 30A-GAL4 > UAS-Rux disc showing RFP-GFP cells in the pouch (arrow). Note the difference in nuclear size from the hinge cells (arrowhead). \((h,l)\) Irradiated discs from \textit{Ankle2} and \textit{PDSW} RNAi larvae illustrate the range of translocation defects seen in the RNAi lines tested. \((j)\) An un-irradiated disc from \textit{Clatherin heavy chain} RNAi larvae shows disc defects. \((k,l)\) DE-cadherin antibody-stained discs expressing RNAi against \textit{Arp2} \((k)\) or \textit{dia} \((l)\) illustrate the lack of cytokinesis failure; we did not observe bi- or multi-nucleate cells among those expressing RNAi \((\text{RFP}+, \text{arrowheads})\). DE-cadherin antibody marks cell boundaries. \((m)\) The temperature shift protocol used to control GAL4.
salivary gland experiments. Furthermore, the top hits in the wing disc such as Ankle2 and past1 produced no effect in the salivary glands. And the effect of depleting Rab35, a hit in the salivary gland screen, was the weakest among the lines with statistical significance. One possible reason why some of the RNAi lines produced a phenotype in the wing disc but not in the embryonic salivary glands is that the gene product is maternally deposited and the RNAi could not counteract this pool of mRNA. Publicly available data, however, do not support this explanation. Ankle2, for example, is maternally deposited at 111 RPKM (Reads Per Kilobase of transcript, per Million mapped reads) in 0–2-hour-old embryos and its levels declined to 45 RPKM in 8–10-hour-old embryos (stage 12), the relevant stage for salivary gland formation (‘gene_rpkm_report_fb_2017_05.tsv’ downloaded from Flybase on 03/29/2019). The corresponding values in the same dataset for PDSW were 131 and 121 RPKM, respectively. The values for MAPk-AK2 were 92 and 57 RPKM, respectively. Yet, PDSW and MAPk-AK2 RNAi produced salivary gland defects while Ankle2 RNAi did not.

While these results suggest that cell movements during salivary gland development and during wing disc regeneration after IR damage depend on largely distinct genetic requirements, the data also suggest commonalities. For example, three genes identified in the salivary gland experiments (rdgA, Rac2 and MAPk-AK2) did show defects in the wing disc with $p$-values in the $10^{-4}$ range. Collectively, these data indicate that cell movements during salivary gland development and during wing disc regeneration after IR damage have common as well as distinct genetic requirements.

One possible mechanism by which MAPk-AK2, RdgA and Rac2 control salivary gland migration and make a partial contribution to cell movements in the wing disc is through regulation of the actin cytoskeleton. We showed previously that known actin regulators, Rac and Rho GTPases, play essential roles in salivary gland migration and that differential distribution of F-actin is important for gland migration and lumen size control [5–8]. RdgA has been shown to regulate fibroblast migration by reorganizing the actin cytoskeleton through activation of Rac1 and Pak1, which dissociates Rac1 from RhoGDI [39]. In migrating endothelial cells MAPk-AK2 activates LIM-kinase 1 to remodel actin through phosphorylation and inactivation of cofilin [40].

The role of Ena and PDSW in the salivary gland could likewise be through the actin cytoskeleton. Ena and its mammalian orthologue, VASP, have been shown in a number of experimental systems to promote actin-based membrane protrusions and cell motility [41,42]. Ena/VASP promotes G-actin incorporation at the growing ends of actin filaments, although the mechanism is still unclear [43]. Although no direct evidence exists for NADH and mitochondrial proteins in regulation of the actin cytoskeleton, interactions between mitochondria and actin have been documented [44]. Regardless of the mechanism, contributions of Ena and PDSW appear less important in the wing disc. By contrast, RNAi against two known regulators of actin, Dia and Arp2, as well as actin (Act5C), produced partial defects in wing disc regeneration but no effect on salivary gland migration. We conclude that, while actin function is important in both systems, how it is regulated may be different.

Of four RNAi lines found to affect salivary gland migration (ena, PDSW, rdgA and MAPk-AK2), only the one

**Figure 6.** Quantification of cell movements during regeneration after IR. (a) The GFP $+$ RFP $-$ area in the pouch (inside the yellow circle in figure 5d) was quantified in ImageJ and expressed as a percentage of the total GFP $+$ RFP $-$ area in the hinge in each disc. The data are from $+$IR samples, $n = 7$ per genotype. (b) The $p$-values for the dataset in (a) were calculated using a two-tailed $t$-test and are shown along with stock information.

