RESEARCH ARTICLE

The Rab11 effectors Fip5 and Fip1 regulate zebrafish intestinal development

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ABSTRACT

The Rab11 apical recycling endosome pathway is a well-established regulator of polarity and lumen formation; however, Rab11-vesicular trafficking also directs a diverse array of other cellular processes, raising the question of how Rab11 vesicles achieve specificity in space, time and content of cargo delivery. In part, this specificity is achieved through effector proteins, yet the role of Rab11 effector proteins in vivo remains vague. Here, we use CRISPR/Cas9 gene editing to study the role of the Rab11 effector Fip5 during zebrafish intestinal development. Zebrafish contain two paralogous genes, fip5a and fip5b, that are orthologs of human FIP5. We find that fip5a- and fip5b-mutant fish show phenotypes characteristic of microvilli inclusion disease, including microvilli defects and lysosomal accumulation. Single and double mutant analyses suggest that fip5a and fip5b function in parallel and regulate trafficking pathways required for assembly of keratin at the terminal web. Remarkably, in some genetic backgrounds, the absence of Fip5 triggers protein upregulation of a closely related family member, Fip1. This compensation mechanism occurs both during zebrafish intestinal development and in tissue culture models of lumenogenesis. In conclusion, our data implicate the Rab11 effectors Fip5 and Fip1 in a trafficking pathway required for apical microvilli formation.

KEY WORDS: FIP5, FIP1, Rip11, RCP, Rab11, Microvilli

INTRODUCTION

Development of many organs, such as the gastrointestinal system, kidneys and respiratory tract, requires morphogenetic remodeling of cells to form a hollow tube, or lumen (Jewett and Prekeris, 2018). Whereas the mechanisms cells use to form a lumen vary by organ, a common feature is that cells adopt a highly polarized configuration including establishment of apical structures such as primary cilia, motile cilia or microvilli (Apodaca and Gallo, 2013). Intestinal epithelia are one of the few vertebrate cell types to lack primary cilia, but their apical cell surface is covered with a brush border composed of actin-rich membrane protrusions called microvilli to aid in nutrient absorption (Apodaca and Gallo, 2013). The molecular basis of cell polarization is well defined, but much less is understood about how trafficking pathways govern formation of these apical structures, especially in vivo.

The Rab11 apical recycling endosome pathway is a well-established regulator of polarity and lumen formation (Jewett and Prekeris, 2018). However, Rab11-directed trafficking events are also implicated in a number of other cellular processes, raising the question of how Rab11 vesicles achieve specificity when involved in numerous cellular functions. In part, Rab specificity is achieved through interaction with effector proteins, and Rab11 in particular interacts with a family of effector proteins called Rab11-family interacting proteins (FIPs) (Horgan and McCaffrey, 2009). There are five FIP family members, all of which contain a coiled-coil region at the C-terminus of the protein, allowing dimerization and binding to two Rab11 molecules, effectively forming a functional heterotetramer. Different FIPs appear to function in unique cellular processes including cytokinesis (FIP3 and FIP4), ciliogenesis (FIP3) and cargo recycling to the cell surface (FIP1, FIP2 and FIP5) (Horgan and McCaffrey, 2009). Previously, our lab implicated FIP5 in apical lumen formation in 3D Madin Darby Canine Kidney (MDCK) cell culture. We and others have shown that Rab11-FIP5 endosomes are required for lumenogenesis and interact with the actin-binding protein MYO5B to traffic cargo to the apical cell surface (Lapiere et al., 2001; Willenborg et al., 2011; Mangan et al., 2016). However, whether FIP5 plays a role in coordinating lumen morphogenesis during development in vivo is unknown.

Polarization is critical for cell function such that polarity disruption results in a number of diseases. Microvillus Inclusion Disease (MVID) is one such example, arising from the inability to form and maintain microvilli at the apical cell surface (Al-Daraji et al., 2010). Patients with MVID suffer from intractable diarrhea and malabsorption due to absent or very sparse microvilli and typically do not live past childhood. At the cellular level, patients display characteristic trafficking defects of lysosome accumulation and microvillus inclusions (Phillips et al., 1985; Phillips and Schmitz, 1992; Ruemmele et al., 2006). Mutations in MYO5B are found in patients with MVID and mutations in the zebrafish ortholog myoVb (also called goosepimples) and result in microvillus inclusions and trafficking defects (Müller et al., 2008; Ruemmele et al., 2010; Sidhaye et al., 2016). Moreover, experiments from intestinal tissue culture models suggest that the interaction between Rab11 and MYO5B is essential for microvilli maintenance (Knowles et al., 2014). Given that FIP5 interacts with MYO5B and is required for lumen formation in tissue culture, we hypothesized that FIP5 regulates intestinal development and microvilli formation in vivo.

