RESEARCH ARTICLE

SNX27–Retromer directly binds ESCPE-1 to transfer cargo proteins during endosomal recycling

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Abstract

Coat complexes coordinate cargo recognition through cargo adaptors with biogenesis of transport carriers during integral membrane protein trafficking. Here, we combine biochemical, structural, and cellular analyses to establish the mechanistic basis through which SNX27–Retromer, a major endosomal cargo adaptor, couples to the membrane remodeling endosomal SNX-BAR sorting complex for promoting exit 1 (ESCPE-1). In showing that the SNX27 FERM (4.1/ezrin/radixin/moesin) domain directly binds acidic-Asp-Leu-Phe (aDLF) motifs in the SNX1/SNX2 subunits of ESCPE-1, we propose a handover model where SNX27–Retromer captured cargo proteins are transferred into ESCPE-1 transport carriers to promote endosome-to-plasma membrane recycling. By revealing that assembly of the SNX27:Retromer:ESCPE-1 coat evolved in a stepwise manner during early metazoan evolution, likely reflecting the increasing complexity of endosome-to-plasma membrane recycling from the ancestral opisthokont to modern animals, we provide further evidence of the functional diversification of yeast pentameric Retromer in the recycling of hundreds of integral membrane proteins in metazoans.

Introduction

A central role of the endosomal–lysosomal network is to determine the sorting of integral proteins and their associated proteins and lipids, collectively termed “cargos,” between 2 fates, either degradation within lysosomes or retrieval from this fate for the promotion of recycling to a variety of organelles that include the cell surface [1]. Efficient and selective endosomal–lysosomal cargo sorting is essential for the function of all eukaryotic cells with increasing evidence pointing to defects in endosomal retrieval and recycling being at the heart of a wide array of human diseases, most notably those associated with neurodegeneration [2]. Establishing the molecular mechanisms and functional significance of endosomal cargo retrieval and recycling is therefore fundamental to understand eukaryotic cell biology.
Mechanism for endosomal sorting by SNX27-Retromer and ESCPE-1

Two general concepts are considered to account for the mechanism of endosomal cargo retrieval and recycling. Sequence-independent sorting posits that sorting can proceed without the need for any sequence information within the intracellular cytoplasmic domains of integral membrane cargos [3]. On the other hand, sequence-dependent sorting is controlled by specific signals in the cytoplasmic domain of cargos. These signals are typically short linear peptide motifs that direct the sorting process through coupling to ancient and evolutionary conserved coat complexes [1,4,5]. In coupling sorting motif recognition with membrane remodeling, these coats coordinate the process of cargo retrieval with the formation of cargo-enriched transport carriers for onward recycling. In mammals, these coats include sorting nexin-3 (SNX3)-Retromer [6–10], SNX27–Retromer [11–14], SNX17–Retriever [15], the CCC (CCDC22/CCDC93/COMMD) [16,17], and endosomal SNX-BAR sorting complex for promoting exit 1 (ESCPE-1) complexes [18,19] and various accessory proteins including the actin polymerizing Wiskott–Aldrich syndrome protein and SCAR homologue (WASH) complex [20,21].

Sorting nexin-27 (SNX27) is an endosome-associated cargo adaptor composed of a central PX (phox homology) domain flanked by an amino-terminal PDZ (PSD95, disks large, and zona occludens) domain and a carboxyl-terminal FERM (4.1/ezrin/radixin/moesin) domain [22,23]. The PDZ domain recognizes a specific sorting motif, termed the PDZ-binding motif (PDZbm), present at the very carboxyl termini of over 400 cargo proteins including receptors, transporters, channels, enzymes, and adhesion molecules [11–13,23–31]. In recognizing these internalized cargos, SNX27 regulates their retrieval from lysosomal degradation and promotes their recycling back to the plasma membrane [11–13].

To facilitate the retrieval process, the SNX27 PDZ domain associates with Retromer, a stable heterotrimer of VPS26 (A and B isoforms expressed in humans [32]), VPS35 and VPS29, to form the SNX27–Retromer [12–14]. Retromer itself assembles to form dimeric arches that may cluster SNX27 captured cargos [9,10,33] and associates with the WASH complex to allow the localized Arp2/3-dependent polymerization of branched filamentous actin [34,35]. In a poorly understood process, these assemblies and actin dynamics are proposed to generate a retrieval subdomain on the endosomal membrane that is enriched in cargo destined for recycling [35–39]. It is from these subdomains that further membrane remodeling ensues to drive the biogenesis of cargo-enriched tubular profiles and tubulovesicular carriers that physically transport cargo back to the cell surface. Importantly, defects in SNX27–Retromer function are associated with the pathobiology of human disease most notable neurological disorders and neurodegenerative disease [40].

A membrane tubulating Bin/Amphiphysin/Rvs (BAR) domain-containing protein associated with SNX27–Retromer tubular profiles is sorting nexin-1 (SNX1) [12]. SNX1, and its functional ortholog SNX2, is part of the ESCPE-1 complex [18,19,41], a membrane tubulating complex that in vitro can reconstitute the formation of membrane tubules with diameters similar to those observed in cellulo [42]. To be functional, ESCPE-1 forms a stable heterodimer of either SNX1 or SNX2 associated with either of SNX5 or SNX6, 2 additional BAR domain-containing orthologs [43]. Importantly, the mechanistic basis of how SNX27–Retromer delivers cargo into forming ESCPE-1 tubular profiles and tubulovesicular carriers remains a major unanswered question.

Here, we establish that a conserved surface in the FERM domain of SNX27 binds directly to acidic-Asp-Leu-Phe (aDLF) sequence motifs present in the disordered amino termini of the SNX1 and SNX2 subunits of ESCPE-1. We show that this interaction is required to handover PDZbm containing cargo proteins retrieved from lysosomal degradation by SNX27–Retromer into ESCPE-1 tubular profiles that promote cargo recycling for subsequent repopulation at the cell surface. Through evolutionary analysis of those protein–protein interactions that assemble
the SNX27-Retromer:ESCPE-1 coat complex, we reveal how this sorting axis has evolved to accommodate the increasing complexities of endosomal sorting observed in animals.

**Results**

**The FERM domain of SNX27 promotes cargo recycling following SNX27–Retromer–mediated retrieval from lysosomal degradation**

Previously, we have established that the interaction of Retromer with the PDZ domain of SNX27 is a crucial step in preventing the routing of internalized cargo into the lysosomal degradative pathway [13,14]. To identify the region(s) of SNX27 required for promoting subsequent cargo recycling to the plasma membrane, we used CRISPR gene editing to generate SNX27 knockout HeLa cells (Fig 1A). Compared to control cells, where endogenous GLUT1 was enriched at the cell surface, in the SNX27 knockout cells, GLUT1 displayed a steady-state enrichment with LAMP1-positive lysosomes, a phenotype that was also observed in a VPS35 knockout HeLa cell line (Fig 1B). Importantly, reexpression of GFP-SNX27 in the SNX27 knockout cells rescued the retrieval of GLUT1 from the lysosomal distribution and promoted its repopulation at the cell surface (Fig 1C). These data mirror those previously published using RNA interference (RNAi)-mediated SNX27 suppression [13] and confirm the SNX27 knockout HeLa cell line as a suitable screening platform through which to dissect the features of SNX27 required for promoting GLUT1 endosome-to-plasma membrane recycling.

