Tumor Protein D54 is a promiscuous Rab effector

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Intracellular membrane traffic is coordinated by Rab GTPases and their effector proteins. Here, we investigated the function in human cells of Tumor Protein D54 (TPD54/TPD52L2), a member of the TPD52-like protein family. Our initial experiments suggested that TPD54 has a role in anterograde membrane trafficking and in the recycling of endocytosed proteins back to the cell surface. To understand how TPD54 controls these diverse functions, we used an inducible method to reroute TPD54 to mitochondria. Surprisingly, this manipulation resulted in the capture of many functional vesicles (30 nm diameter) at the mitochondrial surface. To investigate the identity of captured vesicles, we did a rerouting screen using TPD54 as bait and the Rab GTPases as prey. This screen revealed that TPD54 associates promiscuously with at least 16 out of 43 different Rab GTPases. Our data suggest that TPD54 binds as an effector protein to Rabs involved in anterograde traffic and recycling where it likely promotes vesicle fusion.

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Introduction

Eukaryotic cells are by definition compartmentalized: they contain organelles and membrane-bound domains that have distinct identities. Vesicle transport between these locations is tightly regulated to maintain these identities yet allow exchange of specific materials. The Rab subfamily of small GTPases and their effector proteins are central to the control and regulation of vesicle transport. Rabs exist in two different nucleotide-bound conformations. A GDP-bound Rab is inactive and associates with a GDP-dissociation inhibitor (GDI) so that it may be extracted and transported to the appropriate donor membrane where it is stabilized by a guanine nucleotide exchange factor (GEF) (Wu et al., 2010; Pylypenko et al., 2006; Blümer et al., 2013). The GEF activates the Rab, exchanging GDP for GTP, and the active GTP-bound Rab can now recruit an effector protein (Müller and Goody, 2018). The roles of Rab effector proteins range from the formation of the vesicle, to its transport or its subsequent fusion (Johansson et al., 2007; Semerdjieva et al., 2008; Murray et al., 2016). When the vesicle reaches the acceptor membrane, a GTPase-activating protein (GAP) inactivates the Rab and a GDI extracts the Rab for the cycle to begin again (Müller and Goody, 2018). There are more than 60 Rab proteins in humans, each having a definite set of interactors: GEFs, GAPs and effectors (Barr and Lambright, 2010; Pylypenko et al., 2018). Identifying Rab interactors to understand how Rab GTPases orchestrate membrane traffic is a major goal in cell biology (Kanno et al., 2010; Koch et al., 2016; Gillingham et al., 2014).

In humans, there are four Tumor Protein D52-like proteins (TPD52-like proteins: TPD52, TPD53/TPD52L1, TPD54/TPD52L2 and TPD55/TPD52L3) some of which have been associated with membrane trafficking but the cellular biological roles of the family are not well characterized. TPD52-like proteins are short (140 to 224 residues), have 50% identity and each contain a coiled-coil domain through which they can homo- and heterodimerize. All are ubiquitously expressed with the exception of TPD55, which is restricted to testis (Cao et al., 2006). TPD52 was the first of the family to be identified due to its overexpression in cancer, and it is still the best studied. However, all members have been found to be overexpressed in a series of cancers (Cao et al., 2006; Byrne et al., 1995, 1998; Nourse et al., 1998). Overexpression of TPD52 correlates with poor prognosis in breast cancer patients and in cell models, TPD52 overexpression promotes proliferation and invasion (Byrne et al., 2010, 1996; Li et al., 2017; Dasari et al., 2017).

Rather disparate membrane trafficking functions have been reported for TPD52 and TPD53. First, TPD52 is involved in secretion in pancreatic acinar cells (Thomas et al., 2004, 2010; Messenger et al., 2013). Second, membrane trafficking proteins bind to TPD52, such as the endocytic protein Rab5c (Shahleydari et al., 2014), and the transcytotic protein MAL2 (Wilson et al., 2001). Third, TPD52 has a role in lipid droplet biogenesis at the Golgi (Kamili et al., 2015). Finally, a role in membrane fusion was proposed for TPD53 (Proux-Gillardaux et al., 2003). By contrast, the potential functions of TPD54 remain unexplored.

What is striking about TPD54 is its sheer abundance in cells. Previous quantitative proteomic analyses revealed that TPD54 is one of the most abundant proteins in HeLa cells, ranked 180th out of 8,804 (Hein et al., 2015; Kulak et al., 2014). There are an estimated 3.3 × 10⁶ copies of TPD54 per HeLa cell (2.7 μM), whereas abundant membrane trafficking proteins such as clathrin light chain A or β2 subunit of AP2 total 2.2 × 10⁵ and 1.0 × 10⁵ copies (1.8 μM or 0.4 μM), respectively (Hein et al., 2015). Despite its abundance, there are virtually no published data on the cell biology of TPD54. Due to sequence similarity and heterodimerization properties, we hypothesized that TPD54, like other members of the family, would also be involved in membrane trafficking. We set out to investigate the cell biology of TPD54 and found that it is associated with small transport vesicles where it acts as an effector to at least one-quarter of the Rab GTPase family. Our experiments highlight roles for TPD54 in regulating anterograde membrane traffic as well as recycling.
Results
TPD54 is a membrane trafficking protein. To investigate the subcellular localization of TPD54, we tried to make antibodies to see TPD54 by immunofluorescence microscopy. We did not succeed and therefore we generated a cell line where TPD54 was endogenously tagged with monomeric GFP (Figure 1A and Supplementary Figure S1). GFP-TPD54 fluorescence was diffuse in the cytoplasm, but was also seen at the Golgi apparatus, marked with GaIT-mCherry, and on endosomes, marked by APPL1 and OCRL1. It also colocalized with various membrane trafficking proteins, such as clathrin light chain a and the R-SNARE, VAMP2 (Figure 1A). A similar pattern was seen by overexpression of GFP- or mCherry-tagged TPD54 in parental cells. These observations suggest that TPD54 is a protein associated with membrane trafficking.

As a next step to characterizing TPD54, we investigated the binding partners of TPD54. To do so, we performed an immunoprecipitation (IP) of GFP-tagged TPD54 from HeLa cell lysates and analyzed co-precipitating proteins by mass spectrometry (Figure 1B). We found that two other members of the TPD52-like family, TPD52 and TPD53, were significantly enriched in the TPD54 samples versus control. TPD52, TPD53 and TPD54 have been reported to heterodimerize (Byrne et al., 1998), and this suggested that our assay was able to detect binding partners of TPD54. Among the other significant hits, we found the Rab GTPases Rab14, Rab2a and Rab5c. Rab14 has been identified as a regulator of the transport between the Golgi apparatus and early endosomes (Junutula et al., 2004), as well as from the Golgi apparatus to the plasma membrane (Kitt et al., 2008). Rab2a is on the endoplasmic reticulum (ER) to Golgi pathway (Tisdale et al., 1992), and Rab5c is found on the endocytic pathway (Bucci et al., 1995). Taken together, the results confirm that TPD54 is a protein involved in membrane trafficking.

