Retrograde transport of CDMPR depends on several machineries as analyzed by sulfatable nanobodies

Dominik P Buser, Gaëtan Bader, Martin Spiess

Retrograde transport protein from the cell surface and endosomes to the TGN is essential for membrane homeostasis in general and for the recycling of mannose-6-phosphate receptors (MPRs) for sorting of lysosomal hydrolases in particular. We used a nanobody-based sulfation tool to more directly determine transport kinetics from the plasma membrane to the TGN for the cation-dependent MPR (CDMPR) with and without rapid or gradual inactivation of candidate machinery proteins. Although knockdown of retromer (Vps26), epsinR, or Rab9a reduced CDMPR arrival to the TGN, no effect was observed upon silencing of TIP47. Strikingly, when retrograde transport was analyzed by rapamycin-induced rapid depletion (knocksideways) or long-term depletion by knockdown of the clathrin adaptor AP-1 or of the GGA machinery, distinct phenotypes in sulfation kinetics were observed, suggesting a potential role of GGA adaptors in retrograde and anterograde transport. Our study illustrates the usefulness of derivatized, sulfation-competent nanobodies, reveals novel insights into CDMPR trafficking, and further outlines that the selection of machinery inactivation is critical for phenotype analysis.

DOI 10.26508/lsa.202101269 | Received 22 October 2021 | Revised 8 March 2022 | Accepted 9 March 2022 | Published online 21 March 2022

Introduction

Retrograde transport of proteins from the cell surface and endosomes to the TGN is critical for membrane homeostasis and to retrieve components of anterograde transport machineries. Proteins recycled back to the TGN comprise transport receptors for lysosomal hydrolases, processing enzymes, SNAREs (soluble N-ethylmaleimide–sensitive fusion factor attachment receptors), nutrient transporters, and a subset of other intracellular transmembrane proteins with diverse functions (Bonifacino & Rojas, 2006; Johannes & Popoff, 2008). In addition, extracellular bacterial and plant toxins exploit the retrograde transport machineries of target cells. One of the most thoroughly studied cargoes retrieved from endosomes to the TGN are the cation-dependent and cation-independent mannose-6-phosphate receptors (CDMPR and CIMPR), involved in efficient anterograde transport of lysosomal acid-hydrolases from the TGN to endosomes (Ghosh et al, 2003a). After cargo unloading in the mildly acidic endosomal environment, MPRs are recycled to the TGN apparently by several retrograde transport machineries from both early/recycling and late endosomes (Sandvig & van Deurs, 2002; Bonifacino & Rojas, 2006; Pfeffer, 2009; McKenzie et al, 2012). The estimated copy number of CDMPR and CIMPR in HeLa cells is ~660,000 and ~310,000 per cell, respectively (Itzhak et al, 2016). Recently, cell surface MPR has been recognized as an efficient and potential therapeutic platform for targeted degradation of extracellular and transmembrane proteins using hexasaccharide–anti-target conjugates by shuttling the target to lysosomes, whereas MPR is retrieved to the TGN or plasma membrane (Banik et al, 2020). Understanding transport routes and sorting machineries involved in MPR trafficking is central to modulate this pathway.

The most prominent sorting machinery reported to mediate transport of MPRs to the TGN is the retromer complex, a pentameric protein assembly comprising Vps (vacuolar protein sorting) and SNX-BAR (sorting nexin-Bin/Amphipathysin/Rvs) subunits (Bonifacino & Hurley, 2008; Seaman, 2012; Gallon & Cullen, 2015). The core complex, termed retromer, consists of the heterotrimer Vps26-Vps29-Vps35 that transiently associates with the tubulation subcomplexes composed of SNX1 or SNX2, and SNX5 or SNX6. Other SNX-BAR proteins, including SNX3, have been shown to mediate endosome-to-TGN transport of Wntless (WLS) (Belenkaya et al, 2008; Harterink et al, 2011). The precise sites from which retromer complexes operate remain to be defined: Vps35 of retromer was shown to be recruited by Rab7a, a marker of late endosomes, but SNX-BAR components of the multimeric complex bind via their Phox homology (PX) domain to phosphatidylinositol 3-phosphate (PI3P), a phospholipid enriched on early endosomes (Cozier et al, 2002; Carlton et al, 2004, 2005; Rojas et al, 2008; Seaman et al, 2009). It was thus proposed that MPR sorting by retromer complex is a progressive process coupled to endosomal maturation during the Rab5–to-Rab7 switch (Rojas et al, 2008). Some uncertainty also exists about cargo recognition by Vps and/or SNX-BAR subunits. Previously, it was reported that Vps subunits serve as cargo adaptors for the cytoplasmic domain of CIMPR (Nothwehr et al, 2000; Seaman, 2007; Fjorback et al, 2012; Lucas et al, 2016; Cui et al, 2019; Suzuki et al, 2019). Two recent independent studies rather suggest that SNX1/2 and SNX5/6 mediate cargo recognition and
retrieval of CiMPR (Kvainickas et al, 2017; Simonetti et al, 2017). Their results not only showed that SNX-BAR dimers associate with a WLM motif in the cytoplasmic tail of CiMPR, but also that Vps35 depletion, unlike depletion of SNX-BARs, did not cause receptor misdistribution from juxtanuclear to peripheral compartments. This observation is in disagreement with previous results showing a prominent mislocalization of CiMPR to endosomes upon silencing of Vps26 or Vps35 (Arighi et al, 2004; Seaman, 2004).

A further pathway for retrograde transport of MPRs was described to involve Rab9a and the adaptor TIP47 (tail-interacting protein of 47 kD). Using a cell-free system, it was shown that Rab9a recruits TIP47 to late endosomes and that interference with GTPase-effector function resulted in severe impairment of transport of MPRs (Lombardi et al, 1993; Diaz & Pfeffer, 1998). However, TIP47 was since identified to be a component of lipid droplets involved in their biogenesis (Bulankina et al, 2009) and an additional role in retrograde transport was not independently reproduced (Mediges & Schu, 2003).

Another mechanism for MPR retrieval to the TGN involves clathrin-coated vesicles (CCVs) with the adaptor protein (AP)–1 complex and/or epsinR. AP-1 has a generally accepted role in anterograde transport of MPRs from the TGN to endosomes in cooperation with GGA (Golgi-localized, γ-adaptin ear-containing, Arf-binding) proteins (Doray et al, 2002; Ghosh & Kornfeld, 2004; Sanger et al, 2019). Unlike retromer complexes, AP-1 has a dual distribution both at the TGN and on early endosomes (Le Borgne et al, 1996; Seaman et al, 1996; Meyer et al, 2000). Inactivation of AP-1 was found to result in a dispersed MPRL distribution pattern towards the periphery of cells (Meyer et al, 2000; Robinson et al, 2010), similar to the phenotype of retromer complex inactivation, suggesting a role in retrograde transport. EpsinR, an interactor of AP-1, was also shown to have a role in CiMPR retrieval to the TGN (Hirst et al, 2004; Saint-Pol et al, 2004), although receptor distribution did not significantly change in epsinR-depleted cells. It is puzzling why epsinR and AP-1 have different effects on MPR localization when depleted, whereas they seem to depend on each other for incorporation into CCVs, depletion of one reducing the CCV content of the other (Hirst et al, 2004, 2015).

GGAs also localize to both TGN and endosomes (Boman et al, 2000; Ghosh et al, 2003b). Yet, they have been implicated mainly in anterograde transport. Rapid depletion of GGA2 by knocksideways specifically depleted lysosomal hydrolases and their receptors (MPRs and sortillin) from CCV contents, whereas knocksideways of AP-1 affected also a number of SNAREs and additional membrane proteins (Hirst et al, 2012). Depletion of lysosomal hydrolases from CCVs was more efficient upon inactivation of GGA2 and depletion of their receptors more efficient upon inactivation of AP-1. This result suggested a role of GGAs primarily in anterograde transport and of AP-1 in both directions.

To analyze retrograde transport machinery, most studies used fluorescence microscopy to monitor changes in MPR localization relative to TGN-resident or endosomal markers by statistical steady-state image analysis. A few laboratories imaged antibody uptake to follow retrograde transport from the cell surface to the TGN (e.g., Breusegem and Seaman [2014]), whereas Johannes and colleagues used sulfation as a specific modification of the trans-Golgi/TGN to probe Golgi arrival of Shiga toxin B-chain (STxB) tagged with a sulfation motif or of antibodies derivatized with sulfatable peptides (Saint-Pol et al, 2004; Popoff et al, 2009). However, the disadvantage of conventional divalent antibodies is that they are rather large and can crosslink their antigens and thus potentially alter their trafficking. In contrast, monomeric protein binders, such as nanobodies, are monovalent and small. Nanobodies are easily derivatized with sequence tags and fluorescent or enzymatic protein domains and can be produced in bacteria.

To study retrograde traffic, we have previously established a versatile toolkit of functionalized anti–GFP nanobodies (Buser et al, 2018; Buser & Spiess, 2019). In particular, we generated nanobodies containing tyrosine sulfation (TS) sites to monitor their arrival in the trans-Golgi/TGN. Using cell lines stably expressing EGFP-CiMPR or EGFP-CiMPR, we determined the transport kinetics of these receptors from the cell surface to the TGN. In addition, we used the knocksideways system developed by Robinson et al (2010) to analyze the contribution of AP-1 upon rapid depletion. The system is based on rapamycin-induced heterodimerization between the γ-subunit of AP-1 fused to FKBP12 (FK506-binding protein of 12 kD) and the FKBP-rapamycin-binding domain (FRB) of mammalian target of rapamycin anchored in the outer mitochondrial membrane as a trap (Mitotrap). Upon rapid inactivation of AP-1, a robust reduction of sulfation kinetics by approximately on third was observed, confirming a significant contribution of AP-1/CCVs in retrograde transport of MPRs (Buser et al, 2018).

In the present study, we applied this tool of sulfatable nanobodies to analyze plasma membrane-to-TGN transport kinetics of CiMPR to define the contribution of individual different sorting machineries in parallel on retrograde transport in living cells. We could confirm retrograde transport activity of retromer and epsinR as well as a considerable involvement of Rab9a, but not of TIP47. Unexpectedly, silencing of GGA1-3 using RNAi also reduced CiMPR arrival at the TGN, suggesting a role of GGAs in endosome-to-TGN rather than anterograde transport. Conversely, acute inactivation of GGA2 by knocksideways did not affect CiMPR transport from the cell surface to the compartment of sulfation, suggesting possible indirect effects by RNAi.