In contrast to wild-type SNX27, expression of a SNX27 mutant lacking the loop region essential for Retromer association (GFP-SNX27Δ67–77) [13,14] failed to rescue retrieval from lysosomal colocalization and failed to promote GLUT1 cell surface recycling (Fig 1C). These data further confirm that the association of SNX27 with Retromer is a crucial step in coupling sequence-dependent cargo recognition with cargo retrieval from the lysosomal degradative fate [13,14]. Consistent with this, the expression of a SNX27 deletion mutant lacking the FERM domain (GFP-SNX27ΔFERM), a mutant that retained the association of SNX27 with Retromer, retained the ability to retrieve GLUT1 from lysosomes, as defined by a decreased colocalization between GLUT1 and LAMP1 (Fig 1C). However, the GFP-SNX27ΔFERM mutant failed to promote GLUT1 recycling to the cell surface, rather GLUT1 was enriched in a Retromer-positive endosomal compartment (Fig 1C). The overall levels of GLUT1 were not decreased in the KO and rescue conditions compared to parental cells (S1 Fig). Together, these data are consistent with a model in which the coupling of SNX27 to Retromer is necessary for retrieval from the lysosomal degradative fate, while the SNX27 FERM domain is required to promote the subsequent endosome-to-cell surface recycling of GLUT1.

**SNX1 and SNX2 couple ESCPE-1 to the FERM domain of SNX27**

ESCPE-1 regulates tubular-based endosomal recycling of several cargos, many of which are also dependent on SNX27–Retromer [13,18,19]. ESCPE-1 is a heterodimeric assembly of the functionally redundant SNX1 or SNX2 associated with the functionally redundant SNX5 or SNX6 (Fig 2A) [18,19], while SNX27–Retromer is an assembly of SNX27 with the VPS26A/B/VPS35: VPS29 heterotrimeric Retromer complex (Fig 2B) [13]. To define a possible connection between SNX27 and ESCPE-1, we first performed a series of GFP-nanotrap immunoisolation of SNX27 and various SNX27 mutants expressed in HEK-293T cells and probed for the presence of associated endogenous ESCPE-1. Robust association was observed between full-length GFP-SNX27 and all components of ESCPE-1 (Fig 2C–2E). This association was independent of the SNX27 PDZ domain as binding was retained in the PDZ deletion mutant GFP-SNX27ΔPDZ and was not observed when just the PDZ domain was expressed in HEK-293T cells (Fig 2C and 2D).
Retromer association with SNX27 was not required for ESCPE-1 binding as GFP-SNX27Δ67–77 and GFP-SNX27(L67A/L77A), 2 mutants that fail to bind to Retromer [14], each retained association with ESCPE-1 (Fig 2E). However, ESCPE-1 association was lost upon deletion of the SNX27 FERM domain (SNX27ΔFERM) or upon switching the FERM domain of SNX27 for the corresponding FERM domain from SNX17 (Fig 2C). Moreover, just the isolated FERM domain of SNX27, when expressed in HEK-293T cells, was able to associate with ESCPE-1 (Fig 2D).

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Finally, in the reverse immunoisolations, GFP-tagged SNX1, SNX2, SNX5, or SNX6 revealed that only SNX1 and SNX2 displayed a pronounced association with endogenous SNX27 when expressed in HEK-293T cells (Fig 2F). Moreover, this association did not require the presence of the SNX1 or SNX2 BAR domains, since deletion of these domains failed to affect binding to SNX27 (Fig 2G). Together, these data are consistent with the FERM domain of SNX27 associating with the SNX1 and SNX2 components of ESCPE-1.
The amino termini of SNX1 and SNX2 present tandem acidic-Asp-Leu-Phe motifs that associate with the SNX27 FERM domain

Compared to SNX5 and SNX6, human SNX1 and SNX2 each contain amino-terminal extensions of 142 and 139 amino acids, respectively, which are predicted to be unstructured. Expression in HEK-293T cells of GFP-SNX1(1–279), a construct encoding for the amino-terminal extension plus the SNX1 PX domain, and GFP-SNX1(1–138), a construct encoding for just the amino-terminal extension of SNX1, established that the amino-terminal extension of SNX1 is sufficient for association with endogenous SNX27 (Fig 3A). Likewise, the amino terminus of SNX2 interacted with SNX27, while those of SNX5 and SNX6 failed to do so (S2A Fig). To establish the direct nature of the association, we recombinantly expressed and purified full-length mouse SNX1 and the isolated FERM domain of human SNX27; mouse and human SNX1 are 94.2% identical. Isothermal titration calorimetry (ITC) established that SNX1 bound directly to the FERM domain of SNX27 with an affinity ($K_d$) of 41.0 μM (Fig 3B and S1 Table).

A similar binding affinity (39.1 μM) was observed with a recombinant protein corresponding to the amino-terminal extension of mouse SNX1 (85.9% sequence identity with human SNX1) (Fig 3B). To map down the features of SNX1 and SNX2 required for SNX27 binding, we performed a deletion mutagenesis screen of the amino-terminal extension of human SNX1 (Fig 3C). Walking upstream from a minimal binding region encoded by residues 1 to 138 established that 2 regions were required for maximal SNX27 binding. Deletion of residues 79 to 100 led to a pronounced reduction in binding and subsequent deletion of residues 41 to 54 abolished the interaction. Alignment of these regions of SNX1 revealed 2 similar motifs with DIF or DLF sequences flanked by upstream acidic residues. Comparison with the amino terminus of SNX2, which also binds SNX27 (S2A Fig), showed that it also has 2 similar sequences in its disordered amino-terminal domains, and, altogether, these define a conserved sequence D/E-x-x-D/I/L-F, here termed the aDLF motif (Fig 3D and 3E).

Site-directed mutagenesis targeting the first aDLF motif of SNX1 established that full-length GFP-SNX1(E42K), GFP-SNX1(E44K), GFP-SNX1(D45K), GFP-SNX1(I46K), and GFP-SNX1(F47K) all displayed reduced binding to SNX27 of around 50% (Fig 3F). A similar reduction in binding was also observed when targeting the second aDLF motif of SNX1 as observed with the GFP-SNX1(E84K), GFP-SNX1(D87K), and GFP-SNX1(F89D) mutants (Fig 3F). In a double GFP-SNX1(D45K,D87K) mutant that simultaneously targeted key residues in both aDLF motifs SNX27, binding was reduced to near undetectable levels (Fig 3F). Importantly, all mutants retained their ability to associate with SNX6, establishing that ESCPE-1 was still able to assemble. In addition, the mutant targeting both aDLF motifs in SNX2, GFP-SNX2(D26K,D72K), also reduced SNX27 binding to near undetectable levels (S2B Fig). Next, we used ITC to quantify the binding of aDLF peptides from SNX1 and SNX2 to full-length mouse SNX27. Peptides corresponding to residues 35 to 51 and 75 to 92 encompassing the first and second aDLF motifs of human SNX1, respectively, bound to SNX27 with respective affinities of 33.4 μM and 13.3 μM (Fig 3G and S1 Table). Establishing the redundant nature of the interaction with SNX27, we designed peptides corresponding to the first and second aDLF motifs of SNX2, namely residues 16 to 33 and 62 to 82. These also bound to full-length mouse SNX27 with affinities of 37.8 μM and 22.6 μM, respectively (Fig 3H). Further defining the redundancy between SNX1 and SNX2, introduction of a charge swap mutation into the SNX1 35–51 peptide, SNX1(D45K), and a triple mutation in the SNX2 16–33 peptide, SNX2(DLF-SSS), resulted in mutant peptides that showed no detectable binding to SNX27 (Fig 3G and 3H). Finally, a competitive ITC assay using full-length SNX27 preincubated with excess SNX1 75–92 peptide established that it competed for binding of SNX2 16–33 peptide, consistent with both sequences associating with the same surface within the SNX27 FERM domain (S2C Fig). Overall, these data establish
that through tandem aDLF motifs present within their unstructured amino terminal extensions SNX1 and SNX2 directly associate with the FERM domain of SNX27.
A basic surface in the SNX27 FERM domain is required for SNX1/SNX2 binding

