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Retromer Controls Planar Polarity Protein Levels and Asymmetric Localization at Intercellular Junctions

Highlights
- Retromer regulates levels and asymmetry of Flamingo and Strabismus in the pupal wing
- Retromer regulates Flamingo and Strabismus levels independently of the Wash complex
- The cargo adaptor Snx27 interacts directly with the PDZ-binding motif of Flamingo
- Loss of Snx27 disrupts Flamingo trafficking

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In Brief
Planar cell polarity—the organization of cells within a two-dimensional plane—is an important feature in tissue organization. Strutt et al. show that, in the Drosophila wing, endosomal recycling through retromer and its cargo adaptor Snx27 mediates the cell-surface localization of two key planar polarity proteins, Flamingo and Strabismus.
Retromer Controls Planar Polarity Protein Levels and Asymmetric Localization at Intercellular Junctions

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SUMMARY

The coordinated polarization of cells in the plane of a tissue, termed planar polarity, is a characteristic feature of epithelial tissues [1]. In the fly wing, trichome positioning is dependent on the core planar polarity proteins adopting asymmetric subcellular localizations at apical junctions, where they form intercellular complexes that link neighboring cells [1–3]. Specifically, the seven-pass transmembrane protein Flamingo and the cytoplasmic proteins Dishevelled and Diego localize to distal cell ends, the four-pass transmembrane protein Strabismus and the cytoplasmic protein Prickle localize proximally, and the seven-pass transmembrane spanning atypical cadherin Flamingo localizes both proximally and distally. To establish asymmetry, these core proteins are sorted from an initially uniform distribution; however, the mechanisms underlying this polarized trafficking remain poorly understood. Here, we describe the identification of retromer, a master controller of endosomal recycling [4–6], as a key component regulating core planar polarity protein localization in Drosophila. Through generation of mutants, we verify that loss of the retromer-associated Snx27 cargo adaptor, but notably not components of the Wash complex, reduces junctional levels of the core proteins Flamingo and Strabismus in the developing wing. We establish that Snx27 directly associates with Flamingo via its C-terminal PDZ binding motif, and we show that Snx27 is essential for normal Flamingo trafficking. We conclude that Wash-independent retromer function and the Snx27 cargo adaptor are important components in the endosomal recycling of Flamingo and Strabismus back to the plasma membrane and thus contribute to the establishment and maintenance of planar polarization.

RESULTS AND DISCUSSION

The endosomal sorting of internalized cargo for recycling to the cell surface is a highly regulated process [7]. A master conductor for the recycling of numerous cargoes is retromer, a stable heterotrimer of VPS29, VPS35, and VPS26 [4, 5]. To examine the role of retromer in the establishment and maintenance of planar polarization (Figures 1A and 1B), we performed genetic analysis in Drosophila. As null mutants of the retromer subunit Vps35 are lethal during late larval or early pupal stages [8, 9], we generated Vps35-null mutant clones in the pupal wing. This revealed that functional levels of the transmembrane core proteins Flamingo (Fmi), Strabismus (Stbm), and Frizzled (Fz) were decreased within Vps35-null clones (Figures 1C, 1D, and S1A–S1E). Levels of the cytoplasmic core protein Dishevelled (Dsh) were also slightly reduced, and levels of Prickle (Pk) and other junctional proteins, such as Armadillo, were unaffected (Figures S1B–S1E). In contrast, loss of the retromer-linked SNX-BAR proteins, Snx1 or Snx6, or the core component of the retriever complex C16orf62 (data not shown) did not affect levels of Fmi or Stbm (Figures S1F–S1G). Notably, both core protein asymmetry and polarity coordination between cells (Figures 1E and S1F) were reduced in Vps35 mutant tissue, accompanied by a delay in trichome initiation (Figure 1F). By revealing a role for retromer in regulating the cell surface levels and asymmetry of core planar polarity proteins in the pupal wing, these data extend the role of retromer in specifying polarity through recycling of Crumbs [10, 11] and the Scribble polarity module [12].

In mammalian cells, retromer function is coupled to the actin-polymerizing Wiskott-Aldrich syndrome and SCAR homolog (WASH) complex, a pentameric assembly of WASH (WASHC1), FAM21 (WASHC2), CCDC53 (WASHC3), SWIP (WASHC4), and Strumpellin (WASHC5) [13, 14]. Within this complex, WASH stimulates the ARP2/3 complex to drive polymerization of branched actin networks that aid the organization of endosomal retrieval sub-domains [7]. We used CRISPR/Cas9 editing to generate a Drosophila Fam21-null mutant (see Figure S1J; STAR Methods). Homozygous Fam21-null flies were viable and fertile. In Fam21-null mutant clones in the pupal wing, we observed no effect on the levels of Fmi and Stbm (Figures 1D and 1G) nor was any effect observed in wash-null mutant clones (Figures 1D and 1H) [15]. Therefore, in contrast to Drosophila trachea development,
Figure 1. Vps35 Regulates Levels of Fmi and Stbm at Apical Junctions Independently of the Wash Complex

(A) Diagram illustrating asymmetric localization of the core planar polarity proteins in the pupal wing. Two cells are shown, with Fmi, Fz, Dsh, and Dgo localizing on distal cell edges. This forms an intercellular complex with Fmi, Stbm, and Pk on proximal edges of the neighboring cell.