TPD54 regulates the anterograde and recycling pathways. To investigate potential functions of TPD54, we sought to identify trafficking defects caused by the loss of TPD54. Using RNAi to deplete TPD54 in HeLa cells, we first assessed the transport of cargoes from the ER to the Golgi apparatus, and from the Golgi to the plasma membrane (PM) with the RUSH (retention using selective hooks) system (Boncompain et al., 2012). Briefly, the RUSH system allows the synchronous release of a reporter (here, GFP-tagged E-cadherin with a streptavidin-binding domain) from an ER-localized hook (here, streptavidin fused to a KDEL motif) by addition of biotin. After release, EGFP-E-cadherin is transported from the ER to the PM, via the Golgi apparatus. In control cells, the reporter reached maximal intensity at the Golgi between 14 and 28 minutes after release and then left the Golgi for the PM (Figure 2A and Supplementary Video SV1). By contrast, TPD54-depleted cells had obviously delayed kinetics (Supplementary Video SV2). We quantified the fluorescence of the reporter at the Golgi and expressed it as a fraction of the total cell fluorescence (Figure 2B). The resulting data were best described by a logistic function representing ER-to-Golgi transport and a line fit to describe Golgi-to-PM (Figure 2C-D, see Methods). This automated procedure allowed us to find the half-time (τ₁/₂) for ER-to-Golgi and ER-to-PM transport, and also infer the Golgi transport time as the difference between these times (Figure 2E-G). The data suggest that TPD54-depleted cells have delayed export of E-Cadherin at all stages.

We also wanted to know if TPD54 was required for endocytosis or cargo recycling, since Rab5c was one of our mass spectrometry hits (Figure 1B). To do so, we performed a transferrin uptake and recycling assay in TPD54-depleted and control HeLa cells. The internalization of transferrin was unchanged, but recycling to the plasma membrane was slower in TPD54-depleted cells (Figure 2H). In these experiments, the efficiency of the depletion was checked by western blot analysis, using α-tubulin as a loading control (Figure 2I). These results suggest that TPD54 operates in several steps of the anterograde pathway and in endosomal recycling.

Rerouting TPD54 to mitochondria allows detection of its associated proteins. To better understand the pathways on which TPD54 functions, we wanted to test which proteins co-reroute with TPD54 when it is rerouted to mitochondria using rapamycin-induced heterodimerization. Proteins tagged with an FK506-binding protein (FKBP) domain can be rerouted to MitoTrap, a mitochondrially-targeted FRB domain, by the addition of rapamycin (Robinson et al., 2010). As expected, mCherry-FKBP-TPD54 but not mCherry-TPD54 was efficiently rerouted to mitochondria with this method (Figure 3A). The kinetics of rerouting was reasonably fast, with TPD54 appearing at mitochondria 6 s post-rapamycin (Figure 3B-C and Supplementary Video SV3). The increase in mitochondrial TPD54 was best fit by a single exponential function (χ² = 0.43, τ = 37.98 ± 0.38 s) while the loss in cytoplasmic signal followed similar kinetics (τ = 47.52 ± 0.21 s, Figure 3B). Although the kinetic analysis suggested a single process, close inspection of the movies suggested that the pool of TPD54 which is apparently cytosolic is rerouted first, making visible a small number of endosomes (Figure 3C, arrow), before they also became rerouted leaving no detectable TPD54 away from the mitochondria (Figure 3E, 4 min).

To test if TPD54-binding proteins become co-rerouted to the mitochondria in this assay (Figure 3D), we used TPD52. TPD52 is a TPD52-like protein family member that can heterodimerize with TPD54 (Byrne et al., 1998) and was also detected in our mass spectrometry analysis (Figure 1B). As shown in Figure 3E, TPD52 is efficiently co-rerouted with TPD54, suggesting that the rerouting method can be used as a way to find proteins that associate with TPD54. During our TPD54 rerouting experiments we noticed that once the rerouting was complete, mitochondrial morphology changed and that mitochondria began to aggregate (Figure 3C).

Mitochondrial rerouting of TPD54 results in vesicle capture. To investigate why mitochondrial morphology be-
came altered at later time-points after rerouting TPD54, we used electron microscopy (EM) to examine the ultrastructure of mitochondria at different time-points (Figure 4). Cells expressing MitoTrap and mCherry-FKBP-TPD54 were imaged by EM. The mitochondrial TPD54 signal was fixed at various times after rerouting and the same cells were aged as rapamycin was applied (Figure 4A). They were then expressing MitoTrap and mCherry-FKBP-TPD54 were imaged by EM to examine the ultrastructure of mitochondria at different time-points (Figure 4). Cells were below the resolution limit of the microscope. Close inspection of live cell imaging data showed that the cytoplasmic TPD54 signal flickered, whereas free GFP did not (Supplementary Video SV4), suggesting that the cytoplasmic GFP-TPD54 signal is mainly vesicular. To address this point quantitatively, we used fluorescence recovery after photobleaching (FRAP) to assess diffusion and/or exchange of GFP-TPD54 at sub-resolution structures (summarized in Figure 5 and Table 1). FRAP kinetics were much slower for GFP-TPD54 (either expressed or endogenous) compared to GFP, suggesting diffusion-coupled binding of GFP-TPD54 to membranes. There were two phases of GFP-TPD54 recovery, a small fast process (τ = 2 s) with the majority of recovery via a slow process, which was on the order of tens of seconds. These kinetics were consistent with the majority of TPD54 binding to subcellular structures, with a minor fraction being cytosolic. Analysis of individual FRAP traces showed that in cells expressing higher levels of GFP-TPD54, FRAP was faster and that this was due to a larger fraction recovering via the fast process (Figure 5B). These experiments indicate that the freely diffusing pool of TPD54 is minimal and the majority of TPD54 is associated with small vesicles below the resolution limit. This conclusion is consistent with the vesicle capture by mitochondria during TPD54 rerouting.

Vesicles captured by TPD54 rerouting to mitochondria are functional. The small size of the vesicles captured by TPD54 rerouting to mitochondria raised the question of whether or not they were functional. As a first step to answering this question, we sought to identify the vesicles’ cargo. To do this, we tested five model cargoes where the alpha chain of CD8 is fused to different peptides that bear various endocytic motifs (Kozik et al., 2010). Briefly, CD8-FANPAY, CD8-YAAP or CD8-EAAALL have a single YXXφ, [F/Y]XNPX[Y/F] or [D/E]XXX[L/I/M] motif, respectively. CD8-CIMPR has the tail of the cation-independent mannose-6-phosphate receptor (CIMPR), which contains at least four endocytic motifs including two of the dileucine type. As a control, CD8-8xA was used which has
eight alanines, no endocytic motif and cannot be internalized. We examined the subcellular distribution of these cargoes in cells where TPD54 had been rerouted to mitochondria. In the control condition with no addition of rapamycin, all CD8 constructs were in endosomes or in the case of CD8-8xA, at the plasma membrane. After rerouting, the localization of CD8-8xA, CD8-FANPAY and CD8-YAAL was unaffected, whereas CD8-EAAALL and CD8-CIMPR were co-rerouted with TPD54 to the mitochondria (Figure 6A). To ensure that this co-rerouting was genuine and not a peculiarity of the model cargoes, we confirmed that endogenous CIMPR also co-rerouted with TPD54 (Figure 6B). This suggested that