SNX27 and SNX17 are related endosome associated cargo adaptors [23,45]. While they display entirely distinct functional roles in sequence-dependent endosomal cargo sorting [13,15], they share structurally similar PX and FERM domains [45,46] (Fig 4A). A comparative immunolocalization of GFP-SNX17 and GFP-SNX27 from HEK-293T cells established that SNX17 does not bind to ESCPE-1 (Fig 4B), consistent with the loss of ESCPE-1 binding to the SNX27 chimera containing the SNX17 FERM domain (Fig 2C). To identify potential regions for aDLF motif binding to the SNX27 FERM domain, we initially performed a comparative structural analysis between the X-ray structure of the SNX17 FERM domain [45] and a homology model of the SNX27 FERM domain [46]. In particular, we focused on the distribution and clustering of complementary basic amino acids. This revealed that the FERM domain of SNX27 possesses a basic surface defined in part by R437, R438, K495, K496, R498, and K501, residues that in SNX17 often corresponded to uncharged residues or negatively charged amino acids (Fig 4A and 4C). Indeed, charge swap site-directed mutagenesis of these residues generated full-length GFP-SNX27 mutants that when expressed and immunosolated from HEK-293T cells lacked the ability to associate with ESCPE-1; all mutants, however, retained the ability to associate with Retromer (Fig 4D). In direct ITC-quantified binding assays human SNX27(K495D), SNX27(R498D), and SNX27(K501D) all displayed no detectable binding to the human SNX1 75–92 peptide, while the SNX27(R496D) mutant displayed a reduce binding affinity of 56.4 μM when compared with the 12.7 μM observed for wild-type SNX27 (Fig 4E).

Finally, we modeled the binding of the aDLF sequences from SNX1 and SNX2 with SNX27 using the AlphaFold2 machine learning algorithm [47,48]. In total, we generated 12 models, with 3 models each of SNX27 in the presence of the 4 separate aDLF peptides (Figs 4F and S3A–S3C). Without any prior inputs apart from the SNX27 and peptide sequences, AlphaFold2 produced very high confidence models of the SNX27 FERM domain. Remarkably, in all 12 models, the aDLF peptides associated specifically with the SNX27 F3 subdomain, near the basic surface identified by mutagenesis (Fig 4F). Closer inspection showed that in 8 of the models the core D[L/I]F residues adopted essentially identical bound structures, with the hydrophobic [L/I]F side chains inserting into a hydrophobic pocket and interacting with SNX27 I467, F469, Y490, and V500 and the acidic Asp side chain surrounded by the complementary basic SNX27 residues R437, R438, K439, and R498 (Fig 4F). To further validate the model, we therefore mutated I467, Y490, and V500 to Lys, all of which disrupted the interaction of SNX27 with ESCPE-1 (Fig 4G). Together, these models are entirely consistent and strongly support the mutagenesis data for the SNX27-peptide interactions.

SNX27 association with ESCPE-1 couples cargo retrieval with promotion of cell surface recycling

To define the functional significance of the association of SNX27 to ESCPE-1, we first transfected our SNX27 knockout HeLa cells line with constructs encoding for either GFP-tagged wild-type SNX27 or the ESCPE-1 binding defective mutant GFP-SNX27(R498D) (Fig 5A). Analysis of their relative subcellular distributions established that they each associated with EEA1 and SNX6-positive endosomes (Fig 5B). Previously, we established that by binding to the PDZbm present within the cytoplasmic facing tail of GLUT1, the SNX27–Retromer complex serves to retrieve internalized GLUT1 away from lysosomal mediated degradation ([13], and see Fig 1A). We therefore performed a series of rescue experiments to quantify the SNX27-dependent retrieval of internalized GLUT1 from lysosomal mediated degradation, using the SNX17–Retriever-mediated retrieval of alpha-5 integrin as a negative control (Fig
Fig 4. SNX1 and SNX2 amino termini bind a unique basic surface of the SNX27-FERM domain. (A) Sequence alignment of SNX27 FERM domain with the SNX17 FERM domain. The positively charged residues that constitute a basic surface unique to SNX27 are boxed and labeled in blue. (B) Co-immunoprecipitation of GFP-tagged SNX27 and SNX17. The constructs were transiently transfected in HEK-293T cells, the cell lysates were subjected to GFP trap–based immunoprecipitation, and the immunoprecipitates were blotted for VPS35, SNX1, SNX2, SNX5, SNX6, and GFP. The blot is representative of 3 independent GFP traps. (C) Homology model of the SNX27 FERM domain [46]. The residues that are unique to SNX27 FERM domain and those that may account for the interaction with ESCPE-1 are labeled. (D) Co-immunoprecipitation of GFP-tagged SNX27 WT and a panel of amino acid swap mutants across the FERM domain. The constructs were transiently transfected in HEK293T cells, the cell lysates were subjected to GFP trap–based immunoprecipitation, and the immunoprecipitates were blotted for VPS35, SNX1, SNX2, SNX5, and SNX6. The band intensities were measured from \( n = 3 \) independent experiments using Odyssey software. The band intensity, normalized to GFP expression, is presented as the average fraction of the WT. The binding of SNX27 point mutants to VPS35, SNX1, SNX2, SNX5, and SNX6 was compared with the SNX27 WT using a 2-way ANOVA and Dunnett test. (E) Binding of SNX1\(^{75-92}\) peptide to hSNX27\(^{FL}\) proteins by ITC (hSNX27\(^{R498D}\): dark-blue; hSNX27\(^{K501D}\): orange; hSNX27\(^{K495D}\): lilac; hSNX27\(^{K496D}\): sky-blue; hSNX27 WT: red-orange; ND: no binding detected). The ITC graphs represent the integrated and normalized data fit with 1:1 ratio binding. The binding affinity (\( K_d \)) is given as mean of at least 2 independent experiments. Thermodynamic parameters for the binding are provided in S1 Table. (F) The left panel shows the AlphaFold2 generated model of the SNX27 FERM domain (pink ribbons) bound to the core aDLF sequences (yellow sticks) of SNX1 and SNX2 with identical binding orientations. For clarity, only one SNX27 model is shown with the 8 aDLF peptides overlaid (2 peptide models each of the 4 sequences from SNX1 and SNX2). The 3 subdomains of the SNX27 FERM domain F1, F2, and F3 are indicated. The middle panel shows a close-up view of the modeled SNX1 aDLF core sequence \(^{44}EDIFTGA\)\(^{50}\). The strictly conserved Phe side chain of the aDLF sequences is numbered \( F_0 \) for reference, and other residues are numbered with respect to this. (G) Co-immunoprecipitation of GFP-tagged SNX27 and a panel of
In the SNX27 knockout HeLa cells, GLUT1 displayed an enhanced rate of degradation following 8 hours of treatment with cycloheximide, data that are consistent with the loss of SNX27 expression leading to a reduced ability for transporter retrieval from the lysosomal degradative fate (Fig 5C). This phenotype was rescued by reexpression of wild-type GFP-tagged SNX27 (Fig 5C). Interestingly, expression of GFP-SNX27(R498D), a mutant that fails to associate with ESCPE-1 but retains binding to Retromer (Fig 4D and 4E), also fully rescued the retrieval of GLUT1 from lysosomal degradation (Fig 5C). The coupling of SNX27–Retromer to the ESCPE-1 complex is therefore not required for the retrieval of GLUT1 from lysosomal degradation.