(B) During polarization, complexes sort from a uniform distribution (left), and all the complexes become oriented in the same direction (right). This specifies positioning of trichomes (black in right diagram) to distal cell edges.

(C, F, G, and H) 28-hr after puparium formation (APF) (C, G, and H) or 32-hr APF (F) pupal wings carrying clones of Vps35 (C and F), Fam21 (G), or wash (H), marked by loss of β-gal staining (blue in C, G, and H and green in F). Wings are immunolabeled for Fmi in green and Stbm in red (C, G, and H) or Fmi in blue and phalloidin in red (F). The reduced phalloidin staining in Vps35 mutant tissue (F) indicates a delay in trichome initiation. In older wings, phalloidin-stained trichomes are visible in Vps35 clones (not shown). Scale bar 10 μm.

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where retromer and Wash work together [16, 17], the retromer-mediated asymmetry of Fmi and Stbm in the pupal wing occurs independently of the Wash complex.

A global proteomic analysis, performed in vitro and in non-polarized cultured cells, identified an association between CELSR1 and VANGL1/VANGL2, mammalian equivalents of Drosophila Fmi and Stbm, respectively, with the retromer cargo adaptor sorting nexin 27 (SNX27) [18]. SNX27 directly associates with the VPS26 subunit of retromer through a mechanism that is conserved in Drosophila [19] and promotes endosomal recycling to the plasma membrane. SNX27 contains an amino-terminal PDZ domain that binds to cargoes that contain a C-terminal PDZ binding motif [18, 20]. Drosophila Fmi and Stbm both contain PDZ binding motifs (Figure 2A), and they were both able to associate with human SNX27 and Drosophila Snx27 in a PDZ-binding-motif-dependent manner (Figures 2B–2E).

To establish the role of Drosophila Snx27 in retromer-mediated sorting of Fmi and Stbm, we generated a Snx27 loss-of-function mutant (see Figure S2A; STAR Methods). Snx27-null flies are homozygous viable and fertile. As in Vps35 mutant clones, junctional levels of both Fmi and Stbm were reduced in Snx27 pupal wing clones (Figures 3A and 3B). Western blotting revealed that total levels of Fmi were reduced in Snx27 mutant pupal wings (Figures S2B and S2C), consistent with increased Fmi degradation in the absence of Snx27-retromer-mediated recycling. Levels of Fz at apical junctions were also slightly reduced, and levels of other junctional proteins (Pk, Dsh, and Armadillo) were unaffected (Figures S2D–S2H). Despite the reduced protein levels, core protein asymmetry and trichome polarization correctly. The reduced junctional levels of Fmi and Stbm were fully rescued by expression of a rescue transgene in non-Snx27 mutant tissue (Figures S2I and S2J), confirming that these effects are due to loss of Snx27.

Interestingly, Stbm levels appeared more greatly reduced in Vps35 clones than in Snx27 clones, but Fmi levels were similar in each (compare Figures 1D and 3B). This was confirmed by making overlapping clones of Vps35 and Snx27 in the same wings (Figures S3A, S3B, and S3E). This suggests that, although retromer acts together with Snx27 to regulate Fmi and Stbm levels, Vps35 also acts on Stbm independently of Snx27. Furthermore, this might explain why reduced asymmetry is seen in Vps35 mutants, but not in Snx27 mutants. This Snx27-independent effect of Vps35 is likely to be a retromer-dependent function, as Stbm levels are also decreased more than Fmi levels in Vps26 mutant tissue (Figures S3G and S3H). Interestingly, Fmi also accumulated in intracellular puncta in Vps26 mutant tissue (Figure S3G). Simultaneous loss of Snx27 and Vps35 did not cause a further decrease in levels of Fmi and Stbm, over loss of Vps35 alone (Figures S3C, S3D, and S3F). This suggests that Snx27 acts only through retromer in regulating Fmi and Stbm recycling.

As Snx27 interacts with the PDZ binding motifs of Fmi and Stbm and is required for their retromer-mediated recycling, we examined whether the PDZ binding motifs of these proteins were necessary for their correct junctional levels. First, we generated flies carrying P[acman] rescue constructs, in which either full-length Stbm or Stbm lacking the PDZ binding motif was tagged with EGFP at the N terminus. We then generated pupal wings in which clones of EGFP-Stbm were juxtaposed to clones of EGFP-StbmaΔPDZ binding motif, in a stbm mutant background. No difference in junctional levels of Stbm was seen (Figures S4A and S4B), suggesting that the PDZ binding motif of Stbm is not required for maintenance of Stbm junctional levels and by extension that Snx27 cannot be regulating Stbm levels via a direct interaction between the PDZ binding motif and the PDZ domain. To confirm this, clones of Snx27 were induced in tissue expressing either EGFP-Stbm or EGFP-StbmaΔPDZ binding motif. In both cases, junctional levels were reduced (Figures S4C–S4E), suggesting that the effect of Snx27 on Stbm is independent of the PDZ binding motif.

