VARP Is Recruited on to Endosomes by Direct Interaction with Retromer, Where Together They Function in Export to the Cell Surface

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SUMMARY

VARP is a Rab32/38 effector that also binds to the endosomal/lysosomal R-SNARE VAMP7. VARP binding regulates VAMP7 participation in SNARE complex formation and can therefore influence VAMP7-mediated membrane fusion events. Mutant versions of VARP that cannot bind Rab32:GTP, designed on the basis of the VARP ankyrin repeat/Rab32:GTP complex structure described here, unexpectedly retain endosomal localization, showing that VARP recruitment is not dependent on Rab32 binding. We show that recruitment of VARP to the endosomal membrane is mediated by its direct interaction with VPS29, a subunit of the retromer complex, which is involved in trafficking from endosomes to the TGN and the cell surface. Transport of GLUT1 from endosomes to the cell surface requires VARP, VPS29, and VAMP7 and depends on the direct interaction between VPS29 and VARP. Finally, we propose that endocytic cycling of VAMP7 depends on its interaction with VARP and, consequently, also on retromer.

INTRODUCTION

VARP (VPS9-domain ankyrin repeat protein) is a widely expressed, multidomain protein of 1,050 residues present in most animals (although lost from D. melanogaster and C. elegans), as well as in some species of unicellular organisms closely related to animals, including Capsaspora and Choanofo-

gellates. VARP is a Rab32/38 effector, and, in line with other Rab:Rab-effector systems, it has been assumed that Rab32 would be primarily responsible for recruiting VARP on to membranes (Tamura et al., 2009; Wang et al., 2008). VARP also possesses Rab21 GEF (guanine nucleotide exchange factor) activity (Zhang et al., 2006) and binds to VAMP7 (Figure 1A), an R-SNARE implicated in a variety of endocytic and exocytic fusion events (Bal et al., 2013; Fader et al., 2009; Monteiro et al., 2013; Moreau et al., 2011; Pryor et al., 2004; Rao et al., 2004) (reviewed in Chaineau et al., 2009). VARP binds to VAMP7 in its “closed,” prefusion conformation, and this interaction inhibits VAMP7-mediated SNARE complex formation by clamping its regulatory longin domain on to its SNARE motif (Schäfer et al., 2012). Through associations with GolginA4 and Kif5A, VARP has also been suggested to participate in a molecular network that regulates delivery of VAMP7-containing vesicles toward the cell periphery (Burgo et al., 2012), which could contribute to the process of neurite outgrowth (Burgo et al., 2009).

The Rab family of small GTPases plays a key role in protein sorting and membrane trafficking throughout eukaryotic cells (reviewed in Stenmark, 2009). Rab32 is an endosomal Rab GTPase, thought to have been present in the last eukaryotic common ancestor, with Rab38 arising in vertebrates as an evolutionary gene duplication of Rab32 (Klöpper et al., 2012). Rab32 (together with Rab7A and B, Rab7L1, Rab9A and B, and Rab23) is classified in the late endocytic group III Rabs (Klöpper et al., 2012), a group of Rabs with functions at multiple points along the endocytic system. For example, Rab7A mediates the recruitment of the retromer complex (Rojas et al., 2008; Seaman et al., 2009) and is also important for delivery to lysosomes (reviewed in Stenmark, 2009).

Rab32 is expressed in a variety of cell types (Cohen-Solal et al., 2003) and has been implicated in the biogenesis of, and
transport to lysosome-related organelles (LROs) such as melanosomes (Bultema et al., 2012; Park et al., 2007; Tamura et al., 2009; Wasmeier et al., 2006), autophagosomes (Wang et al., 2012), and platelet dense granules (Ambrosio et al., 2012). Rab38 is expressed in specialized cell types enriched in LROs (Osanai et al., 2005) and serves partially redundant functions with Rab32. Consistent with VARP functioning as a Rab32/38 effector, in melanocytes, VARP colocalizes with Rab32/38 on endocytic organelles and melanosomes (Wasmeier et al., 2006). In other cell types, VARP localizes primarily to endosomes but is also present on vesicular/tubular clusters, lysosomes, and the plasma membrane (Schäfer et al., 2012), a pattern of localization consistent with VARP’s multiple functions.

Figure 1. Structure of the VARP-ANKRD1:Rab32 Complex
(A) Schematic representation of VARP and Rab32 proteins, highlighting the functions of different domains. The domain color scheme depicted is used in all subsequent figures.
(B) Structure of the VARP (residues 451–640): Rab32(Q85L) heterodimer in “ribbon representation,” with GppCp shown as ball and stick.
(C) Orthogonal views of a VARP:Rab32 2:2 cross-dimer formed from two heterodimers that are in different asymmetric units. In the left-hand view, the membrane is located at the bottom, and in the right-hand view, the tetramer is viewed “through the membrane.”
(D and E) Surface representation, in (D), of the cross-dimer “opened out” as indicated to show the residues involved in the two different VARP:Rab32 interactions. The main residues involved are labeled in the zoomed-in views (bottom panels) and are shown highlighted in the same colors in the sequence alignment shown in (E).
See also Figure S1.
We set out to determine the molecular mechanism of the expected recruitment of VARP to endosomal membranes by Rab32/Rab38. We determined the crystal structure of the complex between the first ankyrin repeat domain of VARP with the GTP hydrolysis-deficient forms of Rab32 and Rab38. Unexpectedly, however, we found that mutations based on these structures, which abolished their interaction in vitro, did not prevent endosomal recruitment of VARP in vivo. Subsequently, we have identified the retromer subunit VPS29 as a direct binding partner of VARP. Precise mapping of the VPS29-VARP interaction shows this interaction recruits VARP on to endosomes, and we demonstrate that VARP, retromer, and VAMP7 operate together in a pathway that traffics the known retromer cargo GLUT1 from endosomes to the cell surface.