RESULTS

The mechanisms by which cells polarize and form an apical lumen have been studied extensively in 3D tissue culture, but whether these processes are recapitulated in vivo is unclear, because vertebrate models are inherently more complex and have compensatory mechanisms. Furthermore, intestinal tissue culture models are limited due to a lack of proper microvilli that are subject to the stresses and strains encountered by a functional animal intestine. To
address these limitations, we used zebrafish intestinal development as an in vivo model of lumeneogenesis and microvilli formation. We first examined the degree to which zebrafish Fip5 protein was conserved with human and dog FIP5 protein, as most work on FIP5 during cell polarization has been performed in MDCK cells. Zebrafish contain Fip5a and Fip5b orthologs to mammalian FIP5 with two highly conserved functional domains: a phospholipid-binding domain C2 domain at the N-terminus and a coiled-coil region at the C-terminus of the protein (Fig. S1A, yellow and blue highlight, respectively) required for dimerization and binding to Rab11 (Prekeris et al., 2001). Zebrafish intestinal development begins around 3 days post-fertilization (dpf) when many small lumens develop throughout the intestinal tract and subsequently fuse to form a single continuous lumen from mouth to anus (Ng et al., 2005; Alvers et al., 2014). To determine where fip5a and fip5b were expressed in zebrafish larvae during development, we performed in situ hybridization on 4 dpf larvae. Luminal organs such as the intestine, spinal cord and notochord expressed fip5a and fip5b (Fig. S2A,B). In measuring mRNA levels of fip5a and fip5b, we found that both transcripts showed increased levels around 3 dpf, and high levels of fip5b mRNA persisted throughout 8 dpf (Fig. S2C). We thus focused our efforts first on fip5b.

**Zebrafish intestinal microvilli establishment requires Fip5b**

To study the function of Fip5b, we used CRISPR/Cas9 gene editing. We selected two different fip5b alleles that introduced a premature stop codon right after the C2 domain at the N-terminus (Fig. 1A; Fig. S1B), thereby eliminating the Rab-binding domain (RBD) at the C-terminus essential for Fip5 function. We maintained these fip5b mutant stocks in a heterozygous state and performed intercrosses to generate zygotic mutants for analysis. Stage matched wild-type siblings were used as controls. We performed qRT-PCR to measure fip5b expression in fip5bCO40-homozygous mutant larvae and observed an almost complete loss of fip5b mRNA levels (Fig. 1B), suggestive of nonsense-mediated decay. fip5bCO40-homozygous mutant fish appeared morphologically normal from embryo through adulthood and were homozygous viable as adults. However, to determine if loss of Fip5b affected intestinal development at the cellular level, we performed transmission electron microscopy on fixed sections through the midgut region (Fig. 1C, yellow box) at developmental time points. At 3 dpf, when intestinal lumen morphogenesis initiated, fip5bCO40 mutant larvae formed a single lumen (Fig. 1D), but upon closer examination, we noticed an accumulation of membrane vesicles in the subapical cytoplasm not present in wild-type larvae (Fig. 1E yellow box, F). These structures resembled microvilli inclusions, which are pathological hallmarks of MVID. At 6 dpf when intestinal development was mostly complete (Ng et al., 2005), microvillus inclusion-like structures were no longer evident near the subapical surface. Instead, intestinal cells of homozygous mutant larvae showed an accumulation of small (less than 500 nm) apical vesicles (Fig. 1G, H) and large (greater than 500 nm) organelles that resided medially in the cells (Fig. 1G arrows, I) compared to wild-type cells which did not show an accumulation of intracellular vesicles. Moreover, microvilli were shorter in both the anterior intestinal bulb and posterior midgut of 6 dpf homozygous mutant fish compared to wild-type siblings (Fig. 1G,G’). Finally, the terminal web, an apical cytoskeletal network anchoring microvilli into the cell, was disrupted in mutant fish. Wild-type larvae had a defined, electron-dense line at the base of the microvilli and an organell-free zone just below the apical cell surface which was absent in mutants (Fig. 1G, boxed region magnified in G’, bracket). These data revealed trafficking and microvilli defects in fip5bCO40-mutant larvae.

**Endosome maturation and terminal web keratin organization require Fip5b function**

To investigate the identity of the large organelles observed in fip5bCO40 mutant larval intestinal cells, we performed immunohistochemistry to detect proteins that serve as common endosome markers. Because Fip5 binds Rab11 vesicles, we first examined Rab11 localization. In wild-type intestinal cells, Rab11 vesicles localized just beneath the apical cell surface, as revealed by actin staining (Fig. 2A). In contrast, Rab11 vesicles mislocalized to the basolateral surface of intestinal cells in fip5bCO40 mutant larvae (Fig. 2A). Because the large organelles observed through electron microscopy in mutant tissue were near the apical cell surface, they were unlikely to be Rab11-positive. Intestinal cells of MVID patients accumulate lysosomal granules (Iancu et al., 2007), so we next stained cells to detect the late endosome/lysosome marker Rab7. Notably, Rab7-positive organelles accumulated near the apical cell surface in fip5bCO40 mutant cells, whereas we did not detect these large organelles in wild-type cells (Fig. 2B). These Rab7 endosomes were consistent in size and localization with the structures revealed by electron microscopy (Fig. 2C). Taken together, these data suggested that fip5b is required for Rab11 apical localization and regulation of trafficking processes.

Our electron microscopy analysis also revealed defects in microvilli length and the terminal web in fip5bCO40 mutant cells. The terminal web is composed of actin and intermediate filaments and is located just below the apical cell surface to anchor the base of microvilli into the cell (Moosker et al., 1984). Because actin localized to the apical cell surface of mutant cells similar to wild-type cells (Fig. 2A,B,D), we focused our attention on intermediate filaments. In polarized epithelia, intermediate filaments are composed of keratin polymers, so we stained cells with a pan-cytokeratin antibody to visualize intermediate filaments comprising the terminal web. We found that in wild-type cells, the keratin network resided just below the apical actin network; however, in fip5bCO40 mutant cells, keratin mislocalized to lateral and cytoplasmic regions of the cell (Fig. 2D,E). These observations were consistent with the possibility that Fip5b regulates keratin polymerization and terminal web formation at the apical cell surface.

