Polarized sorting of the copper transporter ATP7B in neurons mediated by recognition of a dileucine signal by AP-1

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ABSTRACT Neurons are highly polarized cells having distinct somatodendritic and axonal domains. Here we report that polarized sorting of the Cu²⁺ transporter ATP7B and the vesicle-SNARE VAMP4 to the somatodendritic domain of rat hippocampal neurons is mediated by recognition of dileucine-based signals in the cytosolic domains of the proteins by the σ1 subunit of the clathrin adaptor AP-1. Under basal Cu²⁺ conditions, ATP7B was localized to the trans-Golgi network (TGN) and the plasma membrane of the soma and dendrites but not the axon. Mutation of a dileucine-based signal in ATP7B or overexpression of a dominant-negative σ1 mutant resulted in nonpolarized distribution of ATP7B between the somatodendritic and axonal domains. Furthermore, addition of high Cu²⁺ concentrations, previously shown to reduce ATP7B incorporation into AP-1–containing clathrin-coated vesicles, caused loss of TGN localization and somatodendritic polarity of ATP7B. These findings support the notion of AP-1 as an effector of polarized sorting in neurons and suggest that altered polarity of ATP7B in polarized cell types might contribute to abnormal copper metabolism in the MEDNIK syndrome, a neurocutaneous disorder caused by mutations in the σ1A subunit isoform of AP-1.

INTRODUCTION Neurons are highly polarized cells having axonal and somatodendritic domains with distinct protein compositions. The steady-state localization of transmembrane proteins to each of these domains depends on their biosynthetic sorting into axonal and somatodendritic transport carriers, subsequently complemented by selective retention and retrieval mechanisms (Lasiecka and Winckler, 2011; Bonifacino, 2014). Somatodendritic sorting is often mediated by discrete signals contained within the cytosolic tails of the transmembrane proteins (West et al., 1997; Jareb and Banker, 1998; Poyatos et al., 2000; Rivera et al., 2003; Farias et al., 2012; Mattera et al., 2014). A subset of somatodendritic sorting signals fitting the tyrosine-based YXXØ motif (where Y is tyrosine, X is any amino acid, and Ø is a bulky hydrophobic amino acid) has been recently shown to interact with the μ1A subunit of the heterotetrameric adaptor protein 1 (AP-1) complex (Farias et al., 2012; Mattera et al., 2014; Figure 1A), a component of clathrin coats associated with the trans-Golgi network (TGN) and recycling endosomes (Robinson, 2004; Traub and Bonifacino, 2013). Mutation of the YXXØ signal or inactivation of AP-1 abrogates somatodendritic sorting of the corresponding proteins, resulting in their nonpolarized distribution in rat hippocampal neurons (Farias et al., 2012; Mattera et al., 2014). This role of AP-1 appears phylogenetically conserved, as mutation of the μ1-subunit orthologue UNC-101 in Caenorhabditis elegans abolishes the somatodendritic polarity of various receptors and transporters (Dwyer et al., 2001; Bae et al., 2006; Margeta et al., 2009). AP-1 also binds, via its γ and σ1 subunits, a subset of dileucine-based signals fitting the canonical motif [DE]XXXL[LI] (Janvier et al., 2003; Doray et al., 2007; Mattera 2011), but it remains to be determined whether these interactions also mediate somatodendritic sorting.
We decided to examine whether [DE]XXX[L][I]–AP-1 interactions participate in polarized sorting in neurons, with a focus on the mammalian Cu\(^{2+}\)-transporting P-type ATPases ATP7A and ATP7B. These transporters are closely related (~60% amino acid identity), multispanning membrane proteins that deliver Cu\(^{2+}\) from the cytosol to the lumen of the secretory pathway or the extracellular space (Wang et al., 2011; Polishchuk and Lutsenko, 2013). This transport is required for the loading of copper into copperoenzymes and the efflux of excess copper from the cells. ATP7A is expressed in most tissues, whereas ATP7B has a limited pattern of expression, with highest levels in the liver and lower levels in the brain and kidney. Mutations in ATP7A cause Menkes disease (Online Mendelian Inheritance in Man [OMIM] 309400), an X-linked recessive disorder characterized by general copper deficiency (Kaler, 2013). Mutations in ATP7B, on the other hand, cause Wilson disease (OMIM 277900), an autosomal recessive disorder leading to copper accumulation in the liver and brain (Kaler, 2013). Under basal Cu\(^{2+}\) conditions, both transporters exhibit steady-state localization to the TGN (Yamaguchi et al., 1996; Hung et al., 1997). In the presence of high Cu\(^{2+}\) levels, however, the transporters redistribute to the plasma membrane and peripheral cytoplasmic vesicles (Petris et al., 1996; La Fontaine et al., 1998; Roelofsen et al., 2000; Guo et al., 2005; Cater et al., 2006; Polishchuk et al., 2014). Subsequent lowering of Cu\(^{2+}\) levels causes the transporters to return to the TGN. This retrieval is dependent on dileucine residues fitting the [DE]XXX[L][I] motif in the C-terminal tails of the transporters (Francis et al., 1998; Petris et al., 1998; Roelofsen et al., 2000; Guo et al., 2005; Cater et al., 2006; Polishchuk et al., 2014). It remains to be determined, however, whether the [DE]XXX[L][I] motifs in these transporters function through direct interaction with AP-1. Moreover, despite the neuropathological

