The understudied links of the retromer complex to age-related pathways

Kenneth A. Wilson

Received: 5 February 2021 / Accepted: 30 July 2021 / Published online: 9 August 2021
© The Author(s) 2021

Abstract  Neuronal aging is associated with numerous diseases resulting in memory impairment and functional decline. A common hallmark of these disorders is the accumulation of intracellular and extracellular protein aggregates. The retromer complex plays a central role in sorting proteins by marking them for reuse rather than degradation. Retromer dysfunction has been shown to induce protein aggregates and neurodegeneration, suggesting that it may be important for age-related neuronal decline and disease progression. Despite this, little is known about how aging influences retromer stability and the proteins with which it interacts. Detailed insights into age-dependent changes in retromer structure and function could provide valuable information towards treating and preventing many age-related neurodegenerative disorders. Here, we visit age-related pathways which interact with retromer function that ought to be further explored to determine its role in age-related neurodegeneration.

Keywords  Retromer complex · Aging · Protein trafficking · Neurodegeneration · Alzheimer’s disease · Parkinson’s disease · Autophagy

Introduction

Neurodegenerative diseases such as Alzheimer’s, Parkinson’s, and Huntington’s are among the most prevalent age-related disorders [1–4]. These lead to loss of memory and motor control, and the associated decline in health is emotionally and financially costly [5]. Despite many hallmarks found for each of these diseases, such as the presence of amyloid-β extracellular plaques and intracellular hyper-phosphorylated Tau neurofibrillary tangles in Alzheimer’s disease or α-synuclein in Parkinson’s disease, there is no known way to prevent or cure many of them [6]. Clinical trials for treatments based on the clearance of these aggregates have proven disappointing [7], sparking investigations to find other means to stave off these age-related detriments.

Proper protein trafficking is essential for neuronal health as well as the processing of proteins which aggregate in neuronal decline [8]. In particular, endocytosed proteins must be sorted for degradation or reuse, and the balance between these two fates is intricate [9]. Proteins which are signaled for degradation are trafficked from the late endosome to the lysosome for autophagy. They can also be Trafficked directly to the plasma membrane or the trans-Golgi network when sorted through the retromer complex [10], named for its function of facilitating retrograde transport in neurons [11]. Dysfunction in the retromer complex has been implicated in numerous defects, including developmental problems in
simple organisms such as *Drosophila melanogaster* [12] and neurodegenerative phenotypes [13, 14]. It is also known to be important for age-related neuronal homeostasis [15], yet nothing is known about how the complex and its function are affected by age. How this mechanism declines, which mechanisms regulate its age-related function, and what it traffics to prevent neuronal decline are all questions that ought to be of paramount importance for understanding the mechanisms of neurodegeneration.

**The retromer mechanism**

The retromer complex localizes at the late endosome, interacting with the late endosomal protein Rab7 (Fig. 1) and other Rab7-interacting proteins such as TBC1D5 [16]. The complex core consists of a heterotrimer of vacuolar protein sorting 35 (VPS35), VPS26, and VPS29 [17]. This complex is responsible for recognizing cargo to be trafficked. Deletion of VPS26, VPS35, and in some contexts VPS29 can eliminate retromer-mediated trafficking altogether [12, 15, 18]. This trimer is conserved across species, and dysfunction of these components have been implicated in lifespan shortening and severe neurological defects in complex organisms [12, 19, 20].

A number of proteins interact directly with this trimer. Most prominently, sorting nexins are responsible for facilitating the trafficking of proteins. There are over 30 sorting nexins (SNXs) which interact with different stages of endocytosis and protein sorting [21]. Specifically, SNX1/2 [22] and SNX5/6 [23] as well as SNX3 [24] directly interact with the retromer to facilitate its activity. Depending on the specific SNXs which bind to the retromer, the cargo is then trafficked to different cellular loci [25, 26]. Proteins responsible for this trafficking include SNX27 [27], Rab4, which binds vesicles and cargo destined for the plasma membrane [27, 28], and Rab11, which mediates the trafficking of vesicles and cargo post-Golgi processing [29, 30].