Next, we generated flies in which the Fmi was tagged with EGFP in its extracellular domain, expressed under the armadillo promoter. Clones of EGFP-Fmi adjacent to clones of EGFP-FmiΔPDZ binding motif were then created, in a fmi mutant background. Deletion of the PDZ binding motif caused a reduction in junctional levels of Fmi (Figures 3E and 3F). Lower levels of Stbm were also observed at junctions in EGFP-FmiΔPDZ binding motif clones (Figures 3E and 3F), consistent with the previous observation that Fmi is required for normal Stbm levels at the plasma membrane [21] and suggesting that Stbm levels decrease in Snx27 clones as a consequence of a decrease in Fmi levels. Deletion of the PDZ binding motif of Fmi, although lowering junctional levels of Fmi, did not reduce overall asymmetry (Figure 3G), similar to what is observed in Snx27 mutants and also in flies with reduced fmi gene dosage [22].

The similar phenotypes observed upon loss of Snx27 and loss of the Fmi PDZ binding motif are consistent with Fmi interacting directly with Snx27 via the PDZ domain of Snx27. Confirming this, we found that EGFP-FmiΔPDZ binding motif was insensitive to loss of Snx27: although junctional levels of EGFP-Fmi were reduced in Snx27 clones (Figures 3H and 3I), levels of EGFP-FmiΔPDZ binding motif did not decrease any further (Figures 3H and 3J).

These data are consistent with the reduced junctional levels of Fmi being caused by reduced recycling, but it is also possible that there is less delivery of newly synthesized Fmi. To distinguish these possibilities, we analyzed Fmi protein dynamics. First, in an antibody internalization experiment, live prepupal

(C) High magnification image of wild-type and mutant regions immunolabeled with Fmi and used to quantitate polarity.

(D) Quantitation of mean intensity of Fmi (red dots) or Stbm (orange dots) membrane labeling in pupal wing clones. Intensity is shown as a ratio of signal in mutant versus wild-type tissue (see C'). Values from the same wing are linked by black bars; mean and SD are listed. Paired t tests were used to compare values in the same wing. **p < 0.001. See also Figure S1 and Data S1 for all statistical comparisons.
wings carrying Snx27 clones were labeled with a pulse of Fmi antibody and then internalization was time resolved after antibody wash off. As expected, extracellular Fmi labeling revealed less junctional Fmi in Snx27 tissue compared to wild-type (0.64 ± 0.10; p < 0.001) at the start of the experiment (Figure 4A). In both wild-type and Snx27 tissue, extracellular Fmi levels decreased over time (Figures 4A–4C and 4G). This was accompanied by the appearance of Fmi in intracellular puncta, which we previously showed were endosomal (Figures 4D–4F) [23]. Interestingly, at 10 min internalization, a similar number of puncta

Figure 2. Drosophila Snx27 Interacts with the PDZ Binding Motifs Present in the C Termini of Fmi and Stbm
(A) Alignment of the C-terminal PDZ binding motifs of Drosophila Fmi and Stbm with the optimized sequence for PDZ binding motif recognition by the PDZ domain of SNX27.
(B–E) GFP-nanotrap immunoprecipitations of HEK293T cells transiently transfected with constructs encoding for GFP-tagged tail regions of Fmi and Stbm (GFP-Fmi wild-type [WT], B and D, and GFP-Stbm WT, C and E) or the corresponding constructs deleted for the last three amino acids of the PDZ binding motif (GFP-FmiΔPDZbm, B and D, and GFP-StbmΔPDZbm, C and E).
(B and C) Samples immunoblotted for the presence of endogenous human SNX27.
(D and E) Cells co-transfected with mCherry-tagged Drosophila Snx27 and samples immunoblotted with anti-mCherry. Data are representative of similar data derived from three independent biological replicates.
Figure 3. Snx27 Regulates Junctional Levels of Fmi and Stbm via the PDZ Binding Motif of Fmi

(A and D) 28-hr APF (A) or 32-hr APF (D) pupal wings carrying clones of Snx27 (Figure S2A), marked by loss of GFP immunolabeling (green in A) or RFP fluorescence (red in D). Wings are immunolabeled for Fmi in red and Stbm in blue (A) or Fmi in blue and phalloidin in green (D). Scale bar 10 μm.

(B) Quantitation of mean intensity of Fmi (red dots) or Stbm (orange dots) membrane labeling in pupal wing clones of Snx27. Intensity is shown as a ratio of signal in Snx27 mutant compared to wild-type in each wing.

(C) Mean polarity and variation in polarity angle of wings immunolabeled for Fmi in wild-type and Snx27 mutant tissue.

(E) 28-hr APF pupal wing with twin clones of arm-PRO-EGFP-fmi next to arm-PRO-EGFP-fmi ΔPDZ binding motif (ΔPDZbm), marked by β-gal immunolabeling in blue, in a fmiE59/fmiE45 mutant background. The wing is immunolabeled for EGFP in green and Stbm in red.

(F) Quantitation of mean intensity of membrane labeling of EGFP-Fmi (red dots) and Stbm (orange dots). Intensity is shown as a ratio of signal in ΔPDZbm compared to full-length protein in each wing.

(legend continued on next page)
were observed in Snx27 and wild-type, and puncta were larger in Snx27 (Figures 4E and 4H). Because less extracellular Fmi labeling was observed in Snx27 tissue than wild-type (Figure 4A), and internalization rates were similar (Figure 4G), it follows that endosomal puncta are more persistent in Snx27 than in wild-type. The number of puncta in Snx27 relative to wild-type was reduced at 30 min internalization, and puncta size was similar (Figures 4F and 4H). This is consistent with a proportion of Fmi being rapidly recycled to the plasma membrane in wild-type, but in Snx27, Fmi persists in endosomes and is subsequently degraded.