RESULTS

Structure of the VARP Ankyrin Repeat-Containing Domain:Rab32 Complex

VARP has been shown to interact with Rab32/38 via its first ankyrin repeat-containing domain (Tamura et al., 2009, 2011; Wang et al., 2008). Two fragments of VARP comprising residues 397–650 and 451–640 of VARP, which contains the first ankyrin repeat-containing domain (ANKRD1; Figure 1A), bind to Rab32 in a GTP-dependent manner with a dissociation constant, K_D, of 7 ± 3 μM and a K_G of 2.5 ± 0.4 μM, respectively (Figure 2 and Figure S1A available online). We were able to crystallize a 1:1 complex of the smaller VARP fragment (residues 451–640) with the GTP hydrolysis-deficient mutant of Rab32 (Q85L) in the presence of the nonhydrolyzable analog of GTP, GppCP. Its structure was solved at 2.8 Å resolution by molecular replacement using VARP ANKRD2 and Rab7:GTP as search models (Figure 1B; Table I). The final model comprises three molecules of the complex in the asymmetric unit, where VARP (residues 452–618), Rab32 (residues 22–198), and Rab32’s bound GppCP:Mg^{2+} are well defined in the electron density map (Figures S1C and S1D). The three complexes in the asymmetric unit all have the same structure, with root-mean-square deviation (rmsd) values of 0.32 Å (for 306 Cα atoms of molecules A and B) and 0.33 Å (for 313 Cα atoms of molecules A and C).

VARP ANKRD1 contains five tandem ankyrin repeats that form a slightly curved stack (Figure 1B). The structure of Rab32 bound to GppCP:Mg^{2+} resembles that of other effector-bound Rabs (reviewed in Lee et al., 2009). The P loop (residues 32–39) stabilizes the Mg^{2+}-coordinated triphosphate moiety of the nucleotide, while Switch-I (residues 48–61) folds over the nucleotide molecule and stabilizes the γ-phosphate of the GppCP. Switch-II (residues 84–97) also stabilizes the γ-phosphate but differs somewhat in conformation as compared to those in other GTP-bound Rab structures (Figure S1B).

The VARP ANKRD1:Rab32 heterodimer buries 1,530 Å^{2} of solvent accessible surface area (calculated with PISA; Krissinel and Henrick, 2007) (Figure 1D). The elongated interface consists of a central hydrophobic patch of residues from helices α2B (L513 and L514) and α3B (Y550 and Y551) of VARP and residues from Switch-I (V60), interswitch (F62, L69, and W80), and Switch-II (M91, V94, K97, and Y95) of Rab32. Flanking the hydrophobic core of the interface are two salt bridges formed through Switch-II of Rab32 (D480{VARP}-R93{Rab32}, H517{VARP}-K97{Rab32}) and a third one through the interswitch of the GTPase (K546{VARP}-D61{Rab32}). The residues involved in these interactions are conserved in other organisms (Figure 1E). Many of the key interacting residues in Rab32 are unique to Rab32/38, thus explaining the selectivity for VARP binding Rab32/38 over other endosomal Rabs, as demonstrated in “glutathione S-transferase (GST) pull-down” experiments (Figure S1E).

Mutational Analysis of the VARP-ANKRD1:Rab32 Complex Interface

Four residues within VARP that contribute to the complex interface, L513VARP, Y550VARP, Q509VARP, and K546VARP were mutated, and their ability to bind Rab32 in its GTP-locked state in solution was investigated (note that Q509VARP and Y550VARP have been previously proposed to participate in the VARP:Rab32/38 interaction; Tamura et al., 2011). Two double mutants, Q509A/Y550A and L513D/K546D, were also generated. The results of GST pull-down assays (Figure 2C) showed that VARP (wild-type [WT]) binds GSTRab32(Q85L) but not GST. VARP(Y550A) and VARP(K546D) showed reduced binding to GSTRab32:GTP, while the mutants Q509A, L513D, L513D/K546D, and Q509A/Y550A showed no detectable binding (Figure 2C). In broad agreement with these data, subsequent isothermal titration microcalorimetry (ITC) showed that the K_Gs for the interaction of these mutants were increased from 2.5 μM for WT VARP451-640 to ~8 μM for K546D, ~22 μM for Y550A, and ~30 μM for Q509A (Figure 2B). The L513D mutation reduced binding to below detectable levels (K_G > 300 μM), as did the double mutant Q509A/Y550A, but the L513D/K546D mutant was unstable in the constantly stirring environment of the ITC cell.

Mutating residues M91/R93 in Switch-II of Rab32, which participate in the interaction with VARP (Figure 2D), resulted in reduced binding as shown by pull down (Figure 2F) and no detectable binding by ITC (Figure 2E).

Dimerization of the Rab32:VARP Heterodimer

Many small GTPase:effector complexes are believed to form dimers, which increases both the apparent affinity and specificity of the recruitment of effectors to membrane-anchored GTPases (Lee et al., 2009; Panic et al., 2003). In the crystal, all VARP:Rab32 heterodimers form identical homodimers, resulting in the heterotetramers shown in Figures 1C and 1D. The majority of the heterodimer/heterodimer interface arises from the VARP molecules contacting the other Rab32 molecule in the tetramer through residues from Switch-I and Switch-II that are different...
from those involved in the VARP:Rab32 heterodimer formation. This second VARP:Rab32 interface buries 1,460 Å² of solvent accessible surface area, making the total solvent accessible buried surface area on tetramer formation around 6,400 Å², which suggests that the tetramer should be stable. Existence of a dimer of heterodimers in solution is supported by analytical gel filtration, which is consistent with the existence of a concentration-dependent equilibrium of a 2:2 tetramer and a 1:1...