Results

Functionalized nanobodies to analyze retrograde transport of CiMPR in cells depleted of candidate machineries

To study retrograde traffic to intracellular compartments including the TGN, we have previously established a versatile toolkit of functionalized anti–GFP nanobodies (Buser et al, 2018; Buser & Spiess, 2019). They can be used to label GFP-tagged proteins of interest at the cell surface and follow their route to endosomes, the TGN, and back to the plasma membrane. Here, we used anti–GFP nanobodies (VHH, variable heavy-chain domain of heavy-chain–only antibody) modified with a hexahistidine tag for purification, a T7 and an HA tag for immunodetection, a biotin acceptor peptide for biotinylation, and sequences conferring tyrosine sulfation (VHH-2xTS) or red fluorescence (VHH-mCherry) (Fig 1A). These functionalized nanobodies were bacterially expressed and isolated to
Functionalized nanobodies to analyze retrograde transport of CDMPR upon silencing of adaptor protein-1, retromer, Rab9a, or TIP47.

(A) Schematic representation of the functionalized nanobodies. The standard nanobody (VHH-std) consists of the GFP-specific VHH domain, T7 and HA epitope tags, a biotin acceptor peptide (BAP), and a hexahistidine (His6) purification tag. Other nanobodies in addition contain two tyrosine sulfation sequences (VHH-2xTS) or mCherry (VHH-mCherry). Scale bar in aa.

(B) Bacterially expressed and purified nanobodies (30 μg) were analyzed by SDS-gel electrophoresis and Coomassie staining (left). Immunoblot analysis of nanobodies (10 ng) with antibodies against the HA, His6, T7, or mCherry epitopes, or with streptavidin-HRP (SA-HRP). Marker proteins with molecular weights in kilodalton are shown on the left. As previously reported (Buser et al., 2018; Buser & Spiess, 2019), mCherry-containing nanobodies are slightly susceptible to clipping between the VHH and mCherry domains.

(C) HeLa cells were transfected with non-targeting siRNA or siRNAs targeting μ1A-adaptin, Vps26, TIP47, or Endosome-to-TGN transport routes of CDMPR Buser et al. https://doi.org/10.26508/lsa.202101269 vol 5 | no 7 | e202101269 3 of 19
high purity, and shown to be efficiently immunodetected using epitope tag antibodies or streptavidin-HRP (Fig 1B).

A well-established phenotype of retrograde transport deficiency on MPR traffic is the redistribution of the receptor from juxtanuclear to more peripheral compartments. To analyze the contribution of individual MPR retrieval routes to the TGN, we depleted machinery components of the AP-1/clathrin-, the retromer-, and the Rab9a/TIP47-dependent pathways by RNA interference using well-established siRNAs. Inactivation of the AP-1 complex was achieved by depleting the μ1A-subunit of the heterotetrameric adaptor complex (Hirst et al, 2003, 2005, 2009). The retromer complex was inactivated by silencing the subunit Vps26 of the cargo–selective complex (Popoff et al, 2007, 2009). Rab9a and TIP47, which do not form a stable complex, were knocked down separately (Ganley et al, 2004; Reddy et al, 2006; Bulankina et al, 2009; Kucera et al, 2016). All these proteins could be robustly depleted by >85% (Fig 1C and D), except TIP47 which was consistently reduced by ~65%, similar to the depletion efficiencies for TIP47 in the literature where they had been reported to produce clear effects on MPR traffic in vitro and in vivo (Diaz & Pfeffer, 1998; Ganley et al, 2004). As previously observed, depletion of μ1A or Vps26 caused a concomitant reduction of its complex partners γ-adaptin or Vps35, respectively (Fig 1C) (Meyer et al, 2000; Arighi et al, 2004), whereas the retromer-associated SNX-BAR proteins SNX1 and SNX2 were not affected upon depletion of Vps26 (Arighi et al, 2004; Rojas et al, 2007). Depletion of components of one pathway did not affect expression levels of proteins associated with other retrograde transport routes to the TGN (Fig 1C).

We silenced the above machinery components in HeLa cells stably expressing EGFP-CDMPR and analyzed its steady-state localization by fluorescence microscopy (Fig 1E) to test for mislocalization to endosomal compartments, a well-documented phenotype thought to result from defective endosome-to-TGN retrieval (Meyer et al, 2000; Arighi et al, 2004; Seaman, 2004; Popoff et al, 2009; Robinson et al, 2010; Hirst et al, 2012). In addition, we added VHH-mCherry nanobodies to the cells for 1 h before fixation to specifically detect the mature pool of the receptor cycling between surface, endosomes, and TGN (similar to previous antibody uptake experiments [Meyer et al, 2000; Robinson et al, 2010]). To measure the extent of mislocalization of EGFP-CDMPR to peripheral compartments, we used a semi-quantitative approach classifying the CDMPR staining patterns of individual cells as “mainly TGN,” “mainly peripheral,” and “fully peripheral” (Fig 1F) as previously applied by Cullen and colleagues (Wassmer et al, 2007; Simonetti et al, 2017). Knockdown of AP-1 or retromer caused a pronounced shift of both receptor and imported nanobody from the TGN to peripheral punctae as compared with control cells. We observed an approximately threefold increase in peripherally dispersed MPR-nanobody distributions in AP-1- and Vps26-depleted cells (Fig 1E and F), in agreement with previous analyses (e.g., Wassmer et al [2007]). In contrast, MPR localization was not significantly affected upon depletion of TIP47 or Rab9a.

Because CDMPR abundance, particularly at the plasma membrane, affects nanobody uptake and sulfation, we determined total and surface levels in silenced cells biochemically or by flow cytometry. Knocking down any of these four components did not cause any apparent change in plasma membrane levels of EGFP-CDMPR as assessed by nanobody (VHH-2xTS) binding only to the surface receptors at 4°C followed by immunoblotting (Fig S1). As a positive control, depletion of clathrin heavy chain (CHC17), which is required for clathrin-mediated endocytosis, produced the expected clear increase in surface EGFP-CDMPR (Fig S1). Interestingly, knockdown of the AP complex 2 (AP-2) did not mimic the phenotype of CHC17 depletion, in line with the observation that an AP-2 knockdown also did not cause cell surface accumulation of CIMPR (Dugast et al, 2005; Keyel et al, 2008; Tobys et al, 2021). To more quantitatively determine total and surface levels of EGFP-CDMPR in machinery-depleted cells, we performed flow cytometry to quantify the mean fluorescence intensity of GFP and of surface-bound VHH-mCherry, respectively (Fig 1G and H). RNAi-mediated depletion of Rab9a and Vps26 caused a significant increase of total EGFP-CDMPR levels by ~60%, probably by reducing lysosomal receptor degradation (Fig 1G). Surface receptor abundance was only augmented in Rab9a-silenced cells by ~40% (Fig 1H). Silencing of TIP47 or μ1A did not affect CDMPR levels.

To make sure that depletion of potential machinery components do not generally affect sulfation efficiency, on which our assay critically depends, HeLa cells stably expressing a sulfatable form of the secretory protein α1-protease inhibitor (A1Pi) were labeled with [35S]sulfate. No change in sulfation of A1Pi was observed for any of these protein knockdowns (Fig S2A and B).

Retrograde transport of CDMPR to the TGN is affected by depletion of Rab9a or Vps26, but not of TIP47

To more directly, more sensitively, and more quantitatively assay endosome-to-TGN transport in control and knockdown cells, we used VHH-2xTS nanobodies containing sites for tyrosine sulfation. This allows us to correlate appearance of nanobody sulfation with TGN arrival and residence time in this compartment. Cells stably expressing EGFP-CDMPR were silenced for one of the candidate retrograde machinery components or, as a control, transfected with Rab9a. 3 d after transfection, the cells were subjected to immunoblot analysis with antibodies against the indicated proteins. (D) To determine the knockdown (kd) efficiency, the residual protein was quantified in percent of the value after control-kd (mean and SD of three independent experiments). (E) HeLa cells stably expressing EGFP-CDMPR were depleted of μ1A-adaptin, Vps26, TIP47, or Rab9a as in (C). Cells were incubated for 1 h at 37°C with full medium containing 5 μg/ml VHH-mCherry (~0.1 μM), fixed, stained for EE1 and nuclei (DAPI, blue), and imaged by fluorescence microscopy. Bar: 10 μm. (F) Quantitation of the percentage of cells displaying the CDMPR localization phenotypes “mainly TGN,” “mainly peripheral,” or “fully peripheral” as in Wassmer et al (2007) and Simonetti et al (2017). For each condition, random frames with a total of 136–160 cells were scored from three independent experiments. (G, H) Total and surface EGFP-CDMPR levels in RNAi-silenced cells were quantified by flow cytometry. Cells were incubated for 30 min at 4°C with VHH-mCherry for exclusive binding to EGFP-CDMPR at the cell surface, washed, dissociated and analyzed for GFP and mCherry fluorescence to determine the levels of total (G) and surface EGFP-CDMPR (H), respectively. Median fluorescence intensities above background of parental HeLa cells without EGFP-CDMPR of each condition were normalized to the average of cells treated with non-targeting control siRNA. For each condition, 50,000 cells were analyzed in each experiment (mean and SD of three independent experiments).
non-targeting siRNA. The cells were then incubated with media containing VHH-2xTS for up to 75 min while labeling with $[^{35}\text{S}]$sulfate. In control cells, nanobody binding to EGFP-CDMPR and uptake reached its maximum within little more than 30 min and 50% after about 15 min (Fig 2A and B, open squares). Sulfation started only after a lag time of ~15 min and had not yet reached saturation after 75 min (Fig 2B, filled squares), in full agreement with our previous report (Buser et al, 2018). The difference between uptake and sulfation curves reflects the time of transport to the TGN.

The most thoroughly analyzed sorting machinery in endosome-to-TGN retrieval of MPRs is the canonical retromer complex (Arighi et al, 2004; Seaman, 2004; Wassmer et al, 2007; Burd & Cullen, 2014). Knocking down Vps26 as a core component of retromer caused a significant reduction of the rate of sulfation, indicating only ~50% of nanobody transport to the TGN after 75 min, whereas uptake was not considerably affected (Fig 2A and B). These results support a contribution of Vps26 in retrograde transport of CDMPR, confirming previous observations by different laboratories for CIMPR using immunofluorescence and antibody uptake assays (Arighi et al, 2004; Seaman, 2004, 2007; Popoff et al, 2007, 2009; McKenzie et al, 2012). Notably, we obtained a similar extent of transport impairment for EGFP-CDMPR as previously reported for STxB using single-time point sulfation experiments (Popoff et al, 2007, 2009).