**FIGURE 1:** Binding of AP-1 to cargo proteins. (A) Schematic representation of the active, membrane-bound conformation of the heterotetrameric (γβ1-μ1-σ1) AP-1 complex indicating the core, hinge, and ear domains. The γ, μ1, and σ1 subunits occur as two or three isoforms encoded by different genes. YXXØ signals bind to a site on the μ1 subunit, whereas [DE]XXX[L][I] signals bind to a site on the γ-σ1 hemicomplex. (B) SDS–PAGE and Coomassie blue staining of recombinant GST-6His-tagged AP-1 core and 6His-tagged Arf1 used in vitro pull-down experiments. The AP-1 core used in these experiments was composed of the N-terminal, “trunk” domains of γ and β1 and the full-length μ1A and σ1C subunits (Ren et al., 2013). (C) Amino acid sequences of portions of the indicated proteins, with YXXØ and [DE]XXX[L][I] signals highlighted in red. For use in pull-down experiments, an N-terminal maltose-binding protein moiety was fused to the mouse VAMP4 cytosolic domain (residues 1–118), the entire HIV-1 Nef (NL4-3 variant), residues 324–353 of the cytosolic tail of rat TGN38, and the C-terminal cytosolic tails of human ATP7A (residues 1405–1500) and ATP7B (residues 1371–1465). (D) Pull down of recombinant GST-His-tagged AP-1 core by MBP-tagged wild-type (WT) or LL/AA (in the [DE]XXX[L][I] signal) or Y/A (in the YXXØ signal) mutant proteins in the absence or presence of GTP-locked Arf1Δ1-16-Q71L. MBP-tagged proteins were immobilized on amylase resin, and bound AP-1 core was analyzed by SDS–PAGE and immunoblotting with antibody to GST. MBP (~) was used as a negative control. The positions of molecular mass markers (in kilodaltons) are indicated on the left.
manifestations of both Menkes disease and Wilson disease, the role of the [DE][XXX][LI] motifs and the AP-1 adaptor in sorting of the transporters in neurons has not been studied.

In the present study, we tested for interactions of AP-1 with the ATP7A and ATP7B C-terminal tails using a recently developed in vitro pull-down assay involving conformational activation of AP-1 by the Arn1 GTPase (Guo et al., 2013; Ren et al., 2013). We detected weak binding of AP-1 to ATP7A, but strong, [DE][XXX][LI]-dependent binding of AP-1 to ATP7B. Analysis of the localization of ATP7B in rat hippocampal neurons in primary culture showed that this protein was mainly localized to the TGN, as well as to the plasma membrane of both the soma and dendrites. Remarkably, the protein was largely excluded from the axon, indicating that its distribution in neurons is polarized. Addition of high Cu2+ concentrations, previously shown to reduce ATP7B incorporation into AP-1–containing clathrin-coated vesicles (Hirst et al., 2012), caused loss of both TGN localization and somatodendritic polarity of ATP7B. Furthermore, mutation of the [DE][XXX][LI] motif or overexpression of α1-subunit mutants incapable of binding [DE][XXX][LI] signals abrogated the somatodendritic polarity of ATP7B. Similar manipulations also caused loss of somatodendritic polarity of another [DE][XXX][LI]-containing protein, the vesicle–soluble N-ethylmaleimide–sensitive factor attachment protein receptor (v-SNARE) VAMP4. These observations demonstrated that interactions of canonical [DE][XXX][LI] signals with the α1 subunit of AP-1 play a general role in protein sorting to the somatodendritic domain. Our findings provide new information relevant to the regulation of brain copper homeostasis under physiological and pathological conditions. In addition, they buttress the proposed explanation for copper metabolism abnormalities in the neurocutaneous disorder MEDNIK syndrome, which is caused by mutation of the α1A subunit isofrom (Montpetit et al., 2008; Martinelli et al., 2013).