![Fig. 1](image_url)

**Fig. 1** Endocytosis and the retromer complex. Proteins which are endocytosed are either degraded at the lysosome or are sorted by the retromer complex at the late endosome. Proteins sorted by the retromer are directed for fast transport or processing by the trans-Golgi network.
The retromer and neurodegenerative disease

The retromer complex has primarily been studied with reference to neurodegenerative diseases. Related to Alzheimer’s disease (AD), β-secretase (BACE) [31–34] and its cleavage target amyloid precursor protein (APP) [32, 35–37] are shuttled through the retromer complex. The cleavage of APP by BACE contributes to the toxic aggregation of amyloid-β. Stabilization of the retromer complex by the chaperone compounds R33 and R55 reduces the presence of amyloid-β [38, 39] as well as Tau neurofibrillary tangles [39]. VPS35 (D620N) mutations are associated with Parkinson’s disease (PD) phenotypes, dopaminergic neuron loss, and hippocampal Tau accumulation [40]. This association has been attributed to an interaction between VPS35 and LRRK2, a key protein which interacts with Parkin [41]. RNAi for VPS26 has also been shown to induce α-synuclein accumulation in fly neurons. Similar to AD phenotypes, this can be prevented with R55 treatment [12]. The retromer has also been implicated in Huntington’s disease (HD), as SorCS2, which is downregulated in R6/2 and zQ175 HD mouse models, which in turn downregulates VPS35 [42].

In addition to proteins which aggregate in specific diseases, the improper retromer-mediated sorting of other proteins has also been found to induce age-related neuronal decline. One of the earliest examples of retromer-mediated sorting was the trafficking of Wnt receptor. Wnt-mediated axon development was found to be severely impaired in nematode worms and frogs with retromer dysfunction [43, 44]. Improper Wnt function has been linked to AD [45] and PD [46], indicating the importance of properly regulating the components in Wnt signaling. Wnt signals induce the mitochondrial unfolded protein response and this process relies on retromer function [47], suggesting an essential role for the retromer in cellular stress response. Retromer function has also been linked to the trafficking of arrestins [48], which regulate signal transduction at GPCRs. Different types of arrestins regulate different cellular signals, including visual systems through photoreceptor signaling. Arrestin inhibition has been shown to alter photoreceptor morphology and light sensitivity [49] and arrestin function is also known to be altered by age [50]. The retromer complex has been specifically shown to be a key player in mediating beta-arrestin 1 function [48]. The relationship between the retromer’s regulation of arrestins with age remains unexplored, but could provide new insights into visual degeneration.

Links to age-related pathways

Despite the clear importance of retromer function in age-related neurodegenerative diseases, little is known about how age influences the retromer. It has been shown that retromer protein levels are reduced in the cortex of 14-month-old transgenic AD mice (Tg2576) [51], but it remains unknown how it is affected by normal aging. There are, however, numerous age-related pathways which have been linked to retromer function. These pathways need to be further characterized to determine how they impact (or are impacted by) retromer activity.

One aspect of age-related retromer function that requires further study is the relationship between the retromer and age-related pathways. One such crossover has been observed between proteins of the Sortilin family. As previously mentioned, SorCS2 has been linked to retromer sorting and HD [42]. Another protein within the same protein family, SORL1, has also been extensively linked to retromer dysfunction-related amyloid-β aggregation and AD phenotypes [52, 53]. SORL1 has previously been implicated in diabetes, providing an interesting direction for future studies comparing the incidence of diabetes and AD [54]. Relating to this, SNX6 has also been found to physically interact with insulin-like growth factor 1 receptor (IGF1R) [55]. This also stimulates the phosphorylation of AKT, influencing growth and aging signals [55]. Additionally, the phosphoinositide binding domain (PX) in SNXs is recruited by phosphatidylinositol-3-phosphate (PtdIns3P, or PI3P) at the endosomal membrane [56, 57]. PI3P activity is known to mediate signals downstream of AKT signaling upstream of retromer function [58]. This creates a challenge, as retromer function is necessary for healthy aging yet is regulated by age-promoting mechanisms. Interestingly, previous work has also demonstrated that insulin can induce retromer dissociation, which results in GLUT4 deficiency due to its over-degradation by the lysosome [59]. Taken together, these findings suggest that while retromer...
function can contribute to cell growth through insulin signaling, over-active insulin signaling can also cause retromer dysfunction. Significantly more dissection of this paradox is necessary to determine how to efficiently maintain retromer function without promoting mechanisms that contribute to aging.