Table 1. Kinetics of fluorescence recovery after photobleaching (FRAP).

| Protein                  | Recovery | $\tau_{fast}(s)$ | $\tau_{slow}(s)$ | Fast (%) | Slow (%) |
|--------------------------|----------|------------------|------------------|---------|----------|
| GFP                      | 0.960    | 0.74             | 6.90             | 62.4    | 37.6     |
| GFP-TPD54                | 0.868    | 2.14             | 13.40            | 40.9    | 60.1     |
| GFP-TPD54 (endogenous)   | 0.703    | 2.41             | 20.25            | 23.1    | 76.9     |
vesicles with cargo harbouring a dileucine motif were preferentially captured by TPD54 rerouting (Figure 6C). Receptors containing dileucine endocytic motifs are internalized at the PM and then recycled either via recycling endosomes or the Golgi apparatus. We therefore labeled CD8-EAAALL at the surface with Alexa 488-conjugated anti-CD8 antibodies, allowed internalization and trafficking to proceed, and at different time-points, performed mitochondrial vesicle capture via TPD54 rerouting (Figure 6D). Capture of surface-labeled CD8-EAAALL occurred at time-points greater than 60 min after internalization (Figure 6E). These experiments indicate that dileucine motif-containing receptors transit via the vesicles which can be captured on mitochondria by TPD54 rerouting, but only at late time points after internalization. The time course of capture is consistent with recycling of endocytic cargo from the Golgi apparatus.

A second test of vesicle functionality is whether the vesicles contain the machinery for fusion. Accordingly, we tested for co-rerouting of endogenous SNAREs in our vesicle capture assay. Generally, vesicle-resident R-SNAREs but not target membrane-resident Q-SNAREs were co-rerouted with TPD54 to mitochondria. We found co-rerouting of the R-SNAREs VAMP2, VAMP3, VAMP7 and VAMP8, but not the Q-SNAREs STX6, STX7, STX8, STX10 or STX16 (Figure 7). There was some evidence of selectivity with the localization of the R-SNARE VAMP4 being unaffected by TPD54 rerouting. Together these experiments indicate that, despite their small size, the vesicles captured by TPD54 rerouting are functional: they contain specific cargo and they have R-SNAREs for fusion. Moreover the presence of different SNAREs suggests that although the captured vesicles appear morphologically homogeneous, they are likely to be a crowd of different vesicle identities.

TPD54 is associated with multiple Rab GTPases. What are the identities of the vesicles captured by TPD54 rerouting? To answer this question, we screened 43 GFP-tagged Rab GTPases for co-rerouting with mCherry-FKB-P-TPD54 to dark MitoTrap. The collection of GFP-Rabs tested covers a range of membrane trafficking pathways (Yoshimura et al., 2007; Zhen and Stenmark, 2015; Wandinger-Ness and Ze- rial, 2014). The results of the screen are presented in Figure 8, with examples of positive and negative hits shown in Supplementary Figure S2. Significant co-rerouting was detected for 16 out of 43 Rabs. These positive hits were: Rab30, Rab25, Rab26, Rab45, Rab14, Rab11a, Rab12, Rab1a, Rab43, Rab1b, Rab10, Rab33b, Rab19, Rab33a, Rab37 and Rab2a (listed by descending effect size, Figure 8C). Some evidence for co-rerouting of Rab38, Rab5c and Rab35 was seen, although in any individual trial no clear difference was observed. The localization of the other 24 Rabs was unaffected by rerouting of TPD54 and was indistinguishable from GFP. Rab30 was the most efficiently co-rerouted Rab, with the post-rerouting sig-
Figure 4. Capture of small vesicles by rerouting TPD54 to mitochondria.
(A) Fluorescence microscopy images of mCherry-FKBP-TPD54 in HeLa cells. Cells expressing mCherry-FKBP-TPD54 and dark MitoTrap were fixed after no rapamycin application (Ctrl) or after 20 s, 5 min or 30 min of rapamycin addition (200 nM). The pictured cell was then imaged by electron microscopy. Scale bar, 10 µm. (B) Sample electron micrographs of the cells shown in A. Scale bar, 200 nm. (C) Segmented view of mitochondria (gray) and vesicles (purple) in the images shown in B. (D) Profiles of segmented vesicles from electron micrographs. All vesicles segmented from the control dataset are shown with a random sample from the treatment groups as a comparison. The sample size is in proportion to the capture of vesicles at the mitochondria (34, 320, 594, 1347 for ctrl, 20 s, 5 min and 30 min respectively). (E) Violin plot to show the diameter of vesicles imaged in each dataset. Spots represent individual vesicles. Marker shows the median. Box plot to show the number of vesicles captured per 1 µm of mitochondrial membrane. Spots show the number per micrograph in each dataset. Box shows the IQR, median and whiskers show 9th and 91st percentiles. Violin plot to show the fraction of mitochondrial membrane that is decorated with vesicles. Spots show individual mitochondria from the dataset. See Methods for details.

Figure 5. Fluorescence recovery after photobleaching (FRAP) analysis of GFP-TPD54.
(A) FRAP data for GFP-TPD54 in knock-in cells (endo), expressed GFP or GFP-TPD54 in parental cells. Lines and shaded areas show mean ± 1 s.d. Dotted line shows a double exponential function fitted to the average data. The fit coefficients are summarized in Table 1. Inset: FRAP data from different cells colored as indicated and displayed on the same axes range. (B) Recovery (mobile fraction) of individual fits to FRAP data plotted against $T_{1/2}$. Inset: Initial cellular fluorescence as a function of $T_{1/2}$. (C) Plot to show FRAP kinetics in individual cells. $\tau_{slow}$ is plotted against $T_{1/2}$, and marker size indicates the fraction recovered by the slow process. Markers represent individual cells and colors indicate the protein being imaged.
Figure 6. TPD54 co-reroutes dileucine motif-containing receptors only.

(A) Representative widefield micrographs of cells co-expressing mCherry-FKBP-TPD54, dark MitoTrap and the indicated CD8 construct. Rerouting was induced by 200 nM rapamycin. Cells were fixed, permeabilized and stained for total CD8. (B) Representative widefield micrographs showing co-rerouting of endogenous CIMPR detected by immunofluorescence with rerouting of mCherry-FKBP-TPD54 to dark MitoTrap by addition of 200 nM rapamycin. (C) Diagram of the revised interpretation of TPD54 rerouting experiments. When TPD54 is rerouted to the mitochondria, it reroutes the small vesicles with which it is associated. These vesicles contain cargo with a dileucine motif.