Next, we turned to examining the promotion of GLUT1 recycling back to the cell surface. Here, we utilized restricted cell surface biotinylation to quantify the amount of GLUT1 at the cell surface [13]. Compared with control HeLa cells, SNX27 knockout HeLa cells displayed a reduced cell surface abundance of GLUT1, a result of missorting of internalized GLUT1 for lysosomal degradation (Fig 5D). Again, this phenotype was rescued by expression of GFP-SNX27 (Fig 5D) [13]. In contrast, expression of GFP-SNX27(R498D) failed to rescue the cell surface level of GLUT1 (Fig 5D), consistent with the association of SNX27 with ESCPE-1 being required to promote the cell surface recycling of internalized GLUT1. Importantly, GFP-SNX27(R498D) exhibited an endosomal localization indistinguishable from wild-type SNX27 (Fig 5B), consistent with the inability to promote endosome-to-plasma membrane recycling of GLUT1 not arising from a loss of endosomal localization. Furthermore, confocal analysis of the steady-state distribution of GLUT1 in GFP-SNX27(R498D) rescue cells confirmed that GLUT1 was retrieved from the lysosomal degradative fate, as shown by reduced co-localization with the LAMP1-labeled lysosome (Fig 5E). However, unlike the rescue with wild-type GFP-SNX27, where GLUT1 was enriched at the cell surface, the GLUT1 in the GFP-SNX27(R498D) rescue displayed an increased level of intracellular puncta and failed to repopulate the cell surface suggestive of the transporter being trapped in an endosomal compartment (Fig 5E), the precise nature of which will require further study. Together, these data are consistent with SNX27 orchestrating cargo retrieval from the degradative fate through its PDZ domain-mediated association with cargo and Retromer, while the SNX27 FERM domain mediates association with ESCPE-1 to promote tubular-based plasma membrane recycling of the retrieved cargo.

**ESCPE-1 association with SNX27 couples cargo retrieval with promotion of cell surface recycling**

To complete the mechanistic dissection of SNX27 coupling to ESCPE-1, we used a previously characterized double SNX1 and SNX2 knockout HeLa cell line [49] to generate lentiviral-transduced stable lines expressing GFP-SNX1 or GFP-SNX1(D45K,D87K) (Fig 6A). When expressed at near to endogenous levels each chimera displayed an indistinguishable association with EEA1 and SNX6-labeled endosomes (Fig 6B). Consistent with ESCPE-1 not being required for GLUT1 retrieval from lysosomal degradation, the rate of internalized GLUT1 degradation following 8 hours of treatment with cycloheximide was equivalent between...
Fig 5. SNX27:ESCPE-1 interaction is required for endosome-to-plasma membrane recycling of SNX27 cargos. (A) Representative western blot showing similar expression levels between transiently expressed GFP-SNX27 and GFP-SNX27(R498D). (B) Representative images of SNX27 KO HeLa cells transiently transfected with GFP-SNX27 or GFP-SNX27(R498D). Moreover, 48 hours after transfection, cells were fixed and immunostained for the endosomal markers EEA1 and SNX6. Magnified views of the white boxes are shown boxed. Cell numbers analyzed for colocalization were 60 GFP–SNX27 and 60 GFP–SNX27 R498D cells across n = 3 independent experiments. The Pearson coefficient values were compared using a 2-tailed unpaired t test; for EEA1 P = 0.2971, SNX6 P = 0.0506. (C) Degradation assay in parental HeLa cells or SNX27 KO HeLa cells that were transiently transfected with GFP-SNX27 or GFP-SNX27(R498D) mutant. Moreover, 48 hours after transfection, cells were incubated with 10 μg/ml cycloheximide and lysed at different time points as indicated. The band intensities of endogenous GLUT1 and
ITGA5 were measured from \( n = 3 \) independent experiments using Odyssey software. The levels of GLUT1 and ITGA5 at 8 hours were compared with the corresponding level at 0-hour time point. Analysis was done using a 2-way ANOVA and Sidak test. GLUT1: \( P = 0.0735 \) (HeLa), 0.0068 (KO), 0.9997 (+GFP-SNX27), 0.9975 (+GFP-R498D). ITGA5: \( P = 0.1151 \) (HeLa), 0.9820 (KO), 0.9999 (+GFP-SNX27), and 0.6899 (+GFP-R498D). (D) Analysis of surface levels of GLUT1 and ITGA5 in parental HeLa cells or SNX27 KO HeLa cells that were transiently transfected with GFP-SNX27 or GFP-SNX27(R498D) mutant. Moreover, 48 hours after transfection, cells were subjected to surface biotinylation followed by streptavidin-based immunosoluation. The immunosoluates were blotted for GLUT1, and the band intensities were measured from \( n = 4 \) independent experiments using Odyssey software and compared to the levels in parental HeLa using 1-way ANOVA and Dunnett test. GLUT1: \( P = 0.0134 \) (KO versus HeLa), 0.9587 (+GFP-SNX27 versus HeLa), and 0.0163 (+GFP-SNX27 R498D versus HeLa). The immunosoluates were blotted for ITGA5, and the band intensities were measured from \( n = 3 \) independent experiments using Odyssey software and compared to the levels in parental HeLa using 1-way ANOVA and Dunnett test. ITGA5: \( P = 0.5616 \) (KO versus HeLa), 0.0733 (+GFP-SNX27 versus HeLa), and 0.2354 (+GFP-SNX27 R498D versus HeLa). (E) Representative images of parental HeLa cells, SNX27 KO HeLa cells transiently transfected with GFP-SNX27 or GFP-SNX27(R498D). Moreover, 48 hours after transfection, cells were fixed and immunostained for the cargo GLUT1, the endosomal marker VPS35, and the late endosome/lysosome marker LAMP1. A total of 60 cells were quantified per condition in 4 micrographs. Molecular masses are given in kilodaltons. Bars, error bars, and symbols represent the mean, SEM, and individual data points, respectively. * \( P < 0.05 \), ** \( P < 0.01 \), *** \( P < 0.001 \), **** \( P < 0.0001 \), ns = not significant. The data underlying the graphs shown in the figure can be found in S1 Data. ESCPE-1, endosomal SNX-BAR sorting complex for promoting exit 1; KO, knockout; SNX6, sorting nexin-6; SNX27, sorting nexin-27; WT, wild-type.

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Evolutionary assembly of the SNX27:Retromer:ESCPE-1 sorting axis

We used phylogenetic analyses to investigate the evolution of the association of SNX27 with Retromer and with the ESCPE-1 complex. The inferred phylogeny of SNX27 suggests the gene was ancestrally present in filozoans (i.e., the group comprising filastereans, choanoflagellates, and metazoans) (S4A Fig). In SNX27, an exposed β-hairpin in the SNX27 PDZ domain, encoded by residues 67 to 79 in human SNX27, engages a groove in the VPS26A (and VPS26B) subunit of Retromer [14] (Fig 7A). An equivalent sequence, including key residues Leu-67 and Leu-74 of the β-hairpin, is present in SNX27 across all choanozoans (this is, the group comprising metazoans and choanoflagellates; Figs 7B and S4A). Key residues in VPS26A required for SNX27 binding include D44 and L154, equivalent residues in VPS26B being D42 and L152 [14]. The inferred phylogeny of VPS26A and VPS26B revealed that VPS26B is ancestrally present across eukaryotes, with a duplication of VPS26B in vertebrates leading to the emergence of VPS26A (Figs 7B and S4B). Interestingly, within VPS26B, the D44 residue is present ancestrally in filozoans, while the L154 residue evolved in choanozoans (Fig 7B). Together, these analyses suggest that the SNX27–Retromer association first appeared in the last common ancestor of choanoflagellates and metazoans.
Fig 6. ESCPE-1 interaction with SNX27 is required for endosome-to-plasma membrane recycling of GLUT1. (A) Representative western blot showing similar expression levels between stably transfected GFP-SNX1 and GFP-SNX1(KK) in SNX1/2KO cells. SNX1 band intensities were measured from 3 independent experiments and normalized to actin. Analysis was done using a 2-tailed unpaired t-test. (B) Representative images of SNX1/2 KO HeLa cell line stably transfected with GFP-SNX1 or GFP-SNX1(KK). Cells were fixed and immunostained for the endosomal markers EEA1 and SNX6. Magnified views of the white boxes are shown boxed. Cell numbers analyzed for colocalization were 50 GFP–SNX1 and 50 GFP–SNX1(KK) cells across 3 independent experiments. The Pearson coefficient values were compared using a 2-tailed unpaired t-test; for EEA1 P = 0.5971, SNX6 P = 0.6459. (C) Degradation assay in parental HeLa cells or SNX1/2 KO HeLa cells that were stably transfected with GFP-SNX1 or GFP-SNX1(KK) mutant. Cells were incubated with 10 μg/ml cycloheximide and lysed at different time points as indicated. The band intensities of endogenous GLUT1 and ITGA5 were measured from 3 independent experiments using Odyssey software. The levels of GLUT1 and ITGA5 at 8 hours were compared with the corresponding level at 0-hour time point. Analysis was done using a 2-way ANOVA and Sidak test. GLUT1: P = 0.6796 (HeLa).
The association between SNX27 and ESCPE-1 depends on the presence of binding motifs in SNX1 and SNX2 and the FERM domain of SNX27. SNX1 is present across eukaryotes; a gene duplication giving rise to SNX1 and SNX2 from a single ancestral protein occurred in the common ancestor of vertebrates (Figs 7B and S4C). Several motifs required in the FERM domain of H. sapiens SNX27 for binding to ESCPE-1 appear to have evolved in the common ancestor of metazoans or early during their radiation (Fig 7B); many of the key residues (R437, R498, and K501) were in place in the last metazoan common ancestor or prior to the divergence of cnidarians and bilaterians (K495 and K496). For SNX1, the first aDLF motif was likely already present in the last metazoan common ancestor. It is present in sequences from bilaterians and Porifera (sponges) but is absent in cnidarians, placazoans, and ctenophorans, which might represent secondary losses of the motif. As for the second aDLF motif, this is present in all vertebrates. This analysis therefore suggests that the SNX27:ESCPE-1 interaction evolved after the SNX27–Retromer link during early metazoan evolution.