Fluorescence recovery after photobleaching (FRAP) experiments using EGFP-fmi knockin flies provide a further confirmation of altered Fmi dynamics in Snx27 wings. Recovery of Fmi levels was quantified by fluorescence intensity measurement (Figure 4G). The mean intensity of EGFP-Fmi was significantly lower in Snx27 tissue than in wild-type at 0 min after antibody internalization (Figure 4G), indicating reduced Fmi levels in Snx27 tissue.

Intracellular Fmi puncta size and number were also quantified using FRAP experiments (Figure 4H). The number of puncta and the size of puncta were reduced in Snx27 compared to wild-type, consistent with the reduced Fmi levels observed in Snx27 tissue.

Figure 4. Loss of Snx27 Disrupts Intracellular Trafficking of Fmi

(A–F) Images of Fmi internalization experiment. (A–C) Extracellular Fmi immunolabeling of 5.5 hr APF prepupal wings, imaged at the apical junctions. (D–F) Total Fmi immunolabeling of the same wings, imaged 1.2 μm below the apical junctions. Images at 0 min (A and D), 10 min (B and E), or 30 min (C and F) after removal of Fmi antibody, in wild-type (A–F) or Snx27 (A’–F’) mutant regions of the same wings are shown. Scale bar 5 μm. (G and H) Quantification of extracellular Fmi immunolabeling (G) or Fmi intracellular puncta size and number (H) in wild-type and Snx27 tissue, after antibody internalization. (G) Intensity at 0 min is normalized to 1.0 for each genotype (but note that Fmi levels in Snx27 tissue were 64% of levels in wild-type). Error bars are SD; ANOVA with Dunnett’s multiple comparisons test was used to compare intensities to 0 min; ***p < 0.001; **p < 0.01. Immunolabeling is more punctate after internalization, as Fmi that is highly clustered is stable, and non-clustered Fmi is rapidly internalized (see [23]). (H) Data are shown as a ratio of puncta size or number in Snx27 mutant compared to wild-type in each wing. Error bars are SD. One-sample t tests were used to determine whether the ratio differed from 1.0; ***p < 0.001; **p < 0.01. Note that puncta intensity was similar in wild-type and Snx27 tissue.

(I) FRAP of EGFP-Fmi, in 5.5-hr APF prepupal wings from wild-type or Snx27 flies. Note that pre-bleach levels of EGFP-Fmi in Snx27 tissue were decreased relative to wild-type (0.56 ± 0.13; p < 0.001), but data on the graph were normalized to a pre-bleach intensity of 1.0 for each genotype. A two-phase exponential curve was fitted; error bars are SD. Recovery in the first phase is not significantly different, but there is proportionally less recovery in the second phase in Snx27 compared to wild-type (p = 0.006). Recovery of neither genotype reaches a plateau in the time of imaging.

See Data S1 and Table S1 for all statistical comparisons.
EGFP-Fmi fluorescence after bleaching fitted to a two-phase exponential curve, where the fast phase of recovery was similar in wild-type and Snx27 wings and the slow recovery was reduced in Snx27 compared to wild-type (Figure 4f; Table S1). These phases likely correspond to recovery by diffusion and endocytosis and recycling, respectively (compare to [24]) and thus suggest reduced recovery via endosomal trafficking in Snx27 wings.

In the present study, we provide biochemical and genetic evidence to establish that retromer and its associated cargo adaptor Snx27 are important components in the in vivo trafficking of the core planar polarity proteins Fmi and Stbm in the Drosophila pupal wing. As far as we are aware, this is the first in vivo demonstration of a functional coupling between the core planar polarity cargo and an endosomal recycling complex. Interestingly, in humans, planar polarity mutations have been shown to contribute to the etiology of Robinow syndrome, a severe skeletal dysplasia characterized by short limbs and craniofacial anomalies [25–31]. It is intriguing to note that the Snx27−/− mouse phenotype is characterized not only by neurological defects but also skeletal dysplasia, including shortened forelimbs and cranial defects [32], which could be attributed to a defect in the robustness of planar polarization.

STAR METHODS

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SUPPLEMENTAL INFORMATION

Supplemental Information includes four figures, one table, and two data files and can be found with this article online at https://doi.org/10.1016/j.cub.2018.12.027.

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AUTHOR CONTRIBUTIONS

Conceptualization, P.J.C. and D.S.; Reagent Generation, Acquisition, and Analysis of Data, H.S., P.F.L., N.P., and K.J.M.; Writing – First Draft, P.J.C., H.S., and N.P.; Writing – Review and Editing, H.S., P.F.L., N.P., K.J.M., D.S., and P.J.C.; Funding Acquisition, Resources, and Supervision, D.S. and P.J.C.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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# STAR★METHODS

## KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| Antibodies          |        |            |
| mouse monoclonal anti-Fmi #74 (immunolabelling of fixed tissues) | DSHB [33] | RRID: AB_2619583 |
| mouse monoclonal anti-Fmi #71 (antibody internalisation) | [33] | n/a |
| rat anti-Stbm       | [34]   | n/a       |
| affinity purified rabbit anti-Fz | [35] | n/a |
| affinity purified rat anti-Pk | [36] | n/a |
| rat anti-Dsh        | [37]   | n/a       |
| mouse monoclonal anti-Arm | DSHB | RRID: AB_528089 |
| mouse monoclonal anti-β-gal 40-1a | DSHB | RRID: AB_2314509 |
| rabbit anti β-gal   | MP Biochemicals/Cappel | cat# 0855976 (old cat# 55976); RRID: AB_2334934 |
| affinity purified rabbit anti-GFP | Abcam | cat# ab6556; RRID: AB_305664 |
| mouse monoclonal anti-SNX27 1C6 | Abcam | cat# AB77799; RRID: AB_10673818 |
| mouse monoclonal anti-GFP | Roche | cat# 11814460001; RRID: AB_390913 |
| mouse monoclonal anti-mCherry | Abcam | cat# AB125096; RRID: AB_11133266 |
| mouse monoclonal anti-Actin AC40 | Sigma | cat# A7400; RRID: AB_476730 |
| Alexa Fluor 680 goat anti-mouse IgG | Thermo-Fisher | cat# A21057; RRID: AB_2535723 |
| Alexa Fluor 800 goat anti-rabbit IgG | Thermo-Fisher | cat# SA535571; RRID: AB_2556775 |
| HRP conjugated goat anti Mouse | DAKO | cat# P0447; RRID: AB_2617137 |
| Chemicals, Peptides, and Recombinant Proteins |        |            |
| Alexa Fluor 568 conjugated Phalloidin | Invitrogen | cat# A12380 |
| FITC conjugated Phalloidin | Molecular Probes | cat# F-432 |
| DMEM                 | Sigma   | cat# D5796 |
| Foetal bovine serum for use with DMEM | Sigma | cat# F7524 |
| GFP trap beads       | Chromotek | cat# gta-20 |
| SuperSignal West Dura Extended duration substrate | Thermo-Fisher | cat# 34075 |
| 16% paraformaldehyde solution (methanol free) | Agar Scientific | cat# R1026 |
| Triton X-100         | VWR     | cat# 28817.295; CAS: 9002-93-1 |
| Normal goat serum    | Jackson Labs | cat# 005-000-121 |
| Glycerol             | VWR     | cat# 284546F; CAS: 56-81-5 |
| DABCO                | Fluka   | cat# 33480; CAS: 280-57-9 |
| Schneider’s medium   | Thermo-Fisher | cat# 21720 |
| Foetal bovine serum for use with Schneider’s medium | Sigma | cat# F4135 |
| Mowiol               | Polysciences | cat# 17951; CAS: 9002-89-5 |
| Methyl cellulose     | Sigma   | cat# 274429; CAS: 9004-67-5 |
| Experimental Models: Cell Lines |        |            |
| HEK293               | ATCC    | CRL-11268  |
| Experimental Models: Organisms/Strains |        |            |
| Vps39Δ [20]          | FlyBase: FBal0221635        |
| Snx1Δ [22]           | FlyBase: FBal0336681        |

(Continued on next page)
**CONTACT FOR REAGENT AND RESOURCE SHARING**

Further information and requests for resources and reagents should be directed to and will be fulfilled by Peter Cullen (Pete.Cullen@bristol.ac.uk).

**EXPERIMENTAL MODEL AND SUBJECT DETAILS**

HEK293T cells were maintained in DMEM (Sigma, cat# D5796) supplemented with 10% (v/v) fetal bovine serum (Sigma, cat# F7524) under standard conditions. *Drosophila melanogaster* flies were grown on standard cornmeal/agar/molasses media at 18°C or 25°C, in plastic vials in a controlled humidity environment, on a 12 hr/12 hr light-dark cycle. For pupal wing dissections, pupae were aged for 28 hr after...
puparium formation (APF) at 25°C, or for 32 hr APF for trichome staining. For experiments using prepupal wings, pupae were aged for 5.5 hr APF at 25°C. Sex of pupae was not determined, unless stated below. The health/immune status cannot be determined for individual pupae. Pupae were not subjected to previous procedures and were drug and test naive.

Fly strains are described in the Key Resources Table. stbm6, fmiE59, fmiE45, Vps35MH20, wash185, Snx112 and Snx61 are putative null mutations. Vps26C is a point mutation in a highly conserved residue in the arrestin-like domain. Mitotic clones were generated using the FLP-FRT system and either Ubx-FLP or hs-FLP.

**Genotypes of experimental models**

**Figure 1**

(C-F) y w Ubx-FLP; FRT42 Vps35MH20 / FRT42 arm-lacZ
(D, G) y w Ubx-FLP; FRT42 Fam21KO / FRT42 arm-lacZ
(D, H) y w Ubx-FLP; FRT42 wash185 / FRT42 arm-lacZ

**Figure 3**

(A-D) y w Snx27KO FRT19A / y w ubi-GFP FRT19A; Ubx-FLP/+ (females)
(E-G) y w Ubx-FLP; arm-PRO-EGFP-fmi FRT40 fmiE59 / arm-PRO-EGFP-fmi.DPDZbm arm-lacZ FRT40 fmiE45
(H, I) y w Snx27KO FRT19A / Ubi-RFP-nls, w hs-FLP FRT19A; arm-PRO-EGFP-fmi.DPDZbm FRT42 fmiE59 / + (females)