RESULTS

Dileucine-dependent binding of the C-terminal tail of ATP7B to AP-1

To search for novel interactions of [DE][XXX][LI] with AP-1 (Figure 1A), we used a recently developed pull-down assay based on the incubation of maltose-binding protein (MBP)–fusion proteins having putative signals with glutathione S-transferase (GST)–hexahistidine (6His)–tagged AP-1 core in the presence of the 6His-tagged, GTP-locked form of Arn1 (Arn1Δ-Q71L), all recombinitely produced in Escherichia coli (Figure 1B and Supplemental Figure S1; Guo et al., 2013; Ren et al., 2013). Addition of this Arn1 construct induced a conformational activation of the AP-1 core, greatly increasing the sensitivity of the assay (Guo et al., 2013; Ren et al., 2013). The MBP–fusion proteins with bound AP-1 core were collected on amylose beads and analyzed by SDS–PAGE and immunoblotting with antibody to GST. As controls, we showed that the cytosolic domain of the vesicle-SNARE VAMP4 (Peden et al., 2001; Ren et al., 2013) and the Nef protein of HIV-1 (Janvier et al., 2003; Doray et al., 2007; Mattera et al., 2011), both having well-validated [DE][XXX][LI] signals (Figure 1C), interacted with AP-1 in a dileucine- and Arn1-dependent manner (Figure 1D). Similarly, the cytosolic tail of the TGN protein TGN38, which has a YXXØ signal (Humphrey et al., 1993; Ohno et al., 1995; Figure 1C), also interacted with AP-1 in a tyrosine- and Arn1-dependent manner (Figure 1D). We then used this assay to test for AP-1 binding to the C-terminal cytosolic tails of the Cu2+ transporters ATP7A and ATP7B, both having putative [DE][XXX][LI] signals (DKHSSL and DKWSSL, respectively; Figure 1C). We found that the ATP7A tail bound AP-1 weakly under all the conditions tested (Figure 1D). In contrast, the ATP7B tail exhibited strong, Arn1-dependent interaction with AP-1 (Figure 1D). This interaction was largely abrogated by mutation of Leu-1454 and Leu-1455 within the [DE][XXX][LI] motif (Figure 1D). On the basis of these results, we focused our subsequent experiments on the role of [DE][XXX][LI]–AP-1 interactions in neuronal trafficking of ATP7B.

Cu2+-regulated, dileucine-dependent localization of ATP7B to the neuronal TGN

ATP7B is mainly expressed in the liver but has also been detected in select areas of the brain, including the developing CNS (Kuo et al., 1997), the hippocampus (Saito et al., 1999), and the cerebellum (Barnes et al., 2005; see also Allen Brain Atlas, www.brain-map.org/). The neurological and psychiatric symptoms of Wilson disease patients are likely secondary to defective copper metabolism in the liver but could be compounded by primary alterations of copper metabolism in brain patients (Wang et al., 2011). The intracellular localization and traffic of ATP7B in neurons, however, have not been previously studied. To address this issue, we expressed green fluorescent protein (GFP)-tagged ATP7B by transfection into rat hippocampal neurons in primary culture. The localization of ATP7B was examined by confocal fluorescence microscopy in comparison to organelar markers. We observed that, under normal culture conditions, ATP7B largely colocalized with the TGN marker TGN38 in the neuronal soma, although some was also present at the plasma membrane (Figure 2A). Addition of the Cu2+ chelator bathocuproine disulfonate (BCS) for either 2 h (Figure 2B) or overnight (Supplemental Figure S2) had no effect on this localization. In contrast, incubation with high Cu2+ concentrations (200 μM) for 2 h caused redistribution of ATP7B from the TGN to the plasma membrane and peripheral vesicles (Figure 2C). Thus the intracellular localization of ATP7B in neurons is subject to regulation by Cu2+ levels, as previously shown in other cell types (Roelofsen et al., 2000; Guo et al., 2005; Cater et al., 2006). Of importance, mutation of the dileucine residues Leu-1454 and Leu-1455 in the [DE][XXX][LI] motif (Figure 1C) caused a similar redistribution of ATP7B to the cell periphery (Figure 2D), consistent with a function of this motif in determining TGN localization (Francis et al., 1998; Petris et al., 1998; Cater et al., 2006; Braiterman et al., 2011; Lalioti et al., 2014).