Terminal web defects result in microvilli abnormalities, which can be exacerbated by physical stress from intestinal activity. We therefore hypothesized that fed mutant larvae would show more severe microvilli phenotypes than unfed 6 dpf larvae still living off the yolk. To test this, we began feeding the larvae daily at 7 dpf and then analyzed larvae at 11 dpf. Mutant larvae showed moderate trafficking defects at 11 dpf (Fig. 2F, arrows, G,H); however, the terminal web defects recovered, and microvilli were now significantly longer than wild-type siblings (Fig. 2F, bracket, I). This phenotypic recovery was unexpected and perhaps explains in part why adult mutant fish were homozygous viable. Importantly, these trafficking and microvilli phenotypes were recapitulated in another fip5b mutant allele fip5bCO62 (Figs S1B and S3A–D) indicating that these phenotypes were specific to fip5b. Taken together, these data provided evidence that fip5b functions in apical trafficking processes and microvilli formation during zebrafish intestinal development.

**fip5a functions similarly to fip5b in endosome maturation and terminal web organization**

Whereas fip5b mutant phenotypes were prominent during early developmental stages, these mutant fish recovered from these defects and were viable as homozygous adults. One possible explanation is a compensatory mechanism, perhaps through upregulation of another trafficking pathway, and an obvious candidate for compensation is the
Fig. 1. See next page for legend.
zebrafish *fip5b* paralog, *fip5a*. To test *fip5a’s* role in intestinal development, we again used CRISPR to create *fip5a*-mutant alleles (Fig. 3A, Fig. S1C). *fip5a*-mutant stocks were maintained in a heterozygous state and intercrossed to generate zygotic mutants for analysis. Stage matched wild-type siblings were used as controls.

Fig. 1. Zebrafish intestinal microvilli establishment requires Fip5b. (A) Domain schematic of zebrafish Fip5b protein containing a C2 domain at the N-terminus and a Rab-binding domain (RBD) at the C-terminus. The red arrowhead with STOP denotes premature termination codon in *fip5b* mutant alleles. (B) qRT-PCR for *fip5b* in wild-type and *fip5b*<sup>CO40</sup> mutant larvae at 6 dpf. n=50 larvae for each condition. (C) 6 dpf larvae expressing Tg(hsp: GFP;Rab11a) labeling the intestine. The intestinal bulb is denoted by a bracket and the midgut by a dashed box. All following images are representative cross sections through the midpoint region. Wild-type siblings are used as controls. For all experiments, three separate animals for each condition were analyzed. (D) Electron micrographs showing 3 dpf wild-type and *fip5b*<sup>CO40</sup> mutant larvae. Luminal space is lighter gray region. (E) High magnification electron micrographs showing 3 dpf wild-type and *fip5b*<sup>CO40</sup> mutant larvae. Yellow box shows zoomed in view on a region with subapical microvillus inclusion-like structures. (F) Quantitation of the mean number of microvillus length in the intestinal bulb and midgut in 6 dpf larvae. Each dot represents a single microvillus length combined across three animals. All plots show mean±s.e.m. A <sup>-test was used for Gaussian data and a Mann-Whitney test for all other statistics. ***<0.0005, **<0.005.

Fig. 2. Endosome maturation and terminal web keratin organization require *fip5b* function. All following images are representative cross sections through the midgut region. Wild-type siblings are used as controls. For all experiments, three separate animals for each condition were analyzed. (A,B,D) Immunohistochemistry on cross sections of 6 dpf wild-type and *Fip5b*<sup>CO40</sup> mutant larvae stained with Hoechst (blue), Phalloidin (red), and Rab11 (A), Rab7 (B), or Cytokeratin (D) (green). (C) Quantitation of Rab7-vesicle diameter. (E) Ratio of fluorescence intensity of apical keratin to cytoplasmic keratin. (F) Electron micrographs showing 11 dpf fed wild-type and *fip5b*<sup>CO40</sup> mutant larvae. Arrows point to larger than 500 nm organelles and brackets mark terminal web. (G) Quantitation of less than 500 nm apical vesicles in 11 dpf larvae. (H) Quantitation of greater than 500 nm organelles in 11 dpf larvae. (I) Quantitation of microvilli length in the intestinal bulb and midgut of 11 dpf larvae. Each dot represents a single microvillus length combined across three animals. All plots show mean±s.e.m. A t-test was used for Gaussian data and a Mann-Whitney test for all other statistics. ***<0.0005, **<0.005, *<0.05.