(D) Pulse label and timed vesicle capture experiments. Cells expressing CD8-EAAALL were surface labeled with Alexa488-conjugated anti-CD8 antibodies for 30 min, then incubated at 37 °C for the indicated time (min), treated with 200 nM rapamycin for 5 min and fixed. (E) Representative widefield micrographs from a pulse label and timed vesicle capture experiment. Inset, 5× zoom. Scale bars, 10 µm.

nal being 2.5 fold higher than before TPD54 rerouting (Supplementary Video SV5). The smallest effect that we could reliably detect was Rab2a, with a 1.4 fold increase.

If the relocalization of Rabs observed in the screen was the result of co-rerouting with TPD54, a correlation between the extent of rerouting for a Rab and TPD54 is predicted (Figure 8D). This was broadly true, with a positive correlation observed for almost all of our positive hits, and a low, flat relationship for negative Rabs. Rab14 was an exception. Here the relationship was high and flat indicating that Rab14 rerouting was maximal even after modest TPD54 rerouting.

We next performed a test of reciprocality, by asking if mCherry-TPD54 was co-rerouted to mitochondria when a GFP-FKBP-Rab was rerouted to dark MitoTrap using 200 nM rapamycin. We tested two positive hits from our screen, Rab11a and Rab25 as well as Rab7a, a negative Rab (Supplementary Figure S3). Rerouting of either Rab11a or Rab25 caused co-rerouting of TPD54, while rerouting Rab7a to mitochondria had no effect on TPD54 localization. Interestingly, we noticed that when Rab11a or Rab25 were...
TPD54 is a multi-Rab effector. We next tested if TPD54 binds directly to Rab GTPases and if it does so as an effector to the active, GTP-bound form (Pylypenko et al., 2018). To do this, we examined binding of recombinant MBP-TPD54-His to GST-Rab1a (positive hit) or GST-Rab6a (negative) in their GDP- or GppNHp-bound form. TPD54 bound to Rab1a, with preferential binding to the GppNHp-form. By contrast, the amount of binding to GST-Rab6a in either form was similar to GST alone (Figure 9A-B).

We also performed similar experiments in cells using the co-rerouting approach. Mutants that lock the Rab in an inactive (GDP-bound) or active (GTP-bound) state were made for GFP-tagged Rab30, Rab1a and Rab6a. These constructs were co-expressed with mCherry-FKBP-TPD54 and dark MitoTrap and their co-rerouting to mitochondria after rapamycin addition was compared to their wild-type counterparts (Figure 9C). Co-rerouting was seen for the active and wild-type forms of Rab30 and Rab1a, but no change in localization was observed for the inactive mutant of either Rab. No co-rerouting was observed for wild-type, inactive or active mutants of Rab6a. We analyzed the kinetics of TPD54 rerouting and Rab co-rerouting and found no evidence for interaction with the GDP-bound forms (Figure 9C). These observations support the idea that TPD54 binds directly to Rab in their active form, as an effector protein.

Discussion

This study shows that TPD54, an abundant protein in mammalian cells, regulates anterograde membrane traffic and the recycling of endocytosed cargo. We discovered TPD54 is found on numerous small functional vesicles throughout the cell and that it associates with many Rab GTPases. This...
Figure 8. A screen to identify Rab GT-Pases that are associated with TPD54.
(A) Quantification of the change in mitochondrial fluorescence intensity of GFP or GFP-Rabs 2 min after rerouting of mCherry-FKBP-TPD54 to dark MitoTrap with 200 nM rapamycin. Multiple independent experiments were completed (dots) across three independent trials. Black bars, mean ± sd. The mean ± sd for GFP (control) is also shown as a black line and gray zone, down the plot. Dunnett’s post-hoc test was done for each trial using GFP as control. Colors indicate if $p < 0.05$ in one, two, three trials or only when all the data were pooled. (B) Effect size and bootstrap 95% confidence interval of the data in A. (C) The plot in B is ranked from highest to lowest effect size. (D) Small multiple plots to show the correlation between the mCherry-FKBP-TPD54 rerouting and GFP-Rabs co-rerouting (gray dots), a line fit to the data (black) and a $y = x$ correlation (white).
Figure 9. TPD54 binds to active, GTP-bound Rab.

(A) Coomassie-stained SDS-PAGE gel of a GST pull-down assay to test for binding of MBP-TPD54-His to GST, GST-Rab1a or GST-Rab6a in GDP- or GppNHp-loaded states. SN, supernatant; B, beads. (B) Quantification of three GST pull-down experiments. MBP-TPD54-His bound to beads, normalized to the GST-Rab1a + GppNHp condition. (C) Testing for co-rerouting of the indicated GFP-Rab with mCherry-FKBP-TPD54 to MitoTrap induced by the addition of 200 nM rapamycin. Wild-type Rab30, Rab1a and Rab6a were tested alongside inactive (serine/threonine to asparagine) or active (glutamine to leucine) mutants. Insets, 3× zoom. Scale bars, 10 µm. Rerouting kinetics for TPD54 (red) and Rabbs (green) to the mitochondria or from the cytoplasm are shown to the right. Line and shaded region indicate mean ± s.e.m. for the following: Rab30 WT n = 14, \( \tau_{\text{red}} = 31.7, \tau_{\text{green}} = 38.8 \); QL n = 12, \( \tau_{\text{red}} = 25.1, \tau_{\text{green}} = 45.4 \); SN n = 11, \( \tau_{\text{red}} = 25.2, \tau_{\text{green}} = \text{n.d.} \); Rab1a WT n = 12, \( \tau_{\text{red}} = 37.1, \tau_{\text{green}} = 55.9 \); QL n = 11, \( \tau_{\text{red}} = 25.7, \tau_{\text{green}} = 59.0 \); SN n = 12, \( \tau_{\text{red}} = 26.6, \tau_{\text{green}} = \text{n.d.} \); Rab6a WT n = 12, \( \tau_{\text{red}} = 28.0, \tau_{\text{green}} = \text{n.d.} \). Single exponential fits to unweighted averaged data quoted in seconds.
promiscuous association of TPD54 with at least one-quarter of the Rabome is likely due to direct binding as an effector to the active, GTP-bound form of the Rab.

We used a knocksideways-based system to detect TPD54 binding partners and the cargoes transported by TPD54-associated vesicles (Robinson et al., 2010). There have been previous reports of vesicle capture at mitochondria following knocksideways of gadkin (Hirst et al., 2015), or by ectopic mitochondrial expression of Golgins (Wong and Munro, 2014). However, the vesicles captured by TPD54 rerouting are smaller, 29.9 ± 9.4 μm in diameter. Few types of vesicles are this small. For example, clathrin-coated vesicles are 50 to 100 μm, COPII-coated vesicles are 60 to 70 μm, and intra-Golgi transport vesicles are 70 to 90 μm (Vigers et al., 1986; Balch et al., 1994; Orci et al., 2000). The closest in size are intralumenal vesicles (ILVs) which range from 20 to 100 μm (Edgar et al., 2014; Raposo and Stoorvogel, 2013). Besides the mismatch in size profile, the inaccessibility and opposite orientation of ILVs makes them an ill-suited candidate for the vesicles captured by TPD54 rerouting. Our interpretation is that the vesicles we have captured are an overlooked class of intracellular vesicle which carry cargo from location A to location B. With no coat to distinguish them and with an unimposing size, these inconspicuous vesicles may have evaded study until now.