Figure 2. Biochemical Analysis and In Vivo Role of the VARP:Rab32 Interaction
(A) Molecular details of the VARP:Rab32 heterodimer interface. Residues mutated in VARP are boxed in color.
(B) Effects on the binding of the VARP:Rab32 interaction of mutating key VARP residues in the interface as measured by ITC. Curves are color coded as in (A).
(C) GST pull-downs using GSTRab32(Q85L) and VARP451-640His6, showing the effect of mutating key residues on VARP in the interaction interface (A).
(D) Molecular details of VARP:Rab32 interface in orthogonal view to (A). Residues mutated in Rab32 are boxed in color.
(E) Effects on the VARP:Rab32 interaction of mutating key Rab32 residues in the interface as measured by ITC. Curves are color coded as in (D).
(F) GST pull-downs using GSTRab32(Q85L) and VARP451-640His6, showing the effect of mutating key residues on Rab32 in the interaction interface (D).
(G) GST pull-downs using GSTRab32(Q85L) and VARP451-640His6 showing the effect of mutating key residues on Rab32 in the interaction interface (D).
(H) IF confocal microscopy of cytosol-extracted HeLa cells stably expressing VARP-GFP constructs: WT, Q509A/Y550A (QY), L513D/K546D (LK), M684D/Y687S (MY), and Q509A/Y550A/M684D/Y687S (QY+MY). GFP, green; VPS26, red; nuclei, blue; merged panel (MERGE). Boxed regions in the merged panels are shown as separate green (top), red (middle), and merged (bottom) channels on the right. Colocalization coefficients for the VPS26 versus GFP signals were as follows: WT, 0.89; QY, 0.75; LK, 0.59; MY, 0.66; QY+MY, 0.64. Scale bar, 20 μm.

In (B) and (E), curves for data showing binding are the mean of a minimum of three experiments ± SD. In (C) and (F), top panels are gel stained with Coomassie blue, and the lower panels are western blots using an anti-His probe. See also Figure S2.
Table 1. Crystallographic Data for VARP ANKRD1:Rab32(Q85L) Crystals

| Parameter                        | Native       | Xenon        |
|----------------------------------|--------------|--------------|
| Space group                      | P3₂ 21       | P3₂ 21       |
| Number of complexes in ASU       | 3            | 3            |
| Unit cell (Å)                    | a = b = 144.4, c = 135.7 | a = b = 144.9, c = 137.0 |
| Wavelength (Å)                   | 0.9763       | 1.7          |
| Resolution range (Å)             | 72.22–2.80   | 72.45–3.30   |
|                                  | (2.91–2.80)  | (3.53–3.30)  |
| Beamline                         | I03          | I03          |
| Number of crystals               | 3            | 1            |
| Rmerge in top intensity bin      | 0.070        | 0.057        |
| Rmerge                           | 0.237 (2.215) | 0.318 (2.091) |
| Mean (|I|/SD|I|)     | 8.8 (0.9)    | 9.8 (2.3)    |
| Half-data set correlation        | 0.989 (0.229) | 0.996 (0.663) |
| coefficient CG₁/2                | 98.8 (94.1)  | 99.4 (98.9)  |
| Completeness (%)                 | 99.6 (99.1)  |              |
| Anomalous completeness (%)       | 7.4 (5.0)    | 17.3 (17.4)  |
| Multiplicity                     | 9.0 (8.9)    |              |
| Anomalous multiplicity           |              |              |
| Δ<sub>corr</sub> Correlation     | 0.146        |              |
| between half-sets                | (inner 0.763) |              |
| Wilson plot B (Å²)               | 54.1         | 79.3         |

Refinement

- Number of atoms: Protein 8,040, Ligand 99, Xe 9
- H₂O 145
- R-factor 0.19
- R<sub>free</sub> 0.24
- Number of reflections (number R<sub>free</sub>) 38,004 (2,011)
- Mean <B> (Å²) 74.2
- rmsd bond length deviation (Å) 0.011
- rmsd angle deviation (°) 1.589
- Ramachandran favored (%) 92.8
- Ramachandran outliers (%) 1.6

Var<sup>†</sup> means isolated in these screening studies, the endosomal localization and central role in endocytic trafficking of retromer led us to consider it as a candidate for the role of recruiting VARP to endosomes. In HeLa cells stably expressing VARP-GFP, small interfering RNA (siRNA)-mediated depletion of the retromer subunits VPS29 or VPS35, but not of Rab32 or VAMP7, resulted in loss of VARP-GFP but not of EEA1 recruitment to endosomes (Figures 3A and 3B; Figures S3A–S3D).

We next examined the subcellular colocalization of stably expressed VARP-GFP and the endogenous retromer and WASH complexes by immunofluorescence (IF) confocal microscopy. WT-VARP-GFP exhibited robust colocalization with the endogenous retromer subunit VPS26 on both vacuolar and tubular domains of endosomes (Figures 3C and 3D). The WASH complex subunit FAM21 (Harbour et al., 2012) colocalized with VARP and retromer on vacuolar domains of endosomes but was largely excluded from tubules (Figures 3C and S3E). In summary, the Rab5 effector EEA1 appeared that, in crystals, VARP ANKRD1:SeMet-Rab32Q85L and VARP ANKRD1:Rab38Q69L contain the same “cross-dimer of heterodimers” across multiple crystal forms, as compared to the wtVARP:Rab32Q85L complex (Tables 1 and S1).