Similar to *fip5b* mutants, *fip5a*<sup>CO38</sup>-homozygous mutant larvae were morphologically normal and viable as homozygous adults. To study the role of *fip5a* during intestinal development, we performed the same transmission electron microscopy analysis on fixed sections through the mid-intestinal region. Notably, *fip5a*<sup>CO38</sup> mutant fish recapitulated phenotypes seen in *fip5b* mutant fish. At 3 dpf, *fip5a*<sup>CO38</sup> mutant larvae formed a lumen, but exhibited subapical organelles resembling microvillus inclusions (Fig. 3B,C). By 6 dpf, inclusion-like structures cleared, and *fip5a*<sup>CO38</sup> mutant cells now accumulated small apical vesicles (Fig. 3D,E) and large organelles (Fig. 3D, arrows, F) not present in wild-type larvae. Additionally, midgut microvilli were shorter (Fig. 3D,G) and the terminal web was also disrupted in mutants compared to wild-type larvae (Fig. 3D, brackets). Mutant larvae had large Rab7-positive organelles and terminal web defects appeared to be the result of mislocalized keratin from the apical cell surface (Fig. 3H–K). Again, similar to *fip5b* mutants at 11 dpf, *fip5a*<sup>CO38</sup> mutants maintained trafficking defects (Fig. 3L, arrows, M,N), but unlike *fip5b* mutants, the terminal web defects and shorter microvilli persisted in *fip5a*<sup>CO38</sup> mutants at 11 dpf (Fig. 3L, brackets, O). Importantly, these trafficking and microvilli phenotypes were recapitulated in another *fip5a* mutant allele, *fip5a*<sup>CO35</sup> (Figs S1C and S3E–H) indicating that these phenotypes were specific to *fip5a*. Collectively, these data implicated *fip5a* in apical trafficking and microvilli formation and suggested a similar function to *fip5b* during zebrafish intestinal development.

*fip5a* and *fip5b* double mutants show severe microvilli and trafficking phenotypes

*fip5a* and *fip5b*-homozygous mutant larvae showed similar phenotypes, but it remained unclear whether *fip5a* and *fip5b*
function in parallel or through a common pathway. To test this, we created a *fip5a*/*fip5b* heterozygous mutant line (*fip5a*<sup>CO35/+;</sup> *fip5b*<sup>CO40/+</sup>). This fish line was maintained in a heterozygous state and intercrossed to generate *fip5a*<sup>CO35/CO35</sup>/*fip5b*<sup>CO40/CO40</sup> homozygous double-mutant embryos for experiments. Wild-type siblings were used as controls. Through electron microscopy analysis at 6 dpf, *fip5a*<sup>CO35</sup>/*fip5b*<sup>CO40</sup> zygotic double-mutant fish showed two classes of phenotypes. The first was a severe microvilli defect where microvilli density was significantly reduced and the microvilli that did form were shorter and more heterogeneous in double mutants compared to wild-type larvae (Fig. 4A and A″, braces, C). The second was a severe trafficking phenotype where the majority of the cell cytosol was filled with giant Rab7-positive organelles (Fig. 4A′ and A‴, arrows, D,E). Double-mutant fish also accumulated small
apical and terminal web defects (Fig. 4A′, bracket, F,G) like those seen in single mutants. These phenotypes were not mutually exclusive, as some mutant larvae displayed both microvilli and trafficking defects. It is worth noting that wild-type siblings also showed mild microvilli, terminal web and trafficking defects (Fig. 4A′-A′, brace, bracket and arrows, respectively), perhaps suggestive of maternal contribution, as stage-matched wild-type AB fish did not show these phenotypes (Fig. 4B). In addition to these intestinal phenotypes, about 50% of the double-mutant larvae had multiple kidney lumens, whereas wild-type siblings or single *fip5a* or *fip5b* mutant larvae always had a single continuous kidney lumen (Fig. S3I). Moreover, double-mutant animals did not live past 2 weeks. Thus, the severity of these double-mutant phenotypes suggested that *fip5a* and *fip5b* function in parallel in microvilli formation during zebrafish intestinal development through apical trafficking pathways that regulate terminal web formation.

**Upregulation of Fip1 rescues *fip5a* and *fip5b* double-mutant phenotypes**

Although larvae deficient for zygotic functions of both *fip5a* and *fip5b* had severe intestinal phenotypes, contribution of wild-type maternal products to the eggs laid by heterozygous females potentially partially suppressed the phenotype. To test this possibility, we removed the maternal contribution of *fip5a* by intercrossing *fip5a*CO38/CO38;*fip5b*CO40/+ mutant larvae. We called these maternal-zygotic double mutants *fip5a*CO38⁎;*fip5b*CO40⁎- adults. We termed these maternal-zygotic double mutants *fip5a*CO38⁎;*fip5b*CO40⁎- to differentiate from zygotic double mutants in Fig. 3 created from a heterozygous intercross (Fig. 5A versus B). Surprisingly, *fip5a*CO38⁎;*fip5b*CO40⁎- larvae, lacking maternal and zygotic functions of *fip5a* and zygotic functions of *fip5b*, had no intestinal phenotypes and could not be discerned morphologically from wild-type larvae (Fig. 5C). Thus, removing maternal *fip5a* function suppressed, rather than enhanced, the phenotype of double-mutant larvae.

Recent literature has posited a role for compensatory mechanisms due to gene knockout when the mutant mRNA undergoes nonsense-mediated decay (Rossi et al., 2015; Eli-Brolosy et al., 2019). One compensatory mechanism included upregulation of transcripts similar in sequence to the mRNA encoded by the mutated gene (Eli-Brolosy et al., 2019). We thus wondered if another FIP family member could be upregulated in the absence of maternal and zygotic functions of *fip5a* and zygotic functions of *fip5b*. Previous work in our lab showed that both FIP5 and FIP1 bind the same Rab11 vesicles and FIP5 proteomics revealed an interaction with FIP1 (Willenberg et al., 2011; Mangan et al., 2016) (Fig. 5D). We thus used a MDCK tissue culture model of lumenogenesis to ask if FIP1 could compensate for FIP5. When MDCK cells were grown in an extracellular matrix, the majority of wild-type cells formed a single, continuous lumen inside the cyst of cells; however, most FIP5 and FIP1 double KO cells showed a multilumenal phenotype and a small percentage showed an inverted polarity phenotype (Fig. 5E,F). These luminal phenotypes were significantly more severe than FIP5 KO alone (Fig. 5F). Correspondingly, western blot analysis demonstrated that FIP1 protein levels were upregulated in FIP5 KO cells (Fig. 5G,H) and immunohistochemistry experiments with a FIP1 antibody confirmed this (Fig. 5I; Fig. S4A). This protein upregulation was specific to FIP1 in FIP5 KO cells, as FIP5 levels did not increase in FIP1 KO cells (Fig. 5G; Fig. S4B).