Proper maintenance of the retromer complex also maintains healthy lysosomal function [18, 60, 61]. Studies using *Drosophila* models of retromer dysfunction have shown an abundance of aberrant lysosomes and photoreceptor necrosis [12, 61]. With fewer possible fates for endocytosed proteins in retromer dysfunction, it is possible that more proteins are being shuttled for autophagy, thus overburdening the lysosomes. This needs to be more extensively tested, but it provides an interesting counterpoint to many anti-aging therapies that aim for pathways which promote autophagy alone, such as dietary restriction and mTOR-targeting therapeutics like rapamycin. Interestingly, the only current link between mTOR activity and the retromer demonstrates that retromer activity is essential for mTOR to respond to dietary amino acids [62]. This suggests that retromer function contributes to mTOR activity, known to advance aging. These results indicate that retromer function surprisingly contributes to aging rather than providing an important function for age-related neuronal homeostasis [62]. Previously, it had also been shown that interaction between VPS35 and the Golgi-related protein GOLPH3 regulate mTOR activity [63], and that deletion of VPS29 and VPS35 influences sensitivity of cells to rapamycin [64].

There is a clear relationship between aging, the retromer, and neuronal health, there is ample room for study to determine how retromer decline takes place and what factors regulate its function. There are likely numerous environmental factors such as age and lifestyle choices which could influence these, and each of these could in turn affect the homeostasis of retromer proteins and the proteins which interact with it.

**Regulators of retromer function**

Retromer dysfunction has numerous outcomes due to it essential function, but it is worth noting that the most common resulting phenotypes are akin to age-related neurodegenerative diseases such as AD and PD. Despite clear links between age-related disease and the retromer complex, few factors are known which regulate retromer activity. One such factor that has been elucidated is phospholipase A2 group VI (PLA2G6), which is also linked to PD [12]. This study showed that VPS26 dysfunction led to accelerated photoreceptor decline with age in flies, and that this response was related to improper clearance of ceramides. They also showed direct interactions between PLA2G6, VPS26, and VPS35. PLA2G6 knockout caused α-synuclein accumulation which was alleviated by retromer stabilization. Similarly, CLN3 regulates retromer function through its interaction with Rab7 [65]. With such a clear relationship between aging, the retromer, and neuronal health, there is ample room for study to determine how retromer decline takes place and what factors regulate its function. There are likely numerous environmental factors such as age and lifestyle choices which could influence these, and each of these could in turn affect the homeostasis of retromer proteins and the proteins which interact with it.

**Conclusions**

The role of the retromer in maintaining homeostasis of proteins linked with age-related diseases is clear, but the means by which age influences this function is not known. Though some work has demonstrated interactions between retromer proteins and known aging-related pathways such as insulin and mTOR, there remain very few findings of how age is related to the maintenance of this mechanism, and there is potentially conflicting evidence which suggests that retromer function may promote age-related pathways. The balance between proper sorting of proteins for reuse or degradation is intricate, but its importance is clear. Understanding how age affects the key players in this process is an essential area of study to understand neuronal homeostasis.

**Acknowledgements** Thank you to Drs. Lisa Ellerby, Pankaj Kapahi, and Hugo Bellen for their mentorship, wisdom, and resources. Fig. 1 was created with BioRender.com.