**Figure 4**

(A-H) y w Snx27KO FRT19A / Ubi-RFP-nls, w hs-FLP FRT19A; Ubx-FLP / + (females)
(I) w1118; EGFP-fmi / + (males) and y w Snx27KO FRT19A; EGFP-fmi / + (males)

**Figure S1**

(A-F) w1118 (males) and y w Snx27KO FRT19A (males)
(G, H) y w Snx27KO FRT19A / y w ubi-GFP FRT19A; Ubx-FLP/+ (females)
(I, J) y w Snx27KO FRT19A / Ubi-RFP-nls, w hs-FLP FRT19A; tub-GFP-Snx27 / Ubx-FLP (females)

**Figure S3**

(A-F) y w Snx27KO FRT19A / Ubi-RFP-nls, w hs-FLP FRT19A; FRT42 Vps35MH20 / FRT42 arm-lacZ (females)
(G, H) y Vps26C w FRT19A / Ubi-RFP-nls, w hs-FLP FRT19A; FRT42 arm-lacZ (females)

**Figure S4**

(A, B) y w Ubx-FLP; P[acman]-EGFP-stbm arm-lacZ FRT40 stbm6 / P[acman]-EGFP-stbm.DPDZbm FRT40 stbm6
(C, E) y w Snx27KO FRT19A / Ubi-RFP-nls, w hs-FLP FRT19A; P[acman]-EGFP-stbm FRT40 stbm6 (females)
(D, E) y w Snx27KO FRT19A / Ubi-RFP-nls, w hs-FLP FRT19A; P[acman]-EGFP-stbm.DPDZbm FRT40 stbm6 (females)

**METHOD DETAILS**

**Cell culture and western blots**

pEGFP-C1-Stbm and pEGFP-C1-Stbm.DPDZbm were generated by amplifying the Stbm C-terminal intracellular tail (encoding the last 285 amino acids for the full-length tail) from genomic DNA (from w1118 flies) and cloning into the BglII and EcoRI sites of pEGFP-C1 (Clontech). pEGFP-C1-Fmi and pEGFP-C1-Fmi.DPDZbm were made by amplifying the C-terminal intracellular tail (encoding the last 527 amino acids for the full-length tail) of Fmi from cDNA and cloning into the Xhol and EcoRI sites of pEGFP-C1. pmCherry-C1-Snx27 was made by amplifying Snx27-RA from cDNA and cloning into the NotI and BamHI sites of pmCherry-C1 (Clontech).

Cells were transfected or transduced with the desired GFP-tagged or mCherry-tagged constructs and immunoprecipitations using GFP traps were carried out as previously described [18]. Western blots were performed using standard procedures. Detection was carried out on a Li-Cor Odyssey Infrared scanning system using fluorescently labeled secondary antibodies. Primary antibodies were mouse anti-SNX27 (Clone 1C6, Abcam, cat# AB77799, 1:500), mouse anti-GFP (Roche, cat# 11814460001, 1:1000) and mouse
anti-mCherry (Abcam, cat# AB125096, 1:1000). Secondary antibodies were Alexa Fluor 680 goat anti-mouse IgG (Invitrogen, cat# A21057, 1:10000) and Alexa Fluor 800 goat anti-rabbit IgG (Invitrogen, cat# SA535571, 1:10000). Immunoprecipitations were performed on three independent biological replicates.

For pupal wing westerns, 28 hr APF pupal wings were dissected directly into sample buffer. One pupal wing equivalent was used per lane. Westerns were probed with mouse monoclonal anti-Fmi 74 (DSHB [33]) and Actin AC-40 mouse monoclonal (Sigma, cat# A4700, 1:5000). Secondary antibody was HRP-conjugated goat anti-mouse (DAKO, cat# P0447, 1:5000), and detection used SuperSignal West Dura Extended Duration Substrate (Thermo Scientific, cat# 34075). A BioRad ChemiDoc XR+ was used for imaging. Western blots were performed on four independent biological replicates. No sample size estimation was carried out, and no data were excluded.

Generation of CRISPR/Cas9 Snx27 and Fam21 mutants, and EGFP-fmi knockin flies

CRISPR/Cas9 Snx27 and Fam21 mutants were designed using protocols from http://www.crisprflydesign.org and [41], and gRNAs were designed using http://www.flyrnai.org/crispr2/. For Snx27 the following gRNAs were used: Snx27 gRNA1 – 5’ GCGAGC CATCCTCACGCC 3’ and Snx27 gRNA2 – 5’ ATCGAAAGAGCCAAATCC 3’. Guide RNAs were separately cloned into the pCFD3 vector (http://www.crisprflydesign.org/gna-expression-vectors/ and http://www.addgene.org/Simon_Bullock/) and co-injected into nos-Cas9 (Bloomington: 54591) embryos by the University of Cambridge Department of Genetics. Progeny were screened by PCR and sequenced, and one mutant allele was obtained, Snx27<sup>os</sup>. For the Fam21 knockout, CRISPR/Cas9 was used to generate a 4699 bp deletion, which removes the Fam21 5’UTR and all of the coding sequence except the last 36 bp. Gibson assembly was used to insert 1 kb homology arms into the 5’ and 3’ multiple cloning sites of pTV3 (a simplified version of pTVcherry described in [42], pTV3 sequence available on request) to generate pCFD4-Fam21. Two CRISPR targets within proximity of the homology arms, and within the sequence to be deleted, were chosen based on their efficiency and lack of off targets. The 5’ CRISPR target sequence was GCAGGCTTAGGGATGAGCGT (on the plus strand) and the 3’ CRISPR target sequence was TTTGATGAATGTAGCAGT (on the minus strand). These protospacer sequences were cloned into the pCFD4 vector [41] to generate pCFD4-Fam21. pTV3-Fam21 and pCFD4-Fam21 were mixed 1:1 and injected at a concentration of 500 ng/ul into nos-Cas9 (Bloomington: 54591) embryos by Bestgene. The resulting ‘founder’ adult males were crossed to y w; G, bc / CyO and the progeny were screened for mCherry fluorescence, which indicates successful targeting. Knockout alleles were verified by sequencing. EGFP was knocked into the endogenous fmi locus by homologous recombination. The sequence for EGFP was inserted into the pRK2 targeting vector [43], such that EGFP is upstream of the LoxP-white-LoxP cassette. Homology arms of approximately 3 kb of genomic DNA for fmi were inserted on either side, inserting EGFP 5 amino acids downstream of the PRO domain and 7 amino acids upstream of the first cadherin repeat, after amino acid Q355.