Moreover, FIP5 and FIP1 double KO cells did not show general defects in apical polarity or tight junction formation when grown in a polarized monolayer (Fig. S4C,D), suggesting that FIP5 and FIP1 function were specific to apical trafficking during lumenogenesis. Given that FIP1 could compensate for FIP5 in epithelial tissue culture, we asked if Fip1 could do the same in vivo. To test this, we performed immunohistochemistry on 6 dpf wild-type, *fip5b*CO40⁎, *fip5a*CO38⁎;*fip5b*CO40⁎, and *fip5a*CO38⁎;*fip5b*CO40⁎- larvae stained for endogenous Fip1 protein. Fip1 staining was mostly absent from wild-type, *fip5a*CO40⁎ mutant, and *fip5a*CO38⁎;*fip5b*CO40⁎- zygotic double-mutant larvae; however, we observed a significant increase in Fip1 signal in *fip5a*CO38⁎;*fip5b*CO40⁎- larvae, especially at the apical cell surface (Fig. 5L,K). This suggested that maternal contribution of wild-type *fip5a* may influence Fip1 protein levels to compensate for maternal and zygotic loss of *Fip5a* together with zygotic loss of *Fip5b*.

**DISCUSSION**

Rab11 specificity for a particular cellular pathway is achieved through interacting with effector proteins, and our work revealed a role for the Rab11 effector paralogs Fip5a and Fip5b in apical trafficking and microvilli establishment during zebrafish intestinal development through apical trafficking pathways that regulate terminal web formation.
maintain apical polarity (Knowles et al., 2014; Vogel et al., 2015; Weis et al., 2016). Our work posits an additional potential explanation in the intermediate filament networks. In polarized epithelia, groups of keratin proteins form polymers at the subapical cell surface, just below the apical actin cortex (Apodaca and Gallo, 2013). These actin and intermediate filament networks together comprise the terminal web, which is responsible for anchoring the microvilli rootlets into the cell. In flip5-mutant zebrafish, we observed loss of keratin localization from the apical cell surface to lateral and cytoplasmic regions. It remains unclear how Fip5 regulates apical keratin localization. Because keratins are cytosolic proteins whose assembly and disassembly into networks is mediated by phosphorylation state (Cooper, 2000), one possibility is that Rab11-Fip5 vesicles traffic a keratin kinase or phosphatase to the site of keratin polymerization thereby regulating network assembly. Alternatively, the effect of Fip5 on intermediate filament

Fig. 4. flip5a and flip5b double mutants show severe microvilli and trafficking phenotypes. All following images are representative cross sections through the midgut region of 6 dpf larvae. (A–A′′′′) Electron micrographs showing wild-type siblings and flip5a\(^{CO35/CO35}\), flip5b\(^{CO40/CO40}\) zygotic mutant larvae. Arrows point to larger than 500 nm organelles, braces point out sparse microvilli and brackets mark terminal web or lack thereof in mutants. N indicates number of representative larvae out of total number of larvae analyzed. (B) Electron micrograph showing wild-type AB larva. (C) Quantitation of microvilli density. Each dot represents the number of microvilli per micron for a field of view across three wild-type and five mutant animals. (D) Immunohistochemistry on cross sections of wild-type and flip5a\(^{CO35}\), flip5b\(^{CO40}\) mutant larvae stained with Hoechst (blue), Phalloidin (red), and Rab7 (green). (E) Quantitation of Rab7-vesicle diameter. (F) Immunohistochemistry on cross sections of wild-type and flip5a\(^{CO35}\), flip5b\(^{CO40}\) mutant larvae stained with Hoechst (blue), Phalloidin (red) and Cytokeratin (green). (G) Ratio of fluorescence intensity of apical keratin to cytoplasmic keratin. Three separate animals for each condition were analyzed. All plots show means±s.e.m. A t-test was used for Gaussian data and a Mann–Whitney test for all other statistics. ***P<0.0005.
polymerization could be a more indirect result of general disruption in intracellular trafficking events as we see an accumulation of a number of vesicles and larger organelles in mutant cells. Because we used an endpoint analysis, we cannot fully differentiate whether Fip5 functions in the formation or maintenance of microvilli; however, our results demonstrate that without Fip5a or Fip5b, zebrafish intestinal microvilli are disrupted and major endosomal trafficking defects are observed in KO animals.