Somatodendritic polarity of ATP7B depends on a dileucine motif

Inspection of the neurons expressing GFP-tagged ATP7B using higher gain settings in the confocal microscope revealed that this protein was restricted to the soma and dendrites (marked by MAP2 staining) and largely excluded from the axon (marked by staining of the axon initial segment for ankyrin G) under normal culture conditions (Figure 3A). Mutation of the dileucine residues in the [DE][XXX][LI] motif resulted in loss of somatodendritic polarity, as evidenced by the appearance of ATP7B in the axons (Figure 3B). Quantification of fluorescence intensities in dendrites and axons from many cells yielded a dendrite/axon “polarity index” of 9.9 ± 2.6 for the normal protein and 1.3 ± 0.4 for the dileucine mutant (Figure 3C). From these experiments, we concluded that the [DE][XXX][LI] motif mediates not only TGN localization (Figure 2) but also somatodendritic polarity of ATP7B (Figure 3).

Characterization of α1 subunit mutants for use as dominant-negative constructs

The atomic structure of the AP-1 core in complex with a [DE][XXX][LI] signal has not yet been solved by x-ray crystallography. However, structural analyses of the unliganded AP-1 core (Heldwein et al., 2004; Ren et al., 2013) and the homologous AP-2 core in complex
endogenous AP-1. Coprecipitation with the endogenous $\gamma$ subunit of AP-1 showed that the A63W, V98S, Y62S, and R155S mutants of $\sigma$1A were all incorporated into AP-1 as efficiently as the wild-type protein (Figure 4C). In contrast, the I103S, E100Y, and V88D $\sigma$1A mutants were not incorporated (Figure 4C), probably because of conformational alterations induced by the mutations. The V98S mutants of $\sigma$1B and $\sigma$1C were also assembled into AP-1 (Figure 4C). Immunofluorescence microscopy of transfected neurons showed that the wild-type and V98S mutant forms of $\sigma$1A, $\sigma$1B, and $\sigma$1C in neurons all exhibited a similar pattern of localization to the TGN (Figure 4D; see also Supplemental Figure S3). ATP7B also colocalized with $\sigma$1B, as well as with the endogenous $\gamma$ subunit at the TGN (Supplemental Figure S4). The V98S $\sigma$1A mutants thus fulfilled the requirements for use in dominant-negative interference experiments.

**Interaction with AP-1 is critical for somatodendritic polarity of ATP7B and VAMP4**

To determine whether interaction with $\sigma$1 is required for somatodendritic sorting of [DE]XXL[LI]-containing proteins, we coexpressed GFP-tagged ATP7B with wild-type or V98S forms of myc- or HA-tagged $\sigma$1A, $\sigma$1B, and $\sigma$1C in hippocampal neurons. We observed that whereas expression of wild-type $\sigma$1A, $\sigma$1B, or $\sigma$1C had no effect on the somatodendritic polarity of ATP7B, expression of the corresponding V98S mutants resulted in nonpolarized distribution of the transporter (Figure 5, A–D). To generalize these findings, we extended our analyses to VAMP4 (Figure 6, A–E). We observed that VAMP4 also exhibited somatodendritic polarity (Figure 6A), which was lost upon mutation of the dileucine motif (Figure 6B). Similar to ATP7B, VAMP4 somatodendritic polarity was not affected by overexpression of wild-type $\sigma$1B (Figure 6C) but was abrogated by overexpression of the V98S $\sigma$1B mutant (Figure 6D). In contrast to these findings, overexpression of the V98S $\sigma$1B mutant had no effect on the somatodendritic sorting of the YXXØ-containing transferrin receptor (TFR; Figure 7, A–C), consistent with the recognition of the YXXØ signal by $\mu$1 and not $\sigma$1 (Ohno et al., 1995). Thus specific recognition of [DE]XXL[LI] signals by the $\sigma$1 subunit of AP-1 mediates sorting of a subset of transmembrane proteins, including ATP7B and VAMP4, to the neuronal somatodendritic domain.