**Funding** KAW is supported by NIH/NIA training grant T32AG000266-21.
Open Access  This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit http://creativecommons.org/licenses/by/4.0/.

References

1. Ballard C, et al. Alzheimer’s disease. Lancet. 2011;377(9770):1019–31.
2. James BD, et al. Dementia from Alzheimer disease and mixed pathologies in the oldest old. JAMA. 2012;307(17):1798–800.
3. Lees AJ, Hardy J, Revesz T. Parkinson’s disease. Lancet. 2009;373(9680):2055–66.
4. Walker FO. Huntington’s disease. Lancet. 2007;369(9557):218–28.
5. Alzheimer’s. A 2016 Alzheimer’s disease facts and figures. Alzheimers Dement. 2016;12(4):459–509.
6. Rafii MS, Aisen PS. Alzheimer’s disease clinical trials: moving toward successful prevention. CNS Drugs. 2019;33(2):99–106.
7. Huang YM, Shen J, Zhao HL. Major clinical trials failed the amyloid hypothesis of Alzheimer’s disease. J Am Geriatr Soc. 2019;67(4):841–4.
8. Ansell-Schutz A, et al. Reduced retromer function results in the accumulation of amyloid-beta oligomers. Mol Cell Neurosci. 2018;93:18–26.
9. Marusz T, et al. Retromer guarantees the degradation of autophagic cargo by maintaining lysosome function in Drosophila. Traffic. 2015;16(10):1088–107.
10. Williams ET, Chen X, Moore DJ. VPS35, the retromer complex and Parkinson’s disease. J Parkinsons Dis. 2017;7(2):219–33.
11. Seaman MN, McCaffery JM, Emr SD. A membrane coat complex essential for endosome-to-Golgi retrograde transport in yeast. J Cell Biol. 1998;142(3):665–81.
12. Lin G, et al. Phospholipase PLA2G6, a Parkinsonism-associated gene, affects Vps26 and Vps35, retromer function, and ceramide levels, similar to alpha-synuclein gain. Cell Metab. 2018;28(4):605-618 e6.
13. Reitz C. Retromer dysfunction and neurodegenerative disease. Curr Genomics. 2018;19(4):279–88.
14. Muzio L, et al. Retromer stabilization results in neuroprotection in a model of amyotrophic lateral sclerosis. Nat Commun. 2020;11(1):3848.
15. Ye H, et al. Retromer subunit, VPS29, regulates synaptic transmission and is required for endolysosomal function in the aging brain. eLife. 2020;9:e51977.
16. Xie J, et al. TBC1D5-catalyzed cycling of Rab7 is required for retromer-mediated human papillomavirus trafficking during virus entry. Cell Rep. 2020;31(10):107750.
17. Kovtun O, et al. Structure of the membrane-assembled retromer coat determined by cryo-electron tomography. Nature. 2018;561(7724):561–4.
18. Cui Y, et al. Formation of retromer transport carriers is disrupted by the Parkinson disease-linked Vps35 D620N variant. Traffic. 2021;22(4):123–136.
19. Wang HS, et al. In vivo evidence of pathogenicity of VPS35 mutations in the Drosophila. Mol Brain. 2014;7:73.
20. Linhart R, et al. Vacuolar protein sorting 35 (Vps35) rescues locomotor deficits and shortened lifespan in Drosophila expressing a Parkinson’s disease mutant of Leucine-Rich Repeat Kinase 2 (LRRK2). Mol Neurodegener. 2014;9:23.
21. Worby CA, Dixon JE. Sorting out the cellular functions of sorting nexins. Nat Rev Mol Cell Biol. 2002;3(12):919–31.
22. Rojas R, et al. 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. 2007;27(3):1112–24.
23. Sun Q, et al. Structural and functional insights into sorting nexin 5/6 interaction with bacterial effector IncE. Signal Transduct Target Ther. 2017;2:17030.
24. McGough IJ, et al. SNX3-retromer requires an evolutionary conserved MON2:DOPEY2:ATP9A complex to mediate Wntless sorting and Wnt secretion. Nat Commun. 2018;9(1):3737.
25. Cullen PJ. Endosomal sorting and signalling: an emerging role for sorting nexins. Nat Rev Mol Cell Biol. 2008;9(7):574–82.
26. Zhang H, et al. The retromer complex and sorting nexins in neurodegenerative diseases. Front Aging Neurosci. 2018;10:79.
27. Temkin P, et al. SNX27 mediates retromer tubule entry and endosome-to-plasma membrane trafficking of signalling receptors. Nat Cell Biol. 2011;13(6):715–21.
28. Gomez-Lamarca MJ, et al. Rme-8 depletion perturbs Notch recycling and predisposes to pathogenic signaling. J Cell Biol. 2015;210(2):303–18.
29. Inoshita T, et al. Vps35 in cooperation with LRRK2 regulates synaptic vesicle endocytosis through the endosomal pathway in Drosophila. Hum Mol Genet. 2017;26(15):2933–48.
30. Redpath GMI, et al. Membrane heterogeneity controls cellular endocytic trafficking. Front Cell Dev Biol. 2020;8:757.
31. Marks N, Berg MJ. BACE and gamma-secretase characterization and their sorting as therapeutic targets to reduce amyloidogenesis. Neurochem Res. 2010;35(2):181–210.
32. Okada H, et al. Proteomic identification of sorting nexin 6 as a negative regulator of BACE1-mediated APP processing. FASEB J. 2010;24(8):2783–94.
33. Finan GM, Okada H, Kim TW. BACE1 retrograde trafficking is uniquely regulated by the cytoplasmic domain of sortilin. J Biol Chem. 2011;286(14):12602–16.
34. Toh WH, et al. GGA1 regulates signal-dependent sorting of BACE1 to recycling endosomes, which moderates Abeta production. Mol Biol Cell. 2018;29(2):191–208.