The captured vesicles certainly appear to be real, functional transport carriers because they had cargo and fusion machinery. There was selectivity in the cargo, which all contained dileucine-type endocytic motifs and in the R-SNARE complement of the vesicle. The reason for co-rerouting SNAREs could also be due of their dileucine motifs, although we saw no co-rerouting of VAMP4, which has one such motif. The captured vesicles are unlikely to be the result of vesiculization of larger membranes since we saw the capture of 30 nm vesicles after only 20 s of TPD54 rerouting and no further change in size at longer timepoints. Is it possible that TPD54 is on a wider variety of vesicles but that rerouting only captures the smallest of all TPD54-positive vesicles? One could imagine that smaller vesicles are captured more efficiently than larger ones. However, larger vesicles and even Golgi cisternae can be captured by mitochondria under different experimental conditions (Hirst et al., 2015; Wong and Munro, 2014; Shin et al., 2017; Dunlop et al., 2017), suggesting that TPD54 is predominantly localized to these small vesicles and that this is the reason for their more efficient capture. This interpretation is supported by live cell imaging and FRAP analyses which suggest that the majority of TPD54 is on the small vesicles.

Using rerouting as a screening method to identify Rabs could be useful for the study of other effector proteins. A previous study used a similar method to demonstrate that the recruitment of Rab5a by its GEF (Rabex-5) depends on GEF activity (Blümer et al., 2013). The recruitment was much slower than in our study. There, co-rerouting was on the timescale of several minutes, whereas TPD54 rerouting and co-rerouting of Rabs occurred in seconds. The slow recruitment might be specific to Rabex-5 or to Rab5 and its localization. We identified Rab5c as a TPD54 interactor by proteomics but it was only a borderline hit in our screen. This might mean that co-rerouting of Rab5-vesicles is inefficient. Our screen certainly only reports on Rabs that have the ability to be co-rerouted, a caveat which may mean that we have underestimated the promiscuity of TPD54. An alternative interpretation to explain the difference is that effectors can influence Rab localization more effectively than GEFs. It is unusual for an effector to bind multiple Rabs. Indeed, the ability of the Rabs to finely tune all parts of the trafficking pathways is in part due to the specificity of their effectors (Stenmark, 2009; Pylypenko et al., 2018). There are a few effectors that can bind more than one Rab GTPase. They can do this either by having several Rab binding sites or by virtue of a single site that can bind several Rabs. For example, Rab Coupling Protein (RCP/RAB11FIP1) has one Rab-binding domain through which it can bind Rabs of the
same subgroup (Rab11a, Rab25 and Rab14 in Group IV) (Lall et al., 2015; Klöpper et al., 2012). Whereas GCC185, a large, 1684 residue Rab effector, binds Rab5 from three different subgroups (I, III and IV), but this occurs via six distinct domains on GCC185 (Hayes et al., 2009). To our knowledge, only OCRL1 binds to multiple Rab5 (six Rab5 from four different subgroups), via a single domain of 123 residues (Hou et al., 2011). Our data suggest that TPD54 follows the OCRL1 precedent of being a promiscuous Rab effector, presumably also via a single domain. TPD54 is a small, 206 residue protein and we found association with at least 16 Rab GTPases from three different sub-groups. Effectors generally bind Rab5 via one or two helices (Pylypenko et al., 2018; Rai et al., 2017). TPD54 has a predicted coiled-coil domain towards the N-terminus, through which it can homodimerize or even heterodimerize with other TPD52-like proteins (Byrne et al., 1998). This could mean that TPD54 as a homodimer binds two Rab5, in a 2:2 configuration described for other effectors such as RCP. These two Rab5 would most likely be of the same type as they would be co-resident on the same membrane. One attractive hypothesis is that, by being able to dimerize, TPD54 allows the aggregation of Rab5 to create functional sub-domains on vesicles and organelles (Sönntichsen et al., 2000; Pfeffer, 2010). Our Golgi dispersal phenotype in cells lacking TPD54 suggests that the effector function of TPD54 is either vesicle tethering or fusion. Vesicle tethering proteins tend to be large and capable of spanning significant distances in the cell, therefore we favor the interpretation that TPD54 promotes fusion. In support of this, TPD53/TPD52L1 was previously enhance the interaction between SNAREs syntaxin 1 and VAMP2 (Proux-Gillardeaux et al., 2003).

TPD52-like proteins were first identified due to their overexpression in cancer and the expression of TPD52 correlates with a more aggressive metastatic breast cancer phenotype (Shehata et al., 2008). Having defined a role for TPD54 in anterograde membrane traffic and cargo recycling through functional studies and by our Rab screen, the next question is if dysregulation of these functions contributes to cancer formation or progression. We note that TPD54 associates with Rab11a and Rab25, two Rab5 which are implicated in cancer cell migration and invasion due to their role in integrin recycling (Caswell et al., 2007; Muller et al., 2009). Pursuing the role of TPD52-like proteins in membrane traffic in a cancer context is therefore an exciting direction for the future.

Methods

Molecular biology. GFP-TPD54, mCherry-TPD54 and FLAG-TPD54 were made by amplifying TPD54 by PCR from human Tumor protein D54 (IMAGE clone: 3446037) and inserted into eGFP-C1, pmCherry-C1 or pFLAG-C1 via XhoI-MfeI. GFP-TPD52 was made by amplifying TPD52 by PCR from Tumor protein D52 (IMAGE clone: 3916870) and inserted into pEGFP-C1 via XhoI-MfeI. GFP-FKBP-TPD54 was made by ligating a XhoI-BamHI fragment from mCherry-TPD4 into pEGFP-FKBP-C1. This plasmid was converted to mCherry-FKBP-TPD54 by cutting at BamHI-MfeI and inserting into pmCherry-C1. MBP-TPD54-His was cloned by inserting TPD54 in pMalPre-His vector via FseI and EagI.