**High Cu$^{2+}$ concentration abrogates the somatodendritic polarity of ATP7B**

Live-cell imaging of neurons expressing GFP-ATP7B under basal conditions showed very few GFP-ATP7B–containing particles entering the axon, indicating that this protein was excluded from the axon at the level of the soma (Figure 8A and Supplementary Video S1). Incubation with 200 μM Cu$^{2+}$ for 2 h proved deleterious for the morphology of the neurons under the conditions of live-cell imaging. A milder treatment with 100 μM Cu$^{2+}$ for 1 h, however,
preserved the morphology of the neurons and revealed movement of a large number of GFP-ATP7B–containing particles from the soma into the axon (Figure 8A and Supplementary Video S1). Quantification of fluorescence intensities in many cells yielded a dendrite/axon polarity index of 8.7 ± 1.8 under basal conditions and 2.5 ± 0.7 upon treatment with 100 μM Cu²⁺ (Figure 8B). Kymographic analysis of the live-cell imaging data showed increased numbers of both anterograde (lines with negative slopes) and retrograde (positive slopes) carriers, as well as stationary foci (vertical lines), in the axons of neurons treated with 100 μM Cu²⁺ (Figure 8C). A recent study showed that elevated Cu²⁺ levels cause movement of ATP7B from the TGN to late endosomes and lysosomes in a hepatoma cell line (Polishchuk et al., 2014). Indeed, we observed that many of the GFP-ATP7B–containing particles that moved into the axon upon treatment with 100 μM Cu²⁺ also contained the late endosomal/lysosomal marker Lamp-1-RFP (Figure 8D). From these experiments, we concluded that elevated Cu²⁺ levels interfere with AP-1–dependent TGN/somatodendritic localization of ATP7B, resulting in its transport to late endosomes or lysosomes capable of moving into the axon.

**DISCUSSION**

The results presented here demonstrate that interactions of [DE]XX[L]L-[I]-type dileucine signals with the α1 subunit of AP-1 underlie the sorting of a subset of transmembrane proteins (e.g., ATP7B, VAMP4) to the somatodendritic domain of neurons. Together with the previous demonstration that interactions of YXXO-type signals with the μ1A subunit of AP-1 mediate somatodendritic sorting of other transmembrane proteins (Farias et al., 2012; Mattera et al., 2014), our findings support a general role for AP-1 in polarized sorting in neurons. Disruption of signal–AP-1 interactions results in nonpolarized distribution of the proteins to both dendrites and axons, indicating that these interactions normally function to exclude somatodendritic proteins from the axon. The exact mechanism by which AP-1 performs this exclusion remains to be elucidated. In the case of ATP7B, somatodendritic sorting correlates with concentration of this protein at the TGN, suggesting that axonal exclusion requires AP-1–dependent localization or retrieval to the TGN. Presumably AP-1 acts by segregating somatodendritic proteins away from TGN or recycling endosomal sites that give rise to axonal carriers.

In addition to shedding light on a general mechanism of polarized sorting, our findings reveal new aspects of the trafficking of a physiologically important protein, ATP7B. Previous studies of ATP7B traffic had been done in liver and epithelial cells but not in neurons. We find that ATP7B predominantly localizes to the TGN under basal conditions. We also observe localization of ATP7B to the plasma membrane of the soma and dendrites but not the axon. This distribution is consistent with cycling between the TGN and the somatodendritic plasma membrane under low Cu²⁺ levels. The somatodendritic restriction of ATP7B likely ensures efficient copper loading of cuproenzymes, such as peptidylglycine α-amidating monooxygenase (PAM; also an AP-1-interacting cargo; Bonnemaison et al., 2014), in the perikaryon, as well as protection from copper toxicity in the axon, as suggested by the deleterious effects of some missense mutations of the related ATP7A in distal motor neuropathies (Kennerson et al., 2010; Yi et al., 2012). High Cu²⁺ levels, however, cause ATP7B redistribution from the TGN to cytoplasmic vesicles that are capable of moving into the axon. Some of these vesicles also contain Lamp-1, identifying them as late endosomal/lysosomal compartments. These observations are in line with the reported high-Cu²⁺–induced relocalization of ATP7B to late endosomes and lysosomes and eventually to the bile canalicular surface in a hepatoma cell line (Polishchuk et al., 2014) and with the ability of lysosomes to undergo axonal transport (Mar et al., 2014; Otero et al., 2014). It thus appears that the mechanisms that regulate transport of ATP7B are conserved in different polarized cell types. Because somatodendritic sorting of ATP7B depends on [DE]XX[L]L-[I]–AP-1 interactions, it would now be of interest to study whether these interactions also mediate polarized sorting of ATP7B in hepatocytes, where this protein is most highly expressed.