Upon depletion of Rab9α, kinetics of nanobody uptake remained unaffected, whereas sulfation was modestly, but significantly reduced (Fig 2C and D). This effect is consistent with previous observations based on image analysis and antibody uptake for chimeric CIMPR and furin (Seaman et al, 2009; Chia et al, 2011). In these previous reports, transport to the TGN was reduced by up to 50% upon Rab9 knockdown, whereas we could only observe a reduction in signal of ~25% after 75 min, a difference that might be due to the method used. Taking the increased total and surface levels of EGFP-CDMPR in silenced cells into account (Fig 1H and G), the observed reduction of ~25% after 75 min might be an underestimate of the effect of Rab9α on retrograde CDMPR transport.

Whereas depletion of Rab9α or Vps26 both significantly affected transport of CDMPR from the plasma membrane to the compartment of sulfation, lysosomal hydrolase delivery—as measured by increased secretion of recombinant myc-tagged procathepsin D into the medium—was only affected by knockdown of Vps26, but not of Rab9α (Fig 2E and F). This is consistent with previous studies showing that CRISPR/Cas9–mediated deletion of Rab9α had no effect on cathepsin delivery to lysosomes, whereas lysosomal delivery of hydrolases was defective in retromer- or Vps26-deficient cells (Bugarcic et al, 2011; Fuse et al, 2015; Cui et al, 2019; Homma et al, 2019). As additional control, we analyzed secretion of transfected His6/myc-tagged PAUF (pancreatic adenocarcinoma up-regulated factor), a cargo of CARTS (carriers of the TGN to the cell surface) that bypass membranes related to Vps26 or Rab9α function (Wakana et al, 2012). Indeed, PAUF secretion was not altered by either knockdown (Fig 2G and H), confirming that the effect seen on procathepsin D is specific for Vps26.

Because TIP47 was proposed to mediate Rab9α-dependent MPR transport, knockdown should produce a similar effect as Rab9α depletion. However, kinetics of nanobody uptake and sulfation by EGFP-CDMPR remained unchanged in TIP47 knockdown cells (Fig 2I and J). Our sulfation experiments add an additional method to those used previously to evaluate retrograde transport kinetics of CDMPR with and without TIP47, again with a negative result.

**AP-1 contributes to both retrograde and anterograde transport of CDMPR**

To investigate the contribution of AP-1 in this process, we have previously analyzed the role of AP-1 in CD- and CIMPR transport to the TGN by rapid inactivation of AP-1 using knocksideways (Buser et al, 2018). Rapid depletion showed a reduction of approximately one third in the rate of sulfation, demonstrating a significant contribution of AP-1/clathrin in endosome-to-TGN transport of the MPRs. To test the outcome with AP-1 silencing in the same manner as applied to analyze the contribution of the other potential machineries above, we also performed the transport assay upon siRNA knockdown of μ1A.

Surprisingly, we did not observe a reduction of the kinetics and the extent of nanobody sulfation as expected from the more peripheral steady-state distribution of CDMPR in long-term AP-1–depleted cells. Instead, we found, after a similar lag phase as in all previous conditions, an ~2.5-fold increase in rate and extent of sulfation (Fig 3A and B). If this reflected transport directly, it would indicate increased retrograde transport activity by other mechanisms that even strongly overcompensated the loss of AP-1/clathrin–mediated transport. Yet, no increase in the levels of other machineries was detectable in μ1A knockdown cells (Figs 1C and S1) and no increase in general tyrosine sulfation (Fig S2A and B). The strong sulfation signal is also not the result of increased uptake of VHH-2xTS because the signals of cell-associated nanobody after loading at 37°C to steady-state or after binding only to cell-surface EGFP-CDMPR at 4°C were not increased in AP-1 knockdown compared with control cells (Figs 1H, 3A and B, and S1).

To rule out artefacts such as off-target effects of siRNA-mediated silencing of μ1A, we generated HeLa AP-1 knockout (ko) cells in which the γ1-adaptin genes were inactivated using CRISPR/Cas9. As expected, knockout cells displayed a complete loss of γ-adaptin staining in immunoblot and immunofluorescence analysis and a concomitant reduction of μ1A and σ1A subunits (Fig 3C and D). HeLa AP-1-γ1-ko cells recapitulated the phenotypes of μ1A knockdown cells. Whereas Golgi morphology remained unchanged, internalized anti-CIMPR antibody (as originally shown by Robinson et al [2010]) showed mainly peripheral localization that was largely rescued to a predominantly perinuclear Golgi localization upon re-expression of a γ-adaptin fusion protein (Fig 3E). Just like in the μ1A knockdown cells, sulfation of nanobody internalized by transfected EGFP-CDMPR was at least twofold higher after 75 min of labeling than in wild-type HeLa cells (Fig 3F and G).

However, we have previously observed that sulfation is not simply a detector of arrival in the compartment. Nanobody sulfation appears not to be sufficiently efficient to immediately and completely modify the sulfation sites as they enter the sulfation compartment. This explains why sulfation per nanobody depended on the target receptor: nanobodies taken up by EGFP-TGN46 showed considerably higher specific sulfation within 1 h than those captured by the EGFP-MPRs, even though maximal nanobody uptake was reached much later (Buser et al, 2018). This most likely reflects the residence time of these proteins in the sulfation...
compartment during which sulfate was continually incorporated into the nanobodies. A potential explanation of the observed hypersulfation is thus an increased residence time of the nanobody–EGFP-CDMPR complexes that still reached the TGN in the absence of AP-1. This is not unlikely because AP-1/clathrin is not only involved in retrograde retrieval to the TGN, but also in anterograde transport of MPRs out of the TGN to endosomes (Doray et al, 2002; Waguri et al, 2003; Ghosh et al, 2003a).

Rapid AP-1 inactivation by knocksideways inhibits TGN exit of CDMPR

The bidirectional function of AP-1/clathrin in MPR traffic thus makes it impossible to directly compare nanobody sulfation kinetics with other knockdown situations. To more directly demonstrate the effect of an anterograde transport block at the TGN on nanobody sulfation, we employed the AP-1 knocksideways cells (HeLa-AP1ks) established previously (Buser et al, 2018). AP-1 rerouting to mitochondria by rapamycin for 1 h shifted the steady-state distribution of EGFP-CDMPR to peripheral compartments as expected (Fig 4A and B). To demonstrate AP-1 dependence of TGN exit, we first loaded HeLa-AP1ks cells expressing EGFP-CDMPR with VHH-2xTS nanobody to steady-state during sulfate starvation, followed by [35S]sulfate labeling for up to 75 min. Upon addition of [35S]sulfate, there is a delay of 2–3 min for uptake and formation of 3′-phosphoadenosine-5′-phosphosulfate (PAPS) before incorporation of radioactivity gradually starts. To avoid these starting effects, rapamycin was added only 15 min after addition of...
Endosome-to-TGN transport routes of CDMPR  Buser et al.

Endosome-to-TGN transport routes of CDMPR  Buser et al.
transport of EGFP-CDMPR by ~50% after 75 min (Fig 5F and G), comparable with the effect of Vps26 depletion (Fig 2A and B). This result confirms a role of epsinR in endosome-to-TGN retrograde transport of CDMPR as previously suggested for CIMPR, TGN46, and STxB by Johannes and colleagues (Saint-Pol et al, 2004). Surprisingly, however, depletion of the GGAs revealed a very similar and significant reduction of VHH-2xTS sulfation kinetics by ~40% with no apparent effect on uptake (Fig 5H and I). This result is not consistent with the expected unique function of GGAs in anterograde transport, but rather supports a role in retrograde traffic.

Rapid GGA2 inactivation by knocksideways has no net effect on sulfation of CDMPR-bound nanobodies

If GGAs mediate anterograde transport of CDMPR out of the Golgi as indicated by previous studies, their silencing is expected to result in an increase in TGN residence time of CDMPR and imported nanobodies and thus in an increase in sulfation. Surprised by the opposite finding upon gradual depletion of GGAs, we also tested the effect of rapid GGA inactivation using the well-characterized GGA2 knocksideways system by Robinson and colleagues (Hirst et al, 2012). Because their GGA2 knocksideways cell line expresses YFP-modified Mitotrap to anchor-away FKBP-tagged GGA2 adaptors, we replaced GFP by mCherry in the CDMPR reporter and used functionalized anti-mCherry nanobodies (LaM4; [Fridy et al, 2014]), constructed in the same way as the VHHGFP4 constructs (Fig 6A). The derivatized anti-mCherry nanobodies were produced at the same high purity, yield, and functionality (Fig 6B), and were specifically endocytosed by mCherry-tagged reporter proteins (shown for mCherry-CDMPR and transferrin receptor [TfR]-mCherry in Fig S4).

We stably expressed mCherry-CDMPR in the HeLaM-GGA2ks cells (Fig 6C). Rapamycin-triggered mitochondrial rerouting of FKBP-GGA2 adaptors for 1 h did not cause detectable changes in the sulfation profile of CDMPR-bound nanobodies (Fig 6D).
Figure 5. Knockdown of the clathrin adaptors epsinR and GGA1–3 reduces retrograde transport of CDMPR.

(A) HeLa cells were transfected with non-targeting siRNA (Control-kd) or siRNAs targeting GGA1, GGA2, and GGA3, individually or combined (GGA1–3-kd), or epsinR. 3 d after transfection, cells were analyzed by immunoblotting with antibodies against the indicated proteins. (B) To determine the knockdown efficiency, the residual protein was quantified in percent of the value after control-kd (mean and SD of three independent experiments). (C) HeLa cells stably expressing EGFP-CDMPR were depleted of epsinR, or all three GGAs (GGA1–3) as in (B). Cells were incubated for 1 h at 37°C with full medium containing 5 μg/ml VHH-mCherry (~0.1 μM), fixed, stained for EEA1 and nuclei (DAPI, blue), and imaged by fluorescence microscopy. Bar: 10 μm. (D) Quantitation of the percentage of cells displaying the CDMPR localization phenotypes “mainly TGN,” “mainly peripheral,” or “fully peripheral” (Wassmer et al, 2007; Simonetti et al, 2017). For each condition, random frames with a total of 137–152 cells were scored from three independent experiments. (E) Normalized levels of total and surface EGFP-CDMPR levels in RNAi-silenced cells were quantified by flow cytometry as in Fig 1G and H. Mean fluorescence intensities of each condition were normalized to the average of cells treated with non-targeting control siRNA. For each knockdown condition,
distribution of mCherry-CDMPR (Fig 6F). To ensure functionality of the modified GGA2 knocksideways cell line, we assayed mis-sorting of myc-tagged procathepsin D precursor into the medium upon addition of rapamycin. Rerouting GGA2 to mitochondria resulted in a significant increase in procathepsin D secretion (Fig 6D and E).