Discussion

Here, in combining biochemical, structural, and cellular analyses, we have established that aDLF motifs in the redundant SNX1 and SNX2 subunits of ESCPE-1 directly associate with the FERM domain of SNX27 within the SNX27–Retromer assembly. Studies of endosomal cargo sorting show that this association is necessary to promote the ESCPE-1-dependent tubulovesicular cell surface recycling of internalized transmembrane proteins. Our data suggest that ESCPE-1-mediated recycling occurs subsequently or in conjunction with SNX27–Retromer-dependent retrieval from lysosomal degradation. Building on previous studies [12–14], these data collectively identify the SNX27:Retromer:ESCPE-1 assembly as a dynamic endosomal coat complex that couples sequence-dependent cargo recognition with membrane remodeling to facilitate the formation of cargo-enriched transport carriers for cargo recycling. In addition, we have begun to describe the functional diversification of Retromer as it has evolved from its tight association with the SNX-BAR homologues Vps5 and Vps17, as originally identified in yeast [50], to a more transient engagement of ESCPE-1 in metazoans and their relatives mediated at least in part through the SNX27 cargo adaptor.

At present, we propose the following model to describe the functional connectivity of the SNX27:Retromer:ESCPE-1 coat complex within the context of cargo retrieval and recycling. Newly endocytosed cargo containing a functional SNX27 PDZbm are recognized by SNX27, associated with the PI(3)P-enriched early endosome, and these interactions together can stabilize the endosomal residency of SNX27 [45]. As the endosome matures, SNX27 binds to...
Retromer through direct association of VPS26A (or VPS26B) to the SNX27 PDZ domain. These interactions are mutually stabilizing, thereby locking down cargo association [14,29]: This assembly is essential for cargo retrieval from the lysosomal degradative fate [13]. Retromer dimerization and the formation of Retromer arches [9,10,33] may aid SNX27 and cargo clustering, thereby forming the retrieval subdomain and initiating weak membrane curvature [51]. These events are supported through the Retromer-mediated recruitment of the WASH complex and the localized production of branched filamentous actin [34,35]. An additional accessory protein, ANKRD50, may serve to stabilize the formation of the SNX27:Retromer:WASH assembly [52]. To this emerging “bud,” ESCPE-1 is recruited through recognition of 3-phosphoinositides and membrane curvature [41,53,54] and the direct binding to SNX27 described in the present study. The timed increase in ESCPE-1 residency and localized concentration provides the handover mechanism to ensure that the captured and enriched cargo enter the forming tubular profile and the ensuing transport carrier. Continued localized WASH-mediated actin polymerization and the recruitment of motor proteins facilitates tubule maturation and fission [55–57] (Fig 8). Additional cargo entry may be achieved through direct recognition by ESCPE-1 and yet to be described mechanisms [18,19]. As with the assembly of other coat complexes [58], association of the SNX27:Retromer:ESCPE-1 coat is driven by avidity through a series of low affinity interactions that evoke the consideration of checkpoints for
monitoring the fidelity of pathway progression. It is important to state that this model is certainly an oversimplification and many key questions remain unanswered. Not least whether such a coat complex can be visualized through membrane-associated cryo-electron tomography of reconstituted SNX27:Retromer:ESCPE-1 perhaps in the presence of ANKRD50.

The importance of the amino-terminal extension of SNX1 and SNX2 has not previously been considered. While we and others [59–60] have independently established that they constitute primary sites of interaction with SNX27, it is tempting to speculate that they could orchestrate a broader set of protein–protein interactions. Interestingly, these unstructured amino termini may extend to over 30 nm in length and could allow the ESCPE-1 oligomeric assemblies to capture peripheral factors for the fine-tuned regulation of tubular maturation and scission. This would resemble the organization of the endocytic clathrin coat where several accessory proteins protrude from the core clathrin lattice into the cytoplasm to enable a regulated function of the endocytic coat during endocytic vesicle formation [61].

Other than mediating the recruitment of accessory proteins, there is some evidence that these amino-terminal extensions could play a role in the assembly and membrane targeting of SNX-BAR dimers. In fact, besides SNX1 and SNX2, SNX4, SNX7, SNX8, and SNX30, all possess unstructured amino-terminal extensions of different length and of unknown function. The low-complexity region of the SNX8 homologue Mvp1 appears to stabilize soluble Mvp1 in a cytosolic tetrameric form where the PX and BAR domains are masked and unable to target membranes [62]. The release of the amino terminus liberates functional dimers of Mvp1 facilitating their membrane association and function [62]. It will be interesting to explore whether a similar mechanism also regulates ESCPE-1 targeting to the endosomal network in mammalian cells during endosomal recycling.

An important aspect of cargo sorting by SNX27 is regulation by posttranslational modification. Phosphorylation of residues within the core carboxyl-terminal PDZbm negatively impairs SNX27 association, but phosphorylation of residues preceding the minimal PDZbm can dramatically enhance affinity for the SNX27 PDZ domain [29]. A well-characterized example of cargo tail phosphorylation modulating SNX27 affinity and subsequent endosomal sorting is the β2AR [29,63]. Furthermore, phosphorylation of SNX27 at Ser-51 alters the conformation of its PDZbm-binding pocket decreasing the affinity for cargo proteins, thereby inhibiting endosomal sorting [64] and Ser-302 and Ser-304 phosphorylation in VPS26B can modulate SNX27 association [65]. Similarly, ESCPE-1 function can be regulated by phosphorylation; in this case, Ser-226 of SNX5 undergoes phosphorylation, which decreases dimerization with SNX1 and SNX2 and reduces endosomal recycling [66]. Therefore, a fascinating question regards whether the SNX27:ESCPE-1 interaction may be orchestrated by posttranslational modification. We have here shown that a series of acidic residues upstream of the DLF motif in the SNX1 and SNX2 amino termini contribute to the binding to SNX27 (Fig 3F). Interestingly, several phosphorylation sites in the aDLF sequences of SNX1 have been annotated, including Ser-39, Ser-41, and Ser-72, as well as Thr-72 [67–69]. It is tempting to speculate that multiple phosphorylations of the ESCPE-1 amino termini may regulate its association with SNX27–Retromer and, in turn, dictate the timing and kinetics of the endosomal recycling route.

