Retromer Binding to FAM21 and the WASH Complex Is Perturbed by the Parkinson Disease-Linked VPS35(D620N) Mutation

Summary

Retromer is a protein assembly that plays a central role in orchestrating export of transmembrane-spanning cargo proteins from endosomes into retrieval pathways destined for the Golgi apparatus and the plasma membrane [1]. Recently, a specific mutation in the retromer component VPS35, VPS35(D620N), has linked retromer dysfunction to familial autosomal dominant and sporadic Parkinson disease [2, 3]. However, the effect of this mutation on retromer function remains poorly characterized. Here we established that in cells expressing VPS35(D620N) there is a perturbation in endosome-to-TGN transport but not endosome-to-plasma membrane recycling, which we confirm in patient cells harboring the VPS35(D620N) mutation. Through comparative stable isotope labeling by amino acids in cell culture (SILAC)-based analysis of wild-type VPS35 versus the VPS35(D620N) mutant interactomes, we establish that the major defect of the D620N mutation lies in the association to the actin-nucleating Wiskott-Aldrich syndrome and SCAR homolog (WASH) complex. Moreover, using isothermal calorimetry, we establish that the primary defect of the VPS35(D620N) mutant is a 2.2 ± 0.5-fold decrease in affinity for the WASH complex component FAM21. These data define the primary molecular defect in retromer assembly that arises from the VPS35(D620N) mutation and, by revealing functional effects on retromer-mediated endosome-to-TGN transport, provide new insight into retromer deregulation in Parkinson disease.

Results and Discussion

Endosome-to-TGN Transport of CI-MPR Is Impaired in VPS35(D620N)-Expressing Cells

We sought to investigate the effect of the VPS35(D620N) mutation on trafficking of two known retromer cargos, the endosome-to-TGN transport of the cation-independent mannose 6-phosphate receptor (CI-MPR) [4–7], and the endosome-to-plasma membrane transport of the glucose transporter GLUT1 [8]. At steady state, GLUT1 is localized at the plasma membrane from where it undergoes continuous rounds of endocytosis and PDZ ligand-dependent endosome-to-plasma membrane recycling [9], the latter being mediated by the SNX27-retromer [5]. In the absence of retromer, GLUT1 accumulates in the lysosome and is degraded [6]. To establish whether retromer-mediated endosome-to-plasma membrane transport was affected by the VPS35(D620N) mutation, we performed a quantitative analysis of GLUT1 surface abundance, lysosomal localization, and kinetics of lysosomal-mediated degradation [8]. In HeLa or RPE1 cells, the depletion of endogenous VPS35 by siRNA-mediated suppression followed by re-expression of either wild-type GFP-VPS35 or GFP-VPS35(D620N) produced cell lines where the GFP-tagged VPS35 transgenes were expressed at near to endogenous levels (Figure 1C). In these cell lines, expression of GFP-VPS35 or GFP-VPS35(D620N) efficiently rescued the lysosomal mis-sorting of GLUT1 observed upon VPS35 suppression (Figures 1A and 1B) (see Figure S1A available online for split channels and Figure S1B for a larger field of view). Furthermore, while the knockdown of VPS35 in RPE1 cells led to a pronounced decrease of GLUT1 surface abundance, re-expression of wild-type or mutant VPS35 rescued GLUT1 surface abundance (Figures 1A and 1C). Finally, an analysis of GLUT1 degradation kinetics in the RPE1 cells revealed that the post-translational stability of GLUT1 was not affected by the VPS35(D620N) mutation (Figures 1Di–1Diii); the degradation of transferrin receptor was also monitored as a negative control and as expected its degradation rate was also unaffected by the VPS35(D620N) mutation. Overall, these data establish that the VPS35(D620N) mutation does not impair retromer-mediated endosome-to-plasma membrane transport of GLUT1.