To monitor the consequences of rapid GGA2 inactivation on retrograde CDMPR transport, cells were silenced for endogenous GGA1–3 by siRNA transfection, before they were incubated with LaM4-3xTS and simultaneously labeled with $^{35}$S sulfate in the presence or absence of rapamycin to rapidly remove the GGA2–FKBP fusion protein. Surprisingly, acute GGA2 depletion barely affected the sulfation kinetics of the nanobody imported by mCherry-CDMPR over 75 min (Fig 7A–C). This result is neither in agreement with GGAs specifically mediating retrograde, nor exclusively anterograde transport at the TGN–endosome interface. Rather, it could be the consequence of simultaneous reduction of transport in both directions.

To determine a potential direct role of GGA2 in TGN exit and thus anterograde transport, we applied the nanobody-preloading strategy (as already applied on AP-1ks cells in Fig 4C). We first loaded HeLa-GGA2ks cells expressing mCherry-CDMPR with LaM4-3xTS nanobody to steady-state during sulfate starvation, followed by $^{35}$S sulfate labeling for up to 75 min and where rapamycin was added 15 min post $^{35}$S sulfate addition (Fig 7D). Inactivation of GGA2 caused at best a mild, but not statistically significant increase in sulfation compared to cells treated with vehicle only (Fig 7E and F). Also this result would be consistent with compensatory effects on retrograde and anterograde transport of CDMPR of GGA2, notably in the absence of GGA1 and GGA3.

As proposed earlier for machinery depletion, the strategy of protein inactivation (gradual versus acute) turns out to be critical for phenotype analysis (Robinson et al, 2010; Hirst et al, 2012, 2015). Whereas knocksideways circumvent cellular adaptation and the up-regulation of potential compensatory mechanisms, long-term silencing by knockdown or knockout might be affected by accumulating indirect effects.

**Discussion**

In the present study, we have systematically analyzed the contribution of various intracellular sorting machineries to retrieval of CDMPR from the cell surface to the TGN by a transport assay. Most of the earlier studies were based on the analysis of changes in steady-state distributions of transported proteins by fluorescence microscopy upon silencing of potential machinery components. To more directly and quantitatively assess transport and its kinetics, we took advantage of functionalized anti-GFP nanobodies in combination with cell lines stably expressing EGFP-CDMPR (Buser et al, 2018; Buser & Spiess, 2019). Nanobodies containing TS sites report arrival and residence in the TGN, the compartment of sulfation, as they are internalized piggyback by EGFP-CDMPR from the cell surface. We previously applied this assay to test the contribution of AP-1 using the knocksideways system for rapid depletion (Buser et al, 2018). Because AP-1 is also implicated in anterograde transport from the TGN to endosomes, rapid inactivation promised less indirect effects resulting from inhibition of TGN exit than long-term silencing by siRNA-mediated knockdown or knockout. A clear reduction of the rate of sulfation by approximately one-third was detected after rapamycin-induced knocksideways, thus confirming a non-exclusive role of AP-1/clathrin in retrograde transport of CDMPR.

Here, we performed the experiment also with and without AP-1 inactivation by knockdown or knockout. The result was indeed strikingly different because hypersulfation was observed (Fig 3). Whereas other, AP-1-independent pathways still mediate significant retrograde transport of nanobody–EGFP-CDMPR complexes from endosomes to the TGN, their exit from the TGN is reduced by the absence of AP-1, extending their residence time in the sulfation compartment and thus the incorporation of $^{35}$S sulfate. We could show that pre-equilibrated nanobody–EGFP-CDMPR at the TGN was more strongly sulfated as soon as AP-1 depletion was triggered by rapamycin addition in knocksideways cells (Fig 4), clear evidence for inhibition of TGN exit and the anterograde role of AP-1-CCVs for CDMPR. Upon long-term depletion of AP-1, the steady-state pool of CDMPR at the TGN is likely higher than immediately after rapamycin-triggered knocksideways. This will further increase the residence time in the TGN and thus sulfation of entering nanobody–EGFP-CDMPR complexes and may account for the strong increase in sulfation after knockdown and knocksideways that overcompensates the reduction in incoming CDMPR.

It cannot be excluded that additional indirect effects caused by gradual and long-term depletion contribute to hypersulfation. Several unexpected and unexplained phenomena have previously been observed upon knockdown of AP-1, but not upon knocksideways: almost no reduction of CIMPR and ARF1 in CCVs, but increased AP-2 levels (Robinson et al, 2010; Navarro Negredo et al, 2017), and GGA2 was still incorporated into CCVs isolated from AP-1 knockdown but not from knocksideways cells (Hirst et al, 2012). However, tyrosine sulfation activity was not affected in AP-1 knockdown cells (Fig S2). Interestingly, in a proteomics search for CCV content dependent on AP-1, SLC35B2, one of the transporters delivering the activated sulfation precursor 3′-phosphoadenosine-5′-phosphosulfate (PAPS) into the TGN lumen scored positive (Hirst et al, 2012). One might thus speculate that lack of retrieval of this and other components of the sulfation machinery specifically by AP-1 might lead to hypersulfation in endosomes. Our observation, however, that localization of Golgi sulfotransferases remained largely unaffected makes this scenario appear unlikely, in particular...
because a concomitant redistribution of sulfotransferases and PAPS transporters is required for proper sulfate incorporation in proteins (Dick et al, 2008). Irrespective of that, one should mention that Johannes and colleagues observed a slight increase in STxB sulfation, when AP-1 was silenced by siRNA (Saint-Pol et al, 2004). Their findings with AP-1 have not been commented.

In any case, using our assay, it is expected that silencing of components involved in retrograde transport machineries for CDMPR causes reduced rates of nanobody sulfation, and depletion of proteins mediating TGN exit causes increased rates of sulfation. Accordingly, we found a clear reduction of nanobody sulfation upon knockdown of Vps26, confirming the role of retromer in retrograde transport.
transport also for CDMPR. There are many reports for a requirement of retromer for CIMPR (Arighi et al, 2004; Seaman, 2004, 2007, 2018; Wassmer et al, 2007; Bulankina et al, 2009; Seaman et al, 2009; Harbour et al, 2010; Bugarcic et al, 2011; Fjorback et al, 2012; Hao et al, 2013; McGough et al, 2014; Hirst et al, 2018; Wang et al, 2018; Chen et al, 2019; Cui et al, 2019). However, the requirement of the trimeric Vps retromer complex for CIMPR retrieval has recently been challenged by reports of the Cullen and Steinberg labs (Kvainickas et al, 2017; Simonetti et al, 2017, 2019; Evans et al, 2020). Using colocalization analyses, they failed to detect mislocalized CIMPR in the absence of rapamycin after 75 min (mean and SD of three independent experiments). Without rapamycin is shown as black squares, with rapamycin as gray circles; uptake as open symbols, sulfation as filled symbols. (E) HeLa-GGA2ks cells stably expressing mCherry-CDMPR were siRNA-silenced for endogenous GGA1–3, followed by starvation for sulfate in the presence of LaM4-3xTS to preload mCherry-CDMPR in the surface/endosome/TGN pool. The cells were then labeled with [35S] sulfate for up to 75 min in the continued presence of LaM4-3xTS, without or with addition of 500 nM rapamycin after 15 min (arrow) to inactivate GGA2 (+Rapa). (F) Nanobody analysis was performed as above (B). (F) Experiments as shown in panel E were quantified and presented as the percentage of the value in the absence of rapamycin after 75 min (mean and SD of three independent experiments). Without rapamycin is shown as black squares, with rapamycin as gray circles; uptake as open symbols, sulfation as filled symbols.

Figure 7. Effect of rapid GGA2 inactivation on retrograde and anterograde CDMPR transport using derivatized anti-mCherry nanobodies. (A, D) Schematic outline of the retrograde (A) anterograde (D) transport sulfation assays, respectively, with HeLa-GGA2ks/mCherry-CDMPR cells. (B) HeLa-GGA2ks cells stably expressing mCherry-CDMPR were siRNA-silenced for endogenous GGA1–3. The cells were starved for sulfate for 1 h, and then labeled for up to 75 min with [35S] sulfate in the presence of LaM4-3xTS nanobodies either in the absence (−) or presence of 500 nM rapamycin to inactivate GGA2-FKBP (+Rapa). The nanobodies were isolated by Ni/NTA beads and subjected to SDS-gel electrophoresis followed by immunoblot analysis (anti-His6) and autoradiography ([35S]). In parallel, aliquots of the cell lysates were immunoblotted for actin as a control for the amount of cells used. (C) Experiments as shown in panel B were quantified and presented as the percentage of the value of the absence of rapamycin after 75 min (mean and SD of three independent experiments). Without rapamycin is shown as black squares, with rapamycin as gray circles; uptake as open symbols, sulfation as filled symbols.

Endosome-to-TGN transport routes of CDMPR

Buser et al.

https://doi.org/10.26508/lsa.202101269 vol 5 | no 7 | e202101269

12 of 19
redistribution of CDMPR was observed and not perinuclear accumulation at the TGN. The results contradict an exclusive role of GGAs in anterograde transport of MPRs in cooperation with AP-1 as previously proposed (Doray et al, 2002, 2021; Ghosh et al, 2003a; Ghosh & Kornfeld, 2004). Contrary to the knockdown experiments silencing all three GGAs, rapid depletion of overexpressed GGA2 in the absence of GGA1 and GGA3 by knocksideways had no significant effect on the sulfation kinetics of CDMPR-imported nanobodies, neither hypersulfation supporting reduced TGN exit, nor hyposulfation consistent with reduced retrograde transport (Fig 7). This result suggests either that GGA2 contributes to neither anterograde nor retrograde transport of CDMPR, or that it is involved in both directions, whereby the opposing effects on nanobody sulfation could cancel each other out. In the latter case, one would have to assume that long-term depletion of the GGAs in the knockdown experiment produced changes/adaptations to specifically compensate the TGN exit defect. To more directly assess a role of GGA2 in anterograde transport, we imported nanobody into steady state before sulfation and acute inactivation. Rapid depletion of GGA2 caused a mild increase in sulfation consistent with a contribution to TGN exit and with the hypothesis of bidirectional functions. Direct comparison of knockdown experiments for all three GGAs with GGA2 knocksideways experiments in the absence of GGA1 and GGA3 is valid only under the assumption of complete redundancy between all three GGAs. This has not been analyzed in detail yet.