YFP-MitoTrap and the CD8-chimeras were gifts from Scottie Robinson (University of Cambridge, UK) and mCherry-MitoTrap was previously described (Cheeseman et al., 2013). The dark MitoTrap (pMito-dCherry-FRB) has a K70N mutation in mCherry (Wood et al., 2017). The GFP-Rab constructs were a gift from Francis Barr (University of Oxford, UK), except for GFP-Rab1a and GFP-Rab5c. GFP-Rab1a and GFP-Rab5c were made by amplifying human Rab1a or Rab5c (Rab1a: Addgene: #46776, Rab5c: GeneArt synthesis) by PCR and inserting the genes in pEGFP-C1 via SacI-KpnI. Rab mutants GFP-Rab1a S25N and Q70L, GFP-Rab6a T27N and Q66L, and GFP-Rab30 T23N and Q68L were generated for GFP-Rab1a, GFP-Rab6a or GFP-Rab30 using site-directed mutagenesis. GFP-FKBP-Rab11 and GFP-FKBP-Rab25 were a gift from Patrick Caswell (University of Manchester, UK), and GFP-FKBP-Rab7a was made by inserting Rab7a in pEGFP-FKBP-C1 via SacI-SalI. The GST-Rab1a and GST-Rab6a constructs were cloned by cutting Rab1a and Rab6a from GFP-Rab1a and GFP-Rab6a with SalI and NotI and inserting into pGEK-6P1. Plasmid to express mCherry-OCRL1 was a gift from Martin Lowe (University of Manchester, UK). GalT-mCherry was made by cutting GalT via BamHI and MfeI from GalT-CFP (gift from Ben Nichols, MRC Laboratory of Molecular Biology, UK) and inserted into pmCherry-N1. GFP-VAMP2 and mCherry-VAMP2 were made by amplifying VAMP2 from synaptopHluorin (gift from James Rothman, Yale School of Medicine, USA) and inserted into pEGFP-C1 or mRFP-C1 via HindIII and EcoRI. FLAG-TACC3 was made by amplifying TACC3 (IMAGE clone: 6148176 BC106071) by PCR and inserted into pFLAG-C1 via XmaI and MluI. Other plasmids such as mCherry-LCa were available from previous work (Hood and Royle, 2009). SBP-EGFP-E-cadherin and APPL1-mCherry were obtained from Addgene (#65292 and #27683, respectively).

Cell culture. HeLa cells (HPA/ECACC #93021013) were maintained in DMEM supplemented with 10% FBS and 100 U/ml penicillin/streptomycin at 37 °C and 5% CO2. RNA interference was done by transfecting 100 nM siRNA (TPD54: GUCCUAACCUGUACCAGAU) with Lipofectamine 2000 (Thermo Fisher Scientific) according to the manufacturer’s protocol. For DNA plasmids, cells were transfected with a total of 300 ng DNA (per well of a 4-well dish) using 0.75 μl Genejuice (Merck Millipore) following the manufacturer’s protocol. Cells were imaged 1 d after DNA transfection and 2 d after siRNA transfection. For two rounds of RNAi, HeLa cells were transfected with the TPD54-targeting siRNA for 48 h and transfected again with the siRNA for an additional 72 h. The GFP-TPD54 CRISPR knock-in HeLa cell line was generated by transfecting the Cas9n D10A nickase plasmid containing the guide pairs (guide 1: 5’ACCGGTGTCCGCGGCGCTCTAT, guide 2: 5’GCCGGAACATG-
GACTCCGC) and the repair template. 9 d post-transfection, GFP-positive cells were selected by FACS and isolated. Clones were validated by western blotting and genome sequencing (sequencing primers: 5’CAGTTTTCGCCC-TATCAGGTTAGTC, 5’GAACACACCTCGGAA CG-GTC, 5’CAGTTTGTGCCAGAGATTTG, 5’CAACTACAAGACCCGCGCCGAG).
The TPD54 knock-out HeLa cell lines were generated by transfecting the Cas9 plasmid containing one of the three guide pairs (guide 1: 5’caccgtTGCCGATTACGAAACCGCG, guide 2: 5’caccgTTTTGTAATTCCGCGATCGCA, guide 3: 5’caccACCCGTGTCGCGGGCGCTAT). The transfected cells were selected with 1 µg/ml puromycin 24 h post-transfection. Clones were isolated and validated by western blot and sequencing genomic DNA.

Biochemistry. For western blot analysis, antibodies used were rabbit anti-TDP54 (Dundee Cell products) 1:1000, mouse anti-α-tubulin (Abcam: ab7291) 1:10000, mouse anti-GFP clones 7.1 and 13.1 (Sigma Roche: 118144600010) 1:1000 and mouse anti-clathrin heavy chain TD.1 (hybridoma) 1:1000.

For the purification of GST, GST-Rab1a and GST-Rab6a, transformed BL21 Escherichia coli were grown and proteins induced by addition of 0.5 mM IPTG at 25 °C overnight. Cells were harvested, resuspended in lysis buffer pH8.0 (50 mM Tris, 150 mM NaCl, protease inhibitor tablet (Roche), 0.2 mM PMSF) and lysed by sonication (6×10 s pulses). The bacterial debris were pelleted at 34000 g for 30 min at 4 °C. Supernatant was loaded onto a GSTrap FF Columns (GE Healthcare) and washed with high-salt wash buffer pH8.0 (50 mM Tris, 500 mM NaCl, protease inhibitor tablet, 0.1 mM PMSF). Proteins were eluted with elution buffer pH 8.0 (50 mM Tris, 150 mM NaCl, 50 mM glutathione).

MBP-TDP54-His was purified similarly except induction was at 37 °C for 4 h. Cleared supernatant was loaded onto a HisTrap Fast Flow Column (GE Healthcare), proteins were washed with buffer A (20 mM Tris pH8.0, 400 mM NaCl, 0.5 mM CaCl2, 10 mM imidazole) and eluted with buffer B (20 mM Tris pH8.0, 400 mM NaCl, 0.5 mM CaCl2, 400 mM imidazole).

For the Rab binding assay, 3 µM of purified GST, GST-Rab1a or GST-Rab6a was pre-loaded with 300 µM GDP or GppNHp for 90 min at 20 °C in binding buffer (25 mM HEPES, 150 mM KCl, 5 mM MgCl2 pH 7.4). These were incubated with 3 µM of purified MBP-TDP54-His for 90 min at 20 °C. The GST/TDP54 mixture was then incubated with glutathione-Sepharose 4B beads for 30 min at 20 °C with rotation and then beads were pelleted. The supernatant was collected and the beads were washed 5× with binding buffer containing 0.1% Tween 20. Proteins were eluted from the beads in Laemmli buffer and boiled. Supernatant and bead fractions were loaded on a 4-15% polyacrylamide gel and visualized by staining with Coomassie brilliant blue.

Immunoprecipitation. Two 10 cm dishes of confluent HeLa cells expressing either GFP or GFP-TPD54 were used for each condition (10 µg DNA transfected per 10 cm dish). Cells were lysed in lysis buffer (10 mM Tris-HCl pH 7.5, 150 mM NaCl, 0.5 mM EDTA, 0.5% NP-40, protease inhibitors (Roche)). The lysate was then incubated for 1 h with GFP-Trap beads (ChromoTek), washed once with exchange buffer (10 mM Tris-HCl pH7.5, 150 mM NaCl, 0.5 mM EDTA) and three times with wash buffer (10 mM Tris-HCl pH7.5, 500 mM NaCl, 0.5 mM EDTA). The immunoprecipitations were run on a 4-15% polyacrylamide gel until they were 1 cm into the gel. The columns were then cut and sent for mass spectrometry analysis to the FingerPrints Proteomics Facility (University of Dundee, UK). Protein scores from four experiments were used to make the volcano plot in IgorPro.