The wing disc assay identified two regulators of the actin cytoskeleton (Arp2, dia) and actin itself (Act5; figure 6). The requirement for actin and its regulators in cell translocation could be due to their direct contribution to cell migration or because actin function is required for the cytokinesis step of cell division, which we know from Rux experiments makes a contribution in this experimental model. Examination of Arp2 or dia RNAi discs, however, did not reveal any evidence of cytokinesis failure (figure 5k,l); RFP $+$ cells that experienced RNAi show discrete nuclei and there is no evidence of bi- or multi-nucleate cells expected from cytokinesis failure (arrowheads). Therefore, we conclude that the role of actin and its regulator in the translocation of hinge cells into the pouch is more likely to be due to their role in cell migration.

### 3. Discussion

In this study, we compared the requirement for collective cell migration during salivary gland development and for cell movements during regeneration of irradiated wing discs, both in *Drosophila*. Of 30 RNAi lines tested in the salivary glands, eight produced defects and included three known (Rac2, Rab35 and Rab40) and four novel regulators. The eighth is Cdk9, whose effects may be indirect. From all the RNAi lines tested in wing disc regeneration, all except two produced statistically significant effects although the magnitude of the effect varied from line to line. The two exceptions are *ena* and *PDSW*, which are precisely two of the eight identified in...
4. Material and methods

4.1. Drosophila techniques

For the analysis of salivary glands, Canton-S flies were used as wild-type controls. ND-PDSW10101, ena02309, ena23, MAPK-AK2G265, rdgA1 and UAS-GFP-NLS were obtained from the Bloomington Stock Center and are described in FlyBase (http://flybase.bio.indiana.edu/). For salivary gland-specific expression of the UAS constructs, we used fork head (fkh)-GAL4. The UAS-GFP-NLS fkh-GAL4 recombinant line was generated using standard genetic techniques.

For the analysis of wing disc, the larvae were raised in Nutri-Fly Bloomington formula food (Genesee Scientific) at room temperature unless otherwise noted. The embryos were collected for 8–24 h at 25°C and reared at room temperature for 72 h from the end of the collection. The vials were monitored regularly for overcrowding (typically seen as dimples in the food) and split to prevent overcrowding. The larvae were then shifted to 29°C for 24 h before irradiation to despress GAL4. The un-irradiated controls remained at 29°C for an additional 38–48 h while the irradiated samples were kept at 29°C for 72 h post-irradiation. For balanced RNAi lines, the absence of CyO balancer-encoded GFP was used to identify the experimental animals.

4.2. Irradiation

The larvae in food were placed in Petri dishes and irradiated in a Faxitron Cabinet X-ray System Model RX-650 (Lincolnshire, IL) at 115 kV and 5.33 rad s⁻¹.

4.3. Immunocytochemistry and in situ hybridization

Embryo fixation and antibody staining were performed as previously described [7]. The following antisera were used at the indicated dilutions: rat dCREB antiserum at 1:10 000; rat antiserum to DE-cadherin at 1:400 (Developmental Studies Hybridoma Bank, DSHB; Iowa City, IA); rabbit aPKC antiserum (Santa Cruz Biotechnology, Dallas, TX) at 1:500; mouse β-galactosidase (β-gal) antiserum (Promega, Madison, WI) at 1:500. Appropriate AlexaFluor 488-, 547- or rhodamine- (Molecular Probes, Eugene, OR) conjugated secondary antibodies were used at a dilution of 1:500 for salivary glands. Anti-mouse-Cy5 secondary antibody was used at 1:200 dilution for wing discs (Jackson, West Grove, PA). Stained embryos were mounted in Aqua Polymount (Polysciences, Inc., Warrington, PA) and thick (1 μm) fluorescence images were acquired on a Zeiss Axioscope microscope (Carl Zeiss) equipped with an LSM 710 for laser scanning confocal microscopy.

To collect wing discs, the larvae were dissected in 1× phosphate-buffered saline (PBS) and fixed in 4% paraformaldehyde for 30 min at room temperature, washed with 1× PBS for 10 min, followed by a 5 min wash with PBTx (0.1% Triton X-100). The wing discs were stained with 10 μg ml⁻¹ Hoechst33342 in PBTx for 2 min, washed three times in PBTx and mounted on glass slides in Fluormount G (SouthernBiotech).

4.4. Image analysis

Salivary gland images were acquired on a Zeiss Axioscope microscope (Carl Zeiss) equipped with confocal laser microscopy.
scanning microscopy (LSM710, Medgar Evers College-CUNY, New York, NY). The number of nuclei surrounding the central lumen and apical domain area and an elongation ratio of salivary gland cells were measured and quantified as previously described [6,7]. Statistical analysis performed in Microsoft Excel using a two-tailed t-test.