Fig. 5. Upregulation of Fip1 rescues fip5a and fip5b double-mutant phenotypes. (A,B) Schematic of genetic crosses resulting in fip5aCO35; fip5bCO40 double zygotic or maternal/zygotic (mat-) mutant offspring generated from two different parental genotypes. (C) Electron micrographs showing cross sections through midgut of 6 dpf wild-type AB and fip5aCO35; fip5bCO40 mat- mutant larvae. Three separate animals for each condition were analyzed. (D) Cartoon schematic showing FIPS and FIP1 bind same Rab11 vesicles. (E) Wild-type and FIPS; FIP1 double KO (DKO) MDCK cells grown in an extracellular matrix to induce 3D lumen formation. Arrows denote multiple lumens in KO cyst. (F) Quantitation of luminal phenotypes from three biological replicates. (G) Western blot on wild-type and KO MDCK cell lysates probed for FIP1, FIPS or tubulin (control) antibodies. (H) Quantitation of FIP1 band intensity for wild-type and FIP5 KO cell lysates from one biological replicate. (I) Quantitation of FIP1 fluorescence intensity in wild-type and FIP5 KO cells grown in polarized monolayers from three biological replicates. Representative images are shown in Fig. S4A. (J) Immunohistochemistry on cross sections through midgut of 6 dpf wild-type, fip5bCO40 mutant, fip5aCO35, fip5aCO40 double mutant and fip5aCO35, fip5bCO40 mat- mutant larvae stained with Hoechst (blue), Phalloidin (red) and Fip1 (green). (K) Quantitation of fluorescence intensity of Fip1. Three separate animals for each condition were analyzed. All plots show mean ± s.e.m. A t-test was used for Gaussian data and a Mann–Whitney test for all other statistics. ***P<0.0005, *P<0.05.
Mutations in MYO5B have been found in patients with MVID. As such, several MYO5B-deficient animal models have been developed to elucidate the cellular mechanism behind MVID. Our fip5 mutant fish show a number of phenotypes consistent with these animal models including accumulation of apical vesicles, mislocalized Rab11, enlarged lysosomes, decreased microvilli length and structures resembling microvillus inclusions, although further work is needed to fully demonstrate that these inclusion-like structures are equivalent to those in MYO5B-deficient animals. It is also interesting to note some differences. In zebrafish myo5b mutant larvae, microvillus inclusions are maintained until 6 dpf (Sidhaye et al., 2016), whereas in our mutants structures resembling inclusions were only present in 3 dpf larvae. This discrepancy could be due to differences in function between Myo5b and Fip5, with Myo5b regulating more diverse trafficking pathways than Fip5. Consistent with this, studies in tissue culture have shown that MYO5B binds FIP2, another Rab11 effector closely related to FIP5 (Hales et al., 2002). Alternatively, our data show that the zebrafish paralogs Fip5a and Fip5b function in parallel, whereas fish only contain one gene for Myo5b, potentially causing myo5b mutants to demonstrate stronger phenotypes than single fip5a or fip5b mutants. Moreover, loss of microvillus inclusions has been observed in MYO5B mutant mice, where inclusions were more pronounced in neonates and disappeared after weaning (Knowles et al., 2014; Weis et al., 2016).

Our analyses of zygotic and maternal/zygotic double-mutant animals raise interesting questions about gene compensation and maternal contribution. Whereas zygotic double-mutant larvae showed severe trafficking and microvilli defects and did not survive past two weeks, it was surprising that the wild-type siblings also past two weeks, it was surprising that the wild-type siblings also past two weeks, it was surprising that the wild-type siblings also past two weeks, it was surprising that the wild-type siblings also.

**MATERIALS AND METHODS**

**Antibodies**
The following antibodies were used in this study: Pan cytokeratin AE1/AE3 (Abcam/ab27988) 1:50, Rab7 (Abcam/ab50333) 1:100, Rab11 (Life Technologies/713500) 1:100, GP135 (DSHB/3F2/D8) 1:100, Cingulin (Prekeris Lab) 1:100, FIP1 (Prekeris Lab) 1:200, FIP5 (Prekeris Lab 1:100), Alexa 488 Anti-Rabbit secondary (Jackson ImmunoResearch/711-545-152) 1:200, Alexa 488 Anti-Mouse secondary (Jackson ImmunoResearch/715-545-150) 1:200, Alexa-568 Phalloidin (Invitrogen/A12380) 1:100.

**Zebrafish husbandry**
All animal work was approved by the Institutional Animal Care and Use Committee at the University of Colorado School of Medicine, CO, USA. All stocks unless otherwise specified were maintained in a heterozygous state and kept according to Standard Operating Procedure defined in ‘The Zebrafish Book’ (M. Westerfield, Institute of Neuroscience, University of Oregon, OR, USA).

**qRT-PCR**
RNA extraction from larvae was performed with TRizol reagent (Invitrogen) followed by cDNA synthesis with (Script cDNA Synthesis Kit (Bio-Rad)). SYBR Green PCR Master Mix (Applied Biosystems) was used for qPCR. All reactions were performed in technical triplicate and a minimum of three biological replicates were performed. Primer sequences are listed in Table S1.

**Protein alignments**
Fip5 protein alignments were generated using T-Coffee and Boxshade 3.2. The following protein accession numbers from NCBI were used for alignments: human NP_056285; dog XP_003639656 (isofrm X5); zebrafish Fip5a XP_009305489 (isofrm X2); zebrafish Fip5b XP_017214658 (rab11 family-interacting protein 5-like isofrm X2).

**Zebrafish immunohistochemistry**
At all timepoints analyzed, animals had not yet developed sex organs, thus experiments were blinded to gender. Larvae were placed in 1–2% Tricaine for 10 min or until they were unresponsive to touch then decapitated immediately posterior to the otic vesicle using a scalpel. The larva body was placed in fix solution (4% paraformaldehyde, 4% sucrose, 0.15 mM CaCl₂, pH 7.3) at 4°C overnight, whereas the head was placed in lysis buffer and genotyped (see genotyping). The fixed larvae were then embedded in a melted agar solution (1.5% agar, 5% sucrose in water) and, after the blocks hardened, they were trimmed and immersed in 30% sucrose in water solution overnight at 4°C. Blocks were then dried with a chempaper, frozen on dry ice for ~15 min, then stored at -80°C until ready to section.