35. Vieira SI, et al. Retrieval of the Alzheimer’s amyloid precursor protein from the endosome to the TGN is S655 phosphorylation state-dependent and retromer-mediated. Mol Neurodegener. 2010;5:40.

36. Sullivan CP, et al. Retromer disruption promotes amyloidogenic APP processing. Neurobiol Dis. 2011;43(2):338–45.

37. Choy RW, Cheng Z, Schekman R. Amyloid precursor protein (APP) traffic from the cell surface via endosomes for amyloid beta (Abeta) production in the trans-Golgi network. Proc Natl Acad Sci U S A. 2012;109(30):E2077–82.

38. Mecozzi VJ, et al. Pharmacological chaperones stabilize retromer to limit APP processing. Nat Chem Biol. 2014;10(6):443–9.

39. Li JG, et al. A pharmacological chaperone improves memory by reducing Abeta and tau neuropathology in a mouse model with plaques and tangles. Mol Neurodegener. 2020;15(1):1.

40. Chen X, et al. Parkinson’s disease-linked D620N VPS35 knockin mice manifest tau neuropathology and dopaminergic neurodegeneration. Proc Natl Acad Sci U S A. 2019;116(12):7565–74.

41. Mir R, et al. The Parkinson’s disease VPS35[D620N] mutation enhances LRRK2-mediated Rab protein phosphorylation in mouse and human. Biochem J. 2018;475(11):1861–83.

42. Ma Q, et al. SorCS2-mediated NR2A trafficking regulates motor deficits in Huntington’s disease. JCI Insight. 2017;2(9):e88995.

43. Prasad BC, Clark SG. Wnt signaling establishes anteroposterior neuronal polarity and requires retromer in C. elegans. Development. 2006;133(9):1757–66.

44. Coudreuse DY, et al. Wnt gradient formation requires retromer function in Wnt-producing cells. Science. 2006;312(5775):921–4.