Finally, a fascinating question relates to the molecular evolution of Retromer [70]. In yeast, Retromer can form a stable pentameric assembly of 2 subcomplexes, the so-called “cargo-selective” Vps26:Vps35p:Vps29p heterotrimer and the membrane remodeling Vps3p and Vps17p heterodimer [50], the latter being considered analogous to ESCPE-1. Two functional processes, sequence-dependent cargo recognition and the biogenesis of tubulovesicular transport carriers, are therefore brought together in a single pentameric assembly [71]. In metazoans, the VPS26:VPS35:VPS29 Retromer and ESCPE-1 appear to no longer form a stable pentameric assembly [72]; rather, they have diverged into 2 separate
complexes and in so doing have expanded their functional roles [18,19,49,72,73]. Our phylogenetic analysis suggests that the SNX27:Retromer:ESCPE-1 assembly observed in metazoans evolved in a stepwise manner, with the SNX27–Retromer interaction evolving prior to the origin of metazoans, in the common ancestor of metazoans and choanoflagellates—the group of unicellular and colonial flagellates that are the closest living relatives of animals. The SNX27:ESCPE-1 interaction evolved later, during early metazoan evolution and likely prior to the divergence of cnidarians and bilaterians. Thus, within the SNX27:Retromer:ESCPE-1 pathway, the retrieval of cargo from the degradative fate appears to have evolved prior to the direct coupling to ESCPE-1 for promotion of tubulovesicular-based recycling to the cell surface. The divergence of the ancestral pentameric Retromer coat complex therefore appears to have occurred to accommodate the expanding complexity of the endomembrane system in metazoans [74]. The ability of metazoan Retromer to associate with SNX27 and the ability of ESCPE-1 to directly bind cargo through a helix–loop–helix region [18,19], features not observed in yeast, have greatly expanded the capacity for sequence-dependent endosomal retrieval and recycling in metazoans. Indeed, it is tempting to speculate that the increased plasticity of the SNX27:Retromer:ESCPE-1 axis, and the
existence of additional Retromer-independent pathways [1], has allowed the metazoan endosomal network to accommodate the evolving diversity and number of cargos undergoing sequence-dependent sorting, leading to an expansion in the number of recycling itineraries that cargos can engage to interface with distinct membrane compartments. Dissecting how these sorting complexes and pathways are integrated and regulated, for example, by posttranslational modifications, to shape sequence-dependent cargo sorting within the metazoan endosomal network will be a major challenge moving forward.

Overall, in defining the mechanistic basis of SNX27–Retromer association with ESCPE-1, we have provided new functional insight into the endosomal sorting of hundreds of internalized PDZbm-containing receptors, channels, transporter, enzymes, and adhesion molecules for repopulation of the cell surface [13]. This increased molecular understanding provides new insight into the role of sequence-dependent endosome sorting in cellular processes ranging from the establishment of polarity and the uptake of nutrients through to synaptic function and cancer cell metastasis in both health and disease [40].

**Material and methods**

**Antibodies**

Antibodies used in this study were mouse monoclonal antibodies GFP (clones 7.1 and 13.1; 11814460001; Roche, Germany) (1:2,000 for WB), LAMP1 (clone H4A3, AB2296838; DSHB, USA) (IF 1:200), SNX1 (clone 51/SNX1; 611482; BD, USA) (1:1,000 for WB, 1:200 for IF), SNX2 (clone 13/SNX2; 5345661; BD) (1:1,000 for WB, 1:200 for IF), SNX6 (clone d-5, 365965; Santa Cruz Biotechnology, USA) (1:1,000 for WB, 1:200 for IF), SNX27 (ab77799, Abcam, UK) (1:1,000 for WB, 1:200 for IF), β-actin (A1978; Sigma-Aldrich, USA) (1:2,000 for WB), ITGA5 (610633, BD) (1:1,000 for WB), VPS29 (clone d-1, 398874; Santa Cruz Biotechnology) (1:200 for WB); rabbit monoclonal antibodies: EEA1 (C45B10; Cell Signaling Technology, USA) (IF 1:200), VPS35 (EPR11501(B); 157220; Abcam) (1:1,000 for WB), GLUT1 (ab115730, Abcam) (1:1,000 for WB, 1:200 for IF); rabbit polyclonal antibodies: GFP (GTX20290; GeneTex, USA) (WB 1:2,000), VPS26A (23892; Abcam) (1:1,000 for WB), VPS35 (97545; Abcam) (1:1,000 for WB, 1:200 for IF), VPS29 (98929; Abcam) (1:100 for WB); and goat polyclonal antibodies to VPS35 (10099; Abcam) (1:200 for IF).

**Cell culture and transfection**

HeLa and HEK-293T cell lines were sourced from ATCC. Authentication was from the ATCC. Cells were grown in DMEM (Sigma-Aldrich) supplemented with 10% (v/v) FCS (Sigma-Aldrich) and penicillin/streptomycin (Gibco, USA) and grown under standard conditions. FuGENE HD (Promega, USA) was used for transient transfection of DNA according to the manufacturer’s instructions. Cycloheximide was used to prevent up-regulation of protein synthesis. In this case, cycloheximide (C7698; Sigma-Aldrich) at 10 μg/ml was added to the cells 48 hours after FuGENE transfection for the indicated time points. For the generation of CRISPR-Cas9 SNX27 knockout HeLa cells, HeLa cells were transfected with pX330 plasmid coding for the gRNA against the gene of interest together with a puromycin resistance-expressing plasmid. Cells were then subjected to puromycin selection for 24 hours. The pooled population of cells was subjected to lysis and western blotting to confirm the knockout. The VPS35 knockout clonal cell line used in this study was characterized previously [73]. For GFP-based immunoprecipitations, HEK293T cells were transfected with GFP constructs using polyethyleneimine (Sigma-Aldrich) and expression was allowed for 48 hours.
Immunoprecipitation and quantitative western blot analysis
For western blotting, cells were lysed in PBS with 1% (v/v) Triton X-100 and protease inhibitor cocktail. The protein concentration was determined with a BCA assay kit (Thermo Fisher Scientific, USA), and equal amounts were resolved on NuPAGE 4% to 12% precast gels (Invitrogen, USA). Blotting was performed onto polyvinylidene fluoride membranes (Immobilon-FL; EMD Millipore, USA) followed by detection using the Odyssey infrared scanning system (LI-COR Biosciences, USA). For GFP-based immunoprecipitations, HEK-293T cells were lysed 48 hours after transfection in immunoprecipitation buffer (50 mM Tris-HCl, 0.5% (v/v) NP-40, and Roche protease inhibitor cocktail) and subjected to GFP trap (ChromoTek, Germany). Immunoblotting was performed using standard procedures. Detection was performed on an Odyssey infrared scanning system (LI-COR Biosciences) using fluorescently labeled secondary antibodies. In using the Odyssey, we routinely performed western blot analysis where a single blot is simultaneously probed with antibodies against 2 proteins of interest (distinct antibody species) followed by visualization with the corresponding secondary antibodies conjugated to distinct spectral dyes.

Biotinylation of cell surface proteins
For surface biotinylation experiments, fresh Sulfo-NHS-SS Biotin (Thermo Fisher Scientific, #21217) was dissolved in ice-cold PBS at pH 7.8 at a final concentration of 0.2 mg/ml. Cells were washed twice in ice-cold PBS and placed on ice to slow down the endocytic pathway. Next, cells were incubated with the biotinylation reagent for 30 minutes at 4˚C followed by incubation in TBS for 10 minutes to quench the unbound biotin. The cells were then lysed in lysis buffer and subjected to Streptavidin beads-based affinity isolation (GE Healthcare, USA).

Immunofluorescence staining
Cells were fixed in 4% (v/v) PFA for 20 minutes and washed 3 times in PBS and permeabilized with 0.1% (v/v) Triton X-100. For LAMP1 staining, cells were instead permeabilized by submersion in liquid nitrogen for 20 seconds. Fixed cells were blocked in 1% (w/v) BSA and incubated in primary antibody and respective secondary antibody (Alexa Fluor; Thermo Fisher Scientific) in 1% (w/v) BSA.