In separate experiments, it had been observed that native immunoprecipitation of VPS29-GFP or GFP-VPS35 followed by mass spectrometry identified VARP as a potential interacting partner of retromer (Figure S3A), in addition to established retromer accessory proteins, e.g., the WASH complex and the RabGAP TBC1D5 (Harbour et al., 2010; Seaman et al., 2009). Although few VARP peptides than TBC1D5 and WASH complex peptides were isolated in these screening studies, the endosomal localization and central role in endocytic trafficking of retromer led us to consider it as a candidate for the role of recruiting VARP to endosomes. In HeLa cells stably expressing VARP-GFP, small interfering RNA (siRNA)-mediated depletion of the retromer subunits VPS29 or VPS35, but not of Rab32 or VAMP7, resulted in loss of VARP-GFP but not of EEA1 recruitment to endosomes (Figures 3A and 3B; Figures S3B–S3D).

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Figure 3. Retromer Recruits VARP on to Endosomes

(A) IF confocal microscopy of cytosol-extracted HeLa cells stably expressing VARP-GFP, which were knocked down using single siRNA oligonucleotides at 100 nM (NT, nontargeting control, VPS29-1, VPS35-1, VAMP7-1, Rab32-1). GFP, green; EEA1, red; nuclei, blue, merged panel (MERGE).

(B) Western blots of the cells imaged in (A) showing successful protein depletion.

(C) IF confocal microscopy of VARP-GFP cells. GFP, green; and either VPS26, Fam21, or EEA1, red; nuclei, blue, merged panels. Cells were either fixed intact (whole cell) or after cytosol extraction (saponin). Boxed regions in the merged panels are shown as separate green (top), red (middle), and merged (bottom) channels on the right. Colocalization coefficients for VPS26, FAM21, and EEA1 versus GFP were as follows, respectively (saponin only): VPS26, 0.56; FAM21, 0.16; EEA1, 0.67.

(D) IF confocal microscopy image of VARP-GFP cells. GFP, green; VPS26, red; the boxed regions (shown in the MERGE column) are enlarged below each image, demonstrating colocalization of VARP-GFP and VPS26 on both vacuolar and tubular domains of endosomes.

Scale bars, 20 μm. See also Figure S3.
to localize in many instances to the same endosomes as VARP, although it was differentially partitioned on those structures when compared with VARP, which colocalizes with retromer and the WASH complex (Figures 3A, 3C, and S3F).

**Identification and Characterization of the VARP:VPS29 Interaction**

We used a combination of yeast 2-hybrid (Y2H) analysis, GST pull-downs with bacterially expressed proteins, and in vivo siRNA knockdown and rescue experiments to identify the retromer subunit VPS29 as the direct binding partner of VARP. By Y2H, strains expressing both VARP and VPS29 exhibited growth, but no growth was observed when VARP was coexpressed with retromer subunits VPS26A, VPS26B, or VPS35 (Figure 4A). In agreement with these data, in GST pull-downs, full-length VARP bound to GST-VPS29 and a retromer subcomplex containing GST-VPS35, VPS26, and VPS29 with near-1:1 stoichiometry but not to a retromer subcomplex of GST-VPS35 and VPS26 (Figure 4B). The interaction was further mapped, using fully folded (Collins et al., 2005) mutant versions of VPS29 in both Y2H analysis and GST pull-downs, to a hydrophobic patch on the surface of VPS29, centered on Leu152 (Figures 4A, 4C, and S4A), which also binds to the RabGAP TBC1D5 (Harbour et al., 2010) and, possibly, very weakly to Snx1 (Collins et al., 2005; Swarbrick et al., 2011). In HeLa cells, endogenous VARP was coimmunoprecipitated with stably expressed WT-VPS29-GFP but not with L152E-VPS29-GFP (Figure S4B). In vivo, the loss of endosomal recruitment of VARP-GFP following VPS29 depletion was rescued by expression of an siRNA-resistant form of VPS29-TagRFP (red fluorescent protein) (red fluorescent protein) (Figures 4B and S4D). Consistent with the Y2H and biochemical interaction data, recruitment of VARP-GFP (or TBC1D5; Figures 4E and S4E) was not rescued on expression of the L152E-VPS29 mutant. The specificity of this effect was demonstrated by the fact that FAM21 localization, which was also lost from endosomes on VPS29 depletion due to destabilization of the retromer complex (Figure S4D), was rescued by both WT and L152E forms of VPS29 (Figure 4E), since FAM21 associates with retromer by binding to VPS35 (Harbour et al., 2010).
VPS29 Interacts with Two Cysteine-Rich Motifs within VARP

The evolution of VARP appears to have involved tandem duplication of a region containing six ankyrin repeats. Two regions of VARP (residues 410–450 and 692–730), which separate the first two ankyrin repeats, each contain a two cysteine-rich consensus motif with the sequence CHPLCXxxC that is conserved across all species (Figure 5A). Each region was shown independently to bind directly to VPS29 by Y2H (Figure 5C). Indeed, in ITC experiments, recombinant VARP fragments containing the Cys-rich motifs (residues 396–460 and 692–746) in isolation were able to bind to VPS29 with $K_D$s in the low micromolar range ($\sim$13 $\mu$M and $\sim$5 $\mu$M, respectively; Figure S5A).