Wing disc images were taken on a Leica DMR compound fluorescence microscope using a Q-Imaging R6 CCD camera and Ocular software. The images were assembled, processed and quantified using ImageJ software. Figure 5j,k (100× images) were taken on a PerkinElmer spinning disc confocal attached to a Nikon inverted microscope, using an SDS aAndor iXon Ultra (DU-897) EM CCD camera. For wing discs, we dissected a minimum of 10 larvae per genotype per condition and mounted all discs onto a slide. We then randomly selected and imaged seven intact, flat discs for quantification. To justify the sample size of seven, we used a simplified resource equation from [90]. Briefly, $E = \frac{\text{total number of animals} - \text{total number of groups}}{\text{value of } 10–20}$ is considered adequate. When we compared two groups (vector versus RNAi, for example), six per group or $E = 11$ would be adequate. Therefore, we chose seven.

Data accessibility. This article does not contain any additional data.

Competing interests. We declare we have no competing interests.

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References

1. Mayor R, Etienne-Manneville S. 2016 The front and rear of collective cell migration. Nat. Rev. Mol. Cell Biol. 17, 97 – 109. (doi:10.1038/nrm.2015.14)
2. Bradley PL, Myat MM, Comeaux CA, Andrew DJ. 2003 Posterior migration of the salivary gland requires an intact visceral mesoderm and integrin function. Dev. Biol. 257, 249 – 262. (doi:10.1016/S0012-1606(03)00103-9)
3. Pirraglia C, Myat MM. 2010 Genetic regulation of salivary gland development in Drosophila melanogaster. Front. Oral Biol. 14, 32 – 47. (doi:10.1159/000313706)
4. Maryama R, Andrew D. 2012 Drosophila as model for epithelial tube formation. Dev. Dyn. 241, 119 – 135. (doi:10.1002/dvdy.22775)
5. Pirraglia C, Jattani R, Myat MM. 2006 Rac function in epithelial tube morphogenesis. Dev. Biol. 290, 435 – 446. (doi:10.1016/j.ydbio.2005.12.015)
6. Pirraglia C, Walters I, Ahn N, Myat M. 2013 Rac1 GTPase acts downstream of aPS1bPS integrin to control collective migration and lumen size in the Drosophila salivary gland. Dev. Biol. 377, 21 – 32. (doi:10.1016/j.ydbio.2013.02.020)
7. Xu N, Bagumian G, Galliano M, Myat MM. 2011 Rh GTPase controls Drosophila salivary gland lumen size through regulation of the actin cytoskeleton and Moesin. Development 138, 5415 – 5427. (doi:10.1242/dev.069831)
8. Xu N, Keung B, Myat M. 2008 Rh GTPase controls invagination and cohesive migration of the Drosophila salivary gland through crumbs and Rhokinase. Dev. Biol. 321, 88 – 100. (doi:10.1016/j.ydbio.2008.06.007)
9. Myat MM, Rashmi R, Mannia D, Xu N, Patel U, Galliano M, Zielinski K, Lam A, Welte MA. 2015 Drosophila KASH-domain protein Klarsicht regulates microtubule stability and integrin receptor localization during collective cell migration. Dev. Biol. 407, 103 – 114. (doi:10.1016/j.ydbio.2015.08.003)
10. Jattani R, Patel U, Kerman B, Myat M. 2009 Deficiency screen identifies a novel role for beta 2 tubulin in salivary gland and myoblast migration in the Drosophila embryo. Dev. Dyn. 238, 853 – 863. (doi:10.1002/dvdy.21899)
11. Worley ML, Setlawan L, Hanahan IK. 2012 Regeneration and transdifferentiation in Drosophila imaginal discs. Annu. Rev. Genet. 46, 289 – 310. (doi:10.1146/annurev-genet-110711-155837)