Immunofluorescence. HeLa cells grown on cover slips were fixed at room temperature (RT) with 3% paraformaldehyde, 4% sucrose in PBS during 15 min and permeabilized at RT in 0.1% saponin for 10 min (for all staining, unless stated otherwise). For LAMP1 staining, cells were fixed and permeabilized with ice-cold methanol at −20 °C for 10 min. For CD8, TGN46 and FLAG staining, cells were fixed in 3% paraformaldehyde, 4% sucrose in PBS during 15 min and permeabilized at RT in 0.5% Triton-X100 in PBS for 10 min. Cells were then blocked in 3% BSA in PBS for 1 h. Cells were incubated for 1 h at RT with primary antibody used as follows: mouse anti-EEA1 (BD Biosciences: 610457) 1 µg/mL, rabbit anti-LAMP1 (Cell Signaling: 9091) 1:200, sheep anti-TGN46 (AbD Serotec: AHP500G) 1.25 µg/mL, mouse anti-CD8 (Biorad: MCA1226GA) 10 µg/mL, rabbit anti-CIMPR (Thermo Fisher: PA3-850) 1:500, mouse anti-FLAG M2 (Sigma: F1804). Anti-SNARE antibodies were a gift from Andrew Peden (University of Sheffield, UK): Rabbit anti-VAMP3 (1:200), Rabbit anti-VAMP4 (1:500), Rabbit anti-VAMP7 (1:50), Rabbit anti-VAMP8 (1:100), Rabbit anti-STX6 (1:200), Rabbit anti-STX7 (1:400), Rabbit anti-STX8 (1:100), Rabbit anti-STX10 (1:50), Mouse anti-STX16 (1:200), described in (Gordon et al., 2010). Cells were washed three times with PBS for 5 min and incubated during 1 hour at RT with AlexaFluor (Invitrogen) secondary antibodies. To co-reroute the CD8-EAAALL chimera after timed incubation with anti-CD8 antibodies, cells were labelled with 10 µg AlexaFluor 488- conjugated anti-CD8 antibodies (AbD Serotec, MCA1226A488) at 4 °C for 30 min. Cells were then incubated at 37 °C in warm growth medium for the indicated time points. Rerouting was done by adding 200 nM rapamycin at 37 °C. Cells were fixed and mounted after 5 min, as described above.

For the transferrin assay, HeLa cells grown on coverslips were serum starved for 30 min at 37 °C, then incubated at 4 °C for 30 min with 25 µg/mL of AlexaFluor 488- conjugated transferrin (Thermo Fisher Scientific, 11550756). The coverslips were then dipped in dH2O, placed in warm growth medium and incubated at 37 °C for 5 to 75 min to allow internalization and recycling, before fixation.

Confocal Microscopy. Cells were grown in 4-well glass-bottom 3.5 cm dishes (Greiner Bio-One) and media ex-
changed for Leibovitz L-15 CO₂-independent medium for imaging at 37 °C on a spinning disc confocal system (Ultra-View Vox; PerkinElmer) with a 100 x 1.4 NA oil-immersion objective. Images were captured using an ORCA-R2 digital CCD camera (Hamamatsu) following excitation with 488 nm and 561 nm lasers. For the RUSH assay, SBP-EGFP-E-cadherin was released from the ER by adding a final concentration of 40 μM D-Biotin (Sigma) in L-15 medium. Images were captured at an interval of 2 min. Rerouting of mCherry-FKBP-TPD54 to the mitochondria (dark MitoTrap) was induced by addition of 200 nM rapamycin (Alfa Aesar). Rerouting kinetics experiments were measured by recording movies of 150 s (1 frame per sec), where rapamycin is added after 10 s. The kinetics of mCherry-FKBP-TPD54 rerouting to mitochondria was similar in cells with or without depletion of endogenous TPD54. For the Rab GTPase co-rerouting experiments, an image pre-rapamycin and an image 2 min post-rapamycin was taken of live cells. For the FRAP experiment, a region of 6.69 μm by 10.76 μm was bleached using a 488 nm for five cycles of 100 ms. Images were captured at the highest frame rate possible (0.1775 s).

**Correlative light-electron microscopy.** Following transfection, cells were plated on gridded dishes (P35G-1.5-14-CGRD, MatTek Corporation, Ashland, MA, USA). Light microscopy was done using a Nikon Ti epifluorescence microscope, a heated chamber (OKOlab) and CoolSnap MYO camera (Photometrics) using NIS elements AR software. During imaging, cells were kept at 37 °C in Leibovitz L-15 CO₂-independent medium supplemented with 10% FBS. Transfected cells were found and the grid coordinate containing the cell of interest was recorded at low magnification. Live cell imaging was done on a cell-by-cell basis at 100x. During imaging, 200 mM (final concentration) rapamycin was added for variable times before the cells were fixed in 3% glutaraldehyde, 0.5% paraformaldehyde in 0.05 M phosphate buffer pH 7.4 for 1 h. Aldehydes were quenched in 50 mM glycine solution and thoroughly washed in H₂O. Cells were post-fixed in 1% osmium tetroxide and 1.5% potassium ferrocyanide for 1 h and then in 1% tannic acid for 45 min to enhance membrane contrast. Cells were rinsed in 1% sodium sulphate then twice in H₂O before being dehydrated in grade series ethanol and embedded in EPON resin (TAAB). The coverslip was removed from the polymerized resin and the grid was used to relocate the cell of interest. The block of resin containing the cell of interest was then trimmed with a glass knife and serial 70 nm ultrathin sections were taken using a diamond knife on an EM UC7 (Leica Microsystems) and collected on formvar coated hexagonal 100 mesh grids (EM resolutions). Sections were post-stained with Reynolds lead citrate for 5 min. Electron micrographs were recorded using a JEOL 1400 TEM operating at 100 kV using iTEM software.

**Image analysis.** For analysis of RUSH movies, a region of interest (ROI) was drawn around the cell and around the Golgi apparatus in FIJI. The area, mean pixel intensity and integrated density was measured from these ROIs to get fluorescence intensity ratios. Data were processed in IgorPro using custom scripts. Briefly, a logistic function (equation 1) was fitted to the data using the start of the movie and two frames after the maximum value as limits for the fit.

\[
\begin{align*}
    f(x) &= y_0 + \frac{(y_{max} - y_0)}{1 + \left( \frac{x - a}{b} \right)^n} \\
\end{align*}
\]

The value for \(x_{1/2}\) in minutes was used for the \(T_{1/2}\) for ER-to-Golgi. The corresponding \(y\) value for \(x_{1/2}\) was used to find the \(T_{1/2}\) for ER-to-PM. A line of best fit from \(y_{max}\) to the end of the trace was found using \(f(x) = a + bx\) and the \(y\) value corresponding to \(y_{x_{1/2}}\) found. The Golgi transit time is taken from the difference between the two \(T_{1/2}\) values.

Rerouting kinetics were quantified by averaging the pixel intensity of 10 ROIs of 5x5 pixels on mitochondria and 4 ROIs of varying size in the cytoplasm per cell, in the red, or red and green channels, throughout the duration of the movies. The images were corrected for photobleaching using the simple ratio method prior to measuring pixel intensity for all ROIs. The intensity and time data was fed into IgorPro, and a series of custom-written functions processed the data.