The presence of a spatially conserved pattern of cysteines in an intracellular protein suggests the involvement of a metal: most likely, zinc. Consequently, elemental analysis by microbeam proton-induced X-ray emission (microPIXE) (Garman and Grime, 2005) was carried out on both bacterially expressed domains to investigate this possibility. The resulting data (Figure 5B and Table S2) indicate stoichiometric levels of Zn$^{2+}$ in both Cys-rich motifs and that no other nonorganic elements are present at levels above 0.02 atoms per motif. Attempts to abrogate the interaction between VARP and VPS29 in vitro by adding EDTA in an effort to chelate the bound Zn$^{2+}$ failed, presumably because of the difficulty of extracting tightly bound Zn$^{2+}$, as has been previously observed for other “double Zn-finger”-containing proteins (J. Deane, personal communication).

Mutation of all four Cys residues to Ser (4C1 or 4C2), or of the conserved His and Leu residues to Ser and Ala (HL1 or HL2), in the context of $\sim$40-residue fragments containing the first or the second Cys-rich motifs, respectively, abolished binding to VPS29 by Y2H (Figure 5C). In ITC experiments using similar fragments, the HL2 mutation reduced binding to undetectable levels, while the 4C2 mutation decreased binding by $\sim$20-fold (Figure S5A). In the context of full-length VARP, the HL1 and HL2 mutations reduced binding to VPS29 to below-detectable levels by Y2H (Figure 5C). However, in GST pull-downs, only simultaneous mutation of both HL1 and HL2 fully abolished VARP binding to VPS29 (Figure 5D). This agrees with data from HeLa cells stably expressing VARP-GFP, where extraction of non-membrane-associated proteins by a brief saponin wash prior to fixation resulted in only a partial loss of VARP-GFP from endosomes upon mutation of individual Cys-rich motifs, whereas simultaneous mutation of both motifs led to complete loss of VARP-GFP from endosomes (Figure 5E). This is in contrast to the effect of simultaneously mutating Rab32 and VAMP7 binding sites within VARP (Figure 2G). When these data are considered along with the small size of the VPS29 hydrophobic patch centered on L152 (Figure S4), the relative band intensities in Figures 4B and 5D, and the observation that a single Cys-rich motif binds to VPS29 with low micromolar $K_D$ affinity (Figure S5), the most likely scenario for the interaction between full-length VARP and VPS29 is a 1:1 stoichiometry with only one Cys-rich motif binding at any one time, with a resulting apparent $K_D$ in the high nanomolar range. In other words, binding is relatively strong because of the presence of two binding sites for VPS29 on the same VARP molecule. Unfortunately, neither the $K_D$ nor the stoichiometry of the binding of full-length proteins could be accurately determined, as all constructs containing both stacks of ankyrin repeats aggregated in ITC experiments, and all cocrystallization attempts have failed.

We therefore conclude that VARP is recruited to endosomes through the direct interaction of its Cys-rich motifs with VPS29, thus placing its ability to bind and regulate the fusogenic potential of free VAMP7, and to interact with Rab32:GTP, on the surface of endosomes. In agreement with this model, GST pull-down experiments (Figure 5F) showed that VARP simultaneously binds to VPS29, Rab32/38, and VAMP7 in vitro.

A VARP/Retromer/VAMP7-Dependent Pathway Plays a Role in GLUT1 Traffic out of Endosomes

Recent studies have suggested that a number of cell surface receptors and transporters recycle from endosomes to the plasma membrane by a retromer-dependent pathway (Piotrowski et al., 2013; Steinberg et al., 2013; Temkin et al., 2011). In light of our results showing that VARP associates with retromer, we hypothesized that deleting VARP and/or VAMP7 (which cycles from endosomes/endolysosomes through the plasma membrane [Pryor et al., 2008] and binds directly to VARP) should also disrupt the trafficking of retromer-dependent cargos. We chose to analyze the recently characterized retromer cargo GLUT1, which redistributes from a predominantly cell surface localization to an intracellular endosomal localization upon

Figure 5. VARP Binds to VPS29 via Two Conserved Zn$^{2+}$ Coordinating Cys-Rich Motifs

(A) The two conserved Cys-rich motifs (conserved residues highlighted in pink with asterisks beneath them) in VARP from different species. Residue numbering is given for the human sequence.

(B) Proton-induced X-ray emission (PIXE) spectra for protein fragments containing the first (residues 396–460, left) and second (residues 692–746, right) Cys-rich motifs, with the Zn and Br peaks labeled above them. Black dots indicate the individual data points, and the red line is the fit.

(C–E) In (C), Y2H analysis is shown of the DNA binding domain (BD) fused to VPS29 and the activation domain (AD) fused to the indicated VARP fragments. Mutations in the Cys-rich motifs are labeled as follows in (C), (D), and (E): (first motif) H432A/L434A = HL1, C431S/C435S/C437S/C440S = 4C1; (second motif) H712A/L714A = HL2, C711S/C715S/C717S/C720S = 4C2. (D) Coomassie-stained gel of GST pull-down experiment showing interaction of VARP-His$_{10}$ WT or the indicated VARP mutants with GST-VPS29 or with GST alone. (E) IF confocal microscopy of cytosol extracted HeLa cells stably expressing VARP-GFP WT or the indicated VARP mutants. GFP, green; VPS26, red; nuclei, blue, merged panels (MERGE). Colocalization coefficients of VPS26 versus GFP were as follows: WT, 0.9; HL1, 0.08; 4C1, 0.09; HL2, 0.07; 4C2, 0.1; HL1+HL2, 0; 4C1+ 4C2, 0. Scale bar, 20 $\mu$m.