45. Palomer E, Buechler J, Salinas PC. Wnt signaling deregulation in the aging and Alzheimer’s Brain. Front Cell Neurosci. 2019;13:227.

46. Marchetti B. Parkinson’s disease, aging and adult neurogenesis: Wnt/beta-catenin signalling as the key to unlock the mystery of endogenous brain repair. Aging Cell. et al2020;19(3):e13101.

47. Zhang Q, et al. The mitochondrial unfolded protein response is mediated cell-non-autonomously by retromer-dependent Wnt signaling. Cell. 2018;174(4):870-883 e17.

48. Nogueras-Ortiz C, et al. Retromer stops beta-arrestin 1-mediated signaling from internalized cannabinoid 2 receptors. Mol Biol Cell. 2017;28(24):3554–61.

49. Levine ES, et al. Rapid light-induced activation of retinal microglia in mice lacking arrestin-1. Vision Res. 2014;102:71–9.

50. Mayer RJ. From neurodegeneration to neurohomeostasis: the role of ubiquitin. Drug News Perspect. 2003;16(2):103–8.

51. Chu J, Pratico D. The retromer complex system in a transgenic mouse model of AD: influence of age. Neurobiol Aging. 2017;52:32–8.

52. Knupp A, et al. Depletion of the AD risk gene SORL1 selectively impairs neuronal endosomal traffic independent of amyloidogenic APP processing. Cell Rep. 2020;31(9):107719.

53. Dumanis SB, et al. Distinct functions for anterograde and retrograde sorting of SORLA in amyloidogenic processes in the brain. J Neurosci. 2015;35(37):12703–13.

54. Lane RF, et al. Diabetes-associated SorCS1 regulates Alzheimer’s amyloid-beta metabolism: evidence for involvement of SorL1 and the retromer complex. J Neurosci. 2010;30(39):13110–5.

55. Bareja A, et al. The proximity-labeling technique BioID identifies sorting nexin 6 as a member of the insulin-like growth factor 1 (IGF1)-IGF1 receptor pathway. J Biol Chem. 2018;293(17):6449–59.

56. Yong X, et al. Mechanism of cargo recognition by retromer-linked SNX-BAR proteins. PLoS Biol. 2020;18(3):e3000631.

57. Dong X, et al. Sorting nexin 5 mediates virus-induced autophagy and immunity. Nature. 2021;589(7842):456–61.

58. Virbasius JV, et al. Activation of the Akt-related cytokine-independent survival kinase requires interaction of its phox domain with endosomal phosphatidylinositol 3-phosphate. Proc Natl Acad Sci U S A. 2001;98(23):12908–13.

59. Ma J, et al. Prolonged insulin stimulation down-regulates GLUT4 through oxidative stress-mediated retromer inhibition by a protein kinase CK2-dependent mechanism in 3T3-L1 adipocytes. J Biol Chem. 2014;289(1):133–42.

60. Ravussin A, et al. The phosphatidylinositol 3-phosphate binding protein SNX4 controls ATG9A recycling and autophagy. J Cell Sci. 2021;134(3):jcs250670.

61. Tang FL, et al. Coupling of terminal differentiation deficit with neurodegenerative pathology in Vps35-deficient pyramidal neurons. Cell Death Differ. 2020;27(7):2099–116.

62. Kvaninickas A, et al. Retromer and TBC1D5 maintain late endosomal Rab7 domains to enable amino acid-induced mTORC1 signaling. J Cell Biol. 2019;218(9):2019–38.

63. Scott KL, et al. GOLPH3 modulates mTOR signaling and rapamycin sensitivity in cancer. Nature. 2009;459(7250):1085–90.

64. Xie MW, et al. Insights into TOR function and rapamycin response: chemical genomic profiling by using a high-density cell array method. Proc Natl Acad Sci U S A. 2005;102(20):7215–20.

65. Yasa S, et al. CLN3 regulates endosomal function by modulating Rab7A-effector interactions. J Cell Sci. 2020;133(6):jcs24047.

Publisher’s note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.