Next, we examined the endosome-to-TGN transport of the CI-MPR. After delivery to endosomes, CI-MPR dissociates from its ligand and is recognized by the retromer complex and retrieved to the TGN for further rounds of ligand binding and transport [4–7]. In the absence of retromer, the efficiency of CI-MPR retrieval is perturbed and an increase in endosomal localization of CI-MPR is observed [4–7]. At steady state in our RPE1 cell line, CI-MPR was predominantly localized to the TGN (Figure 1Ei). Suppression of VPS35 led to an increase in the amount of CI-MPR on dispersed puncta and a decrease in the Pearson’s correlation between CI-MPR and TGN46 (Figures 1Ei and 1F). This dispersal and decrease in Pearson’s correlation was partially rescued by re-expressing GFP-VPS35 but not GFP-VPS35(D620N) (Figures 1Ei and 1F). The punctate CI-MPR staining in the VPS35(D620N)-expressing
cells was positive for VPS35 (Figure 1Eii), and there was an increase in the overlap between CI-MPR and VPS35 in the GFP-VPS35(D620N) cells when compared to GFP-VPS35-expressing cells (Figure 1G), consistent with a defect in endosome-to-TGN transport and a corresponding CI-MPR dispersal. To extend this, we also examined the steady-state distribution of CI-MPR in fibroblasts obtained from a healthy donor and from a patient harboring the VPS35(D620N) mutation. Again an increased dispersal of CI-MPR was observed (Figures 1H and 1I). This dispersal was not due to a fragmented Golgi as both TGN46 and GRASP65 distribution appeared normal in VPS35(D620N) fibroblasts (Figure S1C). The CI-MPR dispersal observed in the VPS35(D620N) fibroblasts could be rescued by suppression of endogenous VPS35 and re-expression of wild-type GFP-VPS35 (Figure 1I), suggesting that the differences in CI-MPR distribution are not due to variability between cell lines isolated from different patients. These data confirm and extend recent findings of an impairment in
The VPS35(D620N) Mutant Does Not Perturb the Formation of the Retromer Heterotrimer or the Endosome Localization of VPS35

To determine the molecular basis of the perturbed endosome-to-TGN trafficking observed with the VPS35(D620N) mutant, we first sought to determine whether the VPS35(D620N) mutation affected formation of the VPS26-VPS29-VPS35 heterotrimer or its endosomal localization. Like GFP-VPS35, GFP-VPS35(D620N) localized predominantly to a sorting nexin-1 (SNX1) positive early-to-late transitional endosomal compartment (Figures 2A and 2B). In agreement with this, VPS35 showed normal distribution and appearance of the retromer-decorated endosomal compartment in the fibroblast cells derived from patient samples as well as comparable colocalization with SNX1 and the early endocytic marker EEA1 (Figures 2C and 2D). Immunoisolation followed by western blotting also established that VPS35(D620N) retained the ability to form a heterotrimeric complex with VPS29 and VPS26A (Figure 2E). These data establish that the molecular defect associated with the VPS35(D620N) mutation lies outside of heterotrimeric retromer assembly and retromer’s endosome association.
Comparative Interactome Analysis of VPS35 and the VPS35(D620N) Mutant

Retromer scaffolds the assembly of a number of accessory proteins that aid its role in endosomal sorting [1]. To identify the effect of the VPS35(D620N) mutant on the assembly of these accessory proteins, we performed an unbiased quantitative analysis of the VPS35 and VPS35(D620N) interactomes. In a triple SILAC setup, RPE1 cell lines stably expressing GFP, GFP-VPS35 or GFP-VPS35(D620N) were grown in light (GFP), medium heavy (GFP-VPS35) and heavy (GFP-VPS35(D620N)) SILAC media until full steady-state labeling was ensured. Cells were then subjected to lysis, GFP trap immunoprecipitation and pooling of the samples for SDS-PAGE, and subsequent quantification by LC-MS/MS after in-gel tryptic digestion. Known VPS35-interacting partners VPS26A and VPS26B, VPS29, TBC1D5 [12], SNX27 [8], and all components of the actin-polymerizing Wiskott-Aldrich syndrome and SCAR homolog (WASH) complex [12–17] were enriched in the wild-type GFP-VPS35 immunoprecipitation (Tables S1 and S2). In addition, a number of previously unidentifed interactors including VARP, SDCCAG3, and ANKRD50 were identified, the functional relevance of which will be described elsewhere.