(F) GST pull-down experiment showing the interaction of the indicated combinations of VARP-His$_{10}$, Rab32, and VAMP7 cytoplasmic domain (CD) (inputs, left three lanes), with either GST alone or GST-VPS29. The pull-down samples were western blotted for Rab32 and VAMP7. See also Figure S5.
VARP Is Recruited to Endosomes by Retromer
retromer depletion (Piotrowski et al., 2013; Steinberg et al., 2013). Consistent with these published findings, depletion of VPS29 in HeLa cells caused an increased intracellular distribution of GLUT1 as compared to control cells (Figure 6A; Figure S6A). A similar redistribution of GLUT1 was also observed upon knockdown of VARP and VAMP7, whereas knockdown of the clathrin adaptor AP-1 had no effect on GLUT1 distribution. We followed up on these initial observations of GLUT1 redistribution by measuring the colocalization of GLUT1 with the late endosomal/lysosomal membrane protein LAMP1, because Steinberg et al. (2013) had previously shown that depletion of the retromer subunit VPS35 increased such colocalization. We observed a marked increase in the colocalization of GLUT1 with LAMP1 after knockdown of VPS29, VARP, or VAMP7 (Figures 6B and 6C; Figures S6B and S6C) but, interestingly, not after depletion of the RabGAP TBC1D5 (Figures S6B and S6D) that binds to VPS29 at the same site as VARP (Harbour et al., 2010). Subsequent experiments with transiently transfected HeLa cells showed that siRNA-resistant WT VPS29-TagRFP, but not the L152E mutant that does not bind VARP, was able to rescue the increase in colocalization of GLUT1 with LAMP1 observed after knockdown of endogenous VPS29 (Figures 6D, 6E, and S6E). There was no statistically significant difference in colocalization of GLUT1 with LAMP1 after knockdown of VPS29 in cells expressing the siRNA-resistant L152E mutant when compared with nearby untransfected cells (Figures 6D and S6E). These data confirmed that the interaction between VARP and VAMP29 was necessary to rescue the GLUT1/LAMP1 colocalization phenotype observed after knockdown of VPS29.

Taken together, our data demonstrate that VARP and VAMP7 act within the retromer-dependent recycling pathway that traffics the glucose transporter GLUT1 from endosomes to the plasma membrane. Our rescue experiments show that the VPS29 requirement for this pathway is dependent on its binding to VARP. Given the role of retromer in mediating traffic out of endosomes, and the knowledge that VARP binds the retromer subunit VPS29 and VAMP7 at different binding sites, we decided to study further whether the intracellular distribution of VAMP7 is affected by its association with VARP/retromer. Stable HeLa cell lines expressing WT VAMP7-HA or the D69A, E71F, S73D (DES) mutant VAMP7-HA, which we have previously shown cannot bind VARP (Schäfer et al., 2012; Kent et al., 2012), were examined by IF confocal microscopy. The DES VAMP7-HA mutant was primarily distributed to the perinuclear region as compared to WT VAMP7-HA, which had a more peripheral endosomal localization in addition to a perinuclear pool (Figure 7A). This difference in localization was quantified by measuring the degree of colocalization between VAMP7-HA and the trans-Golgi network (TGN) marker TGN46, with DES VAMP7-HA exhibiting a higher degree of colocalization (Figure 7B). The increased colocalization of DES VAMP7-HA with TGN46 was lost on treatment of the cells with chloroquine (CQ), which inhibits the trafficking of proteins from endosomes to the TGN (Chapman and Munro, 1994; Kent et al., 2012; Reaves and Banting, 1994) but only weakly affects the distribution of TGN46 (Ponnambalam et al., 1996), unlike the more marked effect on its ortholog TGN38 in rodent cells (Chapman and Munro, 1994). Our data with the DES mutant of VAMP7 imply that the normal cycling of VAMP7 within the endocytic system is dependent on its interaction with VARP, in addition to its interactions with Hrb (Pryor et al., 2008) and AP3-Δ-adaptin (Kent et al., 2012). We also observed an increase in colocalization of WT VAMP7-HA with TGN46 after knocking down the retromer subunits VPS29 or VPS35 (Figures 7C and 7D).

**DISCUSSION**

Retromer drives trafficking from endosomes/endolysosomes via tubular-vesicular carriers to either the Golgi or the cell surface (Cullen and Korswagen, 2012; Seaman, 2012; Seaman et al., 2013). Here, we have shown that VARP is recruited to endosomes through a direct interaction with retromer, where it participates in the retromer-dependent pathway that delivers the glucose transporter GLUT1 to the plasma membrane (Figures 7C and 7D).