GGAs clearly localize both to the TGN and to endosomes (Dell’Angelica et al, 2000; Hirst et al, 2000; Ghosh et al, 2003b; Wahle et al, 2005; D’Souza et al, 2014; Ratcliffe et al, 2016; Uemura et al, 2018; Uemura & Waguri, 2020), like AP-1, and thus might operate at both places. Because GGA depletion had previously been observed to cause redistribution of CIMPR to EEA1-positive compartments, a role also in retrograde transport from endosomes had not been excluded (Ghosh et al, 2003b). Comparative CCV proteomics with GGA2 and AP-1 knocksideways cells pointed towards involvement of GGA7/AP-1 coats in anterograde sorting of MPR–lysosomal hydrolase complexes from the TGN (Hirst et al, 2012), yet the authors did not discount a potential retrograde function. A very surprising finding by Hirst et al (2012) was that Rabaptin5 (RABEP1) was the only known accessory component to be significantly lost from GGA2 knocksideways CCVs. Rabaptin5 is a marker of early endosomes where, as a complex with Rabex5 it activates Rab5 (Kalit et al, 2016). The fact that GGA depletion affects the CCV association of an endosomal protein points towards a role of these adaptors on endosomes, possibly in retrograde transport.

Interestingly, a recent study performed in Schizosaccharomyces pombe demonstrated that GGAs in collaboration with clathrin adaptors indeed contribute to efficient retrograde transport of Vps10, yeast’s MPR homologue, from the prevacuolar endosome to the TGN (Vangus et al, 2019). In addition, despite the functional relationship of GGA2 and AP-1 adaptors, it is surprising to observe that they are spatially segregated from each other to a considerable extent (Huang et al, 2019). Our results showing reduced CDMPR transport to the TGN using a sulfation-based approach strongly support a contribution to retrograde transport by GGAs.

In the present study, we have analyzed the contribution of individual sorting machineries in retrograde endosome-to-TGN transport of CDMPR and found that several machineries contribute, likely from different types of endosomes: retromer, the clathrin adaptors AP-1, epsinR, and—most surprisingly—the GGAs, and Rab9a. Other sorting machinery components that might facilitate endosome-to-TGN transport of MPRs include Rab29, Rab35, SNX-BAR proteins, and AP-5 (Wang et al, 2014; Cauvin et al, 2016; Kvainickas et al, 2017; Simonetti et al, 2017, 2019; Hirst et al, 2018). The two latter machineries have been shown to be important for CIMPR retrieval, not yet for CDMPR. Future systematic and comparative analyses of machinery requirement between CDMPR and CIMPR in retrograde transport for these machineries are important to further understand the coexistence and operation of multiple TGN retrieval pathways for receptors. Although CDMPR and CIMPR have completely different cytosolic tails and sorting motifs, they seem to largely share the same sorting machineries for their transport.

Importantly, our study also highlights the values and caveats of different depletion assays. RNAi silencing may be incomplete and may thus not reveal the full phenotypes. Protein depletion is gradual and thus allows compensatory mechanisms to be activated. Whereas in knockout cells, the protein of interest is completely absent, in both systems the long-term lack of a function may lead to indirect effects, such as the mislocalization of machinery components (e.g., SNAREs) involved in other transport steps. These disadvantages are avoided by rapid depletion by knocksideways, which aims to surprise the cell by the sudden absence of a protein. It requires, however, expression of a fusion protein that is fully functional and ideally expressed at a similar level as the original protein. Completeness of removal upon rapamycin addition is difficult to assess (particularly when highly overexpressed). In the case of the GGAs, the situation is complicated the existence of three isoforms (indicating isoforms?). Their complete redundancy is not established, since depletion of single GGAs already showed effects on cathepsin processing and sorting of (Ghosh et al, 2003a; Mardones et al, 2007). The available knocksideways cell line specifically addresses the effect of rapid depletion of overexpressed GGA2-FKBP in cells lacking GGA1 and GGA3.

The use of sulfatable nanobodies provides a new perspective on the sorting machineries of CDMPR. Our study highlights the critical differences of inactivation strategies (knockout, knockdown, knocksideways), but also provides evidence for a role of GGAs in retrograde transport of CDMPR, consistent with their localization to endosomes.

Materials and Methods

Bacterial expression and purification of functionalized nanobodies

Functionalized nanobodies were bacterially expressed and isolated as previously described (Buser et al, 2018; Buser & Spiess, 2019). Briefly, bacterial expression vectors encoding derivatized VHH or LaM4 nanobodies and myc-BirA (#109424; Addgene) were transformed together into Rosetta DE3 cells (Merck), and plated on LB plates with 50 μg/ml kanamycin and 50 μg/ml carbenicillin. A 20-ml overnight culture of a single colony was diluted into 1 liter LB medium with antibiotics and 200 μM D-biotin and grown to an OD600 of 0.6–0.7 at 37°C. Expression was induced with 1 mM isopropyl-β-D-
thiogalactopyranoside (IPTG) at 16°C overnight (VHH-mCherry and LaM4-EGFP), or at 30°C for 4 h (VHH-std, VHH-2xTS, LaM4-std, LaM4-1xTS, LaM4-2xTS, and LaM4-3xTS). Cells were pelleted at 5,000 g for 45 min and stored at -80°C. Upon thawing, they were resuspended in 30 ml PBS with 20 mM imidazole, 200 μg/ml lysozyme, 20 μg/ml DNase I, 1 mM MgCl₂, and 1 mM PMSF, incubated for 10 min at room temperature and 1 h at 4°C while rotating, followed by mechanical lysis using a tip sonicator for three times 30 s with 1-min cooling periods. The lysate was cleared by centrifugation at 15,000 g for 1 h at 4°C and loaded on a His GraviTrap column (GE Healthcare Life Sciences), washed with 20 mM imidazole in PBS, and eluted with 2 ml PBS with 500 mM imidazole. The purified nanobodies were desalted on PD-10 columns (GE Healthcare Life Sciences), concentrated to 2 mg/ml (VHH-std, VHH-2xTS, LaM4-std, LaM4-1xTS, LaM4-2xTS, and LaM4-3xTS) or 5 mg/ml (for VHH-mCherry and LaM4-EGFP), flash-frozen in liquid nitrogen, and stored at -80°C. Plasmids for nanobody fusion protein expression are deposited with Addgene (#109417, VHH-std; #109419, VHH-2xTS; #109421, VHH-mCherry; Addgene). TS- und EGFPtransfected LaM4 derivatives will be made available by Addgene (#162777–162779, #182641; Addgene).

Cell culture, plasmids, CRISPR/Cas9 gene editing, and RNA interference

HeLa α cell lines were maintained in high-glucose DMEM with 10% FCS, 100 units/ml streptomycin, 2 mM L-glutamine and appropriate selection antibiotics (1 mg/ml puromycin, 1 mg/ml hygromycin B, or 7.5 μg/ml blasticidin) at 37°C in 7.5% CO₂. The HeLa α cell line was cell line authenticated and proven to be human (Microsynth). Phoenix Amphi packaging cells (from the Nolan laboratories, Stanford University) were grown in complete medium supplemented with 1 mM sodium pyruvate.

HeLa cells stably expressing EGFP-CMTPR, and HeLa-AP1ks cells stably expressing the respective EGFP reporter were previously described (Buser et al., 2018). HeLaM-GGA2ks cells were a generous gift of Scottie Robinson and Jennifer Hirst.

To generate HeLa cells stably expressing SHMY-ΔA1Pi, SHMY-PAUF, cathepsin D-YM, TPST1-EGFP, and TPST2-EGFP, Phoenix Amphi packaging cells were transfected pQCXIP-SHMY-ΔA1Pi, pQCXIP-SHMY-PAUF, pQCXIP-cathepsin D-YM, pQCXIP-TPST1-EGFP, and pQCXIP-TPST2-EGFP using FuGENE HD (Promega). The viral supernatant was harvested after 48 h, passed through a 0.45 μm filter, supplemented with 15 μg/ml polybrene, and added to target HeLa α cells. The next day, complete medium with 1.5 μg/ml puromycin was added for selection, and pooled resistant clones were used for experiments. HeLa cells expressing TPST1-EGFP or TPST2-EGFP were further subjected to cell sorting on a FACSAria III (BD Biosciences) to obtain a cell pool with homogeneous expression levels. cDNA sequences of PAUF or cathepsin D were generous gifts by Vivek Malhotra (CRG) or Stuart Kornfeld (WUSM). Reporter plasmids were deposited at Addgene (#182642, EGFP-CMTPR; #182643, mCherry-CMTPR; #182644, cathepsin D-YM; #182645, SHMY-PAUF; #182648, SHMY-ΔA1Pi; Addgene).

A plasmid encoding TPST1-EGFP (#66617; Addgene) or TPST1-EGFP (#66618; Addgene) were kind gifts from David Stephens. The anti-mCherry nanobody sequence (LaM4) based on Rout and colleagues (Fridy et al, 2014) was a kind gift from Kazuhisa Nakayama (#70696; Addgene).

Generation of HeLaM-GGA2ks expressing mCherry-CMTPR (in pQCXIP) were established as outlined above, but selected and propagated with 1.5 μg/ml puromycin, 500 μg/ml hygromycin B, and 500 μg/ml G418. Homogeneous mCherry-CMTPR expression was obtained by cell sorting.

To generate a γ-adaptin knockout HeLa cell line by CRISPR/Cas9, sgRNAs for gene editing were purchased from Santa Cruz Biotechnology (sc-403986). Briefly, parental HeLa cells were transfected with 2 μg plasmid containing a GFP cassette and 4 μl FuGENE HD (Promega) in a six-well cluster. After 24 h of expression, cells were subjected to FACS, and single cells or pooled clones were collected. Transient transfection of AP-1ko cells with pQCXIP-γ-FKB, pQCXIP-TPST1-EGFP, or pQCXIP-TPST2-EGFP was performed using FuGENE HD according to the manufacturer’s instructions. Anti-CMTPR antibody uptake was performed as described previously (Robinson et al., 2010).