Image acquisition and image analysis
Microscopy images were collected with a confocal laser-scanning microscope (SP5 AOBS; Leica Microsystems, Germany) attached to an inverted epifluorescence microscope (DMI6000; Thermo Fisher Scientific). A 63× 1.4 NA oil immersion objective (Plan Apochromat BL; Leica Biosystems, Germany) and the standard SP5 system acquisition software and detector were used. Images were captured at room temperature as z stacks with photomultiplier tube detectors with a photocathode made of gallium-arsenide-phosphide (Leica Microsystems) for collecting light emission. Images were captured using Application Suite AF software (version 2.7.3.9723; Leica Microsystems) and then analyzed with the Volocity 6.3 software (PerkinElmer, USA).

Statistics and reproducibility
All quantified western blot are the mean of at least 3 independent experiments. Statistical analyses were performed using Prism 7 (GraphPad Software, USA). Graphs represent means and SEM For all statistical tests, P < 0.05 was considered significant and is indicated by asterisks.
Molecular biology and cloning for recombinant protein production

For bacterial expression, full-length mouse SNX1 with an amino-terminal His-tag (mSNX1FL hereafter) was cloned into the pMW172Kan vector [75], and the truncated mSNX1 construct containing residues 1 to 139 was cloned into pGEX4T-2 vector for expression as a thrombin-cleavable amino-terminal GST fusion protein. DNA encoding full-length mouse SNX27 (mSNX27FL hereafter) was inserted into the pMCSG7 vector [76] by the ligation-independent cloning (LIC) approach. Human SNX27 constructs, hSNX27FL, hSNX27FERM, and hSNX27FL single-site mutants R437D, K495D, K496D, R498D, and K501D were cloned into the pET-28a vector by GenScript and codon optimized for bacterial protein expression. An additional TEV cleavage site was inserted between the amino-terminal His-tag and hSNX27 coding sequences.

Protein expression and purification

The DNA constructs of mSNX27 and mSNX1 were expressed using *Escherichia coli* BL21 CodonPlus (DE3) competent cells, while the hSNX27 constructs were expressed using *E. coli* BL21 (DE3) competent cells. The cells were grown in Ultra Yield flasks at 37˚C, 180 rpm until the cell density reached OD$_{600}$ approximately 0.8. Protein expression was then induced with 1 mM IPTG and cells incubated for 16 to 18 hours at 18˚C. For SNX27, 0.5 mM IPTG was added for induction to get optimal protein expression yield. To collect cell pellets, the cells were harvested by 10-minute centrifugation (JLA 8.1 rotor) at 6,000 rpm, 4˚C.

For SNX1 purification, the cell pellet was resuspended in lysis buffer containing 50 mM HEPES pH 7.5, 200 mM NaCl, 5% glycerol, 1 mM β-mercaptoethanol (β-ME), 1 mM benzamidine, and 10 μg/mL DNase I dissolved in PBS containing 10% glycerol. For SNX27 proteins, the cell pellet was resuspended in lysis buffer containing 20 mM Tris-Cl pH 8.0, 500 mM NaCl, 5% glycerol, 1 mM β-ME, 1% Triton X-100, 1 mM benzamidine, and 10 μg/mL DNase I dissolved in PBS containing 10% glycerol. Cells were lysed by high-pressure homogenization (35 Kpsi). The homogenate was centrifuged for 30 minutes at 4˚C using a JA 25.50 rotor with 19,000 rpm. The supernatant of His-tagged protein homogenates were loaded onto preequilibrated TALON metal affinity resin (Clontech, USA) or Ni-nitrilotriacetic acid (NTA) resin, while the supernatant of GST-fusion protein homogenates were loaded onto preequilibrated glutathione Sepharose beads (GE Healthcare) and incubated on an orbital rotator for an hour in cold room.

Affinity matrices with bound fusion proteins were loaded into empty glass chromatography columns (Bio-Rad, USA) washed with 100 ml of washing buffer containing either 50 mM HEPES pH 7.5, 200 mM NaCl, 5% glycerol, 1 mM β-ME or 20 mM Tris-Cl pH 8.0, 200 mM NaCl, 5% glycerol, 1 mM β-ME. For His-tagged protein purification, 10 mM and 300 mM imidazole was added to wash and elution buffers, respectively. To collect the GST-tagged mSNX1 constructs, 50 mM reduced glutathione in elution buffer was applied for elution. cOmplete EDTA-free protease inhibitor tablets were added to the eluted SNX1 samples to reduce proteolytic degradation of their disordered amino-terminal region.

The fractions containing eluted proteins were concentrated to approximately 5 ml by Amicon cc ultrafiltration devices (10 kDa or 30 kDa molecular mass cutoff) (Millipore) and centrifuged in a bench centrifuge for 10 minutes at maximum speed to remove any precipitated material before being subjected to size-exclusion chromatography (SEC). For SEC, the HiLoad 16/600 Superdex 200 (GE Healthcare) column was used with proteins eluted in SEC buffer containing 50 mM HEPES pH 7.5, 200 mM NaCl, 5% glycerol, 1 mM β-ME or 20 mM Tris-Cl pH 8.0, 100 mM NaCl, 5% glycerol, and 1 mM β-ME. To check the purity and integrity of the purified fractions, sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) was applied using Bolt 12% Bis-Tris gel with 1x MES running buffer at 190V for 25 minutes.
After the SDS-PAGE, gels were stained with Coomassie Brilliant Blue R250 solution for visualization. For hSNX27FERM, an additional anion exchange chromatography (IEX) was applied using the Mono Q anion exchange chromatography column preequilibrated with IEX buffer A: 20 mM Tris-HCl pH 8.0, 100 mM NaCl, 5% glycerol, 1 mM β-ME. A linear salt gradient from 0% to 40% IEX buffer B (20 mM Tris-HCl pH 8.0, 500 mM NaCl, 5% glycerol, 1 mM β-ME) was conducted. Proteins usually eluted in buffer containing 200 mM NaCl, and fractions collected were analyzed by SDS-PAGE.

**Isothermal titration calorimetry**

Human SNX1 and SNX2 peptide sequences were synthesized by Genscript (USA). To obtain the peptide stock solutions at neutral pH, 100 mM Tris-HCl, pH 10.6 was used to dissolve the peptide to a final concentration of 8 mM. The peptides were SNX1\textsubscript{35-51} EAGDSDEGEDIFTGAA, SNX1\textsubscript{135-145} (D45K) EAGDSDEGEKIFTGAA, SNX1\textsubscript{75-92} NGIHEEQQEPQDLFADA, SNX2\textsubscript{16-33} TDFEDLEDGEDLFTSTVS, SNX2\textsubscript{16-33} (DLF/SSS) TDFEDLEDGESSSTSTVS, and SNX2\textsubscript{62-82} TEVVLDDREDLFAEATEEV.

All microcalorimetry experiments were conducted at 25˚C using a PEAQ ITC (Malvern, UK) in 50 mM HEPES (pH 7.5), 200 mM NaCl, 5% glycerol, and 1 mM β-ME (ITC buffer). All proteins were purified by SEC using the same buffer prior to ITC experiments to avoid buffer mismatches. To measure binding of SNX27 to SNX1 proteins, 800 μM of hSNX27FERM was titrated into 20 μM of GST-mSNX1\textsubscript{1-139} or His-SNX1\textsubscript{FL}. For the binding of SNX1 and SNX2 peptides to SNX27, peptides were diluted 10-fold to 800 μM in ITC buffer peptides and titrated into 20 μM SNX27\textsubscript{FL} proteins containing the same 1:10 ratio of peptide buffer (100 mM Tris-HCl, pH 10.6). The competitive ITC experiments were carried out by titrating 800 μM SNX1\textsubscript{75-92} peptides into 20 μM proteins with 2-fold molar excess SNX1\textsubscript{16-33} peptide preincubated with SNX27 on ice for an hour. For ITC experiment using SNX1\textsubscript{75-92} against hSNX27 wild-type and single-site mutants, 600 μM SNX1\textsubscript{75-92} peptide was titrated into 20 μM hSNX27 proteins.