The current working model for retromer function suggests that, in mammalian cells, the VPS26/35/29 subcomplex associates with endosomal membranes via Rab7A (Rojas et al., 2008; Seaman et al., 2009) and Snx3 (reviewed in Cullen and Korswagen, 2012; Seaman, 2012). Cargo selective functions have been suggested for VPS26 (Fjorback et al., 2012), VPS35 (Arighi et al., 2004; Nothwehr et al., 2000; Seaman, 2007), and the associated sorting nexin, Snx27 (Ghai et al., 2013; Steinberg et al., 2013; Temkin et al., 2011). Retromer also interacts with the WASH complex, which promotes localized actin polymerization to provide a structural and/or force-generating scaffold on which...
Figure 7. The Steady-State Localization of VAMP7 Depends on Its Interaction with VARP
(A and B) HeLa cells stably expressing VAMP7-HA WT or DES were imaged by IF confocal microscopy: HA, green; TGN46, red; nuclei, blue, merged panels (MERGE). Cells were either left untreated or treated with 100 μM CQ for 2 hr prior to fixation. Seven independent fields (>10 cells per field) were imaged for each condition, and the colocalization coefficient measuring the colocalization of VAMP7-HA with TGN46 was obtained for each field (mean ± SEM in (B) with representative cell images in (A)).
endosomal tubular carriers may be formed (Derivery et al., 2009; Gomez and Billadeau, 2009; Harbour et al., 2010; Jia et al., 2012). The known destinations of retromer-derived carriers leaving endosomes are the Golgi and the cell surface (Cullen and Korswagen, 2012; Fjorback et al., 2012; Seaman, 2007, 2012; Steinberg et al., 2013; Temkin et al., 2011). The interaction between retromer and VARP described here places VARP firmly in the trafficking pathway from endosomes to the cell surface and further cements the role of retromer as a hub for recruiting key machinery required for trafficking out of endosomes (Harbour et al., 2010) (Figure 7C). The binding of VARP by retromer is mediated by the VPS29 subunit and involves the same cluster of hydrophobic residues that mediates the association of retromer with the RabGAP TBC1D5. It has been proposed that binding of TBC1D5 by retromer leads to retromer dissociation from membranes through GTP hydrolysis of Rab7A, a Rab key to retromer recruitment (Harbour et al., 2010; Seaman et al., 2009) VARP association with retromer may inhibit the binding of TBC1D5, leading to a more persistent association of retromer with the membrane, thereby facilitating the sorting of cargo proteins into the endosome-to-cell surface pathway. Retromer was first identified in yeast (Seaman et al., 1997, 1998) and was shown to mediate endosome-to-Golgi retrieval. To date, there is no compelling evidence that retromer in yeast can drive endosome-to-cell surface traffic, and VARP is not present in yeast, consistent with the hypothesis that VARP regulates the activity of retromer in the endosome-to-cell surface pathway in mammalian cells (Gerondopoulos et al., 2012; Kloer et al., 2010; Nottingham et al., 2011; Rojas et al., 2008; Seaman et al., 2009; Stenmark, 2009). The reason for the VARP:Rab32 interaction remains unclear. As it is not required for VARP recruitment to endosomal membranes, we did not pursue it further in this study. However, our data are consistent with a model in which VARP is initially recruited to the endosome through its interaction with retromer. Subsequently, the VARP:retromer complex may either be partitioned into or contribute to the formation of a Rab32-defined endosomal subdomain, from which trafficking out of the endosome may occur.

VAMP7 mediates fusion of both endosomes and endolysosomes with lysosomes and fusion of endolysosomes and their derived carriers with the plasma membrane (reviewed in Chaineau et al., 2009). VAMP7 is then returned to its steady-state endolysosomal localization through interactions with Hrb and AP-3 (Boucrot and Kirchhausen, 2007; Chaineau et al., 2008; Kent et al., 2012; Martinez-Arca et al., 2003; Pryor et al., 2004, 2008; Rao et al., 2004). The demonstration that VARP can interact simultaneously with retromer and VAMP7 suggests other potential reasons why the VARP:retromer interaction may be important in trafficking out of endosomes. While Hrb and AP-3 appear to be the major factors determining the endosomal localization of VAMP7 (see Kent et al., 2012), a role for VARP in VAMP7 distribution within the endosomal compartment is supported by our observation that a VARP binding-deficient mutant of VAMP7 is mislocalized away from endosomes in a similar manner to that observed in retromer-knockdown cells. The redistribution of VAMP7 to a TGN localization after loss of VARP interaction suggests that VARP:retromer is necessary to direct VAMP7 away from a retromer-independent endosome-to-Golgi retrieval pathway such as that used by furin (Chia et al., 2011). Alternatively, VAMP7 redistribution to the TGN could be caused, at least in part, by VARP operating at the TGN to sort VAMP7 into vesicles for transport to endosomes (Burgo et al., 2012). However, this is difficult to reconcile with the inability to detect obvious localization of WT VARP at the TGN in nonneuronal cells by IF (Zhang et al., 2006) or immunoelectron microscopy (Schäfer et al., 2012).

The incorporation of VAMP7 into retromer-dependent endosome-derived carriers could facilitate the subsequent fusion of these carriers with their final destinations (see also Burgo et al., 2012), consistent with our observation that both VAMP7 and retromer are required for delivery of the glucose transporter GLUT1 to the plasma membrane. In addition, once VARP is concentrated on an endosomal membrane by retromer, VARP has the potential to regulate the fusogenic ability of VAMP7 (Schäfer et al., 2012). Therefore, it is possible that control of endosome/lysosome fusion could, in part, be regulated by the amount of retromer on the endosome surface through its ability to recruit VARP.

In summary, through its multiple interactions, VARP can link the retromer complex (VPS26/35/29) and its associated proteins (including sorting nexins and the WASH complex) to both a key ancestral R-SNARE protein (VAMP7) (Filippini et al., 2001) involved in many membrane fusion events (Chaineau et al., 2009) and to an endosome-localized Rab (Figure 7). Thus, we propose that VARP is a key component of a protein network that both drives and controls traffic out of endosomes.

**EXPERIMENTAL PROCEDURES**

**Protein Expression and Purification**

Protein expression was performed in E. coli Rosetta 2 pLysS. Cells were grown at 37°C, and expression was induced by the addition of 500 μM IPTG. VARP expression was carried out at 16°C for 20–24 hr, while all other proteins were expressed at 22°C for 14–18 hr. Cells were lysed in 20 mM HEPES, pH 7.4, and 500 mM NaCl (buffer A), supplemented with 5 mM MgCl2, 1 mM MnCl2, 5 mM dithiothreitol (DTT), 4 μg/ml DNase, and protease inhibitors and AEBSF. Seleno-methionine proteins were expressed in E. coli B834 using M9 media supplemented as in Collins et al. (2008).