For RNA interference, cells were reverse-transfected with target siRNA in Opti-MEM I using Lipofectamine RNAIMAX (both Thermo Fisher Scientific) following the manufacturer’s instructions. For a conventional knockdown of AP-1, the sequence 5’-AAGGCAUCAAGAUUCGGAAGATdTdT3’ against the μ1A-subunit of the heterotetrameric complex was used as formerly reported (Hirst et al, 2003, 2005, 2009). For RNA interference with retromer complex, we applied siRNA duplexes with the sequence 5’-AACUCUCUGAAACCUUCUAGGdtdTdT3’ targeting Vps26 as described in previous studies (Popoff et al, 2007, 2009). To specifically silence Rab9 or TIP47, we applied the siRNA sequence 5’-GUUUAAGACCGCUCUCdTdT3’ for Rab9 (Garley et al, 2004; Reddy et al, 2006; Kucera et al, 2016), or 5’-CCCGGGGCUCAUUUCACdTdTdT3’ for TIP47 (Bulinkina et al, 2009). EpsinR was targeted with 5’-AAUACAGAUAUGCCCACAATTdTdT3’, GGA1 with 5’-CACAGAGUGGGGAGGCGGAdTdT3’, GGA2 with 5’-UGAUUAUGGUUUCCGCAATdTdT3’, and GGA3 with 5’-UGUGACGGCCUCAUGAACATTdTdT3’ as previously described (Hirst et al, 2004, 2012). To knockdown AP-2δ and CHC17, we used 5’-AA GACGACUGACGCGCUCCUCATdTdT3’ and 5’-UAAUCAUAGUAGGGGGCAdTdT3’, respectively, as previously described (Motley et al, 2003). We used the non-targeting siRNA 5’-UA GGCACUAGAUGAGGAACdTdT3’ as control siRNA (Salazar et al, 2009). All siRNAs were used at a final concentration of 50 nM, apart from the GGA siRNAs (used at 25 nM each).

To silence γ-adaptin in HeLa-AP1ks cells, the siRNA sequence 5’-GAAGUAUAAUCCUUUU3’ was used as previously described (Robinson et al, 2010; Buser et al, 2018). Cells were transfected twice (day 1 and 3) and used at day 5. siRNA duplexes were purchased from Microsynth.

Gen electrophoresis and immunoblot analysis

Proteins separated by SDS-gel electrophoresis (7.5–15% polyacrylamide) were transferred to Immobilon-P® PVDF membranes (Millipore). After blocking with 5% non-fat dry milk in TBS (50 mM Tris-HCl, pH 7.6, 150 mM NaCl) with 0.1% Tween-20 (TBST) for 1 h, the membranes were probed with primary antibodies in 1% BSA in TBST for 2 h at room temperature or overnight at 4°C, followed by incubation with HRP-coupled secondary antibodies in 1% BSA in TBST.
for 1 h at room temperature. Immobilon Western Chemiluminescent HRP Substrate (Millipore) was used for detection, a Fusion Vilber Lourmat Imaging System for imaging, and Fiji software for quantification.

**Secretion assay**

To monitor effects of machinery depletion on CDMPR-mediated lysosomal sorting, HeLa cells stably expressing cathepsin D-YMH or SHMY-PAUF were reverse-transfected with target siRNA as outlined above. After RNAi, cells were washed and incubated in serum-free medium for 1 h in the presence of 5 mM mannose-6-phosphate to avoid subsequent endocytosis of the secreted cathepsin precursor. Precipitation of recombinant His6-tagged cathepsin precursors from the medium was performed with Ni Sepharose High Performance beads (GE Healthcare Life Sciences) for 1 h at 4°C. Beads were washed three times with lysis buffer containing 20 mM imidazole and boiled in SDS-sample buffer. In addition, a fraction (50–100 μl) of the cell lysate was used for an actin control. Samples were run on an SDS–PAGE and analyzed by immunoblotting.

To assay recombinant cathepsin secretion in RNAi-silenced HeLaM-GGA2ks cells, cells were transiently transfected with pQCXIP-cathepsin D-YMH. After RNAi, cells were harvested with non-enzymatic cell dissociation solution (Millipore), suspended in 1 ml lysis buffer containing 2 mM PMSF and protease inhibitor cocktail, and centrifuged at 10,000 g for 10 min. The supernatant was collected and incubated for 1 h at 37°C with 1% Igepal CA-630 and 7.5% CO2 in the presence of 500 nM rapamycin. Secretion was biochemically assessed as outlined above.

**Fluorescence microscopy**

For immunofluorescence staining, cells were grown on glass coverslips, fixed with 3% PFA for 10 min at room temperature, washed with PBS, quenched with 50 mM NH4Cl in PBS for 5 min, permeabilized with 0.1% Triton X-100 in PBS for 10 min, blocked with 1% BSA in PBS for at least 15 min, incubated with primary antibody in BSA/PBS for 2 h, washed, and stained with fluorescent secondary antibodies in BSA/PBS for 1 h. After a 5 min staining with 5 μg/ml DAPI and three washes with PBS, coverslips were mounted in Fluoromount-G (Southern Biotech). Staining patterns were imaged on a Zeiss Dmi8 Confocal LSM700 or LSM880 microscope in super-resolution mode with the Airyscan detector. Receptor mislocalization analysis was performed by scoring and quantification of the percentage of cells displaying each phenotype in machinery-depleted cells using a Zeiss Axioplan microscope with a Leica DFC420C imaging system as described previously (Wassmer et al., 2007; Simonetti et al., 2017).

**Flow cytometry**

To quantitatively assess changes in total and surface abundance of recombinant CDMPR upon silencing of machinery components, parental HeLa cells as well as HeLa cells stably expressing EGFP-CDMPR, untreated or after treatment with non-targeting control siRNA or various targeting siRNAs, were incubated with 5 μg/ml purified VHH-mCherry in complete medium for 30 min at 4°C to specifically bind surface EGFP-CDMPR. After six washes with ice-cold medium to remove unbound nanobody, cells were gently harvested with non-enzymatic cell dissociation solution (Millipore). Collected cells were resuspended in ice-cold PBS with 0.5% FBS for flow cytometry analysis using a LSR Fortessa Analyzer (BD Biosciences) with a blue (488 nm) and yellow-green (561 nm) laser. Surface CDMPR (mCherry) and total receptor (GFP) levels above the background of parental HeLa cells were quantified, as median fluorescence intensities using FCS Express Research 7 (De Novo Software) and FlowJo 10 (BD), and normalized to the mean value of control knockdown cells.

**Sulfation analysis**

To analyze retrograde transport and kinetics of EGFP-CDMPR to the compartment of sulfation, cells were incubated with 1 ml sulfate-free medium for 1 h at 37°C and 7.5% CO2 before labeling with sulfate-free medium supplemented with 0.5 mCi/ml [35S]sulfate (Hartmann Analytics) and 2 μg/ml purified VHH-2xTS for up to 75 min. For the knocksideways experiment, HeLa-A1Pi cells stably expressing EGFP-CDMPR and mCherry-CDMPR, respectively, were starved with sulfate-free medium for 1 h at 37°C and 7.5% CO2 in the presence of 2 μg/ml purified VHH-2xTS or LaM4-3xTS, followed by two 1 ml washes with sulfate-free medium, before labeling with medium reconstituted with 0.5 mCi/ml [35S]sulfate. 500 nM rapamycin from a 2,000× stock solution in DMSO, or DMSO alone was added. Cells stably expressing SHMY-A1Pi were starved as described above and pulsed with medium reconstituted with 0.5 mCi/ml [35S]sulfate for 75 min. After incubation, cells were washed twice with ice-cold PBS, lysed in 1 ml lysis buffer containing 2 mM PMSF and protease inhibitor cocktail, and centrifuged at 10,000g for 15 min at 4°C. A fraction (50–100 μl) of the postnuclear supernatants was used for immunoblot analysis of total cell-associated nanobody and an actin control. The rest was incubated for 1 h at 4°C with 20 μl Ni Sepharose High Performance beads (GE Healthcare Life Sciences) to isolate the nanobodies. Beads were washed three times with lysis buffer containing 20 mM imidazole and boiled in SDS-sample buffer. Nanobodies were analyzed by SDS-gel electrophoresis and autoradiography using BAS Storage Phosphor Screens and a Typhoon FLA7000 IP phosphorimager (GE Healthcare Life Sciences).

**Antibodies**

For immunofluorescence microscopy, goat anti-α-adaptin (#EB11875; 1:1,000; Everest Biotech), mouse anti-CIMPrab (ab2733; 1:5,000; Abcam), anti-EEA1 (#R60456; 1:1,000; BD Biosciences), rabbit anti-FLAG (#2368; 1:500; Cell Signaling Technology), and rabbit anti-CHC17 (made from TD.1 hybridoma; 1:200) antibodies were used.

For immunoblotting, mouse anti-α-adaptin (#610505; 1:5,000; BD Biosciences), mouse anti-γ-adaptin (#610385; 1:5,000; BD Biosciences), mouse anti-β1/2-adaptin (#610381; 1:5,000; BD Biosciences), mouse anti-γ-adaptin (made from 100/3 hybridoma; 1:5,000), rabbit anti-ζ-adaptin (#A305-396A-M; 1:1,000; Bethyl Laboratories), mouse anti-γ-adaptin (made from 100/3 hybridoma; 1:5,000), mouse anti-actin (#MAB1501; 1:100,000; EMD Millipore), mouse anti-CHC17 (made from TD.1 hybridoma; 1:200), rabbit anti-epsinR (#A301-926A; 1:1,000; Bethyl Laboratories), mouse anti-FLAG (#8146; 1:1,000; Cell Signaling Technology), mouse anti-GFP (#11814460001-Roche;
As secondary antibodies for immunofluorescence microscopy, A568-labeled donkey anti-goat (αA-11057; 1:500; Thermo Fisher Scientific), A647-labeled donkey anti-goat (αA-21447; 1:500; Thermo Fisher Scientific), A633-labeled goat anti-mouse (αA-21052; 1:500; Thermo Fisher Scientific), A633-labeled goat anti-rabbit (αA-21071; 1:500; Thermo Fisher Scientific) immunoglobulin antibodies were used. As secondary antibodies for immunoblotting, HRP-labeled goat anti-rabbit (αA-0168; 1:10,000; Sigma-Aldrich) and goat anti-mouse (αA-0168; 1:10,000; Sigma-Aldrich) immunoglobulin antibodies were used. To detect biotinylated proteins on blots, Streptavidin-HRP (αA4323; 1:10,000; Thermo Fisher Scientific) was used.

Supplementary Information

Supplementary Information is available at https://doi.org/10.26508/lsa.202010269.

Acknowledgements

This work was supported by grant 31003A-182519 from the Swiss National Science Foundation. We particularly thank Nicole Beuret, Janine Bögli from the Biozentrum Imaging Core Facility (IMCF), for their support, and Scottie Robinson and Jennifer Hirst for the HeLaM-GGA2ks cells and advice on the knocksideways system.

Author Contributions

DP Buser: conceptualization, resources, data curation, formal analysis, supervision, validation, investigation, methodology, project administration, and writing—original draft, review, and editing.

G Bader: formal analysis and investigation.

M Spiess: conceptualization, resources, data curation, formal analysis, supervision, funding acquisition, validation, visualization, methodology, project administration, and writing—original draft, review, and editing.

Conflict of Interest Statement

The authors declare that they have no conflict of interest.