Blocks were mounted in OCT and 20 μm sections cut using a Leica CM 1950 cryostat microtome. Sections were placed on FisherBrand charged slides (12-550-15) and rehydrated in PBS for 30 min. Excess liquid was dried, and then a wax pen was used to draw around the edge of the slide. Slides were then blocked with 2% BSA and 5% donkey serum (Thermo Fisher Scientific, NC9624464) in PBS for 1 h, then incubated in primary antibody diluted in block at room temperature for 2–3 h. Slides were then washed four times, for 15 min each with PBS and incubated in secondary antibody diluted in block for 1–2 h at room temperature. Slides were again washed four times, for 15 min each, with PBS, adding Hoescht (Thermo Fisher Scientific, 33342 at 1:500) to the second-to-last wash. Slides were then dried, mounted in Vectorshield (Vector Laboratories, H-100), and sealed with nail polish.

**Widefield microscopy and image analysis**
All slides of fixed fish sections were imaged with an inverted Axiovert 200M microscope (Carl Zeiss) with a 63× oil immersion lens and QE charge-coupled device camera (Sensicam). Images were acquired using Slidebook 6.0 (Intelligent Imaging Innovations) software. Images were processed using a combination of Slidebook 6.0 (Intelligent Imaging Innovations) software, Fiji (PMID 22743772), and Adobe Photoshop. Figures were made in Adobe Illustrator. A minimum of three biological replicates were performed for each experiment and quantitation was performed unblinded.

**Genotyping zebrafish**
Fish tissue was isolated from a fin clip for adult fish or from the heads for larvae. Fish tissue was placed in lysis buffer (10 mM Tris pH 8.3, 50 mM KCl, 1.5 mM MgCl₂, 0.3% Tween-20, 0.3% NP-40 in water) with 2% Proteinase K (Invitrogen, 25530049). Lysis reactions were incubated at 55°C for 4 h, then 95°C for 20 min to inactivate Proteinase K. A PCR/restriction-enzyme-based assay was used to genotype fip5a and fip5b mutant fish lines. For fip5a, a 400 base pair region of genomic DNA surrounding the CRISPR target site was amplified by PCR. The PCR product was then digested with BssHII for 1 h at 50°C, and the resulting product was run on a 2% agarose gel. For fip5b, a similar schematic was used with the BsaWI or Agel restriction enzyme depending on the allele. PCR primer sequences are listed in Table S1. Genotyping was performed prior to experiments and only wild-type and homozygous mutant larvae were selected for analysis.
CRISPR/Cas9 in zebrafish

All primer sequences are listed in Table S1. Guide RNA (gRNA) oligos were designed using ZiFIT Targeter Software for CRISPR/Cas9 Nucleases (Sander et al., 2010, 2007). The gRNA target sites were then blasted (NCBI BLASTn) against the zebrafish genome to look for potential off-target sites. gRNA oligos were annealed and phosphorylated, then cloned into the pDR274 vector (Addgene) using the Bsal-HF restriction site. Positive clones were sequenced to confirm correct insertion. The gRNA-containing vector was linearized using DraI and purified by ethanol precipitation. The gRNA sequence was then transcribed to RNA using T7 polymerase and purified by phenol chloroform extraction. Cas9 mRNA was synthesized from the pT3TS vector (Addgene) using mMESSAGE mMACHINE T3 Transcription Kit (Thermo Fisher Scientific, AM1348) and purified using phenol chloroform extraction. The injection mix was prepared as follows: 0.2 M potassium chloride, 0.15 ng/µl Cas9 mRNA, 70 ng/µl gRNA, and 10% phenol red in DEPC water. Embryos were injected with 1–3 nl of the injection mixture at the one-cell stage. Founder fish were determined by incubating tissue in 500 µl of 1:1 osmium tetroxide to 0.1 M sodium cacodylate for 2 h. Tissue was washed with double-distilled water, then incubating tissue in 500 µl of 1:1 osmium tetroxide to pH 7.4 for 30 min. Larvae turned brown at this point. Larvae were washed again in double-distilled water, then an ethanol dehydroxylation series was performed (50%/75%/100%). Larvae were then incubated in 1:1 Epon to ethanol for 1 h, then 2:1 Epon to ethanol overnight. The following day, larvae were embedded in 100% Epon, which was replaced with fresh 100% Epon, and then baked for 2 days. Larvae were cut in half where the body narrows (see schematic in Fig. 1C), and then 65 nm thick sections were cut and collected on formvar-coated copper slot grids. Sections cut in the anterior direction were designated the intestinal bulb region and in the posterior direction the midgut. Sections were imaged on a FEI Tecnai G2 Biotwin Transmission Electron Microscope, run at 80 kV with a side-mount AMT XR005-B digital camera. For transmission electron microscopy quantitation, a minimum of three biological replicates were used for each experiment and images were blinded prior to analysis.