While there is comparative enrichment of VPS29, VPS26A, and VPS26B between wild-type and VPS35(D620N) (left side of Figure 2E; Tables S1 and S2), there was a clear decrease in the enrichment of all components of the WASH complex within the VPS35(D620N) interactome (Figure 2E; Tables S1 and S2). In addition, there was a decrease in enrichment of the known retromer interactor RME-8 [18, 19] and two of the newly identified interactors SDCCAG3 and ANKRD50 (Figure 2E; Tables S1 and S2). In contrast, there was no loss in binding to other known retromer interactors such as TBC1D5 [20] and SNX27 [8] and the newly identified interactor VARP [12] (Figure 2E; Tables S1 and S2). Validating the mass spectrometric quantification, fluorescence-based quantitative western analysis confirmed the decrease in binding of these proteins to the VPS35(D620N) mutant (Figures 2E and 2F). Given the loss of binding to the WASH complex observed in the GFP-VPS35(D620N) cells, we sought to investigate whether the membrane association of the WASH complex is impaired in the VPS35(D620N)-expressing cells. We saw no obvious impairment of FAM21 endosomal localization or a decrease in Pearson’s correlation between GFP-VPS35(D620N) and FAM21 (Figures 2G and 2H). However, we cannot rule out that the loss of binding to the WASH complex alters turnover of the WASH complex on the endosome or affects the endosomal subdomain organization of the retromer and WASH complexes.

The Binding of ANKRD50, SDCCAG3, and RME-8 to Retromer Is Mediated through the WASH Complex

To identify the primary defect in the VPS35(D620N) mutant, we performed a series of GFP-VPS35 immunosolubilizations from cells treated with RNAi’s targeting FAM21—the component of the WASH complex that directly binds to VPS35 [16], ANKRD50, SDCCAG3, and RME-8. Suppression of FAM21 resulted in a loss in the enrichment of ANKRD50, SDCCAG3, and RME-8 in GFP-VPS35 immunosolubilates (Figures 3A–3C), establishing that these proteins interact indirectly with retromer through the WASH complex. Suppression of ANKRD50, SDCCAG3, and RME-8 did not affect the binding of FAM21 to VPS35 (Figures 3A–3C).

To correlate these data with the previously described endosome association of SDCCAG3 and RME-8 [21, 22], we analyzed parallel experiments using immunofluorescence. Confirming published data, suppression of VPS35 led to the dissociation of FAM21 from endosomes (Figures 3D and 3E) [15, 16]. Consistent with the biochemical analysis, the endosome association of SDCCAG3 required the presence of both retromer and FAM21 (Figures 3D and 3E). In contrast, the endosome association of RME-8 was only partially dependent on retromer and FAM21 (Figures 3D and 3E), which is consistent with the known binding of RME-8 to other endosome-associated proteins including SNX1 [16]. Together, these data establish that the association of ANKRD50, SDCCAG3, and RME-8 to retromer is mediated through binding to the WASH complex, and for SDCCAG3 this is necessary for targeting to retromer-labeled endosomes. Moreover, they provide evidence that the major defect of the VPS35(D620N) mutant lies with the association to the WASH complex.

The VPS35(D620N) mutation has been suggested to inhibit the neuroprotective effect of VPS35 [23], so we sought to investigate whether suppression of VPS35, FAM21, SDCCAG3, RME-8, or ANKRD50 enhanced the toxicity and increased cell death observed in the neuroblastoma cell line SH-SY5Y upon exposure to MPP+, which interferes with complex I of the electron transport chain in mitochondrion, causing cell death. However, no increase in cell death upon suppression of any of these components was observed (Figure S2) when the cells were incubated in 500 μM of MPP+ for 48 hr. This is in line with a previous study that found no effect of VPS35 suppression on cell survival when SH-SY5Y cells were challenged with MPP+ [23]. However, the absence of an effect could also be due to insufficient suppression of these components in SH-SY5Y cells.

VPS35(D620N) Impairs Its Interaction with FAM21 In Vitro

Previous studies have suggested that the interaction between the retromer and FAM21 is mediated by the VPS35 and VPS29 (but not VPS26) subunits of the retromer heterotrimer and multiple LFa motifs from the C terminus of FAM21 [15, 16, 24]. FAM21 contains 21 such LFa motifs with varied affinities to the VPS35/VPS29 complex. The two most C-terminal ones, R20 and R21, bind to the VPS35/VPS29 complex with the highest affinity, and deletion of R20-21 abolished endosomal localization of FAM21 in cells [16]. Thus, to evaluate the effect of the VPS35(D620N) substitution, we examined whether this mutation affects the affinity of the VPS35/VPS29 complex to the R20-21 fragment by using isothermal titration calorimetry (ITC).