12. Vergheze S, Su TT. 2016 Drosophila Wnt and STAT define apoposis-resistant epithelial cells for tissue regeneration after irradiation. PLoS Biol. 14, e1002536. (doi:10.1371/journal.pbio.1002536)
13. Vergheze S, Su TT. 2017 STAT, wingless, and Nurf-38 determine the accuracy of regeneration after radiation damage in Drosophila. PLoS Genet. 13, e1007055. (doi:10.1371/journal.pgen.1007055)
14. D’Ambrosio MV, Vale RD. 2010 A whole genome RNAi screen for Drosophila S2 cell spreading performed using automated computational image analysis. J. Cell Biol. 191, 471 – 478. (doi:10.1083/jcb.201003315)
15. Kiger AA, Baum B, Jones S, Jones MR, Coulson A, Echeverri C, Perrimon N. 2003 A functional genomic analysis of cell morphology using RNA interference. J. Biol. 2, 27. (doi:10.1186/1475-2924-2-27)
16. Liu T, Sims D, Baum B. 2009 Parallel RNAi screens across different cell lines identify genetic and cell type-specific regulators of actin organization and cell morphology. Genome Biol. 10, R26. (doi:10.1186/gb-2009-10-3-r26)
17. Rohn JL et al. 2011 Comparative RNAi screening identifies a conserved core metazoan actinome by phenotype. J. Cell Biol. 194, 789 – 805. (doi:10.1083/jcb.20113168)
18. Eisenberg JC, Shilatifard A, Donokhov N, Michener DE. 2007 Cdks is an essential kinase in Drosophila that is required for heat shock gene expression, histone methylation and elongation factor recruitment. Mol. Genet. Genomics 277, 101 – 114. (doi:10.1007/s00438-006-0164-2)
19. Andrew DJ, Henderson KD, Seshaih P. 2000 Salivary gland development in Drosophila melanogaster. Mech. Dev. 92, 5 – 17. (doi:10.1016/S0925-4773(99)00321-4)
20. Ahern-Djamali SM, Comer AR, Bachmann C, Kastenmeier AS, Reddy SK, Beckerle MC, Walter U, Hoffmann FM. 1998 Mutations in Drosophila enable and rescue by human vasodilator-stimulated phosphoprotein (VASP) indicate important functional roles for Ena/VASP homology domain 1 (EHV1) and EHV2 domains. Mol. Biol. Cell. 9, 2157 – 2171. (doi:10.1091/molc.9.8.2157)
21. Xu N, Myat MM. 2012 Coordinated control of lumen size and collective migration in the salivary gland. Fly (Austin) 6, 142 – 146. (doi:10.4161/fly.20246)
22. Masai I, Okazaki A, Hosoya T, Hotta Y. 1993 Drosophila retinal degeneration A gene encodes an eye-specific diacylglycerol kinase with cysteine-rich zinc-finger motifs and ankyrin repeats. Proc. Natl Acad. Sci. USA 90, 1157 – 1161. (doi:10.1073/pnas.90.23.11157)
23. Bryant PJ. 1970 Cell lineage relationships in the imaginal wing disc of Drosophila melanogaster. Dev. Biol. 22, 389 – 411. (doi:10.1016/0012-1606(70)90160-0)
24. Haynie JL, Bryant PJ. 1977 The effects of X-rays on the proliferation dynamics of cells in the imaginal wing disc of Drosophila melanogaster. Wilhelm Roux’s Arch. Dev. Biol. 183, 85 – 100. (doi:10.1007/BF00488779)
25. James AA, Bryant PJ. 1981 A quantitative study of cell death and mitotic inhibition in gamma-irradiated imaginal wing discs of Drosophila melanogaster. Radiat. Res. 87, 552 – 564. (doi:10.2307/3575520)
26. Herrera SC, Martin R, Morata G. 2013 Tissue homeostasis in the wing disc of Drosophila melanogaster: immediate response to massive damage during development. PLoS Genet. 9, e1003446. (doi:10.1371/journal.pgen.1003446)
27. Evans CJ et al. 2009 G-TRACE: rapid Gal4-based cell lineage analysis in Drosophila. Nat. Methods 6, 603 – 605. (doi:10.1038/nmeth.1356)