RNA in situ hybridization

Sense and antisense RNA probes were designed against both the coding sequence and 3’ UTR region of zebrafish fip5a and fip5b genes. A PCR-based method with T7 sites at the end of the primers was used to amplify the probe DNA sequence from 8 dpf wild-type fish cDNA (see Table S1). The RNA probes were transcribed with the T7 polymerase and labeled using the DIG RNA labeling kit (Roche, 11175025910). After the labeling reaction was complete, the probes were mixed with 1 µl 0.5 M EDTA to stop the reaction, then 2 µl 5 M lithium chloride and 75 µl cold ethanol were added for purification by ethanol precipitation and the probe was resuspended in DEPC water. The RNA probe was checked by agarose gel electrophoresis, then mixed with an equal volume of formamide and stored at -80°C. RNA in situ hybridization assays were conducted based on a modified version of a previously published protocol described by Hauptmann and Gerster (2000). Larvae were fixed in 4% paraformaldehyde in DEPC PBS overnight at 4°C. Larvae were stored in MeOH at -20°C until use, when they were washed twice for five minutes in DEPC-PBSTw (0.5% Tween-20 in PBS made with DEPC water). Pigmentation was bleached in a hydrogen peroxide solution (3% H2O2, 0.5% KOH in DEPC water) until larvae eyes turned brown (15–30 min). Larvae were then washed twice for 5 min in DEPC-PBSTw, fixed for 20 min at room temperature in 4% PFA in DEPC-PBS, and washed again twice for 5 min in DEPC-PBSTw. The larvae were digested with 0.1 mg ml-1 Proteinase K (Invitrogen, 25530049) in DEPC-PBS for 17 min to permeabilize the larvae, then washed twice for 5 min each wash in DEPC-PBSTw, followed by fixation for 20 min in 4% PFA, and again washed twice for 5 min in DEPC-PBSTw. Larvae were incubated in 500 µl hybridization media (HM) block solution (50% formamide, 5x saline-sodium citrate buffer, 10 µl ml-1 tRNA, 50 mg ml-1 heparin, 0.01 M citric acid and 0.5% Tween-20 in DEPC H2O) for 1 h at 70°C. The block was replaced with HM containing 200 ng of the appropriate RNA probe, and the larvae were incubated at 70°C overnight. The following day, a series of progressive washes were performed for 10 min each wash at 70°C: 200 µl 100% HM without probe, 300 µl 66% HM/33% 2x saline-sodium citrate buffer (SSC; Cellgro, Mediatech, Inc., Manassas, VA, USA), 300 µl 33% HM/66% 2x SSC, 1 ml 2x SSC, 1 ml 0.2x SSC, 1 ml 0.1x SSC (this wash was performed twice), 1 ml DEPC-PBSTw. Another 10 min wash with DEPC-PBSTw was performed at room temperature, followed by a 1 h antibody block (2% sheep serum and 2 mg ml-1 BSA in DEPC-PBSTw). Anti-digoxigenin-AP Fab fragments (Roche Applied Science, Indianapolis, IN, USA) was incubated in antibody block overnight at 4°C. Finally, four 15 min washes were conducted at room temperature in DEPC-PBSTw. Larvae were incubated in staining solution (0.1 M Tris, pH 9.5, 0.25 M MgCl2, 0.1 M NaCl, 0.5% Tween-20) for 15 min at room temperature. Thereafter, the larvae were moved to a staining dish, covered with 500 µl precipitating BM Purple AP Substrate (Roche Applied Science, Indianapolis, IN, USA), and incubated at 37°C for 8 h until staining was visible. The larvae were then washed twice for 5 min in PBS and imaged or processed for sectioning immediately.

Cell culture and immunohistochemistry

MDCK II cells (ATCC and negative for mycoplasma contamination) were cultured in DMEM with 10% FBS and penicillin/streptomycin. For polarized MDCK experiments, cells were plated on collagen type I-coated Transwell filters (Corning, 3460) to reach confluence in 24 h. Cells were then grown for a further 3 days before transepithelial resistance measurements or fixation. Cells were fixed with 4% paraformaldehyde for 20 min at room temperature. Cells were blocked for 1–2 h in block buffer (PBS, 0.1% Triton X-100, 10% normal donkey serum). Primary antibodies were diluted in block buffer and incubated overnight at room temperature. Cells were washed with PBSx3 before adding secondary antibodies for 1–2 h at room temperature. Cells were washed again before mounting in VectaShield and sealing with nail polish or mounting in Prolong Gold. Coverslips used for all experiments were #1 thickness.

Trans-epithelial resistance measurements

MDCK cells were grown on collagen-coated Transwell filters (see Cell culture and immunohistochemistry section) and resistance measurements were taken four days after plating with a Millicell ERS-2 Voltohmmeter (Millipore). Three measurements per well, one for each space between the plastic prongs of the filter holder, were averaged and subtracted from the average of the blank well containing a collagen-coated filter without any cells.

Generating MDCK and RPE-1 CRISPR knockout lines

MDCK cells stably expressing Tet-inducible Cas9 (Dharmacon Edit-R inducible lentiviral Cas9 nuclease) were grown in a 12-well dish to about 75% confluency before treatment with doxycycline at a final concentration of 1 µg ml-1 for 24 h to induce Cas9 expression. Cells were then transfected with crRNA:tracrRNA mix as described for DharmaFECT Duo co-transfection protocol (Horizon Discovery, T-2010-xx). Transfected cells were incubated for 24 h before trypsinizing and plating for individual clones. Individual clones were screened through genotyping PCR and Sanger sequencing. All CRISPR gRNAs are listed in Table S1.

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Competing interests
The authors declare no competing or financial interests.

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