It remains to be determined to what extent the neurological manifestations of Wilson disease are an indirect consequence of copper metabolism dysregulation in the liver or a direct result from its alteration in the brain (Emre et al., 2001; Schumacher et al., 2001). In any event, our findings provide important new information on the neuronal trafficking of a key regulator of iron metabolism that may be applicable to its trafficking in other polarized cell types, including hepatocytes. Our findings might also contribute to explaining the pathogenesis of the genetic disease MEDNIK syndrome, which is
Sorting of ATP7B in neurons
caused by mutations in the AP1S1 gene encoding the α1A subunit of AP-1 (Montpetit et al., 2008). MEDNIK is an acronym for the main symptoms of this disease: mental retardation, enteropathy, deafness, neuropathy, ichthyosis, and keratoderma. Of note, these manifestations all involve organs with a core component of polarized cells, including neurons and epithelial cells. Given the function of AP-1 in sorting to the somatodendritic domain of neurons (Farias et al., 2012; Mattera et al., 2014) and the basolateral domain of epithelial cells (Folsch et al., 1999; Carvajal-Gonzalez et al., 2012; Gravotta et al., 2012), it is reasonable to hypothesize that defects in polarized sorting of specific cargo proteins might contribute to the symptoms of MEDNIK syndrome. Why does a defect in only one α1 subunit isoform, α1A, cause this condition? The α1A, α1B, and α1C isoforms recognize overlapping but distinct sets of [DE]XX[L] signals (Mattera et al., 2011). It is thus likely that MEDNIK syndrome results from mis-sorting of a α1A-specific cargo in cells that otherwise express α1B and α1C. Alternatively, each α1 isoform could be expressed in a distinct set of cells, such that loss of α1A would cause a general failure of AP-1–dependent sorting in the corresponding cell population.

MEDNIK patients have been recently reported to have copper metabolism defects characteristic of both Menkes and Wilson diseases, pointing to ATP7A and ATP7B as likely candidates for missorting in this disease (Martinelli et al., 2013). Indeed, in fibroblasts from MEDNIK patients, ATP7A is not localized to the TGN but to peripheral structures irrespective of Cu²⁺ levels (Martinelli et al., 2013). We did detect specific interaction of the ATP7A tail with the AP-1 core (Figure 1D), but this interaction was too weak for detailed characterization. The weakness of this interaction could be explained by the fact that the AP-1 core construct we used in our studies had α1C instead of α1A. Although interaction of this core construct with ATP7B was stronger, it is possible that interaction with a α1A-containing core could have been even stronger. Our studies suggest that in addition to failure to localize to the TGN at low Cu²⁺ levels, altered polarity of ATP7B (and perhaps also ATP7A) in polarized cell types might underlie some of the copper metabolism defects in MEDNIK syndrome. Further studies on the signal-recognition preference and cell-specific expression of the different α1 subunit isoforms should help elucidate the mechanistic basis for the copper metabolism defects and other symptoms in MEDNIK syndrome.

**FIGURE 4:** Mutational analysis of α1 residues involved in recognition of dileucine motifs. (A) Surface representation of the predicted binding site for dileucine signals on the γ-α1A hemichannel based on the crystal structures of the unliganded AP-1 core (Heldwein et al., 2004; Ren et al., 2013) and AP-2 in complex with the dileucine signal from CD4 (Kelly et al., 2008) and previous biochemical analyses (Mattera et al., 2011). Protein Data Bank accession codes are indicated. Residues targeted for mutagenesis are highlighted. (B) Yeast three-hybrid (Y3H) analysis of the interaction of the cytosolic domain (residues 1–118) of VAMP4 with wild-type (WT) α1A and the indicated α1A mutants. Growth of transformants on plates lacking histidine (−His) indicates interactions. (C) Incorporation of transgenic α1 constructs into the endogenous AP-1 complex. HeLa cells were transfected with plasmids encoding the indicated WT or mutant α1A, α1B, or α1C isoforms tagged with the myc or HA epitopes. Cell extracts were subjected to immunoprecipitation (IP) with antibody to the myc or HA epitopes, followed by immunoblotting (IB) with antibody to the endogenous γ subunit of AP-1 or to the myc or HA epitopes, as indicated. Untransfected HeLa cells were used as a negative control (−). (D) Confocal immunofluorescence microscopy of rat hippocampal neurons expressing transgenic wild-type or V98S-mutant α1B tagged with the HA epitope. Cells were immunostained for HA (transgenic α1B subunit; green), for the γ subunit of AP-1 or TGN38 (red), and for MAP2 (blue). Rightmost images are merges of the green (α1B-HA) and red (γ-adaptin or TGN38) channels. Scale bar, 10 μm.
MATERIALS AND METHODS

DNA constructs and mutagenesis

The cDNA encoding human σ1A tagged at the C-terminus with three copies of the myc epitope (σ1A-myc) was cloned into pcDNA3.1 (Hygro+; Life Technologies, Grand Island, NY), and those encoding human σ1B or σ1C tagged at the C-terminus with three copies of the HA epitope (σ1B-HA, σ1C-HA) were cloned into pCI-neo (Promega, Madison, WI). The cDNAs encoding the full-length or cytosolic portion of wild-type and mutant cargo proteins were cloned into a modified pHis2 vector encoding MBP (Ren et al., 2013).