The recombinant VPS35/VPS29 complex was purified by co lysing individually expressing bacterial cells as described previously [16]. The VPS35(D620N) substitution did not affect complex formation with VPS29, further confirming our biochemical and proteomic analysis (Figures 2E and 2F). The ITC experiments for each VPS35/VPS29 complex were repeated three times with two independently prepared protein samples and slightly different injection protocols. Raw data were pooled and subjected to global fitting using the software package SEDPHAT. Similar to our previous studies, VPS35/VPS29 bound R20-21 with a stoichiometry of 1:2 [16]. The affinity of VPS35(D620N)/VPS29 to R20-21 (Kd = 4.6 ± 0.7 μM) was significantly lower (by 2.2 ± 0.5-fold) than that of the wild-type VPS35/VPS29 (Kd = 2.4 ± 0.6 μM) (Figures 4A and 4B). These results establish that the VPS35(D620N) substitution impairs its interaction with FAM21.

In addition to R20-21, we also attempted to measure the affinity for a larger fragment of FAM21, R15-21. Although the ITC thermograms were qualitatively consistent with a reduced
affinity for the VPS35(D620N) mutant, since the R15-21 fragment contains seven LFa motifs potentially interacting with VPS35/VPS29, it was not possible to quantitatively fit and interpret these data (data not shown).

In summary, stemming from an unbiased quantitative proteomic analysis, we have identified that at the molecular level the primary effect of the Parkinson disease-linked VPS35(D620N) mutation is a decrease in the affinity for FAM21 and as a consequence a reduced association with the WASH complex and its associated proteins, including RME-8, and the newly identified retromer-associated proteins SDCCAG3 and ANKRD50. While not perturbing retromer-mediated endosome-to-plasma membrane recycling of GLUT1, this partial loss of FAM21 binding and the WASH complex-associated proteins does lead to a decrease in the efficiency of endosomes-to-TGN retrieval of the CI-MPR. While the primary defect of the VPS35(D620N) mutation is a decrease in affinity for the WASH complex, it is important to stress that the observed phenotypes may arise from a reduced activity of RME8, SDCCAG3, or ANKRD50 or a combination of these and FAM21. This is consistent with the current view that WASH-mediated actin polymerization is required for the organization and maturation of retromer sorting domains on early-to-late transition endosomes [16]. Precisely why the decrease in coupling to the WASH complex, and associated proteins, leads to a selective retromer effect on endosome-to-TGN over endosome-to-plasma membrane transport is unclear and will require further investigation. With evidence linking mutations in the WASH complex component strumpellin to hereditary spastic paraplegia [25], and the recent description of an RME-8(N855S) mutation that segregates with late-onset Parkinson disease [26], the data presented here establish that the association and function relationship between retromer and WASH complexes will be an important area in defining the role of defective endosomal sorting in these neurodegenerative diseases.

During the final stages of reviewing our manuscript Zavodszky and colleagues, using a candidate approach, published work consistent with a defect in WASH binding to the VPS35(D620N) mutant [27]. Our detailed biochemical analysis extends their findings and, importantly, through our unbiased global analysis, highlights that the corresponding loss of...
RME-8, SDCCAG3, and/or ANKRD50 may also contribute to the defect in retromer function in Parkinson disease. As such, the role of these newly identified retromer interactors offers a potentially exciting avenue of research into the causes of this neurodegenerative disease.

Supplemental Information

Supplemental Information includes Supplemental Experimental Procedures, two figures, and two tables and can be found with this article online at http://dx.doi.org/10.1016/j.cub.2014.06.024.

Author Contributions


Acknowledgments

We specifically thank Dr. Kai Erdmann (University of Sheffield) for supplying reagents to study SDCCAG3. I.J.M. was supported by Wellcome Trust 4 Year PhD Studentship awarded through the Dynamic Cell Biology programme (083474). P.J.C. is supported by the Wellcome Trust (089928 and 085743), the Medical Research Council (MR/K018299/1), and the Royal Society. D.D.B. is supported by NIH grant AI065474. M.K.R. and D.J. are supported by the Howard Hughes Medical Institute and grants from the NIH (R01-GM56322) and Welch Foundation (I–1544).

Received: November 24, 2013
Revised: May 16, 2014
Accepted: June 10, 2014
Published: June 26, 2014

References