The ITC experiments were conducted with 1 injection of 0.4 μL followed by a series of 12 injections of 3.2 μL each with 180-second intervals and a stirring speed of 850 rpm. The heat exchange of interactions was obtained through integrating the observed peaks and background correction by subtracting the heats of dilution. Data were analyzed with Malvern software package by fitting and normalized data to a single-site binding model, yielding the thermodynamic parameters $K_d$, $\Delta H$, $\Delta G$, and $-T\Delta S$ for all binding experiments. The stoichiometry was refined initially, and if the value was close to 1, then N was set to exactly 1.0 for calculation. All experiments were performed at least twice to guarantee reproducibility of the data.

**AlphaFold predictions of peptide binding to SNX27**

To generate predicted models of SNX1 and SNX2 amino-terminal peptides associating with the SNX27 FERM domain, we used the AlphaFold2 neural network [47] implemented within the freely accessible ColabFold pipeline [48]. For each peptide-modeling experiment, the sequence of the human SNX27a FERM domain (residues 269 to 531; Q96L92) was modeled with the 4 separate amino-terminal aDLF sequences from SNX1 and SNX2. The specific peptides we modeled were residues 33 to 56 and 75 to 98 from human SNX1 (Q13596) and residues 14 to 37 and 60 to 83 from human SNX2 (O60749). ColabFold was executed using default settings where multiple sequence alignments were generated with MMseqs2 [77], and structural relaxation of final peptide geometry was performed with Amber [78] to generate 3 models per peptide. Sequence conservation was mapped onto the modeled SNX27 FERM domain structure with Consurf [79]. Structural alignments and images were generated with Pymol (Schrodinger, USA).
Phylogenetic analyses

Homologous proteins were identified using NCBI protein–protein BLAST (with default parameters) against all non-redundant GenBank CDS Translations+PDB+SwissProt+PIR+PRF excluding environmental samples from WGS projects for a wider range of species (unaligned sequence data with accession numbers available in S4 Fig). For the ctenophore data, we used the BLAST (with default parameters) from the Mnemiopsis Genome Project Portal (MGP) portal [80–82]. Sequences were aligned using MAFFT v7.480 [83], L-INS-I mode, and then used to infer initial exploratory trees to identify specific duplications (using IQtree 2.1.4 [84] with the LG+F+G model and 1,000 ultrafast bootstraps). After orthologues of each copy of the gene were identified in a range of taxa, we then performed iterative tree inferences to confirm paralogues and to identify species-specific duplication events. Subsequent maximum likelihood phylogenetic analyses were performed using IQtree 2.1.4 with the best fitting model selected using the Bayesian information criterion (BIC), including models which account for site-rate heterogeneity (LG+C10...C60) [85], empirical amino acid frequencies (+F), and accounting for across-site rate variation using either a Gamma distribution (+G) [86] or the free rates model (+R) [87,88] with 10,000 ultrafast bootstrap replicates. The best fitting model according to BIC was LG+C30+F+G for the final SNX1 and SNX2 tree (see S4 Fig) and LG+C20+F+G for SNX27 and VPS26 (alignments and maximum likelihood trees in S4 Fig).

Supporting information

S1 Fig. Western blot showing the levels of GLUT1 in parental HeLa, VPS35 KO, and SNX27KO not transfected or transiently transfected with GFP-SNX27 WT, GFP-SNX27 Δ67–77, and GFP-SNX27 ΔFERM. The blot is representative of 3 independent assays. Molecular masses are given in kilodaltons. Bars, error bars, and symbols represent the mean, SEM, and individual data points, respectively. *P < 0.05, **P < 0.01, ***P < 0.001, ****P < 0.0001, ns = not significant. The data underlying the graphs shown in the figure can be found in S1 Data. KO, knockout; SNX27, sorting nexin-27; WT, wild-type. (TIF)

S2 Fig. (A) Co-immunoprecipitation of GFP-tagged extended amino-terminal regions (NT) of SNX1 and SNX2 alongside the GFP-tagged short amino-terminal regions of SNX5 and SNX6 expressed in HEK293T cells. The cell lysates were subjected to GFP trap–based immunoprecipitation and the immunoprecipitates were blotted for SNX27 and GFP. The blot is representative of 3 independent GFP traps. (B) Co-immunoprecipitation of GFP-tagged SNX2 WT or SNX2 KK (D27K, D72K) expressed in HEK293T cells. The cell lysates were subjected to GFP trap–based immunoprecipitation and the immunoprecipitates were blotted for SNX27, SNX6, and GFP. The blot is representative of 4 independent GFP traps. (C) Competitive ITC assay of SNX216–33 and mSNX27 FL preincubated with 2-fold molar excess of SNX175–92 peptide showing that binding is blocked by the competing SNX175–92 peptide. Molecular masses are given in kilodaltons. Bars, error bars, and symbols represent the mean, SEM, and individual data points, respectively. *P < 0.05, **P < 0.01, ***P < 0.001, ****P < 0.0001, ns = not significant. The data underlying the graphs shown in the figure can be found in S1 Data. SNX1, sorting nexin-1; SNX5, sorting nexin-5; SNX6, sorting nexin-6; SNX27, sorting nexin-27; WT, wild-type. (TIF)

S3 Fig. (A) SNX27 surface colored according to sequence conservation. The peptides are predicted to bind to a highly conserved pocket in SNX27. (B, C) As described in methods, 12 models of SNX27 were generated in total, consisting of 3 models each of SNX27 in association
with the 4 aDLF sequences from human SNX1 and SNX2. B shows the overlay of all 12 models in Ca trace representation colored by the pLDDT score. The pLDDT is a per-residue confidence score between 0 (lowest confidence) and 100 (highest confidence). The majority of the SNX27 structures show very high pLDDT scores and overlay with very high precision. C shows only the best 8 models where the core aDLF sequences have identical binding conformations. aDLF, acidic-Asp-Leu-Phe; pLDDT, predicted Local Distance Difference Test; SNX1, sorting nexin-1; SNX2, sorting nexin-2; SNX27, sorting nexin-27.

(TIF)

S4 Fig. Maximum likelihood phylogenies of (A) SNX27, (B) VPS26A/B and (C) VPS5/SNX1/SNX2. (A) SNX27 (green) with SNX17 (red) used as an outgroup with representatives across Filozoa. Amino acids aligned with Homo sapiens in positions 437, 495, 496, 498, and 501 are written next to the species name. (B) VPS26A, VPS26B, and VPS26C, with VPS26C in red and VPS26B in green; the duplication event occurring in the ancestral vertebrate is in blue. Interestingly, we also find another independent duplication of VPS26B in plants evidenced by Arabidopsis thaliana and Quercus suber in this tree. (C) The SNX1 sequences across choanozoans, with the duplication in vertebrates in blue. The number of aDLF motifs in the amino terminus is written next to the species name. Phylogenies were inferred under the best-fit model as determined by BIC. Branch support values are percentages based on 10,000 ultrafast bootstraps. aDLF, acidic-Asp-Leu-Phe; BIC, Bayesian information criterion; SNX1, sorting nexin-1; SNX27, sorting nexin-27.

(TIF)

S1 Table. Summary of ITC binding parameters. ITC, isothermal titration calorimetry.

(PDF)

S1 Raw Images. Unprocessed images for blots shown in the paper.

(PDF)

S1 Data. Numerical data for Figs 1B, 1C, 2C, 3A, 3C, 3F, 4D, 4G, 5A, 5B, 5C, 5D, 5E, 6A, 6B, 6C, 6D and 6E and S1A and S2B.

(XLSX)

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**References**