Targeting vectors were introduced into the genome by P-element mediated transgenesis, to produce donor strains. Homologous recombination was carried out as described by [43]. Targeted lines on the correct chromosome were verified by PCR of EGFP, and the white marker gene was excised by Cre-Lox recombination, leaving a single LoxP site downstream of EGFP.

Transgenics

Drosophila Snx27 cDNA (LD13361 Berkeley Drosophila Genome Project) was cloned into pEGFP-C1 (Clontech). GFP-Snx27 was then cloned into the tub-MCS vector and tub-GFP-Snx27 was integrated into the genome via P-element mediated transgenesis. P[acman]-EGFP-stbm was made by inserting an in-frame EGFP tag upstream of the stbm ORF in P[acman]-stbm, using standard recombineering methods, leaving a LoxP site between EGFP and the stbm ORF. For P[acman]-EGFP-stbmΔPDZbm, the final three amino acids of the stbm ORF were precisely deleted using recombineering with positive-negative selection. P[acman]-EGFP-stbm and P[acman]-EGFP-stbmΔPDZbm were integrated into the genome via eC31-mediated recombination into the attP40 landing site and recombined with arm-lacZ FRT40 stbm<sup>os</sup> or FRT40 stbm<sup>os</sup> respectively.

To make arm-PRO-EGFP-fmi, cDNA from the stan-PA isoform was used, which contains a PDZbm at its C terminus. PCR was used to insert EGFP in frame after Q355 in the N terminus of Fmi, 5 amino acids downstream of the predicted PRO cleavage site [44], and 7 amino acids upstream of the start of the first cadherin repeat. The ORF was then inserted downstream of the armadillo promoter and upstream of a polyA sequence in a modified pAtTb vector. PCR was used to delete the final three amino acids to make arm-PRO-EGFP-fmiΔPDZbm. arm-PRO-EGFP-fmi and arm-PRO-EGFP-fmiΔPDZbm were integrated into the genome via eC31-mediated recombination into the attP40 landing site and recombined with FRT40 fmi<sup>es9</sup> or arm-lacZ FRT40 fmi<sup>es45</sup> respectively.

Transgenics were made by Bestgene or Genetivision.

Pupal wing immunostainings, imaging, and quantitation

Pupal wings were dissected at 28 hr after pupariation formation (APF) at 25°C, or at 32 hr APF for trichomes [45]. Pupae were removed from their pupal case and fixed for 25-60 min in 4% paraformaldehyde in PBS, depending on antibody combinations. Wings were then dissected and the outer cuticle removed, and were blocked for 1 hr in PBS containing 0.2% Triton X-100 (PTX) and 10% normal goat serum. Primary and secondary antibodies were incubated overnight at 4°C in PTX with 10% normal goat serum, and all washes were in PTX. After immunolabelling, wings were post-fixed in 4% paraformaldehyde in PBS for 30 min. Wings were mounted in 25 μl of PBS containing 10% glycerol and 2.5% DABCO, pH 7.5. Antibodies used for immunolabelling were mouse monoclonal anti-Fmi 74 (DSHB [33]), rat anti-Stbm [34], affinity-purified rabbit anti-Fz [35], affinity-purified rat anti-Pk [36], rat anti-Dsh [37], mouse
monoclonal anti-Armadillo (DSHB), affinity-purified rabbit anti-GFP (Abcam, cat# ab6556), rabbit anti-β-galactosidase (MP Biomedicals/Cappel, cat# 55976) and mouse monoclonal anti-β-galactosidase 40-1a (DSHB). Trichomes were stained using fluorescein Phalloidin (Molecular Probes, cat# F-432) or Alexa Fluor 568 Phalloidin (Invitrogen cat# A12380). Pupal wings were imaged on a Nikon A1R GaAsP confocal microscope using a 60x NA1.4 apochromatic lens. Nine Z-slices separated by 150 nm were imaged. Wings from at least five independent animals were imaged for each experiment. No randomization, blinding or sample size estimation was carried out, and no data were excluded.