royalsocietypublishing.org/journal/rsob
28. Verghese S, Su TT. 2018 Ionizing radiation induces stem cell-like properties in a caspase-dependent manner in Drosophila. PLoS Genet. 14, e1007659. (doi:10.1371/journal.pgen.1007659)
29. Schniedelin J et al. 2012 Fiji: an open-source platform for biological-image analysis. Nat. Methods 9, 676 – 682. (doi:10.1038/nmeth.2019)
30. Baena-Lopez LA, Baonza A, Garcia-Bellido A. 2005 The orientation of cell divisions determines the shape of Drosophila organs. Curr. Biol. 15, 1640 – 1644. (doi:10.1016/j.cub.2005.07.062)
31. Zhou Z, Alegot H, Irvine KD. 2019 Oriented cell divisions are not required for Drosophila wing shape. Curr. Biol. 29, 856 – 864. (doi:10.1016/j.cub.2019.01.044)
32. Li W, Kale A, Baker NE. 2009 Oriented cell division as a response to cell death and cell competition. Curr. Biol. 19, 1821 – 1826. (doi:10.1016/j.cub.2009.09.023)
33. Foley E, Sprenger F. 2001 The cyclin-dependent kinase inhibitor Roughex is involved in mitotic exit of Drosophila. Curr. Biol. 11, 151 – 160. (doi:10.1016/S0960-9822(01)00505-1)
34. Sprenger F, Yakovlevich N, D’Farrell PH. 1997 S-phase function of Drosophila cyclin A and its downregulation in G1 phase. Curr. Biol. 7, 488 – 499. (doi:10.1016/S0960-9822(06)00220-X)
35. Yamamoto S et al. 2014 A Drosophila genetic resource of mutants to study mechanisms underlying human genetic diseases. Cell 159, 200 – 214. (doi:10.1016/j.cell.2014.09.002)
36. Donot O, Steller H, Segal D, Horowitz M. 2017 Past1 modulates Drosophila eye development. PLoS One 12, e0169639. (doi:10.1371/journal.pone.0169639)
37. Koles K, Messelaar EM, Feiger Z, Yu CJ, Frank CA, Rudal AA. 2015 The EHD protein Past1 controls postsynaptic membrane elaboration and synaptic function. Mol. Biol. Cell. 26, 3275 – 3288. (doi:10.1091/mbc.E15-02-0093)
38. Olsowka-Kutz Y, Gertel Y, Benjamin S, Sela O, Pekar A, Arama E, Steller H, Horowitz M, Segal D. 2009 Drosophila Past1 is involved in endocytosis and is required for germline development and survival of the adult fly. J. Cell Sci. 122(Pt 4), 471 – 480. (doi:10.1242/jcs.038521)
39. Abramovici H, Mojtahabi P, Parks RJ, Zhong XP, Koretzky GA, Topham MK, Gee SH. 2009 Diacylglycerol kinase zeta regulates actin cytoskeleton reorganization through dissociation of Rac1 from RhoGDI. Mol. Biol. Cell 20, 2049 – 2059. (doi:10.1091/mbc.e07-12-1248)
40. Kobayashi M, Nishita M, Mishima T, Ohashi K, Mizuno K. 2006 MAPKAPK-2-mediated LIM-kinase activation is critical for VEGF-induced actin remodeling and cell migration. Embo J. 25, 713 – 726. (doi:10.1038/sj.emboj.7600973)
41. Krause M, Gautreau A. 2014 Steering cell migration: lamellipodium dynamics and the regulation of directional persistence. Nat. Rev. Mol. Cell. Biol. 15, 577 – 590. (doi:10.1038/nrm3861)
42. Trichet L, Sykes C, plastino J. 2008 Relaxing the actin cytoskeleton for adhesion and movement with Ena/WASP. J. Cell Biol. 181, 19 – 25. (doi:10.1083/jcb.200710168)
43. Bear JE, Gertler FB. 2009 Ena/WASP: towards resolving a pointed controversy at the barbed end. J. Cell Sci. 122(Pt 12), 1947 – 1953. (doi:10.1242/jcs.038125)
44. Boldogh IR, Pon LA. 2006 Interactions of mitochondria with the actin cytoskeleton. Biochim. Biophys. Acta 1763, 450 – 462. (doi:10.1016/j.bbamcr.2006.02.014)
45. Molinie N, Gautreau A. 2018 The Arp2/3 regulatory system and its deregulation in cancer. Physiol. Rev. 98, 215 – 238. (doi:10.1152/physrev.00006.2017)
46. Hudson AM, Coaley L. 2002 A subset of dynamic actin rearrangements in Drosophila requires the Arp2/3 complex. J. Cell Biol. 156, 677 – 687. (doi:10.1083/jcb.200109065)
47. Lesch C, Jo J, Wu Y, Fish G, Glako M. 2010 A targeted UAS-RNAi screen in Drosophila larval identifies wound closure genes regulating distinct cellular processes. Genetics 186, 943 – 957.
48. Rogers KK, Jou T-S, Guo W, Lipszutz JH. 2003 The Rho family of small GTPases in involved in epithelial cystogenesis and tubulogenesis. Kidney Int. 63, 1632 – 1644. (doi:10.1046/j.1523-1755.2003.00962.x)
49. Castillion DH, Wasserman SA. 1994 Diaphanous is required for cytokinesis in Drosophila and shares domains of similarity with the products of the limb deformity gene. Development 120, 3367 – 3377.
50. Chan J, Kanathara ND. 2013 How to calculate sample size in animal studies? J. Pharmacol. Pharmacother. 4, 303 – 306. (doi:10.4103/0976-500X.119726)
51. Dietzl G et al. 2007 A genome-wide transgenic RNAi library for conditional gene inactivation in Drosophila. Nature 448, 151 – 156. (doi:10.1038/ nature05954)