References

Arighi CN, Hartnell LM, Aguilar RC, Haft CR, Bonifacino JS (2004) Role of the mammalian retromer in sorting of the cation-independent mannose 6-phosphate receptor. J Cell Biol 165: 123–133. doi:10.1083/jcb.200312055

Banik SM, Pedram K, Wisnovsky S, Ahn G, Riley NM, Bertozzi CR (2020) Lysosome-targeting chimeraeas for degradation of extracellular proteins. Nature 584: 291–297. doi:10.1038/s41586-020-2545-9

Belenkaya TY, Wu Y, Tang X, Zhou B, Cheng L, Sharma YV, Yan D, Selva EM, Lin X (2008) The retromer complex influences Wnt secretion by recycling wntless from endosomes to the trans-Golgi network. Dev Cell 14: 120–131. doi:10.1016/j.devcel.2007.12.003

Boman AL, Zhang Cj, Zhu X, Kahn RA (2000) A family of ADP-ribosylation factor effectors that can alter membrane transport through the trans-Golgi. Mol Biol Cell 11: 1241–1255. doi:10.1091/mbc.11.4.1241

Bonifacino JS, Hurley JH (2008) Retromer. Curr Opin Cell Biol 20: 427–436. doi:10.1016/j.jocb.2008.03.009

Bonifacino JS, Rojas R (2006) Retrograde transport from endosomes to the trans-Golgi network. Nat Rev Mol Cell Biol 7: 568–579. doi:10.1038/nrm1985

Breussegem SY, Seaman MNJ (2014) Image-based and biochemical assays to investigate endosomal protein sorting. Methods Enzymol. 534: 155–178. doi:10.1016/B978-0-12-397926-1.00009-3

Bugarcic A, Zhe Y, Kerr MC, Griffin J, Collins BN, Teasdale RD (2011) Vps26a and Vps26b subunits define distinct retromer complexes. Traffic 12: 1759–1773. doi:10.1111/j.1600-0854.2011.01284.x

Bulankina AV, Deggerich A, Wenzel D, Mutenda K, Wittmann JG, Rudolph MG, Burger KN, Höning S (2009) TIP47 functions in the biogenesis of lipid droplets. J Cell Biol 185: 641–655. doi:10.1083/jcb.200812042

Burd C, Cullen PJ (2014) Retromer: A master conductor of endosome sorting. Cold Spring Harb Perspect Biol 6: a016774. doi:10.1101/cshperspect.a016774

Buser DP, Schleicher KD, Prescianotto-Baschong C, Spiess M (2018) A versatile nanobody-based toolkit to analyze retrograde transport from the cell surface. Proc Natl Acad Sci U S A 115: E6227–E6236. doi:10.1073/pnas.1801865115

Buser DP, Spiess M (2019) Analysis of endocytic uptake and retrograde transport to the trans-Golgi network using functionalized nanobodies in cultured cells. J Vis Exp. doi:10.3791/26511

Carlton J, Bujny M, Peter Bj, Oorschot VM, Rutherford A, Mellor H, Klumperman J, McMahon HT, Cullen PJ (2011) Vps26a and Vps26b subunits define distinct retromer complexes. Curr Opin Cell Biol 23: 1791–1800. doi:10.1016/j.cub.2011.10.077

Carlton JG, Bujny NV, Peter Bj, Oorschot VM, Rutherford A, Arkell RS, Klumperman J, McMahon HT, Cullen PJ (2005) Sorting nexin-1 mediates tubular endosome-to-TGN transport through coincidence sensing of high-curve membrane and 3-phosphoinositides. Curr Biol 14: 1800–1806. doi:10.1016/j.cub.2004.09.077

Carlton JG, Bujny M, Peter BJ, Oorschot VM, Rutherford A, Arkell RS, Klumperman J, McMahon HT, Cullen PJ (2005) Sorting nexin-2 is associated with tubular elements of the early endosome, but is not essential for retromer-mediated endosome-to-TGN transport. J Cell Sci 118: 4527–4539. doi:10.1242/jcs.02568

Cauvin C, Rossendale M, Gupta-Rossi N, Rocancourt M, Larrau A, Perrais D, Echard A (2016) Rab35 GTPase triggers switch-like recruitment of the Lowe syndrome lipid phosphatase OCRL on endosomal traficking by Retromer and Retriever. J Cell Sci 129: 1383–1394. doi:10.1242/jcs.175903

Chia PZ, Gnesanaru I, Lieu ZZ, Gleeson PA (2011) Rab9-dependent retrograde transport and endosomal sorting of the endoprotease furin. J Cell Sci 124: 2401–2413. doi:10.1242/jcs.083782

Endosome-to-TGN transport routes of CDMPR Buser et al. https://doi.org/10.26508/lsa.202101269 vol 5 no 7 e202101269
Cozier GE, Carlton J, McGregor AH, Gleeson PA, Teasdale RD, Mellor H, Cullen PJ (2002) The phox homology (PX) domain-dependent, 3-phosphoinositide-mediated association of sorting nexin-1 with an early sorting endosomal compartment is required for its ability to regulate epidermal growth factor receptor degradation. J Biol Chem 277: 48730–48736. doi:10.1074/jbc.M206986200

Cui Y, Carosi JM, Yang Z, Ariotti N, Kerr MC, Parton RG, Sargeant TJ, Teasdale RD (2019) Retromer has a selective function in cargo sorting via endosome transport carriers. J Cell Biol 218: 615–631. doi:10.1083/jcb.201806153

D’Souza RS, Semus R, Billings EA, Meyer CB, Conger K, Casanova JE (2014) Rab4 orchestrates a small GTPase cascade for recruitment of adaptor proteins to early endosomes. Curr Biol 24: 1187–1198. doi:10.1016/j.cub.2014.04.003

Dell’Angelica EC, Puertollano R, Mullins C, Aguilar RC, Vargas JD, Hartnell LM, Bonifacino JS (2000) GGAs: A family of ADP ribosylation factor-binding proteins related to adaptors and associated with the Golgi complex. J Cell Biol 149: 81–94. doi:10.1083/jcb.149.1.81

Díaz E, Pfeffer SR (1998) TIP47: A cargo selection device for mannose 6-phosphate receptor trafficking. Cell 93: 433–443. doi:10.1016/s0092-8674(00)81171-x

Dick G, Gründahl F, Prydz K (2008) Overexpression of the 3-5-phosphoadenosine 5’-phosphosulfate (PAPS) transporter 1 increases sulfation of chondroitin sulfate in the apical pathway of MDCK II cells. Glycobiology 18: 53–65. doi:10.1093/glycob/cwm121

Doray B, Ghosh P, Griffith J, Geuze HJ, Kornfeld S (2002) Cooperation of GGAs and AP-1 in packaging MPRTs at the trans-Golgi network. Science 297: 1700–1703. doi:10.1126/science.1075327

Doray B, Liu L, Lee WS, Jennings BC, Kornfeld S (2021) Inactivation of the three GGA genes in HeLa cells partially compromises lysosomal enzyme sorting. FEBS Open Bio 11: 367–374. doi:10.1111/1753-8585.14304

Dugast M, Toussaint H, Dousset C, Benaroch P (2005) AP2 clathrin adaptor proteins with gamma-adaptin and VHS domains that facilitate trafficking between the trans-Golgi network and the vacuole/lysosome. J Cell Biol 169: 67–80. doi:10.1083/jcb.149.1.67

Hirst J, Miller SE, Taylor MJ, von Mollard GF, Robinson MS (2004) EpsinR is an adaptor for the SNARE protein Vti1b. Mol Cell 15: 5593–5602. doi:10.1016/j.molcel.2004.06.046

Hirst J, Motley A, Harasaki K, Peak Chee SY, Robinson MS (2003) EpsinR: An ENTH domain-containing protein that interacts with AP-1. Mol Biol Cell 14: 625–641. doi:10.1091/mbc.e02-09-0552

Hirst J, Sahlender DA, Robinson MS (2020) A family of proteins with gamma-adaptin and VHS domains that facilitate trafficking between the trans-Golgi network and the vacuole/lysosome. Proc Natl Acad Sci USA 117: 1711–1716. doi:10.1073/pnas.201207.012

Hirst J, Borner GH, Harbour M, Robinson MS (2005) The adapter protein complex in late endosome-to-Golgi retrieval. PLoS Biol 16: e200411. doi:10.1371/journal.pbio.200411

Hirst J, Lux WW, Bright NA, Totty N, Seaman MN, Robinson MS (2000) A family of proteins with gamma-adaptin and VHS domains that facilitate sorting and processing. J Biol Chem 275: 1868–1873. doi:10.1074/jbc.M202570200

Hirst J, Lui WW, Bright NA, Totty N, Seaman MN, Robinson MS (2000) A family of proteins with gamma-adaptin and VHS domains that facilitate sorting and processing. J Cell Sci 133: jcs246033. doi:10.1242/jcs.246033

Hirst J, Miller SE, Taylor MJ, von Mollard GF, Robinson MS (2004) EpsinR is an adaptor for the SNARE protein Vti1b. Mol Cell 15: 5593–5602. doi:10.1016/j.molcel.2004.06.046

Hirst J, Motley A, Harasaki K, Peak Chee SY, Robinson MS (2003) EpsinR: An ENTH domain-containing protein that interacts with AP-1. Mol Biol Cell 14: 625–641. doi:10.1091/mbc.e02-09-0552

Hirst J, Sahlender DA, Choma M, Sinka R, Harbour M, Parkinson M, Robinson MS (2009) Spatial and functional relationship of GGAs and AP-1 in Drosophila and HeLa cells. Traffic 10: 1696–1710. doi:10.1111/j.1600-089x.2009.00983.x

Homma Y, Kinosita R, Kuchitsu Y, Wakao PS, Wawro PS, Marubashi S, Oguchi ME, Ishida M, Fujita N, Fukuda M (2019) Comprehensive knockout analysis of the Rab family GTPases in epithelial cells. J Cell Biol 218: 2035–2050. doi:10.1083/jcb.201810134

Huang Y, Ma T, Lau PK, Wang J, Zhao T, Du S, Loy MMT, Guo Y (2019) Visualization of protein sorting at the trans-golgi network and endosomes through super-resolution imaging. Front Cell Dev Biol 7: 181. doi:10.3389/fcell.2019.00181

Itzhak DN, Tyanova S, Cox J, Borner GH (2016) Global, quantitative and dynamic mapping of protein subcellular localization. Nat Methods 13: 914–915. doi:10.1038/nmeth.4781

Johannes L, Popoff V (2008) Tracing the retrograde route in protein trafficking. Cell 135: 1175–1187. doi:10.1016/j.cell.2008.12.009