Antibody internalisation

5.5 hr APF prepupal wings were dissected in Schneider’s Medium (SM; Thermo-Fisher, cat# 21720) containing 10% fetal bovine serum (FBS; Sigma, cat# F4135) and transferred to a microtiter plate on ice. Medium was replaced with SM/FBS containing mouse monoclonal Fmi antibody 71 [33], and wings were incubated at 4°C for 30 min. Wings were washed briefly in SM/FBS on ice, and chased in 1 mL SM/FBS at room temperature (RT) for various times. Endocytosis was stopped by pipetting wings into SM/FBS at 4°C for 5 min, and wings were then fixed in 4% paraformaldehyde/PBS for 20 min. For detection of extracellular Fmi, tissue was incubated in Cy2-conjugated anti-mouse secondary antibody in the absence of detergent, and post-fixed before adding Alexa 647-conjugated anti-mouse secondary antibody in the presence of 0.1% Triton X-100, for total Fmi staining. Wings were mounted in Mowiol (Polysciences, cat# 17951) containing 2.5% DABCO. Pupal wings were imaged on a Nikon A1R GaAsP confocal microscope using a 60x NA1.4 apochromatic lens. Nineteen Z-slices separated by 150 nm were imaged using constant confocal settings. Wings from ten independent animals were imaged. No randomization, blinding or sample size estimation was carried out, and no data were excluded.

FRAP

For FRAP, 5.5 hr APF prepupal wing were mounted in a small volume of SM containing 1.25% methyl cellulose (Sigma cat# 274429) to reduce sample movement, and imaged within 45 min. Images were 256 × 256 pixels, with a pixel size of 100 nm, and a pinhole of 1.2 AU. ‘Hub-and-spoke’ ROIs of 3-4 μm² were selected, that covered a vertex and 3 half-cell edges. Three pre-bleach images were taken at 2 frames/sec, and ROIs were then bleached using a 488 nm Argon laser at 80% with 8 passes (1 s total time), which resulted in 60%–75% bleaching. Immediately following bleaching, 5 images were taken at 5 s intervals, followed by 10 images at 10 s intervals and 26 images at 15 s intervals. Laser power was adjusted to maintain constant power between different imaging sessions. Wings from ten independent animals were imaged. Based on the mean intensity and standard deviation of previous sets of wings, we determined that imaging at least 6 wings per genotype would allow detection of differences of 20% in the means, in a pairwise comparison, with a power of 0.8 and α = 0.05 (using G*Power). No randomization or blinding was carried out. The last five or six time points from two images were excluded, as the image moved out of focus during imaging.

QUANTIFICATION AND STATISTICAL ANALYSIS

Where the sample size was large enough, a D’Agostino and Pearson normality test was used to determine if the data fitted to a normal distribution. All datasets analyzed had a normal distribution, so parametric statistical tests were performed.

Quantification of western blots

Band intensities from four biological replicates were quantified using ImageJ. Data were compared using unpaired t tests.

Quantification of membrane intensity and polarity in pupal wings

The three brightest slices around apicolateral junctions were selected and averaged for each channel in ImageJ. Membrane masks were generated in Packing Analyzer [46], and MATLAB scripts were used to calculate mean membrane intensity for mutant or wild-type regions of each wing [22]. Polarity magnitude (maximum asymmetry ratio on a cell-by-cell basis) and variation in polarity angle were also calculated using MATLAB [22]. Error bars represent standard deviation. Values were compared between control and mutant regions of the same wings using paired t tests. Alternatively, one sample t tests were used to determine if the ratio of signal between mutant and wild-type regions of the same wing differed from 1.0 (p ≤ 0.05 *, p ≤ 0.01 **, p ≤ 0.001 ***).

Quantification of antibody internalisation

For quantification of extracellular staining, ImageJ was used to select and average the three brightest slices around apicolateral junctions, and to measure total signal in mutant or wild-type regions of each wing. Laser-off background was subtracted, and the readings were normalized to 1.0 at the 0 min time point for each genotype. Error bars represent standard deviation, and ANOVA with Dunnett’s multiple comparisons test was used to compare intensities to the 0 min control (p ≤ 0.05 *, p ≤ 0.01 **, p ≤ 0.001 ***).

Intracellular puncta were quantified from three slices just below the level at which junctional staining was no longer visible. A modified version of our MATLAB script [22] was used to find a threshold value that resulted in 1% of the total area in wild-type wings being identified as puncta. The same threshold value was then applied to mutant tissue of the same wing. The number, size, and intensity of puncta was determined, and error bars represent standard deviations. One sample t tests were used to determine if the ratio of values between mutant and wild-type regions of the same wing differed from 1.0 (p ≤ 0.05 *, p ≤ 0.01 **, p ≤ 0.001 ***).
FRAP processing

ImageJ was used to manually reselect up to six bleached regions in each image for each time point. The laser off background was subtracted, and the values were corrected for acquisition bleaching and normalized against the average of the prebleach values. Data were then plotted on an xy graph using Prism (v7 Graphpad) and bleached regions within the same wing were averaged. A two-phase exponential curve was fitted for each wing, as this was preferred over a one-phase exponential fit in Prism. Ten wings (n = 10) were then combined per genotype and an exponential association curve was fitted. Curves were compared using an extra-sum-of-squares F test (p ≤ 0.05 *, p ≤ 0.01 **, p ≤ 0.001 ***).

DATA AND SOFTWARE AVAILABILITY

A MATLAB script for comparing puncta intensities in two regions of the same wing, using the same threshold value to identify puncta, is supplied as Data S2.