**Protein Purification and VARP:Rab Complex Reconstitution**

Proteins were first purified on GSH-Sepharose, and the GST tag was cleaved off by incubating the beads with PreScission Protease (GE Healthcare).
overnight at 4°C. His-tagged VARP1–1050His10 was further purified on Ni-NTA agarose. All proteins were purified by gel filtration on Superdex 75 in buffer A supplemented with 10 mM DTT and additionally with 5 mM MgCl2 and 10 mM GppCp or GTP–S in the case of Rab32/38. Complexes were formed by mixing VARP with a GTPase in a molar excess of 1:1.25 for 1 hr at 4°C and finally subjecting them to gel filtration on Superdex 75 equilibrated in buffer A containing 5 mM MgCl2, 10 mM GppCp or GTP–S, and 10 mM DTT. Fractions of 1:1 complex as assessed by SDS-PAGE were pooled and concentrated for crystallization. Proteins for ITC were gel filtered on Superdex 75 in 50 mM HEPES, pH 7.4, 200 mM NaCl, 5 mM MgCl2, 1 mM TCEP.

Protein Purification for Pull-Downs

VARP1–1050His10 WT and mutants were purified in 500 mM NaCl, 20 mM Tris pH8.5, buffer B as in (Schäfer et al., 2012). GST-tagged proteins were purified on GSH-sepharose and eluted in buffer B containing 30 mM reduced glutathione, or the GST was removed by overnight incubation with PreScission Protease or thrombin as appropriate. Proteins were finally purified by gel filtration on Superdex 200 in buffer B that, in the case of Rabs, was supplemented with 5 mM MgCl2 and 10 μM of matching nucleotide.

Crystallization and Structure Determination

Crystals of VARP1–1050His10, Rab32–225(Q85L) were grown by sitting drop vapor diffusion at 18°C against well solutions containing 14%–19% (v/v) of polyethylene glycol 3350 and 200 mM sodium citrate, pH 8.0, at an initial complex concentration of 2.5 mg/ml. Cryoprotected crystals exhibited anisotropic diffracting at 100 K and diffracted in the best directions to 2.8 Å. The structure was solved by molecular replacement with Phaser (McCoy et al., 2007), using models based on the structures of Rab7 (Protein Data Bank [PDB] ID 1YHN) and the VARP-ANKR domain 2 structure (PDB ID 4B93). Crystals of VARP-ANKR domain 1:SeMet– Rab32–225(Q85L);Q58/HL/V100M/Q153M/V158M/I192M and of VARP-ANKR domain 1:SeMet–Rab38 (Q89L) diffracted in their best directions to 3 Å and 4.5 Å, respectively. Their structures were solved by molecular replacement using the coordinates of native VARP-ANKR domain 1:Rab32Q85L complex as the search model. For a full description of structure determination, see Supplemental Information.

Protein-Protein Interaction Studies

Y2H for screening and analyzing specific interactions was performed using pGBT9 and pGAD-C vectors and the HF7c reporter strain as previously described for Y2H for screening and analyzing specific interactions was performed using pGBT9 and pGAD-C vectors and the HF7c reporter strain as previously described elsewhere (Weekes et al., 2013). For mass spectrometric analysis, four dishes (140 mm in diameter) of cells were used for each cell line. For western blotting analysis, a single dish of cells was used for each cell line, and the washed protein-A sepharose was dissociated in a SpeedVac prior to resuspension in SDS-PAGE sample buffer omitting the glycine elution and acetone precipitation steps.

Mammalian Cell Culture, Microscopy, and Western Blotting

HeLa cells stably expressing VPS29-EGFP or EGFP-VPS35 were generated using the pLXIN retroviral system (Clontech). When indicated, HeLa cytosol was extracted prior to fixation by rinsing cells briefly with PBS, followed by incubation for 1 min in PBS (with Ca2+ and Mg2+) containing 0.05% saponin, and then immediately formaldehyde fixed. IF confocal microscopy and western blotting were carried out essentially as described elsewhere (Schäfer et al., 2012). For quantitative IF confocal microscopy of GLUT1 and LAMP1 colocalization, cell fields (three independent experiments with five fields each per condition, average of ≥2 cells per field) were randomly selected based on nuclear stain and focused using the LAMP1 signal. Single confocal images were acquired corresponding to 1 Axy unit, and the degree of colocalization of two channels was measured by Manders M1/M2 colocalization coefficients (Manders et al., 1993) using Zen software (Carl Zeiss). To negate the effects of inherent cell-to-cell variability of GLUT1 surface staining, the fraction of LAMP1 signal colocalizing with GLUT1 was measured for each field.

ACCESS NUMBERS

The PDB accession number for the structures presented here is 4CYM, and for structure factors, it is 4cymsf.

SUPPLEMENTAL INFORMATION

Supplemental Information includes Supplemental Experimental Procedures, six figures, and two tables and can be found with this article online at dx.doi.org/10.1016/j.devcel.2014.04.010.

AUTHOR CONTRIBUTIONS

I.P.-D., I.B.S., G.G.H., A.J.M., and P.R.E. carried out work on the Rab32:VARP complex. G.G.H., L.P.J., L.W., I.B.S., S.R.G., M.E.H., M.N.J.S., and D.J.O. carried out work on VARP:Retromer. O.B.Z. and E.F.G. carried out the microPIXE analysis. J.P.L., M.N.J.S., and D.J.O. conceived and initiated the study. All authors contributed to the design of the study and the writing of the manuscript.

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