Kalín S, Buser DP, Spiess M (2016) A fresh look at the function of Rabaptin5 on endosomes. Small GTPases 7: 34–37. doi:10.1080/21541248.2016.1140616

Keyel PA, Thieman JR, Roth R, Erkan E, Everett ET, Watkins SC, Heuser JE, Traub LM (2008) The AP-2 adaptor beta2 appendage scaffolds alternate...
cargo endocytosis. *Mol Biol Cell* 19: 5309–5326. doi:10.1091/mbc.e08-07-0712

Kucera A, Borg Distefano M, Berg-Larsen A, Skjeldal F, Repnik U, Bakke O, Progida C (2016) Spatiotemporal resolution of Rab9 and Cl-MPR dynamics in the endocytic pathway. *Traffic* 17: 211–229. doi:10.1111/tra.12357

Kvainickas A, Jimenez-Orgaz A, Nagele H, Hu Z, Dengjel J, Steinberg F (2017) Cargo-selective SNX-BAR proteins mediate retromer trimer independent endosome-to-TGN transport. *J Cell Biol* 216: 3677–3693. doi:10.1083/jcb.201702137

Le Borgne R, Griffiths G, Hoflack B (1996) Mannose 6-phosphate receptors and ADP-ribosylation factors cooperate for high affinity interaction of the AP-1 Golgi assembly proteins with membranes. *J Biol Chem* 271: 2162–2170. doi:10.1074/jbc.2714.2162

Lombardi D, Soldati T, Riederer MA, Goda Y, Zerial M, Pfeffer SR (1993) Rab9 functions in transport between late endosomes and the trans Golgi network. *EMBO J* 12: 677–682. doi:10.1002/j.1460-2075.1993.tb05701.x

Lucas M, Gershlick DC, Vidaurrezaga A, Rojas AL, Bonifacino JS, Hierro A (2016) Structural mechanism for cargo recognition by the retromer complex. *Cell* 167: 1623–1635.e14. doi:10.1016/j.cell.2016.10.056

Mardones GA, Burgos PV, Brooks DA, Parkinson-Lawrence E, Mattera R, Bonifacino JS (2007) The trans-Golgi network accessory protein p56 promotes long-range movement of GGA/clathrin-containing transport carriers and lysosomal enzyme sorting. *Mol Biol Cell* 18: 3468–3501. doi:10.1091/mbc.e07-02-0190

McGough IJ, Steinberg F, Jia D, Barbuti PA, McMillian KJ, Heesom KJ, Whone AL, Caldwell MA, Billadeau DD, Rosen MK, et al (2014) Retromer binding to FAM21 and the WASH complex is perturbed by the Parkinson disease-linked VPS35(D20N) mutation. *Curr Biol* 24: 1678. doi:10.1016/j.cub.2014.07.004

Mckenzie JE, Raisely B, Zhou X, Naslavsky N, Taguchi T, Caplan S, Sheff D (2012) Retromer guides STxB and CD8-M6PR from early to recycling endosomes, EHD1 guides STxB from recycling endosome to Golgi. *J Cell Biol* 198: 1159–1170. doi:10.1083/jcb.201211005

Metherel GR, Schu P (2003) Characterization of the in vitro retrograde transport of MPRA6. *Traffic* 4: 802–811. doi:10.1111/j.1600-0854.2003.tb01366.x

Meyer C, Zizioli D, Lausmann S, Eskelinen EL, Hamann J, Saftig P, von Figura K, Ratcliffe CD, Sahgal P, Parachoniak CA, Ivaska J, Park M (2016) Regulation of cell migration and β1 integrin trafficking by the endosomal adapter GGA3. *Traffic* 17: 670–688. doi:10.1111/tra.12390

Reddy IV, Burguete AS, Sridive K, Ganley IG, Nottingham RM, Pfeffer SR (2006) A functional role for the GCC185 golgin in mannose 6-phosphate receptor recycling. *Mol Biol Cell* 17: 4333–4343. doi:10.1091/mbc.e06-02-0153

Robinson MS, Sahlender DA, Foster SD (2010) Rapid inactivation of proteins by rapamycin-induced rerouting to mitochondria. *Dev Cell* 18: 324–331. doi:10.1016/j.devcel.2009.12.015

Rojas R, Kamekata T, Haft CR, Bonifacino JS (2007) Interchangeable but essential functions of SNX1 and SNX2 in the association of retromer with endosomes and the trafficking of mannose 6-phosphate receptors. *Mol Cell Biol* 27: 1112–1124. doi:10.1128/MCB.00156-06

Rojas R, van Vlijmen T, Mardones GA, Prabhu Y, Rojas AL, Mohammed S, Heck AJ, Raposo G, van der Sluijs P, Bonifacino JS (2008) Regulation of retromer recruitment to endosomes by sequential action of Rab6 and Rab7. *J Cell Biol* 183: 513–526. doi:10.1083/jcb.200804048

Saint-Pol A, Velamos B, Amessou M, Mills IG, Dugast M, Tenza D, Schu P, Antony C, McMahan HT, Lamaze C, et al (2004) Clathrin adaptor epsinR is required for retrograde sorting on early endosomal membranes. *Dev Cell* 6: 525–538. doi:10.1016/s1534-5807(04)00100-5

Salazar G, Zlatic S, Craigie B, Peden AA, Pohl J, Faundez V (2009) Hermansky-Pudlak syndrome protein complexes associate with phosphatidylinositol 4-kinase type II alpha in neuronal and non-neuronal cells. *J Biol Chem* 284: 1790–1802. doi:10.1074/jbc.M805991200

Sandvig K, van Deurs B (2002) Transport of protein toxins into cells: Pathways used by ricin, cholera toxin and Shiga toxin. *FEBS Lett* 529: 49–53. doi:10.1016/s0014-5793(02)01318-4

Sanger A, Hirst J, Davies AK, Robinson MS (2019) Adaptor protein complexes and disease at a glance. *J Cell Sci* 132: jcs229992. doi:10.1242/jcs.229992

Seaman MN (2004a) Cargo-selective endosomal sorting for retrieval to the Golgi requires retromer. *J Cell Biol* 165: 111–122. doi:10.1083/jcb.200312034

Seaman MN, Harbour ME, Tattersall D, Read E, Bright N (2009) Membrane recruitment of the cargo-selective retromer subcomplex is catalysed by the small GTPase Rab7 and inhibited by the Rab-GAP TBC1D5. *J Cell Sci* 122: 2371–2382. doi:10.1242/jcs.048686

Seaman MN (2007) Identification of a novel conserved sorting motif required for retromer-mediated endosome-to-TGN retrieval. *J Cell Sci* 120: 2378–2389. doi:10.1242/jcs.009654

Seaman MN, Sowerby PJ, Robinson MS (1996) Cytoplasmic and membrane-associated proteins involved in the recruitment of AP-1 adaptors onto the trans-Golgi network. *J Biol Chem* 271: 25446–25451. doi:10.1074/jbc.2541.254546

Seaman MN (2012) The retromer complex - endosomal protein recycling and beyond. *J Cell Sci* 125: 4693–4702. doi:10.1242/jcs.133440

Seaman MN (2018) Retromer and the cation-independent mannose 6-phosphate receptor: Time for a trial separation? *Traffic* 19: 150–152. doi:10.1111/tra.12542

Simonetti B, Danson CM, Heesom KJ, Cullen PJ (2017) Sequence-dependent cargo recognition by SNX-BARs mediates retromer-independent transport of CI-MPR. *J Cell Biol* 216: 3695–3712. doi:10.1083/jcb.201703015

Suzuki SW, Chuang YS, Li M, Seaman MN, Emr SD (2019) A bipartite sorting signal ensures specificity of retromer complex in membrane protein recycling. *J Cell Biol* 218: 2876–2886. doi:10.1083/jcb.201901019
Tobys D, Kowalski LM, Czudaj E, Müller S, Zentis P, Pach E, Zigrino P, Blaeske T, Honing S (2021) Inhibition of clathrin-mediated endocytosis by knockdown of AP-2 leads to alterations in the plasma membrane proteome. Traffic 22: 6–22. doi:10.1111/tra.12770

Uemura T, Kametaka S, Waguri S (2018) GGA2 interacts with EGFR cytoplasmic domain to stabilize the receptor expression and promote cell growth. Sci Rep 8: 1368. doi:10.1038/s41598-018-19542-4

Uemura T, Waguri S (2020) Emerging roles of Golgi/endosome-localizing monomeric clathrin adaptors GGAs. Anat Sci Int 95: 12–21. doi:10.1007/s12565-019-00505-2

Waguri S, Dewitte F, Le Borgne R, Rouillé Y, Uchiyama Y, Dubremetz JF, Hoffack B (2003) Visualization of TGN to endosome trafficking through fluorescently labeled MPR and AP-1 in living cells. Mol Biol Cell 14: 142–155. doi:10.1091/mbc.e02-06-0338

Wahle T, Prager K, Raffler N, Haass C, Famulok M, Walter J (2005) GGA proteins regulate retrograde transport of BACE1 from endosomes to the trans-Golgi network. Mol Cell Neurosci 29: 453–461. doi:10.1016/j.mcn.2005.03.014

Wakana Y, van Galen J, Meissner F, Scarpa M, Polishchuk RS, Mann M, Malhotra V (2012) A new class of carriers that transport selective cargo from the trans Golgi network to the cell surface. EMBO J 31: 3976–3990. doi:10.1038/emboj.2012.235

Wang J, Fedoseienko A, Chen B, Burstein E, Jia D, Billadeau DD (2018) Endosomal receptor trafficking: Retromer and beyond. Traffic 19: 578–590. doi:10.1111/tra.12574

Wang S, Ma Z, Xu X, Wang Z, Sun L, Zhou Y, Lin X, Hong W, Wang T (2014) A role of Rab29 in the integrity of the trans-Golgi network and retrograde trafficking of mannose-6-phosphate receptor. PLoS One 9: e96242. doi:10.1371/journal.pone.0096242

Wassmer T, Attar N, Bujny MV, Oakley J, Traer CJ, Cullen PJ (2007) A loss-of-function screen reveals SNX5 and SNX6 as potential components of the mammalian retromer. J Cell Sci 120: 45–54. doi:10.1242/jcs.03302

Yanguas F, Moscaso-Romero E, Valdivieso MH (2019) Ent3 and GGA adaptors facilitate diverse anterograde and retrograde trafficking events to and from the prevacuolar endosome. Sci Rep 9: 10747. doi:10.1038/s41598-019-47035-5

License: This article is available under a Creative Commons License (Attribution 4.0 International, as described at https://creativecommons.org/licenses/by/4.0/).