For vesicle capture analysis, electron micrographs were manually segmented in IMOD using a stylus by a scientist blind to the experimental conditions. The coordinates corresponding to contours and objects were fed into IgorPro using the output from model2point. All coordinates were scaled from pixels to real-world values and the vesicle diameters calculated using the average of the polar coordinates around the vesicle center for each vesicle, along with other parameters. As a metric for vesicle capture, the length of the mitochondrial perimeter was measured and used to express the vesicle abundance per image (vesicles per 1 μm). The intersection of an area corresponding to the vesicles, dilated by 15 nm and the mitochondrial perimeter was used to express the fraction of mitochondrial perimeter that was decorated with vesicles.

For the Rab screen, co-rerouting of Rab GTPases was quantified by averaging for each cell, the pixel intensity in the green channel in 10 ROIs of 10x10 pixels on the mitochondria, before and after rapamycin. This mitochondrial intensity ratio (\(F_{post}/F_{pre}\)) for every Rab was compared to the ratio of GFP in TPD54 rerouted cells. Estimation statistics were used to generate the difference plot shown in Figure 8B. The mean difference is shown together with bias-corrected and accelerated (BCa) 95% confidence intervals calculated in R using 1 x 10⁵ bootstrap replications.

For FRAP analysis, an ImageJ macro was used to define and measure the GFP intensity (MPD) in the FRAP region, background and whole cell. These data and timestamps from OME were fed into IgorPro for processing. The background-subtracted intensities for the FRAP region and whole cell were used to calculate a ratio (to correct for bleaching of molecules induced by the procedure). These values were paired with the timestamps, scaled so that the intensity after bleach was 0 and an average of the first five images minus was 1, and then an interpolated average was created. Fits to individual traces were also calculated using a script. Double exponential function was used for fitting since this gave better
fits than a single exponential, particularly for GFP-TPD54, and so that all conditions were fitted in the same way for comparison.

For the Rab binding assay, ROIs of the same area were drawn over the MFP-TPD54-His bands and over the background in the same lanes, and the raw integrated density was measured. The background value was subtracted from the band value and the data was normalized to the MFP-TPD54-His band in the GST-Rab1a (GppNHp) condition. To measure Golgi dispersion, the area of a convex hull around the cell TGN46 signal was expressed as a fraction of the area of an ROI drawn around the cell.

The phylogenetic tree was generated with phylogeny.fr, using the "one click" mode (Dereeper et al., 2008), and was visualized with phylo.io.

All figures were made with either FIJI or Igor Pro 8 (WaveMetrics) and assembled using Adobe Illustrator.

**Data and software availability.** The data for proteomics, volcano plot, FRAP data and EM segmentation coordinates are available together with code and scripts for analysis at https://github.com/quantixd/TPD54 | Royle, 2018.

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**COMPETING FINANCIAL INTERESTS**

The authors declare no conflict of interest.

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Figure S1. Characterization of the GFP-TPD54 knock-in cell line

(A) Gene map of TPD52L2 and location of GFP-tagging. (B) FACS plot. GFP-positive cells in the indicated gate were recovered and characterized. (C) Representative confocal micrograph of an example GFP-TPD54 knock-in clone. Scale bar, 10 µm. (D) Clones were validated by western blot. Cells overexpressing GFP-TPD54, parental HeLa cells, TPD54-depleted cells and three different clones are shown. Clone 35 exhibited the desired band profile. A single GFP-TPD54 band detected by blotting for TPD54 and GFP, with no untagged TPD54. Tubulin is shown as a loading control, note that one-tenth of the GFP-TPD54 transfected sample was loaded. (E) Sequencing the TPD52L2 locus in clone 35. Two bands were amplified using primers flanking the integration site. The first sequence shows integration of monomeric GFP between the homology arms, giving GFP-TPD54. The second sequence shows that clone 35 is null at the other allele.
Figure S2. Examples of co-rerouting data from the screen

Representative micrographs showing the co-rerouting of GFP-Rab14, but not GFP or GFP-Rab9, after a rerouting of mCherry-FKBP-TPD54 to dark Mitotrap by addition of 200 nM rapamycin. Note that GFP-Rab14 localization is unaffected by rerouting of mCherry-FKBP. Insets, 3× zoom. Scale bar, 10 µm.
Figure S3. Reciprocal rerouting of Rabs and TPD54

Two positive hits (Rab11a and Rab25) and a negative Rab (Rab7a) were tested for TPD54 co-rerouting. Micrographs of cells before and after rerouting the indicated GFP-FKBP-Rab to dark MitoTrap in cells also expressing mCherry-TPD54. Insets, 3× zoom. Scale bar, 10 μm.
Figure S4. Cellular localization and phylogenetic tree of the Rabs used in the screen
(A) Schematic diagram showing the cellular pathways on which the Rab GTPases operate. Rabs are represented by their number. Red and gray numbers indicate positive and negative Rab hits, respectively. (B) Phylogenetic tree of the Rab GTPases in the screen. Black branches indicate Rabs that interact with TPD54. Colored circles are as described in Figure 8.
Figure S5. Targeted disruption of TPD54 gene in HeLa cells using CRISPR/Cas9

(A) Three guides were designed to target the TPD52L2 locus. (B) Single cell clones were isolated and screened by western blotting. Two clones, 2.2 and 2.4 showed loss of TPD54 expression. (C) Sequencing of PCR amplicons using primers flanking the CRISPR/Cas9 targeting site revealed disruption of the locus in clones 2.2 and 2.4. Sequencing of the top five most similar PAM sequences in the genome showed no change from the parental sequence.

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Supplementary Videos

**Figure SV1. SBP-EGFP-E-Cadherin RUSH imaging in control cells.**
Live cell confocal microscopy of RUSH assay, biotin is added at time 0. Still images from this movie are shown in Figure 2.
Time, hh:mm. Scale bar, 10 µm.

**Figure SV2. SBP-EGFP-E-Cadherin RUSH imaging in TPD54-depleted cells.**
Live cell confocal microscopy of RUSH assay, biotin is added at time 0. Still images from this movie are shown in Figure 2.
Time, hh:mm. Scale bar, 10 µm.

**Figure SV3. Rerouting mCherry-FKBP-TPD54 to mitochondria in cells co-expressing MitoTrap.**
Live cell confocal microscopy of rerouting assay, mCherry-FKBP-TPD54 is rerouted to MitoTrap using rapamycin 200 nM, applied at 10 s. Still images from this movie are shown in Figure 3.
Time, mm:ss. Scale bar, 10 µm.

**Figure SV4. Fluorescence recovery after photobleaching (FRAP).**
Fluorescence recovery after photobleaching (FRAP) of GFP (left) or GFP-TPD54 (right) expressed in HeLa cells. Bleach area is a rectangle inset by 0.8 µm.
Time is indicated. Scale bar, 5 µm.

**Figure SV5. Co-rerouting of a GFP-Rab with mCherry-FKBP-TPD54.**
Live cell confocal microscopy of rerouting assay. Co-rerouting of GFP-Rab30 (left, green) with mCherry-FKBP-TPD54 (middle, red) to dark MitoTrap (not shown) using rapamycin 200 nM, applied at 10 s.
Time, mm:ss. Scale bar, 10 µm.