Plasmids encoding the recombinant AP-1 core, Arf1Δ1-16-Q71L (referred to as His Arf1Δ-Q71L), and VAMP4-GFP have been described previously (Tran et al., 2007; Guo et al., 2013; Ren et al., 2013). Lamp-1-RFP, plasmid #1817, was obtained from Addgene (submitted by W. Mothes, Yale University School of Medicine, New Haven, CT). Plasmids encoding GFP-ATP7B and TfR-mCherry were gifts from S. Lutsenko (Johns Hopkins University, Baltimore, MD) and J. Lippincott-Schwartz (National Institutes of Health, Bethesda, MD), respectively. Mutagenesis was performed using the QuikChange kit (Agilent, Santa Clara, CA).

Antibodies and other reagents

The antibodies used in this study and their sources are as follows: rabbit anti-MAP2 (Santa Cruz Biotechnology, Dallas, TX); goat anti–ankyrin G (Santa Cruz Biotechnology); chicken anti-GFP (Invitrogen, Carlsbad, CA); mouse anti-TGN38 (Thermo Scientific, Rockford, IL); mouse anti-γ-adaptin (BD Biosciences, San Diego, CA); mouse anti-HA (Covance, Dedham, MA); chicken anti-HA (Millipore, Billerica, MA); chicken anti-MAP2 (Abcam, Cambridge, MA); and mouse anti–pan-neurofascin (extraneuronal epitope, Clone A12/18; University of California, Davis/National Institutes of Health NeuroMab Facility, Bethesda, MD). Rabbit anti-HA and rabbit anti-myc were gifts from A. Sharma (National Institutes of Health). The antibody to p230 was a gift from M. Krieger (MIT, Cambridge, MA). Rabbit anti-GST antiserum and mouse anti-myc clone 9E10 have been described before (Dell’Angelica et al., 1998; Mattera et al., 2011). Mix-n-stain-CF640R (Biotium, Hayward, CA) was used to label the antibody to neurofascin.

FIGURE 5: Binding of a dileucine motif to the σ1 subunit of AP-1 is required for somatodendritic sorting of ATP7B. (A–C) Rat hippocampal neurons coexpressing GFP-tagged ATP7B with (A) myc-tagged WT or V98S-mutant σ1A, (B) HA-tagged WT, or (C) V98S-mutant σ1B, HA-tagged WT, or V98S-mutant σ1C were fixed and immunostained for GFP (negative grayscale images, left), the HA epitope, or the myc epitope (red), MAP2 (blue), and ankyrin G (ANK-G; cyan) as indicated. Cells were imaged by confocal fluorescence microscopy. Cyan arrowheads point to the axon initial segment, and red arrowheads mark the axon. Scale bar, 50 μm.

(D) Quantification of the dendrite/axon polarity index for ATP7B when coexpressed with WT or V98S-mutant σ1B, σ1A, or σ1C, as indicated. Values are the mean ± SD from the indicated number (n) of cells. ****p < 0.0001.
Recombinant protein purification
MBP-ATP7A, -ATP7B, -VAMP4, -Nef, and -TGN38 fusion proteins, Arf1∆1-16-Q71L, and AP-1 core were purified from bacterial lysates using nickel-nitriloacetic acid (Qiagen, Valencia, CA), followed by gel filtration (AKTA; GE Healthcare, Piscataway, NJ; Guo et al., 2013; Ren et al., 2013). In the case of the AP-1 core, affinity purification was performed using glutathione–Sepharose 4B (GE Healthcare) before gel filtration. All purified proteins were snap frozen using liquid nitrogen and stored at −80°C.

Recombinant protein purification

FIGURE 7: Somatodendritic polarity of Tfr is independent of the [DE][XXL][LI]-binding function of σ1. (A, B) Rat hippocampal neurons coexpressing Tfr-mCherry (negative grayscale images, left) with HA-tagged WT (A) or V98S-mutant σ1B (B) were fixed and immunostained with antibody to the HA epitope (green), MAP2 (somatodendritic marker; blue), and ankyrin G (ANK-G, marker for the axon initial segment; cyan) as indicated. Cells were imaged by confocal fluorescence microscopy. Cyan arrowheads point to the axon initial segment, and red arrowheads mark the axon. Scale bar, 50 μm (C) Quantification of the dendrite/axon polarity index for Tfr-mCherry when coexpressed with WT or V98S-mutant σ1B. Values are the mean ± SD from the indicated number (n) of cells. **** p < 0.0001.

FIGURE 6: Somatodendritic polarity of VAMP4 is dependent on a dileucine motif and its binding to the σ1 subunit of AP-1. (A, B) Rat hippocampal neurons expressing WT (A) or LL/AA mutant (B) VAMP4, both tagged with GFP, were fixed and immunostained for GFP (negative grayscale images, left), MAP2 (blue), and ankyrin G (ANK-G) (cyan). (C, D) Rat hippocampal neurons coexpressing VAMP4-GFP with HA-tagged WT (C) or V98S-mutant σ1B (D) were fixed and immunostained for GFP (negative grayscale images, left), the HA epitope (red), MAP2 (blue), and ankyrin G (ANK-G) (cyan), as indicated. Cyan arrowheads point to the axon initial segment, and red arrowheads mark the axon. Scale bar, 50 μm (E) Quantification of VAMP4 polarity. The ratio of fluorescence intensities in dendrites and axons (i.e., polarity index) for WT and LL/AA mutant VAMP4, and of VAMP4 when coexpressed with WT or V98S-mutant σ1B, was calculated. Values are the mean ± SD from the indicated number (n) of cells. **** p < 0.0001.
HeLa cells, followed by immunoprecipitation-immunoblotting as previously described (Guo et al., 2013; Mattera et al., 2014).

Yeast three-hybrid analysis
Yeast three-hybrid analysis was performed as previously described (Mattera et al., 2011). The cytosolic domain of VAMP4 (residues 1–118) and the σ1A subunit of AP-1 were cloned in MCS1 and MCS2, respectively, of the same pBridge vector. The plasmids encoding p53, γ, and SV40 T-Ag were described earlier (Mattera et al., 2011).

Pull-down assay
Interaction of cargo proteins with the AP-1 core in the absence or presence of Arf1Δ1-16-Q71L was analyzed using a previously described pull-down assay (Guo et al., 2013; Ren et al., 2013). In brief, MBP-tagged cargo protein (20 μg) immobilized on amylose resin (New England Biolabs, Beverly, MA) was incubated with 0.05 μM AP-1 core pre-cleared on amylose resin in the absence or presence of 10 μM Arf1Δ1-16-Q71L and 1 mM GTP. The pull down was analyzed by SDS–PAGE, followed by immunoblotting.

Cell culture and transfection
Primary rat hippocampal neurons were prepared as previously described (Kaech and Banker 2006; Farias et al., 2012). Transfections of DNA constructs were performed at day in vitro 3 (DIV3) using Lipofectamine 2000 (Invitrogen), and the transfected neurons were analyzed at DIV10. HeLa cells were maintained in DMEM supplemented with 10% fetal bovine serum, glutamine, and penicillin/streptomycin. Transfections of DNA constructs into HeLa cells were done using Lipofectamine 2000.

Immunostaining, confocal microscopy, and image analysis
Immunostaining and immunofluorescence imaging of transfected hippocampal neurons were performed as previously described (Farias et al., 2012; Mattera et al., 2014). Fluorescence images were acquired using a confocal Zeiss LSM 710 microscope fitted with 40×/1.3 numerical aperture (NA) or 63×/1.4 NA objectives. Analysis of the fluorescence images was done using ImageJ (National Institutes of Health, Bethesda, MD). The polarity index was calculated as previously described (Farias et al., 2012; Mattera et al., 2014).

Immunoprecipitation and immunoblotting
The expression and assembly of dominant-negative σ1 mutants tagged with myc or HA was tested by transient transfection in HEK293T cells. Values are represented as the mean ± SD in all the figures. The statistical analysis was done with a nonparametric t test using Prism (GraphPad Software, La Jolla, CA).
Live-cell imaging and kymographic analysis

Hippocampal neurons were transfected on DIV4 with a plasmid encoding GFP-ATP7B. At DIV8, neurons were immunostained with CF640R-conjugated antibody to neurofascin for 30 min at 37°C to identify the axon. Then control neurons and neurons treated with 100 μM CuSO₄ for 1 h were imaged on a spinning-disk confocal microscope (Marianas; Intelligent Imaging, Denver, CO) fitted with a 63×/1.4 NA objective. Digital images were acquired with an electron-multiplying charge-coupled device camera (Evolve; Photometric, Tucson, AZ). For dual-color videos, GFP-ATP7B and NF-CF640R channels were sequentially exposed for 200 and 100 ms, respectively. Neurons were recorded every 500 ms for 60 s. To analyze the movement of vesicles containing GFP-ATP7B, kymographs were generated using ImageJ, version 1.44o. Lines (20 pixels wide, 50 μm long) were traced in the axon and straightened. This was followed by reslicing stacks and construction of Z-projections. Tracks were orientated so that anterograde movement occurred from left to right in 60 s of recording.

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